# Comprehensive Evaluation of Middle Columbia River Public Utility District's Summer Chinook Salmon Hatchery Programs 

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## Executive Summary

The three Public Utility Districts (PUD) of the middle Columbia River strive to achieve no net impact of Salmon and steelhead as a result of construction and operation of five Columbia River dams. One of the three components the PUDs use to achieve no net impact is the production of hatchery fish to replace juvenile fish lost through the project areas. A comprehensive monitoring and evaluation plan is implemented to determine if the performance of the hatchery programs is achieving the goals described in the plan (Hillman et al. 2019). This report is a synthesis of the analyses and results from data collected for Grant, Chelan, and Douglas County PUD's spring Chinook Salmon hatchery programs through 2018. Other covered species (e.g., fall and summer Chinook, Sockeye Salmon, and steelhead) are presented in other reports. Authorship, titles, and abstracts of each of the report chapters are presented below.

1) Pearsons, T. N, T. W. Hillman, and C. C. Willard. The Effects of hatchery supplementation on the abundance and productivity of adult summer Chinook Salmon in the Upper Columbia Basin.

Managers frequently select hatchery supplementation as a tool to increase abundance and harvest of Chinook Salmon (Oncorhynchus tshawytscha), but supplementation can pose risks to naturally spawning populations. We evaluated the effects on naturally spawning summer Chinook Salmon populations of two adaptively managed conservation/harvest augmentation programs operating from 1992-2018 in the Wenatchee and Methow rivers, tributaries to the Upper Columbia River. Comparisons between supplemented populations and a reference (unsupplemented) population were performed using a Before-After-Control-Impact Paired (BACIP) design to evaluate whether supplementation changed adult productivity, density-adjusted adult productivity, and the abundance of total spawners, natural-origin adult recruits (NOR), and natural-origin spawners (NOS). Adults collected and spawned in hatcheries produced considerably more adult recruits per spawner than those that spawned in the natural environment. The BACIP contrasts for total spawner abundance and NOS in the Wenatchee population were significantly negative ( $\mathrm{P}<0.000$ ), which was contrary to management objectives, and significantly positive for productivity $(\mathrm{P}=0.021)$ but not when adjusted for density $(\mathrm{P}=0.091)$. Consistent with management objectives, all BACIP contrasts were positive in the Methow population but only NOR was significantly positive ( $\mathrm{P}=0.017$ ). Mean total spawners, NOS, and NOR increased in both the Methow and reference populations between the before and after periods but the Methow population increased more than did the reference population during that period. The statistical power to detect differences of BACIP contrasts stabilized 10-20 years after supplementation, which was within the timeframe of this evaluation (24 years). Summer Chinook Salmon supplementation in the upper Columbia Basin produced mixed results relative to an un-supplemented reference population outside the Upper Columbia Basin, which may have been a function of density-dependence, but both programs contributed substantially to harvest.
2) Buchanan, R. A., R. L. Townsend, and G. Mackey. Investigations into association between proportion of hatchery spawners and juvenile productivity for spring Chinook, summer Chinook, and summer steelhead in the Wenatchee and Methow river basins.

Conservation hatchery programs implemented in the Wenatchee and Methow river basins are intended to increase the abundance of the target populations. The strategy of the hatchery programs is to return hatchery adults that increase the spawning population, resulting in an increase in the number of offspring that will maintain and recover naturally reproducing populations. However, there is a concern that the hatchery programs may lower the juvenile productivity in supplemented watersheds. The populations are managed under the concept of Proportionate Natural Influence (PNI), a framework that uses genetic modeling to estimate risk of hatchery introgression in natural populations and prescribes management benchmarks intended to allow hatchery programs to operate within an acceptable risk level to the natural population. One of the metrics in the PNI framework is the proportion of hatchery origin spawners ( pHOS ). Our investigation of whether higher pHOS may result in lowered juvenile productivity was largely inconclusive. For most populations studied, no evidence of an effect of pHOS was observed. For the single population that demonstrated a possible negative effect of pHOS on juvenile productivity (Twisp River Summer Steelhead), the evidence was weak: when density dependence was accounted for, the perceived negative association between pHOS and juveniles per redd was no longer observed. Additionally, the significant result was observed only when a possible outlier in emigrant counts was omitted, which may not be justified depending on the cause of the unusual emigrant count for that brood year. Thus, even when some evidence of a negative effect of pHOS was found, the evidence was particularly weak. For all three stockrecruitment models considered, there was often high uncertainty in model parameter estimates even when the models could be fit to the data. This was indicated by wide confidence intervals that often included maximum smolt or emigrant values far beyond the range of observed data, and by the high correlation in model parameter estimates. High correlation lowers the ability to distinguish between different model fits and increases uncertainty in the results; it arises from lack of contrast in the data or violation of modeling assumptions. Given all the complicating factors identified here, only the most extreme effect of pHOS on juvenile productivity could have been detected. Thus, we caution against concluding that such an effect is truly absent. We conducted a power analysis based on the Chiwawa spring Chinook data-the most robust dataset in this study. Simulated power analyses revealed that studies shorter than 70 years will have low power to detect an effect on recruitment for all but the strongest effect sizes of pHOS . Future data-collection efforts could be substantially improved by increasing the contrast in pHOS levels and including suitable in-basin unsupplemented references. More specifically, a planned experiment using deliberately chosen pHOS levels could be implemented in order to achieve the necessary contrast and remove confounding analyses with temporal and age-structured processes. Appropriate power analyses must be performed on the study design prior to commencing work to ensure that a result may be obtained within an acceptable time frame.
3) Mackey, G., T. N. Pearsons, and T. W. Hillman. The effect of hatchery programs on Proportionate Natural Influence (PNI) in the upper Columbia Basin.

Hatchery programs and natural populations in the Wenatchee and Methow subbasins were managed under the Proportionate Natural Influence (PNI) strategy, whereby gene flow between the hatchery and natural populations was manipulated to achieve greater overall genetic influence from the natural population as opposed to the hatchery population. The target PNI value was 0.67 . Steelhead programs in the Wenatchee and Methow had PNI averages or medians below 0.67. The new management regime for steelhead in the Methow was not initiated until 2017. Therefore, data in this report do not reflect the future management of the steelhead in the Methow subbasin for gene flow. Spring Chinook Salmon PNI in the Wenatchee Subbasin was below 0.67 for all populations under the contemporary management strategy but exceeded 0.50 in all cases. Methow subbasin spring Chinook PNI was 0.29 , and has not increased appreciably in the two years of adult returns following reduction in program sizes in 2013. Wenatchee and Methow summer Chinook both had PNI means that exceeded 0.67. Both of these programs appear successful in meeting PNI objectives with the Wenatchee PNI an impressive 0.87. The results of this analysis indicate that the Spring Chinook and steelhead programs require adjustment to meet the PNI targets. In some cases, the PNI values may improve as new management strategies mature. The Methow steelhead and particularly Methow spring Chinook may require substantial management changes in order to achieve the gene-flow objectives. In general, PNI targets were not met for small populations (spring Chinook Salmon and steelhead) but were for large populations (summer Chinook Salmon).
4) Graf, P. J., C C. Willard, T. W. Hillman, T. N. Pearsons, T. H. Kahler, and G. Mackey. Adult migration timing, spawn timing, and spawning distribution of spring Chinook Salmon and summer Chinook Salmon in the Wenatchee and Methow basins.

The migration timing, spawn timing, and spatial spawning distribution of hatchery- and naturalorigin salmon in the natural environment can be important metrics in the evaluation of integrated hatchery programs. The timing of migration and spawning of hatchery- and natural-origin spring Chinook Salmon were generally similar in the Wenatchee and Methow sub-basins. Although the difference in arrival timing was small and not statistically significant, the visual observation and PIT-tag data at Tumwater and Wells dams suggests a tendency for hatchery-origin spring Chinook Salmon to arrive later than natural-origin fish. Differences in migration timing between spring Chinook Salmon populations in the Wenatchee versus Methow sub-basins were observed and may result in differences in survival at the adult life-stage. With summer Chinook Salmon, hatchery-origin fish in the Wenatchee sub-basin consistently passed Dryden Dam later than their natural-origin counterparts whereas the migration timing for Methow and Okanogan sub-basin hatchery-origin fish was a near match with natural-origin fish at Wells Dam. Summer Chinook Salmon spawn timing in both Wenatchee and Methow hatchery-origin fish was later than natural-origin fish, with Methow-origin fish having a larger average difference. For spatial distribution of spawning spring Chinook Salmon, differences between hatchery- and naturalorigin spawner distribution across the historical survey reaches were observed in all programs except the White River program. In general, hatchery-origin females spawned lower in the watershed. However, the proportion of hatchery-origin fish was high (i.e., greater than $50 \%$ ) in
the majority of survey reaches ( 33 out of 51), particularly in those reaches that were the primary spawning areas by natural-origin fish. The proportion of hatchery-origin spring Chinook Salmon exceeded $30 \%$ in 48 of the 51 survey reaches. The distributions of spawning Wenatchee and Methow hatchery-origin summer Chinook Salmon also differed from the spawner distributions of the natural-origin populations, with hatchery-origin females more often spawning lower in the watershed. This difference in spawning distribution was consistent with management objectives. The proportion of hatchery-origin summer Chinook Salmon was greater than $30 \%$ in 6 of the 17 survey reaches.
5) Pearsons, T. N. and R. R. O'Connor. Stray rates of natural-origin Chinook Salmon and steelhead in the upper Columbia Watershed.

Despite the importance of straying in understanding the ecology of salmon and steelhead, most of what is known about salmon and steelhead straying comes from tagged hatchery fish. We provide donor estimates of natural-origin spring, summer, and fall Chinook Salmon Oncorhynchus tshawytscha and steelhead Oncorhynchus mykiss straying at three spatial scales in the upper Columbia watershed using Passive Integrated Transponder (PIT) tags. A total of 823,770 natural-origin spring, summer, and fall Chinook Salmon and summer steelhead were PIT-tagged as juveniles in the Wenatchee, Entiat, Methow, and Okanogan River subbasins and tributaries and the upper Columbia River between 2002 and 2017. Anadromous adults with PIT tags were detected at a variety of antenna arrays in the Columbia River Basin between 2004 and $2018(\mathrm{n}=2,611)$. Mean donor stray rates of each population were less than $1 \%$ at the basin scale (range $0.0 \%-0.7 \%$ ), less than $10 \%$ at the subbasin scale (range $0.0 \%-9.8 \%$ ) and less than $15 \%$ at the tributary scale (range $0.0 \%-14.3 \%$ ). Many of the populations (11 of 28) that were evaluated across all spatial scales did not have any strays detected, and the mean of means of all species stray rates at all spatial scales was generally less than 5\% (range 0.2\%-4.0\%). Chinook Salmon and steelhead strayed at similar rates when originating from the same subbasins and tributaries. Most straying occurred in an upstream direction at the subbasin (84\%) and tributary scales ( $94 \%$ ). Variation in stray rates was most consistently associated with spatial scale and location and was less than $15 \%$ for all species at all spatial scales.
6) Pearsons, T. N. and R. R. O’Connor. Comparisons of donor stray percentages between hatchery- and natural-origin Chinook Salmon and steelhead in the upper Columbia Watershed.

Artificial propagation of salmon Oncorhynchus spp. and steelhead $O$. mykiss is a common strategy that is used to achieve conservation and harvest goals. However, unintended effects of artificial propagation, such as high donor stray percentages, can reduce the number of adults that return to target areas and also contribute spawners to different populations where they are not desired. Until recently, it was difficult to assess if hatchery-origin fish stray rates were atypical because few estimates of stray rates of natural-origin fish were available. We used last PIT-tag detections to estimate and compare donor stray percentages of hatchery-origin and natural-origin Chinook Salmon $O$. tshawytscha and steelhead in the upper Columbia River watershed between

2002-2018. Donor stray percentages of hatchery-origin spring, summer, and fall Chinook Salmon and steelhead were $<0.3 \%$ at the upper-Columbia basin scale and generally not higher than natural-origin donor stray percentages at larger spatial scales but were higher (up to 62\%) at smaller spatial scales. Returning hatchery-origin Chinook Salmon and steelhead generally strayed in an upstream direction and the proportions of fish that strayed upstream were not significantly higher than natural-origin fish. Juvenile spring Chinook Salmon that were moved 14 to 389 river kilometers from centralized hatcheries to tributaries for overwintering or final acclimation, strayed at a much higher rate than those that completed their incubation, rearing, and acclimation at a single location. In contrast, steelhead that were moved for acclimation, including direct releases from trucks, did not stray at higher rates than those that completed their incubation, rearing, and acclimation at a single location. Other adaptive management actions that were implemented to reduce straying produced mixed results. A variety of approaches can be considered to reduce undesirable production of strays, but most of them involve difficult trade-offs.
7) Pearsons, T. N., and M. D. Miller. Stray compositions of hatchery-origin Chinook Salmon Oncorhynchus tshawytscha and steelhead $O$. mykiss in recipient natural populations of the upper Columbia Watershed.

One of the biggest concerns of operating hatchery Salmon and steelhead programs is high straying of returning adults into non-target populations and the possible homogenization of genetic diversity among populations caused by spawning of stray fish. The composition of hatchery-origin stray Chinook Salmon Oncorhynchus tshawytscha and steelhead O. mykiss relative to the natural spawning populations, termed recipient population stray rate, was evaluated in the Upper Columbia Basin. Chinook Salmon carcasses were collected from 19992018 in spawning areas shortly after spawning and carcasses were examined to determine origin. Adipose fin clips and coded-wire-tags were used to distinguish non-target hatchery, target hatchery, and natural-origin fish; coded-wire-tags were read in the lab to determine the origin of hatchery-origin fish. Steelhead strays and spawning escapement were evaluated using passiveintegrated transponder (PIT) tags between 2013-2018. The recipient population stray rates ranged between $0.02-87.35 \%$ and increased with decreasing spatial scale. Recipient stray rates of all taxa at the basin scale were $<3 \%$, and summer Chinook and fall Chinook salmon were $<0.5 \%$. Stray rates in subbasins for all taxa ranged between $0.07-33.04 \%$; spring and summer Chinook Salmon exceeded 5\% in some 10 year periods in the Entiat and Methow subbasins, but stray rates for all Chinook Salmon were $<5 \%$ in the Wenatchee, Okanogan, and Hanford Reach for all periods. All steelhead stray rates exceeded 5\% except for those in the Wenatchee subbasin. Stray rates of spring Chinook Salmon in tributaries (the only taxa that met the tributary criteria) ranged between $0.61 \%-87.35 \%$ and only the Chiwawa, Icicle, and Twisp rivers were consistently below $10 \%$; the Chiwawa River was consistently below $5 \%$. In cases where recipient stray management targets were exceeded, some were the result of single hatchery contributions, but others were the result of cumulative contributions from multiple hatcheries. Options to achieve recipient stray management targets include reducing donor stray rates,
reducing hatchery program size, removing hatchery-origin adults prior to spawning in the natural environment, and increasing the natural-origin population. It is likely that balancing trade-offs among hatchery program size and recipient population stray rate will be necessary in order to achieve management targets in some locations.
8) McKinney, G., S. Brown, A. Louden, M. P. Small, T. R. Seamons, C. C. Willard, T. N. Pearsons, T. H. Kahler, and G. Mackey. Examining the genetic structure of upper Columbia Summer/Fall Chinook Salmon and evaluating the effects of the supplementation program.

We examined baseline (1982-1994) and contemporary (2017-2018) summer and fall Chinook Salmon (Oncorhynchus tshawytscha) from the Upper Columbia River Watershed to determine if hatchery supplementation programs have had any impacts on the genetic diversity and structure of these populations. Baseline collections included both hatchery- and natural- origin samples where available. Contemporary collections exclusively consisted of samples collected at broodstock collection facilities; their origin (hatchery or natural) was only sometimes known. Summer Chinook Salmon populations with paired baseline and contemporary samples included the Methow River, the Wenatchee River, and the Okanogan River. Populations with only contemporary samples included Chelan Falls, Entiat National Fish Hatchery, and Wells Fish Hatchery. Fall Chinook Salmon were represented by collections from the Hanford Reach spawning grounds and Priest Rapids Hatchery. Measures of genetic diversity (allelic richness, heterozygosity, linkage disequilibrium, and effective number of breeders) showed little differentiation among baseline and contemporary populations for either summer or fall Chinook, suggesting that hatchery programs have not led to a decrease in genetic diversity. There was a general pattern where $F_{\mathrm{ST}}$ was higher among baseline than contemporary collections suggesting that genetic drift and homogenization among stocks has occurred over time. Despite these patterns, pairwise comparisons of $F_{\text {ST }}$ were generally statistically non-significant both for baseline and contemporary collections. Similar to previous evaluations, there appears to be little evidence for neutral genetic divergence between contemporary hatchery programs in the upper Columbia watershed and baseline samples collected in the late 1980s and early 1990s. The large population sizes of summer and fall Chinook Salmon relative to the hatchery program sizes in the upper Columbia basin, low recipient population stray rates in natural populations, and the management strategies that were implemented to reduce genetic risk all likely contribute to the lack of neutral genetic change. This evaluation did face two limitations: first, we were not able to evaluate potential differentiation among contemporary hatchery and natural origin individuals due to lack of data on individual origin; second, we were not able to evaluate potential shifts in adaptive genetic diversity using genetic techniques and it is possible for adaptive genetic diversity (i.e., run-timing, age at maturity) to change in response to selection (i.e., domestication) while neutral genetic diversity remains the same. While adaptive genetic variation was not directly monitored, phenotypic metrics measured as part of other portions of the monitoring plan can serve as a proxy for adaptive genetic variation.
9) Graf, P. J., T. W. Hillman, T. H. Kahler, C. C. Willard, and T. N. Pearsons. The effects of hatchery supplementation on size and age at maturity and fecundity of spring and summer Chinook Salmon in the upper Columbia Basin.

A common objective of conservation hatchery programs is to produce fish with phenotypic traits similar to individuals from the natural population. We evaluated the size and age at maturity, and fecundity of five spring Chinook Salmon and three summer Chinook Salmon hatchery programs. Comparisons to natural-origin fish from the targeted populations over multiple generations and during two periods of hatchery production releases (high and low number of smolts) were made for applicable programs. Generally, the hatchery-populations were composed of more, younger fish and fewer older fish, but the predominate age at maturity was similar between hatchery- and natural-origin populations. The size at maturity of returning adults was significantly affected by age and sex, and in some cases origin and period. However, when matched by age and sex, the differences in size at maturity by origin and period that were statistically significant were minor and likely of little biological relevance. For all populations, the fecundity metrics were significantly affected by fish size and weight, but fecundity differed between origins in only two populations. For all metrics, the high/low program release numbers did not substantially affect these phenotypic traits. Future analyses of these metrics should consider changes over time and consider trends between hatchery-origin and natural-origin fish. Declines in age and size at maturity are a range-wide concern for Pacific salmonids and therefore a temporal analysis of the dataset generated for the monitoring of these hatchery programs would benefit the region.
10) Mackey, G., and T. W. Hillman. Numbers and sizes of fish released from hatcheries in the upper Columbia Basin.

Chelan, Douglas, and Grant PUDs produce salmon and steelhead for mitigation under various agreements including the Rock Island Habitat Conservation Plan (HCP), the Rocky Reach HCP, the Wells HCP, and the Priest Rapids Settlement Agreement and associated Biological Opinion. The various hatchery programs are monitored for the number of fish released, as well as metrics for fish size at release and condition. The programs include Summer Chinook Salmon, Spring Chinook Salmon, and summer steelhead produced and released at a variety of facilitates in the Wenatchee River Subbasin, Methow River Subbasin, and Columbia River. Programs generally met or exceeded the release-number targets. Specific programs did not meet the release-number targets in all years, but all programs met or exceeded targets in the majority of years, with the exception of the White River spring Chinook Salmon captive broodstock program. This program was in development for all of its history and tested numerous fish culture and release strategies. Therefore, the program was not able to meet targets that remained consistent across years. Most programs met or were close to meeting the CV target on average. Meeting the CV targets for steelhead programs was more difficult than for the Chinook Salmon programs. Meeting fish-perpound (fpp) targets was more difficult in Spring Chinook Salmon and conservation steelhead programs, potentially because these programs use natural-origin broodstock. Spring Chinook Salmon exhibited near-isometric growth. Steelhead exhibited negative allometric growth, as did Wells and Methow summer Chinook Salmon. However, Wenatchee and Chelan Falls summer

Chinook Salmon exhibited isometric or positive allometric growth. Condition factors across all programs were very close to or exceeded 1 . Trade-offs between post-release survival and age at maturation influence hatchery rearing strategies. Large Chinook Salmon generally survive better than small fish, but they also are more prone to mature at younger ages. Hatchery programs that use natural-origin fish for broodstock may experience higher levels of precocial maturation than those that use hatchery-origin fish for broodstock. In addition, larger fish also pose ecological risks to other fish through mechanisms such as competition or predation. In general, the length targets, and some of the metrics that used a single value (e.g., length and weight) associated with a range of fpp targets, were not useful. Adaptation of targets may occur to achieve a better suite of benefits among the many trade-offs involved in growing fish to a target size or weight.
11) O'Connor, R. R., and T. N. Pearsons. Harvest of Chinook Salmon and steelhead originating from Upper Columbia River hatchery programs.

The objective of this evaluation was to determine if a diversity of upper Columbia Basin Chinook Salmon and steelhead hatchery programs contributed to harvest. More specifically, we were interested in evaluating whether harvest rates were consistent with management objectives and where fish were harvested. Harvest rates were lowest on endangered spring Chinook Salmon with annual brood year means of $5-6 \%$ for Methow, Chewuch, and Twisp spawning aggregates (annual range 0 to $59 \%$ ) and $26 \%$ for the Chiwawa spawning aggregate (annual range 0 to $95 \%$ ). The percent of the population harvested was not correlated with spawning escapement ( $\mathrm{P}>0.05$ ) and the total number of fish harvested was correlated with spawning escapement ( $\mathrm{P}<0.05$ ) in the Chiwawa and Twisp rivers but not in the Methow or Chewuch rivers. Most harvest of spring Chinook Salmon occurred in freshwater. Harvest rates were much higher for the more abundant summer and fall Chinook Salmon programs with annual brood year averages around $53-75 \%$ and annual ranges of 14 to $91 \%$. Percent harvest increased with increasing spawning escapement for summer Chinook in the Methow ( $\mathrm{P}=0.01$ ) and Okanogan ( $\mathrm{P}=0.0002$ ) rivers but not for summer Chinook in the Wenatchee River $(\mathrm{P}=0.49)$, Chelan Falls/Turtle Rock program ( $\mathrm{P}=0.43$ ), and Hanford Reach fall Chinook ( $\mathrm{P}=0.28$ ). The total number fish harvested was not correlated with spawning escapement ( $\mathrm{P}>0.05$ ) for the Wenatchee River, Wells subyearling, Methow River, or Okanogan River programs, but significant correlations were detected $(\mathrm{P}<0.05)$ for the Chelan Falls/Turtle Rock yearling and Wells yearling programs and for fall Chinook Salmon from Priest Rapids Hatchery. Most of the harvest of summer Chinook Salmon occurred in the ocean and harvest of fall Chinook Salmon occurred evenly between freshwater and the ocean. Harvest rates averaged $16 \%$ (range $0-54 \%$ ) for threatened hatchery-origin steelhead and less than 5\% (range 0 to $4 \%$ ) for natural-origin steelhead. The percent of steelhead harvested increased with increasing escapement in the Okanogan River ( $\mathrm{P}=0.006$ ) but was not significantly correlated in the Methow ( $\mathrm{P}=0.29$ ) and Wenatchee rivers ( $\mathrm{P}=0.85$ ). Total harvest of hatchery steelhead was not significantly correlated with spawning escapement in the Methow or Wenatchee rivers ( $\mathrm{P}>0.05$ ) but was correlated in the Okanogan River $(\mathrm{P}=0.006)$. Every hatchery program that was evaluated contributed to harvest and sometimes substantially. The magnitude of harvest generally corresponded to the
status of the population: the lowest harvest occurred on the most imperiled stocks and the highest harvest occurred on the healthiest stocks. However, harvest sometimes hindered meeting broodstock collection goals and harvest management of endangered or threatened species could impede conservation objectives and might be improved by tailoring harvest to abundance, weak stocks, and weak broodyears.

Please read the full chapters for more detail about each of the topics in the abstracts presented above. All data in this report should be considered preliminary until published in a peer-reviewed journal.

## Introduction

The three Public Utility Districts (PUD) of the middle Columbia River strive to achieve no net impact of Salmon and steelhead as a result of construction and operation of five Columbia River dams. This report describes one of the main ways the PUDs achieve no net impact; the production of hatchery fish to replace those lost through the project areas. A comprehensive monitoring plan is implemented to determine the performance of the hatchery programs at achieving their goals (Hillman et al. 2019). This report is a synthesis of the data collected for Grant, Chelan, and Douglas PUD's ssummer Chinook Salmon hatchery programs through 2018. Other covered species (e.g., spring Chinook, fall Chinook, Sockeye Salmon, and steelhead) are presented in other reports.

The Douglas and Chelan PUDs' Habitat Conservation Plans (HCPs), Grant PUD's Settlement Agreement, and the 2008 NMFS Biological Opinion (Biop) for Grant PUD (hereafter referred to collectively as the Agreements) specify certain reporting dates or intervals for hatchery monitoring and evaluation (M\&E). The Endangered Species Act (ESA) incidental take permits and the Monitoring and Evaluation Plan for PUD Hatchery Programs (Hillman et al. 2019) also have reporting requirements. These reporting date requirements were designed to provide timely information to operators and managers and fulfill permitting requirements. Additionally, the reports are used to inform other activities such as updating M\&E plans, recalculation of hatchery production, evaluation of meeting M\&E objectives, status of meeting permit requirements, and adaptive management actions. Past reporting timing has not necessarily met the intent of the Agreements, and has not been orchestrated to align with the various actions that the Hatchery Committees and NMFS require. Subsequently, we have designed a reporting schedule that is consistent with the Agreements, meets reporting requirements under the M\&E Plan, meets ESA Section 10 permit requirements, and optimizes the sequence of reporting and the actions that rely on M\&E information.

Three levels of M\&E reporting have been and will be implemented (Table 1). These reports are consistent with past reporting and the M\&E Plan, but have been restructured to streamline transfer of information and meet the requirements of the Agreements.

Table 1. Monitoring and evaluation report types, frequency, content and function.

| Report <br> type | Frequency | Content | Function |
| :--- | :--- | :--- | :--- |
| Data | Annual | Cumulative description of data (raw <br> and derived) and field methods. <br> Basic statics reported. | Informs annual M\&E <br> implementation plans |
| Statistical | 5 year | Presentation of statistical analyses <br> and description of statistical methods. <br> Addressed in the Program Review <br> when the two would occur in the <br> same year. | Informs 5 year M\&E plan <br> and provides in depth <br> data analysis |
| Program | 10 year | Integrates and interprets information <br> from data and statistical reports and <br> also includes integration from other | Informs recalculation and <br> adaptive management. <br> Determines if programs <br> are meeting objectives. |
|  |  | programs and studies. Written in <br> scientific manuscript format. Fulfills <br> HCP "Program Review" |  |
| requirements. Addresses Statistical |  |  |  |
| Report requirements. |  |  |  |

The Data Report will be produced annually and will provide data collected in the most recent field year. The report will provide tables of cumulative data, including the most recently collected, and provide summary statistics where appropriate (e.g., mean, standard deviation, etc.). The report will provide a concise description of the field methods that could be used in a scientific publication and describe deviations from previous sampling, standard field practices or sampling plans. This report will provide up to date information for managers and operators, fulfill incidental take reporting requirements, and inform annual adjustments to the implementation of the M\&E plan.

The Statistical Report will be produced every ten years on the five year intervals between the Program Review (see below). The report will provide a concise description of the analytical methods used (e.g., similar to a scientific journal article) and the results of the statistical analyses for each objective as described in the M\&E plan. The report will also provide the assumptions of the statistical analyses and note any deviations in expected performance of a given analysis (e.g., issues related to normality, dependency, non-constant variance; etc.). The report is not intended to provide interpretation of the results, but will provide the outcomes of the statistical tests. This will provide managers and operators a periodic update of the performance of the hatchery programs.

The Program Review, also known as the Comprehensive Report, will be produced every ten years and will meet the Program Review as described in the HCPs (Section 8.8 of the Wells HCP, Section 8.7 of the Rocky Reach/Rock Island HCPs) and will address the information reported in the Statistical Report. The report will provide the results of any natural population/hatchery interaction studies (as needed), and determine if the hatchery programs are operating consistent with the goals as outlined in the relevant M\&E Plan. The review will determine if hatchery program goals and objectives, as defined in the Hatchery Plan (HCPs Section 8), Section 10 permits, as further defined in the HCPs, have been met or sufficient progress is being made toward their achievement; and determine if hatchery production objectives are being achieved.

The M\&E reporting schedule is designed to be consistent with the Agreements. However, it also has been designed to provide a logical sequence of information based on significant milestones in the HCPs as well as consistency with Grant PUDs settlement agreement and NMFS Biological Opinion. Reporting was designed to provide the Program Review (ten year interval) prior to recalculation in order to have the most up to date data vetted and organized prior to recalculation. The Statistical Report will be produced every ten years. On the five year intervals between the ten year intervals, the Statistical Report material will be addressed in the Program Review. The Data Report will be produced annually. The PUDs also require advanced knowledge of M\&E and reporting requirements to facilitate timely contracting. The Agreements terminate in 2052.

## Summary

Annual reports have been conducted for decades (e.g., Hillman et al. 2020, Snow et al. 2020, Richards and Pearsons 2019), but there has only been one comprehensive analysis of PUD programs and this did not integrate data from all of the PUD programs nor with relevant literature from other locations (Hillman et al. 2012; Murdoch et al. 2012). Furthermore, many of the data sets were not mature enough to make robust conclusions. The current evaluation attempts to improve upon previous evaluations by: 1) including more data, 2) improving analytical techniques, 3 ) including all PUD programs together, and 4) integration of findings relative to other published work.

We attempted to generate relevant chapter topics that encompassed all of the monitoring and evaluation plan objectives (Hillman et al. 2019). The objectives of the M\&E plan and the associated chapter numbers are in Table 2. Finally, we conducted more analyses that were identified in the plan in order to provide a more comprehensive evaluation of the programs.

Table 2. Hatchery monitoring and evaluation plan objectives contained in Hillman et al. 2019 and the associated chapter numbers in this report that address them.

| Objective | Objective Description | Report Chapter |
| :---: | :---: | :---: |
| 1 | Determine if conservation programs have increased the number of naturally spawning and naturally produced adults of the target population and if the program has reduced the natural replacement rate (NRR) of the supplemented population. | 1 |
| 2 | Determine if the proportion of hatchery fish on the spawning grounds affects the freshwater productivity of supplemented stocks. | 2 |
| 3 | Determine if the hatchery adult-to-adult survival (i.e., hatchery replacement rate, HRR) is greater than the natural adult-to-adult survival (i.e., natural replacement rate, NRR) and the target hatchery survival rate. | 1 |
| 4 | Determine if the proportion of hatchery-origin spawners ( pHOS or PNI) is meeting the management target. | 3 |
| 5 | Determine if the run timing, spawn timing, and spawning distribution of the hatchery component is similar to the natural component of the target population or is meeting program-specific objectives. | 4 |
| 6 | Determine if the stray rate of hatchery fish is below the acceptable levels to maintain genetic variation among stocks. | 5,6.7 |
| 7 | Determine if genetic diversity, population structure, and effective population size have changed in natural spawning populations as a result of the hatchery program. | 8 |
| 8 | Determine if hatchery programs have caused changes in phenotypic characteristics of natural populations. | 9 |
| 9 | Determine if hatchery fish were released at the programmed size and number. | 10 |
| 10 | Determine if appropriate harvest rates have been applied to conservation, safety-net, and segregated harvest augmentation programs to meet the HCP/SSSA goal of providing harvest opportunities while also contributing to population management and minimizing risk to natural populations. | 11 |

One of the topics identified for the hatchery program review has already been published in a peer-review journal, the highest standard of the profession. The citation of the publication is provided below.

Pearsons, T. N. and R. R. O’Connor. 2020. Stray rates of natural-origin Chinook Salmon and Steelhead in the Upper Columbia Watershed. Transactions of the American Fisheries Society 149:147-158. DOI: 10.1002/tafs. 10220

There has been extensive review and adaptation of both the PUD hatchery and monitoring and evaluation programs. The PUD hatchery programs have been reviewed by the PUD Hatchery Committees and the Hatchery Science and Review Group (HSRG). In addition, the PUD hatchery monitoring and evaluation plan has been reviewed by a number of different groups including the PUD Hatchery Committees, the Independent Scientific Advisory Board (ISAB) in 2018, and an expert genetics panel that was assembled in 2019. These reviews and associated adaptations have resulted in high quality hatchery and monitoring and evaluation programs.

The hatchery programs have undergone many operational and in some cases facility changes during the time of monitoring and implementation. This poses challenges to evaluate the many changes that have occurred. For example, hatchery programs were resized in 2013 and will be resized every 10 years based upon mitigation requirements and hatchery programs were revised consistent with hatchery reform principles such as PNI management. In many cases, the programs were not held constant for enough years to statistically evaluate changes such as those associated with resizing the hatchery programs that began with smolt releases in 2014. Therefore, we evaluated the programs as the outcome of adaptive management to achieve longterm program goals, which generally did not change. This is appropriate because, the programs are continually evolving in attempts to improve the probability of achieving overarching management goals.

The committees had an extended period of time to review the chapters contained in this report and their comments are presented as a separate document to this report. The authors responses to those comments are also presented in that document.

This report will help inform a future committee authored summary report. The summary report will include committee approved recommendations that will inform a revision of the monitoring and evaluation plan as well as program operation. The recommendations that are provided in this report are those of the authors and do not necessarily reflect the views of the hatchery committees.

## Acknowledgments

Thanks are due to the many people who have contributed to the collection, analysis, and presentation of the data contained in this report. Specific contributions are mentioned at the end of each chapter of this report. We also thank the PUDs and other funding entities such as the Bonneville Power Administration who have invested in understanding the fishes in the Upper Columbia Region. Finally, we thank the HCP Hatchery Committees and the PRCC Hatchery Subcommittee for their input and review of the hatchery and M\&E programs.

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# The Effects of Hatchery Supplementation on the Abundance and Productivity of Adult Summer Chinook Salmon in the Upper Columbia Basin 

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#### Abstract

Managers frequently select hatchery supplementation as a tool to increase abundance and harvest of Chinook Salmon (Oncorhynchus tshawytscha), but supplementation can pose risks to naturally spawning populations. We evaluated the effects on naturally spawning summer Chinook Salmon populations of two adaptively managed conservation/harvest augmentation programs operating from 1992-2018 in the Wenatchee and Methow rivers, tributaries to the Upper Columbia River. Comparisons between supplemented populations and a reference (unsupplemented) population were performed using a Before-After-Control-Impact Paired (BACIP) design to evaluate whether supplementation changed adult productivity, density-adjusted adult productivity, and the abundance of total spawners, natural-origin adult recruits (NOR), and natural-origin spawners (NOS). Adults collected and spawned in hatcheries produced considerably more adult recruits per spawner than those that spawned in the natural environment. The BACIP contrasts for total spawner abundance and NOS in the Wenatchee population were significantly negative ( $\mathrm{P}<0.000$ ), which was contrary to management objectives, and significantly positive for productivity $(\mathrm{P}=0.021)$ but not when adjusted for density $(\mathrm{P}=0.091)$. Consistent with management objectives, all BACIP contrasts were positive in the Methow population but only NOR was significantly positive ( $\mathrm{P}=0.017$ ). Mean total spawners, NOS, and NOR increased in both the Methow and reference populations between the before and after periods but the Methow population increased more than did the reference population during that period. The statistical power to detect differences of BACIP contrasts stabilized 10-20 years after supplementation, which was within the timeframe of this evaluation ( 24 years). Summer Chinook Salmon supplementation in the upper Columbia Basin produced mixed results relative to an un-supplemented reference population outside the Upper Columbia Basin, which may have been a function of density-dependence, but both programs contributed substantially to harvest.


## Introduction

One of the key uncertainties about using supplementation hatcheries is whether it can be done without negatively affecting natural-origin populations (Pearsons and Hopley 1999; Pearsons 2002; Fast et al. 2015). Well-run hatcheries can be used to increase abundance and harvest because they can produce more adults-per-spawner than can the natural environment (Fast et al. 2015). However, it is less clear whether hatcheries can be used to provide increases to abundance while keeping genetic and ecological impacts within acceptable limits (Ham and Pearsons 2001; Williamson et al. 2010; Chilcote et al. 2011). Mobrand et al. (2005) and Paquet et al. (2011) propose two supplementation strategies for containing risks to natural-origin populations while contributing to abundance and harvest. Their first strategy, termed "segregated," keeps the gene pools of hatchery- and natural-origin populations separate. Guidelines for accomplishing this segregation suggest that hatchery-origin fish should make up less than $5 \%$ of the natural-spawning population (Mobrand et al. 2005; Paquet et al. 2011). The second strategy, and the subject of this article, is to manage the hatchery program so that the gene pool of the hatchery population does not diverge from that of the natural-origin population from which the hatchery population originates. This strategy is termed "integrated."

In cases where it was not possible to achieve the gene-flow guidelines for a segregated program (i.e., gene flow $<5 \%$ ), integrated hatchery strategies were selected or were implemented before identification of a particular hatchery strategy (e.g., integrated or segregated). Many hatcheries had operated for decades before the identification and establishment of riskcontainment strategies. The guidelines for operating an integrated hatchery program promote the management of hatchery recruits (broodstock and hatchery returns that spawn in the natural environment) so that the dominant selection pressures operating within the supplemented population are from the natural environment, not the hatchery environment (Mobrand et al. 2005; Paquet et al. 2011). The combination of selection pressures from the hatchery environment is referred to as domestication selection and it has been indexed as Proportionate Natural Influence (PNI), which is a quantitative measure of the relative contributions of natural-environment and domestication selective pressures on populations subjected to hatchery programs. The relatively recent standard for reducing genetic risks of domestication is to match or exceed a PNI of 0.67 (Mobrand et al. 2005; Paquet et al. 2011; Pearsons et al. 2020). Domestication selection can reduce fitness in the natural environment (Araki et al. 2008; Fritts et al. 2007; Pearsons et al. 2007). Other unintentional effects of integrated harvest augmentation hatcheries can also include demographic changes (Knudsen et al 2006; Larsen et al 2013; Ford et al. 2015), undesirable ecological interactions (Pearsons et al. 2007; Pearsons et al. 2012; Temple and Pearsons 2012), and straying (Pearsons and O'Connor 2020, Keefer and Caudill 2014), and it can be challenging to disentangle the genetic and ecological mechanisms leading to these effects (Chilcote et al. 2011).

Most evaluations of the effects of hatchery supplementation on Chinook Salmon have focused on spring run Chinook Salmon that spawn in small river systems (Williamson et al 2010; Fast et al. 2015; Venditti et al. 2018). Populations that spawn in small rivers are much easier to evaluate than those in large rivers. For example, Chinook Salmon that spawn in large rivers can be more difficult to enumerate because of the difficulty in working in such large, deep, and often turbid environments. Furthermore, there are fewer populations of Chinook Salmon that spawn in large rivers to serve as reference populations for making comparisons between supplemented and non-supplemented populations. Without the ability to compare supplemented populations to
non-supplemented populations, identification of supplementation effects is hampered (Venditti et al. 2018).

The goal of this evaluation is to determine whether hatchery augmentation programs of summer Chinook Salmon in the Wenatchee and Methow rivers have increased the number of spawning adults while maintaining impacts to the natural populations within acceptable limits. More specifically, the purpose of this analyses is to determine whether the hatchery programs have changed the total adult abundance, natural-origin abundance, and productivity of summer Chinook Salmon in the Wenatchee and Methow rivers. We compared adult-to-adult productivities of fish that spawned in the hatchery and in the river to determine whether per capita hatchery production exceeded that in the river. In addition, we used paired Before-After-Control-Impact (BACIP) evaluations to determine whether supplementation increased abundance and productivity in supplemented populations relative to reference populations (Stewart-Oaten and Bence 2001; Pearsons and Temple 2010; Chevalier et al. 2019). BACIP analyses are among the most useful methods to evaluate the effects of long-term field treatments such as supplementation (Pearsons 2012), and recent improvements in analytical metrics have improved the utility of data interpretation from these evaluations (Chevalier et al. 2019).

## Methods

## Background

## Wenatchee River

The goal of summer Chinook salmon supplementation in the Wenatchee Subbasin is to use artificial production to replace Chinook salmon lost because of mortality at Priest Rapids, Wanapum, and Rock Island dams, while not reducing the natural production or long-term fitness of the extant summer Chinook population in the basin. The Rock Island Fish Hatchery Complex began operation in 1989 under funding from Chelan PUD and subsequently Grant PUD began cost-sharing the program in 2012. The Complex operated originally through the Rock Island Settlement Agreement, but since 2004 has operated under the Rock Island Anadromous Fish Agreement and Habitat Conservation Plan (HCP) as well as the Priest Rapids Project Salmon and Steelhead Settlement Agreement.

Adult summer Chinook are collected for broodstock from the Wenatchee River run-atlarge at the right- and left-bank traps at Dryden Dam, and at Tumwater Dam if weekly quotas cannot be achieved at Dryden Dam. Before 2012, the goal was to collect up to 492 natural-origin adults for the Wenatchee program for an annual release of 864,000 yearling smolts. In 2011, the Hatchery Committees reevaluated the amount of hatchery compensation needed to achieve no net impact (NNI). Based on that evaluation, the smolt-production goal of the program was reduced. The current goal (beginning with brood year 2012) was to collect up to 274 adult natural-origin adults for an annual release of 500,001 yearling smolts. The 500,001 smolts were the combined Grant PUD and Chelan PUD smolt production target, with Chelan PUD's obligation at 318,000 and Grant PUD's obligation at 182,001. Broodstock collection occurred from about 1 July through 15 September with trapping occurring up to 24 hours per day, seven days a week at Dryden Dam and up to 16 hours per day, three days per week at Tumwater Dam.

If natural-origin broodstock collection fell short of expectation, hatchery-origin adults were collected to meet the collection quota (see PNI chapter in this report).

Adult summer Chinook are spawned at Eastbank Fish Hatchery, where the majority of juveniles are reared in raceways, and a portion in re-use circular tanks. Juveniles are transferred from the hatchery to Dryden Acclimation Pond on the Wenatchee River, in March of each release year, and they are released from the pond volitionally beginning mid-April and pushed out by the end of April.

Before 2012, the production goal for the Wenatchee summer Chinook supplementation program was to release 864,000 yearling smolts into the Wenatchee River at ten fish per pound. Beginning with the 2012 brood, the revised production goal is to release 500,001 yearling smolts into the Wenatchee River at 18 fish per pound. Targets for fork length and weight are 163 mm $(\mathrm{CV}=9.0)$ and 45.4 g , respectively. Over $95 \%$ of these fish are marked with CWTs. In addition, since 2009, about 20,000 juveniles were PIT tagged annually.

## Methow River

The original goal of summer Chinook salmon supplementation in the Methow Basin was in part to use artificial production to replace Chinook Salmon lost because of mortality at Wells, Rocky Reach, and Rock Island dams, while not reducing the natural production or long-term fitness of summer Chinook in the basin. The Rock Island Fish Hatchery Complex began operation in 1989 under funding from Chelan PUD. The Complex operated originally through the Rock Island Settlement Agreement, but from 2004 to 2012 operated under the Rock Island and Rocky Reach HCPs. Beginning with broodstock collection in 2012, Grant PUD took over funding and operation of the summer Chinook salmon supplementation program in the Methow River basin. Grant PUD constructed a new overwinter acclimation facility adjacent to the Carlton Acclimation Pond and the first release of fish from this facility was in 2014. The first fish that were overwinter acclimated in the facility were released in 2015. The new facility includes eight, 30 -foot diameter dual-drain circular tanks.

Presently, adult summer Chinook are collected for broodstock from early July to midSeptember at the east-ladder trapping facility at Wells Dam. Before 2012, the goal was to collect up to 222 natural-origin adults for the Methow program. In 2011, the Hatchery Committees reevaluated that amount of hatchery compensation needed to achieve NNI. Based on that evaluation, the goal of the program was revised. The current goal (beginning with brood year 2012) is to collect up to 102 natural-origin adults for the Methow program. Broodstock collection occurs from about 1 July through 15 September with trapping occurring no more than 16 hours per day, three days a week. If natural-origin broodstock collection falls short of expectation, hatchery-origin adults can be collected to make up the difference.

Adult summer Chinook were spawned and progeny reared at Eastbank Fish Hatchery. Before the initiation of overwinter acclimation with juveniles from the 2013 brood year, juveniles were transferred from the hatchery to Carlton Acclimation Pond in March. Beginning with brood year 2013, juveniles have been transferred to the Carlton Acclimation Facility in October or November and released from the new facility the following spring in mid-April to early May.

Before 2012, the production goal for the Methow summer Chinook supplementation program was to release 400,000 yearling smolts into the Methow River at 10 fish per pound. Beginning with the 2012 brood, the revised goal is to release 200,000 yearling smolts at 13-17
fish per pound. Targets for fork length and weight are $163 \mathrm{~mm}(\mathrm{CV}=9.0)$ and 45.4 g , respectively. Over $90 \%$ of these fish were marked with CWTs. In addition, since 2009, 5,000 juveniles have been PIT tagged annually.

## Data Collection and Derived Metrics

Summer Chinook Salmon spawning ground surveys have been conducted since at least 1982 within the Wenatchee and Methow subbasins to determine the abundance, distribution, and origin of spawners. Since 1982, various methods have been used to estimate the number of summer Chinook Salmon redds in the Wenatchee and Methow rivers. Within the two rivers, redd counts have been conducted from aerial and ground surveys. In some years, only aerial counts were used and in other years both aerial and ground surveys were conducted. In addition, different ground surveys have been used. Those included (1) peak ground count, which was the maximum number of redds observed during a given survey; (2) total ground count, which involved mapping the location of all redds over the spawning period; and (3) peak expansion count, which was a hybrid of the two methods. The peak expansion count involved paired observations (peak and map counts) within an index area of a reach to provide an expansion factor for that reach. Ground surveys followed methods described in Gallagher et al. (2007) and Murdoch et al. (2010). Surveyors walked or floated the entire distribution of summer Chinook Salmon spawning habitat and identified and counted new redds weekly throughout the spawning season from late September to mid-November. Redds were flagged and locations recorded on a maps and/or Global-Positioning-System devices to avoid recounting in subsequent surveys. Aerial counts were conducted from fixed-wing aircraft and were usually conducted throughout the spawning period.

Peak ground counts and aerial counts only counted the maximum number of visible redds on a given survey and thus they did not distinguish between old or new redds on successive surveys and likely underestimated the true number of redds. Total ground counts mapped out all new redds as they were constructed so new and old redds were accounted for in the survey. Total ground counts were believed to be the most accurate survey method and this method has been used in the Wenatchee River since 2014 and in the Methow River since 1990.

Because redd counts were used to estimate escapements and several other derived metrics, it was important to have a normalized data set across all years that was not interrupted with changes in methodologies. We used expansion factors for the Wenatchee and Methow rivers to adjust the data for years in which aerial counts and peak counts were conducted. Reachspecific expansion factors were developed for each river to expand aerial or peak counts to an estimated ground count (see Miller et al. 2011).

For the Wenatchee River, summer Chinook Salmon redd counts have been estimated using aerial surveys, peak ground surveys, and peak expansion counts. Aerial counts were conducted in the Wenatchee River up to 1996. Peak ground counts began in 1987. In 2006, peak expansion counts were obtained for six of the ten reaches on the Wenatchee River. It was not until 2008 that all ten reaches of the Wenatchee River had separate index areas for map counts. In 2010, a complete data set was created to adjust for differences in survey methods over the years. We used a two-step process to adjust counts: (1) convert aerial counts to peak ground counts and (2) convert peak ground counts to an estimated total count (peak expansion count). To estimate peak ground counts from aerial counts, we used data from years (1990-1996) in which both surveys methods were conducted. Individual expansion factors for each year were
calculated as the aerial count divided by the peak ground count for each reach. Aerial counts from 1981 to 1987 were expanded by dividing reach-specific aerial counts for each reach by the reach-specific mean expansion factor. The reach-expanded aerial counts were then summed to derive an estimated peak ground count.

Peak ground counts were then expanded to estimate total redd counts in the Wenatchee River. This involved a two-step process using map counts to expand peak counts. During the first step, we used data collected during 2008-2010 when both peak ground counts and map counts were conducted in all reaches in the Wenatchee River. The map counts documented new or recently constructed redds within index areas during each survey. Peak counts were estimated within each reach (both index and non-index areas), while map counts only occurred within the index areas. An index area expansion factor was developed based on the ratio of peak to map counts for each index area within a reach. Reach-specific index area peak expansion factors were applied to all non-index areas and the expanded counts were summed along with the map count for the estimated total redd count for a reach. The non-index area peak counts were divided by the index peak expansion factor and then summed to get a reach total peak expansion estimate. The sum of all reach totals provides the total peak expansion redd count. During the second step, we applied the mean of reach-specific peak expansion factors to all years prior to 2006. Thus, we developed a time series of standardized redd counts that were adjusted for the different survey methods used in the Wenatchee River.

In the Methow River, prior to 1990, aerial surveys were used to estimate the number of summer Chinook Salmon redds. Comprehensive total ground counts on the Methow River began in 1990 and have continued since. Both total ground counts (map counts) and peak aerial counts were conducted in 1991 to 1996. In 1996, aerial surveys in the Methow River were discontinued. To adjust the aerial redd counts for the Methow River (redd counts prior to 1990), we used reach-specific expansion factors derived from aerial and total ground counts from 1991 to 1995. Individual expansion factors for each year were calculated as the aerial count divided by the total ground count for each reach.

Carcasses surveys were also conducted during spawning ground surveys in the Wenatchee and Methow rivers. All carcasses found were counted and examined to determine sex, origin (i.e., hatchery or natural origin; nearly all hatchery-origin fish were marked with a tag such as a CWT and/or adipose fin clipped prior to release), size, distribution, and other biological characteristics. In addition, scales collected from carcasses were used to determine fish age and origin.

We expanded spawning ground data into derived estimates of total adult abundance (combined hatchery- and natural-origin adults), natural-origin spawners (NOS) abundance, natural-origin adult recruits (NOR) abundance, adult-to-adult productivity, and adjusted adult-toadult productivity. The latter productivity was adjusted based on population carrying capacity (described below). We assumed that each female made one redd (Murdoch et al. 2009a), female carcass location was a good surrogate for spawning location by origin (Murdoch et al. 2009b), and that redd counts and carcass data could be used to estimate spawning escapement for hatchery- and natural-origin Chinook Salmon (Murdoch et al. 2010).

Spawning escapements for summer Chinook Salmon derived from redd counts were estimated using two different methods. Before 1998, spawning escapements were calculated as the number of redds times 3.1, based on work conducted by Meekin (1967). However, Meekin (1967) did not include jack Chinook Salmon in his estimate of 3.1 adults per redd. Therefore, Washington Department of Fish and Wildlife modified the 3.1 adults per redd ratio by including
jacks. The "modified-Meekin" method adjusted the 3.1 adults to account for the proportion of jacks counted in the summer Chinook Salmon run at Wells Dam (for Methow summer Chinook Salmon), or the difference in jack counts observed at Rock Island and Rocky dams (for Wenatchee summer Chinook Salmon). The modified-Meekin estimate was calculated as follows for Methow summer Chinook Salmon:

$$
\text { Modified Meekin }=3.1\left(1+\left(\frac{\text { Number of Jacks Counted at Wells Dam }}{\text { Total Count of Chinook at Wells Dam }}\right)\right)
$$

and as follows for Wenatchee summer Chinook Salmon:

$$
\text { Modified Meekin }=3.1\left(1+\left(\frac{\text { Rock Island Jack Count }- \text { Rocky Reach Jack Count }}{\text { Rock Island Total Count }- \text { Rocky Reach Total Count }}\right)\right)
$$

A second method was used to adjust estimates from 1998 to present, and it estimated adult escapement based on the annual male-to-female ratio determined from broodstock sampling at Wells Dam (for Methow summer Chinook Salmon) and Dryden Dam (for Wenatchee summer Chinook Salmon). The expansion factor was calculated as one plus the male-to-female ratio:

$$
\text { Sex Ratio Expansion Factor }=1+\left(\frac{\text { Number of Males }}{\text { Number of Females }}\right)
$$

After estimating total adult spawning abundance, we estimated natural-origin adult recruits by summing all adults produced from a brood year including adult spawners, hatchery broodstock, and harvest. Natural-origin spawners were estimated by multiplying the proportion of natural-origin carcasses by the total adult spawning abundance. Productivity was calculated by dividing the natural-origin adult recruits by the total number of spawners for each cohort. A density-adjusted estimate of productivity was calculated by assigning the productivity value at carrying capacity to brood years that exceeded the minimum estimated spawner abundance necessary to generate the asymptotic number of adult recruits as modelled using the Ricker stock-recruitment function (Ricker 1954). We used natural logarithms to transform recruitment data to satisfy modeling assumptions.

Finally, we evaluated whether adult productivity estimates were higher in hatchery or natural environments. Natural replacement rates (NRRs; natural-origin productivity) were calculated as the ratio of natural-origin recruits (NOR) to the parent spawning population (spawning escapement). Natural-origin recruits were naturally produced fish that survived to contribute to harvest, to broodstock, and to spawning grounds. Hatchery replacement rates (HRRs) were the hatchery adult-to-adult returns and were calculated as the ratio of hatcheryorigin recruits (HOR) to the parent broodstock collected. Both NOR and HOR included fish that spawned naturally, fish harvested, and fish used as broodstock in hatchery programs. A pairedsample t-test evaluated the hypothesis that there was no difference between mean HRR and NRR. In addition, annual HRRs were compared to HRRs targets established by the managers of the hatchery programs.

Selection of Reference Populations for BACIP Analyses

Reference populations are an important component of an effectiveness monitoring program because they provide the standard to which treatment conditions are compared (ISRP and ISAB 2005; Galbreath et al. 2008). Selecting appropriate reference populations and maintaining them over long periods of time is needed to assess the effectiveness of supplementation programs using BACIP analyses. Ideal reference populations possess biotic and abiotic characteristics similar to that of the supplemented populations. We developed a three-step process for identifying suitable reference populations for use in evaluating the effectiveness of hatchery supplementation programs.

The first step in selecting suitable reference populations included comparing general attributes of both supplemented and reference populations. Attributes included (1) no or few hatchery-origin spawners in the reference population (proportion of hatchery-origin spawners ( pHOS ) less than $10 \%$ ), (2) similar life-history characteristics (similar run timing, age structure, and spawn timing), (3) similar out-of-basin effects (similar migration and ocean survivals), (4) similar trends in freshwater habitat, (5) harvest estimates available for adjusting escapements, and (6) a long time series of natural-origin abundance and productivity estimates. Reference populations with high ( $>10 \%$ ) pHOS, no harvest data, or short time series were excluded from further consideration.

Reference populations that were deemed similar to supplemented populations based on Step 1 attributes were then evaluated for similar trends and synchronicity during the presupplementation period. For both Wenatchee and Methow summer Chinook Salmon, the presupplementation period was 1981 through 1991 for abundance and 1981 through 1995 for NOS, NOR, and productivity. Abundance, NOS, NOR, and productivity data were transformed with natural logarithms and compared graphically to evaluate trends and synchronicity between reference and supplemented populations before supplementation. In addition to graphic analysis, we calculated the Pearson correlation coefficient, which is an index of the strength of the association between reference and supplemented populations, and compared trends using analysis of covariance. Although there is some redundancy in these methods, in concert they allowed us to evaluate trends, fluctuations, and synchronicity between reference and supplemented populations prior to supplementation, and they improved precision of final analyses. As a final exercise under Step 2, we conducted power analyses to determine minimum differences that could be detected when comparing reference populations to supplemented populations. Because our study used BACIP designs, which includes replication before and after supplementation in both the supplemented and reference populations, we calculated difference scores using natural-log transformed data (Stewart-Oaten et al. 1992; Smith et al. 1993). We calculated minimal detectable differences assuming the use of an independent two-sample t-test with a type-I error rate of 0.05 , power of 0.80 (beta or type-II error rate of 0.20 ), and sample sizes (supplementation years) evaluated at five-year increments out to 100 years during supplementation. The objective was to find reference populations with the lowest minimal detectable differences.

During the final step in selecting suitable reference populations, we developed a method to quantify and rank potential reference populations. In order to reduce subjectivity, we developed a method of scoring and weighting selection criteria. For consistency, criteria were scored from 0 to 1 , with 0 being the worst possible score and 1 being the best. We also weighted each criterion. For example, proportion of hatchery-origin spawners was assigned a higher weight than statistical attributes (e.g., graphic analysis, correlation, and trends). We calculated a total score for each reference population by multiplying the estimated value, which ranged from

0 to 1 , by its weight. The sum of the weighted values provided a total score that ranged from 0 to 100. Based on several simulations, we set the cut-off score at 80 . That is, if the total score for a given reference population exceeded 80 , the population was included as a suitable reference. If the total score was equal to or less than 80 , the population was not considered a suitable reference.

Using the three-step process, we identified one reference population for each of the supplemented populations: Deschutes River fall Chinook Salmon in Oregon. That population had an abundance time series from 1981-2018 and a NOS, NOR, and productivity time series from 1989-2012. There were no NOS, NOR, and productivity data for Deschutes River fall Chinook Salmon before 1989. Therefore, our analyses included pre-supplementation years 1981-1991 and during supplementation years 1992-2018 for abundance and pre-supplementation years 19891995 and during supplementation years 1996-2014 for NOS, NOR, and productivity data. We found no reference populations in the upper Columbia Region that satisfied the criteria mainly because most were supplemented and those that were not supplemented had a high proportion of hatchery-origin strays in the spawning population.

## Data Analysis

We made comparisons between supplemented populations and the reference population using BACIP analyses (Stewart-Oaten and Bence 2001; Downes et al. 2002; Pearsons and Temple 2010; Chevalier et al. 2019). The objective of the comparison of supplemented to reference populations was to determine whether the addition of a hatchery supplementation treatment increased the total spawning abundance, NOS abundance, NOR abundance, and adult-to-adult productivity metrics described above. The after period for total spawning abundance was defined as the period after the year in which age- 3 hatchery-origin adults returned from the first broodstock collections. The after period for NOS, NOR, and productivity metrics was defined as the period beginning three years after the first age- 4 hatchery-origin females from the programs spawned in the natural environment. For example, if the first broodstock for a program was collected in 1989, the after period would begin in 1992 for total spawning abundance and in 1996 for NOS, NOR, and productivity. The analyses included adult returns up to 2018. We did not specifically evaluate the effects of reduced hatchery smolt production that occurred in 2014 on population metrics because the period following smolt reduction was too short to evaluate adequately. Thus, in our analyses, the supplementation (after) period included years both before and after reduction in smolt production, and each program was considered as a single, adaptively managed program rather than programs with multiple supplementation treatments.

To evaluate the effect of the supplementation programs, we calculated three BACIP metrics: BACIP Contrast (effect), Control Impact (CI)-contribution, and CI-divergence (Chevalier et al. 2019). The formulas to calculate the metrics were:

$$
\begin{aligned}
& \text { BACIP Contrast }=\left(\mu_{\mathrm{TA}}-\mu_{\mathrm{TB}}\right)-\left(\mu_{\mathrm{CA}}-\mu_{\mathrm{CB}}\right) \\
& \text { CI-contribution }=\left|\mu_{\mathrm{TA}}-\mu_{\mathrm{TB}}\right|-\left|\mu_{\mathrm{CA}}-\mu_{\mathrm{CB}}\right| \\
& \text { CI-divergence }=\left|\mu_{\mathrm{TA}}-\mu_{\mathrm{CA}}\right|-\left|\mu_{\mathrm{TB}}-\mu_{\mathrm{CB}}\right|
\end{aligned}
$$

where $\mu$ is the mean, T is treatment (supplemented), C is control (reference), A is after (period of supplementation), and B is before (period before supplementation). Evaluation of a combination of BACIP metrics contributes to an understanding of why BACIP contrast values differ (Chevalier et al. 2019).

In this evaluation, we interpret a significant $(\mathrm{P} \leq 0.05)$ BACIP Contrast value as an indication that the dependent variable (e.g., NOS) has become significantly different in the supplemented population relative to the reference population between the before and after time periods. A significant positive value indicated that the dependent variable increased more (or decreased less) in the supplemented population than the reference population, and a significant negative value indicated that the dependent variable decreased more (or increased less) in the supplemented population than the reference population (Chevalier et al. 2019). For BACIP evaluations, we tested differences between the supplemented populations and reference population using an Aspin-Welch unequal-variance test. This test evaluated the hypothesis that the mean difference before supplementation equals the mean difference during supplementation. This test was conducted separately for each supplemented population with the reference population. All analyses were conducted on natural-log-transformed data. In total, this work resulted in the calculation of 10 Aspin-Welch tests. We did not adjust the value for significance ( $\mathrm{P} \leq 0.05$ ) based on the large number of tests conducted. Rather, we reported the actual P -value for each test.

Following the BACIP Contrast evaluations, the relationship between supplemented and reference populations was further evaluated using a CI-contribution value and CI-divergence value (Chevalier et al. 2019). The CI-contribution value was used to identify which population (supplemented or reference) made the larger contribution to the change from the before to the after period, which was indicated by the sign (negative or positive) of the CI-contribution value. For example, a positive value indicated that the observed difference in the BACIP Contrast was largely attributable to changes in the supplemented population, whereas a negative value indicated that the difference was largely attributable to changes in the reference population. The CI-divergence value was used to indicate the change in similarity or dissimilarity between the supplemented and reference populations from the before to the after period. A positive value indicated that the two populations were more different in the after period than the before (i.e., diverged), whereas a negative value indicated that the supplemented and reference populations were more similar in the after period than the before period (i.e., converged). In this analysis, we were only interested in the sign of the value.

## Power Analysis

Power analyses were conducted to determine the length of time necessary to detect different effect sizes in BACIP contrasts. This analysis calculates the minimum detectable difference between mean difference scores (Treatment-Control) before and after treatment and was a different analysis than the one that was done to evaluate the suitability of reference populations describe above. The variance estimates during the pretreatment period and posttreatment period were used to calculate minimum detectable differences. The null hypothesis tested was that the mean difference before treatment equals the mean difference after treatment. The alternative hypothesis is that the mean difference before treatment is different than the mean difference after treatment (two-tail test). This analysis assumes that mean difference scores will be compared using an independent $t$-test with an alpha of 0.05 , beta of 0.20 , and sample sizes (post-treatment years) of 5 to 100 by 5 .

Many other analyses specified in the monitoring and evaluation plan (Hillman et al. 2019) were conducted but were not presented here because of the redundancy of test results and complexity of presentation. Additional statistical tests supported the conclusions in this
evaluation. We prioritized simplicity and statistical power in this report in order to distill the main findings of the analysis. The full suite of analyses is available upon request.

## Results

## HRR vs. NRR

The HRRs of summer Chinook Salmon were significantly higher than corresponding NRRs in the Wenatchee and Methow ( $\mathrm{P}<0.05$; Tables 1 and 2). On average, HRRs were 3.6 times higher than NRRs in the Wenatchee and 2.8 times higher than NRRs in the Methow. In addition, the HRRs exceeded NRRs in most years ( $79 \%$ of years in the Wenatchee and $67 \%$ of years in Methow) and exceeded management targets in each of the last 5 years (Table 2). The HRRs in the Wenatchee were 1.6 times higher than HRRs in the Methow.
Stock-Recruitment Analysis
We successfully fit the Ricker stock-recruitment function to the Wenatchee, Methow, and Deschutes Chinook Salmon populations (Figure 1). The model estimated that 6,250 spawners are needed to produce the maximum number of NOR in the Wenatchee, 2,326 spawners are needed in the Methow, and 9,091 spawners are needed in the Deschutes. These estimates were used to adjust productivity estimates within each population.

Table 1. Productivities of adult natural-origin recruits (NOR) and hatchery-origin recruits (HOR) summer Chinook Salmon in the Wenatchee and Methow rivers.

| Program/Stock | Natural-origin Chinook |  |  |  | Hatchery-origin Chinook |  |  |  |
| :---: | :---: | :---: | :---: | :---: | :---: | :---: | :---: | :---: |
|  | Number <br> of years | Mean <br> spawning <br> escapement | Mean <br> NOR | Mean <br> NRR | Number <br> of years | Mean <br> broodstock <br> collection | Mean <br> HOR | Mean <br> HRR |
|  | 24 | 9,489 | 20,418 | 2.65 | 24 | 405 | 4,015 | 9.58 |
| Methow | 24 | 1,756 | 3,054 | 2.16 | 24 | 219 | 1,285 | 6.01 |

Table 2. Comparisons of productivities of adult natural-origin replacement rates (NRR) and hatchery-origin replacement rates (HRR) summer Chinook Salmon in the Wenatchee and Methow rivers relative to each other and management targets.

| Program/Stock | HRR Target Analysis |  |  |  |  | Paired-Sample T-Test |  |  |
| :---: | :---: | :---: | :---: | :---: | :---: | :---: | :---: | :---: |
|  | HRR Target | No. Years | No. Yrs HRR $\geq$ Target | No. Yrs HRR<Target in last 5 Yrs* | No. Yrs HRR>NRR |  |  |  |
|  |  |  |  |  |  | t-value | P-value | Power |
| Wenatchee | 5.7 | 24 | 13 | 0 | 19 | 3.551 | 0.002 | 0.925 |
| Methow | 3.0 | 24 | 13 | 0 | 16 | 2.730 | 0.012 | 0.744 |

[^1]

Figure 1. Relationships between number of spawners and natural-origin recruits (NORs) for the Wenatchee, Methow, and Deschutes Chinook Salmon populations. Figures show the fit of the Ricker stock-recruitment model; $\mathrm{K}_{\mathrm{sp}}=$ number of spawners need to produce the maximum number of recruits; $K_{R}=$ maximum number of recruits.

## BACIP Analysis

## Wenatchee River

The BACIP contrasts for total spawners and NOS were significantly negative in the Wenatchee ( $\mathrm{P} \leq 0.001$; Table 3). Mean total spawners and NOS decreased slightly from the before to after periods in the Wenatchee but increased in the reference population (Figure 1). Mean total spawners between the Wenatchee and reference populations converged from the before to after periods, while NOS diverged. In both cases, the reference population changed more than did the Wenatchee population from the before to after period.

Table 3. Summary of statistical tests (Aspin-Welch unequal-variance test) and BACIP metrics on LN abundance and productivity for summer Chinook Salmon hatchery programs in the Wenatchee and Methow rivers. Deschutes River fall Chinook Salmon served as the reference population for BACIP analyses.

| Population | Statistic | Abundance | NOS | NOR | Productivity | Adj. <br> Productivity |
| :---: | :---: | :---: | :---: | :---: | :---: | :---: |
|  | P-value | 0.001 | 0.000 | 0.398 | 0.022 | 0.084 |
|  | BACIP Contrast | -0.732 | -0.930 | 0.352 | 0.821 | 0.575 |
|  | CI-contribution | -0.450 | -0.264 | 0.352 | 0.245 | 0.316 |
|  | CI-divergence | -0.274 | 0.092 | -0.352 | -0.536 | -0.190 |
| Methow | P-value | 0.247 | 0.867 | 0.017 | 0.087 | 0.143 |
|  | BACIP Contrast | 0.276 | 0.036 | 1.017 | 0.535 | 0.436 |
|  | CI-contribution | 0.276 | 0.036 | 1.017 | -0.041 | 0.177 |
|  | CI-divergence | -0.276 | -0.036 | -1.017 | -0.535 | -0.436 |

The BACIP contrast for NOR was not significant and both the Wenatchee and reference populations increased slightly from the before to after periods (Figure 2). Mean NOR between the Wenatchee and reference populations converged during the after period (became more similar) with the Wenatchee population changing the most from the before to after periods. The BACIP contrast for productivity was significantly positive $(\mathrm{P}=0.022)$, but after adjusting for density dependence, the adjusted productivity was not significantly different ( $\mathrm{P}=0.084$; Table 3). Adjusted productivity increased in the Wenatchee and decreased in the Deschutes between the before and after periods (Figure 2). Mean productivity between the Wenatchee and reference populations converged from the before to after period, with the Wenatchee population changing most during that period.

## Methow River

All BACIP contrasts were positive in the Methow but only NOR was significantly positive (Table 3). Mean total spawners, NOS, and NOR increased in both the Methow and reference populations between the before and after periods but the Methow population increased more than did the reference population during that period (Figure 2). Thus, mean population metrics converged between the before and after periods. Adjusted productivity increased in the Methow and decreased in the reference population between the before and after periods.


Figure 2. Comparisons of mean total spawning abundance, natural-origin spawners (NOS), natural-origin recruits (NOR), and density-adjusted productivity (with $95 \% \mathrm{CI}$ ) before and after hatchery supplementation of summer Chinook Salmon in the Wenatchee and Methow rivers and in the Deschutes River (reference population). Graphs include BACIP contrasts and their significance (P-values based on Aspin-Welch test), CI-contribution values, and CI-divergence values.

## Power Analysis

The statistical power to detect significant BACIP contrasts for all population metrics stabilized 10-20 years after supplementation, which was within the timeframe of this evaluation (Figure 3). The power to detect BACIP contrasts differed between the Wenatchee and Methow populations, with greater statistical power for detecting supplementation effects in the Wenatchee population than in the Methow population. This is primarily because the BACIP contrast (treatment effect) for the Wenatchee was much larger than the BACIP contrast for the Methow.


Figure 3. Statistical power to detect a BACIP difference in total spawner abundance, naturalorigin spawners (NOS), natural-origin recruits (NOR) and productivity of summer Chinook Salmon spawners in the Wenatchee and Methow rivers. Deschutes River fall Chinook Salmon served as the reference population. Curves were developed based on an independent t-test with an alpha of 0.05 , beta of 0.20 , and sample sizes (post-treatment years) of 5 to 100 by 5 .

## Discussion

Results were somewhat inconsistent between the Wenatchee and Methow subbasins even though the reference population used in the analyses was the same for both. The abundance metrics in the Wenatchee were lower (total spawners, NOS) or indifferent (NOR) relative to the un-supplemented reference population, which was contrary to management objectives of an increase in these metrics. In contrast, the abundance metrics for the Methow River were consistent with management objectives because all BACIP contrast metrics were positive and NOR was significantly higher. Additionally, the density-adjusted productivity did not appear to be negatively influenced by supplementation in either the Wenatchee or Methow rivers, which is also consistent with management expectations.

One possible explanation for the differences in the Wenatchee and the Methow rivers was differences in the performance of the hatcheries (i.e., differences in HRRs). However, the Wenatchee HRRs were only 1.6 times higher than those in the Methow, so this did not appear to explain the difference. It is currently unclear why the Wenatchee River abundance metrics did not increase in the after period as did those metrics in the Methow and Deschutes rivers in the after period. It is also unclear why contrasts of BACIP total spawner abundance and NOS in the Wenatchee River were lower than in the Deschutes River when the Wenatchee River HRRs were significantly higher than the NRRs. In addition, the productivities were not negatively affected by the supplementation programs, which would be one possible mechanism for explaining why we observed reduced total spawner abundance and NOS BACIP contrasts in the Wenatchee River relative to the Deschutes River.

The negative CI-Contribution for Wenatchee abundance suggests that the difference in BACIP contrast was primarily attributable to the increase in the Deschutes abundance, rather than to the slight decrease in Wenatchee abundance. Likewise, the negative CI-Divergence suggests that the mean abundances of the Wenatchee and Deschutes populations were more similar in the after period than during the before period. Supporting analyses indicate that the mean abundance in the Wenatchee for both the before and after periods is similar and both those means exceed the stock size at capacity estimated from the Ricker stock-recruitment functions. Additionally, mean abundance increased from the before to after period in both the Methow and the Deschutes; nevertheless, mean abundance in the after period remains below the stock sizes at capacity for both populations (based on the Ricker stock-recruitment functions). A likely explanation, then, for the differences in observed abundance BACIP contrasts for the Wenatchee and Methow populations, is that both the Methow and the Deschutes had the capacity for increased abundance in the after period, while the Wenatchee did not.

Hatchery programs in both the Wenatchee and Methow rivers contribute significantly to harvest in the ocean and freshwater environments and have some of the highest harvest rates in the Columbia River (O'Connor and Pearsons 2021 this report). The hatchery programs support these high rates of harvest (e.g., 60-70\%) without influencing the sustainability of the integrated populations. The lack of significant differences in adjusted productivity relative to the unsupplemented Deschutes River population suggested that the hatcheries are not contributing to reduced productivity in the natural environment, an important concern when managing to achieve sustainability.

The findings of hatchery supplementation of summer Chinook Salmon in this evaluation were quite different than those for spring Chinook Salmon in the same subbasins. Abundance and productivity metrics for spring Chinook Salmon were generally negative when compared to
reference populations (Pearsons et al. 2021). Summer Chinook Salmon were much more abundant than spring Chinook Salmon and have different life-history strategies (Quinn 2018). Summer Chinook Salmon in the upper Columbia basin have fared better than spring Chinook Salmon both from a hatchery and population perspective, and numerous aspects of their respective life-history strategies contribute to the differential status of these Chinook Salmon stocks.

Including reference populations in the Upper Columbia watershed would have enhanced our ability to definitively evaluate the influence of supplementation. We cannot exclude the possibility that differences in responses, which occurred between the supplemented populations and reference population during supplementation, were caused by regional factors (Dorner et al. 2018) independent of supplementation. For example, habitat improvement projects or other fisheries management in the Deschutes River may have been more effective than those in the Wenatchee or Methow rivers. In addition, fish from the population in the Deschutes River migrate through two dams and those in the upper Columbia migrate through seven to nine dams. Alternatively, there may have been changes in predation that occurred during the evaluation (Fritts and Pearsons 2004, 2006, 2008), which may have resulted in detecting an Upper Columbia watershed effect unrelated to supplementation. Productivity metrics for both supplemented populations may have manifested similar increases relative to those of the reference population, which could suggest an effect either from supplementation or from something specific to the Upper Columbia watershed. Without the addition of a reference population in the Upper Columbia, we cannot be certain that observed effects were related entirely to supplementation or a combination of supplementation and regional effects. A local reference population would have strengthened inferences regarding population metrics.

The results presented were mostly the effects of the original hatchery production targets coupled with recent adaptive management of the programs. Beginning in 2014, the release numbers were greatly reduced for most long-term programs as a result of adjustments in hatchery production required by the Federal Energy Regulatory Commission licenses for the respective hydroelectric projects for which the hatchery programs mitigate. Because of the timing of these changes, we were unable to independently evaluate their effects at this time. Power analyses indicated that minimum detectable differences stabilized approximately 10-20 years after supplementation and therefore it is likely that statistical power to detect differences would not be fully realized in less than 10 years (Ham and Pearsons 2000). In addition, effects on populationlevel metrics resulting from specific modifications to program management (e.g., changes in program size every 10 years) would be difficult to detect because the duration of those modifications to program management rarely exceeds 10 years before the implementation of additional modifications. In contrast, evaluation of the full adaptively managed programs since inception provided statistical power as high as could be expected because the number of treatment years exceeded the time interval necessary to detect differences.

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# Investigations into Association between Proportion of Hatchery Spawners and Juvenile Productivity for Spring Chinook, Summer Chinook, and Summer Steelhead in the Wenatchee and Methow River Basins 

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#### Abstract

Conservation hatchery programs implemented in the Wenatchee and Methow river basins are intended to increase the abundance of the target populations. The strategy of the hatchery programs is to return hatchery adults that increase the spawning population, resulting in an increase in the number of offspring that will maintain and recover naturally reproducing populations. However, there is a concern that the hatchery programs may lower the juvenile productivity in supplemented watersheds. The populations are managed under the concept of Proportionate Natural Influence (PNI), a framework that uses genetic modeling to estimate risk of hatchery introgression in natural populations and prescribes management benchmarks intended to allow hatchery programs to operate within an acceptable risk level to the natural population. One of the metrics in the PNI framework is the proportion of hatchery origin spawners ( pHOS ). Our investigation of whether higher pHOS may result in lowered juvenile productivity was largely inconclusive. For most populations studied, no evidence of an effect of pHOS was observed. For the single population that demonstrated a possible negative effect of pHOS on juvenile productivity (Twisp River Summer Steelhead), the evidence was weak: when density dependence was accounted for, the perceived negative association between pHOS and juveniles per redd was no longer observed. Additionally, the significant result was observed only when a possible outlier in emigrant counts was omitted, which may not be justified depending on the cause of the unusual emigrant count for that brood year. Thus, even when some evidence of a negative effect of pHOS was found, the evidence was particularly weak. For all three stock-recruitment models considered, there was often high uncertainty in model parameter estimates even when the models could be fit to the data. This was indicated by wide confidence intervals that often included maximum smolt or emigrant values far beyond the range of observed data, and by the high correlation in model parameter estimates. High correlation lowers the ability to distinguish between different model fits and increases uncertainty in the results; it arises from lack of contrast in the data or violation of modeling assumptions. Given all the complicating factors identified here, only the most extreme effect of pHOS on juvenile productivity could have been detected. Thus, we caution against concluding that such an effect is truly absent. We conducted a power analysis based on the Chiwawa spring Chinook data-the most robust dataset in this study. Simulated power analyses revealed that studies shorter than 70 years will have low power to detect an effect on recruitment for all but the strongest effect sizes of pHOS. Future data-collection efforts could be substantially improved by increasing the contrast in pHOS levels and including suitable in-basin unsupplemented references. More specifically, a planned experiment using deliberately chosen pHOS levels could be implemented in order to achieve the necessary contrast and remove confounding analyses with temporal and age-structured processes. Appropriate power analyses must be performed on the study design prior to commencing work to ensure that a result may be obtained within an acceptable time frame.


## Introduction

Conservation hatchery programs implemented in the Wenatchee and Methow river basins are intended to increase the abundance of the target populations. The strategy of the hatchery programs is to return hatchery adults that increase the spawning population, resulting in an increase in the number of offspring that will maintain and recover naturally reproducing populations. The assumptions are that 1 ) increasing the number of spawners will increase the naturally reproducing population abundance in future generations, and 2) hatchery-origin spawners have similar fitness to natural-origin spawners and will not decrease the productivity of natural spawning fish. Assessment of the production of juvenile offspring prior to entering the ocean provides information on freshwater productivity and avoids ocean effects on the abundance and productivity of the population. However, there is a concern that the hatchery programs may lower the juvenile productivity in supplemented watersheds. The populations are managed under the concept of Proportionate Natural Influence (PNI; Hatchery Scientific Review Group, 2009), a framework that uses genetic modeling to estimate risk of hatchery introgression in natural populations and prescribes management benchmarks intended to allow hatchery programs to operate within an acceptable risk level to the natural population. PNI is composed of two metrics: Proportion of Hatchery Origin Spawners ( pHOS ; the proportion of hatchery fish in the naturally spawning population), and Proportion of Natural Origin Broodstock ( pNOB ; the proportion of natural-origin fish in a hatchery broodstock). Part of ongoing monitoring and evaluation (M\&E) activities is to assess the relationship between pHOS and measures of juvenile productivity including the number of recruits to the smolt or emigrant populations and the average number of juveniles per redd. This report describes the statistical methods and results used to investigate these questions. In addition, we performed a power analysis to estimate the sample sizes required to detect an effect of pHOS on recruitment at varying pHOS effect strengths.

## Methods

Section 3.1 of the monitoring and evaluation plan for the hatchery programs (Hillman et al. 2019) addresses freshwater juvenile productivity and presents two questions:

Question 1: Has the supplementation program changed the number of juveniles (smolts, parr, and/or emigrants) per redd within the supplemented population?

Question 2: Does the number of juveniles per redd decrease as the proportion of hatchery spawners increases?

The first question could not be assessed because it requires reference populations that are unavailable for the target populations. Furthermore, estimates of juvenile production are not available in a long enough time series to estimate the juvenile productivity before and after a hatchery program was implemented. However, the basic concepts of Question 1 are addressed in Questions 2, to the extent the available data allowed.

The second question was assessed in this report. The data and methods used to assess Question 2 are described below.

## Data

All data were collected under the Monitoring and Evaluation Programs operated to assess the hatchery programs of Douglas County, Chelan County, and Grant County PUDs (Hillman et al. 2020; Snow et al. 2020). The abundance of spawners and redds was assessed by conducting spawning ground surveys. The number of redds was tabulated and for semelparous species, carcass recoveries were used to estimate the proportions of males and females by origin (hatchery and natural origin). Summer steelhead biological data for gender and origin were obtained at sampling points (Wells Dam and Twisp Weir). Abundance of semelparous spawners was estimated by expansion of redd counts based on the estimated portions of males to females, by origin, and assuming the number of redds created by a single female. Abundance of steelhead was estimated by adjusting dam and/or trap counts for proportions of fish estimated to return to the various tributaries in the Upper Columbia based on previous radio telemetry studies. Escapement estimates were adjusted to compensate for fall back and re-ascension at dams and to exclude fish removed from the spawning population through harvest and broodstock collection.

Juvenile abundance was estimated using rotary screw traps to sample migrating juveniles. Estimates were generated by conducting trap efficiency trials across a range of river flows, generating a regression model (flow versus trap efficiency), and using the model to predict the number of emigrants based on mean daily flow and daily captures. The daily estimates were summed for each year (season) to estimate the total abundance of emigrating juveniles per population. Juvenile steelhead were parsed into cohorts based on age determined through scales.

Spring chinook juveniles were yearlings assumed to be emigrating to the ocean. Age $0+$ spring Chinook migrants were not included in the analysis because their life history and survival are not well understood and we preferred to focus on emigrating fish when possible. However, age $0+$ spring Chinook that migrate for tributaries to main stem rivers may be an important component of the population (Copeland, et al. 2014). These age $0+$ migrants would be assessed by the lower rotary screw trap population estimates as age 1 emigrants. Summer Chinook emigrants were subyearlings and assumed to be exhibiting an ocean-type life history. All age classes (excluding age 0 ) of steelhead from each cohort were included in the estimate of cohort emigrants because of the difficulty in knowing how to parse emigrating fish from those that are not leaving the freshwater system. Therefore, steelhead brood year cohorts include the entire range of observed age classes for each cohort. For the sake of simplicity, we refer to the steelhead juveniles caught in the screw traps as "emigrants."

Douglas County PUD compiled the productivity data described above for the populations listed in Table 1. The size of the adult spawning stock, number of redds, and proportion of hatchery origin spawners ( $\mathrm{pHOS} \mathrm{)} \mathrm{were} \mathrm{provided} \mathrm{for} \mathrm{each} \mathrm{brood} \mathrm{year}$, or emigrants. No data were available for Summer Steelhead from the Wenatchee River because juveniles have not been assigned to brood year cohorts in the available data (Josh Williams, WDFW, personal communication).

Table 1. Summary of spawner and recruitment data compiled by Douglas County PUD. The size of the adult spawning stock, number of redds, and proportion of hatchery origin spawners ( pHOS ) were provided for each brood year, in addition to the number of smolts or emigrants.

| Species and Run | Population | Brood <br> Years | Smolts | Emigrants $^{\text {a }}$ | Source | Comment |
| :--- | :--- | :---: | :---: | :---: | :--- | :--- |
| Spring Chinook | Chiwawa | $1991-2017$ | X |  | Hillman et al. 2020 |  |
| Spring Chinook | Nason | $2002-2017$ | X |  | Hillman et al. 2020 |  |
| Spring Chinook | White | $2005-2017$ | X |  | Hillman et al. 2020 |  |
| Spring Chinook | Twisp | $2003-2017$ | X |  | Snow et al. 2020 |  |
| Spring Chinook | Methow | $2002-2017$ | X |  | Snow et al. 2020 |  |
| Summer Chinook | Wenatchee | $1991-2018$ |  | X | Hillman et al. 2020 | No emigrant data: |
| Summer Chinook | Methow | $2006-2018$ |  | X | Snow et al. 2020 | No emigrant data: 2012 |
| Summer Steelhead | Wenatchee |  |  |  | Hillman et al. 2020 | No data |
| Summer Steelhead | Methow | $2003-2015$ |  | X | Snow et al. 2020 |  |
| Summer Steelhead | Twisp | $2003-2015$ |  | X | Snow et al. 2020 |  |

$\mathrm{a}=$ Emigrant data were provided for Spring Chinook Salmon populations from the Chiwawa, Methow, and Twisp rivers but were not used in the analysis. On average, $52 \%$ of Chiwawa and $57 \%$ of Twisp spring Chinook emigrate from these rivers as age $0+$ fish.

## Statistical Methods

Spawner, redd count, and juvenile recruitment data were analyzed together with proportion of hatchery spawners to address Question 2, stated above (Hillman et al. 2019): does the number of juveniles per redd decrease as the proportion of hatchery spawners increases? The following statistical hypotheses were investigated:
$\mathrm{H} 0{ }_{2.2 \text {. 1.1. }}$ : There is no association between the proportion of hatchery-origin spawners ( pHOS ) and the residuals from the smooth hockey stick stock-recruitment curve; $\rho=0$.
$\mathrm{H} 0_{\text {2.2.1.2: }}$ : The slope between proportion of hatchery spawners and juveniles $/$ redd is $\geq 0$.
Three stock-recruitment models were used to account for density dependent mortality in assessment of the two hypotheses. The Smooth Hockey Stick model assumes that recruitment increases quickly at low levels of spawners and asymptotes to a maximum recruitment level as spawners increase:

$$
\begin{equation*}
R=R_{\infty}\left(1-e^{-\left(\frac{\alpha}{R_{\infty}}\right) s}\right) \tag{1}
\end{equation*}
$$

where $\mathrm{R}=$ recruits and $\mathrm{S}=$ spawners for a given brood year, $R_{\infty}$ is the maximum number of recruits (population carrying capacity), and $\alpha$ is the slope at low levels of spawner abundance (Froese 2008).

The Beverton-Holt model also assumes asymptotic growth to a maximum:

$$
\begin{equation*}
R=\frac{\alpha S}{\beta+S} \tag{2}
\end{equation*}
$$

where $\mathrm{R}=$ recruits and $\mathrm{S}=$ spawners for a given brood year, $\alpha$ is the asymptotic maximum number of recruits (population carrying capacity), and $\beta$ is the predicted spawner abundance required to produce half the maximum recruits (Hilborn and Walters 1992).

The Ricker model assumes that recruitment increases at lower levels of spawners and declines at higher spawner abundance:

$$
\begin{equation*}
R=\alpha S e^{-\beta S} \tag{3}
\end{equation*}
$$

where $\mathrm{R}=$ recruits and $\mathrm{S}=$ spawners for a given brood year, $\alpha$ is the increase in recruits per spawner at low levels of spawners and $\beta$ is the intensity of the decrease in recruitment at high levels of spawner abundance (Ricker 1954). The maximum number of recruits is defined as $K=$ $(\alpha / \beta) e^{-1}$.

Modeling assumptions were (based on Hillman et al. 2019):

1. Density-dependent mortality: The brood instantaneous mortality rate is proportional to the number of spawners for the brood year (Ricker 1954).
2. Lognormal errors: The variation in recruitment about the modeled recruitment for a given spawner abundance is lognormally distributed and acts multiplicatively (Quinn and Deriso 1999).
3. Negligible measurement error: Measurement error in spawning stock size and recruitment is small relative to the range of observed spawning stock sizes and variation in recruitment, respectively (Hilborn and Walters 1992).
4. Stationarity: The average stock-recruitment relationship is constant over time, with the possible exception of effects of changes in pHOS (Hilborn and Walters 1992).

The proportion of hatchery origin spawners may have the effect of lowering the maximum recruitment or recruits per spawner. The pHOS variable may be introduced into the Beverton-Holt model to reduce maximum recruitment when pHOS is high as follows:

$$
\begin{equation*}
R=\frac{\alpha e^{-\beta_{2} P_{\text {Hos }} S}}{\beta+S} \tag{4}
\end{equation*}
$$

or equivalently

$$
\begin{equation*}
\ln (R / S)=\ln (\alpha)-\ln (\beta+S)-\beta_{2} P_{\mathrm{HOS}} \tag{5}
\end{equation*}
$$

In Equations (4) and (5), the maximum number of recruits per brood year is $\alpha$ when pHOS is 0 and decreases asymptotically to $\alpha e^{-\beta_{2}}$ when pHOS is 1 .

The pHOS variable may be introduced into the Ricker model in such a way as to lower the slope in the spawner-recruitment curve at low levels of spawner abundance as follows:

$$
\begin{equation*}
R=\alpha e^{-\beta_{2} P_{\text {HOS }}} S e^{-\beta S} \tag{6}
\end{equation*}
$$

or equivalently

$$
\begin{equation*}
\ln (R / S)=\ln (\alpha)-\beta_{2} P_{\text {HOS }}-\beta S . \tag{7}
\end{equation*}
$$

Alternatively, if higher levels of pHOS are expected to increase the intensity of the decrease in recruitment at high spawner levels, then pHOS may be incorporated as follows:

$$
\begin{equation*}
R=\alpha S e^{-\left(\beta+\beta_{2} P_{H O S}\right) S} \tag{8}
\end{equation*}
$$

or equivalently

$$
\begin{equation*}
\ln (R / S)=\ln (\alpha)-\left(\beta+\beta_{2} P_{\text {HOS }}\right) S . \tag{9}
\end{equation*}
$$

Both models in Equation (6) and Equation (8) result in lower maximum recruitment for higher pHOS if $\beta_{2}>0$.

## Hypothesis $\mathrm{H0}_{2.2 .1 .1}$ Effect of pHOS on Recruitment:

Hypothesis $\mathrm{H}_{2.2 .1 .1}$ was investigated by fitting a stock-recruitment model to the available data and regressing the residuals from the model against pHOS. The Smooth Hockey Stick model, Beverton-Holt model, and Ricker model were considered.

Modeling assumption 1 was assessed by investigating the linear relationship between the log of recruits per spawner with the number of spawners; a negative relationship was consistent with density-dependent mortality. Modeling assumption 2 was assessed via quantile-quantile plots and Shapiro-Wilk tests (Shapiro and Wilk 1965) performed on the residuals from the fitted stockrecruitment model on the log scale. Modeling assumption 4 was assessed by plotting model residuals versus brood year and examining for autocorrelation. The available data did not allow for assessment of assumption 3; violation of assumption 3 would result in lower ability to detect a relationship between spawner abundance and recruitment (Hilborn and Walters 1992).

The fits of the three stock-recruitment models in Equations (1), (2), and (3) to the data were ranked using AICc (Burnham and Anderson 2002). Models with $\Delta \mathrm{AICc} \leq 2$ compared to the minimum observed AICc were used to assess the association between residuals and pHOS . The Smooth Hockey Stick model was included regardless of its AICc rank. Additionally, bias, uncertainty, and correlation in estimates of model parameters were estimated using bootstrapping with at least 1,000 bootstrap samples. Point estimates, $95 \%$ bootstrap confidence intervals, and bootstrap correlation coefficients between model parameters were reported for each model. In the event that bootstrapping failed, asymptotic (normal theory) confidence intervals and correlation coefficients were provided. Bias and correlation were considered in assessment of model fit.

For a given model, the residual for brood year $i(i=1, \ldots, n)$ was defined as

$$
\begin{equation*}
r_{i}=R_{i}-\hat{R}_{i}, \tag{10}
\end{equation*}
$$

where $R_{i}$ is the observed number of recruits for brood year $i$, and $\hat{R}_{i}$ is the predicted number of recruits from the model for brood year $i$. The association between the residuals $r_{i}(i=1, \ldots, n)$
and pHOS was investigated using the Pearson correlation coefficient and visual inspection of the fitted linear model

$$
\begin{equation*}
r_{i}=\beta_{0}+\beta_{1} p^{H_{O} O S_{i}}+\epsilon_{i} \tag{11}
\end{equation*}
$$

for regression coefficients $\beta_{0}$ and $\beta_{1}$ and random error $\epsilon_{i} \sim N\left(0, \sigma_{r}^{2}\right)$. Hypothesis H02.2.1.1 was rejected if the Pearson correlation coefficient between $r_{i}$ and pHOS was negative and significantly different from $0(\mathrm{P}<0.05)$. In this event, we concluded that there was evidence that hatchery fish may be reducing the productivity of the wild population.

An additional assessment of the effect of pHOS on juvenile productivity was performed by fitting the pHOS-enhanced stock-recruitment models in Equations (4), (6), and (8) and testing whether $\beta_{2}=0$ using likelihood ratio tests.

## Hypothesis $\mathrm{H0}_{2.2 \text {.1.2 }}$ Effect of pHOS on Juveniles per Redd:

Hypothesis H02.2.1.2 was investigated in two ways. First, the ratio of recruits per redd was modeled using pHOS in a linear model:

$$
\begin{equation*}
\frac{R_{i}}{S_{i}}=\beta_{0}+\beta_{1} p H O S_{i}+\epsilon_{i}, \tag{12}
\end{equation*}
$$

where $R_{i}$ is the observed number of recruits for brood year $i, S_{i}$ is the observed number of redds for brood year $i, \beta_{0}$ and $\beta_{1}$ are regression coefficients, and $\epsilon_{i} \sim N\left(0, \sigma_{r}^{2}\right)$. The estimated regression coefficient $\hat{\beta}_{1}$ was tested against 0 using a one-sided t-test. If $\hat{\beta}_{1}$ was significantly $<0(\mathrm{P}<0.05)$ (equivalently, significant negative Pearson correlation coefficient), we rejected hypothesis H 02.2 .1 .2 and concluded there was evidence that hatchery fish may be reducing the productivity of the wild population. Modeling assumptions were that the error terms $\epsilon_{i}(i=1, \ldots, n)$ are independent normal random variables with mean 0 and common variance $\left(\sigma_{r}^{2}\right)$. The observed error terms from the fitted model in Equation (12) were examined for non-normality using quantilequantile plots and Shapiro-Wilk tests (Shapiro and Wilk 1965). Heteroscedasticity was examined using plots of $\hat{\epsilon}_{i}$ against fitted values of $R_{i} / S_{i}$. In the event of heteroscedasticity and non-normal errors, $R_{i} / S_{i}$ was log-transformed in Equation (12).

The approach described above assumes that there is no density dependence in recruits per redd, that is, that any density dependence observed in analysis of hypothesis H02.2.1.1 occurs during spawning rather than during incubation and rearing. This assumption was assessed by inspection of recruits per redd compared to redd counts: a negative association (i.e., negative Pearson correlation coefficient) was interpreted as evidence of density dependence after spawning. In this event, an additional assessment was implemented in which we modeled recruits per redd as a function of both redd counts and pHOS using the Beverton-Holt model in Equation (5) and the Ricker model in Equation (7), using redd counts in place of spawner counts. A negative association between pHOS and juveniles per redd was assessed by the $95 \%$ bootstrap confidence interval for $\beta_{2}$ using at least 1,000 bootstrap samples. In the event that bootstrapping failed, asymptotic (normal theory) confidence intervals and correlation coefficients were provided. If the $95 \%$ confidence interval was entirely $>0$, it was concluded that there was a negative association between
pHOS and juveniles per redd, and that hatchery fish may result in lowered juvenile productivity of naturally spawning fish.

## Power Analysis

The statistical power was computed to detect a reduction in mean juvenile recruitment with Type I error probability of $\alpha=0.10$ when the actual reduction in mean recruitment associated with a change in the pHOS level was $5 \%$. Power was computed at varying sample sizes, where sample size $=$ number of years of observations, and both with and without measurement error in the number of spawners and pHOS. The stock-recruitment model was a Ricker model with parameters taken from the Chiwawa River spring Chinook Salmon population. The Chiwawa population was selected because it has the longest time series of stock and juvenile recruitment data of the populations available and the stock recruitment curves have fit the juvenile productivity from this population relatively well (Buchanan and Townsend 2021). Power was calculated for sampling sizes from $\mathrm{N}=5$ to $\mathrm{N}=100$ years for varying levels of pHOS effect size, $\beta_{-}$P. (see Appendix B for details on the power analysis).

## Results

## Spring Chinook Salmon

Chiwawa River Spring Chinook salmon adult spawner, smolt recruitment, pHOS , and redd count data were available for the 1991-2017 brood years. Adult stock abundance (spawners) ranged from 33 in 1995 to 2,032 in 2011 and averaged 723, and smolt recruitment ranged from 3,830 in 1995 to 82,845 in 2002 and averaged 36,318. Redd counts averaged 328.2 (range $=13$ to 1,078 ), and pHOS averaged 0.508 (range $=0.00$ to 1.00 ).

Nason Creek Spring Chinook data were available for the 2002-2017 brood years. Spawner abundance ranged from 132 (2017) to 702 (2011) and averaged 398, and smolt recruitment ranged from 930 (2014) to 8,696 (2005) and averaged 4,774. Redd counts averaged 176.9 (range $=68$ to 413 ), and pHOS averaged 0.599 (range $=0.20$ to 0.86 ). The analysis was performed with and without the low smolt count of 930 from the 2014 brood year.

White River Spring Chinook data were available for the 2005-2017 brood years. Spawner abundance ranged from 31 (2017) to 173 (2009) and averaged 94, and smolt recruitment ranged from $580(2014)$ to $11,170(2016)$ and averaged 5,079. Redd counts averaged 44.1 (range $=16$ to 86 ), and pHOS averaged 0.412 (range $=0.00$ to 0.75 ). The analysis was performed with and without the low smolt count of 580 from the 2014 brood year.

Twisp River Spring Chinook data were available for the 2003-2017 brood years. Spawner abundance ranged from 43 (2003) to 341 (2004) and averaged 165, and smolt recruitment ranged from 900 (2003) to 16,415 (2006) and averaged 5,295. Redd counts averaged 79.3 (range $=18$ to 145 ), and pHOS averaged 0.52 (range $=0.28$ to 0.76 ).

Methow River Spring Chinook salmon data were available for the 2002-2017 brood years. Spawner abundance ranged from 417 (2017) to 2,692 (2011) and averaged 1,417, and smolt recruitment ranged from 5,163 (2007) to 51,325 (2010) and averaged 23,624. Redd counts averaged 705 (range $=210$ to 1,366 ), and pHOS averaged 0.77 (range $=0.57$ to 0.97 ).

## Hypothesis $\mathrm{HO}_{2.2 .1 .1}$ Effect of pHOS on Recruitment:

Negative linear relationships between spawners and smolt recruitment indicated the presence of density dependence in mortality in one or more life stages between spawner data collection and smolt data collection in the Chiwawa, Nason, and Twisp populations. Weak negative linear relationships were observed between spawners and smolt recruitment in the White and Methow, but were non-significant, providing inconclusive evidence of density dependence in mortality between the spawning and smolt life stages. Without the 2014 brood year, density dependent mortality was detected in the White River population as well (Figure 1).


Figure 1. Smolts per spawner (log scale) versus spawner abundance for each population, with fitted linear regression line, Pearson correlation coefficient (r), and P-value from one-sided t-test of negative slope. Plots with and without the 2014 brood year (possible outlier) are shown for Nason and White.

The Smooth Hockey Stick, Beverton-Holt, and Ricker stock-recruitment models were attempted to be fit to the spawner and smolt data for each population. All three models were successfully fit to the data from the Chiwawa, White, Twisp, and Methow populations (Figure 2). Only the Ricker model could be fit to the Nason Creek data (Figure 3). For the Chiwawa, White, Twisp, and Methow populations, there was little difference in the model fits among the three models, demonstrated by the fitted recruitment curves, AICc, and correlation coefficients between the model parameters (Figure 2; Table 2). There was high uncertainty in the model fits for the White, Twisp, and Methow populations, as seen by the wide confidence intervals on parameter
estimates (Table 2). For Nason Creek, the pattern of observed spawner and smolt data did not support the assumption of asymptotic growth in recruitment as spawner abundance increases that is required by the Smooth Hockey Stick and Beverton-Holt models, and neither of those models could be fit to the Nason Creek data. The Ricker model was fit to Nason Creek data with and without the 2014 brood year (Figure 3); when the 2014 brood year was included, the model assumption of lognormal errors was not supported.


Figure 2. Stock-recruitment models fit to smolt and spawner data for Spring Chinook salmon populations (plot for White population includes 2014 brood year).

Table 2. Fitted stock-recruitment models for smolts and stock (spawner) data for each Spring Chinook salmon population. Confidence intervals were estimated using bootstrap with at least 1,000 bootstrap samples. Correlation $=$ Pearson correlation coefficient between parameter estimates from bootstrap samples. Model equations are defined in Equations (1), (2), and (3). $\Delta \mathrm{AICc}$ represents change in AICc compared to model with minimum AICc for the population. (Results include 2014 brood year for all populations). *Only the Ricker model could be fit to the Nason Creek data.

| Population | Model* | $\Delta$ AICc | Parameter $^{\mathrm{a}}$ | Parameter Estimates | $95 \% \mathrm{CI}$ | Correlation |
| :--- | :--- | :---: | :---: | :---: | :---: | :---: |
| Chiwawa | Smooth Hockey Stick | 0.00 | $\alpha$ | 168 | $118.2-248.7$ | $\alpha, \mathrm{R}_{\infty}:-0.39$ |
|  |  |  | $\mathrm{R}_{\infty}$ | 44,090 | $35,095-56,732$ |  |
|  | Beverton-Holt | 0.75 | $\alpha$ | 54,825 | $40,678-77,093$ | $\alpha, \beta: 0.87$ |
|  |  |  | $\beta$ | 278 | $134.5-559.0$ |  |
|  | Ricker | 0.71 | $\alpha$ | 141 | $109.2-184.3$ | $\alpha, \beta: 0.80$ |
|  |  |  | $\beta$ | 0.0011 | $0.0008-0.0014$ |  |


| Population | Model* | $\triangle \mathrm{AICc}$ | Parameter ${ }^{\text {a }}$ | Parameter Estimates | 95\% CI | Correlation |
| :---: | :---: | :---: | :---: | :---: | :---: | :---: |
| Nason | Smooth Hockey Stick | NA | K | 47,760 | 40,825-56,520 | $\alpha, \mathrm{R}_{\infty}$ : NA |
|  |  |  | $\alpha$ | NA | NA |  |
|  |  |  | $\mathrm{R}_{\infty}$ | NA | NA |  |
|  | Beverton-Holt | NA | $\alpha$ | NA | NA | $\alpha, \beta$ : NA |
|  |  |  | $\beta$ | NA | NA |  |
|  | Ricker | NA | $\alpha$ | 51.1 | 25.7-89.2 | $\alpha, \beta: 0.89$ |
|  |  |  | $\beta$ | 0.0037 | 0.0022-0.0052 |  |
|  |  |  | K | 5,144 | 3,710-7,011 |  |
| White | Smooth Hockey Stick | 0.08 | $\alpha$ | 79.7 | 44.7-262.1 | $\alpha, \mathrm{R}_{\infty}:-0.24{ }^{\text {b }}$ |
|  |  |  | R | 7,183 | 3,516-60,989 |  |
|  | Beverton-Holt | 0.14 | $\alpha$ | 11,011 | 5,286-185,892 | $\alpha, \beta: 0.96$ |
|  |  |  | $\beta$ | 130 | $32-3,591$ |  |
|  | Ricker | 0.00 | $\alpha$ | 78.9 | $35.5-180.4$ | $\alpha, \beta: 0.90$ |
|  |  |  | $\beta$ | 0.0049 | $<0.0001-0.0137$ |  |
|  |  |  | K | 5,632 | 4,571-9,125 |  |
| Twisp | Smooth Hockey Stick | 0.31 | $\alpha$ | 51.2 | 30.7-146.7 | $\alpha, \mathrm{R}_{\infty}:-0.20^{\text {c }}$ |
|  |  |  | $\mathrm{R}_{\infty}$ | 7,104 | 4,344-36,313 |  |
|  | Beverton-Holt | 0.62 | $\alpha$ | 11,295 | 6,317-101,201 | $\alpha, \beta: 0.99$ |
|  |  |  | $\beta$ | 216 | $75-3,103$ |  |
|  | Ricker | 0.00 | $\alpha$ | 51.2 | 29.7-93.6 | $\alpha, \beta: 0.88$ |
|  |  |  | $\beta$ | 0.0032 | <0.0001-0.0064 |  |
|  |  |  | K | 5,926 | $4,267-\infty$ |  |
| Methow | Smooth Hockey Stick | $0.02$ | $\alpha$ | 19.7 | 14.7 - 37.7 | $\alpha, \mathrm{R}_{\infty}:-0.30^{\mathrm{d}}$ |
|  |  |  | $\mathrm{R}_{\infty}$ | 65,051 | 23,577-549,603 |  |
|  | Beverton-Holt | 0.04 | $\alpha$ | 115,394 | $\begin{aligned} & 32,755- \\ & 1,382,406 \end{aligned}$ | $\alpha, \beta$ : 0.99 |
|  |  |  | $\beta$ | 5,806 | 785-82,314 |  |
|  | Ricker | 0.00 | $\alpha$ | 19.6 | 11.6-32.7 | $\alpha, \beta: 0.88$ |
|  |  |  | $\beta$ | 0.0001 | <0.0001-0.0005 |  |
|  |  |  | K | 50,572 | 22,637-m |  |

$\mathrm{a}=$ Estimated maximum recruitment parameters are: $R_{\infty}$ for the Smooth Hockey Stick model, $\alpha$ for the BevertonHolt model, and $K=(\alpha / \beta) e^{-1}$ for the Ricker model.
$\mathrm{b}=$ Correlation coefficient between $\alpha$ and $R_{\infty}$ is uninformative because parameter estimates were inversely related. The correlation coefficient between $1 / \alpha$ and $R_{\infty}=0.6241$.
$\mathrm{c}=$ Correlation coefficient between $\alpha$ and $R_{\infty}$ is uninformative because parameter estimates were inversely related. The correlation coefficient between $1 / \alpha$ and $R_{\infty}=0.4951$.
$\mathrm{d}=$ Correlation coefficient between $\alpha$ and $R_{\infty}$ is uninformative because parameter estimates were inversely related.
The correlation coefficient between $1 / \alpha$ and $R_{\infty}=0.7281$.


