# High levels of persistent organic pollutants measured in blubber of island-associated false killer whales (*Pseudorca crassidens*) around the main Hawaiian Islands

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Persistent organic pollutants (POPs) have been measured in tissues of marine mammals since the mid 1960s (Holden and Marsden. 1967; Wolman and Wilson, 1970). These compounds include several pesticides (e.g., DDTs, chlordanes) and industrial chemicals (e.g., PCBs) that are ubiquitous, highly lipophilic and not readily degraded or metabolized. As a result, they can biomagnify to high levels in lipid-rich tissues of top-level marine predators. POPs enter marine waters via direct inputs (e.g., sewage outfalls, industrial and agricultural runoff) as well as from indirect sources (e.g., ocean currents) (Friedlander et al., 2005). Exposure to POPs in marine mammals has been linked to a number of biological effects including reproductive impairment (DeLong et al., 1973; Subramanian et al., 1987), reduced reproductive success (Wells et al., 2005), immune suppression (De Swart et al., 1994; Hammond et al., 2005; Ross et al., 1995) and endocrine disruption (reviewed in O'Hara and O'Shea (2001)). Although many POPs, such as PCBs and DDTs, have been banned for production or use in the US for more than thirty years, some of these compounds are still used in other regions of the world (Fielder, 2008; van den Berk, 2009) and continue to be measured in the tissues of marine mammals throughout coastal regions of the US.

Another class of POPs gaining the attention of environmental scientists and managers are the polybrominated diphenyl ethers (PBDEs). Three different PBDE technical mixtures (i.e., penta-BDE, octa-BDE, deca-BDE) have been manufactured and added as flame retardants to plastics, textiles, clothing, electronic circuit boards and other materials in industrial and developing nations (de Wit, 2002). Deca-BDE is the primary commercial product produced and used in the U.S. as a result of the sole manufacturer phasing out the production of penta-BDE and octa-BDE (U.S. EPA, 2006). Similar to PCBs, these compounds are lipophilic, persistent, and tend to bioaccumulate in marine mammal tissues (de Wit et al., 2004; Ikonomou et al., 2002a, 2002b). Some of the highest levels of PBDEs have been measured in tissues of wildlife and humans from North America due to high volume PBDE use in this region of the world (Hites, 2004; Ikonomou et al., 2002b; LeBeuf et al., 2004). Because these compounds can travel over long distances via atmospheric transport, they have been measured in marine organisms throughout the world, including Antarctica and the Arctic (Corsolini et al., 2006; de Wit et al., 2004). Exposure to PBDEs has been associated with a variety of biological effects (e.g., thyroid disruption, neurobehavioral effects) in laboratory animals (de Wit, 2002) but currently no threshold levels for PBDEs have been established for toxicological effects in marine mammals.

The main Hawaiian Islands include eight volcanic islands (i.e., Hawai'i, Kaho'olawe, Kauai, Lana'i, Mau'i, Moloka'i, Ni'ihua, O'ahu) that are located in the middle of the Pacific Ocean, approximately 1500 miles southwest of the contiguous US. Tourism, defense and agriculture (e.g., production of raw sugar, fresh

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pineapple) are the primary contributors to the economy of this region (State of Hawaii, Dept. of Business, Economic Development and Tourism, 2008). In addition, coastal development of the main Hawaiian Islands is ongoing and includes conversion of agricultural lands to residences and resorts, as well as expansion of harbor facilities to accommodate large cargo and cruise ships (Friedlander et al., 2005). Activities related to these industries and development processes can be potential sources of POPs to this region. For example, in the 1970's, elevated levels of chlorinated insecticides used to control agricultural pests and termites were reported in water, sediment and aquatic organisms from the main Hawaiian Islands (Bevenue et al., 1972; Tanita et al., 1976).

Around the main Hawaiian Islands, the highest trophic level cetacean regularly encountered is the false killer whale (*Pseudorca crassidens*). Based on observations of predation, individuals from this population appear to feed primarily on large game fish such as mahimahi (*Coryphaena hippurus*), yellowfin tuna (*Thunnus albacares*) and swordfish (*Xiphias gladius*), some of which can be long-lived (Baird et al., 2008). Population estimates for cetaceans within the Hawaiian Exclusive Economic Zone (EEZ) indicate that false killer whales may have the smallest population size of any odon-tocete within the Hawaiian EEZ (Barlow, 2006). In addition, within this region of Hawai'i there is evidence of population structure for false killer whales, with genetically differentiated insular and off-shore populations (Chivers et al., 2007).

