This manuscript is textually identical with the published paper:

Gergely Boros, Péter Sály, Michael J. Vanni (2015) Ontogenetic variation in the body stoichiometry of two fish species. Oecologia, Volume 179, Issue 2, pp 329-341. DOI 10.1007/s00442-015-3349-8

Ontogenetic variation in the body stoichiometry of two fish species

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Abstract

One of the central questions of ecological stoichiometry theory is to what extent animal
species maintain constant elemental composition in their bodies. Although several recent
studies demonstrate intraspecific variation in animal elemental composition, relatively little is
known about ontogenetic changes in vertebrates, especially during early life stages. We
studied the intraspecific and interspecific ontogenetic variation in the body stoichiometry of
two fish species in two different orders; fathead minnow (Pimephales promelas) and
sheepshead minnow (Cyprinodon variegatus), reared under controlled laboratory conditions.
During ontogeny, we measured the chemical composition of fish bodies, including carbon
(C), nitrogen (N), phosphorus (P), calcium (Ca), and ribonucleic-acid (RNA) contents. We
found that N and RNA contents were relatively high in early life stages and declined
substantially during development. In contrast, body C and C:N ratios were relatively low in
embryos, post-embryos and larvae, and increased remarkably thereafter. Concentrations and
ratios of some elements (e.g., Ca, P, Ca:P) did not exhibit consistent ontogenetic trends, but
fluctuated dynamically between consecutive developmental stages in both species. Specific
growth rates correlated significantly with RNA contents in both species. Analyses of the
relative importance of different P pools at each developmental stage revealed that RNA was a
considerable P pool in post-embryos, while bone-associated P was the dominant body P pool
in later stages. Our results suggest that the elemental composition of fish bodies changes
considerably during ontogeny. Each ontogenetic stage has its own stoichiometric signature,
but the timing, magnitude and direction of ontogenetic changes can vary substantially
between taxa.

Keywords: nutrients, ecological stoichiometry, elemental homeostasis, organismal development, phosphorus pools

Introduction

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Ecological stoichiometry (ES) theory provides a framework for predicting how different species vary in storing and recycling nutrients (Sterner and Elser 2002). ES expresses biological interactions in terms of the balance of energy (carbon; C) and nutrients such as nitrogen (N) and phosphorus (P) (Sterner and Elser 2002; El-Sabaawi et al. 2012a). One of the early tenets of ES theory was that heterotrophic organisms maintain relatively constant elemental composition in their bodies, in the face of variable food nutrient contents or ingestion rates. Moreover, the theory also assumes that to a large extent the nutrient stoichiometry of animals is a genetically determined trait, arising from evolutionary pressures on form and function (Sterner and Elser 2002). The assumption of species-specific and tightly constrained elemental homeostasis generates the conclusion that body nutrient concentrations within a particular species are relatively constant across populations and life stages. However, several recent analyses challenge the notion that animals are as homeostatic in their elemental composition as previously hypothesized (e.g., Pilati and Vanni 2007; Hood and Sterner 2010; Vrede et al. 2011; Boros et al. 2012; El-Sabaawi et al. 2012 a,b; Back and King 2013; Benstead et al. 2014). These studies mandate that we reconsider and refine widespread notions about taxon-specific constancy in elemental composition. As Hendrixson et al. (2007) state, "strict homeostasis is a simplifying assumption about a complex reality, where nutrient content varies with many factors". Nakazawa (2011) argued that assuming a constant body elemental composition is only an approximation and a simplification that has been used for model development and that ecological stoichiometry theory is still incomplete in this sense. Fish have been frequently studied in the context of ecological stoichiometry, as their biomasses often constitute important nutrient pools in aquatic ecosystems (Kitchell et al. 1975; Sereda et al. 2008; Vanni et al. 2013), and they can support a substantial proportion of the demands of primary producers via nutrient recycling (Vanni 2002; McIntyre et al. 2008).

Thus, alterations in fish biomass and community assemblage influence the availability of nutrients to primary producers (McIntyre et al. 2008; Boros et al. 2009). Occupying relatively high trophic positions and being rich in nutrients, fish represent a locus where N and P are concentrated (e.g. Sterner and Elser 2002; Tarvainen et al. 2002; Vanni et al. 2013), which is important because these nutrients play a key role in limiting primary production (Lewis and Wurtsbaugh 2008). Because of their rapid growth and high mortality rates, young-of-the-year fish can be especially important in these processes (Kraft 1992; Lorenzen 2000). Hence, elemental stoichiometry of fish, including both sequestration and release of nutrients, has been of great interest during the recent decades (Kitchell et al. 1975; Parmenter and Lamarra 1991; Vanni 2002; Vrede et al. 2011; Vanni et al. 2013). Several studies demonstrate that the body stoichiometry of fish may vary with ecological and environmental conditions such as habitat, resources, food quality, trophic state, predation pressure and stress (Boros et al. 2012; El-Sabaawi et al. 2012 a,b; Benstead et al. 2014; Dalton and Flecker 2014; Sullam et al. 2015). Thus, significant intraspecific differences in elemental composition may exist among individuals of different populations. However, ontogeny also can play a role in explaining intraspecific differences in the elemental composition of fish. Recent studies report that intraspecific variability in body stoichiometry is greater than previously thought, and that body size, ontogeny and/or morphology can explain a significant part of the variation (Pilati and Vanni 2007; Vrede et al. 2011). The mass and relative proportion of different tissues and biochemicals can change dynamically during organismal development, and this can lead to changes in whole-body elemental composition, because different tissue types and biochemicals contain elements in different quantities. Bones and scales are rich in calcium (Ca) and P, while nucleic acids also contain significant amounts of P (Rønsholdt 1995; Vrede et al. 2004; Hendrixson et al. 2007). Muscle tissue stores considerable amounts of N as protein (Pangle and Sutton 2005; Vrede et

