# YKFP WDFW M\&E Report 

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## 1. Executive Summary

## a. Fish Population RM\&E

The Yakima/Klickitat Fisheries Project's (YKFP) monitoring and evaluation (M\&E) project was established to evaluate critical uncertainties associated with spring Chinook salmon supplementation in the Yakima Basin. The YKFP is co-managed by the Yakama Nation (lead entity) and the Washington Department of Fish and Wildlife with guidance from the Northwest Power Planning Council and is funded predominantly by the Bonneville Power Administration. The M\&E project historically, and is currently collecting information under several disciplines associated with the supplementation of spring Chinook salmon, including ecological interactions and ecological risk containment monitoring, domestication monitoring, genetic monitoring, competition/capacity/habitat saturation monitoring, natural production monitoring, and monitoring the relative reproductive success of fish in the program, consistent with the Columbia Basin’s Fish
and Wildlife Program. Results from the project have been presented in public and professional forums, and are intended to inform others throughout the region on the information learned under the project. The current investigations included in this report provide summarized results of ongoing studies and should be considered preliminary until published in the peer reviewed literature.

Status and trends in abundance, productivity, distribution, and diversity of spring Chinook, and non-target taxa were collected during this contract period. Preliminary results of ongoing studies suggests operating the YKFP's production program has provided a demographic benefit to the population, has not impacted valued fish taxa beyond acceptable levels and the risk containment monitoring program is working as planned, some small levels of domestication in the context of predation vulnerability and competitive dominance may have occurred although the evidence is not compelling, genetic stock partitioning of mixed stock smolt migrants remains a viable method to estimate population specific juvenile chinook smolt abundance and productivity, rearing habitat saturation has likely been met in several years under current conditions, very low levels of naturally produced precociously maturing chinook have been observed on the spawning grounds.

## b. Hatchery RM\&E

The YKFP M\&E project was established to monitor the progress of the Cle Elum Supplementation and Research Facility (CESRF) progress at meeting spring Chinook production and biological objectives established for the YKFP's production program. The objectives were explicitly stated in the YKFP's monitoring and evaluation plan (Busack et al. 1997) and more recently, as Quantitative Objectives for the project. The project Quantitative Objectives provide benchmark values against which the performance of the project can be monitored and evaluated relative to fixed standards. Quantitative objectives have been established for the Spring Chinook supplementation program and include objectives for natural production, harvest, genetics, ecology, habitat, and science. While all of these objectives evaluate the performance of the Yakima/Klickitat fisheries project at some level, we focused on the hatchery RM\&E strategy for natural production and uncertainties research in this report. Monitoring the demographic benefit of the supplementation program has been thoroughly described in the Yakama Nation annual reports. This report extends the findings to cover uncertainty research of factors that may limit supplementation success and the projects performance relative to the natural production, ecological, and genetic quantitative objectives. It should be noted that the program strategies of hatchery RM\&E and fish population RM\&E are highly intertwined under this project and are not necessarily independent from one another. The hatchery RM\&E evaluations included in this report provide summarized results of ongoing studies and should be considered preliminary until published in the peer reviewed literature.

## 2. Introduction

The Yakima-Klickitat Fisheries Project is a cooperatively managed project with the Yakama Nation (YN; lead entity) and the Washington Department and Fish and Wildlife (WDFW) with supported in large part by the Bonneville Power Administration (BPA) with the oversight and guidance from the Northwest Power and Conservation Council (NPCC; Sampson et al. 2013). The Yakima Klickitat Fisheries Project’s (YKFP) Monitoring and Evaluation (M\&E) program has been described as the "Omnibus" scientific component of the broader YKFP (ISRP review 20060831). The M\&E project provides a rigorous assessment of the assumptions of supplementation and the application in the Yakima Basin to increase the natural production of salmon throughout the basin. The YKFP is an adaptively managed supplementation program designed "to test the assumption that new artificial production can be used to increase harvest and natural production while maintaining the long-term genetic fitness of the fish population being supplemented and keeping adverse genetic and ecological interactions with non-target species or stocks within acceptable limits" (BPA 1996). The M\&E project was designed to evaluate the YKFP progress towards addressing these four questions:

1) Can integrated hatchery programs be used to increase long-term natural production?
2) Can integrated hatchery programs limit genetic impacts to non-target Chinook populations?
3) Can integrated hatchery programs limit ecological impacts to non-target populations?
4) Does supplementation increase harvest opportunities?

This contract supports ongoing M\&E activities and research conducted by the Washington Department of Fish and Wildlife under the YKFP. The WDFW previously produced a minimum of 4 technical reports annually as deliverables under this contract (Competition/Capacity monitoring; Ecological Risk Containment monitoring; Genetics; Domestication Research and Monitoring) but with the new streamlined BPA reporting guidance and requirements, the reporting structure (and timelines) have been reduced to the summarized information herein. The work and reporting under each the topics of ecological interactions, domestication selection, competition/capacity, and genetic investigations are in varying stages of development and should be considered preliminary until published in the peer reviewed literature. Finally, this project has produced numerous publications that provide detailed evaluations under each topical research area (Appendix A).

## a. Fish Population RM\&E

F\&W Program Strategy: Assess the status and trend of adult natural and hatchery origin abundance of fish populations for various life stages.

F\&W Program Management Question: What are the status and trend of adult abundance of natural and hatchery origin fish populations?

Adult status and trend data for spring Chinook salmon are collected and presented annually under the Yakama Nation contracts and reports associated with the YKFP (Project 1995-063-25; Sampson et al. 2013).

Adult status and trend population data are also collected for non-target taxa (NTT) under the ecological risk containment monitoring program under this WDFW contract. The status and trends of 15 non-target taxa of concern (NTTOC) are collected annually to ensure the operation of the YKFP's production scale salmon supplementation program does not adversely affect the status of these taxa. Benchmark values were established during the pre-supplementation period and changes in the population status for these NTTOC are judged relative fixed standards termed containment objectives. Acceptable levels of change were established as containment objectives for each NTTOC under the project (Pearsons et al. 1998) and change in the population status are monitored within the risk containment and adaptive management framework (Temple and Pearsons 2012).

F\&W Program Strategy: Assess the status and trend of juvenile abundance and productivity of natural origin fish populations.

F\&W Program Management Question: What are the status and trend of juvenile abundance and productivity of fish populations?

## Non-target taxa of concern monitoring

The general approach for assessing and containing risks to non-target taxa of concern (NTTOC) that may result from supplementation of spring Chinook, and reintroduction of coho salmon, in the upper Yakima River basin was most recently detailed in Temple and Pearsons (2012), culminating from years of development (Pearsons 1998; Pearsons and Hopley 1999; Ham and Pearsons 2000; Pearsons 2002; Temple and Pearsons 2012). Briefly, we use the sieve approach to risk containment monitoring. First, overlap in the distribution between the target taxa (spring Chinook and coho salmon, and NTT) are identified. Distributional overlap that exceed acceptable levels, termed containment objectives (Pearsons 1998), warrant more rigorous evaluation, and we proceed to the next step in the evaluation. Second, when spatial and temporal overlap exceeds the containment objectives, a before/after comparison of monitoring variables (abundance, size, biomass) is evaluated. In cases where before/after comparisons of monitoring variable for a given NTTOC exceed acceptable levels, we proceed with the next step in the evaluation; a causation analysis in attempt to determine the cause of the decline. Causation analysis generally consists of rigorous evaluation of the data, but may include additional experiments or focused research to determine the mechanism for declines in NTTOC monitoring variables. Finally, the risk containment analysis is conducted under the adaptive management framework such that adjustments to the supplementation or reintroduction program will be made to alleviate unacceptable impacts to NTTOC.

## Spring Chinook habitat saturation and limiting factors

The status and trend of juvenile spring Chinook salmon abundance and productivity are collected annually for rearing spring Chinook (target-taxa) under the spring Chinook competition/carrying capacity program under the YKFP’s, M\&E contract. The carrying capacity of a watershed is an important factor in determining whether supplementation is a viable technique of increasing natural production. In the Yakima River Basin, carrying capacity can limit the number of naturally produced spring

Chinook salmon even when supplementation mechanics are operating perfectly (Busack et al. 1997). Preliminary analysis suggests that density dependent mechanisms affecting spring Chinook survival exist in the upper Yakima River after fall spawning and prior to or during the parr stage the following fall (Johnson et al. 2009). If the Yakima River is at capacity for rearing sub-yearlings in some years, then supplementation efforts can only serve to increase the number of naturally produced smolts when natural production is below that capacity. Therefore, identifying the factors that limit natural production is critical if restoration efforts aimed at maintaining or increasing natural production are to achieve their intended biological goals. The spring Chinook habitat saturation and limiting factors work aims to identify juvenile life-stage survival bottlenecks that may limit supplementation success in some years.

## Spring Chinook residual/precocious male monitoring

Artificial propagation of Chinook salmon has the potential to alter the age that fish mature and result in undesirable interactions with natural origin fish (Knudsen et al. 2006). This is a particular concern for conservation hatcheries where the goal is to increase natural production while maintaining the characteristics of the natural population (Mobrand et al. 2005). Although most Chinook salmon are anadromous (Healey 1991), some salmon complete their entire life cycle in freshwater, even when they have access to the ocean. These salmon are generally small, male, precociously mature, short-lived and are referred to as residents, precocious males, or minijacks (Gebhards 1960; Mullan et al. 1992; Zimmerman et al. 2003). The occurrence of precocity in salmon has been credited to genetic factors and environmental and physiological cues (Thorpe 1987; Bohlin et al. 1990; Foote et al. 1991). Age-at-maturation has been shown to be heritable in salmon (Heath et al. 1994; Unwin et al. 1999); and although it has been known for some time that hatcheries can produce large numbers of precocious Chinook salmon (Robertson 1957; Mullan et al. 1992; Larsen et al. 2004a; Beckman and Larsen 2005), there have been relatively few studies that have investigated the abundance and distribution of these fish in rivers during the spawning season. Previous research indicated that the Yakima Supplementation and Research Facility had produced and released an average of 129,249 precocious males/year into the upper Yakima basin between 1999 and 2008 (Larsen et al. 2004a; Larsen et al. 2008; Yakima/Klickitat Fisheries Project, Unpublished data). Our primary objectives are to 1 ) estimate the annual abundance of hatchery origin precocious males on the spawning grounds, and 2) quantify the annual distribution of hatchery precocious males on and away from the spawning grounds. We also present information about the abundance and distribution of natural origin precocious males so that we can determine how hatchery precocious males might differ.

## Spring Chinook reproductive success/spawning channel

Although hatcheries have been extensively utilized in Chinook salmon management for over 100 years, only recently have rigorous experiments been developed to measure the relative reproductive success of hatchery- and natural-origin spawners in a shared natural setting. Some of the difficulty in designing informative studies has stemmed from the challenges of controlling entry to natural spawning areas and collecting representative samples of recently hatched fry. Furthermore, if control could
be established over the potential spawners in the spawning area, the measurement of individual reproductive output still would require a means of associating individual fish captured in one year with individuals that spawned in a previous year. The spawning behavior of Chinook salmon adds to the complexity of quantifying individual reproductive output through behavioral observations: at a redd site, a female might be courted by several males that compete for access to the female, providing opportunities for multiple paternity in a single redd. In areas with moderate to high spawning densities, males might attend females on several adjacent redds. Microsatellites, a class of highly polymorphic, codominant DNA markers, provide a means to quantify individual spawners' reproductive output. A suite of 10 to 15 highly variable microsatellites can resolve individual identity in a moderate to large population, and through a simple inheritance model, can illuminate parent-offspring relationships.

Washington Department of Fish and Wildlife (WDFW) and the Yakama Nation (YN) are cooperating on a study of Chinook salmon reproductive success in a presumably closed access spawning observation channel at the Cle Elum Hatchery. Viewing blinds line the channel, allowing researchers to observe spawning activities.

Chinook salmon carrying visible external marks were released into the spawning channel in September 2012. Hatchery-control line (two generations of hatchery influence) males and females were released into three of six shared spawning areas and supplementation hatchery line (one generation of hatchery influence) males and females were released into the other three shared spawning areas to select and compete for mates. Prior to the release of the potential spawners, researchers collected and preserved samples of fin tissue to enable genetic characterization of the potential spawners and to allow subsequent inference of parent/offspring relationships after juveniles were collected and genotyped. One group of researchers examined morphological characteristics of these potential parents and observed and recorded spawning area behaviors and interactions. The results of the morphological and behavioral work are described in a separate report.

The potential parents' fin tissue samples and the collected progeny (fry) were delivered to the WDFW Molecular Genetics Laboratory in Olympia, Washington for genetic screening and parentage analysis following the same protocols that have been used from 2002 - 2007, 2009 - 2013 (Young and Kassler 2005, Kassler 2005, Kassler 2006, Kassler and Von Bargen 2007, 2008, and 2010, Kassler et al. 2011; Kassler and Peterson 2012, 2013). The genetic analyses provide direct, quantitative estimates of fry production by individual spawning Chinook salmon. This report presents the parentage results for the 2012 - 2013 Cle Elum spawning channel experiments.

## Spring Chinook Genetic stock separation-juveniles

Production and survival of the Yakima River basin spring Chinook stocks (American River, Naches River, and upper Yakima River) are monitored, as part of the Yakima/Klickitat Fishery Project supplementation evaluation program. However, in the lower Yakima River, where the best facilities to collect samples exist, the three spring Chinook stocks are mixed with one another and with the Marion Drain and Yakima River fall Chinook stocks, during downstream juvenile migration. Thus, methodologies for
discriminating stocks in an admixture are vital for development of stock-specific estimates. Domestication monitoring plans require discrimination of the three spring Chinook salmon stocks in the basin, and a complete analysis of migration timing and stock abundance for all Chinook requires discrimination of the two fall stocks as well. Accurate assignments of Chinook smolts captured at the Chandler fish passage facility to population-of-origin will allow researchers and managers to estimate production by the three spring Chinook stocks, assess smolt-to-smolt survival of the three spring Chinook stocks, and could be utilized to evaluate stock-specific environmental condition factors.

F\&W Program Strategy: Assess the status and trend of spatial distribution of fish populations.

F\&W Program Management Question: What are the status and trend of spatial distribution of fish populations?

The spatial distribution of adult Spring Chinook salmon (target taxa) are best described in the Yakama Nation's annual reports (Sampson et al. 2013) where the spawning distribution for spring Chinook is intensively monitored and reported annually.

The spatial distribution of rearing naturally produced spring Chinook in the upper Yakima basin is monitored under the ecological risk containment monitoring program and is of interest in the context of distributional overlap with non-target taxa of concern. Lack of spatial overlap between the target taxa and non-target taxa are thought to preclude negative effects of species interactions. The distribution of spring Chinook is monitored annually in tributary and mainstem Yakima River index monitoring sites. The ecological effects of distributional overlap with NTTOC are currently monitored in the risk containment monitoring framework (Temple and Pearsons 2012).

Spatial distribution of early rearing spring Chinook in the upper Yakima basin is monitored under the spring Chinook Competition/Capacity program. Previous work in the upper Yakima River (Johnson et al. 2009) has suggested density dependent constraints to spring Chinook production prior to fall estimates of abundance. One primary objective of this program is to identify life-stage specific factors limiting to survival and development in the natural environment. Such data can then be used to educate management decisions in selecting actions to most effectively increase natural production.

F\&W Program Strategy: Assess the status and trend of diversity of natural and hatchery origin fish populations.

F\&W Program Management Question: What are the status and trend of diversity of natural and hatchery origin fish populations?

Operating a production scale supplementation program may have unintended effects that alter the diversity of both natural and hatchery origin fish populations through selective forces imposed by the hatchery environment. The domesticating effects of hatchery culture are being intensively monitored for spring Chinook in the Yakima under the Domestication monitoring program. The YKFP's domestication monitoring plan was
developed to determine if the spring Chinook supplementation program affects a large number of phenotypic and morphometric traits of the Yakima population (Busack et al. 2006.

## Domestication Predation and competitive dominance description

Raising fish in hatcheries can cause unintended behavioral, physiological, or morphological changes in Chinook salmon due to domestication selection. Domestication selection is defined by Busack and Currens 1995 as: "changes in quantity, variety, or combination of alleles within a captive population or between a captive population and its source population in the wild as a result of selection in an artificial environment." Selection in artificial environments could be due to intentional or artificial selection, biased sampling during some stage of culture, or unintentional selection (Busack and Currens 1995). Genetic changes can result in lowered survival in the natural environment (Reisenbichler and Rubin 1999). The goal of supplementation or conservation hatcheries is to produce fish that will integrate into natural populations and increase the number of grandchildren relative to fish that live entirely in natural environments. Conservation hatcheries attempt to minimize intentional or biased sampling so that the hatchery fish are similar to naturally produced fish. However, the selective pressures in hatcheries are dramatically different than in the wild, which can result in genetic differences between hatchery and wild fish. The selective pressures may be particularly prominent during the freshwater rearing stage where most mortality of wild fish occurs. We are attempting to evaluate the effects of domestication on the vulnerability of spring Chinook to predators, and on competitive dominance of spring Chinook salmon.

## b. Hatchery RM\&E

F\&W Program Strategy: Evaluate the effectiveness of hatchery safety-net/conservation programs and the effectiveness of hatchery reform actions on the achievement of biological performance objectives.

F\&W Program Management Question: Are hatchery improvement programs and actions achieving the expected biological performance objectives?

The YKFP has a long history built upon a strong foundation of hatchery RM\&E. The larger YKFP was built upon developing responsible hatchery operations and production protocols consistent with many of the general (and specific) hatchery reform actions and recommendations that have recently been advised by the Hatchery Scientific Review Group for many hatchery programs throughout the Columbia basin and much of the Pacific Northwest. Much of the hatchery effectiveness monitoring information is presented in the Yakama Nation's annual technical report of the YKFP (Sampson et al. 2013). The YKFP established a long list of performance measures, termed quantitative objectives, and the project's performance relative to these standards are monitored and reported annually (Fritts 2012; Appendix C).

F\&W Program Strategy: Assess and investigate as appropriate critical uncertainties regarding the effects of artificial propagation on the viability of wild fish populations.

F\&W Program Management Question: What deleterious effects does artificial production have on natural populations of anadromous fish?

This M\&E project was founded upon monitoring and evaluating the effects of artificial production on natural populations and anadromous fish. The monitoring tasks described throughout this report covering the disciplines of domestication, genetics, ecological investigations, and competition/capacity work all strive towards answering critical uncertainties associated with artificial production in the Yakima Basin. Results from this work are intended to inform others throughout the Columbia River Region.

## 3. Methods: Protocols, Study Designs, and Study Area

Protocol Title: Ecological Interactions (1995-063-25) v1.0
Protocol Link: http://www.monitoringmethods.org/Protocol/Details/113
Protocol Title: Genetics (1995-063-25) v1.0
Protocol Link: http://www.monitoringmethods.org/Protocol/Details/115
Protocol Title: Natural Production (1995-063-25) v1.0
Protocol Link: http://www.monitoringmethods.org/Protocol/Details/116

## 4. Results

## a. Fish Population RM\&E

Non-target taxa of concern monitoring

## General approach

The inclusion of 2014 NTTOC monitoring data in the risk containment monitoring sieve evaluation provide similar results to last year. The degree of trout overlap with salmon was highest in main stem areas, intermediate for cutthroat and rainbow trout in tributaries, and absent for bull trout (Figure 1). There was no overlap of salmon and bull trout in our index sites. In fact, the shortest distance between the uppermost distribution of Chinook salmon and the lowermost distribution of bull trout was approximately 8 km . Cutthroat trout and supplemented spring Chinook overlapped in distribution in both tributary and main stem Yakima River areas. The distributional overlap in tributary streams was approximately $11 \%$, confined to relatively moderate elevations, and was less than the $40 \%$ containment objective (Figure 1). Salmon overlapped $100 \%$ of the main stem distribution of cutthroat trout (Figure 1). In tributaries, salmon overlapped $50 \%$ of the distribution of rainbow trout. Overlap was predominately confined to lower portions of tributaries (e.g., Swauk Creek 1 and Umtanum Creek 1) and farther upstream in the North Fork Teanaway River. However, salmon did not overlap rainbow trout in high elevation portions of tributaries.

