



Cite this: *Environ. Sci.: Nano*, 2018, 5, 48

Challenges in characterizing the environmental fate and effects of carbon nanotubes and inorganic nanomaterials in aquatic systems

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The current lack of commonly used protocols for dispersion, characterization, and aquatic toxicity testing of nanomaterials (NMs) has resulted in inconsistent results, which make meaningful comparisons difficult. The need for standardized sample preparation procedures that allow the reproducible generation of relevant test conditions remains a key challenge for studies of the environmental fate and aquatic toxicity of NMs. Together with the further development of optimized and cost-effective analytical techniques for physicochemical characterization that depend on reproducible sample preparation, such methods have the potential to overcome the current uncertainties with regard to NM dispersion properties, effective dose, and particle dissolution. In this review, recent data available on the challenges are summarized, especially those associated with preparing and quantifying NM dispersions, determining NM uptake and accumulation in aquatic organisms, and the transformation of organic and inorganic NM in aquatic species. Additional limitations and challenges that are specific to certain types of NMs are highlighted. The release of highly persistent carbon nanotubes (CNTs) from nanocomposites is determined to be a potential source of environmental contamination. Furthermore, the role of NM dissolution and the contribution of ions *versus* particles to NM toxicity are discussed. A phenomenon of particular relevance for the environment is photoactivation of NMs. This is elucidated with regard to its consequences in complex aquatic ecosystems. Widespread implementation of standardized protocols alongside the consideration of phenomena associated with different life cycle stages of industrial products is crucial to the future establishment of NM environmental risk assessment.

Received 30th June 2017,
Accepted 5th October 2017

DOI: 10.1039/c7en00594f

rsc.li/es-nano

Environmental significance

The environmental fate assessment of nanomaterials lacks standardized analytical methods and scenarios for nanomaterial release. Production of nanomaterials and derived products increases, while potential adverse effects remain unpredictable. Development of suitable analytical methods and consideration of nanomaterials in multicomponent compositions are indispensable prerequisites for the environmental risk assessment of nanomaterials. Before new materials of uncertain effects in aquatic systems are produced in large scale, analytical methods for detection and analysis should be present. The recognition of scenarios that may lead to the release of potentially harmful materials into the aquatic environment contributes to higher risk awareness. Suitable analytical methods and the inclusion of relevant industrial products and their life cycle stages can help to prevent negative effects of nanotechnology on aquatic life.

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Introduction

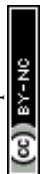
Nanomaterials (NMs) are produced worldwide on a large scale and their applications are steadily increasing.^{1–3} However, there remains considerable concern regarding their release into the environment, fate, behavior, and subsequent potential for eliciting effects in organisms. The potential accumulation of NMs by aquatic organisms has previously been suggested to lead to transfer throughout food chains.^{4–6} While human toxicity and ecotoxicity of NMs have many aspects in common, the study of NM environmental fate and effects presents a number of additional challenges.

Crucially, a lack of suitable standardized methods for NM dispersion, preparation, and characterization has contributed to significant divergences in published data on their ecotoxicity.^{7–9} Parameters such as pH, ionic strength, the presence of biological material such as proteins, and the occurrence of humic acids in environmental media that can adsorb to the NM surface, can have a strong influence on NM agglomeration, persistence, and particle release from nano-enabled products.^{10,11} An increasing number of studies investigated the influence of multiple environmental conditions on the physicochemical properties of NMs, NM environmental behavior, and their subsequent potential for exposure and ecotoxicity to a range of organisms from different environmental matrices. For CeO₂ NMs, it was shown that humic acids prevent agglomeration at low CaCl₂ concentrations and facilitate agglomeration at high concentrations.¹² In the case of multi-walled carbon nanotubes (MWCNTs), agglomeration, physical interaction with cells, and shading were found to contribute to their toxicity toward algae.^{13,14} In a study with the benthic diatom *Nitzschia palea*, a strong increase in MWCNT toxicity by natural organic matter (NOM) was recorded and discussed in relation to the affinity of MWCNTs to biofilms and the potential contribution of MWCNTs to a shading effect.¹⁵ The detection, quantification, and characterization of NM physicochemical properties (e.g., agglomeration, dissolution, shape, chemical reactivity, and adsorbed molecules) in the various environment matrices and relevant exposure scenarios present many challenges and in many cases require new or improved methods and technologies. The current limitations with available sample preparation methods and analytical tools used to address environmental fate and aquatic toxicity of NMs are described in this article. Furthermore, data on NM accumulation, dissolution, and release from nano-enabled products are reviewed with regard to uptake and accumulation in the food chain and potential adverse effects in the aquatic environment. Additional limitations and challenges that are specific to certain types and groups of NM are highlighted. For example, recent findings on NM photoactivation, a phenomenon with specific relevance for aquatic organisms, are summarized.

Dispersion and characterization of nanomaterials in aqueous media

One of the biggest challenges with interpreting and comparing the large volume of published aquatic ecotoxicity data is the wide range of dispersion techniques and conditions employed.¹⁶ Most studies use energy, in the form of shaking, stirring, or sonication, to generate aqueous NM dispersions. However, organic solvents such as tetrahydrofuran, pyridine, methanol, ethanol, propanol, and dimethyl sulfoxide have been used to disperse nanomaterials (e.g., C₆₀), although careful control of the solvent concentration is necessary to avoid negative impacts on organisms employed in nanotoxicity studies.^{17,18} This has led to inconsistent or even contradictory findings in different studies using the same NM and ecotoxicity assay.¹⁹ Reproducibility of NM dispersion in exposure media employed in aquatic ecotoxicity tests, and appropriate methods to characterize NMs in such dispersions, are crucial for obtaining accurate results that can be interpreted meaningfully.^{20–23} A recent study used the same titanium dioxide (TiO₂) NMs (38 ± 10 nm) to investigate the effectiveness of four dispersion protocols that had been developed for past research projects.¹⁹ Variations among the four different protocols included the use of pre-wetting, the type of sonicator, sonication duration and power, cooling, and particle concentration. The results revealed a large degree of variability of the mean TiO₂ particle dispersion diameter among the four protocols with a relative standard deviation of 26%. The authors identified particle concentration as well as sonication conditions (power and duration) as the main parameters influencing the final dispersion characteristics. Although sonication conditions are an important parameter in determining dispersion quality, it is clearly important to consider other parameters, such as particle concentration, age of the dispersion, and subsampling as potential sources of variability.¹⁹

There is currently a lack of internationally-recognized standard dispersion protocols for NMs, but available information on protocols for the preparation of stock suspensions has recently been summarized.²⁴ The need for clear and comprehensive guidelines for preparation of NM dispersions has been highlighted, together with the importance of a well-controlled sonication protocol.^{24–29} Standardizing the energy delivered to the system during dispersion is a key step towards achieving reproducible aqueous NM dispersions.^{30,31} Furthermore, the availability of internationally-accepted reference materials and benchmark data for assessing NM dispersion reproducibility, both within and across laboratories, is required. The generation of individual benchmarked reference datasets for the huge range of aquatic conditions studied in environmental fate and effects assessment is not feasible, yet it is important such studies are conducted using relevant environmental parameters. It is therefore suggested that initial standardization strategies focus on establishing reproducible stock dispersions in deionized water, which can be further diluted when specific media types are required in



ecotoxicity tests or that mimic natural water bodies. This approach could overcome the issue of conducting tests on unnaturally dispersed NMs, which is likely to overestimate the exposure occurring in the natural environment.

