Polygenic discrimination of migratory phenotypes in an estuarine forage fish

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Abstract

Migration is a complex phenotypic trait with some species containing migratory and nonmigratory individuals. Such life history variation may be attributed in part to plasticity, epigenetics, or genetics. Although considered semianadromous, recent studies using otolith geochemistry have revealed life history variation within the critically endangered Delta Smelt. Broadly categorizable as migratory or freshwater residents, we examined Restriction site Associated DNA sequencing data to test for a relationship between genetic variation and migratory behaviors. As previously shown, we found no evidence for neutral population genetic structure within Delta Smelt; however, we found significant evidence for associations between genetic variants and life history phenotypes. Furthermore, discriminant analysis of principal components, hierarchical clustering, and machine learning resulted in accurate assignment of fish into the freshwater resident or migratory classes based on their genotypes. These results suggest the presence of adaptive genetic variants relating to life history variation within a panmictic population. Mechanisms that may lead to this observation are genotype dependent habitat choice and spatially variable selection, both of which could operate each generation and are not exclusive. Given that the population of cultured Delta Smelt are being used as a refugial population for conservation, as a supply for wild population supplementation, and currently represent the majority of all living individuals of this species, we recommend that the hatchery management strategy consider the frequencies of life history-associated alleles and how to maintain this important aspect of Delta Smelt biological variation while under captive propagation.

Keywords: adaptive genetic variation; Delta Smelt; migration; Osmeridae; resident ecotype; semianadromy

Introduction

The causes of migration may be viewed as both proximate and ultimate (Tinbergen 1963). Ultimate causes are concerned with the broader evolutionary mechanisms, origins, and consequences of migration. The scale of these questions allows comparison across taxa and unifying themes to be proposed such as the increasing food availability hypothesis and diadromy in fishes (Gross et al. 1988). Proximate causes are focused on individuals and the expression of migratory (MIG) behavior and associated traits in response to environmental cues, genetic background, and nongenetic parental affects (e.g. Ferguson et al. 2019). These proximate questions are much more limited in the applicability of results across taxonomic levels. In particular, genetic contributions to the control of migration have been widely documented across the literature (e.g. Liedvogel et al. 2011); however, the identification of a shared genomic region associated with MIG timing in 2 divergent species of Pacific salmon (Oncorhynchus) is atypical (Prince et al. 2017).

Studying the causes of migration relies on the characterization of MIG syndromes—a compendium of movement patterns, behavioral, morphological, and physical traits—of individuals within a population (Sih et al. 2004; Dingle 2006). In the most extreme cases, such as partial migration, individuals within the same population exhibit different behaviors, with some individuals (migrants) making movements across habitats, and other nonmigrating individuals (residents) completing their life cycle within a single habitat type (Chapman et al. 2012; Kendall et al. 2015; Hobbs et al. 2019). Individuals of a species exhibiting different MIG syndromes are subject to different selection pressures in the form of suites of predators, physical environmental conditions, trophic resources, growth rates, and anthropogenic disturbances (Gross 1987; Moyle and Cech 2004). The complexity of alternative MIG syndromes may be reflected by a complex genetic background with hundreds of loci covarying between distinct ecotypes (e.g. Pavey et al. 2015). Life history variation and ecotypic differentiation may also be influenced by multiple genes.
(e.g. over 1,000) that are consolidated into supergenes (Thorneycroft 1975; Lowry and Willis 2010; Pearse et al. 2014, 2019; Tuttle et al. 2016), with the possibility of multiple (2 to several) supergene complexes present within a single species (Huang et al. 2020; Campbell et al. 2021). Despite this potential complexity, large differences in behavioral phenotypes result from minor differences in the genome. For example, major differences in migration timing (e.g. spring-run vs fall-run Chinook Salmon) are associated with a simple Mendelian polymorphism (Thompson et al. 2020). Epigenetic control of migration in Rainbow Trout (Oncorhynchus mykiss) has been demonstrated (Baerwald et al. 2016), indicating that heritable nongenetic variation can be considered. In contrast, it is also possible that life history and ecotypic diversity may simply represent phenotypic plasticity without a heritable foundation (Gotthard and Nylin 1995).

