# Freshwater monitoring by nanopore sequencing

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## ABSTRACT

Clean freshwater lies at the heart of human society and monitoring its quality is paramount. In addition to chemical controls, traditional microbiological water tests focus on the detection of specific bacterial pathogens. The direct tracing of all aquatic DNA poses a more profound alternative. Yet, this has hitherto been underused due to challenges in cost and logistics. Here we present a simple, fast, inexpensive and comprehensive freshwater diagnostics workflow centred around portable nanopore DNA sequencing. Using defined bacterial compositions and spatiotemporal microbiota from surface water of an example river in Cambridge (UK), our study shows how nanopore sequencing can be readily integrated for the assessment of aquatic bacterial diversity and pollution. We provide a computational benchmark that features more than ten taxonomic classification tools to derive guidelines for bacterial DNA analyses with nanopore data. Through complementary physicochemical measurements, we find that nanopore metagenomics can depict fine temporal gradients along the main hydrological axis of an urban-rural interface and yield high-resolution pathogen maps that address concerns of public health.

## INTRODUCTION

The global assurance of safe drinking water and basic sanitation has been recognised as a United Nations Millennium Development Goal<sup>1</sup>, particularly in light of the pressures of rising urbanisation, agricultural intensification and climate change<sup>2,3</sup>. These trends enforce an increasing demand for freshwater monitoring frameworks that combine cost effectiveness, fast technology deployability and data transparency<sup>4,5</sup>. Environmental metagenomics, the tracing of organisms present in a substrate through high-throughput DNA sequencing, yields informative measures of relative taxonomic species occurrence and functional diversity<sup>6-8</sup>. Microbial metagenomics studies overcome enrichment biases common to traditional culturing approaches<sup>6</sup>; however, they usually depend on expensive and stationary equipment, highly specialised operational training and substantial time lags between sample preparation, raw data generation and access.

In recent years, these challenges have been revisited with the prospect of 'portable' DNA analysis. The main driver of this is the portable, smartphone-sized MinION device from Oxford Nanopore Technologies (ONT), which enables real-time DNA sequencing using nanopores<sup>9</sup>. Nanopore read lengths can be comparably long (currently up to  $\sim 2*10^6$  bases<sup>10</sup>), which is enabled by continuous electrical sensing of sequential nucleotides along single DNA strands. In connection with a laptop or cloud access for the translation of raw voltage signal into nucleotides (basecalling), nanopore sequencing can be used to rapidly monitor long DNA sequences in remote

locations. Although there are still common concerns about the technology's base-level accuracy, mobile MinION setups have already proven powerful for real-time tracing of bacterial and viral pathogen outbreaks<sup>11-16</sup>.

Here we report a simple, low-cost workflow to assess microbial freshwater communities with nanopore DNA sequencing. Our benchmark involves the design and optimisation of essential experimental steps for multiplexed MinION usage in the context of local environments, together with an evaluation of computational methods for the bacterial classification of nanopore sequencing reads from metagenomic libraries. To showcase the resolution of sequencing-based aquatic monitoring in a spatiotemporal setting, we combine DNA analyses with physicochemical measurements of surface water samples collected at nine locations within a confined ~12 kilometre reach of the River Cam passing through the city of Cambridge (UK) in April, June and August 2018.

## RESULTS

#### Experimental design and computational workflows

Nanopore full-length (V1-V9) 16S ribosomal RNA (rRNA) gene sequencing was performed on all locationbarcoded freshwater samples at each of the three time points (Figure 1; Supplementary Table 1a). Samples were complemented with a negative control (deionised water) and a mock community control composed of eight bacterial species in known mixture proportions (Materials and Methods).

To obtain valid taxonomic assignments from freshwater sequencing profiles using nanopore sequencing, 13 different classification tools were compared through several performance metrics (Supplementary Figure 1, Materials and Methods). Root mean square errors (RMSE) between observed and expected bacteria of the mock community differed slightly across all classifiers. An *Enterobacteriaceae* overrepresentation was observed across all replicates and classification methods, pointing towards a consistent *Escherichia coli* amplification bias potentially caused by skewed taxonomic specificities of the selected 16S primer pair (27f and 1492r)<sup>17</sup>. Robust quantifications were obtained by Minimap2<sup>18</sup> alignments against the SILVA 132 database<sup>19</sup>, for which 99.68 % of classified reads aligned to the expected mock community taxa. Minimap2 classifications reached the second lowest RMSE (excluding *Enterobacteriaceae*), and relative quantifications were highly consistent between mock community replicates. Benchmarking of the classification tools on one aquatic sample further confirmed Minimap2's reliable performance, although other tools such as SPINGO<sup>20</sup>, MAPseq<sup>21</sup>, or IDTAXA<sup>22</sup> also

produced highly concordant results despite considerable variations in processing speed and memory usage (data not shown).

## Diversity analysis and river core microbiome

Using Minimap2 classifications within our bioinformatics consensus workflow (Supplementary Figure 2, Materials and Methods), we then inspected sequencing profiles of three independent MinION runs for a total of 30 river DNA isolates and six controls. This yielded ~8.3 million sequences with exclusive barcode assignments (Figure 2a, Supplementary Table 2). Overall, 55.9% (n = 4,644,194) of raw reads could be taxonomically assigned to the family level (Figure 2b). To account for variations in sample sequencing depth, rarefaction with a cut-off at 37,000 reads was applied across all samples. While preserving  $\sim 90\%$  of the original family level taxon richness (Mantel test, R = 0.814,  $p = 2.1*10^4$ ; Supplementary Figure 3), this conservative thresholding resulted in the exclusion of 14 samples, mostly from June, for subsequent high-resolution analyses. The 16 remaining surface water samples revealed moderate levels of microbial heterogeneity (Figure 2b, Supplementary Figure 3): microbial family alpha diversity ranged between 0.46 (June-6) and 0.92 (April-7) (Simpson index), indicating partially low-level evenness with a few taxonomic families that account for the majority of the metagenomic signal. Hierarchical clustering of taxon profiles showed a dominant core microbiome across all aquatic samples (clusters C2 and C4, Figure 2c). The most common bacterial families observed were Burkholderiaceae (40.0 %), Spirosomaceae (17.7 %), and NS11-12 marine group (12.5 %), followed by Arcobacteraceae (4.8 %), Sphingomonadaceae (2.9 %) and Rhodobacteraceae (2.5 %) (Figure 2d). Members of these families are commonly associated with freshwater environments; for example, Burkholderiaceae reads mostly originate from aquatic genera such as Limnohabitans, Rhodoferax or Aquabacterium, which further validates the suitability of this environmental nanopore sequencing workflow.

Hierarchical clustering additionally showed that two biological replicates collected at the same location and time point (April samples 9.1 and 9.2), grouped with high concordance; this indicates that moderate spatiotemporal trends are discernible within a highly localised context. Besides the dominant core microbiome, microbial profiles showed a marked arrangement of time dependence, with water samples from April grouping more distantly to those from June and August (Figure 2c). Principal component analysis (PCA) (Figure 3a, Supplementary Figure 4) revealed that the strongest differential abundances along the chronological axis of variation (PC3) derived from

the higher abundance of *Carnobacteriaceae* and *Saprospiraceae* in April (Figure 3b). These families are known for their occurrence in waters with lower temperature<sup>23</sup> or high organic biomass availability<sup>24,25</sup>, respectively.

## Hydrochemistry and seasonal profile of the River Cam

While a seasonal difference in bacterial composition can be expected due to increasing water temperatures in the summer months, additional changes may have also been caused by alterations in river hydrochemistry and flow rate (Supplementary Figures 5 and 6, Supplementary Table 1c). To assess this effect in detail, we measured the pH and a range of major and trace cations in all river water samples using inductively coupled plasma-optical emission spectroscopy (ICP-OES), as well as major anions using ion chromatography (Supplementary Figure 5, Materials and Methods). Similarly to the bacterial composition dynamics, we observed significant temporal variation in water chemistry, superimposed on a spatial gradient of generally increasing sodium and chloride concentrations along the river reach. This spatially consistent effect is likely attributed to wastewater and agricultural discharge inputs<sup>26</sup> in and around Cambridge city. A comparison of the major element chemistry in the Cam River transect with the world's 60 largest rivers further corroborates the potential impact of anthropogenic pollution in this fluvial ecosystem<sup>27</sup> (Supplementary Figure 5, Materials and Methods).

