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Research article

Impacts of UVB provision on serum vitamin D₃, pigmentation, growth rates and total body mineral content in Mallorcan midwife toad larvae (*Alytes muletensis*)

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Abstract

Our understanding of captive husbandry for amphibians is rapidly improving as empirical knowledge in this area grows. Early evidence indicates that UVB radiation is an important aspect of captive husbandry for at least some species, and may be critical in combatting nutritional metabolic bone disease and other environmentally linked diseases. However, the limited evidence in this field is restricted to post-metamorphic anurans, and impacts of UVB provision on larvae are even less well known. We measured the effects of ecologically appropriate levels of UVB exposure on growth rates, pigmentation acquisition, serum vitamin levels in the blood plasma, and whole-body mineral content in the Mallorcan midwife toad (*Alytes muletensis*). There were no significant effects of UVB exposure on any parameters measured. We were therefore unable to provide clear evidence that UVB irradiation can be used to synthesise vitamin D₃ in the skin. Our data suggest that when *A. muletensis* larvae have access to a diet containing vitamin D₃ and under husbandry conditions currently recognised as best practice, UVB irradiation is not required to maintain this species successfully. These results do not indicate that UVB provision is unimportant for amphibian larvae in general, however. Further research is needed to elucidate how tadpoles interact with UVB radiation in nature and to examine how UVB radiation is provided in captivity, and to test for effects using a wider variety of species from a range of different habitats.

Introduction

As amphibians are currently the most threatened group of vertebrates (IUCN 2018), captive breeding programmes (CBPs) are considered necessary to ensure the survival of some amphibian populations (Wren et al. 2015). Amphibians have complex and poorly understood captive husbandry requirements, however, and several CBPs have faced many obstacles, including failing to keep animals in good health, and failing to breed them (Gagliardo et al. 2008; King et al. 2011; Tapley et al. 2015a). An important aspect of amphibian husbandry is calcium provision (for a review of the importance of this see Antwis and Browne 2009). Amphibians extract

calcium from their diet and environment, and produce stores in bones and endolymphic sacks (Pilkington and Simkiss 1966), in circulating blood plasma (Stiffler 1993), and under the skin (Bentley 1984; Cheek et al. 1993).

Calcium content of prey items may be increased through dusting (Michaels et al. 2014) or gut loading (Finke 2003). However, provision of calcium alone is often insufficient for maintaining normal calcium levels. Vitamin D₃ acts to regulate the concentration of calcium in the blood plasma by facilitating active calcium uptake from the gut (for a review of this process see Finke 2002; Antwis and Browne 2009; Michaels et al. 2015). Passive uptake of calcium is also possible, but is inefficient

and may only meet requirements in species with slow growth rates and low calcium requirements (Bianchi and Shanes 1959). Vitamin D₃ can either be obtained through the diet or synthesised through cutaneous exposure to UVB radiation in combination with heat (Holick 1981; Norman 1998), or through a combination of both routes. In captivity, dietary supplements often contain both calcium and vitamin D₃. The extent of dependence on cutaneous synthesis versus dietary sources varies between taxa (Allen et al. 1995), but data concerning relative dependence on these routes is yet to be collected in amphibians. However, vitamin D₃ has been shown to increase in some anurans when exposed to UVB radiation (Michaels et al. 2015), indicating a potential role for UV lighting in captive anurans. Diseases associated with poor calcium metabolism, including metabolic bone disease (MBD), may compromise the ability of institutions to successfully maintain amphibians in captivity, including those central to the survival of threatened species (Tapley et al. 2015b). Increasing knowledge of the specific requirements of a species is therefore crucial to the success of CBPs.

In nature, amphibians are often exposed to UVB radiation, either incidentally as they move through their environment (Kats et al. 2012; DeMarchi et al. 2018) or through active basking behaviour (Michaels and Preziosi 2013). Taxa that do not spend the day in lightless environments will be exposed to some UVB radiation on a daily basis. Some authors have found deleterious effects of UVB exposure in amphibians (Blaustein et al. 1994, Grant and Licht 1995; Blaustein et al. 1998; Pandelova et al. 2006), but these are largely the result of either exposure above normal levels in nature, without a paired thermal gradient and/or no opportunity to escape exposure. Several studies in captive amphibians in the context of captive husbandry have shown beneficial effects of UVB radiation (Verschooren et al. 2011; Michaels et al. 2015; Tapley et al. 2015b) and UVB is frequently provided for captive adult amphibians without deleterious effects (C. Michaels; B Tapley pers. obs.). Aquatic amphibian larvae may be exposed to UVB radiation in nature as well as adults, but there is currently no husbandry focused work investigating the UVB requirements of this lifestage.

