Are the adverse effects of stressors on amphibians mediated by their effects on stress hormones?

Caitlin R. Gabor · Sarah A. Knutie · Elizabeth A. Roznik · Jason R. Rohr

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Abstract

Adverse effects of anthropogenic changes on biodiversity might be mediated by their impacts on the stress response of organisms. To test this hypothesis, we crossed exposure to metyrapone, a synthesis inhibitor of the stress hormone corticosterone, with exposure to the herbicide atrazine and the fungal pathogen *Batrachochytrium dendrobatidis* (*Bd*) to assess whether the effects of these stressors on tadpoles and post-metamorphic frogs were mediated by corticosterone. Metyrapone countered atrazine- and *Bd*-induced corticosterone elevations. However, atrazine- and *Bd*-induced reductions in body size were not mediated by corticosterone because they persisted despite metyrapone exposure. Atrazine lowered *Bd* abundance without metyrapone but increased *Bd* abundance with metyrapone for tadpoles and frogs. In contrast, atrazine reduced tolerance of *Bd* infections because frogs exposed to atrazine as tadpoles had reduced growth with *Bd* compared to solvent controls; this effect was not countered by metyrapone. Our results suggest that the adverse effects of atrazine and *Bd* on amphibian growth, development, and tolerance of infection are not mediated primarily by corticosterone. A possible mechanism for these effects is energy lost from atrazine detoxification, defense against *Bd*, or repair from damage caused by atrazine and *Bd*. Additional studies are needed to evaluate how often the effects of anthropogenic stressors are mediated by stress hormones.

Keywords Atrazine · *Batrachochytrium dendrobatidis* · Chytrid · Contaminants · Pathogen

Introduction

We are now in the age of the Anthropocene, a time when human activity is the dominant influence on the environment and biodiversity (Dirzo et al. 2014). Many anthropogenic factors, such as chemical contaminants and introduced pathogens, can function as stressors by elevating or dysregulating glucocorticoid “stress hormones” in vertebrates (Gabor et al. 2015; Larson et al. 1998; McMahon et al. 2017). In turn, these interactions can have profound and enduring effects on the health of organisms, especially when exposure occurs during early life stages (Boekelheide et al. 2012; Martin et al. 2010; Rohr et al. 2013). For example, early life exposure to stress hormones during key developmental stages can permanently alter the functionality of the hypothalamic–pituitary–adrenal (HPA; HPI—interrenal in amphibians) axis, which in turn can alter the immune system into adulthood (Martin et al. 2010; Matthews 2002; Rohr et al. 2013). Thus, even though many physiological responses to stress can be adaptive (Boonstra 2013), many of the adverse effects of the Anthropocene on biodiversity might be mediated by the stress physiology of organisms. However, this hypothesis has
not been thoroughly tested because no studies to date have crossed anthropogenic stressors with compounds that inhibit the synthesis of stress hormones. Such a study would help determine whether stress hormones are largely responsible for adverse effects of anthropogenic stress.

Amphibians are the most threatened class of vertebrates on the planet (Stuart et al. 2004) and represent a taxon extensively impacted by activities dominating the Anthropocene. Anthropogenic factors have been implicated in amphibian declines, including environmental pollutants, infectious diseases, and their interactions (Hayes et al. 2010; Jones et al. 2017; Rohr and McCoy 2010). For instance, chytridiomycosis, the disease caused by the fungal pathogen, *Batrachochytrium dendrobatidis* (*Bd*), has caused major declines and possibly extinctions of hundreds of amphibian species in the last half century (Wake and Vredenburg 2008). Amphibians employ two defense strategies against *Bd* that can be impacted by anthropogenic factors and associated stress hormones. Resistance strategies, such as innate and adaptive immune responses (McMahon et al. 2014; Rollins-Smith et al. 2011), prevent or clear *Bd* infections, whereas tolerance strategies minimize the fitness consequences of infection, such as mechanisms that enhance repair from parasite damage (Råberg et al. 2009; Rohr et al. 2010).

