Uptake, Transport, and Toxicity of Pristine and Weathered Micro- and Nanoplastics in Human Placenta Cells

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BACKGROUND: The first evidence of micro- and nanoplastic (MNP) exposure in the human placenta is emerging. However, the toxicokinetics and toxicity of MNPs in the placenta, specifically environmentally relevant particles, remain unclear.

OBJECTIVES: We examined the transport, uptake, and toxicity of pristine and experimentally weathered MNPs in non-syncytialized and syncytialized BeWo b30 choriocarcinoma cells.

METHODS: We performed untargeted chemical characterization of pristine and weathered MNPs using liquid chromatography high-resolution mass spectrometry to evaluate compositional differences following particle weathering. We investigated cellular internalization of pristine and weathered polystyrene (PS; 0.05–10 μm) and high-density polyethylene (HDPE; 0–80 μm) particles using high-resolution confocal imaging and three-dimensional rendering. We investigated the influence of particle coating with human plasma on the cellular transport of PS particles using a transwell setup and examined the influence of acute MNP exposure on cell viability, damage to the plasma membrane, and expression of genes involved in steroidogenesis.

RESULTS: Chemical characterization of MNPs showed a significantly higher number of unique features in pristine particles in comparison with weathered particles. Size-dependent placental uptake of pristine and weathered MNPs was observed in both placental cell types after 24 h exposure. Cellular transport was limited and size-dependent and was not influenced by particle coating with human plasma. None of the MNPs affected cell viability. Damage to the plasma membrane was observed only for 0.05 μm PS particles in the non-syncytialized cells at the highest concentration tested (100 μg/mL). Modest down-regulation of hsd17b1 was observed in syncytialized cells exposed to pristine MNPs.

DISCUSSION: Our results suggest that pristine and weathered MNPs are internalized and translocated in placental cells in vitro. Effects on gene expression observed upon pristine PS and HDPE particle exposure warrant further examination. More in-depth investigations are needed to better understand the potential health risks of MNP and chemicals associated with them under environmentally relevant exposure scenarios.

Introduction

Micro- and nanoplastics (MNPs) are global environmental pollutants with widespread exposure in the human environment e.g., food,1,2 water,3 air,4,5 and dust.6 Nevertheless, MNP toxicokinetics and toxicity in humans are still largely unexplored.7 Recent animal studies have shown that ingested or inhaled MNPs can translocate across pulmonary and intestinal cell barriers to secondary organs, including placental and fetal tissues.8−11 Moreover, epidemiological research indicates that fine particles (<2.5 μm) from air pollution can reach the placenta in situ and increase the risk of pregnancy complications such as gestational diabetes and preeclampsia.12,13 Preterm birth, or low birth weight.14 These findings have raised concerns for potential adverse health effects of placental MNP exposure during pregnancy, a period of heightened vulnerability when even subtle environmental insults can have long-term effects on fetal development and health later in life.15,16

To date, a limited number of human in vitro placenta cell models17−19 and ex vivo placenta perfusion studies20−22 have demonstrated the transplacental transport of polystyrene MNPs. Recently, a study by Ragusa et al.23 was the first to detect MNPs in human placenta samples. Although this research provided the first indication of placental exposure, understanding of placental toxicokinetics and toxicity of MNPs is limited. Gaining insight into these mechanisms is indispensable for a better understanding of possible adverse effects on placenta development and functioning and how these relate to fetal health.

The placenta is a unique and complex temporary organ that nourishes and protects the fetus. It is a multilayered membranous structure that forms at the interface of maternal and fetal circulation and facilitates the exchange of gases, nutrients, metabolites, and waste products.24 Chorionic villi covered by villous trophoblasts (Figure 1A) are the main placental structures involved in the fetomaternal exchanges.25 The inner layer of the villous trophoblasts is formed by stem cells called cytotrophoblasts.26 These cells proliferate and differentiate by fusion to syncytiotrophoblasts, large multinucleated cells that form the top layer of the villous tree.26 Syncytiotrophoblasts are in direct contact with maternal blood and thus are the main cells to facilitate the fetomaternal exchange.27 Syncytiotrophoblasts also function as important endocrine cells involved in biosynthesis and metabolism of a variety of hormones (e.g., leptin, human chorionic gonadotropin, progesterone, estrogen) crucial for the regulation of maternal and fetal physiology.27,28 Growing evidence shows that disrupted functioning of the villous trophoblasts can lead to impaired endocrine, metabolic, and transport functions of the placenta, resulting in a variety of pregnancy complications.29−31

Although syncytiotrophoblasts are the first placental cells in contact
with maternal blood and are crucial for fetomaternal exchange and placental functioning, to date, no data exist on the uptake or toxicity of MNPs in syncytiotrophoblasts in vitro. The BeWo cell line derived from a human immortalized trophoblast, morphologically and functionally resembles villous cytotrophoblasts and is one of the most widely used placental cell models. The BeWo b30 variant forms a confluent and polarized monolayer and has been successfully used in a variety of transport studies including transport of, e.g., gold, silver, silica, iron, or polystyrene nanoparticles. The intercellular fusion and differentiation (syncytialization) of BeWo b30 can be successfully triggered by forskolin, resulting in the formation of endocrine active cells that resemble in situ human syncytiotrophoblasts.

Particle size, morphology, surface charge and protein corona are factors that can influence translocation of MNP over different cellular barriers. Nevertheless, their role in transplacental MNP transport is still not well understood. Moreover, the key characteristics potentially driving toxicity, such as the presence of inherent or adsorbed contaminants, are largely understudied. Due to the challenges in capturing and isolating environmentally weathered MNPs, the majority of toxicity studies have used pristine, manufactured particles that may not be representative of real environmental exposures. The goal of this study was to examine both pristine and experimentally weathered MNPs at different sizes to investigate the influence of polymer type, size, and weathering on the transport, uptake, and toxicity in both nonsyncytialized (undifferentiated) and syncytialized (differentiated) BeWo b30 cells. The model MNPs selected for this study were polystyrene (PS) and high-density polyethylene (HDPE) particles, two commonly used polymers present in a wide variety of consumer products. We investigated the cellular internalization of MNPs, using high-resolution confocal imaging and three-dimensional rendering, and developed a method for quantitative analysis of cellular uptake in both cell phenotypes. Moreover, we investigated the influence of human plasma protein corona on the cellular transport of MNPs in a transwell setup. Furthermore, we investigated the influence of MNP exposure on several toxicity endpoints, including cell viability, damage to the plasma membrane, and expression of genes important in the placental biosynthesis of steroid hormones. Last, to gain the first insight into particle vs. chemical toxicity, we performed an untargeted, small molecule characterization of pristine and weathered MNPs using liquid chromatography high-resolution mass spectrometry (LC-HRMS). This approach allowed us to test for the presence of co-transported pollutants and to evaluate compositional differences following particle weathering.

