Taxonomic Uncertainty and Phylogenomics: Rescuing a Contentious Species

Complex from the Anomaly Zone (6438)

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Abstract (242/250)

Species are an indisputable unit for biodiversity conservation, yet their delimitation is fraught with both conceptual and methodological difficulties. A classic example is the taxonomic controversy surrounding the *Gila robusta* complex in the lower Colorado River of southwestern North America. Three species were originally defined according to subtle morphological differences with weak diagnostic power, with traditional genetic analyses failing to support the nominal species. Consequently, the complex was re-defined as a single polytypic unit, with the proposed ‘threatened’ status of two putative species withdrawn at the federal level. Here, we utilized dense spatial and genomic sampling (N=387 and >22k loci) to re-evaluate the status of the complex, based on SNP-based coalescent and polymorphism-aware phylogenetic models. In doing so, all three species were supported as evolutionarily independent lineages, despite widespread phylogenetic discordance. To understand this discrepancy with past studies, we categorized evolutionary mechanisms driving discordance. We tested (and subsequently rejected) prior hypotheses suggesting that phylogenetic discord in the complex was hybridization-driven. Instead, we found the *G. robusta* complex to have diverged within the ‘anomaly zone’ of tree space and, as such, have accumulated inconsistent patterns of diversity which have confounded prior studies. After extending these analyses with phylogeographic modeling, we propose that this is reflective of a rapid radiation promoted by Plio-Pleistocene tectonism. Our results not only support resurrection of the three species as distinct entities, but also offer an empirical example of how phylogenetic discordance can be categorized in other recalcitrant taxa.
Introduction (1260)

Complex evolutionary histories remain consistently difficult to disentangle, despite a recent paradigm shift towards the development of increasingly comprehensive datasets (e.g. Edwards 2009; Giarla and Esselstyn 2015). Regardless of these efforts, phylogenetic uncertainty is still prevalent, and with wide-ranging consequences on the study of macroevolutionary patterns (Stadler et al. 2016; Pereira and Schrago 2018), trait evolution (Hahn and Nakhleh 2016; Mendes et al. 2016; Wu et al. 2018), and ecological and biogeographic processes (Rangel et al. 2015; McVay et al. 2017).

Importantly, phylogenetic uncertainty also translates to taxonomic uncertainty. This is because modern systematic taxonomy fundamentally describes homology [i.e. Darwin's (1859) 'propinquity of descent' (Simpson 1961)], which, by definition, requires a phylogenetic context. Phylogenetic uncertainty in this sense can manifest itself as a soft polytomy (= 'honest' uncertainty), the erroneous promotion of non-monophyletic clades, or controversial ‘splitting’ versus ‘lumping.’ Incomplete or biased sampling is often a driver of this disparity (Ahrens et al. 2016; Reddy et al. 2017). Here, narrow taxon sampling may introduce substantial ascertainment bias (=systematic deviations due to sampling). On the other hand, a broader yet sparse sampling regime often fails to sample cryptic lineages (Heath et al. 2008) — with subsequent impacts on both the delimitation of species (Pante et al. 2015; Linck et al. 2019) and study of their traits (Beaulieu and O’Meara 2018).

These sources of uncertainty culminate in topologies that often fluctuate with regard to sampling designs or methodologies, and this results in taxonomic uncertainty [e.g. Ctenophora versus Porifera as sister to all other animals (Pisani et al. 2015; Whelan et al. 2015; Simion et al. 2017)]. Access to genome-scale data has alleviated some of these issues by offering a level of
precision not possible with single-gene phylogenies (Philippe et al. 2005). However, their inherent complexity and heterogeneity introduces new problems, and consequently, additional sources of phylogenetic uncertainty.

Gene tree heterogeneity is a ubiquitous source of discordance in genomic data, and “noise” as a source of this variance must consequently be partitioned from “signal” (where “noise” is broadly categorized as systematic or stochastic error). Large genomic datasets can reduce stochastic error (Kumar et al. 2012), yet it still remains a prevalent issue when individual genes are examined (Springer and Gatesy 2016). On the other hand, systematic error in phylogenomics may represent a probabilistic bias towards incongruence that is inherent to the evolutionary process itself (Maddison 1997). This, in turn, exemplifies the complications introduced by genomic data: As genomic resolution increases, so also does the probability of sampling unmodeled processes (Rannala and Yang 2008; Lemmon and Lemmon 2013). This potential (i.e., simultaneously decreasing stochastic error as systematic error increases) produces the very real possibility of building a highly supported tree that is ultimately incorrect.

Certain demographic histories are more predisposed to systematic error than others. For instance, when effective population sizes are large and speciation events exceptionally rapid, time between divergence events may be insufficient to sort ancestral variation, such that the most probable gene topology will conflict with the underlying species branching pattern. This results in what has been coined an “anomaly zone” of tree space (i.e., dominated by anomalous gene trees (AGTs); Degnan and Rosenberg 2006). Inferring species trees is demonstrably difficult in this region (Liu and Edwards 2009), and exceedingly so if additional sources of phylogenetic discordance, such as hybridization, are also apparent (Bangs et al. 2018).
In clades with such complex histories, it is often unclear where the source of poor support and/or topological conflict resides. Yet, to analytically account for gene tree conflict, it is necessary to categorize these sources and select approaches accordingly. Failure to do so promotes a false confidence in an erroneous topology, as driven by model misspecification (Philippe et al. 2011). The overwhelmingly parametric nature of modern phylogenetics insures that imperative issues will revolve around the processes being modeled, and what they actually allow us to ask from our data (Sullivan and Joyce 2005). However, the selection of methods that model processes of interest requires an *a priori* hypothesis that delimits which processes are involved. Yet, diagnosing prominent processes is difficult in that a phylogenetic context is required from which to build hypotheses. Fortunately, a wealth of information can be parsed from otherwise “non-phylogenetic” signal (*sensu* Philippe et al. 2005). For example, many statistical tests diagnose hybridization via its characteristic signature on the distribution of discordant topologies (e.g. Pease and Hahn 2015). Theoretical predictions regarding AGTs and the parameters under which they are generated are also well characterized (Degnan and Salter 2005; Degnan and Rosenberg 2009). Thus, by applying appropriate analytical approaches that sample many independently segregating regions of the genome, empiricists can still derive biologically meaningful phylogenies, despite the presence of complicated species-histories (McCormack et al. 2009; Kumar et al. 2012).

