Fate of Cerium Oxide Nanoparticles in Natural Waters and Immunotoxicity in Exposed Rainbow Trout

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Abstract

Once released in the environment, engineered nanoparticles (NPs) can undergo important transformation resulting in changed properties under natural conditions. This study investigated the fate, the bioavailability and the immunotoxicity of cerium oxide (CeO$_2$) nanoparticles in fish exposed to CeO$_2$ in representative surface waters differing in pH, organic matter content and conductivity (green and brown waters). Following an incubation period of NP CeO$_2$ in different surface waters, particle size distribution and shape were determined by ultrafiltration and ICP-mass spectrometry, electronic microscopy and dynamic light scattering (DSL). Bioaccumulation and effect biomarkers focusing on the immune system responses (viability of immune cells and phagocytic activity) were also determined. Particle size distributions significantly changed under all types of surface waters where aggregation of NPs was commonly observed. Indeed, >80% of NPs CeO$_2$ were found as aggregates (>450 nm) and large colloids (>100 nm). Less than 1% cerium (Ce) was found in the truly dissolved fraction (<1 kDa) suggesting no evidence of degradation for NP CeO$_2$ in the water samples after 96 h. The NPs CeO$_2$ were preferably accumulated in fish gills and accumulation was the highest in green waters which contained less total organic carbon (TOC), higher conductivity (218 µS/cm) and higher pH (7.8-8.0) than brown waters. The toxic properties (induced phagocytosis) of NP CeO$_2$ also differed when dispersed in brown, green and tap waters. NP CeO$_2$ induced fish mortality at initial concentration of 10 µg/L Ce in both tap and green waters but not in brown waters which have different and high organic matter sources, lower pH and conductivity values. In conclusion, NPs CeO$_2$ lends aggregate in representative freshwater, adsorb on gills and the immunotoxic potential is reduced in the presence of high natural organic matter, mildly acidic pH and low conductivity as found in brown waters.

Keywords: Nanoparticle; Nanotoxicity; Bioaccumulation; Transformation

Introduction

Nanoparticles have unique commercial properties [1] and their growing use will lead to their introduction into the aquatic environment which can threaten ecosystems [2]. Cerium oxide NPs are widely used in biomedical sector as antioxidants in biological systems [3], and engineering industries, as additives in diesel fuel [4] and ceramic applications [5]. Moreover, Ce is considered the most abundant rare earth elements and intensely used by our economy. For these reasons, this metal oxide is on the list of prioritized nanomaterials by the Organization for Economic Cooperation and Development (OECD) for environmental characterization and assessment. Cerium pollution is mainly associated with landfills from leachable of solid wastes of electronic devices and sludge and wastewater discharges from industries such as ceramic plants [6]. Despite the increasing interest in the application of NP CeO$_2$ in the industry, the occurrence in the aquatic environment and potential toxicological effects to organisms still remain to be investigated as a requirement for environmental risk assessment [7]. The positive charge (Zeta potential +30 mV) at the surface of NP CeO$_2$ influences their stabilization and ability of absorption to solids such as sludge. According to a survey/study in treatment plants, up to 6% of CeO$_2$ would escape and release in the plant outflow, reaching effluents and subsequently receiving natural waters [8]. Concentrations of CeO$_2$ in treated wastewater effluents could reach concentrations up to 1 µg/L [6]. Ecological assessments of NP need to address their changing physicochemical properties in pH, organic matter content and conductivity (green and brown waters). Following an incubation period of NP CeO$_2$ in different surface waters, particle size distribution and shape were determined by ultrafiltration and ICP-mass spectrometry, electronic microscopy and dynamic light scattering (DSL). Bioaccumulation and effect biomarkers focusing on the immune system responses (viability of immune cells and phagocytic activity) were also determined. Particle size distributions significantly changed under all types of surface waters where aggregation of NPs was commonly observed. Indeed, >80% of NPs CeO$_2$ were found as aggregates (>450 nm) and large colloids (>100 nm). Less than 1% cerium (Ce) was found in the truly dissolved fraction (<1 kDa) suggesting no evidence of degradation for NP CeO$_2$ in the water samples after 96 h. The NPs CeO$_2$ were preferably accumulated in fish gills and accumulation was the highest in green waters which contained less total organic carbon (TOC), higher conductivity (218 µS/cm) and higher pH (7.8-8.0) than brown waters. The toxic properties (induced phagocytosis) of NP CeO$_2$ also differed when dispersed in brown, green and tap waters. NP CeO$_2$ induced fish mortality at initial concentration of 10 µg/L Ce in both tap and green waters but not in brown waters which have different and high organic matter sources, lower pH and conductivity values. In conclusion, NPs CeO$_2$ lends aggregate in representative freshwater, adsorb on gills and the immunotoxic potential is reduced in the presence of high natural organic matter, mildly acidic pH and low conductivity as found in brown waters.

