Portland State University

PDXScholar

Environmental Science and Management Faculty Publications and Presentations

Environmental Science and Management

11-12-2019

Microplastic Concentrations in Two Oregon Bivalve Species: Spatial, Temporal, and Species Variability

Britta Baechler Portland State University, baechler@pdx.edu

Elise F. Granek Portland State University, graneke@pdx.edu

Matthew V. Hunter Oregon Department of Fish and Wildlife

Kathleen E. Conn United States Geological Survey

Follow this and additional works at: https://pdxscholar.library.pdx.edu/esm_fac

Part of the Environmental Health and Protection Commons, Environmental Indicators and Impact Assessment Commons, and the Environmental Monitoring Commons Let us know how access to this document benefits you.

Citation Details

Baechler, Granek, E. F., Hunter, V. M., & Conn, K. E. (2019). Microplastic Concentrations in Two Oregon Bivalve Species: Spatial, Temporal, and Species Variability. Limnology and Oceanography Letters.

This Article is brought to you for free and open access. It has been accepted for inclusion in Environmental Science and Management Faculty Publications and Presentations by an authorized administrator of PDXScholar. Please contact us if we can make this document more accessible: pdxscholar@pdx.edu.



©pen Access Limnology and Oceanography Letters 2019 © 2019 The Authors. Limnology and Oceanography published by Wiley Periodicals, Inc. on behalf of Association for the Sciences of Limnology and Oceanography. doi: 10.1002/lol2.10124

SPECIAL ISSUE-LETTER

Microplastic concentrations in two Oregon bivalve species: Spatial, temporal, and species variability

Britta R. Baechler¹, ¹* Elise F. Granek, ¹ Matthew V. Hunter, ² Kathleen E. Conn³

¹Department of Environmental Science and Management, Portland State University, Portland, Oregon; ²Oregon Department of Fish and Wildlife, Astoria, Oregon; ³United States Geological Survey, Tacoma, Washington

Scientific Significance Statement

Plastics have innumerable uses and are inextricably tied to daily life in modern society. These plastics begin as or break down into microplastics, which are now found in an array of terrestrial, aquatic, and marine habitats and organisms. These tiny particles may threaten ecosystem balance and natural resource consumers, particularly in the case of seafood. In Oregon, U.S.A., Pacific oysters (*Crassostrea gigas*) and razor clams (*Siliqua patula*) are of commercial, recreational, and cultural importance, yet baseline information on microplastic prevalence in these species and across sites and seasons is absent. Our study is the first to document microplastics in Pacific razor clams and provides important coast-wide data to compare microplastic burden across species, seasons, and sites.

Abstract

Microplastics are an ecological stressor with implications for ecosystem and human health when present in seafood. We quantified microplastic types, concentrations, anatomical burdens, geographic distribution, and temporal differences in Pacific oysters (*Crassostrea gigas*) and Pacific razor clams (*Siliqua patula*) from 15 Oregon coast, U.S.A. sites. Microplastics were present in organisms from all sites. On average, whole oysters and razor clams contained 10.95 ± 0.77 and 8.84 ± 0.45 microplastic pieces per individual, or 0.35 ± 0.04 pieces g⁻¹ tissue and 0.16 ± 0.02 pieces g⁻¹ tissue, respectively. Contamination was quantified but not subtracted. Over 99% of microplastics were fibers. Material type was determined using Fourier-transform infrared spectroscopy. Spring samples contained more microplastics than summer samples in oysters but not razor clams. Our study is the first to document microplastics in Pacific razor clams and provides important coast-wide data to compare microplastic burden across species, seasons, and sites.

Data Availability Statement: Data are available in the PDXScholar data repository at https://doi.org/10.15760/esm-data.1.

Associate editor: Erika Holland

Additional Supporting Information may be found in the online version of this article.

This is an open access article under the terms of the Creative Commons Attribution License, which permits use, distribution and reproduction in any medium, provided the original work is properly cited.

This article is an invited paper to the Special Issue: Microplastics in marine and freshwater organisms: Presence and potential effects Edited by: Dr Elise Granek, Portland State University, Dr Susanne Brander, Oregon State University, and Dr Erika Holland, California State University, Long Beach

^{*}Correspondence: baechler@pdx.edu

Author Contribution Statement: B.R.B. and E.F.G. identified the research question and study approach. M.V.H., E.F.G., and B.R.B. designed the field survey; M.V.H. and B.R.B. conducted the field survey. K.E.C., E.F.G., and B.R.B. designed laboratory methods; B.R.B. conducted the laboratory study. B.R.B. led the manuscript effort; E.F.G., M.V.H., and K.E.C. contributed ideas and edits to the manuscript. B.R.B. and E.F.G. are responsible for manuscript data.

Microplastics, plastics 0.0001–5 mm in any linear direction (UNEP 2016), are found in nearly every environment on earth (Thompson et al. 2004). These tiny fragments, pellets, filaments, and fibers originate from both marine and land-based sources, infiltrating aquatic ecosystems worldwide through pollution, runoff, wastewater, and atmospheric deposition (Zhang 2017). Globally, the overwhelming number of single-use and nondegradable plastic items has led to widespread microplastic pollution. Plastics are manufactured to be durable, so degradation can take hundreds to thousands of years, posing a pervasive and severe problem for ecosystems as well as a human health concern (Cole et al. 2011; Wang et al. 2016).

While spatial distribution of microplastics in the environment is highly complex, areas with high human population, coastal recreation, and tourism pressures generally yield high environmental microplastics (Barnes et al. 2009; Hantoro et al. 2019). Microplastics represent a diverse set of contaminants which encompass infinite combinations of plastic densities, sizes, shapes, surface textures, and chemical properties (Rochman et al. 2019). Once transmitted into the environment, microplastics are subjected to an array of dynamic hydrological, biological, and atmospheric processes, including surface currents, tides, biofouling, mechanical and ultraviolet degradation, precipitation, storm events, and more. While human presence may correlate with microplastic prevalence, it is unclear what specific environmental processes best predict fate and transport of these pernicious particles (Jambeck et al. 2015; Zhang 2017). Density has been thought to ultimately determine environmental fate, with denser plastics like polyvinyl chloride and polyethylene terephthalate (PET) settling to the benthos and low density polymers such as polystyrene, polypropylene, and polyethylene remaining in surface waters; however, a recent review of global surface water and sediment data indicates a mixture of high and low density microplastics in water and sediment samples, attributed to influences of varied environmental and biological processes in coastal areas (Hantoro et al. 2019).

Rivers have been well established as vectors of plastics into coastal and marine environments (Zhang 2017). These dynamic waterways transport between 1.2 and 2.4 million tons of plastic into global oceans each year, with up to 28.8 thousand tons transmitted annually through North and Central American rivers alone (Lebreton et al. 2017). A study investigating microplastic concentrations in surface waters from two Los Angeles, California rivers quantified an input of roughly 2 billion microplastic pieces into coastal waters in the span of just 3 d (Moore et al. 2005). Expanded to an annual output, these two rivers transport over 240 billion microplastics per year to the California coast. Stormwater runoff and wastewater treatment plant (WWTP) effluent also contribute significant microplastic burdens to coastal environments (e.g., Carr et al. 2016; Napper and Thompson 2016; Mintenig et al. 2017). Microfibers, generally broken down from laundered

clothing items or from derelict fishing gear, are the most prevalent form of microplastic in the nearshore environment (Barrows et al. 2018; de Falco et al. 2019).

