Persistent organic pollutants in the green sea turtle

*Chelonia mydas*: nesting population variation, maternal transfer, and effects on development

Jason P. van de Merwe1,5,*, Mary Hodge2, Joan M. Whittier3,6, Kamarruddin Ibrahim4,7, Shing Y. Lee1

1Griffith School of Environment and Australian Rivers Institute, Griffith University Gold Coast campus, Gold Coast, Queensland 4222, Australia
2Queensland Health Scientific Services, Queensland Government, Coopers Plains, Queensland 4108, Australia
3School of Biomedical Sciences, University of Queensland, St. Lucia, Queensland 4072, Australia
4Turtle and Marine Ecosystems Centre, Department of Fisheries Malaysia, Rantau Abang, Terengganu 23050, Malaysia
5Present address: Centre for Marine Environmental Research and Innovative Technology, City University of Hong Kong, Kowloon, Hong Kong SAR China
6Present address: School of Medicine, University of Tasmania, Sandy Bay, Tasmania 7005, Australia
7Present address: Marine Park Department of Malaysia, Ministry of Natural Resources and Environment, Federal Government Administration Centre, Putrajaya 62574, Malaysia

ABSTRACT: Persistent organic pollutants (POPs), such as polychlorinated biphenyls (PCBs), organochlorine pesticides (OCPs) and polybrominated diphenyl ethers (PBDEs), have a wide range of toxic effects on humans and wildlife, and have been reported in a number of endangered sea turtle populations. The present study screened for POPs in a green sea turtle *Chelonia mydas* population in Peninsular Malaysia and investigated the maternal transfer and effects of POPs on embryonic development. At the Ma’Daerah Turtle Sanctuary, blood, eggs and hatchling blood were collected from 11 nesting female *C. mydas*. Samples were analysed for 83 PCBs, 23 OCPs and 19 PBDEs using gas chromatography with tandem mass spectrometry. The chemical profiles of eggs from individual turtles were significantly different, indicating variable contaminant uptake during foraging. There was evidence of maternal transfer of POPs to eggs and hatchlings, with significant correlations in sum of PCBs (ΣPCB), sum of PBDEs (ΣPBDE), γ-hexachlorocyclohexane (γ-HCH), trans-chlordane and mirex concentrations between maternal blood and eggs (p < 0.05, R2 < 0.71), between eggs and hatchling blood (p < 0.05, R2 < 0.83), and between maternal and hatchling blood (p < 0.05, R2 < 0.61). In addition, there was congener-specific transfer of PCBs with less lipophilic congeners (e.g. PCB 99) more readily transferred to hatchlings than the more lipophilic congeners (e.g. PCBs 180 + 193). There was also a significant correlation between increasing egg POP concentration and decreasing hatchling mass:length ratio. POPs may therefore have subtle effects on the development of *C. mydas* eggs, which may compromise offshore dispersal and predator avoidance.

KEY WORDS: *Chelonia mydas* · Persistent organic pollutants · Maternal transfer · Contamination profiles

INTRODUCTION

Persistent organic pollutants (POPs), such as polychlorinated biphenyls (PCBs), organochlorine pesticides (OCPs) and polybrominated diphenyl ethers (PBDEs), have been reported in a number of sea turtle populations. In addition, these chemicals have been correlated with health parameters in free-ranging sea turtles (Keller et al. 2004a, 2006a,b), and embryonic development in other oviparous reptiles (Bergeron et
al. 1994, Guillette et al. 1996, Guillette & Crain 1996, Bishop et al. 1998, Willingham & Crews 1999, de Solla et al. 2008). As sea turtles generally do not feed during the breeding season, the contamination of sea turtles will be influenced by the contamination of their foraging areas (Bjorndal 1985, 1997). Furthermore, due to the lipophilic properties of these compounds and the mobilisation of lipids by sea turtles for egg production (Kwan 1994), it is likely that POPs are transferred from nesting females to eggs and hatchlings during reproduction. This may have consequences for hatching development, due to the high sensitivity of developing sea turtle embryos to disturbance. However, there is currently limited information on POPs in sea turtle populations or the maternal transfer of these chemicals and the effects this may have on the reproductive success of nesting populations.

Green turtle *Chelonia mydas* nesting populations are generally comprised of individuals that have migrated from a wide range of foraging grounds (Liew et al. 1995, Cheng 2000, Godley et al. 2002, Seminoff et al. 2008). During their reproductive years, *C. mydas* show strong fidelity to these foraging and breeding sites, which can be up to thousands of kilometres apart (Carr 1964, Carr & Carr 1972, Limpus et al. 1992, Lohmann et al. 1999). The contamination of marine vertebrates occurs nearly exclusively though feeding (Newman & Unger 2003). Therefore, as nesting female *C. mydas* do not feed during migration or nesting (Bjorndal 1997), their POP concentrations are likely to reflect the contamination in their foraging areas. The contamination of a nesting *C. mydas* population is therefore expected to be variable and presents considerable conservation challenges, in terms of managing contamination in remote foraging areas.

