RESEARCH ARTICLE

Macrophytes shape trophic niche variation among generalist fishes

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Abstract

Generalist species commonly have a fundamental role in ecosystems as they can integrate spatially distinct habitats and food-web compartments, as well as control the composition, abundance and behavior of organisms at different trophic levels. Generalist populations typically consist of specialized individuals, but the potential for and hence degree of individual niche variation can be largely determined by habitat complexity. We compared individual niche variation within three generalist fishes between two comparable lakes in the Czech Republic differing in macrophyte cover, i.e. macrophyte-rich Milada and macrophyte-poor Most. We tested the hypothesis that large individual niche variation among generalist fishes is facilitated by the presence of macrophytes, which provides niches and predation shelter for fish and their prey items. Based on results from stable nitrogen ($\delta^{15}$N) and carbon ($\delta^{13}$C) isotopic mixing models, perch ($Perca fluviatilis$ L.) and rudd ($Scardinius erythrophthalmus$ L.) showed larger individual variation (i.e., variance) in trophic position in Milada as compared to Most, whereas no significant between-lake differences were observed for roach ($Rutilus rutilus$ L.). Contrary to our hypothesis, all the three species showed significantly lower individual variation in the relative reliance on littoral food resources in Milada as compared to Most. Rudd relied significantly more whereas perch and roach relied less on littoral food resources in Milada than in Most, likely due to prevalent herbivory by rudd and prevalent zooplanktivory by perch and roach in the macrophyte-rich Milada as compared to macrophyte-poor Most. Our study demonstrates how the succession of macrophyte vegetation, via its effects on the physical and biological complexity of the littoral zone and on the availability of small prey fish and zooplankton, can strongly influence individual niche variation among generalist fishes with different ontogenetic trajectories, and hence the overall food-web structures in lake ecosystems.
Introduction

Generalist species that feed on multiple trophic levels (cf. [1]) commonly have a fundamental role in ecosystems. Generalists can, for instance, regulate the abundance, composition and niche use of organisms at lower and higher trophic levels and also integrate spatially distinct habitats and food-web compartments [2], thereby affecting the structure and stability of food webs [3,4]. Generalist populations with wide trophic niche typically consist of highly specialized individuals that only use a subset of the entire population niche [5]. Such individual specialization can, in turn, have complex population-, community- and ecosystem-level effects, including speciation processes, intra- and inter-specific interactions as well as nutrient cycles and energy flow ([6–8] and references therein). However, the occurrence and degree of individual niche variation, and hence potential for speciation and habitat coupling, within generalist populations is commonly controlled by niche availability and ecological opportunities [9–11].

In lakes, the niche availability and ecological opportunities are strongly controlled by the presence and extent of macrophyte vegetation, both through the direct effects on habitat complexity and the overall ecosystem productivity and also indirectly through reduced predatory effects (e.g. [12,13] and references therein). Besides affecting the abundance, size and community composition of zooplankton, benthic macroinvertebrates and small fish [12,14,15], macrophytes can also directly contribute to the diet of generalist fishes [13,16]. The effects of macrophyte vegetation on individual niche variation among generalist fishes can largely be shaped by the species’ foraging strategy and ontogenetic trajectories. However, there is limited empirical evidence of how macrophyte vegetation affects individual niche variation and hence the overall population-level niche width of generalist fishes. Here, we use a novel opportunity to compare how different succession stages (i.e., abundance) of macrophyte beds influence trophic niche width of three generalist fishes with contrasting foraging strategies. Perch (Perca fluviatilis L.), roach (Rutilus rutilus (L.)) and rudd (Scardinius erythrophthalmus (L.)) are all generalist and widely spread fishes [17], the former two species commonly dominating the fish communities in European lakes [18–20]. However, the three species show fundamental differences in their foraging strategies. Perch commonly undergo ontogenetic dietary shift from zooplankton to benthic invertebrates and finally to piscivory [21,22], whereas roach and particularly rudd become increasingly herbivorous with increasing size [17,23,24]. Hence, the species can be expected to show markedly different responses to the presence or absence of macrophytes, both in terms of the trophic position occupied by the populations and individuals within, as well as in terms of their use of littoral and pelagic food resources.

The competitive coexistence of perch and roach is one of the most extensively studied examples of intra- and inter-specific interactions among generalist fishes. A general succession from dominance of perch to roach with increasing productivity has been documented for lakes of different trophic states [18,25,26], as well as within reservoirs with pronounced longitudinal gradients [27,28]. Roach are efficient zooplanktivores, but may also feed on macroinvertebrates and non-animal food. Perch are ontogenetic carnivores and start their lives by feeding on zooplankton and later shift to macroinvertebrates to finally become piscivorous [21]. Both species mutually influence each other by way of asymmetrical competitive and predator-prey interactions. Roach are more efficient zooplanktivores in open water and thus force young-of-the-year perch to ingest benthic macroinvertebrates earlier in the season if zooplankton resources are reduced [29,30]. This leads to increased intra-specific competition between perch age classes and thereby to reduced growth and potential to a piscivorous niche shift [30]. In some lakes, perch and roach show marked niche segregation with perch feeding mainly on benthos among macrophyte vegetation and roach feeding on zooplankton in the pelagic areas [29]. However, habitat complexity can fundamentally affect intra- and inter-specific
interactions among perch and roach. Structured habitats offer high biomasses of macroinvertebrates [31,32] and thereby facilitate competitive dominance of the more efficient bentivorous perch [29,33]. Moreover, perch is also more efficient zooplanktivore in structured habitats as compared to roach [34]. Consequently, the presence of structurally complex habitats, such as littoral macrophytes, generally reduces the bottlenecks for ontogenetic niche shifts and increases the growth of perch, even if perch and roach are forced to coexist in these sheltered habitats due to the presence of piscivorous predators [32,35]. Although the perch-roach interactions are well studied, there is limited empirical evidence of how macrophyte vegetation can influence trophic niche width of generalist fishes, including also rudd.

