

appendix A

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## **DETAILED METHODS**

### INTRODUCTION

This appendix contains a summary of the methods used for field sampling, laboratory testing, and data analysis used in the District's Ocean Monitoring Program (OMP). The methods include those for calculations of water quality compliance with California Ocean Plan (COP) criteria, coastal oceanography, water quality monitoring, bacteriology/nutrients, sediment geochemistry, invertebrate and fish community analysis, fish health, and fish tissue contaminants. More detailed methods can be found in the program Quality Assurance Project Plan (QAPP) (OCSD 2008c), Environmental Assessment Division Standard Operating Procedures (OCSD 2008b), and the Environmental Sciences Laboratory Operating Procedures Manual (LOPM) (OCSD 2008a).

For 2010-11, the OMP was conducted under conditions stipulated in the District's 2004 NPDES discharge permit (Order No. R8-2004-0062, NPDES No. CA0110604).

### COASTAL OCEANOGRAPHY

Mooring locations and equipment used for measurement of ocean currents are summarized below.

#### ***Teledyne RD Instruments Workhorse Acoustic Doppler Current Profiler (ADCP).***

##### Field Methods

Three Teledyne/RDI Workhorse Sentinel Acoustic Doppler Current Profilers (ADCP) were deployed across the shelf from the 78-inch outfall (~20 m) to the 120-inch outfall (~60 m) and a fourth ADCP was located downcoast of the 120-inch outfall near Newport Canyon (Table A-1, Figure 3-1). The ADCPs were deployed in trawl resistant bottom mounts (TRBM). Two of the ADCPs had a frequency of 300 kHz and two had a frequency of 600 kHz.

All four ADCPs were equipped to measure water pressure and bottom temperature. Currents were sampled in 1 m depth bins. Data were collected every 6 minutes using a burst sampling interval of 1 minute duration. Compass calibrations and deployment parameters were done using WinSC (2003). A typical deployment lasted about three months. Raw data files were copied from the instruments' compact flash memory cards using WinADCP (2003).

##### Data Analyses

Current meter data were converted into scientific units using WinADCP (2003). Pitch and roll values were checked to see if there was a serious tilt on the instrument during the deployment procedure. If the tilt exceeded 10°, current data was excluded from further analysis, and only temperature data was used. The time base for each instrument was checked for agreement with the projected number of records and for agreement with the logged deployment and recovery times for these sampling events. Pre- and post-deployment periods were truncated from the time series to remove invalid portions of the data when the ADCP was not positioned on the bottom (Figure 3-1).

The remainder of the data processing was conducted in MATLAB (2007). All non-usable data points and outliers were replaced with a "NaN" data flag. These data points were then replaced with linearly interpolated data before applying the low-pass filter. A Finite Impulse Response (FIR) filter was used to perform a zero-phase distortion, forward and reverse digital filter that minimizes start-up and ending transients by matching the initial conditions. A 40-hour low pass cutoff frequency was chosen to remove the tidal and higher frequency signals from the data.

Graphical data were shown as vector plots ("stick plots") with the lines pointing in the direction of the current and the length

**Table A-1. Deployment summary for District's acoustic Doppler current profilers for 2010-11.**

Orange County Sanitation District, California.

Quarter	Station	Deployment Date	Recovery Date	Latitude	Longitude	Depth (m)	Instrument Average Depth (m)	Sampling Interval
Summer	M18	05/05/2010	08/05/2010	33.57690	-118.02470	60	63.7	6 minutes
	M19	05/27/2010	08/19/2010	33.58950	-118.00690	40	41.7	6 minutes
	M20	05/27/2010	08/05/2010	33.60980	-117.97740	20	21.5	6 minutes
	M21	05/27/2010	08/19/2010	33.57960	-117.95900	60	56.7	6 minutes
Fall	M18	08/05/2010	11/23/2010	33.57650	-118.02460	60	63.4	6 minutes
	M19	08/23/2010	11/30/2010	33.59050	-118.00690	40	39.7	6 minutes
	M20	08/05/2010	11/23/2010	33.60980	-117.97840	20	22.0	6 minutes
	M21	08/23/2010	11/30/2010	33.58020	-117.95890	60	55.6	6 minutes
Winter	M18	11/30/2010	03/08/2011	33.57680	-118.02470	60	63.4	6 minutes
	M19	12/01/2010	03/10/2011	33.59030	-118.00620	40	40.0	6 minutes
	M20	11/23/2010	03/08/2011	33.60830	-117.97900	20	23.0	6 minutes
	M21	11/30/2010	03/10/2011	33.57960	-117.95810	60	56.3	6 minutes
Spring	M18	05/25/2011	10/24/2011	33.57330	-117.99540	60	60.5	6 minutes
	M19	05/25/2011	10/24/2011	33.58950	-117.98400	40	42.4	6 minutes
	M20	05/25/2011	10/26/2011	33.60670	-117.97240	20	22.0	6 minutes
	M21	05/25/2011	10/26/2011	33.57920	-117.95900	60	57.3	6 minutes

proportional to the current magnitude. Because current vectors generally tend to follow the local bathymetry, the frame-of-reference for the figures is rotated so that upward-directed “sticks” correspond to upcoast flows and sticks directed to the right correspond to onshore flows. For this presentation, the bathymetric orientation for the primary mooring line is 302°, and for Mooring M21 near Newport Canyon the orientation is 287°. These are consistent with bathymetry orientation values used for prior studies in the District’s study region (OCSD 2004).

## **WATER QUALITY MONITORING**

### ***Field Methods***

#### Offshore Water Quality Monitoring

Permit-specified water quality studies were conducted three days per quarter at 29 stations comprising a fixed-grid pattern (Tables A-2 and A-3, Figure 3-1). Each survey included measurements of pressure (from which depth is calculated), water temperature, conductivity (from which salinity is calculated), dissolved oxygen (DO), pH, water clarity (light transmissivity and Secchi-disc depth), chlorophyll-a, colored dissolved organic matter (CDOM), and photosynthetically active radiation (PAR). Profiling was conducted from the surface (1 m below) to 2 m from the bottom or to a maximum depth of 75 m, when station bottom depths are greater than 75 m. Measurements were conducted using a Sea-Bird Electronics SBE9-03/SBE 11 Deck Unit (SBE9/11) CTD (conductivity-temperature-depth) profiling system. SEASOFT (2010a) software was used for data acquisition, data display, and sensor calibration. A summary of the sampling methods are presented in Table A-4. Light transmittance and PAR were measured for water clarity determinations. PAR is measured in conjunction with chlorophyll-a because of the positive linkage between light intensity and photosynthesis per unit chlorophyll (Hardy 1993).

Visual observations of water clarity (measured as Secchi-disc depth), water color, and floatable materials or grease that might be of sewage origin, were also conducted. Water

color was determined using a Forel/Ule scale at one-half of the Secchi-disc depth. Daily rainfall, sea state, and wind condition data were summarized from Newport Beach Fire and Marine Department and District’s Treatment Plant No. 2 records.

Discrete sampling for ammonium, total coliforms, fecal coliforms, *Escherichia coli* (*E. coli*), and enterococci was conducted using a Sea-Bird Electronics Carousel Water Sampler (SBE32/SBE33) equipped with Niskin bottles. Sample depths are provided in Table A-2. Bacteriology samples were kept on wet ice in coolers and transported to the District’s laboratory within 6-hours of collection for analysis.

#### Central Bight Water Quality Strategic Process Study (SPS)

An expanded grid of water quality stations was sampled quarterly as part of the District’s Central Bight Water Quality Strategic Process Study (SPS). These additional stations were sampled concurrently with the permit specified stations and in conjunction with the Los Angeles County Sanitation District (LACSD), the City of Los Angeles, the City of Oxnard, and the City of San Diego. The northern sampling area extends from the Ventura River in the north to Crystal Cove State Beach in the south. The southern sampling area extends from Mission Bay in the north to south of the United States/Mexico border. Samples were collected using CTDs during a targeted 3- or 4-day period, over which sampling occurred at 300 stations comprising a fixed-grid pattern (see Figure 3-2). Parameters measured included pressure, water temperature, conductivity, dissolved oxygen, pH, chlorophyll-a, CDOM, and water clarity. Profiling was conducted from the surface depth to 2 m from the bottom or to a maximum depth of 75–100 m. Sampling and analytical methods were the same as those presented in Tables A-2 and A-4.