Materials and Methods
An overview of the experimental design is presented in Figure 1B. A full list of reagents can be found in Supplemental Material, “Reagents.”
**Pristine and Weathered MNPs**

Fluorescent and nonfluorescent MNPs of different sizes were used in this study. Fluoresbrite YG polystyrene 0.05 μm (Cat. no. 17149), 0.2 μm (Cat. no. 17151), 1.0 μm (Cat. no. 17154), and 10.0 μm (Cat. no. 18140-2) particles (Fluo-PS) were purchased from Polysciences Europe GmbH, delivered as 2.5% aqueous suspension. Nonfluorescent MNPs, i.e., an HDPE (HDPE Abifor) powder with nonuniformly shaped particles ranging from 0 to 80 μm in size (Gaussian grain size distribution n: <50 μm = 35%–45%, <63 μm = 60%–80%, <80 μm = 98%–100%; Von Moos et al.43) and PS microspheres at 0.05 μm (Cat. no. 08691), 0.2 μm (Cat. no. 07304), 1.0 μm (Cat. no. 07310) and 10.0 μm (Cat. no. 17136; Polysciences Europe GmbH) were kindly provided by Deltares. Stocks of pristine HDPE and PS particles were prepared by washing the particles in hexane and phosphate-buffered saline (PBS), respectively; then the particles were centrifuged for 10 min at 4,000 rpm at room temperature and resuspended in PBS (1% solids w/v). For all exposure studies, if not specified otherwise, serial dilutions of the highest MNP stocks (0.1, 1, 10, 100 μg/mL) were prepared in the exposure medium (culture medium without phenol red) and added to the cells so that the concentration of PBS did not exceed 1%.

**Chemical Characterization of Pristine and Weathered MNPs**

A nontargeted, small molecule characterization was performed on MNP stock suspensions to evaluate chemical differences between pristine and weathered MNPs and to test for the presence of co-transported pollutants following weathering using previously described methods.44 Briefly, MNP suspensions (pristine and weathered PS with 0.05 μm, 0.2 μm, 1 μm, and 10 μm diameters) were prepared for analysis by diluting 50 μL of 10 mg/mL aliquots with 100 μL acetonitrile and centrifuging for 10 min at 16,100 × g. The supernatants were analyzed (in triplicate) using dual-column (HILIC: Waters XBridge BEH Amide XP HILIC, 16,100 × 10 mm) liquid chromatography paired with positive or negative mode electrospray ionization interfaced to a Thermo Scientific Q-Exactive HFX Orbitrap high-resolution mass spectrometer. Analyte separation for HILIC was accomplished using acetonitrile and formic acid; separation for C18 used an acetonitrile and ammonium acetate gradient. The mass spectrometer was operated at a resolution of 120,000 and mass-to-charge ratio (m/z) range of 85–1,275. Uniquely detected ions consisted of m/z, retention time, and ion abundance referred to as m/z features (all descriptors used in the analysis are explained further in Excel Table S1). High-resolution detection of m/z features was accomplished by a maximum injection time of 100 milliseconds and an Automatic Gain Control target of 1 × 10^5. Raw data files were extracted and aligned using apLCMS45 with modifications by xMSanalyzer.46 Detected signals with a fold-change of 5 or higher relative to the PBS buffer solution were annotated using xMSannotate by matching accurate mass m/z to the plastics and plastic-related chemicals from PLASTICSIGNORMAN; Database of Chemicals likely (List A; 906 chemicals) and possibly (List B; 3,377 chemicals) associated with Plastic Packaging (CPPdb).47 Compound annotation confidence was determined based on Schymsanski et al. by matching accurate mass, isotope, and adduct patterns to the combined plastic database; potential compound structures were ranked based on PubMed reference counts.48,49 Initial differences between pristine and weathered MNPs were characterized using partial least squares discriminant analysis (PLS-DA). Next, a variable importance projection (VIP) score >1.5 was used to identify signals that differed due to MNP weathering. To characterize the potential origin of plastic chemicals with VIP >1.5, annotated compounds were assigned a primary use category based on a database of >10,000 plastic-related chemicals (PlasticMap) recently compiled by Wiesinger et al. from industrial, scientific, and regulatory data sources.50 InChIKeys were used to avoid misclassifications due to differences in chemical names.

**Bewo b30 Culture and Syncytialization**

The Bewo b30 choriocarcinoma cell line (AddexBio; Cat. no. C0030002) was kindly provided by Prof. Aldert Piersma and Conny van Oostrom (RIVM, the Netherlands). The cells were maintained in DMEM/F12 supplemented with 1% (v/v) P/S and 10% (v/v) fetal bovine serum (FBS) and incubated at 37°C with 5% CO2 and 95% relative humidity. Every 3 d, when reaching 70%–90% confluency, cells were washed with PBS and subcultured using 0.05% trypsin-EDTA solution. Differentiation and intracellular fusion (syncytialization) were induced by treatment with forskolin 48 h post seeding when cells reached approximately 50% confluency. A stock solution of forskolin (10 mM) was prepared in dimethylsulfoxide (DMSO) and added to the cells in culture medium to the final concentration of 10 μM at 0.1% DMSO. Medium with forskolin was refreshed daily for 3 consecutive days. An equal volume of DMSO was added as a vehicle control to the nonsyncytialized Bewo cells. Two different cell culture media, MEM and DMEM/F12, were tested to ensure optimal conditions for Bewo b30 cell culture and syncytialization. The cells were seeded in a transparent 12-well plate, cultured, and differentiated in each test media; the cells were then stained and imaged as described below (n = 2, with two technical replicates per plate). In general, undifferentiated cells showed a more consistent morphology when cultured with a medium containing high-glucose DMEM/F12; i.e., cells formed a tighter confluent monolayer and showed clearer cellular boundaries, as indicated with β-actin (Sir-actin) and tight junctions (ZO-1) staining (Figure S1). Therefore, DMEM/F12 was used in experiments throughout this study.