Here, we demonstrate an empirical approach that infers species-histories and sources of subtree discordance when conflict originates not only from anomaly zone divergences but also hybridization. To do so, we used SNP-based coalescent and polymorphism-aware phylogenetic methods (Chifman and Kubatko 2014; Leache et al. 2014; De Maio et al. 2015) that bypass the necessity of fully-resolved gene trees. We combine coalescent predictions, phylogenetic network
inference (Solís-Lemus and Ané 2016), and novel coalescent phylogeographic methods (Oaks 2018) to diagnose the sources of phylogenetic discordance and, by so doing, resolve a rather conflicted complex of study-species. We then contextualize our results to demonstrate the downstream implications of ‘problematic’ tree-space for threatened and endangered taxa, as represented by our study complex.

**Gila**

Few freshwater taxa have proven as problematic in recent years as the *Gila robusta* complex (Cyprinoidea: Leuciscidae) endemic to the Gila River basin of southwestern North America (Fig. 1). The taxonomic debate surrounding this complex exemplifies an inherent conflict between the traditional rigidity of systematic taxonomy versus the urgency of decision-making for conservation and management (Forest et al. 2015). Our study system is the Gila River, a primary tributary of the lower basin Colorado River that drains the majority of Arizona and ~11% of New Mexico. The critical shortage of water in this region (Sabo et al. 2010) is a major geopolitical driver for the taxonomic conflict surrounding our study species. As an example, the lower Colorado basin is responsible for approximately half of the total municipal and agricultural water requirements of the state of Arizona, and nearly two-thirds of its total gross state product (GSP) (Bureau of Reclamation 2012; James et al. 2014). This disproportionate regional reliance creates tension between the governance of a resource and its usage (e.g. Huckleberry and Potts 2019) which in turn magnifies the stakes involved in conservation policy (Minckley 1979; Carlson and Muth 1989; Minckley et al. 2006).

We focused on three species (Roundtail chub, *Gila robusta*; Gila chub, *G. intermedia*; and Headwater chub, *G. nigra*) that comprise a substantial proportion of the endemic
ichthyfaunal diversity of the Gila Basin [=20% of 15 extant native species (excluding extirpated

*Gila elegans* and *Xyrauchen texanus*); Minckley and Marsh 2009]. Historically, the focal taxa
have been subjected to numerous taxonomic rearrangements (Fig. 1). Until recently, the

consensus was defined by Minckley and DeMarais (2000) on the basis of morphometric and

meristic characters. These have since proven of limited diagnostic capacity in the field, thus

provoking numerous attempts to re-define morphological delimitations (Brandenburg et al. 2015;

Moran et al. 2017; Carter et al. 2018). Genetic approaches have to date been unproductive

(Schwemm 2006; Copus et al. 2018), leading to a recent taxonomic recommendation that

subsequently collapsed the complex into a single polytypic species (Page et al. 2016, 2017). For

the sake of clarity, we employ herein the nomenclature of Minckley and DeMarais (2000).



**Methods** (1514)

*Taxonomic sampling*

A representative panel of *N*=386 individuals (Table S1; Fig. 2) was chosen from existing

collections (Douglas et al. 2001; Douglas and Douglas 2007), to include broad geographic

sampling of the complex as well as congeners. For the sake of clarity, we retained species-level

nomenclature for all members of the *Gila robusta* complex, and additionally discriminate

between *G. robusta* from the upper and lower basins of the Colorado River ecosystem (Chafin et

al. 2019).

One species, *Gila elegans*, is extirpated from the wild, thus samples were provided by the

Southwestern Native Aquatic Resources and Recovery Center (Dexter, NM). The genus

*Ptychocheilus* served to root the *Gila* clade within the broader context of western leuciscids

(Schönhuth et al. 2012, 2014, 2018).
Reduced-representation sequencing

Genomic DNA was extracted using either PureGene® or DNeasy® kits (Qiagen Inc.) and quantified via fluorometer (Qubit™; Thermo-Fisher Scientific). Library preparations followed the published ddRAD protocol (Peterson et al. 2012). Restriction enzyme and size-selection ranges were first screened using an in silico procedure (Chafin et al. 2018), with the target fragment sizes further optimized by quantifying digests for 15 representative samples on an Agilent 2200 TapeStation. Final library preparations were double-digested using a high-fidelity PstI (5’-CTGCAG-3’) and MspI (5’-CCGG-3’) following manufacturer’s protocols (New England Biosciences). Digests were purified using bead purification (Ampure XP; Beckman-Coulter Inc.), and standardized at 100 ng per sample. Samples were then ligated with customized adapters containing unique in-line barcodes, pooled in sets of 48, and size-selected at 250-350bp (not including adapter length), using a Pippin Prep automated gel extraction instrument (Sage Sciences). Adapters were then extended in a 12-cycle PCR using Phusion high-fidelity DNA polymerase (New England Biosciences Inc.), completing adapters for Illumina sequencing and adding an i7 index. Libraries were pooled to N=96 samples per lane (i.e., 2 sets of 48) for 100bp single-end sequencing on an Illumina HiSeq 2500 at the University of Wisconsin Biotechnology Center (Madison, WI).

Data processing and assembly

Raw Illumina reads were demultiplexed and filtered using the pyRAD pipeline (Eaton 2014). We removed reads containing >1 mismatch in the barcode sequence, or >5 low-quality base-calls (Phred Q<20). Homologs assembly was then performed using de novo clustering in
VSEARCH (Rognes et al. 2016) using an 80% mismatch threshold. Loci were excluded according to following criteria: >5 ambiguous nucleotides; >10 heterozygous sites in the alignment; >2 haplotypes per individual; <20X and >500X sequencing depth per individual; >70% heterozygosity per-site among individuals.

Our ddRAD approach generated 22,768 loci containing a total of 173,719 variable sites, of which 21,717 were sampled (=1/ locus). Mean per-individual depth of coverage across all retained loci was 79X. All relevant scripts for post-assembly filtering and data conversion are available as open-source (github.com/tkchafin/scripts).

Phylogenetic inference

We formulated two simple hypotheses with regards to independent evolutionary sub-units. If populations represented a single polytypic species, then phylogenetic clustering should reflect intraspecific processes (e.g. structured according to stream heirarchy; Meffe and Vrijenhoek 1988). However, if a priori taxon assignments are evolutionarily independent, then they should be recapitulated in the phylogeny. Given well-known issues associated with application of supermatrix/ concatenation approaches (Degnan and Rosenberg 2006; Edwards et al. 2016) and pervasive gene-tree uncertainty associated with short loci (Leaché and Oaks 2017), we also employed SNP-based methods that bypassed the derivation of gene trees (Leaché and Oaks 2017).

