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**Permalink** https://escholarship.org/uc/item/2p471675

**Journal** Molecular Biology and Evolution, 40(10)

# Authors

Wan, Yin Navarrete Méndez, María OConnell, Lauren <u>et al.</u>

Publication Date 2023-10-04

# DOI

10.1093/molbev/msad206

Peer reviewed

# Selection on Visual Opsin Genes in Diurnal Neotropical Frogs and Loss of the SWS2 Opsin in Poison Frogs

Yin Chen Wan (**b**,<sup>†,1,2,3</sup> María José Navarrete Méndez (**b**,<sup>†,1,4</sup> Lauren A. O'Connell (**b**,<sup>5</sup> Lawrence H. Uricchio (**b**,<sup>1,6</sup> Alexandre-Benoit Roland (**b**,<sup>7,8</sup> Martine E. Maan (**b**,<sup>9</sup> Santiago R. Ron (**b**,<sup>4</sup> Mileidy Betancourth-Cundar (**b**,<sup>10</sup> Marcio R. Pie,<sup>11,12</sup> Kimberly A. Howell,<sup>13</sup> Corinne L. Richards-Zawacki (**b**,<sup>13</sup> Molly E. Cummings,<sup>14</sup> David C. Cannatella,<sup>14,15</sup> Juan C. Santos,<sup>\*,16</sup> and Rebecca D. Tarvin (**b**<sup>\*,1</sup>

<sup>1</sup>Museum of Vertebrate Zoology and Department of Integrative Biology, University of California Berkeley, Berkeley, CA, USA <sup>2</sup>Department of Molecular Genetics, University of Toronto, Toronto, Ontario, Canada

<sup>3</sup>School of Biomolecular and Biomedical Science, University College Dublin, Belfield, Dublin, Ireland

<sup>4</sup>Museo de Zoología, Facultad de Ciencias Exactas y Naturales, Pontificia Universidad Católica del Ecuador, Quito, Ecuador

<sup>5</sup>Department of Biology, Stanford University, Palo Alto, CA, USA

<sup>6</sup>Department of Biology, Tufts University, Medford, MA, USA

<sup>7</sup>FAS Center for Systems Biology, Harvard University, Cambridge, MA, USA

<sup>8</sup>Research Centre on Animal Cognition (CRCA), Centre for Integrative Biology (CBI), UMR5169 CNRS, Toulouse University, Toulouse, France

<sup>9</sup>Groningen Institute for Evolutionary Life Sciences, University of Groningen, Groningen, The Netherlands

<sup>10</sup>Departamento de Ciencias Biológicas, Universidad de los Andes, Bogotá, Colombia

<sup>11</sup>Department of Zoology, Universidade Federal do Paraná, Curitiba, Brazil

<sup>12</sup>Biology Department, Edge Hill University, Ormskirk, United Kingdom

<sup>13</sup>Department of Biological Sciences, University of Pittsburgh, Pittsburgh, PA, USA

<sup>14</sup>Department of Integrative Biology, University of Texas at Austin, Austin, TX, USA

<sup>15</sup>Biodiversity Center, University of Texas at Austin, Austin, TX, USA

<sup>16</sup>Department of Biological Sciences, St. John's University, New York City, NY, USA

<sup>†</sup>These authors contributed equally to this work and agree that each has the right to list themselves as first author on their CVs.

\*Corresponding authors: E-mails: rdtarvin@berkeley.edu; santosj@stjohns.edu.

Associate editor: Dr. Belinda Chang

# Abstract

Amphibians are ideal for studying visual system evolution because their biphasic (aquatic and terrestrial) life history and ecological diversity expose them to a broad range of visual conditions. Here, we evaluate signatures of selection on visual opsin genes across Neotropical anurans and focus on three diurnal clades that are well-known for the concurrence of conspicuous colors and chemical defense (i.e., aposematism): poison frogs (Dendrobatidae), Harlequin toads (Bufonidae: *Atelopus*), and pumpkin toadlets (Brachycephalidae: *Brachycephalus*). We found evidence of positive selection on 44 amino acid sites in *LWS*, *SWS1*, *SWS2*, and *RH1* opsin genes, of which one in *LWS* and two in *RH1* have been previously identified as spectral tuning sites in other vertebrates. Given that anurans have mostly nocturnal habits, the patterns of selection revealed new sites that might be important in spectral tuning for frogs, potentially for adaptation to diurnal habits and for color-based intraspecific communication. Furthermore, we provide evidence that *SWS2*, normally expressed in rod cells in frogs and some salamanders, has likely been lost in the ancestor of Dendrobatidae, suggesting that under low-light levels, dendrobatids have inferior wavelength discrimination compared to other frogs. This loss might follow the origin of diurnal activity in dendrobatids and could have implications for their behavior. Our analyses show that assessments of opsin diversification in across taxa could expand our understanding of the role of sensory system evolution in ecological adaptation.

Key words: gene loss, blue-sensitive opsin, Dendrobatidae, Atelopus, Brachycephalus, spectral tuning.

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#### Introduction

Natural selection has favored visual systems that maximize the detection and discrimination of light wavelengths that are relevant to organismal activity patterns and habitats (Warrant and Johnsen 2013). For instance, nocturnal or crepuscular species have visual systems that capture maximal light or permit color discrimination in dim-light conditions (Bowmaker 2008; Gutierrez et al. 2018; Mohun and Davies 2019; Guo et al. 2023). In contrast, diurnal species are active when the light spectrum is the brightest and broadest (e.g., 350-700 nm); therefore, their visual systems may be evolutionarily tuned for improved wavelength discrimination (i.e., color vision; Bowmaker 2008). In vertebrates, visual opsin genes, which encode G-protein-coupled receptors that bind to a retinal chromophore (Bowmaker 2008), have diversified and undergone selection for adaptation to various light environments. For example, diurnal primates underwent duplication and subsequent spectral tuning of the long-wavelength-sensitive opsin twice (Hunt et al. 1998), while fishes have undergone duplication and diversification of opsin genes many times (Hofmann and Carleton 2009). In addition, some groups have lost opsin genes during adaptation to new photic environments including the loss of cone opsins in coelacanths (Yokoyama et al. 1999) and the loss of a short-wavelength-sensitive cone opsin in mammals during adaptation to nocturnal lifestyles (Bowmaker 2008).

The visual opsin genes present in vertebrates include long-wavelength-sensitive (LWS or OPN1LW), middlewavelength-sensitive (RH2, MWS, or RHB; not to be confused with an independent gene duplication that gave rise to a middle-wavelength-sensitive opsin gene in Old World primates also called MWS or OPN1MW [Nathans et al. 1986; Hunt et al. 1998]), short-wavelength-sensitive 1 (SWS1 or OPN1SW), and short-wavelength-sensitive 2 (SWS2 or OPN2SW; Bowmaker 2008; Schott et al. 2022). Other vision-related genes include the rod opsin, rhodopsin (RH1, RHO, RHA, or OPN2), which evolved after a duplication and divergence event from an ancestral opsin shared with RH2 (Okano et al. 1992). Color vision is achieved when two or more photoreceptors with opsins differing in wavelength sensitivity are simultaneously activated, and their signals are compared by the observer's central nervous system (Bowmaker 2008; Gibson 2014). For consistency, we refer to the opsin genes using the following acronyms: LWS, SWS1, SWS2, RH1, and RH2.

The repertoire of visual opsin genes and the spectral sensitivity of retinal photoreceptors have frequently undergone changes to accommodate the evolutionary transition between diurnal and nocturnal activity patterns (Surridge et al. 2003; Hauzman et al. 2017), a shift that has occurred repeatedly throughout the phylogenetic history of animals (Anderson and Wiens 2017; Akiyama et al. 2022). For instance, gene duplication has been observed independently in the common ancestor of catarrhine primates (Old World monkeys, apes, and humans) and in the New World howler monkeys as a consequence of the transition to diurnal ecologies from nocturnal ancestors (Surridge et al. 2003). The transition to diurnality has also resulted in the loss of rod photoreceptors and functional *RH1* genes in some diurnal species of snakes and lizards (Hauzman et al. 2017). Similarly, spectral tuning of opsin proteins has also been involved in adaptation to bright-light environments in several diurnal species of insects (Akiyama et al. 2022), snakes (Hauzman et al. 2017), and birds (Borges et al. 2015). Vision in anuran amphibians is particularly interesting as most species are crepuscular or nocturnal, but several clades have independently evolved diurnal activity (Anderson and Wiens 2017). Some nocturnal frog species (e.g., *Rana temporaria, Xenopus laevis*, and *Rhinella marina* [*Bufo marinus*]) have played critical roles in discovering the biology of vision, yet much less is known regarding vision in diurnal species.

The anuran retina is thought to typically contain three types of cone cells that are active under bright-light conditions: two types of long-wavelength-sensitive cones that express LWS, and a short-wavelength-sensitive cone that expresses SWS1 (Liebman and Entine 1968; Hárosi 1982; Koskelainen et al. 1994; Mohun and Davies 2019; Donner and Yovanovich 2020). Some anuran species such as Rana pipiens (nocturnal) and Oophaga pumilio (diurnal), possess an additional fourth cone with middle-wavelength sensitivity ( $\lambda_{max}$  of ~500 nm), but the pigment contained in this type of photoreceptor cell remains to be determined (Siddigi et al. 2004; Mohun and Davies 2019). In some vertebrates (e.g., fish and reptiles; Bowmaker 2008), the RH2 gene is responsible for green or middle-wavelength sensitivity. To date, the RH2 gene has not been found in any amphibian, and it is believed to have been lost in their last common ancestor (Mohun and Davies 2019; Schott et al. 2022). Thus, it is plausible that RH1 is the gene that is expressed in greensensitive cones in amphibians (Schott et al. 2022), conferring them with the ability to detect middle-wavelength light spectra during the day, as has occurred in at least one species of diurnal snake, Thamnophis proximus (Schott et al. 2016). Although some vertebrates use oil droplets to further tune wavelength sensitivity, oil droplets in anurans are often absent or colorless and thus are unlikely to influence the spectral sensitivity of cone opsin proteins (Toomey and Corbo 2017). For example, oil droplets are present but colorless in O. pumilio (Siddigi et al. 2004). In addition to these four cone cell types (single and double LWS cones, single SWS1 cones, and an MWS cone), the anuran retina has two types of rod cells that are active in dim-light conditions: one that expresses RH1, known as the "red rod," and another that expresses SWS2, known as the "green rod" (Hunt and Collin 2014). The existence of two rod types likely allows some amphibians to have low-light (scotopic) and rodbased color discrimination (Yovanovich et al. 2017).

