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# Safe-by-design assessment of an SiO<sub>2</sub>@ZnO multi-component nanomaterial used in construction†

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Safety aspects of chemicals/materials are transversal in all sustainability dimensions, representing a pillar at the early innovation stages of the European Commission's "safe and sustainable by design" (SSbD) framework for chemicals and materials. The first three of the five SSbD framework steps cover different safety aspects, namely, hazard assessment based on intrinsic properties (step 1), occupational health and safety (including exposure) assessment during the production/processing phase (step 2) and exposure in the final application phase (step 3). The goal of this work was to identify a set of characterization tools/procedures to support the operationalization of the first three safety steps in multi-component nanomaterials (MCNMs), applying the findings to an SiO<sub>2</sub> core–ZnO shell MCNM. The safety of this MCNM, which is used as an additive to silicate/calcium hydroxide mortar to improve air quality through photocatalytic NO<sub>x</sub> removal, was investigated from different perspectives along its value chain. Existing and newly generated data on its hazard profile were collected, the exposure of workers during its synthesis was assessed, and potential exposure to hazardous substances during its final application phase was investigated. In step 1, physico-chemical properties, hazard classification and cytotoxicity assays were considered. In step 2, a three-tiered established methodology for evaluating occupational exposure assessment was performed. Lastly, in step 3, the release of inorganic substances from MCNM-based mortars in the final application phase was investigated. Safety assessment according to the SSbD framework was performed by selecting tools and procedures suitable for application in the early innovation stage, resulting in a preliminary hazard assessment of MCNMs and a suggestion for redesigning a step in the process.

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## Environmental significance

Multicomponent nanomaterials (MCNMs) integrate several functions into one material, leading to innovative applications. SiO<sub>2</sub>@ZnO MCNMs incorporated in mortars increase the mechanical strength of the material while providing photocatalytic activity for NO<sub>x</sub> removal. However, the combination of these multiple features may also lead to unexplored risks, making it necessary to carefully evaluate their human health and environmental aspects before entering the market. In this context, a safety assessment of SiO<sub>2</sub>@ZnO MCNMs using setting tools/procedures to operationalize the safety aspects of the safe and sustainable by design (SSbD) framework is needed for the creation of safer and functional materials. The main goal of this approach, in the early phase of the design process, is to decrease the likelihood of adverse impacts on human health and the environment.

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# 1. Introduction

A wide array of studies has identified that the hazards of engineered nanomaterials (NMs) varies considerably according to their physicochemical characteristics.<sup>1–3</sup> This information has been useful in their risk assessment, regulatory considerations and the development of safe by design (SbD) strategies. However, science and technology are progressing at an increasing speed, developing new unconventional materials with the aim to address key global challenges such as energy provision, climate change, reducing pollution and improving human health. With regard to EU goals defined under the European Green Deal policy, prioritized key enabling technologies include advanced materials, such as multi-component nanomaterials (MCNMs), which consist of at least two components with a size in the nanometer range. MCNMs are widely used in several fields, including targeted drug delivery,<sup>4</sup> multicomponent medical sensing,<sup>5</sup> heterogeneous catalysis<sup>6</sup> and multifunctional coatings in construction. The latter enables enhanced exterior durability and mechanical strength<sup>7</sup> as well as photocatalytic activity using TiO<sub>2</sub>–ZnO MCNMs for NO<sub>x</sub> removal.<sup>8</sup> However, the benefits from their use can be marred by uncertainties about their potential risks to human health and the environment, given that it is difficult to identify and assess these materials owing to their multicomponent nature.<sup>9,10</sup> Indeed, upon contact with biological or environmental systems, the components of MCNMs could exhibit differing rates of degradation or interaction to induce combined effects (*e.g.* antagonistic, synergetic and potentiation effects).