## Maps of potential bacterial pathogens at species-level resolution

In line with these physicochemical trends, we next determined the spatiotemporal enrichment of potentially functionally important bacterial taxa through nanopore sequencing. We retrieved 58 potentially pathogenic bacterial genera through careful integration of species known to affect human health<sup>28,29</sup>, and also 13 wastewater-associated<sup>30</sup> bacterial genera. Of these, 21 potentially pathogenic and eight wastewater-associated genera were detected across all of the river samples (Figure 3c; Materials and Methods). Many of these signals were stronger downstream of urban river sections, within the mooring zone for recreational and residential barges (location 7) and in the vicinity of sewage outflow from a nearby wastewater treatment plant (location 8). The most prolific candidate pathogen genus observed across all locations was *Arcobacter*, which features multiple species implicated in acute gastrointestinal infections<sup>31</sup>.

In general, much of the taxonomic variation across all samples was caused by the isolate of April-7 (Supplementary Figure 4a-b; PC1 explains 27.6 % of the overall variance in bacterial composition). This was characterised by an unusual dominance of *Caedibacteraceae*, *Halomonadaceae* and others (Supplementary Figure

4c). Isolate April-8 also showed a highly distinct bacterial composition, with some families nearly exclusively occurring in this sample (outlier analysis, Materials and Methods). The most predominant bacteria in this sewage pipe outflow are typically found in wastewater sludge or have been shown to contribute to nutrient pollution from effluents of wastewater plants, such as *Haliangiaceae*, *Nitospiraceae*, *Rhodocyclaceae*, and *Saprospiracea*<sup>30,32</sup> (Figure 3c).

Using multiple sequence alignments between nanopore reads and pathogenic species references, we further resolved the phylogenies of three common potentially pathogenic genera occurring in our river samples, *Pseudomonas, Legionella* and *Salmonella* (Supplementary Figure 7, Materials and Methods). While *Legionella* and *Salmonella* diversities only presented negligible levels of known harmful species, a cluster of sequencing reads in downstream sections indicated a low abundance of the opportunistic, environmental pathogen *Pseudomonas aeruginosa* (Supplementary Figure 7).

We also found varying relative abundances of the *Leptospira* genus, which was recently described to be enriched in wastewater effluents in Germany<sup>33</sup> (Figure 3c). This taxonomic group contains several potentially pathogenic species capable of causing life-threatening leptospirosis through waterborne infections<sup>34</sup>. Yet, the genus also features close-related saprophytic and "intermediate" taxa<sup>35</sup>. To resolve its complex phylogeny in the River Cam surface, we aligned *Leptospiraceae* reads from all samples together with various reference sequences assigned to pre-classified pathogenic, saprophytic and other environmental *Leptospira* species<sup>35</sup> (Figure 3d; Materials and Methods). Despite the presence of nanopore homopolymer sequencing errors (Supplementary Figure 8) and correspondingly inflated divergence between reads, we could pinpoint spatial clusters and a distinctly higher similarity between our *Leptospiraceae* amplicons and saprophytic rather than pathogenic *Leptospira* species. These findings were subsequently validated by targeted, *Leptospira* species-specific qPCR (Materials and Methods), confirming that the current nanopore sequencing quality is sufficiently high to yield indicative results for bacterial monitoring workflows at the bacterial species level.

## DISCUSSION

Using a low-cost, easily adaptable and scalable framework, we provide the first detailed nanopore sequencing atlas of bacterial microbiota along a river reach. Beyond the core microbiome of an exemplary fluvial ecosystem, our results suggest that it is possible to robustly assess time changes in accessory bacterial composition in the

context of supporting physical (temperature, flow rate) and hydrochemical (pH, inorganic solutes) parameters. We show that nanopore sequencing can identify human pathogen community shifts along rural-to-urban transitions within a river reach, as illustrated by downstream increases in the abundance of pathogen candidates.

Furthermore, our assessment of popular bioinformatics workflows for taxonomic classification highlights current challenges with error-prone nanopore sequences. We observed differences in terms of taxonomic quantifications, read misclassification rates and consensus agreements between the 13 tested computational methods. In this benchmark, using the SILVA 132 reference database, one of the most balanced performances was achieved with Minimap2 alignments. As nanopore sequencing quality continues to increase through refined pore chemistries and consensus sequencing workflows<sup>36,37</sup>, future bacterial taxonomic classifications are likely to improve as well.

We show that nanopore amplicon sequencing data can resolve the core microbiome of a freshwater body, as well as its temporal and spatial fluctuations. Asides common freshwater bacteria, we find that the differential abundances of *Carnobacteriaceae* and *Saprospiraceae* most strongly contribute to seasonal loadings in the Cam River. As *Carnobacteriaceae* have been associated with cold environments<sup>23</sup>, and we found these to be more abundant in colder April samples (mean 11.3 °C, vs. 15.8 °C in June and 19.1 °C in August), this might further establish the impact of water temperature on the variation of bacterial composition. *Saprospiraceae* are frequently observed in wastewater treatment systems<sup>30,32</sup>, where they likely play a role in heterotrophic polymeric degradation<sup>24,25</sup>. The majority of our *Saprospiraceae* reads (~67.33 %) could indeed be assigned to sewage effluent (outlier sample April-8, Figure 3a), which suggests that this particular sample augments the observed time pattern.

Most routine freshwater surveillance frameworks focus on semi-quantitative diagnostics of only a limited number of target taxa, such as pathogenic *Salmonella, Legionella* and faecal coliforms<sup>38</sup>. While common culture-based, immunological or PCR-based approaches can assist stakeholders with limited assessments of local water quality<sup>39,40</sup>, we show that portable nanopore metagenomics offers the promise of more comprehensive microbial pathogen examinations at similar expense. Our analyses highlight that the combination of full-length 16S rRNA gene amplification and nanopore sequencing can complement hydrochemical controls in pinpointing potentially contaminated sites, some of which had been previously highlighted for their pathogen diversity and abundance of antimicrobial resistance genes<sup>41,42</sup>. Nanopore sequencing allowed for the reliable distinction of closely related pathogenic and non-pathogenic bacterial species of the common *Salomella, Legionella, Pseudomonas* and

*Leptospira*, and future bioinformatics efforts might focus on the automatisation of such assessments across more diverse genera of interest.

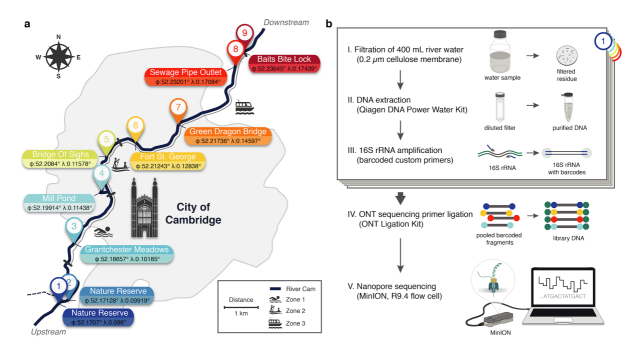
A number of experimental intricacies should be addressed towards future nanopore freshwater sequencing studies, mostly by scrutinising water DNA extraction yields, PCR biases and molar imbalances in barcode multiplexing (Figure 2a, Supplementary Figure 8). Yet, our results show that it would be theoretically feasible to obtain meaningful river microbiota from >100 barcoded samples on a single nanopore flow cell, thereby enabling water monitoring projects involving large collections at a sub-£20 cost per sample (Supplementary Table 3). Barcoded shotgun nanopore sequencing protocols may pose a viable alternative strategy to bypass pitfalls often observed in amplicon-based workflows, namely taxon-specific primer biases<sup>17</sup>, 16S rDNA copy number fluctuations between species<sup>43</sup> and the omission of functionally relevant sequence elements. This could moreover also allow for the monitoring of eukaryotic microorganisms and viruses, when combined with sampling protocol adjustments.