Despite several frog species being key to our early understanding of vision (Liebman and Entine 1968), the vast diversity of anuran visual systems is only just beginning to be unraveled (Donner and Yovanovich 2020). For example, only 108 of the >7,500 currently described anuran taxa (AmphibiaWeb 2022) have information on lens transmission properties (Yovanovich et al. 2020; Thomas et al. 2022), and the spectral sensitivity of retinal photoreceptor cells has been described for only 10 species (Donner and Yovanovich 2020). Moreover, the genetic basis of anuran vision is especially understudied (Mohun and Davies 2019), with genomic resources limited to a handful of well-assembled frog genomes until very recently (Womack et al. 2022) and one study of opsin genes in 33 frogs (Schott et al. 2022). Compared to other vertebrates, the lack of information on the molecular biology of anuran vision contrasts with the relatively large body of research into anuran visual ecology (e.g., Toledo and Haddad 2009; Bell and Zamudio 2012; Rößler et al. 2019). One example is Dendrobatidae (poison frogs), which has become a model system for understanding how natural selection by predators shapes the origin and subsequent diversification of warning signals (e.g., Clough and Summers 2000; Richards-Zawacki and Cummings 2011; Santos and Cannatella 2011; Wang 2011; Cummings and Crothers 2013; Rojas et al. 2014; Lawrence et al. 2019). Chemically defended and brightly colored (aposematic) dendrobatids are also a special focus of molecular work, including published genomes from O. pumilio and Ranitomeya imitator, with several more being assembled. Other frog families that include diurnal and aposematic species include some Neotropical genera of Bufonidae and Brachycephalidae. Atelopus (Harlequin toads, Bufonidae) and Brachycephalus (Pumpkin toadlets, Brachycephalidae) have a handful of studies on color-based intraspecific or visual signaling in brightly colored species (Rebouças et al. 2019; Rößler et al. 2019), but no genomic or transcriptomic work focusing on genes involved in vision in Atelopus and Brachycephalus have been conducted, and no publicly available genomes from either group exist.

Here, we aimed to describe patterns of selection on visual opsin genes in three diurnal, Neotropical frog clades: poison frogs (Dendrobatidae or Dendrobatoidea sensu Grant et al. [2006]), Harlequin toads (Bufonidae: Atelopus), and pumpkin toadlets (Brachycephalidae: Brachycephalus). We sequenced the four visual opsin genes known in amphibians (LWS, SWS1, SWS2, and RH1) using a target-bait capture approach including 116 species with a special emphasis on the three focal clades. We reviewed patterns of positive selection on amino acid sites relevant to the spectral sensitivity of opsin proteins, which may have been modified via substitutions to better absorb wavelengths that are most relevant to the organism's visual ecology, a process known as spectral tuning (Carvalho et al. 2007; Hunt et al. 2007; Osorio and Vorobyev 2008). Using a combination of selection analyses, we determined whether amino acid positions at or near known spectral tuning sites were associated with the transitions to diurnality in each clade and/or were under positive selection.

### Results

#### **Opsin Sequences and Phylogenies**

We retrieved 28 opsin gene sequences of 13 species from GenBank and reconstructed another 49 (representing 19 species) using SRA data deposited in NCBI (see supplementary table S1, Supplementary Material online). From 484,530 Illumina reads from our bait capture experiment, we further reconstructed 289 new opsin gene sequences representing 87 anuran species. In total, we analyzed 366 sequences from 116 anuran species. The latter two-thirds of the reconstructed SWS2 gene had low coverage in many species, so sites after amino acid position 129 were excluded from downstream analyses (see Materials and Methods). Following this exclusion, all assembled sequences had an average read depth of  $>10\times$ , with 82% having coverage  $>50 \times$  (see supplementary table S2, Supplementary Material online). Estimated gene trees were largely concordant with recent family-level phylogenies (Feng et al. 2017; Hime et al. 2021) and hypothesized interspecific relationships (Pyron 2014; Grant et al. 2017; Jetz and Pyron 2018; see supplementary Data S3, Supplementary Material online). The phylogeny inferred using all opsin gene sequences was reciprocally monophyletic for each gene, indicating a low probability of chimeric sequences in our data set.

#### Analyses of Selection

To identify patterns of positive selection that might be associated with diurnality, we used a series of Contrast-FEL analyses to identify sites under differential dN/dS regimes between groups of branches (table 1). We excluded SWS2 from this analysis because no sequences from Dendrobatidae were available (see below). Contrast-FEL estimates site-specific dS values for the entire phylogeny; it then estimates and statistically compares dN values for each site between foreground and background branch sets. For sites identified by Contrast-FEL to be under different selection regimes, we used FEL to determine whether  $dN/dS(\omega)$  was statistically distinguishable from 1, with values less than 1 indicating negative selection and values greater than 1 indicating positive selection. We first compared the dN value estimated for transition branches between nocturnal and diurnal lineages (the stem branches of Dendrobatidae, of Atelopus, and of Brachycephalus) to a dN value estimated for all other branches; no sites were identified as under different patterns of selection (see supplementary Data S1, Supplementary Material online). Then we directly compared dN values between diurnal and nocturnal clades ("DIURNAL" foreground, see Materials and Methods, supplementary table S1, Supplementary Material online, and the supplementary Material for how we determined activity states). This analysis identified two sites in LWS, three sites in SWS1, and four sites in RH1 to be under differential selection regimes in foreground and background lineages (table 1). As six of these sites were also identified by other methods to be under positive selection when the entire tree was included (using FEL, CODEML, FUBAR, or MEME approaches; table 2), we suspected that our large sampling of dendrobatid lineages (all diurnal) might be in part driving this pattern. We then compared *dN* values for Dendrobatidae branches plus its stem branch ("DENDRO" foreground, see table 1) versus other branches. This analysis identified all but one

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Table 1. Results From Contrast-FEL Analyses Comparing Selection Patterns Between Lineage Groups.

Gene	Codon number	Nearby	Location	Foreground	Contrast-FEL		FEL		Interpretation		
	(bovine RH1 numbering)ª	spectral tuning site			dS	dN					
						Fore- ground	Back ground	Foreground (P-value)	Background (P-value)		
LWS	145	None	ECD II	DIURNAL	1.023	1.690	0.093	Neutral (0.385)	Negative (0.007)	Similar results for DIURNAL and DENDRO	
				DENDRO	1.019	2.077	0.086	Neutral (0.224)	Negative (0.006)		
	176 <sup>c</sup>	None	TMD IV	DIURNAL	10.583	0.732	0.000	Negative (<0.001)	Negative (<0.001)	Similar results for DIURNAL and DENDRO	
				DENDRO	10.478	0.901	0.000	Negative (<0.001)	Negative (<0.001)		
	316	None	C-T	DENDRO	1.981	2.304	0.165	Neutral (0.787)	Negative (0.001)	Under negative selectior in DENDRO background	
	318	None	C-T	DIURN-DEN	1.204	3.000	0.000	Neutral (0.323)	Negative (<0.001)	Under negative selectior in DIURN-DEN background	
SWS1	218	None	TMD V	DIURNAL	0.797	1.808	0.096	Neutral (0.120)	Negative (0.017)	Similar results for DIURNAL and DENDRO	
				DENDRO	0.798	2.173	0.089	Neutral (0.058)	Negative (0.013)		
	221	None	TMD V	DIURNAL	0.441	2.135	0.076	Positive (0.024)	Neutral (0.143)	Similar results for DIURNAL and DENDRO	
				DENDRO	0.442	2.628	0.072	Positive (0.010)	Neutral (0.127)		
	342	None	C-T	DIURNAL	0.808	0.838	0.000	Neutral (0.0962)	Negative (0.001)	Similar results for DIURNAL and DENDRO	
				DENDRO	0.802	0.998	0.000	Neutral (0.756)	Negative (0.001)		
RH1	22	None	N-T	DIURN-DEN	1.428	2.026	0.047	Neutral (0.674)	Negative (0.001)	Under negative selectior in DIURN-DEN background	
	87 <sup>c</sup>	None	TMD II	DIURNAL	0.000	1.306	0.092	Positive (0.002)	Neutral (0.370)	Similar results for DIURNAL and DENDRO	
				DENDRO	0.000	1.624	0.082	Positive (0.001)	Neutral (0.577)		
	97 <sup>b,c</sup>	96	TMD II	DENDRO	0.181	0.000	1.429	Neutral (0.251)	Positive (0.009)	Under positive selection in DENDRO background	
	124 <sup>c</sup>	122, 124, 125	TMD III	DIURNAL	1.986	2.204	0.086	Neutral (0.817)	Negative (<0.001)	Similar results for DIURNAL and DENDRO	
				DENDRO	1.988	2.564	0.156	Neutral (0.583)	Negative (<0.001)		
	169 <sup>b,c</sup>	None	TMD IV	DIURNAL	0.433	5.346	1.099	Positive (<0.001)	Neutral (0.110)	Similar results for DIURNAL and DENDRO	
				DENDRO	0.422	6.149	1.197	Positive (<0.001)	Positive (0.066)		
	213 <sup>b,c</sup>	211	TMD V	DIURNAL	1.031	0.702	3.565	Neutral (0.030)	Positive (0.016)	Under positive selection in DIURNAL background	
	277 <sup>b,c</sup>	None	TMD VI	DIURN-DEN	0.549	5.530	0.429	Positive (0.004)	Neutral (0.727)	Under positive selection in DIURN-DEN foreground	

Foreground groups included "DENDRO", that is Dendrobatidae + stem branch, "DIURNAL", that is all diurnal branches as described in methods, or DIURN-DEN, which is all diurnal branches except for Dendrobatidae and its stem branch. A single site-specific synonymous rate (dS or  $\alpha$  in the models) was estimated for each site using maximum likelihood. Two site-specific nonsynonymous rates (dN or  $\beta$  in the models) were estimated separately for the foreground and background and then compared statistically using likelihood ratio tests. We report only the sites that pass the false discovery threshold (q = 0.02). Results from FEL analyses are presented for each site identified to be under different selection regimes by Contrast-FEL. Selection patterns were estimated by FEL for specific subsets of branches using each foreground group (DENDRO, DIURNAL, or DIURN-DEN) or its background (i.e., all branches except DENDRO, DIURNAL, or DIURN-DEN) as foreground lineages. FEL results are reported as Neutral:  $\omega$  not significantly different than 1; Positive:  $\omega$  significantly greater than 1; Negative:  $\omega$  significantly less than 1. We aligned sequences against bovine rhodopsin (NP\_001014890.1) to make comparisons with known spectral tuning sites (see supplementary table S3, Supplementary Material online for references).