There was also extensive overlap between rainbow trout, sucker species, and mountain whitefish and salmon in the main stem (100\%; Figure 1). Salmon overlapped in distribution with longnose dace (59\%) and speckled dace (72\%) in tributaries, although mean overlap was less than the containment objectives for both species. Salmon overlapped sculpin species $17 \%$ in tributaries but this was less than the containment objective. Finally, there was $23 \%$ overlap in distribution between sucker species and salmon in tributary streams, although this was also less than the containment objective.

Data that were collected at similar times and sites by snorkeling and electrofishing methods were consistent with each other. For example, in areas that we found salmon, rainbow trout, cutthroat trout or bull trout, they were detected with both electrofishing and snorkeling methods. In addition, we did not capture any salmon when we electrofished areas where bull trout were present.


Figure 1. Map of species distributions in the upper Yakima Basin. Spring Chinook and coho salmon distributions are shaded grey. The lowest elevation observations of bull trout and cutthroat trout in tributary streams are marked with stars and bars, respectively. Cutthroat trout, suckers and mountain whitefish distribution in the main stem is marked as a dashed line. The Cle Elum hatchery facility is marked with a black square and hatchery acclimation sites are marked with open squares. Rainbow trout are widely distributed throughout the basin and are not marked on the map.

## Before-After Analysis

Rainbow (age 1) and cutthroat trout (<250mm), mountain whitefish, and sucker species in the main stem, and rainbow trout in tributaries (all ages; analog for steelhead) exhibited distributional overlap with salmon that were outside the containment objectives and therefore we compared their abundance, size, and biomass (salmonids) before and after stocking began. The mean abundance and $90 \%$ CL of sympatric rainbow trout (all ages) was $32 \pm 16 \%$ higher in the tributaries and $36 \pm 16 \%$ higher in the main stem (age 1) in the years when supplementation occurred than during the baseline phase (Figure 2). The mean abundance of cutthroat trout ( $<250 \mathrm{~mm}$ ) was $565 \pm 396 \%$ CL higher in the main stem during supplementation than during the baseline phase (Table 1; Figure 2). The mean abundance of sub-adult mountain whitefish increased $109 \pm 40 \%$ CL during supplementation period, while the mean abundance of sucker species adults decreased 44 $\pm 6 \%$ CL and the decrease was significant ( $P<0.01$ ), although it was within our containment objectives (Figure 3). Finally, we observed a $26 \pm 18 \%$ CL increase in subadult sized sucker abundance (analog for mountain sucker) during supplementation and the lower $90 \%$ CL did not exceed our containment objective (Figure 3).

During the supplementation period, the mean and $90 \%$ CL of rainbow trout size (age 1) in the main stem indicated that size decreased by $5 \pm 2 \%$ (Table 1; Figure 2). Slopes between log length-log weight of age 1 rainbow trout in the main stem were not significantly different before and during supplementation ( $P=0.82$ ). An ANCOVA revealed the average weight of fish for a given length was significantly greater during the supplementation period ( $P=0.002$, Figure 4). In addition, biomass increased by $18 \pm$ $15 \%$ CL. Similarly, the mean and $90 \%$ CL of cutthroat trout size ( $<250 \mathrm{~mm}$ ) in the main stem indicated a $1 \pm 3 \%$ CL decrease, and an increase in biomass of $941 \pm 893 \%$ CL (Table 1; Figure 2). The size of rainbow trout in the tributaries (all ages) was similar during both periods ( $1 \pm 2 \%$ CL; Table 2; Figure 2). Slopes between log length-log weight for rainbow trout in tributaries (all ages) were not significantly different before and during supplementation $(P=0.34)$. An ANCOVA indicated the mean weights at each length were slightly greater during the supplementation period than the before period, although not significantly so ( $P=0.06$; Figure 4). Additionally, tributary rainbow trout biomass (all ages) increased by $27 \pm 11 \%$ CL (Table 2; Figure 2). Our index of mountain whitefish size indicated that the proportions of subadults observed increased 10 $\pm 2 \%$ CL during the supplementation period (Figure 3). Our index of sucker species size indicated that the proportion of adults decreased $41 \pm 12 \%$ during supplementation, and although the decrease was significant ( $P<0.001$ ), it was still well within our containment objectives (Figure 3). Our index of mountain sucker size indicated a $26 \pm 8 \%$ CL increase in the proportion of subadults during the supplementation period (Figure 3).

The mean abundance, size, and biomass of catchable sized main stem rainbow trout (>249 mm) did not decrease during supplementation. The mean abundance of rainbow trout greater than 249 mm increased by $13 \pm 14 \%$ (mean $\pm 90 \% \mathrm{CL}$ ), mean size increased by $2 \pm 1 \%$, and biomass increased by $41 \pm 16 \%$ during supplementation when compared to baseline conditions.

The only NTT with parameter estimates outside of the containment objectives was steelhead, which uses rainbow trout as an analog. The lower $90 \%$ CL for age 1
rainbow trout size in the main stem and rainbow trout size (all ages) in the tributaries were exceeded, so we tested whether the decrease was caused by supplementation.

Table 1. Annual abundance (fish/km), size (mm, FL), and biomass (kg/km) estimates and associated $95 \%$ confidence intervals of age 1 rainbow trout and cutthroat trout less than 250 mm fork length in the main stem Yakima River.

| Year | Abundance |  | Size |  | Biomass |  |
| :---: | :---: | :---: | :---: | :---: | :---: | :---: |
|  | RBT | CUT | RBT | CUT | RBT | CUT |
| 1990 |  |  | $210 \pm 33$ | $237 \pm 5$ |  |  |
| 1991 | $189 \pm 67$ | $11 \pm 14$ | $205 \pm 27$ | $237 \pm 11$ | $19 \pm 14$ | $1.6 \pm 3.2$ |
| 1992 | $151 \pm 28$ | 1 | $217 \pm 31$ | 242 | $18 \pm 7$ | 0.1 |
| 1993 | $193 \pm 48$ | $6 \pm 17$ | $232 \pm 36$ | $238 \pm 3$ | $27 \pm 11$ | $0.8 \pm 3.5$ |
| 1994 | $180 \pm 33$ | $2 \pm 1$ | $217 \pm 32$ | $225 \pm 17$ | $21 \pm 8$ | $0.3 \pm 1.4$ |
| 1995 | $190 \pm 54$ | $6 \pm 17$ | $235 \pm 34$ | $239 \pm 6$ | $28 \pm 12$ | $0.9 \pm 3.5$ |
| 1996 | $182 \pm 27$ | $5 \pm 11$ | $217 \pm 32$ | $239 \pm 10$ | $22 \pm 7$ | $0.7 \pm 2.4$ |
| 1997 | $272 \pm 49$ | $10 \pm 44$ | $203 \pm 35$ | $239 \pm 5$ | $27 \pm 10$ | $1.4 \pm 8.9$ |
| 1998 | $130 \pm 20$ | $16 \pm 84$ | $212 \pm 34$ | $230 \pm 5$ | $15 \pm 6$ | $2.0 \pm 16.8$ |
| 1999 | $182 \pm 25$ | $12 \pm 25$ | $217 \pm 33$ | $236 \pm 5$ | $22 \pm 7$ | $1.8 \pm 5.1$ |
| 2000 | $214 \pm 40$ | $13 \pm 1$ | $210 \pm 36$ | $227 \pm 13$ | $24 \pm 10$ | $1.8 \pm 1.4$ |
| 2001 | $384 \pm 81$ | $18 \pm 85$ | $206 \pm 32$ | $238 \pm 7$ | $41 \pm 16$ | $2.5 \pm 17.1$ |
| 2002 | $207 \pm 39$ | $7 \pm 42$ | $203 \pm 31$ | $232 \pm 6$ | $20 \pm 9$ | $0.9 \pm 8.4$ |
| 2003 | $230 \pm 41$ | $10 \pm 34$ | $207 \pm 30$ | $234 \pm 7$ | $24 \pm 9$ | $1.3 \pm 7.0$ |
| 2004 | $275 \pm 19$ | $16 \pm 34$ | $223 \pm 32$ | $234 \pm 5$ | $35 \pm 15$ | $2.3 \pm 6.9$ |
| 2005 | $272 \pm 20$ | $28 \pm 142$ | $213 \pm 32$ | $229 \pm 5$ | $30 \pm 9$ | $3.4 \pm 28.6$ |
| 2006 | $150 \pm 12$ | $16 \pm 11$ | $216 \pm 34$ | $235 \pm 5$ | $17 \pm 7$ | $2.1 \pm 2.5$ |
| 2007 | $233 \pm 17$ | $22 \pm 35$ | $210 \pm 33$ | $233 \pm 5$ | $26 \pm 8$ | $3.1 \pm 7.1$ |
| 2008 | $264 \pm 26$ | $24 \pm 61$ | $204 \pm 33$ | $229 \pm 7$ | $26 \pm 9$ | $3.0 \pm 12.3$ |
| 2009 | $156 \pm 29$ | $44 \pm 138$ | $188 \pm 29$ | $231 \pm 3$ | $12 \pm 3$ | $5.8 \pm 27.8$ |
| 2010 | $233 \pm 48$ | $32 \pm 111$ | $197 \pm 36$ | $230 \pm 5$ | $21 \pm 7$ | $4.1+22.3$ |
| 2011 | $273 \pm 23$ | $39 \pm 63$ | $199 \pm 34$ | $227 \pm 4$ | $26 \pm 9$ | $5.0 \pm 12.8$ |
| 2012 | $270 \pm 30$ | $70 \pm 250$ | $192 \pm 33$ | $226 \pm 5$ | $23 \pm 8$ | $8.7 \pm 50.3$ |
| 2013 | $359+38$ | $237 \pm 335$ | $196 \pm 34$ | $290 \pm 9$ | $32 \pm 10$ | $75.6 \pm 68.0$ |
| 2014 | $342 \pm 46$ | $176 \pm 168$ | $206 \pm 34$ | $276 \pm 8$ | $36 \pm 11$ | $44.4 \pm 41.0$ |

Table 2. Annual abundance (fish/km), size (mm, FL), and biomass (kg/km) estimates and associated $95 \%$ confidence intervals for rainbow trout in Yakima River Basin tributary streams.

| Year | Abundance | Size | Biomass |
| :---: | :---: | :---: | :---: |
| 1990 | $241 \pm 129$ | $136 \pm 8$ | $8 \pm 13$ |
| 1991 | $204 \pm 102$ | $131 \pm 8$ | $6 \pm 8$ |
| 1992 | $375 \pm 240$ | $130 \pm 5$ | $11 \pm 24$ |
| 1993 | $317 \pm 158$ | $131 \pm 7$ | $9 \pm 17$ |
| 1994 | $328 \pm 129$ | $132 \pm 8$ | $11 \pm 15$ |
| 1995 | $213 \pm 118$ | $139 \pm 8$ | $7 \pm 14$ |
| 1996 | $165 \pm 109$ | $133 \pm 8$ | $5 \pm 11$ |
| 1997 | $294 \pm 119$ | $132 \pm 5$ | $8 \pm 11$ |
| 1998 | $442 \pm 174$ | $138 \pm 7$ | $15 \pm 25$ |
| 1999 | $288 \pm 175$ | $135 \pm 8$ | $12 \pm 27$ |
| 2000 | $318 \pm 135$ | $144 \pm 8$ | $11 \pm 21$ |
| 2001 | $464 \pm 178$ | $129 \pm 3$ | $12 \pm 17$ |
| 2002 | $321 \pm 131$ | $132 \pm 6$ | $10 \pm 15$ |
| 2003 | $291 \pm 142$ | $132 \pm 5$ | $8 \pm 14$ |
| 2004 | $243 \pm 135$ | $142 \pm 5$ | $9 \pm 15$ |
| 2005 | $349 \pm 163$ | $127 \pm 5$ | $9 \pm 16$ |
| 2006 | $434 \pm 171$ | $134 \pm 5$ | $13 \pm 20$ |
| 2007 | $368 \pm 153$ | $138 \pm 4$ | $12 \pm 18$ |
| 2008 | $331 \pm 166$ | $138 \pm 7$ | $11 \pm 19$ |
| 2009 | $256 \pm 123$ | $138 \pm 12$ | $9 \pm 19$ |
| 2010 | $548 \pm 243$ | $127 \pm 5$ | $15 \pm 25$ |
| 2011 | $486 \pm 215$ | $124 \pm 7$ | $12 \pm 20$ |
| 2012 | $490 \pm 163$ | $124 \pm 4$ | $13 \pm 15$ |
| 2013 | $571 \pm 232$ | $129 \pm 5$ | $16 \pm 24$ |
| 2014 | $282 \pm 139$ | $134 \pm 5$ | $10 \pm 14$ |



Figure 2. Abundance ( $\mathrm{n} / \mathrm{km}$ ), size ( FL mm ), and biomass ( $\mathrm{kg} / \mathrm{km}$ ) of tributary rainbow trout, main stem Yakima River rainbow trout (age 1) and cutthroat trout ( $<250 \mathrm{~mm}$ ) before and during supplementation. Main stem cutthroat trout abundance, size, and biomass are associated with the right $y$-axis. The horizontal dashed line represents the $0 \%$ containment objective (CO) for steelhead in the main stem and tributaries, and the $10 \%$ CO for mainstem cutthroat trout. The solid horizontal line represents the $10 \%$ CO for main stem rainbow trout and $40 \%$ CO for tributary rainbow trout. Error bars represent $90 \%$ confidence intervals.


Figure 3. Abundance (fish/km) and size (percent by size class) of mountain whitefish, suckers, and mountain suckers before and during supplementation. Error bars represent the $90 \%$ confidence interval. Dashed lines represent the $40 \%$ containment objectives for mountain whitefish, $90 \%$ for sucker species (Spp), and $5 \%$ for mountain suckers.


Figure 4. Mean length-weight relationships of tributary and age 1 main stem Yakima River rainbow trout before (1990-1998) and during (1999-2013) the supplementation period. Each data point represents the mean from a sample site.

## Causation

Since the lower 90\% confidence limit for our steelhead size index was exceeded in both the Yakima River main stem (age 1 rainbow trout) and Yakima Basin tributaries (all ages of rainbow trout), we tested if the changes could be reasonably attributed to supplementation. We did not detect a statistically significant decrease in our steelhead size index (age 1 rainbow trout; BACIP; $P=0.97$ ) in the main stem downstream from the Clark Flats acclimation facility. Interestingly, we did not detect a significant relationship between our steelhead abundance and size index relationship ( $\mathrm{R}^{2}=0.11$; $P=0.11$ ) suggesting density dependence was probably not influencing our steelhead size index. For tributary comparisons, we did not detect significant differences in our steelhead size index in comparisons between the North Fork Teanaway River down stream from the Jack Creek acclimation facility (treatment sites) and the West (BACIP; $P=0.11$ ) and Middle Fork (BACIP; $P=0.52$ ) Teanaway River reference sites. Additional comparisons of our steelhead size index in the main stem Teanaway River relative to the West and Middle Fork Teanaway River reference sites were not consistent with an impact (i.e. all changes were positive). Thus, at this time, the weight-of-evidence suggests declines in our steelhead size index are not likely the result of salmon supplementation activities in the basin.

Although the before vs. after comparisons of rainbow trout abundance did not indicate declines warranting a refined analysis of abundance, we erred on the side of caution and conducted the analysis given our concerns related to the depressed steelhead size index. A comparison of rainbow trout abundance in index monitoring sites located downstream from the Jack Creek Acclimation Facility (e.g., North Fork and Main stem Teanaway Rivers) relative to reference sites in the Middle and West Fork Teanaway Rivers revealed substantial reductions in the abundance of rainbow trout relative to the control streams (BACIP). We attempted to account for factors that may influence abundance such as movement and angler induced mortality. Motion activated cameras mounted in both treatment and reference sites during the open angling season in 2011 indicated that the reduction in abundance was probably not angler induced. In addition, we have not detected large scale movements of tagged rainbow trout between treatment and reference streams that would be consistent with a largescale displacement of trout. However, we do have evidence that the North Fork of the Teanaway River produces a higher proportion of anadromous steelhead smolt migrants than the reference streams and significant migrant production may contribute to reduced resident trout abundance. We will continue this investigation in the coming year in cooperation with the Yakima Steelhead VSP project.

## Spring Chinook habitat saturation and limiting factors

## Post-emergent growth

In a multiple year analysis (2009-2014), observed growth rate differed significantly among years (Homogeny of slopes model: $F_{5,192}=4.90, P<0.01$; Figure 5). Mean length, accounting for sampling date, was also detectably different among years (Separate slopes model: $F_{4,192}=5.22, P<0.01$ ). Post-hoc analysis revealed significantly greater
mean length in 2010 and in 2014 in comparison to 2009, which had the smallest observed mean length within the six-year dataset.


Figure 5. Comparison of mean growth rate among years in the upper Yakima River basin 2009-2012. The observed rate of growth was greater in 2010 (heavy dashed line) when compared to other survey years; 2009 (solid grey line), 2011 (smaller dashed line), 2012 (solid black line), 2013 (dashed grey line), and 2014 (dotted black line).

## Rearing abundance and habitat use

A total of 64 sites were surveyed in the two study reaches between July 7th and August 25th, 2014 (Table 3.) for a total of 128 total replicates. We did not detect a significant difference in sub-yearling Chinook density between study reaches in 2013 ( $t=$ $1.0, \mathrm{df}=126, P=0.31$ ), or in a multi-year analysis (ANOVA: $F_{1,1095}=0.03, P=0.90$ ). Mean density was significantly greater in 2010 when compared all other years (ANOVA: $F_{6,1090}=7.29, P<0.01$; Tukey post-hoc: $P<0.01$; Figure 6.). Summer and fall mean spring Chinook densities trended, but were not significantly correlated with our chosen alpha of $0.05\left(\mathrm{R}^{2}=0.56, P=0.06\right)$. Further, although the relationship is near significant, and the proportion of variation potentially explained is high, it appears as though the 2010 data point is highly influential in the relationship (Cook's D = 1.46). Therefore, additional data points, over a range of fall densities (e.g. between 150 and 200 spring Chinook per kilometer) will be required in order to fully evaluate any potential correlation between summer and fall estimates of spring Chinook rearing density.

We did not detect a significant difference in spring Chinook density among habitat types in 2014 (ANOVA: $F_{5,122}=0.64, P=0.67$; Table 3). However, a multi-year
analysis suggests greater relative densities in pool and deep riffle habitats (Tukey HSD, P < 0.01; Figure 7).

