



Research

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A community genomics approach to natural hybridization

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Hybridization is a complicated, oft-misunderstood process. Once deemed unnatural and uncommon, hybridization is now recognized as ubiquitous among species. But hybridization rates within and among communities are poorly understood despite the relevance to ecology, evolution and conservation. To clarify, we examined hybridization across 75 freshwater fish communities within the Ozarks of the North American Interior Highlands (USA) by single nucleotide polymorphism (SNP) genotyping 33 species ($N = 2865$ individuals; double-digest restriction site-associated DNA sequencing (ddRAD)). We found evidence of hybridization (70 putative hybrids; 2.4% of individuals) among 18 species-pairs involving 73% (24/33) of study species, with the majority being concentrated within one family (Leuciscidae/minnows; 15 species; 66 hybrids). Interspecific genetic exchange—or introgression—was evident from 24 backcrossed individuals (10/18 species-pairs). Hybrids occurred within 42 of 75 communities (56%). Four selected environmental variables (species richness, protected area extent, precipitation (May and annually)) exhibited 73–78% accuracy in predicting hybrid occurrence via random forest classification. Our community-level assessment identified hybridization as spatially widespread and environmentally dependent (albeit predominantly within one diverse, omnipresent family). Our approach provides a more holistic survey of natural hybridization by testing a wide range of species-pairs, thus contrasting with more conventional evaluations.

1. Introduction

Hybridization was once considered ‘exceedingly rare’ [1], but it is now acknowledged as relatively common, primarily because of better detection using modern DNA sequencing [2,3]. A concurrent recognition is the significant role of hybridization and introgression in ecology and evolution [4–6], which diverges sharply from the historical perspective rooted in the biological species concept [7]. But while a growing catalogue of species known to hybridize has helped to draw attention to hybridization (per-species documentation), the prevalence of hybrids within and among communities is poorly understood (per-individual documentation) [2].

The more frequently we encounter hybridization, the more evidence for its influential role in nature [2,8]. Hybridization and introgression are often considered maladaptive threats to biodiversity [6,9]. However, it is becoming more appreciated that the genetic novelty injected into a lineage through introgressive hybridization can be adaptive [8] and even provide evolutionary rescue [10]. Moreover, hybridization is expected to increase in lockstep with global environmental change [11,12]. Therefore, baseline estimates are needed to precisely gauge increases in hybridization and detect where it impacts ecosystems [10,13]. Attempts to understand the geographical patterns and processes of hybridization have not often considered an array of taxa. Doing so will help identify where hybridization occurs more generally and what environmental factors might drive it. These insights are essential because these places/

environments in which hybridization is promoted may disproportionately impact biodiversity and ecosystems [14].

Broad comparative examinations often employ meta-analyses of numerous single-species-pair studies to quantify hybridization rates [15] or assess biogeographic relationships [14,16,17]. Several generalizations have emerged. Hybridization tends to be unevenly distributed across taxonomic groups and occurs more in plants than animals [2]. It has been documented more so in fish than other vertebrates [18], probably due to external fertilization within the aquatic environment [19]. Among fish, evidence of hybridization is more prevalent in fresh waters [1], especially between carps and minnows (Cypriniformes) [19], owing to their high diversity, sympatry and breeding behaviours [20].

The less time since two species have diverged, the less time for pre- and post-zygotic barriers to evolve [20–22]. Similarly, the geographical overlap between two closely related species also controls the pressure to evolve reproductive barriers [23], with most hybridizing lineages first evolving in allopatry [24]. Pre-zygotic barriers are often keyed (directly or indirectly) with the environment [25,26], weakening concurrently with environmental fluctuations, thus facilitating hybridization. For example, many fish rely on coloration to identify conspecific mates, and any environmental alterations impacting visual acuity (e.g. turbidity, siltation) can promote hybridization [27]. Much work has focused on human impacts due to environmental homogenization, sensory cue changes and creating novel environments where hybrids may have advantages [28].

We quantified hybridization across a North American freshwater fish metacommunity by genotyping genome-wide single nucleotide polymorphisms (SNPs) that allowed us to detect hybrids among the individuals we sampled. We addressed the following questions: (i) How many hybrids occur within and among localities, and what is their frequency within species/families? (ii) Is genetic exchange among species (i.e. introgression) occurring as evinced by the presence of backcrossed individuals (e.g. $F_1 \times$ Parental)? (iii) Is the incidence of hybrid individuals within communities predicated on environmental factors which allow the prediction of hybrid occurrence?

2. Methods

(a) Data generation and processing

Our sampling area spanned the White River Basin (71 911 km²; Ozark Plateau, North American Interior Highlands). The area represents an ideal study location because it is: (i) an unglaciated refugium with elevated fish diversity and limited anthropogenic impacts [29–31]; and (ii) an excellent region from which generalizable patterns can be interpreted, because it is central to and representative of the Mississippi system [32]. Sampling procedures were approved by the University of Arkansas Institutional Animal Care and Use Committee (IACUC #17077) and appropriate permitting agencies (electronic supplementary material, S1). Fishes were seined (June 2017 to September 2018) and euthanized via tricaine methanesulfonate (MS-222) and 95% ethanol. Species diagnoses occurred in the laboratory.

Genomic DNA was isolated from fin clips (Qiagen Fast kits; Qiagen Inc.) and quantified by fluorometry (Qubit; ThermoFisher Scientific). Individuals were SNP genotyped using double-digest restriction site-associated DNA sequencing (ddRAD), using modified procedures [33,34], and sequenced as pooled batches ($N=144/1 \times 100$ lane) on the Illumina HiSeq 4000 (electronic supplementary material, S1).

Raw Illumina reads were demultiplexed (ipyrad; [35]), and family-level phylogenies were constructed to verify individual ID to species via an alignment- and assembly-free method (PHYLORAD; [36]; electronic supplementary material, S1). Individuals ($N=3042$) were grouped by family ($N=6$) and processed de novo in IPYRAD to generate separate family-level assemblies. Adapters/primers were removed, and reads with greater than five low-quality bases (Phred < 20) were discarded. Clusters were assembled using an 85% identity threshold, with loci subsequently removed via conditional criteria to ensure high-quality data (electronic supplementary material, S1). Biallelic SNPs were further filtered and visualized using RADIATOR [37] (electronic supplementary material, S1).