Since the commercial launch of the MinION in 2015, a wide set of nanopore sequencing applications like rRNA gene<sup>44-47</sup> and shotgun metagenomics<sup>48-50</sup> have attracted the interest of a growing user community. Although it is to be expected that short-read metagenomics technology continues to provide valuable environmental insights, as shown through recent cataloguing efforts for world's ocean<sup>51</sup> and soil<sup>52</sup> microbiomes, these traditional platforms are cumbersome for monitoring remote environments or low-resource settings. The MinION technology is considerably less challenging to transport, operate and maintain, and our results how that spatiotemporal nanopore sequencing could be readily adapted for multiplexed bacterial pathogen tracing in epidemic contexts. We reason that the low investment costs (Supplementary Table 3), the convenience of MinION handling and data analysis will allow for such endeavours to become increasingly accessible to citizens and public health organisations around the world, ultimately democratising the opportunities and benefits of DNA sequencing.

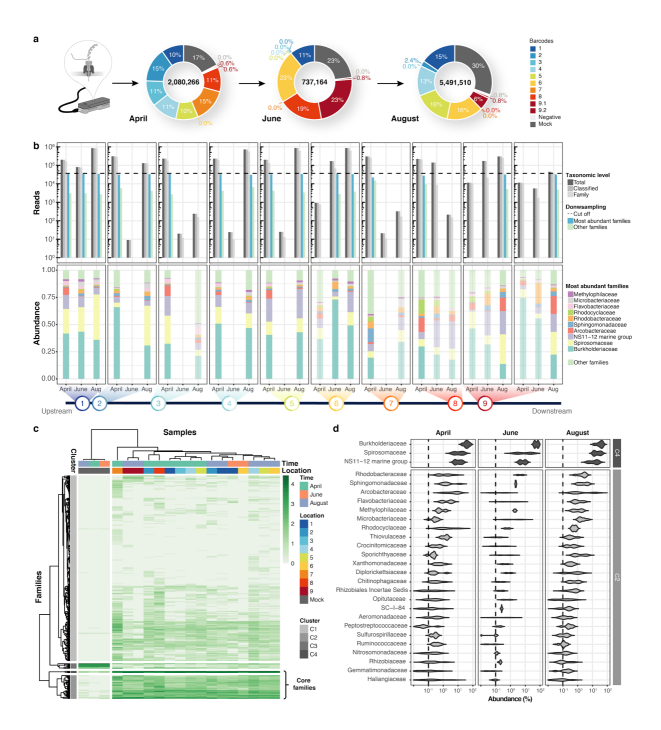
Acknowledgments: We thank Meltem Gürel, Christian Schwall, Jack Monahan, Eirini Vamva and Astrid Wendler for fruitful discussions and water sample collection; Ben Wagstaff, Elliot Brooks, Jennifer Pratscher and Rob Field for providing us with initial test samples; David Seilly and Mervyn Greaves for experimental support with water filtrations and ICP-OES; Tim Brooks and Daniel Bailey from Public Health England for validation tests on *Leptospiraceae*; Jenny Molloy and Michal Filus for help with funding applications; Aleix Lafita, Oana Stroe, Abigail Wood, Paul Saary, Jane Clarke, Fiona Gilsenan and her family for support in public outreach events; Nick Loman, Zamin Iqbal, Rob Finn, Alex Greenwood, Daniela Numberger, Julian Parkhill, Simon Frost, Sam Stubbs, Mark Holmes, Alicja Dabrowska, Alex Patto, Adrien Leger and Kim Judge for advice on nanopore sequencing and environmental pathogen monitoring; Alina Ham (Oxford Nanopore Technologies), Heather Martinez and Gemma Gambrill (Qiagen) for technical support; Víctor de Lorenzo, David Sargan and Lisa Schmunk for helpful comments on the manuscript; Lilo and Manfred Fuchs from the Fuchs Fund for generously supporting LU's conference participation and presentation; Alejandro de Miquel Bleier for statistical advice. Last, we wish to thank our generous supervisors for enabling us to pursue this project in addition to our PhD studies.

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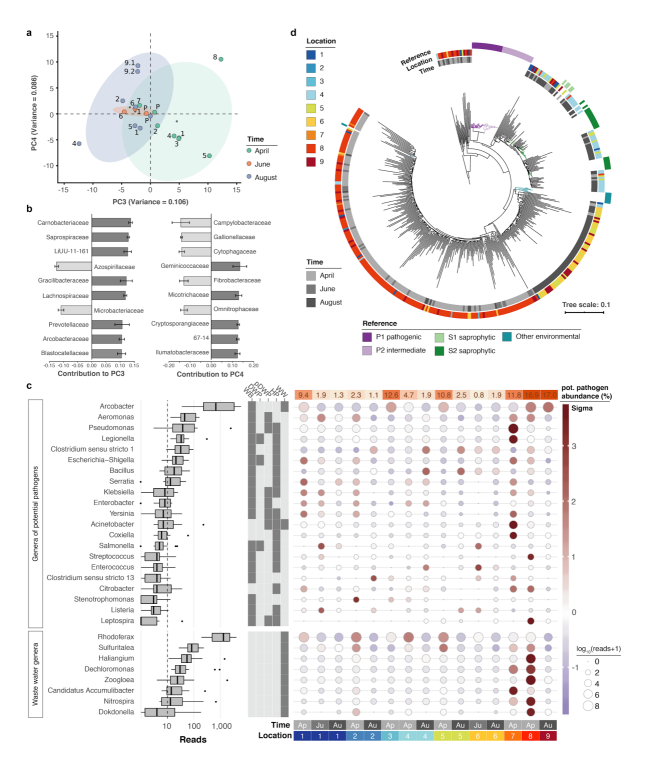


**Figure 1: Freshwater microbiome study design and experimental workflow.** (a) Schematic map of Cambridge (UK) illustrating sampling locations (colour-coded) along the Cam River. Latitude and longitude geographic coordinates are expressed as decimal fractions referring to the global positioning system. (b) Experimental workflow to monitor bacterial communities from freshwater samples using nanopore sequencing (Materials and Methods).



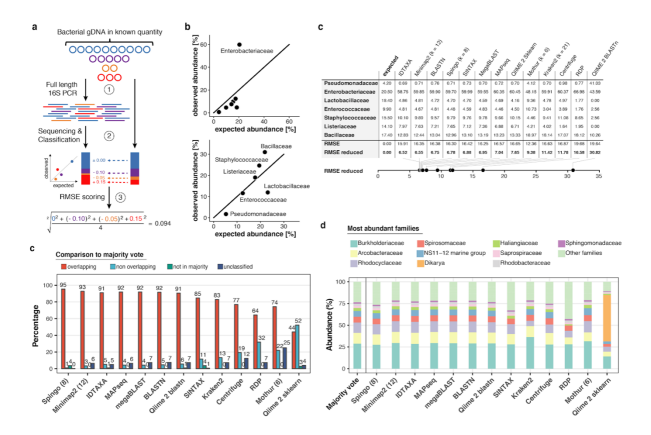
**Figure 2: Bacterial diversity of the River Cam.** (a) Nanopore sequencing output summary. Values within pie charts depict total numbers of classified nanopore sequences per time point. Percentages illustrate representational fractions of locations and control barcodes (negative control and mock community). (b) Read depth and bacterial classification summary. Upper bar plot shows the total number of reads, the number of reads classified to any taxonomic level, to bacterial family level or above, to the ten most abundant bacterial families across all samples or other families. Rarefaction cut-off displayed at 37,000 reads (dashed line). Lower bar plot features fractions of the ten most abundant bacterial families across all samples with less than 37,000 reads are set to transparent. (c) Hierarchical clustering of all bacterial family abundances in

freshwater samples after rarefaction, together with the mock community control. Four major clusters of bacterial families occur, with two of these (C2 and C4) corresponding to the core microbiome of ubiquitously abundant families, one (C3) to the main mock community families and one (C1) to the majority of rarer accessory taxa. (d) Detailed river core microbiome. Violin plots (log<sub>10</sub>) summarise fractional representation of bacterial families from clusters C2 and C4 across the three sampling time points, sorted by median total abundance. Vertical dashed line depicts 0.1 % proportion.