Maternal transfer of POPs to offspring is well documented in marine mammals (Aguilar & Borrell 1994, Debier et al. 2003, Miranda-Filho et al. 2007), birds (Fisk & Johnston 1998) and oviparous reptiles (Kelly et al. 2008). In freshwater snapping turtles *Chelydra serpentina*, egg concentrations of PCBs, DDTs, mirex, octachlorostyrene and HCB were significantly correlated with concentrations in the blood, liver, muscle and adipose tissue of adult females (Hebert et al. 1993, Pagano et al. 1999, Kelly et al. 2008). Similarly, in leatherback sea turtles *Dermochelys coriacea*, significant correlations have been found between blood and egg samples containing DDTs, PCBs and PBDEs (Stewart et al. 2007). These results indicate that nesting female turtles can incorporate POPs into eggs during vitellogenesis and oviposition (Guillette & Crain 1996). However, no studies to date have investigated the further transfer of POPs from eggs to embryos. Furthermore, in mammals there is evidence of selective transfer of POPs to offspring, with the less lipophilic compounds being more readily transferred (Miranda-Filho et al. 2009). However, this remains poorly understood in oviparous reptiles, including sea turtles.

During growth, sea turtle embryos mobilise lipid reserves from the egg yolk to meet developmental requirements (Miller 1985). It is therefore likely that the lipophilic POP contaminants in the yolk are transferred to the hatchlings during this process. In *Chelydra serpentina*, strong correlations have been observed between egg POP concentrations and abnormalities in hatchlings (Bishop et al. 1994, 1998). This association between hatching abnormalities and egg POP concentrations may be due to the disruptive effects of these chemicals on the endocrine system during development (Miller 1985, Guillette & Crain 1996). However, a relationship between POP concentrations and hatching abnormalities has not been previously investigated for *Chelonia mydas* or any other sea turtle species, and warrants further investigation.

The aim of the present study was to analyse POP contamination in a nesting population of *Chelonia mydas*, and investigate the maternal transfer of these chemicals to eggs and embryos and the effects this may have on hatching development.

**MATERIALS AND METHODS**

**Egg and blood collection.** In June and July 2004, eggs and blood were collected from 11 adult female *Chelonia mydas* nesting at the Ma’Daerah Turtle Sanctuary, Terengganu, Malaysia (Table 1). Three eggs were randomly collected from each clutch at the time of oviposition. Care was taken to minimise egg contact with the sand to avoid external contamination. Each egg was wrapped in hexane-rinsed aluminium foil and kept frozen (–20°C) until analysis. Following oviposition, the curved carapace length (CCL) and mass of each nesting female were measured according to Bolten (1999).

At the time of oviposition, 2 to 5 ml of blood was collected from the dorsal cervical sinuses in the neck of the nesting female using a 10 ml syringe with a 21G × 1.25” (3.175 cm) needle (Owens & Ruiz 1980). The eggs not taken for chemical analysis were transferred to a shaded hatchery within 2 h of oviposition and incubated at a depth of 60 cm. Once the hatchlings emerged, a sample of 5 individuals was randomly collected from each nest. Blood samples (300 to 500 µl) were taken from the dorsal cervical sinuses in the neck using an insulin syringe and pooled for each nest. All blood samples were immediately transferred to glass lithium heparin vacutainer tubes and kept frozen (–20°C) until analysis.

**Clutch incubation and hatching morphometrics.** The incubation of clutches in the hatchery was monitored closely around the expected time of emergence.
Immediately following emergence, hatching mass ($\pm 0.01 \text{ g}$) and the straight carapace length (SCL) were measured ($\pm 0.01 \text{ mm}$), and the mass:SCL ratio was calculated for a random sample of 10 hatchlings from each nest. Carapace, plastron and head scutes were also recorded and each hatching was assigned an abnormality index based on the number of deviations from a normal scute pattern, as described by Pritchard & Mortimer (1999) and Miller (1985). The percentage of hatchlings with abnormal scute counts and the mean ($\pm$ SE) of each of hatching morphometric parameter were also calculated for each nest. Seven days after emergence, nests were excavated to determine hatching and emergence success according to Miller (1999).

