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Quantifying trophic interactions and niche sizes of juvenile fishes in an invaded riverine cyprinid fish community

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Abstract

Quantifying feeding interactions between non-indigenous and indigenous fishes in invaded fish communities is important for determining how introduced species integrate into native food webs. Here, the trophic interactions of invasive 0+ European barbel *Barbus barbus* (L.) and the three other principal 0+ fishes in the community, *Squalius cephalus* (L.), *Leuciscus leuciscus* (L.) and *Phoxinus phoxinus* (L.), were investigated in the River Teme, a River Severn tributary in Western England. *Barbus barbus* has been present in the River Teme for approximately 40 years. Analyses of stomach contents from samples collected from three sites between June and September 2015 revealed that, overall, fishes displayed a generalist feeding strategy, with most prey having low frequency of selection. Relationships of diet composition versus body length and gape height were species-specific, with increasing dietary specialisms apparent as the 0+ fishes increased in length and gape height. The trophic niche size of invasive *B. barbus* was always significantly smaller than *S. cephalus* and *L. leuciscus*, and was significantly smaller than *P. phoxinus* at two sites. This was primarily due to differences in the functional morphology of the fishes; 0+ *B. barbus* were generally restricted to foraging on the benthos, whereas the other fishes were able to forage on prey present throughout the water column. Nevertheless, the invasive *B. barbus* were exploiting very similar prey items to populations in their native range, suggesting these invaders were strongly pre-adapted to the River Teme and this arguably facilitated their establishment and invasion.
Introduction

Invasions by non-indigenous fishes can increase inter-specific competition in fish communities, potentially leading to impacted native species having reduced growth and survival rates, and/or being displaced from their original niche (Gozlan et al. 2010). Quantifying feeding interactions between the invasive and extant fishes in the community is thus important for determining the extent of the invasion-mediated shifts in the trophic structure of the food web (Jackson et al. 2012; Cucherousset et al. 2012; Copp et al. 2016).

Ecological theory suggests that these shifts in trophic structure can include the invader occupying an unexploited niche (Shea and Chesson 2002). This will limit their inter-specific competitive interactions and facilitate their integration into the ecological community (Shea and Chesson 2002; Tran et al. 2015). Alternatively, when food resources are more limiting, the niche variation hypothesis suggests that increased inter-specific competition can result in the trophic niches of the competing species to constrict and diverge due to diets becoming more specialised (Van Valen 1965; Olsson et al. 2009; Tran et al. 2015). Conversely, this can result in the trophic niche sizes of competing species to increase, as individuals utilize a wider resource base to maintain their energy requirements (Svanbäck and Bolnick 2007).

When invasive and native species coexist for prolonged periods, high overlaps in their trophic niches can suggest a lack of competitive interactions, perhaps due to resources not being limiting, and so facilitating co-existence (Pilger et al. 2010; Guzzo et al. 2013). However, prolonged co-existence can also result in competitive exclusion, where the invader eventually excludes a native species from its original niche and results in its population decline (Bøhn et al. 2008).
The ability of an introduced fish to develop invasive populations depends on their ability to establish sustainable populations, with reproduction and recruitment being key processes. Consequently, the larval and juvenile life-stages of fishes (‘0+ fishes’) are important in the overall invasion process due to their influence on recruitment (Nunn et al., 2003, 2007a, 2010a). A range of factors influences the growth and survival rates of 0+ fishes, including their ability to capture and ingest the prey items and sizes available (Nunn et al., 2012). If preferred prey items are unavailable, reduced growth rates and/or starvation can occur, with potentially deleterious consequences for that 0+ cohort (Dickmann et al., 2007; Burrow et al., 2011). Where an introduced fish shares food resources with indigenous fishes and these resources become limiting, this can affect 0+ fish food acquisition and assimilation, and growth and survival rates, and so potentially impedes their ability to recruit and, therefore, establish (Gozlan et al., 2010; Dick et al., 2014, 2017).