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al. 2011), while energy-rich lipids are the most important C storage pools in animal bodies (Sterner and Elser 2002; Fagan et al. 2011). During ontogeny, fish may exhibit different strategies and distinct periods of energy accumulation and somatic growth (Post and Parkinson 2001; Biro et al. 2005; Nakazawa 2011), and body stoichiometry of individuals reflects such alterations between life stages. For instance, Deegan (1986) showed that the body composition of young-of-the-year gulf menhaden (Brevoortia patronus) changed considerably during ontogeny owing to a shift in energy allocation away from protein growth to lipid storage. The growth rate hypothesis (Elser et al. 1996, 2003) states that fast-growing animals (which often include those in early life stages) need high quantities of ribonucleic acid (RNA) to achieve and maintain their high specific growth rates, and RNA content of tissues constitutes the most important P-pool of body in early phases of ontogeny (Elser et al. 1996; Vrede et al. 2004). Subsequently, RNA content of tissues declines with decreasing growth rates as ontogeny proceeds (Gillooly et al. 2005); in vertebrates this is accompanied by a gradual ossification (P and Ca allocation) of skeleton (Hendrixson et al. 2007; Pilati and Vanni 2007). For example, Sterner and Elser (2002) and Vrede et al. (2011) pointed out that rapidly growing animals commonly have low C:P and N:P ratios because of the increased P allocation to RNA. However, decreasing C:P and N:P ratios with growth were also reported for later stages of ontogeny in vertebrates, owing to the increasing P allocation to developing skeleton (Pilati and Vanni 2007). This suggests a realignment of P pools in body during ontogeny. Yet, the timing and magnitude of these changes, and how they vary among species, are still largely unknown; for fish we know very little about changes during early life stages. In this study, we explored ontogenetic changes in the body stoichiometry of two fish species in two different orders. We raised fish from embryos to adults under controlled environmental conditions, and assessed their chemical composition (C, N, P, Ca and RNA) at

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several developmental stages. The justification of our experiment is two-fold. First, preceding studies on ontogenetic stoichiometric shifts have been conducted only on a limited number of species and ontogenetic stages; to our knowledge, no previous studies include data on early developmental stages (i.e., embryos and post-embryos) as well as adults of fish. Secondly, intra- and interspecific variation in ontogenetic stoichiometry has not yet been studied in experiments in which feeding and environmental conditions are controlled. Because of these gaps, the factors that contribute to variability in organismal stoichiometry are still poorly understood and warrant more detailed examinations. We had the following objectives:

- 109 (1) To characterize the ontogenetic changes in the body composition of two fish species that belong to different taxonomic orders and that are adapted to different environments.
 - (2) To explore whether the two fish species show the same strategies in allocating nutrients to energy storage and somatic growth during ontogeny, or if the two species exhibit divergent patterns even when environmental conditions are similar.
 - (3) In the light of the growth rate hypothesis, to identify the life stages when RNA is the dominant P pool in fish, and when during development the P stored in RNA becomes negligible compared to the pool in the developing skeletal system.

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Materials and methods

Study species

We studied two fish species from the class of ray-finned fishes (Actinopterygii): fathead minnow (Pimephales promelas) and sheepshead minnow (Cyprinodon variegatus). Fathead minnow (hereafter FM), in the order Cypriniformes, is a widespread fish species across North America, inhabiting all types of freshwater ecosystems. Sheepshead minnow (SM) belongs to the order Cyprinodontiformes and lives in brackish/saltwater environments from the Mid-Atlantic United States to South America. We chose these species because both are omnivorous, are similar in size, have rapid growth to maturity under ideal temperature (22–24 °C) and food supply, and are easily raised under laboratory conditions. Thus, it was possible to conduct an experiment using the same food source and environmental conditions for both species and to raise fish to adulthood in a reasonable time frame. However, they belong to different taxonomic orders and live in different environments (freshwater *vs.* saltwater), allowing us to compare two species with different evolutionary histories.

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Experimental design

Fish were hatched and raised in the aquatic laboratories of the Animal Care Facility of Miami University (Oxford, OH, USA). Light intensity and temperature were controlled in the same manner in freshwater and saltwater rooms, with 12:12 day/night photoperiods and constant 23 °C water temperature. We obtained fish embryos from breeding individuals maintained in the facility and held them in aerated beakers, and then placed in 40 L aguariums after hatching. Both FM and SM cultures were allocated into 3 different replicate groups held in separate aquariums, to maintain a fish density that did not reduce growth. Animal handling and experimental procedures were approved by Miami University's Institutional Animal Care and Use Committee (Protocol No. 860). The experiment lasted for ~4 months, April-August 2012. For comparison of body composition, we divided the ontogeny of fish to the following categories: embryo, postembryo, larva, juvenile and adult. In the "dynamic energy budget" theory, Kooijman (2000) divided the ontogeny of multicellular animals to three basic life stages: embryo (individuals that do not feed or reproduce), juvenile (individuals that feed but do not reproduce), and adult. We elaborated on this classification by including two additional life stages (post-embryonic and larval) to characterize ontogeny at a finer scale. Designation of ontogenetic stages was based on a posteriori growing characteristics, i.e., based on distinct size classes (Table 1), and

on an individual's ability to consume bigger food particles (i.e., a diet shift that corresponded to the beginning of juvenile stage). In addition, fish > 30 mm were considered to be young adults (Van Aerle et al. 2004). Post-embryonic and larval fish were fed two times per day with brine shrimp larvae (*Artemia* sp.), while juveniles and adults consumed TetraMin flake food designed for aquarium fishes, also two times per day. During sampling days, fish were not fed in the morning to avoid the possible effects of consumed food on body chemistry analyses.