**Chemical analysis.** Entire egg contents (excluding shell) and whole blood samples were analysed for 83 PCB congeners, 23 OCPs and 19 PBDE congeners using gas chromatography with tandem mass spectrometry, following methods developed by van de Merwe et al. (2009a). Briefly, samples were extracted in an accelerated solvent extractor (Dionex) with dichloromethane (2009a). Samples were analysed on a Varian 3800 gas chromatograph fitted with a 60 m VF-5MS GC capillary (0.25 mm interior diameter and 0.32 µm film thickness) and a Saturn 2200 mass spectrometer. A 1079 programmable temperature vapourising (PTV) injector was used to inject 20 µl of each sample and ionisation was performed using electron impact. The limit of detection (LOD) was compound- and sample-specific, although for most compounds, it was $<10 \text{ pg g}^{-1}$ for egg samples and $<35 \text{ pg g}^{-1}$ for blood samples. The analysed values for National Institute of Standards and Technology human serum Standard Reference Material (SRM 1589a) were 84.1 $\pm 0.1 \%$ of the certified concentrations (Schantz et al. 2007). Values for avian egg control samples (QC04-ERM1; common murre Uria aalge and thick-billed murre U. lomvia) were 79.5 $\pm 0.1 \%$ of the reference concentrations (Vander Pol et al. 2007). Recoveries of mass-labelled internal standards ranged from 30 to 96%, and were <60% only for the higher-chlorinated PCBs.

**Statistical analysis.** The mean ($\pm$ SE) of each POP compound was calculated for maternal blood, eggs and hatching blood. To investigate differences in POP contamination profiles among clutches, analysis of similarity (ANOSIM) was performed on a Bray-Curtis similarity matrix with no data transformation using Primer v5 (PRIMER-E). Each POP compound was entered as a separate variable and each egg was analysed as an individual sample. Significant differences in the ANOSIM ($p < 0.05$) indicated samples that were different in terms of both the presence and concentration of all 125 POP compounds analysed. A non-metric multi-dimensional scaling (NMDS) plot was also constructed to further illustrate differences in POP contamination profiles between clutches (PRIMER-E).

To investigate transfer of POPs from nesting females to eggs, and from eggs to hatchlings, linear regressions were performed between maternal blood and egg POP concentrations, and between egg and hatching blood POP concentrations, respectively. In addition, linear regressions were performed between maternal blood and hatching blood POP concentrations. To standardise for lipid content, POP concentrations were presented in ng g$^{-1}$ lipid. Only the compounds detected in maternal blood, eggs and hatching blood were analysed in this way. The sum of PCB ($\Sigma$PCB) and PBDE ($\Sigma$PBDE) concentrations were calculated by summing the 83 PCB and 19 PBDE congeners analysed in each sample, respectively. The remaining OCPs were investigated individually.

### Table 1. *Chelonia mydas*. Nesting and incubation details of 11 nesting females. CCL: curved carapace length, SCL: straight carapace length. Data are mean $\pm$ SE (range)

<table>
<thead>
<tr>
<th>Tag</th>
<th>Nesting details</th>
<th>Incubation details</th>
<th>Mass:SCL (g mm$^{-1}$)</th>
</tr>
</thead>
<tbody>
<tr>
<td></td>
<td>CCL (cm)</td>
<td>Eggs Egg mass (g)</td>
<td>Hatchling mass (g)</td>
</tr>
<tr>
<td>MY4133</td>
<td>112.5</td>
<td>130 34.6 $\pm$ 0.8</td>
<td>19.5 $\pm$ 0.2 18.0–21.0</td>
</tr>
<tr>
<td>TF3315</td>
<td>111</td>
<td>131 37.5 $\pm$ 0.8</td>
<td>22.1 $\pm$ 0.2 20.7–23.7</td>
</tr>
<tr>
<td>No tags</td>
<td>99</td>
<td>126 36.5 $\pm$ 0.8</td>
<td>21.6 $\pm$ 0.2 19.2–22.9</td>
</tr>
<tr>
<td>IF2565</td>
<td>95</td>
<td>39   33.5 $\pm$ 0.3</td>
<td>19.8 $\pm$ 0.1 17.8–20.7</td>
</tr>
<tr>
<td>TF3321/TF3319</td>
<td>112</td>
<td>134 37.6 $\pm$ 0.3</td>
<td>23.8 $\pm$ 0.2 21.3–25.2</td>
</tr>
<tr>
<td>TF3351/MY1551</td>
<td>103</td>
<td>106 34.4 $\pm$ 0.8</td>
<td>21.7 $\pm$ 0.2 20.1–23.4</td>
</tr>
<tr>
<td>MY1552</td>
<td>93</td>
<td>67   39.7 $\pm$ 0.4</td>
<td>23.9 $\pm$ 0.1 22.9–24.9</td>
</tr>
<tr>
<td>MY1602</td>
<td>99.5</td>
<td>130 36.4 $\pm$ 0.9</td>
<td>20.1 $\pm$ 0.2 18.5–21.4</td>
</tr>
<tr>
<td>MY0297/MY1502</td>
<td>88.9</td>
<td>101 34.4 $\pm$ 1.3</td>
<td>23.1 $\pm$ 0.2 18.0–21.1</td>
</tr>
<tr>
<td>IF2720</td>
<td>113</td>
<td>143 34.7 $\pm$ 0.8</td>
<td>20.4 $\pm$ 0.2 18.9–21.8</td>
</tr>
<tr>
<td>MY1555</td>
<td>98.4</td>
<td>80   35.7 $\pm$ 0.6</td>
<td>21.8 $\pm$ 0.2 18.6–22.9</td>
</tr>
</tbody>
</table>
To investigate congener-specific maternal transfer of PCBs, the percent of the $\Sigma$PCB concentration was calculated for each of the major congeners and the means for maternal blood, eggs and hatching blood were compared using ANOVA. In cases of ANOVA significance ($p < 0.05$), Tukey’s post hoc test was used to determine which tissues were different.