Independent of their size, NMs of the same mass and chemical composition can have a completely different toxicity per unit mass.³² For toxicity testing of NMs, the metric for dose quantification needs to be carefully chosen. Particle volume, mass, surface area, and number have been used for this purpose.³³ Each metric may provide a useful perspective. Delmaar *et al.*, have outlined a simplified dose metric related to particle diameter for spherical SiO₂ and Ag NMs.³² It appears unlikely that there is a single dose metric that is appropriate for all NMs and test systems.^{32,34} In aqueous suspensions, atomic force microscopy, scanning or transmission electron microscopy, and ultrafiltration are often applied for NM characterization.³⁵ Field-flow fractionation and single particle ICP-MS are increasingly being used for NM size estimation in environmental samples.^{36,37}

Carbon nanotubes

This review emphasizes carbon nanotubes (CNTs) for a detailed assessment of NM dispersion and characterization in aquatic media. CNTs present many of the same challenges observed with metal and metal oxide NMs, but also some additional factors unique to this family of materials. They represent the most commonly studied carbon NM, although environmental studies with fullerene and graphene families have also been reported.^{38–40} The preparation and characterization of CNT dispersions is challenging and has proven difficult to apply accurately and reproducibly in environmental assess-

ment. The inherent physicochemical properties of CNTs mean that their dispersion, quantification, and characterization in aqueous environmental media and ecotoxicity tests require alternative approaches to those routinely employed for many metal and metal oxide NMs. Their hydrophobic nature and relatively large particle size means that CNTs disperse extremely poorly in water, although surface chemistry (*e.g.*, –OH and –COOH functionalities) can reduce hydrophobicity, thereby increasing dispersion concentration (Fig. 1) and stability.^{41–47} CNT dispersibility and stability are further reduced with increasing ionic strength of natural waters and ecotoxicity media.^{38,43,48–50} NOM can significantly increase CNT dispersion concentration and stability in aqueous media.^{38,48,51–53} As such, some forms of NOM are increasingly employed in ecotoxicity studies with CNTs^{53–55} as well as inorganic NMs.^{56,57} Finally, CNT dispersions prepared using sonication will generate higher, more stable concentrations than shaking or stirring,⁵¹ although it is likely to result in breakage of the tubes and generation of artifacts depending on the sonication time and energy.^{18,58}

Many previous studies on CNT ecotoxicity employed specific and diverse sonication methods for dispersion.^{4,59,60} Differences in dispersion methods can lead to significant deviations in the final dispersion concentration, the agglomeration/aggregation state of the CNTs, and the degree of breakage.^{58,61} The issue of CNT breakage is rarely considered in reported CNT ecotoxicity studies, even though such damage has been shown to alter CNT behavior within the context of (eco)toxicological testing. For example, MWCNTs dispersed by sonication have been shown to be more toxic to the fresh water flea *Ceriodaphnia dubia* and the copepod *Tigriopus japonicus* than CNT dispersions prepared by stirring

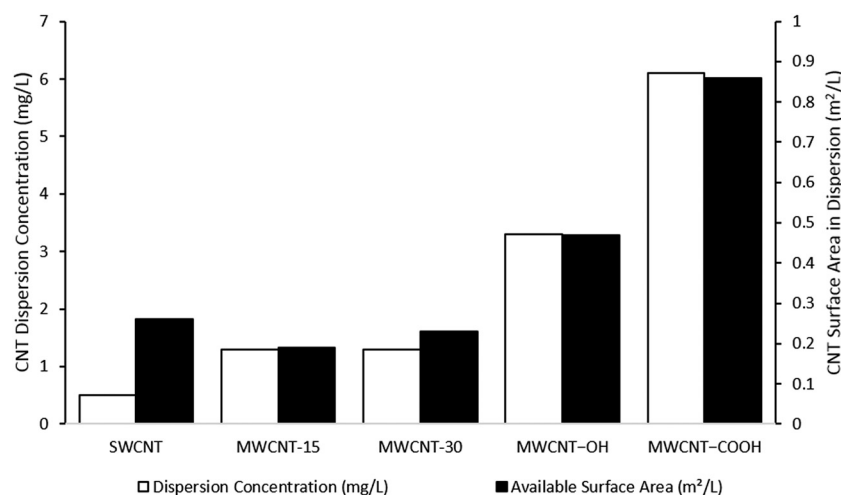


Fig. 1 Carbon nanotube dispersion in algal ecotoxicity media. Dispersion of one single-walled carbon nanotube (SWCNT), two non-functionalized multi-walled carbon nanotubes (MWCNT-15 and MWCNT-30), an OH-functionalized MWCNT and a COOH-functionalized MWCNT in algae ecotoxicity media (OECD technical guideline TG201) containing 20 mg l⁻¹ of dissolved and filtered Suwannee River natural organic matter. The non-functionalized and functionalized MWCNTs have similar diameters (15–20 nm). Data presented as mass- and surface area-based metrics. Reprinted with modifications from B. Glomstad, D. Altin, L. Sørensen, J. Liu, B. M. Jenssen and A. M. Booth, Carbon nanotube properties influence adsorption of phenanthrene and subsequent bioavailability and toxicity to pseudokirchneriella subcapitata, *Environ. Sci. Technol.*, **50**, 2660–2668, Copyright 2016, with permission from ACS.



or shaking.^{62,63} In turn, these parameters affect the final exposure conditions of CNTs, rendering interpretation and comparison of ecotoxicity data generated challenging.

A recent study using a range of non-functionalized MWCNTs investigated the influence of different dispersion preparation techniques on both the comparability of the dispersions and the determination of CNT concentration by UV/vis absorbance.⁶⁴ A sonication process for calibration dispersions using an ultrasonic probe for delivery of acoustic energy was optimized. Verification was conducted by preparing dispersions using an ultrasonic bath, and a procedure to select an appropriate wavelength for each type of MWCNT was also proposed. Results of the study clearly demonstrated that UV/vis absorbance is highly dependent on the dispersion method implemented. Dynamic light scattering was used to determine MWCNT agglomerate size, and revealed that the ultrasonic probe produced dispersions with lower MWCNT agglomeration levels (Fig. 2A) than the ultrasonic bath (Fig. 2B). This study has contributed to the development of standardization in the environmental assessment of CNTs. The European project NANoREG⁶⁵ has also addressed the issue of dispersion reproducibility by developing a standard operating procedure (SOP) for NM dispersion preparation and characterization in environmental fate and ecotoxicity studies.

The separation and characterization of CNT dispersions in both laboratory ecotoxicity tests and in complex environmental/biological matrices represent additional challenges compared to other NMs. Relevant parameters for characterization

in standard ecotoxicity tests include dispersion concentration (especially over time in an exposure), specific surface area (SSA), and the degree of aggregation/agglomeration (*i.e.*, changes in average particle size distribution). Elemental-based quantification techniques (*e.g.*, inductively coupled plasma mass spectrometry) routinely used for inorganic NMs are unsuitable for quantification of CNT dispersion concentrations, and thus alternative approaches are necessary.⁶⁶ UV/vis spectroscopy represents the most common method for CNT quantification in aqueous samples. Despite advantages with respect to time and cost efficiency, this approach has limitations such as potential shading issues, interference from complex environmental matrices, and challenges establishing external calibration curves.^{41,47,67–70} The latter must be based on CNT dispersions with accurately known concentrations.⁵¹ Furthermore, variable wavelengths for absorbance measurements have been reported in the literature indicating that an optimized method has yet to be identified. Thermal analysis techniques (*e.g.*, thermogravimetry, chemothermal oxidation, or thermal optical transmittance) can also be used to quantify CNTs in simple aqueous dispersions,^{51,71–74} but are limited in specificity to particular CNTs or for more complex environmental matrices.^{67,71} The high aspect ratio of CNTs means dynamic light scattering techniques are unsuitable for generating accurate particle sizes in dispersion.^{42,46,75} Scanning and transmission electron microscopy (SEM; TEM) imaging represent better options for determining more relevant particle size data and particle size distributions. However, TEM/SEM imaging of CNTs requires manipulation of the sample prior to analysis, which can significantly change CNT dispersion properties (*e.g.*, aggregation). Vitrification and cryo-analysis have previously been suggested as methods to help overcome this issue,⁷⁶ but both approaches also have the potential to significantly disturb the dispersion. CNTs dispersed in complex aqueous environmental samples may undergo a significant change in SSA due to aggregation and adsorption of NOM.^{44,77,78} However, standard methods for SSA measurement of dry NMs (*e.g.*, BET) cannot be used for direct SSA measurements of CNTs in dispersion, meaning determination of this parameter currently remains unachievable. Thus, determination of SSA currently remains elusive.