Although California’s critically endangered Delta Smelt (Hypomesus transpacificus) has been broadly described as semianadromous (Moyle et al. 1992; Sommer et al. 2011), recent results based on otolith geochemical analyses have identified complex MIG histories in this species (Hobbs et al. 2019). Otoliths are calcium carbonate structures in the inner ear of most bony fishes that form continuously throughout the life of a fish (Campagna 1999). The geochemical analysis of the otoliths can be used to reconstruct movement between different habitats, along rivers and within the different salinities of estuarine environments. As originally described, the Delta Smelt population is dominated in most years by a semianadromous MIG phenotype that spawns in freshwater and rears in brackish waters of the San Francisco Estuary. However, Hobbs et al. (2019) also identified a significant fraction of the population as freshwater residents (FWR) as well as a rarer phenotype that hatches and rears in low-salinity brackish habitats (BWR). Although Delta Smelt can be found in different regions of the estuary, no geographic population structuring has been identified for this species, thus it is managed as a single panmictic population (Fisch et al. 2011).

Given that the Delta Smelt population is now <1% of its former levels (Moyle et al. 2016; Hobbs et al. 2017), the survival of the species now rests largely on hatchery production, which serves as both a genetic-reserve population and a source of fish for supplementation of the wild population. Although culture practices have attempted to follow sound genetic management guidelines, this recently has been hampered by limited introductions of genetic diversity coming from the few wild-caught broodstock in each year. Due to the decline of the naturally reproducing Delta Smelt no natural origin fish contributed to hatchery broodstock in 2020 and 2021. Furthermore, strong hatchery domestication pressures exist on the reserve population (Finger et al. 2018). Any potential genetic diversity associated with life history variation has yet to be considered in the genetic management of the species, thus leaving captive propagation and planned genetic monitoring following the release of cultured individuals into the wild uninformed with respect to life history diversity.

Here, we conduct an interdisciplinary study, combining the results of otolith geochemistry and Restriction Site Associated DNA sequencing (RADseq), to investigate several important questions regarding the potential genetic underpinning for the MIG behaviors of Delta Smelt and related implications for species management and conservation. Specifically, we examine whether there is evidence for a genetic basis of life-history variation within Delta Smelt, how any such genetic basis might be characterized, and how well the MIG phenotype of an individual can be predicted from its genotype. Results of this work illuminate the control of MIG behaviors in Delta Smelt and how such phenotypic diversity might be incorporated into culture and supplementation strategies.

**Materials and methods**

**Samples and phenotypic classification**

The Delta Smelt were collected by the California Department of Fish & Wildlife’s Spring Kodiak Trawl Survey between January and March 2013. Each fish was measured fresh in the field, given a unique individual serial number, and archived in liquid nitrogen. Otoliths were extracted and prepared for geochemical analysis following methods detailed in Hobbs et al. (2019). Otoliths in this study were analyzed in situ at the UC Davis Interdisciplinary Center for Plasma Mass Spectrometry using a multicollector inductively coupled plasma mass spectrometer (Nu Plasma HR from Nu Instrument Inc.) interfaced with a Nd:YAG 213 nm laser (New Wave Research UP213) (LA–MC–ICP–MS). Strontium isotope ratios ($^{87}$Sr/$^{86}$Sr) were obtained from the core to the ventral edge of the otolith, representing the entire lifespan of the fish. Individuals expressing an FWR phenotype exhibit $^{87}$Sr/$^{86}$Sr profiles with values remaining below 0.7075 across the entire profile (entire life span), corresponding with practical salinities <0.5 (i.e. “freshwater”) (Hobbs et al. 2019). In contrast, individuals expressing a semianadromous MIG phenotype exhibit $^{87}$Sr/$^{86}$Sr profiles that begin at values <0.7075 for at least the first 30 days post-hatch, followed by a transition into brackish-water habitats, identified by $^{87}$Sr/$^{86}$Sr values between 0.7075 and 0.7092.

**Genetic analysis**

Total genomic DNA was extracted from tissues using a Qiagen DNEasy extraction kit following the manufacturer’s protocols. RADseq libraries were generated with the SbfI enzyme following the Best Rad protocol (Ali et al. 2016). Libraries were sequenced with 100 base pair paired-end sequencing on an Illumina HiSeq 4000 at the UC Davis Genome Center.

