Time after time: detecting annual patterns in stream bacterial biofilm communities

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Summary
To quantify the major environmental drivers of stream bacterial population dynamics, we modelled temporal differences in stream bacterial communities to quantify community shifts, including those relating to cyclical seasonal variation and more sporadic bloom events. We applied Illumina MiSeq 16S rRNA bacterial gene sequencing of 892 stream biofilm samples, collected monthly for 36-months from six streams. The streams were located a maximum of 118 km apart and drained three different catchment types (forest, urban and rural land uses). We identified repeatable seasonal patterns among bacterial taxa, allowing their separation into three ecological groupings, those following linear, bloom/trough and repeated, seasonal trends. Various physicochemical parameters (light, water and air temperature, pH, dissolved oxygen, nutrients) were linked to temporal community changes. Our models indicate that bloom events and seasonal episodes modify biofilm community changes. Our models indicate that bloom events and seasonal episodes modify biofilm community changes. This is an open access article under the terms of the Creative Commons Attribution-NonCommercial-NoDerivs License, which permits use and distribution in any medium, provided the original work is properly cited, the use is non-commercial and no modifications or adaptations are made.

Introduction
Freshwater streams change over time, including in response to external anthropogenic pressures such as increased urbanization, land-use disturbances and related environmental changes (Jeffries et al., 2015; Labbate et al., 2016; Qu et al., 2017; Kim et al., 2020). Due to stream water and associated microbial communities largely originating from the catchments they drain, the distribution and abundance of stream bacterial communities might provide insights into stream and catchment level ecosystem functional responses to environmental change (Chapin et al., 2000). Indeed, freshwater microbial communities are highly diverse (Battin et al., 2016), play a significant role in ecosystem functioning and are strongly influenced by changes in surrounding landscapes (Lear and Lewis, 2009; Kraemer et al., 2020). Despite this, most studies describing freshwater ecosystem health to date have focused on communities of invertebrates and plants (Sanderson et al., 2007; Wright and Ryan, 2016; Kim et al., 2020). To understand and predict the impact of environmental changes (e.g. abiotic, biotic, anthropogenic, or other processes; Lear et al., 2008) on bacterial community diversity (Loreau et al., 2001) and ecosystem functioning, we first need a baseline understanding of the extent and nature of temporal patterns influencing their distribution and abundance.

Advances in molecular methods, including high-throughput sequencing techniques for determining bacterial community composition and diversity, based on 16S rRNA gene sequences (Good et al., 2018), provide collective evidence that bacteria exhibit substantial spatial and temporal patterns in their distribution and abundance (Lear et al., 2008; Esposito et al., 2016; Gautam et al., 2020). Highly dominant and diverse bacterial phyla reported among freshwater environments include Proteobacteria, Cyanobacteria, Bacteroidetes, Verrucomicrobia, Actinobacteria and Planctomycetes (Besemer et al., 2012; Battin et al., 2016), but their dominance varies greatly over time and geographic location. For example, a greater prevalence of phyla such as the Cyanobacteria is likely to result from warmer temperatures, reduced flows and high light availability (Salmaso et al., 2018). Investigating the extent of repeating annual patterns for multiple bacterial taxa could reveal their potential to serve widely as predictors of environmental change, both over time and among catchments. Furthermore, it may aid in better prediction of harmful cyanobacterial blooms.
We investigated spatial and temporal patterns in freshwater stream bacterial community composition over multiple years. Building on the work of Gautam et al. (2020), we sampled stream biofilm communities monthly from six streams over 3 years (2013–2016), collecting a suite of environmental data from each stream, including (i) instream measurements such as water turbidity, temperature, pH, total suspended solids and dissolved oxygen concentration, (ii) local climate measurements such as air temperature, light and rainfall at each study site and (iii) average soil moisture deficit (i.e. the amount of rainfall, in mm, required to return the catchment soil to field capacity) and concentrations of nutrients such as nitrogen, phosphorus and ammonia in the upstream catchment. First, we used 16S rRNA gene sequencing analysis to reveal bacterial community assemblage patterns. Second, we analysed whether the observed patterns in composition and diversity are repeatable over annual seasonal cycles or if blooms in the abundance of certain taxa occur. Third, we assessed the extent to which environmental factors impact the observed temporal shifts in stream bacterial community dynamics. We set up a model framework to identify bacterial taxa responding to either bloom or seasonal patterns over time. We postulate that freshwater stream bacterial community dynamics can be predicted by seasonal change and provide a priori hypotheses for future experiments, facilitating an efficient approach to understand community assembly and succession in bacterial communities across freshwater ecosystems.

Results

Our analysis generated a total of 123,121 quality-checked 16S rRNA gene sequences. These sequence data were then rarefied to 2000 reads per sample, after which we obtained 774 samples from six different streams. These high-quality sequences were clustered into 21,216 operational taxonomic units (OTUs) at 97% sequence similarity. The bacterial OTUs were classified into 61 phyla, 261 orders, 313 families and 473 genera. The PRIMER RELATE function found a correlation (Rho = 0.8) between amplicon sequence variant (ASV) and OTU resemblance matrices at a significance level of 0.01%. This suggests that both methodologies provide broadly similar results (whether obtained from OTU or ASV files) and hereafter we report only the outputs of OTU table analyses. Diversity index analysis (Supplementary Material 1) showed that streams from highly impacted urban catchment of Oakley Creek had significantly higher relative bacterial richness (OTUs) and diversity compared to the streams draining predominantly native catchments (Tukey HSD multiple pairwise comparisons, p < 0.05).