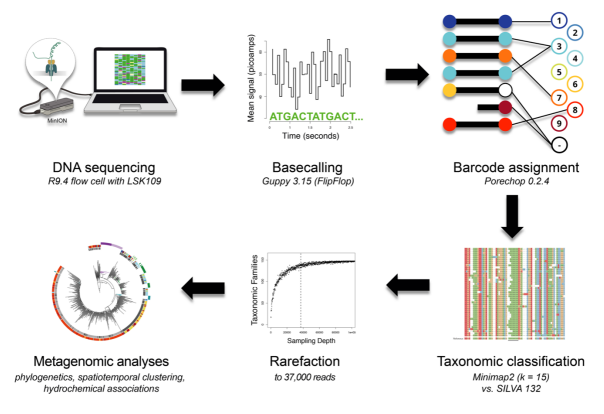


**Figure 3: Rare taxa and potential pathogens of the River Cam.** (a) Principal component analysis indicating community dissimilarities along the main time (PC3) and spatial (PC4) axes of variation. Numbers and coloured dots indicate locations for each time point. (b) Contribution of individual bacterial families to the PCs in (a). (c) Diversity, abundance and distribution of potentially pathogenic bacteria and wastewater treatment related bacteria, at genus level resolution. Species from subsets of genera are categorised as waterborne bacterial pathogens (WB), drinking water pathogens (DWP), potential drinking water pathogens (pDWP), human pathogens (HP) and core

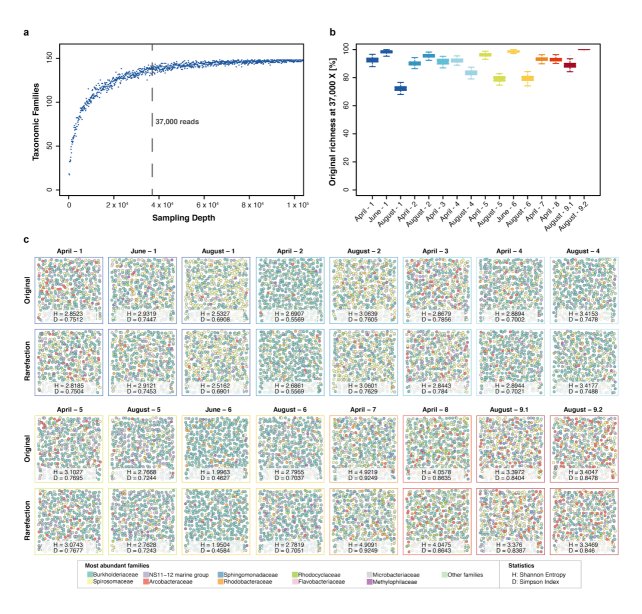
genera from wastewater treatment plants (WW). Circle sizes represent overall read size fractions, while circle colours (sigma scheme) represent the deviation from the observed mean relative abundance within each genus. (d) Phylogenetic tree illustrating the multiple sequence alignment of all river nanopore reads classified as *Leptospira*, together with known *Leptospira* reference sequences ranging from pathogenic to saprophytic species<sup>35</sup>. Branch length tree scale indicates misalignment rates and thus, indirectly, sequencing error rate.



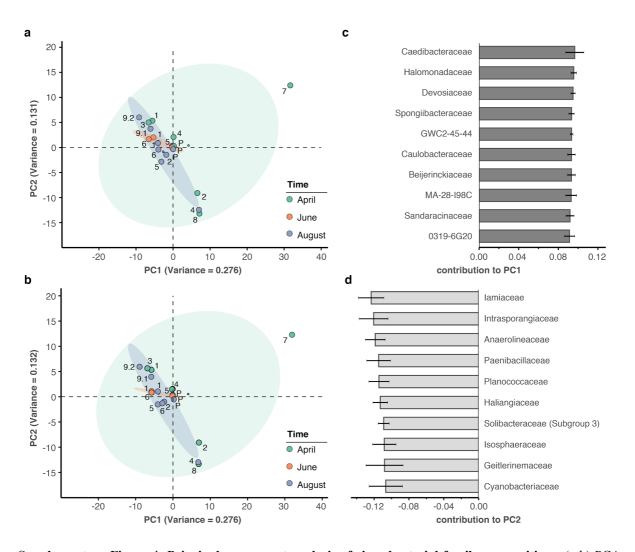
**Supplementary Figure 1: Benchmarking of classification tools with nanopore full-length 16S reads.** (a) Schematic of mock community quantification performance testing. (b) Observed vs. expected read fraction of bacterial families present in 10,000 nanopore reads randomly drawn from mock community sequencing data. Example representation of Minimap2 (kmer length 12) quantifications with (upper) and without (lower) *Enterobacteriaceae* (Materials and Methods). (c) Mock community classification output summary for 13 classification tools tested against the 10,000 nanopore reads. Root mean squared errors observed and expected bacterial read fractions are provided with (RMSE) and without *Enterobacteriaceae* (RMSE reduced). (d) Classification output summary for 10,000 reads randomly drawn from an example freshwater sample (Materials and Methods). 'Overlapping' fractions (red) represent agreements of a classification tool with the majority of tested methods on the same reads, while 'non-overlapping' fractions (light blue) represent disagreements. Green sets highlight rare taxon assignments not featured in any of the 10,000 majority classifications, while dark blue bars show unclassified read fractions. (d) Top 10 represented bacterial taxon families across all 13 classifiers based on the 10,000 reads used in (c).



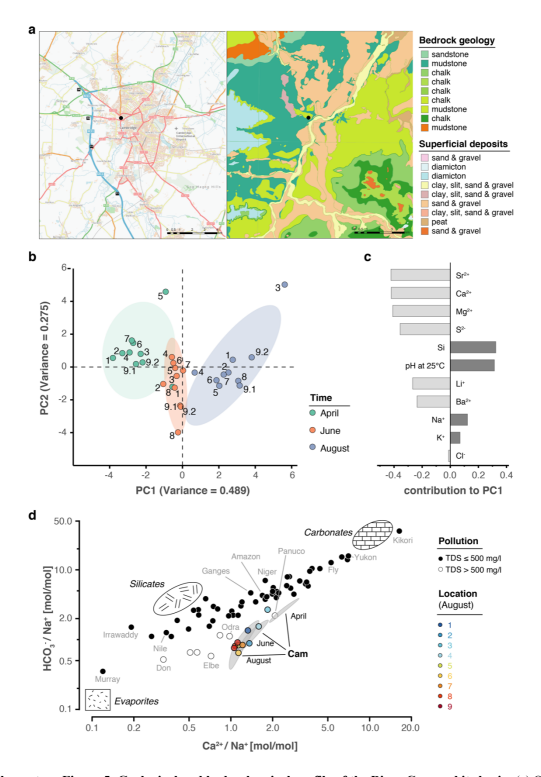
**Supplementary Figure 2: Bioinformatics consensus workflow.** Essential data processing steps, from nanopore sequencing to spatiotemporal bacterial composition analysis (Materials and Methods). After full-length 16S rDNA sequencing with the MinION (R9.4 flow cell), local basecalling of the raw fast5 files was performed using Guppy<sup>53</sup>. Output fastq files were filtered for length and quality (Materials and Methods), and reads assigned to their location barcode using Porechop. We then used Minimap2<sup>18</sup> and the SILVA 132 database<sup>19</sup> for taxonomic classifications. Rarefaction reduced each sample to the same number of reads (37,000), allowing for a robust comparison of bacterial composition across samples in various downstream analyses.



