

# GOPEN ACCESS

**Citation:** Crozier LG, McClure MM, Beechie T, Bograd SJ, Boughton DA, Carr M, et al. (2019) Climate vulnerability assessment for Pacific salmon and steelhead in the California Current Large Marine Ecosystem. PLoS ONE 14(7): e0217711. https://doi.org/10.1371/journal.pone.0217711

Editor: João Miguel Dias, Universidade de Aveiro, PORTUGAL

Received: February 5, 2019

Accepted: May 16, 2019

Published: July 24, 2019

**Copyright:** This is an open access article, free of all copyright, and may be freely reproduced, distributed, transmitted, modified, built upon, or otherwise used by anyone for any lawful purpose. The work is made available under the <u>Creative</u> <u>Commons CC0</u> public domain dedication.

Data Availability Statement: All relevant data are within the manuscript and its Supporting Information files.

**Funding:** ECS Federal, Inc in support of NOAA Fisheries Office of Science and Technology provided salary for author MWN, but did not have any additional role in the study design, data collection and analysis, decision to publish, or preparation of the manuscript. The specific roles of these authors are articulated in the 'author contributions' section. RESEARCH ARTICLE

# Climate vulnerability assessment for Pacific salmon and steelhead in the California Current Large Marine Ecosystem

Lisa G. Crozier<sup>1\*</sup>, Michelle M. McClure<sup>1\*</sup>, Tim Beechie<sup>1</sup>, Steven J. Bograd<sup>2</sup>, David A. Boughton<sup>3</sup>, Mark Carr<sup>4</sup>, Thomas D. Cooney<sup>1</sup>, Jason B. Dunham<sup>5</sup>, Correigh M. Greene<sup>1</sup>, Melissa A. Haltuch<sup>1</sup>, Elliott L. Hazen<sup>2</sup>, Damon M. Holzer<sup>1</sup>, David D. Huff<sup>1</sup>, Rachel C. Johnson<sup>3,6</sup>, Chris E. Jordan<sup>1</sup>, Isaac C. Kaplan<sup>1</sup>, Steven T. Lindley<sup>3</sup>, Nathan J. Mantua<sup>3</sup>, Peter B. Moyle<sup>7</sup>, James M. Myers<sup>1</sup>, Mark W. Nelson<sup>8</sup>, Brian C. Spence<sup>3</sup>, Laurie A. Weitkamp<sup>1</sup>, Thomas H. Williams<sup>3</sup>, Ellen Willis-Norton<sup>4</sup>

 Northwest Fisheries Science Center, National Marine Fisheries Service, National Oceanic and Atmospheric Administration, Seattle, Washington, United States of America, 2 Southwest Fisheries Science Center, National Marine Fisheries Service, National Oceanic and Atmospheric Administration, Monterey, California, United States of America, 3 Southwest Fisheries Science Center, National Marine Fisheries Service, National Oceanic and Atmospheric Administration, Santa Cruz, California, United States of America, 4 Department of Ecology and Evolutionary Biology, University of California, Santa Cruz, California, United States of America, 5 Forest & Rangeland Ecosystem Science Center, U.S. Geological Survey, Corvallis, Oregon, United States of America, 7 Department of Wildlife, Fish and Conservation Biology, University of California, Davis, California, Davis, California, United States of America, 8 ECS Federal, Inc. Under Contract to Office of Sustainable Fisheries, National Marine Fisheries Service, National Oceanic and Atmospheric Administration, Silver Spring, Maryland, United States of America

¤ Current address: Pacific Marine Environmental Laboratory, Seattle, Washington, United States of America

\* lisa.crozier@noaa.gov

# Abstract

Major ecological realignments are already occurring in response to climate change. To be successful, conservation strategies now need to account for geographical patterns in traits sensitive to climate change, as well as climate threats to species-level diversity. As part of an effort to provide such information, we conducted a climate vulnerability assessment that included all anadromous Pacific salmon and steelhead (Oncorhynchus spp.) population units listed under the U.S. Endangered Species Act. Using an expert-based scoring system, we ranked 20 attributes for the 28 listed units and 5 additional units. Attributes captured biological sensitivity, or the strength of linkages between each listing unit and the present climate; climate exposure, or the magnitude of projected change in local environmental conditions; and adaptive capacity, or the ability to modify phenotypes to cope with new climatic conditions. Each listing unit was then assigned one of four vulnerability categories. Units ranked most vulnerable overall were Chinook (O. tshawytscha) in the California Central Valley, coho (O. kisutch) in California and southern Oregon, sockeye (O. nerka) in the Snake River Basin, and spring-run Chinook in the interior Columbia and Willamette River Basins. We identified units with similar vulnerability profiles using a hierarchical cluster analysis. Life history characteristics, especially freshwater and estuary residence times, interplayed with gradations in exposure from south to north and from coastal to interior regions to **Competing interests:** MWN is employed by a commercial contracting company, ECS Federal, Inc in support of NOAA Fisheries Office of Science and Technology. This does not alter our adherence to PLOS ONE policies on sharing data and materials.

generate landscape-level patterns within each species. Nearly all listing units faced high exposures to projected increases in stream temperature, sea surface temperature, and ocean acidification, but other aspects of exposure peaked in particular regions. Anthropogenic factors, especially migration barriers, habitat degradation, and hatchery influence, have reduced the adaptive capacity of most steelhead and salmon populations. Enhancing adaptive capacity is essential to mitigate for the increasing threat of climate change. Collectively, these results provide a framework to support recovery planning that considers climate impacts on the majority of West Coast anadromous salmonids.

# Introduction

Anthropogenic climate change poses a direct threat to existing global biodiversity. In fact, climate-related population extinctions have already occurred in 47% of 976 plant and animal species surveyed in a recent review of the literature [1]. Moreover, local extinction percentages are higher in freshwater (74%) than in terrestrial (46%) or marine habitats (51%) [1]. Such impacts are expected to increase in the future [2–4], and managers are actively seeking information regarding the species or populations most vulnerable to climate change. Information of this kind is needed to prioritize resources for restoration and climate adaptation efforts. Climate vulnerability assessments are an important tool in these efforts because they provide systematic summaries of the relative threat level to a set of species or populations [5-7].

We conducted a comprehensive climate vulnerability assessment for Pacific salmon and steelhead (*Oncorhynchus* spp.) in the U.S. portion of the California Current Large Marine Ecosystem (CCLME) and associated watersheds. Partly as a consequence of natal homing to diverse watersheds, Pacific salmon display significant life history diversity evolved through local adaptation and limited dispersal [8]. In considering the conservation importance of this diversity, NOAA Fisheries applied the concept of evolutionarily significant units [9] to define 52 distinct population segments (DPSs) of Pacific salmon that could potentially be protected under the US Endangered Species Act (ESA). Our analysis focuses primarily on those DPSs that have been identified as species of concern, threatened or endangered (31/52). We also included one chum (*O. keta*) and one pink (*O. gorbuscha*) non-listed DPS to represent these species, which have few or no listed DPSs. In total we compared the relative vulnerability of 33 *Oncorhynchus* DPSs in the CCLME.

Our assessment was based on three components of vulnerability: 1) biological sensitivity, which is a function of individual species characteristics; 2) climate exposure, which is a function of geographical location and projected future climate conditions; and 3) adaptive capacity, which describes the ability of a DPS to adapt to rapidly changing environmental conditions [10]. Objectives were to characterize the relative degree of threat posed by each component of vulnerability across DPSs and to describe landscape-level patterns in specific threats and cumulative vulnerability at the DPS level.

#### Species units, spatial domains, and life histories

Pacific salmon are native to coastal regions of northeastern Asia (Japan, Korea and Russia) and western North America from California to Alaska. Of the seven species of *Oncorhynchus* [11] within the CCLME, we included the six that have primarily anadromous life histories: climate change will profoundly impact both the freshwater and marine life stages for these species

(Table 1). A seventh species, cutthroat trout (*O. clarkii*) has an anadromous component, but is generally considered an inland species. Among the six species included in our analysis, there are 52 DPSs occupying eight recovery domains (Fig 1), or ecoregions with distinct climatic and ecological characteristics.

Table 1. Salmon and steelhead distinct population segments (DPSs) included in the assessment. Species names are shown with number of DPSs in parenthesis. Name, listing status, and recovery domain is also shown for each DPS.

Sp	pecies/distinct population segment name	Listing status	Recovery domain	
Steelhead O. mykiss (11)				
	Northern California steelhead	Threatened	North-Central California Coast	
	California Central Valley steelhead	Threatened	Central Valley	
	Central California Coast steelhead	Threatened	North-Central California Coast	
	South-Central California Coast steelhead	Threatened	S-Central/Southern CA Coast	
	Southern California Coast steelhead	Endangered	S-Central/Southern CA Coast	
	Puget Sound steelhead	Threatened	Puget Sound	
	Upper Columbia River steelhead	Threatened	Interior Columbia	
	Snake River Basin steelhead	Threatened	Interior Columbia	
	Middle Columbia River steelhead	Threatened	Interior Columbia	
	Upper Willamette River steelhead	Threatened	Willamette/-Lower Columbia	
	Lower Columbia River steelhead	Threatened	Willamette/-Lower Columbia	
C	hinook salmon O. <i>tshawytscha</i> (11)			
	Lower Columbia River Chinook	Threatened	Willamette/-Lower Columbia	
	Upper Willamette River Chinook	Threatened	Willamette/-Lower Columbia	
	Puget Sound Chinook	Threatened	Puget Sound	
	Snake River fall-run Chinook	Threatened	Interior Columbia	
	Snake River spring/summer-run Chinook	Threatened	Interior Columbia	
	Middle Columbia River spring-run Chinook	Sensitive*	Interior Columbia	
	Upper Columbia River spring-run Chinook	Endangered	Interior Columbia	
	Central Valley fall/late fall-run Chinook	Species of concern	Central Valley	
	Central Valley spring-run Chinook	Threatened	Central Valley	
	Sacramento River winter-run Chinook	Endangered	Central Valley	
	California Coastal Chinook	Threatened	North-Central California Coast	
C	oho salmon O. kisutch (5)			
	Central California Coast coho	Endangered	North-Central California Coast	
	Southern Oregon/Northern California Coast coho	Threatened	Southern Oregon/Northern CA Coast	
	Oregon Coast coho	Threatened	Oregon Coast	
	Lower Columbia River coho	Threatened	Willamette/Lower Columbia	
	Puget Sound coho	Species of concern	Puget Sound	
C	hum salmon O. keta (3)			
	Columbia River chum	Threatened	Willamette/Lower Columbia	
	Puget Sound chum	Not listed	Puget Sound	
	Hood Canal summer-run chum	Threatened	Puget Sound	
Sc	ockeye salmon O. nerka (2)			
	Lake Ozette sockeye	Threatened	Puget Sound	
	Snake River sockeye	Endangered	Interior Columbia	
Pi	nk salmon O. gorbuscha (1)			
	Odd-year pink	Not listed	Puget Sound	

\* Middle Columbia spring-run Chinook are identified as sensitive by Oregon

https://doi.org/10.1371/journal.pone.0217711.t001



**Fig 1. Salmon recovery domains.** We analyzed patterns in vulnerability across DPSs within geographic recovery domains used to manage West Coast salmon and steelhead populations listed under the ESA [12]. The DPSs inhabiting each domain are listed in <u>Table 1</u>.

https://doi.org/10.1371/journal.pone.0217711.g001

At present, more than half of all anadromous Pacific salmon and steelhead DPSs remaining in the contiguous U.S. are threatened with extinction [13]. Suboptimal climate conditions within the historical range of climate variability have been associated with detectable declines in many of these DPSs, highlighting their sensitivities to climatic drivers [14–17]. In some cases, the synergistic effects of suboptimal climate conditions and intense anthropogenic stressors precipitated the population declines that led to these listing decisions.

There is tremendous life history diversity among and within Pacific salmon species (Fig.2) [18, 19]. Anadromous species hatch in freshwater, migrate to the ocean to feed and grow, and return to freshwater to spawn. Most adults die after spawning, although some steelhead (O. *mykiss*) spawn successfully in multiple years. Juveniles can remain in freshwater anywhere from days to years, with populations that spawn near the ocean typically having shorter freshwater phases [20].

The seasonal timing of the juvenile and adult migrations varies across DPSs and species, as does the extent to which this variation is associated with genetic differentiation [21, 22]. Within the CCLME, Chinook salmon (*O. tshawytscha*) and steelhead exhibit the greatest life-stage variability. For example, some Chinook juveniles spend a full year in freshwater before migrating as yearlings, whereas others enter the marine environment as subyearlings. Adults of different life history types enter freshwater to commence the spawning migration in spring, summer, fall, or winter, with maturation either in the ocean or in freshwater.

Salmon life histories are highly variable within the marine stage as well. In the CCLME, pink salmon (*O. gorbuscha*) characteristically spend 1.5 years at sea, while coho (*O. kisutch*), chum (*O. keta*), Chinook, sockeye (*O. nerka*), and steelhead mature at various ages, with some males and hatchery offspring returning to freshwater within 1 year. Typically, adult coho return after 1.5 years at sea, whereas the other salmon species spend 2-5 years in the ocean. *O. mykiss, O. nerka*, and to a lesser extent other species have some populations or portions of populations that forego the marine migration altogether. Freshwater-resident populations, most notably rainbow trout (*O. mykiss*) and kokanee (*O. nerka*) are generally not included in DPSs.

Anadromous salmonids exhibit a high degree of homing fidelity during the adult migration, which fosters local adaptation to conditions in a particular watershed. Differences in behavior, body shape, thermal tolerance, and disease tolerance reflect genetic adaptations to characteristic patterns in temperature and stream flow (e.g., [23-26]). Differences in environmental cues, habitat conditions, and growth rates also trigger extensive life history variation among and within populations, even in the absence of genetic differentiation (e.g., [27-29]). Many traits appear to have responded to recent climate change, apparently without genetic adaptation [30]. However, to keep pace with climate change, genetic adaptation may be necessary in the long-run [31–33]; thus, maintaining genetic diversity within DPSs and species as a whole is a high priority for salmon conservation [34].

# Methods

Our approach followed the climate vulnerability assessment method developed by Hare et al. [35], which is now being implemented for U.S. marine and anadromous species by NOAA Fisheries [36]. This method was designed for rapid assessment across a wide variety of taxa using available qualitative and quantitative data. It assumes that vulnerability will be periodically re-assessed, and methods refined as status reviews are updated and more information can be considered for individual DPSs.

Hare et al. [35] had four components in their analysis: exposure, sensitivity, probability of directional shift, and net direction of climate effects. They used exposure and sensitivity attributes to calculate total vulnerability, while range shift and net climate effect scores provided

# **PLOS** ONE



Fig 2. Schematic of Pacific salmon life histories for example ecotypes. Colors represent life stages, where yellow indicates adult freshwater migration and holding, red spawning, orange incubation, light blue juvenile freshwater rearing and migration, green estuary and nearshore rearing, and dark blue marine stage.

https://doi.org/10.1371/journal.pone.0217711.g002

supplemental information. Hare et al. [35] intentionally incorporated adaptive capacity attributes into the sensitivity component. Nonetheless, they emphasized specific elements of adaptive capacity that had special relevance by reporting separate scores for range shift and net climate effect.

We used the same analytical structure as Hare et al. [35], but with specific attributes tailored to account for characteristics most relevant to individual life stages and habitats used by salmon and steelhead. This approach allowed us to capture within-species differences at the DPS level, or scale of management units presently used in salmon recovery planning.

We differentiated DPSs by exposure and sensitivity metrics applied to each life stage. Exposure attributes summarized the magnitude of change expected in climate variables with the potential to affect species productivity in a specific region. Sensitivity attributes were based on life history characteristics and proximity to climate thresholds (i.e., sensitivity to climate limits was not assumed to be linear), as well as attributes associated with adaptive capacity, such as population size and artificial breeding programs. We also developed a separate attribute for adaptive capacity that focused on the perceived likelihood of a phenotypic change that could mitigate the effects of climate change. This separate adaptive capacity score was not included in vulnerability ranks but provided additional information useful for conservation planning.

In the following sections, we describe the specific DPSs included in this assessment and the attributes evaluated to determine relative vulnerability. We then detail the process of collecting information on attributes, scoring each attribute, and ranking DPSs into low, moderate, high, and very high vulnerability categories. Finally, we describe additional analyses performed to identify key drivers of climate vulnerability and their likely consequences for species-level diversity if the most vulnerable DPSs are lost.

# Spatial and biological scope

Our assessment included all 28 ESA-listed DPSs of Pacific salmon and steelhead in seven recovery domains (<u>Table 1</u>). Five of these DPSs are listed as endangered and 23 listed as threatened [<u>14</u>, <u>16</u>]. Two additional Chinook and one coho salmon DPSs are considered species of concern by NOAA or sensitive species by state agencies. We also included Puget Sound pink salmon, which combines even-year and odd-year DPSs, because no pink salmon are listed under the ESA. We also analyzed Puget Sound chum salmon because this species has only 2 listed DPSs. Most remaining non-listed DPSs either lacked specific information that could cause their score to differ from that of a neighboring DPS (e.g., Washington Coast Chinook salmon), or were hatchery-dominated to an extent that the effects of climate change will depend more on hatchery management than on the attributes included in our analysis. While hatchery management will also need to adjust to climate change [<u>37</u>], different metrics than those evaluated here are needed to characterize vulnerability in hatchery stocks.

# Sensitivity attributes

Attributes describing biological sensitivity to climate change included life-stage specific metrics that largely reflect the intrinsic biological characteristics and geographic range of each DPS. These attributes also included population-level stressors that reduce natural resilience. All biological sensitivity attributes except *sensitivity to ocean acidification* were modified from species-level assessments [35], which would have produced the same score for all DPSs. Note that we included *exposure to ocean acidification* as a separate attribute to characterize the amount of physical change expected in the CCLME. Salmon generally occupy tributary, mainstem, estuary, and marine habitats sequentially over their life cycle. Therefore, in assessing climate vulnerability, it is important to consider sensitivity at each life stage and corresponding habitat. To standardize scoring across DPSs, we developed a rubric for each sensitivity attribute (Table 2, S1 Appendix).

Life-stage sensitivity. We calculated habitat-specific sensitivity scores associated with five stages of the salmon life cycle. These stages are seasonally and spatially defined, so the particular habitats occupied in each life stage are potentially affected by different exposure attributes. The *early life history* stage included egg incubation and fry emergence; the *juvenile freshwater stage* encompassed the fry-to-smolt transition; and the *estuary* and *marine stages* were distinguished physically by location. The *adult freshwater stage* included freshwater entry, migration, holding, and spawning.

For each life stage, biological sensitivity was scored from low to very high based on the extent of present climate stress within the DPS habitat ( $\underline{S1}-\underline{S8}$  Figs) and on the level of habitat and behavioral diversity within the DPS. For a given life stage and DPS, sensitivity was ranked very high if high mortality had been directly linked to a climate driver in recent history. For example, Sacramento River winter-run Chinook recently experienced high rates of egg

**Table 2. Overview of sensitivity and exposure attributes.** We developed a rubric for each sensitivity and exposure attribute to standardize scoring across DPSs. We included four freshwater and five marine exposure attributes, each considered within the habitat of the respective DPS and life stage. Full descriptions of scoring criteria are included in the <u>S1</u> and <u>S2</u> Appendices.

Attributes	Low vulnerability	Very high vulnerability			
	Sensitivity				
Early life history	Minimal flow & temperature stress in egg/ early fry stage	Flow or temperature stress already apparent			
Juvenile freshwater stage	Flexible subyearling migration strategy	Constrained yearling or stressed subyearling migratory strategy			
Estuary stage	Short estuarine residence or wide window for migration timing	Long estuary residence or climate-related threats in the estuary stage already apparent			
Marine stage Low correlation between marine survival & climate indicators, overlapping cohorts with variable age at return		High correlation between marine survival & climate indicators; simple age structure			
Adult freshwater stage	Adult migration distance & duration short; low climate stress during migration, holding & spawning	Adults encounter peak summer temperatures or flow constraints during migration, holding, or spawning			
Cumulative life- cycle effects	Low risk of loss for defining characteristic of DPS or link between life stages	Imminent climate threshold or life history type already at risk			
Hatchery influence	No hatchery-origin populations released within DPS boundaries	Production hatcheries dominate naturally spawning populations			
Other stressors	Non-climate threats are relatively minor	Multiple threat categories severe relative to other DPSs			
Population viability	Extinction risk low based on viable salmon population criteria	Extinction risk high based on viable salmon population criteria			
Ocean acidification sensitivity	Non-specialist on prey highly sensitive to ocean acidification	The DPS is a sensitive taxon, see text			
	Freshwater exposure				
Stream temperature	Z-score in August mean stream temperature of spawning, rearing, and migration habitats < 0.5	Z-score for August mean stream temperature exceeds 2			
Summer water deficit	Z-score for water balance in summer freshwater habitat $< 0.5$	Z-score for water deficit exceeds 2			
Flooding	Relatively small change projected or freshwater habitat not influenced by floods	Large change in flood events with potentially severe habitat effects expected			
Hydrologic regime	Expected regime change in < 5% of spawning area	Expected regime change in $> 25\%$ of spawning area			
	Marine exposure				
Sea level rise	Sea level rise minimal (projection range includes 0)	Sea level rise > global average			
Sea surface temperature	Z-score in the ocean migration area $<0.5$	Z-score in the ocean habitat exceeds 2			
Ocean acidification exposure	Z-score for pH in ocean range < 0.5	Z-score for ocean pH exceeds 2			
Upwelling	Little projected change in intensity or phenology of upwelling-favorable winds	Significant projected change in intensity or phenology of upwelling-favorable winds			
Ocean currents	Large-scale ocean circulation patterns affecting the northern CCLME are projected to change relatively little	Major changes in ocean circulation are projected			

https://doi.org/10.1371/journal.pone.0217711.t002

mortality due to warm water temperatures [38]. If behavioral or habitat diversity allows a substantial portion of the population to avoid detrimental conditions in a given year, for example by shifted phenology or habitat selection, then sensitivity was ranked lower. We also included an attribute for *cumulative life-cycle effects* to reflect the necessity of completing all stages and maintaining a life history pattern characteristic of the DPS. This attribute accounted for the possibility that individuals might avoid a climate stressor during a given life stage at a cost to subsequent stages. For example, earlier migration in the *juvenile freshwater stage* could increase survival to ocean entry but decrease survival during the *marine stage* because of smaller body size or a mismatch between prey abundance and ocean-entry timing. The *cumulative life-cycle effects* attribute also captured any expert judgment that a given life stage was at such critical risk that reduction in survival at that stage would threaten the entire life cycle or an essential characteristic of the DPS (e.g., anadromy).

