1	The effects of preformed vitamin A and provitamin A carotenoid supplementation on
2	tadpoles of the poison frog <i>Phyllobates vittatus</i>
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39 40

41 Abstract

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43 Understanding the nutritional requirements of captive animals is necessary for proper animal husbandry, however, the specific dietary requirements for many amphibian species 44 commonly kept in captivity are unknown. Like most vertebrates, frogs cannot synthesize 45 carotenoids or vitamin A, and must therefore obtain these essential nutrients through diet. It is 46 unclear if amphibians can cleave provitamin A carotenoids to form vitamin A metabolically 47 48 within the body, so common practice is to supplement their captive diets with both preformed 49 vitamin A and provitamin A carotenoids. We carried out a feeding experiment in tadpoles of 50 *Phyllobates vittatus*, a commonly kept poison frog species, to test the effects of supplementing a fish flake diet with a provitamin A carotenoid (2.5mg/g β -carotene) and vitamin A (0.5 mg/g 51 52 retinyl acetate), both individually and in combination. Contrary to our expectations, 53 supplementation had either no effect or adverse effects on tadpole growth and survivorship. 54 Tadpoles reared under supplemented diets showed higher mortality rates, coupled with symptoms of hypervitaminosis A. Survivors had a smaller body size and mass at metamorphosis. 55 The vitamin A and β -carotene levels in our supplemented diet have been shown to benefit 56 57 tadpoles of other species, yet our results indicate that adding these amounts to what is found in a 58 generalist fish flake mix (5.466 μ g/g and 1.88 μ g/g, respectively) can have detrimental effects on *P. vittatus* tadpoles. More broadly, this study highlights the importance of creating husbandry 59 60 guidelines based on the specific physiological needs of the species (or species groups) being kept in captivity, rather than general ones for all amphibians, as is often done. 61 62 63 64 65 Introduction

The effects of preformed vitamin A and provitamin A carotenoid supplementation on

tadpoles of the poison frog *Phyllobates vittatus*

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As wild populations of a considerable number of amphibian species decline (Campbell et 67 al., 2020), captive populations of some of them are increasing. This is due to both their 68 69 increasing popularity in the pet trade and captive breeding programs (Edmonds, 2021; Tapley et 70 al., 2015). Proper husbandry is required for all captive animals to ensure that their physiological 71 needs are met. An important aspect of proper husbandry is animal nutrition. However, the 72 nutritional requirements of most amphibians are unknown (Tapley et al., 2015). Breeders of 73 captive frogs are aware that feeder insects, like *Drosophila* species, do not contain all of the 74 nutrients that amphibians require and, thus, enhance diets with commercially available 75 supplements. Despite the wide variety of products available for amphibian diet supplementation, 76 there is very little research to inform their use in particular species. 77 Carotenoids are powerful antioxidants that quench and stabilize reactive oxygen species

78 produced during periods of growth and development (Goodwin, 1984; Keogh et al., 2018;

79 McInerney et al. 2019). This limits the damage that reactive oxygen species cause to cells, which 80 is believed to result in increased cell division and differentiation (Keogh et al., 2018; McInerney et al. 2019). In amphibians, antioxidants are especially important during the larval stage and 81 82 metamorphic climax because this is when the production of reactive oxygen species is greatest (Keogh et al., 2018; McInerney et al. 2019). Furthermore, carotenoids and their metabolism are 83 84 involved in coloration in a variety of species across the tree of life (Goodwin, 1984; Toews et al. 2017). Since frogs, like most animals, are unable to produce carotenoids, they must be obtained 85 exclusively through their diet (Densmore and Green 2007; Keogh et al., 2018; McInerney et al. 86 2019). It is therefore common practice amongst frog breeders to add carotenoid supplements to 87 the captive diets of most species. However, breeders must be careful not to over supplement with 88 89 carotenoids as they can have negative effects when given in excess (Keogh et al., 2018; Palozza 90 et al. 2003). For instance, one of the most commonly used carotenoids, β-carotene, can be toxic at high concentrations and has been found to have a pro-oxidant effect that can cause a reduced 91 growth rate (Keogh et al., 2018; Palozza et al. 2003). 92