(b) Hybrid detection

Our initial objective was to detect putative hybrids and determine hybridization rates. We focused the search within families for two reasons: no evidence implicates hybridization among these divergent families within six different orders. Also, a high number of species in RADseq alignments can reduce the power to detect hybrids by diminishing recovery of homologous loci (via allelic dropout from accumulating mutations in restriction enzyme cut-sites) [38]. All analyses were conducted using R 4.1.3 [39]. Individual genetic variation was first visualized using separate principal components analyses (PCAs) on each family-level SNP alignment (ADE4; [40]).

Admixture analysis employing sparse non-negative matrix factorization (sNMF) was used to estimate ancestry coefficients for individuals within families (25 repetitions/ K value (1–25) with regularization parameter (α) = 100 (LEA; [41]). The best K from each run (via cross-validation) was used to impute missing data (*impute* function, method = 'mode' in LEA), with sNMF repeated using imputed data (as above). An individual was flagged as a putative hybrid if the assignment probability for the majority ancestry cluster was less than 0.9.

Results were contrasted with a maximum-likelihood clustering approach considering hybrid categories per expected allele frequencies (SNAPCLUST; [42]). Three models were tested: (i) F_1 only; (ii) F_1 + first-generation backcross; and (iii) F_1 + first- and second-generation backcrosses, with AIC used to determine the best fit [42]. Pairwise analyses were among species within families.

We assessed the power of sNMF and SNAPCLUST to detect hybrids. We first removed putative hybrids following the above analyses. Then, we sampled alleles from the remaining pure individuals for each within-family species-pair to simulate hybridization between parentals (P_1 and P_2). We simulated 10 each of F_1 , F_2 and backcross (5 $P_1 \times F_1$ and 5 $P_2 \times F_1$) (function *hybridize*; ADEGENET; [43]). SNPs for simulated hybrids were combined with parentals before running sNMF and SNAPCLUST (as above). We then quantified the proportion of simulated hybrids detected by each analysis.

(c) Hybrid classification

Hybrids detected with sNMF and SNAPCLUST were classified by assignment as F_1 , F_2 or backcross (NEWHYBRIDS; [44]). Backcrosses underscore hybrid viability and gene exchange among species (i.e. introgression). We implemented HYBRIDDETECTIVE [45] to confirm sufficient statistical power for classification (F_1 , F_2 or BXs). All NEWHYBRIDS analyses were conducted pairwise among species from which hybrids were detected. We first simulated known class hybrids by randomly sampling two alleles/locus from appropriate parental pools, with convergence assessed via three replicates. We then determined an optimal posterior probability threshold (= 0.70) from which hybrid classes could be reliably assigned. The final MCMC was 1 000 000 iterations (250 000 burn-in). Assignments were made using a reduced panel of 200 SNPs exhibiting the greatest among-species

differentiation (Weir and Cockerham's F_{ST} among target species-pair) and lowest linkage disequilibrium ($r^2 < 0.2$) via *getTopLoc*, *HYBRIDDETECTIVE* [45].

(d) Correlates of hybridization

We tested whether hybridizing/non-hybridizing species-pairs differed significantly with respect to genetic differentiation and basin-wide co-occurrence. Weir and Cockerham's F_{ST} (DIVERSITY; [46]) and the number of co-occurrences across communities (COOCCUR; [47]) were calculated among $N = 137$ species-pairs within families. Differences among hybridizing/non-hybridizing pairs were assessed using the Wilcoxon–Mann–Whitney test with a U statistic distribution approximated using Monte Carlo resampling of the data (100 000 permutations).

We evaluated the influence of environmental factors on the incidence of hybrids within communities to determine their predictability. Each community ($N = 75$) was classified as with/without hybrids, with potential predictor variables being: (i) species richness at each site; (ii) drainage identity (eight-digit USGS hydrologic unit); and (iii) high-resolution hydro-environmental descriptors ($N = 281$, RiverATLAS; [48]). These encompassed factors broadly related to hydrology, physiography, climate, land cover, geology and anthropogenic impact. Associations between hybrid incidence and predictive variables were assessed using random forest classification (i.e. supervised machine-learning employing linear/nonlinear relationships among mixed data types [49]). Bootstrapped decision trees (via predictor variables) are aggregated, with each trained on a random two-thirds of samples (= 'in-bag'), with validation via the remaining one-third (= 'out-of-bag').

We implemented the random forest quantile-classifier approach [50] in *RANDOMFORESTSRC* [51] using 100 000 bootstrapped decision trees (= *ntree*), each of which randomly sampled one-third of total predictors (= *mtry*). Variable importance (VIMP; increase in prediction error when a variable is randomized) was quantified via permutation for each predictive variable. VIMPs significantly greater than 0 were assessed by subsampling 1000 trees. Random forest variable selection was executed using minimal depth and a conservative threshold set to 'high', employing only that subset of predictive variables with significant VIMP [52]. A final random forest model was built from this selected subset of predictors. Accuracy was evaluated using G-mean (0–1; the geometric mean of the true negative and positive rates), which replaces the misclassification rate in imbalanced data settings [50] and the normalized Brier score (0–1; mean square difference between true classes and predicted probabilities).

3. Results

Fish collections across 75 communities/sites yielded 72 species ($N = 3605$ individuals), averaging 10.8 species/site—typical for streams across the Mississippi Basin [53]. We genotyped 33 species to represent the fish metacommunity [54] (i.e. 84% of total individuals collected, averaging 9.44 spp./site). Of $N = 137$ intra-familial species-pairs examined, 13 (9.5%) did not co-occur at our sites. Four do have non-overlapping, parapatric distributions, each with one occurring in the upper White River and the other in the Black River: one ictalurid (*Noturus albatere* / *maydeni*); one leuciscid (*Luxilus pilsbryi* / *zonatus*); and two percids (*Etheostoma juliae* / *uniporum*; *E. spectabile* / *uniporum*). Despite range overlap, nine other pairs did not co-occur (six leuciscids and three percids), which may be due to chance/sampling, given that pairs included at least one uncommon species (less than or equal to 10 occurrences).

