

1	Running head: Stream insect diversity and abundance in different climatic regions
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3	Subtropical streams harbour higher genus richness and lower abundance of insects
4	compared to boreal streams, but scale matters
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42 Abstract

44	Aim: Biological diversity typically varies between climatically different regions, and regions							
45	closer to the equator often support higher numbers of taxa than those closer to the poles.							
46	However, these trends have been assessed for a few organism groups, and the existing studies							
47	have rarely been based on extensive identical surveys in different climatic regions.							
48	Location: We conducted standardised surveys of wadeable streams in a boreal (western							
49	Finland) and a subtropical (south-eastern Brazil) region, sampling insects identically from							
50	100 streams in each region and measuring the same environmental variables in both regions.							
51	Taxon: Aquatic insects							
52	Methods: Comparisons were made at the scales of local stream sites, drainage basins and							
53	entire regions. We standardised the spatial extent of the study areas by resampling regional							
54	richness based on subsets of sites with similar extents. We examined differences in genus							
55	richness and assemblage abundance patterns between the regions using graphical and							
56	statistical modelling approaches.							
57	Results: We found that while genus accumulation and rank-abundance curves were relatively							
58	similar at the regional scale between Finland and Brazil, regional genus richness was higher							
59	in the latter but regional abundance much higher in the former region. These regional patterns							
60	for richness and abundance were reproduced by basin and local genus richness that were							
61	higher in Brazil than in Finland, and assemblage abundance that was much higher in Finland							
62	than in Brazil. The magnitude of the difference in genus richness between Brazil and Finland							

63 tended to increase from local through basin to regional scales.

64 Main conclusions: Our findings suggest that factors related to evolutionary diversification

might explain differences in genus richness between these two climatically different regions,

66 whereas higher nutrient concentrations of stream waters might explain the higher abundance

67 of insects in Finland than in Brazil.

68

69 Keywords

- 70 alpha diversity, latitudinal diversity gradient, nutrients, rank abundance, regional diversity,
- 71 stream insects, low-high latitude comparison

73 INTRODUCTION

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Biological diversity typically varies among climatically different regions, with regions closer 75 to the equator often harbouring higher numbers of species than those at higher latitudes 76 (Rosenzweig, 1995; Gaston, 2000). This latitudinal gradient in species richness is evident for 77 terrestrial organisms such as vascular plants, butterflies, birds and mammals (Hillebrand, 78 2004a). It has also been found for marine (Hillebrand, 2004b) and freshwater organisms 79 (Pearson & Boyero, 2009). However, there are also notable exceptions to the latitudinal 80 richness gradient among microbes and invertebrates, to name two highly diverse organism 81 groups (Kouki et al., 1994; Willig et al., 2003; Boulton et al., 2008; Soininen, 2012). Studies 82 83 on many infrequently studied groups of organisms have rarely used extensive standardised surveys (Vinson & Hawkins, 2003; Pearson & Boyero, 2009). These methodological issues 84 85 may complicate the examination of the drivers of regional and local richness, although there are successful classical (Stout & Vandermeer, 1975; Lake et al., 1994) and more recent 86 examples (Dias et al., 2014; McCreadie et al., 2017). In general, regional differences in 87 richness may stem from differences in spatial extent, evolutionary diversification, current and 88 past climates, productivity or history (Willig et al., 2003; Mittelbach et al., 2007), whereas 89 local richness gradients may result from differences in biological interactions and abiotic 90 ecosystem features between geographical regions (Vinson & Hawkins, 2003; Heino, 2011). 91 Stream-dwelling insects are a suitable model group for examining biodiversity 92

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patterns between climatically different regions. They comprise many taxonomic orders, of