In this study, tadpoles of the Mallorcan midwife toad (*Alytes muletensis*) were exposed to levels of UVB radiation appropriate for its natural ecology, to measure its effects on growth and development, serum vitamin levels, and whole-body mineral content, in order to evaluate its relevance to optimising captive husbandry.

Materials and methods

Ethics statement: Methods were approved by the ZSL ethics committee (Project number ZDZ27) and deemed appropriate to carry out in the UK without a Home Office license under the Animals (Scientific Procedures) Act 1984.

Experimental animals

Alytes muletensis was selected for this study, as it is a common non-model organism used in laboratories for amphibian disease studies (Kraaijeveld-Smit et al. 2005; Garner et al. 2009) and represents a species with a long captive husbandry history (Tonge and Bloxham 1989).

All tadpoles were spawned from captive bred *A. muletensis* held at ZSL London Zoo. Thirty tadpoles from six different clutches were used, and these tadpoles were collected from the water dishes of several enclosures immediately after deposition by brooding males. Each clutch was split equally between treatment groups. The tadpoles were photographed (see methods below), and then transferred immediately into individual compartments of the experimental tank (see Figure 1).

Measurement of UVB radiation

The UV Index (UVi) was measured using a Solarmeter 6.5 (Solartech) (Lindgrin et al. 2008). The Ultraviolet index (UVi) is a unitless measure of the intensity of UV radiation from the sun or UV emitting lamp, weighted for the biological relevance of the wavelengths (Fietelov et al. 2009; Tapley et al. 2015b). Field recordings were taken by sealing the Solarmeter 6.5 in a polythene bag (Prima, Mumbai, India) to allow the unit to be submerged. The polythene bag was pulled tight over the sensor. This reduced UVi readings by 0.1 under both T5 and natural sunlight (1300hrs April 2016, clear skies, London, UK). Recordings in the experimental array were taken at surface level with and without a polythene barrier at the surface and with a polythene barrier under water.

Husbandry

Powder free nitrile gloves were always used when servicing the animals. Animals were maintained in a mixture of reverse osmosis water and tap water in a 9:1 ratio respectively. Water was dosed appropriately with tap water conditioner (API/Tetra). Water parameters were maintained at an alkalinity between 30 and 60 mg/L, and at a pH of 7.3. Temperatures were kept at 17–21°C for the duration of the study following EAZA best practice guidelines (Wells et al. 2015). Water quality was maintained using biological and mechanical filtration (Eheim Lav filter media, two Boyu sponge filters, and growth of *Egeria densa* –filtration was not present in the actual experimental compartments; see below) and routine 30% daily water changes as well as detritus siphoning. Filter media was cleaned in tank water as necessary. Ammonia and nitrite were generally maintained at <0.05 mg/L and nitrate at <100 mg/L. Water parameters were monitored weekly using an aquarium test kit (Salifert Nitrate test kit; Nitrate), and a spectrophotometer (Interface photometer 7500, Palintest). Tadpoles were fed on a homogenised diet comprising of 10 parts fish flake, eight parts trout pellet, eight parts grass pellets, three parts cuttlefish bone, two parts dried tubifex, two parts dried river shrimp, and three parts spirulina in one batch (parts measured by weight). Brands varied between batches according to availability, but all tadpoles were fed on same diet at any given time. Food was offered as a sinking pellet to avoid surface interruption of UVB radiation decreasing the amount of UVB reaching the tadpoles. Tadpoles were also able to feed on surface algae within the tank at all times. We did not control for food intake, as in a practical husbandry setting, food is provided ad libitum and is not a limiting factor for growth.