93

94 For many animals, carotenoids also act as vitamin A precursors (Brenes-Soto and Dierenfeld, 2014; Clugston and Blaner, 2014). This means that provitamin A carotenoids can be 95 cleaved to form vitamin A metabolically within the body. Whether this is the case for 96 97 amphibians remains unclear. The few studies that have addressed the problem in vivo have not 98 found evidence for conversion of specific carotenoids to vitamin A (McComb 2010; Collins et 99 al., 1952; Wright, 2006), although there is evidence that at least some species may be able to perform this conversion (Baruah and Goswami 2012, Brenes-Soto and Dierenfeld 2014). 100 Vitamin A plays important roles in gene expression, vision, limb regeneration and embryonic 101 development (McInerney et al. 2019; Clugston and Blaner 2014), and in frogs its deficiency has 102 been linked to squamous metaplasia of the mouth, which is commonly referred to as "short 103 tongue syndrome" (Clugston and Blaner 2014). Frogs affected by short tongue syndrome have 104 difficulty catching prey, resulting in lethargy, weight loss, and eventually death if untreated 105 (Clugston and Blaner 2014). To prevent hypovitaminosis A, a common practice amongst 106 107 breeders today is to co-supplement with carotenoids and preformed vitamin A.

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109 Nevertheless, since vitamin A is fat-soluble, excessive intake poses a risk for toxicity 110 (Densmore and Green 2007). Hypervitaminosis A has been recorded in African clawed frogs 111 (Xenopus laevis) at different stages of their life cycle (Clugston and Blaner 2014). This condition 112 causes weight loss and negatively affects the skin and liver of adult frogs (Pessier 2013), and has 113 been reported to cause abnormal development in tadpoles, with notable observations of 114 hyperpigmented, hemorrhagic skin and chronic diarrhea (Weissmann 1961; Weissman et al. 115 1963). Proper dosages of preformed vitamin A have not been determined for most species, even 116 if under or over-supplementation can lead to hypovitaminosis A or hypervitaminosis A, both of 117 which can be life threatening if untreated (Clugston and Blaner 2014). 118

119 With the goal of improving husbandry practices of captive amphibians, this study 120 investigated the effects of supplementing feed with β -carotene (a commonly used provitamin A 121 carotenoid), vitamin A, and their combination on tadpoles of the poison-dart frog *Phyllobates* 122 vittatus, a species commonly kept in captivity. Through observation and measurement of 123 tadpoles during development, we evaluated the effect of the above supplementation regimes on tadpole survivorship, growth rates, final size at metamorphosis, and time to metamorphosis. Our 124 125 results provide somewhat unexpected initial insights into the dietary carotenoid requirements of *Phyllobates* poison-dart frog tadpoles, and highlight the need for further species-specific research 126 127 on the diets of captive amphibians.

128 Methods

129 Study Species

130 *P. vittatus* is a diurnal, terrestrial species that is endemic to the Golfo Dulce Region of 131 Costa Rica, where it can be found in lowland moist and wet forests. Adults range in size from 132 22.5mm to 31mm, with females being larger than the males. P. vittatus is black with a broad 133 gold, red-orange, or orange dorsolateral stripe that runs continuously from the dorsal base of 134 each thigh to the tip of the snout, and blue-green speckling on its limbs. Their diet is made up of 135 a variety of small arthropods. Females will lay clutches of 7-21 eggs every two to three weeks. The male cares for the eggs by providing them with moisture, and once the eggs hatch, males 136 will transport the tadpoles to a pool of water (Silverstone 1976), where they likely maintain an 137 138 omnivorous diet (Grant et al. 2006). Although this has not been studied in *P. vittatus*, predatory 139 behaviors, including tadpole cannibalism, have been observed in multiple closely related species 140 (Caldwell and de Araujo 1998, Márquez *in review*). Tadpoles are uniformly dark brown on the dorsal side, and are lighter brown ventrally, and gradually develop orange dorsolateral stripes, 141 142 starting around Gosner stage 37 (Gosner 1960). Metamorphosis occurs approximately ten weeks after hatching (Savage 2002; Paitz and Dugas 2021, R. Márquez pers. obs. R. Arkin pers. obs.). 143

