APPENDIX D: Pyrethroids in Southern California Coastal Sediments

Pyrethroids in Southern California Coastal Sediments

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Abstract

Several studies have documented the potential impacts of synthetic pyrethroid pesticides in freshwater systems; however, little is known about their fate and effects in estuarine and marine environments. The goal of this study was to assess the extent and magnitude of pyrethroids in coastal embayments of the southern California Bight (SCB), USA. Using a stratified probabilistic design, 155 sediment samples were collected from 4 embayment habitats (estuaries, marinas, open bays, and ports) and analyzed for 8 common-use pyrethroids. Total pyrethroid concentrations ranged from less than 0.5 to 230 μ g/kg dry weight (area-weighted mean concentration = 5.1 ±3.1 μ g/kg) and were detected in 35% of the total SCB embayment area. Estuaries and marinas had the greatest areal extent of detectable concentrations (up to 65%) and the greatest area-weighted mean concentrations ($22.1 \pm 26.5 \mu g/kg$). Furthermore, sites with the greatest pyrethroid concentrations were located near sources of runoff from urban watersheds. Bifenthrin and cyfluthrin were detected in 32 and 15% of all samples, respectively, whereas the other 6 pyrethroids were detected in <5% of samples. Permethrin and bifenthrin had the highest concentrations at 132 and 65 μ g/kg. Toxic units (TUs) estimated for the marine amphipod Echaustorius estuarius ranged from 0 to 5.8, exceeding unity in 9 and 32% of the total and estuary habitat area, respectively. Although increased mortality of *E. estuarius* was most frequently observed in toxicity tests run on split samples from estuaries compared with other strata, there was no clear correlation between pyrethroid TUs and amphipod mortality. This suggests other mitigating factors may affect the predictive capability of the TU approach resulting from a single test species.

Introduction

Synthetic pyrethroids have become the dominant current-use pesticides in both agricultural and non-agricultural applications since the phase-out of organophosphates over the past 20 years (Amweg *et al.* 2006). Between 1999 and 2008 in five southern California coastal counties, total annual sales of

pyrethroid pesticides for professional (i.e., licensed) application increased from 65 to 153 metric tons (www.cdpr.ca.gov). However, these usage figures underestimate total pyrethroid sales as they do not include direct over-the-counter sales to consumers for residential use. In southern California, with over 17 million residents and 6 million housing units (http://quickfacts.census.gov), approximately 73% of pyrethroids had non-agricultural applications, such as structural pest control and landscape maintenance (http://www.cdpr.ca.gov).

Due to their hydrophobicity (log $K_{ow} = 5-6$) and particle reactive nature (Gan *et al.* 2005, Laskowski 2002), sediments in urban and agricultural runoff are the predominant sink of pyrethroids in California receiving waters (Ahn *et al.* 2005, Weston and Lydy 2010). Pyrethroids have been measured in sediments from California, Texas, and Illinois, and stormwater runoff has been implicated as a primary source (Amweg *et al.* 2005, 2006; Hintzen *et al.* 2009; Domagalski *et al.* 2010; Weston and Lydy 2010). In much of urbanized California, stormwater and wastewater treatment systems are separate, so wet and dry weather runoff receive no treatment before discharging to the coastal ocean. Moreover, the flood control system has been highly modified to reduce flooding (Tiefenthaler *et al.* 2008). As a result, storm flows can change by orders of magnitude in a matter of minutes and carry large quantities of sediment. For example, approximately 700,000 metric tons of suspended solids, with 89% carried by storm flow, were discharged to the coastal ocean of southern California Bight (SCB) in 1994-1995 (Schiff *et al.* 2000).

Pyrethroids can be highly toxic to non-target aquatic species (Amweg *et al.* 2006). In several California urban creeks, bifenthrin concentrations ranged from 2.19 to 219 ng/g dry weight, and most samples with greater than 0.5 toxic units (TUs) exhibited substantial toxicity to the freshwater amphipod *Hyalella azteca* (Holmes *et al.* 2008). In central Texas, pyrethroids were reported as the likely cause of toxicity to *H. azteca* in the sediments of urban streams (Hintzen *et al.* 2009). In Illinois, pyrethroids occurred up to 56 µg/kg with up to 2.9 TUs and were toxic to *H. azteca* in sediments of urban waterways (Ding *et al.* 2010).

Although pyrethroid use in California is widespread, and the potential for environmental impact is substantial, there is a limited number of comprehensive studies evaluating the occurrence, extent, or magnitude of pyrethroids in the coastal environment. Two relatively small-scale, site-specific studies indicating that concern over pyrethroid impacts in marine systems is warranted (Anderson *et al.* 2010, Lao *et al.* 2010) have been conducted in southern California marine habitats. One study in Ballona Creek (Los Angeles, CA), an intensely urbanized coastal estuary, reported pyrethroid concentrations up to 473 µg/kg dry weight and corresponding toxicity to a standard invertebrate toxicity test species *Eohaustorius estuarius* (Lao *et al.* 2010).

The first objective of the current study was to assess the extent and magnitude of pyrethroid concentrations in marine embayments across the entire SCB. The second objective was to assess the influence of receiving water classification and discharge sources by comparing pyrethroid concentrations among different habitats, including: estuaries, marinas, ports, and open bays. The third objective was to determine if a toxic-unit approach was predictive of toxicity observed for *E. estuarius* in sediment toxicity tests utilizing splits of the same samples to quantify pyrethroid concentrations.

Methods

Study Region and Sample Collection

The SCB is delineated by a 400 km length of recessed coastline between Point Conception to the north and the United States-Mexico international border to the south. A stratified-random probability-based design (Stevens, 1997) was used to conduct an unbiased survey of SCB embayments encompassing a total area of 94.1 km² and characterized by salinities greater than 30 parts-per-thousand during the sampling events. Samples were collected at 155 sites in 19 different geographic embayments (see Table SI-1 and Figure SI-1 in Supplemental Information). Sites were classified as belonging to one of four

major strata (estuaries, marinas, ports, and open bays; Bergen 1996), with more than 30 sites for each strata.

Sediment samples were collected during the regional dry season (July through September) in 2008 using a 0.1 m² modified Van Veen grab. The top 5 cm was composited. Subsamples for analysis of pyrethroids, total organic carbon (TOC), and total nitrogen (TN) were placed in pre-cleaned 250 ml amber glass containers with Teflon-lined lids and delivered to the laboratory on dry ice to be frozen at - 20°C until analysis. Subsamples for grain size analysis and toxicity testing were placed in 100 ml and 1 L pre-cleaned plastic containers, respectively, delivered to the laboratory on wet ice, and stored at ~4°C until analysis and testing.