Experimental array

The experimental array comprised of a glass aquarium split into 30 compartments that each housed a single tadpole (Figure 1). A small gap beneath each divide allowed water flow throughout the system whilst also preventing tadpoles from moving between compartments. Two larger compartments, one at each end of the tank, held the filter media and water was transferred through the tank using a small pump (Eheim pump compact+ 3000) at one end, with silicone tubing transporting water to the other end compartment through a spray bar. The tank was illuminated with five 12% T5 Arcadia UVB emitting lamps across each row of compartments, with half of each lamp alternately covered with a UVB filter (CLS200 XSR Clear Ultra Violet Filter) to block UVB radiation whilst still allowing passage of other wavelengths (Michaels et al. 2015). This resulted in 15 compartments being exposed to UVB (UVB+) with the other 15 not (UVB-). Irradiation period was set at four hours on, 20 hours off as *A. muletensis* are often found in steep gorges in mountainous areas (Buley and Garcia 1997), and so direct sunlight would only have been received by most remaining populations for a short period of the day as the sun passes overhead. The array received ambient lighting

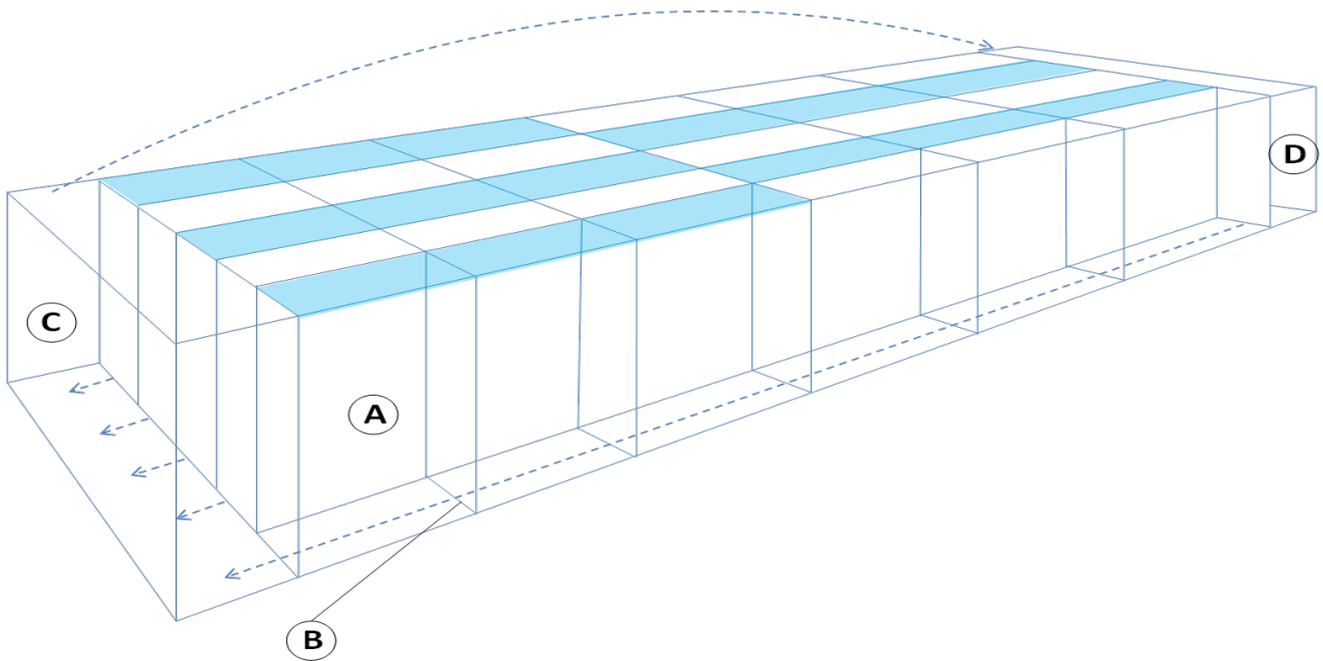


Figure 1. Experimental array: constituting a glass tank (measuring 118 cm x 40 cm x 20 cm) divided into thirty compartments (5 rows and 6 columns each measuring 7.5cm x 17cm x 20cm and filled to a water depth of 17 cm) (A), each housing one tadpole complete with a refugium, and two end compartments (measuring 8cm x 40cm x 20cm) (C and D), each filled with filter media (not shown). Rows of compartments were linked by a small gap underneath each glass division (B). A water pump (not shown) was contained within the left-hand end compartment (C) and established circulation through filter media by pumping water to the other end compartment (D), from which it flowed back (blue dashed arrows) along the tank via the gap underneath each compartment wall (B).

from the room on a 12:12 photoperiod from a non UVB emitting array. Lamps above the experimental array were held in place using tape to prevent movement, and the UVB emitted by these lamps was measured every month from a distance equal to that of halfway through the water column, and lamps were replaced four months into the seven month experiment due to decreased UVB emissions. Each individual compartment was equipped with a refugium comprising an eight cm piece of food grade PVC piping (one-inch diameter; GF Harvel) to allow the tadpole to shelter from either UVB radiation (in UVB+ compartments), or from any perceived danger.

