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GENETIC CHARACTERIZATION OF TWO POPULATIONS OF BONNEVILLE CUTTHROAT TROUT IN GREAT BASIN NATIONAL PARK, USA

Derek D. Houston1, R. Paul Evans2, Jared M. Crowley3, and Dennis K. Shiozawa3

ABSTRACT.—The cutthroat trout (Oncorhynchus clarkii) is a popular sport fish that is native to the waters of western North America. Cutthroat trout comprises many subspecies, each of which has experienced range reduction resulting from anthropogenic activities. Hence, there is a general interest from management agencies to assess the genetic structure of managed populations of cutthroat trout subspecies to ensure that proper conservation plans are implemented. Herein, we utilize microsatellite data to characterize the genetic composition of 2 populations of Bonneville cutthroat trout in Great Basin National Park: Mill Creek and South Fork Big Wash. Mill Creek was used as a source population for reintroduction into South Fork Big Wash in the year 2000, and there is concern that South Fork Big Wash may have experienced a population bottleneck after, or during, the stocking effort. We found that both populations exhibit low genetic diversity, and that the source population, Mill Creek, exhibited mixed signals of having undergone a recent population bottleneck. Structure analysis revealed 4 distinct groups, but those groups did not segregate geographically, although a significant pairwise FST (0.06727, P < 0.00001) between Mill Creek and South Fork Big Wash populations suggests that some genetic differentiation has occurred.

The identification of conservation units (i.e., evolutionary significant units, management units, etc.) within a species is critical to maintaining genetic diversity, an important management objective (Waples 1994, Moritz 1999, 2002, Wang et al. 2002, Reed and Frankham 2003). An essential step in this process is the collection of data that can be used to identify genetically distinct populations (e.g., Peterman et al. 2013). Preserving these populations helps maintain total genetic diversity, which gives species the ability to persist over time (O’Brien et al. 1985, Allendorf and Leary 1988, Spielman et al. 2004). The loss of genetic diversity within a species increases extinction risk by causing a decreased ability to adapt to environmental change and a reduction in fitness due to inbreeding depression (Newman and Pilson 1997, Amos and Balmford 2001, Charlesworth and Willis 2009). In addition to contributing to total genetic diversity, genetically distinct populations can also be important to the evolutionary legacy of a species. Populations at the periphery of a species range are often genetically distinct, and can be where the first steps in speciation occur (Lesica and Allendorf 1995, Demenmoser et al. 2013). Conserving genetically unique populations therefore decreases the likelihood of extinction and simultaneously allows for the continued evolution of the species.

Reduced genetic variation is of general concern in small, isolated populations (Ellstrand and Elam 1993, Lacy 2000), or in populations

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that have experienced founder effects. A reduction in genetic variation can be accompanied by inbreeding and the fixation of deleterious traits (Hedrick and Kalinowski 2000). Such traits can be manifest through fluctuating asymmetry (Leary et al. 1985), reduced fecundity (Kincaid 1976, Su et al. 1996), reduced survivorship, and so forth. In such cases, it can be useful to introduce additional genetic diversity from outside sources to prevent genetic load associated with inbreeding depression. Therefore, it is worthwhile to periodically evaluate conservation populations to ensure that the proper conservation plan is implemented.


The Bonneville cutthroat trout, *Oncorhynchus clarkii utah*, is native to drainages within the Bonneville Basin and was interconnected through tributaries to pluvial Lake Bonneville. Bonneville cutthroat trout populations were fragmented with the desiccation of Lake Bonneville at the end of the Pleistocene Epoch. More recently, many native trout habitats have also been altered by anthropogenic activities including irrigation, grazing of livestock, introduction of nonnative fishes, and changing global climate.

