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Authors: Melander, Scott Lucas, and Mueller, Rachel Lockridge

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Comprehensive Analysis of Salamander Hybridization Suggests a Consistent Relationship between Genetic Distance and Reproductive Isolation across Tetrapods

Scott Lucas Melander¹ and Rachel Lockridge Mueller¹

Hybridization between populations along the path to complete reproductive isolation can provide snapshots of speciation in action. Here, we present a comprehensive list of salamander hybrids and estimate genetic distances between the parental hybridizing species using one mitochondrial and one nuclear gene (MT-CYB and RAG1). Salamanders are outliers among tetrapod vertebrates in having low metabolic rates and highly variable sex chromosomes. Both of these features might be expected to impact speciation; mismatches between the mitochondrial and nuclear genomes that encode the proteins for oxidative metabolism, as well as mismatches in heteromorphic sex chromosomes, can lead to reproductive isolation. We compared the genetic distances between hybridizing parental species across four main tetrapod groups that differ in metabolic rates and sex chromosome diversity: salamanders, lizards, mammals, and birds. Our results reveal no significant differences, suggesting that variation in these traits across vertebrates does not translate into predictable patterns of genetic divergence and incompatible loci in hybrids.

PECIES formation is typically a temporally extended process, occurring over countless generations as one population-level lineage diverges into two independent population-level lineages (de Queiroz, 1998). Because of the long timescales, studying this process is challenging; however, diverging lineages that retain the ability to reproduce can provide snapshots of speciation in action (Harrison and Larson, 2016; Soltani et al., 2017). Classic examples include ring species such as Ensatina eschscholtzii, where different steps of speciation can be examined in a single taxon by looking at how the populations distributed around the geographic "ring" interbreed or hybridize (Pereira and Wake, 2009; Devitt et al., 2011). Specifically, hybrids provide an opportunity to identify the key genetic factors that become reproductive barriers when divergent genomes mix (Arnold, 1997; Toro et al., 2002; Harrison and Larson, 2016; Qvarnström et al., 2016). Typically, populations that come into contact and interbreed after isolation will have alleles interact in new, untested ways (Barton and Hewitt, 1985). If the admixture of alleles is intrinsically harmful, such as the hybrid being sterile, these negative interactions are called Dobzhansky-Muller or Bateson-Dobzhansky-Muller incompatibilities (BDMIs) and are a source of reproductive isolation (Bateson, 1909; Dobzhansky, 1937; Muller, 1942; Orr, 1996).

One special case of BDMIs is when the mitochondria and the nucleus are mismatched after hybridization—i.e., mitonuclear discordance (Ellison and Burton, 2008; Sloan et al., 2017; Hill et al., 2019). When the nuclear genome includes contributions from two species, but the mitochondrial genome comes from only one of the two, the gene products encoded by the two organelles cannot always functionally interact, causing reduced fitness and contributing to reproductive isolation. More specifically, this reduced fitness can be reflected in the metabolic costs of being a hybrid, which include increased respiration rates, increased levels of reactive oxygen species (ROS), and increased metabolic rate (Olson et al., 2010; Gvoždík, 2012; Barreto and Burton, 2013; Borowiec

et al., 2016; McFarlane et al., 2016; Prokić et al., 2018). We hypothesize that if an organism has a low metabolic rate to begin with, this change in OXPHOS functionality might have a smaller effect on overall hybrid fitness. This, in turn, could allow hybridization between species with more severely mismatched mitochondrial and nuclear genomes. Tetrapods are a good model system in which to test this hypothesis because of the wide range of metabolic rates that exists in the clade (Pough, 1980; White el al., 2006; Anderson and Gillooly, 2018). Literature reviews summarizing hybrids in birds, mammals, and lizards have been published previously (Fitzpatrick, 2004; Jančúchová-Lásková et al., 2015); birds have the highest metabolic rates, followed by mammals, and then lizards. Salamanders (order Caudata) are an important clade to incorporate into a comparative analysis of tetrapods because they have the lowest metabolic rates (Pough, 1980; Glatten et al., 1992), and there are many reported cases of salamander hybrids. With their low metabolic requirements, we predict that salamander hybrids can tolerate greater levels of genetic divergence—a proxy for mitonuclear mismatch between parental species before complete reproductive isolation occurs.

Sex chromosomes are also important during speciation and the emergence of reproductive isolation or hybrid incompatibility (Lima, 2014) because of Haldane's rule, where the heterogametic sex is more likely to be infertile by a variety of potential mechanisms, or the related large X/Z effect, where a disproportionate share of hybrid incompatibilities is found on either the X or Z chromosome (Presgraves, 2008; Lavretsky et al., 2015; Janoušek et al., 2019). In hybrid zones, mutations in sex chromosomes have been shown to limit introgression (Cortés-Ortiz et al., 2019) or even cause complete reproductive isolation between lineages (Johnson and Lachance, 2012; Hooper et al., 2019). Compared to other tetrapod clades, salamanders have variable genetic sexdetermining mechanisms with either homomorphic sex chromosomes or heteromorphic sex chromosomes with either ZW or XY systems (Eggert, 2004). ZW and XY systems

Department of Biology, Colorado State University, 251 W Pitkin Street, Fort Collins, Colorado 80523; Email: (SLM) Scott.Melander@colostate. edu; and (RLM) Rachel.Mueller@colostate.edu. Send reprint requests to SLM.
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Table 1. Hybridization in salamanders.

	Parental	species		
Species 1	Sex chromosome type	Species 2	Sex chromosome type	Resources
Ambystomatidae				
Ambystoma dumerilii		Ambystoma mexicanum		Brandon, 1977
Ambystoma dumerilii		Ambystoma rivulare		Brandon, 1977
Ambystoma macrodactylum		Ambystoma macrodactylum		Lee-Yaw et al., 2014
North Central		Rocky Mountains		,
Ambystoma macrodactylum	ZW*	Ambystoma mexicanum	ZW ^{1.2.3}	Brandon, 1977
Ambystoma maculatum	_,,	Ambystoma maculatum	2**	Johnson et al., 2015
Eastern		Western		301113011 Ct di., 2013
Ambystoma mexicanum	ZW ^{1.2.3}	Ambystoma tigrinum	ZW ^{1.2.3}	Woodcock et al., 2017
Ambystoma mexicanum	ZW ^{1.2.3}	Ambystoma againam Ambystoma opacum	ZW*	Brandon, 1977
Ambystoma talpoideum	Z V V	Ambystoma texanum	Z V V	Brandon, 1977 Brandon, 1977
	ZW*		ZW ^{1.2.3}	Brandon, 1977
Ambystoma texanum	ZW ^{1.2.3}	Ambystoma tigrinum		
Ambystoma tigrinum californiense	ZVV	Ambystoma mavortium	ZW*	Fitzpatrick et al., 2009 Fitzpatrick and Shaffer, 2004 Riley et al., 2003
Cryptobranchidae		Andrica innonious		Full was at al. 2015
Andrias davidianus		Andrias japonicus		Fukumoto et al., 2015
Dicamptodontidae	NA/PC	D:	NA/PC	C 1000
Dicamptodon ensatus Hynobiidae	XX^{PC}	Dicamptodon tenebrosus	XX^{PC}	Good, 1989
Hynobius nebulosus		Hynobius nigrescens		Kawamura, 1953
Onychodactylus japonicus		Onychodactylus japonicus		Yoshikawa et al., 2012
S-Tohoku ,		SW-Honshu		
Salamandrella keyserlingii Plethodontidae	XX^{PC}	Salamandrella tridactyla	XX^{PC}	Malyarchuk et al., 2015
Aneides klamathensis		Aneides flavipunctatus		Reilly and Wake, 2019
Batrachoseps gavilanensis		Batrachoseps luciae		Jockusch and Wake, 2002
Bolitoglossa franklini	XX^{PC}	Bolitoglossa lincolni	XX^{PC}	Wake et al., 1980
Desmognathus carolinensis	701	Desmognathus orestes	701	Mead and Tilley, 2000
Desmognathus conanti	XX*	Desmognathus fuscus	XX^{PC}	Bonett, 2002
Desmognathus fuscus	XX ^{PC}	Desmognathus ochrophaeus	XX ^{PC}	Sharbel et al., 1995
Desmognathus fuscus	XX ^{PC}	Desmognathus santeetlah	XX*	Tilley, 1988
Ensatina eschscholtzii croceater	XX ^{PC}	Ensatina eschscholtzii platensis	XX ^{PC}	Pereira and Wake, 2009
Ensatina eschscholtzii eschscholtzii	XX ^{PC}	Ensatina eschscholtzii klauberi	XX ^{PC}	Devitt et al., 2011
	XX ^{PC}	Ensatina eschscholtzii picta	XX ^{PC}	
Ensatina eschscholtzii oregonensis	XX ^{PC}		XXPC	Pereira and Wake, 2009
Ensatina eschscholtzii oregonensis	XX ^{PC}	Ensatina eschscholtzii xanthoptica	XX ^{PC}	Pereira and Wake, 2009
Ensatina eschscholtzii platensis		Ensatina eschscholtzii xanthoptica		Alexandrino et al., 2005 Sweet, 1984
Eurycea bislineata	XXPC	Eurycea cirrigera	XX^{PC}	Guttman and Karlin, 1986
Eurycea cirrigera	XX^{PC}	Eurycea wilderae	XX*	Kozak, 2003
Eurycea neotenes		Eurycea tridentifera		Kozak and Montanucci, 200
Hydromantes ambrosii	XY ^{1.2.3}	Hydromantes italicus	XY ^{1.2.3}	Lunghi et al., 2018 Ficetola et al., 2019
Plethodon aureolus	XX*	Plethodon shermani	XX^{PC}	Highton and Peabody, 2000
Plethodon chattahoochee		Plethodon chlorobryonis		Highton and Peabody, 2000
Plethodon chattahoochee	XX*	Plethodon shermani	XX^{PC}	Highton and Peabody, 2000
Plethodon chattahoochee		Plethodon teyahalee	- -	Highton and Peabody, 2000
Plethodon cheoah		Plethodon teyahalee		Highton and Peabody, 2000
Plethodon chlorobryonis		Plethodon cylindraceus		Highton and Peabody, 2000
Plethodon chlorobryonis	XX*	Plethodon metcalfi	XX^{PC}	Highton and Peabody, 2000
Plethodon chlorobryonis	/V\	Plethodon teyahalee	/V\	Highton and Peabody, 2000
Plethodon cinereus	XX^{PC}	Plethodon electromorphus	XX*	Lehtinen et al., 2016
Plethodon cylindraceus	XX*		XX ^{PC}	
	XX ^{PC}	Plethodon glutinosus		Highton and Peabody, 2000
Plethodon dorsalis	AX ·	Plethodon ventralis	XX*	Highton, 1997
Dothodon ologtrons	VV*	Dlathadan richmandi	XX^{PC}	Duncan and Highton, 1979
Plethodon electromorphus	XX*	Plethodon richmondi		Highton, 1999
Plethodon fourchensis	XX*	Plethodon ouachitae	XX^{PC}	Shepard et al., 2011

Table 1. Continued.