We compared isotopic niche widths of generalist perch, roach and rudd between two deep, post-mining lakes in northern Czech Republic: Lake Milada with abundant macrophyte vegetation and Lake Most where macrophytes are practically absent. In Milada, macrophytes evidently increase the structural complexity of the littoral zone and hence provide niches and shelter against fish predation for various invertebrate taxa [13]. This high physical and biological complexity may hence support a more complex food-web structure in Milada as compared to the macrophyte-poor Most. However, the three generalist fishes likely utilize different trophic niches and may also respond differently to between-lake differences in niche availability. Therefore, we raised three main hypotheses: i) Firstly, due to their different foraging strategies and ontogenetic trajectories, we expected significant niche segregation between the three generalist species, with perch occupying the highest trophic position due to piscivory and rudd occupying the lowest trophic position and relying most heavily on littoral food resources due to herbivory. ii) Secondly, we expected that the abundant macrophyte vegetation in Milada would facilitate higher trophic variability within the generalist fish populations than in the macrophyte-poor Most, as individuals can utilize a wider selection of niches (i.e., prey items and habitats) in the former ecosystem. iii) Thirdly, we expected perch to be particularly responsive to higher niche availability since, unlike more invertivorous and herbivorous roach and rudd, perch can shift to piscivory and utilize small prey fish, which in turn are expected to be more abundant in the macrophyte-rich Milada.

Materials and methods

Study lakes

The study was conducted in two newly created comparable post-mining lakes in northern Czech Republic, Milada and Most (Fig 1), with a permission of the owner of the study sites, Palivový kombinát Ústí, státní podnik. The lakes originate from flooding of restored brown coal opencast mines. Aquatic restoration of Milada was performed in years 2001–2011 and nowadays several species of macrophytes and macroalgae are present in high biomasses to a depth of 12 m (average vegetation cover of > 65% at 0–12 m depth, consisting mainly of Potamogeton sp., Myriophyllum sp. and Chara sp.). Restoration of Most was performed in years 2008–2014, but no abundant macrophyte vegetation has been developed (average vegetation cover of < 1% at 0–6 m depth) (see [36] for more details). In years 2013–2014, the three study species composed in abundance 94–96% and 64–65% of the total catches in Milada and Most, respectively. The relative abundances of the species showed no significant between-lake differences (Wilcoxon matched pairs test: \( p = 0.9, Z_6 = 0.104; \text{S1 Table} \)), with perch and roach being consistently the most abundant and rudd being the least abundant species. Other fishes present in both lakes are ruffe (Gymnocephalus cernua (L.)), pike (Esox lucius L.), European catfish (Silurus glanis L.) and tench (Tinca tinca (L.)), whereas pikeperch (Sander lucioperca (L.)), asp (Leuciscus aspius (L.)) and Prussian carp (Carassius gibelio (Bloch)) are only found in Milada and maraena whitefish (Coregonus maraena (Bloch)) are only found in Most.
Fish sampling

Animal treatment was performed in accordance with the guidelines and permission from the Experimental Animal Welfare Commission under the Ministry of Agriculture of the Czech Republic (Ref. No. CZ 01679). The work was approved by the Ethics Committee of the Czech Academy of Sciences. All sampling procedures and experimental manipulations were approved by the Czech Academy of Sciences. The field study did not involve endangered or protected species.

Fish for stable isotope analysis (SIA) were sampled in September 2013 and 2014 by benthic (height 1.5 m, length 30 m) and pelagic (height 3 m, length 30 m) multi-mesh gillnets (12 mesh sizes ranging from 5 to 55 mm knot-to-knot, [37]) in both lakes. The gillnets were set overnight (2 h before sunset and lifted 2 h after sunrise) at depths of 0–3, 3–6, 6–9 and 9–12 m at two benthic and one pelagic localities in three repetitions for both types of gillnets in both lakes and years. Altogether 36 gillnets (24 benthic and 12 pelagic) were set in each lake and year, summing to a total of 144 gillnet nights and 8,640 m² gillnet area. All captured fish were immediately anaesthetized by a lethal dose of tricainemethanesulfonate (MS–222, Sigma Aldrich Co.), identified, measured, weighed and dissected for gut content analysis. Muscle tissues from randomly chosen perch, roach and rudd individuals were stored frozen at –20°C prior to final preparation for SIA.