Organisms inhabiting coastal environments are subjected to ambient environmental conditions, including microplastic contamination that may exist in surrounding waters, substrata, or in the air. Aquatic filter- and suspension-feeding organisms can encounter microplastics in the marine or freshwater column, mistake them for food items, and ingest them (Hantoro et al. 2019). This transfer of plastics from the environment into aquatic food webs has been documented across diverse taxonomic groups, life histories, habitats, and feeding types (e.g., Cole et al. 2011; Akpan 2014; Rochman et al. 2015; Waite et al. 2018). After uptake, microplastics can adhere to organs or become incorporated into guts, gills, and tissues of organisms, decreasing energy uptake and impairing muscle function and reproduction (e.g., von Moos et al. 2012; Sussarellu et al. 2016; Ribeiro et al. 2017; Kolandhasamy et al. 2018). Microplastics may also sorb harmful contaminants that, once ingested or incorporated in tissues, are released into the organism (Teuten et al. 2007, 2009). In some studies, environmental microplastic concentrations have been directly correlated with microplastic burdens in coastal bivalves (Mathalon and Hill 2014; Li et al. 2016; Qu et al. 2018; Hantoro et al. 2019).

As filter feeders, bivalves are particularly vulnerable to contaminants in the estuarine and open coast environments they inhabit. The Pacific Northwest (PNW) region of North America supports an array of filter-feeding shellfish species, which have been inextricably tied to the natural history and cultural heritage of the area for millennia. Fishery and aquaculture sectors serve as important anchors of the region, with Pacific oysters and razor clams playing particularly significant roles in food security and the economy (Crossett et al. 2013). Pacific oysters (Crassostrea gigas) have been commercially farmed in the PNW since introduction of the species in the early 1900s (Glude and Chew 1982). These filter feeders consume particulates in the water column, such as plankton and other organic material, and reach commercial size (100-150 mm, maximum length 250 mm) over 2-4 years (Pauley et al. 1988; Harris 2008). Pacific razor clams (Siliqua patula) are native to the PNW and are found on intertidal beaches. They have been harvested by first nations and tribal peoples for centuries, and in statemanaged recreational and commercial fisheries since the 1950s. Consuming phytoplankton, razor clams grow rapidly in the first year attaining lengths up to 90 mm, and a maximum length of 16 cm over their 6-yr lifespan (Link 2000).

Microplastic concentrations in field-collected Pacific oysters have been documented worldwide facilitating comparisons between samples grown in Oregon vs. other regions; however, there is no published literature on microplastic prevalence or effects in Pacific razor clams. We initiated this study to answer the question: *What variables predict microplastic concentrations in Oregon Pacific oysters and Pacific razor clams?*

Flowing between the U.S.A. states of Washington and Oregon is the Columbia River, the largest river on the North American continent with a Pacific Ocean terminus. We predicted the Columbia would be a major vector of microplastics, causing elevated burdens in our study species at the northernmost study sites and attenuated burdens with increased distance from the river. Coastal tourism is highest during the summer months (May-October). Tourism results in increased use of beaches and waterways for recreation and an uptick in laundering needs, so we hypothesized that concentration of environmental microplastics in waters and coastal organisms would be higher in summer than spring. We hypothesized that gut tissue would contain more microplastics than nongut tissue due to retention of microplastics in the gut of bivalves observed in previous studies (e.g., Browne et al. 2008; Ward and Kach 2009; Sussarellu et al. 2016; Woods et al. 2018). Because microplastics may become lodged in gills or other organs (Woods et al. 2018), we predicted a positive relationship between organism size and microplastic burden-that larger individuals would contain more microplastics than smaller individuals. We examined these expectations through field-collection of Pacific razor clams and purchase of Pacific oysters at 15 locations, during two seasons, taking biological measurements and investigating whole, gut-tissue, and tissue-only samples.

Methods

Field sites, sample collection, processing, and microplastic enumeration

A total of 141 Pacific oysters and 142 Pacific razor clams were collected from 15 sites during low tides in spring (27–28 April 2017) and summer (21–31 July 2017) (Table 1). Whole oysters were purchased from growers at six sites during both seasons. One oyster grower was selected from each of six Pacific oyster-producing bays. In this report, oyster grower names are withheld and are coded randomly as OY1–OY6. Oyster shell length averaged 125.39 mm (range = 77.67–197.66 mm) and wet tissue weight averaged 30.97 g (range = 8.51-101.67 g; Supporting Information Appendix 1).

Razor clams were collected from nine sandy beach sites stretching from Clatsop in the north, to Gold beach, near the California border, in the south (Fig. 1). Of the nine clam sites, four were sampled in both spring and summer, providing a temporal snapshot of microplastic frequencies. Collection was performed in coordination with Oregon Department of Fish and Wildlife (ODFW) and Oregon Department of Agriculture (ODA), which greatly augmented efforts to achieve desired sample size. Clam sites were selected based on ODFW knowledge of existing clam populations, feasibility of sample collection (access, tides, clam shows), and with a goal of sampling a large swath of the coast. Summer clam sampling was more robust than spring because it corresponded with a coast-wide ODFW survey and coincided with lower tides than spring. Razor clam shell length averaged 113.89 mm (range = 56.00–132.52 mm) and wet tissue weight averaged 55.71 g (range = 5.84–92.11 g; Supporting Information Appendix 1).

All samples were transported on ice to the Applied Coastal Ecology laboratory at Portland State University (PSU) in Portland, Oregon, in clean 2-L glass Mason jars. Shell and tissue measurements were collected with a digital Mitutoyo caliper and Ohaus balance accurate to 0.01 mm and 0.01 g, respectively. Bivalve shells were rinsed with deionized (DI) water to remove sand, mud, and debris, were shucked into clean 120 mL Mason jars and frozen at -20° C.

Samples were thawed and digested for 24 h in a laminar flow fume hood using 10% potassium hydroxide (KOH). Digestion began with the first organism from each site and season, then proceeded to the second organism from each site, until all samples were processed. Samples were poured through a 7.6 cm diameter, 63 µm stainless steel sieve. Material retained on the sieve was rinsed into clean, labeled glass petri dishes. Petri dishes with Petristickers® affixed to the bases were placed in a drying oven at 40°C for 24 h and stored in sealed tubs prior to microscope processing. Due to high levels of organic material and sand granules remaining in clam samples after initial digestion, a second 10% KOH digestion combined with hypersaline density separation (330 g L^{-1} Fisher Chemical Certified ACS Crystalline NaCl) was utilized. Samples were analyzed under a Leica M165C stereomicroscope (×10–120 magnification) connected via a Leica IC80 HD camera to a computer running Leica Application Suite X imaging software. Each suspected microplastic encountered was measured and particle category (fiber, fragment, film, foam, bead, unknown), color, and maximum length were recorded. To determine material type for microplastics, a subset of identified fibers was randomly selected using random number generation to determine: (1) sample dish, then (2) segment of each dish (segment numbers 1-16) from which to extract 26 suspected microplastics. The first fiber visually encountered in the randomly generated dish and segment was selected for validation. Fibers were analyzed using a Thermo Nicolet iS10 Fourier-transform infrared spectrometer (FTIR) equipped with an Attenuated Total Reflectance accessory at the University of New Hampshire Instrumentation Center. Spectra for each microfiber were acquired using 256-1024 scans depending on size and width. Automatic software comparison of microfiber spectra to a set of Thermo Nicolet Omnic™ FTIR spectral libraries was used to generate a best match.