The feeding ecology of mature fishes is relatively well understood, including for temperate riverine cyprinid fishes (e.g. Mann, 1974; Nunn et al., 2012). Extant knowledge includes how diet plasticity can assist the establishment of populations of introduced fishes (Basic et al., 2013; Tran et al., 2015). In contrast, the feeding ecology of 0+ fishes is often poorly understood (Nunn et al., 2012), especially within invaded communities (Britton et al., 2009). This is despite developmental shifts in diet often being important for 0+ fish survival (DeVries et al., 1998). In general, most freshwater fishes are planktivorous at the onset of exogenous feeding, with zooplankton being an important larval prey resource (Nunn et al., 2007b, 2010). Thereafter, diets of juvenile riverine cyprinids in temperate regions tend to consist of a mix of cladocerans, copepods and insect larvae, with some species also exploiting adult dipterans and Aufwuchs (the periphyton and associated microfauna that grow on underwater surfaces) (Nunn et al., 2012). However, as individuals increase in body and gape...
sizes, there is a general shift towards each species developing specific dietary traits that can result in considerable inter-specific diet and niche differences (Nunn et al., 2007b, 2012). As the ability to assimilate adequate energy has important implications for lengths achieved at the end of the first growth year, this can affect over-winter survival, as larger individuals tend to have higher over-winter survival rates (Nunn et al., 2007a,b, 2010).

The aim of this study was to quantify the trophic interactions of a riverine community of 0+ cyprinid fishes invaded by a non-indigenous fish, European barbel *Barbus barbus* (L.). This fish is indigenous to some European rivers but has been widely introduced outside of their natural range for enhancing angling, in countries including Italy and England (Britton & Pegg, 2011). The study system was the River Teme, a River Severn tributary in western England, where *B. barbus* is non-indigenous and invasive (Wheeler & Jordan, 1990; Antognazza et al., 2016). The introduction of *B. barbus* into the River Severn was in 1956, with the species then dispersing through much of the basin (Wheeler & Jordan, 1990). *Barbus barbus* began to be captured by anglers in the River Teme in the 1970s, indicating they have been present in the study river for approximately 40 years (Antognazza et al. 2016). The fish assemblage of the River Teme is relatively species poor; the only other cyprinids present are minnow *Phoxinus phoxinus* (L.), chub *Squalius cephalus* (L.) and dace *Leuciscus leuciscus* (L.). Some salmonid fishes are also present, including grayling *Thymallus thymallus* (L.).

Through application of stomach contents analyses (SCA) (Hyslop, 1980) to quantify 0+ fish diet on samples collected during 2015, the study objectives were to: (1) quantify diet composition across the community of 0+ fishes, with assessment of inter-specific similarity and spatial patterns; (2) identify shifts in the diet composition of each species and in relation to body length and gape size; and (3) quantify trophic niche sizes per species and according
to gape size, with assessment of the extent of inter-specific niche overlap between invasive *B. barbus* and other fishes. Given that invasive *B. barbus* and the other fishes of the study river have co-existed for approximately 40 years, it was predicted that the trophic niches of the 0+ fishes would be divergent through the fishes having developed strong dietary specialisms, as per the niche variation hypothesis that suggests invasions can result in trophic niche constriction and divergence via the development of dietary specialisms resulting from competitive interactions (Van Valen 1965; Olsson et al. 2009).

