Rapture-ready darters: sequence capture outperforms whole-genome data regardless of reference genome in *Etheostoma*

Brendan N. Reid¹*, Rachel L. Moran², Christopher J. Kopack³, Sarah W. Fitzpatrick¹,⁴

¹Kellogg Biological Station, Michigan State University, Hickory Corners, MI, USA
²Department of Evolution, Ecology, and Behavior, University of Minnesota, Saint Paul, MN, USA
³Department of Biology, Colorado State University, Fort Collins, CO, USA
⁴Department of Integrative Biology, Michigan State University, East Lansing, MI, USA

*Corresponding author: reidbre1@msu.edu
Abstract

Researchers studying non-model organisms have an increasing number of methods available for generating genomic data. However, the applicability of different methods across species, as well as the effect of reference genome choice on population genomic inference, are still difficult to predict in many cases. We evaluated the impact of data type (whole-genome vs. reduced representation) and reference genome choice on data quality and on population genomic and phylogenomic inference across several species of darters (subfamily Etheostomatinae), a highly diverse radiation of freshwater fish. We generated a high-quality reference genome and developed a hybrid RADseq/sequence capture (Rapture) protocol for the Arkansas darter (Etheostoma cragini). Rapture data from 1900 individuals spanning four darter species showed recovery of most loci across darter species at high depth and consistent estimates of heterozygosity regardless of reference genome choice. Loci with baits spanning both sides of the restriction enzyme cut site performed especially well across species. For low-coverage whole-genome data, choice of reference genome affected read depth and inferred heterozygosity. For similar amounts of sequence data, Rapture performed better at identifying fine-scale genetic structure compared to whole-genome sequencing and showed promise for detection of loci under selection. Rapture loci also recovered an accurate phylogeny for the study species and demonstrated high phylogenetic informativeness across the evolutionary history of the genus Etheostoma. Low cost and high cross-species effectiveness regardless of reference genome suggest that Rapture and similar sequence capture methods may be worthwhile choices for studies of diverse species radiations.

Keywords: phylogeography, species radiation, bait design, heterologous genome
Introduction

The advent of high-throughput sequencing technology has enabled biologists to generate genome-scale molecular data from a variety of organisms, creating new opportunities for conservation genetics (Shafer et al. 2015), phylogenetics (Lemmon and Lemmon 2013, McCormack et al. 2013), and molecular ecology (Ekblom & Gallindo 2011). As the capacity for high-throughput sequencing has increased, however, repositories of sequence data have become increasingly biased toward sequences from a minority of model organisms (David et al. 2019). Although non-model organisms represent fruitful study systems for answering basic questions in biology (Russell et al. 2017), deciding on appropriate methods for generating and handling genomic data for non-model species remains a challenge.

Whole-genome sequencing may still remain out of reach for large-scale studies of non-model organisms, and as such reduced-representation approaches have grown popular as effective means for answering many questions (da Fonseca et al. 2016, Meek and Larson 2019). Sequence capture or targeted sequence enrichment methods represent an attractive method for generating repeatable, high-coverage sequence data (Grover et al. 2012, Harvey et al. 2016). A hybrid method that uses restriction-associated DNA sequencing (RADseq) combined with targeted enrichment of a user-defined subset of hundreds to thousands of RAD loci, termed ‘Rapture’ (Ali et al. 2016) has great potential as a rapid and efficient method for generating repeatable high-throughput genomic data at low cost and high efficiency. Rapture assays have so far been developed and applied to salmon (Ali et al. 2016), Tasmanian devils (Margres et al. 2018), marine turtles (Komoroske et al. 2019), frogs (Peek et al. 2019), and sea lampreys (Sard et al. 2020). The application of Rapture has mainly focused on population genomics and adaptive
genomics within species, although Rapture loci developed for one species have been shown to be useful for studying hybridization among closely related species (Peek et al. 2019) and across species within slowly-evolving lineages (Komoroske et al. 2019).

For both whole-genome and reduced-representation sequencing, high-quality reference genomes can be used to improve genotype calling accuracy, inference of demographic history, and identification of loci under selection (Manel et al. 2015). For studies of non-model species, however, reference genomes may not be available for the particular species of interest. Assembling NGS data to heterologous genomes of related species is a potential option when such genomes are available. However, simulation studies indicate that even small divergences (0.15% to 2%) between the heterologous reference genomes and the native genome of the species of interest can increase errors in polymorphism calling and in estimates of genetic diversity, particularly when read depths are low (Nevado et al. 2014). Still, the practice of assembling short reads to a reference genome from a closely related species is common, and other empirical studies have concluded that congeneric or confamilial reference genomes may be suitable for SNP discovery (Galla et al. 2019).

Using a native reference genome in every situation is ideal but likely infeasible, especially when studying highly diverse species radiations. Applying high-throughput sequencing to the study of diverse species radiations will be particularly useful for understanding the effects of environmental context on genome evolution and identifying links between genetic variation and adaptive traits. Indeed, whole genome sequencing as well as reduced representation sequencing of adaptive radiations has uncovered signatures of change in genome structure and selection in
African cichlid fish (Brawand et al. 2014) and specific genetic loci associated with beak and body size variation in Darwin’s finches (Chaves et al. 2016). However, the cost of generating separate reference genomes for each species may be prohibitive, and making population genomic comparisons among species often necessitates assembling data to a single reference genome (as in Chaves et al. 2016). If using heterologous reference genomes is unavoidable in studies of diverse species radiations, it is important to quantify the biases that using these genomes will create when working with different types of data.

Darters (subfamily Etheostomatinae) represent a species radiation with great potential for illuminating the biotic and abiotic mechanisms that generate biological diversity. Darters are one of the most diverse clades of freshwater fish in North America, consisting of approximately 250 currently described species that likely shared a common ancestor between 30 and 40 million years ago (Near et al. 2011). Darters exhibit sexually dimorphic coloration that varies substantially among species, and sexual isolation based on divergent sexual selection has likely contributed to diversification in this group (Mendelson 2003, Moran et al. 2017, Moran and Fuller 2018a, Moran and Fuller 2018b). Postzygotic barriers between many sympatric species are not complete and hybridization is common, leading to gene tree discord and detectable signatures of ancient and contemporary introgression (Bossu & Near 2013; Moran et al 2017; Moran et al. 2018). Darters are also dispersal-limited and often restricted to small headwater streams, and as such allopatric diversification due to physical isolation also plays a large role in their diversification (Near and Benard 2004, Hollingsworth and Near 2009). In addition to driving diversification, physical isolation and micro-endemicity, as well as habitat degradation, have
created conservation issues for many darter species, and a substantial proportion of darter species diversity is currently considered threatened or endangered (Jelks et al. 2008).

High-throughput sequencing has great potential for providing insight into the forces controlling diversification in darters as well as for landscape and conservation genomics. Darter research to date has been characterized by a patchwork of molecular methods, making the comparison and integration of data from different studies difficult. Most previous phylogenetic work in darters has focused on Sanger sequencing of a small number of mitochondrial and nuclear genes (Near et al. 2011), while conservation genetics, landscape genetics, and molecular ecology studies have mainly used microsatellite markers developed for single species but with some applicability across the clade (Tonnis 2006, Khudamrongsawat et al. 2007, Switzer et al. 2008, Gabel et al. 2008, Hudman et al. 2008, Saarinen and Austin 2010). Recent work has begun to incorporate high-throughput sequencing methods, employing single-digest RADseq (Moran et al. 2018, MacGuigan et al. 2019, Moran et al. 2020) and double-digest RADseq (ddRAD, Moran et al. 2017, George 2018) to investigate phylogeny, phylogeography, and reproductive barriers among species. While ddRAD and RADseq represent a huge leap forward in terms of the amount of data generated, these methods often increase the number of loci genotyped at the expense of missing data and low coverage (MacGuigan et al. 2019). As such, there is currently no published method for reproducibly generating data for a single consistent set of loci distributed across the genome for darters. Furthermore, a reference genome assembly has only recently become available for a single darter species (the orangethroat darter *Etheostoma spectabile*; Moran et al. 2020).
Here, we describe an efficient and inexpensive Rapture-based method for reliably and repeatably genotyping thousands of loci in darters. This method is based on a capture bait set developed from RADseq data for Arkansas darters (*Etheostoma cragini*), a species of conservation concern found in the Arkansas River and nearby drainages within the Great Plains. Previous work in this species has used microsatellite markers to examine factors influencing population structure and genetic diversity in the western portion of their range (Fitzpatrick et al. 2014). The capture bait set targets over 2000 loci and includes both putatively neutrally-evolving loci as well as loci showing some evidence of selection across this species’ range. We assess two different tiling schemes for these baits, targeting either one or both flanking regions adjacent to a restriction cut site. We assess the performance of this capture bait set in a large set (*n* > 1600) of individual Arkansas darters as well as for individuals of three additional species in the genus *Etheostoma*. We assess the effects of aligning to either the *E. spectabile* or the *E. cragini* genome, and we also compare estimates of genetic diversity and population structure from Rapture to estimates from low-coverage whole-genome sequencing (WGS) data for a subset of *E. cragini* individuals. We ask the following questions to gauge the performance and applicability of the method: 1) How often are loci sequenced using the Rapture baits recovered at high coverage (>20x), and how many reads per individual are needed to attain high coverage?; 2) How much diversity is present within the set of Rapture loci for both the target species and for other darter species?; 3) Can the Rapture loci identify distinct population units within *E. cragini*?; and 4) Do the Rapture loci recover known phylogenetic relationships among and within species? We also demonstrate how the choice of data type (Rapture vs. WGS) and reference genome (heterologous vs native) affects inference of population genetic parameters and population structure.
Methods

Sampling

Dipnetting and electrofishing were used to collect 2,374 *E. cragini* individuals at 216 sites throughout Kansas in 2015-2016. Fin clips were taken from adults (>28mm) and whole specimens were collected for juveniles (<28mm). Samples were stored in 100% ethanol, shipped to Michigan State University (MSU), then stored in a freezer (-20°C) prior to analysis. In addition to the Kansas samples, whole *E. cragini* specimens were collected from six sites in Arkansas by the Arkansas Fish and Game Commission. Tissue samples and isolated DNA from Colorado populations were also available from a previous study (Fitzpatrick et al. 2014).