To investigate the effect of POP contamination on hatching development, regressions were performed between hatching development parameters and the mean egg sum of POP ($\Sigma$POP) concentration for each clutch. Linear regressions were performed between mean egg $\Sigma$POP concentration and hatching success, emergence success and the percentage of abnormal hatchlings, and means of hatchling mass, SCL, mass: SCL ratio or abnormality index. The influence of egg mass on embryonic development was investigated by performing regressions between mean initial egg mass and any hatchling or nest variable that showed significant regression with mean egg $\Sigma$POP concentration.

### RESULTS

#### POP concentrations in nesting females, eggs and hatchlings

The $\Sigma$POP concentrations ranged from 727 to 2835 pg g$^{-1}$ wet mass for nesting female blood, 992 to 1569 pg g$^{-1}$ wet mass for eggs and 1429 to 3321 pg g$^{-1}$ wet mass for hatchling blood (Table 2). The concentrations of PCBs were generally the highest and congeners 99, 118, 128, 138, 153, 180, 183 and 193 were the most abundant. The most abundant PBDE congeners were 47, 99 and 153, and trans-chlordane, mirex and $\gamma$-hexachlorocyclohexane ($\gamma$-HCH) were the only OCPs detected in all 3 sample types. On a wet

![Fig. 1. Chelonia mydas. NMDS plot of egg persistent organic pollutant (POP) profiles for 11 nesting females. Eggs from the same clutch are indicated by the same symbol. Grouping of eggs from the same clutch indicate that within-clutch variation in POP contamination profiles is less than the variation between clutches. There is also some grouping of contamination profiles between clutches, as indicated by clutches with the same symbol shape (either open or filled). Stress = 0.09, indicating the plot is a good representation of the actual differences between samples.](image)

