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8 RUNNING TITLE: Larval cannibalism in *Phyllobates* frogs

9 Keywords: Tadpole cannibalism, Dendrobatidae, Parental care, competition

10 Ethics Statement

11 Frog rearing protocols were approved by the University of Chicago and the University of

Michigan's Institutional Animal Care and Use Committees (UChicago protocol # 72416, UMich
 protocol # PRO00010325).

14 Funding

The rearing of the frog colonies used in this study has been supported by NSF grants DEB-1702014 and IOS-1827333, by the Committee on Evolutionary Biology at the University of Chicago, and by the Michigan Society of Fellows and the Department of Ecology and Evolutionary Biology at the University of Michigan.

19 **Conflicts of interest**

20 The author reports no conflicts of interest.

21 Abstract

22 Cannibalistic behaviour between tadpoles of dendrobatoid poison frogs has been observed in 23 several species with complex parental care dynamics, leading to the idea that it may have played a role in the evolution of parental care. However, the existence or characteristics of this behaviour 24 25 beyond a handful of well-studied species remains largely unknown. I report direct and indirect 26 observations of larval cannibalism in two species of the genus *Phyllobates*, which seem to occur 27 less frequently than in other, better studied species, and possibly associated with body size 28 differences. Beyond expanding the phylogenetic breadth of tadpole cannibalism among poison 29 frogs, my observations point to interesting aspects of this behaviour, such as its apparently plastic 30 and continuous nature, and highlight the importance of studying other lineages to understand the its 31 drivers and effects on poison frog evolution.

32 Keywords: Tadpole cannibalism, Dendrobatidae, Parental care

33 1. INTRODUCTION

34 Predation of conspecifics (i.e. cannibalism) is widespread among animals (Fox 1975; Elgar and 35 Crespi 1992). This behaviour is prevalent among amphibian tadpoles, where it has been associated with the resource-limited environments (e.g. ephemeral pools) in which they develop (Polis and 36 Myers 1985; Crump 1990, 1992). Although tadpole cannibalism has been studied in a variety of 37 contexts, such as kin recognition (Pfennig et al. 1993; Pfennig 1997; Gray et al. 2009), foraging 38 39 behaviour (Caldwell and de Araujo 1998), phenotypic plasticity (Pfennig 1990, 1992), neuroethology (Fischer et al. 2020), and parental care (Summers 1999; Downie et al. 2001; Brown 40 41 et al. 2009; Rojas 2014), its prevalence across the amphibian phylogeny is not well known.

42 Larval cannibalism has received considerable attention in Dendrobatoid poison frogs (families 43 Aromobatidae and Dendrobatidae). Tadpoles of most species in this group develop in small, resource-poor pools such as puddles in the forest floor, or water-filled plant structures, also known 44 45 as phytotelmata (e.g bromeliad axels, tree holes, fallen seed husks and petioles). Dendrobatoids lay eggs outside the water, and exhibit considerable parental care, which involves guarding eggs and 46 transporting tadpoles to water upon hatching. In some species parental care extends up to 47 48 metamorphosis, with females providing infertile eggs to feed their tadpoles (Weygoldt 1980, 1987; 49 Myers and Daly 1983; Summers and McKeon 2004). Within this group tadpole cannibalism has been reported almost exclusively in a group of closely related genera, namely Adelphobates 50 51 (Caldwell and de Araujo 1998), Andinobates (Suárez-Mayorga 1999; Cáceres 2012), Dendrobates 52 (Gómez 2006; Rojas 2014), Oophaga (Dugas et al. 2016), and Ranitomeva (Summers 1999; Poelman and Dicke 2007; Brown et al. 2011; Schulte et al. 2011; Acosta et al. 2013; Fig. 1). Species 53 54 in these genera usually display higher levels of parental investment per individual offspring than other dendrobatoids: They lay smaller clutches, often deposit tadpoles individually in phytotelmata, 55 and in many cases extend parental care until metamorphosis, which has lead to the idea that the 56

evolution of increased parental investment may be related to larval cannibalism in the face of
limited resources (Summers 1999; Summers and McKeon 2004; Carvajal-Castro et al. 2021).