Table 3. Physical parameters of 2014 snorkeling survey sites by sampling reach.

| Habitat Classification | n | Mean site length (m) | SD | n | Site width (m) | SD |
| :--- | :---: | :---: | :---: | :---: | :---: | :---: |
| Easton |  |  |  |  |  |  |
| Deep Riffle | 4 | 35.0 | 4.1 | 4 | 18.9 | 3.2 |
| Glide | 11 | 80.7 | 21.0 | 11 | 20.2 | 7.2 |
| Pool | 5 | 47.4 | 16.0 | 5 | 20.9 | 3.6 |
| Rapid | 1 | 30.0 | $\mathrm{n} / \mathrm{a}$ | 1 | 20.3 | $\mathrm{n} / \mathrm{a}$ |
| Riffle | 6 | 54.8 | 17.4 | 6 | 19.2 | 6.7 |
| Run | 10 | 53.3 | 16.5 | 10 | 17.6 | 5.2 |
|  | Nelson |  |  |  |  |  |
| Deep Riffle | 3 | 53.3 | 23.1 | 3 | 25.9 |  |
| Glide | 11 | 89.4 | 17.7 | 11 | 31.3 | 1.3 |
| Pool | 3 | 43.3 | 12.6 | 3 | 22.1 | 2.5 |
| Rapid | 1 | 45.0 | $\mathrm{n} / \mathrm{a}$ | 1 | 29.0 | $\mathrm{n} / \mathrm{a}$ |
| Riffle | 85.0 | $\mathrm{n} / \mathrm{a}$ | 1 | 36.3 | $\mathrm{n} / \mathrm{a}$ |  |
| Run | 1 | 91.8 | 19.9 | 8 | 31.0 | 6.6 |



Figure 6. Multi-year analysis of observed abundance among years in two upper Yakima River study reaches with similar temperature and flow characteristics (Easton and Nelson). Significantly greater densities of spring Chinook sub-yearlings were detected in 2010.


Figure 7. Mean spring Chinook observed abundance by habitat type, 2008-2014. Error bars represent 95 percent confidence intervals.

Water temperatures during sampling ranged between 12.0 and 18.1 degrees Celsius (mean, 15.4; SD, 1.7). Temperatures at the time of sampling were not detectably different between survey reaches $(t=-0.05, \mathrm{df}=109, P=0.96$, or among habitat classifications (ANOVA: $F_{5,114}=0.01, P=0.96$ ). Overall, temperatures in 2014 were positively correlated with observed abundance of spring Chinook, but explained only five percent of the variation ( $\mathrm{n}=112, \mathrm{R}^{2}<0.05 P<0.02$ ). Visibility while sampling ranged between 0.5 and 3.2 meters (mean, 1.5; SD, 0.5 ) and was not significantly correlated with estimates of abundance ( $\mathrm{R}^{2}<0.01, P=0.87$ ).

Territory size (log transformed) was significantly correlated with fish fork length $(\mathrm{mm})\left(\mathrm{R}^{2}=0.29, P<0.01\right.$; Figure 8 ), and differed significantly among years (ANCOVA: $F_{8,329}=19.2, P<0.01$ ). Mean territory size, adjusted for length was highly correlated with estimates of fall abundance ( $\mathrm{R}^{2}<0.90, P<0.01$ ), and with redd counts (YKFP 2014) from the previous year ( $\mathrm{R}^{2}<0.58, P<0.03$ ). The proportions of feeding strikes were significantly different between categorical distances (1-4 body lengths) from the focal position (Friedman ANOVA: $\chi_{3}^{2}, 694=1166.9 ; P<0.01$; Figure 9). Agonistic strikes were also significantly different between categorical distances (1-4 body lengths) from the focal position (Friedman ANOVA: $\chi_{3}^{2}, 314=170.2, P<0.01$; Figure 9). The observed ratios of agonistic to feeding were not significantly different among grouped distances from the focal position (Friedman ANOVA: $\chi_{9,3}^{2}=7.1, P=0.07$; Figure 10).


Figure 8. Relationship between spring Chinook fork length (mm) and observed territory size in the spring and summer of 2006-2013 (black points) and 2014 (white points).


Figure 9. Proportion of rearing spring Chinook feeding and agonistic strikes with increasing distance from the observed focal position in body lengths 2006-2014.


Figure 10. Mean ratio of agonistic strikes per feeding strike with increasing distance from the observed focal position 2006-2014.

Velocities were higher adjacent to spring Chinook focal positions in 72.1 percent of the observations in 2012, 86.0 percent of the observation in 2013, and 53 percent of the observations in 2014. A summary of microhabitat variable measured around Chinook focal positions is presented in table 4.

Table 4. Summary physical parameters measured at observed spring Chinook focal positions in 2012, 2013 and 2014.

|  | Temp ${ }^{\circ} \mathrm{C}$ | Spc length (mm) | Focal depth (m) | Total depth (m) Focal velocity (m/s) |  |
| :--- | :---: | :---: | :---: | :---: | :---: |
|  | 2012 (n=111) |  |  |  |  |
| Mean | 15.9 | 69.7 | 0.3 | 1.3 | 0.2 |
| SD | 1.0 | 7.4 | 1.1 | 5.7 | 0.1 |
|  |  |  | $2013(\mathrm{n}=43)$ |  |  |
| Mean | 16.3 | 79.5 | 2.3 | 0.9 | 0.2 |
| SD | 0.7 | 9.5 | 9.3 | 4.2 | 0.1 |
|  |  |  | $2014(\mathrm{n}=60)$ |  |  |
| Mean | 16.5 | 78.3 | 0.2 | 0.9 | 0.2 |
| SD | 0.9 | 10.2 | 0.2 | 0.3 | 0.1 |

## Spring Chinook residual/precocious male monitoring

The estimated number of natural origin age 0 , natural origin age 1 , and hatchery precocious males on the spawning grounds during the peak of spawning ranged from 5 to 718, 0 to 92, and 0 to 78 between 1999 and 2014 respectively (Table 5). Differences in the number of observed precocious males on or associated with active redds were detectable among age classes, and origin. Differences in the mean abundance of precociously mature males of different age and origin were detectable among years (ANOVA: $F_{2,45}=14.0, P<0.01$ ). Post-hoc analysis determined that natural production age 0 precocious males were greater in abundance than both natural and hatchery production age- 1 males (Tukey test: $P<0.01$ ). There were no detectable differences in abundance between age 1 natural and hatchery production precocious males (Tukey test: $P=0.96$ ). Among years, age 0 precocious males were found on a greater proportion of redds sampled than either age 1 or hatchery origin (ANOVA: $F_{2,45}=13.2, P<0.01$; Tukey test: $P<0.01$ ), and were greater in number per active redd (ANOVA: $F_{2,45}=19.4$, $P<0.01$; Tukey test: $P<0.01$ ), (Table 6.).

Table 5. Number of observed and estimated totals of natural (age 0 and age 1) and hatchery origin precocious males by age class at the peak of spawning activity in the upper Yakima River. Estimated totals are extrapolations over redds and/or portions of reaches not sampled.

| Survey year | Active redds | (\%) <br> Redds surveyed | (\%) Spawning area sampled | Observed |  |  | Estimated total |  |  |
| :---: | :---: | :---: | :---: | :---: | :---: | :---: | :---: | :---: | :---: |
|  |  |  |  | Age 0 | Age 1 | Hatchery | Age 0 | Age 1 | Hatchery |
| 1999 | 36 | 100 | 87 | 4 | 11 | 17 | 5 | 16 | 19 |
| 2000 | 316 | 66 | 87 | 103 | 42 | 8 | 128 | 42 | 11 |
| 2001 | 276 | 62 | 87 | 336 | 11 | 26 | 555 | 21 | 53 |
| 2002 | 304 | 81 | 87 | 138 | 15 | 8 | 228 | 25 | 14 |
| 2003 | 230 | 78 | 100 | 204 | 25 | 19 | 267 | 35 | 24 |
| 2004 | 1662 | 27 | 100 | 195 | 16 | 21 | 718 | 65 | 78 |
| 2005 | 655 | 99 | 100 | 357 | 17 | 0 | 360 | 17 | 0 |
| 2006 | 198 | 90 | 100 | 148 | 2 | 0 | 177 | 3 | 0 |
| 2007 | 92 | 100 | 100 | 55 | 0 | 0 | 55 | 0 | 0 |
| 2008 | 173 | 82 | 100 | 69 | 55 | 42 | 85 | 67 | 52 |
| 2009 | 105 | 99 | 100 | 87 | 15 | 34 | 88 | 15 | 34 |
| 2010 | 499 | 48 | 100 | 133 | 42 | 12 | 280 | 92 | 21 |
| 2011 | 418 | 73 | 100 | 124 | 40 | 0 | 171 | 55 | 0 |
| 2012 | 243 | 63 | 100 | 44 | 17 | 3 | 70 | 27 | 5 |
| 2013 | 166 | 66 | 100 | 76 | 10 | 3 | 115 | 15 | 5 |
| 2014 | 279 | 191 | 100 | 41 | 1 | 2 | 54 | 1 | 3 |

Table 6. Means of the presence and abundance of natural (age 0 and age 1) and hatchery origin precocious males per active redd at the peak of spawning activity in the upper Yakima River.

| Survey year | Active redds | Presence/Active redd |  |  | Abundance/Active redd |  |  |
| :---: | :---: | :---: | :---: | :---: | :---: | :---: | :---: |
|  |  | Age 0 | Age 1 | Hatchery | Age 0 | Age 1 | Hatchery |
| 1999 | 36 | 0.11 | 0.14 | 0.19 | 0.14 | 0.44 | 0.53 |
| 2000 | 316 | 0.18 | 0.10 | 0.02 | 0.41 | 0.13 | 0.03 |
| 2001 | 276 | 0.31 | 0.03 | 0.04 | 2.01 | 0.08 | 0.19 |
| 2002 | 304 | 0.23 | 0.03 | 0.03 | 0.75 | 0.08 | 0.05 |
| 2003 | 230 | 0.31 | 0.06 | 0.06 | 1.16 | 0.15 | 0.10 |
| 2004 | 1662 | 0.05 | 0.01 | 0.01 | 0.43 | 0.04 | 0.05 |
| 2005 | 655 | 0.24 | 0.02 | 0 | 0.55 | 0.03 | 0 |
| 2006 | 198 | 0.75 | 0.04 | 0 | 0.89 | 0.02 | 0 |
| 2007 | 92 | 0.18 | 0 | 0 | 0.60 | 0 | 0 |
| 2008 | 173 | 0.08 | 0.21 | 0.16 | 0.49 | 0.39 | 0.30 |
| 2009 | 105 | 0.24 | 0.09 | 0.13 | 0.84 | 0.14 | 0.33 |
| 2010 | 499 | 0.15 | 0.05 | 0.03 | 0.56 | 0.18 | 0.05 |
| 2011 | 418 | 0.24 | 0.07 | 0 | 0.41 | 0.13 | 0 |
| 2012 | 243 | 0.13 | 0.08 | 0.02 | 0.29 | 0.11 | 0.02 |
| 2013 | 166 | 0.20 | 0.07 | 0.04 | 0.69 | 0.09 | 0.03 |
| 2014 | 279 | 0.09 | 0.01 | 0.01 | 0.21 | 0.01 | 0.01 |

Hatchery precocious males were distributed differently than natural origin age 0 (G-test; $P=0.02$ ), and nearly when compared to natural origin age 0 and age 1 combined on the spawning grounds (G-test; $P=0.05$ ). A significant difference was not detected between natural origin age 0 and natural origin age 1 fish (G-test; $P=0.69$ ), or between natural origin age 1 and hatchery precocious males (G-test; $P=0.25$; Figure 11). An average of 28 percent of all hatchery precocious males observed on the spawning grounds were in the lowest spawning reach examined, whereas only 7 percent of natural origin age 0 , and 14 percent of natural origin precocious males were observed in this reach (Figure 11.).


Figure 11. Mean proportion $(p)$ of natural and hatchery origin precocious males by reach within the upper Yakima River at the peak of spawning activity 1999-2014. Error bars represent 95 percent confidence intervals with negative boundaries of zero.

Estimated total abundance of hatchery origin spring Chinook salmon away from redds at the time of spawning in 2013 ranged between 0 and $30 \mathrm{fish} / \mathrm{km}$ among sampling reaches (Table 7). The lower and upper Yakima Canyon averaged 59 percent of the estimated number of precocious males away from redds between 1999 and 2013, and the same percentage in 2014 (Figure 12). The annual abundance of hatchery precocious males away from redds was not significantly correlated with the number observed on redds $(P=0.07)$.

Table 7. Estimated abundance of hatchery origin spring Chinook salmon (HSPC) away from redds in the main stem Yakima River in the fall of 2014. The maximum number of fish netted ( n ) in one of two electrofishing surveys completed in consecutive weeks is presented (LCYN is the Lower Canyon, UCYN is the Upper Canyon, EBURG is Ellensburg, THORP is Thorp, and CELUM is Cle Elum). Capture probability was generated using rainbow trout of approximately the same size range as hatchery spring Chinook salmon.

| Section | n | Capture prob. | Section est. | Section km HSPC/km Reach km | Total est. |  |  |
| :--- | :---: | :---: | :---: | :---: | :---: | :---: | :---: |
| LCYN | 13 | 0.11 | 123 | 4.8 | 26 | 19.2 | 490.1 |
| UCYN | 29 | 0.18 | 158 | 5.2 | 30 | 13.4 | 407.0 |
| EBURG | 10 | 0.09 | 107 | 4.2 | 25 | 21.2 | 540.4 |
| THORP | 11 | 0.09 | 128 | 5.7 | 22 | 24.1 | 540.8 |
| CELUM | 0 | 0.09 | 0 | 7.4 | 0 | 16.2 | 0 |
| Total | 63 | $\mathrm{n} / \mathrm{a}$ | $\mathrm{n} / \mathrm{a}$ | $\mathrm{n} / \mathrm{a}$ | 104 | 94.1 | 1978 |



Figure 12. Proportional abundance $(p)$ of hatchery spring Chinook sampled away from redds in the fall of 2014, and the mean proportional abundance between 1999 and 2013.

## Predation Mortality

The mean lengths of the predators were not different between net pens (ANOVA, $P>0.05$ ). The rainbow trout ranged from 162 mm FL to 252 mm FL and the torrent sculpin ranged from 96 mm TL to 143 mm TL (Table 8). No significant differences were found between the mean lengths of the three origins of fry within each net pen at introduction (ANOVA, $P>0.05$ ). Mean lengths never varied more than 0.02 mm (Table 9). The weights of the fry at introduction did not statistically differ. The condition factors were not statistically different between groups in 2014 (ANCOVA, $P>0.05$ ).

Table 8. Dates, predator replicates, and mean lengths (ranges) of the predators for predation challenges (RBT = rainbow trout; TSC = torrent sculpin).

| Date Fry <br> Stocked | Date Fry <br> Removed | Week <br> $\#$ | Predator <br> Set | RBT Length <br> $(\mathrm{mm} \mathrm{FL})$ | TSC Length <br> $(\mathrm{mm} \mathrm{TL})$ |
| :---: | :---: | :---: | :---: | :---: | :---: |
| $3 / 25 / 14$ | $3 / 27 / 14$ | 1 | 1 | $190.1(170-245)$ | $115.7(104-127)$ |
| $3 / 30 / 14$ | $4 / 1 / 14$ | 2 | 2 | $185.6(164-229)$ | $112.8(103-133)$ |
| $4 / 1 / 14$ | $4 / 3 / 14$ | 3 | 3 | $170.5(163-179)$ | $110.8(104-119)$ |
| $4 / 6 / 14$ | $4 / 10 / 14$ | 4 | 4 | $197.5(162-252)$ | $109.6(96-128)$ |
| $4 / 13 / 14$ | $4 / 17 / 14$ | 5 | 5 | $186.3(163-239)$ | $115.3(98-141)$ |
| $4 / 20 / 14$ | $4 / 23 / 14$ | 6 | 6 | $186.2(163-220)$ | $110.6(101-129)$ |
| $4 / 27 / 14$ | $5 / 2 / 14$ | 7 | 7 | $177.7(163-220)$ | $113.2(103-143)$ |
| $5 / 5 / 14$ | $5 / 9 / 14$ | 8 | 8 | $187.8(164-215)$ | $114.3(105-132)$ |

Table 9. Mean fork lengths (standard deviation) of the hatchery (H), supplementation (S), and Naches ( N ) fry upon stocking in each net pen during the predation challenges.

| Origin | Week 1 | Week 2 | Week 3 | Week 4 | Week 5 | Week 6 | Week 7 | Week 8 |
| :---: | :--- | :--- | :--- | :--- | :--- | :--- | :--- | :--- |
| H | 36.35 | 37.47 | 37.89 | 39.46 | 41.04 | 41.83 | 43.68 | 44.03 |
|  | $(0.92$ | $(1.17)$ | $(1.14)$ | $(1.34)$ | $(1.30)$ | $(1.37)$ | $(1.54)$ | $(1.56)$ |
|  | 36.36 | 37.47 | 37.89 | 39.46 | 41.04 | 41.83 | 43.68 | 44.04 |
| S | $(0.92)$ | $(1.17)$ | $(1.14)$ | $(1.34)$ | $(1.28)$ | $(1.37)$ | $(1.54)$ | $(1.56)$ |
|  | 36.34 | 37.46 | 37.88 | 39.45 | 41.04 | 41.83 | 43.68 | 44.02 |
| N | $(0.90)$ | $(1.16)$ | $(1.11)$ | $(1.32)$ | $(1.28)$ | $(1.37)$ | $(1.54)$ | $(1.54)$ |
|  |  |  |  |  |  |  |  |  |

## Predation survival by origin

Overall mean survival between all origins was somewhat similar, with Naches fry survival being slightly higher than both hatchery and supplementation (Table 10). No statistical difference in survival was found between hatchery, supplemental, or Naches fry ( $\mathrm{P}>0.05$, Table 11).

During all years of this study, survival between years has varied considerably (Figure 13). This is likely due to varying lengths of time that the fry were exposed to predation during the trials. Within year differences in survival between groups has been relatively small and in most cases the supplementation line has had a slight survival advantage over the hatchery control line. Unfortunately, the Naches group has not been available for all years but has shown greater variability in relative survival from year to year (Figure 13).