We first explored population trees in SVDQUARTETS (Chifman and Kubatko 2014, 2015; as implemented in PAUP*, Swofford 2002) across 12 variably filtered datasets using four differing occupancy thresholds per SNP locus (i.e., 10, 25, 50, and 75%), along with three differing thresholds per individual (10, 25, and 50%). These filtered datasets ranged from 7357–
21007 SNPs, with 8.48–43.65% missing data and 256–347 individuals. SVDQUARTETS eases computation by inferring coalescent trees from randomly sampled quartets of species (i.e. optimizing among 3 possible unrooted topologies). It then generates a population tree with conflicts among quartet trees minimized via implementation of a quartet-assembly algorithm (Snir and Rao 2012). Given run-time constraints (the longest was 180 days on 44 cores), all runs sampled \( \binom{N_{tips}}{4}/2 \) quartets and were evaluated across 100 bootstrap pseudo-replicates.

We also used a polymorphism-aware method (PoMO; Schrempf et al. 2016) in IQ-TREE (Nguyen et al. 2014). PoMO considers allele frequencies rather than single nucleotides, thus allowing evaluation of change due to both substitution and drift. To provide PoMO with empirical estimates of polymorphism, we used the entire alignment, to include non-variable sequences. We filtered liberally using individual occupancy thresholds of 10% per-locus so as to maximize individual retention and per-population sample sizes. We then deleted populations that contained <2 individuals, and loci with >=90% missing data per-population. This yielded a dataset of 281,613 nucleotides and 40 tips. Non-focal outgroups were excluded due to their disproportionate effect on missing data.

We also calculated concordance factors (CFs) using a Bayesian concordance analysis in BUCKY (Larget et al. 2010), parallelized across all quartets via an adaptation of the TICR pipeline (Stenz et al. 2015). To prepare these data, we sampled all non-monomorphic full gene alignments for which at least 1 diploid genotype could be sampled per population. We excluded outgroups and non-focal Gila so as to maximize number of loci retained. This yielded 3,449 genes across 31 sampled tips. Gene-tree priors were generated using MrBayes v.3.2.6 (Ronquist et al. 2012) with 4 independent chains, each of which was sampled every 10,000 iterations, with a total chain length of 100,000,000 iterations and 50% discarded as burn-in. BUCKY was then
run in parallel to generate quartet CFs across 31,465 quartets, using a chain length of 10,000,000, again with 50% burn-in. Quartet topologies were used to generate a population tree using QUARTETMAXCUT (Snir and Rao 2012), using the get-pop-tree.pl script from TICR (Stenz et al. 2015; https://github.com/nstenz/TICR).

Comparing phylogenies and estimating site-wise conflict

To evaluate the performance of SVDQUARTETS, TICR, and PoMo, we first computed site-wise log-likelihood scores ($SLS$) for each topology by performing a constrained ML search in IQ-TREE. For comparison, we also generated an unconstrained concatenated tree. All ML analyses employed a GTR model with empirical base frequencies and gamma-distributed rates, and were assessed across 1,000 bootstrap pseudoreplicates. Analyses were also reduced to a subset of tips common across all variably filtered datasets. We quantified the phylogenetic signal supporting each resolution as the difference in site-wise log-likelihood scores ($\Delta SLS$) between each population tree and the concatenation tree (Shen et al. 2017). We then calculated site-wise concordance factors ($sCF$) as an additional support metric (Minh et al. 2018).

Tests of hybridization and deep-time reticulation

$D$-statistics (Green et al. 2010; Eaton and Ree 2013) were calculated using COMP-D (Mussmann et al. 2019). To further test hypotheses of reticulation, we used quartet CFs as input for phylogenetic network inference using the SNAQ algorithm in PHYLONETWORKS (Solís-Lemus and Ané 2016; Solís-Lemus et al. 2017). The network was estimated under models of 0–5 hybrid nodes ($h$). Models were evaluated using 48 independent replicates, with the best-fit model
being that which maximized change in pseudolikelihood. Given the computational constraints of network inference, we reduced the dataset to \( N=2 \) populations per focal species (=12 total tips).

Anomaly zone detection

Coalescent theory characterizes the boundaries of the anomaly zone in terms of branch lengths in coalescent units (Degnan and Rosenberg 2006). To test if contentious relationships in our tree fell within the anomaly zone, we first transformed branch lengths using quartet CFs (Stenz et al. 2015, equation 1), then tested if internode branch lengths fell within the theoretical boundary for the anomaly zone (Linkem et al. 2016, equation 1). Code for these calculations are modified from Linkem et al. (2016) and are available as open-source (github.com/tkchafin/anomaly_zone).

Tests of co-divergence

The contemporary course of the Colorado River resulted from the Pliocene erosion of the Grand Canyon (McKee et al. 1967; Minckley 1986). \textit{Gila} in the lower Colorado River basin then differentiated following one or more colonization events (e.g. Rinne 1976). Subsequent work (Douglas et al. 1999) supported this conclusion by examining contemporary phenotypic variation among all three species as a function of historical drainage connectivity, with the conclusion that body shape was most readily explained by Pliocene hydrography.

We tested if divergences were best explained by a model of \textit{in situ} diversification following a single colonization event, or instead by multiple, successive colonizations. We compared divergence models using a Bayesian approach (program \textsc{ecoevolity}, Oaks 2018) that used a coalescent model (Bryant et al. 2012) to update a prior expectation for the number of
evolutionary events across independent comparisons. Four independent MCMC chains were run with recommended settings and a burn-in that maximized effective sample sizes. Event models followed a Dirichlet process, with the concentration parameter exploring four alternative gamma distributed priors (i.e. $\alpha=2.0, \beta=5.70$; $\alpha=0.5, \beta=8.7$; $\alpha=1.0, \beta=0.45$; and $\alpha=2.0, \beta=2.18$).

We randomly sampled 2000 full-locus alignments, then examined potential co-divergences in the lower-basin complex by selecting a series of pairwise comparisons: *Gila elegans* x *G. robusta* (lower); *G. seminuda* x *G. robusta* (lower); *G. jordani* x *G. robusta* (lower); *G. intermedia* x *G. robusta* (lower); and *G. intermedia* x *G. nigra* (lower). These targeted nodes represent H, G, F, I, and N in the SVDQUARTETS topology (Fig. 3A).

Results (1147)

*Phylogenetic conflict in Gila*

We found tree reconstructions across all three population methods relatively congruent (SVDQUARTETS = Fig. 3; TICR, and PoMO = Fig. 4). The concatenated supermatrix tree (Fig. S1) was also largely congruent with the population trees, but with two major disparities (discussed below). Bootstrap support was variable and declined with decreasing node depth in the SVDQUARTETS analysis (Fig. 3), whereas the vast majority of nodes in PoMO were supported at 100% (Fig. 4A).

