Understanding variation in salamander ionomes: A nutrient balance approach

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#### 1 Summary

Ecological stoichiometry uses information on a few key biological elements (C, N, and P) to
 explain complex ecological patterns. Although factors driving variation in these elements are
 well-established, expanding stoichiometric principles to explore dynamics of the many other
 essential elements comprising biological tissues (i.e., the ionome) is needed to determine their
 metabolic relationships and better understand biological control of elemental flows through
 ecosystems.

8 2. In this paper, we report observations of ionomic variation in two species of salamander

9 (Ambystoma opacum and A. talpoideum) across ontogenic stages using specimens from

10 biological collections of two wetlands sampled over a 30-year period. This unique data set

allowed us to explore the extent of ionomic variation between species, among ontogenic stages,between sites, and through time.

3. We found species- and to a lesser extent site-specific differences in C, N, and P along with 13
other elements forming salamander ionomes but saw no evidence of temporal changes.
Salamander ionomic composition was most strongly related to ontogeny with relatively higher
concentrations of many elements in adult males (i.e., Ca, P, S, Mg, Zn, and Cu) compared to
metamorphic juveniles, which had greater amounts of C, Fe, and Mn.

4. In addition to patterns of individual elements, covariance among elements was used to
construct multi-elemental nutrient balances, which revealed differences in salamander elemental
composition between species and sites and changes in elemental proportions across ontogenic
development. These multi-elemental balances distinguished among species-site-ontogenic stage
groups better than using only C, N, and P.

5. Overall, this study highlights the responsiveness of consumer ionomes to life-history and
environmental variation while reflecting underlying relationships among elements tied to
biological function. As such, ionomic studies can provide important insights into factors shaping
consumer elemental composition and for predicting how these changes might affect higher-order
ecological processes.

### 28 Introduction

29 Elements are the fundamental building blocks of living cells and are involved in all metabolic 30 processes. Organisms must take up elements from the environment and, despite vast differences 31 in environmental supplies, maintain their body elemental composition within a relatively narrow 32 range (Persson et al., 2010). Elemental content differs considerably among species due to diversity in classical traits (e.g., life-history, morphological, and physiological traits) that are 33 constructed using different elements or the same elements in differing proportions (Jeyasingh, 34 35 Cothran & Tobler, 2014). Thus, it follows that the elemental composition of an individual is determined by the acquisition, assimilation, and allocation of elements within the organism. This 36 37 abstraction is useful because information on the elemental content of a species enables the 38 application of mass-balance principles to make predictions about sequestration and excretion of elements, both of which have strong impacts on ecological dynamics (Reiners, 1986). This 39 stoichiometric approach has triggered a vibrant area of research integrating data on species 40 elemental composition with cellular processes and organismal life-history to study key 41 ecological phenomena (Sterner & Elser, 2002). 42

43 In addition to species-level differences, it is apparent that there is considerable intraspecific variation in organismal elemental composition (Jeyasingh et al., 2014; Prater, 44 Wagner & Frost, 2017). For example, Bertram et al. (2008) found that variation in the 45 46 phosphorus (P) content of a single cricket species collected across field populations was as extensive as that observed across several orders of insect taxa (Woods et al., 2004). Further, 47 48 strong P plasticity among distinct ontogenic stages in metazoans linked to differences in organismal growth and reproduction has commonly been observed (e.g., Villar-argaiz et al. 49 50 2002, Capps et al. 2015, Tiegs et al. 2016). Besides P, recent work also highlights the potential ecological importance of variation in other elements within organismal tissues, although
metabolic causes of this variation are not always well-known (e.g., Baxter 2010; Jeyasingh et al.
2017). At a time when it is becoming increasingly clear that growth and productivity can be
limited by combinations of elements (Salt, Baxter & Lahner, 2008; Harpole *et al.*, 2011; Parent *et al.*, 2013a), determination of the extent and nature of variation in the content of the other ~20
essential elements found in biological tissues is needed to establish their connections with
organismal metabolism and ecological functions.

58 The ionome is defined as the mineral nutrient and trace element composition of an organism (Salt, Baxter & Lahner, 2008), which underlies its morphological, anatomical, and 59 60 physiological state. Although little is known about the ionomes of phagotrophic metazoans (but 61 see Goos et al. 2017), studies on osmotrophs such as Saccharomyces cerevisiae (Eide et al., 2005) and Arabidopsis thaliana (Baxter et al., 2008) in controlled laboratory conditions clearly 62 show that genetics and resource-supply stoichiometry interact to shape organismal growth and 63 ionomic profiles. Evidence for ionome-wide shifts from the field are also available. For example, 64 Synechococcus cells collected from regions of the Sargasso Sea that vary in nitrogen (N) and P 65 66 supplies exhibited several-fold cell quota differences in a variety of elements [e.g., manganese (Mn), nickel (Ni), and zinc (Zn); Twining et al. 2010]. These studies depict a strong correlative 67 network among elements likely underpinned by a complex web of metabolic connections (Parent 68 69 et al., 2013a; Baxter, 2015). However, our knowledge of factors related to elemental variation in vertebrate consumers, particularly in ecologically sensitive taxa such as amphibians, remains 70 71 limited to a relatively small number of elements.

