

FEATURE

# Ancestry and Adaptation of Rainbow Trout in Yosemite National Park

**Devon E. Pearse\*** and **Matthew A. Campbell** | National Marine Fisheries Service, Southwest Fisheries Science Center, Fisheries Ecology Division, 110 McAllister Way, Santa Cruz, CA 95060 | Ecology and Evolutionary Biology, UCSC/Coastal Biology Building, 130 McAllister Way, Santa Cruz, CA 95060.

\*Corresponding author e-mail: [devon.pearse@noaa.gov](mailto:devon.pearse@noaa.gov)

Photo credit: Tom Koerner/USFWS

California's Central Valley contains an abundance of rivers with historical and potential productivity for anadromous salmonids, which are currently limited by impacts such as dams, water diversions, and high temperatures. We surveyed genetic variation in Rainbow Trout *Oncorhynchus mykiss* within the upper Tuolumne and Merced rivers in and around Yosemite National Park to evaluate both population origins (ancestry) and the evolutionary response to natural and artificial barriers to migration (adaptation). This analysis revealed that despite extensive stocking with hatchery Rainbow Trout strains throughout the study area, most populations retained largely indigenous ancestry. Adaptive genomic variation associated with anadromy was distributed throughout the study area, with higher frequencies observed in populations connected to reservoirs that are known to support adfluvial life history variants. Fish in southern Central Valley rivers experience temperatures near the upper thermal limit for salmonids and represent an important reservoir of genomic diversity for adaptation to climate change. These results highlight the importance of local adaptation as well as the potential for resident Rainbow Trout populations above barrier dams to contribute to the recovery of steelhead (anadromous Rainbow Trout) once migratory connectivity is restored between upstream spawning and rearing habitats and the ocean.

The Central Valley of California is both a productive agricultural region and an important ecosystem in western North America that encompasses two large river systems: the Sacramento River to the north and the San Joaquin River to the south. Together, these rivers and their tributaries are home to the southernmost native populations of Chinook Salmon *Oncorhynchus tshawytscha*, Rainbow Trout *O. mykiss*, and steelhead (anadromous Rainbow Trout; Fisher 1994; Busby et al. 1996; Yoshiyama et al. 1998, 2001; McEwan 2001). However, the construction of barrier dams and water diversions has greatly restricted migratory connectivity on many rivers, resulting in extremely reduced anadromous salmonid populations throughout the Central Valley (Yoshiyama et al. 1998, 2001; May and Brown 2002; Lindley et al. 2006; NMFS 2006, 2014; Katz et al. 2013). Large barrier dams in particular prevent upstream migration of adult salmonids to their spawning habitats, as well as preventing downstream juvenile migration, thus severely impacting anadromous species. As a result, there is an increasing focus on re-connecting migratory anadromous salmonid populations with their historical spawning grounds above impassable dams through dam removal, addition of volitional passageways, or fish passage programs known as “trap and haul” (Anderson et al. 2014; NMFS 2014; Lusardi and Moyle 2017).

The Tuolumne and Merced rivers are tributaries of the San Joaquin River that drain a large portion of Yosemite National Park (hereafter, “Yosemite”) in the central Sierra Nevada as well as Stanislaus National Forest lands and other lands (Figure 1). Both spring-run Chinook Salmon and steelhead historically used these waterways to access the cool refuges of the High Sierra, where they and their offspring could escape the summertime heat and dwindling river flows of the lower elevations. Both species likely spawned in the Merced River throughout Yosemite Valley up to the bases of Half Dome and the spectacular Vernal and Yosemite falls and in the Tuolumne River up to Preston Falls, just downstream of the park's boundary (Figure 1; Tuolumne River above Preston Falls [TAPF]). However, the full extent of their historical migrations is unclear (Yoshiyama et al. 2001; Lindley et al. 2006). Beginning in the mid-1800s, as happened in many Central Valley rivers, construction of a series of dams blocked anadromous salmonids' access between the ocean and the headwaters of the Tuolumne and Merced rivers. Currently, La Grange Dam (completed in 1883) and Crocker-Huffman Dam (completed in 1906) create the upper limits to anadromous migration; above these dams, the much-larger New Don Pedro and New Exchequer dams form major reservoirs on the Tuolumne and Merced rivers,

respectively, providing flood control, water storage, recreation, and power generation (Figure 1). Collectively, these dams and their predecessors have prevented native salmon and steelhead from accessing historic spawning habitats for more than a century. However, even prior to the construction of the major barrier dams, the activities of the California Gold Rush in the 1850s and subsequent development of agricultural infrastructure in the Central Valley had a huge effect on the native fauna, particularly the migratory salmonids (Yoshiyama et al. 1998, 2001), and few naturally spawning anadromous salmonids exist in the reaches below these dams today (Ford and Kirihaara 2010; Cuthbert et al. 2012; NMFS 2014). This situation is further exacerbated by the poor quality of downstream habitat in the Sacramento–San Joaquin Delta for both migratory and nonmigratory native fishes (Moyle et al. 2018). Today, intensive management and hatchery supplementation maintain many salmonid populations in the Central Valley, including the California Central Valley Distinct Population Segment (DPS) of steelhead, which is listed as threatened under the Endangered Species Act (ESA; NMFS 2006); however, inability to access more than 80% of its historical spawning habitat remains a critical issue for the recovery of this DPS (Yoshiyama et al. 2001; Lindley et al. 2006). The National Marine Fisheries Service's (NMFS) Central Valley Recovery Plan identifies the upper Tuolumne River (UTR) and the upper Merced River (UMR) as candidate areas for reintroduction of both steelhead and spring-run Chinook Salmon to support recovery of the southern Sierra Nevada steelhead diversity group through upstream passage of adults and downstream movement of juveniles over the dams (NMFS 2014).

With few exceptions, Chinook Salmon are strictly anadromous (but see Sard et al. 2016; Brenkman et al. 2017), whereas self-sustaining populations of freshwater resident Rainbow Trout commonly persist above barrier dams that block their ability to access the ocean (Kendall et al. 2015). Individuals in above-dam populations may exhibit several life history strategies, including a migratory adfluvial life history utilizing a reservoir as an alternative to a fully anadromous marine migration and returning to spawn in upstream tributaries (e.g., Holecek and Scarnecchia 2013; Leitwein et al. 2017). These populations are typically closely related to the remaining *O. mykiss* found below barriers in the same watershed (Narum et al. 2008; Clemento et al. 2009; but see Pearse and Garza 2015), although stocking of nonnative hatchery Rainbow Trout strains into above-barrier habitats has resulted in partial or complete replacement of the indigenous ancestry in some cases (e.g., Abadía-Cardoso et al. 2016). Importantly, only the anadromous (steelhead) life history is listed under

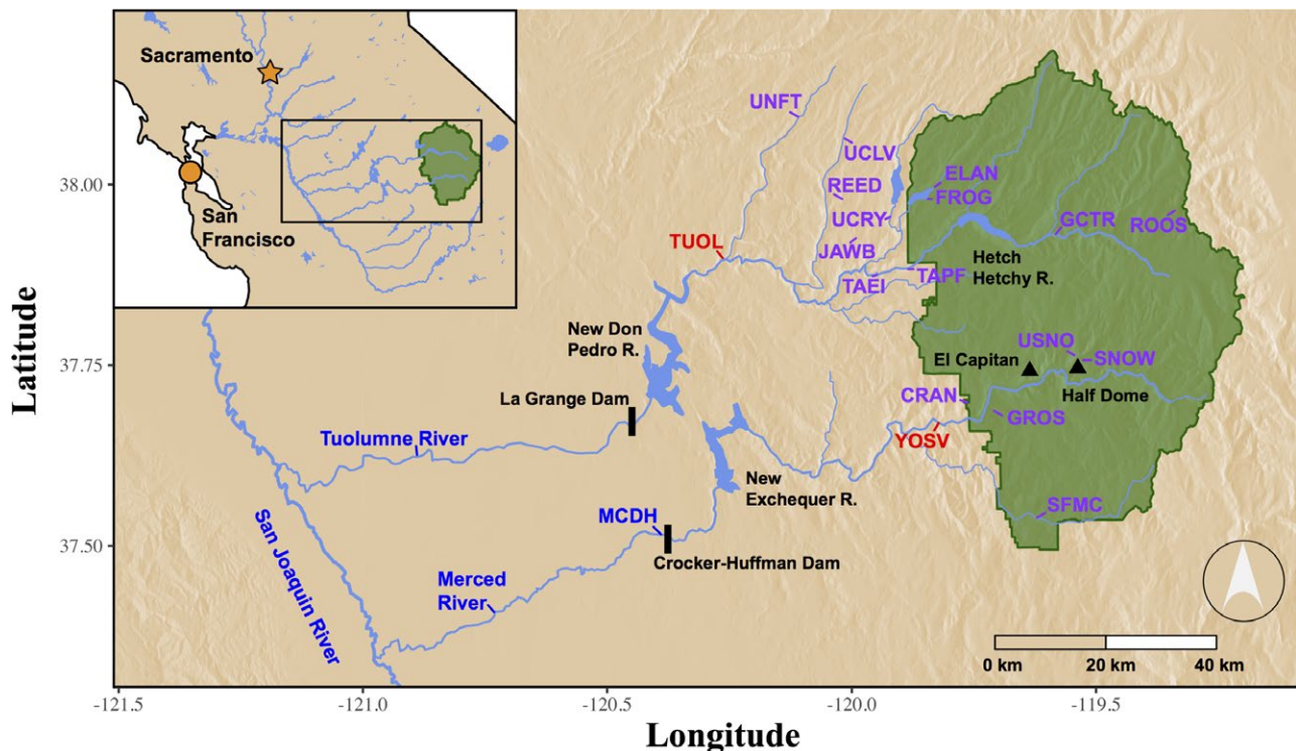


Figure 1. Map of Tuolumne and Merced river sampling locations investigated in this study, showing existing dams and reservoirs. Sampling units as described in the text are labeled following the codes in Table 1; two Pearse and Garza (2015) reference populations are indicated by “Tuolumne River” and “Merced River.” Yosemite National Park is shaded green, with El Capitan and Half Dome indicated by solid black triangles. Population migratory potentials are indicated by color: potentially anadromous (blue), potentially adfluvial (red), and resident Rainbow Trout (purple). La Grange and Crocker-Huffman dams (black bars) serve as the upper limits to anadromy in the Tuolumne and Merced rivers, respectively. Inset depicts the central California region, showing the Sacramento–San Joaquin River system draining to the Pacific Ocean through San Francisco Bay. The inner box in the inset indicates the geographic extent of the main map.

the ESA, whereas even closely related above-barrier Rainbow Trout populations are not protected by the ESA (NMFS 2006). Thus, in considering efforts to reconnect migratory steelhead populations below dams with their historical upriver spawning habitats, an important first step is to evaluate the genetic ancestry and adaptive potential of the Rainbow Trout trapped above the dams (Winans et al. 2010, 2017, 2018).

