Phototoxic effects of two common marine fuels on the settlement success of the coral *Acropora tenuis*

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Coral reefs are at risk of exposure to petroleum hydrocarbons from shipping spills and uncontrolled discharges during extraction. The toxicity of petroleum hydrocarbons can substantially increase in the presence of ultraviolet radiation (UVR), therefore spills in shallow coral reef environments may be particularly hazardous to reef species. Here we investigated the sensitivity of coral larvae (*Acropora tenuis*) to dissolved hydrocarbons from heavy fuel oil (HFO) and diesel in the absence and presence of UVR. Larval settlement success decreased with increasing concentrations of dissolved HFO, and co-exposure to UVR doubled the toxicity: 50% effect concentrations (EC50) decreased from 96 (−UVR) to 51 (+UVR) total petroleum aromatic hydrocarbons (TPAH). Toxic thresholds for HFO were similar to concentrations reported during marine spills: EC10s of 24 (−UVR) and 15 (+UVR) µg l\(^{-1}\). While less toxic, diesel also reduced settlement and exhibited phototoxicity: EC10s of 122 (+UVR) and 302 (−UVR) µg l\(^{-1}\). This study demonstrates that the presence of UVR increases the hazard posed by oil pollution to tropical, shallow-water coral reefs. Further research on the effects of oils in the presence of UVR is needed to improve the environmental relevance of risk assessments and ensure appropriate protection for shallow reef environments against oil pollution.

**Petroleum hydrocarbons in marine environments.** Petroleum hydrocarbons are considered among the most prominent pollution threats to marine environments; however, the risks they pose to coral reef ecosystems remain poorly understood. The environmental effects resulting from oil spills and uncontrolled discharges from extraction vary widely and are dependent on a large number of factors. Hydrocarbon concentrations in marine environments have been measured following large scale spills, with dissolved hydrocarbon concentrations ranging between 22 and 189 µg l\(^{-1}\) total polycyclic aromatic hydrocarbons (PAH) and up to 10,600 µg l\(^{-1}\) total recoverable hydrocarbons (TRH). Hydrocarbons can be retained within an ecosystem during spill events and, despite natural dilution and degradation, can remain detectable long after the spill has ended.

The toxicity of petroleum hydrocarbons to most marine species is predominantly related to the water soluble components, which largely consist of the monoaromatic hydrocarbons (MAH) and PAHs. PAHs, in particular, are considered acutely toxic to aquatic biota, with toxic threshold concentrations for PAHs often orders of magnitude lower than those of MAHs. However, the overall toxicity of dissolved petroleum hydrocarbon from spills is also dependent on the relative concentrations of each component. Aromatic hydrocarbons in petroleum oils are classified as type I narcotic chemicals. Assuming the same mode of toxic action, the total toxicity of the water soluble components of an oil can be predicted using the narcotic target lipid model (NTLM), which sums the expected toxicity and concentration of each aromatic component (see Methods).

**Phototoxicity of PAHs.** The toxicity of the dissolved aromatic mixtures that result from oil spills may increase in the presence of UVR due to the phototoxic contribution of some PAHs. PAH phototoxicity occurs through the formation of radical oxygen species and/or transformation of PAHs into more toxic photoproducts. Co-exposure of PAHs and UVR can increase the toxicity of individual PAHs 1000-fold; however, the...
potential for phototoxicity depends on the compound’s stability, radiation absorbance properties, the type of UVR exposure and the exposed organism’s sensitivity. The potential for harmful effects to marine organisms is higher for UVA (320–400 nm) than UVB (280–320 nm) as the absorption maxima (hence, photoactivation) of many PAHs fall within the UVA range and UVB is more strongly attenuated in seawater. The penetration of UVR in marine environments is also dependent on a range of physical and biological factors, and tropical oligotrophic coral reefs may be at a particularly high risk of PAH phototoxicity as reef organisms are frequently exposed to high solar radiation, including UVR. PAH phototoxicity is not always taken into consideration for risk assessments and management unless the ecological relevance, including UVR exposure and spectral profiles, in the ecosystem have been characterised. However, it is increasingly recognised that a considerable proportion of UVR penetrates to ecologically relevant depths in some marine environments, including coral reefs, and that exposure to cumulative pressures, such as pollution and UVR, can result in increased environmental impacts.

**Petroleum hydrocarbon and PAH toxicity to corals.** Despite a renewed demand for marine hydrocarbon toxicity research following the Deepwater Horizon spill, significant knowledge gaps on the potential effects of hydrocarbon exposure to corals remain, and the sensitivity of tropical marine species to hydrocarbons is relatively understudied. Investigations into the effects of petroleum hydrocarbons, including PAHs, on coral indicate that negative impacts can occur at concentrations as low as 2–20 μg l⁻¹ total hydrocarbons (THC). However, inconsistencies in exposure methodologies, toxic endpoints and reported toxicity values make comparisons between studies problematic. This issue is further compounded by the failure of many studies to present the chemical composition of treatment solutions, in particular the more soluble and toxic MAHs and PAHs. However, the studies that have been conducted show that all life history stages of coral and that larval settlement is generally more sensitive than fertilization, larval survival, or the health of mature coral and that larval settlement process is recognised as an ecologically relevant endpoint due to its importance in the recruitment process and subsequent maintenance of adult populations. The larval life stages of aquatic animals may also be at higher risk than adults to phototoxic effects due to their small size, often transparent bodies, and time spent in shallow waters.