Sampling of food sources for SIA

Littoral and pelagic basal resources (macrophytes and macroalgae, particulate organic matter, POM) and invertebrates (benthic macroinvertebrates, zooplankton) were sampled for SIA to study the overall food-web structure in both study lakes and to estimate the trophic position and the relative reliance of generalist fishes on littoral versus pelagic food resources using two-source isotopic mixing models (cf. [38,39]). Macrophytes, macroalgae, POM and benthic macroinvertebrates were collected in September 2013 and 2014, whereas zooplankton was sampled in September 2013 and in March, May, July, September and November 2014. Macrophytes, macroalgae and benthic macroinvertebrates were collected by two SCUBA divers, using plastic corer in case of macroinvertebrates. Pelagic POM samples were obtained by filtering 20 L of surface water (collected from the uppermost 5 m) through a 50-μm sieve to prevent
contamination by animal matter prior to the filtration under pressure through a glass microfiber filter (GF/C 1.2 μm; Whatham, U.S.A.). Zooplankton was sampled with a 100-μm mesh plankton net by several vertical hauls throughout the uppermost 20 m of the water column. All littoral and pelagic food sources were sorted, cleaned of detritus and other unwanted material, identified to genus or family level and stored frozen at -20°C prior to final preparation for SIA. Only soft body tissue was dissected from molluscs and trichopterans with cases.

Stable isotope analyses

All frozen fish, invertebrate and basal resource SIA samples were dried at 60°C for 48 h and ground into a homogenous powder using a ball-mill Retsch MM 200 (Retsch GmbH, Haan, Germany). Small subsamples (0.520–0.770 mg) were weighed into tin cups for the analysis of δ¹⁵N and δ¹³C. All SIA were conducted using a FlashEA 1112 elemental analyser coupled to a Finnigan DELTA*Plus Advantage mass spectrometer (Thermo Fisher Scientific Corporation, Waltham, MA, U.S.A.) at the University of Jyväskylä, Finland. Stable nitrogen and carbon ratios are expressed as delta values (δ¹⁵N and δ¹³C, respectively) relative to the international standards for nitrogen (atmospheric nitrogen) and carbon (Vienna PeeDeeBelemnite). Analytical precision was always better than 0.20‰ for both isotopes, and was based on standard deviation of repeated analysis of working standards (pike white muscle tissue and birch leaves) inserted in each run after every five samples. As C:N ratios were consistently lower than 3.5 (i.e., 96% of cases), obtained stable isotope values of fish were not lipid corrected [40].

In this study, SIA was applied to compare individual niche variation within the generalist fishes in macrophyte-rich Milada and macrophyte-poor Most. Stable nitrogen (δ¹⁵N) and carbon (δ¹³C) isotopes are widely used to study lake food webs and to estimate the long-term diet of consumers [39] as well as niche width of consumer populations [41,42]. While δ¹⁵N values indicate the trophic position of an organism in the food web, the δ¹³C values reflect the long-term, assimilated carbon sources of consumers, such as the relative proportions of littoral (benthic) versus pelagic (planktonic) food for fish [39]. Hence, SIA provides a powerful tool to study food-web complexity (e.g., food-chain length) and energy flow pathways in the ecosystems. Primary producers and consumers can show marked differences in δ¹⁵N and δ¹³C values between different sites [43,44]. Therefore, the δ¹⁵N and δ¹³C values of higher consumers, such as fish, need to be standardized with the isotopic baselines prior to comparisons across ecosystems. Here, the fish δ¹⁵N and δ¹³C values were standardized across the study lakes by estimating the trophic position (TP) and the relative reliance on littoral versus pelagic carbon sources (hereafter littoral reliance, LR), respectively, using the two-source isotopic mixing models described in [38]. The mean δ¹⁵N and δ¹³C values of macrophytes (i.e. Myriophyllum sp., Potamogeton sp., Elodea sp.) together with macroalgae (i.e. Chara sp.) and of pelagic POM were used as the littoral and pelagic isotopic baselines, respectively. Long-lived primary consumers, such as algae-grazing snails and filtrate feeding zooplankton or clams, are commonly used for estimating isotopic baselines since they often show lower spatial and temporal variation in δ¹⁵N and δ¹³C as compared to primary produces [38]. However, in the present study, it is more justified to use macrophytes and macroalgae as the littoral baseline since some of the study species (i.e., roach and particularly rudd) are facultative herbivores and thus can utilize these plant foods with the most elevated δ¹³C values. For the TP and LR computations, the commonly used trophic fractionation factors of 3.4‰ for δ¹⁵N and 0.4‰ for δ¹³C were applied [38].

Statistical analyses

Depending on normality of the data, either parametric t-tests or non-parametric Mann-Whitney U-tests were used to test between-lake differences in standard length, trophic position...
(TP) and littoral reliance (LR) of the generalist fishes. Analysis of variance (ANOVA) with Tukey’s HSD (honest significant difference) pairwise comparisons were used to test differences in TP and LR between the three species (i.e., inter-specific niche segregation within the lakes). Between-lake differences in the trophic niche width of the generalist species (measured as variance of TP and LR) were tested with Levene’s test for equality of variances, which takes into account differences in sample sizes and is robust to non-normality. All statistical analyses were done in R 3.1.1 [45].