59 Despite the sustained interest in larval cannibalism over the past few decades, its prevalence and 60 phylogenetic distribution within dendrobatoid frogs remains largely unstudied. In fact, the existence and characteristics of this behaviour have only been evaluated in one species outside the clade 61 62 formed by the genera named above (Fig 1): Mannophryne trinnitatis (Downie et al. 2001) Although 63 in some cases cannibalism has been considered absent in a handful of species (Carvajal-Castro et al. 64 2021), this has been based on anecdotal observations of conspecific tadpoles coexisting without 65 apparent aggression, so it remains unclear whether cannibalism occurs in these species. Here I report an instance of larval cannibalism in captive-bred individuals of *Phyllobates aurotaenia*, as 66 well as indirect evidence of the same behaviour in *P. terribilis*, both of which occurred incidentally 67 while conducting research on other aspects these species' biology. 68

69 2. METHODS AND RESULTS

70 Adult frogs were kept in groups of 2-6 animals in terrariums and allowed to breed freely. Tanks 71 were checked daily for new egg clutches, which were removed from the tank and reared in petri 72 dishes kept within a sealed container with a moist paper towel to maintain high humidity. After hatching, tadpoles were transferred to 250ml plastic cups filled with reverse-osmosis (RO) water 73 74 infused with almond leaf extract, and left to acclimate for three days. On the fourth day post-75 hatching, one pellet of Frog and Tadpole Bites (Pisces Pros, upc. 788459100303) dusted with Micron Fry Food (Sera, cat. no. 00720) was offered per tadpole. New food was offered when all 76 pellets in a cup had been consumed, and once a week droppings and leftover food were sucked out 77 with a pipette, and new pellets were added. 78

79 Tadpoles were reared in groups of 2-8, which in most cases were made up of full siblings from the 80 same clutch, although in some cases tadpoles from different clutches were reared together. In one 81 occasion, a newly hatched tadpole of P. aurotaenia (stage 24-25 sensu Gosner, 1960) was moved to 82 a cup with a single stage 34 conspecific that had been last offered food six days prior. 83 Approximately one hour later I observed the older tadpole insistently swimming around and inspecting the hatchling. Soon after, it proceeded attack the hatchling, aggressively biting its tail. 84 85 Initially the hatchling tadpole was able to turn lose, but the large one persisted until the hatchling 86 became motionless and did not display a detectable heartbeat at plain sight. Over the following ~20 87 minutes the large tadpole consumed its prey in its entirety. Figure 2A-E show frames captured from 88 a video of part of the event. In addition to the event described above, in three occasions I noticed 89 that P. terribilis tadpoles being reared in groups with their clutch-mates went missing between daily checkups. Given the daily fcheckups and the apparent good health these individuals. I consider 90 91 cannibalism to be the most likely explanation for their disappearance, as dead tadpoles would have 92 been noticed before decomposing. The missing tadpoles were in all cases between Gosner stages 93 25-32 and were noticeably smaller than the largest of their clutch mates, despite being the same age.

94 **3. DISCUSSION**

95 All instances of cannibalism reported here involved predatory tadpoles that were noticeably larger than their prey, suggesting that size disparity may be a key element in the occurrence of cannibalism 96 97 in *Phyllobates*. This falls in line with previous work on several cannibalistic species (Claessen et al. 98 2004; Ibáñez and Keyl 2010), including other poison frogs (Rojas 2014; Fouilloux et al. 2022), and highlights the facultative, opportunistic nature of larval cannibalism in poison frogs. Phyllobates 99 100 males transport groups of up to ~10-20 tadpoles, which they are though to deposit in groups at 101 water sources on or close to the forest floor, such as fallen palm bracts and petioles or hollowed logs, that may already be inhabited by conspecifics (Silverstone 1976; Myers et al. 1978; R. 102

