

# **Considerations for Development of Entrained Oil Thresholds for Oil Spill Risk Assessments**

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## **Summary**

Oil spill modelling using full oil fate models such as RPS' SIMAP model provides output for two types of water column concentration data: (1) dissolved hydrocarbons and (2) total hydrocarbons and related compounds in petroleum oils (abbreviated as THC) in entrained oil droplets. The purpose of this technical note is to identify appropriate thresholds for use in oil spill risk assessments. The focus is on developing a THC threshold appropriate for modelled entrained oil droplets.

As described in detail herein, toxicity is related to the bioavailability of hydrocarbons and related compounds in petroleum oils (described as “hydrocarbons” herein for brevity). Soluble and semi-soluble components of oil dissolve and therefore become bioavailable (i.e., able to be taken up by aquatic organisms). In relatively fresh oil, some of the hydrocarbons in entrained oil droplets are soluble/semi-soluble that may later dissolve and become bioavailable. However, as oil weathers, these potentially toxic components diminish to the point where the THC in entrained droplets is effectively non-toxic. Therefore, the focus of a risk assessment is typically on the dissolved hydrocarbon exposure.

Measurement of the concentration of oil in water, whether in the field or in laboratory studies, is complicated and not all constituents can be identified and measured individually. Mass recoveries (using the presently recommended methods) for gasoline, kerosene, and gas oil are typically >95%, whereas the mass recoveries for heavier substances in crude oils and heavier refined products are often <30%. Because only some of the compounds in oil are measurable individually, and the desire to be inclusive of effects from hydrocarbons in oil droplets as well as the dissolved phase, aquatic toxicologists have attempted to use THC as a metric for evaluating toxicity of oil exposures. However, given the variable composition of oil as it weathers, development of effects levels or thresholds based on total hydrocarbons (THC) is problematic. The review series (Loughery et al. 2023; Stubblefield et al. 2023; Alloy et al. 2023; Parkerton et al. 2023a,b; Dettman et al. 2023; Bejarano et al. 2023; Lee et al. 2023; French-McCay et al. 2023) provides guidance for aquatic toxicity studies of petroleum oils and their application to risk assessments and modelling-based analyses.

For the dissolved hydrocarbons, thresholds based on acute lethality (LC50s, i.e., lethal concentrations for 50% of test organisms) generally range from about 10 ppb ( $\mu\text{g/L}$ ) for sensitive early life history stages of aquatic species to 300 ppb, or more for less sensitive species and older

life stages. For sub-lethal effects, a threshold (PNEC) of 1 ppb is conservatively protective of aquatic biota.

If THC in entrained oil droplets is to be evaluated as a risk or to define the Environment that May Be Affected (EMBA), 1 ppm (1 mg/L; 1,000 ppb) would be a sufficiently conservative threshold for oil droplets of all oil types and all weathering states, with 3 ppm (3 mg/L; 3,000 ppb) an appropriate threshold for crude oils. For light distillates and condensates, 1 mg/L (1 ppm; 1,000 ppb) is an appropriate threshold.

## Introduction

An objective of oil spill risk assessments is to evaluate whether aquatic habitats and organisms could be exposed to oil or its constituents at concentrations above levels of concern at some time after a spill. Typically, oil transport and fate modelling (e.g., using RPS' Spill Impact Model Application Package [SIMAP], French-McCay 2003, 2004, French-McCay et al. 2021)) is used to estimate over space and time: 1) the spread and concentrations of oil on the water surface, 2) concentrations of oil (i.e., entrained whole oil droplets) in the water column, 3) dissolved concentrations in the water column, and 4) standing of oil on shorelines. Floating and shoreline oil thresholds have been based on observed effects after spills. Water column concentrations have been compared to thresholds based on laboratory toxicity studies performed under controlled conditions designed to provide continuous exposures (e.g., typically without dilution and water column mixing) at sufficiently elevated concentrations to induce a measurable response.

In Australia, recent practice, based on government guidance, has been to identify all areas exceeding conservative threshold concentrations at any instant in time after a spill as the Environment that May Be Affected (EMBA). The proponent is then required to develop a scientific monitoring program for the EMBA's extent, as well as consult with stakeholders regarding potential impacts or concerns.

In determining the EMBA, consideration is to be given to four oil pathways: floating oil, entrained oil droplets, dissolved aromatic hydrocarbons, and stranded oil (on shorelines). The "low" (i.e., conservative) thresholds recommended by NOPSEMA (Environment Bulletin, April 2019) are:

- Floating oil as THC:  $1 \text{ g/m}^2 = 1 \text{ m}^3/\text{km}^2$
- Dissolved aromatic hydrocarbons: 10 ppb
- Entrained oil as THC: 10 ppb
- Shoreline oil as THC:  $10 \text{ g/m}^2 = 10 \text{ mL/m}^2$

The floating and shoreline oil thresholds were based on the literature reviews and analyses described in French McCay (2016) and French McCay et al. (2018). Based on the Bonn Agreement Oil Appearance Code (BAOAC, Bonn Agreement, 2009, 2011), the floating oil threshold is the typical thickness of rainbow sheen. Potential for effects on socioeconomic resources may occur (e.g., fishing may be prohibited) if oil is visible on the water surface, i.e., sheens. Effects on socioeconomic resource uses may occur (e.g., recreation, tourism) above the

shoreline threshold, such that this level of contamination may trigger the need for shoreline clean-up on amenity beaches. Thus, these are reasonable thresholds for establishing an EMBA.

NOPSEMA (2019) indicated that the dissolved and entrained oil thresholds are based on Bridges et al. (2018), French McCay (2016) and French McCay et al. (2018). While the dissolved threshold of 10 ppb is consistent with analyses in French McCay (2016) and French McCay et al. (2018), those publications did not advocate for 10 ppb as an entrained oil threshold. Entrained oil is whole oil mixed into the water column, and French McCay (2016) argued that since the dissolved concentrations derive from compounds that typically represent about 1% of the oil when fresh, an entrained oil threshold should be at least 100 times that for dissolved concentrations. Bridges et al. (2018) developed toxic effects levels based on total polycyclic aromatic hydrocarbons [PAHs] for 50 measurable PAHs), which represented just 2% of the parent (MC252) oil. Crude oils typically contain 1-3%, and condensates contain up to 7% of these measurable PAHs. While the entrained oil threshold was said to be based on potential for exceedance of water quality triggers (NOPSEMA, 2019), it is unclear what the precise basis of the 10-ppb value would have been.