144 Experimental Animals

145 A total of 35 *P. vittatus* eggs, from 6 different clutches, were collected from a single 146 captive colony of 7 adults (4M, 3F) maintained at the University of Michigan. Adults are fed Drosophila hydei and D. melanogaster fruit flies three times a week, supplemented with 147 Repashy Calcium Plus, Rep-Cal Herptivite multivitamin, and Rep-Cal Calcium with Vitamin D 148 (one supplement per feeding). All eggs were laid between 1/20/22 and 2/11/22. They were 149 150 removed from the terrarium and reared in petri dishes until hatching. Thirteen eggs from 2 151 clutches had been de-jellied shortly after laying. We found no differences between these and 152 other eggs in any of the measured parameters (Table S1). All eggs hatched within the time frame typical of our lab colony (mean 16.4 days, s.d. 2.65, range 14-23 days). One tadpole died within
2 days of hatching and before ever being fed, so it was removed from the study.

155 Experimental diets

All four diets were composed of of 99.7% ground fish flakes (75:25 mixture of 156 SeraFlora/SeraSans; SERA, Germany), which was previously analyzed for nutrient content 157 158 (Byrne and Silla, 2017), and contributes 0.01454 mg g⁻¹ total carotenoids and 0.00188 mg g⁻¹ preformed vitamin A The remaining 0.03% varied between four treatments as follows: Diet 1 159 (control) was unsupplemented. Diet 2 was supplemented with 2.5mg g⁻¹ β -carotene (Sigma 160 Aldrich, Cat. No. C9750), Diet 3 was supplemented with 0.5 mg g⁻¹ vitamin A acetate (i.e. retinyl 161 acetate; Sigma-Aldrich Cat. No. R4632), and Diet 4 was supplemented with 2.5mg g⁻¹ β-carotene 162 and 0.5 mg g⁻¹ vitamin A. Cellulose powder (Sigma Aldrich Cat. No. 435236), which is odorless 163 and has been found to have no impact on the growth, survivorship, or development of tadpoles 164 (Keogh et al., 2018; Dias et al., 1998; Deng et al., 2006), was added to treatments 1, 2, and 3 to 165 166 ensure that all experimental diets contained the same amount of feed. Table 1 shows the total 167 amounts of carotenoids and vitamin A in each diet treatment, and Table S2 summarizes the 168 recipes used in each diet. β -carotene was selected as the provitamin A carotenoid due to its 169 commercial availability and presence in popular supplements. Retinyl acetate was chosen 170 because it is the form of vitamin A commonly found in commercial vitamin A supplements for 171 both animals and humans (e.g. Repashy vitamin A plus). All diet mixes were suspended to a 1:10 172 ratio in deionized H₂O, wrapped in aluminum foil and mixed on a rocker for 48 hr at room 173 temperature. Once mixed, diets were loaded onto 3ml syringes, wrapped in foil, and stored at 0 to -4°C until used. 174

175 Feeding experiment

176 After hatching, individuals were randomly assigned to one of four treatments, and housed in individual 230 ml plastic cups filled with approximately 150ml of reverse-osmosis (RO) 177 178 water. They were fed triweekly: Monday, Wednesday, and Friday. Before each feeding, syringes 179 were thawed and the feed was homogenized using a blunt tip needle. Tadpoles received 2 drops 180 of food (~0.1 ml) for the first month. After that time, individuals received 4 drops of feed (~0.2 181 ml). Tadpoles were photographed using a Canon EOS R6 camera with a Canon RF24-105mm F4 182 L IS USM lens after each feeding. The body (from tip of snout to base of tail) and tail length of 183 each individual were measured using image analysis software (ImageJ, version 1.53; Schneider 184 et al. 2012). Feeding and photographing stopped once tadpoles began the metamorphic climax at_ 185 Gosner's stage 42, as evidenced by the emergence of forelimbs (Gosner, 1960). Once both arms

- 186 emerged, tadpoles were moved to a tilted 1L cup with ~75ml of new RO water to facilitate
- 187 leaving the water. Once out of the water, froglets were moved to plastic containers (10 cm x 20
- 188 cm x 10 cm) lined with sphagnum moss and dried magnolia leaves. Froglets were monitored
- 189 every two days until they reached Gosner's stage 46, marked by the full absorption of their tail
- 190 (Gosner, 1960). At that time, individuals were photographed (NIKON D5100 camera with an AF
- 191 Micro-Nikkor 60mm f/2.8D lens) and their body length and weight were recorded. Two tadpoles
- 192 had not yet developed arms after 81 days, so they were put into new 230 ml plastic cups and a
- 193 partial water change was performed. This was done to clear the algae buildup that made it
- 194 difficult to observe and photograph the tadpole.

