Maternal transfer of chlorinated contaminants in the leatherback turtles, *Dermochelys coriacea*, nesting in French Guiana

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species; these adverse effects have been shown for eggs treated experimentally with low dose of pesticides (between 0.25 and 14 ng of trans-Nonachlor, chlordane or pp′-DDE per egg) or for eggs sampled in contaminated areas with concentrations up to 1000 and 3500 ng g⁻¹ of pesticides and PCBs respectively (Woodward et al., 1993; Bishop et al., 1998; Willingham, 2001). Current levels have been described in marine turtles (Supporting Material – SM, Table 1), however toxicokinetic and potentially harmful effects of OCs remain poorly known. Investigations into these types of contaminants could help to assess the potential risk to turtle's health. To assess a baseline of OC concentrations in free-ranging leatherback, non-lethal and adequate sampling is required in the form of blood and egg samples that can be easily collected. Blood is relatively non-invasive and facilitates the repeated collection of larger numbers of samples which improves both monitoring of OC levels and assessment of toxicological effects. As egg formation occurs when females forage in pelagic waters about 4–5 months before arriving at nesting sites (James et al., 2005a) and as it is thought that leatherback females from French Guiana are not eating or eat only a small part of their lipids during migrations facilitating exposure to various environmental toxicants (Miller, 1997) making this species particularly interesting to study in terms of maternal transfer of contaminants. In this study, we assessed this maternal transfer of OCs, described the distribution, patterns and relationships of PCBs and OCs in blood and egg sampled in 38 nesting leatherbacks. We also investigated the temporal variations of this transfer along and between nesting seasons.

2. Materials and methods

2.1. Study site and sample collection

The study was conducted at Yalimapo beach situated within the Amana Natural Reserve in French Guiana. Leatherback females are marked with an internal permanent marker (Passive Integrated Transponder [PIT] tag) located in the shoulder muscle. These coded microchips are used to identify leatherback turtles and to allow temporal monitoring for females during and between nesting seasons. From this information, we were able to know how many nests had occurred between two nesting observations and since how many years this female has nested previously (mainly of 2 or 3 years). From 16th March to 14th May 2006, we patrolled the beach each night around high tide. Nesting females encountered during patrols were scanned for PIT tags and blood and eggs were sampled during egg-laying. We also sampled adipose tissues in the shoulder but biopsies revealed that there were no more adipose reserves here at the beginning of the nesting season. One yolled-egg was collected arbitrarily around the 20th egg laid and whole blood was sampled from the rear flipper and samples were then frozen at −20 °C until analysis. For this study 38 females were sampled including 17 females for two different clutches and three females for three different clutches either consecutive or not.

2.2. Sample preparation

Egg content (yolk and albumen) were homogenized with an Ultra-Turrax. A 4 g sample was lyophilized for 16 h and dry matter was determined gravimetrically. To 500 mg sample of lyophilized egg were added 500 mg of anhydrous sodium sulfate and 50 µL of PCB 112 (100 pg µL⁻¹ in acetone) used as surrogate marker. These spiked samples with surrogate were extracted with a mixture of hexane, dichloromethane and methanol (5:2:1, v:v:v) at 80 °C under a pressure of 1500 psi using an accelerated solvent extractor (Dionex ASE 2000). The solvent with extracted fat were collected in pre-weighted vials and evaporated at 40 °C under a gentle nitrogen flow (Turbovap) and fat content was determined gravimetrically. The residues containing both lipids and OCs of interest were dissolved into 3 mL of hexane and collected into a test tube. The mixture was homogenized by vortexing for 1 min.

Blood samples were prepared by a method modified from Pauwels et al. (1999), Janak et al. (1999) and Sundboerg et al. (2006). Four millilitres of blood were first deproteinised by adding 100 µL of triethylamine and 5 mL of formic acid; 50 µL of PCB 112 (100 pg µL⁻¹ in acetone) were used as surrogate marker. The mixture was stabilized for 30 min in an ultrasound bath. Five millilitres of hexane were added and samples were shaken manually for 2 min and centrifuged for 10 min at 3000 rpm. The organic phase was transferred into a new tube and evaporated under a gentle stream of nitrogen in order to obtain a final volume of 1 mL.