<table>
<thead>
<tr>
<th>POP compound</th>
<th>In egg</th>
<th>Mean ± SE</th>
<th>Range$^a$</th>
<th>In maternal blood</th>
<th>Mean ± SE</th>
<th>Range$^a$</th>
<th>In hatching blood</th>
<th>Mean ± SE</th>
<th>Range$^a$</th>
</tr>
</thead>
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<tr>
<td>PCB 99</td>
<td>17.4 ± 1.4</td>
<td>13.0–27.7</td>
<td>39.4 ± 4.8</td>
<td>22.0–64.3</td>
<td>81.4 ± 8.2</td>
<td>50.0–124.0</td>
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<tr>
<td>PCB 118</td>
<td>16.4 ± 1.3</td>
<td>11.6–23.6</td>
<td>33.9 ± 5.8</td>
<td>18.4–83.6</td>
<td>58.0 ± 7.1</td>
<td>36.0–105.8</td>
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<tr>
<td>PCB 128</td>
<td>14.1 ± 1.2</td>
<td>9.9–20.1</td>
<td>28.7 ± 3.8</td>
<td>16.3–54.4</td>
<td>35.4 ± 2.5</td>
<td>23.1–47.1</td>
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<tr>
<td>PCB 138 (+ 158)$^b$</td>
<td>45.6 ± 8.9</td>
<td>17.0–94.1</td>
<td>62.9 ± 13.9</td>
<td>25.8–174.6</td>
<td>101.3 ± 20.9</td>
<td>46.7–226.7</td>
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<tr>
<td>PCB 153 (+ 132)$^b$</td>
<td>65.8 ± 13.1</td>
<td>25.8–136.4</td>
<td>93.9 ± 24.4</td>
<td>30.6–288.6</td>
<td>171.9 ± 43.1</td>
<td>64.7–462.0</td>
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<tr>
<td>PCB 180 (+ 193)</td>
<td>53.5 ± 7.5</td>
<td>33.9–100.4</td>
<td>114.0 ± 13.5</td>
<td>72.0–202.0</td>
<td>91.1 ± 11.9</td>
<td>35.0–165.2</td>
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<tr>
<td>PCB 183</td>
<td>14.8 ± 1.8</td>
<td>9.7–25.9</td>
<td>52.0 ± 12.1</td>
<td>19.8–134.2</td>
<td>63.8 ± 12.0</td>
<td>28.0–142.4</td>
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<tr>
<td>$\Sigma$PCB</td>
<td>553.6 ± 54.6</td>
<td>392.8–839.4</td>
<td>578.9 ± 85.6</td>
<td>316.4–1206.5</td>
<td>850.8 ± 105.2</td>
<td>559.4–1456.6</td>
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<tr>
<td>trans-chlordane</td>
<td>18.3 ± 0.9</td>
<td>13.8–22.3</td>
<td>21.3 ± 2.7</td>
<td>11.5–33.4</td>
<td>42.5 ± 4.8</td>
<td>17.1–68.4</td>
<td></td>
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<tr>
<td>Mirex</td>
<td>9.4 ± 1.1</td>
<td>&lt;LOD to 12.8 (10)</td>
<td>161 ± 43.0</td>
<td>&lt;LOD to 476.3 (10)</td>
<td>132.4 ± 34.0</td>
<td>&lt;LOD to 340.6 (10)</td>
<td></td>
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<tr>
<td>$\gamma$-HCH</td>
<td>172.3 ± 7.4</td>
<td>137.8–207.8</td>
<td>501.1 ± 59.7</td>
<td>231.3–899.8</td>
<td>939.0 ± 63.4</td>
<td>634.3–1256.2</td>
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<tr>
<td>PBDE 47</td>
<td>21.5 ± 1.7</td>
<td>11.1–28.1</td>
<td>13.3 ± 1.2</td>
<td>7.2–20.1</td>
<td>27.7 ± 2.7</td>
<td>14.9–41.4</td>
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<tr>
<td>PBDE 99</td>
<td>32.0 ± 3.6</td>
<td>12.0–54.8</td>
<td>21.3 ± 3.7</td>
<td>5.0–52.8</td>
<td>55.3 ± 12.1</td>
<td>8.9–131.8</td>
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<tr>
<td>PBDE 153</td>
<td>27.4 ± 1.2</td>
<td>20.8–35.2</td>
<td>86.2 ± 10.3</td>
<td>37.5–151.4</td>
<td>&lt;LOD</td>
<td></td>
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<tr>
<td>$\Sigma$PBDE</td>
<td>129.3 ± 8.1</td>
<td>61.7–163.8</td>
<td>120.8 ± 14.1</td>
<td>57.5–224.3</td>
<td>83.0 ± 14.4</td>
<td>23.8–173.2</td>
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<tr>
<td>$\Sigma$POP</td>
<td>1286.8 ± 66.6</td>
<td>992.3–1568.7</td>
<td>1383.1 ± 190.0</td>
<td>727.1–2834.9</td>
<td>2047.6 ± 175.8</td>
<td>1428.9–3320.6</td>
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<tr>
<td>% Lipids</td>
<td>8.9 ± 0.2</td>
<td>6.8–10.9</td>
<td>1.5 ± 0.1</td>
<td>1.1–2.1</td>
<td>1.6 ± 0.1</td>
<td>1.4–2.0</td>
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</tr>
</tbody>
</table>

$^a$ Number of samples above LOD indicated in parentheses if <11

$^b$ Congeners in parentheses co-elute. However, due to their rarity in environmental samples, they are unlikely to be significantly contributing to the reported concentrations.
mass basis, POP concentrations were generally highest in hatchling blood, followed by maternal blood and eggs. However, the number of POP compounds detected in eggs (54) was higher than in the maternal blood (21) and hatchling blood (20).

The egg POP contamination profiles in the present study were significantly different between clutches (ANOSIM: $R = 0.993$, $p = 0.001$), and the NMDS plot illustrated distinct groups of eggs from the 11 nesting female *Chelonia mydas* (Fig. 1).

**Maternal transfer of POPs and correlations with hatchling parameters**

There were significant correlations between maternal blood and eggs, and between eggs and hatchling blood for $\Sigma$PCBs (Fig. 2; $R^2 > 0.93$, $p < 0.005$), $\Sigma$PBDEs (Fig. 3; $R^2 > 0.71$, $p < 0.005$) and the OCPs $\gamma$-HCH, trans-chlordane and mirex (Fig. 4; $R^2 > 0.83$, $p < 0.005$). This represented all compounds that were detected in all 3 tissue types. In addition, maternal blood was significantly correlated to hatchling blood for all of the POP compounds and groups ($R^2 > 0.61$, $p < 0.005$). Significant differences were also observed in the PCB
congener composition between maternal blood, eggs and hatchling blood (Fig. 5; p < 0.05). PCB 99 made up a significantly greater percentage of the ΣPCB concentration in the hatchling blood (9.9 ± 1.3%), compared to eggs and maternal blood (6.2 ± 1.1% and 7.1 ± 1.3%, respectively), while PCBs 180 + 193 made up a larger percentage of the ΣPCB concentration in the maternal blood and eggs (20.5 ± 2.5% and 17.9 ± 2.4%, respectively), compared to hatchling blood (10.8 ± 1.6%).