The immune system in vertebrates is partly modulated by glucocorticoids, such as corticosterone, the main glucocorticoid related to stress in amphibians. Corticosterone and other stress hormones, such as cortisol, are regularly used to assess overall stress and health of wild animal populations and to direct wildlife management (reviewed by Busch and Hayward 2009; Sheriff et al. 2011). For example, chronically elevated corticosterone can accelerate metamorphosis and decrease amphibian growth, development, and immunity, the latter of which can increase infectious disease risk (Denver 2009; Rollins-Smith 1998; Warne et al. 2011). Chemical contaminants, such as the herbicide atrazine, the second most commonly used pesticide in the US (Kiely et al. 2004), and pathogens, such as *Bd*, can elevate corticosterone (Gabor et al. 2015; McMahon et al. 2017; Peterson et al. 2013; Searle et al. 2014). Contaminants have similar negative effects on amphibians as does chronic corticosterone, such as reduced immunity and growth (Larson et al. 1998; McMahon et al. 2013a; Rohr et al. 2013). This suggests that the negative effects of contaminants might be mediated by corticosterone. For example, early life exposure to atrazine reduces amphibian growth and development and, although it does not affect amphibian resistance to *Bd*, it reduces tolerance of *Bd* infections, thus increasing *Bd*-induced mortality (Rohr et al. 2013).

Using a series of experiments where we inhibited corticosterone synthesis in Cuban tree frog (*Osteopilus septentrionalis*) using the compound metyrapone, we explore whether the effects of atrazine and *Bd* on amphibian growth and development, and the effects of atrazine on amphibian resistance and tolerance of *Bd* were mediated by their effects on amphibian corticosterone. We hypothesized that any effects of atrazine and *Bd* exposure on growth, development, and survival would be at least partly mediated by their effects on corticosterone. We also hypothesized that atrazine would affect amphibian-*Bd* interactions by altering amphibian resistance and/or tolerance of *Bd* and that this too would be at least partly mediated by corticosterone. Importantly, if corticosterone mediates any of these effects, then corticosterone levels should be correlated with these responses and metyrapone should counteract some or all of the effects of atrazine and *Bd*.

### Materials and methods

We collected multiple clutches of tadpoles of *O. septentrionalis* in August 2014 from the Botanical Gardens of the University of South Florida (N 28°03.537’ W 082°25.410’). We maintained them in the lab for at least a week until the majority reached Gosner developmental stage 35 (Gosner 1960). All tadpoles were fed a mixture of fish food and *Spiroulina* suspended in agar ad libitum and were maintained at 21 °C with a 12 h light cycle.

### Baseline corticosterone and stress responses

To determine the physiological range of corticosterone in *O. septentrionalis* tadpoles, we quantified the release rate of water-borne corticosterone from undisturbed (“baseline”) tadpoles (*n* = 18) and we obtained the stress response from naturally stressed tadpoles (*n* = 15) through gentle “agitation”. Given that these methods are already well documented (Gabor et al. 2016), these methods are presented in the Supplemental material.

### Experimental overview

First, tadpoles were exposed to fully crossed atrazine and metyrapone (corticosterone synthesis inhibitor) treatments for 6 days (Fig. 1). Metyrapone has been used in other studies with amphibians to explore the effects of corticosterone on stress responses (Glennemeier and Denver 2002a, b), specifically the stress response to predation (Middlemis Maher et al. 2013; Neuman-Lee et al. 2015), and has not been found to change amphibian behavior (Hossie et al. 2010). Second, after placing tadpoles in fresh untreated water, we challenged half the amphibians with *Bd* or not for 14 days. We obtained water-borne corticosterone release rates from tadpoles after the 6 days of atrazine exposure and again after a week of *Bd* exposure. We swabbed tadpoles for *Bd* on day 21. Third, we challenged the other half of tadpoles,
Once they were post-metamorphic frogs, with Bd or not. We swabbed post-metamorphic frogs on day 98 and obtained mass weekly. We noted tadpole and post-metamorphic frog survival daily. On day 119 we recorded final survival of post-metamorphic frogs.

**Atrazine and metyrapone exposure in tadpoles**

On the first day of our experiment, we filled 41 12-L tanks with 8 L of water from a pond at Trout Creek Park, FL (N 28°09′22.50″, W 082°34′8.083″) that was free of tadpoles and was not exposed to agricultural runoff (i.e., no measurable level of atrazine, see atrazine measurements below). We assigned 16 *O. septentrionalis* tadpoles haphazardly to each tank. We randomly assigned each tank to one of four exposure treatments: (1) the estimated environmental concentration (EEC) of atrazine (200 µg L\(^{-1}\); Chemservice, West Chester, PA; technical grade, purity more than 98%) dissolved in 120 µL of ethanol \((n = 9)\), (2) 110 µM of metyrapone (Sigma Chemical Co. # M2696; St. Louis, MO, USA) dissolved in 120 µL of ethanol \((n = 11)\), (3) the EEC of atrazine and 110 µM of metyrapone jointly dissolved in 120 µL of ethanol \((n = 11)\), and (4) only 120 µL of ethanol as a control \((n = 10)\). We used 110 µM of metyrapone because this level reduced whole body corticosterone in tadpoles by >50% (Glennemeier and Denver 2002b) and, over the short term (weeks), exposure is believed to be non-toxic (Glennemeier and Denver 2002c). Previous work did not detect effects of ethanol (a solvent) on any measured trait, and thus a water control was not included (reviewed by Rohr et al. 2013). Tadpoles were exposed to these treatments for 6 days.