2.3. Sample clean-up and analysis

All prepared samples (egg and blood) were purified by acid and Florisil clean-ups. Sample clean-up were performed as detailed in Debier et al. (2003). The egg and blood purified extracts were analysed by high resolution gas chromatography using a Thermo Quest Trace 2000 gas chromatograph equipped with a NiECD detector and an autosampler for liquids Thermo Quest AS 2000. Analysis of samples is also describes in Debier et al. (2003). Twenty-three PCB pure components (IUPAC 28, 44, 52, 66, 70, 87, 95, 101, 105, 118, 128, 138, 149, 153, 156, 170, 180, 183, 187, 194, 195, 206, 209) and 5 OCs of interest (pp′-DDE, pp′-DDE, p,p′-DDE, α-HCH and γ-HCH) were measured.

2.4. Quality assurance

The quality control was performed by analyses of procedural blanks, by injection of standard and n-hexane blanks. Standard reference materials SRM 1946 (PCBs and OCs in Lake Superior fish tissue) and BCR RM 349 (cod liver), were used to test the whole procedure of the analytical method accuracy (recovery efficiency for certified concentrations between 71% and 128% according to the compounds). Procedural blanks and laboratory made quality control were run with each set of samples to control the extraction and clean-up procedures. Blanks were also used to control lyophilization and ASE steps for egg samples. The laboratory made quality controls (QC) were bovine blood and milk cream, for blood and egg series respectively, spiked with a PCBs and OCs mix at 2.5 and 15 ng g⁻¹ for bovine blood and milk cream respectively. For each PCB congener and OCP, recovery efficiency was calculated on the basis of the concentration of the surrogate marker PCB 112. All results were corrected to obtain 100% recovery. However, the results of the PCBs and OCs analyses were accepted only if the recoveries were between 70% and 130%. In egg samples, concentrations were calculated on a wet-mass basis and on a lipid-mass basis in order to minimize inter-individual variation and to allow comparison of results with other studies. For blood samples, concentrations are only expressed on a wet-mass basis as the lipid fraction in blood is known to be very low (<1%, Keller et al., 2004c). Total PCBs
concentrations (ΣPCBs) were calculated as the sum of all individual quantified congeners, total DDTs (ΣDDTs) as the sum of \( \text{pp}'-\text{DDT} \), \( \text{pp}''-\text{DDE} \) and \( \text{pp}''-\text{DDD} \) concentrations and total HCHs (ΣHCHs) as the sum of \( \alpha \)-HCH and \( \gamma \)-HCH concentrations. For each PCB congeners and OCPs, the limits of quantification (LOQ) were 0.09 ng g\(^{-1}\) and 0.08 ng g\(^{-1}\) for egg and blood samples respectively. The efficiency recovery for PCB congeners and of OCPs was always above 75% for egg and blood samples.

2.5. Statistical analysis

For statistical analysis, OC concentrations that were below the limit of quantification (LOQ) were estimated at half the LOQ. Firstly, general linear models (GLM) with repeated measurements and logit link function were also performed to assess the effect of RI on the proportion of each class of contaminant analyzed. Finally, GLMM were performed to test the relationship between concentrations in eggs when regressed against corresponding concentrations in blood. Simple regression models were then applied to look at potential correlations between concentrations of OCs in eggs and in blood. The normality of the dependent variables was confirmed prior to the analyses. Computations were performed with the SAS package.

3. Results

3.1. Concentrations and patterns

Due to interference during analysis, PCBs 28, 44, 52, 170 and 195 could not have been properly detected and quantified and they were therefore removed from samples profiles. For each congener, the mean percentage contribution to ΣPCBs, called congener pattern, was calculated. Congener pattern is dominated in blood by the more chlorinated congeners whereas congener patterns in eggs were dominated by the less chlorinated congeners (Fig. 1A). When grouping congeners by degree of chlorination (4–5 Cl, 6 Cl and 7–9 Cl, each class having approximately the same number of congeners), the three groups were equally represented in blood (Fig. 1B) but the group's patterns present a tendency for the lower chlorinated group in eggs (Fig. 1B). In blood, PCBs were the predominant OCs found. PCBs 153, which is generally the most persistent and abundant congeners found in biological tissues, is the most prevalent PCB congeners, total DDTs (ΣDDTs) as the sum of \( \text{pp}'-\text{DDT} \), \( \text{pp}''-\text{DDE} \) and \( \text{pp}''-\text{DDD} \) concentrations and total HCHs (ΣHCHs) as the sum of \( \alpha \)-HCH and \( \gamma \)-HCH concentrations. For each PCB congeners and OCPs, the limits of quantification (LOQ) were 0.09 ng g\(^{-1}\) and 0.08 ng g\(^{-1}\) for egg and blood samples respectively. The efficiency recovery for PCB congeners and of OCPs was always above 75% for egg and blood samples.

