

ORIGINAL ARTICLE

Interactions between hydrology and water chemistry shape bacterioplankton biogeography across boreal freshwater networks

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Disentangling the mechanisms shaping bacterioplankton communities across freshwater ecosystems requires considering a hydrologic dimension that can influence both dispersal and local sorting, but how the environment and hydrology interact to shape the biogeography of freshwater bacterioplankton over large spatial scales remains unexplored. Using Illumina sequencing of the 16S ribosomal RNA gene, we investigate the large-scale spatial patterns of bacterioplankton across 386 freshwater systems from seven distinct regions in boreal Québec. We show that both hydrology and local water chemistry (mostly pH) interact to shape a sequential structuring of communities from highly diverse assemblages in headwater streams toward larger rivers and lakes dominated by fewer taxa. Increases in water residence time along the hydrologic continuum were accompanied by major losses of bacterial richness and by an increased differentiation of communities driven by local conditions (pH and other related variables). This suggests that hydrology and network position modulate the relative role of environmental sorting and mass effects on community assembly by determining both the time frame for bacterial growth and the composition of the immigrant pool. The apparent low dispersal limitation (that is, the lack of influence of geographic distance on the spatial patterns observed at the taxonomic resolution used) suggests that these boreal bacterioplankton communities derive from a shared bacterial pool that enters the networks through the smallest streams, largely dominated by mass effects, and that is increasingly subjected to local sorting of species during transit along the hydrologic continuum.

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Introduction

The widespread use of high-throughput sequencing approaches in microbial ecology studies is resulting in unprecedented insight into the structure of prokaryotic communities. We now know that the taxonomic composition of bacterial communities differs greatly between types of ecosystems (Pommier *et al.*, 2006; Lozupone and Knight, 2007; Barberán and Casamayor, 2010) and within ecosystems at different temporal and spatial scales (Gilbert *et al.*, 2012; Jones *et al.*, 2012; Shade *et al.*, 2013; Cottrell *et al.*, 2015). In general, these differences are driven by two major mechanisms: (1) dispersal of microorganisms among communities, which determines the bacteria that can be transported

and potentially colonize a given ecosystem, and (2) selection of species (that is, species sorting) from within this immigrant pool on the basis of their environmental preferences and local conditions (Martiny *et al.*, 2006; Hanson *et al.*, 2012; Lindström and Langenheder, 2012; Székely *et al.*, 2012).

Although there is ample evidence supporting a major role of environmental sorting in shaping bacterial community structure across many different habitats types (Van der Gucht *et al.*, 2007; Logue and Lindström, 2010; Langenheder and Kely, 2011; Logares *et al.*, 2013; Adams *et al.*, 2014; Souffreau *et al.*, 2015), in freshwater ecosystems the influence of mass effects (that is, the structuring of bacterial communities by massive advection of bacteria from other systems) can be particularly important because the flow of water through the landscape often results in strong connectivity and intense exchange of microbes among local communities (Lindström *et al.*, 2006; Crump *et al.*, 2007, 2012; Nelson *et al.*, 2009; Ruiz-González *et al.*, 2015b). It is thus likely that both the environmental and hydrological conditions interact to shape the patterns that we

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observe in freshwater bacterioplankton communities, but how this interaction influences the biogeography of bacterioplankton over large spatial scales remains largely unexplored.

At the catchment scale, the movement of water within the network has been shown to have a key role in modulating the relative importance of species sorting and mass effects in freshwater bacterioplankton community assembly: whereas the rate of water (and thus bacterial) inputs relative to the ecosystem volume determines the intensity of the mass effects, the resulting water residence time (WRT) in turn influences the extent of species sorting by local conditions by bracketing the time available for bacterial growth (Crump *et al.*, 2004, 2007; Lindström and Bergström, 2004; Lindström *et al.*, 2006). In addition, because water and bacteria move directionally, most ecosystems act both as recipients and sources of bacteria, and the immigrant pool arriving to a given ecosystem will also vary depending on its position along the hydrologic continuum (Nelson *et al.*, 2009). The local bacterioplankton community composition within a given aquatic network will thus be determined by the hydrology, the position of the system in the network, and the local environmental conditions, and consequently, the relative importance of the community assembly processes will likely vary along the hydrologic continuum.

Beyond individual catchments and over broader spatial scales, however, bacterial taxonomic composition may further differ between communities as a result of increases in environmental ranges relative to the local conditions, but also due to dispersal limitation among different geographic regions (for example, Lear *et al.*, 2013; Liu *et al.*, 2015), which will determine the composition of the regional species pool. Thus, the large-scale spatial patterns in composition of freshwater bacterioplankton communities likely result from combinations of these local and regional processes, as conceptualized in Figure 1. For example, we may detect differences in the composition of bacterial communities inhabiting ecosystems with long WRTs (for example, lakes) either because they are located in areas that do not share the same regional bacterial pool because of low dispersal (Figure 1a), or because their environmental differences may have selected different taxa from a shared pool of bacteria (Figure 1b). Similarly, communities subjected to strong mass effects, (for example, inhabiting headwater streams with short residence times) may show differences in composition if influenced by distinct bacterial pools (Figure 1a), but may be compositionally similar if derived from a common immigrant pool (Figure 1b), regardless of their local environmental conditions. An important corollary of this conceptual framework is that disentangling the major processes underlying the spatial patterns in bacterial community composition requires incorporating the full range of variation in terms of ecosystem position, hydrologic regime and environmental conditions within and across networks located in distinct regions.

Most studies to date, however, have covered only limited portions of this multidimensional space. For example, the literature abounds in studies focusing on a single type of aquatic ecosystem (for example, only lakes, only rivers; Figure 1—scenarios 2 and 3) varying across spatial scales and environmental gradients (Yannarell and Triplett, 2005; Fierer *et al.*, 2007; Van der Gucht *et al.*, 2007; Heino *et al.*, 2014; Ren *et al.*, 2015). These studies typically conclude that local conditions have a major role in shaping local community composition through species sorting. On the other hand, studies that have included multiple ecosystems (for example, streams, rivers, lakes; Figure 1—scenario 1), which may cover wide ranges of network positioning and hydrologic conditions but that are restricted to small spatial scales (for example, a single catchment, Lindström *et al.*, 2006; Crump *et al.*, 2007, 2012; Nelson *et al.*, 2009; Read *et al.*, 2015), often conclude that hydrology and the associated mass effects are major drivers of community assembly. The framework presented in Figure 1 suggests that the ambiguous and sometimes contradictory conclusions found in the literature concerning the main drivers of the spatial variability in aquatic bacterial community composition are neither incorrect nor mutually exclusive, but rather represent fragments of a more complex biogeographic reality. Understanding the mechanisms underlying this large-scale biogeography thus requires a cross-regional, whole network perspective that has seldom, if ever, been undertaken.