The EMBA is to consider any probability (i.e., even one trajectory out of hundreds of potential pathways) of exceeding the low threshold for any of the four oil pathways at any instant in time (determined by the model time step, assumed 1 hour). Consequently, recent EMBA's have encompassed large areas of the oceans surrounding Australia, owing to use of these conservative thresholds developed without consideration of the dynamic nature of an oil spill (i.e., dilution and dispersion in 3-dimensions and time), duration of exposure, the degree of weathering of the oil, the relative toxicity of different contaminants in oil, and the portion of the water column potentially exposed. An unintended consequence of using conservative thresholds is that resources for assessing impacts (e.g., water sampling, monitoring) would be spread across a large area, potentially diminishing the effectiveness of these efforts.

Of particular concern is the low threshold (10 ppb) for entrained oil recommended by NOPSEMA. The use of this threshold does not consider the bioavailability of constituents in the oil that drive aquatic toxicity, the changing composition of oil as it weathers, or the likelihood and duration of exposure. Oil fate and exposure modelling addresses the complexities of oil composition, weathering, partitioning in the environment, and organisms' behaviors. In view of these points, more appropriate entrained oil thresholds and considerations could be used for risk evaluations and establishing an EMBA, as described below.

## **Bioavailability**

### **Dissolved Compounds in Oil are Bioavailable**

Dissolved compounds are taken up into organisms directly through external surfaces and gills, as well as through the digestive tract. Thus, soluble and semi-soluble hydrocarbons and related compounds in oil are bioavailable, whereas insoluble compounds in oil are not bioavailable to aquatic organisms. Laboratory studies have shown that the dissolved hydrocarbons exert the most effects on aquatic biota (Carls et al. 2008; Nordtug et al. 2011; Redman 2017a; Gobas et al. 2018).

The fraction of oil with the most toxic effects to water-column and benthic organisms are lower-molecular-weight compounds (i.e., PAHs), which are volatile, and soluble or semi-soluble in water. As a result, they persist in the environment long enough for significant exposure to occur (Anderson et al., 1974, 1987; Neff and Anderson, 1981; Malins and Hodgins, 1981; McAuliffe, 1987; NRC 2003, 2005; Nordborg et al. 2018). The monoaromatic hydrocarbons (MAHs), including BTEX (benzene, toluene, ethylbenzene, and xylenes), and the soluble alkanes also contribute to toxicity, but these compounds are highly volatile, so exposures of aquatic biota are minimal or negligible except when light oils are discharged at depth where volatilization does not occur (French-McCay 2002).

Within the soluble and semi-soluble hydrocarbons, toxicity is inversely related to solubility, typically quantified by the octanol-water partition coefficient ( $K_{ow}$ ), a measure of hydrophobicity (Nirmalakhandan and Speece 1988; Hodson et al. 1988; Blum and Speece 1990; McCarty 1986; McCarty et al. 1992a, b; Mackay et al. 1992; McCarty and Mackay 1993; Verhaar et al. 1992, 1999; Swartz et al. 1995; French-McCay 2002; McGrath et al 2009). The range of LC50s varies from ~10 µg/L (ppb) for 3-ring PAHs (which are semi-soluble) to ~10-100 mg/L (ppm) for the highly soluble BTEX compounds (French-McCay 2002). Thus, the toxicity of an oil hydrocarbon mixture is strongly related to the chemical composition, which varies as the oil weathers since the soluble and semi-soluble hydrocarbons are all volatile to varying degree.

### **Effects of Oil Droplets**

While entrained oil droplets could potentially affect organisms by mechanical means (i.e., fouling, interference with feeding), research has not definitively demonstrated direct effects of droplets as separable from the effects of components dissolved from the oil (Parkerton et al. 2023b). Contact of droplets with organisms' surfaces (external or by ingestion) could facilitate direct exposure to hydrocarbons if they become bioavailable (i.e., dissolve; Letinski et al. 2014; Redman et al. 2017a).

### **Oil Composition Affects Toxicity**

Oil is a complex mixture of thousands of hydrocarbons and related compounds of varying physical, chemical, and toxicological characteristics, and therefore, varying environmental fates and effects on organisms. As discussed above, due to differing solubility, oil components vary by orders of magnitude in their potential to cause adverse effects, due to varying bioavailability. A fraction of oil does not readily biodegrade, because the compounds involved are not bioavailable. Oils differ in composition and their fractions of potentially bioavailable components, and so not all oils are equally toxic.

Thus, the use of a simple sum of components to quantify toxicity is problematic, as the composition of dissolved components to which organisms are exposed in the field will differ from that in the laboratory study used to develop the toxicity threshold. Toxic effects levels vary by what is included in the summation and the relative concentrations in the mixture.

The approach used to quantify the additive effects of a mixture of hydrocarbons and related compounds is to use a Toxic Unit (TU) model (Swartz et al. 1995; DiToro et al. 2000; French-McCay 2002; Redman et al. 2017b; NASEM 2020; Parkerton et al. 2023a; see review by French-McCay et al. 2023). A TU for evaluating mortality is defined as the exposure concentration divided by the lethal concentration affecting 50% of exposed organisms (LC50). Thus, at TU=1, mortality is expected. For a mixture, the toxic units are additive. When for a mixture of chemicals at specific concentrations  $\Sigma TU = 1$ , the mixture is predicted to be lethal to 50% of exposed organisms. Use of the TU approach for evaluating effects addresses the varying composition of oil-derived mixtures, including changes in composition due to weathering.

## **Oil Weathering**

As oil weathers, its composition changes due to volatilization and dissolution losses, and biodegradation. When oil is floating, the volatile components evaporate rapidly, and the oil becomes more viscous as a result. Some of the semi-soluble hydrocarbons, which are less volatile, dissolve from the floating oil into the water column. Floating oil may be entrained into the water column by breaking waves, or oil may be released under water. Soluble and semi-soluble hydrocarbons dissolve from subsurface oil droplets, weathering the oil and making them more available to microorganisms. The uptake of hydrocarbons by microorganisms, referred to as biodegradation, reduces water column concentrations, and therefore toxic effects. The volatilization rates of hydrocarbons from surface slicks are faster than the dissolution rates. Thus, dissolution from oil droplets in the water column is the main source of concentrations dissolved in the water.