The targeted nominal concentration of atrazine was 200 µg L\(^{-1}\), the EEC based on US Environmental Protection Agency GENEEC v2 software. The EEC is the concentration estimated to enter a standardized farm pond at a standardized distance from an application site given the chemical properties of the pesticide; thus, it is an ecologically relevant concentration. To quantify actual atrazine concentrations, water samples were taken from each of the 40 tanks 1 h after dosing and atrazine was quantified using the Abraxis ELISA microtiter plate kit (Abraxis LLC, Warminster, PA, USA). Mean (±1 SE) atrazine concentration was 178.2 ± 7.8 µg L\(^{-1}\). All atrazine values for the

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**Fig. 1** Experimental design for exposing tadpoles of *Osteopilus septentrionalis* to atrazine and metyrapone (corticosterone synthesis inhibitor). On day 6, water-borne corticosterone (CORT) was obtained and then moved to new tanks where half were exposed to *Batrachochytrium dendrobatidis* (Bd) and the other half were not. Water-borne CORT was obtained on days 13 and 14. On day 21, remaining tadpoles were swabbed for Bd, and we obtained final mass, snout–vent length, and Gosner stage. Half of the tadpoles from the first chemical exposure were reared in their original tanks to post-metamorphic frogs (day 84), and were then exposed to Bd (or not) and weighed weekly. On day 89, frogs were swabbed for Bd. Survival was recorded up through day 119. Numbers represent sample sizes (i.e., number of individuals/number of tanks) for each time point. Photos by Caitlin Gabor and Betsy Roznik.
non-atrazine-exposed tanks were below the detection limit of 0.06 µg L⁻¹ (this is the level in the pond water). We re-dosed each tank with 110 µM of metyrapone every third day (following Hossie et al. 2010). We did not re-dose with atrazine because its half-life is on the order of weeks and Rohr et al. (2004) found no detectable breakdown of atrazine over 7 days under similar conditions.

After tadpoles were exposed to their respective treatment, we obtained water-borne corticosterone (6 days corticosterone) from two tadpoles per replicate (80 total) by placing tadpoles individually in 250 ml beakers filled with 75 mL of clean water for 1 h (following Gabor et al. 2016). We then removed them and measured their mass and snout–vent length (SVL). Gosner stage was not measured because the same tadpoles were used throughout the experiment but we noted that none had obvious limb buds. Water samples were frozen at −20 °C immediately after collection until ready to be thawed for extraction; this method does not affect corticosterone levels (Ellis et al. 2004).

**Bd exposure in tadpoles**

Immediately following the collection of corticosterone on day 6, eight tadpoles were removed from their original tank and were evenly divided between two 6-L plastic shoeboxes with 2 L of fresh pond water for the *Bd* exposure stage of the experiment. However, some tadpoles were misplaced during this transfer stage so we lost some replicates (Fig. 1). *Bd* isolate SRS812 was cultured following the methods of McMahon et al. (2013a). Half of these shoeboxes received a 6-mL inoculum containing 7 × 10⁴ *Bd* zoospores mL⁻¹ in deionized (DI) water and the other half received an inoculum that was identical to the *Bd* inoculum but was free of *Bd* (i.e., we washed clean agar plates with DI water). We re-exposed all tadpoles to *Bd* (2 mL of 3 × 10⁴ zoospores mL⁻¹) or DI water 3 days later and maintained the tadpoles in these boxes until day 21.

On day 13 and 14 (7 and 8 days after exposure to *Bd*), we collected water-borne corticosterone samples (13, 14 days corticosterone) from two tadpoles of each replicate to examine the effect of *Bd* exposure on individual corticosterone release rates (half the tadpoles each day using the same methods as described above). All other tadpoles were placed in the same size containers with the same amount of water to control for any effects of handling. All tadpoles were returned to their original tank after being placed in individual beakers. Twenty-one days after exposure to *Bd* or not, we swabbed each surviving tadpole for *Bd* by passing a sterile rayon swab along its mouthparts (eight strokes horizontally, and eight strokes vertically). We then euthanized tadpoles with an overdose of MS-222. We recorded their mass, SVL, and Gosner stage, and preserved them in 70% ethanol. We used quantitative PCR (described by Boyle et al. 2004) to quantify *Bd* abundance taken from up to two tadpoles per *Bd*-exposed tank, and a total of 10 tadpoles (each from separate tanks) that were not exposed to *Bd*.

