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Surgical face masks as a source of emergent pollutants in aquatic systems: Analysis of their degradation product effects in *Danio rerio* through RNA-Seq.

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**Highlights**
- Surgical face masks (FMs) can be a source of emergent pollutants in aquatic systems.
- The release of chemicals from FMs to water depends on their degradation stage.
- RNA-Seq analysis of zebrafish larvae exposed to FM degradation products revealed alterations in the transcriptome profile.
- FM degradation stage influenced the biological processes modulated in zebrafish.
- FM degradation products strongly impacted the expression of reproduction-related genes.

**Abstract**
Surgical face masks are the most popularised and effective personal equipment for protecting public health during the COVID-19 pandemic. They are composed of plastic polymer fibres with a large amount of inorganic and organic compounds that can be released into aquatic environments through degradation processes. This source of microplastics and inorganic and organic substances could potentially impact aquatic organisms. In this study, the toxicogenomic effects of face masks at different stages of degradation in water were analysed in zebrafish larvae (*Danio rerio*) through RNA-Seq. Larvae were exposed for 10 days to three treatments: 1) face mask fragments in an initial stage of degradation (poorly degraded masks -PDM- products) with the corresponding water; 2) face mask fragments in an advanced stage of degradation (highly degraded masks -HDM- products) with the corresponding water; and 3) water derived from HDM (W-HDM). Transcriptome analyses revealed that the three treatments provoked the down-regulation of genes related to reproduction, especially the HDM products, suggesting that degradation products derived from face masks could act as endocrine disruptors. The affected genes are involved in different steps of reproduction, including gametogenesis, sperm-egg recognition and binding or fertilisation. Immune-related genes and metabolic processes were also differentially affected by the treatments.

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1. Introduction

The concept of “One Health” proposed by the World Health Organization (WHO) with the COVID-19 pandemic is currently more applicable than ever. One health is the intersection of human, animal and environmental health. Therefore, the synergies and harmony of these three blocks is the cornerstone of wellness and global health. Environmental degradation due to human behaviours and waste mismanagement shows an unsure future for the biosphere and humanity itself.

During the COVID-19 pandemic, the use of personal protective equipment (PPE), such as face masks, has supported significant advancements in public health (Howard, 2020). Currently, face masks are a legal requirement in public spaces in many countries. The reported monthly consumption of face masks has been approximately 129 billion for 7.8 billion people across the globe (Kalina et al., 2020). However, the management of PPE residues as well as uncivil behaviours of the population have already revealed face masks in terrestrial and aquatic compartments. A recent study in Toronto, Canada, showed that 95% of the population have already revealed face masks in terrestrial and aquatic environments as well as uncivil behaviours of the population shows an unsure future for the biosphere and humanity itself.

1.1. Fragmentation and degradation of surgical face masks

The daily use of surgical face masks increases difficulties related to an environmental issue that is far from solved: plastic consumption and therefore plastic pollution. Before the COVID-19 pandemic, different directives and regulations (such as the Basel Convention, United Nations Convention on the Law of the Sea, International Convention for the Prevention of Pollution from Ships, Joint Group of Experts on the Scientific Aspects of Marine Environmental Protection and United Nations Global Partnership on Marine Litter) were advanced to alleviate plastic pollution and therefore the health of the planet. However, surgical face masks are a and large source of plastic pollution and nanofibre delivery due to their fabrication process (Leung and Sun, 2020). Moreover, when face masks reach the natural environment, environmental agents initiate their degradation and fragmentation processes to degrade the mask into smaller pieces called microplastics (MPs < 5 mm; Thompson et al., 2004) and nanoplastics (NPs < 100 nm; Jambeck et al., 2015).

Wastewater treatment plants (WWTPs) are the sink of many emerging pollutants and MPs (de Oliveira et al., 2020; Raju et al., 2020). Nonspecific treatments for the depuration of pollutants of emerging concern like WWTPs a source of pollutants in their surrounding environment, which in most cases is aquatic ecosystems. Due to their small size, MPs are available for a vast group of aquatic animals (Ferreira et al., 2019). Fish are one of the most sensitive groups to MP ingestion, with a frequency of accumulated MPs of approximately 72% (Provencher et al., 2019) and, since ingestion is the main way for MPs accumulation, fish are an animal group of special concern in this sense. The ingestion of MPs by fish is mainly due to their attractive coloration, low density, buoyancy, and resemblance to food (Güven et al., 2017).
2.2. Surgical face mask characterisation and its behaviour in freshwater systems

2.2.1. Size distribution and behaviour

The size distribution of PDM and HDM pieces was analysed by optical microscopy (n:200). In addition, the three layers of the surgical face masks, as well as different zones of face masks, were analysed by Fourier transform infra-red spectroscopy (FTIR) through the Bruker Alpha System. Secondary characteristics of PDM and HDM fragments were analysed by static light scattering (SLS, Malvern Master Sizer 2000, with software version 5.61). The behaviour of the face mask fragments was studied in zebrafish water over time: 0, 24, 48 and 72 h (time between water tank changes). The protocol for zebrafish water preparation is included in the Supplementary materials.

2.2.2. Characterisation of organic compounds

For organic compound characterisation, the elastics of the face masks and aluminium nose clips were also removed. Between 1 and 1.3 g was used to extract the organic compounds with 35 mL of chloroform as the polar dissolvent. The extractions were performed in 100-mL Erlenmeyer flasks and under bath sonication. The liquid obtained from these face masks and the water from the PDM and HDM stored in Section 2.1 were concentrated by a rotavapor to reach a liquid volume of 1 mL. The samples were transferred to vials for organic compound analysis, which was conducted by gas chromatography–mass spectrometry (GC-MS). Details about the GS-MS analysis are provided in the Supplementary materials (Table S1).

2.2.3. Characterisation of inorganic compounds

The elastics of the face masks and aluminium nose clip were also removed, and the face masks were cut with clean scissors. Approximately 0.2 g of face masks was digested by acid digestion in a microwave following the procedures of Rodríguez-Seijo et al. (2015). After digestion, the samples were evaporated and diluted in 10 mL of ultrapure water. The water samples were acidified with 2% of HNO₃. The analysis of the samples was performed by inductively coupled plasma mass spectrometry (ICP-MS; Thermo elemental) in semiquantitative mode to detect the concentration of all the elements of the periodic table.

2.3. Zebrafish larvae treatments

A total of 240 wild-type zebrafish (120 larvae for mortality analysis and 120 for transcriptome experiments) at an age of 15 days post-fertilisation (dpf) were obtained from the facilities of the Instituto de Investigaciones Marinas (Vigo, Spain), where zebrafish are maintained following established protocols (Nusslein-Volhard and Dahm, 2002; Westerfield, 2000). At this stage zebrafish larvae are not sexually differentiated, which allows to avoid the gender bias of the data. Fish care and challenge experiments were conducted according to the guidelines of the CSIC National Committee on Bioethics under approval number ES360570202001/17/FUN.01/INM06/BNG. Zebrafish larvae were maintained under a 12:12-h light-dark cycle at pH 7.0 and temperature 28–29 °C and, from 5 dpf, were fed daily with commercial food (Sera Micron, North Rhine-Westphalia, Germany).