All analyses consistently supported the monophyly of a clade consisting of *G. intermedia*, *G. nigra*, and lower basin *G. robusta* (hereafter the ‘lower basin complex’). This clade had high bootstrap support in both SVDQUARTETS and PoMO, and was universally placed as sister to *G. jordani*. *Gila robusta* was unequivocally polyphyletic in all analyses, forming two distinct clades separated by the Grand Canyon. The lower basin clade was monophyletic in all cases, save the
concatenated tree, where it was paraphyletic (Fig. S1). It was also consistently recovered as sister to a monophyletic *G. nigra* + *G. intermedia*, with the exclusion of a single sample site (Aravaipa Creek) that nested within *G. intermedia* in the PoMo tree. Of note, this population had been previously diagnosed as trending towards *G. intermedia* in terms of morphology (Rinne 1976; Demarais 1986), although hybridization was not supported by *D*-statistics (Table 1).

Topology within the *G. nigra* + *G. intermedia* clade was less consistent. Both were reciprocally monophyletic in the SVDQUARTETS tree (albeit with low support; Fig. 3), whereas PoMo yielded a monophyletic *G. intermedia*, with but one population (Spring Creek) contained within *G. nigra* (Fig. 4A). The PoMo tree also conflicted with the other methods in its paraphyletic placement of upper basin *G. robusta*. We suspect this represents an artefact of well-known hybridization with sympatric *G. cypha* (Dowling and DeMarais 1993; Gerber et al. 2001; Douglas and Douglas 2007; Chafin et al. 2019).

**Discriminating among sources of phylogenetic conflict**

Phylogenetic conflict was variably attributable to either hybridization or rapid divergence. We found support for a single reticulation event connecting *G. seminuda* and *G. elegans*, an hypothesis consistent with prior interpretations (DeMarais et al. 1992). This particular model (i.e., *h*=1) was selected as that which maximized both the first \[L'(h) = L(h) - L(h-1)\] and second order \[L''(h) = L'(h+1) - L'(h)\] rate of change in pseudolikelihood (Fig. S2; following Evanno et al. 2005). Of note, introgression between *G. elegans* and *G. seminuda* was supported by elevated values of *h*, and by *D*-statistics \(\overline{D} = 0.302\) across 86,400 tests; Table 1). Introgression between upper basin *G. robusta* and *G. cypha* was also supported \(\overline{D} = -0.236\) across 45,056 tests). No other introgressions were noted, thus rejecting the hypothesized hybrid
origins for both *G. jordani* (Dowling and DeMarais 1993; Dowling and Secor 1997) and *G. nigra* (Demarais 1986; Minckley and DeMarais 2000).

Multiple internode pairs were observed in the anomaly zone (Fig. 5). In all cases, internode branches separating *G. nigra* and *G. intermedia*, and those separating their constituent lineages, reflected coalescent lengths that would yield anomalous gene trees. Not surprisingly, the internode separating *G. jordani* from the lower basin complex, and that of *G. robusta* from *G. intermedia/G. nigra* (Fig. 5C; tan branches) also fell within the anomaly zone, per TICR and concatenated topology results.

**Relative performance of species-tree methods**

Change in site-likelihoods among constrained and unconstrained IQ-TREE searches in all cases suggested that our recovered species-trees were supported by a minority of sites (Fig. S3), an observation consistent with tree regions being in the anomaly zone. Several discrepancies also reflected idiosyncrasies among the different approaches. For example, the PoMO topology has a paraphyletic upper basin *G. robusta* within which *G. elegans, G. cypha, G. seminuda, G. jordani,* and the lower basin complex were subsumed (Fig. 4A). However, only ~10% of SNPs supported this resolution (Fig. S4), a value far below the theoretical minimum sCF derived from completely random data (Minh et al. 2018). Of note, hybridization is a well-known artefact when a bifurcating tree is inferred from reticulated species (Sosef 1997; Schmidt-Lebuhn 2012), with concatenation or binning approaches using genomic data being demonstrably vulnerable (Bangs et al. 2018). Thus, we tentatively attribute the observed paraphyly as an artefact of documented hybridization between *G. cypha* and *G. robusta* (Chafin et al. 2019), and the inability of PoMO to
model hybridization. Hybridization also potentially drives the lack of monophyly in *G. seminuda*, per TICR and the concatenation tree (Fig. S1).

We also explored the impact of matrix occupancy filters on SVDQUARTETS, and bootstrap support and overall topological consistency declining with increasingly stringent filters (Fig. 3b). This corroborates prior evaluations with regard to the impacts of over-filtering RADseq data (Eaton et al. 2017). In all cases, site-wide concordance was significantly predicted by subtending branch lengths, but not by node depths (Fig. S5). This suggests that site-wise concordance was unbiased in our analyses at either shallower or deeper timescales, but was affected instead by the extent of time separating divergences. Some bioinformatic biases such as ortholog misidentification or lineage-specific locus dropout will disproportionally affect deeper nodes (Eaton 2017). However, we interpret the lack of correlation between node depth and site-wise concordance as an indication that these processes lack substantial bias.

**Biogeographic hypotheses and co-divergence**

ECOEVOLITY used four independent MCMC chains that sampled posterior parameter space over 300,000 iterations. Burn-in was determined as the number of samples removed to optimize effective sample sizes (ESS>500 in all cases following 10-30% burn-in). Model selection was not found to be vulnerable to alternative event priors (Fig. S6). The best-fitting model across all priors consistently demonstrated co-divergence of *G. jordani* with the lower basin complex (*G. robusta x G. intermedia* and *G. intermedia x G. nigra*; Fig. 6). The divergence of *G. elegans* and *G. seminuda* from a theoretical lower basin ancestor pre-dates this putatively rapid radiation, although it is unclear if these estimates were impacted by the aforementioned introgression between *G. seminuda* and *G. elegans*. 
Posterior effective population size \((N_e)\) estimates were large (e.g. >20,000) and consistent with previous estimates (Garrigan et al. 2002). *Gila jordani* was an exception, with a mean posterior \(N_e=6,062\). We find this discrepancy rather unsurprising, given the extremely narrow endemism of this species (Tuttle and Scoppettone 1990), and its recent bottleneck (Hardy 1982), although this is still a rather large estimate given the latter. Posterior divergence time estimates suggested a late-Miocene/ early-Pliocene origin of *G. elegans*. Results for *G. seminuda* and the lower basin radiation indicated Pliocene and early Pleistocene divergences, respectively. These results are supported in the fossil record (Uyeno 1960; Uyeno and Miller 1963), although we note paleontological evaluations of *Gila* have been sparse. Thus, we hesitate to interpret these as absolute dates, given our fixed mutation rate for these analyses and an uncertainty regarding the capacity of RADseq methods to yield an unbiased sampling of genome-wide mutation rate variation (e.g. Cariou et al. 2016).