Supplementary Figure 3: Impact of rarefaction on diversity estimation. (a) Example rarefaction curve for bacterial family classifications of the 'April-1' sample. The chosen cut-off preserves most (~90 %) of the original family taxon richness (vertical line). (b) Difference between original and rarefied family richness at 37,000 reads across all freshwater sequencing runs with quantitative sequencing outputs above the chosen cut-off. Boxplots feature 100 independent rarefactions per sample. Error bars represent Q1 - 1.5\*IQR (lower), and Q3 + 1.5\*IQR (upper), respectively; Q1: first quartile, Q3: third quartile, IQR: interquartile range. (c) Diversity visualisation of the ten most abundant bacterial families across all samples with sequencing outputs >37,000 reads, through 400 "unordered bubbles". Taxonomic proportions and colours are in accordance with Figure 2b. Shannon (H) and Simpson (D) indices for all samples indicate marginal differences between pairs of original and rarefied sets.

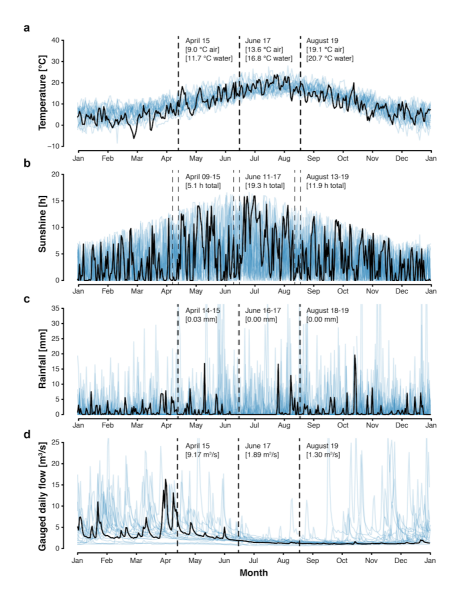


**Supplementary Figure 4:** Principal component analysis of river bacterial family compositions. (a-b) PCA with two independent rarefaction sets to 37,000 reads in all freshwater sequencing samples. Numbers and coloured dots indicate locations for each time point. The first and second principal components (PC1 and PC2, combined variance: ~41 %) robustly capture outlier samples 'April-7' along PC1 and 'April-2', 'August-4' and 'April-8' along PC2. (c-d) Fractional loads of the ten bacterial families most strongly contributing to changes along PC1 (c) and along PC2 (d). Error bars represent standard deviation of these families to the respective PC across four independent rarefactions. Subsequent principal components (PC3 and PC4) are less outlier-driven and depict spatial and temporal metagenomic trends within the River Cam.

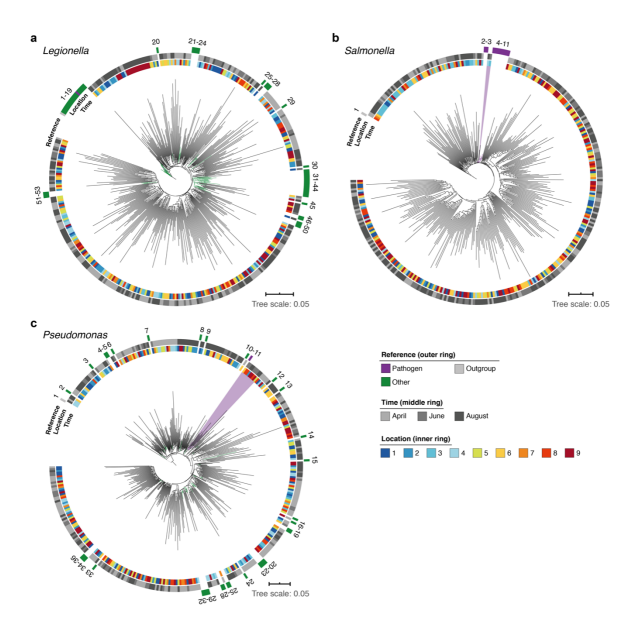


**Supplementary Figure 5: Geological and hydrochemical profile of the River Cam and its basin.** (a) Outline of the Cam River catchment surrounding Cambridge (UK), and its corresponding lithology. Overlay of bedrock geology and superficial deposits (British Geological Survey data: DiGMapGB-50, 1:50,000 scale) is shown as visualised by GeoIndex. Bedrock is mostly composed of subtypes of Cretaceous limestone (chalk), gault (clay, sand) and mudstone. (b) Principal component analysis of measured pH and 13 inorganic solute concentrations of this study's 30 river surface water samples. PC1 (~49 % variance) displays a strong, continuous temporal shift in

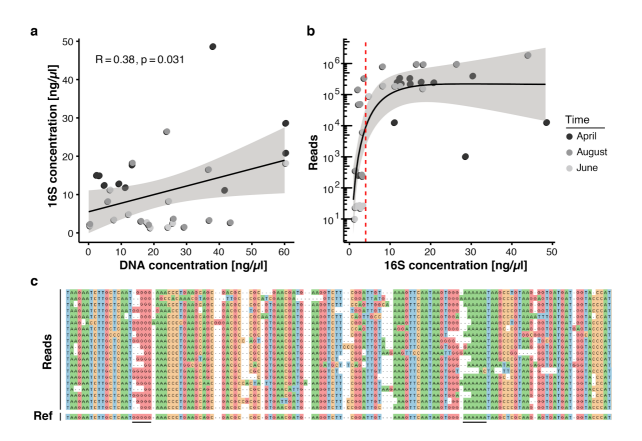
hydrochemistry. (c) Parameter contributions to PC1 in (b), highlighting a reduction in water hardness (Ca<sup>2+</sup>, Mg<sup>2+</sup>) and increase in pH towards the summer months (June and August). (d) Mixing diagram with Na<sup>+</sup>-normalised molar ratios, representing inorganic chemistry loads of world's 60 largest rivers<sup>27</sup>; open circles represent polluted rivers with total dissolved solid (TDS) concentrations >500 mg l<sup>-1</sup>. Cam River ratios are superimposed as ellipses from ten samples per month (50 % confidence, respectively). Separate data points for all samples from August are also shown and colour-coded, indicating the downstream-to-upstream trend of Na<sup>+</sup> increase (also observed in April and June). End-member signatures show typical chemistry of small rivers draining these lithologies exclusively (carbonate, silicate and evaporite).



Supplementary Figure 6: Cambridge weather and Cam River flow rate. (a) Daily air temperature [°C], (b) daily sunshine [hours], and (c) daily rainfall [mm] of Cambridge in 2018 (black trend line) vs. 1998-2017 (blue background trend lines). (d) Cam River gauged daily flow [m<sup>3</sup>s<sup>-1</sup>] in 2018 (black trend line) vs. 1968-2017 (blue background trend lines). Data was compiled from public repositories https://www.cl.cam.ac.uk/research/dtg/weather/ and https://nrfa.ceh.ac.uk/. Gauged daily flow measurements at Jesus Lock, Cambridge (between sampling locations 5 and 6; NRFA #33016) were discontinued in 1983. Yet, contemporary flow rates can be modelled with high accuracy (Pearson's R = 0.9,  $R^2 = 0.8$ ) through linear data integration of three upstream stations already in operation since before 1983: Rhee at Wimpole (NRFA #33027, 70.2 % model weight), Granta at Stapleford (NRFA #33053, 19.6 % model weight) and Cam at Dernford (NRFA #33024, 10.3 % model weight).



Supplementary Figure 7: Phylogenetic clustering of candidate pathogenic bacterial genera in the river Cam. Phylogenetic trees illustrating multiple sequence alignments of exemplary River Cam nanopore reads classified as (a) *Legionella*, (b) *Salmonella* or (c) *Pseudomonas*, together with known reference species sequences ranging from pathogenic to saprophytic taxa within the genes (Table S7b-d). Reads highlighted in light violet background display close clustering with pathogenic isolates of (b) *Salmonella spp.* and (c) *Pseudomonas aeruginosa*.