which mayflies (Ephemeroptera), stoneflies (Plecoptera), caddisflies (Trichoptera), beetles 94 (Coleoptera) and true flies (Diptera) dominate stream insect assemblages in terms of 95 abundance and richness over the world (Vinson & Hawkins, 1998; Lancaster & Downes, 96 2013). Stream insect richness exhibits various patterns along latitudinal gradients, both at 97 regional (Boyero, 2002; Pearson & Boyero, 2009) and local scales (Stout & Vandermeer, 98 1975; Jacobsen et al., 1997; Vinson & Hawkins, 2003; Boyero et al., 2011b). While some 99 100 insect groups (e.g. dragonflies and beetles) are more diverse in tropical than in northern regions (Brown, 1981; Pearson & Boyero, 2009), other groups (e.g. mayflies, stoneflies and 101 102 caddisflies) have also diversified extensively in mid-latitude regions (Illies, 1965; Ward, 1992). The reasons for such anomalous latitudinal patterns may relate to the possibility that 103 many taxa of mayflies, stoneflies and caddisflies originated and thus diversified in cold-water 104 105 streams (Ward, 1992), following the niche conservatism hypothesis (Wiens & Donoghue, 106 2004). Alternatively, their high diversity in temperate regions may stem from high environmental heterogeneity along seasonal and altitudinal gradients (Vinson & Hawkins, 107 2003). The latitudinal gradient in the local diversity of stream insects (i.e. species richness 108 within one stream) may be even more complex than that for regional diversity (i.e. total 109 number of species from several streams in a region), as local abiotic conditions may vary 110 strongly even between neighbouring streams (Hynes, 1970; Allan & Castillo, 2007; Boulton 111 et al., 2008). Such high variation in local conditions dictates that streams close to each other 112 113 may harbour insect assemblages differing highly in richness, abundance and composition (Grönroos & Heino, 2012; Landeiro et al., 2012). Therefore, although regional-scale stream 114 insect diversity may differ (Brown, 1981; Pearson & Boyero, 2009), among-region 115 differences in local diversity may be less clear (Heino et al., 2003; Vinson & Hawkins, 2003; 116

Biological diversity at regional and local scales can be studied by using both graphical 118 pattern-based (Magurran, 2004) and modelling approaches (Legendre & Legendre, 2012). 119 Pattern-based approaches include taxon accumulation curves (Ugland et al., 2003) and rank-120 abundance distributions (Whittaker, 1965), both applicable at regional and local scales. For 121 122 example, there were only slight differences in the taxon accumulation curves among three high-latitude regions (Heino et al., 2015b). Taxon accumulation curves have also been used 123 at local scales, where the accumulation curves of temperate stream insects reached an 124 asymptote much earlier than those of tropical streams (Stout & Vandermeer, 1975). Rank-125 abundance distributions have rarely been applied in stream insect studies at both regional and 126 local scales, but they show a great potential for between-systems comparisons in other 127 contexts (McGill et al., 2007). For example, rank-abundance distributions may reveal 128 whether tropical and boreal regions harbour regional assemblages with different degrees of 129 rarity and commonness. Such differences in rarity and commonness may be due to different 130 degrees of ecological specialization, with tropical taxa being more specialized than those at 131 high latitudes (Mittelbach et al., 2007; Coley & Kursar, 2014). 132

Modelling approaches have been used extensively to examine how different predictor variables affect variation in biological diversity among regions and among sites (Legendre & Legendre, 2012). For example, based on a comparative analysis of 61 datasets from around the world, Heino *et al.* (2015a) found that stream insect assemblages were more strongly related to the environment than to space, and that the explanatory power of these predictors was low with no obvious latitudinal trend. In a global analysis of local stream insect genus

richness, Vinson and Hawkins (2003) found that linear models based on large-scale climatic 139 and energy variables could explain some variation in local richness. They also found no clear 140 latitudinal gradients in the local genus richness of mayflies, stoneflies and caddisflies, but 141 instead that richness showed multiple peaks and often peaked at mid-latitudes. These two 142 studies, although based on large compilations of datasets, are likely to be limited by different 143 sampling methods as well as distinct environmental and spatial extents in different regions. 144 145 This might hinder firm conclusions on whether stream insect assemblages in tropical, temperate and boreal regions show differences in richness and abundance patterns. For 146 147 instance, differences in spatial extent may complicate among-region comparisons because larger study areas exhibit larger environmental heterogeneity, thereby affecting diversity 148 estimates (Chase & Knight, 2013). This issue has been considered in previous local-scale 149 studies by using a long-term sampling approach (Vinson & Hawkins, 2003) and in some 150 regional-scale studies by factoring out the effects of spatial extent (Pearson & Boyero, 2009). 151

152 To overcome some of the problems that have affected comparative studies based on datasets produced by different methods (e.g. Nakagawa & Parker, 2015) and complement 153 previous findings on global gradients (e.g. Pearson & Boyero, 2009), we conducted identical 154 155 surveys of wadeable streams in a boreal (western Finland) and a subtropical (south-eastern Brazil) region. We believe that broad-scale studies (e.g. based on gridded data) and 156 157 comparative studies (i.e. based on standardised sampling protocols) are complementary and offer the possibility to test for patterns using different lines of evidence. Thus, by focusing on 158 two regions, as opposed to previous studies that addressed global gradients, we could take 159 detailed and standardised insect samples from 100 streams in each region and measured the 160 same environmental variables in both regions. We also standardised the extent of the study 161

