Toxicological estimation of mortality of oceanic sea turtles oiled during the Deepwater Horizon oil spill

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ABSTRACT: Using multiple lines of evidence, we estimated the mortality of oceanic sea turtles that were minimally to moderately oiled by the 2010 BP Deepwater Horizon (DWH) oil spill in the Gulf of Mexico. Using estimates of the oil ingested by oceanic sea turtles and comparing them to toxic endpoints following oil ingestion in turtles and other vertebrate species, we derived an estimated percentage of mortality for oil-exposed oceanic sea turtles. Oil ingestion (mg kg⁻¹ BW d⁻¹) in oceanic sea turtles was estimated based on extent of oiling categories assigned by a sea turtle technical working group (STTWG) as follows: non-oiled (0), minimally oiled (1), lightly oiled (2), moderately oiled (3) and heavily oiled (4). Because the STTWG concluded that 100% of heavily oiled turtles would have died from the physical effects of heavy oiling, we limited our assessment of mortality to turtles in categories 0 to 3 and estimated how many of these turtles would have died from ingestion of oil. The estimated mortality was 85% for category 3, 50% for category 2 and 25% for category 1. Visibly non-oiled turtles (category 0) were assigned 0% mortality. To calculate the overall mortality for all turtles, the mortality estimations for categories 0 to 3 were applied to the numbers of turtles observed with different degrees of oiling, as documented by direct capture operations during the DWH spill. We concluded that, overall, approximately 30% of all oceanic turtles in the region affected by the DWH spill that were not heavily oiled would have died from ingestion of oil.

KEY WORDS: Toxicology · Mortality · Oceanic sea turtles · Deepwater Horizon · Oil spill · NRDA

INTRODUCTION

The explosion and sinking of the Deepwater Horizon (DWH) drilling platform in April 2010 caused one of the world’s largest marine oil spills, releasing millions of barrels of MC252 crude oil into the Gulf of Mexico (GoM) before the well was capped in July 2010 (McNutt et al. 2012). Various species of sea turtles occur in the GoM, and all are listed as endangered under the US Endangered Species Act of 1973 (Goodman Hall & Belskis 2012). During the response to the DWH oil spill, oceanic juvenile sea turtles were rescued offshore and sent to rehabilitation facilities for treatment and observation (Stacy 2012). However, the entire GoM was not surveyed, and the search effort for exposed turtles was limited to a period starting a few weeks after the spill (from May 17) to September 21, before the wellhead was capped (Stacy 2012). Therefore, the number of sea turtles dying in the wild during the spill or afterward is un...
known. Furthermore, there is limited knowledge on the sensitivity of turtles to oil (Vargo et al. 1986a,b,c, Bossart et al. 1995, Lutcavage et al. 1995) or the effects of oil spills on reptiles (Hall et al. 1983, Mignucci-Giannoni 1999, Wikelski et al. 2001, Saba & Spotila 2003, Camacho et al. 2013).

To investigate the impact of an oil spill, determining the exposure scenario including the magnitude and the duration of potential exposure is essential. Therefore, critical to our assessment was the data obtained from the sea turtles that were rescued offshore during the DWH incident. Rescued sea turtles were scored based on their extent of external oiling with 5 categories assigned as determined visually by photographs and observations in the field (Stacy 2012, DWH Trustees 2015). These categories represented non-oiled (category 0), minimally oiled (category 1), lightly oiled (category 2), moderately oiled (category 3) and heavily oiled (category 4) sea turtles. The volume of oil in their oral cavity was also visually estimated and was proportional to their external oiling level (Mitchelmore 2012a,b, Stacy 2012, DWH Trustees 2015, Mitchelmore et al. 2015a). These oral oil volume estimations were critical to the first step in our assessment to determine potential internal oil exposure levels. Levels of polycyclic aromatic hydrocarbon (PAH) metabolites in the bile and PAHs in tissues were measured in several field-collected oiled turtles and were positively correlated with the degree of external and internal oiling, demonstrating exposure, uptake and absorption of oil (Ylitalo et al. 2015, 2017 [this Theme Section]).

The objective of this work was to first estimate the potential exposure of oceanic sea turtles to oil and then to use this exposure data to estimate the mortality of oceanic sea turtles due to the DWH oil spill. The exposure of oceanic sea turtles to oil could only be estimated based on the limited information collected during the response phase of the spill, when rescue and rehabilitation of sea turtles was a priority, together with models of the potential exposure that occurred in wild sea turtles during the spill event (Stacy 2012, Stacy & Innis 2015, Stacy et al. 2017 [this Theme Section], Wallace et al. 2017 [this Theme Section], Ylitalo et al. 2017). These 2 sources of information, which provided some estimation of oil exposure, were then used, together with the literature on the toxicity of oil to reptiles and other vertebrates, to estimate mortality. This approach to estimating mortality was novel, and we are not aware of any other mortality estimate assessments in sea turtles that have employed a similar approach.