	Parent	al species		
Species 1	Sex chromosome type	Species 2	Sex chromosome type	Resources
Plethodon glutinosus	XXPC	Plethodon jordani	XX ^{PC}	Hairston et al., 1992
Plethodon glutinosus	XX ^{PC}	Plethodon kentucki	XX*	Kuchta et al., 2016
Plethodon hoffmani		Plethodon virginia		Hairston et al., 1992 Highton, 2009 Dawley, 1987
Plethodon jordani	XX^{PC}	Plethodon metcalfi	XX^{PC}	Chatfield et al., 2010
Plethodon jordani	XX ^{PC}	Plethodon teyahalee	XX*	Chatfield et al., 2010
Plethodon metcalfi	XX ^{PC}	Plethodon teyahalee	XX*	Chatfield et al., 2010
Plethodon shermani	XX ^{PC}	Plethodon teyahalee	XX*	Highton and Peabody, 2000
Proteidae				
Necturus maculosus Salamandridae	XY ^{1.2.3}	Necturus aff. lewisi	XY ^{1.2.3}	Nelson et al., 2017
Chioglossa lusitanica Northern		Chioglossa lusitanica Southern		Sequeira et al., 2005
Cynops pyrrhogaster		Cynops pyrrhogaster		Tominaga et al., 2018
Central	10/23	Western	va/2.3	
Lissotriton helveticus	XY ^{2.3}	Lissotriton vulgaris	χγ ^{2.3} χγ ^{2.3}	Johanet et al., 2011
Lissotriton montandoni	XY*	Lissotriton vulgaris	XY	Babik et al., 2005
Liccotriton vulgaric kocciviai	XY ^{2.3}	Lissotriton vulgaris vulgaris	XY ^{2.3}	Zieliński et al., 2013
Lissotriton vulgaris kosswigi	ΑY		XΥ	Nadachowska and Babik, 2009
Lyciasalamandra antalyana Notophthalmus viridescens		Lyciasalamandra billae Notophthalmus viridescens dorsalis		Johannesen et al., 2006 Takahashi et al., 2011
Ommatotriton ophryticus		Ommatotriton nesterovi		van Riemsdijk et al., 2018
Ommatotriton ophryticus		Ommatotriton vittatus		Yoshikawa et al., 2010
Ommatounton opinyticus		Ommatounon villatas		van Riemsdijk et al., 2018
Pleurodeles nebulosus	ZW*	Pleurodeles poireti	$ZW^{2.3}$	Escoriza et al., 2016
Salamandra salamandra gallaica	XX ^{PC}	Salamandra salamandra bernardezi	XXPC	García-París et al., 2003
Salamandra fastuosa	XX ^{PC}	Salamandra terrestris	XX ^{PC}	Ventura et al., 2015
Salamanara rastassa	701	Salarriara terrestris	701	Canestrelli et al., 2014
				García-París et al., 2003
Salamandrina perspicillata	XX*	Salamandrina terdigitata	XX^{PC}	Hauswaldt et al., 2011
, ,		3		Mattoccia et al., 2011
				Arntzen et al., 2009
Taricha rivularis	XX*	Taricha sierrae	XX^{PC}	Twitty, 1963
Taricha rivularis	XX*	Taricha torosa	XX^{PC}	Twitty, 1963
Taricha torosa	XX^{PC}	Taricha sierrae	XX^{PC}	Kuchta, 2007
Triturus carnifex	$XY^{2.3}$	Triturus cristatus	XY ^{1.2.3}	Arntzen et al., 2014
Triturus carnifex	XY ^{2.3}	Triturus dobrogicus	XY*	Arntzen et al., 2014
Triturus carnifex	XY ^{2.3}	Triturus ivanbureschi	XY*	Arntzen et al., 2014
Triturus carnifex	XY ^{2.3}	Triturus macedonicus	XY*	Arntzen et al., 2014
Triturus cristatus	XY ^{1.2.3}	Triturus dobrogicus	XY*	Arntzen et al., 2014
Triturus cristatus	XY ^{1.2.3}	Triturus ivanbureschi	XY*	Arntzen et al., 2014
Triturus cristatus	XY ^{1.2.3}	Triturus macedonicus	XY*	Arntzen et al., 2014
Triturus cristatus	XY ^{1.2.3}	Triturus marmoratus	XY ^{1.2.3}	Visser et al., 2017
Triturus ivanbureschi		Triturus macedonicus		Vučić et al., 2018

¹ Evans et al., 2012

are both found within multiple families, indicating that sex chromosomes are evolutionarily dynamic in salamanders; this makes the clade a good system for studying the effects of heteromorphic sex chromosome evolution on reproductive isolation (Charlesworth et al., 2005; Evans et al., 2012).

There are numerous studies published on hybrids in salamanders, both from long-term stable hybrid zones and conservation efforts (Fitzpatrick and Shaffer, 2004; Fukumoto et al., 2015). With a few notable exceptions (e.g., Twitty, 1963; Brandon, 1977; Gvoždík, 2012; Prokić et al., 2018),

² Hillis and Green, 1990

³ Perkins et al., 2019

^{*} Predicted sex chromosome type

PC Personal correspondence, Stan Sessions, 2019

there have not been extensive published experimental crosses in salamanders like those done in other vertebrate clades (e.g., toads, Blair, 1972; Malone and Fontenot, 2008). Here, we summarize the known cases of hybridization in salamanders. We then use this dataset to compare the genetic distances across which viable hybrids can form in different tetrapod groups and test whether differences in metabolic rate and sex chromosomes impact hybridization.

MATERIALS AND METHODS

Compilation of salamander hybrids.—The first goal was the establishment of a comprehensive list of published salamander hybrids, which did not exist when we began this research (Table 1). Taxonomy is continually revised to provide scientific names that convey accurate information about species boundaries as well as the evolutionary relationships among species. Making these species designations is a rich discipline, with disagreements among taxonomists requiring different levels of divergence or isolation between populations before they are formally named as species (Highton, 1998; Kuchta and Wake, 2016). For our literature review, we used a general lineage concept of species where a species equates to a segment of a population-level evolutionary lineage (de Queiroz, 1998). We chose to include populations far enough along their own evolutionary trajectories that when secondary contact occurred, the populations did not completely admix into a single population. By doing this, we were able to include several instances of hybridization in which the hybridizing populations have not been formally named as species. We included cases where secondary contact occurred because of human-mediated introductions, and we also included laboratory crosses. Hybridization that resulted in polyploidy was excluded to eliminate the confounding variable of increased ploidy levels. Species with evidence of historical, but not ongoing, hybridization were also excluded. This compilation of hybridizing species represents extant salamanders that are known to have had the opportunity to hybridize and done so successfully; there are undoubtedly pairs of lineages that would be able to hybridize, given the opportunity (either with or without human mediation), as well as undetected instances of natural hybridization. The criteria we chose allowed us to compile the maximum amount of information about hybrid salamanders from the literature, as well as make comparisons with existing literature surveys on birds, mammals, and lizards that applied similar criteria (Fitzpatrick, 2004; Jančúchová-Lásková et al., 2015).

Database searches were performed using the terms salamander, newt, hybrid, and contact zone. Databases used were JSTOR, Web of Science, and Wildlife and Ecology Studies Worldwide. Using the same terms, additional sources were found with the search engine Google Scholar. The searches were undertaken from January 2018 to February 2020 and included research published between 1979–2020. Many taxonomic changes occurred during this 40-year period; whenever there was a conflict in species name, the current listing on AmphibiaWeb (https://amphibiaweb.org) was used to resolve the issue.

Intensity of research on different salamander families.—To check whether or not the reported number of hybrids within each salamander family was a function of the intensity of

publication, a general review was also conducted on how well each family of salamanders is represented in the literature (Fig. 1). Using the Web of Science database, searches were performed using terms based on variations on the salamander family names with two to three search terms used per family. Terms used were: Ambystomatidae (ambystomatid and Ambystoma), Amphiumidae (amphiumid and Amphiuma), Cryptobranchidae (cryptobranchid and Cryptobranchus), Dicamptodontidae (dicamptodontid and Dicamptodon), Hynobiidae (hynobiid and Hynobius), Plethodontidae (plethodontid and Plethodon), Proteidae (proteid), Rhyacotritonidae (rhyacotritonid and Rhyacotriton), Salamandridae (salamandrid and Salamandra), and Sirenidae (sirenid). The genus names Proteus and Siren were excluded due to their use in Greek mythology; because the corresponding salamander families are small, missing papers is unlikely. The number of articles per family was standardized by dividing by the number of species in each family based on AmphibiaWeb as of July 2020 (https://amphibiaweb.org). To test whether the reported number of hybrids within each salamander family was related to the intensity of publication, a Kendall correlation coefficient was calculated comparing papers per species and hybrids per species for each family.

Genetic distances between hybridizing species of salamanders.—

The genetic distances across the parental species for each

hybridizing salamander pair were estimated using the mitochondrial gene cytochrome b (MT-CYB; Fig. 2A). MT-CYB has long been used as a phylogenetic and phylogeographic marker for salamanders, so there is wide coverage for many different species (Johns and Avise, 1998). MT-CYB has also been used as a proxy for overall genetic distance in summaries of hybridizing pairs of species in other vertebrate groups (Fitzpatrick, 2004; Jančúchová-Lásková et al., 2015). For each parental species, the longest high-quality sequence was downloaded from NCBI GenBank (https://www.ncbi. nlm.nih.gov/genbank/). In some cases, this involved extracting the MT-CYB sequence from a complete mitochondrial genome sequence. When multiple equally long, high-quality sequences were available, one was selected at random. Of the 76 salamander species pairs known to hybridize, MT-CYB sequence data were available for 62 (Table 2). For each hybridizing pair, a pairwise alignment was calculated with default ClustalW settings implemented in MEGA X (Kumar et al., 2018). Each alignment was then trimmed to the first and last overlapping nucleotide position. After trimming, the alignments ranged from 345 to 1,141 base pairs. The genetic distances were estimated with PAUP * Version 4.0a (Swofford, 2002) using the HKY85 + Γ nucleotide substitution model

In order to test how representative MT-CYB genetic distances are in describing the overall genetic divergence between species, the substitution rate of mitochondrial genes versus nuclear genes was compared (Fig. 2B). This was accomplished by dividing the genetic distance of the mitochondrial gene MT-CYB by the genetic distance of the nuclear gene recombination activating 1 (RAG1) for the parental species that hybridize (Table 2). RAG1 was selected since the gene has widespread use in phylogenetic studies

(Hasegawa et al., 1985). In order to maximize the amount of

sequence data used, the Γ distribution shape parameter α was

estimated using the MEGA X maximum likelihood model

selector based on a ClustalW alignment of all 81 salamander

MT-CYB sequences used in this study (Kumar et al., 2018).

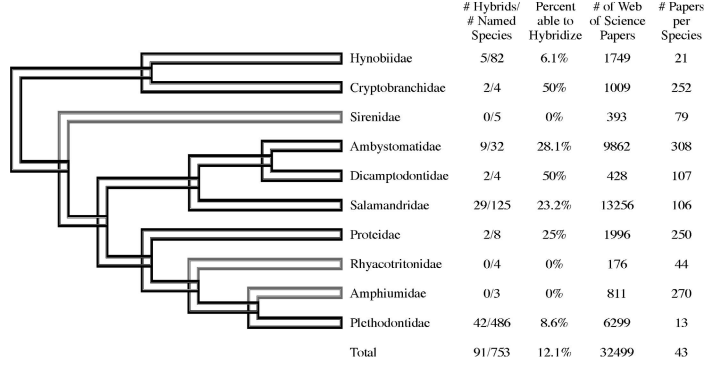
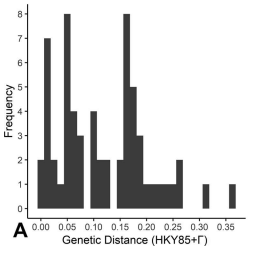


Fig. 1. Salamander hybrids are found in most families (Pyron and Wiens, 2011). About 12.1% of salamanders are known to hybridize with over half belonging to Plethodontidae. There was no significant correlation between the number of papers per species and the proportion of salamanders found to hybridize (Kendall's rank correlation P = 0.236).

and was available for the most taxa (Chiari et al., 2009). For every parental species available, the longest high-quality RAG1 sequence was downloaded from GenBank. When multiple equally long, high-quality sequences were available, one was selected at random. RAG1 genetic distances between parental species that hybridize were then estimated using the same methods as for MT-CYB genetic distances.

