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7	Stable isotopes and gut contents indicate differential resource use by
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24 Abstract

25 Differential use of habitat and prey resources is an important mechanism that may allow 26 coexistence of sympatric species. Unlike interactions between smaller cyprinid and percid 27 fishes, the resource use by coexisting predatory asp (Leuciscus aspius) and pikeperch (Sander 28 *lucioperca*) is relatively unknown. Here, gut content and stable isotope analyses were used to 29 study ontogenetic dietary shifts and interspecific trophic niche overlap between asp and 30 pikeperch coexisting in two reservoirs. The hypothesis that both species show an ontogenetic 31 dietary shift from small invertebrates to large fish prey, but at the same time use different prey 32 resources to reduce potential competitive interactions, was validated. The isotopic niches of 33 the two predators showed no, or only a moderate, degree of overlap (0-65%). The ontogenetic 34 changes in the degree of interspecific isotopic niche overlap were different in the two 35 reservoirs, suggesting that trophic segregation can be dynamic and variable among systems. 36 Gut contents revealed that small (<100 mm standard length) asp consumed mostly terrestrial 37 invertebrates and emerged aquatic insects, whereas small pikeperch foraged on zooplankton, 38 larval and pupal stages of aquatic insects and fish. Larger individuals (>100 mm) of both 39 species were predominantly piscivorous, with asp consuming more cyprinid prey and 40 pikeperch more percid prey. Coexisting asp and pikeperch populations are able to utilise 41 different prey resources, thereby reducing potential negative competitive interactions.

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Keywords: dietary ontogeny, foraging strategy, interspecific competition, piscivory, stableisotopes

46 1 Introduction

47 Differential resource use is perceived as an important mechanism allowing the coexistence of 48 species within ecological communities (Schoener, 1986; Chesson, 2000; Chase & Leibold, 49 2003). This view is based on the competitive exclusion principle (Hardin, 1960), which states that species cannot stably coexist unless the utilisation of limiting resources is well 50 51 differentiated. The segregation of coexisting species can occur along various dimensions such 52 as the time of activity, the habitat used, or the type of prey eaten (Schoener, 1986). A 53 comprehensive review of resource use in fish communities by Ross (1986) suggested that 54 niche segregation among coexisting species is mainly driven by partitioning of available food 55 resources rather than habitat or time segregation. Species coexistence can, however, be 56 influenced also by other mechanisms. For example, stochastic events (e.g., unpredictable 57 environmental fluctuations) that affect demographic attributes of species may result in their 58 coexistence (Sale, 1978; Warner & Chesson, 1985; Grossman et al., 1998). Therefore, one 59 approach to improve our understanding of the ecological mechanisms that determine the 60 coexistence of species is to examine resource overlap among potential competitors. Most 61 studies of dietary segregation between coexisting fish species have focused on adult life 62 stages (e.g., Hodgson et al., 1997; Schulze et al., 2012; Walker et al., 2013; Zaia Alves et al., 63 2017). Ontogenetic variations in the resource use among potential competitors have been 64 examined less frequently (Werner & Gilliam, 1984; Amundsen et al., 2003; Davis et al., 2012) although this knowledge is important to fully understand the structure and functioning 65 66 of fish communities.

Asp (Cyprinidae, *Leuciscus aspius*) and pikeperch (Percidae, *Sander lucioperca*) are important piscivorous fishes in freshwater communities of western Eurasia where they naturally coexist in large rivers, lakes and reservoirs (Kottelat & Freyhof, 2007; Vašek et al., 2013). Adults reach similar sizes (usually up to 1000 mm in total length) and prey on small

71 fish (Baruš & Oliva, 1995; Mittelbach & Persson, 1998). Juveniles of pikeperch forage on 72 aquatic invertebrates whereas juveniles of asp may also feed on terrestrial insects fallen on the water surface (Baruš & Oliva, 1995). The feeding ecology of pikeperch has been explored 73 74 extensively and thus it is well-known that this species usually shifts to piscivory in the first summer of its life (Buijse & Houthuijzen, 1992; van Densen et al., 1996; Mittelbach & 75 76 Persson, 1998). Less is known, however, about the size and age at which asp become 77 piscivorous. Moreover, only limited attempts have been made to quantitatively characterise 78 the diets of coexisting asp and pikeperch populations (Specziár & Rezsu, 2009). In general, 79 similar feeding habits (i.e., invertivory followed by piscivory) suggest that the two species 80 may interact strongly. Sympatric populations of asp and pikeperch thus provide a good 81 opportunity to investigate whether and how the two predators differ in resource use 82 throughout their lives.

