Antagonistic interactions between benzo[a]pyrene and C60 in toxicological response of Marine Mussels

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Abstract:

This study aimed to assess the ecotoxicological effects of the interaction of fullerene (C60) and benzo[a]pyrene (B[a]P) on the marine mussel, Mytilus galloprovincialis. The uptake of nC60, B[a]P and mixtures of nC60 and B[a]P into tissues was confirmed by GC-MS, LC-HRMS and ICP-MS. Biomarkers of DNA damage as well as proteomics analysis were applied to unravel the toxic effect of B[a]P and C60. Antagonistic responses were observed at the genotoxic and proteomic level. Differentially expressed proteins (DEPs) were only identified in the B[a]P single exposure and the B[a]P mixture exposure groups containing 1 mg/L of C60, the majority of which were down-regulated (~52%). No DEPs were identified at any of the concentrations of nC60 (p < 0.05, 1% FDR). Using DEPs identified at a threshold of (p < 0.05; B[a]P and B[a]P mixture with nC60), gene ontology (GO) and Kyoto encyclopedia of genes and genomes (KEGG) pathway analysis indicated that these proteins were enriched with a broad spectrum of biological processes and pathways, including those broadly associated with protein processing, cellular processes and environmental information processing. Among those significantly enriched pathways, the ribosome was consistently the top enriched term irrespective of treatment or concentration and plays an important role in the ribosome.
role as the site of biological protein synthesis and translation. Our results demonstrate the complex multi-modal response to environmental stressors in M. galloprovincialis.

**Keywords**: Trojan Horse effect, B[a]P, nC60, co-exposure, Mussels, DNA damage, proteomics
1. Introduction

There have been concerns regarding the potential for manufactured nanomaterials to cause unpredictable environmental health or hazard impacts, including deleterious effects across differing organismal levels, for over a decade. Despite numerous years of study, it is still unclear at what quantity manufactured nanomaterials can be found in the aquatic environment, along with their fate, potential bioavailability and subsequent hazardous effects to biological systems. This is surprising given the growing concern in the field of aquatic toxicology regarding their availability and potential toxicity [1]. Fullerenes are the smallest known stable carbon nanostructures and lie on the boundary between molecules and nanomaterials, with fullerenes generally exhibiting strong hydrophobicity in aqueous media [2]. Buckminsterfullerene (C60) is the only readily soluble carbon nanostructure, although graphene is dispersible in specific organic solvents [3]. Non-functionalised C60 possesses a measurable, but extremely low solubility in water (1.3 × 10^-11 ng/mL), but can exist in the aqueous phase as aggregates (nC60) [4] and is quantifiable in aqueous environmental samples [5]. nC60 can be formed in water when fullerenes are released into the aquatic environment, increasing the transport and potential risk of this nanomaterials to the ecosystem ecology.

The toxicity associated with C60 is controversial and largely unclear [6]. The ability of C60 to both produce and quench reactive oxygen species (ROS) has recently been recognised as a particularly important property in the interaction of fullerenes with biological systems [7], with many aquatic studies demonstrating that fullerenes are capable of eliciting toxicity via oxidative stress [8–10]. Numerous studies have investigated the beneficial and toxicological effects of fullerenes [11–17]. However, the toxicity of nanomaterials has been shown to be dependent on numerous factors, including surface area, chemical composition and shape [18,19]. In specific cases, such as aqueous fullerenes (nC60), the physiochemical structure is influenced by different preparation methods [15,20,21]. Altered physiochemical properties induced through the different methods of solubilisation have been shown to profoundly influence the observed toxicological effects of fullerene exposure, thus making a consensus assessment of environmental toxicity difficult [20]. While the environmental toxicity of fullerenes is still being investigated, an emerging concern is whether fullerene aggregates can act as contaminant carriers (Trojan Horse effects) in aquatic systems, and whether this confirms the reduction or enhancement of toxicity with these compounds. Current evidence suggests a mixture of effects dependent on chemical properties. Under combined aquatic exposure conditions (viz. nC60 and contaminant), it has been demonstrated that 17α-ethinylestradiol (EE2) has a decreased bioavailability [14], altered toxicity [11,22] and localised increases in mercury bioavailability [23]. Finally, when compared to other anthropogenic contaminants, Velzeboer et al. established that the absorption of polychlorinated biphenyls (PCBs) to nC60 was 3–4 orders of magnitude stronger than to organic matter and polyethylene [24]. This enhanced absorption and modifications to toxicity responses may have significant impacts on the fate, transport and bioavailability of co-contaminants already in the aquatic environment. However, more research is necessary to establish which co-contaminants bioavailability is impacted when co-exposed with nC60.

The aquatic environment is often the ultimate recipient of an increasing range of anthropogenic contaminants, and likely in all probable combinations. Organisms which are exposed to complex mixtures of differing compounds and substances can interact in many ways to induce biological responses be it additively, synergistically or antagonistically. These interactions can and do change the organismal response compared with single compound exposures [2,25,26]. Bivalves have highly developed processes for the cellular internalization of nano- and microscale particles (viz. endocytosis and phagocytosis) that are integral to key physiological functions such as cellular immunity [27]. These organisms are also useful bio-indicators because as suspension feeders they filter large volumes of water which facilitates uptake and bio-concentration of toxic chemicals [28], in addition to microalgae, bacteria, sediments, particulates and natural nanoparticles. This high filtration rate has been shown to be associated with the high potential accumulation of different chemicals in their tissues. A variety of mussel species have been used to elucidate both physiological and molecular mechanisms of action to nanoparticles [29,30] making them an ideal model to investigate how organisms respond to environmental stressors such as chemical mixtures [27].
study aims to evaluate the interactions between nC\textsubscript{60} aggregates and the carcinogen benzo[a]pyrene (B[a]P) using marine mussels. A set of biomarkers or biological responses including proteomic analysis were employed to better understand cellular response to single (viz. B[a]P and C\textsubscript{60} fullerene) and combined exposures.

2. Materials and Methods

2.1. Animal collection and husbandry

Mussels (M. galloprovincialis; 45-50 mm) were collected from the intertidal zone at Trebarwith Strand, Cornwall, UK (50° 38’ 40” N, 4° 45’ 44” S) in October 2016. The site has previously been used as a reference location for ecotoxicological studies and is considered relatively clean with minimum presence of disease [31,32]. Following collection, mussels were transported to the laboratory in cool boxes and placed in an aerated tank at a ratio of 1 mussel L\textsuperscript{-1} with natural seawater from Plymouth Sound (filtered at 10 µm). Mussels were maintained at 15°C, fed with micro-algae (Isochrysis galbana, Interpret, UK) every 2 days with a 100 % water change 2 h post feeding.

2.2. Preparation of stock solutions

2.2.1. Fullerenes (C\textsubscript{60})

C\textsubscript{60} and Er\textsubscript{3}N@C\textsubscript{80} were obtained from Sigma Aldrich UK and Designer Carbon Materials Ltd., respectively. In order to better replicate the conditions of the experiment during analysis, 2 mussels were maintained in 2-L glass beakers for 24 hrs with natural seawater from Plymouth Sound (filtered at 10 µm). Subsequently, fullerenes (1 mg) were added to the mussel-exposed seawater (10 mL) and the suspension homogenised by ultrasonication (Langford Sonomatic 375, 40 kHz) for 1 hr at room temperature. The suspension was allowed to settle for at least 4 hrs at room temperature prior to analysis of the aggregate size. Dynamic light scattering (DLS) was performed using a Malvern Zetasizer Nano-ZS at room temperature. Quoted values are the average of 3 measurements. Bright field transmission electron microscopy (TEM) and dark-field scanning transmission electron microscopy (STEM) were performed using the JOEL 2100+ microscope operated at 200 keV. Energy dispersive X-ray (EDX) spectra were acquired using an Oxford Instruments INCA X-ray microanalysis system and processed using Aztec software. Samples were prepared by casting several drops of the respective suspensions onto copper grid-mounted lacey carbon films.