area by resampling sites from similar-sized areas in both regions. Our specific aims were 162 threefold. We first (i) examined whether genus accumulation curves differed between the 163 boreal and subtropical regions. We expected that there would be faster accumulation of 164 genera in Brazil than in Finland because alpha and beta diversities are typically higher in the 165 tropics (Soininen et al., 2007). We also (ii) studied whether rank-abundance distributions 166 differed between Brazil and Finland, with the expectation that there would be many more 167 168 relatively rare genera in the subtropical than in the boreal region, which is more dominated by a few abundant genera (Brown, 2014). Finally, in addition to these aims, we (iii) modelled 169 170 variation in local genus richness and local assemblage abundance of stream insects in relation to region identity, while controlling for the effects of key local environmental variables. If 171 evolutionary diversification (Willig et al., 2003; Brown, 2014) is contributing to the 172 latitudinal richness gradient, insect genus richness and assemblage abundance should be 173 higher in subtropical than in boreal streams after controlling for the influence of local 174 environmental variables and spatial extent. 175

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177 MATERIALS AND METHODS

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179 Study areas

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Western Finland. Study sites were located between latitudes 60°41'N and 65°12'N, with
spatial extents of ca. 500 km and 300 km in north-south and in east-west directions,

respectively (Supporting Information, Fig. S1). The catchments were covered by different 183 land uses, from ones dominated by agriculture to those covered almost entirely by boreal 184 forests (dominated by Pinus sylvestris and Picea abies). The 100 perennial streams were 185 selected from 20 major river basins, i.e., five separate streams draining into each of 20 boreal 186 rivers were surveyed. Western Finland has the four seasons characteristic of the boreal 187 region: (1) winters lasting from November to March, (2) spring periods in April and May, (3) 188 189 summers between June and August, and (4) autumn periods generally in September and October. Given the large latitudinal extent, temperature and rainfall vary substantially in the 190 191 study area. The stream sites were sampled in September 2014. These streams did not experience heavy floods or droughts just before sampling and ranged from pristine forest 192 streams to agricultural streams with little pollution. 193

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South-eastern Brazil. Study sites were located in the State of São Paulo, between latitudes 195 24°38'S and 23°82'S, with spatial extents of 70 km in north-south and 120 km in east-west 196 directions, respectively (Supporting Information, Fig. S1). The 100 perennial streams, with 197 five streams distributed within 20 different catchments as described above, are located 198 199 between three major Atlantic Forest protected areas (Carlos Botelho, Intervales and Alto Ribeira State Parks). Land cover varied among catchments, with some dominated by 200 agriculture (mainly pastures, and *Eucalyptus* and *Pinus* plantations) to some entirely covered 201 by Atlantic Forest. This region is characterised by two seasons: a drier season from April to 202 August (average rainfall from 45 to 80 mm per month; average temperature from 16 to 20°C) 203 204 and a wetter season from September to March (average rainfall from 105 mm to 180 mm per month; average temperature from 20 to 23°C). The sites were sampled from September to 205

November in 2015. Streams did not suffer from recent floods or droughts and ranged from
pristine forest streams to moderately polluted streams in agricultural areas.

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209 Field surveys

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211 We measured 15 physical and chemical variables that have been found to be important for stream insect distributions in boreal (Malmqvist & Mäki, 1994; Grönroos & Heino, 2012) 212 and tropical (Siqueira et al., 2012; Al-Shami et al., 2013) regions. For physical variables, we 213 measured current velocity (m/s) and depth (cm) at 30 (in Finland) and nine (in Brazil) 214 random spots in a riffle site. Particle size classes (%) were visually estimated in 0.25 m^2 215 squares at 10 (in Finland) and three (in Brazil) random locations in a riffle site. We used a 216 modified Wenthworth's scale of particle size classes: sand (0.25-2 mm), gravel (2-16 mm), 217 pebble (16-64 mm), cobble (64-256 mm) and boulder (256-1024 mm). Based on the mean 218 estimates for each site, we calculated the Shannon diversity of substratum particle sizes for 219 each stream site. We also measured mean stream width of the sampling site based on 10 (in 220 Finland) and three (in Brazil) cross-channel measurements and visually estimated shading 221 222 (i.e. canopy cover) by riparian vegetation at each sampling site. We used satellite images to estimate native forest cover within a 400-m buffer along tracts of the sampled streams. For 223 224 chemical variables, we measured pH and conductivity at each site in the field using YSI device model 556 MPS (YSI Inc., Ohio, USA) in Finland and Horiba device U-50 series in 225 Brazil. Water samples were analysed for total nitrogen (TN) and total phosphorus (TP) 226 following national standards for Finland (Finnish Board of Waters and the Environment 227

228 1981) and Brazil (Golterman *et al.*, 1978; Mackereth *et al.*, 1978).