Sequence data files were aligned to the Delta Smelt reference genome (GCA 021917145.1) with the Burrows–Wheeler Aligner using the MEM algorithm (Li and Durbin 2009). Resulting alignments were sorted, PCR duplicates removed and coverage in terms of the number of aligned reads were calculated with SAMtools (Li et al. 2009). We removed individuals from the analysis that constituted the bottom 25% of coverage in terms of aligned reads and lacked complete phenotype classification and sex metadata. For all analyses, we analyzed assembled contigs of the Delta Smelt genome >500 kb totaling 400,418,546 bases representing 92% of bases in the assembly. The contigs larger than 500 kb in the assembly numbered 73 with a total 376 contigs in the assembly.

We searched for a signal of neutral genetic structure within sampled Delta Smelt with PCAngsd through principal component (PC) analysis (Meisner and Albrechtsen 2018). PCAngsd required a genotype likelihood file that we generated through ANGSD using a SAMtools likelihood model (-GL 1) in the Beagle format (-doGl 2) (Korneliussen et al. 2014). For quality control thresholds, we required a site to be present in 90% of individuals (-minInd 108), a significance value of 1e−6 (-SNP_pval), a minimum mapping quality of 20 (-minMapQ), and a minimum base quality of 20 (-minQ). The resulting genotype likelihoods were analyzed with the default settings of PCAngsd, which includes a minimum minor allele frequency (MAF) of 0.05.

Genetic variants associated with life history variation were identified through a genome-wide association study (GWAS) implemented in ANGSD. We used a generalized linear framework (-doAsso 2) and provided sex as a covariate (-cov). We applied the following options to the program to provide quality control -minMapQ 20, -minQ 20, -SNP_pval 1e6, and -minInd 91. We specified a SAMtools genotype
model (-GL 1) and -minCount 2. To determine a significance threshold, we do not use a Bonferroni correction as sites in the genome are not independent. Instead, we calculated a significance threshold in a Bayesian framework (e.g., Burton et al. 2007). We assumed for the calculation of a prior probability that life history variation in Delta Smelt is polygenic and that closely linked variants to SNPs underlying phenotypic differences would also be detected by GWAS. Following this assumption, we assumed for calculation of a prior probability, \( P(T) \), that there are 25 genetic variants that may contribute to life history variation within the GWAS, \( P(T) = 25/\text{number of variants} \). The significance level can be calculated by the formula: 

\[
\alpha = \text{Prior Odds} \times 1/\text{Posterior Odds} \quad (e.g., \text{Wacholder et al. 2004}), \text{where prior odds is } P(T)/(1 - P(T)), \text{posterior odds is calculated based on 95% certainty of observing a significant effect that is real (0.95/(1 – 0.95)) and an upper bound of power (1) is used.}

We examined the strength of the signal in the genetic data to separate Delta Smelt in FWR and MIG categories with discriminant analysis of PCs (DAPC), hierarchical clustering, and machine learning approaches. With these approaches we used called genotypes generated by ANGSD (-doGeno 2) producing a dataset of 0, 1, 2 coded variation. We applied the same quality control thresholds as used previously, with the following changes: an MAF specified (-minMaf 0.05); posterior cutoff specified (-postCutoff 0.9), and a 90% missing individual threshold (-minInd 109). The genotypes were uploaded into R for further analysis (R Development Core Team 2020).

DAPC identifies the largest axis of variation between predefined groups. We conducted DAPC using the adegenet package in R (Jombart 2008) with MIG phenotype supplied as an a priori grouping variable. We visualized the ability of DAPC to classify the fish into phenotype classes by visualizing the first discriminant function and generating histograms of the posterior probability of assignment to prior classes. We also examined the contributions (loadings) of each site to the separation of phenotypic classes. Separately we investigated the predictive power of genotypes to predict phenotype. We examined the top 200 associated variants identified by GWAS with ANGSD for respective genotype calls and then generated a heatmap with the heatmap.2 function of the gplots package. Clustering of samples and loci was done with Ward’s distance, and missing data were not treated. The same set of 200 most-associated SNPs was then analyzed with a k-nearest neighbor (knn) approach for classification after treating missing data with the na.roughfix function of the randomForest package. We divided our dataset into a training dataset of 30 randomly selected FWR and 30 randomly selected MIG fish. With the training dataset, we identified a best k with a repeated k-fold cross validation using the trainControl and train functions of the caret library. A best k was identified as being most accurate after 100 sampling events and 10-folds for odd values of k from 1 to 29. The selected best k was then applied to all samples for the top associated variants and a cross table of accuracy computed with the CrossTable function of the gmodels library.