2.2.2. Benzo[a]pyrene (B[a]P)

B[a]P (Sigma Aldrich UK) is not water soluble and was previously dissolved in dimethyl sulfoxide (DMSO) after having determined its solubility limit. Chemical solutions were prepared so that the DMSO concentration in the sea water was 0.001%.

2.3. In vivo exposure of M. galloprovincialis to B[a]P and C\textsubscript{60}: Experimental design

Following depuration, mussels were separated (2 per beaker) into 2 L glass beakers containing 1.8 L of seawater and allowed to acclimatize for 48 h. A photoperiod of 12 h light: 12 h dark was maintained throughout the experiment. Oxygenation was provided by a bubbling system. Seawater was monitored in each of the beakers by measuring salinity (36.45 ± 0.19‰). Mussels were exposed for 3 days with no water changes to B[a]P (5, 50 and 100 µg/L), C\textsubscript{60} alone (0.01, 0.1 and 1 mg/L) and a combination of B[a]P (5, 50 and 100 µg/) and C\textsubscript{60} (1 mg/L). Control groups received only DMSO at the same concentrations as used in the other exposure groups (0.001 % DMSO). A total of 26 individuals were used per treatment. Following exposure, tissue samples were collected as follows: gill and digestive gland (DG) tissue was collected from 3 mussels for chemical analysis, digestive tissue was collected from 9 mussels and pooled (3 mussels per one biological replicate) for shotgun proteomics, DG tissue from 10 mussels was collected for comet assay and DNA adducts, with a further 5 DG
collected for DNA oxidation. Water samples from 3 beakers were randomly collected during each
treatment for B[a]P and C60 analyses.

2.4. GC-MS analyses of B[a]P in water and tissue

Water and tissue extracts were analysed using an Agilent Technologies 7890A Gas
Chromatography (GC) system interfaced with an Agilent 5975 series Mass Selective (MS) detector as
described in [33].

2.5. Analyses of C60 in water and tissue

The analyses of C60 were performed on the toluene extracts common to the B[a]P analyses. The
water extracts were analysed with an Agilent 1100 high-performance liquid chromatography–
ultraviolet-visible instrument (HPLC-UV). The separation was performed on a Shimadzu XR-ODS
column (particle size 2.2 µm, 3.0 × 50 mm) using an acetonitrile–toluene gradient starting at 40%
toluene, at a flow rate of 1 mL/min and a column temperature set at 40 °C. The detection wavelength
was set at 330 nm and the fullerene absorption at maximum. Quantification was performed by
external calibration using authentic fullerene standards. Because of their lower concentrations, the
tissue extracts were analysed by ultrahigh performance liquid chromatography coupled with high
resolution mass spectrometry following a protocol adapted from [34].

2.6. Proteomics

2.6.1. Sample collection and quality check

Tissue was removed from the -80 °C, weighed (100 mg) and twice washed in PBS prior to being
homogenised on ice for 60 s in RIPA buffer. The lysed homogenate was centrifuged at 14,000 RPM
for 60 min at 4 °C, the supernatant collected and aliquoted. Protein concentration was determined
using the Pierce BCA protein assay kit (Thermo Scientific) according to manufacturers instructions
with bovine serum albumin as standard. Reproducibility of protein extraction was carried out using
SDS-PAGE. Briefly, 100 µg of protein from each sample was loaded on a polyacrylamide gradient gel
(4-12 %) and stained with Coomassie protein stain (Expedeon, UK) and destained with ELGA water.
Quality checked protein samples were then processed for downstream LC-MS analysis.

2.6.2. Sample preparation for LC-MS

Equal amounts of intestinal protein (100 µg) were processed using the Filter Aided Sample
Preparation (FASP) method as described by [35]. The digested proteins were subsequently purified
using the STAGE tip procedure as previously described [36]. Tryptic peptides were analysed using
liquid chromatography-mass spectrometry (LC-MS).

2.6.3. Mass spectrometry

Peptides were separated on a Dionex Ultimate 3000 RSFC nano flow system (Dionex, Camberly,
UK) and analysed as described in [37].

2.6.4. Analysis

Peptide identification and quantification. Data analysis and quantification was performed using R
(Version 3.5.0) [38]. Thermo .raw files were imported into ProteoWizard [39] and converted to .mzML
format before identification using the MS-GF+ algorithm which is implemented in R via the
MSGFplus package [40]. MS-GF+ was chosen due to its known sensitivity in identifying more
peptides than most other database search tools and its ability to work well with diverse types of
spectra, configurations of instruments and experimental protocols [41]. The protein database utilised
in this study consisted of the UniProt KnowledgeBase (KB) sequences from all organisms from the
taxa Mollusca, sub category Bivalvia (84,410 sequences released 1/10/2018). This was cocatenated
with a common contaminants list downloaded from ftp://ftp.thegpm.org/fasta/cRAP (Version:
January 30th, 2015) using the R package seqRFLP [42]. Searches were carried out using the following criteria: mass tolerance of 10 ppm, trypsin as the proteolytic enzyme, maximum number of cleavage sites = 2 and cysteine carbamidomethylation and oxidation as a fixed modification. Target decoy approach (TDA) was applied as it is the dominant strategy for false discovery rate (FDR) estimation in mass-spectrometry-based proteomics [43]. A 0.1 % peptide FDR threshold was applied in accordance with standard practice, with a 1 % protein FDR applied after protein identification (via aggregation). The resulting .mzid files were converted to MSnSet and quantified using label free spectral counts. The mass spectrometry proteomics data have been deposited to the ProteomeXchange Consortium [44] via the PRIDE [45] partner repository with the dataset identifier PXD013805 and 10.6019/PXD013805.

Data processing and quantification. Data processing was undertaken as follows: each sample was run individually and then regionally combined before all samples were amalgamated into a large dataset. Quantification of proteins occurred via spectral index (SI) [46]. For identification of proteins, the common practice of requiring three peptides per protein was used in order to reduce the number of false positives [47]. Peptides were subsequently aggregated using sum and the protein intensities scaled based on the actual number of proteins summed. Mussel samples were grouped based on biological replicate, exposure and concentration and the resulting data filtered to keep proteins which were identified in more than two biological replicates. To quantitatively describe reliable and biologically relevant protein expression changes based on single exposure to B[a]P, C60 or to a combination of the two, the data analysis was split into three distinct sections. As per recent recommendations, normalisation was carried out first [48]. Based on systematic evaluations of normalisation methods in label free proteomics, normalisation between technical replicates was carried out using variance stabilization normalisation (Vsn) [49]. Based on a study by Lazar et al. [48], it was hypothesised the most likely cause of missing values will be due to a mixture of MAR (missing at random), MCAR (missing completely at random) and MNAR (missing not at random) data. As such, missing value imputation was carried out using a mixed methodology in the form of KNN (K nearest neighbours, biological replicates) and QRILC (left censor method for MNAR data; whole dataset) [50,51]. Following normalisation, differential expression was carried out using msmsTests [52] with p value less than 0.05 considered significant and Q-values (FDR: <1%) calculated for p-value target matches with the Benjamini-Hochberg procedure. Enrichment of function among up- or down-regulated proteins was calculated using GOfuncR using gene ontologies associated with differentially expressed proteins (P-adj = 0.01, calculated using Benjamini-Hochberg method and q-value = 0.05). KEGG analysis was carried out on the identified unique proteins per treatment (p<0.05) using the clusterProfiler package [53]. KEGG annotation was performed using GhostKOALA [54] and pathways with significant enrichment identified using ClusterProfiler (hypergeometric test, q<0.05 following Benjamini correction). Unique and common proteins based on toxicant were graphically represented through Venn diagrams with the software Venny (http://bioinfogp.cnb.csic.es/tools/venny/index.html) [55]. The R script outlining project analysis for this study can be found in supplementary materials (R script S1).