Gut/tissue separation

During summer sampling, three individual organisms from each site (with the exceptions of Bastendorff Beach and Coos Bay) underwent a separation of digestive organs from other tissues. For Pacific oysters, gut-tissue samples included the visceral mass, esophagus, diverticular gland, midgut, and stomach. In razor clams, gut-tissue samples included the stomach, small intestine, and crystalline style. All remaining tissue was

				Sprin	ing				Summer	ner		Bot	Both seasons	S
		# samp	# samples analyzed	۱yzed	Microplastic burden	ic burden	# sampl	samples analyzed	yzed	Microplast	Microplastic burden			
		Whole	Gut	Nongut	Avg # MP per sample	Avg # MP g ⁻¹ tissue	Whole	Gut	Nongut	Avg # MP per sample	Avg # MP g ⁻¹ tissue	Avg MP length in	Min MP Ienath	Max MP Iength
Species	s Site	organisms tissue	s tissue	tissue	(SE) ¹	(SE) ²	organisms	tissue	tissue	(SE) ¹	(SE) ²	mm (SE) ³		(mm) ⁵
Pacific	OYI	10	0	0	13.60 (2.60)	0.55 (0.34)	10	e	°.	9.6 (2.56)	0.49 (0.17)		0.16	5.37
oyste	oyster OY2	12	0	0	10.33 (1.92)	0.35 (0.29)	11	m	°	6.81 (1.58)	0.21 (0.05)	1.32 (0.06)	0.18	5.85
	OY3	10	0	0	14.60 (3.53)	0.62 (0.49)	10	7	3	8.5 (2.13)	0.28 (0.07)	1.24 (0.07)	0.12	6.08
	OY4	10	0	0	17.50 (3.85)	0.39 (0.28)	10	m	3	5.20 (1.54)	0.10 (0.02)	1.23 (0.05)	0.11	5.42
	OY5	10	0	0	16.30 (2.80)	0.85 (0.41)	10	ŝ	2	7.7 (1.48)	0.57 (0.16)	1.24 (0.05)	0.10	5.56
	OY6	10	0	0	10.80 (2.01)	0.31 (0.16)	11	ŝ	2	11.00 (3.03)	0.50 (0.17)	1.31 (0.07)	0.15	5.40
Pacific	Clatsop Beach	10	0	0	9.50 (1.21)	0.18 (0.09)	10	m	3	7.60 (1.01)	0.13 (0.02)	1.30 (0.07)	0.19	5.04
razor	Cannon Beach	5 ו	0	0	10.00 (1.48)	0.18 (0.05)	6	m		9.78 (1.47)	0.17 (0.02)	1.43 (0.09)	0.18	8.19
clam	Cape Meares	10	0	0	8.00 (2.82)	0.19 (0.22)	7	m	3	7.00 (2.57)	0.62 (0.33)	1.19 (0.07)	0.16	4.27
	Agate Beach	0	0	0	N/A	N/A	10	m	3	6.3 (0.91)	0.09 (0.01)	1.32 (0.12)	0.26	5.73
	Newport S. Beach	13	0	0	9.69 (1.49)	0.21 (0.12)	10	ŝ		9.30 (0.84)	0.14 (0.02)	1.44 (0.07)	0.21	7.04
	Coos Bay	12	0	0	10.50 (1.55)	0.23 (0.10)	0	0	0	N/A	N/A	1.46 (0.08)	0.31	4.73
	Bastendorff Beach	0	0	0	N/A	N/A	5	0	0	14.80 (1.24)	0.25 (0.01) 1.38 (0.10) 0.27	1.38 (0.10)	0.27	6.09
	Whiskey Creek 0	0	0	0	N/A	N/A	10	m	3	6.30 (0.87)	0.11 (0.02)	0.11 (0.02) 1.54 (0.13) 0.36	0.36	5.71
	Gold Beach	0	0	0	N/A	N/A	10	m	3	8.70 (1.47)	0.12 (0.02)	(0.02) 1.29 (0.09)	0.26	4.93
Notes: A ¹ Averagi ² Averagi ³ Averagi	Notes: Avg., Average; MP, microplastic; SE, \pm st ¹ Average number of MP per sample (SE). ² Average number of MP per gram of tissue (SE) ³ Average MP length in millimeters (SE). ⁴ Minimum MP lenoth at site in millimeters (SE).	microplastic; er sample (SE er gram of tis llimeters (SE). ite in millimet	SE, ± st. :). :sue (SE). ers (SE).	andard erro	r; 0Y1–0Y6: 0	yster site (ranc	domized). Rej	oorted va	lues incluc	de background	± standard error; OY1–OY6: oyster site (randomized). Reported values include background and processing fiber levels. (SE).	ig fiber levels.		

Table 1. Number of samples analyzed and average microplastic burden in Oregon Pacific oysters and Pacific razor clams by site and season, and average,

Baechler et al.

4

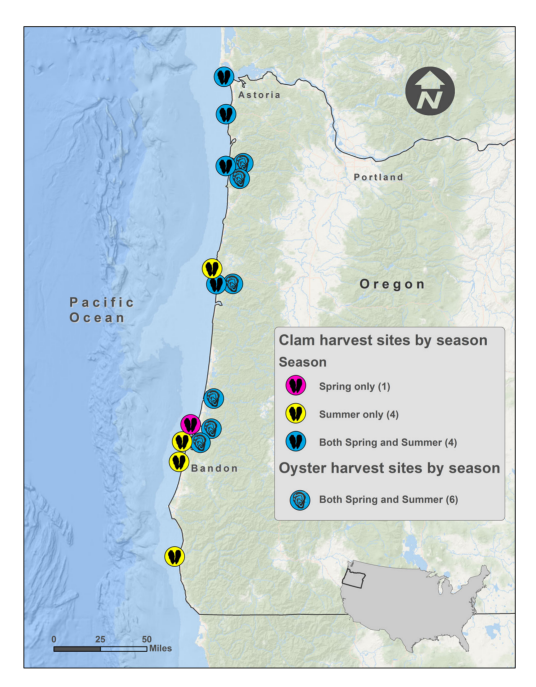


Fig. 1. The 2017 sample collection sites along the Oregon coast delineated for Pacific oysters and Pacific razor clams (Map credit: K. Scully-Engelmeyer; Service Layer Credits: Esri, Garmin, GEBCO, NOAA, NGDC, and other contributors; Sources: Esri, USGS, NOAA).

classified as nongut tissue. Separated gut and nongut tissues underwent the same digestion and microscope analyses as whole organism samples.

Quality control: Contamination quantification and prevention

One hundred percent cotton clothing, cotton lab coats, and nitrile gloves were worn at all times during sample processing, digestion, and analysis procedures. All shucking implements and glassware were rinsed three times with DI water filtered to $0.22 \,\mu$ m. To quantify procedural contamination, 11 replicates of 50 mL filtered DI water were frozen in 4 oz jars and underwent the same digestion and analysis process as organism samples. One procedural blank per week was chemically digested alongside field samples on a randomly generated day. Additionally, three procedural blanks were collected to quantify contamination introduced by the secondary digestion and hypersaline density separation of razor clam samples.

