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Cellular responses of Pacific oyster (*Crassostrea gigas*) gametes exposed *in vitro* to polystyrene nanoparticles

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Abstract:

While the detection and quantification of nano-sized plastic in the environment remains a challenge, the growing number of polymer applications mean that we can expect an increase in the release of nanoplastics into the environment by indirect outputs. Today, very little is known about the impact of nano-sized plastics on marine organisms. Thus, the objective of this study was to investigate the toxicity of polystyrene nanoplastics (NPs) on oyster (*Crassostrea gigas*) gametes. Spermatozoa and oocytes were exposed to four NPs concentrations ranging from 0.1 to 100 mg L-1 for 1, 3 and 5 h. NPs coated with carboxylic (PS-COOH) and amine groups (PS-NH2) were used to determine how surface properties influence the effects of nanoplastics. Results demonstrated the adhesion of NPs to oyster spermatozoa and oocytes as suggested by the increase of relative cell size and complexity measured by flow-cytometry and confirmed by microscopy observations. A significant increase of ROS production was observed in sperm cells upon exposure to 100 mg L-1 PS-COOH, but was not observed with PS-NH2, suggesting a differential effect according to the NP-associated functional group. Altogether, these results demonstrate that the effects of NPs occur rapidly, are complex and are possibly associated with the cellular eco-corona, which could modify NPs behaviour and toxicity.

Highlights

▶ Nanoplastics attach to both oocytes and spermatozoa. ▶ Cellular impacts of NPs was observed on spermatozoa. ▶ PS-COOH exposure generated a dose-response increase in ROS production in spermatozoa. ▶ Higher impact of PS-COOH suggest an influence of particle surface properties.

Keywords: Nanoplastics, Oysters, Gametes, Cellular responses

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1. Introduction

Each year, more than 320 million tons of plastic are produced in the world (Plastic
Europe; 2016), and a large part of this finishes as waste in the oceans: an estimated 4.8-
12.7 million tons in 2010 (Jambeck et al., 2015). There is evidence of small particles of
plastic known as microplastics (MPs, <5mm) in global waters (Barnes et al., 2009).
However, because their identification is difficult, mainly due to limitations of
conventional sampling methods (sampling nets have a mesh size $> 300 \ \mu m$) and lack of
analytical techniques (Koelmans et al., 2015), little attention has been paid to smaller
particles of plastic known as nanoplastics (NPs) (GESAMP, 2016). Plastic particles are
considered nano in the strict sense, (i.e., defined in the same way as for non-polymer
nanomaterials in the field of engineered nanoparticles) if they are <100 nm in at least
two of their dimensions (Klaine et al., 2012; Koelmans et al., 2015). Nevertheless, the
NPs definition was enlarged to all polymeric particles <1000 nm (in at least one of its
dimensions) by the GESAMP (2016) because of their colloidal behaviour. Nanoplastics
can be found as part of many applications such as drug delivery, cosmetics, biosensors,
photonics, nanocomposites, paints, adhesives and coatings between others (Ganajan and
Tijare, 2018; Guterres et al. 2007; Hernandez et al., 2017; Merinska and Dujkova, 2012;
Rogach et al., 2000; Velev and Kaler, 1999). With the number of polymer applications
growing each year, it is estimated that, by 2020, NPs will account for most
nanomaterials on the market (Fabra et al., 2013) making the release of NPs by indirect
outputs an environmental concern. Small microplastics collected at sea have recently
been characterized (Ter Halle et al., 2017) and demonstrated to be increasingly
abundant following a power-law increase with decreasing particle size in sea surface
samples (Erni-Cassola et al., 2017).

59 Environmental effects of NPs are linked to their intrinsic features such as surface 60 charge, size, shape, functionalization and coating (Klaine et al., 2008). In addition, the interaction of NPs with biological cells will be affected by their dispersion, aggregation 61 62 and agglomeration behaviour, which are dependent on the physico-chemical parameters of the surrounding media (pH, temperature and ionic concentration) as well as the 63 presence/absence of natural colloids (Canesi et al., 2017). Thus, it is essential to 64 understand how NPs behave in the marine environment and how they interact with 65 marine organisms. 66 The Pacific oyster, Crassostrea gigas, has a worldwide distribution and has the highest 67 annual production of any aquatic organisms (around 630 000 tons registered worldwide 68 69 in 2014), far exceeding other molluscs (mussels, clams, cockles, etc.) (FAO statistics data, 2016). During spawning, oysters release their gametes into the surrounding waters 70 where fertilization takes place. Consequently, oocytes and spermatozoa are very 71 vulnerable as they will be exposed to a wide range of environmental stressors present in 72 the surrounding water, notably pollutants including micro- and nano-sized plastics, 73 expected to be abundant in the water column (GESAMP, 2016). The effect of MPs on 74 these organisms has recently been published. Sussarellu et al., (2016) showed a 75 significant decrease of oocyte number and spermatozoa and oocyte quality upon 76 77 polystyrene microplastic exposure of adults during gametogenesis, leading to a negative impact on D-larval yield and larval growth of experimental offspring. Working at early 78 79 development stages, Cole and Galloway (2015) showed that C. gigas larvae can readily ingest small plastic particles (<20 µm) and internalized nano-sized plastics (<1 µm) 80 without any significant reported impacts. Both studies illustrated the ingestion of small 81 plastics by adults and early developmental stages. However, there is no information 82 83 available regarding the impact of small plastics, especially nano-sized plastics, on early