Bacterial community variation (assessed using PERMANOVA models)

Multivariate analysis was used to elucidate and display spatial and temporal variations within the stream biofilm bacterial communities. Permutational multivariate analysis of variance (PERMANOVA) model 1 revealed that year is a significant individual factor correlated with the linear change observed in bacterial community composition (p < 0.01) along with stream and months (Supplementary Material 2). The interactions between all the variance factors [ST(CA)xMO(YExSE)] correlate even more strongly with the observed changes in community composition, as shown by their highest square root contribution in this model (Supplementary Material 2). Likewise, PERMANOVA Model 2 (Supplementary Material 3) confirmed significant cyclical patterns in our extended dataset using monthly covariates with variance components (catchment, stream, year). Similar to our previous findings from this model, our extended dataset captured change occurring across repeated annual cycles, by sampling every month, over multiple years. The associations of catchment type, year and stream were also found to correlate significantly with the monthly covariates (Supplementary Material 3; PERMANOVA; p < 0.01). An nMDS (Fig. 1) ordination based on relative abundance data was plotted to illustrate the patterns in these data, where large variability in bacterial community composition is observed over the years compared to among streams and catchment types. There was a clear separation of data collected at different seasons and months, including cyclical patterns in the data (i.e. in average monthly data, Fig. 1).

Fig. 1. nMDS ordination showing relative relationships among the bacterial community data averaged by (green square) month (where the numbers 1–12 represent the months January to December) respectively, (blue circle) year, (pink rhombus) season, (inverted red triangle) catchment and (blue triangle) stream using Bray–Curtis similarities. 2D Stress value: 0.15.
Bacterial community diversity and composition

The highest proportion of OTUs belonged to the phylum Proteobacteria across all samples (32.5%), followed by Planctomycetes (11.2%), Bacteroidetes (9%), Verrucomicrobia (5.5%), Chloroflexi (4.6%), Acidobacteria (4.3%), Actinobacteria (4%), Firmicutes (3.6%), Cyanobacteria (3.6%) and Chlamydiae (1%).

The cyanobacterial order Oscillatoriales was present in all the streams (except for Wairoa and Oakley creek) during summer 2014 and spring 2015, with a high abundance in the rural Mahurangi stream (>90% in October) (Supplementary Material 4). Planctomycetes made up 9% of the community overall (orders Gemmatales, Pirellulales, Planctomycetales) largely from February 2013 to September 2014 and then from May 2015 until the end of the sampling period, with greater representation during autumn and winter. The phylum Firmicutes (orders Bacillales, Lactobacillales, Clostridiales) made up 6.3% of the community. Firmicutes were less abundant in the early sampling period, but their relative abundance increased significantly in the later summer sampling period (December 2015–February 2016). This sudden increase in the order Bacillales appeared to be influenced by a significant increase of pH and dissolved oxygen during that time (Supplementary Material 5). The phylum Bacteroidetes (orders Saprospirales, Bacteroidales, Cytophagales, Flavobacteriales, Sphingobacteriales) contributed 7% of the community and was present in all the streams. Other well-represented phyla present in the dataset were Actinobacteria (4.3%) and Verrucomicrobia (4.2%) (Fig. 2, Supplementary Material 4).

Temporal patterns in bacterial community composition (time series models)

Examination of temporal trends exhibited by the top 50 most abundant bacterial orders using three conceptual time series generalized linear models showed significant differences among the bacterial taxa. In terms of time series model 1 (linear), five bacterial orders fit this model well (Table 1). Twenty-six bacterial orders significantly fit patterns indicative of time series model 2a (bloom) and 2b (trough) occurring sporadically since the beginning of sampling; 19 orders followed significant repetitive seasonal patterns, based on time series model 3 (seasonal). A complete list of orders identified by time series generalized linear model analysis is provided in Table 1.

We further visualized shifts and differences in the abundance of bacterial orders (Fig. 3) relating to each of our time series models by assembling these data into three groups, represented within a shade plot of taxon abundances (Fig. 4). Orders identified from Model 1 (Table 1) were maximally abundant from 2013 to 2014, decreasing in abundance in late 2015–2016 (Supplementary Material 4, Fig. 3A). We plotted subsets of the bacterial communities recognized by model 2 against months to distinguish between communities developing from blooms and troughs over the years (Fig. 3B). Results from time-series models were similar to the PERMANOVA models examined in the previous section, in terms of differences in the relative abundance of bacterial communities over time (years), where some orders were more or less dominant with environmental and seasonal changes. For example, the orders Bacillales, Actinomycetales, Lactobacillales, Pseudomonadales, Deinococcales, SM2F11 and Nostocales were more abundant in 2014–2016, whereas Saprospirales, Myxococcales, Methyllococcales, Stromeniopes were more prominent in 2013 during November, December, January and February. There were distinct differences between autumn and winter bloom communities of 2013, 2014 and 2015 (Fig. 3B). The outcomes of these inferences suggest that large variations exist in the conditions required for different taxa to increase in abundance. Further analysis of seasonal communities from time series model 3 revealed a greater abundance of Flavobacteriales and Enterobacteriales during summer; Acidimicrobiales, Pirellulales, Gemmatales, Clostridiales and Planctomycetales during winter and autumn, and Cyanobacteria, Caulobacteriales and Methylophilales during spring (Fig. 3C).