Genetic distances between hybridizing species of salamanders compared with hybrids in other tetrapod groups.—The MT-CYB genetic distances between hybridizing parental species of salamanders were compared to those of birds, mammals, and lizards (i.e., squamate reptiles excluding snakes). These tetrapod groups were selected because they span part of the range of vertebrate metabolic rates (Pough, 1980; White el al.,

2006; Anderson and Gillooly, 2018) and because of the existence of published summaries of known hybrids that include natural, human-introduced, and lab-crossed hybridization events (Fitzpatrick, 2004; Jančúchová-Lásková et al., 2015). A hybrid toad summary was excluded because it focused only on experimental crosses (Blair, 1972; Malone and Fontenot, 2008). In some tetrapod groups, many closely related species hybridize, or a single species hybridizes several times. This creates a potential bias when looking at many pairwise comparisons as a single taxon becomes overrepresented, and the results become dependent on a few select taxa (Fitzpatrick, 2004). Previously published work in mammals and birds resolved this issue by removing repeated taxa to remove nonindependence (Fitzpatrick, 2004), so the



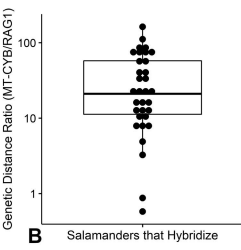


Fig. 2. (A) Histogram of cytochrome b (MT-CYB) genetic distance estimates (HKY85 + Γ) between salamanders that hybridize. Graph bin width is 0.0125. (B) Ratio of mitochondrial MT-CYB genetic distance to nuclear recombination activating 1 (RAG1) genetic distance between parental species of salamanders that hybridize on a log scale. The ratio of genetic distance ranged from 0.6 to 162 with the majority of species pairs having a higher mitochondrial genetic distance (median = 21.1).

Table 2. Hybrid pairs and accession numbers for MT-CYB and RAG1. Lists of bird hybrids and mammal hybrids modified from Fitzpatrick (2004). Lizards modified from Jančúchová-Lásková et al. (2015). Only the highest distance per genus was used in salamanders (*) for the cross-clade comparison (Fig. 3).