**METHODS**

Mortality was estimated only for oceanic sea turtles, which are defined as post-hatchling small juveniles that are essentially surface obligates and are associated with Sargassum and other floating materials in offshore locations (DWH Trustees 2015). For this assessment, it was assumed that all oceanic sea turtles, regardless of species, were equally sensitive to oil exposure during the spill event. It was concluded by a sea turtle technical working group (STTWG) that 100% of heavily oiled sea turtles (i.e. category 4) during the DWH spill would have died but for their capture and rehabilitation, primarily due to the physical effects of being mired in heavy oil in high temperatures (Stacy 2012, DWH Trustees 2015, Wallace et al. 2017). Therefore, the mortality estimates we provide pertain only to oceanic sea turtles that were less than heavily oiled, those in the non-oiled (category 0) through moderately oiled (category 3) categories as assigned by Stacy (2012).

We utilized various sources of information to estimate exposure for oceanic sea turtles. This included data provided by the response team and veterinary assessments of captured and rehabilitated turtles during the DWH spill (Stacy 2012, Stacy & Innis 2015, DWH Trustees 2015, Mitchelmore et al. 2015a, Stacy et al. 2017), particularly the extent of external and internal oiling and, in a small subset of animals, the biliary concentrations of PAH metabolites in sea turtles oiled during the DWH spill (Ylitalo et al. 2015, 2017).

**Estimation of daily and 14 d exposures to oil in DWH-exposed oceanic sea turtles**

To estimate the potential exposure of turtles to DWH oil, we used multiple lines of evidence for estimating internal exposure and uptake of oil in sea turtles following the DWH oil spill. These included the degree of external oiling, an estimation of the extent of oil in the oral cavity and the percentage of oceanic turtles in different oiling categories (see Table 1 and Stacy 2012, DWH Trustees 2015). Detailed descriptions of the criteria used to assign turtles to the different external oiling categories can be found in Stacy (2012), Mitchelmore et al. (2015a) and DWH Trustees (2015). The percentage of turtles with oil in their oral cavity, together with the estimated volume of oil in their oral cavity, is proportional to the extent of external oiling (Table 1). Although our mortality estimations are limited to turtles in categories 0 to 3, cate-
Table 1. Relationship between the extent of external oiling categories, the percentage of turtles with oil in their oral cavity, the estimated oral volume of oil and the bile fluorescent aromatic compounds present as PHN equivalents. Ranges show minimum and maximum values with means and number of turtles (where applicable) in parentheses. ND: not determined. Data from Stacy (2012) and Ylitalo et al. (2015, 2017), and Mitchelmore et al. (2015).

| Extent of visual external oiling (category) | Turtles with oil in oral cavity (%) | Estimated volume of oil in oral cavity (ml) | PHN equivalents (ng g⁻¹ bile, wet weight) |
|-------------------------------------------|-------------------------------------|---------------------------------------------|----------------------------------------|
| 0: non-oiled                              | ND                                  | ND                                          | 5400–30,000 (15 585; n = 20)            |
| 1: minimally oiled                        | 49.2                                | 0.05–0.15 (0.1)                             | ND                                     |
| 2: lightly oiled                          | 76.3                                | 0.25–0.50 (0.38)                            | 333 000–1 466 000 (11 000; n = 1)       |
| 3: moderately oiled                       | 93.2                                | 1.0–3.0 (2.0)                               | ND                                     |
| 4: heavily oiled                          | 96.6                                | 4.0–8.0 (6.0)                               | 110 000–510 000 (370 000; n = 4)        |

The model results are presented in Tables S1 to S3 (in the Supplement at www.int-res.com/articles suppl/n033 p039_supp.pdf) and show the estimated oral oil doses based on the minimum (0.34 kg), mean (1.42 kg) and maximum (4.00 kg) turtle BWs. Table 2 presents the summary of these calculations using the mean turtle BW. For all calculations, a correction factor for oil mass of 1.2 ml oil g⁻¹ (based on an oil specific gravity of 0.833) was used to convert the volume of oil to mass of oil so that the dose data are reported in standard units, i.e. mg oil kg⁻¹ BW turtle⁻¹ d⁻¹. Furthermore, it was assumed that the volume estimated in the mouth and esophagus (oral cavity) by Stacy (2012) was a total daily dose. So, to compare with the other vertebrate daily doses (calculated in the next subsection), estimates of daily oil doses for the 2 lowest oiling categories (using the mean turtle BW) using the minimum, mean and maximum oral estimates would range from 29 to 88 and from 146 to 293 mg oil kg⁻¹ BW turtle⁻¹ d⁻¹ for categories 1 and 2, respectively (see Table 2).