Table 10. Percent of hatchery (H), supplementation (S), and Naches (N) fry surviving predator net pen trials at the end of each week and the overall mean survival and standard deviation.

| Week | Origin | Pen 1 | Pen 2 | Pen 3 | Pen 4 | Pen 5 | Pen 6 | Mean (SD) |
| :---: | :---: | :---: | :---: | :---: | :---: | :---: | :---: | :---: |
| 1 | H | 70.0 | 64.0 | 56.0 | 80.0 | 70.0 | 62.0 | 69.0(10.9) |
|  | S | 82.0 | 54.0 | 66.0 | 72.0 | 60.0 | 68.0 | 71.9(11.1) |
|  | N | 74.0 | 64.0 | 58.0 | 84.0 | 58.0 | 62.0 | 68.8(10.7) |
| 2 | H | 68.0 | 60.0 | 70.0 | 60.0 | 68.0 | 80.0 |  |
|  | S | 58.0 | 66.0 | 64.0 | 52.0 | 42.0 | 76.0 |  |
|  | N | 72.0 | 66.0 | 82.0 | 66.0 | 54.0 | 74.0 |  |
| 3 | H | 68.0 | 70.0 | 64.0 |  |  |  |  |
|  | S | 66.0 | 76.0 | 60.0 |  |  |  |  |
|  | N | 64.0 | 92.0 | 54.0 |  |  |  |  |
| 4 | H | 60.0 | 94.0 | 54.0 | 56.0 | 60.0 | 48.0 |  |
|  | S | 84.0 | 80.0 | 60.0 | 66.0 | 54.0 | 50.0 |  |
|  | N | 80.0 | 90.0 | 64.0 | 62.0 | 74.0 | 56.0 |  |
| 5 | H | 54.0 | 82.0 | 84.0 | 72.0 | 60.0 | 78.0 |  |
|  | S | 58.0 | 86.0 | 76.0 | 70.0 | 74.0 | 82.0 |  |
|  | N | 62.0 | 86.0 | 88.0 | 58.0 | 72.0 | 84.0 |  |
| 6 | H | 56.0 | 72.0 | 68.0 | 56.0 | 60.0 | 70.0 |  |
|  | S | 60.0 | 62.0 | 78.0 | 58.0 | 84.0 | 84.0 |  |
|  | N | 70.0 | 68.0 | 60.0 | 66.0 | 86.0 | 70.0 |  |
| 7 | H | 66.0 | 84.0 | 58.0 | 80.0 | 68.0 | 84.0 |  |
|  | S | 72.0 | 72.0 | 56.0 | 74.0 | 70.0 | 70.0 |  |
|  | N | 66.0 | 86.0 | 82.0 | 74.0 | 72.0 | 68.0 |  |
| 8 | H | 84.0 | 76.0 | 90.0 | 72.0 | 82.0 |  |  |
|  | S | 82.0 | 76.0 | 80.0 | 72.0 | 74.0 |  |  |
|  | N | 80.0 | 64.0 | 96.0 | 74.0 | 82.0 |  |  |

Table 11. Results from Wilcoxon matched pairs tests for survival between the three origins of fry, their relative survival ranking, and absolute and relative differences in the mean survivals.

|  |  |  |  | Relative <br> Survival <br> Ranking | Mean Abs. <br> Difference | Mean Relative Difference |
| :---: | :---: | :---: | :---: | :---: | :---: | :---: |
| Pairing | Z | N | P |  |  |  |
| N vs S | 1.92 | 44 | 0.06 | S=N | $3.14 \%$ | $4.56 \%$ |
| N vs H | 1.60 | 44 | 0.11 | N=H | $2.86 \%$ | $4.15 \%$ |
| S vs H | 0.49 | 44 | 0.63 | S=H | $0.27 \%$ | $0.40 \%$ |



Figure 13. Percent survival by origin for all previous years of this study. Error bars are 95\% confidence intervals.

## Competitive Dominance

Unequal numbers of replicates occurred among pair-wise comparisons because 1) some experiments did not meet the minimum criteria or 2 ) fish died. The replicates that did not meet experimental criteria or cases where fish died were a small percentage of the replicates that were conducted (Table 12).

Dominance was assessed in 164 replicates of supplementation vs. Naches fish during 2014. Naches fry were $1 \%$ more dominant but the difference was not statistically significant (Table 13). The frequencies of the different types of interactions used by supplementation and Naches fish during pair-wise comparisons were not significantly different (Table 14). Differences in aggression between Naches and supplementation fish were not detected. Differences between dominant x dominant and subordinate x subordinate fish trials were also not detected (Table 15). Naches fish grew slightly more and gained slightly less weight than supplementation fish although the differences were not significant (Table 16). Dominant fish regardless of origin grew more length and gained more weight than subordinate fish (Table 19).

Dominance was assessed in 155 replicates of supplementation vs. hatchery fish in 2014. Hatchery fish were 3\% more dominant than supplementation fish in 2014 but the difference was not statistically significant (Table 13). The frequencies of different types of interactions used by supplementation and hatchery fish during pair-wise comparisons were not significantly different (Table 14). Difference in aggression between hatchery fish and supplementation were not detected. Differences between dominant $x$ dominant and subordinate x subordinate fish trials were not detected (Table 15). Hatchery fish grew slightly more and gained slightly less weight than supplementation fish (Table 16). Dominant fish regardless of origin grew more length or lost less weight than subordinate fish (Table 16).

Dominance was assessed in 157 replicates of Naches vs. hatchery fish during 2014. Naches fish were $15 \%$ more dominant than Naches fish in 2104 but the difference was not statistically significant (Table 13). The frequencies of different types of interactions used by Naches and hatchery fish during pair-wise comparisons were not statistically significantly (Table 14). Interaction rates of Naches fish were significantly higher than hatchery fish in 2014. Differences between interaction rates of dominant x dominant fish were not detected. Naches subordinate fish had a significantly higher interaction rate than hatchery subordinate fish in 2014 (Table 15). Naches fish grew slightly more and gained slightly more weight than Naches fish although the differences were not significant (Table 16). Dominant fish regardless of origin grew more length or lost less weight than subordinate fish (Table 19).

Table 12. Factors responsible for eliminating contest replicates from analysis.

| Origin | Died or <br> Missing | $<10$ Pellets | No <br> Interactions | Abnormal <br> behavior | Total |
| :---: | :---: | :---: | :---: | :---: | :---: |
| Supp <br> Naches <br> Both | 6 | 2 | 11 | 1 |  |
| Supp <br> Hatchery <br> Both | 7 | 8 | 8 | 0 | 20 |
| Naches <br> Hatchery <br> Both | 1 |  |  |  | 23 |

Table 13. Comparisons of mean ( $\pm 1$ SD) of the \% food acquisition, \% habitat occupation, \% agonism dominance (dom. interactions), \% total dominance, sum of the scores used to assess dominance, and P values from Wilcoxon matched pairs test in contest competition experiments between supplementation (Supp.), hatchery, and Naches Chinook salmon.

| Origin | n | \% Food | \% Habitat | \% Dom. <br> Interactions | \% Total <br> Dom. | Sum Total <br> Dom. $\%$ | P |
| :--- | :---: | :---: | :---: | :---: | :---: | :---: | :---: |
| Supp | 164 | $48(40)$ | $45(44)$ | $48(49)$ | 49 | $141(128)$ |  |
| Naches | 164 | $52(40)$ | $55(44)$ | $52(49)$ | 51 | $159(128)$ | 0.181 |
|  |  |  |  |  |  |  |  |
| Supp | 155 | $49(38)$ | $48(45)$ | $47(49)$ | 48 | $143(128)$ |  |
| Hatchery | 155 | $51(38)$ | $52(45)$ | $53(49)$ | 52 | $157(128)$ | 0.342 |
|  |  |  |  |  |  |  |  |
| Naches | 157 | $56(40)$ | $56(45)$ | $58(48)$ | 57 | $170(130)$ |  |
| Hatchery | 157 | $44(40)$ | $42(45)$ | $42(48)$ | 43 | $130(130)$ | 0.062 |

Table 14. Comparisons of frequencies of interaction types initiated (mean interaction/fish in each tank ( $\pm 1$ SD)) by supplementation (S), hatchery (H), and Naches $(\mathrm{N})$ fish and total interactions (total ints.) by origin in contest competition experiments.

|  |  |  |  |  |  | Total | P <br> Origin |
| :---: | :---: | :---: | :---: | :---: | :---: | :---: | :---: |
| Crowd | Threat | Chase | Butt | Nip | Ints. | G-test |  |
| S | $0.16(2.67)$ | $2.6(9.84)$ | $8.65(20.20)$ | $0.70(2.76)$ | $0.42(1.48)$ | 2055 |  |
| N | $0.09(1.36)$ | $3.55(11.53)$ | $11.03(23.44)$ | $0.52(1.38)$ | $0.45(1.40)$ | 2565 | 0.998 |
|  |  |  |  |  |  |  |  |
| S | $0.18(2.30)$ | $3.22(14.04)$ | $9.14(18.64)$ | $0.63(2.70)$ | $0.45(2.47)$ | 2106 |  |
| H | $0.11(2.70)$ | $4.55(15.83)$ | $11.45(21.10)$ | $0.81(2.45)$ | $0.59(2.83)$ | 2713 | 0.999 |
|  |  |  |  |  |  |  |  |
| N | $0.14(1.18)$ | $3.98(12.51)$ | $9.86(16.48)$ | $0.58(3.28)$ | $0.44(2.28)$ | 2355 |  |
| H | $0.12(1.27)$ | $2.96(12.99)$ | $7.65(15.66)$ | $0.52(2.83)$ | $0.41(4.09)$ | 1831 | 0.999 |

Table 15. Interaction rates (mean interaction/fish/minute) of agonistic interactions initiated by supplementation (Supp.), hatchery and Naches fish in contest experiments.

| Origin | $\mathrm{n}^{\text {a }}$ | Mean Interaction rate | 1 Standard Deviation | $\mathrm{P}^{\text {b }}$ |
| :---: | :---: | :---: | :---: | :---: |
| Naches | 164 | 0.76 | 1.12 |  |
| Supplementation | 164 | 0.60 | 0.97 | 0.164 |
| Naches Dominant | 94 | 1.29 | 1.22 |  |
| Supp. Dominant | 94 | 1.02 | 1.11 | 0.152 |
| Naches Subordinate | 94 | 0.30 | 0.87 |  |
| Supp. Subordinate | 94 | 0.12 | 0.38 | 0.053 |
| Supplementation | 155 | 0.65 | 0.97 |  |
| Hatchery | 155 | 0.83 | 1.17 | 0.127 |
| Supp. Dominant | 80 | 1.23 | 1.06 |  |
| Hatchery Dominant | 82 | 1.52 | 1.23 | 0.120 |
| Supp. Subordinate | 82 | 0.22 | 0.31 |  |
| Hatchery Subordinate | 80 | 0.06 | 0.75 | 0.045 |
| Naches | 157 | 0.71 | 0.90 |  |
| Hatchery | 157 | 0.56 | 0.90 | 0.006 |
| Naches Dominant | 97 | 1.12 | 0.93 |  |
| Hatchery Dominant | 77 | 1.13 | 1.02 | 0.790 |
| Naches Subordinate | 77 | 0.23 | 0.62 |  |
| Hatchery Subordinate | 97 | 0.09 | 0.42 | 0.012 |
| ${ }^{\text {a }}$ Number of replicates ${ }^{\mathrm{b}} \mathrm{P}$ values from Mann-W dominance, when they | domi | re for comparison and when they w | fish origins $r$ ubordinate. |  |

Table 16. Comparisons of supplementation (Supp.), hatchery, and Naches fish growth in contest competition experiments. Replicate numbers vary a bit because when equal growth occurred, they were not analyzed.

| Origin | $\mathrm{n}^{\mathrm{a}}$ | Average Growth mm Length | Average Growth <br> Mg Weight |
| :---: | :---: | :---: | :---: |
| Supp. | 164 | $0.86(1.25)$ | $89.89(266.61)$ |
| Naches | 164 | $1.04(1.19)$ | $52.90(245.84)$ |
| $\mathrm{P}_{\mathrm{b}}$ |  | 0.206 | 0.258 |
|  |  | $0.86(1.21)$ | $107.63(277.61)$ |
| Supp. | 155 | $0.92(1.30)$ | $104.39(250.17)$ |
| Hatchery | 155 | 0.727 | 0.925 |
| $\mathrm{P}_{\mathrm{b}}$ |  |  |  |
|  |  | $0.94(1.31)$ | $88.97(322.13)$ |
| Naches | 157 | $0.78(1.19)$ | $67.06(252.30)$ |
| Hatchery | 157 | 0.287 | 0.552 |
| $\mathrm{P}_{\mathrm{b}}$ |  |  |  |

${ }^{a}$ number of replicates
${ }^{\mathrm{b}} \mathrm{P}$ values for statistical tests
Note: Numbers in parentheses are 1 standard deviation

Over the years, relative percent dominance between the hatchery and supplementation groups has been variable with a slight indication that the hatchery control fish were increasing in dominance during the first generation but has since leveled out (Figure 14). While not available for comparison for all years of study, the relative dominance between hatchery control and Naches fish have tracked very closely to that of hatchery versus supplementation until the last two years (Figure 14). The relative dominance between the supplementation and Naches fish does not show any clear trend relative to the other two comparisons in the earlier years but has since tracked with the hatchery versus Naches.


Year

Figure 14. The relative percent dominance between the pairings of the three populations from 2005 through 2010 ( $\mathrm{H}=$ hatchery line; S = supplementation line; $\mathrm{N}=$ Naches wild line).

## Spring Chinook reproductive success/spawning channel

Genetic analysis revealed that all 96 fish released or found in the spawning channel had unique genotypes. There were a total of 24 hatchery control line (HC) adult males, 24 HC adult females, 24 supplementation hatchery line (SH) adult males, and 24 SH adult females. Four HC males and four HC females were released into three of the six sections and four SH males and four SH females were released into the other three sections (Table 1, Appendix D). A total of 14 loci were screened and all 14 were used in the analysis (Table 2, Appendix D). Number of alleles ranged from 4-30 (Ots-9 and Omm-1080 respectively) and observed heterozygosity ranged from $0.448-0.958$ (Ots-G474 and Ots201 b respectively). Individual exclusionary power was below $45.4 \%$ for five loci (Ogo2, Ogo-4, Ots-G474, Ots-3M, and Ots-9) and above $61.6 \%$ for the remaining loci when neither parent was known. Exclusionary power was below $42.1 \%$ for three loci (OtsG474, Ots-3M and Ots-9) and above $60.2 \%$ for the remaining loci when one parent was known. Cumulative exclusionary power was 1.000000 for analysis using all loci when one parent was known. Parentage assignments were made when genotype data was available for nine or more loci. All 96 parents were genotyped at 10 or more loci while 2,741 of the 2,784 offspring were successfully genotyped at nine or more loci (Table 3, Appendix D). Parentage analysis was conducted independently for each of the six sections using all 96 adults as possible parents. Each fry was assigned a dam-sire-fry combinations (trios) based on the most likely candidate male parents (sires) and female parents (dams). Those assignments yielded possible. Any fry-sire assignments with more than two mismatching loci were excluded from further consideration. Of the total

2,741 fry included in the analysis a total of 2,545, fry were assigned to a single male and female parent (2,545/2,741 = 92.8\%).

Spring Chinook Genetic stock separation-juveniles
A total of 1,200 unknown Chinook smolts were selected and analyzed from those collected at Chandler Trap. Smolt samples that had data for 10 or more loci were included for analysis. A total of 20 individuals were dropped from statistical analyses. The mixture composition estimates for the entire 2013 smolt outmigration indicated that the largest overall percentage of spring smolts was from the upper Yakima River followed by the Naches River and American River in the first four strata. During the migration from January - May, the proportion of the upper Yakima River stocks was between 66.7 and $76.2 \%$ while the American River and Naches River spring stocks was between 3.3 and 27.5\%. The proportion of the two fall stocks was between $0.0-22.3 \%$ for the first four time strata and $70.6 \%$ in the June - July time stratum (Table 3, Appendix E). A comparison of the morphological assessment to genetic assignment was conducted for all five time strata. A total of 39 smolts in January/February, 82 smolts in March, 724 smolts in April, 140 smolts in May, and 195 in the June/July time strata were scored, and therefore included in the analysis. Results for the time strata were as follows: January/February time stratum - all 39 smolts were assigned identically using morphological and genetic methods ( 39 spring); March stratum - 81 out of 82 smolts were assigned identically using morphological and genetic methods ( 81 spring) the one discrepancy was identified as a fall by the genetic analysis and spring with morphological identification; April time stratum - 724 smolts were assigned identically using morphological and genetic methods ( 724 spring); May time stratum - 136 out of 140 smolts were assigned identically using morphological and genetic methods (100 spring 36 fall), all four of the discrepancies were identified as a spring by the genetic analysis and fall with morphological identification; June/July time stratum - 145 out of 195 smolts were assigned identically using morphological and genetic methods (12 spring and 133 fall), 4 discrepancies were assigned as fall by the genetic analyses while morphological identification was spring, the remaining 46 discrepancies were identified as a spring by the genetic analysis and fall with morphological identification.

## b. Hatchery RM\&E

The performance of the YKFP spring Chinook supplementation program has been documented relative to the project quantitative objectives and has been presented annually in the YKFP M\&E project overview (Fritts 2012). Briefly, the project appears to be meeting or is making progress towards achieving the project's objectives (Appendix C).

## 5. Synthesis of Findings: Discussion/Conclusions

## a. Fish Population RM\&E

Non-target taxa of concern monitoring
We failed to reject the hypothesis that early-middle stages of salmon supplementation have impacted valued trout species in the upper Yakima Basin beyond predetermined containment objectives. There were no impacts of supplementation activities on bull and cutthroat trout that inhabited tributary streams because limited or no overlap with hatchery or naturally produced salmon occurred. However, the potential existed for much overlap between salmon and bull and cutthroat trout in the tributaries of the upper Yakima Basin. For example, hatchery steelhead that were released in 1994 very close to the release site in the North Fork of the Teanaway River, migrated upstream into areas containing bull trout and cutthroat trout (McMichael and Pearsons 2001). Hatchery spring Chinook also migrated upstream of the acclimation site in the North Fork of the Teanaway River, but not nearly as far as hatchery steelhead. This finding is consistent with our earlier work and extends the findings into later stages of supplementation (Pearsons and Temple 2007).

It is possible that some overlap occurred at times and places when/where we did not sample. However, substantial overlap was unlikely because we sampled at times and places that overlap was most likely. There are certainly areas outside the upper Yakima watershed where overlap occurs at the times that we sampled. Furthermore, overlap has been detected using the methods we used (e.g. snorkeling). Salmon and bull and cutthroat trout overlap during the summer in another large tributary in the Yakima Basin that parallels the upper Yakima River. In the Naches Basin, which merges with the upper Yakima River near the city of Yakima, substantial overlap exists between bull and cutthroat trout and naturally produced Chinook salmon (T. Pearsons, unpublished data). Hatchery coho salmon are released into that basin and undoubtedly overlap with bull and cutthroat trout. Other studies have also documented overlap between salmon and cutthroat and bull/Dolly Varden trout (Glova 1984; Bisson et al. 1988; Nakano and Kaeriyama 1995; Thurow et al. 1997).

There are a variety of possible reasons why overlap was not detected in tributaries of the upper Yakima River. First, all but one of the acclimation sites for salmon were located in the main stem and the acclimation site in the tributary was located downstream of bull and cutthroat trout. Risks to bull and cutthroat trout were one of many factors that contributed to acclimation site placement. Second, the distribution of juvenile salmon has not increased substantially even though the abundance of adult salmon has increased. We had expected that the distribution of juvenile salmon would have increased with increasing abundance of spawners. Third, high abundance of rainbow trout in lower elevation portions of tributaries may competitively exclude cutthroat and bull trout to higher elevations that salmon do not occupy. Relaxation of competition could result in broader distributions of bull and cutthroat trout and the possibility of greater overlap with salmon. Fourth, salmon, bull trout, and cutthroat trout have different habitat preferences. Salmon typically occupy streams of lower gradient, lower elevation, and warmer water temperatures than cutthroat and bull trout (Glova 1987; Dunham and Rieman 1999).

Glova (1987) concluded that impacts to cutthroat trout could be reduced by stocking coho in areas with gradients greater than $1 \%$ and ample fast water habitats. Faster water velocities allow for more resource partitioning and competitive dominance by trout. Most of the tributaries in the upper Yakima Basin met these criteria. We did observe overlap between salmon and cutthroat trout in the main stem, where water temperatures were more suitable for both of these species.

Contrary to our previous findings (Pearsons and Temple 2007), we did detect a significant difference in the abundance of rainbow trout in treatment areas in the North Fork Teanaway River and main stem Teanaway River relative to our control sites (Pearsons and Temple 2010). With each additional year of sampling we will have increased power to detect smaller differences (Ham and Pearsons 2000). However, it is important to note that our "Before-After" detection plan would not have triggered the "Causation" analysis that was used to detect the decline and the decline was isolated to a small area and was small relative to the total population size. Furthermore, we do not yet have evidence to support the decline was due to mortality of fish in the treatment area. Other possibilities may include displacement, and perhaps angler harvest, both of which we are currently evaluating.