**Discussion** \((1962 + 313 = 2275)\)

The goal of our study was to determine if extensive geographic and genomic sampling could resolve the taxonomically recalcitrant *G. robusta* complex. We applied diverse phylogenetic models and tests of hybridization and predictions of parameter space within the anomaly zone to diagnose sources of subtree discordance. In so doing, we also tested multiple hypothesized hybrid speciation events. We detected a single reticulation (*G. seminuda*), although other events with a lower component of genomic introgression may have also occurred. We documented rapid co-divergence of lower basin taxa within the anomaly zone and were able to resolve these despite the prevalence of incomplete lineage sorting. This scenario (as outlined
below) is consistent with the geomorphology of the region and seemingly represents an adaptive radiation by our study complex, as facilitated by drainage evolution.

Methodological artefacts and conflicting phylogenetic hypotheses for Gila

Increased geographic and genomic sampling revealed the presence of diagnosable lineages within the *G. robusta* complex, with both rapid and reticulate divergences influencing inter-locus conflict. Phylogenetic hypotheses for our focal group had previously been generated using allozymes (Dowling and DeMarais 1993), Sanger sequencing (Schwemm 2006; Schönhuth et al. 2014), microsatellites (Dowling et al. 2015), and more recently RADseq (Copus et al. 2018). None could resolve relationships within the lower basin complex. To explain these contrasts, we argue that prior studies suffered from systematic artefacts and ascertainment biases that were overcome, at least in part, by our approach.

Incomplete or biased sampling is a familiar problem for biologists (e.g. Hillis 1998; Schwartz and McKelvey 2009; Ahrens et al. 2016), and we suggest it served as a major stumbling block for delineating the evolutionary history of *Gila*. Although insufficient sampling is common in studies of threatened and endangered species, its repercussions are severe with regard to phylogenetic inference (Hillis 1998). This fact is substantiated by the many examples in which increasingly comprehensive geographic sampling spurred a revision of phylogenetic hypotheses (e.g. Oakey et al. 2004; Linck et al. 2019). Likewise, incomplete sampling of genome-wide topological variation (e.g. Maddison 1997; Degnan and Rosenberg 2009) is an additional source of bias, especially when a very small number of markers are sampled. These issues alone may explain the variation among prior studies. For example, Schwemm (2006) sampled extensively, including nearly all of the sites included in this study, but was only able to
examine a handful of genes. Because anomalous gene trees are most probable under a scenario of rapid radiation (as documented herein), the reduced number of loci used by Schwemm (2006) could not recover a consistent species tree. Copus et al. (2016, 2018) examined a dataset containing 6,658 genomic SNP loci (across 1,292 RAD contigs), but only did so across a sparse sample of 19 individuals. A bioinformatic acquisition bias also likely impacted this study, in the form of strict filtering that disproportionately excluded loci with higher mutation rates (Huang and Knowles 2016).

A necessary consideration when validating phylogenetic hypotheses across methods (and datasets) is to gauge compatibility between the underlying evolutionary processes and those actually being modeled. In this sense, the consideration of statistical support metrics alone can be not only misleading, but also promote false conclusions. For example, bootstrapping is by far the most prevalent method of evaluating support in phylogenetic datasets (Felsenstein 1985). While bootstrap concordances may be appropriate for moderately-sized sequence alignments (e.g. Efron et al. 1996), they can be meaningless when applied to sufficiently large datasets (Gadagkar et al. 2005; Kumar et al. 2012). This is apparent in the high bootstrap support displayed for anomalous relationships in our own analysis (Fig. S1). Phylogenetic signal also varies among loci, such that in many instances, relatively few loci drive contentious relationships (Shen et al. 2017). Likewise, not all methods are equal with respect to their simplifying assumptions. Given this, we feel it imperative to consider the biases and imperfections in both our data, and the models we apply.

*Complex evolution and biogeography of the Colorado River*
The taxonomic instability in *Gila* is not uncommon for fishes of western North America, where confusing patterns of diversity were generated by tectonism and vulcanism (Minckley et al. 1986; Spencer et al. 2008). This issue is particularly dramatized when viewed through the lens of modern drainage connections (Douglas et al. 1999). Historic patterns of drainage isolation and intermittent fluvial connectivity not only support our genomic conclusions but also summarize the paleohistory of the Colorado River over temporal and spatial scales.

The earliest record of fossil *Gila* from the ancestral Colorado River is mid-Miocene (Uyeno and Miller 1963), with subsequent Pliocene fossils representing typical ‘big river’ morphologies now associated with *G. elegans*, *G. cypha*, and *G. robusta* (Uyeno and Miller 1965). The modern Grand Canyon region lacked any fluvial connection at the Miocene-Pliocene transition, due largely to regional tectonic uplifts that subsequently diverted the Colorado River (Spencer et al. 2001; House et al. 2005). Flows initiated in early Pliocene (c.a. 4.9 mya; Sarna-Wojcicki et al. 2011), and subsequently formed a chain of downstream lakes associated with the Bouse Formation (Lucchitta 1972; Spencer and Patchett 2002). Evidence suggests ‘spillover’ by a successive string of Bouse Basin paleolakes was episodic, and culminated in mid-Pliocene (House et al. 2008), with an eventual marine connection via the Salton Trough to the Gulf of California (Dorsey et al. 2007). Prior to this, the Gila River also drained into the Gulf (Eberly and Stanley 1978), and sedimentary evidence indicated that it was isolated from the Colorado until at least mid-Pliocene by a northward extension of the Gulf (Helenes and Carreno 2014). This geomorphology is reflected in a broader phylogeographic pattern that underscores marked differences between resident fish communities in the upper and lower basins (Hubbs and Miller 1948).
Intra-basin diversification also occurred as an addendum to hydrologic evolution. Although the course of the pluvial White River is now generally dry, it may have been a Pliocene-early Pleistocene tributary of a paleolake system when the proto-Colorado first extended into the modern-day lower basin (Dickinson 2013). This may represent an initial colonization opportunity for upper basin fishes, an hypothesis that coincidentally aligns well with our rudimentary age estimate for Virgin River chub, *G. seminuda* (Fig. 6). This early isolation, as well as the continued contrast between the spring-fed habitats therein, and the high flows of the ancestral Colorado River, provide an explanation for the unique assemblage of *Gila* and other fishes therein (Hubbs and Miller 1948).

