Influence of Anthropogenic Nutrient Inputs on Rates of Coastal Ocean Nitrogen and Carbon Cycling in the Southern California Bight, USA Southern California Coastal Water Research Project Orange County Sanitation District

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Abstract

Coastal nitrogen (N) enrichment is a global environmental problem that can influence acidification, deoxygenation, and subsequent habitat loss in ways that can be synergistic with global climate change impacts. In the Southern California Bight, an eastern boundary upwelling system, modeling of wastewater discharged through ocean outfalls has shown that it effectively doubles N loading to urban coastal waters. However, effects of wastewater outfalls on biogeochemical rates of primary production and respiration, key processes through which coastal acidification and deoxygenation are manifested, have not been directly linked to observed trends in ambient chlorophyll a, oxygen and pH. In this paper, we compare observations of nutrient concentrations and forms, as well as rates of biogeochemical cycling, in areas within treated wastewater effluent plumes compared to areas spatially distant from ocean outfalls where we expected minimum influence of the plume. We document that wastewater nutrient inputs have an immediate, local effect on nutrient stoichiometry, elevating ammonium and nitrite concentrations by a mean of 4 µM and 0.2 µM, respectively, increasing dissolved nitrogen: phosphorus ratios by a mean of 7 and slightly increasing chlorophyll a by a mean of $1 \mu g$ L-1 in the upper 60 m of the watercolumn, as well as increasing rates of nitrification within the plume by a mean of 17 nmol L-1 day-1 and increasing δ 13C and δ 15N of suspended particulate matter, an integrated measure of primary production, by a mean of 1.3 a significant near plume effect on δ 180 and δ 15N of the dissolved nitrate+nitrite, an indicator of nitrate+nitrite assimilation into the biomass, instantaneous rates of primary production and respiration, or dissolved oxygen concentration, suggesting any potential impact from wastewater on these is moderated by other factors, notably mixing of water masses. These results indicate that a "reference-area" approach, wherein stations within or near the zone of initial dilution (ZID) from the wastewater outfall are compared to stations farther afield (reference areas) to assess contaminant impacts, may be insufficient to document regional scale impacts of nutrients.

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1 Abstract

2 Coastal nitrogen (N) enrichment is a global environmental problem that can influence 3 acidification, deoxygenation, and subsequent habitat loss in ways that can be synergistic with 4 global climate change impacts. In the Southern California Bight, an eastern boundary upwelling 5 system, modeling of wastewater discharged through ocean outfalls has shown that it 6 effectively doubles N loading to urban coastal waters. However, effects of wastewater outfalls on biogeochemical rates of primary production and respiration, key processes through which 7 8 coastal acidification and deoxygenation are manifested, have not been directly linked to 9 observed trends in ambient chlorophyll a, oxygen and pH. In this paper, we compare 10 observations of nutrient concentrations and forms, as well as rates of biogeochemical cycling, in areas within treated wastewater effluent plumes compared to areas spatially distant from 11 12 ocean outfalls where we expected minimum influence of the plume. We document that 13 wastewater nutrient inputs have an immediate, local effect on nutrient stoichiometry, 14 elevating ammonium and nitrite concentrations by a mean of 4 μ M and 0.2 μ M, respectively, 15 increasing dissolved nitrogen: phosphorus ratios by a mean of 7 and slightly increasing chlorophyll α by a mean of 1 µg L⁻¹ in the upper 60 m of the watercolumn, as well as increasing 16 17 rates of nitrification within the plume by a mean of 17 nmol L⁻¹ day⁻¹ and increasing δ^{13} C and 18 δ^{15} N of suspended particulate matter, an integrated measure of primary production, by a 19 mean of 1.3 ‰ and 1 ‰, respectively. We did not observe a significant near plume effect on δ^{18} O and δ^{15} N of the dissolved nitrate+nitrite, an indicator of nitrate+nitrite assimilation into 20 21 the biomass, instantaneous rates of primary production and respiration, or dissolved oxygen 22 concentration, suggesting any potential impact from wastewater on these is moderated by 23 other factors, notably mixing of water masses. These results indicate that a "reference-area" 24 approach, wherein stations within or near the zone of initial dilution (ZID) from the 25 wastewater outfall are compared to stations farther afield (reference areas) to assess 26 contaminant impacts, may be insufficient to document regional scale impacts of nutrients.

1 Introduction

2 Globally, offshore ocean outfalls have been considered an effective and reliable 3 strategy for disposing of treated industrial and domestic wastewater (Wood et al. 1993, 4 Roberts et al. 2010). However, wastewater effluent released through these outfalls are rarely 5 treated to remove nutrients and, consequently, they have been implicated in eutrophication of 6 coastal waters (Roberts et al. 2010, Powley et al. 2016, Valiela et al. 2016). Eutrophication can 7 have impacts on coastal habitats that can be synergistic with global changes, such as increasing 8 frequency and occurrence of algal blooms (Howarth et al. 2002, Glibert et al. 2006), coastal 9 acidification (Borges and Gypens 2010, Wallace et al. 2014), and deoxygenation (Rabalais et al. 10 2014, Breitburg et al. 2018) through enhanced heterotrophic respiration rates. Given 11 appropriate light and temperature conditions, eutrophication initially causes a temporary 12 drawdown of CO₂ concentrations at the surface due to the intense biological productivity of 13 the associated algal bloom (Borges and Gypens 2010). Subsequently, eutrophication lowers pH, because it provides conditions for greater heterotrophic respiration rates (i.e., 14 15 decomposition, also referred to as remineralization) by organisms such as bacteria. This 16 respiration process oxidizes organic matter, draws down local oxygen levels, releases CO₂, 17 and—in extreme cases—can lead to hypoxic dead zones (Diaz and Rosenberg, 2008). The 18 depth of the pH change associated with anthropogenic eutrophication is a function of where 19 organic matter is respired. In many shallow coastal shelf systems, this acidification occurs at or 20 near the bottom sediments, where organic matter is oxidized (Waldbusser et al. 2010, Sunda 21 and Cai 2012). In the absence of global strategies to mitigate coastal ocean changes, local 22 managers are being urged to consider the management of local, land-based nutrient sources 23 to slow the progression of acidification and deoxygenation and their impact on coastal habitats 24 (Kelly et al. 2011, Strong et al. 2014, Chan et al. 2016).

The role of coastal nutrient discharges and coastal eutrophication in exacerbating coastal acidification and deoxygenation (CAD) is well documented, including the East China Sea and the Gulf of Mexico (Cai et al. 2011), the Baltic Sea (Sunda and Cai 2012), the nearshore regions of the North Sea (Provoost et al. 2010), and the Chesapeake Bay (Waldbusser 2011). However, in an Eastern Boundary Upwelling System (EBUS), strong upwelling and vigorous

1 surface currents have been thought to limit the impacts of local anthropogenic nutrient inputs 2 (Chavez and Messié 2009, Capone and Hutchins 2013, Fennel and Testa 2019). Recent 3 evidence from the Southern California Bight (SCB) has challenged this thinking along this highly 4 urbanized coastline. In the SCB, on an annual basis, 92% of terrestrial N flux is wastewater 5 effluent from Publicly Owned Treatment Works (POTWs), most of which is discharged directly 6 into coastal waters via a relatively small number of outfall pipes (Sutula et al. 2020). Modeled outfall N loads are roughly equivalent to those from upwelling in urbanized sections of the 7 8 coast, effectively doubling N loading to the shelf (Howard et al. 2014). These inputs have an 9 immediate, local effect on nitrification rates as high concentrations of wastewater ammonium 10 are rapidly nitrified in the subsurface plumes (McLaughlin et al. 2017). More broadly, 11 observational evidence has shown an increase in the extent of algal blooms in the SCB over the 12 last decade, with chronic blooms documented in areas co-located with major inputs of 13 anthropogenic nutrients (Schnetzer et al. 2007a, Nezlin et al. 2012, Schnetzer et al. 2013). 14 Analysis of a decade of quarterly ocean surveys across the central and northern SCB have 15 shown a significant decrease in dissolved oxygen (DO) concentrations (Bograd et al. 2008), and 16 the rate of decline in the nearshore, proximal to treated wastewater effluent outfalls, has been 17 faster than offshore regions (Booth et al. 2014). However, these regional-scale observations 18 can also be influenced by climate change and Pacific Basin-scale drivers (Booth et al. 2014, 19 Nezlin et al. 2018) and ambient chlorophyll a, DO and pH have not been specifically linked to 20 an effect of wastewater outfalls on biogeochemical rates of primary production and 21 respiration – key processes through which CAD are manifested.

22 In this paper, we quantified nutrient concentrations, forms, and rates of carbon and 23 nitrogen cycling that ultimately influence processes leading to eutrophication and CAD both 24 within wastewater plumes and in areas spatially distant from plumes. We also explored the 25 environmental factors that may be influencing rates, including seasonal and interannual 26 variability linked to the influence of upwelling. Finally, we employed stable isotope tracers to 27 further track relationships and patterns in N and C biogeochemical cycling in these coastal 28 zones as more integrated measurements to compare to the instantaneous rate 29 measurements. We focused on the effect of wastewater N because the SCB is largely N-limited

1 (Thomas et al. 1974, Cullen and Eppley 1981) and wastewater point source discharges 2 dominate anthropogenic sources (Howard et al. 2014) and contain high concentrations of 3 nitrogenous species. As a core part of our study design, observations of ambient ocean state 4 and rates were made in two types of regions defined by the probability of plume impact: 1) 5 "nearfield" regions, defined as the region of active effluent plume mixing and dispersion within 6 10 km of the outfall and 2) "farfield" regions, defined as the region where ambient ocean flow and biogeochemical conditions dominate and the effluent plume is expected to exert 7 8 minimum influence on carbon and nitrogen cycling.

9 The value of such observations extends beyond testing of the hypothesis of an effect of 10 wastewater plumes on watercolumn biogeochemical parameters (pH, dissolved oxygen, 11 chlorophyll a) and rates of nutrient and carbon cycling. To support California's climate action 12 strategies, a spatially-explicit, 3-dimensional numerical ocean model of the SCB was developed 13 and is being applied to examine the relative effects of climate change, natural climate cycles, 14 and local terrestrial and atmospheric carbon and nutrient inputs (Deutsch et al. in review, 15 Kessouri et al. in press). However, in order to support conversations on the utility of nutrient 16 management as a climate change mitigation strategy, such models must be carefully validated 17 against observations of biogeochemical state and rate data. Inadequate numerical model 18 validation has been identified as a significant barrier to effective, evidence-based solutions to 19 coastal eutrophication (Boesch 2019). Models must not only successfully reproduce observed 20 state data, they must also successfully reproduce observed biogeochemical rate data, ensuring 21 that the model is predicting the appropriate state variables for the right reasons. Thus, these 22 observations are part of a coupled observational-numerical modeling approach characterizing 23 the effect of local nutrient and carbon inputs on SCB coastal habitats, laying the foundation for 24 evidence-based solutions for CAD.