<sup>a</sup>The numbering system used in our alignments can be determined by adding 17 to the bovine *RH1* number for *LWS*, adding 9 for *SWS2*, and subtracting 5 for *SWS1*. However, at site 342 in *SWS1*, a subtraction of 6 is required.

<sup>b</sup>Amino acid sites reported by Schott et al. (2022) as under positive selection.

<sup>c</sup>Amino acid sites reported by FUBAR, FEL, or MEME analyses as under positive selection in this study, across the entire tree.

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of the same sites found to be evolving under significantly different regimes in the DIURNAL foreground and background; only site 213 in *RH1* was unique to the DIURNAL analysis. Interestingly site 213 in *RH1* was identified to be under positive selection in the DIURNAL background (i.e., the nocturnal lineages) and neutrally evolving in the foreground, suggesting that it might be under selection in nocturnal lineages. In addition, two sites were found to be evolving under different regimes in the DENDRO foreground and background but were not identified in the DIURNAL analyses (site 316 in *LWS* and site 97 in *RH1*), suggesting that these sites are under different selective conditions in Dendrobatidae compared to other lineages.

To further assess whether the amino acid sites identified to be under selection in diurnal lineages were driven by our biased sampling of Dendrobatidae and to potentially identify amino acid sites under selection in nondendrobatid diurnal lineages, we then created a third group ("DIURN-DEN"), which is the DIURNAL group excluding Dendrobatidae and its stem branch. In these analyses, three sites were identified to be under differential selective regimes (318 in LWS and 22 and 277 in RH1), and they did not overlap with any sites identified to be under differential selective regimes when using the DIURNAL or DENDRO foregrounds (table 1). Two sites were found to be neutrally evolving in the foreground but under negative selection in the background (318 in LWS and 22 in RH1) and one site was found to be under positive selection in foreground but neutrally evolving in background (277 in RH1). Thus, these sites could be related to diurnality in nondendrobatid clades. Based on our Contrast-FEL results, we note that caution should be taken when contrasting branches for selection pattern analyses based on a phylogeny that has multiple independent origins of a trait. The uneven distribution of taxa (e.g., over-representation of clades with the derived character of interest-in our case Dendrobatidae) might drive the analyses to identify sites with a signature of selection exclusive to the overrepresented group as significant in overall selection analyses.

Next, we aimed to determine whether other amino acid sites experienced positive selection along the entire phylogeny, with the goal of identifying sites of functional importance for anurans and potentially for diurnal vision. We conducted four types of selection tests that detect positive selection at specific sites (FEL, FUBAR, CODEML) or branch-site combinations (MEME). In CODEML analyses, the M8 model including positive selection was a better fit than M7 and M8a models (which exclude positive selection) for LWS and RH1 but not for SWS1 or SWS2 (see supplementary table S4, Supplementary Material online). Thus, we only report CODEML results for RH1 and LWS. In combination with Contrast-FEL results, site and branch-site selection tests identified a total of 44 sites across the four opsin genes as having experienced positive selection across all branches (FEL, FUBAR, CODEML) or in a subset of branches (MEME, Contrast-FEL) (tables 1 and 2). Of the identified sites, 14 were in LWS, 6 in SWS1, 4 in SWS2, and 20 in RH1. Based on published data, 3 of the 44 sites are

known spectral tuning sites including site 217 in LWS and sites 124 and 164 in RH1; 12 other sites are within three amino acids from a known spectral tuning site (table 2; supplementary Data S1, Supplementary Material online; supplementary table S3, Supplementary Material online). While results from FEL, FUBAR, and CODEML reflect signatures of positive selection that are strong enough to provide enough signal across the entire phylogeny, MEME detects signals of selection in subsets of branches and thus provides more precision about where in the phylogeny sites are under positive selection. Most sites identified by MEME were found to be under positive selection in only one or two branches, but 13 (sites 49, 162, 205, and 262 in LWS, sites 159 and 221 in SWS1, and sites 87, 107, 124, 168, 169, 213, and 277 in RH1) were found to be under positive selection in three or more branches, suggesting that these sites may be of particular functional importance in frogs (fig. 1 and supplementary S1, Supplementary Material online).

Among the list of sites in table 2, a few present complex characterizations. For example, site 277 in RH1 was reported to be under positive selection by MEME, but we observed that there were no nonsynonymous changes on the branches identified by MEME (Atelopus spurrelli and Node 33, the ancestral branch leading to A. spurrelli, Atelopus glyphus, Atelopus limosus, and Atelopus varius; fig. 1) Instead, these branches have two mutations that do not result in an amino acid substitution (Ser-AGC  $\rightarrow$  Ser-TCC), but likely require a nonsynonymous intermediate that is not recovered in our analysis. Another six sites were found by MEME to be under positive selection in a subset of branches and by FEL to be under negative selection across the entire phylogeny (table 2, denoted with "FEL (-)"). Five of these sites are characterized by high conservation, with an amino acid change present in only one or two branches (fig. 1 and supplementary S1, Supplementary Material online). In contrast, the sixth site, site 124 in RH1, presents an alanine-to-serine replacement in eight branches. Finally, one site in LWS, two sites in SWS1, and one site in RH1 (table 2) were found to be under positive selection by MEME but without any branch identified, suggesting that the signal for selection is diffuse, which occurs when there is enough signal to report positive selection, but no individual branch rises to the level of significance (Spielman et al 2019).

#### Verifying Absence of SWS2 in Poison Frogs

During our exome capture reconstructions, we failed to recover any exon or fragment of the SWS2 gene in dendrobatids. We considered at first that our baiting design might have been unsuccessful. While it is possible that our SWS2 sequence baits (designed using a consensus sequence matrix) were not similar enough to capture this gene in Dendrobatidae, especially given that part of SWS2 was not captured efficiently in other frogs, we had no issues obtaining partial sequences from all other clades including the other aposematic species of *Atelopus* and *Brachycephalus* (see supplementary fig. S1, Supplementary Material online). Therefore, we needed to confirm that this gene was not

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### Table 2. Results From Site and Branch-site Selection Analyses.

Gene	Known spectral tuning sites in vertebrates	Codon number (bovine <i>RH1</i> numbering) <sup>a</sup>	Nearby spectral tuning site	Location	Selection test	P-value/ Posterior Probability	Number of branches (MEME only)
LWS	100, 164, 181, 214, 217, 261, 269, 292, 293	49 <sup>b</sup>	None	TMD I	MEME	P < 0.01	3
					CODEML	PP = 1.00	_
		59	None	TMD I	MEME	P = 0.02	1
					FEL (—)	P = 0.04	_
		162	164	TMD IV	MEME	P < 0.01	7
		166 <sup>b</sup>	164	TMD IV	MEME	P < 0.01	1
					FUBAR	PP = 0.99	_
					CODEML	PP = 0.97	_
		176	None	TMD IV	MEME	P < 0.01	2
		170	None	inte it	FEL (–) <sup>c</sup>	P < 0.01	_
		205	None	TMD V	MEME	P = 0.04	8
		205	None	TMD V	MEME	P < 0.01	1
		200	None	TMD V	MEME	P < 0.01 P = 0.02	2
		209 217 <sup>b</sup>					
			217	TMD V	MEME	P = 0.04	0
		231	None	TMD V	MEME	P = 0.02	1
		2.00	266	TAAD 17	FEL (—)	P < 0.01	-
01110 x		262	261	TMD VI	MEME	P < 0.01	3
SWS1	46, 49, 52, 81, 86, 90, 91, 93, 97, 108, 109, 113,	50	49, 52	TMD I	MEME	P = 0.04	0
	114, 116, 118, 207				FEL (+)	P = 0.03	-
		ь.			FUBAR	PP = 0.97	-
		120 <sup>b</sup>	118	TMD III	MEME	P = 0.04	0
					FEL (+)	P = 0.02	—
		159 <sup>b</sup>	None	TMD IV	MEME	P = 0.01	6
		221	None	TMD V	MEME	P < 0.01	5
SWS2 (Only sites up to	44, 46, 49, 52, 91, 93, 94, 97, 99, 109, 116, 117,	-1	None	N-T	FUBAR	PP = 0.90	_
129 were analyzed)	118, 122, 164, 207, 261, 265, 269, 275, 292, 295	8	None	N-T	FUBAR	PP = 0.98	_
		56	None	TMD I	FUBAR	PP = 0.98	_
		106	None	ECD I	MEME	P = 0.02	1
					FEL (—)	P = 0.01	_
RH1	83, 90, 96, 102, 118, 122, 124, 125, 132, 164,	37	None	TMD I	CODEML	PP = 1.00	_
	183, 189, 194, 195, 207, 208, 211, 253, 261, 265,	39 <sup>b</sup>	None	TMD I	MEME	P = 0.03	0
	269, 289, 292, 295, 299, 300, 308				FEL (+)	P = 0.02	_
					FUBAR	PP = 0.97	_
		82	83	TMD II	MEME	P < 0.01	2
		87	None	TMD II	MEME	P = 0.03	7
					FEL (+)	P = 0.03	_
					FUBAR	PP = 0.95	_
		97 <sup>b</sup>	96	TMD II	FEL (+)	P = 0.05	_
					FUBAR	PP = 0.95	_
		103	102	ECD I	CODEML	PP = 1.00	_
		107 <sup>b</sup>	None	ECD I	MEME	P = 0.04	9
		111	None	TMD III	MEME	P < 0.04	2
			122, 124,	TMD III	MEME		8
		124				P = 0.03 P = 0.04	
		150	125 None		FEL (-)	P = 0.04	-
		159	None	TMD IV	MEME	P < 0.01	2
		164	164	TMD IV	CODEML	PP = 1.00	-
		168	None	TMD IV	MEME	P = 0.03	3
		169 <sup>6</sup>	None	TMD IV	MEME	P < 0.01	12
					FEL (+)	P < 0.01	—
		a h		<b>T</b> 115 · ·	FUBAR	PP = 1.00	-
		213 <sup>b</sup>	211	TMD V	MEME	P < 0.01	9
					FUBAR	PP = 0.97	-
		260	261	TMD VI	MEME	P < 0.01	1
		262	261	TMD VI	MEME	P = 0.01	1
		-			FEL (—)	P = 0.01	_
		277 <sup>b</sup>	None	TMD VI	MEME	P = 0.01	3

(continued)

Gene	Known spectral tuning sites in vertebrates	Codon number (bovine RH1 numbering) <sup>a</sup>	Nearby spectral tuning site	Location	Selection test	P-value/ Posterior Probability	Number of branches (MEME only
		281	None	ECD III	CODEML	PP = 1.00	_
		290	289, 292	TMD VI	MEME	P = 0.02	2
					FUBAR	PP = 0.95	_

Amino acid sites in four opsin genes that were detected to be under positive selection by site (FEL, FUBAR, CODEML) or branch-site combinations (MEME). We aligned sequences against bovine rhodopsin (NP\_001014890.1) to make comparisons with known spectral tuning sites (see supplementary table 53, Supplementary Material online for references). More details regarding the phylogenetic patterns of selection can be reviewed in figure 1 and supplementary 51, Supplementary Material online. We report all significant FEL results that correspond to sites identified by MEME or FUBAR and note each FEL result as indicative of positive (+) or negative (-) selection. Sites identified to be under positive selection in CODEML analyses are reported as posterior probabilities from Bayes Empirical Bayes analysis of M8 models. N-T, N-terminus; C-T, C-terminus; ECD I, extracellular domain I; TMD I–VII, transmembrane domains I–VII; –, does not apply. Reported *P*-values for MEME have been corrected for multiple testing using Holm–Bonferroni. Input files and results are available in supplementary Data S1, Supplementary Material online.