83 In this study, gut content (GCA) and stable isotope (SIA) analyses were used to 84 explore ontogenetic dietary shifts and niche segregation between asp and pikeperch co-85 occurring in two artificial lakes (i.e., reservoirs). It was expected that both species undergo an 86 ontogenetic dietary shift from invertebrates to fish prey, but this shift occurs later (i.e., at a 87 larger body size) for asp due to its higher tendency to feed on invertebrates. It was also 88 hypothesised that coexisting asp and pikeperch use different prey resources, but the degree of 89 trophic segregation diminishes with increasing body size, i.e. when both species become 90 piscivorous.

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93 2 Methods

94 2.1 Study sites

95 The study was carried out in two reservoirs located in South Bohemia, Czech Republic. Lipno 96 Reservoir (hereafter Lipno; 48°37'58"N, 14°14'13"E), situated on the upper Vltava River, is 97 a relatively shallow water body (Table 1). Due to its shallowness and frequent wind action, 98 most of the reservoir area does not thermally stratify during the summer season. In contrast, 99 Římov Reservoir (hereafter Římov; 48°51'00"N, 14°29'28"E), situated on the Malše River, is 100 a deep canyon-type lake (Table 1) that is strongly thermally stratified during the summer 101 season. Both reservoirs have similar water clarity and a moderately eutrophic trophic status 102 (Table 1).

103 Due to seasonal water level fluctuations, the littoral zone vegetation is poorly 104 developed and submerged macrophytes are practically missing in both reservoirs. The adult 105 fish community compositions are similar in Lipno and Římov, with a dominance of cyprinid 106 species (mostly roach Rutilus rutilus, bleak Alburnus alburnus, bream Abramis brama and 107 white bream Blicca bjoerkna) accompanied by perch Perca fluviatilis and ruffe 108 Gymnocephalus cernua (Čech et al., 2009; Vašek et al., 2016). Asp and pikeperch naturally 109 reproduce in both reservoirs (Jůza et al., 2013; Blabolil et al., 2016). In Římov, however, 110 populations of the two predators are also regularly supported by stocking with pond-reared 111 fingerlings in autumn (Vašek et al., 2013).

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113 2.2 Sample collection

Fish sampling and treatment was conducted in compliance with guidelines from the Experimental Animal Welfare Commission under the Ministry of Agriculture of the Czech Republic. Asp, pikeperch and their fish prey were sampled from Lipno in August/September 2012 and 2013, and from Římov in August 2013 and 2014. Sampling was carried out with multi-mesh survey gillnets set overnight in littoral, profundal and pelagic zones at four to five different stations within each reservoir (for details of the gillnet sampling, see Vašek et al.,

2016). Additional samples of young-of-the-year (YOY) asp and pikeperch, as well as prey
fish, were collected from the littoral and pelagic zones of both reservoirs using a beach seine
net and a trawl, respectively (for details of these sampling methods, see Jůza et al., 2014).

123 Each fish was measured for standard length (mm) and a sample of dorsal muscle was 124 dissected and stored at -20° C until processed for stable isotope analysis. The analysed prey 125 fish included YOY perch, ruffe and roach, and one-year old bleak. The digestive tracts of asp 126 and pikeperch were dissected and preserved in a 10% formaldehyde solution for later diet 127 analysis. Scales and otoliths were taken and used for age determination following validated methods described by Ruuhijärvi et al. (1996) and Krpo-Ćetković et al. (2010). To evaluate 128 129 ontogenetic changes in the short-term diets (based on GCA that represents the recently 130 ingested prey items) and long-term diets (based on SIA that represents the assimilated food 131 sources over several weeks to months) of asp and pikeperch, individuals of both species were 132 grouped into <100, 100–199, 200–299, and ≥300 mm size classes that corresponded approximately to age categories 0+, 1+, 2+, and $\geq 3+$, respectively. 133

134 At both reservoirs, invertebrate samples for SIA were collected from three to four 135 sampling stations and three to four times (June to August) during the same summers when the 136 fish were captured. Bulk zooplankton was collected from the pelagic zone by taking several 137 vertical hauls through the upper 5 m (Lipno) or 10 m (Římov) of the water column with a 138 200-µm mesh plankton net. The live zooplankton was immediately sieved through a 350-µm 139 mesh and stored frozen at -20° C. Before preparation for SIA, defrosted samples were visually 140 inspected using a stereomicroscope. Most of the samples were dominated by herbivorous 141 crustaceans, mainly cladocerans (Daphnia) and calanoid copepods. Bulk samples of 142 macroinvertebrates from the littoral zone (<2 m depth) were collected with a kick net (mesh 143 size 0.7 mm) and hand-picked from stones. Only non-predatory organisms (primary 144 consumers) were considered and they included mainly trichopteran, ephemeropteran and