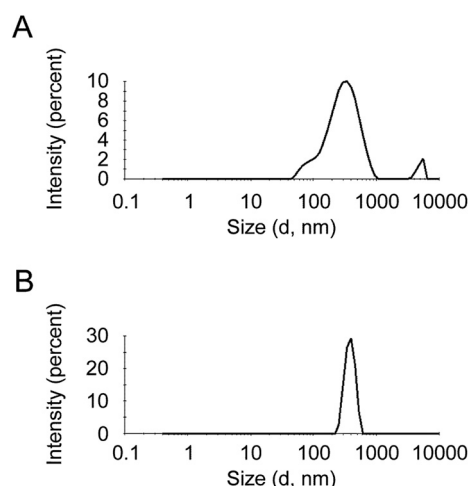


Fig. 2 Multi-walled carbon nanotube distribution following ultrasonic probe and bath dispersion. Size distributions by intensity of MWCNT (nanocyl NC7000) agglomerates in (A) calibration dispersion (5 mg l⁻¹), obtained with ultrasonic probe and (B) verification dispersion (5 mg l⁻¹), obtained with ultrasonic bath. (A) z average, mean 269.5 nm, PDI mean 0.380, (B) z average, mean 678.1 nm, PDI mean 0.604. Reprinted with modifications from C. Cerrillo, G. Barandika, A. Igartua, O. Areitioaurtena, A. Marcaide, G. Mendoza, Ecotoxicity of multiwalled carbon nanotubes: standardization of the dispersion methods and concentration measurements, *Environ Toxicol Chem.*, **34**(8), 1854–1862, Copyright 2016, with permission from John Wiley and Sons Inc.

Release of particles from nanocomposites and coatings

Polymers are frequently enabled with NMs in order to obtain properties such as increased mechanical and barrier strength, biocidal activity, or to repel water and dirt. While materials in which particles, platelets, or fibers dispersed in a polymer matrix are called nano-composites, surface coating represents a further common functionalization technology. Inorganic particles such as nano-Ag and nano-TiO₂ are common components of polymeric materials of textiles, food contact materials, and other products of daily use. CNTs are dispersed in polymer matrices in low amounts (usually <5 weight%) to transfer some of their beneficial properties to



the plastic matrix, mainly mechanical strength,⁷⁹ *e.g.*, for motor helmets or tennis rackets, as well as electrical conductivity, *e.g.*, for flexible electrodes, antistatic coatings, or piezoresistive sensors.⁸⁰ It was assumed that *ca.* 0.2% of the 300 million tons of plastic material produced worldwide each year represent composites with embedded NMs.⁸¹ One of the main questions is whether and under which conditions NMs from such products might reach the aquatic environment. Release of NMs from nanocomposites and coatings can occur during the use phase, *e.g.*, by wear or washing, by drilling or sanding, as well as by environmental weathering of composites after disposal, *e.g.*, by solar irradiation and leaching processes.^{82–86} A study on the release of silver from nano-enabled textiles into artificial sweat demonstrated that coated products release a higher amount compared to nanocomposites, where particles are integrated in the polymer.⁸⁵ Although release of nano-Ag was observed, this was not related to

the primary particle size and chemistry of the silver compounds used for textile functionalization. The finding that NMs may not only originate from direct release of elemental silver but also from reduction of silver ions was confirmed by experiments on textile washing in which metallic Ag NMs were detected in experiments in which AgNO₃ was introduced.⁸⁶

An analysis of NM release due to mechanical treatment was recently conducted with 11 acrylate coatings and 11 polypropylene composites.⁸⁷ The samples were equipped with 10 pigments: TiO₂, carbon black, and 8 nanoscale pigments. High numbers of released NMs were observed, however, no correlation to the primary pigment particle size was found. Standardized scenarios for measurement of NM release to air during occupational and daily life still remain to be developed. A protocol was recently established that reproducibly simulates the drilling of polyamide-6 nanocomposites containing 30 weight% of glass fiber and different

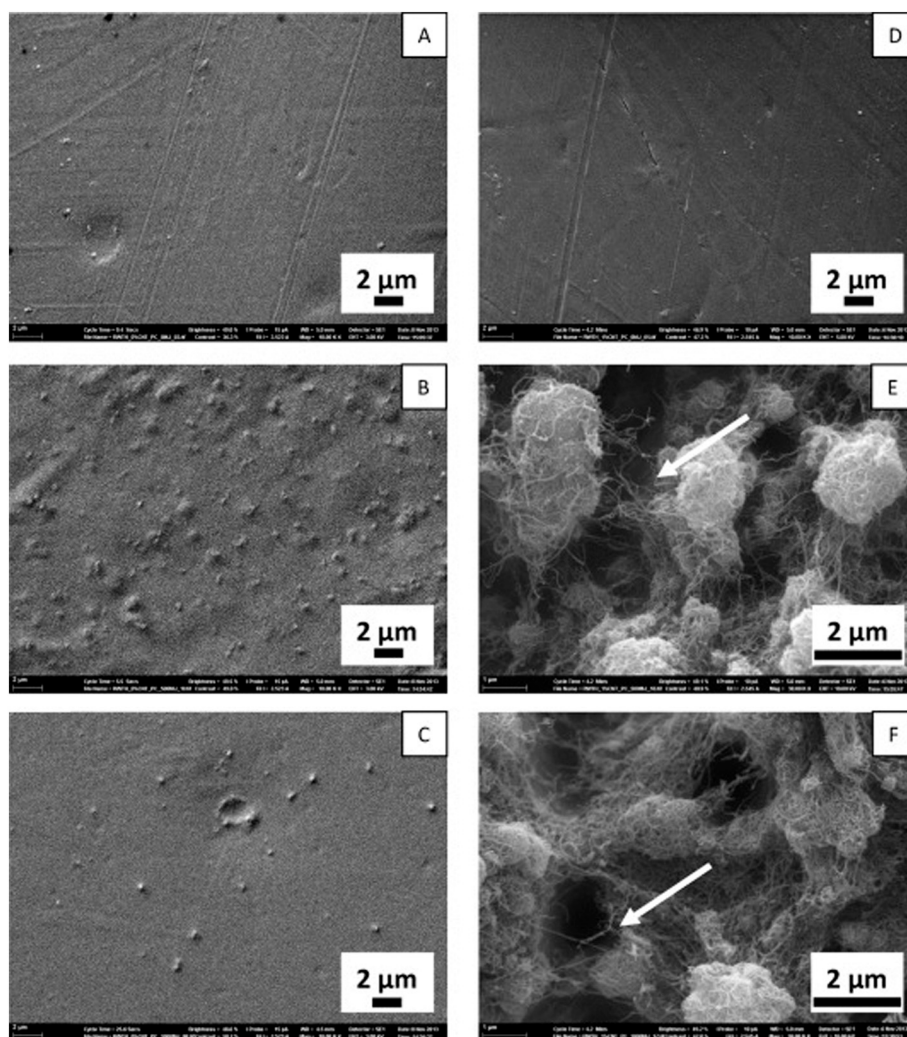


Fig. 3 Scanning electron microscopic pictures of samples of carbon nanotubes (CNT) containing polycarbonate (PC) composites and controls with no CNT material; A: PC without CNT not irradiated; B: PC without CNT after 500 MJ m⁻² irradiation with artificial sunlight; C: PC without CNT after 1000 MJ m⁻² irradiation; D: PC with 1 weight% CNT not irradiated; E: PC with 1 weight% CNT after 500 MJ m⁻² irradiation; F: PC with 1 weight% CNT after 1000 MJ m⁻² irradiation. The arrow points to a single CNT.⁹²