Although we observed decreases in the size of rainbow trout during the postsupplementation period, the decline is unlikely to have been caused by supplementation. If supplementation had changed the size structure or growth of the steelhead size index, we would expect to detect this change in areas with high densities of salmon. We did not detect a reduction in the size of rainbow trout in the high-density areas of the target taxa below the Clark Flats acclimation site or below the release site in the North Fork Teanaway River. These areas are likely to have the greatest potential of detecting an impact. One potential explanation for the observed decrease in main stem rainbow trout size is that intraspecific density dependent mechanisms have altered the size of main stem Yakima River rainbow trout. The abundance of rainbow trout increased by approximately $30 \%$ ( $30 \%$ increase of age 1 fish, and $29 \%$ increase of fish greater than 249 mm ) after stocking began. This information and results from small-scale enclosure experiments (McMichael et al. 1997) leads us to believe that the decline in rainbow trout lengths is most likely the result of intraspecific competition.

With the exception of the BACIP results from the Teanaway basin, the lack of detectable impacts to rainbow trout were consistent with results that were derived from smaller scale enclosure experiments between naturally produced spring Chinook salmon and rainbow trout in high elevation tributaries (McMichael and Pearsons 1998). In these experiments, growth and abundance of rainbow trout were not impacted when the density of salmonids was doubled by the addition of naturally produced spring Chinook salmon parr. However, growth of rainbow trout was suppressed when the density was doubled with rainbow trout (McMichael et al. 1997), which supports the previously mentioned idea of intraspecific impacts to rainbow trout growth in the main stem. The current results extend the findings of McMichael and Pearsons (1998) to smolts, residuals, coho salmon, and to lower elevation waters such as the main stem. Our ability to detect impacts with the BACI design and the longer experimental period in this study (higher statistical power) may explain the differences among the studies. Opportunities for
cumulative impacts to manifest and larger sample sizes may be necessary to detect impacts where high natural variation occurs.

It is possible that our abundance estimates in the main stem and tributaries and the size estimates in the tributaries were influenced by the size breaks that we used in our analysis. The lower size breaks were necessary (e.g., 80 mm in tributaries and 100 mm in the main stem), because we capture very few of these fish due to our low electrofishing efficiencies on small fish and hence cannot calculate valid estimates on these fish. This could result in varying proportions of age 0 and 1 fish in our estimate if the length at age varied across years or sites. However, we do not believe that length truncations significantly affected our conclusions. For example, if fish length was negatively impacted then the distribution of fish size would have become smaller, and more age 1 fish could have been pushed below 100 mm . Regardless of how many fish may have been shifted below 100 mm , if the impact occurred to the whole age class then we should have detected a decrease in size for fish above 100 mm (e.g., the whole length frequency curve would be shifted to smaller sizes). Similarly, if many fish were impacted so that they were less than 100 mm then the abundance of age 1 fish would have been negatively biased. In other words, we would expect to detect less fish than we did prior to supplementation. If we had concluded that impacts had occurred, then our length truncations would be a more serious issue.

We did not detect impacts to non-trout NTT that could be attributed to supplementation. In the tributaries, this was because none of the non-trout NTT overlapped with salmon at high enough levels to exceed the CO. All non-trout NTT in the main-stem overlapped completely, but none exceeded the containment objectives.

With the exception of minimum daily stream discharge in the main stem Yakima River, we did not detect changes in the environmental variables that were measured. We hypothesize the increased minimum daily stream discharge observed would benefit NTT. However, the increased minimum daily discharge was not significantly correlated with our NTT monitoring variables suggesting that it did not confound our results. Average and maximum stream discharge and temperature were heavily regulated by upstream irrigation reservoirs providing a relatively stable environment to conduct risk containment monitoring. Although discharge in tributaries is unregulated, summer base flows have not differed drastically during the time of sampling from year to year. The relatively stable environmental conditions observed in both tributary and main stem areas supports the use of time as a control in our evaluation.

## Post-emergent growth

In a multiple year analysis (2009-2014), observed growth rate differed significantly among years (Homogeny of slopes model: $F_{5,192}=4.90, P<0.01$; Figure 5). Mean length, accounting for sampling date, was also detectably different among years (Separate slopes model: $F_{4,192}=5.22, P<0.01$ ). Post-hoc analysis revealed significantly greater mean length in 2010 and in 2014 in comparison to 2009; which had the smallest observed mean length within the six-year dataset.

The development of a growth model has allowed the detection of annual differences in size and growth rate among years. This information provides insight into upper Yakima spring Chinook population dynamics in the fry-to-parr life stage, and contributes to our understanding of environmental factors and/or behavioral responses which may negatively affect growth or survival in years of high spawner density.

The Yakima Basin experienced flow conditions throughout the 2009-2010 incubation period that were lower and far less variable than average for the system (Johnson et al. 2012). These conditions may have resulted in a relaxation of environmental influences on survival, resulting in uncharacteristically high survival and, in effect, an increase in the system capacity for spring Chinook subyearlings. Our data indicate that greater growth and size were present in the early rearing period in 2010, suggesting such a relaxation in limiting factors was present within or before the spring sampling period.

High observed productivity in the fall of both 2010 and also 2011 may give indications of the time period in which density dependent constraints exist under normative conditions in the upper Yakima. Preliminary results from genetic stock separation analysis of 2011 spring Chinook smolts originating from both the Yakima and Naches River basins (2010 fry-parr; WDFW unpublished data) suggests that high productivity in 2010 was not unique to the Yakima River basin. It is possible that a relaxing of capacity constraints occurred as a result of larger scale environmental conditions affecting multiple basins. If this is true, and the larger scale trends temporary, we might expect to observe a slow decrease in productivity in years following high escapement until the system again returns to its previous capacity for spring Chinook production. If this occurs, data collected from years of unusually high system productivity may give additional insight into the specific factors, again present, affecting survival in the upper basin.

## Rearing abundance and habitat use

Our data suggest a greater abundance of summer rearing spring Chinook in 2010 than in any other survey year. This is consistent with our detection of greater size and growth in the spring, and also abundance and size in the fall of 2010. These findings, along with a nearly significant correlation between estimates of summer and fall abundance, suggest our methods were successful in tracking relative productivity through three distinct subyearling spring Chinook life-stages. Over time, these data should allow identification of the life-stage in which limitations to growth and survival are occurring; a critical first step in identifying the specific factor or factors negatively affecting the population in some years.

Yakima River spring Chinook redd-to-parr productivity observed in the fall of 2010 (WDFW unpublished data) was much higher than that predicted through the use of a Beverton-Holt recruitment curve developed using data from the previous sixteen years (Johnson et al 2009). Environmental conditions in the spring of 2010 appear to have been very conducive to early survival, perhaps due to an uncharacteristically low number of high-flow events during the incubation period (Johnson et al 2012). The absence of a
detectable response through subsequent life-stages when environmental conditions were not notably different (late spring, summer, and fall), suggests that capacity constraints may exist in earlier developmental periods in years where environmental conditions are more normative.

Documenting the existence of density dependent constraints post-emergence is confounded by the fact that this is often a period of high mortality, even when spawner densities are low. A system's capacity for incubating alevin is generally far greater than its capacity for juveniles, which generally results in low spawning densities, high survival to emergence, and post-emergent thinning of the population (Quinn 2005). However, during high return years, when competition exists for preferred spawning habitats, density dependent limitations to growth and survival may ultimately occur prior to first emergence. Such limitations may be attributable to a number of potentially limiting environmental factors such as increased sedimentation, scour, temperature, and/or decreased dissolved oxygen levels in less optimal spawning habitats. Estimates of lifestage specific growth and abundance during years with a high density dependent response will be necessary to identify limiting factors with any degree of certainty. We will continue to monitor summer parr abundance and to investigate the potential relationship between our summer and fall estimates.

Perhaps as important as the documentation of abundance in the summer rearing period is the identification of the existing habitats most heavily utilized by subyearling Chinook. This information may help in the identification of limiting factors, but will also further our understanding of reach specific productivity in years of low density; a critical metric that is often missing from restoration efforts, which often concentrate only on limitations or "bottlenecks" within the population (Mobrand et al. 1997). Although we encountered high variability in abundance among sampling units, we did find higher densities of rearing Chinook in pool and deep riffle type habitats. Therefore, the summer distribution of rearing subyearlings appears to be in-part due to the presence of certain habitat types. In addition to other, larger scale, environmental factors which may affect movement and subsequent survival (e.g. temperature, flow events) the use of habitat type as an explanatory variable should be beneficial in determining relative productivity among reaches of the upper Yakima River for summer rearing spring Chinook salmon.

Territory size continued to be strongly associated with spring Chinook length, which is consistent with the findings of others (Grant and Kramer 1990, Keeley and Grant 1995). These data suggest that territory may be a reasonable microhabitat metric to measure the degree of competition for space. Previous work in the Yakima Basin was unsuccessful in linking calculated territory based on local abundance to fall abundance (Pearsons et al. 2007). However, the spatial scale of those measures may have been either too large to detect changes in territorial behavior, or measured after any subsequent mortality or out-migration had occurred. Subyearling Chinook decreased the frequency of defense and foraging with increased distance from the holding position. This is consistent with our expectations that increased effort would be required to defend and utilize space away from the position of holding. The frequency of defense may be just as important as the size of the observed territory when evaluating limiting factors. For example, if food is a limiting factor, then we may observe highly defended areas of high
food availability and smaller territory size, and areas of low food availability where the individual is forced to defend a larger area. These two scenarios may be energetically equivalent for the individual. Ranges of focal depth, total depth, and focal velocity during our observations were within the ranges of previous years.

## Spring Chinook residual/precocious male monitoring

Despite the large numbers of precocious males that are apparently released from the CESRF annually (Larsen et al. 2004; Beckman and Larsen 2005; Larsen et al. 2006), only a small fraction of these fish appear to reach the spawning grounds. Hatchery precocious males may experience high mortality, migrate out of the study area after release, and/or fail to migrate back to the spawning grounds. Although the occurrence of some of these factors were observed in this or other studies (Larsen et al. 2004; Beckman and Larsen 2005), we do not know the relative contribution of each of these factors towards the low abundance of precocious males on the spawning grounds.

Mortality of hatchery precocious males may be due to high angler exploitation, starvation, or predation. There is considerable angling pressure focused on trout in the Yakima River, and anglers have at times commented on the number of precocious Chinook males caught, particularly in 2001. However, it is illegal to keep Chinook salmon in the upper Yakima River. Furthermore, studies have shown that hatchery origin fish released into the natural environment have lower survival than natural origin fish, presumably because of their inability to find food or avoid predators (White et al. 1995; Weber and Fausch 2003).

It has been documented that some hatchery precocious males move downstream out of the spawning areas and have been detected as far downstream as Bonneville Dam on the Columbia River (Larsen et al. 2004; Beckman and Larsen 2005). In northern Oregon, precocious males were documented to have migrated at least 800 km and past three dams to reach salt water and return to the Umatilla River (Zimmerman et al. 2003). Hatchery precocious males were collected migrating both downstream in the spring and upstream during the summer (Larsen et al. 2004; Beckman and Larsen 2005). The downstream migrations occurred during the smolt out-migration period and the upstream migrations occurred at the time of adult spawning immigration. If precocious males migrate downstream and then environmental conditions turn poor before they are able to migrate back upstream, then they are likely to die. The lower Yakima River becomes lethal for salmonids during many of the hot summer months when precocious males might attempt to ascend the river. If the factors contributing to hatchery fish mortality in the river are reduced or the conditions in the river are favorable for migration back to the spawning grounds (e.g., favorable flows and low angling pressure), then presumably the number of hatchery precocious males on the spawning grounds could increase dramatically. However, the range of conditions that we evaluated in this study, which included both high and low flow years, provide a reasonable range of what can be expected in the future.

Most of the hatchery precocious males that we encountered were located downstream of spawning areas. The lower and upper Yakima Canyon typically contain less than $1 \%$ of the upper Yakima Basin redds (Yakama Nation, unpublished data) and yet averaged $59 \%$ of the estimated number of hatchery precocious males during the spawning season. Many of the hatchery precocious males on the spawning grounds were observed in a reach that had relatively little spawning activity, whereas the natural origin precocious males were mainly in the areas with high spawning activity. The spawning area where many of the hatchery precocious males were observed was at the lower end of
the spawning distribution. It also happens to be located closest to the Yakima Canyon where the highest abundance of precocious males that were not on the spawning grounds was observed. In the Wenatchee River, very few hatchery precocious males were observed on the spawning grounds, but a considerable number were captured migrating upstream at a location downstream of the spawning areas (Murdoch et al. 2007). These fish may have also distributed themselves below the main spawning areas as we observed in the Yakima Watershed. This behavior is in contrast to natural origin precocious males that are rarely observed moving upstream past dams in the Yakima or Wenatchee watersheds, suggesting that natural origin precocious males have adopted a strategy of remaining on or near the spawning grounds and thus conserving energy and promoting growth and testes development. Some hypotheses as to why sexually mature hatchery precocious males, most of which are exuding milt at the time of sampling, are located in areas away from where most of the spawning activity occurs include: lack of energetic capacity to swim back upstream to the spawning grounds; inappropriate downstream migration behavior for their life-history strategy; late migration timing; and inability to locate areas with spawning females after they had migrated downstream of spawning areas. Younger salmon, such as precocious males and jacks, typically migrate back to the spawning grounds later than older salmon (Knudsen et al. 2006; Murdoch et al. 2007) and may migrate during unfavorable environmental conditions.

Cle Elum Hatchery origin fish are only released at age 1, which eliminates the possibility that age 0 hatchery precocious males will have the potential to spawn. In the absence of hatchery releases, age 0 precocious males are generally more abundant in the spawning areas than age 1 precocious males, so the hatchery is skewing the precocious male composition to an older age and larger size. This is in stark contrast to anadromous hatchery fish which typically mature earlier than wild fish and often at a smaller size-atage (Knudsen et al. 2006). It is interesting to note that few incidences of precocious male maturation at age 0 have been observed in the Yakima hatchery (Larsen et al. 2004). In addition, attempts to experimentally produce age 0 precocious males by high feeding rates in the hatchery did not produce any precocious males in 2002 (Farrell 2003). These fish emerged at the average emergence time of the population. It is possible that only the fish that emerge very early and experience good growth have the potential to precociously mature at age 0 (Larsen et al. 2007). However, because precocious males were not used in the broodstock, we cannot eliminate the possibility that genetics also influenced the absence of precocialism (e.g., Heath et al. 1994; Unwin et al. 1999).

Hatchery age 1 fish may be competitively superior to wild precocious males because hatchery precocious males are larger. Larger salmonids typically dominate smaller ones in behavioral contests (McMichael et al. 1999). We have observed a number of instances where hatchery precocious males displaced wild precocious males from redds or from preferred locations on redds. Behavioral dominance is important because dominant fish are more likely to be close to spawning females and hence more able to fertilize eggs (Garant et al. 2003). Dominant fish are better able to choose which locations pose the best chance for spawning success. Our behavioral observations suggest that per capita fertilization rates of hatchery precocious males should be higher than that of wild precocious males. However, sneaking strategies of smaller individuals may also be successful.

We have identified some issues that could potentially contribute to the underestimation of precocious male numbers during our peak snorkel counts. We may have underestimated the number of active redds by spooking adults or by floating at times when adults are temporarily away from their redds. However, we rarely observed precocious males on redds without adults being present and this finding was also supported by work in the Salmon River drainage (Gebhards 1960). Gebhards (1960) concluded that precocious males were generally only found in areas where there was spawning activity and were usually found in the bowl of the redd, and "the yearling males remained constantly within the redd."

Other reasons include the possibility that precocious males may have been hiding away from the redds, were scared off the redds, were moving between redds, or were present in greater numbers before or after our peak count. Additional snorkeling efforts along the banks in 1998 and 2007 did not find hatchery precocious males in hiding areas such as undercut bank in the vicinity of spawning areas, and multiple reach surveys conducted in 2007 and 2008 did not suggest greater numbers of precocious males on the spawning grounds the week before or after our peak of spawning surveys. We have also observed that repeated counts of precocious males at three different times of the day in the same reach were similar. This suggests that either our counts were accurate or that our bias was consistent. However, our estimates of fish away from redds, that were generated from electrofishing were higher in some years than those generated from snorkeling in the Thorp reach, suggesting that snorkeling may underestimate abundance.

In short, if we underestimated the number of precocious males on the spawning grounds then our numbers should be treated as indices.

Our study suggests that hatchery precocious males are unlikely to contribute a high proportion of genes in the Yakima Watershed when the number of anadromous adult returns is high, but contributions could be high when anadromous adult numbers are low. The highest abundance of hatchery precocious males that we estimated on the spawning grounds during any year was 78 . This is a small proportion of the spawners when anadromous spawners number in the thousands, but relatively large when the abundance of spawners is in the hundreds. This range of anadromous fish abundance has been observed in the upper Yakima Watershed. In a separate DNA pedigree study conducted in an artificial spawning channel (Schroder et al. 2006), hatchery and natural origin precocious males of the upper Yakima spring Chinook salmon stock have been documented to sire offspring. In addition, precocious maturation appears to be highly heritable in Yakima spring Chinook salmon (Pearsons et al. 2007d). In short, it appears that the genetic contribution of hatchery precocious males on the spawning grounds is related to anadromous fish abundance and those factors that influence the abundance of precocious males on the spawning grounds. Variation in the precocious male contribution suggests that domestication risks may vary among years.

## Predation Mortality

The wild Yakima River spring Chinook were found to be slightly over two percent more successful at surviving the predation trials than the first generation of the hatchery control population during 2003 and 2004 (Fritts et al. 2007). Beginning in 2005, the supplementation population has generally exhibited an equal to a slight survival advantage over the hatchery control population (Fritts and Stockton 2010).

We speculate that it is possible to detect differences in survival between the hatchery, supplementation, and wild Naches origin fry in some years and not to detect differences in other years due to changes in selection pressures between years. For example, assume that the numbers of adults used for broodstock at the CESRF were to remain relatively constant and the numbers of adults spawning naturally fluctuated by several orders of magnitude. It is theoretically possible for the supplementation and Naches populations to express varying degrees of predation vulnerability from year to year due to density dependent selection pressures in the river environment. Offspring from an abundant run of adults may experience less predation pressure per capita from a constant level of predation, thus more juveniles could survive that do not express traits that are advantageous to avoid predation. If these fish survive to spawn, they could produce more offspring that inherit those traits, which may limit our ability to detect a difference between the two origins. Supplementation may initially decrease the per capita predation pressure on fry because it increases the abundance of fry relative to predators. However, it is likely that the predator population will eventually increase in abundance if more prey continues to be available during the "building stage" of supplementation (Pearsons 2002). The opposite would be true for the offspring of a weaker run of adults and there could be greater differences in the two origins that we would be able to detect. Thus, in some years the per capita predation selection could be very similar in natural and hatchery environments (no selection) and in other years the selection differential could be large. In addition, because we do not want to adversely impact the Naches population, we use far fewer adults as broodstock for the study fry than is used for the hatchery and supplementation fry. This greatly increases the chances that individual differences in the Naches adults that we collect will influence the results of the study and therefore may not be representative of the Naches population in some years.