Supplementary Figure 8: Key challenges of environmental monitoring with nanopore sequencing. (a-b) Correlation analysis between DNA extraction yield, 16S amplification yield and raw sequencing output (Supplementary Table 2). (a) DNA concentrations (x-axis) obtained from 30 freshwater samples after extraction with the DNeasy PowerWater Kit (Materials and Methods) are compared against the DNA concentration of the same samples after full-length 16S PCR amplification (y-axis), as measured by Qubit dsDNA HS. Fitted linear model displays the 95 % confidence interval (R = Pearson correlation coefficient). (b) The DNA concentration obtained for each sample after full-length 16S PCR amplification (x-axis) is compared against the final number of demultiplexed nanopore sequencing reads. Logarithmic fit with 95 % confidence interval indicates that samples with a minimum input concentration measurement of ~5 ng/µl yielded sequencing outputs sufficient to pass the rarefaction threshold of 37,000 reads. (c) Multiple sequence alignment of an example set of related nanopore 16S sequences, displaying increased indel rates at homopolymer reference sites (underlined).

#### MATERIALS AND METHODS

#### **1.1 Freshwater Sampling**

We monitored nine distinct locations along a 11.62 km reach of the River Cam, featuring sites upstream, downstream and within the urban belt of the city of Cambridge, UK. Measurements were taken at three time points, in two-month intervals between April and August 2018 (Figure 1, Supplementary Table 1a). To warrant river base flow conditions and minimise rain-derived biases, a minimum dry weather time span of 48h was maintained prior to sampling<sup>54</sup>. One litre of surface water was collected in autoclaved DURAN bottles (Thermo Fisher Scientific, Waltham, MA, USA), and cooled to 4 °C within three hours. Two bottles of water were collected consecutively for each time point, serving as biological replicates of location 9 (samples 9.1 and 9.2).

#### 1.2 Physical and Chemical Metadata

We assessed various chemical, geological and physical properties of the River Cam (Supplementary Figures 5 and 6, Supplementary Tables 1b and 1c).

*In situ* water temperature was measured immediately after sampling. To this end, we linked a DS18B20 digital temperature sensor to a portable custom-built, grid mounted Arduino nano v3.0 system. The pH was later recorded under temperature-controlled laboratory conditions, using a pH edge electrode (HI-11311, Hanna Instruments, Woodsocket, RI, USA).

To assess the dissolved ion concentrations in all collected water samples, we aerated the samples for 30 seconds and filtered them individually through a 0.22  $\mu$ M pore-sized Millex-GP polyethersulfone syringe filter (MilliporeSigma, Burlington, MA, USA). Samples were then acidified to pH ~2, by adding 20  $\mu$ L of 7M distilled HNO<sub>3</sub> per 3 mL sample. Inductively coupled plasma-optical emission spectroscopy (ICP-OES, Agilent 5100 SVDV; Agilent Technologies, Santa Clara, CA, USA) was used to analyse the dissolved cations Na<sup>+</sup>, K<sup>+</sup>, Ca<sup>2+</sup>, Mg<sup>2+</sup>, Ba<sup>2+</sup>, Li<sup>+</sup>, as well as Si and SO4<sup>2-</sup> (as total S) (Supplementary Table 1b). International water reference materials (SLRS-5 and SPS-SW2) were interspersed with the samples, reproducing certified values within 10 % for all analysed elements. Chloride concentrations were separately measured on 1 mL of non-acidified aliquots of the same samples, using a Dionex ICS-3000 ion chromatograph (Thermo Fisher Scientific, Waltham, MA, USA) (Supplementary Table 1b). Long-term repeat measurements of a USGS natural river water standard T-143 indicated precision of more than 4 % for Cl. However, the high Cl<sup>-</sup> concentrations of the samples in this study were not fully bracketed by the calibration curve and we therefore assigned a more conservative uncertainty of 10 % to Cl concentrations.

High calcium and magnesium concentrations were recorded across all samples, in line with hard groundwater and natural weathering of the Cretaceous limestone bedrock underlying the river catchment (Supplementary Figure 5). There are no known evaporite salt deposits in the river catchment, and therefore the high dissolved Na<sup>+</sup>, K<sup>+</sup> and Cl<sup>-</sup> concentrations in the Cam are likely derived from anthropogenic inputs<sup>26</sup> (Supplementary Figure 5). We calculated bicarbonate concentrations through a charge balance equation (concentrations in mol/L):

 $conc (HCO_{3}^{-}) = conc (Li^{+}) + conc (Na^{+}) + conc (K^{+}) + 2*conc (Mg^{2+}) + 2*conc (Ca^{2+}) - conc (Cl^{-}) - 2*conc (S^{2-}) + 2*conc (S^{2-}$ 

The total dissolved solid (TDS) concentration across the 30 freshwater samples had a mean of 458 mg/L (range 325 - 605 mg/L) which is relatively high compared to most rivers, due to 1.) substantial solute load in the Chalk groundwater (particularly Ca<sup>2+</sup>, Mg<sup>2+</sup>, and HCO<sub>3</sub><sup>-</sup>) and 2.) likely anthropogenic contamination (particularly Na<sup>+</sup>, Cl<sup>-</sup>, and SO4<sup>2-</sup>). The TDS range and the major ion signature of the Cam is similar to other anthropogenically heavily-impacted rivers<sup>27</sup>, exhibiting enrichment in Na<sup>+</sup> (Supplementary Figure 5).

Overall, ion profiles clustered substantially between the three time points, indicating characteristic temporal shifts in water chemistry. PC1 of a PCA on the solute concentrations [µmol/L] shows a strong time effect, separating spring (April) from summer (June, August) samples (Supplementary Figure 5b). We highlighted the 10 most important features (i.e., features with the largest weights) and their contributions to PC1 (Supplementary Figure 5c).

We integrated sensor data sets on mean daily air temperature, sunshine hours and total rainfall from a public, Cambridge-based weather station (Supplementary Figure 6a-c; Supplementary Table 1c). Similarly, mean gauged daily Cam water discharge [m<sup>3</sup>s<sup>-1</sup>] was retrieved through publicly available records from three upstream gauging stations connected to the UK National River Flow Archive (https://nrfa.ceh.ac.uk/), together with historic measurements from 1968 onwards (Supplementary Figure 6d)

## **1.3 DNA Extraction**

Within 24 hours of sampling, 400 mL of chilled freshwater from each site was filtered through an individual 0.22 µm pore-sized nitrocellulose filter (MilliporeSigma, Burlington, MA, USA) placed on a Nalgene polysulfone bottle top filtration holder (Thermo Fisher Scientific) at -30 mbar vacuum pressure. Additionally, 400 mL deionised (DI) water was also filtered. We then performed DNA extractions with a modified DNeasy PowerWater protocol (Qiagen, Hilden, Germany). Briefly, filters were cut into small slices with sterile scissors and transferred to 2 mL Eppendorf tubes containing lysis beads. Homogenization buffer PW1 was added, and the tubes subjected to ten minutes of vigorous shaking at 30 Hz in a TissueLyser II machine (Qiagen). After subsequent DNA binding and washing steps in accordance with the manufacturer's protocol, elution was done in 50 µL EB. We used Qubit dsDNA HS Assay (Thermo Fisher Scientific) to determine water DNA isolate concentrations (Supplementary Table 2a).