145 chironomid larvae, and also waterlouse (Asellus aquaticus) and small snails (Lymnaeidae). 146 All the trichopteran larvae and snails were removed from their cases or shells. Bulk samples 147 of terrestrial insects were collected with a sweep net from the shoreline grasses and shrubs. 148 Adults of the aquatic insects (Trichoptera, Ephemeroptera, Odonata and Diptera) were 149 excluded when present, and thus the samples contained adult insects of purely terrestrial 150 origin, i.e. Hymenoptera, Hemiptera, Coleoptera, Lepidoptera, Brachycera and Orthoptera. 151 All littoral and terrestrial invertebrate samples were stored frozen at -20°C until further 152 processed for SIA.

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154 2.3 Gut content analysis

155 In the laboratory, digestive tracts were opened and the contents were examined under a 156 stereomicroscope. Since asp lack a true stomach, the contents of the entire gut from the 157 oesophagus to the anus were analysed, whereas only stomach contents were analysed from 158 pikeperch. The total gut or stomach fullness was first visually estimated on a percentage scale 159 ranging from empty (0%) to full (100%). The prey items were identified to the lowest feasible 160 taxonomic level, and their contribution to the total gut or stomach fullness was then 161 determined by the indirect volumetric method (Hyslop 1980). In addition, the number of prey 162 fish individuals discernible to species level was recorded for each digestive tract. When 163 possible, characteristic remains (e.g., scales, pharyngeal arches, opercula and other bones of fish prey, and head capsules, thoraxes, tail spines and other exoskeleton parts of invertebrate 164 165 prey) were used for identification of masticated and partially digested prey items. The prey 166 items were subsequently grouped into six categories: (1) crustacean zooplankton, (2) larval 167 and pupal stages of aquatic insects, (3) emerged aquatic insects, (4) terrestrial insects, (5) 168 cyprinid fish, and (6) percid fish.

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170 2.4 Stable isotope analysis

171 Fish muscle and invertebrate samples were dried at 60°C for 48 h and ground to a fine powder using either a porcelain mortar or a mixer mill MM 200 (Retsch GmbH, Haan, Germany). 172 173 Stable carbon and nitrogen isotopes and the element (C, N) composition of all samples were measured using a Europa Scientific elemental analyser interfaced with a Europa Scientific 20-174 175 20 isotope ratio mass spectrometer (Sercon Ltd, Crewe, UK) at the Iso-Analytical Ltd, Crewe, 176 UK. Vienna Pee Dee Belemnite and atmospheric N₂ were used as the international standards 177 for carbon and nitrogen, respectively, while NBS-1577B (powdered bovine liver, $\delta^{13}C_{V-PDB} =$ -21.60%, $\delta^{15}N_{Air} = 7.65\%$) was used as a working standard. NBS-1557B was calibrated in-178 179 house as a secondary reference material and is directly traceable to IAEA-CH-6 (sucrose, $\delta^{13}C_{V-PDB} = -10.43\%$) and IAEA-N-1 (ammonium sulphate, $\delta^{15}N_{Air} = 0.40\%$). Isotope ratios 180 in each sample were expressed in conventional delta notation ($\delta^{13}C$, $\delta^{15}N$) as parts per 181 182 thousand (%) differences from the international standard. The analytical error (standard deviation), estimated from replicated runs of the reference material, was less than 0.1% for 183 both δ^{13} C and δ^{15} N. Every fifth sample was run in duplicate and the mean difference ± 184 185 standard deviation (SD) between replicates was $0.03 \pm 0.04\%$ for δ^{13} C and $0.06 \pm 0.06\%$ for δ^{15} N. The fish muscle δ^{13} C values were not corrected for lipids due to the generally low C:N 186 187 ratios (<3.5) indicating negligible lipid content in the samples (Hoffman et al., 2015).

The relative contributions of different diet sources assimilated by each size class of asp and pikeperch were modelled using the SIAR package in R (Stable Isotope Analysis in R; Parnell et al., 2010). Inputs to the model were the δ^{13} C and δ^{15} N values of the individual consumers (asp and pikeperch) and the reservoir-specific mean \pm SD δ^{13} C and δ^{15} N values of the potential prey resources (Figure A1). In both reservoirs, pelagic zooplankton and littoral macroinvertebrates did not differ in isotope values (*t*-tests, *P* > 0.05 for both δ^{13} C and δ^{15} N) and hence they were grouped as "aquatic invertebrates" for SIAR. Furthermore, the isotope

195 data collected in the subsequent years were pooled because isotope values for major trophic 196 level groups (i.e. predatory fish, prev fish, aquatic invertebrates and terrestrial insects) did not 197 substantially differ between years. Therefore, diet composition for both asp and pikeperch 198 was estimated from three possible diet sources: aquatic invertebrates, terrestrial insects, and 199 fish. Fractionation factors (mean \pm SD) between resources and the consumers were assumed to be 0.91 \pm 1.04% for δ^{13} C and 3.23 \pm 0.41% for δ^{15} N (Vander Zanden & Rasmussen, 200 201 2001). Element concentrations (proportions of C and N) directly measured in the prey 202 resources were included into the model (Phillips & Koch, 2002).