**Cellular Imaging and Quantification of MNP Uptake**

For the visualization of the cells and quantification of the cellular uptake of MNPs, cells were seeded at a density of 2 × 10^4 cells/mL in a transparent 12-well plate containing 18 mm glass covers pre-coated (for 2 h at 37°C) with collagen (type IV) from human placenta (50 μg/mL in 0.25% acetic acid), cultured and differentiated as above. On day 6 post seeding, the nonsyncytialized and syncytialized cells were exposed for 24 h to fluorescent and...
nonfluorescent PS at 0.05, 0.2, 1, and 10 μm size at different concentrations in 1 mL of exposure media. To ensure an optimal nontoxic particle density during imaging, the concentrations used were chosen for each condition separately and kept within the nontoxic concentration range as indicated by cytotoxicity assays in this study. Exposed and not-exposed cells (control) were fixed in 4% formaldehyde (10 min), washed with PBS and treated (for 1 h at room temperature) with blocking buffer (3% BSA in PBS) containing DAPI (1 μM), SiR-actin labeling probe (1 μM), and ZO-1 antibody (1 μM) or Vybrant labeling dye (5 μM), to stain nucleus, β-actin, and tight junction or plasma membrane, respectively. In the case of ZO-1, cells were subjected to the second round of staining with the secondary antibody (Alexa594, 10 μg/mL) for 1 h at room temperature. After staining, the cells were washed with PBS and rinsed in MilliQ, and the glass covers were mounted on a slide for imaging. Cells were imaged using a Leica TCS SP8 STED 3X microscope with a pulsed (80 MHz) white-light laser (WLL), 405 nm DPSS laser, HyD/PMT detectors, and spectroscopic detection. For imaging of 10 μm beads, we used HC PL APO 20x/0.75 IMM CORR CS2 (Leica 15506343) dry objective, and all other conditions were imaged using HC PL APO 93x/1.30 GLYC motCORR STED (Leica 15506417) glycerol-immersion objective with corrective collar. Due to the absence of spectral cross talk, all fluorescent channels were imaged simultaneously. For DAPI imaging, we used 405 nm DPSS laser as the excitation source and detected emission in the 420–460 nm spectral range; for fluorescent beads, we used 488 nm excitation (WLL) and 500–550 nm emission range; for Vybrant dye, we used 561 nm excitation (WLL) and 570–620 nm emission; for ZO-1, we used 580 nm excitation (WLL) and 590–620 nm emission; and for SiR-actin, we used 633 nm excitation (WLL) and 640–750 nm emission. Nonfluorescent beads were imaged using transmitted light PMT with a 488-nm laser as a source. For 10 μm beads, the size of field-of-view (FOV) was ~415 μm, and for all other conditions, it was in the range between 100 and 160 μm. The height of z-stacks was in the range of 10–30 μm, chosen per each sample individually to include the whole thickness of the cell layer and to ensure that the detected MNPs were within the cells and not only present on top of the cell boundary. The number of cells and particles per FOV was counted manually using the Cell Counter plugin for ImageJ (https://imagej.nih.gov/ij/plugins/cell-counter.html). To accurately quantify and compare the particle uptake between the two cell types, the number of particles taken up was calculated per area (square millimeters) occupied by the cells rather than per cell. Mean values of the particle uptake were calculated based on ≥3 FOV per sample from at least two independent experiments (n = 2). To calculate an area occupied by the cells (adjusted cell density) in a FOV, we merged maximum intensity projections of the membrane (Vybrant dye) and actin (SiR-actin) channels, convolved it with a Gaussian filter of 2 pixels (to remove noise), applied threshold manually, and calculated background area using “Analyze Particles” function of ImageJ. Cell occupied area was calculated as the total area of FOV minus background area. Additionally, to image changes in cell morphology during syncytialization, the nonsyncytialized and syncytialized cells were stained with May–Grünewald–Giemsa following a standard protocol and imaged with bright field microscopy.

Transwell Experiments

Nonsyncytialized BeWo b30 cells were seeded at a density of 2 × 10^5 cells/mL on 12-well polycarbonate Transwell® inserts with 12 mm diameter and 3 μm pore size (Corning®, Sigma Aldrich), containing 1.5 mL of culture medium on the basolateral and 0.5 mL on the apical side. The medium was replaced every second day. The formation of the monolayer and the monolayer integrity were monitored using transepithelial electrical resistance (TEER) measurements and sodium-fluorescein (Na-Flu) leakage assay.

**TEER measurements.** Monolayer formation was monitored with a Millicell®-ERS (Electrical Resistance System), a volt-ohm meter that measures membrane potential and resistance of cell monolayers in culture. TEER values were measured for 12 consecutive days in a transwell plate equilibrated to room temperature and in the presence and absence of cells. The final TEER values were determined by subtracting the intrinsic resistance (insert membrane without cells) from the total resistance (insert membrane with cells) and were corrected for the surface area (Ω cm²).

**Na-Flu leakage assay.** The barrier integrity was further evaluated with a fluorescein sodium salt (Na-Flu) leakage assay. A stock solution of Na-Flu (0.5 mM) was prepared in DMSO. At 4, 6, 8, and 11 d post seeding, 0.5 mL of 5 μM Na-Flu (0.1% DMSO) was added to the apical side, and 1.5 mL of medium only was added to the basolateral side. To avoid interference with fluorescence measurements, exposure medium (culture medium without phenol red) was used. After 3 h incubation, 100 μL of the sample was removed in triplicate from the basolateral side, transferred into a black 96-well plate and analyzed using a fluorescent plate reader (POLARstar Galaxy, BMG Labtech GmbH) and 485 nm excitation and 520 nm emission wavelength, respectively. The final concentration of Na-Flu in the basolateral side was calculated using a Na-Flu dilution series (0.1 mM–10 mM), corrected for background fluorescence and volume and expressed as the average basolateral amount of the initial Na-Flu concentration in a percentage, based on three independent experiments (n = 3).