**Sensitivity to ocean acidification.** Salmon *sensitivity to ocean acidification* most likely occurs through ecological mechanisms mediated by changes to the food web [<u>39–41</u>]. Taxa directly affected by declining marine pH include invertebrates such as pteropods, crabs, and krill, which play a significant role in some salmon diets [<u>42</u>]. Physiological effects of acidification may also impair olfaction, which could hinder homing ability [<u>43</u>], along with other developmental effects [<u>44</u>]. Using the criteria of Morrison et al. [<u>45</u>] for scoring, all salmon had low-to-moderate *sensitivity to ocean acidification*. Slight differences among DPSs stemmed from marine diet: sockeye, chum, and pink salmon consume more zooplankton than Chinook, coho and steelhead, which are mostly piscivorous.

**Population viability.** Scores for population viability were based on indices of extinction risk, as evaluated in recent ESA status reviews and viability assessments [14, 16]. As part of each status review, all listed salmon were formally evaluated with respect to 1) present vs. historical population abundance; 2) population growth rate; 3) spatial structure, or the distribution of populations within a DPS; and 4) genetic and phenotypic diversity [34]. For DPSs not included in status reviews, we asked experts to apply these same criteria to the greatest extent possible, given the information available. These population viability criteria were developed by NOAA Fisheries to monitor long-term evolutionary potential [34], and therefore they relate to adaptive capacity. More specifically, evolutionary potential is strongly related to genetic variability and the risk of demographic extinction, both of which are correlated with population size [33, 46] and growth rate [47].

Hatchery influence. Numerous hatcheries release artificially propagated juvenile salmon into freshwater, estuary, or marine habitats to supplement natural production. After completing the ocean stage, these hatchery-origin fish generally return to tributaries concurrently with natural-origin salmon. Unless they are harvested or collected for broodstock or removal, hatchery-origin fish spawn in natural habitat.

Hatcheries may have mixed effects on the resilience of natural populations to climate change. In the best-case scenario, hatcheries provide a temporary demographic buffer for catastrophic declines in abundance [48]. However, hatchery populations could eventually be more susceptible to large-scale climate forcing than natural populations due to the absence of behavioral, physiological, and genetic adaptation in the wild [15, 49]. Although some hatcheries follow careful genetic protocols to minimize loss of genetic variation, many reduce the effective size of wild populations in proportion to their relative abundance [50–52], which reduces adaptive capacity. In this assessment, we assumed that conservation hatcheries practicing bestmanagement procedures and high-quality monitoring posed lower risks to DPSs than production hatcheries. Thus we ranked no hatchery influence as low vulnerability, influence from conservation hatcheries as moderate, and influence from production hatcheries as high or very high, depending on the proportion of natural-origin adults spawning in streams across the DPS.

**Other stressors.** Salmon populations are affected by numerous stressors not directly related to climate but that potentially reduce their ability to cope with climate change. The

most common of these are habitat loss, habitat degradation, toxic chemicals, pathogens endemic to fish culture, displacement by invasive species through competition and predation, and harvest. All DPSs have experienced habitat loss and degradation along with recent changes in ecosystem composition. The highest scores for this attribute were reserved for the most severe cases, in which DPSs were subjected to a combination of multiple stressors.

Primary factors leading to past declines of wild salmon populations have included migration barriers, overfishing, habitat loss and degradation, and negative effects from hatchery production, which are captured in the *hatchery influence* attribute [53]. Although some of these stressors are now less severe than in the past (especially overfishing), they continue to affect population status in all DPSs, and are often exacerbated by climate stressors [54, 55]. We refer to *population viability, hatchery influence*, and *other stressors* as *extrinsic* factors, because they are imposed extrinsically by human activity.

#### **Exposure attributes**

Climate exposure attributes were used to describe the magnitude of projected change in the physical environment by mid-century. Projected climate change was based on the "business-as-usual" trajectory of greenhouse gas emissions, relative concentration pathway 8.5 [2]. It is important to note that our scores for climate exposure reflected physical change relative to a historical reference period but did not assess whether these conditions were stressful for salmon. Thus, some locations that are already extremely arid or hot may be only marginally suitable for salmon, yet did not score high in the exposure component if they were not expected to change much. Proximity to environmental thresholds was captured in the biological sensitivity rather than the exposure attributes.

We included four freshwater and five marine exposure attributes (<u>Table 2</u>), and each attribute was considered within the habitat of the respective DPS and life stage. For freshwater attributes, we quantified the amount of change projected to occur in the spawning and rearing habitat and in migration corridors delineated for each DPS in the *StreamNet* data repository [<u>56</u>] (<u>S1–S8</u> Figs, <u>S1 Table</u>). Scores for marine exposure considered the ocean migration patterns of the respective DPSs (e.g., [<u>57–59</u>]).

**Freshwater attributes.** Temperature and flow patterns affect all aspects of salmon behavior and physiology in freshwater, often with consequences for the marine life stage as well. Freshwater exposure attributes and scoring criteria are summarized in <u>Table 2</u>, and the specific rationale for each exposure metric is detailed in the <u>S2 Appendix</u>.

Briefly, we summarized temperature change by focusing on summer, when lethal temperatures often occur (14–25°C, depending on life stage) [<u>60–62</u>]. Summer low flows and drought reduce available wetted habitat and can sever connections between habitat areas, causing mortality from stranding; during these periods, water quality in remaining habitats is reduced. Low summer flows and warmer temperatures often work together in altering prey composition, riparian vegetation, and stream morphology.

Conversely, high flows can have positive or negative effects, depending on life stage, season, and watershed characteristics such as connected vs. disconnected floodplains and side channels. For example, migrating smolts generally benefit from higher flows, whereas eggs and fry exposed to higher flows can be scoured from their nests, inundated by sedimentation, or flushed out of preferred habitat especially in areas where floodplain connections have been lost or impaired. Maximum flows can result either from large precipitation events, melting of accumulated snow, or a combination of both.

We selected four metrics to capture projected change in these environmental drivers: August mean *stream temperature*, mean *summer water deficit*, extreme precipitation or *flooding* events, and change in *hydrologic regime*, which is determined by the ratio of rain to snow in winter precipitation.

To describe change in summer stream conditions, we used modeled *stream temperatures* [63, 64] and the evapotranspiration differential (potential minus actual), also known as the *summer water deficit* [65]. The latter attribute served as a proxy for low flows and drought stress on riparian vegetation. For both attributes, conditions within the spawning and rearing habitat projected for 2030–2059 were standardized statistically (*z*-score transformed) using means and standard deviations of the reference periods (1993–2011 for *stream temperature* and 1916–2006 for *summer water deficit*). The reference period for *stream temperature* is much more recent than the other metrics simply because long-term historical records are extremely rare. However, given the strong correlation between stream and air temperature [63] and the trends in longer historical records of air temperature, a longer reference period would only have increased the projected change. Because the standardized rates of change for this metric were often in our highest category (z > 2), a longer reference period would likely have had a minimal impact on our conclusions.

Changes in the magnitude of peak flows have been modeled directly for most northern streams [66], but were evaluated by proxy in California and coastal Oregon based on changes in the frequency and intensity of heavy rain events [67]. In western North America, the most extreme rain events stem from narrow corridors of water vapor called atmospheric rivers, which carry moisture over thousands of kilometers of ocean from the tropical mid-Pacific. We focused on changes in extreme events, represented by the 99th percentile in precipitation or *flooding*. Both analyses used 1970–1999 as the historical period, but the projection period for atmospheric rivers was 2070–2099, whereas the projection period for *flooding* was 2040–2069. Changes in extreme events were not amenable to *z*-score transformation; hence peak flow exposures were left as a raw percent change for experts to rank from low to very high.

*Hydrologic regime* reflects the annual pattern of flows and whether they are primarily driven by rainfall, snowmelt, or groundwater. This attribute was designed to provide a holistic description of the watershed characteristics most often correlated with salmon life history traits, and hence those directly relevant to potential loss of diversity [68, 69]. Peak flows occur during fall or winter in rain-dominated basins and during spring or early summer in snowdominated basins. Groundwater-dominated basins are relatively insensitive to either rainfall or snowmelt.

As temperatures warm, the seasonal transition from rainfall to snowfall begins later in the year, producing higher flows in early winter and shrinking cumulative snowpack. Spring/summer snowmelt is also expected to begin earlier in most basins, causing earlier and smaller spring freshets with lower minimum flows in late summer. We characterized projected change in these flow characteristics by quantifying *hydrologic regime* change in areas within and upstream from spawning and rearing habitat. For scoring, we used threshold criteria defined by Hamlet et al. [70] and most recently modeled by Littell et al. [65]. Any change from snow-dominant to transitional or from transitional to rain-dominant regimes increased the *hydrologic regime* score.

**Marine attributes.** Ocean conditions are a major driver of salmon abundance. Marine survival tends to be correlated across stocks and species in the northeast Pacific, generally following patterns in *sea surface temperature* (SST) and large-scale climate indices [71–73]. Ocean distributions of salmon species are strongly correlated with SST [74–76], and various climate indices associated with salmon survival are related to or derived from this attribute. These include the Pacific Decadal Oscillation [77], North Pacific Gyre Oscillation, and various El Niño-Southern Oscillation indices [78]. Each of these indices reflect large-scale patterns of variation in multiple ocean characteristics such as horizontal currents, upper ocean

temperature and stratification, upwelling, and vertical mixing between deep and surface waters [79-81].

In contrast, future warming trends in the north Pacific Ocean are projected to be dominated by thermal forcing associated with increased greenhouse gas concentrations and the thermodynamic feedbacks they trigger [82]. Furthermore, the relative importance of largescale climate indices for salmon tends to change over time [78], making it difficult to determine which index will be most applicable in the future. For these reasons, we elected to focus on SST itself as the exposure factor.

In addition to climate indices, historical variations in west coast salmon marine survival have been associated with the strength of *ocean currents* that alter the proportion of prey from cold, subarctic waters [71, 72, 83–87]. *Upwelling* also impacts salmon prey composition and is a defining feature of the CCLME; thus, the strength and timing of upwelling-favorable winds was included as an exposure attribute. Upwelling intensity is tightly correlated with input and retention of cold, nutrient-rich waters to the euphotic zone, which promote high levels of primary productivity and a lipid-rich food-web in the CCLME [81, 88, 89].

Sea level rise was included as an exposure attribute because many salmon rear in estuaries for months before they complete the transition to marine life stages. For these fish, transitional estuarine rearing periods strongly influence later survival. Sea level rise will alter estuary and nearshore habitats, likely intensifying the impact of high tides, storms, and floods [90]. Sea level rise will also alter estuarine hydrodynamics, with additional implications for salmon habitat quality and abundance [91]. Sea level rise is associated with a net loss of estuary habitat for juvenile salmon in some assessments; [92] however, estuary dynamics are complicated, especially in terms of sand-bar formation and breaching, and we lacked detailed models with which to project these processes. We therefore differentiated DPSs by their relative rates of sea level rise at ocean entry, assuming a higher rate was more detrimental.

Finally, pH levels in the CCLME have been declining, resulting in reduced abundance and increased corrosion in the shells of calcifying organisms [93–95]. Negative effects of lower pH have been shown for many taxa in the CCLME [41], although the cumulative effects of *exposure to ocean acidification* on salmon are still uncertain.

We examined a total of five attributes reflecting ocean conditions: *sea surface temperature* (SST), *ocean acidification* (OA), *sea level rise*, timing and intensity of *upwelling*, and change in *ocean currents*. For SST and OA, we calculated standardized change in grid cells of 1° latitude by 1° longitude based on output from 27 (SST) and 11 (OA) earth system models downloaded from the NOAA Ocean Climate Change Web Portal [96]. In each grid cell, we calculated the magnitude of change as the difference between mean climate projected for 2006-2055 and mean climate from historical simulations during 1955–2005. We normalized the projected change for these exposure attributes by dividing by the historical standard deviation (*z*-score), then taking the average of *z*-scores across grid cells within the migratory range of the DPS. We calculated the mean magnitude of change at both annual and seasonal time steps to account for seasonal variation. Scorers determined the most relevant season and location for individual DPSs.

For exposure to *sea level rise*, we based scores primarily on analyses conducted by the National Research Council [97] on sea levels projected for the U.S. West Coast in the 2050s. This report projected the highest rates of sea level rise at latitudes south of Cape Mendocino, California, with slower increases at higher latitudes (scoring bins detailed in the <u>S2 Appendix</u>).

Projections of change in the timing and intensity of *upwelling* constitute an active area of research, but consensus across studies is weak. Our *upwelling* scores relied primarily on the analyses of Rykaczewski et al. [98]. They compiled output from 21 GCMs for the period 2071-2100 under the representative concentration pathway 8.5 scenario. They then compared

projected oceanic and atmospheric metrics to those from the early industrial period of 1861-1890. Their results can be summarized as a slight "poleward shift" in the seasonal climatological cycle, wherein the average intensity of upwelling increases in the northern and decreases in the central and southern CCLME, and upwelling begins earlier in the year from central California through central Oregon.

Our <u>S2 Appendix</u>, *Exposure attributes*, describes the present state of the literature regarding potential change in *ocean currents* [99–102]. Ultimately, net projections were considered highly uncertain. However, our scoring method explicitly accounted for this type of uncertainty, as explained below (*Scoring process*). Reference and projection periods varied for different exposure metrics, depending on the available information. Experts used qualitative judgements ranging from low to very high to account for these differences.

#### Adaptive capacity

The Intergovernmental Panel on Climate Change defined adaptive capacity as the potential for a system to respond to environmental change by genetic adaptation or by a non-genetic, phenotypic change that mitigates negative environmental impacts (Working Group II Report 2, Table 18.5 in [2]). Adaptive capacity can be characterized in various ways, including genetic richness, life history plasticity, and dispersal ability [6, 103, 104]. Additional work is needed to explore the consequences of different methods used to characterize adaptive capacity. Although differing methods can produce different rank orders in vulnerability, there is no consensus on which methodological approach can best predict responses to climate change [105–107].

Several aspects of adaptive capacity were included in our sensitivity attributes. High scores in extrinsic factors reflected lower available levels of genetic and habitat diversity to cope with climate change [108]. For example, genetic variation is reduced in small populations, simplified habitats, and populations heavily influenced by hatcheries [109, 110]. Furthermore, fish altered by artificial selection in breeding programs may introduce maladaptive genotypes into wild populations, and these may potentially swamp genotypes that have evolved through natural selection [111]. Thus the attributes of *other stressors, population status*, and *hatchery influence* were intended as proxies for evolutionary potential to some extent. Furthermore, the *life cycle complexity* score addressed the likelihood that a present life history would continue to be viable in future climates, and thus whether phenotypic change would be needed to cope with climate change.

In defining a separate attribute for *adaptive capacity*, we sought an index of whether change in a phenotypic trait was considered likely. For example, if a given life history trait appeared optimal in a future climate, did scorers believe the DPS was likely to change adaptively toward this trait? For this attribute, we included behavioral, physiological, and morphological traits. It was not possible to quantify the extent to which change in relevant traits would result from plastic vs. evolutionary processes because many traits were both highly plastic and heritable. Thus, an initial *adaptive capacity* trait response would likely be plastic but would be subject to selection over time.

Rates of evolutionary response depend on the full genetic architecture of all traits under selection, especially their correlation structure, temporal pattern of the selection gradient at different life stages, and existing genetic variation within the optimal phenotype [47, 112, 113]. None of these elements are known at present, making adaptive capacity scores more subjective than those of other attributes. Nonetheless, changes in key traits have important management implications, especially those that define characteristics of a DPS, such as smolt or adult runtiming, anadromy, or spatial distribution. We therefore asked scorers to evaluate the likelihood

that a trait alteration could mitigate negative effects of climate change and to allocate four tallies to low, moderate, or high bins.

Movement or dispersal is an important component of adaptive capacity [104]. We focused on shifts in range or habitat usage within existing geographic boundaries and accessible habitats, although other types of range shift are possible. Salmon DPSs are defined in terms of their watershed boundaries, so dispersal of a DPS outside its existing freshwater domain would likely involve colonization of habitat occupied by another DPS, and we did not address this possibility. Shifts in salmon marine distributions have been projected based on associations with SST [74–76]; however, these projections are not available at the DPS scale. Moreover, they are based largely on Canadian and Alaskan salmon, which have migratory constraints that differ from those of DPSs included in our assessment. For these reasons, we did not attempt to quantify marine range shifts for the *adaptive capacity* attribute.

Overall, the *adaptive capacity* score was intended to capture perceived potential for behavioral, physiological, or other adaptive response to ameliorate climate stress. We assumed that experts would be familiar with a range of possible responses based on their knowledge of diversity across DPSs. Adaptive capacity scores spanned three levels (low, moderate, or high). If adaptation in a critically threatened life stage was deemed unlikely, the DPS received a low score. A moderate score indicated that some adaptive response might occur, although not in the most sensitive or exposed life stage, or that its magnitude might be fairly small. A high score indicated that some adaptive shift was likely in response to climate change. These scores were not formally integrated into relative vulnerability rankings; they provide additional information to help develop management strategies that support a range of life history expressions.

#### Data quality

A crucial component of any vulnerability assessment is the quality and specificity of information on which it is based. We characterized data quality for each sensitivity and exposure attribute based on the type of data used. We scored each data quality attribute from 0, representing qualitative expert judgement alone, to 3, representing quantitative studies focused on the specific DPS being evaluated. When quantitative studies were available, data-quality scores reflected the breadth of analyses synthesized, for example, the number of GCMs included in an ensemble projection or the number of studies documenting a given relationship, as well as the extent of agreement across studies. A score of 3 indicated broad agreement over a relatively large number of GCMs or studies focused specifically on the DPS region or on populations within the DPS [45].

#### Scoring process

We collected information on exposure attributes for the entire CCLME and associated watersheds and conducted a scoring workshop wherein experts discussed data-quality scores for each exposure attribute. For each sensitivity attribute, *profilers*, or scientists familiar with an individual DPS, wrote a description of each life stage, the seasonality, duration and known climate stressors at that stage, and variability within that life stage across the DPS. Behavior and habitat information has been summarized for each DPS in the NOAA Fisheries biological status reviews and their respective 5-year updates. However, this information is often focused on particular populations, tributaries, or time periods, and therefore may not necessarily represent the entire DPS [<u>114</u>]. Additional literature was cited in many of the DPS profiles, and profilers also assigned a data-quality score for each sensitivity attribute.

Once all of the required information was collated, a separate panel of 16 expert *scorers* rated all freshwater and marine exposure, biological sensitivity, and adaptive capacity attributes

based on the guidelines summarized in Table 2 and detailed in the <u>S1</u> and <u>S2</u> Appendices. Each DPS was scored by four experts, with each expert scoring 5–22 salmon and steelhead DPSs, plus other species included in the West Coast Fish Climate Vulnerability Assessment. To ensure consistency across groups of DPSs, scoring groups were rearranged over several sessions. Each expert independently scored their assigned DPSs based on information contained in the profiles as well as their general knowledge, using the pre-defined scoring bins shown in <u>Table 2</u> and detailed in the <u>S1</u> and <u>S2</u> Appendices.

Each scorer allocated five tallies across four bins (low, moderate, high, or very high) for each sensitivity and exposure attribute as described in Morrison et al. [45]. Adaptive capacity was scored by allocating four tallies across three bins (low, moderate, or high). Following preliminary scoring, all experts participated in a second workshop discussion to ensure that common definitions were applied and that all scorers were aware of DPS or location-specific factors affecting vulnerability. Final scores were then submitted.

The bins were assigned a numerical value (low = 1, moderate = 2, high = 3, very high = 4) to calculate a weighted-mean attribute score. The number of tallies in a bin served as the weighting factor. A greater spread of tallies among bins reflected greater uncertainty in scores and was captured by the standard deviation of the mean score for each attribute.

#### Vulnerability categories

We calculated climate vulnerability for each DPS from its attribute scores in three steps [45]. First, we calculated the weighted mean of tallies for each sensitivity and exposure attribute. Second, we applied a logic model to determine cumulative sensitivity and exposure component ranks from their constituent attributes (Table 3). Rankings from the logic model depended on the number of attribute means that exceeded a specified threshold. For example, if at least two attributes in one component had a mean score equal to or above 3.5, that component was ranked very high.

Sensitivity and exposure component ranks were then assigned a numerical value (very high = 4, high = 3, moderate = 2, low = 1), which was used in the final step. Overall vulnerability was determined by multiplying the numeric values for sensitivity and exposure and assigning a total score for each DPS based on the product (Table 3).

We used a bootstrap analysis to characterize uncertainty in the assignment of a climate vulnerability category [45]. The 20 tallies for each attribute (four scorers per DPS with five tallies

Overall sensitivity or exposure score	Numeric score	Logic rule
Very High	4	More than 3 attribute means $\geq$ 3.5
High	3	More than 2 attribute means $\geq$ 3
Moderate	2	More than 2 attribute means $\geq 2.5$
Low	1	All other scores
Cumulative vulnerability	Component product	Component combinations
Very High	≥12	Very high/high or Very high/very high
High	8-11	Very high/moderate or High/high
Moderate	4-6	Very high/low, High/moderate, or Moderate/ moderate
Low	< 3	High/low, Moderate/low, or Low/low

**Table 3. Logic rule for ranking sensitivity and exposure components and cumulative vulnerability.** We used the logic rule across attributes to assign a numeric score and vulnerability category to sensitivity and exposure components (top section). We then used the product of the numeric component scores to assign cumulative vulnerability for each DPS (bottom section).

https://doi.org/10.1371/journal.pone.0217711.t003

each) were randomly sampled with replacement 1,000 times. From the resampled tallies, we calculated new climate vulnerability attribute means and final vulnerability categories using the three steps described above.