To examine the efficacy of Rapture across darter species, we also obtained genetic samples from three additional darter species: rainbow darters collected for a separate population genetic study in southwestern Michigan (Oliveira et al. 2020), *E. spectabile* specimens collected in the Salt Fork of the Vermillion River, Illinois, and fantail darter (*E. flabellare*) specimens collected in Fox Creek, Illinois.

DNA extractions

DNA from a pilot set of 52 *E. cragini* individuals sampled from seven sites was extracted using Qiagen DNeasy Blood & Tissue kits (Qiagen, Hilden, Germany). These extractions were done with a 60 ul elution in Qiagen EB buffer and quantified using a Qubit (Thermo Fisher Scientific, Waltham, MA, USA). For high-throughput extractions, we used a KingFisher Flex DNA extraction system (Thermo Fisher Scientific) to extract DNA from 20 sets of 90 samples (1800 samples total). We included an overnight digestion step in which tissues were lysed in a 96-well
PCR plate at a constant temperature of 55° C on an Eppendorf Mastercycler thermal cycler (Eppendorf, Hamburg, Germany), and we included 10 uL Proteinase K solution, 10 uL enhancer solution, 100 uL Qiagen Buffer EB solution, and approximately 10 mg tissue in each digest. We then used the MagMax whole blood protocol for an input volume of 200 uL and a final elution volume of 60 uL. We quantified DNA yield from high-throughput extractions using a PicoGreen assay (Thermo Fisher Scientific, Waltham, MA), with the six wells left unused in each plate used for assay standards and a negative control. High-throughput extractions included an additional 1635 E. cragini samples from Kansas, 60 E. cragini samples from Arkansas, 20 E. cragini samples from Colorado, and all E. spectabile and E. flabellare samples. DNA extracted for this study from E. cragini covered a total of 232 collection sites (n = 2-10 per site). Nearly all of these samples yielded high-quality DNA and were included in the Rapture genotyping analyses described below.

Pilot RADseq library preparation & Illumina sequencing

Using the pilot set of 52 E. cragini samples, two initial RADseq libraries were prepared and submitted to the MSU core genomics facility for sequencing. We used the ‘BestRad’ protocol following Ali et al. 2016. Briefly, genomic DNA (100 ng) from each sample was digested with a restriction enzyme (Sbfl-HF) and indexed with a biotinylated RAD adapter. Pooled DNA was sheared to 500 bp fragments using a Covaris sonicator (Covaris, Woburn, MA, USA). Shearing efficiency was evaluated with a fragment analyzer. Dynabeads M-280 streptavidin magnetic beads (Thermo Fisher Scientific, Waltham MA, USA) were used to physically isolate the RAD-tagged DNA fragments. The DNA was then eluted in TE buffer and used in NEBNext Ultra DNA Library Prep Kit for Illumina (New England Biosciences, Ipswich, MA, USA) with no
modifications. The two libraries were each sequenced with paired-end 150 bp reads on an Illumina HiSeq 4000 in separate lanes.

Bioinformatic pipeline for pilot dataset

As the BestRad protocol can result in sequences with barcodes on either the forward or reverse reads, we used a Python script (Flip2BeRad, https://github.com/tylerhether/Flip2BeRAD) to flip sequences with barcodes on the reverse read. We then filtered out potential PCR clones and demultiplexed sequences using the clonefilter and process_radtags commands in Stacks v. 2.4 (Catchen et al. 2013; Rochette et al. 2019). Using the demultiplexed forward reads, we identified loci containing single-nucleotide polymorphisms (SNPs) in ipyrad (Eaton and Overcast 2020). Reads were filtered using ipyrad’s default quality thresholds and mapped to an early draft version of the genome for the orangethroat darter (E. spectabile). We retained an initial set of candidate loci that were genotyped in \( \geq 75\% \) of the pilot set of 52 E. cragini individuals and that contained SNPs with a minor allele frequency (MAF) > 0.05. Additionally, we only retained loci with SNPs that were called in at least two E. cragini individuals, which imposed an additional floor on MAF and removed SNPs called in only one individual due to sequencing error. We created a FASTA file for all loci that passed these allele frequency filters and aligned these sequences to the draft E. spectabile genome using bwa (Li and Durbin 2009). As bait capture is optimally efficient as long as sequences are \( >95\% \) similar to baits (Arbor Biosciences, personal communication), any sequences that exhibited \( <95\% \) similarity or aligned to multiple locations on the E. spectabile draft genome were removed from the candidate set. Because the draft genome contained many small scaffolds, we also removed any loci that were located on scaffolds.
smaller than 10kb, as it would be difficult to determine whether these loci were adjacent to any
other loci in the final chromosome-level genome assembly.

To identify potential signatures of selection in this initial set of candidate loci, we used the
program BayeScan (Foll and Gaggiotti 2008) to identify outlier loci with higher or lower $F_{ST}$
values than expected by chance. We conducted an initial analysis using all populations. After this
analysis showed a high average $F_{ST}$ and an overabundance of lower-than-expected $F_{ST}$ outliers,
we removed individuals from populations in the eastern/Ozark portion of the $E. cragonii$ range
that exhibited strong divergence from darters in the rest of the range and re-ran the analysis. We
used a false discovery rate of 0.05 to identify outlier SNPs.

Bait design

From the candidate set of RAD loci, we identified three different categories of potential baits to
be used as targets for Rapture: (1) short loci (n=3176), in which ipyrad identified a locus
containing at least one SNP that was located on one side of the restriction cut site only; (2) long
loci (n=249), consisting of paired loci that both contained a SNP and were located on either side
of the cut site; and (3) outlier loci (n=29) identified by Bayescan. Long loci were initially chosen
to assess stretches of homozygosity or as potentially more phylogenetically informative blocks of
sequence. We obtained BED coordinates for all target loci on the $E. spectabile$ draft genome and
provided these coordinates and the draft genome to Arbor Biosciences (Ann Arbor, MI, USA).
Arbor Biosciences designed and produced a set of 4,966 80-bp baits to capture all long and
outlier target RAD loci, as well as 1841 of the short target RAD loci. While most previous
Rapture study designs have used 120-bp baits (Ali et a. 2016, Komorske et al. 2019, Peek et al.
2019), we used 80-bp baits tiled in an overlapping manner along the target loci to increase capture efficiency (as in Sard 2020). For short and outlier loci, two baits were tiled along each locus (starting at the restriction site), meaning that approximately 40 base pairs in the center of each locus were covered by two baits and the regions flanking this central region were only covered by one bait. For the long loci, five baits were tiled across both regions flanking the restriction site (Figure 1), meaning that a much longer region (approximately 160 bp) was covered by more than one bait.

### E. cragini whole genomes

To compare population genetic statistics generated with Rapture to those generated using WGS, we produced a reference genome for *E. cragini* and conducted low-coverage whole-genome resequencing. We submitted *E. cragini* muscle tissue from young-of-year fish of unknown sex raised at the John W. Mumma Native Aquatic Species Restoration Facility to Dovetail Genomics (Scotts Valley, CA, USA) to produce a high-quality reference genome for this species. Dovetail performed Illumina shotgun library preparation, paired-end 2x150 sequencing on an Illumina HiSeq X, and *de novo* assembly in Meraculous (Chapman et al. 2011) using a kmer size of 79. The assembly was refined using Chicago and Hi-C libraries, and scaffolds were constructed using HiRise (Putnam et al. 2016).

We submitted samples from 24 *E. cragini* samples for low-coverage whole-genome resequencing at the MSU Core Genomics center. Samples were chosen to include several individuals in each of several population clusters identified by Rapture (see below). We used the Illumina Coverage calculator...
(https://support.illumina.com/downloads/sequencing_coverage_calculator.html) to estimate the amount of sequencing needed to achieve \( \geq 5x \) coverage based on genome size and an estimate of 20% duplicate sequences. These samples were submitted in two batches of 12, and each batch also contained four samples from another fish with a similar genome size (\textit{Gambusia affinis}). As such, we used 75% of a lane of sequencing for each batch of 12 samples (1/16\textsuperscript{th} of a lane for each sample). Due to maintenance problems at MSU, the initial batch of sequencing produced fewer reads than expected. MSU sent the first batch of samples to the University of Michigan genomics core for additional sequencing and sent the second batch of samples to the Illumina FastTrack Sequencing Service Center for sequencing. All sequencing was performed on an Illumina HiSeq 4000.