Four studies have investigated phototoxic effects of petroleum hydrocarbons on corals and each indicated that their sensitivity to dissolved aromatics may increase with co-exposure to UVR (summarised in Table 1). However, the majority of laboratory studies exposing corals to hydrocarbons have not included co-exposure to UVR, so the impacts of hydrocarbon pollution on coral reefs may be significantly underestimated in the context of likely high UVR exposure in situ. To assess the potential for UVR to increase the sensitivity of coral larvae to spills of heavy fuel oil (HFO) and diesel, we: (i) assessed UVR irradiance on one inshore and one mid-shelf reef on the Great Barrier Reef (GBR; Australia); (ii) characterised the chemical composition of the two fuels and their water accommodated fractions (WAFs); and (iii) predicted their narcotic toxicity to marine species using the NTLM. We then (iv) exposed larvae of the reef building coral Acropora tenuis to HFO and diesel WAFs in the absence and presence of UVR (+UVR), at intensities similar to those encountered on the GBR, and assessed the ability of exposed larvae to successfully complete settlement and metamorphosis into sessile polyps following each treatment.

**Results**

**Chemical analysis.** Neat HFO consisted primarily of higher molecular weight hydrocarbons while neat diesel contained a higher proportion of BTEX (benzene, toluene, ethylbenzene and xylene) and other lower molecular weight hydrocarbons (Figs S-1 and S-2, Supplementary information). ∑PAH in HFO (constituting 99.6% of TPAH) was almost 10-fold higher than in diesel (79% of TPAH), whereas ∑BTEX was ~8-fold lower in HFO compared to diesel (see summary in Table 2 and detailed results in Table S-1, Supplementary information). Phenols were below the limit of quantitation in both oils (Table S-1, Supplementary information). Freshly prepared 100% HFO and diesel WAFs contained similar TPAH concentrations (930 and 913 μg l⁻¹, respectively), but the proportions of ∑BTEX and ∑PAH varied (Table S-2, Supplementary information). While HFO WAF contained almost equal concentrations of ∑BTEX and ∑PAH (52% and 48% of TPAH, respectively), ∑BTEX dominated the diesel WAF (98% of TPAH). The most abundant PAHs in the WAFs were naphthalene, alkylphenothalenes, fluorene and phenanthrene; the HFO WAF also containedacenaphthene and dibenzothiophene (Table S-2, Supplementary information). The concentration of TPAH in fuel WAFs decreased by up to 34% over the 48 h exposure (Tables 2 and S-2, Supplementary information). Comparisons of the observed and predicted concentrations of 1-, 2- and 3-ring compounds in the freshly prepared, undiluted WAFs (Tables S-2 and S-3, Supplementary information) as per Redman et al. indicated that no oil droplets were present in either HFO or diesel WAFs.

**Ultraviolet radiation intensities.** The spectral profiles for both the Trunk Reef (mid-shelf) and Esk Reef (inshore) sites showed that the largest decrease in irradiance occurred when light passed from air to water (Fig. 1a and b). Attenuation of UVR at Esk Reef was somewhat lower than at Trunk Reef for shallow measurement depths, despite the higher turbidity (0.1 and 0.8 nephelometric turbidity units, NTU, respectively). However, irradiance was attenuated less with depth at Trunk Reef and the irradiance for the deepest measurements was much higher at Trunk Reef than at Esk Reef (Fig. 1a and b). The depth which 10% of UVR irradiance penetrated to (Z₁₀%) for 305 and 340 nm at Trunk Reef was ~6 and 7 m, respectively. On Esk Reef, Z₁₀% was ~2.7 and 2.4 m for 305 and 340 nm, respectively.

Experimental lights, when positioned 170 mm from the sensor, emitted UVA radiation similar to the irradiance observed at 1 m depth on Trunk Reef and Esk Reef (Fig. 1c). Average total UVA and UVB (280–400 nm) radiation was 0.9 mW cm⁻² (±0.16). Attenuation by the glass scintillation vials used in the experiment redused the average absolute irradiance by ~17% for wavelengths between 300–400 nm (Fig. 1c; measured using
Table 1. Summary of previous studies of the phototoxic effects of petroleum hydrocarbons on scleractinian corals. Study methodology, species tested, chemical analysis performed (if applicable), toxic endpoint and main results shown for each study. If no threshold values or concentrations are presented no effect was observed. Seq = sequential, organisms first exposed to pollutant followed by exposure to UVR while kept in clean FSW. GC-MS = gas chromatography-mass spectrometry. Nil = no analysis reported. * Stress-response-related gene expression and enzyme activity were also investigated (see reference for further details).

### Larval settlement assays

Temperature was maintained at 27.6 ± 1.4 °C (mean ± SD) in the experiments while photosynthetically active radiation (PAR) in the + UVR and − UVR treatments averaged 0.95 ± 0.10 and 1.23 ± 0.10 mW cm−2, respectively. Dissolved oxygen concentration averaged 7.9 ± 0.33 mg l−1 with all replicates maintaining concentrations >7.0 mg l−1 for the duration of the exposure period while pH and salinity averaged 8.1 ± 0.06 and 37.0 ± 0.32 psu, respectively. *A. tenuis* larvae in control treatments were observed to actively swim throughout the exposure (Fig. 2a), but swimming behaviour was not assessed in the fuel exposure treatments. In the control treatments, an average of 73% (SE = 4) larvae underwent settlement in the presence of CCA chips within 24 h (Fig. 2c). This level of settlement success did not change in the presence of UVR with 77% (SE = 3.5) of larvae successfully undergoing settlement. Average larval settlement ≥70% in control treatments was considered indicative of a normal response to settlement inducers based upon several previous studies using CCA or extracts of CCA to initiate settlement of *Acropora* spp.28,29,30.