Bioches 1 MIT-CRB RACT A podestion MIT-CRB RACT MIT-CRB RACT MIT-CRB RACT MIT-CRB (MCRB) RACT MIT-CRB (MCRB) <		Accessi	Accession number		Accessio	Accession number	Genetic	distance	Genetic distance (HKY85 $+$ Γ)
spholes solpozeus LTG71506.1 A polasitio	Species 1	MT-CYB	RAG1	Species 2	MT-CYB	RAG1	MT-CYB	RAG1	MT-CYB/RAG1
Tight 1908 Average	Birds								
AF091728.1 CORDES AF091728.1 GLOGG 1931.1 Gross follows AF091728.1 COLOGG 1931.1 GORG 1931.1 GLOGG 1931.1 GORG 1932.4 O.005 O.005 EUSSS23.1 AF001228.1 Colog 200.8	Acrocephalus sairpaceus	LT671508.1		A. palustris	AJ004774.1		0.121		
KR93611.1 CQ505193.1 Melopatements in Melopatements and Melopat	Aegypius monachus	AY987266.1	EF078711.1	Gyps fulvus	AY987261.1	EU496474.1	0.120	0.005	22.0
EUSBSEG211 AF2946872 Codilis goldus LOB376.1 NMO01031188.1 0.366 0.075 EUSPSEG211 Codis canadensis EU166997.1 0.034 0.074 0.074 LU27556.1 Cous canadensis EU166997.1 0.074 0.074 0.074 AV495384.1 EU466229.1 EU7687.1.1 RAP5332.1 0.074 0.072 KV495384.1 EU16699.1 Crigory corrects and control of the control of the correct and corre	Agapornis nigrigenis	AF001328.1	GQ505193.1	Melopsittacus undulatus	DQ467903.1	DQ143354.1	0.265	0.036	7.3
HEISBESCO.11 Coning connectated LI08386.1 0.0375 VG7556.11 Coning connectated LI08386.1 0.0175 VG6541.11 EUJ4552.8.1 Grandenoss EUJ66997.1 0.0185 0.018 K6674.11 EUJ4552.8.1 Emberior catmelia VA79532.1 0.184 0.0184 K4562.2.2.1 Congo control Congo control VA7771.1 0.187 0.008 KG575.19.2.1 AY22876.1 LOST377.1 KA87535.1 0.159 0.035 KG675.19.2.1 AY22876.1 Farodiscen minor LU25737.1 CA6747.1 0.167 0.008 KG674.9.1.1 AY22876.1 Farodiscen minor LU25737.1 CA6748.1 0.140 0.008 LU27.5.5.1.1 AY22876.1 Farodiscen minor LU2573.1 LU275.2 0.140 0.008 LU27.5.5.1 AY22876.1 AY44529.1 AY44529.1 AY44529.1 0.140 0.008 LU27.5.5.1 AY44529.1 AY44529.1 AY44529.1 AY44529.1 0.140 0.008 LU27.5.2.	Alectura lathami	KF833611.1	AF294687.2	Gallus gallus	L08376.1	NM001031188.1	0.366	0.075	4.9
U.77556.11 Cinus connodensis EU166997.1 0.074 NA6741.11 EU34552.8.1 Grus connodensis EU166997.1 0.074 NA6522.2.1 Cingon connodensis VIVATATI.1 KIMB622.1 0.184 KA6522.2.1 Cingon connodensis VIVATATI.1 KIMB622.1 0.0221 KA6522.2.1 Cingon connodensis VIVATATI.1 KIMB622.1 0.0229 KA67519.2.1 AY2287.6.1 Cingops connocent melanoleura AF483327.1 EF373512.1 0.023 MH64565.9.1 Mycrocox mycroc	Branta canadensis	EU585629.1		Cairina moschata	L08385.1		0.235		
AV87511. EU345528.1 Fenis apivous X86758.1 FF078753.1 0.185 0.018 K4495384.1 Chagera criticulal AV495324.1 0.184 0.021 0.021 K456223.1 Chagera criticulal VA795321.1 0.184 0.084 K675192.1 AV22876.8.1 Craggos criticulas AK33377.1 0.163 0.008 K675192.1 AV22876.8.1 Leucosarca melanoleuca AK33377.1 0.167 0.008 MH645659.1 AV22876.8.1 Furniunculus EL53312.1.1 0.167 0.016 DOG19326.1 AV43290.1 Furniunculus EL53321.1.1 0.042 0.004 AR24055.1 AV443290.1 Furniunculus US6042.1 AV056997.1 0.041 AR20876.1 AV443290.1 H. Apymboraca AF07459.1 AV056997.1 0.014 AR20876.1 AV443290.1 H. Apymboraca AV07459.1 AV056997.1 0.084 AR20876.1.1 AV443290.1 H. Apymboraca AV377168.1 AV3443327.1 0.016 AR20876.1.1	Bugeranus carunculatus	U27556.1		Grus canadensis	EU166997.1		0.074		
KA495384.1 Emberiza attrinella AV495332.1 0.221 KA456229.1 Entheriza attrinella UZ7371.1 0.0167 0.008 KC67519.2.1 AV2866.1 Coragya attratos KX53417.1 KM876315.1 0.167 0.008 X74255.1 AV22876.1 AV2877.1 KK87371.1 0.167 0.008 MH64565.9.1 AVA225.1 AVA225.1 AVA225.1 0.167 0.008 MH64565.9.1 AVX425.1 AVX425.1 0.167 0.008 0.015 DQ67449.1.1 XM00504628.8 F. finanticulus KF13329.1 DQ466798.1 0.014 DQ67449.1.1 XM00504628.1 F. finanticulus AVX5055.1 AVX50679.1 0.046 AR2087.6.1 AVX0050462.1 AVX005049.1 AVX50679.1 0.046 0.006 AR2087.6.1 AVX005049.1 AVX005049.1 AVX005049.1 0.016 0.006 AR2087.6.1 AVX005049.1 AVX005049.1 AVX005049.1 0.006 0.017 AR2087.6.1 AVX005049.1 AVX005049.1 AVX00	Buteo buteo	X86741.1	EU345528.1	Pernis apivorus	X86758.1	EF078753.1	0.185	0.018	10.2
K1456229.1 C nigra U72771.1 C 0.164 EU166984.1 K74253.1 C nigra U72771.1 C 0.157 0.167 0.008 KC675192.1 AY228768.1 Eucrosops are arminor AF193823.1 E173512.1 0.159 0.008 MH645659.1 AY228768.1 Eucrosops are minor AF193823.1 0.159 0.0159 MH645659.1 Mycticorax nyctorax AF193823.1 EU233241.1 0.143 0.006 DQ674491.1 XM005046928.1 F inpoletorax LG8042.1 AF03652.1 0.144 0.006 AB242559.1 AW43290.1 F inpoletorax LG60452.1 0.046 0.004 AB24258.1 AW443290.1 Lifibrondes AR02952.1 AV056997.1 0.146 0.004 AB24258.1 AY42290.1 Lifibrondes AR02992.1 AV056997.1 0.014 0.006 AB20088.1.1 AY42290.1 Lifibrondes AR02992.1 AV056997.1 0.014 0.006 AR2547.1.2 AY42280.1 AY42280.1 AY4234.1 AY4324.1 <td>Carduelis chloris</td> <td>AY495384.1</td> <td></td> <td>Emberiza citrinella</td> <td>AY495392.1</td> <td></td> <td>0.221</td> <td></td> <td></td>	Carduelis chloris	AY495384.1		Emberiza citrinella	AY495392.1		0.221		
EU166984 FC078766.1 Coragpse atrentus KV554417.1 KM876515.1 0.167 0.008 X7425.1 AY228768.1 Leucosacion melanoleuca AF48322.1 EC73512.1 0.139 0.036 X7425.1 MH64569.1 Leucosacion milor UZ5737.1 EU33304.1 0.143 0.036 MH64569.1 LU23167.1 F innuncucus EU23312.1 0.143 0.006 DQ674491.1 XM005046928.1 F innuncucus K195052.1 DQ466798.1 0.140 0.006 DQ674491.1 XM005046928.1 F innuncucus K195052.1 DQ466798.1 0.047 0.004 AB208756.1 AP482290.1 H princhens infaustus RAD4352.1 AV056997.1 0.014 0.006 AB208756.1 AP482290.1 AF04829.1 AV056997.1 0.006 0.007 AB208756.1 AV443290.1 AV057028.1 AV056997.1 0.006 0.007 AB208756.1 AV443290.1 AV057028.1 AV057028.1 0.006 0.007 AV5244134.1 AV728802.1 AV067029.1 <td>Ciconia ciconia</td> <td>KJ456229.1</td> <td></td> <td>C. nigra</td> <td>U72771.1</td> <td></td> <td>0.184</td> <td></td> <td></td>	Ciconia ciconia	KJ456229.1		C. nigra	U72771.1		0.184		
KCR5192.1 AY228768.1 Leucosardia melanoleuca AF48337.1 EF373512.1 0.239 0.036 X74255.1 MH64569.1 Rodisagea minor U25373.1 0.143 0.036 MH64569.1 EU233167.1 Finnunculus EU233121.1 EU233241.1 0.143 0.006 EU333049.1 EVIDOSO696928.1 Finpoleuca RP36052.1 0.0466798.1 0.004 0.006 AB24259.1 AW443290.1 H. Pynthomota RP074591.1 AV056997.1 0.0146 0.004 AB24258.1 AV443290.1 H. Pynthomota AF074591.1 AV056997.1 0.0146 0.006 AB20475.1 AV443290.1 H. Pynthomota AF074591.1 AV056997.1 0.0146 0.006 AB2047.2.1 AV60882.1 AV604329.1 AV66992.1 0.006 0.007 AK1974.3.4 AV228802.1 AV66042.1 AV604329.1 0.006 0.007 AK1974.3.4 AV228802.1 AV66042.1 AV66042.1 AV66042.1 0.006 0.007 AK1974.3.4 AV228802.1	Cathartes aura	EU166984.1	EF078766.1	Coragyps atratus	KX534417.1	KM876315.1	0.167	0.008	21.8
X74255.1 Paradiscae minor U35537.1 0.159 MH645659.1 MH645659.1 Mydrocox nycticoxa AF193829.1 0.143 0.043 EU233049.1 EU233167.1 F innuncular EU23312.1 0.143 0.006 DQ674491.1 XM005046928.1 F innuncular RP350522.1 DQ466798.1 0.0140 DQ119526.1 AY43290.1 H pythoneus MF020923.1 AY056997.1 0.0247 DQ119526.1 AY443290.1 H pythoneus AF074591.1 AY056997.1 0.018 HQ60821.1 Lidbundus RA0745321.1 AY056997.1 0.044 0.004 HQ60821.1 AY05028.1 Rhea camericana L78808.1 AY0443227.1 0.044 LY6063.1 AY05028.1 Rhamman AY043232.1 AY056997.1 0.004 AY32947.2.1 AY05028.1 Rhamman AY043232.1 AY443237.1 0.005 AY1078.2.2.2.1 AY22880.2.1 AY0604329.1 AY443237.1 0.005 AY443.1.2.1 AY2280.2.1 AY45327.1 0.013	Columba livia	KC675192.1	AY228768.1	Leucosarcia melanoleuca	AF483327.1	EF373512.1	0.239	0.036	7.4
MH645659.1 EU233167.1 Mydrotox nycticonax AF193829.1 EU23324.1 0.143 0.006 EU23349.1 XM005046928.1 F finnunculus EU23312.1.1 EU23324.1.1 0.140 0.006 DQ67449.1.1 XM005046928.1 F finnunculus EU23312.1.1 EU23324.1.1 0.140 0.006 DQ119556.1 AY442290.1 H. Propleuca AF056997.1 0.047 0.004 DQ11956.1 AY442290.1 H. Propleuca AF056997.1 0.018 0.006 AR208756.1 AY42290.1 H. AF060092.3.1 AY056997.1 0.016 0.017 AR208756.1 AY42290.1 H. AF0600092.1 AY056997.1 0.006 0.007 AY229472.1 AY22800.2 Vanellus americana LY8808.1 0.006 0.007 AK197835.1 AY22800.2 Vanellus vanellus AY495411.1 AY329126.1 0.006 0.001 AF197835.1 AK19048.1 AY495411.1 AY329126.1 0.006 0.001 AF191042.1 AY100000.1 AY495411.1 AY495411.1 <td>Diphyllodes magnificus</td> <td>X74255.1</td> <td></td> <td>Paradisaea minor</td> <td>U25737.1</td> <td></td> <td>0.159</td> <td></td> <td></td>	Diphyllodes magnificus	X74255.1		Paradisaea minor	U25737.1		0.159		
EU2330491 EU233167.1 F. tinnunculus EU233121.1 EU233241.1 0.140 0.006 DQ674491.1 XM0050469281 F. hypothocus intaustus K935052.1 DQ466798.1 0.042 0.004 AB242556.1 AV443290.1 H. pyrrhonota AF074591.1 AV056997.1 0.0181 0.004 AB208756.1 AV443290.1 H. pyrrhonota AF074591.1 AV056997.1 0.0181 0.010 AB208756.1 AV443290.1 H. pyrrhonota AF074591.1 AV056997.1 0.0181 0.010 HQ60882.1 AV60882.1 HAP07926.2 AV443327.1 0.018 0.016 AY329472.1 AV057028.1 K. satrapa AV0432327.1 0.026 0.052 AY329472.1 AY057028.1 AV057028.1 AV37514.1 0.016 0.001 AY329472.1 AY057028.1 AY37514.1 AY37514.1 0.016 0.005 AY329472.1 AY057028.1 AY37514.1 0.016 0.007 0.017 AY329471.2 AY22802.1 AY495411.1 AY37514.1 </td <td>Egretta garzetta</td> <td>MH645659.1</td> <td></td> <td>Nycticorax nycticorax</td> <td>AF193829.1</td> <td></td> <td>0.143</td> <td></td> <td></td>	Egretta garzetta	MH645659.1		Nycticorax nycticorax	AF193829.1		0.143		
DOG/4491.1 XM005046928.1 F. hypoleuca KI930552.1 DQ466798.1 0.042 0.004 AB242559.1 AR243290.1 H. pynthonota U86042.1 AV056997.1 0.014 0.017 DQ119526.1 AY443290.1 H. pynthonota FM20923.1 AV056997.1 0.016 0.017 HQ60882.1.1 AY443290.1 R. ridibundus MH70382.1 0.0146 0.014 U76054.1 AY057028.1 R. satrapa AV04329.1 AY443327.1 0.005 0.017 AY329472.1 AY057028.1 R. satrapa AV04329.1 AY443327.1 0.006 0.017 AY329472.1 AY057028.1 R. satrapa AV04329.1 AY443327.1 0.006 0.017 AY329472.1 AY057028.1 AY22802.1 AY443327.1 0.026 0.003 AY329472.1 AY22802.1 AY443327.1 0.026 0.003 AY329472.1 AY22802.1 AY443327.1 0.026 0.003 AY32941.1.1 AY307214.1 0.106 0.013 AY9481.18.1 <td>Falco columbarius</td> <td>EU233049.1</td> <td>EU233167.1</td> <td>F. tinnunculus</td> <td>EU233121.1</td> <td>EU233241.1</td> <td>0.140</td> <td>900.0</td> <td>24.4</td>	Falco columbarius	EU233049.1	EU233167.1	F. tinnunculus	EU233121.1	EU233241.1	0.140	900.0	24.4
AB242559.1 APR242559.1 Perisoneus infaustus UB6042.1 APD242559.1 AV056997.1 0.247 DQ119526.1 AY443290.1 H. pyrthonotra AF0743291.1 AY056997.1 0.181 0.010 AB208756.1 L. ridburdus L. ridburdus MH079362.1 AY443220.1 0.146 HQ608821.1 R. rochilus Rhea americana L. 78808.1 0.014 0.014 AY329472.1 AY057028.1 R. satrapa AV043327.1 0.002 0.017 KW434134.1 AY228802.1 T. phiomelos AV495411.1 AY339126.1 0.260 0.002 AF197835.1 KC789829.1 T. phiomelos AV495411.1 AY370214.1 0.196 0.013 AF197835.1 KC789829.1 T. phiomelos AV495411.1 AY3707214.1 0.196 0.003 AF197835.1 KC789829.1 T. phiomelos AV495411.1 AY3707214.1 0.196 0.003 AF19793.1 L. principal Conolophus subcristatus AY948122.1 AY307214.1 0.196 0.001 <td< td=""><td>Ficedula albicollis</td><td>DQ674491.1</td><td>XM005046928.1</td><td>F. hypoleuca</td><td>KJ930552.1</td><td>DQ466798.1</td><td>0.042</td><td>0.004</td><td>11.0</td></td<>	Ficedula albicollis	DQ674491.1	XM005046928.1	F. hypoleuca	KJ930552.1	DQ466798.1	0.042	0.004	11.0
DQ119526.1 AV443290.1 H. pyrrhonota AF074591.1 AV056997.1 0.181 0.010 AB208756.1 L. ridbundus L. ridbundus HM209323.1 AV056997.1 0.036 AB208756.1 L. ridbundus L. ridbundus MH079362.1 0.005 0.016 U76054.1 AV057028.1 R. satrapa A004329.1 AV443327.1 0.002 0.017 AV32472.1 AV057028.1 R. satrapa A004329.1 AV329126.1 0.002 0.017 AV32472.1 AV057028.1 R. satrapa AV49571.1 AV339126.1 0.000 0.017 AV328802.1 Vonellus vanellus I. philomelos AV49571.1 AV359126.1 0.000 0.013 AV348118.1 KK7350710.1 Conolophus subcristatus AV349122.1 AV3507214.1 0.196 0.003 BU037682.1 MI112592.1 A trinitatis AV493592.1 JIN112645.1 0.137 0.007 BU037682.1 MK762187.1 I. monitoria AV34932.1 KR350724.9.1 0.137 0.007	Garrulus glandarius	AB242559.1		Perisoreus infaustus	U86042.1		0.247		
AB208756.1 L ridibundus FM209933.1 0.056 HQ608821.1 P. trochilus NH079362.1 0.046 U76054.1 Rhea americana L ridibundus 1.7808.1 0.044 HQ608821.1 AY329472.1 AY057028.1 R satrapa AL004329.1 AY443327.1 0.002 0.017 KW434134.1 AY228802.1 Vanellus vanellus L satrapa AL004329.1 AY443327.1 0.002 0.017 AF197R35.1 AY228802.1 T. philomelos AY494511.1 AY307214.1 0.196 0.013 AY948118.1 KR350710.1 Conolophus subcristatus AY494511.1 AY307214.1 0.196 0.013 H AF181042.1 JN112592.1 A trinitatis AY494512.1 AY307214.1 0.196 0.009 EU057682.1 JN112592.1 A trinitatis AY494512.1 AY307214.1 0.196 0.009 EU037682.1 JN112592.1 L individus AY494512.1 AY307214.1 0.196 0.001 C cubiaris C collaris C collaris EU037482	Hirundo rustica	DQ119526.1	AY443290.1	H. pyrrhonota	AF074591.1	AY056997.1	0.181	0.010	18.0
HOGOB821.1 P. trochilus MHO79362.1 0.146 AV329472.1 AV057028.1 Researmericana L78808.1 AV443327.1 0.0084 AV329472.1 AV228802.1 Vanellus vanellus L78808.1 AV433327.1 0.002 0.017 KM434134.1 AY228802.1 Vanellus vanellus L7867-14802 AV339126.1 0.260 0.052 AF197835.1 KC789829.1 T. philomelos AV495411.1 AY339126.1 0.260 0.003 AF197835.1 KC789829.1 T. philomelos AV495411.1 AY339126.1 0.006 0.001 AF197835.1 KR7850703.1 Arinitadis AF495592.1 MI12645.1 0.286 0.009 HAF181042.1 KR780621.1 Arinitadis AF181056.1 EL05749.1 0.137 0.007 GUS31976.1 C. collaris EU037482.1 EL05749.1 0.137 0.007 GUS31975.1 KY762187.1 L. monicola HQ23497.1 EF632220.1 0.023 KK610607.1 KS610607.1 L. monicola LAF020251.1	Larus canus	AB208756.1		L. ridibundus	FM209923.1		0.056		
U76054.1 Rhea americana L78808.1 0.084 AV329472.1 AV057028.1 R satrapa AU004232.1 0.002 0.017 KM434134.1 AV228802.1 Vanellus vanellus KM577158.1: AV339126.1 0.200 0.0052 KM434134.1 AY228802.1 T. philomelos AV494812.1 AV339126.1 0.200 0.0052 AY948118.1 KR7350710.1 Conolophus subcristatus AY4948122.1 KR350708.1 0.196 0.003 H AF181042.1 LUST A trinidais AF49359.1 INI 12645.1 0.286 0.009 H AF181042.1 LUST A trinidais AF181056.1 AF181056.1 0.229 0.001 H AF181042.1 LUST C collents EU057482.1 AF181056.1 0.137 0.007 CUST C collents EU057482.1 AF181056.1 0.137 0.007 HQ234901.1 KY762187.1 I. monticola HQ234897.1 AF42220.1 0.050 KK510607.1 Liganaa Liganaa AF420502.1 AF420502.1 <td>Phylloscopus collybita</td> <td>HQ608821.1</td> <td></td> <td>P. trochilus</td> <td>MH079362.1</td> <td></td> <td>0.146</td> <td></td> <td></td>	Phylloscopus collybita	HQ608821.1		P. trochilus	MH079362.1		0.146		
AY32942.1 AY057028.1 R. satrapa A0004329.1 AV443327.1 0.002 0.017 KM434134.1 AY228802.1 Vanellus vanellus KM577158.1: AY339126.1 0.026 0.052 AF197835.1 KC789829.1 T. philomelos AY94811.1 AY350714.1 0.196 0.013 AY948118.1 KR350710.1 Conolophus subcristatus AY948122.1 KR350708.1 0.219 0.003 H AF181042.1 AT181056.1 Crubriqularis South AF493592.1 JN112645.1 0.219 0.009 H CUSTAB2.1 MK78062.1.1 C collaris C collaris<	Pterocnemia pennata	U76054.1		Rhea americana	L78808.1		0.084		
KM434134.1 AY228802.1 Vanellus vanellus KM577158.1: AY339126.1 0.260 0.052 AF197835.1 KC789829.1 T. philomelos AY495411.1 AY307214.1 0.196 0.013 AF197835.1 KC789829.1 T. philomelos AY495411.1 AY307214.1 0.196 0.013 AY948118.1 KR350710.1 Conolophus subcristatus AY495411.1 AY307214.1 0.196 0.003 H AF181042.1 Lubrigularis south AF181056.1 AF181056.1 0.226 0.009 H AF181042.1 C. collaris C.	Regulus calendula	AY329472.1	AY057028.1	R. satrapa	AJ004329.1	AY443327.1	0.002	0.017	0.1
AF197835.1 KC789829.1 T. philomelos AY9495411.1 AY307214.1 0.196 0.013 AY948118.1 KR350710.1 Conolophus subcristatus AY948122.1 KR350708.1 0.219 0.008 H AF181042.1 AF181042.1 A trinidatis AF181056.1 INI 12645.1 0.229 0.009 H AF181042.1 C collents C collents C collents C collents C collents 0.029 0.007 EU037370.1 MK780621.1 C collents C collents C collents 0.024 0.007 EU037370.1 MK7762187.1 L montroola HQ234897.1 EF632220.1 0.026 HQ234901.1 KY762187.1 L montroola AF020251.1 0.172 0.172 AF378067.1 K striata JQ639672.1 KY762190.1 0.172 0.223 AF378064.1 Immon lepidus K striata L cevesii L cevesii IM410538.1 0.024 IM410531.1 K striata L evesii IM410538.1 C uotatatatatatatatatatatatatatatatatatata	Scolopax rusticola	KM434134.1	AY228802.1	Vanellus vanellus	KM577158.1:	AY339126.1	0.260	0.052	5.0
AY948118.1 KR789829.1 T. philomelos AY495411.1 AY307214.1 0.196 0.013 AY948118.1 KR350710.1 Conolophus subcristatus AY948122.1 KR350708.1 0.219 0.008 H AF181042.1 Lubrigularis South AF181056.1 JN112645.1 0.229 0.009 H AF181042.1 C. collaris C. collaris C. collaris 0.025 0.009 EU037682.1 MK780621.1 C. collaris C. collaris 0.037482.1 0.137 0.007 EU037370.1 C. similis EU037415.1 EU03749.1 0.137 0.007 HQ234901.1 KY762187.1 I. monticola HQ234897.1 EF632220.1 0.050 0.001 HQ234901.1 KY762187.1 I. monticola AF030251.1 AF372020.1 0.023 0.001 AF373032.1 EF632222.1 I. striata JA639672.1 KY762190.1 0.204 AF378032.1 L. sevesii JN410538.1 EU054275.1 0.004 AF378032.1 L. gracilis JN410538.1 EU568094.1					13657-14802				
AY948118.1 KR350710.1 Conolophus subcristatus AY948122.1 KR350708.1 0.219 0.008 EU557103.1 JN112592.1 A trinitatis A trinitatis AF493592.1 JN112645.1 0.229 0.009 H AF181042.1 C rubrigularis South AF181056.1 EU03748.1 F1356749.1 0.229 0.009 CU331976.1 C similis C similis CU331975.1 EU037492.1 0.137 0.007 CU331976.1 C similis C wislizenii EU037415.1 EL037415.1 0.006 0.001 KX610607.1 KY762187.1 I monticola AF020251.1 0.050 0.001 KX610607.1 KY762187.1 L striata JQ639672.1 XY762190.1 0.223 AF378063.1 K striata JA620502.1 XY762190.1 0.204 NC014179.1 L reevesii JN410538.1 EU054275.1 0.204 NA10553.1.1 L gracilis JNA10538.1 EU568094.1 0.017 NV6581486.1 HM997172.1 R vianatense JN999978.1	Turdus migratorius Lizards	AF197835.1	KC789829.1	T. philomelos	AY495411.1	AY307214.1	0.196	0.013	15.4
PNTH AF181042.1 AF181056.1 JN112645.1 0.286 0.009 PNTH AF181042.1 C rubrigularis South AF181056.1 IN112645.1 0.229 EU037682.1 MK780621.1 C collaris EU037482.1 FJ356749.1 0.137 0.007 EU037370.1 C similis GU331975.1 EV037492.1 EV356749.1 0.184 0.007 HQ23490.1.1 KY762187.1 I. monticola AF024897.1 EF632220.1 0.001 KX610607.1 KX610607.1 I. guana AF37210.1 KY762190.1 0.172 AF373059.1 KX510607.1 KX610607.1 KY762190.1 0.204 0.001 AF378064.1 K7762190.1 KY762190.1 0.310 0.017 0.204 NC014179.1 L reevesii JN410538.1 KY762190.1 0.310 0.177 KC581486.1 HM997172.1 N. pelagicus KC581545.1 EU054275.1 0.081 0.001 KF691634.1 KF691634.1 KF691642.1 KJ195972.1 0.014 0.005 0.	Amblyrhynchus cristatus	AY948118.1	KR350710.1	Conolophus subcristatus	AY948122.1	KR350708.1	0.219	0.008	25.8
PRTH AF181042.1 C. rubrigularis South AF181056.1 0.229 es EU037682.1 MK780621.1 C. collaris EU037482.1 F1356749.1 0.137 0.007 es EU03731976.1 C. similis EU037415.1 EU037415.1 0.184 0.026 EU037370.1 G. wislizenii EU037415.1 EC0324897.1 EF632220.1 0.050 0.001 KX610607.1 KY762187.1 I. monticola AF020251.1 EF632220.1 0.050 0.001 KX610607.1 KY762187.1 I. iguana AF020251.1 KY762190.1 0.172 AF373032.1 AF572032.1 K. striata JQ639672.1 KY762190.1 0.310 AF378964.1 Imon lepidus JX626302.1 JX626302.1 0.204 NC014179.1 L. reevesii EU305052.1 L. reevesii EU305052.1 0.034 KX581486.1 HM997172.1 AV pelagicus JN410538.1 EU568094.1 0.017 JN999970.1 EU568093.1 AV dagaliii KF691642.1 K195972.1	Anolis aeneus	EU557103.1	JN112592.1	A. trinitatis	AF493592.1	JN112645.1	0.286	0.009	30.8
es EU037682.1 MK780621.1 C collaris EU037482.1 FJ356749.1 0.137 0.007 GU331976.1 C similis C similis GU331975.1 0.184 0.0184 EU037370.1 G. wislizenii EU037415.1 0.026 0.001 HQ234901.1 KY762187.1 I. monticola AF020251.1 0.050 0.001 KX610607.1 K. striata JQ639672.1 K7762190.1 0.172 0.017 AF373032.1 EF632222.1 L. schreiberi JX626302.1 0.204 0.024 NC014179.1 L. reevesii JNA10531.1 KY762190.1 0.310 0.024 NC114105.1 L. gracilis JNA10538.1 EU054275.1 0.039 0.077 KC581486.1 HM997172.1 N. pelagicus KC581545.1 EU054275.1 0.0177 JN999970.1 EU568093.1 O. waimatense JN999978.1 EU568094.1 0.014 AF3 KF691634.1 KF691637.1 KF691637.1 C. 0058 0.005	Carlia rubrigularis NORTH	AF181042.1		C. rubrigularis South	AF181056.1		0.229		
GU331976.1 C. similis GU331975.1 0.184 EU037370.1 G. wislizenii EU037415.1 0.026 HQ234901.1 KY762187.1 I. monticola HQ234897.1 EF632220.1 0.050 0.001 KX610607.1 I. iguana AF020251.1 AF020251.1 0.172 0.172 JQ639739.1 K. striata JQ639672.1 KY762190.1 0.223 0.223 AF378964.1 L. schreiberi JX626302.1 KY762190.1 0.310 0.017 NC014179.1 L. gracilis JN410538.1 L. gracilis JN410538.1 0.339 JN410531.1 HM997172.1 N. pelagicus KC581545.1 EU054275.1 0.004 JN999970.1 EU568093.1 O. waimatense JN999978.1 EU568094.1 0.0114 0.001 tai KF691634.1 KC581413.1 P. vlangalii KF691642.1 KJ195972.1 0.058 0.005	Crotaphytus bicinctores	EU037682.1	MK780621.1	C. collaris	EU037482.1	FJ356749.1	0.137	0.007	21.1
EU037370.1 G. wislizenii EU037415.1 0.026 HQ234901.1 KY762187.1 I. monticola AF020251.1 0.050 0.001 KX610607.1 I. iguana AF020251.1 0.172 0.172 JQ639739.1 K. striata JQ639672.1 KY762190.1 0.172 AF373032.1 EF632222.1 L. schreiberi AF372103.1 KY762190.1 0.204 NC014179.1 I. reevesii JNA10538.1 L. reevesii JNA10538.1 0.339 JNA10531.1 L. gracilis KC581545.1 EU054275.1 0.004 KC581486.1 HM997172.1 N. pelagicus JN999978.1 EU568094.1 0.017 JN999970.1 EU568093.1 P. vlangalii KF691642.1 KJ195972.1 0.058 0.005	Ctenosaura bakeri	GU331976.1		C. similis	GU331975.1		0.184		
HQ234901.1 KY762187.1 I. monticola HQ234897.1 EF632220.1 0.050 0.001 KX610607.1 I. iguana AF020251.1 0.172 0.172 JQ639739.1 K. striata JQ639672.1 0.223 AF373032.1 EF632222.1 L. schreiberi AF372103.1 KY762190.1 0.210 AF378964.1 Timon lepidus JX626302.1 0.204 0.204 NC014179.1 L. gracilis JNA10538.1 0.339 JNA10531.1 R. gracilis JNA10538.1 0.177 KC581486.1 HM997172.1 N. pelagicus KC581545.1 EU054275.1 0.004 JN999970.1 EU568093.1 O. waimatense JN999978.1 EU568094.1 0.014 0.005 tai KF691634.1 KC551413.1 P. vlangalii KF691642.1 KJ195972.1 0.058 0.005	Gambelia sila	EU037370.1		G. wislizenii	EU037415.1		0.026		
KX610607.1 Liguana AF020251.1 0.172 JQ639739.1 K. striata JQ639672.1 0.223 AF373032.1 E. schreiberi AF372103.1 KY762190.1 0.310 AF378964.1 Timon lepidus JX626302.1 0.204 NC014179.1 L. reevesii JNA10538.1 0.339 JNA10531.1 L. gracilis JNA10538.1 0.177 KC581486.1 HM997172.1 N. pelagicus KC581545.1 EU054275.1 0.004 JN999970.1 EU568093.1 O. waimatense JN999978.1 EU568094.1 0.0174 0.001 tai KF691634.1 KC551413.1 P. vlangalii KF691642.1 KJ195972.1 0.058 0.005	Iberolacerta galani	HQ234901.1	KY762187.1	l. monticola	HQ234897.1	EF632220.1	0.050	0.001	33.8
JQ639739.1 K. stnata JQ639672.1 0.223 AF373032.1 EF632222.1 L. schreiberi AF372103.1 KY762190.1 0.310 0.017 AF378964.1 Timon lepidus JX626302.1 0.204 0.204 NC014179.1 L. reevesii L. gracilis JN410538.1 0.339 JN410531.1 L. gracilis JN410538.1 0.177 KC581486.1 HM997172.1 N. pelagicus KC581545.1 EU568094.1 0.001 tai KF691634.1 KC551413.1 P. vlangallii KF691642.1 KJ195972.1 0.058 0.005	Iguana delicatissima	KX610607.1		l. iguana	AF020251.1		0.172		
AF373032.1 EF632222.1 L. schreiben AF372103.1 KY762190.1 0.310 0.017 AF378964.1 Timon lepidus JX626302.1 0.204 NC014179.1 L. reevesii EU305052.1 0.339 JN410531.1 L. gracilis JN410538.1 0.177 KC581486.1 HM997172.1 N. pelagicus KC581545.1 EU054275.1 0.081 JN999970.1 EU568093.1 O. waimatense JN999978.1 EU568094.1 0.114 0.001 tai KF691634.1 KC551413.1 P. vlangalii KF691642.1 KJ195972.1 0.058 0.005	Kentropyx calcarata	JQ639739.1		K. striata	JQ639672.1		0.223		,
AF378964.1 Timon lepidus JX626302.1 0.204 NC014179.1 L. reevesii EU305052.1 0.339 JN410531.1 L. gracilis JN410538.1 0.177 KC581486.1 HM997172.1 N. pelagicus KC581545.1 EU054275.1 0.081 JN999970.1 EU568093.1 O. waimatense JN999978.1 EU568094.1 0.114 0.001 tai KF691634.1 KC551413.1 P. vlangalii KF691642.1 KJ195972.1 0.058 0.005	Lacerta agilis	AF373032.1	EF632222.1	L. schreiberi	AF372103.1	KY762190.1	0.310	0.017	18.7
NC014179.1 L. reevesii EU305052.1 0.339 JN410531.1 L. gracilis JN410538.1 0.177 KC581486.1 HM997172.1 N. pelagicus KC581545.1 EU054275.1 0.081 0.004 JN999970.1 EU568093.1 O. waimatense JN999978.1 EU568094.1 0.114 0.001 tai KF691634.1 KC551413.1 P. vlangalii KF691642.1 KJ195972.1 0.058 0.005	Lacerta pater	AF378964.1		Timon lepidus	JX626302.1		0.204		
JN410531.1 L. gracilis JN410538.1 0.177 KC581486.1 HM997172.1 N. pelagicus KC581545.1 EU054275.1 0.081 0.004 JN999970.1 EU568093.1 O. waimatense JN999978.1 EU568094.1 0.114 0.001 tai KF691634.1 KC551413.1 P. vlangalii KF691642.1 KJ195972.1 0.058 0.005	Leiolepis guttata	NC014179.1		L. reevesii	EU305052.1		0.339		
; KC581486.1 HM997172.1 <i>N. pelagicus</i> KC581545.1 EU054275.1 0.081 0.004 JN999970.1 EU568093.1 <i>O. waimatense</i> JN999978.1 EU568094.1 0.114 0.001 tai KF691634.1 KC551413.1 <i>P. vlangalii</i> KF691642.1 KJ195972.1 0.058 0.005	Liolaemus bibronii	JN410531.1		L. gracilis	JN410538.1		0.177		,
JN999970.1 EU568093.1 <i>O. waimatense</i> JN999978.1 EU568094.1 0.114 0.001 tai KF691634.1 KC551413.1 <i>P. vlangalii</i> KF691642.1 KJ195972.1 0.058 0.005	Nactus multicarinatus	KC581486.1	HM997172.1	N. pelagicus	KC581545.1	EU054275.1	0.081	0.004	20.9
KF691654.1 KC551415.1 <i>P. Vlangalii</i> KF691642.1 KJ195972.1 0.058 0.005	Oligosoma otagense	JN9999970.1	EU568093.1	O. waimatense	JN999978.1	EU568094.1	0.114	0.001	96.4
	Phrynocephalus putjatai	KF691654.1	KC551415.1	P. Vlangalli	KF691642.1	KJ195972.1	0.058	0.005	10.6