Here we explore the large-scale spatial patterns in freshwater bacterioplankton community composition across the boreal biome of Québec, and we infer its main underlying drivers and mechanisms by exploring the variability in bacterial taxonomic structure over broad ranges in hydrologic and environmental conditions, network position, and geographic distance. We have characterized the taxonomic composition of 386 bacterioplankton communities inhabiting freshwater ecosystems positioned along the entire aquatic continuum, ranging from the smallest headwater streams to the largest rivers and lakes, which belong to seven different geographic regions that span large gradients in environmental, climate and landscape properties. This sampling design allowed us to understand how local environmental and hydrologic conditions, network position, and dispersal-related regional differences interact to shape bacterial community composition across these complex boreal aquatic networks.

Materials and methods

Study sites and sampling design

Water samples were collected from 198 lakes and 188 rivers across seven boreal regions of Québec (Canada) between 2009 and 2013. The regions sampled (Laurentides, Abitibi, James Bay, Saguenay, Chibougamau, Schefferville, La Côte-Nord; Figure 2a)

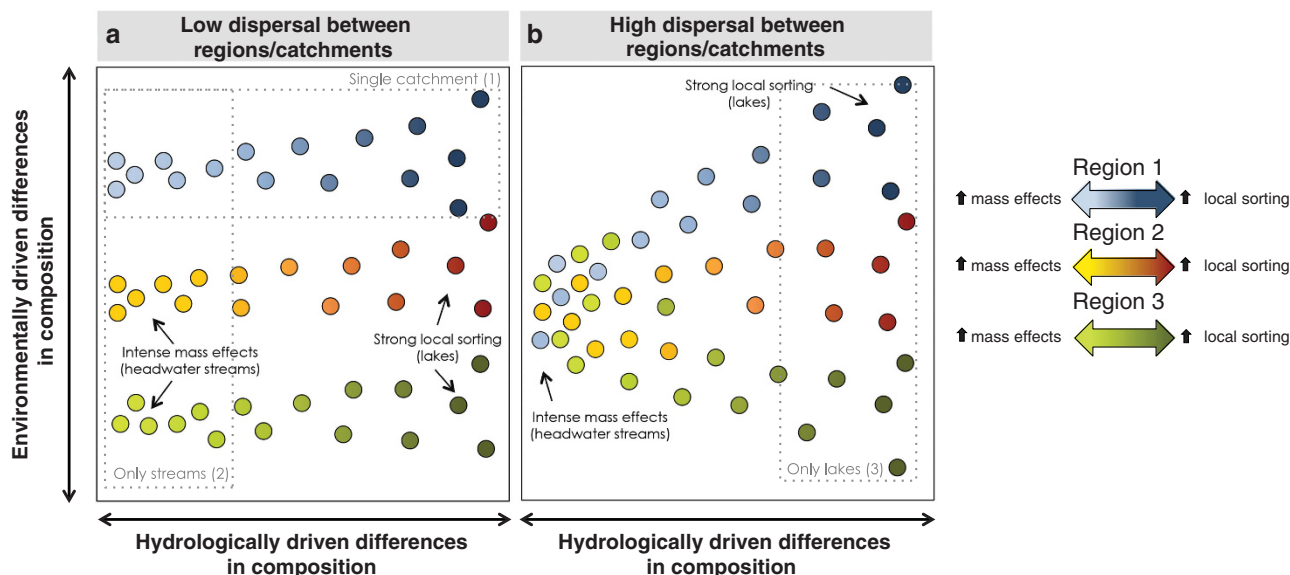


Figure 1 Schematic representation of potential large-scale spatial patterns in bacterioplankton taxonomic composition across freshwater networks, where dots represent individual bacterioplankton communities in an ordination space, based on their compositional dissimilarity, and colors indicate the different catchments or geographic regions to which they belong. Owing to the movement of the water in the landscape, the composition of local bacterioplankton communities within a single catchment or region will differ depending on their position along the hydrologic continuum (horizontal axis), as they will be differentially affected by hydrology that regulates the mass effects versus environmental sorting ratio (see Introduction section for further explanation), represented here by the light to dark color gradient. As a result, the compositional variation because of local conditions (vertical axis) will likely increase along the hydrologic continuum because of a gradual intensification of the local sorting of species with increasing WRT. Over broader spatial scales, the increase in environmental differences will likely result in stronger environmentally driven dissimilarities between communities (that is, along the vertical axis), but the overall large-scale spatial patterns may further differ depending on the degree of dispersal of microbes between regions or catchments: For example, in a case of low dispersal between regions (a), a regional structuring of the communities may be detected even in systems with little or no environmental sorting (that is, headwater streams) because the regional bacterial pools are different due to dispersal limitation. In contrast, under a scenario of high dispersal between regions (b), a common bacterial pool will lead to little environmentally driven compositional differences in the headwaters but to an increasing differentiation of communities toward downstream systems (that is, lakes) because of stronger local sorting of species. From this perspective, whereas communities located at the headwaters of the aquatic continuum will mostly reflect the nature of the regional bacterial pools because of strong mass effects, communities located further downstream in the network will be the result of species sorting by the local aquatic conditions. The interpretation of the factors and mechanisms underlying the observed spatial patterns in bacterial community composition will thus depend on the portion of this complex space that is considered (dotted areas, scenarios 1, 2 and 3). For example, studies based on a narrow spatial scale, as depicted in scenario 1, may conclude that hydrology is the main driver of changes in bacterioplankton community composition. In contrast, studies that cover broader environmental gradients but are limited to certain portions of the hydrologic continuum (scenarios 2 and 3) may conclude that either dispersal limitation (2) or local sorting (3) shape local community assembly. We argue that a cross-regional, whole network approach is necessary to disentangle the influence of the mechanisms and factors that are actually influencing the local assembly of bacterioplankton communities in complex freshwater networks.

cover a total area of *ca* 900 000 km² (44–56°N, 64–80°W) and span large gradients in terms of climate, limnological properties, land cover and geomorphology typical of the boreal zone (for details, see Lapierre *et al.*, 2015; Rasilo *et al.*, 2015; Ruiz-González *et al.*, 2015a). The studied sites covered the entire range in lake area (0.002–4345 km²) and river order (Strahler order 0–8), and were sampled once during the summer (July/August). All the sites in each region were sampled within the same year. Samples were taken at a depth of 0.5 m at the deepest measured spot of lakes and near the shore in rivers. Temperature, dissolved oxygen, pH and conductivity were measured *in situ* with a YSI probe. Samples were filtered *in situ* through 0.45 µm and stored in acid-washed glass vials for dissolved organic carbon (DOC) and optical analyses, or kept in the dark in acid-rinsed bottles for further processing.