Sampling and sample analyses

We sampled fish randomly from aquariums using a hand-net. Embryos were sampled one day before hatching and post-embryos 1–3 days after hatching. All subsequent samplings were performed at 10–12 -day intervals thereafter. Samples of embryos and post-embryos were pooled (15–20 individuals per sample) because individuals in these developmental stages were too small to produce enough material for all analyses. For larvae, juveniles and adults, samples consisted of single individuals. For all samples, subsamples from whole-body homogenates were taken for the various analyses. Carbon, N, P and Ca analyses require dried samples, while RNA content can be measured only from wet tissues; thus, we had to use different fish for elemental analyses and RNA measurements. During samplings, first we randomly selected 3 fish per species for measuring elemental composition (1 fish per aquarium) and then another 3 fish of similar size for RNA analyses.

We anesthetized and sacrificed fish using ice-slurry immersion (Blessing et al. 2010). After death, length and body mass were recorded (except for embryos), and then fish for RNA analyses were immediately immersed in liquid nitrogen and stored in a -80 °C freezer until sample processing. Whole fish samples for C, N, P and Ca analyses were dried to a constant weight at 60 °C and ground to a fine powder with a mortar and pestle, and with a Retsch

ZM100 centrifugal mill (Retsch GmbH, Germany). Carbon and N contents of samples were measured using a CE Elantech Flash 2000 CHN analyser (CE Elantech, USA), while P contents were analysed following ignition at 550 °C and subsequent HCl digestion to convert all P to soluble reactive P, which was assayed with a Lachat QC 8000 FIA autoanalyser (Lachat Instruments, USA). For Ca content analysis, dried and homogenized subsamples were combusted at 550 °C and the produced ash was dissolved in HCl solution. Subsequently, Ca contents were determined with a Perkin-Elmer Optima 7300 DV Optical Emission Spectrometer (Perkin-Elmer Inc., USA). Prior to RNA analyses, deep-frozen and intact samples were homogenized with a sonicator. RNA contents of homogenates were extracted with a Maxwell LEV simplyRNA Blood Kit and Maxwell 16 Nucleic Acid Extraction System (Promega Corporation, USA). The quantity of the extracted RNA was measured with a NanoDrop 2000 spectrophotometer (Thermo Fisher Scientific Inc., USA) in the Center for Bioinformatics and Functional Genomics at Miami University.

Statistical analyses

As the first step in data analyses, we used generalized additive models (GAM) with cubic regression spline smoothers to illustrate the ontogenetic changes of our studied variables for both species (proportions of elements are expressed as percentage of dry mass, proportion of RNA is expressed as percentage of wet mass, and ratios are expressed in molar units). GAMs (Hastie and Tibshirani 1990) are ideal and commonly used for visualizing non-linear statistical relationships (e.g., Guisan et al. 2002; Buisson et al. 2008; Schmera et al. 2012), which frequently occur with ecological variables. We added 95% confidence bands to the GAM plots, in order to indicate the reliability of the predicted values of the models and to provide a visual aid for assessing differences between the two species.

Next, to explore differences between the ontogenetic stages and species, we fitted an analysis of variance (ANOVA) model for each response variable, using species and ontogenetic stage as the grouping variable (FM embryo, SM embryo, etc.). After ANOVAs, Tukey's post-hoc tests were applied to compare all developmental stages in a pairwise manner within a species (e.g., FM embryos *vs.* FM post-embryos) and between the two species (e.g., FM embryos *vs.* SM embryos).

To assess changes in the associations among Ca, RNA and P during the developmental process, we used analyses of covariance models (ANCOVA) with a nested factorial design (i.e., for RNA vs. P and Ca vs. P). In the ANCOVA models, the ontogenetic stage grouping (categorical) variable was nested within the species grouping variable. The slope regression coefficient of the ANCOVA models enabled us to evaluate the statistical relationship between the response variable (RNA and Ca) and the continuous explanatory variable (P) of the model. For the ANCOVA models, we used a contrast matrix to make the *a priori* planned pairwise comparisons of the slope regression parameters between the ontogenetic stages separately for the two species.

Relationships between body component variables and specific growth rate (SGR; Brown 1946) of fish were assessed with Pearson's correlation analysis. Calculation of specific growth rate was based on total body length increments and was calculated as follows:

$$SGR_s(\%) = \frac{(\ln(L_s + 0.01) - \ln(L_{s-1} + 0.01))}{A_s - A_{s-1}} \times 100$$

where s is a given ontogenetic stage, L_s and L_{s-1} are the average total body lengths (mm) at the s and s-1 ontogenetic stages, A_s and A_{s-1} are the average age at days of the s and s-1 ontogenetic stages. Note that calculation of SGR was not possible for embryos, and that addition of an arbitrary constant (0.01) to the formula was necessary because the body length of embryos was undefinable.