**Results**

**Interspecific niche segregation**

The three generalist fishes showed significant niche segregation in terms of their trophic position both in the macrophyte-rich Milada (F<sub>2,423</sub> = 200.1, \( p < 0.001 \)) and in the macrophyte-poor Most (F<sub>2,336</sub> = 73.6, \( p < 0.001 \)). In both lakes, perch occupied on average the highest trophic position whereas rudd occupied the lowest trophic position (Table 1; Fig 2; \( p < 0.001 \) for all pairwise comparisons). The species also showed significant differences in their long-term use of littoral versus pelagic food resources in Milada (F<sub>2,423</sub> = 525.8, \( p < 0.001 \)), but not in Most (F<sub>2,336</sub> = 2.00, \( p = 0.137 \)). In Milada, rudd relied the most whereas perch relied the least on littoral food resources (Table 1; Fig 2; \( p < 0.001 \) for all pairwise comparisons).

**Between-lake differences in food webs and intraspecific niche variation**

Perch collected for SIA were on average smaller (W = 10,167; n = 326; \( p < 0.001 \)) whereas rudd were on average larger (W = 3,476; n = 134; \( p < 0.001 \)) in Milada as compared to Most. No significant between-lake differences were observed in roach size (S2 Table).

All the three generalist fishes, but perch in particular, occupied a significantly higher trophic position in the macrophyte-rich Milada than in the macrophyte-poor Most (Table 1). Rudd relied significantly more whereas perch and roach relied less on littoral food resources in Milada than in Most (Table 1), likely due to prevalent herbivory by rudd and prevalent zooplanktivory by perch and rudd in Milada as compared to Most.

The isotopic niche widths estimated for the three generalist fishes differed significantly between the macrophyte-rich Milada and the macrophyte-poor Most. Perch and rudd showed significantly higher individual variation (i.e., variance) in trophic position in Milada than in Most, whereas no between-lake differences were observed for roach (Table 2; Fig 2).

**Table 1. Mean (SD) and range of trophic position (TP) and littoral reliance (LR) estimates of the generalist fish populations in macrophyte-rich Milada and macrophyte-poor Most.**

|     | Milada |      |     |     | Most |      |     |     |     |
|-----|--------|------|-----|-----|------|------|-----|-----|-----|
|     | n      | Mean (SD) | Range | n      | Mean (SD) | Range | Statistic |     |
| TP  | Perch  | 167   | 4.64 (0.48) | 3.23–5.93 | 159 | 3.79 (0.33) | 3.02–4.49 | t = 18.93; df = 292.82; \( p < 0.001 \) |
|     | Roach  | 183   | 4.23 (0.32) | 3.15–5.14 | 122 | 3.65 (0.35) | 2.54–4.62 | t = 14.82; df = 244.19; \( p < 0.001 \) |
|     | Rudd   | 76    | 3.54 (0.39) | 2.65–4.20 | 58  | 3.18 (0.30) | 2.37–4.05 | W = 3,305; n = 134; \( p < 0.001 \) |
| LR  | Perch  | 167   | 0.39 (0.07) | 0.22–0.63 | 159 | 0.48 (0.17) | 0.10–0.86 | W = 8,209; n = 326; \( p < 0.001 \) |
|     | Roach  | 183   | 0.42 (0.07) | 0.28–0.79 | 122 | 0.53 (0.24) | 0.04–0.97 | W = 7,276; n = 305; \( p < 0.001 \) |
|     | Rudd   | 76    | 0.70 (0.08) | 0.47–0.86 | 58  | 0.49 (0.12) | 0.18–0.69 | W = 4,164; n = 134; \( p < 0.001 \) |

Sample sizes (n) and results from the statistical tests are shown, with significant between-lake differences (\( p < 0.05 \)) marked in bold.

https://doi.org/10.1371/journal.pone.0177114.t001
All the three species showed lower individual variation in the relative littoral reliance in macrophyte-rich Milada as compared to macrophyte-poor Most, with some individuals using mainly littoral benthic \((LR > 0.5)\) and some mainly pelagic planktonic food resources \((LR < 0.5)\) (Table 1; Fig 2). The variance tests were repeated after fixing the ranges in fish standard length between the lakes \((i.e.,)\) after excluding exceptionally small and large individuals but this did not change the results.

**Discussion**

Our study demonstrates how habitat complexity, in terms of macrophyte vegetation, can influence the degree of individual niche variation within and resource competition between coexisting generalist species, but also how generalists with different feeding strategies can respond differently to differences in niche availability. As hypothesized, the three generalist species showed a higher intra-specific niche variation and a more distinct inter-specific niche segregation in the presence of abundant macrophyte vegetation. In essence, the generally higher trophic position of generalists suggests a more complex food-web structure \((i.e.,)\) a longer food chain in the macrophyte-rich Milada as compared to the macrophyte-poor Most. Perch and roach showed higher individual variation in trophic position in Milada than in Most, apparently due to high trophic positions of some specialized piscivorous perch and low trophic positions of some specialized herbivorous roach. At the same time, roach with limited potential to piscivorous or herbivorous specialization showed minor response to between-lake differences in macrophyte vegetation. Contrary to trophic position, the generalists in Milada showed less individual variation in littoral versus pelagic resource use than the conspecifics in Most, possibly due to a general paucity of preferred food resources in the latter ecosystem. Overall, our study gives further insights into the fundamental role of macrophytes in shaping the structure and function of lake food webs via their effects on trophic niche variation among generalist fishes.