Chlorophyll, chemical and optical analyses

Chlorophyll *a* concentration was determined spectrophotometrically in hot ethanol extracts (90%). DOC concentration was measured on an OI 1010, OI Analytical (College Station, TX, USA). Total phosphorus (TP) and nitrogen (TN) concentrations were analyzed after persulfate and alkaline persulfate digestions, respectively, following standard methods (see Rasilo *et al.*, 2015).

The optical properties of DOC were measured as indices of its composition. Colored dissolved organic matter (cDOM) was quantified as the absorbance at 440nm using an Ultrospec 3100 spectrophotometer (Biochrom, Cambridge, UK). DOC composition was described on the basis of fluorescence absorption/emission spectra (EEMS), measured in a Shimadzu RF5301 PC spectrofluorophotometer (Biochrom) across excitation/emission wavelengths of 275–450

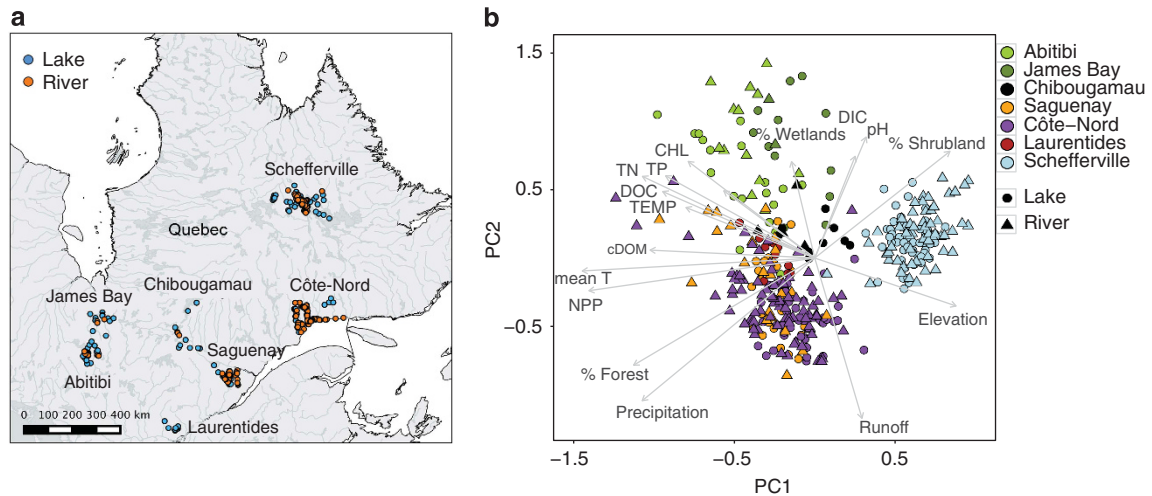


Figure 2 (a) Distribution of sampling sites across the seven sampled regions in Northern Québec (Canada). (b) Principal component analysis (PCA) of the sites based on the measured environmental and geographic parameters. Different regions are indicated by different colors and symbols indicate rivers (triangle) or lakes (circles). The two first axes explain 44.5% of the variance. Chl *a*, chlorophyll *a*; Cond, conductivity; DIC, dissolved inorganic carbon; %C2, percentage of fluorescent component C2 (humic-like DOM); %C5, percentage of fluorescent component C5 (freshly produced labile DOM); Elevation, mean catchment elevation; Mean T, mean annual temperature; NPP, net primary productivity; Precipitation, mean annual precipitation; Runoff, mean annual runoff; %Forest/%Shrubland/%Wetland, %forest/shrubland/wetland covered area in the catchment; Temp, water temperature; TP;TN, total phosphorus and nitrogen.

and 280–600 nm, respectively. Six main fluorescence components related either with refractory-humic (C1–C3) or with biolabile, freshly produced DOM (C4–C6) material were recovered from the EEMS using parallel factor analysis (PARAFAC, for details, see Lapierre and del Giorgio, 2014). The percent contribution of each component was calculated relative to the total fluorescence of the six PARAFAC components.

Geographic analyses

Lake areas, river length and order, catchment areas, the area covered by water in the catchment, and elevation of the sampled sites were derived using the ArcMap 10 and ArcGIS V10 software (ESRI Inc., Redland, CA, USA) applied on the digital elevation models derived from (1:50 000) maps. For each catchment area, we averaged catchment slope based on digital elevation models, as well as various land cover properties obtained from Geobase (2009). Mean annual temperature, precipitation and runoff were extracted at each site location from a long-term climate database (WorldClim, Hijmans *et al.*, 2005). The WRT of lakes was estimated from lake volume (as mean depth \times lake area), catchment area, and mean annual regional runoff. River WRT was calculated based on the measured water velocity at the sampling point and the total upstream distance.

Water isotopes

Samples were collected to determine ^{18}O and ^2H isotopes by laser spectroscopic analysis of liquid water technique (LGR DT-100 Liquid Water Stable Isotope Analyzer, Los Gatos Research Inc., Mountain View, CA, USA). We used water isotopic

composition to calculate the deuterium excess (*d*-excess), as:

$$d - \text{excess} = \delta^2\text{H} - 8 \times \delta^{18}\text{O}$$

which provides an index of the degree of evaporation relative to precipitation, and therefore, the relative proximity to groundwater sources. As headwater systems are strongly influenced by soil groundwater, the water carries an isotopic signature that closely resembles that of precipitation (high *d*-excess). Given that temperature increases and *d*-excess decreases as water transits through the network and is subjected to warming and evaporation (Gibson *et al.*, 2002; Turner *et al.*, 2014), here we use water temperature and *d*-excess as proxies of the groundwater-mediated connectivity with the surrounding catchment.