Decisions about statistical significance were set at P = 0.05 level. All statistical analyses were performed in the R environment (R Core Team 2014). We used the "mgcv" package (Wood 2006) for the GAMs, and the "multcomp" package (Hothorn et al. 2008) for the post-hoc comparisons.

The chemical compositions of both fish species changed considerably during ontogeny,

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Results

often non-linearly. Most parameters showed fluctuating patterns but often with overall increasing or decreasing trends over time (Fig. 1). Nitrogen and RNA contents and N:P ratios of fish bodies were high in early life stages and declined substantially during growth. In contrast, C contents and C:N ratios were relatively low in embryos, post-embryos and larvae, and increased markedly ~60 days after hatching, corresponding to the beginning of the juvenile stage and a diet shift from Artemia larvae (molar C/N/P ratio: 94/19/1) to flake food (C/N/P: 87/13/1). In contrast to the aforementioned relatively consistent temporal trends, P, Ca and Ca:P fluctuated dynamically between consecutive developmental stages with no such definite trends during ontogeny (Fig. 1, Fig. 2). In most cases, the means of these fluctuating variables did not differ significantly between earlier and later stages of development (Table 2). The two species showed somewhat different strategies in allocating biogenic elements to their bodies, especially in their embryonic phase (Fig. 1, Fig. 2). In particular, for many elemental contents (N, P, Ca), fluctuations between developmental stages were much more pronounced in SM than in FM. However, we also observed many commonalities between species in the long-term trends (over 120 days) (Fig. 1). For example, RNA content peaked 3-5 days after hatching and declined markedly thereafter in both species, and RNA contents were very similar between species (Fig. 2). Accordingly, specific growth rates were the

highest in post-embryos and decreased with growth both in FM (post-embryo: 210.4; larval: 248 249 3.3; juvenile: 1.7; adult: 1.1) and SM, respectively (post-embryo: 213.3; larval: 2.3; juvenile: 250 1.7; adult: 1.0). Pearson's correlation analysis revealed significant and positive relationships 251 between SGR-RNA and SGR-N:P in both species. However, we also found significant 252 negative correlations between SGR-P, SGR-Ca and SGR-Ca:P, but only in SM (Table 3). 253 We observed significant differences in many variables between consecutive ontogenetic 254 stages of the same species, as well as between the earlier and later phases of development 255 (e.g., between embryos and adults of the same species) (Table 2). These results show that 256 body composition of both FM and SM changed considerably with growth, but also that the 257 timing, magnitude and direction of these changes were dissimilar in the two species in several 258 cases. For instance, C, C:P and Ca:P ratios differed significantly between the embryonic and 259 post-embryonic stages of FM, but did not differ among these stages for SM. In contrast, N 260 and RNA were significantly different between the embryonic and post-embryonic stages of 261 SM, but were similar to each other in FM. One notable difference between species was that 262 SM embryos had much higher Ca and P than FM embryos (Fig. 2). Thus, for FM, P and Ca 263 differed markedly between earliest stages and adults, while for SM, adults differed only from 264 post-embryos with respect to these parameters. More generally, a lack of significant P and Ca 265 differences between ontogenetic stages was more typical for SM than for FM. Furthermore, 266 C:P and especially Ca:P ratios were also very similar between the different ontogenetic stages 267 of SM, in contrast to FM (Table 2). 268 Comparisons of interspecific differences within the same ontogenetic stage revealed that 269 the body compositions and ratios of elements were the most divergent in embryos, while 270 those of post-embryos and adults were very similar (Fig. 2). FM embryos had ~25% higher C 271 and N contents than SM embryos, and the difference between species was significant for both 272 variables. However, interspecific differences in C and N contents were relatively minimal

compared to differences in RNA content, C:P ratio, and N:P ratio, all of which were 140–150% higher in FM embryos. On the other hand, SM embryos contained significantly higher amounts of P (almost 2-times that of FM embryos), but the largest difference was for Ca, as SM embryos contained 10-times more of this element per unit body mass. Accordingly, molar Ca:P ratios were ~6-times higher in SM embryos than in FM embryos. Interspecific differences in Ca, P and Ca:P were negligible during the post-embryonic and larval stages, but became significant again for juveniles. Note that total lengths of the two species were very similar throughout the experiment (Fig. 1), which facilitated making interspecific comparisons in the chemical composition during ontogeny.

We found mostly positive associations, or no association, between RNA-P and Ca-P within given stages (Fig. 3). RNA and P were significantly correlated (P < 0.001) in postembryos of FM and SM, but there was no significant relationship between these two variables in other developmental stages. Regression coefficients (slope parameters) of FM and SM post-embryos differed significantly (P < 0.001) from the same parameter of their larvae, juveniles and adults. For the Ca-P relationship, we found significant regressions only in larval fishes (P < 0.001 in both species) and in FM adults. Even though the Ca-P regression coefficients appeared to vary between developmental stages (Fig. 3), the only significant difference occurred between larvae and juveniles (P < 0.05 for both species).