Figure 3. Stock and smolt data with fitted LOESS curve and Ricker Model (with and without data from 2014) for Nason Creek Spring Chinook Salmon, 2002-2017.

The residuals from all fitted models were compared to pHOS for each population using correlation analysis. Correlation coefficients between residuals and pHOS were not significantly different from 0 for any population (Figure 4). Additionally, likelihood ratio tests of the effect of pHOS on the Beverton-Holt and Ricker models were non-significant for each population (Table 3). Thus, there was no evidence to suggest that higher proportions of hatchery spawners were associated with lower juvenile productivity for spring Chinook Salmon in the populations. When the 2014 brood year was removed from the Nason Creek and White River data sets, model residuals were again uncorrelated with $\mathrm{pHOS}(\mathrm{r}=0.1614, \mathrm{P}=0.5655$ for Nason Creek; $\mathrm{r}=-0.0184$ to $0.0599, \mathrm{P} \geq 0.8533$ for White River) and likelihood ratio tests of the effect of pHOS remained insignificant (Table 3).

There were a number of limitations in the data and analyses for the majority of the populations of Spring Chinook salmon for hypothesis $\mathrm{H}_{2 \text { 2.2.1.1. }}$. Only the Chiwawa population had as many as 20 years of data; the shorter time series available for the other populations provided lower statistical power to detect an effect of pHOS. Modeling assumptions were not wellsupported by the data for some populations. In particular, the assumption of lognormal errors was not supported for Nason Creek or White River when the 2014 brood year was included in the analysis; in both cases, the lognormal assumption was better supported when 2014 was omitted. For the Twisp population, negative autocorrelation in the model residuals at a time lag of 3 years violated the assumption of non-stationarity and suggested that these stock-recruitment models do not adequately account for the population dynamics. For the White, Twisp, and Methow populations, model parameters had wide confidence intervals that included values far beyond the range of the observed data and parameter estimates were highly correlated; both these conditions reflect poor model fit and increase uncertainty in model predictions and the resulting residual analysis. These limitations preclude firm conclusions and it is possible that a relationship may exist between pHOS and juvenile productivity that was not observable using the available data and analyses.


Figure 4. Proportion Hatchery Origin Spawners verses Residuals from stock recruit model. Pvalue from two-sided t-test and Pearson correlation coefficient (r) are provided in the figure. The stock-recruitment model with the lowest AICc value for each population was chosen for this figure. Plots with the 2014 brood year (possible outlier) are shown for Nason and White.

Table 3. Results from likelihood ratio tests of the effect of pHOS on Beverton-Holt and Ricker Models. Value $=\mathrm{P}$-value from chi-squared test with $\mathrm{df}=1$.

| Population | Beverton-Holt | Ricker |
| :---: | :---: | :---: |
| Chiwawa | 0.9421 | 0.8181 |
| Nason | NA | $0.1420^{\mathrm{a}}$ |
| White | $0.6070^{\mathrm{b}}$ | $0.5518^{\mathrm{b}}$ |
| Twisp | 0.4524 | 0.5256 |
| Methow | 0.7635 | 0.7645 |

$\mathrm{a}=\mathrm{P}=0.5092$ when brood year 2014 was omitted for Nason Creek.
$\mathrm{b}=\mathrm{P}=0.9492$ for the Beverton-Holt model and $\mathrm{P}=0.8331$ for the Ricker model when brood year 2014 was omitted for White river.

## Hypothesis $\mathrm{HO}_{2.2 .1 .2}$ Effect of pHOS on Juveniles per Redd:

The relationship between average smolts per redd and pHOS was investigated for each Spring Chinook salmon population. A weak negative association was found between smolts per redd and pHOS for Chiwawa River Spring Chinook Salmon ( $\mathrm{P}=0.0319$; Figure 5). However, redd count was positively associated with $\mathrm{pHOS}(\mathrm{P}=0.0280)$ and there was a strong pattern of postspawning density dependence apparent from examination of the smolts per redd plotted against the redd counts (Figure 6). Thus, it is possible that the negative association between smolts per redd and pHOS may have resulted from the density dependence. The relationship between smolts per redd and pHOS was examined using the Ricker and Beverton-Holt stock-recruitment models in order to account for density dependence effects, with redds used in place of spawner counts. The Beverton-Holt model had considerably better fit than the Ricker model on the basis of AICc rank ( $\triangle \mathrm{AICc}=5.81$ compared to Ricker model) and examination of model residuals (not shown). The fitted Beverton-Holt model for smolts per redd was (Table 4):

$$
\ln (R / S)=\ln (55,479)-0.0343 P_{\text {HoS }}-\ln (117+S)
$$

The $95 \%$ bootstrap confidence interval for the effect of pHOS (i.e., $\beta_{2}$ in Equation (5)) was ( $-0.5443,0.6421$ ). Because this confidence interval included both positive and negative values, we concluded that there is no evidence of a negative association between pHOS and smolts per redd for Chiwawa Spring Chinook Salmon after adjusting for density dependence.

There was no evidence of a negative linear association between smolts per redd and the proportion of hatchery spawners ( pHOS ) in the other Spring Chinook salmon populations (Figure 5). There was evidence of post-spawning density dependence in Nason and Twisp (Figure 6) and of a linear association of redd count and pHOS in the Methow at the $10 \%$ level ( $\mathrm{P}=0.0919$ ). The relationship between smolts per redd and pHOS was examined using the Ricker and BevertonHolt stock-recruitment models in order to account for any density dependence effects, with redds used in place of spawner counts. In populations where models could be fit to the data, the $95 \%$ bootstrap confidence intervals for the effect of pHOS (i.e., $\beta_{2}$ ) included zero. Because the confidence intervals included both positive and negative values, we concluded that there is no evidence of a negative association between pHOS and smolts per redd for these populations after
adjusting for density dependence (Table 4). Results were unchanged whether or not the 2014 brood year was included for the Nason and White populations.

The analysis of the relationship of smolts per redd to pHOS suffered from the same limitations as the analysis of smolts versus pHOS (hypothesis H02.2.1.1). Sample sizes were small for all populations except the Chiwawa, and there was potentially large measurement error in the data for all populations. The modeling assumption of lognormal errors was not supported by the data for Nason Creek, parameter confidence intervals were wide for all models that could be fit for the Nason, White, and Twisp populations, and there was negative autocorrelation observed in the residuals for the Twisp data. Additionally, the Beverton-Holt model could not be fit for the Nason and Methow populations, and the estimated model parameter values were not consistent with their usual interpretation for the White, Twisp, and Methow data sets. These observations reflect poor model fit of the stock-recruitment models for the Nason, White, Twisp, and Methow populations, which results in lower confidence in the estimated model parameters and lower ability to detection any relationship between pHOS and smolts per redd using the available data.


Figure 5. Smolts per redd versus proportion of hatchery origin spawners ( pHOS ) for each population, with fitted linear regression line, Pearson correlation coefficient (r), and P-value from one-sided $t$-test of negative slope. Plots with and without the 2014 brood year (possible outlier) are shown for Nason and White.


Figure 6. Smolts per redd (log scale) versus redd count for each population, with fitted linear regression line, Pearson correlation coefficient ( r ), and P -value from one-sided t -test of negative slope. Negative slope indicates post-spawning density dependence. Plots with and without the 2014 brood year (possible outlier) are shown for Nason and White.

Table 1. Fitted stock-recruitment models for smolts per redd for each Spring Chinook salmon population, including effect ( $\beta_{2}$ ) of proportion of hatchery origin spawners ( pHOS ). Confidence intervals were estimated using bootstrap with at least 1,000 bootstrap samples. Model equations are defined in Equations (5) and (7). $\Delta \mathrm{AICc}$ represents change in AICc compared to model with minimum AICc for the population. (Results include 2014 brood year for all populations). *Only the Ricker model could be fit to the Nason Creek and Methow River data.

| Population | Model* | $\triangle \mathrm{AICc}$ | Parameter | Parameter Estimates | 95\% CI |
| :---: | :---: | :---: | :---: | :---: | :---: |
| Chiwawa | Beverton-Holt | 0.00 | $\alpha$ | 55,479 | 35,197-95,959 |
|  |  |  | $\beta$ | 117 | 57.9-242.2 |
|  |  |  | $\beta_{2}$ | 0.0343 | -0.5443-0.6421 |
|  | Ricker | 5.81 | $\ln (\alpha)$ | 5.7328 | $5.2646-6.2477$ |
|  |  |  | $\beta$ | 0.0021 | $0.0014-0.0032$ |
|  |  |  | $\beta_{2}$ | 0.0872 | -0.6880-0.9010 |
| Nason | Beverton-Holt | NA | $\alpha$ | NA | NA |
|  |  |  | $\beta$ | NA | NA |
|  |  |  | $\beta_{2}$ | NA | NA |
|  | Ricker | NA | $\ln (\alpha)$ | 3.8891 | 2.2758-4.8782 |
|  |  |  | $\beta$ | 0.0058 | $0.0033-0.0107$ |
|  |  |  | $\beta_{2}$ | -0.7001 | $-2.7076-0.8269$ |
| White | Beverton-Holt | 0.00 | $\alpha$ | 9,478 | 4,270-173,640 |
|  |  |  | $\beta$ | 48.1 | $12.9-1,315$ |
|  |  |  | $\beta_{2}$ | -0.0887 | -1.6125-1.6392 |
|  | Ricker | 0.10 | $\ln (\alpha)$ | 5.0979 | $3.5461-6.3336$ |
|  |  |  | $\beta$ | 0.0096 | -0.0080-0.0228 |
|  |  |  | $\beta_{2}$ | -0.0655 | $-2.2529-1.5695$ |
| Twisp | Beverton-Holt | 0.78 | $\alpha$ | 8,020 | 3,952-39,818 |
|  |  |  | $\beta$ | 103 | 40-724 |
|  |  |  | $\beta_{2}$ | -0.7782 | $-1.9437-0.4321$ |
|  | Ricker | 0.00 | $\ln (\alpha)$ | 4.3238 | $3.6525-5.0395$ |
|  |  |  | $\beta$ | 0.0063 | 0.0011-0.0108 |
|  |  |  | $\beta_{2}$ | -0.7621 | $-2.0433-0.1133$ |
| Methow | Beverton-Holt | NA | $\alpha$ | NA | NA |
|  |  |  | $\beta$ | NA | NA |
|  |  |  | $\beta_{2}$ | NA | NA |
|  | Ricker | NA | $\ln (\alpha)$ | 4.0380 | $2.6075-5.3023$ |
|  |  |  | $\beta$ | $<0.0001$ | < $0.0001-0.0006$ |
|  |  |  | $\beta_{2}$ | 0.7383 | -1.2657-2.3241 |

## Summer Chinook Salmon

Wenatchee River Summer Chinook adult spawner, emigrant, pHOS , and redd count data were available for the 1999-2018 brood years (no emigrant count for 2010 and 2011). Adult stock abundance (spawners) ranged from 3,473 in 2018 to 17,792 in 2006 and averaged 8,695. Emigrant counts ranged from 1,322,383 in 2000 to 20,426,149 in 2003 and averaged 9,118,268. Redd counts averaged $3,600.5$ (range $=1,510$ to 8,896 ), and pHOS averaged 0.176 (range $=0.06$ to 0.31 ). The 2000 and 2003 brood years were identified as possible outliers in emigrant counts, and analysis results were investigated for their sensitivity to inclusion of these brood years.

Methow River Summer Chinook data were available for the 2006-2018 brood years (no emigrant count for 2012). Spawner abundance ranged from 1,364 in 2007 to 3,952 in 2015 and averaged 2,333, and emigrant count ranged from 427,193 in 2017 to $3,465,247$ in 2006 and averaged $1,099,370$. Redd counts averaged 909.5 (range $=591$ to 1,551 ), and pHOS averaged 0.391 (range $=0.11$ to 0.53 ).

## Hypothesis $\mathrm{H0}_{2.2 .1 .1}$ Effect of pHOS on Recruitment:

A positive but non-significant linear relationship was observed between spawners and emigrant count in the Wenatchee population, indicating no evidence of density dependence in mortality between spawner data collection and emigrant data collection (Figure 7). In the Methow population, a slightly negative linear relationship between spawners and emigrant count suggested the presence of density dependence in mortality in one or more life stages between spawner data collection and emigrant data collection, but the relationship was not statistically significant ( $\mathrm{P}=0.4020$; (Figure 7).


Figure 7. Emigrants per spawner (log scale) versus spawner abundance for Wenatchee and Methow Rivers Summer Chinook Salmon, with fitted linear regression line, Pearson correlation coefficient ( r ), and P -value from one-sided t -test of negative slope.

An attempt was made to fit the Smooth Hockey Stick, Beverton-Holt, and Ricker stockrecruitment models to the spawner and emigrant count data from the Wenatchee Summer Chinook
salmon populations, but only the Ricker model could be fit when the full data were used (Figure 8; Table 5). The Ricker model residuals were not correlated with pHOS ( $\mathrm{r}=-0.0985, \mathrm{P}=0.6973$; Figure 9), and the likelihood ratio test of the effect of pHOS on the Ricker model was also nonsignificant $(\mathrm{P}=0.6787)$. When the 2003 brood year was omitted as a possible outlier, all three stockrecruitment models could be fit to the Wenatchee spawner and emigrant data and had common AICc weight ( $\triangle \mathrm{AICc} \leq 0.0072$ ); pHOS was not associated with the residuals from any of the models ( $\mathrm{P} \geq 0.7155$ ). When the 2000 brood year was omitted as a possible outlier, all three models could be fit to the data and the Ricker model was selected ( $\triangle \mathrm{AIC} \mathrm{c} \leq 0.1126$ ); model residuals were again not significantly associated with pHOS levels ( $\mathrm{P} \geq 0.6430$ ), and the likelihood ratio test of the pHOS was also not significant ( $\mathrm{P} \geq 0.6113$ ). Thus, there was no evidence to suggest that higher proportions of hatchery spawners were associated with lower juvenile productivity for Summer Chinook salmon in the Wenatchee River.

The Smooth Hockey Stick, Beverton-Holt, and Ricker stock-recruitment models were each fit to the spawner and emigrant data for the Methow (Figure 8). There was little difference in the model fits among the three models ( $\Delta \mathrm{AICc} \leq 0.02$ ), but the Ricker model had the lowest AICc value and intermediate correlation between parameter estimates (Table 5). The residuals from each of the three models were not correlated to pHOS ( $\mathrm{r}=-0.0059, \mathrm{P}=0.9856$ for each model; Figure 9). Additionally, likelihood ratio tests of the effect of pHOS on the Beverton-Holt and Ricker models were non-significant ( $\mathrm{P} \geq 0.9838$ ). There was no evidence to suggest that higher proportions of hatchery spawners were associated with lower juvenile productivity for Summer Chinook Salmon in the Methow River.


Figure 8. Stock-recruitment models fit to emigrant and spawner data for the Wenatchee and Methow River Summer Chinook Salmon. The Ricker stock-recruitment model is shown for the Wenatchee population.

Table 5. Fitted stock-recruitment models for emigrants and stock (spawner) data from Wenatchee River, 1999-2018 (without 2010, 2011) and Methow River, 2006-2018 (without 2012) Summer Chinook Salmon. Confidence intervals were estimated using bootstrap samples (at least 2,900 Wenatchee and 1,700 Methow). Correlation $=$ Pearson correlation coefficient between parameter estimates from bootstrap samples. Model equations are defined in Equations (1), (2), and (3). $\Delta$ AICc represents change in AICc compared to model with minimum AICc for each population. *Only the Ricker model could be fit to the Wenatchee River data.

|  |  |  | Parameter |  |  |  |
| :--- | :--- | :---: | :---: | :---: | :---: | :---: |
| Population | Model | $\Delta \mathrm{AICc}$ | Parameter $^{\mathrm{a}}$ | Estimates | $95 \% \mathrm{CI}$ | Correlation |
| Wenatchee | Smooth Hockey Stick |  | $\alpha$ | NA | NA | $\alpha, R_{\infty}:$ NA |
|  |  | $R_{\infty}$ | NA | NA |  |  |
|  | Beverton-Holt |  | $\alpha$ | NA | NA | $\alpha, \beta:$ NA |
|  |  |  | $\beta$ | NA | NA |  |
|  | Ricker |  | $\alpha$ | 957.3 | $773.4-1609.5$ | $\alpha, \beta: 0.83$ |
|  |  |  | $\beta$ | $<0.0001$ | $<0.0001-0.0001$ |  |
| Methow | Smooth Hockey Stick | 0.01 | $\alpha$ | $30,446,537$ | $919,213-\infty$ |  |
|  |  |  | $R_{\infty}$ | $4,796,000$ | $889,141-19,840,304$ |  |
|  | Beverton-Holt | 0.02 | $\alpha$ | $9,817,511$ | $874,870-66,701,078$ | $\alpha, \beta: 0.99$ |
|  |  |  | $\beta$ | 21,552 | $-26.7-16,434.1$ |  |
|  | Ricker | 0.00 | $\alpha$ | 461 | $331.4-1032.5$ | $\alpha, \beta: 0.88$ |
|  |  |  | $\beta$ | 0.00005 | $<0.0001-0.0004$ |  |

a = Estimated maximum recruitment parameters are: $R_{\infty}$ for the Smooth Hockey Stick model, $a$ for the Beverton-Holt model, and $K=(\alpha / \beta) e^{-1}$ for the Ricker model.
$\mathrm{b}=$ Correlation coefficient between $a$ and $R_{\infty}$ is uninformative because parameter estimates are not linearly related.


Figure 1. Residuals from Ricker Model versus proportion of hatchery origin spawners (pHOS) with fitted linear regression line, Pearson correlation coefficient (r), and P-value from two-sided $t$-test of slope of linear regression line using emigrant and spawner data from Wenatchee and Methow Rivers Summer Chinook Salmon. The Wenatchee results included the 2000 and 2003 brood years.

## Hypothesis $\mathrm{H0}_{2.2 \text {.1.2 }}$ Effect of pHOS on Juveniles per Redd:

There was a weak but non-significant negative association between emigrants per redd (log scale) and the proportion of hatchery spawners ( pHOS ) for both populations ( $\mathrm{P} \geq 0.3836$; Figure 10). The redd count was not associated with pHOS in either population and there was no indication of post-spawning density dependence apparent from examination of the emigrants per redd plotted against the redd counts ( $\mathrm{P} \geq 0.2944$, Figure 10). Nevertheless, the relationship between emigrants per redd and pHOS was examined using the Ricker and Beverton-Holt stock-recruitment models in order to account for possible density dependence effects, with redds used in place of spawner counts. Both models were equivalent on the basis of AICc rank ( $\Delta \mathrm{AICc} \leq 0.10$ ) and examination of model residuals for both populations (Table 6). However, for both populations, the Beverton-Holt model could not be fit for the bootstrap samples and the $95 \%$ asymptotic confidence intervals for the model parameters $\alpha$ and $\beta$ included both negative values and values far beyond the range of the observed data; these results indicate a lack of fit for the Beverton-Holt model for the two populations. Thus, the Ricker model was preferred. For the Wenatchee population, the fitted Ricker model for emigrants per redd was (Table 6):

$$
\ln (R / S)=8.0322-0.7166 P_{\text {HOS }}-0.00005 * S .
$$

The $95 \%$ bootstrap confidence interval for the effect of pHOS (i.e., $\beta_{2}$ in Equation (7)) was (-2.0006-3.3638). For the Methow population, the fitted Ricker model for emigrants per redd was:

$$
\ln (R / S)=7.0724-0.2448 P_{\text {HOS }}-0.00004 * S
$$

with a $95 \%$ bootstrap confidence interval $(-3.2640,2.4357)$ for the regression coefficient for pHOS. For both populations, the bootstrap confidence interval for the effect of pHOS included both positive and negative values, so we concluded that there is no evidence of a negative association between pHOS and emigrants per redd for Summer Chinook Salmon from either the Wenatchee River or the Methow River. For the Wenatchee population, removing 2000 or 2003 as possible outlier brood years did not significantly change the results.


Figure 10. Emigrants per redd (log scale) versus proportion of hatchery origin spawners (pHOS) (left plots), redd count versus proportion of hatchery origin spawners (pHOS) (middle plots), and emigrants per redd versus redd count (right plots) for Wenatchee and Methow River Summer Chinook Salmon, with fitted linear regression line, Pearson correlation coefficient (r), and P-value from t-test of slope (one-sided test of negative slope for left and right plots, two-sided test for middle plots).

Table 6. Fitted stock-recruitment models for emigrants per redd for Wenatchee River 1999-2018 (without 2010, 2011) and Methow River, 2006-2018 (without 2012), Summer Chinook Salmon, including effect $\left(\beta_{2}\right)$ of proportion of hatchery origin spawners (pHOS). Bootstrap confidence intervals based on 3,000 bootstrap samples (Wenatchee) and 1,000 bootstrap samples (Methow) were provided for the Ricker model, and asymptotic confidence intervals were provided for the Beverton-Holt model. Model equations are defined in Equations (5) and (7). $\Delta \mathrm{AICc}$ represents change in AICc compared to model with minimum AICc for each population.

| Population | Model | $\triangle \mathrm{AICc}$ | Parameter | Parameter Estimates | 95\% CI |
| :---: | :---: | :---: | :---: | :---: | :---: |
| Wenatchee | Beverton-Holt | 0.10 | $\alpha$ | 69,904,888 | - 259,329,930-399,139,700 |
|  |  |  | $\beta$ | 23,464 | -105,680-9,152,607 |
|  |  |  | $\beta_{2}$ | 0.6695 | -3.1075-4.4465 |
|  | Ricker | 0.00 | $\ln (\alpha)$ | 8.0322 | 6.8834-8.6882 |
|  |  |  | $\beta$ | 0.00005 | -0.0002-0.0001 |
|  |  |  | $\beta_{2}$ | 0.7166 | $-2.0006-3.3364$ |
| Methow | Beverton-Holt | 0.001 | $\alpha$ | 31,205,045 | - 881,383,950-943,794,030 |
|  |  |  | $\beta$ | 26,564 | -772,397-825,526 |
|  |  |  | $\beta_{2}$ | 0.2468 | -2.4111-2.9046 |
|  | Ricker | 0.000 | $\ln (\alpha)$ | 7.0724 | $4.6169-8.2810$ |
|  |  |  | $\beta$ | 0.00004 | -0.0016-0.0010 |
|  |  |  | $\beta_{2}$ | 0.2448 | -3.2640-2.4357 |

Summer Steelhead
Methow River Summer Steelhead adult spawner, emigrant, pHOS , and redd count data were available for the 2003-2015 brood years. Adult stock abundance (spawners) ranged from 1,105 in 2012 to 3,680 in 2010 and averaged 2,003, and emigrant recruitment ranged from 9,076 in 2003 to 33,739 in 2007 and averaged 18,154 . Redd counts averaged 1,084 (range $=591$ to 2,019 ), and pHOS averaged 0.81 (range $=0.58$ to 0.89 ). The 2007 brood year stood out as a possible outlier in emigrant count with a much higher count $(33,739)$ than in the other years (range $=9,076-25,845)$.

Twisp River Summer Steelhead adult spawner, emigrant recruitment, pHOS, and redd count data were available from for the 2003-2015 brood years. Spawners ranged from 143 in 2007 to 1,204 in 2003 and averaged 532, and emigrant recruitment ranged from 3,264 in 2008 to 13,669 in 2007 and averaged 6,133 . Redd counts averaged 272 (range $=82$ to 696), and pHOS averaged 0.69 (range $=0.48$ to 0.89 ). The 2007 brood year stood out as a possible outlier in emigrant count, with a much higher emigrant count $(13,669)$ than any of the other years (range $=3,264-7,467$ ). Additionally, pHOS generally declined over the course of the data collection, partly as a result of a policy change in 2010 to target pHOS values of 0.5 .

## Hypothesis $\mathrm{H0}_{2.2 .1 .1}$ Effect of pHOS on Recruitment:

A strong negative linear relationship was observed in both the Methow River and Twisp River between spawners and emigrant recruitment, demonstrating evidence of density dependence in mortality between the spawning and emigrant life stages (Figure 11). Thus, an attempt was made to fit the Smooth Hockey Stick, Beverton-Holt, and Ricker stock-recruitment models to the spawner and emigrant data for each population. However, the Smooth Hockey Stick and BevertonHolt modeling assumption of asymptotic growth in recruitment as spawner abundance increases was not supported by the data for either population, as demonstrated by the locally smoothed curves (LOESS) in Figure 12, and neither the Smooth Hockey Stick model nor the Beverton-Holt model could be fit to the spawner-recruitment data for these populations. The Ricker model was fit to the data for both populations, and estimated maximum recruitment at approximately 1,662 spawners for the Methow population and approximately 410 spawners for the Twisp population (Figure 12). Although the Ricker model could be fit to the spawner and emigrant data from both populations, in each case there were indications of poor model fit and possible bias. For the Methow data, the model residuals demonstrated negative autocorrelation at a time lag of 1 and 4 years, suggesting that the Ricker model does not adequately account for population dynamics. For the Twisp data, the model residuals did not clearly meet the assumption of lognormal errors. In each case, the model parameter estimates may be biased. Additionally, for both populations there was high correlation observed between the model parameter estimates, which increases uncertainty in the model predictions (Table 7).

For the Methow population, the estimated Pearson correlation coefficient between the Ricker model residuals and pHOS was 0.2222 and was not significantly different from 0 ( $\mathrm{P}=0.4656$; Figure 13). Additionally, a likelihood ratio test of the effect of pHOS on the Ricker model was non-significant $(\mathrm{P}=0.3829)$. The Beverton-Holt model could not be fit with pHOS as a variable. Analysis that omitted the possible outlier brood year of 2007 resulted in the same
findings: only the Ricker model was approximately consistent with the data, and there was no evidence of a negative relationship between pHOS and juvenile productivity of emigrants (Figure 14).

For the Twisp population, the estimated Pearson correlation coefficient between the Ricker model residuals and pHOS $(\mathrm{r}=0.2553)$ was not significantly different from $0(\mathrm{P}=0.3999$; Figure 13). A simpler analysis that compared the average Ricker model residuals between the pre-2010 period, when pHOS was not restricted ("control"), and the 2010+ period, when pHOS was targeted at 0.5 ("treatment"), also found no difference in model residuals between the two management periods $(t$-test $=-0.4717, \mathrm{df}=11, \mathrm{P}=0.6463$; Figure 15). Additionally, a likelihood ratio test of the effect of pHOS on the Ricker model was non-significant $(\mathrm{P}=0.3071)$. The Beverton-Holt model could not be fit with pHOS as a variable. When the 2007 brood year was removed as a possible outlier in emigrant count, the results were consistent with the full data set: a non-significant relationship between the Ricker model residuals and $\mathrm{pHOS}(\mathrm{P}=0.2868)$ and also between the Ricker model residuals and the pHOS management periods ( $\mathrm{P}=0.1728$ ), and the Beverton-Holt and Smooth Hockey Stick models could not be fit to the data (Figure 12, Table 7, Figure 14, Figure 15).

Overall, there was no evidence that to suggest that higher proportions of hatchery spawners were associated with lower juvenile productivity for Summer Steelhead in the Methow or Twisp rivers. However, there were only 13 years of data available for either population and the modeling assumptions were not well supported. Thus, it remains possible that there is a relationship between pHOS and juvenile productivity that is not observable using the available data and analyses.


Figure 11. Emigrants per spawner (log scale) versus spawner abundance for Methow and Twisp Rivers Summer Steelhead, with fitted linear regression line, Pearson correlation coefficient (r), and P -value from one-sided t -test of negative slope.


Figure 12. Stock and emigrant data with fitted LOESS curve and Ricker Model for Methow and Twisp Rivers Summer Steelhead.

Table 7. Fitted stock-recruitment models for emigrants and stock (spawner) data from Summer Steelhead in the Methow and Twisp rivers, 2003-2015. Confidence intervals were estimated using bootstrap with at least 2,000 bootstrap samples. Correlation $=$ Pearson correlation coefficient between parameter estimates from bootstrap samples. Model equations are defined in Equations (1), (2), and (3). $\Delta \mathrm{AICc}$ represents change in AICc compared to model with minimum AICc for each population (NA for all models because only one model fit per population). *Only the Ricker model could be fit to the data for Methow and Twisp rivers. The Ricker model was fit both with and without data from the 2007 brood year in the Twisp.

|  |  |  |  | Parameter |  |  |
| :---: | :---: | :---: | :---: | :---: | :---: | :---: |
| Population <br> Methow | Model* | $\triangle \mathrm{AICc}$ | Parameter ${ }^{\text {a }}$ | Estimates | 95\% CI | $\frac{\text { Correlation }}{\alpha, R_{\infty}: \text { NA }}$ |
|  | Smooth Hockey Stick | NA | $\alpha$ | NA | NA |  |
| Methow |  |  | $R_{\infty}$ | NA | NA |  |
|  | Beverton-Holt | NA | $\alpha$ | NA | NA | $\alpha, \beta$ NA |
|  |  |  | $\beta$ | NA | NA |  |
|  | Ricker with 2007 | NA | $\alpha$ | 30.8 | 19.3-48.9 | $\alpha, \beta: 0.90$ |
|  |  |  | $\beta$ | 0.0006 | 0.0004-0.0008 |  |
|  |  |  | $K$ | 18,835 | 15,830-23,693 |  |
|  | Ricker without 2007 | NA | $\alpha$ | 26.2 | 17.1-40.7 | $\alpha, \beta: 0.91$ |
|  |  |  | $\beta$ | 0.0005 | 0.0004-0.0007 |  |
|  |  |  | $K$ | 17,591 | 14,926-21,375 |  |
| Twisp | Smooth Hockey Stick | NA | $\alpha$ | NA | NA | $\alpha, R_{\infty}:$ NA |
|  |  |  | $R_{\infty}$ | NA | NA |  |
|  | Beverton-Holt | NA | $\alpha$ | NA | NA | $\alpha, \beta$ NA |
|  |  |  | $\beta$ | NA | NA |  |
|  | Ricker with 2007 | NA | $\alpha$ | 46.3 | 29.2-76.2 | $\alpha, \beta: 0.87$ |
|  |  |  | $\beta$ | 0.0024 | 0.0016-0.0032 |  |
|  |  |  | $K$ | 6,983 | 5,667-9,221 |  |
|  | Ricker without 2007 | NA | $\alpha$ | 31.8 | 22.9-43.1 | $\alpha, \beta: 0.90$ |
|  |  |  | $\beta$ | 0.0019 | 0.0015-0.0025 |  |
|  |  |  | $K$ | 6,054 | 5,247-6,930 |  |

$\mathrm{a}=$ Estimated maximum recruitment parameters are: $R_{\infty}$ for the Smooth Hockey Stick model, $a$ for the Beverton-Holt model, and $K=(\alpha / \beta) e^{-1}$ for the Ricker model.


Figure 13. Residuals from the Ricker Model versus proportion of hatchery-origin spawners (pHOS) with fitted linear regression line, Pearson correlation coefficient (r), and P-value from two-sided $t$-test of slope of linear regression line using emigrant and spawner data from Methow and Twisp rivers Summer Steelhead (including 2007).


Figure 2. Residuals from the Ricker Model versus proportion of hatchery-origin spawners ( pHOS ) with fitted linear regression line, Pearson correlation coefficient (r), and P-value from two-sided ttest of slope of linear regression line using emigrant and spawner data from Methow and Twisp rivers Summer Steelhead (without 2007).


Figure 15. Distribution of residuals from the Ricker Model categorized by pHOS management period: Control = pre-2010, when pHOS levels were not restricted, and Treatment $=2010-2015$, when pHOS levels were targeted at 0.5 . Data used were emigrant and spawner counts from Twisp River Summer Steelhead, 2003-2015. P-value is from two-sided t-test: t -test $=-0.4717$, $\mathrm{df}=11$ (with brood year 2007), and t-test $=1.4683, \mathrm{df}=10($ without brood year 2007 $)$.

## Hypothesis $\mathrm{H0}_{2.2 \text {.1.2 }}$ Effect of pHOS on Juveniles per Redd

There was no evidence of a negative association between emigrants per redd and the proportion of hatchery spawners ( pHOS ) for Summer Steelhead from either the Methow River or the Twisp River (Figure 16). The linear association between redd count and pHOS was nonsignificant ( $\mathrm{P}=0.2335$ ) in the Methow but significant at the $10 \%$ level ( $\mathrm{P}=0.0723$ ) in the Twisp (Figure 17). Comparison of emigrants per redd with redd count demonstrated strong evidence of post-spawning density dependence in both populations ( $\mathrm{P} \leq 0.0002$; Figure 18). Thus, the relationship between emigrants per redd and pHOS was examined using stock-recruitment models in order to account for density dependence effects, using redds in place of spawner abundance. In both populations the Beverton-Holt model resulted in a negative (i.e., inadmissible) estimate of the $\beta$ parameter, confirmed by examination of the likelihood surface. Thus, results are provided only for the Ricker model.

For the Methow population, the residuals from the Ricker model agreed moderately well with model assumptions. The fitted Ricker model for Methow River emigrants per redd was (Table 8):

$$
\ln (R / S)=3.3547+0.9921 * P_{\text {HOS }}-0.0012 * S
$$

with a $95 \%$ bootstrap confidence interval ( $-3.3407,2.5691$ ) for the effect of pHOS (i.e., $\beta_{2}$ in Equation (7)). Because the confidence interval for the effect of pHOS included both positive and negative values, we concluded that there was no evidence of a negative association between pHOS and emigrants per redd for Summer Steelhead from the Methow River whether or not an adjustment was made for density dependence. We reached the same conclusion when the possible outlier brood year (2007) was omitted from the data analysis (Figure 19, Table 8). However, the small sample size, low degree of variability in the observed pHOS values, and potentially high level of measurement error in the observed data may have lowered the ability to detect a relationship between pHOS and emigrants per redd.

For the Twisp population, the residuals from the Ricker model agreed poorly with the model assumption of lognormal errors. The fitted Ricker model for Twisp River emigrants per redd was (Table 8):

$$
\ln (R / S)=3.9305+0.7680 * P_{\text {HOS }}-0.0045 * S
$$

with a $95 \%$ bootstrap confidence interval of $(-2.3366,2.1443)$ for the effect of pHOS . Because the confidence interval for the effect of pHOS included both positive and negative values, we concluded that there was no evidence of a negative association between pHOS and emigrants per redd for Summer Steelhead from the Twisp River whether or not an adjustment was made for density dependence. However, the failure to meet the model assumptions demonstrates an overall lack of model fit. The small sample size and potentially high level of measurement error in the observed data may have contributed to the lack of model fit, and likewise may have lowered the ability to detect a relationship between pHOS and emigrants per redd using the available data.

When the potential outlier brood year of 2007 was omitted from the Twisp River data set, there was a negative association between emigrants per redd and pHOS when density dependence was not accounted for ( $\mathrm{P}=0.0009$, Figure 19). However, there was significant density dependent mortality observed between the redd count and emigrant count ( $\mathrm{P}<0.0001$, Figure 18), which may have accounted for the perceived association between emigrants per redd and pHOS. When analyzed using the Ricker model, which accounts for density dependence, the estimated effect of pHOS on the redd-emigrant relationship was non-significant, with a $95 \%$ bootstrap confidence interval of ( $-2.3915,1.0898$ ) (Table 8). Without the 2007 brood year, the Beverton-Holt model relating emigrants to redd counts was able to be fit when pHOS was in the model and pHOS was associated with lower emigrants per redd, demonstrated by the $95 \%$ bootstrap confidence interval for $\beta_{2}$ that was entirely greater than 0 (i.e., $(0.0700,1.9685)$; Table 8 ). However, the bootstrap confidence intervals for the other Beverton-Holt model parameters were notably wide: the confidence interval of the asymptotic maximum recruitment $(\alpha)$ included values 5 times the maximum recruitment observed, and the confidence interval for the redd counts predicted to generate half the maximum recruitment ( $\beta$ ) included negative values (Table 8). The apparently poor fit of the Beverton-Holt model raises questions about the validity of its finding of a statistically significant pHOS effect.

Overall, the evidence for a negative association in both populations between the proportion of hatchery origin spawners ( pHOS ) and juvenile productivity in emigrants was not strong. There was no such evidence for the Methow population or for the Twisp population when based on the full data set that included the 2007 brood year; however, in each case, one of the two stockrecruitment models could not be fit to the data because of misalignment of modeling assumptions with the data, and the data set was small (only 13 brood years). When the 2007 brood year was treated as an outlier and omitted from the Twisp analysis, a negative association was observed between emigrants per redd and pHOS , but it disappeared when density dependence was accounted for using the Ricker model. The alternative stock-recruitment model (Beverton-Holt) maintained an effect of pHOS but did not appear to fit the data well, indicating possible bias in the estimate of the pHOS effect. The small sample size available may have lowered the ability to detect an effect of pHOS on juvenile productivity. Additionally, a negative trend in pHOS from nearly 0.90 in 2003 to nearly 0.50 in 2015 raises the possibility that any pHOS effect actually masks a temporal trend in juvenile productivity.


Figure 16. Emigrants per redd (log scale) versus proportion of hatchery origin spawners (pHOS) for Methow and Twisp rivers Summer Steelhead, 2003-2015, with fitted linear regression line, Pearson correlation coefficient (r), and P-value from one-sided $t$-test of negative slope.


Figure 17. Redd count versus proportion of hatchery origin spawners (pHOS) for Methow River and Twisp River Summer Steelhead, 2003-2015, with fitted linear regression line, Pearson correlation coefficient (r), and P-value from two-sided t -test of slope.


Figure 18. Emigrants per redd (log scale) versus redd count for Methow River and Twisp River Summer Steelhead, 2003-2015, with fitted linear regression line, Pearson correlation coefficient (r), and P-value from one-sided t-test of negative slope. Negative slope indicates post-spawning density dependence.

Table 2. Fitted stock-recruitment models for emigrants per redd for Methow River and Twisp River Summer Steelhead, 2003-2015, including effect $\left(\beta_{2}\right)$ of proportion of hatchery origin spawners (pHOS). Confidence intervals were estimated using bootstrap with 3,000 bootstrap samples. Model equations are defined in Equations (5) and (7). $\triangle$ AICc represents change in AICc compared to model with minimum AICc for each population and data set (i.e., with or without 2007; NA if only one model fit). *Only the Ricker model could be fit for the Methow data or for the Twisp data when 2007 was included.

| Population | Model* | $\triangle \mathrm{AICc}$ | Parameter | Parameter Estimates | 95\% CI |
| :---: | :---: | :---: | :---: | :---: | :---: |
| Methow | Beverton-Holt | NA | $\alpha$ | NA | NA |
|  |  |  | $\beta$ | NA | NA |
|  |  |  | $\beta_{2}$ | NA | NA |
|  | Ricker (with 2007) | NA | $\ln (\alpha)$ | 3.3547 | 1.5579-6.1456 |
|  |  |  | $\beta$ | 0.0012 | 0.0006-0.0018 |
|  |  |  | $\beta_{2}$ | -0.9921 | -3.3407-2.5691 |
|  | Ricker (without 2007) | NA | $\ln (\alpha)$ | 3.7302 | 2.3844-6.8984 |
|  |  |  | $\beta$ | 0.0011 | 0.0005-0.0016 |
|  |  |  | $\beta_{2}$ | -0.2687 | -2.2563-3.9353 |
| Twisp | Beverton-Holt (with 2007) | NA | $\alpha$ | NA | NA |
|  |  |  | $\beta$ | NA | NA |
|  |  |  | $\beta_{2}$ | NA | NA |
|  | Ricker (with 2007) | NA | $\ln (\alpha)$ | 3.9305 | 3.3009-5.0401 |
|  |  |  | $\beta$ | 0.0045 | 0.0015-0.0067 |
|  |  |  | $\beta_{2}$ | -0.7680 | $-2.3366-2.1443$ |
| Twisp | Beverton-Holt (without 2007) | $0.24{ }^{\text {a }}$ | $\alpha$ | 12,008 | 5,158-40,066 |
|  |  |  | $\beta$ | 46.4 | -31.2-275.2 |
|  |  |  | $\beta_{2}$ | 0.9259 | 0.0700-1.9685 |
|  | Ricker (without 2007) | $0.00^{\text {a }}$ | $\ln (\alpha)$ | 4.5239 | 3.8033-5.1740 |
|  |  |  | $\beta$ | 0.0027 | 0.0007-0.0049 |
|  |  |  | $\beta_{2}$ | 1.0323 | -2.3915-1.0898 |

$\mathrm{a}=$ compare $\Delta \mathrm{AICc}$ values within Twisp modeling results without 2007 brood year.


Figure 19. Emigrants per redd versus proportion of hatchery origin spawners (pHOS) for Methow and Twisp rivers Summer Steelhead, 2003-2015, omitting brood year 2007, with fitted linear regression line, Pearson correlation coefficient (r), and P-value from one-sided t-test of negative slope.

## Power Analysis

Simulated statistical power to detect a negative effect of pHOS was consistently $\leq 0.32$ for all sample sizes considered ( $\mathrm{N} \leq 100$ years) and for all but the most extreme pHOS effect sizes considered (i.e., $\beta \_\mathrm{P}=-0.068$ to -0.205 ), both with and without measurement error (Figure 20). For the largest effect size ( $\beta$ P $=-0.513$ ), simulated power was as high as 0.54 for a sample size of $\mathrm{N}=$ 40 years without measurement error, and only slightly lower at 0.52 for $\mathrm{N}=40$ years when measurement error was incorporated into the simulations. Achieving power of at least 0.70 required as many as $\mathrm{N}=70$ years of data. Power greater than 0.80 required 90 years of data. These simulations defined detection of a negative effect of pHOS as a $90 \%$ bootstrap confidence interval that was entirely less than 0 .
(i) Without measurement error

(ii) With measurement error


Figure 20. Statistical power of detecting a negative effect of pHOS for various levels of pHOS effect size $\beta_{\mathrm{P}}$ with (i) and without (ii) measurement error in stock and recruitment data. Power based on 1,000 simulations of stock (spawners) and juvenile recruitment data using Ricker stockrecruitment model fit to data from Chiwawa spring Chinook Salmon, 1991-2017: $\mathrm{a}=138, \mathrm{~b}=$ $0.0011, \sigma=0.4386$ (equation (2)). Type I error probability $=0.10$. Measurement error (ii) was incorporated into simulated observations of spawner and recruitment data using coefficient of variation (CV): $\mathrm{CV}=0.07$ for HOS, $\mathrm{CV}=0.06$ for NOS, and $\mathrm{CV}=0.11$ for recruitment. Shaded region $=95 \%$ bootstrap confidence interval using 300 bootstrap samples from the simulated data.

## Discussion

Our investigation of whether a higher proportion of hatchery origin spawners (pHOS) may result in lowered juvenile productivity was largely inconclusive. For most populations studied, no evidence of an effect of pHOS was observed. For the single population that demonstrated a
possible negative effect of pHOS on juvenile productivity (Twisp River Summer Steelhead), the evidence was weak: when density dependence was accounted for, the perceived negative association between pHOS and juveniles per redd was no longer observed. Additionally, the significant result was observed only when a possible outlier in emigrant counts was omitted, which may not be justified depending on the cause of the unusual emigrant count for that brood year. Thus, even when some evidence of a negative effect of pHOS was found, the evidence was particularly weak.

The spring Chinook life history type that emigrates from tributaries and upper reaches to presumably rear in the main stem rivers was not analyzed explicitly. However, the life history type would be included as age 1 emigrants captured by the lower river rotary screw traps in the basin level analyses.

Multiple factors combined to lower the ability to detect an effect, should it exist. The most obvious factor was the small size of most of the data sets. Most of the populations had data for fewer than 20 brood years, resulting in lowered opportunity to observe the full range of variability in stock-recruitment dynamics in response to changes in hatchery proportion of spawners and lower statistical power to detect an effect. A complication in assessing the stock-recruitment data for an effect of pHOS is data quality. The stock-recruitment models all assume low levels of measurement error in both spawners or redd counts and smolts or emigrants. The higher the level of measurement error, the lower the ability to model the dynamics well enough to detect an effect. In some populations (e.g., Twisp River Steelhead, Methow River Steelhead), pHOS has declined almost consistently since the start of data collection, which means any perceived effect of pHOS would be entirely confounded with temporal changes in juvenile productivity. This risk is increased by the observational nature of the pHOS data.

Some data sets studied also demonstrated low contrast in the spawner or redd counts observed across brood years. The stock-recruitment models require observations at both low and high levels of spawners. This requirement means that low variability in spawner (or redd count) data reduces the quality of fit of the stock-recruitment models and in some cases makes them impossible to be fit altogether. A lack of data at low spawner levels may have lowered the ability to fit the Smooth Hockey Stick model, which requires data at low spawner levels to estimate the model parameter associated with depensation $(\alpha)$. Examination of the likelihood surface for the Smooth Hockey Stick model for some populations demonstrated moderate information in maximum recruitment but a complete lack of information in the $\alpha$ parameter, interpreted as the slope of the stock-recruitment curve at very low numbers of spawners (for example, see the flat contour curves in the dimension of the $\alpha$ parameter in Figure 21). Then again, several populations failed to demonstrate a stable maximum (or simply high) juvenile recruitment pattern at high levels of spawners, which is assumed by both the Smooth Hockey Stick and Beverton-Holt models. Again, examination of the likelihood surface in these cases indicated either a lack of information in model parameters or else model parameters optimized at inadmissible values (e.g., negative spawner counts necessary to generate half the asymptotic maximum recruitment, Figure 22). Although the Smooth Hockey Stick stock-recruitment model was the preferred analysis framework, it appeared poorly suited to the stock and juvenile recruitment data available for many populations.

For all three stock-recruitment models considered, there was often high uncertainty in model parameter estimates even when the models could be fit to the data. This was indicated by wide confidence intervals that often included maximum smolt or emigrant values far beyond the range of observed data and by high correlation in model parameter estimates. High correlation lowers the ability to distinguish between different model fits and increases uncertainty in the results; it arises from lack of contrast in the data or violation of modeling assumptions.

In most cases there was no evidence of a time lag in juvenile recruitment per spawner as represented by autocorrelation in the residuals from the stock-recruitment models. However, a negative autocorrelation was observed with a time lag of three years for Twisp River Spring Chinook Salmon and a time lag of one and four years for Methow River Summer Steelhead. Such negative autocorrelations may be observed when a high level of recruitment one year is habitually followed by a low level of recruitment after a consistent delay (e.g., 3 years). Such a pattern may be expected for populations with consistent age structure among spawners, paired with densitydependent mortality between spawning and juvenile recruit surveys. In such cases, the Smooth Hockey Stick, Beverton-Holt, and Ricker models are unlikely to adequately represent all important features of the population dynamics; an age-structured model may be required. The value of pHOS may also be affected by such autocorrelation. Thus, such factors should be accounted for in order to detect a true effect of pHOS on juvenile productivity beyond age- and density-dependent components of the population dynamics.

Simulated power to detect a negative effect of pHOS on juvenile recruitment was low for all but the strongest effect sizes and for studies shorter than approximately 70 years. The low power values resulted from the high variability in residuals from the Ricker stock-recruitment model ( $\sigma=$ 0.4386 ) estimated from its fit to 27 years of data from the Chiwawa Spring Chinook salmon population (1991-2017).

The Chiwawa data set is the largest of the juvenile productivity data sets available for spring Chinook salmon and steelhead. The alterative data sets either failed to meet the modeling assumptions for the Ricker model (e.g., Methow and Twisp steelhead) or else had higher error variance about the fitted Ricker model (e.g., other spring Chinook salmon populations) than seen for the Chiwawa spring Chinook salmon population. Thus, it is expected that the power to detect a negative effect of pHOS on juvenile productivity would be highest for the Chiwawa spring Chinook population. Low power for this population would be compounded for the other populations with shorter time series or more complex population dynamics. As a consequence, it is unlikely that assessment of stock-recruitment curves similar to the Ricker model will be sufficient to detect a negative effect of pHOS in time to mediate any such effect. Other methods of monitoring and assessment are recommended to evaluate the effect of pHOS on juvenile productivity of anadromous salmonids in the upper Columbia River basin.

Given all the complicating factors identified here, only the most extreme effect of pHOS on juvenile productivity could have been detected. Thus, we caution against concluding that such an effect is truly absent. Future data collection efforts could be substantially improved by increasing the contrast in pHOS levels and including suitable in-basin unsupplemented references. More specifically, a planned experiment using deliberately chosen pHOS levels could be implemented in order to achieve the necessary contrast and remove confounding analyses with
temporal and age-structured processes. Appropriate power analyses must be performed on the study design prior to commencing work to ensure that a result may be obtained within an acceptable time frame.


Figure 21. Negative log-likelihood surface and contour curves for the Smooth Hockey Stick model of emigrants and spawner counts. Lighter regions indicate higher likelihood values (lower negative log-likelihood values). The estimated parameter values occur where the negative log-likelihood is minimized. Data set = Twisp River Summer Steelhead, 2003-2015.


Figure 22. Negative log-likelihood surface and contour curves for Beverton-Holt model of emigrants, redd counts, and pHOS , using $\boldsymbol{\beta}_{\mathbf{2}}=-0.1160$. Lighter regions indicate higher likelihood values (lower negative log-likelihood values). The estimated parameter values occur where the negative log-likelihood is minimized. The green triangle (falls below $b=0$ line) indicates the parameter set that optimizes the likelihood. Data set = Twisp River Summer Steelhead, 20032015.

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## Appendix A

Investigations into Association between Proportion of Hatchery Spawners and Juvenile Productivity for M\&E 2021 Update

## COLUMBIA BASIN RESEARCH

## Investigations into Association between Proportion of Hatchery Spawners and Juvenile Productivity for M\&E 2021 Update

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## Introduction

Conservation hatchery programs implemented in the Wenatchee and Methow river basins are intended to increase the abundance of the target populations. The strategy of the hatchery programs is to return hatchery adults that increase the spawning population, resulting in an increase in the number of offspring that will maintain and recover naturally reproducing populations. The assumption are that 1 ) increasing the number of spawners will increase the naturally reproducing population abundance in future generations, and 2) hatchery origin spawners have similar fitness to natural origin spawners and will not decrease the productivity of natural spawning fish. Assessment of the production of juvenile offspring prior to entering the ocean provides information on freshwater productivity and avoids ocean effects on the abundance and productivity of the population. However, there is a concern that the hatchery programs may lower the juvenile productivity in supplemented watersheds. The populations are managed under the concept of Proportionate Natural Influence (PNI; Hatchery Scientific Review Group, 2009), a framework that uses genetic modeling to estimate risk of hatchery introgression in natural populations and prescribes management benchmarks intended to allow hatchery programs to operate within an acceptable risk level to the natural population. PNI is composed of two metrics: Proportion of Hatchery Origin Spawners (pHOS; the proportion of hatchery fish in the naturally spawning population), and Proportion of Natural Origin Broodstock ( pNOB ; the proportion of natural origin fish in a hatchery broodstock). Part of ongoing monitoring and evaluation (M\&E) activities is to assess the relationship between pHOS and measures of juvenile productivity including the number of recruits to the smolt or emigrant populations and the average number of juveniles per redd. This report describes the statistical methods and results used to investigate these questions.

## Methods

The abundance of spawners and redds was assessed by conducting spawning ground surveys. The number of redds was tabulated and for semelparous species, carcass recoveries were used to estimate the proportions of males and females by origin (hatchery and natural origin). Summer steelhead biological data for gender and origin were obtained at sampling points (various traps). Abundance of semelparous spawners was estimated by expansion of redd counts based on the estimated portions of males to females, by origin, and assuming the number of redds created by a single female. Abundance of steelhead was estimated by adjusting dam and/or trap counts for proportions of fish estimated to return to the various tributaries in the Upper Columbia based on previous radio telemetry studies. Escapement estimates were adjusted to compensate for fall back and re-ascension at dams and to exclude fish removed from the spawning population through harvest and broodstock collection.

Juvenile abundance was estimated using rotary screw traps to sample migrating juveniles. Estimates were generated by conducting trap efficiency trials across a range of river flows, generating a regression model (flow versus trap efficiency), and using the model to predict the number of emigrants based on mean daily flow and daily captures. The daily estimates were summed for each year (season) to estimate the total abundance of emigrating juveniles per population. Juvenile steelhead were parsed into cohorts based on age determined through scales.

Spring chinook juveniles were yearlings assumed to be emigrating to the ocean. Age $0+$ spring Chinook migrants were not included in the analysis because their life history and survival are not well understood and we preferred to focus on emigrating fish when possible. Summer Chinook emigrants were subyearlings and assumed to be exhibiting an ocean-type life history. All age classes of steelhead from a cohort were included because of the difficulty in knowing how to parse emigrating fish from those that are not leaving
the freshwater system. Therefore steelhead include the entire range of observed age classes for each cohort. For the sake of simplicity, we refer to the steelhead juveniles caught in the screw traps as "emigrants."

Section 3.1 of the monitoring and evaluation plan for the hatchery programs (Hillman et al. 2019) addresses freshwater juvenile productivity and presents two questions, with additional hypotheses to be tested for each question:

Question 1: Has the supplementation program changed the number of juveniles (smolts, parr, and/or emigrants) per redd within the supplemented population?

Question 2: Does the number of juveniles per redd decrease as the proportion of hatchery spawners increases?

The first question could not be assessed because it requires reference populations, which are unavailable for the target populations. Furthermore, estimates of juvenile production are not available in a long enough time series to estimate the juvenile productivity before and after a hatchery program was implemented.

The second question was assessed in this report. The data and methods used to assess Question 2 are described below.

## Data

Douglas County PUD provided productivity data for the populations listed in Table 1. The size of the adult spawning stock, number of redds, and proportion of hatchery origin spawners ( pHOS ) were provided for each brood year, in addition to the number of smolts or emigrants. No data were available for Summer Steelhead from the Wenatchee River.

Table 1. Summary of spawner and recruitment data provided by Douglas County PUD. The size of the adult spawning stock, number of redds, and proportion of hatchery origin spawners ( pHOS ) were provided for each brood year, in addition to the number of smolts or emigrants.

| Species and Run | Population | Brood <br> Years | Smolts | Emigrants $^{\text {a }}$ | Comment |
| :--- | :--- | :---: | :---: | :---: | :--- |
| Spring Chinook | Chiwawa | $1991-2017$ | X |  |  |
| Spring Chinook | Nason | $2002-2017$ | X |  |  |
| Spring Chinook | White | $2005-2017$ | X |  |  |
| Spring Chinook | Twisp | $2003-2017$ | X |  |  |
| Spring Chinook | Methow | $2002-2017$ | X | X | No emigrant data: |
| Summer Chinook | Wenatchee | $1991-2018$ |  |  | 2010, 2011 |
| Summer Chinook | Methow | $2006-2018$ |  | X | No emigrant data: |
|  |  |  |  |  | 2012 |
| Summer Steelhead | Wenatchee |  |  | X |  |
| Summer Steelhead | Methow | $2003-2015$ |  | X |  |
| Summer Steelhead | Twisp | $2003-2015$ |  | N |  |

$\mathrm{a}=$ Emigrant data were provided for Spring Chinook Salmon populations from the Chiwawa, Methow, and Twisp rivers but were not used in the analysis.

## Statistical Methods

Spawner, redd count, and recruit data were analyzed together with proportion of hatchery spawners to address Objective 2 under Section 3: Juvenile Productivity of the Monitoring and Evaluation Plan for PUD Hatchery Programs: 2019 Update (Hillman et al. 2019):

Objective 2: Determine if the proportion of hatchery fish on the spawning grounds affects the freshwater productivity of supplemented stocks.

Monitoring Question Q2.2.1: Does the number of juveniles per redd decease as the proportion of hatchery spawners increases?

The following statistical hypotheses were investigated:
$\mathrm{H}_{2 \text { 2.2.1.1: }}$ : There is no association between the proportion of hatchery-origin spawners ( pHOS ) and the residuals from the smooth hockey stick stock-recruitment curve; $\rho=0$.
$\mathrm{H} 02_{2.2 .12}$ : The slope between proportion of hatchery spawners and juveniles/redd is $\geq 0$.
Three stock-recruitment models were used to account for density dependent mortality in assessment of the two hypotheses. The Smooth Hockey Stick model assumes that recruitment increases quickly at low levels of spawners and asymptotes to a maximum recruitment levels as spawners increase:

$$
\begin{equation*}
R=R_{\infty}\left(1-e^{-\left(\frac{\alpha}{R_{\infty}}\right) s}\right) \tag{1}
\end{equation*}
$$

where $\mathrm{R}=$ recruits and $\mathrm{S}=$ spawners for a given brood year, $R_{\infty}$ is the maximum number of recruits (population carrying capacity), and $\alpha$ is the slope at low levels of spawner abundance (Froese 2008).

The Beverton-Holt model also assumes asymptotic growth to a maximum:

$$
\begin{equation*}
R=\frac{\alpha S}{\beta+S} \tag{2}
\end{equation*}
$$

where $\mathrm{R}=$ recruits and $\mathrm{S}=$ spawners for a given brood year, $\alpha$ is the asymptotic maximum number of recruits (population carrying capacity), and $\beta$ is the predicted spawner abundance required to produce half the maximum recruits (Hilborn and Walters 1992).

The Ricker model assumes that recruitment increases at lower levels of spawners and declines at higher spawner abundance:

$$
\begin{equation*}
R=\alpha S e^{-\beta S} \tag{3}
\end{equation*}
$$

where $\mathrm{R}=$ recruits and $\mathrm{S}=$ spawners for a given brood year, $\alpha$ is the increase in recruits per spawner at low levels of spawners and $\beta$ is the intensity of the decrease in recruitment at high levels of spawner abundance (Ricker 1954). The maximum number of recruits is defined as $K=(\alpha / \beta) e^{-1}$.

Modeling assumptions were (based on Hillman et al. 2019):

1. Density-dependent mortality: The brood instantaneous mortality rate is proportional to the number of spawners for the brood year (Ricker 1954).
2. Lognormal errors: The variation in recruitment about the modeled recruitment for a given spawner abundance is lognormally distributed and acts multiplicatively (Quinn and Deriso 1999).
3. Negligible measurement error: Measurement error in spawning stock size and recruitment is small relative to the range of observed spawning stock sizes and variation in recruitment, respectively (Hilborn and Walters 1992).
4. Stationarity: The average stock-recruitment relationship is constant over time, with the possible exception of effects of changes in pHOS (Hilborn and Walters 1992).

The proportion of hatchery origin spawners may have the effect of lowering the maximum recruitment or recruits per spawner. The pHOS variable may be introduced into the Beverton-Holt model to reduce maximum recruitment when pHOS is high as follows:

$$
\begin{equation*}
R=\frac{\alpha e^{-\beta_{2} P_{\mathrm{HOS}} S}}{\beta+S} \tag{4}
\end{equation*}
$$

or equivalently

$$
\begin{equation*}
\ln (R / S)=\ln (\alpha)-\ln (\beta+S)-\beta_{2} P_{H O S} \tag{5}
\end{equation*}
$$

In Equations (4) and (5), the maximum number of recruits per brood year is $\alpha$ when pHOS is 0 and decreases asymptotically to $\alpha e^{-\beta_{2}}$ when pHOS is 1 .

The pHOS variable may be introduced into the Ricker model in such a way as to lower the slope in the spawner-recruitment curve at low levels of spawner abundance as follows:

$$
\begin{equation*}
R=\alpha e^{-\beta_{2} P_{H O S}} S e^{-\beta S} \tag{6}
\end{equation*}
$$

or equivalently

$$
\begin{equation*}
\ln (R / S)=\ln (\alpha)-\beta_{2} P_{H O S}-\beta S \tag{7}
\end{equation*}
$$

Alternatively, if higher levels of pHOS are expected to increase the intensity of the decrease in recruitment at high spawner levels, then pHOS may be incorporated as follows:

$$
\begin{equation*}
R=\alpha S e^{-\left(\beta+\beta_{2} P_{H O S}\right) S} \tag{8}
\end{equation*}
$$

or equivalently

$$
\begin{equation*}
\ln (R / S)=\ln (\alpha)-\left(\beta+\beta_{2} P_{H O S}\right) S \tag{9}
\end{equation*}
$$

Both models in Equation (6) and Equation (8) result in lower maximum recruitment for higher pHOS if $\beta_{2}>0$.

Hypothesis $\mathbf{H 0}_{2.2 .1 .1}$
Hypothesis $\mathrm{H}_{2 \text { 2.2.1. }}$ was investigated by fitting a stock-recruitment model to the available data and regressing the residuals from the model against pHOS. The Smooth Hockey Stick model, Beverton-Holt model, and Ricker model were considered.

Modeling assumption 1 was assessed by investigating the linear relationship between the log of recruits per spawner with the number of spawners; a negative relationship was consistent with density-dependent mortality. Modeling assumption 2 was assessed via quantile-quantile plots and Shapiro-Wilk tests (Shapiro and Wilk 1965) performed on the residuals from the fitted stock-recruitment model on the log scale. Modeling assumption 4 was assessed by plotting model residuals versus brood year and examining for autocorrelation. The available data did not allow for assessment of assumption 3; violation of assumption 3 would result in lower ability to detect a relationship between spawner abundance and recruitment (Hilborn and Walters 1992).

The fits of the three stock-recruitment models in Equations (1), (2), and (3) to the data were ranked using AICc (Burnham and Anderson 2002). Models with $\triangle \mathrm{AICc} \leq 2$ compared to the minimum observed AICc were used to assess the association between residuals and pHOS. The Smooth Hockey Stick model was included regardless of its AICc rank. Additionally, bias, uncertainty, and correlation in estimates of model parameters were estimated using bootstrapping with at least 1,000 bootstrap samples. Point estimates, $95 \%$ bootstrap confidence intervals, and bootstrap correlation coefficients between model parameters were reported for each model. In the event that bootstrapping failed, asymptotic (normal theory) confidence intervals and correlation coefficients were provided. Bias and correlation were considered in assessment of model fit.

For a given model, the residual for brood year $i(i=1, \ldots, n)$ was defined as

$$
\begin{equation*}
r_{i}=R_{i}-\hat{R}_{i}, \tag{10}
\end{equation*}
$$

where $R_{i}$ is the observed number of recruits for brood year $i$, and $\hat{R}_{i}$ is the predicted number of recruits from the model for brood year $i$. The association between the residuals $r_{i}(i=1, \ldots, n)$ and pHOS was investigated using the Pearson correlation coefficient and visual inspection of the fitted linear model

$$
\begin{equation*}
r_{i}=\beta_{0}+\beta_{1} p \operatorname{HOS}_{i}+\epsilon_{i}, \tag{11}
\end{equation*}
$$

for regression coefficients $\beta_{0}$ and $\beta_{1}$ and random error $\epsilon_{i} \sim N\left(0, \sigma_{r}^{2}\right)$. Hypothesis H02.2.1.1 was rejected if the Pearson correlation coefficient between $r_{i}$ and pHOS was negative and significantly different from 0 ( $\mathrm{P}<0.05$ ). In this event, we concluded that there was evidence that hatchery fish may be reducing the productivity of the wild population.

An additional assessment of the effect of pHOS on juvenile productivity was performed by fitting the pHOS-enhanced stock-recruitment models in Equations (4), (6), and (8) and testing whether $\beta_{2}=0$ using likelihood ratio tests.

Hypothesis $\mathbf{H} 0_{2.2 .1 .2}$
Hypothesis H02.2.1.2 was investigated in two ways. First, the ratio of recruits per redd was modeled using pHOS in a linear model:

$$
\begin{equation*}
\frac{R_{i}}{S_{i}}=\beta_{0}+\beta_{1} p H O S_{i}+\epsilon_{i} \tag{12}
\end{equation*}
$$

where $R_{i}$ is the observed number of recruits for brood year $i, S_{i}$ is the observed number of redds for brood year $i, \beta_{0}$ and $\beta_{1}$ are regression coefficients, and $\epsilon_{i} \sim N\left(0, \sigma_{r}^{2}\right)$. The estimated regression coefficient $\hat{\beta}_{1}$ was tested against 0 using a one-sided t-test. If $\hat{\beta}_{1}$ was significantly $<0$ ( $\mathrm{P}<0.05$ ) (equivalently, significant negative Pearson correlation coefficient), we rejected hypothesis H02.2.1.2 and concluded there was evidence that hatchery fish may be reducing the productivity of the wild population. Modeling assumptions were that the error terms $\epsilon_{i}(i=1, \ldots, n)$ are independent normal random variables with mean 0 and common variance ( $\sigma_{r}^{2}$ ). The observed error terms from the fitted model in Equation (12) were examined for non-normality using quantile-quantile plots and Shapiro-Wilk tests (Shapiro and Wilk 1965). Heteroscedasticity was examined using plots of $\hat{\epsilon}_{i}$ against fitted values of $R_{i} / S_{i}$. In the event of heteroscedasticity and non-normal errors, $R_{i} / S_{i}$ was log-transformed in Equation (12).

The approach described above assumes that there is no density dependence in recruits per redd, that is, that any density dependence observed in analysis of hypothesis H02.2.1.1 occurs during spawning rather than during incubation and rearing. This assumption was assessed by inspection of recruits per redd compared to redd counts: a negative association (i.e., negative Pearson correlation coefficient) was interpreted as evidence of density dependence after spawning. In this event, an additional assessment was implemented in which we modeled recruits per redd as a function of both redd counts and pHOS using the Beverton-Holt model in Equation (5) and the Ricker model in Equation (7), using redd counts in place of spawner counts. A negative association between pHOS and juveniles per redd was assessed by the $95 \%$ bootstrap confidence interval for $\beta_{2}$ using at least 1,000 bootstrap samples. In the event that bootstrapping failed, asymptotic (normal theory) confidence intervals and correlation coefficients were provided. If the $95 \%$ confidence interval was entirely $>0$, it was concluded that there was a negative association between pHOS and juveniles per redd, and that hatchery fish may result in lowered juvenile productivity of naturally spawning fish.

## Results

## Chiwawa River Spring Chinook Salmon

Adult spawner, smolt recruitment, pHOS , and redd count data were available for the 1991-2017 brood years. Adult stock abundance (spawners) ranged from 33 in 1995 to 2,032 in 2011 and averaged 723, and smolt recruitment ranged from 3,830 in 1995 to 82,845 in 2002 and averaged 36,318 (Figure 1, Figure 2). Redd counts averaged 328.2 (range $=13$ to 1,078 ), and pHOS averaged 0.508 (range $=0.00$ to 1.00 ).


Figure 1. Chiwawa River Spring Chinook Salmon boxplots for Stock (spawner abundance), Smolts, Redd counts, and proportion of hatchery origin spawners (pHOS), 1991-2017. Box indicates interquartile range and thick horizontal bar is median. Whiskers indicate quartile $\pm 1.5 \times$ interquartile range.


Figure 2. Chiwawa River Spring Chinook Salmon annual Smolt counts, Smolt/Spawner (log scale), Smolt/Redd (log scale), and proportion of hatchery origin spawners (pHOS), 1991-2017. Line indicates the linear trend. Shaded area is the estimated $95 \%$ confidence interval of the line.

## Hypothesis H02.2.1.1

A negative linear relationship between spawners and smolt recruitment indicated the presence of density dependence in mortality in one or more life stages between spawner data collection and smolt data collection (Figure 3). Thus, the Smooth Hockey Stick, Beverton-Holt, and Ricker stock-recruitment models were each fit to the spawner and smolt data (Figure 4). There was little difference in the model fits among the three models ( $\Delta \mathrm{AICc} \leq 0.75$ ), but the Smooth Hockey Stock model had the lowest AICc value and the lowest correlation between parameter estimates (Table 2). The residuals from all three models were compared to pHOS using correlation analysis. Correlation coefficients between residuals and pHOS ranged from -0.0132 to 0.0396 ; none was significantly different from 0 ( $\mathrm{P} \geq 0.8445$; Figure 5-Figure 7). Additionally, likelihood ratio tests of the effect of pHOS on the Beverton-Holt and Ricker models were non-significant ( $\mathrm{P} \geq 0.8181$ ). Thus, there was no evidence to suggest that higher proportions of hatchery spawners were associated with lower juvenile productivity for spring Chinook Salmon in the Chiwawa River.


Figure 3. Smolts per spawner (log scale) versus spawner abundance for Chiwawa River Spring Chinook Salmon, 1991-2017, with fitted linear regression line, Pearson correlation coefficient (r), and P-value from one-sided t -test of negative slope.


Figure 4. Stock-recruitment models fit to smolt and spawner data for Chiwawa River Spring Chinook Salmon, 1991-2017.

Table 2. Fitted stock-recruitment models for smolts and stock (spawner) data from Chiwawa River Spring Chinook Salmon, 1991-2017. Confidence intervals were estimated using bootstrap with 3,000 bootstrap samples. Correlation $=$ Pearson correlation coefficient between parameter estimates from bootstrap samples. Model equations are defined in Equations (1), (2), and (3). $\Delta \mathrm{AICc}$ represents change in AICc compared to model with minimum AICc.

| Model | AICc | $\triangle \mathrm{AICc}$ | Parameter ${ }^{\text {a }}$ | Parameter Estimates | 95\% CI | Correlation |
| :---: | :---: | :---: | :---: | :---: | :---: | :---: |
| Smooth Hockey Stick | 35.32 | 0.00 | $\alpha$ | 168 | 118.2-248.7 | $\alpha, R_{\infty}:-0.39$ |
|  |  |  |  |  |  |  |
|  |  |  | $R_{\infty}$ | 44,090 | 35,095-56,732 |  |
| Beverton-Holt | 36.07 | 0.75 | $\alpha$ | 54,825 | 40,678-77,093 | $\alpha, \beta: 0.87$ |
|  |  |  | $\beta$ | 278 | 134.5-559.0 |  |
| Ricker | 36.03 | 0.71 | $\alpha$ | 141 | 109.2-184.3 | $\alpha, \beta: 0.80$ |
|  |  |  | $\beta$ | 0.0011 | 0.0008-0.0014 |  |
|  |  |  | $K$ | 47,760 | 40,825-56,520 |  |

$\mathrm{a}=$ Estimated maximum recruitment parameters are: $R_{\infty}$ for the Smooth Hockey Stick model, $\alpha$ for the BevertonHolt model, and $K=(\alpha / \beta) e^{-1}$ for the Ricker model.


Figure 5. Residuals from Smooth Hockey Stick Model versus proportion of hatchery origin spawners ( pHOS ) with fitted linear regression line, Pearson correlation coefficient ( r ), and P -value from two-sided t test of slope of linear regression line using smolt and spawner data from Chiwawa River Chinook Salmon, 1991-2017. $\mathrm{R}^{2}=0.0008$.


Figure 6. Residuals from Beverton-Holt Model versus proportion of hatchery origin spawners ( pHOS ) with fitted linear regression line, Pearson correlation coefficient (r), and P-value from two-sided t-test of slope of linear regression line using smolt and spawner data from Chiwawa River Chinook Salmon, 1991-2017. $\mathrm{R}^{2}=0.0002$.


Figure 7. Residuals from Ricker Model versus proportion of hatchery origin spawners ( pHOS ) with fitted linear regression line, Pearson correlation coefficient ( r ), and P-value from two-sided t-test of slope of linear regression line using smolt and spawner data from Chiwawa River Chinook Salmon, 1991-2017. $\mathrm{R}^{2}=0.0016$.

## Hypothesis H02.2.1.2

There was a weak negative association between smolts per redd and the proportion of hatchery spawners (pHOS) for Chiwawa River Spring Chinook Salmon (Pearson correlation coefficient $\mathrm{r}=-0.3616, \mathrm{P}=0.0319$ ) (Figure 8). However, redd count was positively associated with $\mathrm{pHOS}(\mathrm{P}=0.0280$, Figure 9 ) and there was a strong pattern of post-spawning density dependence apparent from examination of the smolts per redd plotted against the redd counts (Figure 10). Thus, it is possible that the negative association between smolts per redd and pHOS may have resulted from the density dependence. The relationship between smolts per redd and pHOS was examined using the Ricker and Beverton-Holt stock-recruitment models in order to account for density dependence effects, with redds used in place of spawner counts. The Beverton-Holt model had considerably better fit than the Ricker model on the basis of AICc rank ( $\triangle \mathrm{AICc}=5.81$ compared
to Ricker model) and examination of model residuals (Table 3; Figure A9 to Figure A13). The fitted Beverton-Holt model for smolts per redd was (Table 3):

$$
\ln (R / S)=\ln (55,479)-0.0343 P_{\text {Hо八 }}-\ln (117+S) .
$$

The $95 \%$ bootstrap confidence interval for the effect of pHOS (i.e., $\beta_{2}$ in Equation (5)) was ( -0.5443 , 0.6421 ). Because this confidence interval included both positive and negative values, we concluded that there is no evidence of a negative association between pHOS and smolts per redd for Chiwawa Spring Chinook Salmon after adjusting for density dependence.


Figure 8. Smolts per redd (log scale) versus proportion of hatchery origin spawners (pHOS) for Chiwawa River Spring Chinook Salmon, 1991-2017, with fitted linear regression line, Pearson correlation coefficient (r), and P-value from one-sided t-test of negative slope. $\mathrm{R}^{2}=0.1256$.


Figure 9. Redd count versus proportion of hatchery origin spawners ( pHOS ) for Chiwawa River Spring Chinook Salmon, 1991-2017, with fitted linear regression line, Pearson correlation coefficient (r), and Pvalue from two-sided $t$-test of slope.


Figure 10. Smolts per redd versus redd count for Chiwawa River Spring Chinook Salmon, 1991-2017, with fitted linear regression line, Pearson correlation coefficient (r), and P-value from one-sided t-test of negative slope. Negative slope indicates post-spawning density dependence.

Table 3. Fitted stock-recruitment models for smolts per redd for Chiwawa River Spring Chinook Salmon, 1991-2017, including effect ( $\beta_{2}$ ) of proportion of hatchery origin spawners (pHOS). Confidence intervals were estimated using bootstrap with 3,000 bootstrap samples. Model equations are defined in Equations (5) and (7). $\Delta \mathrm{AICc}$ represents change in AICc compared to model with minimum AICc.

| Model | AICc | $\Delta$ AICc | Parameter | Parameter Estimates | $95 \% \mathrm{CI}$ |
| :--- | :--- | :--- | :---: | :---: | :---: |
| Beverton-Holt | 37.72 | 0.00 | $\alpha$ | 55,479 | $35,197-95,959$ |
|  |  |  | $\beta$ | 117 | $57.9-242.2$ |
|  |  |  | $\beta_{2}$ | 0.0343 | $-0.5443-0.6421$ |
|  |  |  |  |  |  |
| Ricker | 43.54 | 5.81 | $\ln (\alpha)$ | 5.7328 | $5.2646-6.2477$ |
|  |  |  | $\beta$ | 0.0021 | $0.0014-0.0032$ |
|  |  |  | $\beta_{2}$ | 0.0872 | $-0.6880-0.9010$ |

## Nason Creek Spring Chinook Salmon

Adult spawner, smolt recruitment, pHOS , and redd count data were available for the 2002-2017 brood years. Adult stock abundance (spawners) ranged from 132 in 2017 to 702 in 2011 and averaged 398, and smolt recruitment ranged from 930 in 2014 to 8,696 in 2005 and averaged 4,774 (Figure 11, Figure 12). Redd counts averaged 176.9 (range $=68$ to 413), and pHOS averaged 0.599 (range $=0.20$ to 0.86 ). The analysis was performed with and without the low smolt count of 930 from the 2014 brood year.


Figure 11. Nason Creek Spring Chinook Salmon boxplots for Stock (spawner abundance), Smolts, Redd counts, and proportion of hatchery origin spawners (pHOS), 2002-2017. Box indicates interquartile range and thick horizontal bar is median. Whiskers indicate quartile $\pm 1.5 \times$ interquartile range.


Figure 12. Nason Creek Spring Chinook Salmon annual Smolt counts, Smolt/Spawner (log scale), Smolt/Redd (log scale), and proportion of hatchery origin spawners (pHOS), 2002-2017. Line indicates the linear trend. Shaded area is the estimated $95 \%$ confidence interval of the line.

## Hypothesis H02.2.1.1

A negative linear relationship between spawners and smolt recruitment indicated the presence of density dependence in mortality in one or more life stages between spawner data collection and smolt data collection (Figure 13). The low smolt count from 2014 had little effect on the detection of density dependence. An attempt was made to fit the Smooth Hockey Stick, Beverton-Holt, and Ricker stockrecruitment models to the spawner and smolt data. However, the Smooth Hockey Stick and Beverton-Holt modeling assumption of asymptotic growth in recruitment as spawner abundance increases was not supported by the data, as demonstrated by the locally smoothed curve (LOESS) in Figure 14, and neither the Smooth Hockey Stick model nor the Beverton-Holt model could be fit for the Nason Creek spawnerrecruitment data. The Ricker model was fit to the Nason Creek data and demonstrated a maximum predicted recruitment at around 274 spawners (Figure 14), but the residuals demonstrated violation of modeling assumption 2 (lognormal errors) (Figure B1). The residuals from the Ricker model were compared to pHOS using correlation analysis. The estimated Pearson correlation coefficient between the Ricker model residuals and pHOS was 0.3317 and was not significantly different from 0 ( $\mathrm{P}=0.2095$; Figure 15). Additionally, a likelihood ratio test of the effect of pHOS on the Ricker model was non-significant, ( $\mathrm{P}=0.1420$ ). The Beverton-Holt model could not be fit with pHOS as a variable.

The Ricker model was refit without data from year 2014, the year that generated the very large negative residual apparent in Figure 14. Residuals from this model fit appeared to meet the modeling assumption of lognormal errors (Figure B3). The correlation between the Ricker model residuals and pHOS remained
non-significant even without the 2014 brood year data ( $\mathrm{r}=0.1614, \mathrm{P}=0.5655$; Figure 16). A likelihood ratio test of the effect of pHOS on the Ricker model was non-significant ( $\mathrm{P}=0.5092$ ) when the 2014 brood year was excluded. Even excluding data from 2014, the Smooth Hockey Stick model and the BevertonHolt model could not be fit for the Nason Creek data set.

Overall, there was no evidence to suggest that higher proportions of hatchery spawners were associated with lower juvenile productivity for spring Chinook Salmon in the Nason Creek. However, the data did not satisfy the modeling assumptions well, and there may be a relationship that was not observable using the current analysis methods and data quality.


Figure 13. Smolts per spawner (log scale) versus spawner abundance for Nason Creek Spring Chinook Salmon, 2002-2017, with and without brood year 2014, with fitted linear regression line, Pearson correlation coefficient ( r ), and P -value from one-sided t -test of negative slope.


Figure 14. Stock and smolt data with fitted LOESS curve and Ricker Model (with and without data from 2014) for Nason Creek Spring Chinook Salmon, 2002-2017.

Table 4. Fitted stock-recruitment models for smolts and stock (spawner) data from Nason Creek Spring Chinook Salmon, 2002-2017. Confidence intervals were estimated using bootstrap with at least 3,000 bootstrap samples. Correlation $=$ Pearson correlation coefficient between parameter estimates from bootstrap samples. Model equations are defined in Equations (1), (2), and (3). $\Delta \mathrm{AICc}$ represents change in AICc compared to model with minimum AICc. *Only the Ricker model could be fit for Nason Creek data. The Ricker model was fit both with and without data from the 2014 brood year. AICc should not be compared between the two Ricker models.

| Model* | AICc | $\triangle \mathrm{AICc}$ | Parameter ${ }^{\text {a }}$ | Parameter Estimates | 95\% CI | Correlation |
| :---: | :---: | :---: | :---: | :---: | :---: | :---: |
| Smooth Hockey Stick | NA | NA | $\alpha$ | NA | NA | $\alpha, R_{\infty}:$ NA |
|  |  |  | $R_{\infty}$ | NA | NA |  |
| Beverton-Holt | NA | NA | $\alpha$ | NA | NA | $\alpha, \beta$ : NA |
|  |  |  | $\beta$ | NA | NA |  |
| Ricker with 2014 | 35.52 | NA | $\alpha$ | 51.1 | 25.7-89.2 | $\alpha, \beta: 0.89$ |
|  |  |  | $\beta$ | 0.0037 | 0.0022-0.0052 |  |
|  |  |  | K | 5,144 | 3,710-7,011 |  |
| Ricker without 2014 | 20.44 | NA | $\alpha$ | 72.1 | 44.7-108.1 | $\alpha, \beta: 0.90$ |
|  |  |  | $\beta$ | 0.0042 | 0.0032-0.0005 |  |
|  |  |  | $K$ | 6,286 | 4,942-7,995 |  |

$\mathrm{a}=$ Estimated maximum recruitment parameters are: $R_{\infty}$ for the Smooth Hockey Stick model, $\alpha$ for the BevertonHolt model, and $K=(\alpha / \beta) e^{-1}$ for the Ricker model.


Figure 15. Residuals from the Ricker model versus proportion of hatchery origin spawners ( pHOS ) with fitted linear regression line, Pearson correlation coefficient (r), and P-value from two-sided t-test of slope of linear regression line using smolt and spawner data from Nason Creek Chinook Salmon, 2002-2017 (including brood year 2014). $\mathrm{R}^{2}=0.1100$.


Figure 16. Residuals from the Ricker model versus proportion of hatchery origin spawners ( pHOS ) with fitted linear regression line, Pearson correlation coefficient (r), and P-value from two-sided t-test of slope of linear regression line using smolt and spawner data from Nason Creek Chinook Salmon, 2002-2017 without brood year 2014. $\mathrm{R}^{2}=0.0261$.

## Hypothesis H02.2.1.2

There was no evidence of a negative association between smolts per redd and the proportion of hatchery spawners ( pHOS ) for Nason Creek Spring Chinook Salmon (Pearson correlation coefficient $\mathrm{r}=-0.1011$, $\mathrm{P}=0.3547$ ) (Figure 17). Additionally, the linear association between redd count and pHOS was nonsignificant ( $\mathrm{P}=0.2278$; Figure 18). Nevertheless, there was weak evidence of post-spawning density dependence based on comparison of smolts per redd versus redd counts using data from all brood years, including 2014 ( $\mathrm{P}=0.0049$; Figure 19). Thus, the relationship between smolts per redd and pHOS was examined using the Ricker stock-recruitment model in order to account for density dependence effects, using redds in place of spawner abundance. Similar to the modeling using spawner abundance, the Beverton-Holt model could not be fit using redd counts.

The fitted Ricker model (including brood year 2014) was (Table 5):

$$
\ln (R / S)=3.8891+0.7001 P_{\text {HOS }}-0.0058 S
$$

The $95 \%$ bootstrap confidence interval for the effect of pHOS (i.e., $\beta_{2}$ in Equation (7)) was ( -2.7076 , $0.8269)$. When the 2014 brood year was omitted, the regression model was

$$
\ln (R / S)=4.4479+0.0084 P_{\text {HOS }}-0.0060 S
$$

and the $95 \%$ bootstrap confidence interval for the effect of pHOS was ( $-1.6065,1.1149$ ).
Whether or not the 2014 brood year was included, the $95 \%$ confidence interval included both positive and negative values for the effect of pHOS , providing no evidence of a negative association between pHOS and smolts per redd for Nason Creek Chinook Salmon after adjusting for density dependence. However, analysis of residuals from the Ricker model with and without 2014 demonstrated poor agreement with the assumption of lognormal errors and poor model fit (Figures B6, B8), which is also demonstrated by the wide $95 \%$ bootstrap confidence intervals for the model parameters (Table 5). This pattern results in lower
confidence in the estimated model parameters and lowered ability to detect a relationship between pHOS and smolts per redd using the available data.


Figure 17. Smolts per redd versus proportion of hatchery origin spawners (pHOS) for Nason Creek Spring Chinook Salmon, 2002-2017, including brood year 2014, with fitted linear regression line, Pearson correlation coefficient (r), and P-value from one-sided t-test of negative slope. $\mathrm{R}^{2}=0.0102$.


Figure 18. Redd count versus proportion of hatchery origin spawners ( pHOS ) for Nason Creek Spring Chinook Salmon, 2002-2017, including brood year 2014, with fitted linear regression line, Pearson correlation coefficient (r), and P-value from two-sided t -test of slope.


Figure 19. Smolts per redd (log scale) versus redd count for Nason Creek Spring Chinook Salmon, 20022017, with and without 2014, with fitted linear regression line, Pearson correlation coefficient (r), and Pvalue from one-sided t-test of negative slope. Negative slope indicates post-spawning density dependence.

Table 5. Fitted stock-recruitment models for smolts per redd for Nason Creek Spring Chinook Salmon, 2002-2017 (including brood year 2014), including effect $\left(\beta_{2}\right)$ of proportion of hatchery origin spawners (pHOS). Confidence intervals were estimated using bootstrap with 3,000 bootstrap samples. Model equations are defined in Equations (5) and (7). $\triangle \mathrm{AICc}$ represents change in AICc compared to model with minimum AICc. *Only the Ricker model could be fit for Nason Creek data. The Ricker model was fit both with and without data from the 2014 brood year. AICc should not be compared between the two Ricker models.

| Model* | AICc | $\triangle \mathrm{AICc}$ | Parameter | Parameter Estimates | 95\% CI |
| :---: | :---: | :---: | :---: | :---: | :---: |
| Beverton-Holt | NA | NA | $\alpha$ | NA | NA |
|  |  |  | $\beta$ | NA | NA |
|  |  |  | $\beta_{2}$ | NA | NA |
| Ricker with 2014 | 41.22 | NA | $\ln (\alpha)$ | 3.8891 | 2.2758-4.8782 |
|  |  |  | $\beta$ | 0.0058 | $0.0033-0.0107$ |
|  |  |  | $\beta_{2}$ | -0.7001 | $-2.7076-0.8269$ |
| Ricker without 2014 | 33.64 | NA | $\ln (\alpha)$ | 4.4479 | $3.1832-5.0293$ |
|  |  |  | $\beta$ | 0.0060 | 0.0037-0.0100 |
|  |  |  | $\beta_{2}$ | -0.0084 | -1.6065-1.1149 |

## White River Spring Chinook Salmon

Adult spawner, smolt recruitment, pHOS, and redd count data were available for the 2005-2017 brood years. Adult stock abundance (spawners) ranged from 31 in 2017 to 173 in 2009 and averaged 94, and smolt recruitment ranged from 580 in 2014 to 11,170 in 2016 and averaged 5,079 (Figure 20, Figure 21). Redd counts averaged 44.1 (range $=16$ to 86 ), and pHOS averaged 0.412 (range $=0.00$ to 0.75 ).


Figure 20. White River Spring Chinook Salmon boxplots for Stock (spawner abundance), Smolts, Redd counts, and proportion of hatchery origin spawners (pHOS), 2005-2017. Box indicates interquartile range and thick horizontal bar is median. Whiskers indicate quartile $\pm 1.5 \times$ interquartile range.


Figure 21. White River Spring Chinook Salmon annual Smolt counts, Smolt/Spawner (log scale), Smolt/Redd (log scale), and proportion of hatchery origin spawners (pHOS), 2005-2017. Line indicates the linear trend. Shaded area is the estimated $95 \%$ confidence interval of the line.

## Hypothesis H02.2.1.1

A weak negative linear relationship was observed between White River spring Chinook Salmon spawners and smolt recruitment, but it was non-significant, providing inconclusive evidence of density dependence in mortality between the spawning and smolt life stages ( $\mathrm{P}=0.1550$; Figure 22 ). The very low smolt count from 2014 influenced the perceived relationship; without the 2014 brood year, there was a strong negative relationship between spawners and smolt recruitment ( $\mathrm{P}=0.0084$ ). Thus, the Smooth Hockey Stick, Beverton-Holt, and Ricker stock-recruitment models were fit to the spawner and smolt data.

When the 2014 brood year was included in the analysis, there was essentially no difference in the fits among the three stock-recruitment models (Figure 23, Table 6). Correlation coefficients between residuals from those models and pHOS ranged from 0.1408 to 0.1600 ; none was significantly different from 0 ( $\mathrm{P} \geq 0.6015$; Figure 24 - Figure 26). Additionally, likelihood ratio tests of the effect of pHOS on the Beverton-Holt and Ricker models were non-significant ( $\mathrm{P} \geq 0.5518$ ). However, tests of the modeling assumption of lognormal errors were inconclusive for all three models (Figures C1, C7, and C11), and model parameters were highly correlated (Table 6, Figure C3). Additionally, the $95 \%$ bootstrap confidence intervals for maximum recruitment included smolt recruitment levels far beyond the maximum observed smolt recruitment of 11,170 . Thus, results based on these three stock-recruitment models should be interpreted with caution.

Removing the 2014 brood year from the analysis had little effect on the comparisons among the three models (Figure 27, Table 7), although the residuals were more consistently lognormally distributed. There remained high correlation in the model parameters observed from the bootstrap samples (Table 7, Figure C6). The $95 \%$ bootstrap confidence intervals again included maximum recruitment estimates beyond the observed data, but they were considerably lower than when 2014 was included. Nevertheless, even without the 2014 brood year, the residuals were not correlated with pHOS for any of the three models ( $\mathrm{P} \geq 0.8533$; Figure 28-Figure 30). Likelihood ratio tests of the effect of pHOS on the Beverton-Holt and Ricker models were also non-significant ( $\mathrm{P} \geq 0.7763$ ).

Overall, there was no evidence to suggest that higher proportions of hatchery spawners were associated with lower juvenile productivity for spring Chinook Salmon in the White River. However, there were only 13 years of data including the 2014 brood year, and some of the modeling assumptions were not well supported. It is possible that there is a relationship between pHOS and juvenile productivity that was not observable using the available data.


Figure 22. Smolts per spawner (log scale) versus spawner abundance for White River Spring Chinook Salmon, 2005-2017, with and without brood year 2014, with fitted linear regression line, Pearson correlation coefficient (r), and P-value from one-sided $t$-test of negative slope.

White Spring Chinook Salmon


Figure 23. Stock-recruitment models fit to smolt and spawner data for White River Spring Chinook Salmon, 2005-2017, including brood year 2014.

Table 6. Fitted stock-recruitment models for smolts and stock (spawner) data from White River Spring Chinook Salmon, 2005-2017, including brood year 2014. Confidence intervals were estimated using bootstrap with at least 1,000 bootstrap samples. Correlation $=$ Pearson correlation coefficient between parameter estimates from bootstrap samples. Model equations are defined in Equations (1), (2), and (3). $\Delta \mathrm{AICc}$ represents change in AICc compared to model with minimum AICc.
$\left.\begin{array}{lclcccc}\hline \text { Model } & \text { AICc } & \Delta \text { AICc } & \text { Parameter }^{\mathrm{b}} & \begin{array}{c}\text { Parameter } \\ \text { Estimates }\end{array} & 95 \% \text { CI } & \text { Correlation } \\ \hline \text { Smooth Hockey } & 34.06 & 0.08 & \alpha & 79.7 & 44.7-262.1 & \alpha, R_{\infty}:-0.24^{\mathrm{a}} \\ \text { Stick } & & & R_{\infty} & 7,183 & 3,516-60,989 & \\ & & & & & 11,011 & 5,286-185,892\end{array}\right) \alpha, \beta: 0.96$
$\mathrm{a}=$ Correlation coefficient between $\alpha$ and $R_{\infty}$ is uninformative because parameter estimates are not linearly related; see Figure C3. The correlation coefficient between $1 / \alpha$ and $R_{\infty}=0.6241$.
$\mathrm{b}=$ Estimated maximum recruitment parameters are: $R_{\infty}$ for the Smooth Hockey Stick model, $\alpha$ for the Beverton-
Holt model, and $K=(\alpha / \beta) e^{-1}$ for the Ricker model.


Figure 24. Residuals from Smooth Hockey Stick Model versus proportion of hatchery origin spawners ( pHOS ) with fitted linear regression line, Pearson correlation coefficient ( r ), and P -value from two-sided t test of slope of linear regression line using smolt and spawner data from White River Spring Chinook Salmon, 2005-2017, including brood year 2014. $\mathrm{R}^{2}=0.0223$.


Proportion Hatchery Origin Spawners (pHOS)
Figure 25. Residuals from the Beverton-Holt Model versus proportion of hatchery origin spawners ( pHOS ) with fitted linear regression line, Pearson correlation coefficient (r), and P-value from two-sided t-test of slope of linear regression line using smolt and spawner data from White River Spring Chinook Salmon, 2005-2017, including brood year 2014. $\mathrm{R}^{2}=0.0198$.


Figure 26. Residuals from the Ricker Model versus proportion of hatchery origin spawners ( pHOS ) with fitted linear regression line, Pearson correlation coefficient (r), and P-value from two-sided t-test of slope of linear regression line using smolt and spawner data from White River Spring Chinook Salmon, 20052017, including brood year 2014. $\mathrm{R}^{2}=0.0256$.

White Spring Chinook Salmon


Figure 27. Stock-recruitment models fit to smolt and spawner data for White River Spring Chinook Salmon, 2005-2017, without brood year 2014.

Table 7. Fitted stock-recruitment models for smolts and stock (spawner) data from White River Spring Chinook Salmon, 2005-2017, without brood year 2014. Confidence intervals were estimated using bootstrap with at least 3,000 bootstrap samples. Correlation $=$ Pearson correlation coefficient between parameter estimates from bootstrap samples. Model equations are defined in Equations (1), (2), and (3). $\Delta \mathrm{AICc}$ represents change in AICc compared to model with minimum AICc.

| Model | AICc | $\triangle \mathrm{AICc}$ | Parameter ${ }^{\text {b }}$ | Parameter Estimates | 95\% CI | Correlation |
| :---: | :---: | :---: | :---: | :---: | :---: | :---: |
| Smooth Hockey Stick | 20.72 | 0.60 | $\alpha$ | 142.8 | $73.0-364.0$ | $\alpha, R_{\infty}:-0.38^{\text {a }}$ |
|  |  |  | $R_{\infty}$ | 5,903 | 4,348-12,940 |  |
| Beverton-Holt | 20.97 | 0.85 | $\alpha$ | 7,320 | 4,574-24,921 | $\alpha, \beta: 0.99$ |
|  |  |  | $\beta$ | 38.4 | 0.08-340.1 |  |
| Ricker | 20.12 | 0.00 | $\alpha$ | 128 | 73.2-219.4 | $\alpha, \beta: 0.90$ |
|  |  |  | $\beta$ | 0.0084 | 0.0032-0.0135 |  |
|  |  |  | $K$ | 5,632 | 4,571-9,125 |  |

$\overline{\mathrm{a}}=$ Correlation coefficient between $\alpha$ and $R_{\infty}$ is uninformative because parameter estimates are not linearly related; see Figure C6. The correlation coefficient between $1 / \alpha$ and $R_{\infty}=0.5703$.
$\mathrm{b}=$ Estimated maximum recruitment parameters are: $R_{\infty}$ for the Smooth Hockey Stick model, $\alpha$ for the BevertonHolt model, and $K=(\alpha / \beta) e^{-1}$ for the Ricker model.


Figure 28. Residuals from Smooth Hockey Stick Model versus proportion of hatchery origin spawners ( pHOS ) with fitted linear regression line, Pearson correlation coefficient ( r ), and P -value from two-sided t test of slope of linear regression line using smolt and spawner data from White River Spring Chinook Salmon, 2005-2017, without brood year 2014. $\mathrm{R}^{2}=0.0001$.


Figure 29. Residuals from the Beverton-Holt Model versus proportion of hatchery origin spawners ( pHOS ) with fitted linear regression line, Pearson correlation coefficient (r), and P-value from two-sided t-test of slope of linear regression line using smolt and spawner data from White River Spring Chinook Salmon, 2005-2017, without brood year 2014. $\mathrm{R}^{2}=0.0003$.


Figure 30. Residuals from the Ricker Model versus proportion of hatchery origin spawners ( pHOS ) with fitted linear regression line, Pearson correlation coefficient (r), and P-value from two-sided t-test of slope of linear regression line using smolt and spawner data from White River Spring Chinook Salmon, 20052017, without brood year 2014. $\mathrm{R}^{2}=0.0036$.

## Hypothesis H02.2.1.2

There was no evidence of a negative association between smolts per redd and the proportion of hatchery spawners ( pHOS ) for White River Spring Chinook Salmon (Pearson correlation coefficient $\mathrm{r}=-0.1873$, $\mathrm{P}=0.2700$ ) (Figure 31). Additionally, the linear association between redd count and pHOS was nonsignificant ( $\mathrm{P}=0.2795$; Figure 32). There was no evidence of post-spawning density dependence based on comparison of smolts per redd versus redd counts using data from all brood years, including 2014 ( $\mathrm{P}=0.1431$; Figure 33). However, when the 2014 brood year was omitted, density dependence was more
apparent $(\mathrm{P}=0.0109)$. Thus, the relationship between smolts per redd and pHOS was examined using the Ricker and Beverton-Holt stock-recruitment models in order to account for density dependence effects, using redds in place of spawner abundance. The model fits and assessment of residuals were very similar (Table 8, Figure C16 - Figure C23). The fitted Beverton-Holt model for smolts per redd was (including the 2014 brood year; Table 8):

$$
\ln (R / S)=\ln (9,478)+0.0887 P_{\text {HoS }}-\ln (48.1+S)
$$

The $95 \%$ bootstrap confidence interval for the effect of pHOS (i.e., $\beta_{2}$ in Equation (5)) was ( -1.6125 , 1.6392). The fitted Ricker model for smolts per redd was:

$$
\ln (R / S)=5.0979+0.0655 P_{\text {HOS }}-0.0096 S
$$

The $95 \%$ bootstrap confidence interval for the effect of pHOS (i.e., $\beta_{2}$ in Equation (7)) was ( -2.2529 , 1.5695). Similar results were observed when the models were fit without the 2014 brood year (Table 9).

Because the confidence intervals for the effect of pHOS included both positive and negative values regardless of the model and whether the 2014 brood year was included, we concluded that there was no evidence of a negative association between pHOS and smolts per redd for White River Spring Chinook Salmon after adjusting for density dependence. However, the bootstrap confidence intervals for the model parameters were wide and included maximum recruitment levels beyond the range of the observed data (Table 8,

Table 9), demonstrating an overall lack of fit. The small sample size may have contributed to the lack of model fit, and likewise may lower the ability to detect a relationship between pHOS and smolts per redd using the available data.


Figure 31. Smolts per redd versus proportion of hatchery origin spawners ( pHOS ) for White River Spring Chinook Salmon, 2005-2017, with and without brood year 2014, with fitted linear regression line, Pearson correlation coefficient (r), and P-value from one-sided t-test of negative slope. $\mathrm{R}^{2}=0.0351$ with BY 2014 and $\mathrm{R}^{2}=0.0916$ without BY 2014.


Figure 32. Redd count versus proportion of hatchery origin spawners (pHOS) for White River Spring Chinook Salmon, 2005-2017, including 2014 brood year, with fitted linear regression line, Pearson correlation coefficient (r), and P-value from two-sided t -test of slope.


Figure 33. Smolts per redd (log scale) versus redd count for White River Spring Chinook Salmon, 20052017, with and without the 2014 brood year, with fitted linear regression line, Pearson correlation coefficient (r), and P-value from one-sided t-test of negative slope. Negative slope indicates post-spawning density dependence.

Table 8. Fitted stock-recruitment models for smolts per redd for White River Spring Chinook Salmon, 2005-2017, including brood year 2014, including effect $\left(\beta_{2}\right)$ of proportion of hatchery origin spawners (pHOS). Confidence intervals were estimated using bootstrap with 3,000 bootstrap samples. Model equations are defined in Equations (5) and (7). $\Delta \mathrm{AICc}$ represents change in AICc compared to model with minimum AICc.

| Model | AICc | $\Delta$ AICc | Parameter | Parameter Estimates | $95 \% \mathrm{CI}$ |
| :--- | :--- | :--- | :---: | :---: | :---: |
| Beverton-Holt | 38.71 | 0.00 | $\alpha$ | 9,478 | $4,270-173,640$ |
|  |  |  | $\beta$ | 48.1 | $12.9-1,315$ |
|  |  |  | $\beta_{2}$ | -0.0887 | $-1.6125-1.6392$ |
|  |  |  |  |  |  |
| Ricker | 38.82 | 0.10 | $\ln (\alpha)$ | 5.0979 | $3.5461-6.3336$ |
|  |  |  | $\beta$ | 0.0096 | $-0.0080-0.0228$ |
|  |  |  | $\beta_{2}$ | -0.0655 | $-2.2529-1.5695$ |

Table 9. Fitted stock-recruitment models for smolts per redd for White River Spring Chinook Salmon, 2005-2017, omitting brood year 2014, including effect $\left(\beta_{2}\right)$ of proportion of hatchery origin spawners (pHOS). Confidence intervals were estimated using bootstrap with 3,000 bootstrap samples. Model equations are defined in Equations (5) and (7). $\Delta$ AICc represents change in AICc compared to model with minimum AICc.

| Model | AICc | $\Delta$ AICc | Parameter | Parameter Estimates | $95 \% \mathrm{CI}$ |
| :--- | :---: | :--- | :---: | :---: | :---: |
| Beverton-Holt | 24.87 | 0.00 | $\alpha$ | 9,680 | $4,671-39,360$ |
|  |  |  | $\beta$ | 24.9 | $3.0-175.3$ |
|  |  |  | $\beta_{2}$ | 0.3364 | $-0.6880-1.3979$ |
|  |  |  |  |  |  |
| Ricker | 25.26 | 0.38 | $\ln (\alpha)$ | 5.5953 | $4.7824-6.5861$ |
|  |  |  | $\beta$ | 0.0135 | $0.0030-0.0237$ |
|  |  |  | $\beta_{2}$ | 0.3455 | $-0.8663-1.8240$ |

## Twisp River Spring Chinook Salmon

Adult spawner, smolt recruitment, pHOS , and redd count data were available for the 2003-2017 brood years. Adult stock abundance (spawners) ranged from 43 in 2003 to 341 in 2004 and averaged 165, and smolt recruitment ranged from 900 in 2003 to 16,415 in 2006 and averaged 5,295 (Figure 34, Figure 35). Redd counts averaged 79.3 (range $=18$ to 145), and pHOS averaged 0.52 (range $=0.28$ to 0.76 ).


Figure 34. Twisp River Spring Chinook Salmon boxplots for Stock (spawner abundance), Smolts, Redd counts, and proportion of hatchery origin spawners (pHOS), 2003-2017. Box indicates interquartile range and thick horizontal bar is median. Whiskers indicate quartile $\pm 1.5 \times$ interquartile range.


Figure 35. Twisp River Spring Chinook Salmon annual Smolt counts, Smolt/Spawner (log scale), Smolt/Redd (log scale), and proportion of hatchery origin spawners (pHOS), 2002-2017. Line indicates the linear trend. Shaded area is the estimated $95 \%$ confidence interval of the line.

## Hypothesis H02.2.1.1

A weak negative linear relationship was observed between Twisp River Spring Chinook Salmon spawners and smolt recruitment, providing evidence of density dependence in mortality between the spawning and smolt life stages ( $\mathrm{P}=0.0463$; Figure 36). Thus, the Smooth Hockey Stick, Beverton-Holt, and Ricker stockrecruitment models were fit to the spawner and smolt data.

