

1 Comparability of macroinvertebrate biomonitoring indices of river  
2 health derived from semi-quantitative and quantitative  
3 methodologies.

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16 **Abstract:**

17 Aquatic macroinvertebrates have been the basis for one of the primary indicators and a cornerstone  
18 of lotic biomonitoring for over 40 years. Despite the widespread use of lotic invertebrates in  
19 statutory biomonitoring networks, scientific research and citizen science projects, the sampling  
20 methodologies employed frequently vary between studies. Routine statutory biomonitoring has  
21 historically relied on semi-quantitative sampling methods (timed kick sampling), while much  
22 academic research has favoured fully quantitative methods (e.g. Surber sampling). There is an  
23 untested assumption that data derived using quantitative and semi-quantitative samples are not  
24 comparable for biomonitoring purposes. As a result, data derived from the same site, but using  
25 different sampling techniques, have typically not been analysed together or directly compared. Here,  
26 we test this assumption by comparing a range of biomonitoring metrics derived from data collected  
27 using timed semi-quantitative kick samples and quantitative Surber samples from the same sites  
28 simultaneously. In total, 39 pairs of samples from 7 rivers in the UK were compared for two seasons  
29 (spring and autumn). We found a strong positive correlation ( $r_s = +0.84$ ) between estimates of taxa  
30 richness based on ten Surber sub-samples and a single kick sample. The majority of biomonitoring  
31 metrics were comparable between techniques, although only fully quantitative sampling allows the  
32 density of the community (individual  $m^{-2}$ ) to be determined. However, this advantage needs to be  
33 balanced alongside the greater total sampling time and effort associated with the fully quantitative  
34 methodology used here. Kick samples did not provide a good estimate of relative abundance of a  
35 number of species / taxa and, therefore, the quantitative method has the potential to provide  
36 important additional information which may support the interpretation of the biological metrics.

37 **Keywords:**

38 Macroinvertebrate; Species Richness; Biological Monitoring; Biotic Index; River

39

40

41 **1. Introduction:**

42 Rivers and the ecological communities they support comprise some of the most biodiverse habitats  
43 on the globe but are also some of the most degraded as a result of anthropogenic activity (Dudgeon  
44 et al. 2006; Carpenter et al. 2011). River habitats and their ecosystems are threatened by ongoing  
45 human development (Vörösmarty *et al.* 2010), including the modification of channel morphology,  
46 dredging, changes to catchment land-use, pollution from diffuse and point sources, invasion by alien  
47 species, and alterations of the flow regime from abstraction, damming and flood risk management  
48 (Carpenter *et al.* 2011). The historic degradation of rivers has prompted the development of a range  
49 of biological monitoring tools to survey and quantify anthropogenic stressors over the past 40 years  
50 (e.g., Hering et al., 2004) and underpin calls to restore and improve the ecological health of lotic  
51 ecosystems (e.g., Geist, 2011).

52 In order to quantify trends in the health of riverine environments, the response of an organism or  
53 community is often characterised as a metric based on their known tolerances to ‘stressors’.  
54 Biological monitoring, or biomonitoring, can be used to assess the effect of a known change to the  
55 state of a system by comparing the ecological community before and after the change or to  
56 routinely check compliance to nationally / internationally recognised standards, such as the legal  
57 requirement for all waterbodies in the European Union to achieve ‘Good Ecological Status’ under the  
58 Water Framework Directive. The taxonomic resolution of such indices varies from family-level  
59 metrics that give broad indications of water quality (e.g., Walley and Hawkes 1997) to  
60 species/genus-level metrics that can provide information about specific stressors (Hubler et al.,  
61 2016); although some can be used at different taxonomic resolutions (Monk et al., 2012). Other  
62 metrics use higher resolutions; for example, the phenology of species or groups of species can be  
63 used to assess the impacts of climate change (Everall et al. 2015; Thackeray et al. 2016).

64 Aquatic macroinvertebrates are a fundamental component of freshwater ecosystems. Hence,  
65 maintaining macroinvertebrate communities, biodiversity and individual species populations  
66 contributes to the overall ecological integrity of the system (Spänhoff and Arle, 2007). Particular  
67 invertebrates (species, genus or families) have tolerance limits to specific environmental conditions,  
68 such as levels of salinity, pH, organic pollution, suspended sediment concentration, fine sediment  
69 deposition and flow velocity (e.g. Hellowell, 1986). Macroinvertebrate biomonitoring tools and  
70 assessment systems are widely used to assess water quality globally (e.g. North America – Barbour  
71 et al. 1999; Africa – Cummins *et al.* 2004; Asia – Morse *et al.* 2007; South America – Dickens &  
72 Graham, 2002), although there have been recent calls for methods of assessing ecological response  
73 to environmental changes and pressures to be more strongly rooted in ecological and biological

74 theory (e.g. Friberg *et al.* 2011; Johnson and Rice, 2014). In Europe, macroinvertebrate  
75 biomonitoring forms an important part of compliance monitoring within the European Union Water  
76 Framework Directive (WFD). This Directive requires Member States to ensure that all freshwater  
77 bodies are of 'Good Ecological Status (GES) or Good Ecological Potential (GEP) for Heavily Modified  
78 Waterbodies (HMWB) and Artificial Waterbodies (AWB) by 2027 (EU Directive 2000/60/EC).

79 Biomonitoring techniques can be quantitative, semi-quantitative or qualitative, depending on the  
80 technique used. The most common method for sampling invertebrates in rivers is the semi-  
81 quantitative kick sample method, where invertebrates are sampled over a specified time period  
82 (typically three-minutes) supplemented by hand searches of larger substrate clasts; although the  
83 total area or proportion of the community sampled is typically unknown (Murray-Bligh, 1999; ISO  
84 10870, 2012). Most macroinvertebrate biomonitoring indices have been developed to allow  
85 macroinvertebrate community composition to be analysed on a semi-quantitative basis where  
86 sampling effort (time) is standardised (Clements and Newman, 2002). Fully quantitative sampling is  
87 necessary for other forms of analysis that require information regarding the total abundance,  
88 density or diversity of organisms/communities within a specified area. This can be achieved with a  
89 Surber sampler (or other similar devices such as a cylinder sampler, or Hess sampler), where  
90 invertebrates are collected within a specified sampling area.

91 Whilst there is widespread agreement that the macroinvertebrate community provides a valuable  
92 tool to characterise the ecological health of rivers, there is less consensus about the most  
93 appropriate sampling methodologies to employ. Surprisingly, the degree to which biological metrics  
94 derived from semi-quantitative and quantitative samples differ has not been widely assessed in a  
95 systematic way. The largely untested assumption that biomonitoring scores are not comparable  
96 between these methods prevents both historic (e.g. Percival and Whitehead, 1929; Percival and  
97 Whitehead, 1930; Prigg, 2002) and contemporary fully quantitative data from being combined and  
98 used to characterise river health. Hence, the aim of this paper is to compare a semi-quantitative kick  
99 sampling methodology with a quantitative Surber sampling methodology at given sites by cross-  
100 matching: 1) derived biomonitoring scores/indices; 2) inferred water- and habitat-quality; and 3) the  
101 abundance and diversity of the taxa collected by each method.

## 102 **2. Methodology:**

### 103 *2.1. Sampling techniques*

104 Kick sampling is a semi-quantitative method of surveying the invertebrate community, which is  
105 widely used internationally because it is cost effective and results are relatively consistent between  
106 operators (e.g. Carter and Resh, 2001; Metzeling *et al.*, 2003). In this study, a 1 mm<sup>2</sup> mesh net with  
107 an opening 0.25 m wide and 0.22 m deep was held downstream of the operator who kicked the river  
108 bed and swept the net through, for example, submerged macrophytes. This action disturbs sediment  
109 and dislodges benthic invertebrates which are then carried by the river flow into the net. The  
110 duration of kick sampling here followed the Environment Agency of England (EA) best-practice  
111 standard, which requires three-minutes of kick sampling and one-minute hand search of larger  
112 substrates for macro-invertebrates (HMSO, 1985, Murray-Bligh, 1999; Environment Agency, 2009).  
113 The operator moved systematically across and upstream through the river reach being sampled,  
114 ensuring that all main habitat types were sampled (e.g. emergent and submerged macrophyte  
115 stands, woody debris, tree roots, different flow depth/velocities and bed substrate compositions).  
116 The amount of time spent in each designated habitat unit was proportionate to the surface area that  
117 each occupied.

118 To obtain a quantitative comparison, replicate Surber samples were collected. A Surber sampler is a  
119 rectangular quadrat, 0.33 x 0.30 m (area 0.1 m<sup>2</sup>) that is placed on the river bed. The quadrat has a 1  
120 mm<sup>2</sup> mesh net attached, with a 0.29 x 0.34 m opening. The operator disturbs by hand all surface  
121 material within the quadrat area. Total sampling times can vary but in the current study continued  
122 until all of the 0.1 m<sup>2</sup> quadrat area was fully sampled (Surber, 1937; Macan, 1958). Sediment was  
123 disturbed to a maximum depth of 0.1 m. Disturbance dislodges invertebrates that then drift into the  
124 downstream net and, with the aid of side curtains, captures dislodged animals that might otherwise  
125 avoid capture in the net. Traditional Surber net sampling tended to be micro-habitat specific but for  
126 some river types Surber net sampling can form part of a methodology that proportionally samples  
127 different microhabitats (Prigg, 2002; Everall, 2010). In this study, 10 Surber samples, distributed such  
128 that all habitat types within the site were represented, were undertaken at each survey site. As with  
129 kick sampling, the habitats sampled reflected the proportion of the area covered by each habitat  
130 type at the site. For ease of analysis, the 10 individual samples were aggregated into 5 sub-samples  
131 for identification. The data from these 5 sub-sample units were, in turn, aggregated prior to the  
132 calculation of the biomonitoring indices/scores used for comparison between methods.