Recently, the population of false killer whales around the main Hawaiian Islands has been estimated at 123 individuals (CV = 0.72) based on a photographic mark-recapture analysis (Baird et al., 2005). There is evidence that this population may have declined substantially over the last 20 years (Reeves et al., 2009). A number of potential causes have been identified, including mortality in the Hawai'i-based long-line fishery (Baird and Gorgone, 2005; Forney and Kobayashi, 2007), reduction in their prey base, and potential health or reproductive effects due to exposure to high levels of POPs (Reeves et al., 2009). Although the details of life history of false killer whales are poorly known, females appear to reach sexual maturity between 8 and 14 years (Purves and Pilleri, 1978), have long calving intervals (estimated at 6.9 years), and may reach 62.5 years (Kasuya, 1986). Males are thought to mature as much as 10 years later and live to 57.5 years (Kasuya, 1986). As a long-lived upper-trophic level predator, false killer whales are likely to accumulate high levels of POPs. Few individuals have been analyzed for POPs, but high levels have been documented in animals that stranded around British Columbia (Baird et al., 1989; Jarman et al., 1996). In the present study, we report concentrations of PBDEs and other POPs measured in biopsy samples collected from nine individuals from the insular population to determine the baseline levels of these contaminants and assess whether their exposure levels to PCBs may be a risk factor for this population.

Field operations were undertaken as part of ongoing studies of odontocetes around the main Hawaiian Islands (see Baird et al.,

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2008). In July 2008, biopsy blubber samples were collected from nine individual false killer whales from the insular population using a 45 kg pull Barnett RX-150 crossbow and Larsen biopsy tips. measuring 25 mm long and 8 mm wide. A high-density foam collar on the biopsy dart prevented penetration greater than 18 mm. After collection, the biopsy samples were stored in a cooler with ice packs while in the field and transferred to a -20 °C freezer for short-term storage before being stored in a -80 °C freezer. Biopsied individuals were photo-identified, and photographs compared to each other to eliminate duplicate samples, and to the catalog of Baird et al. (2008) to assess population identity and sighting history. Age class (adult/subadult) was assessed in the field based on relative body size and in some cases confirmed based on sighting history. The sex determination of each whale was conducted using zinc finger gene amplification (Chivers et al., 2007). Based on photographs all individuals appeared to be "healthy" (i.e., not emaciated).

Samples were extracted and analyzed for POPs using the gas chromatography/mass spectrometry method of Sloan et al. (2005). Blubber (0.1-0.3 g) was extracted with methylene chloride using an accelerated solvent extractor after the addition of a surrogate standard (PCB 103; 1 ng/ $\mu$ L). This procedure was followed by a clean-up step of the extract on a single stacked, gravity flow silica gel/alumina column to remove any highly polar compounds present in the sample. Using high-performance size exclusion liquid chromatography, the POPs were separated from the bulk lipid and other biogenic material present in each sample, and the cleaned extract was analyzed for POPs using a low-resolution quadrupole GC/MS system equipped with a 60 m DB-5 GC capillary column and a electron impact mass spectrometer in selected ion monitoring mode. The instrument was calibrated using sets of up to ten multi-level calibration standards of known concentrations. Percent lipid and lipid class profiles were determined in biopsy blubber samples using thin-layer chromatography with flame ionization detection (Ylitalo et al., 2005). In this method, each lipid extract sample was spotted on a Type SIII Chromarod and developed in a chromatography tank containing 60:10:0.02 hexane:diethyl ether: formic acid (v/v/v). The lipid classes were separated based on polarity and measured using flame ionization detection. Percent lipid values were calculated by summing the concentrations of five lipid classes (i.e., sterol esters/wax esters, triglycerides, free fatty acids, cholesterol, phospholipids) for each sample, using the mean of two measurements.