#### ***Data Analyses***

Raw CTD data were processed using both SEASOFT, (2010b) and third party (IGODS, 2010) software. The steps include retaining downcast data and identifying outliers by flagging the data if it exceeded specific criteria

**Table A-2. OCSD ocean monitoring program station positions\* and nominal depths.**

Orange County Sanitation District, California.

Station	Latitude	Longitude	Depth (m)	Sample Depths
<b>Offshore Water Quality</b>				
1901 *	33° 33.682' N	117° 49.654' W	10	None
1902 *	33° 33.165' N	117° 49.944' W	60	None
1903 *	33° 32.762' N	117° 50.182' W	100	None
1904 *	33° 31.787' N	117° 50.734' W	405	None
1905 *	33° 30.810' N	117° 51.285' W	510	None
1906 *	33° 29.829' N	117° 51.842' W	550	None
2001 *	33° 35.335' N	117° 51.564' W	10	None
2002 *	33° 34.755' N	117° 51.844' W	60	None
2003 *	33° 34.565' N	117° 52.123' W	100	None
2004 *	33° 33.589' N	117° 52.657' W	345	None
2005 *	33° 32.613' N	117° 53.225' W	410	None
2006 *	33° 31.647' N	117° 53.793' W	470	None
2021 *	33° 35.771' N	117° 52.099' W	10	None
2022 *	33° 35.283' N	117° 52.379' W	53	None
2023 *	33° 34.796' N	117° 52.658' W	165	None
2024 *	33° 33.811' N	117° 53.179' W	300	None
2025 *	33° 32.851' N	117° 53.741' W	390	None
2026 *	33° 31.900' N	117° 54.301' W	432	None
2101 *	33° 36.183' N	117° 55.749' W	10	None
2102 *	33° 35.631' N	117° 56.206' W	26	None
2103 ***	33° 35.089' N	117° 56.678' W	110	1, 5, 10, 15, 30, 45, 60
2104 ***	33° 34.199' N	117° 57.414' W	143	1, 5, 10, 15, 30, 45, 60
2105 **	33° 33.309' N	117° 58.150' W	280	1, 5, 10, 15, 30, 45, 60
2106 **	33° 32.420' N	117° 58.885' W	309	1, 5, 10, 15, 30, 45, 60
2181 *	33° 36.877' N	117° 56.752' W	10	None
2182 *	33° 36.272' N	117° 57.264' W	15	None
2183 ***	33° 35.701' N	117° 57.744' W	36	1, 5, 10, 15, 30, 34
2184	33° 34.811' N	117° 58.480' W	51	None
2185	33° 33.922' N	117° 59.215' W	114	None
2186	33° 33.032' N	117° 59.951' W	247	None
2201 *	33° 37.493' N	117° 57.831' W	10	None
2202 *	33° 36.901' N	117° 58.314' W	16	None
2203 ***	33° 36.313' N	117° 58.810' W	25	1, 5, 10, 15, 23
2204 **	33° 35.423' N	117° 59.546' W	39	1, 5, 10, 15, 30, 37
2205 **	33° 34.534' N	118° 00.282' W	57	1, 5, 10, 15, 30, 45, 55
2206 **	33° 33.644' N	118° 01.018' W	185	1, 5, 10, 15, 30, 45, 60
2221 *	33° 38.099' N	117° 58.908' W	10	None
2222 *	33° 37.522' N	117° 59.374' W	15	None
2223 ***	33° 36.924' N	117° 59.871' W	22	1, 5, 10, 15, 20
2224	33° 36.035' N	118° 00.608' W	31	None

**Table A-2 Continues.**

Table A-2 Continued.

Station	Latitude	Longitude	Depth (m)	Sample Depths
<b>Offshore Water Quality</b>				
2225	33° 35.146' N	118° 01.346' W	47	None
2226	33° 34.257' N	118° 02.083' W	135	None
2301 *	33° 38.572' N	118° 00.064' W	10	None
2302 *	33° 38.053' N	118° 00.495' W	15	None
2303 ***	33° 37.537' N	118° 00.936' W	21	1, 5, 10, 15, 19
2304 **	33° 36.649' N	118° 01.674' W	29	1, 5, 10, 15, 27
2305 **	33° 35.760' N	118° 02.412' W	38	1, 5, 10, 15, 30, 36
2306 **	33° 34.871' N	118° 03.149' W	114	1, 5, 10, 15, 30, 45, 60
2349 *	33° 39.190' N	118° 01.135' W	10	None
2350 *	33° 38.667' N	118° 01.566' W	14	None
2351 ***	33° 38.151' N	118° 02.001' W	21	1, 5, 10, 15, 19
2352	33° 37.262' N	118° 02.739' W	29	None
2353	33° 36.373' N	118° 03.477' W	37	None
2354	33° 35.484' N	118° 04.214' W	123	None
2401 *	33° 39.920' N	118° 02.103' W	10	None
2402 *	33° 39.342' N	118° 02.593' W	16	None
2403 ***	33° 38.765' N	118° 03.072' W	21	1, 5, 10, 15, 19
2404 **	33° 37.875' N	118° 03.808' W	29	1, 5, 10, 15, 28
2405 **	33° 36.986' N	118° 04.544' W	37	1, 5, 10, 15, 30, 35
2406 **	33° 36.096' N	118° 05.280' W	60	1, 5, 10, 15, 30, 45, 58
2451 *	33° 41.475' N	118° 03.944' W	10	None
2452 *	33° 40.739' N	118° 04.584' W	17	None
2453 *	33° 39.987' N	118° 05.204' W	22	None
2454 *	33° 39.098' N	118° 05.946' W	30	None
2455 *	33° 38.210' N	118° 06.675' W	36	None
2456 *	33° 37.318' N	118° 07.411' W	42	None
C2 ***	33° 36.125' N	117° 56.014' W	56	1, 5, 10, 15, 30, 45, 52
<p>* = Central Bight WQ Regional Grid station  ** = Ammonia samples  *** = Ammonia samples and Bacteria (REC-1) samples</p>				

Table A-2 Continues.

Table A-2 Continued.

Station	Latitude	Longitude	Depth (m)
<b>Nearshore (Surfzone) Water Quality</b>			
39N	33° 42.114' N	118° 03.321' W	surf
33N	33° 41.281' N	118° 02.495' W	surf
27N	33° 40.587' N	118° 01.712' W	surf
21N	33° 39.843' N	118° 00.785' W	surf
15N	33° 39.114' N	117° 59.846' W	surf
9N	33° 38.565' N	117° 58.924' W	surf
6N	33° 38.331' N	117° 58.573' W	surf
3N	33° 38.018' N	117° 58.032' W	surf
0	33° 37.764' N	117° 57.598' W	surf
3S	33° 37.619' N	117° 57.264' W	surf
6S	33° 37.337' N	117° 56.704' W	surf
9S	33° 37.033' N	117° 56.283' W	surf
15S	33° 36.342' N	117° 55.459' W	surf
21S	33° 36.059' N	117° 54.213' W	surf
27S	33° 35.646' N	117° 52.910' W	surf
29S	33° 35.559' N	117° 52.508' W	surf
39S	33° 34.700' N	117° 51.946' W	surf
<b>Benthic</b>			
0 *	33° 34.573' N	118° 00.598' W	56
1 *	33° 34.657' N	118° 00.968' W	56
3	33° 34.434' N	118° 00.660' W	60
4 *	33° 34.498' N	117° 59.761' W	56
5	33° 34.749' N	118° 01.612' W	59
7	33° 35.325' N	118° 00.367' W	41
8	33° 35.164' N	117° 59.555' W	44
9 *	33° 34.363' N	117° 59.510' W	59
10	33° 34.902' N	118° 02.081' W	62
12 *	33° 34.385' N	117° 59.054' W	58
13	33° 35.307' N	118° 02.944' W	59
17	33° 33.961' N	118° 00.187' W	91
18	33° 34.064' N	118° 00.750' W	91
20	33° 34.599' N	118° 02.229' W	100
21	33° 35.313' N	118° 01.891' W	44
22	33° 35.204' N	117° 59.028' W	45
23	33° 33.968' N	117° 59.147' W	100
24	33° 33.563' N	118° 01.140' W	200
25	33° 33.924' N	118° 02.176' W	200
27	33° 33.326' N	117° 59.708' W	200
29	33° 35.033' N	118° 03.113' W	100
30	33° 35.493' N	118° 02.899' W	46
33	33° 34.349' N	117° 57.866' W	100

Table A-2 Continues.