133 All samples were collected following the EA best practice guides (Environment Agency, 2009) by an  
134 experienced operator (Everall). Kick and Surber sampling was undertaken on the same day, at the  
135 same site, one immediately after the other. The second sample was taken a few metres upstream of  
136 the first but spatially alternating between kick and Surber net sample reaches at survey sites to

137 reduce any sampling sequence bias. Sample site reaches were selected for their similarity of  
138 instream habitat composition over the sampled reach and were divided into kick and Surber areas  
139 such that each had comparable proportions of the major habitat types.

140

## 141 *2.2. Sampling times and locations*

142 Sampling was undertaken on seven English rivers at a total of 20 sites (Figure 1). These locations  
143 were chosen to provide a range of habitat and climate types (Table 1). Geology and elevation were  
144 obtained from Ordnance Survey maps. Average discharge and average annual maximum discharge  
145 were derived from daily average and daily maximum discharge time-series from the nearest gauging  
146 station on each river available from the Centre for Ecology and Hydrology (CEH). The 1961-1990  
147 average annual precipitation for the area upstream of gauging stations is also included in Table 1.

148 Kick and Surber samples were undertaken in spring (March-May) and autumn (September-October)  
149 at all sites on all rivers except for the River Wye where a kick and Surber sample pair was only taken  
150 in spring (Table 1). In total, 39 paired kick and Surber samples were collected. All samples were  
151 identified by the same laboratory technician to species level where possible. Where not possible,  
152 invertebrates were identified to the highest possible taxonomic level.

153

## 154 *2.2. Biological scoring methods*

155 A set of ecological parameters and biological monitoring scores were calculated for each site (Table  
156 2). These represent commonly applied metrics in the UK that are used to identify water quality and  
157 more specific environmental stressors. The abundance and taxa richness of the whole community  
158 was quantified, as well as the diversity of Ephemeroptera, Plecoptera and Trichoptera (EPT) and  
159 *Gammarus*, which are important sentinels of environmental stressors in the UK. The abundance or  
160 proportion of EPT taxa is widely used and considered to be a good indicator of river health where  
161 salmonid fisheries are economically important (Stanford and Spacie, 1994; Clements and Newman,  
162 2002; Park *et al.* 2003). In addition, the Community Conservation Index (CCI; Chadd and Extence;  
163 2004) provides an indication of exceptionally rich or regionally unusual invertebrate populations by  
164 scoring invertebrates based on their rarity. The CCI can contribute to the overall description of the  
165 condition of an aquatic ecosystem, alongside indices designed to detect, for example, flow variation  
166 or changes in water quality.

167 The Biological Monitoring Working Party (BMWP) score, ranks individual macroinvertebrate families  
168 from 1 to 10 based on their sensitivity to water quality. The sum of the scores of all collected families  
169 is the BMWP score. Given that the BMWP score is affected by the number of families sampled but  
170 not by abundances within those families, the interpretation can be biased as a sample with many  
171 low scoring taxa might score the same as a sample with a few high scoring taxa. Therefore, the  
172 Average Score Per Taxon (ASPT) was introduced, where the BWMP is divided by the total number of  
173 scoring families, to provide an average measure (Armitage et al. 1983). The Whalley Hawkes Paisley  
174 Trigg (WHPT) biometric score (Paisley et al. 2013) was developed as an attempt to integrate the  
175 abundance weighting limitation of the BMWP scoring system. These are indicative of family-level  
176 aggregate and averaged biomonitoring scores and are part of the WFD assessment criteria in the UK,  
177 with similar systems implemented across Europe.

178 Stressor-specific indicators were also deployed. The saprobic index is used to assess organic  
179 pollution by assigning a value (the saprobic value,  $s$ ) to each invertebrate species or family which  
180 indicates their tolerance to organic pollution. Each invertebrate is also given an indicator value ( $G$ ),  
181 that represents the tolerance range of an invertebrate and acts as a weighting value, increasing the  
182 impact of very sensitive organisms on the overall saprobic score ( $S$ ). All saprobic values were  
183 obtained from Schmidt-Kloiber and Hering (2015b). The Saprobic indicator was used here because it  
184 is internationally recognised and used as a good indicator of organic enrichment and pollution and it  
185 was the forerunner for many contemporary systems.

186 Other stressor-specific indices used here include the Proportion of Sediment-sensitive Invertebrates  
187 (PSI), Lotic-invertebrate Index for Flow Evaluation (LIFE) and Total Reactive Phosphorous Index  
188 (TRPI). Both the PSI and LIFE score are regularly applied in the UK, particularly to sites that are in  
189 danger of not achieving WFD requirements. The PSI is used to assess the presence of fine sediment  
190 by calculating the percentage of sediment sensitive taxa present in a sample (Extence et al. 2010).  
191 Similarly, the LIFE score uses the proportion of flow sensitive invertebrates in a sample to describe  
192 the prevailing flow conditions at that site (Extence et al. 1999). Finally, the TRPI (Everall, 2010) uses  
193 the proportion of phosphorous tolerant and intolerant macroinvertebrates in a sample according to  
194 various river types and seasons (Paisley et al., 2003; Paisley *et. al.*, 2011). These scores are good  
195 examples of classification systems where the percentage or proportion of sensitive organisms are  
196 compared to the total community.

197

198 *2.3. Interpretation*

219 To enable interpretation of the indices a ranking system was used, where 1 indicates poor conditions  
200 (highly stressed/impacted conditions) and 5 indicates very good conditions (un-stressed and non-  
201 impacted conditions) (Table 3). The scoring system used herein is based on established knowledge  
202 where available (see references in Table 3). Biomonitoring scores were grouped into each of these  
203 classes and the differences in grouping between kick and Surber sample results were compared. The  
204 WHPT score is interpreted using the River Invertebrate Classification Tool (RICT), a software program  
205 that compares observed WHPT scores to expected scores (see Paisley et al. 2007; UKTAG, 2014) and  
206 therefore simple categorisation is not appropriate for this metric. Given that all the metrics are  
207 continuous and judgement is necessary for data that fall near the boundary of a class, the difference  
208 between kick and Surber samples as a percentage of the category size was also determined. This  
209 indicates the likelihood that a methodological difference would lead to the results falling into a  
210 different category.

211 Where a biomonitoring score has an inconsistent range within categories the average class size was  
212 calculated. For example, in the case of the BMWP, the middle condition (rank 3) has a range of 19  
213 whereas good (rank 4) has a range of 24. Therefore, it is possible for a difference between kick and  
214 Surber sampling to be greater than 100% of a class size but with both samples actually being in the  
215 same category. In addition, where both kick and Surber samples are in the highest category, it is  
216 possible to achieve scores that differ by more than 100% of a class boundary but within the same  
217 class because there is not a higher category.

218

#### 219 2.4. Statistics

220 The statistical significance of differences between sets of biological scores calculated with kick and  
221 Surber sampled data were tested. Shapiro-Wilk tests indicated data was normally distributed with  
222 the exception of the total abundance, abundance of *Gammarus*, species richness, CCI and EPT  
223 diversity. Paired-sample Student t-tests were performed in SPSS v.22 to assess normally distributed  
224 data. In the case of non-normally distributed data, a Wilcoxon Signed Rank test was performed  
225 instead. In addition, Pearson correlation and linear regression analysis was used to compare  
226 normally distributed kick and Surber sampled data. Where data was not normally distributed,  
227 Spearman correlation applied ( $r_s$ ). Initially, this was performed for each biological monitoring score,  
228 incorporating data collected at all sites and seasons ( $n = 39$ ). The data are spatially clustered and in  
229 some instances comprise multiple samples from the same site at different times of year. However,  
230 the regression analysis was not describing relationships between sites or times of year, but between



231 sampling strategies. Therefore, the clustering of data does not affect the robustness of the test. If  
232 the null hypothesis is met and both sampling methods provide identical information, the  $R^2$  should  
233 equal 1 and the data should fall on the 1:1 line (i.e.  $y = x$ ). Subsequently, linear regression analysis  
234 was also performed on spring and autumn data, separately, in a sub-set of cases.

235

236

### 237 **3. Results:**

#### 238 *3.1. Invertebrate abundance, diversity and community measures*

239 In total, 128,129 individual invertebrates were sampled across all sites and techniques (78 samples),  
240 representing 205 different taxa. At sites where Surber samples collected a high abundance of  
241 invertebrates, the equivalent kick sample also tended to collect a high relative abundance. Hence,  
242 the relationship between kick and Surber samples was significantly positively correlated ( $r_s = +0.64$ ;  $p$   
243  $< 0.001$ ). However, there was considerable scatter in the association (Figure 2).

244 In 90% of the samples, the total number of invertebrates collected was higher in the aggregated  
245 Surber samples than in the kick samples. Similarly, the total number of EPT collected was greater in  
246 Surber samples than equivalent kick samples in 85% of cases. The abundance of *Gammarus* sp. in  
247 samples was more similar between sampling methods, with only 62% of sites having greater  
248 abundance in Surber samples. Where Surber samples collected a greater abundance than the paired  
249 kick sample, they contained, on average, twice as many invertebrates as the equivalent kick sample.  
250 In contrast, the kick samples that were more abundant than Surber samples yielded, on average,  
251 only 1.2 times more individuals than the paired Surber samples (Table 4). The total invertebrate  
252 abundance and total EPT abundance for kick and Surber samples were significantly different ( $p <$   
253  $0.01$ , in both cases). The total number of *Gammarus* sp. sampled did not differ statistically between  
254 sampling techniques ( $p = 0.062$ ).