Throughout ontogeny, frog and salamander carbon (C) content generally decreases while
%N, P, calcium (Ca), and sulfur (S) tend to increase (Capps *et al.*, 2015; Luhring, DeLong &

Semlitsch, 2017). These patterns are correlated with developmental and growth rates (Bumpers
et al. 2015; Stephens et al. 2017), arguably due to ontogenic changes in macromolecular
demands that differ in stoichiometric composition (e.g., lipids rich in C, proteins rich in N or S,
and bones rich in Ca and P; Costello and Michel 2013; Liess et al. 2013; Stephens et al. 2017). In
addition to ontogeny, amphibian C:N:P stoichiometry differs considerably across sites at local
scales (Milanovich & Hopton, 2014) due to differences in both bottom-up nutrient supplies
(Stephens, Berven & Tiegs, 2013; Bumpers et al., 2015) and top-down predation pressure
(Costello & Michel, 2013). Furthermore, abiotic factors such as hydroperiod and temperature
have been shown to alter population-level stoichiometry at decadal scales (Capps et al. 2015) and
might explain regional stoichiometric differences through local adaptation of amphibian
populations across latitudinal gradients (Liess et al., 2013). Overall, amphibian composition of a
handful of elements appears to be highly responsive to many ecological factors, and these
changes could result in shifts of other metabolically connected elements ultimately influencing
biogeochemical fluxes through aquatic environments.
Here, we report observations on ionomic variation in two Ambystomatid salamander
species [marbled salamanders (Ambystoma opacum) and mole salamanders (Ambystoma
talpoideum)] using specimens that were collected at two wetlands over a 30-yr period. Based on
stoichiometric theory, we hypothesized that the ionomes of each species should vary due to
differences in life-history and ontogeny, and we sought to identify linkages among elements,
including those already known to vary significantly (i.e., C, Ca, N, P, and S; Capps et al. 2015,
Tiegs et al. 2016, Luhring et al. 2017). Moreover, because hydroperiod can affect salamander
growth and development, we hypothesized that salamander ionomes should vary due to

96 documented differences in hydroperiod between sites. Since hydroperiod differed considerably

over the 30-yr time period during which samples were collected (Daszak *et al.*, 2005; Todd *et al.*,

98 2011), we also expected to see temporal changes in salamander ionomes. In addition to

99 examining variation in individual elements, we also constructed nutrient balances (Parent *et al.*,

100 2013a,b) using knowledge of elemental metabolism and multivariate data from 15 elements to

see how these basic biological factors influence covariation among suites of elements.

### 102 Methods

103 Study Sites, Specimen Collection, and Salamander Life-History: Study animals were collected at the United States Department of Energy's Savannah River Site (SRS) in South Carolina. 104 105 Salamanders were sampled at two ephemeral wetlands, Rainbow Bay (RB: 33.315696; -81.773643) and Ginger's Bay (GB: 33.260035; -81.631320), as part of a long-term monitoring 106 program that began in 1978 and 1986 at each wetland, respectively (Scott, 1990; Pechmann et 107 108 al., 1991). Rainbow Bay and GB are similar in area at full ponding (1-1.5 ha) but are located 14 km apart in different physiographic subregions. Rainbow Bay is found at ~97 m elevation on the 109 110 Aiken Plateau, whereas GB occurs on an old Pleistocene floodplain terrace of the Savannah River (Sunderland Terrace) at 61 m elevation. Both wetlands are surrounded by well-drained 111 sandy loam soils in the Fuguay and Dothan series, but wetlands themselves are located on 112 113 poorly-drained sandy fine loam soils (Davis & Janecek, 1997). Wetland soils also differ in elemental composition from surrounding upland soils as depressional wetland soils on the SRS 114 tend to have elevated aluminum (Al) and barium (Ba) concentrations and total N, P, and C levels 115 compared to upland soils (Looney et al., 1990; Dixon et al., 1997). Wetlands on the Sunderland 116 Terrace (GB) usually hold water for more extended periods (i.e., longer hydroperiod) than Aiken 117 Plateu wetlands (RB), have dissimilar zooplankton communities (Mahoney, Mort & Taylor, 118 119 1990), and may have differences in cation concentrations (Schalles et al., 1989). The mean

120	hydroperiod at RB is $129.6 \pm 98.5$ days (mean $\pm 1$ SD), and it has never held water from one year
121	to the next; GB in contrast has a longer hydroperiod (D.E. Scott and K.A. Capps unpublished
122	data) and remains filled year-round on average once every three years.

Each wetland was encircled by a drift fence with paired 19-L pitfall traps positioned at 123 10-m intervals. Individuals entering and leaving the wetlands were censused daily (RB) or 124 periodically (GB) during the breeding season which occurs from late fall through early spring for 125 our study species (Semlitsch et al., 1993; Scott et al., 2013). When mortality occurred during the 126 127 study (usually due to predation or bucket flooding) deceased salamanders were collected, 128 identified individuals to species, and terrestrial adults were sexed. Partial remains of specimens 129 exposed to predators were used for census counts but excluded from our current study. We 130 classified specimens as one of three ontogenic stages: terrestrial adult male (A. opacum only), recently emerged metamorph (both species), or recently emerged aquatic adult paedomorph (A. 131 *talpoideum* only; defined below) and stored them in a -70°C freezer until processing (Nunziata, 132 Scott & Lance, 2015). A total of 214 organisms were analyzed in our study representing a mix of 133 species, sites, and ontogenic stages collected between 1986-2015 (Supp. Table 1). All study 134 135 specimens were collected under annual renewals of South Carolina Department of Natural Resources collection permits and triannual renewals of animal capture and handling protocols 136 approved by the University of Georgia Institutional Animal Care and Use Committee. 137