Therefore, oil weathering leads to a preferential loss of the toxic components of the oil. The more bioavailable components evaporate and dissolve (and potentially cause acute toxicity in water column), leaving residual oil with lower potential for causing adverse effects. Entrained oil droplets weather rapidly because of their high surface area relative to their volume. Microbes attach to their surfaces and take up dissolved components readily. Highly weathered petroleum residuals have not been shown to have adverse effects on aquatic organisms, i.e., they are effectively non-toxic. Therefore, effects levels for the bioavailable soluble and semi-soluble components should not be applied to whole oil entrained droplets, particularly for weathered entrained oil droplets.

## **Consideration of Duration of Exposure**

### **Time for Uptake of Hydrocarbons**

An additional consideration is that LC50s and other toxicity endpoints vary considerably with exposure duration over the range of several hours to several days (Sprague, 1969; Abel, 1980; Mancini, 1983; Bailey et al., 1985; McAuliffe 1987; McCarty et al. 1992a, b). Effects thresholds are an order of magnitude higher for a few hours of exposure than they are for several days of exposure (McCarty et al. 1992a, b; French-McCay 2002; Bejarano et al. 2014) due to the hydrocarbon accumulation over time up to a critical tissue concentration that causes mortality.

Exponentially higher concentrations are required to achieve equivalent effects over shorter durations (e.g., Figures 1 and 2). The accumulation is slower for more hydrophobic (i.e., less soluble) compounds (Veith et al., 1983; Abernethy et al. 1986, 1988; Verhaar et al., 1992, 1999; DiToro et al. 2000; DiToro and McGrath 2000; see reviews in McCarty et al. 1992a, b; Verhaar et al., 1992; French-McCay 2002; French-McCay et al. 2023). Thus, the LC50s for PAHs are more variable with duration of exposure than are the LC50s for BTEX.

Most bioassays are performed under 48 to 96 hours of exposure to allow concentrations to accumulate in the tissues of the exposed organisms, where they may have adverse effects. Swartz et al. (1995) argued that bioassays for PAHs should be to at least 10 days of exposure in order to allow sufficient time for the PAHs to be taken up into the organisms and reach steady state between uptake and depuration. The focus on Swartz et al.'s (1995) study was PAH contamination in sediments. For long, essentially infinite exposure durations, the measured lethal endpoints are termed incipient LC50s (or other percentage mortality levels; McCarty et al. 1992a, b; Verhaar et al., 1992; French-McCay 2002). The >48-hour and incipient LC50s are conservatively low for shorter-duration exposures (French-McCay 2002). Animals can be unaffected by short-duration exposures to concentrations that could affect them if exposed for days or weeks. Organisms can recover from sublethal exposures, by elimination of toxins from tissues.

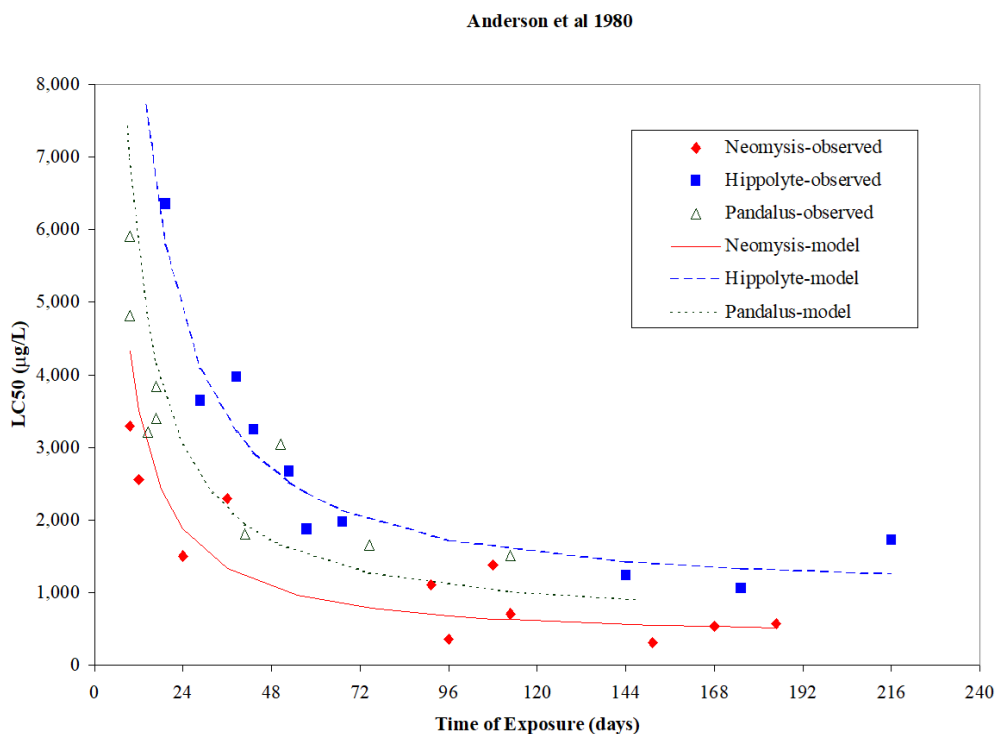


Figure 1. Lethal concentrations (LC50) versus duration of exposure in experiments reported by Anderson et al. (1980).