We examined $N = 2865$ individuals across six families, post-filtering using SNP genotypes (table 1; mean missing data = 21%; mean coverage = 56 \times). SNPs varied by family and were inversely related to the number of species (table 1). Power analyses verified the robustness of panels for detecting/classifying hybrids based on genotype frequencies (electronic supplementary material, S2–S19). Among $N = 30$ simulated hybrids for each unique species-pair assessed ($N = 137$), *sNMF* and *SNAPCLUST* detected 93% and 100%, respectively. Simulated hybrids among pairs targeted for classification via *HYBRIDDETECTIVE* and *NEWHYBRIDS* showed 90–100% assignment accuracy across genotype frequency classes and species-pairs for posterior probability thresholds greater than or equal to 0.70.

(a) Hybrid detection

The number of species-pairs examined varied considerably among the six families/orders, with just one pair tested for two families versus 105 pairs within Leuciscidae (table 2). Hybrids ($N = 70$) were detected within four families (table 2), with hybrid proportions ranging from 0 to 4.4% of individuals. All but four hybrids were minnows (Leuciscidae), with exceptions being: (i) *Micropterus dolomieu* \times *salmoides* (Centrarchidae); (ii) *Etheostoma juliae* \times *zonale* (Percidae); (iii) *Etheostoma spectabile* \times *caeruleum* (Percidae); and (iv) a putative multi-specific hybrid: *Noturus maydeni* \times *albatere* \times *exilis* (Ictaluridae). Thus, we recognized $N = 18$ hybridizing species-pairs and $N = 8$ species-triplets, i.e. multi-species hybrids. We did not detect hybrids within Cottidae or Fundulidae.

Within Leuciscidae, we identified 66 hybrids and 15 hybridizing species-pairs (electronic supplementary material, S20). The most remarkable ($N = 29$ hybrids) involved *Camposotoma anomalum* \times *oligolepis*. All minnow species herein ($N = 15$) appear to hybridize with at least one other species in the study basin. The number of hybrids per species was significantly related to the number of community occurrences ($R^2 = 0.16$, $F = 6.1$, $p = 0.02$).

PCA, *sNMF* and *SNAPCLUST* were largely congruent in detecting hybrids (figures 1 and 2). Both *sNMF* and *SNAPCLUST* detected the same 36 hybrids, each separately identifying additional (26 and 8, respectively). *SNAPCLUST* demonstrated an exceptional ability to identify hybrids based on allele frequencies (100%); however, its discriminatory power was limited. As a result, the same individual was frequently identified as a hybrid in several pairwise tests, regardless of the second parental species involved. In all cases, these could be adjudicated using PCA and *sNMF*; however, low-level multi-specific introgression or that involving unsampled or 'ghost' lineages cannot be conclusively excluded. A list of hybrid individuals and inferences for each are available in electronic supplementary material, S21.

(b) Hybrid classification and introgression

We also found evidence of introgression based on the occurrence of backcrossed hybrid individuals. *NEWHYBRIDS* identified 39% of hybrids as backcrossed (24/62, excluding eight multi-specific hybrids); this corresponds to 55% (10/18) of all putatively hybridizing species-pairs showing evidence of introgressive hybridization (table 3). Approximately 24% of hybrids (15/62) were seemingly F_1 or F_2 . Finally, *NEWHYBRIDS* designated 37% (23/62) of putative

Table 1. Samples ($N = 33$ species) from the White River Basin, USA. N_I , number of individuals/species (post-filtering); N_S , number of collection sites/species (post-filtering); miss, mean proportion of missing data; depth, mean sequencing depth; H_o , observed heterozygosity; N_{FA} , number of individuals/family; SNPs, single nucleotide polymorphisms in family panel (post-filtering). Note: Several taxonomic changes have been adopted in the most recent *Fishes of Arkansas* [30] based on the literature but have yet to be recognized by the American Fisheries Society or Eschmeyers's Catalogue of Fishes.