All blubber contaminant concentrations are reported in ng/g, lipid weight. Sum PCBs ( $\sum$ PCBs) includes the sum of congeners 17, 18, 28, 31, 33, 44, 49, 52, 66, 70, 74, 82, 87, 95, 99, 101/90, 105, 110, 118, 128, 138/163/164, 149, 151, 153/132, 156, 158, 170, 171, 177, 180, 183, 187/159/182, 191, 194, 195, 199, 205, 206, 208 and 209. Sum DDTs ( $\sum$ DDTs) is the sum of *o*,*p*'-DDD, *p*,*p*'-DDD, *o*,*p*'-DDE, *p*,*p*'-DDT and *p*,*p*'-DDT. Sum chlordanes ( $\sum$ CHLDs) is the sum of heptachlor, heptachlor epoxide, oxychlordane, *gamma*-chlordane, nona-III-chlordane, *alpha*-chlordane, *trans*-nona-chlor and *cis*-nonachlor. Sum PBDEs ( $\sum$ PBDEs) is the sum of congeners 28, 47, 49, 66, 85, 99, 100, 153, 154 and 183. Additional POPs analyzed in the current study include hexachlorobenzene (HCB),  $\beta$ -hexachlorocyclohexane ( $\beta$ -HCH), aldrin, dieldrin, mirex and endosulfan I.

As part of a performance-based quality assurance program (Sloan et al., 2006), a method blank and a National Institute of Standards and Technology (NIST) Standard Reference Material (SRM® 1945) were analyzed with the false killer whale blubber samples. Concentrations of individual analytes measured in SRM® 1945 were in excellent agreement with the reference values published by NIST. Other quality control samples met established laboratory criteria. POP concentrations were  $log_{10}(x + 1)$  transformed and percent lipid values were arcsine square root transformed to increase the homogeneity of variance. Analysis of variance (ANOVA) and the Tukey-Kramer honestly significant difference test (HSD) were used to compare mean concentrations of POPs among three age/sex classes (subadult whales (both males and females), adult females, adult males) (Zar, 1999). The level of significance used for all statistical tests was  $a \leq 0.05$ . All statistical analyses were completed using JMP Statistical Software (SAS Institute, Inc., Cary, NC).

The most abundant POPs measured in biopsy blubber of false killer whales from the main Hawaiian Islands were DDTs and PCBs, with concentrations ranging from 1000 to 83000 ng/g, lipid (Table 1). PBDEs, chlordanes,  $\beta$ -HCH, dieldrin, HCB and mirex were also measured in these whales but at much lower concentrations than DDTs and PCBs (Table 1, Fig. 1). Endosulfan I and aldrin, on the other hand, were below the LOQ for all animals analyzed in the current study. Recent studies have reported measuring relatively low levels of PCBs, DDTs, PBDEs and other contaminants in aquatic organisms from the main Hawaiian Island region (Brasher and Wolff, 2004; Kimbrough et al., 2009; Miao et al., 2001; Orazio et al., 2003; Xu et al., 2009; Yang et al., 2008) but no data have been previously available for false killer whales or their presumed prev.

#### Table 1

Concentrations of  $\Sigma$ CHLDs,  $\Sigma$ PDTs,  $\Sigma$ PCBs and  $\Sigma$ PBDEs measured in biopsy blubber samples of false killer whales from the main Hawaiian Islands sampled in July 2008.