Phylogenetic signatures of the anomaly zone (Fig. 5) coupled with co-divergence modeling (Fig. 6) suggest the diversification of lower basin *Gila* occurred rapidly post-colonization. Late Pliocene integration of the two basins provided an opportunity for dispersal into the lower basin tributaries. The Plio-Pleistocene climate of the region was quite different, with a relatively mesic Pliocene as precursor to a protracted monsoonal period extending through early Pleistocene (Thompson 1991; Smith et al. 1993). The latter, in turn, may have resulted in relatively unstable drainage connections (Huckleberry 1996). The potential for climate-driven instability, and the complex history of intra-drainage integration of Gila tributaries during the Plio-Pleistocene (Dickinson 2015), lends support to the ‘cyclical-vicariance’ model proposed by Douglas et al. (1999). These periods of isolation may have promoted an accumulation of ecological divergences that persisted post-contact, and were sufficient to maintain species boundaries despite contemporary sympatric distributions and weak morphological differentiation. This hypothesis is also supported by the non-random mating found among *G. robusta* and *G. nigra*, despite anthropogenically-induced contact (Marsh et al. 2017).
Management implications

A request by the Arizona Game and Fish Department to review the taxonomy of the *Gila robusta* complex prompted the American Fisheries Society (AFS) and the American Society of Ichthyology and Herpetology (ASIH) to recommend the synonymization of *G. intermedia* and *G. nigra* with *G. robusta*, owing in part to their morphological ambiguity and an imprecise taxonomic key (Carter et al. 2018). Given this, a proposal to extend protection to lower basin *G. robusta* and *G. nigra* at the federal level was subsequently withdrawn (USFWS 2017; Fig. 1). Prior to this, *G. intermedia* alone was classified as endangered (USFWS 2005) under the Endangered Species Act (ESA 1973; 16 U.S.C. § 1531 et seq).

This study provides a much needed resolution to this debate by defining several aspects: First, our study reinforced the recognition of *G. robusta* as demonstrably polyphyletic, with two discrete, allopatric clades corresponding to the upper and lower basins of the Colorado River (Dowling and DeMarais 1993; Schönhuth et al. 2014). These data, together with the geomorphic history of the region that promoted endemic fish diversification (as above), clearly reject ‘*G. robusta*’ as a descriptor of contemporary diversity. This underscores a major discrepancy in the taxonomic recommendations for the lower basin complex (Page et al. 2016). Given that the type locality of *G. robusta* is in the upper basin (i.e., the Little Colorado River), we note a pressing need either to determine taxonomic precedence for the lower basin ‘*G. robusta*,’ or to provide a novel designation. The potential resurrection of a synonym is a possibility, necessitating a detailed examinations of the type specimens prior to a formal recommendation. This may be appropriately adjudicated by the AFS-ASIH Names of Fishes panel, as a follow-up to their earlier involvement.
The situation with *G. intermedia* and *G. nigra* is slightly more ambiguous. The short internodes and anomaly zone divergences identified herein explain previous patterns found in population-level studies, with elevated among-population divergence but scant signal uniting species (Dowling et al. 2015). We also unequivocally rejected the previous hypothesis of hybrid speciation for *G. nigra* (Minckley and DeMarais 2000; Dowling et al. 2015).

Rather, intermediacy in the body shape of *G. nigra* reflects differences accumulated during historic isolation (Douglas et al. 1999) and/or the retention of an adaptive ecomorphology (Douglas and Matthews 1992). These hypotheses warrant further exploration, with provisional results employed in future management decisions (Forest et al. 2015). With regards to taxonomy, we confidently recommend that *G. intermedia* be resurrected, and that additional studies be implemented to dissect the potential distinctiveness of *G. nigra*. For management purposes, we echo a conservative, population-centric approach (previously argued for by Dowling et al. 2015; Marsh et al. 2017).

Three primary components of a ‘Darwinian shortfall’ in biodiversity conservation are recognized (Diniz-Filho et al. 2013): (i) The lack of comprehensive phylogenies; (ii) Uncertain branch lengths and divergence times; and (iii) insufficient models linking phylogenies with ecological and life-history traits. Taxonomic uncertainty in *Gila* is severely impacted by the first two of these, with taxonomic resolution prevented by the comingling of sparse phylogenetic coverage with temporal uncertainty. We must now address the relationships between ecology, life history, and phylogeny in *Gila*, so as to understand the manner by which phylogenetic groupings (identified herein) are appropriate as a surrogate for adaptive/ functional diversity. For example: To what degree are *Gila* in the lower basin ecologically non-exchangeable? How do
they vary in their respective life histories? Is reproductive segregation maintained in sympatry (as in Marsh et al. 2018), and if so, by what mechanism?

Conclusion (313)

The intractable phylogenetic relationships in *Gila* were resolved herein through improved spatial and genomic sampling. Our data, coupled with polymorphism-aware methods and contemporary approaches that infer trees, yielded a revised taxonomy for *Gila* in the lower Colorado basin. The geomorphic history of the Colorado River served to explain many anomalous patterns seen in this and previous studies, wherein opportunities for contact and colonization were driven by the tectonism that is characteristic of the region. The signal of rapid diversification is quite clear in our data, as interpreted from patterns inherent to phylogenetic discord. We emphasize that discordance in this sense does not necessarily represent measurement error or uncertainty, but rather an intrinsic component of phylogenetic variance that is not only expected within genomes (Maddison 1997), but also a necessary component from which to build hypotheses regarding the underlying evolutionary process (Hahn and Nakhleh 2016). Ignoring this variance in pursuit of a ‘resolved phylogeny’ can lead to incorrect inferences driven by systematic error. Similarly, insufficient spatial or genomic sampling may also promote a false confidence in anomalous relationships, particularly when character sampling is particularly dense, whereas taxon sampling is sparse.

We reiterate that phylogenetic hypotheses, by their very nature, cannot exhaustively capture the underlying evolutionary process. One approach is to categorize phylogenetic (and “non-phylogenetic”) signals in those regions of the tree that are refractive to certain models (as done herein). We acknowledge that attempting to reconstruct the past using contemporary
observations is a battle against uncertainty and bias, with the evolution of phylogenetic/taxonomic hypotheses expected as additional data are accrued. As such, we urge empiricists that engage in taxonomic controversies (such as this one) to interrogate their results for transparency.

The task of sorting through conflicting recommendations invariably falls to managers, with unreported biases (be they methodological or geopolitical) only confounding those efforts.