<sup>a</sup>The numbering system used in our alignments can be determined by adding 17 to the bovine *RH1* number for *LWS*, adding 9 for *SWS2*, and subtracting 5 for *SWS1*. <sup>b</sup>Amino acid sites reported by Schott et al. (2022) as under positive selection.

present with additional transcriptomic and genomic data. For the transcriptome approach, we extracted mRNA from the *O*. *pumilio* eye, and we were unable to amplify SWS2 cDNA. This suggests that at least in this species, SWS2 is not expressed.

For the genomic approach, we used synteny analyses with BLAST, and we found that SWS2 is located in a syntenic block with LWS and MECP2 in the ancestor of all tetrapods, in the ancestor of all amniotes (fig. 2A), and in five frog families with genomic data that is publicly accessible (fig. 2B: Bufonidae, Leptodactylidae, Eleutherodactylidae, Pipidae, and Ranidae). For dendrobatids, we explored the reassembled O. pumilio scaffolds and found that scaffold70671 contained LWS and was syntenic with other amphibians (two short and highly conserved regions between SWS2 and LWS could be aligned between Nanorana and O. pumilio [supplementary Data S2, Supplementary Material online]), but we could not identify any coding region of SWS2. Then, we explored the more complete scaffold of this region from R. imitator (scaffold934), which included LWS and MECP2, the latter of which is expected to be upstream of SWS2, yet we were unable to find SWS2 between these two genes on this scaffold (fig. 2B). In fact, we could not detect an SWS2 gene or pseudogene on any scaffold in O. pumilio or R. imitator genomes (see supplementary table S5, Supplementary Material online). In contrast, using the same methods, we were able to detect SWS2 upstream of LWS in other hyloid frogs including Bufo, Eleutherodactylus, Engystomops, and Rhinella (fig. 2B, supplementary Data S4, Supplementary Material online). The hyloid clade originated  $\sim$ 70 Ma and the age of the last common ancestor of Dendrobatidae has been estimated to be  $\sim$ 40 My (Feng et al. 2017). Thus, the most parsimonious explanation is that a functional SWS2 gene is not present in living dendrobatids, including O. pumilio or R. imitator, and that the SWS2 gene may have been pseudogenized (to a point beyond recognition) or lost in the ancestor of dendrobatids between 70 and 40 Ma.

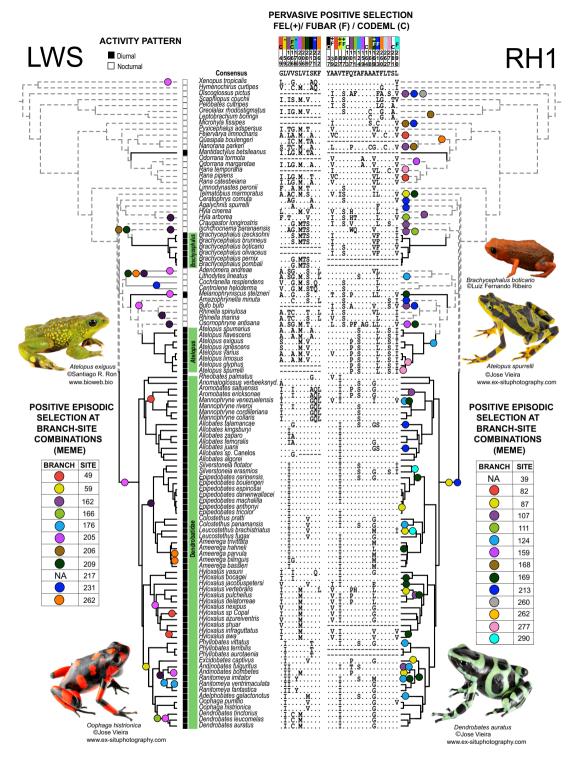
### Discussion

In this study, we reviewed the evolution of the visual opsin gene repertoire (*LWS*, *SWS1*, *SWS2*, and *RH1*) in anurans by

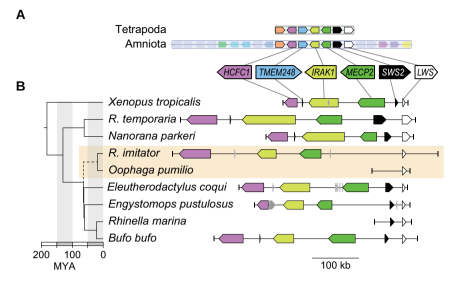
analyzing a dataset of 116 frog species from 20 families, including 78 diurnal and 38 nocturnal species. Our analyses report 44 amino acid sites as under positive selection in our sampling of Anura, including sites in or near locations that have been implicated in visual tuning. Many of these sites are undergoing positive selection in branches within diurnal and aposematic Neotropical frog clades. Moreover, our results provide evidence for the loss of the SWS2 gene in the ancestor of Dendrobatidae between 70 and 40 Ma, an event that coincides with the evolution of diurnal activity patterns in this clade. Diurnality in poison frogs has been hypothesized as a prerequisite for the origin of many of their unique adaptations including aposematism, audiovisual communication, mating choice, and parental care (Santos et al. 2003; Santos and Grant 2011; Yang et al. 2019; Carvajal-Castro et al. 2021).

# Selection Patterns Relating to the Transition to Diurnality in Neotropical Anurans

Initially, we aimed to identify sites and patterns of selection that were associated with the transition to diurnality in the three focal anuran clades sampled for this study (Dendrobatidae, Atelopus and Brachycephalus). However, we found no evidence for unique patterns of selection on the three branches where transitions to diurnality occurred in our dataset compared to other branches in our tree. Nevertheless, we were able to parse apart some of the patterns in Dendrobatidae and other diurnal lineages and propose the following. Seven sites (176 in LWS, 218 and 221 in SWS1, and 87, 124, 169, and 277 in RH1) were found to be under positive selection in diurnal lineages including Dendrobatidae, and two sites (318 in LWS and 22 in RH1) were found to be under positive selection in diurnal lineages excluding Dendrobatidae; these nine sites may be associated with adaptation to diurnality. For example, site 124 in RH1 (a known spectral tuning site) was found to be under positive selection in diurnal lineages and in dendrobatid lineages but under negative selection in other branches. This site was also found to be under positive selection by MEME in eight branches (R. imitator, Adelphobates galactonotus, Andinobates fulguritus, Phyllobates vittatus, Colostethus panamansis,



**FIG. 1.** A summary of activity trait data (see supplementary table S1, Supplementary Material online for references) and results from site and branch-site selection analyses of two opsin genes, *LWS* and *RH1* (see supplementary fig. S1, Supplementary Material online for *SWS1* and *SWS2* and table 2 for details) using bovine *RH1* numbering. Branch-site combinations that were detected by Mixed Effect Models of Evolution (MEME) as evolving under positive selection are mapped onto the phylogenetic tree using filled circles where circle colors correspond to specific sites in the alignment. One site in *LWS* and one in *RH1* were significant in MEME analyses but no specific branch-site combination was reported; these are marked by "NA" in the MEME tables. Amino acid sites marked with "F" or "C" above the alignment were reported to be under positive selection by FUBAR or CODEML, respectively. FEL results are indicated using + to indicate positive selection and – to indicate negative selection (negative selection results are only shown for sites found to be under positive selection by other analyses; see supplementary Data S1, Supplementary Material online for full FEL results). We abbreviated *Anomaloglossus verbeeksnyderorum* as *Anomaloglossus verbeeksnyd*. for brevity. Black branches indicate branches categorized as diurnal.



**Fig. 2.** (A) SWS2 is found in a syntenic block with *LWS* in the ancestors of all tetrapods (block 355) and in the ancestor of all amniotes (block 65). (B) The region in *Xenopus tropicalis* (chromosome 8, NC\_030684:32108538-32360324) shows synteny with *Rana* (*R.*) *temporaria* (chromosome 9, NC\_053497.1:22641469-22170306; Ranidae), *Nanorana parkeri* (unplaced scaffold, NW\_017306743:511492-806940; Ranidae), *Rhinella marina* (contig ctg22529\_RHIMB, ONZH01019223.1:1-80000; Bufonidae), *Bufo bufo* (chromosome 8, NC\_053396.1:25948637-26357822; Bufonidae), *Eleutherodactylus coqui* (chromosome 9, CM034094.1:11013102-11351932; Eleutherodactylidae), *Engystomops pustulosus* (chromosome 10, CM033650.1:68367373-68196643; Leptodactylidae), which also all contain *SWS2*, *MECP2*, *IRAK1*, *TMEM187*, and *HCFC1* upstream of *LWS*. However, *SWS2* is absent in the scaffold containing *LWS* in two dendrobatid genomes (*Oophaga pumilio* [scaffold70671:1-49101] and *Ranitomeya* [*R*.] *imitator* [CAJOBX010072427.1, scaffold934:531527-1]). A dated chronogram (Feng et al. 2017) indicates that *SWS2* may have been lost in the ancestral lineage (dashed line) leading to Dendrobatidae (orange box), between ~70 and 40 Ma. Where coding regions were identified in genome assemblies but not annotated, we added annotations using blastp and blastn. Coding regions in gray had ambiguous blast results and thus were left without annotations.