Exposure data were presented both as a daily instantaneous dose estimate and as a combined 14 d estimate (see Tables S4 & S5 in the Supplement). A daily dose was calculated to allow comparison with literature reporting daily oil doses in other species (see ‘Results and discussion’). Although many potential exposure scenarios are possible for turtles during the approximately 3 mo DWH spill time period, a 14 d exposure was chosen to represent one potential and likely exposure regime during the oil release period. A 14 d exposure regime was also used in a recent DWH oil exposure study in 2 species of freshwater turtles (Mitchelmore et al. 2015b). Therefore, these 14 d exposure calculations are provided in the Supplement as an example of an exposure scenario greater than the daily dose estimate. However, our toxicological comparisons to other published studies in this report are primarily based on the daily dose estimates in Table 2, given the limited amount of data available for comparison to other species’ exposure scenarios. Furthermore, it is unknown exactly how long a single ingested exposure, coating the esophagus, would expose a sea turtle to oil. We provide the 14 d cumulative oil exposure calculation (Table S4 in the Supplement) to highlight the complexity of multiple-day exposure scenarios, which would include a consideration of accumulation and depura-
tion of oil. For example, Table S4 in the Supplement calculates the 14 d retained dose based on repeated daily doses, with excretion occurring after 9 d. The 9 d oil passage time in turtles was estimated by Vargo et al. (1986b), who found excretion occurring after 9 d in juvenile sea turtles that ingested crude oil.

Calculations of daily oil ingestion rates in other vertebrate species

Based on literature searches in Google Scholar, Scopus and Web of Science, few studies have quantified oil exposure and impacts in turtles and other reptiles; therefore, we also considered oil exposure studies in non-reptilian vertebrate species when assessing potential toxicity to sea turtles. This included various bird species (e.g. Lattin & Romero 2014, Lattin et al. 2014) and mink (e.g. Mohr et al. 2008, 2010), as discussed in this section. We compared the estimated exposure in sea turtles to the levels and effects reported in 2 oil dose–response studies in mink (Mohr et al. 2008, 2010). To compare these mink studies to DWH-exposed sea turtles, calculations of daily doses for the mink studies were necessary. In the first study, male mink were fed 48, 520 and 908 ppm of Bunker C fuel oil or mineral oil (control) in their diet for 60 to 62 d (Mohr et al. 2010).

Provided the BWs and ingestion rates of the mink are known or can be estimated, then the dietary concentrations can be converted to a daily dose in units of mg (test agent) kg⁻¹ BW d⁻¹ (USEPA 1993). For mink, conversions of oil concentrations in the diet to a daily dose were based on a normalized ingestion rate of 0.15 kg of food kg⁻¹ BW d⁻¹ (based on assumptions of a food consumption rate of 0.15 kg d⁻¹ and a BW of 1.0 kg; USEPA 1995). BWs of 8 mo old mink range from 1000 to 1500 g (Wood et al. 1965). Therefore, at 48 ppm, oil in the mink diet would represent a daily dose of 7.2 mg oil kg⁻¹ (i.e. a mink ingesting 0.15 kg of a 48 mg oil kg⁻¹ food diet). The highest concentration in the 2 studies was 908 ppm, which equals a daily dose of 136 mg kg⁻¹ d⁻¹. Other doses used in the study included a diet of 420 and 520 ppm, equal to 63 and 78 mg kg⁻¹ d⁻¹, respectively. Therefore, daily doses using a 1 kg BW mink would be 7.2, 136, 63 and 78 mg oil kg⁻¹ BW d⁻¹ for the studies described above.