Discussion

In this study, we followed the ontogenetic changes in the elemental stoichiometry of two, ecologically contrasting fish species, and quantified RNA and Ca to assess the relative importance of different P pools during ontogeny. Based on the homeostasis component of ES theory (Sterner and Elser 2002), our null hypothesis was that elemental composition of an individual is constant throughout its lifespan. We recognize that this is a simplification of ES

theory (Hendrixson et al. 2007; Nakazawa 2011). However, this null hypothesis provided a "baseline" against which we could analyse the extent of deviations from it. Our results showed that elemental composition of fish bodies varied significantly among developmental stages, indicating that the fixed, species-specific elemental homeostasis does not apply to fish when rapidly growing individuals and early stages of ontogeny are included. In fact, intraspecific differences during ontogeny were often as great as variation among the two species. The observed ontogenetic plasticity and interspecific differences suggest different constraints and potentially differential elemental limitation among species, especially in the early developmental stages.

The most remarkable interspecific difference was found between embryos. The much higher (10-fold) Ca levels of SM, compared to FM, may imply that SM start bone formation in the earliest phase of ontogeny, in contrast to the FM individuals, which may have primitive skeleton in this developmental stage. However, the decline in Ca from the embryonic to postembryonic stages of SM suggests that this is not likely to be the explanation. Another possible explanation is that SM lives in saltwater, and higher concentrations of minerals (such as Ca) in the body could facilitate maintaining the osmotic balance in the embryos that have undeveloped osmoregulation.

The scarcity of studies that include all developmental stages of fish (or other animals) reared under controlled conditions renders it difficult to compare our results explicitly with previously published studies. Nevertheless, we can make some comparisons with other studies. Pilati and Vanni (2007) studied the ontogenetic changes in the body stoichiometry of gizzard shad (*Dorosoma cepedianum*) in a lake and zebrafish (*Danio rerio*) reared under controlled laboratory conditions. Even though Pilati and Vanni (2007) did not include embryos and post-embryos in their study, comparison with our results reveals several common traits across fish species. Specifically, they also found that body C increased, while

body N decreased in both gizzard shad and zebrafish, for fish beyond larval stages, similar to both fish species in our study. Increased body C probably indicates increased lipid storage after the larval stage (Fagan et al. 2011). The consistent declines in body N contents do not necessarily imply a loss of muscle mass as ontogeny proceeds, but rather may indicate that C content dominates body elemental composition and any changes in the proportion of C may drive the relative proportions of most other elements, including N. In other words, there may be a "dilution effect" of body C on other elements. Moreover, Pilati and Vanni (2007) reported increasing body P and Ca contents from the larval stage until the beginning of the early juvenile stage, and then relatively constant proportion of these elements in larger fish. We found similar trends in the changes of P and Ca in our study, and presume that variable but generally increasing levels of these elements throughout the observed period of ontogeny indicated that skeletons were developing and ossifying continuously. However, the aforementioned stoichiometric dilution could result in temporal fluctuations in body P and Ca, and a weakening of the relationship between age and elemental concentrations.

The observed positive correlation between N:P ratios and growth rates in both FM and SM

The observed positive correlation between N:P ratios and growth rates in both FM and SM is consistent with the findings of Davis and Boyd (1978), Tanner et al. (2000) and Pilati and Vanni (2007) for various fish species. Another relevant study by Vrede et al. (2011) yielded slightly different results, as they pointed out that size effect was significant on the whole-body C, P, C:N, C:P and N:P of Eurasian perch (*Perca fluviatilis*), while N contents did not change considerably with growth as it was observed in case of FM and SM. However, Vrede et al. (2011) did not include fish from the earliest phases of ontogeny in their analyses (their smallest fish were > 50 mm), thus the comparison with our results must be done with caution. Nevertheless, several similarities can be demonstrated for ontogenetic trends in C, P and C:N. Sterner and George (2000) studied the changes in the body composition of cyprinids, and reported significant negative correlations between body size and P and N contents of fish, and

and George (2000) used fish > 20 mm in their analyses, and no embryos, post-embryos or larvae. If we restrict observations only to fish > 20 mm in our study, we can see similarities with Sterner and George (2000) in C and N trends, but conflicting trends in P. Furthermore, Sterner and George (2000) found increasing N:P ratios with size in cyprinid minnows, a pattern opposite that found by Davis and Boyd (1978), Pilati and Vanni (2007), and our study. Increasing N:P and decreasing C:N ratio could indicate increased N allocation to muscle tissue (Pangle and Sutton 2005; Vrede et al. 2011). In contrast, increasing C:N and C:P ratios could be the consequences of increased lipid storage in fish. Decreasing C:P and N:P ratios along with increasing % Ca values may indicate intensive bone formation (Hendrixson et al. 2007; Pilati and Vanni 2007). Our results suggest that bone formation and the concomitant P and Ca allocation, and muscle formation, both contributed to ontogenetic changes in body N:P. This contrasts somewhat with Pilati and Vanni's (2007) findings for gizzard shad residing in a eutrophic lake, where changes in body P largely drove body N:P dynamics. However, our findings are similar to those for zebra fish grown in the lab (Vanni and Pilati 2007), which showed declining body N and increasing body P during ontogeny. Comparisons of the scant number of studies on ontogenetic variation in fish body stoichiometry reveal some commonalities, but also considerable and apparent variation among species. Opposing trends could arise from differences in the size ranges of fish used. and/or from actual interspecific differences in the dynamics of lipid storage, muscle development and bone formation during ontogeny. It should be noted that in our study, the food supply of fish was optimal, which could result in relatively high lipid storage and consequently a relatively strong dilution of elements in lower proportions (e.g., P). In nature,

the availability of food resources can be highly variable, perhaps leading to different degrees

of stoichiometric dilution. This could represent an important source of variation among

also significant but positive correlation between body length and C content of fish. Sterner