#### ANCESTRY OF YOSEMITE TROUT

There is a rich history of fish stocking in and around Yosemite that has undoubtedly influenced the distribution and genetic composition of its Rainbow Trout. Early visitors took a strong interest in increasing the trout populations, both for food resources and recreation (Caton 1869; Pavlik 1987). Fish planting likely began in the area during the 1870s, initially by settlers moving locally captured fish up into the previously fishless waters above waterfalls and in high alpine lakes. Stocking records describing imported trout first occurred in the 1890s, and by 1895, there was a fish hatchery operating on the South Fork Merced River at Wawona that distributed both indigenous and imported trout throughout the area (Pavlik 1987). A subsequent hatchery was established in 1918 at Happy Isles on the main-stem Merced River in Yosemite Valley, and the importation of eggs from other hatcheries ensured a steady supply of Rainbow Trout as well as nonnative species, such as Lahontan Cutthroat Trout *O. clarkii henshawi*, European Brown Trout *Salmo trutta*, and Brook Trout *Salvelinus fontinalis* (Leitritz 1970). Although

many of the eggs reared at the Happy Isles Fish Hatchery were imported from outside Yosemite, some were collected at an egg taking station on Frog Creek (FROG), a historically fishless tributary of the UTR above Lake Eleanor in the northern part of Yosemite (Figure 1; Pavlik 1987). Thus, over the years, a diverse mixture of both locally sourced Rainbow Trout and fish imported from throughout California have been planted within Yosemite, potentially creating admixed populations with both indigenous and hatchery ancestry. However, management of more recent stocking efforts has changed significantly, and since 2013, most trout planted in California have been sterile triploids, limiting further naturalization and spawning by hatchery fish. The current distribution of Rainbow Trout within Yosemite is therefore composed of self-sustaining populations whose ancestry remains to be evaluated through genetic analysis.

#### ADAPTATION TO RESIDENCY

Despite dramatic differences in traits related to physiology, morphology, and behavior, the diverse life history forms of *O. mykiss* often co-exist and interbreed, forming interrelated populations in nature (Quinn 2011). Consequently, anadromous and resident fish within a drainage basin are typically closely related to each other (Olsen et al. 2006; Narum et al. 2008; Pearse et al. 2009). Although offspring of a particular life history variant may take on an alternative strategy from that of their parents (Courter et al. 2013), there is a great deal of evidence pointing to heritable influences on life

history strategies and associated phenotypes (e.g., Neave 1944; Berejikian et al. 2014; Phillis et al. 2016).

Surveys of genetic variation have found that Rainbow Trout in above-barrier habitats undergo specific genetic changes as populations adapt to residency. In particular, one region of chromosome Omy5 has shown a consistent association with resident (R) and anadromous (A) life histories, although many other genomic regions are also associated with variation in this trait (e.g., Nichols et al. 2008; Hale et al. 2013; Hecht et al. 2013). However, unlike waterfalls, which exert knife-edge selection against downstream migration (Pearse et al. 2009; Northcote 2010), barrier dams create reservoirs above them, allowing Rainbow Trout trapped above the dams to develop an adfluvial migratory life history by utilizing the reservoir as a rearing habitat and spawning in the tributary streams (e.g., Holecek et al. 2012; Holecek and Scarnecchia 2013). Importantly, despite the dramatic difference in osmotic conditions between reservoirs and the ocean, selection for an adfluvial migratory life history appears to affect the same adaptive genomic variants on Omy5 as true anadromous migrations (Pearse et al. 2014; Leitwein et al. 2017). This suggests that adfluvial Rainbow Trout populations isolated above dams and reservoirs could potentially contribute to the recovery of migratory anadromous ecotypes once migratory access to the ocean is restored through dam removal or assisted fish passage (Thrower et al. 2008; Meek et al. 2014; Winans et al. 2017).

The primary goal of this study was to determine the genetic ancestry and current population structure of *O. mykiss* populations in the UTR and UMR (Figure 1). To do so, we investigated the genetic relationships of *O. mykiss* in these rivers relative to (1) other populations above and below barriers to anadromy in the Central Valley, (2) hatchery Rainbow Trout strains commonly used in California, and (3) coastal steelhead populations. In addition, we assayed adaptive genomic variation in the region of chromosome Omy5 known to be associated with anadromous and adfluvial life history traits in *O. mykiss* to estimate the frequencies of alleles associated with migratory behavior relative to the presence of barriers to fish migration (Pearse et al. 2014; Apgar et al. 2017; Leitwein et al. 2017). We use this information to evaluate the potential for UTR and UMR populations to contribute to the recovery of anadromous steelhead below barriers in the southern Central Valley. Together, these data provide a baseline to inform future management of *O. mykiss* populations in these and other Central Valley watersheds and to improve our understanding of the potential to recover anadromous steelhead populations by restoring connectivity with *O. mykiss* populations trapped in habitats upstream of the dams.

## METHODS

### Sampling

Fish were captured in 2015 and 2016 at 37 sites throughout the UTR and UMR watersheds, including both migratory reaches (those historically accessible to migratory steelhead; Lindley et al. 2006) and historically fishless reaches that were isolated above barriers (Figure 1; Table 1). Due to the difficulty of accessing fish in larger rivers as well as the extremely low conductivity of Sierra Nevada streams, many sites were unsuitable for electrofishing, and most fish in the study were captured by hook and line. This “Fly Fishing for Science” had the added benefit of providing a mechanism to allow volunteer fly fishers to contribute to the project as

citizen scientists (Williams et al. 2015). All fish were measured, and fin tissue samples were taken from each individual prior to release at the site of capture. Tissue samples were dried and taken to the NMFS laboratory in Santa Cruz, California, for analysis.

### Genetic data collection

We extracted DNA from dried fin clips by using the DNeasy 96 filter-based nucleic acid extraction system on a BioRobot 3000 (Qiagen, Inc.) in accordance with the manufacturer’s protocols. All DNA extractions were diluted 2:1 with distilled water and used for PCR pre-amplification prior to TaqMan or SNP Type (single-nucleotide polymorphism [SNP]) genotyping with 96.96 Integrated Fluidic Circuit chips (Fluidigm, Inc.). Genotypes were read and scored using Fluidigm SNP Genotyping Analysis software (Fluidigm). All samples were genotyped at total of 92 SNPs for population genetic analysis following Abadía-Cardoso et al. (2013), a gender identification SNP assay (Brunelli et al. 2008; Rundio et al. 2012), and three SNPs on chromosome Omy5 that have been associated with migratory life history traits (Pearse et al. 2014; Pearse and Garza 2015; Abadía-Cardoso et al. 2016; Leitwein et al. 2017).

### Data analysis

The SNP genotype data were combined with published data from 21 representative wild coastal and Central Valley *O. mykiss* populations, three Central Valley steelhead hatcheries, and five hatchery Rainbow Trout strains common in California (Pearse and Garza 2015). The genetic data were analyzed with R version 3.4.1 (R Development Core Team 2017). Genotypes were imported for use in R and converted to a “genind” object for subsequent analyses through the package “pegas” version 0.10 (Paradis 2010). Quality control of individual fish was undertaken with the “missingno” function of “poppr” version 2.5.0 (Kamvar et al. 2014) by specifying that both genotypes and loci were not allowed to have more than 5% missing data. From these filtered data, two separate approaches were implemented: (1) an individual approach; and (2) a population approach in which fish sampled along contiguous reaches without barriers to migration were combined into “sampling units,” resulting in a total of 20 discrete groups of individuals based on local geography and barriers to migration (Figure 1; Table 1).

### Individual approach

For the individual approach, prior population assignment based on collection location was not considered, and individuals were independently assigned to inferred populations. This approach was used to verify the independence or interrelatedness of sampling locations. For example, when hatchery reference populations are included, do sampled individuals show genetic similarities to any hatchery population? We used discriminant analysis of principal components (DAPC) as implemented in R with the package “adegenet” version 2.0.1 (Jombart 2008; Jombart et al. 2010) and STRUCTURE version 2.3.4 (Pritchard et al. 2000; Falush et al. 2003) as complementary individual analyses.