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Because of the relatively low solubility of NP CeO$_2$, this metal oxide NP was reported to be 10 times less accumulated in exposed marine mussels when compared to results for the more soluble NP ZnO [11]. Cerium oxide NPs in water-exposed fish are absorbed through their gastrointestinal tract and gills [19]. However, the tendency of NP CeO$_2$ to aggregate may lead to different exposure routes which can lead to different toxicity in aquatic organisms [20]. Hence, the occurrence of NP CeO$_2$ aggregates could likely influence the immunity which is involved in the recognition and elimination of foreign particles.

In contrast to human and terrestrial toxicological investigations, NP CeO$_2$ aquatic toxicity studies are scarce [5,21-23]. While no toxicity was observed in zebrafish embryos exposed in culture media, exposed adults in the water column led to significant bioaccumulation of Ce in fish liver [19,24]. Exposure of bivalves to NP CeO$_2$ at predicted concentrations (1-100 µg/L) resulted in significant adverse effects impacting the lysosomal system, the catalase activity and the digestive gland functions [25]. Protection against foreign materials such as xenobiotics occurs through two different components of the immune system: the first producing an immediate and nonspecific response (i.e. innate immunity) and the second producing a specific response as well as an immunological memory (i.e. acquired immunity). The immunological defense of most aquatic species relies mainly on the nonspecific immune response as the first line of defense and includes phagocytosis and inflammatory reactions [26]. In mussels that rely only on innate immunity, exposure to CdTe quantum dots aggregates leads to significant effects at the hemocyte viability, phagocytosis and cell lysis potential [27]. Exposure to silver NPs to fish also induced immunosuppression at the innate immunity level as well which suggests that the exposure to NPs and aggregates targets that system in aquatic organisms [28].

Several studies investigated potential impacts on fish as function of NP concentrations of unmodified (i.e., non-transformed) nanoscale metal oxides and pointed out the need to consider the exposure to transformed products as well transformation for a complete risk assessment [12,19,29,30]. The objectives of the study were to evaluate the transformation and behaviour of NP CeO$_2$ in natural waters of different properties (pH, conductivity and natural organic carbon) as key information for environmental impact assessment. The study reports on the bioaccumulation and immunological effects in fish exposed to primary NP CeO$_2$ and their transformation products by taking into account the fate and behavior (including aggregates and dissociated forms) of the nanoparticles in various natural aqueous matrices.

Materials and Methods

Cerium oxide nanoparticle

A stock solution of NP CeO$_2$ from Sigma-Aldrich® chemical company (Ontario, Canada) was used in this study. According to the manufacturer’s specifications, the NP CeO$_2$ suspension has a size ≤25 nm, at a concentration of 10% wt in water. For the exposure regime, rainbow trouts (Oncorhynchus mykiss) were exposed to a nominal concentration of 10 µg/L total Ce as sonicated NP CeO$_2$ in dechlorinated tap water (controls), and two types of surface waters as described below. The NP CeO$_2$ concentration was chosen according to environmental concentrations [31] and to previous standard acute tests with daphnia (Daphnia magna) and NP CeO$_2$ [32].

