Seasonality and response of ocean acidification and hypoxia to major environmental anomalies in the southern Salish Sea, North America (2014–2018)

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Abstract. Coastal and estuarine ecosystems fringing the North Pacific Ocean are particularly vulnerable to ocean acidification, hypoxia, and intense marine heatwaves as a result of interactions among natural and anthropogenic processes. Here we characterize variability during a seasonally resolved cruise time series (2014–2018) in the southern Salish Sea (Puget Sound, Strait of Juan de Fuca) and nearby coastal waters for select physical (temperature, T; salinity, S) and biogeochemical (oxygen, O2; carbon dioxide fugacity, fCO2; aragonite saturation state, Ω_{arag}) parameters. Medians for some parameters peaked (T, Ω_{arag}) in surface waters in summer, while others (S, O2, fCO2) changed progressively across spring–fall, and all parameters changed monotonically or were relatively stable at depth. Ranges varied considerably for all parameters across basins within the study region, with stratified basins consistently the most variable. Strong environmental anomalies occurred during the time series, allowing us to also qualitatively assess how these anomalies affected seasonal patterns and interannual variability. The peak temperature anomaly associated with the 2013–2016 northeast Pacific marine heatwave–El Niño event was observed in boundary waters during the October 2014 cruise, but Puget Sound cruises revealed the largest temperature increases during 2015–2016 timeframe. The most extreme hypoxia and acidification measurements to date were recorded in Hood Canal (which consistently has the most extreme conditions) during the same period; however, they were shifted earlier in the year relative to previous events. During autumn 2017, after the heat anomaly, a distinct carbonate system anomaly with unprecedentedly

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low Ω_{arrag} and high fCO_2 occurred in parts of the southern Salish Sea that are not normally so acidified. This novel "CO₂ storm" appears to have been driven by anomalous<u>ly high</u> river discharge earlier in 2017, which resulted in enhanced stratification and inferred primary productivity anomalies, indicated by persistently and anomalously high O_2 , low fCO_2 , and high chlorophyll. Unusually, this CO_2 anomaly was decoupled from O_2 dynamics compared to past Salish Sea hypoxia and acidification events. The complex interplay of weather, hydrological, and circulation anomalies revealed distinct multiple stressor scenarios that will potentially affect regional ecosystems under a changing climate. Further, the frequencies at which Salish cruise observations crossed known or preliminary species sensitivity thresholds illustrates the relative risk landscape of temperature, hypoxia, and acidification anomalies in the southern Salish Sea in the present-day, with implications for how multiple stressors may combine to present potential migration, survival, or physiological challenges to key regional species in the future. The Salish cruise data product used in this publication is available at https://doi.org/10.25921/zgk5-ep63 (Alin et al., 2022), with an additional data product including all calculated CO_2 system parameters available at https://doi.org/10.25921/zgk5-ep63 (Alin et al., 2023b).

1 Introduction

Northeast Pacific Ocean ecosystems are particularly vulnerable to marine heatwaves, hypoxia, and ocean acidification—the increase in seawater carbon dioxide (CO2) due to ocean uptake of anthropogenic CO2 emissions, which drives declining pH and calcium carbonate saturation states (Ω)—as a result of interactions among natural and anthropogenic processes. Located at the terminus of global oceanic thermohaline circulation, subsurface NE Pacific water masses have low oxygen (O2) and high dissolved inorganic carbon (DIC) content resulting from respiratory processes during isolation from the atmosphere (e.g., Franco et al., 2021 and references therein). Naturally high NE Pacific CO2 levels are enhanced further through the addition of anthropogenic CO₂ (Feely et al., 2004, 2016; Sabine et al., 2004). Eastern boundary current systems accentuate this vulnerability by bringing subsurface, naturally O₂-poor, CO₂-rich waters toward the surface through upwelling (Feely et al., 2008; Chavez and Messié, 2009; Chavez et al., 2017). Estuarine systems such as the Salish Sea are typically lower in buffering capacity and are already rich in CO2 due to dynamic local biological, hydrological, and geochemical processes; this natural estuarine acidification is amplified when oceanic waters acidified by the uptake of anthropogenic CO2 are transported into the estuary via estuarine circulation (Feely et al., 2010, 2018; Wallace et al., 2014; Pacella et al., 2018; Cai et al., 2021; Hunt et al., 2022). Thus, estuaries connected to upwelling coastal systems, particularly in the NE Pacific, receive naturally acidified, low-oxygen marine waters relative to those in other coastal regions (e.g., Windham-Myers et al., 2018). Continually rising CO₂ emissions and other climate change effects on coastal and estuarine processes are expected to increase the spatial and temporal prevalence of acidified estuarine conditions (Pacella et al., 2018; Evans et al., 2019; Jarníková et al., 2022). Further, fjord-like estuaries with entrance sills, like Puget Sound and Hood Canal, retain some of the outgoing waters via mixing over the sills (known as reflux, e.g., MacCready et al., 2021), so anomalies tend to persist longer in these basins (Jackson et al., 2018).

Since 2007, carbonate system observations throughout the water column in coastal and estuarine NE Pacific ecosystems have proliferated, providing insight into the dynamics of ocean acidification parameters, including both measured (dissolved inorganic carbon, DIC; total alkalinity, TA; and sometimes pH on the total scale, pH_T) and calculated (pH_T; CO₂ partial pressure or fugacity, pCO_2 or fCO_2 ; and calcium carbonate saturation states, aragonite: Ω_{arag} , calcite: Ω_{calc}) variables (e.g., Feely et al., 2008, 2010; Alin et al., 2022, 2023a). Hood Canal, having long been known as a hotspot for hypoxia (defined here as oxygen levels below 62 μ mol kg⁻¹ = 2.0 mg L⁻¹ = 1.5 mL L⁻¹) (Newton et al., 2007), was shown to have the most severe argonite undersaturation (Ω_{arae} <1) in the southern Salish Sea during the first direct carbonate system measurements (Feely et al., 2010). Subsequent observations showed aragonite undersaturation to be prevalent throughout most of the water column, most of the time in the northern Salish Sea as well (Ianson et al., 2016; Evans et al., 2019), with numerical models showing that pre-industrial Salish Sea chemistry predisposed it to rapid expansion of undersaturated conditions (Bednaršek et al., 2020a; Jarníková et al., 2022). Surface climatologies of carbonate chemistry in marine surface waters throughout Washington state revealed strong seasonal variability, with particularly high fCO_2 and low pH and Ω values in Puget Sound surface waters during fall and winter months (November-March: Fassbender et al., 2018). Seasonal variability of pCO₂, pH. and Ω_{arag} observed in high-resolution moored surface time-series is among the highest in the world (Sutton et al., 2016), so these waters have a long "time of emergence," i.e., the projected future time when a statistically significant anthropogenic trend in CO₂ content can be detected to emerge from the bounds of natural variability at a location (Sutton et al., 2016, 2019). Moreover, biological modulation of carbonate chemistry or temperature seasonality can obscure or decouple changes in pH

these waters have a long "time of emergence," i.e., the projected future time, when a statistically significant anthropogenic trend, in CO₂ content can be detected to emerge from the bounds of natural variability at a location (Sutton et al., 2016, 2019). Moreover, biological modulation of carbonate chemistry or temperature seasonality can obscure or decouple changes in pH and fCO₂ from those seen in saturation states (Kwiatkowski and Orr, 2018; Lowe et al., 2019; Cai et al., 2020). Estimates of anthropogenic CO₂ content from Salish Sea observations and models point to widespread Ω_{arng} undersaturation having emerged here and in other regional waters since pre-industrial times (Feely et al., 2010; Pacella et al., 2018; Evans et al., 2019; Hare et al., 2020; Jarníková et al., 2022). These factors, in tandem with strong benthic-pelagic coupling of biogeochemical cycles (e.g., high surface productivity contributing to deep respiration hotspots, Hickey and Banas, 2008; Siedlecki et al., 2015), highlight the need for detailed biogeochemical observations throughout the water column in this biologically productive region to understand the atmospheric, terrestrial, and marine processes driving dynamic biogeochemical conditions in the Salish Sea.

5 Here we use the Salish cruise data product (2008–2018; Alin et al., 2022, 2023a, b) to characterize seasonal variability and major anomalies in physical and biogeochemical conditions in Puget Sound and its boundary waters (Strait of Juan de Fuca, coastal waters) during the seasonally resolved part of the time-series (2014–2018). All calculated marine inorganic carbon parameters used in this analysis were calculated from measured dissolved inorganic carbon, total alkalinity, and ancillary hydrographic observations (temperature, salinity, and phosphate and silicate content) described by Alin et al. (2023a). We used temperature, salinity, oxygen (O2), fugacity of carbon dioxide (fCO2), and aragonite saturation state (Ω_{arag}) median conditions and variation to characterize seasonal ocean acidification, hypoxia, and warming conditions across Puget Sound basins and its boundary waters. Major anomalies in large-scale marine and atmospheric temperature, as well as regional precipitation and

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river runoff, occurred during 2013–2018, and we qualitatively relate the timing and magnitude of observed biogeochemical anomalies in the study region to anomalies in regional weather and physical oceanography sometimes driven by these major large-scale anomalies. Cruises prior to the onset of the 2013–2018 anomalies and existing regional climatologies provided the long-term context for the apparent magnitude and duration of physical and biogeochemical anomalies observed during the seasonal sampling period. Finally, we evaluated how the physical and biogeochemical Salish cruise time-series through 2018 reveals the changing landscape of multiple interacting ocean stressors as they are relevant to key ecologically and economically important fish and invertebrate species in this oceanographically dynamic region.

2 Environmental context for the Salish Sea cruise time-series

2.1 Geographic setting

Puget Sound (PS) is the southernmost glacial fjord estuarine system on the North American Pacific Coast and comprises the southern part of the Salish Sea, which also encompasses the Strait of Juan de Fuca (SJdF) and the Strait of Georgia (Figure 1).
 Oceanographic conditions within Puget Sound are determined by a combination of river inputs, marine source waters, vigorous tidal mixing, bathymetric complexity, and local to large-scale climatic influences (Moore et al., 2008; Feely et al., 2010; Banas et al., 2015; MacCready et al., 2021). Puget Sound is comprised of four basins: Main Basin (MB), South Sound (SS), Whidbey
 Basin (WB), and Hood Canal (HC). PS receives direct freshwater input from 14 major and many smaller rivers draining into PS, indirect freshwater input from the Fraser River, which drains into the Strait of Georgia, and carbon and nutrient inputs from urban and agricultural environments surrounding the Salish Sea ecosystem (Mohamedali et al., 2011; Banas et al., 2015).

The northern California Current Ecosystem (CCE) provides the marine source water for deep waters within the southern Salish

Sea and experiences episodic upwelling during spring—early fall (April—September) as a result of northwesterly equatorward winds causing offshore Ekman pumping (Huyer, 1983). Downwelling conditions occur during late fall—early spring (October—March) due to seasonal wind reversal to poleward-dominant winds along the coast. Upwelling conditions bring deep, nutrient-and CO2-rich, and O2-depleted marine water masses into the Strait of Juan de Fuca from the Juan de Fuca Canyon (Figure 1).

This water transits at depth to the glacial sill complex at Admiralty Reach (AR), where it enters PS at depth during episodic marine intrusions. Strong freshwater outflow through SJdF, particularly during summer months when peak Fraser River discharge occurs, contributes to and enhances this estuarine circulation (Davis et al., 2014; Giddings et al., 2014). Strong tides and glacial sills within AR at the entrance to Puget Sound impart strong mixing—of outgoing warmer, fresher surface estuarine waters with colder, saltier marine waters entering PS from SJdF at depth; the strength of tides and resulting mixing influence, the amount and characteristics of incoming marine water that refreshes deep water masses in all PS basins, and some of the outgoing estuarine water is refluxed back into Puget Sound (MacCready et al., 2021; MacCready and Geyer, 2023).

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Glacial sills restrict estuarine circulation throughout the Salish Sea and among the Puget Sound basins as well. limiting marine intrusions and deep-water renewal to episodic occurrences and resulting in long residence and flushing times in some parts of this inland sea, including Hood Canal (Babson et al., 2006; Pawlowicz et al., 2007; MacCready et al., 2021). The Main Basin is the widest, deepest, and most deeply wind-mixed of the Puget Sound basins. Deep waters enter South Sound, the shallowest basin, from MB when they pass over another glacial sill at the Tacoma Narrows and undergo strong tidal mixing again while flowing into South Sound. Thus, the deeply mixed MB and SS share a deep-water transit path. In contrast, Whidbey Basin and Hood Canal have narrower basins than MB and major river inputs emanating from the terminus of each basin, resulting in strong stratification and gradients of physical and biogeochemical conditions between surface and bottom waters. HC is also bounded by a glacial sill and has a long history of study of deep-water oxygen concentrations, as hypoxia and fish kills have been observed there (Newton et al., 2011, 2012 and references therein). Observations and models for the Strait of Georgia 160 suggest that mixing associated with glacial bathymetric features to the north of Puget Sound may afford some protection to deep northern Salish Sea basins, due to more rapid O2 uptake than CO2 outgassing (Johannessen et al., 2014; Ianson et al., 2016); this mechanism does not appear to protect Hood Canal from developing hypoxia. While not bounded by a glacial sill, circulation in WB is severely restricted at its northern outlet, and it receives strong river input in two locations. While WB has side inlets with hypoxia, the mainstem of the basin tends to see only moderately low oxygen values but no hypoxia.

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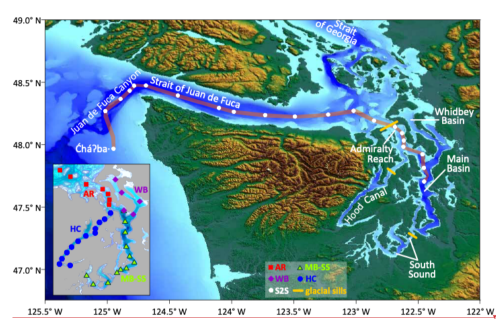
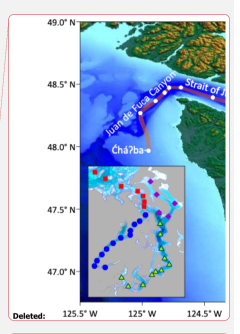


Figure 1: Map of the southern Salish Sea and its boundary waters with all study basins named. The subset of sampling stations tracing a path between the "Chá?ba"—meaning "whale tail" in the language of the Quileute Tribe—mooring on the Washington state (USA) continental shelf to the Main Basin of Puget Sound constitute the Sound-Sea (S2S) transects. Inset map shows station groupings used for analyses of Puget Sound cruises: Admiralty Reach (AR), Main Basin—South Sound (MB—SS), Whidbey Basin (WB), and Hood Canal (HC). Locations of glacial sills that restrict deep-water exchange in Puget Sound are shown. The Fraser River, mentioned in the text, enters the Strait of Georgia from the east, to the north of the map area. We extracted topographic and bathymetric data from the NOAA National Centers for Environmental Grid Extract Coastal Relief Model (3-second resolution, https://www.ncei.noaa.gov/maps/grid-extract/, accessed Nov. 13, 2014), and gridded the data in Surfer using a minimum curve gridding technique.

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Both regional weather and large-scale climate factors play important roles in driving physical, chemical, and biological processes in the Salish Sea and its boundary waters. From 2013 to 2016, an unprecedented marine heatwave (MHW) developed and persisted in the NE Pacific Ocean, followed by a very strong El Niño event in the equatorial Pacific Ocean during 2015–2016, both of which strongly influenced regional weather, oceanography, and ecosystems (e.g., Bond et al., 2015; Jacox et al., 2016; McClatchie et al., 2016; Morgan et al., 2019; N. Bond in Sobocinski, 2021). The NE Pacific heatwave's direct influence on Washington's coastal waters and the Salish Sea ecosystem began when anomalously warm waters from the North Pacific were advected onto the Pacific Northwest coast in mid-September 2014 (Peterson et al., 2017). However, associated strong, large-scale air temperature anomalies strongly influenced the surface Puget Sound system and preceded the arrival of the warmed ocean water masses (Swain et al., 2016), with anomalously warm, dry summer conditions starting in 2013 over the



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southern Salish Sea (Table 1, Figure 2). As a result of these large-scale heat anomalies, Puget Sound and Washington coastal waters also experienced strong precipitation, river discharge, and solar energy flux anomalies during 2013–2018 (Table 1 and references therein). Upwelling anomalies reflect basin-scale climate drivers and influence both the upwelling strength and the depth of marine source waters for deep waters of the southern Salish Sea (e.g., Jacox et al., 2015).