25

26 Materials and Methods

27 Study Region

28 The Southern California Bight (SCB) is the bend in the coastline between Point

29 Conception (~34° 34'N) and the U.S.-Mexico International Border (~32° 32'N). It is situated in

the California Current System on the U.S. Pacific Coast. As an EBUS, the SCB is a biologically 1 2 productive region of high economic and ecological importance. Seasonal spring upwelling of 3 nutrient-rich deep water maintains high rates of biological productivity over broad scales. At 4 the same time, upwelling draws water masses that are low in DO, pH, and carbonate 5 saturation state (Ω_{Ar}) onto the shelf and into the photic zone (Sutton et al. 2017). Southern 6 California has a Mediterranean climate with surface runoff confined mostly to the winter, rainy 7 season. The SCB is home to one of the most densely populated coastal regions in North 8 America, where the discharges of treated (advanced primary or secondary) wastewater from a 9 population of 20 million people are released to the coastal zone via ocean outfalls, along with 10 the urban and agricultural runoff from 72 rivers (Sutula et al. 2020). Modeling efforts have 11 shown that these nutrient sources rival natural upwelling in magnitude (Howard et al., 2014), 12 roughly doubling available N to nearshore coastal waters on an annual basis.

13

14 Study Design

15 The study was designed to characterize effects of wastewater plumes on state and rate 16 variables in regions nearfield (within-plume) and farfield (spatially distant) from ocean outfalls 17 where treated wastewater effluent is discharged to the highly urbanized, central portion of the 18 SCB (Figure 1). We hypothesized that chemical and biological rates and water chemistry would 19 differ between areas near ocean outfalls (nearfield), where treated wastewater plume was 20 detected, compared to farfield, "reference" areas, where the plume could not be detected. 21 Selection of nearfield and farfield locations was informed by previous studies that used 22 colored dissolved organic matter (CDOM) as a tracer of effluent plume (Rogowski et al. 2013, 23 Nezlin et al. 2020); as a rule of thumb, nearfield stations were generally within 10 km of the 24 outfalls, while farfield stations were greater than 10 km away.

We compared water column nutrient profiles and stable isotopic compositions, rates of nitrification, primary production and respiration in nearfield and farfield regions, as well as relationships between parameters and rates and seasonal and interannual differences thereof. Four sub-regions were sampled, including two nearfield sub-regions near Orange County Sanitation District's (OCSD) and Los Angeles County Sanitation District's (LACSD) ocean outfalls,

1 a nearshore, farfield sub-region off the coast of northern San Diego County (Camp Pendleton), 2 and an offshore, farfield sub-region off the coast of Los Angeles and Orange Counties (Table 1 3 and Figure 1). Two stations were sampled in each sub-region for the full suite of 4 measurements and rate analyses. For the nearfield regions, sampling was dynamic and based 5 on currents to enable sampling within the plume. A fixed sampling station was located directly 6 over the outfall pipe and a second site, which varied according to currents, was located a short 7 distance from the outfall, but still within the plume. One additional fixed station was sampled 8 in each effluent-plume sub-region for water column nutrients and stable isotope analysis only 9 (no rate data). For the farfield coastal subregion, one station was located on the continental 10 shelf and the other on the shelf-break. For the offshore farfield subregion, one was located at 11 the San Pedro Ocean Time Series (SPOTS) station (offshore of LACSD's outfall) and the other at 12 the California Cooperative Oceanic Fisheries Investigations (CalCOFI) sample station 90.30 13 (offshore of OCSD's outfall), both within the San Pedro Basin.

Comparisons and relationships were investigated seasonally, both when wastewater discharge was anticipated to constitute a minor fraction of the total N pool (Spring upwelling period) and a major fraction (late Summer, when the water column is stratified). Four seasons were collected over two years: late Summer stratification in 2014 and 2015, and Spring upwelling in 2015 and 2016. Cruises and rate measurements were conducted over a four-week period in each season; one sub-region sampled per week.

20

21 Sample Collection

22 Vertical profiles were collected at each station using a Sea-Bird Electronics, Inc., 23 SBE911plus (24 Hz) or SBE-25 (8 Hz) conductivity-temperature-depth (CTD) system, measuring 24 a suite of oceanographic properties including pressure, temperature, salinity, colored dissolved 25 organic matter (CDOM), dissolved oxygen, pH, and chlorophyll fluorescence and transmissivity. 26 In addition to the experimental stations, CTD casts were also taken at several stations in the 27 nearfield sub-regions to map the location of the plume and regional hydrodynamics at the 28 time of sampling (Figure SI.1). The profiles were averaged to 1 m bins. Regional climatologies 29 of temperature, salinity, dissolved oxygen, chlorophyll a, and CDOM were generated by

interpolation using the Barnes algorithm (Koch et al. 1983) from the R package "oce" (Kelley
 and Richards 2017).

3 Discrete samples were collected from Niskin bottles (1.5 and 3 L) on a rosette deployed 4 with the CTD sensor package. Sampling was adaptive at each site, with sample depths 5 determined from downcast CTD observations to identify key oceanographic features at each 6 station (CDOM to mark the plume and chlorophyll *a* maxima). Generally, samples were collected from 5 depths at the stations > 50 m depth: the surface, subsurface chlorophyll 7 8 maximum layer (depth of highest recorded chlorophyll fluorescence), the center of the 9 thermocline (mixed layer, where the temperature gradient exceeds -0.3 °C m⁻¹), below the 10 thermocline, and at the "bottom" (2-5 m from the sediment surface or the end of the rosette 11 cable, which was either 200 m [OCSD, LACSD] or 300 m [Camp Pendleton]). At shallower 12 stations, four samples were collected from the surface, subsurface chlorophyll maximum, the 13 thermocline, and the bottom. For the nearfield sub-regions, the station over the outfall 14 remained fixed throughout the study, but the "off-outfall/second nearfield" station varied 15 based on currents and was identified using CTD profiles of CDOM from several stations (Figure 16 SI.1). CDOM has been shown to be a reasonable tracer for subsurface effluent plumes 17 (Rogowski et al. 2012, Rogowski et al. 2013, Nezlin et al. 2020).

Field water samples were transferred from the Niskin bottles using acid-washed Tygon tubing into acid-washed 2 L high density polyethylene (HDPE) bottles that were triple rinsed with sample water before filling. Both LACSD and OCSD also supplied samples of effluent for analysis prior to each sampling event in their respective nearfield subregions. These samples were collected as a composite over 24 hours in an acid washed 1 L HDPE bottle. Effluent was stored at 4 °C while the composite was generated.

Sub-samples from both field water and effluent were collected for the suite of dissolved inorganic nutrients (nitrate+nitrite, nitrite, ammonium, and ortho-phosphate), total dissolved nitrogen and phosphorus (TDN, TDP), and dual nitrogen and oxygen isotope ratios for dissolved nitrate+nitrite. In addition, field water was sub-sampled for particulate organic matter concentrations (particulate nitrogen, particulate phosphorus, and particulate organic carbon), stable isotope ratios of carbon and nitrogen of particulate organic matter, and

chlorophyll *a*. Subsamples were frozen immediately. Suspended particulate samples were
collected by vacuum filtration onto pre-combusted (450 °C for 4 hours) glass fiber filters
(Whatman GF/F). Suspended particulate samples were sub-sampled from the whole water
sample and stable isotope analyses and were filtered onboard the cruise and frozen
immediately. Filters were collected into snap-close petri dishes and stored in Ziploc bags on ice
in the dark for transport. Filters in the petri dishes were dried at 50 °C in the dark until
analysis.

8 Sub-samples for nutrient and nitrate stable isotope analysis were hand-filtered through 9 a 0.45 µm polycarbonate filter (Millipore) and collected in triple-rinsed, 60 mL HDPE amber 10 bottles, stored on ice for transport to the laboratory, and frozen until analysis (Wankel et al. 11 2006, Wankel et al. 2007, Santoro et al. 2010). Whole water for primary production and 12 nitrification rate incubations were collected directly into acid-washed, triple-rinsed, 2 L HDPE 13 bottles and stored on ice in the dark until the incubations could begin in the laboratory within 14 6 hours of collection. Whole water for respiration rates was collected into 300 mL borosilicate 15 glass biological oxygen demand bottles that were overfilled with twice the volume of the 16 bottle. Water samples for the rate experiments (primary production, respiration, and 17 nitrification) were stored on ice and in the dark until they were brought back to the lab for 18 experimental preparation.

19

20 Laboratory Analyses

21 Nutrient concentrations. Discrete samples were analyzed for a suite of dissolved 22 nutrients. Nitrate + nitrite, nitrite, soluble reactive phosphate, and ammonium were analyzed 23 using flow injection analyses (FIA, Lachat Instruments, QuikChem 8000 at the Marine Science 24 Institute, at the University of California, Santa Barbara), ammonium was also measured 25 following the protocols of Holmes et al. (1999), total N (TN) and phosphorus (TP) samples were 26 analyzed following persulfate digestion (Patton and Kryskalla 2003) using FIA. A relative assessment of nutrient limitation, N*, was calculated, which represents the deviation in 27 28 "Redfield" N:P stoichiometry due to additional sources and sinks of nitrate (Deutsch et al. 29 2001):

$$N^* = [NO_3^{-1}] - (16^* [PO_4^{3-1}]) + 2.9$$
 (Eq 1).

Positive N* values reflect regions with a source of nitrate (via nitrogen fixation) and
negative N* reflects a sink of nitrate (due to denitrification). Values near zero are consistent
with "Redfieldian" assimilation and nitrification of organic matter or that source and loss terms
are balanced (Gruber and Sarmiento 1997).