For the individual DAPC, we limited our analysis to only new collections from the Tuolumne River and Merced River along with five hatchery trout strains as reference populations to detect hatchery introgression. The DAPC is not based on a

Table 1. Sample information and summary statistics for genetic data analysis of *Oncorhynchus mykiss*. The full name, code, and sample size for each sampling unit are given along with the categorization regarding migratory potential. For each population, the sample size included in genetic analyses ( $n$ ; i.e., samples passing quality control), the expected heterozygosity ( $H_e$ ), and the frequency of the anadromous-type *Omy5* migration-associated region (F[A MAR]) are provided. For each collection that comprised a sampling site, the major drainage basin, dates of collection, sample size ( $N$ ), and World Geodetic System 84 coordinates are indicated.

| Sampling location                  | Code | Total N | Population type         | $N$ | $H_e$ | F(A MAR) | Drainage basin | Date(s)                    | $N$ | Latitude | Longitude |
|------------------------------------|------|---------|-------------------------|-----|-------|----------|----------------|----------------------------|-----|----------|-----------|
| Upper North Fork Tuolumne River    | UNFT | 24      | Above barrier           | 21  | 0.27  | 0.08     | Tuolumne River | Jun 8, 2015                | 24  | 38.10    | -120.11   |
| Tuolumne River                     | TUOL | 150     | Historically anadromous | 145 | 0.36  | 0.31     | Tuolumne River | May 12, 2015               | 11  | 37.90    | -120.07   |
|                                    |      |         |                         |     |       |          | Tuolumne River | Aug 27, 2015               | 3   | 37.90    | -120.26   |
|                                    |      |         |                         |     |       |          | Tuolumne River | Oct 8, 2015                | 41  | 37.84    | -120.06   |
|                                    |      |         |                         |     |       |          | Tuolumne River | May 12, 2015               | 31  | 37.84    | -120.04   |
|                                    |      |         |                         |     |       |          | Tuolumne River | Jun 9, 2015                | 11  | 37.89    | -119.97   |
|                                    |      |         |                         |     |       |          | Tuolumne River | May 15, 2015               | 36  | 37.89    | -119.95   |
|                                    |      |         |                         |     |       |          | Tuolumne River | May 14, 2015               | 17  | 37.88    | -119.97   |
| Upper Clavey River                 | UCLV | 131     | Above barrier           | 129 | 0.30  | 0.08     | Tuolumne River | Jun 9, 2015                | 68  | 37.99    | -120.05   |
|                                    |      |         |                         |     |       |          | Tuolumne River | Jun 8, 2015                | 55  | 38.07    | -120.01   |
|                                    |      |         |                         |     |       |          | Tuolumne River | Jun 8, 2015                | 8   | 38.09    | -120.01   |
| Reed Creek                         | REED | 103     | Above barrier           | 102 | 0.36  | 0.08     | Tuolumne River | May 13, 2015               | 103 | 37.98    | -120.02   |
| Jawbone Creek                      | JAWB | 59      | Above barrier           | 57  | 0.35  | 0.12     | Tuolumne River | May 13, 2015               | 59  | 37.93    | -119.99   |
| Upper Cherry Creek                 | UCRY | 5       | Above barrier           | 5   | 0.31  | NA       | Tuolumne River | Jun 9, 2015                | 5   | 37.96    | -119.92   |
| Eleanor Creek                      | ELAN | 12      | Above barrier           | 12  | 0.30  | 0.08     | Tuolumne River | Jun 19, 2016               | 12  | 38.00    | -119.83   |
| Frog Creek                         | FROG | 25      | Above barrier           | 25  | 0.29  | 0.24     | Tuolumne River | Jun 18, 2016               | 25  | 37.98    | -119.84   |
| Tuolumne River-above Early Intake  | TAEI | 45      | Historically anadromous | 42  | 0.36  | 0.20     | Tuolumne River | May 13, 2015               | 45  | 37.88    | -119.94   |
| Tuolumne River-above Preston Falls | TAPF | 28      | Above barrier           | 23  | 0.32  | 0.20     | Tuolumne River | Jun 10, 2015               | 23  | 37.88    | -119.88   |
|                                    |      |         |                         |     |       |          | Tuolumne River | Jun 11, 2015               | 5   | 37.95    | -119.79   |
| Grand Canyon Tuolumne River        | GCTR | 22      | Above barrier           | 21  | 0.34  | 0.07     | Tuolumne River | Jul 18, 2015               | 22  | 37.93    | -119.58   |
| Roosevelt Lake                     | ROOS | 26      | Above barrier           | 26  | 0.32  | 0.04     | Tuolumne River | Aug 13, 2015               | 26  | 37.96    | -119.34   |
| Merced River Ranch                 | MCRR | 1       | Ocean accessible        | 1   | 0.39  | NA       | Merced River   | Apr 9, 2010                | 1   | 37.52    | -120.40   |
| Merced River Hatchery              | MCDH | 58      | Ocean accessible        | 58  | 0.35  | 0.60     | Merced River   | Dec 9, 2014                | 58  | 37.52    | -120.37   |
| Grouse Creek                       | GROS | 34      | Above barrier           | 33  | 0.23  | 0.00     | Merced River   | Aug 11, 2016               | 34  | 37.69    | -119.70   |
| Crane Creek                        | CRAN | 27      | Above barrier           | 27  | 0.32  | 0.04     | Merced River   | Aug 10, 2016               | 27  | 37.70    | -119.76   |
| South Fork Merced River            | SFMC | 49      | Historically anadromous | 45  | 0.34  | 0.09     | Merced River   | Aug 11, 2016               | 4   | 37.55    | -119.63   |
|                                    |      |         |                         |     |       |          | Merced River   | Aug 11, 2016               | 43  | 37.54    | -119.62   |
|                                    |      |         |                         |     |       |          | Merced River   | Aug 11, 2016               | 2   | 37.52    | -119.66   |
| Yosemite Valley                    | YOSV | 85      | Historically anadromous | 81  | 0.35  | 0.19     | Merced River   | Aug 11, 2016               | 3   | 37.67    | -119.82   |
|                                    |      |         |                         |     |       |          | Merced River   | Jun 30, 2015; Aug 12, 2016 | 47  | 37.68    | -119.74   |
|                                    |      |         |                         |     |       |          | Merced River   | Jun 29, 2016               | 2   | 37.72    | -119.71   |
|                                    |      |         |                         |     |       |          | Merced River   | Aug 11, 2016               | 4   | 37.72    | -119.71   |
|                                    |      |         |                         |     |       |          | Merced River   | Aug 11, 2016               | 15  | 37.72    | -119.56   |
|                                    |      |         |                         |     |       |          | Merced River   | Jun 30, 2015; Aug 10, 2016 | 14  | 37.76    | -119.54   |

(continued)

Table 1. (Continued)

| Sampling location                                    | Code | Total N | Population type         | N   | $H_E$ | F(A MAR) | Drainage basin | Date(s)      | N | Latitude | Longitude |
|--|------|---------|-------------------------|-----|-------|----------|----------------|--------------|---|----------|-----------|
| Snow Creek   | SNOW | 8       | Historically anadromous | 8   | 0.33  | NA       | Merced River   | Aug 10, 2016 | 8 | 37.76    | -119.53   |
| Upper Snow Creek                                     | USNO | 5       | Above barrier           | 5   | 0.34  | NA       | Merced River   | Aug 10, 2016 | 5 | 37.77    | -119.54   |
| <b>Pearse and Garza (2015) Reference Populations</b> |      |         |                         |     |       |          |                |              |   |          |           |
| <i>North Coast</i>                                   |      |         |                         |     |       |          |                |              |   |          |           |
| Mad River  |      | 31      | Ocean accessible        | 31  | 0.35  |          |                |              |   |          |           |
| Eel River (Hollow Tree Creek)                        |      | 28      | Ocean accessible        | 28  | 0.34  |          |                |              |   |          |           |
| Gualala River  |      | 29      | Ocean accessible        | 29  | 0.40  |          |                |              |   |          |           |
| <i>Central Valley</i>                                |      |         |                         |     |       |          |                |              |   |          |           |
| McCloud River (Butcherknife Creek)                   |      | 21      | Historically anadromous | 21  | 0.17  |          |                |              |   |          |           |
| McCloud River (Claiborne Creek)                      |      | 33      | Historically anadromous | 30  | 0.33  |          |                |              |   |          |           |
| Clear Creek  |      | 94      | Ocean accessible        | 86  | 0.34  |          |                |              |   |          |           |
| Deer Creek   |      | 45      | Ocean accessible        | 41  | 0.37  |          |                |              |   |          |           |
| Feather River (Chips Creek)                          |      | 31      | Historically anadromous | 31  | 0.32  |          |                |              |   |          |           |
| Feather River Hatchery                               |      | 30      | Ocean accessible        | 28  | 0.37  |          |                |              |   |          |           |
| Yuba River (Pauley Creek)                            |      | 25      | Historically anadromous | 25  | 0.31  |          |                |              |   |          |           |
| Yuba River   |      | 90      | Historically anadromous | 82  | 0.41  |          |                |              |   |          |           |
| American River (Middle Fork)                         |      | 58      | Historically anadromous | 54  | 0.37  |          |                |              |   |          |           |
| Nimbus Hatchery                                      |      | 98      | Ocean accessible        | 92  | 0.40  |          |                |              |   |          |           |
| Mokelumne Hatchery                                   |      | 162     | Ocean accessible        | 159 | 0.37  |          |                |              |   |          |           |
| Mokelumne River (North Fork)                         |      | 51      | Historically anadromous | 51  | 0.33  |          |                |              |   |          |           |
| Calaveras River                                      |      | 47      | Ocean accessible        | 44  | 0.37  |          |                |              |   |          |           |
| Stanislaus River (upper)                             |      | 52      | Ocean accessible        | 51  | 0.34  |          |                |              |   |          |           |
| Tuolumne River                                       |      | 112     | Ocean accessible        | 106 | 0.39  |          |                |              |   |          |           |
| Tuolumne River (upper)                               |      | 47      | Historically anadromous | 47  | 0.34  |          |                |              |   |          |           |
| Merced River (upper)                                 |      | 35      | Historically anadromous | 35  | 0.35  |          |                |              |   |          |           |
| Merced River   |      | 83      | Ocean accessible        | 81  | 0.29  |          |                |              |   |          |           |
| Mill Flat Creek                                      |      | 26      | Historically anadromous | 26  | 0.36  |          |                |              |   |          |           |
| <i>South Coast</i>                                   |      |         |                         |     |       |          |                |              |   |          |           |
| San Francisquito Creek                               |      | 24      | Ocean accessible        | 24  | 0.36  |          |                |              |   |          |           |
| San Lorenzo River                                    |      | 32      | Ocean accessible        | 32  | 0.37  |          |                |              |   |          |           |
| <i>Hatchery Trout Strains</i>                        |      |         |                         |     |       |          |                |              |   |          |           |
| Kamloops   |      | 47      | NA                      | 47  | 0.23  |          |                |              |   |          |           |
| Mt. Shasta   |      | 92      | NA                      | 83  | 0.32  |          |                |              |   |          |           |
| Eagle  |      | 47      | NA                      | 47  | 0.25  |          |                |              |   |          |           |
| Coleman  |      | 47      | NA                      | 47  | 0.33  |          |                |              |   |          |           |
| Moccasin   |      | 47      | NA                      | 46  | 0.25  |          |                |              |   |          |           |

population genetic model; it relies on the conversion of SNP data into principal components to account for linkage between SNPs and allow generic methods of individual clustering to be used. As opposed to finding axes of maximal variation (e.g., using principal components analysis [PCA]), DAPC maximizes between-population separations and minimizes within-population variation. We identified inferred populations with DAPC by applying the “find.clusters” function of adegenet followed by PCA and discriminant analysis within the “dapc” function of adegenet that utilized the packages “ade4” version 2.7-8 (Chessel and Dufour 2004; Dray and Dufour 2007; Dray et al. 2007) and Modern Applied Statistics with S version 7.3-47 (Venables and Ripley 2002).