Table A-2 Continued.

Station	Latitude	Longitude	Depth (m)
<b>Benthic</b>			
36	33° 35.308' N	117° 57.495' W	45
37	33° 34.832' N	117° 57.369' W	56
38	33° 34.634' N	117° 57.317' W	100
39	33° 33.283' N	117° 58.531' W	200
40	33° 32.496' N	117° 59.775' W	303
41	33° 32.690' N	118° 01.149' W	303
42	33° 33.098' N	118° 02.598' W	303
44	33° 34.586' N	118° 05.422' W	241
55	33° 36.739' N	118° 05.413' W	40
56	33° 35.665' N	118° 05.417' W	100
57	33° 34.970' N	118° 05.418' W	200
58	33° 33.365' N	118° 05.347' W	300
59	33° 36.070' N	118° 03.701' W	40
60	33° 35.532' N	118° 04.017' W	100
61	33° 35.011' N	118° 04.326' W	200
62	33° 34.069' N	118° 04.568' W	300
63	33° 34.173' N	118° 03.407' W	200
64	33° 33.484' N	118° 03.663' W	300
65	33° 33.859' N	117° 57.230' W	200
C *	33° 35.799' N	118° 03.855' W	56
C2	33° 36.125' N	117° 56.014' W	56
C4	33° 35.056' N	117° 55.833' W	187
C5	33° 33.920' N	117° 55.620' W	296
Control 1 *	33° 36.037' N	118° 05.387' W	59
ZB *	33° 34.545' N	118° 00.274' W	56
ZB2 *	33° 34.590' N	118° 00.611' W	56
* = Quarterly Stations			
<b>Trawl</b>			
T0*	33° 37.117' N	117° 59.283' W	18
T1	33° 34.641' N	118° 00.567' W	55
T2	33° 35.688' N	117° 59.561' W	35
T3	33° 34.856' N	117° 57.345' W	55
T6	33° 35.946' N	118° 02.785' W	36
T10	33° 33.771' N	118° 00.250' W	137
T11	33° 36.055' N	118° 05.199' W	60
T12	33° 34.868' N	118° 01.670' W	57
T13	33° 35.535' N	118° 03.637' W	60
T14	33° 34.672' N	118° 03.200' W	137
* = T0 sampled for historical purposes.			



**Table A-3. Sampling dates during 2010-11.**

Orange County Sanitation District, California.

Quarter	Date	Purpose
<b>Water Quality</b>		
Summer	07/15/2010	Water Quality
	08/03/2010	Water Quality
	08/04/2010	Central Bight
	08/10/2010	REC-1 Water Quality
	08/11/2010	REC-1 Water Quality
	08/12/2010	REC-1 Water Quality
	09/16/2010	Water Quality
Fall	10/21/2010	REC-1 Water Quality
	10/26/2010	REC-1 Water Quality
	10/27/2010	Water Quality
	11/02/2010	Water Quality
	11/03/2010	Central Bight
	11/04/2010	REC-1 Water Quality
	12/08/2010	Water Quality
Winter	01/26/2011	Water Quality
	01/27/2011	REC-1 Water Quality
	02/01/2011	REC-1 Water Quality
	02/02/2011	REC-1 Water Quality
	02/08/2011	Water Quality
	02/09/2011	REC-1 Water Quality/Central Bight
	02/10/2011	REC-1 Water Quality/Central Bight
	03/22/2011	Water Quality
Spring	04/26/2011	REC-1 Water Quality Compliance Data
	04/27/2011	Water Quality Compliance Data
	05/03/2011	REC-1 Water Quality Compliance Data
	05/04/2011	REC-1 Water Quality Compliance Data
	05/10/2011	Water Quality Compliance Data
	05/11/2011	Central Bight Water Quality Data Only
	06/09/2011	Water Quality Compliance Data
<b>Sediment and Infauna</b>		
Summer	07/07-14/2010	Summer Quarterly and Annual Stations
Fall	10/05-07/2010	Fall Quarterly Stations
Winter	01/04-05/2011	Winter Quarterly Stations
Spring	04/05-07/2011	Spring Quarterly Stations
<b>Trawls</b>		
Summer	07/17-18,23-25/2010	Trawl Fish Community Analysis
Winter	01/11-12,18-19/2011	Trawl Fish Community Analysis & Fish Tissue Analysis

**Table A-4. Water quality sample collection and analysis methods by parameter.**

Orange County Sanitation District, California.

Parameter	Sampling Method	Method Reference	Preservation	Container	Holding Time	Sampling Interval	Field Replicates
Temperature	<i>in-situ</i> probe	(1)	none	none	not applicable	every 1 m <sup>a</sup>	at least 10% of stations
Salinity (conductivity)	<i>in-situ</i> probe	(2)	none	none	not applicable	every 1 m <sup>a</sup>	at least 10% of stations
pH	<i>in-situ</i> probe	(3)	none	none	not applicable	every 1 m <sup>a</sup>	at least 10% of stations
Dissolved Oxygen	<i>in-situ</i> probe	(4)	none	none	not applicable	every 1 m <sup>a</sup>	at least 10% of stations
Transmissivity	<i>in-situ</i> probe	(5)	none	none	not applicable	every 1 m <sup>a</sup>	at least 10% of stations
Photosynthetically Active Radiation (PAR)	<i>in-situ</i> probe	(6)	none	none	not applicable	every 1 m <sup>a</sup>	at least 10% of stations
Ammonium (NH <sub>4</sub> <sup>+</sup> )	Niskin	EPA Method 350.1B Rev. A	Ice	125 mL HDPE	28 days	Surface, 5m, 10m, 15m, 30m, 45m, 60m, Bottom	at least 10% of samples
Offshore Total Coliforms, Fecal Coliforms, <i>Escherichia coli</i> , and <i>Enterococci</i>	Niskin	Standard Methods 9223 B.	Ice (<6°)	125 mL HDPE (Sterile container)	6 hrs (field) 2 hrs (lab)	Surface, 5m, 10m, 15m, 30m, 45m, 60m, Bottom	at least 10% of samples
Shoreline Total Coliforms, Fecal Coliforms, and <i>Enterococci</i>	grab	Standard Methods 9222 B. Standard Methods 9222 D. EPA Method 1600	Ice (<6°)	125 mL HDPE (Sterile container)	6 hrs (field) 2 hrs (lab)	Ankle deep water	at least 10% of samples
Secchi Depth	visual observations	permit specs.	none	none	not applicable	surface	none
Forel/Ule Color Values	visual observations	permit specs.	none	none	not applicable	½ Secchi depth below surface	none
Surface Observations	visual observations	permit specs.	none	none	not applicable	surface	none

- <sup>a</sup> Sampled continuously but data processed to 1 m intervals.
- (1) Calibrated to reference cells (0.0005°C accuracy) every year.
  - (2) Calibrated to IAPSO Standard and Guildline 8400B Autosol every six months.
  - (3) Referenced and calibrated to NIST buffers of pH 7, 8, and 9 every survey.
  - (4) Referenced and calibrated to saturation table values.
  - (5) Referenced and calibrated to known transmittance in air.
  - (6) Factory calibrated (Biospherical Instruments) once per year.

limits. Flagged data were removed if it was considered to be due to instrument failures, electrical noise (e.g., large data spikes), or physical interruptions of sensors (e.g., by bubbles) rather than by actual oceanographic events. After outlier removal, averaged 1 m depth values were prepared from the downcast scan data; if there were any missing 1 m depths, then the upcast data was used as a replacement. CTD and discrete data were then combined to create a single data file that contained all sampled stations for each survey day.