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studies in the size—element content relationship. We also note that the diet shift to which our fish were subjected, a switch from *Artemia* to flake food in juveniles, could have influenced body stoichiometry. In particular, this represented an increase in dietary C:N from 4.9 (in *Artemia*) to 6.6 (TetraMin flake food), and this was accompanied by increased body C:N in both fish species (Fig. 1, Fig. 2). Thus, ontogenetic changes in body stoichiometry may be partially attributable to changing dietary stoichiometry. However, zebrafish reared on a constant diet of *Artemia* also showed increasing body C:N as they developed (Pilati and Vanni 2007), showing that such changes in body stoichiometry can occur in absence of a change in diet stoichiometry. In general, we know little about how body stoichiometry of vertebrates varies with diet stoichiometry (Benstead et al. 2014), but it is likely that both ontogeny and diet influence body stoichiometry of vertebrates.

The general patterns of changes in RNA contents were similar in both FM and SM, suggesting that RNA production may be a strictly regulated and common trait, i.e., high RNA levels are needed during early ontogeny when specific growth rates are high. This finding is in accordance with the growth rate hypothesis (Elser et al. 1996, 2003). In our study, RNA appeared to be an important P pool in the post-embryonic stage of both species, and proved to be negligible before and after this life stage. Similar trends were described in earlier studies (e.g., Elser et al. 1996; Vrede et al. 2004), assuming that RNA-bound P determines the total body P content only in early life stages, while the beginning of bone formation along with declining RNA levels realign the P pools in the body. Accordingly, correlations between P and Ca became significant in the larvae of both species, indicating the increasing importance of bone-associated P by this developmental stage.

The vast majority of studies dealing with the elemental stoichiometry of fish report body nutrient values obtained from juvenile and/or adult specimens, and do not include larval or embryonic stages. This bias has potential implications when assessing the role of fish as nutrient sinks or sources, because younger fish cohorts often dominate population numbers or feeding rates. Given that the body composition of young individuals differs significantly from that of adults, the quantity of nutrients sequestered by growing fish, or recycled from decomposing carcasses may strongly depend on fish population age structure. Rapidly growing larvae could represent a nutrient sink, and sink strength may be especially high for P given the ontogenetic increase in this element (Kraft et al. 1992; Pilati and Vanni 2007). Natural fish mortality, which ranges between 10% – 67% per year in populations (Chidami and Amyot 2008, and references therein), is inversely proportional to body length in younger fish (Lorenzen 2000), suggesting that embryos, post-embryos or larvae are the most exposed to mortality. Decaying fish carcasses may not function as permanent nutrient sinks in many cases (Parmenter and Lamarra 1991; Boros et al. 2015), and ontogenetic variation in elemental composition could influence the rates and ratios by which carcasses release nutrients. Thus, consideration of age-specific nutrient contents could have important implications not only for ES theory, but also for determining the importance of different fish cohorts in internal nutrient loading.

Time scale is a potentially important factor when considering the importance of elemental homeostasis and the applicability of ES theory. For instance, it is acceptable to assume constant body composition in studies focusing on nutrient excretion on a given day, because body nutrient requirements and storage are not likely to vary significantly within such a short time interval. However, over longer periods and for studies with age-structured populations, ontogenetic changes must be taken into consideration. One key question then is: At what time scale does it become important to incorporate ontogenetic changes in body stoichiometry, to accurately predict nutrient cycling by animals? In terms of mediating nutrient excretion, changes in body stoichiometry are probably more important for larval fish than for older fish, because body composition and specific growth rate change more rapidly for larvae than for

older individuals (Pilati and Vanni 2007). However, much more theoretical and empirical work is needed to resolve this question.

In summary, our results provide evidence that elemental stoichiometry of fish is not simply species-specific. Rather, each ontogenetic stage may have its own stoichiometric signature, which is determined to a great extent by evolved physiological and morphological traits. Fish are excellent vertebrate models for these kinds of studies, but we also need to learn more about ontogenetic variation in animals in general. Thus, we encourage further studies that more extensively explore intraspecific and interspecific variation in body stoichiometry, including all ontogenetic stages of a wide range of aquatic and terrestrial taxa.

432 Acknowledgements

- The Rosztoczy Foundation supported Gergely Boros by providing a postdoctoral
- fellowship to conduct research at Miami University. We acknowledge the support of National
- Science Foundation grant DEB 0743192. We thank E. Mette, L. Porter, Z. Alley, A. Kiss and
- 436 A. Morgan for assistance in the lab, and MU Animal Care Facility and CBFG staff for
- 437 technical support.

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572 Figure legends

582

- 573 Fig. 1 Generalized additive models illustrating the intra- and interspecific variations in body 574 composition during ontogeny (n = 36 per species; for details, see Table 1). Percentage values in the upper left corner show the explained variances (r^2 values, i.e., the goodness of model 575 576 fit). Vertical lines indicate the end of post-embryonic, larval and juvenile stages. 577 Solid line: fathead minnow; dashed line: sheepshead minnow; dotted line: limits of 95% 578 confidence intervals 579 Fig. 2 Box-plots showing the distribution of different variables, according to species and 580 ontogenetic stages (n = 36 per species; for details, see Table 1). Letters above the boxes 581 denote the similarity/difference of the same ontogenetic group of the two species (Tukey's
- Fig. 3 Regression coefficients estimated for each ontogenetic stage from the ANCOVA models (response vs. covariant). R^2 values indicate the general variances explained by the models (i.e., the goodness of fit)

post-hoc test; a – no significant difference; b – P < 0.05; c – P < 0.01; d – P < 0.001)