**Transport study.** The transport of Fluo-PS over the monolayer was measured by exposing cells at day 11 post seeding. Fluo-PS at 0.05, 0.2, and 1 μm were added (50 μg) in exposure medium (0.5 mL) to the apical side of the transwell, and their transport to the basolateral side was monitored at different time points: 1, 2, 5, 24, and 48 h, in three independent experiments (n = 3). At each time point, 50 μL of medium from the basolateral side was transferred into a black 96-well plate and the fluorescence intensity was measured with a plate reader (Tecan Infinite M200; Tecan Trading AG), with excitation and emission wavelength set at 441 and 486 nm, respectively. Fluo-PS in three different conditions were used for the translocation studies: noncoated particles (NC), particles precoated with human plasma (P), and particles precoated with heat-deactivated human plasma (DP). We chose plasma, not serum, because it contains clotting factors and thus resembles more closely maternal blood. A whole blood sample was obtained from a healthy donor with informed consent (University Medical Center Utrecht, the Netherlands). The plasma was isolated by centrifuging the whole blood (EDTA-treated) for 25 min at 500 × g at room temperature. The collected plasma was stored at −20°C. Heat-deactivated plasma was prepared by heating in a water bath at 56°C for 1 h. The coating was done by incubating the particles (5 μL of each highest stock) with plasma (1:4 particles to plasma ratio, v/v) for 2 h at 37°C and constant shaking (at 100 rpm). The particles in the plasma (20 μL) were resuspended in the exposure medium before cell exposure to a final volume of 1 mL and particle concentration of 100 μg/mL and final plasma concentration of 1.6%. The mass transported (∆Qp)/ to the basolateral side at each time point (t) was calculated with a calibration curve (0.01, 0.1, 1, 10, 100 μg/mL) prepared for each of the conditions and corrected for blank sample (fluorescent intensity of nonexposed cells) and volume loss during sampling, using the following equation:
\[ \Delta Q_n = C_n \times V_w + \sum_{j=1}^{n-1} C_n \times V_s, \]  

where \( C_n \) is the concentration of the sample measured at time \( t_n \), \( V_w \) is the volume of the well sampled, \( V_s \) is the sampling volume (50 \( \mu \)L), and the term \( \sum_{j=1}^{n-1} C_n \times V_s \) represents the cumulative mass removed by sampling during all the previous sampling periods. \(^{52}\) The final amount transported was reported as a number of particles transported and as a percentage of the initial mass added to the apical well (50 \( \mu \)g). The method’s limits of detection were calculated for each time point separately as 3 times standard deviation of the signal measured in the exposure medium only (\( n = 3 \)).

**Cell Viability**

The viability of the cells was determined using CellTiter-Glo\(^6\) luminescent assay, which measures adenosine triphosphate (ATP) as an indicator of metabolically active cells. The amount of ATP present is directly proportional to the number of viable cells. Briefly, BeWo b30 cells were seeded at a density of 2 \( \times 10^5 \) cells/mL (100 \( \mu \)L per well) in a black 96-well plate. Cells were differentiated and exposed to MNPs at 0.1, 1, 10, and 100 \( \mu \)g/mL on day 6 post seeding as described above. The concentration range was chosen to encompass a wide range of concentrations used in previous studies\(^{17,19}\) and falls within the quantifiable concentration of plastic particles found recently in human blood (1.6 \( \mu \)g/mL)\(^{83}\). After 24 h exposure, the plate was equilibrated to room temperature for approximately 30 min. After that, 50 \( \mu \)L of medium from each well was transferred to a new 96-well plate, and an equal volume of assay reagent was added. ATP disodium salt was used to generate a calibration curve. A stock solution of ATP (10 mM) was prepared in the exposure medium, and the calibration curve was prepared by serial 10-fold dilutions of the stock (5 nM–5 \( \mu \)M) and added in parallel to each plate. Contents were mixed for 2 min on an orbital shaker to induce cell lysis and then allowed to incubate at room temperature for 10 min to stabilize the luminescent signal. Luminescence was measured using a plate reader (Tecan Infinite M200). Data was corrected for the background luminescence measured in the wells containing medium only and expressed as mean percentage of the control (cells with no exposure), based on three independent experiments (\( n = 3 \)).

**Lactate Dehydrogenase Activity Assay**

Lactate dehydrogenase (LDH) activity assay was used to evaluate the damage to the plasma membrane. In short, 100 \( \mu \)L of BeWo b30 cell suspension (2 \( \times 10^5 \) cells/mL) was added to each well of a transparent 96-well plate. Cells were grown and differentiated as described above. At day 6 post seeding, syncylialized and non-syncylialized cells were exposed to MNPs at 0.1, 1, 10, and 100 \( \mu \)g/mL or a positive control Triton X-100 (0.1\%, v/v) in 100 \( \mu \)L of exposure medium. After 24 h exposure, 10 \( \mu \)L of medium from each well was sampled into another 96-well plate containing 40 \( \mu \)L of LDH assay buffer. Also, a NADH standard curve (2.5–12.5 nmol/well) was added in parallel to each plate. Next, 50 \( \mu \)L of the Master Reaction Mix was added to each well, mixed for 2 min using a horizontal shaker and measured (\( I_{\text{final}} \)) at 490 nm in a plate reader (Tecan Infinite M200). After the initial measurement, the plate was incubated at 37°C. Subsequent measurements were taken every 5 min until the value of the most active sample was greater than the value of the highest standard. Data was corrected for the background absorbance measured in LDH assay buffer only, and the LDH activity was calculated using the following formula:

\[ \text{LDH activity} = \frac{B \times \text{sample dilution factor}}{(\text{reaction time}) \times V}, \]

where \( B \) is the amount of NADH (nanomoles) generated between \( t_{\text{initial}} \) and \( t_{\text{final}} \), the reaction time is \( t_{\text{final}} - t_{\text{initial}} \) (minutes), and \( V \) is the sample volume (milliliters). Mean LDH activity is reported as nanomoles/minute/milliliter (milliunit/milliliter), based on three independent experiments (\( n = 3 \)).