Unlike DAPC, STRUCTURE has an explicit population genetics model and uses the individual genotype data directly. STRUCTURE assigns fractional ancestry ( $q$ -values) to  $K$  inferred populations based on descent from a common ancestral population. For each individual, the  $q$ -values sum to 1.00 and indicate what proportion of each of the  $K$  inferred populations makes up the individual. Migrants and individuals of mixed ancestry can be identified with STRUCTURE without a priori designation of defined populations (Pritchard et al. 2000). We evaluated all individuals in the quality-controlled data set with  $K = \{1, \dots, 12\}$  inferred populations in four independent runs with an initial burn-in of 100,000 steps followed by 1,000,000 Markov-chain Monte Carlo steps. For most parameters, default settings were used. Results from the STRUCTURE runs were visualized with DISTRUCT version 1.1 (Rosenberg 2004).

### Population approach

Sample units were treated as populations for identifying population genetic and phylogenetic relationships, with a minimum required size of 10 individuals per sample unit (Table 1). Population genetic relationships were evaluated with DAPC using the sample unit to predefine population genetic clusters. The same sample units were also evaluated in a neighbor-joining population tree generated through poppr version 2.4.1 by using a chord distance metric (Cavalli-Sforza and Edwards 1967) and the filling of missing data by the mean of that locus. Confidence in nodes of the population tree was assessed by 1,000 bootstrap replicates.

### Signatures of migratory adaptation

Of the three genotyped SNP loci located on chromosome Omy5, one (R04944) is known to accurately identify the R and A haplotypes surveyed in previous studies (Pearse et al. 2014; Leitwein et al. 2017). Based on this locus, we calculated the frequencies of the A allele associated with migratory behavior in all populations. These data were then considered with respect to the migratory potential of each sampling site relative to historical and current barriers and reservoirs, and populations with potential for adfluvial life history variants were identified.

## RESULTS

### Sample Genotyping and Population Statistics

Overall, 897 *O. mykiss* samples from the UTR and UMR were genotyped, and after filtering for missing data and loci under selection linked to the Omy5 inversion, the combined data set of 20 sampling units and 29 reference populations consisted of 2,370 individuals and 88 bi-allelic SNP loci (Table 1). Sample sizes for the UTR and UMR populations

ranged from 2 to 103 individuals per site; sample units smaller than 10 were used for individual-based analyses but were excluded from population-level analysis.

The distribution of neutral genetic diversity among populations showed typical patterns, with most populations isolated above barriers having reduced heterozygosity relative to downstream populations (Table 1). Similarly, most hatchery Rainbow Trout strains had reduced levels of variation, as did populations that were inferred to be of primarily hatchery origin (e.g., Grouse Creek [GROS]). Conversely, larger populations connected by migration (e.g., Tuolumne River [TUOL] and Yosemite Valley [YOSV]) tended to have high levels of heterozygosity, similar to coastal and Central Valley steelhead populations (Table 1; Figure 1).

### Individual Approach

The DAPC of individuals from the 20 UTR and UMR sampling units plus 5 hatchery reference strains supported the inference of eight genetic groups (Figure 2). Three of these inferred clusters constituted fish of natural genetic origin, while the other five contained fish of hatchery origin or individuals with a genetic composition similar to that of hatchery fish (Figure 2). Most fish from the UTR and UMR sampling locations were not placed in clusters with significant hatchery trout contributions. However, the GROS sampling location was entirely placed in inferred group 5 along with the Kamloops Hatchery strain, while many individuals from the upper North Fork Tuolumne River (UNFT) sample were grouped with the Coleman Hatchery strain (group 4; Figure 2). Similarly, most individuals from the Merced River Hatchery (MCDH) sample were placed in group 4 with the Coleman strain, while the

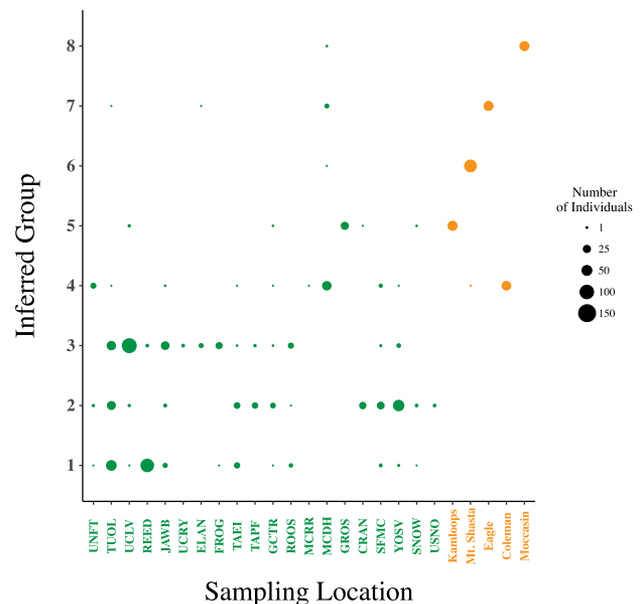


Figure 2. Individual group assignments from discriminant analysis of principal components for individual *Oncorhynchus mykiss* from the 20 upper Tuolumne River/upper Merced River sample units (Yosemite National Park; green) and five hatchery reference strains (orange). The sampling unit is indicated on the x-axis (codes are defined in Table 1), and the hypothesized eight genetic groups of individuals are indicated on the y-axis. Circle sizes indicate the number of individuals from each sampling location that were assigned to a particular genetic group.

rest were grouped with the Mt. Shasta, Eagle, and Moccasin hatchery trout lineages, supporting the mixed hatchery ancestry that had previously been inferred for lower Merced River *O. mykiss* (Pearse and Garza 2015).

STRUCTURE results showed strong convergence, verified by the highly consistent results among all four independent runs (data not shown). The distribution of STRUCTURE *q*-values in and among individuals supported previous findings of relationships between coastal and Central Valley *O. mykiss* and were similar to the individual DAPC results and population genetic analyses (see below). At low values of *K*, there were clear patterns of divergence between coastal steelhead and northern and southern Central Valley-lineage populations (*K* = 4; Figure 3). These patterns remained evident at higher values of *K*, with finer patterns of differentiation consistent with those seen by Pearse and Garza (2015).

### Population Approach

The DAPC of sample units indicated a strong geographical component, with the first and second axes of the DAPC plot roughly encompassing east–west and north–south geography (Figure 4). This pattern of divergence is concordant with previous studies showing a primary division between coastal and Central Valley steelhead and an association between geography and genetic differentiation among *O. mykiss* populations isolated above dams within the Central Valley but not among below-barrier Central Valley steelhead populations (Pearse and Garza 2015).

The phylogenetic tree also supported known patterns of geographic differentiation, although many nodes received less than 50% bootstrap support (Figure 5). Nonetheless, well-supported relationships among several pairs and groups of populations were consistent with previous studies, indicating that the current data set has sufficient power to resolve these relationships (e.g., close similarity of Feather River and Mokelumne Hatchery steelhead, Nimbus Hatchery steelhead

and coastal populations, and the relationships between the new UTR and UMR samples and reference samples from those locations; Pearse and Garza 2015). Among the new UTR and UMR samples, a clade of nine Tuolumne River populations (e.g., TUOL, Reed Creek [REED], upper Clavey River [UCLV], Roosevelt Lake [ROOS], FROG, etc.; Figure 5) was identified with moderate bootstrap support (77%), supporting their common indigenous ancestry. The South Fork Merced River (SFMC) sample appears as sister to this group but without significant support. Other UTR and UMR populations were more widely dispersed in the tree, possibly indicating more diverse sources contributing to these *O. mykiss* populations and also reflecting the limited resolution and low bootstrap support for deeper nodes in the tree. Meaningful support (68% and 98%) was found for the relationships between the GROS population, Kamloops Hatchery strain, and the northern Central Valley population from the McCloud River (Butcherknife Creek), further supporting the complete hatchery origin of the isolated above-barrier GROS population (Figure 5).

### Signatures of Migratory Adaptation

The frequency of the *Omy5* A haplotype in the sampling units within the UTR and UMR ranged from a minimum of 0.00 in GROS to a maximum of 0.31 in TUOL (Figure 6). Given their locations and accessibility to fish migrating from downstream reservoirs, the relatively high frequency of the A haplotype in the TUOL, FROG, and YOSV populations supports the suggestion that they sustain trout with adfluvial life histories. Conversely, the A haplotype exists at relatively low frequency in most other populations, particularly those found above barriers to migration (e.g., REED, Jawbone Creek [JAWB], and Crane Creek [CRAN]; Figures 1, 6). However, there was considerable variability among populations, likely reflecting a combination of selective factors impacting the frequency of adaptive genomic variation on chromosome *Omy5* and other parts of the genome.