If the bootstrap outcome matched the original vulnerability category at least 75% of the time, we considered the score for that DPS to be likely. When 25% or more of the bootstrapped outcomes were either above or below the original climate vulnerability category, we considered the DPS to be borderline between the original and secondary vulnerability categories. Individual bootstrap results are shown in the <u>S3 Appendix</u>.

#### Vulnerability profiles

To explore which attributes were most important in determining overall vulnerability and how specific threats varied across DPSs, we conducted a hierarchical cluster analysis on the full suite of mean scores for all attributes, implemented in the R "cluster" package [115]. These clusters helped visualize differences in specific threats over broad geographical and biological gradients. To group similar DPSs, we applied the Ward's minimum variance algorithm and a Euclidean distance measure. We cut the resulting dendrogram into six groups. We then used a classification and regression tree analysis implemented in the R "tree" package [116] to identify which attributes best predicted cluster assignments.

We characterized vulnerability profiles for each cluster by computing the average score for each attribute across DPSs within each cluster. To show the general source of the threats, we grouped exposure and sensitivity attributes into four categories: freshwater exposure, marine exposure, life-stage sensitivity, and extrinsic sensitivity. We highlighted attributes with a mean cluster score of 3 or greater within each attribute category. We excluded all attributes that did not differ across DPSs, such as exposure and sensitivity to ocean acidification and ocean currents. All analyses were performed in R [117].

#### Highly vulnerable life stages

The overarching principle of this vulnerability assessment is that the most vulnerable DPSs are those most sensitive to climate change *and* most exposed to changing environmental conditions [5]. We applied that same logic to determine which life stages within each DPS were most vulnerable. Because life stages are typically segregated from each other in space and time, they tend to be affected by different exposure attributes. It was thus possible to pair specific sensitivity and exposure attributes. For example, freshwater life stages occurring over fall and winter are most exposed to extreme rain events and *flooding*, whereas those occurring in summer are exposed to *stream temperature* and *summer water deficit*.

The specific attributes most relevant at each life stage varied among DPSs due to differences in life history timing (Fig 2). However, for all DPSs, *hydrologic regime* was paired with the *juve-nile freshwater stage* and *sea level rise* was paired with *estuary stage*. All other marine exposure attributes were paired with the *marine stage*. To identify highly vulnerable life stages, we examined these sensitivity/exposure pairs and identified cases with scores higher than 3 in both attributes.

#### **Results**

#### **Relative vulnerability**

Five Chinook, one coho, and one sockeye salmon DPSs ranked very high in total vulnerability to climate change due to a combination of high and very high scores for sensitivity and exposure (Figs  $\underline{3}$  and  $\underline{4}$ , red boxes). Bootstrap analyses indicated that two additional DPSs, Southern





https://doi.org/10.1371/journal.pone.0217711.g003

Oregon/Northern California Coast coho and Mid-Columbia spring-run Chinook, were borderline between high and very high (<u>S3 Appendix</u>). Among species, Chinook salmon had the highest vulnerability rankings overall (mostly very high and high rankings), followed by coho and sockeye (Fig <u>4</u>). Steelhead and chum DPS scores were generally lower and nearly equally spread across high and moderate vulnerability categories. The only species in the low vulnerability category was pink salmon, which was represented by a single, unlisted DPS. Individual DPS scores are presented in the <u>S2 Table</u> and discussed in the <u>S3 Appendix</u>.

The preponderance of coho DPSs ranked very high in vulnerability to climate change were those occupying regions from southern Oregon to central California. Chinook and sockeye DPSs that ranked very high in vulnerability were concentrated in the two interior recovery domains: Central Valley and Interior Columbia. These results suggest that a combination of life history characteristics and geographic influences (including anthropogenic factors) contributed to high vulnerability for coho near its southern range limit and for Chinook and sockeye throughout the interior domains.



Fig 4. Number of DPSs in each vulnerability rank by species.

https://doi.org/10.1371/journal.pone.0217711.g004

The sensitivity component spanned all vulnerability categories across DPSs and generally aligned with cumulative vulnerability ranks (Fig 3). By contrast, the exposure component of vulnerability was relatively homogeneous across DPSs: of the 33 DPSs evaluated, 29 had high exposure to climate change (Fig 3). This consistency stemmed from exposure scores that were uniformly very high for *exposure to ocean acidification* and mostly high for *sea surface temperature* and *stream temperature*.

Only pink and chum, both typically coastal species, received low or moderate scores for these temperature-related attributes. In the Interior Columbia, exposure scores for both *stream temperature* and *hydrologic regime* were near or above the threshold for very high. Sensitivity to loss of snowpack was generally higher for spring-run Chinook than for steelhead and sockeye due to differences in spawn timing and habitat, respectively. For coho, threats from exposure to *stream temperature*, *flooding*, and *sea level rise* pushed some DPS scores near the edge of the very high category.

To ensure that high scores in multiple categories did not reflect "double counting," we assessed all pairwise correlations between attributes. Attributes that were not strongly correlated were assumed to capture different aspects of climate change, and therefore not double counting. Two pairs of attributes had a Spearman's *rho* correlation coefficient over 0.75: *sea level rise* and *estuary stage*, and *sea surface temperature* and *flooding. Sea surface temperature* and *flooding* reflect independent effects of climate change and hence represented distinct impacts of concern rather than double counting. The *sea level rise* and *estuary stage* pair may reflect some shared impacts; however, the populations most dependent on *the estuary stage* are also those exposed to the highest rates of *sea level rise*. Nonetheless, we confirmed that neither of these correlations affected final vulnerability categories by removing one from each pair and recalculating vulnerability scores.

#### Adaptive capacity

Adaptive capacity scores reflected the opportunity perceived by scorers for trait plasticity to help mitigate the negative effects of climate change (<u>S3 Appendix</u>). Results showed strong geographical patterns (<u>Fig 5</u>). All California Chinook and coho DPSs, the southernmost steelhead

ank	Very High	Snake River spring/summer-run Chinook	Upper Willamette River Chinook	Central California Coast Coho Central Valley fall/late fall-run Chinook Central Valley spring-run Chinook Sacramento River winter-run Chinook Snake River Sockeye
r ra			Hood Canal summer-run Chum	
5	High		Lower Columbia River Coho	
-21		Puget Sound Chinook	Mid-Columbia spring-run Chinook	California Coastal Chinook
ti.		Puget Sound Steelhead	Middle Columbia River Steelhead	Southern California Coast Steelhead
S1		Puget Sound Coho	Oregon Coast Coho	Southern OR/Northern CA Coast Coho
Sn		Snake River fall-run Chinook	Snake River Basin Steelhead	
Se			Upper Columbia River spring-run Chinook	
q			Upper Columbia River Steelhead	
an			Upper Willamette River Steelhead	
ð			Central California Coast Steelhead	
IL	Moderate	Northern California Steelhead	South-Central California Coast Steelhead	
SI		Lower Columbia River Steelhead	Columbia River Chum	Lake Ozette Sockeye
0		Lower Columbia River Chinook	Puget Sound Chum	
X			Central Valley Steelhead	
ΓŢ	Low		Puget Sound Pink	
	High Moderate		Moderate	Low

# Adaptive capacity

Fig 5. Adaptive capacity rank plotted against vulnerability rank. Vulnerability ranks were determined by exposure and sensitivity attributes (Fig 3). Adaptive capacity attribute scores reflected the opportunity perceived by scorers that some trait change would help mitigate the negative effects of climate change.

https://doi.org/10.1371/journal.pone.0217711.g005

DPS, and both sockeye DPSs scored low in adaptive capacity. The southernmost DPSs within each species may already be near tolerance limits, but these DPSs also have some of the most severe anthropogenic impacts and therefore limited scope for potential adaptations to a warmer climate. This explanation applied to Snake River sockeye. In contrast, Lake Ozette sockeye is not climate stressed at present and was simply not expected to change phenotypically.

The DPSs that scored highest in adaptive capacity were Puget Sound Chinook, coho, and steelhead; Lower Columbia Chinook and steelhead; and Snake River spring/summer Chinook and fall Chinook. Northern California steelhead also scored high in adaptive capacity. Higher scores reflected extensive life history diversity in both juvenile and adult stages. Most high-scoring DPSs display extensive juvenile life history variation, such as the subyearling and year-ling Chinook smolts, or 1- to 3-year-old steelhead smolts. Puget Sound and Lower Columbia Chinook display both spring and fall adult migration patterns, and Northern California steelhead migrate over two protracted periods, from late fall to spring for the winter-run and from spring to summer for the summer-run ecotype.

Chinook, coho, and steelhead DPSs had high variation in *adaptive capacity* scores, which ranged from low to high, whereas in other species, all DPSs fell into a single category. For example, all chum and pink DPSs scored moderate, while both sockeye DPSs scored low. There was uncertainty about whether sockeye rearing conditions would become less suitable, but the scorers' best estimate was that smolt age was unlikely to change, and any changes in adult migration timing would not substantially reduce climate stress. Selection for earlier adult

run timing in Snake River sockeye could be occurring at present [<u>31</u>]. However, the long migration through exceptionally warm reaches of the Snake and Salmon River will likely continue to challenge this DPS. The existing population is largely supported by captive broodstock and large hatchery releases; therefore, it is not subjected to the full effects of natural selection. How this might change in the future is uncertain.

#### Vulnerability profiles

Broad geographic trends in both exposure and sensitivity attributes were seen across DPSs, owing to the large spatial scale of climate drivers (Fig 6). In both in the Central Valley and Interior Columbia domains, DPS scores trended higher in both exposure and sensitivity than corresponding scores from their respective adjacent coastal domains (Fig 6, lower panels). In coastal domains, DPSs benefitted from the buffering effects of the Pacific Ocean and California Current, both of which ameliorate climate extremes. Fish in coastal domains also encountered fewer anthropogenic hindrances to migration.

Southern DPSs also tended to score higher in vulnerability than northern DPSs. For example, coho from the southernmost Central California Coast DPS ranked higher in vulnerability than those from the mid-latitude Southern Oregon/Northern California Coast, which in turn ranked higher than the three northernmost coho DPSs. This latitudinal pattern was also evident at the scale of recovery domains, where DPSs of the three coastal domains in California and Oregon were generally more vulnerable than those of the two coastal domains in western Washington. However, exceptions to these general trends were not uncommon.

To better elucidate general patterns of vulnerability, we used a cluster analysis to group DPSs with similar vulnerability characteristics and examined these groupings in relation to geographical gradients and species characteristics. At the highest level of the dendrogram, DPSs clustered into southern and northern groups (Fig 7). *Flooding* was the best predictor of separation between southern and northern branches and the second best between coastal and interior branches. Southern and coastal DPSs faced higher *flooding* due to intensification of atmospheric rivers, which were projected to change more in southern than northern latitudes. Interior Columbia DPSs were less affected by these extreme rain events.

The next level of the dendrogram primarily separated DPSs by species (Fig 7), especially steelhead vs. other species. We noted that both southern and northern steelhead clusters included some fall-run Chinook DPSs. Classification and regression tree results pointed to the sensitivity attributes of *early life history* and *juvenile freshwater stage* as key predictors of separation between clusters at the species level (S9 Fig).

Steelhead spawn in late winter and spring, and hence are less sensitive to changes in fall and winter precipitation than fall-spawning salmon. Heat tolerance and behavioral flexibility also tended to reduce threat levels for steelhead in the *juvenile freshwater* stage. The Chinook DPSs that grouped with steelhead were primarily those with subyearling type juveniles. Their shorter freshwater period produced relatively low vulnerability scores during the *early life history* and *juvenile freshwater* stages. In particular, the fall-run subyearling juvenile type avoids dependency on rearing in freshwater during summer, when thermal impacts, hydrologic regime shifts, and low-flow impacts are expected to be highest. We grouped more similar DPSs into the six groups indicated with rectangles in Fig 7. Group names reflect the predominant species and region of DPSs in each group.

In the northern cluster of the dendrogram, spring-run Chinook and sockeye from the Interior Columbia grouped with Upper Willamette River Chinook. These three DPSs share a temperature-stressed adult migration and summer holding period. Puget Sound pink and both



**Fig 6. Mean exposure and sensitivity scores by species and recovery domain.** Exposure scores are shown at left and sensitivity scores at right by species in upper panels and by recovery domain in lower panels. Because of the small number of DPSs in some domains, in Fig.6 the three recovery domains from southern Oregon to southern California are lumped into a California Coast group, and Oregon Coast is lumped with Lower Columbia. Boxes indicate the interquartile range of the data, whiskers show 1.5 \* the interquartile range, and the black line shows the median value.

https://doi.org/10.1371/journal.pone.0217711.g006

fall-run chum DPSs formed another group (with no listed DPS). The final group included all other DPSs from central and northern Oregon and western Washington.

Each cluster also displayed a unique vulnerability profile (<u>Table 4</u>). Profiles varied widely, from high scores in *ocean acidification* only (pink/fall chum), to high scores for freshwater and marine exposure but not for sensitivity to extrinsic stressors (Western Washington and Oregon), to high scores for freshwater and marine exposure and for extrinsic stressors (northern and southern steelhead), and finally, high scores in freshwater, marine, life stage, and extrinsic attributes (southern Chinook/coho as well as interior Columbia Chinook/sockeye).

In *adaptive capacity*, the southern Chinook/coho cluster had the lowest mean score. The northern steelhead cluster had the highest mean score, although this resulted from high scores for the two Chinook DPSs included in this cluster (see Fig 7). When the Lower Columbia River Chinook DPS was included in the Western Washington/Oregon cluster, that group had





https://doi.org/10.1371/journal.pone.0217711.g007

the highest adaptive capacity. Steelhead DPSs from the northern recovery domains had moderate adaptive capacity on average.

Specific attributes often scored in similar rank order across clusters (Table 4), although regional and biological variations were frequent and provided important insights for recovery planning. Among freshwater exposure attributes, *stream temperature* scored high in most clusters, but *flooding* was high only in the two California clusters, and *hydrologic regime* was high only in the interior cluster. Both of these latter attributes reflected changes in flow and/or precipitation, with increased flooding and drought more relevant in southern locations and loss of snowmelt more relevant in northern locations.

Among marine attributes, *exposure to ocean acidification* and *sea surface temperature* were highest in all clusters, with *sea level rise* second or a close third in both southern clusters as well as the cluster for pink/fall chum. For southern coho, sea level rise may not affect DPSs directly, but may represent a general threat to the freshwater/marine interface, triggering changes in lagoon habitat or sand-bar breaching. The most sensitive life stage differed among clusters, with adult stages more sensitive for interior DPSs, and juvenile stages more sensitive for southern coho/Chinook DPSs. Finally, among extrinsic sensitivity factors, *other stressors* was the

	Vulnerability profile cluster group					
Attribute	Southern Chinook/coho	Interior Columbia Chinook/ sockeye	Southern Steelhead	Northern Steelhead	Western WA/OR	Pink/ Chum
Freshwater Exposure						
Stream temperature	3	3	2	3	3	2
Flooding	3	2	3	2	1	1
Hydrologic regime	2	3	1	2	2	2
Summer water deficit	2	2	2	2	2	1
Marine Exposure						
Ocean currents	1	1	1	1	1	1
Sea level rise	3	1	2	1	2	2
Upwelling	3	2	2	1	1	1
SST	3	3	3	3	2	2
OA exposure	4	4	4	4	4	4
Life Cycle Sensitivity						
Early life history	2	1	1	1	2	2
Adult freshwater	2	3	2	2	1	1
Juvenile freshwater	3	3	2	2	2	1
Cumulative life-cycle	3	3	2	2	2	2
Estuary	3	1	2	1	2	2
Marine	3	2	2	2	2	2
OA sensitivity	1	1	1	1	1	2
Extrinsic Stressors						
Other stressors	3	3	3	3	2	2
Population viability	3	3	2	2	2	2
Hatchery influence	2	2	1	3	2	1
Adaptive Capacity						
Adaptive capacity	1	2	2	3	2	2

Table 4. Vulnerability profiles by cluster. Mean cluster score was the mean attribute score across DPSs within the cluster. Scores were rounded down for each attribute. Red cells indicate a mean score of 3 or higher for exposure and sensitivity or lower than 1.5 for adaptive capacity.

https://doi.org/10.1371/journal.pone.0217711.t004

most common attribute to score high, and paired with *hatchery influence* for the northern steelhead cluster and with *population viability* for the Interior Columbia Chinook/sockeye and California Chinook/coho clusters.

Attributes that varied most across clusters reflected the major factors that differentiated the most vs. least vulnerable DPSs across the entire study (<u>Table 4</u>; <u>Fig 8</u>). Overall, the least sensitive DPSs spent the least amount of time in freshwater (pink and fall chum), while the most sensitive spent more time in freshwater, had long summer adult migrations, or were heavily dependent on estuaries and near-shore coastal rearing habitat. Exposure factors that indicated the highest vulnerabilities to climate change were encountered in both freshwater and marine environments.

Adaptive capacity also differentiated DPS clusters, especially in relation to behavioral flexibility, which relies on both inherent plasticity and habitat heterogeneity. Conceptually, we refer to "natural processes" as the absence of anthropogenic stressors, such as hatcheries and habitat loss (Fig 8). Anthropogenic stressors scored in the sensitivity component are linked to adaptive capacity in a broad sense because they are inherently more malleable than life histories. In general, DPSs with the highest sensitivity and exposure and lowest adaptive capacity were the most vulnerable to climate change, as indicated by the red x in Fig 8.



**Fig 8. Conceptual model of highly influential attributes in final vulnerability ranks.** The most vulnerable DPSs had scores in the intersection of high rates of change in exposure, long periods of sensitivity, and low adaptive capacity attributes (red x), as detailed in <u>Table 4</u>. Natural processes here refer to the absence of other stressors and hatchery influences.

https://doi.org/10.1371/journal.pone.0217711.g008

# Highly vulnerable life stages

Among life stages that scored high in both sensitivity and exposure, regional differences were pronounced. For coho in the two southern recovery domains and for Chinook in the Central Valley, DPS sensitivity scores were high at both the *estuary* and *marine stages*, and exposure scores were high for *sea level rise* and *sea surface temperature* (Table 5). For coho, steelhead, and some fall Chinook in the southern recovery domains, estuary conditions also affect access to freshwater spawning habitat, where watersheds are seasonally blocked by sand bar formation. Specific projections of how estuary and lagoon dynamics will change adult access were not available, and thus did not greatly change scores for the *estuary* or *adult stages* for these DPSs. However, potential obstruction to spawning habitat was noted as a concern by scorers.

Because of the highly modified Sacramento-San Joaquin Delta, all Central Valley DPSs were vulnerable at the *estuary stage*. Diversion of water from the delta supports the largest agricultural economy in the U.S. and provides drinking water to more than 20 million people [118]. Extensive water infrastructure in the estuary has dramatically altered flows and reduced survival of migrating fish. Furthermore, juveniles from all three Central Valley Chinook DPSs migrate predominantly as subyearlings, and as such are greatly dependent on estuary and near-shore habitat during the critical first year. These DPSs are therefore more vulnerable to sea level rise than DPSs with yearling-type juveniles.

**Table 5. Highly vulnerable life stages by DPS with associated exposure attributes.** Criterion for inclusion was a score of 3 or higher in both sensitivity and exposure attributes in each subheading. Additional high exposure scores for upwelling (a), flow regime (b), and flooding (c) also occurred in some DPSs.

Life stage and DPS	Exposure attribute
Early life history	Summer water deficit/Hydrologic regime
Sacramento River winter-run Chinook	Summer water deficit
Puget Sound Chinook	Hydrologic regime
Juvenile freshwater stage	Stream temperature
Mid-Columbia spring-run Chinook <sup>b</sup>	
Snake River spring/summer-run Chinook <sup>b</sup>	
Upper Columbia River spring-run Chinook <sup>b</sup>	
Lower Columbia River coho	
Oregon Coast coho	
Central California Coast coho <sup>c</sup>	
Southern Oregon/Northern California Coast coho <sup>c</sup>	
Puget Sound coho <sup>b</sup>	
Estuary stage	Sea level rise
Central Valley fall/late fall-run Chinook	
Central Valley spring-run Chinook	
Sacramento River winter-run Chinook	
Central California Coast coho	
Southern Oregon/Northern California Coast coho	
Marine stage	Sea surface temperature
Central Valley fall/late fall-run Chinook <sup>a</sup>	
Sacramento River winter-run Chinook	
Central California Coast coho	
Southern Oregon/Northern California Coast coho	
Adult freshwater stage	Stream temperature
Mid-Columbia spring-run Chinook	
Middle Columbia River steelhead	
Snake River Basin steelhead	
Snake River sockeye	
Snake River spring/summer-run Chinook	
Upper Columbia River spring-run Chinook	
Upper Columbia River steelhead	
Upper Willamette River Chinook	
Central Valley spring-run Chinook	
<sup>a</sup> Exposure to upwelling also scored high	

<sup>b</sup>Exposure to hydrologic regime also scored high <sup>c</sup>Exposure to flooding also score high

https://doi.org/10.1371/journal.pone.0217711.t005

All coho were highly vulnerable at the *juvenile freshwater stage* because of its extended duration (1+ years) and corresponding longer exposure to rapidly rising summer *stream temperatures*. In three of the five coho DPSs, *flooding* or *hydrologic regime* posed an additional high exposure at the *juvenile freshwater stage*.