We also used recent studies in house sparrows exposed to oil in their diet for 2 to 4 wk as an additional daily dose comparison, given that birds are a closer taxonomic comparison than a mammalian species. In studies by Lattin & Romero (2014) and Lattin et al. (2014), house sparrows (average weight ~25 g) were fed a diet of 1% oil in their food (i.e. a 10 000 mg oil kg⁻¹ food dose) for 2 to 4 wk. This oil was a weathered GoM Louisiana sweet crude oil, which is more similar to DWH oil than the Bunker C used in the mink studies. Wang & Newman (2013) showed that the food ingestion rate in a similar species, the eastern song sparrow, which is similar in weight to the house sparrow, was estimated to be between 0.1 and 0.4 g food g⁻¹ BW and probably was likely to be between 0.1 and 0.15 g food g⁻¹ BW. Given these estimations, we calculate that the daily ingested dose for these birds would be around 1000 mg (up to 4000) oil kg⁻¹ BW d⁻¹.

Furthermore, we compare sea turtle exposure to a recent 14 d dietary oil exposure in 2 species of freshwater turtles that used daily exposures of 100 and 1000 mg kg⁻¹ BW d⁻¹ for a high- and low-dose exposure, respectively (Mitchelmore et al. 2015b).

RESULTS AND DISCUSSION

Known toxicological effects of oil exposure in turtles and other vertebrate species

There are very few studies addressing the toxicological effects of oil and/or chemical dispersants on
sea turtles (or any turtle species; for a review, see Shigenaka 2003). In the 1980s, a waterborne oil exposure study in loggerhead Caretta caretta and green Chelonia mydas sea turtles examined lethal and sublethal endpoints (Vargo et al. 1986a,b,c, Lutcavage et al. 1995). Juvenile loggerhead and green sea turtles were exposed for 96 h to a simulated oil spill (i.e. waterborne exposure) using weathered South Louisiana crude oil (Vargo et al. 1986b). The turtles ingested the oil during the exposure period and, thereafter, were evaluated up to 2 mo post-exposure. No excretion of oil by the turtles was observed until 9 d post-exposure, a finding used in the 14 d oil exposure estimates (Tables S1–S5 in the Supplement). Physiological assessments found some aspects of blood chemistry and composition were impacted by oil exposure. Oil was observed on the nares and eyes and in the upper portion of the esophagus, as well as in the feces. This demonstrated that external oil exposure of sea turtles can result in coating of epidermal surfaces, excretion in the feces after 9 d and alterations in hematologic and blood chemistry (Vargo et al. 1986b, Lutcavage et al. 1995).

Juvenile loggerhead turtles exposed to South Louisiana crude oil floating on top of the water in their tanks for 96 h at both low exposure (0.5 mm thick oil layer) and high exposure (5.0 mm thick oil layer) exhibited marked and consistent histologic skin changes in both exposure groups, and these effects were more severe in the high-exposure group (Vargo et al. 1986b, Bossart et al. 1995). Skin and exposed mucous membranes began to slough in irregular sheets, beginning after approx. 9 d and continuing to 21 d, but appeared grossly normal by 40 d post-exposure. The potential for the skin dysfunction presented a potential portal of entry by pathogenic organisms, and the authors concluded that the turtles would face substantial risks in and after an oil spill (Bossart et al. 1995).

A recent laboratory study orally exposed juveniles of 2 freshwater turtle species (red-eared slider Trachemys scripta elegans and common snapping turtle Chelydra serpentina serpentina) as surrogates for sea turtles to ‘Slick A’ weathered MC252 oil for 14 d (DWH Trustees 2015, Mitchelmore et al. 2015b). Two oil doses were used to represent minimally to moderately oiled turtles (i.e. 100 and 1000 mg oil kg\(^{-1}\) BW d\(^{-1}\), respectively) based on the oil ingestion estimates and subsequent daily dose estimates detailed in Mitchelmore (2012a,b) and Mitchelmore et al. (2015a). Dose-dependent increases in biliary FACs were observed, but there was no mortality or hemolytic anemia observed after 14 d of exposure (DWH Trustees 2015, Mitchelmore et al. 2015b).

Some oil-dependent physiological abnormalities were seen, including evidence of dehydration, decreasing digestive function and assimilation of nutrients (DWH Trustees 2015, Mitchelmore et al. 2015b). Some other oil-dependent alterations in common PAH biomarker endpoints including oxidative stress and DNA damage were observed (DWH Trustees 2015, Mitchelmore et al. 2015b). Two red-eared sliders that had received a high dose of DHW Slick A oil for 14 d showed dysfunction of the hypothalamic–pituitary–adrenal (HPA) axis. Although there was a decreasing trend in median corticosterone rise following adrenocorticotropic hormone (ACTH) administration, this was not statistically significant, likely due to the low numbers of animals used for the experiment and the high variation in response observed between individuals coupled with only 14 d of exposure (DWH Trustees 2015, Mitchelmore et al. 2015b). In the snapping turtles, no statistically significant differences between control and treatment groups in baseline or ACTH-induced corticosterone levels were observed (DWH Trustees 2015, Mitchelmore et al. 2015b). Therefore, the effects of experimental DWH oil exposure on red-eared sliders and common snapping turtles did not consistently indicate HPA dysfunction, although given the limitations to this study and evidence of dysfunction in the red-eared sliders, impact to the HPA axis and the animals’ responses to future stress events could not be ruled out.