Great Basin National Park is located on the western edge of the Bonneville Basin, the periphery of the range of the Bonneville cutthroat trout. Fourteen conservation populations of Bonneville cutthroat trout occur in Great Basin National Park and the surrounding Mount Moriah Wilderness Area (Snake Range, NV). Two of these populations, South Fork Big Wash and Mill Creek (Fig. 1, Table 1), both in Great Basin National Park, are of particular interest to Great Basin National Park, which oversees the management of these populations. When surveyed by National Park Service and Nevada Department of Wildlife officials in 1999, South Fork Big Wash was unpopulated by cutthroat trout but was restored beginning in the year 2000, using Mill Creek cutthroat trout as a source population. The transfer of fish into the South Fork Big Wash Creek occurred over 2 years, one transfer in July 2000 and the other in July 2001. These populations have not had any additional transfers, nor have they been evaluated genetically since then, but concern exists that the South Fork Big Wash population may have experienced a reduction in genetic diversity resulting from a founder effect with the original stocking effort.

Our objectives were to compare genetic variation within the South Fork Big Wash population, restored in the year 2000, to that of the source population, Mill Creek, and to assess the genetic composition of each stream. We used microsatellite markers to assess the genetic status of 68 individual cutthroat trout samples from these 2 populations to determine whether any signature of a population bottleneck is evident.

We anticipated several possibilities. First, if substantial microsatellite variability exists in the source population, then the restored population should contain at least a subset of this variation. It is unlikely that the entire diversity of the source population (Mill Creek) would have been transferred to the new population (South Fork Big Wash). However, the source population may have also undergone significant bottleneck effects following the desiccation of Lake Bonneville, or as a result of more recent anthropogenic habitat manipulations. Thus, it is possible that genetic diversity of the parent population is also very low. It is also possible that either population (or both) has experienced postrestoration bottlenecks.

**METHODS**

**Sampling and DNA Isolation**

The National Park Service and the Nevada Division of Wildlife provided fin clips of 68 Bonneville cutthroat trout individuals collected in 2010 for genetic analysis: 34 from South
Fork Big Wash and 34 from Mill Creek (Table 1). We extracted whole genomic DNA from fin clips using Qiagen DNeasy tissue kits according to the manufacturer’s recommended protocols. Unused tissues have been catalogued and stored in the Monte L. Bean Life Science Museum ichthyological collection at Brigham Young University (BYU). The DNA extracts were stored at 4°C during the analysis, and have been placed at −80°C for long-term storage.

Polymerase Chain Reaction and Genotyping

Microsatellite regions were amplified via the polymerase chain reaction (PCR) using a 3-primer system that incorporated a fluorescently labeled M13 primer rather than a fluorescently labeled forward primer in each microsatellite primer pair (Schuelke 2000, Blacket et al. 2012). The 5’ end of each microsatellite forward primer was expanded to include an M13 forward sequence (CAC GAC GTT GTA AAA CGA C; Hanna et al. 2012). The extended forward primer was then used in conjunction with an M13 primer that was fluorescently labeled with either 6-FAM or HEX. Hence, each PCR cocktail contained 3 primers: a 5’-augmented microsatellite forward primer, a fluorescently labeled M13
forward primer, and an unmodified microsatellite reverse primer. Microsatellite primers are listed in Table 2. Reactions were 12.5 μL in total volume and contained the following ingredients: 2.3 μL molecular grade H$_2$O, 0.05 μL 5’-augmented microsatellite forward primer, 0.45 μL fluorescently labeled M13 forward primer, 0.45 μL unmodified reverse primer, 6.25 μL of Promega GoTaq master mix, and 3 μL of DNA. The thermal profile consisted of an initial denature at 95 °C for 2 min, followed by 40 cycles of 95 °C for 15 s, primer specific annealing temperature (ranging from 50 °C to 60 °C, see Table 2) for 90 s, and extension at 72 °C for 90 s; a final extension at 72 °C for 10 min ran at the conclusion of the last cycle and that was followed by a 4 °C hold.