Ambystomatid salamander life-history differs considerably between species. *Ambystoma opacum* breeds terrestrially in late summer/autumn (Sept. – Dec.) and oviposits in nests in dry pond beds prior to wetland filling (Scott, 2005). *Ambystoma talpoideum* breeds aquatically during late fall and winter (Nov. - Mar.), and females lay single eggs or small chusters attached to aquatic vegetation (Pechmann et al. 1991, Semlitsch 1987). *Ambystoma talpoideum* eggs typically hatch about two months later than *A. opacum*, which may lead to a predatory or
competitive advantage for *A. opacum* larvae (Boone, Scott & Niewiarowski, 2002). In wetlands
such as GB that occasionally hold water from one year to the next, *A. talpoideum* are
facultatively paedomorphic. Rather than metamorphosing and emigrating to terrestrial uplands,
these individuals may remain in the wetland, retain their gills, become sexually mature aquatic
adults in the winter (Semlitsch, 1987), and usually exit the wetland in spring after one
reproductive bout.

150 Ambystomatid salamander life-history varies with environmental conditions including temperature, predation, food quantity/quality, and population density, but appears to be most 151 strongly tied to pond hydroperiod (Semlitsch 1987, Scott 1990, Pechmann et al. 1991, Daszak et 152 153 al. 2005, Stephens et al. 2017). Yet, despite this extensive phenotypic variation clear differences exist between our study species. Both species appear to have relatively similar larval growth 154 155 rates (Semlitsch, 1987; Scott, 1990), although A. talpoideum typically metamorphoses later and 156 at a larger body size than A. opacum (Pechmann, 1995). Reproductive success (i.e., the production of juveniles that metamorphose and emigrate from wetlands) of both species is tied to 157 hydroperiod, but A. talpoideum requires a later date of pond drying and more frequently faces 158 159 inadequate water at RB, which can result in catastrophic larval mortality (Daszak et al., 2005). Consequently, both species had stable populations at GB over our study period (Nunziata *et al.*, 160 161 2015), but A. opacum populations increased, and A. talpoideum decreased apparently due to a shortened hydroperiod at RB (Daszak et al., 2005). Few mortality events occurred for terrestrial 162 A. talpoideum; as a result, fewer of these species were collected compared to A. opacum. This 163 164 partially explains the uneven sample sizes and complete lack of adult A. talpoideum in our data set (Supp. Table 1). 165

166	Sample Processing and Ionomic Profile Generation: We removed animals from the
167	freezer and placed them individually into trace clean polypropylene grinding vials. Carcasses
168	were then freeze-dried and powdered using a Spex Mill (8000D, Metuchen, NJ) and
169	methacrylate grinding balls, and ground tissues were stored at -20°C. Before elemental analyses,
170	we dried subsamples overnight at 60°C and weighed them to the nearest 0.1 $\mu$ g. Tissue C and N
171	content was measured using an automated vario MICRO cube analyzer (Elementar Americas
172	Inc., Mt. Laurel, NJ). To measure all other elements, we digested between 3-12 mg of
173	homogenized tissues in a 2:1 $v/v$ solution of trace metal grade nitric acid and hydrogen peroxide
174	(BDH Aristar ® Plus, VWR International, Radnor, PA) for 24h in metal-free polypropylene
175	tubes (VWR International, Radnor, PA) followed by dilution to 10 ml with trace metal grade
176	water. Digested samples were then analyzed through inductively coupled plasma optical
177	emission spectrometry (ICP-OES; Thermo Scientific iCAP 7400, Waltham, MA). Sample
178	nutrient concentrations were determined using standard curves from multi-element standards
179	(CCV stds. 1A&B, CPI International, Santa Rosa, CA) and calibrated using an internal Yttrium
180	standard (Peak Performance Inorganic Y Standard, CPI International, Santa Rosa, CA).
181	We measured a total of 28 elements across all organisms. However, measurements for
182	many trace elements fell below the limits of detection (LOD) for many samples (Supp. Table 2).
183	After omitting all elements with analytical uncertainties, a total of 16 elements: Al, Ba, C, Ca,
184	copper (Cu), iron (Fe), potassium (K), lithium (Li), magnesium (Mg), Mn, N, sodium (Na), P, S,
185	strontium (Sr), and Zn were analyzed. For multivariate statistics, any remaining values falling
186	below the LOD's were replaced with 1/2 of the LOD value calculated for each individual element.
187	All elemental concentrations were then converted to percentages by dividing them by the total
188	sample dry mass (Supp. Table 3).

190 To visualize relationships between individual elements in Euclidian space, we first 191 conducted partial least squares (PLS) regressions using ontogeny, species, site, and year as 192 independent variables and log-transformed elemental percentages as dependent variables. We 193 also calculated variable importance (VIP) scores to compare the relative importance of each predictor variable (Wold, Sjöström & Eriksson, 2001; Eriksson et al., 2013). Plots of PLS 194 weights revealed correlations between elements and predictor variables, which differentially 195 196 separated across two factors (Fig. 1). These correlations reflect biologically relevant differences 197 in elemental signatures among organisms. Therefore, we used these partitions (i.e., elemental 198 position in Euclidian space) to construct nutrient balances to further explore differences in 199 elemental profiles among ontogenic stages, species, sites, and years.