229 We took a 2-minute kick-net sample (net mesh size: 0.5 mm) at each of the stream sites surveyed in Finland and Brazil. The sample for each site consisted of four 30-seconds 230 sample units that were obtained in the main microhabitats (i.e. considering variations in 231 232 current velocity, depth, particle size and macrophyte cover) at a riffle site of c. 25 to 50 m². 233 The four sample units were pooled, preserved in alcohol in the field and taken to the laboratory for further processing and identification. All insects were separated from debris, 234 and taxa of our interest (Ephemeroptera, Odonata, Plecoptera, Megaloptera, Trichoptera and 235 Coleoptera) were identified to genus level. 236

Species-level identification is mostly not feasible for aquatic insects in Brazil as many 237 species remain undescribed, particularly their aquatic immature stages, and new species are 238 still being discovered in the region (Cruz et al., 2013). A previous evaluation on the use of 239 240 coarse family-level identification in the same Brazilian region as studied here indicated that it produces reliable assessment of stream fauna (Melo, 2005). Species-level identification is 241 more feasible in Finland, but regional, basin and local aquatic insect faunas of wadeable 242 streams are still inadequately known. However, genus richness portrays adequately patterns 243 in species richness in Finnish streams (Heino & Soininen, 2007). Accordingly, we opted to 244 245 use genus identification to enable direct comparisons between Finland and Brazil.

246

247 Standardization of sampling effort and spatial extent

We developed different procedures to control for variation in sampling effort and differences
in spatial extent, before comparing the two regions. We redirect the reader to the
supplementary material (see Supporting Information and Fig. S2) for a detailed account of
these procedures. These procedures were needed to ensure that between-region differences in
richness were not caused by anomalous sites with low densities and by differences in the
number of streams within drainage basins and spatial extents.

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256 Graphical data analysis at the regional scale

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To describe regional-scale patterns in the stream insect faunas of Finland and Brazil, we used 258 two graphical approaches using mean values from reduced/standardised datasets. First, we 259 drew genus accumulation curves for each region to see if genus richness accumulated 260 similarly with increasing number of stream sites (Coleman et al., 1982). This approach 261 262 followed the method "exact" devised by Ugland et al. (2003). Second, we used genus rank-263 abundance curves at regional level to show which genera were most abundant in Finland and Brazil. In these plots, abundance (as proportions of total regional abundance) is on the y-axis, 264 and genera are ranked from the most abundant to the least abundant on the x-axis (Whittaker, 265 1965). Genus accumulation curves were drawn using the function 'specaccum' in the R 266 package 'vegan' (Oksanen et al., 2013). 267

268

269 Modelling genus richness and assemblage abundance across the regions and along

270 environmental gradients

272	The standardisation of spatial extent (see Supporting Information) and sample sizes					
273	(rarefaction) allowed an improved comparison of genus richness between the regions.					
274	However, the streams also differed on several environmental features between the regions.					
275	We thus estimated differences in abundance, observed genus richness and rarefied genus					
276	richness between the two regions after taking into account the environmental variables					
277	described earlier and known to affect stream insect assemblages.					
278	Physical variables were similar between regions (Table 1). In contrast, chemical					
279	variables differed between the regions, with conductivity, total phosphorous and total					
280	nitrogen being much higher in Finland than in Brazil (Table 1). Accordingly, we standardised					
281	chemical variables by their maximum within each region before analyses using the function					
282	'decostand' in the R package 'vegan' (Oksanen et al., 2013). The standardisation reduced the					
283	correlation with the 'region identity' variable and allowed the evaluation of effects of those					
284	variables within the range of variation within each region.					
285	We removed the variable boulder, as the five substrate classes sum to 1, and they					
286	would thus be correlated to each other. We then removed collinear environmental variables					
287	(i.e. cobble and TN) by sequentially removing those with variance inflation factor (VIF)					
288	higher than 2 (Zuur et al., 2010). We fitted linear models with 14 explanatory variables: the					
289	13 environmental variables described earlier and 'region identity', our categorical variable of					
290	interest. We performed model selection using AIC criterion, although we were most					
291	interested in the effect of 'region identity' after partialling out the effect of environmental					