2.7. DNA damage

2.7.1. Measurement of 8-oxodGuo levels using HPLC/UV–ECD

DNA extraction was performed using 20 mg of digestive gland tissue according to the chaotropic NaI method derived from Helbock et al. [56], slightly modified by Akcha et al. [57]. 8-oxodGuo levels were determined by HPLC (Agilent 1200 series) coupled to electrochemical (Coulochem III, ESA) and UV (Agilent 1200 series) detection as described in [58].

2.7.2. Comet Assay

The comet assay on digestive gland tissue was performed as previously described in [33].
2.7.3. DNA adducts

For each sample, DNA from gills and DG tissues was isolated using a standard phenol-chloroform extraction procedure. We used the nuclease P1 enrichment version of the thin-layer chromatography (TLC) $^{32}$P-postlabelling assay [59] to detect BaP-derived DNA adducts (i.e. 10-(deoxyguanosin-N$^2$-yl)7,8,9-trihydroxy-7,8,9,10-tetrahydro-BaP [dG-N$^2$-BPDE]). The procedure was essentially preformed as described [59]. After chromatography, TLC sheets were scanned using a Packard Instant Imager (Dowers Grove, IL, USA) and DNA adduct levels (RAL, relative adduct labelling) were calculated as reported [60]. An external BPDE-modified DNA standard was used as a positive control [61].

2.8. Confirmation of uptake of fullerenes by mussels

2.8.1. Experimental design

Mussels were exposed to a single treatment, 1 mg/L Er$_3$N@C$_{80}$ for 3 days (static exposure). For each treatment (control and labelled fullerenes), 2 mussels were exposed into 2 L glass beakers containing 1.8 L of seawater.

2.8.2. Bulk spectroscopic analysis

For the determination of erbium concentration in the digestive gland, 2 mussels per treatment were analysed using an X Series II ICP-MS (Thermo Fisher Scientific Inc., Waltham, MA, USA) with PlasmaLab software (Thermo Fisher Scientific Inc., Waltham, MA, USA) as described in [32].

2.8.3. Mussel sectioning and electron microscopy analysis

Following the exposures detailed above, a small piece (~5 mm$^2$) was dissected out of the centre of the digestive gland and fixed in 2% paraformaldehyde, 2.5% glutaraldehyde, 2.5% NaCl, 2mM CaCl$_2$ in 0.1M PIPES, pH 7.2 for 3h. The tissue was then stored in 2.3 M sucrose (in 0.1M PIPES) until analysis. Two mussels were analysed per treatment. Electron transparent sections for STEM analysis were prepared by cutting ~1 mm$^2$ pieces from the washed whole tissues and sectioning to a thickness of ~180-200 nm at -80 °C using the RMC Products PowerTome with the CR-X cryochamber. The cross-sections were transferred onto copper-grid mounted graphene oxide films using the Tokuyasu technique and imaged in dark field STEM using the JOEL 2100+ microscope operating at 200 keV.

2.9. Statistical analysis

Statistical tests were conducted using R software [62]. Normality and variance homogeneity were evaluated using Lilliefor’s test and Bartlett’s test, respectively. When necessary, raw data were mathematically transformed (Ln) to achieve normality before proceeding with an ANOVA. When significant, a posteriori Tukey test was performed. When data could not be normalized, statistical differences between treatments were tested using the non-parametric Kruskal–Wallis test.

2.9.1. Analysis of interactions

Further analysis of the combined effects of C$_{60}$ and BaP on DNA Damage (based on Comet Assay) was performed by calculating the Interaction Factor (IF) in order to test for evidence of additivity, synergism and antagonism [63–65]:

\[
IF = \left( G_{(C_{60} + BaP)} - C \right) - \left( G_{(C_{60})} - C \right) - \left( G_{(BaP)} - C \right) \\
= G_{(C_{60} + BaP)} - G_{(C_{60})} - G_{(BaP)} + C \quad \text{(Equation 1)}
\]

\[
SEM_{(IF)} = \sqrt{(SEM^2_{(C_{60} + BaP)}) + (SEM^2_{(C_{60})}) + (SEM^2_{(BaP)}) + (SEM^2_{(C)})} \quad \text{(Equation 2)}
\]

Where IF is the interaction factor: negative IF denotes antagonism, positive IF denotes synergism, and zero IF denotes additivity. $G$ is the mean cell pathological reaction to toxicants (BaP,
C₆₀ and BaP + C₆₀), C is the mean cellular response under control conditions. SEM(χ) is the standard error of the mean for group X. Results were expressed as IF, and the 95% confidence limits were derived from the SEM values.

In order to test the mixture IF values against predicted additive values (assumed to have an IF = 0), the predicted additive mean values (A) were calculated:

\[ A = (G_{C₆₀} - C) + (G_{BaP} - C) \]  
(Equation 3)

The Pythagorean theorem method for combining standard errors was used to derive combined standard errors for the predicted mean additive values (A) of C₆₀ and BaP (http://mathbench.org.au/statistical-tests/testing-differences-with-the-t-test/6-combining-sds-for-fun-and-profit/). The standard errors for the three C₆₀ and BaP treatments (predicted additive) were derived using the following equation:

\[ SEM_{(add)} = \sqrt{SEM^2(C₆₀) + SEM^2(BaP) + SEM^2(C)} \]  
(Equation 4)

This enabled the 95% confidence limits to be derived for the predicted additive values. The confidence limits were used to test the predicted additive values having an IF = 0 against the IF values for the mixtures.

3. Results

3.1. Characterization of C₆₀ in seawater

Dynamic light scattering and electron microscopy analysis (Figure S1-3 and Table S1) of C₆₀ dispersed in mussel-exposed seawater (~100 µg/mL) with brief ultrasonication followed by equilibration indicates the formation of stable aggregates measuring 653±87 nm (nC₆₀ where n=2x10⁸) in mean hydrodynamic diameter. No significant change in the size of nC₆₀ aggregates was observed upon addition of B[a]P.

3.2. Concentration and uptake of B[a]P and C₆₀ in seawater and tissue

3.2.1. B[a]P

Regarding analyses of B[a]P in seawater, nominal concentrations were matched to stock concentrations (Table S2). No difference was observed between the presence or absence of C₆₀. As already established, there was a rapid disappearance of B[a]P over time in seawater and B[a]P accumulated preferentially in the digestive gland tissue. The results are consistent with previous observations in our lab group with a corresponding dose/response profile [33]. There are no significant differences in B[a]P bio-accumulation depending on the presence/absence of C₆₀, except in the gills where a significantly higher uptake is observed in the presence of C₆₀ at the highest concentration of B[a]P (Figure 1). In general, high variability would conceal subtle changes.
Figure 1. GC-MS analyses of B[a]P in (a) gills and (b) digestive gland of M. galloprovincialis. Asterisks indicate the statistical differences observed between control and exposed groups. (*) p < 0.05, (**) p < 0.01, (***) p < 0.001.

3.2.2. Fullerenes (C60)

A rapid decline in the concentration of C60 in seawater was observed with time, with no quantifiable amounts after day 1 (Table S3). At t0, the measured water concentrations are in reasonable agreement with the nominal concentrations (427.6 ± 45.3, 63.8 ± 11.9 and 7.3 ± 1.8 µg L⁻¹ for nominal concentrations of 1000, 100 and 10 µg L⁻¹ respectively).

Low but quantifiable amounts of C60 in M. galloprovincialis tissues indicate active uptake, with adsorption on the outside of the tissue ruled out due to external washes with toluene prior to analysis (Figure 2). High variability in C60 concentrations in gills and DG makes difficult to detect difference in accumulation between treatments.

Figure 2. LC-MS analyses of C60 in M. galloprovincialis (a) gills and (b) digestive gland (means ± SE). Data marked with different letters differed significantly (Tukey post-hoc test; p < 0.05). An analytical problem led to the loss of two samples of the gills from mussels exposed to Mix100 explaining the absence of standard error.