84	free living cells like oyster gametes. This is crucial information considering that the
85	quality of spermatozoa and oocytes is essential for successful fertilisation (Boulais et
86	al., 2015; 2017; Suquet et al., 2010). Upon spawning, fertilisation can occur within a
87	fairly long period (a few hours) as spermatozoa movement can be maintained up to 24 h
88	(Suquet et al., 2010) making this gametogenic phase very sensitive to waterborne
89	pollutants, especially in estuarine and coastal marine habitats. In these ecosystems,
90	environmental degradation is substantial, waters are greatly influenced by increased
91	human expansion, and we may expect peaks of nano-sized plastics close to industrial
92	sources, as already been observed for microplastics (Filella, 2015; Lambert and Wagner,
93	2016).
94	The objective of this study was to investigate interactions between polystyrene
95	nanoplastics and oyster gametes. Nanoplastics exhibiting different surface
96	functionalization with carboxylic groups (COOH) or primary amine (NH ₂), as anionic
97	and cationic NPs, were used to assess the effects of particle surface properties, NPs
98	behaviour in seawater and ultimate toxicity. Due to the lack of information on
99	concentrations of nano-size particles in the environment, spermatozoa and oocytes were
100	exposed separately to a wide range of concentrations (0.1, 1, 10 and 100 mg L ⁻¹ of PS-
101	NH ₂ or PS-COOH) in order to identify a possible toxicity threshold. Nano-PS size,

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2. Materials and methods

cytometry at 1, 3 and 5 h of NPs exposure.

monitored by Dynamic Light Scattering (DLS) during the experiment. Spermatozoa

motility and cellular responses of spermatozoa and oocytes in terms of viability, cellular

characteristics and Reactive Oxygen Species (ROS) production were monitored by flow

109	2.1. Measurements	of nanoplastic	characteristics
		- I	

110	Fluorescent-green 100 nm amino (PS-NH ₂) and carboxylic (PS-COOH) polystyrene
111	nanoparticles were purchased from Micromod laboratories (Germany) with an
112	Excitation/Emission: 475 nm/510 nm. To avoid commercial artefacts, nanobeads were
113	purchased from the same company and only differed in their functional groups. For
114	Dynamic Light Scattering analysis (DLS) the stock solutions (10 g L ⁻¹) were diluted in
115	0.2-µm filtered seawater (FSW) to a concentration of 100 mg L ⁻¹ . Size (Z-average),
116	charge and aggregation state (polydispersity index, PDI) of nanoplastics were
117	determined using a Zetasizer NanoZS (Malvern, United Kingdom). , in MilliQ water in
118	triplicate, each replicate corresponding to 13 runs for the Z-average and 40 runs for Z-
119	potential, following the protocol described in Della Torre et al. (2014). Data were
120	analyzed using Zetasizer Nano Series software, version 6.20. The nanoplastics were also
121	observed by transmission electronic microscopy (JeolJEM 100 CX II) to verify particle
122	size. In brief, nanoplastics were diluted in MilliQ (100 mg L ⁻¹) and placed on a copper
123	grid (400 nm mesh) with a carbon-coated Formvar film (Polysciences) and marked with
124	2% (wt/vol) uranyl acetate to be measured.
125	To check the behaviour of NPs under experimental conditions, DLS measurements were
126	also performed on particles incubated in FSW and 0.2 µm filtered seminal media in the
127	same conditions as those used during the experiment, i.e., kept in continuous movement
128	using a Stuart rotator SB3A. Filtered seminal media were prepared by filtering
129	spermatozoa (1×10^7 cells mL ⁻¹ in FSW) and oocyte (2×10^5 cells mL ⁻¹ in FSW) solutions
130	on a 0.2 μm filter. Nano-PS concentration was adjusted to 100 mg L ⁻¹ in both FSW and
131	seminal media. Measurements were performed after 1, 3 and 5 h using a Zetasizer Nano
132	as described above.