We used FastQC (http://www.bioinformatics.babraham.ac.uk/projects/fastqc/) to assess sequencing quality for individuals. We used BWA v. 0.7.17-r1188 (Li and Durbin 2009) to align sequences to either the native \textit{E. cragini} genome or the heterologous \textit{E. spectabile} genome. We used samtools v.1.9 (Li et al. 2009) to filter out low-quality sequences and improperly paired reads, remove duplicates, and compute average coverage over the whole genome and over all covered sites for alignment to either the native or heterologous genomes.

\textit{Rapture library preparation, sequencing, data processing pipeline, and quality control}

We used the BestRAD protocol described above along with a sequence capture step that incorporated the Rapture bait sequences to conduct reduced-representation library preparation for 1900 individuals (1855 \textit{E. cragini}, 28 \textit{E. caeruleum}, 8 \textit{E. flabellare} and 9 \textit{E. spectabile}). We aimed for a target DNA mass of 200 ng in 10 uL for the starting material in each reaction. For
DNA samples with concentrations of 15-20ng/uL of DNA, we used 10uL total DNA. For samples with concentrations <15 ng/uL, we used a ThermoSavant DNA120 Speedvac (Thermo Fisher, Waltham, MA, USA) to dry down a sample volume containing 200ng and then resuspended in 10 uL 1x TE buffer. We performed library preparation in batches of four 96-well plates (containing 95 samples and one 1X TE blank), using the BestRAD barcode sequences and a plate-specific Illumina adapter for each plate. After BestRAD library preparation, we pooled all four plates and performed sequence capture using the protocol provided by Arbor Biosciences. Briefly, this involved performing a hybridization step at 65°C for at least 16 hours, isolating bait-target hybrids using streptavidin-coated magnetic beads and washing to remove non-target DNA, and performing PCR amplification of captured DNA for sequencing. We submitted these libraries for sequencing at the MSU Genomics Core facility in five batches of 380 samples each using paired-end 2x150 bp reads on an Illumina HiSeq 4000, using a single lane of sequencing for each batch. We used the steps described above for BestRAD to process the raw data, and we used BWA to align reads to both the E. spectabile (final version) and the E. cragini reference genomes, and used samtools v. 1.7 (Li et al. 2009) to remove improperly paired reads. We generated two updated bed files by aligning the baits to each genome and merging all loci together into a single file. We estimated individual-level coverage of Rapture loci using samtools. We assessed two metrics of coverage: (1) the number of reads with any overlap for each locus; and (2) per-base coverage of each locus for a subset of individuals. For the latter metric, we examined a buffered region (+500 bp from the 3’ end of the baits for short and outlier loci, +/- 500 bp on either side of the baits for long loci ) to see whether we achieved significant coverage from reverse reads. We filtered all BAM files using these buffered regions before performing population genetic and phylogenetic analyses.
Population genomics, population structure, and selection

We used ANGSD v.0.928 (Korneliussen et al. 2014) to calculate genotype likelihoods for single-nucleotide polymorphisms (SNPs) based on aligned BAM files for Rapture and low-coverage WGS data. As we only had WGS data from *E. cragini*, we excluded the other species from this analysis. To create subsetted datasets with comparable numbers of total reads for comparing population genetic inferences between Rapture and WGS, we randomly selected 570 individuals (corresponding to the number of individuals sequenced on 1.5 lanes) from the Rapture dataset. We retained all sites for which ANGSD detected a SNP with a *p*-value of < 1x 10^-6. We then used the program PCAngsd v.0.981 (Meisner and Albrechtsen 2018) to conduct downstream population genomic analyses. We first conducted principal component analyses (PCA) and calculated genotype probabilities on each of six datasets (WGS, full Rapture, and subsetted Rapture aligned to either the *E. spectabile* or *E. cragini* genome), with the optimum number of principal components determined by PCAngsd using a minimum average partial (MAP) test. To obtain estimates of heterozygosity, we used PCAngsd to call genotypes using a probability threshold of 0.9. We compared matched individual heterozygosity values between data types (WGS or Rapture) and between data aligned to either the *E. spectabile* or the *E. cragini* reference genome. We also used PCAngsd to estimate individual admixture proportions for each individual and to perform a PCA-based scans for loci potentially under selection (i.e. loci exhibiting greater differentiation along PCs than expected by drift; Galinsky et al. 2016). We calculated *p*-values for the test statistics generated by PCAngsd selection scan using a one-tailed chi-squared test with one degree of freedom. To account for the large number of tests conducted for the selection scan, we set a conservative significance threshold for each dataset (Rapture and WGS) of 0.05.
divided by the number of SNPs in the dataset. We compared within-species population structure and selection scan results along the first principal component axis between data types and reference genomes for all *E. cragini* individuals.

**Phylogenetics and phylogenetic informativeness**

We compiled filtered BAM files aligned to the *E. cragini* genome from a subset 56 individuals (two individuals from *E. spectabile* and *E. flabellare*, two individuals from each of two sites for *E. caeruleum*, and two individuals from each of 19 sites covering the full distribution of *E. cragini*) and ran the ref_map.pl script in Stacks using the “populations: phylip_var_all” option and default parameter values to call SNPs and output PHYLIP-formatted concatenated multiple sequence alignments for each individual. We then used IQTREE (Nguyen et al. 2015) to construct a phylogenetic tree of all sequences. We used the default maximum likelihood model selection and tree search methods in IQTREE with 1000 bootstraps to calculate support values.

We converted this tree into a time-calibrated ultrametric tree using the R package ape (Paradis and Schliep 2019). We set estimated branching times for three splits based on a published study of darter evolution (Kelly et al. 2012) using the makeChronosCalib function to calibrate ranges of potential branching times for three interspecific splits. We set the root of the tree, identified here as the common ancestor of the clades *Oligocephalus, Psychromaster,* and *Catonotus,* to 24-34 million years ago. We also set the root of *Oligocephalus* (corresponding to the *E. caeruleum – E. spectabile* split in our tree) to 17.5-27.5 million years ago, and the common ancestor of *Psychromaster* and *Catonotus* (corresponding the *E. flabellare – E. cragini* split in our tree) to 16.5 – 26.5 million years ago. We then used the function chronos to construct a time-calibrated
tree under three clock models (correlated, relaxed, and discrete). The correlated model had the highest likelihood and we used the tree calibrated using this model in all further analyses. We plotted the time-calibrated tree in ape and plotted the tips of the tree in space using the R package phytools (Revell 2012).

To calculate phylogenetic informativeness, we created separate PHYLIP files for each set of loci (short, long, and outlier) and concatenated all three into a single dataset and exported as a Nexus file using ape. We then input this alignment and the time-calibrated tree into the PhyDesign web interface (López-Giráldez and Townsend 2011). We examined the inferred net phylogenetic informativeness for each set of baits over the time period covered by the phylogeny (30 million years ago – present).

**Results**

*E. cragini whole genomes*

The *E. cragini* genome was similar in terms of contiguity and completeness compared to other published percid reference genomes, although it was smaller, contained less repetitive content, and exhibited a number of chromosomal rearrangements, especially relative to *E. spectabile* (Supporting Information 1). Coverage for resequenced *E. cragini* individuals varied based on the number of reads generated and the reference genome used. Shotgun sequencing for low-coverage WGS generated between 20.5 – 37.8 million read pairs per individual. Between 8.1%-15.5% of sequences were duplicates. Average read depth for covered sites (i.e. all sites with at least 1x coverage) and for all sites in each genome increased with the number of reads (Figure 2). Average read depth and depth of covered sites were highest when reads were aligned to the *E.*
cragini genome and were almost identical, indicating that nearly all sites in the E. cragini genome assembly were covered at least 1x. Read depth progressively decreased by approximately 20% for covered sites and by approximately 44% for all sites when reads were aligned to the E. spectabile genome (Figure 2). While some of the decline in coverage over all sites may be attributable to the 30% greater length of the E. spectabile assembly (which is suggestive of a reduction in genome size for E. cragini relative to E. spectabile), lower coverage at sites with at least 1x coverage (presumably present in both genomes) also suggests loss of sequencing information resulting from poor alignment to the heterologous reference genome.

Rapture quality control and coverage across species

Most of the samples (96%) sequenced using Rapture generated >10,000 read pairs, and 93.7% generated >100,000 read pairs. For E. cragini samples mapped to the E. cragini genome, an average of 39.7% of read pairs following bait capture (range 25.8% - 55.3%) overlapped the Rapture loci. A similar proportion of the E. caeruleum reads overlapped the rapture loci (average 43.1%, range 39.4% - 47.3%), while a lower proportion of reads from E. flabellare (26.0% average, range 24.4% - 28.2%) and E. spectabile (29.0% average, 27.6% - 30.8%) overlapped the Rapture loci. Statistics were similar for reads mapped to the E. spectabile genome. The average proportion of E. cragini read pairs overlapping the Rapture loci was 37.8% (range 26.3% - 52.9%). A similar proportion of the E. caeruleum reads overlapped the rapture loci (average 42.8%, range 39.3% - 46.9%), while a lower proportion of reads from E. flabellare (25.6% average, range 24.0% - 27.8%) and E. spectabile (30.0% average, range 28.7% - 31.5%) overlapped the Rapture loci. As
alignments were filtered to include only the Rapture loci with a 500bp buffer added, all results below use only the reads overlapping these loci.