292	variables. The model selection employed aimed to provide a simpler model. We ran separate
293	multiple regressions on rarefied richness, observed richness and assemblage abundance using
294	14 predictor variables. Finally, we ran a commonality analysis in the context of multiple
295	regression (Seibold & McPhee, 1979; Ray-Mukherjee et al., 2014) to infer the unique,
296	common and total contributions of each predictor variable to response variables. VIF
297	analyses were done using the function 'vif' in the 'car' R package (Fox & Weisberg, 2011).
298	Modelling analyses were run using the functions 'lm' and 'step' in the R package 'stats' (R
299	Core Team, 2017) and 'commonalityCoefficients' in the R package 'yhat' (Nimon et al.,
300	2015).
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202	
302	NESULIS
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304	Patterns of regional-scale richness and abundance
305	
306	Stream insect abundance and richness showed interesting differences between Finland and
307	Brazil. First, only four genera were shared between Finland and Brazil, these being the
308	mayfly genus Caenis and the caddisfly genera Hydroptila, Oecetis and Oxyethira. Second,
309	total regional abundance was much higher in Finland (total number of insect individuals in
310	100 streams: 86,048) than in Brazil (16,113), yet regional genus richness ($n = 100$ streams)
311	was slightly higher in Brazil (83) than in Finland (77).
312	The 97 paired standardised sets of streams included (see Supporting Information and

Fig. S2), on average, 24.5 (min = 17, max = 29) streams in each region, and the average 313 spatial distance among them was 56.0 km (min = 44.1, max = 65.7) in Brazil and 57.6 km314 (min = 45.3, max = 68.8) in Finland. These results indicate the effectiveness of our 315 standardization protocol. For all standardised pairs, total abundance in the set of streams in 316 Finland was higher than in Brazil (average paired difference = 16,752 individuals). In 317 contrast, observed genus richness was always higher in Brazil and included 40% more genera 318 319 than in their paired counterparts in Finland (average paired difference = 18.66 genera; Fig. 1A). This difference increased to 76% after taking differences in the number of sampled 320 321 individuals into account (within-pair rarefaction; average paired difference = 28.54 genera; Fig. 1B). 322

323 Sample-based accumulation curves were similar between Finland and Brazil, although
324 genus richness was much higher in Brazil (Fig. S3A, B). In contrast, individual-based
325 accumulation curves indicated that many more genera would be detected in Brazil with
326 additional sampling (Fig. S3C, D).

Rank-abundance curves for Finland and Brazil differed mostly in two aspects (Fig.
S4). The two most abundant genera in Finland tended to include much more of the total
number of individuals (on average, 32.03 and 19.78%) than the two most abundant in Brazil
(14.45 and 12.16%). In contrast, the proportions of rare genera were similar between Brazil
(74.5 and 81.6% of the genera with less than 1 and 2% of the total abundance) and Finland
(74.5 and 81.8%).

333

334 Patterns of basin-scale richness and abundance

Samples from the 17 basins in Brazil, each including four or five streams, included a total of
15,471 individuals. The 19 basins in Finland included 85,050 individuals. Observed genus
richness was 31% higher in Brazil than in Finland (37.23 and 28.42, respectively; Fig. 2A).
The basin with the lowest abundance was found in Brazil, with 295 individuals. Rarefied
genus richness for 295 individuals was 61.3% higher in Brazil than in Finland (30.17 and
18.51, respectively; Fig. 2B).
Patterns of local-scale richness and abundance
The reduced set of streams in Brazil, after removing 12 streams with low insect abundance,
included on average 181.5 individuals and 17.84 genera per stream riffle site. The reduced set
of streams in Finland included many more individuals (mean = 886.57) and fewer genera
(14.01) per stream. Observed mean genus richness per stream was 27% higher in Brazil than
in Finland (Fig. 3A). Rarefied genus richness per stream was 64.3% higher in Brazil than in
Finland (11.52 and 7.41, respectively; Fig. 3B).
Differences in genus richness and assemblage abundance between the regions

49.6% of the variation in rarefied genus richness (Table 2). Region had the strongest effect on 355 rarefied genus richness, with Brazil possessing more genera per stream after controlling for 356 the effects of local environmental variables. Rarefied genus richness was also higher in wider 357 streams and those with a high proportion of native forest vegetation cover. Proportion of sand 358 in the stream bottom was negatively related to rarefied richness. A similar reduced model for 359 observed richness explained 37.6% of variation, and the most important predictor variables 360 361 were region and five local variables. Finally, assemblage abundance was best explained by region and seven local variables, and the model accounted for 61% of variation in assemblage 362 abundance (Table 2). Full models can be found in Supporting Information (Table S1). 363

Commonality analysis strengthened the findings of basic linear models by showing 364 that 'region' typically had the highest unique (14.1% to 18.8%) and total (8.5% to 44.2%) 365 effects on rarefied richness, observed richness and assemblage abundance (Table 2). The only 366 exception was for observed richness as stream width had the highest total effect (11.4%). For 367 rarefied richness, shading and forest cover were also relatively important. For observed 368 richness, stream width and forest cover were important in addition to the region effect. 369 Finally, for assemblage abundance, the second and third most important predictors were 370 371 shading and velocity, respectively.