- 133 2.2. Gamete collection
- Mature oysters from the north coast of Brittany (Aber Benoit: 48° 27′ 31″ N, 4° 20′ 42″
- W) were harvested and acclimated for 1 week before gamete collection in laboratory
- conditions [1 µm filtered seawater, 18°C, continuous feeding on a mixed diet of two
- 137 microalgae Chaetoceros gracilis (UTEX LB2658), T-Isochrysis (clone: T-iso;
- 138 CCAP927/14)] to provide good physiological conditions prior spawning. The natural
- seawater used for gamete resuspension and exposure to nanoplastics was UV-treated,
- 140 filtered at 0.2 µm and maintained at T = 18°C, salinity 38, pH 8.32 and conductivity 6
- 141 S/m.
- Spermatozoa and oocytes were collected from individual males (n = 6; length: $110.9 \pm$
- 9.7 mm) and individual females (n = 6; length: 129.5 ± 8.9 mm) by stripping according
- to Steele and Mulcahy (1999). Sperm motility was checked under a microscope. Final
- concentration of 1×10^7 cells mL⁻¹ and 2×10^5 cells mL⁻¹ was adjusted with FSW for each
- individual male and female, respectively.
- 2.3. Gamete exposure to nanoplastics
- Oyster' gametes from males (n=6) and females (n=6) were individually exposed to NPs.
- Four different concentrations of PS-COOH or PS-NH₂ were tested separately on oocytes
- and spermatozoa: Control (no NPs), 0.1, 1, 10 and 100 mg L⁻¹ of PS-COOH or PS-NH₂
- which correspond to 0, 1.9×10^1 , 1.9×10^2 , 1.9×10^3 and 1.9×10^4 particles spermatozoa⁻¹;
- and 1.9×10^3 , 1.9×10^4 , 1.9×10^5 and 1.9×10^6 particles oocyte⁻¹, respectively. Samples
- were kept in continuous movement (program 3) using a Stuart rotator SB3 (Cole-
- Parmer, UK) in order to prevent cell sedimentation, in a dark room at 18°C. Sampling
- was performed in fresh samples after 1, 3 and 5 h exposure to NPs. Additionally, after

- 3h exposure, control and exposed samples of oocytes and spermatozoa were fixed in
- formaldehyde (3% final) for later microscopy observations.
- 158 2.4. Analyses
- 159 2.4.1. Microscopy
- 160 2.4.1.1. Spermatozoa motility
- 161 To measure spermatozoa motility, fresh sperm solution was mixed with FSW
- 162 containing pluronic acid (PA 1g L⁻¹) (vol:vol) and transfer to a FastRead cell. The
- percentage of motile spermatozoa, their movement linearity and velocity (VAP:
- Velocity of the Average Path; μm sec⁻¹) were measured using CASA plug-in for the
- 165 Image J software adapted for Pacific oyster spermatozoa according to Suquet et al.
- 166 (2014) under a microscope (dark field, Olympus B×51, ×20 magnification), connected
- to a video camera (Qicam Fast, 60 frames sec⁻¹).
- 168 2.4.1.2. Visualization of NP-gamete interactions
- Once fixed in formaldehyde, both cell types emitted green fluorescence coming from
- the fixative (formaldehyde). In order to better localize cells and discriminate the
- 171 fixative' green fluorescence from the diffuse green fluorescence of NPs, cell nucleus
- were labelled in blue using DAPI ((4',6-diamidino-2-phenylindole), which emits blue
- 173 fluorescence upon binding to DNA. Cells were visualized using a confocal laser
- scanning microscope Zeiss Axio Observer Z1 coupled to a ZEISS LSM780 confocal
- laser module, with a Plan-Apochromat 63x/1.40 oil DIC M27 objective. Images were
- made using two channels: ChS1-T1 (Ex/Em: 488/599 nm) and Ch1-T2 (Ex/Em:
- 177 405/449 nm). Images of spermatozoa 16.6×16.6 µm in size (X-scaling × Y-scaling)
- were acquired at a sampling speed of 6.7 µs/pixel and a zoom of 8.1. Images of oocytes
- 179 $112.4 \times 112.4 \,\mu m$ in size (X-scaling × Y-scaling) were acquired at a sampling speed of

1.8 µs/pixel and a zoom of 1.2. The images were obtained using ZEN 2012 SP2 180 software (Carl Zeiss Microscopy GmbH, Germany). 181 2.4.2. Flow cytometry 182 Analyses were performed using an EasyCyte Plus cytometer (Guava Technologies, 183 Millipore, Billerica, MA), equipped with a 488-nm argon laser and three fluorescence 184 185 detectors: green (525/30 nm), yellow (583/26 nm) and red (680/30 nm). Flow cytometry analysis were performed in fresh samples with or without fluorescent probes as 186 described below. To run the different tests, cytometer' setting were established using 187 unlabelled cell population (negative control) for which was assigned a relative (low) but 188 non-zero fluorescence value. 189 2.4.2.1. Cell number, relative size and complexity 190 Spermatozoa and oocyte cell numbers were recorded in fresh samples without adding 191 fluorescent probe with and without NPs, over 30 sec at 0.12 µL sec⁻¹ flow rate for 192 spermatozoa and at 0.59 µL sec⁻¹ for oocytes. Cells were detected and described on the 193 194 flow-cytometer according to their optical characteristics obtained at small angle 195 (Forward Scatter: FSC) and large angle (Side Scatter: SSC) giving respectively information on relative size and complexity of cells. For analysis of sperm samples, two 196 regions were designed to discriminate single and aggregate spermatozoa. 197 2.4.2.2. Cell mortality 198 Viability was measured after 10 minutes of incubation with SYBR-14 for spermatozoa 199 and SYBR-green for oocytes (final concentration: 1/100 commercial solution) together 200 with propidium iodide (PI, final concentration: 10mg L⁻¹) following the protocols 201 202 established by Le Goïc et al. (2013; 2014) for spermatozoa and oocytes, respectively. PI penetrates cells that have lost membrane integrity and are considered to be dead 203