PAR data for each station were normalized to represent the percent of the respective surface PAR values at that station. The percent of light available to phytoplankton for photosynthesis is derived from the normalized PAR data (Hardy 1993). Spatial and seasonal patterns in water quality data are summarized in 2- and 3-dimensional color plots of temperature, salinity, DO, pH, transmissivity, and two-dimensional displays of PAR and chlorophyll-a. The 3-dimensional plots were produced using IGODS (2010) software.

### Compliance Calculations

Water quality compliance is assessed based on (1) specific numeric criteria for dissolved oxygen and pH; and (2) narrative (non-numeric) criteria for transmissivity, floating particulates, oil and grease, water discoloration, beach grease, and excess nutrients. The sampling approach comprises a 2 x 2 km fixed-grid pattern with 7 nearshore-offshore transects of 4 stations each (Table A-2). Transect 7 also includes the historically relevant Newport Canyon Station C2. Station locations were defined as either Zone A (stations are along and inshore of the 3-mile limit) or Zone B (beyond the 3-mile limit) as shown in Figure A-1. Stations numbers ending in 03 and 04 comprise the Zone A stations; those ending in 05 and 06 are the Zone B stations. Compliance evaluations are based on statistical comparisons to the corresponding upcurrent Zone A or Zone B reference station (OCSD 1999). This matching of Zone A or Zone B stations allows comparisons of data from similar water depths. Water quality sampling dates for 2010-11 are shown in Table A-3.

The depth of the pycnocline layer is then calculated for each station using temperature and salinity data. The pycnocline is defined as the depth layer where stability is greater than  $0.05 \text{ kg/m}^3$  (Officer 1976). Data for each station and numeric compliance parameter (transmissivity, dissolved oxygen, and pH) are then binned by water column stratum: above, within, or below the pycnocline; or top, middle, or bottom third when a pycnocline is absent. Mean values for each parameter are calculated by stratum and station. All binned observations (1 m intervals) within each stratum are used in the calculations. The number of observations usually differs from station to station due to different water and pycnocline depths.

The selections of the appropriate reference stations (i.e., upcoast or downcoast) for each survey day are determined based on current measurements (if available) and/or on the presence or absence of ammonium at stations upcoast or downcoast from the outfall diffuser. Once reference stations are determined, the data are analyzed using in-house MATLAB (2007) routines to calculate out-of-range occurrences (OROs) for each sampling date and parameter. These OROs are based on comparing the mean data by stratum and test station with the corresponding reference station data to determine whether the following COP criteria were exceeded:

- Dissolved oxygen: cannot be depressed >10% below the mean;
- pH: cannot be greater than +/- 0.2 pH units different than the mean; and
- Natural light, defined as transmissivity: shall not be significantly reduced, where statistically different from the mean is defined as the 95% confidence limit.

In accordance with permit specifications, the outfall station (2205) is not included in the comparisons because it is within the zone of initial dilution (ZID).

To determine whether an ORO is out-of-compliance (OOC), distributional maps are created that identify the reference stations for each sampling date and location of each ORO,

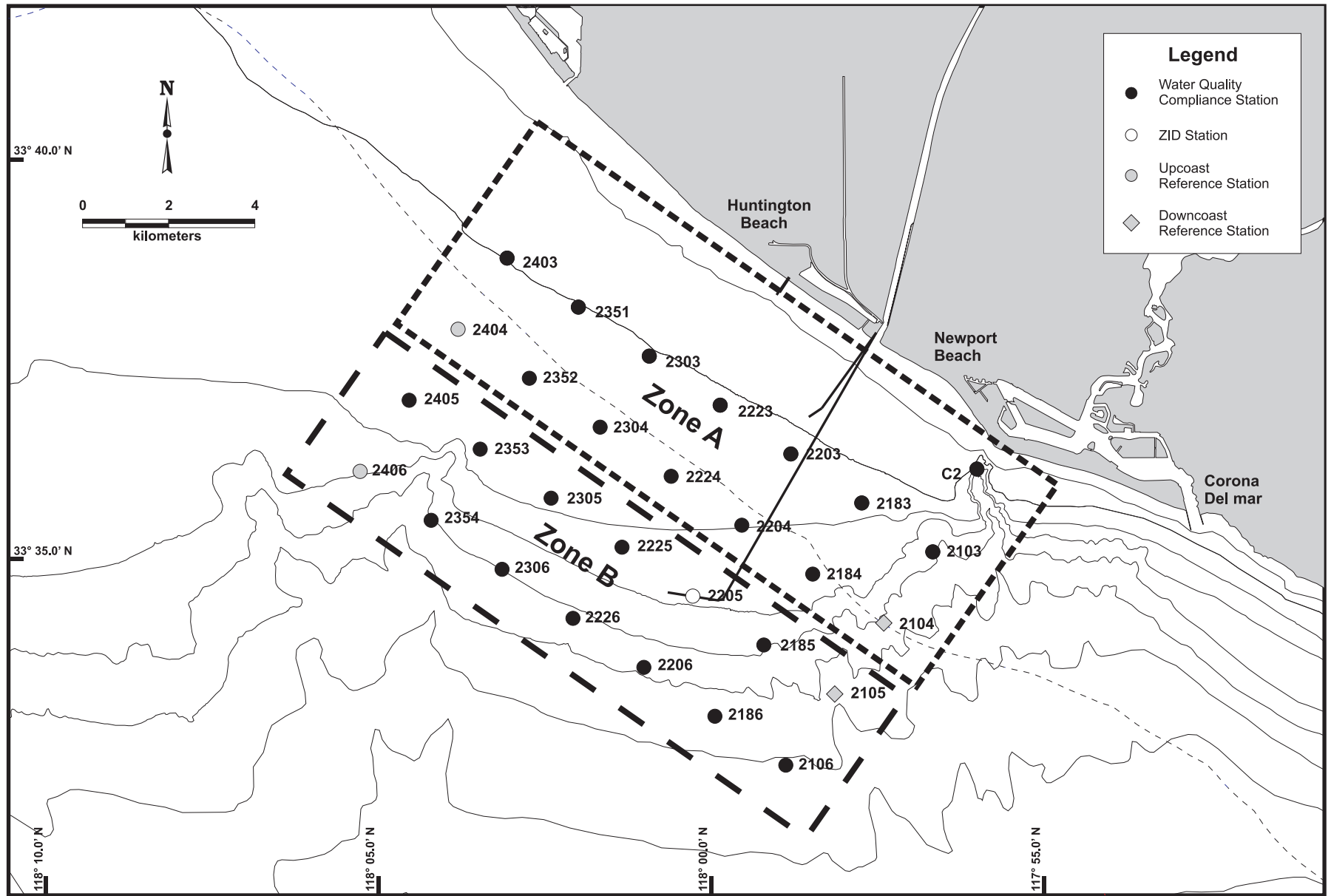


Figure A-1. Water quality monitoring stations and zones used for compliance determinations.

including which stratum is out of range. Each ORO is then evaluated to determine if it represents a logical OOC event. These evaluations are based on (A) evaluation of the wastewater plume location relative to depth using a combination of temperature, salinity, and CDOM (colored dissolved organic matter); (B) evaluation of features in the water column relative to naturally occurring events (i.e., high chlorophyll-a due to red tide); and (C) unique characteristics of some stations that may not be comparable with permit-specified reference stations (2104/2105 or 2404/2406) due to differences in water depth and/or variable oceanographic conditions. For example, Zone A stations (2103, 2203, 2303, and 2403) are located at shallower depths than reference station 2104. Waves and currents can cause greater mixing and resuspension of bottom sediments at shallower stations under certain conditions (e.g., winter storm surges). This can result in naturally decreased water clarity (transmissivity) that is unrelated to the wastewater discharge. An ORO can be in-compliance if, for example, a downcurrent station is different from the reference, but no intermediate (e.g., nearfield) stations exhibit OROs.