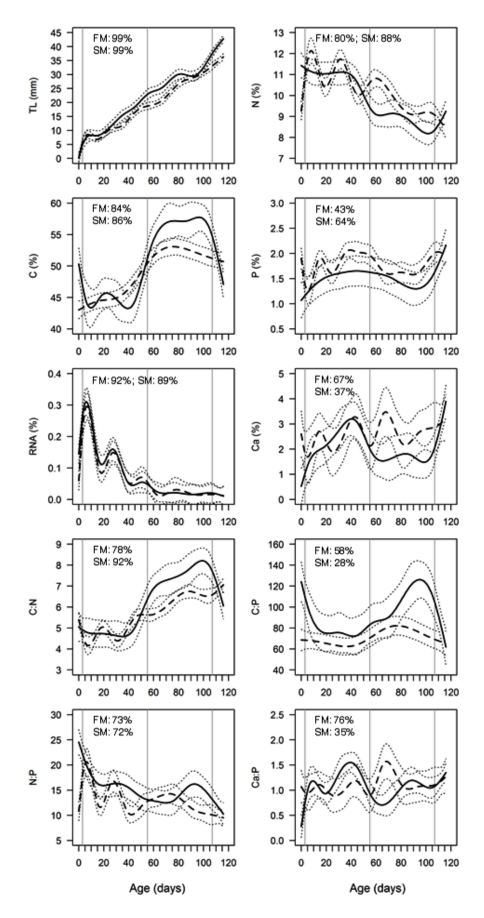
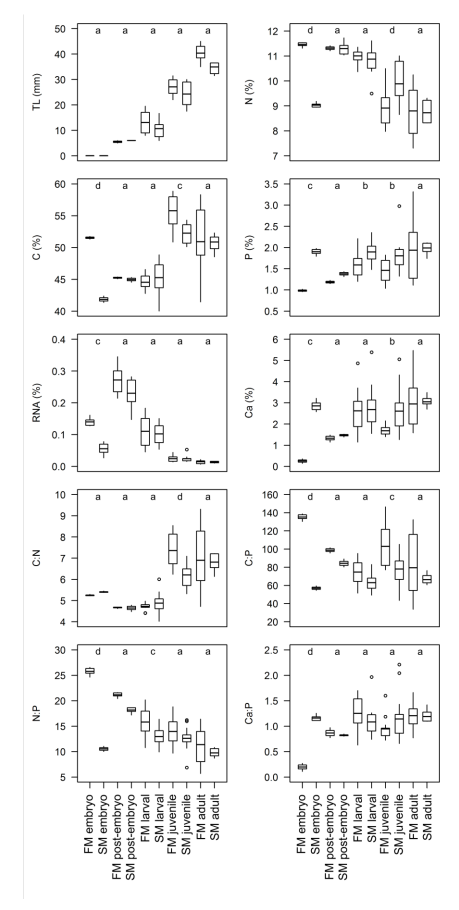
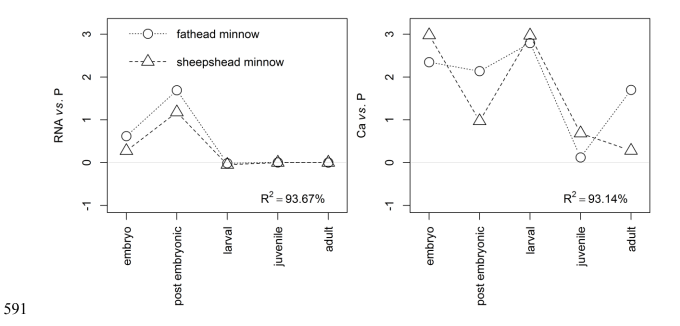


Fig. 1



588

589 Fig. 2



592 Fig. 3

Table 1 Total body lengths, sample sizes and age of fathead (FM) and sheepshead (SM) minnows at different developmental stages. Mean \pm SD denotes the arithmetic average and standard deviation; range is showed as an interval between the minimum and maximum values. Note that body length of embryos was not measurable.

		Embryo	Post-embryo	Larval	Juvenile	Adult
FM total length (mm)	mean \pm SD	-	5.5±0.5	13.0±4.4	27.1±3.3	40.3±3.4
	range	-	5.0-6.0	8.0-19.5	22.0-30.0	35.5-45.0
SM total length (mm)	$mean \pm SD$	-	6.0 ± 0.0	10.7±3.7	24.2±4.7	34.9 ± 2.3
	range	-	6.0-6.0	7.0-16.7	17.5-30.0	31.5-36.5
Number of samples/species		3	3	12	12	6
Days elapsed after hatching		0	1-3	4-55	56-107	108-120

Table 2 Tukey's pairwise comparisons from the ANOVA models, indicating several significant intraspecific differences during ontogeny. Numbers in the table denote differences in the means of variables at each ontogenetic state, while asterisks mark the significant differences (*P < 0.05; **P < 0.01; ***P < 0.001). FM – fathead minnow; SM – sheepshead minnow