### Heavy fuel oil toxicity

HFO WAF inhibited larval settlement in both the absence and presence of UVR (Fig. 3a,c and Table 3). Little or no effect on settlement success was observed at low concentrations (<10 µg l−1 TPAH); above which, settlement decreased with increasing TPAH concentration (Fig. 3). The toxicity of HFO WAF was enhanced in the presence of UVR resulting in a ~50% decrease of the EC50 from 96 to 51 µg l−1 TPAH (95% confidence intervals did not overlap; Table 3). The toxic threshold value (EC50) also reduced in the presence of UVR (Table 3). Obvious mortality (disintegrating cell membranes) was only observed in the highest TPAH treatment (890 µg l−1), in the presence of UVR, at the end of the 48 h exposure. No other visible effects on larvae, in either control or HFO WAF treatments, were observed and surviving larvae in the highest HFO treatments exhibited normal morphology (Fig. 2a and b). At 72 h (following the 24 h settlement period) unattached larvae and attached juvenile polyps in control treatments retained normal morphology (Fig. 2c). At this point the frequency and severity of abnormalities increased with increasing TPAH concentration in attached juvenile polyps (asymmetrical or underdeveloped recruits; Fig. 2d) and non-settled larvae (bumps, deformities and necrosis; Fig. 2e) for both ± UVR treatments. Severe deformities were observed in unattached larvae at TPAH concentrations as low as 28 µg l−1 (+ UVR) and a substantial proportion of larvae were immobile and/or dead in the highest concentration treatment (890 µg l−1). Most of the successfully attached juvenile polyps underwent complete metamorphosis in the absence of UVR; however, larvae exposed to HFO WAFs appeared to develop more slowly than expected and some had only undergone partial metamorphosis in higher concentration treatments at the time of assessment.
Table 2. Time-averaged concentrations of $\sum\text{BTEX}$, $\sum\text{PAH}$ and TPAH in undiluted fuel WAFs, $\sum\text{BTEX}$, $\sum\text{PAH}$ and TRH in neat fuels, and toxic units (narcosis) calculated from predicted and observed fuel WAF concentrations. Time-averaged concentrations per light treatment calculated from concentrations measured at $t_0$ and $t_{48h}$ for HFO and diesel WAFs; % change in concentrations after 48 h indicated in brackets. TUs calculated using an average CTLBB value (86.8 μmol g$^{-1}$ octanol; n = 15) and aqueous concentrations of BTEX and PAH observed in fuel WAFs ($\text{TU}_{\text{WAF}}$) or predicted from neat fuel oil concentrations ($\text{TU}_{\text{Neat fuel}}$).

|        | $\sum\text{BTEX}$ | $\sum\text{PAH}$ | TPAH | $\sum\text{BTEX}$ | $\sum\text{PAH}$ | TPAH | $\sum\text{BTEX}$ | $\sum\text{PAH}$ | TPAH | $\sum\text{BTEX}$ | $\sum\text{PAH}$ | TPAH |
|--------|-------------------|-------------------|------|-------------------|-------------------|------|-------------------|-------------------|------|-------------------|-------------------|------|
| HFO    |                   |                   |      |                   |                   |      |                   |                   |      |                   |                   |      |
| - UVR  | 498 (+4%)         | 440 (+11%)        | 938 (+2%) | 190              | 50494            | 50684 | 0.03              | 0.43              | 0.46 | 0.01              | 0.91              | 0.92 |
| + UVR  | 478 (+4%)         | 414 (+13%)        | 892 (+8%) | 190              | 50494            | 50684 | 0.03              | 0.40              | 0.43 | 0.02              | 0.92              | 0.92 |
| Diesel |                   |                   |      |                   |                   |      |                   |                   |      |                   |                   |      |
| - UVR  | 745 (−33%)        | 14 (−57%)         | 759 (−34%) | 1491             | 5723            | 7214  | 0.04              | 0.01              | 0.05 | 0.07              | 0.12              | 0.19 |
| + UVR  | 767 (−28%)        | 16 (−41%)         | 783 (−29%) | 1491             | 5723            | 7214  | 0.05              | 0.02              | 0.06 | 0.05              | 0.10              | 0.19 |

**Discussion**

**Co-exposure to UVR considerably enhanced the negative impacts of the water-soluble fractions of two petroleum fuels on the settlement success of coral larvae.** This phototoxicity was evident under UVR intensities equal to or lower than expected exposures in shallow reef habitats, and the toxic thresholds ($\text{EC}_{50}$) for dissolved aromatics occurred at concentrations below those detected after major accidental spills or releases. These results indicate that by ignoring phototoxicity, the hazards posed by oil spills to coral larvae may be substantially underestimated in shallow-water tropical reef systems.