There are very few reports of the effects of oil spills on wild sea turtles. Vargo et al. (1986a) estimated that about 3.2% of sea turtle strandings reported to the Florida Department of Natural Resources were petroleum-related, and notably 37.7% were of unknown cause. Among the petroleum-affected sea turtles, the most often reported strandings were from southeastern Florida, and primarily green sea turtles, both adults and juveniles, were affected. Other brief reports describe occurrences of oiled sea turtles and rehabilitation and release efforts of sea turtles during oil spills in the central west coast of Florida and in the Persian Gulf (Anonymous 1991, 1994, Thomas & Robinson 1991). Computer models of the impact of oil spills on Atlantic loggerhead turtles Caretta caretta indicate that an oil spill accelerates population decline (Leung et al. 2012). Oil exposure has been implicated as the principle cause of strandings of juvenile sea turtles in the Canary Islands, Spain, between 1998 and 2011 (Camacho et al. 2013).

The blowout of the Ixtoc 1 oil well in Mexico’s Bay of Campeche in 1979 resulted in the daily release of
10,000 to 15,000 barrels of oil for a period of several months, with the oil drifting northward in the GoM. Green sea turtles and Kemp’s ridley sea turtles Lepidochelys kempii were found dead and had petroleum in their upper alimentary systems; there was no evidence that the oil had caused alimentary lesions and no sign of pulmonary aspiration (Hall et al. 1983).

During an oil spill in Puerto Rico in 1994, Mignucci-Giannoni (1999) found that rehabilitated sea turtles (Chelonia mydas and Eretmochelys imbricata), when compared with rehabilitated sea birds, had higher chances of survival after being exposed to an oil spill. Survival estimates were based on the relative health condition of released animals. But as noted by Saba & Spotila (2003), that does not infer that the sea turtles had no crude oil residuals present in their tissues following rehabilitation (Mignucci-Giannoni 1999). However, survival and post-rehabilitation behavior of 4 species of freshwater turtles exposed to a crude oil spill in February 2000 in a US National Wildlife Refuge indicates that rehabilitation of oil-exposed freshwater turtles can be effective in restoring these animals to normal behavior, based on home range sizes and water temperature preferences (Saba & Spotila 2003). Modeling from oil exposure to impact in turtles is not entirely novel. For example, population mortality estimates for freshwater diamondback terrapins Malaclemys terrapin were assigned by Michel et al. (2001) following an oil spill in Maryland. However, these estimates were based only on the oiling of the shoreline of the Patuxent River, as no measured levels or estimates of internal exposure were available for the turtles, thereby limiting comparisons to DWH oil-exposed turtles and laboratory oil exposure studies in turtles and other vertebrate species discussed in this subsection (Michel et al. 2001). In the weeks following the Patuxent spill, the extent and degree of shoreline oiling were defined, and mortality estimates associated with different levels of shoreline oiling were as follows: (1) heavily oiled, with 10% mortality; (2) moderately oiled, with 2% mortality; and (3) lightly oiled, with 0.5% mortality within the subpopulations of the turtles that lived in those areas (Michel et al. 2001). This risk assessment approach was similar to ours except that we were able to base our estimates on semi-quantitative observations of external and ingested oil of sea turtles, allowing us to compare findings with laboratory oil exposures in turtles and other vertebrate species.

Mortality estimates for DWH MC252-exposed sea turtles were made based on the variety of clinical and medical parameters assessed (blood chemistry and hematological endpoints) during the rehabilitation of captured sea turtles (Stacy & Innis 2015, Stacy et al. 2017). Using a series of models that have previously been used in turtles (e.g. see Stacy et al. 2013), an average of 14% (range 4 to 22%) mortality for all of the turtles irrespective of level of oiling would have occurred had they not been rescued (Stacy & Innis 2015). This was deemed to be a conservative measure, as it does not take into account long-term, sub-lethal exposures or delayed effects of oil. In a follow-up study, focusing only on Kemps ridley turtles, a different approach was used which resulted in an estimate of 25% mortality for this species (Stacy et al. 2017).