255 The total diversity of invertebrates collected in Surber samples was positively correlated with the  
256 taxa richness of equivalent kick samples ( $r_s = +0.84$ ,  $p < 0.001$ ). Correlations for taxa richness were  
257 stronger than for measures of abundance, but there was still considerable scatter (Figure 3a, b). In  
258 general samples collected following the Surber sample methodology were more taxa rich than  
259 equivalent kick samples, with 70% of samples having more taxa in the Surber than the kick. The  
260 difference in species richness and EPT richness between kick and equivalent Surber samples was  
261 statistically significant in both cases ( $p < 0.001$ , in both cases).

262 The CCI calculated from Surber and kick net samples are positively correlated ( $r_s = +0.81$ ;  $p < 0.001$ )  
263 and are statistically similar ( $p = 0.499$ ) (Figure 3c), indicating similarity in the collection of rarer taxa  
264 between methods.

265

### 266 3.2. Biomonitoring scores

267 Paired-sample Student t-tests indicate that the differences between the BMWP, ASPT and WHPT  
268 calculated from kick and Surber sampled data were not statistically different for any metric ( $p = 0.06$ ;  
269  $p = 0.955$ ,  $p = 0.08$ , respectively). BMWP, ASPT and WHPT displayed strong, statistically significant  
270 correlations between Surber and kick sampled results (BMWP  $r = +0.85$ ,  $p < 0.001$ ; ASPT  $r = +0.88$ ,  $p$   
271  $< 0.001$ ; WHPT  $r = +0.93$ ,  $p < 0.001$ ). There was scatter in each relationship, but slightly more  
272 variance was explained for WHPT ( $R^2 = 0.87$ ) than for the ASPT ( $R^2 = 0.78$ ) and BMWP ( $R^2 = 0.74$ )  
273 (Figure 4).

274 The difference between each of the four stress-sensitive metrics when calculated on Surber and kick  
275 sampled data were statistically indistinguishable (Saprobic  $p = 0.656$ ; TRPI  $p = 0.147$ ; PSI  $p = 0.143$ ;  
276 LIFE  $p = 0.166$ ) (Figure 5). All four metrics showed a strong relationship between Surber and kick  
277 sampled data, and were all significantly positively correlated ( $p < 0.001$  in all cases). The strongest  
278 association between kick and Surber sampled data was for the PSI and LIFE scores, both of which are  
279 based on the proportion of sensitive invertebrates to all sampled invertebrates.

280 The TRPI score displayed the lowest  $R^2$  of the stress specific metrics, although the  $R^2 = 0.78$  still  
281 suggests a strong relationship between kick and Surber sampled results. The TRPI was affected by  
282 two outliers where the Surber sample scored 100% whereas the equivalent kick sample scored  
283 substantially less. When these two outliers were removed,  $R^2$  increases to 0.90.

284 Comparing kick and Surber methods taken in the spring with those collected during the autumn  
285 indicated that spring samples were generally more consistent between sampling methods (Table 5).  
286 There was more variation between the two sampling methods in autumn for all biological metrics,  
287 with the exception of the BMWP, ASPT and Saprobic index, which were slightly more consistent in  
288 the autumn.

289

### 290 3.3. Score interpretation

291 Differences between biomonitoring scores calculated on Surber and kick sampled data are sufficient  
292 to alter the resulting classification of 35 (15%) of the biometric scores (Table 6). In 17 cases, the kick  
293 samples returned a higher class category than the Surber sample method, whereas the reverse was  
294 true in 18 cases. On average, the BMWP calculated using the Surber sample methodology was 63%  
295 of a class boundary greater than the kick sampled equivalent. The ASPT differed by an average of  
296 22% of a class boundary and the saprobic index by 15% of a class boundary.

297 The LIFE score differed by 19% and the PSI by 19% of a class boundary and the equivalent value for  
298 the TRPI was 23% (Table 7). In general, kick samples returned higher ranking of the PSI and Saprobic  
299 Index. As the scoring systems were continuous, rankings could be altered by small increments in  
300 score if they fall close to the class boundary. To assess the likelihood that a difference in sampling  
301 method would lead to different class interpretation, the difference between kick and Surber sample  
302 methodology scores was presented as a percentage of the number within each class (Table 7).

303

#### 304 3.4. Preferential sampling of particular species

305 Across all aggregated sites, some species of invertebrate were consistently more likely to be caught  
306 using the Surber sample than by the equivalent kick sample method and, to a lesser extent, the  
307 opposite was observed for a small number of taxa. Some invertebrates, such as *Gammarus pulex*  
308 and *Baetis* sp., were recorded at much greater abundances in the Surber sample than the kick  
309 sample method (Figure 6). For example, nearly twice as many *Agapetus* sp. caddisfly and three-times  
310 as many Simuliid blackfly larvae were found in total across all Surber samples. In contrast, kick  
311 samples caught more *Limnephilus lunatus* (cased caddisfly larvae) and the amphipod shrimp  
312 *Crangonyx pseudogracilis* than equivalent Surber samples (Figure 6). Whilst more abundant, these  
313 invertebrate taxa were not found at more sites and, consequently, any sampling bias did not alter  
314 biological metrics between methods. However, some invertebrates were found at more sites, with  
315 potential implications for biomonitoring scores (Figure 7). Notable examples were the bivalve  
316 *Pisidium* sp. and the caseless caddisfly *Lype reducta* which were both recorded in more kick samples  
317 than equivalent Surber samples (6 and 5 more sites, respectively; Figure 7). In contrast, the leeches  
318 *Piscicola geometra* and *Helobdella stagnalis* were recorded in 8 more Surber samples than kick  
319 samples. There were 27 taxa only recorded in Surber samples in contrast to 7 taxa only found in kick  
320 samples (Supplementary A). Those only found in kick samples were only ever observed at one site  
321 whereas some of the invertebrates only recorded in Surber samples were sometimes found at  
322 multiple sites.

323

324 **4. Discussion:**

325 *4.1. Sensitivity of biomonitoring scores to sampling method*

326 Differences in the invertebrate community collected using the Surber and kick sample methods  
327 affect the biomonitoring scores that are derived to varying degrees and levels of significance. The  
328 BMWP was most affected, because this is calculated by aggregating the score associated with  
329 identified families. Hence, any increased diversity of Surber samples leads to higher BMWP scores.  
330 The effect of different sample sizes was reduced to some extent by the ASPT score, which was more  
331 similar between sampling methods. The WHPT was most consistent, with this method both  
332 averaging the score by the total abundance, as well as standardising invertebrate scores by  
333 individual family abundance within the sample.

334 Stress-specific scores were similar for data collected via Surber and kick sampling. Kick and Surber  
335 sample LIFE and PSI scores were both highly correlated and statistically similar. However, for the  
336 sites sampled here, the PSI was consistently higher for the kick sample, probably because the kick  
337 sample was not as effective at collecting sediment-dwelling invertebrates which tend to reduce the  
338 score. One explanation for this is that hand disturbance of surface grains and the aim to disturb  
339 sediment to 10 cm depth in Surber samples is likely to dislodge more subsurface material.  
340 Furthermore, the Surber net has a wider mouth for sample collection and hand sampling causes less  
341 hydrodynamic disturbance than kick sampling (which may drive some animals around the net  
342 entrance). The Surber net also has retention sides or curtains at the mouth to aid sample capture  
343 which the kick-sweep net does not.

344 The saprobic and TRPI were also consistent between kick and Surber sampling, although the latter  
345 was affected by an outliers. These are important findings for the Saprobic index since loss and gain  
346 of species numbers have indicated a strong mechanistic link with elevated and declining organic  
347 enrichment (BOD levels) across UK rivers with differing Surber and kick sampling techniques  
348 employed in recent years (Clews and Ormerod, 2009; Durance and Ormerod, 2009; Overall, 2010).

349

350 *4.2. Sensitivity of water- and habitat-quality to sampling method*

351 Variance between biomonitoring scores calculated with kick and Surber sampled data can lead to  
352 different interpretations if a ranking classification is used. In the current investigation, all scores  
353 differed on average by less than a single class, although the BMWP does differ on average by 63% of

354 a class boundary. However, this was largely associated with very high scoring Surber samples where  
355 the equivalent kick sample was already in the top class. Hence, the BMWP was actually the metric  
356 where boundary classifications were most consistent between the two methods examine. The least  
357 consistent was the PSI, despite being very highly correlated, statistically similar and with an average  
358 difference of only 19% of a class boundary. This is likely because many of the sites fell close to class  
359 boundaries and even a slight but consistent bias in kick sampled data was sufficient to under-  
360 represent sediment dwelling invertebrates.

361 Although not explicitly tested here, it is likely that the uncertainty due to the choice of Surber or kick  
362 sampling method is comparable to the uncertainty when comparing between different operators, at  
363 different times of year, in different areas. For example, there are natural seasonal variations in all  
364 biometrics because of temporal changes in macroinvertebrate community structure, life cycle stages  
365 and the concomitant response of the seasonally resident invertebrate communities to the  
366 ephemeral toxicity of contaminants (Hynes, 1970; Hellowell, 1989; Clements and Newman, 2002).  
367 Overall, metrics were more similar between kick and Surber samples in spring but this was  
368 dependent on the biomonitoring score used. Data presented here suggest that the difference in  
369 metrics at the same site between spring and autumn can be twice as great as the difference when  
370 comparing metrics collected using kick and Surber sampling techniques. This underlines the  
371 importance of sampling across known natural variations in invertebrate community structures and  
372 seasonal impacts of pollution to fully characterise water quality.