Steps were taken to ensure any differences that are detected in survival can be attributed to genetic differences. The great care that is taken to size match the fry is important to ensure the results are not affected by size-influenced predation. Smaller fish may be more vulnerable to predators because of slower swimming speed (Taylor and McPhail 1985) or less likely to be gape limiting to a predator (Pearsons and Fritts 1999). Studies have shown that smaller salmonids are more vulnerable to predators than larger salmonids (Patten 1977; Hargreaves and LeBrasseur 1986), such as was evident all but one year (2012) in the small and large fry that were individually marked during the trials. The sizes of the predators in each net pen were similar in order to decrease the chances of differential size selective predation. Alternating the mark type each origin of fish received between net pens ensured that any marking effect would not influence our
conclusions. Although, it is unlikely, we cannot exclude the possibility that these findings were influenced by a maternal effect (Heath and Blouw 1998). However, if maternal effects were occurring one would expect to see significant changes in survival as fish get older because it is believed that maternal effects are most pronounced in young fish (Heath et al. 1999). We did not detect changes in survival through time suggesting that maternal effects were not prevalent in our experiment. Finally, we found very small to no difference in the background mortality of the three groups.

The results of this experiment are also more likely to be representative of the whole population than other studies because we tested the offspring of far more families than any other study of predation and domestication that we are aware of. Only testing a few families increases the chances that any differences would be due to a single adult that had genetic attributes that made them exceptionally good or poor at avoiding predators. Using two types of predators also ensured that the fry would require a more complete suite of predator avoidance tactics. During short observations immediately after introduction of the fry into to net pens, the fry were observed to form a single school and swim along the bottom of the net pens where two or three predatory attacks by the sculpins would be witnessed within the first five minutes. When the fry were recovered on the last day of the trials, they were generally higher in the water column beneath the overhead cover where they were safe from the sculpin but still vulnerable to the trout. Qualitative observations of the stage of decomposition of fry in the stomach contents of the predators showed that both species consumed several fry and that the sculpins consumed most of the fry early during the trial while the trout consumed fry throughout the duration of the trial.

Because the prey fish were treated identically, any differences found should be due to genetic differences and not abnormal behavior that is learned in the hatchery environment. This means that any differences that we find could be expressed in the natural environment. However, because the experiments were conducted in an artificial environment, we do not know how differences will be manifested in the natural environment. For example, in years of low predation pressure, no differences in survival of the offspring may occur..

This study that has shown a diversity of results thus far. Of the peer-reviewed literature that have found predation differences due to origin, those studies have only lasted one or two years and have generally represented a smaller number of families (Table 17). It is important to evaluate behavioral studies for multiple years using high numbers of families because of the annual differences in selection pressures and variability between individuals within populations.

Table 17. A comparison of studies that have tested the effects of domestication on predation vulnerability. Species tested, origins compared, number of generations under hatchery culture, founding stock, rearing environment, years tested, number of families tested, and the metric used to asses vulnerability are compared.

| Study | Species | Comparison | $\begin{gathered} \mathrm{H} \\ \text { gen } \end{gathered}$ | Stock ${ }^{\text {a }}$ | Rearing ${ }^{\text {b }}$ | Yrs | Families | Metric ${ }^{\text {c }}$ |
| :---: | :---: | :---: | :---: | :---: | :---: | :---: | :---: | :---: |
| 1 | Brown trout | wild vs. hatchery | 1-2 | S? | D | 1 | 5-7 | B |
| 2 | Steelhead | wild vs. hatchery | 1-7 | S | S | 1 | 7-10 | M |
| 3 | Brown <br> trout | wild vs. hatchery | 5 | S | S | 1 | 9 | B |
| 4 | Atlantic salmon | wild vs. farmed | 7 | S | S | 1 | 8 | B |
| 5 | Steelhead rainbow trout | wild vs. wild/farmed hybrid | 5+ | D | S | 1 | 11 | B |
| 6 | Brown trout | wild vs. hatchery/wild hybrid | 5 | S | S | 1 | Up to 64 (mixture) | B |
| 7 | Atlantic salmon | wild vs. farmed | 7 | S | S | 2 | ? | B |
| 8 | Masu salmon | wild vs. hatchery vs. farmed | 7+ | D | D | 1 | ? | B |
| This study | Chinook salmon | Supplemented vs. hatchery w/ wild control | 1-2 | S | S | 8 | 12-59 | M |

${ }^{1}$ Alvarez and Nicieza (2003); ${ }^{2}$ Berejikian (1995); ${ }^{3}$ Ferno and Jarvi (1998); ${ }^{4}$ Fleming and Einum (1997); ${ }^{5}$ Johnsson and Abrahams (1991); ${ }^{6}$ Johnsson et al. (1996); ${ }^{7}$ Johnsson et al. (2001);
${ }^{8}$ Yamamoto and Reinhardt (2003); *Present study
${ }^{\text {a }}$ Same (S) or different (D) founder stock.
${ }^{\text {b }}$ Same (S) or different (D) rearing environment.
${ }^{\mathrm{c}}$ Behavior (B) or mortality (M).

The first two years of the study (2003-2004) were the last two years where we had the opportunity to use offspring of truly wild spring Chinook from the upper Yakima River because the first adult returns from the Cle Elum Hatchery spawned naturally in 2001. There was a slight chance that a naturally produced jack used for 2003 brood (2004 study population) could have been sired by a hatchery jack in 2000 but we consider that unlikely given the small proportion of hatchery jacks in 2000 relative to the wild population. The hatchery control population began with the spawning of returning hatchery origin fish in 2002, our 2003 study population. Brood year 2013 completed the third generation of the hatchery control population and the final year of this study. We will perform a comprehensive analysis of the three generations of data and produce a final report of this work to be included in the next annual report.

## Competitive dominance

We have observed the full range of possible outcomes in dominance between supplementation and hatchery fish. Supplementation fish dominated hatchery fish in 2005, opposite results were found in 2006, 2008 and 2013, and neither was dominant in 2007, 2009, 2010, 20112012 or 2014. At this time we cannot think of any compelling reason why offspring of wild (2003-2004, Pearsons et al. 2007) and the early supplementation population (2005, with minimal natural spawning first generation hatchery influence) appeared to dominate the hatchery population during the first three years of this study. Since that time, there has been no obvious trend of one group becoming more dominant. Lynch and O’Hely (2001) predict that it typically takes 10 to 20 generations for a supplemented population to reach $50 \%$ equilibrium in terms of the genetic load from captive breeding depending on strengths of selection in the hatchery and natural environments and proportion of hatchery fish spawning in the wild. If this is the case, then it seems reasonable that the hatchery and supplementation fish could exhibit this flip-flopping of dominance between years for quite some time as deleterious alleles are expressed at different rates depending on environmental pressures and the proportion of hatchery fish on the spawning grounds until they begin reaching equilibrium.

The differences in dominance and aggression that we have observed were likely due to an interaction between genetic changes that occurred from fish culture, differences in stocks, and a year effect. However, we cannot exclude the possibility that changes were caused by a maternal rearing environment effect (e.g., not a genetic effect). This might occur if hatchery rearing caused phenotypic differences in females that were passed on to progeny. We believe that this was unlikely to have had much of an effect on our experiments because 1) egg sizes of hatchery and wild fish were not significantly different (Knudsen 2005), and 2) fish were tested approximately 4 months after hatching. Most studies that have reported maternal effects in fish have documented relationships between female size and progeny size (Heath and Blouw 1998). We attempted to control for size effects by size matching our fish. Maternal effects are more likely to occur when fish are very young. In a review of maternal effects in fish, Heath and Blouw (1998) concluded "maternal effects in fishes are usually negligible beyond the early juvenile life stages."

With the exception of this study, annual differences in competitive dominance associated with domestication have generally not been evaluated. Most studies that have evaluated this topic are based on one year of study and none have been longer than two years (Table 18). The study presented in this report combined with the work presented in Pearsons et al. (2007) represent twelve years of study. We have seen considerable annual differences in our results. If we had restricted our study to a single year, then we may have concluded that domestication positively, negatively, or neutrally influenced competitive dominance. This finding suggests that we should use caution when interpreting dominance results that do not evaluate multiple years of study.

In comparison to our observations, juvenile coho salmon reared in hatcheries have been documented to be more aggressive than wild fish (Swain and Riddell 1990; Berejikian et al. 1999) or less aggressive (Berejikian et al. 1996). Furthermore, Einum
and Fleming's (2001) meta-analysis of aggression revealed that hatchery fish were more aggressive than wild fish. We suspect that the differences in findings are caused by 1) the duration and type of hatchery practices, and 2 ) differences in the rearing environment of the fish tested. Most, if not all, of the studies that have previously been conducted outside of the Yakima Basin have used hatchery fish that have been under culture for several generations and frequently these are of non-local origin (Table 21). If genetic changes or maternal effects are additive, then it is likely that larger differences in aggression will be detected with each additional generation of fish culture. Furthermore, fish that are collected from natural environments and compared to fish reared in hatchery environments are likely to produce differences because of the differences in rearing conditions. For example, in another study, we found that spring Chinook smolts reared in the hatchery dominated salmon smolts that were reared in the Yakima River. Larger fish generally dominated smaller fish, but the size difference did not have to be as great for hatchery fish to dominate as wild fish (Pearsons et al.,WDFW, unpublished data). In short, hatchery fish were dominant over wild fish in contest competition experiments unless wild fish were sufficiently larger than hatchery fish. In a study of coho salmon, Rhodes and Quinn (1998) reported similar findings.

Table 18. Comparison of dominance studies that relate to domestication selection of salmonids of varying origins.

| Study | Species | Comparison ${ }^{\text {a }}$ | Hatch. gener. | Stock | Yrs | Number of families | Trial $\text { type }^{\text {b }}$ | Replicates | Metric ${ }^{\text {c }}$ |
| :---: | :---: | :---: | :---: | :---: | :---: | :---: | :---: | :---: | :---: |
| 1 | Steelhead | W vs. H | 4-7 | Same | 1 | $13 \mathrm{~W} ; 18 \mathrm{H}$ | C | 16 | A, P, C |
| 2 | Coho | W vs. C | 1 | Different | 1 | 15 | C | 44 | A, P |
| $3{ }^{\text {e }}$ | Atlantic | W vs. F | 6-10 | Different | 1 | ? | C | 218 | P |
| 4 | Grayling | W vs. W <br> H vs. H | 2 | Same | 2 | ? | ? | 30 ? | A |
| 5 | Coho | $\begin{aligned} & \text { W vs. W } \\ & \text { H vs. H } \end{aligned}$ | 5 | Different | 1 | $\begin{aligned} & 10 \& 13 \mathrm{~W} \\ & 11 \& 191 \mathrm{H} \end{aligned}$ | M | 21? | A |
| $6^{\text {f }}$ | Chinook | H vs. H | 1 vs. 5 | Same | 1 | 5 | C | 40 | P, F, A |
| $7^{8}$ | Chinook | W vs. H | 1 | Same | 1 | 6 | C | 89 | P, F, A |
| 8 | Chinook | W vs. H | 1 | Same | 2 | 54-59 | C, S | 229, 276 | P, F, A |
| 9 | Atlantic | W vs. F | 7 | Different ${ }^{\text {d }}$ | 1 | $6+\mathrm{W}, 8 \mathrm{~F}$ | C | 30 stream | P, F, A |
| 9 | Atlantic | W vs. F | 7 | Different ${ }^{\text {d }}$ | 1 | $6+\mathrm{W}, 8 \mathrm{~F}$ | C | 15 tank | A |
| 10 | Brown trout | W vs. H | 10 | Same | 1 | Up to 64 (mixture) | C | 12 | A |
| This study | Chinook | S vs. H <br> w/ W control | 1-2 | Same | 8 | 23-52 | C, S | 157-299 | P, F, A |

${ }^{1}$ Berejikian et al. (1996); ${ }^{2}$ Berejikian et al. (1999); ${ }^{3}$ Metcalfe et al. (2003); ${ }^{4}$ Salonen and Peuhkuri (2004); ${ }^{5}$ Swain and Riddell (1990); ${ }^{6}$ Wessel et al. (2006); ${ }^{7}$ Farrell et al. (2003); ${ }^{8}$ Pearsons et al. (2007); ${ }^{9}$ Fleming and Einum (1997); ${ }^{10}$ Johnsson et al. 1996; *Present study.
${ }^{\text {a }}$ Offspring of wild (W), supplementation (S), hatchery or sea-ranched (H), farmed (F), and captive brood (C).
${ }^{\mathrm{b}}$ Contest (C), mirror image (M), and Scramble (S).
${ }^{\mathrm{c}}$ Metrics used to assess dominance are aggression (A), position (P), color (C), and food (F).
${ }^{\mathrm{d}}$ Farmed population founded, in part, from wild population.
${ }^{\text {e}}$ Subjects were not size matched.
${ }^{\mathrm{f}}$ Subjects were within $\pm 3 \mathrm{~mm}$ FL.
${ }^{\text {g }}$ Subjects were within 4\% FL.
The results presented in this chapter are part of a long-term study that attempts to evaluate if hatchery supplementation alters competitive dominance relative to an unsupplemented reference population and a hatchery population. Now that the third
generation of the hatchery control population is complete, we will perform an analysis of the entire dataset and produce a final report on this work.

## Spring Chinook reproductive success/spawning channel

Approximately 93 percent successes were achieved at inferring parent-offspring relationships. Examination of Table 4 reveals a very uneven pattern of reproductive success among the candidate parents. Based on the subsample of 2,545 fry that were successfully assigned parents, the range of inferred reproductive output among males was $0-370$ fry; the range for the same period in reproductive output among females was 0 197 fry. Some of the dam-sire matings we inferred are well supported (there were a lot of fry assigned to them) and some are weakly supported (not many fry were assigned to them). Caution should be used when interpreting dam-sire-fry combinations that were inferred rarely. Future integration of fecundity estimates for spawners will enrich the interpretation of these estimates of reproductive output.

Interpretation of the inferred parental reproductive output based on parentage assignments by genetic analysis requires the consideration and analysis of individual fish attributes, including fecundity and body size, the closed nature of the experimental environment in which sub-dominant males had a more limited number of alternative females to court than they might have had in an open system, and relative stocking levels and synchronicity of spawning.

## Spring Chinook Genetic stock separation-juveniles

Collection of smolts at the Chandler Trap in 2013 utilized a sampling design intended to yield a sample that was proportional to the number of smolts passing the Chandler Trap. Sampling a proportional number of smolts was important to determine an accurate percentage of smolts from each stock that were outmigrating from the basin. Developing the sampling strategy for identifying a "standard" versus "peak" day of smolts that were in the trap and applying a sampling goal for those days allowed for a proportional sample. Subsampling the smolts collected for genetic analysis provided a best fit to the actual passage of smolts for a given day.

Monitoring the relative abundances of Chinook smolts in the Yakima River from the three different populations of spring Chinook (upper Yakima River, American River, and Naches River) and the two populations of fall Chinook (Marion Drain and lower Yakima River) requires the ability to estimate population composition of smolts outmigrating past Chandler trap. Because all five Chinook populations are intermingled when they pass Chandler trap, and the vast majority are unmarked and untagged, the only way to determine population-of-origin is by genetic analysis. This method requires that sufficient genetic differences exist among these populations in the Yakima River basin.

A baseline of 19 individual collections from the five populations in the Yakima River basin was used for the population-of-origin assignments of the outmigrating smolts. The baseline collections as a whole had higher genotyping failure compared to the Chandler smolt samples. Scales were taken from carcasses on spawning grounds for
most baseline collections; therefore, DNA quality was presumably poorer than the Chandler smolt collection where tissue was collected from live fish. The upper Yakima River tissue collections were also taken from live fish at the hatchery and, therefore, genotyping success was higher for this collection than the other baseline collections.

Assessment of spring or fall smolts by morphological and genetic analysis revealed agreement with 55 individuals being identified differently between the two methods. Identification as a spring or fall smolt was the same for 1,125 smolts collected during the January - February, March, April, May, and June - July time strata.

The majority of the assignments between January and May were from the three spring stocks. The upper Yakima River spring stock accounted for the highest average percentage ( $76.2 \%$ ) of smolts present in that period. Rank in abundance of the three spring stocks was the same in the three time strata (January-February, March, April, and May) with upper Yakima River spring stock having the most. The June-July time stratum was predominately composed of the fall Chinook stocks, accounting for over $70.6 \%$ of the total number of smolts.

## Assessment of DNA Mixture Assignments from 2000-2013

Mixed stock analysis has been conducted on Chandler smolts since 2000 (Young 2004, Kassler et al. 2005, Kassler 2006, Kassler and VonBargen 2007, 2008, 2009 and 2010, Kassler and Peterson 2011, Kassler and Bell 2012, Kassler and Bowman 2013); however the sampling design for samples collected in 2000-2003 was not proportionalized during the run. The yearly assignments are therefore not comparable from those years. Beginning in 2004, staff at the Chandler trap utilized a sampling protocol to provide a number of smolts that was relative to the percentage of smolts passing that day. Samples were then subsampled at WDFW to provide a proportional number of samples that would represent the overall passage to be analyzed.

## b. Hatchery RM\&E

The performance of the YKFP spring Chinook supplementation program has been documented relative to the project quantitative objectives and has been presented annually in the YKFP M\&E project overview (Fritts 2012). Briefly, the project appears to be meeting or is making progress towards achieving the project's objectives (Appendix C).

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## Appendix A: M\&E Project Publication List

The following publication list includes technical reports and peer reviewed publications that have been produced from the work under the Yakima/Klickitat Fisheries Project's monitoring and evaluation program.

Amaral, S. V., F. C. Winchell, and T. N. Pearsons. 2001. Reaction of Chinook salmon, northern pikeminnow, and smallmouth bass to behavioral guidance stimuli. Pages 125-144 in C. C. Coutant, editor. Behavioral technologies for fish guidance. American Fisheries Society, Symposium 26, Bethesda, Maryland.

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## Appendix B: Use of Data \& Products

Raw electronic data files (Database) are secured on the WDFW Corporate server in Olympia, WA, as well as on WDFW district 8 field office personal computers. Data housed on personal computers are duplicated on the local office server which is in turn backed up on the WDFW corporate server in Olympia, WA nightly.

## Appendix C. Performance measures relative to project quantitative objectives

| Performance <br> Measure | Goal | Performance | Comments |
| :---: | :---: | :---: | :---: |
| Natural <br> Production of Target Species | Increase while maintaining the longterm fitness of the target population (see quantitative objectives; Pearsons et al. 2006) | Quantitative objectives for adults and smolts are being achieved. Differences in traits of hatchery and natural origin fish are a concern | - Too early to evaluate conclusively, but strategies to reduce genetic risk are being implemented. <br> - Hatchery has increased the number and distribution of adult spawners on the spawning grounds. Quantitative management objectives for natural production of upper Yakima and basin total spring Chinook adults and smolts are being achieved. <br> - Significant but small changes in many demographic and reproductive success traits indicate cause for concern. <br> - Predation and competition may be limiting natural production objectives and may constrain the benefits of supplementation. |
| Harvest | Increase (see quantitative objectives; Pearsons et al. 2006) | Increased, and objectives are being met | - Tribal subsistence fisheries occurred on both hatchery and naturally produced fish in all years. Sport fisheries on hatchery fish have also occurred in the Yakima River in 10 of the 14 years since 2001. <br> - Quantitative harvest objectives for the upper Yakima stock and all Yakima basin stocks combined are |


|  |  |  | being met for the Columbia or <br> Yakima Rivers |
| :--- | :--- | :--- | :--- |
| Genetics | Minimize genetic <br> impacts to non-target <br> taxa | Achieved to <br> date | Stray rates are very low |
| Ecology | Keep impacts to non- <br> target taxa within <br> containment <br> objectives | Achieved for <br> most taxa to <br> date | Impacts for most species are <br> within containment objectives or <br> are currently not attributable to <br> supplementation. |
| Habitat | Protect the most <br> productive stream <br> reaches and increase <br> productivity/capacity <br> of freshwater <br> environment so that <br> quantitative <br> objectives can be <br> achieved. | Progress | Habitat protection, restoration, <br> and tributary passage efforts are <br> ongoing, with incremental <br> progress each year. |
| Science | Disseminate <br> important findings for <br> use throughout the <br> Yakima Basin, <br> Columbia Basin, and <br> world | Achieved to <br> date <br> the benefits of supplementation, <br> especially over the long-term. |  |

# Appendix D. DNA-Based Parentage Assignments of Chinook Salmon from the Cle Elum Spawning Channel in 2013 

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#### Abstract

We used a maximum likelihood parentage assignment procedure to estimate the reproductive output of Chinook salmon spawners from the hatchery-control line (two generations of hatchery influence) and the supplementation hatchery line (SH - one generation of hatchery influence) in the Cle Elum experimental spawning channel for the 2012 brood year. The assignments were based on offspring genotypes at 14 microsatellite loci. The probabilities of exclusion (inferring non-parentage by randomly picked adults) assuming neither parent was known were estimated to be 0.999999 . Two thousand five hundred and forty-eight of 2,741 fry from the 2012 brood that were genotyped at nine or more loci were assigned to a parental pair with $95 \%$ confidence. The number of progeny attributed to individual potential parents was quite variable, ranging from 0 to 370 for all males and from 0 to 197 for females. The sum of progeny attributed to the hatchery-control line males and females was 766, while the sum of progeny attributed to supplementation hatchery line males and females was 1,779.