concentrations of nano-SiO₂ or organically modified montmorillonite.⁸⁸

Degradation of CNT-containing composites induced by photooxidation under artificial sunlight has already been described.^{89–91} CNTs were uncovered during degradation of the embedding matrix and formed a network on the weathered surface resulting in potential release from the composite (Fig. 3).

However, exact quantification of the released amounts upon environmental weathering is not possible without using a labelling approach. Rhiem *et al.*⁹² used ¹⁴C-labelling of the carbon skeleton of uncoated CNTs to establish a mass balance. They prepared a black CNT-polycarbonate composite sample (on average 138 mg per sample) containing 1 weight% of multi-walled ¹⁴C-CNT. After irradiation with simulated sunlight (lamp 220 W m⁻², cut-off at 300 nm; total irradiation 1000 MJ m⁻²) and gentle tapping of the sample to remove loose material, further environmental scenarios were consecutively simulated: (a) 21 days shaking in water with six water exchanges during this period, (b) temperature stress (one week 70 °C, a temperature that black plastic material can achieve under natural sunlight), (c) freeze–thawing for one week (–22 °C, once per day thawed at 20 °C), (d) one week shaking in humic acid in water (25 ppm, pH 11.3), one week shaking in artificial acid rain (pH 2.6), and (e) one week shaking in artificial disposal site effluent prepared according to Rhiem *et al.* and Kalbe *et al.*^{92,93} The results in Table 1 were obtained with respect to released material (A: sample irradiated (conditions see above), B: sample not irradiated).

Referring to the initial amount of 1 weight% CNT per composite sample, *ca.* 1.0 ± 0.2% of the CNTs were released from irradiated samples – equivalent to about 64 mg CNT equivalents per square meter composite surface – and only 0.03 ± 0.02% from non-irradiated control samples after all environmentally-relevant treatments. SEM analysis revealed that single CNTs, CNT agglomerates, and CNTs embedded in small matrix particles were released under these scenarios. In line with this release rate,⁹⁴ approximately 100 mg CNT per m² per year would be released at the surface of polyoxymethylene composites containing 5 weight% CNT.

CNTs released from the composites will have a very long half-life in the environment. In experiments with horseradish peroxidase, an enzyme known to catalyze the oxidation of aromatic compounds, a half-life of 80 years was determined by

the slow mineralization of ¹⁴C-labelled CNT and quantification of formed ¹⁴C-CO₂.⁹⁵

Uptake, accumulation, and transformation of nanomaterials in the environment

Development of analytical methods. One of the key challenges in the qualification and quantification of NM uptake and accumulation in aquatic organisms is confirming that internalization has occurred. Another issue is the measurement of specific physicochemical properties of NMs after they have been taken up and/or accumulated by aquatic organisms. Qualitative identification and analysis of internalized NMs can be achieved using techniques such as TEM or SEM. These approaches allow the visualization of NM location at the tissue, organ, and cellular level.⁹⁶ However, sample preparation for EM, as well as the imaging of the sample, requires that the sample be under vacuum, which may introduce artifacts that can alter the true environmental state of the NM.^{14,23} For translucent organisms such as *Daphnia magna*, high resolution light microscopy can be readily used to detect a wide range of NMs as large agglomerates.^{97–100} Time-of-flight secondary ion mass spectrometry allows for the chemical imaging of internalized NMs and the surrounding biomolecules.¹⁰¹ This technique provides a spatial resolution down to 60 nm on nanoscale depth.¹⁰² Synchrotron X-ray radiation tools also appear very promising for investigating the internal fate of NMs in tissue samples, as well as NM behavior in different media such as water or sediment.^{103,104} Other visualization techniques include the use of fluorescent particles together with confocal laser scanning microscopy, hyperspectral imaging, and coherent anti-Stokes Raman scattering microscopy.^{99,105–109} The advantage of these newer techniques compared to TEM and SEM is that particles can be tracked inside whole organisms, diminishing the artefacts related to sample preparation.¹⁰⁴ Although being able to image particles *in vivo* increases our qualitative understanding of NM accumulation within tissues, methods for accurate quantification remain more challenging. Also, a limitation of TEM and SEM is that only a tiny fraction of the sample can be analyzed. Importantly, approaches that minimize sample preparation and analysis artefacts are necessary. Studies on the trophic transfer of NMs remain scarce, and factors leading to this transfer are poorly understood.¹⁰⁴

Accumulation and transformation of CNTs. Uptake of single-walled carbon nanotubes (SWCNTs) and MWCNTs and functionalized MWCNTs has been demonstrated for a

Table 1 Release of radioactivity (single carbon nanotubes (CNTs), agglomerated CNTs, and matrix particles with embedded CNTs) from nanocomposite samples which were (A) previously irradiated by simulated sunlight or (B) untreated control samples during different consecutive degradative treatments (see text)

	Tapping	Shaking (water)	70 °C	Freeze–thaw	Shaking (humic acid)	Shaking (acid rain)	Shaking (disposal effluent)	Tissue wiping	Sum
A (μg)	0.01 ± 0.01	2.6 ± 1.6	0.07 ± 0.004	0.77 ± 0.57	1.87 ± 0.76	2.50 ± 0.85	2.75 ± 2.57	4.12 ± 1.53	14.70 ± 1.86
B (μg)	—	0.1 ± 0.1	0.004 ± 0.004	0.002 ± 0.002	0.03 ± 0.03	0.04 ± 0.01	0.19 ± 0.27	0.02 ± 0.02	0.40 ± 0.27



number of aquatic species.^{9,110–113} Owing to the propensity of many CNT types (and NMs in general) to aggregate and settle in aquatic systems, however, there is concern that they might affect benthic organisms and sediment systems more strongly than pelagic species, despite the latter being more frequently studied.¹¹⁴ Despite strong evidence for CNT uptake, there is currently limited data available with regard to their potential bioaccumulation.¹¹⁵

Some recent studies on functionalized and non-functionalized MWCNT accumulation have employed single-celled organisms such as algae,³⁹ bacteria, and protozoa.¹¹⁶ In the case of bacteria and protozoa, an approach employing separation of ¹⁴C-labeled functionalized MWCNTs by density gradient centrifugation was described.¹¹⁷ The bacterium *Pseudomonas aeruginosa* was found to adsorb the functionalized MWCNTs at concentrations of 0.18 ± 0.04 and 21.9 ± 4.2 μg per mg dry mass at respective nominal concentrations of 0.01 and 1 mg L⁻¹. In the same study, an accumulation of up to 0.9 ± 0.3 μg of functionalized MWCNT per mg dry mass was recorded in the protozoan *Tetrahymena thermophila* following trophic transfer via MWCNT-encrusted *Pseudomonas aeruginosa* cells, while up to 3 ± 1 μg of functionalized MWCNT per mg dry mass was detected following direct uptake. Thus, protozoa were identified as a potential vector for the transfer of functionalized MWCNTs to the next trophic level.¹¹⁶ The alga *Desmodesmus subspicatus* was found to contain mean concentrations of 1.3 ± 0.5 , 2 ± 2 , and 5 ± 2 μg non-functionalized MWCNTs per mg dry weight following exposure to a suspension of 1 mg L⁻¹ of MWCNTs for 24, 48, and 72 h respectively.⁸¹