**Materials and Methods**

**Sampling sites and methodology**

Three sampling sites were used in the non-indigenous range of *B. barbus* in the River Teme (Fig. 1). Due to negligible off-channel habitat throughout the river, each sampling site consisted of areas of reduced flow rates within the river channel. Each site was separated by at least 5 km of river length was thus were considered as independent from each other, with the 0+ fish unable to intentionally move between them. Site 1 was the furthest upstream, located at Tenbury Wells (52°19’N, -2°24’W) (Fig. 1). The sampled areas were located immediately downstream of a road bridge at the downstream end of a large gravel island, near to the right-hand bank. Riparian vegetation included overhanging trees (*Salix* spp.) and, within the river, there was minimal in-stream vegetation, with the river generally running over gravel at depths of < 1m. Sampling areas comprised of large patches of minimal/ negligible flow in marginal areas where depths were generally < 1 m. Site 2 was located at Knightwick (52°12’N, -2°23’W) (Fig. 1), with samples generally collected at the downstream end of an exposed gravel beach where there were shallow patches (< 1 m depth) of low flow over gravel that created nursery habitat for 0+ fishes, but where instream vegetation was
minimal. Site 3 was the most downstream site (52°10’N, -2°14’W) (Fig. 1), with the sampling area located at the downstream end of a gravel riffle used by spawning *B. barbus* and, again, where there were shallow (< 1 m) patches of low and negligible flow over gravel, but with instream vegetation absent. Samples were collected on up to five occasions per site between July and October 2015 (Supplementary material: Table S1), with samples not collected thereafter due to elevated river levels throughout the winter period that prevented safe access to sampling sites.

Due to the restricted 0+ fish habitat of the River Teme and poor riparian access, point-abundance sampling by electric fishing was not an appropriate sampling method (Copp 2010). Micro-mesh seine netting was used instead, with acknowledgement that this would limit the proportion of larval fishes <15 mm in samples (Cowx et al. 2001; Copp 2010). On each sampling occasion, the 0+ fish were collected between 07.00 and 11.00, euthanised (MS222) and then preserved in 70 % IMS. Samples were unable to be collected at night for access and safety issues. These samples were then stored at 5 °C prior to their processing in the laboratory. All samples were processed in the laboratory within six months of sampling to minimise issues associated with shrinkage of body lengths related to preservation (Leslie & Moore, 2001).

Sample processing and data collection

There were four 0+ fish species, all of the Cyprinidae family, that were captured in sufficient numbers to enable subsequent dietary analyses: *B. barbus, S. cephalus, minnow Phoxinus phoxinus* (L.) and dace *Leuciscus leuciscus* (L.) (Table S1). In the laboratory, following identification to species level (Pinder, 2001), a maximum of 30 non-indigenous *B. barbus* and 20 individuals of the other fishes per site and per sample date were analysed. These numbers
of analysed fishes were achieved by sub-sampling within the collected samples, with this stratified to ensure the size ranges of fish present in each sample were covered. This involved their measurement using digital callipers (standard length, \(L_s\), to 0.01 mm). The majority of the fishes were already at juvenile stages (a consequence of the sampling method) and thus subsequent dietary analyses focused on these, rather than larval stages (Krupka, 1988; Pinder, 2001). Gape size was measured as the height of the mouth when open at its widest angle, using a stage micro-meter (Lukoschek & McCormick, 2001; Nunn et al., 2007b). The intestine (‘gut’) was then dissected, with gut fullness (%) estimated and the total gut contents extracted, mounted on a glass slide and fixed using Polyvinyl alcohol-lactic acid-glycerol (PVLG). Prey items were then identified to their lowest practicable taxonomic level using microscopy (to x100 magnification), with their number then counted to provide data on abundance. Periphytic biota (diatoms and similar material that was too small to classify more precisely) were classed as ‘Aufwuchs’. The amount of Aufwuchs in each gut was estimated on the basis of their percentage cover on the slide area and converted to a number (0 to 5 scale), similar to other studies (Garner 1996; Mann 1997), so that it was comparable to enumerated prey. As the majority of fishes had low proportions of Aufwuchs in the gut, this scale focused on slide coverage of below 55 % to allow greater discrimination between individual diets and thus greater precision in analyses. Thus, the scale used was: 0 (0 to 1 % coverage), 1 (2 to 3 %), 2 (4 to 7 %), 3 (8 to 20 %), 4 (21 to 55 %) and 5 (56 to 100 %).