373 Previous research shows that inherent uncertainties in sampling and identifying macroinvertebrate  
374 samples can substantially exceed those described here, associated with sample collection. For  
375 example, Haase *et al.* (2010) audited river macroinvertebrate biomonitoring as part of an EU WFD  
376 requirement. A subset of samples processed by government agencies were re-processed by auditors  
377 who found that 29% of specimens and 21% of all taxon were overlooked when sorting and that  
378 individuals successfully selected in the sorting processes were correlated to body-size. Over 30% of  
379 taxa were identified differently between individuals and auditors, which was not biased towards  
380 harder to identify individuals. As a result of these differences, 34% of samples were categorised into  
381 a different quality classes. Similarly, Carter and Resh (2001) found in the USA that different methods  
382 of data collection, sub-sampling and sorting were commonly used yet these were known to yield  
383 different results. Here, leeches and flatworms were recorded preferentially when using the Surber  
384 sampler method which could be because of more limited detritus present in Surber samples, making  
385 these animals easier to distinguish than in the paired kick samples.

386

387 4.3. Sensitivity of invertebrate community to sampling method

388 The Surber sample method collected significantly more invertebrates (abundance) and a significantly  
389 greater diversity of invertebrate species than the kick sample method, both in spring and autumn.  
390 This is particularly true of the EPT taxa. For example, the Surber sample method collected twice as  
391 many *Ephemera danica* mayfly larvae when aggregated across all samples than equivalent kick  
392 samples (Figure 6). Similarly, invertebrates that attach themselves to the sediment were more  
393 prevalent in samples using the Surber sampler method (e.g. Simuliidae blackfly larvae) (Figure 6).  
394 This was expected given the increased sampling effort when compared to the three-minute kick  
395 sample method. The kick samples were limited to 3-minutes but Surber samples continued until all  
396 the surface area had been disturbed, resulting in a longer overall sampling time than kick samples.

397 Invertebrates that were found preferentially by one method over the other will potentially alter  
398 biomonitoring scores. An example is the cased caddisfly larvae, *Glossosoma* sp., which were  
399 recorded at seven sites using Surber sampling in comparison to only two kick sample sites. Other  
400 organisms more likely to be recorded using the Surber sampler than the kick sample method  
401 included the leeches *Helobdella stagnalis* and *Piscicola geometra* which were found in 15 and 17  
402 Surber samples, but only 5 and 10 kick samples, respectively. Similarly, the flatworm *Polycelis felina*  
403 was found in six more Surber samples than equivalent kick samples. It may be that these sediment-  
404 dwelling animals are caught more efficiently in Surber samples where sampling is attempted to a  
405 depth of 10 cm, ensuring that sub-surface material is thoroughly disturbed.

406 The only two organisms identified that were consistently observed in more kick samples than Surber  
407 samples, was the caseless caddisfly *Lype reducta*, which was found in seven of the 39 kick samples in  
408 comparison to only two of the equivalent number of Surber samples, and individuals in the bivalve  
409 genus *Pisidium*, which were found in 6 more kick samples than Surber samples. The reason for this is  
410 not clear, but in the case of *Lype reducta* it could possibly be because they are xylophagous and have  
411 a close association with coarse wood on the river bed.

412 These results are consistent with the study of Gillies *et al.* (2009) who found kick samples collected  
413 only 63% of taxa that were collected in quantitative Surber samples in New South Wales, Australia.  
414 Gillies *et al.* (2009) also found that kick samples were biased towards sampling large, abundant and  
415 widely distributed taxa, with those missed generally being smaller in size and rarer in the wider  
416 environment. In the current study, individual samples using the kick sample method were not  
417 obviously biased towards larger species, because even large invertebrates such as *Ephemera danica*  
418 (body length > 20 mm) were under-represented in kick samples. Similarly, Storey *et al.* (1991) found

419 that Surber and kick samples in south-western Australia were broadly similar, but with key  
420 differences represented by Sorensen's similarity coefficients of 66% in June and 61% in September.

421

#### 422 *4.4. Added value of a quantitative sample*

423 There is a great deal of data held in records that have been used to generate biomonitoring scores,  
424 which could provide additional, valuable information. However, where qualitative or semi-  
425 quantitative measures have been used, the comparability of data is not readily assessed given the  
426 lack of information about the proportion of the river bed or invertebrate population that has been  
427 sampled. Although kick samples here did generally under-represent some taxa, they did provide a  
428 sufficiently good estimate of the invertebrate diversity to provide statistically similar biomonitoring  
429 scores to the quantitative Surber sample. However, the kick sample did not provide a good estimate  
430 of the relative abundance of many species. Given that this information is not required for many  
431 biomonitoring scores, this does not affect the value of biological metrics calculated. However,  
432 without a good estimate of total abundance, it is difficult to make ecological assertions about the  
433 community. In addition, not quantifying the abundance of taxa may lead to loss of important  
434 information, such as changing abundance / occurrence through time which may be indicative of a  
435 chronic issue but which would not be identified by most biomonitoring scores unless species are also  
436 concurrently impacted from the community. The Surber sampling method used here provides a  
437 quantitative measure of population (e.g. the abundance / m<sup>2</sup>), so it provides added value over semi-  
438 quantitative methods, allowing a more thorough investigation of the data, which may lend support  
439 or add detail to the information gained from the use of biological metrics.

440

#### 441 **5. Conclusions:**

442 This study set out to establish the extent to which community, biomonitoring scores, and inferred  
443 environmental conditions, are sensitive to the choice of invertebrate sampling method. Our analysis  
444 was based on an English data set covering 20 sites, 205 taxa and 128,129 identified organisms. We  
445 found that the overall taxa richness of aquatic invertebrates that were collected in quantitative  
446 Surber samples were greater than semi-quantitative kick sample equivalents, although the two were  
447 positively correlated. Surber samples enable additional ecological information and analysis to be  
448 undertaken and, at least at the sites studied here, gave a more complete overview of the abundance  
449 and diversity of macroinvertebrates. However, biomonitoring scores did not differ significantly in  
450 most cases and, therefore, a semi-quantitative kick sample methodology provided a suitable

451 estimate of the river health of the chosen sites. In particular, specific pressure based biomonitoring  
452 scores which use an abundance weighting (ratio of sensitive to total invertebrate abundance), such  
453 as the LIFE, PSI and TRPI scores, yielded very similar results, regardless of the sampling method.

454 The comparability of biometric indices from Surber and kick-sweep net sampling raises the  
455 possibility of using historical Surber net sample data to assess longer-term trends in biological stress  
456 signatures. Based upon the findings here, a wider use of replicated Surber net sampling is proposed,  
457 particularly where it is necessary to detect rare taxa that may be endangered or for 'one-off'  
458 quantitative and statistically testable benchmarking of ecological condition in river reaches,  
459 additional to routine regulatory monitoring programmes.

460

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464 to thank two anonymous reviewers for their helpful comments.