Daphnia magna has been shown to internalize functionalized MWCNTs. Following a 48 h exposure to a suspension of 0.4 μg per ml, 63 ± 15 μg of the test substance was recorded per mg dry mass.⁹⁸ Previously, a value of approximately 29 μg mg⁻¹ dry mass was reported, following a 48 h exposure of *D. magna* to a 30 μg L⁻¹ fullerene suspension.¹¹⁸ Interestingly, no facilitation of functionalized MWCNT excretion by NOM was observed, which has been described to facilitate suspension of CNTs.⁹⁸ The mean level of non-functionalized MWCNTs in whole fish, following 48 or 168 h exposure to 1 mg L⁻¹ test medium, was determined to be 73 ± 93 ng per mg dry weight, indicating a low bioaccumulation.¹¹⁹ Factors controlling CNT accumulation and retention are poorly understood, but the role of NOM appears to be important. For example, π - π interaction and hydrogen bonding have been suggested as the predominant interactions between CNTs and negatively-charged biocolloids.¹²⁰ These forces were further found to regulate the interaction of CNTs with humic acids, natural biopolymers, and model solid-phase polymers in a systematic comparison by Zhao *et al.*,¹²¹ who described aromaticity and surface polarity as the most positive factors for CNT retention. A recent review by Hu *et al.*³⁹ emphasized the need to consider the complete range of techniques for extraction, isolation, and characterization of CNTs in order to follow their transformation in natural environments and organisms. However, in the case of CNTs, few data on struc-

tural degradation in the environment have been reported. A strain of the bacterial species *Trabusiella guamensis* isolated from soil was shown to cause surface oxidation and structural changes in MWCNTs.¹²² Furthermore, a lignin peroxidase isolated from the mushroom *Sparassis latifolia* was shown to biodegrade SWCNTs.¹²³

Current methodologies for isolating CNTs and their transformed forms from complex matrices are still limited, whilst techniques for accurately characterizing and quantifying CNTs also remain underdeveloped.^{23,98,117,124–126} Furthermore, the presence of low CNT concentrations in environmental matrices and the difficulty in distinguishing natural sources of carbon from those corresponding to CNTs compound the challenges of separation and analysis. The challenge of determining CNT uptake and accumulation in organisms and environmental samples highlights the necessity of using radiolabeling in laboratory studies.¹²⁷

Accumulation and transformation of inorganic nanomaterials. As an example of inorganic granular biopersistent NMs, CeO₂ NMs were shown to adsorb to phytoplankton, an important food source for marine organisms, within <1 h of exposure at 0, 1, 2, and 3 mg L⁻¹. Electrostatic attraction between the charged surfaces was assumed as a mechanism rather than active uptake or a chemical process.¹²⁸ In a study performed with the freshwater algae *Pseudokirchneriella subcapitata*, up to 38% of the total cerium became directly associated with the cells during the 72 h exposure to concentrations between 15 and 200 μg L⁻¹ polyacrylic acid stabilized CeO₂. At the same time, the concentration of dissolved cerium in the exposure suspensions was very low (0.5–5.6 μg L⁻¹), suggesting that toxicological effects derived from dissolved ions are minimal.¹²⁹ Both results confirm an affinity of CeO₂ for algal cells; however, this may depend on exposure concentration and particle modification. The freshwater microcrustacean, *Daphnia pulex*, was shown to accumulate in a water suspension of 10 mg L⁻¹ of CeO₂ NMs a total amount of $24 \pm 5\%$ Ce per g dry weight when fed with algae (*Chlorella pseudomonas*).¹³⁰ In contrast, a reduced uptake of $7 \pm 3\%$ Ce uptake per g dry weight was recorded when no algae were given. This demonstrates the importance of CeO₂/algae interactions for NM uptake by daphnids. It was further demonstrated by X-ray fluorescence and SEM that the shedding of the chitinous exoskeleton is the main mechanism governing the release of CeO₂ by *D. pulex* independent of the feeding regimen. This confirms the crucial role of chitin for transfer of NMs in aquatic environments.¹³⁰ There have also been reports of a relationship between metal ion uptake and NM biokinetics in *D. magna*.^{103,131,132} Exposure of daphnids to TiO₂ NMs at a concentration of 1 mg L⁻¹ was shown to increase the uptake of cadmium and zinc ions from the dissolved phase.¹³¹ The effect, which declined following TiO₂ NM clearance from the gut, may be linked with the creation of sorption sites for metals within *D. magna* by TiO₂. In contrast, it has also been shown that calcium ions (Ca²⁺) affect polyacrylate-coated TiO₂ NM (PAA-TiO₂-NM) bioaccumulation by *Daphnia magna*.¹⁰³ At low ambient Ca²⁺ concentrations, the



well-dispersed NMs were found to be taken up by endocytosis and passive drinking, which revealed the highest accumulation in the abdominal zone and in the gut. On the other hand, there was an active uptake of micrometer-sized aggregates observed at high concentrations that were concentrated in the gut exclusively. Furthermore, PAA-TiO₂-NMs were shown to adsorb cadmium independently of the Ca²⁺ concentration.¹³² There was a distribution of the element throughout the daphnids observed at low Ca²⁺ concentrations, associated with subsequent dissociation of metal-NP complexes and reduced accumulation of cadmium. In contrast, there was concentration of both PAA-TiO₂-NMs and Cd in the daphnid gut induced at high Ca²⁺ concentrations.¹³² Similar to TiO₂ NMs at low concentration,¹⁰³ nano-ZnO and ZnSO₄ were also shown by TEM analysis to distribute within gut, epithelial cells, and tissues behind the epithelial barrier of *D. magna* at a concentration of 0.3 mg Zn per L.¹³³ While an accumulation was observed between 48 h and 9 d of the exposure period, the particle number remained constant between days 9 and 21 of the experiment, a phenomenon that was tentatively attributed to NM dissolution and translocation. Interestingly, cellular uptake of ZnSO₄ NM could be shown. This finding, together with the occurrence of empty vacuoles with disordered membranes following ZnSO₄ treatment, indicates the occurrence of particle specific effects.¹³³

With respect to NM accumulation in fish, TiO₂ particles were mainly identified in the kidney, liver, gills, and to a lesser extent in muscle of the marine species *Trachinotus carolinus* following intraperitoneal injection of 1.5 and 3.0 µg g⁻¹ bodyweight.¹³⁴ Animals were exposed for 24, 48, or 72 h; the size of the employed TiO₂-NM was estimated by TEM as being 11–40 nm. As no size distribution was reported for the injected suspension, which was described as containing primary NMs and larger agglomerates/aggregates, no conclusion on a relationship between size and distribution is possible from this study.¹³⁴ Facilitated uptake of larger-sized NMs is indicated in a study by Chen *et al.*,¹³⁵ who observed enhanced accumulation of agglomerating nano-Fe₃O₄ in medaka fish (*Oryzias latipes*) when compared to the stable colloid Fe⁰. Fe₂O₃ NMs were further reported to adhere to the surface of zebrafish (*Danio rerio*) embryos during toxicity testing.¹³⁶ In line with these results suggesting a facilitated uptake of larger particle sizes, an apparent sorption of nano-Ag of a primary particle size of 50 nm into exposed carp embryos was described for 400 nm, but not for 200 nm, agglomerates.¹³⁷ In the same study, a relationship between NM agglomeration and toxicity has been described for common carp embryos. While there was no dose–response relationship observed in nano-Ag supplemented media with Ag concentrations of 5, 10, and 25 µM comprising aggregates up to a size of 200 nm, there was a clear dose–response observed for concentrations of 10, 25, and 50 µM when agglomerates up to a maximum size of 400 nm were present.