**Gene Expression Analysis**

The effects of MNPs on placental hormone synthesis were investigated with real-time quantitative PCR (RT-qPCR) using syncylialized BeWo b30 cells and a set of genes directly involved in the placental steroidogenesis (cyp19, cyp11a1, hsdd3b1, hsdd17b1, st3; see Table S1 for a detailed primer list). BeWo b30 cells were seeded in 24-well plate (2 \( \times 10^5 \) cells/mL), cultured, differentiated, and exposed to MNPs at 0.1, 1, 10, and 100 \( \mu \)g/mL for 24 h in three independent experiments (\( n = 3 \)), with three technical replicates per sample per plate. After 24 h, the cells were washed with PBS, and the RNA was extracted using Tri-reagent following the manufacturer’s protocol. The quality and concentration of the extracted RNA were determined with Nanodrop 2000\(^{®}\) (Thermo Scientific). RNA (0.6 \( \mu \)g) was reverse transcribed using an iScript cDNA synthesis kit, and the cDNA was stored at -20°C until further use. RT-qPCR was performed with a CFX96 real-time PCR Detection System (BioRad) using SYBR Green universal supermix, 10 times diluted cDNA and 400 nM of each primer in a final reaction volume of 15 \( \mu \)L. qPCR reaction conditions were as follows: 3 min at 95°C followed by 40 cycles of 15 s at 95°C and 45 s at 60°C. A melting curve was included in each run to check for primer dimer formation and/or formation of secondary products. The Cq (determined by regression) was normalized to two reference genes \( \beta \)-actin and YWHAZ using the BioRad CFX analysis software, and the log2 fold change in expression (\( \Delta \Delta Cq \)) was calculated relative to the control group (nonexposed cells).

**Statistical Analyses**

Differences in the rate of transport between NC, P, and DP Fluos particles over the nonsyncylialized BeWo b30 monolayer were investigated using the mixed-effects model and Tukey’s multiple comparisons test. For any other statistical analysis, the one-way analysis of variance (ANOVA) followed by Dunnett’s multiple comparisons test was used. All statistical analyses were performed using GraphPad (version 8.3; GraphPad Software, Inc.), with \( p \leq 0.05 \) considered statistically significant.

**Results**

**MNP Chemical Profiling**

To mimic environmental weathering processes and to generate representative MNPs for the toxicological studies, PS and HDPE particles were weathered using the procedures described in the “Materials and Methods” section. This process was hypothesized to alter the surface chemistry of the MNPs and potentially increase the adsorption of organic compounds present in surface water. To gain insight into possible chemical changes occurring to the MNPs from the weathering processes, pristine and weathered PS were characterized for small molecule profiles using non-targeted HRMS. Initial analysis of nontargeted results showed 4,141 and 2,150 chemical signals, with peak intensities 5-fold or higher relative to the PBS buffer, detected for pristine PS and...
A comparison of the two different PS formulations showed that pristine PS had the largest number of unique compounds, 301 were unique to the weathered PS particles, and 1,849 chemical signals were overlapping between the two formulations (Figure 2A). Potential identities of chemicals detected in the pristine and weathered particles were evaluated by annotating mass spectral signals using chemical databases that included plastic-related compounds, followed by PLS-DA to identify which annotated chemicals showed the greatest difference between pristine and weathered PS. Accurate mass annotation included matching based on coeluting adducts and isotopes to reduce false positives and identified 1,062 plastic-related chemicals from which 182 were high confidence (level 3) annotations and 880 low confidence (level 4) annotations (Excel Table S2). The 182 plastic-related chemicals detected in pristine and weathered PS corresponded to 282 unique compounds (Excel Table S2). The number of compounds was higher than the number of detected signals because isomers could not be differentiated using this HRMS approach. Annotated compounds included multiple plastic additives, intermediates in polymer synthesis, hydrolysis products, and environmental chemicals (Excel Table S2). PLS-DA analysis showed significant differences in compound levels associated with pristine and weathered PS and suggested that for pristine PS particles the differences could be mediated by particle surface area (Figure 2B). Primary use categories of compounds annotated with high confidence (level 3) were determined by comparison to a database of >10,000 plastic-related chemicals (PlasticMap). The primary use categories that differed between pristine and weathered particles (with VIP >1.5) are listed in Figure 2C and included, e.g., catalysts, colorants, biocides, antioxidants, plasticizers, and monomers. Use categories for all annotated compounds are provided in Excel Table S3. Compounds present in the PLASTICS|NORMAN and Plastics Packaging Database that were not included in PlasticMap may represent plastic transformation products or environmental chemicals not directly linked to plastic manufacturing.

**BeWo b30 Syncytialization**

Forskolin treatment resulted in a differentiated morphological phenotype evident after 72 h, i.e., progressive nuclear aggregation followed by intercellular fusion (resulting in large multinucleated cells), loss of epithelial cell–cell junctions, and thus loss of barrier integrity of the confluent monolayer (Figure 3 and S1). The tight-junction protein ZO-1, a clear cell boundary marker in the nonsyncytialized cells, was notably less pronounced in the syncytialized cells (Figure S1). Therefore, for further imaging, a plasma membrane marker (Vybrant) was used instead. The syncytialized cells were distinct from the nonsyncytialized cells in morphology and size, i.e., consisted of large often multinucleated cells without clear cell boundaries (Figure 3). The adjusted cell density, calculated as the total area of FOV minus background area (Figure 4; Excel Table S4), showed that the syncytialized cells were on average approximately 3 times less dense...
than the nonsyncytialized cells, with a mean adjusted cell density of 1.964 and 5.559 / mm², respectively.

Uptake of MNPs

Representative images (still frames) of the internalization of Fluo-PS in the nonsyncytialized BeWo b30 cells are presented in Figure 5 (also see Supplemental Video files for the representative 3D videos). Only Fluo-PS particles at 0.2, 1, and 10 μm were investigated, because it was impossible to quantify individual 0.05 μm particles due to their low fluorescent signal. Nonfluorescent MNPs were also investigated, using transmitted light PMT. Here, only 1 μm and 10 μm pristine and weathered PS particles were quantifiable, i.e., clearly visible and distinguishable from other cellular compartments (Figure 6). Moreover, for the nonfluorescent HDPE and PS particles, we observed weak auto-fluorescence in the red part of the spectrum (approximate excitation/emission wavelength of 590/650 nm, Figure S2). However, the auto-fluorescence intensity was too low in comparison with the background cellular auto-fluorescence in the same spectral range to be used for quantification of cellular uptake. Additional imaging using polarized light microscopy (PLM) indicated that HDPE particles were also in contact with the placenta cells and possibly internalized (Figure S3); however, cellular uptake was not quantifiable using this method.