Thus, to guide the innovation process for chemicals and materials, including MCNMs, the Joint Research Centre (JRC)<sup>11</sup> proposed a safe and sustainable by design (SSbD) framework, which forms the basis of the recommendation by the European Commission (EC).<sup>12</sup> This framework was developed by considering SbD methods and frameworks to diagnose potential risks from the use of NMs in different commercial products.<sup>13,14</sup> Recently, Sudheshwar *et al.*<sup>15</sup> reviewed the existing studies prior to the SSbD framework, showing that the SbD concept is originally linked to the nanotechnology sector. They also noted that the 89 SbD studies examined predominantly addressed human safety aspects over environmental issues, with only 14 of them being relevant to the SSbD framework. These studies were compared based on aspects such as the tools used/developed, the applicability domain, suggested guidance, life cycle stages addressed, case study presence/absence, and link to the SSbD framework. With respect to previous SbD frameworks, the main novelty of the SSbD by the EC is the integration of safety and sustainability aspects of chemicals and materials with a desirable function (or service) as early as possible in the innovation process. This premarket approach considers safety, environmental, social and economic aspects along the entire life cycle of a chemical/material and consists of 5 steps, as follows: i) hazard assessment; ii) human health and

safety aspects in the production and processing phase; iii) human health and environmental aspects in the final application phase; iv) environmental sustainability assessment; and v) social and economic sustainability assessment. The framework proposes the integral assessment of all steps in an iterative manner as the innovation proceeds. This means that all steps should be addressed together in the different innovation stages, which leads to an iterative approach of the framework as the innovation proceeds and more data become available. In this work, we focused on safety as a transversal aspect of sustainability, and specifically steps 1–3 of the SSbD framework. In the case of MCNM, this is already challenging and not straightforward and requires specific attention to the physical, chemical and toxicological characterization procedures to be used and the exposure assessments to be performed for MCNMs.

Given the current lack of knowledge on the safety aspects of nano-enabled products and the need for effective characterization procedures and suitable exposure assessments for MCNMs according to the SSbD approach, it is crucial to identify physical, chemical and toxicological testing strategies for the safety assessment of new materials throughout the design phase before they enter the market. These strategies must be able to carefully evaluate their hazard profile in line with the policy initiatives launched by the European Green Deal and regulatory requirements.

The case study used is an SiO<sub>2</sub> core–ZnO-shell MCNM embedded in a cement mortar for the photocatalytic decontamination of NO<sub>x</sub> in air. In this study, according to the safety pillar of the SSbD framework, a safety assessment of the MCNM was performed in the very early stages of the design process. This work was performed in the frame of the European H2020 SUNSHINE project (<https://h2020sunshine.eu>), which strives to develop an overarching approach for the SSbD of MCNMs, including demonstration *via* industrially relevant case studies.<sup>16</sup>