Table 1. Major environmental anomalies occurring during 2013–2018 and regional environmental drivers affecting the southern Salish Sea.

	circulation driver anomalies and effects											
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Year	Air temperature (°C) ^a	Solar energy flux ^b	Precipitation ^a	River discharge (Q) ^c	$Upwelling^d$	Puget Sound circulation and stratification ^e	Sources ^f					
2013	+0.5-0.9 May-Sep.	Higher than normal May– Aug.	Apr. and Sep. very wet, Oct.–Dec. drier than normal	FR: high early peak Q in May, low fall Q; PS: higher Q in Mar.–Jun. and early Oct.	Below normal Aug.– Sep.	Stratification started earlier; normal marine intrusion timing (fall)	Bumbaco ^a ; Albertson et al. ^{b,d} ; Dzinbal ^c ; Ruef et al. ^e					
2014	0.9+ (5 th warmest year on record)	Near theoretical maximum May-Sep.	119% (wettest Mar.)	FR: high early peak Q in May, low summer Q, high late fall Qs; PS: higher than normal in spring and fall, average in summer	Early fall transition to downwelling (stronger than normal in SepOct.)	Deep mixing driven by cold, dry, windy conditions led to persistent (Feb.—Oct,) high O ₂ , low T conditions in deep HC; stronger than normal stratification in parts of MB	Bumbaco ^a ; Albertson et al. ^{b,d} ; Dzinbal ^c ; Mickett et al. ^e ; Stark ^e					
2015	1.4+ (warmest year on record)	Near theoretical maximum May-Sep.; gloomy fall	107%, snowpack deficit, summer drought (3 rd wettest Dec.)	FR: record high spring Qs, high/early peak, record low summer Qs; PS: extremely high Jan.— Mar., Nov.—Dec.; extremely low May— Sep./Oct.	Stronger May–Jun. upwelling	Strong early stratification in MB, reduced by summer drought; HC deep-water renewal six weeks early (deep water >2.5 °C above climatology)	Bumbaco ^a ; Albertson et al. ^{b,d} ; Dzinbal ^c ; Bos et al. ^e ; Ruef et al. ^e					
2016	1.0+ (3 rd warmest year on record)	Below average winter and fall; above average spring— summer	113% (wet Feb.–May, normal summer, wettest Oct.)	FR: early low peak (4–6 weeks early), very low summer, very high Nov.; PS: very high Qs mid-Jan.—Mar., Oct.—Nov., low May—Sep. Qs	Stronger downwelling JanMar., Oct-Nov.; stronger May upwelling	Stronger stratification than normal in spring and fall; longer residence time in MB during summer drought; annual flushing of deep HC water four weeks early	Bond and Bumbaco ^a ; Albertson et al. ^{b,d} ; Burks ^c ; Bos et al. ^c ; Ruef et al. ^c ; Albertson et al. ^e					
2017	Normal (warmest August)	Below average FebMay, Nov.; above average Jun Oct. (JulSep. periods of haze to wildfire smoke)	112% (wettest FebApr., driest Jul Sep., wet fall, normal snowpack)	FR: early higher peak Q, lower Q Jul.—Oct., high Nov.—Dec.; PS: high peaks in Jan.—Mar., low end of normal Qs through summer, event peaks in Oct., Nov.	Stronger downwelling in Apr., Nov.	Stronger, more persistent stratification than normal due to high spring Q anomaly (resulting in sustained MB phytoplankton blooms, persistent low water column S in MB); normal HC deep flushing, historic salinity minima (2 SD below normal before summer)	Bumbaco and Bond ^a ; Albertson et al. ^{b.d.c} ; Burks ^c ; Bos et al. ^c ; Ruef et al. ^c					
2018	0.6+	Above average except Jan., Feb., Apr. (AugSep.	98% (2 nd wettest Apr., driest May– Aug.)	FR: runoff peak one month early, highest peak of these years, low Qs JunNov.; PS: very	Stronger downwelling in Jan.; upwelling in	Strong springtime density stratification, but decreased in summer (delayed bloom); more favourable	Bumbaco and Bond ^a ; Albertson et al. ^{b,d,e} ;					

reduced due to wildfire smoke)

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high Q peaks in winterspring (early) and fall, low-to-very-low summer Feb. (spring period transition two months early) since for de

periods for intrusions across AR than any year since 2013; normal timing for deep HC renewal (by end Sep.)

^a Annual and monthly average air temperature and precipitation anomalies are relative to 1981–2010 normal. Record monthly, seasonal, or annual anomalies up to 2018 noted in parentheses. ^b Daily solar energy flux, an indicator of sunniness, measured in Seattle and compared to the highest theoretical solar energy for the latitude and time of year and fully overcast conditions. ^c River flows for the Fraser River (FR)—with a single summer discharge (Q, m³ s³) peak—are described relative to the median discharge from 1912 to the year reported. Puget Sound rivers (PS)—with an early summer snowmelt discharge peak, and a late fall rainfall and winter storm discharge peak—are reported relative to the full length of their U.S. Geological Survey observational records, ranging from 47 to 104 years. ^d Upwelling anomalies are reported as monthly average upwelling index values (m³ s⁻¹ per 100 m of coastline) that fall outside the interquartile range (25–75%) for 48° N, 125° W by NOAA's Pacific Fisheries Environmental Laboratory, Baseline period is 1967 to the year of each annual report. ^c Anomalies in Puget Sound stratification or deep-water renewal events were reported in temperature and salinity water quality narratives of annual marine conditions reports. ¹ Sources for observations in previous columns are listed by authors of the relevant sections in the PSEMP Marine Waters Workgroup annual overview of marine conditions published in the following year (i.e., listed as citations in the relevant year's report: PSEMP Marine Waters Workgroup, 2014, 2015, 2016, 2017, 2018, 2019).

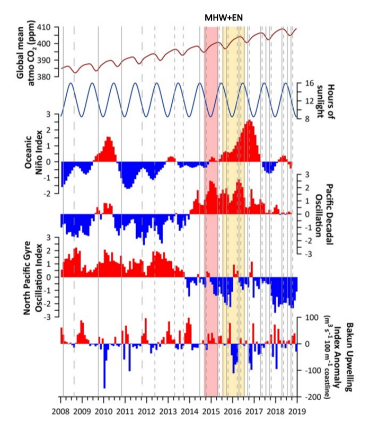


Figure 2: Monthly time-series for 2008–2018 for the Oceanic Niño Index anomaly (NOAA Climate Prediction Center, 2019), Pacific Decadal Oscillation (Mantua, N., 2019), North Pacific Gyre Oscillation (Di Lorenzo, 2019), and Bakun Upwelling Index Anomaly for 48° N (NOAA Pacific Fisheries Environmental Laboratory, 2019). Positive anomalies for all climate indices are shown in red, negative in blue. Superimposed on this are the durations of the maximum intensity of the northeast Pacific marine heatwave (MHW), shaded in red during its peak manifestation in Washington marine waters (September 2014–April 2015) and in yellow for its later moderate intensity window, which overlapped with the 2015–2016 El Niño event (EN, July 2015–July 2016). Duration and intensity ranges were inferred from Gentemann et al. (2017) and the OSU MODIS water temperature anomaly climatology tool (NANOOS, 2019). Overlap between the 2014–2016 NE Pacific heatwave and warm waters that may be the result of the El Niño can be seen by comparing the Oceanic Niño Index positive anomalies to the yellow shading, based on satellite sea surface temperature analysis associated with the marine heatwave (i.e., Gentemann et al., 2017). Seasonality is shown using hours of sunlight per day as a proxy, shown at top (timeanddate.com, 2019). Finally, the timing of all Salish cruises is indicated by vertical lines, with Sound-to-Sea (S2S) cruises in short-dashed lines, Puget Sound (PS) cruises in solid lines, and the two cruises that encompassed both sets of stations in long-dashed lines.

3 Methods: Observations, calculations, and data visualization

3.1 Salish cruise time-series

For this analysis of seasonal variability and oceanographic anomalies, we used the Salish cruise data product, comprising 35 consistently formatted and quality controlled cruises data sets collected throughout the study region from 2008 to 2018 (Alin et al., 2022, 2023a). The Salish cruise data product includes CTD and discrete measurements collected on each cruise, including temperature, salinity, and oxygen measurements collected by sensors during CTD casts, and discrete water samples measured for oxygen, nutrients (phosphate, silicate, nitrate, nitrite, and ammonium), dissolved inorganic carbon (DIC), and total alkalinity (TA) content. Measurement uncertainties were ±0.01 °C_{*}±0.02, and ±2% of saturation for temperature, salinity, and oxygen sensor observations, respectively. For laboratory analyses on discrete water samples, uncertainties were ±1% for

and oxygen sensor observations, respectively. For laboratory analyses on discrete water samples, uncertainties were ±1% for oxygen (precision), ±2% for nutrients, and ±0.1% (~2 µmol kg_s⁻¹) for DIC and TA content. Data collection methods are explained in detail in the companion paper (Alin et al., 2023a), with quality control following methods developed for identifying outliers in the Coastal Data Analysis Product in North America (CODAP-NA, Jiang et al., 2021).

260 The timing of both Puget Sound (PS) cruises and Sound-to-Sea (S2S) cruises has been particularly consistent and frequent since 2014, with 24 of the 35 cruises having taken place between July 2014 and October 2018. Thus, our analysis of seasonal patterns in ocean conditions spanning the upwelling season focuses on the period from 2014 to 2018 (Figure 2), and will provide the oceanographic context for numerous biological oceanography studies that have also been conducted on the S2S and PS cruises. While each cruise represents a snapshot of conditions along a transect from the coast into the southern Salish

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Sea or across the basins within the southern Salish Sea, collectively this cruise time-series illuminates typical spatial patterns throughout the southern Salish Sea, throughout the water column, and through the seasonal cycle.

3.2 Calculated parameters and uncertainty

DIC and TA measurements from the Salish cruise data product were used to calculate the full suite of inorganic carbon parameters, although our discussion of seasonal ocean acidification conditions and anomalies focuses on the calculated parameters fCO_2 and Ω_{amg} , as these are two of the inorganic carbon parameters most familiar to our science and resource management end users. Of the carbonate system parameters, fCO_2 is most directly relatable to atmospheric values and trends and thus has intuitive value, and Ω_{amg} is critical to many of the important calcifying species in the Salish Sea. For species and investigators for whom pH_T and Ω_{calc} are more relevant (e.g., Dungeness crab), complementary figures are provided in the Supplemental Material. A regression between Ω_{arng} and Ω_{calc} had a coefficient of determination (R^2) >0.99, with Ω_{calc} being 1.59 times Ω_{arag} in the Salish cruise data package, which is a slightly larger coefficient than reported for the relationship between the two parameters by Mucci (1983), presumably to different temperature and salinity characteristics of our regional water masses.

We used the R package seacarb function carb to calculate all carbonate system parameters (Gattuso et al., 2023). Input parameters from the Salish cruise compiled data set comprised DIC (DIC UMOL KG), TA (TA UMOL KG), phosphate (PHOSPHATE UMOL KG), and silicate (SILICATE UMOL KG) content values from bottle samples analyzed in the laboratory, along with CTD measurements in the field of temperature (CTDTMP DEG C ITS90), salinity (CTDSAL PSS78), and pressure (CTDPRS DBAR). Within seacarb, we used the TEOS-10 thermodynamic seawater 285 equations option (IOC, SCOR, and IAPSO, 2010). For equilibrium constants (K_1 and K_2), we did two sets of calculations, using Lueker et al. (2000, "L00") for one and Waters et al. (2014, "W14") for the other, with all other seacarb options the same. We adopted the total scale for pH (pH_T), the Uppstrom (1974) formulation for deriving total boron concentration from salinity, the seacarb default option for K_f (Perez and Fraga, 1987 for temperatures above 9 °C; Dickson and Goyet, 1994 for those below), and the Dickson (1990) option for K_s (following results of Orr et al., 2015). All input content data were first divided by 106 to convert from µmol kg⁻¹ to mol kg⁻¹, and pressure (dbar) was divided by 10 to convert to bar, to conform with the default units of seacarb. Calculated values of fCO2 shown in figures here are the seacarb "in situ" CO2 fugacity values, referenced to in situ temperature and pressure, rather than atmospheric pressure as the "standard" and "potential" options are computed (Gattuso et al., 2023), because in situ results are more germane to understanding the environmental conditions confronting populations of marine organisms in the wild. Total uncertainties on calculated values using high-quality DIC and 295 TA measurements as input parameters are $\pm 3.5\%$ for fCO₂ (± 14 µatm at 400 µatm and ± 70 µatm at 2000 µatm) and $\pm 4.9\%$ for Ω_{arag} (± 0.025 at $\Omega_{arag} = 0.5, \pm 0.049$ at $\Omega_{arag} = 1$, and ± 0.075 at $\Omega_{arag} = 1.5$) (per Orr et al., 2018).

In order to facilitate comparison with results from West Coast Ocean Acidification (WCOA) cruise publications (e.g., Feely et al., 2008, 2016), we used in L00 results in figures, statistics, and discussion. However, for end users using dissociation constants explicitly targeted to the broader salinity ranges seen in the Salish Sea (i.e., W14 constants) for their own measurements, we did a comparison between calculated values using L00 and W14 dissociation constants. This comparison showed that the average differences in calculated values using L00 rather than W14 constants (i.e., L00 – W14) were – 0.003 ± 0.002 for pH_T, $+5.0\pm1.3$ µatm for fCO₂ ($+5.1\pm1.4$ for pCO₂), and -0.001 for Ω_{arag} and Ω_{calc} (±0.003 and ±0.004 , respectively). Only two samples (of 4021) of those with good analytical values for all analytical input parameters had salinity <19—the lower end of the salinity range used by Lueker et al. (2000), and thus generated warning flags. Thus, our use of the L00 constants would not have a discernible effect on the results discussed here. For the L00 vs. W14 comparison, we used the TEOS-10 thermodynamic seawater equations with both sets of carbonate system dissociation constants. To facilitate use of this data set, including the calculated values discussed here, we created a new multi-stressor Salish cruise data product that includes all of the highest quality *seacarb* input data (i.e., temperature, salinity, and DIC, TA, phosphate, and silicate content with quality flags of 2, indicating "acceptable" data quality), along with high-quality recommended O₂ values, and the most commonly used CO₂ system parameters (pH_T, fCO₂, pCO₂, pCO₂, pCO₃, pCO₄ and pCoalculated using both L00 and W14 dissociation constants (Alin et al., 2023b).

However, prior WCOA cruise publications used EOS-80 seawater equations, whereas we used the more recent TEOS-10 equations, in keeping with current recommendations (IOC, SCOR, and IAPSO, 2010; Jiang et al., 2022). The differences between paired calculated carbonate system parameters caused by using EOS-80 vs. TEOS-10 thermodynamic seawater equations with the L00 dissociation constants were similar in magnitude, with average offsets (i.e., EOS-80 – TEOS-10) of – 0.003 for pH_T, +5.2 μatm for fCO₂, 0.000 for Ω_{arag}, and –0.001 for Ω_{cale}.

320 Finally, we use fCO₂ rather than pCO₂ as it provides the most accurate estimate for in situ gas-phase CO₂, because it accounts for interactions between CO₂ and other molecules in seawater (per recommendations in Jiang et al., 2022). To facilitate comparisons between fCO₂ values and pCO₂ values from other sources, the average difference between the two values (i.e., pCO₂ – fCO₂ because pCO₂ is always larger) in this data compilation is 22.5 µatm across all pairs of seawater and carbonate system thermodynamic constants. The difference increased with fCO₂ level, with fCO₂ values averaging 2.8 µatm lower than pCO₂ at 0–499 µatm, 17.7 µatm at 500–999 µatm, 43.4 µatm at 1000–1999 µatm, and 55.7 µatm at >2000 µatm.

3.4 Data visualization

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Raincloud plots display both raw data and percentile distributions to provide transparent statistical data summaries (Allen et al., 2021). We use them to visually summarize the 2014–2018 statistical distributions of temperature, salinity, oxygen, fCO₂, and Ω _{arag} observations from May and October boundary water (S2S) cruises in Figures 3A–4A and 6A–8A and for April, July, and September PS cruises in Figures 3B–4B and 6B–8B. Raincloud statistical summaries for potential density anomaly (sigma

theta, σ_{θ}), Ω_{calc} , and pH_T are provided in Figures S1–S3 for readers interested in this information. Raincloud plots were created using R code by Cédric Scherer (Scherer, 2021) but modified extensively for use with Salish cruise data. To characterize differences in median and extreme values for all parameters with depth, we used 20 dbar as the boundary between surface and subsurface depth categories throughout the region, although we acknowledge that mixing depths vary across the study region and seasons, such that the upper mixed layers occupy different depth ranges through space and time.