Sample Analysis

Bifenthrin, cyfluthrin, cypermethrin, deltamethrin, esfenvalerate, fenpropathrin, λ -cyhalothrin and permethrin were targeted for analysis in this study. Sediment samples for pyrethroid measurement were extracted with dichloromethane using a microwave extraction system according to EPA method 3546. Elemental sulfur was removed from extracts using acid-activated copper powder. The extracts were further cleaned using silica gel/alumina column chromatography and analyzed using an Agilent 6890 gas chromatograph coupled to a 5973 quadrupole mass spectrometer (GC-MS) with a DB-5MS column (60 m × 0.25 mm × 0.25 µm; Agilent Technologies, Palo Alto CA). The carrier gas was helium at a flow rate of 1.5 ml/minute. The oven temperature programmed 45°C for 5 minutes, then ramped at 25°C/minute to 150°C, then ramped at 2.5°C/minute to 285°C and held for 16.8 minutes. The MS was operated in fullscan (m/z 45-500) negative chemical ionization mode at 1.67 scans/second. Quantitation was based on a 5-point internal standard calibration curve (with the lowest concentration at the reporting limit) with 2,2',5,5'-tetrabromobiphenyl as the internal standard (the quantitation and confirmation ions are listed in Table SI-2). Procedural blanks, matrix spikes, matrix spike duplicates, and replicate samples were

analyzed to validate the analytical protocol. Mean recovery of target pyrethroids in matrix spikes/duplicates were 98.6 ±11.0% (n = 14, 95% confidence interval), and 97.6 ±5.5% (n = 14), respectively. The relative percent difference (mean ± standard deviation) for target pyrethroids in sample duplicates was 8.74 ±2.95% (n=14). The method detection and reporting limits were 0.5 and 2.0 μ g/kg for individual target pyrethroids with the exception of permethrin (5.0 and 25 μ g/kg). Total organic carbon and TN were determined using a Carlo Erba 1108 CHN Elemental Analyzer, while grain size analyses were conducted using a Horiba LA900 instrument (Maruya and Schiff 2009).

Bulk sediment toxicity was measured by exposing the estuarine amphipod *E. estuarius* to split samples from SCB embayment sediments following previously published protocols (USEPA 1994). The amphipods were collected from a non-contaminated estuarine location (Beaver Creek, OR). Sediments were passed through a 2 mm sieve prior to testing to remove debris and any indigenous organisms. The exposures were conducted in 1 L glass chambers containing ~2 cm of sediment and 800 ml of filtered ($\leq 20 \mu$ m) seawater at a salinity of 32 psu. Twenty amphipods were added to each lightly aerated beaker and exposed for 10 days under constant light and 15°C. Five replicates were tested for each station. At the end of the test, the surviving amphipods were counted to determine percentage mortality relative to control exposures. A negative control (amphipod collection site sediment) was included with each testing batch of samples (Bay *et al.* 2011).

Data Analysis

Data analysis was comprised of four general steps: 1) spatial extent of detectable pyrethroids; 2) magnitude of concentrations; 3) mass inventories; and 4) toxicity potential. Pyrethroid concentration in sediment was expressed on a dry weight basis. For individual pyrethroids that were detected at estimated concentrations below the reporting limit but above the method detection limit, 1/2 of the reporting limit was used for calculation. A value of zero was used for individual pyrethroids that were not detected (i.e.,

below the method detection limit). Total pyrethroid concentration was the sum of the eight target compounds.

The extent of pyrethroid concentration was described using a cumulative distribution function (CDF), the total detectable area, percent of total area, and percent of detectable areas for individual pyrethroid. The CDF of concentration by area was calculated according to Equation 1:

$$CDF_j = \frac{\prod_{i=1}^{j} AW_i}{\prod_{i=1}^{n} AW_i}$$
Eq. 1

where CDF_j is the cumulative distribution frequency for station *j* in ascending order, AW_i is the area weight (km²) for station *i*, and *n* denotes the total number of the stations (Zeng *et al.* 2005).

The total detectable area was the sum of area weight of stations from which any targeted pyrethroid was detected. The percent of total area was the ratio of the total detectable area to the total area. Percent of detectable area for individual pyrethroid was the ratio of its detectable area to the total detectable area. Permethrin was excluded in the CDF calculation due to its substantially higher reporting limit. The area and stratum calculations for total pyrethroids were not affected by the exclusion of permethrin because the other targeted pyrethroids were also detected at relatively high levels (>15 μ g/kg) in the same sediments.

Area weighted mean (AWM) concentration and associated 95% confidence interval (95% CI) were computed according to Equations 2 and 3, respectively (Thompson 2000):

$$AWM = \frac{\prod_{i=1}^{n} (p_i \times AW_i)}{\prod_{i=1}^{n} AW_i}$$
Eq. 2

95% CI = 1.96 ×
$$\frac{\frac{n}{i=1}[(p_i - AWM) \times AW_i)]^2}{(\frac{n}{i=1}AW_i)^2}$$
 Eq. 3

where p_i is pyrethroid concentration at station *i*; AWM were calculated by stratum.

The total mass inventory of pyrethroids in surficial sediments by embayment stratum was calculated according to Equation 4:

$$Mass = AWM \times \delta \times A \times T$$
 Eq. 4

where δ is dry density of sediment (estimated 1.5 g/ml; Maruya and Schiff 2009), *A* is total area of stratum represented by the samples, and *T* is the thickness of sediment (2 cm here).

A disproportionate accumulation factor (DAF) of pyrethroids for each stratum was estimated by Equation 5:

$$DAF = \frac{\%Mass}{\%Area}$$
 Eq. 5

A DAF = 1 indicates mass accumulation in direct proportion with the stratum area.

Toxicity potential was calculated using a toxic-unit approach (Amweg *et al.* 2005). Pyrethroidspecific TUs were calculated for a given sample by dividing the organic carbon normalized pyrethroid concentration by its organic carbon normalized median lethal concentration (LC_{50}). Total pyrethroid TU was the sum of individual TU. Ten-day sediment organic carbon normalized LC_{50} values for *E. estuarius* were bifenthrin = 1.03 µg/g OC, cypermethrin = 1.41 µg/g OC, permethrin = 17.9 µg/g (Anderson *et al.* 2008), and cyfluthrin = 0.33 µg/g OC (Bay *et al.* 2010). No LC_{50} values are currently available for the remaining target pyrethroids. AWM and 95% CI TUs were calculated by substituting TU for pyrethroid concentration in Equations 2 and 3. Associations among pyrethroid concentration, TOC, TN and grain size, percentage mortality and TU were assessed using Spearman's rank correlational analyses (SigmaState; V2.03, SPSS).

Results and Discussion

Total pyrethroid concentrations were correlated with TOC ($r_s = 0.59$; p < 0.01; n = 52) and weakly correlated with TN ($r_s = 0.30$; p < 0.05; Table SI-3). This is not unexpected because pyrethroids are hydrophobic and have a strong affinity for organic phase (Gan *et al.* 2005, Laskowski 2002). There was no significant relationship between pyrethroid sediment concentrations and percent fine grained sediments ($< 63 \mu$ m), in contrast to a previous study (Lao *et al.* 2010) that observed targeted stations located a short distance from the mouth of Ballona Creek. This inconsistency could be the result of proximity to runoff sources and/or rapid transformation of many pyrethroids under estuarine/marine conditions, where hydrolysis rates are orders of magnitude greater than in freshwater systems (Laskowski 2002). For example, the eight target pyrethroids were stable (half-life >600 days) at a pH of ~5, while their half-lives ranged from ~4 to ~35 days in seawater with a pH of ~8 (W. Lao, unpublished data).