Once the total number of OOC events is summed by parameter, the percentage of OROs and OOCs is calculated according to the total number of observations. In a typical year, Zone A, has a total of 504 possible comparisons if 14 stations (not including the reference station) and three strata over 12 survey dates per year are used. For Zone B, 432 comparisons are possible from 12 stations (not including the reference station), three strata, and 12 sampling dates. The total combined number of ORO and OOC events is then determined by summing the Zone A and Zone B results. If not all of the strata are present or additional surveys are conducted, the total number of comparisons in the analysis may be more or less than the target number of comparisons possible (936).

## BACTERIOLOGY/NUTRIENTS

### Field Methods

#### Offshore Monitoring

Quarterly surveys were conducted in August and November 2010, and February and May 2011. Primary water quality sampling occurred three times per quarter at 29 sites that form a grid around the OCSD diffuser (see Figure 3-1). Samples were collected for ammonium at a subset of 20 stations and for bacteriology at 9 REC-1 stations (Tables A-2, Figure 3-1) located within state waters (within 3 miles of the shoreline) for the purposes of determining compliance with Receiving Water Limitation C.2.a.1. Discrete samples for ammonium and bacteria (total coliform, *E. coli*, and enterococci) were collected at 5 and 15 m depth intervals from the surface (1 m below) to 2 m above the ocean floor or a maximum depth of 60 m (Table A-2). Two additional surveys are conducted to collect bacteriology and ammonium samples at the nine REC-1 water quality stations within 30 days of the three primary water quality sampling days in order to calculate a 30-day geometric mean. During most surveys, additional bacteriological samples were collected at the outfall (Station 2205) and two nearfield stations (1 and 9).

#### Nearshore Monitoring

Nearshore (surfzone) samples for analysis of total and fecal coliform and enterococci bacteria were collected at 17 required stations, plus an additional 3 stations at a frequency ranging from 3-5 days per week (Table A-2, Figure 3-1). The permit requires that beginning Memorial Day to Labor Day, samples are collected five times per week. From Labor Day through October 31, and April 1 to Memorial Day, samples are collected three times per week and from November 1 through March 31; samples are collected twice per week”

Samples were collected in ankle-deep waters, on an incoming wave, with the sampler downstream and away from the bottle, and the mouth of the bottle facing into the current. Sterile sample bottles were used and the sampler used aseptic techniques, making certain that the bottle does not touch the ocean bottom. After the sample was taken, the bottle

was tightly capped and promptly stored on ice in the dark. Laboratory analysis began within 6 hours of sample collection. The occurrence and size of any grease particles at the high tide line was recorded 2 days per week at the same sampling locations.

## Laboratory Methods

Laboratory analyses of ammonium and bacteriology samples followed standard EPA guidelines, as listed in Table A-4. Quality assurance/quality control (QA/QC) procedures included analysis of laboratory blanks and duplicates. All data underwent at least three separate reviews prior to being included in the final database used for statistical analysis, comparison to standards, and data summaries.

## Data Analyses

### Offshore Bacteriology Monitoring

Spatial and seasonal patterns of several water quality parameters are summarized graphically in 2- and 3-dimensional color figures. The 3-dimensional plots were produced using IGODS (2010) software. Prior to disinfection (see Chapter 1), these analysis included offshore bacteriology data (total coliform and *E. coli*); however, since the onset of disinfection in August 2002, bacteriological values have become negligible and only the ammonium data are considered relative to other water quality parameters.

The permit identifies water contact standards for water within the nearshore zone, which correspond to the depth of the water column extending out 3 miles from shore (REC-1). Samples that do not meet these standards are considered out of compliance and can result in civil penalties. Compliance evaluations of offshore bacteriological samples were conducted in accordance with the following Basin Plan criteria for recreational waters (REC-1):

- Total coliform bacteria shall not exceed a geometric mean of 1000/100 mL MPN over a 30-day period provided that less than 20% of the samples exceed 1000 MPN/100 mL and no single sample can exceed

10,000/100 mL MPN when verified within 48 hours; and

- Fecal coliform bacteria shall not exceed a geometric mean of 200/100 mL MPN over a 30-day period, and not more than 10% of the samples in a 60-day period shall exceed 400/100 mL MPN.

Determinations of offshore bacteriology compliance with the fecal coliform water contact standard were accomplished by multiplying Colilert® *E. coli* data by 1.1 to obtain a calculated fecal coliform value.

## SEDIMENT GEOCHEMISTRY

### Field Methods

Sediment samples were collected for the ocean monitoring program during July and October 2010, and January and April 2011, at 10 quarterly stations located along the 60 m bottom contour (Tables A-2, Figure 4-1). Single samples were collected at all quarterly stations in each of the four surveys (Table A-5). Single samples were collected during July at 39 additional annual stations that ranged in depth from 40 to 303 m. In addition, 3 L of sediment were collected from each of the quarterly stations for sediment toxicity testing in October 2010 and April 2011.

Bottom sediments were collected using paired 0.1 m<sup>2</sup> Van Veen grab samplers. The top 2 cm of the sample was collected for individual chemical and toxicity analyses using a stainless steel scoop. The sampler and scoop were rinsed thoroughly with filtered seawater prior to sample collection. Sample storage, preservation, and holding times followed specifications in the District's QAPP, as well as guidance based on EPA/301(h) protocols. All sediment chemistry samples (metals, organics, TOC, grain size, and dissolved sulfides) were transferred to the laboratory for analysis. All sediment grain size samples were subsequently transferred to Weston Solutions, Inc. (Carlsbad, CA) for analysis. Sediment TOC samples were transferred to Columbia Analytical Services, Inc. (Kelso, WA) for analysis.

**Table A-5. Sediment composition and chemistry sampling summary.**

Orange County Sanitation District, California.

Stations	Sampling Frequency	Sampling Interval	Field Replicates	Parameters
0, 1, 4, 5, 9, 12, C, ZB, ZB2, CON	Quarterly (Summer)	0-2 cm; from undisturbed grab sample	1 replicate per station	Metals Total Organic Carbon Dissolved Sulfides Grain size PAHs LABs
0, 1, 4, 5, 9, 12, C, ZB, ZB2, CON	Quarterly (Fall, Winter, Spring)	0-2 cm; from undisturbed grab sample	1 replicate per station	Metals Total Organic Carbon Dissolved Sulfides Grain size PAHs Chlorinated Pesticides Polychlorinated Biphenyls
3, 7, 8, 10, 13, 17, 18, 20, 21, 22, 23, 24, 25, 27, 29, 30, 33, 36, 37, 38, 39, 40, 41, 42, 44, 55, 56, 57, 58, 59, 60, 61, 62, 63, 64, 65, C2, C4, C5	Annually (Summer)	0-2 cm; from undisturbed grab sample	1 replicate per station	Metals Total Organic Carbon Dissolved Sulfides Grain size PAHs LABs

All sample transfers were conducted and documented using required chain-of custody protocols through LIMS.

## Laboratory Methods

Sediment chemistry and grain size samples were processed and analyzed using performance-based and EPA-recommended methods (EPA 1986) that are listed in Table A-6. The measured sediment chemistry parameters are listed in Table A-7. Samples for dissolved sulfide were analyzed in accordance with procedures outlined in Green and Schnitker (1974) and Standard Methods 20<sup>th</sup> Edition (1998).

Sediment toxicity, following EPA-recommended methods (EPA 1994), was tested in October 2010 and April 2011, at the 10 quarterly stations using whole sediments for the 10-day *Eohaustorius estuarius* amphipod survival test. Amphipods were exposed to test and control sediments and the percent survival in each were determined. Toxicity threshold criteria were selected to be consistent with the Water Quality Control Plan for Enclosed Bays and Estuaries – Part 1 Sediment Quality (State Water Resources Control Board, California Environmental Protection Agency, 2009; Bay et al. 2009). Stations which were significantly different when compared to the control, determined by a two sample t-test, were categorized as nontoxic when survival was 90-100% of the control, lowly toxic when survival was 80-89% of the control, and moderately toxic when survival was 59-81% of the control. Stations which were not significantly different when compared to the control were categorized as nontoxic when survival was 82-100% of the control and lowly toxic when survival was 59-81% of the control. All stations exhibiting survival less than 59% of the control were categorized as highly toxic.