**Bd exposure in post-metamorphic frogs**

Individuals that were not used in the *Bd* experiment as tadpoles remained in their original tanks until they metamorphosed. After the 6-day chemical treatment, freshwater, free of atrazine, was provided every other week. Tadpole survival and day of metamorphosis (all four limbs had emerged; Gosner stage 42) were recorded daily. Upon metamorphosis, individuals were removed from the tanks and placed in cups (6 cm high × 12 cm diameter) with moist *Sphagnum* sp. moss. The post-metamorphic frogs were maintained in the laboratory (12 h light cycle, 22 °C) and fed ad libitum vitamin- and mineral-dusted crickets twice per week. Frog survival was recorded daily. A week prior to *Bd* exposure, body mass was recorded for each individual. On day 84 of the experiment and approximately 1 month after most of the tadpoles metamorphosed, half the surviving post-metamorphic frogs from each tank were randomly assigned to receive *Bd* (isolate SRS812) and the other half received the control (each tank had 1–2 frogs exposed to each treatment depending on survival). Post-metamorphic frogs were exposed to *Bd* or a control solution (everything but *Bd*) by pipetting 1 mL of 6 × 10⁴ zoospores mL⁻¹ onto the frog’s dorsal side. Excess inoculum remained in each frog’s plastic container, which contained moist sterile *Sphagnum* moss. Survival was monitored daily for 5 weeks. Frogs were also weighed weekly and swabbed (entire ventral surface and all four feet twice) at 2 and 3 weeks after *Bd* exposure. *Bd* from the swabs was quantified using the methods described above. We then euthanized individuals with an overdose of MS-222 and preserved them in 70% ethanol for further processing.

**Hormone extraction and validation**

We obtained all water-borne hormone samples between 0830–1500 h. We extracted water-borne hormones following (Gabor et al. 2016). We re-suspended the dried hormone residue in 260 µL enzyme-immunoassay (EIA) buffer (provided by Cayman Chemicals Inc., Ann Arbor, MI, USA) and we further diluted all samples to 1:2. We measured corticosterone in duplicates using a corticosterone EIA kit (Cayman Chemicals Inc.) on a spectrophotometer plate reader set to 405 nm (BioTek ELX800). Because tadpoles were placed in clean water, no chemicals from the exposure stage interacted with the corticosterone assay. See Supplemental material, Methods for validation of the water-borne corticosterone collection method from *O. septentrionalis* on EIA plates and intra-plate and plate-independent variation.
**Statistical analyses**

Statistical analyses were conducted using R statistical software (R Core Team 2013) and all probability values were calculated using type II sums of squares. We used a general linear mixed-effects model (GLMM: nlme package) to test for all main and interactive effects of atrazine and metyrapone exposure on log-transformed 6 days corticosterone release rates adjusted for mass (pg g⁻¹ h⁻¹) with tank as a random factor and EIA test plate and SVL as covariates. We also used the nlme package to test for all main and interactive effects of atrazine, metyrapone, and *Bd* treatments on 13 and 14 days corticosterone release rates (pg g⁻¹ h⁻¹) using least trimmed squares with EIA test plate and SVL as covariates and tank as a random effect. We used a GLMM to test for the main and interactive effects of atrazine and metyrapone on the number of days to metamorphosis, treating tank as a random effect. To evaluate the effects of treatments on frog growth after metamorphosis, we conducted two-way MANOVAs (atrazine, metyrapone, atrazine x metyrapone) using log-transformed mass and SVL as response variables. We only present the mass data because mass and SVL were highly correlated. To determine the effect of treatments on tadpole and post-metamorphic frog survival, we conducted a mixed-effects Cox proportional hazards survival analysis using the coxme function in the coxme R package.