UVB+ compartments were irradiated with a maximum UVI of 2-3.7 (1.9-3.6 with the polythene barrier). Using a deeper chamber to allow for the height of the solarmeter, UVB+ chambers had a minimum UVI of c. 1 on the substrate at the bottom of the compartment. A zone of UVI 0 was available inside the PVC tube hide. These UVI are broadly concordant with a UVI of c. 2.4 collected underwater (at a range of depths) at an *A. muletensis* breeding site at Coco de Bovas, Mallorca in July 2017 (J. Bosch pers. comm.). UVB- compartments received a UVI of 0, which was achieved through the placement of lamps, reflectors and glass panels and was confirmed using the Solarmeter 6.5. T5 fluorescent tubes produce markedly less UVB radiation towards the ends of the tubes; the array and light selection was designed to avoid any compartments being under the lower-UVB parts of the lamps; this was confirmed using the Solarmeter 6.5.

Data collection

Tadpoles were photographed next to a scale at the beginning of the experiment and weekly thereafter. Each tadpole was removed from its enclosure using a net and photographed in a moist petri dish against the same scale, before being returned to its compartment. The Gosner stage of development (Gosner 1960) of the tadpole was recorded weekly. Photographs were analysed using ImageJ (1.48v, NIH). Tail length (from tail tip to vent), head length (snout to vent length), and total tadpole length (sum of tail length and head length (McDiarmig and Altig 1999) was measured using this software. The tail:body ratio of the tadpoles was also calculated. Each tadpole was given a colour score from one to five with one being the lightest colour, and five being the darkest. This was to determine if UVB had any effect on colouration (Blaustein and Belden 2003). Non-experimental larvae were used as standards of each colour category and experimental larvae were matched to the closest standard by eye.

Once the tadpoles had reached Gosner stage 39 (their maximum length before shrinking during metamorphosis), or at seven months post introduction to the experimental array (whichever came sooner), they were euthanased using a buffered methocaine trisulphate (MS222) solution. A blood sample was taken from the heart of each tadpole immediately after euthanasia. This was done by making an incision in the ventral skin of the tadpole to expose the heart, removing any excess interstitial fluid that may dilute the blood sample using a cotton bud, puncturing the heart using a

Table 1. Statistical outcomes of growth data and colouration data analysis

Measured value	Test statistic and degrees of freedom	UV treatment*age interaction effect
Total length	$F_{1,28} = 0.33$	0.5711
Tail length	$F_{1,27} = 0.1$	0.7507
Head length	$F_{1,26} = 1.3$	0.2644
Head to tail length ratio	$F_{1,20} = 0.375$	0.55
Colouration score	$F_1 = 0.385$	0.755

scalpel, and extracting the blood that filled the body cavity using a heparinised capillary tube (Vitrex Medical BRIS microhematocrit tubes). Blood drops were then dropped onto test blotting strips obtained from City Assays (City Hospital, Birmingham, Sandwell and West Birmingham Hospital NHS trust; www.vitaminDtest.org.uk). These were returned to this laboratory to measure 25-hydroxy vitamin D₃ by tandem mass spectrometry after derivatisation and liquid-liquid extraction, using in house calibrators and quality control material (Jaffe et al., 2019). The bodies of these tadpoles were then stored in Eppendorf tubes (50 ml capacity) at -20°C and sent for calcium and phosphorus analysis using inductively coupled plasma atomic emission spectroscopy (ICP-AES) (Michaels et al. 2014) at Manchester Metropolitan University.