Types of water

Three types of water, with initial contrasting water chemistry, were sampled in November 2013; green water, organic-rich brown water and UV/charcoal-treated, filtered tap water. Total organic carbon (TOC) concentrations, as well as pH and conductivity were measured at the beginning and at the end of the exposure (Table 1). Rainbow trouts were then exposed to NP CeO$_2$ in tap water, green and brown freshwater. Tap water consisted of dechlorinated and UV/charcoal-treated green water from the Saint-Lawrence River near the city of Montréal. Green waters were surface waters with low organic carbon content (3.2 mg/L), relatively high conductivity (218 µS/cm) and pH (7.6). Brown waters sampled in the Ottawa River were different with lower pH (6.7) and conductivity (112 µS/cm), but higher TOC (6.4 mg/L).

Fish exposure

Juvenile rainbow trouts (Oncorhynchus mykiss) (mean size 122.5 ± 9.2 mm; mean weight 26.3 ± 5.5 g) were provided by a local hatchery (Pisciculture des Arpents-Vert, Ste-Edwidge, Qc), maintained in 1000-L tanks at 15°C, fed daily with a commercial trout chow for 2 weeks and held under a natural photoperiod (12 h light: 12 h dark) before the initiation of exposure. The following week of water sampling, five trouts were placed in 10 L containers lined with polyethylene bags and exposed to 10 µg/L total Ce as NP CeO$_2$ in each type of unfiltered water for 96 h at 15°C. The exposure experiment was repeated twice. The water was not renewed during the experiment. Fish were not fed and monitored daily for any signs of distress or changes in behaviour. Dissolved oxygen was maintained above 95%, pH between 7.2-7.9, and temperature at 15°C during the exposure period. After the exposure period, fish were ethically euthanized with 0.1% of MS-222 (Sigma-Aldrich®, ON, Canada) using approved protocol by the Canadian Council on Animal Care Committee. The pronephros was dissected out and kept on ice for immunocompetence assessments on the same day. Liver and gills were immediately collected, weighed and stored at -80°C for subsequent chemical and biochemical analyses.

Characterization of NP CeO$_2$

Transmission electron microscope (TEM) observation and electron-dispersive X-ray analysis: A sub-sample (50 mL) of each type of water was collected in duplicate after 96 h and kept at 4°C for further NP CeO$_2$ characterization. The samples as well as the stock solution were observed by TEM no longer than 3 days after the end of exposure. A drop of exposure medium was placed on a copper grid capped with a lacey carbon film for TEM analysis. Once the sample was dehydrated, it was examined by TEM (JEOL, 2100-F model) operated at 200 kV for image capture in clear bottom. For each TEM picture, an electron-dispersive X-ray analysis (EDS) was performed for element composition of targeted particles. Maximal length (L) and width (W) were measured on the TEM pictures for the three types of water and the suspension stock in order to calculate an eccentricity ratio (e=L/W) using the software ImageJ 1.51 K (Wayne Rasband, National Institutes of Health, USA). For the three types of water, all TEM pictures were considered in order to obtain an important event number (brown

<table>
<thead>
<tr>
<th>Exposure time</th>
<th>Parameters</th>
<th>Tap water</th>
<th>Green water</th>
<th>Brown water</th>
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</thead>
<tbody>
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<tr>
<td></td>
<td>Conductivity (µS/cm)</td>
<td>347</td>
<td>377</td>
<td>161</td>
</tr>
</tbody>
</table>

TOC: Total Organic Carbon.

Table 1: Characterization of waters before and after incubation.
water n=20, green water n=11 and tap water n=20). The picture was separated in 4 equal squares and one of them was randomly chosen, all NPs CeO₂ were measured in this square (n=45).

Dynamic light scattering: Nanoparticle hydrodynamic size and Zeta potential were measured in duplicate using dynamic light scattering (DLS) (BrookHaven Instrument®, ZetaPlus/Bi-PALS) in the three types of water. Each water sample was previously filtered on a 450 nm membrane prior measurements to remove large aggregates. The NP CeO₂ stock solution was also diluted 1:10 with milli-Q water before the measurement on the DLS to improve the reading.