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**Table 2. Individual niche variation within the generalist fish populations in macrophyte-rich Milada and macrophyte-poor Most, measured as the variance in trophic position (TP) and littoral reliance (LR) estimates.**

|       | Variance |       |       |       |       |       |
|-------|----------|-------|-------|-------|-------|-------|
|       | n        | Milada| Most  | F, p  |       |       |
| TP    |          |       |       |       |       |       |
| Perch | 326      | 0.228 | 0.104 | 19.563; 0.001 |
| Roach | 305      | 0.104 | 0.122 | 152; 0.219 |
| Rudd  | 134      | 0.152 | 0.092 | 4.377; 0.038 |
| LR    |          |       |       |       |       |       |
| Perch | 326      | 0.0050| 0.0283| 80.323; 0.001 |
| Roach | 305      | 0.0048| 0.0576| 102.450; 0.001 |
| Rudd  | 134      | 0.0063| 0.0137| 13.892; 0.001 |

Results from the Levene’s test of equality of variances are shown, with significant between-lake differences \((p < 0.05)\) marked in bold.
The results from stable isotope analyses support our first hypothesis predicting significant long-term trophic niche segregation between the three generalist species. Perch, roach, and rudd are all generalist species, but they show markedly different ontogenetic trajectories, with perch shifting diet from zooplankton to benthic macroinvertebrates and finally to fish [46,47] and rudd commonly shifting diet from zooplankton to benthic macroinvertebrates and finally to macrophytes [23,36]. Roach typically prefer zooplankton diet throughout ontogenesis, but may shift to benthic macroinvertebrates and plant matter if the zooplankton resources are scarce or absent [48]. The present study supports this conception of niche segregation between the three generalists as perch generally occupied the highest whereas rudd occupied the lowest trophic position. Herbivory is particularly common among rudd and hence this foraging strategy likely reduces resource competition with the more zooplanktivorous and benthivorous roach [16,49]. In the present study, roach and rudd showed significant differences in relative littoral reliance only in Milada. There, the abundant macrophyte vegetation apparently provide a profitable food resource for herbivorous rudd, but also a predation shelter for large-sized zooplankton, which was one of the main prey items for roach (S3 Table), as observed elsewhere [49,50]. Hence, our results suggest that macrophyte vegetation can reduce niche overlap, and hence interspecific resource competition, between coexisting generalist species [33,51,52]. The three generalist species likely compete for zooplankton resources at early life-stages, but the niche overlap apparently decreases with the individuals' age and size due to increased piscivory by perch [21,33] and herbivory by rudd [53]. At the same time, piscivory by large perch generates novel intra- and inter-specific interactions as these individuals can predate upon smaller conspecifics as well as upon early life stages of other, potentially competitive fish species [21,33,46,54].

Macrophyte vegetation increases the overall physical complexity of the lake littoral zone, but also affects competitive and predatory interactions between different species and ontogenetic stages of fish [18,33,55]. Moreover, macrophytes can have direct effects on the lake total production via both increased primary production and increased density of benthic macroinvertebrates [15], large-sized zooplankton [12] as well as juvenile fish [56], but also indirectly via reducing resuspension of nutrients and fine particles from the sediment [13,57,58]. The effects of macrophytes on physical and biological complexity of the littoral zone can ultimately shape the overall structure and function of lake food webs [15,59]. While most studies have focused on macrophyte effects on competitive and/or predator–prey interactions [13,15,33], there has been limited empirical evidence of how macrophytes affect individual specialization among generalist species. The present study hence gives novel insights into how macrophyte vegetation can influence trophic niche variation within, as well as inter-specific resource competition between, generalist fishes with different foraging strategies. In particular, we found elevated and more variable trophic position of generalists in Milada, suggesting increased food-web complexity under the presence of macrophytes. This perception is supported by previous studies indicating that food-chain length generally increases with increasing resource availability [60,61], and it also supports our second hypothesis predicting higher intraspecific niche variation in the macrophyte-rich Milada. Our results also demonstrate clear between-lake differences in use of littoral and pelagic food resources, with generalists showing markedly larger individual variation in the macrophyte-poor Most. This high individual variation in littoral and pelagic resource use of generalists might indicate increased intra-specific competition due to a general paucity of food, forcing individuals to specialized foraging. This argument is consistent with the theory of trophic adaptability [62] and supported by previous studies from estuarine fishes where a scarcity of preferred food resources has led to population-level niche expansion [63]. This could explain why for instance some roach in Most have apparently specialized to zooplanktivory (LR < 0.5) and some other individuals have specialized to
herbivory ($LR > 0.5$) instead of utilizing generalist diet consisting largely of preferred, large-sized benthic macroinvertebrates and terrestrial insects.