The mass:SCL ratio of hatchlings was significantly correlated with the mean egg ΣPOP concentration (Fig. 6; R² = 0.65, p = 0.02). However, there were no relationships between mean egg ΣPOP concentrations and hatching success, emergence success, percentage of abnormal hatchlings, hatchling mass, hatchling SCL or hatchling abnormality index (p > 0.05).

**DISCUSSION**

**POPs in green turtles of Peninsular Malaysia**

The mean ΣPOP concentrations in maternal blood (1287 ± 67 pg g⁻¹ wet mass) and eggs (1383 ± 190 pg g⁻¹ wet mass) in the present study were similar to levels observed in concurrent studies on *Chelonia mydas* eggs from Peninsular Malaysia (1097 ± 433 pg g⁻¹ wet mass; van de Merwe et al. 2009b) and blood from rehabilitating *Chelonia mydas* subadults from Australia (929 ± 170 pg g⁻¹ wet mass; van de Merwe et al. 2009c). In relation to other sea turtle species, lipid-normalised maternal and hatchling blood concentrations of mirex in the present study (6 to 23 and 2 to 18 ng g⁻¹ lipid, respectively) were similar to concentrations in the blood of free-ranging juvenile loggerhead turtles *Caretta caretta* from North Carolina, USA (<LOD to 20.5 ng g⁻¹ lipid; Keller et al. 2004b). However, the concentrations of *trans*-chlordane (0.7 to 2.2 and 1.1 to 4.2 ng g⁻¹ lipid for maternal and hatchling blood, respectively) and ΣPCB (26 to 57 and 37 to 84 ng g⁻¹ lipid for maternal and hatchling blood, respectively) in the present study were up to 100 times lower than concentrations in *Caretta caretta* juveniles (LOD to 12.8 and 1020 to 2810 ng g⁻¹ lipid, respectively; Keller et al. 2004b). This could be expected due to the higher trophic level occupied by *Caretta caretta* and the vast geographical separation of the populations sampled. In addition, the ΣPCB wet mass concentrations in the eggs of the present study (0.39 to 0.84 ng g⁻¹ wet mass) were considerably lower than *Chelonia mydas* eggs from Ascension Island (20 to 220 ng g⁻¹ wet mass; Thompson et al. 1974). Similarly, lipid-normalised egg ΣPCB concentrations (2.5 to 5.9 ng g⁻¹ lipid) were significantly lower than *Caretta caretta* eggs sampled in Florida, USA (7 to 3930 ng g⁻¹ lipid; Alava et al. 2006).

The relative concentrations of the different POP compounds in the present study were also comparable with the contamination profiles of mussels and fish analysed within the Southeast Asian region (Kannan et al. 1995, Tanabe et al. 2000). These studies found regional variation in these concentrations, although PCBs and DDTs were generally highest, while chlordanes and HCHs were lower. The absence of DDTs in the present study is a conspicuous discrepancy from these previous studies on mussels and fish. However, the detection of DDE in *Chelonia mydas* eggs from a concurrent study (J. P. van de Merwe unpubl.) in Peninsular Malaysia and the reliability of the analytical method indicate that the absence of DDE is unlikely to be due to analytical error. Satellite tracking studies on the Ma’Daerah and adjacent *C. mydas* populations indicate that these turtles often migrate to foraging areas within Southeast Asia far removed from
industrialised cities and agricultural areas (Liew et al. 1995, van de Merwe et al. 2009d). This may reduce the exposure of the *C. mydas* in the present study to DDTs, resulting in the absence of these chemicals in eggs and blood of the turtles analysed. In addition, the presence of *trans*-chlordane in the present study without *cis*-chlordane or the nonachlors is unusual for sea turtle samples (e.g. Keller et al. 2004b,c). However, detection of chlordanes at low concentrations is difficult using electron-impact ionisation due to extensive fragmentation. The concentration of *trans*-chlordane in the present study may therefore represent an underestimation of the total chlordanes in this *C. mydas* population.