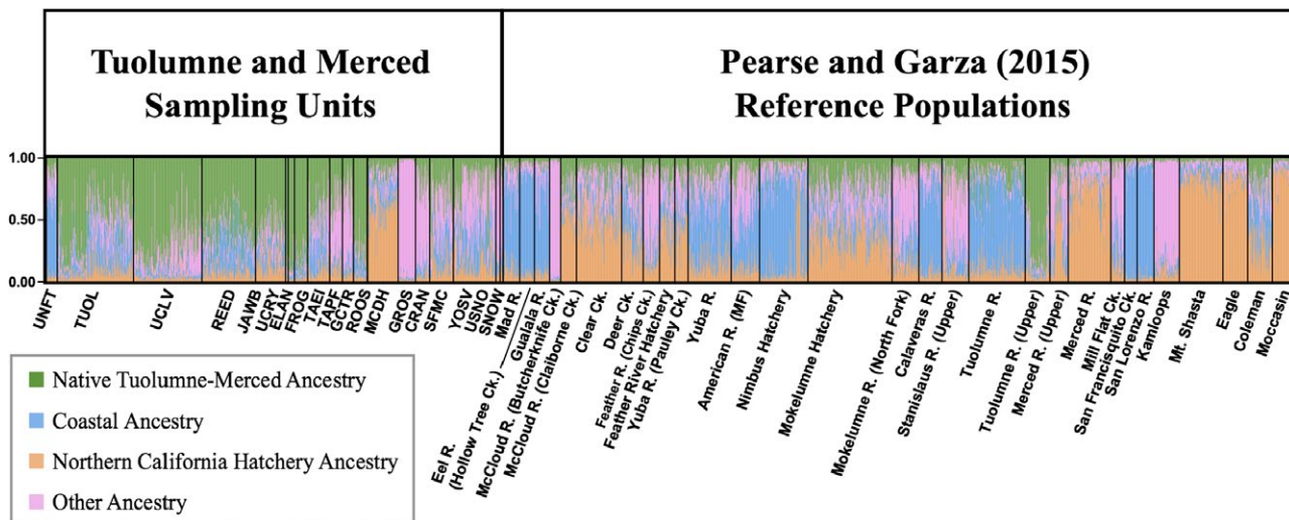


Figure 3. Individual-based plot of fractional ancestry from a hypothesized number of distinct genetic groups (*K* = 4) for *Onchorhynchus mykiss*. Sampling units and reference populations (as described and ordered in Table 1) are indicated along the bottom of the plot. Each individual is represented by a vertical line, and the proportion of estimated ancestry from each of the hypothetical genetic groups is represented by the proportionate amount of color within the vertical column (inferred ancestry: green = Yosemite National Park ancestry; blue = coastal ancestry; orange = northern Central Valley hatchery ancestry; pink = other ancestry; see text for details).

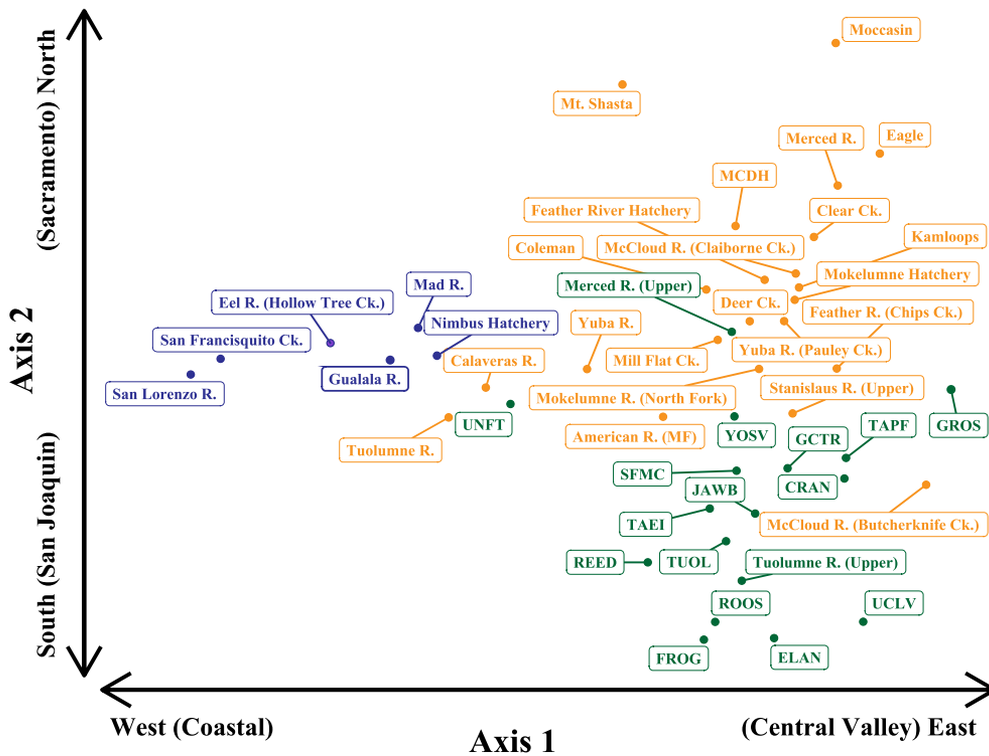


Figure 4. Population discriminant analysis of principal components plot, showing genetic relationships among *Oncorhynchus mykiss* from the upper Tuolumne River and upper Merced River sampling units (green; codes are defined in Table 1) relative to coastal reference populations (blue) and Central Valley reference populations and hatchery trout strains (orange). The central value for each population is indicated.

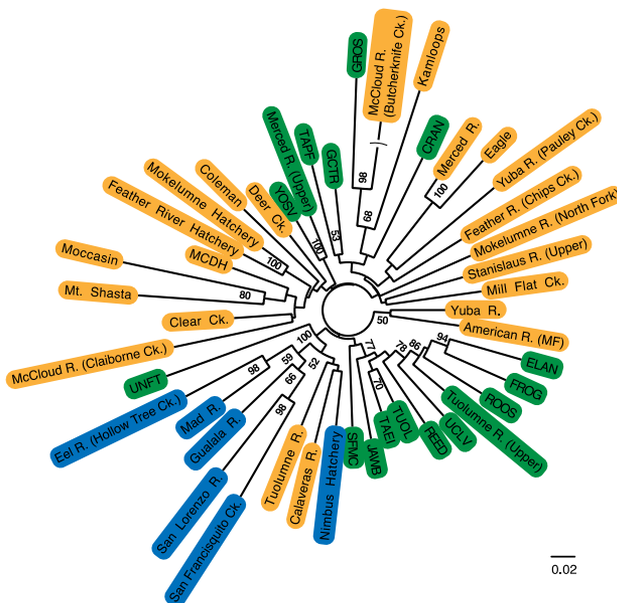


Figure 5. Neighbor-joining phylogenetic tree based on population chord distances, showing relationships among *Oncorhynchus mykiss* sampling units within the upper Tuolumne River and upper Merced River relative to other Central Valley and coastal *O. mykiss* populations. Colors highlight Yosemite National Park sampling units (green; codes are defined in Table 1), coastal reference populations (blue), and Central Valley reference populations and hatchery trout strains (orange). Bootstrap support from 1,000 replicates is depicted (values <50% are not shown). The branch to the McCloud River (Butcherknife Creek), indicated by a bisecting curve, has been shortened to one-third of the original length for visual presentation.

## DISCUSSION

Overall, the observed genetic relationships between Rainbow Trout in the UTR and UMR and other Central Valley *O. mykiss* populations and hatchery trout strains indicate that a mixture of lineages exists in these Yosemite watersheds. However, despite the extensive stocking with nonindigenous hatchery trout strains throughout the region, native ancestry appears to remain as the primary component of most sampling units examined in this study, with primarily indigenous southern Central Valley–San Joaquin River ancestry in reaches that were historically accessible to migratory salmonids. This includes the Clavey River, which has been designated as one of the “Heritage and Wild Trout Waters” by the California Department of Fish and Wildlife ([wildlife.ca.gov/fishing/inland/trout-waters](http://wildlife.ca.gov/fishing/inland/trout-waters)). These results support the hypothesis that local adaptation has played a key role in the persistence of these lineages.

In terms of ancestry, the primary division between coastal and Central Valley *O. mykiss* that has previously been documented (Nielsen et al. 2005; Pearse and Garza 2015) was also clear in multiple analyses of the present data set (Figures 3, 4, 5). This is important because it confirms that unlike some below-barrier populations in the southern Central Valley, including *O. mykiss* sampled in the lower Tuolumne River (Pearse and Garza 2015), the trout populations in the UTR and UMR do not show evidence of introgression from the coastal-origin steelhead propagated at Nimbus Hatchery. However, the close evolutionary relationships among all Central Valley *O. mykiss*—including most hatchery Rainbow Trout strains commonly used in California—hinder precise inference of population relationships and admixture within the Central Valley, and the weak

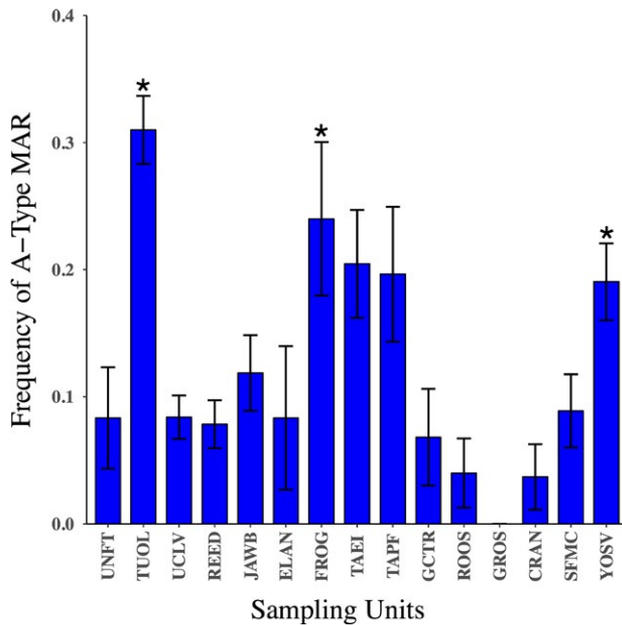


Figure 6. Frequency of the anadromous-type *Omy5* migration-associated region haplotype (A-Type MAR) estimated from upper Tuolumne River and upper Merced River sampling units of *Oncorhynchus mykiss* examined in this study (codes are defined in Table 1). Potentially adfluvial populations with access to reservoirs are indicated by asterisks. Standard errors for allele frequency estimates are shown (Hartl and Clark 1997).

signal of genetic differentiation among these populations likely reflects biological reality rather than limited resolution. Nonetheless, the relative proximities of populations shown in the DAPC revealed a clear pattern of geographic divergence among populations, with axis 2 highlighting the north–south divergence within the Central Valley (Figure 4). This is consistent with the hypothesis that Rainbow Trout populations isolated above dams in the Sierra Nevada better reflect their historical geographic origins than the scrambled steelhead populations that persist below barriers to migration (Pearse and Garza 2015).