## 2. Materials and methods

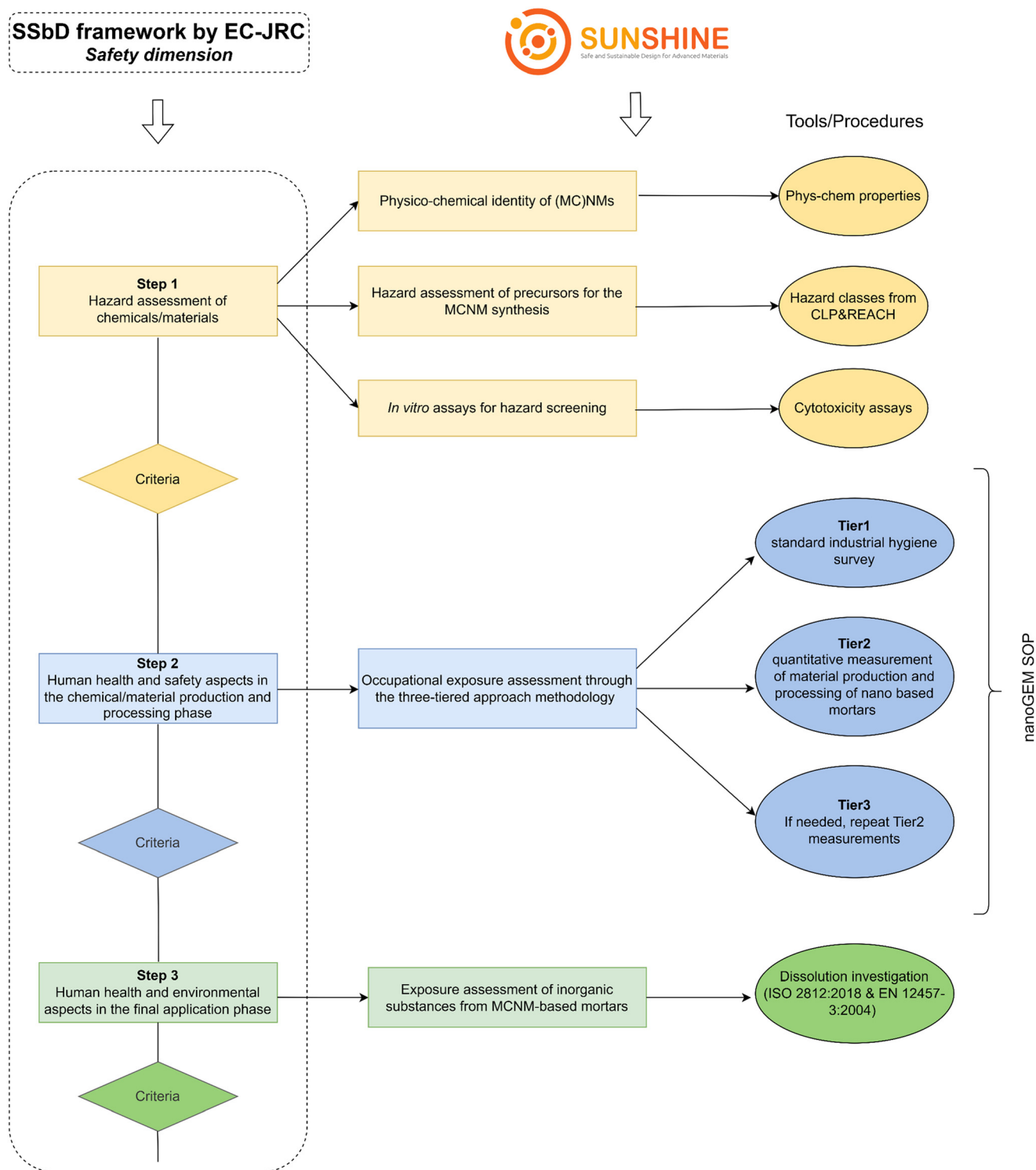
### 2.1 SUNSHINE implementation of the safety dimension within the SSbD framework

The first three steps of the SSbD framework cover different aspects of chemicals/materials safety, for which indicators, criteria and an evaluation system need to be defined. In detail, step 1 aims at investigating the intrinsic properties of a chemical/material to determine its hazard profile before further assessing its safety during use. This is based, when available, on hazard classes and categories established within the Classification, Labelling and Packaging (CLP) Regulation 1227/2008 as well as Registration, Evaluation, Authorisation and Restriction of Chemicals (REACH) Regulation 1907/2006. Step 2 deals with the human health and safety aspects in the chemical/material production and processing phases, investigating whether the chemical/material of interest can pose any risk to workers before application. This step involves the use of available models to assign scores/levels to



the related variables including hazard of chemicals, frequency and duration of exposure, amount of chemical used/present, physico-chemical properties, operational conditions, and type of existing risk management measures. Based on the results, the risk estimation is performed. Lastly,

step 3 covers the human health and environmental aspects in the final application phase, identifying the likelihood of the exposure and the potential exposure routes and related toxicity impacts. Within SUNSHINE, the safety aspects for the advanced  $\text{SiO}_2\text{@ZnO}$  MCNM were covered, as summarized



**Fig. 1** Safety dimension assessment of the SSbD framework (in dashed rectangle) and corresponding implementation within SUNSHINE. Tools/procedures for each step are displayed in ellipses.