195 Statistical analyses

To compare mortality over time between treatments, we computed survival curves using the
Kaplan-Meier method (Kaplan & Meier 1958) and tested for differences between survival curves

Trapian Meler menoa (Trapian & Meler 1930) and tested for anterenees between survival earves

using the chi-squared approach described by Harrington & Fleming (1982). Survival curves were

- 199 fitted using the survfit() function and comparisons were made using the survdiff() function, both
- 200 implemented in the R package survival (Therneau 2022).

We then evaluated the effect of diet treatment of tadpole growth as a function of age by fittinglogistic curves to each tadpole's growth trajectory and comparing the best-fitting growth rate and

203 maximum size parameters between treatments. The logistic equation was used in the following

204 form:

205
$$S_t = \frac{M}{1 + \left(\frac{M - m}{m}\right)e^{-m}}$$

206 where S_t is the size at a given age, M is the maximum size asymptote, m is the minimum size asymptote, r is the growth rate, and t is the age. Growth curves were fit to each individual using 207 208 non-linear least squares as implemented in the function nlsLM() of the minpack.lm R package 209 (Elzhov et al. 2022). We tested for differences in maximum size and growth parameter estimates 210 between treatments using bi-variate anovas with treatment and whether tadpoles survived to 211 metamorphosis as fixed factors. To further explore differences between treatments and 212 survivorship categories, we conducted post-hoc Tukey-Kramer tests. Finally, we tested for 213 between-treatment differences in our empirical measurements of size, weight and age at 214 metamorphosis also using anovas and Tukey-Kramer tests. The data and code used for all 215 analyses and visualizations are available on the dryad repository associated with this paper.

216 Results

217 *Effects of diet on survivorship*

Overall, 82.35% (28/34) of tadpoles survived to complete metamorphosis. Tadpoles that 218 219 were fed diets supplemented with vitamin A (i.e. diets 3 and 4) had higher mortality rates, 220 especially the treatment group that was supplemented with both vitamin A and β -carotene (i.e. 221 Diet 4), which showed a clear excess of observed deaths relative to the expectation under the Kaplan-Meier model ($\chi^2 = 9.67$, df = 3, p = 0.02; Fig 1A-B). Remarkably, tadpoles that died in 222 the treatment groups supplemented with vitamin A (1 in treatment 3, 4 in treatment 4), were 223 224 observed to have skin and gastrointestinal symptoms (persistent diarrhea) that reflect what was 225 described in X. laevis tadpoles with hypervitaminosis A (Fig 1C; Weissmann 1961; Weissman et 226 al. 1963).

227 Effects of diet on growth

228 All of the tested tadpoles displayed growth trajectories characterized by initial rapid growth that gradually slows down towards metamorphosis (Fig. 2A, S1). We found significant 229 effects of treatment, survivorship, and their interaction on both maximum body size (Fig. 2B; 230 231 treatment: $F_{3,27} = 32.5$, p = 4.2e-9; survivorship: $F_{1,27} = 171.3$, p = 3.3e-13; treatment × survivorship: $F_{2,27} = 16.9$, p = 1.7e-5) and growth rate (Fig. 2C; treatment: $F_{3,27} = 15.9$, p = 3.7e-5) 232 6; survivorship: $F_{1,27} = 67.4$, p = 8.2e-9; treatment × survivorship: $F_{2,27} = 12.4$, p = 1.5e-4). 233 234 However, most of the signal in these tests is due to the tadpoles that did not survive to 235 metamorphosis, most of which displayed faster initial growth spurts but stopped growing soon 236 after, at considerably smaller sizes, where they remained for several days before dying (Fig. 2A-237 C; Fig. S1). The only significant post-hoc comparisons were either between non-surviving tadpoles of different diet treatments, or between surviving and non-surviving tadpoles, both 238 239 between and within treatments (Table 2).