## Data Analyses

Linear regression was used to determine depth related factors and Pearson correlation for relationships to sediment physical attributes and chemical concentrations (e.g., total linear alkylbenzene-tLAB). Regression and correlation analyses were conducted using

MINITAB 15 (2007) statistical software. Principal components analysis (PCA) is an ordination technique used to map stations in multiple-dimensions based on the similarity of their samples (i.e., sediment chemical concentrations). Correlation-based PCA was performed using the PRIMER (2001) statistical software package on samples collected in July 2010. Test parameters incorporated the sediment physical and chemical measures determined to be the best effluent footprint indicators. These were determined statistically by Dr. Kerry Ritter (Biostatistician) of the Southern California Coastal Water Research Project (Costa Mesa, CA) using District historical marine monitoring data. The parameters used were percent fine sediments (silt+ clay), tLAB, cadmium, and zinc. Temporal trends were assessed graphically (qualitatively).

Total dichlorodipheynltrichloroethane (tDDT) represents the summed concentrations of o,p'- and p,p'- [2,4- and 4,4'-] isomers of DDD, DDE, and DDT and p,p'-DDMU, total polychlorinated biphenyls (tPCB) represents the summed concentrations of 45 congeners, and total chlorinated pesticides (tPest) represents the sum of alpha- and cis-chlordane, cis- and trans-nonachlor, hexachlorobenzene, aldrin, dieldrin, endrin, gamma-BHC, heptachlor, heptachlor epoxide, and Mirex. For summed concentrations, undetected components (i.e., concentrations below the analytical detection limits) were treated as zero. When all component concentrations were undetected, the corresponding total concentrations were assumed to be zero. Non-detected single analytes (e.g., individual metals) are assigned a value of one-half the detection limit for statistical analysis.

## BENTHIC INFAUNA

### Field Methods

The infaunal community was monitored by collecting marine sediments using paired 0.1 m<sup>2</sup> modified Van Veen sediment grab samplers. Samples were collected concurrent with sediment geochemistry samples (Tables A-2 and A-3, Figure 5-1). In July 2010, single samples were collected at each of the 10



**Table A-6. Sediment handling and analysis summary.**

Orange County Sanitation District, California.

<b>Parameter</b>	<b>Method Reference</b>	<b>Preservation</b>	<b>Container</b>	<b>Holding Time</b>
Dissolved Sulfides	Green and Schnitker (1974) Standard Methods 20 <sup>th</sup> Ed.	Freeze	Plastic container	6 months
Chlorinated Pesticides	NS&T (NOAA 1993)	Freeze	Amber glass jar	6 months (analyze within 40 days of extraction)
Grain Size	EPA 3-284 and 3550	4° C	Plastic bag	6 months
Linear Alkyl Benzenes	Eganhouse et al. (1983)	Freeze	Glass jar	12 months
Metals	NS&T (NOAA 1993)	Freeze	Amber glass jar	6 months
Polychlorinated Biphenyls	NS&T (NOAA 1993)	Freeze	Amber Glass jar	6 months (analyze within 40 days of extraction)
Polycyclic Aromatic Hydrocarbons	NS&T (NOAA 1993)	Freeze	Glass jar	6 months (analyze within 40 days of extraction)
Total Organic Carbon	EPA 415	Freeze	Glass jar	6 months

**Table A-7. Parameters measured in sediments.**

Orange County Sanitation District, California.

<b>Metals</b>		
Aluminum	Copper	Nickel
Arsenic	Iron	Selenium
Beryllium	Lead	Silver
Cadmium	Mercury	Zinc
Chromium		
<b>Other Metrics</b>		
Dissolved Sulfides	Total Organic Carbon	Grain Size
<b>Chlorinated Pesticides</b>		
Aldrin	Endosulfan sulfate	Mirex
<i>alpha</i> -BHC	Endrin	<i>trans</i> -Nonachlor
<i>alpha</i> -Chlordane	Endrin aldehyde	2,4'-DDD (o,p'-DDD)
<i>beta</i> -BHC	<i>gamma</i> -BHC	2,4'-DDE (o,p'-DDE)
<i>cis</i> -Nonachlor	<i>gamma</i> -Chlordane	2,4'-DDT (o,p'-DDT)
<i>delta</i> -BHC	Heptachlor	4,4'-DDD (p,p'-DDD)
Dieldrin	Heptachlor epoxide	4,4'-DDE (p,p'-DDE)
Endosulfan 1	Hexachlorobenzene	4,4'-DDT (p,p'-DDT)
Endosulfan 2	Methoxychlor	4,4'-DDMU
<b>PCB Congeners</b>		
PCB 8	PCB 110	PCB 167
PCB 18	PCB 114	PCB 168
PCB 28	PCB 118	PCB 169
PCB 37	PCB 119	PCB 170
PCB 44	PCB 123	PCB 177
PCB 49	PCB 126	PCB 180
PCB 52	PCB 128	PCB 183
PCB 66	PCB 138	PCB 187
PCB 70	PCB 149	PCB 189
PCB 74	PCB 151	PCB 194
PCB 77	PCB 153	PCB 195
PCB 81	PCB 153/168	PCB 200
PCB 87	PCB 156	PCB 201
PCB 99	PCB 157	PCB 206
PCB 101	PCB 158	PCB 209
PCB 105		
<b>PAH Compounds</b>		
1,6,7-Trimethylnaphthalene	Benzo[a]pyrene	Dibenzothiophene
1-Methylnaphthalene	Benzo[b]fluoranthene	Fluoranthene
1-Methylphenanthrene	Benzo[e]pyrene	Fluorene
2,6-Dimethylnaphthalene	Benzo[g,h,i]perylene	Indeno[1,2,3-c,d]pyrene
2-Methylnaphthalene	Benzo[k]fluoranthene	Naphthalene
Acenaphthene	Biphenyl	Perylene
Acenaphthylene	Chrysene	Phenanthrene
Anthracene	Dibenz[a,h]anthracene	Pyrene
Benz[a]anthracene		

**Table A-7 Continues.**

**Table A-7 Continued.**

<b>PAH Alkylated Homologues</b>		
C1-Chrysenes	C1-Fluorenes	C3-Naphthalenes
C2-Chrysenes	C2-Fluorenes	C4-Naphthalenes
C3-Chrysenes	C3-Fluorenes	C1-Phenanthrenes/Anthracenes
C4-Chrysenes	C1-Fluoranthenes/Pyrenes	C2-Phenanthrenes/Anthracenes
C1-Dibenzothiophenes	C1-Naphthalenes	C3-Phenanthrenes/Anthracenes
C2-Dibenzothiophenes	C2-Naphthalenes	C4-Phenanthrenes/Anthracenes
C3-Dibenzothiophenes		
<b>LAB Compounds</b>		
2-Phenyldecane	2-Phenyltetradecane	4-Phenylundecane
3-Phenyldecane	3-Phenyltetradecane	5-Phenylundecane
4-Phenyldecane	4-Phenyltetradecane	6-Phenylundecane
5-Phenyldecane	5-Phenyltetradecane	2-Phenyl-dodecane
2-Phenyltridecane	6-Phenyltetradecane	3-Phenyl-dodecane
3-Phenyltridecane	7-Phenyltetradecane	4-Phenyl-dodecane
4-Phenyltridecane	2-Phenylundecane	5-Phenyl-dodecane
5-Phenyltridecane	3-Phenylundecane	6-Phenyl-dodecane
7+6-Phenyltridecane		

quarterly stations and at each of the 39 annual stations (Table A-8). Three replicate samples were collected at each of the 10 quarterly stations in October 2010 and January and April 2011.

All infauna sediment samples were qualitatively and quantitatively assessed for acceptability prior to collection and processing. Samples were defined as acceptable if they had a volume of at least 4 L and a relatively undisturbed surface. It can be difficult to obtain a 4 L sediment sample at some stations due to sediment conditions (i.e., compact sediments). If three samples with volumes <4 L are encountered at a station, then samples of at least 3 L are considered acceptable and are retained for analysis. These samples were gently washed with filtered seawater through a 1.0-mm sieve. Retained organisms were placed in one liter plastic containers and anesthetized with 7% magnesium sulfate for approximately 30 minutes. The samples were then fixed in a 10% buffered formalin solution and returned to the laboratory.