Yearling Chinook, which are characteristic of many populations in the Interior Columbia recovery domain, were also highly vulnerable at the *juvenile freshwater* stage because of the year-round reliance during this stage on freshwater habitat. Although specific habitat preferences differ, both coho and Chinook are sensitive to changes in summer flow and *stream* 

*temperature*. Most Interior Columbia Chinook DPSs were also highly vulnerable to temperature in the *adult freshwater stage* due to long adult migrations in spring and summer through highly modified rivers, along with exposure to high summer *stream temperatures* during the holding period prior to spawning. Upper Willamette spring-run [119, 120] and Central Valley spring-run Chinook [121] face similar thermal challenges and high mortality between adult migration and spawning. Snake River fall-run Chinook did not score high in sensitivity to *stream temperature*, although adults do encounter high temperatures during late-summer migrations and have experienced compromised fecundity as a result [122, 123].

For most DPSs, sensitivity attributes were not scored high at the *early life history* stage. Puget Sound Chinook was an exception due to sedimentation and scour during flood events. Sacramento River winter-run Chinook also scored high in sensitivity at this stage. This DPS incubates eggs over summer, when *stream temperature* can be high if cold water in the Shasta reservoir is insufficient to cool the upper Sacramento River throughout the incubation period [<u>38</u>]. A recent analysis of Upper Willamette River spring-run Chinook indicated high temperatures are projected to increase mortality in the egg stage for this DPS as well, because spawning habitat is constrained by dams to the lower river reaches [<u>124</u>].

#### Data quality

Most evidence used for scoring was based on quantitative data specific to each DPS, although DPSs were often represented by only a few index populations that were monitored consistently. In some cases, information on one DPS had to be inferred from a similar DPSs. Still, relative to information available for most marine fish, data quality was quite strong.

With the exceptions of the highly quantified projections for *exposure to ocean acidification* and *sea surface temperature*, sensitivity attributes tended to be based on higher-quality data than exposure attributes. Likewise, data for freshwater attributes was generally of higher quality than that for marine attributes (S10 Fig, top panel). Specifically, the freshwater life-stage sensitivity attributes of *early life history* and of *juvenile* and *adult freshwater stage* had relatively high data quality, as did assessments of *population viability* and *other stressors*. The weakest data for sensitivity attributes concerned sensitivity to *ocean acidification* and survival during the *marine stage*. In both of these cases, impacts on salmon were mediated by numerous potential food web interactions, which made net effects difficult to predict.

Some of the highest data-quality scores among the exposure attributes were from hydrological and stream temperature models. *Flooding* and *water deficit* exposures were less certain, and this was also reflected in high standard deviations in scores (wider spread across bins, <u>S10</u> Fig, bottom panel). In the marine environment, data for *exposure to ocean acidification* and *sea surface temperature* were of very good quality, with consistent results across many models. However, projected changes in *ocean currents* and *upwelling* were inconsistent across models. Of all attributes, *upwelling* had the highest mean standard deviation of tallies across bins, indicating the largest uncertainty.

# Discussion

#### Spatial and biological patterns in vulnerability

Patterns in climate vulnerability have important implications for Pacific salmon across the Pacific Coast, both in terms of total variation in life history diversity and in the likelihood of southern or interior range-edge contractions. The DPSs most vulnerable to climate change were those with life history types presently rare in the CCLME but prevalent further north, such as spring-run Chinook, and those unique to the species as a whole, such as late-fall and winter-run Chinook and summer-run chum. For Chinook, the highest vulnerability scores

were for DPSs of the Central Valley and Interior Columbia recovery domains. For sockeye and steelhead as well, DPSs of the Interior Columbia scored higher than those of the coastal domains. This geographical pattern suggested a potential range contraction toward the coast for anadromous life histories unless access to higher-elevation habitats is restored and habitat quality in rearing areas and migration corridors is improved [108].

For coho, which have been extirpated from interior basins, vulnerability was very high in the entire southern portion of the range throughout California and southern Oregon. Finally, for steelhead, the southernmost salmonid in the CCLME, low *adaptive capacity* (potential loss of anadromy) and proximity to critical thresholds in the present climate raise the possibility of impending range contraction. Lower exposure scores for southern coastal DPSs suggest such a contraction could be coastward rather than northward. Resident forms of *O. mykiss* may persist in the inland areas, although these populations may become increasingly isolated [125].

**Species-level results and similarities with other vulnerability assessments.** Although we employed a rapid-assessment, our findings were of sufficient detail to provide conclusions similar to those of more geographically or ecologically focused studies [126–130]. Among species we considered, Chinook and coho had the greatest proportion of highly vulnerable DPSs. Climate vulnerability for the two sockeye DPSs was split between very high and moderate (Fig 4), while steelhead and chum DPSs were intermediate between high and moderate vulnerability. Puget Sound pink salmon scored lowest in vulnerability.

This species-level ordering was consistent with results from the West Coast Fish Climate Vulnerability Assessment (M. Haltuch, NOAA Fisheries, personal communication), which ranked Chinook salmon vulnerability *very high*, and coho, sockeye, steelhead, and chum *high*. In the Eastern Bering Sea Climate Vulnerability Assessment, all five salmon species scored high in sensitivity but low in exposure (P. Spencer, NOAA Fisheries, Seattle, personal communication). Lower exposure further north as well as increasing abundance and apparent range expansion of Chinook [131, 132] and Atlantic salmon (*Salmo salar*) [133] suggest that salmon species may shift the centroid of their respective ranges northward, as predicted by other niche-mapping studies [75, 76].

Several other groups have conducted vulnerability assessments that included Pacific salmon and steelhead. Both the NOAA Fisheries Multi-species Recovery Plan [126] and Moyle et al. [127] ranked California salmonids. Pacific Northwest steelhead were ranked by Wade et al. [128, 129], and all species were included in an assessment by the Washington Department of Fish and Wildlife [130]. Our relative ranks were similar to ranks from other studies, especially for Chinook and coho salmon, unlike recent reviews where systematic comparisons of vulnerability assessment results for terrestrial species found poor congruence [107, 134]. Salmon assessments may be more similar to each other both in the data that is analyzed and the categories of threats that are considered than across the broad spectrum of terrestrial taxa; a comparison of lizards produced a similar congruence [106]. Nonetheless, variation in spatial resolution and criteria for vulnerability do produce different results, which should be kept in mind when using these results in management decisions.

Steelhead vulnerability was somewhat more variable across studies than Chinook and coho, partly depending on whether authors rated loss of ecotypes vs. loss of the DPS as a whole. Studies with finer spatial and temporal resolution had greater differentiation of risk and generally higher vulnerability scores, potentially resulting from severe local stressors and specialized ecotypes. For example, certain parts of the Northern California steelhead DPS, specifically the summer-run ecotype, were scored as critically vulnerable in the *Multi-Species Recovery Plan*. On the other hand, studies with coarser spatial and biological resolution placed nearly all salmon and steelhead in a single, moderate-high or high-very high risk category [130].

Overall, the factors that caused highest vulnerability ratings among salmon DPSs are the same factors that caused higher vulnerability in many diadromous species compared with marine species [35]. As a functional group, diadromous species (e.g., sturgeon, *Acipenser* spp, Blueback herring, alewife and American shad, *Alosa* spp., and Atlantic salmon), had the highest proportion of vulnerable species in the Northeast Climate Vulnerability Assessment. The risk to Atlantic salmon was considered very high. Thus, salmon populations on both coasts are likely to contract northward for similar reasons [135]. Diadromous species rely on sequential freshwater, estuarine, and marine habitats; therefore, these species face a diverse suite of threats from climate change throughout their complex life cycles.

**Specific climate threats.** High exposure ratings throughout our results stemmed from a relatively consistent suite of exposure attributes (<u>Table 4</u>, <u>Fig 8</u>). Nearly all populations face high exposure to changes in sea surface temperature and ocean acidification, and most will confront considerable increases in summer stream temperatures. Accordingly, scores for these attributes were generally quite high (<u>Table 4</u>).

In freshwater and estuarine environments, other impacts varied by latitude and proximity to the coast. Exposure scores were generally higher for southern than for northern DPSs in *sea level rise, flooding*, and *upwelling*. Sea level is projected to rise more slowly in the northern CCLME, where geological uplift compensates somewhat for an expanding ocean [97, 136]. Dramatic increases in projected flooding along the West Coast stem from intensification of atmospheric rivers—a consequence of warmer temperatures over the Pacific Ocean [67]. Among present global models, California is projected to experience the greatest change in atmospheric rivers [67, 137–140]. Changes in the intensity and timing of *upwelling* are less certain. Nonetheless, present models suggest that the largest changes will manifest off the coast of California [98, 141], where relatively mild summer stream temperatures depend fundamentally on upwelling and the closely associated fog regime.

Salmon and steelhead in interior regions, as well as those in Puget Sound, had generally high DPS exposure scores for *hydrologic regime* due to loss of snowpack in mid- and high-ele-vation watersheds. Snowpack is already declining in response to warmer winters throughout the western U.S. [4, 142, 143]. In mountainous regions, warmer winters will transform snow-dominated hydrographs with low winter flows followed by a protracted spring snowmelt to systems characterized by rapid snowmelt and high-flow events during the incubation period [142]. In western Washington, salmon populations may soon lose snow-dominated water-sheds entirely [144]. Such losses are expected to reduce life history diversity within these DPSs [68]. While these DPSs may be buffered from outright extinction by their existing behavioral diversity, losses of habitat diversity and cooling influences of snowmelt may increase vulnera-bility [104].

Interior Columbia DPSs face the largest percentage loss of snow-dominated habitat [144]. These populations are dominated by life history types specifically adapted to elevated flows in spring, which expedite juvenile migrations of up to 1500 km. Summer stream temperatures are also cooler in snow-dominated basins. Characteristic life history strategies in these regions, such as summer juvenile rearing and adult holding depend on these cooling influences. Hence, these genetically distinct life histories are perhaps most threatened by loss of snow cover.

California steelhead tended toward more moderate exposure scores for stream temperature because of a weaker link between rising atmospheric and *stream temperature* in coastal California. Heat-moderating factors such as coastal fog, riparian evapotranspiration [145], and groundwater inputs, are especially relevant in some locations and contribute to a decoupling of stream and air temperatures [146]. This decoupling suggests a potential capacity for thermal refuges from rising air temperatures, all else being equal. However, the buffering capacity of

mitigating factors such as fog could diminish in a warmer climate [147], increasing exposure to *stream temperature* well beyond the moderate levels we scored.

Decoupling of air and water temperatures can also result from a high frequency of intermittent streams in rain-dominated basins such as those in southern California [148, 149], and elsewhere [150]. Climate change could entail an end to this decoupling process as well, limiting future habitat to a greater extent than reflected in our stream temperature exposure scores [151, 152]. In short, coastal steelhead in California may be somewhat protected by thermal refuges, but the factors maintaining those refuges themselves likely have climatic thresholds beyond which they cease to operate.

**Most vulnerable life stages.** Salmon life history types are closely tied to hydrological conditions, so the geographical patterns in exposure factors parallel trends in highly vulnerable life stages. At the scale of this assessment, Chinook demonstrated these patterns most clearly because DPSs differ systematically in the duration of freshwater stages [20, 68, 153]. Southern Chinook DPSs currently lack access to snow-cooled juvenile habitat, so they characteristically smolt as subyearlings. Subyearling juveniles are more vulnerable to near-shore development, *sea level rise*, and *upwelling*. Thus for southern Chinook DPS juveniles, the *estuary* and *marine stages* were highly vulnerable (Table 5). For yearling Chinook and coho migrants, sensitivity scores were higher at the *juvenile freshwater stage* than the *estuary stage* because of their extended freshwater rearing strategies. These strategies, however, make them more vulnerable to *stream temperature* increases and loss of snowpack (*hydrologic regime shift*).

All of the DPSs with a highly vulnerable *adult freshwater stage* migrate in spring or summer, so they are exposed to high *stream temperatures* and pre-spawning mortality. The interior populations also confront long migrations. For southern coastal species, sensitivity in the adult stage might have scored higher due to difficulties accessing freshwater habitat. However, we could not quantify this difficulty owing to uncertainty regarding net change in sand-bar breaching. Our results primarily reflected the fact that longer migrations and freshwater phases expose salmon to more numerous freshwater climate threats and anthropogenic stressors.

Nevertheless, the steelhead we considered, including those with extended freshwater phases and migrations up to 1500 km (Table 1), tended to score lower in sensitivity than Chinook in the same region. Greater resilience in steelhead stems from several factors. First, steelhead inhabit streams warmer than those used by Chinook or coho salmon [61, 154, 155]. Compared to spring-run Chinook and sockeye salmon, steelhead also display greater mobility during migration, utilizing high-elevation, high-velocity, and hard-to-reach or ephemeral and intermittent stream habitats, as well as cool-water tributaries for temporary staging [156]. Despite these advantages, steelhead access to freshwater habitats can be intermittent and hindered by changes in storm frequency [157]. Second, although both species have a strong genetic component in life history traits, *O. mykiss* typically expresses more life history strategies within DPSs, so the DPS as a whole appears less vulnerable than Chinook DPSs [22, 158].

Relatively few DPSs appeared highly vulnerable in their *marine stage*. However, this was also the stage with the greatest uncertainty in scores. While physical conditions in freshwater are clearly and directly linked to salmon survival, factors that influence ocean survival are more complex [159]. Physical processes in the ocean affect salmon through their influence on prey availability and abundance, as well as through the spatial distribution of competitors and predators. Ocean food webs contrast sharply in cold vs. warm years [88, 160, 161]. The combination of increasing *sea surface temperature* and *ocean acidification* heightens the risk for a major, novel reorganization of marine ecosystems.

Marine biological regime shifts of the past are well documented  $[\underline{162}-\underline{164}]$ , and demonstrate widespread ecological responses to change in ocean conditions. These regime shifts were associated with climate changes much more subtle than those projected over the next few

decades; hence they provide only hints of potential of future impacts. For salmon especially, specific consequences of ocean regime change are hard to predict, owing in part to the general non-linearity of marine ecosystem dynamics, along with the numerous possible fish communities that could establish themselves [40]. Nonetheless, prolonged periods of poor ocean survival have been observed during generally warm decades [165]. In recent warm years, a high proportion of empty stomachs were observed in juvenile salmon, as well as poor body conditions, despite an abundance of prey biomass [166]. Thus although we have highlighted risks in freshwater stages, these findings suggest that warmer oceans could be catastrophic for salmon populations throughout the CCLME, as has also been suggested for Atlantic salmon [135].

#### Adaptive capacity

Although the *adaptive capacity* score was not as formalized as the rest of the assessment, results are consistent with larger patterns in habitat and life history diversity. Among DPSs with similar life history diversity, those that scored higher in *adaptive capacity* occupied habitat that was climatically diverse but generally closer to optimal for salmon. Such habitats featured moderate temperatures and wetter overall climates—conditions that support a large range of salmon life histories. Northern California steelhead occupy the interface between the more xeric southern and interior eco-climatic zones and the wetter zones of coastal Oregon and Washington, and they exhibit a wide range of juvenile and adult behaviors. Lower Columbia River Chinook and steelhead and Puget Sound Chinook also display various life histories at multiple life stages. Puget Sound and Lower Columbia DPSs benefit from steep elevation gradients of the Cascade and Olympic Mountains, which provide cool water without the hazards of a lengthy migration. This advantage was reflected in the lower life-stage sensitivity scores for the western Washington/Oregon cluster (Table 4).

For some DPSs, high *adaptive capacity* scores reflected direct evidence of adaptive change. For example, for Snake River fall-run Chinook, a shift in the proportion of fish adopting yearling vs. subyearling juvenile life history strategies has been observed [7, 167]. Increased proportions of yearling type fish appear to have evolved in response to anthropogenic habitat modifications. If climate change favors a reversal of this trend, then this DPS may be expected to continue such adaptive responses. Shifts in adult run timing have also been observed for multiple DPSs in the Interior Columbia recovery domain. Evolutionary changes in run timing were associated with environmental change [31], as well as hatchery supplementation [168]. Some of these DPSs inhabit heavily modified areas; for example, most Snake River salmon must pass eight large hydroelectric dams during both the juvenile and adult migration. Puget Sound DPSs inhabit an area of rapidly expanding human population, with a projected increase of 42% by 2050 [169]. However, unusual behaviors have emerged under altered conditions [170–172], suggesting that adaptive responses to climate change will continue to arise.

Most DPSs that scored high in adaptive capacity benefit from complex terrain that includes snow-cooled streams. However, these snow-dominated hydrological regimes have been consistently projected to disappear during the present century [144, 173], potentially causing a net contraction in life history variability. Thus, the selective landscape could shift to favor a different balance of traits, including some that are not exhibited now. Other DPSs scored moderate in *adaptive capacity*, either due to life history constraints in the case of chum and pink salmon, or due to habitat loss and degradation in the case of interior steelhead and other Chinook.

*Adaptive capacity* was ranked *low* for the three Central Valley Chinook DPSs, along with Central California Coast coho and Snake River sockeye. These ranks were associated with *high* scores for extrinsic exposure attributes and cumulative *life-cycle complexity* (Fig 5 and Table 4). Chinook, coho, and steelhead DPSs in the two southern clusters had the lowest

population viability scores and highest exposure to anthropogenic impacts (Table 4). These impacts included significant *hatchery influence* and *other stressors* such as water withdrawals/ diversions, habitat degradation, loss of access to higher elevation (cooler) spawning and rearing habitats [53, 108], and potential competition or predation from invasive species. Many of these stressors are expected to increase with climate change, as human demand for water increases [3, 4], warm-water invasive predators expand their range [174–180] and the virulence of some diseases intensifies [181–184].

Reductions in abundance, genetic and phenotypic variation, along with proximity to environmental tolerance limits, has brought many DPSs to a threshold of critical impairment to life history types. Low *adaptive capacity* and high *cumulative life cycle effect* scores reflected the fact that without access to historical habitats [108, 185, 186], southern DPSs have fewer options for behavioral mitigation of climate impacts [187] than their conspecifics to the north.

In part, low adaptive capacity scores for Central Valley Chinook resulted from its various life history types that have differentiated over evolutionary time and are considered distinct from one another at the DPS level. Each of the three Central Valley Chinook run types is specialized to a particular aspect of the hydrologic profile, and thus each is especially vulnerable to hydrologic change. In contrast, summer and winter steelhead run types are less genetically distinct and currently considered part of the same DPS [158]. Moreover, anadromy itself is more variable in steelhead than in Chinook. Many steelhead populations interbreed with resident forms of *O. mykiss*, with the frequency of alleles relating to anadromy fluctuating over time [125, 188]. Climate risks to steelhead include loss of the anadromous life history type as a major component of the DPS.

#### Historical trends in loss of diversity

We identified patterns of climate vulnerability that mirrored patterns of extinction estimated for all six species we assessed. Gustafson et al. [189] enumerated loss of historical populations and DPSs in the western U.S., with the concurrent loss of ecological, genetic, and life history diversity. Although overall estimated losses were considerable (29%), they found evidence of fewer extinctions along the northern coastal regions (<20%) compared with southern California (35%), the Central Valley (57%), and the interior Columbia Basin (35-62%). These patterns typified the north-to-south and coast-to-interior gradients in our vulnerability scores.

Greater losses in the interior domains were primarily due to large, impassable dams, which eliminated many populations simultaneously. Gustafson et al. [189] also found that for Chinook salmon and steelhead, extinction rates of stream-maturing populations with longer freshwater phases were higher than those of ocean-maturing populations that reside in freshwater for shorter periods. They found greater extinction rates in sockeye and coho compared with pink and chum salmon, also reflecting the predominant patterns seen in our assessment. Such similarities supported our conclusion that freshwater-dependent life history types are more vulnerable, and that climate change will likely continue the direction of anthropogenic pressures that have accumulated over the past two centuries.

Overall, both historical and future losses of diversity pose a critical challenge for all Pacific salmon species. At both the DPS and species level, the most fundamental components of *adap-tive capacity* are life history diversity, physiological performance, behavioral and morphological plasticity, and genetic variability. However, for West Coast salmon populations, some of the most distinctive and rare characteristics are those at greatest risk. At the same time, large proportions of stream habitat that could provide refuges to help sustain these populations have been lost to anthropogenic barriers [190].

#### Methods of increasing climate resilience

Most of the DPSs we evaluated are listed as species of concern, threatened or endangered under the ESA or are considered by states as sensitive, almost entirely as a result of anthropogenic stressors. Reducing anthropogenic stressors could greatly improve responses to climate change by improving the overall status of these DPSs in terms of abundance, productivity, spatial structure, and diversity.

A robust DPS has greater resilience by virtue of strong population dynamics that make stochastic extinction less likely. Such strengths rely on population spatial structures that provide refuge from disturbances and can allow adaptation to occur at fine scales, as well as diversity in genetic makeup, life history, behavior, and morphology [34, 108]. These processes provide the needed raw material to respond to climate change, allowing for a "portfolio effect" that reduces volatility and risk to the larger demographic unit [191–196]. Increasing synchrony in both climate [152, 197] and salmon population responses [198] indicates declining inter-population diversity and presents a major threat to DPS persistence.

Climate change presents an array of specific threats that can act synergistically with other threats, dramatically increasing the impacts of each [108]. In particular, the loss of population spatial structures, as well as habitat heterogeneity and connectivity, removes the means by which salmon have historically persisted through frequent disturbances and climate extremes. Recent analyses in terrestrial environments found a correlation between habitat loss and climate stress [199]. An analysis of bull trout (*Salvelinus confluentus*) also found that genetic richness is lower in habitats with the highest climate exposure [110]. Thus, due to past adaptation or recent stressors, *adaptive capacity* may already be at its lowest levels precisely where salmon need it most. In prioritizing conservation actions, it is therefore worth exploring specific interactions between existing threats and climate drivers.