240 Among the tadpoles that survived to metamorphosis, diet treatment had moderate effects on body length ($F_{3,24}$ = 3.41, p = 0.034) and body mass ($F_{3,24}$ = 2.68, p= 0.069) at metamorphosis, 241 242 but not on age at metamorphosis ($F_{3,24} = 1.06$, p = 0.38;)Fig 2D-F). Individuals fed the control 243 diet (Diet 1) were on average the largest and heaviest at metamorphosis, while those 244 supplemented with just vitamin A (Diet 3) were smaller and weighed less than those in other treatments. Individuals supplemented with β-carotene (Diet 2) and β-carotene combined with 245 246 vitamin A (Diet 4) were found to have intermediate sizes and weights (Fig 2D-E). However, the 247 only statistically significant pairwise comparisons in both metamorphic size and weight were

between the vitamin A and control treatments (Diet 3 vs. Diet 1; Table 3), possibly due to ourmoderate sample sizes.

250

251 Discussion

Our goal in this study was to examine the effects of supplementing the diets of P. vittatus 252 253 tadpoles with additional carotenoids (β -carotene) and preformed vitamin A. This practice is 254 commonly carried out by captive amphibian owners and caretakers, but the safety, benefits, and 255 appropriate dosages of these supplements have only been studied in a limited number of species. 256 The results of our feeding experiment suggest that, at least in the case of *P. vittatus*, supplementing with additional β -carotene and/or vitamin A than what is found in our base fish 257 258 flake feed (5.466 μ g/g β -carotene and 0.00188 mg/g preformed vitamin A, respectively) does not 259 provide any benefits, and can have detrimental effects in terms of tadpole mortality, and 260 probably also on pre-metamorphic growth.

The most severe negative effects were observed in the tadpoles fed Diet 4, which 261 contained both supplemental β-carotene and vitamin A. This treatment resulted in a significantly 262 263 lower survival rate than the other tested diets, and the affected individuals displayed clearly 264 different growth trajectories, as well as symptoms reminiscent of descriptions of 265 hypervitaminosis A in tadpoles of other frog species (Fig 1C; Weissmann 1961; Weissman et al. 266 1963), suggesting that these tadpoles died due to a diet-associated disease, most likely hypervitaminosis A. The same was true for the single tadpole that died under Diet 3 (only 267 268 vitamin A supplement). The fact that tadpoles supplemented with only vitamin A were much less 269 likely to die or show symptoms than those supplemented with both β -carotene and vitamin A could be due to some of the β -carotene being converted endogenously to vitamin A. While it is 270 271 commonly assumed that amphibians are not capable of cleaving provitamin A carotenoids to 272 form vitamin A, there is evidence that suggests that at least some species can do so (Brenes-Soto 273 and Dierenfeld 2014, Baruah and Goswami 2012). For instance, false tomato frogs (Dyscophus 274 guineti)supplemented with a mixture of carotenoids showed increased vitamin A plasma levels, 275 indicating they are likely able to convert carotenoids to vitamin A (Brenes-Soto and Dierenfeld 276 2014). In vitro experiments on the β -carotene metabolism of Asian common toad (*Duttaphrvnus* melanostictus) and Indian bullfrog (Haplobatrachus tigerinus) tadpoles also suggest that they are 277 278 able to convert β-carotene to vitamin A (Baruah and Goswami 2012). However, it is unknown if 279 this is the case for *P. vittatus* tadpoles, so further research is necessary to isolate the 280 physiological causes of this result. In the end, if a species is able to convert carotenoids into

vitamin A, then proper carotenoid supplementation could eliminate the need for additional
preformed vitamin A supplements, highlighting the importance of species-specific research for
husbandry practices.

We based the composition of the experimental diets on previous work with other 284 (distantly related) frog species (e.g. Pseudophryne corroboree, Byrne and Silla 2017; Keogh et 285 286 al. 2018; Weissmann 1961; Weissman et al. 1963), in an attempt to avoid over supplementation . 287 Yet, it seems clear that tadpoles in this study did experience adverse effects associated with over 288 supplementation. This highlights the importance of knowledge on the biology of focal species for 289 adequate captive breeding. The lack of research on the nutritional requirements of most 290 amphibian species kept in captivity (or their close relatives) is concerning, considering the 291 increasing number of species kept in the pet trade, in zoos, and in ex-situ conservation efforts. Understanding the nutritional requirements of captive amphibians is especially important for 292 293 endangered species since adequate husbandry is the keystone of captive breeding programs. 294 Implementing the proper diet with proper levels of supplementation leads to larger, healthier, 295 more genetically diverse populations for release, which is essential for reintroduction success 296 (McInerney et al. 2019).