Lithodytes lineatus, Mannophryne venezuelensis, and the branch leading to the Atelopus clade). The amino acid type at site 124 of those species is serine instead of alanine as seen in other anuran species. Mutagenesis experiments in bovine RH1 indicate that alanine-to-threonine and alanine-to-serine substitutions at this position provide a slight blue shift (Lin et al. 1998; Castiglione and Chang 2018). In addition, A124S in combination with L119F significantly increases the stability of the active conformation of bovine RH1 (Castiglione and Chang 2018). Therefore, it is possible that an anuran RH1 with the A124S substitution has a blue-shifted  $\lambda_{max}$  and that such changes may be associated with adaptation to diurnality as has been suggested in species of diurnal snakes (Schott et al. 2016; Hauzman et al. 2017). The loss of the cone opsin RH2 in the ancestor of all amphibians resulted in a decreased sensitivity to a significant portion of the visual spectrum, mainly due to the absence of overlap between LWS and SWS1. However, a blue shift in the absorption spectrum of RH1 would coincide with the wavelength detection range previously covered by RH2. This shift would enhance chromatic discrimination in diurnal clades by providing more uniform coverage across the color spectrum (Schott et al. 2016). Previous studies have suggested the expression of RH1 in the green-sensitive cones of amphibians (Schott et al. 2022), supporting a potential role for such a blue shift.

Our failure to identify sites under differential selection in branches where the transition to diurnality occurred suggests either that we may have limited power to detect such sites and/or that relatively stronger selection occurred following the transition rather than during it. We identify many sites under selection within the three diurnal clades on which we focused (i.e., poison frogs, Atelopus and Brachycephalus), even if these were not found to be specifically associated with diurnality in Contrast-FEL assessments. As many diurnal animals rely on color-based signals (i.e., detecting ripe fruits; [Melin et al. 2009]) or perceiving sexual signals (Carleton et al. 2005), they are expected to have a suite of opsins that are tuned to perceive the relevant signal under the corresponding light environment while balancing other aspects of protein function (e.g., kinetics; Hauser et al. 2017). Thus, our selection results suggest that the transition to diurnality is complex and idiosyncratic such that each lineage differently accumulated changes to accommodate the diel habit. In other words, the opsin genes in each clade experienced the process of visual tuning independently and may have been influenced by other confounding factors (e.g., mating visual signal recognition); not surprisingly, this resulted in some parallel and some different substitution patterns.

## Identification of Putative Functional Sites in Anuran Opsin Genes

Tables 1 and 2 provide an extensive list of opsin amino acid sites putatively involved in frog vision. Changes at these sites may work alone or together to shift the wavelength sensitivity of anuran opsin proteins as they adapt to diverse light environments, including diurnality. The sites inferred to be under positive selection occur predominantly in the transmembrane domains. Such domains impact the tertiary structure, thermal stability, and properties of the retinal-binding pocket (Andrés et al. 2001; Yokoyama et al. 2006). Spectral sensitivity is related directly to interactions between the amino acid residues in the transmembrane domains and the chromophore (Yokoyama et al. 2006). Although it is possible that replacements at these sites might influence spectral tuning in frogs, without more experimental data, their implications are difficult to interpret. For this reason, we conservatively discuss their possible role in anuran vision below.

Of the 44 sites we found to be under positive selection, only three are previously known spectral tuning sites in vertebrates. Site 124 in RH1 is discussed above. Site 217 in LWS (233 in the human LWS) has been previously shown to play a role in the differentiation of the spectral sensitivities between the long-wave-sensitive and the middlewave-sensitive pigments in humans and other mammals (Yokoyama and Yokoyama 1990; Asenjo et al. 1994; Fasick and Robinson 1998; Yokoyama and Radlwimmer 2001, but see Hiramatsu et al. 2004). In our study, MEME detected positive selection on this site without indicating any specific branches, suggesting a diffuse signal of selection at this site across the tree (fig. 1; table 2). Moreover, this site was also reported by Schott et al. (2022) to be under positive selection, confirming that this is a key site that could be responsible for shifting the sensitivity of LWS in frogs, including the diurnal taxa explored in this study. In our alignment, amino acid substitutions at site 217 are present in several species including several diurnal clades such as the dendrobatid genera Mannophryne (whose females have a colorful yellow collar), Rheobates, Aromobates, and toxic Phyllobates, as well as two species of Atelopus. However, this site was not identified in Contrast-FEL analyses as under positive selection specifically in diurnal lineages.

The third known spectral tuning site, 164 in RH1, was found by CODEML to be under positive selection; other analyses did not identify this site to be under selection. In our data set, 102 species at this position had an alanine, while just Osornophryne antisana expressed a glycine. Substitutions at this site are known to contribute to a red shift in the absorption maxima of bovine RH1 (Chan et al. 1992). Although an A164S substitution seems to result in a rather small (2 nm) red shift in absorption, it has been shown that an additive effect is achieved when it is in combination with other substitutions including F261S and A269T (Chan et al. 1992). Thus, it is difficult to predict how changes at this site might impact the spectral sensitivity of O. antisana or other frog species, and our analyses do not offer strong support for the involvement of site 164 in RH1 in adaptation to diurnality.

Considering the general lack of data on spectral tuning sites in frog opsins, it is possible that other sites we identified to be under positive selection (tables 1 and 2) could be directly or indirectly involved in spectral tuning in anurans.

Eleven sites found to be under positive selection are not known spectral tuning sites but are located within three amino acids of a known spectral tuning site: 162, 166, and 262 in *LWS*; 50 and 120 in *SWS1*; 82, 97, 103, 213, 260, 262, and 290 in *RH1*. Additionally, 13 sites, including three at or near spectral tuning sites, were found to be under selection in three or more branches: 49, 162, 205, and 262 in *LWS*; 159 and 221 in *SWS1*; 87, 107, 124, 168, 169, 213, and 277 in *RH1* (table 2, fig. 1). We speculate that some of these sites may be of functional importance in the vision of frogs. For example, site 169 in *RH1* was found to be under positive selection in 12 lineages. Experiments using site-directed mutagenesis could help verify which sites are important for spectral tuning in amphibians.

A prior study (Schott et al. 2022) identified 16 sites in opsin genes of anurans to be under positive selection; we found 11 of these to be under positive selection in our study. We failed to identify five sites (65, 212, and 270 for *RH1*; 154 for *LWS*; -2 for *SWS2*), which might reflect differences in the number of taxa in our studies (i.e., 116 in our study vs. 33 in theirs) or the specific focus on diurnal species in this article. Schott et al. (2022) also discussed additional variation in other known spectral tuning sites. We exclude them from our discussion because we could not find evidence of positive selection on these sites. Our comparison depicts the conserved nature of opsins but also reports new amino acid changes that contribute to opsin spectral sensitivity.

## Visual Signals in Focal Frog Clades

Due to the selective pressure of diurnal and visually guided predators, many diurnal frogs are adorned with colorful visual signals that evolved as part of a defensive strategy known as aposematism, in which conspicuous signals warn predators of prey defenses (Bell and Zamudio 2012). Each of our three focal clades (i.e., poison frogs, Atelopus and Brachycephalus) contain species that possess aposematic signals. Although aposematic signals initially evolve under selective pressure from predators and might intensify over evolutionary time (Mappes et al. 2005; Sherratt 2008; Loeffler-Henry et al. 2023), warning signals can also become entangled with mating behaviors and further diversify under sexual selection pressures (Cummings and Crothers 2013; Rojas et al. 2018), if the mate recognition and visual sensory system is appropriately tuned to the visual cues. We hypothesize that some of the sites identified to be under selection are related to intraspecific signaling within these aposematic clades. For instance, color perception seems to be key for mate recognition and territorial display in dendrobatids. In the aposematic species O. pumilio, tadpoles imprint on their mother's color (Yang et al. 2019), females show assortative mate preferences (Summers et al. 1999; Reynolds and Fitzpatrick 2007; Maan and Cummings 2008; Yang et al. 2016), male-male competition relies on color-mediated aggressive behaviors towards phenotypically similar rivals (Yang et al. 2018; Yang and Richards-Zawacki 2020), and there may be directional sexual selection on male coloration brightness (Maan and Cummings 2009). Assortative mating by coloration and pattern is present but less pronounced in other dendrobatid species including additional *Oophaga* species (*Oophaga histrionica* and *Oophaga lehmanni*; Medina et al. 2013), species of *Mannophryne* with sexual dichromatism where throat coloration in males (gray) and females (bright yellow) plays a role in territoriality and mate choice (Greener et al. 2020), and in many species of *Allobates* that are sexually dimorphic for throat color that is visible in vocalizing males.

In our other focal clades, visual signaling has been investigated in only a few species. For example, Atelopus zeteki appears to use limb motions as visual signals to conspecifics and this might be related to their lack of tympanic middle ear (Lindquist and Hetherington 1998); yet, it is unknown if other Atelopus use their conspicuously colored soles for intraspecific communication or aposematic signals (Rößler et al. 2019). Likewise, at least one species of Brachycephalus is inferred to be aposematic based on experimental evidence (Goutte et al. 2019; Reboucas et al. 2019); Brachycephalus ephippium and Brachycephalus pitanga also have fluorescent dermal bones visible through their skins with a potential function as a signal (Goutte et al. 2019). While territorial males of some Brachycephalus species display a foot-waving behavior to warn off encroaching males similar to some species of Atelopus (Pombal et al. 1994), it is unclear how important a role color and/or contrast play in the display. Future studies could investigate species- or population-level variation in opsin sequences to assess whether the sites listed in tables 1 and 2 have any effect on the recognition or evaluation of intraspecific signals.

# Loss of SWS2 in Dendrobatidae and the Implications for Their Visual Ecology

Based on our inability to sequence SWS2 from an O. pumilio eye transcriptome, the complete absence of SWS2 from bait-capture data from all sequenced dendrobatids (60 species representing all major lineages), and our failure to identify any trace of an SWS2 gene or pseudogene in the O. pumilio and R. imitator genomes, we hypothesize that dendrobatids lost this gene early in their history. Such gene-loss events are not surprising as other comparative genomic studies have shown (Borges et al. 2015; Xu et al. 2021). The loss of short-wavelength photopigment genes has occurred multiple times in the evolutionary history of vertebrates and coincides with shifts in activity patterns, habitat occupancy, and the evolution of other aspects of the sensory capacity of animals (Bowmaker 2008; Jacobs 2013). For example, the loss of SWS2 in therian mammals and coelacanths is considered an adaptation to nocturnality and deep-sea environments, respectively. The loss of a functional SWS1 pigment in several bat species coincides with the origin of a specialized form of echolocation (i.e., high-duty-cycle echolocation; Jacobs 2013).

In dendrobatids, it remains unclear how the loss of SWS2 may have impacted their vision and visual ecology. To answer such a general question, we can explore some specific ones. First, how has SWS2 loss potentially impacted the physiology of dendrobatid visual systems? The limited empirical data from anurans suggest that SWS2 expression is restricted to green rods (Yovanovich et al. 2017; Mohun and Davier 2019). However, green

et al. 2017; Mohun and Davies 2019). However, green rods are absent from the *O. pumilio* retina (Siddiqi et al. 2004). Thus, the loss of SWS2 and of the green rod cell type suggests that under low-light levels, dendrobatids might be less capable of discriminating color (scotopic vision) than other diurnal frogs.