103 Márquez pers. obs.). In view of this, I suggest two main situations that may trigger cannibalism in 104 *Phyllobates* tadpoles: First, as has been suggested for other species (e.g. Rojas 2014), if tadpoles are deposited in a pool already inhabited by older, larger conspecifics, they are at risk of being 105 106 consumed. Second, when competition for resources between similarly-aged tadpoles inhabiting the 107 same water body (including clutch mates) leads to some individuals growing more slowly than others, and eventually being cannibalised by their larger conspecifics. Clutches reared communally 108 109 in captivity often display marked body size variation among tadpoles (Fig 2F), suggesting that this 110 is an ecologically relevant scenario. In both cases food scarcity is likely to promote cannibalism, both by directly motivating larger tadpoles to attack their smaller conspecifics, and by promoting 111 112 stronger intraspecific competition, which in turn leads to greater variance in body size among the tadpoles inhabiting a pool, and increases the likelihood of cannibalism. Further research on the 113 reproductive behaviour of adults and the ecological interactions between tadpoles inhabiting the 114 115 same pool should help illuminate the relative frequency of both situations, as well as the relationship between cannibalism and the evolution of parental care strategies in *Phyllobates*. 116

117 The existence of larval cannibalism in Mannophryne trinitatis (Downie et al. 2001), Phyllobates 118 aurotaenia, and P. terribilis (this study) could, at first glance, appear to challenge to the idea that larval cannibalism is a driver of increased parental investment, since these three species are thought 119 120 to invest less in parental care than all others where larval cannibalism has been reported, and similarly to species where this behaviour has not been observed (Summers et al. 1999; Summers 121 122 and McKeon 2004; Carvajal-Castro et al. 2021). However, considering that most examined 123 dendrobatoid tadpoles have omnivorous or predaceous diets (Lehtinen et al. 2004; Ryan and Barry 2011), which in some cases appear indiscriminate (Caldwell and de Araujo 1998), it is possible that 124 125 tadpoles of most species possess the anatomical and neural machinery required to attack and 126 consume conspecifics. Instead of the discrete presence/absence of cannibalism, species may mostly vary in the conditions that trigger this behaviour (e.g. size differences, food scarcity), and the ease 127

with which it is triggered. The fact that I have observed at most four cannibalistic interactions over
eight years of *Phyllobates* captive breeding efforts, while observations of this behaviour have been
amply reported in other groups such as *Ranitomeya* (Summers 1999; Poelman and Dicke 2007;
Schulte et al. 2011; Acosta et al. 2013; Fischer et al. 2020) or *Dendrobates* (Gómez 2006; Rojas
2014; Fouilloux et al. 2022), both in the lab and in the field, falls in line with this idea.

133 Larval cannibalism has been recognised as an important factor in the ecology and evolution of a 134 group of closely-related poison frog genera with more specialised parental care than their relatives. My observations of this behaviour outside this group not only expand the phylogenetic breadth of 135 136 this behaviour (Fig. 1), but also suggest interesting avenues for future research and reflection around it, such as the degree to which cannibalism can be modelled as a continuous or threshold 137 trait (sensu Wright 1934), rather than a binary one, or the environmental and ecological contexts 138 139 that trigger its expression. Further inquiries of larval cannibalism in other branches of the dendrobatoid phylogeny will certainly help illuminate these and other aspects of the ecology and 140 141 evolution of larval cannibalism in poison frogs.

142 Acknowledgements

I thank members of the Animal Resource Center at the University of Chicago and the Unit for
Laboratory Animal Medicine at the University of Michigan for assistance with frog rearing,
Marcus Kronforst and Bibiana Rojas for encouragement to publish these observations, and Bibiana
Rojas for feedback on the manuscript.



Figure 1. Genera of dendrobatoid frogs where tadpole cannibalism has been observed, with *Phyllobates* highlighted in bold. The topology follows (Grant et al. 2017), and divergence times follow Santos et al. (2014), Guillory et al. (2019), and Douglas, Márquez & Tarvin (i*n prep*).



Figure 2. Larval cannibalism in *Phyllobates aurotaenia* (A-E) and within-clutch body size variation in *P. terribilis* (F). Some time after being placed in the same cup, a stage 34 *P. aurotaenia* inspected (A), attacked (B, C), and pursued (D) a recently hatched conspecific (stage 24-25), before eventually killing and consuming it (E). Panel F illustrates body size variation between communally-reared *P. terribilis* tadpoles of the same clutch, presumably due to intraspecific competition. Red arrows indicate two individuals noticeably smaller than the rest, despite being the same age.

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