(orange/red fluorescence), whereas SYBR-green, which binds DNA, penetrates both

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205	dead and living oocytes (green fluorescence, SYBRgreen fluorescence later in the text)
206	and SYBR-14 binds DNA of live spermatozoa only. The percentage of dead oocytes
207	and spermatozoa was calculated by the ratio between the number of cells showing
208	orange/red fluorescence (PI) and the total number of cells \times 100. Results were
209	expressed as percentages of dead cells.
210	A strong decrease in viability was observed in control oocytes at 5 h incubation, which
211	is consistent with the short life cycle of oocytes. In consequence, after 5 h exposure all
212	samples of oocytes were excluded from the data.
213	2.4.2.3. Reactive Oxygen Species (ROS) production
214	ROS production by cells was measured using 2,7-dichlorofluorescein diacetate (DCFH-
215	DA) after 50 min of incubation at 18°C. DCFH is hydrolysed (esterase) intracellularly to
216	form DCF, which turns fluorescent green upon oxidation with ROS. The green
217	fluorescence measured is quantitatively related to the ROS production in cells and was
218	expressed in arbitrary units (A.U.).
219	2.3.2.4. Flow cytometry data normalization
220	After exposure to both NPs, green fluorescence level of unlabelled cells were higher in
221	the presence of nanoparticles (exposed cells) than in their absence (control cells). Thus,
222	fluorescence values of negative controls from both, spermatozoa and oocytes, were
223	subtracted from the labelled cells' fluorescence data obtained after addition of the
224	different fluorescent probes: DCFH-DA (ROS production) and SYBRgreen/SYBR14
225	(viability). This was done to avoid any bias due to an increase in the cells' background
226	green fluorescence observed during PS-COOH and PS-NH ₂ exposure.
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228	2.5 Statistical analyses

Statistical analyses were performed using STATGRAPHICS Centurion XVII and were carried out separately for PS-COOH and PS-NH $_2$ conditions treatments. When the requirements of normality and homogeneity of variances were met, one-way ANOVAs were performed to establish significant differences between treatments at each sampling time. Tukey's post-hoc test was used to test for differences among the different exposure treatments with P < 0.05 as the significance level for all analyses. Arcsine or log transformation was performed when necessary to meet the normality and homoscedasticity criteria. Non-parametric analysis (Kruskal-Wallis test) was performed when the variables did not meet the requirements of ANOVA. In addition, simple regressions using the coefficient of determination (R^2) were used to investigate the relationship between spermatozoa motility parameters and ROS production, with P < 0.05 as the significance level for all analyses.

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3. Results

- 3.1. Nanoplastic size, charge and aggregation under experimental conditions
- DLS analysis confirmed the outlined size of NPs in MilliQ water, with a Z-average of
- 245 131.82 ± 0.23 nm (mean \pm SD) and 130.83 ± 0.32 nm, for PS-COOH and PS-NH₂,
- respectively (Figure 1A-B). The expected size of NPs was also observed in FSW with
- 140.11 \pm 0.92 nm and 141.13 \pm 1.50 nm for PS-COOH and PS-NH₂ respectively.
- 248 Aggregation was negligible as suggested by a PdI < 0.2 for both particle types in MilliQ
- 249 and FSW conditions. Nanoparticle geometric diameter was also confirmed using
- 250 transmission electron microscopy (Fig. 1C-D).
- 251 Results of NPs size and aggregation state monitored under constant rotation (to
- reproduce the incubation conditions) in FSW and in seminal media did not show any

- 253 changes in size or aggregation, regardless of the particle type, media or incubation time
- 254 (PdI < 0.2).
- Both NPs resuspended in FSW showed a negative charge: -2.89 ± 7.21 mV and $-5.69 \pm$
- 3.65 mV for PS-COOH and PS-NH₂, respectively. Similar behaviour was observed in
- oocyte and spermatozoa media, with charges of -8.17 ± 4.66 and -13.60 ± 2.92 for PS-
- 258 COOH and values of -8.26 \pm 3.20 and -6.95 \pm 1.37 mV for PS-NH₂, respectively.
- 3.2. Visualization of interactions between NPs and oyster gametes
- 260 Images of spermatozoa and oocytes exposed to the highest concentration of PS-COOH
- are shown in Figure 2 alongside the controls. Only particles displaying significant
- 262 biological effects (PS-COOH) were measured by microscopy. In order to localize
- 263 cellular structures and discriminate from fixative' fluorescence (green also) and
- 264 nanoplastics' fluorescence (green), cells were labelled with DAPI which allow staining
- nucleus DNA in blue (Fig. 2A; 2B). As a result, it was possible to localize NPs
- aggregates attached to the cells (Fig. 2C and 2D). In spermatozoa, aggregates of NPs
- were found attached mainly to the acrosome (Fig. 2C). Interaction of NPs with oocytes
- is shown in Fig. 2D, with oocytes covered by a semi-transparent structure formed by
- 269 colloids and debris present in the medium. The aggregates of NPs appeared to be
- entrapped inside this structure and not directly attached to the cell.
- 3.3. Cellular responses of gametes exposed to NPs
- 272 3.3.1 *Cell number, relative size and complexity*
- 273 After 3 h and 5 h of exposure to 100 mg L⁻¹ PS-COOH, a significant decrease in the
- 274 number of single spermatozoa was observed: 32% and 24% lower than the control (P
- 275 <0.01 and P <0.05), respectively. This decrease was directly correlated with the increase</p>