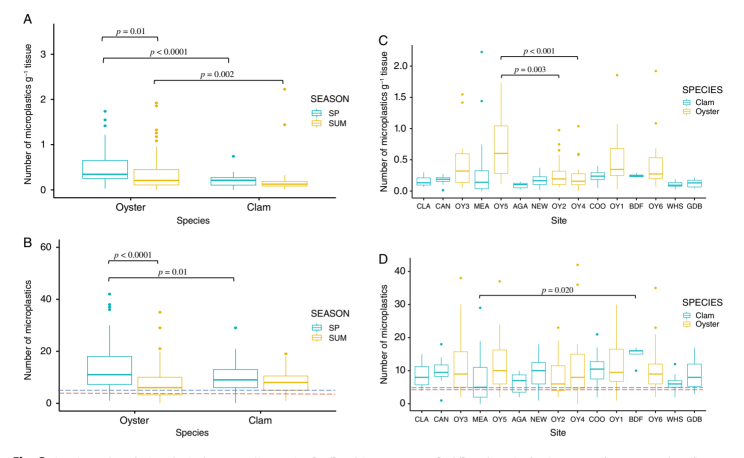


Fig. 2. (**A**, **B**) Number of microplastics by season (SP = spring [teal] and SUM = summer [gold]) and species for Oregon Pacific oysters and Pacific razor clams: (**A**) per gram of whole-organism tissue, and (**B**) per whole organism. Welch's *t*-tests were run to determine seasonal intraspecies and interspecies differences in log transformed values. *p* values show significant differences in microplastic burdens for seasons and/or species pairs indicated. (**C**, **D**) Number of microplastics in Oregon Pacific oysters (gold) and razor clams (teal): (**C**) per gram of whole-organism tissue, and (**D**) per whole organism. Dashed blue line indicates average contamination level for razor clams (6.11 microplastics per sample); dashed red line indicates contamination level for oysters (5.11 microplastics per sample). ANOVA and post hoc Tukey tests were run for each species to determine significance. *p* values show significant differences in microplastic burdens for site pairs indicated. Data are combined for both spring and summer sampling periods. Reported values include background and processing fiber levels. Sites are arranged north to south by latitude. OY1–OY6, randomized oyster sites; CLA, Clatsop Beach; CAN, Cannon Beach; MEA, Cape Meares; AGA, Agate Beach; NEW, Newport; COO, Coos Bay; BDF, Bastendorff Beach; WHS, Whiskey Run; GDB, Gold Beach.

During microscope analysis, a petri dish containing filtered DI water was placed adjacent to each sample on the microscope base and left open to the air to quantify airborne contaminants. After sample analysis, the control petri dish was analyzed for microplastics; any particles detected were assumed to be contamination and were measured and categorized.

Data analysis and availability

To identify differences between sample sites, seasons, and anatomical burdens, ANOVA and Welch's *t*-tests were conducted in the R statistical program (v1.2.1335) using the aov and t.test functions (R Core Team 2019). Linear regression models were used to examine relationships between biological parameters (shell length, body weight) and microplastic burdens. Microplastic concentrations are expressed as number of microplastics per sample or mean number of particles g⁻¹ tissue (wet weight; whole organisms only). Number of microplastics per sample and number of microplastics per gram of tissue variables were log transformed (log x + 1) prior to statistical analysis. The statistical cutoff (alpha) for all tests was 0.05 with standard error (SE) reported. Data and metadata are available in the Portland State University PDXScholar data repository.

Results

Quality control

Numerous measures were taken to minimize procedural contamination, but as with other studies (e.g., Li et al. 2015; Davidson and Dudas 2016; Qu et al. 2018; Su et al. 2018) it was not completely eliminated. Contamination in procedural controls (4.91 ± 1.11), microscope blanks (0.20 ± 0.03), and, for razor clams, a secondary digestion and separation step

ific oysters and Pacific razor clams (in number of microplastics per whole individual and per gram of whole	ated in bold.	
Table 2. Microplastic burden of Oregon Pacific oysters and Pacific razor clams (in number o	organism tissue). Significant differences are indicated in bold.	

		Ŧ	# Microplastics per gram of tissue	s per gram	of tissue	#	# Microplastics per whole organism	s per whole	organism
Species	Microplastic burden comparison	Test type	Test statistic	df	Significance	Test type	Test statistic	df	Significance
Both species (Pacific oyster and	Whole oyster to whole razor clam (both spring and summer)	t-test	<i>t</i> = -6.43	df = 199	<i>p</i> = <0.0001	t-test	<i>t</i> = −1.16	df = 235	<i>p</i> = 0.25
Pacific razor clam)	Whole oyster to whole razor clam (spring only)	t-test	<i>t</i> = -6.21	df = 89	<i>p</i> = <0.0001	t-test	t = -2.63	df = 103	<i>p</i> = <0.001
	Whole oyster to whole razor clam (summer only)	t-test	t = -3.24	df = 103	<i>p</i> = 0.002	t-test	t = -1.29	df = 112	<i>p</i> = 0.19
Pacific oyster	Gut vs. tissue	N/A				t-test	t = 0.48	df = 31	p = 0.63
	Spring vs. summer	t-test	t = 2.57	df = 121	p = 0.01	t-test	t = 4.41	df = 121	<i>p</i> = <0.0001
Pacific razor clam	Gut vs. tissue	N/A				t-test	t = -0.55	df = 39	p = 0.59
	Spring vs. summer	t-test	t = -0.29	df = 50	p = 0.77	t-test	t = 0.09	df = 71	p = 0.93
<i>Notes</i> : Number of micro log transformed data. Tc pled in both seasons (Cla	Notes: Number of microplastics per gram of tissue was not compared for gut and tissue samples, as mass was recorded for whole organism samples only. <i>E</i> -tests were conducted on log transformed data. To test temporal difference in Pacific oysters, data from all six sites sampled in both spring and summer were compared; for razor clams, only the four sites sampled in both seasons (Clatsop Beach, Cannon Beach, Cape Meares, Newport South Beach) were compared.	ed for gut data from Newport S	and tissue san all six sites sam outh Beach) w	nples, as mass npled in both s ere compared	was recorded for v spring and summer	vhole orga were com	nism samples c pared; for razoi	only. <i>t</i> -tests we r clams, only tl	ere conducted on he four sites sam-

Microplastics in Oregon bivalves

 (1.0 ± 0.0) was quantified (Supporting Information Appendix 2). From these controls and procedural blanks, total contamination in oyster and clam samples was estimated at 5.11 and 6.11 microplastics per sample, respectively. Average microplastic length detected as contamination (n = 124) for all sample types was 1.67 ± 0.11 mm, and most frequently detected colors in blanks and controls were colorless (79%) and blue (10%). As with multiple other studies (e.g., Li et al. 2015, 2016, 2018a; Davidson and Dudas 2016; Qu et al. 2018; Su et al. 2018; Rochman et al. 2019), we report microplastics detected in blank samples (Supporting Information Appendix 2), rather than performing a blank-subtraction on environmental results since controls were intended to provide a range of possible contamination levels introduced through laboratory procedures. As such, our reported numbers are estimated maximum possible microplastic concentrations.

Microplastic occurrence in study species

A total of 3,053 suspected microplastics were isolated from 320 whole-organism, gut-tissue, and nongut tissue samples. Over 99% of particles were microfibers (n = 3,026) averaging 1.34 mm in length (range = 0.10–8.72 mm). The remaining < 1% of microplastics were categorized as fragments (n = 12), beads (n = 5), films (n = 5), foams (n = 2), or unknown (n = 3). Colorless, blue, gray, and black were the most commonly observed fiber colors at 62%, 21%, 7%, and 4%, respectively.

Microplastics were present in organisms at all sites during both sampling periods and across the entire geographic range sampled with some discernible patterns (Table 1, Fig. 2).