A total of 37 distinct prey items were detected across the 0+ fish diets and thus, for some analytical purposes, these were categorised into the following 16 groups according to their taxonomy and functional ecology: Chironomid larvae, Aufwuchs, amphipods, winged insects, chalcid wasp, copepods, Cladocera, nymphs (stonefly and mayfly), Arachindae, Hemipteroids, saucer bugs, caddis larvae, beetles, beetle larvae, springtail (hexapods), seed/
spore/ plant material, and fish. The largest prey item in the gut of each individual fish was then measured; for Chironomid larvae this always consisted on measuring the width of the head.

Data analysis

Differences in fish standard length between the sites were tested initially using one-way ANOVA with a Tukey post-hoc test. The vacuity index (%Iv) (i.e. the proportion of fish with empty guts) was calculated from: \( %I_v = \frac{S_0}{S_1} \), where \( S_0 \) is the number of fish with empty guts and \( S_1 \) is the total number of larval and juvenile fish stomachs examined (Hyslop, 1980).

Frequency of occurrence of prey categories (\( F_i \)) represented the proportion of all guts that contain that prey category and was determined from: \( F_i = \frac{N_i}{N} \), where \( N_i \) is the number of guts in which that prey item i occurred and \( N \) is the total number of guts with prey present (Caillet, 1977). Relative abundance of a given prey category (%A_i) represented the proportion of total gut contents from all fish that comprised that prey category and was calculated from: \( %A_i = \frac{100(\Sigma S_i S_i)}{\Sigma S_i} \), where \( S_i \) is the number of prey items comprising prey i and \( S_{ti} \) is the total number of prey in all guts regardless of whether they contained prey item i (Macdonald & Green, 1983). Prey-specific abundance (\( P_i \)) represented the proportion of all prey that comprised of a specific prey category and was determined from data from only the guts in which prey items in that category were encountered. It was calculated from: \( P_i = \frac{100(\Sigma S_i S_{ii})}{\Sigma S_{ii}} \) here \( P \) is the number of prey items comprising prey i and \( S_{ii} \) is the total number of prey items in guts that contained prey item i (Amundsen et al., 1996).

The calculation of frequency of occurrence and prey-specific abundance enabled feeding strategy plots to be produced (Costello, 1990). These plots provided information about the importance of prey categories and feeding strategies of each species via examination of the...
distribution of points along the diagonals and the axes of the plot according to: prey importance (represented in the diagonal from the lower left (rare prey) to upper right (dominant prey), feeding strategy (represented in the vertical axis from the bottom (generalization) to top (specialization)), and the relationship between feeding strategy and the between or within-phenotype contributions to the niche width (represented in the diagonal from the lower right (high within-phenotype component, WPC) to upper left (high between-phenotype component, BPC)) (Amundsen et al., 1996; Leunda et al., 2008).

To test whether fish with larger body sizes consumed different prey items to smaller conspecifics, linear regression was used, with standard length as the independent variable and the percentage of specific prey items as the dependent variable. Where assumptions for the test were not met, the percentages of prey data were square-root transformed. Differences in gape height and standard length of the fishes were tested using general linear models, where gape height (µm) or standard length (mm) was the dependent variable and the independent variables were site and species. Differences in the maximum prey size per species were also tested using a general linear model; maximum prey size was the dependent variable, species was the independent variable and standard length was the covariate. This model structure was also used to test differences in maximum prey sizes according to sampling year and site. All general linear models were interpreted with regards to the significance of the independent variable on the dependent variable, the significance of covariates, and the estimated marginal means (i.e. mean values per group, adjusted for effect of covariate) and the significance of their differences according to independently linear pairwise comparisons with Bonferroni adjustment for multiple comparisons. To identify how body length, gape height and their interaction influenced the maximum prey size of each species, multiple regression was used. The outputs were the standardised β coefficients of each independent variable, where higher
values (irrespective of whether they were positive or negative) indicated a stronger
correlative effect on the dependent variable, plus their $R^2$ values and significance.