Dry PDM and HDM fragments were mixed with their corresponding filtered water (described in Section 2.1) to obtain the same concentration in the treatments. Three replicates (10 larvae per replicate) were used for each treatment: a) control, b) PDM (10 mg·L⁻¹ of the PDM fragments with the corresponding water), c) HDM (10 mg·L⁻¹ of the HDM fragments with the corresponding water) and d) water derived from HDM (W-HDM); Fig. 1. The water volume of each tank was 0.5 L, and it was renewed every 3 days with the corresponding treatment. After 10 days of treatment exposure, seven zebrafish larvae were collected from each tank and pooled for RNA-Seq analysis. The larvae were euthanized by thermal shock on ice, and RNA isolation was immediately carried out after zebrafish collection. A parallel identical experiment was conducted to analyse whether the treatments had some impact on the survival of the zebrafish larvae.

2.4. RNA isolation and Illumina sequencing

Total RNA was isolated using the Maxwell 16LEV Simply RNA Tissue Kit (Promega) according to the manufacturer’s instructions. The quantity of RNA was measured in a NanoDrop ND-1000 (NanoDrop Technologies, Inc.), and RNA integrity was analysed in an Agilent 2100 Bioanalyzer (Agilent Technologies Inc.) according to the manufacturer’s instructions.

Fig. 1. Procedures and experimental design for PDM, HDM and W-HDM treatments and exposure to zebrafish.
instructions. All the samples passed the quality control tests and were used for Illumina library preparation.

Three biological replicates per treatment (control, PDM, HDM and W-HDM), composed of each replicate by 7 pooled larvae, were used for mRNA sequencing. Double-stranded cDNA libraries were constructed using the TruSeq Stranded mRNA LT Sample Prep Kit (Illumina, USA), and sequencing was performed using Illumina NovaSeq 6000 technology at Macrogen Inc. (Seoul, Republic of Korea). The raw read files (in FASTQ format) were deposited in the Sequence Read Archive (SRA) (https://www.ncbi.nlm.nih.gov/sra) under BioProject accession number PRJNA770826.

2.5. Trimming, mapping, RNA-Seq and differential expression analysis of mRNAs

CLC Genomics Workbench v. 20.0.4 software (CLC Bio, Qiagen, Denmark) was used to filter and trim raw reads and perform the RNA-Seq and differential expression analyses. Raw reads were trimmed to remove adapter sequences and low-quality reads (Phred score limit of 0.05). RNA-Seq analyses were carried out by mapping the filtered reads against the last version of the zebrafish genome (GRCz11) with the following parameters: length fraction = 0.8, similarity fraction 0.8, mismatch cost = 2, insertion cost = 3 and deletion cost = 3. The expression values were set as transcripts per million (TPMs). Finally, a differential expression analysis test was used to compare gene expression levels and to identify differentially expressed genes (DEGs) among the treatments. Transcripts with absolute fold change (FC) values ≥ 2 and p values < 0.05 were retained for further analyses.

To identify and quantify the directions of variability in the data, a principal component analysis (PCA) plot was constructed using the TPM expression values with the ClustVis tool (https://biit.cs.ut.ee/clustvis/; Metsalu and Vilo, 2015). Heatmaps were created to show a hierarchical clustering of gene expression (TPM values) using the average linkage method with the Euclidean distance with the ClustVis tool.

To illustrate the common and exclusive DEGs among the treatments compared to the control larvae, Venn diagrams were performed using the Venny 2.1 tool (https://bioinfogp.cnb.csic.es/tools/venny/).

2.6. Gene Ontology (GO) terms, Kyoto Encyclopaedia of Genes and Genomes (KEGG) pathways and protein domain enrichment analyses

GO enrichment analysis of biological processes (BP), KEGG pathway analysis and protein domain (PD) enrichment analysis were conducted using DAVID software (https://david.ncifcrf.gov/; Huang et al., 2007). In all cases, a Bonferroni correction was applied (p < 0.05), and a minimal count of 5 was established. For domain enrichment, the InterPro database (Apweiler et al., 2001) was used. The representation of the different categories was based on the fold-enrichment value.

2.7. Protein–protein interaction networks

The interactions of the proteins encoded by the DEGs in each treatment were analysed with STRING v11.0 software (https://string-db.org/; Szklarczyk et al., 2021). The clusters formed in the STRING software indicated that the expressed proteins had more interactions among themselves than would be expected for a random set of proteins of similar size drawn from the genome. Such an enrichment indicated that the proteins were at least partially biologically connected as a group.

3. Results

3.1. Primary chemical characterisation of surgical face masks and secondary chemical characterisation of face mask degradation products in freshwater

The commercial face masks used in this study were composed of three layers, with polypropylene as a plastic polymer (Fig. 2A). The nominal sizes of the fragments used for the PDM and HDM treatments were 3.10 ± 1.6 mm and 1.25 ± 0.52 mm (mean ± standard deviation), respectively (Fig. 2B). Both PDM and HDM fragments showed continuous fragmentation/degradation behaviour after 72 h in zebrafish water (Fig. 2C). After this period, the face mask fragments reached a homogeneous population with only a peak between 17.37 and 12.22 μm for the PDM and a heterogeneous population with four peaks between 1.44 and 954.90 μm for the HDMI (Fig. 2C).

The organic and inorganic compounds were analysed in the face masks and in the water derived from PDM and HDMI. The chromatography analysis of the face masks showed a profile of waxes and hydrocarbons (Fig. 2A). With respect to inorganic compounds, the face masks showed high concentrations of Li (49,695 mg kg⁻¹), Ti (10,475.19 mg kg⁻¹), Al (6197.64 mg kg⁻¹), Si (3904.01 mg kg⁻¹) and Mg (340.88 mg kg⁻¹). Values of Cr, Mn, Fe, Co, Ni, Cu, Zn and Ga were also present, and low values were found for In, Ba, La, Ce, Pb, Bi and Th (Fig. 2E). However, these profiles of organic and inorganic compounds found in the surgical face masks were not the same as those found in the zebrafish water after a week of UV-C irradiation (degradation conditions). Waxes and hydrocarbons found in the face masks were not released into the water. However, other compounds, such as pyridine 2,4,6-trimethyl, 1-dodecanamine N,N-dimethyl and alanine N-methyl-N-methoxyacarbonyl-undecyl ester, were found in the water derived from PDM and HDMI degradation (Fig. 2D). No peaks for these compounds were found in the control zebrafish water under the same exposure conditions; therefore, they could be byproducts formed under surgical face mask degradation conditions.

Low concentrations of inorganic compounds, such as Na, B, Al, Ti, Fe and Cu, were only released into the water of HD-M. Conversely, Zn and Sr were found in both the PDM and HDMI water samples (Fig. 2F).

3.2. Transcriptome modulation of zebrafish larvae exposed to the different face mask degradation products

The survival of the larvae was higher than 91% over the experiment for all the tanks. Although no significant mortalities were observed in the larvae treated with the different face mask products (data not shown), RNA-Seq analyses were conducted to evaluate the impact of the different face mask degradation products on the overall transcriptome of zebrafish larvae.

A total of 321,832,678 million reads were obtained from the twelve sequenced samples, with a mean of 26,819,390 million reads per sample. From these, 99.9% of the raw reads passed the quality control, and these high-quality reads were mapped against the zebrafish genome. For the different samples, between 95% and 97% of the reads were successfully mapped against the genome (Table S2).