There was little difference in the model fits among the three stock-recruitment models, with the only perceptible differences in model predictions observed for spawner abundance greater than approximately 240 (Figure 37). The Ricker model had the lowest AICc value but the Beverton-Holt and Smooth Hockey Stick models both had $\Delta \mathrm{AICc} \leq 0.62$ (Table 10). However, all three models produced wide bootstrap confidence intervals for the model parameters. The maximum recruitment estimates in particular had confidence intervals that extended far beyond the maximum observed smolt recruitment $(16,415)$. High correlation was observed between the model parameter estimates, increasing uncertainty in the model predictions (Table 10, Figure D3). Additionally, analysis of residuals from each of the models demonstrated a negative autocorrelation at a time lag of 3 years (Figure D2, Figure D5, Figure D7), further suggesting that these models do not adequately account for the population dynamics and that the model parameters may be biased.

Correlation analysis found no association between the residuals from these stock-recruitment models and pHOS, with correlation coefficients ranging from 0.1612 to 0.1921 ( $\mathrm{P} \geq 0.4927$; Figure 38 - Figure 40 ).

Additionally, likelihood ratio tests of the effect of pHOS on the Beverton-Holt and Ricker models were non-significant ( $\mathrm{P} \geq 0.4470$ ). Thus, there was no evidence to suggest that higher proportions of hatchery spawners were associated with lower juvenile productivity for spring Chinook Salmon in the Twisp River. However, there were only 15 years of data available, and the modeling assumptions were not well supported. Thus, it remains possible that there is a relationship between pHOS and juvenile productivity that are not observable using the available data and analyses.


Figure 36. Smolts per spawner (log scale) versus spawner abundance for Twisp River Spring Chinook Salmon, 2003-2017, with fitted linear regression line, Pearson correlation coefficient (r), and P-value from one-sided t-test of negative slope.

Twisp Spring Chinook Salmon


Figure 37. Stock-recruitment models fit to smolt and spawner data for Twisp River Spring Chinook Salmon, 2003-2017.

Table 10. Fitted stock-recruitment models for smolts and stock (spawner) data from Twisp River Spring Chinook Salmon, 2003-2017. Confidence intervals were estimated using bootstrap with at least 2,000 bootstrap samples. Correlation $=$ Pearson correlation coefficient between parameter estimates from bootstrap samples. Model equations are defined in Equations (1), (2), and (3). $\Delta \mathrm{AICc}$ represents change in AICc compared to model with minimum AICc.

| Model | AICc | $\Delta$ AICc | Parameter $^{\mathrm{b}}$ | Parameter <br> Estimates | $95 \%$ CI | Correlation |
| :--- | :---: | :--- | :---: | :---: | :---: | :---: |
| Smooth Hockey 30.39 0.31 $\alpha$ 51.2 $30.7-146.7$ <br> Stick   $R_{\infty}$ 7,104 $4,344-36,313$ | $\alpha, R_{\infty}:-0.20^{\mathrm{a}}$ |  |  |  |  |  |
|  |  |  |  |  |  |  |
| Beverton-Holt | 30.69 | 0.62 | $\alpha$ | 11,295 | $6,317-101,201$ | $\alpha, \beta: 0.99$ |
|  |  |  | $\beta$ | 216 | $75-3,103$ |  |
| Ricker | 30.07 | 0.00 | $\alpha$ | 51.2 | $29.7-93.6$ | $\alpha, \beta: 0.88$ |
|  |  |  | $\beta$ | 0.0032 | $<0.0001-0.0064$ |  |
|  |  |  | $K$ | 5,926 | $4,267-\infty$ |  |

$\mathrm{a}=$ Correlation coefficient between $\alpha$ and $R_{\infty}$ is uninformative because parameter estimates are not linearly related; see Figure D3. The correlation coefficient between $1 / \alpha$ and $R_{\infty}=0.4951$.
$\mathrm{b}=$ Estimated maximum recruitment parameters are: $R_{\infty}$ for the Smooth Hockey Stick model, $\alpha$ for the Beverton-
Holt model, and $K=(\alpha / \beta) e^{-1}$ for the Ricker model.


Figure 38. Residuals from Smooth Hockey Stick Model versus proportion of hatchery origin spawners ( pHOS ) with fitted linear regression line, Pearson correlation coefficient (r), and P-value from two-sided ttest of slope of linear regression line using smolt and spawner data from Twisp River Spring Chinook Salmon, 2003-2017. $\mathrm{R}^{2}=0.0324$.


Figure 39. Residuals from the Beverton-Holt Model versus proportion of hatchery origin spawners ( pHOS ) with fitted linear regression line, Pearson correlation coefficient (r), and P-value from two-sided t-test of slope of linear regression line using smolt and spawner data from Twisp River Spring Chinook Salmon, 2003-2017. $\mathrm{R}^{2}=0.0369$.


Figure 40. Residuals from the Ricker Model versus proportion of hatchery origin spawners ( pHOS ) with fitted linear regression line, Pearson correlation coefficient (r), and P-value from two-sided t-test of slope of linear regression line using smolt and spawner data from Twisp River Spring Chinook Salmon, 20032017. $\mathrm{R}^{2}=0.0260$.

## Hypothesis H02.2.1.2

There was no evidence of a negative association between smolts per redd and the proportion of hatchery spawners (pHOS) for Twisp River Spring Chinook Salmon (Pearson correlation coefficient $\mathrm{r}=0.2925$, $\mathrm{P}=0.8549$ ) (Figure 41). Additionally, the linear association between redd count and pHOS was nonsignificant ( $\mathrm{P}=0.8313$; Figure 42). However, comparison of smolts per redd with redd count demonstrated
evidence of post-spawning density dependence ( $\mathrm{P}=0.0142$; Figure 43 ). Thus, the relationship between smolts per redd and pHOS was examined using the Ricker and Beverton-Holt stock-recruitment models in order to account for density dependence effects, using redds in place of spawner abundance. The model fits and assessment of residuals were very similar (Table 11, Figure D9 - Figure D13). The fitted BevertonHolt model for smolts per redd was (Table 11):

$$
\ln (R / S)=\ln (8,020)+0.7782 P_{\text {HOS }}-\ln (103+S)
$$

The $95 \%$ bootstrap confidence interval for the effect of pHOS (i.e., $\beta_{2}$ in Equation (5)) was (-1.9437, 0.4321 ). The fitted Ricker model for smolts per redd was:

$$
\ln (R / S)=4.3238+0.7621 P_{\text {HOS }}-0.0063 S
$$

The $95 \%$ bootstrap confidence interval for the effect of pHOS (i.e., $\beta_{2}$ in Equation (7)) was (-2.0433, $0.1133)$.

Because the confidence intervals for the effect of pHOS included both positive and negative values regardless of the model, we concluded that there was no evidence of a negative association between pHOS and smolts per redd for Twisp River Spring Chinook Salmon after adjusting for density dependence. However, the bootstrap confidence intervals for the model parameters were wide and included maximum recruitment levels beyond the range of the observed data (Table 11), demonstrating an overall lack of fit. There was moderate evidence of autocorrelation in the residuals (Figure D11, Figure D13), suggesting a failure to account for some feature of the population dynamics. The small sample size may have contributed to the lack of model fit, and likewise may lower the ability to detect a relationship between pHOS and smolts per redd using the available data.


Figure 41. Smolts per redd (log scale) versus proportion of hatchery origin spawners (pHOS) for Twisp River Spring Chinook Salmon, 2003-2017, with fitted linear regression line, Pearson correlation coefficient (r), and P-value from one-sided t-test of negative slope. $\mathrm{R}^{2}=0.0855$.


Figure 42. Redd count versus proportion of hatchery origin spawners (pHOS) for Twisp River Spring Chinook Salmon, 2003-2017, with fitted linear regression line, Pearson correlation coefficient (r), and Pvalue from two-sided $t$-test of slope.


Figure 43. Smolts per redd (log scale) versus redd count for Twisp River Spring Chinook Salmon, 20032017, with fitted linear regression line, Pearson correlation coefficient (r), and P-value from one-sided ttest of negative slope. Negative slope indicates post-spawning density dependence.

Table 11. Fitted stock-recruitment models for smolts per redd for Twisp River Spring Chinook Salmon, 2003-2017, including effect ( $\beta_{2}$ ) of proportion of hatchery origin spawners (pHOS). Confidence intervals were estimated using bootstrap with 3,000 bootstrap samples. Model equations are defined in Equations (5) and (7). $\triangle \mathrm{AICc}$ represents change in AICc compared to model with minimum AICc.

| Model | AICc | $\Delta$ AICc | Parameter | Parameter Estimates | $95 \% \mathrm{CI}$ |
| :--- | :---: | :--- | :---: | :---: | :---: |
| Beverton-Holt | 28.58 | 0.78 | $\alpha$ | 8,020 | $3,952-39,818$ |
|  |  |  | $\beta$ | 103 | $40-724$ |
|  |  |  | $\beta_{2}$ | -0.7782 | $-1.9437-0.4321$ |
|  |  |  |  |  |  |
| Ricker | 27.79 | 0.00 | $\ln (\alpha)$ | 4.3238 | $3.6525-5.0395$ |
|  |  |  | $\beta$ | 0.0063 | $0.0011-0.0108$ |
|  |  |  | $\beta_{2}$ | -0.7621 | $-2.0433-0.1133$ |

## Methow River Spring Chinook Salmon

Adult spawner, smolt recruitment, pHOS , and redd count data from spring Chinook Salmon were available from the Methow River for the 2002-2017 brood years. Adult stock abundance (spawners) ranged from 417 in 2017 to 2,692 in 2011 and averaged 1,417, and smolt recruitment ranged from 5,163 in 2007 to 51,325 in 2010 and averaged 23,624 (Figure 44, Figure 45). Redd counts averaged 705 (range $=210$ to 1,366 ), and pHOS averaged 0.77 (range $=0.57$ to 0.97 ).


Figure 44. Methow River Spring Chinook Salmon boxplots for Stock (spawner abundance), Smolts, Redd counts, and proportion of hatchery origin spawners (pHOS), 2002-2017. Box indicates interquartile range and thick horizontal bar is median. Whiskers indicate quartile $\pm 1.5 \times$ interquartile range.


Figure 45. Methow River Spring Chinook Salmon annual Smolt counts, Smolt/Spawner (log scale), Smolt/Redd (log scale), and proportion of hatchery origin spawners (pHOS), 2002-2017. Line indicates the linear trend. Shaded area is the estimated $95 \%$ confidence interval of the line.

## Hypothesis H02.2.1.1

A weak negative linear relationship was observed between Methow River Spring Chinook Salmon spawners and smolt recruitment but it was non-significant, providing inconclusive evidence of density dependence in mortality between the spawning and smolt life stages ( $\mathrm{P}=0.2101$; Figure 46). Nevertheless, the Smooth Hockey Stick, Beverton-Holt, and Ricker stock-recruitment models were fit to the spawner and smolt data.

There was little difference in the model fits among the three stock-recruitment models, with no perceptible difference in model predictions for any range of observed spawner abundance (Figure 47). The Ricker model had the lowest AICc value but $\Delta \mathrm{AICc} \leq 0.04$ for all models (Table 12). Despite the agreement in model predictions among the three models, all models showed considerable uncertainty in the estimated parameters, with particularly wide confidence intervals. The maximum recruitment estimates in particular had confidence intervals that extended far beyond the maximum observed smolt recruitment $(51,325)$. High correlation was observed between the model parameter estimates, increasing uncertainty in the model predictions (Table 12, Figure E3). However, there was no severe violation of the assumption of lognormal errors and no evidence of lack of stationarity or autocorrelation. (Figures E1, E2, Figure E4 - Figure E7).

Correlation analysis found no association between the residuals from these stock-recruitment models and pHOS, with correlation coefficients ranging from 0.0697 to 0.0704 ( $\mathrm{P} \geq 0.7955$; Figure 48 - Figure 50). Additionally, likelihood ratio tests of the effect of pHOS on the Beverton-Holt and Ricker models were
non-significant ( $\mathrm{P} \geq 0.7635$ ). Thus, there was no evidence to suggest that higher proportions of hatchery spawners were associated with lower juvenile productivity for spring Chinook Salmon in the Methow River. However, there were only 16 years of data available, and the modeling assumptions were not well supported. Thus, it remains possible that there is a relationship between pHOS and juvenile productivity that are not observable using the available data and analyses.


Figure 46. Smolts per spawner (log scale) versus spawner abundance for Methow River Spring Chinook Salmon, 2002-2017, with fitted linear regression line, Pearson correlation coefficient (r), and P-value from one-sided t -test of negative slope.


Figure 47. Stock-recruitment models fit to smolt and spawner data for Methow River Spring Chinook Salmon, 2002-2017.

Table 12. Fitted stock-recruitment models for smolts and stock (spawner) data from Methow River Spring Chinook Salmon, 2002-2017. Confidence intervals were estimated using bootstrap with at least 2,000 bootstrap samples. Correlation $=$ Pearson correlation coefficient between parameter estimates from bootstrap samples. Model equations are defined in Equations (1), (2), and (3). $\Delta \mathrm{AICc}$ represents change in AICc compared to model with minimum AICc.

| Model | AICc | $\Delta$ AICc | Parameter $^{\mathrm{b}}$ | Parameter <br> Estimates | $95 \% \mathrm{CI}$ | Correlation |
| :--- | :---: | :---: | :---: | :---: | :---: | :---: |
| Smooth Hockey | 26.64 | 0.02 | $\alpha$ | 19.7 | $14.7-37.7$ | $\alpha, R_{\infty}:-0.30^{\mathrm{a}}$ |
| Stick |  |  | $R_{\infty}$ | 65,051 | $23,577-549,603$ |  |
|  |  |  |  |  | 115,394 | $32,755-1,382,406$ |
| Beverton-Holt | 26.66 | 0.04 | $\alpha$ | $\alpha, \beta: 0.99$ |  |  |
|  |  |  | $\beta$ | 5,806 | $785-82,314$ |  |
| Ricker | 26.62 | 0.00 | $\alpha$ | 19.6 | $11.6-32.7$ | $\alpha, \beta: 0.88$ |
|  |  |  | $\beta$ | 0.0001 | $<0.0001-0.0005$ |  |
|  |  |  | $K$ | 50,572 | $22,637-\infty$ |  |

$\overline{\mathrm{a}}=$ Correlation coefficient between $\alpha$ and $R_{\infty}$ is uninformative because parameter estimates are not linearly related; see Figure D3. The correlation coefficient between $1 / \alpha$ and $R_{\infty}=0.7281$.
$\mathrm{b}=$ Estimated maximum recruitment parameters are: $R_{\infty}$ for the Smooth Hockey Stick model, $\alpha$ for the BevertonHolt model, and $K=(\alpha / \beta) e^{-1}$ for the Ricker model.


Figure 48. Residuals from Smooth Hockey Stick Model versus proportion of hatchery origin spawners ( pHOS ) with fitted linear regression line, Pearson correlation coefficient ( r ), and P -value from two-sided t test of slope of linear regression line using smolt and spawner data from Methow River Spring Chinook Salmon, 2002-2017. $\mathrm{R}^{2}=0.0050$.


Figure 49. Residuals from the Beverton-Holt Model versus proportion of hatchery origin spawners ( pHOS ) with fitted linear regression line, Pearson correlation coefficient (r), and P-value from two-sided t-test of slope of linear regression line using smolt and spawner data from Methow River Spring Chinook Salmon, 2003-2017. $\mathrm{R}^{2}=0.0049$.


Figure 50. Residuals from the Ricker Model versus proportion of hatchery origin spawners ( pHOS ) with fitted linear regression line, Pearson correlation coefficient (r), and P-value from two-sided t-test of slope of linear regression line using smolt and spawner data from Methow River Spring Chinook Salmon, 20022017. $\mathrm{R}^{2}=0.0049$.

## Hypothesis H02.2.1.2

There was no evidence of a negative association between smolts per redd and the proportion of hatchery spawners (pHOS) for Methow River Spring Chinook Salmon (Pearson correlation coefficient $\mathrm{r}=-0.2409$, $\mathrm{P}=0.1844$ ) (Figure 51). Although the linear association between redd count and pHOS was significant at the $10 \%$ level ( $\mathrm{P}=0.0919$; Figure 52), comparison of smolts per redd with redd count demonstrated no
evidence of post-spawning density dependence ( $\mathrm{P}=0.3761$, Figure 53). Nevertheless, the relationship between smolts per redd and pHOS was examined using stock-recruitment models in order to account for density dependence effects, using redds in place of spawner abundance. The Beverton-Holt model could not be fit to the data; examination of the likelihood indicated a flat likelihood surface and high correlation between the $\alpha$ and $\beta$ model parameter estimates which could not be overcome when pHOS was included in the model. The Ricker model was able to be fit and the residuals agreed moderately well with model assumptions (Figures E9, Figure E10); however, the estimate for the $\beta$ parameter was essentially 0 , which is not consistent with interpretation of model parameters (Table 13). The fitted Ricker model for smolts per redd was:

$$
\ln (R / S)=4.0380-0.7383 * P_{\text {HoS }}+(<0.0001) * S .
$$

The $95 \%$ bootstrap confidence interval for the effect of pHOS (i.e., $\beta_{2}$ in Equation (7)) was $-1.2657-$ 2.3241). Because the confidence interval for the effect of pHOS included both positive and negative values, we concluded that there was no evidence of a negative association between pHOS and smolts per redd for Methow River Spring Chinook Salmon whether or not an adjustment was made for density dependence. However, the model parameters were not consistent with their usual interpretation, demonstrating an overall lack of fit. The small sample size, low degree of variability in the observed pHOS values, and potentially high level of measurement error in the observed data may have contributed to the lack of model fit, and likewise may lower the ability to detect a relationship between pHOS and smolts per redd using the available data.


Figure 51. Smolts per redd versus proportion of hatchery origin spawners (pHOS) for Methow River Spring Chinook Salmon, 2002-2017, with fitted linear regression line, Pearson correlation coefficient (r), and Pvalue from one-sided $t$-test of negative slope. $\mathrm{R}^{2}=0.0580$.


Figure 52. Redd count versus proportion of hatchery origin spawners (pHOS) for Methow River Spring Chinook Salmon, 2002-2017, with fitted linear regression line, Pearson correlation coefficient (r), and Pvalue from two-sided $t$-test of slope.


Figure 53. Smolts per redd (log scale) versus redd count for Methow River Spring Chinook Salmon, 20022017, with fitted linear regression line, Pearson correlation coefficient (r), and P-value from one-sided ttest of negative slope. Negative slope indicates post-spawning density dependence.

Table 13. Fitted stock-recruitment models for smolts per redd for Methow River Spring Chinook Salmon, 2002-2017, including effect ( $\beta_{2}$ ) of proportion of hatchery origin spawners (pHOS). Confidence intervals were estimated using bootstrap with at least 2,000 bootstrap samples. Model equations are defined in Equations (5) and (7). $\triangle$ AICc represents change in AICc compared to model with minimum AICc. *Only the Ricker model could be fit for Methow River Spring Chinook Salmon data.

| Model $^{*}$ | AICc | $\Delta$ AICc | Parameter | Parameter Estimates | $95 \% \mathrm{CI}$ |
| :--- | :--- | :--- | :---: | :---: | :---: |
| Beverton-Holt | NA | NA | $\alpha$ | NA | NA |
|  |  |  | $\beta$ | NA | NA |
|  |  |  | $\beta_{2}$ | NA | NA |
| Ricker | 23.56 | NA | $\ln (\alpha)$ | 4.0380 | $2.6075-5.3023$ |
|  |  |  | $\beta$ | $<0.0001$ | $<0.0001-0.0006$ |
|  |  |  | $\beta_{2}$ | 0.7383 | $-1.2657-2.3241$ |

## Wenatchee River Summer Chinook Salmon

Adult spawner, emigrant, pHOS , and redd count data were available for the 1999-2018 brood years (no emigrant count for 2010 and 2011). Adult stock abundance (spawners) ranged from 3,473 in 2018 to 17,792 in 2006 and averaged 8,695. Emigrant counts ranged from 1,322,383 in 2000 to 20,426,149 in 2003 and averaged 9,118,268 (Figure 54, Figure 55). Redd counts averaged 3,600.5 (range $=1,510$ to 8,896 ), and pHOS averaged 0.176 (range $=0.06$ to 0.31 ). The 2000 and 2003 brood years were identified as possible outliers in emigrant counts (Figure 55), and analysis results were investigated for their sensitivity to inclusion of these brood years.


Figure 54. Wenatchee River Summer Chinook Salmon boxplots for Stock (spawner abundance), Emigrant counts, Redd counts, and proportion of hatchery origin spawners (pHOS), 1999-2018, without 2010 and 2011. Box indicates interquartile range and thick horizontal bar is median. Whiskers indicate quartile $\pm 1.5$ $\times$ interquartile range.


Figure 55. Wenatchee River Summer Chinook Salmon annual Emigrant counts, Emigrant/Spawner (log scale), Emigrant/Redd (log scale), and proportion of hatchery origin spawners (pHOS), 1999-2018 (without 2010, 2011). Line indicates the linear trend. Shaded area is the estimated $95 \%$ confidence interval of the line.

## Hypothesis H02.2.1.1

A positive but non-significant linear relationship was observed between spawners and emigrant count, indicating no evidence of density dependence in mortality between spawner data collection and emigrant data collection (Figure 56). The Smooth Hockey Stick and Beverton-Holt stock-recruitment models could not be fit to the spawner and emigrant data, so only the Ricker model is presented (Figure 57 with parameter estimates (Table 14). The residuals from the Ricker model was compared to pHOS using correlation analysis. The correlation coefficient between residuals and pHOS was -0.0985 and was not significantly different from 0 ( $\mathrm{P}=0.6973$; Figure 58). Additionally, the likelihood ratio test of the effect of pHOS on the Ricker model was non-significant ( $\mathrm{P}=0.6787$ ). There was no evidence to suggest that higher proportions of hatchery spawners were associated with lower juvenile productivity for Summer Chinook Salmon in the Wenatchee River.

When the 2003 brood year was omitted as a possible outlier, the Smooth Hockey Stick and Beverton-Holt models could be fit to the spawner and emigrant data but had equivalent weight with the Ricker model ( $\Delta \mathrm{AICc} \leq 0.0072$ ); pHOS was not associated with the residuals from any of the models ( $\mathrm{P} \geq 0.7155$ ). When the 2000 brood year was omitted as a possible outlier, all three models could be fit to the data but the Ricker model was again selected ( $\triangle \mathrm{AICc} \leq 0.1126$ ). Model residuals were again not significantly associated with
pHOS levels without the 2000 brood year ( $\mathrm{P} \geq 0.6430$; Figure F3 - Figure F6), and the likelihood ratio test of the pHOS was also not significant ( $\mathrm{P} \geq 0.6113$ ).


Figure 56. Emigrants per spawner ( $\log$ scale) versus spawner abundance for Wenatchee River Summer Chinook Salmon, 1999-2018 (without 2010, 2011), with fitted linear regression line, Pearson correlation coefficient (r), and P-value from one-sided $t$-test of negative slope.

Wenatchee Summer Chinook Salmon


Figure 57. The Ricker stock-recruitment model fit to emigrants and spawner data for Wenatchee River Summer Chinook Salmon, 1999-2018 (without 2010, 2011).

Table 14. Fitted stock-recruitment models for emigrants and stock (spawner) data from Wenatchee River Summer Chinook Salmon, 1999-2018 (without 2010, 2011). Confidence intervals were estimated using bootstrap with at least 2,900 bootstrap samples. Correlation $=$ Pearson correlation coefficient between parameter estimates from bootstrap samples. Model equations are defined in Equations (1), (2), and (3). $\Delta$ AICc represents change in AICc compared to model with minimum AICc. *Only the Ricker model could be fit to the Wenatchee River data.

| Model* | AICc | $\triangle \mathrm{AICc}$ | Parameter ${ }^{\text {a }}$ | Parameter <br> Estimates | 95\% CI | Correlation |
| :---: | :---: | :---: | :---: | :---: | :---: | :---: |
| Smooth Hockey Stick |  |  | $\alpha$ | NA | NA | $\alpha, R_{\infty}:$ NA |
|  |  |  | $R_{\infty}$ | NA | NA |  |
| Beverton-Holt |  |  | $\alpha$ | NA | NA | $\alpha, \beta$ NA |
|  |  |  | $\beta$ | NA | NA |  |
| Ricker | 31.72 |  | $\alpha$ | 957.3 | 773.4 - 1609.5 | $\alpha, \beta: 0.83$ |
|  |  |  | $\beta$ | <0.0001 | $<0.0001-0.0001$ |  |
|  |  |  | $K$ | 30,446,537 | 919,213-m |  |

$\mathrm{a}=$ Estimated maximum recruitment parameters are: $R_{\infty}$ for the Smooth Hockey Stick model, $a$ for the BevertonHolt model, and $K=(\alpha / \beta) e^{-1}$ for the Ricker model.


Figure 58. Residuals from Ricker Model versus proportion of hatchery origin spawners ( pHOS ) with fitted linear regression line, Pearson correlation coefficient ( $r$ ), and P-value from two-sided $t$-test of slope of linear regression line using emigrant and spawner data from Wenatchee River Summer Chinook Salmon, 19992018 (without 2010, 2011). $\mathrm{R}^{2}=0.0097$.

## Hypothesis H02.2.1.2

There was a weak and non-significant negative association between emigrants per redd (log scale) and the proportion of hatchery spawners ( pHOS ) for Wenatchee River Summer Chinook Salmon (Pearson correlation coefficient $\mathrm{r}=-0.0750, \mathrm{P}=0.3836$ ) (Figure 59). The redd count was not associated with pHOS ( $\mathrm{P}=0.4460$, Figure 60 ) and there was no indication of post-spawning density dependence apparent from examination of the emigrants per redd plotted against the redd counts (Figure 61). Nevertheless, the relationship between emigrants per redd and pHOS was examined using the Ricker and Beverton-Holt stock-recruitment models in order to account for density dependence effects, with redds used in place of spawner counts. Both models were equivalent on the basis of AICc rank ( $\triangle \mathrm{AICc}=0.01$ compared to Ricker model) and examination of model residuals (Table 15; Figure F8 - Figure F12). However, the BevertonHolt model could not be fit for the bootstrap samples and the $95 \%$ asymptotic confidence intervals for the model parameters $\alpha$ and $\beta$ included both negative values and values far beyond the range of the observed data; both these results indicate a lack of fit for the Beverton-Holt model. Thus, the Ricker model was preferred. The fitted Ricker model for emigrants per redd was (Table 15):

$$
\ln (R / S)=8.0322-0.7166 P_{\text {HOS }}-0.00005 * S
$$

The $95 \%$ bootstrap confidence interval for the effect of pHOS (i.e., $\beta_{2}$ in Equation (7)) was ( -2.0006 3.3638). Because this confidence interval included both positive and negative values, we concluded that there is no evidence of a negative association between pHOS and emigrants per redd for Wenatchee Summer Chinook Salmon after adjusting for density dependence. Removing 2000 or 2003 as possible outlier brood years did not significantly change the results (e.g., Figure F13).


Figure 59. Emigrants per redd ( $\log$ scale) versus proportion of hatchery origin spawners ( pHOS ) for Wenatchee River Summer Chinook Salmon, 1999-2018 (without 2010, 2011), with fitted linear regression line, Pearson correlation coefficient (r), and P-value from one-sided t-test of negative slope. $\mathrm{R}^{2}=0.0056$.


Figure 60. Redd count versus proportion of hatchery origin spawners (pHOS) for Wenatchee River Summer Chinook Salmon, 1999-2018 (without 2010, 2011), with fitted linear regression line, Pearson correlation coefficient (r), and P-value from two-sided t -test of slope.


Figure 61. Emigrants per redd versus redd count for Wenatchee River Summer Chinook Salmon, 19992018 (without 2010, 2011), with fitted linear regression line, Pearson correlation coefficient (r), and P-value from one-sided $t$-test of negative slope. Negative slope indicates post-spawning density dependence.

Table 15. Fitted stock-recruitment models for emigrants per redd for Wenatchee River Summer Chinook Salmon, 1999-2018 (without 2010, 2011), including effect $\left(\beta_{2}\right)$ of proportion of hatchery origin spawners (pHOS). Bootstrap confidence intervals based on 3,000 bootstrap samples were provide for the Ricker model, and asymptotic confidence intervals were provided for the Beverton-Holt model. Model equations are defined in Equations (5) and (7). $\Delta \mathrm{AICc}$ represents change in AICc compared to model with minimum AICc.

| Model | AICc | $\Delta$ AICc | Parameter | Parameter Estimates | $95 \% \mathrm{CI}$ |
| :--- | :---: | :---: | :---: | :---: | :---: |
| Beverton-Holt | 38.41 | 0.10 | $\alpha$ | $69,904,888$ | $-259,329,930-399,139,700$ |
|  |  |  | $\beta$ | 23,464 | $-105,680-9,152,607$ |
|  |  |  | $\beta_{2}$ | 0.6695 | $-3.1075-4.4465$ |
| Ricker | 38.31 | 0.00 | $\ln (\alpha)$ | 8.0322 |  |
|  |  |  | $\beta$ | 0.00005 | $-0.8834-8.6882$ |
|  |  |  | $\beta_{2}$ | 0.7166 | $-2.0002-0.0001$ |
|  |  |  |  |  |  |

## Methow River Summer Chinook Salmon

Adult spawner, emigrant, pHOS , and redd count data were available for the 2006-2018 brood years (no emigrant count for 2012). Adult stock abundance (spawners) ranged from 1,364 in 2007 to 3,952 in 2015 and averaged 2,333, and emigrant count ranged from 427,193 in 2017 to 3,465,247 in 2006 and averaged 1,099,370 (Figure 62, Figure 63). Redd counts averaged 909.5 (range $=591$ to 1,551 ), and pHOS averaged 0.391 (range $=0.11$ to 0.53 ).


Figure 62. Methow River Summer Chinook Salmon boxplots for Stock (spawner abundance), Emigrant counts, Redd counts, and proportion of hatchery origin spawners ( pHOS ), 2006-2018. Box indicates interquartile range and thick horizontal bar is median. Whiskers indicate quartile $\pm 1.5 \times$ interquartile range.


Figure 63. Methow River Summer Chinook Salmon annual Emigrant counts, Emigrant/Spawner (log scale), Emigrant/Redd (log scale), and proportion of hatchery origin spawners (pHOS), 2006-2018 (without 2012). Line indicates the linear trend. Shaded area is the estimated $95 \%$ confidence interval of the line.

## Hypothesis H02.2.1.1

A slightly negative linear relationship between spawners and emigrant count indicated the presence of density dependence in mortality in one or more life stages between spawner data collection and emigrant data collection (Figure 64). Thus, the Smooth Hockey Stick, Beverton-Holt, and Ricker stock-recruitment models were each fit to the spawner and emigrant data (Figure 65). There was little difference in the model fits among the three models ( $\triangle \mathrm{AICc} \leq 0.02$ ), but the Ricker model had the lowest AICc value and the intermediate correlation between parameter estimates (Table 16). The residuals from all three models were compared to pHOS using correlation analysis. Correlation coefficients between residuals and pHOS were all non-significant ( $\mathrm{r}=-0.0059, \mathrm{P}=0.9856$ for each model; Figure 66 - Figure 68). Additionally, likelihood ratio tests of the effect of pHOS on the Beverton-Holt and Ricker models were non-significant ( $\mathrm{P} \geq 0.9838$ ). There was no evidence to suggest that higher proportions of hatchery spawners were associated with lower juvenile productivity for Summer Chinook Salmon in the Methow River.


Figure 64. Emigrants per spawner (log scale) versus spawner abundance for Methow River Summer Chinook Salmon, 2006-2018 (without 2012), with fitted linear regression line, Pearson correlation coefficient (r), and P-value from one-sided t-test of negative slope.


Figure 65. Stock-recruitment models fit to emigrants and spawner data for Methow River Summer Chinook Salmon, 2006-2018 (without 2012).

Table 16. Fitted stock-recruitment models for emigrants and stock (spawner) data from Methow River Summer Chinook Salmon, 2006-2018 (without 2012). Confidence intervals were estimated using bootstrap with at least 1700 bootstrap samples. Correlation $=$ Pearson correlation coefficient between parameter estimates from bootstrap samples. Model equations are defined in Equations (1), (2), and (3). $\Delta \mathrm{AICc}$ represents change in AICc compared to model with minimum AICc.

|  |  |  | Parameter |  |  |  |
| :--- | :--- | :--- | :---: | :---: | :---: | :---: |
| Model | AICc | $\Delta$ AICc | Parameter $^{\mathrm{b}}$ | Estimates | $95 \% \mathrm{CI}$ | Correlation |
| Smooth | 26.85 | 0.01 | $\alpha$ | 459 | $352.6-1344.6$ | $\alpha, R_{\infty}:-0.23^{\mathrm{a}}$ |
| Hockey Stick |  |  | $R_{\infty}$ | $4,796,000$ | $889,141-19,840,304$ |  |
|  |  |  |  |  |  |  |
|  |  | $\alpha$ | $9,817,511$ | $874,870-66,701,078$ | $\alpha, \beta: 0.99$ |  |
| Beverton-Holt | 26.86 | 0.02 |  | 21,552 | $-26.7-16,434.1$ |  |
|  |  |  |  |  |  |  |
| Ricker | 26.84 | 0.00 | $\alpha$ | 461 | $331.4-1032.5$ | $\alpha, \beta: 0.88$ |
|  |  |  | $\beta$ | 0.00005 | $<0.0001-0.0004$ |  |
|  |  |  | $K$ | $3,457,540$ | $919,213-\infty$ |  |

$\mathrm{a}=$ Correlation coefficient between $a$ and $R_{\infty}$ is uninformative because parameter estimates are not linearly related; see Figure D3. The correlation coefficient between $1 / \alpha$ and $R_{\infty}=0.2984$.
$\mathrm{b}=$ Estimated maximum recruitment parameters are: $R_{\infty}$ for the Smooth Hockey Stick model, $a$ for the Beverton-Holt model, and $K=(\alpha / \beta) e^{-1}$ for the Ricker model.


Figure 66. Residuals from Smooth Hockey Stick Model versus proportion of hatchery origin spawners ( pHOS ) with fitted linear regression line, Pearson correlation coefficient (r), and P-value from two-sided ttest of slope of linear regression line using emigrant and spawner data from Methow River Summer Chinook Salmon, 2006-2018 (without 2012). $\mathrm{R}^{2}<0.0001$.


Figure 67. Residuals from Beverton-Holt Model versus proportion of hatchery origin spawners ( pHOS ) with fitted linear regression line, Pearson correlation coefficient (r), and P-value from two-sided $t$-test of slope of linear regression line using emigrant and spawner data from Methow River Summer Chinook Salmon, 2006-2018 (without 2012). $\mathrm{R}^{2}<0.0001$.


Figure 68. Residuals from Ricker Model versus proportion of hatchery origin spawners ( pHOS ) with fitted linear regression line, Pearson correlation coefficient ( r ), and P-value from two-sided $t$-test of slope of linear regression line using emigrant and spawner data from Methow River Summer Chinook Salmon, 2006-2018 (without 2012). $\mathrm{R}^{2}<0.0001$.

## Hypothesis H02.2.1.2

There was a very weak and non-significant negative association between $\ln$ (emigrants per redd) and the proportion of hatchery spawners ( pHOS ) for Methow River Summer Chinook Salmon (Pearson correlation coefficient $\mathrm{r}=-0.0641, \mathrm{P}=0.4216$ ) (Figure 69). The redd count was not associated with $\mathrm{pHOS}(\mathrm{P}=0.6767$, Figure 70) and there was no indication of post-spawning density dependence apparent from examination of the emigrants per redd plotted against the redd counts (Figure 71). Nevertheless, the relationship between emigrants per redd and pHOS was examined using the Ricker and Beverton-Holt stock-recruitment models in order to account for density dependence effects, with redds used in place of spawner counts. Both models were equivalent on the basis of $\mathrm{AICc} \operatorname{rank}(\triangle \mathrm{AICc}=0.001$ compared to Ricker model) and examination of model residuals (Table 17; Figure G8 - Figure G12). However, the Beverton-Holt model fit was not robust to variability in the data, as demonstrated by the inability to fit it to bootstrap samples and the wide and inadmissible asymptotic confidence intervals on the model parameters (Table 17). Thus, the Ricker model was preferred. The fitted Ricker model for emigrants per redd was (Table 17):

$$
\ln (R / S)=7.0724-0.2448 P_{\text {HоS }}-0.00004 * S
$$

The $95 \%$ bootstrap confidence interval for the regression coefficient for pHOS (i.e., $\beta_{2}$ in Equation (7)) was ( $-3.2640,2.4357$ ). Because this confidence interval included both positive and negative values, we concluded that there is no evidence of a negative association between pHOS and emigrants per redd for Methow Summer Chinook Salmon after adjusting for density dependence.


Figure 69. Emigrants per redd ( $\log$ scale) versus proportion of hatchery origin spawners ( pHOS ) for Methow River Summer Chinook Salmon, 2006-2018 (without 2012), with fitted linear regression line, Pearson correlation coefficient (r), and P-value from one-sided t-test of negative slope. $\mathrm{R}^{2}=0.0041$.


Figure 70. Redd count versus proportion of hatchery origin spawners (pHOS) for Methow River Summer Chinook Salmon, 2006-2018 (without 2012), with fitted linear regression line, Pearson correlation coefficient (r), and P-value from two-sided $t$-test of slope.


Figure 71. Emigrants per redd versus redd count for Methow River Summer Chinook Salmon, 2006-2018 (without 2012), with fitted linear regression line, Pearson correlation coefficient (r), and P-value from onesided $t$-test of negative slope. Negative slope indicates post-spawning density dependence.

Table 17. Fitted stock-recruitment models for emigrants per redd for Methow River Summer Chinook Salmon, 2006-2018 (without 2012), including effect ( $\beta_{2}$ ) of proportion of hatchery origin spawners ( pHOS ). Bootstrap confidence intervals based on $\geq 1,000$ bootstrap samples were provide for the Ricker model, and asymptotic confidence intervals were provided for the Beverton-Holt model. Model equations are defined in Equations (5) and (7). $\triangle \mathrm{AICc}$ represents change in AICc compared to model with minimum AICc.

| Model | AICc | $\Delta$ AICc | Parameter | Parameter <br> Estimates | $95 \%$ CI |
| :--- | :---: | :---: | :---: | :---: | :---: |
| Beverton-Holt | 31.64 | 0.001 | $\alpha$ | $31,205,045$ | $-881,383,950-943,794,030$ |
|  |  |  | $\beta$ | 26,564 | $-772,397-825,526$ |
|  |  |  | $\beta_{2}$ | 0.2468 | $-2.4111-2.9046$ |
| Ricker | 31.64 | 0.000 | $\ln (\alpha)$ | 7.0724 | $4.6169-8.2810$ |
|  |  |  | $\beta$ | 0.00004 | $-0.0016-0.0010$ |
|  |  |  | $\beta_{2}$ | 0.2448 | $-3.2640-2.4357$ |

## Methow River Summer Steelhead

Adult spawner, emigrant recruitment, pHOS , and redd count data from Summer Steelhead were available from the Methow River for the 2003-2015 brood years. Adult stock abundance (spawners) ranged from 1,105 in 2012 to 3,680 in 2010 and averaged 2,003, and emigrant recruitment ranged from 9,076 in 2003 to 33,739 in 2007 and averaged 18,154 (Figure 72, Figure 73). Redd counts averaged 1,084 (range $=591$ to 2,019 ), and pHOS averaged 0.81 (range $=0.58$ to 0.89 ). The 2007 brood year stood out as a possible outlier in emigrant count with a much higher count $(33,739)$ than in the other years (range $=9,076-25,845$ ) (Figure 73a).


Figure 72. Methow River Summer Steelhead boxplots for Stock (spawner abundance), Emigrants, Redd counts, and proportion of hatchery origin spawners (pHOS), 2003-2015. Box indicates interquartile range and thick horizontal bar is median. Whiskers indicate quartile $\pm 1.5 \times$ interquartile range.


Figure 73. Methow River Summer Steelhead annual Emigrant counts, Emigrant/Spawner (log scale), Emigrant/Redd (log scale), and proportion of hatchery origin spawners (pHOS), 2003-2015. Line indicates the linear trend. Shaded area is the estimated $95 \%$ confidence interval of the line.

## Hypothesis H02.2.1.1

A strong negative linear relationship was observed between Methow River Summer Steelhead spawners and emigrant recruitment, demonstrating evidence of density dependence in mortality between the spawning and emigrant life stages ( $\mathrm{P}=0.0002$; Figure 74). Thus, an attempt was made to fit the Smooth Hockey Stick, Beverton-Holt, and Ricker stock-recruitment models to the spawner and emigrant data. However, the Smooth Hockey Stick and Beverton-Holt modeling assumption of asymptotic growth in recruitment as spawner abundance increases was not supported by the data, as demonstrated by the locally smoothed curve (LOESS) in Figure 75, and neither the Smooth Hockey Stick model nor the Beverton-Holt model could be fit for the Methow River spawner-recruitment data. The Ricker model was fit to the Methow River data and estimated maximum recruitment at approximately 1,662 spawners (Figure 75). Although the Ricker stock-recruitment model could be fit to the Methow River spawner and emigrant data, the model residuals demonstrated negative autocorrelation at a time lag of 1 and 4 years (Figure H2), suggesting that the Ricker model does not adequately account for the population dynamics and that the model parameters may be biased. Additionally, high correlation was observed between the model parameter estimates, increasing uncertainty in the model predictions (Table 18).

The residuals from the Ricker model were compared to pHOS using correlation analysis. The estimated Pearson correlation coefficient between the Ricker model residuals and pHOS was 0.2222 and was not
significantly different from $0(\mathrm{P}=0.4656$; Figure 76). Additionally, a likelihood ratio test of the effect of pHOS on the Ricker model was non-significant ( $\mathrm{P}=0.3829$ ). The Beverton-Holt model could not be fit with pHOS as a variable. Analysis that omitted the possible outlier brood year of 2007 resulted in the same findings: only the Ricker model was approximately consistent with the data, and there was no evidence of a negative relationship between pHOS and juvenile productivity of emigrants.

Overall, there was no evidence that to suggest that higher proportions of hatchery spawners were associated with lower juvenile productivity for Summer Steelhead in the Methow River. However, there were only 13 years of data available, and the modeling assumptions were not well supported. Thus, it remains possible that there is a relationship between pHOS and juvenile productivity that are not observable using the available data and analyses.


Figure 74. Emigrants per spawner ( $\log$ scale) versus spawner abundance for Methow River Summer Steelhead, 2003-2015, with fitted linear regression line, Pearson correlation coefficient (r), and P-value from one-sided t -test of negative slope.


Figure 75. Stock and emigrant data with fitted LOESS curve and Ricker Model for Methow River Summer Steelhead, 2003-2015.

Table 18. Fitted stock-recruitment models for emigrants and stock (spawner) data from Methow River Summer Steelhead, 2003-2015. Confidence intervals were estimated using bootstrap with 3,000 bootstrap samples. Correlation $=$ Pearson correlation coefficient between parameter estimates from bootstrap samples. Model equations are defined in Equations (1), (2), and (3). $\triangle \mathrm{AICc}$ represents change in AICc compared to model with minimum AICc. *Only the Ricker model could be fit for Methow River data.

| Model* | AICc | $\Delta$ AICc | Parameter $^{\text {a }}$ | Parameter <br> Estimates | $95 \%$ CI | Correlation |
| :--- | :---: | :---: | :---: | :---: | :---: | :---: |
| Smooth Hockey <br> Stick | NA | NA | $\alpha$ | NA | NA | $\alpha, R_{\infty}:$ NA |
|  |  |  | $R_{\infty}$ | NA | NA |  |
| Beverton-Holt |  |  | NA | NA | $\alpha$ | NA |
|  |  |  | $\beta$ | NA | NA | $\alpha, \beta:$ NA |
| Ricker |  |  |  |  |  |  |
|  | 17.22 | NA | $\alpha$ | 30.8 | $19.3-48.9$ | $\alpha, \beta: 0.90$ |
|  |  |  | $\beta$ | 0.0006 | $0.0004-0.0008$ |  |

$\mathrm{a}=$ Estimated maximum recruitment parameters are: $R_{\infty}$ for the Smooth Hockey Stick model, $\alpha$ for the Beverton-Holt model, and $K=(\alpha / \beta) e^{-1}$ for the Ricker model.


Figure 76. Residuals from the Ricker Model versus proportion of hatchery origin spawners ( pHOS ) with fitted linear regression line, Pearson correlation coefficient (r), and P-value from two-sided t-test of slope of linear regression line using emigrant and spawner data from Methow River Summer Steelhead, 20032015. $\mathrm{R}^{2}=0.0494$.

## Hypothesis H02.2.1.2

There was no evidence of a negative association between emigrants per redd and the proportion of hatchery spawners ( pHOS ) for Methow River Summer Steelhead (Pearson correlation coefficient $\mathrm{r}=-0.1628$, $\mathrm{P}=0.2976$ ) (Figure 77). Also, the linear association between redd count and pHOS was non-significant ( $\mathrm{P}=0.2335$; Figure 78). However, comparison of emigrants per redd with redd count demonstrated strong
evidence of post-spawning density dependence ( $\mathrm{P}=0.0002$, Figure 79). Thus, the relationship between emigrants per redd and pHOS was examined using stock-recruitment models in order to account for density dependence effects, using redds in place of spawner abundance. The Beverton-Holt model resulted in an inadmissible (i.e., negative) estimate of the $\beta$ parameter, confirmed by examination of the likelihood surface. Thus, no results are provided for the Beverton-Holt model. The Ricker model was able to be fit and the residuals agreed moderately well with model assumptions (Figure H4, Figure H5). The fitted Ricker model for emigrants per redd was (Table 19):

$$
\ln (R / S)=3.3547+0.9921 * P_{\text {HOS }}-0.0012 * S
$$

The $95 \%$ bootstrap confidence interval for the effect of pHOS (i.e., $\beta_{2}$ in Equation (7)) was (-3.3407, 2.5691). Because the confidence interval for the effect of pHOS included both positive and negative values, we concluded that there was no evidence of a negative association between pHOS and emigrants per redd for Methow River Summer Steelhead whether or not an adjustment was made for density dependence. We came to the same conclusion when the possible outlier brood year (2007) was omitted from the data analysis. However, the small sample size, low degree of variability in the observed pHOS values, and potentially high level of measurement error in the observed data may have lowered the ability to detect a relationship between pHOS and emigrants per redd.


Figure 77. Emigrants per redd (log scale) versus proportion of hatchery origin spawners ( pHOS ) for Methow River Summer Steelhead, 2003-2015, with fitted linear regression line, Pearson correlation coefficient ( r ), and P -value from one-sided t -test of negative slope. $\mathrm{R}^{2}=0.0265$.


Figure 78. Redd count versus proportion of hatchery origin spawners ( pHOS ) for Methow River Summer Steelhead, 2003-2015, with fitted linear regression line, Pearson correlation coefficient (r), and P-value from two-sided $t$-test of slope.


Figure 79. Emigrants per redd (log scale) versus redd count for Methow River Summer Steelhead, 20032015, with fitted linear regression line, Pearson correlation coefficient (r), and P-value from one-sided ttest of negative slope. Negative slope indicates post-spawning density dependence.

Table 19. Fitted stock-recruitment models for emigrants per redd for Methow River Summer Steelhead, 2003-2015, including effect ( $\beta_{2}$ ) of proportion of hatchery origin spawners ( pHOS ). Confidence intervals were estimated using bootstrap with 3,000 bootstrap samples. Model equations are defined in Equations (5) and (7). $\triangle$ AICc represents change in AICc compared to model with minimum AICc. *Only the Ricker model could be fit for Methow River Summer Steelhead data.

| Model $^{*}$ | AICc | $\Delta$ AICc | Parameter | Parameter Estimates | $95 \%$ CI |
| :--- | :---: | :---: | :---: | :---: | :---: |
| Beverton-Holt | NA | NA | $\alpha$ | NA | NA |
|  |  |  | $\beta$ | NA | NA |
|  |  |  | $\beta_{2}$ | NA | NA |
|  |  |  |  |  |  |
| Ricker | 19.92 | NA | $\ln (\alpha)$ | 3.3547 | $1.5579-6.1456$ |
|  |  |  | $\beta$ | 0.0012 | $0.0006-0.0018$ |
|  |  |  | $\beta_{2}$ | -0.9921 | $-3.3407-2.5691$ |

## Twisp River Summer Steelhead

Adult spawner, emigrant recruitment, pHOS , and redd count data from Summer Steelhead were available from the Twisp River for the 2003-2015 brood years. Adult stock abundance (spawners) ranged from 143 in 2007 to 1,204 in 2003 and averaged 532, and emigrant recruitment ranged from 3,264 in 2008 to 13,669 in 2007 and averaged 6,133 (Figure 80, Figure 81). Redd counts averaged 272 (range $=82$ to 696), and pHOS averaged 0.69 (range $=0.48$ to 0.89 ). The 2007 brood year stood out as a possible outlier in emigrant count, with a much higher emigrant count $(13,669)$ than any of the other years (range $=3,264-7,467$ ) (Figure 81a). Additionally, pHOS generally declined over the course of the data collection, partly as a result of a policy change in 2010 to target pHOS values of 0.5 (Figure 81d).


Figure 80. Twisp River Summer Steelhead boxplots for Stock (spawner abundance), Emigrants, Redd counts, and proportion of hatchery origin spawners (pHOS), 2003-2015. Box indicates interquartile range and thick horizontal bar is median. Whiskers indicate quartile $\pm 1.5 \times$ interquartile range.


Figure 81. Twisp River Summer Steelhead annual Emigrant counts, Emigrant/Spawner (log scale), Emigrant/Redd (log scale), and proportion of hatchery origin spawners (pHOS), 2003-2015. Line indicates the linear trend. Shaded area is the estimated $95 \%$ confidence interval of the line.

## Hypothesis H02.2.1.1

A strong negative linear relationship was observed between Twisp River Summer Steelhead spawners and emigrant recruitment, providing evidence of density dependence in mortality between the spawning and emigrant life stages ( $\mathrm{P}=0.0001$; Figure 82). Thus, an attempt was made to fit the Smooth Hockey Stick, Beverton-Holt, and Ricker stock-recruitment models to the spawner and emigrant data. However, the Smooth Hockey Stick and Beverton-Holt modeling assumption of asymptotic growth in recruitment as spawner abundance increases was not supported by the data, as demonstrated by the locally smoothed curve in Figure 83, and neither the Smooth Hockey Stick model nor the Beverton-Holt model could be fit for the Twisp River spawner-recruitment data. The Ricker model was fit to the Twisp River data and estimated maximum recruitment at approximately 410 spawners (Figure 83). Although the Ricker stock-recruitment model could be fit to the Twisp River spawner and emigrant data, the model residuals did not clearly meet the assumption of lognormal errors (Figure I1), suggesting that the model parameters may be biased.

The residuals from the Ricker model were compared to pHOS using correlation analysis. The estimated Pearson correlation coefficient between the Ricker model residuals and pHOS ( $\mathrm{r}=0.2553$ ) was not significantly different from 0 ( $\mathrm{P}=0.3999$; Figure 84). A simpler analysis that compared the average Ricker model residuals between the pre-2010 period, when pHOS was not restricted ("control"), and the 2010+ period, when pHOS was targeted at 0.5 ("treatment"), also found no difference in model residuals between the two management periods $(t$-test $=-0.4717, \mathrm{df}=11, \mathrm{P}=0.6463$; Figure 85 ). Additionally, a likelihood
ratio test of the effect of pHOS on the Ricker model was non-significant ( $\mathrm{P}=0.3071$ ). The Beverton-Holt model could not be fit with pHOS as a variable.

The 2007 brood year was noted as a possible outlier in emigrant count, so the analysis was redone without the 2007 brood year. The results from the Ricker model were consistent with the full data set (i.e., nonsignificant relationship between residuals and $\mathrm{pHOS}, \mathrm{P}=0.2868$, and also between residuals and pHOS management periods, $\mathrm{P}=0.1728$ ), and the Beverton-Holt and Smooth Hockey Stick models could not be fit to the data (Figure 83, Table 20, Figure 86, Figure 87).

Overall, there was no evidence that to suggest that higher proportions of hatchery spawners were associated with lower juvenile productivity for Summer Steelhead in the Twisp River. However, with only 13 years of data, it remains possible that there is a relationship between pHOS and juvenile productivity that are not observable using the available data and analyses.


Figure 82. Emigrants per spawner (log scale) versus spawner abundance for Twisp River Summer Steelhead, 2003-2015, with and without brood year 2007, with fitted linear regression line, Pearson correlation coefficient ( r ), and P -value from one-sided t -test of negative slope.


Figure 83. Stock and emigrant data with fitted LOESS curve and Ricker Model for Twisp River Summer Steelhead, 2003-2015, with and without brood year 2007.

Table 20. Fitted stock-recruitment models for emigrants and stock (spawner) data from Twisp River Summer Steelhead, 2003-2015. Confidence intervals were estimated using bootstrap with at least 2,000 bootstrap samples. Correlation $=$ Pearson correlation coefficient between parameter estimates from bootstrap samples. Model equations are defined in Equations (1), (2), and (3). $\Delta$ AICc represents change in AICc compared to model with minimum AICc. *Only the Ricker model could be fit for Twisp River data. The Ricker model was fit both with and without data from the 2007 brood year. AICc should not be compared between the two Ricker models.

| Model* | AICc | $\Delta$ AICc | Parameter $^{\mathrm{a}}$ | Parameter <br> Estimates | $95 \%$ CI | Correlation |
| :--- | :---: | :---: | :---: | :---: | :---: | :---: |
| Smooth Hockey | NA | NA | $\alpha$ | NA | NA | $\alpha, R_{\infty}:$ NA |
| Stick |  |  | $R_{\infty}$ | NA | NA |  |
|  |  |  |  |  |  |  |
| Beverton-Holt | NA | NA | $\alpha$ | NA | NA | $\alpha, \beta:$ NA |
|  |  |  | $\beta$ | NA | NA |  |
| Ricker with 2007 | 22.50 | NA | $\alpha$ | 46.3 | $29.2-76.2$ | $\alpha, \beta: 0.87$ |
|  |  |  | $\beta$ | 0.0024 | $0.0016-0.0032$ |  |
| Ricker without 2007 | 9.44 | NA | $\alpha$ | 6,983 | $5,667-9,221$ |  |
|  |  |  | $\beta$ | 31.8 | $22.9-43.1$ | $\alpha, \beta: 0.90$ |
|  |  |  | $K$ | 6,054 | $0.0015-0.0025$ |  |

$\mathrm{a}=$ Estimated maximum recruitment parameters are: $R_{\infty}$ for the Smooth Hockey Stick model, $\alpha$ for the BevertonHolt model, and $K=(\alpha / \beta) e^{-1}$ for the Ricker model.


Figure 84. Residuals from the Ricker Model versus proportion of hatchery origin spawners ( pHOS ) with fitted linear regression line, Pearson correlation coefficient (r), and P-value from two-sided t-test of slope of linear regression line using emigrant and spawner data from Twisp River Summer Steelhead, 20032015 , including brood year 2007. $\mathrm{R}^{2}=0.0652$.

Ricker Model Residuals vs. pHOS Period


Figure 85. Distribution of residuals from the Ricker Model categorized by pHOS management period: Control $=$ pre-2010, when pHOS levels were not restricted, and Treatment $=2010-2015$, when pHOS levels were targeted at 0.5 . P -value is from two-sided t -test: $t$-test $=-0.4717, \mathrm{df}=11$. Data used were emigrant and spawner counts from Twisp River Summer Steelhead, 2003-2015, including brood year 2007.


Figure 86. Residuals from the Ricker Model versus proportion of hatchery origin spawners ( pHOS ) with fitted linear regression line, Pearson correlation coefficient (r), and P-value from two-sided t-test of slope of linear regression line using emigrant and spawner data from Twisp River Summer Steelhead, 20032015, without brood year 2007. $\mathrm{R}^{2}=0.1124$.

Ricker Model Residuals vs. pHOS Period


Figure 87. Distribution of residuals from the Ricker Model categorized by pHOS management period: Control $=$ pre-2010, when pHOS levels were not restricted, and Treatment $=2010-2015$, when pHOS levels were targeted at 0.5 . P-value is from two-sided $t$-test: $t$-test $=1.4683, \mathrm{df}=10$. Data used were emigrant and spawner counts from Twisp River Summer Steelhead, 2003-2015, without brood year 2007.

## Hypothesis H02.2.1.2

There was no evidence of a negative association between emigrants per redd and the proportion of hatchery spawners ( pHOS ) for Twisp River Summer Steelhead using the full data set that included brood year 2007 (Pearson correlation coefficient $\mathrm{r}=-0.3180, \mathrm{P}=0.1449$ ) (Figure 88). The linear association between redd count and pHOS was significant at the $10 \%$ level ( $\mathrm{P}=0.0723$; Figure 89 ), and comparison of emigrants per redd with redd count demonstrated strong evidence of post-spawning density dependence ( $\mathrm{P}=0.0001$,

Figure 90). The relationship between emigrants per redd and pHOS was examined using stock-recruitment models in order to account for density dependence effects, using redds in place of spawner abundance. The Beverton-Holt model could not be fit to the data; examination of the likelihood indicated an optimum model fit for a negative value of the $\beta$ model parameter, which is inconsistent with model interpretation (Figure I6). The Ricker model was able to be fit but the residuals agreed poorly with model assumptions (Figure I4, Figure I5). The fitted Ricker model for emigrants per redd was (Table 21):

$$
\ln (R / S)=3.9305+0.7680 * P_{\text {НоS }}-0.0045 * S
$$

The $95 \%$ bootstrap confidence interval for the effect of pHOS (i.e., $\beta_{2}$ in Equation (7)) was ( -2.3366 , 2.1443). Because the confidence interval for the effect of pHOS included both positive and negative values, we concluded that there was no evidence of a negative association between pHOS and emigrants per redd for Twisp River Summer Steelhead whether or not an adjustment was made for density dependence. However, the model assumptions were not well met, demonstrating an overall lack of fit. The small sample size and potentially high level of measurement error in the observed data may have contributed to the lack of model fit, and likewise may have lowered the ability to detect a relationship between pHOS and emigrants per redd using the available data.

When the potential outlier brood year of 2007 was omitted from the data set, there was a negative association between emigrants per redd and pHOS when density dependence was not accounted for ( $\mathrm{P}=0.0009$, Figure 91 ). However, there was significant density dependent mortality observed between the redd count and emigrant count ( $\mathrm{P}<0.0001$, Figure 90 ), which may have accounted for the association between emigrants per redd and pHOS . The Ricker model estimated that the effect of pHOS on the reddemigrant relationship was non-significant, with a $95 \%$ bootstrap confidence interval of ( $-2.3915,1.0898$ ) (Table 22). Without the 2007 brood year, the Beverton-Holt model relating emigrants to redd counts was able to be fit when pHOS was in the model and pHOS was associated with lower emigrants per redd, demonstrated by the $95 \%$ bootstrap confidence interval for $\beta_{2}$ that was entirely greater than 0 (i.e., ( 0.0700 , 1.9685); Table 22). However, the bootstrap confidence intervals for the other Beverton-Holt model parameters were notably wide: the confidence interval of the asymptotic maximum recruitment $(\alpha)$ included values 5 times the maximum recruitment observed, and the confidence interval for the redd counts predicted to generate half the maximum recruitment ( $\beta$ ) included negative values (Table 22). The apparently poor fit of the Beverton-Holt model raises questions about the validity of its finding of a statistically significant pHOS effect.

Overall, the evidence for a negative association between the proportion of hatchery origin spawners ( pHOS ) and juvenile productivity in emigrants was not strong. There was no such evidence based on the full data set that included the 2007 brood year; however, one of the two stock-recruitment models could not be fit to the data because of misalignment of modeling assumptions with the data, and the data set was small (only 13 brood years). When the 2007 brood year was treated as an outlier and omitted from analysis, a negative association was observed between emigrants per redd and pHOS , but it disappeared when density dependence was accounted for using the Ricker model. The alternative stock-recruitment model (BevertonHolt) maintained an effect of pHOS but did not appear to fit the data well, indicating possible bias in the estimate of the pHOS effect. The small sample size available may have lowered the ability to detect an effect of pHOS on juvenile productivity. Additionally, a negative trend in pHOS from nearly 0.90 in 2003 to nearly 0.50 in 2015 raises the possibility that any pHOS effect actually masks a temporal trend in juvenile productivity (Figure 81).


Figure 88. Emigrants per redd (log scale) versus proportion of hatchery origin spawners (pHOS) for Twisp River Summer Steelhead, 2003-2015, including brood year 2007, with fitted linear regression line, Pearson correlation coefficient (r), and P-value from one-sided $t$-test of negative slope. $\mathrm{R}^{2}=0.1011$.


Figure 89. Redd count versus proportion of hatchery origin spawners ( pHOS ) for Twisp River Summer Steelhead, 2003-2015, with and without brood year 2007, with fitted linear regression line, Pearson correlation coefficient (r), and P-value from two-sided t -test of slope.


Figure 90. Emigrants per redd (log scale) versus redd count for Twisp River Summer Steelhead, 20032015, with and without brood year 2007, with fitted linear regression line, Pearson correlation coefficient (r), and P-value from one-sided t -test of negative slope. Negative slope indicates post-spawning density dependence.

Table 21. Fitted stock-recruitment models for emigrants per redd for Twisp River Summer Steelhead, 20032015, including brood year 2007 and the effect ( $\beta_{2}$ ) of proportion of hatchery origin spawners ( pHOS ). Confidence intervals were estimated using bootstrap with 3,000 bootstrap samples. Model equations are defined in Equations (5) and (7). $\triangle$ AICc represents change in AICc compared to model with minimum AICc. *Only the Ricker model could be fit for Twisp River Summer Steelhead data.

| Model $^{*}$ | AICc | $\Delta$ AICc | Parameter | Parameter Estimates | $95 \%$ CI |
| :--- | :--- | :--- | :---: | :---: | :---: |
| Beverton-Holt | NA | NA | $\alpha$ | NA | NA |
|  |  |  | $\beta$ | NA | NA |
|  |  |  | $\beta_{2}$ | NA | NA |
| Ricker | 25.81 |  | NA | $\ln (\alpha)$ | 3.9305 |
|  |  | $\beta$ | 0.0045 | $3.3009-5.0401$ |  |
|  |  |  | $\beta_{2}$ | -0.7680 | $0.0015-0.0067$ |
|  |  |  |  |  | $-2.3366-2.1443$ |



Figure 91. Emigrants per redd versus proportion of hatchery origin spawners (pHOS) for Twisp River Summer Steelhead, 2003-2015, omitting brood year 2007, with fitted linear regression line, Pearson correlation coefficient (r), and P-value from one-sided t-test of negative slope. $\mathrm{R}^{2}=0.6417$.

Table 22. Fitted stock-recruitment models for emigrants per redd for Twisp River Summer Steelhead, 20032015, omitting brood year 2007 and including the effect $\left(\beta_{2}\right)$ of proportion of hatchery origin spawners (pHOS). Confidence intervals were estimated using bootstrap with 3,000 bootstrap samples. Model equations are defined in Equations (5) and (7). $\Delta \mathrm{AICc}$ represents change in AICc compared to model with minimum AICc.

| Model | AICc | $\Delta$ AICc | Parameter | Parameter Estimates | $95 \% \mathrm{CI}$ |
| :--- | :---: | :--- | :---: | :---: | :---: |
| Beverton-Holt | 10.55 | 0.24 | $\alpha$ | 12,008 | $5,158-40,066$ |
|  |  |  | $\beta$ | 46.4 | $-31.2-275.2$ |
|  |  |  | $\beta_{2}$ | 0.9259 | $0.0700-1.9685$ |
|  |  |  |  |  |  |
| Ricker | 0.31 | NA | $\ln (\alpha)$ | 4.5239 | $3.8033-5.1740$ |
|  |  |  | $\beta$ | 0.0027 | $0.0007-0.0049$ |
|  |  |  | $\beta_{2}$ | 1.0323 | $-2.3915-1.0898$ |

## Discussion

Our investigation of whether a higher proportion of hatchery origin spawners (pHOS) may result in lowered juvenile productivity was largely inconclusive. For most populations studied, no evidence of an effect of pHOS was observed. For the single population that demonstrated a possible negative effect of pHOS on juvenile productivity (Twisp River Summer Steelhead), the evidence was weak: when density dependence was accounted for, the perceived negative association between pHOS and juveniles per redd was no longer observed. Additionally, the significant result was observed only when a possible outlier in emigrant counts was omitted, which may not be justified depending on the cause of the unusual emigrant count for that brood year. Thus, even when some evidence of a negative effect of pHOS was found, the evidence was particularly weak.

The lack of an observed effect of pHOS does not necessarily mean that such an effect does not exist, however. Multiple factors combined to lower the ability to detect an effect, should it exist. The most obvious factor was the small size of most of the data sets. Most of the populations had data for fewer than 20 brood years, resulting in lowered opportunity to observe the full range of variability in stock-recruitment dynamics
in response to changes in hatchery proportion of spawners and lower statistical power to detect an effect. Then too, many populations showed low contrast in the observed pHOS values. A wider range of hatchery proportion values would make detecting an effect more feasible. In some populations (e.g., Twisp River Steelhead, Methow River Steelhead), pHOS has declined almost consistently since the start of data collection, which means any perceived effect of pHOS would be entirely confounded with temporal changes in juvenile productivity. This risk is increased by the observational nature of the pHOS data. An experiment that purposely manages hatchery origin spawners at planned pHOS levels would improve the ability to detect an effect on juvenile productivity.

Some data sets studied also demonstrated low contrast in the spawner or redd counts observed across brood years. The stock-recruitment models require observations at both low and high levels of spawners. This requirement means that low variability in spawner (or redd count) data reduces the quality of fit of the stock-recruitment models and in some cases makes them impossible to be fit altogether. A lack of data at low spawner levels may have lowered the ability to fit the Smooth Hockey Stick model, which requires data at low spawner levels to estimate the model parameter associated with depensation ( $\alpha$ ). Examination of the likelihood surface for the Smooth Hockey Stick model for some populations demonstrated moderate information in maximum recruitment but a complete lack of information in the $\alpha$ parameter, interpreted as the slope of the stock-recruitment curve at very low numbers of spawners (for example, see the flat contour curves in the dimension of the $\alpha$ parameter in Figure 92). Then again, several populations failed to demonstrate a stable maximum (or simply high) juvenile recruitment pattern at high levels of spawners, which is assumed by both the Smooth Hockey Stick and Beverton-Holt models. Again, examination of the likelihood surface in these cases indicated either a lack of information in model parameters or else model parameters optimized at inadmissible values (e.g., negative spawner counts necessary to generate half the asymptotic maximum recruitment, Figure 93). Although the Smooth Hockey Stick stock-recruitment model was the preferred analysis framework, it appeared poorly suited to the stock and juvenile recruitment data available for many populations.

For all three stock-recruitment models considered, there was often high uncertainty in model parameter estimates even when the models could be fit to the data. This was indicated by wide confidence intervals that often included maximum smolt or emigrant values far beyond the range of observed data and by high correlation in model parameter estimates. High correlation lowers the ability to distinguish between different model fits and increases uncertainty in the results; it arises from lack of contrast in the data or violation of modeling assumptions.

In most cases there was no evidence of a time lag in juvenile recruitment per spawner as represented by autocorrelation in the residuals from the stock-recruitment models. However, a negative autocorrelation was observed with a time lag of three years for Twisp River Spring Chinook Salmon and a time lag of one and four years for Methow River Summer Steelhead. Such negative autocorrelations may be observed when a high level of recruitment one year is habitually followed by a low level of recruitment after a consistent delay (e.g., 3 years). Such a pattern may be expected for populations with consistent age structure among spawners, paired with density-dependent mortality between spawning and juvenile recruit surveys. In such cases, the Smooth Hockey Stick, Beverton-Holt, and Ricker models are unlikely to adequately represent all important features of the population dynamics; an age-structured model may be required. The value of pHOS may also be affected by such autocorrelation. Thus, such factors should be accounted for in order to detect a true effect of pHOS on juvenile productivity beyond age- and density-dependent components of the population dynamics.

A final complication in assessing the stock-recruitment data for an effect of pHOS is data quality. The stock-recruitment models all assume low levels of measurement error in both spawners or redd counts and smolts or emigrants. The higher the level of measurement error, the lower the ability to model the dynamics well enough to detect an effect. Given all the complicating factors identified here, only the most extreme effect of pHOS on juvenile productivity could have been detected. Thus, we caution against concluding
that such an effect is truly absent. Instead, we recommend that future data collection efforts attempt to minimize measurement error and increase the contrast in pHOS levels. We recommend that a planned experiment using deliberately chosen pHOS levels be implemented in order to achieve the necessary contrast and remove confounding with temporal and age-structured processes. Finally, we recommend that additional stock-recruitment models be considered that may better match the observed patterns in stock and juvenile recruitment data for populations of Chinook Salmon and Steelhead in tributaries affected by hatchery origin spawners.


Figure 92. Negative log-likelihood surface and contour curves for the Smooth Hockey Stick model of emigrants and spawner counts. Lighter regions indicate higher likelihood values (lower negative loglikelihood values). The estimated parameter values occur where the negative log-likelihood is minimized. Data set = Twisp River Summer Steelhead, 2003-2015.


Figure 93. Negative log-likelihood surface and contour curves for Beverton-Holt model of emigrants, redd counts, and pHOS , using $\beta_{2}=-0.1160$. Lighter regions indicate higher likelihood values (lower negative log-likelihood values). The estimated parameter values occur where the negative log-likelihood is minimized. The green triangle (falls below $\mathrm{b}=0$ line) indicates the parameter set that optimizes the likelihood. Data set = Twisp River Summer Steelhead, 2003-2015.

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## Appendices: Evaluation of Modeling Assumptions

## Appendix A: Chiwawa River Spring Chinook Salmon



Figure A1. Normal quantile-quantile plot of residual from Smooth Hockey Stick model of smolt recruitment as a function of spawner abundance (stock) for Chiwawa River Spring Chinook, 1991-2017. Points should lie on the line under assumption of normal errors. Shapiro-Wilk test of normality of residuals: $\mathrm{W}=0.9643$, $\mathrm{P}=0.4604$.


Figure A2. Autocorrelation plot of residuals for Smooth Hockey Stick model of smolt recruitment as a function of spawner abundance (stock) for Chiwawa River Spring Chinook, 1991-2017. Blue lines mark $95 \%$ confidence interval under assumption of no autocorrelation.


Figure A3. Correlation plot of bootstrap parameter estimates from the Smooth Hockey Stick model of smolt recruitment as a function of spawner abundance (stock) for Chiwawa River Spring Chinook, 1991-2017, based on 3,000 bootstrap samples.


Figure A4. Normal quantile-quantile plot of residual from Beverton-Holt model of smolt recruitment as a function of spawner abundance (stock) for Chiwawa River Spring Chinook, 1991-2017. Points should lie on the line under assumption of normal errors. Shapiro-Wilk test of normality of residuals: $\mathrm{W}=0.9672$, $\mathrm{P}=0.5293$.

## Chiwawa Spring Chinook: Beverton-Holt Model



Figure A5. Autocorrelation plot of residuals for Beverton-Holt model of smolt recruitment as a function of spawner abundance (stock) for Chiwawa River Spring Chinook, 1991-2017. Blue lines mark 95\% confidence interval under assumption of no autocorrelation.


Figure A6. Normal quantile-quantile plot of residual from Ricker model of smolt recruitment as a function of spawner abundance (stock) for Chiwawa River Spring Chinook, 1991-2017. Points should lie on the line under assumption of normal errors. Shapiro-Wilk test of normality of residuals: $\mathrm{W}=0.9788, \mathrm{P}=0.8347$.

## Chiwawa Spring Chinook: Ricker Model



Figure A7. Autocorrelation plot of residuals for Ricker model of smolt recruitment as a function of spawner abundance (stock) for Chiwawa River Spring Chinook, 1991-2017. Blue lines mark $95 \%$ confidence interval under assumption of no autocorrelation.


Figure A8. Residual plots for linear model of smolts per redd as a function of pHOS for Chiwawa River Spring Chinook, 1991-2017. Shapiro-Wilk test of normality of residuals: $\mathrm{W}=0.9672, \mathrm{P}=0.5302$.

Table A1. Pearson correlation coefficient (below diagonal) and two-sided P-value (above diagonal) for Beverton-Holt stock-recruitment model of smolts per redd for Chiwawa River Spring Chinook, 1991-2017. Model form is: $\ln (R / S)=\ln (\alpha)-\beta_{2} P_{\text {Hos }}-\ln (\beta+S)$. Correlation computed from bootstrap (3,000 bootstrap samples).

| Parameter | $\alpha$ | $\beta$ | $\beta_{2}$ |
| :---: | :--- | :--- | :--- |
| $\alpha$ |  | $<0.0001$ | $<0.0001$ |
| $\beta$ | 0.7954 |  | $<0.0001$ |
| $\beta_{2}$ | 0.7776 | 0.3714 |  |

Table A2. Pearson correlation coefficient (below diagonal) and two-sided P-value (above diagonal) for Ricker stock-recruitment model of smolts per redd for Chiwawa River Spring Chinook, 1991-2017. Model form is: $\ln (R / S)=\beta_{0}-\beta S-\beta_{2} P_{\text {HOS }}$. Correlation computed from bootstrap (3,000 bootstrap samples).

| Parameter | $\beta_{0}$ | $\beta$ | $\beta_{2}$ |
| :---: | :--- | :--- | :--- |
| $\beta_{0}$ |  | $<0.0001$ | $<0.0001$ |
| $\beta$ | 0.3198 |  | $<0.0001$ |
| $\beta_{2}$ | 0.7427 | -0.3097 |  |

Chiwawa Spring Chinook: Beverton-Holt Model - Juveniles Per Redd


Figure A9. Residual plots for Beverton-Holt model of smolts per redd as a function of redd count and pHOS for Chiwawa River Spring Chinook, 1991-2017. Shapiro-Wilk test of normality of residuals: $\mathrm{W}=0.9608$, $\mathrm{P}=0.3864$.


Figure A10. Normal quantile-quantile plot of residual from Beverton-Holt model of smolts per redd as a function of redd count and pHOS for Chiwawa River Spring Chinook, 1991-2017. Points should lie on the line under assumption of normal errors. Shapiro-Wilk test of normality of residuals: $\mathrm{W}=0.9608, \mathrm{P}=0.3864$.


Figure A11. Autocorrelation plot of residuals for Beverton-Holt model of smolts per redd as a function of redd count and pHOS for Chiwawa River Spring Chinook, 1991-2017. Blue lines mark 95\% confidence interval under assumption of no autocorrelation.


Figure A12. Residual plots for Ricker model of smolts per redd as a function of redd count and pHOS for Chiwawa River Spring Chinook, 1991-2017. Shapiro-Wilk test of normality of residuals: W=0.9801, $\mathrm{P}=0.8656$.


Figure A13. Autocorrelation plot of residuals for Ricker model of smolts per redd as a function of redd count and pHOS for Chiwawa River Spring Chinook, 1991-2017. Blue lines mark 95\% confidence interval under assumption of no autocorrelation.

## Appendix B: Nason Creek Spring Chinook Salmon



Figure B1. Normal quantile-quantile plot of residual from Ricker model of smolt recruitment as a function of spawner abundance (stock) for Nason Creek Spring Chinook, 2002-2017, including brood year 2014. Points should lie on the line under assumption of normal errors. Shapiro-Wilk test of normality of residuals: $\mathrm{W}=0.8152, \mathrm{P}=0.0044$.


Figure B2. Autocorrelation plot of residuals for Ricker model of smolt recruitment as a function of spawner abundance (stock) for Nason Creek Spring Chinook, 2002-2017, including brood year 2014. Blue lines mark $95 \%$ confidence interval under assumption of no autocorrelation.


Figure B3. Normal quantile-quantile plot of residual from Ricker model of smolt recruitment as a function of spawner abundance (stock) for Nason Creek Spring Chinook, 2002-2017 without brood year 2014. Points should lie on the line under assumption of normal errors. Shapiro-Wilk test of normality of residuals: $\mathrm{W}=0.9478, \mathrm{P}=0.4906$.


Figure B4. Autocorrelation plot of residuals for Ricker model of smolt recruitment as a function of spawner abundance (stock) for Nason Creek Spring Chinook, 2002-2017 without brood year 2014. Blue lines mark $95 \%$ confidence interval under assumption of no autocorrelation.


Figure B5. Residual plots for linear model of smolts per redd as a function of pHOS for Nason Creek Spring Chinook, 2002-2017 (including brood year 2014). Shapiro-Wilk test of normality of residuals: $\mathrm{W}=0.8923$, $\mathrm{P}=0.0606$.

Table B1. Pearson correlation coefficient (below diagonal) and two-sided P-value (above diagonal) for Ricker stock-recruitment model of smolts per redd for Nason Creek Spring Chinook, 2002-2017, with and without brood year 2014. Model form is: $\ln (R / S)=\beta_{0}-\beta S-\beta_{2} P_{H O S}$. Correlation computed from bootstrap ( 3,000 bootstrap samples).

| With BY 2014 |  | $\beta$ |  |
| :---: | :--- | :--- | :--- |
| Parameter | $\beta_{0}$ | $\beta$ | $\beta_{2}$ |
| $\beta_{0}$ | 0.3600 | $<0.0001$ | $<0.0001$ |
| $\beta$ | 0.8629 | -0.0925 | $<0.0001$ |
| $\beta_{2}$ |  |  |  |
|  |  | $\beta$ |  |
| Without BY 2014 | $\beta_{0}$ | $<0.0001$ |  |
| Parameter |  |  | $\beta_{2}$ |
| $\beta_{0}$ | 0.3509 | -0.2354 | $<0.0001$ |
| $\beta$ | 0.7692 |  |  |
| $\beta_{2}$ |  |  |  |



Figure B6. Residual plots for Ricker model of smolts per redd as a function of redd count and pHOS for Nason Creek Spring Chinook, 1991-2017, with brood year 2014. Shapiro-Wilk test of normality of residuals: $\mathrm{W}=0.8831, \mathrm{P}=0.0434$.

Nason Spring Chinook: Ricker Model - Juveniles Per Redd


Figure B7. Autocorrelation plot of residuals for Ricker model of smolts per redd as a function of redd count and pHOS for Nason Creek Spring Chinook, 2002-2017, with brood year 2014. Blue lines mark 95\% confidence interval under assumption of no autocorrelation.


Figure B8. Residual plots for Ricker model of smolts per redd as a function of redd count and pHOS for Nason Creek Spring Chinook, 1991-2017, without brood year 2014. Shapiro-Wilk test of normality of residuals: $\mathrm{W}=0.8533, \mathrm{P}=0.0193$.


Figure B9. Autocorrelation plot of residuals for Ricker model of smolts per redd as a function of redd count and pHOS for Nason Creek Spring Chinook, 2002-2017, without brood year 2014. Blue lines mark 95\% confidence interval under assumption of no autocorrelation.

## Appendix C: White River Spring Chinook Salmon



Figure C1. Normal quantile-quantile plot of residual from the Smooth Hockey Stick model of smolt recruitment as a function of spawner abundance (stock) for White River Spring Chinook, 2005-2017, including brood year 2014. Points should lie on the line under assumption of normal errors. Shapiro-Wilk test of normality of residuals: $\mathrm{W}=0.8887, \mathrm{P}=0.0936$.

White Spring Chinook: Smooth Hockey Stick Model


Figure C2. Autocorrelation plot of residuals for the Smooth Hockey Stick model of smolt recruitment as a function of spawner abundance (stock) for White River Spring Chinook, 2005-2017, including brood year 2014. Blue lines mark $95 \%$ confidence interval under assumption of no autocorrelation.


Figure C3. Correlation plot of bootstrap parameter estimates from the Smooth Hockey Stick model of smolt recruitment as a function of spawner abundance (stock) for White River Spring Chinook, 2005-2017, including brood year 2014, based on 3,291 bootstrap samples.


Figure C4. Normal quantile-quantile plot of residual from the Smooth Hockey Stick model of smolt recruitment as a function of spawner abundance (stock) for White River Spring Chinook, 2005-2017 without brood year 2014. Points should lie on the line under assumption of normal errors. Shapiro-Wilk test of normality of residuals: $\mathrm{W}=0.9484, \mathrm{P}=0.6144$.


Figure C5. Autocorrelation plot of residuals for the Smooth Hockey Stick model of smolt recruitment as a function of spawner abundance (stock) for White River Spring Chinook, 2005-2017 without brood year 2014. Blue lines mark $95 \%$ confidence interval under assumption of no autocorrelation.


Figure C6. Correlation plot of bootstrap parameter estimates from the Smooth Hockey Stick model of smolt recruitment as a function of spawner abundance (stock) for White River Spring Chinook, 2005-2017, without brood year 2014, based on 3,618 bootstrap samples.


Figure C7. Normal quantile-quantile plot of residual from Beverton-Holt model of smolt recruitment as a function of spawner abundance (stock) for White River Spring Chinook, 2005-2017, including brood year 2014. Points should lie on the line under assumption of normal errors. Shapiro-Wilk test of normality of residuals: $\mathrm{W}=0.8893, \mathrm{P}=0.0954$.

White Spring Chinook: Beverton-Holt Model


Figure C8. Autocorrelation plot of residuals for Beverton-Holt model of smolt recruitment as a function of spawner abundance (stock) for White River Spring Chinook, 2005-2017, including brood year 2014. Blue lines mark $95 \%$ confidence interval under assumption of no autocorrelation.


Figure C9. Normal quantile-quantile plot of residual from Beverton-Holt model of smolt recruitment as a function of spawner abundance (stock) for White River Spring Chinook, 2005-2017 without brood year 2014. Points should lie on the line under assumption of normal errors. Shapiro-Wilk test of normality of residuals: $\mathrm{W}=0.9501, \mathrm{P}=0.6384$.

White Spring Chinook: Beverton-Holt Model


Figure C10. Autocorrelation plot of residuals for Beverton-Holt model of smolt recruitment as a function of spawner abundance (stock) for White River Spring Chinook, 2005-2017 without brood year 2014. Blue lines mark $95 \%$ confidence interval under assumption of no autocorrelation.


Figure C11. Normal quantile-quantile plot of residual from Ricker model of smolt recruitment as a function of spawner abundance (stock) for White River Spring Chinook, 2005-2017, including brood year 2014. Points should lie on the line under assumption of normal errors. Shapiro-Wilk test of normality of residuals: $\mathrm{W}=0.8826, \mathrm{P}=0.0774$.


Figure C12. Autocorrelation plot of residuals for Ricker model of smolt recruitment as a function of spawner abundance (stock) for White River Spring Chinook, 2005-2017, including brood year 2014. Blue lines mark $95 \%$ confidence interval under assumption of no autocorrelation.


Figure C13. Normal quantile-quantile plot of residual from Ricker model of smolt recruitment as a function of spawner abundance (stock) for White River Spring Chinook, 2005-2017 without brood year 2014. Points should lie on the line under assumption of normal errors. Shapiro-Wilk test of normality of residuals: $\mathrm{W}=0.9632, \mathrm{P}=0.8287$.


Figure C14. Autocorrelation plot of residuals for Ricker model of smolt recruitment as a function of spawner abundance (stock) for White River Spring Chinook, 2005-2017 without brood year 2014. Blue lines mark $95 \%$ confidence interval under assumption of no autocorrelation.


Figure C15. Residual plots for linear model of smolts per redd as a function of pHOS for White River Spring Chinook, 2005-2017 (including brood year 2014). Shapiro-Wilk test of normality of residuals: $\mathrm{W}=0.9443, \mathrm{P}=0.5150$.

Table C1. Pearson correlation coefficient (below diagonal) and two-sided P-value (above diagonal) for Beverton-Holt stock-recruitment model of smolts per redd for White River Spring Chinook, 2005-2017, with and without brood year 2014. Model form is: $\ln (R / S)=\ln (\alpha)-\beta_{2} P_{H O S}-\ln (\beta+S)$. Correlation computed from bootstrap ( $\geq 3,000$ bootstrap samples).

| With BY 2014 |  | $\beta$ |  |
| :---: | :--- | :--- | :--- |
| Parameter | $\alpha$ | $\beta$ | $\beta_{2}$ |
| $\alpha$ |  | $<0.0001$ | $<0.0001$ |
| $\beta$ | 0.9107 | 0.0175 | 0.3015 |
| $\beta_{2}$ | 0.1240 |  |  |
| Without BY 2014 |  |  |  |
| Parameter $\alpha$ | $\alpha$ | $<0.0001$ | $\beta_{2}$ |
| $\alpha$ |  |  | $<0.0001$ |
| $\beta$ | 0.9669 | 0.1259 | $<0.0001$ |
| $\beta_{2}$ | 0.2922 |  |  |

Table C2. Pearson correlation coefficient (below diagonal) and two-sided P-value (above diagonal) for Ricker stock-recruitment model of smolts per redd for White River Spring Chinook, 2005-2017, with and without brood year 2014. Model form is: $\ln (R / S)=\beta_{0}-\beta S-\beta_{2} P_{H O S}$. Correlation computed from bootstrap ( $\geq 3,000$ bootstrap samples).

| With BY 2014 |  | $\beta$ |  |
| :---: | :--- | :--- | :--- |
| Parameter | $\beta_{0}$ | $\beta$ | $\beta_{2}$ |
| $\beta_{0}$ | 0.6155 | $<0.0001$ | $<0.0001$ |
| $\beta$ | 0.7044 | -0.0652 | 0.0003 |
| $\beta_{2}$ |  |  |  |
|  |  | $\beta$ |  |
| Without BY 2014 | $\beta_{0}$ | $<0.0001$ |  |
| Parameter |  |  | $\beta_{2}$ |
| $\beta_{0}$ | 0.5568 | -0.0755 | $<0.0001$ |
| $\beta$ | 0.7439 |  | $<0.0001$ |
| $\beta_{2}$ |  |  |  |

White Spring Chinook: Beverton-Holt Model - Juveniles Per Redd


Figure C16. Residual plots for Beverton-Holt model of smolts per redd as a function of redd count and pHOS for White River Spring Chinook, 2005-2017, with 2014. Shapiro-Wilk test of normality of residuals: $\mathrm{W}=0.8585, \mathrm{P}=0.3681$.


Figure C17. Normal quantile-quantile plot of residual from Beverton-Holt model of smolts per redd as a function of redd count and pHOS for White River Spring Chinook, 2005-2017, with 2014. Points should lie on the line under assumption of normal errors. Shapiro-Wilk test of normality of residuals: $\mathrm{W}=0.8585$, $\mathrm{P}=0.3681$.


Figure C18. Autocorrelation plot of residuals for Beverton-Holt model of smolts per redd as a function of redd count and pHOS for White River Spring Chinook, 2005-2017, with 2014. Blue lines mark 95\% confidence interval under assumption of no autocorrelation.


Figure C19. Residual plots for Beverton-Holt model of smolts per redd as a function of redd count and pHOS for White River Spring Chinook, 2005-2017, without 2014. Shapiro-Wilk test of normality of residuals: $\mathrm{W}=0.9549, \mathrm{P}=0.7099$.


Figure C20. Normal quantile-quantile plot of residual from Beverton-Holt model of smolts per redd as a function of redd count and pHOS for White River Spring Chinook, 2005-2017, without 2014. Points should lie on the line under assumption of normal errors. Shapiro-Wilk test of normality of residuals: $\mathrm{W}=0.9549$, $\mathrm{P}=0.7099$.


Figure C21. Autocorrelation plot of residuals for Beverton-Holt model of smolts per redd as a function of redd count and pHOS for White River Spring Chinook, 2005-2017, without 2014. Blue lines mark 95\% confidence interval under assumption of no autocorrelation.


Figure C22. Residual plots for Ricker model of smolts per redd as a function of redd count and pHOS for White River Spring Chinook, 2005-2017, including brood year 2014. Shapiro-Wilk test of normality of residuals: $\mathrm{W}=0.8682, \mathrm{P}=0.0495$.


Figure C23. Autocorrelation plot of residuals for Ricker model of smolts per redd as a function of redd count and pHOS for White River Spring Chinook, 2005-2017, including brood year 2014. Blue lines mark $95 \%$ confidence interval under assumption of no autocorrelation.


Figure C24. Residual plots for Ricker model of smolts per redd as a function of redd count and pHOS for White River Spring Chinook, 2005-2017, omitting brood year 2014. Shapiro-Wilk test of normality of residuals: $\mathrm{W}=0.9540, \mathrm{P}=0.6956$.


Figure C25. Autocorrelation plot of residuals for Ricker model of smolts per redd as a function of redd count and pHOS for White River Spring Chinook, 2005-2017, omitting brood year 2014. Blue lines mark $95 \%$ confidence interval under assumption of no autocorrelation.

## Appendix D: Twisp River Spring Chinook Salmon



Figure D1. Normal quantile-quantile plot of residual from the Smooth Hockey Stick model of smolt recruitment as a function of spawner abundance (stock) for Twisp River Spring Chinook, 2003-2017. Points should lie on the line under assumption of normal errors. Shapiro-Wilk test of normality of residuals: $\mathrm{W}=0.9707, \mathrm{P}=0.8678$.


Figure D2. Autocorrelation plot of residuals for the Smooth Hockey Stick model of smolt recruitment as a function of spawner abundance (stock) for Twisp River Spring Chinook, 2003-2017. Blue lines mark 95\% confidence interval under assumption of no autocorrelation.


Figure D3. Correlation plot of bootstrap parameter estimates from the Smooth Hockey Stick model of smolt recruitment as a function of spawner abundance (stock) for Twisp River Spring Chinook, 2003-2017, based on 2,813 bootstrap samples.


Figure D4. Normal quantile-quantile plot of residual from Beverton-Holt model of smolt recruitment as a function of spawner abundance (stock) for Twisp River Spring Chinook, 2003-2017. Points should lie on the line under assumption of normal errors. Shapiro-Wilk test of normality of residuals: $\mathrm{W}=0.9700$, $\mathrm{P}=0.8579$.

Twisp Spring Chinook: Beverton-Holt Model


Figure D5. Autocorrelation plot of residuals for Beverton-Holt model of smolt recruitment as a function of spawner abundance (stock) for Twisp River Spring Chinook, 2003-2017. Blue lines mark 95\% confidence interval under assumption of no autocorrelation.


Figure D6. Normal quantile-quantile plot of residual from Ricker model of smolt recruitment as a function of spawner abundance (stock) for Twisp River Spring Chinook, 2003-2017. Points should lie on the line under assumption of normal errors. Shapiro-Wilk test of normality of residuals: $\mathrm{W}=0.9815, \mathrm{P}=0.8831$.

## Twisp Spring Chinook: Ricker Model



Figure D7. Autocorrelation plot of residuals for Ricker model of smolt recruitment as a function of spawner abundance (stock) for Twisp River Spring Chinook, 2003-2017. Blue lines mark 95\% confidence interval under assumption of no autocorrelation.


Figure D8. Residual plots for linear model of smolts per redd (log scale) as a function of pHOS for Twisp River Spring Chinook, 2003-2017. Shapiro-Wilk test of normality of residuals: $\mathrm{W}=0.9575, \mathrm{P}=0.6489$.

Table D1. Pearson correlation coefficient (below diagonal) and two-sided P-value (above diagonal) for Beverton-Holt stock-recruitment model of smolts per redd for Twisp River Spring Chinook, 2003-2017. Model form is: $\ln (R / S)=\ln (\alpha)-\beta_{2} P_{\text {HoS }}-\ln (\beta+S)$. Correlation computed from bootstrap (2,798 bootstrap samples).

| Parameter | $\alpha$ | $\beta$ | $\beta_{2}$ |
| :---: | :--- | :--- | :--- |
| $\alpha$ |  | $<0.0001$ | $<0.0001$ |
| $\beta$ | 0.9153 |  | 0.3039 |
| $\beta_{2}$ | 0.1298 | -0.0194 |  |

Table D2. Pearson correlation coefficient (below diagonal) and two-sided P-value (above diagonal) for Ricker stock-recruitment model of smolts per redd for Twisp River Spring Chinook, 2003-2017. Model form is: $\ln (R / S)=\beta_{0}-\beta S-\beta_{2} P_{\text {Hоs }}$. Correlation computed from bootstrap (3,000 bootstrap samples).

| Parameter | $\beta_{0}$ | $\beta$ | $\beta_{2}$ |
| :---: | :--- | :--- | :--- |
| $\beta_{0}$ | 0.5750 | $<0.0001$ | $<0.0001$ |
| $\beta$ | 0.7243 | -0.0372 | 0.0418 |
| $\beta_{2}$ |  |  |  |

Twisp Spring Chinook: Beverton-Holt Model - Juveniles Per Redd


Figure D9. Residual plots for Beverton-Holt model of smolts per redd as a function of redd count and pHOS for Twisp River Spring Chinook, 2003-2017. Shapiro-Wilk test of normality of residuals: W=0.9345, $\mathrm{P}=0.3185$.


Figure D10. Normal quantile-quantile plot of residual from Beverton-Holt model of smolts per redd as a function of redd count and pHOS for Twisp River Spring Chinook, 2003-2017. Points should lie on the line under assumption of normal errors. Shapiro-Wilk test of normality of residuals: $\mathrm{W}=0.9345, \mathrm{P}=0.3185$.

Twisp Spring Chinook: Beverton-Holt Model - Juveniles Per Redd


Figure D11. Autocorrelation plot of residuals for Beverton-Holt model of smolts per redd as a function of redd count and pHOS for Twisp River Spring Chinook, 2003-2017. Blue lines mark $95 \%$ confidence interval under assumption of no autocorrelation.


Figure D12. Residual plots for Ricker model of smolts per redd as a function of redd count and pHOS for Twisp River Spring Chinook, 2003-2017. Shapiro-Wilk test of normality of residuals: W=0.9284, $\mathrm{P}=0.2587$.

Twisp Spring Chinook: Ricker Model - Juveniles Per Redd


Figure D13. Autocorrelation plot of residuals for Ricker model of smolts per redd as a function of redd count and pHOS for Twisp River Spring Chinook, 2003-2017. Blue lines mark $95 \%$ confidence interval under assumption of no autocorrelation.

## Appendix E: Methow River Spring Chinook Salmon



Figure E1. Normal quantile-quantile plot of residual from the Smooth Hockey Stick model of smolt recruitment as a function of spawner abundance (stock) for Methow River Spring Chinook, 2002-2017. Points should lie on the line under assumption of normal errors. Shapiro-Wilk test of normality of residuals: $\mathrm{W}=0.9376, \mathrm{P}=0.3202$.

Methow Spring Chinook: Smooth Hockey Stick Model


Figure E2. Autocorrelation plot of residuals for the Smooth Hockey Stick model of smolt recruitment as a function of spawner abundance (stock) for Methow River Spring Chinook, 2002-2017. Blue lines mark $95 \%$ confidence interval under assumption of no autocorrelation.


Figure E3. Correlation plot of bootstrap parameter estimates from the Smooth Hockey Stick model of smolt recruitment as a function of spawner abundance (stock) for Methow River Spring Chinook, 2002-2017, based on 3,182 bootstrap samples.


Figure E4. Normal quantile-quantile plot of residual from Beverton-Holt model of smolt recruitment as a function of spawner abundance (stock) for Methow River Spring Chinook, 2002-2017. Points should lie on the line under assumption of normal errors. Shapiro-Wilk test of normality of residuals: $\mathrm{W}=0.9382$, $\mathrm{P}=0.3275$.

Methow Spring Chinook: Beverton-Holt Model


Figure E5. Autocorrelation plot of residuals for Beverton-Holt model of smolt recruitment as a function of spawner abundance (stock) for Methow River Spring Chinook, 2002-2017. Blue lines mark 95\% confidence interval under assumption of no autocorrelation.


Figure E6. Normal quantile-quantile plot of residual from Ricker model of smolt recruitment as a function of spawner abundance (stock) for Methow River Spring Chinook, 2002-2017. Points should lie on the line under assumption of normal errors. Shapiro-Wilk test of normality of residuals: $\mathrm{W}=0.9370, \mathrm{P}=0.3135$.

Methow Spring Chinook: Ricker Model


Figure E7. Autocorrelation plot of residuals for Ricker model of smolt recruitment as a function of spawner abundance (stock) for Methow River Spring Chinook, 2002-2017. Blue lines mark 95\% confidence interval under assumption of no autocorrelation.


Figure E8. Residual plots for linear model of smolts per redd as a function of pHOS for Methow River Spring Chinook, 2002-2017. Shapiro-Wilk test of normality of residuals: $\mathrm{W}=0.9467, \mathrm{P}=0.4400$.

Table E1. Pearson correlation coefficient (below diagonal) and two-sided P-value (above diagonal) for Ricker stock-recruitment model of smolts per redd for Methow River Spring Chinook, 2002-2017. Model form is: $\ln (R / S)=\beta_{0}-\beta S-\beta_{2} P_{\text {Hos }}$. Correlation computed from bootstrap (3,000 bootstrap samples).

| Parameter | $\beta_{0}$ | $\beta$ | $\beta_{2}$ |
| :---: | :--- | :--- | :--- |
| $\beta_{0}$ | -0.2102 | $<0.0001$ | $<0.0001$ |
| $\beta$ | 0.9366 | -0.5241 | $<0.0001$ |
| $\beta_{2}$ |  |  |  |



Figure E9. Residual plots for Ricker model of smolts per redd as a function of redd count and pHOS for Methow River Spring Chinook, 2002-2017. Shapiro-Wilk test of normality of residuals: W=0.9794, $\mathrm{P}=0.9589$.


Figure E10. Autocorrelation plot of residuals for Ricker model of smolts per redd as a function of redd count and pHOS for Methow River Spring Chinook, 2002-2017. Blue lines mark 95\% confidence interval under assumption of no autocorrelation.

## Appendix F: Wenatchee River Summer Chinook Salmon



Figure F1. Normal quantile-quantile plot of residual from Ricker model of emigrant count as a function of spawner abundance (stock) for Wenatchee River Summer Chinook, 1999-2018 (without 2010, 2011). Points should lie on the line under assumption of normal errors. Shapiro-Wilk test of normality of residuals: $\mathrm{W}=0.9015, \mathrm{P}=0.0611$.

Wenatchee Summer Chinook: Ricker Model


Figure F2. Autocorrelation plot of residuals for Ricker model of smolt recruitment as a function of spawner abundance (stock) for Wenatchee River Summer Chinook, 1999-2018 (without 2010, 2011). Blue lines mark $95 \%$ confidence interval under assumption of no autocorrelation.

Wenatchee Summer Chinook Salmon


Figure F3. The Smooth Hockey Stick, Beverton-Holt, and Ricker stock-recruitment model fit to emigrants and spawner data for Wenatchee River Summer Chinook Salmon, 1999-2018 (without 2000, 2010, 2011).


Figure F4. Residuals from the Smooth Hockey Stick Model versus proportion of hatchery origin spawners ( pHOS ) with fitted linear regression line, Pearson correlation coefficient ( r ), and P-value from two-sided ttest of slope of linear regression line using smolt and spawner data from Wenatchee River Summer Chinook Salmon, 1999-2018 (without 2000, 2010, 2011). $\mathrm{R}^{2}=0.0141$.


Figure F5. Residuals from the Beverton-Holt Model versus proportion of hatchery origin spawners (pHOS) with fitted linear regression line, Pearson correlation coefficient ( r ), and P-value from two-sided t-test of slope of linear regression line using smolt and spawner data from Wenatchee River Summer Chinook Salmon, 1999-2018 (without 2000, 2010, 2011). $\mathrm{R}^{2}=0.0134$.


Figure F6. Residuals from Ricker Model versus proportion of hatchery origin spawners (pHOS) with fitted linear regression line, Pearson correlation coefficient ( r ), and P -value from two-sided t -test of slope of linear regression line using smolt and spawner data from Wenatchee River Summer Chinook Salmon, 1999-2018 (without 2000, 2010, 2011). $\mathrm{R}^{2}=0.0147$.


Figure F7. Residual plots for linear model of emigrants (log scale) per redd as a function of pHOS for Wenatchee River Summer Chinook, 1999-2018 (without 2010, 2011). Shapiro-Wilk test of normality of residuals: $\mathrm{W}=0.8877, \mathrm{P}=0.0353$.

Table F1. Pearson correlation coefficient (below diagonal) for Beverton-Holt stock-recruitment model of smolts per redd for Wenatchee River Summer Chinook, 1999-2018 (without 2010, 2011). Attempts to bootstrap for variance failed, and correlation significance could not be estimated. Model form is: $\ln (R / S)=\ln (\alpha)-\beta_{2} P_{\text {HOS }}-\ln (\beta+S)$. Correlation computed from Hessian matrix from nonlinear model fitting.

| Parameter | $\alpha$ | $\beta$ | $\beta_{2}$ |
| :---: | :---: | :---: | :---: |
| $\alpha$ |  |  |  |
| $\beta$ | 0.9887 |  |  |
| $\beta_{2}$ | -0.0548 | -0.1913 |  |

Table F2. Pearson correlation coefficient (below diagonal) and two-sided P-value (above diagonal) for Ricker stock-recruitment model of smolts per redd for Methow River Summer Chinook, 1999-2018 (without 2010, 2011). Model form is: $\ln (R / S)=\beta_{0}+\beta_{1} S-\beta_{2} P_{\text {HOS }}$. Correlation computed from bootstrap ( $>3000$ bootstrap samples).

| Parameter | $\beta_{0}$ | $\beta_{1}$ | $\beta_{2}$ |
| :---: | :---: | :---: | :---: |
| $\beta_{0}$ |  | $<0.0001$ | $<0.0001$ |
| $\beta_{1}$ | -0.8055 |  | $<0.0001$ |
| $\beta_{2}$ | 0.4975 | 0.3896 |  |



Figure F8. Residual plots for Beverton-Holt model of emigrants per redd as a function of redd count and pHOS for Wenatchee River Summer Chinook, 1999-2018 (without 2010, 2011). Shapiro-Wilk test of normality of residuals: $\mathrm{W}=0.8845, \mathrm{P}=0.0311$.


Figure F9. Normal quantile-quantile plot of residual from Beverton-Holt model of emigrants per redd as a function of redd count and pHOS for Wenatchee River Summer Chinook, 1999-2018 (without 2010, 2011). Points should lie on the line under assumption of normal errors. Shapiro-Wilk test of normality of residuals: $\mathrm{W}=0.8845, \mathrm{P}=0.0311$.

Wenatchee Summer Chinook: Beverton-Holt Model - Juveniles Per Redd


Figure F10. Autocorrelation plot of residuals for Beverton-Holt model of emigrants per redd as a function of redd count and pHOS for Wenatchee River Summer Chinook, 1999-2018 (without 2010, 2011). Blue lines mark $95 \%$ confidence interval under assumption of no autocorrelation.


Figure F11. Residual plots for Ricker model of emigrants per redd as a function of redd count and pHOS for Wenatchee River Summer Chinook, 1999-2018 (without 2010, 2011). Shapiro-Wilk test of normality of residuals: $\mathrm{W}=0.8822, \mathrm{P}=0.0284$.


Figure F12. Autocorrelation plot of residuals for Ricker model of smolts per redd as a function of redd count and pHOS for Wenatchee River Summer Chinook, 1999-2018 (without 2010, 2011). Blue lines mark $95 \%$ confidence interval under assumption of no autocorrelation.


Figure F13. Emigrants per redd (log scale) versus proportion of hatchery origin spawners ( pHOS ) for Wenatchee River Summer Chinook Salmon, 1999-2018 (without 2000, 2010, 2011), with fitted linear regression line, Pearson correlation coefficient (r), and P-value from one-sided t-test of negative slope. $\mathrm{R}^{2}$ $=0.0030$.

## Appendix G: Methow River Summer Chinook Salmon



Figure G1. Normal quantile-quantile plot of residual from Smooth Hockey Stick model of smolt recruitment as a function of spawner abundance (stock) for Methow River Summer Chinook, 2006-2018 (without 2012). Points should lie on the line under assumption of normal errors. Shapiro-Wilk test of normality of residuals: $\mathrm{W}=0.9192, \mathrm{P}=0.2793$.

## Methow Summer Chinook: Smooth Hockey Stick Model



Figure G2. Autocorrelation plot of residuals for Smooth Hockey Stick model of smolt recruitment as a function of spawner abundance (stock) for Methow River Summer Chinook, 2006-2018 (without 2012). Blue lines mark $95 \%$ confidence interval under assumption of no autocorrelation.


Figure G3. Normal quantile-quantile plot of residual from Beverton-Holt model of emigrant count as a function of spawner abundance (stock) for Methow River Summer Chinook, 2006-2018 (without 2012). Points should lie on the line under assumption of normal errors. Shapiro-Wilk test of normality of residuals: $\mathrm{W}=0.9206, \mathrm{P}=0.2910$.