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Table 2. Continued.	230220	Accession number		, and a very distribution of the second of t	Accession number	tono	ic distance	Constit distance (HKV85 T)
Species 1	MT-CYB	RAG1	Species 2	MT-CYB	RAG1	MT-CYB	RAG1	MT-CYB/RAG1
Phrynosoma cornutum Plestiodon japonicus Podarcis melisellensis Zootoca vivipara carniolice	AY141087.1 EU203134.1 AY185036.1 AY714929.1	DQ385423.1 HM161196.1	P. coronatum P. latiscutatus P. sicula Z. vivipara vivipara	AY141097.1 EU203035.1 AY770890.1 AY714913.1	FJ356738.1 HM161203.1	0.291 0.175 0.292 0.069	0.030	9.8
Mammals Alcelaphus buselaphus Arctocephalus pusillus Bos bison Camelus bactrianus Capra caucasica Cervus elaphus	A)222681.1 AM181018.1 AF036273.1 X177500.1 AF034738.1 AB001612.1	XM010964664.1	Damaliscus lunatus Zalophus californianus B. indicus C. dromedarius C. sibirica Odocoileus hemionus	AF016635.1 AM422164.1 AF419237.2 KU509220.1 KF990328.1 AF091630.1	XM011000597.1	0.161 0.088 0.089 0.170 0.110	0.004	48.2
Chinchilla brevicaudata Equus caballus Eulemur macaco Halichoerus grypus Hylobates agilis Kobus kob	AF464/56.1 DQ223535.1 AF081049.1 GU167293.1 AJ010583.1 AF052939.1 U06428.1	NM001256901.1 HM759153.1	C. Ianigera E. grewi E. mongoz Phoca hispida H. hoolock K. megaceros Vicugna vicugna	AF122820.1 X56282.1 AF081051.1 X82304.1 Y13304.1 AJ222686.1 U06430.1	AY239184.1 EU342315.1	0.069 0.118 0.140 0.043 0.178 0.083	0.009	13.1 27.4
Lepus europaeus Macaca fascicularis	AY/45112.1 AF295584.1	XM005578172.2	L. timidus Mandrillus sphinx	A2/9424.1 JQ068151.1	HM759046.1	0.328	0.001	353.2
Martes martes Mastomys natalensis Microtus californicus Ovis aries	AF154975.1 JX292865.1 AF163891.1 AF034730.1	DQ023475.1 KC953523.1 XM012134165.2	M. putorius Mus musculus M. montanus O. canadensis	X94925.1 AY057804.1 AF119280.1 EU365985.1	NM009019.2 KC953524.1 AY239177.1	0.218 0.317 0.189 0.089	0.055 0.0173 0.006	5.7 10.9 14.1
Panthera leo Pecari tajacu Peromyscus difficilis Spermophilus major Ursus americanus Vulpes lagopus	X82300.1 DQ179055.1 AF155394.1 AF157903.1 U23556.1 LT559489.1	AB109364.1 DQ240717.1 GU167551.1	P. pardus Tayassu pecari P. truei S. pygmaeus U. arctos V. vulpes	JF720058.1 AY534303.1 FJ800579.1 AF157907.1 U18870.1 AY928669.1	XM019470137.1 XM026511863.1 XM026012934.1	0.107 0.073 0.221 0.158 0.172	0.002	58.2 21.0 1.0
Salamanders Ambystoma dumerilii Ambystoma macrodactylum Ambystoma mexicanum Ambystoma talpoideum	AY659994.1 EF036633.1 AY659991.1 AY659991.1 MG822788.1	AY323752.1	A. mexicanum A. mexicanum A. tigrinum A. opacum A. texanum	AY659991.1 AY659991.1 AY69992.1 AY691730.1 GU078471.1	AY650130.1	0.053 0.208 0.075 0.256*	0.030*	8 *4.
Ambystoma texanum Andrias davidianus Batrachoseps gavilanensis Bolitoglossa franklini Desmognathus carolinensis Desmognathus fuscus	GUU78471.1 KU131042.1 KM203055.1 MK165231.1 EU314306.1 KY659020.1	MH106790.1 KM202898.1 KC614439.1 KR732369.1 KR732370.1 KR732372.1	A. rigrinum A. japonicus B. luciae B. lincolni D. orestes D. fuscus	AY659992.1 AB208679.1 KM203053.1 GU725464.1 EU314288.1 AY728227.1	AY583346.1 KM202878.1 KC614440.1 KR827015.1 KR732372.1 KR732377.1	0.172 0.099* 0.187* 0.030* 0.161 0.236*	0.004* 0.011* 0.003* 0.003 0.011*	21.1* 17.1* 12.0* 57.6 20.7*