The problems with hybrids in conservation have long been recognized (Allendorf et al. 2001), and the potential conservation value of hybrid populations remains an active area of discussion (Wayne and Shaffer 2016). Within the UTR and UMR, many populations show at least some evidence of mixed ancestry, as is common in studies of *O. mykiss* above dams (e.g. Winans et al. 2017), but we did not find the complete replacement of indigenous ancestry that has been observed in some regions of California subjected to intensive hatchery trout stocking (e.g., southern California; Abadía-Cardoso et al. 2016). Although admixed populations do not represent pure indigenous lineages, they often have high genetic diversity and should not be entirely discounted when considering source populations for recovery efforts (Abadía-Cardoso et al. 2016).

Most fish sampled at sites that were historically fishless due to their positions above barriers or at high elevation represented a mixture of indigenous and imported ancestries, with some having largely indigenous ancestry (e.g., FROG) while others appeared entirely descendent from hatchery Rainbow Trout strains (e.g., GROS). The UNFT sample showed

variable associations in different analyses, with genetic similarity to both the Coleman Hatchery trout strain and coastal lineage populations. The UNFT site has a long history of intensive hatchery stocking due to its location near a major road (California Highway 108), and both UNFT and GROS had low levels of heterozygosity, consistent with hatchery-strain ancestry. In contrast, the REED and JAWB populations had high heterozygosity and were genetically similar to other nearby populations within the UTR genetic group, despite being isolated above very large natural barrier waterfalls.

#### Adaptive Variation and Migratory Potential

It is important to note that adaptive genomic variation like that documented on chromosome *Omy5* is subject to the same factors that affect the distribution of neutral genetic variation among all natural populations, including drift due to small population sizes and introgression by nonnative lineages with highly divergent patterns of variation (Pearse 2016). In the case of hatchery Rainbow Trout, *Omy5* haplotype frequencies vary widely among strains, so their influence on introgressed wild populations is difficult to determine. However, to the extent that they reflect ongoing selection, the frequencies of alleles in this genomic region provide information about the relative fitness of alternative life history patterns in a given set of populations.

Within the UTR and UMR, the distribution of *Omy5* haplotype variation suggests that the populations most likely to express an adfluvial life history—and therefore to retain the potential to express anadromy—are those with unimpeded migratory access to Don Pedro and McClure reservoirs (e.g., TUOL and YOSV), as well as the FROG population tributary to Lake Eleanor (Figures 1, 6). Although the maximum frequency of migration-associated alleles among the UTR and UMR populations (0.31 in TUOL) was low relative to that of coastal anadromous and adfluvial populations (typically >0.60; Pearse et al. 2014; Leitwein et al. 2017), it was similar to that seen in potentially adfluvial populations of *O. mykiss* in the upper American River (0.33; DEP, unpublished data). In addition, the genomic region of *Omy5* associated with migratory life history patterns has also been associated with differences in temperature-specific development rates (Miller et al. 2012). The additional influence of temperature could contribute to the elevated frequency of resident-associated alleles in the colder, high-elevation populations, but further research is needed to better understand the factors that may influence the distribution of this adaptive genomic variation. Together, these results suggest that the UTR and UMR populations that now occupy river reaches between the reservoirs and the historical barriers to upstream migration are the most likely to express migratory adfluvial behavior and retain adaptive genomic variation associated with anadromy (Holecek et al. 2012; Holecek and Scarnecchia 2013; Leitwein et al. 2017).

#### Conservation Implications

Efforts to restore salmonid populations and the watersheds they inhabit will require a diverse set of approaches, investment, and cooperation among stakeholders (Phillis et al. 2013; Penaluna et al. 2016; Lackey 2017; Warren et al. 2017), particularly for migratory anadromous forms like steelhead (NMFS 2014). From an evolutionary genetics perspective, this study has several implications for the potential restoration of connectivity between the UTR and UMR populations and the California Central Valley steelhead DPS below the dams.

First, the present study was based on a data set with a modest number of SNPs by today's standards and thus has relatively low power to estimate population genetic parameters; a large genomic data set based on the thousands of loci generated by high-throughput sequencing could undoubtedly refine the results observed in the present study. For example, more than 230,000 SNP loci were recently used to accurately estimate the proportions of European, African, and Native American ancestry in admixed human populations in Colombia (Conley et al. 2017). It is also possible that hatchery trout strains that were not included in the present study have been stocked in these watersheds, so their contributions could not be specifically detected. However, the basic conclusions regarding the distribution of indigenous Rainbow Trout within the Tuolumne and Merced river watersheds and their implications for management are unlikely to change in biologically significant ways. Similarly, further characterization of the distribution of adaptive genomic variation on chromosome Omy5 and other parts of the genome will provide insight into the evolutionary processes affecting trout populations above dams. However, such information would not necessarily impact conservation planning because the basic principles of conservation genetic management to preserve genetic diversity remain the same (Pearse 2016). Nonetheless, as more examples of adaptive genomic variation associated with life history traits are identified in *O. mykiss* and other salmonid species (Barson et al. 2015; Hess et al. 2016), fisheries managers will need to carefully consider the most appropriate ways to conserve and protect this important biodiversity (Pearse 2016).

Second, Pearse and Garza (2015) detected introgression by coastal-origin steelhead propagated at Nimbus Hatchery into the limited populations of *O. mykiss* that remain in the ocean-accessible river reaches below dams in the Calaveras, Tuolumne, and Stanislaus rivers and found that *O. mykiss* captured in the lower Merced River were primarily descended from hatchery trout, especially the Eagle Lake strain. The signal of hatchery ancestry observed in the sample of 59 lower Merced River fish analyzed in the present study further confirms this result. However, recent data on the physiology of steelhead in the lower Tuolumne River have shown that they have a much higher thermal tolerance than populations from northern latitudes, demonstrating their local adaptation to the high temperatures of the southern Central Valley (Verhille et al. 2016). Thus, the *O. mykiss* currently inhabiting below-barrier reaches of the Tuolumne and Merced rivers likely represent a mixture of indigenous, hatchery, and coastal ancestry, and both admixture and local adaptation have likely influenced their current genetic composition, including the frequencies of Omy5 haplotypes and other adaptive genomic variation.

Third, although our data show that the Rainbow Trout trapped above these dams have both ancestry and adaptive genomic variation that are representative of indigenous migratory populations, the development of an anadromous steelhead population from these stocks through fish passage via two-way trap-and-haul operations or other means presents many challenges (Lusardi and Moyle 2017). Re-establishing gene flow between formerly connected populations above and below barrier dams has many potential benefits in terms of maintenance of genetic diversity and facilitating adaptation, but these must be evaluated against the possible risks and constraints within the larger reintroduction and recovery framework (Anderson

et al. 2014). Nonetheless, anadromous salmonid life histories can emerge rapidly from formerly adfluvial populations after dam removal, demonstrating that such populations are capable of re-establishing their dormant ability to complete an ocean migration (Quinn et al. 2017). In this context, migratory adfluvial individuals in the Tuolumne River, Merced River, and other Central Valley watersheds could be considered as potential contributors to future fish passage programs and reintroduction efforts (Thrower et al. 2008) provided that the logistical issues associated with re-establishing connectivity can be overcome (NMFS 2014). Thus, in considering the potential for passage of migratory fish above New Don Pedro and New Exchequer dams, directed studies are needed to determine the potential for trapping downstream migrants, among other considerations, as has been undertaken in similar situations (e.g., Clancey et al. 2017; Winans et al. 2018).

Finally, it should be noted that the populations of steelhead in the southern Central Valley are likely among the most vulnerable to the impacts of climate change, so their continued persistence is far from certain. Therefore, in the context of protecting and restoring anadromous fish populations in California, genetic factors should be considered as secondary to the basic need for access to appropriate habitat to support all phases of the migratory life cycle. This includes access to suitable spawning and rearing habitats, as provided by removal of large barrier dams or via carefully monitored two-way trap-and-haul fish passage programs (Anderson et al. 2014), as well as modification or removal of smaller migration barriers (Apgar et al. 2017), adjustments to flow regimes, and other improvements in downstream habitats to support native fishes and restore viable migratory connectivity with the ocean for both out-migrating juveniles and returning adult salmonids (NMFS 2014). In the absence of these changes, the existence of migratory salmonid populations in the Central Valley will continue to depend on hatchery propagation and other interventions until the dams that block their migratory paths are modified or removed (Katz et al. 2013; Quiñones et al. 2015).

#### ACKNOWLEDGMENTS

We thank John Wooster and Steve Edmondson (NMFS West Coast Region, California Central Valley Office, Hydro team) and the members of the NMFS Southwest Fisheries Science Center's Molecular Ecology and Genetic Analysis Team, particularly Cassie Columbus, Ellen Campbell, Elena Correa, and Mary Ables Ray, for their support on this project. Diana Baetscher, William Foster, John Carlos Garza, Cyril Michel, and Erin Strange provided valuable comments that improved the manuscript. We also thank Rob Grasso for facilitating access and assisting with sampling in Yosemite National Park. Samples were collected under Scientific Collecting Permit 13029 from the California Department of Fish and Wildlife. Special thanks are extended to all of the fly fishers who contributed their time and skills to obtain samples for this project, including Ben Burford, Rob Grasso, Sean Hayes, Tom Holley, Steve Lindley, Michael Martin, Alex McHuron, Cyril Michel, Jeremy Notch, David Swank, and Larry Thompson. There is no conflict of interest declared in this article.