To test for the effects of treatments on *Bd* resistance, we used the nlme package and lme function to test for the fully crossed effects of life stage (tadpole vs frog), atrazine, and metyrapone on log-transformed *Bd* abundance with sampling time (individuals sampled before and after metamorphosis) nested in tank as random effects. To test for effects on tolerance of *Bd*, we again used the lme function to test for the fully crossed effects of life stage, atrazine, metyrapone, and log *Bd* abundance on the percent mass change of each amphibian (standardized) with sampling time nested in tank as random effects. As a reminder, tolerance is measured as the slope of the relationship between pathogen load and a fitness proxy. Thus, a treatment with a more negative slope is less tolerant of infection (Råberg et al. 2007). When higher order interactions, covariates, or blocking factors were not significant, they were dropped from the statistical model. Finally, we performed Pearson tests to explore the correlations between 6 days corticosterone and mass, development time to metamorphosis, *Bd*, survival, resistance and tolerance for tadpoles and post-metamorphic frogs (all based on tank means). All parametric analyses met the underlying assumptions and we applied an alpha of 0.05.

**Results**

**Baseline corticosterone and stress responses**

We found that baseline corticosterone release rates were significantly lower than the corticosterone release rates of agitated (stress response), and atrazine-exposed tadpoles. See Supplemental material, Results.

**Effects on tadpoles and metamorphosis**

Atrazine elevated corticosterone on experimental day 6 and *Bd* alone elevated corticosterone on experimental day 13 and 14, but metyrapone countered these increases in corticosterone induced by both factors (metyrapone x atrazine: $\chi^2 = 8.20$, $df = 1$, $P = 0.004$; metyrapone x *Bd*: $\chi^2 = 3.87$, $df = 1$, $P = 0.049$; atrazine x *Bd*: $\chi^2 = 3.87$, $df = 1$, $P = 0.068$; Fig. 2a, b, c). Tadpoles exposed to atrazine alone had higher corticosterone than all other treatments ($P < 0.05$, Tukey’s HSD). However, metyrapone plus atrazine did not have significantly different amounts of corticosterone than the control treatment (no atrazine or metyrapone) ($P > 0.05$, Tukey’s HSD; Fig. 2a), indicating that the metyrapone successfully inhibited corticosterone synthesis. Tadpoles exposed to *Bd* alone had higher corticosterone than the control treatment (not exposed to *Bd*) and those exposed to *Bd* plus metyrapone ($P < 0.05$, Tukey’s HSD), but metyrapone alone did not have significantly different amounts of corticosterone than these two treatments (Fig. 2b).

Exposure to atrazine reduced tadpole body size 1 week into the experiment (MANOVA on SVL and mass: $F_{1,36} = 3.60$, $P = 0.04$; Fig. 3a), while metyrapone had no effect on body size at this time (metyrapone: $F_{1,36} = 0.00$, $P = 1.00$; atrazine x metyrapone: $F_{1,36} = 1.32$, $P = 0.25$; Fig. 3b). *Bd* exposure had no significant effects on mass or SVL ($P > 0.05$). Tadpole survival (% mortality) was not significantly affected by atrazine ($\chi^2 = 0.30$, $df = 1$, $P = 0.86$; Fig. 3d) or metyrapone ($\chi^2 = 0.30$, $df = 1$, $P = 0.58$; Fig. 3c). Tadpole survival was weakly affected by an interaction between *Bd* and metyrapone; metyrapone tended to decrease days alive (survival) in the absence of *Bd* but increased it slightly in the presence of *Bd* ($\chi^2 = 3.87$, $df = 1$, $P = 0.05$; Fig. 4a). Time to metamorphosis was not significantly affected by atrazine,
metyrapone, or their interaction (atrazine: $\chi^2 = 0.00$, $df = 1$, $P = 0.99$, metyrapone: $\chi^2 = 0.10$, $df = 1$, $P = 0.75$, atrazine x metyrapone: $\chi^2 = 0.00$, $df = 1$, $P = 0.95$; Supplemental Fig. 1). Similarly, $Bd$ had no effect on time to metamorphosis ($P > 0.05$). See Supporting material, Results for effects of atrazine, metyrapone, and $Bd$ on Gosner stage. Corticosterone on experimental day 6 was not correlated significantly with tadpole mass, time to metamorphosis, or survival ($P > 0.05$; see Supplemental Table 1).

Effects on post-metamorphic frogs

For post-metamorphic frogs, prior exposure to atrazine did not have a significant effect on survival ($\chi^2 = 0.02$, $df = 1$, $P = 0.88$) or body size (MANOVA, $F_{2,64} = 1.82$, $P = 0.17$) and there were no interactive effects of atrazine and metyrapone on these responses ($F_{2,64} = 0.07$, $P = 0.93$). Thus, atrazine reduced size before metamorphosis when measured soon after the chemical exposure but this size difference did not persist post-metamorphosis (stage x atrazine: $F_{1,61} = 5.48$, $P = 0.02$; Fig. 3a). In contrast, metyrapone reduced mass ($F_{2,64} = 4.03$, $P = 0.02$) and survival ($\chi^2 = 15.37$, $df = 1$, $P < 0.001$) after metamorphosis, but did not significantly affect these responses before metamorphosis (Fig. 3b, c), indicating that there were delayed effects of metyrapone. Exposure to $Bd$ did not significantly affect frog survival (days alive) after metamorphosis ($P > 0.05$ for all effects including $Bd$). Once again, corticosterone in tadpoles at experimental day 6 did not significantly correlate with post-metamorphic mass or survival ($P > 0.05$, see Supplemental Table 1).