Figure G4. Autocorrelation plot of residuals for Beverton-Holt model of smolt recruitment as a function of spawner abundance (stock) for Methow River Summer Chinook, 2006-2018 (without 2012). Blue lines mark $95 \%$ confidence interval under assumption of no autocorrelation.


Figure G5. Normal quantile-quantile plot of residual from Ricker model of emigrant count as a function of spawner abundance (stock) for Methow River Summer Chinook, 2006-2018 (without 2012). Points should lie on the line under assumption of normal errors. Shapiro-Wilk test of normality of residuals: $\mathrm{W}=0.9181$, $\mathrm{P}=0.2707$.

Methow Summer Chinook: Ricker Model


Figure G6. Autocorrelation plot of residuals for Ricker model of smolt recruitment as a function of spawner abundance (stock) for Methow River Summer Chinook, 2006-2018 (without 2012). Blue lines mark 95\% confidence interval under assumption of no autocorrelation.


Figure G7. Residual plots for linear model of smolts per redd as a function of pHOS for Methow River Summer Chinook, 2006-2018 (without 2012). Shapiro-Wilk test of normality of residuals: W=0.9284, $\mathrm{P}=0.3632$.

Table G1. Pearson correlation coefficient (below diagonal) and two-sided P-value (above diagonal) for Beverton-Holt stock-recruitment model of smolts per redd for Methow River Summer Chinook, 2006-2018 (without 2012). Model form is: $\ln (R / S)=\ln (\alpha)-\beta_{2} P_{H O S}-\ln (\beta+S)$. Correlation computed from Hessian matrix from nonlinear model fitting.

| Parameter | $\alpha$ | $\beta$ | $\beta_{2}$ |
| :---: | :---: | :---: | :---: |
| $\alpha$ |  | NA | NA |
| $\beta$ | 0.9993 |  | NA |
| $\beta_{2}$ | 0.1676 | 0.1333 |  |

Table G2. Pearson correlation coefficient (below diagonal) and two-sided P-value (above diagonal) for Ricker stock-recruitment model of smolts per redd for Methow River Summer Chinook, 2006-2018 (without 2012). Model form is: $\ln (R / S)=\beta_{0}+\beta_{1} S-\beta_{2} P_{H O S}$. Correlation computed from bootstrap ( $>$ 1500 bootstrap samples).

| Parameter | $\beta_{0}$ | $\beta_{1}$ | $\beta_{2}$ |
| :---: | :--- | :--- | :--- |
| $\beta_{0}$ | -0.7893 | $<0.0001$ | $<0.0001$ |
| $\beta_{1}$ | 0.8198 |  | $<0.0001$ |
| $\beta_{2}$ | -0.3334 |  |  |



Figure G8. Residual plots for Beverton-Holt model of emigrants per redd as a function of redd count and pHOS for Methow River Summer Chinook, 2006-2018 (without 2012). Shapiro-Wilk test of normality of residuals: $\mathrm{W}=0.9255, \mathrm{P}=0.3352$.


Figure G9. Normal quantile-quantile plot of residual from Beverton-Holt model of emigrants per redd as a function of redd count and pHOS for Methow River Summer Chinook, 2006-2018 (without 2012). Points should lie on the line under assumption of normal errors. Shapiro-Wilk test of normality of residuals: $\mathrm{W}=0.9255, \mathrm{P}=0.3352$.

Methow Summer Chinook: Beverton-Holt Model - Juveniles Per Redd


Figure G10. Autocorrelation plot of residuals for Beverton-Holt model of emigrants per redd as a function of redd count and pHOS for Methow River Summer Chinook, 2006-2018 (without 2012). Blue lines mark $95 \%$ confidence interval under assumption of no autocorrelation.


Figure G11. Residual plots for Ricker model of emigrants per redd as a function of redd count and pHOS for Methow River Summer Chinook, 2006-2018 (without 2012). Shapiro-Wilk test of normality of residuals: $\mathrm{W}=0.9251, \mathrm{P}=0.3313$.


Figure G12. Autocorrelation plot of residuals for Ricker model of smolts per redd as a function of redd count and pHOS for Methow River Summer Chinook, 2006-2018 (without 2012). Blue lines mark 95\% confidence interval under assumption of no autocorrelation.

## Appendix H: Methow River Summer Steelhead



Figure H1. Normal quantile-quantile plot of residual from Ricker model of emigrant recruitment as a function of spawner abundance (stock) for Methow River Summer Steelhead, 2003-2015. Points should lie on the line under assumption of normal errors. Shapiro-Wilk test of normality of residuals: $\mathrm{W}=0.9445$, $\mathrm{P}=0.5183$.


Figure H2. Autocorrelation plot of residuals for Ricker model of emigrant recruitment as a function of spawner abundance (stock) for Methow River Summer Steelhead, 2003-2015. Blue lines mark 95\% confidence interval under assumption of no autocorrelation.


Figure H3. Residual plots for linear model of emigrants per redd (log scale) as a function of pHOS for Methow River Summer Steelhead, 2003-2015. Shapiro-Wilk test of normality of residuals: W=0.9579, $\mathrm{P}=0.7205$.

Table H1. Pearson correlation coefficient (below diagonal) and two-sided P-value (above diagonal) for Ricker stock-recruitment model of emigrants per redd for Methow River Summer Steelhead, 2003-2015. Model form is: $\ln (R / S)=\beta_{0}-\beta S-\beta_{2} P_{\text {Hos }}$. Correlation computed from bootstrap (3,000 bootstrap samples).

| Parameter | $\beta_{0}$ | $\beta$ | $\beta_{2}$ |
| :---: | :--- | :--- | :--- |
| $\beta_{0}$ |  | 0.0003 | $<0.0001$ |
| $\beta$ | -0.0656 |  | $<0.0001$ |
| $\beta_{2}$ | 0.9637 | -0.3156 |  |



Figure H4. Residual plots for Ricker model of emigrants per redd as a function of redd count and pHOS for Methow River Summer Steelhead, 2003-2015. Shapiro-Wilk test of normality of residuals: W=0.9671, $\mathrm{P}=0.8578$.


Figure H5. Autocorrelation plot of residuals for Ricker model of emigrants per redd as a function of redd count and pHOS for Methow River Summer Steelhead, 2003-2015. Blue lines mark $95 \%$ confidence interval under assumption of no autocorrelation.

## Appendix I: Twisp River Summer Steelhead



Figure I1. Normal quantile-quantile plot of residual from Ricker model of emigrant recruitment as a function of spawner abundance (stock) for Twisp River Summer Steelhead, 2003-2015, including brood year 2007. Points should lie on the line under assumption of normal errors. Shapiro-Wilk test of normality of residuals: $\mathrm{W}=0.9011, \mathrm{P}=0.1383$.

Twisp Summer Steelhead: Ricker Model


Figure I2. Autocorrelation plot of residuals for Ricker model of emigrant recruitment as a function of spawner abundance (stock) for Twisp River Summer Steelhead, 2003-2015, including brood year 2007. Blue lines mark $95 \%$ confidence interval under assumption of no autocorrelation.


Figure I3. Normal quantile-quantile plot of residual from Ricker model of emigrant recruitment as a function of spawner abundance (stock) for Twisp River Summer Steelhead, 2003-2015, without brood year 2007. Points should lie on the line under assumption of normal errors. Shapiro-Wilk test of normality of residuals: $\mathrm{W}=0.8429, \mathrm{P}=0.0300$.

## Twisp Summer Steelhead: Ricker Model



Figure I4. Autocorrelation plot of residuals for Ricker model of emigrant recruitment as a function of spawner abundance (stock) for Twisp River Summer Steelhead, 2003-2015, without brood year 2007. Blue lines mark $95 \%$ confidence interval under assumption of no autocorrelation.


Figure I5. Residual plots for linear model of emigrants per redd (log scale) as a function of pHOS for Twisp River Summer Steelhead, 2003-2015, including brood year 2007. Shapiro-Wilk test of normality of residuals: $\mathrm{W}=0.8461, \mathrm{P}=0.0254$.


Figure I6. Negative log-likelihood surface and contour curves for Beverton-Holt model of emigrants, redd counts, and pHOS , using $\beta_{2}=-0.1160$. Lighter regions indicate higher likelihood values. The green triangle (falls below $\mathrm{b}=0$ line) indicates the parameter set that optimizes the likelihood. Data set $=$ Twisp River Summer Steelhead, 2003-2015, including brood year 2007.

Table I1. Pearson correlation coefficient (below diagonal) and two-sided P-value (above diagonal) for Ricker stock-recruitment model of emigrants per redd for Twisp River Summer Steelhead, 2003-2015, including 2007. Model form is: $\ln (R / S)=\beta_{0}-\beta S-\beta_{2} P_{\text {Hos }}$. Correlation computed from bootstrap ( 3,000 bootstrap samples).

| Parameter | $\beta_{0}$ | $\beta$ | $\beta_{2}$ |
| :---: | :--- | :--- | :--- |
| $\beta_{0}$ | -0.6298 | $<0.0001$ | $<0.0001$ |
| $\beta$ | 0.9280 |  | $<0.0001$ |
| $\beta_{2}$ | -0.8606 |  |  |



Figure I7. Residual plots for Ricker model of emigrants per redd as a function of redd count and pHOS for Twisp River Summer Steelhead, 2003-2015, including 2007. Shapiro-Wilk test of normality of residuals: $\mathrm{W}=0.8740, \mathrm{P}=0.0592$.

Twisp Summer Steelhead: Ricker Model - Juveniles Per Redd


Lag
Figure I8. Autocorrelation plot of residuals for Ricker model of emigrants per redd as a function of redd count and pHOS for Twisp River Summer Steelhead, 2003-2015, including 2007. Blue lines mark 95\% confidence interval under assumption of no autocorrelation.


Figure 19. Residual plots for linear model of emigrants per redd as a function of pHOS for Twisp River Summer Steelhead, 2003-2015, omitting brood year 2007. Shapiro-Wilk test of normality of residuals: $\mathrm{W}=0.9230, \mathrm{P}=0.3116$.

Table I2. Pearson correlation coefficient (below diagonal) and two-sided P-value (above diagonal) for Beverton-Holt stock-recruitment model of smolts per redd for Twisp River Summer Steelhead, 2003-2015, omitting brood year 2007. Model form is: $\ln (R / S)=\ln (\alpha)-\beta_{2} P_{H O S}-\ln (\beta+S)$. Correlation computed from bootstrap ( 2,909 bootstrap samples).

| Parameter | $\alpha$ | $\beta$ | $\beta_{2}$ |
| :---: | :--- | :--- | :--- |
| $\alpha$ |  | $<0.0001$ | $<0.0001$ |
| $\beta$ | 0.8976 |  | $<0.0001$ |
| $\beta_{2}$ | 0.8437 | 0.6772 |  |

Table I3. Pearson correlation coefficient (below diagonal) and two-sided P-value (above diagonal) for Ricker stock-recruitment model of emigrants per redd for Twisp River Summer Steelhead, 2003-2015, omitting brood year 2007. Model form is: $\ln (R / S)=\beta_{0}-\beta S-\beta_{2} P_{\text {HOS }}$. Correlation computed from bootstrap ( 3,000 bootstrap samples).

| Parameter | $\beta_{0}$ | $\beta$ | $\beta_{2}$ |
| :---: | :--- | :--- | :--- |
| $\beta_{0}$ |  | $<0.0001$ | $<0.0001$ |
| $\beta$ | -0.6094 |  | $<0.0001$ |
| $\beta_{2}$ | 0.9213 | -0.8469 |  |



Figure I10. Residual plots for Beverton-Holt model of smolts per redd as a function of redd count and pHOS for Twisp River Summer Steelhead, 2003-2015, omitting brood year 2007. Shapiro-Wilk test of normality of residuals: $\mathrm{W}=0.9728, \mathrm{P}=0.9377$.


Figure I11. Normal quantile-quantile plot of residual from Beverton-Holt model of smolts per redd as a function of redd count and pHOS for Twisp River Summer Steelhead, 2003-2015, omitting brood year 2007. Points should lie on the line under assumption of normal errors. Shapiro-Wilk test of normality of residuals: $\mathrm{W}=0.9728, \mathrm{P}=0.9377$.

Twisp Summer Steelhead: Beverton-Holt Model - Juveniles Per Redd


Figure I12. Autocorrelation plot of residuals for Beverton-Holt model of smolts per redd as a function of redd count and pHOS for Twisp River Summer Steelhead, 2003-2015, omitting brood year 2007. Blue lines mark $95 \%$ confidence interval under assumption of no autocorrelation.


Figure I13. Residual plots for Ricker model of emigrants per redd as a function of redd count and pHOS for Twisp River Summer Steelhead, 2003-2015, omitting brood year 2007. Shapiro-Wilk test of normality of residuals: $\mathrm{W}=0.9588, \mathrm{P}=0.7658$.

Twisp Summer Steelhead: Ricker Model - Juveniles Per Redd


Figure I14. Autocorrelation plot of residuals for Ricker model of emigrants per redd as a function of redd count and pHOS for Twisp River Summer Steelhead, 2003-2015, omitting brood year 2007. Blue lines mark $95 \%$ confidence interval under assumption of no autocorrelation.

## COLUMBIA BASIN RESEARCH

## Appendix B

Power Analysis for Effect of pHOS on Juvenile Productivity

## COLUMBIA BASIN RESEARCH

## Power Analysis for effect of pHOS on juvenile productivity

## 5 August 2021

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## Introduction

There is concern that a higher proportion of hatchery-origin spawners ( pHOS ) on the spawning grounds may have a negative effect on juvenile productivity. Analysis of existing data has shown little to no evidence of a negative effect for Chinook Salmon and steelhead in the Upper Columbia tributaries but is limited by short time series and high observation error. A power analysis is warranted to determine the ability to detect a negative effect of pHOS , should it exist, and how many years of data collection are expected to achieve a useful level of power.

## Statistical Methods

The statistical power was computed to detect a reduction in mean juvenile recruitment with Type I error probability of $\alpha=0.10$ when the actual reduction in mean recruitment associated with a change in the pHOS level was $5 \%$. Power was computed at varying sample sizes, where sample size $=$ number of years of observations, and both with and without measurement error in the number of spawners and pHOS . The stock-recruitment model was a Ricker model with parameters taken from the Chiwawa River spring Chinook Salmon population. The Chiwawa population was selected because it has the longest time series of stock and juvenile recruitment data of the populations available and the stock recruitment curves have fit the juvenile productivity from this population relatively well (Buchanan and Townsend 2021). Power was calculated for sampling sizes from $N=5$ to $N=100$ years for varying levels of pHOS effect size, $\beta_{P}$.

## Ricker Stock-Recruitment Model

The effect of pHOS is assessed after accounting for density dependence by either regressing the residuals from a stock-recruitment curve onto pHOS or else incorporating pHOS as a term in a stock-recruitment model. For the purposes of this power analysis, we used the latter approach and incorporated pHOS into the Ricker model as follows:

$$
\begin{equation*}
R=a e^{\beta_{P} P_{H o s}} \times S e^{-b S} \tag{1}
\end{equation*}
$$

where $R=$ recruits, $S=$ spawner abundance, $a=$ increase in recruits per spawner at low levels of spawners, and $b=$ intensity of the decrease in recruitment at high levels of spawners, and lognormal errors are assumed. The parameter $\beta_{P}$ is the fixed effect of pHOS on recruitment. The maximum number of recruits is defined as $K=(a / b) e^{-1} e^{\beta_{P} P_{\text {ноя }}}$. This formulation of the model assumes that pHOS may lower the slope of the spawner-recruitment curve at low levels of spawner abundance if $\beta_{P}<0$, which will also result in a lower maximum recruitment. Model (1) may be rewritten as a linear model as follows:

$$
\begin{equation*}
Y=\ln (a)-b S+\beta_{P} P_{H O S}+\varepsilon \tag{2}
\end{equation*}
$$

where $Y=\ln (R / S)$ and $\varepsilon \sim N\left(0, \sigma^{2}\right)$. A negative effect of pHOS is consistent with $\beta_{P}<0$ and may be concluded when the upper limit of the $(1-\alpha) \times 100 \%$ bootstrap confidence interval is $<0$ (i.e., the confidence interval is entirely below 0 ) for $\alpha=$ probability of a Type I error. Power was computed using simulations. Briefly, the number of spawners ( $S$ ), pHOS, and smolt recruits were simulated from model (2) according to the existing data for the Chiwawa River spring Chinook Salmon population and for a specific value of the pHOS effect $\left(\beta_{P}\right)$. Measurement error was added to the spawners and pHOS values, and the Ricker model in model (2) was fit to the resulting simulated data set. Statistical power was computed by the proportion of the simulated data sets that yielded a $90 \%$ bootstrap confidence interval that was completely $<0$. One thousand $(1,000)$ simulations were used. Details and results are provided below.

## Simulations

For each simulation $i=1, \ldots, 1000$, the total number of spawners $S_{i y}$ and the number of natural origin spawners $\operatorname{NOS}_{i y}$ were simulated for each year $y(y=1, \ldots, N)$ from independent negative binomial distributions whose parameters were estimated from observed Chiwawa spring Chinook Salmon data from 1991-2017 (Table 1 and Figure 1):

$$
\begin{gathered}
S_{i y} \sim \operatorname{NBin}(n=1.30, \mu=722.81) \\
\operatorname{NOS}_{i y} \sim \operatorname{NBin}(n=1.36, \mu=287.57)
\end{gathered}
$$

where $\mu$ is the mean and $n$ is the dispersion parameter. Any simulated NOS value $>S$ was replaced by a new simulation to ensure $N O S_{i y} \leq S_{i y}$ for each year $y$ and simulation $i$. For each simulation $i$ and year $y$, the simulated pHOS value was calculated as $p \operatorname{HOS}_{i y}=\operatorname{HOS}_{i y} / S_{i y}$, where $\operatorname{HOS}_{i y}=\left(1-N O S_{i y}\right) S_{i y}$ is the simulated number of hatchery origin spawners.

For a given value of the pHOS effect $\beta_{P}$, the number of juvenile recruits was calculated according to

$$
\begin{equation*}
R_{i y}=S_{i y} \times \exp \left(\ln (a)-b S_{i y}+\beta_{P} P_{H O S, i y}+\varepsilon_{i y}\right) \tag{3}
\end{equation*}
$$

where $\varepsilon_{i y} \sim N\left(0, \sigma^{2}\right)$. The parameters $a, b$, and $\sigma^{2}$ were estimated from the Chiwawa spring Chinook salmon population data from 1991-2017: $a=138, b=0.0011$, and $\sigma^{2}=0.1924$.

Measurement error was incorporated to the data set by simulating the size of measurement error for each component and adding it to the simulated "true" value. The data collected each year consisted of the number of natural origin spawners (NOS) and the number of hatchery origin spawners (HOS); the data reported are $S=N O S+H O S$ and $p H O S=H O S / S$. Thus, measurement error was simulated for NOS and HOS using reported precision levels of coefficient of variation (CV) $=0.07$ for HOS and CV $=0.06$ for NOS, based on Murdoch et al. (2019). Measurement error was also simulated for juvenile recruitment $R$ using $\mathrm{CV}=0.11$, based on the mean of the stock-averaged CV values reported for spring Chinook Salmon from the Twisp and Methow populations (rotary screw trap data provided by WDFW). This yielded the "observed" values for each simulation $i$ and year $y$ as follows:

Observed NOS: $\widetilde{N O S}_{i y}=N O S_{i j}+e_{N O S, i y}$ where $e_{N O S, i y}$ is randomly generated from the $N\left(0, \sigma_{N O S, i y}^{2}\right)$ distribution with $\sigma_{N O S, i y}^{2}=N O S_{i y}^{2} \times C V_{N O S}^{2}$

Observed HOS: $\widetilde{H O S}_{i y}=\operatorname{HOS}_{i j}+e_{H O S, i y}$ where $e_{H O S, i y}$ is randomly generated from the $N\left(0, \sigma_{H O S, i y}^{2}\right)$ distribution with $\sigma_{H O S, i y}^{2}=H O S_{i y}^{2} \times C V_{H O S}^{2}$.

Observed S: $\tilde{S}_{i y}=\widetilde{N O S}_{i y}+\widetilde{H O S}_{i y}$
Observed pHOS: $\widetilde{p H O S}_{i y}=\frac{\widetilde{H O S}_{i y}}{\tilde{S}_{i y}}$

Observed R: $\tilde{R}_{i y}=R_{i j}+e_{R, i y}$ where $e_{R, i y}$ is randomly generated from the $N\left(0, \sigma_{R, i y}^{2}\right)$ distribution for $\sigma_{R, i y}^{2}=R_{i y}^{2} \times C V_{R}^{2}$.

For each simulated data set, the Ricker model in equation (2) was fit to the simulated observations of spawner, recruit, and pHOS data both with and without measurement error in the spawners and pHOS values. For each simulation, the bootstrap was used to calculate a $90 \%$ bootstrap confidence interval for the pHOS regression coefficient $\beta_{P}$ using 1,000 bootstrap samples. The statistical power was computed as the proportion of the simulated data sets for which the upper limit of the $90 \%$ bootstrap confidence interval for $\beta_{P}$ was $<0$. Because the power estimates were the result of simulations, the bootstrap was again used on the simulated data sets to generate $95 \%$ confidence intervals on the power estimates.

Table 1. Spawner and juvenile recruit data for Chiwawa Spring Chinook Salmon. Value of HOS was calculated as $\mathrm{S}^{*} \mathrm{pHOS}$, and value of NOS was calculated as S-HOS.

| Brood Year | Spawners $(\mathrm{S})$ | Smolts $(\mathrm{R})$ | pHOS | HOS | NOS |
| :--- | :--- | :--- | :--- | :--- | :--- |
| 1991 | 242 | 42,525 | 0.00 | 0 | 242 |
| 1992 | 676 | 39,723 | 0.00 | 0 | 676 |
| 1993 | 233 | 8,662 | 0.01 | 2 | 231 |
| 1994 | 184 | 16,472 | 0.33 | 61 | 123 |
| 1995 | 33 | 3,830 | 1.00 | 33 | 0 |
| 1996 | 58 | 15,475 | 0.29 | 17 | 41 |
| 1997 | 182 | 27,555 | 0.67 | 122 | 60 |
| 1998 | 91 | 19,257 | 0.35 | 32 | 59 |
| 1999 | 94 | 10,931 | 0.07 | 7 | 87 |
| 2000 | 346 | 39,812 | 0.33 | 114 | 232 |
| 2001 | 1,725 | 79,814 | 0.71 | 1,225 | 500 |
| 2002 | 707 | 82,845 | 0.64 | 452 | 255 |
| 2003 | 270 | 16,559 | 0.38 | 103 | 167 |
| 2004 | 851 | 67,491 | 0.32 | 272 | 579 |
| 2005 | 599 | 58,833 | 0.77 | 461 | 138 |
| 2006 | 529 | 41,951 | 0.78 | 413 | 116 |
| 2007 | 1,296 | 23,766 | 0.88 | 1,140 | 156 |
| 2008 | 1,158 | 32,849 | 0.84 | 973 | 185 |
| 2009 | 1,347 | 32,979 | 0.78 | 1,051 | 296 |
| 2010 | 1,094 | 47,511 | 0.62 | 678 | 416 |
| 2011 | 2,032 | 37,185 | 0.61 | 1,240 | 792 |
| 2012 | 1,478 | 34,334 | 0.61 | 902 | 576 |
| 2013 | 1,378 | 39,396 | 0.69 | 951 | 427 |
| 2014 | 975 | 37,170 | 0.46 | 449 | 526 |
| 2015 | 967 | 53,344 | 0.65 | 629 | 338 |
| 2016 | 546 | 31,300 | 0.29 | 158 | 388 |
| 2017 | 431 | 39,015 | 0.63 | 272 | 159 |

(a) NOS

Empirical and theoretical CDFs: NOS

(b) S

Empirical and theoretical CDFs: S


Figure 1. Comparison of empirical and fitted negative binomial cumulative distribution functions (CDFs) for natural origin spawners (NOS) data (a) and total spawners (S) data (b). Black dots represent observed data, and horizontal lines represent the distance between observations in empirical step-wise distribution.

## pHOS Effect Size

Figure 2 demonstrates how the value of pHOS is expected to lower predicted recruitment across a wide range of spawner levels for a hypothetical population for two fixed values of the effect size, $\beta_{P}$, according to the model in equation (2). When $\beta_{P}$ is farther from 0 (e.g., $\beta_{P}=-0.5$, Figure 2 a ), a small change in pHOS is expected to produce a larger reduction in recruitment than when $\beta_{P}$ is closer to 0 (e.g., $\beta_{P}=-0.1$, Figure $2 b)$. Larger effects are detectable with smaller sample sizes than smaller effects.

Statistical power is defined in this setting to be the probability of detecting a negative effect of pHOS given that pHOS actually has a negative effect. Calculating it requires specifying both the true effect size, i.e., the true value of $\beta_{P}$, and the change in pHOS value for which the effect is expected to occur. The value of the pHOS effect that is of interest is the effect necessary to reduce recruitment by $5 \%$ for a given level of spawners or stock. Over time, such a reduction will result in a smaller population that will be at increased risk of extinction.

Although any reduction in recruitment is a concern, a reduction in recruitment that occurs for a small change in pHOS values will be harder to avoid than the same reduction that occurs only for a large change in pHOS values. Thus, it is also necessary to specify the difference in pHOS values over which the recruitment reduction is expected to occur. For example, a $5 \%$ reduction recruitment that is associated with a difference in pHOS of only 0.1 (e.g., an increase in pHOS from 0.2 to 0.3 ) represents a stronger pHOS effect than a $5 \%$ reduction in recruitment associated with a difference in pHOS of 0.5 (e.g., an increase in pHOS from 0.2 to 0.7 ). Figure 3 demonstrates how the level of pHOS and the size of the effect $\left(\beta_{P}\right)$ combine to lower the predicted recruitment at a fixed level of spawner. In Figure 3, the predicted recruitment from the Ricker model achieved at $S=1,000$ spawners is reduced by $5 \%$ from the baseline setting (i.e., no hatchery origin spawners, $\mathrm{pHOS}=0$ ) at a lower value of pHOS when $\beta_{P}$ is more negative, indicated by the colored curves crossing the dashed line. When $\beta_{P}$ is closer to 0 , it requires a higher value of pHOS to result in a $5 \%$ reduction in recruitment from a setting without hatchery origin spawners. Thus, values of $\beta_{P}$ that are more negative may be considered more extreme effects of pHOS than values of $\beta_{P}$ that are closer to 0 .


Figure 2. Predicted juvenile recruitment from Ricker model (2) at various levels of spawners S and pHOS for (a) $\beta_{\mathrm{P}}=-0.5$ and (b) $\beta_{\mathrm{P}}=-0.1$. Ricker model parameters: $\mathrm{a}=138, \mathrm{~b}=0.0011$.


Figure 3. Predicted juvenile recruitment from Ricker model at $S=1,000$ spawners for various levels of pHOS and $\beta_{\mathrm{P}}$, the effect of pHOS on recruitment. Dashed line indicates $5 \%$ reduction in recruitment compared to $\mathrm{pHOS}=0$. Ricker model parameters: $\mathrm{a}=138, \mathrm{~b}=0.0011$.

The relationship between the reduction in mean recruitment, the change in pHOS values, and the effect size $\beta_{P}$ can be derived as follows: Let $R_{1}$ be the mean recruitment for a given number of spawners at a baseline pHOS value of $P_{1}$, and $R_{2}$ be the mean recruitment for a given number of spawners for a treatment pHOS value of $P_{2}$; that is:

$$
R_{1}=a e^{\beta_{P} P_{1}} S e^{-b S}
$$

and

$$
R_{2}=a e^{\beta_{P} P_{2}} S e^{-b S}
$$

$R_{2}$ can be expressed in terms of $R_{1}$ by taking the ratio of $R_{2} / R_{1}$ as follows:

$$
\frac{R_{2}}{R_{1}}=\frac{a e^{\beta_{P} P_{2}} S e^{-b S}}{a e^{\beta_{P} P_{1}} S e^{-b S}}=\frac{e^{\beta_{P} P_{2}}}{e^{\beta_{P} P_{1}}}=e^{\beta_{P}\left(P_{2}-P_{1}\right)}
$$

which leads to:

$$
R_{2}=R_{1} e^{\beta_{P}\left(P_{2}-P_{1}\right)}
$$

or

$$
R_{2}=R_{1} e^{\beta_{P} \Delta P_{\text {HOS }}}
$$

where $\Delta P_{H O S}=P_{2}-P_{1}$. Then if $R_{2}$ is a $5 \%$ reduction from $R_{1}$, we have:

$$
\begin{equation*}
\beta_{P}=\frac{\ln (0.95)}{\Delta P_{\text {HOS }}} \tag{6}
\end{equation*}
$$

Thus, for a $5 \%$ reduction in recruitment for a change in pHOS of $\Delta P_{H O S}=0.1$, the pHOS effect size is $\beta_{P}=-0.513$, whereas for $\Delta P_{\text {HOS }}=0.5$, the pHOS effect size is only $\beta_{P}=-0.103$. We calculated power to detect a $5 \%$ reduction in recruitment for a change in pHOS of $\Delta P_{\text {HOS }}=0.1,0.25,0.50$, and 0.75 . These settings are consistent with a pHOS effect of $\beta_{P}=-0.513,-0.205,-0.103$, and -0.068 , respectively.

## Results

Simulated statistical power to detect a negative effect of pHOS was consistently $\leq 0.32$ for all sample sizes considered ( $N \leq 100$ years) and for all but the most extreme pHOS effect sizes considered (i.e., $\beta_{P}=$ -0.068 to -0.205 ), both with and without measurement error (Figure 4). For the largest effect size ( $\beta_{P}=$ -0.513 ), simulated power was as high as 0.54 for a sample size of $N=40$ years without measurement error, and only slightly lower at 0.52 for $N=40$ years when measurement error was incorporated into the simulations. Achieving power of at least 0.70 required as many as $N=70$ years of data. Power greater than 0.80 required 90 years of data. These simulations defined detection of a negative effect of pHOS as a $90 \%$ bootstrap confidence interval that was entirely less than 0 .
(i) Without measurement error

(ii) With measurement error


Figure 4. Statistical power of detecting a negative effect of pHOS for various levels of pHOS effect size $\beta_{\mathrm{P}}$ with (i) and without (ii) measurement error in stock and recruitment data. Power based on 1,000 simulations of stock (spawners) and juvenile recruitment data using Ricker stock-recruitment model fit to data from Chiwawa spring Chinook Salmon, 1991-2017: $\mathrm{a}=138, \mathrm{~b}=0.0011, \sigma=0.4386$ (equation (2)). Type I error probability $=0.10$. Measurement error (ii) was incorporated into simulated observations of
spawner and recruitment data using coefficient of variation (CV) $=0.07$ for HOS, $\mathrm{CV}=0.06$ for NOS, and $\mathrm{CV}=0.11$ for recruitment. Shaded region $=95 \%$ bootstrap confidence interval using 300 bootstrap samples from the simulated data.

## Conclusions

Simulated power to detect a negative effect of pHOS on juvenile recruitment was low for all but the strongest effect sizes and for studies shorter than approximately 70 years. The low power values resulted from the high variability in residuals from the Ricker stock-recruitment model $(\sigma=0.4386)$ estimated from its fit to 27 years of data from the Chiwawa Spring Chinook salmon population (1991-2017).

The Chiwawa data set is the largest of the juvenile productivity data sets available for Chinook salmon and steelhead. The alterative data sets either failed to meet the modeling assumptions for the Ricker model (e.g., Methow and Twisp steelhead) or else had higher error variance about the fitted Ricker model (e.g., other spring Chinook salmon populations) than seen for the Chiwawa spring Chinook salmon population. Thus, it is expected that the power to detect a negative effect of pHOS on juvenile productivity would be highest for the Chiwawa spring Chinook population. Low power for this population would be compounded for the other populations with shorter time series or more complex population dynamics. As a consequence, it is unlikely that assessment of stock-recruitment curves similar to the Ricker model will be sufficient to detect a negative effect of pHOS in time to mediate any such effect. Other methods of monitoring and assessment are recommended to evaluate the effect of pHOS on juvenile productivity of anadromous salmonids in the upper Columbia River basin.

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## COLUMBIA BASIN RESEARCH

# The Effect of Hatchery Programs on Proportionate Natural Influence (PNI) in the Upper Columbia Basin 

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#### Abstract

Hatchery programs and natural populations in the Wenatchee and Methow subbasins were managed under the Proportionate Natural Influence (PNI) strategy, whereby gene flow between the hatchery and natural populations was manipulated to achieve greater overall genetic influence from the natural population as opposed to the hatchery population. The target PNI value was 0.67 . Steelhead programs in the Wenatchee and Methow had PNI averages or medians below 0.67 . The new management regime for steelhead in the Methow was not initiated until 2017. Therefore, data in this report do not reflect the future management of the steelhead in the Methow subbasin for gene flow. Spring Chinook Salmon PNI in the Wenatchee Subbasin was below 0.67 for all populations under the contemporary management strategy but exceeded 0.50 in all cases. Methow subbasin spring Chinook PNI was 0.29 , and has not increased appreciably in the two years of adult returns following reduction in program sizes in 2013. Wenatchee and Methow summer Chinook both had PNI means that exceeded 0.67. Both of these programs appear successful in meeting PNI objectives with the Wenatchee PNI an impressive 0.87 . The results of this analysis indicate that the Spring Chinook and steelhead programs require adjustment to meet the PNI targets. In some cases, the PNI values may improve as new management strategies mature. The Methow steelhead and particularly Methow spring Chinook may require substantial management changes in order to achieve the gene-flow objectives. In general, PNI targets were not met for small populations (spring Chinook Salmon and steelhead) but were for large populations (summer Chinook Salmon).


## Introduction

Conservation hatchery programs integrate the spawning of hatchery- and natural-origin fish into a single gene pool. However, the gene flow among origins has the potential to impose genetic risk associated with domestication selection. Hatchery programs and natural populations in the Wenatchee and Methow subbasins were managed under the Proportionate Natural Influence (PNI; HSRG 2009) strategy, whereby gene flow between the hatchery and natural populations is manipulated to achieve greater overall genetic influence from the natural population as opposed to the hatchery population. Two components may be manipulated to achieve a desired PNI: the proportion of hatchery-origin spawners ( pHOS ) in nature and the proportion of natural-origin broodstock ( pNOB ) in the hatchery. The PNI is an index of domestication selection that is relatively easy to measure and ranges from 0 to 1 . A PNI of 0 indicates that all of the selection is from gene flow among hatchery-origin fish. In contrast, a PNI of 1 indicates that all of the selection is from gene flow among natural-origin fish.
Management targets of $>0.67$ have been established for PNI for the various programs under the Endangered Species Act and for many unlisted programs of conservation importance (Mobrand et al. 2005; Paquet et al. 2011; Pearsons et al. 2020). These targets are intended to minimize the risk of domestication selection which could decrease the fitness of fish spawning in the natural environment. We estimated PNI for the spring Chinook Salmon, summer Chinook Salmon, and summer steelhead populations and programs in the Wenatchee and Methow subbasins that have PNI targets of 0.67.

## Study Area

The study examined PNI values in the Wenatchee and Methow subbasins. In the Wenatchee, PNI for spring Chinook populations in the Chiwawa River, Nason Creek, and White River was estimated separately, plus PNI was estimated for the population upstream of Tumwater Dam. In the Methow Subbasin, PNI for spring Chinook was evaluated at the subbasin level, according to terms in the ESA permits. PNI for summer steelhead was evaluated at the subbasin level in the Wenatchee and Methow subbasins. Similarly, PNI for summer Chinook was evaluated at the subbasin level in the Wenatchee and Methow subbasins.

## Methods

## Data Collection and Derived Metrics

Information needed to estimate PNI included the number of natural-origin fish in hatchery broodstock (NOB), the number of hatchery-origin fish in hatchery broodstock (HOB), the number of hatchery-origin spawners (HOS), and the number of natural-origin spawners (NOS). From these, we calculated the proportion of natural-origin fish in hatchery broodstock $(\mathrm{pNOB}=\mathrm{NOB} /(\mathrm{NOB}+\mathrm{HOB}))$, the proportion of hatchery-origin fish in hatchery broodstock $(\mathrm{pHOB}=\mathrm{HOB} /(\mathrm{NOB}+\mathrm{HOB}))$, the proportion of hatchery-origin spawners $(\mathrm{pHOS}=\mathrm{HOS} /$ $($ NOS +HOS$)$ ), and the proportion of natural-origin spawners (pNOS $=$ NOS $/(\mathrm{NOS}+\mathrm{HOS})$ ). For all programs, pNOB and pHOB were estimated based on the total number of adults collected for broodstock.

Methods used to estimate NOS and HOS varied depending on stock. Spawning-ground surveys were used to estimate NOS and HOS for spring and summer Chinook Salmon in the Wenatchee and Methow subbasins. For most stocks, these surveys have been conducted since at least 1981 to determine the abundance, distribution, and origin of spring and summer Chinook Salmon spawners. Field methods were consistent with those described in Gallagher et al. (2007) and Murdoch et al. (2010). Surveyors walked or floated the entire distribution of Chinook Salmon spawning habitat and identified and counted new redds weekly throughout the spawning season. Redds were flagged and locations recorded on a Global-Positioning-System device to avoid recounting in subsequent surveys. Chinook Salmon carcasses were also counted and examined to determine sex, origin (e.g., hatchery or natural origin; nearly all hatchery-origin fish were marked with a tag such as a coded wire tag (CWT) and/or adipose fin clipped prior to release), size, distribution, and other biological characteristics. In addition, scales collected from carcasses were used to determine fish age and confirm origin.

We expanded spawning-ground data into derived estimates of total adult abundance (combined hatchery- and natural-origin adults), NOS abundance, and HOS abundance. We assumed that each female made one redd (Murdoch et al. 2009a), female carcass location was a good surrogate for spawning location by origin (Murdoch et al. 2009b), and that redd counts and carcass data could be used to estimate spawning escapement for hatchery- and natural-origin Chinook Salmon (Murdoch et al. 2010). Total adult spawning abundance was estimated by multiplying redd counts by the male:female sex ratio estimated at Dryden Dam (for Wenatchee River summer Chinook Salmon), Tumwater Dam (for Wenatchee River spring Chinook Salmon), or at Wells Dam (for Methow programs). Adult fish removed from the natural spawning population by gene-flow management and broodstock-collection activities were subtracted from the data used to estimate the sex ratios. NOS and HOS abundances were estimated by multiplying the respective proportion of natural-origin carcasses and hatcheryorigin carcasses by the total adult spawning abundance. From these data and the equations described above, we calculated pNOS and pHOS for each Chinook salmon program.

Several methods and combinations of methods have been used to estimate steelhead NOS and HOS abundances within the Wenatchee and Methow subbasins. Within the Wenatchee River Subbasin, spawning escapements have been estimated based on run reconstruction and markrecapture (Passive Integrated Transponder (PIT) tag) models. Prior to 2014, only run reconstruction was used to estimate steelhead escapements within the Wenatchee River. Steelhead run reconstruction was based on the number of hatchery- and natural-origin steelhead observed at Priest Rapids and Wells dams and apportioned to Upper Columbia subbasins based on previously conducted radio-telemetry studies (English et al. 2001; 2003) and differences in dam counts. Run escapement to each of the subbasins was then adjusted for adult management, harvest, broodstock collection, and an assumed $10 \%$ pre-spawn mortality to estimate spawning escapement. Beginning in 2014, steelhead escapements in tributaries were estimated using PITtag mark-recapture techniques (Truscott et al. 2017), while observer-efficiency-expanded redd counts were used to estimate escapements in the mainstem Wenatchee River (See 2021). Total redd counts were also used to estimate escapements in the lower portions of the main tributaries (downstream from the PIT-tag interrogation sites). Redd counts were expanded by multiplying redd counts by the male:female sex ratio estimated based on detections of PIT-tagged males and females within the Wenatchee River Subbasin.

Steelhead spawning escapements in the Methow River Subbasin were based on four population components: (1) conservation programs including the Twisp River and Winthrop National Fish Hatchery (WNFH), (2) a safety-net program that included Methow River releases from Wells Hatchery, (3) annual spawning component of the upper Methow River and tributaries (also included Beaver, Gold, and Libby creeks), and (4) annual spawning component of the mainstem Methow River downstream from the Methow Fish Hatchery. Overall NOS and HOS abundances were estimated based on PIT-tag detections at the lower Methow River instream PIT-tag array (LMR) expanded by estimated efficiency of the array and the PIT-tag rate of hatchery- and natural-origin steelhead tagged at Priest Rapids Dam within each return year since 2014. All adipose fin-clipped steelhead lacking a CWT were assumed to be from the Wells safety-net program, while those with a CWT were assumed to be from the WNFH conservation program. Steelhead with a CWT but without an adipose fin-clip were assumed to be from the Twisp River conservation program. Some returning fish from previous marking strategies (e.g., yellow elastomer tags) were included as conservation program fish if their parental origin was greater than or equal to 0.5 natural-origin steelhead. Similarly, known conservation program fish from other subbasins (e.g., Omak Creek in the Okanogan River Subbasin) were pooled with other conservation program returns in the Methow River Subbasin. All other hatchery-origin steelhead were considered safety-net program fish for modeling purposes.

After estimating the total steelhead escapement to the Methow River Subbasin, escapement estimates for the upper Methow River and tributaries were subtracted from the LMR-generated subbasin estimate. This provided a separate escapement estimate for the conservation (upper Methow River and tributaries) and management (lower Methow River mainstem) areas. From the management area estimate, we subtracted all known steelhead removed during broodstock collection or adult-management activities, including sport fisheries, broodstock collections, or management (removal) of excess hatchery-origin fish at the Methow Fish Hatchery and WNFH. From the conservation area estimate, we subtracted all known steelhead removed at the Twisp River weir for broodstock or to reduce escapement of hatcheryorigin fish. The proportion of natural-origin steelhead in the broods from which the returning conservation program adults originated in each spawn year was estimated as an average of the pNOB for each returning age class and program (i.e., Twisp 1- and 2-salt, WNFH 1- and 2-salt). The genetic parentage of safety-net program adults was assumed to be $75 \%$ safety-net program fish and $25 \%$ conservation program fish for broods prior to 2018.

## Data Analysis

Although PNI is estimated using pNOB and pHOS , different methods can be used to calculate PNI. According to authorized annual take permits, PNI is calculated using the PNI approximate equation 11 (HSRG 2009). However, in this report, we used Ford's (2002) equations 5 and 6 with a heritability of 0.3 and a selection strength of three standard deviations to calculate PNI for all Wenatchee River stocks. This approach is more accurate than using the PNI approximate equation. For Methow River stocks, PNI was calculated using a multi-population model developed by Busack (2015).

The interpretation of PNI is straightforward. The larger the PNI value, the greater the strength of selection in the natural environment relative to that of the hatchery environment. For
the natural environment to dominate selection, PNI should be greater than 0.50, and integrated populations should have a PNI of at least 0.67 (HSRG/WDFW/NWIFC 2004; HSRG 2009; Mobrand et al. 2005; Paquet et al. 2011). For the Wenatchee and Methow River steelhead programs, PNI criteria were implemented in accordance with permits to achieve a subbasin-wide, five-year running average of $\mathrm{PNI} \geq 0.67$. In years when the natural-origin steelhead escapement is low (i.e., $<500$ fish in the Methow and $<433$ fish in the Wenatchee), the populations will be managed to meet escapement goals rather than PNI.

## Results

## Wenatchee River Steelhead

As described above, for the Wenatchee River steelhead program, PNI criteria are implemented in accordance with Permit 18583 to achieve a subbasin-wide, five-year running average of $\mathrm{PNI} \geq 0.67$. In years when the natural-origin escapement is low (i.e., $<433$ fish), the Wenatchee River steelhead population will be managed to meet escapement goals rather than PNI.

For brood years 2001-2013, prior to the reduction in smolt production, PNI values were consistently less than 0.67 and the five-year running average ranged from 0.49 to 0.53 (Table 1). For brood years 2014-2018, the period after reduction in smolt production, PNI values were generally less than 0.67 and the five-year running average was 0.55 (Table 1). Because of low escapement in 2017, the Wenatchee steelhead population was managed to meet escapement goals rather than PNI.

Table 1. PNI values for the Wenatchee steelhead supplementation program for brood years 2001-2018. PNI estimates for the period 2001-2013 are based on estimates of spawners upstream from Tumwater Dam; PNI estimates for the period 2014-present are based on mark-recapture modeling for the entire Wenatchee River Subbasin.

| Brood Year | Spawners ${ }^{\text {a }}$ |  |  | Broodstock |  |  | PNI | $\begin{gathered} \hline \text { PNI (5- } \\ \text { yr } \\ \text { mean) } \\ \hline \end{gathered}$ |
| :---: | :---: | :---: | :---: | :---: | :---: | :---: | :---: | :---: |
|  | NOS | HOS | pHOS | NOB | HOB | pNOB |  |  |
| 2001 | 158 | 127 | 0.45 | 51 | 103 | 0.33 | 0.45 | -- |
| 2002 | 731 | 542 | 0.43 | 96 | 64 | 0.60 | 0.59 | -- |
| 2003 | 355 | 350 | 0.50 | 49 | 90 | 0.35 | 0.43 | -- |
| 2004 | 371 | 445 | 0.55 | 75 | 61 | 0.55 | 0.51 | -- |
| 2005 | 690 | 862 | 0.56 | 87 | 104 | 0.46 | 0.47 | 0.49 |
| 2006 | 253 | 210 | 0.45 | 93 | 69 | 0.57 | 0.57 | 0.51 |
| 2007 | 145 | 115 | 0.44 | 76 | 58 | 0.57 | 0.58 | 0.51 |
| 2008 | 168 | 279 | 0.62 | 77 | 54 | 0.59 | 0.50 | 0.53 |
| 2009 | 171 | 545 | 0.76 | 86 | 73 | 0.54 | 0.43 | 0.51 |
| 2010 | 524 | 970 | 0.65 | 96 | 75 | 0.56 | 0.48 | 0.51 |
| 2011 | 351 | 472 | 0.57 | 91 | 70 | 0.57 | 0.51 | 0.50 |
| 2012 | 381 | 209 | 0.35 | 59 | 65 | 0.48 | 0.59 | 0.50 |
| 2013 | 322 | 148 | 0.31 | 49 | 68 | 0.42 | 0.59 | 0.52 |
| Average $^{\text {b }}$ | 355 | 406 | 0.51 | 76 | 73 | 0.51 | 0.52 | 0.51 |
| Median ${ }^{\text {b }}$ | 351 | 350 | 0.50 | 77 | 69 | 0.55 | 0.51 | 0.51 |
| 2014 | 901 | 477 | 0.35 | 62 | 66 | 0.48 | 0.59 | -- |
| 2015 | 988 | 711 | 0.42 | 58 | 52 | 0.53 | 0.57 | -- |
| 2016 | 587 | 372 | 0.39 | 64 | 66 | 0.49 | 0.57 | -- |
| 2017 | 198 | 232 | 0.54 | 56 | 63 | 0.47 | 0.48 | -- |
| 2018 | 324 | 165 | 0.34 | 70 | 75 | 0.48 | 0.52 | 0.55 |
| Average $^{\text {c }}$ | 524 | 391 | 0.41 | 62 | 64 | 0.49 | 0.55 | 0.55 |
| Median ${ }^{\text {c }}$ | 456 | 272 | 0.39 | 62 | 66 | 0.48 | 0.57 | 0.55 |

${ }^{a}$ The presence of eroded fins or missing adipose fins was used to distinguish hatchery fish from wild fish during video monitoring at Tumwater Dam. Unknown-origin fish (i.e., undetermined by scale analysis, no elastomer, no CWT, no fin clips, and no additional hatchery marks) were considered naturally produced. Therefore, because not all hatchery fish have eroded fins or missing adipose fins, it is likely we are underestimating WxW-cross hatchery-origin returns based on video monitoring. The PNI estimates are appropriate for steelhead spawning upstream from Tumwater Dam but may not represent PNI for steelhead spawning downstream from Tumwater Dam.
${ }^{\mathrm{b}}$ Descriptive statistics using escapements estimated upstream from Tumwater Dam.
${ }^{\text {c }}$ Descriptive statistics using escapement estimates based on mark-recapture modeling.

## Methow River Steelhead

For the Methow River steelhead program, PNI criteria are implemented in accordance with Permit 23163 to achieve a subbasin-wide, five-year running average of $\mathrm{PNI} \geq 0.67$ by 2022. The Methow Subbasin is managed in two zones: 1) the upper Methow River and primary tributaries are to achieve pHOS of $\leq 0.25$, and 2 ) the remainder of the subbasin is managed to achieve an overall subbasin PNI of 0.67. Expected PNI from 2018 to 2021 is estimated to be $\geq$ 0.45 (NMFS, 2017). In years when the natural-origin escapement is low (i.e., $<500$ fish), the Methow River steelhead population will be managed to meet 500 total spawners rather than PNI.

For brood years 2014-2018, PNI has been below 0.67 and ranged from 0.43 to 0.59 with a five-year running average of 0.50 (Table 2). The proportion of hatchery-origin spawners within conservation areas has ranged from 0.36 to 0.61 , while the proportion of hatchery-origin spawners within management areas has ranged from 0.51 to 0.83 .

Table 2. PNI and proportion of hatchery-origin fish on spawning grounds ( $\mathrm{pHOS} \mathrm{)} \mathrm{calculated} \mathrm{by}$ spawn year in the Methow River conservation and management areas based on expanded PIT-tag observations. PNI was estimated using the model described by Busack (2015) using the proportion of hatchery-origin returns in each area that derive from conservation program returns (HOR-c), safety-net program returns (HOR-sn) and natural-origin returns (NOR). The net proportion of natural-origin fish in the broods from which the HOR-c returns originated (pNOB HOR-c) was calculated as a mean value from contributing adult broods. The genetic contribution of the returning HOR-sn component (not shown) was estimated as being $75 \%$ from safety-net adults, and $25 \%$ from conservation program adults prior to 2019 .

| Brood <br> Year | Conservation areas |  |  |  | Management area |  |  |  | Overall | pNOB <br> PON |
| :---: | :---: | :---: | :---: | :---: | :---: | :---: | :---: | :---: | :---: | :---: |
|  | 0.61 | 0.13 | 0.26 | 0.39 | 0.29 | 0.34 | 0.36 | 0.71 | 0.50 | 0.53 |
| 2015 | 0.54 | 0.14 | 0.32 | 0.46 | 0.17 | 0.11 | 0.71 | 0.83 | 0.46 | 0.56 |
| 2016 | 0.64 | 0.14 | 0.22 | 0.36 | 0.35 | 0.15 | 0.50 | 0.65 | 0.53 | 0.56 |
| 2017 | 0.39 | 0.18 | 0.44 | 0.61 | 0.25 | 0.19 | 0.56 | 0.75 | 0.43 | 0.71 |
| 2018 | 0.62 | 0.14 | 0.24 | 0.38 | 0.49 | 0.46 | 0.06 | 0.51 | 0.59 | 0.83 |
| Average | $\mathbf{0 . 5 6}$ | $\mathbf{0 . 1 5}$ | $\mathbf{0 . 3 0}$ | $\mathbf{0 . 4 4}$ | $\mathbf{0 . 3 1}$ | $\mathbf{0 . 2 5}$ | $\mathbf{0 . 4 4}$ | $\mathbf{0 . 6 9}$ | $\mathbf{0 . 5 0}$ | $\boldsymbol{0 . 6 4}$ |
| Median | $\mathbf{0 . 6 1}$ | $\mathbf{0 . 1 4}$ | $\mathbf{0 . 2 6}$ | $\mathbf{0 . 3 9}$ | $\mathbf{0 . 2 9}$ | $\mathbf{0 . 1 9}$ | $\mathbf{0 . 5 0}$ | $\mathbf{0 . 7 1}$ | $\mathbf{0 . 5 0}$ | $\boldsymbol{0 . 5 6}$ |

## Chiwawa River Spring Chinook Salmon

For brood years 1989-2011, prior to the reduction in smolt production, PNI values ranged from 0.26 to 1.00 (Table 3). PNI values for brood years 1989-1994 were greater than or equal to 0.67 . For brood years 2012-2018, the period after reduction in smolt production, PNI values were generally less than 0.67 , except for brood year 2016, which was 0.68 (Table 3).

Table 3. PNI values for the Chiwawa spring Chinook supplementation program for brood years 1989-2018.

| Brood Year | Spawners |  |  | Broodstock |  |  | PNI ${ }^{\text {a }}$ |
| :---: | :---: | :---: | :---: | :---: | :---: | :---: | :---: |
|  | NOS | HOS | pHOS | NOB | HOB | pNOB |  |
| 1989 | 713 | 0 | 0.00 | 28 | 0 | 1.00 | 1.00 |
| 1990 | 571 | 0 | 0.00 | 18 | 0 | 1.00 | 1.00 |
| 1991 | 242 | 0 | 0.00 | 27 | 0 | 1.00 | 1.00 |
| 1992 | 676 | 0 | 0.00 | 78 | 0 | 1.00 | 1.00 |
| 1993 | 231 | 2 | 0.01 | 94 | 0 | 1.00 | 0.99 |
| 1994 | 123 | 61 | 0.33 | 8 | 4 | 0.67 | 0.68 |
| 1995 | 0 | 33 | 1.00 | No Program |  |  |  |
| 1996 | 41 | 17 | 0.29 | 8 | 10 | 0.44 | 0.62 |
| 1997 | 60 | 122 | 0.67 | 32 | 79 | 0.29 | 0.32 |
| 1998 | 59 | 32 | 0.35 | 13 | 34 | 0.28 | 0.47 |
| 1999 | 87 | 7 | 0.07 | No Program |  |  |  |
| 2000 | 233 | 113 | 0.33 | 9 | 21 | 0.30 | 0.50 |
| 2001 | 506 | 1219 | 0.71 | 113 | 259 | 0.30 | 0.32 |
| 2002 | 254 | 453 | 0.64 | 20 | 51 | 0.28 | 0.33 |
| 2003 | 168 | 102 | 0.38 | 41 | 53 | 0.44 | 0.55 |
| 2004 | 574 | 277 | 0.33 | 83 | 132 | 0.39 | 0.56 |
| 2005 | 139 | 460 | 0.77 | 91 | 181 | 0.33 | 0.32 |
| 2006 | 114 | 415 | 0.78 | 91 | 224 | 0.29 | 0.29 |
| 2007 | 155 | 1141 | 0.88 | 43 | 104 | 0.29 | 0.27 |
| 2008 | 190 | 968 | 0.84 | 83 | 220 | 0.27 | 0.26 |
| 2009 | 297 | 1050 | 0.78 | 96 | 111 | 0.46 | 0.39 |
| 2010 | 419 | 675 | 0.62 | 77 | 98 | 0.44 | 0.43 |
| 2011 | 801 | 1231 | 0.61 | 80 | 93 | 0.46 | 0.45 |
| 2012 | 574 | 904 | 0.61 | 66 | 45 | 0.59 | 0.50 |
| 2013 | 422 | 956 | 0.69 | 68 | 2 | 0.97 | 0.59 |
| 2014 | 523 | 452 | 0.46 | 58 | 12 | 0.83 | 0.65 |
| 2015 | 337 | 630 | 0.65 | 64 | 0 | 1.00 | 0.61 |
| 2016 | 389 | 157 | 0.29 | 57 | 42 | 0.58 | 0.68 |
| 2017 | 160 | 271 | 0.63 | 50 | 18 | 0.74 | 0.55 |
| 2018 | 166 | 456 | 0.73 | 30 | 57 | 0.34 | 0.34 |
| Average | 307 | 407 | 0.48 | 55 | 66 | 0.57 | 0.56 |
| Median | 238 | 274 | 0.61 | 58 | 44 | 0.45 | 0.53 |

${ }^{\text {a PNI }}$ was calculated previously using PNI approximate equation 11 (HSRG 2009; their Appendix A). All PNI values presented here were recalculated by iterating Ford's (2002) equations 5 and 6 to equilibrium using a heritability of 0.3 and a selection strength of three standard deviations. C. Busack, NOAA Fisheries, 21 March 2016, provided the model for calculating PNI.

## Nason Creek Spring Chinook Salmon

For brood years 1989-2012, when no brood stock were collected for the Nason Creek Program, the PNI values ranged from 0.28 to 1.00 (Table 4). During this period, PNI values varied over time because of Chiwawa spring Chinook straying into Nason Creek. For brood years 2013-2018, a period when broodstock were collected for the Nason Creek Program, PNI values for the Nason Creek Program ranged from 0.38 to 0.79 (Table 4).

Table 4. PNI values of hatchery spring Chinook spawning in Nason Creek, brood years 19892018. See notes below the table for description of each metric.

| Brood Year | Spawners |  |  |  |  | Broodstock |  |  | PNI |
| :---: | :---: | :---: | :---: | :---: | :---: | :---: | :---: | :---: | :---: |
|  | NOS | $\mathrm{HOS}_{N}$ | HOSs | $\mathrm{pHOS}_{\mathrm{N}}$ | pHOS ${ }_{\text {N }+\mathrm{s}}$ | $\mathrm{NOB}_{\mathrm{N}}$ | $\mathrm{HOB}_{\mathrm{N}}$ | pNOB |  |
| 1989 | 288 | 0 | 0 | 0.00 | 0.00 | 0 | 0 | 1.00 | 1.00 |
| 1990 | 235 | 0 | 0 | 0.00 | 0.00 | 0 | 0 | 1.00 | 1.00 |
| 1991 | 156 | 0 | 0 | 0.00 | 0.00 | 0 | 0 | 1.00 | 1.00 |
| 1992 | 181 | 0 | 0 | 0.00 | 0.00 | 0 | 0 | 1.00 | 1.00 |
| 1993 | 430 | 0 | 61 | 0.00 | 0.12 | 0 | 0 | 1.00 | 0.90 |
| 1994 | 60 | 0 | 0 | 0.00 | 0.00 | 0 | 0 | 0.67 | 1.00 |
| 1995 | 18 | 0 | 0 | 0.00 | 0.00 | 0 | 0 | 0.00 | 1.00 |
| 1996 | 58 | 0 | 25 | 0.00 | 0.30 | 0 | 0 | 0.44 | 0.61 |
| 1997 | 67 | 0 | 55 | 0.00 | 0.45 | 0 | 0 | 0.29 | 0.42 |
| 1998 | 61 | 0 | 3 | 0.00 | 0.05 | 0 | 0 | 0.28 | 0.86 |
| 1999 | 22 | 0 | 0 | 0.00 | 0.00 | 0 | 0 | 0.00 | 1.00 |
| 2000 | 189 | 0 | 81 | 0.00 | 0.30 | 0 | 0 | 0.30 | 0.52 |
| 2001 | 257 | 0 | 341 | 0.00 | 0.57 | 0 | 0 | 0.30 | 0.37 |
| 2002 | 313 | 0 | 290 | 0.00 | 0.48 | 0 | 0 | 0.28 | 0.39 |
| 2003 | 152 | 0 | 50 | 0.00 | 0.25 | 0 | 0 | 0.44 | 0.65 |
| 2004 | 297 | 0 | 210 | 0.00 | 0.41 | 0 | 0 | 0.39 | 0.51 |
| 2005 | 81 | 0 | 266 | 0.00 | 0.77 | 0 | 0 | 0.33 | 0.32 |
| 2006 | 117 | 0 | 154 | 0.00 | 0.57 | 0 | 0 | 0.29 | 0.36 |
| 2007 | 83 | 0 | 380 | 0.00 | 0.82 | 0 | 0 | 0.29 | 0.28 |
| 2008 | 139 | 0 | 425 | 0.00 | 0.75 | 0 | 0 | 0.27 | 0.29 |
| 2009 | 163 | 0 | 371 | 0.00 | 0.69 | 0 | 0 | 0.46 | 0.42 |
| 2010 | 59 | 0 | 349 | 0.00 | 0.86 | 0 | 0 | 0.44 | 0.35 |
| 2011 | 250 | 0 | 452 | 0.00 | 0.64 | 0 | 0 | 0.46 | 0.43 |
| 2012 | 220 | 0 | 474 | 0.00 | 0.68 | 0 | 0 | 0.66 | 0.50 |
| Average* | 159 | 0 | 166 | 0.00 | 0.36 | 0 | 0 | 0.48 | 0.63 |
| Median* | 154 | 0 | 71 | 0.00 | 0.36 | 0 | 0 | 0.42 | 0.52 |
| 2013 | 70 | 0 | 339 | 0.00 | 0.83 | 20 | 5 | 0.80 | 0.50 |
| 2014 | 165 | 0 | 66 | 0.00 | 0.29 | 21 | 0 | 1.00 | 0.78 |


| Brood Year | Spawners |  |  |  |  | Broodstock |  |  | PNI |
| :---: | :---: | :---: | :---: | :---: | :---: | :---: | :---: | :---: | :---: |
|  | NOS | $\mathrm{HOS}_{\mathrm{N}}$ | HOS ${ }_{\text {s }}$ | $\mathrm{pHOS}_{\mathrm{N}}$ | pHOS ${ }_{\text {N }+\mathrm{s}}$ | $\mathrm{NOB}_{\mathrm{N}}$ | $\mathrm{HOB}_{\mathrm{N}}$ | pNOB |  |
| 2015 | 130 | 0 | 21 | 0.00 | 0.14 | 60 | 63 | 0.49 | 0.79 |
| 2016 | 120 | 11 | 18 | 0.07 | 0.19 | 70 | 66 | 0.51 | 0.74 |
| 2017 | 61 | 32 | 39 | 0.24 | 0.54 | 70 | 64 | 0.52 | 0.51 |
| 2018 | 21 | 70 | 78 | 0.41 | 0.88 | 53 | 54 | 0.50 | 0.38 |
| Average** | 95 | 19 | 94 | 0.12 | 0.48 | 49 | 42 | 0.64 | 0.62 |
| Median** | 95 | 6 | 53 | 0.04 | 0.42 | 57 | 59 | 0.52 | 0.63 |

$\mathbf{H O S}_{\mathbf{N}}=$ hatchery-origin spawners in Nason Creek from the Nason Creek spring Chinook Supplementation Program.
$\mathbf{p H O S}_{\mathbf{N}}=$ proportion of hatchery-origin spawners from Nason Creek spring Chinook Supplementation Program.
$\mathbf{H O S}_{\mathbf{s}}=$ stray hatchery-origin spawners in Nason Creek.
$\mathbf{p H O S}_{\mathbf{s}}=$ proportion of stray hatchery-origin spawners.
$\mathbf{N O B}_{\mathbf{N}}=$ natural-origin broodstock spawned in the Nason Creek spring Chinook Supplementation Program.
$\mathbf{H O B}_{\mathbf{N}}=$ hatchery-origin broodstock spawned in the Nason Creek spring Chinook Supplementation Program.
pNOB = proportion of hatchery-origin broodstock. Because of the high incidence of strays to Nason Creek from the Chiwawa River spring Chinook program, pNOB values from the Chiwawa program were used to estimate PNI values during the period from 1989 to 2012 (italicized). The weighting for those years was $100 \%$ based on the Chiwawa program broodstock selection, because there had been no hatchery returns from the Nason Creek spring Chinook program.
$\mathbf{P N I}_{\mathbf{N}}=$ Proportionate Natural Influence for Nason Creek spring Chinook calculated using the gene-flow model for multiple programs.

* Average and median for the period 1989-2012, a period when no brood stock were collected for the Nason Creek Program.
** Average and median for the period 2013-present, a period when brood stock was collected for the Nason Creek Program.


## White River Spring Chinook Salmon

For brood years 1989-2000, PNI values ranged from 0.95 to 1.00 (Table 5). For brood years 2001-2013, PNI during the White River Program averaged 0.60 (range, 0.33-1.00) and most of the hatchery spawners originated from the Chiwawa River Hatchery Program (Table 5). The captive brood program ended with brood year 2013.

Table 5. PNI values for hatchery spring Chinook spawning in the White River, brood years 1989-2013. See notes below the table for description of each metric.

| Brood Year | Spawners |  |  |  |  | Broodstock |  |  | PNI |
| :---: | :---: | :---: | :---: | :---: | :---: | :---: | :---: | :---: | :---: |
|  | NOS | $\mathrm{HOS}_{w}$ | HOSs | pHOS ${ }_{\text {w }}$ | pHOSs | $\mathrm{NOB}_{\mathrm{N}}$ | $\mathrm{HOB}_{\mathrm{N}}$ | pNOB |  |
| 1989 | 145 | 0 | 0 | 0.00 | 0.00 | 0 | 0 | 1.00 | 1.00 |
| 1990 | 49 | 0 | 0 | 0.00 | 0.00 | 0 | 0 | 1.00 | 1.00 |
| 1991 | 49 | 0 | 0 | 0.00 | 0.00 | 0 | 0 | 1.00 | 1.00 |
| 1992 | 78 | 0 | 0 | 0.00 | 0.00 | 0 | 0 | 1.00 | 1.00 |
| 1993 | 138 | 0 | 7 | 0.00 | 0.05 | 0 | 0 | 0.99 | 0.95 |
| 1994 | 7 | 0 | 0 | 0.00 | 0.00 | 0 | 0 | 0.67 | 1.00 |
| 1995 | 5 | 0 | 0 | 0.00 | 0.00 | 0 | 0 | 1.00 | 1.00 |
| 1996 | 30 | 0 | 0 | 0.00 | 0.00 | 0 | 0 | 0.60 | 1.00 |
| 1997 | 33 | 0 | 0 | 0.00 | 0.00 | 0 | 0 | 0.30 | 1.00 |
| 1998 | 11 | 0 | 0 | 0.00 | 0.00 | 0 | 0 | 0.44 | 1.00 |


| Brood Year | Spawners |  |  |  |  | Broodstock |  |  | PNI |
| :---: | :---: | :---: | :---: | :---: | :---: | :---: | :---: | :---: | :---: |
|  | NOS | $\mathrm{HOS}_{\mathrm{w}}$ | HOSs | pHOS ${ }_{\text {w }}$ | pHOSs | $\mathrm{NOB}_{\mathrm{N}}$ | $\mathrm{HOB}_{\mathrm{N}}$ | pNOB |  |
| 1999 | 3 | 0 | 0 | 0.00 | 0.00 | 0 | 0 | 1.00 | 1.00 |
| 2000 | 22 | 0 | 0 | 0.00 | 0.00 | 0 | 0 | 0.48 | 1.00 |
| Average* | 48 | 0 | 1 | 0.00 | 0.00 | 0 | 0 | 0.79 | 1.00 |
| Median* | 32 | 0 | 0 | 0.00 | 0.00 | 0 | 0 | 1.00 | 1.00 |
| 2001 | 111 | 0 | 55 | 0.00 | 0.33 | 5 | 0 | 1.00 | 0.50 |
| 2002 | 60 | 0 | 26 | 0.00 | 0.30 | 18 | 0 | 1.00 | 0.51 |
| 2003 | 31 | 0 | 5 | 0.00 | 0.14 | 7 | 0 | 1.00 | 0.77 |
| 2004 | 54 | 0 | 12 | 0.00 | 0.18 | 6 | 0 | 1.00 | 0.70 |
| 2005 | 38 | 11 | 106 | 0.07 | 0.68 | 103 | 73 | 0.59 | 0.33 |
| 2006 | 41 | 5 | 9 | 0.09 | 0.16 | 191 | 135 | 0.59 | 0.61 |
| 2007 | 62 | 23 | 7 | 0.25 | 0.08 | 254 | 6 | 0.98 | 0.67 |
| 2008 | 20 | 2 | 30 | 0.04 | 0.58 | 116 | 0 | 1.00 | 0.34 |
| 2009 | 81 | 29 | 63 | 0.17 | 0.36 | 238 | 0 | 1.00 | 0.53 |
| 2010 | 27 | 22 | 23 | 0.31 | 0.32 | 90 | 0 | 1.00 | 0.50 |
| 2011 | 83 | 0 | 0 | 0.00 | 0.00 | 306 | 0 | 1.00 | 1.00 |
| 2012 | 89 | 10 | 45 | 0.07 | 0.31 | 390 | 0 | 1.00 | 0.73 |
| 2013 | 44 | 55 | 5 | 0.53 | 0.05 | 383 | 0 | 1.00 | 0.64 |
| Average** | 57 | 12 | 30 | 0.12 | 0.27 | 162 | 16 | 0.94 | 0.60 |
| Median** | 54 | 5 | 23 | 0.07 | 0.30 | 116 | 0 | 1.00 | 0.61 |

HOSw = hatchery-origin spawners in White River from the White River spring Chinook Supplementation Program.
$\mathbf{p H O S} \mathbf{w}=$ proportion of hatchery-origin spawners from White River spring Chinook Supplementation Program.
HOSs $_{\mathbf{s}}=$ stray hatchery-origin spawners in the White River.
$\mathbf{p H O S}_{\mathbf{s}}=$ proportion of stray hatchery-origin spawners.
NOBw = natural origin broodstock spawned for the White River spring Chinook Supplementation Program.
$\mathbf{H O B}_{\mathbf{w}}$ = hatchery-origin broodstock spawned in the White River spring Chinook Supplementation Program.
$\mathbf{p N O B}=$ proportion of hatchery-origin broodstock. Because of the high incidence of strays to the White River from the Chiwawa River spring Chinook program, pNOB values from the Chiwawa program were used to estimate PNI values during the period from 1989 to 2000 (italicized). The weighting for those years was $100 \%$ based on the Chiwawa program broodstock selection, because there have been no hatchery returns from the White River spring Chinook program during this period.
PNI = Proportionate Natural Influence for White River spring Chinook calculated using the gene-flow model for multiple programs.

* Average and median for the period 1989-2000.
** Average and median for the period 2001-2013.


## Wenatchee Subbasin Spring Chinook Salmon

For brood years 1989-2018, PNI values ranged from 0.34 to 1.00 in the portion of the Wenatchee Subbasin upstream of Tumwater Dam (Table 6; see the sections on Chiwawa River, Nason Creek, and White River for details as to how program changes affected PNI). PNI values for brood years 1989-1994, 2014, and 2016 were greater than or equal to 0.67 . However, there were no hatchery origin spawners in brood years 1989-1992 (pre-dated the first hatchery program returns).

Table 6. PNI values for the Wenatchee Subbasin (upstream of Tumwater Dam) spring Chinook supplementation programs for brood years 1989-2018.

| Brood Year | Spawners |  |  | Broodstock |  |  | PNI ${ }^{\text {a }}$ |
| :---: | :---: | :---: | :---: | :---: | :---: | :---: | :---: |
|  | NOS | HOS | pHOS | NOB | HOB | pNOB |  |
| 1989 | 1461 | 0 | 0.00 | 28 | 0 | 1.00 | 1.00 |
| 1990 | 1003 | 0 | 0.00 | 18 | 0 | 1.00 | 1.00 |
| 1991 | 585 | 0 | 0.00 | 27 | 0 | 1.00 | 1.00 |
| 1992 | 1098 | 0 | 0.00 | 78 | 0 | 1.00 | 1.00 |
| 1993 | 935 | 257 | 0.22 | 94 | 0 | 1.00 | 0.82 |
| 1994 | 214 | 66 | 0.24 | 8 | 4 | 0.67 | 0.74 |
| 1995 | 23 | 36 | 0.61 | 0 | 0 | NA | NA |
| 1996 | 139 | 43 | 0.24 | 8 | 10 | 0.44 | 0.66 |
| 1997 | 211 | 177 | 0.46 | 32 | 79 | 0.29 | 0.41 |
| 1998 | 149 | 35 | 0.19 | 13 | 34 | 0.28 | 0.62 |
| 1999 | 123 | 10 | 0.08 | 0 | 0 | NA | NA |
| 2000 | 484 | 278 | 0.36 | 9 | 21 | 0.30 | 0.48 |
| 2001 | 979 | 1977 | 0.67 | 118 | 259 | 0.31 | 0.34 |
| 2002 | 732 | 863 | 0.54 | 38 | 51 | 0.43 | 0.46 |
| 2003 | 383 | 212 | 0.36 | 48 | 53 | 0.48 | 0.59 |
| 2004 | 1010 | 591 | 0.37 | 89 | 132 | 0.40 | 0.54 |
| 2005 | 304 | 1169 | 0.79 | 194 | 254 | 0.43 | 0.37 |
| 2006 | 304 | 636 | 0.68 | 282 | 359 | 0.44 | 0.41 |
| 2007 | 388 | 1619 | 0.81 | 297 | 110 | 0.73 | 0.48 |
| 2008 | 362 | 1778 | 0.83 | 199 | 220 | 0.47 | 0.38 |
| 2009 | 585 | 1610 | 0.73 | 334 | 111 | 0.75 | 0.52 |
| 2010 | 539 | 1220 | 0.69 | 167 | 98 | 0.63 | 0.49 |
| 2011 | 1213 | 1778 | 0.59 | 386 | 93 | 0.81 | 0.59 |
| 2012 | 931 | 1580 | 0.63 | 456 | 45 | 0.91 | 0.60 |
| 2013 | 614 | 1408 | 0.70 | 471 | 7 | 0.99 | 0.59 |
| 2014 | 779 | 575 | 0.42 | 79 | 12 | 0.87 | 0.68 |
| 2015 | 616 | 775 | 0.56 | 124 | 63 | 0.66 | 0.55 |
| 2016 | 615 | 226 | 0.27 | 127 | 108 | 0.54 | 0.68 |
| 2017 | 259 | 369 | 0.59 | 120 | 82 | 0.59 | 0.51 |
| 2018 | 221 | 661 | 0.75 | 83 | 111 | 0.43 | 0.38 |
| Average | 575 | 665 | 0.45 | 131 | 77 | 0.64 | 0.60 |
| Median | 562 | 472 | 0.50 | 86 | 52 | 0.61 | 0.57 |

${ }^{\text {apNI }}$ was calculated previously using PNI approximate equation 11 (HSRG 2009; their Appendix A). All PNI values presented here were recalculated by iterating Ford's (2002) equations 5 and 6 to equilibrium using a heritability of 0.3 and a selection strength of three standard deviations. C. Busack, NOAA Fisheries, 21 March 2016, provided the model for calculating PNI.

## Methow River Spring Chinook Salmon

For brood years 2003-2018, PNI has consistently been below 0.67 and ranged from 0.06 to 0.50 (Table 7). During this time, the mean proportion of natural-origin spring Chinook Salmon on spawning grounds was higher in the Twisp River than in the Methow or Chewuch rivers. However, mean Methow River Subbasin PNI values are low and indicate that most genetic selection pressure on progeny produced from naturally spawning adults comes from the hatchery environment (Table 7).

Table 7. PNI calculated for specific broods of spawning spring Chinook Salmon in the Methow River Subbasin. PNI was calculated using a three-population model incorporating the proportion of hatchery fish from conservation programs (HC; e.g., Methow Hatchery), safety-net programs (HSN; e.g., Winthrop National Fish Hatchery), and natural-origin (Wild) fish on the spawning grounds within each tributary and spawning year. Stray hatchery-origin fish were included in the HC or HSN categories based on the known or assumed parentage of broodstock.

| $\begin{array}{\|c\|} \hline \text { Brood } \\ \text { Year } \end{array}$ | Chewuch |  |  | Methow |  |  | Twisp |  |  | Methow Subbasin |  |  |  |
| :---: | :---: | :---: | :---: | :---: | :---: | :---: | :---: | :---: | :---: | :---: | :---: | :---: | :---: |
|  | HC | HSN | Wild | HC | HSN | Wild | HC | HSN | Wild | HC | HSN | Wild | PNI |
| 2003 | 0.92 | 0.03 | 0.05 | 0.65 | 0.34 | 0.01 | 0.42 | 0.00 | 0.58 | 0.76 | 0.19 | 0.05 | 0.2 |
| 2004 | 0.83 | 0.03 | 0.14 | 0.56 | 0.20 | 0.24 | 0.23 | 0.06 | 0.71 | 0.54 | 0.13 | 0.33 | 0.21 |
| 2005 | 0.52 | 0.05 | 0.43 | 0.56 | 0.14 | 0.30 | 0.28 | 0.00 | 0.72 | 0.52 | 0.09 | 0.38 | 0.50 |
| 2006 | 0.54 | 0.20 | 0.26 | 0.61 | 0.27 | 0.12 | 0.60 | 0.01 | 0.39 | 0.59 | 0.23 | 0.19 | 0.06 |
| 2007 | 0.42 | 0.31 | 0.27 | 0.28 | 0.50 | 0.22 | 0.62 | 0.00 | 0.38 | 0.35 | 0.41 | 0.25 | 0.30 |
| 2008 | 0.37 | 0.29 | 0.34 | 0.30 | 0.41 | 0.29 | 0.76 | 0.00 | 0.24 | 0.39 | 0.31 | 0.30 | 0.2 |
| 2009 | 0.46 | 0.19 | 0.35 | 0.43 | 0.42 | 0.15 | 0.67 | 0.08 | 0.25 | 0.45 | 0.34 | 0.21 | 0.20 |
| 2010 | 0.51 | 0.18 | 0.31 | 0.50 | 0.32 | 0.18 | 0.38 | 0.00 | 0.62 | 0.49 | 0.25 | 0.25 | 0.0 |
| 2011 | 0.42 | 0.16 | 0.42 | 0.50 | 0.26 | 0.24 | 0.16 | 0.18 | 0.66 | 0.45 | 0.22 | 0.33 | 0.19 |
| 2012 | 0.67 | 0.05 | 0.28 | 0.76 | 0.10 | 0.14 | 0.67 | 0.01 | 0.32 | 0.72 | 0.07 | 0.20 | 0.2 |
| 2013 | 0.67 | 0.05 | 0.28 | 0.76 | 0.06 | 0.18 | 0.75 | 0.00 | 0.25 | 0.73 | 0.05 | 0.22 | 0.40 |
| 2014 | 0.53 | 0.09 | 0.38 | 0.63 | 0.19 | 0.18 | 0.62 | 0.01 | 0.37 | 0.60 | 0.15 | 0.25 | 0.40 |
| 2015 | 0.41 | 0.12 | 0.47 | 0.59 | 0.24 | 0.17 | 0.31 | 0.02 | 0.67 | 0.52 | 0.19 | 0.29 | 0.39 |
| 2016 | 0.19 | 0.19 | 0.62 | 0.28 | 0.36 | 0.36 | 0.33 | 0.00 | 0.67 | 0.24 | 0.30 | 0.46 | 0.29 |
| 2017 | 0.38 | 0.19 | 0.43 | 0.32 | 0.34 | 0.34 | 0.40 | 0.13 | 0.47 | 0.35 | 0.27 | 0.38 | 0.38 |
| 2018 | 0.33 | 0.20 | 0.47 | 0.16 | 0.34 | 0.50 | 0.25 | 0.06 | 0.69 | 0.22 | 0.25 | 0.53 | 0.44 |
| Avg. | 0.51 | 0.15 | 0.34 | 0.49 | 0.28 | 0.23 | 0.47 | 0.04 | 0.50 | 0.50 | 0.22 | 0.29 | 0.29 |
| Median | 0.49 | 0.17 | 0.35 | 0.53 | 0.30 | 0.20 | 0.41 | 0.01 | 0.53 | 0.51 | 0.23 | 0.27 | 0.29 |

## Wenatchee River Summer Chinook Salmon

For brood years 1989-2018, the PNI value has been greater than 0.67 (Table 8). For those brood years, PNI ranged from 0.68 to 1.00 and averaged 0.87 .

Table 8. PNI values for the Wenatchee summer Chinook supplementation program for brood years 1989-2018.

| Brood year | Spawners |  |  | Broodstock |  |  | PNI |
| :---: | :---: | :---: | :---: | :---: | :---: | :---: | :---: |
|  | NOS | HOS | pHOS | NOB | HOB | pNOB |  |
| 1989 | 14,331 | 0 | 0.00 | 290 | 0 | 1.00 | 1.00 |
| 1990 | 10,861 | 0 | 0.00 | 57 | 0 | 1.00 | 1.00 |
| 1991 | 10,168 | 0 | 0.00 | 105 | 0 | 1.00 | 1.00 |
| 1992 | 11,652 | 0 | 0.00 | 274 | 0 | 1.00 | 1.00 |
| 1993 | 8,842 | 608 | 0.06 | 406 | 44 | 0.90 | 0.94 |
| 1994 | 8,476 | 1,678 | 0.17 | 333 | 54 | 0.86 | 0.84 |
| 1995 | 6,854 | 901 | 0.12 | 363 | 16 | 0.96 | 0.89 |
| 1996 | 6,000 | 168 | 0.03 | 263 | 3 | 0.99 | 0.97 |
| 1997 | 5,408 | 505 | 0.09 | 205 | 13 | 0.94 | 0.92 |
| 1998 | 4,707 | 645 | 0.12 | 299 | 78 | 0.79 | 0.87 |
| 1999 | 3,997 | 1,342 | 0.25 | 242 | 236 | 0.51 | 0.68 |
| 2000 | 4,466 | 1,046 | 0.19 | 275 | 180 | 0.60 | 0.77 |
| 2001 | 8,356 | 1,691 | 0.17 | 210 | 136 | 0.61 | 0.79 |
| 2002 | 11,846 | 3,740 | 0.24 | 409 | 10 | 0.98 | 0.81 |
| 2003 | 10,064 | 1,736 | 0.15 | 337 | 7 | 0.98 | 0.87 |
| 2004 | 8,044 | 1,070 | 0.12 | 424 | 2 | 1.00 | 0.90 |
| 2005 | 6,869 | 1,834 | 0.21 | 397 | 3 | 0.99 | 0.83 |
| 2006 | 15,405 | 3,188 | 0.17 | 432 | 4 | 0.99 | 0.86 |
| 2007 | 2,764 | 1,806 | 0.40 | 263 | 3 | 0.99 | 0.72 |
| 2008 | 8,061 | 3,699 | 0.31 | 376 | 71 | 0.84 | 0.74 |
| 2009 | 7,754 | 1,330 | 0.15 | 449 | 8 | 0.98 | 0.86 |
| 2010 | 6,253 | 1,671 | 0.21 | 388 | 5 | 0.99 | 0.83 |
| 2011 | 8,107 | 1,681 | 0.17 | 375 | 7 | 0.98 | 0.86 |
| 2012 | 6,963 | 1,150 | 0.14 | 267 | 1 | 1.00 | 0.88 |
| 2013 | 6,798 | 2,412 | 0.26 | 234 | 2 | 0.99 | 0.80 |
| 2014 | 9,901 | 772 | 0.07 | 261 | 2 | 0.99 | 0.94 |
| 2015 | 4,033 | 240 | 0.06 | 248 | 0 | 1.00 | 0.95 |
| 2016 | 5,700 | 509 | 0.08 | 259 | 0 | 1.00 | 0.93 |
| 2017 | 7,620 | 906 | 0.11 | 252 | 1 | 1.00 | 0.90 |
| 2018 | 2,606 | 656 | 0.20 | 205 | 5 | 0.98 | 0.83 |
| Average | 7,764 | 1,233 | 0.14 | 297 | 30 | 0.93 | 0.87 |
| Median | 7,687 | 1,058 | 0.15 | 275 | 5 | 0.99 | 0.87 |

## Methow River Summer Chinook Salmon

For brood years 1993-2003, the PNI values were generally less than 0.67 (Table 9). Since brood year 2003, PNI has generally been equal to or greater than 0.67 . For the entire time series, PNI has ranged from 0.32 to 1.00 and averaged 0.71 .

Table 9. PNI values for the Methow summer Chinook supplementation program for brood years 19892018.

| Brood <br> year | Spawners |  |  | Broodstock |  |  | PNI |
| :---: | :---: | :---: | :---: | :---: | :---: | :---: | :---: |
|  | 492 | 0 | 0.00 | 1,297 | 312 | 0.81 |  |
| 1990 | 1,421 | 0 | 0.00 | 828 | 206 | 0.80 | 1.00 |
| 1991 | 566 | 0 | 0.00 | 924 | 314 | 0.75 | 1.00 |
| 1992 | 460 | 0 | 0.00 | 297 | 406 | 0.42 | 1.00 |
| 1993 | 314 | 194 | 0.38 | 681 | 388 | 0.64 | 0.64 |
| 1994 | 596 | 489 | 0.45 | 341 | 244 | 0.58 | 0.58 |
| 1995 | 596 | 618 | 0.51 | 173 | 240 | 0.42 | 0.47 |
| 1996 | 435 | 180 | 0.29 | 290 | 223 | 0.57 | 0.67 |
| 1997 | 529 | 168 | 0.24 | 198 | 264 | 0.43 | 0.71 |
| 1998 | 435 | 240 | 0.36 | 153 | 211 | 0.42 | 0.56 |
| 1999 | 570 | 415 | 0.42 | 224 | 289 | 0.44 | 0.53 |
| 2000 | 862 | 338 | 0.28 | 164 | 339 | 0.33 | 0.56 |
| 2001 | 1,108 | 1,659 | 0.60 | 91 | 266 | 0.25 | 0.32 |
| 2002 | 2,591 | 2,039 | 0.44 | 247 | 241 | 0.51 | 0.55 |
| 2003 | 2,318 | 1,612 | 0.41 | 381 | 101 | 0.79 | 0.67 |
| 2004 | 1,641 | 548 | 0.25 | 506 | 16 | 0.97 | 0.80 |
| 2005 | 1,672 | 889 | 0.35 | 391 | 9 | 0.98 | 0.74 |
| 2006 | 1,685 | 1,048 | 0.38 | 500 | 10 | 0.98 | 0.73 |
| 2007 | 656 | 708 | 0.52 | 456 | 17 | 0.96 | 0.66 |
| 2008 | 1,197 | 750 | 0.39 | 404 | 41 | 0.91 | 0.71 |
| 2009 | 1,044 | 714 | 0.41 | 507 | 0 | 1.00 | 0.72 |
| 2010 | 1,325 | 1,168 | 0.47 | 484 | 8 | 0.98 | 0.68 |
| 2011 | 1,487 | 1,430 | 0.49 | 467 | 26 | 0.95 | 0.67 |
| 2012 | 1,596 | 1,351 | 0.46 | 98 | 1 | 0.99 | 0.69 |
| 2013 | 1,707 | 1,876 | 0.52 | 97 | 4 | 0.96 | 0.66 |
| 2014 | 1,450 | 175 | 0.11 | 96 | 0 | 1.00 | 0.90 |
| 2015 | 3,142 | 809 | 0.20 | 97 | 1 | 0.99 | 0.84 |
| 2216 | 1,466 | 775 | 0.35 | 103 | 0 | 1.00 | 0.75 |
| 2017 | 1,039 | 369 | 0.26 | 111 | 0 | 1.00 | 0.80 |
| 2018 | 675 | 692 | 0.51 | 130 | 1 | 0.99 | 0.67 |
| Average | $\mathbf{1 , 1 6 9}$ | 708 | $\mathbf{0 . 3 4}$ | 358 | $\mathbf{1 3 9}$ | $\boldsymbol{0} 9.76$ | $\mathbf{0 . 7 1}$ |
| Median | $\mathbf{1 , 0 7 6}$ | $\mathbf{6 5 5}$ | $\mathbf{0 . 3 8}$ | 294 | 71 | $\mathbf{0 . 8 6}$ | $\mathbf{0 . 6 9}$ |

## Discussion

Target PNI values in the upper Columbia Basin were generally not met for species with low population size such as endangered spring Chinook Salmon and threatened steelhead, but were exceeded for populations of summer Chinook Salmon that were relatively large. Recently, PNI targets were exceeded for fall Chinook Salmon in the Hanford Reach, another large upper Columbia population (Pearsons et al. 2020). When hatchery production was large relative to the natural spawning population, then PNI targets were difficult to achieve without significant management of returning adults to control pHOS. In some areas, such as upstream of Tumwater Dam in the Wenatchee Subbasin, the management of hatchery-origin fish was possible. In other areas, such as the Methow Subbasin, the management of hatchery-origin adults was more challenging because it relied upon removal at hatchery traps and recreational angling. It was also difficult to achieve high pNOB when populations were small. Federal Section 10 permits require that the proportion of natural-origin fish that can be collected for hatchery broodstock be less than $33 \%$ of the run. When run sizes are small, then the pNOB will be low because a greater proportion of hatchery-origin fish will have to be used for broodstock to meet hatchery production goals. In short, operating large integrated hatchery programs relative to the naturalorigin spawning population creates difficult trade-offs and challenges.

Reliance upon adult management to achieve pHOS goals poses risk of mining naturalorigin fish from the spawning population, which could pose greater risk to the population than domestication. If hatchery-origin fish were killed to manage pHOS , and natural-origin fish were used as broodstock to produce those hatchery-origin fish, then the natural-origin fish were mined to support the hatchery program. This poses a demographic risk to the population and also removes parental natural-origin fish from contributing to natural production. The risks of mining the natural-origin population could be evaluated relative to the risks of domestication and the size of hatchery programs to determine what is most optimal to achieve program goals.

Straying of non-target hatchery fish is another factor that contributed to lowering PNI of small populations. In some cases, such as in Nason Creek and the White River, strays from other hatcheries influenced PNI more than target hatcheries (Pearsons and Miller, see chapter in this report). Stray hatchery-origin fish increase domestication risk but also pose risk of decreasing between-population genetic variation. Larger populations such as summer Chinook Salmon were able to absorb strays without large increases in PNI that occurred in smaller populations.

Managing to achieve conservation PNI targets could follow guidelines whereby pNOB should be larger than pHOS , but pNOB should not be increased in order to achieve the PNI target. Rather, pHOS should be decreased to the point where the PNI target can be achieved, and should be less than $30 \%$ (HSRG 2009; [typically the ESA permitted pHOS upper limit is $25 \%$ in conservation programs]). Increasing pNOB above $50 \%$ offers minor genetic benefit, while reducing pHOS allows lower pNOB (HSRG 2009). However, controlling pHOS may be difficult or impossible in systems where the opportunity for origin-selective adult removal is limited. Such cases necessitate the re-evaluation of management objectives and program structure to provide a broader suite of approaches for constraining the number of hatchery-origin fish on the spawning ground to a suitable number for meeting the PNI objective. In cases where a population is at risk of functional extirpation, the PNI guidelines described by the HSRG
(2009) may be altered to encourage more fish spawning in the wild. In these cases, clearly defined guidelines could be established and the population carefully monitored (HSRG 2009). In general, pNOB must exceed pHOS (HSRG 2009); however, when pHOS is high, increasing pNOB to compensate risks of mining the natural population of spawners, exacerbating pHOS while continuing to produce more hatchery origin fish than are warranted under a PNI management regime. Management targets have been established for PNI for the various programs listed under the Endangered Species Act. Steelhead programs in the Wenatchee and Methow had PNI averages or medians below 0.67. Assessment of mean PNI can be complicated by management for escapement, as opposed to PNI, when run sizes are small, as happened in the Wenatchee in 2017. The new steelhead management regime in the Methow did not initiate until 2017. Therefore, data in this report do not reflect the current or future management of the Methow River Subbasin for gene flow.

The results of this analyses indicate that the Spring Chinook and steelhead programs would benefit from adjustment to meet the PNI targets. In some cases, the PNI values may improve as the reduction in hatchery release numbers and new management strategies mature. Evaluation of the interaction of adult management, hatchery program size, and stray management could help to achieve PNI targets. Furthermore, assessment of genetic and demographic risks could be evaluated to determine trade-offs between how these factors contribute to long-term fitness. The Methow steelhead and particularly spring Chinook may require substantial management changes in order to achieve the gene flow objectives.

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# Adult Migration Timing, Spawn Timing, and Spawning Distribution of Spring Chinook Salmon and Summer Chinook Salmon in the Wenatchee and Methow Basins 

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[^4]
#### Abstract

The migration timing, spawn timing, and spatial spawning distribution of hatchery- and naturalorigin salmon in the natural environment can be important metrics in the evaluation of integrated hatchery programs. The timing of migration and spawning of hatchery- and natural-origin spring Chinook Salmon were generally similar in the Wenatchee and Methow sub-basins. Although the difference in arrival timing was small and not statistically significant, the visual observation and PIT-tag data at Tumwater and Wells dams suggests a tendency for hatchery-origin spring Chinook Salmon to arrive later than natural-origin fish. Differences in migration timing between spring Chinook Salmon populations in the Wenatchee versus Methow sub-basins were observed and may result in differences in survival at the adult life-stage. With summer Chinook Salmon, hatchery-origin fish in the Wenatchee sub-basin consistently passed Dryden Dam later than their natural-origin counterparts whereas the migration timing for Methow and Okanogan sub-basin hatchery-origin fish was a near match with natural-origin fish at Wells Dam. Summer Chinook Salmon spawn timing in both Wenatchee and Methow hatchery-origin fish was later than natural-origin fish, with Methow-origin fish having a larger average difference. For spatial distribution of spawning spring Chinook Salmon, differences between hatchery- and naturalorigin spawner distribution across the historical survey reaches were observed in all programs except the White River program. In general, hatchery-origin females spawned lower in the watershed. However, the proportion of hatchery-origin fish was high (i.e., greater than $50 \%$ ) in the majority of survey reaches ( 33 out of 51 ), particularly in those reaches that were the primary spawning areas by natural-origin fish. The proportion of hatchery-origin spring Chinook Salmon exceeded $30 \%$ in 48 of the 51 survey reaches. The distributions of spawning Wenatchee and Methow hatchery-origin summer Chinook Salmon also differed from the spawner distributions of the natural-origin populations, with hatchery-origin females more often spawning lower in the watershed. This difference in spawning distribution was consistent with management objectives. The proportion of hatchery-origin summer Chinook Salmon was greater than $30 \%$ in 6 of the 17 survey reaches.


## Introduction

Hatchery programs, through hatchery practices that may induce physiological or behavioral consequences such as unintentional straying, can reduce life-history variability resulting in the reduction of the portfolio effect of multiple aggregate populations (Schindler et al. 2010; Carlson and Satterthwaite 2011). For example, the release timing of juvenile hatcheryorigin fish is typically highly condensed in space and time relative to the natural-origin juvenile migration patterns. The condensed release of juvenile hatchery fish, when timed with preferred environmental conditions (e.g., high flows, darkness, low visibility), may increase juvenile survival. However, this artificial narrowing of the emigration temporal window may also reduce the buffering effect of a more widely distributed migration pattern (Kovach et al. 2013). The diversity and variability observed in populations of natural-origin fish is made possible by a combination of genetic, phenotypic, and habitat heterogeneity, all of which can be reduced by both the loss or simplification of habitat and the introduction of hatchery-reared fish. For example, the diversity of Chinook Salmon in the Central Valley of California, the only location where four distinct runs of Chinook Salmon occurred, was reduced first by construction of impassable dams and then by the temporally condensed release of genetically homogeneous hatchery-origin fish in space and time as mitigation for those losses (Carlson and Satterthwaite 2011; Huber and Carlson 2015).

The spatial spawning distribution of hatchery- and natural-origin salmon in the natural environment can be an important metric in the evaluation of integrated hatchery programs (Knudsen et al. 2006; Hoffnagle et al. 2008; Murdoch et al. 2009; Hughes and Murdoch 2017; Hillman et al. 2019). Integrated hatchery programs incorporate naturally produced fish into the hatchery broodstock and hatchery-origin fish are intended to spawn on the spawning grounds with natural-origin fish, with the goal of increasing spawner abundance (Mobrand et al. 2005) and the assumption that this will result in a larger population overall. Spatial similarity in the spawning distributions of hatchery- and naturally produced fish on the spawning grounds indicates effective integration of the spawning population unless assortative mating occurs. Differences in the spatial distribution of spawning hatchery- and naturally produced salmon may simply be an artifact of the low abundance of natural- or hatchery-produced fish (i.e., vacant habitat), sub-optimal smolt release locations (Murdoch et al. 2007; Hoffnagle et al. 2008), or inadequate imprinting and homing of hatchery-origin fish (Murdoch et al. 2007). Alternatively, it can be the result of an intentional management objective where differences in spatial and temporal spawning distribution are desirable (Mackey et al. 2001). Furthermore, strays from other hatchery programs can influence the distribution of hatchery-origin spawners in non-target streams. The Hatchery Scientific Review Group (HSRG) recommends that the proportion of hatchery-origin spawners ( pHOS ) should not exceed 0.30 for integrated hatchery programs in order to reduce the risk of domestication selection (Mobrand et al. 2005; Paquet et al. 2011).

The spawning distribution objective for the spring Chinook Salmon programs evaluated in these analyses are for hatchery-origin and natural-origin fish to spawn in similar locations, in contrast to the summer Chinook Salmon hatchery programs in which replication of the spawning distribution between hatchery-origin and natural-origin fish is not the objective. The upper range of summer Chinook Salmon spawning distribution has the potential to overlap with ESA-listed spring Chinook Salmon, which may pose an unknown and potentially adverse impact to spring Chinook Salmon; therefore, summer Chinook Salmon acclimation sites are located lower in the sub-basin in an effort to minimize this overlap in the upper range of the spawning distribution of
summer Chinook Salmon. Exact replication of the spawning distribution for summer Chinook Salmon is not the management expectation (Hillman et al. 2019).

Using a combination of observational data, we explored qualitative and quantitative differences in migration timing, spawn timing, and spawning distribution between hatchery- and natural-origin spring and summer Chinook Salmon within the Wenatchee Sub-basin and Methow Sub-basin (WA). From these data, we assessed whether the respective hatchery programs are achieving the goal of similar migration and spawn timing of adult hatchery- and natural-origin fish, and whether there is evidence of a temporal shift of migration or spawning timing in either hatchery- or natural-origin fish over time. Additionally, we assessed whether the spatial distribution of redds is similar for hatchery-origin Chinook Salmon compared to natural-origin Chinook Salmon.

## Methods

## Adult Run Timing

Using PIT-tag recapture data from the PTAGIS database (Pacific States Marine Fisheries Commission, https://www.ptagis.org/), we compared the arrival timing of hatchery- and naturalorigin Upper Columbia spring Chinook Salmon populations to various locations within the Columbia Basin. The comparisons included age 3, 4, and 5 fish from return years 2005-2018. The run timing of these fish was evaluated by origin, age, and release location as identified in PTAGIS.

Using data reported from the Public Utility Districts' (PUD) hatchery monitoring and evaluation reports (Hillman et al. 2020; Snow et al. 2020), we evaluated mean arrival timing of hatchery- and natural-origin spring Chinook Salmon from 1998-2018 by calculating mean day of the year (DOY) (days since January 1) that $10 \%, 50 \%$, and $90 \%$ of the fish passed Tumwater Dam (Wenatchee populations) and Wells Dam (Methow populations). All age groups were pooled together in the analysis, because these data were based on video monitoring and trapping. The origin of adults (hatchery or natural) was determined by adipose fin presence or absence or the presence of coded-wire tags. Migration timing at Tumwater Dam was based on video sampling; migration timing at Wells Dam was based on stock-assessment and broodstock trapping in the Wells Dam fishways. Data for 1998 through 2003 were based on video and broodstock trapping and may not reflect the actual number of hatchery-origin spring Chinook Salmon that passed the dams. Most spring Chinook Salmon were visually examined during trapping from 2004 to 2018; however, enumeration errors may still exist because of misidentified run-type assignment (i.e., spring or summer). The mean differences in mean arrival timing were compared with paired t-tests on the $10 \%, 50 \%$ (median), and $90 \%$ DOY.

Due to the limited number of natural-origin PIT-tagged summer Chinook Salmon, evaluations of adult run timing for summer Chinook Salmon were limited to visual observations at dam fishways or broodstock collection locations. As with spring Chinook Salmon, we compared the mean migration timing of hatchery- and natural-origin summer Chinook Salmon by comparing the mean DOY on which $10 \%, 50 \%$, and $90 \%$ of the fish passed Dryden Dam and Wells Dam. These data were based on stock assessment and broodstock collection at Dryden Dam (Wenatchee population) and Wells Dam during the migration period 2007-2018. The natural populations observed at Wells Dam consist of fish produced in the Methow and Okanogan sub-basins, plus potentially in the Columbia River. The hatchery populations consist
of fish released in the Methow, Okanogan, and Columbia rivers, plus potentially strays from downstream locations.

## Spawn Timing

Differences in spawn timing between hatchery- and natural-origin spring Chinook Salmon and summer Chinook Salmon were evaluated by comparing the temporal distributions of female carcasses recovered on the spawning grounds. These data were assessed with all years pooled and year-by-year by sub-basin or major spawning tributary. The mean differences in spawn timing between hatchery- and natural-origin fish were evaluated with paired t-tests on the 10th, 50th, and 90th percentile carcass-recovery date. Additionally, we evaluated the relationship between the DOY and elevation of carcass recoveries. Carcass recovery is an indirect measurement of spawn timing and dependent on several other independent variables that may affect an evaluation of timing. Therefore, the results of differences or similarities in timing should be evaluated in the context of carcass recovery date as a proxy for spawn timing and the relationship between those two metrics.

## Spawning Distribution

Differences in the distribution of hatchery- and natural-origin Chinook Salmon spawners were assessed by examining the location (RKm) where female carcasses were observed in spawning streams. The focus was on female carcasses because they are a better indicator of spawning location than are male carcasses (Murdoch et al. 2009). During weekly spawning ground surveys (described in Hillman et al. 2019), crews recorded the location of female carcasses (recorded to the nearest 0.1 RKm ) as measured from the confluence ( $\mathrm{RKm} \mathrm{0.0} \mathrm{)} \mathrm{of} \mathrm{each}$ stream. The entire spawning distribution of Chinook Salmon was inventoried (Hillman et al. 2019).

We evaluated differences in spawning locations at two different spatial scales: at the historical reach scale and at the $0.5-\mathrm{km}$ (in tributaries) or $1.0-\mathrm{km}$ (in mainstem rivers) scale. Historical reaches are the stream reaches defined by individuals conducting redd surveys over the history of the respective monitoring programs and are generally related to river accessibility and the length of stream that one can reasonably survey in given amount of time. At the historical reach scale, we evaluated the proportion of natural- and hatchery-origin spawners among all the reaches and within each reach. That is, we calculated the fraction of all naturaland hatchery-origin spawners within a stream that were observed within each reach of that stream. We also calculated pHOS and pNOS (proportion of natural-origin spawners) within each reach by dividing the number of hatchery-origin or natural-origin spawners within a reach by the total number of hatchery- and natural-origin spawners within that reach. We then constructed frequency-distribution plots to compare the spawning distribution of hatchery- and natural-origin fish within and among reaches. We performed two-way Yates' Chi-square tests on count data to determine whether there were statistical differences in spawning distributions between hatcheryand natural-origin fish at the historical-reach scale. We were unable to evaluate differences in distribution of hatchery- and natural-origin fish "before" and "after" a reduction in Chinook Salmon hatchery production. Based on the year when the last age- 5 fish return from the "before" period would provide at most only two years of "after" data. Two years of "after" data are too few to make reasonable comparisons.

## Results

## Migration Timing of Wenatchee and Methow Spring Chinook Salmon

Adult Upper Columbia River spring Chinook Salmon typically began arriving at Bonneville Dam by April and their migration continued through July (Figure 1). Generally, fish from the Leavenworth National Fish Hatchery and the Methow River sub-basin were the earliest to arrive. From 2005-2018, the median arrival date to Bonneville Dam of natural-origin fish from the Methow Sub-basin was approximately 15 days earlier than natural-origin fish from the Chiwawa River (Figure 1). The arrival timing of hatchery-origin fish to Bonneville Dam followed the same pattern as natural-origin fish; fish from the Methow Sub-basin arrived earlier than Wenatchee Sub-basin fish and the median passage date of fish from the Methow Hatchery was approximately 20 days earlier than fish released from the Chiwawa Acclimation Facility.

In the Wenatchee Sub-basin, the arrival timing at Tumwater Dam of PIT-tagged hatchery- and natural-origin spring Chinook Salmon was nearly identical with some variability due to small sample sizes (Figure 2). The mean arrival DOY of natural-origin fish was 182 versus 184 for hatchery-origin fish (median $=183$ vs 185). Similarly, at Wells Dam, PIT-tagged natural-origin and hatchery-origin spring Chinook Salmon from the Methow Sub-basin arrived at similar times (average $=151$ for natural-origin, versus 150 for hatchery-origin, median $=148 \mathrm{vs}$ 149 , respectively; Figure 3). Both the Methow and Wenatchee sub-basins were consistent in the pattern of return date by age where age- 3 fish returned later than age- 4 and age- 5 fish (Figure 4).

The observed patterns described above were consistent with cumulative frequency plots of migration timing of PIT-tagged hatchery- and natural-origin fish (Figure 5) and statistical tests of differences between hatchery- and natural-origin fish visually observed at Tumwater Dam and Wells Dam (Table 1). In the evaluation of PIT-tag returns, hatchery- and natural-origin arrival timing are a near match at both sampling locations (Figure 5). At Wells Dam, hatchery-origin fish arrived slightly earlier during the second half of the return distribution (e.g., 6 days earlier at the 75th percentile).