Transect plots of all calculated ocean acidification parameters (fCO_2 , Ω_{arag} , Ω_{calc} , and pH_T) were also prepared in *Surfer* can be found in Figures S4–S8. Comparable plots of temperature, salinity, and oxygen, DIC, TA, and nutrient content are shown in Figures 4–9 and S1–S4 of the companion article to this paper (Alin et al., 2023a).

Bubble cloud plots are scatterplots where additional statistics can be represented by the size and colour of each data point. We used the R ggplot2 package ggpubr functions $facet_grid$ and $geom_point$ to create bubble plots summarizing seasonal changes in median values and ranges of T, S, O_2 , fCO_2 , and Ω_{arag} across the study region and by depth during 2014–2018 (Figure 5).

4 Results: Seasonal variability of physical and biogeochemical parameters across depths and basins during 2014-2018

In this section, we describe seasonal oceanographic variation within and across basins for the latter half of the time-series (2014–2018), including differences between surface and subsurface water masses, with ranges serving as our metric of variability. We note the timing and magnitude of apparent anomalies in median or variability for each parameter here and relate these physical and biogeochemical Salish cruise anomalies to the major 2013–2018 weather, hydrological, and circulation anomalies (Table 1) in Section 5, using existing climatologies to provide longer-term context. We refer to cruises occurring in April and May as "spring" cruises, July cruises as "summer," and September—October cruises as "fall." To denote specific cruises, we abbreviate the cruise by the first letter of the month (A, M, J, S, and O, respectively) and the two-digit year (e.g., October 2014 becomes O14). "Spring" cruises also reflect early upwelling season conditions, while July—September cruises represent late upwelling season conditions. "Coastal" refers to stations outside the mouth of the Strait of Juan de Fuca (SJdF), sampling either deep Juan de Fuca Canyon (JdFC) stations or the Ćhá?ba· station in shallower water on the continental shelf (Figure 1). All observations in this compiled cruise data product reflect open basin conditions throughout Puget Sound and its boundary waters, which may be quite different circulation, freshwater influence, and retention times.

4.1 Physical oceanographic seasonality across basins

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During 2014–2018, coastal surface temperature and salinity are largely dominated by seasonal upwelling/downwelling dynamics, as expected. Coastal surface temperatures spanned similar ranges in the early and late upwelling season, whereas deep coastal water had more of a seasonal contrast between early and late upwelling season, with warmer water and wider T

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ranges in the fall (Figure 3A, Table 2). Both surface and subsurface temperatures were warmer by nearly 3 °C during O14, when the MHW was strongest in coastal waters, with less elevated temperatures in surface waters in M16 during the El Niño. Surface salinities were lower in spring than fall, with somewhat fresher anomalies during M17 and O15. October cruises during 2016–2018 had higher surface salinities than O14 and O15 (Figure 4A). However, deep fall salinities during 2014–2015 occupied larger ranges (Table 2). Potential density anomaly values (sigma theta, σ_{θ}) track salinity quite closely throughout this region (Moore et al., 2008) and are not discussed further here but are represented in Figure S1. Depth distributions of physical parameters can be seen in more detail in Figures 4–6 in Alin et al. (2023a).

Seasonal patterns in Strait of Juan de Fuca surface waters were similar, with somewhat narrower surface temperature and salinity ranges than at coastal stations across the upwelling season (Figures 3A, 4A). As seen at coastal stations, subsurface temperature, and salinity, in SJdF showed stronger variability, particularly in fall (Table 2). Temperature anomalies manifested as the widest ranges and highest medians during O14 across the water column, with some residual heat persisting in the form of wider ranges and higher medians across the water column in O15 and O16 relative to O17 and O18 when medians were lower. Subsurface salinity had higher medians than at the surface on all cruises and occupied wider ranges except during M18. Salinity had lower median values across the water column during O14, O15, and O16 than O17 and O18.

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AR showed a seasonal progression of warming between April and July-September cruises, with the median temperatures warming by 1–2° C and variability increasing across depths (Figure 3B). April temperatures were warmer across depths during 2015–2016 relative to 2017–2018, and a similar magnitude of warming was observed across depths between J14 and J15 (Table 2). The seasonal span of salinities at AR overlapped across depths, with greater variability at depth in all seasons (Figure 4B). Median salinity increased across depths by a few salinity units from April to September each year as upwelled deep coastal waters arrive at AR by the end of the upwelling season. Median 2017 AR salinities decreased by ~1 across the water column compared to other years, except at depth in S17.

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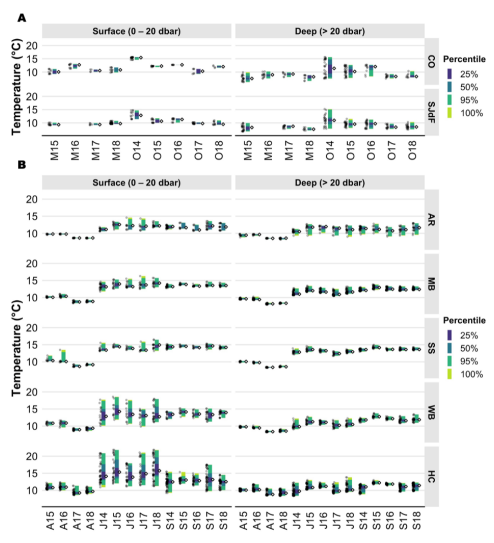


Figure 3: A) Raincloud plots for CTD temperature in coastal (upper row) and Strait of Juan de Fuca (lower row) surveys in the early and late upwelling season beginning in the fall of 2014. Cruise timing is indicated with a one letter month (M=May, O=October) and a two-digit year. Surface observations are in the left column, and subsurface observations are in the right column. Percentiles for observations are

reflected by the colours of the vertical bars, similarly to a box plot, with the median displayed to the right of each bar as an unfilled black diamond and individual observations plotted to the left of each vertical bar as transparent grey circles. Note that all panels in this figure have the same scale bar, but differ from those in the corresponding Puget Sound figure. B) Raincloud plots for CTD temperature in Puget Sound regions—Admiralty (top row), Main Basin (second row), South Sound (third row), Whidbey Basin (fourth row), and Hood Canal (bottom row)—in April, July, and September beginning in July 2014. Cruise timing is indicated with a one letter month (A=April, J=July, S=September) and a two-digit year. Surface observations are in the left column, and subsurface observations are in the right column. Percentiles for observations are reflected by the colours of the vertical bars, similarly to a box plot, with the median displayed to the right of each bar as an unfilled black diamond and individual observations plotted to the left of each vertical bars as transparent grey circles.

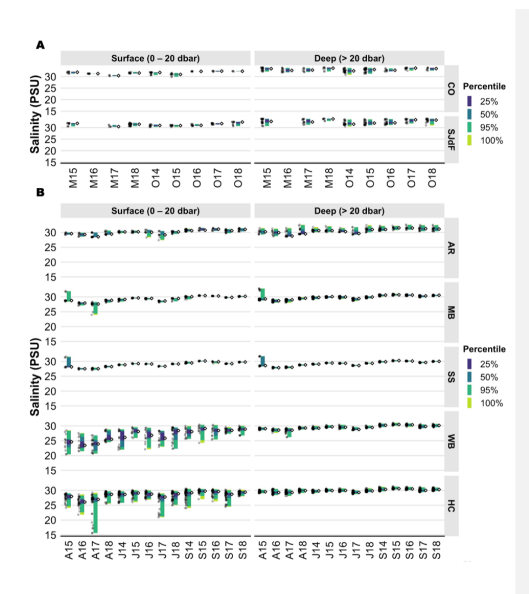


Figure 4: A) Raincloud plots for CTD salinity in coastal and Strait of Juan de Fuca surveys in the early and late upwelling season beginning in the fall of 2014. Figure organization is the same as in Fig. 3A. B) Raincloud plots for salinity in Puget Sound surveys in April, July, and September beginning in July 2014. Figure organization is the same as in Fig. 3B.

MB and SS are the least strongly stratified basins within Puget Sound; they show relatively weak gradients and narrow ranges in temperature and salinity from surface to deep waters, despite MB bottom depths reaching ~225 dbar (Figures 3B, 4B, Table 2). Due to deep mixing, MB and SS are often the warmest at depth of all basins. Temperatures increased by 2–4 °C between April and July–September cruises in both surface and deep water, with typically 1–3 °C difference between surface and deep median temperatures (Figure 3B). Salinity across depths showed progressive increases of ~1–3 across April–September cruises in a given year in both basins, with low variability across the water column (difference of ~1 or less between surface and deep medians, Figure 4B). April temperatures were ~2 °C higher across depth in both basins during 2015–2016 than 2017–2018. Median surface temperatures in J15 and J18 were elevated by ~1–2 °C across MB–SS, and deep J15 and O15 median temperatures were <1 °C higher than in other years (Figure 3B). High salinity outliers (by >3 salinity units) were seen across depth in both basins during A15, and relatively low surface salinity medians were observed during A17 (outliers to ~24) and J17 in MB and SS (Figure 4B, Table 4).

WB and HC have the strongest stratification and gradients of physical and biogeochemical conditions between surface and bottom waters in PS (Figures S5–S8 here and Figures 5–9 and S1–S4 in Alin et al., 2023a). At the entrance to HC, deep-water replacement is constrained by an additional glacial sill and influenced by further mixing of surface with deep water. Subsurface waters in both basins warmed continuously from April through September cruises each year, as they did in MB (Table 2). Deep-water salinity also increased steadily between April and September across WB and HC, with the only obvious anomaly among years being lower salinity in deep WB waters during A17 (lower by ~2). Surface waters in WB and HC were the most variable of all regions for temperature and salinity. The widest ranges of surface temperature occurred in WB and HC during Julys, though median surface temperatures were similar between Julys and Septembers of each year in WB and warmest during July cruises in HC, with cooling by Septembers. Surface salinity tended to be highest and ranges narrower in Septembers, with considerable interannual variability during 2014–2018. Surface temperature anomalies were seen in higher medians in J15 and J18 in HC and J15 in WB, and at depth in both basins in S15, while A15 and A16 had medians ~2 °C warmer across depth in both basins than A17 and A18 (Figure 3B). A16 and A17 had lower median salinities in both WB and HC than A15 or A18 (Figure 4B). All 2017 cruises had anomalously wide ranges and low outliers for salinity in HC.

To summarize across years, regions, and parameters, Figure 5 shows that Puget Sound temperature ranges were mostly shifted up relative to the boundary waters across depths, with higher medians and wider ranges everywhere in summer and fall than spring (see also Table 2). As expected, median salinity was consistently fresher in PS than boundary waters, with substantially wider overall ranges (Figure 5). Vertical temperature and salinity gradients (surface median – deep median) were weakest at AR and SS stations. Temperature gradients were weakest in spring and strongest in summer, while salinity gradients were

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strongest in spring and weakest in fall. HC summertime temperature gradients were the strongest (4.5° C), with WB usually having the strongest salinity variability and surface-deep gradients (4.5 salinity units). While surface variability in T and S tended to be higher in PS surface waters, boundary waters typically showed greater subsurface variability.

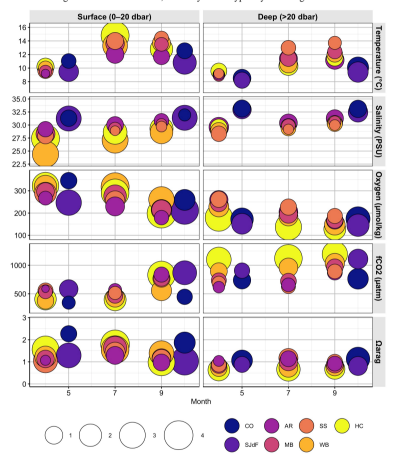


Figure 5: Bubble plots of summary statistics for all five parameters—temperature, salinity, oxygen content, fugacity of CO_2 (fCO_2), and aragonite saturation state (Ω_{arag}) in surface (left column) and deep (right column) water. Bubbles are plotted by the magnitude of mean monthly medians for each parameter taken across Washington Ocean Acidification Center (WOAC, April, July, and September) and Sound-to-Sea (S2S, May and October) cruises during 2014–2018. Bubbles are filled with colours representing the basin the observations were

derived from (CO=Coast, SJdF=Strait of Juan de Fuca, AR=Admiralty Reach, MB=Main Basin, SS=South Sound, WB=Whidbey Basin, HC=Hood Canal). The area of the bubble represents the average range width for that parameter across 2014–2018 WOAC or S2S cruises. The bubble sizes in the legend represent the upper ends of four bins of average range widths (i.e., maximum — minimum) for each parameter: temperature range bins are 0.0-2.5 °C, 2.5-0.0°C, 0.0-7.5°C, and 0.0-1.0°C, salinity range bins are 0.0-2.5 4.6, and 0.0-1.0°C, 0.0-1.0°C, 0.0-1.0°C, and 0.0-1.0°C, 0.0-1.0°C,

Table 2. Ranges of surface (\leq 20 dbar) and deep (>20 dbar) temperature (T, ITS-90), salinity (S, PSS-78), oxygen (O₂), and calculated values of CO₂ fugacity (fCO₂) and aragonite saturation state (Ω_{arag}) for all regions.

		0 6	n m				- n		p 600	0.6	
Regio		Surface	Deep T	Surface	Deep S	Surface O ₂	Deep O ₂	Surface	Deep fCO ₂	Surface	Deep
mon	tn	T (° C)	(° C)	S		(μmol kg ⁻¹)	(µmol kg ⁻¹)	fCO ₂ (µatm)	(µatm)	$\Omega_{ m arag}$	$\Omega_{ m arag}$
Coast											
May											
	2015	9.3-11.2	6.2-9.9	31.3-32.1	32.1-34.0	206–387	77–250	221-713	635–1183	1.07-2.70	0.66-1.16
	2016 2017	11.3-13.1 10.1-10.9	7.6–10.2 8.2–9.9	31.3-31.3 30.2-30.7	31.9-33.8 32.0-33.7	319–391 311–332	129-220 133-293	189-364 281-289	507-888 306-855	1.86-3.19 2.06-2.18	0.89-1.43 0.94-2.20
	2017	9.8–11.7	6.7-8.8	31.3-32.1	32.0-33.7	280-408	82–176	171–525	360–1177	1.37-3.25	0.67-1.82
October		0.7-0.0	31.3-32.1	32.2-34.0	200 400	02-170	171-323	300-1177	1.57-5.25	0.07-1.02	
Octobe	2014	14.6-15.6	7.6-15.6	30.7-31.9	30.8-33.8	241-250	78-248	365-432	371-1399	1.83-2.27	0.58-2.18
	2015	11.9–12.4	7.8–12.7	29.9–31.6	31.3–33.8	254-275	106-261	340-378	358-1005	1.87-2.02	0.79-2.04
	2016	12.7-12.7	8.3-12.7	32.3-32.3	32.3-33.6	265–267	118-268	379–396	376-991	1.95-2.02	0.82-2.03
	2017	9.3-11.2	7.6-9.2	32.1-32.5	32.5-33.9	141-268	52 ^b -171	754-1042	668-2376	0.87-1.16	0.38-1.19
	2018	11.6-12.5	8.0-10.4	32.3-32.4	32.6-33.9	281-326	82-257	300-359	321-1101	2.04-2.40	0.75 - 1.96
Strait a	of Ivan	de Fuca									
May	y ouun	uc I aca									
iviay	2015	9.1-10.1	6.8-9.9	30.5-32.0	30.7-33.9	193-221	96-220	636–777	669-1058	0.96-1.11	0.74-1.07
	2017	9.2–9.9	7.8–9.0	30.3-31.2	31.2-33.8	262-429	120-245	157-528	559-942	1.23-3.09	0.85-1.27
	2018	9.6-10.7	7.1-8.3	30.6-32.1	32.9-33.9	222-357	90–144	310-993	712-1115	0.78-2.15	0.72-1.09
Octobe											
	2014	11.4-14.7	7.8-15.2	30.3-31.3	30.5-33.6	201-248	90-245	394-840	405-1121	0.96-1.94	0.66 - 1.92
	2015	10.0-11.8	8.1-11.5	30.2-31.2	31.0-33.6	230-260	113-239	436-751	512-1044	0.97-1.55	0.77 - 1.38
	2016	10.1-11.7	7.8 - 11.3	30.7-31.6	31.1-33.7	186-257	80-235	432-878	578-1269	0.89-1.67	0.65-1.30
	2017	9.6-10.2	7.7–9.7	31.4-31.9	31.7-33.9	168-195	89-158	654a-2168	1130-2820	0.40-1.20a	0.31 - 0.74
	2018	9.3-11.0	7.6–10.3	30.7-32.6	30.9–33.9	133–209	65–178	803-1099	887–1338	0.74-0.94	0.62-0.86
Admira	ıltv Red	ach									
April											
	2015	9.6-9.9	8.8-9.8	29.2-30.0	29.3-31.5	247-263	199-260	609-650	581-674	1.01-1.08	1.03-1.16
	2016	9.6-9.9	9.3-9.8	28.6-29.9	28.7-313	248-279	221-271	500-544	528-627	1.15-1.19	1.10-1.16
	2017	8.6-8.7	8.4-8.6	28.4-30.0	28.6-31.5	258-299	210-283	425-573	473-679	1.09-1.28	1.02-1.18
	2018	8.5-8.7	8.2-8.7	29.2-30.6	29.3-32.6	243-270	168-265	709-735	670-884	0.84-0.92	0.85 - 0.93
July											
	2014	10.6-11.8	9.3-11.3	29.7-30.5	30.1-31.8	200-241	161-220	569-810	578-823	0.92-1.25	0.93 - 1.24
	2015	11.1-13.4	9.3-12.7	29.8-30.6	30.1-32.1	192-246	156-228	512-684	576-790	1.10-1.47	0.98-1.32
	2016	11.1-14.6	10.1-12.2	28.3-30.9	30.1–31.5	198-296	172-227	443-684	583-773	1.10-1.60	0.98-1.26
	2017 2018	10.8–14.3 11.7–13.5	9.4–12.0	27.4-30.5	29.2–31.8	212–349	153-252	215–559	437–785 596–945	1.28-2.48	0.96-1.51
Septem		11./-13.5	9.0-12.3	29.5-30.5	30.0-32.5	208-262	130-221	470–595	396-943	1.25-1.53	0.84-1.25
Septem	2014	11.1-12.5	9.7-12.1	30.3-31.2	30.5-32.2	173-192	139-188	630-879	756-1018	0.92-1.21	0.80-1.04
	2014	11.1–12.3	10.0–12.1	30.6-31.5	30.7-32.2	178-210	149–200	418–848	745–972	0.96-1.76	0.84-1.10
	2015	10.7–12.5	9.1–11.8	30.5-31.5	30.9–32.6	157–184	122–171	648–1363	791–1563	0.59-1.23	0.51-1.02
	2017	11.1–13.6	9.3–12.3	29.7–31.1	30.4–32.5	151-202	118-175	683–930	799–1067	0.84-1.11	0.76-0.99
	2018	10.8-13.0	9.5–12.9	30.3-31.4	30.4–32.5	157–196	128–190	694–870	765–997	0.92-1.15	0.82-1.06
Mais P											
Main B April	usin										
Арги	2015	9.9-10.4	9.4-10.0	28.4-31.9	28.8-32.8	235-289	206-260	554-709	546-827	0.90-1.13	0.79-1.14
	2015	9.9-10.4	9.4–10.0	27.1–28.2	28.1-29.2	285–289	232–303	293–526	469–826	1.12-1.70	0.79-1.14
	2016	9.9–11.0 8.4–9.2	9.3–10.3 8.0–8.5	24.0-28.3	28.1-29.2	285–349 271–385	242-294	223-658	534-831	0.87-1.70	0.76-1.24
	2017	8.6-9.2	8.2–8.6	28.1-29.1	29.1–29.9	269–287	235–264	462-751	657–1140	0.82-1.23	0.72-1.02
July	2010	0.0-7.2	0.2 -0.0	20.1 -27.1	27.1 27.7	207-207	233-204	102-751	037 1140	0.02-1.23	0.50-0.75
,	2014	11.8-14.6	10.7-12.7	28.4-29.4	29.2-30.2	229-337	184-243	315-627	426-833	1.16-2.01	0.87-1.52