**Functional analysis**

Annotated genes located near SNPs distinguished by GWAS were identified to provide insight into the functional relevance of the polymorphisms to MIG phenotypes. We determined first the extent to which linkage disequilibrium (LD) operates in the Delta Smelt genome given our data, and then obtained annotated features within that window. Descriptions of genes, Gene Ontology (GO) terms, GO names, and GO definitions were obtained with the biomaRt package in R by querying the zebrafish dataset (drerio_gene_ensemble) (Durinck et al. 2005, 2009).

**Results**

After filtering for coverage and complete metadata, our dataset contained 60 FWR and 61 MIG individuals closely split between sexes (Table 1). Sample metadata is reported in Supplementary Document 1 with FWR individuals largely restricted to the northern region of the Sacramento-San Joaquin Delta while MIG individuals were more-widely distributed (Fig. 1a). Based on the reference assembly, a theoretical number of RAD loci based on the Sfj restriction site is 16,578. Genotype likelihoods were identified from 11,638 RAD loci (63%, Supplementary Appendix 1). PC analysis of 18,765 SNPs did not suggest that phenotype classification and genetic structuring were closely related overall (Fig. 1b). We identified 5 significantly associated variants from 13,376 sites examined by GWAS located on 4 linkage groups (Fig. 2 and Table 2). Two of the most-associated variants were found in close proximity, only 153 bp apart, on l002. The complete association test results are provided as Supplementary Document 2.

Calling genotypes produced 9,068 variants (provided as an R data binary as Supplementary File 1) composed of 7,274 independent SNPs that do not exhibit genetic structuring between FWR and MIG fish and represent ~11,090 RAD loci (Supplementary Appendix 1). Within DAPC, we specified 110 PCs (n.pca = 110) and a single axis for discriminant analysis (n.da = 1). The samples largely were separated into 2 groups with high posterior probability of assignment (Fig. 3). FWR individuals had a mean posterior assignment probability of 0.999 to the FWR class and 6.77 × 10⁻⁴ to the MIG class. The mean posterior assignment for the MIG to MIG class was 0.983, with 1.70 × 10⁻² of MIG fish to the FWR class. A single MIG individual had a FWR posterior assignment of 1.00, causing a large overall reduction in MIG posterior assignment probability. The sites contributing more than 0.0005 to the loadings are provided in Supplementary Document 3.

Hierarchical clustering with the 200 genotypes having the strongest phenotypic association creates 2 major groupings each representing a majority FWR or a majority MIG (Supplementary Fig. 1). Most, 88% (53/60), of FWR compose a cluster and most, 95% (58/61), of MIG individuals compose the other cluster. The same 200 genotypes exhibited the highest accuracy with knn with k = 13 neighbors (94%, Fig. 4a). Subsequently, we were able to successfully assign 88% of FWR individuals and 98% of the MIG individuals for an overall accuracy of 94% (Fig. 4b).

| Phenotype | Sex     | Average aligned reads | Sample size | Total N for phenotype |
|-----------|---------|-----------------------|-------------|-----------------------|
| FWR       | Male    | 1,080,559             | 34          | 60                    |
| FWR       | Female  | 1,022,383             | 26          |                       |
| MIG       | Male    | 892,302               | 32          | 61                    |
| MIG       | Female  | 1,091,578             | 29          |                       |

Table 1. Summary of Delta Smelt examined in this study reported by phenotype and sex.
Fig. 1. a) The geographic distribution of Delta Smelt samples examined in this study, facets are split between FWR and MIG individuals. b) A PC analysis of individuals examined in this study. The facets are split between a plot of PC1 vs PC2 and PC1 vs PC3.

Fig. 2. Manhattan plot of genome-wide association testing contrasting FWR and MIG Delta Smelt individuals. Sex was provided as a covariant and the linkage groups of the Delta Smelt genome assembly are shown. Sites exceeding the significance threshold are indicated with larger filled circles and the significance level $P = 9.9 \times 10^{-5}$, $-\log_{10}(P) = 4.00$. 