Coral larval settlement was very sensitive to HFO WAF with threshold concentrations ($\text{EC}_{10}$ and $\text{EC}_{50}$) lower than or similar to previously reported concentrations during and after oil spills (42–189 μg l$^{-1}$ $\sum\text{PAH}$ and 22 μg l$^{-1}$ TPAH). The $\text{EC}_{10}$ and $\text{EC}_{50}$ values (24 and 96 μg l$^{-1}$ TPAH, respectively; −UVR) were also lower than those reported for inhibition of larval settlement in the same species in 24 h exposures to natural gas condensate in the absence of UVR (103 and 339 μg l$^{-1}$ TPAH, respectively)\textsuperscript{29}. Differences in sensitivity are likely to be due to the longer duration of the current exposure and the higher proportion of more toxic PAHs in the HFO WAF. Conversely, the sensitivity of A. tenuis larvae to diesel WAF in the absence of UVR was less than that reported for either HFO or condensate, potentially due to the lower concentrations of total dissolved PAHs (14 μg l$^{-1}$) in the undiluted diesel WAF compared to HFO (440 μg l$^{-1}$; Table 2) and condensate (107 μg l$^{-1}$)\textsuperscript{29} WAFs. The sensitivity of coral larvae observed here is difficult to compare with other studies, which reported different exposure types, endpoints and measures of petroleum hydrocarbon concentration in WAFs (summarised in Negri, et al.\textsuperscript{29} and Turner, et al.\textsuperscript{7}). Nevertheless, studies exposing coral larvae to hydrocarbons in the absence of UVR reported negative effects on coral settlement at THC concentrations between 82 and 620 μg l$^{-1}$\textsuperscript{17,33,46} and the sensitivity of coral larvae is likely to be greater than that of other life stages\textsuperscript{2}.
Figure 1. Penetration of ultraviolet radiation (UVR) on Trunk Reef (a) and Esk Reef (b) as well as a comparison (c) of exposure intensity and spectrum of artificial UVR used during settlement toxicity assays and UVR observed in situ on the Great Barrier Reef (GBR, Australia) during spring. Full spectrum measurements of UVR in air and at 0.1, 1, 2, 3, and 3.8 m depth at a mid-shelf (Trunk reef) and inshore (Esk reef) reef on the GBR in October 2016. Total irradiance values calculated using the percentage reductions in light intensity recorded for each wavelength at each depth (in relation to measurements in air) and total irradiance measurements in air on a clear day (cloud coverage <5%). Comparison of UVR intensity and spectrum emitted from fluorescent tubes used in settlement toxicity assays, calculated UVR exposure inside scintillation vials and UVR exposures observed at 1 m depth on the GBR during spring.
The toxicity of both HFO and diesel WAFs doubled (EC50: 95 to 51 and 1300 to 494 µg l−1 TPAH ± UVR, respectively) in the presence of ultraviolet radiation. This observed phototoxicity is consistent with the 2.5-fold increase in toxicity of natural gas condensate WAF in the presence of UVR to the same larval species in 24 h WAF exposures, based on measured TPAH concentrations. This toxicity increase is similar between studies despite differences in UVR exposure profiles, where Negri, et al.29 applied a single, higher intensity UVR dose for a shorter period of time (2 h, summing to 39 W cm−2) compared to this study (see Methods). Overmans et al.18 recently reported that the inhibition of coral larval metamorphosis by the PAH anthracene

![Figure 2](image-url). Photomicrographs of A. tenuis planulae larvae exposed to water accommodated fractions (WAFs) of heavy fuel oil (HFO) in the absence (−UVR) or presence (+UVR) of ultraviolet radiation as well as juvenile polyps (following settlement). Larvae exposed to (a) filtered seawater (0 µg TPAH l−1), (b) approximately 900 µg TPAH l−1 after 48 h of exposure as well as juvenile polyps, attached and unattached larvae treated with (c) FSW (0 µg TPAH l−1), (d) 115 µg TPAH l−1 and (e) approximately 900 µg TPAH l−1 HFO WAF after 48 h of exposure, introduction of settlement inducer and a 24 h settlement period (a total ~72 h after experiment start). All concentrations in µg TPAH l−1. TPAH = total petroleum aromatic hydrocarbons.

![Figure 3](image-url). Concentration-response curves for coral larval settlement following exposure to heavy fuel oil (a and c) and diesel (b and d) water accommodated fractions (WAF) in the presence (blue) and absence (green) of ultraviolet radiation (µg TPAH l−1). Model mean (solid line) and 95% confidence intervals (shaded area) for quasibinomial GLMs fitted for the settlement success data of each treatment combination as well as observed settlement success for each replicate (open ring) used in model fitting. All concentrations in µg l−1. n_{HFO−/−UVR} = 63, n_{diesel−UVR} = 65, n_{diesel+UVR} = 64. TPAH = total petroleum aromatic hydrocarbons.
increased 7-fold in the presence of UVR (EC_{50}: 45 – UVR to 6.3 + UVR µg l^{-1}), while there was little influence of UVR exposure on the toxicity of phenanthrene. The impacts of PAH and UVR on larval metamorphosis were also detected at lower PAH concentrations than other sub-lethal biomarkers investigated following the 48 h exposures (including 10 h UVR exposure per day)\textsuperscript{18}. Peachey and Crosby\textsuperscript{17} observed 100% mortality of UVR exposure on the toxicity of phenanthrene. The impacts of PAH and UVR on larval metamorphosis were also shown for each treatment combination. 95% confidence intervals in brackets where applicable. All concentration values in µg TPAH l^{-1}. TPAH = total petroleum aromatic hydrocarbons. *Extrapolated from fitted model.