465



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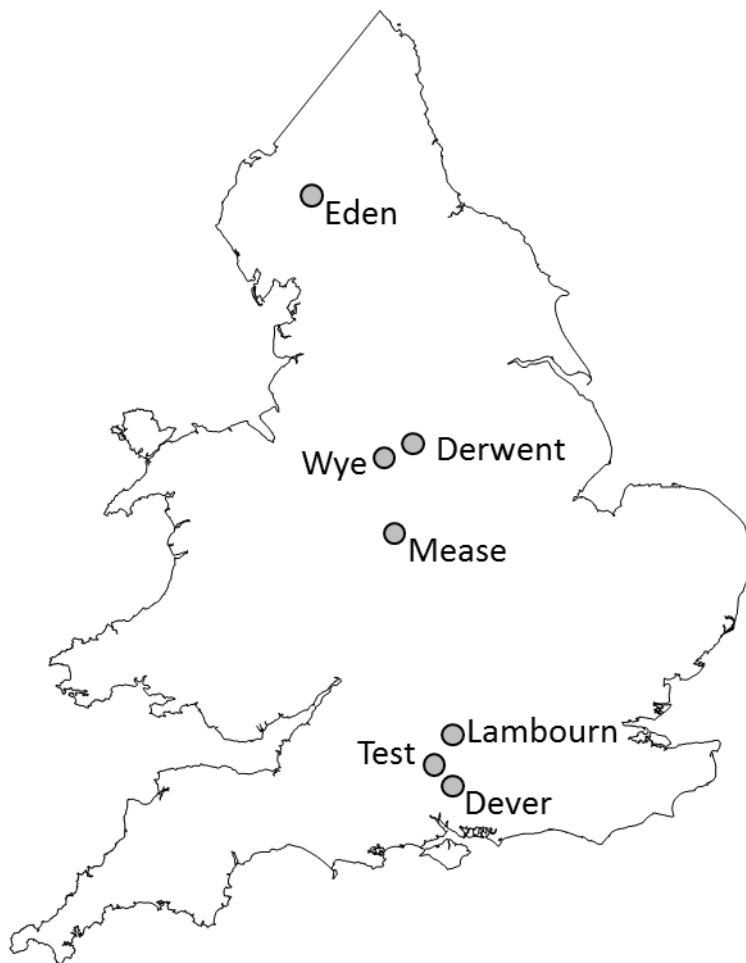
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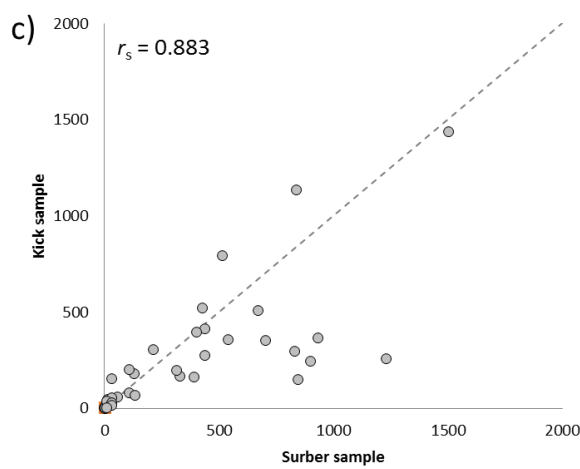
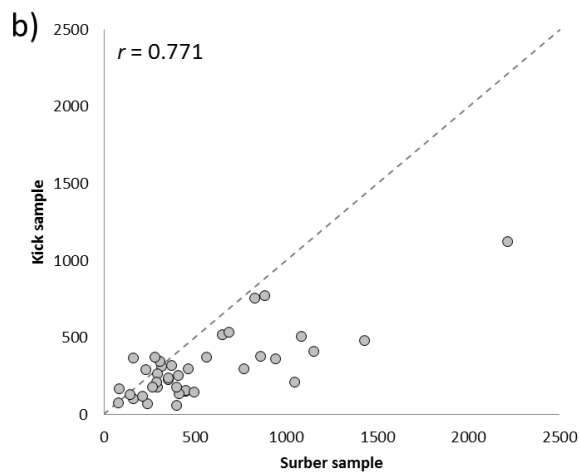
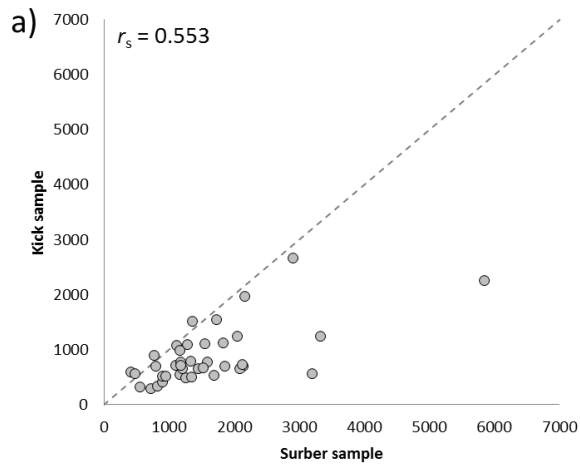
628 **Figure 1:** A map of England and Wales with the 7 sampled rivers with circles.



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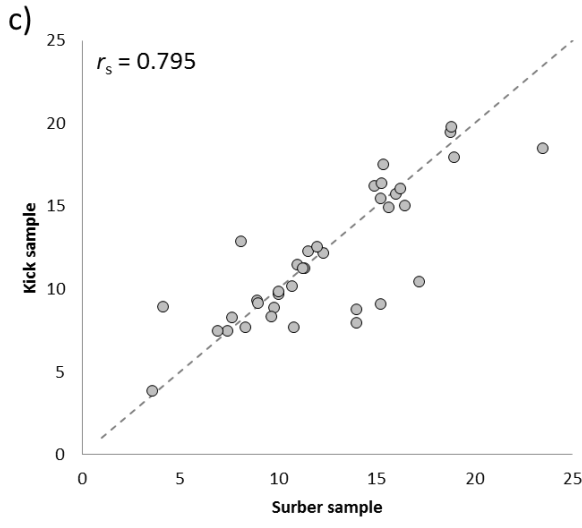
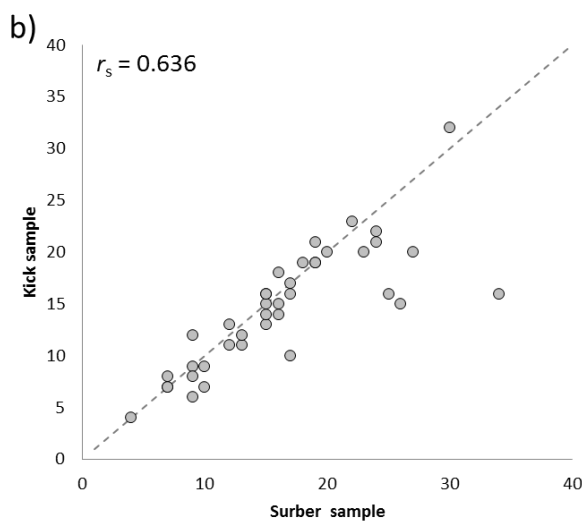
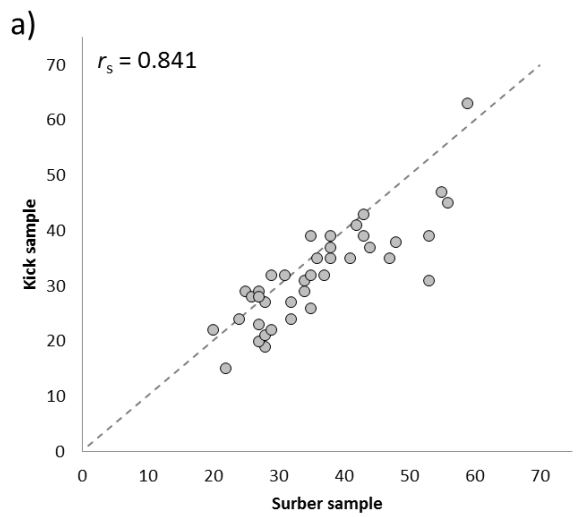
631 **Figure 2:** Relationship between the a) total invertebrate abundance, b) total EPT and c) total  
632 *Gammarus* collected in Surber samples versus kick samples, taken on the same day and at the same  
633 site. Pearson ( $r$ ) and Spearman ( $r_s$ ) correlation coefficients are included for normal and non-  
634 parametric data, respectively.



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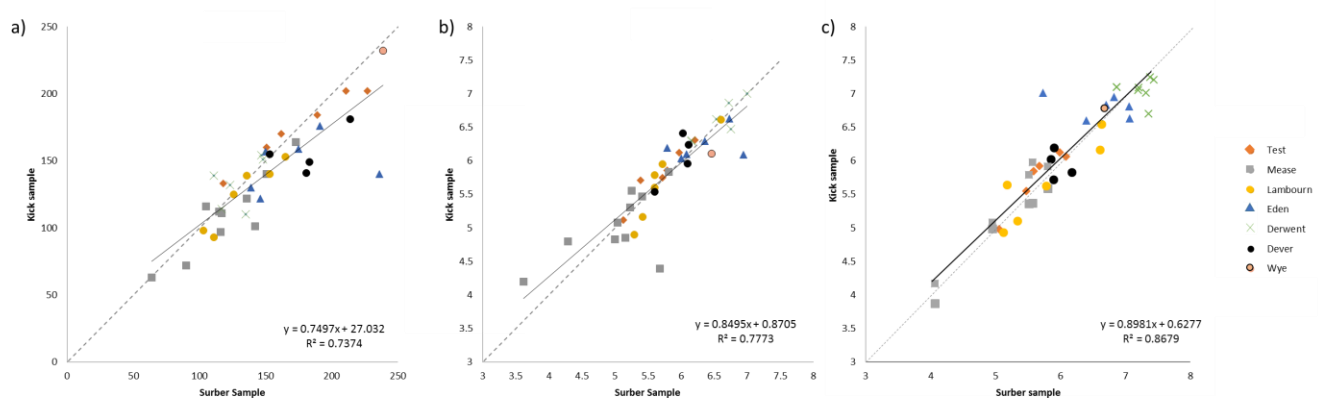


636 **Figure 3:** Relationship between a) taxa richness, b) EPT richness (e.g. mayfly, stonefly and caddisfly;  
637 EPT) and c) the CCI collected in Surber and kick samples, taken on the same day and at the same site.  
638 Spearman ( $r_s$ ) correlation coefficients are included.