below. With regards to step 1 of the SSbD framework, physico-chemical characterization of both individual NMs and MCNM, hazard assessment of the MCNM precursors and preliminary *in vitro* assays with NMs and MCNM were performed (*cf.* 3.2). In step 2, an occupational exposure assessment was carried out through an air monitoring campaign (*cf.* 3.3). Concerning step 3, the potential leaching of inorganic substances from both the pristine powders and the MCNM-based mortars was assessed by dissolution experiments (*cf.* 3.4). Fig. 1 shows the relation between the safety dimension assessment of the SSbD framework and the corresponding implementation within the SUNSHINE project.

## 2.2 Case study materials

The pristine  $\text{SiO}_2\text{@ZnO}$  MCNMs were synthesized by the CIAC Company (Cordoba, Spain), starting from commercial mesoporous  $\text{SiO}_2$  NM powder with an average particle diameter of 20 nm (99.5% purity, NanoAmor, TX, USA) and  $\text{ZnAc}_2\cdot 2\text{H}_2\text{O}$  (purity 99.5%, ITW Reagents S.r.l.). In brief, zinc acetate dihydrate was deposited on the  $\text{SiO}_2$  NM core in a 1:2 ratio through a calcination process at 600 °C (heating rate of 100 °C h<sup>-1</sup>) for 2 h, and then hand ground with agate mortar for 15 min, to form the desired core-shell structure with ZnO NM as the outer shell. In addition, a ZnO NM was also synthesized as a comparable individual component (benchmark) material by following the same calcination procedure at 600 °C from zinc acetate dihydrate in the absence of  $\text{SiO}_2$ .

Concerning the MCNM-enabled product, the mortar used for testing was Morcemsec® Capa Fina CR CSIV W2 (Grupo PUMA S.L., Cordoba, Spain) based on Portland cement (between 9 and <40 wt%) and calcium dihydroxide (<5 wt%). The formulation of the MCNM-based mortars was performed by adding the  $\text{SiO}_2\text{@ZnO}$  MCNM powder to commercial mortar, obtaining homogeneous dust, and then mixing it with water, following the procedure described in EN 196-1:2016 (*ref.* 17) before placing in a mold for 48 h. Afterwards, according to the curing conditions of EN 1015-11:2019,<sup>18</sup> the samples were exposed to 95 ± 5% relative humidity and 20 ± 2 °C for 5 days and 65 ± 5% relative humidity and 20 ± 2 °C for an additional 21 days.

## 2.3 Hazard assessment of the chemicals/materials (step 1)

**2.3.1 Physicochemical characterization.** The physicochemical identity of the two individual NMs and the MCNM (henceforth called (MC)NMs) was performed by selecting the minimum set of both chemical and physical properties according to expert judgment and the available literature.<sup>19</sup> The properties examined were constituent particle size,<sup>20</sup> morphology, and chemical composition by transmission electron microscopy coupled with energy dispersive X-ray (TEM-EDX and STEM), crystalline structure by X-ray diffraction (XRD) and elemental composition by inductively coupled plasma optical emission spectroscopy

(ICP-OES). TEM characterization was carried out in a JEOL Model JEM-1200 EXII transmission electron microscope equipped with an electron emission source operated at 200 kV and STEM characterization was carried out using a JEOL 2100F TEM/STEM equipped with an electron emission source at 200 kV. EDX (X Max 80, Oxford Instruments). For the TEM and STEM observations, the samples were dispersed in ethanol and sonicated for 5 min at 80 W using a bath sonication (Elma S30; Elmasonic). Then, a few drops were placed on a holey carbon-coated copper grid, allowing the solvent to evaporate in air before TEM/STEM observation. EDX was used to collect information on the elemental identity of the MCNMs.