The cellular uptake of 1 μm and 10 μm PS, calculated as a number of particles per area occupied by cells (square millimeters), was concentration-dependent and in general lower in syncytialized than in nonsyncytialized cells regardless of the concentration, type (pristine/weathered), or size of PS (Table 1; Excel Table S4). In general, higher uptake was observed at lower exposure concentrations (e.g., for 10 μm PS, 0.83% and 0.46% of the initial mass were taken up at 0.5 and 5 μg/mL, respectively), indicating that the cell capacity for particle uptake was limited and might have reached saturation levels at the higher concentrations tested. Particle weathering seemed to affect the uptake of PS, resulting in a higher uptake of 1 μm but lower uptake of 10 μm in both nonsyncytialized and syncytialized cells (Table 1; Excel Table S4).

Translocation of MNPs

The development and integrity of the nonsyncytialized BeWo b30 monolayer formed in transwell inserts were monitored with TEER and Na-Flu leakage assay for 12 consecutive days (Figure 7). TEER values showed a steady increase over 9 d, stabilizing at 120 Ω · cm² on day 10 post seeding (Figure 7A). Simultaneously, the leakage of Na-Flu decreased over time, with 1.6% (0.08 μM) of the initial concentration added to the apical well measured in the basolateral compartment at day 11 post seeding (Figure 7B). TEER values and the retardation of Na-Flu transport confirmed completion of the monolayer formation by day 10 post seeding.

As the transwell experiments depend on the measurement of fluorescence, the transport of MNPs was investigated only with the Fluo-PS particles. Fluo-PS particles at 0.05 μm, 0.2 μm, and 1 μm were added (50 μg) to the apical well at day 11 post seeding. Additionally, NC, P, and DP Fluo-PS particles were used to investigate whether the precoating with human plasma would influence the transport kinetics. The biggest size particles (10 μm) were excluded from the transport study due to the use of transwell with a 3 μm pore size, i.e., the biggest pore size that allowed for uniform cell growth without cell migration to the basolateral side. The transport of the particles was monitored over 48 h at six different time points (1, 2, 3, 5, 24, and 48 h). All three sizes of Fluo-PS were transported through the nonsyncytialized cell monolayer (Figure 7C–F; Table S2). The quickest transport over the monolayer was observed for 0.05 μm Fluo-PS with the fluorescent signal (>LOD; Table S2A) measured in the basolateral compartment already 1 h after exposure, followed by 0.2 μm (5 h) and 1 μm particles (24 h) (Figure 7C–F; Table S2B). After 48 h of exposure,
the average number of NC particles transported through the monolayer was $6.0 \times 10^{10}$ (4.13 μg), $4.6 \times 10^7$ (0.2 μg), and $1.4 \times 10^6$ (0.08 μg) for 0.05 μm, 0.2 μm, and 1 μm Fluo-PS, which represented 8%, 0.4%, and 0.16% of the initial mass (50 μg) added to the apical chamber, respectively (Figure 7C–F; Table S2C). For 0.05 μm and 0.2 μm particles, even after 48 h exposure, no flattening off of the number of particles transported was observed, which suggests that longer incubation times would result in higher translocation. No significant differences in the rate of transport were observed between NC, P, and HDP particles at any of the time points measured ($p > 0.05$).

**Cytotoxicity, LDH activity, and effects on steroidogenesis.**

The cytotoxicity of MNPs was investigated using CellTiter-Glo luminescent assay, which measures the amount of ATP present in culture as an indicator of metabolically active cells. None of the pristine or weathered PS (0.05, 0.2, 1, 10 μm) and HDPE particles showed cytotoxic effects in the nonsyncytialized or syncytialized cells, after 24 h exposure to 0.1, 1, 10, and 100 μg/mL concentration (Figure 8A; Excel Table S5). Plasma membrane integrity, quantified by measuring the release of the intracellular LDH into the cell culture medium upon damage to the plasma membrane, was also not affected by any of the conditions tested, except for 0.05 μm PS particles at the highest concentration (100 μg/mL), in the nonsyncytialized cells only (Figure 8B; Excel Table S5). In general, the LDH concentration measured in the cell culture medium was two times higher in the syncytialized cells (~200 mU/mL) in comparison with the nonsyncytialized cells (~100 mU/mL; Excel Table S5).

We also examined the expression of five genes involved in the placental biosynthesis of steroid hormones in syncytialized BeWo b30 cells to investigate the effects of pristine and weathered MNP and associated chemicals on placental steroidogenesis. Of the five investigated genes, only hsd17b1 was down-regulated in cells exposed to pristine MNPs and to a lesser degree weathered MNPs, whereas hsd3b1 was marginally down-regulated only after exposure to pristine particles (Figure 9; Figure S4; Excel Table S6). This trend was observed regardless of the type of size of MNP tested; nevertheless, it was not statistically significant ($p \leq 0.05$).

**Discussion**

Human *in vitro* cell models of the placenta barrier are indispensable for understanding the placental uptake of MNPs and their possible adverse effects on placental functioning and consequently fetal and maternal health. Villous cytотrophoblasts and syncytiotrophoblasts are important cells of placenta villi, with the latter being the main cells responsible for maternal/fetal exchange and other specialized placental functions, such as hormone synthesis and metabolism.28,54 Nevertheless, to our knowledge, the toxicokinetics and toxicity of MNPs in syncytiotrophoblasts *in vitro* have not been reported. This study set out to investigate the uptake, transport, and toxicity of MNPs in both nonsyncytialized and syncytialized BeWo b30 cells, i.e., well-established *in vitro* placenta cell models representative of villous cytотrophoblasts and syncytiotrophoblasts, respectively.

In this study, all PS particles tested, including the large 10-μm particles, were taken up by both nonsyncytialized and...
syncytialized cells. To our knowledge, only one other study, by Cartwright et al., investigated PS uptake in the nonsyncytialized BeWo cells, whereas uptake in syncytialized cells has never been investigated. Overall, the nonsyncytialized cells showed higher uptake than the syncytialized cells. The smaller particles (0.05 µm) were taken up more efficiently than the larger particles, confirming observation by Cartwright et al., and the total uptake was higher at lower exposure concentrations, indicating that the cell capacity for particle uptake was limited and might have reached saturation levels at the higher concentrations and/or larger sizes. This data is supported by ex vivo placental perfusion studies where a size-dependent maternal-to-fetal translocation of PS particles (up to 0.5 µm in size) was observed, with a lower transfer rate corresponding to a higher accumulation of larger PS particles in the placental tissue, specifically in the syncytiotrophoblast layer. Although the staining of cellular compartments together with z-stack confocal imaging and spectroscopic detection allowed for confirmation of cellular internalization of PS particles, the irregularly shaped, nonfluorescent HDPE were not distinguishable from other cellular compartments, and their internalization could not be confirmed with this method. Imaging using PLM indicated that HDPE particles were indeed in contact with the placenta cells and possibly internalized (Figure S3). Further studies are needed to confirm HDPE uptake using other methods or labeled particles.