Habitat restoration is especially important in allowing salmon to express their intrinsic life history diversity. Salmon are highly adapted to disturbance regimes, but they need access to a wide variety of physical and thermal conditions within a watershed if they are to respond to increasing climate variability, such as frequent flooding or persistent droughts. Three main themes have emerged from recent literature (e.g., [55, 108, 200]). First, reconnection of habitats blocked by artificial barriers, either longitudinally or laterally (floodplains), can be highly effective in expanding the effective climate space of a watershed. Reconnected habitats restore natural processes and provide refuges from extremes in both temperature and flow. Second, amelioration of temperature or flow constraints can actively reduce climate stress, for example, through hypolimnetic releases from reservoirs, reconnection to historical sources of cool water, riparian restoration, and other techniques. Finally, identifying and improving access to food-rich environments can improve tolerance of climate stress by reducing bioenergetic constraints and mortality risks that are often lower for larger fish.

Projects focused on restoration and protection of accessible habitat are underway in numerous river systems, although the scope of work needed for species recovery sometimes involves nearly all existing habitat [201]. Nonetheless, when estuarine and freshwater habitats and processes are restored, natural environmental complexity provides a buffer against climate impacts in some cases [202]. Model results show that restoration can mitigate for declines that would otherwise result from climate change [203–205]. Guidelines to identify habitat restoration actions that will have a climate benefit have been developed [55] and are being used to realign priorities in some watersheds [206], but have not become the norm [207]. Management of freshwater stream temperatures and flows to support a diversity of salmon life history strategies, as well as to improve survival (and thus abundance and productivity) will be a crucial tool for increasing resilience to climate change [108, 208, 209]. Large, impassable dams block access to large areas that could serve as climate refuges as well as supporting more diversity and larger populations in general [208, 209]. There have been major improvements in fish passage at dams on the mid and lower Columbia and Snake Rivers [210], and reintroduction of coho to the interior Columbia is currently underway [211]. Furthermore, removal of dams has become much more frequent in recent years, including dams on the Elwha, Rogue, White Salmon, Sandy, and Carmel Rivers [212–215]. Salmon responded rapidly when multiple dams were removed in the Rogue [216], Sandy [217] and Elwha River basins [218–220], as did other salmonids, including re-establishment of the anadromous life history in bull trout (*Salvelinus confluentus*) [221].

Nonetheless, a large fraction of historical salmon habitat is still completely inaccessible [190, 222]. Pilot efforts to establish experimental populations above some dams are underway. For example, reintroduction of winter-run Chinook to historical habitat in the Sacramento River Basin has involved removal of migration barriers and restoration of more natural flow (Battle Creek [223]), as well as transport above barriers that will continue to be impassable (McCloud River, above Shasta Dam [224]). Similar projects exist for coho, Chinook, and sockeye salmon and steelhead in the Columbia River Basin [225–228]. However, because certain dams will not be removed in several of these plans, assisted migration using trap-and-haul operations will continue to be essential [185, 186, 224], adding uncertainty for long-term population viability. In other cases, the inadequacy of existing dams to cope with new extremes of flow and sediment movement may support removal as a tool to mitigate climate change impacts.

Hatchery supplementation can reduce fitness in wild salmon populations both through introducing maladaptive genotypes and reducing the effective population size of wild populations [229, 230]. Therefore, reducing the number of hatchery-origin fish in general can be expected to improve the adaptive capacity of wild populations in the face of increasing exposure to climate change. In the case of highly endangered populations, however, hatcheries can provide a short-term buffer from extinction risks [231], which is the primary risk for salmon during adaptation to climate change. Criteria for limiting introgression between hatchery and natural-origin fish have been developed to reduce the risks of domestication [232]. Furthermore, improvements in hatchery spawning techniques, mating designs, incubation and rearing protocols, may reduce the potential for inbreeding and domestication selection [233].

Harvest practices also could be adjusted based on periods and conditions when populations are less stressed. For example, catch-and-release fisheries or fishing closures are used to restrict angling to cool temperature periods. Such practices mitigate the interaction between handling and temperature stress [234], but run the risk of accidentally selecting on run timing [235] and other traits [236, 237]. Consideration of how all anthropogenic factors exacerbate or possibly mitigate for climate stressors is much needed [238]. For example, fisheries typically select for smaller body size and shorter generation time, which could also be advantageous in a warming climate [239, 240]. However, these traits also reduce fecundity and population stability, which is ultimately disadvantageous for both humans and salmon population viability [239–243].

More active proposals of assisted gene flow and gene editing are being proposed to introduce more heat tolerant genotypes into hatchery programs [244] and wild populations [245– 247]. However, as with any new technology, risks are difficult to quantify and there are many factors that need to be considered [248–255]. In many cases, humans have intentionally or unintentionally caused traits to shift in direction or variability that are maladapted for climate change [238, 256–258], putting some DPSs at additional risk. More research is needed to identify best practices in relation to anthropogenic selection (e.g., [242]). Though many uncertainties remain to be addressed, all of these avenues can potentially improve opportunities for local adaptation and overall survival.

# Conclusion

Loss of the southernmost populations within a species' range is widely predicted with climate change [259], but our assessment also highlighted that unique life histories are at high risk. Both the late-fall and winter-run Chinook ecotypes exist only at the southern end of the species range, and both face extinction without continued intensive management. Similarly, for chum salmon, the summer-run is rare and faces relatively greater vulnerability than the more common fall or winter-run life history types in northern regions. Local adaptations to distinct flow and temperature conditions are the characteristics that contribute to high vulnerability for these life history types and make them particularly sensitive to climate change.

In addition to southern range contractions, we found that interior losses may be even greater, due in part to greater change to interior climates and anthropogenic constraints on migration pathways. Some life histories ranked highly vulnerable by us or others, such as spring-run Chinook and northern California summer-run steelhead, will still be represented further north. However, Chinook salmon and steelhead that evolved distinct lineages in interior basins [21, 22] are at risk of losing some of their unique life histories not only in the Columbia River Basin but also in the neighboring Fraser River Basin in Canada [32, 260, 261]. The evolution of early adult migration (spring-run Chinook and summer-run steelhead) appears to reflect a rare event that would be quickly lost if these migratory pathways are selected against [158]. Declines in these life histories could entail significant loss of diversity in these species as a whole.

The highest scores for extrinsic effects (anthropogenic stressors) occurred in interior and southern regions (Table 4), exactly where climate is expected to change the most. A similar pattern in smaller-scale genetic analyses [110] suggests this could be a widespread phenomenon. Efforts to promote resilience to climate change are similar to those that increase viability more generally and have been part of historical conservation practices. However, our assessment indicates that more intense and perhaps novel efforts will be needed to compensate for the added pressure from climate change. Additional research to refine this assessment and explore adaptive capacity would be especially valuable. For DPSs that scored high in adaptive capacity, particular care is warranted to avoid loss of life history diversity and thus maintain the flexibility to continue adapt to climate change in the future. Resource managers should expect changes in fish characteristics, such as run timing and body size, but also other responses which have unknown consequences for population viability.

By pointing to the most vulnerable DPSs, identifying the most vulnerable life stages within each DPS, and assessing where life histories are most likely to change, these results provide a framework to support recovery planning for climate change impacts on West Coast salmon. This assessment considered present conditions, and therefore present risks confronted by Pacific salmonids that are related to climate change. Most, if not all, Pacific salmonid habitat in the western U.S. has diverged significantly from historical conditions and processes. Where dams block passage and interrupt ecological and physical processes, dam removals will likely result in habitat that diverges less from those seen historically. This is likely to reduce impacts of climate change for fish at all life stages. As demonstrated by recent dam removals and restoration activities that reconnect floodplains, physical and ecological responses can be rapid and can effectively reduce habitat constraints on these systems [217, 218]. Thus, we may be able to provide some relief to the extensive climate change risks highlighted in this vulnerability analysis.

# Supporting information

**S1 Appendix. Biological sensitivity attributes.** (PDF)

S2 Appendix. Climate exposure factors. (PDF) S3 Appendix. Distinct population segment scores and narratives. (PDF) S1 Table. Exposure factor data. (XLSX) S2 Table. Distinct population segment scores. (XLSX) S3 Table. Adaptive capacity scores. (XLSX) S1 Fig. Chinook salmon habitat. (JPG) S2 Fig. Fall Chinook salmon habitat. (JPG) S3 Fig. Winter Chinook salmon habitat. (JPG) S4 Fig. Chum salmon habitat. (JPG) S5 Fig. Coho salmon habitat. (JPG) S6 Fig. Pink salmon habitat. (JPG) S7 Fig. Sockeye salmon habitat. (JPG) S8 Fig. Steelhead habitat. (JPG) S9 Fig. Classification and regression tree results. (PDF) S10 Fig. Data quality scores and standard deviation of tallies for each attribute. (DOCX)

#### Acknowledgments

We thank all who contributed information for our species profiles and consulted on attributes, particularly Martin Liermann, Flora Cordoleani, and Michael Beakes. Guillaume Mauger, Michael Warner, Eric Salathé and Dan Isaak provided information on freshwater attributes. Rick Gustafson, Robin Waples, and Jeff Jorgensen provided valuable comments on early versions of the manuscript. Use of trade, product, or firm names is for descriptive purposes only and does not imply endorsement by the U.S. Government.

# **Author Contributions**

**Conceptualization:** Lisa G. Crozier, Michelle M. McClure, David A. Boughton, Melissa A. Haltuch, Elliott L. Hazen, Nathan J. Mantua, Mark W. Nelson.

Data curation: Lisa G. Crozier, Tim Beechie, Correigh M. Greene.

Formal analysis: Lisa G. Crozier, Mark W. Nelson.

Funding acquisition: Michelle M. McClure, Mark W. Nelson.

- Investigation: Lisa G. Crozier, David A. Boughton, Mark Carr, Thomas D. Cooney, Jason B. Dunham, Melissa A. Haltuch, Elliott L. Hazen, David D. Huff, Chris E. Jordan, Isaac C. Kaplan, Nathan J. Mantua, Peter B. Moyle, James M. Myers, Mark W. Nelson, Brian C. Spence, Laurie A. Weitkamp, Thomas H. Williams, Ellen Willis-Norton.
- Methodology: Lisa G. Crozier, David A. Boughton, Melissa A. Haltuch, Nathan J. Mantua, Mark W. Nelson, Ellen Willis-Norton.
- **Project administration:** Lisa G. Crozier, Michelle M. McClure, Steven J. Bograd, Melissa A. Haltuch, Elliott L. Hazen, Mark W. Nelson.

Supervision: Michelle M. McClure, Steven J. Bograd, Steven T. Lindley.

- Visualization: Lisa G. Crozier, Correigh M. Greene, Damon M. Holzer.
- Writing original draft: Lisa G. Crozier, Tim Beechie, David A. Boughton, Thomas D. Cooney, Correigh M. Greene, David D. Huff, Rachel C. Johnson, Isaac C. Kaplan, Nathan J. Mantua, Peter B. Moyle, James M. Myers, Mark W. Nelson, Brian C. Spence, Thomas H. Williams, Ellen Willis-Norton.
- Writing review & editing: Lisa G. Crozier, Michelle M. McClure, Tim Beechie, Steven J. Bograd, David A. Boughton, Mark Carr, Thomas D. Cooney, Jason B. Dunham, Correigh M. Greene, Melissa A. Haltuch, Elliott L. Hazen, David D. Huff, Rachel C. Johnson, Isaac C. Kaplan, Steven T. Lindley, Nathan J. Mantua, Peter B. Moyle, James M. Myers, Mark W. Nelson, Brian C. Spence, Laurie A. Weitkamp, Thomas H. Williams, Ellen Willis-Norton.

#### References

- 1. Wiens JJ. Climate-related local extinctions are already widespread among plant and animal species. PLoS Biol. 2016; 14(12):e2001104. <u>https://doi.org/10.1371/journal.pbio.2001104</u> PMID: 27930674
- IPCC, Intergovernmental Panel on Climate Change. Climate change 2014: Synthesis report. Contribution of working groups I, II and III to the fifth assessment report of the Intergovernmental Panel on Climate Change. (eds.) RKPaLAM, editor. Geneva, Switzerland: IPCC; 2014. 151 p.
- 3. Carter S, Peterson J, Lipton D, Crozier L, Fogarty M, Gaichas S, et al. Ecosystems, Ecosystem Services, and Biodiversity. Climate Change Impacts in the United States: The Fourth National Climate Assessment: U.S. Global Change Research Program; 2018.
- USGCRP, U.S. Global Change Research Program. Climate Science Special Report: Fourth National Climate Assessment, Volume I. Wuebbles DJ, Fahey D.W., Hibbard K.A., Dokken D.J., Stewart B.C., and Maycock T.K., editor. Washington, DC: U.S. Global Change Research Program; 2017. 470 p.
- Pacifici M, Foden WB, Visconti P, Watson JFM, Butchart SHM, Kovacs KM, et al. Assessing species vulnerability to climate change. Nature Climate Change. 2015; 5(3):215–25. <u>https://doi.org/10.1038/nclimate2448</u> WOS:000350327800015.
- 6. Glick P, Stein BA, Edelson NA, editors. Scanning the conservation horizon: a guide to climate change vulnerability assessment. Washington, D.C.: National Wildlife Federation 2011.
- Williams JG, Zabel RW, Waples RS, Hutchings JA, Connor WP. Potential for anthropogenic disturbances to influence evolutionary change in the life history of a threatened salmonid. Evol Appl. 2008; 1 (2):271–85. <u>https://doi.org/10.1111/j.1752-4571.2008.00027.x</u> PMID: 25567631
- Healey M, Price A. Scales of variation in life history tactics of Pacific salmon and the conservation of genotype and phenotype. Am Fish Soc Symp. 1995; 17:176–84.
- Waples RS. Pacific salmon, Oncorhynchus spp., and the definition of "species" under the Endangered Species Act. Marine Fisheries Review. 1991; 53(11–22).
- Foden EWB, Young BE. IUCN SSC guidelines for assessing species' vulnerability to climate change Version 1.0. Cambridge, UK: Occasional paper of the IUCN Species Survival Commission, 2016.

- 11. Behnke RJ. Native trout of western North America. Bethesda, Maryland, USA1992.
- NMFS, National Marine Fisheries Service. Northwest Region. Status of ESA listings and critical habitat designations for West Coast salmon and steelhead, 2016. Available from <u>www.westcoast.fisheries.</u> <u>noaa.gov/publications/gis\_maps/maps/salmon\_steelhead/critical\_habitat/wcr\_salmonid\_ch\_esa\_july2016.pdf</u> (November 2018). 2016.
- 13. NMFS, National Marine Fisheries Service. West Coast salmon & steelhead listings: National Marine Fisheries Service, West Coast Region. Available at <a href="http://www.westcoast.fisheries.noaa.gov/protected\_species/salmon\_steelhead/salmon\_and\_steelhead\_listings/salmon\_and\_steelhead\_listings.html">www.westcoast.fisheries.noaa.gov/protected\_species/salmon\_steelhead/salmon\_and\_steelhead\_listings/salmon\_and\_steelhead\_listings.html</a> 2016.
- 14. Ford MJ, Barnas K, Cooney T, Crozier LG, Diaz M, Hard JJ, et al. Status review update for Pacific salmon and steelhead listed under the Endangered Species Act: Pacific Northwest. Seattle, WA. Available from <u>www.nwfsc.noaa.gov/publications/scipubs/searchdoc.cfm</u>: U.S. Dep Commerce NOAA Fisheries Northwest Fisheries Science Center, 2015.
- Lindley ST, Grimes CB, Mohr MS, Peterson W, Stein J, Anderson JT, et al. What caused the Sacramento River fall Chinook stock collapse? Santa Cruz, CA: U.S. Bur Commerce NOAA-TM-NMFS-SWFSC-447, 2009 NOAA-TM-NMFS-SWFSC-447.
- Williams TH, Spence BC, Boughton DA, Johnson RC, Crozier LG, Mantua NJ, et al. Viability assessment for Pacific salmon and steelhead listed under the Endangered Species Act: Southwest Santa Cruz, CA: U.S. Dep Commerce NOAA Tech Memo NMFS SWFSC 564, 2016.
- Ward EJ, Anderson JH, Beechie TJ, Pess GR, Ford MJ. Increasing hydrologic variability threatens depleted anadromous fish populations. Glob Chang Biol. 2015; 21(7):2500–9. Epub 2015/02/04. <u>https://doi.org/10.1111/gcb.12847</u> PMID: 25644185.
- Riddell BE, Brodeur RD, Alexander V. Bugaev, Moran P, Murphy JsM, Orsi JhA, et al. The ocean ecology of Chinook salmon In: Beamish RJ, editor. The ocean ecology of Pacific salmon and trout. Bethesda, MD: American Fisheries Society; 2018. p. 555–696.
- Quinn TP. The Behavior and Ecology of Pacific Salmon and Trout. 2nd ed. Seattle, Washington: University of Washington Press; 2018.
- Taylor EB. Environmental correlates of life-history variation in juvenile chinook salmon, Oncorhynchus tshawytscha (Walbaum). J Fish Biol. 1990; 37(1):1–17.
- Busby PJ, Wainwright TC, Bryant GJ, Lierheimer LJ, Waples RS, Waknitz FW, et al. Status review of west coast steelhead from Washington, Idaho, Oregon, and California. Seattle, WA: U.S. Dep Commerce NOAA Tech Memo NMFS-NWFSC-27, 1996.
- Waples RS, Teel DJ, Myers JM, Marshall AR. Life-history divergence in Chinook salmon: historic contingency and parallel evolution. Evolution. 2004; 58(2):2004.
- Drinan DP, Zale AV, Webb MAH, Shepard BB, Kalinowski ST. Evidence of local adaptation in westslope cutthroat trout. Trans Am Fish Soc. 2012; 141(4):872–80. <u>https://doi.org/10.1080/00028487.</u> 2012.675907 WOS:000306462500002.
- 24. Hess JE, Narum SR. Single-nucleotide polymorphism (SNP) loci correlated with run timing in adult Chinook salmon from the Columbia River basin. Trans Am Fish Soc. 2011; 140(3):855–64. <u>https://doi.org/10.1080/00028487.2011.588138</u> WOS:000291813800027.
- Larson WA, Seeb JE, Dann TH, Schindler DE, Seeb LW. Signals of heterogeneous selection at an MHC locus in geographically proximate ecotypes of sockeye salmon. Molecular Ecology. 2014; 23 (22):5448–61. <u>https://doi.org/10.1111/mec.12949</u> WOS:000345572300005. PMID: <u>25283474</u>
- Narum SR, Campbell NR, Meyer KA, Miller MR, Hardy RW. Thermal adaptation and acclimation of ectotherms from differing aquatic climates. Molecular Ecology. 2013; 22(11):3090–7. <u>https://doi.org/ 10.1111/mec.12240</u> WOS:000319675300019. PMID: <u>23452191</u>
- Berejikian BA, Campbell LA, Moore ME. Large-scale freshwater habitat features influence the degree of anadromy in eight Hood Canal *Oncorhynchus mykiss* populations. Can J Fish Aquat Sci. 2013; 70 (5):756–65. <u>https://doi.org/10.1139/cjfas-2012-0491</u> WOS:000319257700010.
- Keefer ML, Peery CA, Caudill CC. Migration timing of Columbia River spring Chinook salmon: Effects of temperature, river discharge, and ocean environment. Trans Am Fish Soc. 2008; 137(4):1120–33. <u>https://doi.org/10.1577/t07-008.1</u> ISI:000258163500016.
- Reed TE, Martinek G, Quinn TP. Lake-specific variation in growth, migration timing and survival of juvenile sockeye salmon *Oncorhynchus nerka*: Separating environmental from genetic influences. J Fish Biol. 2010; 77(3):692–705. <u>https://doi.org/10.1111/j.1095-8649.2010.02711.x</u> ISI:000280643900016. PMID: <u>20701648</u>
- Crozier LG, Hutchings JA. Plastic and evolutionary responses to climate change in fish. Evol Appl. 2014; 7(1):68–87. <u>https://doi.org/10.1111/eva.12135</u> PMID: <u>24454549</u>