The XRD patterns were recorded using an X'Pert PRO diffractometer by PANalytical and data were collected in the 2θ range of 5° to 55° at a time step of 50 s. The quantification of the elemental composition was performed using an ICP-OES 5100 – Vertical Dual View apparatus coupled with a OneNeb nebulizer (Agilent Technologies, Santa Clara, CA, USA), using a mixture of 10% v/v  $\text{HNO}_3$ , 10% v/v  $\text{H}_2\text{SO}_4$  and 1% v/v HF (ultrapure analytical grade from Merck, Darmstadt, Germany) to ensure complete digestion. Calibration curves were obtained with 0.1, 1, 10 and 100 mg L<sup>-1</sup> of Si and Zn standards prepared in ultrapure water using the same procedure applied to the samples.

The reactivity of both individual and MCNMs and the degree of coverage of  $\text{SiO}_2$  by ZnO was assessed *via* temperature programmed reaction (TPRx) using methanol as the probe molecule. The potential reaction products include carbon dioxide ( $\text{CO}_2$ ) for materials with basic sites, methyl formate ( $\text{HCOOCH}_3$ ) for materials possessing bifunctional basic-redox sites, formaldehyde ( $\text{HCHO}$ ) for materials with redox sites, dimethoxymethane ( $(\text{CH}_3\text{O})_2\text{CH}_2$ ) for materials featuring bifunctional acid-redox sites, and dimethyl ether ( $\text{CH}_3\text{OCH}_3$ ) for materials with acidic sites. The reaction products were detected using a Pfeiffer OmniStar mass spectrometer. The experimental procedure involved four phases including pretreatment, initial purge, methanol chemisorption, and TPRx. Pretreatment involved heating the sample from 100 °C to 450 °C at a rate of 10 °C min<sup>-1</sup> with the introduction of 150 mL min<sup>-1</sup> of synthetic air to eliminate surface moisture and impurities. Following pretreatment, an initial purge with inert gas (150 mL min<sup>-1</sup> of Ar) at 100 °C was conducted until impurities were no longer detected in the mass spectra. Subsequently, methanol chemisorption was carried out with a methanol/argon mixture at a constant temperature of 100 °C, concluding when a stable methanol signal was observed in the mass spectrum ( $m/z = 31$ ). The final phase, TPRx, involved maintaining the methanol/argon flow, while increasing the temperature from 100 °C to 450 °C at a rate of 10 °C min<sup>-1</sup>. The experiment utilized a sample of 0.1 g (mesh size: 25–100 μm) diluted with 0.5 g of SiC (black 180, Navarro SiC S. A.) to ensure a uniform temperature distribution.





**2.3.2 Hazard assessment of the MCNM precursors.** The information gathered from the physico-chemical characterization of (MC)NMs was used as the basis to assess their potential hazard. The safety concept of the SSbD approach aims to avoid hazardous chemicals/materials or decrease their use to avoid and minimize risks to humans and the environment.<sup>11</sup> Accordingly, the first hazard evaluation of the chemicals used for the synthesis of the SiO<sub>2</sub>@ZnO MCNM was performed based on CLP and REACH regulations.<sup>21,22</sup>

**2.3.3 *In vitro* toxicity testing.** Due to the lack of hazard information, an assessment of the precursors, NM and MCNM was performed using *in vitro* (non-animal) methods. A common mechanism of toxicity induced by particles in the lung is inflammation (activation of the immune cells such as macrophages), which can lead to health effects such as lung fibrosis (scarring preventing breathing) and tumors. Therefore, the measurement of the potential for NMs or MCNMs to induce inflammation was used as an indicator of toxicity, which has been included in many hazard assessment strategies.<sup>23,24</sup> The *in vitro* indicators used to assess the potential to induce inflammation include the production by cells of proteins known as cytokines. In particular, the release of the cytokine interleukin 1-beta (IL-1β) indicates the activation of mechanisms in macrophages considered to play an important role in NM-induced toxic responses.<sup>25</sup>