Successful PCR amplifications of the microsatellite loci were verified qualitatively by viewing bands following gel electrophoresis. PCR products were diluted 1:50 with nuclease-free water, then 10 μL of each diluted sample was added to the appropriate well of a new 96-well plate and dried prior to genotyping. Samples were genotyped on an ABI 3100 automated sequencer after 0.3 μL of ROX 500 size standard and 12 μL HiDi Formamide were added to each dried sample. Peaks were initially scored automatically using PEAKSCANNER (Applied Biosystems), and final scoring of microsatellite alleles was performed by eye using the software program GENEIOUS v.6.1.4 (Biomatters; www.geneious.com).

### Population Genetic Analyses

We used MICRO-CHECKER v.2.3.4 (Van Oosterhout et al. 2004) to check for possible genotyping errors (i.e., stuttering, large allele dropout, and null alleles). We used ARLEQUIN v.3.5.1.3 (Excoffier and Lischer 2010) to calculate the number of alleles, the size range of alleles, the expected and observed heterozygosities (needed to perform Hardy–Weinberg equilibrium tests), and a pairwise F$_{ST}$ estimate between the 2 cutthroat trout populations. The input files for ARLEQUIN were created using the file conversion option in GENEPOP v.4.2 (Raymond and Rousset 1995, Rousset 2008).

We used BOTTLENECK v.1.2.02 to determine whether populations had undergone genetic bottlenecks (Cornuet and Luikart 1996). The null hypothesis is that the populations are in mutation-drift equilibrium (i.e., there is no signature of a recent population bottleneck). However, we infer that populations exhibiting significant heterozygosity excesses have experienced a recent bottleneck (Cornuet and Luikart 1996). Such a signal results from a Hardy–Weinberg imbalance following a recent bottleneck and dissipates over time as generations of random mating return the population to equilibrium, albeit with reduced genetic variation. We performed sign tests, standardized differences tests, and Wilcoxon rank tests in BOTTLENECK under the infinite alleles model and the stepwise mutational model based on 1000 iterations.

We used STRUCTURE (Pritchard et al. 2000) and STRUCTURE-HARVESTER (Earl and vonHoldt 2012) to assess whether the Mill Creek and South Fork Big Wash populations were genetically distinct. The STRUCTURE analysis was evaluated 30 times for each K (with K ranging from 1 to 8), with 1,000,000 repetitions per run after discarding the first 100,000 repetitions as burn-in.

### RESULTS

**Polymerase Chain Reaction and Genotyping**

Thirteen primer sets successfully amplified microsatellite loci in Bonneville cutthroat trout samples from South Fork Big Wash and Mill Creek. One locus that did amplify reliably, Fgt3, was not possible to score because of multiple peaks from pronounced stuttering; thus it was excluded from further analyses. The other 12 microsatellite loci showed no evidence of genotyping errors resulting from stuttering, null alleles, or large allele dropouts.

Twelve microsatellite loci showed varying levels of allelic diversity in the Mill Creek and South Fork Big Wash Bonneville cutthroat trout populations (Table 3). Some loci were monomorphic despite being variable in test runs on other Bonneville cutthroat trout populations (DDH unpublished data). The Mill Creek samples contained 4 monomorphic loci (H118, J14, J132, Oc18), and the South Fork Big Wash samples contained 3 monomorphic loci (H118, J14, J132). The number of alleles per locus in each population ranged from 1 to 9 (Table 3). The aforementioned monomorphic loci accounted for the low end of that range, and the loci with the highest amount of allelic
Table 2. Microsatellite primers that successfully amplified Bonneville cutthroat trout microsatellite loci from South Fork Big Wash and Mill Creek.