3.3. Genotoxicity of B[a]P and C60 in the digestive gland of M. galloprovincialis

3.3.1. DNA strand breaks
B[a]P exposure induced DNA damage in the digestive gland at the intermediate and highest concentrations (50 µg L⁻¹ and 100 µg L⁻¹) after 3 days of exposure (Figure 3). No effect was observed at the lowest concentration. Regarding exposure to C₆₀ only, higher DNA strand breaks compared to the controls were observed only at the highest concentration (1 mg L⁻¹, p < 0.001). Lower C₆₀ concentration did not appear to have any genotoxic effects on mussel digestive gland at the concentrations tested. In mussels exposed to B[a]P + C₆₀, significant higher DNA damage compared to control were observed at all the tested concentrations.

![Image of DNA strand break level following 3 days of exposure to C₆₀, B[a]P and mixture of both in the digestive gland. Asterisks indicate the statistical differences observed between control and exposed groups. (*) p < 0.05, (**) p < 0.01, (***) p < 0.001.](image)

**Figure 3.** DNA strand break level following 3 days of exposure to C₆₀, B[a]P and mixture of both in the digestive gland. Asterisks indicate the statistical differences observed between control and exposed groups. (*) p < 0.05, (**) p < 0.01, (***) p < 0.001.

**Interactions.** Interactions between C₆₀ and B[a]P on DNA damage (Comet assay) are shown in Table 1. There is evidence of an antagonistic interaction between C₆₀ and B[a]P at the highest concentration between C₆₀ and B[a]P (Table 1).

**Table 1.** Analysis of combined effects of B[a]P and C₆₀ on DNA damage based on Interaction Factors (IF)

| Treatments                           | DNA Damage (Comet assay)   |
|--------------------------------------|---------------------------|
| BₐP 5 µg L⁻¹ + C₆₀ 1 mg L⁻¹           | -7.48 ± 6.63              |
| BₐP 50 µg L⁻¹ + C₆₀ 1 mg L⁻¹          | -10.39 ± 3.50             |
| BₐP 100 µg L⁻¹ + C₆₀ 1 mg L⁻¹         | -12.69 ± 6.05*            |

3.3.2. DNA oxidation

A significant increase (p = 0.00108) in 8-oxo-dGuo levels was detected in the digestive gland of mussels exposed to C₆₀ (15.3 ± 2.3) compared to control (5.9 ± 1.3) (Figure 4). Despite a higher level of oxidative DNA damage in other treatments compared to control, no significant difference was observed (p > 0.05).
Figure 4. 8-oxodGuo levels in the digestive gland of mussels. Asterisks indicate the statistical differences observed between control and exposed groups. (*) p < 0.05, (**) p < 0.01, (***) p < 0.001.

3.3.3. DNA adducts

Whatever the exposure concentration of B[a]P and mixture of B[a]P and C₆₀, no DNA adducts were detectable in DNA samples from the digestive gland of *M. galloprovincialis*.

3.4. Proteomics

3.4.1. Identification of differentially expressed proteins

In order to identify differentially expressed proteins in the digestive gland proteome of controls, B[a]P, nC₆₀ and mixture (B[a]P and 1 mg/L nC₆₀), a label free LC-MS/MS approach was used with trypsinised tissue homogenates. Following removal of common contaminants in each dataset, peptide mapping quantified 3125, 3428 and 3475 unique proteins following identification from UNIPROT database distinct to B[a]P, C₆₀ and mixture (B[a]P and 1 mg/L C₆₀) treatments, respectively. Irrespective of treatment, protein sequences from the Pacific oyster *Crassostrea gigas* (Organism ID = 94323) were highly represented in the samples at approximately 38 %, followed by Japanese scallop *Mizuhopecten yessoensis* (Organism ID = 6573) at 34 %. Surprisingly, sequences from the genus *Mytilus* were less represented in the search at approximately 3 % with the Mediterranean mussel *Mytilus galloprovincialis* (Organism ID = 29158) representing approximately 1 % of identified sequences. This may be due to a lack of genomic information available for this genus in the UNIPROT database, even though a genome sequence is available [66].

Differentially expressed proteins (DEPs) was determined using a quasi-likelihood GLM. Comparison of each dose per treatment (B[a]P: 5, 50 and 100 μg/L, nC₆₀: 0.01, 0.1 and 1 mg/L, and a mixture: 5, 50 and 100 μg/L B[a]P and 1 mg/L nC₆₀) with the control group was visualised using Venn diagrams (Figure 5). Minimal overlap between varying concentrations was observed for the mixture treatment (average of 2 %) (Figure 5c) when compared to B[a]P (Figure 5a, 9 %) or nC₆₀ (Figure 5b, 8 %). Volcano plots were used to visualise statistically significant changes in protein abundance for varying concentrations of the above treatments following comparison to controls (Figure 6). Applying a 1 % FDR threshold, 401 differentially expressed proteins were identified following B[a]P treatment (all concentrations) and 297 differentially expressed proteins were identified following treatment with the mixture of B[a]P and nC₆₀. No differentially expressed proteins (p < 0.05) were identified in C₆₀ treated samples. The identified DEPs can be further broken down based on treatment
with 42, 50 and 164 DEPs identified at 5, 50 and 100 µg/L B[a]P. Following exposure to a mixture solution, 95, 108 and 94 DEPs were identified at each concentration respectively (1 mg/L of C60 and 5, 50 and 100 µg/L of B[a]P) with Figure 7 representing a visual comparison of commonalities between single exposure versus combined exposure. A subset of DEP based on the top 3 unique proteins per concentration is displayed in Table 2, with the full list of unique proteins and associated p-value and FDR correction (Spreadsheet S1). The majority of differentially expressed proteins detected in this study (B[a]P and mixture exposure) were down-regulated (52%) between the treatment and control conditions irrespective of concentration.

3.4.2. GO functional enrichment

Gene ontologies were directly annotated using a custom annotation database derived from UNIPROTKB (bivalvia) with enrichment carried out using GOfuncR. This provides a controlled vocabulary to describe gene product characteristics in three independent ontologies viz. biological process, molecular function and cellular components. Based on the R package GOfuncR, 31, 35 and 23 GO nodes were found enriched at a threshold of \( p < 0.05 \) (FWER correction) following treatment with B[a]P, C60 or co-mixtures (5 - 100 µg L\(^{-1}\) B[a]P and 1 mg L\(^{-1}\) C60). The top GO terms are listed in Table 3 (threshold set FWER = 0.01), while the full list separated by treatment and concentration can be found in supplemental material (Spreadsheet S1). Irrespective of treatment, biological process records the majority of enriched terms.

**Figure 5.** Venn diagram visualising the overlap between the control sample and varying concentrations of B[a]P **(a)**, C60 **(b)** or a mixture of the two (5-50-100 µg/L B[a]P, 1 mg/L C60) **(c)** following exposure for 3 days. Note that overlap is based on a threshold of \( p < 0.05 \) and does not include FDR correction.
Table 2. Significantly expressed proteins of B[a]P, C60 and mixture (5-100 µg/L and 1 mg/L C60).
Species id’s are as follows: 6573 = Mizuhopecten yessoensis, 6551 = Mytilus trossulus, 29159 = Crassostrea gigas and 94323 = Crassostrea ariakensis.