Mean microplastic concentrations in whole organisms was 10.95 \pm 0.77 in Pacific oysters (range = 0–42) and 8.84 \pm 0.45 in razor clams (range = 0-38). Mean microplastic burden per gram of tissue in whole organisms was significantly different between ovsters $(0.35 \pm 0.04 \text{ g}^{-1} \text{ tissue})$ and razor clams $(0.16 \pm 0.02 \text{ g}^{-1} \text{ tissue; } t = -6.43, \text{ df} = 199; p \le 0.0001), \text{ but}$ number of microplastics per whole organism was not significantly different (Table 2; t = -1.16, df = 235, p = 0.25). FTIR analysis of 26 individual fibers extracted from whole organisms indicates material types of PET (n = 8), acrylic (n = 2), aramid (n = 1), zein (n = 1), and cellophane, a cellulose-based material (n = 10). Because cellophane exhibited a low spectral match percentage (20-67%) relative to other materials (aramid: 68%; all others: 80-95%), we believe the cellophanecharacterized fibers should be more broadly deemed cellulosebased material types. Additional fibers (n = 4) were run but no material type matches were determined, most likely due to small fiber width and concomitant low signal to noise data.

Temporal differences

Significant intraspecies and interspecies differences in microplastic burdens were detected during the two sampling periods (Fig. 2A,B). Spring Pacific oysters contained significantly more microplastics than summer; on average, whole spring oysters contained 13.74 ± 1.16 microplastics (0.45 \pm 0.05 g⁻¹ tissue) whereas summer oysters contained

 8.16 ± 0.88 (0.26 ± 0.05 g⁻¹ tissue; whole organism: *t* = 4.41; df = 121; p < 0.0001; MP g⁻¹ tissue: t = 2.57; df = 121; p = 0.01). There was no significant temporal difference in microplastic burden for clams when the four sites sampled in spring and summer were compared (Clatsop Beach, Cannon Beach, Cape Meares, Newport South Beach). Spring razor clams contained 9.54 ± 0.81 microplastics per whole individual (0.19 \pm 0.02 g⁻¹ tissue) whereas summer had 8.35 \pm 0.51 $(0.14 \pm 0.04 \text{ g}^{-1} \text{ tissue; whole organism: } t = 0.09; \text{ df} = 71;$ p = 0.93; MP g⁻¹ tissue: t = -0.29; df = 50; p = 0.77). When comparing spring oysters to spring razor clams, spring oysters contained more microplastics g^{-1} tissue (Table 2; t = -6.21; df = 89; $p \le 0.0001$) and more microplastics per whole sample (Table 2; t = -2.63; df = 103; p = 0.01). Summer oysters contained more microplastics g^{-1} tissue than summer razor clams (Table 2; t = -3.24; df = 103; p = 0.002), but not more plastics per whole organism (Table 2; t = -1.29; df = 112; p = 0.19).

Site differences

ANOVA and post hoc Tukey tests revealed site-specific differences in microplastic burdens per gram of whole oyster tissue from two site pairings (Fig. 2C; p < 0.001 and p = 0.003). Site-specific differences in number of microplastics per individual were not detected in oysters (Fig. 2D; F = 0.56; df = 5; p = 0.73). For razor clams, site-specific differences in microplastics per gram of tissue were not detected (Fig. 2C), but were for microplastics per individual in one site pairing (Fig. 2D; F = 2.54; df = 8; p = 0.020).

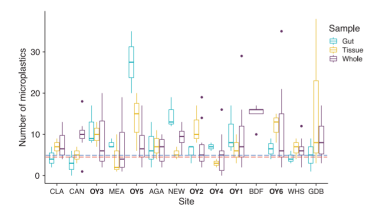


Fig. 3. Comparison of sample types for summer-collected Pacific oysters (site names bolded) and Pacific razor clams. Reported values include background and processing fiber levels. Gut = gut-tissue; Tissue = nongut tissue; Whole = whole organism. Dashed blue line indicates average contamination level for razor clams (6.11 microplastics per sample); dashed red line indicates contamination level for oysters (5.11 microplastics per sample). Sites are arranged north to south by latitude. OY1–OY6, randomized oyster sites; CLA = Clatsop Beach; CAN = Cannon Beach; MEA = Cape Meares; AGA = Agate Beach; NEW = Newport; BDF = Bastendorff Beach; WHS = Whiskey Run; GDB = Gold Beach.

Anatomical burdens

Microplastics were detected in whole organism, gut-tissue, and nongut tissue samples in both species from all sites sampled in the summer, except at Bastendorff Beach where sample size precluded separate gut and tissue analyses, and Coos Bay, which was not sampled in summer (Fig. 3). Microplastic burden (number of plastics per sample) did not differ between gut-tissue and nongut tissue in either species (Table 2; Oysters: t = 0.48; df = 31; p = 0.63; Clams: t = -0.55; df = 39; p = 0.59). In oysters, average microplastic burden was 10.69 ± 2.01 in gut-tissue and 9.41 ± 1.30 in nongut tissue samples. In razor clams, average microplastic burden was 6.57 ± 1.02 in gut-tissue and 7.43 ± 1.64 in nongut tissue samples.

Shell length, body weight, and microplastic burden

Regression analyses revealed shell length (in mm) was not significantly correlated with number of microplastics per whole organism in oysters (F = 0.081, df = 122, $R^2 = -0.008$, p = 0.777) or razor clams (F = 0.421, df = 118, $R^2 = -0.005$, p = 0.518). Similarly, body weight (in g) was not significantly correlated with number of microplastics per whole organism in oysters (F = 0.430, df = 122, $R^2 = -0.005$, p = 0.514) or razor clams (F = 1.355, df = 118, $R^2 = 0.003$, p = 0.247).

Discussion

Microplastics were present in both Pacific ovsters and Pacific razor clams collected from all 15 Oregon coast sample sites in both spring and summer 2017. All whole organisms (n = 245) except one oyster and one razor clam contained at least one plastic particle. Microplastic concentrations varied significantly by season in oysters but not razor clams. Limited site-specific differences in microplastic burden were detected. Contamination in our samples combined with relatively small sample size may have influenced the lack of site differences during statistical analyses. No anatomical microplastic burden differences were detected between gut and nongut tissues in either species, and organism size did not correlate with microplastic burden. Both Pacific oysters and razor clams are low trophic level species important to both the ecology of Oregon's nearshore and estuarine environments and humans who culture or consume them. To our knowledge, this is the first study to document microplastics in Pacific oysters and razor clams harvested in Oregon. Various edible ovster and clam species have been found to contain microplastics elsewhere in the world, including Asia, British Columbia, and Europe (e.g., Mathalon and Hill 2014; Van Cauwenberghe and Janssen 2014; Li et al. 2015; Davidson and Dudas 2016; Su et al. 2018). In this study, the average number of microplastics found in Pacific oysters and razor clams $(0.35 \pm 0.04 \text{ g}^{-1} \text{ tissue and } 0.16 \pm 0.02 \text{ g}^{-1} \text{ tissue)}$ was low compared to average concentrations in Pacific oysters in France, China, and Tunisia of 0.47, 0.62, and 1.5 items g^{-1} tissue (Van Cauwenberghe and Janssen 2014; Abidli et al. 2019;

Teng et al. 2019), mussels from China of 2.2–-2.4 items g^{-1} tissue (Li et al. 2015, 2016), and manila clams from British Columbia, Canada of 0.9–1.7 items g^{-1} tissue (Davidson and Dudas 2016). Concentrations from this study are in the range of those found in blue mussels in France and Belgium of 0.23 and 0.26 items g^{-1} tissue (De Witte et al. 2014; Phuong et al. 2018), and are the low end of concentrations found in manila clams in China of 0.3–4.9 items g^{-1} tissue (Su et al. 2018). These patterns may result from the relatively small human population residing on the Oregon coast.