PIT-tag-based, within-year data are limited by sample size, particularly for natural-origin PIT-tagged fish; however, we provide Figure 6 as an example of the variability in migration timing that may occur on a year-by-year basis. Even with this variability, the general trends discussed above (i.e., hatchery- and natural-origin overlap and later arriving age- 3 fish) are evident.

Based on visual observations, arrival timing at both dams showed similar results with only slight differences from the PIT-tag based evaluation. The differences in arrival timing between hatchery- and natural-origin fish at the 10th, 50th, and 90th percentiles were again minor (range $=-2.28-1.14$ days). The paired differences were significant on the 10th percentile $(\mathrm{P}$-value $=0.0228)$ at Tumwater Dam and the 10th $(\mathrm{P}$-value $=0.0179)$ and 50 th percentile $(\mathrm{P}-$ value $=0.0206$ ) at Wells Dam; however, the magnitude of the differences was small and in a different temporal direction than the differences observed with PIT-tag data. As described above, where there was a difference in the PIT-tagged fish evaluation, hatchery-origin fish arrived earlier than natural-origin fish. Using visual observations, hatchery-origin fish consistently
arrived slightly later than natural-origin fish. In either case, the differences were small relative to the variability observed within and between years.

When viewed year-by-year, the differences between hatchery- and natural-origin arrival timing in the visual observation data show that hatchery-origin fish arrived later than naturalorigin fish in most years (Figure 7). However, the temporal trend in differences at both Tumwater Dam and Wells Dam is generally towards no difference and the only significant trend was at Wells Dam for the 50th percentile arrival date.


Figure 1. Return timing of PIT-tagged natural-origin (W, red) and hatchery-origin (H, blue) spring Chinook Salmon (age-3, 4, and 5) from the Methow and Wenatchee sub-basins to Bonneville Dam, return years 2006-2018. Data grouped by basin, origin, and river or tributary of release.


Figure 2. Return timing to Tumwater Dam of PIT-tagged natural-origin (W, red) and hatcheryorigin (H, blue) spring Chinook Salmon (age-3, 4, and 5) from the Wenatchee Sub-basin tagged upstream from Tumwater Dam, return years 2006-2018. Data grouped by basin, origin, and river or tributary of release.


Figure 3. Return timing of PIT-tagged natural-origin (W, red) and hatchery-origin (H, blue) spring Chinook Salmon (age-3, 4, and 5) from the Methow Sub-basin to Wells Dam, return years 2006-2018. Data grouped by basin, origin, and river or tributary of release.


Figure 4. Return timing of PIT-tagged natural-origin (W, red) and hatchery-origin (H, blue) spring Chinook Salmon by age at return from the Methow and Wenatchee sub-basins to Priest Rapids Dam, return years 2006-2018. Data grouped by basin, origin, and river or tributary of release.


Figure 5. Cumulative frequency plots of migration timing of adult hatchery- and natural-origin spring Chinook Salmon passing Tumwater Dam (TUF) and Wells Dam (WEA). Migration timing was based on PIT-tagged fish detected during 2005-2018 migration years. Sample sizes at Tumwater Dam were 821 hatchery-origin and 490 natural-origin fish, and at Wells Dam 1,258 hatchery- and 137 natural-origin fish.

Table 1. Results of paired t-tests and $95 \%$ confidence limits on the $10 \%$, $50 \%$ (median), and $90 \%$ day of the year that hatchery- and natural-origin spring Chinook Salmon migrated over Tumwater Dam (1998-2018) and Wells Dam (2006-2018). Migration timing was based on video monitoring and visual observations at both dams.

| Location | Statistic | 10th percentile | 50th percentile | 90th percentile |
| :---: | :--- | ---: | ---: | ---: |
| Tumwater Dam | Mean Difference | -2.00 | -1.19 | 1.14 |
|  | Upper 95\% Mean | -0.31 | 0.22 | 3.14 |
|  | Lower 95\% Mean | -3.69 | -2.60 | -0.85 |
|  | N | 21 | 21 | 21 |
|  | Test Statistic | -2.47 | -1.76 | 1.19 |
|  | Prob > \|t| | 0.023 | 0.093 | 0.246 |
| Wells Dam | Mean Difference | -2.48 | -1.88 | -0.75 |
|  | Upper 95\% Mean | -0.51 | -0.34 | 1.17 |
|  | Lower 95\% Mean | -4.46 | -3.43 | -2.66 |
|  | N | 13 | 13 | 13 |
|  | Test Statistic | -2.74 | -2.66 | -0.84 |
|  | Prob > \|t| | 0.018 | 0.021 | 0.412 |



Figure 6. Return timing of PIT-tagged natural-origin (W, red) and hatchery-origin (H, blue) spring Chinook Salmon by age at return from the Wenatchee Sub-basin to Tumwater Dam by year (2009-2018).


Figure 7. The difference in days (natural-origin minus hatchery-origin) that $10 \%, 50 \%$ (median), and $90 \%$ of natural-origin and hatchery-origin spring Chinook Salmon passed Tumwater Dam (1998-2018) and Wells Dam (2006-2018) and the mean difference and trend of that time period. Negative values indicate that hatchery-origin fish passed later in the year. Passage timing was based on visual observations at each dam (data from Table 1, used for paired t-test).

Spawn Timing (Carcass Recovery) of Spring Chinook Salmon in the Wenatchee and Methow Sub-basins
From 1992-2018, carcass recovery of female spring Chinook Salmon in the Wenatchee and Methow sub-basins generally began in early August and extended to early October (Figure 8). Across their range in both sub-basins, hatchery- and natural-origin fish generally had similar and overlapping spawn timing. Across all years, spawn timing was consistent with median carcass recovery dates generally varying between one and two weeks over more than 20 years of sampling (Figure 9). On a year-by-year basis, hatchery-origin fish generally matched the timing of natural-origin spawning (Figure 9 and Figure 10).

The means of paired differences at the 10th, 50th, and 90th timing percentile were small relative to the variation in natural-origin spawn timing within years and, in most cases, not significant at the 0.05 level (Table 2 and Table 3). At the 50th percentile (median), only the Twisp River had a significant difference between hatchery- and natural-origin spawning. However, this analysis and deriving percentiles from distributions was limited by small sample sizes (Table 2 and 3.). For example, in years with fewer than ten fish, percentile metrics (e.g., 10th, 50th, and 90th) should be viewed with caution. Similar limitations exist in the White River. In the Chiwawa River, at the 10th percentile hatchery-origin fish spawned on average 3.29 days earlier than natural-origin fish and this difference was significant. Figure 9 illustrates this difference, and the annual variation associated with it.


Figure 8. Spawn timing of hatchery- (H, blue) and natural-origin (W, red) female spring Chinook Salmon in the Wenatchee (top, 1992-2018) and Methow (bottom, 1993-2018) sub-basins by stream or tributary. Spawn timing was based on the day of the year that female carcasses were recovered on the spawning grounds. Each dot represents one female. Boxes cover the 25-75\% percentile; dots beyond the whiskers are outliers. Sample sizes on right border.



Figure 9. The 10th, 50th (median), and 90th percentile spring Chinook Salmon spawn day of the year in the Wenatchee (top) and Methow (bottom) sub-basins by year and stream. Estimates are based on recovery of female carcasses on spawning grounds.


Figure 10. The distributions of spawn timing of hatchery- and natural-origin spring Chinook Salmon by year in the Wenatchee (top) and Methow (bottom) sub-basins based on female carcass recovery. Sample size by origin above the x-axis. Boxes span the $25-75$ th percentiles. Line inside box $=50$ th percentile $($ median $)$. Dashes outside of boxes $=10$ th $/ 90$ th, 2.5 th $/ 97.5$ th, and minimum/maximum.

Table 2. Results of paired t-tests and $95 \%$ CIs on the $10 \%, 50 \%$ (median), and $90 \%$ day of spawn timing (based on female carcass recovery) of hatchery- and natural-origin spring Chinook Salmon in the Wenatchee Sub-basin during the period 1993-2018.

| Location | Statistic | 10th percentile | 50th percentile | 90th percentile |
| :---: | :---: | :---: | :---: | :---: |
| Chiwawa River | Mean Difference | 3.29 | 0.59 | 2.08 |
|  | Upper 95\% Mean | 6.55 | 3.11 | 5.87 |
|  | Lower 95\% Mean | 0.04 | -1.93 | -1.71 |
|  | N | 23 | 23 | 23 |
|  | Test Statistic | 2.09 | 0.48 | 1.13 |
|  | Prob $>\|t\|$ | 0.048 | 0.634 | 0.268 |
| Nason Creek | Mean Difference | -1.75 | -1.92 | -0.45 |
|  | Upper 95\% Mean | 1.28 | 0.62 | 2.05 |
|  | Lower 95\% Mean | -4.79 | -4.45 | -2.95 |
|  | N | 24 | 24 | 24 |
|  | Test Statistic | -1.20 | -1.56 | -0.37 |
|  | Prob > $\|t\|$ | 0.244 | 0.132 | 0.714 |
| White River | Mean Difference | -3.55 | -1.84 | 2.38 |
|  | Upper 95\% Mean | 3.88 | 0.54 | 8.03 |
|  | Lower 95\% Mean | -10.98 | -4.23 | -3.28 |
|  | N | 16 | 16 | 16 |
|  | Test Statistic | -1.02 | -1.65 | 0.90 |
|  | Prob $>\|t\|$ | 0.324 | 0.120 | 0.385 |

Table 3. Results of paired t-tests and $95 \%$ CIs on the $10 \%, 50 \%$ (median), and $90 \%$ day of spawn timing (based on female carcass recovery) of hatchery- and natural-origin spring Chinook Salmon in the Methow Sub-basin during the period 1999-2018.

| Location | Statistic | 10th percentile | 50th percentile | 90th percentile |
| :---: | :---: | :---: | :---: | :---: |
| Chewuch River | Mean Difference | -1.87 | -0.76 | -0.25 |
|  | Upper 95\% Mean | 1.64 | 0.37 | 2.60 |
|  | Lower 95\% Mean | -5.39 | -1.89 | -3.10 |
|  | N | 19 | 19 | 19 |
|  | Test Statistic | -1.12 | -1.42 | -0.18 |
|  | Prob > $\|t\|$ | 0.277 | 0.174 | 0.858 |
| Methow River | Mean Difference | -2.53 | -1.00 | 1.02 |
|  | Upper 95\% Mean | 0.07 | 0.26 | 3.58 |
|  | Lower 95\% Mean | -5.13 | -2.26 | -1.55 |
|  | N | 20 | 20 | 20 |
|  | Test Statistic | -2.03 | -1.66 | 0.83 |
|  | Prob > $\|t\|$ | 0.056 | 0.113 | 0.417 |
| Twisp River | Mean Difference | -5.95 | -2.97 | -0.76 |
|  | Upper 95\% Mean | -1.58 | -0.06 | 2.44 |
|  | Lower 95\% Mean | -10.32 | -5.89 | -3.96 |
|  | N | 18 | 18 | 18 |
|  | Test Statistic | -2.87 | -2.15 | -0.50 |
|  | Prob $>\|t\|$ | 0.011 | 0.046 | 0.622 |

Spawning Distribution of Spring Chinook Salmon in the Chiwawa River
Both hatchery- and natural-origin spring Chinook Salmon spawned throughout the Chiwawa River and within each reach; however, there was a significant difference in the distribution of hatchery- and natural-origin spawners among historical survey reaches for years 1993-2018 (Yates' Chi-square $=371.914 ; \mathrm{P}=0.000 ;$ Effect Size $=0.342$ ) (Figures 11 and 12). The proportion of natural-origin spawners (pNOS) was highest within reaches C3 through C6 ( $\mathrm{pNOS}=0.30-0.56$ ); the proportion of hatchery-origin spawners ( $\mathrm{pHOS} \mathrm{)} \mathrm{was} \mathrm{highest} \mathrm{within}$ reaches $\mathrm{C} 1(\mathrm{pHOS}=0.89), \mathrm{C} 2(\mathrm{pHOS}=0.70)$, and $\mathrm{C} 7(\mathrm{pHOS}=0.75)$ (Figure 13). In all reaches, pHOS exceeded 0.30 and it exceeded 0.50 within five of the seven reaches. When comparing spawn distribution at the $0.5-\mathrm{RKm}$ scale, the greatest proportion of hatchery-origin spring Chinook Salmon spawned in the lower 25 km of the river, while the greatest proportion of natural-origin fish spawned between km 30 and 55 (Figure 14).

## Spawning Distribution of Spring Chinook Salmon in Nason Creek

The Nason Creek Hatchery Program began producing adult returns in 2016. Prior to 2016, hatchery-origin spring Chinook Salmon that spawned in Nason Creek were strays, mainly from the Chiwawa spring Chinook Salmon Hatchery Program. Hatchery- and natural-origin spring Chinook Salmon spawned throughout Nason Creek; however, there was a significant
difference in the distribution of hatchery- and natural-origin spawners among historical survey reaches for years 1993-2018 (Yates' Chi-square $=158.721 ; \mathrm{P}=0.000$; Effect Size $=0.296$ ) (Figures 11 and 15). The pHOS exceeded 0.30 in all reaches and was highest within reaches N1 $(\mathrm{pHOS}=0.79)$ and $\mathrm{N} 2(\mathrm{pHOS}=0.61)$. The pHOS and pNOS were close to equal within reaches N 3 and N 4 where $\mathrm{pHOS}=0.46$ and $\mathrm{pNOS}=0.54$ for both reaches (Figure 13). When comparing spawn distribution at the $0.5-\mathrm{RKm}$ scale, the highest proportion of hatchery-origin spring Chinook Salmon spawned in the lower 7 km of Nason Creek; the highest proportion of naturalorigin spring Chinook Salmon spawned upstream from RKm 16 (Figure 14).

## Spawning Distribution of Spring Chinook Salmon in the White River

There are three historical reaches $(\mathrm{H} 2, \mathrm{H} 3$, and H 4$)$ in the White River and most of the hatchery and natural-origin Chinook Salmon spawned in reach H3. Many of the hatchery-origin spawners were strays from the Chiwawa Hatchery program. There was no significant difference in the distribution of hatchery- and natural-origin spawners among the three historical survey reaches for years 1993-2018 (Yates' Chi-square $=0.155 ; \mathrm{P}=0.925$; Effect Size $=0.49$ ) (Figures 11 and 16). The pNOS was higher than pHOS within all three reaches and pNOS was highest in the uppermost reach ( $\mathrm{pNOS}=0.71$ ) (Figure 13). The pHOS exceeded 0.30 in two of the three reaches. When comparing spawn distribution at the $0.5-\mathrm{RKm}$ scale, the highest proportion of hatchery- and natural-origin spring Chinook Salmon spawned between RKm 20 and 24 (Figure 14).

Chiwawa River


Figure 11. Proportion of hatchery- and natural-origin spring Chinook Salmon spawners within a stream that were observed within each historical reach on the Chiwawa River (top), Nason Creek (middle), and on the White River (bottom) during the period 1993-2018. Here, the proportion of natural-origin or hatchery-origin spawners across historical reaches sum to " 1.00 ." The line represents the proportion of hatchery-origin spawners within each historical reach.


Figure 12. Map of the Chiwawa River showing locations of historical spring Chinook Salmon survey reaches.


Figure 13. Proportion of hatchery- and natural-origin spring Chinook Salmon spawners within each of the historical sampling reaches on the Chiwawa River (top), Nason Creek (middle), and the White River (bottom) during the period 1993-2018. Here, pNOS and pHOS sum to " 1.00 " within each reach.


Figure 14. Proportion of natural- and hatchery-origin spring Chinook Salmon spawners distributed along the length of the Chiwawa River (top), Nason Creek (middle), and White River (bottom) during the period 1993-2018. Distribution was based on $0.5-\mathrm{km}-l o n g$ reaches. Chiwawa River sample sizes $=716$ natural- and 1,686 hatchery-origin fish, Nason Creek sample sizes $=$ 411 natural- and 808 hatchery-origin fish, and White River sample sizes $=94$ natural- and 83 hatchery-origin fish.


Figure 15. Map of Nason Creek showing locations of historical spring Chinook Salmon survey reaches.


Figure 16. Map of the White River showing locations of historical spring Chinook Salmon survey reaches.

Spawning Distribution of Spring Chinook Salmon in the Chewuch River
Both hatchery- and natural-origin spring Chinook Salmon spawned throughout the Chewuch River and within each reach; however, there was a significant difference in the distribution of hatchery- and natural-origin spawners among historical survey reaches for years 1996-2018 (Yates' Chi-square $=205.772 ; \mathrm{P}=0.000$; Effect Size $=0.329$ ) (Figures 17 and 18). The pHOS was higher than or equal to the pNOS within 10 out of 13 reaches (Figure 19) and, in general, pHOS declined in an upstream direction. The pHOS was greater than 0.30 in all reaches ( pHOS range $=0.31-0.92$ ). When comparing spawn distribution at the $0.5-\mathrm{RKm}$ scale, the highest proportions of both hatchery- and natural-origin spring Chinook Salmon spawned in the lower 55 km of the Chewuch River (Figure 20). A greater proportion of hatchery-origin spring Chinook Salmon spawned in the lower 20 km of the river, while a larger proportion of naturalorigin spring Chinook Salmon spawned upstream from RKm 25.

Spawning Distribution of Spring Chinook Salmon in the Methow River
Both hatchery- and natural-origin spring Chinook Salmon spawned throughout the Methow River and within each reach; however, there was a significant difference in the distribution of hatchery- and natural-origin spawners among historical survey reaches for years 1996-2018 (Yates' Chi-square $=370.839 ; \mathrm{P}=0.000$; Effect Size $=0.314$ ) (Figures 17 and 21). The pHOS was higher than the pNOS within 12 of the 15 reaches (Figure 19). The pHOS exceeded 0.30 in 13 of the 15 reaches. When comparing spawning distribution at the $0.5-\mathrm{RKm}$ scale, a larger proportion of hatchery-origin fish spawned between RKm 80 and 90, while a larger proportion of natural-origin fish spawned upstream from RKm 90 (Figure 20).

Spawning Distribution of Spring Chinook Salmon in the Twisp River
Both hatchery- and natural-origin spring Chinook Salmon spawned throughout the Twisp River and within each reach; however, there was a significant difference in the distribution of hatchery- and natural-origin spawners among historical survey reaches for years 1996-2018 (Yates' Chi-square $=29.803 ; \mathrm{P}=0.000$; Effect Size $=0.244$ ) (Figures 17 and 22). The pHOS was higher than the pNOS within reaches T1-T5 and lower within reaches T6-T9 (Figure 19). The pHOS exceeded 0.30 in all reaches. When comparing spawn distribution at the $0.5-\mathrm{RKm}$ scale, a larger proportion of hatchery-origin fish spawned downstream from RKm 25, while a larger proportion of natural-origin fish spawned upstream from RKm 20 (Figure 20).

Chewuch River


Methow River


Twips River


Figure 17. Proportion of hatchery- and natural-origin spring Chinook Salmon spawners within a stream that were observed within each historical reach on the Chewuch River (top), the Methow River (middle), and the Twisp River (bottom) during the period 1996-2018. Here, the proportion of natural-origin or hatchery-origin spawners across historical reaches sum to " 1.00 ." The line represents the proportion of hatchery-origin spawners within each historical reach.


Figure 18. Map of the Chewuch River showing locations of historical spring Chinook Salmon survey reaches.

Chewuch River


Twisp River


Figure 19. Proportion of hatchery- and natural-origin spring Chinook Salmon spawners within each of the historical sampling reaches on the Chewuch River (top), Methow River (middle), and the Twisp River (bottom) during the period 1996-2018. Here, pNOS and pHOS sum to " 1.00 " within each reach.


Figure 20. Proportion of natural- and hatchery-origin spring Chinook Salmon spawners distributed along the length of the Chewuch River (top), Methow River (middle), and Twisp River (bottom) during the period 1996-2018. Distribution was based on $0.5-\mathrm{km}$-long reaches. Chewuch River sample sizes $=393$ natural- and 782 hatchery-origin fish, Methow River sample sizes $=366$ natural- and 1,677 hatchery-origin fish, and Twisp River sample sizes $=164$ naturaland 234 hatchery-origin fish.


Figure 21. Map of the Upper Methow River showing locations of historical spring Chinook Salmon survey reaches.


Figure 22. Map of the Twisp River showing locations of historical spring Chinook Salmon survey reaches.

## Migration Timing of Wenatchee and Methow Summer Chinook Salmon

At Dryden Dam, based on 12 years of sampling, there were significant differences in the migration timing of hatchery- and natural-origin summer Chinook Salmon at the 10th and 50th percentiles (Table 4). At most, the average difference in migration timing between hatchery- and natural-origin fish was 2.5 weeks. This later arrival date of hatchery-origin fish at Dryden Dam was generally consistent year-by-year and throughout the arrival distribution (Figure 23). At Wells Dam, migration timing between hatchery- and natural-origin fish were a near match; only at the tail end of the migration (90th percentile) was there a small measurable difference ( 1 week) and this was not significant. However, the run of hatchery-origin summer Chinook at Wells Dam consists of a mix of fish released in the Methow sub-basin, Okanogan sub-basin, and Columbia River, plus stray fish from downstream locations.

Table 4. Results of paired t-tests and $95 \%$ CIs on the $10 \%, 50 \%$ (median), and $90 \%$ day of migration timing (weeks) of hatchery- and natural-origin summer Chinook Salmon at Dryden Dam and Wells Dam during the period 2007-2018.

| Location | Statistic | 10th percentile | 50th percentile | 90th percentile |
| :---: | :--- | ---: | ---: | ---: |
| Dryden Dam | Mean Difference | -1.08 | -2.50 | -0.91 |
|  | Upper 95\% Mean | -0.13 | -0.61 | 1.17 |
|  | Lower 95\% Mean | -2.04 | -4.39 | -2.99 |
|  | N | 12 | 12 | 12 |
|  | Test Statistic | -2.49 | -2.92 | -0.96 |
|  | Prob > $\|\mathrm{t}\|$ | 0.030 | 0.014 | 0.358 |
| Wells Dam | Mean Difference | -0.18 | 0.08 | 1.00 |
|  | Upper 95\% Mean | 0.20 | 0.51 | 2.33 |
|  | Lower 95\% Mean | -0.53 | -0.34 | -0.33 |
|  | N | 12 | 12 | 12 |
|  | Test Statistic | -1.00 | 0.432 | 1.66 |
|  | Prob > $\|\mathrm{t}\|$ | 0.339 | 0.674 | 0.126 |



Figure 23. The difference in weeks (natural-origin minus hatchery-origin) that $10 \%, 50 \%$ (median), and $90 \%$ of natural-origin and hatchery-origin summer Chinook Salmon passed Dryden Dam and Wells Dam from 2007-2018 and the mean difference of that time period. Negative values indicate that hatchery-origin fish passed later in the year. Passage timing was based on run composition sampling and broodstock collection.

Spawn Timing (Carcass Recovery) of Summer Chinook Salmon in the Wenatchee and Methow Sub-basins

From 1993-2018, female carcass recovery of summer Chinook Salmon in the Wenatchee and Methow sub-basins generally began in mid-September and extended to mid-November (Figure 24). Summer Chinook Salmon in the Wenatchee Sub-basin generally spawned earlier than fish in the Methow Sub-basin and in both sub-basins hatchery-origin fish consistently spawned later than natural-origin fish (Figure 24). However, the difference in timing between hatchery- and natural-origin fish was generally small (seven days or less $90 \%$ of the time at the 50th percentile spawn date) and hatchery-origin fish generally followed the timing pattern of natural-origin fish at the 10th, 50th, and 90th percentiles (Figure 25).

The mean of paired differences at the 10th, 50th, and 90th timing percentile reflected the consistent later date of spawning of hatchery-origin fish in both sub-basins (Table 5). In the Wenatchee Sub-basin, hatchery-origin fish spawned significantly later then natural-origin fish at the 50th and 90th percentile with a mean difference of -2.4 days at the 50th percentile. In the Methow Sub-basin, hatchery-origin fish were significantly later statistically at all three levels of measurement with a mean difference of -3.6 days at the 50th percentile. There was no significant directional temporal trend in the differences between hatchery- and natural-origin fish in either sub-basin (Figure 26); however, over the period of data collection, spawn timing for both hatchery- and natural-origin fish shifted to significantly earlier in the year (Figure 27).


Figure 24. The temporal distribution of spawn timing of summer Chinook Salmon in the Wenatchee and Methow sub-basins from 1993-2018. Sample size by sub-basin and origin on right margin. Boxes span the 25-75th percentiles. Line inside box $=50$ th percentile (median). Dashes outside of boxes $=10 \mathrm{th} / 90 \mathrm{th}, 2.5 \mathrm{th} / 97.5 \mathrm{th}$, and minimum $/$ maximum.



Figure 25. The 10th, 50th (median), and 90th percentile spawn day of the year of summer Chinook Salmon in the Wenatchee (top) and Methow (bottom) sub-basins by year.

Table 5. Results of paired t-tests and $95 \%$ CIs on the $10 \%, 50 \%$ (median), and $90 \%$ day of spawn timing of hatchery- and natural-origin summer Chinook Salmon in the Wenatchee and Methow sub-basins during the period 1993-2018.

| Location | Statistic | 10th percentile | 50th percentile | 90th percentile |
| :---: | :---: | :---: | :---: | :---: |
| Wenatchee Sub-basin | Mean Difference | -0.73 | -2.42 | -1.75 |
|  | Upper 95\% Mean | 0.20 | -1.28 | -0.580 |
|  | Lower 95\% Mean | -1.68 | -3.57 | -2.92 |
|  | N | 26 | 26 | 26 |
|  | Test Statistic | -1.60 | -4.36 | -3.078 |
|  | Prob > $\|t\|$ | 0.120 | 0.000 | 0.005 |
| Methow Sub-basin | Mean Difference | -2.69 | -3.62 | -2.72 |
|  | Upper 95\% Mean | -0.36 | -2.02 | -1.29 |
|  | Lower 95\% Mean | -5.02 | -5.21 | -4.15 |
|  | N | 26 | 26 | 26 |
|  | Test Statistic | -2.37 | -4.66 | -3.91 |
|  | Prob > $\|t\|$ | 0.0255 | 0.000 | 0.001 |



Figure 26. The difference in day (natural-origin minus hatchery-origin) that $10 \%, 50 \%$ (median), and $90 \%$ of hatchery- and natural-origin summer Chinook Salmon spawned from 1993-2018 and the mean difference of that time period. Negative values indicate that hatchery-origin fish spawned later in the year. Spawn timing was based on carcass recovery date.


Figure 27. The mean, interquartile range, and trend of summer Chinook Salmon carcass recoveries in the Wenatchee and Methow sub-basins by year.

## Spawning Distribution of Summer Chinook Salmon in the Wenatchee River

Both hatchery- and natural-origin summer Chinook Salmon spawned throughout the Wenatchee River and within each reach; however, there was a significant difference in the distribution of hatchery- and natural-origin spawners among historical survey reaches for years 1993-2018 (Yates' Chi-square $=2551.446 ; \mathrm{P}=0.000$; Effect Size $=0.345$ ) (Figures 28 and 29). The pNOS was higher than the pHOS in all 10 reaches (Figures 30). The pHOS exceed 0.30 in 3 of the 10 reaches. When comparing spawn distribution at the $1.0-\mathrm{RKm}$ scale, the highest proportions of hatchery-origin summer Chinook Salmon spawned in the lower 45 km of the Wenatchee River, while the highest proportions of natural-origin summer Chinook Salmon spawned upstream from RKm 35 (Figure 31).

Spawning Distribution of Summer Chinook Salmon in the Methow River
Both hatchery- and natural-origin summer Chinook Salmon spawned throughout the Methow River and within each reach; however, there was a significant difference in the
distribution of hatchery- and natural-origin spawners among historical survey reaches for years 1993-2018 (Yates' Chi-square $=829.375 ;$ P=0.000; Effect Size $=0.339$ ) (Figures 28 and 32). The pNOS was higher than the pHOS within six of seven reaches (Figure 30). The pHOS exceeded 0.30 in three of the seven reaches. When comparing spawn distribution at the $1.0-\mathrm{RKm}$ scale, the highest proportions of hatchery-origin summer Chinook Salmon spawned in the lower 65 km of the Methow River, while the highest proportion of natural-origin fish spawned upstream from RKm 60 (Figure 31).


Figure 28. Proportion of hatchery- and natural-origin summer Chinook Salmon spawners within a river that were observed within each historical reach on the Wenatchee River (top) and Methow River (bottom) during the period 1993-2018. Here, the proportion of natural-origin or hatchery-origin spawners across historical reaches sum to " 1.00 ." The line represents the proportion of hatchery-origin spawners within each historical reach.


Figure 29. Map of the Wenatchee River showing locations of historical summer Chinook Salmon survey reaches.


Figure 30. Proportion of hatchery- and natural-origin summer Chinook Salmon spawners within each of the historical sampling reaches on the Wenatchee River (top) and Methow River (bottom) during the period 1993-2018. Here, pNOS and pHOS sum to " 1.00 " within each reach.


Figure 31. Proportion of natural- and hatchery-origin summer Chinook Salmon spawners distributed along the length of the Wenatchee River (top) and Methow River (bottom) during the period 1993-2018. Distribution was based on 1.0-km-long reaches. Wenatchee River sample sizes $=8,044$ natural- and 2,073 hatchery-origin fish and Methow River sample sizes $=2,567$ natural- and 1,774 hatchery-origin fish.


Figure 32. Map of the Methow River showing locations of historical summer Chinook Salmon survey reaches.

## Discussion

The timing of migration and spawning of hatchery- and natural-origin spring Chinook Salmon were generally similar in the Wenatchee and Methow subbasins. In both sub-basins, the differences between hatchery- and natural-origin fish were small relative to the inter- and intraannual variability of natural-origin migration and spawn timing. Although the difference in arrival timing was small and not statistically significant, the visual observation data at Tumwater and Wells dams suggest a tendency for hatchery-origin spring Chinook Salmon to arrive later than natural-origin fish. Such differences could be explained by differences in age distribution between the two groups; hatchery-origin fish tend to mature at younger ages (Graf et al. 2020; Knudsen et al. 2008) and, as observed in Figure 4 and Figure 6, younger fish tend to arrive later. In addition, spring Chinook Salmon generally spawn earlier at higher elevations and hatcheryorigin fish generally spawn at lower elevations than natural-origin fish (Williamson et al. 2010; Fast et al. 2015). Spawner distribution may also explain differences in spawn timing. As we described, the distribution of hatchery-origin fish is strongly influenced by the location of acclimation facilities which are located at lower elevations, thus resulting in an elevation-caused spawn timing difference as well. If migration timing is correlated with spawn timing, then this could be another mechanism explaining the small difference in migration timing. Overall, from these data, it is likely that hatchery- and natural-origin fish are generally experiencing similar selective pressures during their adult freshwater migration period and during spawning.

Conversely, the differences in migration timing between spring Chinook Salmon populations in the Wenatchee versus Methow sub-basins likely has resulted in differences in survival at the adult life-stage. Mortality events such as predation and harvest are uneven within a year, and in the case of predation by pinnipeds downstream from Bonneville Dam, earlier arriving fish experience higher mortality rates (Wargo Rub et al. 2019). Based on PIT-tag arrival timing at Bonneville Dam, this suggests that fish from the Methow Hatchery, Winthrop National Fish Hatchery, and Leavenworth National Fish Hatchery may have experienced the highest rates of predation by pinnipeds (Sorel et al. 2020).

Differences in hatchery- and natural-origin summer Chinook migration timing varied between the Wenatchee and Methow/Okanogan sub-basins. Hatchery-origin fish observed at Wells Dam include fish released in the Methow and Okanogan sub-basins, plus the Columbia River. Natural populations consist of the Methow and Okanogan sub-basins, plus fish produced in the Columbia River. Thus, the comparison at Wells Dam consists of mixed stocks of both hatchery- and natural-origin fish. Differences in spawn timing of summer Chinook also varied between the Wenatchee and Methow sub-basins. For migration timing, hatchery-origin fish in the Wenatchee sub-basin consistently passed Dryden Dam later than their natural-origin counterparts; whereas, the migration timing for Methow/Okanagon hatchery-origin fish were a near match with natural-origin fish at Wells Dam. Conversely, spawn timing in both Wenatchee and Methow hatchery-origin fish was later than natural-origin fish, with Methow-origin fish having a larger average difference. This pattern of arrival timing differences at Dryden Dam versus Wells Dam may be due to the relative proximity of the sampling location to the spawning grounds. In both sub-basins, hatchery- and natural-origin summer Chinook Salmon have different spawning spatial distributions where natural-origin fish tend to spawn farther upstream. This spatial difference likely manifests in a temporal difference at Dryden Dam, where hatcheryorigin fish have nearly 'arrived' at their spawning destination. The consistent later spawn date of Wenatchee and Methow hatchery-origin fish may also be explained by differences in spawning
spatial distribution. Environmental conditions higher in the sub-basin (e.g., colder temperatures and lower river flows in later summer) may be driving earlier spawn timing of natural-origin fish. Lastly, hatchery- and natural-origin summer Chinook Salmon from both sub-basins have trended towards earlier spawn dates. This consistent signal may suggest regional environmental conditions that are driving or favoring earlier spawn dates (e.g., lower flows in later summer and/or warmer river temperatures). Should this trend continue, temporal segregation of summer and spring Chinook Salmon spawning populations will diminish, increasing opportunity for these populations to mix.

Hatchery-origin spring Chinook Salmon spawned throughout the range of natural-origin spring Chinook Salmon and generally spawned in the reaches where most natural-origin fish spawned. However, differences between hatchery- and natural-origin spawner distribution across historical survey reaches were observed during some years in all spring Chinook Salmon programs except the White River program. In general, hatchery-origin females spawned farther downstream than natural-origin females, which is consistent with Hughes and Murdoch (2017), who concluded that over a 10-year period, a greater proportion of hatchery-origin females spawned in lower reaches of the Chiwawa River which is likely the result of the acclimation facilities being located in the lower survey reaches. Nevertheless, the greater abundance of hatchery-origin relative to natural-origin fish resulted in exceedances of HSRG recommended pHOS objectives for reducing the risk of domestication selection (Mobrand et al. 2005; Paquet et al. 2011) in nearly every spawning reach ( 13 of 14 reaches in the Wenatchee Sub-basin; 35 of 37 reaches in the Methow Sub-basin). Thus, even though more hatchery-origin fish spawned lower in each sub-basin than natural-origin fish, returns of hatchery-origin fish to each sub-basin were sufficient to dominate or substantially influence nearly the entire spawning distribution.

The distribution of hatchery-origin fish is likely strongly influenced by the location of acclimation facilities. For example, in the Chiwawa River, juvenile hatchery-origin spring Chinook Salmon are overwinter acclimated at the Chiwawa Acclimation Facility and released volitionally into the river. Most of the returns of hatchery-origin fish released from this facility spawn within the lower reaches of the Chiwawa River. Hughes and Murdoch (2017) found that hatchery-origin fish that spawned farther downstream in the Chiwawa River sub-basin near the program's overwinter acclimation site had lower relative reproductive success compared to natural-origin spring Chinook Salmon. Hughes and Murdoch (2017) believe the mechanism for the reduced relative reproductive success between hatchery- and natural-origin Chiwawa spring Chinook Salmon is related to the lower-quality spawning habitat used by hatchery-origin females in the lower reaches of the Chiwawa River compared to the middle and upper reaches. The Upper Columbia Regional Technical Team of the Upper Columbia Salmon Recovery Board ranked the lower Chiwawa River reaches as high-priority reaches for restoring reach functions to improve spawning and rearing habitat (Upper Columbia Salmon Recovery Board 2021).

The spawning distribution of spring Chinook Salmon in Nason Creek was likely influenced by strays from the Chiwawa Hatchery program, the reduction in hatchery smolt production in the Chiwawa River, and the initiation of the Nason Creek Hatchery program. Prior to the Nason Hatchery program, strays from the Chiwawa Hatchery program spawned mostly in the downstream sections of Nason Creek whereas greater proportions of the natural-origin fish spawned upstream. At the time the Nason hatchery program started, the number of strays from the Chiwawa program decreased as a result of lowered production and this resulted in fewer fish spawning in lower reaches. At the same time, returns from the Nason Creek hatchery program
returned to upstream areas and the spawning distribution more closely matched that of the natural-origin distribution. The difference in distribution of the Chiwawa and Nason hatcheryorigin fish was likely a result of the locations of the acclimation sites; with the Nason Creek Acclimation Facility located upstream in Nason Creek and the Chiwawa Acclimation Facility located downstream in the Chiwawa River.

The spawning distribution of natural- and hatchery-origin spring Chinook Salmon in the White River was mainly confined to the middle spawning reach. Hatchery-origin fish from the White River captive broodstock program and strays from the Chiwawa Hatchery program all spawned primarily in the reach that was used most by the natural-origin fish and this reach was upstream of where both of these hatchery-origin groups were acclimated. Hatchery-origin fish migrated past portions of the lower White River presumably because spawning habitat was unsuitable in lower reaches. A similar phenomenon was observed for spring Chinook Salmon in the upper Yakima River where habitat suitability was low adjacent to the Clark Flats acclimation site (Cram et al. 2013).

The spawning distributions of spring Chinook Salmon in the Chewuch, Methow, and Twisp rivers were likely influenced by the location of acclimation sites. In all three streams, the distribution of spawning spring Chinook Salmon differed significantly between natural- and hatchery-origin fish with more hatchery-origin fish spawning within the lower survey reaches than natural-origin fish. A greater proportion of natural-origin fish spawned in the upper survey reaches in all three streams. On the Methow River, a large proportion of hatchery-origin spring Chinook Salmon spawned between RKms 83 and 87, which is where the Methow Salmon Hatchery is located. Most of the natural-origin spring Chinook Salmon on the Methow River spawned upstream of RKm 90.

The proportion of hatchery-origin spawners was high within nearly all spring Chinook Salmon survey reaches on Nason Creek and the Chiwawa, White, Chewuch, Methow, and Twisp rivers. The pHOS exceeded $50 \%$ within 33 of the 51 survey reaches. The HSRG recommended pHOS below $30 \%$ for integrated populations (Mobrand et al. 2005; Paquet et al. 2011), but this recommendation was met in only 3 of the 51 spring Chinook Salmon survey reaches. Although the HSRG recommendation for pHOS is at the population scale, we evaluated pHOS at a reach base scale to better understand pHOS distribution within each of the spawning reaches. Managing for pHOS while managing for similar spawn distribution between hatchery- and natural-origin fish can create competing management objectives. In the majority of the lower and middle reaches, pHOS is too high (i.e., greater than $50 \%$ ). Implementing adult management can likely reduce pHOS at both the population and reach levels; however, due to the recent years of low adult returns, the requirements to implement adult management have not been met. Increasing homing of hatchery-origin spawners to the upper reaches, where they are often limited, compared to natural-origin spawners, would increase the similarity of the spawner distributions. However, concurrently, pHOS would likely increase to greater than $50 \%$ in all reaches because the number of natural-origin spawners in the upper reaches is so low, that an increase of only a few hatchery-origin spawners would dramatically elevate pHOS in those reaches. Furthermore, because the natural-origin abundance is so low in the upper reaches and thus able to accommodate only very few additional hatchery-origin fish and still maintain pHOS less than $30 \%$, we cannot expect that increasing abundance of hatchery-origin fish in those upstream reaches would increase natural production appreciably. Increasing natural-origin
spawners in all reaches would improve overall and reach-level pHOS values and provide scope for extending the distribution of hatchery-origin spawners into underutilized spawning reaches.

As stated earlier, the management objective for the summer Chinook Salmon hatchery programs analyzed was hatchery-origin spawners to not replicate the spawning distribution of natural-origin fish. Returns from both Wenatchee summer Chinook Salmon and Methow summer Chinook Salmon hatchery programs achieved this objective by exhibiting spawner distributions significantly different than natural-origin spawners; the hatchery-origin fish spawned lower in the rivers than the natural-origin fish. It is also important that the hatchery programs do not create overlap in the spawning distributions of spring and summer Chinook Salmon. In the Entiat River, summer Chinook Salmon superimposed spring Chinook Salmon redds and some inbreeding occurred (Fraser et al. 2020; Fraser and Cooper 2022).

The pNOS was higher in 16 of the 17 Wenatchee and Methow summer Chinook Salmon spawn survey reaches, which is likely the result of the large number of natural-origin spawners and a popular recreational fishery that targets hatchery-origin summer Chinook Salmon. The pHOS exceeded $30 \%$ in 6 of the 17 summer Chinook Salmon survey reaches and exceeded $50 \%$ in only one reach.

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# Stray Rates of Natural-Origin Chinook Salmon and Steelhead in the Upper Columbia Watershed 

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#### Abstract

Despite the importance of straying in understanding the ecology of salmon and steelhead, most of what is known about salmon and steelhead straying comes from tagged hatchery fish. We provide donor estimates of natural-origin spring, summer, and fall Chinook Salmon Oncorhynchus tshawytscha and steelhead Oncorhynchus mykiss straying at three spatial scales in the upper Columbia watershed using Passive Integrated Transponder (PIT) tags. A total of 823,770 natural-origin spring, summer, and fall Chinook Salmon and summer steelhead were PIT-tagged as juveniles in the Wenatchee, Entiat, Methow, and Okanogan River subbasins and tributaries and the upper Columbia River between 2002 and 2017. Anadromous adults with PIT tags were detected at a variety of antenna arrays in the Columbia River Basin between 2004 and $2018(\mathrm{n}=2,611)$. Mean donor stray rates of each population were less than $1 \%$ at the basin scale (range $0.0 \%-0.7 \%$ ), less than $10 \%$ at the subbasin scale (range $0.0 \%-9.8 \%$ ) and less than $15 \%$ at the tributary scale (range $0.0 \%-14.3 \%$ ). Many of the populations ( 11 of 28 ) that were evaluated across all spatial scales did not have any strays detected, and the mean of means of all species stray rates at all spatial scales was generally less than $5 \%$ (range $0.2 \%-4.0 \%$ ). Chinook Salmon and steelhead strayed at similar rates when originating from the same subbasins and tributaries. Most straying occurred in an upstream direction at the subbasin (84\%) and tributary scales ( $94 \%$ ). Variation in stray rates was most consistently associated with spatial scale and location and was less than $15 \%$ for all species at all spatial scales.


## Introduction

Straying by salmon and steelhead is an important mechanism for colonizing new habitats (Quinn 2005; Keefer and Caudill 2014; Westley et al. 2015). However, it can also reduce the spawning population of donor populations and disrupt local adaptation of recipient populations if it occurs at high rates (Ford 2002; Mobrand et al. 2005; Brenner et al. 2012). Most of what is known about salmon and steelhead straying comes from studies of tagged hatchery fish (Dittman et al. 2010; Westley et al. 2013; Keefer and Caudill 2014). Access to large numbers of fish in controlled environments and high tag rates provide great opportunities to learn about straying (Dittman et al. 2010; Westley et al. 2013; Bond et al. 2017). Although estimates of hatcheryorigin fish straying are informative, they may be very different from estimates of natural-origin salmon and steelhead (Keefer and Caudill 2014; Dittman et al. 2015).

Surprisingly few estimates of natural-origin Chinook Salmon Oncorhynchus tshawytscha and steelhead Oncorhynchus mykiss straying have been published despite the importance to understanding the metapopulation dynamics of these fish and how these estimates might inform expectations about stray rates of hatchery-origin salmon and steelhead (Quinn 2005; Keefer and Caudill 2014; Fullerton et al. 2016). Dispersal rate was found to be very important in metapopulation structure of modelled Chinook Salmon populations in the Snake River Basin, however they acknowledged that they had few empirical data to estimate dispersal rates among populations (Fullerton et al. 2016). Because of the difficulty of capturing, tagging and recapturing sufficient numbers of wild juveniles there are a lack of studies on stray rates of natural-origin fish. This is particularly true for species with low survival rates following tagging because more fish have to be collected to generate reasonable estimates. Shapovalov and Taft (1954) performed one of the earliest studies of stray rates of natural-origin fish involving more than one species. They studied stray rates of tagged Coho Salmon and steelhead in two coastal California creeks that were less than 8 km apart. Other creeks were not evaluated for strays beyond the two nearby creeks; thus, their stray rates should be considered minimums. The minimum stray rate of Coho Salmon was $14.9 \%$ for Coho Salmon originating from Waddell Creek and $26.8 \%$ from Coho Salmon originating from Scott Creek. The minimum stray rate for steelhead was $1.9 \%$ for steelhead originating from Waddell Creek and $2.9 \%$ from steelhead originating from Scott Creek. It is likely that environmental conditions influenced access to home tributaries and influenced stray rates, particularly for Coho Salmon.

More recently, Ford et al. (2015a) estimated stray rates of natural-origin spring Chinook Salmon in the upper Wenatchee watershed of the Columbia River in Washington using genetic techniques. Stray rates were $4.1 \%$ for fish originating from the Chiwawa River, $17.5 \%$ for fish originating from the Little Wenatchee River, $9.0 \%$ for fish originating from Nason Creek, $1.3 \%$ for fish originating from the White River, and $100 \%$ for fish originating from the upper Wenatchee River (Ford et al. 2015a). Variation in spring Chinook Salmon stray rates were related to origin (e.g., hatchery and natural) and tributary location. They also suggested that the difference in stray rates between origins could be a genetic or environmental effect. Finally, a maximum recipient population stray rate of natural-origin fish into the Columbia River was less than $0.1 \%$ using genetic methods (Hess et al. 2014).

Data from the studies described above indicated that stray rates of natural-origin fish at various scales ranged between $0 \%$ and $100 \%$ but all but one estimate was below $30 \%$. Additional estimates of natural-origin stray rates would contribute to understanding the
magnitude of straying and the distribution of stray rates among species, populations, and environments. Knowing the magnitude of straying is important to understanding metapopulation dynamics, interpreting genetic data, informing scale of management units, and placing stray rates of hatchery origin fish into context (Keefer and Caudill 2014; Fullerton et al. 2016; Bett et al. 2017). Furthermore, discovering patterns related to natural-origin fish stray rates may contribute to identifying mechanisms associated with the variation in stray rates and also where fish may stray to. For example, adult salmon and steelhead have been shown to undershoot (Bond et al. 2017) and overshoot their natal area (Weigel et al. 2013; Richins and Skalski 2018) when they migrate home, in part because of access to cold water refugia.

In this paper, we provide estimates of donor natural-origin spring, summer, and fall Chinook Salmon and steelhead straying in the upper Columbia Watershed using PIT tags. The term of this type of straying is donor straying (Keefer and Caudill 2014). The upper Columbia watershed has one of the largest network of PIT tag antenna arrays in the United States which provides great opportunities to look at stray rates at a variety of scales. Three spatial scales of straying were evaluated: the upper Columbia basin, subbasins of the upper Columbia basin, and tributaries of upper Columbia subbasins (Figure 1; also see definition in Methods). These scales were selected because they were important homing targets for management, recovery, and understanding of population dynamics. We also looked for patterns in the data to identify whether there is a tendency for natural-origin spawners to stray in an upstream or downstream direction. We hypothesized that: 1) stray rates would increase as spatial scale decreased, 2) stray rates of steelhead would be higher than Chinook Salmon, and 3) stray rates would be similar in an upstream and downstream direction. We also hypothesized that stray rates would be towards the lower end of the range of stray rates that have been reported for natural origin Salmon and steelhead (0-100\%).


FIGURE 1. Release locations (green bullseye) and final PIT tag detection locations (yellow bullseye) of Chinook Salmon and steelhead originating from the upper Columbia River Basin. Other PIT tag detection sites are displayed as shaded dots for reference. Hydropower dams are denoted with triangles. The subbasins are the Okanogan, Methow, Entiat, and Wenatchee rivers and the Hanford Reach of the Columbia River. Collectively, these named subbasins represent the Upper Columbia Basin. Numbered tributaries indicate locations of straying individuals at the basin and tributary scales. The tributaries are (1) Little White Salmon River, (2) Deschutes River, (3) Snake River, (4) Peshastin Creek, (5) Nason Creek, (6) Little Wenatchee River, (7) White River, (8) Lost River.

## Methods

## Study Area

This study was conducted in the Columbia River watershed, USA, and most of the work was conducted in the upper Columbia Basin above the confluence with the Snake River (Figure 1). Three races of Chinook Salmon and one race of steelhead inhabit this area and are the focus of this study. Races are defined by the timing that they enter freshwater. Sockeye and Coho salmon also inhabit the upper Columbia, but there were insufficient numbers of natural-origin fish that were PIT tagged to include them in the analysis. Fall Chinook Salmon spawn in one of the few free flowing reaches of the Columbia River downstream of Priest Rapids Dam, are one of the largest Chinook Salmon populations in the United States, and contribute large numbers of fish to harvest in the Pacific Ocean and Columbia River, making this population economically very important (Harnish et al. 2014; Langshaw et al., 2017; Pearsons et al. in press). Summer Chinook Salmon spawn primarily in the mainstems of four subbasins of the upper Columbia River (e.g., Wenatchee, Entiat, Methow, and Okanogan) and support considerable fisheries in the Pacific Ocean and Columbia River. The naturally produced juveniles of summer and fall run Chinook Salmon migrate to the sea as sub-yearlings. Spring Chinook Salmon spawn in tributaries to mainstem subbasins and in upper portions of mainstem subbasins (Williamson et al. 2010; Murdoch et al. 2010; Ford et al. 2015a). The naturally produced juveniles of spring Chinook Salmon migrate to the sea as yearlings. They are listed under the Endangered Species Act as endangered (McClure et al. 2008). Summer steelhead spawn throughout subbasins and are listed as threatened (Ford et al. 2016). Naturally produced juvenile steelhead migrate to the sea at ages 1-7, but most migrate at ages 2 and 3 (Peven et al. 1994). All races of Chinook Salmon and steelhead have a long history of interactions with hatchery programs and hatcheryand natural-origin fish overlap in much of their spawning distributions (e.g., Williamson et al 2010; Pearsons et al. 2012; Ford et al. 2015a; Ford et al. 2016; Johnson et al. 2018).

Tagging and detection
Natural origin spring, summer, and fall Chinook Salmon and summer steelhead were PIT-tagged as juveniles in the upper Columbia River basin between 2002 and 2017. Chinook Salmon races and steelhead were only found, and later released, in portions of the upper Columbia River basin in which they historically spawn (See Methods: Study Area). Fish were collected with a variety of methods and for various purposes unrelated to straying. Fish were collected with rotary screw traps in subbasins and their tributaries, electrofishing in tributaries, fish bypasses at dams, and seining in the Columbia River (Johnson et al. 2007; Hillman et al. 2018). Fish were at least 50 mm FL when tagged (range 50 to 267 mm FL ) but less than $4 \%$ of fish were less than 60 mm FL to minimize potential effects of tag burden (Brown et al. 2010), and were released at the location of tagging or in the near vicinity. Fish were anesthetized and identified as natural-origin based upon absence of hatchery specific marks (e.g., adipose fin clip) and tags ((e.g., Coded Wire Tag (CWT)), the timing of collections (e.g., before hatchery fish are released), and the condition of fish (e.g., size, fin condition). Except for fall Chinook Salmon produced at Priest Rapids Hatchery, almost all of the hatchery-origin fish were tagged and/or marked. Tagging of natural origin fall Chinook Salmon in the Hanford Reach generally occurred prior to the release of hatchery origin fall Chinook Salmon in the Hanford Reach, and were also selected based upon size differences between hatchery and natural origin fish. PIT tags were 12
mm long, 2.1 mm diameter, and cylindrically shaped and were injected into the coelomic cavity of juveniles with syringes. In most cases, fish were allowed to recover before they were released. Short-term tag retention was generally high (e.g., >99\%) and mortality was low (e.g., <2\%) (Caisman 2018).

Anadromous adults with PIT tags were detected at a variety of antenna arrays in the Columbia River Basin between 2004 and 2018 (Figure 1). Antennas were able to read PIT tags in fish as they swam close enough to the antenna. Arrays were located in the fish ladders of many dams as well as the mouths of subbasins and their tributaries. Subbasin and tributary arrays were typically anchored to the bottom of rivers or streams. The efficiency of adult detections in most mainstem Columbia River dams was near 100\% (Pearsons et al. 2016). The efficiencies of subbasin and tributary arrays were less certain but likely varied with flow and fish migration behavior. Efficiencies were likely to be lower at high flows and when fish migrate high in the water column. Recent work suggest that efficiencies of subbasin and tributary arrays exceed $90 \%$ for steelhead (methods described by Connolly et al. 2008) and that stray estimates using CWT, that do not rely upon arrays, were similar to estimates using PIT tags for hatchery spring and fall Chinook Salmon (Grant County Public Utility District, unpublished data). Data from fish that passed arrays were uploaded to a centralized database.

## Analysis

The PIT Tag Information System (PTAGIS) maintained by the Pacific States Marine Fisheries Commission (PSMFC) was queried for adult salmon and steelhead returns to the Upper Columbia Basin. Individuals with known locations of tagging and release as juveniles were included in the analysis. Release quantities and detection records were used to create datasets for analysis. All detection records for natural-origin spring, summer, and fall Chinook Salmon and summer steelhead that were PIT-tagged as juveniles and originated from the Wenatchee, Entiat, Methow, and Okanogan River subbasins and the upper Columbia River were included in the analysis (Figure 1). Fish with last detections at hatcheries were excluded because these fish did not have an opportunity to self-correct and therefore inclusion of these detections would overestimate straying, however we only detected two fish with last detections at a hatchery so this rule was rarely implemented. Occurrence of straying was evaluated at three spatial scales; fish that originated from and returned to: (1) the upper Columbia River Basin (e.g., basin scale; all rivers and creeks above the confluence with the Snake River); (2) a subbasin within the Upper Columbia (e.g., subbasin scale; Wenatchee, Entiat, Methow, or Okanogan River subbasins and their tributaries; and the mainstem of the Columbia River); and (3) a tributary of a subbasin (e.g., tributary scale; Chiwawa River or Nason Creek, which are tributaries to the Wenatchee River).

A combination of time gaps and behavior, as determined by detection history, were used to exclude or include fish in the analyses. The time gap between release and final detection was used to generate a list of potential fish to include in the analysis. Chinook with at least 1.0 year and steelhead with at least 3 months between release and final detection were further evaluated to determine if the behavior of tagged individuals was consistent with that of anadromous salmonids. In this way, we attempted to eliminate fish that precociously matured and completed their life in freshwater (Pearsons et al. 2009). Detections of PIT tagged individuals in fish ladders at mainstem Columbia River dams were used to assess adult migration behavior. Fish detected at consecutive mainstem Columbia River dam fish ladders (i.e., Bonneville, McNary,
and Priest Rapids dams) were further evaluated to determine the occurrence of straying at the basin, subbasin, and tributary scales (Figure 1).

Fish that displayed behavior consistent with returning adults were further evaluated to determine final detection locations within the upper Columbia River. The occurrence of straying was determined using both brood year and return year for Chinook Salmon and return year only for steelhead. Brood year of spring Chinook was determined by tagging date within the calendar year. Fish tagged between January 1 and June 30 were classified as yearlings with brood year two years prior to tagging year. Fish tagged between July 1 and December 30 were classified as subyearlings with brood year one year prior to tagging year. This method aligned with trends observed in length of fish at tagging (Hillman et al. 2018). Fall Chinook were all collected and tagged in the upper Columbia River as subyearlings. Steelhead brood year was unknown because the age at migration was variable (e.g., 1 to 7 years) and length was not a good indicator of migration age because age-classes overlapped substantially (Peven et al. 1994). There were minor differences between stray estimates using brood year and return year (return year stray rates were minimally higher than brood year stray rates), however we present only return year results to allow comparison among all races of Chinook and between Chinook and steelhead.

We assumed that the last PIT detection in the database was the most likely spawning location. However, tagged individuals with final detections at mainstem Columbia River fish ladders were excluded from stray assignment at the subbasin and tributary scale, because it is unlikely that these fish spawned in the Columbia River. Fish with final detections within the subbasin where they were released, as determined by the river kilometer (RKM) of the subbasin, were assigned as homing to that subbasin. Fish with final detections in another subbasin in the upper Columbia River were assigned as straying to that subbasin. At the tributary scale, fish that originated from and had a final detection within a tributary were assigned as homing to that tributary. Fish with a final detection in another tributary of the same or different subbasin of origin were assigned as tributary strays. Only steelhead with final detections that corresponded with the spring spawning period (March through June) were included to exclude wandering behaviors from spawning behaviors.

Stray occurrence was calculated by summing the quantity of fish that strayed. The overall proportion of strays was calculated by dividing the stray total by the return total. Finally, the average stray occurrence was calculated by averaging the yearly stray occurrence when the quantity of returning fish was five or greater. Years with fewer than five returning fish were excluded from the calculation. We did not evaluate mechanisms of straying using mathematical models because of the low number of strays detected and because the main focus of this work was to document the magnitude of straying.

## Results

## Stray rate

A total of 823,770 PIT tags were injected into natural-origin fish and later evaluated to determine stray rates of natural-origin salmon and steelhead in the upper Columbia Watershed (Table 1). Despite a massive PIT tagging effort, the low survival rates between tagging of juveniles and returning adults resulted in low sample sizes for some years, species, and locations.

A total of 2,611 adults returned to the Columbia Basin and met our analytical criteria and were included in this analysis.

Table 1. Quantities (Qty) of PIT-tagged natural-origin Chinook Salmon and steelhead that homed to and strayed from the upper Columbia River basin, 2002-2018. Spring Chinook (SPC), summer Chinook (SUC), and steelhead (STH) that homed were detected at Priest Rapids or Rock Island dam fishways and locations upstream. Fall Chinook (FAC) that originated from the Hanford Reach of the Columbia River and were last detected at McNary or Priest Rapids dam fishways were assigned as home. Individuals assigned as strays were last detected outside the upper Columbia River. When more than one stray location is listed, the quantity of individuals is displayed in parentheses.

| Species/race | Qty PIT <br> Released | Qty <br> Home | Qty <br> Stray | Stray <br> rate | Stray Location |
| :--- | :---: | :---: | :---: | :---: | :--- |
| SPC | 352,109 | 1,000 | 0 | $0.0 \%$ |  |
| SUC | 100,273 | 98 | 0 | $0.0 \%$ |  |
| FAC | 140,114 | 286 | 2 | $0.7 \%$ | (1) Deschutes River, (1) Little |
|  |  |  |  |  | White Salmon River <br> STH |
| Total | 8231,274 | 1,223 | 2 | $0.2 \%$ | Snake River |
| Mean |  |  |  | $0.2 \%$ |  |

The mean stray rates of spring, summer, and fall Chinook Salmon and steelhead originating in the upper Columbia Basin were below $15 \%$ at all spatial scales. Stray rates were lowest at the basin scale and highest at the tributary scale. Mean stray rates of each population were less than $1 \%$ at the basin scale (range $0.0 \%-0.7 \%$, Table 1 ), less than $10 \%$ at the subbasin scale (range $0.0 \%-9.8 \%$, Table 2), and less than $15 \%$ at the tributary scale (range $0.0 \%-14.3 \%$, Table 3). Many of the populations that were evaluated across all spatial scales did not have any strays detected ( 11 of 28) and the mean of means of all species stray rates at all spatial scales was generally less than $5 \%$ (range $0.2 \%-4.0 \%$ ). Summer and fall Chinook Salmon were never detected straying into tributaries. Stray rates of Chinook Salmon and steelhead were similar when compared from the same subbasins and tributaries (Figure 2).

Table 2. Release, homing, and straying quantities (Qty) of PIT-tagged natural-origin spring Chinook (SPC), summer Chinook (SUC), fall Chinook (FAC), and steelhead (STH), originating from the upper Columbia River and its subbasins from 2002-2018. The mean stray rate excludes years with $<5$ homing adults. When more than one stray location is listed, the quantity of individuals is displayed in parentheses.

| Species/race | Qty PIT released | Qty <br> Home | $\begin{gathered} \text { Qty } \\ \text { Stray }^{1} \end{gathered}$ | Mean Stray <br> Rate | Stray Location |
| :---: | :---: | :---: | :---: | :---: | :---: |
| Columbia River |  |  |  |  |  |
| FAC ${ }^{2}$ | 140,114 | 286 | 2 | 0.7\% | (1) Deschutes, (1) Little White Salmon |
| Wenatchee |  |  |  |  |  |
| SPC | 230,770 | 497 | 4 | 1.2\% | (2) Entiat, (2) Methow |
| SUC | 476 | 0 | 0 | 0.0\% |  |
| STH | 58,960 | 241 | 2 | 0.5\% | Entiat |
| Entiat |  |  |  |  |  |
| SPC | 72,759 | 250 | 5 | 2.0\% | (1) Wenatchee, (2) Entiat, (2) Methow |
| SUC | 86,401 | 51 | 6 | 9.8\% | (1) Wenatchee, (4) Methow, (1) Okanogan |
| STH | 80,570 | 241 | 12 | 3.7\% | Methow |
| Methow |  |  |  |  |  |
| SPC | 48,580 | 67 | 3 | 5.2\% | (1) Wenatchee, (2) Okanogan |
| SUC | 6,676 | 2 | 0 | 0.0\% |  |
| STH | 73,773 | 175 | 9 | 5.3\% | (2) Snake, (7) Okanogan |
| Okanogan |  |  |  |  |  |
| SUC | 6,720 | 6 | 0 | 0.0\% |  |
| STH | 17,971 | 20 | 0 | 0.0\% |  |
| Total | 823,770 | 1,836 | 43 |  |  |
| Mean |  |  |  | 2.8\% |  |

${ }^{1}$ Strays were last detected outside the subbasins from which they originated.
${ }^{2}$ Fall Chinook were released into the Hanford Reach of the Columbia River and not into the Wenatchee, Entiat, Methow, or Okanogan rivers.


Figure 2. The percent of PIT-tagged natural-origin spring Chinook Salmon and steelhead that strayed away from their subbasins and tributaries of origin.

## Stray direction

The small number of fish that strayed at the subbasin and tributary scales generally strayed upstream of their capture location. It was not possible for fish to stray upstream of the basin scale because there is no basin above the upper Columbia for fish to stray into. At the basin scale, only 4 fish strayed ( 2 steelhead and 2 fall Chinook), and all of them strayed into locations downstream of the upper Columbia River (Table 1). Two steelhead strayed into the Snake River and two Fall Chinook Salmon were detected in subbasins well downstream of Priest Rapids Dam (Deschutes River and the Little White Salmon River). Fall Chinook Salmon originating in the Hanford Reach below Priest Rapids dam were not detected in upper Columbia River subbasins.

At the subbasin scale, spring, summer, and fall Chinook Salmon and steelhead strays were generally detected in subbasins upstream of the home subbasin, however, there were instances of straying to a downstream subbasin within the upper Columbia (e.g., a spring Chinook Salmon that originated from the Entiat River but returned to the Wenatchee River). Of the 43 salmon and steelhead that strayed, $84 \%$ (36) were last detected in a subbasin upstream of home (Tables 2). One hundred percent (4 of 4) of spring Chinook Salmon from the Wenatchee subbasin, $80 \%$ from the Entiat subbasin (4 of 5), and $67 \%$ ( 2 of 3 ) from the Methow subbasin strayed upstream Eighty-three percent (5 of 6) of summer Chinook Salmon from the Entiat River strayed upstream. One hundred percent (2 of 2) of steelhead from the Wenatchee subbasin,
$100 \%$ (12 of 12) from the Entiat subbasin, and $78 \%$ (7 of 9) from the Methow subbasin strayed upstream. One hundred percent (2 of 2) of fall Chinook strayed downstream. At the tributary scale, $94 \%$ of spring Chinook Salmon and Steelhead strayed upstream. Only 9 spring Chinook Salmon strayed and 8 of them strayed to an upstream tributary ( $89 \%$ ) while $100 \%$ ( 9 of 9) steelhead strayed upstream (Table 3). Despite the tendency for Salmon and steelhead to stray upstream, the stray rates of fish originating from locations upstream (e.g., Methow subbasin) appeared higher than those originating from downstream locations (e.g., Wenatchee subbasin; Figure 2).

Table 3. Quantities (Qty) of PIT-tagged natural-origin spring Chinook Salmon (SPC) and steelhead (STH) originating from upper Columbia River subbasins (Wenatchee $=\mathrm{W}$, Entiat $=\mathrm{E}$, Methow $=\mathrm{M}$, Okanogan $=\mathrm{O}$ ) with homing and straying totals at the tributary scale 2002-2018. The mean stray rate excludes populations with < 5 homing adults. When more than one stray location is listed, the quantity of individuals is displayed in parentheses.

| Species/race | Tributary | Qty PIT Released | Qty <br> Home | $\begin{aligned} & \text { Qty } \\ & \text { Stray } \end{aligned}$ | Stray Rate | Stray Location |
| :---: | :---: | :---: | :---: | :---: | :---: | :---: |
| SPC | Chiwawa [W] | 167,953 | 216 | 5 | 2.3\% | (2) Little Wenatchee [W], (1) Nason Cr [W], <br> (1) Peshastin Cr [W], (1) White River [W] |
| SPC | Nason [W] | 26,656 | 42 | 3 | 6.7\% | (1) Little Wenatchee [W], (1) White River [W], (1) Twisp River [M] |
| SPC | White [W] | 3,275 | 2 | 0 | 0.0\% |  |
| SPC | Twisp [M] | 23,391 | 31 | 1 | 3.1\% | Lost River [M] |
| SPC | Chewuch [M] | 11,425 | 16 | 0 | 0.0\% |  |
| STH | Nason [W] | 15,808 | 21 | 0 | 0.0\% |  |
| STH | Chiwawa [W] | 15,065 | 25 | 0 | 0.0\% |  |
| STH | Mad [E] | 9,538 | 16 | 1 | 5.9\% | Libby Creek [M] |
| STH | Chewuch [M] | 9,672 | 17 | 1 | 5.6\% | Salmon Creek [O] |
| STH | Beaver/Gold/ <br> Libby [M] | 14,284 | 18 | 3 | 14.3\% | Twisp River [M] |
| STH | Twisp [M] | 28,075 | 61 | 4 | 6.2\% | (1) Loup Loup Creek <br> [O], (1) Bonaparte Cr <br> [O], (1) Tunk Cr [O], (1) <br> Hancock Springs [M] |
| STH | Omak [O] | 10,462 | 13 | 0 | 0.0\% |  |
| Total |  | 335,604 | 478 | 18 |  |  |
| Mean |  |  |  |  | 4.0\% |  |

## Discussion

Our results indicated that mean stray rates of natural-origin Chinook Salmon and steelhead were below $15 \%$ at all three spatial scales and were at the low end of estimates that were previously published for natural-origin steelhead and spring Chinook Salmon (Shapovalov and Taft 1954; Ford et al. 2015a). Stray rates of natural-origin PIT tagged spring Chinook Salmon in the upper Wenatchee Basin were about $56-74 \%$ of those reported using genetic techniques in the same tributaries (Ford et al. 2015a). For instance, stray rates for spring Chinook Salmon originating from the Chiwawa River were $2.3 \%$ using PIT tags and $4.1 \%$ using genetic techniques. Furthermore, stray rates for spring Chinook Salmon originating from Nason Creek were $6.7 \%$ using PIT tags and $9.0 \%$ using genetic techniques. These differences may be within sample size and measurement error or be due to differences in the years included in the different studies. Alternatively, it is possible that the efficiency of the PIT antenna arrays was less than $100 \%$ and our methodology underestimated straying. However, recent work suggests that efficiencies of subbasin and tributary arrays exceed $90 \%$ for steelhead and that stray estimates using CWT, that do not rely upon arrays, were similar to estimates using PIT tags for hatchery-origin spring Chinook Salmon (Grant County Public Utility District, unpublished data). There is also a possibility of overestimating strays using the method of last PIT tag detections. This could occur if fish temporarily stray or wander (e.g. Bond et al. 2017; Richins and Skalski 2018) and then are not detected at a different antenna. Preliminary information from comparisons of hatchery-origin summer Chinook Salmon stray rates derived from CWT and PIT tags suggested PIT tag estimates were correlated with, but higher than CWT estimates (Grant County Public Utility District, unpublished data). This suggests that natural-origin stray rates of summer Chinook Salmon at the subbasin and tributary scales may be overestimates.

Unfortunately, we could not make comparisons to spring Chinook Salmon spawning populations with high stray rates reported by Ford et al. (2015a) ( $100 \%$ for fish originating from the upper Wenatchee River and $17.5 \%$ for fish originating from the Little Wenatchee River) because we didn't have sufficient PIT tags from those locations. However, PIT tag estimates for spring Chinook Salmon in five upper Columbia tributaries were substantially lower than these high stray rates (e.g., $<7 \%$ ). Estimating stray rates of small populations will likely be a challenge in the future, particularly using methods such as we described in this work. Another alternative method to estimate straying is to evaluate otolith chemistry in cases where water chemistry is sufficiently different (Brenner et al. 2012; Budnik et al. 2018; Watson et al., 2018). Differences in water chemistry signatures have been found in tributaries of the upper Wenatchee and there was ability to discriminate juvenile spring Chinook Salmon that resided in tributaries prior to migration as yearlings using chemical differences in fin rays (Linley et al. 2016). Thus, it may be possible to evaluate straying using fin rays or otoliths, but different emigration times of juveniles from tributaries may decrease discrimination of adults (Linley et al. 2016) and decrease the utility of stray estimates using this method.

The stray rates of natural-origin fish that we report may be higher than what occurred prior to habitat degradation and the large inputs of hatchery-origin fish (see descriptions in Williamson et al. 2010; Ford et al. 2015a; Johnson et al. 2018). Ford et al. (2015a) found that natural born offspring of spring Chinook Salmon with hatchery-origin parentage had higher stray rates than those from natural-origin parents. None of the natural born fish from natural-origin spring Chinook Salmon were detected as strays in that study. The natural-origin juveniles from our study were likely produced from a variety of matings of both hatchery and natural-origin
parents which may have increased the stray rate when compared to systems without hatcheryorigin spawners. In addition, it has been speculated that degraded spawning habitat has contributed to increased stray rates (Ford et al. 2015a, Cram et al. 2012) and there has been habitat degradation in the upper Columbia Basin such as passage impediments, warming water temperature, and stream channelization. Furthermore, management actions that disrupt sequential imprinting or homing, such as barging or routing of water through irrigation canals and tributaries, can also increase straying (Keefer and Caudill 2014; Bond et al. 2017).

Stray rates were different depending upon the spatial scale of evaluation. Mean stray rates of each population were less than $1 \%$ at the basin scale, less than $10 \%$ at the subbasin scale, and less than $15 \%$ at the tributary scale. These findings highlight the importance of spatial scale in evaluations and the necessity of defining spatial scales when making comparisons and communicating results (Keefer and Caudill 2014). We could not generate a good estimate of stray rates at the Columbia River Basin scale because of insufficient PIT detection in other Basins. However, estimates of natural-origin strays into the Columbia River suggests that straying between large river Basins may be low (Hess et al. 2014) such as we found at the largest spatial scale we examined in this study. Many studies have evaluated straying of hatchery-origin fish at the subbasin and larger scales (Westley et al. 2013, 2015, Bond et al. 2017). Ford et al. (2015a) presented stray rate information at a finer spatial scale (e.g., within tributaries) than this study using genetic methods; something we could not do with the PIT tag methods that were used in this study.

Other studies may detect different patterns of stray rates depending upon the dendricity and spatial positioning of spawning habitats. It is also possible that the magnitude of naturalorigin fish straying could differ depending upon differences in hatchery-origin fish abundance and spawning success, habitat degradation, barging, and water withdrawals. Hatchery-origin fall Chinook Salmon that were collected in the Snake River and barged downstream strayed at higher rates than those that were not barged (Bond et al. 2017). Similarly, the likelihood of straying increased during years of warmer river temperatures. If natural origin fish encounter these conditions, then it is likely that they would stray at higher rates than what we presented for the upper Columbia basin.

Our results do not support the reputation that steelhead have for high straying propensity (Richins and Skalski 2018, Budnik et al. 2018). The mean stray rates at all scales were relatively low and Chinook Salmon strayed at similar rates as steelhead at the scales that we examined. Furthermore, in another study Coho Salmon had dramatically higher rates of straying than steelhead in two coastal California streams (Shapovalov and Taft 1954). Perhaps steelhead have received their reputation for straying based upon their wandering behavior before spawning and because most of what is known about steelhead straying comes largely from hatchery-origin fish (Richins and Skalski 2018, Budnik et al. 2018). However, Westley et al. (2013) reported that hatchery Chinook Salmon strayed more than hatchery steelhead. The differences in straying that occurs among species may differ between regions depending upon the myriad of factors that influence straying, such as imprinting, hatchery influence, barriers to migration, water temperature, irrigation routing, and spawning habitat conditions (Keefer and Caudill 2014; Cram et al. 2012), and the relative frequency of those factors in the different regions. For instance, steelhead may stray more than Chinook Salmon in some regions but not in others.

Directionality

Most of the spring and summer Chinook Salmon and steelhead strays strayed in an upstream direction. This is interesting because the opportunities for straying in a downstream direction were much higher than for straying in an upstream direction. The further upstream a fish migrates the fewer opportunities it has to stray in an upstream direction. Salmon and steelhead pass many subbasins and tributaries as they migrate up the Columbia River and yet they tend to stray upstream of their natal rearing area. This may be a result of sequential imprinting errors (Dittman et al. 2015) or an adaptation to colonize new upstream habitats such as when glaciers retreat, volcanic eruptions cease, flood waters recede, or migration barriers are removed (Leider 1989; Pearsons et al. 1992; Weigel et al. 2013). For some species that migrate when water temperatures are relatively warm, such as steelhead and fall Chinook, fish may overshoot (Richins and Skalski 2018) or undershoot (Bond et al. 2017) natal areas in search of cold water refugia. As such, there are likely multiple factors that influence the direction of straying and the stray direction may be different in other locations outside the upper Columbia basin.

Management implications
The low stray rates that we observed in this study are consistent with the development of genetic differentiation among populations at various spatial scales in the upper Columbia Basin (McClure et al. 2008). However, even low stray rates can result in significant interbreeding with non-target populations and result in increased homogenization of spawning populations (Bett et al. 2017). This is particularly true: (1) when the donor populations are large, (2) when donor straying is frequent, and (3) when the recipient population is small (Bett et al. 2017).
Furthermore, hatchery programs can disrupt patterns of natural-origin stray rates and decrease genetic differentiation (Ford et al. 2015a, b; Ford et al. 2016). This study focused on donor stray rates, but estimates of recipient population stray rate are more relevant when evaluating potential genetic effects on natural spawning populations and yet estimates of recipient population stray rate are relatively rare (Keefer and Caudill 2014). Until recipient population stray rates are available at multiple spatial scales, managers can use donor population stray rates to help inform management actions.

Estimates of natural-origin fish stray rates, such as those in this study, could be used to inform management targets for hatchery programs. However, the variation in donor population stray rates that have been observed for natural-origin salmonids has been highly variable ranging from $0-100 \%$ and can vary between species, geographic location, environmental condition, and spatial scale (Shapovalov and Taft 1954; Ford et al. 2015a). Some authors have suggested that universal management targets for donor strays are not appropriate (Quinn 2005; Brenner et al. 2012; Keefer and Caudill 2014). In contrast, recipient population stray compositions have been recommended based upon genetic and ecological risk toleration and have ranged between $2 \%$ 10\% (Ford 2002; Mobrand et al. 2005; Brenner et al. 2012; Paquet et al. 2011; Hillman et al. 2018). It is likely that more information is necessary before donor population stray rate targets can be set and that site specific information will be needed to inform management targets. In addition, the objectives of a hatchery program will influence what donor stray rate targets are appropriate. For example, in cases of large-scale reintroduction, such as above Chief Joseph and Grand Coulee dams (Johnson et al. 2018), high stray rates may be desirable in order to colonize large areas. Furthermore, managers should consider whether estimates of donor stray rate targets of natural-origin fish are realistic to achieve for hatchery-origin fish that are cultured under dramatically different conditions. It remains to be seen whether natural- and hatchery-origin fish
stray rates differ at a variety of spatial scales and in different regions, however Ford et al. (2015a) indicated that stray rates of hatchery-origin spring Chinook Salmon were higher than natural-origin spring Chinook Salmon in the Chiwawa River.

The tendency for natural-origin fish to stray in upstream directions can be used to predict what groups of fish are likely to populate newly created habitats within subbasins and tributaries and also be candidates for reintroduction. Newly created habitats include removal of passage impediments such as culverts and also include locations exposed to floods, droughts, volcanic eruptions, and other disturbances (Pearsons et al. 1992, Leider 1989; Weigel et al. 2013). Selecting candidate populations for reintroduction, such as above Chief Joseph and Grand Coulee dams, might also be informed based upon what populations would likely colonize the area naturally. Natural-origin fish that stray might have some traits that make them particularly suitable for colonizing new habitats, although we are not aware of data that supports this idea.

## Conclusion

We demonstrated that PIT tags can be an effective means to estimate the magnitude of natural-origin salmon and steelhead straying and can also be used to evaluate factors associated with straying. Unfortunately, massive efforts for PIT tagging and deployment of antenna arrays are necessary to generate estimates. One weakness of using PIT tags to estimate straying is there is no confirmation that a fish spawned within the area that it was last detected. We found that stray rates of natural-origin spring, summer, and fall Chinook Salmon and steelhead at three spatial scales were less than $15 \%$ and there was variation in stray rates between spatial scales. Furthermore, most of the fish that strayed into non-natal subbasins and tributaries strayed in an upstream direction. There continues to be a lack of studies that have evaluated stray rates of natural-origin fish, and further work would contribute to our understanding of the magnitude of straying by different populations in a variety of different habitats.

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# Comparisons of Donor Stray Percentages Between Hatchery- and Natural-Origin Chinook Salmon and Steelhead in the Upper Columbia Watershed 

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#### Abstract

Artificial propagation of salmon Oncorhynchus spp. and steelhead $O$. mykiss is a common strategy that is used to achieve conservation and harvest goals. However, unintended effects of artificial propagation, such as high donor stray percentages, can reduce the number of adults that return to target areas and also contribute spawners to different populations where they are not desired. Until recently, it was difficult to assess if hatchery-origin fish stray rates were atypical because few estimates of stray rates of natural-origin fish were available. We used last PIT-tag detections to estimate and compare donor stray percentages of hatchery-origin and natural-origin Chinook Salmon O. tshawytscha and steelhead in the upper Columbia River watershed between 2002-2018. Donor stray percentages of hatchery-origin spring, summer, and fall Chinook Salmon and steelhead were $<0.3 \%$ at the upper-Columbia basin scale and generally not higher than natural-origin donor stray percentages at larger spatial scales but were higher (up to 62\%) at smaller spatial scales. Returning hatchery-origin Chinook Salmon and steelhead generally strayed in an upstream direction and the proportions of fish that strayed upstream were not significantly higher than natural-origin fish. Juvenile spring Chinook Salmon that were moved 14 to 389 river kilometers from centralized hatcheries to tributaries for overwintering or final acclimation, strayed at a much higher rate than those that completed their incubation, rearing, and acclimation at a single location. In contrast, steelhead that were moved for acclimation, including direct releases from trucks, did not stray at higher rates than those that completed their incubation, rearing, and acclimation at a single location. Other adaptive management actions that were implemented to reduce straying produced mixed results. A variety of approaches can be considered to reduce undesirable production of strays, but most of them involve difficult trade-offs.


## Introduction

Hatcheries are frequently used to increase harvest and conserve natural populations of salmon and steelhead but the large-scale production of salmon and steelhead in hatcheries poses a variety of unintended ecological and genetic risks to natural-origin populations (Busack and Currens 1995; Pearsons 2008; Pearsons et al. 2012) and straying is among the most significant concerns (Ford 2002; Mobrand et al. 2005; Paquet et al. 2011). Unusually high incidence of strays from hatchery programs are undesirable for a number of reasons. First, stray fish do not come back to the intended target area and therefore are not available for location specific harvest or conservation purposes (Keefer and Caudill 2014; Sturrock et al. 2019). Second, hatcheryorigin strays that spawn with other recipient populations, may reduce genetic diversity among natural-origin populations (Quinn 2005; Mobrand et al. 2005; Brenner et al. 2012). Straying can be estimated as either the percentage of a source spawning population that strays (i.e., donor stray percentage) or the percentage of a recipient spawning population that is composed of nonnatal spawners (i.e., recipient stray percentage) (Keefer and Caudill 2014). Stray fish that spawn with non-target populations can pose risks to both donor and recipient populations. The spatial scale of straying is also an important consideration (Keefer and Caudill 2014; Pearsons and O'Connor 2020) because long-distance straying is likely to pose more undesirable risks to harvest and conservation objectives than short-distance straying.

Salmon and steelhead are hypothesized to home by sequentially imprinting as juveniles and then following imprinted cues in reverse when returning as adults (Hasler and Scholz 1983; Dittman et al. 2010; 2015). Other factors such as habitat quality, pheromones of conspecifics, and geographic complexity can influence homing, particularly at finer scales (Cram et al. 2012; Keefer and Caudill 2014; Bett et al. 2017). Much uncertainty remains about how hatchery practices influence homing and straying, but some hatchery practices are generally thought to increase straying compared to naturally produced fish (Keefer and Caudill 2014) and achieving acceptably high homing is one of the greatest challenges for fish culturists (Westley et al. 2013; 2015; Ford et al. 2015a).

A variety of fish-husbandry methods are currently used to reduce straying of hatcheryorigin fish and to return fish to target areas. For example, acclimation sites are used to imprint juvenile fish on surface water in specific areas prior to release in the hopes that they will return to the target area around the acclimation site (Dittman et al. 2010; Clarke et al. 2012; Keefer and Caudill 2014). The length of time that fish are acclimated can vary from a few weeks in the spring to over six months spanning the winter for yearling smolt programs (Dittman et al. 2010; Clarke et al. 2012; Ford et al. 2015a). Also, fish are generally released when they are undergoing smoltification, the time that fish have a very strong spike in the hormone thyroxine, which is thought to be associated with chemical imprinting (Scholz 1980; Hasler and Scholz 1983; Westley et al. 2013). Embryonic imprinting, where fish are exposed to natal water at the alevin to fry life stages, has been proposed for hatchery programs that incubate eggs and embryos at locations far from release locations (Dittman et al. 2015). Although embryonic imprinting has not been evaluated in cases where fish are transported prior to release, it does occur in locations where all life-stages are raised and released at the same location, however the water is often local ground water instead of surface water in order to reduce disease risk.

Most of what is known about salmon and steelhead straying is derived from studies of hatchery-origin fish (Westley et al. 2013, 2015; Keefer and Caudill 2014). It has been difficult to determine whether hatchery-origin fish stray rates are unusually high or low when compared
to natural-origin fish because natural-origin fish stray rate estimates were not available from the same area where hatchery-origin fish are released, and because observed natural-origin stray rates have been highly variable, ranging between 0 and 100\% (Shapovalov and Taft 1954; Ford et al. 2015a; Keefer and Caudill 2014). Recently, estimates of natural-origin stray rates have been developed using genetic (Ford et al. 2015a) and passive integrated transponder tag (PIT tag) (Pearsons and O'Connor, 2020) methods. Mean donor stray percentages for natural-origin Chinook Salmon and steelhead in the Upper Columbia watershed were less than $1 \%$ at the upper Columbia basin scale, less than $10 \%$ at the subbasin scale, and less than $15 \%$ at the tributary scale (Pearsons and O’Connor, 2020). Most of the populations that were evaluated across all spatial scales did not have any strays detected. Chinook Salmon strayed at higher rates than steelhead. Straying mostly occurred in an upstream direction at both the subbasin and tributary scales. The directionality of straying is important because it provides information about which recipient populations are likely to be affected by strays as well as what new habitats may be colonized by strays.

In this paper, we used similar methods to estimate donor stray percentages of hatcheryorigin salmon and steelhead in the upper Columbia watershed as we did to estimate donor stray percentages of natural-origin salmon and steelhead in the same area (Pearsons and O'Connor 2020), and made comparisons between natural-origin and hatchery-origin donor stray percentages. We focused our efforts on 'permanent strays' as opposed to adult wandering prior to spawning (Keefer and Caudill 2014) and also focused on 'management strays' which was defined as adults that did not return to spawn near the juvenile release location. We formed hypotheses that were informed by what we observed in natural-origin adults in the upper Columbia watershed as well as previously published information about straying by hatcheryorigin adults (Pearsons and O’Connor 2020; Keefer and Caudill 2014). We hypothesized that: 1) donor stray percentages of hatchery-origin fish would increase with decreasing spatial scale similar to the pattern we observed for natural-origin fish (Pearsons and O’Connor 2020), 2) donor stray percentages of hatchery-origin fish would be higher than donor stray percentages of natural-origin fish, particularly at smaller spatial scales such as was suggested by other published studies (Keefer and Caudill 2014), 3) hatchery-origin fish stray direction would depend upon release location such as would be supported by the sequential imprinting hypothesis (Keefer and Caudill 2014), and 4) donor stray percentages would decline after management actions intended to reduce straying were implemented. We also evaluated the quality of PIT-tag-based stray estimates by comparing them to estimates generated using coded-wire tags (CWT).

## Methods

## Study Area

This study was conducted in the Columbia River, USA and most of the work was conducted in the upper Columbia watershed upstream of the confluence with the Snake River and downstream of Chief Joseph Dam, with fish from hatchery programs in the Wenatchee, Entiat, Methow, and Okanogan River subbasins and the upper Columbia River (Figure 1). The upper Columbia River watershed has an abundance of hatchery facilities as a result of mitigation for the construction and operation of hydropower dams (Figure 1). These hatcheries produce fall, summer, or spring Chinook Salmon, Coho Salmon O. kisutch, Sockeye Salmon O. nerka,


Figure 1. Release locations (green bullseye) and final detection locations (orange bullseye) of spring, summer, and fall Chinook Salmon and steelhead from the upper Columbia River Basin. Other points along rivers indicate PIT tag detection arrays. Numbered boxes represent locations of subject hatcheries including: (1) Little White Salmon, (2) Priest Rapids, (3) Eastbank, (4) Leavenworth, (5) Entiat, (6) Chelan, (7) Wells, (8) Chief Joseph, (9) Cassimer Bar, (10) Methow, and (11) Winthrop.
and steelhead for harvest, conservation, or a combination of both; but Chinook Salmon and steelhead are the only species considered here (Table 1). There were insufficient numbers of natural-origin Sockeye and Coho salmon that were PIT tagged to include these species in this comparative analysis. Some of the hatchery programs incubate, rear, and release fish from a single hatchery location, whereas other programs transport parr or smolts to acclimation sites for subsequent release (Table 1). The study area and biological background was previously described by Pearsons and O'Connor (2020) and is also briefly described below.

Fall Chinook Salmon spawn in the Hanford Reach, one of the few free-flowing reaches of the Columbia River downstream of Priest Rapids Dam, comprising one of the largest Chinook Salmon populations in the United States, and contribute large numbers of fish to harvest in the Pacific Ocean and Columbia River, making this population economically very important (Harnish et al. 2014; Langshaw et al. 2017; Pearsons et al. 2020). Summer Chinook Salmon spawn primarily in the mainstems of four subbasins of the upper Columbia River (e.g., Wenatchee, Entiat, Methow, and Okanogan) and support considerable fisheries in the Pacific Ocean and Columbia River. The naturally produced juveniles of summer and fall run Chinook Salmon generally migrate to the sea as sub-yearlings. Spring Chinook Salmon spawn in tributaries to mainstem subbasins and in upper portions of mainstem subbasins (Williamson et al. 2010; Murdoch et al. 2010; Ford et al. 2015a). Upper Columbia River spring Chinook Salmon are listed under the Endangered Species Act (ESA) as endangered (McClure et al. 2008). The naturally produced juveniles of spring Chinook Salmon migrate to the sea as yearlings. Summer steelhead spawn throughout upper Columbia subbasins and are ESA listed as threatened (Ford et al. 2016). Naturally produced juvenile steelhead migrate to the sea at ages 1-7, but most migrate at ages 1, 2 and 3 (Peven et al. 1994). All life history types of Chinook Salmon and steelhead have a long history of interactions with hatchery programs and hatchery- and natural-origin fish overlap in much of their spawning distributions (e.g., Williamson et al 2010; Pearsons et al. 2012; Ford et al. 2015a; Ford et al. 2016; Johnson et al. 2018).

## Analytical Framework and Definitions

We used information from PIT tags and PIT-tag detection arrays deployed throughout the region for various purposes to evaluate donor stray percentages of hatchery-origin salmon and steelhead. The analytical methods and years used for these analyses were similar to those described for estimation of natural-origin donor stray percentages in the same geographic area of Pearsons and O'Connor (2020). We assumed that the last PIT detection in the database was the most likely spawning location. However, tagged individuals with final detections at mainstem Columbia River fish ladders were excluded from stray assignment at the subbasin and tributary scale, because it is unlikely that these fish spawned in the Columbia River, except fall Chinook Salmon in the Hanford Reach of the Columbia River. Fish with final detections within the subbasin where they were released, were assigned as homing to that subbasin. Fish with final detections in another subbasin in the upper Columbia River were assigned as straying to that subbasin. At the tributary scale, fish that originated from and had a final detection within a tributary were assigned as homing to that tributary. Fish with a final detection in another tributary of the same or different subbasin of origin were assigned as tributary strays. Only steelhead with final detections that corresponded with the spring spawning period (March through June) were included to exclude wandering behaviors from spawning behaviors.

Wandering behaviors included temporary residency in a subbasin or tributary during migration or overwinter periods. Final detections that aligned with spawning periods were assumed to be

Table 1. Locations of hatchery activities and PIT tag quantities (Qty) for hatchery programs in the upper Columbia Basin. All fish were released as yearlings except for fall Chinook Salmon and some summer Chinook Salmon which were released as subyearlings into the Okanogan and Columbia rivers. PIT-tagged juvenile summer Chinook Salmon reared at Wells Hatchery and released into the Methow and Okanogan rivers in 2010 for survival studies were included in basin-scale analyses but not for subbasin stray results.

| Incubation and Rearing | Final Acclimation | Release | Years of release | Quantities (Qty) of PIT-tagged juvenile Chinook Salmon and steelhead |
| :---: | :---: | :---: | :---: | :---: |
| Spring Chinook Salmon |  |  |  |  |
| Eastbank | Nason | Nason Creek | 2015-2017 | 35,243 |
| Eastbank | Chiwawa | Chiwawa River | 2007-2017 | 99,940 |
| Little White | White River and | White River, | 2008-2015 | 277,729 |
| Salmon | Lake Wenatchee | Lake |  |  |
|  |  | Wenatchee, Wenatchee |  |  |
|  |  | River |  |  |
| Leavenworth | Leavenworth | Icicle Creek | 2000-2017 | 995,661 |
| Methow | Twisp | Twisp River | 2004, 2012- | 40,503 |
|  |  |  | 2017 |  |

## Summer Chinook Salmon

| Eastbank | Dryden | Wenatchee River | 2007-2017 | 126,765 |
| :---: | :---: | :---: | :---: | :---: |
| Eastbank | Carlton | Methow River | 2007-2017 | 34,740 |
| Eastbank | Similkameen | Similkameen <br> River | 2011, 2013 | 10,125 |
| Entiat | Entiat | Entiat River | 2010-2017 | 89,710 |
| Wells | Wells | Columbia River | 2000-2017 | 152,400 |
| Wells | Wells | Methow River | 2010 | 30,343 |
| Wells | Wells | Okanogan River | 2000, 2010 | 11,030 |
| Chief Joseph | Omak | Similkameen and Okanogan rivers | 2015-2017 | 24,718 |
| Chief Joseph | Chief Joseph | Columbia River | 2015-2017 | 29,971 |
| Fall Chinook Salmon |  |  |  |  |
| Priest Rapids | Priest Rapids | Columbia River | 2000-2017 | 357,808 |
|  |  | Steelhead trout |  |  |


| Eastbank and Chelan | Turtle Rock | Chiwawa River, <br> Nason Creek, <br> Wenatchee <br> River | $\begin{gathered} 2005,2009- \\ 2011 \end{gathered}$ | 235,451 |
| :---: | :---: | :---: | :---: | :---: |
| Eastbank and Chelan | Chiwawa | Chiwawa River, <br> Nason Creek, <br> Wenatchee <br> River | $\begin{gathered} 2003,2005 \\ 2009,2011, \\ 2012-2017 \end{gathered}$ | 118,507 |
| Eastbank and Chelan | Turtle Rock (Columbia River) and Chiwawa | Various throughout Wenatchee River subbasin | $\begin{aligned} & \text { 2003-2005, } \\ & 2007-2017 \end{aligned}$ | 314,077 |
| Eastbank and Chelan | Blackbird Island | Wenatchee River | 2010-2016 | 20,769 |
| Eastbank | Nason (Rolfing) | Wenatchee River | 2010 | 20,211 |
| Wells | Wells | Columbia River | $\begin{gathered} 2000,2003 \\ 2012-2017 \end{gathered}$ | 161,954 |
| Wells | Twisp | Twisp River | $\begin{aligned} & 2003-2005, \\ & 2010-2017 \end{aligned}$ | 198,334 |
| Wells | Methow <br> Hatchery | Methow River | $\begin{aligned} & 2003-2005, \\ & 2010-2017 \end{aligned}$ | 275,839 |
| Wells | Chewuch | Chewuch River | $\begin{aligned} & 2003-2005, \\ & 2010-2011 \end{aligned}$ | 123,312 |
| Winthrop | Winthrop | Methow River | $\begin{aligned} & \text { 2003-2005, } \\ & 2008-2017 \end{aligned}$ | 380,202 |
| Winthrop | Winthrop | Chewuch River | 2010, 2012 | 996 |
| Cassimer Bar | Cassimer Bar | Omak Creek | $\begin{gathered} 2005,2007- \\ 2011 \end{gathered}$ | 88,332 |
| Cassimer Bar | Cassimer Bar | Stapaloop Creek | 2004, 2006 | 23,334 |
| Wells | Saint Mary's | Omak Creek | $\begin{aligned} & 2003-2005, \\ & 2012-2017 \end{aligned}$ | 90,249 |
| Wells | Wells | Salmon Creek | 2012, 2017 | 11,310 |
| Wells | Wells | Similkameen River | $\begin{aligned} & 2003-2005, \\ & 2012,2017 \end{aligned}$ | 93,613 |
| Total |  |  |  | 4,379,563 |

spawning fish. The donor stray percentages of natural-origin fish presented previously were used for comparisons to hatchery-origin fish (Pearsons and O’Connor 2020).

We defined donor straying as a fish that did not return to the location of release, which was the management intent of acclimation or location of release. Furthermore, we were interested in permanent rather than temporary straying, which is why we use last PIT detections in our evaluation. However, adults that returned to a hatchery or adjacent location where
juveniles had earlier rearing experience such as during embryonic development may have homed correctly, but were not consistent with the management objective. We did not include fish that were detected at hatcheries in this evaluation because they did not have the opportunity to escape once they entered a facility, facilities were not always equipped with a PIT detector, and fish were not always scanned for PIT tags at hatcheries.

A representative sample of fish were PIT tagged (typically 5,000-10,000 annually) at central hatcheries or acclimation sites between 2000 and 2017 and allowed to recover prior to release (Table 1). The timing of tagging varied depending upon the size of fish and the objective of the tagging. In general, fish were tagged in the fall or spring prior to release. Fish were PIT tagged when they were at least 60 mm FL and were anesthetized prior to tagging. The PIT tags were Biomark ${ }^{\text {TM }}$ model, 12 mm long, 2.1 mm diameter, and cylindrically shaped and were injected into the coelomic cavity of juveniles with syringes. Short-term tag retention was generally high (e.g., >99\%) and mortality was low (e.g., <2\%) (Hillman et al. 2019).

Two major hatchery management modifications to fish acclimation occurred during this study to reduce straying. We compared the donor stray percentages of fish before and during the modification to determine whether the modification reduced straying. The expectation was that the donor stray percentages would decrease substantially after the management action was implemented. First, we evaluated whether a new overwinter acclimation facility decreased summer Chinook Salmon donor stray percentages when compared to spring acclimation at the same site. It was hypothesized that longer periods of acclimation may improve imprinting and homing. Summer Chinook Salmon were raised at Eastbank Hatchery on the Columbia River and then transferred to the Carlton acclimation site in the Methow River subbasin in the spring for final acclimation and release in 2010 and 2011. A new overwinter acclimation facility was subsequently built on the same property with the first release in 2014. The fish released in 2014 were spring acclimated, but from 2015 through 2017 fish were overwinter acclimated. We compared donor stray percentages of summer Chinook Salmon that were spring acclimated (2010, 2011, 2014) and overwinter acclimated (2015-2017). Second, a change in hatchery and acclimation facilities for steelhead from a) Turtle Rock Hatchery on the Columbia River and using trucks to plant steelhead throughout the Wenatchee River subbasin (release years 20062008) to b) Eastbank hatchery and an overwinter acclimation facility and release on the Chiwawa River in the Wenatchee River subbasin (release years 2014, 2016, 2017). This change increased exposure to water from the Wenatchee River subbasin, where fish were targeted to return.

## Analysis

The PIT Tag Information System (PTAGIS) maintained by the Pacific States Marine Fisheries Commission (PSMFC) was queried for hatchery-origin adult salmon and steelhead returns to the Upper Columbia Basin. Individuals with known locations of tagging and release as juveniles were included in the analysis. Release quantities and detection records were used to create datasets for analysis. All detection records for hatchery-origin spring, summer, and fall Chinook and summer steelhead that were PIT-tagged as juveniles and originated from the Wenatchee, Entiat, Methow, and Okanogan river subbasins and the upper Columbia River were included in the analysis (Figure 1). Occurrence of straying was evaluated at three spatial scales that include fish originating (released) from and returning to: (1) the upper Columbia River basin (e.g., above the confluence with the Snake and Yakima rivers); (2) a subbasin within the Upper Columbia (e.g., Wenatchee, Entiat, Methow, or Okanogan River subbasins or the Hanford

Reach of the Columbia River); and (3) a tributary of a subbasin (e.g., Chiwawa River, Nason Creek). These scales generally conform to management units of the Evolutionarily Significant Unit (Basin), the major spawning population (subbasin), and the spawning aggregate (tributary) (McClure et al. 2008). Summer Chinook Salmon reared at Wells Hatchery and released in the Methow and Okanogan rivers for survival studies in 2010 were included for upper Columbia River basin analyses but excluded from subbasin stray results because they were not acclimated consistent with the approved hatchery programs. Methods for assigning homing and straying are described in Pearsons and O'Connor (2020), but brief descriptions are provided below.

Donor stray percentage was calculated by summing the annual quantity of adults that strayed and dividing the annual stray total by the annual return total of the strayed and homed adults of the donor population. The average stray percentage was calculated by averaging the yearly stray percentages when the quantity of returning fish was five or greater. Years with fewer than five returning fish were excluded from the calculation because of potential extreme annual effects of low sample size.

We compared donor stray percentages using two different methods to evaluate the quality and consistency of the estimates made using PIT tags on return year and to reduce the number of metrics that were evaluated in this study. First, we compared return-year and brood-year donor stray percentages estimated using PIT tags. Brood-year donor stray percentages included all return years from a single brood and may reduce the influence of interannual environmental conditions on straying of adults when they migrate home. Second, we compared return-year donor stray percentages estimated using PIT with brood-year donor stray percentages estimated with CWT for Chinook Salmon only. Donor stray percentages derived from CWT were compiled from technical reports or generated from a United States Fish and Wildlife Service CWT database for upper Columbia River basin hatcheries (data accessed August 2019). Due to limited PIT tag samples for some programs, all spatial scales for CWT stray estimates were combined in order to make comparisons with PIT tag estimates. Only CWT stray estimates with temporal and spatial overlap for the PIT-based estimates were included. A correlation analysis was implemented to evaluate similarities among return- and brood-year estimates of donor stray percentages, and between PIT and CWT estimates of donor stray percentages.

Comparisons between donor stray percentages of hatchery- and natural-origin fish were made using the counts of PIT-tagged fish that homed and those that strayed at each spatial scale with all years pooled in a non-parametric contingency test (Fisher's Exact Test, Agresti 2002). Comparisons of the stray direction of hatchery- and natural-origin fish were made using Fisher's Exact contingency tests of the pooled counts of PIT-tagged fish that strayed downstream or upstream at each spatial scale. Donor stray percentages of fish that were moved to remote acclimation sites in the spring or fall were compared in a contingency test to those that were incubated, reared, acclimated and released from a single facility by pooling the years of each treatment for each facility. A one-tailed Fisher's Exact test p-value was used to test significance at an alpha of 0.05 . A one-tailed test was used because we were interested in detecting whether hatchery-origin stray rates were higher than natural-origin stray rates.

## Results

There were $5,652,887$ PIT tags injected into hatchery-origin juvenile fish and later evaluated to determine donor stray percentages of hatchery-origin salmon and steelhead in the
upper Columbia Basin. These included tags from specific hatchery programs (4,379,563; Table 1) and tags that were part of studies or tagged at collection sites in the natural environment where origin was known based upon fin clips, tags, and geographic location $(1,273,374)$. From those releases, 27,261 PIT tagged adult salmon and steelhead returned to the upper Columbia River Basin. Homing and straying totals for basin, subbasin, and tributary scales are presented in Table 2.

Table 2. Homing and straying of adult hatchery-origin PIT-tagged upper Columbia Watershed Chinook Salmon and steelhead 2000-2018. The range represents annual donor stray percentage.

| Scale | Location | Total N Home | Total N Stray | Range |
| :---: | :---: | :---: | :---: | :---: |
| Spring Chinook Salmon |  |  |  |  |
| Basin | Upper Columbia River | 5,378 | 3 | 0.06\% |
| Subbasin | Wenatchee River | 1,138 | 20 | 0-4.6\% |
| Tributary | Nason Creek | 93 | 3 | 0-7.3\% |
| Tributary | Chiwawa River | 241 | 104 | 8.3-55.6\% |
| Tributary | White River | 66 | 108 | 49.1-79.5\% |
| Subbasin | Methow River | 926 | 23 | 0-8.8\% |
| Subbasin | Okanogan River | 32 | 2 | 0-12.5\% |
| Summer Chinook Salmon |  |  |  |  |
| Basin | Upper Columbia River | 9,149 | 4 | 0.04\% |
| Subbasin | Wenatchee River | 190 | 57 | 6.1-35.0\% |
| Subbasin | Entiat River | 334 | 25 | 0-19.0\% |
| Subbasin | Methow River | 204 | 7 | 0-23.1\% |
| Subbasin | Okanogan River | 131 | 0 | 0\% |

## Fall Chinook Salmon

Upper Columbia River/Hanford
$\begin{array}{lllll}\text { Basin/Subbasin Reach } & 1,776 & 3 & 0.17 \%\end{array}$

## Steelhead trout

| Basin | Upper Columbia River | 11,178 | 3 | $0.03 \%$ |
| :--- | :--- | ---: | ---: | ---: |
| Subbasin | Wenatchee River | 978 | 131 | $0-31.4 \%$ |
| Tributary | Nason Creek | 103 | 74 | $21.7-61.1 \%$ |
| Tributary | Chiwawa River | 46 | 34 | $28.6-54.5 \%$ |
| Subbasin | Methow River | 173 | 25 | $0-25.0 \%$ |
| Tributary | Twisp River | 38 | 5 | $7.1-16.7 \%$ |
| Tributary | Chewuch River | 6 | 7 | $0-28.6 \%$ |
| Subbasin | Okanogan River | 466 | 7 | $0-15.8 \%$ |
| Tributary | Omak Creek | 335 | 16 | $0-21.3 \%$ |
| Tributary | Salmon Creek | 2 | 1 | - |

PIT-tag-based donor stray percentages by return year and brood year were highly correlated and were similar in magnitude for spring and summer Chinook Salmon and steelhead (Figure 2). In addition, the stray estimates generated from PIT tags and CWT were highly correlated and similar in magnitude for spring Chinook Salmon and highly correlated but different in magnitude for summer Chinook Salmon (Figure 2). Donor stray percentages of summer Chinook Salmon were about three times higher when estimated with PIT tags ( $<22 \%$ using PIT tags and $<8 \%$ using CWT). Only one fall Chinook hatchery (Priest Rapids Hatchery) in the upper Columbia River was available to estimate straying and the CWT estimate (3.3\%) was about 10 times higher than the PIT tag estimate ( $0.2 \%$ ). Stray estimates using CWT were not available for steelhead so they could not be compared to PIT estimates. In summary, both methods were highly correlated and produced similar results for spring Chinook Salmon, return year and brood year estimates for steelhead were highly correlated, PIT estimates were higher than CWT estimates for summer Chinook Salmon, and lower for fall Chinook Salmon. Other than the results we describe above, we present only return year results using PIT tags to allow comparison among all life history types of Chinook Salmon and between Chinook Salmon and steelhead using the same metric and to facilitate clarity and efficiency of the presentation. The implications of using return year estimates on our findings are presented in the discussion section.