An *in vitro* barrier model was recently suggested to elucidate NM fate in the intestinal epithelial cells of fish. For this purpose, intestinal cells of rainbow trout (*Oncorhynchus*

mykiss) were grown as monolayers on permeable membranes, establishing an upper (apical) and a lower (basolateral) compartment. The created polarized epithelium was shown to efficiently prevent the translocation of polystyrene-NMs between the apical and basolateral compartments.¹³⁸

Bivalves are considered to be key organisms in the fate and transport of NMs in aquatic habitats because of their ability to filter and concentrate water-suspended particles. The influence of nano-CeO₂ association to phytoplankton on the internalization of the compound by the marine mussel *Mytilus galloprovincialis* was investigated by Conway *et al.*¹²⁸ While direct exposure to 3 mg L⁻¹ CeO₂ caused a mean dry tissue cerium concentration of 33 ± 9 µg g⁻¹, a value of 28 ± 5 µg g⁻¹ was measured following exposure to the same concentration sorbed to phytoplankton. Clearance rates and pseudofeces production were shown to increase with increasing exposure concentrations. In a further study, *M. galloprovincialis* was shown to accumulate 79 ± 13 µg Cu per g dry weight following exposure to 3 mg L⁻¹ CuO NMs for four weeks. Biodeposits excreted by the animals were found to contain 110 mg Cu per g. This finding demonstrates that there is a potential for copper magnification in sediments and a possible contribution of mussels to trophic transfer of NMs to predators such as crabs and fish.¹³⁹

The results achieved so far on NM accumulation in the aquatic environment demonstrate the need to further characterize NM interactions with organic substances such as humic acids and chitin. Both are suspected to influence trophic transfer and thus potential adverse effects. Moreover, dissolution is a key parameter for NM accumulation that is likely influenced by environmental compartments.

The role of nanoparticle dissolution in their ecotoxicity

Toxicity of inorganic NMs in organisms and complex ecological systems is poorly understood despite the widespread production, use, and release of these substances into the environment.¹⁴⁰ Most profound effects are to be expected in ecosystems dominated by organisms highly sensitive to these compounds.^{140,141} Microorganisms in natural ecosystems are prime candidates because several NMs, such as nano-Ag or nano-ZnO, are deliberately designed and applied to exert antimicrobial effects. The causes of toxicity of such NMs to microorganisms are still largely unclear. Nevertheless, release of the ionic form of the metal from NMs has been suggested to play a key role.^{142,143} For instance, Navarro *et al.*¹⁴² have shown that the inhibitory effect of nano-Ag on the photosynthesis of algae was abolished when an Ag⁺ chelator was added to the culture medium. Similarly, results from a study by Xiu *et al.*¹⁴⁴ indicated a lack of negative effects on bacteria when anaerobic conditions prevented the oxidative dissolution of nano-Ag. Consequently, the importance of the released amount of ions into the test medium has aroused great interest. A study by Rohder *et al.*¹⁴⁵ showed that toxicity of nano-CeO₂ to a green freshwater alga could be attributed to Ce³⁺ ions released in the NM suspension. Furthermore,



the results of this study indicated that the measured toxicity increased with the amount of bioavailable Ce^{3+} ions in the media. In contrast, Rodea-Palomares *et al.*¹⁴⁶ suggested that toxicity of nano- CeO_2 to algal and bacterial cells was not caused by the released ions in the test media but by the attachment or sorption of the NMs to the outer surface of the microorganisms. Despite the importance of using single species microorganisms in understanding mechanisms of toxicity and ensuring comparability due to rigorous standardization, evaluation of the risks posed by soluble inorganic NMs should also consider the complex biodiversity, functions, and response dynamics in ecosystems.

Few studies have investigated the role of dissolution in the ecotoxicity of NMs at the community level.¹⁴⁷ Gil-Allué *et al.*¹⁴⁸ and Tlili *et al.*¹⁴⁹ selected two microbial communities that play key roles in fresh waters through primary production and organic matter decomposition, respectively, to provide convenient model systems to assess nano-Ag effects on complex ecological interactions and ecosystem processes. The first community of heterotrophic fungi and bacteria colonizes decomposing leaf litter; the second is an autotrophic community commonly referred to as periphyton, which consists of algae and bacteria attached to submerged surfaces. Together, these two systems capture much of the essence of stream ecosystem functioning. The studies showed that acute exposures to citrate coated nano-Ag (size 25 ± 13 nm; zeta potential -36.6 ± 3.2 mV) affected the majority of the functional endpoints tested. Most importantly, acute nano-Ag toxicity was mainly driven by dissolved Ag^+ ions; however, this was not the case for all activities measured, such as bacterial and fungal growth.¹⁴⁹ In the experiments with stream periphyton, the observed inhibition of leucine aminopeptidase, an extracellular enzyme involved in nutrient acquisition, by nano-Ag was attributed to a putative, physically-mediated particle effect independent of silver ions (Ag^+).¹⁴⁸ Indeed, addition of an Ag^+ chelator failed to prevent the inhibitory effect of nano-Ag on the measured enzyme. Consequently, the hypothesis that toxicity of nano-Ag is caused by dissolved Ag^+ could not fully explain the results obtained in short-term toxicity tests. Nano-Ag was described as exerting some particle-related effects on fungi, algae and bacteria, independent of the effect of Ag^+ , when complex microbial communities were considered. A study in zebrafish embryos and larvae on genes involved in detoxifying processes and oxidative stress responses revealed that Ag^+ and nano-Ag induced similar gene responses and affected the same target tissues, pointing to an ion-mediated effect of nano-Ag.¹⁵⁰

This notwithstanding, there is a debate whether particle-related effects, which can be the stimulation or inhibition of microbial activities, are mainly associated with changes in metal dissolution kinetics when particles are in contact with microorganisms.¹⁴⁹ Indeed, NMs near microbial cells are susceptible to being oxidized by metabolic products such as H_2O_2 .¹⁵¹ Such oxidation can lead to high local concentrations of ions, which are then rapidly taken up, and therefore cannot be measured in the test media. Firm conclusions cannot

yet be drawn on the role of ion dissolution in the toxicity of NMs to microorganisms. However, these findings highlight the importance of investigating the physicochemical state of NMs (*i.e.*, NMs *vs.* dissolved ions, primary particles *vs.* agglomerates) when ecotoxicity is assessed. They also underline the need to consider the complexity of ecosystems for studies on fate and toxicity of NMs in the environment and to focus on multiple functional endpoints for such investigations.

The impact of non-ionizing radiation on nanomaterial ecotoxicity

Ultraviolet radiation significantly influences the environmental toxicity of photoactive materials. Due to its redox ability and chemical stability, TiO_2 is most frequently used as a photocatalyst.^{152–154} The impact of different particle sizes on TiO_2 phototoxicity was compared in a study with *D. magna*. For all test materials the toxicity was significantly enhanced when organisms were simultaneously exposed to simulated sunlight irradiation in comparison to exposures under laboratory light. While intermediate-sized particles (NM-102, 20–25 nm) showed the highest potency followed by small particles (NM-101, 7–10 nm), larger particles (NM-100, 200–220 nm) were less toxic (Fig. 4).¹⁵⁵ Median effective concentrations (EC_{50}) for the three uncoated NM-types were 0.53, 1.28, and 3.88 mg L^{-1} at 48 h, respectively.