2.5. Statistical analysis

For statistical analysis, OC concentrations that were below the limit of quantification (LOQ) were estimated at half the LOQ. Firstly, general linear models (GLM) with repeated measurements were carried out to investigate the variations in concentration of OCs and lipid percentage during the nesting season; the dependent variable was the sum of concentrations for a class of OCs or the percentage of egg lipid content, and the independent variable was the nest laying interval between clutches, in days, (the time 0 corresponded to the day when we observed the first clutch for each female). For both egg and blood concentrations, repeated measurements were used to compare data from the same female at different nesting events. Differences in OC concentrations and proportions between 2-year and 3-year RI females were then investigated. General linear mixed models (GLMM) for each tissue (egg and blood) and each sum of class of OCs (the dependent variables of each model) were carried out. We used mixed models because values for the same female at different times (representing different laying events) were correlated; this covariance structure was handled by introducing the individual females as a random effect into the GLMM. GLM with binomial error distribution and logit link function were also performed to assess the effect of RI on the proportion of each class of contaminant analyzed. Finally, GLMM were performed to test the relationship between concentrations in eggs when regressed against corresponding concentrations in blood. Simple regression models were then applied to look at potential correlations between concentrations of OCs in eggs and in blood. The normality of the dependent variables was confirmed prior to the analyses. Computations were performed with the SAS package.

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### Table 1

<table>
<thead>
<tr>
<th></th>
<th>ΣPCBs</th>
<th>ΣDDTs</th>
<th>ΣHCHs</th>
<th>% Lipids</th>
</tr>
</thead>
<tbody>
<tr>
<td><strong>Blood (ng mL(^{-1}))</strong></td>
<td>Mean ± SD</td>
<td>1.26 ± 0.71</td>
<td>0.31 ± 0.22</td>
<td>0.15 ± 0.16</td>
</tr>
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<td></td>
<td>Range</td>
<td>0.86–4.04</td>
<td>0.09–1.04</td>
<td>0.09–0.90</td>
</tr>
<tr>
<td></td>
<td>n</td>
<td>43</td>
<td>44</td>
<td>14</td>
</tr>
<tr>
<td></td>
<td>% of ΣOCs</td>
<td>72.9</td>
<td>18.1</td>
<td>9</td>
</tr>
<tr>
<td><strong>Egg (ng g(^{-1}) wet mass)</strong></td>
<td>Mean ± SD</td>
<td>6.98 ± 5.02</td>
<td>1.44 ± 1.26</td>
<td>0.41 ± 0.26</td>
</tr>
<tr>
<td></td>
<td>Range</td>
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<td>0.08–5.82</td>
<td>0.08–1.06</td>
</tr>
<tr>
<td></td>
<td>n</td>
<td>46</td>
<td>43</td>
<td>44</td>
</tr>
<tr>
<td></td>
<td>% of ΣOCs</td>
<td>79</td>
<td>16.3</td>
<td>4.7</td>
</tr>
<tr>
<td><strong>Egg (ng g(^{-1}) lipid)</strong></td>
<td>Mean ± SD</td>
<td>55.14 ± 43.17</td>
<td>11.03 ± 9.04</td>
<td>3.43 ± 2.13</td>
</tr>
<tr>
<td></td>
<td>Range</td>
<td>4.40–189.87</td>
<td>0.12–37.35</td>
<td>0.08–9.87</td>
</tr>
<tr>
<td></td>
<td>n</td>
<td>46</td>
<td>43</td>
<td>44</td>
</tr>
<tr>
<td></td>
<td>% of ΣOCs</td>
<td>79.2</td>
<td>15.9</td>
<td>4.9</td>
</tr>
</tbody>
</table>
PCBs 153 + 105 are detected in all egg samples (the proportion of congener 105 being estimated to 10% of the peak 153 + 105). ΣDDTs were the second most abundant OCs class measured, represented mainly by the major metabolite pp’-DDE which occurred in 86% samples contrasting with pp’-DDD and pp’-DDE which occurred only in 4.4% and 6.7% of samples respectively.