family	common name	scientific name	N_I	N_S	miss	depth	H_o	N_{FA}	SNPs
Centrarchidae	bluegill sunfish	<i>Lepomis macrochirus</i>	69	23	0.15	46	0.028	375	1926
	longear sunfish	<i>Lepomis megalotis</i>	239	54	0.09	47	0.036		
	smallmouth bass	<i>Micropterus dolomieu</i>	43	30	0.47	39	0.011		
	largemouth bass	<i>Micropterus salmoides</i>	24	17	0.46	41	0.013		
Cottidae	banded sculpin	<i>Cottus carolinae</i> ^a	33	18	0.08	64	0.043	75	5344
	Ozark sculpin	<i>Cottus hypselurus</i> ^b	42	10	0.16	63	0.057		
Fundulidae	northern studfish	<i>Fundulus catenatus</i>	105	28	0.12	37	0.008	226	2366
	blackspotted topminnow	<i>Fundulus olivaceus</i>	121	34	0.20	33	0.024		
Ictaluridae	Ozark madtom	<i>Noturus albater</i>	10	5	0.12	65	0.033	31	2744
	slender madtom	<i>Noturus exilis</i>	16	12	0.09	51	0.042		
	Black River madtom	<i>Noturus maydeni</i>	5	3	0.28	34	0.034		
Leuciscidae	central stoneroller	<i>Campostoma anomalum</i> ^c	128	44	0.27	58	0.010	1507	343
	largescale stoneroller	<i>Campostoma oligolepis</i>	135	41	0.29	52	0.008		
	southern redbelly dace	<i>Chrosomus erythrogaster</i>	38	10	0.41	38	0.001		
	whitetail shiner	<i>Cyprinella galactura</i>	75	16	0.19	59	0.012		
	steelcolor shiner	<i>Cyprinella whipplei</i>	30	8	0.27	65	0.010		
	striped shiner	<i>Luxilus chrysocephalus</i>	63	18	0.18	59	0.013		
	duskystrip shiner	<i>Luxilus pilsbryi</i>	258	33	0.09	76	0.023		
	bleeding shiner	<i>Luxilus zonatus</i>	100	17	0.09	85	0.013		
	redfin shiner	<i>Lythrurus umbratilis</i>	24	5	0.26	47	0.004		
	bigeye shiner	<i>Notropis boops</i>	226	31	0.10	89	0.019		
	Ozark minnow	<i>Notropis nubilus</i>	193	35	0.11	76	0.016		
	carmine shiner	<i>Notropis percobromus</i>	67	15	0.20	66	0.010		
	telescope shiner	<i>Notropis telescopus</i>	83	15	0.14	75	0.006		
	bluntnose minnow	<i>Pimephales notatus</i>	55	24	0.25	73	0.007		
	creek chub	<i>Semotilus atromaculatus</i>	32	14	0.32	51	0.001		
Percidae	greenside darter	<i>Etheostoma blennioides</i>	62	26	0.24	55	0.006	651	687
	rainbow darter	<i>Etheostoma caeruleum</i>	348	53	0.09	54	0.026		
	fantail darter	<i>Etheostoma flabellare</i>	26	11	0.33	38	0.001		
	yoke darter	<i>Etheostoma juliae</i> ^d	63	15	0.24	58	0.011		
	orangethroat darter	<i>Etheostoma spectabile</i> ^e	50	10	0.24	48	0.023		
	current darter	<i>Etheostoma uniporum</i>	18	7	0.25	41	0.005		
	banded darter	<i>Etheostoma zonale</i>	84	26	0.25	51	0.018		

^a*Uranidea carolinae*.

^b*Uranidea immaculata* (knobfin sculpin).

^c*Campostoma plumbeum* (plains stoneroller).

^d*Nothonotus juliae*.

^e*Etheostoma* sp. cf. *spectabile* (Ozark darter).

hybrids as pure parentals, although possibly introgressed individuals poorly classified, i.e. late-generation hybrids [42].

(c) Correlates of hybridization

Genetic differentiation among species (table 4; electronic supplementary material, S22) was significantly lower for hybridizing versus non-hybridizing pairs (mean $F_{ST} = 0.86$

versus mean $F_{ST} = 0.93$; $Z = 2.87$; $p = 0.004$). The number of co-occurrences between hybridizing (mean = 8.4) and non-hybridizing pairs (mean = 7.1) did not differ significantly ($Z = -0.71$; $p = 0.49$).

At least one hybrid individual occurred within 42/75 communities (56%). We identified 12 environmental variables with a significant ability to reduce predictive error (i.e. VIMP) based on random permutation of predictors in the random

Table 2. Hybridization within six families sampled across the White River Basin, USA. Columns include: *N* species = number of species analysed; unique pairs = number of unique intra-familial; pairs not co-occurring = number of pairs within a family not found together at the same site; *N* indiv. = number of individuals examined; *N* hybrids = number of hybrids detected; per cent indiv. = hybrid percentage. Unique species-pairs with hybrids are only considered for ancestry between two species (i.e. species triplets were not counted).

family	<i>N</i> species	unique pairs	pairs not co-occurring	<i>N</i> indiv.	<i>N</i> hybrids	per cent indiv. (%)	species w/ hybrids	per cent species w/ hybrid (%)	unique pairs w/ hybrid	per cent pairs w/ hybrid (%)
Fundulidae	2	1	0	226	0	0.0	0	0	0	0
Cottidae	2	1	0	75	0	0.0	0	0	0	0
Ictaluridae	3	3	1	31	1	3.2	3	100	0	0
Centrarchidae	4	6	0	375	1	0.3	2	50	1	17
Percidae	7	21	5	651	2	0.3	4	57	2	10
Leuciscidae	15	105	7	1507	66	4.4	15	100	15	14
overall	33	137	13	2865	70	2.4	24	73	18	13

forest (electronic supplementary material, S23 and S24). From these, just four were used to maximize the predictive capacity of the final model based on minimal depth selection (figure 3), including species richness (VIMP = 0.09), protected area extent (pac_pc_use; VIMP = 0.04), mean precipitation in May (pre_mm_c05; VIMP = 0.03), and mean annual precipitation (pre_mm_cyr; VIMP = 0.03). The accuracy of the model was satisfactory: *G*-mean = 0.73 and normalized Brier score = 0.78. A similar misclassification error occurred for communities with and without hybrids (0.26 versus 0.27, respectively).

Neither species richness nor protected area extent were strongly correlated with other predictor variables ($r > 0.70$; Pearson correlation). The correlation coefficient between mean May and annual precipitation was $r = 0.61$. Both precipitation variables were strongly and positively correlated with other climatic variables related to temperature, evapotranspiration and soil water content. Annual precipitation was also strongly but negatively correlated with river network position (distance from network outlet; DIST_DN_KM). Although not selected for the final model, eight additional variables were significantly associated with hybridization based on VIMP, including indices of human impact ($N = 2$), road density ($N = 1$), annual snow cover ($N = 1$) and vegetative cover ($N = 4$) (electronic supplementary material, S23 and S24).

4. Discussion

Hybridization and introgression can impact fitness, facilitate gene exchange among species or generate new lineages [5,6,55–57]. While hybridization has been widely documented at the per-species level, its occurrence within communities is expected to be rare; otherwise, species boundaries would seriously deteriorate [2]. Despite these expectations, hybrid prevalence at the community level is largely unknown, given that most studies to date focus on single species-pairs (or but a few closely related pairs) [55].

This study quantified hybridization at the community level without *a priori* assumptions of putative hybridization, thus serving as an appropriate broad-scale model without species-specific biases. It provided a more holistic survey of hybridization across the riverscape and thus stood in contrast to more conventional studies based on single species-pair evaluations.