As this is the first study to document prevalence of microplastics in Pacific razor clams (*S. patula*), no comparisons are possible between our Oregon samples and other areas. Due to the importance of the recreational, commercial, and tribal razor clam fisheries in the broader PNW, microplastic burden data from other states and territories in the region should be collected to help elucidate possible larger-scale patterns in prevalence.

In estuarine-grown oysters, collection season appears to influence microplastic burden more than harvest location. Additional research is needed to identify the environmental or anthropogenic factors driving higher microplastic burden in ovsters in the spring. Seasonal microplastic differences in oysters but not razor clams may be a function of habitat. Oysters inhabit estuarine environments, which receive land-based stormwater and wastewater inputs before ocean-facing beaches do; therefore, pulse inputs of microplastics may be more concentrated in estuaries than along the open coast. Precipitation was at least 100% higher than normal in all coastal counties and up to three orders of magnitude higher in some coastal areas in April 2017 compared to July 2017, which was characterized by at least 50% lower than normal precipitation in all coastal counties (NOAA 2017). Therefore, seasonal differences in ovsters may be driven, in part, by seasonal precipitation and resultant stormwater fluctuations. Another possibility is that the nature of clothing laundered in the spring-cold weather clothes, possibly dominated by insulating synthetic materials-may increase microfiber levels in WWTP outputs when compared to clothing items laundered in the summer. Other potential seasonal factors include temperature-associated influences on metabolic and feeding rates, which may be depressed during colder seasons, and life history events like spawning and associated physiological responses. Differences in aquaculture techniques, such as degree of plastic use by oyster growers, may contribute to variation in oyster microplastic burdens between sites and over time; however, grower-specific culture techniques were not assessed in this study and previous studies in the PNW have failed to find a connection between aquaculture and microplastic burden in cultured Pacific oysters and manila clams when compared to wild-grown organisms (Davidson and Dudas 2016; Covernton et al. 2019). Temporal differences identified in this study indicate oysters may be able to clear

microplastics from their system over time, as previously shown in laboratory studies where manila clams and blue mussels (29–40 mm in length) eliminated microplastics in feces and pseudofeces when depurated in clean water, with up to 60% of particles cleared from the body in as little as 9 h (Xu et al. 2017; Woods et al. 2018). However, elimination of microplastics was not detected in blue mussels (50–55 mm length) during a depuration period of 2 h (Rist et al. 2018). While, in these examples, depuration was studied in bivalves smaller than our study organisms (Supporting Information Appendix 1), the results are promising and warrant further research. Depuration of oysters or razor clams in freshwater for some period of time prior to sale may be a fruitful avenue for reducing anthropogenic debris in those seafood items.

Visual microscopy is routinely used in microplastics research due to the relatively wide availability of microscopes, but it likely introduces error to microplastic counts. Recent studies indicate visual microscopy can either overestimate or underestimate microplastic counts depending on particle shape and size (Song et al. 2015); thus, additional validation methods should be used to supplement visual analysis methods. In this study, FTIR techniques were used to ground truth material composition of a randomly selected subset (n = 26) of the 2428 microfibers found in whole samples, which were subsequently identified as PET (n = 8), acrylic (n = 2), aramid (n = 1), zein (n = 1), and cellophane, a cellulose-based material (n = 10). Our low percentage of validated fibers was due to funding limitations and lack of on-site equipment. Polyethyelene terephthalate, acrylic, and aramid fibers have been previously found in organisms (e.g., Li et al. 2015, 2018a; Nelms et al. 2018), and zein (a corn-based protein used in bioplastics) has been isolated from WWTP sludge (Bayo et al. 2016). Fibrous cellophane, the putative material type comprising the largest proportion of successfully validated fibers (n = 10), is made of heavily modified cellulose but has previously been categorized as a microplastic in studies that identified cellophane fibers in bivalves (Li et al. 2016, 2018a; Ding et al. 2018). Due to low spectral match percentage of cellophane (20-67%) relative to other materials matched to known spectra (aramid: 68%; all others: 80-95%), we believe the cellophane-characterized fibers should be more broadly deemed cellulose-based material types.

Our lower size limit of detection for microplastics was 0.063 mm owing to the mesh size of the sieve used, so microplastics smaller than 0.063 mm in length may be underestimated using these methods. Microplastics between 0.10 and 8.72 mm in length were included in this report, as they are of equal interest as microplastics fitting the conventional 0.0001–5 mm definition. Future studies on these and other bivalve species should include methods capable of detecting both micro and nanoplastics $(1 \times 10^{-6} \text{ to } 1 \times 10^{-4} \text{ mm})$, as particles between 1×10^{-4} and 2×10^{-2} mm can penetrate internal organ barriers (Lusher et al. 2017).

In this study, we found that all whole organisms (n = 245)except one ovster and one razor clam contained at least one microplastic, though we acknowledge that some of these detections may have been influenced by contamination in the laboratory. For this reason, we ran several types of blank and control samples during processing and analysis to quantify it. Microfiber contamination may have been due to presence in KOH pellets used for chemical digestion, fibers shed from clothing, laboratory furniture (chairs), or airborne particles. Average contamination represented 46.7% of the average microplastic burdens reported for whole oysters, and 69.1% of average microplastic reported for whole clams. While contamination in this study appears high, it is consistent with similar studies that report between 51% and 94% of detected microplastic values in mussels and clams may represent contamination (Mathalon and Hill 2014; Davidson and Dudas 2016). Contamination documented in this and other microplastic investigations highlights the ubiquity of anthropogenic microfibers in the environment.

The degree to which microplastics pose a threat to coastal marine ecology or bivalve predators (including humans) is still unclear; however, this study provides valuable insights about spatial and temporal variability in microplastic prevalence in important commercial species, sheds light on potential ecological concerns related to microplastic contamination, and serves as a baseline from which future microplastic studies in the region can draw comparisons. Future research on extent of microplastic encounter rates, consumption, and effects on biological endpoints are critical to better understand potential population-level effects on bivalves and marine organisms around the world.

References

- Abidli, S., Y. Lahbib, and N. Trigui El Menif. 2019. Microplastics in commercial molluscs from the lagoon of Bizerte (Northern Tunisia). Mar. Pollut. Bull. **142**: 243–252. doi:10. 1016/j.marpolbul.2019.03.048
- Akpan, N. 2014. Earth & environment: Microplastics lodge in crab gills, guts: Creatures absorb particles through food and via respiration. Sci. News 186: 9. doi:10.1002/scin.2014. 5591860307
- Barnes, D. K. A., F. Galgani, R. C. Thompson, and M. Barlaz. 2009. Accumulation and fragmentation of plastic debris in global environments. Philos. Trans. R Soc. B Biol. Sci. 364: 1985–1998. doi:10.1098/rstb.2008.0205
- Barrows, A. P. W., S. E. Cathey, and C. W. Petersen. 2018. Marine environment microfiber contamination: Global patterns and the diversity of microparticle origins. Environ. Pollut. 237: 275–284. doi:10.1016/j.envpol.2018.02.062
- Bayo, J., S. Olmos, J. López-Castellanos, and A. Alcolea. 2016. Microplastics and microfibers in the sludge of a municipal wastewater treatment plant. Int. J. Sustain. Dev. Plan. 11: 812–821. doi:10.2495/SDP-V11-N5-812-821