#### REFERENCES

- Abadía-Cardoso, A., E. C. Anderson, D. E. Pearse, and J. C. Garza. 2013. Large-scale parentage analysis reveals reproductive patterns and heritability of spawn timing in a hatchery population of steelhead (*Oncorhynchus mykiss*). *Molecular Ecology* 22:4733–4746.

- Abadía-Cardoso, A., D. E. Pearse, S. Jacobson, J. Marshall, D. Dalrymple, F. Kawasaki, G. Ruiz-Campos, and J. C. Garza. 2016. Population genetic structure and ancestry of steelhead/Rainbow Trout (*Oncorhynchus mykiss*) in southern California and Baja California coastal rivers and streams. *Conservation Genetics* 17:675–689.
- Allendorf, F. W., R. F. Leary, P. Spruell, and J. K. Wenburg. 2001. The problems with hybrids: setting conservation guidelines. *Trends in Ecology and Evolution* 16:613–622.
- Anderson, J. H., G. R. Pess, R. W. Carmichael, M. J. Ford, T. D. Cooney, C. M. Baldwin, and M. M. McClure. 2014. Planning Pacific salmon and steelhead reintroductions aimed at long-term viability and recovery. *North American Journal of Fisheries Management* 34:72–93.
- Apgar, T. M., D. E. Pearse, and E. P. Palkovacs. 2017. Evolutionary restoration potential evaluated through the use of a trait-linked genetic marker. *Evolutionary Applications* 10:485–497. <https://doi.org/10.1111/eva.12471>.
- Barson, N. J., T. Aykanat, K. Hindar, M. Baranski, G. H. Bolstad, P. Fiske, C. Jacq, A. J. Jensen, S. E. Johnston, S. Karlsson, M. Kent, T. Moen, E. Niemela, T. Nome, T. Naesje, P. Orell, A. Romakkaniemi, H. Saegrov, K. Urdal, J. Erkinaro, S. Lien, and C. Primmer. 2015. Sex-dependent dominance at a single locus maintains variation in age at maturity in salmon. *Nature* 528:405–408.
- Berejikian, B. A., R. A. Bush, and L. A. Campbell. 2014. Maternal control over offspring life history in a partially anadromous species, *Oncorhynchus mykiss*. *Transactions of the American Fisheries Society* 143:369–379.
- Brenkman, S. J., K. T. Sutton, and A. R. Marshall. 2017. Life history observations of adfluvial Chinook Salmon prior to reintroduction of anadromous salmonids. *North American Journal of Fisheries Management* 37:1220–1230.
- Brunelli, J. P., K. J. Wertzler, K. Sundin, and G. H. Thorgaard. 2008. Y-specific sequences and polymorphisms in Rainbow Trout and Chinook Salmon. *Genome* 51:739–748.
- Busby, P. J., T. C. Wainwright, G. J. Bryant, L. J. Lierheimer, R. S. Waples, F. W. Waknitz, and I. V. Lagomarsino. 1996. Status review of West Coast steelhead from Washington, Idaho, Oregon, and California. National Oceanic and Atmospheric Administration, Northwest Fisheries Science Center, Seattle, Washington.
- Caton, J. D. 1869. Trout fishing in the Yosemite Valley. *American Naturalist* 3:519–522.
- Cavalli-Sforza, L. L., and A. W. F. Edwards. 1967. Phylogenetic analysis: models and estimation procedures. *American Journal of Human Genetics* 19:233–257.
- Chessel, D., and A. Dufour. 2004. The ade4 package I: one-table methods. *R News* 4:5–10.
- Clancey, K., L. Saito, K. Hellmann, C. Svoboda, J. Hannon, and R. Beckwith. 2017. Evaluating head-of-reservoir water temperature for juvenile Chinook Salmon and steelhead at Shasta Lake with modeled temperature curtains. *North American Journal of Fisheries Management* 37:1161–1175.
- Clemente, A. J., E. C. Anderson, D. Boughton, D. Girman, and J. C. Garza. 2009. Population genetic structure and ancestry of *Oncorhynchus mykiss* populations above and below dams in south-central California. *Conservation Genetics* 10:1321–1337.
- Conley, A. B., L. Rishishwar, E. T. Norris, A. Valderrama-Aguirre, L. Mariño-Ramírez, M. A. Medina-Rivas, and I. K. Jordan. 2017. A comparative analysis of genetic ancestry and admixture in the Colombian populations of Chocó and Medellín. *G3: Genes, Genomes, Genetics* 7:3435–3447.
- Courter, I. I., D. B. Child, J. A. Hobbs, T. M. Garrison, J. J. G. Glessner, and S. Duery. 2013. Resident Rainbow Trout produce anadromous offspring in a large interior watershed. *Canadian Journal of Fisheries and Aquatic Sciences* 70:701–710.
- Cuthbert, R., C. Becker, and A. Fuller. 2012. Fall/winter migration monitoring at the Tuolumne River Weir, 2011 annual report. Prepared for the Turlock and Modesto Irrigation Districts by FISHBIO Environmental, Oakdale, California.
- Dray, S., and A. B. Dufour. 2007. The ade4 package: implementing the duality diagram for ecologists. *Journal of Statistical Software* 22:1–20.
- Dray, S., A. Dufour, and D. Chessel. 2007. The ade4 package II: two-table and K-table methods. *R News* 7:47–54.
- Falush, D., M. Stephens, and J. K. Pritchard. 2003. Inference of population structure using multilocus genotype data: linked loci and correlated allele frequencies. *Genetics* 164:1567–1587.
- Fisher, F. W. 1994. Past and present status of Central Valley Chinook Salmon. *Conservation Biology* 8:870–873.
- Ford, T., and S. Kirihaara. 2010. Tuolumne River *Oncorhynchus mykiss* monitoring report. Prepared for the Federal Energy Regulatory Commission by the Turlock and Modesto Irrigation Districts and Stillwater Sciences, Berkeley, California.
- Hale, M. C., F. P. Thrower, E. A. Berntson, M. R. Miller, and K. M. Nichols. 2013. Evaluating adaptive divergence between migratory and non-migratory ecotypes of a salmonid fish, *Oncorhynchus mykiss*. *G3: Genes, Genomes, Genetics* 3:1273–1285.
- Hartl, D. L., and A. G. Clark. 1997. *Principles of population genetics*, 3rd edition. Sinauer Associates, Sunderland, Massachusetts.
- Hecht, B. C., N. R. Campbell, D. E. Holecek, and S. R. Narum. 2013. Genome-wide association reveals genetic basis for the propensity to migrate in wild populations of Rainbow and steelhead trout. *Molecular Ecology* 22:3061–3076.
- Hess, J. E., J. S. Zandt, A. R. Matala, and S. R. Narum. 2016. Genetic basis of adult migration timing in anadromous steelhead discovered through multivariate association testing. *Proceedings of the Royal Society B* 283:20153064.
- Holecek, D. E., and D. L. Scarnecchia. 2013. Comparison of two life history strategies after impoundment of a historically anadromous stock of Columbia River Redband Trout. *Transactions of the American Fisheries Society* 142:1157–1166.
- Holecek, D. E., D. L. Scarnecchia, and S. E. Miller. 2012. Smoltification in an impounded, adfluvial Redband Trout population upstream from an impassable dam: does it persist? *Transactions of the American Fisheries Society* 141:68–75.
- Jombart, T. 2008. Adegenet: a R package for the multivariate analysis of genetic markers. *Bioinformatics* 24:1403–1405.
- Jombart, T., S. Devillard, and F. Balloux. 2010. Discriminant analysis of principal components: a new method for the analysis of genetically structured populations. *BMC Genetics* 11:94.
- Kamvar, Z. N., J. F. Tabima, and N. J. Grünwald. 2014. Poppr: an R package for genetic analysis of populations with clonal, partially clonal, and/or sexual reproduction. *PeerJ* 2:e281.
- Katz, J. V. E., P. B. Moyle, R. M. Quiñones, J. Israel, and S. Purdy. 2013. Impending extinction of salmon, steelhead, and trout (*Salmonidae*) in California. *Environmental Biology of Fishes* 96:1169–1186.
- Kendall, N. W., J. R. McMillan, M. R. Sloat, T. W. Buehrens, T. P. Quinn, G. R. Pess, K. V. Kuzishchin, M. M. McClure, and R. W. Zabel. 2015. Anadromy and residency in steelhead and Rainbow Trout (*Oncorhynchus mykiss*): a review of the processes and patterns. *Canadian Journal of Fisheries and Aquatic Sciences* 342:319–342.
- Lackey, R. T. 2017. Science and salmon recovery. Pages 69–94 *in* E. P. Weber, D. H. Lach and B. S. Steel, editors. *New strategies for wicked problems: science and solutions in the 21st century*. Oregon State University Press, Corvallis.
- Leitritz, E. 1970. A history of California's fish hatcheries, 1870–1960. *California Department of Fish and Game Fish Bulletin* 150:1–125.
- Leitwein, M., J. C. Garza, and D. E. Pearse. 2017. Ancestry and adaptive evolution of anadromous, resident, and adfluvial Rainbow Trout (*Oncorhynchus mykiss*) in the San Francisco Bay area: application of adaptive genomic variation to conservation in a highly impacted landscape. *Evolutionary Applications* 10:56–67.
- Lindley, S. T., R. S. Schick, A. Agrawal, M. Goslin, T. E. Pearson, E. Mora, J. J. Anderson, B. May, S. Greene, C. Hanson, A. Low, D. McEwan, R. B. MacFarlane, C. Swanson, and J. G. Williams. 2006. Historical population structure of Central Valley steelhead and its alteration by dams. *San Francisco Estuary and Watershed Science* 4(1):3. Available: <http://escholarship.org/uc/item/1ss794fc>.
- Lusardi, R. A., and P. B. Moyle. 2017. Two-way trap and haul as a conservation strategy for anadromous salmonids. *Fisheries* 42:478–487.
- May, J. T., and L. R. Brown. 2002. Fish communities of the Sacramento River basin: implications for conservation of native fishes in the Central Valley, California. *Environmental Biology of Fishes* 63:373–388.
- McEwan, D. R. 2001. Central Valley steelhead. *Fish Bulletin* 179:1–43.
- Meeke, M. H., M. R. Stephens, K. M. Tomalty, B. May, and M. R. Baerwald. 2014. Genetic considerations for sourcing steelhead reintroductions: investigating possibilities for the San Joaquin River. *San Francisco Estuary and Watershed Science* 12(1). Available: <http://escholarship.org/uc/item/gwn5q90h>