6 Stable Isotope Analyses. The stable isotopic compositions of dissolved nitrate and nitrite ($\delta^{15}N_{NO2+NO3}$ and $\delta^{18}O_{NO2+NO3}$), ammonium ($\delta^{15}N_{NH4}$), and suspended particulate matter 7 $(\delta^{15}N_{PN} \text{ and } \delta^{13}C_{PN})$ are natural tracers of N sources and cycling in the ocean. Variation in the 8 9 isotopic composition is attributable to distinct source signatures and the mass dependent 10 isotopic discriminations associated with various biogeochemical transformations that 11 constitute the marine N cycle. Because each pathway causes a characteristic shift in isotope 12 composition of the products and reactants, the isotopic composition of the dissolved and 13 particulate pools can provide useful information on the mechanism of these transformations 14 (Sigman et al. 2005, Wankel et al. 2007, Sugimoto et al. 2009).

The preparation and isotope analysis ($\delta^{15}N_{NO2+NO3}$, $\delta^{18}O_{NO2+NO3}$) of dissolved 15 16 nitrate+nitrite in discrete water samples was performed using a bacterial denitrification assay 17 (Sigman et al. 2001, Casciotti et al. 2002). Samples with at least 0.002 mg/kg as N were 18 analyzed by bacterial conversion of nitrate to nitrous oxide and sub-sequent measurement on 19 a continuous flow isotope ratio mass spectrometer (Sigman et al., 2001; Casciotti et al., 2002; Coplen et al., 2007). Isotope ratios of ¹⁵N/¹⁴N and ¹⁸O/¹⁶O were measured using a 20 21 ThermoFinnigan GasBench + PreCon trace gas concentration system interfaced to a 22 ThermoScientific Delta V Plus isotope-ratio mass spectrometer at the Stable Isotope 23 Laboratory at the University of California, Riverside. Dissolved ammonium was extracted from 24 250 mL of wastewater effluent onto glass fiber filter "traps" (Holmes et al. 1998, Hannon and Böhlke 2008) and the isotope ratios of ¹⁵N/¹⁴N were measured using a coupled Costech 25 Elemental Analyzer with a Finnigan Delta Plus Advantage in Continuous Flow Mode at the 26 27 Stable Isotope Laboratory at the University of California, Riverside. The isotope ratios of ¹⁵N/¹⁴N and ¹³C/¹²C from suspended particulate matter collected on precombusted Whatman 28 29 GF/F were measured using a coupled Costech Elemental Analyzer with a Finnigan Delta Plus

Advantage in Continuous Flow Mode at the Stable Isotope Laboratory at the University of 1 2 California, Riverside. Isotope ratios are reported relative to standards: N₂ in air for δ^{15} N, 3 Vienna Standard Mean Ocean Water (VSMOW) for δ^{18} O, and Vienna Pee Dee Belemnite (VPDB) for δ^{13} C. The standard deviation of replicate standards for particulate material δ^{13} C 4 was 0.084 ‰ and δ^{15} N was 0.165 ‰, and the relative percent difference between measured 5 6 standards and reference values was 0.32% and 0.88%, respectively. The standard deviation of replicate standards for dissolved nitrate δ^{18} O was 0.17 ‰ and δ^{15} N was 0.15 ‰, and the 7 8 relative percent difference between measured standards and reference values was 0.46% and 9 0.04%, respectively.

10

11 Rate Measurements

12 Primary production rate measurements. Short-term incubations of natural plankton 13 communities were conducted to determine rates of primary production using radioactive ¹⁴C labeled compounds, expressed as mg C m⁻³ day⁻¹ or integrated vertically to units of mg C m⁻² 14 day⁻¹. Primary production was assessed at multiple depths bracketing the euphotic zone, 15 16 collecting four depths from light levels from 95% to 1%: including the surface, the mixed layer, 17 the chlorophyl maximum layer, and the 1% light level. Primary production was estimated from 18 ¹⁴C uptake using a simulated *in situ* technique in which the assimilation of dissolved inorganic 19 carbon by phytoplankton yields a measure of the rate of photosynthetic primary production in 20 the euphotic zone (Anderson et al. 2006, Brzezinski and Washburn, 2011). Water was collected from Niskin bottles into 2 L HDPE bottles wrapped in black electrical tape and kept in darkened 21 22 coolers until transported to the laboratory. Depths included: surface, the chlorophyll 23 maximum, the center of the thermocline, and the 1% light level (three times the Secchi depth). Subsamples from each depth were separated into two acid-cleaned 250 mL polycarbonate 24 25 bottles; one bottle was wrapped in black electrical tape to measure dark ¹⁴C uptake and the 26 other was left open to light but darkened to expected ambient light levels using a neutral 27 density screen. Bottles were incubated for 24 hours after inoculation with ¹⁴C sodium 28 bicarbonate in incubators placed outside in ambient light in the Southern California Coastal 29 Water Research Project facility, held at recorded in situ seawater temperatures by circulating

1 incubator water through chillers. At the end of incubations total radioactivity in each sample 2 was determined by adding 250 μ L of incubated seawater to 250 μ L of β -phenethylamine in a 3 20 mL glass scintillation vial followed by 10 mL of Ultima Gold XR scintillation cocktail. Each vial 4 was shaken vigorously for 30 s and ¹⁴C activity assayed after bubbles had cleared and 5 chemoluminescence had subsided (~2 hours) by a liquid scintillation counter using an internal 6 quench curve. A 50 mL subsample of incubated water was also filtered through a precombusted 25 mm Whatman GF/F filter. Filters were placed in individual 20 mL glass 7 8 scintillation vials and 0.25 mL of 0.5 N HCl was pipetted onto each filter in a fume hood to 9 drive off excess tracer. Vials were left uncapped for a minimum of 7 hours, and 10 mL of 10 Ultima Gold XR scintillation cocktail was added to each vial, the vials capped and shaken, and 11 ¹⁴C activity was analyzed by scintillation counting on a scintillation counter using an internal 12 quench curve. Primary productivity rates in both the light and dark incubations were 13 calculated as described by Anderson et al. (2006) and Brzezinski and Washburn (2011). Daily primary production (g C m⁻³ d⁻¹) was calculated as the difference in productivity between light 14 15 and dark bottles. Productivity-to-biomass ratio was calculated as the daily primary productivity (g C m⁻³ d⁻¹) per unit chlorophyll a (g chl a m⁻³) with units of gC (g chl a)⁻¹ d⁻¹. Integrated rates of 16 17 productivity were calculated using the trapezoidal method.

18 *Respiration rate measurements.* Respiration rates were measured at four depths: the 19 chlorophyll maximum, the middle of the thermocline, below the thermocline, and 2 m above 20 the bottom/end of the CTD cable. Whole water was collected from Niskin bottles into 300mL 21 glass bottles, each of which was wrapped in black electrical tape, using acid-cleaned Tygon 22 tubing, overfilling each bottle with at least twice the volume. Three bottles were collected for 23 respiration rate measurements at each depth. An additional 50 mL syringe of whole water was 24 collected from each Niskin bottle and held in a cooler. An initial dissolved oxygen 25 concentration (mg O L^{-1}) and percent oxygen saturation was recorded in the field for each 26 bottle using a YSI ProDO optical dissolved oxygen probe, immediately after bottles were filled. 27 Because the probe displaced some water from each bottle, water was replaced with whole 28 water from the syringe to eliminate headspace in each bottle. Bottles were then stoppered 29 and placed in refrigerated units held at in situ water temperatures. Bottles were incubated for 12

24-48 hours and a final dissolved oxygen concentration (mg L⁻¹) and percent oxygen saturation
 in each bottle was recorded. Respiration rates were calculated for each depth at in situ
 temperatures as the difference between time zero oxygen concentrations and final incubated
 oxygen concentrations.

5 *Nitrification rate measurements:* Nitrification is the sequential oxidation of NH₄⁺ to 6 NO_3^- via NO_2^- . Nitrification rates were determined by measuring the accumulation of ¹⁵N in the dissolved nitrate pool following addition of isotopically-enriched ammonium to bottle 7 8 incubations (Santoro et al. 2010). Water was collected from Niskin bottles into 2 L HDPE 9 bottles at four depths: the chlorophyll maximum layer, the middle of the thermocline, below 10 the thermocline, and 2 m above the bottom/end of the CTD cable. Subsamples from this initial 11 sample were separated into three 500 mL acid-washed polycarbonate bottles wrapped in black tape. An enriched (99%) tracer of ¹⁵N-ammonium chloride was added to a final 12 13 concentration of 100 nM to two of the bottles and a third bottle without the tracer served as a 14 control. Not knowing ammonium concentrations beforehand, we targeted this concentration 15 to minimize impact on nitrogen cycling within the bottles. The range of percentage labeled 16 ammonium from 0.01 to 1.5%, with a median value of 0.1%. Bottles were incubated in the 17 dark to minimize N uptake by phytoplankton and as close to in situ temperature conditions as 18 possible (within $\pm 2^{\circ}$ C) in a series of refrigerated incubator units. For reference, Q₁₀ values for 19 nitrification are on the order of 2-3 (Henriksen 1988). Subsamples of 50 mL each were 20 collected at four-time points (approximately 0, 12, 24, and 36 hours post spike addition), syringe filtered through 0.45 µm filters and frozen until analysis for dissolved nitrate 21 concentration and the isotopic composition of nitrate+nitrite ($\delta^{15}N_{NO2+NO3}$) as described above. 22 Potential nitrification rates were determined by modeling the ¹⁵N and ¹⁴N contents of the 23 24 combined nitrate and nitrite pool with inputs from the labeled ammonium pool and outputs 25 through nitrate and nitrite uptake as described in Santoro et al. (2010). Data fitting for the ¹⁵N 26 and ¹⁴N values measured at each time point was performed by non-linear least squares 27 regression method using MATLAB 8.2 and Statistics Toolbox 8.3 (The MathWorks, Inc.).

28

1 Statistical Analysis

2 To understand relationships between state variables and rates we employed several 3 statistical techniques: ANOVA, Kruskal Wallis (when ANOVA assumptions failed), Spearman 4 Rank, and Random Forest regression. To characterize relationships between variables, we used 5 ANOVA for normally distributed data and Kruskal Wallis and Spearman Rank for nonparametric 6 measures of rank correlation using the R packages (R-core Team, 2019) "Tidyverse" (Wickham et al. 2019) and "Hmisc" (Harrell, 2014). We used Random Forest modeling to characterize 7 8 variable importance in constraining the variance of rate measurements. Random forest 9 modeling, a machine-learning statistical method which combines many classification trees to 10 produce a more accurate classification (Breiman 2001, Cutler et al. 2007) and has been found 11 to be a robust tool for interpreting ecological datasets (Prasad et al. 2006, Cutler et al. 2007). 12 Random forests provide several metrics that aid in interpretation of multivariable datasets. 13 Variable importance can be evaluated based on how much worse the prediction would be if 14 the data for that predictor were permuted randomly. The resulting tables can be used to 15 compare relative importance among predictor variables (Prasad et al. 2006). Random Forests 16 were conducted using the R package "randomForest" (Law and Weiner 2012).