PCA of the twelve samples revealed a poor separation of the samples belonging to the four different treatments (Fig. 3A). However, despite the low differentiation of the overall transcriptomes, an important group of genes were significantly modulated after exposure to the treatments. RNA-Seq and further differential expression analyses revealed 138, 287 and 105 differentially expressed genes (DEGs) in zebrafish larvae exposed to PDM, HDMI and W-HDMI products, respectively, compared with the control larvae (Fig. 3B; Tables S3–S5). Whereas a similar number of up-regulated DEGs was observed for the three treatments (80, 90 and 82 for the larvae exposed to PDM, HDMI and W-HDMI products, respectively), the number of down-regulated genes was much higher for those larvae treated with the HDMI products (58, 197 and 23 for the larvae exposed to PDM, HDMI and W-HDMI products, respectively) (Fig. 3B). The expression profiles of the DEGs in TPM values were represented in heatmaps where the three biological replicates of each treatment clustered together, showing good consistency among the replicates corresponding to the same treatment (Fig. 3C).

A Venn diagram of the DEGs for each treatment compared with the control revealed 18 common DEGs among the three treatments.
Fig. 2. Surgical face mask characterisation and its behaviour in the aquatic matrix. A) FTIR spectra of the different parts of the surgical face mask and different layers. B) Nominal sizes of fragments from PDM and HDM treatments. C) Behaviour of both size fragments of face mask in zebrafish water over 72 h (time between zebrafish water changes). D) Spectrum of the organic substances released into the water from the PDM and HDM treatments. E) Concentration of different elements present in the surgical face masks. For some elements the concentration was zero. F) Inorganic elements released into water from PDM and HDM treatments.
(Fig. 3D), 15 of which were up-regulated and the other 3 down-regulated. The common up- and down-regulated genes are shown in Fig. 3E and F, respectively. No significant biological processes or KEGG pathways were found to be enriched for these common DEGs. However, for the common down-regulated genes among the three treatments, two of them (zp3a.1 and si:zfos-223e1.2) were linked to the reproductive process, since zp3a.1 (zona pellucida glycoprotein 3a, tandem duplicate 1) encodes a component of the oocyte’s zona pellucida and si:zfos-223e1.2 (cfa61; cilia- and flagella-associated protein 61) is required for sperm flagellum formation and male fertility (Huang et al., 2020). Conversely, the common up-regulated DEGs seemed to be involved in a variety of processes, with the most affected genes across the three treatments being si:dky-242g16.2 (ppp1r3c2a; protein phosphatase 1 regulatory subunit 3C2, duplicate a) and si:ch211-121a2.2 (dusp29; dual specificity phosphatase 29), which are involved in glucose metabolism, among other functions.
3.3. Main processes and pathways affected by the different treatments

3.3.1. Zebrafish larvae exposed to PDM products

The top 25 most up- and down-regulated DEGs in zebrafish larvae exposed to PDM products are summarised in Table 1. Among the top 25 down-regulated DEGs, genes involved in reproductive processes, such as several genes encoding zona pellucida glycoproteins (zp2.1, zp3, zp2.5, zp2.2, zp3e, zp3.2.1 and zp2.3), were found. These down-regulated genes were inhibited more than 10-fold compared with the controls. GO biological process enrichment analysis of the down-regulated DEGs in zebrafish larvae exposed to PDM products revealed several domains linked to the immune response, such as “P-type trefoil”, “zona pellucida, conserved site”, “zona pellucida domain”, “C-type lectin”, “C-type lectin fold” and “C-type lectin-like” (Fig. 4A).

Conversely, the top 25 up-regulated genes after PDM treatment were related to i) innate immune response, such as *daph1b_2* (death associated protein 1b), *itn3* (interleukin 3), *itn2_1* (interleukin 1), *sdc2* (selectin C2 domain), *sdc1* (selectin C1 domain), *liver-expressed antimicrobial peptide 2*, *sdc* (sodium-dependent vitamin transporter), *itln2_1* (C-type lectin fold), *itln3* (interlectin 3), *itln2* (interlectin 2) and ii) isoprenoid/sterol biosynthesis, such as *cyp2r1* (cytochrome P450, family 2, subfamily R, gene), *cyp7a1* (cytochrome P450, family 7, subfamily A). (Table 1).

| Name     | Blast2 Go UnitProt/SwissProt Description | FC  | Name     | Blast2 Go UnitProt/SwissProt Description | FC  |
|----------|------------------------------------------|-----|----------|------------------------------------------|-----|
| *daph1b_2* | death associated protein 1b             | 15.01 | *zp2.1* | zona pellucida glycoprotein 2, tandem duplicate 1 | -40.75 |
| *itn3*    | interluekin 3                           | 12.74 | *sdc2* | selectin C2 domain                       | -37.98 |
| *itn2_1*  | dual specificity phosphatase 29         | 10.26 | *zc:171445* | serine-type endopeptidase inhibitor activity | -33.53 |
| *sdc1*    | interluekin 2                           | 8.61  | *zc:2211-14610.7* | sterol metabolic process | -27.78 |
| *klf2a*   | kruuppel-like factor 2a                 | 8.33  | *zc:173856* | sterol metabolic process | -21.26 |
| *sdc2*    | liver-expressed antimicrobial peptide 2  | 6.72  | *wul1702e0* | sterol metabolic process | -20.0 |
| *zc:2211-113d11.8* | protein phosphatase 1 regulatory subunit 3 C B | 5.1  | *zc:173544* | sterol metabolic process | -19.92 |
| *cyp2r1*  | cytochrome P450, family 2, subfamily R, | 5.05  | *zc:173544* | sterol metabolic process | -19.99 |
| *itln2_1* | interlectin 2                           | 5.03  | *sdc2* | sterol metabolic process | -19.65 |
| *itln3*   | interlectin 3                           | 4.97  | *zc:2211-14610.8* | sterol metabolic process | -18.44 |
| *zc:153642* |......| 4.31  | *zp2.5* | zona pellucida glycoprotein 2, tandem duplicate 5 | -17.75 |
| *zc:174917* |......| 3.97  | *zc:22* | zona pellucida glycoprotein 2, tandem duplicate 2 | -17.77 |
| *sdc2*    | liver-expressed antimicrobial peptide 2  | 3.67  | *zc:136254* | zona pellucida glycoprotein 2, tandem duplicate 1 | -16.87 |
| *pnc*     | .........                               | 3.63  | *sdc2* | zona pellucida glycoprotein 2, tandem duplicate 2 | -14.56 |
| *spd*     | collagen alpha-1(XI) chain             | 3.51  | *zc:171474* | zona pellucida glycoprotein 2, tandem duplicate 1 | -14.29 |
| *sdc2*    | .........                               | 3.47  | *zc:171474* | zona pellucida glycoprotein 2, tandem duplicate 1 | -14.18 |
| *sdc2*    |......                               | 3.34  | *zc:2211-14722.4* | zona pellucida glycoprotein 2, tandem duplicate 1 | -14.1 |
| *sdc2*    |......                               | 3.33  | *zc:2211-14722.4* | zona pellucida glycoprotein 2, tandem duplicate 1 | -13.46 |
| *sdc2*    |......                               | 3.31  | *zc:2211-14722.4* | zona pellucida glycoprotein 2, tandem duplicate 1 | -12.49 |
| *sdc2*    |......                               | 3.25  | *zc:2211-14722.4* | zona pellucida glycoprotein 2, tandem duplicate 1 | -12.46 |
| *sdc2*    |......                               | 3.19  | *zc:2211-14722.4* | zona pellucida glycoprotein 2, tandem duplicate 1 | -11.34 |
| *sdc2*    |......                               | 3.15  | *zc:2211-14722.4* | zona pellucida glycoprotein 2, tandem duplicate 1 | -11.02 |
| *sdc2*    |......                               | 3.09  | *zc:2211-14722.4* | zona pellucida glycoprotein 2, tandem duplicate 1 | -10.67 |
| *sdc2*    |......                               | 3.03  | *zc:2211-14722.4* | zona pellucida glycoprotein 2, tandem duplicate 1 | -10.06 |
| *sdc2*    |......                               | 3.02  | *zc:2211-14722.4* | zona pellucida glycoprotein 2, tandem duplicate 1 | -8.9 |