(a) Frequency of natural hybridization

The relevance of hybridization is underscored in our study by the detection of hybrids within four (of six) families (67%), involving 24 (of 33) species (73%). We documented hybridization among 18 unique species-pairs, 10 of which have been reported previously [20,58–62]. Hybridization across our study region was overwhelmingly within leuciscids, a ubiquitous family across Ozark stream communities [63]. The unbalanced number of representatives from each family reflects that and should not necessarily be interpreted as differences in the propensity to hybridize among the families more generally. Surprisingly, few hybrids were detected within Centrarchidae and Percidae, despite an extensively documented presence [19,64,65].

Hybridization was encountered more frequently than anticipated, whether viewed per-individual (2.4%) or per-species (73%). Previous estimates in plants and animals are

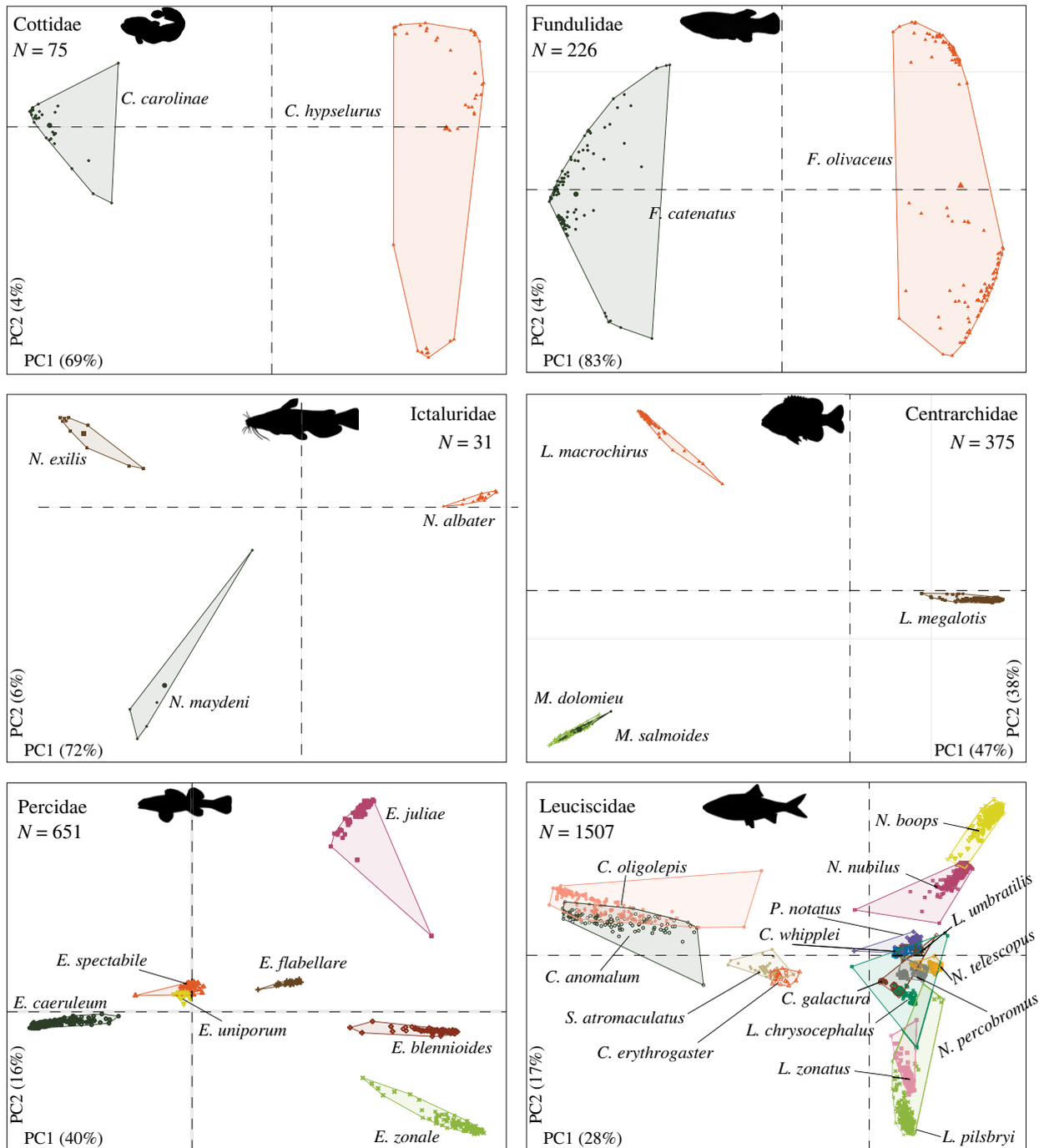


Figure 1. First two principal components derived from SNP genotypes in six fish families collected across the White River Basin, USA. The variance explained by each component is in the bottom left or right corner of each plot. *N*, numbers of individuals/family.

lower than found herein (0.002–0.06%) [15,66,67]. If rates among our study species were comparable to the previous (e.g. 0.1%), we would have expected but two–three hybrids. Earlier estimates from the literature relied on morphological identification, a less sensitive method, especially for detecting later-generation hybrids [19,62]. Additionally, rates herein also reflect both our breadth of individuals evaluated, as well as potential comparisons so tested. While our study is but a subsample of individuals from the region and lacks representatives from every extant species, we nevertheless attempted to reduce unsampled lineages that could bias inferences and allow undetected or misclassified hybrids [68,69].

Hybridization is seemingly greater in fishes than in other vertebrates due to their external fertilization within aqueous environments [1,19,70]. Similar per-individual rates (0–4%)

were identified in marine fishes [71], with even higher rates (22.5%) among invasive Mississippi River carp [72]. Published per-species estimates varied from 1 to 10% across animals and approximately 25% for plants [2,73]. Although differences in the former are apparent from literature and museum records, one could ask if it reflects an actual biological signal or variance in diversity and/or research effort. For example, once adjustments were made for diversity and research intensity, rates were mostly homogeneous among taxonomic groups compiled in a meta-analysis, but with the caveat that rates for fishes were still demonstrably higher than expected [18].