297 More generally, our findings highlight that the dietary requirements of frogs vary 298 between species. This is not surprising considering the clade made up by anurans shared a common ancestor approximately 221.8 million years ago (Kumar et al. 2022). Despite their 299 300 degree of evolutionary divergence, it is common for all frog species to be grouped as a single 301 category in husbandry contexts, which leads to and this belief is reflected by assumed similarities 302 in husbandry practices, which may be unwarranted. This frame of thinking is not usually applied 303 to more "mainstream" species kept in captivity. For instance cats (Felis catus) and dogs (Canis 304 *lupus familiaris*), for which clearly distinct husbandry practices are in place, diverged 305 approximately 55 million years ago (Kumar et al. 2022). In contrast, dendrobatids and 306 mantellids, two families of frogs commonly kept in captivity, diverged approximately 150 307 million years ago (Kumar et al. 2022) but are assumed to have very similar dietary and other 308 husbandry needs. Considering the phylogenetic diversity of amphibians there is a clear need for 309 further research to establish adequate husbandry and veterinary guidelines and practices that 310 reflect the specific needs of each species (or at least taxonomic group). This information can 311 improve the success of ex-situ conservation efforts, as well as the health of amphibians kept as pets, in zoos, and in laboratories. Our study is a step in this direction for *Phyllobates* poison-dart 312 313 frogs, and possibly other dendrobatid frogs kept in captivity.

315 Author Contributions

- 316 RA and RM conceived and designed the study. RA obtained the data. RA and RM analyzed the
- 317 data. RA wrote the manuscript with contributions from RM. RM obtained funding. Both authors
- 318 read and approved the manuscript.

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- 322 Evolutionary Biology at the University of Michigan.

323 Tables

- 324 Table 1. Total carotenoid, β-carotene, and Vitamin A contents of the four tested diets. Baseline contents
- 325 were obtained from Byrne and Silla's (2017) liquid chromatography analyses of the same commercially-
- **326** available feeds used for the control diet (i.e. Diet 1).

	Total Carotenoids	β-carotene	Vitamin A
Treatment 1	0.01454 mg/g	0.005466 mg/g	0.00188 mg/g
Treatment 2	2.51454 mg/g	2.505466 mg/g	0.00188 mg/g
Treatment 3	0.01454 mg/g	0.005466 mg/g	0.50188 mg/g
Treatment 4	2.51454 mg/g	2.505466 mg/g	0.50188 mg/g

327

- 329 Table 2. Pairwise comparisons between diet treatments and survivorship categories in growth curve
- 330 parameters obtained through a Tukey-Kramer post-hoc test. Maximum size and growth rate comparisons
- 331 correspond to panels B and C of Figure 2, respectively.

	Maximum size (mm)				Growth Rate			
Comparison	Difference	Lower 2.5%	Upper 97.5%	p-value	Difference	Lower 2.5%	Upper 97.5%	p-value
D2 Surv. – D1 Surv	-0.03	-0.68	0.62	1.00	0.01	-0.04	0.06	1.00
D3 Surv. – D1 Surv	-0.37	-1.03	0.28	0.58	0.01	-0.04	0.06	1.00
D4 Surv. – D1 Surv	-0.12	-0.86	0.62	1.00	0.01	-0.05	0.06	1.00
D3 Surv. – D2 Surv	-0.34	-0.98	0.29	0.64	0.00	-0.05	0.04	1.00
D4 Surv. – D2 Surv	-0.09	-0.81	0.63	1.00	0.00	-0.06	0.05	1.00
D4 Surv. – D3 Surv	0.26	-0.46	0.98	0.93	0.00	-0.05	0.05	1.00
D3 Died – D1 Died	-1.84	-3.63	-0.05	0.04	0.05	-0.08	0.18	0.86
D4 Died – D1 Died	-2.89	-4.31	-1.48	8.34e-6	0.17	0.06	0.27	3.20e-4
D4 Died – D3 Died	-1.05	-2.47	0.36	0.26	0.11	0.01	0.21	0.026
D1 Surv. – D1 Died	0.65	-0.70	2.00	0.76	-0.02	-0.11	0.08	1.00
D2 Surv. – D1 Died	0.62	-0.72	1.96	0.79	-0.01	-0.10	0.09	1.00
D3 Surv. – D1 Died	0.28	-1.06	1.62	1.00	-0.01	-0.10	0.09	1.00
D4 Surv. – D1 Died	0.54	-0.85	1.92	0.90	-0.01	-0.11	0.09	1.00
D1 Surv. – D3 Died	2.49	1.14	3.85	4.56e-5	-0.07	-0.17	0.03	0.31
D2 Surv. – D3 Died	2.46	1.12	3.81	4.87e-5	-0.06	-0.16	0.04	0.49