372

373 **DISCUSSION**

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We found that, for the entire datasets, the regional difference in genus richness was

surprisingly small between Brazil (83 genera) and Finland (77 genera) considering that 376 regions closer to the equator usually harbour more diversity than those closer to the poles 377 (Rosenzweig, 1995; Willig et al., 2003). While differences between tropical and boreal 378 regions are expected for various groups of organisms, many freshwater organisms do not 379 obey such general 'laws' (Vinson & Hawkins, 2003; Pearson & Boyero, 2009; Heino, 2011; 380 Soininen, 2012). Rather, aquatic insects, such as mayflies, stoneflies and caddisflies, do not 381 382 often follow typical latitudinal trends in regional diversity, whereas some other aquatic insects, such as dragonflies and beetles, do so (Brown, 1981; Pearson & Boyero, 2009). In 383 384 our study, mayflies, dragonflies and beetles were regionally more diverse in Brazil than in Finland, whereas the opposite was true for stoneflies and caddisflies (Supporting Information, 385 Fig. S5). This finding suggests that some insect orders may be more diverse closer to the 386 poles than to the equator. We emphasise, however, that the difference in genus richness 387 between Brazil and Finland was much larger after accounting for spatial extent. Indeed, in the 388 reduced standardised datasets, regional genus richness was clearly higher in Brazil than in 389 Finland. This finding underscores the importance of standardising spatial extent when 390 comparing regional richness estimates. Such standardisation is important because larger 391 spatial extent typically incorporates more environmental heterogeneity, resulting in higher 392 richness (Chase & Knight, 2013) and leading to biased between-region comparisons in the 393 case of different spatial extents. 394

In general, richness tends to be correlated with abundance especially if sampling effort varies between regions (Gotelli & Collwell, 2001; Evans *et al.*, 2005). However, this was not the case in our study with identical field sampling designs. Hence, it seems that the very high regional abundance does not lead to high genus richness in Finland, and tropical

faunas remain more diverse even when having much lower regional abundances than boreal 399 faunas. Previous studies have detected increasing abundance of stream detritivorous 400 invertebrates with increasing altitude (Yule et al., 2009) and latitude (Boyero et al., 2011), 401 suggesting that stream insect abundance may be lower in higher temperatures. Also, other 402 differences between regions, such as local stream productivity and other environmental 403 features, affect stream insect abundance (Jacobsen et al., 1997; Vinson & Hawkins, 2003). In 404 405 our study, nutrient concentrations were much higher in Finnish than Brazilian streams (Table 1), which might have contributed to higher insect abundance in Finland. This difference 406 407 might stem from the generally nutrient-poor soils of tropical landscapes (Reich & Oleksyn, 2004), as catchment features typically determine stream water chemistry (Soininen et al., 408 2015). However, in the linear models, the effect of nutrients was masked by collinearity with 409 the 'region' variable, the latter of which was the variable of our main interest and forced into 410 all models. 411

412 Genus accumulation curves differed slightly between the two regions. While the curves in both regions nearly reached an asymptote, the increase of genera was faster in 413 Brazil than in Finland (Fig. S3). Such steeper accumulation curves in the subtropical than in 414 415 the boreal region might be related to the slightly higher local genus richness in Brazil than in Finland. Working with data from three high-latitude regions (66°N to 70°N), Heino et al. 416 417 (2015b) found only minor differences in species accumulation curves among the regions, suggesting that environmental differences or species pool characteristics did not contribute to 418 differences in species accumulation. The situation should be different between geographically 419 distant regions, harbouring almost completely different stream biotas, such as those in Brazil 420 421 and Finland.

Despite differences in regional genus richness and total abundance between Brazil 422 and Finland, the rank-abundance curves were rather similar between the two regions (Fig. 423 424 S4). The curves were characterised by few common genera and several uncommon to very uncommon genera, a pattern that has previously been found for tropical streams (Siqueira et 425 al., 2012). However, the two most common genera in Finland were much more abundant than 426 their counterparts in Brazil. In both study regions, the most common genus belonged to the 427 428 beetle family Elmidae (Elmis in Finland and Heterelmis in Brazil). These riffle beetles are typical inhabitants of streams over most of the world, being often common in fast-flowing 429 430 sites (Elliott, 2008). The second and third most common genera in Finland belonged to mayflies (Baetis) and stoneflies (Nemoura), whereas those in Brazil belonged to caddisflies 431 (Smicridea) and mayflies (Farrodes). These insect orders typically dominate stream insect 432 faunas in many parts of the world (Lancaster & Downes, 2013). In general, these results 433 suggest an interesting avenue for further investigations: the existence of a "latitudinal 434 gradient" in niche packing (see also Willig et al., 2003). 435