| Treatment                  | Species  | Protein Name                              | UNIPROTKB     | GO annotation                                                                 | Regulation |
|----------------------------|----------|-------------------------------------------|---------------|--------------------------------------------------------------------------------|------------|
| B[a]P (5 µg/L)             | 6573     | Arrestin domain-containing protein 3      | A0A210PE39    |                                                                                 | Up         |
|                            | 6573     | Orexin receptor type 2                    | A0A210PSC6    | GO:0004930, GO:0016021                                                          | Up         |
|                            | 6573     | Ran-specific GTPase-activating protein     | A0A210Q6H5    | GO:0005622, GO:0046907                                                          | Up         |
|                            | 6573     | 5-hydroxytryptamine receptor 1A-alpha     | A0A210R4M3    | GO:0004993, GO:0005887, GO:0008283, GO:0042310, GO:0046883, GO:0050795           | Down       |
| B[a]P (50 µg/L)            | 6573     | Adenylate kinase isoenzyme 5              | A0A210QMB2    | GO:0005524, GO:0006139                                                          | Up         |
|                            | 6573     | Uncharacterised protein                    | A0A210Q912    |                                                                                 |            |
| B[a]P (100 µg/L)           | 6573     | Helicase with zinc finger domain 2         | A0A210PQ46    | GO:0004386, GO:0030374                                                          | Up         |
|                            | 29159    | Peroxiredoxin-4                           | K1QLH0        | GO:0005623, GO:0045454, GO:0051920                                               | Up         |
|                            | 29159    | Hypoxia up-regulated protein 1             | K1QBF7        | GO:0005524                                                                      | Up         |
| B[a]P (5 µg) + C60 (1 mg/L)| 94323    | Ras-like GTP-binding protein RHO           | HP9LJA2       | GO:0003924, GO:0005525, GO:005622, GO:0007264                                    | Up         |
|                            | 29159    | Zinc finger CCCH domain-containing protein 13| K1PKC9        | GO:00046872                                                                     | Up         |
| B[a]P (5 µg) + C60 (1 mg/L)| 29159    | Myosin heavy chain, non-muscle (Fragment) | K1QXX7        | GO:0003774, GO:0003779, GO:005524, GO:0016459                                    | Up         |
|                            | 29159    | Ribosomal protein S20                     | A0A077H0N2    | GO:0003723, GO:0003735, GO:005622, GO:0015935                                    | Down       |
| B[a]P (50 µg) + C60 (1 mg/L)| 6573    | Nucleolar and coiled-body phosphoprotein 1 | A0A210Q9W0    | GO:0005730                                                                       | Down       |
| B[a]P (50 µg) + C60 (1 mg/L)| 29159    | Tripartite motif-containing protein 2      | K1QHD4        | GO:0005622, GO:0008270                                                          | Up         |
| B[a]P (100 µg) + C60 (1 mg/L)| 6573    | Ran-specific GTPase-activating protein     | A0A210Q6H5    | GO:0005622, GO:0046907                                                          | Down       |
| B[a]P (100 µg) + C60 (1 mg/L)| 29159    | Uncharacterized protein                    | K1RS43        |                                                                                 |            |
3.4.3. KEGG pathway enrichment

To further analyse the identified proteins per treatment, KEGG pathway analysis was performed. Using the bioconductor package clusterProfiler, protein sequences were assigned to DEPs (p<0.05) and submitted to GhostKoala to obtained KEGG Orthology numbers (KO). In general, 52-56% of entries were successfully annotated with approximately 92% of annotations associated with the mollusca taxonomy. Variation between enrichment was described per treatment and concentration as follows:

**B[a]P**: at 5 µg/L exposure, 52 enriched processes were identified and include ribosome processes (26 genes), thermogenesis (19 genes), protein processing in endoplasmic reticulum (13 genes) and mTOR signalling pathway (9 genes). At 50 µg/L exposure, 38 pathways were enriched and ribosome (26 genes), protein processing in the endoplasmic reticulum (17 genes) and phagosome (13 genes). Finally, at 100 µg/L, 26 enriched processes were identified including ribosome (26 genes), RNA transport (16 genes), protein processing in the endoplasmic reticulum (16 genes), biosynthesis of amino acids (16 genes) and endocytosis (15 genes). The mTOR signalling pathway was not enriched at either 50 or 100 µg/L.
**Figure 6.** Volcano plots representing the differentially expressed proteins with exposure to B[a]P (a), C₆₀ (b) or a mixture of the two (5-50-100 µg/L B[a]P 1 mg/L C₆₀) (c). Red dots represent DEPs ($p < 0.05$) with an FDR of 5% while blue dots represent DEPs with an FDR of 1%. Black dots represent unique proteins which are not differentially expressed.
Figure 7. Venn diagram visualising the overlap between 5 µg/L (a), 50 µg/L (b) and 100 µg/L (c) of B[a]P with a mixture solution containing the same B[a]P concentrations in addition to 1 mg/L of C60 following 24 h exposure. Overlap is based on $p < 0.05$ and FDR set at 1%.

Table 3. Subset of enriched Gene Ontology (GO) terms with an family wise error (FWER) threshold of 1% (or 0.01) following B[a]P (5-100 µg/L), C60 (0.01 – 1 mg/L) and a mixture of B[a]P (5-100 µg/L) and C60 (1 mg/L) treatments. Cellular component and biological processes are abbreviated to CC and BP respectively.

| Treatment | Ontology | GO-ID | GO-ID Name | FWER |
|-----------|----------|-------|------------|------|
| B[a]P (100 µg) | BP | GO:0006139 | Nucleobase-containing compound metabolic process | 0.01 |
| B[a]P (100 µg) | BP | GO:0006725 | Cellular aromatic compound metabolic process | 0.01 |
| B[a]P (100 µg) | BP | GO:0034641 | Cellular nitrogen compound metabolic process | 0.01 |
| B[a]P (100 µg) | BP | GO:0046483 | Heterocycle metabolic process | 0.01 |
| B[a]P (100 µg) | BP | GO:0090304 | Nucleic acid metabolic process | 0.01 |
| B[a]P (100 µg) | BP | GO:1901360 | Organic cyclic compound metabolic process | 0.01 |
| C60 (0.01 mg/L) | BP | GO:0000226 | Microtubule cytoskeleton organization | 0.01 |
| C60 (0.1 mg/L) | CC | GO:0031974 | Membrane-enclosed lumen | 0.01 |
| C60 (0.1 mg/L) | CC | GO:0043233 | Organelle lumen | 0.01 |
| C60 (0.1 mg/L) | CC | GO:0070013 | Intracellular organelle lumen | 0.01 |

The majority of enriched pathways identified can be grouped under genetic information processing, cellular processes, environmental information processing and metabolism. The top enriched pathways identified per concentration were plotted to identify commonalities and differences between differing concentrations of B[a]P (Figure 8a) based on genes identified in that pathway. Interestingly, unique pathways appear to be activated dependent on exposure concentration, with only the ribosome pathway consistently present and enriched at all
concentrations potentially indicating the high degree of translation which may be occurring as a consequence of PAH exposure.

C_{60}: at 0.01 mg/L exposure, 33 enriched pathways were identified while 12 enriched pathways were identified at 0.1 mg/L exposure and 35 enriched pathways identified at 1 mg/L exposure (p < 0.05, FDR = 5%). The top enriched pathways were illustrated in Figure 8b, with an absence of enrichment of certain pathways dependent on treatment concentration. For example, thermogenesis was only enriched at the highest concentration of 1 mg/L with 12 genes identified in the pathway. The ribosome is the top enriched pathway at all concentrations of C_{60} with 19 genes enriched at 0.01 mg/L exposure, 24 genes enriched at 0.1 mg/L exposure and 35 genes enriched at the highest concentration of 1 mg/L. This is closely followed by protein processing in endoplasmic reticulum, which is broadly comparable in terms of genes between 0.01 mg/L (17 genes), 0.1 mg/L (11 genes) and 1 mg/L (16 genes, Figure 9) exposure. The enriched pathways can be broadly grouped into predominantly genetic information processing, metabolism and cellular processes.
Figure 8. Dotplot of enriched KEGG pathways for DEGs ($p < 0.05$) that were common between concentrations of B[a]P (a), C60 (b) and a mixture of 5, 50 and 100 µg/L with 1 mg/L C60 (c). Along the x-axis, genes represent the number of genes identified as enriched in this particular pathway. The size and colour of each dot represents the gene number and adjustment $p$ based on FDR correction.