Extent by Area and Stratum

Pyrethroids were detected in 34.5% (32.5 km²) of the sediments in SCB embayments (Figure 1). Approximately 9% of the sediments in SCB embayments had concentrations >10 μ g/kg. While pyrethroids were detected in all 19 of the embayments sampled, the extent of detectable pyrethroid concentrations was dissimilar among strata (Table 1). The areal extent of detectable sediment pyrethroid concentrations was greatest in the marina stratum (65.0%), followed by estuaries (49.4%), open bays (35.8%), and ports (16.2%).

The greatest pyrethroid concentrations were observed at the mouths of urban watersheds, including: Ballona Creek near Marina del Rey in the Los Angeles metropolitan area; Dominguez Channel and the Los Angeles River which both empty into the Los Angeles/Long Beach Harbor complex; and San Diego and Chollas Creeks which empty into the San Diego Bay (Figures 2 and SI-2). Decreasing concentrations were often observed moving away from these creek mouths into their respective embayments. This is consistent with findings from a previous field survey at the Ballona Creek estuary (Lao *et al.* 2010) where pyrethroids discharged by the creek accumulated in the estuarine sediments during the dry season. In addition, the marinas contaminated by pyrethroid pesticides (e.g., Marina del Rey, Consolidated Slip in LA Harbor, Dana Point Marina, and San Diego South Bay Marina) were located adjacent to discharges from urban watersheds. Because of the absence of major point sources (e.g., industrial or municipal plants) in these watersheds, urban runoff is the most likely source of pyrethroids to SCB embayments.

Bifenthrin was the most widely distributed pyrethroid; it was present in over 95% of the samples that had detectable pyrethroid concentrations (Table 1). For samples with detectable pyrethroids, bifenthrin was present for 100% of the bay and estuary strata, 97.8% for marinas, and 69.3% for ports. Cyfluthrin was the next most prevalent pyrethroid (44% for the SCB embayment area). As with bifenthrin, cyfluthrin was most frequently detectable in estuaries and marinas. In contrast, cypermethrin was less prevalent in estuaries and marinas and not detected in the port stratum. Permethrin was detected in only 1.9% of the SCB embayment area; however, it should be noted that the 25 μ g/kg reporting limit was 10 times higher than reporting limits for the other targeted pyrethroids. Still, this low detection percentage is in contrast with a recent study (Lao *et al.* 2010) suggesting that permethrin distribution can be wide with concentrations up to two times higher than those of the next most abundant pyrethroid. The remaining four pyrethroids (deltamethrin, esfenvalerate, fenpropathrin, and λ -cyhalothrin) were detected in <2% of the SCB embayment area.

Magnitude and Relative Abundance

Sediment pyrethroid concentrations ranged from not detectable to 230 µg/kg. The overall AWM concentration ($\pm 95\%$ CI) for SCB embayments was 5.15 ± 3.09 µg/kg. The greatest pyrethroid AWM concentrations were observed in the estuarine (22.1 ± 26.5 µg/kg) and marina (20.1 ± 17.5 µg/kg) strata; AWM concentrations in open bays (2.80 ± 3.31 µg/kg) and ports (0.229 ± 0.177 µg/kg) were one to two

orders of magnitude lower (Figure 3). The spatial distribution of pyrethroids among embayment strata was dissimilar from the distribution of other persistent organic contaminants such as dichloro-diphenyl-trichloroethane (DDT) and its degradation products, polychlorinated biphenyl (PCB) congeners, and polycyclic aromatic hydrocarbons (PAHs). For example, significant differences in AWM concentrations of total DDTs, PCBs, or PAHs were observed between estuaries and marinas (Maruya and Schiff 2009).

Bifenthrin (37.5% of the pyrethroid AWM concentration) and permethrin (47.3%) were the most abundant compounds in SCB embayments (Table 2). Bifenthrin was the most abundant pyrethroid in open bays, ports and estuaries, but permethrin comprised the majority of the total pyrethroid concentration in marinas. Cyfluthrin comprised a large portion (38%) of the pyrethroid AWM concentration in the port stratum. Contribution from each of the remaining four pyrethroids was <1% in the SCB embayment area.

The relative composition of the eight target pyrethroids did not correspond to the expected distribution based on application rates in SCB watersheds and assuming a 1:1 relationship between pyrethroid sales and usage (Table 2). California Department of Pesticide Regulation (CDPR) data on active ingredient application rates for agricultural and pest control applications indicate that bifenthrin comprised 9% of the eight target pyrethroids sales in the SCB region, yet this study found that bifenthrin comprised 38% of the estimated total pyrethroids in SCB sediments. This disparity between application and occurrence of bifenthrin was amplified within individual strata (i.e., ports). Cyfluthrin presented similar but less significant differences. Occurrence to application rate agreement was closest for permethrin (35 to 59% AWM vs. 62% usage), noting that occurrence was likely underestimated due to higher variability in its reporting limit. In contrast, the relative contribution of the remaining four pyrethroids in sediment was smaller than corresponding application rates.

There were several potential reasons for the differences between reported application rates and occurrence in SCB embayment sediments (Table 2). First, pyrethroids are highly hydrophobic and have a

strong affinity for solid phases (i.e., soils, sediments, and concrete), which may result in retention in the upper parts of the watershed. Only a small fraction of the total sediment-associated pyrethroid pool is available to be mobilized and subsequently transported out of the watershed via stormwater runoff (Gan *et al.* 2005, Ortiz-Pérez *et al.* 2005, Jiang *et al.* 2011). Further, the sediment-associated pyrethroids could be retained in detention ponds, storm drains, or other catchments in the watershed (Budd *et al.* 2009). Second, the individual pyrethroids targeted in this study exhibit differential persistence in aquatic systems, have different hydrolysis and photolysis rates (Laskowski 2002), and undergo rapid hydrolysis in alkaline aqueous media, such as seawater, at different rates (Camilleri 1984; Takahashi *et al.* 1985a,b). For example, the half-lives for bifenthrin and cyfluthrin have been estimated at 35.0 \pm 0.08 and 4.44 \pm 1.7 days in seawater, respectively (W. Lao, unpublished data). Third, the reported commercial sales rates may not reflect actual usage, because direct over-the-counter sales of pyrethroids to consumers are not reported.

Bifenthrin and permethrin concentrations from this study were similar to concentrations in marine sediments, but lower than concentrations measured in freshwater sediments reported previously. The highest concentrations of bifenthrin (64.8 µg/kg) and permethrin (132 µg/kg) in sediments from the Ballona Creek estuary (this study) were comparable to those reported for the same estuary by Lao *et al.* (2010). Estuarine sediments from Switzer Creek in San Diego Bay had bifenthrin and permethrin concentrations of 23.9 µg/kg and 135 µg/kg, respectively (Anderson *et al.* 2010). In contrast, higher bifenthrin concentrations were reported in freshwater sediments, for instance, an urban wetland in Los Angeles (Sims Pond, 610 µg/kg; Brown *et al.* 2010), San Diego Creek in Orange county (542 µg/kg; Budd *et al.* 2007), and California's Central Valley (Clover Creek, 219 µg/kg; Holmes *et al.* 2008). In addition, sediment from a residential runoff drain near Sacramento, CA, contained 744 µg/kg bifenthrin and 539 µg/kg permethrin (Weston *et al.* 2009). Relative to freshwater sediments, the lower pyrethroid concentrations in marine sediments are partly due to localized differences in loading and persistence,

losses during transport, and dilution and flushing due to tidal exchange (Camilleri 1984, Takahashi *et al.* 1985b, Gan *et al.* 2005, Lao *et al.* 2010, Weston and Lydy 2010).