<table>
<thead>
<tr>
<th>Locus</th>
<th>Annealing temperature</th>
<th>Primer sequencea</th>
<th>Reference for unmodified primers</th>
</tr>
</thead>
<tbody>
<tr>
<td>Fgt3</td>
<td>54 °C</td>
<td>F: CACGACGTTGTAAACGACCAAGAAATTGTGGAGCCGG&lt;br&gt;R: GAAGCCCTGTTTCACTTATTAGC</td>
<td>Sakamoto et al. 1994</td>
</tr>
<tr>
<td>H18</td>
<td>57 °C</td>
<td>F: CACGACGTGTGAAAAACGACCAAAACATTGCCTCTCTGTAC&lt;br&gt;R: CCAACCTGTTAAATTTGCTAC</td>
<td>Pritchard et al. 2007</td>
</tr>
<tr>
<td>H118</td>
<td>57 °C</td>
<td>F: CACGACGTGTGAAAAACGACCTGCTCAATTTGATCTTG&lt;br&gt;R: GGCACTCTTTTATGAAAGCC</td>
<td>Pritchard et al. 2007</td>
</tr>
<tr>
<td>H126</td>
<td>57 °C</td>
<td>F: CACGACGTGTGAAAAACGACCAACCCCTCCCTCAATTC&lt;br&gt;R: ACCACACCTCCAGTC</td>
<td>Pritchard et al. 2007</td>
</tr>
<tr>
<td>J14</td>
<td>57 °C</td>
<td>F: CACGACGTGTTAAAACGACAGAGACTGCAGAGCTAC&lt;br&gt;R: GCAAAAAAGACAGAGAAGG</td>
<td>Pritchard et al. 2007</td>
</tr>
<tr>
<td>J132</td>
<td>57 °C</td>
<td>F: CACGACGTGTTAAAACGACAGAGACTGCAGAGCTAC&lt;br&gt;R: GCAAAAAAGACAGAGAAGG</td>
<td>Pritchard et al. 2007</td>
</tr>
<tr>
<td>K216</td>
<td>57 °C</td>
<td>F: CACGACGTGTTAAAACGACAGAGACTGCAGAGCTAC&lt;br&gt;R: GCAAAAAAGACAGAGAAGG</td>
<td>Pritchard et al. 2007</td>
</tr>
<tr>
<td>K222</td>
<td>57 °C</td>
<td>F: CACGACGTGTTAAAACGACAGAGACTGCAGAGCTAC&lt;br&gt;R: GCAAAAAAGACAGAGAAGG</td>
<td>Pritchard et al. 2007</td>
</tr>
<tr>
<td>Oc18</td>
<td>60 °C</td>
<td>F: CACGACGTGTTAAAACGACAGAGACTGCAGAGCTAC&lt;br&gt;R: GCAAAAAAGACAGAGAAGG</td>
<td>Pritchard et al. 2007</td>
</tr>
<tr>
<td>Omm1036</td>
<td>60 °C</td>
<td>F: CACGACGTGTTAAAACGACAGAGACTGCAGAGCTAC&lt;br&gt;R: GCAAAAAAGACAGAGAAGG</td>
<td>Rexroad et al. 2002</td>
</tr>
<tr>
<td>Omm1241</td>
<td>60 °C</td>
<td>F: CACGACGTGTTAAAACGACAGAGACTGCAGAGCTAC&lt;br&gt;R: GCAAAAAAGACAGAGAAGG</td>
<td>Rexroad et al. 2002</td>
</tr>
<tr>
<td>Ots107</td>
<td>54 °C</td>
<td>F: CACGACGTGTTAAAACGACAGAGACTGCAGAGCTAC&lt;br&gt;R: GCAAAAAAGACAGAGAAGG</td>
<td>Nelson and Beacham 1999</td>
</tr>
</tbody>
</table>