For plots of trophic niche size versus gape height per species, gape heights were classified
into five size groups: 0.8 to 1.4, 1.5 to 2.2, 2.3 to 3.1, 3.2 to 3.9 and 4.0 to 4.8 mm. These
groupings were based on the conversion of the stage micro-meter units to the actual gape
height of the fishes (in mm). In all analyses, gape heights above 4.8 mm were excluded from
analyses as the maximum for *B. barbus* was 3.1 mm. Trophic niche sizes were expressed as
standard deviation ellipses (40%), calculated using detrended correspondence analysis with
basic reciprocal averaging that was completed using the ‘decorana’ function in ‘vegan’
package v2.4 in R (R Core Team, 2016; Oksanen et al. 2017). This was completed within a
Bray-Curtis similarity matrix where all data were square root transformed for normality.
Ellipse areas then compared across the gape height classes for each species to determine their
influence on the size of the trophic niche.

Finally, to determine the differences in trophic niche sizes between species and sites, an
ANOVA was carried out using a permutational approach. This analysis was carried out in R
(R Core Team, 2017) using the vegan package (Oksanen et al. 2017), with the adonis
function used to complete a PERMANOVA analysis. All vacuous guts and guts containing
only diatoms were removed from the dataset prior to these analyses, plus three dietary items
that only occurred once. As the dietary composition data were expressed as percentages, they
were square-root transformed, followed by construction of a resemblance matrix with Bray-
Curtis similarity that enabled the PERMANOVA analysis to be calculated between species
and sites. To identify inter-specific differences, pairwise comparisons were carried out to
identify the significance of differences in niche sizes (Martinez Arbizu 2017). Drivers of
inter-specific difference by site were determined using a SIMPER analysis (PRIMER 7).

Results

Sample sizes, stages and lengths

Across the four 0+ fishes, SCA was performed on 878 individuals (B. barbus: n = 431; S.
cephalus: n = 174; L. leuciscus: n = 81; P. phoxinus: n = 192). Across the samples, no fish
were present at larval stage 1 and, as there was only one fish at larval stage 2, this individual
was removed from subsequent analyses (Table S1). As there were low numbers of fish
sampled at larval stages 3 to 5, and relatively high numbers of juvenile fishes (juvenile stages
6 to 9), these fish were all grouped together as ‘juveniles’ for analytical purposes (Table S1).
The minimum, maximum and mean lengths of these juveniles per species are provided in
Table 1. The low number of larvae in samples also meant that testing of ontogenetic diet
changes used fish lengths instead of larval stage.

Across the dataset, the standard length of B. barbus differed significantly between sites
(ANOVA: F_{2,428} = 3.97, P = 0.02), with fish at Site 1 being significantly larger than those at
Site 2 (Table 2). Similarly, S. cephalus at Site 2 were significantly smaller than the other sites
(ANOVA: F_{2,156} = 8.87, P < 0.01; Table 2). Phoxinus phoxinus were significantly smaller at
Site 3 than the other sites (ANOVA; F_{2,174} = 17.9, P < 0.01). As L. leuciscus was only sampled
at Site 3, no spatial comparisons were possible. Vacuity indices were generally low, with the
highest values in S. cephalus (up to 6 %) and lowest in B. barbus (0 to 0.6 %) (Table 2).
Relative frequency of prey and feeding strategies

Chironomid larvae were the most important prey item across the species, with values ranging between 44 % (S. cephalus) and 83 % (B. barbus) of diet, with Aufwuchs also a prominent item for all fishes (Table 2). There was variability in the contributions of prey categories between the fishes with, for example, Hemipteroids comprising of 7 % and 24 % of the diet of S. cephalus and L. leuciscus respectively, but less than 1 % for both B. barbus and P. phoxinus. Spatially, there was low variability in the relative frequencies of prey items in B. barbus diet, with Chironomid larvae being the dominant prey at all sites. In contrast, there was greater spatial variability in S. cephalus diet, for example in the proportion of hemipteroids (1 % at Site 3, > 10 % at other sites). For P. phoxinus, the major spatial differences were in the proportions of Chironomid larvae and Aufwuchs, although when combined, these prey categories still comprised between 85 and 94 % of their diet (Table 2).