Bacterial community composition

In all, 300–500 ml water samples were filtered onto 0.22 μm pore-size filters, and genomic DNA was extracted from the filters using the MoBio PowerWater DNA extraction kit (Mo Bio, Carlsbad, CA, USA) following the manufacturer's protocol. Libraries of the V3–V4 region of the 16S ribosomal RNA were made using the primers 515F and 806R, and sequenced on an Illumina MiSeq2000 (Illumina, San Diego, CA, USA) following a paired-end approach (Caporaso *et al.*, 2012). Paired-end reads were assembled with FLASH (Magoč and Salzberg, 2011) and sequences between 250 and 290 bp were used for downstream analyses in QIIME to remove primers and low-quality, archaeal and chloroplast reads (Caporaso *et al.*, 2010). After the detection and removal of chimeric sequences with ChimeraSlayer (Haas *et al.*, 2011), quality sequences were aligned with Mothur aligner (Schloss *et al.*, 2009) and clustered into operational

taxonomic units (OTUs, $\geq 97\%$ similarity) using UCLUST (Edgar, 2010). In order to assign the taxonomy to the representative sequences, we used the RDP classifier and the Ribosomal Database Project taxonomy (Wang *et al.*, 2007). We discarded all OTUs that were represented in < 10 samples and/or by < 10 sequences. To enable comparisons between samples, the OTU table was randomly subsampled to ensure an equal number of sequences per sample, based on the sample with the least number of reads (50 323 sequences). The sequences have been deposited in the European Nucleotide Archive, under accession number PRJEB11530.

Statistical analyses

We used nonmetric multidimensional scaling (Vegan metaNMDS, Oksanen *et al.*, 2015) with Bray–Curtis distances to ordinate the samples based on their dissimilarity in community composition, and the Vegan envfit function (Oksanen *et al.*, 2015) to fit environmental vectors onto the ordination space. Differences in environmental and taxonomic composition among categories (that is, regions, ecosystems) were tested using PerMANOVA with 999 permutations (Anderson, 2006; Anderson, 2011). To estimate the relative contribution of environmental and hydrological factors in explaining the variation of the large-scale patterns of bacterial community composition, we performed a variation partitioning analysis using the non-metric multidimensional scaling (NMDS) scores as the response variables. We grouped the variables as environmental or hydrological factors, which, after a forward selection procedure, were used as the explanatory groups of variables (Borcard *et al.*, 2011). All the analyses were made using the R 3.0.0 software (R Core Team, 2013).

Results

Large-scale patterns in bacterial community composition across boreal aquatic ecosystems

The studied lakes and rivers segregated into clearly differentiated groups of sites based on their environmental, climatic and landscape properties (Figures 2a and b). These groups roughly corresponded to the different geographic regions (PerMANOVA_{BY_REGION} $R^2 = 0.43$, $P < 0.01$), although systems located in Laurentians, Côte-Nord and Saguenay tended to overlap. Sites were segregated among these regional groups based on differences in water chemistry (pH, DOC, dissolved inorganic carbon and cDOM) and trophic status (TP, TN and chlorophyll *a*), and lakes and rivers from a given region tended to cluster together. Briefly, the sites varied along a gradient from alkaline and more productive waters in Abitibi and Bay-James to acidic and less productive waters in Saguenay and Côte-Nord. These environmental gradients were also related with changes in mean annual temperature and regional net primary productivity (as illustrated by the PC1 axis), as well as mean annual precipitation and runoff (PC2).

This environmental segregation of sites was not as clearly reflected in terms of the composition of bacterial communities, and the NMDS analysis showed much smaller differences between the geographical regions on the basis of bacterial composition (Figure 3a, PerMANOVA_{BY_REGION} $R^2 = 0.10$, $P < 0.01$), and a significant overlap in composition between sites from different regions (Figure 3a). River and lake communities, in contrast, showed a clearer segregation, although there was some degree of overlap between them (Figure 3, PerMANOVA_{BY_ECOSYSTEM} $R^2 = 0.11$, $P < 0.01$), as the largest rivers (Strahler order > 4) clustered together with lakes (details not

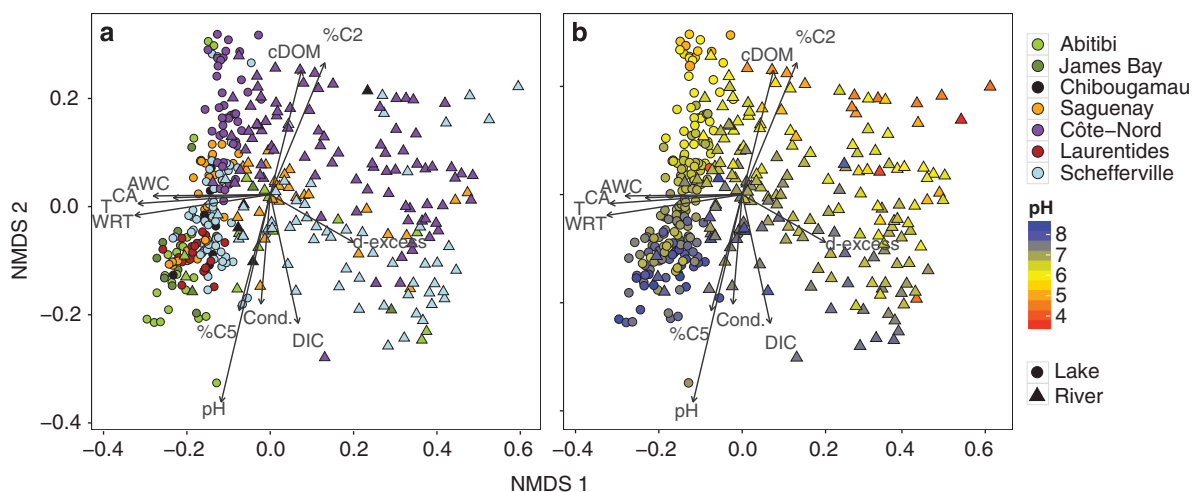


Figure 3 NMDS ordination of bacterioplankton communities based on the Bray–Curtis dissimilarity of community composition (stress = 0.13). Shape indicate rivers (triangles) or lakes (circles), and sites are colored according to geographical region (a) and water pH (b), which was the variable that best fitted the ordination space ($R^2 = 0.61$, see Table 1). The arrows indicate the direction at which the environmental and hydrological vectors fit the best (using envfit function) onto the NMDS ordination space. The size of the arrow is proportional to the strength of the correlation of each variable. AWC, area of water in the catchment; CA, catchment area; Cond, conductivity; DIC, dissolved inorganic carbon; %C2, percentage of fluorescent component C2; %C5, percentage of fluorescent component C5 (freshly produced labile DOM); T, water temperature.

shown). Interestingly, river communities were much more dispersed along the NMDS1 axis than lake communities (Figure 3).

Drivers of bacterial community composition across freshwater ecosystems

In order to identify the drivers explaining the spatial patterns in community composition depicted by the NMDS in Figure 3, we fit the measured environmental and landscape variables onto the ordination. pH, WRT and water temperature were the variables most strongly correlated to the two NMDS axes (Figure 3, Table 1). NMDS1 was strongly correlated to hydrologic and landscape variables such as water temperature, WRT, d-excess, catchment area and the area of water in the catchment. This NMDS axis also represented to some extent network positioning, with scores declining with increasing river order along the hydrologic continuum, and also with increasing distance to the headwaters, thus separating small headwater streams from rivers, and lumping together the largest rivers with lakes (details not shown). NMDS2 was mostly related to water chemistry, and particularly to pH (but also to cDOM, dissolved inorganic carbon and specific DOM fluorescence components; Figure 3, Table 1).