We also found differences in local genus richness and local assemblage abundance 436 between Finland and Brazil. However, the richness difference between Finland and Brazil 437 was not as strong at the local scale as at the regional scale (compare Fig. 1 and 2 with Fig. 3), 438 which was evidenced by both the entire dataset and the reduced standardised datasets. While 439 440 local genus richness was higher in subtropical than in boreal streams, local assemblage abundance showed the opposite pattern. However, both richness and abundance also showed 441 considerable variation within each region (Table 1, Fig. 3). These findings suggest that local 442 richness and local abundance in streams are determined by both regional and local factors, 443 and it seems that genus richness benefits from tropical conditions, whereas assemblage 444

abundance benefits from boreal ecological features. Higher genus richness in the tropics 445 might result from ice age history and temperature-related evolutionary rates (Mittelbach et 446 447 al., 2007), which should be less important and more important, respectively, in the tropical than in boreal regions (Brown & Lomolino, 1998). For example, previous evidence indicates 448 that Neotropical areas can act both as "museums" (i.e. with old persistent lineages) and as 449 "cradles" (i.e. with new species continuously originating) of insect diversity (Moreau & Bell, 450 451 2013). In addition, regional factors set the upper limits to local genus richness and assemblage abundance, which are further affected by more localised variation in stream 452 453 environmental variables (Poff, 1997; Vinson & Hawkins, 2003). These environmental variables may vary within regions (e.g. pH) or differ between regions (e.g. TN), as in our 454 455 case.

An interesting finding was that the between-region richness difference decreased from 456 regional through basin to local scales (40, 31 and 27% for observed genus richness and 76, 63 457 and 55% for rarefied genus richness; Figs 1, 2 and 3). This finding suggests that evolutionary 458 history and climatic influences are strong in setting up differences in regional genus richness 459 (standardized spatial extents), while local genus richness differences are slightly diminished 460 due to the possible operation of catchment variables (acting on catchment-scale) and local 461 environmental variables (acting on stream-scale) with decreasing spatial grain of a study. 462 463 While previous studies that were based on intensive site-based inventories (Vinson & Hawkins, 2003) or more extensive regional inventories (Pearson & Boyero, 2009) did not 464 adopt standardized methods, they were able to provide broad generalizations on global 465 richness gradients. However, our study adds to previous findings by providing more precise 466 evidence from explicit consideration of variation of richness and abundance across spatial 467

scales. Hence, our findings emphasize the need to consider both spatial extent and grain of
the study when making comparisons of between-region differences in biodiversity patterns
(Vinson & Hawkins, 1998; Heino, 2011).

471

472 Conclusions

473

We found that stream insect genus richness and assemblage abundance differed between our 474 subtropical and boreal regions. Our main finding was that genus richness was higher in Brazil 475 than in Finland (and more so at regional and basin than local scales), yet assemblage 476 abundance was much higher at all spatial scales in Finland than in Brazil. These patterns hold 477 in both entire datasets and in reduced datasets based on resampling of the stream sites from 478 similar-sized areas. Differences in nutrient concentrations and temperature of stream waters 479 might explain the higher abundance of insects in Finland than in Brazil, whereas possibly 480 factors related to evolutionary diversification might explain differences in genus richness 481 between these two regions. Our further studies should shed more light on these issues by 482 focusing on the functional structure and phylogenetic diversity of stream insect faunas in 483 484 these geographically distinct regions.

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666	
667	DATA ACCESSIBILITY
668	The data used in this work are stored in PANGAEA (<u>https://www.pangaea.de/)</u> . The datasets
669	can also be found in Supporting Information of this paper.
670	
671	BIOSKETCH
672	Our Finnish-Brazilian team is interested in biodiversity patterns in general and those from
673	stream ecosystems in particular. This research originated during a joint research project
674	funded by the Academy of Finland and FAPESP. Author contributions: JH, JS, TS, ASM,
675	VLL and LMB conceived the ideas; JJM, DKP, KTT, VP and TSFS collected the data; DKP,
676	VSS and KTT identified the insects; ASM analysed the data; and JH and TS led the writing.
677	All authors contributed to the ideas and approved the final version of the manuscript.
678	

679 SUPPORTING INFORMATION

680 Additional Supporting Information is available in the online version of this paper.

681 **Tables and Figures**

682

Table 1. Descriptive statistics of local genus richness, local rarefied genus richness and local

assemblage abundance, of stream insects, as well as of selected physical and chemical

variables measured in streams in Finland (N = 97) and Brazil (N = 88). These values are

based on the reduced datasets after removal of anomalous sites. SD = standard deviation, IQR

687 = interquartile range, CV = coefficient of variation.