### Table 3. Concentrations of TPAH in fuel WAFs which inhibited 50% (EC_{50}) and 10% (EC_{10}) of larval settlement in the absence (− UVR) and presence (+ UVR) of ultraviolet radiation. Effect concentrations calculated from settlement success data fitted with quasibinomial GLMs with approximate r^2, deviance and degrees of freedom (df) shown for each treatment combination. 95% confidence intervals in brackets where applicable. All concentration values in µg TPAH l^{-1}.

|            | EC_{50} | EC_{10} | Approximate r^2 | Deviance df |
|------------|---------|---------|-----------------|-------------|
| HFO − UVR  | 24 (9–39) | 96 (75–116) | 0.748 | 146.12 | 61 |
| + UVR      | 15 (6–23) | 51 (41–62)  | 0.767 | 137.03 | 61 |
| Diesel − UVR | 302 (51–552) | −1300\textsuperscript{*} (253–2410) | 0.062 | 196.07 | 63 |
| + UVR      | 122 (50–194) | 494 (371–617) | 0.423 | 165.97 | 62 |

The results of this study clearly demonstrate the strong phototoxic effects of the water-soluble fractions of common marine fuels on coral larval settlement in the presence of environmentally realistic UVR exposures. The current study applied UVR intensities slightly lower than the intensities measured at 1 m depth on a mid-shelf (Trunk Reef) and inshore reef (Esk reef) on the Central GBR. These intensities are also consistent with previous measurements on the GBR\textsuperscript{23}, and other coastal and oceanic locations\textsuperscript{22,24}. The penetration of UVR in marine environments is primarily affected by chromophoric dissolved organic matter (CDOM) and, for some systems, particulate matter such as plankton and detritus\textsuperscript{22,23}. Attenuation is therefore likely to be lowest in clear-water oligotrophic coral reef environments, as measured at the mid-shelf site at Trunk Reef, compared to the more turbid waters of the inshore Esk Reef. Barron et al.\textsuperscript{20} found that at depths greater than 0.5 m attenuation of light varies between habitats, emphasising the need to accurately estimate the intensity and wavelengths experienced by each species or ecosystem when investigating the influence of UVR on the toxicity of pollutants. Coral gametes, embryos and larvae developing at the water surface may also be exposed to substantially higher UVR intensities in situ than applied in the present study, potentially reducing the toxic threshold values further. Moreover, these results suggest that the hazard (hence risk) posed by aqueous petroleum hydrocarbons to shallow-water tropical coral reefs will be underestimated if phototoxic activation by UVR is not taken into consideration.

The NTLM for predicting narcotic mortality underestimated the impacts of HFO WAF in comparison to the observed effects on larval settlement. 50% larval mortality was predicted for HFO WAFs by the NTLM at ~900 µg l^{-1} TPAH (TU_{Neat fuel}); however, coral larval settlement was reduced by 50% at far lower concentrations (EC_{50} 96 to 51 µg l^{-1} TPAH ± UVR). It has been reported that the NTLM can underestimate the toxicity of some hydrocarbon mixtures\textsuperscript{14}, and these underestimations may result from the assumption of a simple narcotic mode of action for all aromatic components. The observed discrepancies could partially result from the contribution of other toxic mechanisms, especially those that specifically affect coral metamorphosis from pelagic planula to sessile juveniles, as suggested by Negri et al.\textsuperscript{20}. Furthermore, the sub-lethal endpoint of settlement inhibition (while ecologically relevant\textsuperscript{36}) is by definition more sensitive than mortality, which is typically used to generate the CTLBB values used in the NTLM. Determining CTLBB values for both larvae and adults of key coral reef species would improve species-specific toxicity predictions at different life stages and assist in ranking the potential risks posed by crude oils and petroleum products\textsuperscript{25}. It is also possible that other components (e.g. photoproducts) that were not measured may have influenced the toxicity. Although a phototoxic target lipid model was recently
presented\(^{30}\), its use is currently limited to estimation of the acute toxicity of individual PAHs, and further development is necessary before it can be applied to complex hydrocarbon mixtures or chronic exposures.

Even in well managed and protected areas, groundings of large vessels and petroleum hydrocarbon releases from offshore extraction facilities have occurred in the last decade\(^{41-45}\). Large spills as well as chronic hydrocarbon pollution can lead to degradation of adult coral health and changes to reef composition\(^{46,47}\) with recovery taking more than 10–20 years\(^{5}\). The slow recovery of these reefs is likely to be at least partially due to long term effects of hydrocarbon contamination on recruitment processes, including larval settlement\(^{48}\) used as a sensitive toxicity endpoint in this study. The overlap between a large oil spill and the coral recruitment window can be substantial, with water from some spill sites remaining phototoxic to invertebrate embryos for up to 13 d\(^{49}\) and larvae of some coral species remaining in the water column for up to 3 months (as reviewed by Jones, et al.\(^{30}\)). The increase in toxicity of dissolved aromatics from HFO by UVR exposure resulted in low toxic thresholds, underscoring the potential hazard to corals posed by phototoxic compounds found in petroleum oils and fuels. Previous assessments may therefore have substantially underestimated the risks posed by oil and petroleum product spills on shallow-water, tropical coral reefs by not accounting for interactions with environmental factors such as UVR. Further research into the effects of petroleum hydrocarbons on more tropical reef organisms, including potential interactions with UVR and other stressors, is needed to more effectively quantify these risks.