Mass Inventories

The total mass of the eight target pyrethroids in SCB embayment sediments was estimated to be 36 kg, assuming identical conditions of surface sediment deposition and conservation in the embayment. This mass represents a miniscule fraction $(0.036\% \pm 0.003\%)$ of the quantity of pyrethroid sales in the SCB region, which were estimated to be 101 ± 10 mt in 2007-08 (www.cdpr.ca.gov). To estimate pyrethroid loading to SCB embayments via stormwater runoff, annual stormwater runoff volumes were multiplied by average stormwater pyrethroid concentrations determined by a regional consortium of stormwater agencies (Table SI-4). The mass of pyrethroids estimated to reside in SCB embayment sediments was only 27% of the total annual loading estimated from stormwater runoff, assuming no previous accumulation of pyrethroids in the sediments. Even without assuming degradation, a large fraction of pyrethroids likely resides in freshwater sediments upstream of SCB embayments, which is consistent with previous reports of higher concentrations in freshwater sediments (Brown *et al.* 2010, Jiang *et al.* 2011). Another explanation for the lack of mass balance in SCB embayments is the advection of stormwater inputs out of estuaries and bays, as evidenced by the presence of particulate-laden freshwater plumes extending as far as 30 km offshore (Ahn *et al.* 2005).

Of the estimated 36 kg of pyrethroids that reside in SCB embayment sediments, the greatest mass (46%) occurred in marinas and the least mass (1%) in ports (Table 3). However, a disproportionately greater accumulation of pyrethroids was observed in both the marina and estuary strata relative to their areas. The DAF values indicated that four times the mass resides in marina and estuary strata relative to their area. In contrast, bays accumulated approximately half of the mass expected based on an equal distribution according to area.

Toxicity Potential

Toxic units based on individual pyrethroid LC50s derived for *E. estuarius* ranged from 0 to 5.8 (Figure 4), and 14 of the 155 samples had TUs greater than unity. The maximum TU was calculated for a sample from the Ballona Creek estuary. In a sediment sample from a previous study in San Diego Bay (Anderson *et al.* 2010), calculated pyrethroid TU (sum of bifenthrin, permethrin and cyfluthrin) based on *E. estuarius* was 21.7. In another pilot study of sediments from Ballona Creek (Lao *et al.* 2010), the calculated pyrethroid TU (sum of bifenthrin, cyfluthrin and cypermethrin) based on *H. azteca* ranged from 1.1 to 29.8 and exhibited high spatial and temporal variations. Thus, using these TUs and *H. azteca* LC50 data, the pyrethroid TU based on *E. estuarius* could be estimated ranging from 0.59 to 31.2. Notably, the maximum TU for the current study was lower than that for the two earlier site-specific studies, reflecting spatial and temporal variability in coastal environments.

Approximately 9% of the total SCB embayment area, and a relatively large extent of the estuarine (32.2%) and marina (26.2%) strata compared to port (0.3%) and open bay (7.9%) strata, had a TU greater than unity. Furthermore, TUs exceeding unity were in 26.7% total pyrethroid detectable-area. Similarly, within the pyrethroid-detectable area of each stratum, TUs exceeding unity were in 64.4, 40.3, 22.1, and 2.1% for estuary, marina, bay, and port strata, respectively. The estuary stratum had the greatest AWM TU (0.98 \pm 0.82) of the four strata (Figure 4).

Bifenthrin was the dominant contributor to the pyrethroid TUs in SCB embayment sediments (Table 2). In contrast to being a dominant component in total pyrethroid concentration, permethrin had negligible contribution to the TUs due to its relative low toxicity (17 and 54 times less toxic than bifenthrin and cyfluthrin, respectively, based on LC50 values for *E. estuarius*). Cyfluthrin was the second highest contributor to TUs and approximately three times more toxic than bifenthrin.

The Ballona Creek estuary sediment sample with the highest TU also exhibited a high degree of toxicity based on *E. estuarius* 10-day mortality (97%). However, the overall predicted toxicity based on TUs did not correlate to observed sediment toxicity (% mortality) for *E. estuarius* ($r_s = 0.129$; p = 0.110; n = 155; Figure 5). Amphipod mortality was $\leq 25\%$ relative to controls for 13 sediments with TUs that ranged from 1.0 to 3.2 with no clear relationship between total pyrethroid TUs and *E. estuarius* toxicity. In contrast, *E. estuarius* mortality ranged from 0 to 41% relative to controls when pyrethroids were not detectable (TU = 0).

Several factors may have contributed to the lack of correlation between predicted and observed toxicity in this study. Where toxicity was observed, but not predicted based on pyrethroid TUs, the presence of other contaminants may have acted as toxicants. Estuarine and marina sediments in the SCB are known to have a complex mixture of organic and inorganic toxicants, including petroleum, chlorinated hydrocarbons, and trace metals (Tiefenthaler *et al.* 2008). Anderson *et al.* (2010) reported that pyrethroids only partially contributed to the sediment toxicity observed for *E. estuarius* in San Diego Bay based on toxicity identification evaluation procedures. Other investigators have found toxicity in SCB embayments prior to the onset of widespread pyrethroid applications (Fairey *et al.* 1998).

Both chemical and toxicological factors may have contributed to instances where toxicity was predicted but not observed. Uncertainties in both the chemical and toxicological test methods may have contributed to variability as great as $\pm 30\%$. Although a comprehensive quality assurance and quality control approach that included analysis of blanks and matrix spikes was instituted, the lack of an appropriate standard or certified reference sediment material makes it difficult to fully validate any pyrethroid method, particularly at low (μ g/kg) concentrations. The LC₅₀ values used to generate TUs may also contribute uncertainty due to the limited scope of sediments utilized. For example, LC₅₀ values of bifenthrin, permethrin and cypermethrin for *E. estuarius* were determined in formulated sediment (salinity = 20%; TOC = 0.78%) that may not accurately represent chemical partitioning among dissolved and particulate phases (Anderson *et al.* 2010) or the diversity of natural sediment encountered in this study.

While this is somewhat accounted for by TOC normalization, there may be other ameliorating factors in natural sediment (i.e., presence of pore water dissolved organic carbon) that were not present during the initial dosing experiments with *E. estuarius*. Furthermore, calculating LC_{50} values from freely dissolved porewater concentration estimates has been shown to yield results with lower variability (Hawthorne *et al.* 2005, Xu *et al.* 2007). Another complicating factor in using laboratory-derived LC_{50} 's for predicting *in situ E. estuarius* toxicity may be temperature or the presence of antagonistic/synergistic compounds (i.e., piperonyl butoxide) that are known to alter the toxicity of pyrethroids (Weston *et al.* 2006). Moreover, variability could be increased based on differences in potency for chiral enantiomers of the same parent pyrethroid compound (Liu *et al.* 2005).