The impact of different particle sizes of TiO_2 was shown to depend on the generation of different reactive oxygen species (free and surface-adsorbed OH radicals) and the particle-*D. magna* interaction area, both influenced by particle size.¹⁵⁵ The effect of photoactivation on toxicity of NMs has been described for several additional species. In the case of the nematode *Caenorhabditis elegans*, an increased toxicity of photoactivated nano- TiO_2 associated with reactive oxygen species could be demonstrated with a median effect concentration decreasing from more than 100 mg L^{-1} to 53 mg L^{-1} .¹⁵⁶ Phototoxicity of nano- TiO_2 was further confirmed for algae.¹⁵⁷ Similar to TiO_2 , light-induced toxicity of nano- ZnO was described for algae,¹⁵⁷ nematodes,¹⁵⁸ and fish.¹⁵⁹ In a further study using zebrafish embryos and a cell line originating from the caudal fin tissue of the bluegill sunfish (*Lepomis macrochirus*), TiO_2 and Ag NMs were reported to express enhanced toxicity upon sunlight exposure, which was attributed to hydroxyl radical generation and Ag^+ -release, respectively.¹⁶⁰

While sunlight-promoted particle toxicity is assumed in the case of TiO_2 , facilitated ion release is discussed as a potential reason for the increased effects of irradiated nano- ZnO .¹⁶¹ Since the toxicity of the tested NMs was significantly enhanced when organisms were simultaneously exposed to solar irradiation in comparison to exposures under laboratory light, this study gives further evidence that standard ecotoxicity tests such as *D. magna* immobilization underestimate the environmental hazard associated with photoactive NMs. For Ag NMs with a similar coating, a lower toxicity to *D. magna* was observed at high pH, in the presence of DOM



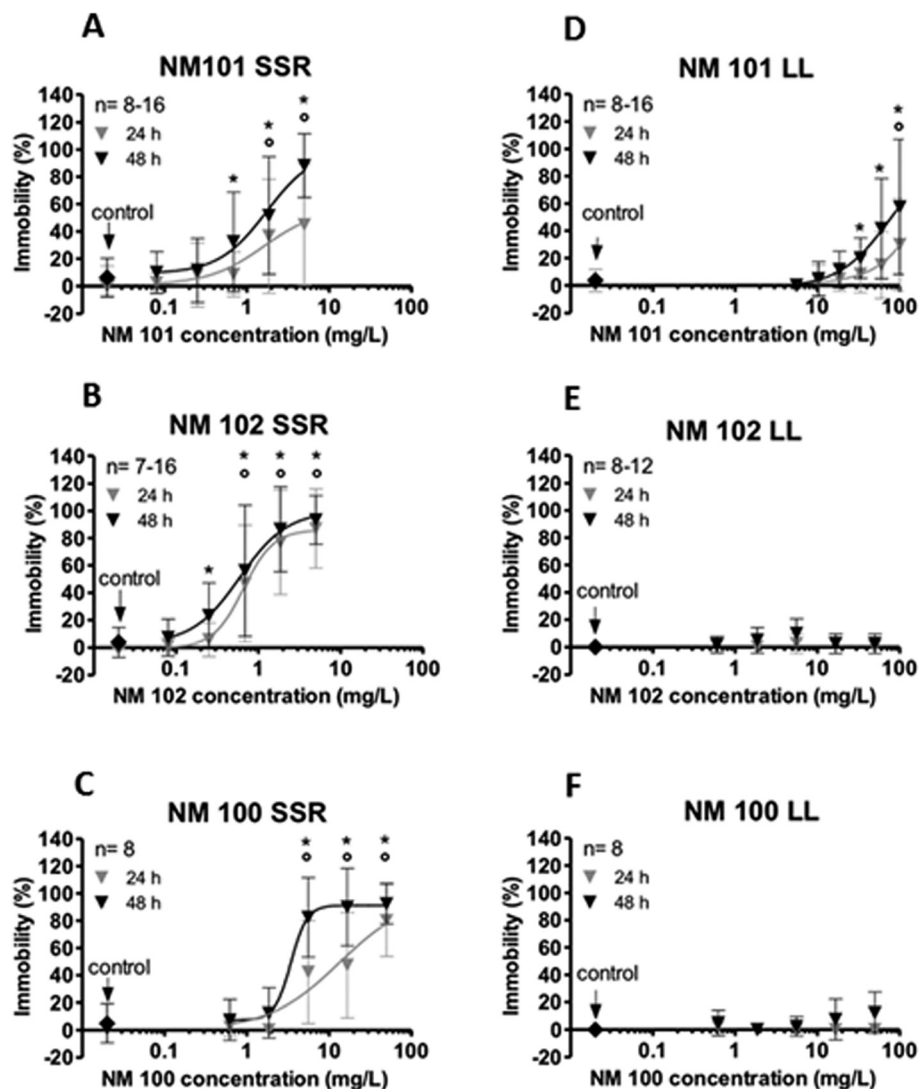


Fig. 4 Ecotoxicological investigation of two titanium dioxide (TiO_2) nanomaterials (NM-101, ϕ 7–10 nm, NM-102, ϕ 20–25 nm) and of a non-nano-sized TiO_2 material (NM-100, ϕ 200–220 nm). The immobilization of daphnids as a function of TiO_2 concentrations is shown under simulated sunlight irradiation (SSR; left; A–C) and under laboratory light (LL; right; D–F). Simulated sunlight irradiation was obtained from a mercury-vapor lamp which emitted a spectrum comparable to sunlight (280–800 nm, uva 2.36 mW cm^{-2} and uvb 0.15 mW cm^{-2}). The spectrum of this light source is given in Wyrwoll *et al.* (2016).¹⁵⁵ Laboratory light was emitted by a laboratory bulb (Osram Lumilux Cool White, HO 49W/840). Experiments were conducted in 10-fold diluted iso medium. Error bars represent standard deviations of the mean of the replicates from at least two independently conducted experiments. Circles (24 h of exposure) and asterisks (48 h of exposure) indicate significant differences from the control ($p < 0.05$). Reprinted with modifications from, A. J. Wyrwoll, P. Lautenschlager, A. Bach, B. Hellack, A. Dybowska, T. A. Kuhlbusch, H. Hollert, A. Schäffer, H. M. Maes, Size matters-the phototoxicity of TiO_2 nanomaterials, *Environ Pollut.*, **208**, 859–867, Copyright 2016, with permission from Elsevier.

and with increasing particle size, thereby supporting the major contribution of Ag species below 2 nm, including Ag^+ ions, to toxicity.¹⁶²

To date, mostly simple NM forms have been tested for phototoxicity. However, there are multiple approaches to improve the photocatalytic properties of commonly studied TiO_2 . As the molecule is only excited by UV light with a wavelength below 388 nm, sunlight absorption is limited to ca. 3%. Moreover, the electron–hole pair recombination reduces its photocatalytic activity.^{152,163,164} Multiple approaches, comprising noble metal deposition,^{165,166} semiconductor compounding,¹⁶⁷ dye sensitizing,¹⁶⁸ carbon compo-

sition,^{169,170} and doping^{171,172} were undertaken to extend the photocatalytic activity of TiO_2 from UV to visible light range.