Methods
Coral collection and larval cultures. Gravid colonies of Acropora tenuis, a reef-building coral common throughout the Pacific Ocean, were collected by hand on SCUBA from Magnetic Island (October 2016, 19.157°S, 146.861°E), GBR, under Great Barrier Reef Marine Park Authority Permit G12/35236.1. Colonies were placed in flow-through seawater of ambient temperature and transported to the National Sea Simulator, Townsville, within 24 hours. On arrival colonies were transferred to 70% shaded flow-through outdoor holding tanks and kept at temperatures equivalent to the collection site (27°C) until spawning. When showing signs of setting colonies were isolated and gametes collected by gentle scooping.

Larval cultures (6 parental colonies, 95% fertilisation) were initiated on the 19th October 2016. Cultures were maintained at 27°C and densities <500 larvae per l at ~27°C\(^{38}\) in round, 500 l fiberglass rearing tanks with cone-shaped bases. Flow-through seawater (1.5 turnovers per day) was filtered seawater\(^{<}\)through cylindrical mesh filter (15 h cm\(^{-2}\)). Flow-through seawater of ambient temperature and transported to the National Sea Simulator, Townsville, within 24 hours. On arrival colonies were transferred to 70% shaded flow-through outdoor holding tanks and kept at temperatures equivalent to the collection site (27°C) until spawning. When showing signs of setting colonies were isolated and gametes collected by gentle scooping.

Preparation of fuel water accommodated fractions. Heavy residual fuel oil (HFO, International Bunker Supplies Pty Ltd, Gladstone, Australia) and automotive diesel (Puma Energy Australia, Fortitude Valley, Australia) WAFs were prepared in capped, solvent rinsed aspirator bottles (5 l) using 0.45 µm filtered seawater (FSW; pH 8.1, salinity 37.0 psu) loaded at 20 g fuel l\(^{-1}\) with a 20% headspace\(^{31}\). Solutions were protected from light, stirred at 50 rpm for 18 h without a vortex and allowed to settle for 30 min prior to use\(^{51}\). Ten dilutions (0, 0.4, 0.8, 1.6, 3.125, 6.25, 12.5, 25, 50 and 100% WAF) were prepared from the fresh undiluted (100%) WAF using 0.45 µm FSW\(^{29,52}\) and used within 3 h in the assays below.

Fuel phototoxicity assays. Acropora spp. larvae reach competency to settle and undergo metamorphosis after ~4 d\(^{40}\). HFO and diesel static non-renewal exposure experiments over 48 h were performed on 7- and 8-d old A. tenuis larvae, respectively. 10–14 larvae and 20 ml of WAF were gently added to 6 replicate glass vials for each WAF concentration and ±UVR treatment combination (12 replicate vials for 0% WAF controls). Vials were tightly capped with approximately 10% headspace\(^{51}\) to allow for oxygen exchange. Vials were randomly placed on each W AF concentration and old W AFs were prepared in capped, solvent rinsed aspirator bottles (5 l) using 0.45 µm filtered seawater (FSW; pH 8.1, salinity 37.0 psu) loaded at 20 g fuel l\(^{-1}\) with a 20% headspace\(^{31}\). Solutions were protected from light, stirred at 50 rpm for 18 h without a vortex and allowed to settle for 30 min prior to use\(^{51}\). Ten dilutions (0, 0.4, 0.8, 1.6, 3.125, 6.25, 12.5, 25, 50 and 100% WAF) were prepared from the fresh undiluted (100%) WAF using 0.45 µm FSW\(^{29,52}\) and used within 3 h in the assays below.

Fuel phototoxicity assays. Acropora spp. larvae reach competency to settle and undergo metamorphosis after ~4 d\(^{40}\). HFO and diesel static non-renewal exposure experiments over 48 h were performed on 7- and 8-d old A. tenuis larvae, respectively. 10–14 larvae and 20 ml of WAF were gently added to 6 replicate glass vials for each WAF concentration and ±UVR treatment combination (12 replicate vials for 0% WAF controls). Vials were tightly capped with approximately 10% headspace\(^{51}\) to allow for oxygen exchange. Vials were randomly placed on their side in trays inside temperature controlled, orbital shaker incubators (Thermoline Scientific, Australia) at 80 rpm to ensure larvae did not settle during the exposure period. One set of vials (−UVR), n = 66, was placed in an incubator fitted with actinic LEDs emitting 1.23 mW cm\(^{-2}\) photosynthetically active radiation (PAR; Aqualina Blue 450 nm, 10,000 K and 420 nm Actinic LED strips, The Aquatic Life Product Company, Willawong, Australia). The second set of vials (n = 66) was placed in a second incubator in the presence of both actinic light and UVR (+UVR). This incubator (+UVR) was fitted with identical actinic LEDs as the first incubator (PAR: 0.95 mW cm\(^{-2}\)) as well as three sets of UV-emitting fluorescent tubes (each set consisting of one DeluxLight Blacklight Blue 18 W and one Reptile One UVB 5.0 18 W T8 fluorescent tube) emitting 0.75 mW cm\(^{-2}\) UVR. Fluorescent tubes emitting predominantly in the UVA spectrum were chosen due to the high UVB attenuation of seawater\(^{49}\) (See Ultraviolet radiation intensities section below on how fluorescent tube irradiance was characterised, and Fig. 1c for comparison of experimental lights, light attenuation of vials used for experimental exposures and in situ intensities of UVR on the GBR). PAR was provided on a 12:12 h L:D cycle and UVR on a 6:18 h L:D cycle (total irradiance 16.1 W cm\(^{-2}\)). An additional 30 vials containing undiluted WAF, but no larvae, were also placed in each incubator to allow for the collection of chemical samples at the end of the 48 h exposure. The positions of vials within each incubator were exchanged randomly throughout each experiment to minimize variation in light exposure. Temperature was continuously logged (Onset HOBO temperature logger, Onset Computer Corp., Massachusetts, USA) while pH, salinity and dissolved oxygen concentrations were measured at the beginning and end of each experiment. At the end of the 48 h exposure approximately 600 ml of undiluted WAF was pooled, from the 30 additional vials (not containing larvae), for chemical analysis (see Chemical analysis below).