Our study demonstrates the fundamental role of habitat complexity (i.e., the presence/absence of macrophyte vegetation) in shaping individual variation in and resource use of generalist fishes. Such individual- and population-level differences in trophic niches of generalists can have complex effects on the community composition, through top-down and bottom-up trophic cascades, as well as on the entire ecosystems through altered energy-flow pathways and habitat coupling [5,7,64,65]. For example, the partial population-level niche specialization by herbivorous rudd, zooplanktivorous roach and piscivorous perch suggests high predation pressure on lower trophic levels and also limited habitat coupling by the species in the macrophyte-rich Milada. However, besides habitat (spatial) complexity, the intra- and inter-specific interactions within lake food webs can be fundamentally shaped by seasonal (temporal) variation in prey availability and environmental conditions [66]. Moreover, the other coexisting fishes, including piscivorous pike, catfish, pikeperch and zooplanktivorous whitefish, may have unexplored impacts on the niche width and resource use of the studied generalist species. Hence, seasonal studies and more controlled mesocosm experiments could give valuable insights into how macrophytes shape individual niche variation as well as intra- and inter-specific interactions among generalist fishes. Telemetry studies, in conjunction with trophic studies, could also provide important information about how macrophyte vegetation affects habitat use and activity levels of different generalist species and their ontogenetic stages, as well as individuals with different personalities [67].

Supporting information

S1 Table. Relative abundances (%) of the generalist species caught with survey gillnets from macrophyte-rich Milada and macrophyte-poor Most in 2013 and 2014.
(PDF)

S2 Table. Means (SD) and ranges of standard length (mm) and wet mass (g) of perch, roach and rudd sampled from macrophyte-rich Milada and macrophyte-poor Most in 2013–2014.
(PDF)

S3 Table. Relative proportions of main prey items in the gut contents of perch, roach and rudd caught from macrophyte-rich Milada and macrophyte-poor Most in 2013–2014.
(PDF)

S4 Table. Data for PONE-D-16-38804.
(PDF)

Acknowledgments

We thank all FishEcU members for their help during gillnet sampling and Luboš Kočvara for help with preparation of stable isotope samples.

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References
1. Pimm SL, Lawton JH. On feeding on more than one trophic level. Nature 1978; 275: 542–544.
2. Polis GA, Anderson WB, Holt RD. Toward an integration of landscape and food web ecology: the dynamics of spatially subsidized food webs. Annu Rev Ecol Syst 1997; 28: 289–316.
3. Schindler DE, Scheuereell MD. Habitat coupling in lake ecosystems. Oikos 2002; 98: 177–189.
4. Kratina P, LeCraw RM, Ingram T, Arnhold BR. Stability and persistence of food webs with omnivory: is there a general pattern? Ecosphere 2012; 3: 1–18.
5. Bolnick DI, Svanbäck R, Fordyce JA, Yang LH, Davis JM, Hulsey CD, et al. The ecology of individuals: incidence and implications of individual specialization. Am Nat 2003; 161: 1–28. https://doi.org/10.1086/343878 PMID: 12650459
6. Araújo MS, Bolnick DI, Layman CA. The ecological causes of individual specialization. Ecol Lett 2011; 14: 948–956. https://doi.org/10.1111/j.1461-0248.2011.01662.x PMID: 21790933
7. Bolnick DI, Amarasekare P, Araújo MS, Bürger R, Levine JM, Novak M, et al. Why intraspecific trait variation matters in community ecology? Trends Ecol Evol 2011; 26: 183–192. https://doi.org/10.1016/j.tree.2011.01.009 PMID: 21367482
8. Mittelbach GG, Bailieu NG, Kjelvik MK. Fish behavioural types and their ecological consequences. Can J Fish Aquat Sci 2014; 71: 1–18.
9. Schluter D. The ecology and origin of species. Trends Ecol Evol 2001; 16: 372–380. PMID: 11403870
10. Svanbäck R, Persson L. Individual specialization, niche width and population dynamics: implications for trophic polymorphisms. J Anim Ecol 2004; 73: 973–982.
11. Losos JB. Adaptive radiation, ecological opportunity, and evolutionary determinism. Am Nat 2010; 175: 623–639. https://doi.org/10.1086/652433 PMID: 20412015
12. Schriver P, Bøggestrand J, Jeppesen E, Søndergaard M. Impact of submerged macrophytes on fish-zooplankton-phytoplankton interactions: large-scale enclosure experiments in a shallow eutrophic lake. Freshw Biol 1995; 33: 255–270.
13. Jeppesen E, Søndergaard M, Søndergaard M, Christoffersen K. The structuring role of submersed macrophytes in lakes. New York: Springer, 1992.
14. Newman RM. Herbivory and detritivory on freshwater macrophytes by invertebrates: a review. J N Am Benthol Soc 1991; 10: 89–114.
15. Warfe DM, Barmuta LA. Habitat structural complexity mediates food web dynamics in a freshwater macrophyte community. Oecologia 2006; 150: 141–154. https://doi.org/10.1007/s00442-005-0505-1 PMID: 16932971
16. Specziári A, Rezsu ET. Feeding guilds and food resource partitioning in a lake fish assemblage: an ontogenetic approach. J Fish Biol 2009; 75: 247–267. https://doi.org/10.1111/j.1095-8649.2009.02283.x PMID: 20738494
17. Kottelat M, Freyhof J. Handbook of European freshwater fishes. Berlin: Publications Kottelat, Cornol and Freyhof, 2007.
18. Persson L. Behavioral response to predators reverses the outcome of competition between prey species. Behav Ecol Sociobiol 1991; 28: 101–105.
19. Kubečka J. Succession of fish communities of Central and East European reservoirs. In: Comparative Reservoir Limnology and Water Quality Management (Eds. Straškraba M., Tundisi JS. & Duncan A.). Dordrecht: Kluwer Academic Publishers, 1993; pp. 159–168.
20. Blabolil P, Logez M, Ricard D, Prchalova M, Ríha M, Sagouis A, et al. An assessment of the ecological potential of Central and Western European reservoirs based on fish communities. Fish Res 2016; 173: 80–87.