Mineral analysis was carried out at Manchester Metropolitan University. Frozen tadpoles were dried out at 60°C in a GenLab incubator for 72 hours in a 50 ml glass vial, then washed with ultrapure water. A dry weight of all tadpoles was taken. The tadpoles were then transferred to digestion vessels and two ml Hydrogen peroxide (>30% w/v) was added to break down the organic matter. After 30 minutes, 8 ml of nitric acid (>68% PrimerPlus- trace analysis grade) was added and left to digest for two hours. The vessels (plus six blanks) were then transferred to the microwave (CEM Mars Xpress 5) and run for five minutes at 90°C, then 10 minutes at 170°C. This was then repeated to ensure digestion was complete. The volume of each sample was then made up to 50ml with distilled water, and 13 ml of each sample was added to 15 ml centrifuge tubes (Fisherbrand) for the ICP-AES analysis (Thermo scientific iCAP6300 Duo). Five repeats were run for each sample, and an average was taken to be used for analysis. Levels of calcium and phosphorus were measured.

Statistical Analysis

All statistics carried out used a significance threshold of P=0.05. From 10 randomly chosen images, the total length of each tadpole was measured three times using ImageJ (1.48v, NIH). Between each measurement, the scale was recalibrated, and repeatability (r) for each measurement was calculated using the following formula:

$r = \frac{S_{(2A)}}{S_{(2 + S_{2A})}}$ where S_{2A} is the among-groups variance and S₂ is the within-groups variance (Lessels and Boag, 1987; Verschooren et al. 2011; Michaels et al. 2015).

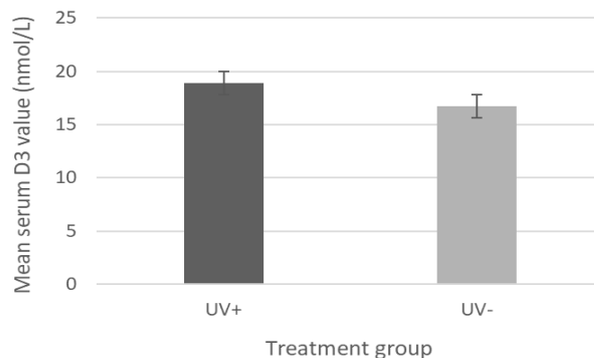


Figure 2. Serum D₃ levels: Average levels of vitamin D₃ taken from tadpole blood samples for both treatment groups

Growth data and colouration

Data were plotted as histograms and no deviation from normal distribution was noted. Plotting residuals against the data showed no evidence of heterogeneity of variance, therefore parametric statistics were deemed appropriate for analysis. A longitudinal linear mixed effects model was used to assess the effects of UV treatment on the rate of tadpole growth over time. Fixed effects included UV treatment, and linear and quadratic polynomials of age and the interactions of UV treatment and two age polynomials to control for repeated measures. In addition to the random clutch effect, intercept, linear and quadratic polynomials of age for the tadpole were also modelled as random effects using unstructured covariance type. Analysis was carried out using Statistics software R (3.4.0) and the package <lme4>. A generalized estimating equation using a cumulative logit link function was used to assess the effect of UV treatment, age, and their interactions on the tadpole colouration. The order in which tadpoles was added to the experimental array was accounted for. First-degree autoregressive working correlation structure was used to account for the repeated measures from the same tadpole. This analysis was done using SPSS (Version 23).

Vitamin D₃ levels in blood serum

Serum was successfully recovered and tested for 25 tadpoles, 13 from UVB+ and 12 from UVB-. A Mann Whitney U test (two tailed) was carried out on serum 25(OH)D₃ levels for both treatment groups after anomalous results were omitted. Two test runs, each representing the mean average of four tests are conducted per sample. Three individual results were omitted as they were several times higher than the other repeats within their test runs. Therefore, data were available from all 25 animals from which samples were recovered. This analysis was done using Graphpad Prism (7.03).

ICP-AES mineral content analysis

Calibration curves were calculated for each mineral. All calibration curves had a correlation of 0.999. A Mann Whitney U test (two tailed) was carried out on both calcium and phosphorus content of all tadpoles from both treatment groups. This analysis was done using Graphpad Prism (7.03).