Table 1
Top 25 most up- and down-regulated DEGs in zebrafish larvae exposed to PDM treatment for 10 days.
Fig. 4. Biological processes (BP), KEGG pathways, protein domains (PD) and protein–protein interactions enriched for the genes modulated by the PDM treatment. A) Enrichment analyses of the down-regulated genes. B) Enrichment analyses of the up-regulated genes. C) STRING analysis (protein–protein interaction) of the DEGs under PDM treatment.
interconnected with the previous clusters and corresponded to the main down-regulated DEGs, which are related to different reproductive processes, such as cumulus cells and zona pellucida (zp2.1, zp2.3, zp2.5, zp3b, zp3a.1, zp3b) and chromosome condensation in primordial germ cells (h1m).

3.3.2. Zebrafish larvae exposed to HD3 treatment

The top 25 most up- and down-regulated DEGs in zebrafish larvae exposed to HD3 products are shown in Table 2. Among the top 25 down-regulated genes were also genes related to reproductive processes, such as the zona pellucida glycoproteins zp2.1, zp3, zp2.3, zp2.5, zp2.2, zp2.6, zpax4, zp3a.1 and zp3a.2.1, which were inhibited more than 50-fold compared with the control larvae. Although not restricted therein, GO biological process enrichment analyses of the downregulated DEGs revealed numerous terms related to reproductive processes (“P granule organization”, “DNA methylation involved in gamete generation”, “egg coat formation”, “positive regulation of acrosome reaction”, “binding of sperm to zona pellucida”, “gamete generation”, “meliotic nuclear division”, “oogenesis”, “germ cell development”, “spermatid development”, “meliotic cell cycle” and “spermatogenesis”) and RNA silencing (“negative regulation of gene silencing by miRNA”, “piRNA metabolic process”, “regulation of translation” and “gene silencing by RNA”) (Fig. 5A). Similar to the results obtained for the GO terms, the enriched protein domains showed a high representation of terms linked to reproduction (“Fetuin-A-type cystatin domain”, “Fetuin-B-type cystatin domain”, “P-type trefoil”, “DNA binding HORMA”, “zona pellucida domain”) or RNA silencing during gametogenesis (“OST-HTH/LOTUS domain”, “stem cell self-renewal protein Piwi”, “arginate/Dicer protein”, “Tudor domain”) (Fig. 5A). Interestingly, only one KEGG pathway was significantly enriched, which corresponded to “dorsal-ventral axis formation” (Fig. 5A).

The top 25 up-regulated DEGs in zebrafish larvae exposed to HD3 products are shown in Table 2. Among them, three members of the heat shock protein 70 family were shown in Table 2. Among them, three members of the heat shock protein 70 family, hspa1a.1, hspa1b.1 (haemoglobin, alpha adult 1), hspa1a.1 (haemoglobin, beta alpha 1) were also genes related to reproductive processes, such as ischaemia, inflammation, osmotic and oxidative stress (Franzellitti and Fabbri, 2005). Up-regulated genes involved in the immune response were recognised, such as dapan1b.2 (death-associated protein 1b), efna5b (ephrin-A5b), fosb (FB3 murine osteosarcoma viral oncogene, si:ch211–181d7.1_2 (nlp3; NACH, LRR and PYD domain-containing protein 3), si:ch211–181d7.3 (nlp12; NACH, LRR and PYD domain-containing protein 12), si:cdk4-204a24.11 (gv1; interferon-induced very large GTPase 1) and macir (macrophage immunometabolism regulator). GO biological process enrichment analysis of the up-regulated DEGs revealed terms involved in the response to stimulus (“response to heat”, “response to cadmium ion” and “intracellular signal transduction”) and developmental process (“camera-type eye morphogenesis”) (Fig. 5B). The KEGG pathways significantly enriched for the up-regulated genes were “spliceosome”, “protein processing in endoplasmic reticulum”, “MAPK signalling pathway” and “endocytosis” (Fig. 5B). Due to the high representation of HSP70-encoding genes, the most enriched protein domains corresponded to “heat shock protein 70, conserved site” and “heat shock protein 70 family” (Fig. 5B).

For the HD3 product treatment, the protein-protein interactions were higher than those observed for PDM products (Fig. 5C). In this case, the main cluster formed by the DEGs corresponded to genes that were also involved in cumulus cells and zona pellucida (zp2.1, zp2.3, zp2.5, zp2.6, zp3.1, zp3a.1, zp3a.2, zp3b), but some additional genes related to gametogenesis and embryogenesis (patl2, sarl1, bsc, gsf31, dazl, kpn7) were also observed, as well as DEGs involved in cell adhesion and tight junctions (clndn, clng), cell proliferation (s100a1, npm2), RNA transcription and protein translation (pabpc1, lsm14b, gsf3ab, habp4, elf4e1b), among others (Fig. 5C). This cluster was connected with a second cluster, also mainly composed of genes encoding proteins with a role in germ cell development, proliferation and integrity (smc1b, vasa, fkbp6, rnf17, tdrd1, tdrd5, tdrd7, dnd1, piwil1, ago3b, pld6, hermad1). The third cluster included several genes with a role in coagulation (cph2, habp2, ahsg, krnl, spp2, fgf) but also with iron (hpx, cp), vitamin D (gc) and lipids (apob2.2), transport, and hormones and receptors linked to glucose metabolism (gqga, stz2, vpr1b) (Fig. 5C).

3.3.3. Zebrafish larvae exposed to W-HDM treatment

The top 25 most up- and down-regulated DEGs in zebrafish larvae exposed to W-HDM treatment are shown in Table 3. Among the top 25 down-regulated genes were also DEGs related to reproductive processes, such as span1 (sperm adhesion molecule 1), zp3a.1 and zspf223e1.2 (cspf61), and two members of the HSP70 family, but these DEGs were different from those differentially expressed with the HD3 products hspa12a.1 and hspa12a.2. For the down-regulated genes after W-HDM treatment, no GO biological processes or KEGG pathways were significantly enriched, and only the protein domain “Zinc finger, B-box” was overrepresented (Fig. 6A).

With regard to the top 25 up-regulated DEGs after W-HDM treatment (Table 3), an important insight is the number of DEGs encoding different haemoglobin types: hbaa1.2 (haemoglobin, beta adult 1), hbaa1.1 (haemoglobin, alpha adult 2), hbaa1 (haemoglobin, alpha adult 1) and hbaa5 (haemoglobin, alpha embryonic 5). The high representation of these haem-binding proteins explains the GO biological process enriched terms “oxygen transport”, “response to hypoxia” and “haemopoiesis” for the up-regulated DEGs (Fig. 6B); different immune, response to stimulus, metabolism and development processes were also significantly enriched (Fig. 6B). The KEGG pathways enriched under H-HDM exposure were “RNA degradation”, “MAPK signalling pathway”, “herpes simplex infection” and “protein processing in endoplasmic reticulum” (Fig. 6B). As expected, among the most enriched protein domains were those related to the haemoglobin proteins (“haemoglobin, alpha”, “haemoglobin, pi”, “globin”, “globin-like”) but also, as observed for the up-regulated genes after HD3 treatment, protein domains linked to the HSP70 family (“heat shock protein 70” and “heat shock protein 70 family”) (Fig. 6B).