One final confounding factor may be the bioavailability of pyrethroids in urban sediments (You *et al.* 2008, Hunter *et al.* 2009). Bondarenko *et al.* (2007) reported that the freely dissolved (or "bioavailable") concentration of pyrethroids using solid-phase microextraction (SPME) was only a small fraction of the total pore-water concentration determined by liquid–liquid extraction in a marine sediment. Slightly negative correlations between pyrethroid availability and black carbon content in sediment have also been observed (Yang *et al.* 2009). In general, little is currently known of the true pyrethroid bioavailability across aquatic sediments, particularly sediments that have been profoundly impacted by human activity.

The inconsistencies observed in the current study's attempts to correlate predicted pyrethroid toxicity to laboratory-measured toxicity have also been observed by other studies. For example, both high (>80%) and low (<30%) *E. estuarius* mortality was observed in urban estuarine sediments with pyrethroid TUs less than 5, possibly due to the presence of other toxicants (Lao *et al.* 2010). Other earlier studies observed poor relationships between toxicity to the freshwater amphipod, *H. azteca* and predicted sediment toxicity with pyrethroid TUs between 1 and 5 (Amweg *et al.* 2005, 2006; Weston and Lydy 2010). Further, in Hintzen *et al.* (2009) <30% *H. azteca* mortality was observed after exposure to sediments from an urban stream in central Texas with pyrethroid TUs ranging between 1 and 3. On the

other hand, most studies to date have found that sediments with pyrethroid TUs >5 are nearly always highly toxic to amphipods (Weston *et al.* 2005; Amweg *et al.* 2005, 2006; Hintzen *et al.* 2009; Domagalski *et al.* 2010; Lao *et al.* 2010).

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Figure 1. Cumulative distribution function of total pyrethroid concentration (excluding permethrin) versus area in Southern California Bight embayments (94.1 km²).





Figure 2. Geographical distribution of total pyrethroid concentrations in sediments from embayments of the Southern California Bight.



Figure 3. Area-weighted mean (AWM) concentrations of total pyrethroid concentration and associated 95% confidence intervals by stratum.



Figure 4. Box plots of toxic units (TUs) for total pyrethroid concentrations (sum of bifenthrin, cyfluthrin, cypermethrin and permethrin) by stratum based on the estuarine amphipod *E. estuarius*. Boxes represent the median, 25^{th} and 75^{th} percentiles, 10^{th} and 90^{th} percentiles, and individual sites beyond the 10^{th} and 90^{th} percentile of TUs. Also shown in square symbols is the area-weighted mean TU of total pyrethroid concentration for each stratum. The dashed reference line represents TU=1.



Figure 5. Percent mortality of *E. estuarius* in 10-day whole sediment toxicity tests versus TUs for total pyrethroid concentration (sum of TUs for bifenthrin, cyfluthrin, cypermethrin and permethrin) from sediments collected in embayments of the Southern California Bight.

Stratum	Somela ciza	Total area	Pyrethroi	ds detectable area ^a	Bif ^b	Cyf	Сур
Stratum	Sample size	(km ²)	$(\mathrm{km}^2)^{\mathrm{c}}$	(% of total area)	(% of c	letectabl	e area)
Bay	29	51.7	18.5	35.8	100	33.1	22.0
Port	42	26.3	4.26	16.2	69.3	32.8	0
Marina	35	11.1	7.21	65.0	97.8	72.7	30.2
Estuary	49	4.99	2.47	49.4	100	63.0	51.8
Total	155	94.1	32.5	34.5	95.5	44.1	23.2

Table 1. Extent of detectable pyrethroids in SCB embayment sediments.

^a Sum of the seven pyrethroids; ^b Bif = bifenthrin, Cyf = cyfluthrin, Cpy = cypermethrin; ^c Used for calculating % of detectable area. Note: permethrin was excluded in this analysis due to its higher reporting limit

		Percentage of total pyrethroids (%)											
	Ва	Bay		rt	Mar	ina	Estu	ary	Entire	area	Usage ^a		
	AW M	TU s	AW M	TU s	AW M	TU s	AW M	TU s	AW M	TU s			
Bifenthrin	46	55	62	40	28	47	45	86	38	57	9.2 ± 2.5		
Cyfluthrin	9.7	36	38	60	7.9	44	3.7	10	7.8	38	4.9 ± 1.7		
Cypermethrin	9.0	8.1	0	0	4.2	4.7	5.0	2.4	5.8	3.8	16 ± 5.0		
Permethrin ^b	35	1.9	0	0	59	4.6	42	1.5	47	2.0	62 ± 4.4		
Other pyrethroids	0		0		1.4		4.4		1.7		8.4 ± 3.4		

Table 2. Relative percentage contributions of targeted pyrethroids to the total average weighted mean(AWM) concentration and toxic units (TUs) in SCB embayments sediments.

^a Average annual (1999-2008) use (\pm standard deviation) in the five coastal counties (Santa Barbara, Ventura, Los Angeles, Orange, and San Diego) of southern California (USA) (www.cdpr.ca.gov); ^b reporting limit for permethrin was ten times higher than for the other targeted pyrethroids. ^c Sum of deltamethrin, esfenvalerate, fenpropathrin, λ -cyhalothrin.

 Table 3. Disproportionate accumulation factors (DAF) for total pyrethroid mass in sediments from

 embayments in the Southern California Bight.

	Area	Mass	
	(%)	(%)	DAF
Bay	55.0	29.9	0.54
Port	27.9	1.2	0.04
Marina	11.8	46.1	3.91
Estuary	5.3	22.8	4.29

SUPPLEMENTAL INFORMATION

Pyrethroids in Southern California Coastal Sediments

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The following is the list of supporting materials for the article

Table SI-1. Sampling information and sample characteristics

Table SI-2. Quantitation and confirmation ions for pyrethroid analysis on GC-NCI/MS

Table SI-3. Pyrethroids concentrations correlation with other parameters in 52 sediments (Spearman Rank Order correlation coefficient and p value)

Table SI-4. Comparison of pyrethroid found and usage

Figure SI-1. Map of study region and sampling sites.

Figure SI-2. Sum of pyrethroid concentration in each region.