To more thoroughly investigate sources of variation in salamander ionomes, we 200 201 constructed a series of isometric log-ratio balances. These balances represent unbiased estimates of multivariate relationships between elements, which avoid violating common statistical 202 assumptions and can be used to describe and interpret elemental interactions in organismal 203 204 tissues (Parent *et al.*, 2013a). Specifically defined in the context of our study and mathematically defined below, nutrient balances represent orthogonal log contrasts of elements derived from 205 binary partitions of multivariate elemental data projected into Euclidean space. To construct 206 207 these balances (hereafter referred to as nutrient or elemental balances), we first separated all elements into bulk and trace elements based on the classification of Frausto da Silva and 208 209 Williams (2001). Then we constructed one bulk and one trace elemental balance to reflect 210 elemental partitions along each PLS axis using absolute PLS weight values of 0.1 as a cutoff for 211 including elements in each balance. Following conventions (Sterner & Elser, 2002), we ordered

elements from high to low in each balance as a function of their percentage of dry mass. Nutrientbalances were then calculated using the equation:

Balance= SQRT  $(rs/r+s) \ln [g(c^+)/g(c^-)]$  (Parent *et al.*, 2013a)

where r and s represent the number of elements on the left- and right-hand side of the balance 215 216 and  $g(c^{-})$  and  $g(c^{-})$  minus represent the geometric mean of elemental percentages on the left- and 217 right-hand side of the balance, respectively. To calculate a [C | Ca, P] balance for an adult salamander with an elemental composition of 43% C, 5% Ca, and 3% P for example, we would 218 219 use the formula  $[C | Ca, P] = SQRT [(1x2)/(1+2)] \ln [(43/g(5,3)] = 1.97$ . Balances constructed in 220 this way can then be used to be used to examine proportional differences in elemental 221 composition between groups of organisms such as adults and juveniles that are related to 222 underlying biological factors. For instance, we would expect that [C | Ca, P] balance differences between these two ontogenic stages should reflect proportionally higher investment in C-rich 223 224 lipids in juveniles and higher Ca and P investment into bone for adults resulting in relatively higher juvenile values (Fig. 2). 225

226 In total, we constructed 4 novel balances: bulk [C | Ca, P, S, Mg]; [N | Ca, Na, Mg] and 227 trace [Fe, Mn | Zn, Ba, Cu, Sr, Li]; [Fe, Zn, Al, Ba | Cu, Sr, Li] in addition to two traditional stoichiometric balances [C | P] and [N | P]. These balances represent a combination of 1) 228 knowledge of well-known elemental relationships related to organismal macromolecular 229 230 composition (C, N, P, Ca) and 2) statistically identified biologically relevant relationships related to our study variables (ontogeny, species, site, and time). As such their formation represents a 231 232 blend of metabolically grounded expert knowledge and exploratory biplot analyses (Parent *et al.*, 2013b), which while not exhaustive may serve as a starting point for future functional studies. 233

To compare differences in individual elemental balances, we first separated the data into 234 235 unique species, site, and ontogenic stage groups (n=7) and conducted one-way analysis of 236 variance (ANOVAs) to examine differences across groups. After identifying significant group differences, we designed contrasts to compare differences across species at the same ontogenic 237 stage (metamorphs averaged across sites), sites (separate comparisons of metamorphs and adult 238 239 males averaged across species), and stages (A. opacum males and metamorphs averaged across sites and A. *talpoideum* metamorphs and paedomorphs from GB) using post-hoc Tukey's tests. 240 241 Specific comparisons were made rather than full factorial comparisons as complete species, site, 242 stage combinations were not available. All P-values were adjusted using Bonferroni corrections 243 to provide conservative estimates of elemental differences for comparison to multivariate 244 statistics.

In addition to univariate comparisons, we first used Wilks' Lambda tests from 245 multivariate ANOVA (MANOVA) to determine multivariate differences in nutrient balances 246 247 among each species, ontogenic stage, and site groups. We used similar methods to examine temporal patterns in salamander balances by conducting MANOVAs on salamanders separated 248 249 into species(site) data sets for each ontogenic stage, which were grouped by decade to 250 compensate for uneven sample collection and inadequate samples sizes in certain years across 251 the 30-year period. Finally, we conducted two separate discriminant analyses including all 252 balances and one including only [C | P] and [N | P] balances to: a) visualize multivariate trait 253 differences across groups and b) examine the classification accuracy of classical stoichiometric balances vs. all elemental balances. Cross-validation of these models was conducted using leave-254 255 one-out methods. Overall, these complimentary univariate and multivariate analyses were designed to quantify ecological sources of variance in individual balances (ANOVAs) and 256

multivariate phenotypes (MANOVAs & discriminant analyses) and reduce the likelihood of
statistical artifacts associated with multivariate data analyses.