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	Accession	Accession number		Accession	Accession number	Ceneti	ic distance (Genetic distance (HKY85 $+$ Γ)
Species 1	MT-CYB	RAG1	Species 2	MT-CYB	RAG1	MT-CYB	RAG1	MT-CYB/RAG1
Desmoanathus fuscus	AY728227.1	KR732372.1	D. santeetlah	EU314270.1	KR732384.1	0.196	0.012	16.3
Dicamptodon ensatus	AY734600.1	EF107335.1		AAW70411.1	AY650132.1	0.044*	0.005*	*0.8
Ensatina eschscholtzii croceater	L75796.1		E. eschscholtzii platensis	FJ151995.1		0.063		
Ensatina eschscholtzii eschscholtzii	FJ151951.1			L75801.1		0.181		
Ensatina eschscholtzii oregonensis	FJ151696.1			FJ151670.1		0.059		
Ensatina eschscholtzii oregonensis	FJ151696.1		E. eschscholtzii xanthoptica	FJ151887.1		0.223		
Ensatina eschscholtzii platensis	FJ151995.1		E. eschscholtzii xanthoptica	FJ151887.1		0.177		
Eurycea bislineata	AY728217.1	AY691706.1	E. cirrigera	NC035494.1	FJ750236.1	0.124*	0.017*	7.5*
Eurycea cirrigera	NC035494.1	FJ750236.1	E. wilderae	JQ920621.1	JQ920766.1	0.101	0.021	4.9
Eurycea neotenes	AY528400.1	AY650122.1	E. tridentifera	AY014848.1	KF562669.1	0.009	0.010	0.9
Hydromantes ambrosii	FJ602258.1	FJ602321.1	H. italicus	FJ602299.1	EU275791.1	0.059*	0.0008*	71.9*
Hynobius nebulosus	HM036356.1	AY650144.1	H. nigrescens	JQ929922.1	KJ715356.1	0.309*	0.010*	32.0*
Lissotriton helveticus	DQ821238.1		L. vulgaris	EU880339.1		0.360		
Lissotriton montandoni	DQ821254.1		L. vulgaris	EU880339.1		0.010		
Ommatotriton nesterovi	KX682106.1		O. ophryticus	DQ821267.1		0.011		
Ommatotriton ophryticus	DQ821267.1		O. vittatus	EU880338.1		0.014		
Plethodon aureolus	DQ994914.1	DQ995012.1	P. shermani	DQ994985.1	DQ995063.1	0.166	0.002	73.9
Plethodon chattahoochee	DQ994919.1	DQ995014.1		DQ994990.1	DQ995068.1	0.047	0.003	13.5
Plethodon chattahoochee	DQ994919.1	DQ995014.1		DQ994923.1	DQ995016.1	0.051	0.003	15.3
Plethodon chattahoochee	DQ994919.1	DQ995014.1		DQ994985.1	DQ995063.1	0.051	0.002	24.7
Plethodon cheoah	DQ994921.1			DQ994990.1		0.046		
Plethodon chlorobryonis	DQ994923.1		P. cylindraceus	DQ994928.1		0.016		
Plethodon chlorobryonis	DQ994923.1		P. teyahalee	DQ994990.1		0.011		
Plethodon chlorobryonis	DQ994923.1		P. metcalfi	DQ994956.1		0.160		
Plethodon cinereus	AY378042.1	AY691703.1	P. electromorphus	AY378060.1	DQ995024.1	0.151	0.013	11.3
Plethodon cylindraceus	DQ994928.1	DQ995022.1	P. glutinosus	DQ994937.1	DQ995027.1	0.169	0.003	56.1
Plethodon dorsalis	GQ464404.1	DQ995023.1		DQ994993.1	DQ995071.1	0.034	0.003	983
Plethodon electromorphus	AY378060.1	DQ995024.1	P. richmondi	AY378072.1	DQ995051.1	0.081	0.002	39.2
Plethodon fourchensis	FJ611481.1	DQ995026.1	P. ouachitae	FJ266744.1	AY691704.1	0.179	0.002	86.9
Plethodon glutinosus	DQ994937.1	DQ995027.1	P. jordani	DQ994947.1	DQ995032.1	0.184	0.002	82.0
Plethodon glutinosus	DQ994937.1	DQ995027.1		DQ994948.1	DQ995033.1	0.186*	0.005*	34.9*
Plethodon hoffmani	AY378047.1	DQ995029.1		AY378049.1	DQ995072.1	0.047	0.0007	68.4
Plethodon jordani	DQ994947.1	DQ995032.1	P. metcalfi	DQ994956.1	DQ995040.1	0.076	0.0007	111.2
Plethodon jordani	DQ994947.1	DQ995032.1		DQ994990.1	DQ995068.1	0.162	0.002	84.6
Plethodon metcalfi	DQ994956.1		P. teyahalee	DQ994990.1		0.156		
Plethodon shermani	DQ994985.1	DQ995063.1		DQ994990.1	DQ995068.1	0.003	0.001	3.3
Salamandra salamandra bernardezi	DQ092219.1			KX094979.1		0.024		
Salamandra salamandra tastuosa	UQ221254.1			AY222505.1		0.006		;
Salamandrella keyserlingii	AY593141.1	AY650145.1		AB363608.1	KJ855096.1	0.153*	0.004*	41.6*
Salamandrina perspicillata	DQ821207.1	HQ915345.1	S. terdigitata	EU880332.1	HQ915218.1	0.004*	0.007*	*9.0
Taricha rivularis	EU880334.1	AY650133.1	T. torosa	DQ196247.1	EF107340.1	0.161*	0.007*	21.9*
laricha rivularis T: L	EU880334.1		l. sierrae	DQ196282.1		0.167		
i aricna sierrae	UQ 196282.1		I. torosd	UQ196247.1		0.125		