### Laboratory Methods

After 3–10 days in formalin, samples were transferred to 70% ethanol for laboratory processing. Samples were sent to Weston Solutions, Inc. to be sorted to major taxonomic groups (polychaetes, molluscs, crustaceans, echinoderms, and minor phyla). Removal of the organisms from the sediments was monitored to insure that at least 95% of all organisms were successfully separated from the sediment matrix. QA/QC generally found that over 99% of all organisms had been recovered. Upon completion of sample sorting, the major taxonomic groups were distributed to qualified taxonomists for enumeration and speciation. As part of insourcing the taxonomy, OCSD's taxonomists did a portion of the identifications. These included all the July 2010 annual stations, the first replicate of each July 2010 quarterly station and all the January 2011 stations. Weston's taxonomists did all the remaining samples. Also as a result of insourcing, 10% of the January 2011 quarterly samples were sent for reanalysis by Weston's taxonomists. Taxonomic differences were resolved and the database was edited

accordingly. In general all samples met or exceeded the 90% threshold (see Appendix C). After completion of the counting and identifications, each taxonomic group was wet-weighted to the nearest 0.1 g.

### Data Analyses

Infaunal community data was analyzed to determine if populations outside the ZID were affected by the outfall discharge. Nine measures are used to assess infaunal community health and function: (1) total number of species, (2) total abundance of individuals, (3) total biomass, (4) Shannon-Wiener Diversity ( $H'$ ), (5) Margalef Species Richness (SR), (6) Schwartz' 75% Dominance Index (Dominance), (7) Species Evenness ( $J'$ ), (8) Infaunal Trophic Index (ITI; Word 1978, 1990), and (9) Benthic Response Index (BRI; Bergen et al. 1999). Shannon-Wiener Diversity and Evenness, which is the ratio of the observed Shannon-Wiener Diversity to the maximum given the same number of taxa, were calculated using  $\log_e$  (Pielou 1969). Dominance was calculated as the minimum number of species with combined abundance equal to 75% of the individuals in the sample (Swartz *et al.* 1986). Biomass measurements are sometimes influenced by the occurrence of occasional, large organisms, so they tend to be much more variable than other community measures. For that reason, organisms having large biomass (e.g., sea stars and large mollusks) are removed from the sample calculation.

Diversity values are based upon the number of species and the equitability of their distribution, but these two attributes vary independently. Consequently, a large number of diversity indices have been developed. Shannon-Wiener diversity (Shannon and Weaver, 1962) Pielou's evenness index (Pielou 1966) and Dominance (Swartz *et al.* 1986) are more sensitive to the distribution of species within a sample, while Margalef's Species Richness is more sensitive to the number of species (Tetra Tech 1985).

The presence/absence of certain pollution sensitive and pollution tolerant indicator species were also determined to further assess sediment quality in the monitoring area. The

**Table A-8. Benthic infauna sampling summary.**

Orange County Sanitation District, California.

Stations	Sampling Frequency	Sampling Volume	Field Replicates	Parameters
0, 1, 4, 5, 9, 12, C, CON, ZB, ZB2	Quarterly (Summer)	4 liters; from undisturbed grab sample	1 replicates per station	Infauna
0, 1, 4, 5, 9, 12, C, CON, ZB, ZB2	Quarterly (Fall, Winter, Spring)	4 liters; from undisturbed grab sample	3 replicates per station	Infauna
3, 7, 8, 10, 13, 17, 18, 20, 21, 22, 23, 24, 25, 27, 29, 30, 33, 36, 37, 38, 39, 40, 41, 42, 44, 55, 56, 57, 58, 59, 60, 61, 62, 63, 64, 65, C2, C4, C5	Annually (Summer)	4 liters; from undisturbed grab sample	1 replicate per station	Infauna

pollution sensitive species included the red brittlestar (*Amphiodia urtica*) and amphipods in the genera *Ampelisca* and *Rhepoxynius*, which are commonly used in whole-sediment toxicity testing. Two pollution tolerant species, *Capitella* “*capitata*” complex (polychaete) and *Euphilomedes carcharodonta* (ostracod) were examined because their presence may indicate stressed, polluted, or organically enriched environments.

Spatial pattern analysis was conducted with multivariate statistical analyses using the PRIMER (2001) multivariate statistical software package. Hierarchical cluster analysis was performed on the July 2010 species abundance data. Abundance data was 4<sup>th</sup> root transformed, a Bray-Curtis similarity matrix constructed, and a dendrogram generated using a group average cluster mode. As a confirmatory step, ordination clustering using non-metric multidimensional scaling (MDS) analysis was performed on the same data set. The data were 4<sup>th</sup> root transformed in order to down-weight the highly abundant species and incorporate the importance of the less common species (Clarke and Warwick 2001). The SIMPER (“similarity percentages”) routine was also used to determine inter- and intra-group species differences.

Multivariate cluster analysis was used to analyze spatial patterns to define similar habitats (i.e., station clusters). This technique groups those stations (habitats) having the most similar species/abundance relationships (assemblages). Cluster analysis was performed on data from July 2010 using a hierarchical agglomerative clustering method. Abundance data from the first replicate from the quarterly stations and data from the 22 shallow-, mid-, and outer-shelf annual stations were used in this analysis. The slope and basin stations were excluded from this analysis because cluster analysis should not be used when there are known environmental gradients (Clarke and Warwick 2001). Previous analyses have consistently shown that the slope, basin, and submarine canyon stations cluster apart from the shelf stations (OCSD 2010).

Correlation and regression analyses were used to test for relationships among factors (i.e., community measures vs. station depth and

sediment characteristics). Prior to analysis, the data were transformed as necessary (e.g. square-root, log<sub>10</sub>, rank, or arcsine). Where appropriate, significance was set at  $p \leq 0.05$ . Regression analysis was used to determine depth-related factors. Regression and correlation analyses were conducted using MINITAB (2007) statistical software. Temporal trends were assessed graphically (qualitatively).

## DEMERSAL FISH COMMUNITIES

### Field Methods

Demersal fish and epibenthic macroinvertebrates (EMI) species were collected in the summer (August) of 2010 and the winter (January) of 2011. Sampling was conducted at nine stations: inner shelf (36 m) stations T2 and T6; middle shelf (60 m) stations T1, T3, T11, T12, and T13; and outer shelf (137 m) stations T10 and T14 (Tables A-2 and A-3, Figure 6-1). Two replicates were conducted at the inner and outer shelf stations and three replicates were conducted at the middle shelf stations.

Trawling was conducted using a 7.6 meter (25 ft) wide, Marinovich, semi balloon otter trawl (2.54 cm mesh) with a 0.64 cm mesh cod-end liner, an 8.9-m chain-rigged foot rope, and 23 m long trawl bridles following regionally adopted methodology (Mearns and Allen 1978). The trawl wire scope varied from a ratio of approximately 5:1 at the shallowest stations to approximately 3:1 at the deepest station. To minimize catch variability due to weather and current conditions, which may affect the bottom-time duration of the trawl, trawls generally were taken along a constant depth at each station, and usually in the same direction.

Established survey methods for southern California require that a portion of the trawl track must pass within a 100-m circle that originates from the nominal sample station position and be within 10% of the station’s depth. The speed of the trawl should range from 0.77 to 1.00 m/s or from 1.5 to 2.0 kts. Since 1985, the District has trawled a set distance of 450 m (the distance that the net is actually on the bottom collecting fish and invertebrates). Station locations and trawling paths were determined using Global

Positioning System (GPS) navigation. GPS was also used to control the speed of the trawl (2.0–2.5 knots over the bottom) and determine the distance sampled (450 m). Trawl depths and time on the bottom were determined using an attached pressure sensor on the trawl boards.