| Sample ID                     | Sex/age class         | Collection date | Percent lipid | Lipid weight (            | ng/g)                    |                    |                           |
|-------------------------------|-----------------------|-----------------|---------------|---------------------------|--------------------------|--------------------|---------------------------|
|                               |                       |                 |               | $\sum$ CHLDs <sup>a</sup> | $\sum$ DDTs <sup>a</sup> | ∑PCBs <sup>a</sup> | $\sum$ PBDEs <sup>a</sup> |
| RWB2008Jul26.02               | Subadult – female (S) | 7/26/2008       | 16            | 2900                      | 16,000                   | 14,000             | 2400                      |
| RWB2008Jul26.03 <sup>b</sup>  | Subadult – male       | 7/26/2008       | 41            | 3200                      | 23,000                   | 24,000             | 2900                      |
| -                             |                       | Mean ± SD       | 29 ± 18       | $3100 \pm 210$            | 20,000 ± 4900            | 19,000 ± 7100      | 2700 ± 350                |
| RWB2008Jul16.01               | Adult male (M)        | 7/16/2008       | 18            | 4100                      | 83,000                   | 33,000             | 780                       |
| RWB2008Jul16.04               | Adult male            | 7/16/2008       | 16            | 4900                      | 43,000                   | 33,000             | 1600                      |
|                               |                       | Mean ± SD       | 17 ± 1.4      | $4500 \pm 570$            | 63,000 ± 28,000          | 33,000 ± 0         | 1200 ± 580                |
| RWB2008Jul16.03               | Adult female (F)      | 7/16/2008       | 23            | 190                       | 1200                     | 1000               | 26                        |
| RWB2008Jul16.05               | Adult female          | 7/16/2008       | 36            | 1300                      | 8300                     | 11000              | 1700                      |
| RWB2008Jul16.06               | Adult female          | 7/16/2008       | 35            | 430                       | 2500                     | 2200               | 120                       |
| RWB2008Jul16.07 <sup>b</sup>  | Adult female          | 7/16/2008       | 12            | 140                       | 1200                     | 1100               | <loq<sup>c</loq<sup>      |
| RWB2008Jul26.06               | Adult female          | 7/26/2008       | 16            | 310                       | 1800                     | 2100               | 260                       |
|                               |                       | Mean ± SD       | $24 \pm 11$   | $470 \pm 480$             | 3000 ± 3000              | $3500 \pm 4200$    | $420 \pm 720$             |
| p Value <sup>d</sup>          |                       |                 | 0.6393        | 0.0074                    | 0.0029                   | 0.0101             | 0.1832                    |
| Tukey-Kramer HSD <sup>e</sup> |                       |                 | -             | M,F; S,F                  | M,F; S,F                 | M,F; S,F           | -                         |

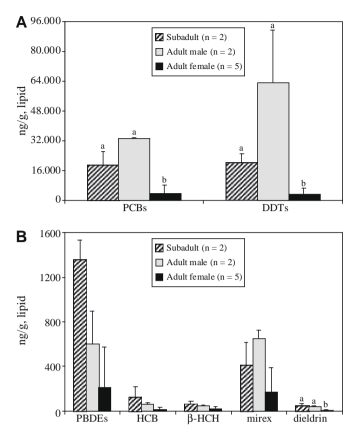
<sup>a</sup> Individual compounds summed are reported above.

<sup>b</sup> Mother/offspring pair.

<sup>c</sup> <LOQ for the sum indicates concentrations of all compounds included in the sum were below their individual limits of quantitation. For each <LOQ, a value of zero was used to calculate the mean and standard deviation of the mean.

<sup>d</sup> Significant differences (ANOVA, p < 0.05) in POPs and lipid concentrations based on age class are shown in bold.

<sup>2</sup> Unlike letters indicate significant differences using Tukey–Kramer honestly significant difference (HSD) test (p < 0.05).



**Fig. 1.** Mean (±SD) concentrations of  $\sum$ PCBs and  $\sum$ DDTs (A) and  $\sum$ PBDEs HCB,  $\beta$ -HCH, mirex and dieldrin (B) measured in biopsy blubber samples of adult and subadult false killer whales from the main Hawaiian Islands. Bars with unlike letters differ significantly; Tukey–Kramer HSD test, *p* < 0.05.

Consumption of contaminated prey is the primary route of exposure for marine mammals (Aguilar et al., 1999).

Age class and sex appeared to influence the concentrations of POPs measured in the false killer whales. Mean levels of  $\sum$ CHLDs,  $\sum$ DDTs and  $\sum$ PCBs were significantly different among the three age/sex classes of whales, with adult females having lower values than those measured in adult males and subadults (Table 1). Similarly, the mean PBDE concentration in adult females was lower than those in subadults and adult males, but these differences were not significant at a = 0.05 level (Table 1). Examination of POPs measured in blubber samples of a mother-offspring pair (RWB2008Jul16.07 and RWB2008Jul26.03) also showed that the levels of  $\sum$ CHLDs,  $\sum$ DDTs,  $\sum$ PCBs (Table 1) and mirex (data not shown) were at least an order of magnitude higher in the subadult male offspring than those measured in his mother. A number of contaminant studies on odontocetes have also reported lower POP levels in adult females compared to those measured in blubber of adult males and juveniles (subadult) (Hansen et al., 2004; Tilbury et al., 1999; Wells et al., 2005; Westgate et al., 1997; Ylitalo et al., 2001) due to the transfer of lipids and the POPs associated with these lipids from mother to calf during gestation and lactation (Aguilar and Borrell, 1994; Gardner et al., 2007). In contrast, males continue to accumulate these compounds throughout their lives.