**Mixtures:** Under mixture scenario, C60 at a constant concentration of 1 mg/L was mixed with 5 µg/L, 50 µg/L and 100 µg/L of B[a]P resulting in 50, 38 and 54 enriched pathways respectively. At the lower mixture concentration of 5 µg/L B[a]P and C60, the top 3 enriched descriptive terms were related to the ribosome (29 genes), protein processing in endoplasmic reticulum (20 genes) and pathways in cancer (23 genes). At 50 µg/L B[a]P and C60, the top 3 enriched descriptive terms were related to the ribosome (23 genes), carbon metabolism (23 genes) and protein processing in endoplasmic reticulum (19 genes).

Finally, at 100 µg/L B[a]P and C60, the top 3 enriched descriptive terms were related to the ribosome (25 genes), pathways in cancer (23 genes) and MAPK signalling pathway (17 genes). Key genes consistently identified in the protein processing in the endoplasmic reticulum (irrespective of treatment) include Hsp70, Hsp90, TRAP, PDIs and OSTs. At the highest concentration of B[a]P and C60, genes identified in pathways in cancer include GSTs, CASP3 and Wnt. The top pathways based on quantity of genes present in the pathway were presented in Figure 8 with clear trends towards an absence of enrichment in certain pathways based on mixture concentration, e.g. MAPK signalling which is only present at the top exposure concentration combination.

3.5 Analysis of fullerenes (C60) uptake in mussels
To provide further insight into the uptake of fullerenes by marine mussels, it was necessary to use a form labelled with a diagnostic marker. In our experiments, we explored the application of the endohedral fullerene Er3N@C80, fabricated using the trimetallic nitride template (TNT) process, as it represents a good structural analogue to C60, possessing similar surface chemistry, and contains a rare earth element, shielded from the external environment within the fullerene cage, that is not found in nature. The presence of erbium in the mussel digestive gland, as diagnostic of the uptake of labelled fullerenes, was thus quantified using ICP-MS and found with a mean concentration of 151.5 µg/kg (236.5 µg/kg and 66.4 µg/kg for each mussel). However, despite an exhaustive electron microscopy investigation of whole and cross-sectioned DG tissues (Figures S4, Supplementary materials), no direct visualisation of labelled fullerenes was observed.

4. Discussion

Bivalves are ideal organisms for evaluating the adverse effects caused by various environmental stressors including polycyclic aromatic hydrocarbons (PAHs) and nanomaterials. PAHs such as B[a]P have a ubiquitous aquatic distribution and are known to cause several adverse effects in a diverse range of aquatic organisms. While single exposure studies are more common, various bivalve species have already been used as biological models in proteomics to assess the effects of complex mixtures [22,67,68] in addition to other aquatic species [69,70]. Nanomaterials, both as solids and colloids, are ingested by many organisms and bio-accumulate in large quantities, especially in molluscs. The mussel digestive gland is one of the principal detoxification organs with an acknowledged concentration of phase I detoxification enzymes [71]. As such, it is unsurprising that mussel digestive gland has been used as model tissue for eco-toxicological studies of various NPs [72–74], with Di et al. reporting that the digestive gland in *Mytilus edulis* accumulates more C60 than other tissues [72].

There is considerable debate in the literature regarding the actual toxicological impact of nanomaterials in the aquatic environment, with fullerene toxicity controversial. In the aquatic system, Kahrur et al. compiled fullerene toxicological data for fourteen organisms and classified C60 as "very toxic" [75]. Using mouse and human cell lines, Isakovic et al. demonstrated that pristine C60 and aqueous suspensions of C60 are more toxic than its hydroxylated derivatives [76]. In marked contrast, other studies have demonstrated that pristine C60 has low or limited toxicity to cells and various organisms [10,77–79]. The lack of consensus regarding C60 toxicity may be partly due to limited studies which incorporate both a physiological and ecological approach. As a consequence, little is still known about NP bioavailability, mode of uptake, ingestion rates and actual internal concentrations related to Absorption, Distribution, Metabolism and Excretion (ADME) [27]. Despite the contradictory reports, there is consensus that some nanomaterials may potentially affect biological systems directly but also through interactions with other compounds which may be available in the environment (reviewed in [6]). Studies which investigate co-exposure with carbon-based nano-compounds, such as nanotubes or fullerenes are limited, especially in aquatic systems. Using *Danio rerio* (zebrafish) hepatocytes, Ferreira et al. investigated the co-exposure of C60 with B[a]P and provided evidence of toxicological interactions whereby C60 increased the uptake of B[a]P into cells, decreased cell viability and impaired detoxification responses [69]. While, Baun et al. reported that co-exposure with fullerene C60 enhanced toxicity of phenanthrene to *Daphnia magna* and *Pseudokirchneriella subcapitata* [22]. With respect to B[a]P and C60, Di et al demonstrated organ specific response to both single and combined mixtures with no observation of cytotoxicity and duration dependent and condition specific genotoxic response in *M. galloprovincialis* [72]. Importantly, the observed genotoxic response was reversible after a recovery period. In this study, we add to the growing evidence that toxicity associated with C60 may in fact be related to the nanomaterials ability to act as a vector for other contaminants, and in its aqueous form is not inherently toxic itself.

4.1. Chemistry support

4.1.1. Accumulation of B[a]P
Comparable B[a]P tissue concentrations in the presence or absence of C60 indicate that despite the expected strong sorption of B[a]P on C60 [80], C60-sorbed B[a]P remains bioavailable to *M. galloprovincialis*. Changes in the bioavailability of contaminants co-exposed with carbon nanomaterial has been reported, from a decrease in bioavailability [81,82] to its enhancement, also called the “carrier effect” [83,84]. It has been demonstrated that carbon nanopowder helps BaP uptake by zebrafish embryos and very interestingly also affected the distribution of the pollutant in the organism [85]. However, in the same species, in zebrafish larvae it has been shown that bioavailability of 17α-ethynylestradiol (EE2) was reduced with increasing concentration of nC60 nanoparticles [14]. It appears that the bioavailability of nanomaterials and their co-contaminants depends on many factors such as their size, shape, surface coating and aggregation state and on the metabolism of the species investigated [81,86].

4.1.2. Accumulation of C60

In a previous study in *M. galloprovincialis* [87], mussels exposed to C60 alone showed higher accumulation of C60 in the digestive gland compared to the gill. Interestingly, co-exposure to fluoranthene modified accumulation of C60, with higher accumulation of C60 when animals are exposed to C60 alone compared to combined exposure. When comparing water and tissue concentrations for B[a]P and C60, the bioconcentration observed in our conditions was much lower for C60 compared to B[a]P: the uptake in the DG of mussels exposed to a similar aqueous concentration of B[a]P and C60 was about 2000 times more important for B[a]P. However, non-constant concentrations in the aqueous phase, attributed to sorption and/or sedimentation, did not allow the calculation of bioaccumulation factors, which also requires reaching a steady-state in the tissues. The difference between B[a]P and C60 tissue concentration could also be attributed to different kinetics of uptake, which could only be explored through longer exposure periods and regular sampling. Recent work indicated a continuous increase of C60 concentrations in whole mussels over at least 3 weeks [88].

Further confirmation of the accumulation of fullerenes within mussels was afforded by ICP-MS analysis of digestive gland tissues extracted from mussels exposed to Er-labelled fullerenes. However, no evidence for the presence of labelled fullerene aggregates within tissue sections using our novel STEM-EDX approach was afforded, indicating that the fullerenes are likely distributed within the tissues at the near molecular level (i.e. highly dispersed) and therefore below the sensitivity of either microscopy or *in situ* spectroscopy approaches in complex materials such as these.

4.2. DNA damage

Most of the observed DNA damage will result from oxidative injury by ROS generated by futile cycling of BaP, and also produced by C60 and lipofuscin associated iron [89,90]. According to a review by Johnston et al., fullerene toxicity has been suggested to involve oxidant-driven response and suggests evaluating toxicity by including oxidative stress and related consequences including inflammation or genotoxicity [91].