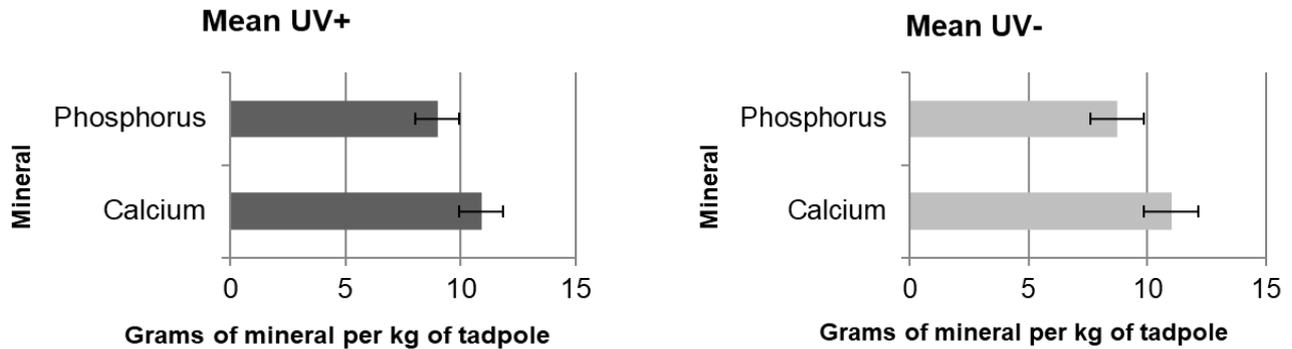


Figure 3. Mineral content: The average calcium and phosphorus content of tadpoles from the two treatment groups

Results

Repeatability analysis

All measures taken were found to be highly repeatable with the lowest r value being 0.96.

Growth data and colouration

As the effects of the linear polynomial of tadpole age were significant, this was used to control for repeated measures and so the quadratic polynomial was not used. We report P values for the UVB treatment*age interaction effect, which account for repeated measures. The order in which tadpoles were added to the experimental array had no significant effect on any results ($P > 0.05$ for all measures). There was no effect of UV treatment on any growth parameters or coloration score measured in this study (Table 1).

Vitamin levels in blood serum

There was no significant difference in serum vitamin D_3 results between treatment groups ($U_{12}=59$, $P=0.3134$) (Figure 2).

ICP-AES mineral content analysis

There was no significant difference in calcium content ($U_{14}=85$, $P=0.5860$), or phosphorus content ($U_{14}=76$, $P=0.3389$) (Figure 3). The average calcium to phosphorus ratio for the UVB+ treatment was 1.20:1, and the average calcium to phosphorus ratio for the UVB- group was 1.26:1 respectively.

Discussion

Michaels et al. (2015) found that captive post-metamorphic fire-bellied toads (*Bombina orientalis*) that were exposed to ecologically and physiologically relevant levels of UVB had significantly, although only slightly, more robust skeletal development, a significantly increased rate of adult colour acquisition, and a significantly increased level of serum vitamin D_3 than animals that were not exposed. Verschooren et al. (2011) also showed that UVB exposure in recently metamorphosed treefrogs (*Trachycephalus resinifictrix*) increased growth and development of skeletal structures. These species differ greatly in their ecology, with *T. resinifictrix* inhabiting the canopy of tropical

rainforests (La Marca et al. 2010) and *B. orientalis* inhabiting a wide range of water bodies within lowland temperate and subtropical forests (Kuzmin et al. 2004), as well as their phylogeny. However, both species actively bask in sunlight or artificial light sources given the opportunity. As these two studies (the only two experimental studies using controls so far focusing on amphibian UVB provision from a husbandry perspective) have demonstrated that UVB provision is arguably beneficial in both these species of basking anuran from otherwise differing ecologies, it is reasonable to assume that at least anurans that exhibit basking behaviour are likely to have a physiological requirement for UVB radiation. These studies only focus on UVB provision post-metamorphosis, and so the present study is the first to focus on provision of quasi-natural levels of UVB in larval amphibians from the perspective of captive husbandry.