For obtaining genes near the 5 SNPs identified by GWAS, we determined that LD in Delta Smelt did not extend widely in the genome (Supplementary Document 4). A conservative 26,500 bp range was searched around identified SNPs, except in the case of the 2 SNPs occurring in close proximity on lg02, where a single region was searched but extended by the distance between those 2 SNPs. Nine genes are annotated and named nearby to highly associated SNPs (Table 3). GO names and descriptions are reported in Supplementary Document 4.

**Discussion**

**Summary of results/conclusions**

In this study we examined genotype-phenotype associations with respect to the life history of Delta Smelt by combining otolith Sr isotope geochemistry and genetic sequencing. Delta Smelt exhibit a complex life history known as partial migration where some portion of the population spawn in freshwaters of the Sacramento-San Joaquin Delta and disperse downstream to rear in estuarine habitats within the SFE before returning to spawn in freshwater (semianadromous migrants, MIG); while others complete their life cycle entirely within freshwater habitats, foregoing migration (FWR) (Hobbs et al. 2019). The presence of a genetic foundation for this observed variation in MIG behaviors and adaptive genetic variation relevant to these alternative selective regimes is an important gap in our understanding of the molecular mechanisms underlying life history diversity in this species. Overall, we found no apparent population-level genetic structure in terms of genome-wide nucleotide diversity as has been previously reported (Fisch et al. 2011) (Supplementary Appendix 1), however, we discovered a significant association between phenotypic and genotypic variation, suggesting that life history complexity in Delta Smelt exhibits a heritable genetic foundation, indicative of evolutionary processes that select for diverse traits. Outwardly, the MIG fish were larger than the FWR as measured through fork length, which is in agreement with the increased foraging success of MIG fish (Hammock et al. 2017). Though more work is needed, translation of this adaptive genetic variation into management-relevant tools remains an under-explored and likely valuable option for improving conservation and recovery efforts for this imperiled endemic species.

**Evolutionary origin of Delta Smelt and Delta Smelt life history diversity**

Although the observed life history diversity in Delta Smelt has likely existed for a long time, it remains unclear whether semianadromy has always been favored, or if it is only currently dominant due to recent natural or anthropogenic forcings. The
evolutionary origins of Delta Smelt point to a mid-Pliocene to early Pleistocene divergence from Surf Smelt (H. pretiosus), a widely distributed marine species (Ivlevs and Taylor 2007). A likely scenario is that Delta Smelt diverged through glacial isolation in a freshwater basin in western California, such as in the Pleistocene lakes of the southern San Joaquin Valley (Norriss and Webb 1990). Thus, historically, freshwater inputs from rain, springs, and snowmelt likely provided large expanses of cool freshwater habitats for FWR Delta Smelt, and postglaciation, Delta Smelt were reconnected with the estuary and likely thrived in both freshwater and estuarine habitats.

However, warming temperatures, hydrological alterations, and habitat destruction have dramatically altered Delta Smelt habitats [Cloern et al. 2011; San Francisco Estuary Institute-Aquatic Science Center (SEFIA-ASC) 2014; Hobbs et al. 2017; Hutton et al. 2017]. Contemporary FWR Delta Smelt are now largely reconnected to the Sacramento Deep Water Ship Channel in the northern region of the Sacramento-San Joaquin Delta and are abundant only in the coolest of years; whereas MIG Delta Smelt are found more widely throughout the SFE and are more abundant in years with average and above average temperatures (Hobbs et al. 2019; Lewis et al. 2022). The complex life history of Delta Smelt is likely a result of pre-existing adaptation to a wide variety of habitats historically available in the Sacramento-San Joaquin Delta (Hobbs et al. 2019), and its ancient origin constitutes an important aspect of the species’ biology that is critical for informing the long-term management of the species.

### Genetic association vs causative polymorphisms

Species may respond to heterogeneous environments through local adaptation or phenotypic plasticity. Evidence for population genetic structure and associated neutral genetic divergence indicating local adaptation is lacking in Delta Smelt (Fig. 1b). The alternative view, that phenotypic plasticity is underlying life history variation in Delta Smelt is undercut by our results indicating that there is a genetic association with MIG phenotypes in Delta Smelt that is polygenic (Fig. 2). Two nonexclusive hypotheses may be proposed to explain the observed patterns in Delta Smelt: genotype-dependent habitat choice, and intragenerational spatially varying selection that results in ecotypic differentiation but not large genetic differentiation (e.g. Pavey et al. 2015). These two hypotheses would recreate generationally the observed patterns within Delta Smelt in terms of phenotype and genotype and have been observed in other organisms (e.g. Bourret et al. 2014; Soria-Carrasco et al. 2014; Pavey et al. 2015).