- Browne, M. A., A. Dissanayake, T. S. Galloway, D. M. Lowe, and R. C. Thompson. 2008. Ingested microscopic plastic translocates to the circulatory system of the mussel, *Mytilus edulis* (L.). Environ. Sci. Technol. **42**: 5026–5031. doi:10. 1021/es800249a
- Carr, S. A., J. Liu, and A. G. Tesoro. 2016. Transport and fate of microplastic particles in wastewater treatment plants. Water Res. 91: 174–182. doi:10.1016/j.watres.2016.01.002
- Cole, M., P. Lindeque, C. Halsband, and T. S. Galloway. 2011. Microplastics as contaminants in the marine environment: A review. Mar. Pollut. Bull. 62: 2588–2597. doi:10.1016/j. marpolbul.2011.09.025
- Covernton, G. A., B. Collicutt, H. J. Gurney-Smith, C. M. Pearce, J. F. Dower, P. S. Ross, and S. E. Dudas. 2019. Microplastics in bivalves and their habitat in relation to shellfish aquaculture proximity in coastal British Columbia, Canada. Aquacult. Environ. Interact. 11: 357–374. doi:10. 3354/aei00316
- Crossett, K., B. Ache, P. Pacheco, and K. Haber. 2013, *National coastal population report, population trends from 1970 to 2020*. NOAA state of the coast report series. US Department of Commerce. Available from http://oceanservice.noaa.gov. proxy.lib.pdx.edu/facts/coastal-population-report.pdf
- Davidson, K., and S. E. Dudas. 2016. Microplastic ingestion by wild and cultured Manila clams (*Venerupis philippinarum*) from Baynes Sound, British Columbia. Arch. Environ. Contam. Toxicol. **71**: 147–156. doi:10.1007/s00244-016-0286-4
- de Falco, F. D., E. D. Pace, M. Cocca, and M. Avella. 2019. The contribution of washing processes of synthetic clothes to microplastic pollution. Sci. Rep. 9: 1–11. doi:10.1038/ s41598-019-43023-x
- De Witte, B., L. Devriese, K. Bekaert, S. Hoffman, G. Vandermeersch, K. Cooreman, and J. Robbens. 2014. Quality assessment of the blue mussel (*Mytilus edulis*): Comparison between commercial and wild types. Mar. Pollut. Bull. **85**: 146–155. doi:10.1016/j.marpolbul.2014. 06.006
- Ding, J.-F., J.-X. Li, C.-J. Sun, C.-F. He, F.-H. Jiang, F.-L. Gao, and L. Zheng. 2018. Separation and identification of microplastics in digestive system of bivalves. Chin. J. Anal. Chem. 46: 690–697. doi:10.1016/S1872-2040(18)61086-2
- Glude, J. B., and K. K. Chew. 1982. Shellfish aquaculture in the Pacific Northwest. University of Alaska, Anchorage. Alaska Sea Grant Report 82-2:291-304. *In* G. B. Pauley, B. van der Raay, and D. Troutt [eds.], *1988. Species profiles: Life histories and environmental requirements of coastal fishes and invertebrates (Pacific Northwest), Pacific Oyster.* Washington Cooperative Fishery Research Unit. Available from https:// apps.dtic.mil/dtic/tr/fulltext/u2/a203409.pdf
- Hantoro, I., A. J. Löhr, F. G. A. J. V. Belleghem, B. Widianarko, and A. M. J. Ragas. 2019. Microplastics in coastal areas and seafood: Implications for food safety. Food Addit. Contam. Part A Chem. Anal. Control Expo. Risk Assess. 36: 674–711. doi:10.1080/19440049.2019.1585581

- Harris, J. 2008. Pacific oyster, *Crassostrea gigas* (Thunberg, 1793). *In, Aquatic invasive species profile*. Aquatic Invasion Ecology.
- Jambeck, J. R., R. Geyer, C. Wilcox, T. R. Siegler, M. Perryman, A. Andrady, R. Narayan, and K. L. Law. 2015. Plastic waste inputs from land into the ocean. Science **347**: 768–771. doi:10.1126/science.1260352
- Kolandhasamy, P., L. Su, J. Li, X. Qu, K. Jabeen, and H. Shi. 2018. Adherence of microplastics to soft tissue of mussels: A novel way to uptake microplastics beyond ingestion. Sci. Total Environ. 610–611: 635–640. doi:10.1016/j.scitotenv. 2017.08.053
- Lebreton, L. C. M., J. van der Zwet, J.-W. Damsteeg, B. Slat, A. Andrady, and J. Reisser. 2017. River plastic emissions to the world's oceans. Nat. Commun. 8: 15611. doi:10.1038/ ncomms15611
- Li, H.-X., and others. 2018a. Microplastics in oysters *Saccostrea cucullata* along the Pearl River Estuary, China. Environ. Pollut. **236**: 619–625. doi:10.1016/j.envpol.2018.01.083
- Li, J., D. Yang, L. Li, K. Jabeen, and H. Shi. 2015. Microplastics in commercial bivalves from China. Environ. Pollut. **207**: 190–195. doi:10.1016/j.envpol.2015.09.018
- Li, J., X. Qu, L. Su, W. Zhang, D. Yang, P. Kolandhasamy, D. Li, and H. Shi. 2016. Microplastics in mussels along the coastal waters of China. Environ. Pollut. **214**: 177–184. doi:10.1016/j.envpol.2016.04.012
- Li, J., C. Green, A. Reynolds, H. Shi, and J. M. Rotchell. 2018. Microplastics in mussels sampled from coastal waters and supermarkets in the United Kingdom. Environ. Pollut. 241: 35–44. doi:10.1016/j.envpol.2018.05.038
- Link, T. 2000. History and status of Oregon's Pacific razor clam resources, p. 1–25. Information reports 2000-06. Oregon Department of Fish and Wildlife, Marine Resources Program. Available from https://www.dfw.state.or.us/MRP/ publications/docs/razorclams200006.pdf
- Lusher, A., P. Hollman, and J. Mendoza-Hill. 2017. Microplastics in fisheries and aquaculture: Status of knowledge on their occurrence and implications for aquatic organisms and food safety. FAO Fisheries and Aquaculture technical paper. Available from http://www.fao.org/3/a-i7677e.pdf
- Mathalon, A., and P. Hill. 2014. Microplastic fibers in the intertidal ecosystem surrounding Halifax Harbor, Nova Scotia. Mar. Pollut. Bull. **81**: 69–79. doi:10.1016/j.marpolbul. 2014.02.018
- Mintenig, S. M., I. Int-Veen, M. G. J. Löder, S. Primpke, and G. Gerdts. 2017. Identification of microplastic in effluents of waste water treatment plants using focal plane arraybased micro-Fourier-transform infrared imaging. Water Res. 108: 365–372. doi:10.1016/j.watres.2016.11.015
- Moore, C., G. Lattin, and A. Zellers. 2005, A brief analysis of organic pollutants sorbed to pre and post production plastic particles from the Los Angeles and San Gabriel river watersheds. Algalita Marine Research Foundation. Available from https://www.researchgate.net/publication/253305700_A_

Brief_Analysis_of_Organic_Pollutants_Sorbed_to_Pre_and_ PostProduction_Plastic_Particles_from_the_Los_Angeles_ and_San_Gabriel_River_Watersheds