## Introduction

Although hatcheries have been extensively utilized in Chinook salmon management for over 100 years, only recently have rigorous experiments been developed to measure the relative reproductive success of hatchery- and natural-origin spawners in a shared natural setting. Some of the difficulty in designing informative studies has stemmed from the challenges of controlling entry to natural spawning areas and collecting representative samples of recently hatched fry. Furthermore, if control could be established over the potential spawners in the spawning area, the measurement of individual reproductive output still would require a means of associating individual fish captured in one year with individuals that spawned in a previous year. The spawning behavior of Chinook salmon adds to the complexity of quantifying individual reproductive output through behavioral observations: at a redd site, a female might be courted by several males that compete for access to the female, providing opportunities for multiple paternity in a single redd. In areas with moderate to high spawning densities, males might attend females on several adjacent redds. Microsatellites, a class of highly polymorphic, codominant DNA markers, provide a means to quantify individual spawners' reproductive output. A suite of 10 to 15 highly variable microsatellites can resolve individual identity in a moderate to large population, and through a simple inheritance model, can illuminate parent-offspring relationships.

Washington Department of Fish and Wildlife (WDFW) and the Yakama Nation (YN) are cooperating on a study of Chinook salmon reproductive success in a presumably closed access spawning observation channel at the Cle Elum Hatchery. Viewing blinds line the channel, allowing researchers to observe spawning activities.

Chinook salmon carrying visible external marks were released into the spawning channel in September 2012. Hatchery-control line (two generations of hatchery influence) males and females were released into three of six shared spawning areas and supplementation hatchery line (one generation of hatchery influence) males and females were released into the other three shared spawning areas to select and compete for mates. Prior to the release of the potential spawners, researchers collected and preserved samples of fin tissue to enable genetic characterization of the potential spawners and to allow subsequent inference of parent/offspring relationships after juveniles were collected and genotyped. One group of researchers examined morphological characteristics of these potential parents and observed and recorded spawning area behaviors and interactions. The results of the morphological and behavioral work are described in a separate report.

The potential parents' fin tissue samples and the collected progeny (fry) were delivered to the WDFW Molecular Genetics Laboratory in Olympia, Washington for genetic screening and parentage analysis following the same protocols that have been used from 2002 - 2007, 2009 - 2013 (Young and Kassler 2005, Kassler 2005, Kassler 2006, Kassler and Von Bargen 2007, 2008, and 2010, Kassler et al. 2011; Kassler and Peterson 2012, 2013). The genetic analyses provide direct, quantitative estimates of fry production by individual spawning Chinook salmon. This report presents the parentage results for the 2012 - 2013 Cle Elum spawning channel experiments.

## Materials and Methods

Collection of potential spawners - 2012
Fin tissue was collected from a total of 48 adult females and 48 adult males (Table 1) prior to their release into each of six sections in the spawning channel during September 2012. The genetic analysis program CERVUS (version 3.0; Marshall et al. 1998) was used to check for identical multilocus genotypes among the potential parents. Data recorded for each released fish included gender, and whether it was of hatchery-control line origin or supplementation hatchery line origin (Table 1).

## Collection of Fry

Fry collections occurred from November 30, 2012 to April 25, 2013. Fry samples were collected from each section daily when fry were present. During that period a total of 2,979 fry were collected.

## DNA Extraction Methods

Genomic DNA was extracted by digesting a small piece of fin tissue using the nucleospin tissue kits obtained from Macherey-Nagel following the recommended conditions in the user manual. Extracted DNA was eluted with a final volume of $100 \mu \mathrm{~L}$.

## PCR Methods

Potential spawners and offspring from 2013 were genotyped at 14 loci (Table 2).
Potential spawners were screened twice and scored independently at all 14 loci by two biologists to minimize potential genotyping error of the parents.

The polymerase chain reaction mixture contained the following for a $10 \mu \mathrm{l}$ reaction: approximately 25 ng template DNA, 1X Promega buffer, $1.5 \mathrm{mM} \mathrm{MgCl} 2,200 \mu \mathrm{M}$ each of dATP, dCTP, dGTP, and dTTP, approx. $0.1 \mu \mathrm{M}$ of each oligonucleotide primer, and 0.05 units GoTaq Flexi DNA polymerase (Promega). Amplification was performed using MJ Research PTC-200 and AB 9700 thermocyclers. The thermal profile was as follows: an initial denaturation step of 2 minutes at $94^{\circ} \mathrm{C} ; 40$ cycles of 15 seconds at $94^{\circ} \mathrm{C}, 30$ seconds at $49-58^{\circ} \mathrm{C}$, and 1 minute at 72 oC ; plus a final extension step at $72^{\circ} \mathrm{C}$ for 10 minutes, followed by a final indefinite holding step at $4^{\circ} \mathrm{C}$.

Microsatellite DNA loci (Table 2) were amplified via the polymerase chain reaction (PCR) using fluorescently labeled primers (obtained from Applied Biosystems or Integrated DNA Technologies). Loci were combined into multiplexes to increase efficiency and decrease costs.

Data were collected using an AB-3730 Genetic Analyzer. Applied Biosystems GENEMAPPER v.3.7 software was used to collect and analyze the raw data and to determine genotypes at each locus (based on estimated allele sizes in base pairs using an internal size standard). Alleles were binned in GENEMAPPER using the standardized allele sizes established for the Chinook coastwide standardization efforts (Seeb et. al. 2007).

## Parentage Assignments

The dataset included 37,716 single-locus genotypes. A genotyping error rate in that dataset of $1.0 \%$ would result in 377 incorrect single-locus genotypes. Our error rate is unknown, but possibly greater than $1 \%$. Since parentage analyses involve comparing genotypes of candidate parental pairs with offspring genotypes, genotyping errors can produce parent-offspring genotype mismatches and suggest exclusion of true parentoffspring pairings from consideration. Alternatively, genotyping errors can lead to failure to exclude parent-offspring pairings that are incorrect. We used a maximum likelihood procedure, implemented in CERVUS (version 3.0; Marshall et al. 1998) to infer parent-offspring relationships. The procedure uses allele frequency data to assign likelihoods to parent-offspring combinations, and allows mismatching genotypic data to be evaluated concurrently with matching genotype data.

Genotyping error is not the only potential source of mismatches between the genotypes of fry and their putative parents. We would expect allele misidentification to be randomly distributed throughout the genotype dataset and not to occur in clusters. Parent-offspring mismatches can result also from germ-line mutation in which a parent passes a changed allele to its offspring or from the inadvertent exclusion of one or more contributing parents from the parental dataset. These mismatches are due to correctly assigned but unexpected genotypes, and we expect that those genotypes should cluster in families. Distinguishing between mutation-based mismatches and mismatches that result from reproductive participation by un-genotyped parents is difficult. Assuming that all dams in the experimental channel are represented in the parental data set, we might suspect reproductive participation by one or more unrepresented sires if groups of fry that are assigned to a dam-offspring relationship with no mismatching loci, have multiple locus mismatches with all candidate sires, and no more than four alleles at a locus within the group. The data set was carefully examined for evidence of reproductive contributions by such un-genotyped parents (because evidence of ungenotyped parents had been observed in previous years).

## Results

## Parents

Genetic analysis revealed that all 96 fish released or found in the spawning channel had unique genotypes. There were a total of 24 hatchery control line (HC) adult males, 24 HC adult females, 24 supplementation hatchery line (SH) adult males, and 24 SH adult females. Four HC males and four HC females were released into three of the six sections and four SH males and four SH females were released into the other three sections (Table 1).

## Loci Screened

A total of 14 loci were screened and all 14 were used in the analysis (Table 2). Number of alleles ranged from 4-30 (Ots-9 and Omm-1080 respectively) and observed heterozygosity ranged from $0.448-0.958$ (Ots-G474 and Ots-201b respectively). Individual exclusionary power was below 45.4\% for five loci (Ogo-2, Ogo-4, Ots-G474, Ots-3M, and Ots-9) and above $61.6 \%$ for the remaining loci when neither parent was known. Exclusionary power was below 42.1\% for three loci (Ots-G474, Ots-3M and Ots9) and above $60.2 \%$ for the remaining loci when one parent was known. Cumulative exclusionary power was 1.000000 for analysis using all loci when one parent was known.

## Parentage Assignments

Parentage assignments were made when genotype data was available for nine or more loci. All 96 parents were genotyped at 10 or more loci while 2,741 of the 2,784 offspring were successfully genotyped at nine or more loci (Table 3).

Parentage analysis was conducted independently for each of the six sections using all 96 adults as possible parents. Each fry was assigned a dam-sire-fry combinations (trios) based on the most likely candidate male parents (sires) and female parents (dams). Those assignments yielded possible. Any fry-sire assignments with more than two mismatching loci were excluded from further consideration.

Of the total 2,741 fry included in the analysis a total of 2,545 , fry were assigned to a single male and female parent (2,545/2,741 = 92.8\%).

## Discussion

Approximately 93 percent successes were achieved at inferring parent-offspring relationships. Examination of Table 4 reveals a very uneven pattern of reproductive success among the candidate parents. Based on the subsample of 2,545 fry that were successfully assigned parents, the range of inferred reproductive output among males was $0-370$ fry; the range for the same period in reproductive output among females was 0 197 fry. Some of the dam-sire matings we inferred are well supported (there were a lot of fry assigned to them) and some are weakly supported (not many fry were assigned to them). Caution should be used when interpreting dam-sire-fry combinations that were inferred rarely. Future integration of fecundity estimates for spawners will enrich the interpretation of these estimates of reproductive output.

Interpretation of the inferred parental reproductive output based on parentage assignments by genetic analysis requires the consideration and analysis of individual fish attributes, including fecundity and body size, the closed nature of the experimental environment in which sub-dominant males had a more limited number of alternative females to court than they might have had in an open system, and relative stocking levels and synchronicity of spawning.

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Table 1. Potential Chinook salmon spawners in the six section of the Cle Elum experimental spawning channel in 2012. Origin is identified as hatchery-control (HC) or supplementation hatchery (SH).

| Origin | Section $1-1 \mathrm{~A}$ Females | Section $1-2 \mathrm{~A}$ Females | Section 1 - 3A Females | Section $2-1 \mathrm{~A}$ Females | Section $2-2 \mathrm{~A}$ Females | Section 2 - 3A Females |
| :---: | :---: | :---: | :---: | :---: | :---: | :---: |
| SH | 4 | -- | 4 | -- | 4 | -- |
| HC | -- | 4 | -- | 4 | -- | 4 |
|  | Males | Males | Males | Males | Males | Males |
| SH | 4 | -- | 4 | -- | 4 | -- |
| HC | -- | 4 | -- | 4 | -- | 4 |
|  | Section 1-1B | Section 1-2B | Section 1-3B | Section 2-1B | Section 2-2B | Section $2-3 B$ |
| Origin | Females | Females | Females | Females | Females | Females |
| SH | 4 | -- | 4 | -- | 4 | -- |
| HC | -- | 4 | -- | 4 | -- | 4 |
|  | Males | Males | Males | Males | Males | Males |
| SH | 4 | -- | 4 | -- | 4 | -- |
| HC | -- | 4 | -- | 4 | -- | 4 |

Table 2. Locus summary.
Exclusionary power

| Locus | \# alleles | \# parents genotyped | $\mathrm{H}_{\mathrm{O}}$ (observed) | HE (expected) | neither parent | one parent |
| :---: | :---: | :---: | :---: | :---: | :---: | :---: |
| Oki-100 | 19 | 95 | 0.926 | 0.904 | 0.669 | 0.802 |
| Ots-201b | 21 | 95 | 0.905 | 0.901 | 0.657 | 0.793 |
| Ots-208b | 25 | 95 | 0.958 | 0.944 | 0.780 | 0.876 |
| Ssa-408 | 17 | 95 | 0.726 | 0.900 | 0.652 | 0.790 |
| Ogo-2 | 7 | 96 | 0.865 | 0.795 | 0.423 | 0.602 |
| Ssa-197 | 21 | 96 | 0.927 | 0.915 | 0.694 | 0.819 |
| Ogo-4 | 10 | 96 | 0.760 | 0.815 | 0.454 | 0.629 |
| Ots-213 | 21 | 96 | 0.917 | 0.927 | 0.728 | 0.843 |
| Ots-G474 | 8 | 96 | 0.448 | 0.443 | 0.108 | 0.263 |
| Omm-1080 | 30 | 96 | 0.938 | 0.944 | 0.782 | 0.877 |
| Ots-3M | 7 | 96 | 0.667 | 0.658 | 0.249 | 0.421 |
| Ots-211 | 23 | 86 | 0.977 | 0.924 | 0.721 | 0.838 |
| Ots-212 | 20 | 92 | 0.891 | 0.887 | 0.616 | 0.763 |
| Ots-9 | 4 | 92 | 0.696 | 0.661 | $0.233$ | 0.389 |

Table 3. Summary of genotyping efficiency in potential parents and offspring.

| Loci genotyped | Parents (12IQ) | Offspring (13MO) |
| :---: | :---: | :---: |
| 14 | 82 | 1,879 |
| 13 | 9 | 322 |
| 12 | 3 | 199 |
| 11 | 1 | 180 |
| 10 | 1 | 85 |
| 9 | 0 | 76 |
| 8 | 0 | 3 |
| 7 | 0 | 8 |
| 6 | 0 | 6 |
| 5 | 0 | 2 |
| 4 | 0 | 1 |
| 3 | 0 | 1 |
| 2 | 0 | 0 |
| 1 | 0 | 1 |
| 0 | 0 | 21 |

Table 4. Total number of offspring assigned to females and males from each of the six sections in the spawning channel and the life stage (HC - hatchery control line; SH - supplementation hatchery line) for each fish.

| Females | Section | HC or SH | Total Offspring | Males | Section | HC or SH | Total Offspring |
| :---: | :---: | :---: | :---: | :---: | :---: | :---: | :---: |
| 12IQ0003 | 1-1A | SH | 18 | 12IQ0001 | 1-1A | SH | 4 |
| 12IQ0004 | 1-1A | SH | 30 | 12IQ0002 | 1-1A | SH | 144 |
| 12IQ0005 | 1-1A | SH | 56 | 12IQ0006 | 1-1A | SH | 33 |
| 12IQ0008 | 1-1A | SH | 77 | 121Q0007 | 1-1A | SH | 0 |
| 12IQ0009 | 1-2A | HC | 14 | 12IQ0011 | 1-2A | HC | 34 |
| 12IQ0010 | 1-2A | HC | 42 | 12IQ0012 | 1-2A | HC | 5 |
| 12IQ0013 | 1-2A | HC | 103 | 12IQ0015 | 1-2A | HC | 0 |
| 12IQ0014 | 1-2A | HC | 0 | 12IQ0016 | 1-2A | HC | 118 |
| 12IQ0017 | 1-3A | SH | 72 | 12IQ0021 | 1-3A | SH | 289 |
| 12IQ0018 | 1-3A | SH | 197 | 12IQ0022 | 1-3A | SH | 0 |
| 12IQ0019 | 1-3A | SH | 93 | 12IQ0023 | 1-3A | SH | 59 |
| 12IQ0020 | 1-3A | SH | 11 | 12IQ0024 | 1-3A | SH | 25 |
| 12IQ0025 | 2-1A | HC | 56 | 12IQ0026 | 2-1A | HC | 86 |
| 12IQ0027 | 2-1A | HC | 0 | 12IQ0028 | 2-1A | HC | 73 |
| 12IQ0029 | 2-1A | HC | 60 | 12IQ0030 | 2-1A | HC | 0 |
| 12IQ0031 | 2-1A | HC | 43 | 12IQ0032 | 2-1A | HC | 0 |
| 12IQ0033 | 2-2A | SH | 0 | 121Q0037 | 2-2A | SH | 0 |
| 12IQ0034 | 2-2A | SH | 10 | 12IQ0038 | 2-2A | SH | 145 |
| 12IQ0035 | 2-2A | SH | 17 | 12IQ0039 | 2-2A | SH | 0 |
| 12IQ0036 | 2-2A | SH | 118 | 12IQ0040 | 2-2A | SH | 0 |
| 12IQ0045 | 2-3A | HC | 0 | 12IQ0041 | 2-3A | HC | 0 |
| 12IQ0046 | 2-3A | HC | 4 | 12IQ0042 | 2-3A | HC | 0 |
| 12IQ0047 | 2-3A | HC | 92 | 12IQ0043 | 2-3A | HC | 101 |
| 12IQ0048 | 2-3A | HC | 5 | 12IQ0044 | 2-3A | HC | 0 |
| 12IQ0049 | 1-1B | SH | 0 | 12IQ0053 | 1-1B | SH | 118 |
| 12IQ0050 | 1-1B | SH | 6 | 12IQ0054 | 1-1B | SH | 83 |
| 12IQ0051 | 1-1B | SH | 108 | 12IQ0055 | 1-1B | SH | 0 |
| 12IQ0052 | 1-1B | SH | 87 | 12IQ0056 | 1-1B | SH | 0 |
| 12IQ0061 | 1-2B | HC | 64 | 12IQ0057 | 1-2B | HC | 0 |
| 12IQ0062 | 1-2B | HC | 0 | 12IQ0058 | 1-2B | HC | 0 |
| 12IQ0063 | 1-2B | HC | 0 | 12IQ0059 | 1-2B | HC | 0 |
| 12IQ0064 | 1-2B | HC | 0 | 12IQ0060 | 1-2B | HC | 64 |
| 12IQ0066 | 1-3B | SH | 56 | 12IQ0065 | 1-3B | SH | 9 |
| 12IQ0067 | 1-3B | SH | 87 | 12IQ0068 | 1-3B | SH | 0 |
| 12IQ0069 | 1-3B | SH | 105 | 121Q0070 | 1-3B | SH | 14 |
| 12IQ0071 | 1-3B | SH | 43 | 12IQ0072 | 1-3B | SH | 268 |
| 12IQ0077 | 2-1B | HC | 86 | 12IQ0073 | 2-1B | HC | 0 |
| 121Q0078 | 2-1B | HC | 0 | 12IQ0074 | 2-1B | HC | 0 |
| 12IQ0079 | 2-1B | HC | 52 | 12IQ0075 | 2-1B | HC | 174 |
| 12IQ0080 | 2-1B | HC | 36 | 12IQ0076 | 2-1B | HC | 0 |
| 12IQ0085 | 2-2B | SH | 89 | 12IQ0081 | 2-2B | SH | 90 |
| 12IQ0086 | 2-2B | SH | 141 | 12IQ0082 | 2-2B | SH | 370 |
| 12IQ0087 | 2-2B | SH | 168 | 12IQ0083 | 2-2B | SH | 0 |
| 12IQ0088 | 2-2B | SH | 190 | 12IQ0084 | 2-2B | SH | 127 |
| 12IQ0093 | 2-3B | HC | 17 | 12IQ0089 | 2-3B | HC | 92 |
| 12IQ0094 | 2-3B | HC | 0 | 12IQ0090 | 2-3B | HC | 1 |
| 12IQ0095 | 2-3B | HC | 36 | 12IQ0091 | 2-3B | HC | 0 |
| 12IQ0096 | 2-3B | HC | 56 | 12IQ0092 | 2-3B | HC | 19 |
|  |  |  | 2545 |  |  |  | 2545 |

Appendix E.