We detected $25(OH)D_3$ in the serum of tadpoles in this study. This suggests that this species may use vitamin D_3 in active absorption of calcium from the gut, and that it does not rely solely on passive absorption. This underpins the importance of providing at least a diet containing vitamin D_3 . Wild *A. muletensis* larvae are deposited in large, warm, shallow water bodies from May to September, where they develop and metamorphose in the autumn or following spring (Pinya and Pérez-Mellado 2014). Developing larvae are exposed to UVB and are active during the day (Spence 2002) when ambient UVB levels are higher. Although *A. muletensis* tadpoles appear to avoid the most intense sun exposure by hiding beneath the overhangs of rocky pools (Schley et al. 1998), even in these microhabitats tadpoles are exposed to a level of UVB similar to those used in this study (J. Bosch pers. comm.; see Methods). It is therefore surprising that we found no effect of UVB exposure on blood serum $25(OH)D_3$ levels, growth rates, or colouration of the tadpoles, as it would be reasonable to expect an organism to utilise a free source of energy such as UVB radiation from the sun (Michaels and Preziosi 2013). It is important to note that comparisons with serum $25(OH)D_3$ levels in wild individuals would inform the establishment of a normal baseline. Without these data it is unclear if tadpoles in this study were exhibiting normal, equally high, or equally low concentrations of serum vitamin D_3 (although no deleterious effects were noticed either way). However, no clinical signs of hypovitaminosis D_3 were noted.

The absence of a statistically significant difference in vitamin D₃ between treatments may indicate that larval *A. muletensis* are incapable of photobiosynthesis of vitamin D₃ and instead rely entirely upon dietary sources. However, there are several alternative explanations for our data, which mean that photobiosynthesis cannot be ruled out. The increase of 25(OH)D₃ in the serum does not rise in direct proportion to vitamin D₃ intake or synthesis, but slows under homeostatic control once optimal levels are reached (e.g. Heaney et al. 2008). In our animals, excess D₃ may have been stored in tissues, excreted through kidneys or destroyed by continued exposure to UVB radiation in the skin if carrier proteins were not available to transport it into tissues. It is therefore possible that the dietary vitamin D₃ present in the food, which could not be eliminated for ethical and legislative reasons, provided sufficient vitamin D₃ to saturate serum levels of 25(OH)D₃. Figure 2 shows a non-significant increase in 25(OH)D₃ levels, which may be indicative of this dynamic. We were not able to include treatments with no UVB irradiation or dietary vitamin D₃ in order to address this issue, as this would have fallen into the remit of the Animals (Scientific Procedures) Act 1986 in UK legislation, which would require licenses and facilities beyond the scope of this study.

Photobiosynthesis of vitamin D₃ in the skin requires both UVB irradiation and appropriate temperatures to enable biochemical reactions (see Michaels et al. 2015). Although the temperatures used in this study fall within the c. 17-24°C range recorded in a study of free-living tadpoles of this species (Schley et al. 1998), and within the range determined as preferred temperature and optimum for growth (Kadel and Hemmer 1984; Martens 1984) larvae in this study did not have access to a temperature gradient and therefore may not have become sufficiently warm to utilise UVB radiation in the synthesis of vitamin D₃. The duration of UVB irradiation per day was relatively short. Tadpoles of this species often occur in rocky gorges, where direct insolation may be limited to a short period. However, light scatter from gorge sides may extend UVB irradiation in nature beyond the period of direct insolation (Caldwell et al 1980). Free living tadpoles modulate their use of deeper and shallower water and rock overhangs throughout the day, and aggregate under rock shelves in deeper water during periods of most intense sunlight (Schley et al. 1998). This means that the parameters used in the experimental array may have mimicked larval exposure to UVB and temperature in the field reasonably well, with a short period of UVB irradiance and relatively constant temperatures, but further studies looking at UVB exposure in wild tadpoles over time are required to inform future work. The inclusion of thermal as well as UVB gradients in further experiments would also be useful.

The experimental array consisted of a shared water body between all 30 tadpoles. This allowed maintenance of stable water quality, which would not have been possible if tadpoles were housed separately. Consistently good water quality is impossible to maintain in small, isolated unfiltered containers, which require frequent 100% water changes and results in fluctuation of parameters including nitrogenous waste, bacterial load and pH. Poor and fluctuating water quality causes stress, which influences growth and development rates in amphibians in a complex fashion (e.g. Denver 2009). As stocking density has been shown to influence growth rates (Dash and Hota 1980; Wilbur 1997), the order in which the tadpoles were added to the array was statistically controlled for as a proxy for stocking density, in addition to the systematic paired addition of animals under both treatments.