In this study, a frequently used cell line was employed (THP-1 monocyte cells; ATCC (TIB 202), passage number 15–20), which was treated to differentiate the cells to behave like mature macrophages. A standard operating procedure (SOP) was utilised to culture the cells and treat them with controls, NMs or MCNMs to enhance the reproducibility of the results and comparison with previous studies using the same SOP (<https://www.patrols-h2020.eu/publications/sops/index.php>).<sup>26</sup>

SiO<sub>2</sub>@ZnO MCNM, SiO<sub>2</sub> NM or ZnO NM were suspended at a concentration of 1 mg mL<sup>-1</sup> in phosphate buffered saline with 0.05% w/v bovine serum albumin (BSA) and dispersed by sonication using an ultrasonication water bath for 16 min. Dispersion was followed immediately by dilution in RPMI medium (without serum) to obtain the required final concentrations (final concentration 0–100 μg mL<sup>-1</sup>). The treatments were added to each well of a 96-well plate containing cells prior to incubation at 37 °C in the presence of 5% CO<sub>2</sub> for 24 h.

The deposited dose was not assessed as this was not deemed appropriate for hazard screening at such an early innovation stage given that it is important to keep the experimental work manageable and affordable.

Cytotoxicity was assessed as a decrease in cell viability to indicate short term hazard, but more importantly it allowed the selection of appropriate concentrations to investigate the production of IL-1β at sublethal concentrations.<sup>23</sup> Cell viability was assessed using the Alamar blue (ThermoFisher, catalog number: DAL1025) assay, which measures the ability of cells to reduce the non-fluorescent dye 7-hydroxy-3H-phenoxazin-3-one 10-oxide (resazurin) to the fluorescent product resorufin. A decrease in the ability to generate the fluorescent product (excitation/emission wavelengths of 560/590 nm) was used as the measure of cytotoxicity. Data were expressed as % cell viability. The control cells were treated with RPMI medium (without FCS) without particles added. The production of IL-1β was measured in the cell culture supernatant, according to the R&D Human IL-1β/IL-1F2 DuoSet protocol (catalog numbers: DY201-05).

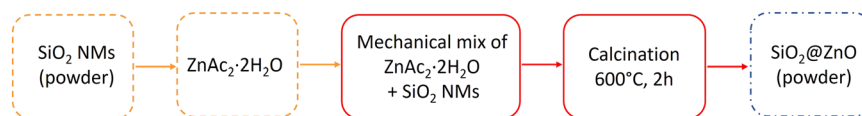
**2.3.4 Statistical analysis.** For the statistical analysis, each well of the plate was considered an individual experimental unit. Three replicates were conducted per experiment, and the experiment was repeated 4–5 times independently, resulting in 12–15 measurements in total. The fluorescence of each well was expressed as a percentage of the mean untreated control on the same day of experimentation. The mean and standard error of the mean of the percentage of control values from all experiments for each exposure material and dose were calculated and plotted graphically.

## 2.4 Human health and safety aspects in the chemical/material production and processing phase (step 2)

The NMs/chemicals used and the steps of the process to synthesize the SiO<sub>2</sub>@ZnO MCNM are displayed in Scheme 1. Regarding the evaluation of the occupational safety and health (OSH) aspects in the production and processing of (MC)NMs, a three-tiered methodology, already recommended by different authors or institutions,<sup>27–31</sup> was applied, as depicted in Fig. 2.