17

18 Results

19 Wastewater Effluent Characterization

20 Throughout the study, the effluent nutrient concentrations, isotope ratios, and 21 discharge rates at each outfall were slightly variable (Table 2). Over the study period, both 22 agencies' effluent discharge rate decreased slightly, a part of a long-term trend related to 23 increasing water use efficiency and reduced water consumption in California. During this same time, LACSD discharged an average of 4.1×10^7 (± 4.5×10^6) grams of inorganic nitrogen per 24 25 day and OCSD an average of 1.5×10^7 (± 3.2×10^6) grams of inorganic nitrogen per day. The 26 characteristics of effluent from OCSD and LACSD were different, likely due to differences in 27 their treatment systems. OCSD wastewater treatment is secondary with advanced 28 nitrification/denitrification (NDN) and LACSD is secondary treatment with no advanced 29 nutrient removal. Ammonium accounted for >99% of LACSD's dissolved inorganic nitrogen

(DIN) load and between 55 and 67% of OCSD's DIN load. OCSD's effluent contains more 1 2 phosphate compared to LACSD, with an average N:P molar ratio of 100:1 for OCSD and 678:1 3 for LACSD (Table 2). Dissolved inorganic carbon and dissolved organic carbon were highly 4 variable in both agencies' effluent throughout the study period. The stable N isotopic 5 composition of ammonium ($\delta^{15}N_{NH4}$) and nitrate+nitrite ($\delta^{15}N_{NO2+NO3}$) were different for both plants. OCSD had relatively consistent $\delta^{15}N_{NH4}$, with values ranging between 8.6 and 9.1 ‰ and 6 $\delta^{15}N_{NO2+NO3}$ with values ranging between 7.5 and 9.1 ∞ . LACSD had $\delta^{15}N_{NH4}$ values ranging 7 8 from 5.1 to 12.4 %. The ammonium isotopic composition in LACSD's effluent was linearly 9 related to the concentration of ammonium in the wastewater (R^2 for LACSD is 0.907, P = 0.049, 10 and R² for OCSD is 0.364, P = 0.505). LACSD also had variable $\delta^{15}N_{NO2+NO3}$ with values ranging 11 between -9.6 and -2.3 ‰.

12

13 Seasonal Hydrography, Chlorophyll, and Dissolved Oxygen

14 There were clear seasonal differences in temperature and salinity throughout the study 15 region (Figures 2, SI.2). Summer surveys were characterized by high surface water 16 temperatures, shallow thermoclines and haloclines (~10-15 m) and large vertical temperature 17 gradients, indicating strong thermal stratification. Spring surveys were characterized by 18 weaker vertical thermal gradients and deeper thermoclines and haloclines (~20-30 m). The 19 wastewater plume was identifiable by both low salinity and high CDOM in both LACSD's and 20 OCSD's nearfield sub-regions at mid-depth. During Spring 2016 there was a cold, high salinity, 21 low DO water mass intruding into the region from depth throughout the Palos Verdes Shelf 22 region, and notably in the OC Offshore station (CALCOFI station 9030).

There were no statistically significant differences between DO in nearfield versus far field areas (Kruskal Wallis). Nearfield stations generally had shallower oxyclines compared to the farfield stations, but that is likely due to close proximity to shore compared to most farfield stations, except for CP1 (Figure 2). DO profiles varied seasonally (Figures 2, SI.2). In the Summer, all regions had similar profiles and a narrower range of values in the upper 60 m. During the Spring surveys, DO had a larger range of values in the upper 60 meters, with surface ocean values similar to Summer, but much lower at depth. Spring also had greater variability in

the depth to the oxycline among different stations. The lowest DO was in deeper waters and
shoaled closer to the surface during Spring (particularly Spring 2016), indicative of upwelling of
cold, deep waters, low in DO.

4 Chlorophyll fluorescence in nearfield areas was slightly, but significantly, higher in 5 nearfield areas compared to farfield within the first 60 m of the water column, which captures 6 the chlorophyll maximum depth at all stations, with a mean chlorophyll *a* concentration of 2 μ g L⁻¹ in the nearfield compared to 1 μ g L⁻¹ in the farfield (P <2e⁻¹⁶). Daytime, chlorophyll 7 8 florescence had a similar vertical structure to DO, with a clear subsurface maximum layer in all 9 seasons (Figure 2). The depth to and magnitude of the subsurface chlorophyll maximum was 10 variable among stations and among seasons. The lowest chlorophyll was during Summer 2015 11 and the highest during Spring 2015. The highest values during Spring 2015 were associated 12 with the LACSD and OCSD nearfield areas, appearing as a thin subsurface chlorophyll layer 13 (Figure 2). Spring 2016 had a shallow, narrow chlorophyll layer, with no chlorophyll 14 fluorescence associated with the deep, salty water mass present in Spring 2016. Summer 2014 15 had middling water column chlorophyll that was deeper offshore and shallower nearshore, 16 whereas Summer 2015 had relatively low chlorophyll overall with the most consistency among 17 stations.

Sampling occurred over a four-week period for each season and this may have
introduced some variability among stations; however, there was no clear pattern within any
sampling season of the nearfield stations having consistently different hydrographic conditions
relative to the farfield stations that might create bias in the dataset (Figure SI.2).

22

23 Water Column Nutrient Concentrations

Dissolved Inorganic Nutrients. Despite the continuous N-load from both wastewater
 treatment plants, surface dissolved inorganic nitrogen (DIN) concentrations were generally low
 in all subregions (Figure 3), but not completely depleted. Surface DIN values ranged from not detected to 26.9 μM with a mean value of 3.6 μM in farfield areas and 4.7 μM in nearfield
 areas. Nutrient profiles generally showed lower concentrations in surface waters with
 increasing concentrations at depth (Figure 3). Mean DIN concentrations for samples collected

1 below the thermocline ranged from 0.6 to 53.9 μ M, with a mean value of 18.7 μ M in farfield 2 areas and 17.0 μM in nearfield areas). Excluding the bottom sample (which was collected at 3 significantly greater depths offshore compared to the nearshore sites), ammonium and nitrite 4 were significantly higher in the nearfield stations compared to farfield, non-plume stations (Pvalues of 0.000542 and 1.30e⁻⁵, respectively). High concentrations of ammonium (maximum 5 value of 18.8 μ M and a mean value of 5 μ M) and nitrite (maximum value of 1.1 μ M and mean 6 7 value of 0.4 μ M) were associated with the plume mixing zone (as detected by CDOM, Figure 8 SI.1) in the nearfield regions, the depth where the plume reaches neutral buoyancy in the 9 water column (between 20 and 60 m depth). Nitrate and phosphate were not significantly 10 different in the nearfield compared to the farfield. All dissolved inorganic nutrients showed significant differences by season ($P = 3.86e^{-6}$ for nitrate, $P = 7.69e^{-05}$ for ammonium, and P =11 $1.03e^{-10}$ for phosphate), except for nitrite (P = 0.15). 12

13

14 Suspended Particulate Nutrients. Water column suspended particulate nutrients had 15 similar profiles, with the highest concentrations at the surface and decreasing with depth 16 (Figure SI.3). Mean surface particulate nitrogen was 3.5 μ M at the surface and 2.5 μ M below 17 the thermocline, mean surface particulate phosphorus was 0.18 μ M at the surface and 0.12 18 μ M below the thermocline, and mean surface particulate carbon was 19.0 μ M and 13.2 μ M 19 below the thermocline. There were no statistically significant differences by station type for 20 particulate nitrogen or phosphate, but particulate carbon was slightly, significantly higher in 21 nearfield stations (Kruskal Wallis, P = 0.03), particularly those within the plume mixing zone 22 detected by CDOM (with particulate carbon concentrations 21 μ M for depths within the plume 23 in nearfield stations compared to 16.9 μ M for similar depths in farfield stations). There were 24 also significant differences by season for all particulate parameters (Kruskal Wallis, P = 0.0008, 25 0.02, 2.2e⁻¹⁶, for PC, PN and PP respectively). Spring 2016 was slightly anomalous from the 26 other three seasons, exhibiting higher concentrations in particulate carbon, nitrogen and 27 phosphorus in surface samples relative to the other seasons, perhaps related to the cold, salty 28 water mass intrusion during that season.

1 Nutrient Ratios. Nutrient concentrations for particulate matter and dissolved inorganic 2 nutrients were highly correlated, following the Redfield relationship of C:N:P 106:16:1 (Figure 3 4, Figure SI. 4). The mean N:P ratio for dissolved nutrients was 15.4 and was significantly lower 4 (P = 0.028) in the farfield (mean DIN:P of 13.4) compared to the nearfield (mean DIN:P of 5 17.5). The highest mean DIN:P ratios were in the nearfield stations at depths associated with 6 the plume (mean DIN:P of 18.5). The lowest mean DIN:P ratios were recorded at the chlorophyll maximum depth in the farfield (9.9). N* showed strong seasonality, with water 7 8 column values close to 0 during the Spring upwelling periods, whereas the Summer fluctuated 9 between positive N* values (Summer 2014), indicative of a source of nitrate, and negative N* 10 values (Summer 2015), indicative of a sink for nitrate (Figure 5, Figure SI. 5). N* values near 11 zero suggest that there were no large sources of nitrate (e.g., from nitrogen fixation) or sinks 12 (e.g., due to denitrification).

13 Particulate matter, particulate C and N were highly correlated, with 90% of C:N ratios between 3.8 and 20.0, with a mean regional C:N of 6.7. There were some differences between 14 15 particulate C:N ratios in the nearfield and farfield (with mean values of 7.1 and 6.3, 16 respectively). C:N ratios less than 6 were found throughout the water column and are atypical 17 of phytoplankton and often associated with bacteria (Goldman et al. 1987). C:P and N:P 18 relationships were close to Redfield, with occasional deviations. These deviations were not 19 related to depth (Figure SI.4) and were more closely tied to station type, with nearshore 20 stations (nearfield stations and farfield, coastal stations at Camp Pendleton) having greater 21 variability than farfield, offshore stations. Relatively high particulate P concentrations were not 22 matched by high concentrations in N and C during Spring 2016.