The number of Rapture loci covered increased with number of reads for a given individual and tended to reach an asymptote above 10,000-100,000 reads (Figure 3, Supporting Figure 2). The maximum number of loci covered varied between species and between types of loci. For *E. cragini*, nearly all of the 2119 Rapture loci were covered at each read depth. For the other species, a maximum of 1700-1800 of the Rapture loci were covered (Figure 3). The reduction in covered loci mainly came from a loss of short loci, of which only ~1500 of 1841 (~80%) were covered. A higher proportion of long loci (88%-95%) were sequenced at high coverage, and almost all of the outlier loci were sequenced at high coverage as well (Figure 3). Coverage for Rapture loci was nearly identical when the heterologous *E. spectabile* reference genome was used for alignment (Supporting Figure 2). Per-base read depth was high for the portion of each locus covered by the capture baits for both long and short loci, representing large numbers of forward reads starting from the cut site overlapping the same region, although short loci had lower read depth beyond the capture baits compared to long loci (Supporting Figure 3).

**Polymorphism and heterozygosity**

For the Rapture dataset, there were ~16,401 SNPs (for the alignment to *E. cragini*) and ~24,903 SNPs (for the alignment to *E. spectabile*), indicating the presence of multiple SNPs per locus. For the WGS dataset, there were 5,774,040 SNPs for the alignment to *E. cragini* and 16,003,116 SNPs for the alignment to *E. spectabile*. Individual SNP heterozygosities were highly correlated across datasets – however, estimated heterozygosities were higher for the WGS datasets, and
heterozygosity was higher for the *E. spectabile* WGS dataset than the *E. cragini* WGS dataset. (Figure 6).

**Population structure and selection**

We compared the results of population structure and selection in analyses for *E. cragini* among different datasets (full and subsetted Rapture datasets versus WGS) to evaluate how data type and reference genome affected downstream population genetic inferences. The admixture analysis in PCAngsd indicated that the best population delineation included 16 different populations for the full Rapture dataset aligned to either reference genome (Figure 5a and 5b), and 14 and 11 populations for the subsetted Rapture dataset aligned to the native and heterologous reference genomes (Supporting Figure 4), respectively. For WGS data, however, PCAngsd found 3 populations for the data aligned to the *E. cragini* reference, and 2 populations for the data aligned to the *E. spectabile* reference (Figure 5c and 5d). The populations resolved for Rapture datasets broadly corresponded to major river drainages. The populations resolved for WGS lumped together populations in the major northern and southern drainages.

Selection scans in PCAngsd using the full Rapture dataset aligned to the native reference genome identified 66 SNPs showing significant evidence for selection along the first principal component after correction for multiple tests, although no loci crossed the significance threshold in analyses of the subsetted data (Figure 6, Supporting Figure 5). Interestingly, none of the SNPs in the outlier loci that initially showed evidence of selection in the pilot dataset were identified as significant in these scans after correction for multiple testing, although several exhibited p-values < 1x10^-4. The selection scan of Rapture data aligned to the heterologous genome identified many
more loci with p-values lower than the significance threshold (Figure 6, Supporting Figure 5). For the WGS datasets, PCAngsd did not identify any loci with significant evidence for selection when aligned to either reference genome after correction for the larger number of tests (Figure 6).

Phylogeny

Maximum likelihood phylogenetic analyses indicated that the Rapture loci were capable of resolving phylogenetic relationships with fairly strong support. The ML analysis produced 100% bootstrap support for correctly grouping *E. spectabile* with *E. caeruleum* and for grouping *E. flabellare* with *E. cragini*, as well as for grouping all individuals within their respective species (Figure 7a). Several deep phylogenetic splits (approximately 2.5-6 million years old) within *E. cragini* also received high support, and individuals within sites and within drainages were often grouped together with high support. Within *E. cragini*, populations showed a nested phylogeographic structure, with Arkansas populations basal to all populations to the east, and populations in east Kansas basal to populations further west. There was also a strongly supported split between populations in the mainstem Arkansas River and its tributaries and populations in drainages to the south of the mainstem Arkansas River (Figure 7b).

Per-site phylogenetic informativeness profiles for the three categories of loci showed similar overall patterns from 30 million years ago to 2-3 million years ago, with a slightly convex but relatively stable informativeness profile over time (Supporting Figure 6). Informativeness dropped rapidly from 2 million years ago to the present for the long and short loci, but outlier loci exhibited a secondary peak from 1-2 million years ago for the outlier loci followed by a
steep decline. Long loci tended to have lower per-site phylogenetic informativeness than short or outlier loci.

**Discussion**

There are a number of common questions any researcher involved in the design and implementation of a population genomic or phylogenomic study in a non-model organism will have to address. These include: how many loci and how many individuals do I need to include? Should I sequence loci over the entire genome or should I use sequence capture to target a smaller number of loci at high depth? Should I generate a reference genome for my species or will I be able to use a reference genome from a closely related species, and how will this choice affect the interpretation of my data? Will one methodology work equally well across all target populations and species? And how cost-effective are these alternative methods? All of these questions are perhaps even more relevant for projects aimed at diverse species radiations, as such projects by their nature encompass a number of closely related species. Based on the work described here, we discuss how these questions can be addressed and which methods are most appropriate for different applications.

**To Rapture or not to Rapture (and how to Rapture)**

A number of sequence capture methods exist, ranging from anchored probes (Lemmon et al. 2012) and ultraconserved elements (Faircloth et al. 2012) developed for use across a wide variety of taxa, to more focused methods that develop and use a bait set for a single species (Margres et
Previous work with the Rapture method in marine turtles demonstrated that baits developed for a single species work well in related species that diverged tens of millions of years ago (Komoroske et al. 2019), and we confirm in this work that Rapture loci developed for a single darter species can also be used in other species from the same group. Rapture loci were recovered with highest coverage from the target species (E. cragini) but a majority of loci were recovered from all four species. Long loci spanning both sides of the restriction site were recovered with higher frequency than loci that did not span the restriction site (short loci) across species, and we obtained higher coverage in regions flanking the RAD locus for long loci as well. This is possibly because of a greater possibility of bait capture for more dissimilar sequences with more baits per locus (5 baits for long loci compared to 2 baits for short loci). We also used 80-bp tiled baits as opposed to 120-bp baits used in previous Rapture studies, which may have improved the likelihood of capture as well. These results suggest overall that using loci that span both sides of the restriction site and using 80-bp tiled baits will likely lead to the most consistent recovery of Rapture loci across related species. Incorporating multiple reference genomes or creating pseudo-reference genomes from pilot RAD data for other species of interest may be useful in designing bait sets that will function best across species radiations.

Native vs heterologous reference genomes

Although mapping sequence reads generated from one species to the genome of a closely related species is still common practice, the effects of mapping reads to a heterologous genome versus a native genome are still relatively understudied. Galla et al. (2019) mapped RADseq and low-coverage WGS data to either a conspecific, congeneneric, confamilial, or conordinal genome and
found a decreasing alignment rate with increasing phylogenetic distance, as well as less consistency in estimates of genetic diversity when reads were mapped to a more distantly-related genome. Our WGS results generally agree with these findings. Mapping reads generated from low-coverage WGS of *E. cragini* individuals to the *E. cragini* reference genome was generally more successful than mapping to the *E. spectabile* genome. Lower read depth and allelic dropout could contribute to different estimates of genetic diversity from native and heterologous reference genomes. However, even though we inferred population structure using genotype likelihoods, which should mitigate the effects of lower read depth associated with mapping to a heterologous reference genome (Nevado et al. 2014), admixture results were also affected by the choice of reference genome. The existence of multiple rearrangements among darter genomes observed in this study and others (Moran et al. 2020) likely aggravates the effects of using a heterologous reference genome. For our Rapture dataset, however, the effects of mapping to a heterologous reference were much reduced, and downstream inferences regarding diversity and population structure were similar, regardless of which reference genome was used for mapping. This suggests that sequence capture may reduce biases associated with the absence of a closely-related reference genome, possibly because the RAD loci targeted by sequence capture in this case were fairly conserved across genomes. However, selection scans using Rapture data were still impacted by choice of reference genome. The larger number of loci detected with very low p-values when using the *E. spectabile* reference suggests that using a heterologous reference genome may introduce biases not present when using the native reference genome, and further work with known loci under selection would be useful to determine whether the use of a heterologous reference genome increases false discovery rates.
Different data types can be differentially suited to different analyses. We found that Rapture was much better at identifying fine-scale population structure than WGS. This is likely partially due to the much greater spatial coverage and the greater number of individuals we were able to sequence via this method. Higher coverage overall for the Rapture data may also alleviate allelic dropout and decreased sensitivity for calling heterozygotes associated with low-coverage WGS, although as PCAngsd uses genotype likelihoods rather than called genotypes this issue may not have strongly affected these analyses. While previous work has asserted that WGS is typically better suited to detecting evidence of selection using genome scan methods than RAD-based approaches which target relatively small portions of the genome (Lowry et al. 2016), we found weaker signals of selection in our WGS data compared to Rapture. This may be due again to the lower number of individuals sampled for WGS, and less accurate estimation of population structure may also have confounded detection of loci under selection using WGS. RADseq-based methods can detect selection if marker density is high relative to the size of linkage disequilibrium blocks (Catchen et al. 2017), and Rapture workflows designed to detect selection with these factors in mind may be comparable to WGS. Alternatively, Rapture methods can also include loci with known \textit{a priori} effects on fitness (such as loci associated with disease susceptibility). While our Rapture loci identified \textit{a priori} as under selection by genome scans in the pilot dataset did not show strong evidence of selection in the larger datasets, Rapture panels designed to include high marker density as well as immune-associated loci constituted an effective means identifying loci associated with survival in female Tasmanian devils \textit{(Sarcophilus harrisii)} with a transmissible cancer (Margres et al. 2018).
Phylogenetic informativeness of Rapture loci