Our per-species rate was highest within our most specious group (Leuciscidae; table 2). This parallels previous meta-analyses identifying disproportionately high

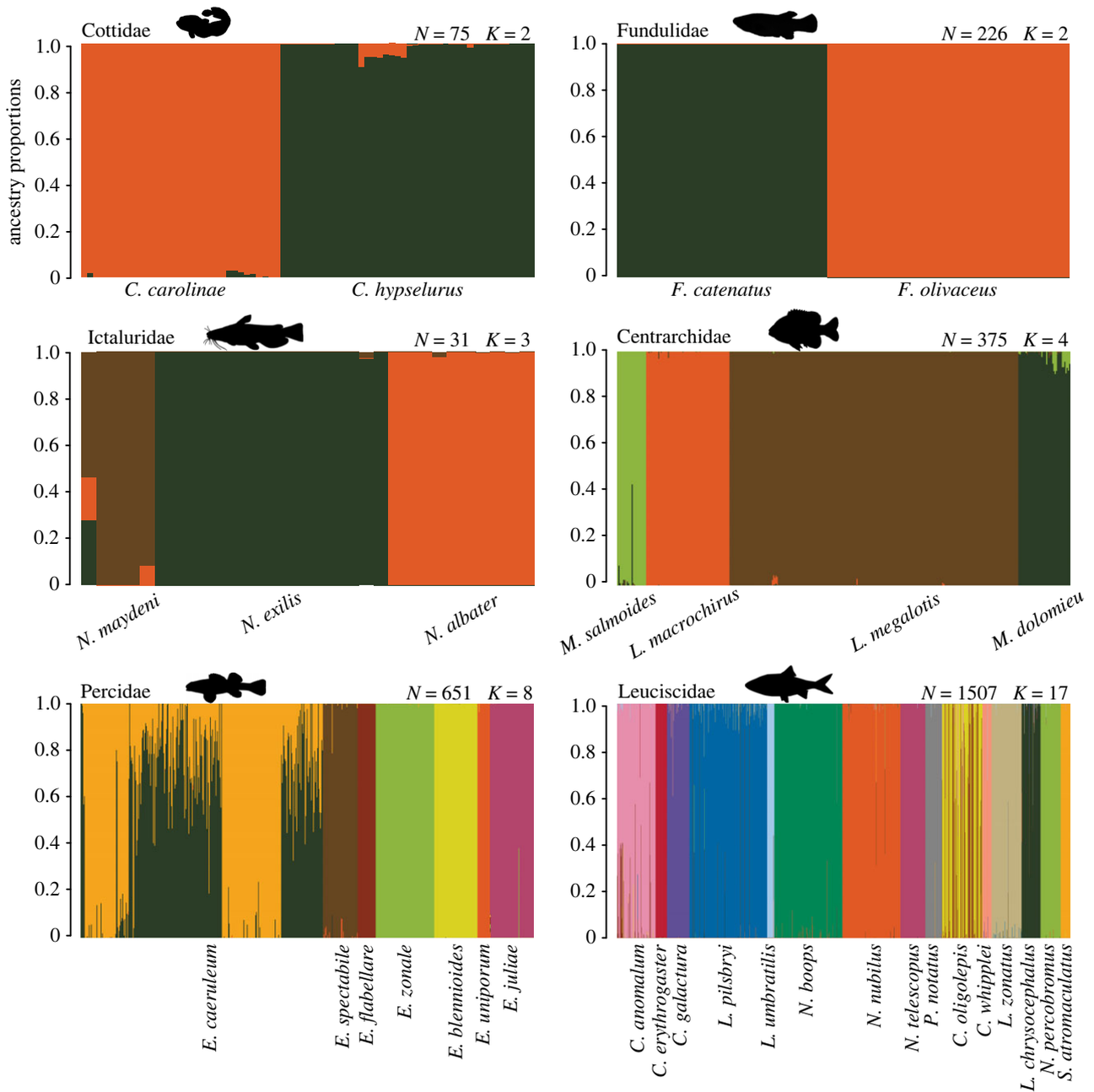


Figure 2. Calculated ancestral proportions for $N = 2865$ individuals from $N = 33$ fish species grouped by family collected from the White River Basin, USA. N , number of individuals, K , optimum number of clusters.

hybridization rates within minnows [1,19,20]. Interestingly, leuciscid breeding behaviours—especially nest building and association—significantly predicted hybridization rates across the clade [20]. Many minnows broadcast gametes widely or affix them onto substrates, often shared among species [20,74], with as many as six simultaneously employing the same gravel substrate at once [1]. In addition, minnows comprise the most diverse and widely distributed North American stream fish family and numerically dominate stream communities [75]. They thus encounter broad environmental heterogeneity, demonstrate a high degree of sympatry and exhibit uneven species abundances, which promote hybridization [19].

(b) Introgression

The evidence of backcrossing between hybrids and parentals suggests that hybrids can be viable and fertile, thus

facilitating introgression [67]. This genetic exchange can provide a source of novelty for evolutionary forces to act upon [73], and the prevalence found here provides some indication as to how probable an extraneous genetic contribution can be. We identified 24 individuals bred from parental/hybrid backcrosses spanning 10 species-pairs and two families (Leuciscidae, Percidae). It is intriguing that the most diverse group of North American freshwater fish—leuciscids—has established a reputation for frequent hybridization and is also a prolific contributor to gene exchange among species in our study. Introgressive hybridization has played a crucial role in cyprinid evolutionary history [67], and the evidence gathered here suggests it may continue to do so.