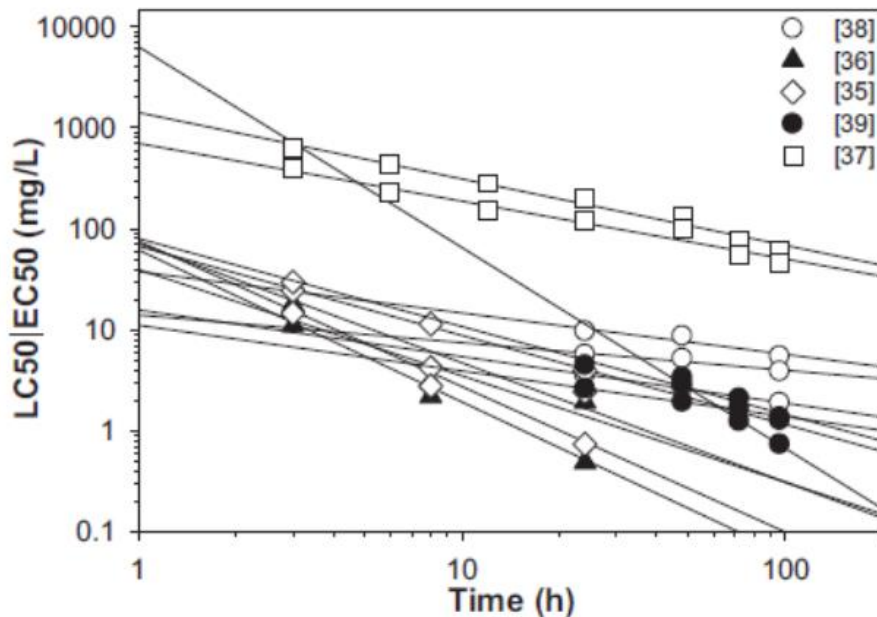


Figure 2. Lethal and effects concentrations affecting 50% of test organisms (LC50, EC50) versus duration of exposure in experiments reviewed by Bejarano et al. (2014; see source for original citations).

### Oil Dynamics

For most oil spills, exposures of water column biota to concentrations above potential thresholds of concern are typically on time scales of minutes to hours, even for spills lasting weeks or months because of the varying movements of the oil in the water, dilution and losses to biodegradation and volatilization. Furthermore, the concentrations vary in time over the short exposure periods (McAuliffe et al. 1980, 1981; McAuliffe 1987; Lunel 1994; French McCay 2002, 2004; Bejarano et al. 2014; French-McCay et al. 2023).

Marine waters are not static systems, as the oil rapidly spreads and moves through space over time. Oil is concentrated floating on the surface, along shorelines, and as it settles to sediments, while it is dispersed in three-dimensional space in open water. When floating oil is entrained, a portion refloats immediately, more so for larger droplets originating from more viscous oils. Entrained oil in smaller droplets than those that refloat, rapidly disperses away from the surface. Currents and dispersion in the water column, combined with differential buoyancy of oil droplets of various sizes, rapidly dilute concentrations of entrained droplets. Therefore, oil exposures in open water are typically of short duration.

Figure 3 provides an example of the changing composition of dissolved hydrocarbons and the short duration of exposure to high concentrations in a tank containing floating crude oil held in place by the containment (Payne et al. 1984). If the floating oil were in the ocean, the dissolved components would be spread by the movement of the oil over the surface and by current shear, such that concentrations would be much lower, more ephemeral, and patchy. Bejarano et al.

(2014) reviewed measurement data from field trials finding oil concentrations decreased rapidly (in minutes) after oil release (Figure 4).

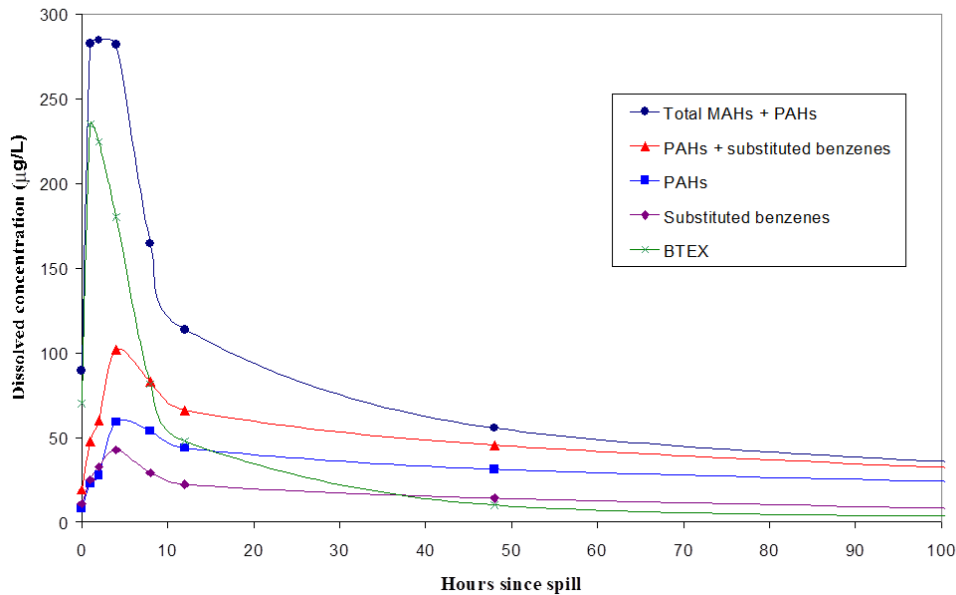


Figure 3. Dissolved concentrations versus time in a tank (data from Payne et al. 1984)

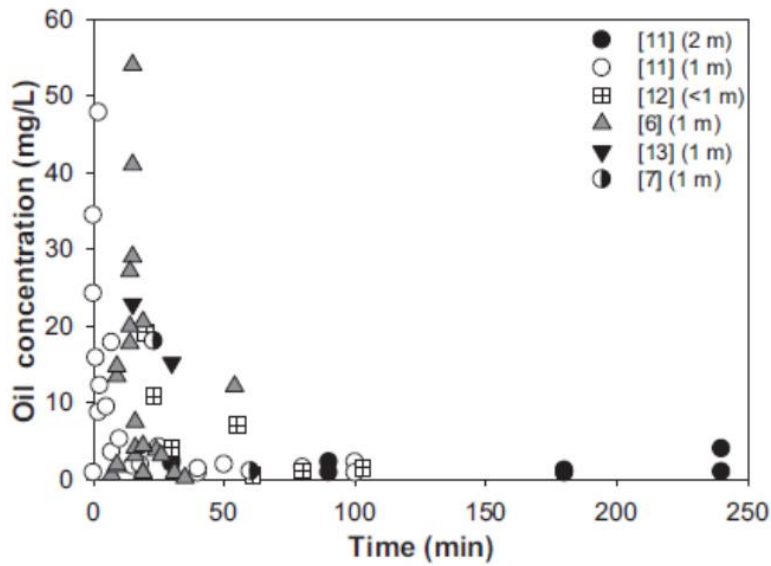


Figure 1. The relationship between measured oil concentrations in the upper water column ( $\leq 2$  m) versus time from field trials at sea. Data represent oil concentrations from physically ([11], black circles) and chemically enhanced dispersion of oil ([11], white circles; [6,7,12,13]).