Sequence capture strategies targeting ultra-conserved elements (UCEs) and protein-coding genes have been evaluated in the past for percomorph fishes (Gilbert et al. 2015). UCE flanks and protein-coding genes in general showed great utility for resolving deeper split but a loss of phylogenetic signal for more recent epochs, with per-locus phylogenetic informativeness for UCE flanks and protein-coding genes peaking between 20-40 million years ago and exhibiting rapid decline from 20 million years ago to the present. The phylogenetic informativeness of Rapture loci was fairly constant over time and potentially more useful for examining relatively recent splits between closely related species. However, phylogenetic informativeness for Rapture declined rapidly for very recent epochs. This is also potentially reflected in support values estimated here for relationships within the *E. cragini*. We obtained 100% bootstrap support for older splits between populations in Arkansas versus populations further east, as well as high support for a sister relationship between populations in eastern Kansas and all other populations to the west and a split dating to approximately 3 million years ago between populations in drainages associated with the mainstem Arkansas river and populations in drainages to the south of the Arkansas River. For more recent splits, support values were overall fairly high (95-100%) but much lower for some nodes, indicating ambiguous support for some relationships. This likely represents both a true lack of phylogenetic informativeness (i.e. substation rates too low to allow for reliably distinguishing among alternative relationships) as well as potentially other confounding factors, such as gene flow and maintenance of ancestral polymorphisms.
Alternative methods that incorporate gene flow and demographic modeling (Jackson et al. 2017, Scott et al. 2018) could allow for more reliable inferences for recently diverged populations.

Costs and benefits of different sequencing methods

With limited funding, cost will always be a consideration. Rapture has a somewhat costly initial investment but is still highly cost-effective ($13.42 sample including bait design and production for 1900 individuals in our study, or <$10 per sample if baits are already available; Table 3) in terms of cost per sample when compared to either BestRAD or low-coverage WGS. Given these low costs, Rapture is a very attractive method for conducting future work in the darter system, especially when extensive individual-level and spatial sampling are important components of the project design. The data produced by Rapture can be supplemented by low-coverage WGS if this is needed for the study, and the relatively high cost per individual of WGS in this study (~$275 per sample) could potentially be reduced by using poolseq (Schlötterer et al. 2014).

Overall, the Rapture method outlined here represents a potentially powerful methodology for phylogenomics and population genomics, both in darters and in diverse radiations of non-model organisms more generally. We have also shown several potential pitfalls associated with using heterologous genomes. While targeted sequence capture seems to mitigate some of these pitfalls, choosing to use a heterologous reference genome still has consequences that should be carefully considered during study design. As more reference genomes and sequence capture methods become available, Rapture will become an increasingly attractive option, especially when large sample sizes, extensive spatial coverage, or high read depth are important.
References

Ali, O. A., O’Rourke, S. M., Amish, S. J., Meek, M. H., Luikart, G., Jeffres, C., & Miller, M. R. (2016). Rad capture (Rapture): Flexible and efficient sequence-based genotyping. *Genetics, 202*(2), 389–400. https://doi.org/10.1534/genetics.115.183665

Bossu, C. M., & Near, T. J. (2013). Characterization of a contemporaneous hybrid zone between two darter species (*Etheostoma bison* and *E. caeruleum*) in the Buffalo River System. *Genetica, 141*(1–3), 75–88. https://doi.org/10.1007/s10709-013-9707-8

Brawand, D., Wagner, C. E., Li, Y. I., Malinsky, M., Keller, I., Fan, S., … Di Palma, F. (2015). The genomic substrate for adaptive radiation in African cichlid fish. *Nature, 513*(7518), 375–381. https://doi.org/10.1038/nature13726

Catchen, J. M., Hohenlohe, P. A., Bernatchez, L., Funk, W. C., Andrews, K. R., & Allendorf, F. W. (2017). Unbroken: RADseq remains a powerful tool for understanding the genetics of adaptation in natural populations. *Molecular Ecology Resources, 17*(3), 362–365. https://doi.org/10.1111/1755-0998.12669

Catchen, J., Hohenlohe, P. A., Bassham, S., Amores, A., & Cresko, W. A. (2013). Stacks: an analysis tool set for population genomics. *Molecular Ecology, 22*(11), 3124–3140. https://doi.org/10.1111/mec.12354

Chapman, J. A., Ho, I., Sunkara, S., Luo, S., Schroth, G. P., & Rokhsar, D. S. (2011). Meraculous: De novo genome assembly with short paired-end reads. *PLoS ONE, 6*(8). https://doi.org/10.1371/journal.pone.0023501

Chaves, J. A., Cooper, E. A., Hendry, A. P., Podos, J., De León, L. F., Raeymaekers, J. A. M., … Uy, J. A. C. (2016). Genomic variation at the tips of the adaptive radiation of Darwin’s finches. *Molecular Ecology, 25*(21), 5282–5295. https://doi.org/10.1111/mec.13743

da Fonseca, R. R., Albrechtsen, A., Themudo, G. E., Ramos-Madrigal, J., Sibbesen, J. A., Maretty, L., … Pereira, R. J. (2016). Next-generation biology: Sequencing and data analysis approaches for non-model organisms. *Marine Genomics, 30*, 3–13. https://doi.org/10.1016/j.margen.2016.04.012

David, K. T., Wilson, A. E., & Halanych, K. M. (2019). Sequencing Disparity in the Genomic Era. *Molecular Biology and Evolution, 36*(8), 1624–1627. https://doi.org/10.1093/molbev/msz117

Eaton, D. A. R., & Overcast, I. (2020). ipyrad: Interactive assembly and analysis of RADseq datasets. *Bioinformatics*. https://doi.org/10.1093/bioinformatics/btz966

Ekblom, R., & Galindo, J. (2011, July 8). Applications of next generation sequencing in molecular ecology of non-model organisms. *Heredity, Vol. 107*, pp. 1–15. https://doi.org/10.1038/hdy.2010.152
Faircloth, B. C., McCormack, J. E., Crawford, N. G., Harvey, M. G., Brumfield, R. T., & Glenn, T. C. (2012). Ultraconserved Elements Anchor Thousands of Genetic Markers Spanning Multiple Evolutionary Timescales. Systematic Biology, Vol. 61, pp. 717–726. https://doi.org/10.2307/41677973

Fitzpatrick, S. W., Crockett, H., & Funk, W. C. (2014). Water availability strongly impacts population genetic patterns of an imperiled Great Plains endemic fish. Conservation Genetics, 15(4), 771–788. https://doi.org/10.1007/s10592-014-0577-0

Foll, M., & Gaggiotti, O. (2008). A genome-scan method to identify selected loci appropriate for both dominant and codominant markers: A Bayesian perspective. Genetics, 180(2), 977–993. https://doi.org/10.1534/genetics.108.092221

Gabel, J. M., Dakin, E. E., Freeman, B. J., & Porter, B. A. (2008). Isolation and identification of eight microsatellite loci in the Cherokee darter (Etheostoma scotti) and their variability in other members of the genera Etheostoma, Ammocrypta, and Percina. Molecular Ecology Resources, 8(1), 149–151. https://doi.org/10.1111/j.1471-8286.2007.01903.x

Galinsky, K. J., Bhatia, G., Loh, P. R., Georgiev, S., Mukherjee, S., Patterson, N. J., & Price, A. L. (2016). Fast Principal-Component Analysis Reveals Convergent Evolution of ADH1B in Europe and East Asia. American Journal of Human Genetics, 98(3), 456–472. https://doi.org/10.1016/j.ajhg.2015.12.022

Galla, S. J., Forsdick, N. J., Brown, L., Hoeppner, M., Knapp, M., Maloney, R. F., … Steeves, T. E. (2018). Reference Genomes from Distantly Related Species Can Be Used for Discovery of Single Nucleotide Polymorphisms to Inform Conservation Management. Genes, 10(1), 9. https://doi.org/10.3390/genes10010009

George, M. (2018). Phylogeny of the Orangethroat Darter (Etheostoma spectabile) species complex in the Ozark Highlands of Arkansas. Animal Science Undergraduate Honors Theses. Retrieved from https://scholarworks.uark.edu/anscuht/22

Gilbert, P. S., Chang, J., Pan, C., Sobel, E. M., Sinsheimer, J. S., Faircloth, B. C., & Alfaro, M. E. (2015). Genome-wide ultraconserved elements exhibit higher phylogenetic informativeness than traditional gene markers in percomorph fishes. Molecular Phylogenetics and Evolution, 92, 140–146. https://doi.org/10.1016/j.ympev.2015.05.027

Grover, C. E., Salmon, A., & Wendel, J. F. (2012). Targeted sequence capture as a powerful tool for evolutionary analysis. American Journal of Botany, 99(2), 312–319. https://doi.org/10.3732/ajb.1100323