One finding of interest in the current study was that subadult whales had the highest mean level of  $\sum$ PBDEs measured in these animals; however, these differences were not significant (*p* = 0.1832) among the age classes of whales (Table 1). The subadults also had elevated mean HCB,  $\beta$ -HCH and dieldrin concentrations (Fig. 1) compared to those measured in adult males and females but only dieldrin levels were significantly different

(p = 0.0496) among the age classes. In recent killer whale (*Orcinus orca*) studies, higher levels of  $\sum$ PBDE, HCB and  $\sum$ HCHs have been reported in blubber of juvenile fish-eating individuals ("Southern Residents") compared to those determined in adult males and females from the same population (Krahn et al., 2007b, 2009). The elevated levels of PBDEs, HCB,  $\beta$ -HCH and dieldrin measured in the blubber of the subadult false killer whales may be due to differences in prey items or feeding rates, as well as variations in metabolism and excretion of these lipophilic compounds compared to adults (Aguilar et al., 1999). These findings of elevated contaminant levels in subadult whales are a concern as these animals are still developing biologically and may be at higher risk to deleterious effects associated with exposure to these compounds than adults in the same population.

The percent lipid measured in the biopsy blubber samples of the false killer whales ranged from 12–41% and contained primarily triglycerides (>84%) and wax esters (<16%). These percent lipid values are comparable to those reported in biopsy blubber samples of Eastern North Pacific killer whales that were analyzed by the same quantitation method (Herman et al., 2005; Krahn et al., 2007a,b, 2009; Ylitalo et al., 2001). Similar to our findings, Litchfield et al. (1975) reported that blubber of false killer whales contained both wax esters (4%) and triglycerides (96%).

The mean concentrations of  $\sum$  PCBs,  $\sum$  DDTs and  $\sum$  CHLDs measured in biopsy blubber of adult male false killer whales in the present study are much lower than those reported previously in blubber of two adult male false killer whales (Table 2) that stranded in British Columbia in the late 1980s (Baird et al., 1989; Jarman et al., 1996). These findings may be due to differences in analytical methodologies or variations in contaminant levels in presumably "healthy" wild-ranging whales (current study) vs. stranded animals that may have been in poor health. In addition, it is also probable that differences in contaminant levels in feeding ranges (waters of main Hawaiian Islands vs. west coast of North America) and sample collection years (2008 vs. 1987/1989) as well as variations in the ages of animals sampled also contributed to these differences in POPs levels observed for the false killer whales (Aguilar et al., 1999). In contrast to  $\sum PCBs$ ,  $\sum DDTs$  and  $\sum CHLDs$ , the levels of mirex - a POP used as a fire retardant and insecticide in the US until 1978 (ASTDR, 1995) - measured in whales in the current study were more than two times higher than those found in the false killer whales that stranded in British Columbia (Baird et al., 1989; Jarman et al., 1996). These differences in mirex levels between the two whale groups may be due to the more extensive use of this pesticide in Hawai'i (used to control mealy bugs in pineapple fields) compared to the west coast of North America (ASTDR, 1995; UNEP/FAO, 2005).

We compared the POPs levels measured in the current study with those reported recently in other fish-eating marine mammal species from the west coast of North America (Blasius and Goodmanlowe, 2008; Ikonomou et al., 2002b; Krahn et al., 2007a,b, 2009; Meng et al., 2009) because no contemporary POP data have been published on cetaceans from the Hawaiian Island region (O'Shea et al., 1980). Mean  $\sum$ CHLDs,  $\sum$ PCBs and  $\sum$ DDTs concentrations measured in the Hawaiian false killer whales were lower than the values reported for California sea lions (Zalophus californianus) from southern California (Blasius and Goodmanlowe, 2008), offshore killer whales sampled in Alaska (Krahn et al., 2007a) and "Southern Resident" killer whales (Krahn et al., 2007b, 2009) but were higher than those measured in Alaskan resident killer whales except for  $\sum$ CHLDs (Krahn et al., 2007a). The mean level of  $\sum$ PBDEs (Table 2) measured in the Hawaiian false killer whales was higher or comparable to those reported in Alaskan resident killer whales (Krahn et al., 2007a) and harbor porpoises (Phocoena phocoena) that stranded in urban harbors of British Columbia (Ikonomou et al., 2002b) but was lower than