The significantly different POP profiles in the *Chelonia mydas* clutches gives an indication of the variation in POP contamination of this nesting population. As discussed in the ‘Introduction’, the contamination of adult female green turtles will be strongly influenced by the contamination of the areas in which they forage. A recent study on this Ma’Daerah nesting population found that individual *C. mydas* migrate to a number of foraging grounds (in Vietnam, Indonesia and Borneo Malaysia) that may be under threat from varying levels of chemical contamination (van de Merwe et al. 2009d). The different POP profiles observed in the present study may therefore reflect different foraging grounds used by this nesting population. However, the amount of variation in POP profiles that represents animals using different foraging areas remains unclear. There are a number of factors that may complicate this interpretation, such as the age of the animals (an indication of duration of exposure) and the variations in contamination of food sources within foraging areas. Nevertheless, the variation in POP profiles observed in the present study indicates that animals are being exposed to different types and concentrations of POP compounds, and highlights the challenge of managing contamination in *C. mydas* nesting populations.

**Maternal transfer of POPs**

The correlations between maternal blood, eggs and hatchling blood POP concentrations indicates that these chemicals are being transferred from nesting female *Chelonia mydas* to eggs and hatchlings. Sea turtles mobilise lipids to meet the metabolic demands of migration and egg production (Kwan 1994, Hamann et al. 2002). Therefore, due to the lipophilic properties of POPs, it is likely that these chemicals are transferred from nesting females as lipids are mobilised for yolk production. The transfer of POPs from nesting females to eggs and hatchlings may be a mechanism of chemical off-loading for adult female *C. mydas*. However, due to the vulnerability of developing embryos to disturbance, the transfer of chemicals may also have conservation implications for *C. mydas* populations.

The results of the present study also indicate that the transfer of POPs from eggs to hatchlings may be compound-specific, related to lipophilicity. PCB 99 made up a larger percentage of the EPCB concentration in hatchling blood, compared to congeners 180 and 193, indicating that the less lipophilic congeners may be preferentially transferred from eggs to hatchlings. Alternatively, the lower proportion of PCB 180 and 193 in the blood of hatchlings may indicate that the highly lipophilic congeners are more quickly accumulated into the fatty tissue of hatchlings and/or that hatchlings have mechanisms for metabolising these congeners. These results are supported by previous studies on seals that found strong inverse correlations between the log *K*<sub>ow</sub> (i.e. lipophilicity) and transfer rate from dams to pups (Miranda-Filho et al. 2009). This congener-specific transfer of PCBs may have implications for hatching development and survival. Generally speaking, less-chlorinated PCBs have higher toxic equivalency factors (TEFs) than more-chlorinated PCBs, although the position of the chlorine atoms also influences congener toxicity (Van den Berg et al. 1998). Preferential transfer of the less lipophilic, more toxic PCB congeners may therefore increase the risks of PCB exposure in *Chelonia mydas* hatchlings.

The strong correlations between maternal blood and egg POP concentrations also support the use of egg samples to predict POP contamination of adult female *Chelonia mydas*. Furthermore, due to the low variability in POP concentrations within clutches, a small sample (e.g. 3 eggs, as in the present study) would be sufficient for this analysis. The collection of eggs for contamination screening is a much simpler sampling method than collecting blood from nesting females. Nesting under bushes and shrubs, as well as the short window of opportunity between oviposition and when the front flippers are engaged in nest filling, limits the access for blood sampling of nesting female sea turtles. Furthermore, eggs (30 to 50 g) represent a relatively higher sample mass compared to blood samples (10 to 15 g), and have significantly higher lipid content. This allows lower LODs to be reached under current methods and therefore increases the likelihood of identifying more compounds in each sample. Although egg sampling is destructive as it prevents the incubation of the eggs sampled, 3 eggs from a clutch represents <0.5% of the reproductive output for each adult female *C. mydas* over a nesting season. Alternatively, the use of undeveloped eggs at the end of incubation could be explored to further reduce the destructive nature of egg sampling. However, further studies would need to be performed to determine if the POP concentrations in the eggs change over the duration of incubation and
whether there were any external nest influences on egg POP concentrations during this period.

Effects of POPs on hatching development

The significant negative correlation between egg POP concentration and mass:SCL ratio (a measure of body condition) of hatchlings (see Fig. 6) may indicate an effect of POPs on the embryonic development of *Chelonia mydas*. A previous study on juvenile loggerhead turtles *Caretta caretta* reported negative correlations between organochlorine blood concentrations and body condition, as well as indicators of immune function, anaemia, kidney function, metabolism and glucose regulation (Keller et al. 2006b). The mass:SCL ratio may therefore be an early indicator of the effects of POPs on sea turtle development, although direct effects of POPs on sea turtles have not been established. Podreka et al. (1998) exposed *Chelonia mydas* eggs to 1,1-bis-(4-chlorophenyl)-2,2-dichloroethene (p,p′-DDE), but found no effect on sex reversal, hatch success or hatchling development for predicted egg concentrations of up to 543 ng g⁻¹ wet mass. However, they did not investigate more subtle effects of DDE on hatchling growth and development. Further manipulative experiments may be warranted to investigate non-lethal effects of POPs on the development and survival of sea turtle hatchlings.