Figure 2. PIT-tag-based return-year (RY, spawn year) stray estimates versus brood-year (BY) stray estimates using either PIT tags or BY coded wire tags (CWT) for upper Columbia watershed a) spring Chinook Salmon, b) summer Chinook Salmon, and c) steelhead.

## Spatial scale and taxa

Donor stray percentages of hatchery-origin fall, summer, and spring Chinook Salmon and steelhead were generally not higher than natural-origin donor stray percentages at larger spatial scales but were higher at smaller spatial scales. Donor stray percentages of hatchery-origin fall Chinook Salmon ( $\mathrm{P}=0.98$ ), summer Chinook Salmon ( $\mathrm{P}=0.96$ ), spring Chinook Salmon $(\mathrm{P}=0.60)$, and steelhead ( $\mathrm{P}=0.99$ ) were not significantly higher than natural-origin donor stray percentages at the basin scale and were $<0.3 \%$ (Figure 3).


Figure 3. Donor stray percentages of hatchery- and natural-origin a) spring Chinook Salmon, b) summer Chinook Salmon (SUC) and fall Chinook Salmon, (FAC) and c) steelhead at basin, subbasin, and tributary scales.

Hatchery-origin spring Chinook Salmon donor stray percentages were $<3 \%$, hatcheryorigin donor stray percentages of summer Chinook Salmon were $<22 \%$, hatchery-origin donor stray percentages of fall Chinook Salmon from the Hanford Reach was $<1 \%$, and hatchery-origin donor stray percentages of steelhead was $<11 \%$ at the subbasin scale (Figure 3). At the subbasin scale, donor stray percentages of hatchery-origin fall Chinook Salmon ( $\mathrm{P}=0.77$ ), summer Chinook Salmon ( $\mathrm{P}=0.45$ ), and spring Chinook Salmon ( $\mathrm{P}=0.16$ ), were not significantly higher than natural-origin donor stray percentages, but donor stray percentages of hatchery-origin steelhead were significantly higher than natural-origin donor stray percentages ( $\mathrm{P}<0.0001$ ) (Figure 3). Results for spring, summer, and fall Chinook Salmon were consistent across individual subbasins ( $\mathrm{P}>0.05$ ), but donor stray percentages of hatchery-origin steelhead in the Okanogan subbasin were not significantly higher than natural-origin donor stray percentages $(\mathrm{P}=0.75)$ despite the other subbasins being different $(\mathrm{P}<0.05)$.

At the tributary scale, donor stray percentages of hatchery-origin spring Chinook Salmon ( $\mathrm{P}<0.001$ ), were significantly higher than natural-origin donor stray percentages (Figure 3). There was some variation in differences within each of the taxa and in some tributaries. For example, donor stray percentages of hatchery-origin spring Chinook Salmon in Nason Creek and the White River were not significantly higher than natural-origin donor stray percentages in those tributaries ( $\mathrm{P}>0.05, \mathrm{n}=2$ natural-origin spring Chinook Salmon at White River). Donor stray percentages of hatchery-origin steelhead in the Twisp River were not significantly higher than natural-origin donor stray percentages ( $\mathrm{P}=0.25$ ). Hatchery-origin spring Chinook Salmon donor stray percentages were as high as $62 \%$ and 3 of 4 tributary hatchery-origin donor stray percentages were numerically higher than natural-origin donor stray percentages from the same tributary (Figure 3).

## Stray direction

Hatchery-origin Chinook Salmon and steelhead generally strayed in an upstream direction (i.e., overshot the target destination such as a target tributary or subbasin as opposed to a location within a tributary or subbasin) and the proportions of hatchery fish that strayed upstream was not significantly different than natural-origin fish ( $\mathrm{P}>0.05$; Figure 4). In general, of those fish that strayed over $80 \%$ of hatchery-origin Chinook Salmon and steelhead strayed in an upstream direction and some hatchery populations only strayed in an upstream direction (Figure 4). The few exceptions to this pattern were cases with limited opportunities to stray in an upstream direction, such as fish released in the Okanogan subbasin.


Figure 4. Direction of hatchery- and natural-origin straying. Abbreviations: SPC spring Chinook Salmon, SUC summer Chinook Salmon, FAC fall Chinook Salmon, STH steelhead. W Wenatchee River, M Methow River, E Entiat River, HR Hanford Reach, NC Nason Creek, CW Chiwawa River, LV Leavenworth Nation Fish Hatchery, TW Twisp River, CE Chewuch River, MH Methow Hatchery, WH Winthrop National Fish Hatchery, EH Entiat National Fish Hatchery, OC Omak Creek.

Movement for remote acclimation

Only spring Chinook Salmon and steelhead met the criteria for comparing donor stray percentages of fish that were moved between facilities for acclimation and those that were not. Spring Chinook Salmon that were moved to other tributaries for acclimation strayed at much higher percentages than those that completed their incubation, rearing, and acclimation at a single location ( $\mathrm{P}<0.0001$; Figure 5). In contrast, steelhead that were moved for acclimation did not stray at higher percentages than those that completed their incubation, rearing, and acclimation at a single location ( $\mathrm{P}=0.69$; Figure 5).


Figure 5. Mean donor stray percentages of hatchery-origin fish that were moved as juveniles among facilities prior to release (moved) or those that were incubated, reared, acclimated and released from a single facility (not moved). Error bars represent the range of values when more than one value was available. Abbreviations: SPC spring Chinook Salmon, SUC summer Chinook Salmon, FAC fall Chinook Salmon, STH steelhead. W Wenatchee River, M Methow River, E Entiat River, O Okanogan River, HR Hanford Reach.

## Management changes

The management actions that were implemented to reduce straying produced mixed results. Donor stray percentages were not significantly different for summer Chinook Salmon released into the Methow subbasin in the years when they were both overwinter and spring acclimated (2015-2017) than when they were just spring acclimated (2010, 2011, 2014), $\mathrm{P}=0.19$; (Figure 6). Overwinter acclimation of steelhead in the Wenatchee subbasin did result in lower donor stray percentages at the subbasin scale ( $\mathrm{P}<0.0001$, Figure 6).


Figure 6. Mean donor stray percentages in periods of differing management regimes for a) summer Chinook Salmon released from the Carlton Acclimation Facility on the Methow River, and b) steelhead released into the Wenatchee River after being raised at either Turtle Rock or Eastbank Hatchery. Bars represent the mean of annual estimates and error bars are ranges.

## Discussion

Spatial scale (size of the target) and risk
Hatchery-origin fish were able to return to the largest target (basin) with high accuracy and were as accurate as natural-origin fish. As the target size became smaller and more numerous, such as subbasins and tributaries, the accuracy decreased for both returning hatcheryand natural-origin fish, but more so for hatchery-origin fish (Figure 7). There are many factors that may contribute to increasing homing accuracy with increasing spatial scale. The most obvious factor is that it is easier to find a big target than a small one and that there are more opportunities to miss targets at the tributary level because there are more tributaries than basins or subbasins. Another factor that likely contributed to the basin accuracy was that most juvenile

## TARGET AREA

FLOW


Figure 7. Comparisons of homing rates (minimums in black) and straying (maximums in grey) between hatchery- and natural-origin Salmon and steelhead at three spatial scales in the upper Columbia subregion. The size of the targets was scaled to the fall discharges of the upper Columbia River (basin), the mean of the Wenatchee, Entiat, Methow, and Okanogan subbasins (subbasin), and the mean of tributaries to subbasins (tributary).
fish were not moved outside of the basin: all of the PIT tagged fish that were released into the upper Columbia basin were spawned, incubated, reared and released into the upper Columbia basin except for White River spring Chinook Salmon. As such, with one exception, the fish were exclusively imprinted on upper Columbia basin water and oriented on upper Columbia basin geography. In contrast, many of the fish released into subbasins and tributaries were moved between two hatchery facilities prior to release which likely contributed to reduced homing by hatchery-origin fish (discussed below).

The demographic and genetic risks of hatchery-origin salmon and steelhead straying varied dramatically with spatial scale but risks to harvest were universally low. At the upper

Columbia basin scale over $99.7 \%$ of returning hatchery-origin fish homed to the basin of origin and the donor stray percentages were similar to natural-origin fish. In contrast, returning hatchery-origin fish donor stray percentages were as high as $62 \%$ at the tributary scale and the donor stray percentages were higher than natural-origin fish in many, but not all, tributaries. Straying posed little risk to harvest objectives at the spatial scales considered because fisheries occurred downstream of areas where fish stray, such as in the ocean, Columbia River, and subbasins; and not in tributaries (Hillman et al. 2019; Pearsons et al. 2020).

For conservation hatchery programs, straying had the potential to result in demographic risks at the tributary scale for spring Chinook Salmon and at tributary and subbasin scales for steelhead (excluding tributaries of the Wenatchee River) because strays did not contribute to target spawning populations in all cases and therefore may not contribute to population recovery of these ESA listed species. However, these strays might have also contributed to the demographics of other nearby non-target spawning aggregates or populations. For example, spring Chinook Salmon released in the Chiwawa River contributed substantial numbers of strays to the adjacent Nason Creek spawning aggregate and these fish contributed to natural production (Williamson et al. 2010; Ford et al. 2015a). In other cases, the scale of population management can influence whether a fish is characterized as a stray or not and management zones can influence the magnitude of demographic or genetic risks. In short, a portion of the hatcheryorigin returns had the potential to contribute to target spawning aggregates while others strayed nearby and potentially contributed to the larger population at the subbasin and basin scale. Total numbers of fish produced naturally from hatchery-origin fish that homed or strayed away from target spawning locations in tributaries or subbasins may produce the same numbers of offspring in the basin as if they all spawned in target locations. However, among other things, this assumes that density-dependent mortality is equal among spawning and rearing locations and that the genetic characteristics of hatchery-origin fish does not influence the reproductive success in non-target areas. Both of these assumptions are unlikely to be true (Williamson et al. 2010; Ford et al. 2015a; Ford et al. 2016).

Finally, genetic risks to nearby spawning aggregates occur when strays potentially disrupt local adaptation (McClure et al. 2008; Keefer and Caudill 2014). These genetic risks are most likely to occur within spawning aggregates of a subbasin for spring Chinook Salmon, and for some spawning aggregates and major population groups for steelhead. The degree of risk is likely influenced by the amount of reproductive success that is influenced by genetic differentiation. Fish that stray into populations that are genetically similar to one another pose lower risk than those that are very different. In general, adjacent populations are genetically more similar than those that are geographically separated by longer distances (Hillman et al. 2019), so adjacent populations are also less likely to dramatically influence local adaptation. The genetic risks of straying are better evaluated by estimating recipient population stray percentage than donor stray percentage because recipient population stray percentage also incorporates the size of the recipient population relative to the abundance of strays (Keefer and Caudill 2014; Bett et al. 2017). For example, high donor population stray percentages may pose low genetic risks to large recipient populations but high genetic risks to small recipient populations.

The patterns and magnitudes of hatchery-origin fish straying that we present in this study were within the range of those presented by others that work in the Columbia Basin and elsewhere. For example, Ford et al. (2015a) found that donor stray rates of hatchery-origin spring Chinook Salmon in the Chiwawa River using CWT were higher than those estimated for natural-origin fish using genetic methods and that approximately $5 \%$ strayed to other subbasins.

Westley et al. (2013) assessed donor straying at the subbasin scale in the Columbia River Watershed and observed a wide range of stray rates of hatchery-origin Chinook Salmon and steelhead from $0.11 \%-54.9 \%$. Donor stray rates of fish at the subbasin scale in the upper Columbia Basin (1.6-21.6\%) were within the range of other parts of the Columbia Basin (Westley et al. 2013). Donor stray rates of returning hatchery-origin Chinook Salmon released as yearlings in the Yakima Basin were very low at the Yakima Basin and subbasin scale and relatively high at the tributary scale (Dittman et al. 2010; Fast et al. 2015). Over 55\% of returning hatchery-origin spring Chinook Salmon were recovered over 25 km from their acclimation release site and donor stray rates of fish released from the Jack Creek Acclimation site were approximately 76\% (Dittman et al. 2010; Cram et al. 2012). Finally, donor stray rates of returning hatchery-origin fall Chinook Salmon in California's Central Valley ranged between $0 \%$ and $89 \%$ (Sturrock et al. 2019).

The lack of differences that we found in at least one of our comparisons was likely the result of low sample size and associated low statistical power (Ham and Pearsons 2000). We did not detect a difference in donor stray percentages of hatchery- and natural-origin spring Chinook Salmon in the White River even though the estimated donor stray percentages were $62 \%$ and $0 \%$, respectively, and was the highest donor stray percentage of hatchery-origin fish that we evaluated. The sample size of the natural-origin population was only two fish, which was lower than the standard we used for hatchery-origin fish ( $n>4$ ), and was the reason why the statistical test did not result in a statistically significant result. The donor stray percentages of naturalorigin spring Chinook Salmon in other tributaries of the upper Columbia with higher sample sizes has been below 7\% (Pearsons and O'Connor 2020) and it is likely that even with a larger sample size, these rates also apply to natural-origin spring Chinook Salmon in the White River. Thus, it is likely that hatchery-origin donor stray percentages in the White River were substantially higher than natural-origin donor stray percentages and we simply couldn't detect it because of the low sample size that was used to estimate natural-origin donor stray percentages. It is possible that lack of detectable differences occurred for other comparisons in our evaluation, but visual examinations of the graphs (Figures) do not indicate dramatic omissions in detectable differences such as occurred in the White River.

## Factors influencing straying

There are multiple factors that may influence hatchery-origin spring Chinook Salmon and steelhead to stray at higher percentages than natural-origin fish in tributaries. In addition to the transportation of fish from incubation and rearing sites to release and/or acclimation sites described below, the hatchery rearing environment may also be a factor that affects homing success (Ford et al. 2015a). In a review of straying, Keefer and Caudill (2014) reported that hatchery-origin fish were widely believed to have reduced imprinting compared to natural-origin fish, in part because of reduced stimuli in the hatchery environment and lower olfactory activity and reduced brain development compared to natural-origin fish. In addition, Westley et al. (2013) found that the hatchery practice of rearing ocean-type Chinook Salmon as yearlings rather than the subyearlings (the natural age at migration) was associated with increased straying. The hatchery management approach of extended rearing is used in the upper Columbia for summer Chinook Salmon and results in a possible trade-off between increased post-release survival and increased straying (see Unwin and Quinn 1993). Without addressing the trade-offs of survival inside and outside hatcheries, rearing conditions in hatcheries that may be responsible for
reduced imprinting at finer scales of resolution, and straying, it may not be possible to achieve management objectives of homing.

Responses of hatchery- and natural-origin fish to factors outside of the hatcheries may also explain the variation in straying we observed. For example, barrier weirs for collecting broodstock near acclimation sites may increase straying and also result in a higher propensity for hatchery-origin fish to be displaced than natural-origin fish (Bugert 1998; Clarke et al. 2012) because they are often located closer to hatchery acclimation sites than natural spawning sites farther upstream which may result in less drive to negotiate a barrier if they are near their homing target site (Hoffnagle et al. 2008). In addition, thermal attractants, or thermal or physical barriers may increase wandering behavior and ultimately straying (Leider 1989; Bond et al. 2017; Richins and Skalski 2018), but it is unclear how this would influence hatchery-origin fish differently than natural-origin fish, unless run and spawn timing differed between origins (Hoffnagle et al. 2008). Finally, poor habitat quality in areas near acclimation sites may increase straying outside of a tributary by hatchery-origin fish returning to the area around the acclimation facility (Cram et al. 2012; Fast et al. 2015; Ford et al. 2015a). In short, using best practices for imprinting hatchery-origin fish may not result in achieving management objectives because factors outside of the hatchery can influence straying too. Therefore, management actions inside and outside hatcheries should be considered in order to increase the potential of meeting management objectives for homing. Alternatively, managers could shape objectives for homing in accordance with the physical constraints of the river systems and facility infrastructure, and the biological characteristics of the supplemented species.

## Stray direction

Contrary to our hypothesis that the direction (upstream vs. downstream) of donor stray percentages would vary depending upon hatchery locations, hatchery-origin fish generally strayed in an upstream direction similar to natural-origin fish (Pearsons and O'Connor, 2020). There may be fitness advantages to stray in an upstream direction if there is a higher probability of colonizing new habitats that are more productive than target or downstream habitats (Pearsons and O'Connor, 2020). In contrast, Dittman et al. (2010) found that hatchery-origin spring Chinook Salmon in the Yakima Basin spawned upstream of their acclimation site when the acclimation site was low in the system and downstream of their acclimation sites when they were located high in the system. This result may have been confounded by limited spawning habitat upstream of acclimation sites because of the presence of a dam and reservoir or because of an increase in stream gradient. In addition, differences between studies may be the result of differences in the spatial scales that were assessed. In our work we did not evaluate straying direction within a specific spatial scale such as a subbasin such as was done by Dittman et al. (2010), but rather between tributary and subbasin junctures. Similar to our findings, straying between spawning aggregations in the Yakima Basin was in an upstream direction. Knowing the direction of straying can be used to assess risks to nearby populations and to plan appropriate management actions to reduce impacts and achieve acceptable escapement goals. For example, genetic risks to upstream populations would be assessed to be higher than to downstream populations if suitable spawning areas were available upstream. One approach to reduce straying is to locate hatcheries or to release fish far upstream of where populations of concern exist and where upstream straying could be contained. It is also possible that locating releases far upstream in a tributary might reduce wandering behavior to other tributaries that could occur in
the absence of embryonic imprinting. This strategy is not without risk because the farther upstream fish are released the greater migration distance and lower migration survival as well as the potential for increased ecological risks (Pearsons and Hopley 1999; McMichael et al. 1999; Pearsons et al. 2012).

## Moved vs. non-moved

Donor stray percentages of hatchery-origin spring Chinook Salmon that were transported for acclimation and/or release (but not mainstem Columbia River truck or barge transport) had greater deviations from donor stray percentages of natural-origin fish than those that were not transported. This observation is consistent with the sequential imprinting hypothesis (Scholz 1980; Hasler and Scholz 1983; Dittman et al. 2015) and also with evaluations of downstream transportation during spring outmigration (Bond et al. 2017; Sturrock et al. 2019). It appears that fish were able to find their way back to the subbasin of release, but then fish strayed possibly because they were searching for the location of their birth and that location was far from the release location. It is not clear whether the transportation of spring Chinook Salmon results in straying because of imprinting on another water source at an earlier life stage or because of disruption of the appropriate geographic cues or some other factor. If imprinting on another water source is the primary factor contributing to straying, then transportation of water to a centralized hatchery facility or exposure to unique odors could be used to imprint fish, particularly if it can be done when fish are embryos (Dittman et al. 2015). If disruption of appropriate geographic cues caused by transportation is the primary factor contributing to straying, then it is not clear what could be done to reduce donor straying if fish must be transported. High straying of hatchery-origin spring Chinook Salmon at the tributary scale also occurred in a Yakima Basin tributary, North Fork Teanaway River, even though the tributary was located within the same subbasin as the central hatchery facility and fish were moved during the spring for acclimation in the North Fork Teanaway River (Dittman et al. 2010; Cram et al. 2012); a scenario very similar to spring Chinook Salmon management within the Methow subbasin. It is unclear why steelhead did not exhibit the same patterns of differences associated with movement between hatchery facilities that spring Chinook Salmon did.

The highest donor stray percentage that we observed occurred in the White River Captive Broodstock spring Chinook Salmon program. The fish that were released for this program were founded from local broodstock and incubated, hatched, and reared to yearlings at the Little White Salmon National Fish Hatchery located on the Little White Salmon River, a tributary that enters the Columbia River hundreds of kilometers downstream of the upper Columbia Basin (Ford et al. 2015b). During the spring, spring Chinook Salmon yearling parr were trucked to the White River or Lake Wenatchee for at least six weeks of acclimation (Figure 1). Most fish were acclimated in streamside tanks or in net pens in the lake and released in those locations or trucked and released in the Wenatchee River below Lake Wenatchee to avoid low migration survival in the lake. The convoluted sequence of transportation and acclimation these fish experienced likely contributed to the highest donor stray percentages we observed.

The management action with the highest potential to reduce donor stray percentages is to reduce or eliminate the transportation of fish after the eyed-egg stage. However, this action is problematic for a variety of reasons. First, there is often not enough water to complete incubation, rearing, and acclimation at many remote sites such as in smaller tributaries. Some sites that do not have enough water to provide single-pass water through rearing vessels may
have enough water to consider high amounts (e.g., >95\%) of water recirculation, but this might pose other fish-culture risks such as disease and poor fish quality. However, these risks have not been manifested for summer Chinook Salmon reared with $60 \%$ reuse water at Eastbank Hatchery or Wenatchee steelhead reared with reuse water at Chiwawa Acclimation Facility (Chelan Public Utility District, unpublished data). Second, the cost of building and operating new infrastructure for existing programs may be prohibitive and there is potential that additional infrastructure within spawning and rearing areas could reduce habitat quality for salmon and steelhead. New hatchery programs should consider ways to minimize fish transportation if donor stray percentages between tributaries are an important consideration for program success.

Responses to management actions
The management actions that were implemented to reduce straying had mixed results, suggesting that there is much to learn about the factors governing straying (Keefer and Caudill 2014). Management actions at the Carlton overwinter acclimation facility for summer Chinook Salmon assumed that additional imprinting occurred during the winter. The lack of detectable stray differences in this program suggests that imprinting may not be important during the winter for these yearling Chinook Salmon. In addition, donor stray rates of returning Chinook Salmon released as yearlings were not different in a study involving two- and four- month acclimation prior to release in early March into the Umatilla River, Oregon (Clarke et al. 2012), suggesting further that acclimation during the winter period may not be a strong factor influencing straying of returning Chinook Salmon released as yearlings.

One of the main reasons for acclimating fish during both the winter and into spring is to reduce straying (Clarke et al. 2012). However, our results indicate that homing at the tributary and larger scales was not improved by providing overwinter acclimation of Chinook Salmon at satellite sites. It was hypothesized that longer periods of acclimation may improve imprinting, however, the length of time may be less important than specific periods when salmon are known to imprint such as during embryonic development and smoltification (Scholz 1980; Dittman et al. 2015). Overwinter acclimation can provide other benefits to fish besides the potential for improved homing by exposing them to more natural water temperatures that modulates fish growth (Clarke et al. 2012; Larsen et al 2013). However, overwinter acclimation may also result in undesirable impacts to fish. For example, acclimation at remote sites is typically more challenging than at centralized locations because of higher exposure to pathogens downstream of decaying carcasses, higher turbidity, and debris and icing risks to intakes that compromise access to water and these factors may result in high on-site juvenile mortality. It is unclear whether the high financial costs and additional ecological and demographic risks associated with overwinter acclimation is sufficient to outweigh the benefits of overwinter acclimation if the acclimation does not reduce straying enough to meet management objectives. Assessments of the risks and benefits of overwinter acclimation are likely to be idiosyncratic for each hatchery program, resulting in the need to evaluate them on a case-by-case basis. For example, spring Chinook Salmon that were reared at the same central hatchery facility and then acclimated at three different satellite sites in the Yakima River had significantly different homing patterns relative to their acclimation site (Dittman et al. 2010); a phenomenon that was also observed in our study.

In contrast to the Chinook Salmon example, the management action to reduce steelhead straying appeared to be successful in the Wenatchee subbasin. Multiple factors may have contributed to improved homing of steelhead in the Wenatchee River including longer term
acclimation and changing water sources during rearing. In addition, summer steelhead overwinter as adults so multiple cues over a longer time may benefit steelhead homing more than Chinook Salmon. The variability in success of management actions prompt us to recommend that the mechanisms of straying be better understood before making large investments in costly management actions.

Tag methods and influence on conclusions
Our data suggests that donor stray percentages estimated by run year or brood year could be used interchangeably for all species but that the type of tag used resulted in varying differences of estimates depending upon the species. For example, donor stray percentages estimated using PIT and CWT were similar for spring Chinook Salmon, PIT estimates were three times higher than CWT for summer Chinook Salmon, and ten times lower than CWT for fall Chinook Salmon (although estimates using each method were $<5 \%$ for fall Chinook Salmon). These differences could result from the logistics of tag detection for each tag type. PIT tag detections at in-stream arrays were easier to reconcile with physically recovered CWT tags at the tributary scale where spring Chinook Salmon spawn, less so at the subbasin scale because the larger flows where summer Chinook Salmon primarily spawn could make it more difficult to both detect PIT tags and recover CWT, and most difficult at the basin scale (mainstem Columbia River) where fall Chinook Salmon spawn because PIT tags are nearly always detected at mainstem dams while CWT recovery is more challenging. The physical placement of PIT detection arrays throughout the upper Columbia Watershed allowed us to assess whether the locations of the last detection were in the vicinity of expected spawning locations and limit which fish were included at each spatial scale. We used consistent methods in this study by only using PIT tags so the comparisons between hatchery- and natural-origin fish should not be influenced by our methods. However, the magnitude of stray estimates could be influenced by the methodology associated with the different type of tag that we used. In general, the magnitude of spring Chinook Salmon estimates are likely accurate, summer Chinook Salmon are likely overestimates, and fall Chinook Salmon are likely underestimates.

## Applications

All hatchery programs are unique and therefore the findings we describe for the Upper Columbia Basin may differ in other locations. Indeed, substantial variation in donor stray percentages between hatcheries occurred within the Upper Columbia Basin. Furthermore, the hatchery programs in the Upper Columbia are well funded and managed with a high degree of oversight and hatchery programs that deviate from the practices used in the Upper Columbia may generate different results.

This work focused on donor population strays, but recipient population stray evaluations should also be considered. Recipient population stray rates are influenced by factors such as the size of the hatchery program, the size of the recipient population, and the donor stray rate (Bett et al. 2017). Large hatchery programs that are adjacent to small natural populations must have very low donor population stray rates in order to meet recipient population stray rates of 5 or $10 \%$. In some cases, difficult trade-offs will be necessary to achieve potentially competing stray, survival, and program size objectives. In some cases, fish culture techniques such as raising summer Chinook Salmon to yearlings, moving fish to remote acclimation sites, and maintaining large
program sizes will need to be evaluated relative to the amount of straying that occurs. In still other cases, the only way to comply with mandated recipient population stray rates will likely be to reduce hatchery program size or change release locations.

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# Stray Compositions of Hatchery-Origin Chinook Salmon Oncorhynchus tshawytscha and Steelhead O. mykiss in Recipient Natural Populations of the Upper Columbia Watershed 

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#### Abstract

One of the biggest concerns of operating hatchery Salmon and steelhead programs is high straying of returning adults into non-target populations and the possible homogenization of genetic diversity among populations caused by spawning of stray fish. The composition of hatchery-origin stray Chinook Salmon Oncorhynchus tshawytscha and steelhead O. mykiss relative to the natural spawning populations, termed recipient population stray rate, was evaluated in the Upper Columbia Basin. Chinook Salmon carcasses were collected from 19992018 in spawning areas shortly after spawning and carcasses were examined to determine origin. Adipose fin clips and coded-wire-tags were used to distinguish non-target hatchery, target hatchery, and natural-origin fish; coded-wire-tags were read in the lab to determine the origin of hatchery-origin fish. Steelhead strays and spawning escapement were evaluated using passiveintegrated transponder (PIT) tags between 2013-2018. The recipient population stray rates ranged between 0.02-87.35\% and increased with decreasing spatial scale. Recipient stray rates of all taxa at the basin scale were $<3 \%$, and summer Chinook and fall Chinook salmon were $<0.5 \%$. Stray rates in subbasins for all taxa ranged between $0.07-33.04 \%$; spring and summer Chinook Salmon exceeded 5\% in some 10 year periods in the Entiat and Methow subbasins, but stray rates for all Chinook Salmon were $<5 \%$ in the Wenatchee, Okanogan, and Hanford Reach for all periods. All steelhead stray rates exceeded 5\% except for those in the Wenatchee subbasin. Stray rates of spring Chinook Salmon in tributaries (the only taxa that met the tributary criteria) ranged between $0.61 \%-87.35 \%$ and only the Chiwawa, Icicle, and Twisp rivers were consistently below $10 \%$; the Chiwawa River was consistently below $5 \%$. In cases where recipient stray management targets were exceeded, some were the result of single hatchery contributions, but others were the result of cumulative contributions from multiple hatcheries. Options to achieve recipient stray management targets include reducing donor stray rates, reducing hatchery program size, removing hatchery-origin adults prior to spawning in the natural environment, and increasing the natural-origin population. It is likely that balancing trade-offs among hatchery program size and recipient population stray rate will be necessary in order to achieve management targets in some locations.


## Introduction

Hatcheries are frequently used to increase abundance of Chinook Salmon Oncorhynchus tshawytscha and steelhead $O$. mykiss for harvest and conservation, but because of the large numbers of fish produced and the manner in which they are produced, unintended consequences can occur that pose genetic risks to natural populations that are not the target of the production (Keefer and Caudill 2014; Bett et al. 2017; Pearsons and O’Connor 2021). Hatchery-origin Chinook Salmon stray at higher rates than natural-origin fish at some spatial scales, and they are often more abundant than natural-origin fish on the spawning grounds (Keefer and Caudill 2014; Pearsons and O'Connor 2021). In addition, migration and spawning habitats have been altered by humans, which can increase the magnitude of straying (Cram et al. 2012; Ford et al. 2015; Bett et al. 2017). These factors can result in large numbers of stray fish spawning with fish that were not the intended target of hatchery augmentation. Furthermore, many naturally spawning populations of salmon and steelhead have declined from historic levels and therefore hatcheryorigin strays can make up large proportions of the spawning population even when the stray rate is low (Bett et al. 2017; Sturrock et al. 2019).

One of the main concerns with straying of hatchery-origin Chinook Salmon and steelhead is the reduction of local adaptation that occurs through inter-breeding of hatchery- and naturalorigin fish in the natural environment (Keefer and Caudill 2014; Bett et al. 2017). This could occur through mechanisms such as outbreeding depression and domestication (Busack and Currens 1995). Local adaptation can be reduced if sufficient numbers of hatchery-origin fish stray into non-target populations and if they reproduce successfully. This can further result in a reduction in genetic diversity between populations, which can increase extinction risk. Alternatively, straying can result in demographic or genetic rescue in cases of high disturbance or low population size (Bett et al. 2017; Pearsons and O’Connor 2020).

The best stray metric to assess the risk of straying to genetic diversity is referred to as recipient population stray rate (Keefer and Caudill 2014). Recipient population stray rate is quantified as the proportion of the total spawning population that is composed of non-target hatchery-origin strays (Bett et al. 2017). It is distinguished from supplementation programs that intentionally produce fish to contribute to the natural production of a target population (Mobrand et al. 2005; Paquet et al. 2011; Fast et al. 2015). Recipient population stray rates are underrepresented in the literature compared to donor rates, the stray rates of contributing hatcheries, despite the higher management importance of recipient stray rates (Keefer and Caudill 2014; Bett et al. 2017). In addition, relatively few studies have evaluated recipient population straying from multiple hatcheries, species, and spatial scales (Bett and Hinch 2015).

Fisheries managers set recipient stray rate targets for hatchery programs in efforts to maintain local adaptation and trigger management actions to control excessively high stray rates. These targets were informed by genetic modelling of how much gene flow could occur without losing important genetic diversity of recipient populations (Craig Busack, NOAA Fisheries, personal communication). Targets generally range between 2 and 10\%, and can vary depending upon management objectives and risks to local adaptation (Ford 2002; Mobrand et al. 2005; Paquet et al. 2011; Brenner et al. 2012; Hillman et al. 2018). Strays from distant locations are generally regarded as higher risk than those from adjacent locations because they are hypothesized to be less adapted to local conditions than nearby populations (Fraser et al. 2011). For example, the recipient stray management targets for the upper Columbia Basin are: 1)
hatchery-origin strays make up less than $5 \%$ of the spawning escapement within non-target recipient populations, and 2) hatchery-origin strays from a spawning aggregate within a population make up less than $10 \%$ of the non-target spawning aggregate within the same population (Hillman et al. 2019).

Recent work has estimated donor population stray rates for both natural- and hatcheryorigin Salmon and steelhead in the upper Columbia Basin (Pearsons and O'Connor 2020, Pearsons and O'Connor 2021). This work demonstrated that stray rates of hatchery- and naturalorigin fish increased with decreasing spatial scale but the disparity was more pronounced by hatchery-origin fish, particularly at the tributary scale. Furthermore, the magnitude of hatcheryorigin fish straying posed risks to the genetic diversity of the populations and warranted estimation of recipient population straying. This paper fills that gap for the upper Columbia subregion. More specifically we: 1 ) assess the magnitude and composition of recipient population stray rates of spring, summer, and fall Chinook Salmon and summer steelhead spawning populations at three spatial scales, 2) assess (i.e., basin, subbasin and tributary) factors that influence recipient population stray rates, and 3) discuss trade-offs of achieving recipient stray management targets.

## Methods

## Study Area

This study was conducted in the Columbia River, USA, and most of the work was conducted in the upper Columbia Basin upstream of the confluence with the Snake River and downstream of Chief Joseph Dam, with fish from hatchery programs in the Wenatchee, Entiat, Methow, and Okanogan subbasins and the upper Columbia River (Figure 1). The upper Columbia River Basin has an abundance of hatchery facilities as a result of mitigation for the construction and operation of hydropower dams (Figure 1). These hatcheries produce fall, summer, or spring Chinook Salmon, Coho Salmon O. kisutch, Sockeye Salmon O. nerka, and steelhead for harvest, conservation, or a combination of both; but Chinook Salmon and steelhead are the only species considered for this assessment. Some of the hatchery programs incubate, rear, and release fish from a single hatchery location, whereas other programs transport parr or smolts to acclimation sites for subsequent release. The study area and biological background was previously described by Pearsons and O'Connor $(2020,2021)$ and is also briefly described below.


Figure 1. Map of the study area of the upper Columbia Basin and areas outside of the upper Columbia Basin that contributed strays to recipient populations within the upper Columbia Basin.

Fall Chinook Salmon spawn in the Hanford Reach, the only free-flowing reach of the Columbia River between Grand Coulee and Bonneville dams These spawners comprise one of the largest Chinook Salmon populations in the United States, and contribute large numbers of fish to harvest in the Pacific Ocean and Columbia River, making this population economically very important (Harnish et al. 2014; Langshaw et al. 2017; Pearsons et al. 2020). Summer Chinook Salmon spawn primarily in the mainstems of four subbasins of the upper Columbia River (e.g., Wenatchee, Entiat, Methow, and Okanogan) and support considerable fisheries in the Pacific Ocean and Columbia River. The naturally produced juveniles of summer and fall run Chinook Salmon generally migrate to the sea as sub-yearlings. Spring Chinook Salmon spawn in tributaries to mainstem subbasins and in upper portions of mainstem subbasins (Williamson et al. 2010; Murdoch et al. 2010; Ford et al. 2015a). Upper Columbia River spring Chinook Salmon are listed under the Endangered Species Act (ESA) as endangered (McClure et al. 2008). The naturally produced juveniles of spring Chinook Salmon migrate to the sea as yearlings. Summer steelhead spawn throughout upper Columbia subbasins and are ESA listed as threatened (Ford et al. 2016). Naturally produced juvenile steelhead from the upper Columbia migrate to
the sea at ages 1-7 years, but most migrate at ages 1-3 (Peven et al. 1994). All races of Chinook Salmon and steelhead in the upper Columbia have a long history of interactions with hatchery programs and hatchery- and natural-origin fish overlap in much of their spawning distributions (e.g., Williamson et al 2010; Pearsons et al. 2012; Ford et al. 2015a; Ford et al. 2016; Johnson et al. 2018).

## Hatchery description and tagging

Hatchery-origin fish were produced in a variety of hatcheries throughout the Upper Columbia River Basin; see Pearsons and O'Connor, (2021) for information about hatchery programs and release and recovery locations. Most hatchery-origin Chinook Salmon were marked and tagged to facilitate identification to identify their hatchery of origin when recovered as carcasses on the spawning grounds. Fish were tagged with coded-wire tags (CWTs) as juveniles. Tags were generally placed in the snout and each CWT was specific to a hatchery. In a few instances, CWTs or blank wire tags were placed in the caudal peduncle near the adipose fin. Fish were tagged months before release and then released during the spring as subyearlings or yearlings. Steelhead were tagged with PIT tags to identify the hatchery of origin because of the inability to collect carcasses on the spawning grounds (Pearsons and O'Connor 2021). Fish migrated to the ocean and then returned to spawn 1 to 5 years later.

Spawning escapement and composition
A variety of field methods were used to estimate the two derived metrics needed for calculating recipient population stray rate; spawning escapement, and origin composition of spawners. Spawning escapement of spring and summer Chinook Salmon was estimated by multiplying the number of redds by the number of fish per redd (Hillman et al. 2019). The number of fish per redd was estimated at dams or hatcheries by dividing the total abundance by the number of males (Murdoch et al. 2010) and assuming one female per redd (Murdoch et al. 2008). Spawning escapement of fall Chinook Salmon was estimated by counting the number of fall Chinook at McNary Dam and subtracting the number of fish counted at Ice Harbor dam as well as harvest and hatchery returns (Basin estimate; Richards and Pearsons 2019). The escapement of fish to the Hanford Reach of the Columbia River also involved subtracting counts of fall Chinook from the Yakima subbasin and Priest Rapids Dam. Spawning escapement of steelhead was estimated by a mark-recapture method (Hillman et al. 2019).

About $15 \%$ of returning adult steelhead passing Priest Rapids Dam were PIT tagged and subsequently detected or "recaptured" at upstream antennas located in subbasins and tributaries throughout the upper Columbia Basin. A mathematical model was used to estimate escapement to subbasins based upon the number of steelhead PIT tagged at Priest Rapids Dam and the detection of fish at PIT tag antenna arrays within each subbasin (Hillman et al. 2019). We subtracted the number of steelhead harvested, collected for broodstock, or removed for other management purposes in each subbasin from the modelled subbasin escapement to estimate spawning escapement for each subbasin. Spawning escapement estimates for all tributaries could not be generated using available data and procedures.

The composition of spawners on the spawning grounds was estimated using CWTs (Chinook Salmon) and PIT tags (steelhead). Coded-wire tags were collected from Chinook Salmon carcasses. The CWTs were expanded based upon hatchery-specific marking rate (e.g., proportion of the hatchery production that was tagged) and the sampling rate. Hatchery-origin
fish that were not part of the target spawning population were classified as strays. The abundance of the natural-origin population was estimated by subtracting the number of target and non-target hatchery-origin fish by the total escapement.

Field Methods

Spawning-ground surveys were conducted annually throughout the upper Columbia Basin to estimate the number of Chinook Salmon redds and the composition of spawners (Murdoch et al. 2009 a, b; 2010; Hillman et al. 2019; Richards and Pearsons 2019; Snow et al. 2019). Surveys were conducted by foot, raft, and motorized boat throughout the entire spawning distribution and season. In general, spring run Chinook Salmon surveys were conducted by foot, summer Chinook Salmon by raft, and fall Chinook Salmon by motorized boat. Carcass surveys were conducted weekly and carcasses were measured, sexed, evaluated for marks and tags, scales removed for age determination, and presence of the adipose fin was noted. Heads were removed from adipose fin clipped fish and CWTs were extracted and decoded in a laboratory or office to determine hatchery release information. Females were cut open to determine the proportion of eggs retained by the females. The sampling goal for carcasses was approximately $20 \%$ of the spawning population.

Spring Chinook Salmon redd and carcass surveys were conducted during August through September in all of the spawning areas of the Wenatchee, Entiat, and Methow subbasins (Figure 1). Summer Chinook redd and carcass surveys were conducted from September through November throughout the entire spawning distributions of the Wenatchee, Entiat, Methow, Chelan and Okanogan rivers. Fall Chinook Salmon carcass surveys were conducted from October through the beginning of December in the Hanford Reach of the Columbia River.

## Analysis

The recipient population stray rate for each spawning population was estimated by dividing the annual number of strays by the total annual spawning escapement regardless of fish origin (Bett et al. 2017). This was done for each non-target hatchery program that contributed strays to the recipient population. All non-target hatchery contributions were then summed annually to derive a total recipient population stray rate. Donor strays originated from a large number of hatcheries, so some were grouped with others based on similar regions of the Columbia River basin to facilitate a clear presentation of results. Stray rates were assessed at the tributary, subbasin, and basin levels for spring Chinook Salmon and at the subbasin and basin levels for summer Chinook, fall Chinook, and steelhead. Mean stray rates of Chinook Salmon were calculated for 1999-2018, 2009-2018, and 2014-2018. Mean stray rates of steelhead were 2013-2018 and 2014-2018 because reliable PIT tag analyses were not available prior to 2013. These periods were selected to correspond to modifications and maturity of hatchery programs so that temporal changes could be assessed. In addition, all periods were inclusive of latter years to reveal the potential of long-term influence. Mean stray rates for each period were compared to the management targets of $5 \%$ and $10 \%$. The causes of variation in recipient stray rates for each run type were evaluated by examining the number of hatcheries contributing strays, recipient population size, proximity to non-target hatcheries, and spatial scale.

## Results

The recipient population stray rates for all time periods ranged between $0.02-87.35 \%$ and increased with decreasing spatial scale (Tables 1-3). Recipient stray rates of all taxa at the basin scale were $<3 \%$ and summer Chinook and fall Chinook salmon were $<0.5 \%$ (Table 1). Recipient stray rates in subbasins ranged between $0.07-33.04 \%$ and spring and summer Chinook Salmon exceeded $5 \%$ in some periods in the Entiat and Methow subbasins, but stray rates for all Chinook were $<5 \%$ in the Wenatchee, Okanogan, and Hanford Reach for all periods (Table 2). All steelhead recipient stray rates exceeded $5 \%$ for all periods except for those in the Wenatchee subbasin (Table 2). Recipient stray rates of spring Chinook Salmon in tributaries (the only taxa that met the tributary criteria) ranged between $0.61 \%-87.35 \%$ and only the Icicle, Chiwawa, and Twisp rivers were consistently below $10 \%$, and only the Chiwawa River was consistently below 5\% (Table 3).

Table 1. Mean percent strays of non-target spring Chinook Salmon, summer Chinook Salmon, fall Chinook Salmon and steelhead hatchery-origin recruits to the Upper Columbia River basin for the periods 1999-2018, 2009-2018, and 2014-2018. Steelhead includes the time period from 20132018 and 2014-2018. The percent natural and hatchery-origin fish is a mean calculated over multiple years for each time period.

| Spawn year | $\qquad$ | Natural-origin recruits |  | Target <br> Hatchery-origin recruits |  | Non-target strays Hatchery-origin recruits |  |
| :---: | :---: | :---: | :---: | :---: | :---: | :---: | :---: |
|  |  |  |  |  |  |  |  |
|  |  | Number | Percent | Number | Percent | Number | Percent |
| Spring Chinook |  |  |  |  |  |  |  |
| Mean (1999-2018) | 3,929 | 1,915 | 45.07 | 1,959 | 53.55 | 54 | 1.38 |
| Mean (2009-2018) | 3,735 | 1,440 | 40.92 | 2,236 | 57.79 | 59 | 1.29 |
| Mean (2014-2018) | 2,473 | 1,081 | 45.92 | 1,367 | 53.25 | 25 | 0.83 |
| Summer Chinook |  |  |  |  |  |  |  |
| Mean (1999-2018) | 20,240 | 15,292 | 75.66 | 4,944 | 24.32 | 4 | 0.02 |
| Mean (2009-2018) | 20,353 | 15,698 | 77.35 | 4,647 | 22.62 | 8 | 0.03 |
| Mean (2014-2018) | 19,594 | 16,569 | 84.21 | 3,020 | 15.77 | 5 | 0.02 |
| Fall Chinook |  |  |  |  |  |  |  |
| Mean (1999-2018) | 131,807 | 122,587 | 83.49 | 8,643 | 6.58 | 578 | 0.47 |
| Mean (2009-2018) | 172,991 | 161,663 | 94.01 | 10,644 | 5.62 | 685 | 0.37 |
| Mean (2014-2018) | 192,989 | 181,155 | 93.20 | 11,100 | 6.39 | 734 | 0.41 |
| Steelhead |  |  |  |  |  |  |  |
| Mean (2013-2018) | 4,043 | 2,024 | 48.82 | 1,906 | 48.48 | 113 | 2.70 |
| Mean (2014-2018) | 4,009 | 2,073 | 50.14 | 1,840 | 47.57 | 96 | 2.30 |

Table 2. Mean percent strays of non-target spring Chinook Salmon, summer Chinook Salmon, fall Chinook Salmon and steelhead hatchery-origin recruits to the Hanford Reach of the Columbia River, Wenatchee, Entiat, Methow, Chelan and Okanogan river subbasins of the Upper Columbia River basin for the periods 1999-2018, 2009-2018, and 2014-2018. Steelhead includes the time period from 2013-2018 and 2014-2018. The percent natural and hatchery-origin fish is a mean calculated over multiple years for each time period.

| Spawn year | Subbasin escapement | Natural-origin recruits |  | Target |  | Non-target strays |  |
| :---: | :---: | :---: | :---: | :---: | :---: | :---: | :---: |
|  |  |  |  | Hatchery-origin |  | Hatchery-origin |  |
|  |  | Number | Percent | Number | Percent | Number | Percent |
| Wenatchee Spring Chinook Salmon |  |  |  |  |  |  |  |
| Mean (1999-2018) | 1,740 | 644 | 37.83 | 1,084 | 61.53 | 12 | 0.65 |
| Mean (2009-2018) | 1,876 | 747 | 39.28 | 1,123 | 60.45 | 6 | 0.28 |
| Mean (2014-2018) | 1,198 | 441 | 38.35 | 756 | 61.58 | 1 | 0.07 |
| Entiat Spring Chinook Salmon |  |  |  |  |  |  |  |
| Mean (1999-2018) | 292 | 228 | 79.44 | 14 | 4.91 | 50 | 15.65 |
| Mean (2009-2018) | 320 | 244 | 80.75 | 8 | 2.09 | 68 | 17.16 |
| Mean (2014-2018) | 260 | 239 | 92.62 | 0 | 0.00 | 21 | 7.38 |
| Methow Spring Chinook Salmon |  |  |  |  |  |  |  |
| Mean (1999-2018) | 1,897 | 1,047 | 43.28 | 798 | 52.71 | 52 | 4.01 |
| Mean (2009-2018) | 1,539 | 452 | 35.31 | 1,017 | 59.34 | 70 | 5.35 |
| Mean (2014-2018) | 1,015 | 405 | 46.23 | 570 | 47.13 | 40 | 6.63 |
| Wenatchee Summer Chinook Salmon |  |  |  |  |  |  |  |
| Mean (1999-2018) | 8,695 | 7,427 | 84.92 | 1,234 | 14.69 | 34 | 0.40 |
| Mean (2009-2018) | 7,597 | 6,501 | 86.28 | 1,078 | 13.47 | 18 | 0.25 |
| Mean (2014-2018) | 6,315 | 5,804 | 91.02 | 500 | 8.77 | 10 | 0.20 |
| Entiat Summer Chinook Salmon |  |  |  |  |  |  |  |
| Mean (1999-2018) | 391 | 330 | 83.91 | 19 | 3.64 | 43 | 12.45 |
| Mean (2009-2018) | 447 | 367 | 78.49 | 37 | 7.28 | 43 | 14.24 |
| Mean (2014-2018) | 524 | 439 | 83.33 | 72 | 13.94 | 12 | 2.73 |
| Chelan Summer Chinook Salmon |  |  |  |  |  |  |  |
| Mean (1999-2018) | 796 | 420 | 53.49 | 160 | 13.47 | 216 | 33.04 |
| Mean (2009-2018) | 1,128 | 637 | 58.67 | 319 | 26.94 | 172 | 14.39 |
| Mean (2014-2018) | 1,053 | 624 | 58.26 | 365 | 35.40 | 64 | 6.33 |
| Methow Summer Chinook Salmon |  |  |  |  |  |  |  |
| Mean (1999-2018) | 2,430 | 1,625 | 67.89 | 462 | 19.25 | 343 | 12.87 |
| Mean (2009-2018) | 2,429 | 1,636 | 67.77 | 558 | 23.00 | 235 | 9.23 |
| Mean (2014-2018) | 2,119 | 1,612 | 74.36 | 374 | 20.04 | 132 | 5.60 |
| Okanogan Summer Chinook Salmon |  |  |  |  |  |  |  |
| Mean (1999-2018) | 7,929 | 5,479 | 69.42 | 2,260 | 28.15 | 190 | 2.43 |
| Mean (2009-2018) | 8,752 | 6,529 | 74.55 | 2,112 | 24.11 | 111 | 1.34 |
| Mean (2014-2018) | 9,585 | 8,050 | 85.58 | 1,457 | 13.73 | 77 | 0.69 |
| Hanford Reach Fall Chinook Salmon |  |  |  |  |  |  |  |
| Mean (1999-2018) | 85,180 | 76,806 | 90.47 | 7,820 | 8.83 | 554 | 0.70 |
| Mean (2009-2018) | 111,820 | 101,049 | 91.23 | 10,129 | 8.25 | 643 | 0.53 |
| Mean (2014-2018) | 137,369 | 126,614 | 91.76 | 10,098 | 7.76 | 657 | 0.48 |
| Wenatchee Steelhead |  |  |  |  |  |  |  |
| Mean (2013-2018) | 1,323 | 770 | 59.54 | 541 | 38.10 | 13 | 2.36 |
| Mean (2014-2018) | 1,176 | 736 | 62.31 | 425 | 34.86 | 15 | 2.83 |
| Entiat Steelhead |  |  |  |  |  |  |  |
| Mean (2013-2018) | 395 | 333 | 80.30 | 0 | 0.00 | 63 | 19.70 |
| Mean (2014-2018) | 400 | 350 | 83.21 | 0 | 0.00 | 50 | 16.79 |


| Methow Steelhead |  |  |  |  |  |  |  |
| :---: | :---: | :---: | :---: | :---: | :---: | :---: | :---: |
| Mean (2013-2018) | 1,574 | 674 | 42.89 | 778 | 49.42 | 123 | 7.69 |
| Mean (2014-2018) | 1,587 | 713 | 45.15 | 784 | 49.40 | 90 | 5.45 |
| Okanogan Steelhead |  |  |  |  |  |  |  |
| Mean (2013-2018) | 752 | 248 | 32.69 | 328 | 43.25 | 175 | 24.06 |
| Mean (2014-2018) | 846 | 274 | 30.66 | 378 | 46.29 | 194 | 23.04 |

Table 3. Mean percent strays of non-target spring Chinook Salmon hatchery-origin recruits to tributaries of the Wenatchee, Entiat, and Methow river subbasins of the Upper Columbia River basin for the periods 1999-2018, 2009-2018, and 2014-2018. The percent natural and hatcheryorigin fish is a mean calculated over multiple years for each time period.

| Spawn year | Escapement | Natural-origin recruits |  | Target |  | Non-target strays |  |
| :---: | :---: | :---: | :---: | :---: | :---: | :---: | :---: |
|  |  |  |  | Hatchery-origin recruits |  | Hatchery-origin recruits |  |
|  |  | Number | Percent | Number | Percent | Number | Percent |
| Chiwawa River Spring Chinook Salmon |  |  |  |  |  |  |  |
| Mean (1999-2018) | 922 | 347 | 40.51 | 560 | 57.25 | 16 | 2.24 |
| Mean (2009-2018) | 1,087 | 418 | 37.72 | 663 | 61.66 | 6 | 0.61 |
| Mean (2014-2018) | 708 | 248 | 36.32 | 456 | 63.03 | 3 | 0.65 |
| Nason Creek Spring Chinook Salmon |  |  |  |  |  |  |  |
| Mean (1999-2018) | 371 | 160 | 46.85 | 7 | 4.35 | 204 | 48.80 |
| Mean (2009-2018) | 358 | 145 | 40.42 | 13 | 8.70 | 200 | 50.88 |
| Mean (2014-2018) | 166 | 62 | 40.75 | 26 | 17.40 | 78 | 41.85 |
| White River Spring Chinook Salmon |  |  |  |  |  |  |  |
| Mean (1999-2018) | 82 | 59 | 76.98 | 2 | 1.97 | 21 | 21.05 |
| Mean (2009-2018) | 90 | 68 | 75.70 | 4 | 3.94 | 18 | 20.37 |
| Mean (2014-2018) | 64 | 50 | 74.80 | 7 | 7.88 | 7 | 17.32 |
| Little Wenatchee River Spring Chinook Salmon |  |  |  |  |  |  |  |
| Mean (1999-2018) | 65 | 40 | 67.61 | 0 | 0.00 | 25 | 32.40 |
| Mean (2009-2018) | 68 | 44 | 67.66 | 0 | 0.00 | 24 | 32.34 |
| Mean (2014-2018) | 35 | 26 | 73.33 | 0 | 0.00 | 9 | 26.67 |
| Upper Wenatchee River Spring Chinook Salmon |  |  |  |  |  |  |  |
| Mean (1999-2018) | 99 | 18 | 24.56 | 0 | 0.00 | 81 | 75.77 |
| Mean (2009-2018) | 55 | 11 | 16.10 | 0 | 0.00 | 45 | 84.57 |
| Mean (2014-2018) | 46 | 9 | 13.99 | 0 | 0.00 | 38 | 87.35 |
| Icicle Creek Spring Chinook Salmon |  |  |  |  |  |  |  |
| Mean (1999-2018) | 155 | 20 | 14.38 | 119 | 76.46 | 15 | 9.16 |
| Mean (2009-2018) | 202 | 37 | 14.14 | 143 | 77.21 | 23 | 8.65 |
| Mean (2014-2018) | 172 | 13 | 5.22 | 153 | 92.82 | 6 | 1.95 |
| Peshastin Creek Spring Chinook Salmon |  |  |  |  |  |  |  |
| Mean (1999-2018) | 46 | 29 | 67.47 | 0 | 0.00 | 17 | 32.53 |
| Mean (2009-2018) | 16 | 15 | 72.92 | 0 | 0.00 | 2 | 27.08 |
| Mean (2014-2018) | 6 | 6 | 75.00 | 0 | 0.00 | 1 | 25.00 |
| Entiat River Spring Chinook Salmon |  |  |  |  |  |  |  |
| Mean (1999-2018) | 292 | 229 | 79.69 | 14 | 4.91 | 50 | 15.40 |
| Mean (2009-2018) | 320 | 244 | 80.75 | 8 | 2.09 | 68 | 17.16 |
| Mean (2014-2018) | 260 | 239 | 92.62 | 0 | 0.00 | 21 | 7.38 |
| Methow River Spring Chinook Salmon |  |  |  |  |  |  |  |
| Mean (1999-2018) | 1,219 | 652 | 40.51 | 387 | 39.96 | 180 | 19.53 |
| Mean (2009-2018) | 982 | 209 | 28.89 | 563 | 50.74 | 210 | 20.38 |
| Mean (2014-2018) | 655 | 201 | 40.32 | 375 | 46.55 | 78 | 13.13 |
| Chewuch River Spring Chinook Salmon |  |  |  |  |  |  |  |
| Mean (1999-2018) | 479 | 298 | 54.65 | 116 | 28.41 | 65 | 16.95 |
| Mean (2009-2018) | 394 | 160 | 44.92 | 143 | 33.62 | 91 | 21.46 |
| Mean (2014-2018) | 231 | 128 | 56.49 | 53 | 22.14 | 50 | 21.37 |
| Twisp River Spring Chinook Salmon |  |  |  |  |  |  |  |
| Mean (1999-2018) | 199 | 132 | 59.52 | 56 | 34.35 | 11 | 6.13 |
| Mean (2009-2018) | 163 | 83 | 50.89 | 63 | 39.26 | 17 | 9.86 |
| Mean (2014-2018) | 130 | 75 | 57.49 | 46 | 34.74 | 9 | 7.77 |

Recipient stray rates for each taxa were negatively associated with the abundance of spawners (Figure 2). That is, stray rates increased as total spawner abundance decreased. For example, large populations such as summer Chinook Salmon in the Okanogan and Wenatchee and fall Chinook Salmon in the Hanford Reach had stray rates $<5 \%$. The Chiwawa River was the only spring Chinook population with stray rates $<5 \%$ and is the largest of the spring Chinook spawning aggregates (Table 3). The two largest steelhead populations were the only steelhead populations with stray rates $<10 \%$ (Table 2). In contrast, small populations such as Entiat spring and Summer Chinook, and upper Wenatchee River, Little Wenatchee River, and White River spring Chinook Salmon had high stray rates (Tables 2-3). The highest stray rate was for spring Chinook in the upper Wenatchee River, in which almost all the spawners were stray hatchery fish (Table 3).


Figure 2. The relationship between recipient population size and mean stray rate for summer Chinook Salmon, steelhead, and spring Chinook Salmon.

Proximity to non-target hatcheries or the location of a non-target hatchery relative to the migration sequence of an adult returning to a target location seemed also to influence recipient stray rates. For example, although the spring Chinook spawning aggregate in Nason Creek had a fairly large population size, it had high recipient stray rates from the nearby Chiwawa Acclimation Facility. Similarly, the Chewuch River confluence with the Methow River is between and within one kilometer of two hatcheries that release spring Chinook to the Methow River, one of which also releases spring Chinook to the Chewuch River, and stray rates of spring Chinook Salmon to the Methow and Chewuch rivers were high even though population sizes were among the highest evaluated (Table 3).

The contribution of strays from multiple hatcheries increased the cumulative stray rate in many populations, but in some instances a single hatchery was the primary contributor to stray rate (Figures 3-8). In some cases, an individual hatchery would not result in exceedance of recipient stray targets, but because multiple hatcheries contributed strays, a target was exceeded. At the Basin scale between 2014-2018, all of the strays originated from the Snake River Basin and Middle Columbia River subbasins (Figure 3). At the subbasin scale between 2014-2018, no single spring, summer, or fall Chinook Salmon hatchery contributed $>5 \%$ of the stray rate, but when the contributions of all hatcheries were combined the total stray rate exceeded $5 \%$ (e.g., Entiat and Methow spring Chinook Salmon, and Chelan and Methow summer Chinook Salmon), it was the result of multiple hatchery contributions (Figure 4-5). In contrast, steelhead recipient stray rates in the Entiat and Okanogan had multiple hatcheries exceeding contributions of 5\% stray rate (Figure 6).

Only spring Chinook Salmon met the criteria for evaluating recipient strays at the tributary scale. All of the recipient strays in spawning aggregates of the Wenatchee subbasin originated from within the Wenatchee subbasin. Almost every spawning aggregate in tributaries of the Wenatchee subbasin exceeded $10 \%$ because of strays from the Chiwawa River (Figure 7). In Entiat and Methow river spawning aggregates, no single hatchery exceeded the 5\% or 10\% target criteria, but when all hatcheries were combined, the targets were exceeded (Figure 8).


Figure 3. Mean percent hatchery stray Chinook Salmon and steelhead observed in the Upper Columbia River Basin from other regions of the Columbia River Basin from 2014 to 2018.


Figure 4. Mean percent hatchery stray spring Chinook Salmon observed in the Upper Columbia River subbasins from other regions of the Columbia River Basin from 2014 to 2018. The management target is <5\%.


Figure 5. Mean percent hatchery stray summer and fall Chinook Salmon observed in subbasins of the Upper Columbia River basin from 2014 to 2018. The management target is <5\%.


Figure 6. Mean percent hatchery stray steelhead observed in subbasins of the Upper Columbia River basin from 2014 to 2018. The management target is <5\%.


Figure 7. Mean percent hatchery stray spring Chinook Salmon observed in tributaries of the Wenatchee subbasin from 2014 to 2018. The management target is < $10 \%$.


Figure 8. Mean percent hatchery stray spring Chinook Salmon observed in the Entiat River and Methow River and tributaries (Chewuch and Twisp rivers) from 2014 to 2018. The management target is < $10 \%$ except for the Entiat River which is $<5 \%$.

Spatial variation in mean recipient stray rates was substantially higher than temporal variation in mean recipient stray rates although annual variation in both could be quite high for some taxa and locations. Spatial variation ranged from 0.02-87.35\%, a 4,368-fold difference, across all taxa and maximum temporal variation ranged from 0.07-0.65 within a taxa a 9.29 -fold difference (Wenatchee spring Chinook Salmon; Table 2). Recipient stray rates were relatively stable for most populations particularly at large spatial scales and when changes occurred most of them decreased between 1999 and 2018 (Tables 1-3). There were some notable decreases in recipient stray rates between 1999 and 2018 (e.g., Entiat and Chelan Summer Chinook, Icicle Creek spring Chinook) and these were likely the result of reductions in hatchery program size, tributary acclimation, other program modifications, and possibly reductions in donor stray rates (Tables 2-3).

## Discussion

It is clear that recipient population stray rates exceeded management targets (e.g., >5$10 \%$ ) in: 1) many upper Columbia Basin populations of spring Chinook Salmon and steelhead and 2) some summer Chinook Salmon at subbasin and tributary scales, but fall Chinook was lower than management targets. In some cases, this exceedance is the result of many different
hatcheries contributing spawners to a non-target population, while in others it is the result of a single hatchery. Most management targets are structured around the stray contribution of single hatcheries (e.g., Hillman et al. 2018), but cumulative influences of all hatcheries are more biologically relevant because they represent the total spawning population. The complexity of managing strays from multiple hatcheries, some of which are in different states and operated by different organizations with different objectives, is a difficult socio-political challenge. For example, should strays from harvest augmentation hatcheries be considered similarly as those produced to aid in species recovery or should greater leeway be given to hatcheries used to recover species? Should stray rates be managed based upon donor stray rates (e.g., \% of a hatchery population that strays) or the total number of strays contributed to a recipient population?

Recipient population straying has the potential to reduce between-population genetic diversity at the levels that we observed in this study (e.g., $>5-10 \%$ ). However, this assumes that stray fish contribute towards natural production. Relative reproductive success studies indicate that hatchery-origin fish generally produce fewer offspring than natural-origin counter parts (Williamson et al 2010. Ford et al. 2016). Genetic risks of straying are increased if strays successfully spawn and nullified if they do not spawn. This can be evaluated by examining whether female strays void their eggs, an index of spawning success. Stray fish that retain their eggs and die, pose low genetic risks to recipient populations. Upper Columbia Chinook Salmon have very high rates of egg voidance often exceeding $95 \%$, suggesting that they successfully spawned in the areas where carcasses were collected (Murdoch et al. 2009; Richards and Pearsons 2019). An early evaluation of the hatchery effects on genetic diversity in the upper Columbia Basin did not reveal decreases in genetic diversity (Hillman et al. 2019). A more current genetic evaluation that incorporates the time periods of this study is currently in progress.

Recipient population stray rates can be managed in three primary ways (Bett et al. 2017). The first is to manage donor population stray rate through improved fish-culture approaches. This might include techniques to improve imprinting such as raising fish on natal target waters to the greatest extent practicable during the time of imprinting (Dittman et al. 2015, Pearsons and O'Connor 2021). However, even low donor-stray rates can result in high recipient population stray rates if the hatchery program is large and the recipient population is small. Furthermore, donor population stray rates can be influenced by factors other than fish culture such as migration and spawning habitat quality (Cram et al. 2012; Bond et al. 2017; Pearsons and O'Connor 2021), so improvements in fish culture alone may not result in desired management outcomes. The second approach is to manage the number of adults that could potentially stray by reducing hatchery program size, removal at weirs, and removals through harvest. The adult removal approach may not be effective if the intent is to supplement a population because available control measures are often downstream of the target population and it is unclear which fish should be removed and which fish should be allowed to spawn. This may be the case for most listed species. The third approach is to increase natural escapement because escapement is an important factor influencing stray rates. Most large populations in this study met targets and small populations typically did not. Escapements are influenced by many factors beyond the specific hatchery; for example, harvest and natural production as influenced by factors such as ocean conditions, and habitat conditions. In short, multiple factors influence recipient population stray rates, and changes in hatchery practices alone may not achieve dual objectives of increasing abundance and keeping recipient stray rates below target levels.

Trade-offs will have to be made in some cases where hatchery improvements such as improvements in imprinting are limited or unfeasible. For example, acclimation sites are used to cause fish to return to particular locations; however fish that are transported from a downstream hatchery and acclimated at remote sites may stray at higher rates (Pearsons and O'Connor 2021) than those that are not transported. In addition, transportation is likely necessary to get fish to return to the target location for supplementation programs. The value of returning fish to a target location will have to be weighed against the cost of straying. In some cases, the supplementation value will be lower than the cost of straying necessitating a reduction in hatchery-program size to achieve management optima.

Some straying of hatchery-origin fish may occur between spawning aggregates but because the tributaries were part of the same genetic management zone, they were not considered a genetic concern. For example, straying of spring Chinook Salmon occurred between the Methow and Chewuch River, but because they were part of the same genetic management zone, the genetic strays were acceptable to managers from a genetic perspective. However, there was concern that these strays did not return to the target location and therefore were a demographic shortfall to the target population.

In some cases, high recipient stray rates may be keeping a population from extinction. For example, the Nason Creek and White River spring Chinook Salmon spawning aggregates regularly experience recipient stray rates of $30-50 \%$. The upper Wenatchee River is likely a sink population because it has regularly comprised over $85 \%$ strays and none of the progeny of naturally produced fish that spawned there returned to spawn there (Ford et al. 2015). It is possible that some genetic diversity may have been lost from this high gene flow; however, it is also possible that these contributions have maintained some level of genetic differentiation as opposed to losing the population altogether because of unsustainably low survival rates.

In summary, recipient population stray rates of Salmon and steelhead varied dramatically in the upper Columbia Basin and some exceeded management targets at subbasin and tributary scales. In some cases, this was the result of many different hatcheries contributing spawners to a non-target population while in others it was the result of a single hatchery. Targets for recipientstray rates were never exceeded in large recipient populations but were often exceeded in small recipient populations. This was likely one of the reasons why recipient population stray rates increased with decreasing spatial scale because the smaller the scale the lower the population size. Difficult management trade-offs between increasing abundance and minimizing recipient stray rates to within acceptable limits are likely. Some solutions to minimize recipient stray rates will likely involve a combination of changes to hatchery, harvest, and habitat management.

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# Examining the Genetic Structure of Upper Columbia Summer/Fall Chinook Salmon and Evaluating the Effects of the Supplementation Program 

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#### Abstract

We examined baseline (1982-1994) and contemporary (2017-2018) summer and fall Chinook Salmon (Oncorhynchus tshawytscha) from the Upper Columbia River Watershed to determine if hatchery supplementation programs have had any impacts on the genetic diversity and structure of these populations. Baseline collections included both hatchery- and natural- origin samples where available. Contemporary collections exclusively consisted of samples collected at broodstock collection facilities; their origin (hatchery or natural) was only sometimes known. Summer Chinook Salmon populations with paired baseline and contemporary samples included the Methow River, the Wenatchee River, and the Okanogan River. Populations with only contemporary samples included Chelan Falls, Entiat National Fish Hatchery, and Wells Fish Hatchery. Fall Chinook Salmon were represented by collections from the Hanford Reach spawning grounds and Priest Rapids Hatchery. Measures of genetic diversity (allelic richness, heterozygosity, linkage disequilibrium, and effective number of breeders) showed little differentiation among baseline and contemporary populations for either summer or fall Chinook, suggesting that hatchery programs have not led to a decrease in genetic diversity. There was a general pattern where $F_{\text {ST }}$ was higher among baseline than contemporary collections suggesting that genetic drift and homogenization among stocks has occurred over time. Despite these patterns, pairwise comparisons of $F_{\text {ST }}$ were generally statistically non-significant both for baseline and contemporary collections. Similar to previous evaluations, there appears to be little evidence for neutral genetic divergence between contemporary hatchery programs in the upper Columbia watershed and baseline samples collected in the late 1980s and early 1990s. The large population sizes of summer and fall Chinook Salmon relative to the hatchery program sizes in the upper Columbia basin, low recipient population stray rates in natural populations, and the management strategies that were implemented to reduce genetic risk all likely contribute to the lack of neutral genetic change. This evaluation did face two limitations: first, we were not able to evaluate potential differentiation among contemporary hatchery and natural origin individuals due to lack of data on individual origin; second, we were not able to evaluate potential shifts in adaptive genetic diversity using genetic techniques and it is possible for adaptive genetic diversity (i.e., run-timing, age at maturity) to change in response to selection (i.e., domestication) while neutral genetic diversity remains the same. While adaptive genetic variation was not directly monitored, phenotypic metrics measured as part of other portions of the monitoring plan can serve as a proxy for adaptive genetic variation.


## Introduction

One of the main concerns associated with using artificial propagation to supplement natural populations and to increase harvest is the reduction in long-term fitness associated with interbreeding and loss of local adaptation in naturally spawning populations (Ford 2002, Mobrand et al. 2005, Paquet et al. 2011). Interbreeding can result in changes to the genetics of progeny and result in changes to the productivity of populations. Interbreeding can be intentional and substantial such as when the objective of the hatchery program is to increase natural production (Williamson et al. 2010, Ford et al. 2015a). Interbreeding can also be an unintentional byproduct of operating a hatchery program when hatchery-origin fish stray to nontarget spawning populations (Keefer and Caudill 2014, Ford et al. 2015b, see Pearsons and O'Connor and Pearsons and Miller chapters in this report). Despite the long-term risks of interbreeding between hatchery- and natural-origin fishes, we are not aware of standardized methods for long-term monitoring of the effects of hatcheries on naturally spawning populations.

The long-term fitness of natural populations is related to their genetic diversity. However, hatchery programs select a subset of individuals whose probability of passing on genetic material to the next generation is increased by reducing mortality associated with freshwater rearing and development. This subset is often a relatively small number of individuals that then produce a large number of adult offspring and thus these programs can change allele frequencies and reduce effective population size (Ryman and Laikre 1991). Therefore, it is important to monitor the genetic status of the natural populations to determine if there are signs of changes in genetic distance among populations, changes in allele frequencies, linkage disequilibrium, and to estimate effective population size.

## Background

Construction of the Grand Coulee Dam in 1939 eliminated upstream migration of anadromous salmon past the dam. To mitigate the effects of this habitat loss, the Grand Coulee Fish Maintenance Project (GCFMP) was initiated. This project aimed to maintain fish runs by improving salmon habitat and establishing hatchery operations (Fish and Hanavan 1948). From 1939 to 1943, all adult fish passing Rock Island Dam were collected and either spawned artificially at U.S. Fish and Wildlife Service (USFWS) hatcheries on the Wenatchee or Methow rivers, or transported and allowed to spawn naturally in fenced reaches of the Wenatchee or Entiat rivers. "Early-run" fish, presumably spring run, were separated from "late-run fish", presumably summer and fall run, however no effort was made to separate summer or fall components of the run (Waknitz et al. 1995). There are relatively few 6-year old adult Chinook Salmon in the Columbia River ( $<1 \%$ ), and the Rock Island Dam collection program lasted five years. As a result, nearly all contemporary late-run Chinook Salmon above Rock Island Dam are progeny of mixed Chinook Salmon stocks and mixed summer and fall run-timings collected during this program (Waknitz et al. 1995).

The Chinook Salmon population below Rock Island Dam did not experience the same machinations associated with the GCFMP. The primary spawning and rearing area of the Chinook Salmon spawning below Rock Island Dam is the Hanford Reach of the Columbia River which spans $\sim 51$ miles from the Priest Rapids dam (constructed in 1956) to the town of

Richland, WA. Early operations of the dam resulted in large variations in discharge, which lead to dewatering sections with redds, and stranding of Chinook Salmon individuals (Langshaw et al. 2018). Ultimately, the Hanford Reach Fall Chinook Protection Program Agreement (HRFCPPA) was implemented to mitigate and further protect Hanford fall Chinook. Fall Chinook Salmon that spawn in the upper Columbia River Basin are considered to be one of the few remaining "robust" stocks in the basin (Langshaw et al. 2018). The Hanford Reach of the upper Columbia River supports the largest spawning population of fall Chinook Salmon in the Pacific Northwest (Huntington et al. 1996, Dauble and Watson 1997, Harnish et al. 2012, Langshaw et al. 2018)).

The Public Utility Districts (PUDs) of Chelan, Douglas, and Grant counties agreed to implement hatchery programs to mitigate for unavoidable mortality caused by construction and operation of hydroelectric projects on the Columbia River. Initial hatchery program sizes were negotiated with fisheries managers and later refined using estimates of mortality caused by hydropower projects and survival of hatchery fish. Committees consisting of a representative from the USFWS, National Marine Fisheries Service (NMFS), Washington State Department of Fish and Wildlife (WDFW), Yakama Nation (YN), Confederated Tribes of the Colville Reservation (CCT), and each PUD were formed to oversee the implementation of PUD hatchery programs. These hatchery committees were tasked with developing long-term plans for monitoring the hatchery programs and with adaptive management of the programs as new information became available.

## Hanford fall Chinook at Priest Rapids Hatchery

Natural-origin fall Chinook Salmon produced in the Hanford Reach emerge from the substrate in the spring and rear there until outmigration in the summer. Egg-to-fry survival and egg-to-pre smolt survival of natural production within the Hanford Reach have been estimated to be $\sim 71 \%$ and 40.2-63.4\%, respectively (Harnish et al. 2012, Oldenburg et al. 2012, Harnish 2017).

The Priest Rapids Hatchery (PRH) was constructed at the top end of the Hanford Reach to mitigate for losses associated with the inundation of the portions of the Columbia River caused by the construction of Priest Rapids (1959) and Wanapum dams (1963). The PRH has evolved from a spawning channel initially constructed downstream from Priest Rapids Dam in 1963 to a state-of-the-art hatchery facility completed in 2014. While operating as a spawning channel from 1963 through 1971, summer/fall Chinook Salmon adults trapped in the east ladder of Priest Rapids Dam were used as broodstock. This practice was generally ineffective at producing juveniles because of a variety of factors leading to mortality of both adult broodstock and in eggs deposited in redds. Artificial propagation of fall Chinook Salmon at the site began in 1972 with the collection and spawning of broodstock derived from adults returning to the spawning channel. In 1978, use of the spawning channel was terminated and all fish released from PRH were derived from artificial production at that facility (Chapman et al. 1994). A major rebuild of the facility was completed in 2014 including a renovated trapping facility, new adult holding ponds, new adult sorting capabilities, a new incubation building, 30 new raceways, and five renovated acclimation ponds.

The annual release of fall Chinook Salmon smolts from PRH has ranged considerably since the initial release of roughly 150,625 smolts from the 1977 brood year to over roughly 10.30 million from the 1982 brood year. From 1977 to 2013 the release goal of the PRH program was 5 million subyearling smolts and additional production was produced for USACE. In 2013, the target number of fish to release at PRH was revised to 7,299,504 (5,599,504 combined with the ongoing USACE's John Day mitigation of $1,700,000$ smolts). In addition to production released by PRH, the United States Army Corps of Engineers (USACE) also released subyearling fall Chinook Salmon from Ringold Springs Hatchery (RSH) into the lower end of the Hanford Reach beginning in 1994. The smolts released by RSH were derived from adult salmon returning to Bonneville Hatchery prior to 2009 and PRH during years afterwards to collect eggs sufficient to release 3.5 million subyearling smolts. Thus, a total annual release goal of $10,799,504$ hatchery reared subyearling smolts was planned for the Hanford Reach from 2014 to present.

The age at maturity for naturally produced fish in the Hanford Reach varies between age1 mini-jack and age- 6 adults: albeit recoveries of age- 1 and 6 fish are generally rare. The abundance of mini-jacks maturing as age-1 males is currently not known. Age-2 male fall Chinook Salmon (jacks) return to the Hanford Reach after spending roughly one year in the ocean. The majority of the natural-origin adults return after spending three to four years in the ocean (age-4 and 5). A small portion, typically less than $2 \%$, will spend up to five years in the ocean and return as age-6. Adults return to the mouth of the Columbia River between August and October and spawn in large cobble substrate between October and December (Langshaw et al. 2018, Richards and Pearsons 2019).

Wenatchee Summer Chinook
The goal of summer Chinook Salmon supplementation in the Wenatchee Subbasin is to use artificial production to replace Chinook Salmon lost because of mortality at Priest Rapids, Wanapum, and Rock Island dams, while not reducing the natural production or long-term fitness of the extant summer Chinook Salmon population in the basin. The Rock Island Fish Hatchery Complex began operation in 1989 under funding from Chelan PUD and subsequently Grant PUD began cost-sharing the program in 2012. The Complex operated originally through the Rock Island Settlement Agreement, but since 2004 has operated under the Rock Island Anadromous Fish Agreement and Habitat Conservation Plan (HCP) as well as the Priest Rapids Project Salmon and Steelhead Settlement Agreement.

Adult summer Chinook Salmon are collected for broodstock from the Wenatchee River run-at-large at the right- and left-bank traps at Dryden Dam, and at Tumwater Dam if weekly quotas cannot be achieved at Dryden Dam. Before 2012, the goal was to collect up to 492 natural-origin adults for the Wenatchee program for an annual release of 864,000 yearling smolts. In 2011, the Hatchery Committees reevaluated the amount of hatchery compensation needed to achieve no net impact (NNI). Based on that evaluation, the smolt-production goal of the program was reduced. The current goal (beginning with brood year 2012) was to collect up to 274 adult natural-origin adults for an annual release of 500,001 yearling smolts. The 500,001 smolts were the combined Grant PUD and Chelan PUD smolt production target, with Chelan PUD's obligation at 318,000 and Grant PUD's obligation at 182,001. Broodstock collection
occurred from about 1 July through 15 September with trapping occurring up to 24 hours per day, seven days a week at Dryden Dam and up to 16 hours per day, three days per week at Tumwater Dam. If natural-origin broodstock collection fell short of expectation, hatchery-origin adults were collected to meet the collection quota.

Adult summer Chinook Salmon are spawned at Eastbank Fish Hatchery, where the majority of juveniles are reared in raceways, and a portion in circular tanks. Juveniles are transferred from the hatchery to Dryden Acclimation Pond on the Wenatchee River, in March of each release year, and they are released from the pond volitionally beginning mid-April and pushed out by the end of April.

Before 2012, the production goal for the Wenatchee summer Chinook salmon supplementation program was to release 864,000 yearling smolts into the Wenatchee River at ten fish per pound. Beginning with the 2012 brood, the revised production goal is to release 500,001 yearling smolts into the Wenatchee River at 18 fish per pound. Targets for fork length and weight are $163 \mathrm{~mm}(\mathrm{CV}=9.0)$ and 45.4 g , respectively. Over $95 \%$ of these fish are marked with CWTs. In addition, since 2009, about 20,000 juveniles were PIT tagged annually.

## Entiat National Fish Hatchery Summer Chinook

Entiat National Fish Hatchery (NFH) operates a segregated harvest program that currently produces summer Chinook Salmon for commercial, sport, and tribal harvest while attempting to minimize adverse impacts to the environment. The United States Fish and Wildlife Service (USFWS) operates the facility with funds provided by the Bureau of Reclamation (BOR). Summer Chinook Salmon from this program are not intended to spawn naturally, supplement, or support any summer Chinook Salmon populations. The release target of 400,000 yearling adipose-clipped summer Chinook Salmon was established after discussion with the relevant co-managers and is described in the U.S. v. Oregon Columbia River Management Agreement.

Entiat NFH is a mitigation hatchery originally established by the Grand Coulee Fish Maintenance Project (1937) and began operations in 1942 as partial mitigation for the loss of anadromous fish production due to the construction and operation of Grand Coulee Dam. Since 1942, Entiat NFH has released a variety of species from multiple stocks however spring and summer Chinook Salmon (Oncorhynchus tshawytscha) have been the primary stocks reared to meet mitigation requirements. The hatchery began rearing spring Chinook Salmon that originated from mixed upriver stocks intercepted at Rock Island Dam in 1942 and 1944. No spring Chinook Salmon were reared from 1945 to 1974. In 1974, spring Chinook Salmon production resumed and egg sources included: Cowlitz River (1974), Carson NFH (1975-1982), Little White Salmon NFH (1976, 1978, 1979, 1981), Leavenworth NFH (1979-1981, 1994), and Winthrop NFH (1988). Returning adults that voluntarily entered the hatchery were the primary broodstock in 1980 and from 1982 to 2006. The spring Chinook Salmon rearing program was terminated in 2006 to reduce the impact of Entiat NFH-origin spring Chinook Salmon on ESAlisted natural-origin spring Chinook Salmon in the Entiat River. The last on-station release of spring Chinook Salmon to the Entiat River occurred in 2007 and the last adults returned in 2010.

In the fall of 2009, the hatchery began a new program propagating summer Chinook Salmon with broodstock captured at Wells Fish Hatchery. Wells Fish Hatchery (Wells stock) was selected as the broodstock because they are genetically part of the upper Columbia River summer Chinook Salmon stock (Kassler et al. 2011). Additionally, a genetic evaluation of the existing natural-origin stock in the Entiat River determined the population to not be genetically distinct from the Wells stock or the upper Columbia River summer Chinook Salmon population (Smith et al. 2011). Entiat NFH reared and released juvenile summer Chinook Salmon into the Entiat River from 1941-1964, and in 1976 (Mullan 1987). Summer Chinook Salmon egg sources have included: mixed upriver stocks intercepted at Rock Island Dam (1939-1943), Methow River (1944), Carson NFH (1944), Entiat River (1946-1965), Spring Creek NFH (1964), and Wells Hatchery (1974, 2009-2013). Adult summer Chinook Salmon returning to Entiat NFH have been the primary brood source since 2014 (Fraser et al. 2020).

## Chelan Falls Hatchery Summer Chinook

The Chelan Falls summer Chinook program is a segregated harvest program. Adult returns spawn in the lower Chelan River; however, there is no escapement goal for natural spawning. Chelan Falls summer Chinook are available for harvest in ocean and Columbia River commercial, tribal, and recreational fisheries. The Chelan Falls summer Chinook program (formerly the Turtle Rock program) included the production of 200,000 fish for No Net Impact (NNI) compensation for passage mortalities associated with Rocky Reach Dam and a 400,000 subyearling/yearling program for compensation for lost spawning habitat as a result of the construction of Rocky Reach Dam. In 2011, as part of the periodic recalculation of NNI for Rocky Reach Dam (inundation), the previous 200,000 NNI program was reduced to 176,000 fish. This reduced the combined Chelan Falls summer Chinook production from 600,000 to 576,000 beginning with the 2012 brood.

The original program consisted of both subyearling (normal and accelerated groups) and yearling releases. Subyearlings were transferred to Turtle Rock Acclimation Facility for acclimation in May. These fish were released in June after about 30 days of acclimation on Columbia River water. The production goal of this program was to release $1,620,000$ subyearling summer Chinook ( 810,000 normal and 810,000 accelerated subyearlings) into the Columbia River. In 2010, the subyearling program was converted to a 400,000 -yearling program. The production goal of the yearling program was to release 200,000 summer Chinook smolts into the Columbia River from the Turtle Rock Acclimation Facility. Beginning with the 2006 brood year, yearling summer Chinook were acclimated at both Turtle Rock Acclimation Facility and the Chelan River net pens. With the conversion of the subyearling program to a yearling program and the reduction of the NNI component to 176,000 , the current goal is to release 576,000 yearling summer Chinook smolts ( 176,000 from the NNI program plus 400,000 from the converted subyearling program). Beginning in 2012, the 576,000 yearlings are acclimated overwinter at the Chelan Falls Acclimation Facility on Chelan River water. In 2012, the Turtle Rock program officially became the Chelan Falls summer Chinook program and all fish were overwinter-acclimated at the Chelan Falls Acclimation Facility.

Before 2012, broodstock were collected at the Wells Dam volunteer trap (WDVT). Summer Chinook were spawned at Wells Fish Hatchery and fertilized eggs were then transferred
to Eastbank Fish Hatchery for hatching and rearing. In 2012, adults were collected at the WDVT and then transferred to Eastbank Fish Hatchery for spawning, hatching, and rearing. Beginning in 2013, broodstock collection was initiated at the Eastbank Fish Hatchery Outfall. With returns to the Outfall diminishing, a pilot broodstock collection program was initiated in 2016 at the outlet structure of the water conveyance canal for the Chelan Tailrace Pump Station (Chelan Falls Canal Trap) and continued through 2018. Concurrently, while collection of broodstock from the Chelan Falls Canal Trap was evaluated, the Entiat National Fish Hatchery and WDVT were used as backup broodstock collection sites. Beginning in 2019, a weir was installed in the habitat channel adjacent to the conveyance canal as another pilot location for broodstock collection. The WDVT was used once again as a backup to this pilot effort. The Chelan Falls summer Chinook program collects hatchery-origin broodstock; approximately 390 adults are necessary for the program. Over $90 \%$ of yearling summer Chinook have been marked with CWTs and 85 to $100 \%$ were ad-clipped. In addition, juvenile summer Chinook were PIT tagged within each of the circular and standard raceways.

## Wells Fish Hatchery Summer Chinook

The goal of the summer Chinook artificial propagation program at Wells Hatchery is to mitigate for the loss of summer Chinook salmon adults and associated fishing opportunity (harvest) that would have been available in the region in the absence of the construction of the Wells Hydroelectric Project (Wells Project). Wells Hatchery began operation in 1967 and is located on the Columbia River west bank of the Wells Dam tailrace. This facility was constructed and is funded by Douglas PUD to mitigate for loss of summer Chinook salmon spawning habitat inundated by Wells Dam. Originally built as a spawning channel, it was reprogrammed to serve as an extended rearing facility in 1977. Since brood year 1993, the program has included two components: 1) a yearling program that releases 320,000 smolts (at 10 fish per pound) annually, and 2) a subyearling program that releases 484,000 fish (at 50 fish per pound) annually, directly to the Columbia River in mid-April (yearlings) and late-May (subyearlings).

The Wells Hatchery summer Chinook program is a segregated harvest program, although up to $10 \%$ of the broodstock may be composed of natural-origin fish. Adult returns are not intended to spawn naturally; therefore, there is no escapement goal for natural spawning areas. However, the goal for the stray rate of Wells Hatchery summer Chinook to natural spawning areas is to comprise less than $5 \%$ of the naturally spawning population. Thus, management of adult returns is necessary to meet program objectives. Wells Hatchery summer Chinook are available for harvest in ocean and Columbia River commercial, tribal, and recreational fisheries. Also, other summer Chinook hatchery programs, including Turtle Rock, Chelan Falls, Entiat, and Yakima, have relied on returns from the Wells Hatchery program. Returns to the Wells Hatchery in excess of broodstock needs for this program or other programs, are collected and surplussed by WDFW to authorized recipients, primarily tribes.

The Wells Hatchery summer Chinook program collects hatchery-origin broodstock with up to $10 \%$ natural-origin broodstock at the Wells Hatchery volunteer channel. Approximately 602 adults are necessary for the two program components, with 230 adults needed for the yearling program, and 372 adults for the subyearling program. Broodstock collection has
historically began on July 1, and continued as late as August 31, but typically most broodstock are collected in July. The spawning facilities at Wells Hatchery are integrated into the broodstock-holding facilities, allowing the sorting of broodstock for sexual maturity followed immediately by spawning. Fertilization, incubation, and rearing also occur at the Wells Hatchery, with final rearing in very large ponds.

For each brood year included in this analysis, 100 percent of the summer Chinook produced were adipose clipped and marked with CWTs (multi-year mean CWT mark-retention rate of $97.6 \%$ for subyearlings and $95.8 \%$ for yearlings), and 6,000 of the subyearlings were PIT tagged. Beginning in brood year 2018, 5,000 of the yearlings were also PIT tagged. Most (63\%) Wells Hatchery subyearlings return as age-4 adults, and most (51\%) yearlings as age-5 adults.

## Methow Summer Chinook

The original goal of summer Chinook Salmon supplementation in the Methow Basin was in part to use artificial production to replace Chinook Salmon lost because of mortality at Wells, Rocky Reach, and Rock Island dams, while not reducing the natural production or long-term fitness of summer Chinook Salmon in the basin. The Rock Island Fish Hatchery Complex began operation in 1989 under funding from Chelan PUD. The Complex operated originally through the Rock Island Settlement Agreement, but from 2004 to 2012 operated under the Rock Island and Rocky Reach HCPs. Beginning with broodstock collection in 2012, Grant PUD took over funding and operation of the summer Chinook Salmon supplementation program in the Methow River basin. Grant PUD constructed a new overwinter acclimation facility adjacent to the Carlton Acclimation Pond and the first release of fish from this facility was in 2014. The first fish that were overwinter acclimated in the facility were released in 2015 . The new facility includes eight, 30 -foot diameter dual-drain circular tanks.

Presently, adult summer Chinook Salmon are collected for broodstock from the run-atlarge at the east-ladder trapping facility at Wells Dam. Before 2012, the goal was to collect up to 222 natural-origin adults for the Methow program. In 2011, the Hatchery Committees reevaluated that amount of hatchery compensation needed to achieve NNI. Based on that evaluation, the goal of the program was revised. The current goal (beginning with brood year 2012) is to collect up to 102 natural-origin adults for the Methow program. Broodstock collection occurs from about 1 July through 15 September with trapping occurring no more than 16 hours per day, three days a week. If natural-origin broodstock collection falls short of expectation, hatchery-origin adults can be collected to make up the difference.

Adult summer Chinook Salmon were spawned and progeny reared at Eastbank Fish Hatchery. Before the initiation of overwinter acclimation with juveniles from the 2013 brood year, juveniles were transferred from the hatchery to Carlton Acclimation Pond in March. Beginning with brood year 2013, juveniles have been transferred to the Carlton Acclimation Facility in October or November and released from the new facility the following spring in midApril to early May.

Before 2012, the production goal for the Methow summer Chinook Salmon supplementation program was to release 400,000 yearling smolts into the Methow River at 10
fish per pound. Beginning with the 2012 brood, the revised goal is to release 200,000 yearling smolts at 13-17 fish per pound. Targets for fork length and weight are $163 \mathrm{~mm}(\mathrm{CV}=9.0)$ and 45.4 g , respectively. Over $90 \%$ of these fish were marked with CWTs. In addition, since 2009, 5,000 juveniles have been PIT tagged annually.

## Okanogan Summer Chinook

The Chief Joseph Hatchery (CJH) program was designed to increase the abundance, productivity, distribution, and diversity of naturally spawning populations of summer/fall Chinook salmon (Oncorhynchus tshawytscha) in the Okanogan River and in the Columbia River above Wells Dam. Program operations began in 2013 and consists of integrated and segregated summer/fall Chinook programs that release up to 2 million smolts to meet conservation and harvest objectives to partially fulfill Federal and Public Utility District mitigation obligations for Columbia River Dam impacts to anadromous salmonids. The integrated summer/fall Chinook program expanded on, and now incorporates the previous Chelan PUD and WDFW Similkameen Pond program. The previous Similkameen program was in operation from 1989 to 2012 and released up to 576,000 smolts that originated from a natural origin brood collected at Wells Dam. Since 2010, the Colville Tribes have been collecting brood at the confluence of the Okanogan and Columbia using a purse seine, as a means of avoiding the previous Methow and Okanogan composite brood collection at Wells Dam. The integrated summer/fall Chinook program uses a high proportion of Okanogan natural-origin broodstock while management actions (e.g., selective harvest and weir removals) maintain a low proportion of hatchery-origin spawners to achieve population objectives for conservation (i.e. PNI $>0.67$; $\mathrm{pHOS}<0.30$ ) that ensure that the natural environment has the majority of influence on local adaptation. The smolt release targets at full program for the integrated program are 800,000 yearling smolts from the Omak and Similkameen acclimation ponds and 300,000 subyearlings from the Omak acclimation pond. The integrated program is $100 \%$ adipose fin clipped and coded-wire tagged with 10,000 PIT tags. The segregated summer/fall Chinook program is intended for harvest and uses primarily first generation returns from the integrated program to minimize multi-generation hatchery affects. The segregated program smolt release goals are 500,000 yearlings and 400,000 subyearlings from the Chief Joseph Hatchery on the Columbia River (upstream of the confluence with the Okanogan River). The segregated program is $100 \%$ adipose fin clipped and includes 200,000 coded-wire tags and 10,000 PIT tags.

## Objectives

In response to the need for evaluation of the supplementation program, both a monitoring and evaluation plan (Murdoch and Peven 2005) and the associated analytical framework (Hays et al. 2006) were developed for the Habitat Conservation Plans Hatchery Committee through the joint effort of the fishery co-managers (CCT, NMFS, USFWS, WDFW, and YN) and Chelan County, Douglas County, and Grant County PUDs. This plan was updated in 2019 (Hillman et al. 2020) and includes twelve objectives to be applied to various species assessing the impacts of hatchery operations mitigating the operation of Rock Island and Rocky Reach Dams. This report pertains to Upper Columbia summer and fall Chinook Salmon and the Chinook Salmon supplementation program as addressed by Objective 7, evaluating population genetics to determine if genetic diversity, population structure, and effective population size have changed
in Upper Columbia summer and fall Chinook Salmon as a result of the conservation and safetynet hatchery programs and assess genetic changes of hatchery-origin returns.

To address Objective 7, the WDFW Molecular Genetics Lab (MGL) obtained baseline and contemporary tissue or genotype collections and samples, surveyed genetic variation with SNP markers using our standard laboratory protocols, and calculated the relevant genetic metrics and statistics. Genotypes from baseline and contemporary hatchery and natural origin collections were analyzed to evaluate differences between baseline and contemporary and between hatchery and natural origin collections. In most cases, baseline sample collections consisted of the oldest samples available from each population and contemporary sample collections were from spawn years 2017 and 2018.

## Methods

## Sample collections

Baseline sample collections consisted of the oldest samples found for each population. Baseline collections had all been used in previous monitoring and evaluation projects (Blankenship et al. 2007, Small et al. 2007, Kassler et al. 2011), except for Hanford fall Chinook Salmon. Baseline for the Hanford fall Chinook Salmon came from scales from 1982 and 1988 that were found in the WDFW scale lab archive. Although some salmon were externally marked in the 1980s, marks were typically made for research purposes and could have been either hatchery- or natural- origin. Origin of Hanford fall Chinook Salmon samples was inferred from sample location. Natural-origin fish were thought to enter the Priest Rapids hatchery trap at low levels, therefore the hatchery-origin baseline was drawn from among those fish that swam into the trap because they were likely of hatchery-origin. Hatchery-origin fish were thought to make up a small proportion of the fish on the spawning grounds, and when they did, it was likely they were mainly found near the upstream hatchery or dam areas. Thus, Hanford fall Chinook Salmon samples taken from carcasses found near Locke Island and White Bluffs spawning grounds were considered natural-origin.

Contemporary Chinook Salmon samples consisted of broodstock spawned in 2017 and 2018 from each of the upper Columbia Chinook Salmon programs. All other genotypes from contemporary Chinook Salmon collections were genotyped by CRITFC and were obtained from the FishGen.net online data repository. Contemporary samples for Hanford fall Chinook, and Wenatchee and Methow summer Chinook Salmon were mixed hatchery and natural origin, and data on individual origin were not available. However, broodstocks targeted certain origins and these targets were usually met. It is likely that the summer Chinook Salmon samples were of natural-origin and the fall Chinook Salmon samples were a mix of hatchery and natural-origin fish.

Genetic sample processing
Briefly, at WDFW Molecular Genetics Laboratory, genomic DNA was extracted using silica membrane column extraction kits following manufacturers protocols. We used an amplicon sequencing procedure, Genotyping in Thousands (GTseq, Campbell et al. 2015), to
assay 332 Chinook SNPs (Appendix A). GTseq amplifies pools of targeted SNPs in a highly multiplexed PCR reaction, attaching sequence adapters that assign amplicons to an individual sample and primer. After we sequenced the pooled library, we used a series of custom Perl scripts (c.f., Campbell et al. 2015) to separate the sequences by sample identifiers. A Perl script in the bioinformatics pipeline assigned genotypes based on allele ratios by counting allelespecific amplicons at each locus. The MGL-specific GTseq protocol is described in more detail in Appendix B.

## Data processing

All data processing and analysis were completed using a series of custom R markdown scripts (G.M. - WDFW; R Core Team 2019). All genotype data, baseline and contemporary, were evaluated for missing data and species ID. Species ID was determined using diagnostic markers and homozygosity (non-target species typically have very high homozygosity). Samples with more than $30 \%$ missing genotypes were removed as were samples identified as non-target species.

Only neutral loci were used in further analysis. SNP marker designations, neutral or adaptive, were established by testing in multiple laboratories, including CRITFC and WDFW laboratories, during development of the SNP panel or by designation as adaptive by CRITFC for markers CRITFC ascertained (Jeff Stephenson - CRITFC, pers. comm.). Neutral loci were evaluated for missing data, deviations from Hardy-Weinberg expectations (HWE), and diversity. Loci 1) with more than $30 \%$ missing data across the entire dataset, 2) that were invariant across the entire dataset, or 3) with deviations from HWE in most collections were excluded from further analysis.

## Data analysis

The monitoring and evaluation plan calls for evaluation of four general questions: 1) are contemporary allele frequencies different from baseline allele frequencies (Q7.1.1 and Q7.1.2); 2) is linkage disequilibrium (LD) in contemporary collections different from baseline LD (Q7.2.1 and Q7.2.2); does genetic distance among subpopulations change over time (Q7.3.1); and 4) does the ratio of effective population size $\left(N_{\mathrm{e}}\right)$ to census population size $(\mathrm{N})$ change over time (Q7.4.1)? All analyses were conducted using R markdown scripts using many different R packages (R Core Team 2019). R scripts are available upon request.

Question 1, Allele frequency - To visualize structure among collections associated with allele frequencies, we performed Principal Component Analysis (PCA) on allele frequencies of collections and graphed the first two axes and separately calculated and graphed average allelic richness among all loci within a collection. We statistically evaluated allele frequency similarity by performing pairwise AMOVA analyses, comparing heterozygosity of baseline and contemporary samples, and by evaluating changes in allelic richness. Comparisons of observed and expected heterozygosity were evaluated with a two-sided permutation test where individuals were permuted to obtain the reference distribution.

Question 2, Linkage Disequilibrium - Linkage Disequilibrium (LD) is the correlation of alleles among loci within an individual. Loci may be in LD because they are physically linked (near one another on a chromosome and as such are inherited together) or they may be statistically linked (e.g., alleles are correlated because of relatedness among individuals within a population). No minimum or maximum allowable LD target exists. Because increased LD indicates a reduction in diversity, advice is generally to avoid increasing LD. Hatchery activities may increase the amount of LD present, in particular due to relatedness among individuals. We evaluated LD two ways. First, we calculated allelic correlation coefficients for all pairwise locus comparisons within collection using PLINK (Purcell 2007, Purcell et al. 2007). Second, we performed a probability test of LD for all pairwise locus comparisons within collection using GENEPOP with default parameters (Rousset 2008). Comparisons of baseline and contemporary collections were made by counting the number of statistically significant $(\alpha=0.05)$ pairwise tests before and after correction for multiple tests. At $\alpha=0.05, \sim 5 \%$ of all pairwise tests should have a $P$ value $<0.05$, before correction for multiple tests. Collections with frequencies of $P$ values $<$ $5 \%$ greater than 5\% were inferred to have high levels of LD (Waples 2015). Differences among collections in the frequency of significant pairwise tests of LD within collection were tested using Mann-Whitney rank tests. Correction for multiple testing achieved a table-wide $\alpha=0.05$ for each collection via false discovery rate (Verhoeven et al. 2005).

Question 3, Genetic Distance - To estimate genetic distance among collections we calculated pairwise $F_{\mathrm{ST}}$ and $95 \%$ confidence intervals with the R package hierfstat using default parameters (Goudet 2005). No minimum viable genetic distance has been identified. Instead, the goal is to avoid reducing genetic distances among populations. Increased genetic distance between a hatchery and natural collection of the same population is an indication that the hatchery broodstock were not a representative sample of the population.

Question 4, Effective Population Size - The effective population size $\left(N_{\mathrm{e}}\right)$ of a population is an important metric for populations that roughly indicates the amount of within-population genetic variation that exists because genetic variation generally increases with the effective number of spawners. There is disagreement among experts on minimum viable $N_{\mathrm{e}}$ values, and as such the recommendation is generally to avoid reductions in $N_{\mathrm{e}}$. Effective population size ( $N_{\mathrm{e}}$ ) for each collection separately was estimated using the LDNE algorithms employed by the software NE Estimator (Do et al. 2014). Using this method with the available tissue collections, LDNE is estimating $N_{\mathrm{b}}$, the effective number of breeders, rather than $N_{\mathrm{e}} ; N_{b}$ is a better metric for monitoring (Luikart et al. 2021). Because hatchery programs are integrated programs, hatchery and natural fish belong to the same population. Thus, we also estimated $N_{b}$ with the contemporary hatchery and natural components combined for each of the two years of samples. Loci with very low minor allele frequencies (MAF; in particular, loci where only one copy of the minor allele exists) cause an upward bias in $N_{\mathrm{b}}$ estimates using LDNE (Waples and Do 2008). Inclusion or exclusion of such loci is accomplished by setting a MAF critical value. Because of variable sample sizes and missing data, problem loci have different MAFs. To choose a critical value, for several collections we evaluated the MAF and counted the number of loci that would be dropped at various critical values. Setting the critical value at 0.02 eliminated all or nearly all problem loci, whereas significantly higher numbers of loci that had higher MAFs were dropped when the critical value was set at 0.05 . Thus, we report results based on the critical value of 0.02. We report the jack-knife $95 \%$ confidence interval (CI) for each collection. Statistical
significance of comparisons was evaluated by overlapping CIs. All previous generations impact $N_{\mathrm{b}}$ estimates to some degree and $N_{\mathrm{b}}$ estimates may be biased due to overlapping generations (Waples et al. 2014). To calculate unbiased $N_{\mathrm{b}} / N$ ratios, we estimated the impacts of multiple generations of influence and corrected bias due to overlapping generations (Waples et al. 2014, c.f. Waters et al. 2015) using escapement estimates for as many spawn years prior to the spawn years of our collections as were available in the WDFW SCoRE database
(https://fortress.wa.gov/dfw/score/score/). We assumed a 5-year generation time for natural origin adults and a 4-year generation time for hatchery-produced adults.