Feeding strategy plots for each species suggested they were all generalists, with the majority of prey items having prey specific abundances of < 50 % with relatively low frequency of occurrences (Fig. 2). The relative high proportion of Chironomid larvae across the diet of each species was, however, strongly reflected in the feeding strategy plots, where their prey specific abundances ranged between 52 and 83 %. The most varied diet was in L. leuciscus, although the majority of prey categories had low frequency of occurrences and low prey specific abundances (Fig. 2). Spatially, there was little variability in the feeding strategy plots for B. barbus (Fig. S1), but with greater variability apparent for P. phoxinus and S. cephalus (Fig. S2, S3).
Fish length and gape height influences on diet

The relationship of gape height versus fish length was significant for each species (B. barbus: $R^2 = 0.81$, $F_{1,515} = 2247.0$, $P < 0.01$; S. cephalus: $R^2 = 0.86$, $F_{1,185} = 1095.0$, $P < 0.01$; L. leuciscus: $R^2 = 0.89$, $F_{1,106} = 738.4$, $P < 0.01$; P. phoxinus: $R^2 = 0.73$, $F_{1,158} = 435.4$, $P < 0.01$). Between the species, there were significant differences in gape height (GLM: Wald $\chi^2 = 1080.84$, df = 3, $P < 0.01$), with standard length a significant covariate ($P < 0.01$). Pairwise comparisons revealed the mean adjusted gape height of Barbus barbus (mean 2.02 ± 0.03 mm) was significantly smaller than the other three fishes (S. cephalus: 2.81 ± 0.05 mm; L. leuciscus: 2.38 ± 0.07 mm; P. phoxinus: 2.82 ± 0.05 mm; $P < 0.01$ in all cases).

Maximum prey sizes differed significantly between the fishes (GLM: Wald $\chi^2 = 197.12$, df = 3, $P < 0.01$), where the covariate of standard length was significant ($P < 0.01$). The mean maximum prey size of B. barbus (0.51 ± 0.02 mm) was significantly smaller than for S. cephalus (0.67 ± 0.05 mm; $P < 0.01$), was not significantly different to L. leuciscus (0.53 ± 0.06 mm; $P = 0.47$), and was significantly larger than P. phoxinus (0.35 ± 0.03 mm; $P < 0.01$). Multiple regression revealed that for B. barbus, standard length and gape height, and their interaction, were all significant variables, but with length explaining most the variation in the prey size ($P < 0.01$ in all cases) (Table 3). For S. cephalus, although gape height and standard length were both non-significant ($P > 0.05$), their interaction was a significant predictor of maximum prey size ($P < 0.01$). In L. leuciscus, standard length was the only significant predictor ($P < 0.01$), and none of the variables were significant predictors of maximum prey size in P. phoxinus ($P > 0.05$ in all cases), with individuals generally consuming much smaller prey than was possible for their gape height (Table 3).
Increases in gape height did not necessarily result in the development of a larger trophic niche across the 0+ fishes (Fig. 3). In *B. barbus* and *S. cephalus*, whilst the size of their trophic niches altered with gape height, it was largest *S. cephalus* at gape height of 2.5 to 3.1 mm and for *B. barbus* at 1.6 to 2.2 mm, with reductions thereafter (Fig. 3). For *P. phoxinus*, their largest trophic niches occurred in the two smallest gape height classes, suggesting their diet became more specialised as their gape height increased (Fig. 3).