aM13 sequence in bold
<table>
<thead>
<tr>
<th>Locus</th>
<th>Repeat unit</th>
<th>N</th>
<th>N_A</th>
<th>Allele size range</th>
<th>H_O</th>
<th>H_E</th>
<th>HW</th>
</tr>
</thead>
<tbody>
<tr>
<td></td>
<td></td>
<td>BW</td>
<td>MC</td>
<td>BW</td>
<td>MC</td>
<td>BW</td>
<td>MC</td>
</tr>
<tr>
<td>H18</td>
<td>(TAGA)_{25}</td>
<td>68</td>
<td>68</td>
<td>2</td>
<td>2</td>
<td>189–193</td>
<td>189–193</td>
</tr>
<tr>
<td>H118</td>
<td>(TAGA)_{16}TAGA</td>
<td>68</td>
<td>68</td>
<td>1</td>
<td>1</td>
<td>181</td>
<td>181</td>
</tr>
<tr>
<td>H126</td>
<td>(GATA)_{21}</td>
<td>66</td>
<td>68</td>
<td>2</td>
<td>2</td>
<td>216–218</td>
<td>216–218</td>
</tr>
<tr>
<td>J14</td>
<td>(GATA)_{13}</td>
<td>68</td>
<td>66</td>
<td>1</td>
<td>1</td>
<td>223</td>
<td>223</td>
</tr>
<tr>
<td>J132</td>
<td>(GATA)_{10}</td>
<td>68</td>
<td>68</td>
<td>1</td>
<td>1</td>
<td>181</td>
<td>181</td>
</tr>
<tr>
<td>K216</td>
<td>(CATA)<em>{8}(CGTA) (CATA)</em>{8}(CGTA)</td>
<td>66</td>
<td>56</td>
<td>4</td>
<td>9</td>
<td>179–254</td>
<td>181–254</td>
</tr>
<tr>
<td>K222</td>
<td>(CATA)<em>{10}(TATA) (CATA)</em>{8}(TATA) (CATA)_{24}</td>
<td>66</td>
<td>66</td>
<td>9</td>
<td>4</td>
<td>121–134</td>
<td>121–134</td>
</tr>
<tr>
<td>Oc18</td>
<td>(GT)_{16}</td>
<td>58</td>
<td>66</td>
<td>2</td>
<td>1</td>
<td>180–184</td>
<td>184</td>
</tr>
<tr>
<td>Onm1036</td>
<td>(TATC)_{38}</td>
<td>68</td>
<td>66</td>
<td>2</td>
<td>2</td>
<td>217–229</td>
<td>217–229</td>
</tr>
<tr>
<td>Onm1241</td>
<td>(TAGA)<em>{11}(GAA)</em>{31}</td>
<td>64</td>
<td>66</td>
<td>2</td>
<td>2</td>
<td>154–158</td>
<td>154–158</td>
</tr>
<tr>
<td>Ots107</td>
<td>(GATA)(GT)(GTCT)</td>
<td>68</td>
<td>66</td>
<td>4</td>
<td>6</td>
<td>196–222</td>
<td>196–222</td>
</tr>
<tr>
<td>Ssa55</td>
<td>(GT)_{14}</td>
<td>66</td>
<td>66</td>
<td>2</td>
<td>2</td>
<td>145–159</td>
<td>145–159</td>
</tr>
</tbody>
</table>
Table 4. Results of bottleneck analysis: tests using the infinite alleles model (I.A.M.) and the stepwise mutational model (S.M.M.) are summarized herein. Statistically significant values are shown in bold font.