## Results

## Sample collections

From 518 baseline samples from 1982-1994, 471 remained after data filtering. Baseline samples were identified as both hatchery- and natural-origin. Genotypes from 17,514 contemporary hatchery- and natural-origin summer and fall Chinook Salmon from 2017 and 2018 were available. To maintain consistent population sizes in baseline and contemporary samples, between 50 and 100 samples were randomly sampled from each population, resulting in 1,106 contemporary samples analyzed. All contemporary samples came from fish that had been used as broodstock for summer and fall Chinook Salmon hatchery programs.

| Analysis Unit | Year | Collection Code | Origin | $\begin{gathered} \text { Run } \\ \text { Timing } \end{gathered}$ | Collection Category | Year | Population | $\underset{\text { available }}{\mathrm{N}}$ | $\underset{\text { used }}{\mathrm{N}}$ | N fixed Loci | AvgRich | Het_obs | Het_exp | Avg $F_{\text {IS }}$ | $\begin{gathered} \% \\ \hline \text { HWE } \\ p< \\ 0.05 \end{gathered}$ | $\begin{gathered} \text { \% Pair } \\ \text { LD } \\ p<0.05 \end{gathered}$ | $\mathrm{N}_{\mathrm{b}}{ }^{1}$ | Jackknife | on samples | $\mathrm{N}^{2}$ | $\mathrm{N}_{b} / \mathrm{N}$ |
| :---: | :---: | :---: | :---: | :---: | :---: | :---: | :---: | :---: | :---: | :---: | :---: | :---: | :---: | :---: | :---: | :---: | :---: | :---: | :---: | :---: | :---: |
| ChelanFalls-Hatchery-Summer-2017 | 2017 | OtsPBT17GHSu | Hatchery | Summer | contemporary | 2017 | Chelan Falls | 333 | 100 | 9 | 1.26 | 0.26 | 0.26 | 0.0064 | 0 | 1514 | 201.7 | 144.4 | 317.3 | NA | NA |
| ChelanFalls-Hatchery-Summer-2018 | 2018 | OtsPBT18FESuFa | Hatchery | Summer | contemporary | 2018 | Chelan Falls | 380 | 100 | 9 | 1.27 | 0.27 | 0.27 | 0.0063 | 0 | 1403 | 389.8 | 228.9 | 1090.9 | NA | nA |
| Entiat_NFH-Hatchery-Summer-2017 | 2017 | OtsPBT17-EntiatNFH | Hatchery | Summer | contemporary | 2017 | Entiat NFH | 273 | 100 | 14 | 1.26 | 0.27 | 0.26 | 0.0223 | 1 | 1516 | 229.9 | 162.2 | 375.6 | NA | NA |
| Entiat_NFH-Hatchery-Summer-2018 | 2018 | OtsPBT18-EntiatNFH | Hatchery | Summer | contemporary | 2018 | Entiat NFH | 269 | 100 | 14 | 1.26 | 0.27 | 0.26 | -0.028 | 0 | 1923 | 93.3 | 69.2 | 134.3 | NA | NA |
| Hanford_Reach-Hatchery-1982 | 1982 | 82 AAB | Hatchery | Fall | baseline | 1982 | Hanford_Reach | 49 | 46 | 21 | 1.26 | 0.29 | 0.26 | -0.111 | 0 | 1016 | 303.4 | 111.7 | Infinite | 30969 | 0.01 |
| Hanford_Reach-Hatchery-1988 | 1988 | 88AAC | Hatchery | Fall | baseline | 1988 | Hanford_Reach | 42 | 39 | 24 | 1.26 | 0.29 | 0.26 | 0.1119 | 0 | 918 | 403.1 | 107.8 | Infinite | 84,299 | 0.00 |
| Hanford_Reach-Natural-1982 | 1982 | 82 AAA | Natural | Fall | baseline | 1982 | Hanford_Reach | 77 | 57 | 20 | 1.26 | 0.27 | 0.26 | 0.0139 | 0 | 1120 | 539.2 | 129.3 | Infinite | 30969 | 0.02 |
| Hanford_Reach-Natural-1988 | 1988 | 88AAB | Natural | Fall | baseline | 1988 | Hanford_Reach | 65 | 53 | 17 | 1.26 | 0.28 | 0.26 | 0.0599 | 0 | 1038 | 1167.1 | 194.5 | Infinite | 84,299 | 0.01 |
| Priest_Rapids_Fish_Hatchery-Hatchery-Fall-2017 | 2017 | OtsPBT17-PRHFa | Unknown | Fall | contemporary | 2017 | Hanford Reach | 6441 | 58 | 14 | 1.27 | 0.27 | 0.27 | 0.0197 | 1 | 1121 | 16246 | 795.3 | Infinite | 174,841 | 0.09 |
| Priest_Rapids_Fish_Hatchery-Hatchery-Fall-2018 | 2018 | OtsPBT18-PRHFa | Unknown | Fall | contemporary | 2018 | Hanford Reach | 6418 | 58 | 11 | 1.27 | 0.27 | 0.27 | 0.0129 | 1 | 1252 | 619.6 | 335.3 | 3380.9 | 183,759 | 0.00 |
| Methow_River-baseline-1994 | 1994 | 94 EJ | Mixed | Summer | baseline | 1994 | Methow_River | 60 | 58 | 19 | 1.26 | 0.25 | 0.25 | 0.0292 | 0 | 1090 | 331.7 | 175.4 | 1935.1 | 1,421 | 0.23 |
| Methow_River-Natural-1993 | 1993 | 93 EC | Natural | Summer | baseline | 1993 | Methow_River | 29 | 29 | 23 | 1.26 | 0.26 | 0.26 | 0.0163 | 0 | 856 | 220.5 | 95.6 | Infinite | 495 | 0.45 |
| Methow-Summer-contemporary-2017 | 2017 | OtsPBT17GMSu | Unknown | Summer | contemporary | 2017 | Methow | 109 | 50 | 8 | 1.26 | 0.26 | 0.26 | 0.0051 | 0 | 1082 | 4197 | 450.2 | Infinite | 3,582 | 1.17 |
| Methow-Summer-contemporary-2018 | 2018 | OtsPBT18FDSuFa | Unknown | Summer | contemporary | 2018 | Methow | 131 | 50 | 16 | 1.26 | 0.26 | 0.26 | 0.0148 | 0 | 1090 | 17663.8 | 767.1 | Infinite | 1,625 | 10.87 |
| Okanogan-natural-baseline-1992 | 1992 | 92FM | Natural | Summer | baseline | 1992 | Similkameen_River | 48 | 46 | 18 | 1.26 | 0.27 | 0.26 | 0.0225 | 0 | 1053 | 451.7 | 129.6 | Infinite | 1,392 | 0.32 |
| Okanogan-natural-baseline-1993 | 1993 | $\begin{gathered} 93 \mathrm{ED} \\ \text { OtsPBT17-CJHInt- } \end{gathered}$ | Natural | Summer | baseline | 1993 | Similkameen_River | 49 | 46 | 16 | 1.27 | 0.27 | 0.27 | 0.0158 | 0 | 1092 | 339.5 | 141.9 | Infinite | 1,719 | 0.20 |
| Okanogan-summer-contemporary-2017 | 2017 | $\begin{gathered} \text { SuFa } \\ \text { OtsPBT18-CJHInt- } \end{gathered}$ | Hatchery | Summer | contemporary | 2017 | Chief Joseph | 682 | 50 | 13 | 1.26 | 0.26 | 0.26 | 0.0268 | 1 | 1052 | 557.2 | 268.5 | Infinite | 5,267 | 0.11 |
| Okanogan-summer-contemporary-2018 | 2018 | SuFa | Hatchery | Summer | contemporary | 2018 | Chief Joseph Wells Fish | 737 | 50 | 14 | 1.27 | 0.27 | 0.27 | 0.013 | 0 | 1142 | 725.1 | 241.7 | Infinite | 10,407 | 0.07 |
| Wells_Fish_Hatchery-Hatchery-Summer-2017 | 2017 | OtsWellsSuPbT17 | Hatchery | Summer | contemporary | 2017 | Hatchery Wells Fish | 534 | 50 | 17 | 1.27 | 0.27 | 0.27 | 0.0087 | 0 | 1137 | 292.5 | 126.1 | Infinite | NA | NA |
| Wells_Fish_Hatchery-Hatchery-Summer-2018 | 2018 | OtsWellsSuPbT18 | Hatchery | Summer | contemporary | 2018 | Hatchery | 752 | 50 | 17 | 1.26 | 0.27 | 0.26 | 0.0021 | 0 | 1192 | 331.8 | 167.6 | 3428.3 | NA | NA |
| Wenatchee_River-Natural-1993 | 1993 | 93 DE | Natural | Summer | baseline | 1993 | Wenatchee_River | 99 | 97 | 6 | 1.26 | 0.26 | 0.26 | 0.0297 | 4 | 1260 | 720 | 279.7 | Infinite | 14,331 | 0.05 |
| Wenatchee-contemporary-Summer-2017 | 2017 | OtsPbT17GLSu | Unknown | Summer | contemporary | 2017 | Wenatchee | 248 | 100 | 8 | 1.26 | 0.26 | 0.26 | 0.0081 | 1 | 1235 | 1531.9 | 726.9 | Infinite | 9,210 | 0.17 |
| Wenatchee-contemporary-Summer-2018 | 2018 | OtsPBT18GLSu | Unknown | Summer | contemporary | 2018 | Wenatchee | 207 | 100 | 11 | 1.26 | 0.26 | 0.26 | 0.0141 | 0 | 1250 | 1918.6 | 802.1 | Infinite | 10,673 | 0.18 |

[^5]
## Evaluation of loci

In total, genotypes from 381 loci were compiled, this includes the 332 SNP loci amplified by the WDFW and an additional 49 SNPs that were included in some of the previously genotyped datasets. Of 381 SNP loci compiled, 82 were identified as adaptive markers and were removed from further analysis as was the sex ID SNP. Of 298 neutral loci, 255 were used in the final analysis. Removed loci included invariant loci ( $n=2$ ), loci with too much missing data $(\mathrm{n}=35)$, and loci with excess deviations from Hardy-Weinberg equilibrium ( $\mathrm{n}=4$ ) and that were excessively negative $F_{\text {IS }}(\mathrm{n}=2)$.

## Data analysis

## Allele frequencies - Question 7.1.1 and 7.1.2 <br> Upper Columbia Summer Chinook Salmon collections

The PCA based on allele frequencies showed three outlier samples (Figure 1). Two of these samples had high heterozygosity relative to other samples in the dataset, suggesting potential contamination. The third outlier sample exhibited no unusual characteristics in terms of missing data or heterozygosity. Minor allele frequencies (MAF) of SNP loci ranged from 0.00 to 0.50 in Upper Columbia summer Chinook Salmon hatchery and natural adult collections. Average MAF of natural baseline and contemporary hatchery collections were similar ( $\sim 0.190$ ). Allelic richness of baseline natural and contemporary hatchery collections was also similar ( $\sim 1.26$ ). AMOVA based on allele frequencies showed a single significant difference between baseline natural-origin collections and contemporary hatchery collections within a population (Methow baseline 1994 vs Methow contemporary 2018). Observed heterozygosity ranged from 0.26 to 0.27 in contemporary HOR adult collections and from 0.25 to 0.27 in NOR baseline collections. No significant difference in average expected heterozygosity was detected among baseline and contemporary collections.

popName

- Chelan Falls Hatchery contemporary 2017
- Chelan Falls Hatchery contemporary 2018
- Chief Joseph Hatchery contemporary 2017
- Chief Joseph Hatchery contemporary 2018
- Entiat NFH Hatchery contemporary 2017
- Entiat NFH Hatchery contemporary 2018
- Methow Hatchery contemporary 2017
- Methow Hatchery contemporary 2018
- Methow River Mixed baseline 1994
- Methow River Natural baseline 1993
- Similkameen River Natural baseline 1992
- Similkameen River Natural baseline 1993
- Wells Fish Hatchery Hatchery contemporary 2017
- Wells Fish Hatchery Hatchery contemporary 2018
- Wenatchee Hatchery contemporary 2017
- Wenatchee Hatchery contemporary 2018
- Wenatchee River Natural baseline 1993

Figure 1. Graph of the first two axes of Principal Components Analysis (PCA) of Upper Columbia summer Chinook Salmon allele frequencies. There is no apparent structure in any of the collections. These graphs identify three outliers on Axis 1 ; two of these outliers have very high heterozygosity suggesting contamination.

## Hanford fall Chinook Salmon collections

The PCA based on allele frequencies showed some structure, along axis two, which appeared to be driven by at least two outliers (Figure 2A). These individuals belonged to the 2018 Contemporary OtsPBT18-PRHFa-0451, and OtsPBT18-PRHFa-2601. In order to determine the origin of these outliers, we completed an assignment test of all individuals against a broad Chinook Salmon baseline. Analysis revealed individual OtsPBT18-PRHFa-0451 to assign with the lower Columbia fall collection ( $99 \%$ assignment probability). The other individual did not assign to a population outside of the Upper Columbia, and thus remained in the analysis. Once the Lower Columbia fall individual was removed, the scatter plot moved to the center of the plot, and no obvious structure was observed (Figure 2B). Minor allele frequencies (MAF) of SNP loci ranged from 0.00 to 0.50 in Hanford fall Chinook Salmon hatchery and natural adult baseline and contemporary collections. Average MAF of baseline and contemporary collections were similar ( $\sim 0.17$ ). Allelic richness of contemporary hatchery collections and baseline hatchery and natural populations were also similar (average $N_{\mathrm{A}} \sim 1.27$ ). AMOVA based on allele frequencies showed the 1982 baseline collections (natural and hatchery) were significantly different from the 2017 contemporary hatchery collections. The other collections did not show significant differences. Observed heterozygosity ranged from $0.10-0.27$ in the baseline natural/hatchery collections $0.13-0.33$ and in the contemporary collections. Significant differences in average expected heterozygosity were detected among baseline (hatchery and natural) and contemporary collections.


Figure 2. Graph of the first two axes of Principal Components Analysis (PCA) of Hanford fall Chinook Salmon allele frequencies (A) with all individuals included, and (B) outlier removed. There is no apparent structure across baseline and contemporary collections.

Linkage Disequilibrium - Question 7.2.1 and 7.2.2
Upper Columbia Summer collections
Weaker linkage disequilibrium existed within hatchery contemporary collections than within baseline natural collections (natural collections average $r^{2}=0.017$, hatchery collections average $r^{2}=0.026$ ). The Wenatchee, Chelan Falls, and Entiat summer Chinook also had lower $r^{2}$ on average than other summer Chinook populations ( 0.012 vs 0.024 ). Surprisingly, there tended to be a negative relationship between $r^{2}$ and the number of significant pairwise tests of LD, with populations with higher $r^{2}$ having fewer locus pairs in significant LD. Mann-Whitney tests of the distribution of $P$ values showed that all pairwise comparisons of those distributions were statistically significant.

## Hanford fall Chinook collections

Stronger linkage disequilibrium existed in baseline hatchery collections (1982 $r^{2}=0.027$; $\left.1988 r^{2}=0.032\right)$ than in the baseline natural collections (1982 $r^{2}=0.021 ; 1988 r^{2}=0.021$ ). Contemporary hatchery collections exhibited similar levels of linkage disequilibrium to the baseline natural collections (2017 $\left.r^{2}=0.18 ; 2018 r^{2}=0.019\right)$. The collections with any significant pairwise tests of LD (one locus pair per each collection, for the same locus pair) were in the contemporary hatchery collections, and in the 1988 natural baseline collection. MannWhitney tests of the distribution of $P$ values showed that all pairwise comparisons of the baseline and contemporary distributions were statistically significant, except for the comparison of the contemporary collections.

## Genetic Distance - Question 7.3.1

Upper Columbia Summer collections
For summer Chinook Salmon, contemporary HOR adults were generally not significantly genetically different from baseline NOR adults as estimated by $F_{\text {ST }}$ (Figure 3, Figure 4); however, there was a general trend where $F_{\mathrm{ST}}$ was elevated in baseline vs contemporary collections. In addition, $F_{\mathrm{ST}}$ among contemporary collections was generally lower than among baseline collections, suggesting homogenization among stocks.


Figure 3. Plot of pairwise $F_{\mathrm{ST}}$ among summer and fall Chinook Salmon populations. $95 \%$ confidence intervals were estimated. Those $F_{\text {ST }}$ estimates whose $95 \%$ confidence intervals lower bound was larger than zero were deemed significantly different from zero and are shown with red outline ( $1=$ significant $), F_{\text {ST }}$ estimates not different than zero have a white outline $(0=$ not significant $)$.


Figure 4. Heatmap of pairwise $F_{\mathrm{ST}}$ among baseline and contemporary summer Chinook Salmon populations. 95\% confidence intervals were estimated. Those $F_{\mathrm{ST}}$ estimates whose $95 \%$ confidence intervals lower bound was larger than zero were deemed significantly different from zero and are shown with red outline ( $1=$ significant $), F_{\text {ST }}$ estimates not different than zero have a white outline $(0=$ not significant $)$.

## Hanford fall Chinook collections

Baseline collections were not significantly different from one another nor were they significantly different from almost all contemporary collections (Figure 5). The lone exception was the comparison of 1982 hatchery baseline to 2017 contemporary, which had a small $F_{\text {ST }}$ ( $F_{\text {ST }}$ $=0.006$ ) but was statistically significant. All pairwise values indicated little differentiation between all collections.


Figure 5. Heatmap of pairwise $F_{\mathrm{ST}}$ among baseline and contemporary Hanford fall Chinook Salmon collections. $95 \%$ confidence intervals were estimated. Those $F_{\text {ST }}$ estimates whose $95 \%$ confidence intervals lower bound was larger than zero were deemed significantly different from zero and are shown with red outline ( $1=$ significant $), F_{\text {ST }}$ estimates not different than zero have a white outline $(0=$ not significant $)$.

Effective Population Size $\left(\mathrm{N}_{\mathrm{e}}\right)$ - Question 7.4.1

## Upper Columbia Summer collections

There was no clear pattern in baseline vs contemporary $N_{\mathrm{b}}$ or $N_{\mathrm{b}} / N$ ratios (Figure 6). Estimates of $N$ were not available for several baseline populations, preventing estimation of $N_{\mathrm{b}} / N$ ratios.


Figure 6. Estimated effective number of breeders ( $N_{\mathrm{b}}$; top) and ratio of $N_{\mathrm{b}}$ to abundance ( $N_{\mathrm{b}} / N$; bottom) for Upper Columbia Summer/Fall Chinook Salmon baseline and contemporary collections. No clear pattern in baseline $N_{\mathrm{b}}$ or $N_{\mathrm{b}} / N$ ratios compared to contemporary was evident (right).

Error bars extending past the graph boundary were infinite indicating not enough linkage disequilibrium existed to estimate the upper bound, i.e., $N_{\mathrm{b}}$ was large. The $N_{\mathrm{b}}$ was estimated using LDNE (Do et al. 2014). Abundance was escapement estimates of hatchery and natural origin fish as found in the WDFW SCoRE database. Since $N_{\mathrm{b}}$ estimates refer to parental generations, abundance from one generation prior was used assuming 5-year generation for natural origin and 4-year generation for hatchery origin.

## Hanford fall Chinook collections

Estimates of $N_{\mathrm{b}}$ for baseline collections were slightly lower than those of contemporary collections, but confidence intervals overlapped greatly. The contemporary 2017 collection had a high estimate in relation to the other collections ( $N_{\mathrm{b}}=16,246$ ), however, the confidence intervals
overlapped those of other collections (Figure 6). Baseline hatchery and natural collections, and the contemporary 2018 collection had low estimates of $N_{\mathrm{b}}$, which were significantly different from the contemporary 2017 collection. There appeared to be no differences in all baseline and contemporary collection $N_{\mathrm{b}} / N$ ratios (Figure 6), except for the 2017 contemporary collection.

## Discussion

To evaluate genetic impacts of hatchery programs on Upper Columbia summer/fall Chinook Salmon populations, we compared genetic data from baseline and contemporary natural and hatchery collections and evaluated genetic metrics. For summer Chinook Salmon, contemporary hatchery collections and baseline natural collections were similar for the genetic metrics measured (allelic richness, allele frequencies, levels of linkage disequilibrium, and $N_{\mathrm{b}}$ ). This suggests that there has not been a significant loss of neutral genetic diversity in contemporary collections relative to baseline collections. For populations with both baseline and contemporary collections, patterns of pairwise $F_{\text {ST }}$ showed slightly higher differentiation among baseline collections than among contemporary collections, and highest $F_{\text {ST }}$ when comparing baseline vs contemporary. It is possible that contemporary populations have become homogenized, and that contemporary populations have differentiated from baseline populations, likely due to genetic drift; however, most of the $F_{\text {ST }}$ comparisons were very low ( $<0.01$ ) and non-significant.

In general Hanford fall Chinook Salmon contemporary and baseline collections displayed similar levels for the genetic metrics measured (allele frequencies, allelic richness, $F_{\mathrm{ST}}$, and $N_{\mathrm{b}}$ ). Baseline hatchery collections (1982 and 1988) showed slightly higher levels of linkage disequilibrium than the other collections. This pattern is likely due to a relatively low number broodstock used in these years, at least in comparison to natural collections, and to the use of a high proportion of hatchery-origin broodstock which tends to increase LD as a result of higher relatedness. The patterns observed in the Hanford fall Chinook Salmon suggest little population differentiation between baseline and contemporary collections, which is most likely due to the large number of broodstock used. The low level of differentiation among contemporary and baseline collections for summer and fall Chinook Salmon is consistent with a previous evaluation that compared samples collected ~2006-2009 with the same baseline individuals, but using different genetic markers (Kassler et al. 2011).

The maintenance of genetic diversity in contemporary summer/fall Chinook Salmon hatchery programs is in contrast to Upper Columbia steelhead which show a reduction in genetic diversity relative to baseline natural populations. This difference is likely due to the larger census size and larger number of broodstock collected for summer and fall Chinook Salmon hatchery programs relative to steelhead. The larger broodstock numbers would help to prevent the loss of alleles due to genetic drift and promote genetic diversity. All of the natural populations of summer Chinook Salmon (Wenatchee, Methow, and Okanogan) and fall Chinook Salmon (Hanford Reach) are large and genetic changes from mechanisms such as genetic drift are less likely in large populations. In addition, large natural populations are less likely to be influenced by hatchery-origin strays because strays would make up a smaller percent of the total spawning population (Pearsons and Miller, see chapter in this report).

Mann-Whitney tests showed significant differences among all pairwise collections in the amount of linkage disequilibrium, which was not informative. This was likely a power issue. With 255 SNP loci we have a lot of power to detect small differences between collections that are not likely to be biologically significant. In the future, different methods of evaluating LD may need to be developed to obtain meaningful results.

Hatcheries can alter among-population genetic structure, and though the monitoring plan did not specifically call for evaluating among-population structure, we were able to evaluate it. In the upper Columbia, hatchery broodstock for the Grand Coulee Fish Maintenance Project beginning in 1939 were once collected in traps at mainstem Columbia River dams, spawned, and were spread throughout all populations we examined, promoting genetic homogenization. Our $F_{\mathrm{ST}}$ results suggest that very little genetic structure currently exists among the Upper Columbia summer or fall Chinook Salmon populations (Figures 3, 4, 5). While the magnitude of $F_{\text {ST }}$ was greater in the baseline collections relative to contemporary samples, the values were still very low and consistent with genetic homogenization prior to the collection of baseline samples. Pearsons and O'Connor (2020) measured donor stray rates (rate of fish originating in a population that stray to a different location) of natural origin Chinook Salmon in the UCR ESU. Donor stray rates among basins was very low; most movement was within basin and from downstream to upstream locations. Straying only impacts among population diversity if strays successfully interbreed with the recipient population (effective strays). For hatchery-origin fish, recipient stray rates among basins were very low for populations of Upper Columbia River summer and fall Chinook Salmon (Pearsons and Miller, see chapter in this report). Since 1999, mean recipient stray rates from non-target hatcheries were $<0.5 \%$ for Wenatchee summer Chinook Salmon, <13\% for Methow summer Chinook Salmon, <2.5\% for the Okanogan, and < $0.71 \%$ for the Hanford Reach. With the exception of the Methow summer Chinook Salmon, all of the recipient populations stray rates were lower than the widely used target of $5 \%$ that is used to reduce the risk of loss of between population genetic diversity (Mobrand et al. 2005, Paquet et al. 2011, Hillman et al. 2020). Recipient stray rates of populations that may have been created by human actions had higher stray rates: $14 \%$ for the Entiat River and $33 \%$ for the Chelan River. Recipient stray rates are currently unknown for natural-origin fish. Effective stray rates are currently unknown but likely lower than estimated recipient stray rates due to local adaptation reducing stray fish reproductive success, which may also drive divergence. Here, however, we used putatively neutral markers, so any divergence observed is likely due to random changes in allele frequencies, i.e., genetic drift.

The monitoring plan and the current implementation of the monitoring plan have limitations, but we are not aware of any other large-scale monitoring of hatchery genetic effects on natural populations that has been developed or implemented. The monitoring plan has been extensively reviewed by science and genetic experts (e.g., ISAB and genetic expert panel) and has been adapted based on evaluation of reviews. One of the challenges associated with longterm genetic monitoring is changes in genetic techniques. Over the years, upper Columbia hatchery evaluations have utilized allozyme, microsatellite, and SNP markers making direct comparisons of results problematic. Adding larger sample sizes to the M\&E program may be appropriate when final analyses or specific issues need resolution, but interim evaluations may not need such level of precision, particularly if new and more powerful techniques are available
for future monitoring work. The monitoring plan also lacks monitoring of adaptive genetic diversity. The SNP markers associated with adaptive traits have been discovered and developed for run-timing (Prince et al. 2017, Narum et al. 2018, Thompson et al. 2020) and male age at maturity (McKinney et al. 2020, McKinney et al. 2021) and could be used to evaluate the impacts of hatchery propagation on allele frequencies at these markers. Surprisingly, instead of having polygenic associations with important traits, some of these adaptive traits are associated with single gene regions with only a few variant alleles (Ford et al. 2020). Under this simpler system, variability can be rapidly lost from a population. Monitoring of allele frequencies of these few available marker-trait associations may be important for those traits but also would serve as model data for other undiscovered marker-trait associations that may have a similar genetic architecture. The full PUD monitoring and evaluation plan includes many additional metrics that help evaluate adaptive traits (see other chapters in this report), including straying, productivity, age at maturity, size at age, run and spawn time, spawn distribution, and PNI. Evaluating genetic and other metrics together would provide the most comprehensive means of evaluating the hatchery programs, but it is unclear how additional metrics would influence adaptive management decisions.

This monitoring evaluation was the second timestep of the monitoring plan following initial baseline sampling (e.g., a total of 3 collection times representing multiple broodyears) and represented a large effort in genotyping and analysis to characterize patterns of genetic variation in hatchery and natural origin samples from different time points. However, this evaluation should be considered within the larger context of the long-term monitoring and evaluation plan. Future assessments will occur at 10 year intervals and will result in an increase in sample size and the broodyears included (e.g., evaluations in 2031, 2041, 2051). Definitive conclusions may not be possible during each 10 year timestep, but cumulative assessments should provide useful information to adaptively manage the hatchery programs. As mentioned above, one of the biggest challenges in this timestep and future timesteps is how to compare data and findings from evaluations that use different genetic and analytical techniques. We were able to compare only a subset of data and findings from the last genetic evaluation and we also reran the baseline samples using the most current techniques so that we could make direct comparisons to baseline samples. Future work would benefit by developing technical approaches that would facilitate more direct comparisons across 10 year timesteps. Another potential improvement would be to evaluate the sample sizes, cohorts, and number of timesteps necessary to detect specified genetic changes of interest to managers. The sample sizes and cohorts used in this evaluation were chosen by attempting to balance standard genetic sample sizes, inclusion of the most recent cohorts to provide maximum contrast, and cost. Power analyses could be conducted to refine sample sizes and cohorts necessary to detect changes at a reasonable cost. The power to detect declines in $N_{\mathrm{b}}$ is in part determined by the number of cohorts analyzed (Luikart et al. 2021). Improvements in the power to detect changes in $N_{\mathrm{b}}$ could be achieved by including future 10year timesteps. Luikart et al. (2021) found that the power to detect a declining trend in $N_{\mathrm{b}}$ nearly doubled when the number of consecutive cohorts analyzed went from five to ten. Up to a point, power also increased when the number of samples and loci used increased. However, changing loci in different timesteps will make it more difficult to compare data across the duration of the monitoring program. While $N_{\mathrm{b}}$ estimates were high in this study, early detection of any future negative trends in $N_{\mathrm{b}}$ would allow for changes to be made to hatchery programs preventing further decline. This might be particularly important if census sizes, which are
monitored annually, decrease substantially. Finally, some of the contemporary collections used in this evaluation (Hanford Reach, Wenatchee, Methow) likely contained both natural- and hatchery-origin individuals; however, data on individual origin was not available, limiting our ability to evaluate whether contemporary natural- and hatchery-origin individuals show different genetic trends, as was clearly seen in upper Columbia steelhead. Obtaining data on individual origin should be considered for the next round of monitoring.

Summary - In agreement with previous analyses, the baseline and contemporary Upper Columbia River summer and fall Chinook Salmon collections showed similar levels of neutral genetic diversity suggesting that hatcheries have not led to a reduction in this genetic diversity within this time frame. These findings are consistent with observed recipient population stray rates and with current management strategies that are intended to minimize genetic risks. However, it is possible that differences in some variables were not detected because of low statistical power. Other assessments of phenotypic variables such as run and spawn timing and age-at-maturity were also evaluated in other chapters of this report. In addition, contemporary collections were composed of either hatchery-origin or a mix of hatchery- and natural-origin individuals, and in the latter case data on origin was not available for individual fish, preventing comparison of genetic patterns in natural- vs hatchery-origin fish in contemporary samples. The lack of genetic differentiation among contemporary and baseline samples is likely explained by a combination of the large population sizes of summer and fall Chinook Salmon relative to the hatchery program sizes in the upper Columbia basin, low recipient population stray rates in natural populations, and the management strategies that were implemented to reduce genetic risk.

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Appendix A. List of adaptive and neutral diploid single nucleotide polymorphic (SNP) loci used in this study. Primer and probe sequences for unpublished loci available by request.

| Locus Name | Purpose | Allele 1 | Allele 2 | Reference |
| :---: | :---: | :---: | :---: | :---: |
| Ots_101770-82 | Neutral | G | T | (Janowitz-Koch et al. 2019) |
| Ots_104048-194 | Neutral | C | T | (Janowitz-Koch et al. 2019) |
| Ots_105897-124 | Neutral | C | T | (Janowitz-Koch et al. 2019) |
| Ots_111312-435 | Neutral | C | T | (Janowitz-Koch et al. 2019) |
| Ots_98409-850 | Neutral | C | T | (Janowitz-Koch et al. 2019) |
| Ots_98683-796 | Neutral | A | T | (Janowitz-Koch et al. 2019) |
| Ots_aldb-177M | Neutral | T | A | (Janowitz-Koch et al. 2019) |
| Ots_Chin30up-211 | Neutral | C | T | (Janowitz-Koch et al. 2019) |
| Ots_crRAD12711-37 | Neutral | C | T | (Janowitz-Koch et al. 2019) |
| Ots_crRAD26541-47 | Neutral | G | A | (Janowitz-Koch et al. 2019) |
| Ots_crRAD28677-65 | Neutral | C | T | (Janowitz-Koch et al. 2019) |
| Ots_crRAD292-21 | Neutral | C | T | (Janowitz-Koch et al. 2019) |
| Ots_crRAD30341-48 | Neutral | T | A | (Janowitz-Koch et al. 2019) |
| Ots_crRAD33054-62 | Neutral | A | T | (Janowitz-Koch et al. 2019) |
| Ots_crRAD3758-51 | Neutral | T | C | (Janowitz-Koch et al. 2019) |
| Ots_crRAD38095-29 | Neutral | A | T | (Janowitz-Koch et al. 2019) |
| Ots_crRAD38746-36 | Neutral | T | A | (Janowitz-Koch et al. 2019) |
| Ots_crRAD42058-48 | Neutral | A | T | (Janowitz-Koch et al. 2019) |
| Ots_crRAD48459-74 | Neutral | T | C | (Janowitz-Koch et al. 2019) |
| Ots_crRAD5061-27 | Neutral | T | C | (Janowitz-Koch et al. 2019) |
| Ots_crRAD57537-24 | Neutral | A | C | (Janowitz-Koch et al. 2019) |
| Ots_FGF6A | Neutral | G | T | (Janowitz-Koch et al. 2019) |
| Ots_hsc71-5'-453 | Neutral | C | T | (Janowitz-Koch et al. 2019) |
| Ots_LEI-292 | Neutral | G | A | (Janowitz-Koch et al. 2019) |
| Ots_MHC1 | Neutral | G | A | (Janowitz-Koch et al. 2019) |
| Ots_SERPC1-209 | Neutral | A | T | (Janowitz-Koch et al. 2019) |
| Ots_Tnsf | Neutral | A | G | (Janowitz-Koch et al. 2019) |
| Ots_u07-17.373 | Neutral | A | Deletion | (Janowitz-Koch et al. 2019) |
| Ots_u07-19.260 | Neutral | C | T | (Janowitz-Koch et al. 2019) |
| Ots_u1004-117 | Neutral | C | T | (Janowitz-Koch et al. 2019) |
| Ots_u1006-171 | Neutral | C | T | (Janowitz-Koch et al. 2019) |
| Ots_USMG5-67 | Neutral | C | T | (Janowitz-Koch et al. 2019) |
| Ots_zP3b-215 | Neutral | G | T | (Janowitz-Koch et al. 2019) |
| Ots_110495-380 | Neutral | G | C | (Janowitz-Koch et al. 2019) |
| Ots_ARNT | Neutral | G | T | (Janowitz-Koch et al. 2019) |
| Ots_crRAD18289-33 | Neutral | T | C | (Janowitz-Koch et al. 2019) |
| Ots_crRAD55400-59 | Neutral | C | T | (Janowitz-Koch et al. 2019) |
| Ots_crRAD57376-68 | Neutral | T | C | (Janowitz-Koch et al. 2019) |
| Ots_100884-287 | Neutral | T | C | (Janowitz-Koch et al. 2019) |
| Ots_101119-381 | Neutral | T | C | (Janowitz-Koch et al. 2019) |


| Locus Name | Purpose | Allele 1 | Allele 2 | Reference |
| :---: | :---: | :---: | :---: | :---: |
| Ots_101554-407 | Neutral | C | G | (Janowitz-Koch et al. 2019) |
| Ots_101704-143 | Neutral | T | G | (Janowitz-Koch et al. 2019) |
| Ots_102213-210 | Neutral | A | G | (Janowitz-Koch et al. 2019) |
| Ots_102414-395 | Neutral | A | G | (Janowitz-Koch et al. 2019) |
| Ots_102457-132 | Neutral | A | G | (Janowitz-Koch et al. 2019) |
| Ots_102801-308 | Neutral | C | A | (Janowitz-Koch et al. 2019) |
| Ots_102867-609 | Neutral | A | G | (Janowitz-Koch et al. 2019) |
| Ots_103041-52 | Neutral | G | A | (Janowitz-Koch et al. 2019) |
| Ots_103122-180 | Neutral | T | C | (Janowitz-Koch et al. 2019) |
| Ots_104063-132 | Neutral | C | T | (Janowitz-Koch et al. 2019) |
| Ots_104415-88 | Neutral | C | T | (Janowitz-Koch et al. 2019) |
| Ots_105105-613 | Neutral | C | G | (Janowitz-Koch et al. 2019) |
| Ots_105132-200 | Neutral | G | T | (Janowitz-Koch et al. 2019) |
| Ots_105385-421 | Neutral | A | G | (Janowitz-Koch et al. 2019) |
| Ots_105401-325 | Neutral | G | T | (Janowitz-Koch et al. 2019) |
| Ots_105407-117 | Neutral | T | A | (Janowitz-Koch et al. 2019) |
| Ots_106313-729 | Neutral | A | G | (Janowitz-Koch et al. 2019) |
| Ots_106419b-618 | Neutral | G | T | (Janowitz-Koch et al. 2019) |
| Ots_106499-70 | Neutral | C | G | (Janowitz-Koch et al. 2019) |
| Ots_106747-239 | Neutral | C | A | (Janowitz-Koch et al. 2019) |
| Ots_107074-284 | Neutral | A | T | (Janowitz-Koch et al. 2019) |
| Ots_107285-93 | Neutral | T | A | (Janowitz-Koch et al. 2019) |
| Ots_107607-315 | Neutral | A | C | (Janowitz-Koch et al. 2019) |
| Ots_107806-821 | Neutral | T | A | (Janowitz-Koch et al. 2019) |
| Ots_108007-208 | Neutral | A | T | (Janowitz-Koch et al. 2019) |
| Ots_108390-329 | Neutral | G | C | (Janowitz-Koch et al. 2019) |
| Ots_108735-302 | Neutral | C | T | (Janowitz-Koch et al. 2019) |
| Ots_108820-336 | Neutral | G | A | (Janowitz-Koch et al. 2019) |
| Ots_109525-816 | Neutral | C | T | (Janowitz-Koch et al. 2019) |
| Ots_109693-392 | Neutral | T | G | (Janowitz-Koch et al. 2019) |
| Ots_110064-383 | Neutral | C | T | (Janowitz-Koch et al. 2019) |
| Ots_110201-363 | Neutral | A | T | (Janowitz-Koch et al. 2019) |
| Ots_110381-164 | Neutral | A | G | (Janowitz-Koch et al. 2019) |
| Ots_110551-64 | Neutral | C | A | (Janowitz-Koch et al. 2019) |
| Ots_110689-218 | Neutral | T | G | (Janowitz-Koch et al. 2019) |
| Ots_111084b-619 | Neutral | C | A | (Janowitz-Koch et al. 2019) |
| Ots_111681-657 | Neutral | G | T | (Janowitz-Koch et al. 2019) |
| Ots_112208-722 | Neutral | C | A | (Janowitz-Koch et al. 2019) |
| Ots_112301-43 | Neutral | T | C | (Janowitz-Koch et al. 2019) |
| Ots_112419-131 | Neutral | A | T | (Janowitz-Koch et al. 2019) |
| Ots_112820-284 | Neutral | C | T | (Janowitz-Koch et al. 2019) |
| Ots_112876-371 | Neutral | C | A | (Janowitz-Koch et al. 2019) |


| Locus Name | Purpose | Allele 1 | Allele 2 | Reference |
| :---: | :---: | :---: | :---: | :---: |
| Ots_113242-216 | Neutral | C | T | (Janowitz-Koch et al. 2019) |
| Ots_113457-40R | Neutral | C | T | (Janowitz-Koch et al. 2019) |
| Ots_115987-325 | Neutral | T | G | (Janowitz-Koch et al. 2019) |
| Ots_117242-136 | Neutral | A | G | (Janowitz-Koch et al. 2019) |
| Ots_117259-271 | Neutral | T | G | (Janowitz-Koch et al. 2019) |
| Ots_117370-471 | Neutral | G | T | (Janowitz-Koch et al. 2019) |
| Ots_117432-409 | Neutral | A | G | (Janowitz-Koch et al. 2019) |
| Ots_118175-479 | Neutral | C | T | (Janowitz-Koch et al. 2019) |
| Ots_118205-61 | Neutral | T | C | (Janowitz-Koch et al. 2019) |
| Ots_118938-325 | Neutral | C | T | (Janowitz-Koch et al. 2019) |
| Ots_120950-417 | Neutral | A | T | (Janowitz-Koch et al. 2019) |
| Ots_122414-56 | Neutral | C | T | (Janowitz-Koch et al. 2019) |
| Ots_123048-521 | Neutral | A | C | (Janowitz-Koch et al. 2019) |
| Ots_123921-111 | Neutral | A | G | (Janowitz-Koch et al. 2019) |
| Ots_124774-477 | Neutral | T | C | (Janowitz-Koch et al. 2019) |
| Ots_126619-400 | Neutral | C | T | (Janowitz-Koch et al. 2019) |
| Ots_127236-62 | Neutral | T | A | (Janowitz-Koch et al. 2019) |
| Ots_127760-569 | Neutral | C | T | (Janowitz-Koch et al. 2019) |
| Ots_128302-57 | Neutral | C | T | (Janowitz-Koch et al. 2019) |
| Ots_128693-461 | Neutral | C | T | (Janowitz-Koch et al. 2019) |
| Ots_128757-61R | Neutral | A | Deletion | (Janowitz-Koch et al. 2019) |
| Ots_129144-472 | Neutral | C | A | (Janowitz-Koch et al. 2019) |
| Ots_129170-683 | Neutral | C | A | (Janowitz-Koch et al. 2019) |
| Ots_129458-451 | Neutral | T | C | (Janowitz-Koch et al. 2019) |
| Ots_129870-55 | Neutral | A | T | (Janowitz-Koch et al. 2019) |
| Ots_130720-99 | Neutral | A | G | (Janowitz-Koch et al. 2019) |
| Ots_131460-584 | Neutral | T | C | (Janowitz-Koch et al. 2019) |
| Ots_131802-393 | Neutral | C | T | (Janowitz-Koch et al. 2019) |
| Ots_131906-141 | Neutral | A | T | (Janowitz-Koch et al. 2019) |
| Ots_94857-232R | Neutral | T | C | (Janowitz-Koch et al. 2019) |
| Ots_94903-99R | Neutral | G | T | (Janowitz-Koch et al. 2019) |
| Ots_95442b-204 | Neutral | A | T | (Janowitz-Koch et al. 2019) |
| Ots_96222-525 | Neutral | C | T | (Janowitz-Koch et al. 2019) |
| Ots_96500-180 | Neutral | G | T | (Janowitz-Koch et al. 2019) |
| Ots_96899-357R | Neutral | T | A | (Janowitz-Koch et al. 2019) |
| Ots_97077-179R | Neutral | G | T | (Janowitz-Koch et al. 2019) |
| Ots_97660-56 | Neutral | A | T | (Janowitz-Koch et al. 2019) |
| Ots_99550-204 | Neutral | C | T | (Janowitz-Koch et al. 2019) |
| Ots_afmid-196 | Neutral | G | C | (Janowitz-Koch et al. 2019) |
| Ots_AldB1-122 | Neutral | C | T | (Janowitz-Koch et al. 2019) |
| Ots_AldoB4-183 | Neutral | T | A | (Janowitz-Koch et al. 2019) |
| Ots_arp-436 | Neutral | A | T | (Janowitz-Koch et al. 2019) |


| Locus Name | Purpose | Allele 1 | Allele 2 | Reference |
| :---: | :---: | :---: | :---: | :---: |
| Ots_AsnRS-60 | Neutral | T | C | (Janowitz-Koch et al. 2019) |
| Ots_aspat-196 | Neutral | G | C | (Janowitz-Koch et al. 2019) |
| Ots_BMP2-SNP1 | Neutral | C | T | (Janowitz-Koch et al. 2019) |
| Ots_brp16-64 | Neutral | T | C | (Janowitz-Koch et al. 2019) |
| Ots_Cath_D141 | Neutral | T | C | (Janowitz-Koch et al. 2019) |
| Ots_CCR7 | Neutral | C | T | (Janowitz-Koch et al. 2019) |
| Ots_CD59-2 | Neutral | G | A | (Janowitz-Koch et al. 2019) |
| Ots_CD63 | Neutral | A | C | (Janowitz-Koch et al. 2019) |
| Ots_cgo24-22 | Neutral | T | C | (Janowitz-Koch et al. 2019) |
| Ots_CirpA | Neutral | C | T | (Janowitz-Koch et al. 2019) |
| Ots_cox 1-241 | Neutral | C | T | (Janowitz-Koch et al. 2019) |
| Ots_CRB211 | Neutral | A | C | (Janowitz-Koch et al. 2019) |
| Ots_crRAD10447-25 | Neutral | C | T | (Janowitz-Koch et al. 2019) |
| Ots_crRAD11620-55 | Neutral | C | T | (Janowitz-Koch et al. 2019) |
| Ots_crRAD12037-39 | Neutral | A | G | (Janowitz-Koch et al. 2019) |
| Ots_crRAD13725-51 | Neutral | C | A | (Janowitz-Koch et al. 2019) |
| Ots_crRAD16540-50 | Neutral | C | T | (Janowitz-Koch et al. 2019) |
| Ots_crRAD17527-58 | Neutral | C | T | (Janowitz-Koch et al. 2019) |
| Ots_crRAD18492-65 | Neutral | C | T | (Janowitz-Koch et al. 2019) |
| Ots_crRAD18937-60 | Neutral | G | A | (Janowitz-Koch et al. 2019) |
| Ots_crRAD20262-46 | Neutral | A | G | (Janowitz-Koch et al. 2019) |
| Ots_crRAD20376-66 | Neutral | G | A | (Janowitz-Koch et al. 2019) |
| Ots_crRAD20887-70 | Neutral | G | A | (Janowitz-Koch et al. 2019) |
| Ots_crRAD21115-24 | Neutral | C | T | (Janowitz-Koch et al. 2019) |
| Ots_crRAD22960-32 | Neutral | C | T | (Janowitz-Koch et al. 2019) |
| Ots_crRAD23631-48 | Neutral | G | A | (Janowitz-Koch et al. 2019) |
| Ots_crRAD24807-74 | Neutral | A | T | (Janowitz-Koch et al. 2019) |
| Ots_crRAD25367-50 | Neutral | T | G | (Janowitz-Koch et al. 2019) |
| Ots_crRAD255-59 | Neutral | T | C | (Janowitz-Koch et al. 2019) |
| Ots_crRAD26081-28 | Neutral | T | G | (Janowitz-Koch et al. 2019) |
| Ots_crRAD26165-69 | Neutral | C | T | (Janowitz-Koch et al. 2019) |
| Ots_crRAD27164-55 | Neutral | A | T | (Janowitz-Koch et al. 2019) |
| Ots_crRAD27515-69 | Neutral | T | A | (Janowitz-Koch et al. 2019) |
| Ots_crRAD2806-42 | Neutral | C | A | (Janowitz-Koch et al. 2019) |
| Ots_crRAD33491-71 | Neutral | C | T | (Janowitz-Koch et al. 2019) |
| Ots_crRAD34397-33 | Neutral | C | G | (Janowitz-Koch et al. 2019) |
| Ots_crRAD35313-66 | Neutral | A | G | (Janowitz-Koch et al. 2019) |
| Ots_crRAD36072-29 | Neutral | T | C | (Janowitz-Koch et al. 2019) |
| Ots_crRAD36152-44 | Neutral | C | T | (Janowitz-Koch et al. 2019) |
| Ots_crRAD44588-67 | Neutral | C | T | (Janowitz-Koch et al. 2019) |
| Ots_crRAD46081-56 | Neutral | C | T | (Janowitz-Koch et al. 2019) |
| Ots_crRAD46751-42 | Neutral | C | T | (Janowitz-Koch et al. 2019) |


| Locus Name | Purpose | Allele 1 | Allele 2 | Reference |
| :---: | :---: | :---: | :---: | :---: |
| Ots_crRAD47297-55 | Neutral | T | C | (Janowitz-Koch et al. 2019) |
| Ots_crRAD55475-26 | Neutral | T | G | (Janowitz-Koch et al. 2019) |
| Ots_crRAD57520-66 | Neutral | T | G | (Janowitz-Koch et al. 2019) |
| Ots_crRAD57687-34 | Neutral | T | G | (Janowitz-Koch et al. 2019) |
| Ots_crRAD60614-46 | Neutral | G | T | (Janowitz-Koch et al. 2019) |
| Ots_crRAD60620-51 | Neutral | A | G | (Janowitz-Koch et al. 2019) |
| Ots_crRAD61523-71 | Neutral | A | G | (Janowitz-Koch et al. 2019) |
| Ots_crRAD66330-60 | Neutral | G | T | (Janowitz-Koch et al. 2019) |
| Ots_crRAD69327-53 | Neutral | G | T | (Janowitz-Koch et al. 2019) |
| Ots_crRAD73823-60 | Neutral | T | A | (Janowitz-Koch et al. 2019) |
| Ots_crRAD74766-28 | Neutral | G | A | (Janowitz-Koch et al. 2019) |
| Ots_crRAD75581-70 | Neutral | A | G | (Janowitz-Koch et al. 2019) |
| Ots_crRAD76512-28 | Neutral | A | T | (Janowitz-Koch et al. 2019) |
| Ots_crRAD78968-46 | Neutral | C | T | (Janowitz-Koch et al. 2019) |
| Ots_crRAD92420-25 | Neutral | G | T | (Janowitz-Koch et al. 2019) |
| Ots_crRAD9615-69 | Neutral | T | C | (Janowitz-Koch et al. 2019) |
| Ots_DDX5-171 | Neutral | C | T | (Janowitz-Koch et al. 2019) |
| Ots_E2-275 | Neutral | A | G | (Janowitz-Koch et al. 2019) |
| Ots_EndoRB1-486 | Neutral | G | A | (Janowitz-Koch et al. 2019) |
| Ots_EP-529 | Neutral | A | G | (Janowitz-Koch et al. 2019) |
| Ots_Est1363 | Neutral | A | T | (Janowitz-Koch et al. 2019) |
| Ots_Est740 | Neutral | T | C | (Janowitz-Koch et al. 2019) |
| Ots_ETIF1A | Neutral | A | C | (Janowitz-Koch et al. 2019) |
| Ots_FARSLA-220 | Neutral | G | A | (Janowitz-Koch et al. 2019) |
| Ots_FGF6B_1 | Neutral | A | C | (Janowitz-Koch et al. 2019) |
| Ots_GCSH | Neutral | C | T | (Janowitz-Koch et al. 2019) |
| Ots_GDH-81x | Neutral | C | Deletion | (Janowitz-Koch et al. 2019) |
| Ots_GH2 | Neutral | A | T | (Janowitz-Koch et al. 2019) |
| Ots_GnRH-271 | Neutral | C | T | (Janowitz-Koch et al. 2019) |
| Ots_GPDH-338 | Neutral | G | A | (Janowitz-Koch et al. 2019) |
| Ots_GPH-318 | Neutral | C | T | (Janowitz-Koch et al. 2019) |
| Ots_GST-207 | Neutral | G | A | (Janowitz-Koch et al. 2019) |
| Ots_GST-375 | Neutral | C | T | (Janowitz-Koch et al. 2019) |
| Ots_GTH2B-550 | Neutral | C | G | (Janowitz-Koch et al. 2019) |
| Ots_HFABP-34 | Neutral | C | T | (Janowitz-Koch et al. 2019) |
| Ots_HMGB1-73 | Neutral | G | T | (Janowitz-Koch et al. 2019) |
| Ots_hnRNPL-533 | Neutral | A | T | (Janowitz-Koch et al. 2019) |
| Ots_hsc71-3'-488 | Neutral | C | T | (Janowitz-Koch et al. 2019) |
| Ots_hsp27b-150 | Neutral | G | A | (Janowitz-Koch et al. 2019) |
| Ots_Hsp90a | Neutral | G | C | (Janowitz-Koch et al. 2019) |
| Ots_HSP90B-100 | Neutral | C | T | (Janowitz-Koch et al. 2019) |
| Ots_IGF-I.1-76 | Neutral | A | T | (Janowitz-Koch et al. 2019) |


| Locus Name | Purpose | Allele 1 | Allele 2 | Reference |
| :---: | :---: | :---: | :---: | :---: |
| Ots_Ikaros-250 | Neutral | G | A | (Janowitz-Koch et al. 2019) |
| Ots_IL11 | Neutral | T | C | (Janowitz-Koch et al. 2019) |
| Ots_IL8R_C8 | Neutral | C | T | (Janowitz-Koch et al. 2019) |
| Ots_IsoT | Neutral | T | C | (Janowitz-Koch et al. 2019) |
| Ots_LWSop-638 | Neutral | T | C | (Janowitz-Koch et al. 2019) |
| Ots_mapK-3'-309 | Neutral | T | G | (Janowitz-Koch et al. 2019) |
| Ots_mapKpr-151 | Neutral | A | T | (Janowitz-Koch et al. 2019) |
| Ots_MetA | Neutral | T | A | (Janowitz-Koch et al. 2019) |
| Ots_MHC2 | Neutral | T | G | (Janowitz-Koch et al. 2019) |
| Ots_mybp-85 | Neutral | C | T | (Janowitz-Koch et al. 2019) |
| Ots_Myc-366 | Neutral | T | C | (Janowitz-Koch et al. 2019) |
| Ots_myo1a-384 | Neutral | A | C | (Janowitz-Koch et al. 2019) |
| Ots_myoD-364 | Neutral | T | G | (Janowitz-Koch et al. 2019) |
| Ots_NAML12-SNP1 | Neutral | A | G | (Janowitz-Koch et al. 2019) |
| Ots_nelfd-163 | Neutral | A | G | (Janowitz-Koch et al. 2019) |
| Ots_NFYB-147 | Neutral | C | T | (Janowitz-Koch et al. 2019) |
| Ots_nkef-192 | Neutral | C | T | (Janowitz-Koch et al. 2019) |
| Ots_NOD1 | Neutral | C | G | (Janowitz-Koch et al. 2019) |
| Ots_nramp-321 | Neutral | G | A | (Janowitz-Koch et al. 2019) |
| Ots_ntl-255 | Neutral | T | A | (Janowitz-Koch et al. 2019) |
| Ots_Ostm1 | Neutral | C | G | (Janowitz-Koch et al. 2019) |
| Ots_OTALDBINT1SNP1 | Neutral | T | C | (Janowitz-Koch et al. 2019) |
| Ots_OTDESMIN19SNP1 | Neutral | C | A | (Janowitz-Koch et al. 2019) |
| Ots_Ots 311-101x | Neutral | A | Deletion | (Janowitz-Koch et al. 2019) |
| Ots_OTSMTA-SNP1 | Neutral | C | T | (Janowitz-Koch et al. 2019) |
| Ots_OTSTF1-SNP1 | Neutral | G | T | (Janowitz-Koch et al. 2019) |
| Ots_P450-288 | Neutral | A | G | (Janowitz-Koch et al. 2019) |
| Ots_P450 | Neutral | T | A | (Janowitz-Koch et al. 2019) |
| Ots_P53 | Neutral | G | A | (Janowitz-Koch et al. 2019) |
| Ots_parp3-286 | Neutral | A | G | (Janowitz-Koch et al. 2019) |
| Ots_PEMT | Neutral | C | T | (Janowitz-Koch et al. 2019) |
| Ots_PGK-54 | Neutral | T | A | (Janowitz-Koch et al. 2019) |
| Ots_pigh-105 | Neutral | A | Deletion | (Janowitz-Koch et al. 2019) |
| Ots_pop5-96 | Neutral | T | C | (Janowitz-Koch et al. 2019) |
| Ots_ppie-245 | Neutral | C | A | (Janowitz-Koch et al. 2019) |
| Ots_Prl2 | Neutral | A | G | (Janowitz-Koch et al. 2019) |
| Ots_RAD4543-52 | Neutral | T | C | (Janowitz-Koch et al. 2019) |
| Ots_RAG3 | Neutral | C | T | (Janowitz-Koch et al. 2019) |
| Ots_RAS1 | Neutral | C | T | (Janowitz-Koch et al. 2019) |
| Ots_redd1-187 | Neutral | A | G | (Janowitz-Koch et al. 2019) |


| Locus Name | Purpose | Allele 1 | Allele 2 | Reference |
| :---: | :---: | :---: | :---: | :---: |
| Ots_RFC2-558 | Neutral | A | Deletion | (Janowitz-Koch et al. 2019) |
| Ots_S7-1 | Neutral | T | C | (Janowitz-Koch et al. 2019) |
| Ots_SClkF2R2-135 | Neutral | A | T | (Janowitz-Koch et al. 2019) |
| Ots_sept9-78 | Neutral | G | A | (Janowitz-Koch et al. 2019) |
| Ots_SL | Neutral | A | G | (Janowitz-Koch et al. 2019) |
| Ots_slc7a2-71 | Neutral | G | T | (Janowitz-Koch et al. 2019) |
| Ots_stk6-516 | Neutral | C | A | (Janowitz-Koch et al. 2019) |
| Ots_SWS1op-182 | Neutral | T | A | (Janowitz-Koch et al. 2019) |
| Ots_TAPBP | Neutral | C | T | (Janowitz-Koch et al. 2019) |
| Ots_TCTA-58 | Neutral | C | T | (Janowitz-Koch et al. 2019) |
| Ots_TGFB | Neutral | C | T | (Janowitz-Koch et al. 2019) |
| Ots_Thio | Neutral | T | C | (Janowitz-Koch et al. 2019) |
| Ots_TLR3 | Neutral | C | T | (Janowitz-Koch et al. 2019) |
| Ots_TNF | Neutral | C | T | (Janowitz-Koch et al. 2019) |
| Ots_tpx2-125 | Neutral | C | T | (Janowitz-Koch et al. 2019) |
| Ots_trnau1ap-86 | Neutral | G | T | (Janowitz-Koch et al. 2019) |
| Ots_txnip-321 | Neutral | T | C | (Janowitz-Koch et al. 2019) |
| Ots_u07-07.161 | Neutral | C | T | (Janowitz-Koch et al. 2019) |
| Ots_u07-17.135 | Neutral | A | G | (Janowitz-Koch et al. 2019) |
| Ots_u07-18.378 | Neutral | A | T | (Janowitz-Koch et al. 2019) |
| Ots_u07-20.332 | Neutral | A | C | (Janowitz-Koch et al. 2019) |
| Ots_u07-25.325 | Neutral | T | C | (Janowitz-Koch et al. 2019) |
| Ots_u07-49.290 | Neutral | G | A | (Janowitz-Koch et al. 2019) |
| Ots_u07-53.133 | Neutral | C | T | (Janowitz-Koch et al. 2019) |
| Ots_u07-57.120 | Neutral | A | T | (Janowitz-Koch et al. 2019) |
| Ots_u07-64.221 | Neutral | G | C | (Janowitz-Koch et al. 2019) |
| Ots_u1002-75 | Neutral | T | C | (Janowitz-Koch et al. 2019) |
| Ots_u1007-124 | Neutral | A | G | (Janowitz-Koch et al. 2019) |
| Ots_u1008-108 | Neutral | T | A | (Janowitz-Koch et al. 2019) |
| Ots_u202-161 | Neutral | T | A | (Janowitz-Koch et al. 2019) |
| Ots_u211-85 | Neutral | C | T | (Janowitz-Koch et al. 2019) |
| Ots_U212-158 | Neutral | G | A | (Janowitz-Koch et al. 2019) |
| Ots_U2305-63 | Neutral | T | Deletion | (Janowitz-Koch et al. 2019) |
| Ots_U2362-227 | Neutral | A | T | (Janowitz-Koch et al. 2019) |
| Ots_U2362-330 | Neutral | A | G | (Janowitz-Koch et al. 2019) |
| Ots_U2446-123 | Neutral | C | A | (Janowitz-Koch et al. 2019) |
| Ots_U2567-104 | Neutral | G | A | (Janowitz-Koch et al. 2019) |
| Ots_u4-92 | Neutral | T | C | (Janowitz-Koch et al. 2019) |
| Ots_U5049-250 | Neutral | G | T | (Janowitz-Koch et al. 2019) |
| Ots_U5121-34 | Neutral | A | G | (Janowitz-Koch et al. 2019) |
| Ots_u6-75 | Neutral | C | T | (Janowitz-Koch et al. 2019) |
| Ots_unk1104-38 | Neutral | C | T | (Janowitz-Koch et al. 2019) |


| Locus Name | Purpose | Allele 1 | Allele 2 | Reference |
| :---: | :---: | :---: | :---: | :---: |
| Ots_unk1832-39 | Neutral | C | T | (Janowitz-Koch et al. 2019) |
| Ots_unk3513-49 | Neutral | C | T | (Janowitz-Koch et al. 2019) |
| Ots_unk526 | Neutral | A | G | (Janowitz-Koch et al. 2019) |
| Ots_unk7936-50 | Neutral | C | G | (Janowitz-Koch et al. 2019) |
| Ots_unk9480-51 | Neutral | G | C | (Janowitz-Koch et al. 2019) |
| Ots_vatf-251 | Neutral | G | Deletion | (Janowitz-Koch et al. 2019) |
| Ots_zn593-346 | Neutral | A | T | (Janowitz-Koch et al. 2019) |
| Ots_ZR-575 | Neutral | G | A | (Janowitz-Koch et al. 2019) |
| Ots28_11073102 | Adaptive | T | A | (Narum et al. 2018) |
| Ots28_11202863 | Adaptive | C | A | (Narum et al. 2018) |
| Ots28_11186543 | Adaptive | A | T | (Narum et al. 2018) |
| Ots28_11033282 | Adaptive | G | A | (Narum et al. 2018) |
| Ots28_11202400 | Adaptive | C | T | (Narum et al. 2018) |
| Ots28_11062192 | Adaptive | C | G | (Narum et al. 2018) |
| Ots28_11025336 | Adaptive | A | C | (Narum et al. 2018) |
| Ots28_11095755 | Adaptive | A | T | (Narum et al. 2018) |
| Ots28_11077576 | Adaptive | A | G | (Narum et al. 2018) |
| Ots28_11202190 | Adaptive | T | C | (Narum et al. 2018) |
| Ots28_11077172 | Adaptive | G | A | (Narum et al. 2018) |
| Ots28_11160599 | Adaptive | G | T | (Narum et al. 2018) |
| Ots28_11205993 | Adaptive | C | T | (Narum et al. 2018) |
| Ots28_11075712 | Adaptive | C | T | (Narum et al. 2018) |
| Ots28_11072994 | Adaptive | C | T | (Narum et al. 2018) |
| Ots28_11164637 | Adaptive | C | A | (Narum et al. 2018) |
| Ots28_11201129 | Adaptive | T | G | (Narum et al. 2018) |
| Ots28_11073668 | Adaptive | T | A | (Narum et al. 2018) |
| Ots28_11023212 | Adaptive | A | G | (Narum et al. 2018) |
| Ots28_11206740 | Adaptive | T | C | (Narum et al. 2018) |
| Ots28_11143508 | Adaptive | G | A | (Narum et al. 2018) |
| Ots28_11070757 | Adaptive | A | G | (Narum et al. 2018) |
| Ots28_11071377 | Adaptive | T | C | (Narum et al. 2018) |
| Ots28_11077016 | Adaptive | C | T | (Narum et al. 2018) |
| Ots28_11207428 | Adaptive | T | G | (Narum et al. 2018) |
| Ots28_11210919 | Adaptive | C | T | (Narum et al. 2018) |
| Ots28_11205423 | Adaptive | A | G | (Narum et al. 2018) |
| Ots28_11075348 | Adaptive | G | A | (Narum et al. 2018) |
| Ots37124-12267397 | Adaptive | C | T | SWFSC - Clemento unpubl. |
| Ots37124-12272852 | Adaptive | C | T | SWFSC - Clemento unpubl. |
| Ots37124-12277401 | Adaptive | T | A | SWFSC - Clemento unpubl. |
| Ots37124-12281207 | Adaptive | A | T | SWFSC - Clemento unpubl. |
| Ots37124-12310649 | Adaptive | A | T | SWFSC - Clemento unpubl. |
| Ots19_46172427 | Adaptive | G | A | (Narum et al. 2018) |


| Locus Name | Purpose | Allele 1 | Allele 2 | Reference |
| :---: | :---: | :---: | :---: | :---: |
| Ots19_46172133 | Adaptive | C | T | (Narum et al. 2018) |
| Ots17_22360456 | Adaptive | T | G | (Narum et al. 2018) |
| Ots14_5453033 | Adaptive | G | A | (Narum et al. 2018) |
| Ots4_42378741 | Adaptive | C | T | (Narum et al. 2018) |
| Ots5_70908626 | Adaptive | T | C | (Narum et al. 2018) |
| Ots11_32418659 | Adaptive | A | T | (Narum et al. 2018) |
| Ots18_3550047 | Adaptive | A | G | (Narum et al. 2018) |
| Ots3_57055518 | Adaptive | T | C | (Narum et al. 2018) |
| Ots4_41638710 | Adaptive | G | A | (Narum et al. 2018) |
| Ots29_18791740 | Adaptive | T | G | (Narum et al. 2018) |
| Ots9_16115048 | Adaptive | G | A | (Narum et al. 2018) |
| Ots29_23344676 | Adaptive | T | C | (Narum et al. 2018) |
| Ots4_40942276 | Adaptive | G | A | (Narum et al. 2018) |
| Ots30_17330688 | Adaptive | T | C | (Narum et al. 2018) |
| Ots22_32650802 | Adaptive | G | A | (Narum et al. 2018) |
| Ots3_34894254 | Adaptive | T | C | (Narum et al. 2018) |
| Ots30_17330452 | Adaptive | G | C | (Narum et al. 2018) |
| Ots7_50997124 | Adaptive | G | T | (Narum et al. 2018) |
| Ots18_3426299 | Adaptive | T | A | (Narum et al. 2018) |
| Ots6_10904949 | Adaptive | C | T | (Narum et al. 2018) |
| Ots18_29943476 | Adaptive | A | G | (Narum et al. 2018) |
| Ots4_64978818 | Adaptive | C | A | (Narum et al. 2018) |
| Ots10_21244146 | Adaptive | A | C | (Narum et al. 2018) |
| Ots17_885364 | Adaptive | C | A | (Narum et al. 2018) |
| Ots2_38264269 | Adaptive | A | C | (Narum et al. 2018) |
| Ots33_19359879 | Adaptive | T | C | (Narum et al. 2018) |
| Ots6_33505144 | Adaptive | T | A | (Narum et al. 2018) |
| Ots5_44795073 | Adaptive | C | T | (Narum et al. 2018) |
| Ots18_32088284 | Adaptive | T | C | (Narum et al. 2018) |
| Ots15_18157381 | Adaptive | C | T | (Narum et al. 2018) |
| Ots12_23066874 | Adaptive | A | G | (Narum et al. 2018) |
| Ots2_42405643 | Adaptive | G | T | (Narum et al. 2018) |
| Ots7_51409415 | Adaptive | T | C | (Narum et al. 2018) |
| Ots1_72858599 | Adaptive | A | G | (Narum et al. 2018) |
| Ots7_53291035 | Adaptive | G | A | (Narum et al. 2018) |
| Ots7_53631522 | Adaptive | A | G | (Narum et al. 2018) |
| Ots18_30099101 | Adaptive | C | T | (Narum et al. 2018) |
| Ots11_11925999 | Adaptive | G | T | (Narum et al. 2018) |
| Ots18_3541813 | Adaptive | T | C | (Narum et al. 2018) |
| Ots9_28975221 | Adaptive | A | T | (Narum et al. 2018) |
| Ots_CHI06048618_5222 | Adaptive | T | G | Chen unpublished |
| Ots_CHI06105101_18523 | Adaptive | A | G | Chen unpublished |


| Locus Name | Purpose | Allele 1 | Allele 2 | Reference |
| :--- | :---: | :---: | :---: | :---: |
| Ots_CHI06105101_16717 | Adaptive | C | T | Chen unpublished |
| Ots_CHI06035945_4547 | Adaptive | C | T | Chen unpublished |
| Ots_CHI06027687_14347 | Adaptive | G | A | Chen unpublished |
| 7 | Adaptive | A | C | (Narum et al. 2018) |
| Ots18_3417174 | Adaptive | G | C | (Narum et al. 2018) |
| Ots11_32468959 | Adaptive | T | A | (Narum et al. 2018) |
| Ots7_54212944 | Sex ID | X | Y | (Janowitz-Koch et al. 2019) |
| Ots_SEXY3-1 |  |  |  |  |

Janowitz-Koch, I., Rabe, C., Kinzer, R., Nelson, D., Hess, M.A., and Narum, S.R. 2019. Longterm evaluation of fitness and demographic effects of a Chinook Salmon supplementation program. Evol. Appl. 12: 456-469. doi:10.1111/eva. 12725.
Narum, S.R., Genova, A.D., Micheletti, S.J., and Maass, A. 2018. Genomic variation underlying complex life-history traits revealed by genome sequencing in Chinook salmon. Proc. R. Soc. B 285(1883): 20180935. doi:10.1098/rspb.2018.0935.

## Appendix B. WDFW GTseq genotyping protocol details

The genotyping was done using a cost-effective method based on custom amplicon sequencing called Genotyping in Thousands (GTseq) (Campbell et al. 2015). GTseq is an efficient genotyping method that amplifies pools of targeted SNPs and then indexes individual samples. The pools are sequenced, de-multiplexed, and genotyped by generating a ratio of allele counts for each individual. The entire process can be broken down into four segments; extraction, library preparation, sequencing, and genotyping.

Genomic DNA was extracted for all samples by digesting a small piece of fin tissue with a Macherey-Nagel 96 column NucleoSpin kit, following the manufacturers recommendations (Macherey-Nagel GmbH \& Co. KG, Duren, Germany). The DNA was then concentrated 2.5 times before proceeding to library preparation. Next, the multiplexed pool of targeted loci was amplified. The multiplex PCR consisted of 2 uL of cleaned DNA extract, 3.5uL of Qiagen Multiplex PCR Plus mix (Qiagen, 10672201), and 1.5 uL pooled primer mix (IDT, Appendix A, final volume $=7 \mathrm{uL}$; final primer concentrations at each locus $=54 \mathrm{nM})$. Thermal cycling conditions were as follows: $95^{\circ} \mathrm{C}-15 \mathrm{~min} ; 5$ cycles $\left[95^{\circ} \mathrm{C}-30 \mathrm{~s}, 5 \%\right.$ ramp down to $57^{\circ} \mathrm{C}-30 \mathrm{~s}$, $\left.72^{\circ} \mathrm{C}-2 \mathrm{~min}\right] ; 10$ cycles $\left[95^{\circ} \mathrm{C}-30 \mathrm{~s}, 65^{\circ} \mathrm{C}-30 \mathrm{~s}, 72^{\circ} \mathrm{C}-30 \mathrm{~s}\right] ; 4^{\circ} \mathrm{C}$ hold. Following the multiplex PCR, the amplified samples were diluted 20 -fold. 3 uL of diluted multiplex PCR product was then used in the barcoding PCR. The barcoding PCR adds indexes that identify each sample by well and by plate. For the barcoding PCR, 1 uL of 10 uM well-specific i5 tagging primer (IDT) and 1 uL of 10 uM plate-specific i7 tagging primer were added to the 3 uL of amplified sample. 5uL of Qiagen Multiplex PCR Plus mix (Qiagen, 10672201) was then added for a final reaction volume of 10 uL . Thermal cycling conditions were: $95^{\circ} \mathrm{C}-15 \mathrm{~min} ; 10$ cycles $\left[98^{\circ} \mathrm{C}-10 \mathrm{~s}, 65^{\circ} \mathrm{C}-30 \mathrm{~s}, 72^{\circ} \mathrm{C}-30 \mathrm{~s}\right] ; 72^{\circ} \mathrm{C}-5 \mathrm{~min} ; 4^{\circ} \mathrm{C}$ hold. Following the barcode PCR, each plate of samples (library) was normalized using the SequalPrepTM Normalization Plate Kit (Applied Biosystems, A1051001) according to the manufacturer's instructions. Upon completion of normalization, 10 uL of each sample per 96 -well plate was pooled into a 1.5 mL tube constituting a library. A purification step was then performed on each library with Agencourt AMPure ${ }^{\circledR}$ XP magnetic beads (Agencourt, A63881) according to the manufacturer's instructions for size selection with a $2: 1$ and 1.43:1 ratio of library to beads. The purified libraries were then eluted with 15 uL of TE pH 8.0 . In order to complete the final process of library preparation, each library was quantified and normalized. The libraries were quantified using a Qubit 3 Fluorometer (Invitrogen) and QubitTMdsDNA HS Assay Kit reagents (Invitrogen, Q32854) according to the manufacturer's instructions. Following the quantification, the concentration of each library was calculated using the molecular weight specific to the multiplex pool used (i.e. One.382). Then each library was normalized to 4 nM and pooled with other libraries that were sequenced on the same sequencing run. Pooled libraries were then sequenced at a 2.5 pM loading concentration on an Illumnia NextSeq 500 instrument of a single-end read flow cell using 111 cycles with dualindex reads of six cycles each. To genotype the samples, a bioinformatics pipeline was used. This pipeline is explained and available online at https://github.com/GTseq/GTseq-Pipeline (Campbell et al. 2015). Essentially, there are a series of custom perl scripts that ultimately count amplicon-specific sequences for each allele. Allele ratios are then used to generate genotypes.

Campbell, N.R., Harmon, S.A., and Narum, S.R. 2015. Genotyping-in-Thousands by sequencing (GT-seq): A cost effective SNP genotyping method based on custom amplicon sequencing. Mol. Ecol. Res. 15(4): 855-867. doi:10.1111/1755-0998.12357.

# The Effects of Hatchery Supplementation on Size and Age At Maturity and Fecundity of Spring and Summer Chinook Salmon in the Upper Columbia Basin 

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#### Abstract

A common objective of conservation hatchery programs is to produce fish with phenotypic traits similar to individuals from the natural population. We evaluated the size and age at maturity, and fecundity of five spring Chinook Salmon and three summer Chinook Salmon hatchery programs. Comparisons to natural-origin fish from the targeted populations over multiple generations and during two periods of hatchery production releases (high and low number of smolts) were made for applicable programs. Generally, the hatchery-populations were composed of more, younger fish and fewer older fish, but the predominate age at maturity was similar between hatchery- and natural-origin populations. The size at maturity of returning adults was significantly affected by age and sex, and in some cases origin and period. However, when matched by age and sex, the differences in size at maturity by origin and period that were statistically significant were minor and likely of little biological relevance. For all populations, the fecundity metrics were significantly affected by fish size and weight, but fecundity differed between origins in only two populations. For all metrics, the high/low program release numbers did not substantially affect these phenotypic traits. Future analyses of these metrics should consider changes over time and consider trends between hatchery-origin and natural-origin fish. Declines in age and size at maturity are a range-wide concern for Pacific salmonids and therefore a temporal analysis of the dataset generated for the monitoring of these hatchery programs would benefit the region.


## Introduction

Conservation hatchery programs should produce fish that maintain and match the phenotypic characteristics of the targeted natural population. The objective of maintaining phenotypic characteristics is to limit introgression and domestication effects while the objective of matching is intended to produce fish with any fitness benefits conferred with the phenotypic characteristics of natural-origin fish. In the Upper Columbia River Basin, conservation hatchery programs have collected data on phenotypic variables including size at maturity, age at maturity, and fecundity of natural-origin and hatchery-origin Chinook Salmon for nearly 30 years. These data provide the opportunity to evaluate how these hatchery programs have performed in matching these phenotypic traits to their targeted population over multiple generations. The question of maintaining, or not affecting, natural-origin phenotypic traits is more complex due to the non-static nature of phenotypic variables and multiple factors that can affect the size and age at maturity, and fecundity of natural-origin fish. For example, throughout the range of Pacific Salmon and in all five species there has been a well-documented decline in size and age at maturity (Ricker 1981, Jeffrey et al. 2017, Losee et al. 2019). Hatchery-origin fish introgression, as well as competition among hatchery- and natural-origin fish, are two of several theories for the observed changes. Other theories with supporting data include size-selective harvest, environmental changes such as ocean temperature and productivity, inter-species competition, and marine mammal predation (for a review of each theory see Ohlberger et al. 2018).

In this chapter, we test whether the age and size at maturity, and fecundity of returning hatchery-origin spring and summer Chinook Salmon from the Upper Columbia River Basin matches those phenotypic characteristics in their targeted natural-origin populations, and answer three programmatic monitoring questions: (1) Is the age at maturity of hatchery- and naturalorigin fish similar at the time of spawning? (2) Is the size at maturity of a given age and sex of hatchery-origin fish similar to the size at maturity of the same age and sex of natural-origin fish? and, (3) Is the size versus fecundity relationship of hatchery- and natural-origin fish similar?

## Methods

## Data Collection and Derived Metrics

Spring and summer Chinook Salmon spawning ground surveys were used to collect data to estimate origin, age and size at maturity, and sex of carcasses within the Wenatchee and Methow subbasins. Spawning ground surveys were also used to collect data on summer Chinook Salmon in the lower Chelan River. Field methods were consistent with those described in Gallagher et al. (2007) and Murdoch et al. (2010). Surveyors walked or floated the entire distribution of spring and summer Chinook Salmon spawning habitat and sampled carcasses weekly throughout the spawning season. Chinook Salmon carcasses were counted and examined to determine sex, origin (e.g., hatchery- or natural-origin), size at maturity, distribution, and other biological characteristics. In addition, scales collected from carcasses were used to determine fish age at maturity and verify origin. The sampling goal for carcasses was at least $20 \%$ of the spawning population.

Age at maturity was estimated from scales collected from each carcass sampled. At least four scales were removed from the left side of the fish about two rows above the lateral line and
on the diagonal row from the posterior insertion of the dorsal fin to the anterior insertion of the anal fin. Following methods in Quist et al. (2012), cellulose acetate impressions of each scale were made and read to determine the total age (i.e., egg-to-spawning adult) and ocean (salt) age of each fish. For spring Chinook Salmon, we evaluated differences between hatchery- and natural-origin fish using total age. For summer Chinook Salmon, we used ocean age to evaluate differences between hatchery- and natural-origin fish, because hatchery-origin summer Chinook Salmon were generally released after one year of rearing, while natural-origin summer Chinook Salmon migrate primarily as age-0 fish. In this report, we only analyzed hatchery-origin summer Chinook Salmon that were released as age-1 fish, because those releases had a natural-origin component for comparison. There were no natural-origin comparisons for the age-0 summer Chinook releases (e.g., Wells Hatchery and Turtle Rock Acclimation Facility age-0 releases).

Size at maturity was estimated by measuring the post-orbital to hypural length ( $\mathrm{POH} ; \mathrm{cm}$ ) of all carcasses sampled. Because of erosion of the caudal fin and secondary morphological changes to the head of Chinook Salmon, POH provides a more robust measure of fish size than does fork length or total length. Sex was determined by examination of gonads. For each carcass sampled, origin was determined by examining tags (e.g., CWT and PIT tags), marks (missing adipose fin), and scale analysis. Nearly all hatchery-origin fish were marked with a tag such as a CWT and/or adipose fin clipped prior to release.

Fecundity at size data for both spring and summer Chinook Salmon were collected from broodstock. To enumerate fecundity, an optical counter enumerated eyed eggs and the number of eyed eggs was added to the number of dead eggs enumerated during incubation. To determine gonadal mass, a representative sample of 50 female Chinook Salmon from across the spawning period were sampled for numbers and weight of green eggs. For each female selected, the entire gamete mass was extracted, ovarian fluid drained, and weighed to the nearest 0.1 g . A subsample of 100 green eggs were counted and weighed to within 0.01 g to estimate individual egg weight for each female. Total fecundity of each female was estimated by dividing the weight of the total egg mass by the calculated mean individual egg weight. Because each sample of the total egg mass likely contained small amounts of ovarian fluid, fecundity estimates may be overestimated slightly. Females were also measured for length ( POH ), weighed (post-spawning; kg), and aged following methods described earlier. Origin was determined by examining tags, marks, and scale analysis. Females were stratified by fork-length categories to obtain a representative fecundity sample for all sizes of fish. This was needed to better estimate the relationship between size and fecundity. Complete fecundity data only exist for brood years 2014 to present.

## Data Analysis

We conducted separate analyses for each program. For spring Chinook Salmon, we evaluated differences between hatchery- and natural-origin age and size at maturity, and fecundity for the Chiwawa River, Nason Creek, White River, Methow/Chewuch River, and Twisp River programs. For summer Chinook Salmon, we conducted separate analyses for the Wenatchee River, Methow River, and Chelan River programs. We only analyzed fecundity data for Wells summer Chinook Salmon because there were no age at maturity or size at maturity data for natural-origin summer Chinook Salmon to compare with Wells summer Chinook Salmon. For Nason Creek spring Chinook Salmon we evaluated two different program effects. First, we examined the differences in age and size of maturity of hatchery-origin strays and natural-origin spring Chinook Salmon in Nason Creek. In this case, the "before" period (1993-2000)
represented the time period when the proportion of hatchery-origin spawners ( $\mathrm{pHOS} \mathrm{)} \mathrm{in} \mathrm{Nason}$ Creek was low (mean $<0.10$ ) and the "after" period (2001-2018) represented the time period when pHOS was high (mean $>0.50$ ). Second, we evaluated the effects of the Nason Creek Spring Chinook Hatchery Program on age and size at maturity of spring Chinook Salmon in Nason Creek. In this case, the "before" period (1993-2015) represents the time period before the implementation of the Nason Creek spring Chinook Salmon program and the "after" period (2016-2018) represents the time period following the initiation of the hatchery program. For White River spring Chinook Salmon, the "before" period (1993-2007) represents the time before the influence of the spring Chinook Salmon captive brood program, while the "after" period (2008-2018) represents the time during the influence of the program.

We evaluated differences between hatchery- and natural-origin spring and summer Chinook Salmon for age and size at maturity as a complete time series (1993-2018) and an interrupted time series for four of the programs. The interrupted time series included the period "before" the reduction in hatchery production (1993-2014) and the period "after" the reduction in hatchery production (2015-2018) for most hatchery programs. For summer Chinook Salmon in the Chelan River, we used data collected from 2000 to 2018, because that is the period when intensive spawning ground surveys were conducted and there were no subyearling releases. There was no change in hatchery production of Wells summer Chinook Salmon; therefore, there was no interrupted time series to evaluate for Wells summer Chinook Salmon. The specific years for the Nason Creek and White River spring Chinook Salmon programs did not follow the same before and after dates as other programs (see below). Two-way Yates’ Chi-square determined whether age at maturity of hatchery- and natural-origin Chinook Salmon differed significantly. Because of different age-at-migration characteristics, we evaluated male and female Chinook Salmon separately. For size at maturity, we evaluated the length ( POH ) of hatchery- and naturalorigin Chinook Salmon of the same age (total age for spring Chinook Salmon and ocean age for summer Chinook Salmon). Three-way, unbalanced, general linear models, analysis of variance (GLM ANOVA) tested differences in sizes of hatchery- and natural-origin fish. This analysis included sex, origin, and age as fixed factors in the model. Where there were enough data, a four-way, unbalanced, GLM ANOVA tested differences in sizes of hatchery- and natural-origin fish before and after changes in programs. For all analyses, we interpreted a significant difference ( $\mathrm{P} \leq 0.05$ ) as an indication of differences between hatchery- and natural-origin fish.

For both hatchery- and natural-origin spring and summer Chinook Salmon, we used simple linear regression to describe fecundity as a function of female size (both fork length and weight), and mean egg weight and total egg weight as a function of female size. Analysis of covariance (ANCOVA) examined fecundity relationships between hatchery- and natural-origin Chinook Salmon. In this case, ANCOVA tested whether the slopes of the regression lines were the same for hatchery- and natural-origin fish. Here, we focused only on the F-test and its Pvalue for the origin-by-size interaction term in the analysis of variance table. Because we have fecundity data only for 2014-2018, we cannot evaluate changes in relationships before and after reduction in fish production numbers for each supplementation program. Thus, all available years were pooled together in the analysis.

## Results

Spring Chinook Salmon

## Chiwawa River Program

Age-at-Maturity Analysis-For the complete time series (1993-2018), age at maturity differed significantly between hatchery- and natural-origin female (Yates' Chi-square $=204.070$, $\mathrm{P}=0.000$ ) and male spring Chinook Salmon (Yates' Chi-square $=138.640, \mathrm{P}=0.000$ ) in the Chiwawa River (Figure 1). Most females and males returned at age-4; however, a larger proportion of hatchery-origin fish returned at younger ages than did natural-origin fish. For example, $94 \%$ of hatchery-origin females and $78 \%$ of the natural-origin females returned at age4. About $22 \%$ of natural-origin females and $5 \%$ of hatchery-origin females returned at age-5. A similar pattern was observed with males: about $27 \%$ of the hatchery-origin males and $9 \%$ of natural-origin males returned at age-3. In contrast, about $21 \%$ of natural-origin males and $7 \%$ of hatchery-origin males returned at age- 5 .

There was little change in age at maturity before (1993-2014) and after (2014-2018) reduction in hatchery production numbers, although a larger fraction of hatchery-origin males returned as age-4 fish following the reduction in program production than before the reduction (Figure 1). During both the before and after periods, the age at maturity differed significantly between hatchery- and natural-origin female (Before: Yates' Chi-square $=167.543, \mathrm{P}=0.000$; After: Yates' Chi-square $=34.387, \mathrm{P}=0.000$ ) and male spring Chinook Salmon (Before: Yates' Chi-square $=119.657, \mathrm{P}=0.000$; After: Yates' Chi-square $=15.282 ; \mathrm{P}=0.000$ ) in the Chiwawa River.

Size-at-Maturity Analysis-For the complete time series (1993-2018), size at maturity differed significantly between hatchery- and natural-origin spring Chinook Salmon in the Chiwawa River (GLM ANOVA F = 6.666, $\mathrm{P}=0.010$ ) (Figure 2). Size at maturity was significantly affected by age, origin, and sex (GLM ANOVA F = 5.025, P = 0.007). For females, age- 3 hatchery-origin fish were larger on average than age- 3 natural-origin fish (mean difference $=9 \mathrm{~cm}$ ). However, age- 5 natural-origin fish were on average larger than age- 5 hatchery-origin fish (mean difference $=2 \mathrm{~cm}$ ). Likewise, for age- 5 males, natural-origin fish were slightly larger than hatchery-origin fish (mean difference $=1 \mathrm{~cm}$ ). In contrast, age-3 and age-4 hatchery-origin males were slightly larger on average than natural-origin males of the same age.

There were differences in size at maturity before and after reduction in hatchery production numbers, although sample sizes for some age groups were small (Figure 2). The significant four-way interaction term (GLM ANOVA F $=3.928, \mathrm{P}=0.048$ ), which only included age-4 and 5 fish because of sample sizes, indicated that differences in sizes between hatcheryand natural-origin spring Chinook Salmon were affected by age, sex, and period. This was most apparent for males. During the before period, age-4 hatchery-origin males were slightly larger than age-4 natural-origin males, while age- 5 hatchery-origin males were slightly smaller than age- 5 natural-origin males. This reversed during the after period. Differences in average sizes were small ( $\leq 2 \mathrm{~cm}$ ), however.

Fecundity Analysis-There were significant linear relationships between female size and fecundity, mean egg weight, and gonadal mass for both hatchery-origin and natural-origin spring Chinook Salmon in the Chiwawa River for the combined years 2014-2018 (Figure 3). Fecundity relationships between hatchery-origin and natural-origin spring Chinook Salmon differed significantly for fork length and fecundity (ANCOVA F $=4.644, \mathrm{P}=0.032$ ) and fork length and mean egg weight (ANCOVA F $=4.307, \mathrm{P}=0.039$ ). There were no differences in relationships
between hatchery- and natural-origin females for weight and fecundity (ANCOVA F $=3.359, \mathrm{P}$ $=0.068)$ and fork length and gonadal mass (ANCOVA $\mathrm{F}=0.513, \mathrm{P}=0.474)$.

## Chiwawa River Spring Chinook Salmon



Figure 1. Proportion of hatchery- and natural-origin female and male spring Chinook Salmon spawners of different ages sampled on the spawning grounds in the Chiwawa River basin for the combined years 1993-2018 (top row), before reduction (1993-2014; middle row), and after reduction (2015-2018; bottom row) in hatchery smolt production. Sample sizes for females $=956$ natural-origin and 2,322 hatchery-origin fish and for males $=647$ natural-origin and 1,243 hatchery-origin fish for the combined analysis. For the before analysis, sample sizes for females $=748$ natural-origin and 1,969 hatchery-origin fish and for males $=522$ natural-origin and 1,091 hatchery-origin fish; and for the after analysis, sample sizes for females $=208$ natural-origin and 353 hatchery-origin fish and for males $=125$ natural-origin and 151 hatchery-origin fish.

Chiwawa River Spring Chinook Salmon


Figure 2. Mean POH length $(\mathrm{cm})$ and $95 \% \mathrm{CI}$ of hatchery- and natural-origin female and male spring Chinook Salmon spawners of different ages sampled on the spawning grounds in the Chiwawa River basin for the combined years 1993-2018 (top row), before reduction (1993-2014; middle row), and after reduction (2015-2018; bottom row) in hatchery smolt production. Sample sizes are shown within each bar.

## Chiwawa River Spring Chinook Salmon



Figure 3. Relationships between fecundity (number of eggs, mean weight of eggs, and gonadal mass) and size ( POH length and weight) for hatchery- (dashed line) and natural-origin (solid line) Chiwawa River spring Chinook Salmon for return years 2014-2018. Simple linear regression statistics are shown in each figure.

## Effects of Hatchery-origin Strays on Natural-origin Spawners in Nason Creek

Age-at-Maturity Analysis-For the complete time series (1993-2018), age at maturity differed significantly between hatchery-origin and natural-origin female (Yates' Chi-square $=$ $133.741, \mathrm{P}=0.000$ ) and male spring Chinook Salmon (Yates' Chi-square $=232.691, \mathrm{P}=0.000$ ) in Nason Creek (Figure 4). Most female and male spring Chinook Salmon returned at age-4; however, a larger proportion of hatchery-origin fish returned at younger ages than did naturalorigin fish. For example, $94 \%$ of hatchery-origin females and $77 \%$ of natural-origin females returned at age-4. About $23 \%$ of natural-origin females, but only $5 \%$ of hatchery-origin females returned at age-5. A similar pattern was observed with male spring Chinook Salmon. About 44\% of the hatchery-origin males, and only $7 \%$ of natural-origin males returned at age-3. In contrast, about $19 \%$ of natural-origin males, and only $4 \%$ of hatchery-origin males returned at age- 5 .

There were changes in age at maturity before (1993-2000) and after (2001-2018) the increase in pHOS in Nason Creek and this was most apparent in males (Figure 4). For males, the proportion of age- 3 returns decreased from the before to after period, while age- 4 males
increased from the before to after period. During both the before and after periods, age at maturity differed significantly between hatchery- and natural-origin female (Before: Yates' Chisquare $=5.061, \mathrm{P}=0.024$; After: Yates' Chi-square $=119.625, \mathrm{P}=0.000$ ) and male spring Chinook Salmon (Before: Yates' Chi-square $=28.501, \mathrm{P}=0.000$; After: Yates' Chi-square $=$ 204.031; $\mathrm{P}=0.000$ ) in Nason Creek.

Size-at-Maturity Analysis-For the complete time series (1993-2018), size at maturity did not differ significantly between hatchery- and natural-origin female or male spring Chinook Salmon in Nason Creek (GLM ANOVA F $=2.384, \mathrm{P}=0.123$ ) (Figure 5). There were, however, significant differences in size at maturity among age classes (GLM ANOVA F $=677.8, \mathrm{P}=$ 0.000 ) and sex (GLM ANOVA F = 7.956, $\mathrm{P}=0.005$ ). Regardless of origin, older fish were larger than younger fish and older males were larger than older females.

Size at maturity did not differ significantly between hatchery- and natural-origin female or male spring Chinook Salmon from before (1993-2000) to after (2001-2018) the increase in pHOS in Nason Creek (GLM ANOVA F $=1.737, \mathrm{P}=0.188$ ) (Figure 5). There were significant differences in size at maturity between male and female fish (GLM ANOVA F $=4.064, \mathrm{P}=$ 0.044 ) and among ages (GLM ANOVA $\mathrm{F}=120.678, \mathrm{P}=0.000$ ), but those differences did not change from the before to after period (GLM ANOVA F $=1.478, \mathrm{P}=0.224$ ).

Fecundity Analysis-There were significant linear relationships between female size and fecundity, mean egg weight, and gonadal mass for both hatchery- and natural-origin spring Chinook Salmon in Nason Creek for the combined years 2014-2018 (Figure 6). There were no differences in relationships between hatchery- and natural-origin fish for fork length and fecundity (ANCOVA $\mathrm{F}=1.959, \mathrm{P}=0.163$ ), female weight and fecundity ( $\mathrm{ANCOVAF}=3.273$, $\mathrm{P}=0.072$ ), fork length and mean egg weight (ANCOVA $\mathrm{F}=0.135, \mathrm{P}=0.714$ ) or fork length and gonadal mass (ANCOVA F $=2.219, \mathrm{P}=0.138$ ).

## Nason Creek Spring Chinook Hatchery Program

Age-at-Maturity Analysis—Results for the complete time series (1993-2018) in Nason Creek were described above under the Effects of Hatchery-origin Strays on Natural-origin Spawners in Nason Creek section.

There was little change in age at maturity of spring Chinook Salmon before (1993-2015) and during (2016-2018) the implementation of the Nason Creek Supplementation Program (Figure 7). For both the before and during supplementation periods, the age at maturity differed significantly between hatchery- and natural-origin female (Before: Yates' Chi-square $=124.694$, $\mathrm{P}=0.000$; After: Yates' Chi-square $=7.788, \mathrm{P}=0.005$ ) and male spring Chinook Salmon (Before: Yates' Chi-square $=213.035, \mathrm{P}=0.000$; After: Yates' Chi-square $=15.032 ; \mathrm{P}=0.000$ ) in Nason Creek.

Size-at-Maturity Analysis-Results for the complete time series (1993-2018) in Nason Creek were described above under the Effects of Hatchery-origin Strays on Natural-origin Spawners in Nason Creek section.

Because of a lack of data for the during supplementation period, we could not evaluate statistically the effects of period, age, sex, and origin on size at maturity in Nason Creek. However, there was a difference in size among age classes, with older fish being larger than younger fish (Figure 8). In addition, there were differences in sizes between males and females, with older males generally larger than older females.

Fecundity Analysis-These results were described above under the Straying Effects on Nason Creek section.

Nason Creek Spring Chinook Salmon (pHOS)


Figure 4. Proportion of hatchery- and natural-origin female and male spring Chinook Salmon spawners of different ages sampled on the spawning grounds in Nason Creek for the combined years 1993-2018 (top row), before increase in pHOS (1993-2000; middle row), and after increase in pHOS (2001-2018; bottom row). Sample sizes for females $=705$ natural-origin and 1,086 hatchery-origin fish and for males $=495$ natural-origin and 883 hatchery-origin fish for the combined analysis. For the before analysis, sample sizes for females $=69$ natural-origin and 33 hatchery-origin fish and for males $=59$ natural-origin and 42 hatchery-origin fish; and for the after analysis, sample sizes for females $=636$ natural-origin and 1,053 hatchery-origin fish and for males $=436$ natural-origin and 841 hatchery-origin fish.

Nason Creek Spring Chinook Salmon (pHOS)


Figure 5. Mean POH length (cm) and $95 \%$ CI of hatchery- and natural-origin female and male spring Chinook Salmon spawners of different ages sampled on the spawning grounds in Nason Creek for the combined years 1993-2018 (top row), before increase in pHOS (1993-2000; middle row), and after increase in pHOS (2001-2018; bottom row). Sample sizes are shown within each bar.

## Nason Creek Spring Chinook Salmon



Figure 6. Relationships between fecundity (number of eggs, mean weight of eggs, and gonadal mass) and size ( POH length and weight) for hatchery- (dashed line) and natural-origin (solid line) Nason Creek spring Chinook Salmon for return years 2014-2018. Simple linear regression statistics are shown in each figure.

Nason Creek Spring Chinook Salmon (Supplementation)


Figure 7. Proportion of hatchery- and natural-origin female and male spring Chinook Salmon spawners of different ages sampled on the spawning grounds in Nason Creek for the combined years 1993-2018 (top row), before supplementation (1993-2015; middle row), and during supplementation (2016-2018; bottom row). Sample sizes for females $=705$ natural-origin and 1,086 hatchery-origin fish and for males $=495$ natural-origin and 883 hatchery-origin fish for the combined analysis. For the before analysis, sample sizes for females $=647$ natural-origin and 1,008 hatchery-origin fish and for males $=433$ natural-origin and 814 hatchery-origin fish; and for the during analysis, sample sizes for females $=58$ natural-origin and 78 hatchery-origin fish and for males $=62$ natural-origin and 69 hatchery-origin fish.

Nason Creek Spring Chinook Salmon (Supplementation)


Figure 8. Mean POH length $(\mathrm{cm})$ and $95 \% \mathrm{CI}$ of hatchery- and natural-origin female and male spring Chinook Salmon spawners of different ages sampled on the spawning grounds in Nason Creek for the combined years 1993-2018 (top row), before supplementation (1993-2015; middle row), and during supplementation (2016-2018; bottom row). Sample sizes are shown within each bar.

## White River Program

Age-at-Maturity Analysis-For the complete time series (1993-2018), age at maturity differed significantly between hatchery- and natural-origin females (Yates' Chi-square $=24.659$, $\mathrm{P}=0.000$ ) but not between hatchery- and natural-origin male spring Chinook Salmon (Yates’ Chi-square $=5.119, \mathrm{P}=0.077$ ) in the White River (Figure 9). Most females and males returned at age- 4 ; however, a larger proportion of hatchery-origin fish returned at younger ages than did natural-origin fish. For example, $95 \%$ of hatchery-origin females and $72 \%$ of natural-origin
females returned at age-4. About $28 \%$ of natural-origin females and 5\% of hatchery females returned at age- 5 . A similar pattern was observed with males: about $22 \%$ of natural-origin males returned at age-5, while no hatchery males returned at age- 5 .

There was little change in age at maturity before (1993-2007) and during (2008-2018) the implementation of the captive-brood program in the White River (Figure 9). For both the before and during periods, the age at maturity differed significantly between hatchery- and naturalorigin females (Before: Yates' Chi-square $=12.891, \mathrm{P}=0.000$; After: Yates' Chi-square $=$ $10.315, \mathrm{P}=0.001$ ) but not for male spring Chinook Salmon (Before: Yates' Chi-square $=5.584$, $\mathrm{P}=0.061$; After: Yates' Chi-square $=0.359 ; \mathrm{P}=0.359$ ).

Size-at-Maturity Analysis-Sample sizes were too small to evaluate statistically the effects of origin, sex, and age on size at maturity of spring Chinook Salmon in the White River. Over the complete time series (1993-2018), however, there were small differences in sizes between hatchery- and natural-origin fish within each age class (Figure 10). There were clear differences in sizes among ages with older fish larger than younger fish.

Because of a lack of data, we could not evaluate statistically the effects of period, age, sex, and origin on size at maturity in the White River. Differences in sizes among age classes were consistent during the before and during supplementation periods and differences in sizes between hatchery- and natural-origin fish were small during both periods (Figure 10).

Fecundity Analysis-No comparative fecundity data similar to what was done for other programs were collected for the White River program. This was because broodstock were collected as eyed eggs or fry from the White River (Ford et al. 2015).

White River Spring Chinook Salmon



Female Before Captive Brood





Figure 9. Proportion of hatchery- and natural-origin female and male spring Chinook Salmon spawners of different ages sampled on the spawning grounds in the White River basin for the combined years 1993-2018 (top row), before the captive brood program (1993-2007; middle row), and during the captive brood program (2008-2018; bottom row). Sample sizes for females $=151$ natural-origin and 125 hatchery-origin fish and for males $=65$ natural-origin and 29 hatchery-origin fish for the combined analysis. For the before analysis, sample sizes for females $=87$ natural-origin and 69 hatchery-origin fish and for males $=44$ natural-origin and 15 hatchery-origin fish; and for the during analysis, sample sizes for females $=64$ natural-origin and 56 hatchery-origin fish and for males $=21$ natural-origin and 14 hatchery-origin fish.

White River Spring Chinook Salmon


Figure 10. Mean POH length (cm) and 95\% CI of hatchery- and natural-origin female and male spring Chinook Salmon spawners of different ages sampled on the spawning grounds in the White River basin for the combined years 1993-2018 (top row), before the captive brood program (1993-2007; middle row), and during the captive brood program (2008-2018; bottom row). Sample sizes are shown within each bar.

## Methow-Chewuch Program

Age-at-Maturity Analysis-For the complete time series (1993-2018), age at maturity differed significantly between hatchery- and natural-origin female (Yates' Chi-square $=143.960$, $\mathrm{P}=0.000$ ) and male spring Chinook Salmon (Yates' Chi-square $=263.369, \mathrm{P}=0.000$ ) in the Methow and Chewuch rivers (Figure 11). Most females and males returned at age-4; however, a larger proportion of hatchery-origin fish returned at younger ages than did natural-origin fish. For example, $90 \%$ of hatchery-origin females and $77 \%$ of the natural-origin females returned at
age-4. About $23 \%$ of natural-origin females and $10 \%$ of hatchery-origin females returned at age5. A similar pattern was observed with males. About $33 \%$ of the hatchery-origin males and $7 \%$ of natural-origin males returned at age-3. In contrast, about $20 \%$ of natural-origin males and $9 \%$ of hatchery-origin males returned at age- 5 .

There was little change in age at maturity before (1993-2014) and after (2014-2018) reduction in hatchery production numbers, although a larger fraction of hatchery- and naturalorigin males returned as age-4 fish following the reduction in program production (Figure 11). During both the before and after periods, age at maturity differed significantly between hatcheryand natural-origin females (Before: Yates' Chi-square $=142.913, \mathrm{P}=0.000$; After: Yates' Chisquare $=10.169, \mathrm{P}=0.001$ ). Age at maturity differed significantly between hatchery- and natural-origin males before reduction (Yates' Chi-square $=250.188, \mathrm{P}=0.000$ ) but not after reduction in hatchery production (Yates' Chi-square $=3.773 ; \mathrm{P}=0.152$ ).

Size-at-Maturity Analysis-For the complete time series (1993-2018), size at maturity was significantly affected by sex and age (GLM ANOVA F $=82.040, \mathrm{P}=0.000$ ) but not by origin (GLM ANOVA $\mathrm{F}=3.807, \mathrm{P}=0.051$ ) (Figure 12). Older fish of both sexes were larger than younger fish and older males were generally larger than older females. Except for age-3 females, size differences between hatchery- and natural-origin fish within each age group were small, generally less than 0.5 cm on average.

There were significant differences in the size at maturity before and after reduction in hatchery production (GLM ANOVA F $=7.129, \mathrm{P}=0.008$ ) (Figure 12). For most age groups, both hatchery- and natural-origin fish were smaller following the reduction in hatchery production; however, the change in size was only about 1 cm on average. During both time periods for both hatchery- and natural-origin fish, older males were larger on average than females of the same age (GLM ANOVA F $=22.459, \mathrm{P}=0.000$ ).

Fecundity Analysis-There were significant linear relationships between female size and fecundity, mean egg weight, and gonadal mass for both hatchery- and natural-origin spring Chinook Salmon in the Methow and Chewuch rivers for the combined years 2014-2018 (Figure 13). There were no differences in relationships between hatchery- and natural-origin fish for fork length and fecundity (ANCOVA $\mathrm{F}=0.418, \mathrm{P}=0.518$ ), female weight and fecundity (ANCOVA $\mathrm{F}=0.261, \mathrm{P}=0.610$ ), fork length and mean egg weight ( $\mathrm{ANCOVAF}=0.316, \mathrm{P}=0.574$ ) and fork length and gonadal mass (ANCOVA $\mathrm{F}=2.123, \mathrm{P}=0.146$ ).

Methow/Chewuch Spring Chinook Salmon


Figure 11. Proportion of hatchery- and natural-origin female and male spring Chinook Salmon spawners of different ages sampled on the spawning grounds in the Methow and Chewuch rivers for the combined years 1993-2018 (top row), before reduction (1993-2014; middle row), and after reduction (2015-2018; bottom row) in hatchery smolt production. Sample sizes for females $=1,163$ natural-origin and 4,351 hatchery-origin fish and for males $=923$ natural-origin and 3,263 hatchery-origin fish for the combined analysis. For the before analysis, sample sizes for females $=946$ natural-origin and 3,873 hatchery-origin fish and for males $=772$ natural-origin and 3,050 hatchery-origin fish; and for the after analysis, sample sizes for females $=217$ naturalorigin and 478 hatchery-origin fish and for males $=151$ natural-origin and 213 hatchery-origin fish.

Methow/Chewuch Spring Chinook Salmon


Figure 12. Mean POH length (cm) and 95\% CI of hatchery- and natural-origin female and male spring Chinook Salmon spawners of different ages sampled on the spawning grounds in the Methow and Chewuch rivers for the combined years 1993-2018 (top row), before reduction (1993-2014; middle row), and after reduction (2015-2018; bottom row) in hatchery smolt production. Sample sizes are shown within each bar.

## Methow/Chewuch Spring Chinook Salmon



Figure 13. Relationships between fecundity (number of eggs, mean weight of eggs, and gonadal mass) and size (POH length and weight) for hatchery- (dashed line) and natural-origin (solid line) Methow/Chewuch spring Chinook Salmon for return years 2014-2018. Simple linear regression statistics are shown in each figure.

## Twisp River Program

Age-at-Maturity Analysis-For the complete time series (1993-2018), age at maturity differed significantly between hatchery- and natural-origin female (Yates' Chi-square $=64.722$, $\mathrm{P}=0.000$ ) and male spring Chinook Salmon (Yates' Chi-square $=45.211, \mathrm{P}=0.000$ ) in the Twisp River (Figure 14). Most females and males returned at age-4; however, a larger proportion of hatchery-origin fish returned at younger ages than did natural-origin fish. For example, $96 \%$ of hatchery-origin females and $73 \%$ of the natural-origin females returned at age- 4 . About $27 \%$ of natural-origin females and $4 \%$ of hatchery-origin females returned at age- 5 . A similar pattern was observed with males: about $38 \%$ of the hatchery-origin males and $13 \%$ of natural-origin males returned at age-3. In contrast, about $14 \%$ of natural-origin males and $2 \%$ of hatchery males returned at age-5.

There was little change in age at maturity before (1993-2014) and after (2014-2018) reduction in hatchery production numbers, although a larger proportion of hatchery- and naturalorigin males returned as age 4 fish following the reduction in program production (Figure 14). During the before period, age at maturity differed significantly between hatchery- and natural-
origin females (Yates' Chi-square $=68.527, \mathrm{P}=0.000$ ) and males (Yates' Chi-square $=40.255$, $\mathrm{P}=0.000$ ). During the after period, however, age-at maturity did not differ significantly between hatchery- and natural-origin females (Yates' Chi-square $=0.820, \mathrm{P}=0.365$ ) and males (Yates' Chi-square $=0.962, \mathrm{P}=0.327$ ) .

Size-at-Maturity Analysis-For the complete time series (1993-2018), size at maturity was significantly affected by sex and age (GLM ANOVA F $=13.922, \mathrm{P}=0.000$ ) but not by origin (GLM ANOVA $\mathrm{F}=1.424, \mathrm{P}=0.233$ ) (Figure 15). Older fish of both sexes were larger than younger fish and older males were generally larger than older females. Size differences between hatchery- and natural-origin fish within each age group were small, generally less than 1.5 cm on average.

Because of a lack of data, we could not evaluate statistically the effects of period, age, sex, and origin on size at maturity in the Twisp River. Differences in sizes among age classes were consistent during the before and after reduction periods and differences in sizes between hatchery- and natural-origin fish were small during both periods (Figure 15).