There are few available data regarding serum vitamin D₃ levels in anurans (Michaels et al. 2015), and the dry blood spot method of quantifying this has not been validated for amphibians (or any other non-human animal), and in the only non-human

animal where validation has been attempted (corncrakes; Jaffe et al., 2019), calibration was not favourable enough to permit interchangeability of methods. In our dataset, there was some variation between repeats, and some repeats were omitted due to substantially different results. Some blood spots developed uneven serum distribution on the blotting paper, with a 'halo' developing around the central spot. This has also been noted in *Bombina orientalis* (Michaels et al., 2014) and is suggested as a potential cause of variation (N Barlow, pers. comm.). Therefore, these data are used tentatively and, although they are more likely to be comparable within a study, they may not be comparable with data from other species (Michaels et al. 2015; Jaffe et al., 2019). However, for small animals, where analysis of liquid serum is not possible due to minimum sample volume requirements, the dry blood spot approach is the only viable method available,

Bearing these caveats in mind, Michaels et al. (2015) found a range of serum levels ranging from 73.59–88.68 nM/L across UV- and UV+ treatment groups. However, the raw data collected in our study ranged from 3.8–52.5 nM/L across both treatment groups. This could result from differing physiological requirements, different relative impacts of captivity on each species, methodological error or a host of other reasons. Methodological validation and the acquisition of a larger data set from a variety of species is needed in order to better understand the implications of these data.

Whole body mineral analysis, including calcium and phosphorus, of adult Cuban tree frogs (*Osteopilus septentrionalis*) was carried out by Allen et al. (1993). Frogs were fed on either a high or a low calcium diet (no UVB exposure provided), and results showed much higher calcium:phosphorus ratios (approximately 2:1 in females and approximately 2.5:1 in males for both treatment groups) than were found in this study (approximately 1.2:1 in the UV+ group and 1.26:1 in the UV- group). This is probably because metamorphosed frogs with ossified skeletons will contain more mineralised calcium within these structures, compared to the tadpoles in this study which lack mineralised skeletons. However, figures from this study meet the recommended requirement of a 1:1 or 2:1 ratio of calcium to phosphorus in the diet respectively (Finke 2002), indicating that no signs of ill health would be expected as a result of this, and that the levels of 25(OH)D₃ that we detected are likely to be sufficient for normal calcium homeostasis.

Increased pigmentation with melanin may be used by amphibians as protection against harmful doses of UVB radiation (Brenner and Hearing 2008). However, we found no significant difference in pigmentation score between treatments, and neither did we observe any negative effects of UVB exposure (as found in other studies; Worrest and Kimmeldorf 1976; Grant and Licht 1995; Blaustein et al. 1998, 2003, 2005). Three tadpoles from the UVB negative group died of natural causes unrelated to UVB provision (confirmed by post mortem examination). No animals from the UVB+ group died during this study. This is unsurprising, as tadpoles were provided with an ecologically appropriate level of UVB with both a gradient, and refugia to allow exposure to be avoided if chosen. Whilst the tadpoles used the refugia provided, there seemed to be no increased usage amongst UVB positive animals, and many times animals from both treatment groups were observed swimming at the top of the water column where UVB exposure would have been greatest. The numbers of tadpole deaths were too small for statistical analysis.

Overall, our results indicate that the integration of ecologically appropriate UVB radiation into existing husbandry protocols for larval *A. muletensis* had no effect on measured parameters while vitamin D₃ was available in the diet. Although this suggests that existing husbandry practices which omit UVB provision may be appropriate for this species, this alone is not sufficient to inform general recommendations regarding UVB provision for captive

anuran tadpoles. Further work exploring the impacts of diet, thermal gradients, and UVB dosage and exposure length may be required to fully understand UVB requirements in *A. muletensis* larvae. Moreover, UVB requirements may vary between taxa depending on their evolutionary and ecological context (Michaels et al. 2014).

This study focuses on UVB provision for calcium homeostasis, but irradiation may have other important effects. For example, UVB exposure has also been shown to kill bacteria found on human skin (Jekler et al. 1992; Fluhr and Gloor 1997; Yoshimura-Mishima et al. 1999). Should the same principles apply to amphibians, then UVB radiation may have effects on host-pathogen interactions, both by directly inhibiting bacterial growth on the skin, and through interactions with bacterial floras that may mediate infection. UVB provision could therefore potentially be used to help reduce the incidence of disease in CBPs, and its provision may still be useful for rearing tadpoles in capacities unrelated to calcium metabolism. This study provides a basis upon which further research methods into tadpole UV provision studies can be refined and improved to ultimately determine the role of UV provision for anuran larvae in captivity.

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