Influence of physicochemical parameters on bacterial community composition

Distance-based linear models (DistLM) analysis was performed to examine the relative influence of physicochemical parameters on subgroups of the bacterial communities identified from analysis of the three time series GLM models. Light, pH, nitrates, total phosphorus and water temperature (Table 2) combined were able to explain 8% (adjusted $R^2$) of the total variation in the composition of the subset of taxa identified by model 1 (linear). The subset of the bacterial community identified by model 2 (bloom/trough) was found to be significantly affected by light, pH, deficit, total nitrogen, water temperature, turbidity, total phosphorus, concentration of dissolved oxygen, total suspended solids, nitrates, ammonia and maximum air temperature (Table 2), which explained 12% (adjusted $R^2$) of the total observed variation. Similarly, light, deficit, pH, total nitrogen, ammonia, soluble phosphorus, nitrates, TKN (total Kjeldahl nitrogen), turbidity, total suspended solids, water temperature and dissolved oxygen were significantly correlated with the subset of the bacterial community identified by model 3 (seasonal), explaining 10% (adjusted $R^2$) of the total variation (Table 2).
Discussion

Long-term investigations and temporal predictions of stream microbiology require new and improved statistical tools to comprehend temporal variability in freshwater bacterial community composition and potential environmental drivers that influence these patterns. Here, we provide a robust statistical approach for analysing bacterial community compositional shifts over time in freshwater stream biofilms, using data collected monthly. We developed and tested three time series generalized linear models by exploiting stream bacterial communities to investigate temporal patterns (month, year) and seasonal shifts from our monthly sampling period (February 2013–February 2016) for 3 years. Our results indicated that bacterial taxa respond to monthly and seasonal environmental changes, and some may be categorized as bloom and seasonally responsive taxa. Our models may be used to reduce the complexity of identifying differences in bacterial community composition, based on their responsiveness to the changing environment and assist with the prediction of community patterns over the years.

**Freshwater stream biofilm bacterial community composition**

PERMANOVA models used previously (Gautam et al., 2020), but incorporating substantial additional data, confirmed that the bacterial community composition varied between streams and over time, supporting observations from previous studies (Lear et al., 2008; Hassell et al., 2018). Proteobacteria formed the largest relative fraction of the bacterial community in our dataset; they are recognized as the most common dominant phylum in benthic and hyporheic biofilms overall (Battin et al., 2016). Abundant cyanobacterial communities (mainly Oscillatoriales) in our dataset were correlated with changes in light intensity and nutrients, as similarly observed by others (McCall et al., 2017). We also observed a relatively high abundance of the phylum Firmicutes (order Bacillales) during spring and summer 2015 and 2016 when dissolved oxygen, water temperature and pH were higher, creating ideal conditions for Firmicutes growth (Zhao et al., 2017), since dissolved oxygen is typically elevated during summer (Schmidt et al., 2017). We observed a considerable relative abundance of Planctomycetes (11.2%) in our community dataset despite them being recognized as a less-abundant bacterial phylum in freshwater ecosystems. It has been speculated that composition of Planctomycetes in freshwater ecosystems is associated with seasonal variations (Brümmer et al., 2004) and influenced by parameters including water temperature, pH and nutrient concentrations (Pollet et al., 2011).

**Temporal patterns in bacterial community diversity**

PERMANOVA models and time series generalized linear models designed and tested in this study provide...
evidence that stream bacterial community composition is highly responsive to seasonal and environmental changes. The time series model 1, which identified bacterial taxa responding to linear trends over time, comprised the least taxa but included some common Proteobacteria and Bacteroidetes orders (Battin et al., 2016). With increasing temperature and pH, the order Sphingomonadales (phylum proteobacteria) and Cytophagales (phylum Bacteroidetes) were observed at increased abundance. Members of the phylum Bacteroidetes are reported to break down biopolymers that contribute to the high molecular weight fraction of dissolved organic debris in streams (Battin et al., 2016) and so increased water temperature may have significant impacts on Bacteroidete-associated transformations of organic matter within freshwater systems.