in spermatozoa' aggregates for this NP condition (P < 0.05 and P < 0.01 after 3h and 5h 276 exposure to 100 mg L⁻¹ PS-COOH). No significant formation of spermatozoa' 277 aggregates was observed in treatments exposed to PS-NH₂, and consequently, no 278 differences in single spermatozoa number was reported. Spermatozoa exposed to the 279 highest concentration of PS-COOH and PS-NH₂ showed a 4-5% increase in relative size 280 compared with controls after 1, 3 and 5 h exposure. Spermatozoa also showed higher 281 cellular relative complexity upon exposure to 10 and 100 mg L⁻¹. An increase of 24%, 282 25% and 25% cellular complexity at 10 mg L⁻¹ and 53%, 24% and 52% increase at 100 283 mg L⁻¹ was observed after 1, 3 and 5 h exposure to PS-COOH. Similar tendency was 284 observed upon exposure to PS-NH₂ that revealed a 14%, 15% and 19 % increase at 10 285 mg L⁻¹ whereas increased values of 59%, 37% and 62% were observed at 100 mg L⁻¹ 286 after 1, 3 and 5 h exposure, respectively. By contrast, no significant changes in oocyte 287 288 cell number, relative size or complexity were observed by flow cytometry in the whole experiment. 289

290 3.3.2 *Cell mortality*

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After exposure to the highest NP concentration (100 mg L⁻¹), oocytes exposed to both NPs types showed a decrease in their SYBRgreen fluorescent signal for both exposure times 1 and 3 h (P < 0.001 and P < 0.01 for PS-COOH and PS-NH₂, respectively at both time points). Due to the significant decrease of SYBRgreen fluorescent observed in all females (e.g. Supplementary figure 1 for one female), fluorescent value of PI cannot be considered either. Thus, viability test for this NP condition is not conclusive and cannot be considered. For the other 3 NP exposures (0.1, 1 and 10 mg L⁻¹) where the values of SYBRgreen and PI fluorescence were validated, no significant differences in the percentage of dead cells were observed in exposed oocytes regardless of exposure duration.

- No significant differences were observed in the percentage of dead cells in spermatozoa
- exposed to NPs among sampling times (from 1 to 5 h exposure) for none of the tested
- 303 NPs.
- 304 3.3.3 ROS production
- Control spermatozoa showed 17.1 \pm 5.7 (CV=33.8), 24.3 \pm 8.2 (CV=33.7) and 22.4 \pm
- 9.7 (CV=43.55) A.U. of ROS production at 1, 3 and 5 h, respectively. After 1 h of
- 307 exposure to PS-COOH, a dose-response increase in ROS production was observed in
- spermatozoa exposed to 1, 10 and 100 mg L^{-1} (P <0.001) that was 17.4%, 59.4% and
- 309 121% higher respectively than in the control. After 3 and 5 h, significant increases were
- only observed in spermatozoa exposed to 10 and 100 mg L^{-1} (P < 0.001 and P < 0.01 for
- 311 3 and 5 h, respectively) with ROS values around 30% and 70% higher than in control
- 312 (Figure 3). Spermatozoa exposed to PS-NH₂ were not significantly affected.
- 313 ROS production in oocytes was not significantly affected by NP exposure. Levels of
- ROS production in control oocytes ranged from 8.7 to 44.3 A.U. (24.9 \pm 11.2 A.U.; CV
- $= 121.5 \text{ and } 20.6 \pm 12.8 \text{ A.U.}$; CV = 126.3 after 1 and 3 h of incubation, respectively)
- showing a high inter-individual variability in stress response.
- 3.3.4. Spermatozoa motility
- No significant differences were observed in the percentage of motile spermatozoa, VAP
- or movement linearity after exposure to NPs. Nevertheless, when spermatozoa were
- 320 exposed to PS-COOH, a significantly positive correlation was found between the
- 321 percentage of motile spermatozoa and spermatozoa' ROS production through the
- 322 experiment ($R^2 = 0.12$, P < 0.001), particularly after 5 h exposure ($R^2 = 0.24$, P < 0.001).