Size fraction distribution by filtration and ultrafiltration: Before the exposure, the NP CeO₂ suspensions were sonicated and characterized for their size fraction distribution (i.e., from truly dissolved to aggregate class) by a procedure previously described in Bruneau et al. [33]. Briefly, NP CeO₂ solutions were fractionated by microfiltration and ultrafiltration using a parallel decreasing membrane porosity size gradient. A subsample of 250 mL of NP CeO₂ suspension in each type of water was first filtered on a membrane of 450 nm porosity (FHLC04700, EMD-Millipore®) and 40 mL was then sampled for total Ce determination. The 450 nm filtrate was passed through membranes of three different pore sizes in parallel: 100 nm (VCTP04700, EMD-Millipore®) and 50 nm (VMWP04700, Millipore) and 25 nm (VMWP04700, EMD-Millipore®).

An ultrafiltration cell with constant agitation was used (Amicon® 400 system, EMD-Millipore®) for the ultrafiltration with 1 kDa cut-off (about 1.5 nm) (YM1 76 mm diameter, EMD-Millipore®) to determine the potential release of low molecular weight of Ce ion and its complexes. The pressure in the system was maintained constant at 60 psi and the sample at room temperature. The flow rate was near to 1 mL/min. This ultrafiltration step was considered to provide the “truly” dissolved ion Ce fraction. Total Ce concentrations were evaluated with an inducted-coupled plasma mass spectrometry (XSERIES 2 ICP-MS, Thermo Fisher Scientific, USA). Calibration of procedures and accuracy of the measurement were assessed with five replicates of SLRS-5 reference material (River water reference material for trace metals, National Research Council, Canada). Lanthanide element analyses in SLRS-5 were comparable to those reported in the study of Rousseau et al. [34]. In our study, the calculated limit of detection (LOD) was 0.3 ng/L and the limit of quantification (LOQ) was 0.9 ng/L. Concentrations in exposure media were expressed as total Ce in μg/L.

Cerium bioaccumulation in fish tissues

To determine Ce loading in fish tissues, livers and gills were individually sampled, weighed and frozen at -80°C until analysis. Tissues were digested with high purity 8 mL of concentrated HNO₃ (Seastar™ Chemical, BC, Canada), 1 mL of concentrated HCl (Seastar™ Chemical, BC, Canada), and 2 mL of concentrated H₂O₂ (Seastar™ Chemical, BC, Canada) added in that order. The tissues were then digested during 2 h with increasing temperature gradient (maximum 180°C) using a microwave digestion system (Ethos EZ, Milestone Scientific®, ON, Canada). Each digested tissue sample was then placed in a 15 mL-polyethylene tube and the final volume was adjusted to 12 mL with deionized water. Total Ce concentration was determined by ICP-MS (XSERIES 2 ICP-MS, Thermo Fisher Scientific, USA). For each digestion series, triplicates of dogfish liver certified reference material for trace metals (DOLT-5, National Research Council, Canada) were used to insure the reproducibility of the extraction method. The calculated LOD is 0.012 μg/g and the LOQ is 0.03 μg/g. Concentrations in tissues were expressed as total Ce in μg/mg wet tissues.

Immune parameters

The effects of Np CeO₂ were determined at the innate immunity level in exposed fish [28,35,36]. Immunocompetence was determined in duplicate in freshly prepared leucocytes using flow cytometry [26,37]. Briefly, leucocytes were extracted from the pronephros and isolated by centrifugation on a 51% Percoll™ gradient at 400 g, 30 min and 20°C (Sigma-Aldrich®, ON, Canada). The leucocyte fraction, which partitioned at the Percoll-media interface, was collected and resuspended in phosphate buffered saline (PBS: 140 mM NaCl, 1 mM KH₂PO₄, and 1 mM NaHCO₃, pH 7.4) and centrifuged at 400 g for 10 min at 20°C. The cells were then resuspended in 1 mL of RPMI cell culture media for cell counting and viability determination using trypan blue (dead cells remain blue). Observations and cell counting were done on a hemocytometer at 200x enlargement.