As already observed in [72], mixture did not increase the formation of DNA strand breaks in the digestive gland of *M. galloprovincialis*. Interestingly, the analysis of interactions performed on the comet assay results revealed an antagonistic effect at the highest concentration in the co-exposure treatments. The analysis of oxidative DNA damage through the analysis of 8-oxodGuo confirmed the induction of oxidative damage by C60. The small increase (not significant) of 8-oxodGuo observed for B[a]P treatment could be due to the short exposure time (3 days). In [57], an increase in the level of 8-oxodGuo was observed after 10 days of B[a]P exposure in the digestive gland of *M. galloprovincialis*. In our study, the antagonistic effect observed in the co-exposure treatment at the highest concentration may be caused by a reduction in ROS generation, or more effective scavenging of ROS by C60, when C60 and BaP are present together in close association, as previously described by [9,72]. C60 fullerenes are both scavengers and generators of reactive oxygen species (ROS) [92]; and when C60 and BaP are closely associated or bound together within the lysosomal compartment of the mussel digestive cells, their ROS scavenging and generating properties may be altered.
4.3. Protein expression profiles

Investigations into proteome responses of marine organisms to various stressors is comparatively small when compared to other model laboratory organisms, both aquatic and terrestrial. Proteomic analysis represents a fundamental step in extending understanding of the physiological processes involved in organismal responses to environmental stressors. In addition, proteomics also provides better qualitative data on post-translational modifications without interference from mRNA instability [93]. A major limitation in the field has been the lack of available annotated genomes for a broad diversity of marine organisms. As a consequence, it has been considered a widely under utilised tool [94]. The lack of genome information has not stopped studies on proteome characterisation in bivalvia/mollusca species using broad protein databases limited to either the phylum, class or specific combination of species [95–98]. However, studies investigating proteome response to environmental stressors or injury are less abundant [30,68,99]. In the current study, a label free shot-gun proteomics approach was performed to investigate proteome alterations in the digestive gland of M. galloprovincialis following treatment with B[a]P, C₆₀ and a combination of B[a]P with 1 mg/L of C₆₀. In the identification of proteome changes between conditions and contaminants, a default threshold for fold change was not set a priori in order to appreciate moderate protein changes. Instead, strict statistical criterion for significance was adopted. Whilst thousands of unique proteins were identified in each treatment, differentially expressed proteins were only identified following treatment with B[a]P and B[a]P in combination with C₆₀. Focusing on B[a]P exposure, proteomic analysis of mussel digestive gland revealed significant differences at all exposure concentrations when compared to the control. Statistically, 42, 195 and 164 proteins were differentially expressed after a 3-day exposure at 5, 50 and 100 µg/L of B[a]P (1% FDR, p < 0.05). In comparison, no differentially expressed proteins were identified following a 3 day C₆₀ exposure. However, when B[a]P was co-exposed to C₆₀ (1 mg/L), differentially expressed proteins at the two highest concentrations in the mixture exposure decreased when compared to single exposure (Figure 7). This trend towards higher protein alterations in single exposures versus co-exposures suggests a non-additive combine effect and is in agreement with prior studies which reported generally higher protein alterations of B[a]P and Cu under single exposure then when co-exposed together [68]. The data in this study suggests that an interaction occurs between B[a]P and C₆₀ whereby the effect of the mixture is different from the presumption of additivity (were by dose response relationships of mixtures are enhanced in comparison to the individual components) as outlined in Rosa et al [100]. In this case, the data suggests an antagonistic relationship between B[a]P and C₆₀ at the higher concentrations of 50 and 100 µg/L. This observation has previously been observed in Mytilus edulis digestive gland [72]. However, this trend is not replicated at the lowest concentration of 5 µg/L whereby mixture exposure resulted in higher DEPs than single exposure. This difference in DEPs may potentially be related to reduced accumulation of B[a]P at the higher concentrations due to saturation of mussel tissue and thereby limiting protein changes. In previous studies, increased impact and accumulation of B[a]P at lower concentrations in M. galloprovincialis has been attributed to tissue saturation [101]. The increase in differentially expressed proteins at the lower concentration may also reflect the inability of membrane transporters such as p-glycoprotein to efflux this particular nanoparticle [102] and as such acts to bypass typical protective mechanisms initiated to protect the organism from PAH stress.

The ability of a stress organism to adjust its cellular processes via transcriptional and subsequently proteomic processes allows it where possible to minimise cellular damage, which may lead to organism death. GO analysis revealed 30 enriched proteins following B[a]P exposure, 42 following C₆₀ exposure and 31 in the mixture exposure. The response of M. galloprovincialis to B[a]P is characterised by a predominant enrichment of Biological processes (67 % or 20 GO’s) with the majority of these occurring at 100 µg/L. When compared to the mixture model at the same concentration, 7 terms are absent in the mixture model compared to the single exposure viz. DNA metabolic processes (GO:0006259), DNA repair (GO:0006281), Cellular response to DNA damage stimulus (GO:0006974), cellular response to stress (GO:0033554), metabolic processes (GO:0008152), cellular metabolic processes (GO:0044237) primary metabolic processes (GO:0044238) and organic...
substance metabolic processes (GO:0071704). The absence of these enriched terms at the highest mixture concentration of B[a]P and C_{60} in association with the reduction in differentially expressed proteins (when compared to single exposure and 50 µg/l) suggest an antagonistic interaction between the two common contaminants. This may be explained by known properties of the chemicals. nC_{60} is an exceptional free radical scavenger [103,104], while B[a]P has been shown to produce free radicals under a variety of conditions [105]. B[a]P contributes approximately 50 % of the total carcinogenic potential of the PAH group [106]. Transcriptomic alterations related to B[a]P are likely to be related to genotoxic mechanisms in addition to other biological processes such as mitochondrial activities and immune response as outlined previously [33]. In contrast, Zhang et al demonstrated that aqueous C_{60} aggregates induced apoptosis of macrophage by changing the mitochondrial membrane potential [107]. As predicted by the literature, enriched GO terms following single nC_{60} exposure are predominantly related to changes to the membrane-enclosed, organelle and intracellular lumen, while mixed exposure resulted in enrichment of mitochondrial components (viz. matrix, ribosome and protein complex). This enrichment of organelle cellular components correlates with enrichment of the ribosome KEGG pathway (ko03010, 35 proteins at 1 mg/L C_{60}), suggesting an increase in the production of newly synthesised organelle proteins which must find its way from site of production in the cytosol to the organelle where it functions. It was not feasible to quantify changes in cellular components in the digestive gland during this study however we can postulate from prior studies that observed changes may be linked to changes in the mitochondria. Mitochondria are essential eukaryotic organelles required for a range of metabolic, signalling and development processes. Using fullerenol, a polyhydroxylated fullerene derivative, Yang et al. demonstrated significant changes to isolated mitochondria via mitochondrial swelling, collapse of membrane potential, decreased of membrane fluidity and alterations to the ultrastructure [108]. The increase in protein production via the ribosome at the highest concentration may reflect the activation of a repair mechanism for damage to this structure. In a recent review, the main negative molecular and cellular responses associated with carbon nanotube (CNTs) in mammals were associated with oxidative stress which can promote inflammation, mitochondrial oxidation and activation of apoptosis [109]. Additionally, Zhang et al. reported on a loss in mitochondrial membrane potential in a mouse in vitro model, in association with increase in cellular ROS suggesting mitochondria associated apoptosis [107]. In a typical aquatic NPs exposure, uptake is followed by localisation into the endosomes, lysosomes and digestive associated cells as well as the lumen of digestive tubules [22,27,110]. This NP exposure response can be followed by disruption or modification to mitochondrial activity [30]. Although the current study would support the hypothesis of mitochondrial damage/repair, further work will need to be carried out to verify.