Effects of atrazine and metyrapone on resistance and tolerance of $Bd$ in both life stages

Of the $Bd$-exposed tadpoles, 21 of 70 (30%) tested positive for $Bd$ with loads ranging from 625 to 52,045 zoospores; we confirmed that control tadpoles were not infected with $Bd$. Of the post-metamorphic frogs exposed to $Bd$, 14 of 34 (41%) became infected, with loads ranging from 154 to 10,010 zoospores.

Atrazine and metyrapone significantly affected resistance to $Bd$. Atrazine lowered $Bd$ abundance in the absence of metyrapone, but elevated $Bd$ abundance in the presence of metyrapone (atrazine x metyrapone $\chi^2 = 8.59$, $df = 1$, $P = 0.003$; Fig. 4b), and this effect was consistent across life stages (all effects including life stage had $P > 0.22$).
Most frogs survived the atrazine and Bd treatments and thus we focused on the ability of frogs to maintain or increase body size while facing the infection as our measure of tolerance. Both atrazine and metyrapone significantly affected tolerance of Bd, but this relationship was dependent on life stage of the individual frog. Metyrapone did not significantly affect tolerance of Bd as tadpoles, but reduced tolerance of post-metamorphic frogs. Frogs previously exposed to metyrapone lost more weight per Bd zoospore than frogs not previously exposed to metyrapone (stage x metyrapone x Bd load: \( \chi^2 = 3.93, df = 1, P = 0.04; \) Fig. 5a, b). This is consistent with the delayed adverse effects of metyrapone observed for growth and survival (Fig. 3b, c). Atrazine reduced tolerance of Bd both before and after metamorphosis. In the full statistical model, there was no evidence of any interactions between atrazine and life stage or atrazine and metyrapone, indicating that the effect of atrazine was consistent across life stages and levels of the metyrapone treatment. Thus, these effects were dropped from the statistical model. The resulting simplified model revealed that frogs with early life exposure to atrazine were less tolerant of Bd infections later in life than frogs that were not exposed to atrazine (atrazine x Bd load: \( F_{1,34} = 4.50, P = 0.04; \) Fig. 5c). Corticosterone on experimental day 6 was not correlated significantly with resistance or tolerance of infections (\( P > 0.05, \) see Supplemental Table 1).

**Discussion**

We explored whether the negative effects of atrazine and Bd on growth, development, and survival of Cuban tree frogs were mediated by the stress hormone corticosterone. We found that corticosterone levels in Cuban tree frogs were elevated after exposure to both an ecologically relevant concentration of atrazine and the fungal pathogen Bd. Importantly, exposure to metyrapone prevented these elevations in corticosterone (Fig. 2), demonstrating that it was successful in inhibiting corticosterone synthesis. However, we found little support for the hypothesis that the adverse effects of atrazine and Bd on growth, development, and tolerance to infection were mediated by corticosterone because corticosterone was not significantly correlated with any of these response variables (Supplemental Table 1) and because the observed effects of atrazine and Bd were not counteracted by exposure to metyrapone. Further studies are required to determine whether atrazine, Bd, and their effects on
corticosterone mediate other events that we did not measure, such as metabolic regulation and oxidative balance.

Similar to our findings, several previous studies have shown that atrazine exposure can disrupt the HPA axis of vertebrates, dysregulating the production of the vertebrate stress hormones cortisol and corticosterone. For example, several studies showed that exposure to ecologically relevant concentrations of atrazine was associated with an increase in circulating corticosterone of small mammals (Fraites et al. 2009; Pruett et al. 2009; Riffle et al. 2014; Rogers et al. 2014) and cortisol of fish (Cericato et al. 2009; Koakoski et al. 2014). Exposure of salamanders and frogs to atrazine increased their circulating corticosterone levels (Larson et al. 1998; McMahon et al. 2017). Hernandez et al. (2014) showed that atrazine competitively inhibits corticosterone from binding with corticosterone-binding globulin in both amphibians and mammals, further indicating that atrazine disrupts corticosterone regulation in these two vertebrate groups.