Viability of immune cells was performed according to an adapted method from Brouseau et al. [26,37]. The leucocyte cell fraction was diluted at a concentration of 2 million viable cells/ml in RPMI (Roswell Park Memorial Institute) media containing 10% fetal bovine serum (FBS), and 100 U/mL penicillin, 100 µg/mL streptomycin and 10 mM HEPES (4-(2-Hydroxyethyl) piperazine-1-ethanesulfonic acid), pH 7.4, and incubated in the dark for an additional 18 h at 15°C in duplicate. After this incubation, cells were washed and resuspended in RPMI as described above. Viability was observed by flow cytometry using propidium iodide (PI) (Sigma-Aldrich®, ON, Canada). A 4 µL aliquot of PI (100 µg/mL) was added to 200 µL of each cell suspension for 5 min on ice before measurement. Propidium iodide fluorescence was analyzed with a flow cytometer equipped with an argon laser excitation (λ=488 nm ± 10 nm) (Guava® EasyCyte, EMD-Millipore®, USA). Fluorescence for each sample was measured in duplicate at 625 nm with 42 nm bandwidth, and 5000 events were registered. The proportion of lymphocytes and monocytes, which are the main sub-populations of leucocytes with granulocytes, were determined on the basis on the forward (cell volume) and side (cell internal complexity) scatter dot plots using the software instrument. Lymphocytes are generally smaller cells and have a more homogeneous cytoplasm than monocytes.

Phagocytosis activity was measured following the protocol of Brouseau et al. [26,37]. Briefly, 1 ml of adjusted cell concentration (2 million/ml) was added to 24 well culture-coated well plates in duplicate. Cells were incubated with a ratio 100:1 of fluorescent latex beads (Polysciences®, PA, USA) in order to observe the phagocytosis capacity of the cells. After an incubation period of 18 h at 15°C, the cell suspensions were overlaid on 3% serum bovine albumin in PBS and centrifuged at 400 g for 5 min to remove loosely adhered beads on the cell surface. Cell pellets were then resuspended and fixed in 0.5% formaldehyde and 0.2% sodium azide in PBS. Cells containing Latex bead-fluorescence were measured using an argon laser flow cytometer at 530 nm emission as described above, and at least 10 000 events were registered. The immunoactivity was defined as the number of cells containing at least one bead and the immunoefficiency as the number of cells containing three beads or more.

Data analysis

Differences between the biomarkers were examined using a one way ANOVA when data normality was confirmed using Kruskal Wallis normality test. There were 6 treatments in the present study design: 1-3) fish exposed to control water (3 types) only; 4) fish exposed to tap water and 10 μg/L Ce as CeO₂, NP, 5) fish exposed to green water and 10 μg/L Ce as CeO₂, NP and 6) fish exposed to brown water and...
10 µg/L Ce as CeO₂ NP. A post-hoc Tukey test was used to determine the differences between the groups. When the data were not normal, a Kruskal Wallis non-parametric ANOVA was used instead and critical difference between treatments were appraised using the Mann-Whitney rank test. Significance was set at p<0.05.

Results and Discussion

Characterization of NP CeO₂

In stock suspension, NPs CeO₂ were retrieved in small aggregate forms (Figure 1). Dynamic light scattering (DLS) analyses revealed that NPs CeO₂ left after filtration were smaller (mean 16.9 nm) in tap water, than NP CeO₂ in brown (32.0 nm) and in green water (26.1 nm) (Table 2). The eccentricity ratios were 1.50 (±0.35), 1.50 (±0.50), 1.78 (±0.85), for brown, green and tap water respectively, revealing no significant difference in shape among the three types of matrix and confirming the stability of NP CeO₂ in natural waters after short-term exposure. In the stock suspension the measured eccentricity ratio was 1.53 (±0.44) and was close to the eccentricity ratios measured in brown and green waters. As observed on TEM images, NPs CeO₂ were retrieved as small aggregates and maintained their geometric form (Figure 2). No significant modification of the NP shape was observed with each water types. This is in contrast with other reported observations with NP silver [28]. These results suggest that NPs CeO₂ were not likely degraded during the exposure period and were bioavailable as NP forms.

Size distribution of NP CeO₂

At the beginning of the exposure, the nominal Ce concentrations

![Figure 1](image1.png)
![Figure 2](image2.png)

Table 2: Mean diameter size and Zeta potential of NP CeO₂ in water after exposure. These results were observed with a DLS after filtration through a membrane of 0.45 µm in order to remove large aggregates and measure the mean diameter size of the remaining particles.
were relatively constant at 7.5, 11 and 8 µg/L for tap, green and brown waters respectively. After 96 h, Ce concentrations in water were 3.4, 2.9 and 3.6 µg/L in tap, green and brown waters respectively, confirming the equilibrium of the exposure concentrations and observed settlement. After the exposure period, TOC concentrations were increased for each type of water indicating a release of carbon from the fish during the exposure. During the exposure, pH increased in tap and green waters and decreased in brown water (Table 1). Initial NP CeO₂ suspension became homogenous during the exposure as small NPs still available in the water column and aggregates settled to the bottom of the tank. This hypothesis was in agreement with the TEM picture and the DLS sizes (Figure 2).