<table>
<thead>
<tr>
<th>Test</th>
<th>Assumptions: all loci fit I.A.M., mutation-drift equilibrium</th>
<th>Assumptions: all loci fit S.M.M., mutation-drift equilibrium</th>
</tr>
</thead>
<tbody>
<tr>
<td>Sign test</td>
<td>Expected number of loci with heterozygote excess: 3.55</td>
<td>Expected number of loci with heterozygote excess: 3.56</td>
</tr>
<tr>
<td></td>
<td>2 loci with heterozygosity deficiency, 5 loci with heterozygosity excess</td>
<td>3 loci with heterozygosity deficiency, 4 loci with heterozygosity excess</td>
</tr>
<tr>
<td>Probability</td>
<td>Probability: 0.23419</td>
<td>Probability: 0.51792</td>
</tr>
<tr>
<td></td>
<td>Assumptions: all loci fit S.M.M., mutation-drift equilibrium</td>
<td>Assumptions: all loci fit S.M.M., mutation-drift equilibrium</td>
</tr>
<tr>
<td></td>
<td>Expected number of loci with heterozygote excess: 3.75</td>
<td>Expected number of loci with heterozygote excess: 3.78</td>
</tr>
<tr>
<td></td>
<td>4 loci with heterozygosity deficiency and 3 loci with heterozygosity excess</td>
<td>5 loci with heterozygosity deficiency and 2 loci with heterozygosity excess</td>
</tr>
<tr>
<td>Probability</td>
<td>Probability: 0.42345</td>
<td>Probability: 0.16479</td>
</tr>
<tr>
<td>Standardized differences test</td>
<td>Assumptions: all loci fit I.A.M., mutation-drift equilibrium</td>
<td>Assumptions: all loci fit S.M.M., mutation-drift equilibrium</td>
</tr>
<tr>
<td>T2: 2.338 (P = 0.00969)</td>
<td>T2: 0.972 (P = 0.16549)</td>
<td>T2: 1.119 (P = 0.13148)</td>
</tr>
<tr>
<td>Wilcoxon test</td>
<td>Assumptions: all loci fit I.A.M., mutation-drift equilibrium</td>
<td>Assumptions: all loci fit I.A.M., mutation-drift equilibrium</td>
</tr>
<tr>
<td>Probability (1 tail for H deficiency): 0.98828</td>
<td>Probability (1 tail for H deficiency): 0.85156</td>
<td>Probability (1 tail for H deficiency): 0.28906</td>
</tr>
<tr>
<td>Probability (1 tail for H excess): 0.01953</td>
<td>Probability (1 tail for H excess): 0.18750</td>
<td>Probability (1 tail for H excess): 0.81250</td>
</tr>
<tr>
<td>Probability (2 tails for H excess and deficiency): 0.03906</td>
<td>Probability (2 tails for H excess and deficiency): 0.37500</td>
<td>Probability (2 tails for H excess and deficiency): 0.46875</td>
</tr>
</tbody>
</table>
diversity were K216 in the Mill Creek population and K222 in the South Fork Big Wash population. The expected heterozygosity ranged from 0.187 to 0.766, and the observed heterozygosity ranged from 0.040 to 0.900 (excluding the monomorphic loci for which both observed and expected heterozygosity was zero). The population pairwise $F_{ST}$ estimate between Mill Creek and South Fork Big Wash was 0.06727 ($P < 0.00001$).

Population Genetic Analyses

Tests for Hardy–Weinberg equilibrium could not be performed on monomorphic loci, but significant deviations from Hardy–Weinberg equilibrium were detected for some of the polymorphic loci (Table 3). Mill Creek had 3 loci that were not in Hardy–Weinberg equilibrium (H18, H126, K216). South Fork Big Wash also had 3 loci that were not in Hardy–Weinberg equilibrium (H126, K222, Oc18). Lower South Fork Big Wash had one locus that was not in Hardy–Weinberg equilibrium (H126). Hendrys Creek had 2 loci that were not in Hardy–Weinberg equilibrium (H126, K222).

The results of BOTTLENECK analysis revealed that the Mill Creek population shows some signs of experiencing a recent population bottleneck, whereas the South Fork Big Wash population does not. The Wilcoxon one-tailed test for heterozygote excess significantly deviated from mutation-drift equilibrium under the infinite alleles model ($P = 0.01953$), and the Wilcoxon 2-tailed test for heterozygote excess and deficiency also revealed a significant departure from mutation-drift equilibrium ($P = 0.03906$) under the infinite alleles model. The standardized differences test also showed a significant departure from mutation-drift equilibrium ($P = 0.00969$) under the infinite alleles model. However, this pattern did not hold for tests under the stepwise mutational model, as none of those results showed significant departure from mutation-drift equilibrium for the Mill Creek population.

DISCUSSION

The results of STRUCTURE and STRUCTURE-HARVESTER revealed 4 distinct populations of Bonneville cutthroat trout. However, these 4 groups of alleles were relatively evenly distributed in the Mill Creek and South Fork Big Wash populations (Fig. 2). This observed pattern was likely driven by the monomorphic loci that occur in these 2 populations.