# DNA-Based Population-of-Origin Assignments of Chinook Salmon Smolts Outmigrating Past Chandler Trap at Prosser Dam (Yakima River) in 2013 

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#### Abstract

A population-of-origin assignment procedure was used to estimate the percentages of unknownorigin smolts from each of five stock groups outmigrating past Chandler Trap (Yakima River) from December 2012 - July 2013. Mixture analysis was conducted on a proportional subsample of 1,200 smolts collected during the outmigration at Chandler Trap. Assignment of each individual to a population-of-origin was determined if the posterior probability of the assignment was greater than $90.0 \%$. The largest percentage of outmigrating smolts in the January/February, March, April, and May time strata was from the upper Yakima River stock while the June - July time stratum was dominated by the fall stocks with $70.6 \%$ of the total assignments. Comparison of morphological assessment and genetic assignment as a spring or fall Chinook smolt conducted for all time strata indicated agreement for 1,125/1,180 (95.3\%) of the smolts.


## Introduction

Production and survival of the Yakima River basin spring Chinook stocks (American River, Naches River, and upper Yakima River) are monitored, as part of the Yakima/Klickitat Fishery Project supplementation evaluation program. However, in the lower Yakima River, where the best facilities to collect samples exist, the three spring Chinook stocks are mixed with one another and with the Marion Drain and Yakima River fall Chinook stocks, during downstream juvenile migration. Thus, methodologies for discriminating stocks in an admixture are vital for development of stock-specific estimates. Domestication monitoring plans require discrimination of the three spring Chinook salmon stocks in the basin, and a complete analysis of migration timing and stock abundance for all Chinook requires discrimination of the two fall stocks as well. Accurate assignments of Chinook smolts captured at the Chandler fish passage facility to population-of-origin will allow researchers and managers to estimate production by the three spring Chinook stocks, assess smolt-to-smolt survival of the three spring Chinook stocks, and could be utilized to evaluate stock-specific environmental condition factors.

The methodology used in this study to estimate the population-of-origin for individual fish in a mixture followed a Bayesian approach by Rannala and Mountain (1997). This approach assumes linkage equilibrium among loci and uses the multilocus genotype of an individual to compute the probability of that genotype belonging to a population in the baseline. Others have used the methodology developed by Rannala and Mountain (1997) to provide robust population-of-origin assignments of unknown individuals (Hauser et al. 2006, Taylor and Costello 2006, and Waples and Gaggiotti 2006).

Calculation of population-of-origin for Chinook smolts trapped at Chandler trap throughout the entire outmigration (January through July) was hindered in the first few years of analysis for several reasons: non-representative temporal sampling of the downstream migration, past omission of the Marion Drain fall and lower Yakima River mainstem fall Chinook stocks from the DNA baseline, and by maintenance and other shutdowns of trap operations in December and January in many years. In the analyses of samples from 2004-2010, attempts were made to eliminate the problems present in previous analyses. A new sampling design was initiated to provide a proportional sample of smolts outmigrating past Chandler trap and a larger number of smolts were analyzed. Repeated multi-year samples of all five baseline stocks were used to characterize the potential sources of smolts in the Yakima River basin.

This report presents the population-of-origin assignments for outmigrating smolts collected at the Chandler trap during 2013.

## Materials and Methods

## Collections

There were no collections added to the Yakima River baseline this year. Since 1989, sampling crews from the Yakama Nation and WDFW have collected adult spawning ground tissue samples to be included in the baseline. The tissue samples consisted of dry-mounted scales or fin tissue preserved in 100\% ethanol from five baseline stocks collected across multiple years (American River spring, Naches River spring, upper Yakima River spring, Marion Drain fall, and lower Yakima River fall; Table 1 and Figure 1).

An estimated total of 818,968 smolts passed the lower Yakima River at Chandler from January 2 - July 12, 2013. This estimate was based on expansion of the total number of smolts counted at the Chandler trap $(87,988)$ to account for trap efficiency, etc. Unknown-origin smolts were collected at Prosser Dam (Chandler Trap) following a sampling design that would identify a proportional number of smolt samples that represents the entire smolt outmigration. The following five time strata (January - February, March, April, May, and June - July) were used for analysis. Samples were collected from January 2 - July 13, 2013. These samples were genetically analyzed to get reliable estimates of population proportions. Each day, the total number of smolts at the trap was visually estimated before any processing occurred. If that number was below a predetermined threshold then a "standard" day's sample was taken (e.g. 10 fish). If the number of smolts was above the threshold then a "peak" day's sample was taken (e.g. 30 fish). The threshold for "standard" and "peak" days and the numbers of samples to be taken on each day varied for each of the time strata. These values were determined by analyzing the number of "peak" and "standard" days counted during four years of smolt outmigration monitoring. Based on this sampling design, 2,679 Chinook smolt samples were collected for genetic analysis.

The total estimated numbers of smolts passing the Chandler Trap each day were plotted with the total number of genetic samples that had been collected. A process was then employed to proportionalize the available genetic samples with the daily counts to provide a representative number of smolts that were outmigrating from January - July. A total of 1,200 smolts were identified for analysis.

## DNA Extraction Methods

Genomic DNA was extracted by digesting a small piece of fin tissue (all smolt and some adult baseline collections) or scales (most adult baseline collections) using the nucleospin tissue kits obtained from Macherey-Nagel following the recommended conditions in the user manual. Extracted DNA was eluted with a final volume of $100 \mu \mathrm{~L}$.

## PCR Methods

The polymerase chain reaction mixture contained the following for a $10 \mu \mathrm{~L}$ reaction: approximately 25 ng template DNA, 1X Promega buffer, $1.5 \mathrm{mM} \mathrm{MgCl}_{2}, 200 \mu \mathrm{M}$ each of dATP, dCTP, dGTP, and dTTP, approx. $0.1 \mu \mathrm{M}$ of each oligonucleotide primer, and 0.05 units GoTaq Flexi DNA polymerase (Promega). Amplification was performed using MJ Research PTC-200 and Applied Biosystems 9700 thermocyclers. The thermal profile was as follows: an initial denaturation step of 2 minutes at $94^{\circ} \mathrm{C} ; 40$ cycles of 15 seconds at $94^{\circ} \mathrm{C}, 30$ seconds at $50-60^{\circ} \mathrm{C}$, and 1 minute at $72^{\circ} \mathrm{C}$; plus a final extension step at $72^{\circ} \mathrm{C}$ for 10 minutes, followed by a final indefinite holding step at $10^{\circ} \mathrm{C}$.

Eleven microsatellite DNA loci (Table 2) were amplified via the polymerase chain reaction (PCR) using fluorescently labeled primers (obtained from Applied Biosystems or Integrated DNA Technologies). Loci were combined in multiplexes to increase efficiency and decrease costs.

Data were collected using an AB-3730 Genetic Analyzer. Applied Biosystems GENEMAPPER v.3.7 software was used to collect and analyze the raw data and to determine genotypes at each locus (based on estimated allele sizes in base pairs using an internal size standard). Alleles were binned in GENEMAPPER using the standardized allele sizes established for the Chinook coastwide standardization efforts (Seeb et. al., 2007).

## Population-of-origin Analysis

The program ONCOR (Kalinowski et al. 2008) was used to assign each individual to one of the baseline collections. ONCOR uses conditional maximum likelihood to estimate mixture proportions (Millar 1987) and genotype probabilities are calculated using a partial Bayesian procedure method of Rannala and Mountain (1997). This Rannala and Mountain (1997) method uses the expectation-maximization (EM) algorithm to calculate the population-source probabilities (posterior probabilities) for each sample. All assignments with a posterior probability greater than or equal to $90 \%$ were accepted.

## Comparison of Morphological ID and Genetic Assignment

Smolts were categorized as spring or fall Chinook when they were intercepted at the Chandler Trap based on morphological characteristics. Three morphological features (length, size of the eye, and snout shape) were used to identify smolts as spring or fall (Mark Johnston, Yakama Nation; pers. comm.).

## Results

## Collections

A total of 1,200 unknown Chinook smolts were selected and analyzed from those collected at Chandler Trap. Smolt samples that had data for 10 or more loci were included for analysis. A total of 20 individuals were dropped from statistical analyses.

## Population-of-origin Analysis

The mixture composition estimates for the entire 2013 smolt outmigration indicated that the largest overall percentage of spring smolts was from the upper Yakima River followed by the Naches River and American River in the first four strata. During the migration from January May, the proportion of the upper Yakima River stocks was between 66.7 and $76.2 \%$ while the American River and Naches River spring stocks was between 3.3 and 27.5\%. The proportion of the two fall stocks was between $0.0-22.3 \%$ for the first four time strata and $70.6 \%$ in the June July time stratum (Table 3).

## Comparison of Morphological ID and Genetic Assignment

A comparison of the morphological assessment to genetic assignment was conducted for all five time strata. A total of 39 smolts in January/February, 82 smolts in March, 724 smolts in April, 140 smolts in May, and 195 in the June/July time strata were scored, and therefore included in the analysis. Results for the time strata were as follows: January/February time stratum - all 39 smolts were assigned identically using morphological and genetic methods ( 39 spring); March stratum - 81 out of 82 smolts were assigned identically using morphological and genetic methods ( 81 spring) the one discrepancy was identified as a fall by the genetic analysis and spring with morphological identification; April time stratum - 724 smolts were assigned identically using morphological and genetic methods ( 724 spring); May time stratum - 136 out of 140 smolts were assigned identically using morphological and genetic methods ( 100 spring 36 fall), all four of the discrepancies were identified as a spring by the genetic analysis and fall with morphological identification; June/July time stratum - 145 out of 195 smolts were assigned
identically using morphological and genetic methods (12 spring and 133 fall), 4 discrepancies were assigned as fall by the genetic analyses while morphological identification was spring, the remaining 46 discrepancies were identified as a spring by the genetic analysis and fall with morphological identification.

## Discussion

Collection of smolts at the Chandler Trap in 2013 utilized a sampling design intended to yield a sample that was proportional to the number of smolts passing the Chandler Trap. Sampling a proportional number of smolts was important to determine an accurate percentage of smolts from each stock that were outmigrating from the basin. Developing the sampling strategy for identifying a "standard" versus "peak" day of smolts that were in the trap and applying a sampling goal for those days allowed for a proportional sample. Subsampling the smolts collected for genetic analysis provided a best fit to the actual passage of smolts for a given day.

Monitoring the relative abundances of Chinook smolts in the Yakima River from the three different populations of spring Chinook (upper Yakima River, American River, and Naches River) and the two populations of fall Chinook (Marion Drain and lower Yakima River) requires the ability to estimate population composition of smolts outmigrating past Chandler trap. Because all five Chinook populations are intermingled when they pass Chandler trap, and the vast majority are unmarked and untagged, the only way to determine population-of-origin is by genetic analysis. This method requires that sufficient genetic differences exist among these populations in the Yakima River basin.

A baseline of 19 individual collections from the five populations in the Yakima River basin was used for the population-of-origin assignments of the outmigrating smolts. The baseline collections as a whole had higher genotyping failure compared to the Chandler smolt samples. Scales were taken from carcasses on spawning grounds for most baseline collections; therefore, DNA quality was presumably poorer than the Chandler smolt collection where tissue was collected from live fish. The upper Yakima River tissue collections were also taken from live fish at the hatchery and, therefore, genotyping success was higher for this collection than the other baseline collections.

Assessment of spring or fall smolts by morphological and genetic analysis revealed agreement with 55 individuals being identified differently between the two methods. Identification as a
spring or fall smolt was the same for 1,125 smolts collected during the January - February, March, April, May, and June - July time strata.

The majority of the assignments between January and May were from the three spring stocks. The upper Yakima River spring stock accounted for the highest average percentage (76.2\%) of smolts present in that period. Rank in abundance of the three spring stocks was the same in the three time strata (January-February, March, April, and May) with upper Yakima River spring stock having the most. The June-July time stratum was predominately composed of the fall Chinook stocks, accounting for over $70.6 \%$ of the total number of smolts.

Assessment of DNA Mixture Assignments from 2000-2013
Mixed stock analysis has been conducted on Chandler smolts since 2000 (Young 2004, Kassler et al. 2005, Kassler 2006, Kassler and VonBargen 2007, 2008, 2009 and 2010, Kassler and Peterson 2011, Kassler and Bell 2012, Kassler and Bowman 2013); however the sampling design for samples collected in 2000 - 2003 was not proportionalized during the run. The yearly assignments are therefore not comparable from those years. Beginning in 2004, staff at the Chandler trap utilized a sampling protocol to provide a number of smolts that was relative to the percentage of smolts passing that day. Samples were then subsampled at WDFW to provide a proportional number of samples that would represent the overall passage to be analyzed.

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Table 1. Nineteen Chinook salmon collections assembled into a baseline and used for the analysis of the known-origin and unknown-origin smolts. "*" the 05LU collection from Marion Drain was not used in the baseline, but is listed here as a collection from Marion Drain. The percentage of single locus genotypes missing are shown for each collection.

| Baseline Collections | Collection Code | \# <br> Processed | \# <br> Analyzed | \% Single Locus Genotypes Missing |
| :---: | :---: | :---: | :---: | :---: |
| American River - spring | 89AG | 80 | 77 | 10.4\% |
|  | 91DQ | 102 | 87 | 9.8\% |
|  | 93DO | 18 | 17 | 3.2\% |
|  | 03EH | 100 | 70 | 6.6\% |
|  |  | 300 | 251 | 8.6\% |
| ches River | 89AC | 76 | 74 | 11.4\% |
|  | 89AI | 26 | 22 | 7.0\% |
|  | 93DQ | 50 | 45 | 6.3\% |
|  | 93DR | 32 | 25 | 7.3\% |
|  |  |  |  |  |
| little Naches River - spring | 04BI | 42 | 41 | 2.2\% |
|  | 04EM | 56 | 45 | 9.9\% |
|  |  | 282 | 252 | 7.9\% |
|  |  |  |  |  |
| upper Yakima River - spring | 92DN | 24 | 23 | 5.9\% |
|  | 97DA | 123 | 115 | 3.9\% |
|  | 03GO | 99 | 99 | 1.4\% |
|  |  | 246 | 237 | 3.0\% |
|  |  |  |  |  |
| Marion Drain - fall | 89BX | 100 | 92 | 8.3\% |
|  | 92FQ | 92 | 92 | 5.4\% |
|  | 93DY | 8 | 8 | 8.0\% |
|  | 05LU* | 65 | 47 | 15.3\% |
|  |  | 265 | 239 | 8.6\% |
|  |  |  |  |  |
| lower Yakima River - fall | 90DF | 109 | 104 | 12.6\% |
|  | 93DW | 82 | 80 | 9.8\% |
|  | 98FB | 61 | 50 | 8.7\% |
|  |  | 252 | 234 | 10.8\% |
|  |  |  |  |  |
| Chandler Trap Smolts - 2013 | 13AP | 1,200 | 1,180 | 0.7\% |

Table 2. Microsatellite locus information (number alleles/locus and allele size range) for multiplexed loci used in the analysis of Chinook from five stocks in the Yakima River Basin. Also included are the percent missing genotypes for both the baseline and smolt collections and heterozygosity (observed $\left(\mathrm{H}_{0}\right)$ and expected $\left(\mathrm{H}_{\mathrm{e}}\right)$ ) for each locus.

| expected ( $\mathrm{H}_{\mathrm{e}}$ )) for each locus |  |  |  |  |  |  |  |  |
| :---: | :---: | :---: | :---: | :---: | :---: | :---: | :---: | :---: |
|  |  |  |  |  |  |  | Heterozygosity |  |
| Multiplex | Locus | Annealing temp ${ }^{\circ} \mathrm{C}$ | \# Alleles/ <br> Locus | Allele Size <br> Range (bp) | \% missing genotypes baseline $N=1,166$ | \% missing genotypes smolts $\mathrm{N}=1,180$ | $\mathrm{H}_{0}$ | $\mathrm{H}_{\mathrm{e}}$ |
| Ots-M | Oki-100 ${ }^{\text {a }}$ | 50 | 41 | 164-365 | 11.6\% | 0.5\% | 0.913 | 0.940 |
|  | Ots-201b ${ }^{\text {a }}$ | 50 | 42 | 137-310 | 7.1\% | 0.4\% | 0.916 | 0.936 |
|  | Ots-208b ${ }^{\text {b }}$ | 50 | 52 | 158-342 | 9.7\% | 0.6\% | 0.943 | 0.954 |
|  | Ssa-408 ${ }^{\text {c }}$ | 50 | 32 | 184-308 | 3.5\% | 1.5\% | 0.827 | 0.934 |
| Ots-N | Ogo-2 ${ }^{\text {d }}$ | 60 | 19 | 202-256 | 3.7\% | 0.8\% | 0.756 | 0.854 |
|  | Ssa-197 ${ }^{\text {e }}$ | 60 | 38 | 181-318 | 11.8\% | 0.3\% | 0.915 | 0.940 |
| Ots-O | Ogo-4 ${ }^{\text {d }}$ | 56 | 17 | 132-164 | 15.2\% | 0.5\% | 0.776 | 0.884 |
|  | Ots-213 ${ }^{\text {b }}$ | 56 | 40 | 182-362 | 9.3\% | 0.6\% | 0.908 | 0.940 |
|  | Ots-G474 ${ }^{\text {f }}$ | 56 | 15 | 152-212 | 3.0\% | 1.2\% | 0.507 | 0.697 |
|  |  |  |  |  |  |  |  |  |
| Ots-R | Ots-3M ${ }^{g}$ | 53 | 15 | 128-158 | 2.5\% | 0.3\% | 0.601 | 0.672 |
| Ots-S | Ots-9 ${ }^{g}$ | 60 | 8 | 99-113 | 5.1\% | 1.4\% | 0.668 | 0.709 |
| ${ }^{\mathrm{a}}=$ Unpublished |  |  |  |  |  |  |  |  |
| ${ }^{\mathrm{b}}=$ Greig et al. 2003 |  |  |  |  |  |  |  |  |
| ${ }^{\text {c }}$ = Cairney et al. 2000 |  |  |  |  |  |  |  |  |
| d $=$ Olsen et al. 1998 |  |  |  |  |  |  |  |  |
| ${ }^{\text {e }}=$ Oreilly et al. 1996 |  |  |  |  |  |  |  |  |
| ${ }^{\mathrm{f}}=$ Williamson et al. 2002 |  |  |  |  |  |  |  |  |
| ${ }^{\mathrm{g}}=$ Banks et al. 1999 |  |  |  |  |  |  |  |  |

Table 3. Stock-of-origin assignments for five stocks of Chinook in the Yakima River Basin using ONCOR.


Figure 1. Geographic location of the Chandler trap on the Yakima River, Washington and the primary streams in the basin.