	2015	12.6-15.3	11.6-13.4	29.4-29.8	29.7-30.2	219-322	194-235	443-952	591-998	0.91-1.67	0.80-1.27
	2016	12.9-15.5	11.3-12.8	29.1-29.6	29.3-30.3	224-347	183-253	272-566	495-793	1.29-2.43	0.94 - 1.39
	2017	12.6-16.3	10.5-12.5	28.1-28.7	28.6-29.6	261-419	173-265	155-502	416-842	1.34-3.45	0.82 - 1.54
_	2018	13.0-15.7	11.1-13.2	28.5-29.5	29.5-30.2	243-397	166-242	404-865	578-1133	0.90-1.79	0.67 - 1.28
Septe		120 120		20.2.20.2	20 1 20 0	100.262	156 220	251 (25	5/5 0/0		004 125
	2014 2015	12.8-13.9 13.6-14.3	11.8-13.1 12.3-13.8	29.2-30.3 30.4-30.5	30.1-30.9 30.4-31.1	188-362 197-304	156-220 165-227	271-697 379-780	567–960 495–1084	1.14-2.33 1.09-1.96	0.84-1.35 0.79-1.57
	2015	13.0-14.3	12.3-13.8	30.4-30.3	30.3–30.8	177–204	148–190	711–844	821–990	0.97-1.18	0.79-1.37
	2017	13.2-14.6	12.0-13.3	29.7–29.9	29.9–30.5	183-241	152-191	687–1540	770–1887	0.56-1.14	0.45-1.02
	2018	13.1–14.4	12.2-13.3	29.9–30.3	30.3-30.7	182-239	157–191	577-853	765–968	0.97-1.39	0.84-1.06
South	Sound										
April											
•	2015	10.2-11.8	9.8 - 10.3	28.0-31.6	28.3-31.8	255-417	194-265	169-609	617-964	1.02-2.92	0.70-1.05
	2016	9.8-13.5	9.5-10.0	27.3-27.8	27.6-28.2	278-432	241-294	171-524	535-774	1.09-2.82	0.77 - 1.06
	2017	8.5-9.2	8.3-8.5	27.1–27.9	27.7–28.3	264–315	256-267	445-706	651–769	0.78-1.20	0.74-0.87
July	2018	8.9–9.4	8.4-8.9	28.0-28.5	28.6-29.0	283-386	248-304	247-659	496-814	0.90-2.02	0.75-1.16
July	2014	13.1-15.6	12.2-13.9	28.6-29.0	28.8-29.3	249-304	217-260	351-556	468-685	1.29-2.04	1.06-1.53
	2015	14.1–15.0	12.9-14.6	29.1–29.4	29.3–29.6	241–278	202-252	480-590	554-786	1.32-1.58	0.99-1.41
	2016	13.5-15.2	12.7-13.8	29.0-29.3	29.2-29.6	228-282	191–229	332-589	612-746	1.27-1.98	1.00-1.22
	2017	13.0-15.9	11.7-13.1	28.2-28.5	28.5-28.8	244-347	188-246	322-566	180-788	1.21-2.10	0.88-2.69
	2018	13.2-16.5	12.6-13.7	29.0-29.2	29.3-29.6	236-411	196-247	267-595	587-822	1.23-2.52	0.91-1.26
Septe											
	2014	13.6-15.2	13.0-13.9	29.3-29.9	29.6-30.1	185-333	167-208	356-805	501-792	1.02-2.02	1.03-1.52
	2015	14.3-15.1	13.8-14.9	30.0-30.2	30.0-30.4	136-206	159-202	690–1069	715–1041	0.83-1.24	0.84-1.20
	2016	13.9–14.7	13.3-14.1	29.3-30.0	30.0-30.3	176-235	154-192	680–985	802-1033	0.85-1.20	0.81-1.03
	2017 2018	13.6–15.3 13.7–14.7	13.2–13.4 13.4–14.2	29.1-29.4 29.6-29.9	29.4-29.8 29.8-30.1	160-255 168-288	169-202 173-209	830–2070 471–863	984–1564 647–850	0.45-1.04 0.96-1.63	0.55-0.89 0.97-1.24
	2010	15.7 1	15.1 11.2	25.0 25.5	25.0 50.1	100 200	173 207	171 003	017 030	0.90 1.03	0.57 1.21
Whid	bey Basi	n									
April		<u></u>									
	2015	10.2-11.3	9.5-10.1	20.4-28.5	28.6-29.5	217-410	172-268	146-778	704-1156	0.77-1.97	0.59-0.92
	2016	9.6-11.6	9.3-10.1	21.5-27.6	27.7-29.2	242-376	180-285	140-714	463-1173	0.81-2.09	0.56-1.16
	2017	8.6-9.4	8.2-8.6	20.7-26.8	26.2-29.3	306-389	223-296	110-368	463-922	1.26-2.13	0.66-1.14
	2018	8.6-9.6	8.3-9.1	24.5-28.8	28.8-29.8	204-294	175-263	549-1072	781-1218	0.59-0.86	0.54-0.79
July											
	2014	10.1–17.7	9.1–12.2 10.4–12.7	22.0-28.9 25.9-29.6	28.8-30.0 29.3-30.2	217–454 224–415	159-248	113-763	500-1405	0.86-2.94 1.06-2.67	0.49-1.36 0.53-1.28
	2015 2016	11.9–18.6 10.6–17.6	10.4-12.7	22.1–29.4	28.8-30.4	168-458	134–231 146–195	183–667 269–1212	583-1366 759-1278	0.42-2.02	0.55-0.96
	2017	9.9–15.0	9.2–11.8	22.9-28.4	28.5-29.6	172–384	145-227	143-947	569-1353	0.67-2.55	0.48-1.17
	2018	11.2-17.9	9.7–12.3	22.3-29.5	29.2-30.1	204-441	140-206	119-847	777–1377	0.81-3.12	0.51-0.95
Septe	mber										
	2014	12.1-14.2	11.0-12.4	25.6-30.0	29.6-30.8	157-258	93-176	480-1007	685-1581	0.72-1.27	0.50-1.14
	2015	12.7-15.2	12.2-13.7	24.3-30.4	30.0-31.1	101-365	89-252	210-1529	825-1609	0.54-2.15	0.52-1.03
	2016	12.3-14.9	11.7-12.8	25.4–30.3	29.7–30.9	137–380	97–170	180-1135	890-1549	0.67-2.69	0.51-0.91
	2017 2018	11.3–15.1 12.4–14.6	10.9-12.9 11.0-13.4	27.4–29.6 26.8–30.1	29.2-30.5 29.5-30.7	146-259 200-379	107-179 100-204	514–1324 188–1008	786–1437 730–1692	0.55-1.37 0.76-2.78	0.51-0.99 0.45-1.11
	2010	12.4-14.0	11.0-15.4	20.0-30.1	27.5-30.7	200-377	100-204	100-1000	750-1072	0.70-2.70	0.45-1.11
Hood	Canal										
April											
·	2015	10.2-12.2	9.7-10.6	24.1-29.0	29.0-30.2	201-481	46-321	113-879	369-2439	0.75-3.15	0.32-1.60
	2016	10.0-12.4	9.6-11.8	21.8-28.6	28.4-30.5	268-413	64–293	148-1800	416-2100	0.41-2.60	0.38-1.37
	2017	8.6-11.3	8.5-10.5	15.6-29.2	28.1-30.3	225-467	94-284	68-886	375-1747	0.70-2.42	0.43 - 1.42
	2018	9.0-11.1	8.4-10.5	25.5-29.7	29.1-30.6	142-282	98-269	538-1597	614–1812	0.47 - 1.12	0.44 - 1.03
July											
	2014	10.9-20.9	8.4–12.3	25.8-29.8	29.1–30.2	230–386	84–258	231-709	456-2274	1.02-2.74	0.31-1.54
	2015 2016	12.0-21.8 11.2-17.9	10.2-13.0 10.7-12.3	26.1-30.1 26.9-30.0	29.6-30.6 29.4-30.7	199–388 89–327	12-247 22-210	220-1450 252-2185	478-3460 615-2814	0.57-2.87 0.39-2.57	0.23-1.57 0.28-1.21
	2016	11.5-21.1	9.1–12.4	20.9-30.0	28.7-30.0	223–394	63-255	223–940	408-2259	0.39-2.57	0.28-1.21
	2018	11.2-21.8	9.5–13.8	25.0-29.9	29.4–30.8	99-474	63-246	212-1625	466-2309	0.48-2.85	0.33-1.57
Septe											
•	2014	9.3-15.4	8.8-11.9	24.1-30.5	29.6-31.2	66-389	62-180	292-2611	665-2475	0.28 - 2.02	0.30 - 1.16
	2015	12.0-15.3	11.8-13.0	27.0-30.5	30.3-31.3	79-259	88-213	707-1822	686-1755	0.46 - 1.14	0.48 - 1.15
	2016	11.4–15.0	11.4-12.2	26.3-30.4	30.1-31.2	53-259	75–176	293-2120	919–1769	0.38-2.01	0.46-0.88
	2017	9.7–17.5	9.6–12.1	24.5-30.2	29.4–31.0	106–317	58-175	285-3122	460-3372	0.25-2.29	0.23-1.47
	2018	10.1–14.5	9.8–12.3	28.0-30.2	29.8-31.0	46-291	49–181	156-2641	789–2612	0.29-2.93	0.29-1.02

4.2 Biogeochemical seasonality across basins

4.2.1 Dissolved oxygen

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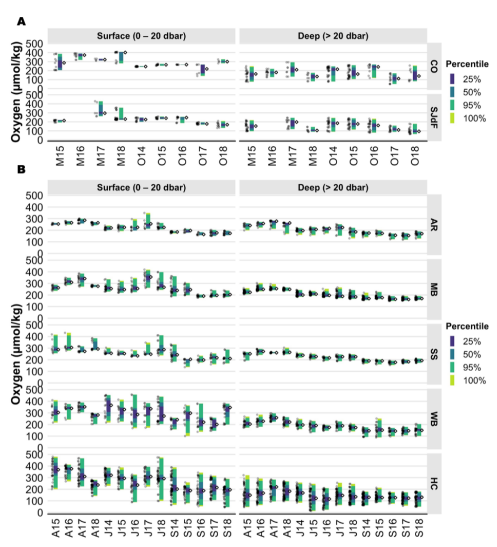
Wide ranges of oxygen content were seen at coastal and SJdF stations, at surface and subsurface depths. Deep-water oxygen observations frequently had higher variability than surface waters, with strong interannual variability in O_2 medians, which were all above the hypoxia threshold (i.e., $62 \mu mol \ kg^{-1} = 2.0 \ mg \ L^{-1} = 1.5 \ mL \ L^{-1}$), and ranges that occasionally dipped into hypoxic conditions (Figure 6A, Table 4). No clear seasonal difference in deep O_2 content emerged between early and late in the upwelling season, but deep-water O_2 medians were lower during O17, M18, and O18 in boundary waters. Boundary water surface O_2 medians were consistently higher than subsurface values during any single cruise, although ranges sometimes overlapped across depths. Surface ranges were wider at SJdF stations, but subsurface ranges were widest at the coastal stations.

Oxygen content in deep AR waters decreased in median, minimum, and maximum values from April to September each year (Figure 6B). Surface O2 also showed decreasing median values April–September, but surface ranges were narrower than at depth in spring and fall, such that surface O2 observations ranges in April and September did not overlap. Wider O2 ranges and higher outliers were observed during J16 and J17 at the surface, and wider ranges with low outliers were seen at depth during A17, J17, A18, and J18.

Open waters of MB and SS were consistently well-oxygenated to the bottom (Figure 6B, Table 4). MB bottom water oxygen was >220 μ mol kg⁻¹ in spring and declined to median values of ~160–180 μ mol kg⁻¹ during September surveys (Figure 6B, cf. Figure 9 in Alin et al., 2023a). Deep waters in SS only fell below 180 μ mol kg⁻¹ twice, with minimum values of <160 μ mol kg⁻¹ during S16 and S17. O₂ content in MB and SS surface waters was always >200 μ mol kg⁻¹ during April and July cruises, and occasionally medians were >300 μ mol kg⁻¹ (e.g., A16 in both basins and A17 and J17 in MB). Across seasons, variability was higher in surface than deep waters, with generally decreasing median values throughout the water column from April to September.

Surface waters in WB and HC had wide ranges of O₂ through spring–fall cruises, typically with lower median O₂ content in Septembers compared with Aprils or Julys, particularly in HC (Figure 6B). Subsurface O₂ variability was lower in WB, but O₂ variability remained high in HC deep water (Table 2). A progressive April–September decline in subsurface O₂ medians was observed in WB, although in HC variability declined more consistently than medians across seasons. Lower surface O₂ medians were observed during J16, A18, and J18 in WB and HC, with higher medians in A15, S15, A16, and S18 in WB. Deep WB waters had higher median values in A17 and high outliers in S15. In deep HC waters, median O₂ values in A17 appear higher than normal, while J15 and J16 O₂ medians and minima were ~25–50 µmol kg⁻¹ lower than other Julys. The

only measurements of hypoxic conditions in Puget Sound were taken in HC, at depth during S14, A15, J15, J16, S17, and S18 cruises, and in surface waters during S16 and S18. 22



495 Figure 6: A) Raincloud plots for adjusted CTD oxygen in coastal and Strait of Juan de Fuca surveys in the early and late upwelling season beginning in the fall of 2014. Figure organization is the same as in Fig. 3A. B) Raincloud plots for adjusted CTD oxygen in Puget Sound surveys in April, July, and September beginning in July 2014. Figure organization is the same as in Fig. 3B.