Table 2. Continued.								
	Accession number	ımber		Accession number	mber	Gene	Genetic distance (HKY85 $+\Gamma$)	$KY85 + \Gamma)$
Species 1	MT-CYB	RAG1	Species 2	MT-CYB	RAG1	MT-CYB	RAG1	MT-CYB/RAG1
Triturus carnifex	NC 015788.1		T. cristatus	HQ697273.1		0.102		
Triturus carnifex	NC 015788.1		T. dobrogicus	HQ697274.1		0.104		
Triturus carnifex	NC 015788.1		T. macedonicus	HQ697278.1		0.058		
Triturus cristatus	HQ697273.1		T. dobrogicus	HQ697274.1		0.114		
Triturus cristatus	HQ697273.1		T. marmoratus	HQ697279.1		0.263		
Triturus cristatus	HQ697273.1		T. macedonicus	HQ697278.1		0.115		
Triturus marmoratus	HQ697279.1		T. pygmaeus	HQ697280.1		0.048		

list of hybrids for salamanders and lizards was also pared down to make results comparable. For salamanders, the dataset was pared down to a single pair that can hybridize per genus; all genera are monophyletic. When a genus had multiple hybridizing species pairs, the pair with the greatest MT-CYB genetic distance was used. The list of hybrids for lizards was also pared down to the largest MT-CYB genetic distance per genus based on previously published work (Jančúchová-Lásková et al., 2015). MT-CYB genetic distances between the parental species for each hybridizing species pair were calculated for birds, mammals, and lizards using the same methods as for salamanders (Fig. 3, Table 2). For each of the tetrapod groups, a different Γ distribution shape parameter α was estimated using all MT-CYB sequences included in this study (46 birds, 50 mammals, and 40 lizards; Kumar et al., 2018). A Kruskal-Wallis test was then used to determine if the genetic distance across parental species that hybridize differs among the tetrapod groups. Effect size was calculated and reported in terms of η^2 .

In order to test how representative MT-CYB genetic distances are in describing the overall genetic divergence between species, the substitution rate of mitochondrial genes relative to the substitution rate of nuclear genes was compared across the four vertebrate clades (Fig. 4). This was accomplished by dividing the genetic distance of the mitochondrial gene MT-CYB by the genetic distance of the nuclear gene RAG1 for the parental species that hybridize (Table 2). RAG1 genetic distances between the parental species for each hybridizing species pair were calculated for birds, mammals, and lizards using the same methods as for salamanders. A Kruskal-Wallis test was used to determine if the ratio of MT-CYB/RAG1 genetic distances differed among the four tetrapod groups. A Kruskal-Wallis test was also used to determine if RAG1 genetic distance differed among the four tetrapod groups. Effect size was calculated for both tests and reported in terms of η^2 .

Relationship between sex chromosomes and hybridization in salamanders.—A list of the salamanders that have heteromorphic sex chromosomes was generated to see if the presence of heteromorphic sex chromosomes impacted hybridization. Salamanders with heteromorphic sex chromosomes were identified by using two published reviews and a recently published amphibian karyotype database (Hillis and Green, 1990; Evans et al., 2012; Perkins et al., 2019). In the available literature, it is not always apparent whether a salamander species is known to have homomorphic sex chromosomes or if the species was merely never confirmed as having heteromorphic sex chromosomes (Perkins et al., 2019). To further clarify, additional unpublished information on salamanders that are known to have homomorphic sex chromosomes was obtained (S. Sessions, pers. comm., 19 June 2019). MT-CYB genetic distances between parental species with heteromorphic sex chromosomes that hybridize were compared to the genetic distances between parental species with homomorphic sex chromosomes that hybridize using a t-test (Fig. 5). A comparison was also made between the overall percentage of salamanders known to hybridize and the percentage of salamanders with heteromorphic sex chromosomes known to hybridize using a one-sided exact test of goodness-of-fit. The overall results were similar whether the analyses were performed using only the species confirmed to have homomorphic sex chromosomes (based

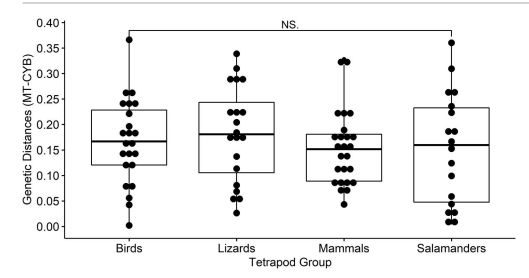


Fig. 3. Cytochrome b (MT-CYB) genetic distances between parental species that hybridize in different tetrapod groups. Kruskal-Wallis H test showed no significant differences across the groups (P=0.661). Salamanders are not hybridizing across greater genetic distances.

on personal communication) or using the larger dataset that assumed that all salamanders that have not been reported to have heteromorphic sex chromosomes have homomorphic sex chromosomes.

RESULTS

Hybrids in salamanders.—Our literature review found 81 pairs of genetically distinct salamander lineages that hybridize in nature (Table 1). These lineages represent 12.1% of named salamander species and seven out of ten salamander families (Fig. 1). The seven families represented include Plethondontidae (42 species), Salamandridae (29 species), Ambystomatidae (9 species), Hynobiidae (5 species), Cryptobranchidae (2 species), Dicamptodontidae (2 species), and Proteidae (2 species; Fig. 1). Many pairs that hybridize come from Plethondontidae and Salamandridae, consistent with these being the two largest families of salamanders (486 and 125 species, respectively; https://amphibiaweb.org). The third largest salamander family (Hynobiidae with 82 species) contains proportionally fewer hybrids with only three recorded pairs that can hybridize (Fig. 1). The families Amphiumidae, Rhyacotritonidae, and Sirenidae have no recorded cases of hybrids. These families of salamanders contain very few species with the largest of the three, Sirenidae, only containing five species. There is no case of hybridization occurring between species of different genera. Overall, these results are consistent with those of another recently published study (Patton et al., 2020).

Intensity of research on different salamander families.—We identified a total of 32,499 papers published using the salamander family related terms. Out of the large salamander families, the family with the highest number of papers per species, Ambystomatidae, did not have the highest number of reported hybrids (Fig. 1). However, Ambystoma mexicanum is a contributor to the high papers-per-species count because it is a model lab animal and makes up a quarter (2,694) of the Web of Science results for Ambystomatidae (Fig. 1; Voss et al., 2010). Two of the families with the highest numbers of species have the lowest intensity of publication: Hynobiidae (21 papers per species) and Plethondontidae (13 papers per species). There are proportionally more papers published for families with fewer species: Amphiumidae (270 papers per species), Cryptobranchidae (252 papers per species), Proteidae (249.5 papers per species), Dicamptodontidae (107 papers per species), Sirenidae (79 papers per species), and Rhyacotritonidae (44 papers per species). There was no significant correlation between the number of papers per species and the number of salamanders found to hybridize per family (Kendall's rank correlation P = 0.236). We

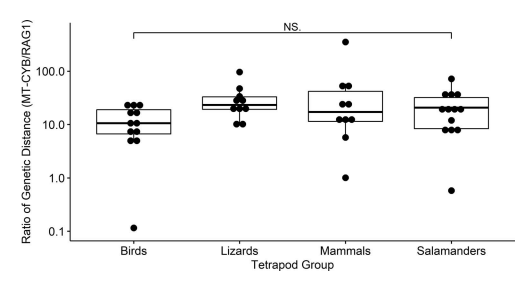


Fig. 4. Genetic distance ratio of mitochondrial cytochrome b (MT-CYB) to nuclear recombination activating 1 (RAG1) for pairs of species that hybridize in major tetrapod clade on a log scale. Kruskal-Wallis H test showed no strong significant differences across the four groups (P = 0.108).

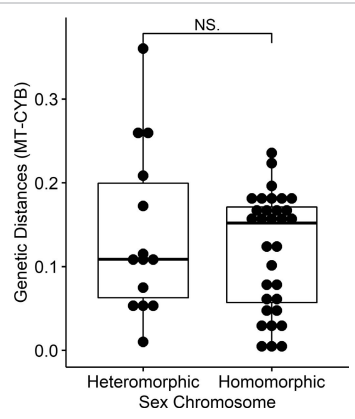


Fig. 5. Genetic distances between salamanders that hybridize with heteromorphic sex chromosomes (median = 0.109) and salamanders that hybridize without sex chromosomes (median = 0.152). There is no significant difference (P = 0.5293) between salamanders with sex chromosomes and salamanders without sex chromosomes.

acknowledge that studies of salamanders address far more biological questions than only those that are relevant to hybrid identification. Further analyses based on refined literature searches that target only the most pertinent disciplines (e.g., phylogeography, systematics) would provide a stronger test of the relationship between research effort and hybrid identification.

Salamander hybrid genetic distances.—For the 62 parental pairs of salamanders that have published MT-CYB sequence data, the mean genetic distance was 0.117 (Table 2). Lissotriton vulgaris X Lissotriton helveticus had the largest genetic distance of 0.360, while Plethodon teyahalee X Plethodon shermani had the smallest distance of 0.003 (Table 2). For the 33 parental pairs of salamanders that have published RAG1 data, the mean genetic distance was 0.00657 (range 0.000683 to 0.0306; Table 2). The genetic distance ratio of the mitochondrial gene MT-CYB over the nuclear gene RAG1 between salamanders that hybridize was highly variable (Fig. 2B). The ratio ranged from 0.6x to 162x with most species pairs having a higher mitochondrial genetic distance (median ratio = 21.1).