Following exposure to fuel WAFs in the presence or absence of UVR, larvae were transferred with 10 ml treatment solution directly to individual wells in 6-well cell culture plates (Nalge Nunc Int., Denmark). Larvae were presented with a settlement inducer consisting of 5 × 5 mm live chips of the crustose coralline algae (CCA) Porolithon onkodes\(^{53}\). Settlement success was assessed as percentage of larvae which attached and flattened into a disc shape\(^{53}\) after ~24 h incubation at 27°C. Average settlement success >70% in controls was considered indicative of a normal response to settlement.
inducers based on several previous studies using CCA or extracts of CCA to initiate settlement\(^{29,33,37,54,55}\). Additional notes regarding deformities, ratio of fully (as outlined by Heyward & Negri\(^{53}\)) to partially metamorphosed recruits as well as failure to attach were also made for each treatment combination.

**Settlement data analysis.** Settlement data was fitted with binomial generalized linear models (GLMs) with a logit link function using the R stats-package (R version 3.4.1\(^{30}\)) to model the settlement success of *A. tenuis* larvae in response to treatment concentration (fixed numerical factor) for each light treatment and fuel type combination. The fitted models were validated by plotting the simulated residuals against fitted values\(^{57}\) and ensuring no individual values influenced the model fits’ disproportionally. Quasibinomial GLMs were fitted where diagnostics indicated overdispersion. An \(r^2\) analogue was calculated using the deviance of the fitted models with and without the fixed numerical factor (equation (1)) to approximate the goodness-of-fit for each model\(^{58}\).

\[
\text{Approximate } r^2 = 1 - \left( \frac{\text{deviance}_{\text{Full model}}}{\text{deviance}_{\text{Null model}}} \right)
\]  

EC\(_{10}\) and EC\(_{50}\) values with 95% confidence intervals (CI) were interpolated from model mean values and 95% CI (adapted from Venables & Ripley\(^{59}\)). Predicted model mean values and 95% CI were exported and graphical outputs produced using GraphPad Prism (version 7.02, GraphPad Software Inc., CA, USA). The high levels of replication used allowed the identification of outliers in the datasets which were excluded. These comprised three HFO FSW controls and one diesel low concentration replicate where CCA chips induced 0% and 7% settlement, respectively, likely due to misidentification of a few of the live CCA chips.

**Chemical analysis.** Samples of freshly prepared, undiluted WAF (“100% WAF”) were collected for chemical analysis at the beginning of each experiment (t\(_0\)). Undiluted WAF was also added to 30 vials per UVR treatment and incubated simultaneously with vials containing larvae to ensure that the undiluted WAF was exposed to the same experimental conditions as the test solutions. At the end of the exposure period (t\(_{48\text{h}}\)), the undiluted WAF, and incubated simultaneously with vials containing larvae to ensure that the undiluted WAF was exposed to the same experimental conditions as the test solutions. At the end of the exposure period (t\(_{48\text{h}}\)), the undiluted WAF, in vials containing no larvae, was pooled and sampled for chemical analysis. For BTEX analysis, samples (40 ml) were collected in amber glass vials with open hole caps and PTFE septa. For all other analyses, single samples (500 ml) were collected in amber glass bottles with PTFE-lined caps. All samples were acidified to pH 2 using 6 M hydrochloric acid and stored at 4 °C until shipped to ChemCentre (Perth, Australia) for analysis as previously described by Negri et al.\(^{39}\). Briefly, WAFs were analysed directly for BTEX using Purge and Trap GC-MS in full scan mode (USEPA method 8260). The 500 ml WAF samples were extracted three times with dichloromethane (DCM) and the combined extracts analysed for PAH and alkylated PAH using GC-MS in SIM or scan mode, and TRH using GC-FID (USEPA method 8270). Neat HFO and diesel were diluted in DCM and analysed for TRH, BTEX, PAH/alkylated PAH and phenols using the same methodology, except additional surrogate (2-fluorophenol, phenol-d5 and 2,4,6-tribromophenol) and internal (1,4-dichlorobenzene-d4) standards were added to the samples prior to analysis. To profile their general composition, the neat oils (1 μl) were subjected to whole oil analysis using GC-MS and hydrocarbons were identified through comparison with a pre-characterised reference oil.