Looking across basins, surface oxygen content occupied similar overall range widths in spring and fall, while medians declined seasonally by 35–125 µmol kg⁻¹ everywhere (Figure 5, Table 2). Surface O₂ variability was highest across seasons in HC, WB, and SJdF and lowest in AR. Surface variability was highest in summer in PS basins except SS where it was lowest. WB surface O₂ medians and range width peaked and were highest among PS basins in summer, approaching spring coastal surface O₂ median values. Everywhere else, surface O₂ medians decreased from spring to fall. At depth, O₂ content decreased monotonically by 52–94 µmol kg⁻¹ across all PS basins from April to September, with the largest decrease at AR and the smallest decrease across seasons in HC. In contrast, median O₂ in deep boundary waters remained roughly the same from spring to fall, with variability often exceeding surface variation. Ranges in deep O₂ content were widest in HC during spring–summer, followed by coastal and SJdF stations. While we observed hypoxic conditions in surface waters and near-anoxia at depth in HC, O₂ concentrations below the hypoxia threshold have not been observed elsewhere in PS during these cruises. O₂ concentrations were consistently second lowest at the river end of the WB basin, with lowest O₂ conditions occurring consistently in September, even during heatwave years when deep HC O₂ was lowest during Julys.

4.2.2 Carbon dioxide fugacity (fCO2)

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Subsurface CO₂ fugacity values typically had lower highs, lows, and medians at coastal than SJdF stations (Figures 7A and S4, Table 2). Coastal surface median values were thus often undersaturated with respect to atmospheric fCO₂, whereas SJdF medians were often above atmospheric values. As for O₂, no clear seasonal fCO₂ difference was evident, either at the surface or at depth. Coastal stations had higher surface median fCO₂ in M15, but the most notable variation was the anomalously wide fCO₂ ranges observed during O17, with deep-water medians >1000 μatm at coastal stations (high=2376 μatm) and ~1900 μatm at SJdF stations (high=2820 μatm). O17 SJdF surface fCO₂ values were unprecedented, with a median of 1575 μatm and highs up to 2168 μatm. Coastal surface fCO₂ anomalies were also notable during O17, with the only observations of fCO₂ >1000 μatm occurring then. SJdF M18 subsurface and O18 median fCO₂ observations across depth were also somewhat elevated compared to other cruises, suggesting possible carryover of the anomalously acidified water masses from the previous fall.

AR fCO₂ medians and ranges increased between April and September, with most cruises having narrow ranges and little depth structure (Figure 7B, S5, Table 2). The relatively wide overall AR fCO₂ ranges reflect low surface outliers in J17 and high outliers across depths in S16. The latter co-occurred with the lowest O₂ median in AR surface waters in this cruise time-series (cf. Figure 6B).

In MB and SS, fCO₂ variability across depths was lower during April–July than September (Figures 7B, S5, Table 2). The majority of the MB-SS observations >1000 μatm were associated with extremely high fCO₂ anomalies in S17. Extreme fCO₂ conditions in S17 were preceded by anomalously low fCO₂ in MB surface waters in J17 and to a lesser extent in A17, which

Deleted:

we interpret as reflecting a protracted season of biological drawdown due to the co-occurrence of high O_2 , low fCO_2 , and sustained high chlorophyll (PSEMP Marine Waters Workgroup, 2018).

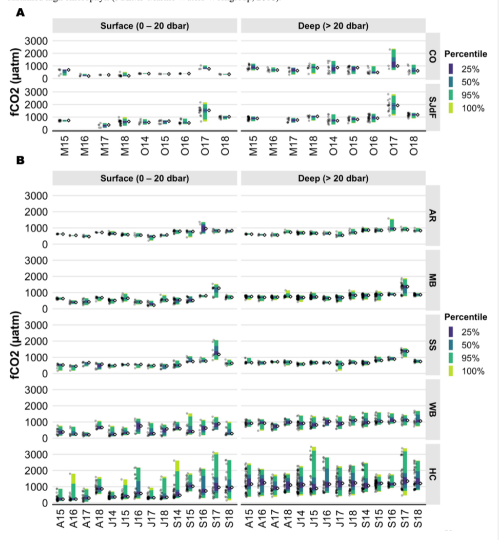


Figure 7: A) Raincloud plots for the fugacity of carbon dioxide in coastal and Strait of Juan de Fuca surveys in the early and late upwelling season beginning in the fall of 2014. Figure organization is the same as in Fig. 3A. B) Raincloud plots for the fugacity of carbon dioxide in Puget Sound surveys in April, July, and September beginning in July 2014. Figure organization is the same as in Fig. 3B.

Variability in fCO₂ was highest across regions in HC and second highest in WB, with similar range widths in surface and subsurface waters in both basins (Figures 7B, S5, Table 2). Surface fCO₂ medians tended to be lower in Aprils and Julys in both basins. HC had higher high fCO₂ values across depths and months than WB. Cruise medians were within ±230 μatm between WB and HC surface waters in Aprils and Julys, but HC surface water medians were more supersaturated with fCO₂ in all Septembers except S14. HC surface waters had particularly high fCO₂ outliers during S14, S17, and S18, with the highest deep water HC values seen during J15 and S17. WB surface fCO₂ had the highest outliers during S15 and an anomalously high median during A18, which could reflect persistence of acidified conditions from fall 2017.

The majority of the water column was supersaturated with respect to atmospheric fCO₂ values (i.e., >400 μatm) in most places and times for the duration of this time-series (Tables 4–5), reflecting the importance of respiration processes in Salish Sea CO₂ chemistry, although moored time-series show that surface undersaturation prevails in spring–summer (Alin et al. in PSEMP Marine Waters Workgroup, 2021; Fassbender et al., 2018). Average surface fCO₂ medians and ranges were lower and narrower, respectively, during spring and summer months than fall across regions. As for temperature, salinity, and oxygen, HC, WB, and SJdF had the largest surface variability (Figure 5). Median surface fCO₂ values were lowest across coastal, WB, and HC stations in spring, and in HC and WB among PS basins in summer. The highest spring surface median values were in SJdF and AR, reflecting vigorous mixing bringing deep CO₂-rich waters to the surface. In fall, HC had comparably high surface median fCO₂ values to SJdF and AR, with the highest variability in HC and SJdF. In deep water, median fCO₂ values were higher than at the surface across seasons by ~30–400 μatm at AR, MB–SS, and boundary water stations, and by ~350–725 μatm in WB and HC. Surface and deep fCO₂ levels overlapped most at AR in all seasons and in HC during fall, when local upwelling brings deep water enriched in fCO₂ to near-surface depths as it is flushed out of the basin via estuarine circulation (Figures 7 and S5). Median deep fCO₂ values were highest in HC across seasons, although SJdF and WB approached HC levels at depth during fall cruises. Coastal deep median fCO₂ values were comparable to MB and SS and higher than AR values in spring but lowest across regions in fall.

560 4.2.3 Aragonite saturation state (Ω_{arag})

Both surface and subsurface boundary water Ω_{arag} spanned undersaturated to quite supersaturated (> 1) values, with strong surface variability and lower highs and lows in SJdF (Figure 8A, Table 2). Spring surface Ω_{arag} ranges were wider in both regions than fall ranges, with more variable median values, particularly at the coast. The highest deep medians occurred in O14 in SJdF and O16 at coastal stations, and the lowest medians were seen in O17 in both regions. Notably high surface median Ω_{arag} values were observed in M16 and M18 at coastal stations and M17 at SJdF stations, with notable surface low medians during O17 at both coastal and SJdF stations, as well as during M15 at coastal stations.

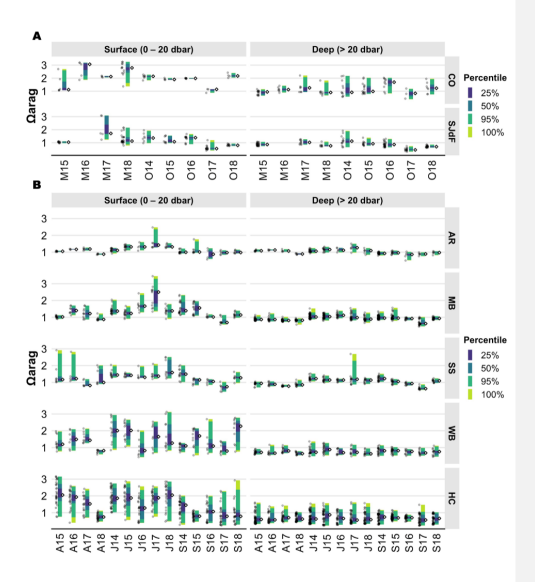


Figure 8: A) Raincloud plots for aragonite saturation state in coastal and Strait of Juan de Fuca surveys in the early and late upwelling season beginning in the fall of 2014. Figure organization is the same as in Fig. 3A. B) Raincloud plots for aragonite saturation state in Puget Sound surveys in April, July, and September beginning in July 2014. Figure organization is the same as in Fig. 3B.

In contrast to O_2 and fCO_2 , surface Ω_{arag} medians at AR tended to be highest in Julys, followed by Aprils, with Septembers having the lowest values. The same pattern was evident but weaker at depth at AR. Ranges for Ω_{arag} were typically widest in July (Table 4). Anomalously wide ranges in Ω_{arag} were seen in surface waters with high anomalies in S15 and J17 and with a lower median and lows in S16.

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Whereas fCO_2 variability was similar across depths in MB and SS, Ω_{arag} generally had substantially higher variability in surface waters than deep (Figure 8B, Table 2). Both surface and subsurface MB and SS waters had highest Ω_{arag} medians in Julys, with lower medians and significant interannual variability across Aprils and Septembers. Median deep SS Ω_{arag} observations were above saturation during Julys and some Septembers, whereas MB subsurface medians were typically below the saturation threshold ($\Omega_{arag} = 1$), reflecting the greater volume of deep water in MB. Exceptions occurred during J14, J15, and J16, when MB deep waters had medians >1, and J17 in SS, which had high deep-water outliers. Median surface Ω_{arag} values were supersaturated for all cruises except A18 in MB, A17 in SS, and S17 in both basins. Notably high Ω_{arag} outliers in surface waters were observed during A15, A16, and J18 in SS and J17 in MB, which also had the highest median observed across all basins and seasons in PS. Notably low Ω_{arag} medians occurred in S17 in both basins and across depths.

In WB and HC, Ω_{arag} ranges were much wider in surface than subsurface waters, with wider ranges at both depths in HC than WB (Table 4). Ω_{arag} ranges tended to be widest in WB surface waters in July, with no clear seasonal pattern in surface medians (Figure 8B). Notable WB surface Ω_{arag} medians were >2 in J14, J15, and S18 and <1 in A18, J16, and S17. Surface HC Ω_{arag} medians were notably low in A18 and J16 and high in S14 (Figures 8B and S6). Deep-water Ω_{arag} medians were more stable across months and consistently undersaturated.

In summary, median aragonite saturation states were substantially higher in surface coastal waters in spring and fall than throughout SJdF and PS basins (Figure 5). HC had the highest surface variability across seasons, with the highest PS medians in spring and summer and the lowest in fall. AR had the least inter- and intra-seasonal variability of all regions in both surface and deep water. Deep HC medians were lowest across months, with consistently high variability reflecting considerable alongbasin gradients at depth (Figure S6). Deep WB Ω_{arag} medians fell between those for PS and HC, but the range widths were more similar to PS basins than HC. Notably low Ω_{arag} anomalies occurred in fall 2017, with indications that acidified conditions were held over until A18 observed in multiple basins.

4.3 The role of distinct seasonality across parameters and basins in driving severity of acidification and hypoxia

Average ocean conditions from the coast through Puget Sound are summarized in bubble plots of each parameter for each month, region, and depth across all 2014–2018 cruises (Figure 5). Seasonal variation of median values and ranges across basins was not consistent across parameters. For instance, surface salinity seasonality was different from temperature seasonality across PS basins. Surface water temperatures peaked in some PS basins in summer, following solar radiation (Figure 2), while deep-water temperatures continued to rise until fall, except at AR (Figure 3B). In contrast, PS salinities progressively increased and ranges contracted from spring to fall, tracking seasonal precipitation and river discharge patterns (Figure 4B, references in Table 1, and Banas et al., 2015). The continued increase of deep temperature and salinity until fall reflects a combination of surface conditions mixing to depth and the influence of upwelling bringing colder, saltier, lower oxygen water masses into the Salish Sea, and displacing the fresher, warmer, more oxygenated water masses that are present in MB and SS in winter. Moored time-series provide a longer-term, more temporally resolved context for the seasonal variability across parameters and PS basins observed in the Salish cruise time-series and confirm that water mass properties do not vary consistently across PS basins. Specifically, HC deep-water seasonal peaks and troughs for temperature lag those from other basins by a few months (https://nwem.apl.washington.edu/prod_PS_ClimateTrends.shtml). Salinities reach their peaks and nadirs 1–2 months after temperature in both surface and deep waters across all PS moorings.

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 O_2 medians in most basins and across depths declined steadily from spring to fall, whereas Ω_{arag} medians across depths usually peaked in summer and fCO_2 levels typically increased most substantially by fall (Figure 5). Variability in O_2 and Ω_{arag} was markedly lower in subsurface than surface waters, although deep-water HC ranges were still wide (Figures 6, 8, Table 2). However, subsurface fCO_2 ranges were typically as wide as surface ranges, and in the case of HC, often wider (Figure 7, Table 2), which likely reflects the amplification of fCO_2 variability that occurs when buffering capacity declines (Pacella et al., 2018; Kwiatkowski and Orr, 2018).

Oxygen and inorganic CO_2 dynamics often mirror each other within a local water mass because they are linked by the stoichiometry of biological production and respiration processes, but these can be decoupled across depth (e.g., Cai et al., 2011; Feely et al., 2010). Surface fCO_2 and O_2 levels dominantly reflect phytoplankton bloom activity, which peaks from spring to summer throughout the study region (e.g., PSEMP Marine Waters Workgroup, 2019 and earlier years). Organic matter from phytoplankton blooms subsequently drives the regional spring-to-fall O_2 decrease and fCO_2 increase via respiration in both surface and subsurface waters. While CO_2 drawdown also affects saturation states, Ω_{arng} peaked in summer in many basins, reflecting stronger temperature influence on carbonate ion concentrations than fCO_2 in surface waters (Figure 1 in Cai et al., 2020). In deep water masses, biogeochemical parameters tended to follow more monotonic seasonal trajectories, with T, S, and fCO_2 increasing and O_2 decreasing across spring—fall as a result of longer residence times and respiration contributing to higher fCO_2 (e.g., Feely et al., 2010, 2023). Thus, seasonal decoupling across metrics of ocean acidification and oxygenation

reflects the relative importance of physical versus biological control on each parameter, which have strong gradients across the estuarine to coastal to open ocean continuum (Kwiatkowski and Orr, 2018; Lowe et al., 2019; Cai et al., 2020).

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Interpreting Salish cruise seasonal patterns in the context of the moored climatologies across PS shows that deep climatological temperature and salinity minima tend to co-occur with maximum annual deep O2 values and vice versa from AR to SS, whereas annual low and high values of T, S, and O2 occur synchronously in HC. The lag in temperature and salinity seasonality in deep HC waters is consistent with the longer residence and transit times along the deep axis of HC compared to other basins (Babson et al., 2006; MacCready et al., 2021). While deep-water renewal in MB, SS, and WB basins follows mixing of incoming upwelled water with outgoing surface water at AR, deep-water renewal in HC requires incoming marine waters to pass over a second glacial sill before transport along the long deep axis of HC, contributing to lags in temperature and salinity seasonality in deep HC waters. Consequently, mid-summer to fall bottom water in HC was often colder and fresher than in MB-SS, while in winter-spring, bottom water toward the southern end of the HC basin was frequently warmer and saltier (Figures 5-6 in Alin et al., 2023a). Peak surface O2 values occur earliest in HC, followed by SS and AR, with MB peaking latest, spanning spring to summer. Climatological low surface O₂ values span fall to winter, occurring earliest at AR, followed by MB, SS, and finally HC. Collectively, these observations suggest that the earlier surface O2 peak (fCO2 nadir) in HC surface waters, which along with higher chlorophyll values imply earlier seasonal onset and peak of primary production (PSEMP Marine Waters Workgroup, 2018), translates to earlier O₂ depletion in deep waters driven by remineralization of sinking phytoplankton. The earlier peaks in production and respiration in HC thus effectively offset the deep-water lag in O2 minimum (and fCO2 maximum) timing relative to T and S seasonality seen in other basins. This is important because the interaction and timing of seasonal peaks in physical and biogeochemical processes drives in the formation of hypoxic, corrosive conditions in HC deep waters. Thus, future changes in deep-water renewal timing—as observed during the MHW—and the phenology of surface biological processes may influence deep-water conditions differently across the distinct basins of this complex ecosystem and could result in new acidification or hypoxia hotspots emerging.