203 Trophic position (TP) of individual asp and pikeperch was estimated from stable
204 isotope data, using the equation described by Cabana and Rasmussen (1996):

205
$$TP_{consumer} = (\delta^{15}N_{consumer} - \delta^{15}N_{baseline}) / 3.23 + 2$$

where $\delta^{15}N_{consumer}$ is the $\delta^{15}N$ value of asp or pikeperch, $\delta^{15}N_{baseline}$ is the $\delta^{15}N$ value of the baseline organisms (calculated as the average value from aquatic invertebrates), 3.23 is the assumed diet-tissue enrichment in $\delta^{15}N$ per trophic level (Vander Zanden & Rasmussen, 2001), and the constant 2 refers to the trophic position of the baseline organisms.

Finally, the isotopic niche widths of each size class of asp and pikeperch were calculated as sample size-corrected standard ellipse areas (SEA_C) using the SIBER package in R (Stable Isotope Bayesian Ellipses in R; Jackson et al., 2011). SEA_C was also used to determine the degree of isotopic niche overlap between the two species, using the equation of Stasko et al. (2015):

215 % Overlap = [{(area of overlap between SEA_{C1} and SEA_{C2}) x 2} / (SEA_{C1} + SEA_{C2})] x 100

where SEA_{C1} and SEA_{C2} are the ellipse areas calculated from asp and pikeperch samples,
respectively.

219 2.5 Statistical analysis

220 Non-parametric one-way analysis of similarities (ANOSIM) was run in PAST ver. 3.19 221 (Hammer et al., 2001) to compare volumetric proportions of different prey categories in the 222 digestive tracts of different size classes of asp and pikeperch in the Lipno and Římov 223 reservoirs. ANOSIM was based on Bray-Curtis similarity index and the one-tailed 224 significance was computed by permutation of group membership with 9,999 replicates. The 225 size at piscivorous shift was compared between the species by using binomial data of prey 226 fish presence in gut contents (0 = no fish remains in gut, 1 = fish remains in gut) as the 227 response variable and fish length and species as the predictor variables in logit-regression 228 models. Furthermore, the ontogenetic (i.e., size-related) changes in asp and pikeperch TP 229 were analysed by fitting asymptotic regression models using the SSasymp function in R (Ritz 230 et al., 2015). The differences in TP between asp and pikeperch of each size class in each 231 reservoir were also compared using *t*-test. Finally, the likelihood test in the SIBER (Jackson et 232 al., 2011) was used to test for between-species differences in isotopic niche widths of asp and 233 pikeperch size-classes. All statistical analyses except ANOSIM were performed in the R 234 computing programme ver. 3.4.1 (R Core Team, 2017).

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237 3 Results

Both GCA and SIA data demonstrated clear ontogenetic dietary shifts and differential use of the prey resources by coexisting asp and pikeperch. The GCA results indicated significant between-species differences in the prey compositions (ANOSIM: R = 0.457, P < 0.001), but the diets of asp and pikeperch became more similar with increasing size (Table 2). Small (<100 mm) asp fed on terrestrial and emerged aquatic insects, whereas larger asp consumed mostly fish (Fig. 1). Correspondingly, small (<100 mm) pikeperch foraged on zooplankton,

larval and pupal stages of aquatic insects and fish, whereas larger pikeperch were mainly 244 245 piscivores (Fig. 1). Contrary to asp, no terrestrial insects or emerged aquatic insects were 246 found in pikeperch stomachs. The two species showed contrasting prey fish compositions, 247 with asp feeding more often on cyprinid prey fish and pikeperch feeding mostly on percid fish 248 (Fig. 1). When only prey fish discernible to species level were considered, the most abundant 249 species found in asp guts were ruffe in Lipno and bleak in Římov, whereas the most abundant 250 species observed in pikeperch stomachs were perch and conspecifics in Lipno and ruffe in 251 Římov (Table 3). The logit-regression models (Fig. 2) demonstrated that pikeperch shifted to 252 piscivory at a smaller size than asp, both in Lipno (parameter estimate \pm SE for species effect: 253 2.4 ± 0.9 ; Z = 2.6, P = 0.009) and Římov (2.3 ± 0.6; Z = 3.7, P < 0.001).