Given the paucity of studies of the effects of oil exposure in sea turtles, literature for other marine and freshwater reptiles, mammals and birds was also considered when deriving our mortality estimates. In 2001, an oil spill occurred in the Galapagos Islands, Ecuador, releasing diesel and bunker oil throughout the archipelago of San Cristóbal Island in the Galápagos National World Heritage Site. Marine iguanas Amblyrhynchus cristatus had been studied on that site since 1981, and a few animals died immediately (Wikelski et al. 2001). Of the 170 individuals examined, 70% had oil residue on their skin (Wikelski et al. 2001). Plasma levels of corticosterone were highly elevated in animals sampled after the oil spill compared to animals sampled 10 d prior to the spill. This was true for baseline (within 3 min of capture) and stress-induced levels (stress was a manual restraint of the animals). Corticosterone levels were indistinguishable between animals that had externally visible oil blotches and those without, suggesting that animals in the vicinity of the oil spill were affected even without visible oil exposure. Corticosterone levels in the marine iguana population were linearly related to survival rates (Wikelski et al. 2001).

In contrast, adrenal insufficiency following oil exposure, albeit in longer-term exposures, has been demonstrated in other vertebrate species. For example, within the footprint of the DWH oil spill, bottlenose dolphins Tursiops truncatus exhibited adrenal insufficiency and anemia, low blood-sugar levels and symptoms of liver and lung disease associated with mortality (Schwacke et al. 2014). Because the HPA response to stress appears to be conserved across vertebrate species, and because there is little information on how oil exposure affects this critical pathway (Moore & Jessop 2003), we also considered studies on birds and mammals.
exposed orally to oil as potential indicators of toxicity in sea turtles. In a laboratory study of oil-exposed mink, effects on blood chemistry, hematology and cytochrome P450 were observed, in addition to adrenal hypertrophy (Mazet et al. 2000, Beckett et al. 2002, Schwartz et al. 2004, Mohr et al. 2008). Also in mink, dietary exposure to Bunker C fuel oil for 60 to 62 d resulted in adrenal hypertrophy in males at all doses tested, including the lowest dose of 48 ppm and up to the highest dose of 908 ppm (i.e. 7.2 and 136 mg oil kg\(^{-1}\) BW d\(^{-1}\), respectively; Mohr et al. 2008). However, those chronic oral exposures to low concentrations of Bunker C oil did not alter serum or fecal glucocorticoid concentrations. In a second related study, mink were fed artificially weathered fuel oil in their diet for 60 to 62 d, and adrenal hypertrophy was also found at all doses and in both sexes (Mohr et al. 2010).

Recent studies in house sparrows demonstrated that stress levels of corticosterone were reduced following chronic exposure to crude oil for 2 to 4 wk (i.e. oil doses ~1000 mg oil kg\(^{-1}\) BW d\(^{-1}\); Lattin et al. 2014). Ingestion of crude oil also showed a decrease in stress-induced corticosterone levels in mallard ducks (Gorsline & Holmes 1982). In a similar study, levels of glucocorticoid receptors were decreased in the liver of the oil-exposed birds relative to controls, highlighting that oil may impact the HPA axis at multiple levels and through multiple physiological mechanisms (Lattin & Romero 2014).

The 14 d oil exposure laboratory study in 2 species of freshwater turtles using oil doses representative of the low to moderate exposures in DWH-oiled turtles (i.e. 100–1000 mg oil kg\(^{-1}\) BW d\(^{-1}\)) also provided some evidence of adrenal dysfunction, including decreased levels of corticosterone following a stress response (discussed earlier in this subsection; see Mitchelmore et al. 2015b). However, conclusions were limited, given the limited number of animals, the large variability in response and the shorter exposure duration period compared with the mink and bird studies.

The results of this literature review confirmed that there is evidence, in reptiles and other vertebrates, that exposure to oil (including chronic low doses and continued impacts post-exposure) adversely affects the adrenal glands and/or their function. It has been suggested that chronic low-dose ingestion of oil can affect not only stress-induced corticosterone levels but also glucocorticoid receptor density, demonstrating that oil can act at multiple levels to disrupt responses in organisms and their ability to survive stressors, including extreme events or predator attack (Lattin et al. 2014). Indeed, an overall general suppression of the stress response has been put forth to at least partially explain why oil-exposed animals show elevated mortality after exposure to subsequent stressors (see Holmes et al. 1979).