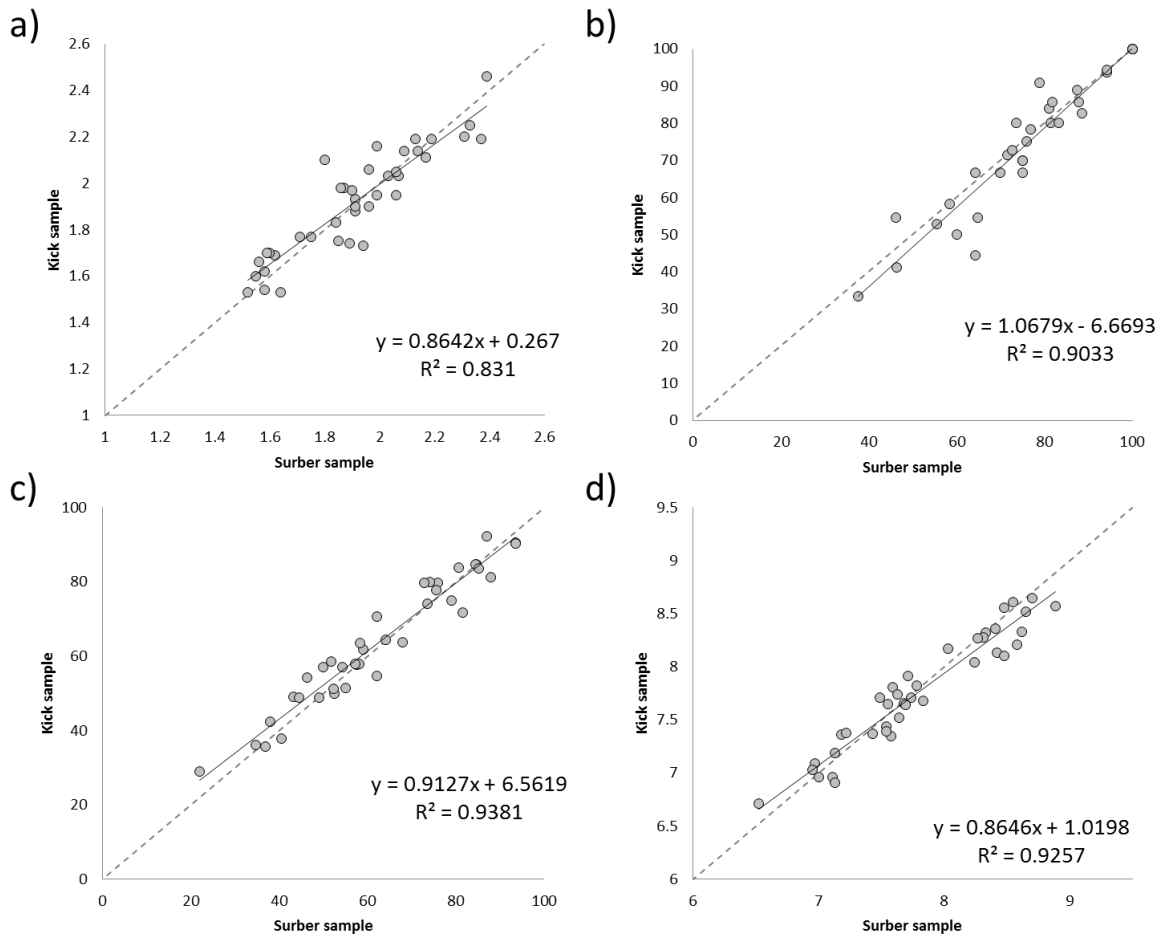
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640 **Figure 4:** Relationships between the (a) BMWP, (b) ASPT and (c) WHPT calculated using data from  
 641 Surber versus kick samples, taken on the same day and at the same site. Points are colour-coded to  
 642 designate the river where the sample was taken.



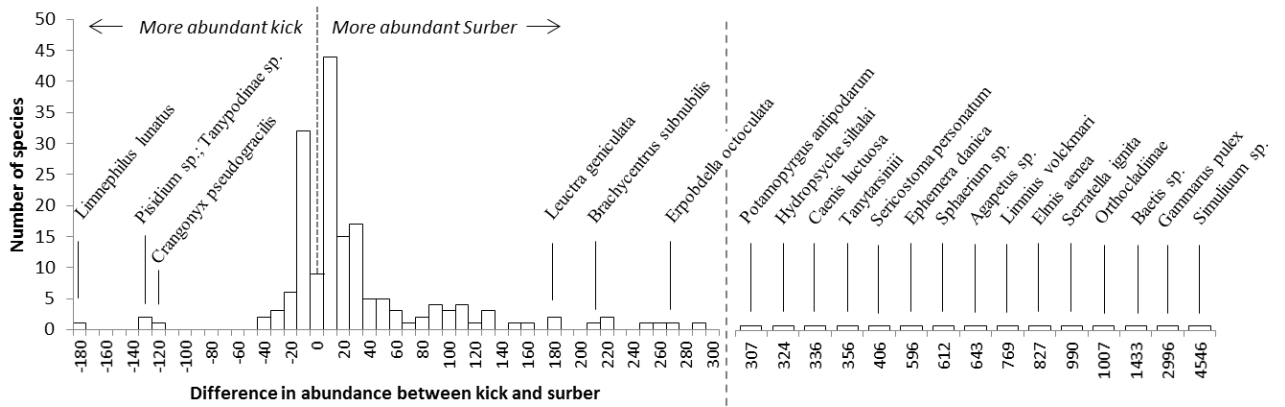
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645 **Figure 5:** Relationships between a) the Saprobic index, b) the TRPI, c) PSI and d) LIFE score calculated  
646 on Surber and kick samples, taken on the same day and at the same site.



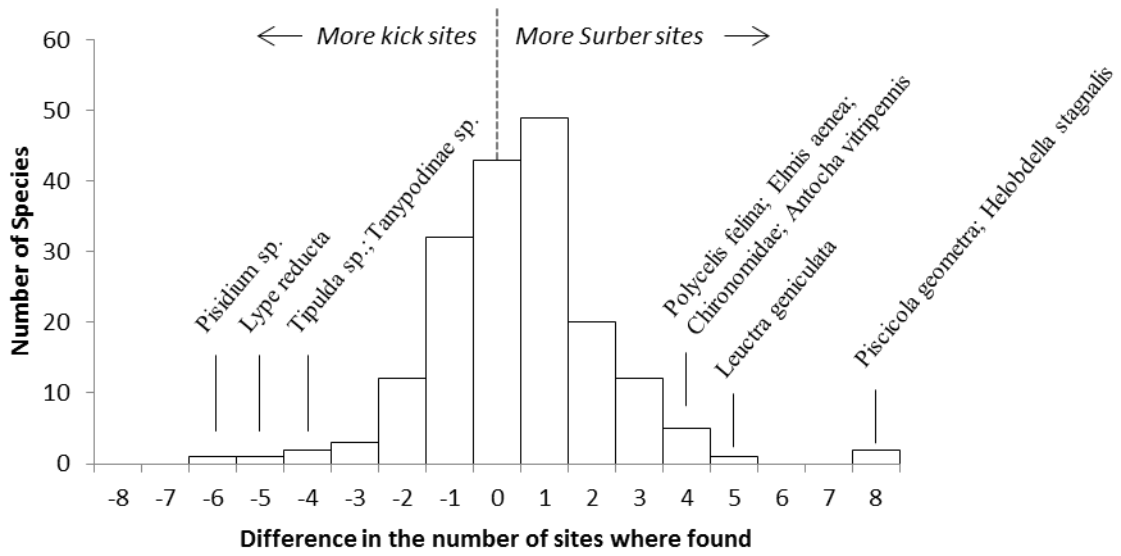
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648 **Figure 6:** The difference in abundance between kick and Surber samples for invertebrate taxa,  
 649 aggregated across all sites. Note that the right-hand grey dashed line marks a transition in the  
 650 horizontal axis from categorical values to absolute values. Taxa of note due to large differences  
 651 between kick and Surber samples are labelled. Note that in some cases taxa were grouped to genus  
 652 level (e.g. Baetis sp.) because differences in the proportion of individuals successfully identified to  
 653 species level (as opposed to genus level) could otherwise have biased results.



654  
655

656 **Figure 7:** The difference in the number of sites where taxa were caught between kick and Surber  
 657 samples. Taxa that were found at four or more additional sites for one method are labelled. Zero  
 658 indicates the taxa was found in the same number of kick and Surber samples.



659

660 **Table 1:** The dates and locations of sampling sites with representative geographic, climatic and  
661 hydrologic information for the 7 rivers studied. Land cover proportions were derived from LCM2007  
662 imagery; precipitation information is taken from the UK Met Office 30 year average and discharge  
663 information is derived from a 44 year record of gauged flow from the National River Flow Archive.

	<b>Derwent</b>	<b>Dever</b>	<b>Eden</b>	<b>Lambourn</b>	<b>Mease</b>	<b>Test</b>	<b>Wye</b>
<b>Number of Sites</b>	3	2	3	3	5	3	1
<b>Grid Ref</b>	SK 24671 74452	SU 43300 41999	NY 55831 36050	SU 43371 70208	SK 22166 11370	SU 34838 21355	SK 24367 65787
<b>Date: Spring</b>	19/04/2015	24/04/2015	24/04/2015	14/04/2015	17/05/2013	05/03/2013	22/05/2013
<b>Date: Autumn</b>	14/10/2015	29/09/2015	09/09/2015	01/10/2015	12/09/2013	24/09/2013	
<b>Upstream catchment (km<sup>2</sup>)</b>	203	122	616	176	167	453	154
<b>Geology</b>	<i>Carboniferous sandstone</i>	<i>Cretaceous Chalk</i>	<i>Permian &amp; Triassic Sandstones</i>	<i>Cretaceous Chalk</i>	<i>Triassic sandstone/ Mercia mudstone</i>	<i>Cretaceous chalk / Paleogene clay</i>	<i>Carboniferous Mudstone</i>
<b>Arable / Grassland (%)</b>	53	57	81	52		46	84
<b>Woodland cover (%)</b>	10	10	5	9		15	4
<b>Urban cover (%)</b>	0.2	0.5	0.4	0.4		1.6	2.3
<b>Site elevation (masl)</b>	139	50	92	96		10.1	139
<b>Annual Precipitation (mm)</b>	1325	780	1146	745		790	1166
<b>Average discharge (m<sup>3</sup> s<sup>-1</sup>)</b>	5.0	1.11	15.0	1.04	N/A	11.2	1.0
<b>Q<sub>10</sub> (m<sup>3</sup> s<sup>-1</sup>)</b>	11.4	2.0	34.8	1.8	N/A	17.5	6.2

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667 **Table 2:** Definitions of ecological parameters and biological monitoring scores tested.