Genetic distances between parental species that hybridize in different tetrapod groups.—After paring down the salamanders and lizards to the pair that can hybridize with the highest genetic distance per genus, the across-tetrapod comparison included 23 pairs of hybrids for birds, 20 pairs for lizards, 25 pairs for mammals, and 18 pairs for salamanders (Table 2). The median MT-CYB genetic distances

for the groups were: lizards (0.181), birds (0.167), salamanders (0.160), and mammals (0.152; Fig. 3). There were no significant differences in distributions of genetic distance among the four groups ($\chi^2 = 1.5922$, P = 0.66, $\eta^2 = -0.0172$). Across all tetrapods, birds had both the lowest and highest genetic distances with Regulus calendula X Regulus satrapa (0.002) and Gallus gallus X Alectura lathami (0.366). Using the pared down list of hybrids, RAG1 sequences were available for 12 pairs of hybrids for birds, 10 pairs for lizards, 10 pairs for mammals, and 13 pairs for salamanders (Table 2). The median RAG1 genetic distances for the four groups were: birds (0.0151), mammals (0.00722), salamanders (0.00651), and lizards (0.00600). There were no significant differences among the four tetrapod groups ($\chi^2 = 5.8748$, P = 0.12; $\eta^2 =$ 0.0701). The median MT-CYB/RAG1 genetic distance ratio for the four groups were: lizards (23.4), salamanders (20.7), mammals (17.6), and birds (10.6; Fig. 4). There were no significant differences among the four tetrapod groups (χ^2 = 6.0734, P = 0.11; $\eta^2 = 0.075$).

Relationship between sex chromosomes and hybridization in salamanders.—Forty-eight species of salamanders in seven families have been discovered with heteromorphic sex chromosomes (Ambystomatidae, Hynobiidae, Plethodontidae, Proteidae, Salamandridae, Sirenidae, and Proteidae; Table 3). Salamander families are not characterized by a single type of sex chromosome; different members of Salamandridae and Plethodontidae have ZZ/ZW and XX/ XY systems. Out of the salamanders with known heteromorphic sex chromosomes, 12/48 are known to hybridize with a different species (Table 3). There is no difference between the percentage of salamanders that hybridize with known heteromorphic sex chromosomes and the known overall salamander hybridization rate (P = 0.99). Also, the genetic distances between parental salamanders that hybridize with known heteromorphic sex chromosomes (median = 0.109) and parental salamanders that hybridize with homomorphic sex chromosomes (median = 0.152) are not significantly different (P = 0.53; Fig. 5). Based on these analyses, there is no indication that the presence of heteromorphic sex chromosomes hinders hybridization in salamanders.

DISCUSSION

Overall, we found that hybridization is widespread throughout the salamander clade, occurring in both the biggest and smallest families (Fig. 1). The largest four families (Ambystomatidae, Hynobiidae, Plethodontidae, and Salamandridae) account for nearly 96% of all salamander species and account for over 91% of the hybrids (Fig. 1). With respect to geographic location, known hybrids are widespread in the Northern temperate zone, occurring in North America, Europe, and Asia. In contrast, there is only one reported case of salamanders hybridizing in the tropics, occurring between Bolitoglossa franklini and B. lincolni (Table 1). Bolitoglossa is the largest genus of salamanders (134 species), and it occurs exclusively in the tropics. More generally, close to half of all salamander diversity occurs in the tropics, suggesting that the low reported number of tropical hybrids is not a result of low tropical species diversity (https:// amphibiaweb.org). If the low reported number of hybrids is an accurate reflection of the underlying biological reality, it

Table 3. Heteromorphic sex chromosomes in salamanders.

Species	Family	Sex chromosome type	Hybrid (Yes/No
Ambystoma jeffersonianum ³	Ambystomatidae	ZW	No
Ambystoma laterale ^{1,2,3}	Ambystomatidae	ZW	No
Ambystoma mexicanum ^{1,2,3}	, Ambystomatidae	ZW	Yes
Ambystoma tigrinum ^{1,2,3}	, Ambystomatidae	ZW	Yes
Hynobius hidamontanus ^{1,3}	Hynobiidae	ZW	No
Hynobius quelpaertensis ^{1,3}	Hynobiidae	ZW	No
Hynobius tokyoensis ³	Hynobiidae	ZW	No
Aneides ferreus 1,2,3	Plethodontidae	ZW	No
Bolitoglossa subpalmata 1,2,3	Plethodontidae	XY	No
Chiropterotriton dimidiatus ^{1,2,3}	Plethodontidae	ZW	No
Cryptotriton veraepacis 1,2,3	Plethodontidae	XY	No
Dendrotriton bromeliacius ^{2,3}	Plethodontidae	XY	No
Dendrotriton cuchumatanus ²	Plethodontidae	XY	No
Dendrotriton rabbi ^{1,2,3}	Plethodontidae	XY	No
Dendrotriton xolocalcae 1,2,3	Plethodontidae	XY	No
Hydromantes ambrosii ^{1,2,3}	Plethodontidae	XY	Yes
Hydromantes flavus ^{1,2,3}	Plethodontidae	XY	No
Hydromantes imperialis ^{1,2,3}	Plethodontidae	XY	No
Hydromantes italicus ^{1,2,3}	Plethodontidae	XY	Yes
Hydromantes genei ³	Plethodontidae	XY	No
Hydromantes supramontis ^{1,3}	Plethodontidae	XY	No
Nototriton abscondens ^{1,3}		XY	
Notatritan picadai 1,2,3	Plethodontidae Plethodontidae	XY XY	No
Nototriton picadoi ^{1,2,3}		XY	No
Nototriton richardi	Plethodontidae		No
Oedipina cyclocauda 1	Plethodontidae	XY	No
Oedipina parvipes ^{1,2}	Plethodontidae	XY	No
Oedipina poelzi 1,2,3	Plethodontidae	XY	No
Oedipina pseudouniformis 1	Plethodontidae	XY	No
Oedipina uniformis ^{2,3}	Plethodontidae	XY	No
Thorius dubitus ^{1,2,3}	Plethodontidae	XY	No
Thorius pennatulus ^{2,3}	Plethodontidae	XY	No
Necturus alabamensis ³	Proteidae	XY	No
Necturus beyeri 1,2,3	Proteidae	XY	No
Necturus lewisi 1,2,3	Proteidae	XY	Yes
Necturus maculosus 1,2,3	Proteidae	XY	Yes
Necturus punctatus 1,2,3	Proteidae	XY	No
Ichthyosaura alpestris 3	Salamandridae	XY	No
Lissotriton boscai ³	Salamandridae	XY	No
Lissotriton helveticus ^{2,3}	Salamandridae	XY	Yes
Lissotriton italicus ^{2,3}	Salamandridae	XY	No
Lissotriton vulgaris ^{2,3}	Salamandridae	XY	Yes
Pleurodeles poireti 1,2,3	Salamandridae	ZW	Yes
Pleurodeles waltl 1,2,3	Salamandridae	ZW	No
Triturus carnifex ^{2,3}	Salamandridae	XY	Yes
Triturus cristatus 1,2,3	Salamandridae	XY	Yes
Triturus karelinii ^{1,3}	Salamandridae	XY	No
Triturus marmoratus ^{1,2,3}	Salamandridae	XY	Yes
Siren intermedia ^{1,2}	Sirenidae	ZW	No

¹ Evans et al., 2012

may reflect general differences in mechanisms underlying the origin and maintenance of species in temperate versus tropical areas worthy of further investigation (Kozak and Wiens, 2010). However, low reported numbers of hybrids could also mean that additional cases have yet to be discovered and described. Additional research will allow discrimination between these two possibilities.

Both the mitochondrial genetic distances (estimated with MT-CYB) and the nuclear genetic distances (estimated with RAG1) between parental species of salamanders that hybridize spanned two orders of magnitude (0.003 to 0.360 and 0.0007 to 0.020, respectively). The ratio of genetic distances between these two genes provides an estimate of the difference in substitution rates of the two genomes. In

² Hillis and Green, 1990

³ Perkins et al., 2019

vertebrates, the mitochondrial genome typically has a higher substitution rate than the nuclear genome (Brown et al., 1979). However, this ratio is variable across genes and species in all of the major vertebrate clades including amphibians, where the majority of the estimates are from frogs (Allio et al., 2017). Our salamander estimates (0.6 to 162, median = 21.1) are consistent with existing estimates for other vertebrate taxa, suggesting similar relative evolutionary rates between the two genomes. Because the MT-CYB and RAG1 sequences used are short (345–1,141 bp and 556–1,510 bp, respectively) and represent single loci, some of the variation in genetic distance estimates reflects sampling error.

The ratio between mitochondrial and nuclear genetic distances can also be used to identify candidate examples where loci have introgressed from one parental species to another. In our dataset, potential cases of mitochondrial introgression would show a ratio much less than the median (i.e., <<21.1) These potential cases include *Salamandrina perspicillata X Salamandrina terdigitata* (MT-CYB / RAG1 ratio = 0.6) and *Eurycea neotenes X Eurycea tridentifera* (MT-CYB / RAG1 ratio = 0.9; Table 2). In phylogeographic studies, hybridization between *Salamandrina perspicillata* and *S. terdigitata* was identified to have limited mitochondrial introgression (Mattoccia et al., 2011). Additional cline-based analysis of mitochondrial and nuclear introgression across the hybrid zones of salamanders with extreme mitochondrial/nuclear genetic distance ratios would be informative.

Within each tetrapod group, the genetic distance between parental species that hybridize is highly variable, but we found no significant differences in genetic distances among the groups. In addition, there were no significant differences in the ratio of mitochondrial to nuclear genetic distances among the groups. Taken together, these results suggest that speciating lineages of salamanders retain the ability to hybridize at similar overall genetic divergence levels as speciating lineages in other tetrapod groups. Thus, the differences in metabolic rates across tetrapods do not appear to translate into predictable patterns of overall genetic divergence and incompatible loci in hybrids. This pattern may reflect a similar evolved match between metabolic "supply and demand" (i.e., ATP use and ATP synthesis; Darveau et al., 2002) in all tetrapod clades, irrespective of overall metabolic rates, such that increased metabolic costs in hybrids have equal detrimental effects on fitness. Cline analyses of OXPHOS-specific loci, combined with OXPHOS functional data in parental species and hybrids, would allow more rigorous testing of this hypothesis.

The presence of heteromorphic sex chromosomes does not appear to deter salamanders from hybridizing (Fig. 3). This pattern suggests that, despite evolutionary lability in amphibian sex chromosomes (Hillis and Green, 1990; Evans et al., 2012; Sessions et al., 2016), the underlying genetic sexdetermining factors might be sufficiently conserved to not be a barrier to reproduction. Consistent with this explanation, for some salamanders, the difference between the heteromorphic sex chromosomes is extremely small (Schartl et al., 2016; Keinath et al., 2018), showing that the selective pressures might be similar on each chromosome or that the region of suppressed recombination on the X or Z chromosomes is small (unlike in birds or mammals; Schartl et al., 2016). Additionally, evidence that cryptobranchid salamander homomorphic sex chromosomes have been conserved for nearly 60 million years (Hime et al., 2019) suggests that the underlying genes for determining sex may be highly conserved. Finally, we note that the relationship between sex chromosome morphology and reproductive isolation is complex; in some cases, even homomorphic sex chromosomes show a large X/Z effect (e.g., *Hyla arborea* X *H. orientalis*, Dufresnes et al., 2016).

In conclusion, this comprehensive analysis of salamander hybrids revealed hybrids throughout the clade with a wide range of genetic divergences between parental species. Despite being outliers among tetrapods in metabolic rate and sex chromosome diversity, both of which were predicted to impact the relationship between genetic divergence and reproductive isolation, our analyses suggest that salamanders hybridize across similar levels of genetic divergence to the other examined tetrapod groups. This study further demonstrates the power of comparative studies of hybridization as an approach to understanding the process of speciation.

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