After a first filtration step on 450 nm, only 10%, 3% and 7% of Ce from NP CeO₂ were measured in tap, green and brown water respectively; indicating that more than 90% of Ce was preferentially linked to large particles ≥ 450 nm (Figure 3). After a filtration step on 50 nm, less than 1% of the Ce was measured in waters, confirming that NPs CeO₂ were preferentially retrieved in aggregates. Large aggregates were observed for natural waters compared to the reference tap water.

Smaller aggregates (<100 nm) were found in the reference tap water. The aggregation of NP CeO₂ in natural green and brown waters was observed, but to a lesser extent for the latter (Figures 2 and 3). Brown waters from the Ottawa River reduced aggregation, where more Ce was found in the filterable (<450 nm) fraction compared to the green water. No dissolved Ce, however, was detected in the 1 kDa (≈1.5 nm) fraction suggesting low degradation of the NP CeO₂ during the exposure with the three types of water.

Bioaccumulation of NP CeO₂ in fish tissues

Significant increases in Ce concentrations compared to the control were observed in trout gills exposed to each type of water (Figure 4A). This result could indicate that NP CeO₂ has a direct interaction with fish gills and was adsorbed via water contact through gills. Most Ce was accumulated or adsorbed in gills (one order of magnitude higher) compared to the liver. Cerium adsorption in gills was more important in green water but not significantly different (Kruskal-Wallis test) from tap and brown waters. In addition, some trout mortality (unpublished data) was observed in green water suggesting that the green water would promote adsorption of NP CeO₂ and induce harmful effect on fish compared to the two other types of water. A significant bioaccumulation of Ce was only observed in the trout liver exposed to NPs CeO₂ in brown water compared with fish exposed to NPs in other waters, but at one order of magnitude less than the levels found in gills (Figure 4B). Uptake of NP CeO₂ was also observed in the liver of zebrafish via the water column [19]. Our findings confirm that NP CeO₂ were bioavailable to fish external tissues (gills) as well as low distribution in internal organs such as the liver. For such nano-sized material and larger particles, the gastrointestinal tract was suggested as the main uptake pathway for fish [38]. It would have been of interest to analyze Ce in the fish gut to confirm whether fish ingested Ce aggregates. In mussels exposed to Ce and Zn oxide nanoparticles, substantial amounts of these elements in the pseudofeces were detected [11]. Mussels exposed to 10 mg/L Ce or Zn oxide nanoparticles rejected 21 mg/g and 63 mg/g of Ce and Zn in the pseudofeces while they were accumulated in tissues at 62 and 880 µg/g on a dry weight basis suggesting that Ce was bioavailable to mussels albeit one order of magnitude less than Zn.

Biomarkers

Viability: Significant decreases in both lymphocyte and monocyte viability were observed between control and NP CeO₂ treatments for
the three types of water, indicating a cytotoxic effect of NP during short exposure (Figure 5A). This suggests that the main driver of cytotoxicity was NP CeO₂, concentration and not the surface water types. To best of our knowledge, this is the first study on the immunotoxicity of NP CeO₂ on fish leukocyte populations. Cerium oxide NPs were found to induce apoptosis and autophagy in human peripheral blood monocytes [39]. This response was induced at high concentrations (1-10 mg/L; 40 h) due to aggregation by the high salt contents in the culture medium. In another study with human hepatoma cells, decreased viability was produced at 20 µg/L NP CeO₂ (20 nm diameter) after 70 h incubation time [22]. A different exposure to large bulk of CeO₂ particles (5 µm) was equally toxic to these cells suggesting that the primary interaction of NP CeO₂ is similar to large particles and surface interactions at the outer cytoplasmic membrane are at play.