The reduced representation sequence data used in this study are unlikely to sample the causative polymorphisms observed but should be able to sample regions that are in LD with a causative genome region. As a result, the variants identified are likely not causative polymorphisms, but may exhibit some linkage to causative polymorphisms. Evaluation of the functional differences reflected by observed genotypic differences rests on further evaluation with more complete genome coverage and understanding of the Delta Smelt’s genome. Very strong associations and extremely high assignment accuracy are not expected given the data type; however, we did identify statistically significant associations with life history variation and had high classification success. Overall, we did not identify a clear single region of association; rather at least 4 separate chromosomal regions were implicated by GWAS. The strongest association identified on lg02 (site 11230464) also contributed the greatest to the separation of FWR and MIG fish through DAPC (Supplementary Document 3).

As a whole, the genetic signatures of FWR and MIG fish permitted classification with >90% accuracy into phenotypic class with simple algorithms (Ward’s distance clustering, km). Several fish were not categorized correctly (7 FWR individuals and 1 MIG). All of these fish came from the same sampling location where most FWR were sampled (Supplementary Appendix 1). The more geographically dispersed samples of FWR fish; however, were assigned correctly.

### Candidate genes for MIG phenotypes

The functional analysis conducted did identify several genes that may relate to the observed phenotypes. The mostly high-associated SNP on lg02 is located near solute carrier family 6 member 3 (sc6b3), which functions in transmembrane sodium transport (GO: 0035725, Supplementary Document 4). Members of the solute carrier family 6 including sc6b3 have been previously associated with adaptation to salinity variation in three-spine sticklebacks (Gasterosteus aculeatus) as well as a sc6b3 specifically being identified as a candidate gene from freshwater and marine divergence (Hohenlohe et al. 2010; Guo et al. 2015).

The transcription factor nuclear factor of kappa light polypeptide gene enhancer in B-cells 1 (nkb1) is located near a highly associated SNP and is a key regulator of immune response (Oechkinghaus and Ghosh 2009). MIG fishes inhabit different environments and thus are exposed to alternative parasites and pathogens in those alternative habitats. Furthermore, immunity is an energy intensive process and may result in trade-offs with other energy intensive processes such as fecundity and movement (Colgan et al. 2021). Between FWR and MIG phenotypes, immune-related genes may be under strong selective processes. A second transcription factor was also identified in our functional analysis: transcription factor CP2-like 1 (tfcp2l1). This transcription factor is not well-characterized in fishes, however, the mouse ortholog is known to function in the development of kidneys and the diversification of both intercalated and principal cells regulating acid-base and salt-water homeostasis (Werth et al. 2021).
variation through whole-genome sequencing and corroboration of genetic or structural variants more closely linked to life history genetic variation would be best served through the identification brood year (2012). Monitoring of Delta Smelt life history adaptive coverage and the Delta Smelt in this study are all from a single per are useful for these purposes, there are limitations in genome also advisable. Although the genetic variants identified in this pa-

Management implications
The Sacramento-San Joaquin Delta is now unable to sustain large numbers of Delta Smelt, with captive Delta Smelt comprising nearly all existing members of the species (Lessard et al. 2018). As our results indicate both FWR and MIG genetic backgrounds in this population, the preservation of life history-associated vari-
dants during captive propagation is needed in order to maintain the full portfolio of Delta Smelt life histories in the cultured pop-
ulation. Many diadromous species have evolved diverse portfo-
lios of MIG behaviors that allow them to persist within stochastic environments. Partial migration (co-occurring MIG and resident phenotypes), for example, can enhance the stability and resil-
ience of populations to natural and anthropogenic disturbances (Lundberg 1988; Greene et al. 2010) by serving as a bet-hedging strategy that spreads risk and enhances resilience to environ-
mental stochasticity (Roff 1993; Rochet 2000; Schindler et al. 2010; Moore et al. 2014; Hodge et al. 2016; Brennan et al. 2019). Once quantified, this intraspecific phenotypic variation in MIG behav-
iors can be translated into management-relevant tools that are key to effective management and conservation (Schindler et al. 2010). Management practices should account for and incorporate such variation, as focusing on a single life history phenotype may not maximize population stability in the long-term (Hilborn et al. 2003; Greene et al. 2010; Schindler et al. 2010; Brennan et al. 2019).