3.2. Fluctuations during the nesting season

Concentrations in blood remained constant during the nesting season ($P = 0.209$ and $P = 0.610$ for ΣPCBs and ΣDDTs respectively) except for ΣHCHs for which concentrations slightly increased ($P = 0.010$) (SM, Fig. 1). In contrast, concentrations in eggs on a wet-mass basis decreased significantly during the nesting season from one clutch to the next ($P = 0.003$, $P = 0.003$, $P = 0.001$ for ΣPCBs, ΣDDTs and ΣHCHs respectively). The mean percentage of lipid was 12.9% of the egg content (SD = 5.1%, range 3.8–25.5%) but a significant decrease in the percentage of lipid was observed for eggs collected later in the nesting season ($P = 0.005$) (SM, Fig. 1). After lipid normalization, concentrations in eggs still decreased ($P = 0.042$ and $P = 0.007$ for ΣPCBs and ΣDDTs respectively) but become constant for ΣHCHs ($P = 0.192$) (SM, Fig. 1).

3.3. Variation in concentrations with remigration interval

No significant difference was observed between the blood concentrations of the 2 and 3 years remigrant females for ΣPCBs and ΣHCHs ($P = 0.104$ and $P = 0.899$ respectively). Concerning ΣDDTs, a significant difference was observed ($P = 0.017$) with the 3 years remigrant females having higher ΣDDTs blood concentrations than the 2 years females. Concerning eggs, the RI had a significant effect on ΣPCBs and ΣDDTs ($P = 0.017$ and $P = 0.001$ respectively) but not for ΣHCHs ($P = 0.173$) (Fig. 2A). When using chlorination groups as dependent variables, higher chlorinated groups of congeners presented higher concentrations in the 3 years remigrant females with a significant difference for the 6 Cl group ($P = 0.016$) (Fig. 2A). Finally, when comparing the proportion of ΣOCs according to the RI, it appears that the 3 years remigrant females present a higher proportion of high chlorinated groups of congeners (6 Cl group: $P = 0.001$ and 7–9 Cl group: $P < 0.001$) and a significant lower proportion of low chlorinated group of congeners (4–5 Cl group: $P < 0.001$) compared to the 2 years remigrant females (Fig. 2B).

3.4. Relationship between egg and blood concentrations

Significant relationships were found for ΣDDTs and pp’-DDE ($P = 0.0009$ and $P = 0.0001$ respectively) for which concentrations in eggs were positively correlated with their corresponding concentrations in blood (wet-mass basis). For ΣPCBs and ΣHCHs, relationship were not significant ($P = 0.1602$ and $P = 0.0626$, respectively). When taking the most prevalent congeners alone (congeners 153 + 105, 180 and 138) to display the correlation, the relationships were highly significant ($P = 0.0060$, $P = 0.0034$ and $P = 0.0001$, respectively). Simple linear regressions for pp’-DDE and PCB 153 + 105 in eggs compared with concentrations in blood are presented in Fig. 3 and confirmed the statistically significant relationship.