Figure 4. Measured oil concentrations versus time after release from field trials (from Bejarano et al. 2014; see source for original citations).



## Organism Behavior

Organisms are not distributed evenly in the environment and are often more concentrated in coastal areas and structural habitats such as coral reefs, rocky reefs, seagrass beds, and seaweeds. Early life history stages of fish and invertebrates are concentrated in these habitats and also in surface waters. In addition, animals move under their own volition and passively (as plankton) in currents, while animals that are attached to structures or otherwise stationary, would be exposed to oil components in plumes that would pass by them. Thus, the duration of exposure to hydrocarbons depends on the distribution of the organisms and their entrainment or movements in relation to the dynamics (changes in) the oil contaminants plume.

A recent modelling study by French-McCay et al. (2024, in review) provides examples of exposure patterns planktonic and stationary organisms might experience for different oil types and spill situations. For example, Figure 5 shows exposures over time for stationary (non-mobile) and planktonic organisms for a modelled 1000 bbl (159 m<sup>3</sup>) spill of a light crude (Hibernia) in 7 m/s winds.

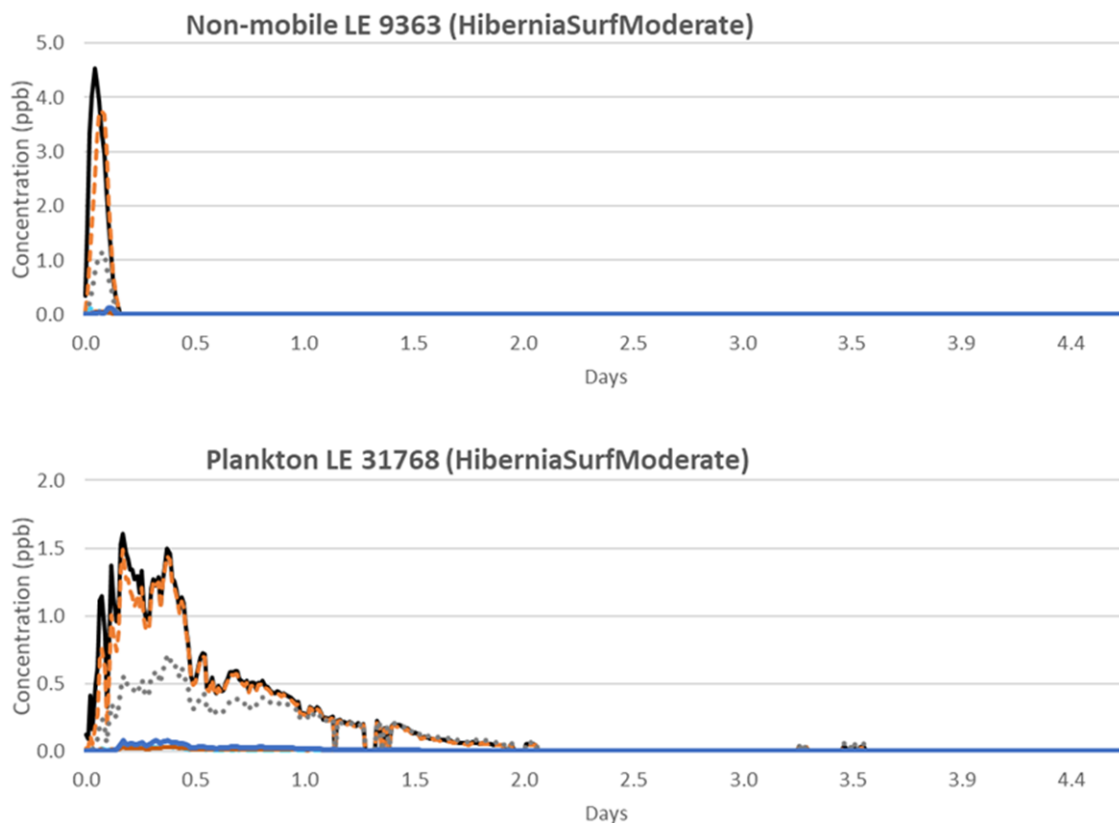


Figure 5. Example exposures to various dissolved oil components over time for stationary (non-mobile) and planktonic organisms after a 1000-bbl spill of light crude.

Note that organisms would need to encounter dissolved concentrations or entrained droplets for any adverse effects to occur, but many are not present in the same depth range as the oil

contamination (e.g., surface waters). Other organisms might move through the area of contamination, where their exposure would be for shorter duration than for plankton or stationary organisms. Thus, a 2-dimensional map of maximum concentrations in the vertical dimension, a typical summary presentation of oil modelling results, would not apply to the entire water column.

## **Measurements of Oil Concentrations as Total Hydrocarbons**

Measurement of the concentration of oil in water, whether in the field or in laboratory studies, is complicated and not all constituents can be identified and measured individually. There are many analytical chemistry techniques used to estimate total hydrocarbons in water, and each of these only measures a portion of the hydrocarbons in oil (Redman et al. 2012; Redman and Parkerton 2015; Yang et al. 2017, Parkerton et al. 2023a). Recent guidance (Redman and Parkerton 2015; Dettman et al. 2023) recommends measuring BTEX and C5–C9 aliphatics (“volatiles”) using purge-trap GC–MS (Gas Chromatograph–Mass Spectrometer), analysis of parent and alkyl PAHs using GC–MS (with target analytes including C0–C4 alkyl decalins and C0–C4 alkyl phenols), and C9–C40 saturate hydrocarbons by GC/FID. Older studies often did not include this detail and range of analyses (see review in Bejarano et al. 2023).

Measurements of C8 to C50 hydrocarbons in oils using these techniques by Yang et al. (2017) identified only 20-30% of the compounds, the remainder of the total petroleum hydrocarbon (TPH) being the unresolved complex mixture (UCM). Mass recoveries (using the presently recommended methods) for gasoline, kerosene, and gas oil are typically >95%, whereas the mass recoveries for heavier substances in crude oils and heavier refined products are often <30%. Mass recoveries of condensates would be somewhat higher than for crude oils but <95%.