Harvey, M., Smith, B., Glenn, T., … B. F.-S., & 2016, undefined. (n.d.). Sequence capture versus restriction site associated DNA sequencing for shallow systematics. Academic.Oup.Com. Retrieved from https://academic.oup.com/sysbio/article-abstract/65/5/910/2223860
Hoffberg, S. L., Kieran, T. J., Catchen, J. M., Devault, A., Faircloth, B. C., Mauricio, R., & Glenn, T. C. (2016). RADcap: sequence capture of dual-digest RADseq libraries with identifiable duplicates and reduced missing data. *Molecular Ecology Resources, 16*(5), 1264–1278. https://doi.org/10.1111/1755-0998.12566

Hollingsworth Jr, P. R., & Near, T. J. (2009). Temporal patterns of diversification and microendemism in eastern highland endemic barcheek darters (Percidae: Etheostomatinae). *Evolution, 63*(1), 228–243. https://doi.org/10.1111/j.1558-5646.2008.00531.x

Hudman, S. P., Grose, M. J., Landis, J. B., Skalski, G. T., & Wiley, E. O. (2008). Twenty-three microsatellite DNA loci for population genetic studies and parentage assignment in orangethroat darter. *Molecular Ecology Resources, 8*(6), 1483–1485. https://doi.org/10.1111/j.1755-0998.2008.02312.x

Jackson, N. D., Morales, A. E., Carstens, B. C., & O’Meara, B. C. (2017). PHRAPL: Phylogeographic Inference Using Approximate Likelihoods. *Systematic Biology, 66*(6), 1045–1053. https://doi.org/10.1093/sysbio/syx001

Jelks, H. L., Walsh, S. J., Burkhead, N. M., Contreras-Balderas, S., Diaz-Pardo, E., Hendrickson, D. A., … Warren, M. L. (2008). Conservation Status of Imperiled North American Freshwater and Diadromous Fishes. *Fisheries, 33*(8), 372–407. https://doi.org/10.1577/1548-8446-33.8.372

Kelly, N. B., Near, T. J., & Alonzo, S. H. (2012). Diversification of egg-deposition behaviours and the evolution of male parental care in darters (Teleostei: Percidae: Etheostomatinae). *Journal of Evolutionary Biology, 25*(5), 836–846. https://doi.org/10.1111/j.1420-9101.2012.02473.x

Khudamrongsawat, J., Heath, L. S., Heath, H. E., & Harris, P. M. (2007). Microsatellite DNA primers for the endangered vermilion darter, *Etheostoma chermocki*, and cross-species amplification in other darters (Percidae: *Etheostoma*). *Molecular Ecology Notes, 7*(5), 811–813. https://doi.org/10.1111/j.1471-8286.2007.01712.x

Komoroske, L. M., Miller, M. R., O’Rourke, S. M., Stewart, K. R., Jensen, M. P., & Dutton, P. H. (2019). A versatile Rapture (RAD-seq/Capture) platform for genotyping marine turtles. *Molecular Ecology Resources, 19*(2), 497–511. https://doi.org/10.1111/1755-0998.12980

Korneliussen, T. S., Albrechtsen, A., & Nielsen, R. (2014). ANGSD: Analysis of Next Generation Sequencing Data. *BMC Bioinformatics, 15*(1), 356. https://doi.org/10.1186/s12859-014-0356-4

López-Giráldez, F., & Townsend, J. P. (2011). PhyDesign: An online application for profiling phylogenetic informativeness. *BMC Evolutionary Biology, 11*(1), 152. https://doi.org/10.1186/1471-2148-11-152
Lemmon, A. R., Emme, S. A., & Lemmon, E. M. (2012). Anchored Hybrid Enrichment for Massively High-Throughput Phylogenomics. *Syst. Biol.*, 61(5), 727–744. https://doi.org/10.1093/sysbio/sys049

Lemmon, E. M., & Lemmon, A. R. (2013). High-Throughput Genomic Data in Systematics and Phylogenetics. *Annual Review of Ecology, Evolution, and Systematics*, 44(1), 99–121. https://doi.org/10.1146/annurev-ecolsys-110512-135822

Li, H., Bioinformatics, R. D.-, & 2009, undefined. (n.d.). Fast and accurate short read alignment with Burrows–Wheeler transform. *Oxford University Press*.

Li, H., Handsaker, B., Wysoker, A., Fennell, T., Ruan, J., Homer, N., … Durbin, R. (2009). The Sequence Alignment/Map format and SAMtools. *Bioinformatics*, 25(16), 2078–2079. https://doi.org/10.1093/bioinformatics/btp352

Lowry, D. B., Hoban, S., Kelley, J. L., Lotterhos, K. E., Reed, L. K., Antolin, M. F., & Storfer, A. (2017). Breaking RAD: an evaluation of the utility of restriction site-associated DNA sequencing for genome scans of adaptation. *Molecular Ecology Resources*, 17(2), 142–152. https://doi.org/10.1111/1755-0998.12635

MacGuigan, D. J., & Near, T. J. (2019). Phylogenomic Signatures of Ancient Introgression in a Rogue Lineage of Darters (Teleostei: Percidae). *Systematic Biology*, 68(2), 329–346. https://doi.org/10.1093/sysbio/syy074

Manel, S., Perrier, C., Pratlong, M., Abi-Rached, L., Paganini, J., Pontarotti, P., & Aurelle, D. (2016). Genomic resources and their influence on the detection of the signal of positive selection in genome scans. *Molecular Ecology*, 25(1), 170–184. https://doi.org/10.1111/mec.13468

Margres, M. J., Jones, M. E., Epstein, B., Kerlin, D. H., Comte, S., Fox, S., … Storfer, A. (2018). Large-effect loci affect survival in Tasmanian devils (*Sarcophilus harrisii*) infected with a transmissible cancer. *Molecular Ecology*, 27(21), 4189–4199. https://doi.org/10.1111/mec.14853

Meek, M. H., & Larson, W. A. (2019). The future is now: Amplicon sequencing and sequence capture usher in the conservation genomics era. *Molecular Ecology Resources*, 19(4), 795–803. https://doi.org/10.1111/1755-0998.12998

Meisner, J., & Albrechtsen, A. (2018). Inferring population structure and admixture proportions in low-depth NGS data. *Genetics*, 210(2), 719–731. https://doi.org/10.1534/genetics.118.301336

Mendelson, T. C. (2003). Sexual isolation evolves faster than hybrid inviability in a diverse and sexually dimorphic genus of fish (Percidae: *Etheostoma*). *Evolution*, 57(2), 317–327. https://doi.org/10.1111/j.0014-3820.2003.tb00266.x
Moran, R. L., Catchen, J. M., & Fuller, R. C. (2020). Genomic resources for darters (Percidae: Etheostominae) provide insight into postzygotic barriers implicated in speciation. *Molecular Biology and Evolution, 37*(3), 711–729. https://doi.org/10.1093/molbev/msz260

Moran, R. L., & Fuller, R. C. (2018). Male-driven reproductive and agonistic character displacement in darters and its implications for speciation in allopatry. *Current Zoology, 64*(1), 101–113. https://doi.org/10.1093/cz/zox069

Moran, R. L., & Fuller, R. C. Agonistic character displacement of genetically based male colour patterns across darters. *Proceedings of the Royal Society B: Biological Sciences, 285*(1884). https://doi.org/10.1098/rspb.2018.1248

Moran, R. L., Zhou, M., Catchen, J. M., & Fuller, R. C. (2018). Hybridization and postzygotic isolation promote reinforcement of male mating preferences in a diverse group of fishes with traditional sex roles. *Ecology and Evolution, 8*(18), 9282–9294. https://doi.org/10.1002/ece3.4434

Moran, R. L., Zhou, M., Catchen, J. M., & Fuller, R. C. (2017). Male and female contributions to behavioral isolation in darters as a function of genetic distance and color distance. *Evolution, 71*(10), 2428–2444. https://doi.org/10.1111/evo.13321

Near, T. J., & Benard, M. F. (2004). Rapid allopatric speciation in logperch darters (Percidae: *Percina*). *Evolution, 58*(12), 2798–2808. https://doi.org/10.1111/j.0014-3820.2004.tb01631.x

Near, T. J., Bossu, C. M., Bradburd, G. S., Carlson, R. L., Harrington, R. C., Hollingsworth, P. R., … Etnier, D. A. (2011). Phylogeny and temporal diversification of darters (Percidae: Etheostomatinae). *Systematic Biology, 60*(5), 565–595. https://doi.org/10.1093/sysbio/syr052

Nevado, B., Ramos-Onsins, S. E., & Perez-Enciso, M. (2014). Resequencing studies of nonmodel organisms using closely related reference genomes: optimal experimental designs and bioinformatics approaches for population genomics. *Molecular Ecology, 23*(7), 1764–1779. https://doi.org/10.1111/mec.12693

Nguinkal, J. A., Brunner, R. M., Verleih, M., Rebl, A., de los Ríos-Pérez, L., Schäfer, N., … Goldammer, T. (2019). The First Highly Contiguous Genome Assembly of Pikeperch (*Sander lucioperca*), an Emerging Aquaculture Species in Europe. *Genes, 10*(9), 708. https://doi.org/10.3390 GENES10090708

Nguyen, L.-T., Schmidt, H. A., von Haeseler, A., & Minh, B. Q. (2015). IQ-TREE: A Fast and Effective Stochastic Algorithm for Estimating Maximum-Likelihood Phylogenies. *Molecular Biology and Evolution, 32*(1), 268–274. https://doi.org/10.1093/molbev/msu300
Oliveria, D.A., Reid, B.R., Fitzpatrick, S. W. (2020). Genome-Wide Diversity Underlies Fine-Scale Thermal Variation in the Rainbow Darter (*Etheostoma caeruleum*). *Evolutionary Applications*, in review.