- Napper, I. E., and R. C. Thompson. 2016. Release of synthetic microplastic plastic fibres from domestic washing machines: Effects of fabric type and washing conditions. Mar. Pollut. Bull. **112**: 39–45. doi:10.1016/j.marpolbul. 2016.09.025
- Nelms, S. E., T. S. Galloway, B. J. Godley, D. S. Jarvis, and P. K. Lindeque. 2018. Investigating microplastic trophic transfer in marine top predators. Environ. Pollut. 238: 999–1007. doi:10.1016/j.envpol.2018.02.016
- NOAA. 2017. Northwest River Forecast Center. [accessed 2019 January 1]. Available from https://www.nwrfc.noaa. gov/rfc/
- Pauley, G. B., B. Van Der Raay, and D. Troutt. 1988, Species profiles: Life histories and environmental requirements of coastal fishes and invertebrates (Pacific Northwest), Pacific Oyster. Washington Cooperative Fishery Research Unit. Available from https://apps.dtic.mil/dtic/tr/fulltext/u2/a203409.pdf
- Phuong, N. N., L. Poirier, Q. T. Pham, F. Lagarde, and A. Zalouk-Vergnoux. 2018. Factors influencing the microplastic contamination of bivalves from the French Atlantic coast: Location, season and/or mode of life? Mar. Pollut. Bull. **129**: 664–674. doi:10.1016/j.marpolbul.2017. 10.054
- Qu, X., L. Su, H. Li, M. Liang, and H. Shi. 2018. Assessing the relationship between the abundance and properties of microplastics in water and in mussels. Sci. Total Environ. 621: 679–686. doi:10.1016/j.scitotenv.2017.11.284
- R Core Team. 2019. R: A language and environment for statistical computing. R Foundation for Statistical Computing. [accessed 2019 January 1] Available from https://www.Rproject.org/
- Ribeiro, F., A. R. Garcia, B. P. Pereira, M. Fonseca, N. C. Mestre, T. G. Fonseca, L. M. Ilharco, and M. J. Bebianno. 2017. Microplastics effects in *Scrobicularia plana*. Mar. Pollut. Bull. **122**: 379–391. doi:10.1016/j. marpolbul.2017.06.078
- Rist, S., I. M. Steensgaard, O. Guven, T. G. Nielsen, L. H. Jensen, L. F. Møller, and N. B. Hartmann. 2018. The fate of microplastics during uptake and depuration phases in a blue mussel exposure system. Environ. Toxicol. Chem. **38**: 99–105. doi:10.1002/etc.4285
- Rochman, C. M., and others. 2015. Anthropogenic debris in seafood: Plastic debris and fibers from textiles in fish and bivalves sold for human consumption. Sci. Rep. 5: 14340. doi:10.1038/srep14340
- Rochman, C. M., and others. 2019. Rethinking microplastics as a diverse contaminant suite. Environ. Toxicol. Chem. 38: 703–711. doi:10.1002/etc.4371
- Song, Y. K., S. H. Hong, M. Jang, G. M. Han, M. Rani, J. Lee, and W. J. Shim. 2015. A comparison of microscopic and spectroscopic identification methods for analysis of

microplastics in environmental samples. Mar. Pollut. Bull. **93**: 202–209. doi:10.1016/j.marpolbul.2015.01.015

- Su, L., H. Cai, P. Kolandhasamy, C. Wu, C. M. Rochman, and H. Shi. 2018. Using the Asian clam as an indicator of microplastic pollution in freshwater ecosystems. Environ. Pollut. 234: 347–355. doi:10.1016/j.envpol.2017.11.075
- Sussarellu, R., and others. 2016. Oyster reproduction is affected by exposure to polystyrene microplastics. Proc. Natl. Acad. Sci. USA **113**: 2430–2435. doi:10.1073/pnas. 1519019113
- Teng, J., and others. 2019. Microplastic in cultured oysters from different coastal areas of China. Sci. Total Environ. 653: 1282–1292. doi:10.1016/j.scitotenv.2018.11.057
- Teuten, E. L., S. J. Rowland, T. S. Galloway, and R. C. Thompson. 2007. Potential for plastics to transport hydrophobic contaminants. Environ. Sci. Technol. 41: 7759–7764. doi:10.1021/es071737s
- Teuten, E. L., and others. 2009. Transport and release of chemicals from plastics to the environment and to wildlife. Philos. Trans. R Soc. B Biol. Sci. 364: 2027–2045. doi:10. 1098/rstb.2008.0284
- Thompson, R. C., Y. Olsen, R. P. Mitchell, A. Davis, S. J. Rowland, A. W. G. John, D. McGonigle, and A. E. Russell. 2004. Lost at sea: Where is all the plastic? Science **304**: 838–838. doi:10.1126/science.1094559
- UNEP. 2016, Marine plastic debris and microplastic technical report. United Nations Environmental Programme.
- Van Cauwenberghe, L., and C. R. Janssen. 2014. Microplastics in bivalves cultured for human consumption. Environ. Pollut. **193**: 65–70. doi:10.1016/j.envpol.2014.06.010
- von Moos, N., P. Burkhardt-Holm, and A. Köhler. 2012. Uptake and effects of microplastics on cells and tissue of the Blue Mussel *Mytilus edulis* L. after an experimental exposure. Environ. Sci. Technol. **46**: 11327–11335. doi:10. 1021/es302332w
- Waite, H. R., M. J. Donnelly, and L. J. Walters. 2018. Quantity and types of microplastics in the organic tissues of the eastern oyster *Crassostrea virginica* and Atlantic mud crab *Panopeus herbstii* from a Florida estuary. Mar. Pollut. Bull. **129**: 179–185. doi:10.1016/j.marpolbul.2018.02.026

- Wang, J., Z. Tan, J. Peng, Q. Qiu, and M. Li. 2016. The behaviors of microplastics in the marine environment. Mar. Environ. Res. **113**: 7–17. doi:10.1016/j.marenvres.2015.10.014
- Ward, J. E., and D. J. Kach. 2009. Marine aggregates facilitate ingestion of nanoparticles by suspension-feeding bivalves. Mar. Environ. Res. 68: 137–142. doi:10.1016/j.marenvres. 2009.05.002
- Woods, M. N., M. E. Stack, D. M. Fields, S. D. Shaw, and P. A. Matrai. 2018. Microplastic fiber uptake, ingestion, and egestion rates in the blue mussel (*Mytilus edulis*). Mar. Pollut. Bull. **137**: 638–645. doi:10.1016/j.marpolbul.2018. 10.061
- Xu, X.-Y., W. T. Lee, A. K. Y. Chan, H. S. Lo, P. K. S. Shin, and S. G. Cheung. 2017. Microplastic ingestion reduces energy intake in the clam *Atactodea striata*. Mar. Pollut. Bull. **124**: 798–802. doi:10.1016/j.marpolbul.2016.12.027
- Zhang, H. 2017. Transport of microplastics in coastal seas. Estuar. Coast. Shelf Sci. **199**: 74–86. doi:10.1016/j.ecss. 2017.09.032

Acknowledgments

We thank E. Foster for procedural and analytical guidance; A. Marquardt, M. Rogers, S. Rumrill of ODFW and N. Jensen of ODA for assistance with razor clam sample collection and coordination; P. Wilkinson at the University of New Hampshire for her FTIR work; K. Scully-Engelmeyer for mapping assistance; J. Dorsheimer of Thermo Scientific for spectral matching assistance; A. Strecker for use of her microscope; A. Bolm, S. Holland, C. Orr, J. Adelsheim, M. Rollins, C. Homer, M. Jauregui, H. Smiley, D. Van Leuven, and C. Tenorio for assistance in the lab and field; and the Applied Coastal Ecology lab for valuable input and support. We also thank the two anonymous peer reviewers and editorial team for providing comments that greatly improved the manuscript. Funding for this project was provided by Oregon Sea Grant (Award NA14OAR4170064), the Edward D. & Olive C. Bushby Scholarship, the OR American Fisheries Society, and the Portland State University Institute for Sustainable Solutions.

> Submitted 20 March 2019 Revised 04 September 2019 Accepted 22 September 2019