- Miller, M. R., J. P. Brunelli, P. A. Wheeler, S. Liu, C. E. Rexroad, Y. Palti, C. Q. Doe, and G. H. Thorgaard. 2012. A conserved haplotype controls parallel adaptation in geographically distant salmonid populations. *Molecular Ecology* 21:237–249.
- Moyle, P. B., J. A. Hobbs, and J. R. Durand. 2018. Delta Smelt and water politics in California. *Fisheries* 43:42–51.
- Narum, S. R., J. S. Zandt, D. Graves, and W. R. Sharp. 2008. Influence of landscape on resident and anadromous life history types of *Oncorhynchus mykiss*. *Canadian Journal of Fisheries and Aquatic Sciences* 65:1013–1023.
- Neave, F. 1944. Racial characteristics and migratory habits in *Salmo gairdneri*. *Journal of the Fisheries Research Board of Canada* 6:245–51.
- Nichols, K. M., A. F. Edo, P. A. Wheeler, and G. H. Thorgaard. 2008. The genetic basis of smoltification-related traits in *Oncorhynchus mykiss*. *Genetics* 179:1559–1575.
- Nielsen, J. L., S. A. Pavey, T. Wiacek, and I. Williams. 2005. Genetics of Central Valley *O. mykiss* populations: drainage and watershed scale analysis. *San Francisco Estuary and Watershed Science* 3(2):3. Available: <http://escholarship.org/uc/item/6sc3905g>
- NMFS (National Marine Fisheries Service). 2006. Endangered and threatened species: final listing determinations for 10 distinct population segments of West Coast steelhead, final rule. *Federal Register* 71(January 5, 2006):834–862.
- NMFS (National Marine Fisheries Service). 2014. Recovery plan for the evolutionarily significant units of Sacramento River winter-run Chinook Salmon and Central Valley spring-run Chinook Salmon and the distinct population segment of California Central Valley steelhead. NMFS, California Central Valley Area Office, Sacramento.
- Northcote, T. G. 2010. Controls for trout and char migratory/resident behaviour mainly in stream systems above and below waterfalls/barriers: a multidecadal and broad geographical review. *Ecology of Freshwater Fish* 19:487–509.
- Olsen, J. B., K. Wuttig, D. Fleming, E. J. Kretschmer, and J. K. Wenburg. 2006. Evidence of partial anadromy and resident-form dispersal bias on a fine scale in populations of *Oncorhynchus mykiss*. *Conservation Genetics* 7:613–619.
- Paradis, E. 2010. Pegas: an R package for population genetics with an integrated-modular approach. *Bioinformatics* 26:419–420.
- Pavlik, R. C. 1987. The trout hatcheries of Yosemite. *Yosemite* 49:8–9.
- Pearse, D. E. 2016. Saving the spandrels? Adaptive genomic variation in conservation and fisheries management. *Journal of Fish Biology* 89:2697–2716.
- Pearse, D. E., and J. C. Garza. 2015. You can't unscramble an egg: population genetic structure of *Oncorhynchus mykiss* in the California Central Valley inferred from combined microsatellite and SNP data. *San Francisco Estuary and Watershed Science* 13(4). Available: <http://escholarship.org/uc/item/8dk7m218>
- Pearse, D. E., S. A. Hayes, M. H. Bond, C. V. Hanson, E. C. Anderson, R. B. Macfarlane, and J. C. Garza. 2009. Over the falls? Rapid evolution of ecotypic differentiation in steelhead/Rainbow Trout (*Oncorhynchus mykiss*). *Journal of Heredity* 100:515–525.
- Pearse, D. E., M. R. Miller, A. Abadía-Cardoso, and J. C. Garza. 2014. Rapid parallel evolution of standing variation in a single, complex, genomic region is associated with life history in steelhead/Rainbow Trout. *Proceedings of the Royal Society B* 281:20140012.
- Penaluna, B. E., A. Abadía-Cardoso, J. B. Dunham, F. J. García-Dé León, R. E. Gresswell, A. R. Luna, E. B. Taylor, B. B. Shepard, R. Al-Chokhachy, C. C. Muhlfeld, K. R. Bestgen, K. Rogers, M. A. Escalante, E. R. Keeley, G. Temple, J. E. Williams, K. Matthews, R. Pierce, R. L. Mayden, R. P. Kovach, J. C. Garza, and K. D. Fausch. 2016. Conservation of native Pacific trout diversity in western North America. *Fisheries* 41:286–300.
- Phillis, C. C., J. W. Moore, M. Buoro, S. A. Hayes, J. C. Garza, and D. E. Pearse. 2016. Shifting thresholds: rapid evolution of migratory life histories in steelhead/Rainbow Trout, *Oncorhynchus mykiss*. *Journal of Heredity* 107:51–60.
- Phillis, C. C., S. M. O'Regan, S. J. Green, J. E. B. Bruce, S. C. Anderson, J. N. Linton, and B. Favaro. 2013. Multiple pathways to conservation success. *Conservation Letters* 6:98–106.
- Pritchard, J. K., M. Stephens, and P. Donnelly. 2000. Inference of population structure using multilocus genotype data. *Genetics* 155:945–959.
- Quinn, T. P. 2011. *The behavior and ecology of Pacific salmon and trout*. University of British Columbia Press, Vancouver.
- Quinn, T. P., M. H. Bond, S. J. Brenkman, R. Paradis, and R. J. Peters. 2017. Re-awakening dormant life history variation: stable isotopes indicate anadromy in Bull Trout following dam removal on the Elwha River, Washington. *Environmental Biology of Fishes* 100:1659–1671. Available: <https://doi.org/10.1007/s10641-017-0676-0>
- Quiñones, R. M., T. E. Grantham, B. N. Harvey, J. D. Kiernan, M. Klasson, A. P. Wintzer, and P. B. Moyle. 2015. Dam removal and anadromous salmonid (*Oncorhynchus* spp.) conservation in California. *Reviews in Fish Biology and Fisheries* 25:195–215.
- R Development Core Team. 2017. R: a language and environment for statistical computing. R Foundation for Statistical Computing, Vienna, Austria.
- Rosenberg, N. A. 2004. DISTRUCT: a program for the graphical display of population structure. *Molecular Ecology Resources* 4:137–138.
- Rundio, D. E., T. H. Williams, D. E. Pearse, and S. T. Lindley. 2012. Male-biased sex ratio of nonanadromous *Oncorhynchus mykiss* in a partially migratory population in California. *Ecology of Freshwater Fish* 21:293–299.
- Sard, N. M., D. P. Jacobson, and M. A. Banks. 2016. Grandparentage assignments identify unexpected adfluvial life history tactic contributing offspring to a reintroduced population. *Ecology and Evolution* 6:6773–6783.
- Thrower, F. P., J. E. Joyce, A. G. Celewyc, and P. W. Malecha. 2008. The potential importance of reservoirs in the western United States for the recovery of endangered populations of anadromous steelhead. Pages 309–324 in M. S. Allen, S. Sammons, and M. J. Maceina, editors. *Balancing fisheries management and water uses for impounded river systems*. American Fisheries Society, Symposium 62, Bethesda, Maryland.
- Venables, W., and B. Ripley. 2002. *Modern Applied Statistics with S*, 4th edition. Springer, New York.
- Verhille, C. E., K. K. English, D. E. Cocherell, A. P. Ferrell, and N. A. Fangue. 2016. High thermal tolerance of a Rainbow Trout population near its southern range limit suggests local thermal adjustment. *Conservation Physiology* 4:1–12.
- Warren, R. F., T. Reeve, and J. S. Arnold. 2017. Reimagining watershed restoration: a call for new investment and support structures for greater resiliency and long-term impact. *WIREs Water* 4:e1174. <https://doi.org/10.1002/wat2.1174>.
- Wayne, R. K., and H. B. Shaffer. 2016. Hybridization and endangered species protection in the molecular era. *Molecular Ecology* 25:2680–2689.
- Williams, S. M., B. J. Holmes, and J. G. Pepperell. 2015. The novel application of non-lethal citizen science tissue sampling in recreational fisheries. *PLoS ONE* 10:e0135743. Available: <https://doi.org/10.1371/journal.pone.0135743>
- Winans, G. A., M. B. Allen, J. Baker, E. Lesko, F. Shrier, B. Strobel, and J. Myers. 2018. Dam trout: genetic variability in *Oncorhynchus mykiss* above and below barriers in three Columbia River systems prior to restoring migrational access. *PLoS ONE* 13(5):e0197571. Available: <https://doi.org/10.1371/journal.pone.0197571>
- Winans, G. A., M. C. Baird, and J. Baker. 2010. A genetic and phenetic baseline before the recolonization of steelhead above Howard Hanson Dam, Green River, Washington. *North American Journal of Fisheries Management* 30:742–756.
- Winans, G. A., J. Baker, M. McHenry, L. Ward, and J. Myers. 2017. Genetic characterization of *Oncorhynchus mykiss* prior to dam removal with implications for recolonization of the Elwha River watershed, Washington. *Transactions of the American Fisheries Society* 146:160–172.
- Yoshiyama, R. M., F. W. Fisher, and P. B. Moyle. 1998. Historical abundance and decline of Chinook Salmon in the Central Valley region of California. *North American Journal of Fisheries Management* 18:487–521.
- Yoshiyama, R. M., E. R. Gerstung, F. W. Fisher, and P. B. Moyle. 2001. Historical and present distribution of Chinook Salmon in the Central Valley drainage of California. *California Department of Fish and Game Fish Bulletin* 179:71–176. [AFS](#)