- Crozier LG, Scheuerell MD, Zabel RW. Using time series analysis to characterize evolutionary and plastic responses to environmental change: A case study of a shift toward earlier migration date in sockeye salmon. Am Nat. 2011; 178(6):755–73. <u>https://doi.org/10.1086/662669</u> ISI:000297503800009. PMID: 22089870
- Reed TE, Schindler DE, Hague MJ, Patterson DA, Meir E, Waples RS, et al. Time to evolve? Potential evolutionary responses of Fraser River sockeye salmon to climate change and effects on persistence. PLoS ONE. 2011; 6(6):e20380. <u>https://doi.org/10.1371/journal.pone.0020380</u> WOS:000292142800002. PMID: 21738573
- Gomulkiewicz R, Holt RD. When does evolution by natural selection prevent extinction? Evolution. 1995; 49(1):201–7. <u>https://doi.org/10.1111/j.1558-5646.1995.tb05971.x</u> PMID: <u>28593677</u>
- 34. McElhany P, Ruckelshaus MH, Ford MJ, Wainwright TC, Bjorkstedt EP. Viable salmonid populations and the recovery of Evolutionarily Significant Units. Seattle, WA: National Marine Fisheries Service, Northwest Fisheries Science Center, 2000 Technical Memorandum NMFS-NWFSC 42.
- Hare JA, Morrison WE, Nelson MW, Stachura MM, Teeters EJ, Griffis RB, et al. A vulnerability assessment of fish and invertebrates to climate change on the northeast U.S. continental shelf. PLoS ONE. 2016; 11(2):e0146756. <u>https://doi.org/10.1371/journal.pone.0146756</u> PMID: 26839967
- Link JS, Griffis R, Busch S. NOAA Fisheries climate science strategy. Silver Spring, MD: National Marine Fisheries Service, 2015 NOAA Technical Memorandum NMFS-F/SPO-155.
- Hanson KC, Peterson DP. Modeling the potential impacts of climate change on Pacific salmon culture programs: An example at Winthrop National Fish Hatchery. Environ Manage. 2014; 54(3):433–48. <u>https://doi.org/10.1007/s00267-014-0302-2</u> WOS:000340538100005. PMID: 24993792
- Martin BT, Pike A, John SN, Hamda N, Roberts J, Lindley ST, et al. Phenomenological vs. biophysical models of thermal stress in aquatic eggs. Ecol Lett. 2016; 20(1):50–9. <u>https://doi.org/10.1111/ele.</u> 12705 PMID: 27891770
- Mathis JT, Cross JN, Evans W, Doney SC. Ocean acidification in the surface waters of the Pacific-Arctic boundary regions. Oceanography. 2015; 28(2):122–35. <u>https://doi.org/10.5670/oceanog.2015.36</u> WOS:000357231700014.
- Busch DS, Harvey CJ, McElhany P. Potential impacts of ocean acidification on the Puget Sound food web. ICES J Mar Sci. 2013; 70(4):823–33. <u>https://doi.org/10.1093/icesjms/fst061</u>
- 41. Busch DS, McElhany P. Estimates of the direct effect of seawater pH on the survival rate of species groups in the California Current ecosystem. PLoS ONE. 2016; 11(8):e0160669. Epub 2016/08/12. <u>https://doi.org/10.1371/journal.pone.0160669</u> PMID: <u>27513576</u>; PubMed Central PMCID: PMC4981315.
- 42. Wells BK, Santora JA, Field JC, MacFarlane RB, Marinovic BB, Sydeman WJ. Population dynamics of Chinook salmon Oncorhynchus tshawytscha relative to prey availability in the central California coastal region. Mar Ecol Prog Ser. 2012; 457:125–37. https://doi.org/10.3354/meps09727
- Munday PL, Dixson DL, Donelson JM, Jones GP, Pratchett MS, Devitsina GV, et al. Ocean acidification impairs olfactory discrimination and homing ability of a marine fish. Proceedings of the National Academy of Sciences of the United States of America. 2009; 106(6):1848–52. <u>https://doi.org/10.1073/</u> pnas.0809996106 ISI:000263252500033. PMID: <u>19188596</u>
- Ou M, Hamilton TJ, Eom J, Lyall EM, Gallup J, Jiang A, et al. Responses of pink salmon to CO2induced aquatic acidification. Nature Climate Change. 2015; 5(10). <u>https://doi.org/10.1038/ nclimate2694</u> WOS:000361840600017.
- 45. Morrison W, Nelson M, Howard J, Teeters E, Hare JA, Griffis R. Methodology for assessing the vulnerability of fish stocks to changing climate. Natinoal Marine Fisheries Service, Office of Sustainable Fisheries, 2015 Accessed 1/1/2017. Report No.: NOAA Technical Memorandum NMFS-OSF-3.
- 46. Bell G. Evolutionary rescue and the limits of adaptation. Philosophical Transactions of the Royal Society B: Biological Sciences. 2013; 368(1610):20120080. <u>https://doi.org/10.1098/rstb.2012.0080</u> PMID: 23209162
- Kopp M, Matuszewski S. Rapid evolution of quantitative traits: theoretical perspectives. Evol Appl. 2014; 7(1):169–91. <u>https://doi.org/10.1111/eva.12127</u> WOS:000332699400012. PMID: <u>24454555</u>
- Kalinowski ST, Van Doornik DM, Kozfkay CC, Waples R. Genetic diversity in the Snake River sockeye salmon captive broodstock program as estimated from broodstock records. Conserv Genet. 2012; 13:1183–93.
- Knudsen CM, Schroder SL, Busack CA, Johnston MV, Pearsons TN, Bosch WJ, et al. Comparison of life history traits between first-generation hatchery and wild upper Yakima river spring Chinook salmon. Trans Am Fish Soc. 2006; 135(4):1130–44. <u>https://doi.org/10.1577/t05-121.1</u> WOS:000240325200025.

- Waples RS, Hindar K, Karlsson S, Hard JJ. Evaluating the Ryman–Laikre effect for marine stock enhancement and aquaculture. Current Zoology. 2016; 62(6):617–27. <u>https://doi.org/10.1093/cz/ zow060</u> %J Current Zoology. PMID: <u>29491949</u>
- Christie MR, Marine ML, French RA, Waples RS, Blouin MS. Effective size of a wild salmonid population is greatly reduced by hatchery supplementation. Heredity. 2012; 109(4):254–60. <u>https://doi.org/ 10.1038/hdy.2012.39</u> WOS:000309109700008. PMID: <u>22805657</u>
- Gall GAE, Bartley D, Bentley B, Brodziak J, Gomulkiewicz R, Mangel M. Geographic variation in population genetic structure of Chinook salmon from California and Oregon. Fish Bull. 1992; 90(1):77–100. WOS:A1992HT86200004.
- National Research Council, Committee on Protection and Management of Pacific Northwest Anadromous Salmonids. Upstream: salmon and society in the Pacific Northwest. Washington, D.C.: National Academy Press; 1996.
- 54. Waples RS, Beechie T, Pess GR. Evolutionary history, habitat disturbance regimes, and anthropogenic changes: What do these mean for resilience of Pacific salmon populations? Ecol Soc. 2009; 14 (1):3. 3. ISI:000267846300044.
- 55. Beechie T, Imaki H, Greene J, Wade A, Wu H, Pess G, et al. Restoring salmon habitat for a changing climate. River Res Appl. 2013; 29(8):939–60. <u>https://doi.org/10.1002/rra.2590</u>
- 56. StreamNet. Fish data for the Northwest. <u>http://www.streamnet.org/</u>, accessed January 2010: The Pacific State Marine Fisheries Comission, 2010 Contract No.: Jan 2005.
- Myers KW, Aydin KY, Walker RV, Fowler S, Dahlberg ML. Known ocean ranges of stocks of Pacific salmon and steelhead as shown by tagging experiments, 1956–1995. Fisheries Research Institute, University of Washington School of Fisheries, 1996 FRI-UW-9614.
- Weitkamp LA. Marine distributions of Chinook salmon from the West Coast of North America determined by coded wire tag recoveries. Trans Am Fish Soc. 2010; 139(1):147–70. <u>https://doi.org/10.1577/T08-225.1</u>
- 59. Beamish RJ, editor. The Ocean Ecology of Pacific Salmon and Trout: American Fisheries Society; 2018.
- 60. McCullough DA. A review and synthesis of effects of alterations to the water temperature regime on freshwater life stages of salmonids, with special reference to Chinook salmon. Seattle, Washington: U.S. Environmental Protection Agency, Region 10, 1999.
- Richter A, Kolmes SA. Maximum temperature limits for chinook, coho, and chum salmon, and steelhead trout in the Pacific Northwest. Rev Fish Sci. 2005; 13(1):23–49. ISI:000228086500002.
- Strange JS. Migration strategies of adult Chinook salmon runs in response to diverse environmental conditions in the Klamath River basin. Trans Am Fish Soc. 2012; 141(6):1622–36. <u>https://doi.org/10. 1080/00028487.2012.716010</u> WOS:000315216000018.
- Isaak DJ. Norwest: Regional database and modeled stream temperature. Data tool of the U.S. Department of Agriculture and U.S. Forest Service. Available from <u>wwwfsfedus/rm/boise/AWAE/projects/</u> NorWeSThtml (March 2016). 2016.
- Pike A, Danner E, Boughton D, Melton F, Nemani R, Rajagopalan B, et al. Forecasting river temperatures in real time using a stochastic dynamics approach. Wat Res Res, 2013; 49(9):5168–82. <u>https:// doi.org/10.1002/wrcr.20389</u> WOS:000325991100005.
- 65. Littell JS, Mauger GS, Salathé EP, Hamlet AF, Lee S, Stumbaugh MR, et al. Uncertainty and extreme events in future climate and hydrologic projections for the Pacific Northwest: Providing a basis for vulnerability and core/corridor assessments. Seattle, WA: College of the Environment Climate Impacts Group, the University of Washington 2014 Available from <a href="http://cses.washington.edu/db/pubs/abstract825.shtml">http://cses.washington.edu/db/pubs/abstract825.shtml</a>. Report No.: Final report fot the U.S. for Department of the Interior Pacific Northwest Climate Science Center.
- Salathé EP Jr., Hamlet AF, Mass CF, Lee S-Y, Stumbaugh M, Steed R. Estimates of twenty-first-century flood risk in the Pacific Northwest based on regional climate model simulations. Journal of Hydrometeorology. 2014; 15(5):1881–99. <u>https://doi.org/10.1175/jhm-d-13-0137.1</u> WOS:000342964400010.
- Warner M, Mass CF, Salathé EP. Changes in winter atmospheric rivers along the North American west coast in CMIP5 climate models. J Hydrometeorology. 2015; 16:118–28. <u>https://doi.org/10.1175/ JHM-D-14-0080.1</u>
- Beechie T, Buhle E, Ruckelshaus M, Fullerton A, Holsinger L. Hydrologic regime and the conservation of salmon life history diversity. Biol Conserv. 2006; 130(4):560–72. <u>https://doi.org/10.1016/j.biocon.</u> 2006.01.019
- Waples RS, Gustafson RG, Weitkamp LA, Myers JM, Johnson OW, Busby PJ, et al. Characterizing diversity in salmon from the Pacific Northwest. J Fish Biol. 2001; 59:1–41. ISI:000173541100001.

- Hamlet AF, Elsner MM, Mauger GS, Lee SY, Tohver I, Norheim RA. An overview of the Columbia Basin climate change scenarios project: Approach, methods, and summary of key results. Atmosphere-Ocean. 2013; 51(4):392–415. <u>https://doi.org/10.1080/07055900.2013.819555</u> WOS:000323508500005.
- Kilduff DP, Botsford LW, Teo SLH. Spatial and temporal covariability in early ocean survival of Chinook salmon (*Oncorhynchus tshawytscha*) along the west coast of North America. ICES Journal of Marine Science: Journal du Conseil. 2014; 71(7):1671–82. <u>https://doi.org/10.1093/icesjms/fsu031</u>
- 72. Stachura MM, Mantua NJ, Scheuerell MD. Oceanographic influences on patterns in North Pacific salmon abundance. Can J Fish Aquat Sci. 2014; 71(2):226–35. <u>https://doi.org/10.1139/cjfas-2013-0367</u> WOS:000331451100005.
- **73.** Mueter FJ, Boldt JL, Megrey BA, Peterman RM. Recruitment and survival of Northeast Pacific Ocean fish stocks: temporal trends, covariation, and regime shifts. Can J Fish Aquat Sci. 2007; 64(6):911–27. https://doi.org/10.1139/f07-069 ISI:000248572900007.
- 74. Abdul-Aziz OI, Mantua NJ, Myers KW. Potential climate change impacts on thermal habitats of Pacific salmon (*Oncorhynchus spp.*) in the North Pacific Ocean and adjacent seas. Can J Fish Aquat Sci. 2011; 68(9):1660–80. <u>https://doi.org/10.1139/f2011-079</u> WOS:000296169300012.
- 75. Cheung WWL, Brodeur RD, Okey TA, Pauly D. Projecting future changes in distributions of pelagic fish species of Northeast Pacific shelf seas. Prog Oceanogr. 2015; 130:19–31. <u>https://doi.org/10. 1016/j.pocean.2014.09.003</u>
- 76. Morley JW, Selden RL, Latour RJ, Frölicher TL, Seagraves RJ, Pinsky ML. Projecting shifts in thermal habitat for 686 species on the North American continental shelf. PLoS ONE. 2018; 13(5):e0196127. <u>https://doi.org/10.1371/journal.pone.0196127</u> PMID: <u>29768423</u>
- 77. Mantua NJ, Hare SR, Zhang Y, Wallace JM, Francis RC. A Pacific interdecadal climate oscillation with impacts on salmon production. Bull Am Meteorol Soc. 1997; 78:1069–79.
- 78. Kilduff DP, Di Lorenzo E, Botsford LW, Teo SLH. Changing central Pacific El Niños reduce stability of North American salmon survival rates. Proceedings of the National Academy of Sciences of the United States of America. 2015; 112(35):10962–6. <u>https://doi.org/10.1073/pnas.1503190112</u> WOS:000360383200051. PMID: <u>26240365</u>
- 79. Johnstone JA, Mantua NJ. Atmospheric controls on northeast Pacific temperature variability and change, 1900–2012. Proceedings of the National Academy of Sciences of the United States of America. 2014; 111(40):14360–5. <u>https://doi.org/10.1073/pnas.1318371111</u> WOS:000342633900029. PMID: 25246555
- Newman M, Alexander MA, Ault TR, Cobb KM, Deser C, Lorenzo ED, et al. The Pacific decadal oscillation, revisited. Journal of Climate. 2016; 29(12):4399–427. <u>https://doi.org/10.1175/jcli-d-15-0508.1</u>
- Di Lorenzo E, Combes V, Keister J, Strub PT, Thomas A, Franks P, et al. Synthesis of Pacific Ocean climate and ecosystem dynamics. Oceanography. 2013; 26(4):68–81. <u>https://doi.org/10.5670/ oceanog.2013.76</u>
- Overland JE, Wang M. Future climate of the north Pacific Ocean. Eos, Transactions American Geophysical Union. 2007; 88(16):178–82. <u>https://doi.org/10.1029/2007EO160003</u>
- Burke BJ, Peterson WT, Beckman BR, Morgan C, Daly EA, Litz M. Multivariate models of adult Pacific salmon returns. PLoS ONE. 2013; 8(1):e54134. <u>https://doi.org/10.1371/journal.pone.0054134</u> PMID: <u>23326586</u>; PubMed Central PMCID: PMC3543311.
- Malick MJ, Cox SP, Mueter FJ, Dorner B, Peterman RM. Effects of the North Pacific current on the productivity of 163 Pacific salmon stocks. Fish Oceanogr. 2017; 26(3):268–81. <u>https://doi.org/10.1111/ fog.12190</u>
- Rupp DE, Wainwright TC, Lawson PW, Peterson WT. Marine environment-based forecasting of coho salmon (*Oncorhynchus kisutch*) adult recruitment. Fish Oceanogr. 2012; 21(1):1–19. <u>https://doi.org/ 10.1111/j.1365-2419.2011.00605.x</u> WOS:000297794600001.
- Sydeman WJ, Thompson SA, Field JC, Peterson WT, Tanasichuk RW, Freeland HJ, et al. Does positioning of the North Pacific current affect downstream ecosystem productivity? Geophys Res Lett. 2011; 38:L12606. <u>https://doi.org/10.1029/2011GL047212</u>
- Miller JA, Teel DJ, Peterson WT, Baptista AM. Assessing the relative importance of local and regional processes on the survival of a threatened salmon population. PLoS ONE. 2014; 9(6). <u>https://doi.org/ 10.1371/journal.pone.0099814</u> WOS:000338701300096. PMID: <u>24924741</u>
- Ruzicka JJ, Brink KH, Gifford DJ, Bahr F. A physically coupled end-to-end model platform for coastal ecosystems: Simulating the effects of climate change and changing upwelling characteristics on the Northern California Current ecosystem. Ecol Model. 2016; 331:86–99. <u>https://doi.org/10.1016/j. ecolmodel.2016.01.018</u> WOS:000376831700009.

- Wells BK, Santora JA, Schroeder ID, Mantua N, Sydeman WJ, Huff DD, et al. Marine ecosystem perspectives on Chinook salmon recruitment: A synthesis of empirical and modeling studies from a California upwelling system. Mar Ecol Prog Ser. 2016; 552:271–84. <u>https://doi.org/10.3354/meps11757</u> WOS:000379812200019.
- Glick P, Clough J, Nunley B. Sea-level rise and coastal habitat in the Pacific northwest: An analysis for Puget Sound, southwestern Washington, and northwestern Oregon. Washington, D.C.: National Wildlife Federation, 2007.
- Yang Z, Wang T, Voisin N, Copping A. Estuarine response to river flow and sea-level rise under future climate change and human development. Estuarine Coastal and Shelf Science. 2015; 156:19–30. <u>https://doi.org/10.1016/j.ecss.2014.08.015</u> WOS:000354151700003.
- McClure MM, Alexander M, Borggaard D, Boughton D, Crozier L, Griffis R, et al. Incorporating climate science in applications of the US endangered species act for aquatic species. Conserv Biol. 2013; 27 (6):1222–33. https://doi.org/10.1111/cobi.12166 PMID: 24299088.
- 93. Bednarsek N, Feely RA, Reum JCP, Peterson B, Menkel J, Alin SR, et al. *Limacina helicina* shell dissolution as an indicator of declining habitat suitability owing to ocean acidification in the California Current Ecosystem. Proceedings of the Royal Society B-Biological Sciences. 2014; 281(1785). <u>https://doi.org/10.1098/rspb.2014.0123</u> WOS:000335812100011. PMID: <u>24789895</u>
- Bednarsek N, Ohman MD. Changes in pteropod distributions and shell dissolution across a frontal system in the California Current System. Mar Ecol Prog Ser. 2015; 523:93–103. <u>https://doi.org/10.3354/meps11199</u> WOS:000351452700008.
- 95. Feely RA, Alin SR, Carter B, Bednarsek N, Hales B, Chan F, et al. Chemical and biological impacts of ocean acidification along the west coast of North America. Estuarine Coastal and Shelf Science. 2016; 183:260–70. <u>https://doi.org/10.1016/j.ecss.2016.08.043</u> WOS:000390726900023.
- NOAA, National Oceanic and Atmospheric Administration, NOAA Earth System Research Laboratory, Climate Change web portal, CMIP5 maps. Available from <u>esrl.noaa.gov/psd/ipcc/ocn/ccwp.html</u> (November 2018). 2018.
- 97. National Research Council. Sea-level rise for the coasts of California, Oregon, and Washington: Past, present, and future. Committee on Sea Level Rise in California O, and Washington, Sciences BoE, Board ROS, Studies DoEL, Council NR, editors: The National Academies Press 2012.
- Rykaczewski RR, Dunne JP, Sydeman WJ, Garcia-Reyes M, Black BA, Bograd SJ. Poleward displacement of coastal upwelling-favorable winds in the ocean's eastern boundary currents through the 21st century. Geophys Res Lett. 2015; 42(15):6424–31. <u>https://doi.org/10.1002/2015gl064694</u> WOS:000360414900036.
- 99. Bakun A, Black BA, Bograd SJ, García-Reyes M, Miller AJ, Rykaczewski RR, et al. Anticipated effects of climate change on coastal upwelling ecosystems. Current Climate Change Reports. 2015; 1(2):85– 93. https://doi.org/10.1007/s40641-015-0008-4
- Francis RC, Hare SR, Hollowed AB, Wooster WS. Effects of interdecadal climate variability on the oceanic ecosystems of the NE Pacific. Fish Oceanogr. 1998; 7(1):1–21.
- King JR, Agostini VN, Harvey CJ, McFarlane GA, Foreman MG, Overland JE, et al. Climate forcing and the California Current ecosystem. ICES J Mar Sci. 2011; 68(6):1199–216.
- 102. Stouffer RJ, Yin J, Gregory JM, Dixon KW, Spelman MJ, Hurlin W, et al. Investigating the causes of the response of the thermohaline circulation to past and future climate changes. Journal of Climate. 2006; 19(8):1365–87.
- 103. Dawson TP, Jackson ST, House JI, Prentice IC, Mace GM. Beyond predictions: biodiversity conservation in a changing climate. Science. 2011; 332(6025):53–8. <u>https://doi.org/10.1126/science.1200303</u> WOS:000289000000035. PMID: <u>21454781</u>
- 104. Nicotra AB, Beever EA, Robertson AL, Hofmann GE, O'Leary J. Assessing the components of adaptive capacity to improve conservation and management efforts under global change. Conserv Biol. 2015; 29(5):1268–78. <u>https://doi.org/10.1111/cobi.12522</u> WOS:000363729100003. PMID: <u>25926277</u>
- 105. Wade AA, Hand BK, Kovach RP, Luikart G, Whited DC, Muhlfeld CC. Accounting for adaptive capacity and uncertainty in assessments of species' climate-change vulnerability. Conserv Biol. 2017; 31 (1):136–49. <u>https://doi.org/10.1111/cobi.12764</u> WOS:000394428100015. PMID: <u>27214122</u>
- 106. Ofori BY, Stow AJ, Baumgartner JB, Beaumont LJ. Influence of adaptive capacity on the outcome of climate change vulnerability assessment. Sci Rep. 2017; 7:12979. <u>https://doi.org/10.1038/s41598-017-13245-y</u> WOS:000412781300047. PMID: <u>29021590</u>
- 107. Lankford AJ, Svancara LK, Lawler JJ, Vierling K. Comparison of climate change vulnerability assessments for wildlife. Wildlife Society Bulletin. 2014; 38(2):386–94. <u>https://doi.org/10.1002/wsb.399</u> WOS:000351431200021.