| Order            | Model 1 (linear) AIC 0.58 | Model 2 (bloom) AIC 0.34 | Model 3 (seasonal) AIC 0.08 |
|------------------|---------------------------|--------------------------|-----------------------------|
| Cytophagales     | 724.04                    | 724.85                   | 727.51                      |
| Sphingomonadales | 952.1                     | 953.96                   | 954.06                      |
| Sphingobacterales| 465.4                     | 467.16                   | 465.62                      |
| Pedosphaerales   | 287.13                    | 287.42                   | 288.03                      |
| Oceanospirillales| 209.21                    | 209.83                   | 210.11                      |
| Rhizobiales      | 1131.7                    | 1123.3                   | 1127.4                      |
| Bacillales       | 1357.1 0.01               | 1330.4 1               | 1347.5                      |
| Verrucomicrobiales| 760.11                    | 751.75                   | 760.27                      |
| Sapropiraeles    | 774.78 0.06               | 769.36 0.86              | 773.88 0.08                 |
| Actinomycetales  | 680.14 0.03               | 673.15 0.95              | 681.1 0.02                  |
| Xanthomonadales  | 632.07 0.05               | 626.13 0.94              | 635.44 0.01                 |
| Pseudanaebaeales | 692.91 0.01               | 681.38 1               | 696.37 0.01                 |
| Synechoococcales | 797.42 0.07               | 792.04 0.97              | 800.32 0.01                 |
| Chroococcales    | 697.75 0.02               | 689.66 1               | 698.83 0.01                 |
| SM2F11           | 580.45 0.14               | 576.61 0.84              | 583.88 0.02                 |
| Myxococcales     | 185.37 0.27               | 183.54 1               | 186.33 0.13                 |
| Lactobacillales  | 220.22 0.02               | 202.42 1               | 220.1 0.01                  |
| Rhodobacterales  | 974.18 0.09               | 970.33 0.53              | 970.77 0.08                 |
| Pseudonanaebaeles| 1257.3 0.01               | 1249.1 0.69              | 1249.5 0.03                 |
| Legionellales    | 384.56 0.02               | 377.77 0.55              | 378.05 0.43                 |
| Oscillatoriataes | 1342 0.42                 | 1341.2 0.54              | 1346 0.04                   |
| Rhodospirillales | 357.49 0.34               | 356.23 0.57              | 359.83 0.08                 |
| Deinococcales    | 471.66 0.32               | 470.12 0.62              | 474.45 0.06                 |
| Methylococcales  | 655.06 0.33               | 653.61 0.61              | 658.09 0.06                 |
| Nostocales       | 639.18 0.29               | 637.66 0.56              | 640.03 0.15                 |
| Bacteria         | 156.51 0.34               | 155.28 0.57              | 158.68 0.09                 |
| Bacteroidetes    | 350.48 0.32               | 348.93 0.62              | 353.39 0.06                 |
| Ellin6067        | 116.8 0.23                | 114.4 0.68              | 118.29 0.09                 |
| SM2F09           | 382.05 0.29               | 380.33 0.61              | 383.69 0.15                 |
| OD1              | 219.86 0.27               | 217.87 0.64              | 221.48 0.09                 |
| Stramenopiles    | 1290.6 0.33               | 1289.5 0.52              | 1291.7 0.15                 |
| Flavobacteria    | 913.24 0.16               | 914.33 0.08              | 909.71 0.75                 |
| Acidimicrobiales | 600.99 0.00               | 602.74 0.00              | 583.64 1                    |
| Burkholderiales  | 978.72 0.12               | 979.28 0.08              | 974.43 0.8                   |
| Pirellulales     | 923.75 0.02               | 924.22 0.02              | 915.76 0.96                 |
| Gemmatales       | 788.95 0.01               | 786.16 0.03              | 778.78 0.97                 |
| RB41             | 389.19 0.00               | 386.69 0.00              | 374.15 1                    |
| Chthoniobacterales| 309.13                    | 302.66 0.18              | 299.33 0.82                 |
| SJC184           | 472.2 0.01                | 471.87 0.01              | 461.54 0.99                 |
| WD2101           | 538.32 0.00               | 540.21 0.00              | 525.88 1                    |
| Ellin6529        | 422.7 0.09                | 423.1 0.07              | 417.72 0.85                 |
| Cyanobacteria    | 244.26 0.13               | 245.46 0.06              | 240.1 0.81                  |
| Clostridiales    | 258.44 0.11               | 259.65 0.06              | 253.98 0.83                 |
| Aeromonaediales  | 182.38 0.06               | 183.37 0.03              | 176.42 0.91                 |
| MN1D             | 15.202 0.23               | 16.045 0.12              | 12.698 0.64                 |
| Planctomycetales | 379.52 0.02               | 381.33 0.01              | 370.78 0.98                 |
| Enterobacteria   | 1065.3 0.04               | 1060.9 0.31              | 1059.2 0.65                 |
| Caulobacteria    | 261.72 0.31               | 263.62 0.11              | 260.03 0.58                 |
| Methylophilales  | 198.53 0.17               | 197.45 0.26              | 195.57 0.58                 |
| Streptophyta     | 248.25 0.23               | 247.52 0.3              | 246.37 0.47                 |
The relative abundance of cyanobacterial and proteobacterial orders adhered most closely to a blooming pattern of temporal change (time series model 2). Indeed, the progression of diatoms in spring and a more diverse community dominated by chlorophytes and cyanobacteria in spring and/or autumn is a widely identifiable trend in freshwater streams (Salmaso, 2000). With increase in the relative abundances of cyanobacteria particularly in spring 2014 (Fig. 2), the selection of cyanobacteria by time series model 2 is not surprising.
Representative cyanobacterial orders such as Pseudanabaenales, Synechococcales, Chroococcales, Oscillatoriales and Nostocales dominate during spring and autumn blooms, indicating favourable growth conditions for cyanobacterial blooms in reduced flows and increased residence time with high temperature (Steffen et al., 2012). Nostocales' ability to fix nitrogen is thought to be a crucial component in their dominance during high turbidity and bloom conditions, indicative of regional changes in nutrient loads, as well as due to changes in various biotic conditions (Wu et al., 2012). Nostocales has been identified as an indicator of global warming and is regarded as an effective nutrient competitor (Sukenik et al., 2012) while being toxic to humans and animals (Stewart et al., 2011). Pseudanabaenales and Chroococcales, on the other hand, possess the ability to create a favourable environment for their growth and metabolic activities by exchanging specialized chemical signals (Rasmussen and Givskov, 2006; Muñoz-Garcia and Ares, 2016). For example, orders Chroococcales (also regarded as *Microcystis* species), Oscillatoriales and Synechococcales are positively associated with nutrient enrichment. Warming of surface waters, promoting more substantial and longer thermal stratification periods (Kormas and Lymperopoulou, 2013; Ma et al., 2016; Salmaso, 2000; Zhu et al., 2019), can also increase the abundance of Oscillatoriales in late summer, perhaps as they occupy new ecological niches along vertically segregated environmental gradients (Salmaso, 2000). Environmental disturbances, for instance, elevated temperature, high light and eutrophic conditions (Olappede and Leff, 2005; Simonin et al., 2019; Zhu et al., 2019) provide ample opportunities that can be easily and competitively utilized by opportunistic bacterial taxa, including bloom-forming bacterial communities (Piehler et al., 2009). Research shows that cyanobacteria interact with their biotic environment in various ways, from impacting predator–prey interactions to forming mutualistic relationships with micro, macroalgae and non-photosynthetic protists (Bauer and Forchhammer, 2021; Ma et al., 2017; Mutalipassi et al., 2021; Usher et al., 2007). Cyanobacteria and proteobacteria coexist in