In addition to monitoring life history associated variants in the hatchery population, continued monitoring of highly associ-
ated variants through the course of the reintroduction effort is also advisable. Although the genetic variants identified in this pa-
per are useful for these purposes, there are limitations in genome coverage and the Delta Smelt in this study are all from a single brood year (2012). Monitoring of Delta Smelt life history adaptive genetic variation would be best served through the identification of genetic or structural variants more closely linked to life history variation through whole-genome sequencing and corroboration across more brood years of Delta Smelt.

Future research
Though informative, here we only examined a single cohort of Delta Smelt. The importance and prevalence of these associa-
tions through time are unknown and genome coverage was not complete. Furthermore, patterns in the relative abundance of each phenotype are likely to covary with changes in environmen-
tal conditions that may exert phenotype-specific effects on sur-

Data availability
Sample metadata and scripts for analysis and generation of fig-
ures in this manuscript are available at https://github.com/ MacCampbell/delta-smelt. Sequence data have been deposited with NCBI into the Sequence Read Archive under BioProject PRJNA834474. Otolith microchemistry profiles and assignments are available as part of the Bureau of Reclamation Directed Outflow Technical Report #3 (https://www.usbr.gov/mp/bdo/di-

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Conflicts of interest
None declared.

Literature cited
Ali OA, O’Rourke SM, Amish SJ, Meek MH, Luikart G, Jeffres C, Miller MR. RAD capture (capture): flexible and efficient sequence-based genotyping. Genetics. 2016;202(2):389–400. doi:10.1534/genet-
ics.115.183665.
Fisch KM, Henderson JM, Burton RS, May B. Population genetics and conservation implications for the endangered delta smelt in the San Francisco Bay-Delta. Conserv Genet. 2011;12(6):1421–1434. doi:10.1007/s10592–011–0240-y.

Gotthard K, Nylin S. Adaptive plasticity and plasticity as an adaptation: a selective review of plasticity in animal morphology and life history. Oikos. 1995;74(1):3–17. doi:10.2307/3545669.

Greene CM, Hall JE, Guilbault KR, Quinn TP. Improved viability of populations with diverse life-history portfolios. Biolett. 2010; 6(3):382–386. doi:10.1098/rstb.2009.0780.

Gross MR. The evolution of diadromy in fishes. Am Fish Soc Symp. 1987;1:14–25.

Gross MR, Coleman RM, McDowell RM. Aquatic productivity and the evolution of diadromous fish migration. Science. 1988;239(4845):1291–1293. doi:10.1126/science.239.4845.1291.

Guo B, DeFaveri J, Sotelo G, Nair A, Merilä J. Population genomic evidence for adaptive differentiation in Baltic Sea three-spined sticklebacks. BMC Evol Biol. 2015;15:19. doi:10.1186/s12862–015–0130-8.

Hammock BG, Slater SB, Baxter RD, Fangue NA, Cochrell D, Hennessy A, Kurobe T, Tai CY, Teh SJ. Foraging and metabolic consequences of semi-anadromy for an endangered estuarine fish. PLoS ONE. 2017;12(3):e0173497. doi:10.1371/journal.pone.0173497.

Hammock BG, Hartman R, Dahlgren RA, Johnston C, Kurobe T, Lehman PW, Lewis LS, Van Nieuwenhuyse E, Ramirez-Duarte WF, Schultz AA, et al. Patterns and predictors of condition indices in a critically endangered fish. Hydrobiologia. 2022;849(3):675–695. doi:10.1007/s10750–021–04738-z.

Hilborn R, Quinn TP, Schindler DE, Rogers DE. Biocomplexity and fisheries sustainability. Proc Natl Acad Sci USA. 2003;100(11):6564–6568. doi:10.1073/pnas.1037274100.

Hobs JA, Lewis LS, Willmes M, Denney C, Bush E. Complex life histories discovered in a critically endangered fish. Sci Rep. 2019,9(1):16772. doi:10.1038/s41598–019–52273–8.