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Table 1: Four-taxon $D$-statistic tests for admixture. Results are reported across $N$ separate quartet samples per four-taxon test, randomly sampled without replacement. Significance is reported as the proportion of tests at $p<0.05$ ($\text{nSig}/N$) using chi-squared ($\chi^2$), $Z$-test$^1$, and $Z$-test with Bonferroni correction$^2$. Positive and negative values of $D$ suggest introgression of the P3 lineage with either P2 or P1, respectively. Results in bold were also supported by the phylogenetic network.

| P3            | P2            | P1           | Mean $D$ | $N$       | $\text{nSig}/N$ ($\chi^2$) | $\text{nSig}/N$ ($Z^1$) | $\text{nSig}/N$ ($Z^2$) |
|---------------|---------------|--------------|----------|-----------|--------------------------|--------------------------|--------------------------|
| cypha jordani | lower basin   | 0.175        | 86400    | 0.033     | 0.072                    | 0.001                    |
| cypha seminuda| lower basin   | 0.099        | 86400    | 0.102     | 0.130                    | 0.002                    |
| elegans jordani| lower basin  | -0.063       | 84800    | 0.029     | 0.050                    | 0.000                    |
| elegans robusta (lower) nigra/int. | -0.026 | 413600 | 0.014 | 0.047 | 0.001 |
| elegans robusta (upper) cypha | -0.236 | 45056 | 0.380 | 0.415 | 0.045 |
| elegans seminuda | lower basin | 0.302 | 86400 | 0.654 | 0.674 | 0.251 |
| jordani robusta (lower) nigra/int. | 0.087 | 601600 | 0.042 | 0.072 | 0.001 |
| jordani robusta (lower) nigra/int. | 0.091 | 212800 | 0.041 | 0.067 | 0.005 |
| nigra int. (Salt) int. (Verde) | 0.086 | 126976 | 0.057 | 0.082 | 0.001 |
| robusta (lower) intermedia nigra | 0.041 | 793600 | 0.001 | 0.002 | 0.000 |
| robusta (upper) jordani robusta (lower) | 0.165 | 168000 | 0.050 | 0.081 | 0.001 |
| robusta (upper) robusta (lower) nigra/int. | -0.009 | 601600 | 0.011 | 0.031 | 0.000 |
| robusta (upper) seminuda lower basin | -0.017 | 180800 | 0.030 | 0.053 | 0.004 |
| seminuda jordani lower basin | -0.204 | 81920 | 0.107 | 0.152 | 0.000 |
| seminuda robusta (lower) nigra/int. | 0.054 | 212800 | 0.011 | 0.031 | 0.001 |
| atraria robusta (upper) cypha | 0.082 | 57344 | 0.064 | 0.095 | 0.033 |
| nigrescens robusta (lower) nigra/int. | -0.075 | 485472 | 0.023 | 0.079 | 0.002 |
| nigrescens robusta (upper) cypha | -0.039 | 53248 | 0.040 | 0.066 | 0.005 |
| pandora robusta (lower) nigra/int. | -0.123 | 225600 | 0.012 | 0.105 | 0.010 |
| pandora robusta (upper) cypha | -0.047 | 24576 | 0.031 | 0.057 | 0.003 |
Figure 1: Timeline of the conservation status of *Gila* species endemic to the lower Colorado River basin [*See Copus et al (2018) for a detailed overview of taxonomic synonymies; † ‘The Center’ refers to the Center for Biological Diversity (501c3), Tuscon, AZ; ‡ ‘DPS’ = Distinct Population Segment as referenced in the Endangered Species Act (ESA 1973; 16 U.S.C. § 1531 et seq), here referring specifically to a lower basin sub-unit of *G. robusta*. Note that timeline is not to scale.

- **Descriptions:**
  - *G. robusta*
  - *G. grahamii*
  - *G. nigra*
  - *G. intermedia* + synonyms

- **Rinne:**
  - *G. robusta robusta*
  - *G. robusta grahamii*
  - *G. nigra (=grahamii)*
  - *G. intermedia*

- **Minckley & DeMarais:**
  - *G. robusta*
  - *G. nigra (=grahamii)*
  - *G. intermedia*

- **The Center** petitions USFWS to list *nigra* and *robusta* DPS[^1]

- **The Center** challenges USFWS decision on *nigra* and *robusta*

- **Proposed threatened:**
  - *G. robusta DPS[^1]*
  - *G. nigra*
  - [80 FR 60753]

- **This study:**
  - 306 individ.
  - > 229k RAD I

- **Various taxonomic shuffling[^*]:**
  - All three species listed as ‘cat. 2 candidate species’ [47 FR 58454]
    - discontinued 1996

- **Proposed endangered:**
  - *G. intermedia* [70 FR 66663]

- **USFWS classifies *intermedia* endangered [70 FR 66663] and rejects proposals for *robusta* DPS and *nigra* [71 FR 26007]

- **USFWS gives DPS[^1] status to *robusta* [74 FR 32352]

- **Page et al:**
  - All species synonymized under “*G. robusta*”