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**Table 2.** Summary of the relationship between bacterial community composition and physicochemical variables using DistLM analysis of subsets of the community data, as selected by comparison of three time series GLM models.

| Model 1 (variable)       | Adjusted $R^2$ (cumulative) | SS (trace) | Pseudo-$F$ | $p$  |
|--------------------------|-------------------------------|-----------|------------|------|
| Light                    | 0.040462                      | 8732.6    | 8.3795     | 0.0003 |
| pH                       | 0.07145                       | 6864.5    | 6.8068     | 0.0006 |
| Nitrate + nitrite        | 0.086744                      | 3865.4    | 3.8971     | 0.0095 |
| Total phosphorus         | 0.088667                      | 1349      | 1.363      | 0.2401 |
| Water temperature        | 0.089744                      | 1188.5    | 1.2022     | 0.2849 |

**Model 2 (variable)**

| Light       | 0.041976 | 14,320 | 8.6676 | 0.0001 |
| pH          | 0.068711 | 9628.5 | 5.9953 | 0.0003 |
| Deficit     | 0.080587 | 5128.4 | 3.2345 | 0.0067 |
| Total N     | 0.092735 | 5167.9 | 3.3031 | 0.0056 |
| Water temp. | 0.10319  | 4630   | 2.9938 | 0.012  |
| Turbidity   | 0.11018  | 3584.4 | 2.3959 | 0.0339 |
| Total P     | 0.11478  | 2866.4 | 1.8777 | 0.0817 |
| DO % Sat    | 0.11663  | 2057.8 | 1.3508 | 0.2101 |
| Total solids| 0.11813  | 1954.6 | 1.2853 | 0.2446 |
| Nitrate + nitrite | 0.11877 | 1701.7 | 1.1198 | 0.3265 |
| Ammonia     | 0.12112  | 2184.1 | 1.441   | 0.1782 |
| Mean Tmax   | 0.12117  | 1530.3 | 1.0097 | 0.3954 |

**Model 3 (variable)**

| Light       | 0.047729 | 12,414 | 9.7712 | 0.0001 |
| pH          | 0.067584 | 5853   | 4.7052 | 0.0005 |
| Deficit     | 0.080412 | 4187.6 | 3.4133 | 0.005  |
| Total N     | 0.084612 | 2185   | 1.7892 | 0.0985 |
| Ammonia     | 0.087952 | 1978.7 | 1.6262 | 0.1338 |
| Soluble P   | 0.090824 | 1864.3 | 1.537  | 0.1723 |
| Nitrate + nitrite | 0.093122 | 1727.9 | 1.4282 | 0.1946 |
| TKN         | 0.097542 | 2194.7 | 1.8229 | 0.0557 |
| Turbidity   | 0.098554 | 1428.2 | 1.1875 | 0.2856 |
| Total solids| 0.10125  | 1795.3 | 1.4973 | 0.1754 |
| Water temp. | 0.10166  | 1290.3 | 1.0766 | 0.3466 |
| DO % Sat    | 0.10221  | 1318.3 | 1.1007 | 0.3444 |

A forward section approach was utilized to identify environmental variables correlated significantly with observed changes in bacterial community composition (based on Bray–Curtis dissimilarity). These variables were fitted sequentially in order, with each subsequent variable being included in the model based on those which precede it in the table. $p$ values were obtained using 999 permutations of the data. Mean Tmax refers to average daily maximum air temperature.
>90% of bloom-forming bacterial communities, contributing equally to the overwhelming majority of functional genes detected during bloom formation (Steffen et al., 2012). Indeed, mesocosm studies confirm associations of various proteobacterial orders, such as Legionellales, Rhizobiales and Rhodobacterales, within freshwater cyanobacterial blooms (especially Microcystis) (Li et al., 2011). Such interactions make sense because variables like grazing and viral lysis can cause blooms to decline (Riemann et al., 2000; Yager et al., 2001; Löder et al., 2011; Buchan et al., 2014), thereby supplying a diverse range of nutrient resources and organic matter that promotes proteobacterial growth and production (Bird and Karl, 1999). Firmicutes are typically underrepresented in freshwater habitats and exhibit low abundance during cyanobacterial blooms. Nevertheless, they possess numerous biodegradation competencies that are shown to be associated with cyanobacterial degradation (Kormas and Lymperopoulou, 2013). With these findings, we suggest that future bloom predictions should be based on an improved knowledge of bacterial community interactions, as well as physicochemical parameters such as differences in temperature, light, pH and nutrients, all of which have been shown to influence bloom-forming bacterial communities in freshwater. To investigate potential bloom trends in freshwater streams, regular sampling throughout time is essential, as changes in the bacterial populations may precede bloom events.

Our “seasonal” time series model (model 3) was defined mainly by Planctomycetes (Pirellulales, Gemmatales, WD2101, Planctomycetales) and Proteobacterial (Burkholderiales, Enterobacteriales, SC-I-84, Aeromonadales, Enterobacteriales, Caulobacterales, Methylophilales) orders. Orders Planctomycetes Pirellulales and Gemmatales, as well as orders WD2101, Planctomycetales, Methylophilales, Chthoniobacterales and Acidimicrobiales, were abundant in winter and autumn, presumably due to seasonal succession from the ground up bacterial community composition. In contrast, Pollet et al. (2011) and Jijun et al. (2006) discovered a greater abundance of Planctomycetes during the spring and summer seasons. Here, we confirmed that Caulobacterales, Flavobacteriales, Aeromonadales and Enterobacteriales dominated summer and spring communities; high Enterobacteriales abundances were positively linked with bloom-forming Bacillales and Pseudomonadales orders. Bacillales and Enterobacteriales are facultative anaerobes that have been previously detected in heavy metal and faecal contaminated sites and can survive extreme weather conditions (Humayoun et al., 2003). We propose that many changes in bacterial community composition are not continuous over time and that continual alteration in community composition is induced by slow environmental change and disparities in resource availability.