Second, when and how did SWS2 disappear from the genome of ancestral dendrobatids? Based on our results, we could not find a SWS2 pseudogene (i.e., nonfunctional segments of DNA that resemble functional SWS2) in the available genomic data for the poison frog species R. imitator and O. pumilio. In contrast, close relatives of dendrobatids do have a functional SWS2 (fig. 2). Thus, our data suggest that only dendrobatids lost SWS2. Dendrobatidae is nested within the superfamily Hyloidea, whose last ancestor lived about  $\sim$ 70 Ma (Feng et al. 2017; Hime et al. 2021). As the approximate age of the crown clade of Dendrobatidae is ~40 Ma (Feng et al. 2017), SWS2 must have been lost between 40 and 70 Ma. Two alternative explanations exist for the loss of SWS2 during this 30-My timeframe: (1) The ancestor of dendrobatids had a functional gene which pseudogenized and changed beyond recognition during that period or (2) SWS2 was lost without pseudogenization. Given the lack of evidence of SWS2 pseudogene relics, the most parsimonious explanation is a complete deletion of the gene. Additional data may show that other hyloid groups have lost SWS2, but until then, we consider the loss of SWS2 as a synapomorphy of Dendrobatidae (i.e., Aromobatidae + Dendrobatidae sensu Grant et al. [2006]).

Third, was the loss of SWS2 consequential in that it affected other aspects of dendrobatid vision? Other diurnal hyloid clades in our dataset do not appear to have lost SWS2: all Atelopus and Brachycephalus frogs maintain this gene. Some uncertainty remains regarding SWS2 functionality in nondendrobatids because we were unable to recover the entire gene from many species using target-bait capture, yet complete SWS2 sequences are available in assembled genomes (e.g., fig. 2, Eleutherodactylus). Whether diurnal hyloid species have the green rod cell type (which expresses SWS2) remains unknown. Nevertheless, dendrobatids are known to differ in at least two visual properties from other hyloids. First, the lenses of some dendrobatid species (O. pumilio, Epipedobates tricolor, Dendrobates auratus, Dendrobates leucomelas, Allobates femoralis, and Adelphobates castaneoticus) transmit less short-wavelength light (lens  $\lambda_{t50}$  413–425 nm; except for D. leucomelas lens  $\lambda_{t50}$ 326 nm) than those of bufonids (R. marina, R. icterica, R. ornata, Bufo bufo, Rhaebo guttatus, Sclerophis maculata, and Atelopus varius lens  $\lambda_{t50}$  331–365 nm) and brachycephalids (Brachycephalus rotenbergae, Ischnocnema parva,

Ischnocnema henseli lens  $\lambda_{r50}$  314–356 nm) (Donner and Yovanovich 2020; Yovanovich et al. 2020; Thomas et al. 2022). As the typical peak wavelength sensitivity of SWS2 rods (~430 nm; Yovanovich et al. 2017) is much closer to the limit of lens transmission in dendrobatids than other frogs, it is plausible that lens transmission properties were adjusted in some species following the loss of SWS2, or vice versa. Additionally, in O. pumilio, the short-wavelength-sensitive cone (likely SWS1), which absorbs light wavelengths  $\sim$ 430 nm in other anurans (Yovanovich et al. 2017), was found to absorb light at 466 nm (Siddigi et al. 2004), suggesting that it has undergone a shift in spectral sensitivity. Further investigation of these patterns will clarify whether loss of SWS2 in dendrobatids in combination with a diurnal lifestyle led to altered lens transmission properties and a change in SWS1 wavelength sensitivity, or vice versa.

Despite the apparent loss of SWS2 in dendrobatid frogs and the lack of RH2 pigment in amphibians (Mohun and Davies 2019), microspectrophotometry data revealed an MWS photoreceptor in the O. pumilio retina (Siddiqi et al. 2004). In accordance with prior literature (Schott et al. 2022), we speculate that these frogs are instead using RH1 in those photoreceptors as the microspectrophotometry absorbance graphs for rods and MWS cones appear nearly identical (Siddigi et al. 2004). Whether this is the case and if there are any modifications to the RH1 pigment or to any proteins in the O. pumilio MWS cone phototransduction pathway required to make the rod protein function in cone cells requires further investigation. We did find that a large number of amino acid sites (i.e., 20) are under selection in RH1 within frogs, which could be a result of this dual function, though only a few of these sites were noted to be specifically under selection in Dendrobatidae or experiencing dendrobatid-specific patterns of selection. Further, our genome mining, transcriptome, and synteny analyses do not provide any evidence suggesting that RH1 has been duplicated in frogs, but at least one species (Pyxicephalus adspersus) has a duplication of LWS (Schott et al. 2022). Empirical data of spectral sensitivity and opsin protein function in frogs are sparse, and further studies using microspectrophotometry of isolated frog rods and cones will be necessary to recapitulate the evolution of vision in frogs (Donner and Yovanovich 2020).

### Conclusion

Our results fill gaps in our knowledge by illustrating how diversification in ecology and life history may have affected opsin evolution in amphibians. We uncover evidence of new putative tuning sites in frogs and show strong evidence suggesting that poison frogs have lost the *SWS2* gene. Further work is needed to elucidate the functional consequences of its loss, and of the potential role of *RH1* in facilitating color vision in some frogs. Like Donner and Yovanovich (2020), we expect that additional studies of opsin evolution in amphibians will reveal noncanonical visual adaptations and broaden our understanding of the many ways in which animals adapt to diverse light environments.

## **Materials and Methods**

# Bait Capture Design, Library Preparation, and Sequencing

Using publicly available sequences (see supplementary table S1, Supplementary Material online) and previously generated transcriptomes (Santos et al. 2018), we designed a custom bait-capture array with myBaits (Arbor Biosciences). The 120-bp baits were synthesized at  $>10 \times$  tiling across all exons of the genes of interest. Tissues from representative species of Dendrobatidae, Atelopus, and Brachycephalus and their relatives were obtained from the field or from museum collections (see supplementary table S1, Supplementary Material online for source information), under approved protocols (UT AUP-2012-00032 and AUP-0709210, Austin STRI 200715122207, 2015-00205 Tulane #0453) and collection (001-13 IC-FAU-DNB/MA and permits 001-11 IC-FAU-DNB/MA [Ecuador], IBD0359 Res 1177-2014 and Isla Gorgona PIDB DTPA 020 - 16 Res 061-16 [Colombia], and SE/A-47-07 and CITES export permit SE/A-47-07 [Panama]). Tissue samples from Ecuador and Peru were loaned by Luis A. Coloma (Fundación Otonga and Centro Jambatu de Investigación y Conservación de Anfibios; Ecuador) and César Aguilar (Universidad Nacional Mayor de San Marcos; Lima, Peru). We note there is some controversy over the elevation of subgenera such as Oophaga to genera (Santos et al. 2009; Grant et al. 2017), but we have followed taxonomy following (AmphibiaWeb 2022).

DNA was extracted using Qiagen DNeasy kits (Germantown, MD). DNA guality was reviewed with a 0.8% electrophoresis gel and highly degraded samples were excluded. RNA was removed from extractions with RNAse A (E1008, Zymo Research, Irvine, CA), and extractions were further purified and concentrated with Genomic DNA Clean and Concentrate (D4011, Zymo Research) and then guantified using a Qubit 3.0 fluorometer (Thermo Fisher Scientific, Waltham, MA) following manufacturer protocols. Given the large size of amphibian genomes, we used 400–1600 ng of starting DNA for each sample for library preparation. DNA was sheared to  $\sim$ 300 bp using a Covaris S2 Focused-ultrasonicator (Covaris, Inc, Woburn, MA; settings as Intensity: 5; Duty Cycle: 10%; Cycles per Burst: 200; Time: 50 s; Temp: 7C; Water Level: 12; Sample Volume: 50 µl). Whole-genome libraries were prepared from sheared samples using the KAPA Hyper Plus library preparation kit (KK8514, Roche Diagnostics, Santa Clara, CA), NEBNext Multiplex Oligos for Illumina (E7600, New England BioLabs, Ipswich, MA), home-made SPRI beads (Rohland and Reich 2012) (Lydia Smith, personal communication), and manufacturer protocols. Uniquely barcoded libraries from 4-10 closely related species were pooled to a total of 1.6-5 µg DNA and then size-selected for an insert size of  $250 \pm 25$  bp (total length with 120-bp adaptors:  $345 \pm 25$  bp) using a Blue Pippin (Sage Science, Beverly, MA) and 2% gel cassette. For samples with low concentrations, a second batch of libraries were constructed with the same protocol, except that these were size-selected to  $250 \pm 50$  bp with a 1.5% gel cassette (total length with 120-bp adaptors:  $370 \pm 50$  bp). All size-selected libraries were cleaned with Qiagen MinElute PCR purification kits, eluted in 10 µl, and quantified with Qubit 3.0.

Seven microliter of each pooled library set (50-275 ng/ pool) was then hybridized with custom myBaits biotinylated RNA, Cot-1 DNA, and xGen Universal Blockers-TS Mix (1075474, Integrated DNA Technologies, Redwood City, CA) according to myBaits v4.01 recommended protocols. As amphibian genomes are large and contain many repetitive sequences, we used a large amount of starting DNA (1.6-5 µg at size selection) and a larger quantity of blocking oligos than described in the manufacturer's protocol (8  $\mu$ g human C<sub>0</sub>t-1 and 8  $\mu$ g salmon C<sub>0</sub>t-1 per hybridization reaction). We also used the xGen blockers rather than those provided in the myBaits kit. Following hybridization with  $C_0$ t-1, universal blockers, and baits for 36 h, pooled libraries were washed and cleaned with DynaBeads MyOne Streptavidin C1 beads (65002, Thermo Fisher Scientific). 12-17 µl of resulting libraries were amplified using the NEBNext Illumina primers (that came with the multiplex oligos) and the KAPA Library Amplification Kit (KK2611, Roche Diagnostics) in two separate PCR reactions with 15-17 cycles. These were purified and eluted in 10 µl with the Qiagen MinElute purification kit. Duplicate PCR reactions and libraries from all 97 samples were normalized by concentration and according to the number of samples per pool and then sequenced across two lanes of the Illumina HiSeq4000 at the Genome Sequencing and Analysis Facility (GSAF) at University of Texas at Austin, yielding approximately 4 M reads per sample.