Transcellular transport of Flu-PS across the nonsyncytialized BeWo b30 monolayer was limited (<10%) and size dependent, with higher transport rates observed for particles with a smaller diameter. Transcellular transport studies could only be performed with nonsyncytialized BeWo b30 cells that formed a tight, polarized monolayer on the transwell insert. Syncytialization resulted in cell fusion and loss of barrier integrity; therefore, MNP translocation using syncytialized BeWo cells could not be investigated. To our knowledge, in vitro transplacental transport of MNPs in nonsyncytialized BeWo b30 monolayers has been previously investigated in four studies, all of which used Flu-PS particles as model MNPs. These studies reported size-dependent and limited transport within 24 h of exposure. It should be noted that there are discrepancies in the scientific literature regarding the timing of BeWo monolayer formation in the transwell setup (ranging from 4 to 10 d post seeding). These differences might stem from different insert types, pore size, pore density, passage number, seeding density, medium type, and volume used, among others. It is, therefore, crucial to monitor the monolayer formation on a day-to-day basis, using more than one indicator, such as a combination of TEER, retardation of Na-Flu transport, or/and confocal imaging. Moreover, it is important to note that the transwell inserts can, to various degrees, retain particles on the apical side, which might be dependent on the pore sizes and insert type. For example, polyester insert membranes have been shown to retain particles with higher efficiency than, e.g., polycarbonate inserts. In this study, the translocation of MNPs through the polycarbonate insert membrane only was significantly higher than when measured with the presence of nonsyncytialized BeWo cells (Table S2), which further confirmed the limited cellular transport of MNPs. In our study, the 3-µm pore size was the largest pore size that allowed BeWo b30 monolayer formation without cellular migration to the basolateral side; nevertheless, it still limits the MNP size range that can be investigated in the transwell setup to <3 µm. However, using imaging, we showed that MNPs up to 10 µm in diameter were internalized by the nonsyncytialized and syncytialized BeWo cells.

In our study, the transport of 0.05 µm and 0.2 µm particles did not reach a plateau at 48 h (Figure 7C and D), suggesting that longer incubation periods may result in higher transport. Different size-dependent transport mechanisms have been proposed for MNPs, including passive diffusion, facilitated diffusion, and energy-dependent transport such as endocytosis. An interesting aspect is that trophoblastic cells also have the capacity for phagocytosis. Nevertheless, the extent to which MNPs are internalized in the placenta has not been investigated to date. Furthermore, this is the first study to consider the transplacental transport of MNPs in the context of placental health. Additional investigations are needed to study how the placenta may respond to MNPs and whether MNPs may cross the placenta or not, which could provide critical insight into potential placental toxicity and fetal exposure.

Table 1. Comparison of the cellular uptake of pristine and weathered polystyrene (PS) particles between the nonsyncytialized and syncytialized BeWo b30 cells.

| Size  | Condition | µg/mL | Nonsyncytialized | Syncytialized |
|-------|-----------|-------|-----------------|--------------|
| µm    |           | Mean (SD) | No. of FOV | Mean (SD) | No. of FOV |
| 1     | Pristine  | 0.1 | 234 (138) | 8 | 50 (38) | 8 |
| 1     | 1         | | 950 (381) | 7 | 337 (246) | 3 |
| 1     | Weathered | 1 | 3,143 (1,917) | 15 | 519 (253) | 15 |
| 10    | Pristine  | 0.5 | 8 (4) | 10 | 4 (5) | 10 |
| 10    | 5         | | 41 (25) | 15 | 26 (14) | 10 |
| 10    | Weathered | 5 | 30 (16) | 15 | 4 (4) | 15 |

Note: Cellular uptake was quantified based on high-content imaging as described in the “Materials and Methods” section. Cells were exposed to pristine (including fluorescent) and experimentally weathered PS particles at 1 µm and 10 µm in size and varying concentrations (micrograms per milliliter). The mean number of particles taken up was calculated per area occupied by the cells (square millimeters), based on ≥3 FOV per sample from at least two independent experiments, ± SD (Excel Table S4). FOV, fields of view; SD, standard deviation.

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which these mechanisms influence MNP transplacental kinetics is still unknown. The size-dependent, limited transport of Fluo-PS observed in vitro is generally in agreement with ex vivo placenta perfusion studies. However, in placenta perfusions, PS particles accumulated mainly in the syncytiotrophoblasts layer, suggesting that syncytiotrophoblasts and not the cytotrophoblasts play a major role in regulating transplacental particle transport. The use of syncytialized BeWo cells may therefore provide more accurate estimations of the uptake or transplacental transport of MNPs. It should be noted, however, that the expression profiles of, e.g., transporter proteins, such as ATP-binding cassette (ABC) transporters and solute carriers (SLC), are not fully comparable between syncytialized BeWo cells and primary human trophoblast cells, likely in part due to the different developmental stages they represent. Overall, carefully validated experimental models are needed to study different aspects of trophoblast function. Furthermore, the use of placental co-cultures, although technically challenging, may more accurately reflect complex cell-to-cell interactions and provide a better understanding of the placental toxicokinetics of MNPs.

The protein corona has been shown to play an important role in the transport of metallic nanoparticles in in vitro cellular models and ex vivo placenta perfusion models (reviewed by Bongaerts et al.) however, the influence of the protein corona on transplacental transport of MNPs is still not well understood. In our study, prior incubation of Fluo-PS with human plasma or DP did not enhance particle transport across BeWo b30 barrier. In contrast, Gruber et al. found that the protein corona formed during incubation of PS with human plasma enhanced placental transfer in a human ex vivo placenta perfusion study. In particular, the protein corona formed by human albumin was found to significantly induce transplacental transfer of drugs as well as PS particles. Moreover, serum heat-deactivation has been shown to affect PS protein corona composition, resulting in enhanced nanoparticle uptake in, for example, the human lung barrier.
In our study, the presence of FBS in the exposure medium might explain why we did not observe enhanced transport. It has been previously suggested that the complex interaction between particles and protein-rich media may lead to progressive displacement of proteins with lower affinity in favor of those with higher. This could change corona composition and dominate the overall properties of the particles, possibly neglecting the influence of prior plasma coating on cellular transport. Further studies are needed to corroborate this hypothesis.