4. Discussion

Leatherbacks in this study present low OC concentrations in blood and eggs probably resulting from their low trophic level with a diet based mainly on jellyfish (Davenport, 1998). These low levels
may also result from the pollution status of the region where turtles are living: nesting grounds in French Guiana and oceanic foraging grounds mainly used by leatherbacks, remote from coastal anthropogenic sources (James et al., 2005b; Doyle et al., 2007). OC concentrations in leatherback eggs are close to those observed in the herbivorous green turtle and lower than those encountered in omnivorous loggerhead turtles (SM, Table 1). In oviparous organisms, early stages of life often exhibit a greater sensitivity to contaminants (Russell et al., 1999). But concentrations of OCs in eggs of this study remain very low and below concentrations that have been described for deleterious effects: decreased viability in clutches of alligators for egg treated with OCPs (11 075 ng g⁻¹ yolk for DDE, Rauschenberger et al., 2004); altered reproduction in sea birds with adverse effects established for 2300 ng g⁻¹ wet mass for ΣPCBs in egg (Fisk et al., 2005); and increased developmental abnormalities in embryos and hatchlings in aquatic turtles eggs from contaminated lakes (3575 ng g⁻¹ and 389 ng g⁻¹ wet mass for ΣPCBs and DDE respectively from Lake Ontario, Bishop et al., 1998). Further investigations would be however needed to assess toxic thresholds at which adverse effects occur in marine turtle eggs. OCs may also impact marine reptiles at later stages. Correlations between clinical health parameters and OCs in juvenile loggerhead sea turtles have indeed suggested that sea turtles may be relatively sensitive to sub lethal effects of OCs, like modulation of the immune system or alteration of protein, ion and carbohydrate homeostasis. This negative effect could occur even at much lower concentrations than those in studies listed above (blood concentration for ΣPCBs and DDE respectively 5.56 ng g⁻¹ and 0.65 ng g⁻¹ wet mass (Keller et al., 2004a), concentrations close to those encountered in leatherbacks from our study). Ward and Lafferty (2004) reported indeed an increasing trend in disease occurrence in marine turtles and environmental pollutants are expected to be partly responsible for these infectious diseases but their exact effect on sea turtle health is little known and need therefore further studies. OC concentrations are expected to vary in blood during periods of high energy requirements because of a release of OCs stored in adipose tissue into the bloodstream after lipid mobilization. OC fluctuations have been effectively shown in blood of turtles and seals after periods of high mobilization of stores during fasting or low food intake (Lydersen et al., 2002; Keller et al., 2004b; Debier et al., 2006). In the leatherback females, lipid mobilization is also likely to occur during migration or vitellogenesis. The nesting season could also be a phase of high energy requirement because of the numerous massive clutches to lay (Girondot and Fretey, 1996) and because females may go through this period with little or no food intake (Fossette et al., 2007; Caut et al., 2008). Blood concentrations for ΣPCBs and ΣDDTs remain constant suggesting no increase of release of this class of OCs from lipid reserves into the bloodstream during the nesting season. A hypothesis is that lipid reserves have already been mobilized during migration or egg production, periods during which the costs are larger than those during the nesting season (Wallace et al., 2005). The fact that sea turtles display an important weight loss between foraging grounds and nesting sites (depletion of 33% of body mass, James et al., 2005b) and the fact that no adipose tissue were found in the flipper of these females at the beginning of nesting season support this hypothesis.

In oviparous species, females often transfer a part of their burden to their eggs due to the lipophilic nature of OCs (Rauschenberger et al., 2002). Indeed, all of the substances detected in leatherback blood were also detected in eggs confirming that maternal transfer. The absence of parallelism between congener patterns in females and their progeny have already been described in marine vertebrates (turtles, birds, mammals; McKenzie et al., 1997, 1999; Miao et al., 2001; Verreault et al., 2006). This suggests a preferential reproductive transfer of lower chlorinated groups from blood to eggs based on the lipophilic nature of the compound being transferred, the more chlorinated and lipophilic compounds being the more difficult to transfer. In our study, maternal transfer for ΣDDTs and the most prevalent PCB congener was depending on concentrations in female blood (the lack of correlation for ΣPCBs is certainly due to the very low occurrence of the majority of congeners in blood samples); eggs concentrate pollutants proportionally to those accumulated by female. In turtles, lipids and proteins in egg-yolk are the primary reserves that will provide energy and building materials to facilitate embryogenesis (Wallace et al., 2006). Eggs are provided with those nutrients and energy during vitellogenesis during which lipophilic OCs are also mobilized and transferred into the developing ova (Wu et al., 2000). Intra-clutch variation is not likely to occur in turtles, since the supply of yolk is simultaneous and equal in follicles forming one clutch (Bowden et al., 2004). But in contrast inter-clutch variation could occur as vitellogenesis of all clutches to be laid does not necessarily happen at the same time. A decreasing trend occurred in OCs concentrations in clutches throughout the nesting season. This could suggest a progressive offloading of the female through its clutches: the first clutch being the most contaminated as the female burden is higher at the beginning of the nesting season. Then the female burden decreases with each clutch produced. A similar decline has been documented in birds that exhibit sequential ovulation for a clutch leading to a decreasing trend in OCs concentrations in eggs of a clutch in relation to laying order (Van den Steen et al., 2009). Our results show also a significant decrease in egg-lipid percentage between clutches suggesting that yolk deposition in follicles is not equal for all clutches to be laid and that the reproductive investment from females into their subsequent clutches decreases. These results are in contrast with other results on green turtles in which the lipid concentrations are constant throughout the nesting season (Hamann et al., 2002). This difference may be due to the particularity of leatherback to have lipid reserves already largely depleted at their arrival on nesting site. Indeed, adipose tissue could not have been sampled in that study probably resulting from the large depletion of body mass (James et al., 2005b). In sea turtles, once vitellogenesis is initiated follicles grow continuously prior to the nesting season (Miller, 1997) but if female lipid reserves decrease, lipid yolk deposition would decrease too: the follicles provisioned with lipids while female fat reserves decrease are likely to have decreased lipid content.