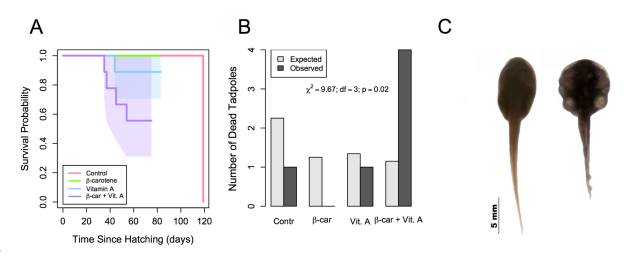
D3 Surv. – D3 Died	2.12	0.78	3.46	4.36e-4	-0.06	-0.16	0.04	0.45
D4 Surv. – D3 Died	2.38	0.99	3.76	1.37e-4	-0.06	-0.17	0.04	0.44
D1 Surv. – D4 Died	3.55	2.75	4.34	5.89e-13	-0.18	-0.24	-0.12	1.89e-9
D2 Surv. – D4 Died	3.52	2.74	4.29	4.12e-13	-0.17	-0.23	-0.12	3.80e-9
D3 Surv. – D4 Died	3.17	2.40	3.95	4.86e-12	-0.17	-0.23	-0.12	2.99e-9
D4 Surv. – D4 Died	3.43	2.58	4.28	6.64e-12	-0.18	-0.24	-0.11	1.52e-8

333	Table 3. Pairwise comparisons between	diet treatments in size an	d weight of surviving tadpoles at
	1		

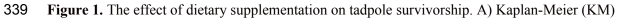
334 metamorphosis. Weight and length comparisons correspond to panels E and F of Figure 2, respectively.

	Weig	ht at Meta	morphosis	(g)	Body Length at Metamorphosis (mm)			
	Difference	Lower 2.5%	Upper 97.5%	p-value	Difference	Lower 2.5%	Upper 97.5%	p-value
D2 – D1	0.00	-0.02	0.01	0.82	-0.10	-0.53	0.33	0.93
D3 - D1	-0.02	-0.03	0.0001	0.052	-0.45	-0.88	-0.02	0.038
D4 - D1	-0.01	-0.03	0.01	0.55	-0.32	-0.80	0.17	0.31
D3 – D2	-0.01	-0.03	0.005	0.24	-0.35	-0.77	0.06	0.11
D4 - D2	-0.004	-0.02	0.01	0.94	-0.22	-0.69	0.26	0.59
D4 – D3	0.01	-0.01	0.02	0.68	0.14	-0.34	0.61	0.86

337 Figures







340 survivorship curves for each of the four treatments. Translucent polygons represent 95% confidence

341 intervals. B) Observed and expected mortality by treatment under a KM model with equal survivorship by

across treatments. C) Examples of a healthy tadpole (left), and one exhibiting the symptoms associated

343 with A hypervitaminosis (right). Photos were taken around Gosner stage 30, 24 days after eclosion.