259 **Results** 

260 Individual Elemental Profiles:

261 Ontogeny was the strongest predictor of elemental variation according to PLS regressions followed by species and site, which were moderate and weak predictors, respectively (Fig. 1A). 262 Temporal effects were the weakest by far (VIP< 0.2) indicating that salamander elemental 263 content was relatively invariant through time. Elemental composition was most divergent along 264 the first PLS factor, which separated strongly between terrestrial adult males (hereby referred to 265 266 as males) and juvenile metamorphic ontogenic stages with aquatic adult paeodomorphs showing intermediate phenotypes (Fig. 1B). Species and sites also separated out to a lesser extent along 267 268 this axis with A. opacum and individuals from GB falling on the left-hand side and A. talpoideum 269 and specimens from RB located on the right. Bulk elemental concentrations on the male side of the axis were generally higher for all elements other than C, which was higher in metamorphs 270 271 and N, Na, and K, which did not differ greatly along this axis. Trace elements were were also 272 higher in males for all elements except for Fe and Mn. Compared to the first factor, factor two explained far less variation but seemed to separate paedo- and metamorphic ontogenic stages 273 along with species and site differences. Relatively fewer bulk elements separated on this axis as 274 N was positively correlated and Ca, Mg, and Na were negatively correlated with factor two, 275 276 respectively. In contrast, all trace elements differed along the second axis with higher 277 concentrations of Fe and Zn associated with A. opacum and RB and A. talpoideum and 278 salamanders from GB showing higher amounts of Cu.

280	In addition to individual elements, nutrient balances differed across species, sites, and
281	ontogenic stages (Fig. 3). Metamorphs of A. opacum and A. talpoideum differed significantly in
282	their body $[N   P]$ and $[C   P]$ but did not differ in other balances (Table 1). In contrast,
283	salamander stoichiometric balances did not differ between sites, but we found significant site-
284	specific differences for all other multi-element balances in metamorphs and for [N   Ca, Na, Mg]
285	balances in males. All balances differed significantly among A. opacum males and metamorphs,
286	but only [N   P] differed between A. talpoideum metamorphs and paedomorphs.
287	Balances for unique species-site-ontogenic stage group combinations also differed in
288	multivariate space ( $P$ <0.001), but we found no evidence of significant temporal changes ( $P$ >0.1).
289	According to the discriminant analysis, male A. opacum phenotypes diverged strongly from other
290	groups along the first discriminant axis (Fig. 4), as males showed lower [C   P] and [N   Ca, Na,
291	Mg] values and other stages had higher [C   Ca, P, S, Mg] and [Fe, Mn   Zn, Ba, Cu, Sr, Li]
292	balances. Species differences separated out on the second discriminant axis as A. opacum
293	generally showed lower [C   P] and [Fe, Zn, Al, Ba   Cu, Sr, Li] concentrations and A. talpoideum
294	had higher $[N   P]$ and $[C   Ca, P, S, Mg]$ . Elemental balances showed relatively greater overlap
295	between sites in A. talpoideum but were more negatively correlated with the second discriminant
296	axis for each ontogenic stage in A. opacum from RB. Dual balances explained nearly half of the
297	elemental variation among all groups but were inadequate for distinguishing between
298	metamorphs of each species (Table 2). Including additional multi-elemental balances improved
299	classification accuracy from 49 to 65% among all groups. Despite our conservative analyses, it is
300	important to note that elemental correlations and nutrient balances in our data set could still
301	possibly contain information attributable to artifacts of high-dimensional analyses. Although we

302 cannot quantify the potential for these artifacts, consistency between uni- and multivariate
303 statistics suggests that their likelihood of influencing our findings was minimal.

## 304 Discussion

In this study, we found ecologically relevant differences in traditionally studied stoichiometric 305 306 elements (i.e., C, N, and P) along with a suite of other elements comprising salamander ionomes. In contrast, we saw no significant temporal changes in the elemental composition of either 307 species. Salamander ionomic composition was most strongly related to ontogeny with relatively 308 309 higher concentrations of many elements in males (Ca, P, S, Mg, Zn, and Cu) compared to recently emerged metamorphic juveniles, which had greater amounts of C, Fe, and Mn. In 310 311 addition to identifying correlations among single elements, we also found differences among 312 species, site, and ontogenic stages in multivariate space. We observed systematic differences in elemental relationships between species and sites and evidence of suites of changes in elemental 313 proportions across ontogenic stages. Together, these results demonstrate the utility of ionomic 314 approaches for studying environmental and biological sources of consumer elemental variation 315 316 and their ecological effects.

317 Similar to previous amphibian studies (Milanovich, Maerz & Rosemond, 2015; Luhring et al., 2017), we found strong differences in the elemental composition of our study species. 318 Consistent with stoichiometric predictions, the species with the smaller body size and faster 319 320 developmental rates, A. opacum, had higher body P content than A. talpoideum. The largerbodied A. talpoideum also had higher body %C, which supports previous work showing positive 321 322 relationships between body size, juvenile lipid content, and body %C in ambystomatid salamanders from this study region (Scott et al., 2007; Luhring et al., 2017). In addition to these 323 324 traditional stoichiometric elements, we also confirmed species differences in salamander Ca and

S content found in previous work (Milanovich *et al.*, 2015; Luhring *et al.*, 2017) and identified differences in several other essential elements. Altogether, these patterns validate the focus on well-studied elements (i.e., C and P) for determining the ecological importance of interspecific variation in body stoichiometry. Our results also suggest that expanding the range of study elements could yield further insights into the ecological roles and nutritional ecology of these species.