Parameter	Definition
<i>Community Parameters</i>	
Total abundance (A)	The total number of all collected invertebrate taxa
Total diversity (R)	The total number / richness of taxa collected
EPT abundance	The total number/ abundance of all collected Ephemeroptera; Plecoptera, Trichoptera taxa
EPT diversity	The total number / richness of all Ephemeroptera; Plecoptera, Trichoptera taxa
<i>Gammarus</i> abundance	The total number of all shrimp ( <i>Gammarus</i> sp.) collected
Community Conservation Index (CCI)	The national and regional rarity and therefore conservation value of the species community profile
<i>Water Framework Directive Assessment Tools</i>	
Biological Monitoring Working Party Score (BMWP)	The BMWP score calculated with family-level data. No metric for species level
Average Score Per Taxon (ASPT)	The ASPT calculated with family-level data. No metric for species level. It is the BMWP divided by the number of scoring families
Whalley Hawkes Paisley Trigg (WHPT) method	The WHPT is calculated with family-level data. No metric for species level. It uses BMWP scoring system, but scores are dependent on abundance of each scoring family.
<i>Specific Stressor Indicators</i>	
Saprobic Index (S)	The weighted average sensitivity of the invertebrate species community to organic pollution
Proportion of Sediment-sensitive Invertebrates (PSI)	The proportion of sediment-sensitive invertebrates at species level (PSI)
Lotic–invertebrate Index for Flow Evaluation (LIFE)	The proportion of flow-sensitive invertebrates at species level (LIFE)
Total Reactive Phosphorous Index (TRPI)	The proportion of invertebrates sensitive to total reactive phosphorous impact at family level (TRPI)

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675 **Table 3:** Class rankings for each biological parameter used, where 1 indicates highly  
 676 impacted/polluted conditions and 5 indicates un-impacted conditions. An indication of whether the  
 677 classification is based on the authors' judgement or established knowledge is also given.

Rank	1 (v. poor)	2	3	4	5 (v. good)	
A	≤ 99	100 - 249	250 - 999	1000 - 4999	> 5000	Judgement
R	< 14	15 - 24	25 - 34	35 - 44	> 44	Judgement
EPT	≤ 1	2 - 9	10 - 19	20 - 29	> 30	Judgement
CCI	0 - 5	5 - 10	10 - 15	15 - 20	> 20	Chadd and Extence 2004
BMWP	0 - 35	36 - 50	51 - 70	71 - 95	> 96	Hellawell, 1986
ASPT	< 5	< 5	5 - 6	6 - 6.5	> 6.5	Hellawell, 1986; Wright et al. 2000
S	3.2 - 4	2.7 - 3.19	2.3 - 2.69	1.81 - 2.29	1.0 - 1.8	Schmidt- Kloiber and Hering 2015b
PSI	0 - 20	21 - 40	41 - 60	61 - 80	81 - 100	Extence et al. 2011
LIFE	< 6	6 - 6.49	6.5 - 6.99	7 - 7.99	> 8	Extence et al. 1999
TRPI	0 - 20	21 - 40	41 - 60	61 - 80	81 - 100	Everall 2010

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680 **Table 4:** The number of sites where either Surber or kick samples were more abundant in terms of  
 681 total abundance, Gammarus abundance, and EPT abundance. The average, maximum and minimum  
 682 difference in abundance, between Surber samples and kick samples is also given.

	Surber samples more abundant			Kick samples more abundant		
	Total (A)	<i>Gammarus</i>	EPT	Total (A)	<i>Gammarus</i>	EPT
Average	2.08	2.27	2.17	1.22	2.14	1.50
Max	5.71	5.68	6.73	1.41	4.88	2.27
Min	1.03	1.02	1.10	1.11	1.07	1.00
<i>n</i>	36	24	33	5	15	7

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685 **Table 5:** The gradient, intercept and amount of explained variance ( $R^2$ ) by linear regression between  
 686 biological monitoring scores derived from Surber and kick samples when performed on spring data,  
 687 autumn data, and spring combined with autumn data. All regressions were significant at  $p < 0.01$ .  
 688 Associated graphs can be seen as Supplementary Material B.

Score	Season	Gradient	Intercept	$R^2$
BMWP	<i>Spring</i>	0.944	23.615	0.60
	<i>Autumn</i>	0.929	14.88	0.79
ASPT	<i>Spring</i>	0.828	1.062	0.71
	<i>Autumn</i>	0.975	0.089	0.86
WHPT	<i>Spring</i>	0.929	0.376	0.93
	<i>Autumn</i>	0.854	0.924	0.80
PSI	<i>Spring</i>	1.062	6.712	0.97
	<i>Autumn</i>	0.984	1.459	0.90
LIFE	<i>Spring</i>	1.095	0.749	0.95
	<i>Autumn</i>	1.069	0.445	0.90
Saprobic	<i>Spring</i>	0.936	0.132	0.80
	<i>Autumn</i>	1.041	0.105	0.89
TRPI	<i>Spring</i>	1.095	0.749	0.95
	<i>Autumn</i>	0.996	7.011	0.63

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690 **Table 6:** Number of cases where differences in biomonitoring score calculated using kick and Surber  
691 samples results in that site being assigned to a different class. A score of 1 indicates the kick sample  
692 is one class above the equivalent Surber and -1 indicates the kick sample is one class below the  
693 equivalent Surber. The table also shows the difference in biomonitoring score as a percentage of the  
694 average class boundary. Values are shaded when the percentage difference is more than 50% of a  
695 class boundary. All sites on all rivers are included for samples taken in spring (Sp) and autumn (Au).

River	Site and Season		Number of classes different						% difference of class boundaries					
			BWMP	ASPT	PSI	S	TRPI	LIFE	BWMP	ASPT	PSI	S	TRPI	LIFE
Derwent	1	Sp	0	0	0	0	0	0	28	0	-14.6	14	1.6	-6.7
	1	Au	0	-1	0	0	0	0	-100	-28	0.8	10	0	-17.3
	2	Sp	0	0	0	0	0	0	112	9	-16.2	-22	0	-42.7
	2	Au	0	0	0	0	0	0	-12	14	1.5	8	0	-38.7
	3	Sp	0	0	0	0	0	0	12	14	26.8	20	-1.9	10.7
	3	Au	0	0	0	0	0	0	36	14	-33.8	20	0	-49.3
Dever	2	Sp	0	0	0	0	0	0	-132	13	-20.5	4	-5	-4.0
	2	Au	0	0	0	0	-1	0	-136	-14	14.0	-2	6.7	5.3
	3	Sp	0	-1	1	0	0	0	8	-6	0.3	-8	-41.7	-16.0
	3	Au	0	0	0	0	0	0	-160	38	42.4	-6	7.0	26.7
Eden	1	Sp	0	0	0	1	0	0	-64	-10	18.6	-30	-10.9	-38.7
	1	Au	0	0	0	0	-1	0	-96	2	-20.1	34	-50.8	-50.7
	2	Sp	0	-1	0	1	1	0	-384	-85	29.5	-42	60.3	-1.3
	2	Au	0	1	0	0	0	0	-36	40	33.9	0	11.9	18.7
	6	Sp	0	0	0	0	-1	0	-60	-7	2.2	-8	-7.4	-6.7
	6	Au	0	0	0	0	0	0	28	4	10.9	4	-25	-4.0
Lambourn	1	Sp	0	0	1	1	0	0	12	19	25.7	-20	19.5	14.7
	1	Au	0	0	0	0	0	0	-20	-26	3.8	-12	-25	13.3
	2	Sp	0	0	0	0	0	0	-4	23	1.5	22	-0.6	-20.0
	2	Au	-1	0	0	0	-1	0	-72	-39	-18.0	23	-99.3	-20.0
	3	Sp	0	0	0	0	0	0	-48	1	16.4	2	0	8.0
	3	Au	0	0	1	0	-1	0	-52	0	-49.5	14	-183.3	-26.7
Mease	1	Sp	0	0	0	0	0	0	-36	6	29.6	-8	-13.1	24.0
	1	Au	0	0	0	0	0	0	-44	2	-12.7	10	0	-8.0
	2	Sp	0	0	0	0	0	0	-164	-129	-0.7	0	31.8	-20.0
	2	Au	0	-1	0	0	0	-1	44	-17	-37.0	20	0	-30.7
	3	Sp	0	0	0	1	0	0	-72	51	35.2	-16	42.0	25.3
	3	Au	0	0	0	1	0	0	-4	58	39.8	-22	0	8.0
	4	Sp	-1	-1	1	0	0	1	-76	-31	22.6	14	-20.9	16.0
	4	Au	0	0	-1	0	0	0	-24	30	-6.3	0	0	-5.3
	5	Sp	0	0	0	1	0	1	-56	4	6.8	-36	-25.8	10.7
5	Au	0	0	-1	0	0	0	-12	7	22.5	02	0	21.3	
Test	1	Sp	0	1	1	0	0	0	-100	15	33.3	22	14.4	29.3
	1	Au	0	0	0	0	0	0	32	7	-5.3	-12	0	-13.3
	2	Sp	0	0	1	0	0	0	36	32	35.7	-22	-29.2	29.3
	2	Au	0	0	0	0	0	0	-20	3	-1.3	-2	-16.7	-2.7
	3	Sp	0	0	1	-1	-1	0	-36	10	14.0	60	-16.7	-6.7
	3	Au	0	0	-1	0	0	-1	60	-1	-13.5	12	-50.0	-29.3
Wye	1	Sp	0	0	0	0	0	0	-28	-35	-7.2	12	-62.5	0

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698 **Table 7:** Percentage difference between samples taken in spring and autumn, using both a kick and  
 699 Surber method. The percentage difference between kick and Surber samples in spring and kick and  
 700 Surber samples in autumn are also shown.