The uncharacterized mass (UCM) is due to high-molecular-weight alkanes, asphaltenes, resins, and other large molecules that are either not eluted from chromatographic columns or not well-resolved by gas chromatographic techniques that are available (Redman et al 2012; Yang et al. 2017). In the discussion below THC will refer to the true concentration of total hydrocarbons (and related compounds), whereas TRH will be used to refer to the measured (i.e., recoverable) portion of the THC.

Because only some of the compounds in oil are measurable individually, and the desire to be inclusive of effects from hydrocarbons in oil droplets (e.g., the entrained droplets in SIMAP model outputs) as well as the dissolved phase, aquatic toxicologists have attempted to use THC as a metric for evaluating toxicity of oil exposures. However, given the complexities described above, among others related to experimental design and conditions, development of lethal toxicity values (e.g., LC50s) or sublethal thresholds based on total hydrocarbons (THC) is problematic. The hydrocarbon compositions in exposure media are highly variable from experiment to experiment, leading to estimated toxicity thresholds that range over many orders of magnitude, as is evident from reviews of these data (e.g., see NRC 2003; NASEM 2020).

## **Oil Bioassays**

Toxicity studies often have used experimental systems containing media preparations made by mixing oil in water long enough to bring the composition of oil components into equilibrium, i.e., Water Accommodated Fractions (WAFs) of oil. WAFs are purposely static, and thus, do not represent variable field compositions and changing concentrations. WAFs can be made with fresh or weathered oil, but bioassays using a particular WAF do not account for weathering changes in the field. Further, as the concentrations of individual compounds in WAFs change from test to test and the individual compounds vary in toxicity, effects levels based on total PAH or TRH are highly variable. LC50s or other toxicity metrics based on nominal additions of oil to media (in an attempt to measure a THC effects level) are unreliable because some compounds evaporate, adsorb to test containers, bioaccumulate, and degrade in test chambers, and the oil component mixture composition is unknown (Hodson et al. 2019). The 2023 Special Issue of Aquatic Toxicology (Loughery et al. 2023; Stubblefield et al. 2023; Alloy et al. 2023; Parkerton et al. 2023a,b; French-McCay et al. 2023; Dettman et al. 2023; Bejarano et al. 2023; Lee et al. 2023) contains reviews of toxicity test methods, including the issues summarized here. Loughery et al. (2023), Stubblefield et al. (2023), Alloy et al. (2023), Parkerton et al. (2023a,b), Dettman et al. (2023), Bejarano et al. (2023), and Lee et al. (2023) provide guidance for aquatic toxicity studies, building on and updating the standardized toxicity test protocols developed by the Chemical Response to Oil Spills: Ecological Effects Research Forum (CROSERF; Aurand and Coelho 2005, Singer et al. 2000).

Thus, toxicity parameters developed using WAFs should not be applied directly to field conditions to evaluate effects. TU modelling is used to evaluate the toxicity in the bioassay mixture (Parkerton et al. 2023a), and the toxicity of the mixture in the field, for which the composition and toxicity will differ. Whole oil bioassays using WAFs or other preparations exposing organisms to dissolved components are useful and important for validating the predictions of toxicity models based on additive effects of oil components (French-McCay et al. 2023).

WAF-based effects levels have been used to develop concentration thresholds above which effects might be expected. Even if the experimental oil composition of a test were representative of the field under the specific conditions tested, the experimental effects level would over-estimate toxicity to the extent it is applied to modelled oil, as the model accounts for the entire mass of oil, as opposed to what was measurable in the test.

## **Recommendations for Thresholds**

### **PAH-Based Threshold for Dissolved Hydrocarbons**

Because most of the toxicity from dissolved hydrocarbons and related compounds is due to PAHs and related compounds, particularly in surface waters, PAH-based lethal and sublethal thresholds are typically used to evaluate the risks of oil contamination. In many bioassays, measured total PAH concentrations have been used to quantify toxicity endpoints, with the understanding that the measured PAH concentrations reflect the effects of any dispersants

applied (which increase concentrations in the water, Bejarano et al. 2014) and other unmeasured constituents associated with the PAHs are contributing to the observed toxicity (Bejarano et al. 2017; Forth et al. 2017; Parkerton et al. 2023a), although with less effect (French-McCay 2002).

PAH concentrations on the order of tens to hundreds of micrograms per liter ( $\mu\text{g/L}$ , i.e., ppb) have been shown to be acutely toxic to aquatic biota (French-McCay 2002, 2016). French-McCay (2002) reviewed available laboratory oil and PAH acute bioassay data, finding that 95% of species and life stages exhibited 50% mortality between 6 and 400 ppb ( $\mu\text{g/L}$ ) total PAH concentration. Based on this result, the 6 ppb total PAH value would be protective of 97.5% of species and life stages. Early life history stages of fish appear to be more sensitive than older fish stages and invertebrates. Bioassay data compiled by French-McCay (2002) indicate that 96+-hour LC50s (Lethal Concentrations to 50% of exposed biota) of juvenile and adult fish are >100 ppb. Bejarano et al. (2017) developed species sensitivity distributions (SSDs) for acute bioassay tests using WAFs. Bejarano et al (2017) found that the 5<sup>th</sup> percentile sensitivity (Hazardous Concentration for 5%, HC5, Posthuma et al. 2002, which is protective of 95% of species and life stages) for lethality ranged from 28 to 282 ppb total PAH concentration for fish and invertebrates from a wide variety of latitudes and habitats, a range similar to that predicted by French-McCay (2002). Note that in recent years the 5<sup>th</sup> percentile for a range of species sensitivities in toxicity tests has come to be used in practice for developing thresholds, and this is termed the HC5 (Barron et al. 2013; Fox et al. 2021; Stubblefield et al. 2023). Thus, for acute lethality and to the nearest order of magnitude (given uncertainties), total PAH LC50s generally range from about 10 ppb for sensitive early life history stages to 300 ppb or more for less sensitive species and older life stages.