In relation to the STRING analysis, the DEGs predominantly grouped into a main cluster composed of genes encoding proteins related to cell proliferation (fos, fosl1a, bg2), immunity and inflammation (irg11, socs1, socs3a, cebp, mmp9), extracellular matrix degradation (mmp9, timp2b) and haem binding (hbaa1, hbaa1, hbaa1, hbaa1), among other functions (Fig. 6C).

3.4. Face mask degradation products differentially impact biological processes

Based on the enrichment analyses, we wanted to examine in a more detailed manner different processes that were significantly affected by at least one of the treatments. To achieve this goal, we constructed heatmaps representing the TPM values of the genes across the different treatments. Although the initial face mask material was the same for the three treatments, the stage of degradation of the face masks provoked significant differences in the zebrafish transcriptome response.

Since several enriched terms linked to the immune response were found to be significantly enriched after the face mask treatments, we separately analysed groups of genes that were well-defined for their main immune function (complement pathway, coagulation, type I interferon system, antigen presentation and inflammation). As expected, due to complement-coagulation cross-talk (Kenawy et al., 2015), both processes followed a similar tendency, with most of the genes involved in these pathways being strongly inhibited in zebrafish larvae exposed to the HD3 products and with the lower impact caused by W-HDM products (Fig. 7A and B). Similar to that observed for the complement and coagulation genes, HD3 treatment inhibited certain antiviral genes.
Table 2
Top 25 most up- and down-regulated DEGs in zebrafish larvae exposed to HDM for 10 days.

| Name          | Complete name or Predicted description | FC  | Name          | Complete name or Predicted description | FC  |
|---------------|---------------------------------------|-----|---------------|---------------------------------------|-----|
| dap1b_2       | death associated protein 1b            | 16.61 | zp2.1        | zona pellucida glycoprotein 2, tandem   | -562.51 |
| rsf1b_1,2     | remodelling and spacing factor 1b      | 16.56 | zp3          | zona pellucida glycoprotein 3           | -394.92 |
| BX005305.1    | legumain-like                          | 12.71 | zp2.3        | zona pellucida glycoprotein 2, tandem duplicate 3 | -321.14 |
| zgc:101699    | cytosolic phospholipase A2 gamma       | 11.04 | zp2.5        | zona pellucida glycoprotein 2, tandem duplicate 5 | -290.81 |
| mrf1026.1     | zinc finger protein 1026               | 10.32 | zgc:173856   | zona pellucida glycoprotein 2, tandem duplicate 5 | -282.11 |
| stdkey-920.3  | stdkey-920.3                           | 10.15 | stdkey-14610.7 | zona pellucida glycoprotein 2, tandem duplicate 2 | -203.1 |
| sichi211-121a2.2 | dual specificity phosphatase 29       | 10.13 | stdkey-14610.7 | zona pellucida glycoprotein 2, tandem duplicate 2 | -199.07 |
| zgc:171534    | zgc:171534                             | 8.7  | stdkey-2410.8 | zona pellucida glycoprotein 2, tandem duplicate 6 | -193.35 |
| opn4a         | opsin 4a (melanopsin)                  | 6.19  | stdkey-14610.7 | zona pellucida glycoprotein 2, tandem duplicate 6 | -167.48 |
| hpy70.1       | heat shock cognate 70-kd protein, tandem | 6.01  | stdkey-2410.7 | zona pellucida glycoprotein 2, tandem duplicate 6 | -167.48 |
| nif1          | nitrilase 1 [Source:ZFIn;Acc:ZDB-GENE-040912-65] | 5.73  | stdkey-2410.7 | zona pellucida glycoprotein 2, tandem duplicate 6 | -167.48 |
| msh4          | mutS homologue 4                       | 5.24  | stdkey-2410.7 | zona pellucida glycoprotein 2, tandem duplicate 6 | -167.48 |
| dnah11.2      | dynein, axonemal, heavy chain 11       | 4.96  | stdkey-2080.4 | zona pellucida glycoprotein 2, tandem duplicate 6 | -167.48 |
| cyp230.2       | cytochrome P450, family 2, subfamily K | 4.77  | stdkey-2410.7 | zona pellucida glycoprotein 2, tandem duplicate 6 | -167.48 |
| hpy70.2       | heat shock cognate 70-kd protein, tandem | 4.33  | stdkey-2290.4 | zona pellucida glycoprotein 2, tandem duplicate 6 | -167.48 |
| kcnh6b        | potassium voltage-gated channel, subfamily | 4.08  | stdkey-2290.4 | zona pellucida glycoprotein 2, tandem duplicate 6 | -167.48 |
| efn5a2b       | ephrin-A5b                             | 4.03  | stdkey-2290.4 | zona pellucida glycoprotein 2, tandem duplicate 6 | -167.48 |
| stdkey-2420.16 | Protein phosphatase 1 regulatory subunit 3 C-B | 3.7  | stdkey-2290.4 | zona pellucida glycoprotein 2, tandem duplicate 6 | -167.48 |
| hpy70.1       | heat shock cognate 70-kd protein, like  | 3.66  | stdkey-2290.4 | zona pellucida glycoprotein 2, tandem duplicate 6 | -167.48 |
| fosb          | FBJ murine osteosarcoma viral oncogene | 3.65  | stdkey-2290.4 | zona pellucida glycoprotein 2, tandem duplicate 6 | -167.48 |
| sichi73-191k20.3 | sichi73-191k20.3                  | 3.63  | stdkey-2290.4 | zona pellucida glycoprotein 2, tandem duplicate 6 | -167.48 |
| sichi211-181d7.1,2 | NACHT, LRR and PYD domains-containing protein 3 | 3.59  | stdkey-2290.4 | zona pellucida glycoprotein 2, tandem duplicate 6 | -167.48 |
| sichi211-181d7.3 | NLR family CARD domain-containing protein 3 | 3.58  | stdkey-2290.4 | zona pellucida glycoprotein 2, tandem duplicate 6 | -167.48 |
| stdkey-204a24.11 | interferon-induced very large GTase 1 | 3.53  | stdkey-2290.4 | zona pellucida glycoprotein 2, tandem duplicate 6 | -167.48 |
| MACIR         | macrophage immunometabolism regulator | 3.48  | stdkey-2290.4 | zona pellucida glycoprotein 2, tandem duplicate 6 | -167.48 |
Fig. 5. Biological processes (BP), KEGG pathways, protein domains (PD) and protein–protein interactions enriched for the genes modulated by HDM treatment. A) Enrichment analyses of the down-regulated genes. B) Enrichment analyses of the up-regulated genes. C) STRING analysis (protein–protein interaction) of the DEGs under HDM treatment.
Table 3
Top 25 most up- and down-regulated DEGs in zebrafish larvae exposed to W-HDM for 10 days.