	Total Abundance	EPT	Gammarus	R	EPT	BMWP	ASPT	WHPT	PSI	Sap	TRPI	LIFE
<i>Between spring and autumn</i>												
Kick	71.7	137.6	704.4	25.3	37.7	20.8	9.1	6.4	14.6	8.9	19.9	3.4
Surber	46.3	88.8	709.4	29.2	36.7	26.4	9.0	5.3	11.8	7.8	46.3	3.0
<i>Between kick and Surber</i>												
Spring	99.4	107.3	81.0	19.9	20.5	15.3	4.7	3.4	6.7	5.1	5.0	1.6
Autumn	95.1	99.1	103.0	16.4	15.0	9.1	3.4	4.6	6.1	2.9	12.7	2.0

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706 **Supplementary Material A:** Taxa that preferentially occur in either kick or Surber samples. The  
707 difference in the number of samples between kick and Surber samples is presented, along with the  
708 percentage difference between kick and Surber samples. Only those taxa where the percentage  
709 difference is >50% are included.

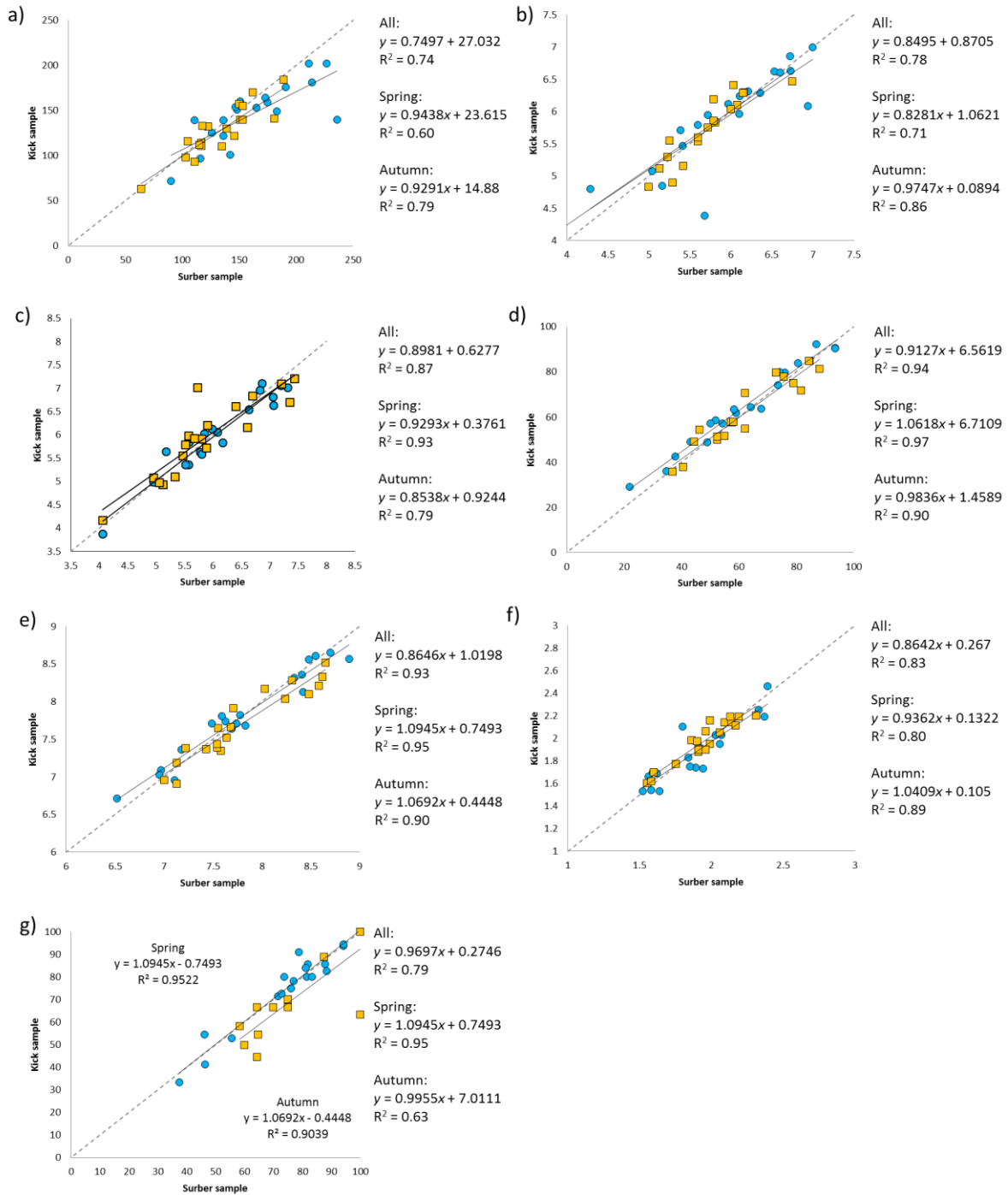
Phylum/ Class	Order	Family	Species name	% difference between kick and surber	Difference in number of samples
Insecta	Coleoptera	Dytiscidae	<i>Agabus didymus</i>	100% in kick	1
Insecta	Coleoptera	Dytiscidae	<i>Scirtes</i> sp.	100% in kick	1
Insecta	Diptera	Muscidae	<i>Limnophora</i> sp.	100% in kick	1
Insecta	Diptera	Ptychopteridae		100% in kick	1
Annelida	Arhynchobdellida	Erpobdellidae	<i>Erpobdella testacea</i>	100% in kick	1
Platyhelminthes	Tricladida	Planariidae	<i>Polycelis tenuis</i>	100% in kick	1
Crustacea	Decapoda	Astacidae	<i>Austropotamobius pallipes</i>	100% in kick	1
Insecta	Trichoptera	Psychomyiidae	<i>Lype reducta</i>	71% in kick	5
Mollusca	Veneroida	Sphaeriidae	<i>Pisidium</i> sp.	50% in kick	6
Insecta	Plecoptera	Perlidae	<i>Dinocras cephalotes</i>	100% in Surber	1
Insecta	Ephemeroptera	Baetidae	<i>Centroptilum luteolum</i>	100% in Surber	1
Insecta	Ephemeroptera	Heptageniidae	<i>Ecydonurus dispar</i>	100% in Surber	1
Insecta	Ephemeroptera	Ephemeridae	<i>Ephemera vulgata</i>	100% in Surber	1
Insecta	Trichoptera	Hydroptilidae	<i>Agraylea multipunctata</i>	100% in Surber	2
Insecta	Trichoptera	Leptoceridae	<i>Ceraclea nigronervosa</i>	100% in Surber	1
Insecta	Trichoptera	Glossosomatidae	<i>Glossosoma</i> spp.	100% in Surber	3
Insecta	Trichoptera	Limnephilidae	<i>Hydatophylax infumatus</i>	100% in Surber	1
Insecta	Trichoptera	Limnephilidae	<i>Limnephilus marmoratus</i>	100% in Surber	1
Insecta	Trichoptera	Leptoceridae	<i>Mystacides azurea</i>	100% in Surber	1
Insecta	Trichoptera	Hydroptilidae	<i>Oxyethira</i> spp.	100% in Surber	1
Insecta	Trichoptera	Phryganeidae	<i>Phryganea grandis</i>	100% in Surber	1
Insecta	Trichoptera	Limnephilidae	<i>Potamophylax</i> spp.	100% in Surber	1
Insecta	Trichoptera	Leptoceridae	<i>Ylodes conspersus</i>	100% in Surber	1
Insecta	Trichoptera	Hydropsychidae	<i>Hydropsyche angustipennis</i>	100% in Surber	1
Insecta	Trichoptera	Hydropsychidae	<i>Hydropsyche contubernalis</i>	100% in Surber	2
Insecta	Trichoptera	Hydropsychidae	<i>Hydropsyche</i> sp.	100% in Surber	2
Insecta	Trichoptera	Polycentropodidae	<i>Plectrocnemia conspersa</i>	100% in Surber	2
Insecta	Trichoptera	Polycentropodidae	<i>Polycentropus irroratus</i>	100% in Surber	1
Insecta	Coleoptera	Halipidae	<i>Brychius elevatus</i>	100% in Surber	2
Insecta	Coleoptera	Halipidae	<i>Halipus</i> spp.	100% in Surber	1
Insecta	Diptera	Empididae	<i>Chelifera</i> sp.	100% in Surber	1
Insecta	Diptera	Ptychopteridae	<i>Ptychoptera</i> sp.	100% in Surber	2
Mollusca		Bithyniidae	<i>Bithynia leachi</i>	100% in Surber	1
Mollusca		Planorbidae	<i>Planorbis carinatus</i>	100% in Surber	1
Mollusca	Veneroida	Sphaeriidae	<i>Pisidium nitidum</i>	100% in Surber	2
Insecta	Plecoptera	Leuctridae	<i>Leuctra geniculata</i>	83% in Surber	5
Insecta	Diptera	Muscidae	<i>Lispe</i> spp.	80% in Surber	4
Insecta	Diptera	Chironomidae		67% in Surber	4
Annelida	Rhynchobdellida	Glossiphoniidae	<i>Helobdella stagnalis</i>	67% in Surber	8

Annelida	Rhynchobdellida	Piscicolidae	<i>Piscicola geometra</i>	53% in Surber	8
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712 **Supplementary Material B: Linear regressions of Surber versus kick samples, collected on the same**  
 713 **day and at the same site for a) BMWP, b) ASPT, c) WHPT, d) PSI, e) LIFE, f) Saprobic and g) TRPI**  
 714 **scores. Blue circles were taken in spring and orange squares in autumn.**



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