Aquatic microbial communities have been demonstrated to be temporally dynamic, with repeating patterns of community structure (Fuhrman et al., 2006; Shade et al., 2007; Cram et al., 2015; Fuhrman et al., 2015; Yan et al., 2017; Capo et al., 2019). Various studies have used culture-independent sequencing approaches and high-throughput 16S amplicon sequencing to confirm repeated changes of microbial communities in bloom-affected freshwater bodies (Eiler et al., 2012; Li et al., 2015; Woodhouse et al., 2016; Tomas et al., 2017). However, it is difficult to extrapolate the predictions from bloom and seasonal effects robustly if the studies are carried out for only short periods of time. Therefore, information on whether freshwater bacterial population dynamics are repeatable and predictable at different spatial and temporal scales, including incorporating samples collected over several consecutive years is highly advantageous.

Light, temperature and nutrients are growth-limiting factors for freshwater bacterial communities, essentially bloom-forming and seasonal bacterial communities (Olpadale and Leff, 2005; Paerl et al., 2011; Zhu et al., 2019). Here, supporting the results of previous studies, water temperature and average maximum daily air temperature, along with concentrations of ammonia, nitrates, phosphorus, dissolved oxygen, light, pH, moisture deficit, total suspended solids and turbidity were found to be significantly associated with bloom and seasonal patterns (time series model 2, 3) explaining 12% and 10% of the observed variation [adjusted R² (cumulative)] respectively. Water temperature, light, pH, total phosphorus and nitrates were significantly correlated with abundances of bacterial orders selected by the time series model 1, explaining 8% of the total variation observed. These results suggest that seasonal changes in freshwater, as well as differences in nutrient levels, provide ideal conditions for the growth of specific bacterial communities to form blooms that can dominate for short periods while also influencing competition for available nutrients among other organisms (Mur et al., 1999; Stewart et al., 2011; Zhu et al., 2019). We demonstrate that the relative abundances of some bacterial community members are cyclically repetitive (Table 2). However, as we can only explain 12% or less of the total observed variation, more investigation is required to understand the underlying cause of these variations, for example relating to spatiotemporal changes in environmental drivers. Unmeasured abiotic factors such as water conductivity, salinity, anthropogenic activities and other biotic factors such as competition, mutualism and succession could explain the rest of the variation. Additionally, stochastic
factors, dispersal limitation, mass immigration effects, as well as deterministic factors such as species sorting likely influence bacterial community differences over time. Future research is needed to confirm community responses to a greater diversity of environmental, seasonal and successional changes and thus to incorporate them into conceptual models. An increased sampling frequency including measurements of biotic and abiotic environmental drivers might enable more accurate predictions of bloom, pre-bloom and seasonal specific communities. This would further help to predict repetitive patterns within stream biofilms, and the extent to which these predictors are stream specific.

Conclusions

Given that there are unexplained drivers of differences in freshwater stream bacterial community composition, we recognized that spatially and temporally heterogeneous bacterial communities cannot be adequately defined by any one simple statistical model. Using multiple time series models, we have made first attempts to conceptualize the role of local and regional environmental drivers of stream bacterial community composition to provide a more flexible methodology for freshwater ecologists to understand bacterial community assembly and succession. Overall, we demonstrate a conceptual framework to better understand bacterial community assembly, learn the reasons for changes in the relative abundance of specific taxa, and quantify their response to environmental changes.

Experimental procedures

Sample collection, processing and DNA extraction

The six principal sampling locations for this study were based in the Auckland region of New Zealand, draining three different catchment types. These consisted of streams draining two urban, two rural and two native forest catchments. Cascades and Wairua streams (Supplementary Material 6) drain densely vegetated native forest reserves and have consistently good water quality (Foley et al., 2018; Buckthought et al., 2020). Oakley and Otara Stream catchments drain mostly urban, bare and lightly vegetated surfaces with significant human populations and residential and commercial activities. Mahurangi and Ngakaroa streams are rural streams and drain catchments dominated by improved exotic grassland and other herbaceous vegetation used for grazing livestock. A summary of each stream’s catchment area, land cover distribution and location is provided in Supplementary Material 7.

We collected five stream biofilm samples within ~10 m reach of each stream, monthly for 36 months (February 2013–February 2016). Biofilm biomass was removed from five submerged rocks by scraping the upper surface of each using a separate, sterile Speci-Sponge™ (Nasco, USA) before placement into individual Whirl-Pak™ bags (Nasco, USA). Bags were sealed and held at 4°C before being transported back to the laboratory and frozen (~20°C) until further analysis (Lear and Lewis, 2009). Auckland Council collected physicochemical stream water attributes, adhering to clear data collection standards (APHA, 1998) and collated them in a National Institute of Water and Atmospheric Research (NIWA; https://cliffo.niwa.co.nz/) database. At each location and sample date, the data for this study were collated as described by Gautam et al. (2020).

Each biofilm sample was thawed and thoroughly macerated to separate the biofilm samples from the sponges using a Lab Stomacher 400 device (Seward, Norfolk, UK). We extracted DNA from each pelleted biofilm sample using the method described by Miller et al. (1999). This method utilizes a bead-beating methodology with chloroform–isoamyl alcohol extraction (Miller et al., 1999). The protocol has been described previously for biofilm sample processing and DNA extraction (Gautam et al., 2020).