**Ultraviolet radiation intensities.** *In situ* UVR irradiance on the Central GBR during spring was assessed through full spectrum measurements of UVR in air and at five depths at two reefs. Esk Reef (18.775°S, 146.522°E), located in the Palm Islands-group (QLD), was selected as a representative inshore reef site while Trunk Reef (18.329°S, 146.846°E) was selected as a representative clear-water, mid-shelf reef site. Three replicate light intensity measurements for wavelengths between 300–400 nm were performed on SCUBA using a Jaz handheld spectrometer (Ocean Optics, Inc., Florida, USA) and a 5 m fibre optic cable (QP600-025-UV, Ocean Optics, Inc., Florida, USA) aimed directly towards the sun. Z10% values at 305 and 340 nm were estimated by calculating the irradiance corresponding to 10% of surface irradiance (in air) for each reef site and the negative linear relationship between irradiance and measurement depth of measurements made at Trunk Reef and Esk Reef.

Full spectrum measurements of radiation emitted by the UVR fluorescent tubes (Deluxlite Black Light Blue 18W; Reptile One UVB 5.0 18W), used in + UVR larval settlement assays, were performed using the same calibrated Jaz spectrometer and 250 mm UVR compatible fibre optic cable used to quantify the UV radiation of natural sunlight. Full spectrum measurements were made at approximately the same distance as sample vials during experimental exposures (170 mm) in five separate positions relative to the three sets of fluorescent tubes. Total UVA and B radiation measurements were also performed (Solarmeter Model 5.0 UVA + UBV meter, Solartech Inc., Pennsylvania, USA) for the five replicates. Additionally, the attenuation of UVR, between 300–400 nm, emitted from fluorescent tubes by scintillation vial glass was estimated (calibrated Jaz spectrometer). Measurements
were made 200 mm from UVR fluorescent tubes (Deluxlite Black Light Blue 18W; Reptile One UVB 5.0 18W) through the base of a 20 ml scintillation vial. The average total UVA and UBV exposure of larvae inside scintillation vials was calculated using measurements performed with the Solarmeter model 5.0 and the average attenuation of scintillation vial glass between 300–400 nm.

Toxic unit calculations for narcotic toxicity. The toxicity of aromatics to aquatic organisms is depend-
ent on the partitioning of dissolved compounds between water and lipids. The narcotic toxicity of hydrocarbon mixtures can be estimated using the narcotic target lipid model (NTLM) which combines the octanol-water partitioning coefficients (K_{ow}) of all dissolved components in the solution, their measured (or expected) aqueous concentrations and the critical lipid body burden (CTLBB) of the organism of interest (where the CTLBB is the amount of the compound dissolved in the organism's tissues which causes a specific toxic effect; e.g. 50% mortality)\(^\text{14}\). The NTLM is useful for assessing the relative risks posed by different oil and fuel types to aquatic organisms, but may be less useful if the CTLBB is unknown or may underestimate toxicity if other toxic modes of action are important for a given species\(^\text{80}\).

The dissolved composition of petroleum hydrocarbons in a WAF can be measured (Table S-2, Supplementary information) or modelled by applying an oil solubility model (e.g. PETROTOX) to an oil of known composition\(^\text{99}\). Toxic units (TU) for each constituent in a WAF are defined as the ratio between the concentration (C\(_{WAF,i}\)) and the 50% critical effect level (LC\(_{50,i}\) or EC\(_{50,i}\)) of each constituent (i) in a solution, and can be used to enable comparisons of studies using differing experimental designs and hydrocarbon compositions\(^\text{97}\). Assuming the same mode of action (narcosis) for all MAHs and PAHs, the TUs for each component of a WAF containing a mixture of PAHs and MAHs, are considered additive and can be summed to estimate the TU of the WAF\(^\text{95}\) according to equation (2).

\[
TU_{WAF} = \sum \frac{C_{WAF,i}}{EC_{50,i}} + \frac{C_{WAF,2}}{EC_{50,2}} + \frac{C_{WAF,3}}{EC_{50,3}} + \cdots + \frac{C_{WAF,n}}{EC_{50,n}}
\] (2)

A TU of 1 indicates that the mixture is predicted to be toxic and will affect 50% of exposed organisms\(^\text{92}\). In this study, expected TUs (TU\(_{Neat\ fuel}\)) were calculated as per Redman and Parkerton\(^\text{45}\), by applying the observed concentrations of individual compounds in neat HFO and diesel and the average LC\(_{50}\) CTLBB (86.8 \(\mu\)mol g\(^{-1}\) octanol) of 15 marine and estuarine organisms\(^\text{42,61}\). An average marine CTLBB was used as no CTLBB currently exist for acroporid corals. Additionally, observed TUs (TU\(_{WAF}\)) for each light treatment were calculated using measured concentrations of individual compounds in undiluted HFO and diesel WAFs and the same CTLBB value.

Data availability. Data are available upon request.

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Author Contributions
F.M.N. and A.P.N. designed the experiment with input from F.F., D.L.B. and S.A. F.M.N., F.F., D.L.B. and A.P.N. performed experiments. F.M.N. analysed the data. F.M.N. wrote the manuscript with input from A.P.N. as well as F.F., D.L.B. and S.A.

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