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4. Discussion

325	Nanoplastics had similar size and charge in seawater and seminal media over time
326	The most important feature of nanoplastics is their high surface area-to-volume ratio
327	(Mattsson et al., 2015), which makes them highly reactive and prone to strongly interact
328	with biological membranes (Rossi et al., 2014). In general, nanoplastics, as hydrophobic
329	particles, are not thermodynamically stable and tend to aggregate very easily, forming
330	agglomerates. However, they can become soluble if they are charged since electrostatic
331	repulsions fight against attractive van der Waals forces as explained by the classic
332	Derjaguin-Landau-Verwey-Overbeek (DLVO) theory (Derjaguin, 1941; Verwey et al.,
333	1999). In the present work, oyster gametes were exposed to free single NPs of 100 nm,
334	as nanoplastics did not aggregate in seawater or seminal media over time. Our results
335	are in agreement with Cai et al. (2018) which recently reported absence of aggregation
336	in 100 nm polystyrene nanoplastics in NaCl and CaCl ₂ solutions charged with humic
337	substances. This behaviour was explained by the fact that electrostatic repulsive forces
338	increased in presence of organic matter in these solutions. Nevertheless, Cai and co-
339	authors highlighted that aggregation behaviour of nanoplastics is influenced by complex
340	environmental factors and further research is necessary to understand these interactions.
341	In the present study, a higher impact was observed with PS-COOH than with PS-NH ₂ ,
342	contrary to previous works which revealed a higher impact of positively charged nano-
343	polystyrene particles on different marine organisms: Crustacea (Bergami et al., 2016;
344	2017; Nasser and Lynch, 2016) Bivalvia (Balbi et al., 2017) Equinodea (Della Torre et
345	al., 2014) and Chlorophyceae (Bergami et al., 2017). Positively-charged NPs have been
346	suggested to interact more strongly with biological membranes, as demonstrated by in
347	vitro studies with mammalian cells (Anguissola et al., 2014; Varela et al., 2012). In the

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present study, both PS-COOH and PS-NH2 showed negative charges when they were suspended in seawater. Changes on nanoplastics charge is not surprising as, once in seawater, NPs can interact rapidly with the high number of charged ions (H⁺ and OH⁻) present (Cole and Galloway, 2015). Additionally, in natural environments, NPs can interact with natural colloids, sediment and soils, formatting an eco-corona coating that potentially affects the fate of NPs in the water column, bioavailability, toxicity and uptake (Canesi et al., 2017; Galloway et al., 2017). Afshinnia et al. (2018) reported that natural colloids from the media (e.g., humic substances) inverted the charge in silver nanoparticles (AgNPs), suppressing the positive charge and enhancing the negative one, depending on point zero charge of the particles and the solution pH. Nevertheless, positive charges of 50 nm polystyrene amino coated (PS-NH₂) in seawater has been reported in other studies (Bergami et al., 2017; Della Torre et al., 2014; Manfra et al., 2017). This obliges us to ask whether the size or the commercial supplier has an impact on particles characteristics in seawater. More information about physico-chemical characteristics of nanoplastics is needed for better interpretation of NPs behave in different solutions. In addition, more powerful techniques should be used considering the limitations of DLS technique which, according to ZETASIZER manual, cannot provide accurate Z-potential between -10 to +10 mV. Overall, in our study, NPs were from the same commercial brand and size, differing only in coated functional group and charge. As consequence, the differences in toxicity observed between PS-COOH and PS-NH₂ are probably not related to the size, aggregation state or even charge of the particles, which were similar between the two particle types. Instead, the nature of the coating (carboxylic or amine) may have interacted differently with the cell membranes, either directly or through interaction with specific biomolecules present in each seminal medium.

3/3	Nanoplastics attachment to spermatozoa and micro-scale aggregate formation
374	around oocytes
375	Spermatozoa consist of a head (between 2-3 µm length), which contains the acrosome,
376	an intermediate part and the flagellum ($\sim 33~\mu m$ in mature spermatozoa) (Demoy-
377	Schneider et al., 2013). The increase in relative size and complexity observed upon
378	exposure to NPs is likely explained by an adhesion of NPs mostly to the spermatozoa
379	head, as observed for PS-COOH using confocal microscopy. Similarly, an increase in
380	relative size and relative complexity of algal cells was previously reported under
381	exposure to metallic nanoparticles, but the localization of these particles was not
382	confirmed by microscopy (Sendra et al., 2017).
383	Regarding the oocytes, NPs formed hetero-aggregates with the organic matter from
384	seminal media that were entrapped to the cells. These aggregates were not present in all
385	observations, which is probably due to the high variability observed in cell responses to
386	NP exposure. Also as microscopy observations were performed on fixed samples,
387	addition of fixative could have led to modifications regarding NPs adhesion.
388	Nevertheless, slight modifications are expected since fixative was added after exposure
389	of oocytes to NPs.
390	Nanoplastics exposure affects spermatozoa more than oocytes
391	Differential effects of NPs observed between spermatozoa and oocytes could be
392	explained by differences in their membrane characteristics (Kline, 1991). In our study,
393	we observed NPs attached to both, spermatozoa and oocytes. NPs exposure favour
394	spermatozoa aggregation on the PS-COOH conditions. The increase of spermatozoa
395	aggregates was concomitant to the decrease of single spermatozoa cell number observed
396	upon exposure to PS-COOH. Conversely, no aggregation was observed in oocytes.