*Spatial and inter-specific dietary comparisons*

There was a significant difference in niche size between the four species (PERMANOVA: *P* < 0.01) and across the three sites (PERMANOVA: *P* < 0.01) (Table 4). According to their niche sizes (as 40 % ellipse areas), *S. cephalus* had the largest niche of all species, with this significantly larger than *B. barbus* in all cases (Fig. 4; Table 5). The size of the *B. barbus* niche was significantly smaller than *L. leuciscus* at Site 3, and *P. phoxinus* at Site 2 and 3 (Table 5).

At Site 1, the niches of the three fishes present were generally discrete with low overlap (Fig. 4). At Site 2, the large niche of *S. cephalus* did not overlap with *B. barbus*, but the *B. barbus* niche sat within the larger niche of *P. phoxinus* (Fig. 4). At Site 3, the only site with all four fishes present, the niche of *B. barbus* had some overlap with all the other species, but with the niches of the other fishes having some differences, especially between *S. cephalus* and *L. leuciscus* (Fig. 4).
Discussion

This study successfully described the diet composition of 0+ fishes in a cyprinid fish community of low species richness that has been invaded by non-indigenous *B. barbus*. Overall, the 0+ fishes displayed a generalist feeding strategy, with most (but not all) prey categories having low selectivity according to feeding strategy plots. For some prey items in the diet, there were strong relationships with fish length, indicating the importance of increasing body size as a driver of dietary changes. There were, however, some differences in how the effects of body length and gape height manifested on diet composition, with dietary shifts in *B. barbus* and *S. cephalus* influenced strongly by their interaction, whereas in *L. leuciscus*, increased length was the only significant explanatory variable in their dietary changes.

The prediction was that the trophic niches of the 0+ fishes would be divergent, with this divergence developing according to the dietary specialisms of fishes. The results suggested some consistency with this prediction. Although the diets of all the fishes were described as generalist, they became more specialised as their body length and gape height increased. The prediction also included that the inter-specific niche divergence would be driven by competitive interactions, as per the niche variation hypothesis (Van Valen 1965; Olsson et al. 2009). Although this was difficult to test, it was considered unlikely, given the increasing and significant ontogenetic differences in the gape size of the fishes, plus their general functional morphological differences (De Silva et al., 1979). For example, the increased dietary specialisations apparent in *B. barbus* versus *L. leuciscus* were likely to be strongly driven by *B. barbus* having an inferior mouth that was primarily suited for only feeding on the benthos, with *L. leuciscus* having a terminal mouth and larger gape that enabled their exploitation of a
greater diversity of prey (e.g. by also exploiting drifting aerial insects). *Squalius cephalus* also has a terminal mouth that enabled their foraging throughout the water column, and they correspondingly had a very generalist diet and the largest niche of all the fishes at all sites. Given these results, there was no evidence to suggest the prolonged cohabitation of *B. barbus* with the other fishes in the study river had resulted in the competitive exclusion of a native species from its original niche (Bøhn et al. 2008). This is a contrast to invasive *B. barbus* in Italy where data suggest they have displaced endemic *Barbus* fishes in invaded river systems via competitive interactions, although dietary data on the fishes are currently absent (Carosi et al., 2017).

Across the 0+ fishes, trophic niche sizes and composition were most similar between *B. barbus* and *P. phoxinus*. The main driver of their trophic similarity was their high dietary proportions of Chironomid larvae. Given that *P. phoxinus* were the most abundant 0+ fish at each site, this suggests some potential for high inter-specific competition for resources with invasive *B. barbus* (Chase et al., 2016). However, both fishes had other items in their diet, suggesting that had intense competitive interactions resulted in reduced food intake rates, they could have switched to alternative prey (Dill, 1983). Moreover, with *P. phoxinus* the most numerically abundant 0+ fish at all sites and sampling occasions (their analysed sample sizes here of n = 20 per site and sampling occasions were derived via sub-sampling), there was no evidence to suggest their high dietary similarity with invasive 0+ *B. barbus* was having negative consequences at the population level, given their high abundance.