|   | Collection region   | Collection region Collection year(s) $n$ Percent lipid Lipid weight $(ng/g)$ | и  | Percent lipid | Lipid weight (n       | g/g)              |   |                      |                 |                                |
|---|---------------------|--|----|---------------|-----------------------|-------------------|---|----------------------|-----------------|--------------------------------|
| Species                                   |                     |  |    |               | Mirex                 | ∑chlds            | <b>DDTs</b>                             | $\sum PCBs$          | ∑PBDEs          | Reference                      |
| False killer whale                        | Hawai'i             | 2008   | 2  | 17 ± 1.4      | $610 \pm 510$         | $4500 \pm 570$    | 63,000 ± 28,000                         | 33,000 ± 0           | $1200 \pm 580$  | Current study                  |
| False killer whale                        | British Columbia    | 1987, 1989   | 7  | 91 ± 2.8      | 260 ± 99              | $15,000 \pm 710$  | $1,000,000 \pm 1400,000$                | $45,000 \pm 7500$    | Not analyzed    | Jarman et al. (1996)           |
| Harbor porpoise                           | British Columbia    | 1991/1993  | 5  | $80 \pm 10$   | Not analyzed          | Not analyzed      | Not analyzed                            | Not analyzed         | $810 \pm 740$   | Ikonomou et al. (2002b)        |
| California sea lion                       | Southern California | 1994/2006  | 5  | 38 ± 41       | Not detected          | $4900 \pm 7700$   | 2300,000 ± 2900,000                     | 290,000-440,000      | 55,000 ± 79,000 | Blasius and Goodmanlowe (2008) |
|   |                     |  |    |               |                       |                   |   |                      |                 | and Meng et al. (2009)         |
| Offshore killer whale                     | Alaska              | 2003/2004  | 4  | 18 ± 3.7      | $760 \pm 180^{a}$     | $16,000 \pm 2300$ | $16,000 \pm 2300$ $420,000 \pm 100,000$ | $110,000 \pm 22,000$ | $3300 \pm 940$  | Krahn et al. (2007a)           |
| Resident killer whale                     | Alaska              | 2003/2004  | 40 | 25 ± 11       | $170 \pm 64^{a,b}$    | $6200 \pm 2400$   | $21,000 \pm 12,000$                     | $13,000 \pm 5900$    | 76 ± 70         | Krahn et al. (2007a)           |
| Resident killer whale Puget Sound, WA     | Puget Sound, WA     | 2004/2007  | 10 | 22 ± 6.1      | 190 ± 72 <sup>a</sup> | 8600 ± 3300       | 82,000 ± 38,000                         | 56,000 ± 46,000      | $4400 \pm 1800$ | Krahn et al. (2007b, 2009)     |
| <sup>a</sup> Data from I Bolton Pers Comm | hers Comm           |  |    |               |                       |                   |   |                      |                 |                                |

n = 37.

Concentrations of persistent organic pollutants measured in blubber of adult male marine mammals from the west coast of North America.

Table

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the mean values measured in blubber of California sea lions that stranded in southern California (Meng et al., 2009), as well as Southern Resident (Krahn et al., 2007b, 2009) and offshore killer whales (Krahn et al., 2007a). These findings were expected as offshores and "Southern Residents", as well as California sea lions, appear to spend a portion of their time feeding on fish from highly urbanized areas (e.g., Puget Sound, Washington, central and southern California coasts) based on observational field data and contaminant levels and/or ratios, whereas the false killer whales, harbor porpoises and Alaskan resident killer whales primarily consume prey from less contaminated regions of the eastern North Pacific (e.g., main Hawaiian Islands, Vancouver, British Columbia, Eastern Aleutian Islands). A possible source of PBDEs to the Hawaiian coastal region is effluent from wastewater treatment plants as a number of plants discharge to the coastal ocean in Hawai'i (Friedlander et al., 2005) and appreciable levels of these compounds have been measured previously in wastewater effluents (de Boer et al., 2003; North, 2004). However, PBDEs may be entering this marine ecosystem via other sources that have not yet been identified.