23

24 Water Column Isotope Ratios

25 Stable Isotopic Composition of Dissolved Nitrate. $\delta^{15}N_{NO2+NO3}$ values were generally 26 lower in surface waters (mean of 7.5 ‰), increased to a subsurface maximum (mean of 9.5 27 ‰), before decreasing again (to mean of 8.7 ‰). $\delta^{15}N_{NO2+NO3}$ was significantly higher in the 28 farfield, non-plume stations compared to nearfield stations (P = 0.02). The $\delta^{18}O_{NO2+NO3}$ was 29 highest in surface waters (mean of 14.6 ‰) and decreased with depth (mean of 5.2 ‰) with 1 no obvious maximum or minimum (Figure 6). The shapes of profiles for both $\delta^{15}N_{NO2+NO3}$ and 2 $\delta^{18}O_{NO2+NO3}$ were correlated with the vertical distribution of nitrate, DO, and chlorophyll. There 3 were also significant differences in $\delta^{15}N_{NO2+NO3}$ and $\delta^{18}O_{NO2+NO3}$ values between sampling 4 seasons (P-values of 3.07e⁻¹⁰ and 6.99e⁻⁹, respectively).

At low nitrate concentrations $\delta^{15}N_{NO2+NO3}$ values were variable, but asymptote towards 5 6 values between 7 and 9 ‰ for all sampling periods as the concentration of nitrate, the fraction of nitrate in the DIN pool, and as nitrification rate all increase (Figure 5, Figure SI.5), similar to 7 what has been seen in urban coastal environments (Sugimoto et al. 2009, McLaughlin et al. 8 9 2017). There was no significant relationship between $\delta^{15}N_{NO2+NO3}$ and N^{*}, largely because N^{*} 10 was typically very close to 0 for most sampling stations and events (Figure 5B, Figure SI.5). 11 Stable Isotopic Composition of Suspended Particulate Matter. The stable isotopic composition of suspended particulate organic carbon ($\delta^{13}C_{PM}$) and particulate organic nitrogen 12 $(\delta^{15}N_{PM})$ were lower at the surface (-25.9 ‰ and 4.6 ‰, respectively), increased to a 13 14 subsurface maximum (-25.5 ‰ and 5.4 ‰, respectively), and decreased at depth (-27.2 ‰ and 4.2 ‰, respectively; Figure 6, Figure SI.6). Both $\delta^{13}C_{PM}$ and $\delta^{15}N_{PM}$ were significantly higher in 15 the nearfield areas relative to the farfield ($P = 1.68e^{-06}$ and 0.00107, respectively). There were 16 also significant temporal differences in $\delta^{13}C_{PM}$ and $\delta^{15}N_{PM}$ (P = < 2e⁻¹⁶ and P = 8.22e⁻⁰⁶, 17 respectively). $\delta^{13}C_{PM}$ in Summer 2014 (mean value of -26.7 ‰) and Spring 2015 (-28.9 ‰) were 18 lower than Summer 2015 (-26.0 %) and Spring 2016 (-23.8 %) and $\delta^{15}N_{PM}$ was highest in 19 20 Spring 2015 (mean 5.7 ‰), and lowest in Spring 2016 (3.4 ‰), with both summer periods having similar mean values, Summer 2014 (4.4 ‰) and Summer 2015 (4.9 ‰) (Figure 6, Figure 21 SI.6). $\delta^{15}N_{PM}$ showed an asymptote to values between 4 and 10 ‰ with increasing particulate 22 N concentration and chlorophyll *a* concentration (Figure 5I, Figure SI.7). By contrast, no 23 significant relationship between $\delta^{15}N_{PM}$ and particulate N nor $\delta^{15}N_{NO2+NO3}$ was observed (Figure 24 25 5G,H, Figure SI.7). $\delta^{13}C_{PM}$ was also positively correlated with $\delta^{15}N_{PM}$ for most stations and seasons and showed an asymptote between -26 and -20 ‰ with increasing particulate carbon 26 concentration and chlorophyll a (Figure 5F, Figure SI.8). $\delta^{13}C_{PM}$ was also negatively correlated 27 28 with water column DIN (Figure SI.8).

1 Primary Production and Respiration

2 There was large spatial/temporal variability in both primary production and respiration 3 rates (Figure 7A and 7B, Figure SI.9). Primary production rates ranged from 129 to 2842 mg C m⁻² d⁻¹ and respiration rates ranged from 80.55 to 1521 mg O m⁻³ d⁻¹. There was seasonality in 4 5 primary production for the two farfield offshore stations, with higher production during the 6 Spring, due to upwelling. However, for the nearshore stations (nearfield stations and the farfield, coastal stations at Camp Pendleton), there were no statistically significant differences 7 8 between station type (nearfield vs farfield) or season in primary production. Similarly, 9 respiration rates offshore were typically higher than nearshore values, but there were no 10 significant differences between the nearfield stations and farfield, coastal stations (Figure 7B). 11 Primary productivity values were similar to what has been reported for the region (Smith et al. 12 1982, Eppley 1992) with similar seasonality (Mantyla et al. 2008). Respiration rates are within 13 the range of other coastal areas and slightly higher than what has been reported for the Pacific 14 coastal environment (Robinson 2019). There were no significant differences between the 15 LACSD nearfield and the OCSD nearfield (Figure SI.9). There was no significant relationship 16 between primary production and respiration for any season (Figure SI.10). It should be noted 17 that one integrated rate for both primary production and respiration was provided per station, 18 so sample sizes were small (n = 12 nearfield and farfield sites, 24 total primary production rate 19 estimates; n = 16 nearfield and farfield sites, 32 total respiration rate estimates).

20

21 Nitrification

22 Nitrification rates were highly variable throughout the study ranging from 0.001 to 325 23 nmol L⁻¹ day⁻¹, with no clear seasonal patterns. Rates were significantly higher in samples 24 collected below the thermocline (mean rate of 40.0 nmol L⁻¹ day⁻¹) compared to samples 25 collected in the mixed layer (16.5 nmol L⁻¹ day⁻¹) and deep chlorophyll *a* maximum (24.6 nmol L^{-1} day⁻¹) (P = 0.01; Figure 7C), similar to profiles measured by others (Ward 1987, 2005, 26 27 Santoro et al. 2010, Smith et al. 2014) and attributed to light inhibition of nitrification in 28 surface waters. Rates were highest in nearshore stations (mean rate of 35.2 nmol L⁻¹ day⁻¹ in 29 the nearfield stations and the farfield, coastal stations at Camp Pendleton) compared to the

two offshore stations (mean rate of 21.1 nmol L⁻¹ day⁻¹ offshore) and were significantly higher
in nearfield areas (36.2 nmol L⁻¹ day⁻¹) relative to farfield areas (23.6 nmol L⁻¹ day⁻¹) (P = 0.04).
There were no significant differences between LACSD and OCSD nearfield stations (Figure SI.9).

5 Discussion

6 Anthropogenic nutrient discharges into coastal waters can drive significant 7 biogeochemical changes and local managers are being urged to consider whether nutrient 8 management strategies can slow the progression of acidification and deoxygenation and their 9 impact on coastal habitats (Kelly et al. 2011, Strong et al. 2014, Chan et al. 2016). Globally, 10 wastewater discharge, 80% of which is untreated (WWAP 2017), represents a significant 11 pathway of nutrient enrichment and eutrophication of coastal waters. However, the relative 12 impact of these inputs compared to global change on N and C biogeochemical cycling is poorly 13 characterized (Kelly et al. 2011, Strong et al. 2014), particularly in EBUS. In this study, we 14 document that wastewater nutrient inputs may have an immediate, local effect on nutrient 15 stoichiometry, elevating ammonium and nitrite concentrations and increasing dissolved 16 nitrogen: phosphorus ratios, as well as increasing rates of nitrification within the plume and 17 slightly increasing chlorophyll a concentrations. We did not observe a consistent, near plume 18 effect on primary production, respiration, or DO, suggesting any potential impact from 19 wastewater on these processes might be moderated by regional factors, notably mixing of 20 water. Furthermore, a regional assessment of aragonite saturation state conducted at the 21 same time as this study, indicated that there was no clear pattern in reduced Ω_{Ar} near ocean 22 outfalls (McLaughlin et al. 2018). This suggests that further study of implications of these 23 changes on the SCB lower trophic ecosystem through ocean numerical modeling studies is 24 warranted, given the difficulty in disentangling local versus regional versus global drivers 25 through observations alone.

26

2

Anthropogenic nutrient inputs impact the biogeochemical cycling of nitrogen in the immediate, local vicinity of outfalls.

The doubling of coastal ocean annual nitrogen loads from wastewater outfalls (Howard et al. 2014) has important implications for nitrogen cycling in the nearshore. Wastewater nitrogen appears to be altering the composition of the N pool within the plumes; ammonium and nitrite concentrations were elevated in the nearfield regions compared to farfield regions (Figure 3). Furthermore, there was no seasonal difference in nitrite concentration, suggesting a source of nitrite independent of seasonal upwelling.

9 The N:P ratio in coastal waters also appears to be altered by the presence of 10 wastewater plumes. The ratio of N:P is a nearly constant 16:1 throughout the world's oceans, 11 in both plankton biomass and in dissolved nutrient pools (Redfield 1958), and the farfield 12 stations adhere closely to this ratio (Figure 4). However, the nearfield stations had elevated 13 N:P ratios, which can be attributed to the relatively low concentrations of P compared to N in wastewater (Table 2, effluent N:P ~115:1). This increase in P-limitation near outfalls may have 14 15 important implications for the planktonic community compositions near these discharges 16 (Grosse et al. 2017, Moreno and Martiny 2018, Fagan et al. 2019). For example, harmful algal 17 bloom species (HABs), as P-limitation has been linked to increased toxin production in *Pseudo*-18 nitzschia (Fehling et al. 2004), a common HABs species in the SCB (Schnetzer et al. 2007b).