### 1.4 Bacterial Full-Length 16S rDNA Sequence Amplification

DNA extracts from each sampling batch and DI water control were separately amplified with V1-V9 full-length (~1.45 kbp) 16S rDNA gene primers, and respectively multiplexed with an additional sample with a defined bacterial mixture composition of eight species (*Pseudomonas aeruginosa, Escherichia coli, Salmonella enterica, Lactobacillus fermentum, Enterococcus faecalis, Staphylococcus aureus, Listeria monocytogenes, Bacillus subtilis*; D6305, Zymo Research, Irvine, CA, USA) (Supplementary Figure 1b-c), which was previously assessed using nanopore shotgun metagenomics<sup>48</sup>. We used common primer binding sequences<sup>17</sup> 27f and 1492r, both coupled to unique 24 bp barcodes and a nanopore motor protein tether sequence (Supplementary Table 4). Full-length 16S PCRs were performed with the following conditions:

30.8 μL DI water
6.0 μL barcoded primer pair (10 μM)
5.0 μL PCR-buffer with MgCl<sub>2</sub> (10x)
5.0 μL dNTP mix (10x)
3.0 μL freshwater DNA extract
0.2 μL Taq (Qiagen)

94 °C - 2 minutes

94 °C - 30 seconds, 60 °C - 30 seconds, 72 °C - 45 seconds (35 cycles)

72 °C - 5 minutes

## **1.5 Nanopore Library Preparation**

Amplicons were purified from reaction mixes with a QIAquick purification kit (Qiagen). Two rounds of alcoholic washing and two additional minutes of drying at room temperature were then performed, prior to elution in 30 µL 10 mM Tris-HCl pH 8.0 with 50 mM NaCl. After concentration measurements with Qubit dsDNA HS, twelve barcoded extracts of a given batch were pooled in equimolar ratios, to approximately 300 ng DNA total (Supplementary Table S2b). We used KAPA Pure Beads (KAPA Biosystems, Wilmington, MA, USA) to concentrate full-length 16S products in 21 µL DI water. Multiplexed nanopore ligation sequencing libraries were then made by following the SQK-LSK109 protocol (Oxford Nanopore Technologies, Oxford, UK).

### 1.6 Nanopore Sequencing

R9.4 MinION flow cells (Oxford Nanopore Technologies) were loaded with 75  $\mu$ l of ligation library. The MinION instrument was run for approximately 48 hours, until no further sequencing reads were collected. Fast5 files were basecalled using Guppy (version 3.15) and output DNA sequence reads with Q>7 were saved in fastq files. Various output metrics per library and barcode are summarised in Supplementary Table 2c.

## 1.7 Leptospira Validation

In collaboration with Public Health England, raw Cam River water DNA isolates from each location and time point were subjected to the UK reference service for leptospiral testing. This is based on quantitative real-time PCR (qPCR) of 16S rDNA and *LipL32*, implemented as a TaqMan assay for the detection and differentiation of pathogenic and non-pathogenic *Leptospira* spp. from human serum. Briefly, the assay consists of a two-component PCR; the first component is a duplex assay that targets the gene encoding the outer membrane lipoprotein *LipL32*, which is reported to be strongly associated with the pathogenic phenotype. The second reaction is a triplex assay targeting a well conserved region within the 16S rRNA gene (*rrn*) in *Leptospira* spp. Three different genomic variations correlate with pathogenic (PATH probe), intermediate (i.e., those with uncertain pathogenicity in humans; INTER probe) and non-pathogenic *Leptospira* spp. (ENVIRO probe), respectively.

#### 2. DNA Sequence Processing Workflow

Data processing and read classification was developed using the Snakemake workflow management system<sup>55</sup> and is available on Github - together with all necessary downstream analysis scripts to reproduce the results of this manuscript (https://github.com/d-j-k/puntseq). De-multiplexed and processed reads, separated by month and location, are available through the European Nucleotide Archive (PRJEB34900).

### 2.1 Read Data Processing

Reads were demultiplexed and adapters trimmed using Porechop (version 0.2.4, https://github.com/rrwick/porechop). The only non-default parameter changed was '--check\_reads' (set to 50,000), to increase the subset of reads to search for adapter sets. Next, we removed all reads shorter than 1.4 kbp and longer than 1.6 kbp with Nanofilt (version 2.5.0).

We gathered read statistics such as quality scores and read lengths using NanoStat (version 1.1.2, https://github.com/wdecoster/nanostat), and used Pistis (https://github.com/mbhall88/pistis) to create quality control plots. This allowed us to assess GC content and Phred quality score distributions, which appeared consistent across and within our reads. Overall, we obtained 2,080,266 reads for April, 737,164 for June, and 5,491,510 for August, with a mean read quality of 10.0 (Supplementary Table 2c).

## 2.2 Benchmarking of Bacterial Taxonomic Classifiers using Nanopore Reads

We used 13 different computational tools for bacterial full-length 16S rDNA sequencing read classification (section 2.2.1).:

Tool	Version	Commands
BLASTN <sup>56</sup>	v.2.9.0+	<pre># build database makeblastdb -in silva.fna -parse_seqids -blastdb_version 5 -title "2019-08-24_SILVA_BLASTdatabase" -dbtype nucl # run BLASTN blastn -db silva.fna -query Cam16S.fa -out Cam16S.out -outfmt '6'</pre>
Centrifuge <sup>57</sup>	v.1.0.4	<pre># build database centrifuge -x centrifuge_16s_database -U Cam16S.fathreads config["centrifuge_16s"]["threads"]report-file Cam16S_report.tsv -S Cam16S.tabmet-stderr centrifuge-kreport -x centrifuge_16s_database Cam16S.tab {input} &gt; Cam16S.kreport</pre>

IDTAXA <sup>22</sup>	Implemented in R DECIPHER v.2.10.2	load("SILVA_SSU_r132_March2018.RData") IdTaxa(Cam16S.fa, trainingSet, strand = "both", threshold = 0)
Kraken2 <sup>58</sup>	v.2.0.7	# build database kraken2db kraken2_16s_databaseoutput Cam16S,out report Cam16S.kreportgzip-compressedthreads 1 Cam16S.fa
MAPseq <sup>21</sup>	v.1.2.3	mapseq Cam16S.fa silva_ref.fa > Cam16S.mseq
MegaBLAST <sup>59</sup>	v.2.9.0+	<pre># build database makeblastdb -in silva.fna -parse_seqids -blastdb_version 5 -title "2019-08-24_SILVA_BLASTdatabase" -dbtype nucl # run megaBLAST blastn -task "megablast" -db silva.fna -query Cam16S.fa -out Cam16S.out -outfmt '6'</pre>
Minimap2 <sup>18</sup>	v.2.13-r852-dirty	minimap2 -k 15 -d silva_k15.mmi silva.fna minimap2 -ax map-ont -L silva_k15.mmi Cam16S.fa > Cam16S.sam
Mothur <sup>60</sup>	v.1.43.0	align.seqs(candidate=Cam16S.fa, template=mothur.silva.nr_v132.align, processors=1, ksize=6, align=needleman)
QIIME 2 blastn <sup>61</sup>	v.2019.7	qiime feature-classifier classify-consensus-blasti-query Cam16S.qzai-reference-reads silva.qzai-reference- taxonomy silva_tax.qzao-classification Cam16S.qza output-dir /Qiime2/Cam16S_blastn
QIIME 2 sklearn <sup>61</sup>	v.2019.7	qiime feature-classifier classify-sklearni-reads Cam16S.qza - -i-classifier silva-132-99-nb-classifier.qzao-classification Cam16S.qzap-n-jobs 8output-dir /Qiime2/Cam16S _sklearn
RDP <sup>62</sup>	Implemented in R DADA2 v.1.12.1 <sup>63</sup>	assignTaxonomy(seqs = Cam16S.fa, refFasta = silva_nr_v132_train_set.fa.gz", tryRC = T, outputBootstraps=T,minBoot=0)
SINTAX <sup>64</sup>	Implemented in VSEARCH v.2.13.3 <sup>65</sup>	vsearch -makeudb_usearch silva_tax.fa -output silva_tax.udb vsearch -sintax Cam16S.fa -db silva_tax.udb -tabbedout Cam16S.sintax -strand both -sintax_cutoff 0.5
SPINGO <sup>20</sup>	v.1.3	spindex -k 8 -p 1 -d silva_spingo_orig.fa spingo -d silva_spingo_orig.fa -k 8 -a -i Cam16S.fa > Cam16S.spingo

#### 2.2.1 Datasets

We used nanopore sequencing data from our mock community and freshwater amplicons for benchmarking the classification tools. We therefore subsampled (a) 10,000 reads from each of the three mock community sequencing replicates (section 1.4), and (b) 10,000 reads from an aquatic sample (April-8; three random draws served as replicates). We then used the above 13 classification tools to classify these reads against the same database, SILVA v.132<sup>19</sup> (Supplementary Figure 1).