The diet composition of these invasive 0+ *B. barbus* in the River Teme was relatively similar to their diets in rivers in their indigenous range. For example, in the River Seig, Germany, larvae of Chironomids, caddisfly and mayfly were also all present in 0+ *B. barbus* diet
Similarly, in the River Trent, Eastern England, the diet of *B. barbus* in their late larval stages was also strongly dependent on Chironomid larvae (Nunn et al., 2007b). In the River Lee, England, Copp et al. (2005) also reported 0+ *B. barbus* predating upon similar items, including larvae of caddis fly and Chironomid larvae. Thus, there appears to be high similarity in *B. barbus* diet between their indigenous and non-indigenous ranges. When coupled with their diet similarities with the indigenous and highly abundant *P. phoxinus*, these results suggest some consistency with the pre-adaptation hypothesis of invasion biology. This hypothesis suggests that the probability of invasion by an introduced species is elevated when they share similar ecological traits and behaviours with indigenous species (Duncan & Williams, 2002). These similar traits and behaviours can include similar abilities to acquire resources (Duncan & Williams, 2002; Ricciardi & Mottiar, 2006). Invasion probability is also increased when the introduced species expresses their traits and behaviours in a similar manner to populations in their natural range (Duncan & Williams, 2002; Ricciardi & Mottiar, 2006; Buoro et al., 2016). The results here suggest that 0+ *B. barbus* underwent minimal shifts in their foraging behaviours to adapt to the River Teme, given their diet similarities to both their natural range and the other species in their new range. It is suggested that these factors assisted their establishment in, and invasion of, the River Teme.

There was a very low proportion of small-bodied (< 15 mm) and early larval stages in the 0+ fish samples. This was likely to have related to sampling bias resulting from the micromesh seine net, with it being inefficient to capture fishes of these lengths and life-stages (Cowx et al., 2001). If future studies require increased numbers of larval fishes in their analyses then an alternative sampling method would be required, such as point abundance sampling using electric fishing. This method can potentially sample larvae as small as 5 mm length (Copp, Bischoff & Freyhof, 1998).
Notwithstanding, at the free embryo stage and when they emerge from within spawning gravels, *B. barbus* larvae can be between 8 and 13 mm (Vilizzi & Copp, 2013). Thus, to capture early larval stages might require sampling methods capable of catching fish within the spawning gravels. Although the use of preservation of fish samples enabled enhanced dietary analyses in the laboratory, this can potentially result in shrinkage of body lengths (Fox, 1996). However, Leslie & Moore (2001) suggested shrinkage effects are relatively low when using similar preservation methods, providing samples are processed within a year of collection, as was completed here. Consequently, the relationships between diet and fish lengths in our study were considered valid. Finally, in our study, spatial comparisons were made in diet of each species, with differences between sites likely to have related to differences in food availability. However, the food availability of each site was not quantified accurately (given the presence of 37 items across the diets), preventing further analysis. Although these data on resource availability might also have assisted more precise testing of whether diets were generalist or specialist, assumptions on this were made from the feeding strategy plots (Amundsen et al. 1996). From these plots, all the fishes were described as generalists. However, across the four species, there was variation in the extent of this dietary generalism. *Barbus barbus* generally had the narrowest diet and smallest niche, and so they have also been described as being the species with the most specialist diet of the analysed fishes.

In summary, these results indicated how invasive 0+ *B. barbus* had successfully integrated into a 0+ cyprinid fish community via their diet and feeding ecology. The results highlighted that the 0+ *B. barbus* were consuming similar items to conspecifics in their indigenous range, suggesting some consistency with the pre-adaptation hypothesis of invasion biology. As the 0+ fishes all increased in their lengths and gape sizes, their diets became increasingly
dissimilar, especially between *B. barbus* and other fishes. This was primarily due to
differences in their functional morphology and resulted in the *B. barbus* niche sizes generally
being significantly smaller than the other fishes. This invaded fish community thus represents
a strong case study of how the invasion of a river system by a non-indigenous fish was
facilitated by the utilisation of their pre-adapted foraging behaviours.

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