- Herbold B, Carlson SM, Henery R. Managing for salmon resilience in California's variable and changing climate. San Fran Est Wat Sci. 2018; 16(2). <u>https://doi.org/10.15447/sfews.2018v16iss2art3</u>
- Allendorf FW, Luikart G. Conservation and the Genetics of Populations. Malden, USA: Blackwell; 2007.
- Kovach RP, Muhlfeld CC, Wade AA, Hand BK, Whited DC, DeHaan PW, et al. Genetic diversity is related to climatic variation and vulnerability in threatened bull trout. Global Change Biol. 2015; 21 (7):2510–24. <u>https://doi.org/10.1111/gcb.12850</u> WOS:000356422500006. PMID: <u>25656972</u>
- 111. Ford MJ, Murdoch AR, Hughes MS, Seamons TR, LaHood ES. Broodstock history strongly influences natural spawning success in hatchery steelhead (*Oncorhynchus mykiss*). PLoS ONE. 2016; 11(10): e0164801. <u>https://doi.org/10.1371/journal.pone.0164801</u> WOS:000385505800132. PMID: <u>27737000</u>
- 112. Gomulkiewicz R, Kingsolver JG, Carter PA, Heckman N. Variation and evolution of function-valued traits. In: Futuyma DJ, editor. Annual Review of Ecology, Evolution, and Systematics, Vol 49. Annual Review of Ecology Evolution and Systematics. 492018. p. 139–64.
- 113. Pelletier F, Coltman DW. Will human influences on evolutionary dynamics in the wild pervade the Anthropocene? BMC Biol. 2018; 16. <u>https://doi.org/10.1186/s12915-017-0476-1</u> WOS:000422724000001. PMID: 29334950
- 114. NMFS, National Marine Fisheries Service. West Coast salmon recovery planning and implementation. NOAA Fisheries West Coast Region website. Available from <u>www.westcoast.fisheries.noaa.gov/</u> <u>protected\_species/salmon\_steelhead/recovery\_planning\_and\_implementation</u> (November 2018). 2018.
- 115. Maechler M, Rousseeuw P, Struyf A, Hubert M, Hornik K. Cluster: Cluster analysis basics and extensions. R package version 2.0.6. 2017.
- 116. Ripley B. Tree: Classification and regression Trees. R package version 1.0–39. CRAN2018.
- **117.** R Core Team. R: A language and environmental for statistical computing. Version 3.0.1.R Vienna, Austria: Foundation for Statistical Computing; 2013.
- Brown LR, Bennett WA, Wagner RW, Morgan-King T, Knowles N, Feyrer F, et al. Implications for future survival of Delta Smelt from four climate change scenarios for the Sacramento–San Joaquin Delta, California. Estuaries and Coasts. 2013; 36(4):754–74. <u>https://doi.org/10.1007/s12237-013-9585-4</u>
- Keefer ML, Clabough TS, Jepson MA, Naughton GP, Blubaugh TJ, Joosten DC, et al. Thermal exposure of adult Chinook salmon in the Willamette River basin. Journal of Thermal Biology. 2015; 48:11–20. <a href="https://doi.org/10.1016/j.jtherbio.2014.12.002">https://doi.org/10.1016/j.jtherbio.2014.12.002</a> WOS:000350189200002. PMID: 25660625
- 120. Bowerman T, Roumasset A, Keefer ML, Sharpe CS, Caudill CC. Prespawn mortality of female Chinook salmon increases with water temperature and percent hatchery origin. Trans Am Fish Soc. 2017; 147:31–42. <u>https://doi.org/10.1080/00028487.2017.1377110</u>
- 121. Thompson LC, Escobar MI, Mosser CM, Purkey DR, Yates D, Moyle PB. Water management adaptations to prevent loss of spring-run Chinook salmon in California under climate change. Journal of Water Resources Planning and Management. 2012; 138(5):465–78. <u>https://doi.org/10.1061/(asce)wr.</u> 1943-5452.0000194
- 122. Connor WP, Tiffan KF, Chandler JA, Rondorf DW, Arnsberg BD, Anderson KC. Upstream migration and spawning success of Chinook salmon in a highly developed, seasonally warm river system. Rev Fish Sci Aqua. 2018; <u>https://doi.org/10.1080/23308249.2018.1477736</u>
- 123. Keefer ML, Caudill CC. Estimating thermal exposure of adult summer steelhead and fall Chinook salmon migrating in a warm impounded river. Ecol Freshwat Fish. 2015; 25(4):599–611. <u>https://doi.org/10.1111/eff.12238</u>
- 124. Myers J, Jorgensen J, Sorel M, Bond M, Nodine T, Zabel R. Upper Willamette River life cycle modeling and the potential effects of climate change. U.S. Dep Commerce, NOAA Fisheries Northwest Fisheries Science Center, Seattle, Washington, 2018 Report to the U.S. Army Corps of Engineers.
- 125. Pearse DE, Campbell MA. Ancestry and adaptation of rainbow trout in Yosemite National Park. Fisheries. 2018; 43(10):472–84. <u>https://doi.org/10.1002/fsh.10136</u>
- 126. NMFS National Marine Fisheries Service. Coastal multispecies recovery plan. National Marine Fisheries Service, West Coast Region, Santa Rosa, California, 2016.
- 127. Moyle PB, Kiernan JD, Crain PK, Quinones RM. Climate change vulnerability of native and alien freshwater fishes of California: a systematic assessment approach. PLoS ONE. 2013; 8(5):e63883. <u>https://doi.org/10.1371/journal.pone.0063883</u> WOS:000320362700075. PMID: <u>23717503</u>
- Wade A, Beechie T, Fleishman E, Mantua N, Wu H, Kimball J, et al. Steelhead vulnerability to climate change in the Pacific Northwest. J Appl Ecol. 2013; 50:1093–104.

- 129. Wade AA, Hand BK, Kovach RP, Luikart G, Whited DC, Muhlfeld CC. Accounting for adaptive capacity and uncertainty in assessments of species' climate-change vulnerability. Conserv Biol. 2016; 31 (1):136–49. <u>https://doi.org/10.1111/cobi.12764</u> WOS:000394428100015. PMID: <u>27214122</u>
- **130.** WDFW, Washington Department of Fish and Wildlife. Washington's State wildlife action plan: 2015 update. Olympia, Washington: Washington Department of Fish and Wildlife, 2015.
- 131. Logerwell E, Busby M, Carothers C, Cotton S, Duffy-Anderson J, Farley E, et al. Fish communities across a spectrum of habitats in the western Beaufort Sea and Chukchi Sea. Progress in Oceanography. 2015; 136:115–32. <u>https://doi.org/10.1016/j.pocean.2015.05.013</u> WOS:000358626900008.
- 132. Nielsen JL, Ruggerone GT, Zimmerman CE. Adaptive strategies and life history characteristics in a warming climate: Salmon in the Arctic? Environmental Biology of Fishes. 2013; 96(10–11):1187–226. https://doi.org/10.1007/s10641-012-0082-6 WOS:000324559700005.
- 133. Jensen AJ, Karlsson S, Fiske P, Hansen LP, Ostborg GM, Hindar K. Origin and life history of Atlantic salmon (*Salmo salar*) near their northernmost oceanic limit. Canadian Journal of Fisheries and Aquatic Sciences. 2014; 71(11):1740–6. <u>https://doi.org/10.1139/cjfas-2014-0169</u> WOS:000344604800013.
- 134. Wheatley CJ, Beale CM, Bradbury RB, Pearce-Higgins JW, Critchlow R, Thomas CD. Climate change vulnerability for species—Assessing the assessments. Global Change Biol. 2017; 23(9):3704–15. <u>https://doi.org/10.1111/gcb.13759</u> WOS:000406812100023. PMID: <u>28660715</u>
- **135.** Hayes SA, Kocik JF. Comparative estuarine and marine migration ecology of Atlantic salmon and steelhead: blue highways and open plains. Rev Fish Biol Fish. 2014; 24(3):757–80.
- **136.** Sweet WV, Kopp RE, Weaver CP, Obeysekera J, Horton RM, Thieler ER, et al. Global and regional sea level rise scenarios for the United States. National Ocean Service, Center for Operational Oceano-graphic Products and Services, 2017 NOAA Technical Report NOS CO-OPS 083.
- 137. Gao Y, Lu J, Leung R, Yang Q, Hagos S, Qian Y. Dynamical and thermodynamical modulations on future changes of landfalling atmospheric rivers over western North America. Geophys Res Lett. 2015; 42(17):7179–86. <u>https://doi.org/10.1002/2015gl065435</u> WOS:000363411200039.
- 138. Payne AE, Magnusdottir G. An evaluation of atmospheric rivers over the North Pacific in CMIP5 and their response to warming under RCP 8.5. J Geophys Res-Atmos. 2015; 120(21):11173–90. <u>https:// doi.org/10.1002/2015jd023586</u> WOS:000367823600008.
- 139. Barth NA, Villarini G, Nayak M, White K. Mixed populations and annual flood frequency estimates in the western United States: The role of atmospheric rivers. Wat Res Res, 2016; 53:257–69. <u>https://doi.org/10.1002/2016WR019064</u>
- Shields CA, Kiehl JT. Atmospheric river landfall-latitude changes in future climate simulations. Geophys Res Lett. 2016; 43(16):8775–82. <u>https://doi.org/10.1002/2016gl070470</u> WOS:000384443800053.
- Brady RX, Alexander MA, Lovenduski NS, Rykaczewski RR. Emergent anthropogenic trends in California Current upwelling. Geophys Res Lett. 2017; 44(10):5044–52. <u>https://doi.org/10.1002/ 2017gl072945</u> WOS:000404131900079.
- 142. Mote P, Snover AK, Capalbo S, Eigenbrode SD, Glick P, Littell J, et al. Northwest. In: Melillo JM, Richmond TC, Yohe GW, editors. Climate change impacts in the United States: The third national climate assessment: U.S. Global Change Research Program; 2014. p. 487–513.
- Stoelinga MT, Albright MD, Mass CF. A new look at snowpack trends in the Cascade mountains. Journal of Climate. 2010; 23(10):2473–91.
- 144. Vano JA, Nijssen B, Lettenmaier DP. Seasonal hydrologic responses to climate change in the Pacific Northwest. Wat Res Res, 2015; 51(4):1959–76. <u>https://doi.org/10.1002/2014wr015909</u> WOS:000354733500005.
- 145. Boughton DA, Hatch C, Mora E. Identifying distinct thermal components of a creek. Wat Res Res,. 2012; 48(9). https://doi.org/10.1029/2011wr011713
- 146. Witiw MR, LaDochy S. Cool PDO phase leads to recent rebound in coastal southern California fog. Die Erde. 2015; 146(4):232–44. WOS:000368657600002.
- 147. Johnstone JA, Dawson TE. Climatic context and ecological implications of summer fog decline in the coast redwood region. Proc Natl Acad Sci. 2010; 107(10):4533–8. <u>https://doi.org/10.1073/pnas.0915062107</u> PMID: 20160112
- 148. Boughton DA, Fish H, Pope J, Holt G. Spatial patterning of habitat for Oncorhynchus mykiss in a system of intermittent and perennial streams. Ecol Freshwat Fish. 2009; 18(1):92–105. <u>https://doi.org/10.1111/j.1600-0633.2008.00328.x</u>
- Woelfle-Erskine C. Collaborative approaches to flow restoration in intermittent salmon-bearing streams: Salmon Creek, CA, USA. Water. 2017; 9(3). <u>https://doi.org/10.3390/w9030217</u>
- Soria M, Leigh C, Datry T, Bini LM, Bonada N. Biodiversity in perennial and intermittent rivers: a metaanalysis. Oikos. 2017; 126(8):1078–89. <u>https://doi.org/10.1111/oik.04118</u>

- 151. Fritz KM, Hagenbuch E, D'Amico E, Reif M, Wigington PJ, Leibowitz SG, et al. Comparing the extent and permanence of headwater streams from two field surveys to values from hydrographic databases and maps. Journal of the American Water Resources Association. 2013; 49(4):867–82.
- 152. Arismendi I, Safeeq M, Johnson SL, Dunham JB, Haggerty R. Increasing synchrony of high temperature and low flow in western North American streams: double trouble for coldwater biota? Hydrobiologia. 2013; 712(1):61–70.
- 153. Brannon EL, Powell MS, Quinn TP, Talbot A. Population structure of Columbia River Basin Chinook salmon and steelhead trout. Rev Fish Sci. 2004; 12(2–3):99–232. <u>https://doi.org/10.1080/</u> 10641260490280313
- **154.** Kammerer BD, Heppell SA. The effects of semichronic thermal stress on physiological indicators in steelhead. Trans Am Fish Soc. 2013; 142(5):1299–307. <u>https://doi.org/10.1080/00028487.2013.</u> 806349 WOS:000324613800012.
- **155.** Sloat MR, Osterback AMK. Maximum stream temperature and the occurrence, abundance, and behavior of steelhead trout (*Oncorhynchus mykiss*) in a southern California stream. Can J Fish Aquat Sci. 2013; 70(1):64–73. <u>https://doi.org/10.1139/cjfas-2012-0228</u> WOS:00031483300008.
- **156.** Keefer ML, Peery CA, High B. Behavioral thermoregulation and associated mortality trade-offs in migrating adult steelhead (*Oncorhynchus mykiss*): variability among sympatric populations. Can J Fish Aquat Sci. 2009; 66(10):1734–47. https://doi.org/10.1139/f09-131 ISI:000271832800009.
- 157. Boughton DA, Pike AS. Floodplain rehabilitation as a hedge against hydroclimatic uncertainty in a migration corridor of threatened steelhead. Conserv Biol. 2013; 27(6):1158–68. <u>https://doi.org/10.1111/cobi.12169</u> WOS:000327564300005. PMID: <u>24299082</u>
- Prince DJ, O'Rourke SM, Thompson TQ, Ali OA, Lyman HS, Saglam IK, et al. The evolutionary basis of premature migration in Pacific salmon highlights the utility of genomics for informing conservation. 2017; 3(8):e1603198. <u>https://doi.org/10.1126/sciadv.1603198</u> %J Science Advances. PMID: 28835916
- 159. Wainwright TC, Weitkamp LA. Effects of climate change on Oregon coast Coho salmon: Habitat and life-cycle interactions. Northwest Sci. 2013; 87(3):219–42. <u>https://doi.org/10.3955/046.087.0305</u> WOS:000325271800003.
- Fiechter J, Huff DD, Martin BT, Jackson DW, Edwards CA, Rose KA, et al. Environmental conditions impacting juvenile Chinook salmon growth off central California: An ecosystem model analysis. Geophys Res Lett. 2015; 42(8):2910–7. <u>https://doi.org/10.1002/2015gl063046</u> WOS:000354560800044.
- Ruzicka JJ, Brodeur RD, Emmett RL, Steele JH, Zamon JE, Morgan CA, et al. Interannual variability in the northern California Current food web structure: changes in energy flow pathways and the role of forage fish, euphausiids, and jellyfish. Prog Oceanogr. 2012; 102:19–41. <u>https://doi.org/10.1016/j. pocean.2012.02.002</u> WOS:000307421600003.
- 162. Beaugrand G, Conversi A, Chiba S, Edwards M, Fonda-Umani S, Greene C, et al. Synchronous marine pelagic regime shifts in the Northern Hemisphere. Philosophical Transactions of the Royal Society B-Biological Sciences. 2015; 370(1659). <u>https://doi.org/10.1098/rstb.2013.0272</u> WOS:000346147200010.
- 163. Koenigstein S, Mark FC, Gossling-Reisemann S, Reuter H, Poertner HO. Modelling climate change impacts on marine fish populations: process-based integration of ocean warming, acidification and other environmental drivers. Fish and Fisheries. 2016; 17(4):972–1004. <u>https://doi.org/10.1111/faf.</u> 12155 WOS:000386938900004.
- 164. Vert-pre KA, Amoroso RO, Jensen OP, Hilborn R. Frequency and intensity of productivity regime shifts in marine fish stocks. Proc Natl Acad Sci. 2013; 110(5):1779–84. <u>https://doi.org/10.1073/pnas.</u> <u>1214879110</u> PMID: 23322735
- 165. Zabel RW, Scheuerell MD, McClure MM, Williams JG. The interplay between climate variability and density dependence in the population viability of Chinook salmon. Conserv Biol. 2006; 20(1):190–200. PMID: <u>16909672</u>
- 166. Daly EA, Brodeur RD, Auth TD. Anomalous ocean conditions in 2015: impacts on spring Chinook salmon and their prey field. Mar Ecol Prog Ser. 2017; 566:169–82. <u>https://doi.org/10.3354/meps12021</u> WOS:000396051300013.
- 167. Waples RS, Elz A, Arnsberg BD, Faulkner JR, Hard JJ, Timmins-Schiffman E, et al. Human-mediated evolution in a threatened species? Juvenile life-history changes in Snake River salmon. Evol Appl. 2017; 10(7):667–81. Epub 2017/07/19. <u>https://doi.org/10.1111/eva.12468</u> PMID: <u>28717387</u>; PubMed Central PMCID: PMC5511361.
- 168. Ford MJ, Fuss H, Boelts B, LaHood E, Hard J, Miller J. Changes in run timing and natural smolt production in a naturally spawning coho salmon (Oncorhynchus kisutch) population after 60 years of intensive hatchery supplementation. Can J Fish Aquat Sci. 2006; 63(10):2343–55. <u>https://doi.org/10. 1139/f06-119</u> WOS:000241585400019.

- 169. Puget Sound Regional Council. Vision 2050: Draft 2050 forecast of people and jobs. Seattle, WA: 2018.
- 170. Parkinson E, Perrin C, Ramos-Espinoza D, Taylor E. Evidence for freshwater residualism in coho salmon, *Oncorhynchus kisutch*, from a watershed on the North Coast of British Columbia. Can Field-Nat. 2017; 130(4):336–43. <u>https://doi.org/10.22621/cfn.v130i4.1928</u>
- 171. Perales KM, Rowan J, Moyle PB. Evidence of landlocked Chinook salmon populations in California. N Am J Fish Manage. 2015; 35:1101–5. <u>https://doi.org/10.1080/02755947.2015.1082518</u>
- 172. Romer JD, Monzyk FR. Adfluvial life history in spring Chinook salmon from Quartzville Creek, Oregon. N Am J Fish Manage. 2014; 34(5):885–91.
- 173. Sproles EA, Roth TR, Nolin AW. Future snow? A spatial-probabilistic assessment of the extraordinarily low snowpacks of 2014 and 2015 in the Oregon Cascades. The Cryosphere. 2017; 11(1):331–41. <u>https://doi.org/10.5194/tc-11-331-2017</u>
- 174. Kuehne LM, Olden JD, Duda JJ. Costs of living for juvenile Chinook salmon (*Oncorhynchus tshawytscha*) in an increasingly warming and invaded world. Canadian Journal of Fisheries and Aquatic Sciences. 2012; 69(10):1621–30. <u>https://doi.org/10.1139/f2012-094</u> WOS:000309959900006.
- 175. Lawrence DJ, Olden JD, Torgersen CE. Spatiotemporal patterns and habitat associations of smallmouth bass (*Micropterus dolomieu*) invading salmon-rearing habitat. Freshwater Biology. 2012; 57 (9):1929–46. <u>https://doi.org/10.1111/j.1365-2427.2012.02847.x</u> WOS:000306736200015.
- 176. Sharma S, Jackson DA, Minns CK. Quantifying the potential effects of climate change and the invasion of smallmouth bass on native lake trout populations across Canadian lakes. Ecography. 2009; 32 (3):517–25. https://doi.org/10.1111/j.1600-0587.2008.05544.x ISI:000267659400017.
- 177. Sanderson BL, Barnas KA, Rub AMW. Nonindigenous species of the Pacific Northwest: an overlooked risk to endangered salmon? BioScience. 2009; 59:245–56.
- 178. Hinrichsen RA, Hasselman DJ, Ebbesmeyer CC, Shields BA. The Role of Impoundments, Temperature, and Discharge on Colonization of the Columbia River Basin, USA, by Nonindigenous American Shad. Trans Am Fish Soc. 2013; 142(4):887–900. <u>https://doi.org/10.1080/00028487.2013.788553</u> WOS:000324613900001.
- 179. Demetras NJ, Huff DD, Michel CJ, Smith JM, Cutter GR, Hayes SA, et al. Development of underwater recorders to quantify predation of juvenile Chinook salmon (*Oncorhynchus tshawytscha*) in a river environment. Fish Bull. 2016; 114(2):179–85. <u>https://doi.org/10.7755/fb.114.2.5</u> WOS:000374724100005.
- Rahel FJ, Olden JD. Assessing the effects of climate change on aquatic invasive species. Conservation Biology. 2008; 22(3):521–33. <u>https://doi.org/10.1111/j.1523-1739.2008.00950.x</u> PMID: <u>18577081</u>
- Ray RA, Holt RA, Bartholomew JL. Relationship between temperature and *Ceratomyxa shasta*induced mortality in Klamath River salmonids. Journal of Parasitology. 2012; 98(3):520–6. <u>https://doi.org/10.1645/JP-GE-2737.1</u> WOS:000305926200011. PMID: 22746389
- 182. Jokinen IE, Salo HM, Markkula E, Rikalainen K, Arts MT, Browman HI. Additive effects of enhanced ambient ultraviolet B radiation and increased temperature on immune function, growth and physiological condition of juvenile (parr) Atlantic Salmon, *Salmo salar*. Fish Shellfish Immunol. 2011; 30(1):102–8. <u>https://doi.org/10.1016/j.fsi.2010.09.017</u> WOS:000286905800013. PMID: 20883792
- 183. Lõhmus M, Björklund M. Climate change: what will it do to fish-parasite interactions? Biological Journal of the Linnean Society. 2015; 116(2):397–411. <u>https://doi.org/10.1111/bij.12584</u> WOS:000361198500013.
- 184. Miller KM, Teffer A, Tucker S, Li S, Schulze AD, Trudel M, et al. Infectious disease, shifting climates, and opportunistic predators: cumulative factors potentially impacting wild salmon declines. Evol Appl. 2014; 7(7):812–55. <u>https://doi.org/10.1111/eva.12164</u> WOS:000341637100009. PMID: <u>25469162</u>
- **185.** Lusardi RA, Moyle PB. Two-way trap and haul as a conservation strategy for anadromous salmonids. 478–487. 2017.
- 186. Lindley ST, Schick R, Mora E, Adams P, Anderson J, Greene S, et al. Framework for assessing the viability of threatened and endangered Chinook salmon and steelhead in the Sacramento-San Joaquin basin. San Fran Est Wat Sci. 2007; 5:1.
- 187. Beever EA, Hall LE, Varner J, Loosen AE, Dunham JB, Gahl MK, et al. Behavioral flexibility as a mechanism for coping with climate change. Front Ecol Environ 2017; 15:299–308.
- 188. Pearse DE, Miller MR, Abadia-Cardoso A, Garza JC. Rapid parallel evolution of standing variation in a single, complex, genomic region is associated with life history in steelhead/rainbow trout. Proceedings Biological sciences. 2014; 281(1783):20140012. Epub 2014/03/29. <u>https://doi.org/10.1098/rspb.2014.</u> 0012 PMID: 24671976; PubMed Central PMCID: PMC3996610.