The geographic location (latitude and longitude) was not significantly related to either of the NMDS axes, and there was no significant relationship between the spatial distance and the taxonomic dissimilarity between sites (Partial Mantel $R = 0.06$ $P = 0.1$; Figure 4a), thus indicating that the observed regional segregation described above (Figure 3a) was not primarily driven by dispersal limitation. Rather, the community dissimilarity was significantly correlated to both the environmental and hydrologic distances between pairs of sites (Figures 4b and c, respectively). Environmental and hydrologic distances were only weakly

Table 1 Envfit results and range of the environmental and hydrological variables included in the analysis shown in Figure 3

	NMDS1	NMDS2	R ²	P-value	Range
Temp (°C)	-0.99881	-0.0488	0.401	0.001	6.24 to 32.3
WRT (days)	-0.99124	-0.13205	0.4185	0.001	0.01 to 10 523
geographic km ²	-0.99853	-0.05429	0.3162	0.001	0.002 to 12 753
CA (km ²)	-0.99484	-0.10143	0.2177	0.001	16 to 55 148
d-excess	0.92625	-0.3769	0.1889	0.001	-5.44 to 37
pH	-0.29484	-0.95555	0.6178	0.001	3.3 to 8.95
cDOM	0.31115	0.95036	0.2318	0.001	0.01 to 58.8
DIC	0.29063	-0.95684	0.2285	0.001	0.282 to 42.2
%C2	0.48359	0.8753	0.2873	0.001	0.13 to 0.48
%C5	-0.32545	-0.94556	0.2047	0.001	0.05 to 0.25

Abbreviations: CA, catchment area; cDOM, colored dissolved organic matter; %C2, percentage of fluorescent component C2 (humic-like DOM); %C5, percentage of fluorescent component C5 (freshly produced labile DOM); DIC, dissolved inorganic carbon; T, water temperature; WRT, water residence time.

The degree of fit within the whole ordination space (R^2) and for each ordination axis (NMDS1, NMDS2) are shown.

correlated to each other (details not shown), such that any given range of hydrologic conditions was associated to the entire range in environmental variability, suggesting that our sampling scheme captured the full range in both dimensions that exists across the boreal biome.

Changes in taxonomic diversity and composition along the hydrologic continuum

In order to further understand the observed changes in community composition along the hydrologic continuum, we divided the sampled sites into seven groups (G1–G7) that were equally spaced along the

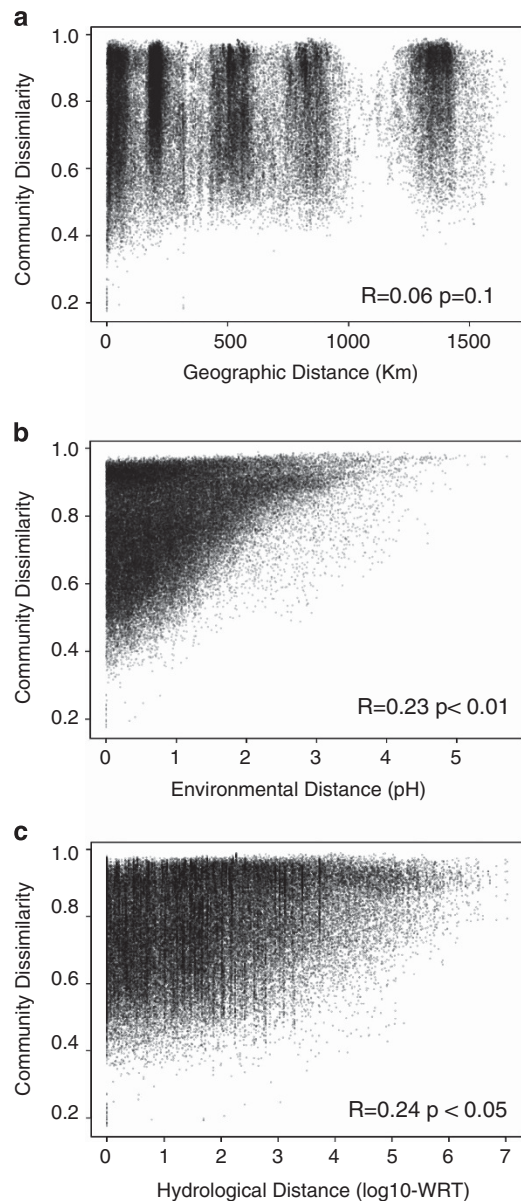


Figure 4 Relationship between geographic (a), environmental (b) and hydrological (c) distances and pairwise bacterial community dissimilarity (Bray–Curtis). pH and WRT (\log_{10} -WRT) were used as proxies of environmental and hydrological distances, respectively. Mantel correlations (R) and the probabilities are provided for each case.

NMDS1 axis. These groups were associated to different average network position and WRT within the hydrologic continuum: most lakes were contained within groups G1 and G2, the smallest headwater streams were mostly included in G7 and the remainder of the rivers and some lakes were distributed among the other groups. For each of the seven groups, we calculated the average OTU richness, average Pielou's evenness index and the percentage of sequences within the major bacterial Phyla. This analysis revealed that sites with the highest scores on the NMDS1 axis (G7, smallest streams) showed the highest average OTU richness (Figure 5a) and evenness (Figure 5b), and that there was a striking decline in both indices toward sites with the lowest average NMDS1 scores (G1, larger rivers and lakes; Figures 5a and b): on average, 6000 OTUs per community were lost between these two groups of sites, and the evenness decreased from 0.86 to 0.66. The taxonomic composition also changed gradually along this NMDS1 axis, resulting in highly differentiated communities at the extremes of this hydrologic continuum even at the phylum level (Figure 5c). Overall, although Actinobacteria, Bacteroidetes and Cyanobacteria increased toward larger rivers and lakes, Betaproteobacteria increased toward larger rivers but decreased again in lakes. The remaining classes of Proteobacteria and dominant soil groups such as Acidobacteria systematically decreased from the smallest streams to lakes.