	FM					SM			
C		Post-embryo	Larval	Juvenile	Adult	Post-embryo	Larval	Juvenile	Adult
	Embryo	6.30 **	7.00 ***	4.23 *	0.63	3.11	3.40	10.40 ***	9.00 ***
	Post-emb	oryo	0.70	10.53 ***	5.67 **		0.29	7.29 ***	5.90 **
	Larval			11.23 ***	6.37 ***			6.99 ***	5.61 ***
	Juvenile				4.86 ***				1.40
N		Post-embryo	Larval	Juvenile	Adult	Post-embryo	Larval	Juvenile	Adult
	Embryo	0.14	0.45	2.54 ***	2.66 ***	2.27 ***	1.85 ***	0.86 *	0.31
	Post-emb	oryo	0.31	2.40 ***	2.52 ***		0.42	1.41 **	2.57 ***
	Larval			2.09 ***	2.21 ***			0.99 ***	2.15 ***
	Juvenile				0.12				1.16 ***
P		Post-embryo	Larval	Juvenile	Adult	Post-embryo	Larval	Juvenile	Adult
	Embryo	0.20	0.60 *	0.48	0.95 ***	0.52	0.01	0.10	0.09
	Post-emb	oryo	0.40	0.28	0.75 **		0.52 *	0.42	0.61 *
	Larval			0.13	0.35 *			0.09	0.01
	Juvenile				0.48 **				0.19
RNA		Post-embryo	Larval	Juvenile	Adult	Post-embryo	Larval	Juvenile	Adult
	Embryo	0.13 ***	0.03	0.12 ***	0.13 ***	0.17 ***	0.05 *	0.03	0.04
	Post-emb	oryo	0.16 ***	0.25 ***	0.26 ***		0.13 ***	0.21 ***	0.22 ***
	Larval			0.09 ***	0.10 ***			0.08 ***	0.09 ***
	Juvenile				0.01				0.01

Ca		Post-embryo	Larval	Juvenile	Adult	Post-embryo	Larval	Juvenile	Adult
	Embryo	1.08	2.37 ***	1.43 *	2.70 ***	1.39	0.17	0.25	0.19
	Post-emb	oryo	1.29 *	0.36	1.62 **		1.22 *	1.14	1.58 *
	Larval			0.94 *	0.33			0.08	0.36
	Juvenile				1.26 **				0.44
C:N		Post-embryo		Juvenile	Adult	Post-embryo		Juvenile	Adult
	Embryo	0.58	0.52	2.11 ***	1.65 ***	0.76	0.53	0.80	1.41 **
	Post-emb	oryo	0.06	2.69 ***	2.23 ***		0.23	1.56 **	2.17 ***
	Larval			2.63 ***	2.18 ***			1.33 ***	1.94 ***
	Juvenile				0.45				0.61
~ ~				,				,	
C:P		Post-embryo		Juvenile	Adult	Post-embryo		Juvenile	Adult
	Embryo	36.54 *	60.60 ***	32.27 *	55.88 ***	27.41	6.00	21.13	9.41
	Post-emb	oryo	24.06	4.27	19.34		21.41	6.27	18.00
	Larval			28.33 ***	4.71			15.13 *	3.41
	Juvenile				23.62 **				11.73
N:P		Post-embryo	Lorvol	Juvenile	Adult	Post-embryo	Lorvol	Juvenile	Adult
IV:I	F1	,	9.97 ***		14.41 ***	•			
	Embryo					7.03	2.42	2.06	0.79
	Post-emb	oryo	5.36 ***	7.21 ***	9.79 ***		5.21 **	5.56 ***	8.41 ***
	Larval			1.85	4.44 ***			0.36	3.21 **
	Juvenile				2.58 *				2.85 *
Ca:P		Post-embryo	Larval	Juvenile	Adult	Post-embryo	Larval	Juvenile	Adult
	Embryo	0.67 *	1.05 ***	0.75 ***	1.01 ***	0.34	0.08	0.02	0.03
	Post-emb	oryo	0.38	0.08	0.34		0.26	0.32	0.37
	Larval			0.31 *	0.05			0.06	0.11
	Juvenile				0.26				0.05

Table 3 Pearson's correlation tests of specific growth rates and body component variables. The "r" denotes the correlation coefficient, lower and upper 95% are the limits of the confidence intervals of the correlation coefficient, and "P" is the value of significance for the test with a null hypothesis of r = 0. Significant correlations are marked with an asterisk.

Fathead minnow

	r	lower 95%	upper 95%	P	
TL	-0.82	-0.99	0.68	0.18	_
C	-0.59	-0.99	0.86	0.41	
N	0.78	-0.73	0.99	0.22	
P	-0.83	-0.99	0.66	0.17	
RNA*	0.98	0.35	0.99	0.02	
Ca	-0.71	-0.99	0.79	0.29	
C:N	-0.71	-0.99	0.79	0.29	
C:P	0.41	-0.91	0.98	0.59	
N:P*	0.96	0.04	0.99	0.03	
Ca:P	-0.65	-0.99	0.83	0.35	

Sheepshead	minnow
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	r	lower 95%	upper 95%	P
TL	-0.75	-0.99	0.75	0.25
C	-0.66	-0.99	0.82	0.34
N	0.74	-0.76	0.99	0.26
P*	-0.97	-0.99	-0.22	0.02
RNA*	0.96	-0.06	0.99	0.04
Ca*	-0.98	-0.99	-0.43	0.01
C:N	-0.73	-0.99	0.78	0.27
C:P	0.75	-0.76	0.99	0.25
N:P*	0.96	-0.03	0.99	0.04
Ca:P*	-0.99	-0.99	-0.64	0.01