Phagocytosis: Significant stimulations were observed between control and exposed treatments for tap and green waters. However, non-significant immunoinactivity and immunoefficiency (as per definition in the methods section) was observed with NP CeO₂ in brown water (Figure 5B). Similarly, ecotoxicological studies using different biomarkers classified NP CeO₂ in natural waters as pollutants with low in vivo toxicity [2,23,38]. Data on the immunotoxicity in fish of CeO₂ as powder or nanoparticles are rather scarce. The exposure concentration used in the present study (10 µg/L) was kept at reported levels of Ce in wastewater and more than one order of magnitude below the estimated probable no-effect concentration of NP CeO₂ between 3-5000 mg/L depending on the species [40]. Algae were the most sensitive species towards NP CeO₂ compared to daphnia (Daphnia magna), beaver-tail fairy shrimp (Thamnocephalus platyurus) and zebra fish (Danio rerio) embryos. According to the study of van Hoecke [40], the acute toxicity of NP CeO₂ would be reached at concentration >5000 mg/L in aquatic ecosystems but mechanisms such as altered embryos hatching could be observed following 72 h exposure to 200 mg/L of NP CeO₂ of 20 mm and chronic toxicity for green algae at lower concentrations (2.6-5.4 mg/L).

In another study, NPs CeO₂ were not considered toxic compared with bulk CeO powder at 0-10 mg/L concentration range for daphnia (Daphnia magna) and 0.01-0.1 mg/L for common carp (Cyprinus carpio) [38]. Nevertheless, in the present study exposure to NP CeO₂ at 10 µg/L decreased cell viability in lymphocytes and monocytes and increased phagocytosis in macrophages for tap and green waters for the latter. The innate immune system involves the recognition of foreign particles (such as NP and their aggregates) and ingestion by macrophages through the process of phagocytosis [4]. Ingested NPs would be then degraded in phagolysosomes with the production of reactive oxygen species (ROS), and hydrolytic enzymes [17]. Since phagocytosis involves the generation of ROS, the presence of NP CeO₂ aggregates could stimulate the phenomenon at first by scavenging radicals during the oxidative burst. Moreover, this metal oxide NP was reported as an active redox catalyst with potential for oxidative stress [41,42]. Several studies have shown that exposure to NP CeO₂ can result in oxidative stress, inflammation and DNA damage to organisms [18,20]. During the phagocytosis, the degradation of NPs and release of elements from which the NPs are derived occur and could lead to cytotoxicity. In marine mussels, hemocytes exposed in vitro to NP CeO₂ decreased viability and phagocytosis were observed in hemocytes but at concentrations (7 mg/L) much higher than in the present study [43]. The oxidative effect of NP CeO₂ would be coupled to the reduction of Ce(IV) to Ce(III) at high NP concentrations in the mg/L range [20,24,44]. In NP CeO₂, Ce atoms could have valence states Ce (III/IV) allowing the storage and the release of oxygen by the nanoparticle [45]. Various studies demonstrated that the redox state plays a large role in determining the characteristics and behaviour of NP CeO₂ [9,40,46,47]. In agreement with this statement, we hypothesized that the source of the organic matter could play a role in the behaviour of NP CeO₂ in natural media affecting the defense mechanism of the aquatic organisms such as phagocytosis and the level of the ROS (reactive oxygen species) and inducing negative impacts on aquatic biota health. Further experiments are required to consolidate this statement.

Conclusion

This study raises new concerns about NP CeO₂ fate and toxicity to aquatic organisms. According to our observations, the commercial NP suspension was mostly found as NP aggregates in natural waters. We found that 90% of NP CeO₂ occurred as aggregates (>450 nm) regardless of the three types of water (tap, green, and brown). Cerium oxide NPs were preferentially adsorbed or bioaccumulated in gills when fish were exposed to NPs in natural waters. After 96 h, the only significant increase in Ce concentration in liver was observed in trouts exposed to NP in brown water. Cerium oxide NP induced phagocytosis in fish maintained in tap and green waters. More investigation is required on the fate of NPs CeO₂ of different sizes in natural media with variable properties (e.g., organic matter concentrations and quality) and their subsequent effects to aquatic organisms.

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