The Mill Creek and South Fork Big Wash populations of Bonneville cutthroat trout have low genetic diversity among the 12 microsatellite loci that were included in this study. The 2 populations differ in the number of alleles at only 4 of the 12 loci that were successfully genotyped, and allele size ranges were identical for all but 2 of the 12 loci (Table 3). The 2 loci that did differ in allele size ranges had unique alleles in the South Fork Big Wash population that were not detected in Mill Creek. The sharing of alleles between the 2 populations is not surprising considering that Mill Creek was used as a source for the establishment of the South Fork Big Wash population. However, the significant $F_{ST}$ value between the Mill Creek and South Fork Big Wash populations does indicate a lack of gene flow, suggesting that some genetic differentiation has occurred since the transplant took place, possibly resulting from genetic drift. This genetic differentiation is likely manifest in subtle differences in alleles and allele frequencies. It was somewhat surprising that, of the 4 loci that exhibited differences in the number of alleles between the populations, 2 loci (K222 and Oc18) each had unique alleles in the South Fork Big Wash population that were additional to the alleles in Mill Creek, the source population for the transplant (Table 3). Typically, transplanted populations experience a founder effect, wherein individuals exhibit lower genetic diversity than those in the source population (Mayr 1942).

It is possible that some alleles in either or both populations were not sampled for this study. In part, this could be further understood if the founding population for South Fork Big Wash Creek came from a broader area in Mill Creek than was sampled for the Mill Creek microsatellite study. That does not appear to be the case, since the founding South Fork Big
Wash population was transferred from the same location in Mill Creek that was sampled for this study (Mark Pepper, Great Basin National Park, personal communication).

Differential sampling of alleles may be associated with temporal differences in the stocking of fish into South Fork Big Wash Creek. The transfer of fish into the South Fork Big Wash Creek occurred over 2 years, one in July 2000 and the other in July of 2001. Different genotypes may have been present in the sampled area of Mill Creek during that period of time.

It is also possible that the source population in Mill Creek has undergone a population bottleneck after, or even because of, the removal of individuals to populate South Fork Big Wash. Mill Creek was the only population that exhibited a signature of a population bottleneck, although that result occurred only under the infinite alleles model, not the stepwise mutational model (Table 4). Both populations had numerous monomorphic microsatellite loci, which is typical for populations that have experienced high levels of inbreeding and/or genetic drift. These fixed loci could not be included in the BOTTLENECK analysis because their expected heterozygosity was zero, but they still indicate that the populations may have reduced genetic variation in comparison to other Bonneville cutthroat trout populations where these loci were polymorphic.

The Mill Creek and South Fork Big Wash Creek populations do show high similarity, as expected. They also show fixation at numerous loci, as expected in populations with reduced genetic variability. However, it is unclear whether inbreeding has occurred in these populations, or if so, whether the inbreeding is significant enough to raise concern about the onset of inbreeding depression. While much has been discussed regarding the problems with intense inbreeding (Wright 1921, Charlesworth and Charlesworth 1987, Mills and Smouse 1994, Hedrick and Kalinowski 2000, García-Fernández et al. 2012), relatively few cases of population loss due to such conditions have been documented in wild populations (e.g., Crnokrak and Roff 1999, Rosner 2012). The risk of loss of Bonneville cutthroat trout populations in Great Basin National Park can easily be assessed via regular population estimates in the streams, which is consistent with the idea that proper management is best handled on a case-by-case basis (Houde et al. 2011).

ACKNOWLEDGMENTS

We thank Mark Pepper (National Park Service) and Chris Crookshanks (Nevada Department of Wildlife) for providing the Bonneville cutthroat trout samples used in this study. Ed Wilcox prepared samples for sequencing once they were submitted to the BYU DNA Sequencing Center. The National Park Service provided the funding for this study.
LITERATURE CITED


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