#### Assembly of Opsin Genes

Read quality was checked with FastQC (Andrews 2010), barcodes were excluded with Trimmomatic (Bolger et al. 2014) and, because combining multiple assembly methods better characterize multigene families (Holding et al. 2018), the reads were assembled with default parameters for de novo with MEGAHIT (Li et al. 2015), Trinity (Grabherr et al. 2011), and SPAdes (Bankevich et al. 2012). Final assemblies were reduced using CD-HIT set to >98% similarity. Reduced assemblies were annotated using a custom library generated with the opsin sequences used to design the baits and BLASTX. Reference sequences for BLASTX were derived from amphibian opsin genes available in GenBank: X. laevis (Pipidae), Nanorana parkeri (Dicroglossidae) and other species with opsin sequences (e.g., Rana catesbeianus [Ranidae]). Any sequence matching one of the reference opsin genes with an e-value  $<10^{-6}$  was pulled out for downstream analyses.

Given that we de novo assembled genomic DNA that was captured using baits designed from mRNA, the sequences were often only partially assembled, and some shorter sequences were identified erroneously by BLASTX. We used the program BLAT v.36  $\times$  2 (Kent 2002) with reference sequences from the Nanorana parkeri genome (LWS, XM\_018560714.1; SWS1, NW\_017306744.1; SWS2, NW 017307939.1; RH1, NW 017306456.1) to verify sequences from our dataset that were putatively identified as opsins using BLASTX as described above. A BLAT server was prepared using N. parkeri reference sequences with the "-trans" option to translate the database into protein for all six reading frames. Then BLAT was run with options "-t = dnax" and "-q = dnax" to specify that the format of the database and query were both DNA sequences translated in six frames to protein. Sequences that matched N. parkeri references were pulled out from the BLAT results file generated using "-o = pslx". These query sequences were then aligned to the reference genome using MAFFT v47.19 (Katoh and Standley 2013) with the options "-auto" and "-adjustdirection". The smaller fragments whose sequences did not align to the exons were manually removed, and exons from the same individual of each species were merged.

For most of these genes, some edges of exons and several short exons were still missing from the assemblies. To recover the sequences or fill these gaps, we implemented an in silico target capture using MITObim (Hahn et al. 2013). For this procedure, we used the recovered exon sequences as bait sequences and the raw exome capture as target data. We included the following parameters "-quick" (starts process with initial baiting using provided fasta reference), "-mismatch" (% number of allowed mismatches in mapping), and "-kbait" (set kmer for baiting stringency). For the last two parameters we used low stringency (12-17 and 10-15, respectively) to bait more raw reads beyond those obtained from standard assemblers. This approach allowed MITObim to progressively expand the extremes of the bait sequences beyond the original gene reconstructions. The resulting sequences were then aligned with the original sequence matrices for each gene and full sequences from NCBI references (e.g., Xenopus, Nanorana, and Rana). In most cases, these extended sequences overlapped and were merged with the original ones to recover the missing regions. Lastly, we generated a clean alignment containing only coding regions that we used for the subsequent analyses. Raw sequencing coverage for SWS2 past position 389 was low (perhaps because probes designed for that region did not effectively pull-down exons compared to other probes), so we excluded the data past position 389 to avoid any potential errors in its reconstruction.

#### Gene Tree Construction

To check for any sequence contaminants, misidentifications, or assembly errors, we estimated a gene tree for each opsin, partitioned by codon position, using raxml-ng v.0.9.0 (Kozlov et al. 2019) implemented with the GTR + G model and 200 bootstrap replicates to assess support. We found several sequences of different species to be identical for each gene, although in each case the identical sequences were from closely related species. For LWS, there were two sets of identical sequences: Atelopus glyphus and A. spurrelli; Brachycephalus boticario and B. olivaceus. For SWS1 and SWS2, there was one set of identical sequences for each gene: Atelopus glyphus, A. limosus, and A. varius for SWS1; Brachycephalus pernix and B. pombali for SWS2. For RH1, we found two sets of identical species: B. pernix and B. pombali; Phyllobates aurotaenia and P. terribilis. Because there are no genomes for these species, it was not possible to verify the sequences, so we conservatively excluded all identical sequence sets from the following analyses. Finally, to check for chimeras, we aligned all four gene sequence sets to each other using MAFFT (Katoh and Standley 2013) and estimated a gene tree using raxml-ng v.0.9.0 (Kozlov et al. 2019), the GTR + G model, and 1000 bootstrap replicates to assess support (with bootstrapping enabled).

#### Determination of Diel Habits

Data on time of activity were compiled from primary and secondary literature, including species descriptions, taxonomic revisions, and books (see supplementary table S1, Supplementary Material online for details and Supplementary Information for references). Activity patterns were categorized as diurnal or nocturnal, with the latter category conservatively including species with crepuscular or mixed activity patterns. When not clearly stated, the time of activity was inferred from reports on time of calling, breeding activity, and behavior (asleep or active) at the time of collection. Diurnal groups include all of Dendrobatidae, Atelopus, and Brachycephalus (and the stem branches leading to each of these clades), as well as Mantella baroni, Mantidactylus betsileanus, and Melanophryniscus stelzneri. Species of the following genera were classified as nocturnal: Adenomera, Agalychnis, Amazophrynella, Amietia, Bufotes, Centrolene, Ceratophrys, Cochranella, Craugastor, Discoglossus, Fejervarya, Hyla, Hymenochirus, Ischnocnema, Leptobrachium, Limnodynastes, Lithodytes, Microhyla, Nanorana, Odorrana, Oreolalax, Osornophryne, Pelobates, Pyxicephalus, Quasipaa, Rana, Rhinella, Scaphiopus, Telmatobius, and Xenopus.

#### Analyses of Selection

An updated phylogeny of the focal taxa was derived from alignments provided in the two largest phylogenetic reconstructions of amphibians (Pyron 2014; Jetz and Pyron 2018). Both alignments were appended, taxa not included in our analysis were removed, and duplicate taxa were removed by choosing the one with more sequence data. Taxonomic nomenclature was updated following AmphibiaWeb (2022). The final alignment was realigned with MAFFT (Katoh and Standley 2013), then reviewed and adjusted manually, particularly the mitochondrial ribosomal gene sequences. Multiple alignment programs provide a good starting point, but they usually need to be examined and adjusted by eye (Baum and Smith 2013). The optimized alignment was then used for phylogenetic estimation. A maximum likelihood tree was estimated using IQ-TREE2 v2.1.3 (Minh et al. 2020) with five replicate runs. The tree was constrained so that the topology among the families of Hyloidea matched that found by Feng et al. (2017) and Hime et al. (2021). This method was preferred because the trees presented in Pyron (2014) and Jetz and Pyron (2018) are largely dictated by the abundance of data from mitochondrial genes. In contrast, the trees found by Feng et al. (2017) and Hime et al. (2021) are based on 100-1000 s of nuclear genes. The sequences were partitioned by gene and codon and the best-partitioned model was determined using -TESTMERGE option (Chernomor et al. 2016; Kalyaanamoorthy et al. 2017) within IQTREE2. Ultrafast bootstrap values (Hoang et al. 2018) were calculated using 50,000 replicates and were plotted on the best likelihood tree. The data matrix, IQTREE2 scripts, constraint tree, and analysis log files are included in supplementary Data S3, Supplementary Material online in the folder IQTree.

For each selection analysis, we pruned the tree using the "ape" R package (Paradis and Schliep 2019) to contain the subset of species available for each opsin gene. In some cases, we replaced a tip in the tree with a closely related species that was used in our study (see supplementary table S1, Supplementary Material online for details). We then used the resulting tree as the backbone for three types of site-based selection analyses (with P-value set to 0.05) in HyPhy v2.5.14 (Kosakovsky Pond et al. 2005). The type of selection is determined by the value of omega  $(\omega)$ , which is calculated by rate of nonsynonymous substitutions (dN) divided by rate of synonymous substitutions (dS).  $\omega$  values significantly less than 1 indicate negative selection, while neutral selection is indicated by  $\omega$  values not significantly different from 1, and positive selection is detected by  $\omega$  values significantly greater than 1. To conduct hypothesis testing, we first used Contrast-FEL, which statistically compares dN substitution rates for each site between two sets of branches (foreground and background) to determine whether different selection regimes occur in each set of branches (Kosakovsky Pond et al. 2021). Because there were no SWS2 sequences for Dendrobatidae, we did not include this gene in Contrast-FEL analyses. We created five sets of foreground lineages to compare with background lineages in Contrast-FEL: "DENDRO", that is all Dendrobatidae plus its stem branch, "DIURNAL", that is all diurnal branches as described above, and "DIURN-DEN", which is all diurnal branches except for the Dendrobatidae lineages and its stem branch. We also compared substitution rates between transition branches (the stem branches of Dendrobatidae, of Atelopus, and of Brachycephalus) and all other branches, but no sites were identified to be under different selection regimes in this foreground (see supplementary Data S1, Supplementary Material online). Once sites under different selection regimes were identified with Contrast-FEL, we used FEL (see below) with the same sets of foreground and background lineages to determine whether  $\omega$  values at sites of interest were significantly different from 1. For these analyses, we designated foreground groupings according to the list above (DENDRO, DIURNAL, and DIURN-DEN). We repeated FEL analyses using the inverse of each grouping (e.g., all lineages except DENDRO) to estimate background  $\omega$  values (reported in table 1).

We then conducted a series of complementary approaches to detect site-specific and branch-site combinations of selection patterns across the entire phylogeny using methods in HyPhy and PAML. Fixed effects likelihood (FEL) estimates the rate of synonymous (dS) and nonsynonymous (dN) substitutions per site with maximum likelihood and compares them using likelihood ratio tests to determine if  $\omega$  is greater or less than 1 (Kosakovsky Pond and Frost 2005); this test provides an estimate of  $\omega$  for the entire phylogeny (or only for foreground branches when specified) for each site, which can be used to show pervasive positive or negative selection at specific sites. Fast, Unconstrained Bayesian AppRoximation (FUBAR) is similar to FEL in that it compares single estimates of dN and dS values for each site but uses a hierarchical Bayesian method rather than maximum likelihood (Murrell et al. 2013); this test is more sensitive than FEL to weaker signatures of positive selection (i.e., when  $\omega$  is close to 1). Mixed effects model of evolution (MEME) is a branch-site model that uses maximum likelihood to test for positive selection at each site (Murrell et al. 2012); this method does not assume a single  $\omega$  value across the tree and thus can be used to detect selection at a specific site in a subset of branches on a tree. To compare our results to those of Schott et al. (2022), we also conducted CODEML analyses comparing the M7, M8, and M8a models in the PAML software suite (Yang 1997, 2007). The fit of M7 and M8 models for each gene were compared using a likelihood ratio test with two degrees of freedom; the fit of M8 and M8a models were compared using a likelihood ratio test with one degree of freedom.