Questions remain regarding whether the introgression seen in our region is evolutionarily adaptive, maladaptive or neutral. Answers may ultimately hinge upon differences among species and environments [76], yet they would help predict and mitigate adverse outcomes of hybridization

Table 3. Observed genotypic frequency classes of hybrid individuals inferred from NewHybrids analysis for 18 hybridizing fish species-pairs collected across the White River basin, USA. Note that eight multi-specific hybrids were not included. Putative hybrids were assigned to a genotype frequency class (F_1 , F_2 , Bx [= backcross], pure) based on Bayesian posterior probability greater than 0.70.

no.	species A	species B	<i>N</i> indiv.	F_1	F_2	Bx	pure
1	<i>Campostoma anomalum</i>	<i>Campostoma oligolepis</i>	29	5	—	11	13
2	<i>Campostoma anomalum</i>	<i>Chrosomus erythrogaster</i>	1	1	—	0	—
3	<i>Campostoma anomalum</i>	<i>Luxilus pilsbryi</i>	1	—	1	0	—
4	<i>Campostoma oligolepis</i>	<i>Notropis telescopus</i>	1	1	—	0	—
5	<i>Cyprinella galactura</i>	<i>Cyprinella whipplei</i>	2	1	—	1	—
6	<i>Cyprinella whipplei</i>	<i>Lythrurus umbratilis</i>	1	—	—	1	—
7	<i>Luxilus chrysocephalus</i>	<i>Luxilus zonatus</i>	2	1	—	0	1
8	<i>Luxilus chrysocephalus</i>	<i>Semotilus atromaculatus</i>	1	—	—	1	—
9	<i>Luxilus pilsbryi</i>	<i>Lythrurus umbratilis</i>	6	—	1	5	—
10	<i>Luxilus pilsbryi</i>	<i>Notropis percobromus</i>	2	1	—	1	—
11	<i>Luxilus pilsbryi</i>	<i>Luxilus zonatus</i>	8	—	—	0	8
12	<i>Luxilus pilsbryi</i>	<i>Luxilus chrysocephalus</i>	1	—	—	1	—
13	<i>Luxilus zonatus</i>	<i>Pimephales notatus</i>	1	1	—	0	—
14	<i>Notropis boops</i>	<i>Notropis nubilus</i>	2	—	—	1	1
15	<i>Pimephales notatus</i>	<i>Semotilus atromaculatus</i>	1	—	1	0	—
16	<i>Micropterus salmoides</i>	<i>Micropterus dolomieu</i>	1	1	—	0	—
17	<i>Etheostoma spectabile</i>	<i>Etheostoma caeruleum</i>	1	—	—	1	—
18	<i>Etheostoma juliae</i>	<i>Etheostoma zonale</i>	1	—	—	1	—
Totals	—	—	62	12	3	24	23

Table 4. Mean, minimum and maximum values of Weir and Cockerham's pairwise F_{ST} calculated among species within families collected across the White River Basin, USA. Families Cottidae and Fundulidae were represented by only two species each, hence one value of F_{ST} .

family	mean F_{ST}	min F_{ST}	max F_{ST}
Centrarchidae	0.94	0.85	0.98
Cottidae	—	0.82	0.82
Fundulidae	—	0.98	0.98
Ictaluridae	0.81	0.66	0.91
Leuciscidae	0.92	0.65	0.99
Percidae	0.93	0.78	0.98

[77]. Climate-mediated environmental shifts will predictably exacerbate hybridization, especially for fishes whose life histories are sensitive to spawning temperatures and streamflow [12]. Yet we recognize that hybridization is not necessarily negative. For example, adaptive introgression between generalist and specialist rainbow fishes has seemingly diminished climate change vulnerability in admixed individuals compared with pure populations [10]. It is thus viewed as a valuable component of 'evolutionary rescue' and an underappreciated conservation tool [10].

(c) Correlates of hybridization

Our data concur with the premise that hybridization is strongly influenced by both divergences among species and

also environmental factors. We focus our discussion on the latter, per the novelty of our data and the previously established support for the former [20–22]. In synopsis, communities were more likely to harbour hybrids when: (i) greater species richness existed; (ii) protected area within the catchment was limited; and (iii) habitats were prone to more precipitation.

While hybridizing and non-hybridizing species-pairs did not differ significantly in their co-occurrence, we did note that communities where more species occur together (greater richness) also possessed more hybrids (figure 3). Although seemingly intuitive, the opposite is suggested at larger spatial and temporal scales where greater niche availability for hybrids might be expected in areas of lower species richness [16,23]. Empirical results are few, and conclusions varied: for example, the number of plant hybrids across US counties was significantly related to species richness [17], yet uncorrelated among coral reef fishes [14,16].

The extent of protected areas within the total watershed upstream of the reach was negatively associated with basin-level hybridization (figure 3). Similarly, anthropogenic impacts were positively associated with hybridization (electronic supplementary material, S24). Environmental perturbations in general, and specifically those anthropogenic, loom large in the hybridization literature [19,28]. Stable, more pristine environments are expected to harbour fewer hybrids than those perturbed, due mainly to a breakdown of reproductive isolating mechanisms facilitated by translocations, habitat modifications and ongoing climate change [28,78].

Greater mean precipitation within the reach catchment (annually and in May) was also associated with elevated