5 Discussion: How did major environmental anomalies of 2013-2018 affect physical and biogeochemical conditions?

The 2014–2016 part of the Salish cruise time-series coincided with the protracted, intense heat anomaly initiated by the 2013–2015 NE Pacific marine heatwave (MHW) and extended by the 2015–2016 El Niño event (Bond et al., 2015; Jacox et al., 2016; Gentemann et al., 2017; Peterson et al., 2017). The combination of the Pacific Ocean heat anomalies and a protracted atmospheric heat anomaly affecting the western U.S. during 2014–2015 (Swain et al., 2016) altered atmospheric and seawater temperatures in the Salish Sea (Table 1 and references therein). Record precipitation occurred during at least one month per year throughout 2014–2018, with river discharge timings showing strong anomalies tending toward earlier, higher peak runoff and lower summer flows than in the historical baseline (Table 1 and references therein). Below we connect how the timing of these environmental anomalies contributed to the oceanographic anomalies observed in Washington's marine waters, focusing

on understanding the genesis of summer and fall deep-water anomalies, because the strongest hypoxic and acidified conditions typically occur then. We focus on the oceanographic responses of boundary waters, weakly stratified MB, and strongly stratified HC to examine whether the 2013–2018 environmental anomalies caused changes in timing or decoupling of physical or biogeochemical processes such as those described in Section 4.3, and in so doing, worsened or ameliorated ocean acidification and hypoxia at new locations or times.

5.1 Physical oceanographic anomalies during 2014-2018

higher temperatures within the estuary.

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Temperature and salinity anomalies observed in the 2014–2018 Salish cruise time-series did not always occur synchronously in space or time. The intense NE Pacific marine heat anomalies manifested as surface temperature medians in O14 that were mostly outside the envelope of other 2014–2018 boundary water temperatures (Figure 3). At depth, the widest boundary water temperature ranges were observed in O14. Wider deep-water ranges and higher medians persisted through 2015–2016. A larger-than-normal volume of warm coastal surface waters was also observed during the O16 cruise (Figure 4 in Alin et al., 2023a), but temperatures were lower than O14. Nearby coastal time-series showed that warm anomaly events of up to 4.5 °C lasting for 10–20 days and affecting the water column to bottom depths of 40 m during 2014 and 2015 (Koehlinger et al., 2023).

680 Comparison of summer-fall PS cross-sections before and after the arrival of heatwave-warmed waters on the coast in September 2014 (Peterson et al., 2017; Koehlinger et al., 2023) provides perspective on the timeline of MHW warming across PS basins (Figure 5 in Alin et al., 2023a). Within PS basins, surface warming reflected regionally elevated air temperatures and greater than average solar radiation (Table 1), with surface warm extremes of ≥21° C during J15, J17, and J18 in southern HC being >2.5° C above monthly averages from Fassbender et al. (2018) (Figure 5 in Alin et al., 2023a, Tables 2-3). Peak 685 MHW surface temperatures were cooler overall in MB, but still 1.8-3.1°C above average (Fassbender et al., 2018). Subsurface water temperatures did not reflect the strong MHW warming seen on the coast during 2014, but temperature increases were observed throughout the water column in most basins throughout 2014-2016, with some deep-water warming evident through 2018 (cf. August-October cruises 2008-2014 in Alin et al. 2023a Figure 5). For instance, deep waters in southern HC warmed by ~2 °C during Julys and Septembers of 2015–2016 relative to the same months in 2014, subsequently cooling by ~1 °C in 2017-2018. A deep mixing event in HC during anomalously cold weather in February 2014 caused cooler deep water to persist in HC through 2014; thus, it is unclear whether deep HC September temperatures in 2017-2018 reflect a return to normal (PSEMP Marine Waters Workgroup, 2015). For comparison, heat from the NE Pacific marine heatwave of 2013–2015 lingered at depth until at least early 2018 in a fjord to the north of the Salish Sea, indicating pronounced persistence of the MHW warming signature in other deep, stratified basins in the region (Jackson et al., 2018). Fjords with bathymetric sills are known to retain waters due to refluxing over the sills (e.g., MacCready et al., 2021), which would serve to prolong the anomalously Boundary water observations in O14 reflected a fresher water column at the time when the MHW arrived in these waters than during any other late upwelling season cruise in this time-series (Table 2), consistent with a warmer, fresher water mass from the NE Pacific being advected to the Washington and Oregon coastline in fall 2014 (Peterson et al., 2017; Koehlinger et al., 2023). Octobers salinities during 2014–2016 remained fresher by ~1, overlapping more with spring salinities than O17 and O18 and illustrating the longevity of the fresher water masses associated with the MHW at the coast. The combination of higher temperatures and lower salinities in boundary waters during O14 and O16 manifested as substantially less dense waters to 100 m depth (Figure S4).

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Within PS, salinity anomalies related to precipitation and winter–spring river discharge anomalies should manifest as freshening anomalies during spring. The lowest surface salinities were observed in WB and HC during Aprils of 2015–2017, and Julys of 2014 and 2017–2018 (Figure 6 in Alin et al., 2023a, Table 2). Lower A17 and J17 salinities in MB and HC reflect the wettest February–April on record and strong PS river runoff January–March (Figure 4B, Tables 1–2), with salinities two standard deviations below average also observed in PS moored time-series (Ruef et al. in PSEMP Marine Waters Workgroup, 2018). Lower salinities persisted throughout the water column in all PS basins across seasons in 2017 (Figure 6 in Alin et al., 2023a). Spring 2017 surface salinities were fresher in boundary waters and at depth on the Washington shelf (the latter observed by Mickett and Newton in PSEMP Marine Waters Workgroup, 2018), but the freshening signature had disappeared from boundary waters by O17. Other data sources show that the Salish Sea experienced one of the top five annual total freshwater inflow years since 1999 during 2017 (Khangaonkar et al., 2021).

Conversely, summer drought conditions increased in severity and duration across 2015, 2017, and 2018 (Table 1 and references therein) and could be reflected by higher-than-normal salinity anomalies during September cruises (Figure 6 in Alin et al., 2023a). The lowest surface fall salinities in WB and HC were observed in S14 and S16, whereas in drought years, minimum surface salinities were ~1–2 salinity units higher in September (Table 2). Compared to monthly average surface salinities from moorings, the lowest salinity values seen in HC in S15 and S17 were up to 2.5 salinity units higher than average (Fassbender et al., 2018). In MB, the persistent fresher conditions (by ~0.5 salinity unit) in the upper water column during S17 suggest that low river discharges during the summer 2017 drought may have slowed estuarine circulation, allowing the fresher, more stratified conditions from earlier in 2017 to persist (Table 1; *cf.* Fassbender et al., 2018).

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Table 3. Distribution of temperature (T), oxygen (O₂), calcite and aragonite saturation states (Ω_{calc} , Ω_{arag}), and carbon dioxide fugacity (fCO₂) observations in the Salish cruise data package relative to thresholds with potential implications for altering carbon cycle fluxes or affecting physiological processes or survival in Salish Sea species. Boundary waters includes both coastal and Strait of Juan de Fuca stations. The total number of observations (Total n), number of surface observations (Surface n, \leq 20 dbar), and percent of each parameter's observations (%) in the Salish cruise data package crossing the threshold for each parameter are given in the three respective righthand columns, with total observations by basin broken down in columns to the left.

Tri 1 11	Boundary	Admiralty	Main Basin	South	Whidbey Basin	Hood Canal	Total na	Surface	%
Threshold	waters	Reach		Sound				n	
T ≥ 21 °C ^b	_	_	_	_	_	9	9	9	0.1
T ≥ 19 °C ^b	_	_	_	_	_	38	38	38	0.5
$T \ge 15$ °Cg	28	1	10	25	31	114	209	203	2.8
$O_2 \le 62 \mu mol \ kg^{-1(c)}$	36	_	_	_	_	64	100	19	1.3
$O_2 \le 110 \mu mol \ kg^{-1(b,c)}$	220	2	_	_	26	386	634	76	8.4
$O_2 \le 155 \mu mol \ kg^{-1(b,c)}$	471	38	15	5	165	925	1624	181	21.7
$\Omega_{\rm calc}$ < 1 d	70	9	13	17	78	441	628	131	14.3
$\Omega_{ m arag}$ < 1 $^{ m d}$	599	330	444	226	234	873	2706	695	61.2
$\Omega_{\rm arag}$ $< 1.2^{\rm e}$	800	379	571	364	253	1073	3440	984	78.6
$pH_T < 7.80^{f}$	626	380	486	254	218	890	2861	722	64.3
$pH_T < 7.70^{f}$	331	95	94	56	172	728	1476	286	33.2
$pH_T < 7.65 g$	190	21	40	35	121	631	1038	187	23.3
$pH_T < 7.52^{f}$	23	1	6	15	27	323	395	106	8.9
$fCO_2 < 400$ μ atm ^h	123	3	46	27	59	160	412	392	9.3
$fCO_2 = 401 - 1000 \mu atm^{i}$	674	426	675	451	140	517	2898	1143	65.1
$fCO_2 = 1001 - 2000 \mu atm^{i}$	243	25	49	38	123	526	1010	167	22.7
$fCO_2 = 2001 - 3000 \mu atm^{-1}$	7	_		_	_	116	124	32	2.8
$fCO_2 > 3000$ μ atm ⁱ	_	_	_	_	_	5	5	1	0.1

⁷³⁵ Total numbers of data points in the Salish cruise data package are 7526 for temperature, 7492 for oxygen, and 4449 for inorganic carbon measurements. b Migration blockages for adult salmonids occur at 19-23 °C, particularly in combination with oxygen levels below 3.5 mg L-1 (~110 μmol kg-1) to 5 mg L-1 (~155 μmol kg-1) (McCullough et al., 2001; U.S. Environmental Protection Agency, 2003). Thresholds for sublethal to lethal hypoxia impacts range from 0.7-2.5 mg L⁻¹ for various invertebrate taxa to 1.5-4.4 mg L⁻¹ for fish (Vaquer-Sunyer and Duarte, 2008); the threshold of 2.0 mg L⁻¹ is commonly used to delineate hypoxic conditions (\sim 62 μ mol kg⁻¹ = 1.4 mL L⁻¹), with 0.7 740 mg L⁻¹ (~31 µmol kg⁻¹=0.5 mL L⁻¹) as a threshold used for "severe hypoxia" in the oceanographic literature (e.g., Grantham et al., 2004; Chan et al., 2008). d Thermodynamic saturation threshold for calcium carbonate saturation states (e.g., Zeebe and Wolf-Gladrow, 2001; Dickson et al., 2007). e Represents severe dissolution and growth exposure thresholds for calcifying pteropods (14 days at Ω_{arag} =1.20 and 7 days at Ω_{arag} =1.15, respectively; Bednaršek et al., 2019). Decapod sensitivity thresholds from Bednaršek et al. (2021b) as described in text. g A multi-stressor vulnerability analysis specific to Dungeness crab used temperature, oxygen, and pH_T thresholds of 15 °C, 62 µmol kg⁻¹, 745 and 7.65, respectively, after testing a range of values for each parameter (Berger et al., 2021). h The 400 µatm fCO2 value represents the approximate atmospheric CO2 mole fraction (xCO2) during 2008-2018 (the range of mean annual global marine boundary layer atmospheric xCO₂ across 2008-2018 is 385-408 ppm, per NOAA Global Monitoring Laboratory, 2022). Below atmospheric levels, uptake of atmospheric CO2 by surface water air-sea exchange is favoured; above atmospheric levels, outgassing or evasion from surface waters to the atmosphere is favoured. Values of xCO₂ are ~2.5% higher and pCO₂ values are ~0.4% higher than fCO₂ at seawater surface atmospheric pressure and are different as a result of considering relative humidity and molecular interactions within the measured sample (Dickson et al.,

2007). ⁱ fCO₂ levels above atmospheric values have been divided into broad bins based on thresholds used in the literature to project hypercapnia impacts in fish and invertebrates (McNeil and Sasse, 2016) and behavioural to gene expression responses observed in Pacific salmon (Williams et al., 2019).

5.2 Biogeochemical anomalies during 2014-2018

When the atmospheric and marine heatwaves that warmed Salish Sea surface and deep waters began, a simplistic expectation would have been that increased temperatures would drive increased rates of respiration rates in deep waters, assuming adequate supply of organic matter. If a temperature-driven increase in respiration had dominated, we would have expected to see lower-than-normal O₂ and Ω and higher fCO₂ in deep water masses during high temperature anomalies. In contrast, either surface warming or stronger-than-normal freshwater input could have limited primary production by cutting off surface nutrient supply under stronger stratification, resulting in reduced supply of organic matter and higher-than-normal O₂ and Ω and lower fCO₂ at depth. Additionally, input of NE Pacific source waters at depth with a stronger surface water signal (due to the reduced deep mixing during the MHW) may have also contributed to the higher O₂ and Ω and lower fCO₂ at depth. However, biogeochemical anomalies observed in this time-series did not show a simple temperature-dependent response to the heat anomaly, but rather reflected a combination of heat and river discharge influences.

765 5.2.1 Hypoxia anomalies

We know from observations and models that hypoxia often occurs in hotspots further south than Salish cruise stations on the Washington shelf during late summer (Connolly et al., 2010; Peterson et al., 2013; Siedlecki et al., 2015, 2016b; Barth et al., 2024), and the resulting hypoxic water masses can be advected northward into our study area (Mickett and Newton in PSEMP Marine Waters Workgroup, 2018). Deep boundary waters anomalies observed during the heat anomaly were the opposite of the temperature-driven expectation, with a better oxygenated water column during O14 than earlier cruises this time of year (cf. Au08, O11, S13 in Figure 4 in Alin et al., 2023a). Fall boundary water cruises after O14 never captured hypoxic waters in JdFC and measured hypoxic conditions at one shelf station during O17, although deep waters were closer to the hypoxia threshold in O17 and O18 than O14 and O16. The presence of deep, well-oxygenated water to greater depths than normal in boundary waters during O14 is consistent with the advection of fresher, warmed, well-oxygenated water from the NE Pacific gyre that moved on shore and dominated the upper water column during 2014–2015 (Siedlecki et al., 2016a; Peterson et al., 2017).

Hypoxic conditions were not observed in MB in this cruise time-series or in moored near-bottom time-series (Figure 9 in Alin et al., 2023a and https://nwem.apl.washington.edu/prod_PS_ClimateTrends.shtml, respectively), likely due to the degree of mixing and shorter residence times (e.g., Babson et al., 2006; MacCready et al., 2021). Minimum deep O2 values in MB during S15 and S18 were somewhat higher than during pre-MHW fall cruises, while in S16 and S17, they were slightly lower. Surface O2 levels were particularly high in MB during A17 and J17, and to a lesser extent in A16 and J16 (Figure 6, Table 4, and Figure 9 in Alin et al., 2023a). In combination with observations of sustained MB phytoplankton blooms during April–August

2017 (PSEMP Marine Waters Workgroup, 2018), these high surface O₂ anomalies suggest that high phytoplankton biomass during spring of 2016 and spring–summer 2017 provided stronger inputs of organic matter to deep MB waters in both years, resulting in lower O₂ levels due to enhanced respiration at depth by the fall of each year.

In contrast, deep HC waters develop hypoxia during some years, typically in August–September in southern HC (Feely et al., 2010; Newton et al., 2011). Hypoxic water masses were observed during O11 and S14 cruises in the process of being circulated out of the deep southern HC basin by the fall marine intrusion (Figure 9 in Alin et al., 2023a). O2 content in HC deep water was not exceptionally low in S14 compared to earlier hypoxia years. However, during continued higher temperatures through 2015–2016, anomalously low O2 conditions in deep southern HC waters were apparent as early as April of both years (compared to A17, A18, and https://nwem.apl.washington.edu/prod_PS_ClimateTrends.shtml) and worsened considerably by J15 and J16. The most extensive and severe hypoxic conditions captured in southern HC during the 2008–2018 Salish cruise time-series were observed during J15 and J16 MHW cruises (Figure 6, Table 2). Unexpectedly, by S15 and S16, deep-water O2 minima were higher than in J15 and J16 as well as during S14, S17, and S18 cruises. In 2015 and 2016, early marine intrusions flushed out deep waters and improved conditions sufficiently by September that smaller volumes of low-O2 water remained than during a relatively good pre-MHW year (cf. S09 in Figure 9 in Alin et al., 2023a; references in Table 1). In contrast, S17 and S18 O2 levels in deep HC water reflected hypoxic conditions equivalent to a bad pre-MHW year, in both timing and magnitude.