Compared to species, we found relatively weaker evidence of spatio-temporal effects on 331 332 salamander ionomes. Site was a poor predictor of stoichiometric variation in males and metamorphs despite strong differences in genetic structure for A. opacum between these 333 334 populations (Nunziata et al., 2015). Male multi-elemental balances were also similar across sites, 335 but metamorphs showed strong contrasts between RB and GB. These patterns seem to indicate that while male nutrient uptake (i.e., diet based) appeared to be similar in the surrounding 336 337 uplands, metamorph elemental acquisition either from dietary sources or epithelial diffusion directly from the water column (e.g., metals or ions; Motais and Garcia-Romeau 1972, Handy et 338 al. 2002) likely differed across wetlands separated by only a short distance (~14 km). In addition 339 340 to differences between sites, there was also extensive annual variation in breeding season temperature and hydroperiod within each site (Pechmann et al., 1991; Daszak et al., 2005; Todd 341 et al., 2011). This variation may have limited our ability to identify temporal changes in 342 343 salamander elemental profiles and likely resulted in a lack of consistent selection pressures on 344 salamander life-history traits tied to their elemental composition. Indeed, previous genetic work examining a subset of animals from our study (Nunziata et al., 2017) along with our phenotypic 345 346 results seem to preclude the role of evolutionary change in shaping observed ionomic patterns in our study populations. However, as longer-term hydroperiod trends have been tied to shifts in 347

species abundances at RB (Daszak *et al.*, 2005), hydroperiod could nevertheless still modify
nutrient cycles in these wetlands by causing local species extinctions and thus shaping the
elemental profiles of the remaining amphibian communities. However, these ecological changes
would almost certainly interact with organismal life-history and ontogeny to ultimately control
these dynamics.

Salamander ontogenic stage had the strongest effect on elemental variation in our study 353 animals. Consistent with a previous amphibian study, we found that larger-bodied terrestrial 354 355 adult males had considerably higher body %P than juvenile metamorphs (Tiegs et al., 2016). Higher %P along with %Ca is typically associated with increased bone development and 356 ossification in adults (Kemp & Hoyt, 1969; Milanovich et al., 2015; Luhring et al., 2017) and 357 358 may explain the lower adult [C | P] and [N | Ca, Na, Mg] balances found in our study. In addition to these elements, we measured higher concentrations of many other essential elements in adult 359 tissues (Zn, S, Li, Cu, and Mg). Our results also confirm previous measurements of higher 360 juvenile body %C and no difference in body N content between terrestrial adults and 361 metamorphic amphibians (Tiegs et al., 2016) while showing differences in Fe and Mn 362 concentrations and in related [Fe, Mn | Zn, Ba, Cu, Sr, Li] nutrient balances between these 363 groups for the first time. This suggests that in addition to traditionally examined stoichiometric 364 elements (e.g., N and P; Stephens et al. 2017), ontogenic variation in organismal elemental 365 366 composition may reflect unique stage-specific nutritional physiologies and elemental requirements. However, it also appears that environmental nutrient supplies may potentially 367 override these differences as we found substantial overlap in elemental profiles between pond-368 369 dwelling A. talpoideum metamorphs and paedomorphic adults in GB, which only differed in their %N content out of all elements surveyed (Supp. Table 3). As such, it is clear that we should 370

371 consider moving beyond studies of individual or pairs of elements and examine integrated
372 changes in suites of metabolically connected elements by refining tools such as nutrient balances
373 (Parent *et al.*, 2013a; Baxter, 2015) to better understand the complex relationships between
374 environmental nutrient supplies and organismal life-history traits and ionomes.

Traditional stoichiometric ratios relate nutrient interactions to biological functions (e.g., 375 N:P ratios to ribosomal protein translation; Loladze and Elser 2011), and nutrient balance 376 concepts extend this approach to encompass the entire organismal ionome (Parent et al., 2013a). 377 378 While it is beyond the scope of this paper to functionally relate changes in salamander balances to specific metabolic pathways and physiological process, our study demonstrates the potential 379 380 of using nutrient balance principles to further our understanding of relationships between 381 consumer ecology and elemental composition. We found evidence for novel nutrient balances [C | Ca, P, S, Mg] and [Fe, Mn | Zn, Ba, Cu, Sr, Li] that differed across ontogenic stages, and 382 similar to plant-based studies, we documented systematic differences in elemental combinations 383 384 between species and sites (Watanabe et al., 2007; Parent et al., 2013a). Previously constructed balances inherently differ from ours due to their focus on specific aspects of plant physiology, 385 386 but the substantial overlap between elemental relationships in our study and previous research is perhaps indicative of fundamental elemental relationships governed by biological processes 387 operating at the cellular level (Sterner & Elser, 2002; Watanabe et al., 2007; Baxter et al., 2008; 388 389 Parent *et al.*, 2013a).

In this study, we examined correlations between salamander ionomic profiles and
biological and environmental variables. As previously demonstrated, traditional stoichiometric
balances (C | P and N | P) were useful for highlighting species differences in biogeochemically
important elements (Sterner & Elser, 2002). However, we also showed that focusing solely on

traditional stoichiometric elements alone could neglect other important elements, which might be 394 differentially required for optimal metabolic functioning in terrestrial adult males (Ca, S, Mg, 395 396 Zn, & Cu) and juveniles (Fe & Mn), respectively. Ionomic data analyzed here using balance techniques could thus be used to generate a greater understanding of the nutritional ecology of 397 amphibian development. Such data are vital for providing a starting point for designing 398 399 manipulative experiments to not only better understand the nutritional ecology of threatened vertebrates such as salamanders but also to illuminate the entire suite of ecological functions that 400 401 they might influence.