19 Nitrification of ammonium to nitrate has been found to play an important role in 20 coastal ocean nitrogen cycling (Ward 1987, 2005) and can support a significant fraction of 21 productivity in surface waters (Wankel et al. 2007, Santoro et al. 2010). Nitrification rates were 22 significantly higher in nearfield stations compared to farfield areas, suggesting the ammonia 23 from wastewaters may be increasing these rates within the plumes. Though it should be noted 24 that the high concentrations of ammonium in the nearfield and the relatively small percentage 25 of labeled ammonium added to these samples during the incubation experiments (less than 26 1% spike) may have resulted in an underestimation of nitrification in the nearfield samples. 27 Therefore, the differences in nitrification rates in the nearfield compared to the farfield may be greater than what is described here. Offshore (farfield) rates were consistent with those 28 29 measured in similar locations in the Southern California Bight (Ward 1987) and coastal

1 locations in Monterey Bay, California (Ward 2005, Smith et al. 2014), but slightly lower than 2 those measured within the California Current (Santoro et al. 2010). The rate of nitrification is 3 light inhibited and related to ammonium concentration (Ward 2008). Our results are 4 consistent with this, where nitrification rates show positive correlations with water column N 5 species and negative correlations with DO, temperature (Spearman rank analysis, Table 3). 6 Random forest regressions explained a relatively low percent of the variance for nitrification 7 rates, 20% (Figure SI.10). Phosphate and station type, were most predictive, followed by 8 temperature, nitrate and dissolved oxygen, likely related to the fact that nitrifying bacteria are 9 typically light inhibited and thus more abundant below the eutrophic zone, where water 10 temperatures were colder and dissolved oxygen is lower (Ward 1987, 2005, Santoro et al. 11 2010, Smith et al. 2014). The low percentage of the variability explained by the random forest 12 models suggests that there were other factors which affect the nitrification rates, such as the 13 composition of the bacterial community (Ward 2005) or possibly the added uncertainty of the 14 underestimation of the rates due to insufficient spike as described above.

15 Wastewater N is predominantly ammonium and is discharged at depth, rising in a 16 buoyant plume and generally trapped below the mixed layer (Figure 3) (Nezlin et al. 2020). 17 Because wastewater ammonium is discharged at depth below the photic zone, wastewater 18 ammonium was expected to be rapidly nitrified. Indeed, elevated nitrite concentrations and 19 higher nitrification rates were associated with OCSD's effluent plume (McLaughlin et al. 2017), 20 and effluent discharges in other coastal regions have seen similarly elevated nitrification rates 21 in sediments near outfalls (Axelrad et al. 1981, Nowicki 1994). Furthermore, the OCSD 22 diversion study was able to track changes in nitrification rates when the plume is "turned-off" 23 relative to "turned-on" and results suggested that wastewater ammonium was nitrified on 24 relatively short time scales; hours to days (McLaughlin et al. 2017). Thus, the contribution of 25 nitrate from the nitrification of wastewater effluent has the potential to support significant 26 productivity in coastal areas. Ammonium concentrations were elevated in nearfield stations 27 relative to farfield stations, particularly at the depths associated with the plume (Figure 3). 28 These high ammonium concentrations were associated with increased nitrification rates, both 29 directly over the outfall and at a distance from the outfall but still within the plume, suggesting

a local impact on N cycling from wastewater effluent discharges. Generally, LACSD's 1 2 nitrification rates were higher than OCSD's, particularly below the thermocline, which is 3 consistent with the higher concentrations of ammonium in effluent (Figure SI.9). However, 4 OCSD had much greater variability in nitrification rates near its outfall compared to LACSD, 5 with relatively high nitrification rates within the chlorophyll maximum and within the mixed 6 layer, as well as the highest recorded nitrification rate in a bottom sample during Spring 2015, demonstrating the variability present in the rates both within a site and across the region. 7 8 Concentrations of nitrite, likely from oxidation of ammonium, were elevated within the plume 9 over both outfalls and, as noted above, did not show any seasonal variability suggesting a 10 continuous source (Figure 3). Nitrate concentrations were not significantly elevated in the 11 nearfield relative to farfield stations; however, the concentrations of nitrate were 2-3 times 12 higher than ammonium and 10 times higher than nitrite, thus the additional contribution from 13 nitrification if diluted over a larger area may not have been distinguishable from local 14 heterogeneity in concentrations. Nitrification rates were not higher in the "fresh" plume 15 compared to "older" plume stations, suggesting that the time-scales of nitrification of plume 16 ammonium were on the order of days and that the elevation in nitrification rates 17 encompasses, at minimum, plume areas as defined by this study (i.e., detectable by CDOM). 18

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Regional factors play a more important role in nitrate assimilation compared to localized impacts of wastewater plume.

21 Nitrate assimilation (incorporation of nitrate into the biomass) was not directly 22 measured in this study; however, stable isotopes were employed as tracers of this process in the water column. The relationship between $\delta^{15}N_{NO2+NO3}$ and $\delta^{18}O_{NO2+NO3}$ was generally 23 24 consistent with nitrate assimilation in the water column and was not significantly different 25 between seasons and station location. The relative enrichments in $\delta^{15}N_{NO2+NO3}$ and $\delta^{18}O_{NO2+NO3}$ generally fell along a line with a slope of 1 (Figure 5C, Figure SI.5), particularly during the 26 Spring 2016 event; however, the low surface values in $\delta^{15}N_{NO2+NO3}$ resulted in poor or negative 27 correlation between $\delta^{15}N_{NO2+NO3}$ and $\delta^{18}O_{NO2+NO3}$ for some stations. A linear relationship 28 between $\delta^{15}N_{NO2+NO3}$ and $\delta^{18}O_{NO2+NO3}$ with a slope of 1 would be predicted if phytoplankton 29

1 assimilation or denitrification were the dominant processes controlling the isotopic 2 composition of nitrate (Granger et al. 2004). However, because concentrations of oxygen were 3 relatively high (always greater than 2 mg L⁻¹), denitrification is not likely to be a significant 4 process in these waters and the isotope affect is likely to be attributed to assimilation (Wankel 5 et al. 2007). It should be noted that because our dissolved nitrate+nitrite oxygen isotope ratios 6 were measured using the denitrifier method, there may be some errors in the apparent 7 $\delta^{18}O_{NO2+NO3}$ values reported. Casciotti et al. (2007) noted that when nitrite was 2% or more of 8 the total nitrate plus nitrite present in the sample, the denitrifier method would give a 9 detectable error in the apparent $\delta^{18}O_{NO2+NO3}$ in the sample, underestimating the value. This is a 10 concern for 30% of our samples, particularly the nearfield stations (38%) where mean nitrite 11 fraction of the nitrate+nitrite pool is 3.8%, though it also affects 20% of the farfield stations 12 where the mean nitrite fraction of 1.2%. This effect only applies to the oxygen isotope values 13 and, assuming the offset would add 25‰ to the δ^{18} O of nitrite (as estimated by Cassioti et al. 2007) would increase $\delta^{18}O_{NO2+NO3}$ values of samples 0.08 - 13.8‰ with a mean increase of 2.69 14 ‰, which would slightly reduce the spread of samples along the $\delta^{18}O_{NO2+NO3}$ axis of Figure 5C 15 16 (Figure SI.12).

As indicated by patterns in $\delta^{15}N_{\text{NO2+NO3}}$ and $\delta^{18}O_{\text{NO2+NO3}}$, nitrate+nitrite assimilation was 17 18 not significantly different in nearfield areas versus farfield areas, but had clear seasonal 19 patterns, suggesting regional mixing and dilution of wastewater N likely spreads the impact on 20 assimilation over a much wider area thereby creating a potential temporal lag in effect. During 21 upwelling periods, particularly during the Spring of 2016, the effect of assimilation on the 22 isotopic composition of the dissolved nitrate pool is evident from the measurements of 23 $\delta^{18}O_{NO2+NO3}$ and falling along a 1:1 line (Figure 5C, Figure SI.5). However, during the Summer sampling periods, the isotope effect from assimilation of "new" nitrate from upwelling is 24 25 diluted and the relationship between the two isotopes appears more influenced by nitrification (low $\delta^{15}N_{NO2+NO3}$ and high $\delta^{18}O_{NO2+NO3}$). In Monterey Bay, in northern California, 26 27 nitrification of "natural" ammonium from regenerated organic matter was found to contribute 28 ~30% of nitrate based primary production (Wankel et al. 2007). The isotopic composition of 29 dissolved nitrate at all stations increased with nitrate concentration and the fraction of nitrate

in the DIN pool. This is consistent with nitrification as a significant driver defining the
composition of the DIN pool and the isotopic composition thereof, region-wide (Sugimoto et
al. 2009, McLaughlin et al. 2017). However, because nitrification is largely happening below
the euphotic zone, the importance of this source for regional primary productivity will be
directly related to physical mixing in the region. Time scales of when this subsurface nitrate
source mixes into the surface waters could result in impacts of this source in the farfield
regions and not necessarily near the discharge location.

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Impacts of anthropogenic nutrient inputs on the biogeochemical cycling of carbon and oxygen are regional in scale.

11 Regional factors such as upwelling and Pacific Basin-scale changes in circulation have 12 been shown to be the primary drivers for the concentrations of chlorophyll biomass and the 13 depth of the chlorophyll maximum layer (Mantyla et al. 2008, Nezlin et al. 2012, Nezlin et al. 14 2018). This work highlights the importance of regional-scale influences on carbon and oxygen 15 cycling. In this study, no statistically significant differences in instantaneous rates of primary 16 productivity and respiration were found in nearfield areas versus farfield (Figure 7). This is 17 consistent with an investigation of pH and aragonite saturation state during the same time 18 period as this study, which found no significant differences between these parameters on the 19 continental shelf (near anthropogenic discharges) compared to offshore of these influences 20 (McLaughlin et al. 2018). While there were some regional differences in chlorophyll a and 21 dissolved oxygen, and mean chlorophyll fluorescence was increased in proximity to the 22 wastewater plume, there was a high amount of variability among stations and seasons (Figure 23 2, Table 3). This is consistent with other research in the area which has shown that physical 24 mixing processes, such as internal waves which can create thin bands of high productivity 25 along the coast and have a significant effect on primary production over relatively small spatial 26 scales (Lucas et al. 2011, Omand et al. 2011, 2012). The alteration of the N:P ratios in nearfield 27 regions could impact phytoplankton community productivity as noted above (Grosse et al. 28 2017, Moreno and Martiny 2018, Fagan et al. 2019), resulting in a dilution of the impact of 29 increased N over a larger area. Furthermore, we saw evidence of seasonal upwelling bringing

increased chlorophyll biomass and intrusions of deep, cold waters that are low in dissolved
oxygen into surface waters; impacting both nearfield and farfield regions. This supports the
hypothesis that regional mixing of water masses dilutes the impact of wastewater N on the
coastal environment, potentially enhanced by outfall diffuser systems which discharge effluent
over large areas, spreading the effect on primary production and respiration over larger areas,
and by extension carbon and oxygen cycling in the SCB (Mantyla et al. 2008, Nezlin et al. 2012,
Nezlin et al. 2018).