5.2.2 Acidification anomalies

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Late summer–early fall fCO₂ values in surface boundary waters ranged from <400 μatm to >800 μatm in surface waters before the heat anomaly, with waters below 100–150 m consistently >1200 μatm (Figure S4). Comparable pre-MHW Ω_{arag} values spanned ~1 to >2, with the lowest deep values being <0.6 (Au08, O11, S13 in Figures 7 and S4). During O14 and particularly in O16, smaller volumes of >1200 μatm water and the deepest aragonite saturation horizon (depth where Ω_{arag} =1) were observed in boundary waters. These observations are consistent with NE Pacific source waters with a lower respiration signal having been advected to the Washington coast and persisting in the water column to significant depth during late 2014–2016 (cf. Franco et al., 2021). In contrast, the highest fCO₂ values measured in boundary waters to date were observed during O17, from coastal to AR stations, with fCO₂ extremes of >1600 μatm and >2400 μatm measured at the surface and at depth, respectively. Contemporaneous Ω_{arag} values were ~0.35, corresponding to Ω_{calc} values <0.6 and pH_T values <7.3 (Figures S7–S8). While these values are not unprecedented within Puget Sound, they had not been observed previously in these boundary waters or the northern CCE.

Summer and fall cruises 2014–2018 showed most surface fCO₂ observations <600 μatm throughout MB, SS, and AR, with spatially limited surface areas below atmospheric saturation levels (~400 μatm CO₂), and most of the water column having >800 μatm fCO₂ by Septembers (Figure S5). While the lowest Ω_{arag} values typically co-occur with hypoxia, Ω_{arag} values fell

below the thermodynamic threshold (Ω_{arag} =1) most times and places in Puget Sound, whereas the occurrence of hypoxic conditions was typically limited to a few months per year in southern HC (Figures 8 and S6, Table 3). In MB, surface conditions during S14 were similar to conditions in Au08, whereas during S15 and S16, surface Ω_{arag} conditions were somewhat lower, but bottom waters had fewer low Ω_{arag} measurements. In contrast, MB surface waters had larger volumes of low-CO₂ water than usual in A17 and J17, corresponding to the particularly high O₂ observations and reflecting a protracted MB bloom (PSEMP Marine Waters Workgroup, 2018). These lower fCO₂ conditions were followed by unprecedented high fCO₂ and low Ω_{arag} observations throughout the water column in MB during S17 (Figures 7–8 and S5–S6, Table 2).

825 HC fCO $_2$ and Ω_{arag} temporal dynamics roughly followed the trajectory described for O $_2$. The most harmful pre-MHW conditions were observed during S09 and O11, with comparable extreme values during S14 (Figures S5-S7). More severe and widespread fCO_2 and Ω_{arag} conditions occurred during J15 and to a lesser extent J16, with fCO_2 values up to 3460 μ atm, Ω_{arag} as low as 0.23 (Ω_{calc}=0.36), pH down to 7.13, and O₂ reaching 12 μmol kg⁻¹ (Table 2). High fCO₂ values in S17 were similarly severe but less widespread than in J15, and S18 conditions were as widespread but less severe than S09, which had the most widespread pre-MHW high fCO2 values. The 12 fCO2 observations higher than the pre-MHW record came from the J15 and S17 cruises. Relationships between Ω values and O_2 content in HC ($R^2 > 0.7$) indicate that aragonite and calcite saturation thresholds (i.e., Ω_{arag}=1 and Ω_{calc}=1) occur at O₂ levels of 225 μmol kg⁻¹ and 145 μmol kg⁻¹, respectively, indicating that even calcite undersaturation occurs more frequently and is more widespread than hypoxia within the southern Salish Sea. The early, extreme J15 fCO2 and Ω_{arag} conditions in HC were likely caused by heatwave-driven subsurface respiration. In contrast, the 835 elevated S17 conditions appear to reflect spring-summer runoff anomalies that strengthened stratification and early seasonal blooms, which in turn would have enhanced deep respiration. Notably, the lowest minima fCO2 across the global surface CO2 observing network have been observed in HC during early spring and highest maxima in fall-winter (Alin et al. in PSEMP Marine Waters Workgroup, 2021; cf. Sutton et al., 2018).

5.2.3 Implications of hydrological anomalies for future biogeochemistry in the southern Salish Sea

While the most severe biogeochemical conditions observed in the Salish Sea through 2018 occurred in J15 and J16, coincident with the maximum expression of the MHW in Puget Sound, biogeochemical response to higher temperatures was not simple in driving the observed low O₂, low Ω_{arag}, high fCO₂ conditions. Rather, indirect effects of increased temperature on biogeochemistry via effects of regional atmospheric warming and precipitation anomalies on watershed hydrology (e.g., river discharge volume and timing) and circulation (stratification and deep-water renewal timing) proved to be as important in shaping the observed biogeochemical anomalies. Prior results have shown that drought as well as enhanced runoff can exert strong influence on circulation and oceanographic conditions in the Salish Sea (Newton et al., 2003 and references in Table 1). The combination of increased atmospheric heat during most of 2013–2018, combined with record-setting precipitation anomalies and earlier snowmelt in the region, caused earlier and higher river discharges that appear to have enhanced estuarine circulation earlier in the year, resulting in bottom water renewal earlier in 2015 and 2016 than typical. In this case, while HC

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bottom waters were more hypoxic, corrosive (low Ω), and high in CO₂ in J15 and J16, conditions in the deep basin had improved by S15 and S16 as a result of these early flushing events.

An independent example of the importance of hydrology to regional biogeochemical conditions came in 2017, the sole year with normal annual air temperature between 2013 and 2018. High river discharge in spring delivered nutrient and stratification conditions suitable for unusually protracted phytoplankton blooms MB and left a fresher salinity signature that persisted until September (Table 2). The blooms led to high surface O₂ and Ω_{arag} and low fCO₂ conditions in MB during spring–summer 2017, but were translated to unprecedented high fCO₂, low Ω_{arag} conditions through the water column by S17 as a result of the subsequent respiration of the high spring–summer phytoplankton biomass. However, while fCO₂ and Ω_{arag} experienced striking anomalies in 2017 compared to 2016 or 2018, deep MB O₂ levels in S17 were unexpectedly slightly higher than in S16 or S18, presumably because the water column started the growing season better oxygenated at depth in A17 than during A16 or A18 (Figure 9 in Alin et al., 2023a). HC also experienced some of its most extreme fCO₂ and Ω_{arag} conditions during S17, indicating that runoff-enhanced biological processes were likely stimulated across Puget Sound basins, however, the conditions were less obviously anomalous in HC where hypoxia and extremely acidified conditions are known to recur. This interpretation is consistent with numerical simulation results indicating that freshwater inflow and estuarine exchange anomalies exerted a stronger influence on the biomass of primary producers in the Salish Sea than increased heat associated with the 2014–2016 heat anomaly (Khangaonkar et al., 2021).

The strongest biogeochemical anomaly seen in the boundary waters was the high fCO₂ and low Ω_{amg} values observed throughout the SJdF water column the following month (O17). Comparing observations across October cruises indicates that O₂ minima were also higher in O17 than O16 or O18 in SJdF, while O17 fCO₂ maxima and Ω_{amg} minima were substantially higher and lower, respectively, than observed in O16 or O18. In fact, oxygen concentrations at a southern HC location known for hypoxia were described as "the least hypoxic on record over the last several years" (Ruef et al. in PSEMP Marine Waters Workgroup, 2018). Extreme O17 acidified conditions in SJdF thus showed the same pattern as in MB of having not particularly low₆O₂ levels compared to either O16 or O18, simultaneous with fCO₂ and Ω_{amg} values completely outside the range of previous observations (Table 2). Widespread low O₂ and Ω_{amg} conditions are known to have occurred during summer 2017 on the Washington shelf (Olympic Coast National Marine Sanctuary and S.R. Alin, unpublished data). The driver of these shelf conditions could have been higher coastal productivity and respiration, although no pronounced coastal anomalies in temperature, salinity, or upwelling were observed during Salish cruises earlier in 2017 to support this interpretation (Figures 2–4, and S4, Table 1). However, lower coastal salinities at depth suggest a potential role for anomalous river input in coastal hypoxia in 2017 as well (Mickett and Newton in PSEMP Marine Waters Workgroup, 2018). It remains unclear whether the strongly acidified water mass in SJdF in O17 was the same acidified water mass observed in MB and SS during S17, subsequently circulated out of PS via estuarine circulation, or the acidified shelf water mass observed in unpublished data.

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Either way, the acidified conditions were likely distributed throughout the SJdF water column by a downwelling wind event that occurred during or just before the O17 cruise.

Because 2017 revealed an anomaly during which O_2 and CO_2 dynamics were decoupled, manifesting as strong acidification not accompanied by particularly low oxygen or marine heatwave conditions, we dubbed this novel biogeochemical anomaly a " CO_2 storm," because it was characterized by particularly high fCO_2 , and accompanied by low Ω_{grow} , Ω_{gale} , and pH_I. The "storm" terminology follows from the description of "carbonate weather" by Waldbusser and Salisbury (2014). Notably, we observed these CO_2 storm conditions in September 2017 in Puget Sound's MB (Figures 7B, 8B, S5–S8) and October 2017 in SJdF (Figures 7A, 8A, and S4), which are locations not previously identified as acidification and hypoxia hotspots. With the higher level of background acidification in HC, what appears to be a CO_2 storm in other basins may manifest as less anomalous carbonate weather there, much as a rainstorm appears less unexpected in a rainforest than a desert. The background acidification and hypoxia gradient across the region—low in MB, moderate in SJdF, and high in HC—thus afforded us the opportunity to observe novel biogeochemical responses to environmental anomalies.

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2021; Hunt et al., 2023).

Collectively, the environmental anomalies of 2013–2018 yielded distinct types of biogeochemical anomalies in the Salish Sea. Marine heat anomalies may drive coupled O₂ and CO₂ system anomalies, whereas terrestrial runoff anomalies driven by atmospheric heat or precipitation anomalies can lead to decoupled O₂ and CO₂ anomalies, as shown here. To understand how future CO₂ storms may affect estuarine and coastal organisms and ecosystems, it is critical to have coupled O₂ and CO₂ system observations because a proxy approach to estimating carbonate chemistry from O₂ and physical parameters will not accurately predict CO₂ storm conditions (e.g., Juranek et al., 2009; Alin et al., 2012). For organisms with sensitivity to high CO₂ or low Ω or pH conditions, direct observation of extreme CO₂ events like those described here would be the only way to know that these anomalous conditions had occurred, which may in turn provide insight for interpreting observed ecological changes (e.g., changes in abundance of sensitive species or onset of a marine disease outbreak). Further, maintaining and enhancing these coupled observations will improve the ability of coupled three-dimensional physical-biogeochemical ocean simulations, such as Regional Ocean Modeling System (ROMS) models, to better differentiate and attribute the roles of and complex interactions among atmospheric, terrestrial, and marine processes in influencing estuarine and coastal acidification (Khangaonkar et al.,

6 Biological importance of understanding changing conditions in the Salish Sea and its boundary waters

915 The iconic marine biota of the Salish Sea—Pacific salmon, Dungeness crab, shellfish—are important resources supporting cultural well-being, livelihoods, and food security for Pacific Northwest communities. Many regional species are vulnerable to direct effects of hypoxic and acidified conditions that naturally occur in this region but are worsening due to climate change. Pink and coho salmon experience changes in their response to olfactory signals that may impair appropriate predator-avoidance

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behaviour during freshwater and early ocean life phases under elevated CO₂ conditions that are currently found in some Salish Sea marine environments (Ou et al., 2015; Williams et al., 2019). Chinook salmon are sensitive to warming temperatures and hypoxia (Crozier et al., 2019, 2021) and may be sensitive to direct ocean acidification impacts as well.

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Dungeness crab are the U.S. West Coast's most economically important marine species (e.g., Alin et al., 2015) and an important recreational and tribal fishery in Puget Sound (e.g., Froehlich et al., 2017). Recent closures of the Dungeness crab fishery in HC and SS have limited regional access to this marine resource (e.g., Washington Department of Fish and Wildlife, 2020). Field studies show that early life stages experience sublethal dissolution damage to carapaces and mechanoreceptors with sensory and behavioural functions under current conditions (Bednaršek et al., 2020b). Further, Dungeness crab are sensitive to increased temperature and declining oxygen and pH across life stages, with population-level vulnerability to projected warming, hypoxia, and acidification levels from surface to benthic habitats by the year 2100 (Hodgson et al., 2016; Berger et al., 2021). Pacific oysters, another regionally important commercial shellfish species, show a high proportion of defects in larval shell development (Waldbusser et al., 2015) in response to aragonite saturation levels currently present at most times, depths, and places in Puget Sound (Figure 8, Table 3).

Warming, hypoxia, and acidification also affect the prey and predators for these important species via trophic linkages. Euphausiids (krill) are a dominant food source for finfish and seabirds in the CCE whose larval development and survival are impaired under current pH conditions (McLaskey et al., 2016). Pteropods are another abundant prey source (Bednaršek et al., 2019 and references therein), and calcifying pteropods in the Salish Sea showed severe dissolution effects during 2014–2016 (Bednaršek et al., 2021a). Pteropods experience synergistic effects of high temperature (>11 °C), low oxygen, and low Ω_{arag} or high pCO₂ on their abundance, shell dissolution, and oxidative stress biomarkers (Bednaršek et al., 2018, 2021a; Engström-Öst et al., 2019). Even the regional apex predator, Southern resident orca whales, whose population has been in decline for decades, may be affected if their dominant food source, Chinook salmon, declines further in abundance as a result of increasingly stressful multi-stressor conditions in the region (Hanson et al., 2021).

Many of the oceanographic variables in this observational data package passed thresholds known to be potentially harmful to regionally important species. Below we briefly describe the frequencies at which individual and multiple stressors pass known biological thresholds for salmon, crab, and pteropods in order to provide insight to marine resource managers about present-day ecosystem conditions facing Salish Sea resources. These examples of potential combined biological impacts of ocean acidification, hypoxia, and warming in the Salish Sea illustrate how a complex ecosystem like Puget Sound manifests as a mosaic of environmental stressors occurring at different frequencies through space and time.

6.1 Species' thresholds to and occurrence frequencies of individual stressors

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Thermal barriers to Pacific Northwest salmonid migration are known to occur at temperatures of 19–23 °C (McCullough et al., 2001). During most July 2014–2018 cruises in HC, temperatures ≥19 °C were observed in the upper 10 m of the water column, with several exceeding 21 °C (Table 3). These observations comprise 0.5% and 0.1% of the compiled Salish cruise data product, respectively, and occurred only in HC, although July temperatures also approached stressful levels for salmon in WB. Dungeness crabs have substantial hatch mortality at temperatures >15 °C (Rasmuson, 2013). Temperatures crossed this threshold in 2.8% of observations, across all basins and dominantly in the upper 20 m, where brooding females aggregate (Pauley et al., 1989; Rasmuson, 2013).

Oxygen levels harmful to salmonids can be as high as 3.5–5 mg L⁻¹ (~110–155 μmol kg⁻¹) (e.g., Table 3 in U.S. Environmental Protection Agency, 2003 and references therein). Oxygen levels ≤155 μmol kg⁻¹ comprised 21.7% of Salish cruise observations (Table 3). Dungeness crab experience feeding cessation in adults at 62 μmol kg⁻¹ (Rasmuson, 2013). Oxygen fell below this hypoxia threshold in 1.3% of observations, at depths of 5–335 m on both pre- and post-MHW cruises, with a majority at HC stations and 20% in surface waters. Hypoxic bottom waters in HC have also been shown to drive Dungeness crab to shallower habitat, which may affect their catchability, predation, competition, and cannibalism, and thus potentially future population numbers (Froehlich et al., 2014, 2017). The remaining hypoxic measurements in SJdF and coastal waters (Table 2) do not represent the southern Washington coastal waters where the frequency and severity of hypoxic conditions during the upwelling season is higher (Connolly et al., 2010; Peterson et al., 2013).

As noted previously, aragonite and calcite fall below their saturation thresholds at higher O_2 levels, so the frequencies at which Ω_{arag} and Ω_{calc} undersaturation occur are much higher than hypoxia. Ω_{arag} was undersaturated in 61% of observations and Ω_{calc} 14% (Table 2), including hundreds of surface observations (Table 3). Biological thresholds for severe dissolution or growth in calcifying pteropods ($\Omega_{arag} = 1.15-1.20$; Bednaršek et al., 2019) are higher than the thermodynamic threshold, thus yielding >78% of all Salish cruise observations below this threshold, of which a third were in surface waters. While Ω_{arag} and Ω_{calc} thresholds were crossed in all seasons and basins within our study area, the threshold exceedance frequencies are much higher for calcite in HC than elsewhere, and much higher for aragonite across all basins than for calcite, temperature, or oxygen conditions becoming harmful.