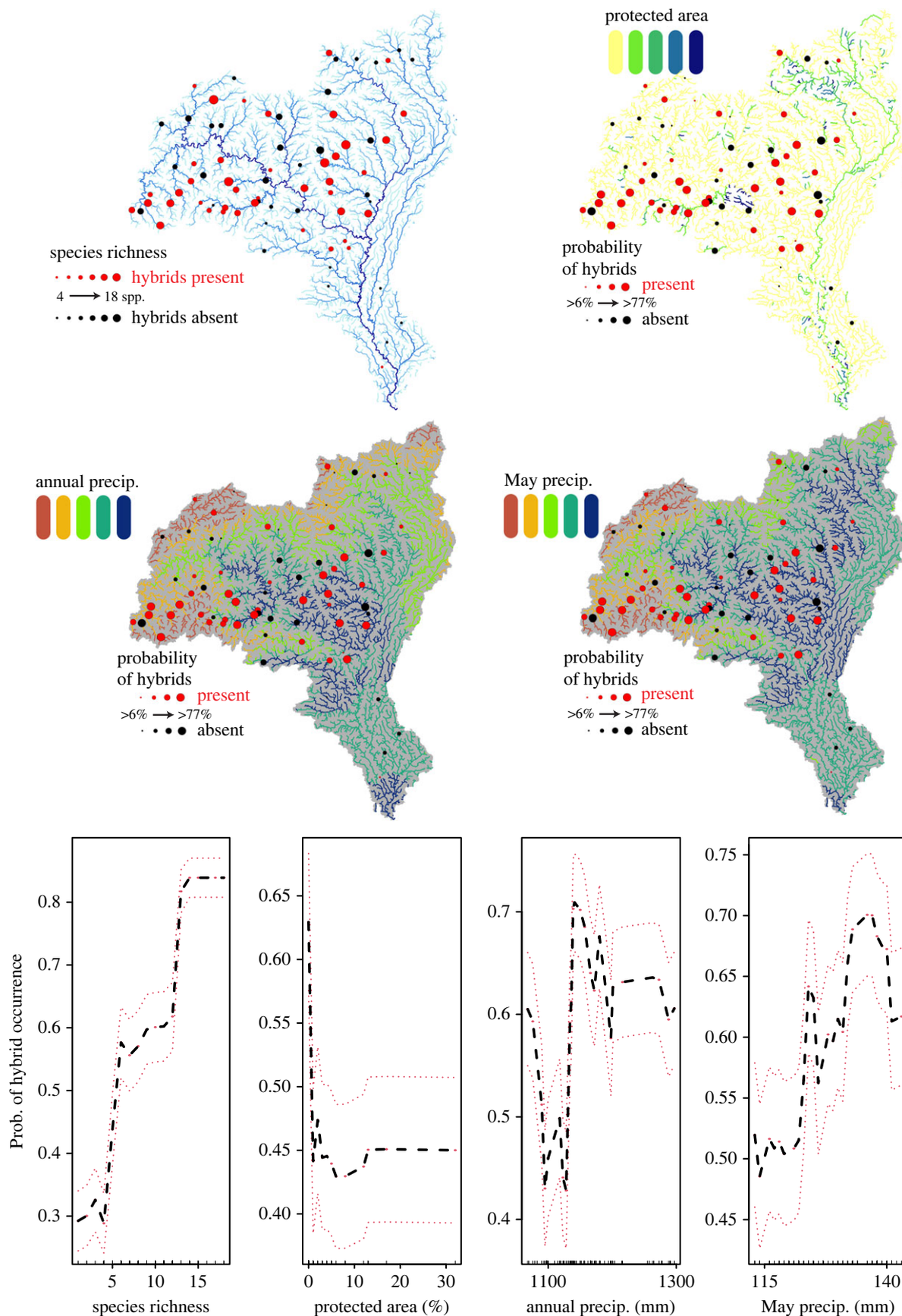


Figure 3. Significant predictors of hybridization across the White River Basin, USA. Maps depict collection sites with circles (red indicates the presence of a hybrid). In the first map, the site diameter is scaled by species richness, while in the remaining maps, it is scaled by the probability of hybrid occurrence based on the model. Basin backgrounds differ for enhanced visibility. Below the maps are partial marginal effects plots showing relationships between probability of hybrid individuals occurring per four predictive variables. The mean (black) and 95% confidence intervals (red) are shown, along with distribution of variable values across sites (black tick marks above x-axis).

hybridization (figure 3). More precipitation is associated with warmer and more downstream communities in our system (i.e. more flow). Elevated levels of hybridization in these

communities are potentially driven by flooding magnitude, particularly given the combination of higher precipitation and lower network position. Horton stream order and

catchment size promote the magnitude of flooding [79], as compounded by the precipitation gradient of our study system [80]. Additionally, the hydrologic disturbance index (a compendium of several anthropogenic impacts) is more significant in streams with larger drainage areas and lower gradients (i.e. further downstream), which in turn has been shown to promote variance in fish community composition [31].

Flooding can affect fish spawning in several ways: promotion of spawning activity [80]; concentration of fishes within refugia [53,81]; disturbance of spawning habitats and nests [82]; displacement of oviposited eggs [53]; and elevation of discharge/turbidity, which weakens those sensory cues (e.g. visual, olfactory, environmental) that sustain breeding isolation [25–27]. The inclusion of May precipitation in our most predictive models supports the above, in that study species (save two cottids) spawn in May/June [83].

5. Conclusion

Hybridization occurs more frequently than expected in the White River Basin and is predictable based on specifics of the environment. Although recognized as a creative evolutionary force, hybridization is also considered a maladaptive threat. Further research may blueprint an even more complex scenario and demonstrate that groups (such as minnows) thrive proportional to their prolific gene exchange. Moreover, hybridization is predicted to increase in frequency with global environmental change [11,12], a recognition consistent with our finding of hybrid occurrence in lockstep with climate-related variables. Therefore, baseline estimates are required to gauge the increase in hybridization, predict which ecosystems will be so impacted (and how severely), and promote a more robust conservation and

management strategy that allows those impacts to be understood and adjudicated (if so needed) [10]. Future studies like ours will be performed at the whole-genome level, with a resolution more robust for detecting and untangling hybridization and its genomic consequences [24].

Data accessibility. Raw sequence files are accessioned in the NCBI GenBank Sequence Read Archive (SRA) BioProject: PRJNA809538 [84]. SNP alignments and R code are archived on Open Science Framework [85].

The data are provided in electronic supplementary material [86].

Authors' contributions. Z.D.Z.: conceptualization, data curation, formal analysis, investigation, methodology, validation, visualization, writing—original draft, writing—review and editing; M.R.D.: funding acquisition, project administration, supervision, writing—review and editing; T.K.C.: data curation, methodology, resources, software, writing—review and editing; M.E.D.: funding acquisition, project administration, supervision, writing—review and editing.

All authors gave final approval for publication and agreed to be held accountable for the work performed therein.

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