Shifts in the relative influence of the environment versus hydrology on bacterial community composition

The pronounced loss of OTUs downstream the hydrologic continuum, together with the decline in taxonomic evenness associated to the dominance of certain bacterial groups (Figure 5c), suggest that the relative importance of mass effects and environmental sorting of species may change along the hydrologic continuum. To test this hypothesis, we calculated the correlation between taxonomic and environmental dissimilarity matrices for each of the seven groups of sites along the NMDS1 axis (G1 to G7, described above; Figure 6a). We used differences in pH as a proxy of environmental distance because it was the variable most strongly correlated to changes in taxonomic composition along the NMDS2 axis (Figure 3, Table 1). The correlation between the environmental and the compositional dissimilarities between sites were weakest in the groups with the shortest average WRTs and highest position within the network (that is, smallest headwater streams, G7, Mantel $R=0.25$), and increased gradually toward larger rivers and lakes, where changes in pH explained a much larger portion of the variability in taxonomic composition (G1, Mantel $R=0.49$; Figure 6a).

To explore whether there are thresholds along the hydrologic continuum that modulate the relative importance of hydrology and environment sorting,

we analyzed the changes in NMDS1 scores as a function of WRT (Figure 6b), and found that above a 10-day threshold, an increase in WRT did not trigger further changes in community composition along the NMDS1 axis. Thus, in order to compare the relative influence of the environmental and hydrologic conditions on the dissimilarity between communities within the NMDS ordination space, we performed a variation partitioning analysis dividing the data set into two groups of sites based on whether their estimated WRT was >10 or <10 days (Figure 6c). This exercise clearly shows that environmental variables explained most of the variation (65%) in bacterial taxonomic composition at those sites with high WRT (>10 days), whereas hydrological variables explained a larger share of the variation (33%) than the environment (14%) in sites with shorter WRT (Figure 6c).

Discussion

Our results show that environmental sorting is a major driver of bacterial community assembly across the boreal biome, but further indicate that hydrology and network position, and the associated mass effects, also have a fundamental role in shaping bacterioplankton communities in complex aquatic networks. Although previous studies had identified these two processes as major mechanisms of freshwater bacterial community assembly (Crump *et al.*, 2007; Fierer *et al.*, 2007; Nelson *et al.*, 2009; Ren *et al.*, 2015; Souffreau *et al.*, 2015), no study to date had explored how they interact to shape the large-scale biogeography of freshwater bacterioplankton communities. Assessing this interaction necessarily requires expanding the conceptual framework proposed by Martiny *et al.* (2006) to incorporate, in addition to the environmental gradients and geographic distance (as a proxy for dispersal limitation), a hydrologic dimension that controls the relative importance of the mass effects on community assembly, as we conceptualize in Figure 1.

Our results show no significant relationship between geographic distance and the taxonomic dissimilarity among the studied bacterioplankton communities (Figure 4a), suggesting that dispersal limitation between aquatic bacterial communities is negligible across boreal Québec, at least at the taxonomic resolution of our analysis. Rather, there was an overall positive relationship between environmental and taxonomic distances (Figure 4b), and communities appeared to be strongly structured by local water chemistry (mostly pH; Figure 3b), pointing to a major role of environmental sorting in structuring the large-scale biogeography of boreal freshwater bacterioplankton. We should note that although pH emerged as the most important environmental variable, in agreement with previous studies (Fierer *et al.*, 2007; Ren *et al.*, 2015; Ruiz-González *et al.*, 2015b), it is likely that besides