- **USFWS withdraws proposed protection for *nigra* and *robusta* [82 FR 16681]
Figure 2: Sampling localities for *Gila* within the Colorado River Basin, southwestern North America. Locality codes are defined in Table S1. Sympatric locations (R14 and C2) are slightly offset for visibility purposes. Map insert increases the viewing scale for sampling sites within the lower basin ‘complex’ (Bill Williams and Gila Rivers).
Figure 3: (A) Majority-rule consensus cladogram of SVDQUARTETS across 12 variably filtered datasets. (B) Binned bootstrap concordance values are reported for each dataset, coded by the matrix occupancy threshold per individual (“i”) and per column (“c”; e.g. i50_c50 = 50% occupancy required per individual and per column). Dashed terminal branches indicate positions for taxa missing from >50% of datasets.
Figure 4: (A) PoMo phylogram with branch lengths as the number of substitutions and inferred number of drift events per site, with branch supports (as values <100%) representing concordance among 1000 bootstrap replicates; (B) TICR phylogram reporting branch lengths in coalescent units.
Figure 5: Internode pairs within the anomaly zone, as determined using coalescent-unit transformed branch lengths mapped onto the (A) SVDQUARTETS, (B) PoMo, (C) TICR, and (D) concatenated trees (displayed here as cladograms). Paired internodes are color-coded, with those bicolored indicating multiple anomalous divergences.
Figure 6: Posterior estimates for divergences times and effective populations sizes \( (N_e) \) derived from ECOEVOLITY. Branches are annotated with mean (std. dev.) \( N_e \) and posterior probabilities for divergence times are plotted on corresponding nodes. Units are in years, using a static mutation rate of \( 1.2 \times 10^{-8} \) substitutions per year. Posterior probabilities for divergence models (insert) suggest the co-divergence of *Gila jordani*, *G. robusta*, *G. nigra*, and *G. intermedia*.
| Site        | Major Drainage     | Location                                      | N  |
|-------------|--------------------|-----------------------------------------------|----|
| *Gila atraria* |                   |                                               |    |
|              | Bonneville, Snake  | Multiple localities                           | 14 |
| *Gila cypha*  |                   |                                               |    |
| C1          | Yampa              | Yampa Canyon, Moffat Co., CO                  | 1  |
| C2          | Colorado           | Black Rocks & Westwater Canyons, Colorado     | 2  |
| C3          | Colorado           | Grand Canyon, Coconino Co., AZ                | 9  |
| C4          | Little Colorado    | Little Colorado R. mainstem, Coconino Co., AZ | 4  |
| *Gila ditaenia* | de la Concepcion |                                               | 2  |
| *Gila elegans* | Hatchery         |                                               | 11 |
| *Gila eremica* | Rio Sonora        |                                               | 2  |
| *Gila intermedia* |                   |                                               |    |
| I1          | Verde              | Spring Creek, Yavapai Co., AZ                 | 10 |
| I2          | Agua Fria          | Silver Creek, Yavapai Co., AZ                 | 8  |
| I3          | Agua Fria          | Indian Creek, Yavapai Co., AZ                 | 9  |
| I4          | Agua Fria          | Sycamore Creek, Yavapai Co., AZ               | 10 |
| I5          | Gila               | Mineral Creek, Pinal Co., AZ                  | 10 |
| I6          | Gila               | Eagle Creek, Greenlee Co., AZ                 | 10 |
| I7          | Gila               | Hardin-Cienega Creek, Greenlee Co., AZ        | 10 |
| I8          | San Pedro          | Turkey Creek, Santa Cruz Co., AZ              | 10 |
| *Gila jordani* | Puhranagat/ White | Key-Pittman WMA refuge population (brood    | 21 |
|              |                   | stock sourced from Ash Springs), NV           |    |
| *Gila minacae* | Rio Yaqui         |                                               | 7  |
| *Gila nigra* |                   |                                               |    |
| N1          | Verde              | Fossil Creek, Yavapai Co., AZ                 | 11 |
| N2          | Verde              | Weber Creek, Gila Co., AZ                     | 8  |
| N3          | Verde              | Verde River headwaters, Gila Co., AZ          | 5  |
| N4          | Salt               | Gordon Creek, Gila Co., AZ                   | 10 |
| N5          | Salt               | Gun Creek, Gila Co., AZ                       | 11 |
| N6          | Salt               | Marsh Creek, Gila Co., AZ                     | 10 |
| *Gila nigrescens* | Multiple        | Multiple localities                           | 5  |
| *Gila pandora* | Rio Grande        | Palomas Creek, Sierra Co., NM                 | 6  |
| *Gila pulchra* | Rio Yaqui         | Rio Tomochic, Chihuahua, MX                   | 5  |
| *Gila purpurea* | Rio Yaqui         | San Bernadino NWR, Cochise Co., AZ            | 2  |
| *Gila robusta* |                   |                                               |    |
| R1          | Bill Williams      | Trout Creek, Mohave Co., AZ                   | 9  |
| R2          | Bill Williams      | Francis Creek, Yavapai Co., AZ                | 8  |
| R3          | Verde              | Verde River mainstem, Yavapai Co., AZ         | 8  |
| R4          | Verde              | Verde River mainstem, Yavapai Co., AZ         | 4  |
| R5          | Verde              | Verde River mainstem, Yavapai Co., AZ         | 3  |
| R6          | Verde              | West Clear Creek, Yavapai Co., AZ             | 5  |
| R7          | Salt               | Salt River mainstem, Maricopa Co., AZ         | 5  |
| R8          | Salt               | Cherry Creek, Gila Co., AZ                    | 8  |
| R9          | San Pedro          | Aravaipa Creek, Pinal Co., AZ                 | 10 |
| R10         | Green              | Upper Green River tributaries, Wyoming         | 21 |
| R11         | Yampa              | Little Snake River, Wyoming                   | 5  |
| R12         | Yampa              | Upper Yampa River tributaries, Moffat Co., CO | 5  |
| R13         | Green              | San Rafael River, Utah                       | 4  |
| R14         | Colorado           | Upper Colorado River mainstem, Colorado       | 4  |
| R15         | San Juan           | Navajo River, Colorado                       | 2  |
| R16         | Little Colorado    | East Clear Creek, Coconino Co., AZ           | 16 |
| V1          | Virgin             | Muddy (Moapa) River, Clark Co., NV            | 19 |
| V2   | Virgin        | Virgin River mainstem, Washington Co., UT | 17 |
|------|---------------|-------------------------------------------|----|
| Ptychocheilus |               |                                           |    |
| *P. grandis* | Eel River     | South Fork Eel River, Humboldt Co., CA    | 2  |
| *P. lucius*  | Colorado      | Yampa River, Moffat Co., Colorado          | 8  |

1Lower Colorado River basin (below Grand Canyon)
2Upper Colorado River basin (above Grand Canyon)
Figure S1: Phylogram showing results from an unconstrained search using concatenated SNP data in IQ-TREE. Focal nodes are annotated with bootstrap support (values for shallow nodes omitted for clarity).
Figure S2: Model selection results for SNAQ/PHYLONETWORKS; \( h \) = maximum number of hybrid edges allowed per model; (A) \( L(h) = -\log \) likelihood for the best network of \( N=48 \) replicate runs per value of \( h \); (B) \( L'(h) = 1^{st} \) order change in \( L(h) = L(h) - L(h-1) \); (C) \( L''(h) = 2^{nd} \) order change in \( L(h) = L'(h+1) - L'(h) \); and (D) \( \Delta h = L''(h) / s(h) \) where \( s(h) \) is the standard deviation in \( L(h) \) among replicates.
Figure S3: Site-wise log-likelihood differences ($\Delta SLS$) for (A) SVDQUARTETS, (B) PoMo, and (C) TICR topologies as compared to an unconstrained concatenated tree. $\Delta SLS$ values are transformed as signed square-roots, with positive values indicating increased site-likelihood under the constrained model, and negative values having increased likelihood under the unconstrained concatenated model.
Figure S4: Site-wise concordance factors (sCF) for lineage trees produced in IQ-TREE under topological constraints for the (A) SVDQUARTETS, (B) PoMo, and (C) TICR results.
Figure S5: Characterization of site-wise concordance ($s$CF) factors for SVDQUARTETS, PoMo, and TICR phylogenies. Panels show (left to right): Linear regression of subtending branch lengths (log-transformed) with $s$CF; node height (cumulative branch lengths from root to focal node); and densities of $s$CF across nodes as compared to the discordance factors for the two conflicting quartet resolutions ($s$DF1 and $s$DF2).
Figure S6: Prior and posterior probabilities for number of independent divergence events in EcoEvolity co-divergence models for *Gila*. Parameters across all runs were identical, except for the shape ($\alpha$) and scale ($\beta$) of the gamma-distributed prior on the Dirichlet process concentration.

\begin{align*}
\alpha &= 2.0; \beta = 5.70 \\
\alpha &= 0.5; \beta = 8.70 \\
\alpha &= 1.0; \beta = 0.45 \\
\alpha &= 2.0; \beta = 2.18
\end{align*}