StationID	SampleDate	Stratum	Region	Latitude	Longitude	Area- weight	Depth	тос	TN	%Fines
						Km2	m	(%)	(%)	(%)
6462	9/10/2008	Вау	LA/LB harbor	33.75095	-118.17611	2.041	11	1.75	0.076	72.4
6448	9/9/2008	Вау	LA/LB harbor	33.74448	-118.16908	2.041	13	1.18	0.082	89.3
6437	9/9/2008	Вау	LA/LB harbor	33.73968	-118.1718	2.041	14	1.52	0.092	90.5
6432	9/18/2008	Вау	LA/LB harbor	33.7358	-118.17995	2.041	17	0.45	0.055	61.8
6416	7/17/2008	Вау	LA/LB harbor	33.73126	-118.2237	2.041	12	0.26	0.028	56.3
6404	7/24/2008	Вау	LA/LB harbor	33.72446	-118.22395	2.041	17	0.96	0.098	85
6387	7/22/2008	Вау	LA/LB harbor	33.71345	-118.24183	2.041	23	1.3	0.108	82
6386	7/22/2008	Вау	LA/LB harbor	33.7122	-118.25826	2.041	16	1.29	0.085	82.4
6384	7/23/2008	Вау	LA/LB harbor	33.70903	-118.26183	2.041	5	0.44	0.044	70.1
6383	7/23/2008	Вау	LA/LB harbor	33.70733	-118.26903	2.041	4	0.09	0.022	35.2
6223	8/5/2008	Вау	Mission bay	32.79425	-117.22039	0.02	1.52	3.33	0.169	66.9
6219	8/5/2008	Вау	Mission bay	32.78738	-117.20924	0.02	2.89	2.24	0.166	79.1
6217	8/5/2008	Вау	Mission bay	32.78437	-117.21529	6.231	3.56	2.62	0.157	89.6
6212	8/5/2008	Вау	Mission bay	32.76779	-117.24127	0.567	6.21	0.08	0.01	8.4
6152	8/7/2008	Вау	San Diego bay	32.71488	-117.18292	2.009	12.28	0.43	0.056	52.3
6136	8/8/2008	Вау	San Diego bay	32.69962	-117.16085	0.032	14.05	0.56	0.1	43
6134	8/8/2008	Bay	San Diego bay	32.69827	-117.15841	0.032	13.74	0.51	0.06	40.4

Table SI-1. Sampling information and sample characteristics

6093	8/19/2008	Вау	San Diego bay	32.67539	-117.14381	2.009	4.6	0.54	0.051	39.5
6090	8/19/2008	Bay	San Diego bay	32.67347	-117.13642	2.609	4.54	0.84	0.07	44.7
6084	8/19/2008	Bay	San Diego bay	32.67038	-117.13647	2.009	4.81	0.22	0.046	31.5
6083	8/21/2008	Bay	San Diego bay	32.67027	-117.15476	2.609	4.41	0.8	0.076	81.7
6080	8/21/2008	Bay	San Diego bay	32.66492	-117.1498	2.609	4.57	0.55	0.055	53.2
6071	8/21/2008	Bay	San Diego bay	32.65832	-117.14422	2.609	5.09	0.99	0.093	78.7
6041	8/22/2008	Bay	San Diego bay	32.6475	-117.11686	0.041	10.75	0.8	0.05	55.8
6040	8/21/2008	Bay	San Diego bay	32.64724	-117.11783	0.041	11.21	0.79	0.064	57.4
6039	8/21/2008	Bay	San Diego bay	32.64686	-117.11961	0.041	12.64	0.68	0.046	61.4
6031	8/22/2008	Bay	San Diego bay	32.63248	-117.13566	2.609	1.79	1.12	0.062	65.3
6017	8/22/2008	Вау	San Diego bay	32.60837	-117.1114	2.609	1.52	0.35	0.02	50.3
6015	8/22/2008	Вау	San Diego bay	32.60756	-117.12241	2.609	1.82	1.82	0.124	76.7
6363	8/8/2008	Estuary	Upper Newport bay	33.64682	-117.88411	0.484	3	1.1	0.086	82.1
6362	8/8/2008	Estuary	Upper Newport bay	33.64574	-117.8888	0.484	3.9	1.47	0.117	96.2
6282	7/8/2008	Estuary	Agua Hedionda lagoon	33.1447	-117.33563	0.179	2.4	0.61	0.043	20.7
6280	7/8/2008	Estuary	Agua Hedionda lagoon	33.14456	-117.32811	0.248	2.74	0.57	0.065	49.3
6271	7/8/2008	Estuary	Agua Hedionda lagoon	33.14016	-117.3251	0.228	2.36	1.01	0.098	85.6
6270	7/8/2008	Estuary	Agua Hedionda lagoon	33.1396	-117.3186	0.17	0.6	0.64	0.049	38.6
6269	7/8/2008	Estuary	Agua Hedionda lagoon	33.13921	-117.3377	0.219	6.7	0.52	0.061	48.5
6264	7/29/2008	Estuary	Batiquitos lagoon	33.0906	-117.2872	0.18	0.45	1.11	0.094	92.8
6252	7/29/2008	Estuary	Batiquitos lagoon	33.0885	-117.2726	0.139	1.52	1.69	0.167	95.5

6253	7/29/2008	Estuary	Batiquitos lagoon	33.0885	-117.30495	0.216	1.61	0.43	0.045	29.2
6251	7/29/2008	Estuary	Batiquitos lagoon	33.08806	-117.3105	0.186	1.68	0.03	0.008	0.1
6250	7/29/2008	Estuary	Batiquitos lagoon	33.08775	-117.29285	0.206	0.57	0.32	0.039	70.5
6200	7/11/2008	Estuary	Mission bay	32.76053	-117.21014	0.015	1.7	0.45	0.075	56.4
6197	7/11/2008	Estuary	Mission bay	32.76008	-117.22071	0.034	0.54	0.23	0.046	33.9
6192	7/11/2008	Estuary	Mission bay	32.7579	-117.22693	0.166	0.53	1.3	0.065	60.7
6189	7/11/2008	Estuary	Mission bay	32.75766	-117.242	0.084	0.3	0.08	0.024	19
6181	7/11/2008	Estuary	Mission bay	32.75679	-117.23502	0.172	1.1	2.49	0.173	67.7
6069	7/10/2008	Estuary	Sweetwater river	32.65664	-117.08712	0.014	1.5	0.24	0.116	2.9
6065	7/10/2008	Estuary	Sweetwater river	32.65503	-117.09129	0.023	1.2	1.12	0.085	60.7
6060	7/10/2008	Estuary	Sweetwater river	32.65368	-117.09492	0.038	1.03	1.15	0.111	68.8
6057	7/10/2008	Estuary	Sweetwater river	32.65211	-117.10048	0.035	1.67	1.77	0.122	78.4
6052	7/10/2008	Estuary	Sweetwater river	32.65109	-117.1031	0.041	2.28	1.29	0.086	71.5
6047	8/22/2008	Estuary	Sweetwater river	32.64925	-117.11	0.026	4.57	1.05	0.089	75.3
6049	8/21/2008	Estuary	Sweetwater river	32.64889	-117.11284	0.026	10.75	1.05	0.084	67.7
6046	8/22/2008	Estuary	Sweetwater river	32.64879	-117.11388	0.026	11.82	0.72	0.024	56.8
6045	8/22/2008	Estuary	Sweetwater river	32.64831	-117.11611	0.026	10.97	1.09	0.081	67.1
6553	9/3/2008	Estuary	Channel island	34.18464	-119.23089	0.309	3.81	2.16	0.19	92.4
6520	9/11/2008	Estuary	Ballona creek	33.97125	-118.43955	0.309	2.3	0.09	0.543	58.3
6508	9/29/2008	Estuary	Ballona creek	33.96281	-118.45421	0.309	3.3	5.07	0.074	48.3
6317	8/29/2008	Estuary	Oceanside north	33.23557	-117.40565	0.014	0.91	0.02	0.004	3.4