688

Finland			Brazil					
Variable	Mean	SD	IQR	CV	Mean	SD	IQR	CV
Observed richness	14.01	5.07	8.00	0.36	17.84	7.46	11.25	0.42
Rarefied richness	7.41	2.39	2.80	0.32	11.53	3.60	4.46	0.31
Abundance	886.57	700.73	852.00	0.79	181.50	111.38	190.00	0.61
рН	7.13	0.57	0.86	0.08	6.60	0.51	0.70	0.08
Conductivity (μ S/cm)	88.17	84.19	86.90	0.95	55.79	58.68	29.00	1.05
TN (µg/L)	852.84	507.46	537.50	0.60	119.60	34.59	41.67	0.29
TP (µg/L)	72.14	58.52	65.67	0.81	15.33	12.47	9.30	0.81
Stream width (m)	3.84	2.53	2.95	0.66	2.60	1.75	0.84	0.67
Shading (%)	37.93	23.03	37.75	0.61	72.70	20.39	25.00	0.28
Mean velocity (m/s)	0.25	0.11	0.14	0.43	0.21	0.08	0.12	0.38
Mean depth (m)	0.18	0.07	0.07	0.37	0.13	0.07	0.07	0.54
Substratum diversity	1.06	0.33	0.39	0.31	1.29	0.20	0.25	0.16

689

691 Table 2. Reduced multiple regression models showing the effects of 'region' and local

environmental variables on rarefied richness (a; $F_{13,177} = 24.86$, p < 0.001), observed richness

- 693 (b; $F_{13,178} = 17.88$, p < 0.001) and assemblage abundance (c; $F_{13,176} = 34.41$, p < 0.001) of
- 694 stream insects in Finland and Brazil. Also shown are unique, common and total effects of
- 695 each predictor variable from commonality analysis. See Supporting Information Table S1 for
- 696 full models.

a) Rarefied richness									
	Estimate	SE	t	р	Unique	Common	Total	\mathbb{R}^2	adj. R ²
(Intercept)	5.504	0.792	6.946	< 0.001					
region	4.506	0.542	8.309	< 0.001	0.188	0.130	0.318		
width	0.439	0.095	4.639	< 0.001	0.072	-0.050	0.022		
forest cover	1.654	0.679	2.438	0.016	0.020	0.085	0.105		
sand	-0.027	0.013	-2.156	0.032	0.009	0.001	0.010		
total P	-2.388	1.204	-1.983	0.049	0.013	0.016	0.029		
shading	0.016	0.009	1.716	0.088	0.008	0.121	0.129		
								0.496	0.476

b) Observed richness

c) Abundance

	Estimate	SE	t	р	Unique	Common	Total	
(Intercept)	-0.609	5.394	-0.113	0.910				
region	5.428	0.852	6.369	< 0.001	0.142	-0.057	0.085	
width	1.046	0.187	5.607	< 0.001	0.110	0.004	0.114	
forest cover	4.368	1.382	3.160	0.002	0.035	0.057	0.092	
sand	-0.069	0.025	-2.723	0.007	0.026	0.026	0.052	
pH	11.654	6.060	1.923	0.056	0.013	-0.012	0.001	
total P	-4.493	2.395	-1.876	0.062	0.012	0.042	0.054	

0.376 0.355

,									
	Estimate	SE	t	р	Unique	Common	Total		
(Intercept)	3.328	0.770	4.319	< 0.001					
region	-1.186	0.149	-7.971	< 0.001	0.141	0.301	0.442		
shading	-0.009	0.002	-3.547	< 0.001	0.028	0.265	0.293		
pH	2.820	0.801	3.518	< 0.001	0.027	0.018	0.045		
velocity	2.097	0.599	3.498	< 0.001	0.027	0.116	0.144		
sand	-0.011	0.003	-3.469	< 0.001	0.027	0.103	0.129		
pebble	0.011	0.003	3.100	0.002	0.021	-0.003	0.018		
particle diversity	0.286	0.194	1.472	0.143	0.005	0.045	0.049		
forest cover	0.274	0.189	1.455	0.147	0.005	-0.002	0.002		
								0.610	0.592



Fig. 1. Differences in regional stream insect genus richness between Brazil and Finland based
on similar-sized areas after random resampling in both regions. Note that a large number of
resampling analyses show the same pattern, i.e., regional genus richness is much higher in
Brazil than in Finland.



Fig. 2. Boxplots denoting differences in basin-scale stream insect genus richness between the
two regions, Brazil (N = 17 basins) and Finland (N = 19 basins). Shown are observed (A) and
rarefied (B) genus richness values.



Fig. 3. Boxplots denoting differences in local stream-scale genus richness between the two regions, Brazil (N = 88 streams) and Finland (N = 97 streams). Shown are observed (A) and rarefied (B) genus richness values.