## 2.2.2 Comparison of Mock Community Classifications

For the mock community classification benchmark, we assessed the number of unclassified reads, misclassified reads (i.e. sequences not assigned to any of the seven bacterial families), and the root mean squared error (RMSE) between observed and expected taxon abundance of the seven bacterial families. Following the detection of a strong bias towards the *Enterobacteriaceae* family across all classification tools, we also analysed RMSE values after exclusion of this particular family (Supplementary Figure 1b-c).

## 2.2.3 Comparison of River Community Classifications

For the aquatic sample, the number of unclassified reads were counted prior to monitoring the performance of each classification tool in comparison with a consensus classification, which we defined as majority vote across classifications from all computational workflows. We observed stable results across all three draws of 10,000 reads from the same dataset (data not shown), indicating a robust representation of the performance of each classifier.

#### 2.2.4 Overall Classification Benchmark

Minimap2 performed second best at classifying the mock community (lowest RMSE), while also delivering freshwater bacterial profiles in line with the majority vote of other classification tools (Supplementary Figure 1de), in addition to providing rapid speed (data not shown). Yet, the application of this software to our entire dataset caused insufficient memory errors (at ~150 Gb RAM with kmer length 12), likely due to major sequence redundancies within the SILVA v.132 reference fasta file. Hence, to run each of our full samples within a reasonable memory limit of 50 Gb, it was necessary to reduce the number of threads to 1, raise the kmer size ('-k') to 15 and set the minibatch size ('-K') to 25M (i.e., the number of query bases that are processed at any time), prolonging the runtime of several samples to ~three days.

#### 2.3 Bacterial Analyses

## 2.3.1 General Workflow

After applying Minimap2 to the processed reads as explained above (section 2.2.4), we processed the resulting SAM files by firstly excluding all header rows starting with the '@' sign and then transforming the sets of read IDs, SILVA IDs, and alignment scores to TSV files of unique read-bacteria assignments either on the bacterial genus or family level. All reads that could not be assigned to the genus or family level were discarded, respectively. In the case of read assignment to multiple taxa with the same alignment score, we determined the lowest taxonomic level in which these multiple taxa would be included. If this level was above the genus or family level, respectively, we discarded the read.

#### 2.3.2 Estimating the Level of Misclassifications and Contaminants

Across three independent sequencing replicates of the same linear bacterial community standard (section 2.2.1), we found that the fraction of reads assigned to unexpected genus level taxa resides at  $\sim 1$  % when using the Minimap2 classifier and the SILVA 132 database.

Raw quantified DNA, PCR amplicons and sequencing read counts were considerably less abundant in DI water negative controls, as compared to actual freshwater specimens (Supplementary Table 2a). Only the negative control of the most prolific flow cell run (August 2018) passed our high confidence threshold of 37,000 sequencing reads on the family level (Figure 2b, Supplementary Figure 3, section 2.4). Further inspection of these negative control reads revealed that their metagenomic profile closely mimicked the taxonomic classification profiles of river samples within the same sequencing batch, in addition to low-level kit contaminants like alphaproteobacteria of the *Bradyrhizobium* and *Methylobacterium* genus<sup>66</sup> which were otherwise nearly completely absent in any of the true aquatic isolates (Supplementary Table 5).

## 2.4 Rarefaction and High-Confidence Samples

Sample-specific rarefaction curves were generated by successive subsampling of sequencing reads classified by Minimap2 against the SILVA 132 database (section 2.2.1). For broader comparative data investigations, we chose to only retain samples that passed a conservative minimum threshold of 37,000 reads. Family and genus-level

species richness was hence mostly kept at ~90 % of the original values, in accordance with stable evenness profiles across a series of 100 bootstrap replicates (Supplementary Figure 3, section 2.4.1). Although we mainly present a single example rarefied dataset for our final downstream analyses, we repeated each analysis, including PCAs, hierarchical clustering and Mantel tests, based on additional rarefied datasets to assess the stability of the analyses.

#### 2.4.1 Mantel Test

We performed Mantel tests (using scikit-bio version 0.5.1) to compare rarefied datasets with the full dataset. We therefore compared the Euclidean distance based on Z-standardised bacterial genera between all samples with more than 37,000 reads (two-sided test, 99,999 permutations). This resulted in a Pearson correlation of 0.814 (p =  $2.1*10^{-4}$ ) for our main rarefied dataset (results of the Mantel test applied to the remaining three other rarefied datasets: R = 0.819 and p =  $1.0*10^{-4}$ , R = 0.828 and p =  $8.0*10^{-5}$ , R = 0.815 and p =  $1.4*10^{-4}$ , respectively). Results of the Mantel tests applied to the genus-level bacterial classifications were also similar for all four subsampled datasets (R = 0.847 and p =  $1.0*10^{-5}$ , R = 0.863 and p =  $1.0*10^{-5}$ , R = 0.851 and p =  $1*10^{-5}$ , R = 0.856 and p =  $1.0*10^{-5}$ ).

## 2.5 Meta-Level Bacterial Community Analyses

All classification assessment steps and summary statistics were performed in R or python (https://github.com/dj-k/puntseq). We used the python package 'scikit-bio' for the calculation of the Simpson index and the Shannon's diversity as well as equitability index.

#### 2.6 Data Processing for Hierarchical Clustering, Principal Component and Outlier Analysis

Rarefied read count data was subjected to  $log_{10}(x+1)$  and Z-transformations. For the final PCA, negative and mock control samples were initially removed. Mock community samples were then aligned to the eigenspace determined by the water samples and added to plots displaying the main principal components (PCs explaining >10 % variance, respectively). For each of these relevant PCs, we further highlighted the 10 most important features (i.e., taxa with the largest weights) and their contributions to the PCs in barplots, and added the standard error across the three additional rarefied datasets.

For detecting outlier bacterial families per sample, we chose bacteria which were 1.) identified by more than 500 reads, and 2.) which were at least five times more abundant in any single sample than in the mean of all samples combined.

#### 2.7 Pathogen Candidate Assessments

A list of 88 known bacterial pathogenic species, respectively spanning 32 families and 45 genera, was compiled for targeted sequence testing. This was done through the careful integration of curated databases and online sources, foremost using PATRIC<sup>28,29</sup> (Supplementary Table 6a). Additionally, we integrated known genera from a large wastewater reference collection<sup>30</sup> (Supplementary Table 6b).

To identify if DNA reads assigned to *Leptospiraceae* were more similar to sequence reads of previously identified pathogenic, intermediate or environmental *Leptospira* species, we built a neighbour-joining tree of *Leptospiraceae* reads classified in our samples data, together with sequences from reference databases (Figure 3d; species names and NCBI accession numbers in clockwise rotation around the tree in Supplementary Table 7a). We matched the orientation of our reads, and then aligned them with 68 *Leptospira* reference sequences and the *Leptonema illini* reference sequence (DSM 21528 strain 3055) as outgroup. We then built a neighbour-joining tree using Muscle v.3.8.31<sup>67</sup> (excluding three reads in the "Other Environmental" clade that had extreme branch lengths >0.2). The reference sequences were annotated as pathogenic and saprophytic clades P1, P2, S1, S2 as recently described<sup>35</sup>. Additional published river water *Leptospira* that did not fall within these clades were included as "Other Environmental"<sup>68</sup>. Similarly, we constructed phylogenies for the *Legionella*, *Salmonella* and *Pseudomonas* genus, using established full-length 16S reference species sequences from NCBI (Supplementary Table 7b-d).

## 3. Total Project Cost

This study was designed to enable freshwater microbiome monitoring in budget-constrained research environments. Although we had access to basic infrastructure such as pipettes, a PCR and TissueLyser II machine, as well a high-performance laptop, we wish to highlight that the total sequencing consumable costs were held below £4,000. Here, individual costs ranged at ~£75 per sample. With the current MinION flow cell price of £720, we estimate that per-sample costs could be further reduced to as low as ~£15 when barcoding and pooling ~£100 samples in the same sequencing run (for details, see Supplementary Table 3). Assuming near-equimolar amplicon

pooling, flow cells with an output of  $\sim$ 5,000,000 reads can yield well over 37,000 sequences per sample and

thereby surpass this conservative threshold applied here for comparative river microbiota analyses.

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