**Description of the approaches used in defining mortality estimates in oceanic sea turtles exposed to DWH oil**

Our estimates of mortality in less than heavily oiled sea turtles (categories 0–3) are based on the limited information available for oil-related impacts in turtles coupled with evidence derived from other vertebrate species (mink and birds). Based on the estimated daily ingested doses of oil presented in this report, turtles during the DWH spill would be exposed to concentrations of oil that are associated with toxicological effects in turtles and other vertebrate species. For example, using our calculated estimated daily doses of oil (Table 2 and Tables S1–S5 in the Supplement), the exposures estimated for oceanic sea turtles during the DWH oil spill are equal to or substantially higher than a known dose in mink of 7.2 mg kg\(^{-1}\) d\(^{-1}\), that specifically impacts the HPA axis (Mohr et al. 2008, 2010) for all exposure scenarios. However, it should be noted that the mink experiments were conducted with Bunker C fuel oil and not the DWH MC252 oil. House sparrows exposed to oil for 2 to 6 wk also showed negative impacts to the HPA axis (i.e. reduced stress corticosterone levels and reduced or unchanged [not up-regulated as expected] glucocorticoid receptor levels) using an oil similar to the DWH oil at daily dose levels equivalent (i.e. 1000 mg oil kg\(^{-1}\) d\(^{-1}\)) to the moderately exposed sea turtles (see Table 2).

Although it is difficult to compare across different species (species sensitivity issues), oils, doses and exposure periods, based on our daily estimates, it appears that even minimally oiled turtles (category 1) would be at risk for adrenal insufficiency, which would have consequences for long-term survival and fitness of the sea turtle population. This finding is substantiated by the findings of the literature review, including the marine iguana study that indicates that reptiles exposed to oil spills will respond through the HPA axis and that elevated corticosterone responses can be indicators and correlates of reduced survival (Holmes et al. 1979, Wikelski et al. 2001). Hypo- or hyper-adrenal response has been identified as an effect of concern across taxa, including marine igua-
nas, birds and mammals (Gorsline & Holmes 1982, Wikelski et al. 2001, Mohr et al. 2008, 2010, Lattin & Romero 2014, Lattin et al. 2014, Schwacke et al. 2014).

From the evidence in the literature and our modeling, we concluded that sea turtles exposed to the DWH oil spill would likely experience adrenal dysfunction leading to mortality. Adrenal insufficiency can impact turtles in several ways that can reduce population viability through increased mortality and reduced reproduction. For turtles in oiling categories 2 and 3, the estimated daily ingested dose of oil greatly exceeds the dose that would be expected to cause adrenal insufficiency. Turtles in category 4 would also be in this group, but turtles in this group were already considered to have a mortality of 100% and so are not considered further here in our mortality estimates. For turtles in category 3, where 93% were observed to have oil in their mouths, a daily dose of oil of 208 to 7353 mg kg\(^{-1}\) d\(^{-1}\) was estimated, which is orders of magnitude greater than the known dose which specifically impacts the HPA axis in mink of 7.2 mg kg\(^{-1}\) d\(^{-1}\). Therefore, we initially estimated that all turtles that ingested this amount of oil would have died. However, given the potential for a shorter acute exposure period in sea turtles after the DWH oil spill compared with the mink studies, we concluded that there would have been a minimum of 90% mortality in category 3 oiled turtles (range of 90–100%; Table 3). Given that this mortality is only applicable for turtles that have ingested oil (i.e. 93%), we estimated an overall mortality for category 3 turtles of 85% (90% of 93%; mortality estimates are rounded to the nearest 5%; Table 3).

Using similar reasoning for category 2 turtles, where 76% had oil in their mouths at high daily dose levels (52–1225 mg kg\(^{-1}\) BW d\(^{-1}\)), and given the same caveat about exposure time, we determined a range of 65 to 80% mortality in category 2 oiled turtles. Using the lowest estimate of 65% mortality for only the 76% of turtles that had ingested oil, 50% mortality (i.e. 65% of 76%; Table 3) was estimated for category 2. For turtles in category 1, with visible oil in their oral cavity in 49% of cases, where the minimum estimated 14 d dose (10 mg kg\(^{-1}\) d\(^{-1}\); range 10–36 mg kg\(^{-1}\) d\(^{-1}\)) was similar to the lowest exposure in mink that induced adrenal insufficiency, and again given the possibility that exposure duration may be lower in turtles compared to the 62 d mink study, a 50% mortality estimate was assigned.