The SIA results confirmed the ontogenetic dietary shifts of asp and pikeperch to 254 255 piscivory, as illustrated by the positive non-linear relationship between size and TP (Fig. 3, 256 Table 4) and by the SIAR estimates showing a shift from invertebrate to fish prey with 257 increasing predator size (Fig. 4). In Lipno, small (<100 mm) asp had a significantly lower TP 258 than similar-sized pikeperch, whereas no between-species differences were observed among 259 larger size classes (Table 5). An opposite pattern was observed in Římov, where no between-260 species differences in TP were observed for small asp and pikeperch, whereas larger asp had 261 consistently lower TP as compared to similar-sized pikeperch (Table 5). The results from 262 SIAR isotopic mixing model suggested that terrestrial insects contributed only little to the 263 long-term diet of all size classes of asp and pikeperch in Římov but had a relatively high 264 contribution to the long-term diet of both small (<100 mm) asp (54%) and pikeperch (32%) in 265 Lipno (Fig. 4).

The isotopic niche widths generally did not differ between the coexisting asp and pikeperch populations (Table 5). In Lipno, there was no overlap between isotopic niches (SEA_c) of small (<100 mm) asp and pikeperch, whereas the isotopic niches of larger asp and pikeperch overlapped partially (i.e., 49–65%; Table 5). An opposite pattern was observed in \tilde{R} ímov, where the interspecific SEA_C overlap was highest (65%) in the <100 mm size class, intermediate (42%) in the 100–199 mm size class, and none in the 200–299 and \geq 300 mm size classes (Table 5).

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275 4 Discussion

276 Both GCA and SIA suggested that there were clear ontogenetic dietary shifts and interspecific 277 niche segregation between asp and pikeperch in the two reservoirs examined. So far, only 278 limited information on resource use has been available for co-occurring asp and pikeperch 279 populations (Specziár & Rezsu, 2009), and the current study is also the first that applied SIA 280 approach (together with conventional GCA method) to characterise dietary niches of the two 281 sympatric predators. Consequently, findings of this study provide important insights into the 282 trophic ecology of coexisting asp and pikeperch populations and their roles in freshwater food 283 webs.

284

285 4.1 Ontogenetic dietary shifts in asp and pikeperch

286 Gut content and stable isotope data both showed that asp and pikeperch consumed more fish 287 prey with increasing body size. According to the GCA and SIAR results, fish prey 288 overwhelmingly dominated in the short- and long-term diets of large- and medium-sized 289 predators, whereas they contributed only around 50% or less to the diets of small-sized (<100 290 mm) asp and pikeperch. Our results demonstrate that asp and pikeperch can begin feeding on 291 fish as early as their first summer (i.e., as YOY), although shifting to piscivory was completed 292 in their second summer of life (i.e., at length >100 mm). Both the logit-regression models 293 (based on absence/presence of prey fish in predators' digestive tracts) and SIA-based TP

294 estimates consistently indicated that pikeperch shifted to piscivory at a smaller size than asp. These results confirmed the expectation that juvenile asp have a higher tendency to feed on 295 296 invertebrates and shift to piscivory somewhat later (i.e., at a larger size) than pikeperch. 297 However, although piscivory occurred later for asp, the TP estimates suggest that both species 298 accomplished shifting to predominantly piscivorous feeding in their second summer of life 299 since individuals of the 100–199 mm size class attained mean TP values of \geq 3.5, indicating 300 piscivory. Consequently, both species can be characterised as "specialist piscivores" (sensu 301 Keast, 1985) because they shift to piscivory relatively early in life.

302 GCA indicated that small-sized (<100 mm) asp consumed mainly terrestrial and 303 emerged aquatic insects in both reservoirs. SIAR results suggested that terrestrial insects 304 dominated (54%) the long-term diet of small-sized (<100 mm) asp in Lipno, whereas small 305 asp in Římov showed a greater reliance upon prey fish. Hence, the SIAR results indicate that 306 small-sized Římov asp probably consumed more fish than suggested by GCA which reflects only recently ingested prey items (e.g., Paradis et al., 2008). According to GCA, small-sized 307 308 (<100 mm) pikeperch fed on zooplankton, aquatic insects and fish, but not on terrestrial and 309 emerged aquatic insects. Although terrestrial insects were absent in pikeperch digestive tracts, 310 we always included terrestrial invertebrates as a third prey source (besides aquatic 311 invertebrates and fish) to make the SIAR analysis consistent between asp and pikeperch. 312 Consequently, the SIAR results confirmed that terrestrial insects generally represented an 313 unimportant prey source for pikeperch of all size classes. Small-sized (<100 mm) Lipno 314 pikeperch were, however, an exception because the SIAR suggested that terrestrial insects 315 might be a substantial diet source (32%) for these fish. This apparent bias (i.e., overestimation 316 of the terrestrial prey contribution to pikeperch diet) can be explained by the fact that stable 317 isotope values of terrestrial and aquatic invertebrates partially overlapped (particularly in 318 terms of δ^{13} C) in Lipno, hindering assessment of the relative contributions of these prev sources to higher trophic levels. Nevertheless, the SIAR results appropriately revealed theontogenetic niche shift from feeding on invertebrates to piscivory in both species.