8 Random forest regression characterized a low percent of variance for both primary 9 production and respiration (30% and 18%, respectively Figure SI.11), thus the primary drivers 10 for variability in these processes were either not well accounted for in the observed 11 parameters or the relationships between factors are diluted regionally. Season was the most 12 predictive variable of primary production in random forest analysis, indicating the importance 13 of regional variables on primary productivity, as the SCB is subject to seasonal upwelling. This 14 is shown in the seasonal differences in temperature, salinity and chlorophyll a during the study 15 (Figure 2). Station type was not very predictive for either primary production or respiration, 16 suggesting any impact of the wastewater plume is diluted across the region. This dilution 17 effect is also apparent in the isotopic signature of the assimilation of "new" nitrate on $\delta^{15}N_{NO2+NO3}$ is also mixed throughout the study area, particularly during Summer, stratified, 18 19 periods (Figure 5C). Furthermore, concentrations of suspended particulate matter (carbon, 20 nitrogen and phosphorus) and nutrient ratios within that suspended matter were not 21 significantly different in the nearfield versus the farfield (Figures 4 and 5). This decoupling of 22 the presence of the wastewater plume, and its associated nutrients, from primary production, 23 suspended organic matter, and respiration can be explained by regional dilution of effluent 24 nitrogen due to water mass mixing. Both LACSD and OCSD outfalls have multiport diffusers to 25 disperse and increase dilution of the plume over a large area to minimize immediate local 26 impacts of the discharge on the coastal ocean environment (Koh and Brooks 1975). The 27 relative buoyancy of the plume to local seawater traps the plume in the subsurface near the base of the euphotic zone where nutrients could be utilized by primary producers, but the 28 29 impact is designed to be diluted over a larger area dictated by the local hydrodynamics of

plume mixing as well as seasonal, basin-scale changes in currents and ocean state. Given that
 no difference was observed in primary production and respiration in plume affected areas
 relative to farfield areas, it is likely that effect of effluent nutrients on these processes are
 either diluted throughout the region into farfield areas or is too small to detect.

5 While most of the patterns in carbon and oxygen cycling were regional in scope, there was some evidence of local influence. Isotopic signatures of particulate matter ($\delta^{15}N_{PM}$ and 6 $\delta^{13}C_{PM}$) were both slightly, but significantly higher the nearfield compared to farfield, 7 potentially reflecting a small local influence on carbon and nitrogen cycling (Figure 5). Higher 8 9 $\delta^{15}N_{PM}$ and $\delta^{13}C_{PM}$ have been associated with higher primary productivity in coastal areas 10 (Oczkowski et al. 2014, 2016) and may be a more integrated metric of primary productivity 11 differences than an instantaneous rate measurement. The isotopic composition of nitrogen in suspended material ($\delta^{15}N_{PM}$) is slightly lower than dissolved nitrate in the water column 12 13 $(\delta^{15}N_{NO2+NO3})$ indicating the presence of a small isotopic discrimination associated with uptake of nitrate into the biomass (Figure 5H) (Ostrom et al. 1997). The lighter isotope is preferentially 14 15 utilized by phytoplankton and, as N becomes limiting, this isotopic discrimination becomes less 16 and less until the isotopic composition of the biomass is the same as that of the N being 17 utilized (Ostrom et al. 1997), as such higher values are typically associated with more 18 competition for N and higher primary production (Oczkowski et al. 2014, 2016). Both $\delta^{15}N_{PM}$ and $\delta^{13}C_{PM}$ increased with chlorophyll fluorescence (Figure 5I, F). Using chlorophyll *a* as a proxy 19 for phytoplankton biomass, an increase in $\delta^{13}C_{PM}$ with biomass is consistent with lowering of 20 21 dissolved inorganic carbon and dissolved carbon dioxide (Ostrom et al. 1997). Furthermore, 22 chlorophyll a fluorescence was also slightly significantly higher in the nearfield, which may 23 reflect increased biomass in the nearshore. This has important implications for regional 24 variability in acidification. Nutrient enrichment of coastal waters has been linked to impacts on 25 the aragonite saturation state, inducing waters into undersaturated conditions when they 26 otherwise might not have been (Cai et al. 2011, Wallace et al. 2014, Rheuban et al. 2019). 27 Regionally, the SCB is exposed to waters with aragonite saturations states below thresholds 28 thought to be important for marine calcifers (McLaughlin et al. 2018), and further investigation

into potential enhancement of acidification associated with increased primary production is
 therefore warranted.

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The failings of a "reference-area" approach to document impacts of point sources on nutrient and carbon cycling.

6 The study design was based upon the hypothesis that areas near wastewater outfalls 7 would be more impacted from nutrient discharges than areas spatially distant from the 8 outfalls. Results suggest that such a concept is has limitations. While nitrification, which 9 occurs at depths directly associated with detectable plume, showed significant differences 10 between nearfield and farfield stations, instantaneous rates of primary production and 11 respiration, an immediate, local effect was not observed. While a slight lag in the timing (and 12 thus distance) of elevated primary production and chlorophyll *a* from wastewater N 13 discharged by ocean outfalls at depth was expected, the observations suggest that levels of 14 advection, stirring, and eventual mixing in the region were sufficient to transport 15 anthropogenic nutrients to presumed "minimally disturbed" reference areas in the farfield. 16 Thus, the concept of a "reference area" (Nezlin et al., 2020) as implemented here in the SCB is 17 flawed for nutrient impacts on these processes. This is supported by numerical ocean model 18 simulations from the Regional Ocean Modeling System (ROMS, www.myroms.org; 19 (Shchepetkin and McWilliams 2005)) run in particle tracking mode to track Lagrangian flow of 20 "plume particles" (Figure 8). When particles were released from the outfalls, they were 21 transported to farfield areas used for this study on timescales of weeks to months, suggesting 22 that wastewater N is mixed over larger regional scales. In such cases, impacts of 23 anthropogenic nutrients on primary production, related elevation in respiration, and impacts 24 on carbon cycling/acidification, are not easily deciphered with observational data alone. Thus, 25 the monitoring scheme to assess nutrient impacts should be reevaluated to include modeling 26 approaches that can account for the complexity of mixing and farfield transport in the region. 27

1 Conclusion

2 In this study, we found that wastewater nutrient inputs have an immediate, local effect 3 on nutrient stoichiometry and nitrogen concentrations and elevated rates of nitrification 4 within the plume. Impact of wastewater plumes on nitrate assimilation, primary production, 5 chlorophyll a, respiration, and DO was moderated by regional mixing of water masses. This 6 lends strong support for further study of local anthropogenic forcing through ocean numerical 7 modeling studies, given the difficulty of disentangling local to global scale drivers through 8 observations alone (Chan et al. 2016). While observational studies such as this are costly, they 9 provide key data needed to evaluate ocean numerical models. These data should not only 10 include state variables, which can be used to characterize the uncertainty in model output (is 11 the model getting the right answers), but also rate variables, which can be used to determine if 12 the model configuration accurately represents the underlying biogeochemical processes 13 creating that output (is the model getting the right answers for the right reasons).

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1 Contributions

Karen McLaughlin was the co-lead on this project with Meredith Howard. McLaughlin
contributed to study design, led the nutrient and stable isotope analysis and nitrification rates
study, conducted the data analysis and drafted the final report.

5 Meredith D.A. Howard was the co-lead on the project with Karen McLaughlin. She

6 contributed to study design, managed project budget, coordinated the field work and led the

7 primary production and respiration portions of the study. She contributed to primary

8 production and respiration sections of the report and reviewed the final draft.

George Robertson coordinated field sampling with the Sanitation Districts, contributed
to study design, reviewed data analysis plan and preliminary data, and reviewed the
manuscript on behalf of management agencies.

12 Carly D.A. Beck was the lead research technician on the project. She contributed to 13 study design, managed field staff in the field and coordinated data collection and quality 14 assurance and reviewed the final report.

15 Minna Ho is a partner on the modeling effort. She provided the particle tracking model 16 output and interpretation of the observational data.

17 Fayçal Kessouri is the lead modeler working on nutrient impacts on coastal

18 environments. Kessouri engineered the ROMS model used for particle tracking model output

19 and interpretation of the observational data.

Nikolay P. Nezlin contributed data analysis for the nitrification study and analysis of
water column profiles.

Martha Sutula is the Principal Investigator of the Biogeochemistry Department at
 SCCWRP, she contributed project management, study design, and drafted portions of the final
 report.

Stephen B. Weisberg is the Executive Director at SCCWRP, he contributed project
 management, coordination with stakeholders, and reviewed the final draft.

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- 28

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14 Dr. Martha Sutula is an editor in the Ecology and Earth Systems Domain and the lead author's husband, Dr. Steven Allison, is Editor-in chief for that same Domain. 15

16 **Data Accessibility**

17 This study was done in collaboration with the Southern California Bight Regional 18 Marine Monitoring Program, a collaborative monitoring program whose goal is to collect 19 regional-scale data on the health of coastal habitats that can be used to make better 20 management decisions. Pursuant to this goal, all data will be made publicly available on 21 SCCWRP's website: www.sccwrp.org

1 Table 1. Site information.

Station ID	Site Description	Region Category	Periods Sampled	Latitude/ Longitude	Depth (m)
2903	LACSD Ocean Outfall	Nearfield	All	33.698/ -118.336	60
3053	LACSD Off-Outfall (northern current)	Nearfield	Summer 2014	33.730/ -118.402	60
3003	LACSD Off-Outfall (northern current)	Nearfield	Spring 2015	33.757/ -118.441	60
2803	LACSD Off-Outfall (southern current)	Nearfield	Summer 2015 Spring 2016	33.668/ -118.297	60
2602	Long Beach Harbor Shelf (LA County)	Nearfield	All	33.694/ -118.191	23
2205	OCSD Ocean Outfall	Nearfield	All	33.576/ -118.005	57
2306	OCSD Off-Outfall (northern current)	Nearfield	Summer 2014	33.581/ -118.052	114
2103	OCSD Off-Outfall (southern current)	Nearfield	Spring 2015 Summer 2015 Spring 2016	33.585/ -117.945	110
1903	Orange County Southern Transect Line	Nearfield	All	33.546/ -117.836	100
CP1	Camp Pendleton- on shelf	Farfield	All	33.215/ -117.481	65
CP2	Camp Pendleton- continental slope	Farfield	All	33.184/ -117.523	430
SPOTS	San Pedro Ocean Time Series (LA County Offshore)	Farfield	All	33.607/ -118.409	730
90.30	CALCOFI station 90 30 (Orange County Offshore)	Farfield	All	33.419/ -117.912	580