### Table 3. Summary of mortality estimates calculated for DWH-exposed oceanic turtles. STTWG: sea turtle technical working group; HPA: hypothalamic–pituitary–adrenal

| Oiling category | Min. mortality estimate (%) | Description and rationale used in ascribing mortality value |
|-----------------|-----------------------------|----------------------------------------------------------|
| 4               | 100\(^a\)                   | The STTWG determined that 100% of heavily oiled turtles would have died but for their capture and rehabilitation, primarily due to the physical effects of being mired in heavy oil in high temperatures. Therefore, this report does not assess mortality in this category, although it does model potential exposure in this group. |
| 3               | 85                          | This mortality estimate is calculated using our estimated daily oil doses for 14 d that far exceed the dose of oil shown to cause adrenal insufficiency in other vertebrate species. Based on the impact to the HPA axis alone, 90 to 100% mortality is expected in the 93% of turtles that had ingested oil. However, given the potential for reduced exposure time to reduce impacts, a minimum estimate of 85% mortality (90% of 93%) was concluded. |
| 2               | 50                          | The rationale for the estimates for this category is as described for category 3, except mortality via HPA axis impact was reduced to a range from 65 to 80%; therefore, the minimum 65% estimate was used in these analyses. Using the lowest estimate of 65% mortality for only the 76% of turtles that had ingested oil, 50% mortality (i.e. 65% of 76%) was estimated. |
| 1               | 25                          | The rationale for the estimates for this category is the same as that described for category 3, except mortality via HPA axis impact ranges from 50 to 65%; using the minimum 50% mortality estimate applied to the 50% of turtles in this category that had oil in their oral cavity, an overall minimum mortality estimate of 25% (i.e. 50% of 50%) was assigned to this group. |
| 0               | 0                           | No oil exposure externally or internally visible.\(^b\) |

\(^a\)Previously estimated by the STTWG (e.g. Stacy 2012, DWH Trustees 2015)

\(^b\)Does not necessarily mean that these turtles were not exposed to and not impacted by oil, given non-visible chronic oiling and other potential exposure routes.
mortality (range 50–65%) was estimated. Therefore, the mortality estimate for only the turtles that had ingested oil is 25% for category 1 turtles (50% of 50% that ingested oil; Table 3).

Mortality was assumed to be zero for sea turtles that had no visible oil exposure externally or internally (Table 3). However, it is quite possible that these turtles may have been exposed (or could be delayed or sublethal effects from physical impairment (Fig. 2) other than adrenal dysfunction were not considered, e.g. heat stress, reduced fitness and implications for effects on behavior affecting foraging and predator avoidance. Impacts resulting from exposure to chemical dispersants or chemically dispersed oil represent an additional unknown toxicity concern.

![Fig. 1. Potential routes of oil exposure to oceanic sea turtles. Redrawn from Mitchelmore (2012a). GI: gastrointestinal](image)

![Fig. 2. Potential direct and indirect effects of oil (including toxicological endpoints) on turtles. Redrawn from Mitchelmore (2012a)](image)
Calculation of mortality of sea turtles in categories 0 to 3

To calculate the number of dead sea turtles in less than heavily oiled turtles (i.e. categories 0 to 3), our percentage estimations were applied to the known numbers of turtles documented by direct capture operations during the DWH spill (Stacy 2012, DWH Trustees 2015, Wallace et al. 2017 and Table 4) that were assigned to each of the oiling categories. This was calculated as 25% of 266 turtles in category 1 (i.e. 66.5), 50% of 87 turtles in category 2 (i.e. 43.5) and 85% of 47 turtles in category 3 (i.e. 40) = 150 dead turtles, which is approximately 30% of the 510 turtles examined (Table 4). Therefore, we project an overall mortality of 30% for all oceanic turtles within the footprint of the DWH oil spill in addition to those already presumed to have died from heavy oiling (i.e. category 4). This mortality estimate has been used by other researchers to estimate the total number of turtles that may have died in the DWH footprint (e.g. DWH Trustees 2015, McDonald et al. 2017 [this Theme Section]).

CONCLUSIONS AND SUMMARY

Using field observations of oil ingestion by sea turtles following the DWH oil spill, our oil exposure estimates, a review of existing scientific literature and results from a laboratory study on a surrogate turtle species, we concluded that there is strong potential for substantial biological perturbation, reduced fitness and subsequent mortality in any sea turtle exposed to oil from the DWH oil spill. The mortality estimates provided here are minimum estimates, especially considering the potential for additional effects from other exposure routes and the currently unknown potential for delayed effects that may manifest at individual and population levels for years to come.

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