397	Oocytes appeared to be covered by a transparent structure, probably from the molecules
398	and colloids present in the medium that were absent from the spermatozoa samples. In
399	general, the presence of proteins reduces the free energy of the NP surface, thus
400	reducing unspecific adhesion to the membrane (Lesniak et al., 2013). Gao et al. (2017)
401	recently reported sex-specific differences in the protein composition of the nanoparticle
402	corona in fish and suggested that some specific proteins related to vitellogenin coated
403	nanoparticles in developing oocytes. Such a mechanism may explain why, in our study,
404	we observed a differential interaction of NPs between spermatozoa and oocytes. Other
405	factors such as (i) the high inter-individual variability in oocyte quality, as previously
406	observed for this species (Boudry et al., 2002), potentially masking the effects of NPs,
407	(ii) the way the oocytes were collected, i.e., because the gonads were stripped, the
408	samples would have included potentially immature oocytes (Le Goïc et al., 2014), or
409	(iii) natural characteristics of each cell type such as the size, 40 times higher oocytes
410	than spermatozoa and functional structures (spermatozoa with a motile flagella vs
411	oocytes not motile and with higher lipid content) could explain the difference in NPs
412	effects between spermatozoa and oocytes.
413	During the viability test in oocytes, NPs exposure (100 mg L ⁻¹) promote a significant
414	decrease of SYBRgreen fluorescence after exposure to both PS-COOH and PS-NH ₂ .
415	This effect may be likewise caused by several factors (i) interaction of NPs with the
416	fluorescent dye (SYBRgreen), (ii) a reduction in DNA content related to chromosome
417	anomalies or chromosome deletions (Haberkorn et al., 2010), or even (iii) changes in
418	the membrane permeability which may also impair the entry or retention of the
419	SYBRgreen into the cells. Cellular membrane hyperpolarization has been demonstrated
420	in other organisms after exposure to pollutants (Seoane et al., 2017). Unfortunately,
421	Pacific oyster oocyte ultrastructure remains poorly described and its envelope thickness

is not precisely known (Suquet, et al., 2007), which hinders a thorough understanding of the biological response observed in exposed gametes in this work. Further research is needed to explore the interaction between NPs and gamete membranes in greater depth, which will require challenging technological developments in microscope visualization.

Increase in ROS production by spermatozoa upon exposure to PS-COOH

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The dose-response generation of intracellular ROS observed in spermatozoa exposed to PS-COOH calls into question their subsequent ability to fertilize oocytes. Increase in ROS production and oxidative damage upon exposure to micro- and nanoplastics has been observed in other species, such as the lugworm (Arenicola marina; Browne et al., 2013), a planktonic crustacean (Daphia magna; Ma et al., 2016; Nasser and Lynch, 2016), algae (Chlorella and Scenedesmus; Bhattacharya et al., 2010), mussels (Mytilus spp.; Paul-Pont et al., 2016) and zebrafish larvae (Danio rerio; Chen et al., 2017). ROS generation and the subsequent oxidative stress have been described as the predominant mechanism leading to nano-toxicity, including DNA damage, unregulated cell signalling, changes in cell motility, cytotoxicity, apoptosis and cancer initiation and promotion (Fu et al., 2014). However, in the present study, the increase in ROS did not lead to any effects on cell motility or viability, suggesting that the oxidative stress produced was not sufficient to cause irreversible damage. From another perspective, ROS play important roles in spermatozoa physiological functions in different species including capacitation, hyperactivation and the acrosome reaction (Kothari et al., 2010; Zilli and Schiavone, 2016 in Aitken, 2017). In the present study, the positive correlation observed between the percentage of motile spermatozoa and their ROS production after being exposed to PS-COOH, particularly after 5 h exposure, may suggest the stimulation of spermatozoa capacitation as previously reported in mammals after an increase in intracellular ROS content (Aitken, 2017). Nevertheless, when ROS

production overcomes the oxidative balance of spermatozoa, a decrease in spermatozoa motility may occur as it has been reported in other studies carried out with mouse sperm exposed to metallic nanoparticles (Hong et al., 2015; Kumar et al., 2001). Further studies using biochemical markers of oxidative stress (MDA formation, lipid peroxidation, etc.) would be useful to clarify the impacts of ROS production induced by NP exposure observed here.

5. Conclusion

The present study evaluated the effects of nanoplastics on oyster gametes that are in close contact with water-borne pollution and thus constitute very sensitive life stages. Our results demonstrated adhesion of NPs to oyster spermatozoa leading to an increase in relative cell size and complexity. A significant dose-response increase in ROS production by spermatozoa was demonstrated upon exposure to PS-COOH, but not with PS-NH₂. Conversely, oocytes were less impacted after exposure to both NPs.

Differences in effects between the NPs and the two gamete cells were not related to particle size, charge or aggregation state, as tested NPs displayed similar features over time both in seawater and seminal media. Instead, results suggest that interactions with seminal biomolecules and/or cell membranes differed according to particle coating and gametes' type. It would be of further interest to investigate the nature of the eco-corona formed on nanoplastics and how this modifies particle toxicity toward oyster gametes. Subsequent impacts of gamete exposure to NPs on oyster reproductive success could be assessed by studying the fertilization rate as proxy of gamete quality (Boulais et al. 2015), as well as embryo-larval development.