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## 413 Conflict of Interest Statement

414 The authors declare no conflicts of interest.

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**Table 1.** Trait differences between species, sites, and ontogenic stages. Degrees of freedom (*df*), f-ratio of mean squares (F), and *P*-values (*P*) are reported for 1-way main effects ANOVAs and multiple comparisons. Significant differences are shown in bold for *P*-values adjusted using Bonferroni corrections (Meta: 0.05/3=0.017; Adult: 0.05/2=0.025). Specific contrast details include: averaged across sites<sup>1</sup>, averaged across species<sup>2</sup>, *Ambystoma opacum* only<sup>3</sup>, average across sites *A. opacum* only<sup>4</sup>, and *A. talpoideum* from Ginger's Bay only<sup>5</sup>. All elements are reported in standard scientific notation. Other abbreviations include: metamorph (Meta) and paedomorph (Paedo).

		[C	[C P] [N P]		[C Ca,P,S,Mg]		[N Ca,Na,Mg]		[Fe,Mn Zn,Ba,Cu,Sr,Li]		[Fe,Zn,Al,H	Ba Cu,Sr,Li]	
Model	df	F	Р	F	Р	F	Р	F	Р	F	Р	F	Р
Group	6	44.88	<0.001	34.86	<0.001	53.56	<0.001	19.52	<0.001	64.52	<0.001	27.07	<0.001
$\mathbf{R}^2$	R <sup>2</sup> 0.565		565	0.5	0.503 0.608		0.361		0.652		0.440		
Contrast	df	F	Р	F	Р	F	Р	F	Р	F	Р	F	Р
<sup>1</sup> Species (Meta)	1	7.12	0.008	8.26	0.005	0.85	0.358	0.02	0.877	1.53	0.217	0.00	0.998
<sup>2</sup> Site (Meta)	1	5.02	0.026	3.36	0.068	6.02	0.015	9.93	0.002	10.40	0.002	20.66	<0.001
<sup>3</sup> Site (Male)	1	0.54	0.463	1.21	0.272	2.11	0.148	14.54	<0.001	0.95	0.332	0.49	0.484
<sup>4</sup> Stage (Male vs. Meta)	1	165.07	<0.001	122.92	<0.001	221.84	<0.001	60.10	<0.001	259.22	<0.001	89.82	<0.001
<sup>5</sup> Stage (Meta vs. Paedo)	1	0.04	0.838	9.33	0.003	0.01	0.906	3.26	0.072	0.26	0.611	4.60	0.033

**Table 2.** Comparison of discriminant analysis classification accuracy based on A) stoichiometric balances only and B) all elemental balances. Cross-validation results are reported as percentages, and percentages correctly assigned to each group are shown in bold.

 Abbreviations include: *Ambystoma opacum* (Amop), *A. talpoideum* (Amta), metamorph (Meta), paedomorph (Paedo), Ginger's Bay (GB), and Rainbow Bay (RB).

# A. [C|P] & [N|P]

Species	Ontogenic Stage	Site	Group	1	2	3	4	5	6	7	Total
Amop	Male	GB	1	61.5	12.8	0	25.6	0	0	0	49.1
Amop	Male	RB	2	43.2	35.1	0	18.9	0	0	2.7	
Amop	Meta	GB	3	0	0	0	100	0	0	0	
Amop	Meta	RB	4	0	0	0	<b>98.4</b>	0	1.6	0	
Amta	Meta	GB	5	0	5.6	0	83.3	0	11.1	0	
Amta	Meta	RB	6	0	0	0	71.4	0	9.5	19	
Amta	Paedo	GB	7	0	0	0	50	0	0	50	

# B. All Balances

Species	Ontogenic Stage	Site	Group	1	2	3	4	5	6	7	Total
Amop	Male	GB	1	76.9	17.9	0	0	2.6	2.6	0	64.5
Amop	Male	RB	2	35.1	62.2	0	2.7	0	0	0	
Amop	Meta	GB	3	0	0	33.3	48.1	11.1	7.4	0	
Amop	Meta	RB	4	0	3.2	8.1	87.1	1.6	0	0	
Amta	Meta	GB	5	0	0	16.7	16.7	38.9	22.2	5.6	
Amta	Meta	RB	6	0	0	4.8	19	23.8	33.3	19	
Amta	Paedo	GB	7	0	0	10	0	10	0	80	

## **Figure Captions**

**Figure 1.** Relationships between ontogeny, species, site, date, and salamander elemental composition. A) Variable importance predictor scores show the relative influence of predictor variables on organismal ionomic profiles. Scores <0.8 are considered weakly important, those between 0.8-1 are moderately important, and scores >1 are strong predictors. B) Partial least squares regression (PLS) loadings plot of variables weights demonstrates correlations between independent (x) and dependent (y) variables where relationships between variables are directly proportional to the sign and distance of all other variables in Euclidian space. Abbreviations of x-variables include: metamorph (meta), paedomorph (Paedo), *Ambystoma opacum* (Amop), *A. talpoideum* (Amta), Ginger's Bay (GB), and Rainbow Bay (RB).