Increases in fCO_2 levels may affect regionally important species through hypercapnia—the metabolic challenge of too much CO_2 rather than too little O_2 . A 1000 μ atm pCO_2 threshold for hypercapnia has been used for a range of fish and invertebrate studies (McNeil and Sasse 2016), although fCO_2 exposure thresholds are not well established for regionally important species. Using the 1000 μ atm threshold, >25% of Salish cruise observations represent conditions potentially conducive to hypercapnia, making them more prevalent than even the 155 μ mol kg⁻¹ oxygen threshold for salmon (cf. U.S. Environmental Protection

Agency, 2003; Vaquer-Sunyer and Duarte, 2008). While a discrete threshold was not identified for these changes, behavioural, neural, and gene expression responses have been observed in ocean-phase Pacific salmon between treatment *p*CO₂ levels of 700 μatm and 2700 μatm (Williams et al., 2019). 65% of Salish cruise CO₂ observations are higher than 700 μatm and 0.4% are higher than 2700 μatm, implying that ocean-phase Pacific salmon may encounter challenging CO₂ levels in the present-day southern Salish Sea.

A synthesis of decapod species' sensitivity to ocean acidification identified pH_T thresholds of 7.80 for egg hatching success, 7.70–7.74 for adult respiration and haemolymph pH, and 7.52 for larval survival (Bednaršek et al., 2021b and references therein). Within the Salish cruise data, 9–64% of pH_T observations crossed these thresholds, with pH conditions below the larval mortality threshold occupying much of the water column in HC during summer–fall (Figure S8, Table 3). Broader pH_T survival threshold estimates for larval, juvenile, and adult decapod life stages span 7.40–7.80 (Bednaršek et al., 2021b); these thresholds were crossed at <10% to >60% frequencies in the Salish cruise data. A multi-stressor vulnerability analysis on Washington and Oregon coastal Dungeness crab populations used a pH_T=7.65 threshold across life stages to assess exposure levels under present and future conditions (Hodgson et al., 2016; Berger et al., 2021); >23% of Salish cruise observations exceeded this threshold. Dungeness thresholds for pH_T sensitivity were crossed with the highest frequency and severity in HC (Figure S8). This chronic exposure in subsurface HC may be sublethal (Berger et al., 2021 and references therein), but the effects of acidified conditions on population distributions and crab catchability during Washington's tribal and state fisheries is currently unknown (cf. Froehlich et al., 2017).

1010 6.2 Co-occurrence or interactions of multiple stressors in the present-day southern Salish Sea

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Populations of valuable Pacific salmon and trout, including some classified as Threatened under the U.S. Endangered Species Act, have native habitat in Puget Sound and its watersheds (NOAA Fisheries, 2022). The combination of high temperature and low oxygen can be particularly disruptive for salmonid migrations. Temperatures ≥ 19 °C never co-occurred in the same sample with $O_2 \leq 155$ µmol kg⁻¹. However, a 21.8 °C temperature occurred at 2.7 m depth with O_2 levels of 99.6 µmol kg⁻¹ and fCO_2 levels of 1437 µatm at 10.9 m at the same HC station during J18, putting three known salmon stressors in close physical proximity, with the risk of combined physiological effects and habitat compression (Table 4). Similar combinations of temperature, oxygen, and fCO_2 conditions were also recorded during J16 and J17 in southern HC. Several salmonid runs enter a nearby river for the metabolically challenging breeding migration during summer, when these conditions formed near the river mouth (Gray, 2022). If heatwave conditions observed during summer 2015 are representative of the future marine stressor landscape, harmful levels of temperature, oxygen, and fCO_2 may be more likely to co-occur earlier in the salmon run season, rather than occurring separately, with peak temperatures in July and minimum O_2 /peak fCO_2 values occurring in September, as was typical during pre-MHW years.

Closer inspection of when and where harmful conditions co-develop in the region reveals that low pHT and O2 co-occurred at 1025 23 stations across 10 cruises, with four cruises and 15 of the stations sampled prior to the onset of strong, recurring atmospheric and surface water temperature anomalies beginning in 2013 (Tables 1 and 4). Of cruises with co-occurring low O2 and pH, 14 stations were in HC and nine in boundary waters, and these conditions predominantly affected the bottom waters that juvenile and adult Dungeness crabs inhabit. In contrast, the combination of stressfully warm surface temperatures with low pH_T near below was observed mostly after the onset of heat and other anomalies and during cruises from July into October. This combination occurred during 11 cruises at 61 stations—occupying a wider geographic distribution across PS and boundary waters—with 10 stations in August 2008 being the only cruise prior to September 2013 showing this combination of stressors (Table 4). While these conditions are unlikely to occur in the present-day during Dungeness hatching season (winter-spring), anomalously low salinity (<15) can also prevent egg hatching, interfere with larval progression, and be lethal for adults (Pauley et al., 1989; Rasmuson, 2013). Thus, nearshore hatching habitats may thus have been affected by the major 2013-2018 precipitation and river runoff anomalies (Table 1), providing another example of how hydrological anomalies may have profound ecosystem effects in PS, beyond effects on carbonate chemistry. We note that low salinity anomalies should also be considered as part of the multiple stressors management landscape.

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A triad of Dungeness crab stressors—high temperature (>15 °C), low O₂ (<62 μmol kg⁻¹), and low pH_T (<7.65) (Berger et al., 2021)—co-occurred six times at stations in southern HC during the J15, J16, and S17 cruises (Table 4). Stressful conditions 1040 for Dungeness spanned much or all of the water column at five stations during J15 and J16 cruises, and occupied the surface and a less extensive part of the subsurface water column during S17. This newly observed co-occurrence of all three crab stressors in Hood Canal during 2015-2017 reveals a possible future path for how the multi-stressor marine environment will evolve in this region under warmer climate conditions. Potential biological ramifications include the disappearance of suitable 1045 habitat for all life stages of Dungeness crab from some areas of the southern Salish Sea during the pelagic larval and settlement seasons. Summer crabbing seasons across 2008-2019 in HC typically spanned July-August (Washington Department of Fish and Wildlife data), making it likely that habitat compression due to combined T, pH, and O2 stress affected crab depth distributions during this time-series and may warrant management consideration. However, parsing whether the combined effects of multiple interacting stressors across crab life stages and habitats has contributed to recent PS Dungeness fisheries closures would require a more complex modelling analysis like those done in coastal waters (Hodgson et al., 2016; Berger et al., 2021).

Table 4. Multiple stressor events relevant to Dungeness crab, and their temporal and spatial occurrence in the Salish cruise data package. Thresholds to determine event occurrence were those used by Berger et al. (2021). Depth ranges affected by combinations of hypoxic (oxygen content <62 μmol kg⁻¹), low pH_T (pH_T <7.65), or high temperature (>15 °C) conditions are indicated in the "Depth range" columns. Cruises that occurred before the beginning of major atmospheric and surface seawater temperature anomalies in the summer of 2013 are in italics.

Stressor	Number of stations ^a				Depth range (dbar)b	
Cruise	Boundary	South	Whidbey	Hood	Boundary	Puget
	waters	Sound	Basin	Canal	waters	Sound
pH _T , O ₂						
Aug. 2008	7	_	_	1	50-335	85-90
Sep. 2009	_	_	_	1	_	10-45
Nov. 2010	_	_	_	2	_	$5-15^{c}$
Oct. 2011	_	_	_	4	_	$5-30^{\circ}$
Sep. 2013	1	_	_		120-330 ^d	_
Oct. 2014	_	_	_	1	_	10-20
Apr. 2015	_	_	_	2	_	120-150 ^d
Sep. 2016	_	_	_	1	_	10-15
Oct. 2017	1	_	_	_	90-100	_
Sep. 2018	_	_	_	2	_	5–55
T, pH_T						
Aug. 2008	_	_	4	6	_	5–175
Sep. 2013	1	_	_		$120-290^{d}$	_
Jul. 2014	_	_	2	9	_	20-175e
Sep. 2014	_	_	_	1	_	20-50
Oct. 2014	2	_	_		$250-325^{d}$	_
Jul. 2015	_	_	2	3	_	10-170
Sep. 2015	_	_	2	1	_	5-145
Jul. 2016	_	_	_	4	_	10-165
Jul. 2017	_	_	_	7	_	20-170
Sep. 2017	_	1	1	5	_	$5-170^{\rm f}$
Jul. 2018	_	1	3	6	_	$10-165^{\rm f}$
T, pH_T, O_2						
Jul. 2015	_	_	_	2	_	20-120
Jul. 2016	_	_	_	3	_	20-115
Sep. 2017	_	_	_	1	_	50-80
Multiple stressors never occurred at a single station simultaneously in Main Basin or Admiralty Reach, so the						

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^a Multiple stressors never occurred at a single station simultaneously in Main Basin or Admiralty Reach, so they are not included in this table. ^b Depth ranges affected by pH_T or O₂ threshold exceedance are rounded to most inclusive 5 dbar intervals, although sampling resolution was sparse enough that the depth ranges affected by low pH_T or O₂ conditions may have been larger for any given cruise. ^c The depths at which hypoxic and low pH conditions were seen on these cruises were near-surface, not near-bottom; bottom conditions were less harmful. ^d Depth ranges affected were deeper than habitats where Dungeness crab are typically most abundant on the Washington coast (e.g., Berger et al., 2021). ^c The largest number of stations affected by two independent stressors were observed during this cruise. ^f Combined temperature and pH_T stressor conditions were the most widespread across basins during these cruises.

7 Conclusions: What we have learned about multi-stressor dynamics from the Salish cruise time-series so far

Ocean acidification variables indicate stressful conditions throughout the region, though with substantial regional variation. Aragonite undersaturation was pervasive throughout the region, and even calcite undersaturation occurred more widely and frequently than hypoxia. fCO₂ levels were above atmospheric values in >90% of observations, often by hundreds of μ atms. Hypoxia was observed most frequently in southern Hood Canal and occasionally in boundary waters during 2008–2018. This seasonally resolved cruise time-series co-occurred with several major environmental anomalies, giving us the opportunity to observe the impacts of atmospheric and marine heatwaves, precipitation, and river discharge anomalies on Salish Sea physical and biogeochemical conditions. The strongest heat anomalies manifested earlier in boundary water cruises (October 2014) but in a delayed and more protracted way within Puget Sound, spanning at least 2015–2016. Increased temperatures throughout the water column likely contributed to the most extreme O₂, fCO₂, and Ω _{arag} conditions seen in southern Hood Canal during July 2015 and July 2016, developing earlier in the season than the typical late-summer timing for previous hypoxia events. However, anomalously early deep-water renewal events in Hood Canal in 2015 and 2016 resulted in late-summer conditions that resembled a less hypoxic, pre-heatwave year in deep southern Hood Canal. The effects of the heat anomaly thus underscore the importance of physical oceanographic conditions (i.e., temperature, stratification, and circulation timing) in setting the stage for the severity and duration of ocean acidification and hypoxia in local hotspots.

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Both seasonal and spatial variation was strong. The decade-long Salish cruise time-series illuminated several differences in the seasonality across parameters and across Puget Sound and its boundary waters. Some parameters peaked at the surface during summer (temperature, Ω_{arag}), with others rising (salinity, fCO₂) or falling (O₂) from spring to fall, while all parameters showed more monotonic seasonal progression at depth. Within Puget Sound, the largest overall variability was observed in the most stratified basins. Notably, deep-water seasonality in Hood Canal was different from other basins in that seasonal O₂ lows and highs co-occurred with temperature and salinity minima and maxima, whereas in other basins, salinity and temperature were inversely correlated with O₂ content. The lags likely result from the combination of more complex deep-water renewal and earlier spring blooms in Hood Canal, which appear to decouple its physical and biogeochemical seasonality relative to other deep <u>Puget Sound</u> basins and may predispose it to more hypoxic, acidified conditions. To the extent that climate change alters the coupling between physical and biogeochemical seasonality in other regional waters, changes in the frequency, severity, and duration of ocean acidification and hypoxia may be expected.

During fall 2017, we observed a novel carbonate system anomaly in Puget Sound and boundary waters. This " CO_2 storm" was characterized by unprecedented high fCO_2 and low Ω_{arag} values, which crossed sensitivity thresholds for regionally important species. This extreme event occurred independently of heatwave or particularly low oxygen conditions and instead appears to have resulted from major river discharge anomalies reflected by low salinity anomalies observed earlier in the year. The CO_2 storm was most obvious in basins with less acidified baseline conditions, but was still detectable in the most acidified basin.

Hood Canal. These observations showed decoupling of carbonate chemistry from that of oxygen, which was unusual in this time series. This result underscores the need for on-going seasonal monitoring of both carbon and oxygen across Salish Sea basins, which showed variable responses to anomalies in physical ocean conditions, river input, and local weather. The Salish cruise time-series has thus illustrated how the arc of major environmental anomalies and their ecological impacts depends on the biogeochemical metric, species of interest, and baseline conditions of the basins in which they occur.

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The frequencies at which Salish cruise observations crossed known or preliminary species sensitivity thresholds illustrates the relative risk landscape of temperature, hypoxia, and acidification anomalies in the southern Salish Sea in the present-day. Interactions between marine heat and other environmental anomalies during the 2014–2018 cruises revealed how multiple stressors can combine to present potential migration, survival, or physiological challenges to key regional species. Collectively, the occurrence frequencies of these combinations of stressors for Dungeness crab, salmon, and pteropods in the Salish cruise data package illustrate how increasingly frequent and severe marine and atmospheric heatwaves may alter the future co-occurrence of multiple stressors in the southern Salish Sea. Specifically, while low pH_T and O₂ co-occurred regularly before the heat anomalies in mostly subsurface Hood Canal and boundary waters, the co-occurrence of high surface temperature and low pH conditions appeared to increase sharply across the marine heatwave, with a broader spatial distribution than where low pH and O₂ previously co-occurred. Future changes in the seasonality of when harmful conditions develop and co-occur may cause sensitive species' thresholds to be crossed more frequently due to changes in timing even without changes in severity. The novel forms and combinations of extreme events observed in the 2014–2018 Salish cruise time-series have thus provided insight into the potential evolution of the future marine stressor risk landscape in the Salish Sea.

In summary, observing assets deployed throughout coastal and estuarine environments in this region allowed us to detect unexpected biogeochemical anomalies in this region, including decoupling between hypoxia and acidification conditions, that would not have been discovered without high-quality measurements of both oxygen and carbonate system variables. The ability of 3D ocean simulations to accurately project complex ecosystem multistressor events and their impacts on marine species will benefit from comparison with rich observational data sets such as this cruise time-series, which is capable of resolving seasonal events throughout the water column and diverse basins of the southern Salish Sea. Further synthesis of the rich information from this data product alongside that from the moored time-series in this region, with its relatively high temporal resolution (sub-daily), may facilitate increased understanding of mechanisms driving variation, which is needed to further develop recommendations for multistressor monitoring approaches and analyses.

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$\boldsymbol{8}$ Data availability and data use

1130 This analysis is focused on a subset of the Salish cruise data product archived in Alin et al. (2022, https://doi.org/10.25921/zgk5-ep63) and described in Alin et al. (2023a). To facilitate use of the combined multi-stressor data

product at the core of this paper, we created a novel data product that includes only the highest-quality measured parameters, along with the calculated CO_2 system parameters pH_T , fCO_2 , pCO_2 , Ω_{arag} , and Ω_{cale} , which were calculated using two sets of dissociation constants as described in Section 3.2; this multi-stressor data product is archived at Alin et al. (2023b, https://doi.org/10.25921/5g29-q841). The data (filename= "SalishCruises_2008to2018_MeasCalcParams_NCEIdataProduct_09262023.csv") and metadata (filename= "SalishCruise_2008to2018_MeasCalcParams_metadata_09262023.xlsx") can be accessed by clicking the "Download Data" button and downloading the files from the Index page.

Author contributions

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SA led analysis of inorganic carbon samples, assembly and analysis of data and metadata, interpretation of data analyses, and manuscript drafting. JN led organization and execution of all cruises, oxygen and nutrient measurements, and provided input on data analysis and interpretations at all stages of the work. RF contributed to the development and implementation of this project and the writing of the manuscript. SS was our research partner on many projects over the duration of these cruises and contributed to understanding the physical and biogeochemical dynamics in the region. DG created the map and transect profile graphics and made major contributions to data wrangling. All authors contributed to editing the manuscript.

50 Competing interests. The authors declare they have no competing interests.

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Canal salmon species migration timing. We thank two anonymous reviewers for constructive comments that improved the clarity of the article for a broader audience. This is PMEL contribution number 5297.

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