- 189. Gustafson RG, Waples RS, Myers JM, Weitkamp LA, Bryant GJ, Johnson OW, et al. Pacific salmon extinctions: quantifying lost and remaining diversity. Conserv Biol. 2007; 21(4):1009–20. <u>https://doi.org/10.1111/j.1523-1739.2007.00693.x</u> PMID: <u>17650251</u>
- 190. Sheer MB, Steel EA. Lost watersheds: barriers, aquatic habitat connectivity, and salmon persistence in the Willamette and Lower Columbia River basins. Trans Am Fish Soc. 2006; 135(6):1654–69.
- 191. Sturrock AM, Wikert JD, Heyne T, Mesick C, Hubbard AE, Hinkelman TM, et al. Reconstructing the migratory behavior and long-term survivorship of juvenile Chinook salmon under contrasting hydrologic regimes. PLoS ONE. 2015; 10(5):e0122380. <u>https://doi.org/10.1371/journal.pone.0122380</u> PMID: 25992556
- 192. Anderson SC, Moore JW, McClure MM, Dulvy NK, Cooper AB. Portfolio conservation of metapopulations under climate change. Ecol Appl. 2015; 25(2):559–72. <u>https://doi.org/10.1890/14-0266.1</u> WOS:000350556400020. PMID: 26263675
- 193. Moore JW, Yeakel JD, Peard D, Lough J, Beere M. Life-history diversity and its importance to population stability and persistence of a migratory fish: steelhead in two large North American watersheds. J Anim Ecol. 2014; 83(5):1035–46. <u>https://doi.org/10.1111/1365-2656.12212</u> WOS:000340877700005. PMID: 24673479
- 194. Greene CM, Hall JE, Guilbault KR, Quinn TP. Improved viability of populations with diverse life-history portfolios. Biology Letters. 2010; 6(3):382–6. <u>https://doi.org/10.1098/rsbl.2009.0780</u> WOS:000277559000028. PMID: 20007162
- 195. Erkinaro J, Czorlich Y, Orell P, Kuusela J, Falkegård M, Länsman M, et al. Life history variation across four decades in a diverse population complex of Atlantic salmon in a large subarctic river. Can J Fish Aquat Sci. 2018; 76(1):42–55. https://doi.org/10.1139/cjfas-2017-0343
- 196. Thorson JT, Scheuerell MD, Buhle ER, Copeland T. Spatial variation buffers temporal fluctuations in early juvenile survival for an endangered Pacific salmon. J Anim Ecol. 2014; 83(1):157–67. <u>https://doi.org/10.1111/1365-2656.12117</u> WOS:000328615300017. PMID: 23919254
- 197. Black BA, van der Sleen P, Di Lorenzo E, Griffin D, Sydeman WJ, Dunham JB, et al. Rising synchrony controls western North American ecosystems. Global Change Biol. 2018; 24(6):2305–14.
- 198. Isaak DJ, Thurow RF, Rieman BE, Dunham JB. Temporal variation in synchrony among chinook salmon (*Oncorhynchus tshawytscha*) redd counts from a wilderness area in central Idaho. Can J Fish Aquat Sci. 2003; 60(7):840–8. https://doi.org/10.1139/f03-073 ISI:000185215700008.
- 199. Mantyka-Pringle CS, Martin TG, Rhodes JR. Interactions between climate and habitat loss effects on biodiversity: A systematic review and meta-analysis. Global Change Biol. 2012; 18(4):1239–52. https://doi.org/10.1111/j.1365-2486.2011.02593.x WOS:000301533100004.
- 200. Williams TH, Reeves GH. Ecological diversity and extinction risk of Pacific salmon and trout. In: Wainwright T, MacCall A, editors. Assessing extinction risk for West Coast: NOAA Tech. Memo NMFS-NWFSC-56, National Marine Fisheries Service, Northwest Fisheries Science Center 2003. p. 107–15.
- 201. Roni P, Pess G, Beechie T, Morley S. Estimating changes in Coho salmon and steelhead abundance from watershed restoration: How much restoration is needed to measurably increase smolt production? N Am J Fish Manage. 2010; 30(6):1469–84. <u>https://doi.org/10.1577/m09-162.1</u> WOS:000286421000014.
- 202. Kurylyk BL, MacQuarrie KTB, Linnansaari T, Cunjak RA, Curry RA. Preserving, augmenting, and creating cold-water thermal refugia in rivers: concepts derived from research on the Miramichi River, New Brunswick (Canada). Ecohydrology. 2014; 8(6):1095–108. Epub October 29, 2014. <u>https://doi.org/10.1002/eco.1566</u>
- Justice C, White SM, McCullough DA, Graves DS, Blanchard MR. Can stream and riparian restoration offset climate change impacts to salmon populations? Journal of Environmental Management. 2017; 188(Supplement C):212–27. <u>https://doi.org/10.1016/j.jenvman.2016.12.005</u>.
- 204. Battin J, Wiley MW, Ruckelshaus MH, Palmer RN, Korb E, Bartz KK, et al. Projected impacts of climate change on salmon habitat restoration. Proc Natl Acad Sci 2007; 104(16):6720–5. <u>https://doi.org/ 10.1073/pnas.0701685104</u> PMID: <u>17412830</u>; PubMed Central PMCID: PMC1871852.
- 205. Honea JM, Jorgensen JC, McClure MM, Cooney TD, Engie K, Holzer DM, et al. Evaluating habitat effects on population status: Influence of habitat restoration on spring-run Chinook salmon. Freshwat Biol. 2009; 54(7):1576–92. <u>https://doi.org/10.1111/j.1365-2427.2009.02208.x</u> ISI:000266637400015.
- 206. Klein S, Herron H, Butcher D. EPA Region 10 climate change and TMDL pilot–South Fork Nooksack River, Washington. Washington, DC: U.S. Environmental Protection Agency, 2017 Contract No.: EPA/600/R-17/281.
- 207. Kemp KB, Blades JJ, Klos PZ, Hall TE, Force JE, Morgan P, et al. Managing for climate change on federal lands of the western United States: perceived usefulness of climate science, effectiveness of adaptation strategies, and barriers to implementation. Ecol Soc. 2015; 20(2).

- 208. Anderson JH, Pess GR, Carmichael RW, Ford MJ, Cooney TD, Baldwin CM, et al. Planning Pacific salmon and steelhead reintroductions aimed at long-term viability and recovery. N Am J Fish Manage. 2014; 34(1):72–93. <u>https://doi.org/10.1080/02755947.2013.847875</u> WOS:000331600200008.
- 209. McClure MM, Carlson SM, Beechie TJ, Pess GR, Jorgensen JC, Sogard SM, et al. Evolutionary consequences of habitat loss for Pacific anadromous salmonids. Evol Appl. 2008; 1(2):300–18. <u>https://doi.org/10.1111/j.1752-4571.2008.00030.x</u> PMID: <u>25567633</u>
- 210. Williams JG. Mitigating the effects of high-head dams on the Columbia River, USA: experience from the trenches. Hydrobiologia. 2008; 609:241–51. <u>https://doi.org/10.1007/s10750-008-9411-3</u> WOS:000256928400020.
- 211. Galbreath PF, Bisbee MA, Dompier DW, Kamphaus CM, Newsome TH. Extirpation and tribal reintroduction of Coho salmon to the interior Columbia River basin Fisheries. 2014; 39(2):77–87. <u>https://doi.org/10.1080/03632415.2013.874526</u>
- 212. Bellmore J, Duda J, Craig L, L. Greene S, E. Torgersen C, J. Collins M, et al. Status and trends of dam removal research in the United States. WIREs Water 2016; 4. https://doi.org/10.1002/wat2.1164
- Noda K, Hamada J, Kimura M, Oki K. Debates over dam removal in Japan. Water and Environment Journal. 2018; 32(3):446–52. <u>https://doi.org/10.1111/wej.12344</u> WOS:000443144600015.
- 214. Hammersley MA, Scott C, Gimblett R. Evolving conceptions of the role of large dams in social-ecological resilience. Ecol Soc. 2018; 23(1). <u>https://doi.org/10.5751/es-09928-230140</u> WOS:000432464800032.
- 215. Quinones RM, Grantham TE, Harvey BN, Kiernan JD, Klasson M, Wintzer AP, et al. Dam removal and anadromous salmonid (*Oncorhynchus* spp.) conservation in California. Rev Fish Biol Fish. 2015; 25 (1):195–215. <u>https://doi.org/10.1007/s11160-014-9359-5</u> WOS:000349767000011.
- 216. Tullos DD, Finn DS, Walter C. Geomorphic and ecological disturbance and recovery from two small dams and their removal. PLoS ONE. 2014; 9(9):e.0108091. <u>https://doi.org/10.1371/journal.pone.</u>0108091 WOS:000342921200081. PMID: 25233231
- 217. Foley MM, Bellmore JR, O'Connor JE, Duda JJ, East AE, Grant GE, et al. Dam removal: Listening in. Wat Res Res, 2017; 53(7):5229–46. <u>https://doi.org/10.1002/2017wr020457</u> WOS:000407895000004.
- 218. Shaffer JA, Juanes F, Quinn TP, Parks D, McBride T, Michel J, et al. Nearshore fish community responses to large scale dam removal: implications for watershed restoration and fish management. Aquatic Sciences. 2017; 79(3):643–60. <u>https://doi.org/10.1007/s00027-017-0526-3</u> WOS:000405798900017.
- Warrick JA, Bountry JA, East AE, Magirl CS, Randle TJ, Gelfenbaum G, et al. Large-scale dam removal on the Elwha River, Washington, USA: Source-to-sink sediment budget and synthesis. Geomorphology. 2015; 246:729–50. <u>https://doi.org/10.1016/j.geomorph.2015.01.010</u> WOS:000360869400055.
- 220. Foley MM, Warrick JA, Ritchie A, Stevens AW, Shafroth PB, Duda JJ, et al. Coastal habitat and biological community response to dam removal on the Elwha River. Ecol Monogr. 2017; 87(4):552–77. <u>https://doi.org/10.1002/ecm.1268</u> WOS:000414241400002.
- 221. Quinn TP, Shaffer JA, Brown J, Harris N, Byrnes C, Crain P. Juvenile Chinook salmon, *Oncorhynchus tshawytscha*, use of the Elwha river estuary prior to dam removal. Environmental Biology of Fishes. 2014; 97(6):731–40. <u>https://doi.org/10.1007/s10641-013-0173-z</u> WOS:000334913000009.
- 222. Yoshiyama RM, Gerstung ER, Fisher FW, Moyle PB. Historical and present distribution of Chinook salmon in the Central Valley. In: Brown R, editor. Contributions to the biology of Central Valley salmonids: CDFG Fish Bulletin 179; 2001. p. 71–176.
- 223. International ICF. Battle Creek winter-run Chinook salmon reintroduction plan. Sacramento, CA: Prepared for California Department of Fish and Wildlife, 2016 ICF00148.15.
- 224. NMFS, National Marine Fisheries Service. Recovery plan for the evolutionarily significant units of Sacramento River winter-run Chinook Salmon and Central Valley spring-run Chinook salmon and the distinct population segment of California Central Valley Steelhead. National Marine Fisheries Service, California Central Valley Area Office: 2014.
- 225. NMFS, National Marine Fisheries Service. Endangered and threatened species: Designation of a nonessential experimental population for middle Columbia River steelhead above the Pelton Round Butte Hydroelectric Project in the Deschutes River Basin, OR Federal Register. 2013; 78(10):2893–907.
- 226. NMFS, National Marine Fisheries Service. Steelhead population to be reintroduced to Oregon's Deschutes River Basin. National Marine Fisheries Service, Northwest Region, 2018.
- 227. Campbell NR, Kamphaus C, Murdoch K, Narum SR. Patterns of genomic variation in coho salmon following reintroduction to the interior Columbia River. Ecology and Evolution. 2017; 7(23):10350–60. https://doi.org/10.1002/ece3.3492 PMC5723619. PMID: 29238560

- 228. Giorgi C, Baldwin CM. Phase 1 Habitat Assessments. Lake Roosevelt Forum, 25 April 2018, Spokane, Washington. Availble at <u>http://lrf.org/conference-presentations/2018/4.25.18–9.40cGiogi-BaldwinPhase1HabitatAssessments-LRF2018v4.pdf</u>: Spokane Tribe and Colville Confederated Tribes, 2018.
- 229. Naish K, Taylor J, Levin P, Quinn T, Winton J, Huppert D, et al. An evaluation of the effects of conservation and fishery enhancement hatcheries on wild populations of salmon. Advances in Marine Biology 2008; 53:61–193.
- Christie MR, Ford MJ, Blouin MS. On the reproductive success of early-generation hatchery fish in the wild. Evol Appl. 2014; 7(8):883–96. <u>https://doi.org/10.1111/eva.12183</u> PMID: <u>25469167</u>; PubMed Central PMCID: PMC4211718.
- 231. Fagan WF, Holmes EE. Quantifying the extinction vortex. 2006; 9(1):51–60. <u>https://doi.org/10.1111/j.</u> 1461-0248.2005.00845.x PMID: 16958868
- **232.** Hatchery Scientific Review Group. Report to Congress on Columbia River Basin Hatchery Reform. 2009.
- 233. Chittenden CM, Biagi CA, Davidsen JG, Davidsen AG, Kondo H, McKnight A, et al. Genetic versus rearing-environment effects on phenotype: hatchery and natural rearing effects on hatchery- and wild-born coho salmon. PLOS ONE. 2010; 5(8):e12261. <u>https://doi.org/10.1371/journal.pone.0012261</u> PMID: 20808853
- 234. Gale MK, Hinch SG, Donaldson MR. The role of temperature in the capture and release of fish. Fish and Fisheries. 2013; 14(1):1–33. <u>https://doi.org/10.1111/j.1467-2979.2011.00441.x</u> WOS:000314804600001.
- 235. Tillotson MD, Quinn TP. Selection on the timing of migration and breeding: A neglected aspect of fishing-induced evolution and trait change. Fish and Fisheries. 2018; 19(1):170–81. <u>https://doi.org/10. 1111/faf.12248</u> WOS:000418927100010.
- 236. Hard JJ, Gross MR, Heino M, Hilborn R, Kope RG, Law R, et al. Evolutionary consequences of fishing and their implications for salmon. Evol Appl. 2008; 1(2):388–408. <u>https://doi.org/10.1111/j.1752-4571.</u> 2008.00020.x WOS:000262783500015. PMID: 25567639
- 237. Law R. Fishing, selection, and phenotypic evolution. ICES J Mar Sci. 2000; 57(3):659–68. <u>https://doi.org/10.1006/jmsc.2000.0731</u>%J ICES Journal of Marine Science.
- 238. Waples RS, Zabel RW, Scheuerell MD, Sanderson BL. Evolutionary responses by native species to major anthropogenic changes to their ecosystems: Pacific salmon in the Columbia River hydropower system. Molecular Ecology. 2008; 17(1):84–96. <u>https://doi.org/10.1111/j.1365-294X.2007.03510.x</u> PMID: <u>18268786</u>
- 239. Waples RS, Audzijonyte A. Fishery-induced evolution provides insights into adaptive responses of marine species to climate change. Front Ecol Environ. 2016; 14(4):217–24. <u>https://doi.org/10.1002/ fee.1264</u>
- 240. Rosenberger AE, Dunham JB, Neuswanger JR, Railsback SF. Legacy effects of wildfire on stream thermal regimes and rainbow trout ecology: an integrated analysis of observation and individual-based models. Freshwater Science. 2015; 34(4):1571–84. <u>https://doi.org/10.1086/683338</u>
- 241. Swain DP, Sinclair AF, Hanson JM. Evolutionary response to size-selective mortality in an exploited fish population. Proceedings of the Royal Society B: Biological Sciences. 2007; 274(1613):1015–22. <u>https://doi.org/10.1098/rspb.2006.0275</u> PMID: <u>17264058</u>
- 242. Piou C, Taylor MH, Papaix J, Prevost E. Modelling the interactive effects of selective fishing and environmental change on Atlantic salmon demogenetics. J Appl Ecol. 2015; 52(6):1629–37. <u>https://doi.org/10.1111/1365-2664.12512</u> WOS:000367095400024.
- Kendall NW, Quinn TP. Quantifying and comparing size selectivity among Alaskan sockeye salmon fisheries. Ecol Appl. 2012; 22(3):804–16. WOS:000303312000006. PMID: 22645812
- 244. Crichigno SA, Becker LA, Orellana M, Larraza R, Mirenna G, Battini MA, et al. Rainbow trout adaptation to a warmer Patagonia and its potential to increase temperature tolerance in cultured stocks. Aquaculture Reports. 2018; 9:82–8. <u>https://doi.org/10.1016/j.aqrep.2017.11.001</u> WOS:000452094700012.
- 245. Hamilton JA, Miller JM. Adaptive introgression as a resource for management and genetic conservation in a changing climate. Conserv Biol. 2016; 30(1):33–41. <u>https://doi.org/10.1111/cobi.12574</u> WOS:000368938000005. PMID: <u>26096581</u>
- 246. Aitken SN, Whitlock MC. Assisted gene flow to facilitate local adaptation to climate change. 2013; 44 (1):367–88. <a href="https://doi.org/10.1146/annurev-ecolsys-110512-135747">https://doi.org/10.1146/annurev-ecolsys-110512-135747</a>
- 247. Esvelt KM, Smidler AL, Catteruccia F, Church GM. Concerning RNA-guided gene drives for the alteration of wild populations. Elife. 2014; 3. <u>https://doi.org/10.7554/eLife.03401</u> WOS:000209690800001. PMID: <u>25035423</u>

- 248. Webber BL, Raghu S, Edwards OR. Opinion: Is CRISPR-based gene drive a biocontrol silver bullet or global conservation threat? 2015; 112(34):10565–7. <u>https://doi.org/10.1073/pnas.1514258112</u> %J Proceedings of the National Academy of Sciences. PMID: <u>26272924</u>
- 249. Allendorf FW, Hohenlohe PA, Luikart G. Genomics and the future of conservation genetics. Nature Reviews Genetics. 2010; 11(10):697–709. <u>https://doi.org/10.1038/nrg2844</u> WOS:000281911300010. PMID: <u>20847747</u>
- **250.** Dana GV, Cooper AM, Pennington KM, Sharpe LM. Methodologies and special considerations for environmental risk analysis of genetically modified aquatic biocontrol organisms. Biological Invasions. 2014; 16(6):1257–72. <u>https://doi.org/10.1007/s10530-012-0391-x</u> WOS:000338444500006.
- 251. Fitzpatrick BM, Johnson JR, Kump DK, Smith JJ, Voss SR, Shaffer HB. Rapid spread of invasive genes into a threatened native species. Proceedings of the National Academy of Sciences of the United States of America. 2010; 107(8):3606–10. <u>https://doi.org/10.1073/pnas.0911802107</u> WOS:000275130900060. PMID: <u>20133596</u>
- 252. Hayes KR, Hosack GR, Dana GV, Foster SD, Ford JH, Thresher R, et al. Identifying and detecting potentially adverse ecological outcomes associated with the release of gene-drive modified organisms. Journal of Responsible Innovation. 2018; 5:S139–S58. <u>https://doi.org/10.1080/23299460.2017.1415585</u> WOS:000434459000009.
- 253. Lu XM, Siemann E, He MY, Wei H, Shao X, Ding JQ. Climate warming increases biological control agent impact on a non-target species. Ecol Lett. 2015; 18(1):48–56. <u>https://doi.org/10.1111/ele.12391</u> WOS:000346464400005. PMID: <u>25376303</u>
- 254. McLachlan JS, Hellmann JJ, Schwartz MW. A framework for debate of assisted migration in an era of climate change. 2007; 21(2):297–302. <u>https://doi.org/10.1111/j.1523-1739.2007.00676.x</u> PMID: <u>17391179</u>
- 255. Supple MA, Shapiro B. Conservation of biodiversity in the genomics era. Genome Biology. 2018; 19. https://doi.org/10.1186/s13059-018-1520-3 WOS:000444255800002. PMID: <u>30205843</u>
- 256. Flagg TA, Waknitz FW, Maynard DJ, Milner GB, Mahnken CVW. The effects of hatcheries on native coho salmon populations in the lower Columbia River. In: Schramm J H. L., Piper RG, editors. Uses and effects of cultured fishes in aquatic ecosystems. Symposium 15. Bethesda, Maryland.: American Fisheries Society; 1995. p. 366–75.
- 257. Quinn TP, Peterson JA, Gallucci VF, Hershberger WK, Brannon EL. Artificial selection and environmental change: Countervailing factors affecting the timing of spawning by coho and Chinook salmon. Trans Am Fish Soc. 2002; 131(4):591–8.
- 258. Ford JKB, Ellis GM. Selective foraging by fish-eating killer whales Orcinus orca in British Columbia. Marine Ecology-Progress Series. 2006; 316:185–99. ISI:000239512600016.
- 259. Staudinger MD, Grimm NB, Staudt A, Carter SL, III FSC, Kareiva P, et al. Impacts of climate change on biodiversity, ecosystems, and ecosystem services: Technical input to the 2013 National Climate Assessment. Cooperative Report to the 2013 National Climate Assessment, 2012.
- 260. Healey M. The cumulative impacts of climate change on Fraser River sockeye salmon (*Oncorhynchus nerka*) and implications for management. Can J Fish Aquat Sci. 2011; 68(4):718–37. <u>https://doi.org/10.1139/f11-010</u> WOS:000291183500012.
- 261. Martins EG, Hinch SG, Patterson DA, Hague MJ, Cooke SJ, Miller KM, et al. Effects of river temperature and climate warming on stock-specific survival of adult migrating Fraser River sockeye salmon (*Oncorhynchus nerka*). Global Change Biol. 2011; 17(1):99–114. <u>https://doi.org/10.1111/j.1365-2486.2010.02241.x</u> WOS:000284851500009.