6314	8/29/2008	Estuary	Oceanside north	33.2354	-117.40745	0.021	1.06	0.04	0.008	5.9
6311	8/29/2008	Estuary	Oceanside north	33.2344	-117.409	0.024	0.91	1.26	0.122	48.9
6308	8/29/2008	Estuary	Oceanside north	33.23342	-117.41152	0.033	0.76	1.06	0.023	44
6303	8/29/2008	Estuary	Oceanside north	33.23226	-117.4132	0.083	0.61	0.01	0.003	0
6245	7/15/2008	Estuary	San Elijo lagoon	33.01426	-117.2769	0.008	2.28	0.14	0.013	7.1
6244	7/15/2008	Estuary	San Elijo lagoon	33.01413	-117.27967	0.017	0.95	0.14	0.009	1
6243	7/15/2008	Estuary	San Elijo lagoon	33.0113	-117.27323	0.01	0.68	0.58	0.067	24
6242	7/22/2008	Estuary	San Elijo lagoon	33.0106	-117.26359	0.011	2.43	2.86	0.22	80.7
6239	7/15/2008	Estuary	San Elijo lagoon	33.008	-117.27063	0.012	0.4	0.26	0.02	18
6236	7/17/2008	Estuary	Soledad creek	32.93378	-117.25683	0.018	1.4	1.13	0.093	46.1
6232	7/17/2008	Estuary	Soledad creek	32.9328	-117.2579	0.016	0.64	0.04	0.006	3.3
6230	7/17/2008	Estuary	Soledad creek	32.9321	-117.25336	0.009	0.39	1.51	0.142	79.1
6229	7/17/2008	Estuary	Soledad creek	32.93155	-117.25097	0.01	1.52	1.18	0.128	81.1
6228	7/17/2008	Estuary	Soledad creek	32.93043	-117.24855	0.007	0.45	0.62	0.096	74.9
6012	7/16/2008	Estuary	Tijuana river estuary	32.56244	-117.10841	0.011	0.8	0.35	0.047	4
6010	7/16/2008	Estuary	Tijuana river estuary	32.55935	-117.11118	0.012	0.6	0.06	0.008	3
6009	7/16/2008	Estuary	Tijuana river estuary	32.5592	-117.11603	0.012	0.64	0.38	0.046	9.5
6004	7/16/2008	Estuary	Tijuana river estuary	32.55737	-117.12239	0.025	0.79	0.18	0.03	13.7
6001	7/16/2008	Estuary	Tijuana river estuary	32.55659	-117.128	0.048	0.85	0.11	0.003	0
6560	9/3/2008	Marina	Ventura west marina	34.24868	-119.26406	0.726	4.2	1	0.076	98.7
6549	9/3/2008	Marina	Channel island	34.17122	-119.22351	0.726	4.2	2.47	0.153	90.7

6530	9/29/2008	Marina	Marina del Rey	33.98308	-118.45068	0.726	3.8	1.42	0.121	99.2
6527	9/29/2008	Marina	Marina del Rey	33.98048	-118.4422	0.726	4.4	1.55	0.101	99.5
6649	9/17/2008	Marina	Marina del Rey	33.97773	-118.45265	0.726	3	1.21	0.068	85.9
6518	9/29/2008	Marina	Marina del Rey	33.97023	-118.44804	0.726	6.8	2.16	0.109	82.8
6513	9/29/2008	Marina	Marina del Rey	33.96469	-118.45333	0.726	7	3.07	0.112	67.6
6489	7/30/2008	Marina	LA/LB harbor	33.76671	-118.2485	0.726	4	3.71	0.139	88.8
6482	7/30/2008	Marina	LA/LB harbor	33.76326	-118.25095	0.726	14	1.42	0.076	86.7
6328	8/4/2008	Marina	Dana point marina	33.46185	-117.70265	0.006	3.13	1.41	0.073	77.8
6327	8/4/2008	Marina	Dana point marina	33.46133	-117.70205	0.078	3.65	1.45	0.083	74.7
6325	8/4/2008	Marina	Dana point marina	33.46061	-117.70591	0.089	5.6	0.67	0.072	63.5
6320	8/4/2008	Marina	Dana point marina	33.45881	-117.6972	0.234	3.04	1.95	0.141	97.7
6294	8/4/2008	Marina	Oceanside	33.2078	-117.39731	0.044	7.86	0.61	0.067	55.5
6291	8/4/2008	Marina	Oceanside	33.20688	-117.39355	0.055	6.03	1.5	0.095	78
6288	8/4/2008	Marina	Oceanside	33.20486	-117.39073	0.118	4.51	1.45	0.124	78.5
6216	8/5/2008	Marina	Mission bay	32.78083	-117.24934	0.448	3.77	1	0.115	61.8
6213	8/5/2008	Marina	Mission bay	32.76826	-117.24723	0.246	5.27	1	0.069	52.2
6211	8/5/2008	Marina	Mission bay	32.76753	-117.23543	0.138	2.56	1.79	0.222	65.9
6204	8/5/2008	Marina	Mission bay	32.76254	-117.23619	0.138	6.49	1.27	0.145	66.7
6180	8/7/2008	Marina	San Diego bay	32.72844	-117.20187	0.293	2.49	0.58	0.056	53.9
6179	8/7/2008	Marina	San Diego bay	32.72811	-117.20838	0.293	2.49	0.49	0.052	51.9
6177	8/7/2008	Marina	San Diego bay	32.72714	-117.20218	0.16	3.84	0.92	0.116	74.5

6173	8/7/2008	Marina	San Diego bay	32.72486	-117.18351	0.293	4.66	0.27	0.044	
6171	8/6/2008	Marina	San Diego bay	32.72438	-117.22485	0.16	4.69	1.43	0.112	76.5
6165	8/6/2008	Marina	San Diego bay	32.72166	-117.22155	0.16	4.57	0.38	0.038	33.8
6159	8/6/2008	Marina	San Diego bay	32.71836	-117.23055	0.16	3.32	0.97	0.062	69.2
6161	8/6/2008	Marina	San Diego bay	32.71825	-117.22586	0.16	4.99	1.42	0.122	90.6
6157	8/6/2008	Marina	San Diego bay	32.71676	-117.2248	0.16	3.59	1.59	0.125	90.5
6153	8/6/2008	Marina	San Diego bay	32.71581	-117.23093	0.321	5.76	1.11	0.118	83.3
6151	8/6/2008	Marina	San Diego bay	32.71428	-117.2297	0.16	4.66	0.6	0.083	51.4
6148	8/6/2008	Marina	San Diego bay	32.71265	-117.23031	0.16	4.81	0.32	0.042	28.8
6145	8/6/2008	Marina	San Diego bay	32.71148	-117.23221	0.16	7.01	1.82	0.192	88.8
6027	8/22/2008	Marina	San Diego bay	32.6265	-117.13465	0.16	3.53	0.99	0.069	63.4
6025	8/22/2008	Marina	San Diego bay	32.62351	-117.13374	0.16	3.65	0.9	0.055	57.4
6546	9/4/2008	Port	Port Hueneme	34.15285	-119.20983	1.307	8.4	1.35	0.085	75.4
6493	7/31/2008	Port	LA/LB harbor	33.76918	-118.21711	1.307	15	1.12	0.072	79
6487	7/30/2008	Port	LA/LB harbor	33.76595	-118.27751	1.307	17	1.09	0.056	82
6467	7/29/2008	Port	LA/LB harbor	33.75306	-118.22368	1.307	15	1.28	0.068	82.7
6466	7/29/2008	Port	LA/LB harbor	33.7526	-118.21785	1.307	23	1.09	0.08	81.1
6460	7/29/2008	Port	LA/LB harbor	33.75013	-118.22486	1.307	16	0.32	0.019	34
6450	7/28/2008	Port	LA/LB harbor	33.74571	-118.21563	1.307	18	0.36	0.054	58.2
6449	7/29/2008	Port	LA/LB harbor	33.74495	-118.23841	1.307	10	0.72	0.041	74.4
6446	7/31/2008	Port	LA/LB harbor	33.74311	-118.20475	1.307	18	1.34	0.114	84.8