The mechanism(s) behind the possible effects of POPs on mass:SCL ratios of *Chelonia mydas* hatchlings can only be speculative. In a recent study on glaucous gulls *Larus hyperboreus*, females contaminated with POPs produced eggs with reduced lipid content (although no reduction in overall egg mass), indicating a disruption to resource allocation during egg production (Verboven et al. 2009). This may be due to effects on endocrine processes, such as production of estrogen in the ovaries, which stimulates the liver to produce vitellogenin, a precursor to egg yolk formation (Etches 1996). If similar mechanisms occur in sea turtles, then less lipids in eggs of contaminated nesting females would be available to hatchlings for growth. Alternatively, hatchlings developing in contaminated eggs may be less able to utilise the yolk for growth because they have other metabolic demands associated with dealing with toxic stress. These factors may lead to reduced mass of hatchlings, reflected by the reduced mass:SCL ratios observed in the present study. Although there are a number of environmental factors, such as nest temperature and moisture, that can also influence the growth of oviparous reptile embryos (Booth & Astill 2001, Packard & Packard 2001), the hatchery incubation of the eggs in the present study ensured that environmental factors remained similar between clutches. Furthermore, there was no significant correlation between hatching mass:SCL and initial egg mass (linear regression: $R^2 = 0.22, p = 0.15$), reducing the likelihood of maternal factors being responsible for the differences in mass:SCL ratios observed in the present study.

In sea turtle hatchlings, a reduced mass:SCL ratio may compromise the duration of offshore dispersal. Sea turtle hatchlings emerge from nests in an energetic frenzy and do not feed in the first 3 to 5 days as they swim continuously to the safer open ocean waters (Dial 1987, Wyneken & Salmon 1992). The energy demands of offshore dispersal must therefore be met by the store of residual yolk that they have at the time of emergence (Miller 1985). A decrease in mass:SCL ratio of sea turtle hatchlings could indicate reduced residual yolk and hence reduce the duration of offshore dispersal. In addition to compromising offshore dispersal, reduced mass:SCL ratio may also indicate increased susceptibility to predation. There is evidence that larger turtle hatchlings survive better than their smaller conspecifics due to superior locomotion and increased mobility and agility, which reduce bird and fish predation (Haskell et al. 1996, Janzen et al. 2000a,b).

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LITERATURE CITED

- Bishop CA, Ng P, Pettit KE, Kennedy SW, Stegeman JJ, Norstrom RJ, Brooks RJ (1998) Environmental contamina-
tion and developmental abnormalities in eggs and hatchlings of the common snapping turtle (*Chelydra serpentina*) from the Great Lakes-St Lawrence River basin (1989–91). Environ Pollut 101:143–156


> Carr A (1964) Transoceanic migrations of the green turtle. Bioscience 14:49–52


Miranda-Filho KC, Metcalfe CD, Metcalfe TL, Muelbert MM and others (2009) Lactational transfer of PCBs and chlorinated pesticides in pups of southern elephant seals (*Mirounga leonina*) from Antarctica. Chemosphere 75:610–616

Pagano JJ, Rosenbaum PA, Roberts RN, Summer GM, Williamson LV (1999) Assessment of maternal contamina
tmental contaminant DDE fails to influence the outcome of sexual differentiation in the marine turtle Chelo
nia mydas. Environ Health Perspect 106:185–188
Pritchard PCH, Mortimer JA (1999) Taxonomy, external mor
phology, and species identification. In: Eckert KL, Bjorn
dal KA, Abreu-Grobois FA, Donnelly M (eds) Research and management techniques for the conservation of sea
turtles. IUCN/SSC Marine Turtle Specialist Group, Washing
ton, DC, p 21–40
gos green turtles Chelonia mydas in relation to oceano
graphic conditions: integrating satellite telemetry with remotely sensed ocean data. Endang Species Res 4:57–72

Thompson NP, Rankin PW, Johnston DW (1974) Polychlori
nated biphenyls and p,p’-DDE in green turtle eggs from Ascension Island, South Atlantic Ocean. Bull Environ Contam Toxicol 11:399–403
Van den Berg M, Birnbaum L, Bosveld ATC, Brunstrom B and others (1998) Toxic equivalency factors (TEFs) for PCBs, PCDDs, PCDFs for humans and wildlife. Environ Health Perspect 106:775–792
ging glaucous gulls (Larus hyperboreus) exposed to persist
ent organic pollutants. Auk 126:123–133
Willingham E, Crews D (1999) Sex reversal effects of environ
tmentally relevant xenobiotic concentrations on the re
dered slider turtle, a species with temperature-dependent sex determination. Gen Comp Endocrinol 113:429–435
Wyneken J, Salmon M (1992) Frenzy and post-frenzy swim
ning activity in loggerhead, green and leatherback hatch

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