Upon retrieval of the trawl net, the contents were emptied into a large flow-through water tank and all specimens were sorted by species into separate containers. First the bioaccumulation specimens were counted, recorded, and removed for processing; then the remaining specimens were measured to the nearest millimeter (standard length) and weighed to the nearest gram. A minimum of 30 (randomly selected) specimens of each species were weighed and measured individually. If a sample contained substantially more than 30 individuals of a species, the excess specimens were enumerated in 1 cm size classes and a bulk weight was recorded. All specimens were examined for external tumors, other lesions, and parasites. Specimens retained for laboratory identification were weighed and measured in the laboratory.

## Laboratory Methods

Specimens for the voucher collection and any animals that could not be identified in the field were preserved in 10% buffered formalin for subsequent laboratory analysis. A representative voucher collection of fish and macroinvertebrates is maintained for reference and verification.

## Data Analyses

Fish and EMI populations were summarized in terms of total abundance and species, percent abundance, frequency of occurrence, and mean abundance per haul. In addition, mean number of species, number of individuals, biomass, and diversity indices including Shannon-Wiener ( $H'$ ), Margalef's Species Richness (SR), and Swartz's 75% Dominance were calculated for both fish and EMI. In some analyses, stations were grouped into the following categories to assess spatial or depth-related patterns: "outfall" (Stations T1 and T12); "shallow" (Stations T2 and T6); "deep" (Stations T10 and

T14); "farfield downcoast" (Station T3); and "farfield upcoast" (Stations T11 and T13).

PRIMER (2001) multivariate statistical software was used to examine the spatial patterns of the fish assemblages in the District's monitoring area (Clarke 1993, Warwick 1993). Analyses included hierarchical clustering with group-average linking based on Bray-Curtis similarity indices, and ordination clustering of the data using non-metric multidimensional scaling (MDS). Data were truncated to include only the middle shelf (60 m) stations since depth is a strong environmental factor in delineating species clusters (OCSD 2004, 2010). Clarke and Warwick (2001) warn that clustering is less useful and may be misleading where there is a strong environmental forcing, such as depth. Prior to the calculation of the Bray-Curtis indices, the data were square-root transformed in order to down-weight the highly abundant species and incorporate the importance of the less common species (Clarke and Warwick 2001). The SIMPER ("similarity percentages") routine was also used to determine inter- and intra-group species differences.

Community measures from Stations T1 and T11 were evaluated for long-term temporal and spatial patterns, and compared with regional reference conditions, such as 1994 Southern California Bight Pilot Project (SCBPP), the Bight'98, and Bight'03 regional monitoring programs (Allen *et al.* 1998, 2002, 2007, respectively).

Fish biointegrity in the District's monitoring area was assessed using the fish response index (FRI). The FRI is a multivariate weighted-average index produced from an ordination analysis of calibrated species abundance data (see Allen *et al.* 2001, 2006). The FRI was calculated for all 9 stations in 2010-11. For a historical perspective, FRI was calculated from 1985 to 2010 for outfall Station T1 and upcoast reference Station T11.

## FISH TISSUE CONTAMINANTS

The District's permit lists three target fish species for analysis of muscle and liver tissue chemistry: English sole (*Parophrys vetulus*),

hornyhead turbot (*Pleuronichthys verticalis*), and bigmouth sole (*Hippoglossina stomata*). Whole fish analysis was performed on one species, the Pacific sanddab (*Citharichthys sordidus*) to allow for comparison of the District's fish tissue data to data from other areas in the SCB. These species were selected because they are common demersal fish of the SCB, important to recreational fisheries, and used for other bioaccumulation studies. Muscle tissues were analyzed because they reflect the effects of chronic contaminant exposures and are typically consumed by humans. Liver tissues were analyzed because they typically have high lipid content and may accumulate relatively high concentrations of lipid-soluble contaminants that have been linked to pathological conditions, and therefore reflect fish health effects.

## Field Methods

Fish were collected during trawl surveys using an otter trawl (described earlier). The sampling objective was to collect 10 individuals of each of three target species for muscle and liver tissue analysis at both outfall (T1/T12) and farfield (T11/T13) sites. Individual fish were weighed and measured in the field, placed in clean, plastic, resealable bags, and stored on wet ice in insulated coolers. Pacific sanddabs were separated into three age classes (0, 1, and 2) based upon centimeter size ranges 5–8, 9–13, and 14–16, respectively. Six individuals per size class per haul were collected for compositing in the laboratory. All samples were subsequently transferred under chain-of-custody protocols to the laboratory.

## Laboratory Methods

Individual fish were dissected and whole-fish tissues were homogenized using a blender in the laboratory under clean conditions. Muscle, liver, and whole fish tissues were analyzed for various parameters listed in Table A-9, including DDT and metabolites, chlorinated pesticides, PCBs (individual congeners), mercury, and lipids using methods consistent with NOAA National Status and Trends (NS&T) protocols (NOAA 1993). Method blanks, analytical quality control samples (duplicates,

matrix spikes, and blank spikes), and standard reference materials were prepared and analyzed with each sample batch. Mercury was quantified by cold vapor atomic absorption spectrophotometry, organochlorines were measured using dual column gas chromatography with an electron capture detector, and lipids were determined gravimetrically. All concentrations are reported on a wet weight basis.

Total dichlorodiphenyltrichloroethane (tDDT) represents the summed concentrations of o,p'- and p,p'- [2,4- and 4,4'-] isomers of DDD, DDE, and DDT and p,p'-DDMU, total polychlorinated biphenyls (tPCB) represents the summed concentrations of 45 congeners, and total chlorinated pesticides (tPest) represents the sum of alpha- and cis-chlordane, cis- and trans-nonachlor, hexachlorobenzene, aldrin, dieldrin, endrin, gamma-BHC, heptachlor, heptachlor epoxide, and Mirex. For summed concentrations, undetected components (i.e., concentrations below the analytical detection limits) were treated as zero. When all component concentrations were undetected, the corresponding total concentrations were assumed to be zero. Non-detected single analytes (e.g., individual metals) are assigned a value of one-half the detection limit for statistical analysis.

## Data Analyses

Data analysis consisted of summary statistics and qualitative comparisons only.

## FISH HEALTH

### Field Methods

Assessment of the overall health of the fish population is also required by the NPDES permit. Consequently, all fish were visually inspected for large non-mobile external parasites, lesions, tumors, and other signs of disease (e.g., skeletal deformities).

### Data Analyses

Data analysis consisted of summary statistics and qualitative comparisons only.



**Table A-9. Parameters measured in fish tissue.**

Orange County Sanitation District, California.

<b>Metals</b>		
Mercury		
<b>Chlorinated Pesticides</b>		
Aldrin	Endosulfan sulfate	Mirex
<i>alpha</i> -BHC	Endrin	<i>trans</i> -Nonachlor
<i>alpha</i> -Chlordane	Endrin aldehyde	2,4'-DDD (o,p'-DDD)
<i>beta</i> -BHC	<i>gamma</i> -BHC	2,4'-DDE (o,p'-DDE)
<i>cis</i> -Nonachlor	<i>gamma</i> -Chlordane	2,4'-DDT (o,p'-DDT)
<i>delta</i> -BHC	Heptachlor	4,4'-DDD (p,p'-DDD)
Dieldrin	Heptachlor epoxide	4,4'-DDE (p,p'-DDE)
Endosulfan 1	Hexachlorobenzene	4,4'-DDT (p,p'-DDT)
Endosulfan 2	Methoxychlor	4,4'-DDMU
<b>PCB Congeners</b>		
PCB 8	PCB 110	PCB 167
PCB 18	PCB 114	PCB 168
PCB 28	PCB 118	PCB 169
PCB 37	PCB 119	PCB 170
PCB 44	PCB 123	PCB 177
PCB 49	PCB 126	PCB 180
PCB 52	PCB 128	PCB 183
PCB 66	PCB 138	PCB 187
PCB 70	PCB 149	PCB 189
PCB 74	PCB 151	PCB 194
PCB 77	PCB 153	PCB 195
PCB 81	PCB 153/168	PCB 200
PCB 87	PCB 156	PCB 201
PCB 99	PCB 157	PCB 206
PCB 101	PCB 158	PCB 209
PCB 105		
<b>Other Parameters</b>		
Lipids		

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