PCR amplification and sequencing

To characterize the bacterial communities present in each sample, we amplified V3/V4 regions of bacterial 16S rRNA genes in each biofilm DNA extract using modifications of the Universal 16S rRNA amplicon primers 341F (5’-TCGTCGCAAGTCATGGTATCAAGAGACGCTTAGAGTCCCTACGGGGNGGCWGCAG-3’) and 785R (5’-GTCTCGTGGGCTCGAGATGTGTATAAGAGACAGTGTTGAGACAGGACTACHVGGGTATCTAATCC-3’) (Edgar, 2013). The primers include Illumina adapter sequences (bold and underlined) that are required for downstream sequencing. PCR amplification and library preparation were performed as described in Gautam et al. (2020). Sequencing was conducted by Auckland Genomics (University of Auckland, New Zealand) on an Illumina MiSeq instrument using 2 × 300 bp chemistry. Before sequencing, the sequencing provider attached a unique combination of Nextera XT dual indices (Illumina, USA) to the DNA from each sample to allow multiplex processing (Illumina, 2013). We deposited all raw amplicon sequence data in the NCBI Sequence Read Archive under accession number PRJNA643645.

Bioinformatics and statistical analyses

DNA sequence data were processed as described previously (Hermans et al., 2020) using the VSEARCH v 7.0 (Edgar, 2013) pipeline. Forward and reverse reads were
merged using the `fastq_mergepairs` command. Sequences with quality scores lower than 20 were trimmed and those with a length shorter than 150 bp were filtered out of all the merged sequences using the `fastq_filter` command. Finally, we dereplicated sequence data (`derep_fulllength`) and removed singleton reads (-sort by size). The remaining sequences were then clustered into species-level OTUs at 97% similarity cut-off in QIIME (Quantitative Insights into Microbial Ecology, version 1.8) and classified against the Greengenes reference database v13.8 (McDonald et al., 2012). We also used DADA2 in R (Callahan et al., 2016) to construct taxon tables of exact ASVs for comparison with our OTU data outputs.

Statistical analyses were performed in Primer-E with the additional add-on PERMANOVA+ (Version 7; Plymouth, UK; Clarke and Gorley, 2015) and using the R software environment for statistical computing and graphics (Version 1.3.959; R Core Team, 2020). The OTU table was rarefied to a sequencing depth of 2000 sequences per sample to maintain the highest number of samples possible after removing those poor numbers of DNA sequence reads, achieved using the ‘rarefy’ function in the ‘vegan’ package (Oksanen et al., 2013) in R to achieve an equal and comparable sequence depth across all samples. After rarefaction, there were 34 440 OTUs across all the samples. The ‘vegan’ package was also used to compute a Bray–Curtis dissimilarity matrix to compare bacterial communities’ alpha diversity (OTU richness, Shannon, Simpson) among all sites. We used the RELATE routine in PRIMER7 to compare the resemblance matrix obtained from both ASV and OTU tables, based on Spearman rank correlation coefficients, using 9999 permutations.

A total of 892 samples were analysed before rarefaction, but we removed 117 samples from our analysis because they had fewer than 2000 DNA sequence reads. After this processing, any site/timepoint combinations represented by less than three replicates were excluded to achieve a fair comparison of data collected among different sites and times. Variations in bacterial community composition were evaluated using pairwise similarities among samples by calculating Bray–Curtis dissimilarities from the OTU table. Dissimilarity matrices were visualized using non-metric multidimensional scaling (nMDS), generating a graphical representation of bacterial community composition’s spatial and temporal patterns. The Bray–Curtis dissimilarity data were then statistically scrutinized using PERMANOVA models in PRIMER as described previously (Gautam et al., 2020). Factors were nested where appropriate and these factors are highlighted in parentheses [e.g. where ST(CA) denotes

Fig. 5. Conceptual diagrams of three time series generalized linear models used to investigate temporal cyclicity in stream bacterial community dynamics across multiple seasons. (A) Model 1 is based on a linear trend as shown by a straight line; (B, C) Model 2 is based on blooms/troughs as interpreted by a curve; (D) Model 3 is based on repetitive seasonal trends represented by periodic waves.

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the factor ‘stream’ is nested within the factor ‘catchment’]. The dynamics of the bacterial taxonomy structure and composition across streams and years were plotted according to their relative abundance (percentage) and visualized using the package ‘ggplot’ in R, at phylum and order levels.

A shade plot/heat map based on the Bray– Curtis similarity matrix was constructed to delineate the abundance and distribution of the top 50 most abundant bacterial orders from our dataset across streams and time. The relative abundances of bacterial community data (orders) were standardized by log \((n + 1)\) before clustering them via a Bray– Curtis similarity matrix, done using Primer-E, v.7.