| Up-regulated genes | Complete name or Predicted description | FC | Down-regulated genes | Complete name or Predicted description | FC |
|--------------------|----------------------------------------|----|-----------------------|----------------------------------------|----|
| blast2 Go Unit     | SH3 domain and nuclear localisation   | 83.14 | Blast2 Go Unit        | AAA - ATPases associated with a variety of cellular activities | -54.27 |
| blast2 Go Unit     | eukaryotic translation initiation factor 3 subunit J | 70.51 | blast2 Go Unit        | rhamnose-binding lectin | -34.9 |
| blast2 Go Unit     | haemoglobin, beta adult 1              | 32.21 | blast2 Go Unit        | cytohesin-3 | -16.82 |
| blast2 Go Unit     | haemoglobin, beta adult 2              | 31.58 | blast2 Go Unit        | karyopherin alpha 7 | -9.09 |
| blast2 Go Unit     | haemoglobin, alpha adult 1             | 21.35 | blast2 Go Unit        | heat shock protein family A (Hsp70) member 12 A.1 | -7.35 |
| blast2 Go Unit     | ladderlectin                           | 20.51 | blast2 Go Unit        | heat shock protein family A (Hsp70) member 12 A.2 | -4.99 |
| blast2 Go Unit     | BX548011.4                             | 20.37 | blast2 Go Unit        | sperm adhesion molecule 1 | -4.91 |
| blast2 Go Unit     | haemoglobin subunit alpha              | 15.69 | blast2 Go Unit        | zona pellucida glycoprotein 3a, tandem duplicate 1 | -4.69 |
| blast2 Go Unit     | death-associated protein 1b            | 13.1 | blast2 Go Unit        | LSM family member 14B | -4.54 |
| blast2 Go Unit     | interlectin 2                          | 12.51 | blast2 Go Unit        | acid-sensing ion channel 1 | -4.17 |
| blast2 Go Unit     | dual specificity phosphatase 29        | 11.54 | blast2 Go Unit        | olfactomedin-like domain | -3.94 |
| blast2 Go Unit     | sgc:112408                             | 9.45 | blast2 Go Unit        | lymphocyte antigen 6 family member M6 | -3.38 |
| blast2 Go Unit     | Stomatlin                              | 8.04 | blast2 Go Unit        | ankyrin repeat, SAM and basic leucine | -3.31 |
| blast2 Go Unit     | kruppel-like factor 2a                 | 7.17 | blast2 Go Unit        | sgc:17581 | -3.08 |
| blast2 Go Unit     | BX088712.5                             | 6.92 | blast2 Go Unit        | phospholipase D family, member 6 | -2.77 |
| blast2 Go Unit     | nitrate 1                              | 6.37 | blast2 Go Unit        | mitogen-activated protein kinase kinase | -2.61 |
| blast2 Go Unit     | haemoglobin, alpha embryonic 5         | 6.21 | blast2 Go Unit        | sgc:61p9.7 | -2.47 |
| blast2 Go Unit     | elastin microfibril interface 3a       | 5.81 | blast2 Go Unit        | cilia- and flagella-associated protein 61 | -2.44 |
| blast2 Go Unit     | mutt homologue 4                       | 5.32 | blast2 Go Unit        | tripartite motif containing 35-27 | -2.28 |
| blast2 Go Unit     | cytochrome P450, family 2, subfamily K, | 5.22 | blast2 Go Unit        | zgc:17581 | -2.28 |
| blast2 Go Unit     | POS-like antigen 1a                    | 4.89 | blast2 Go Unit        | glutamate receptor, ionotropic, N-methyl D-aspartate 2B | -2.22 |
| blast2 Go Unit     | protein phosphatase 1 regulatory subunit 3 C-B | 4.31 | blast2 Go Unit        | |
| blast2 Go Unit     | interferon-induced very large GTPase 1 | 4.21 | blast2 Go Unit        | |
| blast2 Go Unit     | oestrogen-responsive finger protein     | 3.95 | blast2 Go Unit        | |
**Fig. 6.** Biological processes (BP), KEGG pathways, protein domains (PD) and protein–protein interactions enriched for genes modulated by the W-HDM treatment. A) Enrichment analyses of the down-regulated genes. B) Enrichment analyses of the up-regulated genes. C) STRING analysis (protein–protein interaction) of the DEGs under W-HDM treatment.
involved in the type I interferon system (Fig. 7C). In relation to the inflammation process, some genes were up- and down-regulated according to the face mask treatment, and a clear pattern was not observed; however, for the selected genes, the W-HDM treatment clustered together with the control, reflecting an overall lower impact of this treatment on inflammation than the other treatments (Fig. 7D). However, the analysis of the DEGs involved in antigen presentation and modulation at least in one of the treatments revealed that, in this case, W-HDM products had the highest effect, with an up-regulation of these genes after the treatment (Fig. 7E). An unexpected result was also the
high representation of haem-binding protein overexpression in the zebrafish larvae treated with the W-HDM products, which were minimally affected by the HDM treatment (Fig. 7F).

Because terms related to isoprenoid/sterol biosynthesis and redox processes were highly enriched for the PDM treatment, a heatmap representing the genes included in those terms was constructed. In this case, the genes were also almost unaffected by the HDM treatment, whereas some of them were induced by the W-HDM product, and a strong induction was observed in those larvae exposed to PDM products (Fig. 7G).

Fig. 8. Analysis of the DEGs involved in reproduction. A) Heatmap representing the mean TPM expression values of the DEGs with a role in reproduction across the different treatments. B) Schematic representation of the different reproduction steps with genes affected by the face mask treatments.
3.5. Face mask degradation products clearly impair the expression of genes involved in the reproductive process

Although only zebrafish larvae exposed to face mask fragments (PDM and HDM) showed an enrichment of reproductive processes for the down-regulated genes in the GO enrichment analyses, genes associated with reproduction were significantly regulated by the three treatments (PDM, HDM and W-HDM); (Fig. 8A)). The magnitude of the impact on the reproductive process showed an increasing tendency from the W-HDM, followed by PDM, to the HDM treatment, with more than 30 genes significantly affected by the HDM products (Fig. 8A).

The exposure of zebrafish larvae to the different face mask degradation products induced a generalised inhibition of a multitude of genes with a pivotal role in different stages of reproduction (gametogenesis, sperm-egg recognition, fertilisation and embryonic development) (Fig. 8B). Only efna5b (ephrin-a5), a gene involved in fertilisation and embryonic development, was significantly up-regulated by HDM treatment, although the protein encoded by this gene exerts a variety of functions independent of embryogenesis (Worku et al., 2018).

Regarding gametogenesis, genes involved in both male (spermatogenesis) and female (oogenesis) gamete formation were affected; this included genes with a direct role in meiosis (hom1a and smc1b), specific translational regulation in gametocytes (dazl and patl2), oocyte axis specification (buc) or sperm flagella development (cfap61), but most of the affected genes were involved in PIWI-interacting RNA (piRNA) biogenesis (piwil1, piwil2, mlf17, gsf11, fkp61, ddz4, tdr1, tdr3, tdr7a), which protect the germline from invasive transposable elements (Ishizu et al., 2012). Conversely, the DEGs belonging to the sperm-egg recognition process encode a variety of zona pellucida glycoproteins (zp2.1, zp2.2, zp2.3, zp2.5, zp3, zp3a.1, zp3a.2, zp3b, zp3f.1, zp3e, zp3.2_1, zpax1, zpax4), also known as sperm-binding proteins, since they mediate sperm binding to oocytes. Finally, genes with a role in fertilisation and embryogenesis were also down-regulated by the face mask degradation products.