321 Previous studies have shown that, under favourable growth conditions (i.e., higher 322 optimum temperature and food availability), pikeperch become piscivorous during their first 323 summer and reach sizes well above 100 mm (Buijse & Houthuijzen, 1992; van Densen et al., 324 1996). In contrast, under less suitable conditions, YOY pikeperch either remain invertivorous 325 and reach generally small sizes (Specziár, 2005; Vinni et al., 2009, Ginter et al., 2011) or 326 develop a bimodal size distribution with a minor group becoming piscivorous and a majority 327 staying invertivorous (van Densen, 1985; Frankiewicz et al., 1996). Information on 328 ontogenetic dietary shifts in asp is limited. Yet, the data available from Lake Balaton 329 (Specziár & Rezsu, 2009) correspond well with the current study: the <40 mm asp were 330 invertivorous, the 41-120 mm asp had a diet containing both invertebrates and fish, and the 331 121–500 mm fish were entirely piscivorous. The fact that transition to piscivory in our study 332 systems was not completed during the first summer implies growth-limiting conditions for 333 juvenile stages of both species. Persson & Brönmark (2002) highlighted the importance for 334 YOY predators to be synchronised with fluctuations in resource availability. Hence, we can 335 speculate that discontinuous availability of suitable food resources might reduce growth and 336 delay shifting to piscivory in our study systems. However, growth rates of YOY predators 337 might have also been restricted by water temperatures. Římov is a deep reservoir situated in a 338 canyon and hence it warms slowly in spring which may delay the spawning period and 339 shorten the first-year growth season (cf., Wysujack et al., 2002; Jůza et al., 2013). In contrast, 340 Lipno is a shallow reservoir, but because of its location at a higher altitude, thermal 341 conditions may delay spawning and reduce the growth of juvenile stages similarly to that in 342 Římov.

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344 4.2 Interspecific niche segregation between asp and pikeperch

345 The isotopic niche widths generally did not differ between the coexisting asp and pikeperch 346 populations, indicating a similar extent of trophic specialisation in both species. Moreover, 347 the isotopic niches of the two predators showed no or only a moderate degree of overlap (i.e., 348 0-65%). Interestingly, overlap between the isotopic niches of asp and pikeperch increased 349 with increasing predator size in Lipno but decreased in Římov. These findings do not support 350 the hypothesis that the degree of trophic segregation between asp and pikeperch should 351 diminish with increasing body size (i.e., with a shift to piscivory). Instead, the results suggest 352 that size-related trophic segregation between asp and pikeperch may be dynamic and variable 353 among systems, probably reflecting varying availability of prey sources.

354 The results also demonstrate that coexisting asp and pikeperch used rather different 355 prey resources both at small and large sizes. The GCA and SIA data both indicated that the 356 interspecific trophic segregation in the smallest (<100 mm) predator size class was likely due to the exclusive utilisation of terrestrial invertebrates and emerged aquatic insects by asp, 357 358 whereas pikeperch used zooplankton, larval and pupal stages of aquatic insects, and small 359 fish. Similarly, Specziár & Rezsu (2009) observed that small (16-40 mm) asp foraged mostly 360 on adult Chironomidae, whereas co-occurring similar-sized pikeperch relied on zooplankton. 361 Moreover, the GCA results showed that piscivorous stages of the two predators consumed the 362 same fish species, but in different proportions. The fact that asp used relatively more cyprinid 363 prey fish while pikeperch consumed more percid fish might be another reason for the 364 observed segregation of the isotopic niches of the two predators, particularly those of 365 medium- and large-sized classes. Because pikeperch ingest prey intact, with no mastication, it 366 was usually possible to identify (at least to family level) most of prey fish. In contrast, 367 ingested prey fish in asp were often strongly masticated by pharyngeal teeth and digested 368 beyond recognition. Hence, we suppose that the relative contribution of cyprinid prey fish in the diet of asp might even be higher than suggested by the GCA, because small and softcyprinid species such as bleak were probably under-represented due to their rapid digestion.