**Figure 2.** Hypothetical differences in salamander elemental composition between adult and juvenile salamanders. Panel A) depicts univariate differences in body elemental composition between an adult and a juvenile salamander. Panel B) represents a mobile-and-fulcrum plot (*sensu* Parent et al. 2013) showing multivariate differences in [C | Ca, P] balances between these two ontogenic stages. The black circle represents an equilibrium point where salamander C and [Ca, P] content is equal for a given specimen, and the other circles correspond to balance points for each stage.

**Figure 3.** Variation in nutrient balances among species, sites, and ontogenic stages. Boxplots depict medians,  $25^{\text{th}}$  and  $75^{\text{th}}$  percentiles (boxes), and  $10^{\text{th}}$  and  $90^{\text{th}}$  percentiles (error bars) for each balance. All balances differed significantly among species (left- and right-hand panels), sites, and ontogenic stages according to Wilks's lambda scores (*P*< 0.001). Abbreviations include: metamorph (Meta), paedomorph (Paedo), Ginger's Bay (GB; Panels A&B), and Rainbow Bay (RB; Panels C&D).

**Figure 4.** Multivariate relationships among species, sites, and ontogenic stages. Discriminant analysis (DA) loadings plots of centroids and 95% confidence intervals are shown for each group, and the two highest positive and negative standardized canonical discriminant function coefficient scores out of six nutrient balances are reported for each axis. All balances differed significantly among groups according to Wilks's lambda scores (P< 0.001). Abbreviations include: metamorph (Meta), paedomorph (Paedo), *Ambystoma opacum* (Amop), *A. talpoideum* (Amta), Ginger's Bay (GB), and Rainbow Bay (RB).



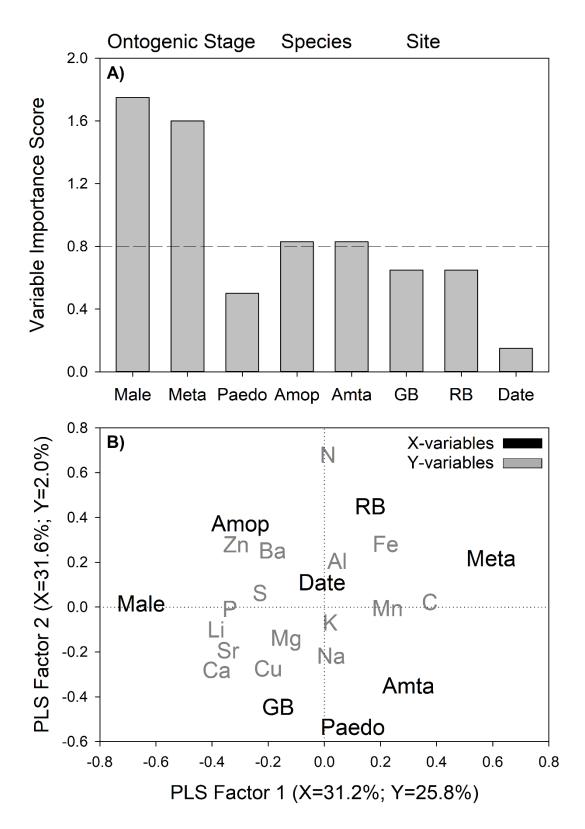
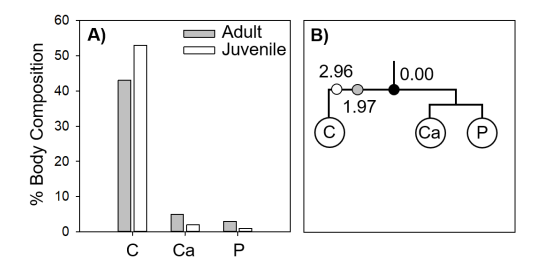


Figure 2





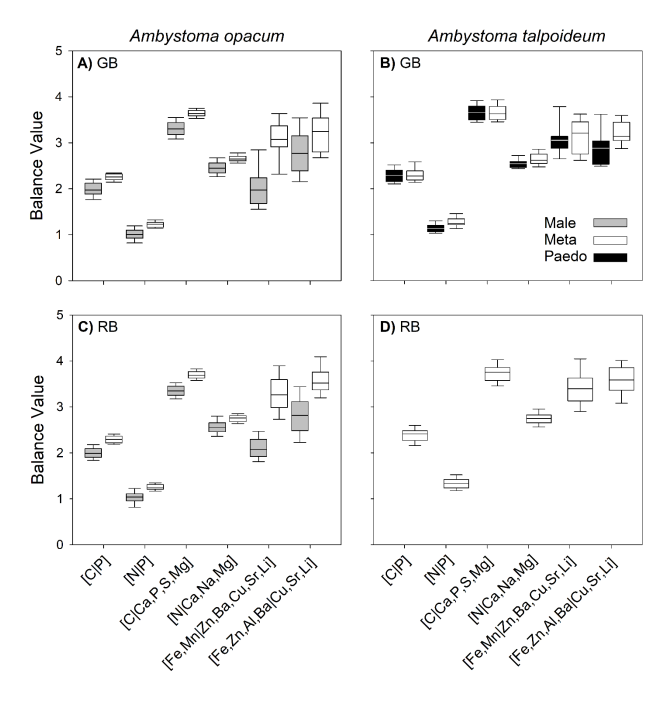


Figure 4