6443	7/31/2008	Port	LA/LB harbor	33.74165	-118.20536	1.307	20	0.33	0.039	69.6
6428	7/24/2008	Port	LA/LB harbor	33.73446	-118.23158	1.307	11	0.53	0.029	53.1
6424	7/22/2008	Port	LA/LB harbor	33.73266	-118.25316	1.307	17	0.92	0.055	68.9
6419	7/31/2008	Port	LA/LB harbor	33.73098	-118.19175	1.307	15	0.78	0.039	57.7
6413	7/17/2008	Port	LA/LB harbor	33.72913	-118.23406	1.307	12	0.89	0.058	72.3
6405	7/17/2008	Port	LA/LB harbor	33.7266	-118.23256	1.307	13	1.13	0.065	76
6402	7/22/2008	Port	LA/LB harbor	33.72411	-118.26233	1.307	26	1.34	0.108	90.4
6154	8/7/2008	Port	San Diego bay	32.71596	-117.17483	1.007	9.9	1.32	0.096	64.6
6155	8/7/2008	Port	San Diego bay	32.71594	-117.17591	0.458	10.85	0.47	0.07	51.6
6140	8/8/2008	Port	San Diego bay	32.70223	-117.16168	0.11	9.38	1.33	0.101	80.7
6133	8/8/2008	Port	San Diego bay	32.69618	-117.15301	0.11	12.52	1.7	0.157	84
6130	8/20/2008	Port	San Diego bay	32.69424	-117.23779	0.458	12.89	1.24	0.157	65.8
6129	8/21/2008	Port	San Diego bay	32.69163	-117.15291	0.458	12.58	0.13	0.02	18
6128	8/20/2008	Port	San Diego bay	32.69141	-117.23823	0.458	14.44	1.92	0.18	80.4
6127	8/18/2008	Port	San Diego bay	32.68995	-117.14003	0.11	12.55	2.33	0.157	91.3
6125	8/18/2008	Port	San Diego bay	32.68814	-117.13819	0.11	7.92	1.86	0.165	91.6
6120	8/18/2008	Port	San Diego bay	32.68704	-117.13414	0.03	8.22	2.24	0.164	73
6119	8/18/2008	Port	San Diego bay	32.68667	-117.13372	0.03	9.99	1.35	0.086	57.6
6116	8/18/2008	Port	San Diego bay	32.68555	-117.13396	0.03	10.75	2.03	0.126	80.6
6115	8/18/2008	Port	San Diego bay	32.6853	-117.13645	0.03	9.99	0.58	0.071	24.4
6113	8/18/2008	Port	San Diego bay	32.68473	-117.13543	0.03	10.51	1.21	0.094	73.8

609	4 8/19/2008	Port	San Diego bay	32.67536	-117.12882	0.11	11.43	1.19	0.084	81.6
608	7 8/19/2008	Port	San Diego bay	32.67163	-117.12618	0.11	12.55	1.24	0.077	83.4
608	5 8/19/2008	Port	San Diego bay	32.67104	-117.12384	0.11	12.22	1.14	0.099	84.5
657	2 8/18/2008	Port	San Diego bay	32.66887	-117.12869	0.458	8.01	1.43	0.136	88.1
607	5 8/19/2008	Port	San Diego bay	32.65987	-117.12241	0.11	6.58	1.16	0.103	48.6
607	2 8/19/2008	Port	San Diego bay	32.65887	-117.11933	0.11	5.57	0.71	0.066	61.7
657	0 8/19/2008	Port	San Diego bay	32.65812	-117.12241	0.11	11.55	1.39	0.115	82.9
666	1 8/21/2008	Port	San Diego bay	32.65759	-117.12275	0.11	11.97	0.89	0.073	72.2
666	0 8/21/2008	Port	San Diego bay	32.65607	-117.12254	0.11	13.25	0.74	0.064	64.5
605	4 8/21/2008	Port	San Diego bay	32.65143	-117.12288	0.458	11.76	0.89	0.077	72.2
665	9 8/22/2008	Port	San Diego bay	32.64782	-117.12026	0.11	6.49	0.79	0.048	46.2
604	2 8/21/2008	Port	San Diego bay	32.64753	-117.12127	0.03	12.22	0.54	0.041	65

Compound	Quantitation	Confir	mation
2,2',5,5'-			
tetrabromobiphenyl	79		
Bifenthrin	386	387	241
Fenpropathrin	141	206	
L-Cyhalothrin	241	205	243
Permethrin	207	209	171
Cyfluthrin	207	209	171
Cypermethrin	207	209	171
Esfenvalerate	211	213	294
Deltamethrin	297	295	299

Table SI-2. Quantitation and confirmation ions for pyrethroid analysis on GC-NCI/MS

Table SI-3. Pyrethroids concentration correlate with other parameters in 52 sediments (Spearman Rank Order correlation coefficient and p value)

	TOC	TN	%Fines	
Pyrethroid	0.585	0.295	0.202	
	(<i>p</i> =0.000)	(<i>p</i> =0.034)	(<i>p</i> =0.151)	
тос		0.598	0.436	
		(p=0.000)	(p=0.001)	
TN			0.419	
			(<i>p</i> =0.002)	

Ave sales in 2007- 2008	Runoff Loading	B'08 Found	Found / Sale (%)	Found / Runoff (%)	Runoff / Sale (%)
(kg)	(kg)	(kg)			
101242	134	36.3	0.036	27.2	0.13
(±10442)	(maximum 508)		(0.033-0.040)	(minimum 7.14)	(maximum 0.46)

Table SI-4. Comparison of pyrethroid found and usage

* Estimated from storm runoff pyrethroid concentrations (average 97.5 μ g/kg, standard deviation 126 μ g/kg) and average wet season storm runoff volume (1.37± 9.06 × 10⁹ m³) of two water years (October-September), i.e., 2007-2008, 2008-2009.



Figure SI-1. Map of study region and sampling sites.

Figure SI-2. Sum of pyrethroid concentration in each region.