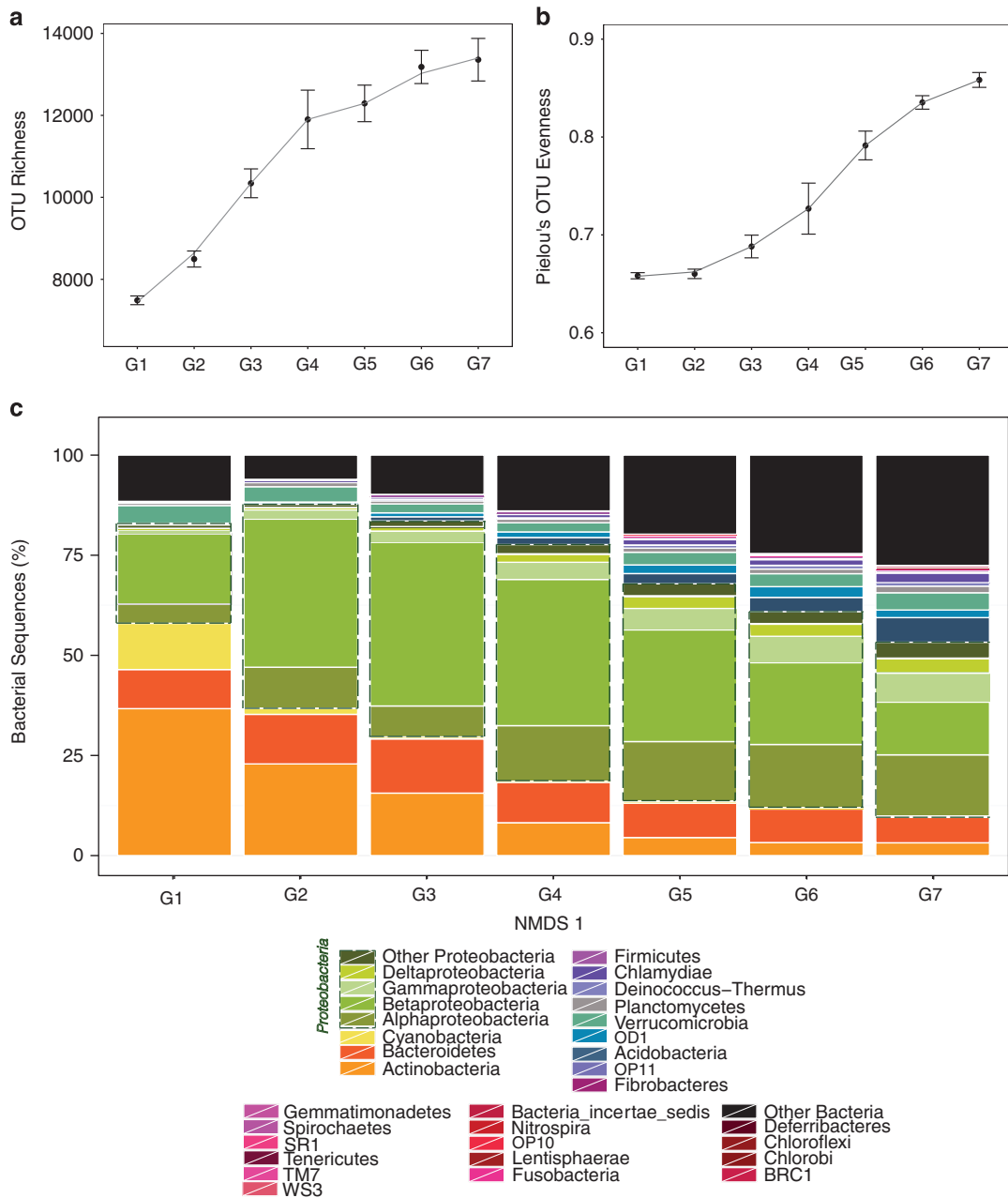


Figure 5 Changes in OTU richness (a), Pielou's evenness (b) and taxonomic composition (at the phylum level, c) for sites grouped according to their position on the NMDS1 axis (G1 to G7, $n = 174, 74, 37, 18, 26, 39, 18$), which mostly represents an hydrologic continuum from the smallest headwater streams (G7) to larger rivers and lakes (G1, for details see Results section). Dots are means and error bars represent the standard error of the values for the sites within each NMDS1 group. Colors indicate different phyla (and different classes within the Phylum Proteobacteria, indicated by the dashed line) and the heights of the bars represent the percentage of sequences associated to each taxonomic rank relative to the total number of sequences within each group of sites.

directly influencing cell physiology, pH is also an integrator of other variables of relevance for bacteria (Fierer *et al.*, 2007). In particular, we found that changes in water pH were strongly correlated to shifts in features linked to the amount and quality of DOM (details not shown), also known to shape bacterial community structure and functioning (Fujii *et al.*, 2012; Ruiz-González *et al.*, 2015a; Wilhelm *et al.*, 2015).

Despite this strong effect of pH, the fact that in environmentally similar sites we observed the entire

range of community dissimilarity (Figure 4b) suggests that mechanisms other than environmental sorting are involved in shaping the large-scale biogeography of boreal freshwater bacterioplankton communities. In this regard, our results show that WRT alone explains as much variability in composition as pH (Figures 4b and c), and that bacterial communities change largely along a hydrologic continuum from headwater streams toward large rivers and lakes, following variations in WRT and network position (Figures 3 and 5).

We hypothesize that this directional structuring is caused by a gradient of decreasing mass effects and increasing environmental sorting toward downstream ecosystems as conceptualized in Figure 1. In a companion study where we characterized the soil communities surrounding aquatic systems within a single boreal region (Ruiz-González *et al.*, 2015b), we show that whereas headwater streams are strongly dominated by bacteria washed from soils, which gradually disappear along the hydrologic continuum, downstream ecosystems are numerically dominated by a few of these soil-derived taxa that have the potential to grow in the aquatic

environment. The high OTU richness found in our headwater streams across the boreal biome (Figure 5a) thus likely results from the massive immigration of terrestrial bacteria (Besemer *et al.*, 2013; Savio *et al.*, 2015). Moreover, we found that variations in d-excess (our proxy of groundwater-mediated connectivity with land) also explained changes in community composition (Figure 3b), supporting that these mass effects depend on the degree of connectivity to the source of immigrants (that is, the soils). In turn, the observed loss of bacterial taxa toward downstream ecosystems was accompanied (and probably caused) by gradual increases in the relative abundances of some common freshwater bacterial groups (*sensu* Newton *et al.*, 2011, see examples in Supplementary Figure S1), suggesting that environmental sorting progressively increases along the hydrologic continuum. In support of this idea, and in accordance to our conceptual framework (Figure 1), we observed that the spatial patterns of the bacterial assemblages inhabiting systems with longer WRT were much more strongly influenced by differences in local aquatic conditions than those from communities inhabiting headwater streams (Figure 6a).

Interestingly, the relative influence of hydrology and the associated mass effects on community composition was not continuous: we observed a clear threshold of WRT of around 10 days, beyond which increases in WRT did not result in further hydrologically driven changes in taxonomic composition (Figure 6b). Our variance partitioning analysis supports this hypothesis, demonstrating that hydrology explains a much larger fraction of the variance in taxonomic composition in systems below this WRT threshold, whereas environmental factors were