We developed three time series models to identify significant correlations and temporal patterns of bacterial community distribution and abundance following environmental change over time using Generalized linear modelling (GLM) (Myers and Montgomery, 1997) in R. In these models, both time and space (i.e. streams) were applied as covariables and as fixed factors. Model 1 (linear) was constructed to assess linear trends (either increasing or decreasing) in the data based on ‘days_since’ the start of the experiment. Model 2 (blooms/troughs) is based on ‘days_since + I(days_since^2)’ to identify blooms or non-linear relationships in the data since first sampling. Model 3 (seasonal trends) is based on ‘days_since +sin(2π× days_since/365) + cos(2π× days_since/365)’ to recognize seasonal patterns since first sampling, using sine and cosine terms. To visually inspect the outcomes of these GLM functions, each bacterial order was compared to our conceptual models (Fig. 5) and visualized using the function ‘ggplot’. These models enabled us to identify statistically significant patterns in bacterial community data over time, quantified by \(R^2\) adjusted values; ggpplots provided a visual inspection of their relationship. Selection of the most plausible model for each order was carried out by calculating and comparing Akaike’s information criterion (AIC) (Akaike, 1998) followed by model averaging (multimodel inference) using AICmodavg (Mazerolle, 2020). Later, we used these outcomes to understand the differential patterns in individual orders (i.e. to which of the three distinct model categories data corresponding to these taxa most closely adhere to). Various physicochemical factors were recorded in-stream at the time of sampling (i.e. turbidity, water temperature, pH, total suspended solids, nitrate + nitrite, ammonia as N, TKN (Total Kjeldahl Nitrogen), TN (Total Nitrogen), DO% sat (Dissolved Oxygen, saturated), TP (Total Phosphorus) and SP (Soluble Phosphorus). We obtained average daily measurements of mean Tmax (mean air temperature maximum), mean Tmin (mean air temperature minimum), light, rain and soil moisture deficit from a NIWA Cliffo database, which were averaged per month to use as physicochemical factors. We also coupled our bacterial data with detailed upstream catchment information. Catchment boundaries upstream of each river sampling location were delineated within ARCGIS (ESRI, 2010) (Supplementary Material 6) based on a river layer from the New Zealand River Environment Classification system (REC) (Snedler et al., 2004).

A non-parametric DistLM analysis (Anderson et al., 2008; Chu et al., 2009; Botwe et al., 2015) was performed to explore the relationships between the various physicochemical variables and bacterial community composition based upon Bray–Curtis dissimilarities among the latter using PRIMER. Each of these environmental variables was tested for significant correlation with the subgroups of bacterial community data identified from the analysis of each of the three time series GLM models. We used a forward selection procedure and adjusted \(R^2\) to identify variables that explain the greatest amount of variation in the bacterial community, using 999 permutations of the data.

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**Supporting Information**

Additional Supporting Information may be found in the online version of this article at the publisher’s web-site:

**Supplementary Material 1.** Box plots for OTU richness, Shannon and Simpson diversity indices across streams. Each colour represents samples from an individual stream. Native streams: CS (Cascades) and WB (Wairoa); Moderately impacted: MH (Mahurangi) and NK (Ngakaroa); Highly impacted: OTR (Otara) and OAK (Oakley). A Tukey post hoc test revealed significantly different data groups, represented by different letters assigned to each stream; the same letter represents no significant difference between streams (two factor ANOVA, Tukey HSD; \( p < 0.05 \)). The centre line in the boxplot represents median data value, top and bottom lines in the box represent the 25th and 75th percentiles; the top and bottom whisks mark the 10th and 90th percentiles respectively. The points at the top and bottom of the box indicate outlying data points

**Supplementary Material 2.** PERMANOVA table of results of model ‘1’ (see Fig. 1), based on Bray–Curtis dissimilarity distance showing the partitioning of variance factors and tests for the factors of (CA) Catchment, (ST) Stream, (YE) Year, (MO) Month and (SE) Season and, where possible, their interactions.

**Supplementary Material 3.** PERMANOVA table of model ‘2’ results generated by periodic regression analysis of Bray–Curtis dissimilarity distance comparisons of bacterial community data and accompanying tests for seasonality on the bacterial community. Covariates (CoV 1 and CoV2) generated from the regression model of the 12-month cycle were used as ANOVA estimators in this model. Factors used were (CA) Dominant catchment land-use, (YE) Year, (ST) Stream, CoV1 and CoV2, and their interaction terms where possible.

**Supplementary Material 4.** Relative abundances of the top 40 most abundant bacterial orders across our stream biofilm dataset. Barplot is partitioned into six small plots (rows) representing six different streams from the beginning of sampling in February 2013 until February 2016. The x-axis represents time and each bar represents the relative abundance of the top 40 most abundant orders in each stream.

**Supplementary Material 5.** Shade plot of measured environmental factors from Feb. 2013–Feb. 2016 from our stream biofilm sites. Data were normalized for each site and constrained by month/year before making the shade plot. Each data point was normalized by subtracting the mean (across all samples) and dividing by the standard deviations of that variable using Euclidean distance, so that all variables have values within a similar range. The normalization scale is shown on the upper right of the plot. Shading intensity within the matrix indicates the transformed value of each parameter as represented by the legend on the right side of the plot. On the y-axis, streams are indicated by symbols, CS (▲), MH (▼), NK (●), OAK (●), OTR (●), WB (●) from the beginning of sampling February 2013 until February 2016. Lines are added to the figure to separate data from each stream.

**Supplementary Material 6.** Map showing the study region and locations of sampling sites within the Auckland Region of New Zealand. The geographic location of each stream is as follows: Cascades Stream, Latitude: -36.89, Longitude: 174.51; Wairoa Stream, Latitude: -37.08, Longitude: 174.95; Mahurangi Stream, Latitude: -36.47, Longitude: 174.73; Ngakaroa Stream, Latitude: -37.11, Longitude: 174.94; Oakley Creek, Latitude: -36.88, Longitude: 174.70; Otara Stream, Latitude: -36.96, Longitude: 174.87.

**Supplementary Material 7.** Site description, location, catchment area, land use distribution of the upstream catchment area (%), and closest climate stations to each sampling site in the Auckland Region (according to the climate database www.climo.niwa.co.nz)

**Supplementary Material 8.** nMDS plot showing annual and seasonal trends followed by bacterial communities using significant physicochemical parameters highlighted in Table 2. Bacterial community data averaged by (a) year (Model 1-linear; Year 2016 was removed due to having only two data points), (b) month and year (Model 2-bloom), (c) month (Model 3-seasonal).