Hobs JA, Moyle PB, Fangue N, Connon RE. Is extinction inevitable for Delta Smelt and Longfin Smelt? An opinion and recommendations for recovery. San Franc Estuay Watershed Sci. 2017;15(2):1–19. doi:10.15447/sfews.2017v15iss2art2.

Hodge BW, Wilzbach MA, Duffy WG, Quinones RM, Hobs JA. Life history diversity in Klamath River steelhead. Trans Am Fish Soc. 2016;145(2):227–238.

Hohenlohe PA, Bassham S, Etter PD, Stiffler N, Johnson EA, Cresko WA. Population genomics of parallel adaptation in threespine stickleback using sequenced RAD tags. PLoS Genet. 2010;6(2):e1000862. doi:10.1371/journal.pgen.1000862.

Huang K, Andrew RL, Owens GL, Ostevik KL, Rieseberg LH. Multiple chromosomal inversions contribute to adaptive divergence of a dune sunflower ecotype. Mol Ecol. 2020;29(14):2535–2549. doi:10.1111/mec.15428.

Hutton PH, Rath JS, Roy SB. Freshwater flow to the San Francisco Bay-Delta estuary over nine decades (Part 2): change attribution. Hydro Process. 2017;31(14):2516–2529.

Ilves KL, Taylor EB. Evolutionary and biogeographical patterns within the smelt genus Hypomesus in the North Pacific Ocean. J Biogeogr. 2007;35:48–64. doi:10.1111/j.1365–2699.2007.01782.x.

Jassby AD, Kimmerer WJ, Monismith SG, Armor C, Colenn JE, Powell TM, Schubel JR, Vendilinskt T. Isosalinie position as a habitat indicator for estuarine populations. Ecol Appl. 1995;5(1):272–289. doi:10.1111/j.1051–0761.1995.00181.x.

Jeffries KM, Connon RE, Davis BE, Komoroske LM, Britton MT, Sommer T, Toegham AE, Fangue NA. Effects of high temperatures on threatened estuarine fishes during periods of extreme drought. J Exp Biol. 2016;219(Pt 11):1705–1716. doi:10.1242/jeb.134528.
Soria-Carrasco V, Gompert Z, Comeault AA, Farkas TE, Parchman TL, Johnston JS, Buerkle CA, Feder JL, Bast J, Schwander T, et al. Stick insect genomes reveal natural selection’s role in parallel speciation. Science. 2014;344(6185):738–742. doi:10.1126/science.1252136.

Swanson C, Reid T, Young PS, Cech JJ. Comparative environmental tolerances of threatened delta smelt (Hypomesus transpacificus) and introduced wakasagi (H. nipponensis) in an altered California estuary. Oecologia. 2000;123(3):384–390. doi:10.1007/s004420051025.

Thompson NF, Anderson EC, Clemento AJ, Campbell MA, Pearse DE, Hearsey JW, Kinziger AP, Garza JG. A complex phenotype in salmon controlled by a simple change in migratory timing. Science. 2020;370(6516):609–613. doi:10.1126/science.aba9059.

Thorneycroft HB. A cytogenetic study of the White-Throated Sparrow, Zonotrichia albicollis (Gmelin). Evolution. 1975;29(4):611–621. doi:10.1111/j.1558-5646.1975.tb00853.x.

Tinbergen N. On aims and methods of ethology. Zeitschrift Tierpsychol. 1963;20(4):410–433.

Tuttle EM, Bergland AO, Korody ML, Brewer MS, Newhouse DJ, Minx P, Stager M, Betuel A, Cheviron ZA, Warren WC, et al. Divergence and functional degradation of a sex chromosome-like supergene. Curr Biol. 2016;26(3):344–350. doi:10.1016/j.cub.2015.11.069.

Wacholder S, Chanock S, Garcia-Closas M, El ghormli L, Rothman N. Assessing the probability that a positive report is false: an approach for molecular epidemiology studies. J Natl Cancer Inst. 2004;96(6):434–442. doi:10.1093/jnci/djh075.

Werth M, Schmidt-Ott KM, Leete T, Qiu A, Hinze C, Viltard M, Paragas N, Shawber CJ, Yu W, Lee P, et al. Transcription factor TFCP2L1 patterns cells in the mouse kidney collecting ducts. eLife. 2017;6:e24265. doi:10.7554/eLife.24265.

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