Paradis, E., & Schliep, K. (2019). *ape 5.0*: an environment for modern phylogenetics and evolutionary analyses in R. *Bioinformatics*, 35(3), 526–528. https://doi.org/10.1093/bioinformatics/bty633

Peek, R. A., Bedwell, M., O’Rourke, S. M., Goldberg, C., Wengert, G. M., & Miller, M. R. (2019). Hybridization between two parapatric ranid frog species in the northern Sierra Nevada, California, USA. *Molecular Ecology*, 28(20), 4636–4647. https://doi.org/10.1111/mec.15236

Putnam, N. H., O’Connell, B. L., Stites, J. C., Rice, B. J., Blanchette, M., Calef, R., … Green, R. E. (2016). Chromosome-scale shotgun assembly using an in vitro method for long-range linkage. *Genome Research*, 26(3), 342–350. https://doi.org/10.1101/gr.193474.115

Revell, L. J. (2012). phytools: An R package for phylogenetic comparative biology (and other things). *Methods in Ecology and Evolution*, 3(2), 217–223. https://doi.org/10.1111/j.2041-210X.2011.00169.x

Rochette, N. C., Rivera-Colón, A. G., & Catchen, J. M. (2019). Stacks 2: Analytical methods for paired-end sequencing improve RADseq-based population genomics. *Molecular Ecology*, 28(21), 4737–4754. https://doi.org/10.1111/mec.15253

Russell, J. J., Theriot, J. A., Sood, P., Marshall, W. F., Landweber, L. F., Fritz-Laylin, L., … Brunet, A. (2017). Non-model model organisms. *BMC Biology*, 15(1), 1–31. https://doi.org/10.1186/s12915-017-0391-5

Saarinen, E. V., & Austin, J. D. (2010). When Technology Meets Conservation: Increased Microsatellite Marker Production Using 454 Genome Sequencing on the Endangered Okaloosa Darter (*Etheostoma okaloosae*). *Journal of Heredity*, 101(6), 784–788. https://doi.org/10.1093/jhered/esq080

Sard, N. M., Smith, S. R., Homola, J. J., Kanefsky, J., Bravener, G., Adams, J. V., … Scribner, K. T. (2020). RAPTURE (RAD capture) panel facilitates analyses characterizing sea lamprey reproductive ecology and movement dynamics. *Ecology and Evolution*, 10(3), 1469–1488. https://doi.org/10.1002/ece3.6001

Schlötterer, C., Tobler, R., Kofler, R., & Nolte, V. (2014, November 25). Sequencing pools of individuals-mining genome-wide polymorphism data without big funding. *Nature Reviews Genetics*, Vol. 15, pp. 749–763. https://doi.org/10.1038/nrg3803

Scott, P. A., Glenn, T. C., & Rissler, L. J. (2018). Resolving taxonomic turbulence and uncovering cryptic diversity in the musk turtles (*Sternotherus*) using robust demographic
modeling. *Molecular Phylogenetics and Evolution, 120*, 1–15.  
https://doi.org/10.1016/j.ympev.2017.11.008

Shafer, A. B. A., Wolf, J. B. W., Alves, P. C., Bergström, L., Bruford, M. W., Brännström, I., … Zieliński, P. (2015, February 1). Genomics and the challenging translation into conservation practice. *Trends in Ecology and Evolution, Vol. 30*, pp. 78–87.  
https://doi.org/10.1016/j.tree.2014.11.009

Switzer, J. F., Welsh, S. A., & King, T. L. (2008). Microsatellite DNA primers for the candy darter, *Etheostoma osburni* and variegate darter, *Etheostoma variatum*, and cross-species amplification in other darters (Percidae). *Molecular Ecology Resources, 8*(2), 335–338.  
https://doi.org/10.1111/j.1471-8286.2007.01946.x

Tonnis, B. D. (2006). Microsatellite DNA markers for the rainbow darter, *Etheostoma caeruleum* (Percidae), and their potential utility for other darter species. *Molecular Ecology Notes, 6*(1), 230–232.  
https://doi.org/10.1111/j.1471-8286.2005.01203.x

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**Data accessibility statement**  
The *E. cragini* Whole Genome Shotgun project has been deposited at DDBJ/ENA/GenBank under the accession JAAVJE000000000. The version described in this paper is version JAAVJE010000000. Short-read data have been uploaded to the NCBI as BioProject PRJNA611833. Analysis scripts and capture bait sequences are available at https://github.com/nerdbrained/darter_rapture.
Table 1. Cost of Rapture and WGS methods used in this study.

| Method          | # samples | Pilot dataset | Baits | Library preparation | Sequencing | Total cost | Cost/sample | Cost/sample after bait design |
|-----------------|-----------|---------------|-------|---------------------|------------|------------|-------------|-----------------------------|
| BestRAD (pilot) | 52        | N/A           | N/A   | $733.00             | $2,661.00  | $3,394.00  | $65.27      |                             |
| Rapture         | 1900      | $3,394.00     | $3,600.00 | $5,190.00         | $13,305.00 | $25,489.00 | $13.42      | $9.73                       |
| WGS             | 24        | N/A           | N/A   | $2,616.00           | $3,991.50  | $6,607.50  | $275.31     |                             |
Figure 1. Flow chart showing procedures used to design *E. cragini* Rapture baits from BestRAD RADseq data, test the Rapture baits using a subsequent round of BestRAD RADseq, and compare the results of population genomic analyses using Rapture versus WGS data.

**a. Bait design**

- **Initial BestRAD**
- **Identify Short/Outlier Loci**
- **Identify Long Loci**

- **SbfI cut site**
- **120 bp locus**
- **240 bp locus**
- **80-bp Baits**
- **120 bp locus**
- **240 bp locus**

**b. Rapture**

- **1900 individuals**
- **BestRAD**
- **Bait Capture**
- **Sequencing**
- **Quality Filtering**
- **Alignment**
- **Population genetics**

- **Native genome**
- **Heterologous genome**

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54 individuals

80-bp Baits: Bait1, Bait2, Bait3, Bait4, Bait5

b. Rapture

1900 individuals

BestRAD, Bait Capture

Sequencing, Quality Filtering, Alignment

Native genome, Heterologous genome

Shotgun library preparation

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Figure 2. Read depth and number of read pairs for whole genome sequencing data. Average depth is shown for either a subset of sites with at least 1x coverage or for all sites in the genome, with alignment to either the native *E. cragini* genome or the heterologous genome of a closely related species (*E. spectabile*).
Figure 3. Coverage for Rapture loci (either all loci combined or short, long, and outlier loci taken separately) mapped to the *E. cragini* genome across four *Etheostoma* species.
Figure 4. Heterozygosity estimates for different datasets. 1:1 dotted line (expected for complete agreement across datasets) is shown.
Figure 5. Admixture plots and mapped ancestry proportions. Each line in barplots represents an individual, and colors represent proportion of ancestry for each individual assigned to a given population. For the maps, pie charts represent either ancestry proportions aggregated for all individuals at a given site (for Rapture data) or admixture proportions for a single individual (for WGS data). Text on barplots indicates drainage of origin.

5a. Rapture loci, full dataset, *E. cragini* reference.
5b. Rapture loci, full dataset, *E. spectabile* reference
5c. WGS data, E. cragini reference
5d. WGS, E. spectabile reference
Figure 6. Manhattan plot of PCA-based selection scan results for Rapture data. X-axis represents position of each SNP in the genome, with chromosomes delimited by dotted lines. p-values for selection scan are shown on the Y axis on a -log(10) scale, and the significance threshold for each dataset is shown as a horizontal dotted line. For Rapture data, outlier loci are shown as black dots.
Figure 7a. Time-calibrated maximum likelihood phylogeny for Rapture data. Eca = *E. caeruleum*, Esp = *E. spectabile*, Efl = *E. flabellare*, Ecr = *E. cragini*. Node labels for *E. caeruleum* individuals include a site identifier, and node labels for *E. cragini* individuals include a metapopulation identifier followed by a site identifier. Time (on the x-axis) is expressed in millions of years ago.
7b. Phylogeny plotted in space.
Supporting Information 1.