Fig. 4 a The interactive effect of 6 days of metyrapone (corticosterone synthesis inhibitor) and subsequent Batrachochytrium dendrobatidis (Bd) exposure (15 days) on mean days alive (± 1 SE) for tadpoles. The Bd main effect and metyrapone x Bd interaction are significant ($P < 0.05$). b The interactive effect of metyrapone and atrazine on mean resistance (± 1 SE) to the fungal pathogen Bd (measured as log Bd abundance averaged across the tadpole and post-metamorphic frog exposure periods is significant ($P < 0.05$) for Osteopilus septentrionalis. Because there was no interaction with life stage we combined data across life stages. Numbers indicate number of replicates. See “Results” for statistics

Fig. 5 The relationship between tolerance (standardized mass change) and log Batrachochytrium dendrobatidis (Bd) load for Osteopilus septentrionalis: a 15 days (day 21) after Bd exposure tadpoles, b 5 weeks after Bd exposure post-metamorphic frogs [all individuals were previously exposed to metyrapone (corticosterone synthesis inhibitor) for 6 days], and c after 6 days of atrazine exposure across life stages. The life stage x metyrapone x Bd load interaction in panel (a, b) and the atrazine x Bd load interactions in panel (c) are significant ($P < 0.05$). See “Results” for statistics
corticosterone levels have also been found in wood frog tadpoles (*Rana sylvatica*) infected with ranavirus (Warne et al. 2011), and in lizards infected with parasites compared to non-infected controls (Dunlap and Schall 1995; Oppliger et al. 1998). However, some studies failed to find an effect of infections on circulating corticosterone in birds and frogs (Eggert et al. 2010; Kindermann et al. 2012; Knutie et al. 2013). Importantly, none of these studies experimentally tested whether corticosterone was mediating these responses to disease by inhibiting corticosterone synthesis.

The corticosterone synthesis inhibitor that we used, metyrapone, had different effects than atrazine on survival and development. Similar to Rohr et al. (2013), we did not find an effect of atrazine on short-term or long-term survival of Cuban tree frogs. Additionally, we found that early life exposure to atrazine reduced growth rates, consistent with several previous studies (Rohr and McCoy 2010; Rohr and Palmer 2013). However, this effect on body size before metamorphosis was not significant after metamorphosis. In contrast, metyrapone had no significant effect on survival or body size before metamorphosis, close to when the actual metyrapone exposures occurred, but significantly decreased survival and body size after metamorphosis (Fig. 3b, c). Thus, the effects of metyrapone were persistent but delayed to later in life. These effects are likely caused by either the direct effect of metyrapone exposure or the indirect effect of low baseline levels of corticosterone on growth and survival.

We also note that metyrapone did not reduce corticosterone in the absence of stressors. This might be because the non-stressed tadpoles (such as the controls) could already be approaching the lower bound levels of baseline corticosterone necessary for homeostasis or because metyrapone cannot reduce circulating levels of corticosterone, only new corticosterone synthesis. Thus, in the absence of a stressor, we might not expect metyrapone to reduce baseline levels of circulating corticosterone if the half-life of corticosterone is reasonably long (see Supplemental material, Discussion of corticosterone half-lives). Similar to our finding, Glenneimeier and Denver (2002b) found that metyrapone lowered corticosterone when individuals were stressed but not when they were not stressed.

We found that both metyrapone and atrazine affected amphibian defenses against *Bd*. Metyrapone exposure was associated with reduced tolerance to *Bd* when infections occurred after, but not before, metamorphosis (Fig. 5b), which is consistent with the delayed effects of metyrapone on growth and survival. While it is possible that reduced tolerance to *Bd* was mediated by dysregulation of corticosterone during *Bd* exposure, this finding is more likely caused by the accumulation of negative effects of metyrapone exposure through time and differences in susceptibility to *Bd* between the two life stages (Fisher et al. 2009; Rohr et al. 2013). Unlike our finding for metyrapone, we found that atrazine lowered tolerance of *Bd* infections similarly for both tadpoles and post-metamorphic frogs (Fig. 5c), results that match those of Rohr et al. (2013). Early life exposure to atrazine reduced mass in *Bd*-infected tadpoles and post-metamorphic frogs, indicating that early life exposure to atrazine had both short- and long-term effects on tolerance to *Bd*.