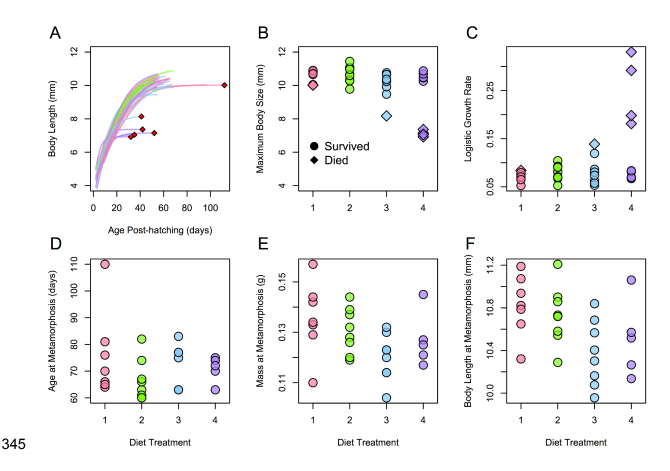


Figure 2. The effect of diet supplementation on tadpole growth. A) Best-fitting logistic growth curves for the 34tadpoles used in the experiment. Tadpoles that did not survive to metamorphosis are marked with a red diamond. B-

348 C) Best-fitting maximum size and growth rate parameters for the curves in panel A grouped by treatment and

349 survivorship status. D-F) Metamorphic age, mass, and body length of individuals that successfully survived past

350 metamorphosis, grouped by treatments. Colors in all figures match those in Figure 1.

352 Supplementary Materials

- 353 Table S1. Survivorship and growth parameters compared between tadpoles born from jellied and de-
- 354 jellied eggs. Mortality was compared using Fisher's exact test, and all other parameters were compared
- 355 using one-way anovas. OR = Odds Ratio.

356

Parameter	Test Statistic	df	p-value
Mortality	OR = 0.565	-	0.642
Maximum Size	F = 0.512	1, 32	0.479
Growth Rate	F = 1.699	1, 32	0.202
Time to Metamorphosis	F = 0.095	1,26	0.760
Mass at Metamorphosis	F = 0.714	1,26	0.406
Body Length at Metamorphosis	F = 0.664	1, 26	0.422

357

Table S2. Recipes for the four tested diets. The flake mix was a 75:25 by weight mixture of Sera Floraand Sera Sans fish flakes.

Prepared Diet					
	Flake Mix	β-carotene	Vitamin A	Cellulose	Total
Control (Diet 1)	3.988 g	0 mg	0 mg	12.0 mg	4 g
β-carotene (Diet 2)	3.988 g	10.0 mg	0 mg	2.0 mg	4 g
Vitamin A (Diet 3)	3.988 g	0 mg	2.0 mg	10 mg	4 g
β-carotene and Vitamin A (Diet 4)	3.988 g	10.0 mg	2.0 mg	0 mg	4 g

360

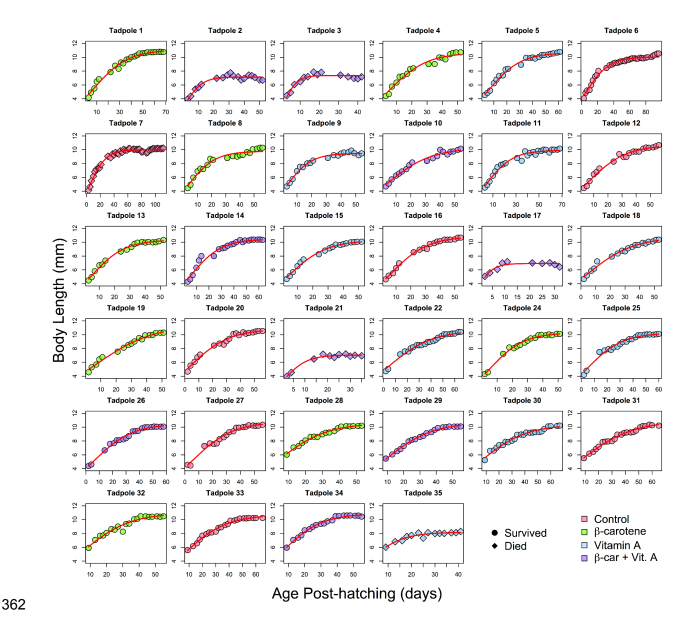


Figure S1. Individual growth trajectories for all tadpoles used in the study. Points represent rawmeasurements, and red lines are the best-fitting logistic growth curves for each individual.

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