Following convention, the amino acid sites are numbered based on the bovine rhodopsin sequence (NP 001014890.1); sequences from each opsin gene were aligned with bovine rhodopsin using MAFFT v7.419, and the numbering of the amino acid site was determined by referring to the numbering of bovine rhodopsin starting with the start codon as 1 (see supplementary Data S1, Supplementary Material online). Table 1 indicates how bovine RH1 positions compare to positions in our alignments. We report our selection results referring to the location of each amino acid according to the 3D structure of opsins, which encompasses seven transmembrane domains (TMD I-VII), three extracellular domains (ECD I-III), and the amino- and carboxyl- termini (N and C) (Palczewski et al. 2000; tables 1 and 2, fig. 1 and supplementary S1, Supplementary Material online).

Attempts to find SWS2 in O. pumilio and R. imitator. As we did not recover SWS2 sequences from any dendrobatid species in our bait-capture data set, we attempted to verify whether the SWS2 gene had been lost in this clade using transcriptomics and genome skimming.

First, we attempted to determine which opsin genes are expressed in eye tissue of O. pumilio. As part of another project, one eye from each of 11 O. pumilio populations **Bastimentos** (Aguacate, Almirante, "Cemetery", Bastimentos "Green", Bastimentos "Orange", Cavo Agua, Colón, Pastores, Popa, San Cristóbal, Solarte; see Maan and Cummings (2012) for details) was taken out of RNAlater and placed immediately in Trizol (Life Technologies, Grand Island, NY). RNA was extracted according to manufacturer instructions. Equal concentrations of total RNA from each of the 11 samples were pooled into one sample from which poly-adenylated RNA was isolated using the Poly(A) purist kit (Life Technologies) and manufacturer instructions. Lack of contaminating rRNA was confirmed using an Agilent 2100 Bioanalyzer. Strand-specific libraries for 100-bp paired-end sequencing were prepared and sequenced on the Illumina HiSeq 2000 according to manufacturer library kit instructions. A total of 83,168,029 reads were obtained. We preprocessed the reads using Trimmomatic (Bolger et al. 2014) by removing adapter sequence and sequence artifacts as well as trimming low-quality nucleotides based on the Phred score (Ewing et al. 1998) greater than 20, which corresponds to a 1% sequencing error rate.

De novo sequence assembly was then completed using Trinity (Grabherr et al. 2011) on the Odyssey cluster supported by the FAS Science Division Research Computing Group at Harvard University. We remapped reads to the raw assembly using BWA (Li and Durbin 2009) and then used eXpress (http://bio.math.berkeley.edu/eXpress/) to generate FPKM (fragments per kilobase per million mapped) scores for each contig. Low-confidence contigs that had an FPKM value of less than one were removed from the draft assembly. After removing contigs with low confidence, we used CD-HIT-EST (Li and Godzik 2006) to remove contig redundancy. Given that redundant contigs can represent alternative splice variants, polymorphisms among the pooled individuals, or sequencing errors, we used a conservative threshold of 98% sequence similarity. The final assembly was annotated using Trinotate (Bryant et al. 2017); it was found to contain only three opsin genes: RH1, LWS, and SWS1. The SWS2 gene was absent. Thus, we hypothesized that SWS2 might have been lost in the ancestor of dendrobatids.

Genomics-based approaches included a synteny analysis with LWS and mining of publicly available dendrobatid genomes. For the first approach, we predicted that remnants of SWS2 might be detectable in genomic data as a pseudogene either near its expected location or translocated elsewhere in the genome. We thus reviewed the genomes of *O. pumilio* (Rogers et al. 2018; Rodríguez et al. 2020) and *R. imitator* (Stuckert et al. 2021) for evidence of SWS2 using synteny and genome mining. We searched for the syntenic block containing LWS using Genomicus v.100.01 (Nguyen et al. 2018) with search term *opn1lw* and *Xenopus tropicalis* as the focal species.

To visualize gene order in the syntenic block, we used BLAST to locate SWS2, LWS, and the gene predicted to be directly upstream of LWS, MECP2 (methyl-CpG binding transcription factor) in N. parkeri, and X. tropicalis genomes. We hypothesized that a degraded ancestral SWS2 sequence might be maintained in O. pumilio and R. imitator upstream of LWS. We mined the published genome assembly of O. pumilio (Rogers et al. 2018) and a rescaffolded version of this genome (Rodríguez et al. 2020), as well as the recently published R. imitator genome for LWS using BLAST v2.10.0 (Altschul et al. 1990). We then attempted to align the scaffolds containing LWS against SWS2 sequences from other amphibians using MAFFT v7.453 (Katoh and Standley 2013). We note that the O. pumilio scaffold contains two sections of Ns upstream of LWS (2093 and 2391 nucleotides long), which may inflate our perceived length of coverage of this chromosome. Likewise, it is possible that this scaffold might be misassembled and require further refinement and deeper coverage. The rescaffolded version (Rodríguez et al. 2020) of the O. pumilio scaffold containing LWS is identical to the original and thus did not alter our conclusions. As a positive control for this analysis, we also ran our pipeline on the draft genome of R. marina, which is known to contain SWS2 based on our data. This analysis is implemented in the Dryad folder named "synteny" (see supplementary Data S4, Supplementary Material online).

Lastly, we also mined two dendrobatid genomes for SWS2. Because we were unable to detect SWS2 upstream of the O. pumilio and R. imitator LWS scaffolds, we then hypothesized that the syntenic relationship between LWS and SWS2 might have been broken in O. pumilio and R. imitator. Therefore, we used tblastn v2.10.0 to screen all scaffolds from the rescaffolded genome assembly of O. pumilio and the draft assembly of R. imitator for any potential SWS2 orthologs. This approach uncovered several candidate sequences with low E-values (<1e-10). We screened the top candidates using tblastn against Nanorana parkeri, a well-annotated frog genome more closely related to dendrobatids than Xenopus. None of the candidate sequences returned SWS2 as the most likely ortholog; the highest scoring sequences were annotated as related proteins (e.g., pinopsin, rhodopsin; supplementary table S5, Supplementary Material online). As a positive control for this analysis, we also ran tblastn on 43 known SWS2 sequences from other frog species against Nanorana parkeri. As SWS2 is known to be present in Nanorana, we would expect our pipeline to correctly detect and annotate these sequences. We found that the top hit from N. parkeri correctly identified SWS2 as the most likely ortholog for all 43 known (see supplementary table SWS2 sequences S5, Supplementary Material online). As another positive control, we ran an identical search for LWS, which identified LWS as the top hit for 101 other frog sequences and for one of the O. pumilio scaffolds (see supplementary table S5, Supplementary Material online). Lastly, we ran the reciprocal best-hit pipeline for SWS2 using the R. marina

genome as the query. Since *R. marina* is known to contain *SWS2*, we would expect that our pipeline should identify *SWS2* in the *R. marina* genome. We found that the pipeline identifies the same contig in *R. marina* that is identified in our synteny-based analysis (ONZH01019223.1) as a likely *SWS2* ortholog, while other sequences with low E-values were annotated with other related opsin genes (see supplementary table S5, Supplementary Material online). This analysis is implemented in the Dryad folders named "SWS2\_search" and "LWS\_search" (see supplementary Data S4, Supplementary Material online).

#### **Supplementary Material**

Supplementary data are available at *Molecular Biology and Evolution* online.

#### Acknowledgments

This research was funded by NSF DEB-1556967 to D.C.C.; NSF GROW, NSF DDIG (DEB-1404409), National Geographic Society 2014 Young Explorer Grant, and UC Berkeley start-up to RDT; NIH 5P50GM068763, NSF IOS-1822025, and William F. Milton Fund from Harvard Medical School to L.A.O.; L.A.O. is a New York Stem Cell Foundation—Robertson Investigator, J.C.S. was supported by SJU start-up funds and NSF DEB-2016372. M.J.N. was supported by the American Association of University Women. We thank Roberto Ibañez and Brian Gratwicke (Amphibian Rescue and Conservation Project), Vicky Poole, and Kevin Barrett (The Maryland Zoo), Luis A. Coloma (Fundación Otonga and Centro Jambatu de Investigación y Conservación de Anfibios; Quito, Ecuador), César Aguilar (Universidad Nacional Mayor de San Marcos; Lima, Peru), and Travis LaDuc (Biodiversity Center, UT Austin) for facilitating access to tissues and specimens. We thank two anonymous reviewers for critiques that improved the quality of this article. We also acknowledge and express gratitude for the lives of the animals involved in this study. The publication of this article was made possible in part by support from the Berkeley Research Impact Initiative (BRII) sponsored by the UC Berkeley Library.

### **Author Contributions**

R.D.T. and J.C.S. conceptualized the study; Y.C.W., R.D.T., J.C.S., L.H.U., L.A.O., M.J.N., A.R., and M.E.C. collected data; Y.C.W., J.C.S., L.H.U., and R.D.T. performed analyses; D.C.C., L.A.O., R.D.T., M.E.C., and J.C.S. acquired funding; R.D.T., J.C.S., M.B.C., and M.E.M. conducted fieldwork; M.R.P., M.E.C., M.E.M., C.L.R.-Z., S.R.R., M.B.C., R.D.T., and J.C.S. contributed samples and helped with permits; Y.C.W., M.J.N., L.H.U., J.C.S., L.H., and R.D.T. prepared the original draft; R.D.T. coordinated the project; R.D.T. and J.C.S. supervised the study; all authors reviewed and revised the manuscript.

# **Data Availability**

Raw Illumina sequencing data are deposited in the Sequence Read Archive (BioProject PRJNA856227). Specimens are available in QCAZ at Pontificia Universidad Católica del Ecuador (Quito, Ecuador), Centro Jambatu de Investigación y Conservación de Anfibios (Quito, Ecuador), Museo de Historia Natural de la Universidad Nacional Mayor de San Marcos (MUSM, Lima, Peru), Coleção de Herpetologia, Departamento de Zoologia, Universidade Federal do Paraná, (Paraná, Brazil), or Museo de Historia Natural ANDES at the Universidad de los Andes. All scripts and other data are available in the supporting material on DataDryad (DOI: 10.5061/dryad.zw3r2289j).

**Conflict of interest:** This publication is based in part on work by D.C.C. while serving at the National Science Foundation. Any opinion, findings, and conclusions or recommendations expressed in this material are those of the author(s) and do not necessarily reflect the views of the National Science Foundation or United States government.

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