Although atrazine reduced mass in *Bd*-infected individuals of both life stages, exposure to metyrapone did not significantly counter these reductions in *Bd* tolerance, and thus, atrazine-induced elevations in corticosterone levels could not account for atrazine-induced reductions in tolerance of *Bd*. Although metyrapone had adverse effects later in life, these adverse effects could not account for the fact that metyrapone did not counteract the adverse effects of atrazine early in life (when it did not have detectable effects). Thus, it seems unlikely that any adverse effects of metyrapone could be masking any corticosterone-mediated effects. Given that corticosterone does not appear to primarily mediate the effects of atrazine or *Bd* on amphibian growth or development, or the effects of atrazine on tolerance of *Bd* infections, these effects are more likely caused by direct effects of atrazine and *Bd* exposure, such as through energy lost from atrazine detoxification, defense against *Bd*, or repair from damage caused by atrazine or *Bd* (McMahon et al. 2013b; Voyles et al. 2009). Alternatively, these effects could be from indirect effects of atrazine and *Bd* on unmeasured hormones, such as thyroxine or steroidal sex hormones. For example, in larval tiger salamanders, atrazine exposure elevated thyroxine, another hormone associated with amphibian growth and metamorphosis (Larson et al. 1998). Additionally, several meta-analyses have revealed a negative association between testosterone and immunity, with sexually mature male vertebrates often exhibiting greater susceptibility to infection and higher parasite burdens in the field (Zuk 1996; Zuk and McKeans 1996). Moreover, recent studies have revealed a causal relationship between testosterone, reduced immunity, and increased parasite loads in mice (Krucken et al. 2005; Lotter et al. 2013).

In contrast to the effects of atrazine on *Bd* tolerance, the effect of atrazine on resistance to *Bd* infections (i.e., lowered *Bd* load) did depend significantly on exposure to metyrapone (Fig. 4b). Resistance to *Bd* was higher when frogs were exposed to atrazine alone than to a combination of atrazine and metyrapone. However, our data do not strongly support the hypothesis that corticosterone was mediating this altered resistance to *Bd*. First, the pattern of highest corticosterone for atrazine alone and lower corticosterone for all other treatments (Fig. 2a) is not parallel to the patterns of resistance across atrazine and metyrapone treatments (Fig. 4b). Specifically, resistance was highest for atrazine and metyrapone alone, and was lowest for the solvent control and when atrazine and metyrapone were combined. Second, we did not find a significant interaction
between atrazine, metyrapone, and \( Bd \) exposure on corticosterone, suggesting that these together did not mediate the observed resistance pattern. Third, there was no significant correlation between day 6 corticosterone and \( Bd \) abundance on tadpoles and post-metamorphic frogs (Supplemental Table 1). There was also no significant correlation between day 13, 14 corticosterone and \( Bd \) abundance on tadpoles (Pearson’s \( R = -0.070 \); \( P = 0.70 \)) or post-metamorphic frogs (Pearson’s \( R = 0.040 \); \( P = 0.87 \)). Hence, our results regarding the relationship between corticosterone and \( Bd \) are equivocal, much like the literature on this topic. For example, some research suggests that corticosterone increases resistance to \( Bd \) (Murone et al. 2016; Tatiersky et al. 2015), whereas other research suggests that it has no effects (Searle et al. 2014). Additional work is needed to more thoroughly grasp the generality of effects of corticosterone on resistance to infections.

In conclusion, we found that there are costs of exposure to atrazine and \( Bd \), which supports the results of other studies (Rohr and McCoy 2010; Rohr et al. 2013). However, by inhibiting corticosterone production with metyrapone, we found that increased corticosterone from atrazine and \( Bd \) exposure may not be the main factor mediating the observed decreases in growth, development, and tolerance to infection in Cuban tree frogs. Instead, our findings might largely be caused by repair from any damage caused by atrazine and \( Bd \) or indirect effects of atrazine and \( Bd \) on hormones other than corticosterone. Increased exposure to contaminants and pathogens are just two of many examples of how human activities are adversely affecting biodiversity and more studies are required to understand the mechanisms driving these effects (Dirzo et al. 2014). In particular, given that measurements of stress hormones are regularly being used to direct the management of wildlife populations (reviewed by Busch and Hayward 2009; Sheriff et al. 2011), additional studies are required to evaluate whether the adverse effects of the Anthropocene on biodiversity are often mediated by the effects of anthropogenic factors on the stress responses of organisms and their subsequent impacts on fitness.

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Compliance with ethical standards

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Conflict of interest The authors declare that they have no conflict of interest.

Ethical approval All applicable institutional and/or national guidelines for the care and use of animals were followed. This project was approved by the Texas State University Animal Care and Use Committee # 201485314.

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