In our study, eggs from females that spent 3 years on the foraging grounds present higher OC concentrations in their eggs and higher proportions of high chlorinated congeners than do the 2 years remigrant females. This difference may be linked first to the length of the RI. Leatherbacks are indeed capital breeders generally spacing out consecutive breeding seasons every 2 or 3 years (Rivalan et al., 2005). Females spending more years on foraging grounds may have higher contaminant burden resulting from a longer trophic contamination and they may therefore transfer more contaminants to their eggs. But the concentrations encountered in the 3 years remigrant females, almost twice as those of the 2 years remigrant females, cannot simply be explained by an additional year of trophic contamination. The difference in egg concentrations and patterns between remigrant females may also be linked to the location of the foraging grounds. Feeding locations have already been shown to be responsible for the inter-individual difference in OC concentrations among females and to influence the levels and patterns of OCs transferred to eggs (Alava et al., 2006). Turtles foraging in offshore areas have been shown to be less exposed than turtles foraging closer to the coast (Day, 2003) and the distributions of PCBs found in the open sea are known to be oriented towards less chlorinated congeners which are relatively more volatile than highly chlorinated congeners found in.
continental shelf waters (Dachs et al., 1997). As congeners patterns in eggs of the 2 years remigrant females is oriented toward lower chlorinated congeners, therefore more volatile congeners, and that their OCs concentrations are lower than the 3 years remigrant females, the 2 years remigrant females are thought to forage in more oceanic area compared to the 3 years remigrant females. Foraging areas for leatherbacks are located all around the North Atlantic Ocean (Houghton et al., 2006; Doyle et al., 2007) in both offshore and more coastal areas likely to differ in levels of contamination due to anthropogenic activity in coastal zones. The different concentrations between remigrant females may finally be the result of complex interaction between length of the RI and location of foraging grounds. In a previous study, leatherback females have indeed been shown to present different isotopic signatures according to their RI suggesting that remigrant females use different foraging grounds. Isotope signatures could provide information on both the latitude, and the pelagic vs. neritic nature of the foraging grounds. Isotope signatures could provide information on both the latitude, and the pelagic vs. neritic nature of the foraging ground (Kelly, 2000; Takai et al., 2000) and it has been possible to reveal that the 2 years remigrant females seem to feed in a more northern and/or offshore area and the 3 years remigrant females in a more southern and/or coastal areas (Caut et al., 2008).

5. Conclusion

This study provides the first ecotoxicological data for OCs in living leatherbacks. The results show important differences in term of contamination level between females according to their RI. This highlights the importance of the time spent on foraging grounds and the level of contamination and location of these foraging grounds. The level of contamination remains however low in all females’ samples but despite these low contaminations, a maternal transfer of OCs occurs and eggs are contaminated. In oviparous organisms, early stages of life often exhibit a greater sensitivity to contaminants but despite the fact that level of hatchling success on Yalimapo beach has been shown to be low, the causal link between OC and embryonic mortality is unlikely on this site.

Acknowledgements

We thank DIREN-Guyane for the use of the facilities of the Amana Nature Reserve. Samples were obtained during a joint field mission with the CEPE-CNRS UPR 9010, 67087 Strasbourg and we thank all those who helped in the field work: X. Desespe (ESR, Paris), J.Y. Georges and S. Foissette (CEPE, Strasbourg). The field work was conducted under legal permit from the French government, via a collaborative agreement with the group of Dr. J.Y. Georges. The subsequent transport of the field samples was authorized by a permit from the French government (Préfecture de la Guayante) granted in 2006. The authors are grateful to M. Louvet for valuable technical assistance in organochlorine analysis. K. Das is supported by FRS-FNRS. This is a MARE publication 186.

Appendix A. Supplementary material


References