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Figure 1. Nanoplastics characterization using Dynamic Light Scattering (A, B) and Transmission Electronic Microscopy (TEM) imaging (C, D). Size distribution of 100 nm PS-COOH (A) and 100 nm PS-NH₂ (B) measured in triplicates (each replicate is marked in blue, red and green) using a Zetasizer Nano Series software 6.2. TEM observations of 100 nm PS-COOH (C) and 100 nm PS-NH₂ (D) are also presented. Scale bar: 50 nm.

Figure 2. Confocal fluorescence microscopy carried out in fixed samples of control spermatozoa and oocytes (A, B) and spermatozoa/ oocytes exposed to 100 mg L⁻¹ PS-COOH for 3h exposure (C, D), stained with DAPI. Control cells (A, B) showed blue fluorescence coming from the DAPI dye and they has lifelike fixative' green fluorescence due to the fixation with formaldehyde. Exposed spermatozoa (C) evidenced high green fluorescence in the acrosome and in some points along the tale, coming from the aggregates of nanoplastics attached to the cell. Exposed oocytes (D) head fluorescence green coming from nanoplastics aggregates wrapped into the organic debris.

Figure 3. Intracellular reactive oxygen production (ROS) of individual' (n=6) spermatozoa after 1 h, 3 h and 5 h exposure to 4 different concentrations of PS-COOH. Significant differences between treatments are marked with lowercase letters.

SUPPLEMENTARY MATERIAL

Supplementary Figure 1 Histograms for SYBR green staining analysis of one female exposed to PS-COOH (1A) and PS-NH₂ (1B) for 1 h. Levels of green fluorescence is represented in X-axis in arbitrary units (A.U) and cell count (Y-axis). Each concentration is given as a type line. The fluorescence of the highest NPs concentration (100 mg L⁻¹), marked with a red area, was significantly different of the rest of treatments for both, PS-COOH and PS-NH₂ (P< 0.001 and P < 0.001; ANOVA) respectively. Control treatment are marked with a green area. Statistical analyses were performed on all females (n=6), only one females was presented as example.

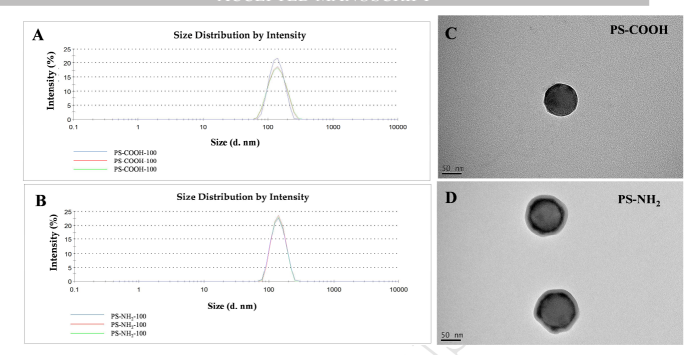
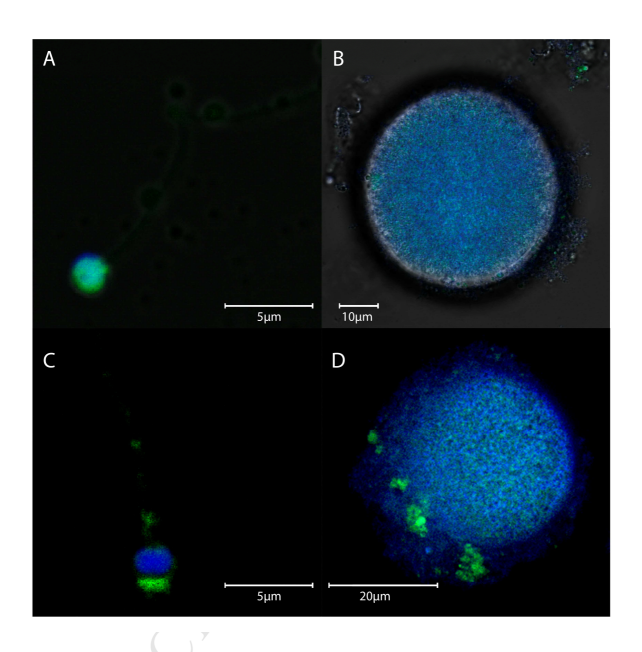


Figure 1



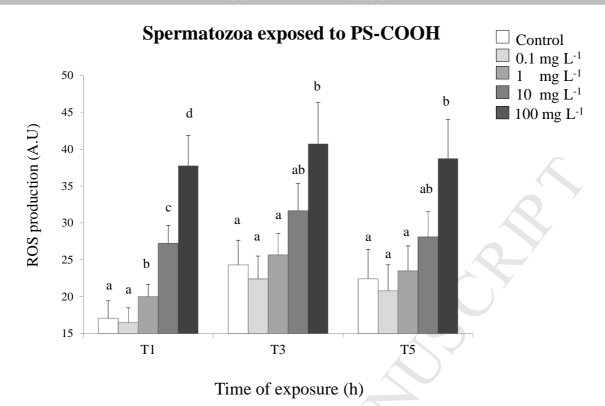


Figure 3.