4. Discussion

4.1. Surgical face masks are a material of emerging concern in aquatic environments

After face mask use, wastes are mismanaged on land (Aragaw, 2020; Dharmaraj et al., 2021), and their degradation can occur by means of different processes, such as weathering, ageing, and microbial degradation (Hasan et al., 2021). After fragmentation and degradation, small pieces of face masks are released into aquatic systems and can release microplastics and other substances, such as B, Al, Ti, Fe, Cu and Sr. The concentration of these compounds after 10 days of degradation with UVC in water did not increase much with respect to the controls, although the concentrations of the cited inorganic elements were dependent on the face mask degradation stage. Although the leaching of inorganic compounds is characteristic of the type of face mask (Sullivan et al., 2021), the degradation processes, the time in the aquatic media and the size of the face mask fragments are relevant in the leaching of chemical compounds from face masks into aquatic systems. According to the leached organic substances, only three compounds were identified as secondary metabolites of face mask degradation, which were not found in intact face masks. Therefore, these compounds are probably byproducts of the degradation of face masks in water. Two of the identified substances corresponded to heterocyclic aromatic compounds (2,4,6-trimethyl-pyridine and N,N-dimethyl 1-dodecanamine) that can be found in creosote-contaminated sites formed through thermal processes related to coal and fossil fuel (Johansen et al., 1997; Zemanek et al., 1997). In the case of 2,4,6-trimethyl-pyridine, this compound is found in various man-made compounds, such as dyes, industrial solvents, herbicides, pesticides, polymers and many natural metabolites (Stankeviciute et al., 2016). Most likely, 2,4,6-trimethyl-pyridine occurs from the degradation of olefin polymers. Polyolefins are commercial polymers that are more common and cheaper than others on the market. The trimethylpyridine and dimethylpyridine mixture represents an important part of the composition of polyolefins (Zhang et al., 2020).

The use of similar chemical substances, such as 2,4,6-trimethyl-triazine, in olefins offers highly crystalline honeycomb-like structures and displays well-defined nanofibrillar morphologies with uniform diameters (Zhang et al., 2020). Therefore, this structure could be perfect for an efficient face mask design (Ullah et al., 2020). In the case of the second organic compound identified, N,N-dimethyl 1-dodecanamine is identified in alkanes and olefins (Zhang et al., 2020). Luryldimethylamine oxide (lauryl amine oxide), also known as dodecyl dimethylamine oxide (DDAO), is an amine oxide with a C12 alkyl tail and is one of the most frequently used surfactants (Varade et al., 2005). The third compound found was alanine, N-methyl-N-methoxy carbonyl-undecyl ester, which is probably a byproduct of degradation or a propane compound, since its industrial application has not been assigned but could include anti-bacterial compounds or dyes. A recent paper showed that single-use surgical face masks can act as dye carriers (methylene blue, crystal violet and malachite green) in aquatic environments (Anastopoulos and Pashalidis, 2021).

4.2. Toxicogenomic impacts of face mask degradation products in zebrafish larvae

Although fragments of face masks do not seem to be a high source of inorganic and nonpolar pollutants leaching into aquatic systems, the face mask fragments and fibres (mainly composed of plastic polymers) released from their degradation could be a source of emerging pollutants. Although face masks have not yet been of special concern, they could have environmental and health impacts on aquatic organisms exposed to these materials in specific locations. Currently, the concentration of face mask degradation products in aquatic ecosystems is not known; therefore, this study is based on the microplastic concentration already known in natural compartments (Koutnik et al., 2021) and used in previous works (Sendra et al., 2019a,b, 2021b).

The degradation stage of the surgical face masks influences their effects, since smaller fragments are more bioavailable for organisms (Wang et al., 2020). This finding could explain the higher gene modulation induced by HDM products on zebrafish larvae as determined by RNA-Seq analysis. However, differential responses were observed for a variety of processes, with certain gene groups remaining unaffected by HDM treatment but being strongly regulated by the PDM and W-HDM treatments. This result could probably be a consequence of the presence of different byproducts generated during the degradation process of the face mask fragments, which could undergo modifications over time. These modifications are probably accelerated with the HDM and W-HDM treatments compared with the PDM due to the higher initial fragmentation. However, the presence of face mask fragments in the HDM and PDM treatments compared with the W-HDM treatment could allow an accumulation of microfibres and probably a higher variety of pollutants directly ingested by larvae. Each treatment in this study showed a particular profile of biological processes and pathways affected, offering a specific footprint in gene regulation. However, a common effect among the three treatments was the down-regulation of genes involved in reproductive processes. Under environmental conditions, the three stages of face mask degradation could be found in nature; therefore, more complex biological processes could be found compared with the results obtained in this work.

In this work it was revealed that under the selected conditions, exposure of zebrafish larvae to face mask degradation products could provoke effects on the immune response since, as determined by RNA-Seq analysis, several immune-related genes were affected by the three treatments. Xenobiotics frequently found in natural compartments can significantly suppress or enhance immune responsiveness, provoking immunotoxicity (Kreitinger et al., 2016). While W-HDM and PDM were...
able to induce the expression of genes involved in certain aspects of the immune response, such as antigen presentation or inflammation-related genes. HDM treatment suppressed the expression of genes involved in complement, coagulation and antiviral responses and dysregulated the expression of several genes with a role in inflammation. Moreover, numerous xenobiotics can also bind haemoglobin members via non-covalent interactions and can affect their function through alteration of the equilibrium between oxygenated and deoxygenated haemoglobin (Gaurav et al., 2021), which could explain the up-regulation of different haemoglobin members in the W-HDM treatment.

A mechanism to fight against xenobiotics is the activation of the KEAP1-NRF2 system, which is involved in the antioxidant enzyme activity of phase II transformation of xenobiotics. In the larvae treated with PDM products, induction of certain genes related to the immune response was observed, which could be in accordance with an up-regulation of genes involved in redox metabolism. The anti-inflammatory gene irg11 (immunoreponsive gene 1-like), which transforms cis-aconitate into aconitate, is involved in the activation of the KEAP1-NRF2 system, and this gene showed a higher expression in larvae treated with PDM, especially in larvae exposed to W-HDM products. Therefore, the higher expression of redox-related genes and sterol biosynthesis (which requires oxygen radicals for their correct functioning) in those treatments could be related to the modulation of the immune genes observed in those groups of treatments (Spann and Glass, 2013). Moreover, some of the enzymes involved in sterol biosynthesis (cytochrome P450 enzymes) have a role in xenobiotic metabolism (Esteves et al., 2021; Lepesheva et al., 2008; Stravoravdis et al., 2021). Previous works have reported specific plastic additives as plasticizers, flame retardants and biocides as endocrine disruptors that mimic natural steroid hormones (Howdeshell et al., 2007; Sendra et al., 2020), which could also be a reasonable explanation for the modulation of genes involved in steroid biosynthesis by PDM treatment. Interestingly, although endocrine disruptors can interfere with the synthesis and activity of sex steroid hormones and, consequently, lead to reproductive disorders at different levels (Tyler et al., 1998), the treatment showing the lowest alteration of genes involved in steroid biosynthesis (HDM) was the same, inducing the highest effect on reproduction-related genes. Therefore, other routes, not only sterol pathways, could induce impairments in the expression of genes related to reproductive processes.