2 Table 2. Nutrient properties of OCSD and LACSD wastewater effluent. Isotope samples for

3 Aug 2015 and Mar 2016 were not analyze	3	Aug 2015 and Mar 2016 were not analyzed
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Agency	LACSD			OCSD				
Season	Aug-14	Mar-15	Aug-15	Mar-16	Aug-14	Mar-15	Aug-15	Mar-16
Discharge								
Rate	268	263	257	256	125	119	92	93
(MGD)								
PO₄ (μM)	5.5	5.3	3.8	3.5	23.4	34.6	26.5	24.2
NO₂ (μM)	3.9	6.1	15.1	6.2	196	238	152	170
NO₃ (μM)	5.1	4.2	10.5	6.5	883	852	784	850
NH₄ (μM)	2690	3460	2670	1260	1890	1460	1800	2990
DIN	2699	3470	2695	1273	2969	2550	2736	4010
DIN	2 8	1 8	27	17	1 0	16	1 2	2 0
Loading	v10 ⁷	×10 ⁷	1.5 v10 ⁷	×10 ⁷				
(g/day)	×10	×10	×10	×10		×10	×10	×10
N:P	490	652	1540	852	127	74	103	94
DOC (μM)	1107	1174	1379	1918	1186	1129	1497	2241
DIC (μM)	4900		4361	2750	5076	6398	4404	2909
δ ¹⁵ N _{NO2+NO3} (‰)	-9.6	-2.3			9.1	7.5		
δ ¹⁸ Ο _{NO2+NO3} (‰)	-15.2	-15.6			-5.8	-3.5		
δ ¹⁵ Ν _{NH4} (‰)	7.5	5.1	9.6	12.4	9.1	8.9	8.6	9.1
enit	-17.1	-7.4			0	-1.9		

1 Table 3. Spearman rank correlations between dissolved and particulate variables.

- 2 Color scale indicates the correlation coefficient, where red indicates positive correlation, blue
- 3 negative, white is 0 (no correlation). Stars indicate significance where * is p < 0.05, ** where p
- 4 < 0.01, and *** where p < 0.001. Parameters were ordered using hierarchical clustering.





- 2 Figure 1. Station Locations. The nearfield sub-regions are shown in red and include LACSD
- and OCSD grid stations. The farfield (reference) sub-regions spatially distant from the
- 4 outfalls are shown in blue and include 2 offshore stations (SPOTS and CalCOFI 90.30) and 2
- 5 nearshore stations located by Camp Pendleton, in northern San Diego County (CP1 and CP2).
- 6 CTD casts were collected at additional stations within the regulatory monitoring grids
- 7 around both LACSD's and OCSD's ocean outfalls to determine the location of the plume at
- 8 the time of sampling to select the second plume station.
- 9







3 Figure 2. Spatial and temporal patterns in temperature, salinity, dissolved oxygen, and

4 chlorophyll during each sampling event. Figure is faceted by nearfield stations and farfield

5 stations for each parameter and by season. Each subregion was sampled in a different week

- 6 in a single month, introducing some temporal variability into the dataset within a season.
- 7 Plume is generally located between 20 and 60 m depth in the nearfield sites (Figure SI.1).
- 8
- 9
- 10



2 Figure 3. Depth profiles of dissolved inorganic phosphate, ammonia, nitrite, and nitrate.

3 Plume is generally located between 20 and 60 m depth in the nearfield sites (Figure SI.1).



Figure 4. Property-property plots for dissolved and particulate nutrients. Particulate carbon

- as a function of particulate nitrogen and particulate phosphorus, particulate nitrogen as a
- function of particulate phosphorus, and dissolved inorganic nitrogen (DIN) as a function of
- dissolved phosphate.





- Figure 5. Property-property plots of $\delta^{15}N_{NO3}$ and nitrate concentration (A), $\delta^{15}N_{NO2+NO3}$ and 2 N^{*} (B), and $\delta^{15}N_{NO2+NO3}$ and $\delta^{18}O_{NO2+NO3}$, where the black line represents 1:1 (C), $\delta^{13}C_{PM}$ and 3 particulate carbon concentration (D), $\delta^{13}C_{PM}$ and $\delta^{15}N_{PM}$ (E), $\delta^{13}C_{PM}$ and chlorophyll *a* (F), 4 $\delta^{15}N_{PM}$ and particulate nitrogen concentration (G), $\delta^{15}N_{PM}$ and $\delta^{15}N_{NO2+NO3}$ (H), $\delta^{15}N_{PM}$ and 5 chlorophyll a (I). Colors represent the seasons; shapes represent the station types. The size 6 7 of the points only applies to the nitrite fraction in panel (C) where increased fraction nitrite in the nitrite+nitrate may underestimate $\delta^{18}O_{NO2+NO3}$. Supplemental Figure SI.12 shows how 8 9 this underestimation may effect results.
- 10
- 11



2 Figure 6. Depth profiles of dissolved $\delta^{15}N_{NO3}$ and $\delta^{18}O_{NO3}$, and particulate $\delta^{13}C_{PM}$ and $\delta^{15}N_{PM}$.

- 3 Plume is generally located between 20 and 60 m depth in the nearfield sites (Figure SI.1).
- 4



2 Figure 7. Depth integrated primary production rates (A), average subsurface respiration rates

- 3 (B), and nitrification rates faceted by depth layer (C) by station type. Black points represent
- 4 individual data points. There was no data collected of primary production in Summer 2014.
- 5 Error bars for indicate the standard deviation.
- 6
- 7



- Figure 8. Downscaled Regional Ocean Model (350 m scale) run in particle tracking mode demonstrating the fate of wastewater effluent released from ocean outfalls. Color bar represents the Coriolis-normalized vorticity (a measure of ocean stirring). Black circles represent particles released from outfalls (green ovals). Red boxes are nearfield areas and blue boxes represent farfield, offshore areas. Panel A shows particle distribution 9 hours from the start of the simulation on January 1 at 01:00). Two weeks into the simulation, particles are nearly completely mixed in the nearshore areas (Panel B). Panel C shows the
- 8 end of the simulation, two months, particles released from the outfalls have been mixed to
- 9 the offshore regions (Dauhajre et al. 2019).

1		
2	SUPPLEMENTAL MATERIALS	
3		
4		
5		



- 2 Figure SI.1. Spatial extent of the effluent plumes during each of the sampling events as
- 3 measured by CDOM (mg m⁻³). Note colored scale on each graphic is different, scaling was
- 4 free on each figure to demonstrate the extent of the plume. Scales are plotted from the
- 5 minimum reported value to the maximum for each site to identify the plume locations over
- 6 each outfall. Each CTD cast location is represented by a black vertical line, multiple casts
- 7 were taken to determine the direction of the plume during sampling events and not all
- 8 stations correspond to sampling locations on Figure 1. The station sampled over each outfall
- 9 is highlighted in red.
- 10
- 11



2 Figure SI.2. Timing of sampling and ranges of data for upper 60 m at each site. Black dot

- 3 represents the mean and bar the standard deviation of all values in upper 60 m.
- 4



- 2 Figure SI.3. Depth profiles of particulate carbon, nitrogen and phosphorus. Plume is
- 3 generally located between 20 and 60 m depth in the nearfield sites.



- 2 Figure SI.4. Depth profiles of particulate C:N, C:P, and N:P. Plume is generally located
- 3 between 20 and 60 m depth in the nearfield sites.



Figure SI.5. The $\delta^{15}N_{NO2+NO3}$ as a function of nitrate concentration, N*, nitrification rate, and

- $\delta^{18}O_{NO2+NO3}$. The black line in the final column represents a 1:1 relationship between the
- relative enrichment of the two isotopes. The size of the points represents the fraction of
- nitrite in the nitrite+nitrate measured in each sample; larger points have higher fraction of nitrite.



- 2 Figure SI.6. Depth profiles of the stable isotopic composition of particulate carbon ($\delta^{13}C_{PM}$)
- and nitrogen ($\delta^{15}N_{PM}$). Plume is generally located between 20 and 60 m depth in the
- 4 nearfield sites.
- 5



 $2 \qquad \mbox{Figure SI.7. The $\delta^{15}N_{PM}$ as a function of particulate nitrogen concentration, dissolved}$

3 inorganic nitrogen (DIN), $\delta^{15}N_{NO2+NO3}$, and chlorophyll *a* concentration.

4





4 Figure SI.8. δ^{13} C_{PM} as a function of particulate carbon, DIN, δ^{15} N_{PM}, and chlorophyll *α*

- 5 concentration.
- 6



- Figure SI.9. Depth integrated primary production rates (A), average subsurface respiration
- 3 rates (B), and nitrification rates faceted by depth layer (C) by station. There was no data
- 4 collected of primary production in Summer 2014. Error bars for indicate the standard
- 5 deviation.
- 6



2 Figure SI.10 Relationship between depth integrated primary production and average

3 subsurface respiration for the three-time periods when both parameters were measured.

4



- 3 Figure SI.11. Variable importance from random forest regression models predicting primary
- 4 production (A), respiration (B), and Nitrification (C). Predictor variables were plotted with
- 5 their increase in percent mean squared error (%IncMSE) when the variable is randomly
- 6 permuted in the model, higher values indicate variables were more important to the
- 7 regression. The overall percent variance explained using the random forest models was 30%
- 8 for primary production, 18% for respiration, and 20% for nitrification.
- 9

10

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Figure SI.12. Estimation of the impact of high nitrite on the $\delta^{18}O_{NO2+NO3}$. For samples with high 2 3 nitrite concentrations, the denitrifier method employed in this study for oxygen isotope 4 analysis of dissolved nitrate + nitrite is expected to underestimate the oxygen isotopic 5 composition of nitrite by 25-30‰. We calculated the fraction of nitrite in each sample and estimated the impact of a 25‰ underestimation of the $\delta^{18}O_{NO2}$ on the final $\delta^{18}O_{NO2+NO3}$ (Adj 6 7 $\delta^{18}O_{NO2+NO3}$ right panel). Values were increased an average of 2.7 ‰ with a maximum increase of 13.8 ‰. Samples with the highest fraction of nitrite (indicated by the size of the point), had 8 9 the greatest increase.

11