We generated assembly statistics with QUAST v4.3 (Gurevich et al. 2013) to evaluate the quality of the *E. cragini* genome compared to the congeneric *E. spectabile* genome (v. UIUC_Espe_1.0, downloaded from NCBI; Moran et al. 2020) and to the confamilial *Perca flavescens* (v. PFLA_1.0, downloaded from NCBI; Feron et al. 2020), *Perca fluviatalis* (v. GENO_Pfluv_1.0; Ozerov et al. 2018), and *Sander lucioperca* (v. SLUC_FBN_1, downloaded from NCBI; Nguinkal et al. 2019) genomes. To assess completeness of the *E. cragini* assembly, we determined the number of Actinopterygii-specific Benchmarking Universal Single-Copy Orthologs (BUSCOs) identified with BUSCO v2.0 (Simão et al. 2015) using the actinopterygii_odb10 lineage dataset. We also used RepeatModeler (Smit et al. 2015) v.1.0.11 and RepeatMasker v.4.0.5 (Smit and Hubley 2015) to identify repetitive elements in the *E. cragini* assembly and in the most recent version of the *P. fluviatalis* genome. We combined this information with published data to compare the proportion of repetitive elements among percid fish genomes. To compare genomic synteny and homology among genomes, we aligned the Dovetail sequence with confamilial genomes that contained chromosome-level assemblies (*E. spectabile, P. flavescens,* and *P. fluviatalis*) with Minimap2 (Li 2018) using the D-Genies web interface (dgenies.toulouse.inra.fr; Cabanettes and Klopp 2018). We visualized the alignment in D-Genies with a dotplot, using the “strong precision” setting to remove small matches. We converted the annotations for the *E. spectabile* genome into ENSEMBL format using a script (https://github.com/NBISweden/EMBLmyGFF3) and transferred annotations from the *E. spectabile* genome to the *E. cragini* genome using RATT v.0.95 (Otto et al. 2011).
The *E. cragini* reference genome generated by Dovetail included data from 461,647,761 2x150bp shotgun read pairs for the initial *de novo* genome assembly and approximately 194 million and 228 million 2x150bp read pairs for the Chicago and HiC libraries, respectively. The final assembly had an estimated physical coverage of 16,456.85x across 4,667 scaffolds, an L50 of 11, an N50 of 27.59 Mb, and a total length of 643.1 Mb (Supporting Table 1). A total of 97.6% of the genome was assembled into 24 chromosomes. The final *E. cragini* assembly size was smaller than predicted by initial estimates based on short-reads (31-mer estimated size of = 736.9 Mb), and was also smaller than all other percid genomes sequenced to date. However, the number of complete BUSCOs observed in the *E. cragini* assembly was comparable to previously published percid assemblies (Supporting Table 2), despite being 68-75% of the size of these other assemblies. Within the *E. cragini* assembly, 172 Mb (26.7% of total genome size) was classified as repetitive, representing both a lower total amount of repetitive sequence and a lower proportion of the genome classified as repetitive compared to other percids (Supporting Table 2).

Genome alignments showed many long regions of synteny between the two genomes, with 24 long scaffolds of the *E. cragini* genome mostly aligning to single chromosomes in the published genomes (Supporting Figure 1). There was also extensive evidence, however, for inversions and chromosomal rearrangements in all 24 chromosomes between the species, often near the ends of chromosomes. Inversions and rearrangements were more pronounced in the *E. cragini – E. spectabile* alignment, and relatively few sequences from the *E. cragini* assembly aligned to the unplaced scaffolds in the *E. spectabile* assembly compared to the chromosome-level scaffolds, although the *E. cragini – E. spectabile* alignment had higher identity overall (Supporting Figure 1). RATT successfully transferred 99% of annotation elements (12,497 of 12,849 possible
elements) across genomes, although some elements (14%) that were contiguous on the *E. spectabile* genome were split among several genomic regions or only partially transferred onto the *E. cragini* genome. All 740 gene models from the *E. spectabile* genome were transferred to the *E. cragini* genome, although 147 gene models (80%) were partially transferred with 161 exons found in the *E. spectabile* genome not found in *E. cragini*.

References:

Gurevich, A., Saveliev, V., Vyahhi, N., & Tesler, G. (2013). QUAST: quality assessment tool for genome assemblies. *Bioinformatics*, 29(8), 1072–1075.

Li, H. (2018). Minimap2: Pairwise alignment for nucleotide sequences. *Bioinformatics*, 34(18), 3094–3100.

Cabanettes, F., & Klopp, C. (2018). D-GENIES: Dot plot large genomes in an interactive, efficient and simple way. *PeerJ*, 2018(6), e4958.

Feron, R., Zahm, M., Cabau, C., Klopp, C., Roques, C., Bouchez, O., … Guiguen, Y. (2020). Characterization of a Y-specific duplication/insertion of the anti-Mullerian hormone type II receptor gene based on a chromosome-scale genome assembly of yellow perch, *Perca flavescens*. *Molecular Ecology Resources*, 20(2), 531–543.

Nguinkal, J. A., Brunner, R. M., Verleih, M., Rebl, A., de los Ríos-Pérez, L., Schäfer, N., … Goldammer, T. (2019). The First Highly Contiguous Genome Assembly of Pikeperch (Sander lucioperca), an Emerging Aquaculture Species in Europe. *Genes*, 10(9), 708.

Ozerov, M. Y., Ahmad, F., Gross, R., Pukk, L., Kahar, S., Kisand, V., & Vasemägi, A. (2018). Highly continuous genome assembly of eurasian perch (perca fluviatilis) using linked-read sequencing. *G3: Genes, Genomes, Genetics*, 8(12), 3737–3743.

Simão, F., Waterhouse, R., Ioannidis, P., Kriventseva, E., & Zdobnov, E. (2015). BUSCO: assessing genome assembly and annotation completeness with single-copy orthologs. *Bioinformatics*, 31(19), 3210–3212.

Smit, A.F.A. & Hubley, R. (2015) *RepeatModeler Open-4.0*. 2015 http://www.repeatmasker.org

Smit, A.F.A., Hubley, R., & Green, P. (2015) *RepeatMasker Open-4.0*. 2015
http://www.repeatmasker.org
## Supporting Table 1

| Species       | Complete | Single-copy | Duplicated | Fragmented | Missing | Total |
|---------------|----------|-------------|------------|------------|---------|-------|
| *E. cragini*  | 3449 (94.8%) | 3425 (94.1%) | 24 (0.7%)  | 46 (1.3%)  | 145 (3.9%) | 3640  |
| *E. spectabile* | 3442 (94.6%) | 3347 (92%)  | 95 (2.6%)  | 18 (0.5%)  | 180 (4.9%) | 3640  |
| *P. flavescens* | 3539 (97.2%) | 3506 (96.3%) | 33 (0.9%)  | 8 (0.2%)   | 93 (2.6%)  | 3640  |
| *P. fluviatilis* | 3509 (96.4%) | 3477 (95.5%) | 32 (0.9%)  | 16 (0.4%)  | 115 (3.2%) | 3640  |
| *S. lucioperca* | 3537 (97.1%) | 3499 (96.1%) | 38 (1%)    | 11 (0.3%)  | 92 (2.6%)  | 3640  |

## Supporting Table 2

| Species       | Total Length | N50 | L50 | % Gaps | %GC | % Repetitive |
|---------------|--------------|-----|-----|--------|-----|--------------|
| *E. cragini*  | 643,100,042  | 27,593,271 | 11  | 4,667  | 0.520 | 40.52        | 26.7 |
| *E. spectabile* | 854,790,067  | 30,497,795 | 13  | 3,118  | 0.470 | 40.91        | 30.9 |
| *P. flavescens* | 877,456,336  | 37,412,490 | 11  | 268    | 0.047 | 40.84        | 41.7 |
| *P. fluviatilis* | 951,362,726  | 39,550,354 | 11  | 304    | 0.031 | 40.90        | 40.6 |
| *S. lucioperca* | 900,477,756  | 4,929,547  | 52  | 1,313  | 0.003 | 40.92        | 39  |
Supporting Figure 1. Dotplots of whole-genome alignments between *E. cragini* and other chromosome-level percid fish genome assemblies after removing weak-precision alignments. The *E. cragini* genome is shown on the y-axis in each case. A summary of the proportion of total matches at various levels identity is shown below each dotplot.

(a) *E. spectabile* vs. *E. cragini*. 

- No match: 25.12 %
- < 25 %: 9.56 %
- < 50 %: 61.03 %
- < 75 %: 3.83 %
- > 75 %: 0.04 %
(b) *P. flavescens* vs. *E. cragini*.
(c) P. fluviatilis vs. E. cragini.
Supplemental Figure 2. Expanded coverage statistics at multiple read depths for different categories of Rapture loci.

(a) Short loci aligned to *E. cragini* reference genome.
(b) Short loci aligned to *E. spectabile* reference genome.

![Graphs showing total reads mapped to baits for different coverages](image-url)
(c) Long loci aligned to *E. cragini* reference genome.
(d) Long loci aligned to *E. spectabile* reference genome.
(e) Loci putatively under selection aligned to *E. cragini* reference genome.
(f) Loci putatively under selection aligned to *E. spectabile* reference genome.
Supplemental Figure 3. Coverage for either long (top) or short (bottom) Rapture loci for a single *E. cragini* individual. Overlapping blue lines on the left panels indicate per-base coverage for all bases within the Rapture locus and a 500-bp buffer around the locus. The right panels show the number of loci for which each base in this region is covered at >20x. The locations of the restriction cut site and the baits are shown in the right panels as well. Read depth beyond the portions of the sequence covered by the capture baits shows a hump-shaped distribution corresponding to reverse reads with varying degrees of overlap created by the random shearing step. Read depth tends to drop off rapidly for short loci, although some bases up to 500 bp from the restriction site were represented at high coverage. For the long loci, however, coverage was consistently high at all bases up to approximately 500 bp in either direction from the restriction site.)
Supporting Figure 4. Inferred population structure for subsetted Rapture data.  
(a) *E. cragini* reference
(b) *E. spectabile* reference
Supporting Figure 5. Selection scan statistics for subsetted Rapture datasets.

Rapture subset aligned to *E. cragini*

Rapture subset aligned to *E. spectabile*
Supporting Figure 6. Phylogenetic informativeness profiles for short (NS), long (NL), and selected (SB) baits.