Despite all the effects provoked by face mask products at transcriptome level, reproductive processes showed the most remarkable changes, as determined by RNA-Seq analysis of zebrafish larvae exposed to the three treatments (PDM, HDM and W-HDM) and compared with the control larvae. Although enriched GO terms related to reproduction were only observed for the down-regulated genes in the PDM and HDM treatments, some of the genes included in those categories were also down-regulated in the W-HDM treatment. A clear increasing effect on reproduction was observed with the different treatments as follows: W-HDM<PDM<HDM. In addition to gametogenesis (with representative genes for both oogenesis and spermogenesis) and embryogenesis, genes involved in fertility and reproductive processes were significantly down-regulated in the three steps of fertilisation Step 1; species-specific binding of sperm to eggs, Step 2; zona pellucida proteins, Ca²⁺, G proteins and the acrosome reaction, and Step 3; penetration of the egg zona pellucida by sperm (Wassarman et al., 2001).

Gametogenesis-related genes seem to be highly impacted by face mask treatments, with down-regulated genes involved in meiosis, transcription regulation in gametogenesis, and axis specification of sperm flagella development. However, most gametogenesis-affected genes are related to PIWI-interacting RNA (piRNA) biogenesis. piRNAs are a class of small noncoding RNAs that form the piRNA-induced silencing complex (piRISC) in the animal germ line (Siomi et al., 2011), which protects the integrity of the genome from invasive transposable elements (Ishizu et al., 2012; Siomi et al., 2011). piRNAs are tightly bound to PIWI proteins of the Argonaute family, and these can interact with many other proteins required for PIWI function, regulating piRNA-mediated silencing (Siomi et al., 2011). This phenomenon is observed for Tudor-domain-containing proteins (TDRDs) (Ishizu et al., 2012), but several non-PIWI, non-TDRD proteins are also involved in piRNA biogenesis, such as DDX4 (VASA), FKBP6 or GTSF1 (Donertas et al., 2013; Ishizu et al., 2012). In this work, we observed a drastic down-regulation of PIWI-like genes (piwil1, piwil2), TDRD genes (tdrd1, tdrd5, tdrd7a, rnf17 -tdrd-4), ddx4, fkbp6 and gtsfl genes, which suggested a severe impairment of piRNA-mediated silencing function. Consequently, based on gene expression analyses, gametogenesis disorders could be expected in zebrafish exposed to face mask degradation products. Nevertheless, further investigations are needed to validate this finding.

Among the genes involved in the reproductive process and affected by the treatments, we found a multitude of zona pellucida glycoproteins. Whereas mammals possess four genes encoding different types of zona pellucida proteins (ZP1, ZP2, ZP3 and ZP4), the copy number of these genes has suffered an expansion in teleosts. This evolutionary process, called concerted evolution, is probably a consequence of the reproductive strategy of fish, which need to produce eggs with high frequency and in large numbers (Mold et al., 2009). ZP proteins have been conserved in mammalian and nonmammalian eggs over the course of several hundred million years of evolution. They play vital roles during oogenesis, fertilisation and preimplantation development (Wassarman, 2008), but they also participate in several steps in the fertilisation pathway, such as receptors for sperm, inducers of sperm exocytosis and the acrosome reaction (Florman et al., 2010; Wassarman et al., 2001). Although some of the genes mentioned above also have a role in fertilisation, certain genes with a specific function in embryogenesis were also affected by the treatments, including zar1 (zygote arrest protein 1), an oocyte-specific gene that encodes a protein that is thought to function in the initiation of embryogenesis (Wu et al., 2003).

In mammals, most of the abovementioned proteins are produced in reproductive organs; however, in teleosts and in many other oviparous vertebrates, the major constituents of the eggs are synthesised in the liver and transported to the oocyte for uptake (Arukke and Goksøyr, 2003). The liver is the main organ for detoxification of xenobiotics; therefore, damage to this organ due to face masks with plastic additives as endocrine disruptors (Sendra et al., 2020) could provoke impairments in reproductive, metabolic and signalling processes (Qiao et al., 2016), as observed in the transcriptome of zebrafish larvae exposed to PDM and HDM treatments. The potential impairment of reproduction by face mask products could include effects at the individual but also at the population level, since fertility or reproductive success can compromise the population (Mathieu-Denoncourt et al., 2015). Furthermore, these changes could be transferred to offspring. From the beginning of this century, there is evidence that epigenetic processes play a pivotal role in regulating many genes involved in the endocrine system. There were modulated genes in this work belonging to the cytochrome P450 family, which are regulated by DNA methylation (an epigenetic mark); consequently, they could impact the major endocrine pathways in which they are involved (Jacobs et al., 2017; Rosenfeld, 2019). Disturbances in meiotic epigenetic inheritance were observed in enriched biological processes in zebrafish exposed to HDM, which could impact further generations owing to the involvement of the germ line. Environmentally induced alterations in the germline, whether genetic or epigenetic, will be transmitted to subsequent generations and could induce altered phenotypes in progeny and therefore exert effects at the population level (Jacobs et al., 2017). Alterations in the reproduction process and fitness were observed in natural populations of different animals exposed to plastic waste (Patrício Silva et al., 2021), which could be an indicative of similar effects for face mask degradation products. Indeed, reproduction is one of the sublethal effects induced by toxic chemicals more commonly extrapolated to a population level, since reproduction integrates the outcome of many interacting biological process (Martin et al., 2014).

This work only shows the complexity and effects of face mask...
products as emergent pollutants in aquatic organisms at transcriptome level. However, face mask fragments and their leaching products coexist with other environmental xenobiotes, such as metals, antibiotics, nanoparticles, pesticides, veterinary drugs, chemical compounds used in industry and personal care products. Therefore, the additive and synergistic effects of all these pollutants have been unknown until now. Knowing the behaviour and biological effects of face mask products in combination with other xenobiotes could provide more real insight into the processes occurring in natural environments.

5. Conclusions

This study reveals, at the transcriptome level, the impact of the most commonly used PPE over the COVID-19 pandemic (face mask products) in the model organism zebrafish. As hypothesised, surgical face masks can release inorganic and organic compounds in aquatic systems, and the release of these compounds could be related to the degradation stage of face masks. The degradation stages are associated with the size of face mask fragments, showing different toxicogenomic effects according to each treatment. Despite face mask fragments coming from degradation, water that has been in contact with face masks could leave a footprint in aqueous media with unwanted effects in aquatic organisms. Although W-HDM showed significant toxicogenomic effects in zebrafish larvae, RNA-Seq analysis confirmed that those treatments containing face mask fragments (PDM and HDM) induced a more severe effect. Although not restricted to, reproductive process was the most remarkable function affected in zebrafish larvae exposed to face mask degradation products. Among the different treatments, HDM products induced the most powerful effect, with the down-regulation of a multitude of genes with a pivotal role in different steps of the reproductive process, from gametogenesis to embryogenesis.

The results suggest that the biological processes affected by face mask fragments could not only occur at the individual but also at the population level through fertility and reproductive impairments. However, these effects will depend on the stage of degradation of the surgical face masks.

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CRediT authorship contribution statement

Marta Sendra: Investigation, Writing – original draft. Patricia Pereiro: Investigation, Writing – review & editing. Pilar Yeste: Investigation, Writing – review & editing. Beatriz Nova: Conceptualization, Supervision, Writing – review & editing, Funding acquisition. Antonio Figueras: Conceptualization, Supervision, Methodology, Software, Writing – review & editing, Funding acquisition.

Declaration of Competing Interest

The authors declare that they have no known competing financial interests or personal relationships that could have appeared to influence the work reported in this paper.

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Appendix A. Supplementary material

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