371 This study provides novel empirical data on piscivorous diets of coexisting asp and 372 pikeperch populations. Previous single-species studies indicated that the piscivorous diets of 373 both asp and pikeperch are dominated by cyprinid (Wysujack et al., 2002; Krpo-Ćetković et 374 al., 2010; Specziár, 2011) and by percid prey fish (Vostradovský & Váša, 1981; Frankiewicz 375 et al., 1999, Keskinen & Marjomäki, 2004). Hence, both predators can behave rather 376 opportunistically and consume the most abundant fish species. However, in sympatry, asp and 377 pikeperch can differentiate prey fish resources as illustrated by this study. In summary, by 378 using a combination of GCA and SIA, our study indicates that coexisting asp and pikeperch 379 populations can use different prey resources at both juvenile and adult life stages, thereby 380 reducing the potential negative competitive interactions (Vanni et al., 2009).

381

382 4.3 Conclusion and recommendation for future studies

The present study demonstrates that coexisting asp and pikeperch forage at the top of the food webs and thereby play similar functional roles in lake ecosystems. Notably, the trophic niches of the two predators were relatively well separated, both at juvenile and older life stages. The observed niche segregation may help to reduce potential interspecific resource competition between coexisting asp and pikeperch populations.

Asp and pikeperch can induce top-down cascading impacts on lower trophic levels (Benndorf, 1990; Brabrand & Faafeng, 1993; Donabaum et al., 1999). They are also popular game fishes for anglers and therefore often stocked into various systems (e.g., Ruuhijärvi et al., 1996; Wysujack et al., 2002; Vašek et al., 2013). Our results are relevant to fisheries management, because they indicate that different use of the prey resources may potentially mitigate interspecific competition between co-occurring asp and pikeperch populations. In

future studies, comparison of trophic niches of the two species under conditions of allopatry and sympatry could help to determine whether relatively low overlap in resource use is the consequence of interspecific competition or different foraging strategies that evolved in the past.

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- 563 species after the removal of a geographic barrier. *Hydrobiologia*, 797, 57–68.

- 565 Tables
- 566 Table 1. Basic environmental characteristics of the two reservoirs studied. Mean values for
- 567 the growing season (May-September) are shown for Secchi depth, total phosphorus and
- 568 chlorophyll-*a*.

Characteristic	Lipno	Římov
Year of filling	1960	1978
Surface altitude (m a.s.l.)	725	471
Surface area (km ²)	48.7	2.1
Mean depth (m)	6	16
Maximum depth (m)	22	43
Hydraulic retention time (days)	244	85
Secchi depth (m)	1.9	2.6
Total phosphorus ($\mu g L^{-1}$)	25	27
Chlorophyll- <i>a</i> (μ g L ⁻¹)	14	19

Table 2. Sample sizes in gut content analysis (GCA) (n) and results from pairwise one-way analysis of similarities (ANOSIM) comparisons of volumetric prey proportions in digestive tracts of asp and pikeperch in the Lipno and Římov reservoirs. Statistically significant

Reservoir	Size class (mm)			ANOSIM P
Lipno	<100	16	17	<0.001
	100-199	4	7	0.024
	200-299	6	11	0.262
	≥300	7	6	0.217
Římov	<100	12	20	<0.001
	100-199	14	10	0.043
	200-299	15	11	<0.001
	≥300	7	5	0.176

differences (P < 0.05) are shown in bold.

576 Table 3. List of fish species preyed upon and their total numbers found in all digestive tracts

Draw fish anaging (fourily)	As	sp	Pikeperch	
Prey fish species (family)	Lipno	Římov	Lipno	Římov
Perch (Percidae)	3	2	14	8
Pikeperch (Percidae)	1	-	11	2
Ruffe (Percidae)	7	-	6	13
Bleak (Cyprinidae)	3	9	1	-
Bream (Cyprinidae)	-	-	-	1
Roach (Cyprinidae)	1	1	-	6
Total number of prey fish discernible to species level	15	12	32	30

577 of asp and pikeperch collected from the Lipno and Římov reservoirs.

Table 4. Parameter estimates and corresponding *t*- and *P*-values for the non-linear (asymptotic) regression models with trophic position (TP) as a response variable and standard length (mm) as a predictor variable, given for asp and pikeperch in Lipno and Římov reservoirs. The models are fitted using SSasymp function in R (R Core Team 2017), producing estimates for the horizontal asymptote on the right side (Asym), the response value (i.e., TP) when length is zero (R0), and the natural logarithm of the rate constant (lrc). See Fig. 3 for sample sizes and predicted regression curves.