KEGG pathway analysis can provide physiological pathway information for various experiments with prior studies using it to aid in identification of mode of action of environmental contaminants [68]. In the current study, irrespective of exposure conditions or concentrations, the top enriched pathway identified using KEGG was the Ribosome with 19-39 genes identified in the pathway dependent on treatment and concentration. This was followed by protein processing in the endoplasmic reticulum and carbon metabolism. The ribosome is a large complex molecule made of RNA and proteins that perform the essential task of protein synthesis in the cell. They also serve as the initiation point for several translation-associated functions including protein folding and degradation of defective or nonstop mRNAs. Previous studies have demonstrated a change in regulation of genes which encode ribosomal protein subunits following B[a]P exposure, with the suggestion that mRNA directed protein synthesis is reduced in mussels exposed to higher B[a]P loads [33]. Additionally, M. Galloprovincialis has been shown to response to B[a]P exposure via changes in abundance of proteins related to synthesis and degradation, energy supply (via ATP) and structural proteins [68]. Proteomic results for B[a]P exposure to digestive gland tissue are in agreement with prior studies and support the observed trends identified using transcriptomic methodologies. In the second most enriched pathway (viz. protein processing in endoplasmic reticulum), three heat shock proteins viz. HSP70, HSP90 and HSP40 and other molecular chaperones were identified dependent on exposure conditions. This is not surprising given that many HSPs function as molecular...
chaperones to protect damaged proteins from aggregation, unfold protein aggregates or refold
damaged proteins or target them for efficient removal [111]. These proteins regulate cell response to
oxidative stress with HSP70 strongly up-regulated by heat stress and toxic chemicals. HSP70 plays
several essential roles in cellular protein metabolism [112,113] while HSP40 facilitates cellular
recovery from adverse effects of damaged or misfolded proteins (proteotoxic stress). Changes in
HSPs, in addition to up/down regulation of HSP40, HSP70 and HSP90 have typically been reported
in response to thermal stress in bivalves [98,114,115] and other environmental contaminants such as
B[a]P [33]. In general, the consistent enrichment of genes involved in the Endoplasmic-reticulum
associated protein degradation (ERAD) pathway suggest that aqueous fullerene exposure targets the
cellular pathway involved in targeting misfolding proteins for ubiquitination (post-translational
modification) and subsequent degradation by proteasomes (protein degrading complex, breaks
peptide bonds). It is interesting to note the overlap between organismal response to fullerene
exposure and that of organismal response to thermal stress. Observed enrichment pathways in the
current study viz protein processing in endoplasmic reticulum, apoptosis, ubiquitin mediated
proteolysis, endocytosis, spliceosome, and MAPK signalling pathway have been observed as
differentially enriched in oysters as a response to thermal stress [114].

3.4. Notes
The lack of consensus regarding C₆₀ toxicity may be partly due limited studies which incorporate
both a physiological and ecological approach. As a consequence, little is still known about NP
bioavailability, mode of uptake, ingestion rates and actual internal concentrations related to ADME
[27]. Generally, the greater the water solubility of fullerene aggregates (through e.g. stirring, surface
modifications, sonication), the less the toxicity associated with the exposure [91]. Gomes et al
highlights that while mussels represent a target for environmental exposure to nanoparticles,
exposure duration may significantly contribute to NPs mediated toxicity [116]. As such, it is possible
that the lack of differentially expressed proteins identified in this study is a factor of limited exposure
duration. Limited exposure duration in the region of days or hours is common in the literature, and
it would be of interest to explore long term exposure to NPs to look at the long term impact and
adaptation of mussels in the marine environment. Species specific responses to C₆₀ are abundant in
the literature and it would be remiss to not discuss how our results align with other marine
invertebrates. Exposure to ROS can cause a range of reversible and irreversible modifications of
protein amino acid side-chains which has been reviewed by Ghezzi and Bonetto [117]. Within the
field of aquatic ecotoxicology, the toxic impact and potential mechanisms of single contaminant
exposures have been extensively studied via laboratory experiments (in vivo, in vitro and in silico) and
field monitoring. However, harder to predict is the effects of mixtures of pollutants in the
environment. Biological damage observed cannot simply be linked to the actual environmental
condition as mixtures of contaminants are known to exist in the aquatic ecosystem. This is further
complicated with respect to nanomaterials due to their inherent properties which can amplify or
negate the toxic effects of other compounds [69]. Complicated interactions may occur which make
interpretation complex. For example, proteomic analysis of Mytilus galloprovincialis revealed that
single Cu and B[a]P exposure in addition to a combination of the two generate different protein
profiles with a non-additive profile [68]. Differences in mixture response compared to single exposure
are likely to be related to individual chemical properties and toxicity mechanisms of B[a]P and C₆₀,
as has been noted in B[a]P co-exposed with various metals [118]. C₆₀ concentration was kept constant
with increasing concentrations of B[a]P in an experimental design that has been previously carried
out using algae and crustacean species [22]. This may reflect limited proteome changes at the
exposure concentrations, with concentrations of C₆₀ in the range of 10 - 500 ppb have been reported
to be 10 fold below the no observable adverse effect level (NOAEL) [119,120]. At 1 mg/L, an increase
in GST activity in the digestive gland has been reported [110] C₆₀ is known to bind to minor grooves
of double stranded DNA and trigger unwinding and disruption of the DNA helix [102] C₆₀ adsorbs
onto cell-membrane P-glycoprotein through hydrophobic interactions, but the stability and
secondary structure of the protein are barely affected [121]. P-glycoprotein is present in Mytilus
galloprovincialis [122] C60 and its derivatives are known to impact DNA and RNA in terms of stability, replication and reactivity in addition to structural stabilisation [123,124]. In a recent study, Canesi et al determined that C60 fullerene exposure to *Mytilus galloprovincialis* hemocytes did not induce significant cytotoxicity, and instead stimulated immune and inflammatory parameters such as lysozyme release, oxidative burst and NO production [10]. Nanomaterial suspensions can induce inflammatory processes in bivalve hemocytes akin to those observed in vertebrate cells [10]. Results from mammalian studies suggest that C60 fullerene exposure results predominantly in inflammatory responses [125].

5. Conclusions

This study has demonstrated for the first time the interaction between two ubiquitous environmental contaminants with an apparent antagonistic relationship at the genotoxic and the proteome expression level. No Trojan horse effects were observed for uptake or toxicity of the co-contaminants B[a]P in interaction with C60. Proteome profile is dependent on concentration and treatment. The exposure to the three conditions had overlap and common mechanisms of response irrespective of differences in mode of action. The provided list of condition specific differentially expressed proteins and enriched pathways (Spreadsheet S1) may represent a step towards definitively identifying mode of action of these compounds in bivalves when combined with other OMICs based approaches. It should be noted that the antagonistic proteome response observed in the current study between B[a]P and C60 is based on a single concentration of the fullerene and as such represents a general overview of toxicological behaviour. It is possible that that this antagonistic interaction will change when another dose range is selected [93]. Gomes et al previously highlighted that while mussels represent a target for environmental exposure to nanoparticles, exposure duration may significantly contribute to NPs mediated toxicity [116]. As such, further work must be carried out to explore mixture effects at different concentrations and over differing exposure duration.

Supplementary Materials: The following are available online, Figure S1: Representative particle size distribution showing the intensity-weighted hydrodynamic diameter (dH) of nC60 in mussel-exposed seawater as determined by DLS (653±87 nm), Figure S2: (a) Bright-field TEM and (b) point EDX spectroscopy analysis of Er3N@C80, Figure S3: Dark-field STEM and EDX spectroscopy mapping analysis of Er3N@C80, confirming the necessity for spectroscopy to confirm the presence of labelled fullerenes, using the characteristic X-rays emitted from Er upon electron irradiation, Figure S4: (a,c,e) Dark-field STEM and (b,d,f) corresponding point EDX spectroscopy analysis of cross-sections of mussel digestive gland exposed to Er3N@C80, Table S1: The influence of benzo(a)pyrene (B[a]P) of the hydrodynamic diameter (dH) of nC60 in mussel-exposed seawater as determined by DLS, Table S2: The concentration of B[a]P in seawater at T0, day 1 and day 3, Table S3: The concentration of nC60 in seawater at T0, day 1 and day 3, Spreadsheet S1: Full list of DEPs, enriched Gene Ontology (GO) terms and KEGGS pathways, R script S1: R script used for proteomics analysis.

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