Doping attempts comprise rare earth metals,¹⁷³ combined use of iron and nitrogen,¹⁷⁴ as well as self-doping with Ti^{3+} .¹⁷⁵ A mineralization of 4-chlorophenol and the azo dye remazol was described for carbon-doped TiO_2 , while the base material was nearly inactive.¹⁶⁴ Ag-doped TiO_2 was shown to possess a 5-fold higher efficacy against bacteriophages in aqueous media when compared to the base material.¹⁷⁶ However, the environmental impact of such procedures still requires investigation. Although Ag doping increases photocatalytic virus inactivation of TiO_2 in the first instance by



increased hydroxyl radical production, leaching of antimicrobial Ag^+ was also concluded to account for antiviral activity.¹⁷⁶ Ion-mediated toxicity is known as a main factor for the biocidal activity of Ag .¹⁷⁷ Similarly, it was shown that the antibacterial activity of Cu-doped TiO_2 is driven by Cu^{2+} release. While Fe-doping of CuO and ZnO NMs was described to have the potential of decreasing the release of toxic ions,^{178,179} the finding of rheotaxis dysfunction and DNA damage in zebrafish due to exposure to S-doped TiO_2 underpins that toxic effects due to particle surface alterations, independent from ion release, cannot be ruled out.¹⁸⁰ In addition to doping, the composition of TiO_2 and carbon has received attention for remediation purposes.¹⁷⁰ A TiO_2 (P25)-graphene photocatalyst showed a narrowed band gap and an improved adsorption of the test substance methylene blue in comparison to straight TiO_2 P25.¹⁶⁹ Later on, a 2.3-fold increased photocatalytic generation of reactive oxygen species in visible light was recorded for the graphene- TiO_2 nanocomposite compared to the base material TiO_2 P25. Together with the enhanced photoreactivity of modified nano- TiO_2 and its changed aquatic toxicity, its reactivity in the respective medium of application needs to be considered, as several environmental factors may affect agglomeration and quenching.¹⁶³ In a study by Quik *et al.*¹⁸¹ utilizing water samples from the European rivers Rhine and Meuse, homoaggregation of CeO_2 NMs was suggested to occur after sample spiking at initial high NM concentrations. Subsequent heteroaggregation or deposition of NMs on natural colloids resulting in NM removal according to first order kinetics was identified as the main mechanism leading to sedimentation.

The physicochemical properties of NMs depend on their functionalization. Coating-dependent degrees of degradation were recorded for three rutile TiO_2 nanomaterials of a primary crystallite size of 30 nm. Following 10 days incubation in ultrapure water or in the dark, there was a stronger degradation observed for particles coated with SiO_2 or SiO_2 and polydimethylsiloxane (PDMS) in comparison to those possessing a capping of SiO_2 , Al_2O_3 , and stearic acid.¹⁸² Aging of a TiO_2 -based nanocomposite used in sunscreen cosmetics, consisting of a TiO_2 core and a coating of $\text{Al}(\text{OH})_3$ and PDMS, revealed a complete desorption of PDMS after 18 h of aging in ultrapure water, while Al remained at the surface of the altered composite.¹⁸³ Coated TiO_2 NMs were shown to undergo larger changes in the light compared to dark conditions.^{182,183}

These factors need to be considered in future testing guidelines.

Outlook

Aquatic ecotoxicity assessment needs to employ standardized protocols for the reproducible and common preparation of aqueous NM dispersions. The European project NANoREG⁶⁵ recently developed harmonized and benchmarked SOPs for probe sonication-based preparation of NM dispersions for use in *in vitro* and ecotoxicity studies. However, the adoption

and widespread implementation of these standardized protocols within the scientific community and relevant regulatory authorities is a necessary next step for risk assessment. Furthermore, such standardized dispersion methods should be accompanied by a full physicochemical characterization of the resulting NM dispersion, when further diluted into specific test media and, importantly, at regular intervals throughout the duration of the selected test. This is of particular relevance for NMs that exhibit a high rate of dissolution or a tendency to agglomerate and/or settle. These properties significantly influence the actual exposure to test organisms. These parameters need to be determined to gain a more accurate assessment of potential toxicity in aquatic environments as is true for human toxicity testing.^{34,184}

The development of analytical techniques suitable for determination of NM uptake, accumulation, and transformation in complex environmental and biological matrices is increasingly necessary to gain a complete picture of NM behavior and fate, transfer through the food chain, and for interpreting toxicity data. Standardization efforts already started in projects such as VAMAS,¹⁸⁵ NanoDefine,¹⁸⁶ or ACEnano,¹⁸⁷ need to be continued and transferred into routine testing. As the toxicity of NMs may be modulated by the varying availability of ions depending on the material being tested, it is therefore important to include several toxicological endpoints to fully assess their aquatic ecotoxicity. This is in addition to the need for conducting multiple toxicity tests to inform environmental guidelines for NMs. In principal, the existing OECD test guidelines are suitable. However, some nanospecific test modifications are necessary²⁹ and were recently proposed for the growth inhibition with the green algae *Raphidocelis subcapitata* (TG 201),¹⁸⁸ acute toxicity with the crustacean *Daphnia magna* (TG 202),¹⁸⁹ development toxicity with the fish *Danio rerio* (TG 210),¹⁹⁰ and reproduction of the sediment-living worm *Lumbriculus variegatus* (TG 225).¹⁹¹

As previously described, the stability of test dispersions remains a key issue for NM aquatic toxicity testing. No additional synthetic organic solvents or dispersants should be used. If natural dispersants such as NOM are added, the lowest concentration suitable to achieve a stable distribution should be applied.¹⁹² We must consider NMs with unique physicochemical properties, such as photoactivity in the case of TiO_2 NMs. This property can have a significant impact on the aquatic toxicity of NMs. These important factors need to be considered and incorporated into current regulatory standard ecotoxicity tests conducted for use in environmental risk assessment. Newly emerging NMs with complex multicomponent compositions, increased functionalization, and surface activities will present additional challenges for characterization and environmental assessment. Furthermore, the degradation of nanocomposites and subsequent release of embedded NMs require attention as these may exert adverse effects on the environment and human health. As we understand more about the possible interactions of NMs with other organic and inorganic substances present in aquatic



matrices, there is an additional need for analytical tools to support studies in this field.

Abbreviations

CNT	Carbon nanotube
MWCNT	Multi-walled carbon nanotube
NM	Nanomaterial
NOM	Natural organic matter
PAA	Polyacrylate-coated
PDMS	Polydimethylsiloxane
SEM	Scanning electron microscopy
SSA	Specific surface area
SOP	Standard operating procedure
SWCNT	Single-walled carbon nanotube
TEM	Transmission electron microscopy
UV/vis	Ultraviolet-visible
P25	Aeroxide TiO ₂

Conflicts of interest

There are no conflicts of interest to declare.

Acknowledgements

The authors acknowledge support from the EU FP7 project 'NANOREG' (Grant Agreement number 310584). The authors wish to thank this project for financial support of their research and for periodic teleconferences and meetings. Acknowledgements (A. M. Booth): Supported by Polish-Norwegian Research Programme (Project Contract No. Pol-Nor/237761/98/2014) and the Research Council of Norway (Contract No. 239199/O70). Acknowledgements (C. Cerrillo): Gemma Mendoza, Olatz Areitioaurtena, Amaya Igartua, Gotzone Barandika, and the ZABALDUZ Program for financing her contract with UPV/EHU in collaboration with IK4-TEKNIKER. Acknowledgements (A. Tlili): Participants in this research include Renata Behra, Julien Cornut, Carmen Gil-Allué, Mark. O. Gessner and Jeremy Jabiol. The project was supported by the Swiss National Science Foundation (SNF, grant no. 200020_134750/1). Acknowledgements (A. Schäffer): Participants in this research included Hanna Maes, Anne Wyrwoll, Alexander Bach, Henner Hollert, Maximilian Gruber, Petra Lautenschläger, Anne Meister-Werner and Ralph Petto.

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