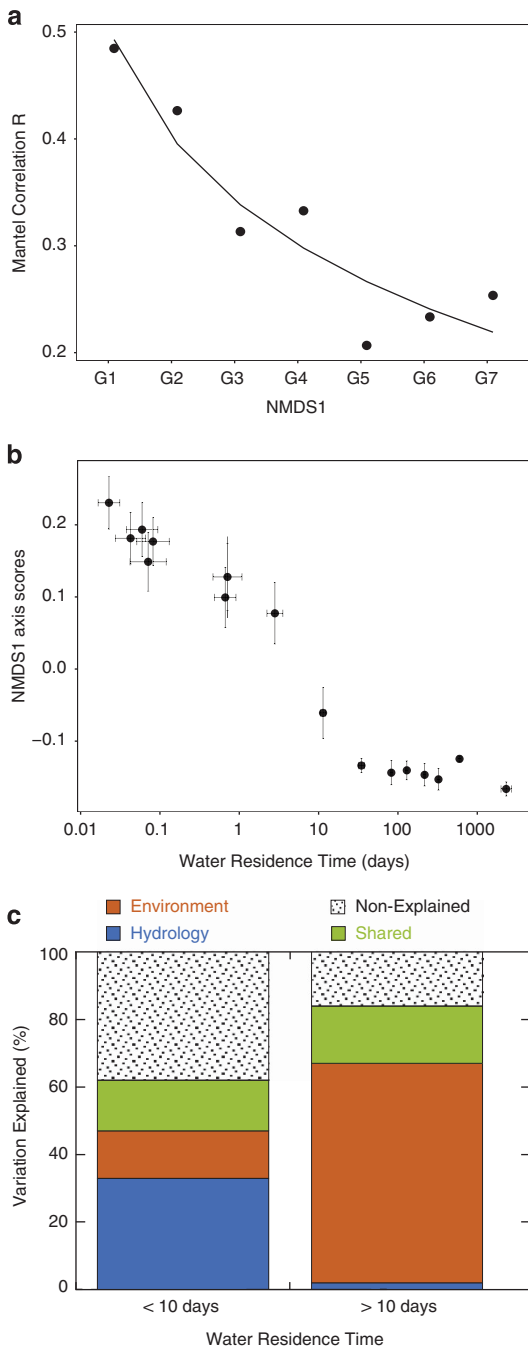


Figure 6 (a) Variation in the R coefficient of the Mantel correlations between the taxonomic and the environmental (pH) dissimilarity matrices for sites grouped according to their position on the NMDS1 axis (G1 to G7). Note that G1 includes most lakes and the largest rivers, and G7 (G1 to G7, $n=174, 74, 37, 18, 26, 39, 18$) contains mostly small headwater streams (for details see Results section). All the correlations were significant ($P < 0.01$). (b) Changes in bacterial community composition along the NMDS1 axis as a function of WRT. The NMDS1 scores of the sites were binned into 16 equal groups ($n=20$) based on ranked WRT. The dots are the average NMDS1 scores within each bin and represent the mean position of the sites within a given bin on the NMDS1 axis. Error bars are the standard errors for either WRT and scores within each bin. Note that increases in WRT result in gradual changes in taxonomic composition along the NMDS1 axis until a WRT of 10 days is reached, above which increases in WRT do not translate into further changes in taxonomic composition along the NMDS1 axis. (c) Percentages of variation in the NMDS ordination space explained by the environment and hydrology, for sites with WRTs above and below 10 days (WRT > 10 and WRT < 10, respectively). Non-explained and shared variation (environment+hydrology) are also shown. The 'environment' category included the variables DIC, dissolved inorganic carbon; %C2, percentage of fluorescent component C2 (humic-like DOM); %C5, percentage of fluorescent component C5 (freshly produced labile DOM) and conductivity. T, water temperature; CA, catchment area; AWC, area of water in the catchment; WRT, and d-excess were included within the 'hydrology' category.

responsible for most of the variability in ecosystems with longer WRT (Figure 6c). In the latter systems, pH seemed to strongly influence OTU richness, which peaked at pH levels close to neutrality (Supplementary Figure S2a), a pattern that has been observed before (Fierer and Jackson, 2006; Ren *et al.*, 2015). This suggests that extreme environmental conditions may impose a stronger filter for bacteria, and that neutral pH may favor the co-existence of taxa with overlapping ranges of tolerance. This relationship was lost, however, at sites with WRT > 10 days (Supplementary Figure S2b), indicating that intense mass effects may overwhelm species sorting even under extreme environmental conditions. Remarkably, we identified a similar WRT within a regional aquatic network (Ruiz-González *et al.*, 2015b), beyond which there was no further selection or growth of soil-derived bacteria along the hydrologic continuum. This threshold is likely determined by the balance between the potential for *in situ* growth of the resident bacterial communities relative to the rate of dilution by allochthonous bacteria (for example, Crump *et al.*, 2007), but this issue needs further investigation.

Considering that the headwater systems are largely subjected to mass effects, we argue that the bacterial composition in these communities must reflect the nature of the terrestrial source communities that feed these networks. The question then becomes whether there are distinct pools of immigrant bacteria separated by dispersal barriers, or whether all aquatic communities derive from a common pool of taxa, at least at the scale of our study. The lack of a relationship between community dissimilarity and geographic distance discussed above suggests that the boreal biome represents a single bacterial province *sensu* Martiny *et al.* (2006), and thus that there is low dispersal limitation of taxa between regions. We are aware, however, that there could be dispersal limitation at finer taxonomic resolution, although in a previous study we have shown that the level of taxonomic aggregation used here (97%) resulted in the clearest spatial and temporal patterns in community composition from boreal and temperate lakes (Cottrell *et al.*, 2015).

In our conceptual framework, a common bacterial pool across all regions should lead to small differences in community structure between systems that are overwhelmingly dominated by mass effects (Figure 1b), a scenario that is not supported by our results. Rather, we observed that the taxonomic differences between systems with the lowest WRT were as large as those between lake communities (Figure 3), a pattern that resembles that of Figure 1a, where dispersal limitation has a major role. As our data suggest that this is not the case, we hypothesize that the observed differences in taxonomic composition between small headwater streams reflect a sorting that occurs in the soils before the loading of terrestrial bacteria to the aquatic network. This may indicate the occurrence of a shared pool of taxa that undergoes

re-arrangements driven by local landscape conditions. Thus, although there is evidence that even in headwater streams there is already some degree of aquatic sorting of terrestrially derived taxa (Ruiz-González *et al.*, 2015b), we argue that the communities from headwater aquatic systems that are subjected to strong mass effects must largely reflect differences in the composition of the surrounding soil assemblages. In this regard, the regional signature observed in headwater streams (Figure 3a) could easily be interpreted as evidence of dispersal limitation (Figure 1a). However, the fact that low-pH headwater streams from Schefferville clustered together with equally acidic sites in Côte-Nord (Figures 3a and b), for example, suggests that the observed patterns are the result of a regional structuring of pH (and likely other covarying factors), and therefore, of pH-driven differences in soil bacterial taxonomic composition (Fierer and Jackson, 2006). This could be considered analogous to historical or legacy effects (Martiny *et al.*, 2006) because it occurs outside the aquatic network and before the aquatic sorting, but is nevertheless based on contemporary environmental ranges and not necessarily on past events linked to dispersal limitation.

Once in the aquatic network, these migrant communities undergo further selection by local conditions as they move downstream, and especially in systems with long WRTs (Crump *et al.*, 2012; Ruiz-González *et al.*, 2015b), such that the composition of these communities will be mainly a reflection of the local aquatic environment (for example, Yannarell and Triplett, 2005; Fujii *et al.*, 2012; Ren *et al.*, 2015; Souffreau *et al.*, 2015). This process renders lake or large river communities clearly distinct from those of headwater streams (Figures 3 and 5) within a given region, even when they may share similar ranges in key environmental factors. We recognize that our results provide a midsummer snap-shot and do not incorporate the potential temporal variability within communities, but when the taxonomic composition was analyzed for a subset of rivers and lakes sampled on three occasions (spring, summer and fall, $n = 331$, data not shown), the same pattern was observed: irrespective of the season, aquatic communities were structured along an hydrologic and a pH gradient, highlighting the consistency and strength of the spatial patterns presented here.

Overall, we show that hydrology and network position interact with environmental gradients to shape the spatial patterns of bacterioplankton community composition across boreal freshwater networks. Our results indicate that hydrology and network position modulate the relative role of environmental sorting and mass effects on community assembly along the aquatic continuum, likely by defining both the time frame for bacterial growth and the composition of the immigrant pool. We found no evidence of dispersal limitation even at the large spatial scale of our study, but we observed a certain regional signature likely caused by the spatial

structuring of environmental conditions in the landscape. We suggest that this landscape structuring influences the headwater communities through a pre-sorting of the immigrant pool of bacteria, and the communities within systems with longer WRT by determining the local aquatic conditions that bacteria will face. Despite that these two processes can generate similar spatial patterns, they represent fundamentally different mechanisms, highlighting the fact that interpretations based on partial fragments of this complex biogeographic reality may lead to a biased understanding of the factors shaping bacterioplankton communities over large spatial scales.

Conflict of Interest

The authors declare no conflict of interest.

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