Fecundity Analysis-Except for the relationship between fork length and mean egg weight for natural-origin spring Chinook Salmon, all the other fecundity relationships for both hatchery- and natural-origin spring Chinook Salmon in the Twisp River were significant for the combined years 2014-2018 (Figure 16). Fecundity relationships between hatchery- and naturalorigin fish differed significantly for fork length and fecundity (ANCOVA F $=5.362, \mathrm{P}=0.025$ ) and fork length and mean egg weight (ANCOVA $\mathrm{F}=5.252, \mathrm{P}=0.027$ ). There were no differences in relationships between hatchery- and natural-origin fish for female weight and fecundity (ANCOVA F $=3.895, \mathrm{P}=0.055$ ) and fork length and total egg weight $($ ANCOVA $\mathrm{F}=$ $0.001, \mathrm{P}=0.982$ ).

## Twisp River Spring Chinook Salmon



Figure 14. Proportion of hatchery- and natural-origin female and male spring Chinook Salmon spawners of different ages sampled on the spawning grounds in the Twisp River for the combined years 1993-2018 (top row), before reduction (1993-2014; middle row), and after reduction (2015-2018; bottom row) in hatchery smolt production. Sample sizes for females $=314$ natural-origin and 311 hatchery-origin fish and for males $=264$ natural-origin and 229 hatcheryorigin fish for the combined analysis. For the before analysis, sample sizes for females $=253$ natural-origin and 268 hatchery-origin fish and for males $=221$ natural-origin and 205 hatcheryorigin fish; and for the after analysis, sample sizes for females $=61$ natural-origin and 43 hatchery-origin fish and for males $=43$ natural-origin and 24 hatchery-origin fish.

## Twisp River Spring Chinook Salmon



Figure 15. Mean POH length (cm) and 95\% CI of hatchery- and natural-origin female and male spring Chinook Salmon spawners of different ages sampled on the spawning grounds in the Twisp River for the combined years 1993-2018 (top row), before reduction (1993-2014; middle row), and after reduction (2015-2018; bottom row) in hatchery smolt production. Sample sizes are shown within each bar.

## Twisp River Spring Chinook Salmon



Figure 16. Relationships between fecundity (number of eggs, mean weight of eggs, and gonadal mass) and size (POH length and weight) for hatchery- (dashed line) and natural-origin (solid line) Twisp River spring Chinook Salmon for return years 2014-2018. Simple linear regression statistics are shown in each figure.

## Summer Chinook Salmon

## Wenatchee River Program

Age-at-Maturity Analysis-For the complete time series (1993-2018), age at maturity differed significantly between hatchery- and natural-origin female (Yates' Chi-square $=$ $1,050.133, \mathrm{P}=0.000$ ) and male summer Chinook Salmon (Yates' Chi-square $=976.779, \mathrm{P}=$ 0.000 ) in the Wenatchee River (Figure 17). Most female and male summer Chinook Salmon returned at ocean-age-3; however, a larger proportion of hatchery-origin fish returned at younger ocean ages than did natural-origin fish. Specifically, a higher proportion of hatchery-origin females returned as ocean-age-2 and 3 fish than did natural-origin females. In contrast, a higher proportion of natural-origin females returned as ocean-age-4 fish than did hatchery-origin females. Similarly, a higher proportion of hatchery-origin males returned as ocean-age-1 and 2 fish than did natural-origin males, while a higher proportion of natural-origin males returned as ocean-age- 3 and 4 fish than did hatchery-origin males.

There was a change in age at maturity before (1993-2014) and after (2014-2018) reduction in hatchery production numbers, although comparisons are affected by the small number of hatchery-origin fish sampled during the after period (Figure 17). During the before period, age at maturity differed significantly between hatchery and natural-origin female (Yates' Chi-square $=889.751, \mathrm{P}=0.000$ ) and male (Yates' Chi-square $=889.751, \mathrm{P}=0.000$ ) summer Chinook Salmon. During the after period, hatchery-origin females returned at a younger ocean age compared to the before period (Figure 17). There was less of a change in the age at maturity of natural-origin females between the before and after periods. In addition, there was little change in the age at maturity for hatchery- and natural-origin males between the before to after periods. Because of small sample sizes, statistical analyses could not be conducted on data collected during the after period.

Size-at-Maturity Analysis-For the complete time series (1993-2018), size at maturity differed significantly between hatchery- and natural-origin summer Chinook Salmon in the Wenatchee River (GLM ANOVA F $=4.346, \mathrm{P}=0.037$ ) (Figure 18). Size at maturity was significantly affected by age, origin, and sex (GLM ANOVA F $=4.225, \mathrm{P}=0.002$ ). For females, ocean-age-2 and 5 natural-origin fish were larger on average than ocean-age-2 and 5 hatcheryorigin fish, respectively (mean differences $=4$ and 3 cm , respectively). For males, ocean-age-1, 2 , and 5 natural-origin fish were larger on average than ocean-age-1, 2 , and 5 hatchery-origin fish, respectively (mean differences $=2,2$, and 4 cm , respectively). For both sexes, samples sizes are low for ocean-age-5 fish.

Because of a lack of data, we could not evaluate statistically the effects of period, age, sex, and origin on size at maturity of summer Chinook Salmon in the Wenatchee River. Differences in sizes among ocean-age classes were consistent during the before and after reduction periods and differences in sizes between hatchery- and natural-origin fish were relatively small during both periods (Figure 18). A notable difference was observed with size at maturity for ocean-age-2 females. During the before period, ocean-age-2 natural-origin females were on average 4 cm larger than ocean-age- 2 hatchery-origin females. During the after period, ocean-age- 2 natural-origin females were on average 4 cm smaller than ocean-age-2 hatcheryorigin females.

Fecundity Analysis-Because only natural-origin summer Chinook Salmon are collected for broodstock for the Wenatchee program, there are no comparisons of fecundity relationships between hatchery- and natural-origin summer Chinook Salmon in the Wenatchee River. Based on examination of natural-origin summer Chinook Salmon, there were significant linear relationships between female size and fecundity, mean egg weight, and gonadal mass for the combined years 2014-2018 (Figure 19).

## Wenatchee River Summer Chinook Salmon



Figure 17. Proportion of hatchery- and natural-origin female and male summer Chinook Salmon spawners of different ocean (salt) ages sampled on the spawning grounds in the Wenatchee River basin for the combined years 1993-2018 (top row), before reduction (1993-2014; middle row), and after reduction (2015-2018; bottom row) in hatchery smolt production. Sample sizes for females $=15,026$ natural-origin and 3,725 hatchery-origin fish and for males $=11,960$ naturalorigin and 2,664 hatchery-origin fish for the combined analysis. For the before analysis, sample sizes for females $=13,525$ natural-origin and 3,529 hatchery-origin fish and for males $=11,060$ natural-origin and 2,582 hatchery-origin fish; and for the after analysis, sample sizes for females $=1,501$ natural-origin and 196 hatchery-origin fish and for males $=900$ natural-origin and 82 hatchery-origin fish.

## Wenatchee River Summer Chinook Salmon



Figure 18. Mean POH length (cm) and 95\% CI of hatchery- and natural-origin female and male summer Chinook Salmon spawners of different ocean (salt) ages sampled on the spawning grounds in the Wenatchee River basin for the combined years 1993-2018 (top row), before reduction (1993-2014; middle row), and after reduction (2015-2018; bottom row) in hatchery smolt production. Sample sizes are shown within each bar.

## Wenatchee River Summer Chinook Salmon



Figure 19. Relationships between fecundity (number of eggs, mean weight of eggs, and gonadal mass) and size ( POH length and weight) for natural-origin Wenatchee River summer Chinook Salmon for return years 2014-2018. No hatchery-origin fish are collected for the Wenatchee Summer Chinook Salmon program. Simple linear regression statistics are shown in each figure.

## Methow River Program

Age-at-Maturity Analysis-For the complete time series (1993-2018), age at maturity differed significantly between hatchery- and natural-origin female (Yates' Chi-square $=805.448$, $\mathrm{P}=0.000$ ) and male summer Chinook Salmon (Yates' Chi-square $=897.500, \mathrm{P}=0.000$ ) in the Methow River (Figure 20). Most females returned at ocean-age-3; however, a larger proportion of hatchery-origin fish returned at younger ocean ages than did natural-origin fish. Specifically, a higher proportion of natural-origin females returned as ocean-age-4 fish than did hatchery-origin females. A higher proportion of hatchery-origin males returned as ocean-age-2 fish, while a higher proportion of natural-origin males returned as ocean-age- 3 fish. As with female summer Chinook, hatchery-origin males returned at a younger ocean age than did natural-origin males.

There was a change in age at maturity before (1993-2014) and after (2014-2018) reduction in hatchery production numbers (Figure 20). During both the before and after periods, age at maturity differed significantly between hatchery and natural-origin female (Before: Yates' Chi-square $=578.136, \mathrm{P}=0.000$; After: Yates' Chi-square $=222.104, \mathrm{P}=0.000$ ) and male
(Before: Yates' Chi-square $=708.532, \mathrm{P}=0.000$; After: Yates' Chi-square $=177.916, \mathrm{P}=0.000$ ) summer Chinook Salmon. Compared to the before period, nearly equal proportions of naturalorigin females returned as ocean-age- 3 and 4 fish during the after period, while there was little change in age at maturity for hatchery-origin females between the before and after periods. A larger proportion of hatchery-origin males returned as ocean-age- 2 fish during the after period, while a larger proportion of natural-origin males returned as ocean-age-4 fish during the after period compared to the before period.

Size-at-Maturity Analysis-For the complete time series (1993-2018), size at maturity differed significantly between hatchery- and natural-origin summer Chinook Salmon in the Methow River (GLM ANOVA F = 6.796, $\mathrm{P}=0.009$ ) (Figure 21). Size at maturity was significantly affected by age, origin, and sex (GLM ANOVA F = 10.503, $\mathrm{P}=0.000$ ). For females, ocean-age-2 natural-origin fish were larger on average than ocean-age-2 hatchery-origin fish (mean differences $=3 \mathrm{~cm}$ ). For males, ocean-age-0 and 1 natural-origin fish were larger on average than ocean-age-0, and 1 hatchery-origin fish, respectively (mean differences $=6$ and 3 cm , respectively). Sample sizes for ocean-age-5 fish for both sexes are too small to evaluate origin differences adequately.

There were significant differences in the size at maturity before and after reduction in hatchery production numbers ( GLM ANOVA $\mathrm{F}=123.000, \mathrm{P}=0.000$ ) (Figure 21). For most age groups, both hatchery- and natural-origin fish were smaller following the reduction in hatchery production, with the change in size ranging from $2-5 \mathrm{~cm}$ on average. During both time periods for both hatchery- and natural-origin fish, older males were larger on average than females of the same age (GLM ANOVA F $=4.363, \mathrm{P}=0.013$ ).

Fecundity Analysis-Because only natural-origin summer Chinook Salmon are collected for broodstock for the Methow program, there are no comparisons of fecundity relationships between hatchery and natural-origin summer Chinook Salmon in the Methow River. Based on examination of natural-origin fish, there were significant linear relationships between female size and fecundity, mean egg weight, and gonadal mass for the combined years 2014-2018 (Figure 22).

In contrast, both hatchery and natural-origin summer Chinook Salmon fecundity data are available for the Wells summer Chinook Salmon program, although sample sizes of naturalorigin fish are small. Analyses of these fish indicated significant linear relationships between female size and fecundity, mean egg weight, and gonadal mass for both hatchery and naturalorigin summer Chinook Salmon for the combined years 2014-2018 (Figure 23). There were no differences in relationships between hatchery and natural-origin fish for fork length and fecundity (ANCOVA $\mathrm{F}=0.725, \mathrm{P}=0.395$ ), female weight and fecundity ( $\mathrm{ANCOVA} \mathrm{F}=0.617$, $\mathrm{P}=0.433$ ), fork length and mean egg weight (ANCOVA $\mathrm{F}=0.308, \mathrm{P}=0.579$ ), and fork length and gonadal mass (ANCOVA F $=0.006, \mathrm{P}=0.940$ ).

## Methow River Summer Chinook Salmon



Figure 20. Proportion of hatchery- and natural-origin female and male summer Chinook Salmon spawners of different ocean (salt) ages sampled on the spawning grounds in the Methow River basin for the combined years 1993-2018 (top row), before reduction (1993-2014; middle row), and after reduction (2015-2018; bottom row) in hatchery smolt production. Sample sizes for females $=3,935$ natural-origin and 2,774 hatchery-origin fish and for males $=3,927$ naturalorigin and 2,537 hatchery-origin fish for the combined analysis. For the before analysis, sample sizes for females $=3,186$ natural-origin and 2,489 hatchery-origin fish and for males $=3,310$ natural-origin and 2,262 hatchery-origin fish; and for the after analysis, sample sizes for females $=749$ natural-origin and 285 hatchery-origin fish and for males $=617$ natural-origin and 275 hatchery-origin fish.

## Methow River Summer Chinook Salmon



Figure 21. Mean POH length (cm) and 95\% CI of hatchery- and natural-origin female and male summer Chinook Salmon spawners of different ocean (salt) ages sampled on the spawning grounds in the Methow River basin for the combined years 1993-2018 (top row), before reduction (1993-2014; middle row), and after reduction (2015-2018; bottom row) in hatchery smolt production. Sample sizes are shown within each bar.

## Methow River Summer Chinook Salmon



Figure 22. Relationships between fecundity (number of eggs, mean weight of eggs, and gonadal mass) and size ( POH length and weight) for natural-origin Methow River summer Chinook Salmon for return years 2014-2018. No hatchery-origin fish are collected for the Methow Summer Chinook Salmon program. Simple linear regression statistics are shown in each figure.

## Wells Summer Chinook Salmon



Figure 23. Relationships between fecundity (number of eggs, mean weight of eggs, and gonadal mass) and size ( POH length and weight) for hatchery- (dashed line) and natural-origin (solid line) Wells summer Chinook Salmon for return years 2014-2018. Simple linear regression statistics are shown in each figure.

## Chelan River Program

Age-at-Maturity Analysis-For the complete time series (2000-2018), age at maturity differed significantly between hatchery- and natural-origin female (Yates' Chi-square $=432.131$, $\mathrm{P}=0.000$ ) and male summer Chinook Salmon (Yates' Chi-square $=98.590, \mathrm{P}=0.000$ ) in the Chelan River (Figure 24). A larger proportion of hatchery-origin fish returned at younger ocean ages than did natural-origin fish. Most hatchery-origin females returned at ocean-age-3, and nearly all natural-origin females returned at ocean-ages 3 or 4 , in nearly equal proportions. A higher proportion of hatchery-origin males returned as ocean-age- 1 and 2 fish, while a higher proportion of natural-origin males returned as ocean-age-3 and 4 fish.

Because hatchery-origin summer Chinook Salmon from several different hatchery programs have spawned in the Chelan River since before the 1990s, there is no comparison of before and after periods for Chelan River summer Chinook Salmon.

Size-at-Maturity Analysis-For the complete time series (2000-2018), size at maturity differed significantly between hatchery and natural-origin summer Chinook Salmon in the

Chelan River (GLM ANOVA F $=55.285, \mathrm{P}=0.000$ ) (Figure 25). Size at maturity was significantly affected by age, origin, and sex (GLM ANOVA F $=4.055$, $\mathrm{P}=0.017$ ). For both females and males, hatchery-origin fish were generally larger than natural-origin fish across most age groups. Those differences in sizes were on average $2-5 \mathrm{~cm}$ depending on age group.

Because hatchery-origin summer Chinook Salmon from several different hatchery programs have spawned in the Chelan River since before the 1990s, there is no comparison of before and after periods for Chelan River summer Chinook Salmon.

Fecundity Analysis - Because only hatchery-origin summer Chinook Salmon are collected for broodstock for the Chelan River summer Chinook Salmon program, there are no comparisons of fecundity relationships between hatchery- and natural-origin summer Chinook Salmon in the Chelan River. Based on examination of hatchery-origin summer Chinook Salmon, there were significant linear relationships between female size and fecundity, mean egg weight, and gonadal mass for the combined years 2014-2018 (Figure 26).

## Chelan River Summer Chinook Salmon



Figure 24. Proportion of hatchery- and natural-origin female and male summer Chinook Salmon spawners of different ocean (salt) ages sampled on the spawning grounds in the Chelan River for the combined years 2000-2018. Sample sizes for females $=891$ natural-origin and 1,262 hatchery-origin fish and for males $=273$ natural-origin and 444 hatchery-origin fish for the combined analysis.

## Chelan River Summer Chinook Salmon



Figure 25. Mean POH length (cm) and 95\% CI of hatchery- and natural-origin female and male summer Chinook Salmon spawners of different ocean (salt) ages sampled on the spawning grounds in the Chelan River for the combined years 2000-2018. Sample sizes are shown within each bar.

## Chelan River Summer Chinook Salmon



Figure 26. Relationships between fecundity (number of eggs, mean weight of eggs, and gonadal mass) and size ( POH length and weight) for hatchery-origin Chelan River summer Chinook Salmon for return years 2014-2018. No natural-origin fish are collected for the Chelan Summer Chinook Salmon program. Simple linear regression statistics are shown in each figure.

## Discussion

We observed similar patterns for most hatchery- and natural-origin adult Spring Chinook and Summer Chinook Salmon for all hatchery programs and these patterns were generally consistent with published studies from other locations inside and outside the Columbia River Basin (Knudsen et al. 2006; 2008). As such, similar mechanisms are likely influencing hatchery programs at a broad scale. For age at maturity, hatchery-origin fish universally returned at a younger age than natural-origin fish, and with a few exceptions, there were no changes in programs with a before and after period. While the difference in age structure between hatcheryand natural-origin populations were statistically significant, in all cases the predominate age at maturity was the same. For example, in Spring Chinook the large majority of male and female hatchery- and natural-origin fish returned at age-4. For Summer Chinook, the majority of hatchery- and natural-origin females returned at salt age-3, with the majority of males distributed between salt age-2 and age-3. All of the Spring Chinook hatchery programs produced few age- 5 returns relative to the proportions of natural-origin age- 5 returns in the respective populations
and likewise for the age- 4 returns in the summer Chinook programs. A notable change from the before to after periods was the reduction in age-3 fish returning to Nason Creek. All hatcheryorigin fish in Nason Creek in the before period were strays. With the reduction in release number in the after period from nearby hatchery programs, there were likely fewer strays, which are often age- 3 fish, returning to Nason Creek.

The size at maturity of returning adult fish was significantly affected by age and sex and in some cases origin and period. However, when matched by age and sex, the differences in size by origin and period that were statistically significant were minor and likely of little biological relevance. For example, the largest difference of all the age and sex matched comparisons was for age- 3 females in the Chiwawa River, where hatchery-origin fish were on average 9 cm larger than natural-origin fish, and where the sample size for natural-origin fish was $n=3$. In most cases, the differences in size by origin were on the order of 2-3 cm for significant and nonsignificant tests.

For all populations the fecundity metrics were significantly affected by fish size and weight. In the Chiwawa River and Twisp River, differences in fecundity between hatchery- and natural-origin fish were significant for length vs. fecundity and egg weight. A visual examination of the relationships suggests that in the Chiwawa River the differences between the hatchery- and natural-origin fork length-fecundity relationship was driven by a larger number of smaller hatchery-origin females with low fecundity. The difference in the length to egg-weight relationship, however, was explained by higher egg weight of smaller hatchery-origin females. In the Twisp, conclusions were more tenuous due to the small sample size (e.g., $\mathrm{n}=10$ ). As with size at maturity, the few differences in the fecundity metrics between hatchery- and naturalorigin fish were small and may have little biological relevance.

Generally, and beyond the Upper Columbia Basin, hatchery-origin fish return at an early age (Tillotson et al. 2021). This is likely explained by the relatively large size of smolts at release, which leads to earlier maturation and thus an early return to fresh water (Vøllestad et al. 2004). Managers of hatchery programs in the upper Columbia River and in most other areas have chosen to produce bigger smolts than their natural-origin counterparts in order to achieve high post-release survival of these fish (Feldhaus et al. 2016). Improvements in hatchery management techniques have been implemented at the hatchery programs analyzed here to reduce early maturation, specifically aimed at reducing mini-jacks (Harstad et al. 2018). These efforts will likely have cascading effects through the age classes and future evaluations may show improvements to the overall age-at-return structure. However, trade-offs associated with postrelease survival and age at maturation will occur. One of the main consequences of deviating from the natural-origin fish growth and size template is the deviation of some phenotypic characteristics of returning fish. Larger smolts typically have higher post-release survival (Gosselin et al. 2019; Vøllestad et al. 2004; but see Feldhaus et al. 2016) but also mature at younger ages and smaller sizes, which are traits associated with lower fecundity (Claiborne et al. 2011; Tattam et al. 2015). Yet, hatchery fish typically have higher adult-to-adult survival rates than natural-origin fish (Pearsons et al., Chapter 1 of this report), which on a broodyear-basis could result in a larger hatchery-origin egg escapement despite lower fecundities on a per-fish basis.

Managers are in the position of considering the many complicated, and often mutually exclusive trade-offs between hatchery-origin fish survival and matching the phenotypic traits of natural-origin fish (Larsen et al. 2019). For example, they must consider the potential influences of parental effects in the broodstock on age and size at return (Church et al. 2021; Forest et al.
2017), phenotypic and phenologic consequences of synergistic interactions between rearing practices and parental phenotype (Winsor et al. 2021), and local adaptations potentially mediating age and size at return (McKinney et al. 2020). Additionally, managers must also weigh the risks to population fitness posed by program objectives (Waters et al. 2018).

Future analyses of these metrics should consider changes over time and consider trends between hatchery-origin and natural-origin fish. Declines in age at maturity and size at maturity are a Pacific-wide concern (Oke et al. 2020; Ohlberger et al. 2018) and therefore a temporal analysis of the dataset generated for the monitoring of these hatchery programs would benefit the region. Finally, data used in these analyses did not accurately account for the contribution of minijacks to cohort age structure, when minijacks have been found to comprise a substantial component of total "returns" from local programs, influencing apparent HRRs and SARs (Harstad et al. 2018).

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# Numbers and Sizes of Fish Released from Hatcheries in the Upper Columbia Basin 

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[^6]
#### Abstract

Chelan, Douglas, and Grant PUDs produce salmon and steelhead for mitigation under various agreements including the Rock Island Habitat Conservation Plan (HCP), the Rocky Reach HCP, the Wells HCP, and the Priest Rapids Settlement Agreement and associated Biological Opinion. The various hatchery programs are monitored for the number of fish released, as well as metrics for fish size at release and condition. The programs include Summer Chinook Salmon, Spring Chinook Salmon, and summer steelhead produced and released at a variety of facilitates in the Wenatchee River Subbasin, Methow River Subbasin, and Columbia River. Programs generally met or exceeded the release-number targets. Specific programs did not meet the release-number targets in all years, but all programs met or exceeded targets in the majority of years, with the exception of the White River spring Chinook Salmon captive broodstock program. This program was in development for all of its history and tested numerous fish culture and release strategies. Therefore, the program was not able to meet targets that remained consistent across years. Most programs met or were close to meeting the CV target on average. Meeting the CV targets for steelhead programs was more difficult than for the Chinook Salmon programs. Meeting fish-per-pound (fpp) targets was more difficult in Spring Chinook Salmon and conservation steelhead programs, potentially because these programs use naturalorigin broodstock. Spring Chinook Salmon exhibited near-isometric growth. Steelhead exhibited negative allometric growth, as did Wells and Methow summer Chinook Salmon. However, Wenatchee and Chelan Falls summer Chinook Salmon exhibited isometric or positive allometric growth. Condition factors across all programs were very close to or exceeded 1. Trade-offs between post-release survival and age at maturation influence hatchery rearing strategies. Large Chinook Salmon generally survive better than small fish, but they also are more prone to mature at younger ages. Hatchery programs that use natural-origin fish for broodstock may experience higher levels of precocial maturation than those that use hatcheryorigin fish for broodstock. In addition, larger fish also pose ecological risks to other fish through mechanisms such as competition or predation. In general, the length targets, and some of the metrics that used a single value (e.g., length and weight) associated with a range of fpp targets, were not useful. Adaptation of targets may occur to achieve a better suite of benefits among the many trade-offs involved in growing fish to a target size or weight.


## Introduction

Chelan, Douglas, and Grant PUDs produce salmon and steelhead for mitigation under various agreements including the Rock Island Habitat Conservation Plan (HCP), the Rocky Reach HCP, the Wells HCP, and the Priest Rapids Settlement Agreement. The targets for number of fish to be produced and fish per pound were established in the Habitat Conservation Plans, Hatchery and Genetic Management plans, hatchery recalculation implementation plans, Hatchery Monitoring and Evaluation Plan, or were subsequently adjusted by the HCP Hatchery Committees or Priest Rapids Hatchery Sub-Committee through formal processes. The various hatchery programs are monitored for the number of fish released, as well as metrics for fish size at release and condition. The programs include Summer Chinook Salmon, Spring Chinook Salmon, and summer steelhead, produced and released at a variety of facilitates in the Wenatchee River Subbasin, Methow River Subbasin, and Columbia River.

## Methods

Study Area
The study area encompasses the Wenatchee River Subbasin, the Methow River Subbasin, and the Columbia River. The hatchery programs, fish production facilities, and release locations are provided in Table 1.

Table 1. Fish production facilities and release locations.

| Program | Type | Facility | Release Location |
| :--- | :--- | :--- | :--- |
| Chiwawa River Spring <br> Chinook | Conservation | Eastbank Hatchery and Chiwawa <br> Acclimation Facility | Chiwawa River |
| Nason Creek Spring <br> Chinook | Conservation and <br> Safety-Net | Eastbank Hatchery and Nason <br> Creek Acclimation Facility | Nason Creek |
| White River Spring <br> Chinook | Conservation | Aquaseed; Little White Salmon <br> National Fish Hatchery; various <br> acclimation sites | White River; Lake <br> Wenatchee; <br> Wenatchee River |
| Methow River Spring <br> Chinook | Conservation | Methow Hatchery and Goat Wall <br> Acclimation Pond | Methow River |
| Chewuch River Spring <br> Chinook | Conservation | Methow Hatchery and Chewuch <br> Acclimation Pond | Chewuch River |
| Twisp River Spring <br> Chinook | Conservation | Methow Hatchery and Twisp <br> Acclimation Pond | Twisp River |
| Wenatchee River <br> Summer Steelhead | Conservation and <br> Safety-Net | Eastbank Hatchery | Wenatchee River |
| Twisp River Summer <br> Steelhead | Conservation | Winthrop National Fish Hatchery <br> and Wells Hatchery | Twisp River, Methow <br> River |
| Methow River <br> Summer Steelhead | Safety-Net | Wells Hatchery | Methow River |
| Columbia River <br> Summer Steelhead | Safety-Net | Wells Hatchery | Columbia River |
| Wenatchee River <br> Summer Chinook | Conservation/Harvest | Eastbank Hatchery and Dryden <br> Acclimation Facility | Wenatchee River |
| Chelan Falls Summer <br> Chinook | Harvest | Chelan Falls Hatchery | Chelan River |
| Wells Yearling <br> Summer Chinook | Harvest | Wells Hatchery | Columbia River |
| Wells Subyearling <br> Summer Chinook | Harvest | Conservation/Harvest | Eastbank Hatchery and Carlton <br> Acclimation Facility |
| Methow River <br> Summer Chinook | Methow River |  |  |

## Hatchery Production Targets

The targets for number of fish to be produced and fish per pound were established in the Habitat Conservation Plans and Hatchery and Genetic Management plans, hatchery recalculation implementation plans, Hatchery Monitoring and Evaluation Plan, or formally adjusted by the

Hatchery Committees or Priest Rapids Coordination Committee's Hatchery Sub-Committee. The Committees also adopted targets for coefficient of variation. Fish weight is described by the fish-per-pound (fpp) targets. The Committees adopted the concept of using Condition Factor targets, but such targets have not yet been identified. Targets for mean fish length have not been formally adopted in the Monitoring and Evaluation (M\&E) Plan but are reported in the M\&E Annual Reports. Previous fish length and weight targets obtained from Piper (1952) were not appropriate based on empirical data for Upper Columbia populations and were abandoned. Length targets in the M\&E Reports are either the same as, or similar to, the Piper targets and are not compatible with the fpp targets. Meeting these fpp targets is typically mutually exclusive of meeting the length target. All the rearing targets and metrics are presented in Table 2. Metrics that do not have quantitative targets are presented in summary statistics.

Table 2. Hatchery program production and fish size targets.

| Program | Production <br> Target | Length Target <br> $(\mathbf{m m})$ | CV | Weight <br> $(\mathbf{g})$ | Fish/Pound |
| :--- | :---: | :---: | :---: | :---: | :---: |
| Chiwawa River Spring <br> Chinook | 144,026 | 155 | 9 | 37.8 | 18 |
| Nason Creek Spring Chinook | 223,670 | 155 | 9 | 37.8 | $18-24$ |
| White River Spring Chinook | 150,000 | NA | 9 | NA | $18-24$ |
| Methow River Spring Chinook | 133,249 | 137 | 9 | 30.2 | 15 |
| Chewuch River Spring <br> Chinook | 60,516 | 136 | 9 | 30.2 | 15 |
| Twisp River Spring Chinook | 30,000 | 135 | 9 | 30.2 | 15 |
| Wenatchee River Summer <br> Steelhead | 247,300 | 191 | 9 | 75.6 | 6 |
| Twisp River Summer <br> Steelhead | 48,000 | 191 | 9 | 75.6 | 6 |
| Methow River Summer <br> Steelhead | 100,000 | 191 | 9 | 75.6 | 6 |
| Columbia River Summer <br> Steelhead | 160,000 | 191 | 9 | 75.6 | 6 |
| Wenatchee River Summer <br> Chinook | 500,001 | 163 | 9 | $30.0-45.4$ | $10-18$ |
| Chelan River Summer <br> Chinook | 576,000 | 161 | 9 | $20.0-45.4$ | $10-22$ |
| Wells Yearling Summer <br> Chinook | 320,000 | 168 | 7 | 45.4 | 10 |
| Wells Subyearling Summer <br> Chinook | 484,000 | NA | 7 | 9.1 | 50 |
| Methow River Summer <br> Chinook | 200,000 | 163 | 9 | 45.4 | $13-18$ |

## Data Collection and Derived Metrics

For each program, fish were enumerated during coded wire tagging and/or marking. Because marking and tagging often occurred weeks to several months before fish were released, hatchery managers kept track of fish that died before release and subtracted those fish from the number marked. Thus, the total number of fish released from a specific program was the number of fish marked minus the number of marked fish that died before release (and any live fish retained for early maturation studies or other studies).

To estimate fish size, a sample of about 200 fish was randomly collected from each stock using cast nets or dip nets when fish were crowded. Each fish was measured for fork length $(\mathrm{mm})$ and weighed to the nearest 0.1 g . Arithmetic means were calculated for both length (L) and weight (W), and weight was also converted to fish per pound (fpp). The coefficient of variation $(\mathrm{CV})$ for length was calculated as the ratio of the sample standard deviation to the sample mean. Because this estimate is generally biased low, the ratio was corrected by multiplying it by ( $1+$ $1 / 4 n$ ), where $n=$ sample size. For each fish we also calculated Fulton Condition Factors using the equation, $\mathrm{K}_{\mathrm{FL}}=\left(\mathrm{W} / \mathrm{L}^{3}\right) \times 100,000$, where $\mathrm{K}_{\mathrm{FL}}$ is the condition factor based on fork length (Neumann et al. 2012).

## Data Analysis

No statistical analyses were needed to evaluate numbers and sizes (lengths and weights) of fish released from each hatchery program. Here, we simply compared the numbers and sizes of fish released to established management targets. In contrast, we used simple linear regression to evaluate the relationship between fish length and weight for each program. We used common logarithms (log with base 10) to transform both length and weight data for analysis. We examined the $b$ parameter (slope of the length-weight relationship) to determine the allometric growth of fish (Neumann et al. 2012). Values of $b$ less than 3 indicated that fish body form became slimmer with increasing length (negative allometric growth), values greater than 3 indicated that fish body form became deeper and/or wider with increasing length (positive allometric growth), and values equal to 3 indicated isometric growth.

## Results

Wenatchee River Conservation and Safety-Net Summer Steelhead
Number Released-The recent goal of the supplementation program is to release 247,300 $( \pm 10 \%)$ juvenile Steelhead into the Wenatchee River Subbasin annually. During the six-year period (2012-2017) under the recent goal of the program, the program achieved that goal for five brood years (Figure 1). Numbers released ranged from 195,344-264,758 (average $=241,351$ ) juveniles.

Wenatchee Steelhead


Figure 1. Number of juvenile Steelhead released in the Wenatchee River Subbasin for brood years 1998-2017. The dashed horizontal line represents the target release number (400,000 from 1998-2011 and 247,300 from 2012-2017).

Size at Release-The current goal of the supplementation program (since brood year 2012) is to release juvenile Steelhead in the Wenatchee River Subbasin that average 191-mm long (fork length) with a CV of 9.0, and 75.6 g ( 6 fish/pound). During the six-year period under the recent goal of the program, the length target has not been reached and the CV target was exceeded in all those years (Figure 2). Mean lengths of fish released ranged from 127-180 mm (average $=154 \mathrm{~mm}$ ), while CVs ranged from 9-22 (average = 16). In addition, the mean weight target and the fish per pound target was not achieved during the recent six-year period a (Figure 2). Throughout the recent six years, mean weights ranged from $27-71 \mathrm{~g}$ (average $=45 \mathrm{~g}$ ) and fish per pound ranged from 6-17 (average $=11$ fish/pound).

## Wenatchee Steelhead Hatchery Releases



Figure 2. Average fork length (mm), coefficient of variation (CV) of fork length, average weight (g), and number of fish per pound of juvenile Steelhead released in the Wenatchee River Subbasin for brood years 1998-2017. The dashed horizontal lines represent the target length (198 mm from 1998-2011 and 191 mm from 2012-2017), length CV (9), weight ( 75.6 g ), and fish per pound ( 6 fish per pound). $\mathrm{W}=$ wild (natural-origin fish), $\mathrm{H}=$ hatchery-origin fish, $\mathrm{R}=$ raceway, and $\mathrm{C}=$ recirculating aquatic system.

Length-Weight Relationship-There was a significant relationship between the length and weight of juvenile hatchery Steelhead (Figure 3). Length explained $97 \%$ of the variation in weight of juvenile fish. In addition, the relationship indicated a mean condition 2.83 for juvenile hatchery steelhead, suggesting negative allometric growth. The Fulton condition factor averaged 1.12 (range, 0.36-3.55).


Figure 3. Relationship between fork length (mm) and weight (g) of juvenile Wenatchee River Steelhead sampled before release during 2003-2018.

## Methow River Safety-Net Summer Steelhead

Number Released-The recent goal of the supplementation program is to release 100,000 ( $\pm 10 \%$ ) juvenile Steelhead into the Methow River annually. During the six-year period under the recent goal of the program, the program achieved that goal in five brood years, and did not achieve it in one brood year (Figure 4). Numbers released ranged from 72,768-106,716 (average $=96,823$ ) juveniles.

## Methow Steelhead



Figure 4. Number of juvenile steelhead released in the Methow River for brood years 1992-2017. The dashed horizontal line represents the target release number (320,000 from 1992-2011 and 100,000 from 2012-2017).

Size at Release-The current goal of the safety-net program (since brood year 2012) is to release juvenile Steelhead in the Methow River that average 191-mm long (fork length) with a CV of 9.0, and 75.6 g ( 6 fish/pound). During the six-year period under the recent goal of the program, the length target has been reached in nearly all years (Figure 5). The CV target, however, was exceeded in all those years. Mean lengths of fish released ranged from 181-202 mm (average $=190 \mathrm{~mm}$ ), while CVs ranged from 10-14 (average $=12$ ). In addition, the mean weight and fish-per-pound targets were achieved in most years (Figure 5). Throughout the recent six years, mean weights ranged from $61-81 \mathrm{~g}($ average $=73 \mathrm{~g})$ and fish per pound ranged from 68 (average $=6$ fish/pound).

## Methow Steelhead Hatchery Releases



Figure 5. Average fork length (mm), coefficient of variation (CV) of fork length, average weight (g), and number of fish per pound of juvenile steelhead released in the Methow River for brood years 2011-2017. The dashed horizontal lines represent the target length ( 191 mm ), length CV (9), weight ( 75.6 g ), and fish per pound ( 6 fish per pound).

Length-Weight Relationship-There was a significant relationship between the length and weight of juvenile hatchery steelhead (Figure 6). Length explained 94\% of the variation in weight of juvenile fish. In addition, the relationship indicated a mean condition 2.94 for juvenile hatchery steelhead, suggesting negative allometric growth. The Fulton condition factor averaged 0.99 (range, 0.66-1.56).

## Methow Steelhead



Figure 6. Relationship between fork length (mm) and weight (g) of juvenile Methow River steelhead sampled before release during 2013-2018.

## Twisp River Conservation Summer Steelhead

Number Released-The recent goal of the conservation program is to release 48,000 ( $\pm 10 \%$ ) juvenile steelhead into the Twisp River annually. During the six-year period under the recent goal of the program, the program achieved that goal in four brood years and exceeded it in two brood years (Figure 7). Numbers released ranged from 50,787-59,226 (average $=54,280$ ) juveniles.

## Twisp Steelhead



Figure 7. Number of juvenile Steelhead released in the Twisp River for brood years 1997-2017. The dashed horizontal line represents the target release number (99,666 from 1997-2011 and 48,000 from 2012-2017).

Size at Release-The current goal of the supplementation program (since brood year 2012) is to release juvenile Steelhead in the Methow River that average 191-mm long (fork length) with a CV of 9.0 , and 75.6 g ( 6 fish/pound). During the six-year period under the recent goal of the program, lengths were generally below the target while CV exceeded the target (Figure 8). Mean lengths of fish released ranged from 155-182 mm (average $=167 \mathrm{~mm}$ ), while CVs ranged from 10-15 (average $=13$ ). In addition, the mean weight fell below the target and fish per pound exceeded the target in all years (Figure 8). Throughout the recent six years, mean weights ranged from 44-68 $\mathrm{g}($ average $=54 \mathrm{~g})$ and fish per pound ranged from 7-11 (average $=9$ fish/pound).

## Twisp Steelhead Hatchery Releases



Figure 8. Average fork length (mm), coefficient of variation (CV) of fork length, average weight (g), and number of fish per pound of juvenile steelhead released in the Twisp River for brood years 2011-2017. The dashed horizontal lines represent the target length ( 191 mm ), length CV (9), weight ( 75.6 g ), and fish per pound ( 6 fish per pound).

Length-Weight Relationship-There was a significant relationship between the length and weight of juvenile hatchery steelhead (Figure 9). Length explained 94\% of the variation in weight of juvenile fish. In addition, the relationship indicated a mean condition 2.94 for juvenile hatchery Steelhead, suggesting negative allometric growth. The Fulton condition factor averaged 1.12 (range, 0.77-1.74).

Twisp Steelhead


Figure 9. Relationship between fork length (mm) and weight (g) of juvenile Twisp River steelhead sampled before release during 2011-2018.

## Columbia River (Wells) Safety-Net Summer Steelhead

Number Released-The recent goal of the safety-net program is to release 160,000 $( \pm 10 \%)$ juvenile steelhead in the Columbia River from the Wells Hatchery annually. During the five-year period when the Columbia Safety-Net Program began implementation (2013-2017), the release goal of the program was not achieved in one brood year, was achieved in two brood years, and exceeded in two brood years (Figure 10). Numbers released (2011-2017) ranged from $55,541-210,328$ (average $=152,535$ ) juveniles.

## Wells Steelhead



Figure 10. Number of juvenile steelhead released from the Wells Hatchery to the Columbia River for brood years 2011-2017. Note that the current Columbia Safety-Net program $(160,000)$ began releases with brood year 2013. The dashed horizontal line represents the target release number (429,000 before 2012 and 160,000 from 2012-2017).

Size at Release-Information on size at release are presented under the Methow River steelhead section.

Length-Weight Relationship-Information on length-weight relationship is presented under the Methow River steelhead section.

## Chiwawa River Spring Chinook Salmon

Number Released-The recent goal of the supplementation program is to release 144,026 $( \pm 10 \%)$ juvenile spring Chinook Salmon into the Chiwawa River annually. During the five-year period under the recent goal of the program, the program achieved that goal for five brood years (Figure 11). Numbers released ranged from 144,360-163,411 (average $=152,661$ ) juveniles.

## Chiwawa Spring Chinook



Figure 11. Number of juvenile spring Chinook Salmon released in the Chiwawa River for brood years 1989-2017. The dashed horizontal line represents the target release number (672,000 from 1989-2010, 298,000 in 2011, 205,000 in 2012, and 144,026 from 2013-2017).

Size at Release-The current goal of the supplementation program (since brood year 2012) is to release juvenile spring Chinook Salmon in the Chiwawa River that average $155-\mathrm{mm}$ long (fork length) with a CV of 9.0, and 37.8 g ( 18 fish/pound). During the six-year period under the recent goal of the program, lengths were below the target while CV generally exceeded the target (Figure 12). Mean lengths of fish released ranged from 127-141 mm (average $=132 \mathrm{~mm}$ ), while CVs ranged from 7-16 (average $=10$ ). In addition, the mean weight fell below the target and fish per pound met or fell below the target during the six-year period (Figure 12). Throughout the recent six years, mean weights ranged from $25-35 \mathrm{~g}$ (average $=28 \mathrm{~g}$ ) and fish per pound ranged from 13-18 (average $=17$ fish/pound).

Chiwawa Spring Chinook Salmon Hatchery Releases


Figure 12. Average fork length (mm), coefficient of variation (CV) of fork length, average weight (g), and number of fish per pound of juvenile spring Chinook Salmon released in the Chiwawa River for brood years 1989-2017. The dashed horizontal lines represent the target length ( 176 mm from 1989-2011 and 155 mm from 2012-2017), length CV (9), weight ( 37.8 g ), and fish per pound ( 12 fish per pound from 1989-2011 and 18 fish per pound from 2012-2017).

Length-Weight Relationship-There was a significant relationship between the length and weight of juvenile hatchery spring Chinook Salmon (Figure 13). Length explained $95 \%$ of the variation in weight of juvenile fish. In addition, the relationship indicated a mean condition 3.01 for juvenile hatchery spring Chinook Salmon, suggesting near isometric growth. The Fulton condition factor averaged 0.99 (range, $0.66-1.56$ ).

## Chiwawa Spring Chinook



Figure 13. Relationship between fork length (mm) and weight (g) of juvenile Chiwawa River spring Chinook Salmon sampled before release during 2003-2018.

## Nason Creek Spring Chinook Salmon

Number Released-The goal of the supplementation program is to release 223,670 $( \pm 10 \%)$ juvenile spring Chinook Salmon into Nason Creek annually (combined conservation and safety-net programs). During the five-year period of the program, the program reached that goal in Nason Creek for three of those brood years or four of the five years if the fish released into the Chiwawa River as part of Grant PUDs production in 2016 were included (Figure 14). Numbers released into Nason Creek ranged from 32,215-243,127 (average $=156,751$ ) juveniles. For brood year 2014, most of the Nason Creek program was transferred to the Chiwawa Acclimation Facility and 196,866 of these fish were released in the Chiwawa River because of a water intake problem at the Nason Creek Acclimation Facility. The Nason Creek release was 32,215 in brood year 2014 but totaled 229,081 with the fish that were released into the Chiwawa River.

## Nason Creek Spring Chinook



Figure 14. Number of juvenile spring Chinook Salmon released in Nason Creek for brood years 2013-2017. The dashed horizontal line represents the target release number $(223,670)$. For brood year 2014, most of the Nason Creek program was transferred to the Chiwawa Acclimation Facility and 196,866 of these fish were released in the Chiwawa River because of a water intake problem at the Nason Creek Acclimation Facility. The total released for brood year 2014 was 229,081.

Size at Release-The goal of both the conservation (WxW) and safety net (HxH) programs is to release juvenile spring Chinook Salmon in Nason Creek that average $155-\mathrm{mm}$ long (fork length) with a CV of 9.0 , and 37.8 g (18-24 fish/pound). During the five-year period of the program, both lengths and CVs were generally below their respective targets (Figure 15). Mean lengths of fish released ranged from 119-129 mm (average $=122 \mathrm{~mm}$ ) for the conservation program and 115-134 mm (average $=122$ ) for the safety-net program. CVs ranged from 7-8 (average $=7$ ) for the conservation program and 6-13 (average $=9$ ) for the safety-net program. The mean weight for both programs fell below the target and fish per pound generally exceeded the target (Figure 15). Mean weights ranged from 21-28 $g$ (average $=23$ ) for the conservation program and $19-29 \mathrm{~g}$ (average $=22 \mathrm{~g}$ ) for the safety-net program. Fish per pound ranged from 16-22 (average $=20$ fish/pound) for the conservation program and 16-24 (average $=21$ fish/pound) for the safety-net program.

## Nason Creek Spring Chinook Salmon Hatchery Releases



Figure 15. Average fork length (mm), coefficient of variation (CV) of fork length, average weight (g), and number of fish per pound of juvenile spring Chinook Salmon released in Nason Creek for brood years 2013-2017. The dashed horizontal lines represent the target length (155 mm ), length CV (9), weight ( 37.8 g ), and fish per pound (18-24 fish per pound). WxW represents the conservation program while HxH represents the safety-net program.

Length-Weight Relationship-There was a significant relationship between the length and weight of juvenile hatchery spring Chinook Salmon (Figure 16). Length explained 95\% of the variation in weight of juvenile fish. In addition, the relationship indicated a mean condition 2.99 for juvenile hatchery spring Chinook Salmon, suggesting near isometric growth. The Fulton condition factor averaged 1.19 (range, 0.61-2.23).


Figure 16. Relationship between fork length (mm) and weight (g) of juvenile Nason Creek spring Chinook Salmon sampled before release during 2013-2018.

## White River Spring Chinook Salmon

Number Released-The goal of the supplementation program was to release 150,000 ( $\pm 10 \%$ ) juvenile spring Chinook Salmon into various release locations annually, including the White River, Lake Wenatchee, and the Wenatchee River. The release strategies were highly variable across years (Table 3). During the twelve-year period of the program, the program reached that goal for two of those brood years (Figure 17). Numbers released ranged from 1,639 to 281,677 (average $=78,716$ ) juveniles.

Table 3. White River Spring Chinook Salmon releases from Brood Years 2002-2013.

| Brood <br> Year | Release <br> Year | Number <br> Released | Acclimation <br> Site | Acclimation <br> Vessel | Release Scenario |
| :---: | :---: | :---: | :---: | :---: | :--- |
| 2002 | 2004 | 2,589 | WR RM 11.5 | Tanks | White River |
| 2003 | 2005 | 2,096 | WR RM 11.5 | Tanks | White River |
| 2004 | 2006 | 1,639 | WR RM 11.5 | Tanks | White River |
| 2005 | 2007 | 69,032 | Lake <br> Wenatchee | Net Pens | Lake Wenatchee |
| 2006 | 2008 | 139,644 | NA | NA | White River |
| 2006 | 2008 | 142,033 | NA | NA | White River |
| 2007 | 2009 | 87,671 | Lake <br> Wenatchee | Net Pens | Lake Wenatchee |
| 2007 | 2009 | 44,172 | None | None | Lake Wenatchee |
| 2008 | 2010 | 10,156 | WR Bridge | Eddy Pen | Escape |
| 2008 | 2010 | 38,400 | Lake <br> Wenatchee | Net Pens | Mouth of Lake |
| 2009 | 2011 | 12,000 | WR RM 11.5 | Side Channel | Escape |
| 2009 | 2011 | 10,000 | WR RM 11.5 | Tanks | White River |
| 2009 | 2011 | 28,000 | WR Bridge | Tanks | White River |
|  | 2009 | 2011 | 14,596 | WR Bridge | Eddy Pen |
| 2009 | 2011 | 48,000 | Lake <br> Wenatchee | Net Pens | Wenatchee River |
| 2010 | 2012 | 18,850 | WR Bridge | Tanks | Wenatchee River |
| 2011 | 2013 | 42,000 | WR Bridge | Tanks | Wenatchee and White <br> Rivers |
| 2011 | 2013 | 105,000 | Lake <br> Wenatchee | Net Pens | Wenatchee River |
| 2012 | 2014 | 42,000 | WR Bridge | Tanks | Wenatchee River |
| 2012 | 2014 | 55,713 | Lake <br> Wenatchee | Net Pens | Wenatchee River |
| 2013 | 2015 | 31,000 | WR Bridge | Tanks | Wenatchee River |

## White River Spring Chinook



Figure 17. Number of juvenile spring Chinook Salmon released in White River, Lake Wenatchee, and Wenatchee River for brood years 2002-2013. The dashed horizontal line represents the target release number $(150,000)$.

Size at Release-The goal of the captive broodstock program was to release juvenile spring Chinook Salmon that average 18-24 fish/pound (see Figure 18). Mean lengths of fish released ranged from 125-207 mm (average $=145 \mathrm{~mm})$. CVs ranged from 8-12 $($ average $=9)$. Mean weights ranged from 23-118 g (average $=40 \mathrm{~g}$ ). Fish per pound ranged from 4-31 (average $=18$ fish/pound).

White River Spring Chinook Salmon Hatchery Releases


Figure 18. Average fork length (mm), coefficient of variation (CV) of fork length, average weight (g), and number of fish per pound of juvenile spring Chinook Salmon released in White River for brood years 2002-2013. The dashed horizontal lines represent the target length CV (9; provided for reference - no CV target was formally identified for this program) and fish per pound (18-24 fish per pound range shown on graph).

## Methow River Spring Chinook Salmon

Number Released-The recent goal of the conservation program is to release 133,249 ( $\pm 10 \%$ ) juvenile spring Chinook Salmon into the Methow River annually. During the six-year period under the recent goal of the program, the program exceeded that goal in five brood years and did not achieve it in one brood year (Figure 19). Numbers released ranged from 59,260196,711 (average $=146,810)$ juveniles.

## Methow Spring Chinook



Figure 19. Number of juvenile spring Chinook Salmon released in the Methow River for brood years 1993-2017. The dashed horizontal line represents the target release number (183,334 from 1993-2011 and 133,249 from 2012-2017).

Size at Release-The current goal of the conservation program (since brood year 2012) is to release juvenile spring Chinook Salmon in the Methow River that average 137-mm long (fork length) with a CV of 9.0 , and 30.2 g ( 15 fish/pound). During the six-year period under the recent goal of the program, lengths were near the target while CVs were at or below the target (Figure 20). Mean lengths of fish released ranged from 131-141 mm (average $=134 \mathrm{~mm}$ ), while CVs ranged from 7-9 (average $=8$ ). Both mean weights and fish per pound fluctuated above and below their respective targets during the six-year period (Figure 20). Throughout the recent six years, mean weights ranged from 27-34 g (average $=29 \mathrm{~g}$ ) and fish per pound ranged from 14-17 (average $=16$ fish $/$ pound) .

## Methow Spring Chinook Salmon Hatchery Releases



Figure 20. Average fork length (mm), coefficient of variation (CV) of fork length, average weight (g), and number of fish per pound of juvenile spring Chinook Salmon released in the Methow River for brood years 1993-2017. The dashed horizontal lines represent the target length ( 137 mm ), length CV (9.0), weight ( 30.2 g ), and fish per pound ( 15 fish per pound).

Length-Weight Relationship-There was a significant relationship between the length and weight of juvenile hatchery spring Chinook Salmon (Figure 21). Length explained $94 \%$ of the variation in weight of juvenile fish. In addition, the relationship indicated a mean condition 3.02 for juvenile hatchery spring Chinook Salmon, suggesting near isometric growth. The Fulton condition factor averaged 1.16 (range, 0.64-1.91).

## Methow Spring Chinook



Figure 21. Relationship between fork length (mm) and weight (g) of juvenile Methow River spring Chinook Salmon sampled before release during 2003-2018.

## Chewuch River Spring Chinook Salmon

Number Released-The recent goal of the conservation program is to release 60,516 $( \pm 10 \%)$ juvenile spring Chinook Salmon into the Chewuch River annually. During the six-year period under the recent goal of the program, the program achieved that goal for three and exceed the goal for two of the five brood years with program releases (Figure 22). There was no Chewuch program for brood year 2012. Numbers released ranged from 60,860-72,000 (average $=67,131$ ) juveniles.

## Chewuch Summer Chinook



Figure 22. Number of juvenile spring Chinook Salmon released in the Chewuch River for brood years 1994-2017. The dashed horizontal line represents the target release number (183,333 from 1994-2011 and 60,516 from 2012-2017).

Size at Release-The current goal of the conservation program (since brood year 2012) is to release juvenile spring Chinook Salmon in the Chewuch River that average $136-\mathrm{mm}$ long (fork length) with a CV of 9.0, and 30.2 g ( 15 fish/pound). During the six-year period under the recent goal of the program, lengths were near the target while CVs were both above and below the target (Figure 23). Mean lengths of fish released ranged from 126-134 mm (average $=132$ mm ), while CVs ranged from 6-13 (average $=10$ ). Both mean weights and fish per pound fluctuated above and below their respective targets during the six-year period (Figure 23).
Throughout the recent six years, mean weights ranged from $24-32 \mathrm{~g}$ (average $=28 \mathrm{~g}$ ) and fish per pound ranged from 14-19 (average $=16$ fish/pound).

Chewuch Spring Chinook Salmon Hatchery Releases


Figure 23. Average fork length (mm), coefficient of variation (CV) of fork length, average weight (g), and number of fish per pound of juvenile spring Chinook Salmon released in the Chewuch River for brood years 1992-2017. The dashed horizontal lines represent the target length ( 137 mm ), length CV (9.0), weight ( 30.2 g ), and fish per pound ( 15 fish per pound).

Length-Weight Relationship-There was a significant relationship between the length and weight of juvenile hatchery spring Chinook Salmon (Figure 24). Length explained $95 \%$ of the variation in weight of juvenile fish. In addition, the relationship indicated a mean condition 3.13 for juvenile hatchery spring Chinook Salmon, suggesting positive allometric growth. The Fulton condition factor averaged 1.19 (range, 0.75-2.09).

## Chewuch Spring Chinook



Figure 24. Relationship between fork length (mm) and weight (g) of juvenile Chewuch River spring Chinook Salmon sampled before release during 2003-2018.

## Twisp River Spring Chinook Salmon

Number Released-The recent goal of the conservation program is to release 30,000 $( \pm 10 \%)$ juvenile spring Chinook Salmon into the Twisp River annually. During the six-year period under the recent goal of the program, the program achieved the goal for four brood years and exceeded the goal for two brood years (Figure 25). Numbers released ranged from 29,33348,924 (average $=36,137$ ) juveniles.

## Twisp Spring Chinook



Figure 25. Number of juvenile spring Chinook Salmon released in the Twisp River for brood years 1992-2017. The dashed horizontal line represents the target release number (183,333 from 1992-2011 and 30,000 from 2012-2017).

Size at Release-The current goal of the conservation program (since brood year 2012) is to release juvenile spring Chinook Salmon in the Twisp River that average $135-\mathrm{mm}$ long (fork length) with a CV of 9.0 , and 30.2 g ( 15 fish/pound). During the six-year period under the recent goal of the program, lengths were near the target while CVs were both above and below the target (Figure 26). Mean lengths of fish released ranged from 125-138 mm (average $=132 \mathrm{~mm}$ ), while CVs ranged from 6-11 (average $=9$ ). Mean weights were generally at or below the target while fish per pound fluctuated above and below the target during the six-year period (Figure 26). Throughout the recent six years, mean weights ranged from 25-31 g (average $=28 \mathrm{~g}$ ) and fish per pound ranged from 15-18 (average $=16$ fish/pound).

## Twisp Spring Chinook Salmon Hatchery Releases



Figure 26. Average fork length (mm), coefficient of variation (CV) of fork length, average weight ( g ), and number of fish per pound of juvenile spring Chinook Salmon released in the Twisp River for brood years 1992-2017. The dashed horizontal lines represent the target length ( 135 mm ), length CV (9.0), weight ( 30.2 g ), and fish per pound ( 15 fish per pound).

Length-Weight Relationship-There was a significant relationship between the length and weight of juvenile hatchery spring Chinook Salmon (Figure 27). Length explained 97\% of the variation in weight of juvenile fish. In addition, the relationship indicated a mean condition 2.97 for juvenile hatchery spring Chinook Salmon, suggesting near isometric growth. The Fulton condition factor averaged 1.19 (range, 0.45-2.92).

## Twisp Spring Chinook



Figure 27. Relationship between fork length (mm) and weight (g) of juvenile Twisp River spring Chinook Salmon sampled before release during 2003-2018.

Wenatchee River Summer Chinook Salmon

Number Released-The recent goal of the supplementation program is to release 500,001 ( $\pm 10 \%$ ) juvenile summer Chinook Salmon into the Wenatchee River annually. During the sixyear period under the recent goal of the program, the program achieved that goal for each of the six brood years (Figure 28). Numbers released ranged from 470,570-550,877 (average $=$ 509,522 ) juveniles.


Figure 28. Number of juvenile summer Chinook Salmon released in the Wenatchee River for brood years 1989-2017. The dashed horizontal line represents the target release number $(864,000$ from 1989-2011 and 500,001 from 2012-2017).

Size at Release-The current goal of the supplementation program (since brood year 2012) is to release juvenile summer Chinook Salmon in the Wenatchee River that average 163mm long (fork length) with a CV of 9.0, and 30.0-45.4 g (10-18 fish/pound). During the six-year period under the recent goal of the program, lengths were below the target while CVs were generally above the target (Figure 29). Mean lengths of fish released ranged from 139-158 mm (average $=148 \mathrm{~mm}$ ), while CVs ranged from 7-13 (average $=10$ ). Mean weights were within the target range (brood years 2012-2014) or below the target (brood years 2015-2017). Likewise, fish per pound was within the target range (brood years 2012-2014) or below the target (brood years 2015-2017) (Figure 29). Throughout the recent six years, mean weights ranged from 29-41 $\mathrm{g}($ average $=34 \mathrm{~g})$ and fish per pound ranged from 11-16 (average $=14$ fish/pound).

## Wenatchee Summer Chinook Salmon Hatchery Releases



Figure 29. Average fork length (mm), coefficient of variation (CV) of fork length, average weight (g), and number of fish per pound of juvenile summer Chinook Salmon released in the Wenatchee River for brood years 1989-2017. The dashed horizontal lines represent the target length ( 135 mm from 1989-2011 and 163 from 2012-2017), length CV (9.0), weight ( 45.4 g with a range of 30.0-45.4 g for brood years 2012-2014), and fish per pound ( 10 fish per pound for brood years 1989-2011, a range of 10-15 fish per pound for brood years 2012-2014, and 18 fish per pound for brood years 2015-2017).

Length-Weight Relationship-There was a significant relationship between the length and weight of juvenile hatchery summer Chinook Salmon (Figure 30). Length explained 94\% of the variation in weight of juvenile fish. In addition, the relationship indicated a mean condition 2.99 for juvenile hatchery summer Chinook Salmon, suggesting near isometric growth. The Fulton condition factor averaged 1.04 (range, 0.37-2.90).


Figure 30. Relationship between fork length (mm) and weight (g) of juvenile Wenatchee River summer Chinook Salmon sampled before release during 2003-2018.

## Chelan Falls Summer Chinook Salmon

Number Released-The goal of the supplementation program is to release 576,000 $( \pm 10 \%)$ juvenile summer Chinook Salmon into the Chelan River annually. During the eight-year period of the program, the program achieved the goal for six brood years, and was below the goal for two brood years (Figure 31). Numbers released ranged from 442,063-600,894 (average $=543,629)$ juveniles.

## Chelan Falls Summer Chinook



Figure 31. Number of juvenile summer Chinook Salmon released in the Chelan River for brood years 2010-2017. The dashed horizontal line represents the target release number $(576,000)$.

Size at Release-The goal of the supplementation program is to release juvenile summer Chinook Salmon in the Chelan River that average 161 -mm long (fork length) with a CV of 9.0, and 20.0-45.4 g (10-22 fish/pound) depending on brood year. During the eight-year period of the program, lengths were below the target while CVs were above the target (Figure 32). Mean lengths of fish released ranged from 129-148 mm (average $=140 \mathrm{~mm}$ ), while CVs ranged from 10-27 (average $=16$ ). Mean weights were within the target range (brood years 2012-2014) or below the target (brood years 2010-2011 and 2015-2017). Fish per pound was within the target range (brood years 2012-2014) or near the target (brood years 2010-2011 and 2015-2017) (Figure 32). Throughout the recent eight years, mean weights ranged from 25-43 g (average $=34$ g) and fish per pound ranged from 11-19 (average $=14$ fish/pound).

## Chelan Falls Summer Chinook Salmon Hatchery Releases



Figure 32. Average fork length (mm), coefficient of variation (CV) of fork length, average weight (g), and number of fish per pound of juvenile summer Chinook Salmon released in the Chelan River for brood years 2010-2017. The dashed horizontal lines represent the target length $(161 \mathrm{~mm})$, length CV $(9.0)$, weight ( 45.4 g with a range of 20.0-45.4 g for brood years 20122014), and fish per pound ( 13 fish per pound for brood years 2010-2011, a range of 10-22 fish per pound for brood years 2012-2014, and 13 fish per pound for brood years 2015-2017).

Length-Weight Relationship-There was a significant relationship between the length and weight of juvenile hatchery summer Chinook Salmon (Figure 33). Length explained 95\% of the variation in weight of juvenile fish. In addition, the relationship indicated a mean condition 3.12 for juvenile hatchery summer Chinook Salmon, suggesting positive allometric growth. The Fulton condition factor averaged 1.09 (range, 0.44-1.99).

## Chelan Falls Summer Chinook



Figure 33. Relationship between fork length (mm) and weight (g) of juvenile Chelan Falls summer Chinook Salmon sampled before release during 2010-2018.

## Methow River Summer Chinook Salmon

Number Released-The recent goal of the supplementation program is to release 200,000 ( $\pm 10 \%$ ) juvenile summer Chinook Salmon into the Methow River annually. During the six-year period under the recent goal of the program, the program achieved that goal for three brood years and was below the goal for three brood years (Figure 34). Numbers released ranged from 143,594-209,490 (average $=180,781$ ) juveniles.

Methow Summer Chinook


Figure 34. Number of juvenile summer Chinook Salmon released in the Methow River for brood years 1989-2017. The dashed horizontal line represents the target release number (400,000 from 1989-2011 and 200,000 from 2012-2017).

Size at Release-The current goal of the supplementation program (since brood year 2012) is to release juvenile summer Chinook Salmon in the Methow River that average $163-\mathrm{mm}$ long (fork length) with a CV of 9.0 , and 45.4 g (13-18 fish/pound). During the six-year period under the recent goal of the program, lengths were below the target while CVs fluctuated above and below the target (Figure 35). Mean lengths of fish released ranged from 125-158 mm (average $=136 \mathrm{~mm}$ ), while CVs ranged from 8-13 (average $=10$ ). Mean weights were below the target while fish per pound was within the target range (Figure 35). Throughout the recent six years, mean weights ranged from 23-42 g (average $=29 \mathrm{~g}$ ) and fish per pound ranged from 11-20 (average $=16$ fish/pound).

## Methow Summer Chinook Salmon Hatchery Releases



Figure 35. Average fork length (mm), coefficient of variation (CV) of fork length, average weight $(\mathrm{g})$, and number of fish per pound of juvenile summer Chinook Salmon released in the Methow River for brood years 1989-2017. The dashed horizontal lines represent the target length ( 176 mm from 1989-2011 and 163 from 2012-2017), length CV (9.0), weight ( 45.4 g ), and fish per pound ( 10 fish per pound for brood years 1989-2011 and a range of 13-18 fish per pound for brood years 2012-2017).

Length-Weight Relationship-There was a significant relationship between the length and weight of juvenile hatchery summer Chinook Salmon (Figure 36). Length explained 95\% of the variation in weight of juvenile fish. In addition, the relationship indicated a mean condition 2.83 for juvenile hatchery summer Chinook Salmon, suggesting negative allometric growth. The Fulton condition factor averaged 1.12 (range, 0.54-2.82).

## Methow Summer Chinook



Figure 36. Relationship between fork length (mm) and weight (g) of juvenile Methow River summer Chinook Salmon sampled before release during 2003-2018.

## Wells Subyearling Summer Chinook Salmon

Number Released-The goal of the supplementation program is to release 484,000 $( \pm 10 \%)$ subyearling summer Chinook Salmon from the Wells Fish Hatchery annually. During the 25 -year period of the program, the program did not achieve that goal in 10 brood years, it achieved that goal in 14 brood years, and exceeded the goal in 1 brood year (Figure 37). Numbers released ranged from 187,382-541,923 (average $=433,474$ ) juveniles.

Wells Subyearling Summer Chinook


Figure 37. Number of subyearling summer Chinook Salmon released from the Wells Fish Hatchery for brood years 1993-2017. The dashed horizontal line represents the target release number (484,000 except for 1998-1999).

Size at Release-The current goal of the supplementation program (since brood year 2012) is to release subyearling summer Chinook Salmon from the Wells Fish Hatchery that average $9.1 \mathrm{~g}(50$ fish/pound) and have a fork length CV of 7.0 (there was not fork length goal from 2012-2017). The Wells subyearling program is reared to achieve release by a certain date and not to achieve a specific size. During the six-year period under the recent goal of the program, CVs fluctuated above and below the target (Figure 38). CVs ranged from 6-11 (average $=7$ ). Mean weights were below the target while fish per pound was above the target (Figure 38). Throughout the recent six years, mean weights ranged from 6-8 g (average $=7 \mathrm{~g}$ ) and fish per pound ranged from 55-78 (average $=65$ fish/pound).

## Wells Subyearling Summer Chinook Salmon Hatchery Releases



Figure 38. Average fork length (mm), coefficient of variation (CV) of fork length, average weight $(\mathrm{g})$, and number of fish per pound of subyearling summer Chinook Salmon released from the Wells Fish Hatchery for brood years 1998-2017. The dashed horizontal lines represent the target length ( 116 mm from 1998-2011; there was no target from 2012-2017), length CV (9.0 from 1998-2011 and 7.0 from 2012-2017), weight ( 22.7 g from 1998-2011 and 9.1 from 20122017), and fish per pound ( 20 fish per pound for brood years 1998-2011 and 50 fish per pound for brood years 2012-2017).

Length-Weight Relationship-There was a significant relationship between the length and weight of juvenile hatchery summer Chinook Salmon (Figure 39). Length explained 99\% of the variation in weight of juvenile fish. In addition, the relationship indicated a mean condition 2.84 for juvenile hatchery summer Chinook Salmon, suggesting negative allometric growth. The Fulton condition factor averaged 1.12 (range, 0.62-2.91).

## Wells Summer Chinook



Figure 39. Relationship between fork length (mm) and weight (g) of juvenile Wells summer Chinook Salmon sampled before release during 2003-2018.

## Wells Yearling Summer Chinook Salmon

Number Released-The goal of the supplementation program is to release 320,000 ( $\pm 10 \%$ ) yearling summer Chinook Salmon from the Wells Fish Hatchery annually. During the 26 -year period of the program, the program achieved that goal in 18 brood years, exceeded it in 7 brood years, and did not achieve it in 1 brood year (Figure 40). Numbers released ranged from $185,200-457,770$ (average $=330,558$ ) juveniles.

Wells Yearling Summer Chinook


Figure 40. Number of yearling summer Chinook Salmon released from the Wells Fish Hatchery for brood years 1992-2017. The dashed horizontal line represents the target release number $(320,000)$.

Size at Release-The current goal of the supplementation program (since brood year 2012) is to release yearling summer Chinook Salmon from the Wells Fish Hatchery that average $168-\mathrm{mm}$ long (fork length with a CV of 7.0 ), and 45.4 g ( 10 fish/pound). During the six-year period under the recent goal of the program, lengths were near the target while CVs fluctuated above and below the target (Figure 41). Mean lengths of fish released ranged from 153-168 mm (average $=163 \mathrm{~mm}$ ), while CVs ranged from 6-9 (average $=8$ ). Mean weights were generally above the target while fish per pound was generally below the target during the six-year period (Figure 41). Throughout the recent six years, mean weights ranged from 38-50 g (average $=46$ g) and fish per pound ranged from 9-12 (average $=10$ fish/pound).

## Wells Yearling Summer Chinook Salmon Hatchery Releases



Figure 41. Average fork length (mm), coefficient of variation (CV) of fork length, average weight ( g ), and number of fish per pound of yearling summer Chinook Salmon released from the Wells Fish Hatchery for brood years 1997-2017. The dashed horizontal lines represent the target length ( 162 mm from 1997-2011 and 168 mm from 2012-2017), length CV (9.0 from 1997-2011 and 7.0 from 2012-2017), weight ( 45.4 g ), and fish per pound (10).

Length-Weight Relationship-The length-weight results for Wells Fish Hatchery summer Chinook Salmon are shown under the Wells Subyearling Summer Chinook Salmon section.

## Discussion

Programs generally met or exceeded the program release number targets. Specific programs did not meet the release number targets in all years, but all programs met or exceeded targets in the majority of years, with the exception of the White River spring Chinook Salmon program. This program was in development for all of its history and tested numerous fish culture and release strategies. Therefore, the program was not able to meet targets that remained consistent across years. There are a variety of reasons why release targets were not met in some years. The Hatchery Committees developed and approved hatchery implementation plans that specify the number of broodstock to collect to meet target production goals. These plans attempted to predict the number of eggs produced per female based upon mean fecundities during the previous 5 years. In addition, the mean hatchery survival was used to determine how
many eggs are needed to produce the desired number of smolts to release. Lower than average fecundities or hatchery survivals was the main reason for failing to meet hatchery release targets.

In general, the length targets, and some of the metrics that used a single value (e.g., length and weight) associated with a range of fpp targets, were not useful. When programs meet the fish per pound metric the length metric cannot be met. This metric should be discarded or realistic length targets need to be developed. Most programs met or were close to meeting the CV target on average. Meeting the CV targets in steelhead programs is more difficult than in the Chinook Salmon programs. Meeting fish-per-pound targets was more difficult in spring Chinook Salmon and conservation steelhead programs. This may be because these programs use natural-origin broodstock and also because attempts are made to reduce precocious maturation by slowing growth or producing fish that do not exceed a size threshold. Spring Chinook Salmon tended to be slightly smaller than the target. Fish were reared to balance achieving threshold size for release while hedging against larger fish that may exhibit higher rates of precocity. In addition, meeting the 6 fpp target for conservation steelhead was difficult, and likely not desirable, due to the short rearing time with spring-spawned steelhead programs. Those Chinook Salmon programs that target a range of fpp at release were sometimes paired with single values for length and weight (e.g., Nason and White spring Chinook, Methow summer Chinook). This was likely a legacy of past targets that were not updated when the fpp ranges were updated to reduce unnaturally high proportions of precocious maturation. The programs that have an updated range for target fpp have been managed primarily to hit the fpp target rather than individual length or weight targets. Targets for programs with a range of target fpps are good candidates for revision.

Spring Chinook Salmon exhibited near-isometric growth. Steelhead exhibited negative allometric growth, as did Wells and Methow summer Chinook Salmon. However, Wenatchee and Chelan Falls summer Chinook Salmon exhibited isometric or positive allometric growth. Condition factors across all programs were very close to or exceeded 1.

Trade-offs between post release survival and age at maturation influence the hatchery rearing strategies. Large Chinook Salmon generally survive better than small fish, but they also are more prone to maturation at younger ages. Hatchery programs that use natural-origin fish for broodstock may also experience higher levels of precocious maturation than if hatchery-origin fish were used for broodstock (Larsen et al. 2020). Attempts to reduce precocious maturation of Chinook Salmon released into Nason Creek has also resulted in smaller fish that don't meet some of the size targets. In addition, larger fish also pose ecological risks to other fish through mechanisms such as competition or predation. Fish that are released from hatcheries continue to be substantially larger than natural-origin fish, even when they fail to meet the size targets. Evaluations of survival and age at maturation that are contained in other portions of the monitoring plan will inform whether current size at release targets are achieving the acceptable trade-offs desired by fishery managers. Adaptation of targets may occur to achieve a better suite of benefits among the many trade-offs involved in growing fish to a target size or weight.

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# Harvest of Chinook Salmon and Steelhead Originating from Upper Columbia River Hatchery Programs 

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#### Abstract

The objective of this evaluation was to determine if a diversity of upper Columbia Basin Chinook Salmon and steelhead hatchery programs contributed to harvest. More specifically, we were interested in evaluating whether harvest rates were consistent with management objectives and where fish were harvested. Harvest rates were lowest on endangered spring Chinook Salmon with annual brood year means of $5-6 \%$ for Methow, Chewuch, and Twisp spawning aggregates (annual range 0 to $59 \%$ ) and $26 \%$ for the Chiwawa spawning aggregate (annual range 0 to $95 \%$ ). The percent of the population harvested was not correlated with spawning escapement ( $\mathrm{P}>0.05$ ) and the total number of fish harvested was correlated with spawning escapement ( $\mathrm{P}<0.05$ ) in the Chiwawa and Twisp rivers but not in the Methow or Chewuch rivers. Most harvest of spring Chinook Salmon occurred in freshwater. Harvest rates were much higher for the more abundant summer and fall Chinook Salmon programs with annual brood year averages around $53-75 \%$ and annual ranges of 14 to $91 \%$. Percent harvest increased with increasing spawning escapement for summer Chinook in the Methow ( $\mathrm{P}=0.01$ ) and Okanogan ( $\mathrm{P}=0.0002$ ) rivers but not for summer Chinook in the Wenatchee River ( $\mathrm{P}=0.49$ ), Chelan Falls/Turtle Rock program ( $\mathrm{P}=0.43$ ), and Hanford Reach fall Chinook ( $\mathrm{P}=0.28$ ). The total number fish harvested was not correlated with spawning escapement $(\mathrm{P}>0.05)$ for the Wenatchee River, Wells subyearling, Methow River, or Okanogan River programs, but significant correlations were detected ( $\mathrm{P}<0.05$ ) for the Chelan Falls/Turtle Rock yearling and Wells yearling programs and for fall Chinook Salmon from Priest Rapids Hatchery. Most of the harvest of summer Chinook Salmon occurred in the ocean and harvest of fall Chinook Salmon occurred evenly between freshwater and the ocean. Harvest rates averaged $16 \%$ (range $0-54 \%$ ) for threatened hatchery-origin steelhead and less than 5\% (range 0 to $4 \%$ ) for natural-origin steelhead. The percent of steelhead harvested increased with increasing escapement in the Okanogan River ( $\mathrm{P}=0.006$ ) but was not significantly correlated in the Methow ( $\mathrm{P}=0.29$ ) and Wenatchee rivers $(\mathrm{P}=0.85)$. Total harvest of hatchery steelhead was not significantly correlated with spawning escapement in the Methow or Wenatchee rivers ( $\mathrm{P}>0.05$ ) but was correlated in the Okanogan River $(\mathrm{P}=0.006)$. Every hatchery program that was evaluated contributed to harvest and sometimes substantially. The magnitude of harvest generally corresponded to the status of the population: the lowest harvest occurred on the most imperiled stocks and the highest harvest occurred on the healthiest stocks. However, harvest sometimes hindered meeting broodstock collection goals and harvest management of endangered or threatened species could impede conservation objectives and might be improved by tailoring harvest to abundance, weak stocks, and weak broodyears.


## Introduction

One of the main functions of salmon and steelhead hatcheries is to increase the opportunity for harvest. However, there are a diversity of harvest objectives associated with different types of hatcheries. In some cases, the sole objective of hatcheries is to produce maximal harvest. These hatcheries are often segregated from naturally spawning populations and the goal of harvesters is to harvest all the fish produced by the hatchery except for those needed for the next brood cycle (Mobrand et al. 2005; Paquet et al. 2011). In other cases, the main objective of a hatchery is to aid in the recovery of depressed populations and harvest is
incidental to natural production objectives. These hatcheries are often referred to as conservation or integrated, and harvest is intentionally negligible so that returns from these programs can contribute to natural production. Finally, other hatcheries fall on a continuum between the two extremes described above, sharing both harvest and conservation objectives within the same hatchery. Harvest from such programs is largely determined by what the population can sustain into the future as well as constraining impacts to non-target populations within acceptable levels.

Harvest rates and allocations are set within complicated processes and agreements among fisheries co-managers. Harvest rates can be determined based upon maximum sustainable yield (MSY), allowable take of ESA listed species or weak stocks, desired escapement objectives, need for removal of hatchery-origin fish for conservation purposes, and a variety of other approaches (Maier 2020). In some cases, fisheries managers focus on selectively harvesting hatchery-origin fish so that the natural-origin fish escape to the spawning grounds. One of the main assumptions of science-based harvest management is that harvestable surplus increases with increasing population sizes particularly when carrying capacity is exceeded.

Harvest of upper Columbia River Chinook Salmon and steelhead occurs across three primary fisheries: ocean commercial (treaty and non-treaty, reported together), Columbia River commercial (treaty and non-treaty, reported separately), and recreational fishing. The timing of each fishery is set to target stocks intended for harvest. For example, ocean commercial fisheries typically begin in early summer to avoid harvest of Upper Columbia spring Chinook Salmon, which primarily enter the river from March through June, and instead focus on summer and fall Chinook Salmon stocks. In the upper river, conservation fisheries for recreational anglers are timed to remove hatchery-origin adults to prevent them from reaching spawning areas when that outcome is desired. Some fisheries are mark-selective, meaning that only hatchery-origin fish with a visible external mark (i.e. a clipped adipose fin) may be retained. The goal of markselective fisheries is to allow unmarked fish to be released to continue migration and reach spawning areas. Non-selective fisheries allow harvest of all stocks but are timed to reduce impacts to non-target and/or natural-origin fish.

Most, but not all, hatchery programs mark or tag some portion of annual releases. This practice necessitates an expansion calculation to estimate overall harvest from monitoring data collected from each fishery. In addition to visible external marks, other common methods include coded-wire tags (CWT) implanted in the snout of juvenile fish allowing identification of fish origin and brood year, and passive integrated transponder tags (PIT) implanted in the body cavity of juvenile fish or dorsal musculature of adults that provide a unique identification code. Coded-wire tags must be recovered from dead fish to be read, while PIT-tags can be read by transponders located in mainstem Columbia River dams and throughout the Columbia River watershed as fish move throughout the system (Pearsons and O’Connor 2020). Both CWT and PIT-tag records are aggregated in regional databases for the purpose of analysis.

Harvesting fish can produce undesirable unintended consequences. For example, overharvest is one factor that has contributed to species or population declines. It can also result in changes to population demographics resulting in reduced population productivity and difficulty in evaluating hatchery effects on natural populations. For example, non-random harvesting of the hatchery- and natural-origin components of the population can skew sex ratios, decrease age at maturity, or influence run and spawn timing, resulting in changes in these metrics through time. In addition, selective harvest of hatchery-origin fish can result in differences in these metrics within a year. The size of Chinook Salmon has decreased during the past decades and one possible mechanism for this reduced size is harvest (Ohlberger et al. 2018, 2020).

The upper Columbia River Public Utility Districts’ (Grant, Chelan, and Douglas PUDs) hatchery programs are guided by harvest monitoring indicators described in the Monitoring and Evaluation Plan for PUD Hatchery Programs (Hillman et al. 2019). The plan states that "Harvest will be applied to different types of programs in an effort to achieve the management objectives of those programs. Programs designed to augment harvest should routinely contribute to harvest at a rate that greatly reduces the incidence of straying to natural spawning grounds, but also allows the program to be sustained. Safety-net programs may be harvested as part of an adult management strategy to minimize excessive escapement of hatchery-origin fish to spawning grounds. Similarly, conservation programs may undergo harvest to manage returning adults, but the emphasis for these programs should be to achieve escapement goals. In all cases, harvest effort should not have the unintended consequence of removing excessive numbers of conservation or natural-origin fish. In years when the expected returns of hatchery adults are above the level required to meet program goals (i.e., supplementation of spawning populations and/or brood stock requirements), surplus fish may be available for harvest." The plan broadly captures the differences in harvest goals of each hatchery program and sets forth monitoring questions to "determine if appropriate harvest rates have been applied to conservation, safety-net, and segregated harvest programs to meet the Habitat Conservation Plan (HCP)/Salmon and Steelhead Settlement Agreement (SSSA) goal of providing harvest opportunities while also contributing to population management and minimizing risk to natural populations".

The objective of this analysis was to determine whether a diversity of upper Columbia Basin salmon and steelhead hatchery programs contributed to harvest. More specifically our objective was to determine whether harvest levels were consistent with management objectives of the hatchery programs. To evaluate these goals we report spawning escapement, number of fish harvested, percent of brood year harvested, and the proportion harvested in various fisheries for each hatchery program.

## Methods

Spawning escapement, number of fish harvested, percent of brood year harvested, and fishery proportion data were aggregated from Grant, Chelan, and Douglas PUD hatchery monitoring and evaluation reports (Richards and Pearsons 2019; Hillman et al. 2020; Snow et al. 2020). The quantities of harvested Chinook Salmon and percent of brood year harvested represent the totals from the hatchery program and exclude natural-origin stocks. Creel survey data for natural-origin steelhead were included in our analyses. We compared among conservation and safety-net hatchery programs for spring Chinook Salmon and steelhead as well as harvest-augmentation programs for summer and fall Chinook Salmon. We also compared percent of brood year harvested with spawning escapement abundance to assess trends when there was a range of spawning escapement. For all Chinook Salmon comparisons, the spawning escapement data were reported for return years (spawn year) and harvest data were reported for brood years. Both spawning escapement and harvest data for steelhead were reported as the span of return migration year and spawn year (i.e. 2002-2003). The plots of spawning escapement versus percent of brood year harvested and total number harvested show a line of best fit, equation of the fit, the $\mathrm{R}^{2}$ value, and F -test results. Other plots used actual values from the annual reports and means of fishery proportions for the included brood years.

As described in the PUD hatchery monitoring and evaluation reports, the Regional Mark Information System (RMIS) database was used to estimate harvest of coded-wire tagged hatchery stocks using an expanded sample rate during the data collection event and the tag-codespecific mark rate for the population. Percent of brood year harvested for Chinook Salmon represents the sum of all harvest in fisheries divided by sum of all harvest in fisheries plus spawning escapement and broodstock collection. Local creel sampling was used to estimate steelhead harvest.

Table 1. Types of harvest that occurred for spring Chinook Salmon (SPC), summer Chinook Salmon (SUC), fall Chinook Salmon (FAC), and steelhead (STH) in the upper Columbia River Public Utility District's conservation and harvest-augmentation hatchery programs. Salmon harvest results were reported for brood years (BY) and steelhead results were reported for return years (RY).

| Species <br> / race | Program | Program Type | Years | Ocean <br> Commercial | Columbia River Tribal | Columbia River Commercial | Recreational |
| :---: | :---: | :---: | :---: | :---: | :---: | :---: | :---: |
| SPC | Chiwawa | Conservation | BY 1989-2012 | x | x | x | x |
| SPC | Methow | Conservation | BY 1993-2012 | X | x | x | x |
| SPC | Twisp | Conservation | BY 1992-2012 | no data | x | x | x |
| SPC | Chewuch | Conservation | BY 1992-2012 | x | x | x | x |
| SUC | Wenatchee | Harvestaugmentation | BY 1989-2012 | x | x | x | x |
| SUC | Chelan Falls/ <br> Turtle Rock | Harvestaugmentation | BY 1995-2012 | x | x | x | x |
| SUC | Wells Hatchery subyearling | Harvestaugmentation | BY 1993-2012 | x | x | x | x |
| SUC | Wells Hatchery yearling | Harvestaugmentation | BY 1993-2012 | x | x | x | x |
| SUC | Methow | Harvestaugmentation | BY 1989-2012 | x | x | x | x |
| SUC | Okanogan | Harvestaugmentation | BY 1989-2012 | x | x | x | x |
| FAC | Priest Rapids Hatchery | Harvestaugmentation | BY 1997-2012 | x | x | x | x |
| STH | Wenatchee | Conservation | RY 2007-2019 |  | x |  | x |
| STH | Methow | Conservation | RY 2002-2019 |  | x |  | x |
| STH | Okanogan | Conservation / safety net | RY 2003-2019 |  | x |  | x |

## Results

## Hatchery Spring Chinook Salmon

Annual spawning escapement of upper Columbia River hatchery-origin Spring Chinook Salmon to the Methow, Twisp, and Chewuch rivers was typically fewer than 1,000 individuals and average harvest was less than $10 \%$ of brood year production (Figure 1). Chiwawa River spawning escapement was generally 1,000-2,000 individuals and harvest averaged $25.6 \%$ of brood year production between 2003-2012. The percent of brood year harvested was as high as $95 \%$ for the Chiwawa and $60 \%$ for some brood years in the Methow, and these high harvest rates occurred when spawning escapement was relatively low. The percent of harvest was not significantly correlated with spawning escapement ( $\mathrm{P}>0.05$; Figure 2 ). The total number of fish harvested was correlated with spawning escapement $(\mathrm{P}<0.05)$ in the Chiwawa and Twisp rivers but not in the Methow or Chewuch rivers. The bulk of harvest occurred in tribal ( $\bar{x}=47 \%$ ) and sport ( $\bar{x}=31 \%$ ) fisheries (Figure 3). Commercial fisheries in the ocean ( $\bar{x}=9 \%$ ) and lower Columbia River ( $\bar{x}=13 \%$ ) accounted for the remaining harvest.

Spawning escapement for Chiwawa River spring Chinook Salmon was low enough in the late 1980's and throughout the 1990's that the broodstock collection goal of 379 individuals was rarely met. Beginning in brood year 2000, spawning escapement improved, and broodstock collection goals were met in most years. Broodstock collection was revised down to 74 individuals beginning in 2009 and spawning escapement has been well above that number since then. Ocean and non-treaty Columbia River commercial harvest was low for these fish; however, tribal harvest exceeded 100 individuals in 7 of 24 years and recreational harvest exceeded 100 individuals in 14 out of 24 years.

The spawning escapement for the aggregated Methow River Basin spring Chinook Salmon programs, which includes production in the Twisp and Chewuch rivers, followed a pattern similar to the Chiwawa River program. The broodstock collection goal of 104 individuals was rarely met in the 1990s but since brood year 2000 the goal has generally been met. While ocean and non-treaty Columbia River harvest was low, there were two years when tribal and recreational harvest of Methow River hatchery-origin Spring Chinook salmon both exceeded 100 individuals.


Figure 1. Spawning escapement, total harvest, and percent of brood year harvested for hatchery spring Chinook Salmon from the Chiwawa ( $\bar{x}=25.6 \%$ ), Methow ( $\bar{x}=5.1 \%$ ), Chewuch ( $\bar{x}=$ $5.8 \%$ ), and Twisp ( $\bar{x}=4.6 \%$ ) rivers (averages represent percent of brood year harvested over brood years 2004-2012).




Figure 2. Spawning escapement versus percent of brood year harvested and total number of hatchery fish harvested for spring Chinook Salmon from the Chiwawa, Methow, Chewuch, and Twisp rivers.


Figure 3. Fishery proportions (mean values) for spring Chinook Salmon harvested from the Chiwawa, Methow, Twisp, and Chewuch rivers.

Hatchery Summer and Fall Chinook Salmon
Annual spawning escapement of Upper Columbia River hatchery summer and fall Chinook Salmon to the Hanford Reach of the Columbia River, Wenatchee, Chelan, Methow, and Okanogan rivers was highly variable among programs and ranged from fewer than 100 individuals for releases directly into the Columbia River from Wells Hatchery to over 90,000 individuals in a single year for Priest Rapids Hatchery fall Chinook Salmon released into the Hanford Reach (summer Chinook Salmon Figure 4, fall Chinook Salmon Figure 5). The average escapement for most programs was fewer than 10,000 individuals. The annual brood year harvest of summer Chinook Salmon ranged from $25.4-80.2 \%$ in the Wenatchee, 17.6-75.6\% in the Methow, 14.0-89.4\% in the Okanogan, 42.9-91.4\% for subyearlings from Wells Hatchery, 24.5$89.5 \%$ for yearlings from Wells Hatchery, 50.2-84.3\% for yearlings from Chelan Falls Hatchery, and 33.8-72.5\% for fall Chinook Salmon from Priest Rapids Hatchery. The percent of brood year harvested increased with increasing spawning escapement for summer Chinook Salmon in the Methow ( $\mathrm{P}=0.01$ ) and Okanogan ( $\mathrm{P}=0.0002$ ) rivers but not for summer Chinook Salmon in the Wenatchee River ( $\mathrm{P}=0.49$ ) and Hanford Reach fall Chinook ( $\mathrm{P}=0.28$ ) (Figure 6). The total number of fish harvested was not correlated with spawning escapement $(\mathrm{P}>0.05)$ for the Wenatchee River, Wells subyearling, Methow River, or Okanogan River programs, but significant correlations were detected $(\mathrm{P}<0.05)$ for the Chelan Falls/Turtle Rock yearling and Wells yearling programs and for fall Chinook Salmon from Priest Rapids Hatchery. Harvest of Wells Hatchery summer Chinook Salmon was generally high, averaging $67 \%$ but uniformly small escapement numbers precluded our ability to assess trends in harvest. Ocean commercial
fisheries accounted for an average of $61 \%$ of observed harvest for all populations (Figure 7). Tribal ( $\bar{x}=21 \%$ ), recreational ( $\bar{x}=14 \%$ ), and lower Columbia commercial fishing ( $\bar{x}=4 \%$ ) accounted for the remaining harvest.

The upper Columbia River hatchery augmentation programs for summer and fall Chinook Salmon have sustained harvest rates often exceeding 50\% of brood year production since the late 1990s. The Methow, Chelan Falls/Turtle Rock, and Wells programs are segregated hatchery programs and returning adults are not intended for spawning in the natural environment. As such, spawning escapement was fewer than 5,000 individuals. Spawning escapement was fewer than 10,000 in the Wenatchee and Okanogan rivers. From 1989-1999, the broodstock collection goal for Wenatchee River summer Chinook Salmon (n=492 individuals) was met only once. From 2000-2011, collection was met or within $10 \%$ of the goal in all but two years as escapement improved. The broodstock collection goal was revised down to 262 individuals in 2012 and the goal has been met each year since. The percent of brood year harvested was at least $60 \%$ twice during the period of 1989-1999 when escapement was low. The broodstock collection goals for the Methow/Okanogan ( $\mathrm{n}=222$ ), Wells age-0 $(\mathrm{n}=284)$ and age-1 $(\mathrm{n}=178)$ programs were met in all years. Broodstock collection for the Chelan Falls/Turtle Rock summer Chinook Salmon program ranged from 318-591 fish from brood year 2013 to brood year 2019 but no specific collection goal is specified. The broodstock collection goals for fall Chinook Salmon at Priest Rapids Hatchery have varied since 1991 but the goal was met each year except for an unusually low return year in 2007.


Figure 4. Spawning escapement, total harvest, and percent of brood year harvested for hatcheryorigin summer Chinook Salmon from the Wenatchee River ( $\bar{x}=67.8 \%$ ), Chelan Falls/Turtle Rock yearling program ( $\bar{x}=74.6 \%$ ), Wells Hatchery yearling program ( $\bar{x}=67.6 \%$ ), Wells Hatchery subyearling program ( $\bar{x}=67.3 \%$ ), Methow River ( $\bar{x}=62.4 \%$ ), and Okanogan River ( $\bar{x}$ $=70.4 \%$ ) programs (averages represent percent of brood year harvested over brood years 20042012).


Figure 5. Spawning escapement, total harvest, and percent of brood year harvested for hatchery fall Chinook Salmon from Priest Rapids Hatchery ( $\bar{x}=52.5 \%$ ) program (average represent percent of brood year harvested over brood years 2004-2012).



—Percent of brood year harvested —Total number harvested


Figure 6. Spawning escapement versus percent of brood year harvested and number harvested for hatchery-origin summer Chinook Salmon from the Wenatchee River, Chelan Falls/Turtle Rock yearling program, Wells Hatchery yearling program, Wells Hatchery subyearling program, Methow River, Okanogan River, and fall Chinook Salmon from Priest Rapids Hatchery.


Figure 7. Fishery proportions (mean values) for summer Chinook Salmon harvested from the Wenatchee River, Chelan Falls/Turtle Rock yearling program, Wells Hatchery subyearling program, Wells Hatchery yearling program, Methow River, Okanogan River, and fall Chinook Salmon from Priest Rapids Hatchery.

Steelhead
Escapement and harvest of hatchery steelhead was greatest in the Methow River, followed by the Okanogan and Wenatchee rivers (Figure 8). Escapement and percent harvest of hatchery steelhead peaked from 2010-2012 with 6,000-11,000 individuals escaped to the Okanogan and Methow rivers respectively, and harvest rates of $40-50 \%$. Harvest ranged from $5.3-53.9 \%$ in the Methow, $4.5-47.4 \%$ in the Okanogan, and from $8.0-12.5 \%$ in the Wenatchee River. Origin-based escapement estimates for steelhead returning to the Wenatchee River were not available prior to the 2011-2012 return year, but since then, hatchery-origin escapement was consistently below the Methow and Okanogan rivers, with a peak of around 2,000 individuals. Percent harvest increased with increasing escapement in the Okanogan $(\mathrm{P}=0.006)$ river but was not significantly correlated with escapement in the Methow ( $\mathrm{P}=0.29$ ) and Wenatchee rivers ( $\mathrm{P}=0.85$ ) (Figure 9). Total harvest of hatchery steelhead was not significantly correlated with spawning escapement in the Methow or Wenatchee rivers ( $\mathrm{P}>0.05$ ) but was correlated in the Okanogan River ( $\mathrm{P}=0.006$ ).

Escapement of natural-origin steelhead was greatest in the Methow and Wenatchee rivers, with peaks of greater than 1,200 individuals in the Methow River during the 2009-2010 and 2015-2016 return years and peaks of similar magnitude in the Wenatchee River during the 2011-2012, 2012-2013, and 2015-2016 return years (Figure 10). Origin-based escapement estimates were not available for the Wenatchee River prior to the 2011-2012 return year. Escapement to the Okanogan River was typically 200-400 individuals and was consistently lower than the Wenatchee and Methow rivers. Reported harvest of natural-origin steelhead was less than $6 \%$ of escapement. Harvest was greatest in the Methow and Okanogan rivers (up to 5\% of escapement in return year 2011-2012), and lower in the Wenatchee (range 1-2\% of escapement). Harvest increased with increasing escapement in the Methow ( $\mathrm{P}=0.004$ ) and Okanogan ( $\mathrm{P}=0.09$ ) but did not in the Wenatchee ( $\mathrm{P}=0.89$ ) (Figure 11). Total harvest of naturalorigin steelhead was correlated with spawning escapement in the Methow and Okanogan rivers ( $\mathrm{P}<0.05$ ) but not in the Wenatchee River ( $\mathrm{P}=0.44$ ).

Spawning escapement for hatchery-origin Wenatchee River steelhead has exceeded the broodstock collection goal of 140 individuals since return year 2011-2012, when origin-based escapement data were available. Escapement of hatchery-origin steelhead to the Methow and Okanogan was more than the 170 individuals required for the Douglas PUD safety-net program for all years examined. Escapement of natural-origin steelhead to the Methow River was well above the 28 individuals required for the Twisp River conservation program. Escapement was sufficient to allow harvest of steelhead in the Wenatchee River in 8 of the last 12 return years. Harvest in the Methow and Okanogan rivers occurred in 13 of the last 17 return years.


Figure 8. Escapement, total number harvested, and percent harvest of hatchery-origin steelhead escapement to the Wenatchee ( $\bar{x}=8.7 \%$ ), Methow ( $\bar{x}=20.2 \%$ ), Okanogan ( $\bar{x}=18.6 \%$ ) rivers (averages represent return years 2003-2017 for the Methow and Okanogan rivers and 2011-2016 for the Wenatchee River).

— Percent of hatchery escapement harvested — Total number harvested


Figure 9. Escapement versus percent of escapement harvested and number harvested of hatcheryorigin steelhead from the Wenatchee, Methow, and Okanogan rivers.


Figure 10. Escapement, total number harvested, and percent of escapement harvested of naturalorigin steelhead for the Wenatchee ( $\bar{x}=1.36 \%$ ), Methow ( $\bar{x}=2.17 \%$ ), and Okanogan ( $\bar{x}=$ $2.16 \%$ ), rivers (averages represent returns years 2003-2017 for the Methow and Okanogan rivers and 2011-2016 for the Wenatchee River).



Figure 11. Escapement versus percent of escapement harvested and total number harvested of natural-origin steelhead from the Wenatchee, Methow, and Okanogan rivers.

## Discussion

The Chinook Salmon and steelhead hatchery programs of the upper Columbia River contributed to treaty and non-treaty commercial fisheries in the ocean and Columbia River as well as recreational fishing. For the programs examined here, harvest rates for upper Columbia River hatchery Chinook Salmon and steelhead were generally in line with the goals of each program. Conservation and safety-net programs for spring Chinook Salmon and steelhead sustained lower multi-year average rates of harvest ( $5-26 \%$ for spring Chinook Salmon, 5-54\% for steelhead) than augmentation programs for summer and fall Chinook Salmon (53-75\%). Every hatchery program that was evaluated contributed to harvest and sometimes substantially. The magnitude of harvest generally corresponded to the status of the population: the lowest harvest occurred on the most imperiled stocks and the highest harvest occurred on the healthiest stocks. However, harvest sometimes hindered meeting broodstock collection goals, particularly during earlier years of the programs, and harvest management of endangered or threatened species could impede achieving conservation objectives.

Spawning escapement of listed species would have been higher if harvest was lower than what occurred. However, it is difficult to evaluate how harvest of hatchery-origin fish influenced population recovery without considering the factors that can influence natural production such as spawner abundance, domestication selection, and recipient stray proportions. In some years, the number of natural-origin recruits was limited by the number of spawners and any harvest likely
reduced the number of natural-recruits. In other years, the proportion of hatchery origin spawners ( pHOS ) was higher than management objectives and targeted harvest may have benefitted natural production by reducing the effects of domestication selection (e.g., steelhead in the Methow River). However, even in cases where fisheries targeted harvest augmentation programs, fisheries were not efficient enough to remove the desired number of hatchery-origin fish particularly in years of very large abundance or when weak stock fisheries limited the allowable harvest under the Endangered Species Act. Finally, higher harvest of hatchery-origin fish may have aided managers achieve targeted recipient population stray percentages (see recipient stray chapter in this report). However, most fisheries occur in areas downstream of what would be desirable locations to manage stray rates. Uncertainty remains about the effects of harvest on individual brood years and resulting viability of endangered or threatened populations of Chinook Salmon and steelhead. Mixed and weak stock fisheries in the ocean and mainstem Columbia River pose challenges to achieving conservation goals in the upper Columbia Watershed.

Abundance of all races of Chinook Salmon were limited by several factors including smolt-to-adult return survival (SAR), which has collapsed in recent years to around $1 \%$ along the entire Pacific coast (Welch 2020). Steelhead are also likely affected by this trend. While hatcheries can compensate for some of the effects of poor survival, opportunities for harvest, conservation, and recovery will be limited if SARs remain low.

## Spring Chinook Salmon

The harvest rates of spawning aggregates within the Upper Columbia River were variable which suggests that some spawning aggregates may be affected by harvest more than others. Among the spring Chinook Salmon hatchery conservation programs examined here, the Chiwawa River program had the highest percent of brood year harvested and the highest spawning escapement. Spawning escapement was sufficient to reach broodstock collection goals in most years since the population began to recover from the low numbers of the 1990's. Since the early 2000's there has been more harvest on the Chiwawa program than the Methow Basin spring Chinook Salmon conservation programs. The difference was greatest in 2009-2012 when harvest for the Methow programs, including the Twisp and Chewuch rivers, ranged between 5$22 \%$ (and were trending together) while harvest of the Chiwawa program ranged from 10-40\% over the last 10 brood years and as high as $95 \%$ in years previous. During this same period the spawning escapement for the Methow Basin programs remained consistently low (around 1,000 individuals) while escapement in the Chiwawa was generally greater, reaching a peak of almost 2,500 individuals in 2011. The combined tribal and recreational fisheries regularly harvest more than 100 adult Spring Chinook Salmon (up to $40 \%$ of escapement) from the Chiwawa program, but rarely harvest greater than 100 individuals (up to $25 \%$ of escapement) from the combined Methow River spring Chinook programs. This difference in exploitation rate may result from differences in return timing (Sorel et al. 2020), or other potential behavioral differences between Methow and Chiwawa program fish. More Chiwawa program fish may overlap with summer Chinook Salmon fisheries in the Upper Columbia if they tend to arrive later than Methow fish. Further investigation of differences between harvest of spring Chinook Salmon returning to the Wenatchee versus the Methow river basins may be useful for fisheries managers and provide insight into appropriate rates of exploitation. Furthermore, mixed stock fisheries pose challenges
to providing sustainable harvest rates for weak stocks or spawning aggregates within an Evolutionary Significant Unit (ESU).

The poor returns of upper Columbia River spring Chinook in the 1990's were apparent in the escapement numbers for the Chiwawa, Methow, Twisp, and Chewuch programs. Escapement improved by brood year 2000 and broodstock collection goals for the Chiwawa were reduced in 2009. Broodstock collection goals for the Methow Basin were reduced in 2012 following hatchery production recalculation. Since reduced broodstock collection goals were adopted, upper Columbia spring Chinook Salmon hatchery programs have typically met broodstock collection goals. Despite attempts by fishery managers to structure seasons to reduce harvest of Upper Columbia spring Chinook Salmon, harvest rates have averaged 12\% (range 9.3-13.8\%) since 2008 (Maier 2020).

Summer and fall Chinook Salmon
By design, all hatchery summer and fall Chinook Salmon programs in the upper Columbia have sustained relatively high rates of harvest compared with spring Chinook Salmon. While all anadromous salmonids in the upper Columbia declined significantly in the 1990's, the recovery of summer and fall Chinook Salmon since 2000 has led to robust fisheries, particularly in the ocean. Summer and fall Chinook Salmon in the Upper Columbia support some of the highest harvest rates in the Columbia River Basin and yet the populations continue to be relatively healthy. Upper Columbia River summer and fall Chinook Salmon tend to move north to forage after leaving the Columbia River estuary and are harvested in the Gulf of Alaska, the southeast Alaska coast, and off the coast of British Columbia including around Vancouver Island (Weitkamp 2010).

## Steelhead

In contrast with upper Columbia River Chinook Salmon, steelhead harvest is uncommon in the ocean. Because steelhead are harvested primarily in recreational fisheries in the spawning tributaries, impacts on natural-origin stocks are closely monitored and the fisheries are closed upon reaching a predetermined impact limit (e.g. 5\% of escapement, determined by local creel sampling). This also means that steelhead are not reliably available for harvest because the fisheries open only when a surplus of hatchery-origin fish are available. Escapement of hatcheryorigin steelhead in the upper Columbia River has been trending down since return year 2011 and as such, recreational fisheries have been uncommon in recent years, last occurring in return years 2015-2016 for the Wenatchee and return years 2016-2017 for the Methow. Even with decreasing escapement, broodstock collection goals have generally been met for all hatchery programs.

## Summary

In summary, PUD hatchery programs in the upper Columbia Basin have consistently provided opportunities for harvest in a variety of ocean and freshwater locations. Fall and summer Chinook Salmon were harvested at high levels and the populations continue to thrive. In contrast, relatively low but uneven harvest rates occurred on ESA listed spring Chinook Salmon and steelhead and the populations struggle to persist. Differences in population status among salmon and steelhead pose challenges to manage mixed stock fisheries in ways that
protect weak stocks, achieve harvest goals, and achieve other conservation objectives such as straying and pHOS management.

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[^0]:    ${ }^{1}$ Grant County Public Utility District
    ${ }^{2}$ BioAnalysts
    ${ }^{3}$ Chelan County Public Utility District

[^1]:    * Colors indicate varying degrees of management action: Green = no action necessary; Red = action may be necessary (See Appendix 2 in Hillman et al. 2017).

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[^3]:    ${ }^{2}$ Public Utility District No. 1 of Douglas County

[^4]:    ${ }^{1}$ Public Utility District Number 2 of Grant County, Post Office Box 878, Ephrata, Washington 98823, USA
    ${ }^{2}$ Public Utility District Number 1 of Chelan County, Post Office Box 1231, Wenatchee, Washington 98801, USA
    ${ }^{3}$ BioAnalysts, Inc., 4725 North Cloverdale Road, Suite 102, Boise, Idaho 83713, USA
    ${ }^{4}$ Public Utility District Number 1 of Douglas County, 1151 Valley Mall Parkway, East Wenatchee, Washington 98802, USA

[^5]:    ${ }_{2}{ }^{-}$escapement estimates for natural origin fish, broodstock counts for hatchery origin fish

[^6]:    ${ }^{1}$ Public Utility District No. 1 of Douglas County, East Wenatchee, WA 98802
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