Sublethal effects may occur at lower concentrations. For PAHs, USEPA (2003, 2008) has derived and used a factor 10 to estimate a sublethal effects threshold (Final Chronic Value, FCV, or Predicted No Effect Concentration, PNEC) based on a lowest acute toxicity-based endpoint such as the HC5 based on 96-hour LC50 tests. In a recent study by McGrath et al. (2018), the acute-to-chronic ratio (ACR) for hydrocarbons was found to average 5.22 (range 1.7-12 for the 10<sup>th</sup> and 90<sup>th</sup> percentiles of the observations, including consideration of one high outlier), therefore, the factor 10 is conservatively protective. Thus, the sublethal effects threshold (PNEC) for PAHs would be 1  $\mu\text{g/L}$  (ppb). Further details are available in French-McCay (2009) and French-McCay et al. (2018). The 1 ppb ( $\mu\text{g/L}$ ) and 10 ppb ( $\mu\text{g/L}$ ) total PAH thresholds have been used in many environmental risk assessments (e.g., French McCay et al., 2005, 2012, 2018; NOAA, 2013).

In addition, photo-chemical reactions upon exposure of oil or contaminated organisms to sunlight produces photoproducts (new compound structures, by photo-modification) or free radicals in organisms (photosensitization to activated PAHs and related compounds) that could adversely affect exposed aquatic organisms. Photosensitization can be significant for early life stages of fish and invertebrates that are translucent or transparent and vulnerable since they are undergoing development (Alloy et al. 2023). Specific thresholds have not been developed that account for photo-toxicity. However, studies indicate the effects threshold accounting for phototoxicity would be approximately 1  $\mu\text{g/L}$  (ppb). Further details are available in French-McCay et al. (2018).

## THC-Based Threshold for Entrained Droplets

The concentrations of entrained droplets output by the SIMAP model represent hydrocarbons that are not bioavailable. The soluble and semi-soluble fractions dissolve from the droplets over time, and a potential effects analysis based on the dissolved hydrocarbons characterizes their risk. However, to satisfy concerns that entrained droplet hydrocarbons might infer some risk, THC-based thresholds have been used.

In addition to considering total PAHs, Bejarano et al. (2017) developed SSDs using *measured* TPH (i.e., TRH; aromatic and aliphatic hydrocarbons [C9-C44], which includes parent and alkylated homologue PAHs and/or parent naphthalene) as a metric. The range of HC5s was found to be 1 to 560 ppm (mg/L) as TRH for WAF preparations from crude oils with API densities ranging from 25 to 44. The HC5 for light refined Group 2 fuel oil (API 35 – 45) was 0.5 mg/L TRH. The TRH HC5 decreased with higher API (lower oil density), i.e., with oils that have a higher percentage of light hydrocarbons (C9-C44) including PAHs. WAFs included aqueous exposure media prepared by physical (low energy water accommodated fraction, LEWAF; and moderate energy WAF or MEWAF) or chemically enhanced oil dispersion (chemically enhanced water accommodated fraction, CEWAF).

Note that the TPH measurements were on WAF media, which are preparations that purposely select for the soluble hydrocarbons and remove the insoluble larger hydrocarbons in the oil (Redman and Parkerton 2015). Thus, the TPH measurements of the WAF media are on a portion of the oil, not the full oil (see also Parkerton et al. 2023a for further detail). This is demonstrated by the ratios of measured TPH to PAHs, where they were both measured on the same WAFs. The ratios were 35-40 for LEWAF and MEWAF preparations and 14 for CEWAF preparations (Table 1). These ratios are far lower than the typical percentage of PAH in crude oils, which is 1% for PAH/THC (or ratio of 100; French-McCay 2002; Forth et al. 2017). Therefore, these TPH measurements reflect only a small percentage of THC, less than 40% of the oil, and for CEWAFs about 15% of the oil. Other analyses of crude oils indicate similar percentages (e.g., Forth et al. 2017). Therefore, the TRH HC5s derived by Bejarano et al. (2017) should be multiplied by a factor 3-7, based on the measured TPH/PAH ratios, to correct for the missing hydrocarbons in the oil since the SIMAP model outputs are for all the hydrocarbons in oil, not just the measurable ones. This yields THC lethal thresholds of 3-28 ppm (mg/L, Table 1), based on the Bejarano et al. (2017) review. Similar adjustments should be made for TRH-based thresholds, depending upon the methods used.

**Table 1.** Measured total PAH and TPH for lethal effects thresholds based on WAF preparations of oil (Bejarano et al. 2017) where 95% of species would be protected. The loss corrected TPH accounts for the fraction of the parent oil not included in the WAF preparation.

Media	PAH (ppb, $\mu\text{g/L}$ )	Measured TPH (ppb, $\mu\text{g/L}$ )	TPH/PAH	Loss-Corrected TPH (i.e., THC) (ppm, mg/l)
WAF + MEWAF light crude oils	27.6	956	35	2.7
WAF + MEWAF medium crude oils	74.9	2,935	39	7.5
CEWAF medium crude oils	282	3,907	14	27.9

This limitation also exists for measurable TPH (i.e., TRH) thresholds presented in Smit et al. (2009), which has been used as a proxy THC threshold for oil spill modelling in the past. However, the measured TPH/PAH ratio is not provided in that paper, so the same correction calculations cannot be carried out. Furthermore, the thresholds developed in Smit et al. (2009) are for no-observed-effect concentrations (NOECs) based on sublethal effects of chronic exposures of 7 to 183 days (e.g., the lowest test NOEC was for a 33-day exposure). As discussed above (and in French-McCay et al. 2023), such long exposure times do not occur in open waters after oil spills and effects concentrations are much higher for short-term exposures.