21. Persson L. Asymmetries in competitive and predatory interactions in fish populations. In: Size-structured populations Ecology and Evolution (Eds. Ebenman B. & Persson L.), Heidelberg: Springer-Verlag, 1988; pp. 203–218.

22. Craig FG. Percid Fishes–Systematics, Ecology and Exploitation. Oxford: Blackwell Science, 2000.

23. Lodge DM. Herbivory on freshwater macrophytes. Aquat Bot 1990; 41: 195–224.

24. Kapuscinski KL, Farrell JM, Wilkinson MA. Feeding patterns and population structure of an invasive cyprinid, the rudd Scardinius erythrophthalmus (Cypriniformes, Cyprinidae), in Buffalo Harbor (Lake Erie) and the upper Niagara River. Hydrobiologia 2012; 693: 169–181.

25. Jeppesen E, Sandegaard M, Lauridsen T, Landkildehus F. Trophic structure, species richness and biodiversity in Danish lakes: Changes along a phosphorus gradient. Freshw Biol 2000; 45: 201–218.

26. Olin M, Rask M, Ruuhijärvi J, Kurkilaiti M, Ala-Opas P, Ylönen O. Fish community structure in mesotrophic and eutrophic lakes of southern Finland: the relative abundances of percids and cyprinids along a trophic gradient. J Fish Biol 2002; 60: 593–612.

27. Vašek M, Kubec̆ka J, Peterka J, Čech M, Drašťík V, Hládek M, et al. Longitudinal and vertical spatial gradients in the distribution of fish within a canyon-shaped reservoir. Int Rev Hydrobiol 2004; 89: 352–362.

28. Vašek M, Kubec̆ka J, Matě̆na J, Seda J. Distribution and diet of 0+ fish within a canyon-shaped European reservoir in late summer. Int Rev Hydrobiol 2006; 91: 178–194.

29. Persson L. Competition-induced switch in young of the year perch, Perca fluviatilis: an experimental test of resource limitation. Environ Biol Fish 1987; 19: 235–239.

30. Persson L, Greenberg LA. Juvenile competitive bottlenecks: the perch (Perca fluviatilis)-roach (Rutilus rutilus) interaction. Ecology 1990; 71: 44–56.

31. Diehl S. Fish predation and benthic community structure: the role of omnivory and habitat complexity. Ecology 1992; 73: 1646–1661.

32. Persson L. Predator-mediated competition in prey refuges: the importance of habitat dependent prey resources. Oikos 1993; 68: 12–22.

33. Diehl S. Foraging efficiency of three freshwater fishes: effects of structural complexity and light. Oikos 1988; 53: 207–214.

34. Winfield IJ. The influence of simulated aquatic macrophytes on the zooplankton consumption rate of juvenile roach, Rutilus rutilus, rudd, Scardinius erythrophthalmus, and perch, Perca fluviatilis. J Fish Biol 1986; 29: 37–48.

35. Persson L, Eklöv P. Prey refuges affecting interactions between piscivorous perch and juvenile perch and roach. Ecology 1995; 76: 70–81.

36. Vejříková I, Vejřík L, Syvärenta J, Kiljunen M, Čech M, Blabolil, et al. Distribution of Herbivorous Fish Is Frozen by Low Temperature. Sci Rep 2016; 6: 39600. https://doi.org/10.1038/srep39600 PMID: 28004804

37. CEN. Water Quality–Sampling of fish with multi-mesh gillnets. EN–14757. 2015.

38. Post DM. Using stable isotopes to estimate trophic position: models, methods, and assumptions. Ecology 2002; 83: 703–718.

39. Layman CA, Araujo MS, Boucek R, Hammerschlag-Peyer CM, Harrison E, Jud ZR, et al. Applying stable isotope to examine food-web structure: an overview of analytical tools. Biol Rev 2012; 87: 545–532. https://doi.org/10.1111/j.1469-185X.2011.00208.x PMID: 22051097

40. Post DM, Layman CA, Arrington DA, Takimoto G, Quattrochi J, Montana CG. Getting to the fat of the matter: models, methods and assumptions for dealing with lipids in stable isotope analyses. Oecologia 2007; 152: 179–189. https://doi.org/10.1007/s00442-006-0630-x PMID: 17225157

41. Bearhop S, Waldron S, Volier SC, Furness RW. Factors that influence assimilation rates and fractionation of nitrogen and carbon stable isotopes in avian blood and feathers. Physiol Biochem Zool 2002; 75: 451–8. https://doi.org/10.1086/342800 PMID: 12298464

42. Newsome SD, Martinez del Río C, Bearhop S, Phillips DL. A niche for isotopic ecology. Front Ecol Environ 2007; 5: 429–436.

43. France RL. Critical examination of stable isotope analysis as a means for tracing carbon pathways in stream ecosystems. Can J Fish Aquat Sci 1995; 52: 651–656.