The MNPs tested in this study showed no acute cytotoxicity. No effects on cellular ATP production were observed, whereas plasma membrane integrity was disrupted only after exposure to 50-nm PS particles at 100 μg/mL and only in the nonsyncytialized cells. At this concentration, both weathered and nonweathered particles disrupted the integrity of the plasma membrane; however, such a high concentration might not be relevant, considering environmental exposures. Although human exposure levels are still not well characterized, recently plastic particles were found in human blood at a mean concentration of 1.6 μg/mL.

![Figure 8](image_url)

**Figure 8.** Cell viability (A) and plasma membrane integrity (B) of the nonsyncytialized and syncytialized BeWo b30 cells exposed for 24h to 0.1, 1, 10, and 100 μg/mL pristine and weathered HDPE, PS particles (0.05, 0.2, 1, and 10 μm), and Triton X (TX; 0.1%) as a positive control, measured with CellTiter-Glo® viability assay and LDH activity assay, respectively, as described in the “Materials and Methods” section. Data points represent the mean percentage of ATP present ± SD (n = 3; Excel Table S5) and LDH concentration (mU/mL) ± SD (n = 3; Excel Table S5). Control (C) represents nonexposed cells (one-way ANOVA; \( p \leq 0.05 \)). Note: ANOVA, analysis of variance; ATP, adenosine triphosphate; HDPE, high-density polyethylene; LDH, lactate dehydrogenase; PS, polystyrene; SD, standard deviation.

![Figure 9](image_url)

**Figure 9.** Heat map representing the log2 fold change values in the expression of genes involved in steroidogenesis measured in the syncytialized BeWo b30 cells after 24 h exposure to 10 μg/mL of pristine and weathered HDPE and PS particles (0.05, 0.2, 1, and 10 μm). Values were normalized against the endogenous reference genes β-actin and YWHAZ. Red implies higher expression, white no difference, and blue lower expression relative to the nonexposed cells (n = 3; Excel Table S6). Note: HDPE, high-density polyethylene; PS, polystyrene.
suggesting that the lower concentration range tested in this study (0.1–10 µg/mL) might be relevant with regard to environmental exposures. In the in vitro placenta cell models published to date, PS particles generally do not show cytotoxicity. However, a study by Kloet et al. showed that positively charged 0.05 µm PS particles induced significant cytotoxicity at concentrations ≥20 µg/mL, whereas negatively charged particles did not. The authors argue that the electrostatic attraction between the positively charged particle and the negatively charged cell membrane could explain the cytotoxicity observed. The protein corona formed on the surface of the particle incubated with FBS could dominate the charge of that particle making it slightly negative, which could explain why in most in vitro studies that used media supplemented with FBS, no cytotoxicity is generally observed. Studies with various other human cell models link MNPs exposure with apoptosis, necrosis, genotoxicity and proinflammatory responses (recently reviewed by González-Acedo et al.)). Toxicity endpoints, other than cell viability, should therefore be considered, as well as longer exposure periods that would be more reflective of the possible long-term accumulation of MNPs in syncytiotrophoblasts in situ.

An important function of the placental syncytiotrophoblasts is the production/metabolism of hormones crucial for the maintenance of pregnancy and fetal development. Accordingly, dysregulation of syncytiotrophoblasts has been shown to impact placenta functioning and fetal health. In this study, the effects of MNPs on the expression of genes essential in steroid hormone biosynthesis were tested using syncytialized BeWo b30 cells. From the five genes tested, hsd17b1 was modestly down-regulated in cells exposed to pristine MNPs and to a lesser degree to weathered MNPs, whereas hsd3b1 was marginally down-regulated only after exposure to pristine particles. The enzyme hsd17b1 catalyzes the final step in estrogen biosynthesis by reducing the weak estrogen estrone to yield the potent estrogen 17β-estradiol, whereas hsd3b1 catalyzes the conversion of pregnenolone to progesterone, among others. Though not significantly different relative to controls, the down-regulation of these genes by MNPs was observed in both PS and HPDE at all sizes tested. This finding warrants further investigation using longer exposure periods, because inhibition of hsd17b1 and hsd3b1 has been associated with a reduction of bioavailable levels of these hormones, which may lead to various adverse pregnancy outcomes.

Chemical profiling of PS particles revealed compositional differences between pristine and weathered particles, with the former showing a higher number of unique features. A recent review, based on industrial, scientific, and regulatory data sources, revealed >10,000 plastic monomers, additives, and processing aids associated with plastics. From the plastic-related substances of potential concern (i.e., substances that met one or more of the persistence, bioaccumulation, and toxicity criteria), 58 of the substances were annotated using features detected in the pristine or weathered PS in this study (Figure 2C; Excel Table S3). The primary categories of the plastic-related compounds differing between pristine and weathered particles included 14 use categories (Figure 2C). The presence of these compounds is surprising because the PS particles were single polymer particles, and a large number of impurities was not expected. In-depth chemical characterization of pristine and environmental weathering processes when performing toxicological studies with MNPs.

Recent evidence demonstrates the presence of MNPs in the human placenta and meconium, which suggests that maternal exposure to MNPs may result in placental uptake, transplacental transport, and fetal exposure. However, the underlying toxicokinetics and toxicity are still not fully understood. In this study, cells central to the maternal–fetal interface showed limited uptake of pristine and weathered MNPs in vitro, which was associated with a lack of acute toxicity and subtle effects on placental gene expression. Although plastic-associated compounds or co-contaminants might be relevant for the effects observed in this study, more in-depth investigations are needed to better understand the potential health risks of MNPs and chemicals associated with them under real-life conditions and environmentally relevant exposure scenarios. In vitro placenta cell models together with the development of analytical tools and workflows for detection and characterization of MNPs in maternal samples and explorative cohort studies are indispensable to understanding the potential risk associated with MNPs to maternal and fetal health.

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