The ANZECC 2000 guidelines and derivation of a TPH-based sublethal threshold include a review of a few studies available at the time but depend on the analysis by Tsvetnenko (1998) for a threshold. Tsvetnenko (1998) followed US EPA methods (Stephan et al. 1985; US EPA 1994) to develop SSDs and HC5s but did not recognize that TPH measurements of Water-Soluble Fractions (WSF) and WAFs do not represent the same composition as the THC in the field. They are biased towards a mix of soluble and semi-soluble components that are the toxic components of the oil (as explained above). Tsvetnenko (1998) developed a lethal HC5 for TPH of 168  $\mu\text{g/L}$  (ppb), and then applied an ACR of 25 to obtain a sub lethal threshold of 7  $\mu\text{g/L}$  (ppb) as TPH. Note that recent reviews do not support this ACR. An ACR=10 would be sufficiently protective for chronic exposures. Even if it were assumed that TPH measurements on WSF/WAF media were representative of THC (which they are not), the HC5 of 168  $\mu\text{g/L}$  should be corrected (upward) for the fraction of the THC of the source oil actually measured by the TPH analyses used and should not be corrected (downward) for the volatilization loss (in table 8.3.24, ANZECC 2000 appears to have lowered the measured TPH toxicity values by a factor of 10, based on Hamoda et al. 1989). The needed data are not available, but likely the correction would be an order of magnitude, given the time and nature of those TPH measurements and apparent volatilization in the experiments. However, it is more appropriate to use recent data where better analytical methods were used, if THC is to be the basis of a threshold.

Because PAHs are the most toxic components of oil and crude oils typically contain about 1% PAHs by mass (French-McCay 2002; Forth et al. 2017), the sublethal concentration threshold (PNEC) expressed as total hydrocarbon concentration (THC, not TPH) based on the most toxic components would be  $\sim 100 \mu\text{g/L}$  (100 ppb) for fresh oil. However, as oil weathers, PAHs are lost to volatilization and degradation. Thus, the whole-oil threshold of 1,000 ppb (1 ppm) is conservative (low, and highly protective of aquatic resources). Based on the review of toxicity studies by Bejarano et al. (2017), THC lethal effects levels of 3-28 mg/L (or 3-30 mg/L [3-30 ppm] with rounding, given uncertainties) would be appropriate for a range of oils and states of weathering for species from all geographical areas globally. An exposure concentration of 1,000 ppb (1 ppm or 1 mg/L) of measurable TPH was deemed a low level of concern for sensitive life stages in marine organisms by Kraly et al. (2001). In reviews by NRC (2005) and NASEM (2020), 1,000 ppb was found to be at the low end of the range where sub-lethal impacts from acute exposure have been observed. Correcting for fraction measurable, this indicates a threshold of 3,000 ppb would be appropriate for modelled entrained oil.

Thus, a THC threshold of 3 mg/L (3 ppm; 3,000 ppb) is an appropriate threshold for crude oils for use in risk assessments and for defining an EMBA. For light distillates and condensates, 1 mg/L (1 ppm; 1,000 ppb) is an appropriate threshold.

### **Consideration of Duration of Exposure for Developing Thresholds**

The use of LC50s for >48 hours of exposure, or chronic endpoints for longer exposure times, as thresholds for oil spills is highly conservative. Acute aquatic toxicity thresholds would be sufficiently conservative for oil spills in open water systems (as opposed to ponds or other contained systems). There is no need for an ACR correction for evaluating acute toxicity to aquatic biota from oil spills in open waters.

In the ANZECC 2000 guidelines, no observable effects concentrations (NOECs) after long exposure durations are recommended. This is highly protective for chronic steady state concentrations of hydrocarbons. Again, since oil spill water column exposures are acute, and durations of exposure are for hours and not days, acute toxicity data (e.g., Final Acute Value, FAV) should be used and there should be no ACR conversion.

To in some part compensate for the effect of exposure duration on effects levels, a higher threshold appropriate for short duration exposures could be used. Alternatively, the modelled exposure concentrations could be time averaged over 1 to several hours.

Another approach would be to use a dose metric as a threshold. The LC50s quoted above are typically for 96 hours of exposure. Thus, the 10-ppb dissolved PAH threshold would be equivalent to 960 ppb-hours of exposure. The 1,000 ppb THC threshold would be equivalent to 96 ppm-hours. Model results could be integrated to calculate the ppb-hours of exposure, therefore addressing the duration of exposure issue.

## **Conclusion**

### **Thresholds**

For the dissolved hydrocarbons, the 1 ppb PAH threshold is conservatively protective of aquatic biota. If THC in oil droplets is to be evaluated as a risk, 1 ppm (1,000 ppb) would be sufficiently conservative for oil droplets of all oil types and all weathering states, with 3 ppm (3,000 ppb) an appropriate threshold for crude oils.

It should be noted that toxicity is determined by the specific mix of hydrocarbon concentrations in the exposure medium and the toxicity of each component. Use of a THC threshold does not address the complexities of the mixture of hydrocarbons of various toxicities. For these reasons, recent evaluations of the aquatic effects of oil have focused on the most toxic and bioavailable components (i.e., dissolved hydrocarbons) and have not used measured TPH or THC as a metric. The review series (Loughery et al. 2023; Stubblefield et al. 2023; Alloy et al. 2023; Parkerton et al. 2023a,b; Dettman et al. 2023; Bejarano et al. 2023; Lee et al. 2023; French-McCay et al.

2023) provides guidance for aquatic toxicity studies and their application to risk assessments and modelling-based analyses. The focus of a risk assessment should be on the dissolved hydrocarbon exposure, along with floating oil and shoreline oil, as opposed to oil droplets.

### **Consideration Regarding EMBA**

It should be noted that exceedance of threshold does not infer adverse effects would occur. This has been understood by NOPSEMA (2019) and others. If a threshold is very conservative, use of the EMBA delineation would not focus monitoring on locations where there is a potentially high risk of consequences. Near the outer bounds of an EMBA, defined by an entrained oil threshold of 10 ppb (as recommended by NOPSEMA 2019), there is a very low probability of concentrations exceeding 10 ppb, the exposure durations are typically minutes to a few hours, and the oil is highly weathered (and so less toxic). Thus, a more realistic threshold would be appropriate, and it need not be protective of every single individual organism that might possibly encounter one droplet of entrained oil, regardless of likelihood or its ability to cause any adverse effect.

Monitoring should focus on locations most at risk of consequences, as opposed to those with marginal, if any, risk of consequences. Such locations include near the release site, in shallow waters, in sensitive habitats, and in areas where there are species with protection status. Monitoring to measure any change is challenging even for very large spills due to natural variability and the large volumes of oceanic waters needing to be sampled. Sampling effort, experimental design and analytical chemistry capabilities need to be carefully considered, otherwise resources would be wasted. Thus, even higher thresholds than those recommended here could be used to identify areas where monitoring would be appropriate and successful in detecting change or effects.

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