Reservoir	Species	Parameter	Estimate	SE	t	Р
Lipno	Asp	Asym	3.85	0.08	45.79	<0.001
		R0	0.22	0.88	0.25	0.806
		lrc	-4.29	0.27	-16.00	< 0.001
Lipno	Pikeperch	Asym	3.77	0.05	81.09	<0.001
		R0	2.03	0.26	7.93	<0.001
		lrc	-4.16	0.20	-21.28	<0.001
Římov	Asp	Asym	3.91	0.07	54.77	<0.001
		R0	2.50	0.39	6.35	<0.001
		lrc	-4.45	0.35	-12.90	<0.001
Římov	Pikeperch	Asym	4.31	0.08	51.30	<0.001
		R0	2.86	0.18	16.07	< 0.001
		lrc	-4.66	0.24	-19.16	< 0.001

Table 5. Mean trophic position (TP; standard deviation in parentheses), isotopic niche width measured as standard ellipse area (SEA_C) and niche overlap for asp and pikeperch of four size classes collected from the Lipno and Římov reservoirs. P^{\dagger} and P^{\ddagger} refer to statistical significances for TP comparisons and for niche width comparisons, respectively, between asp and pikeperch in each size class. Statistically significant differences (P < 0.05) are highlighted in bold. For number of predators analysed for stable isotopes in each size class see Fig. 4.

D	Size class	TP (SD)		P^{\dagger}	SEA _C		P^{\ddagger}	Overlap
Reservoir	(mm)	Asp	Pikeperch	P^{+}	Asp	Pikeperch	P^{*}	(%)
Lipno	<100	2.5 (0.3)	3.1 (0.2)	<0.001	2.6	3.9	0.88	0.0
	100-199	3.5 (0.2)	3.6 (0.3)	0.64	3.0	2.9	0.38	48.8
	200-299	3.7 (0.3)	3.7 (0.2)	0.94	3.2	1.3	0.03	54.9
	≥300	3.8 (0.2)	3.8 (0.2)	0.64	2.1	1.1	0.08	64.7
Římov	<100	3.3 (0.3)	3.5 (0.3)	0.07	2.7	2.7	0.51	65.2
	100-199	3.7 (0.2)	3.9 (0.1)	<0.001	1.0	1.2	0.74	42.3
	200-299	3.8 (0.1)	4.2 (0.2)	<0.001	1.2	1.1	0.28	0.0
	≥300	3.9 (0.2)	4.2 (0.2)	<0.001	1.4	0.9	0.18	0.0

594 Figure Legends

Figure 1. Mean volumetric proportion of different prey categories in the digestive tracts of different size classes of asp and pikeperch in the Lipno and Římov reservoirs. Predators that contained unidentified prey fish in their digestive tracts are not shown to better illustrate interspecific differences in piscivorous foraging on percid and cyprinid fish. The number of examined digestive tracts with discernible prey items is indicated above the bars.

600

Figure 2. Logistic regressions showing the probability of finding fish remains in gut contents as a function of asp (n = 43 & 65) and pikeperch (n = 56 & 69) standard length in the Lipno and Římov reservoirs.

604

Figure 3. Trophic positions of asp (n = 43 & 77) and pikeperch (n = 62 & 88) in the Lipno and \check{R} ímov reservoirs as a function of standard length. Lines indicate predicted values for the nonlinear (asymptotic) regression models (see parameter estimates in Table 4).

608

Figure 4. Proportional contribution (mean \pm 95% credibility intervals) of different prey types (aquatic invertebrates, terrestrial insects and fish) to the diet of different size classes of asp and pikeperch in the Lipno and Římov reservoirs, estimated using SIAR mixing model. The number of predators analysed for stable isotopes in each size class is indicated across the top of the graphs.

615 Appendix

616 Figure A1

Bi-plots of δ^{13} C and δ^{15} N values for individual asp (circles; n = 43 & 77) and pikeperch 617 (triangles; n = 62 & 88), and their principal diet sources in the Lipno and Římov reservoirs. 618 619 Filled squares represent mean ± standard deviation for pelagic zooplankton, littoral 620 macroinvertebrates, terrestrial insects and prey fish. All diet sources were corrected for trophic fractionation using values ($\delta^{13}C = 0.91$, $\delta^{15}N = 3.23$) from Vander Zanden & 621 622 Rasmussen (2001). Because pelagic zooplankton and littoral macroinvertebrates did not differ 623 in their isotope values, they were merged as "aquatic invertebrates" for the purpose of SIAR 624 estimates (see Fig. 4).