44. Cabana G, Rasmussen JB. Comparison of aquatic food chains using nitrogen isotopes. P Natl Acad Sci USA 1996; 93: 10844–10847.
45. R Core Team. R: a language and environment for statistical computing. R Foundation for Statistical Computing. Vienna, Austria, 2014.

46. Persson L, Byström P, Wahlström E. Cannibalism and competition in Eurasian perch: population dynamics of an ontogenetic omnivore. Ecology 2000; 81: 1058–1071.

47. Vejčík L, Matějíčková I, Jůza T, Frouzová J, Seďa J, Blabolil P, et al. Small fish use the hypoxic pelagic zone as a refuge from predators. Freshw Biol 2016; 61: 899–913.

48. Persson L, Diehl S, Johansson L, Andersson G, Hamrin S. Shifts in fish communities along the productivity gradient of temperate lakes–patterns and the importance of size-structured interactions. J Fish Biol 1991; 38: 281–293.

49. Vašek M, Jarolím O, Čech M, Kubečka J, Peterka J, Prchalová M. The use of pelagic habitat by cyprinids in a deep riverine impoundment: Římov Reservoir, Czech Republic. Folia Zool 2008; 57: 324–336.

50. Jarolím O, Kubečka J, Čech M, Vašek M, Peterka J, Matána J. Sinusoidal swimming in fishes: the role of season, density of large zooplankton, fish length, time of the day, weather condition and solar radiation. Hydrobiologia 2010; 654: 253–265.

51. Horppila J, Nurminen L. Food niche segregation between two herbivorous cyprinid species in a turbid lake. J Fish Biol 2009; 75: 1230–1243. https://doi.org/10.1111/j.1095-8649.2009.02359.x PMID: 20738611

52. Estlander S, Nurminen L, Olin M, Vinni M, Immonen S, Rask M, et al. Diet shifts and food selection of perch Perca fluviatilis and roach Rutilus rutilus in humic lakes of varying water colour. J Fish Biol 2010; 77: 241–256. https://doi.org/10.1111/j.1095-8649.2010.02682.x PMID: 20646150

53. Prejs A, Jackowska H. Lake macrophytes as the food of roach (Rutilus rutilus) and Rudd (Scardinius erythrophthalmus L.). I. Species composition and dominance relations in the lake and food. Ekol Pol 1978; 26: 429–438.

54. Persson A, Hansson LA. Diet shift in fish following competitive release. Can J Fish Aquat Sci 1998; 56: 70–78.

55. Hargeby A, Blom H, Blindow I, Andersson G. Increased growth and recruitment of piscivorous perch, Perca fluviatilis, during a transient phase of expanding submerged vegetation in a shallow lake. Freshw Biol 2005; 50: 2053–2062.

56. Grenouillet G, Pont D. Juvenile fishes in macrophyte beds: influence of food resources, habitat structure and body size. J Fish Biol 2001; 59: 939–959.

57. Dennison WC, Orth RJ, Moore KA, Stevenson JC, Carter KS, Bergstrom PW, et al. Assessing water quality with submerged aquatic vegetation. Bioscience 1993; 43: 86–94.

58. Lacoul P, Freedman B. Environmental influences on aquatic plants in freshwater ecosystems. Environ Rev 2006; 14: 89–136.

59. Bode A, Alvarezo-Osorio MT, Varela M. Phytoplankton and macrophyte contributions to littoral food webs in the Galician upwelling estimated from stable isotopes. Mar Ecol Prog Ser 2006; 318: 89–102.

60. Post DM. The long and short of food-chain length. Trends Ecol Evol 2002; 17: 269–277.

61. Takimoto G, Post DM. Environmental determinants of food-chain length: a meta-analysis. Ecol Res 2013; 28: 675–681.

62. Gerking SD. Feeding ecology of fish. USA: Academic Press, 1994.

63. Feyrer F, Herbold B, Matern SA, Molye PB. Dietary shifts in a stressed fish assemblage: consequences of a bivalve invasion in the San Francisco Estuary. Environ Biol Fish 2003; 67: 277–288.

64. Quevedo M, Svanbäck R, Eklöv P. Intrapopulation niche partitioning in a generalist predator limits food web connectivity. Ecology 2009; 90: 2263–2274. PMID: 1973998

65. McMeans BC, McCain KS, Tunney TD, Fisk AT, Muir AM, Lester N, Shuter B, Rooney N. The adaptive capacity of lake food webs: from individuals to ecosystems. Ecol Monogr 2016; 86: 7–19.

66. Hayden B, Harrod C, Sonninen E, Kahilainen KK. Seasonal depletion of resources intensifies trophic interactions in subarctic freshwater fish communities. Freshw Biol 2015; 60: 1000–1015.

67. Castanheira MF, Herrera M, Costas B, Conceição LEC, Martins CIM. Can we predict personality in fish? Searching for consistency over time and across contexts. PLOS ONE; 8: e62037. https://doi.org/10.1371/journal.pone.0062037 PMID: 23614007

68. Jackson AL, Inger R, Parnell AC, Bearhop S. Comparing isotopic niche widths among and within communities: SIBER–Stable Isotope Bayesian Ellipses in R. J Anim Ecol 2011; 80: 595–602. https://doi.org/10.1111/j.1365-2656.2011.01806.x PMID: 21401589