Accumulation of high tissue levels of POPs has been associated with biological and physiological effects in marine mammals (O'Hara and O'Shea, 2001). For example, Kannan et al. (2000) recommended a safe upper PCB threshold concentration of 17,000 ng/g, lipid for PCBs in blubber based on a number of studies that measured various toxicological endpoints (e.g., thyroid hormone concentrations) and PCB concentrations. Three out of nine animals sampled in the current study had  $\sum$ PCBs that exceeded this threshold value. Our findings indicate that some of these animals are exposed to PCB levels that may affect their health. In addition to PCBs, these animals are also exposed to other classes of toxic POPs that may increase their risk to adverse effects.

The current study is the first to report blubber concentrations of POPs, including PBDEs, in free-ranging false killer whales, and the first for any free-ranging cetaceans from the Hawaiian Islands. Wide ranges of POP concentrations were measured in these animals, with DDTs and PCBs being the most abundant. Similar to previous cetacean studies, age class and sex influenced the levels of POPs measured in the whales. Interestingly, subadult false killer whales had higher levels of some classes of POPs (e.g.,  $\sum$ PBDEs, dieldrin, HCB) compared to the other sampled animals. Although the POP concentrations measured in the false killer whales in the current study were generally equal to or lower than those reported for false killer whales that stranded in British Columbia or fish-eating marine mammals from the west coast of North America, some of the animals in the current study were exposed to PCB levels that could potentially affect their health. Due to the small size of this whale population and their life history strategies (e.g., long-lived, time to maturation), continued monitoring of POPs is essential in assessing the health and viability of these animals.

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# Riverine input of particulate material and inorganic nutrients to a coastal reef ecosystem at the Caribbean coast of Costa Rica

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Reasons for the alarming global coral reef destruction can often be found on land (ISRS, 2004). Agricultural activities accompanied by land clearing, fertilization, use of pesticides, and urbanization or tourism expansion along with enhanced sewage and waste production are of concern (ISRS, 2004). Rivers and groundwater carry high loads of sediment, nutrients and other pollutants to the sea, where they can have serious impacts on nearshore ecosystems such as coral reefs (Cortés and Risk, 1985; Guzmán and Jiménez, 1992; Rogers, 1990; Fabricius, 2005).

While nutrients enhance coral growth in lower amounts, they inhibit it when highly concentrated (Tomascik and Sander, 1985; Koop et al., 2001), and accelerate the progress and severity of coral disease (Bruno et al., 2003; Voss and Richardson, 2006). Nutrients fuel algal growth and, combined with reduced herbivory, can be responsible for shifts from coral- to algal-dominated reefs (Díaz-Pulido and McCook, 2003; Hughes et al., 2003). Also bioeroders such as algae, sponges, worms or bivalves profit from nutrient and organic matter increase (Risk and MacGeachy, 1978). Bored sediments and corals are less resistant to storms and waves, resulting in reef erosion (Hallock, 1988; Chazottes et al., 2002).

Suspended matter in the water column decreases transparency and light availability. While organic material may initially be used as an additional food source by corals, this benefit is outweighed in turbid water, where photosynthesis and calcification are reduced (Rogers, 1983; Anthony and Fabricius, 2000). Smothering by particulate material forces the coral to clean its surface using energy needed for growth or reproduction (Tomascik and Sander, 1987; Edmunds and Davies, 1989). Terrestrial runoff can become a serious threat for reef communities and even small rivers have been shown to influence reefs within a few kilometers distance to their mouths (West and van Woesik, 2001; Fabricius, 2005).

The aim of this study was to evaluate the present influence of a heavily anthropogenic impacted river on the distribution of particulate material and dissolved inorganic nutrients in the waters of a nearby coral reef area in Costa Rica.

The Caribbean coast of Costa Rica is characterized by humid, hot climate with year-round rains of about 6000 mm (Cortés and Jiménez, 2003). Precipitation between December and February and between June and August is higher compared to the rest of the year. The largest, best-developed and most diverse reef in the area is found in the Cahuita National Park (Cortés and León, 2002;

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