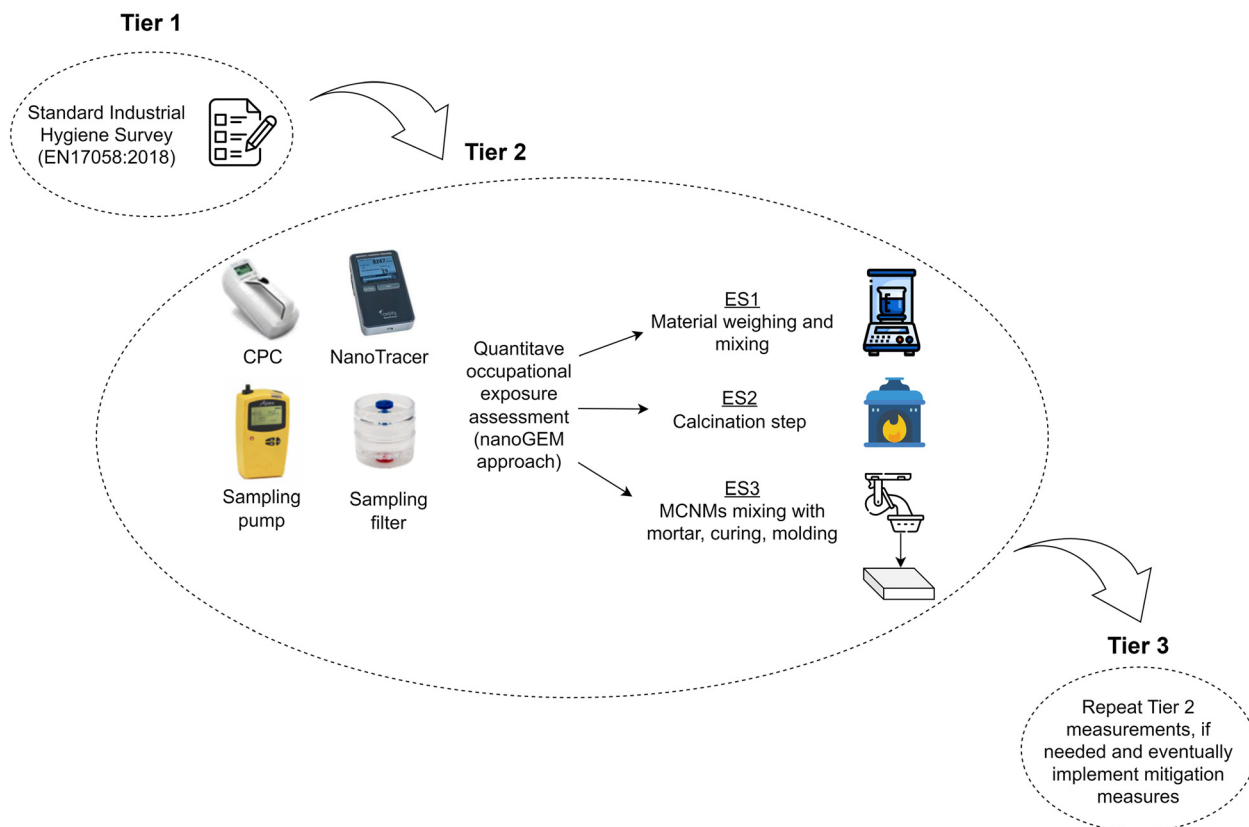
In the first tier, a standard industrial hygiene survey through questionnaires following the EN 17058:2018 (ref. 32) “workplace exposure – assessment of exposure by inhalation of nano-objects and their aggregates and agglomerates” was filled in by CIAC Foundation and included in the ESI† (Tables S1 and S2), for gathering information on potential sources/points of (MC)NMs emission. The target areas, processes or tasks from which any (MC)NMs can be released were identified.

Afterwards, tier two covered the occupational exposure assessment at different stages of the material production



**Scheme 1** Steps of the process to synthesize SiO<sub>2</sub>@ZnO MCNM as powder. NMs/chemicals used are displayed in rectangles with dashed lines, processes with solid lines and final product with dashed-solid lines.





**Fig. 2** Schematic of the three-tiered methodology approach applied to assess the occupational safety and health (OSH) aspects in the production and processing of (MC)NMs. ES = exposure scenario.

and processing of mortars with embedded  $\text{SiO}_2\text{@ZnO}$  MCNM, performing a quantitative exposure assessment. An overview of the exposure scenarios (ES) and the contributing exposure scenarios (CES), which refer to specific activities where the release of (MC)NMs may take place, were identified and listed in Table 1, according to ECHA guidance documents.<sup>33,34</sup> The maps of each ES are displayed in Fig. S1–S3.†

In particular, the potential exposure to (MC)NMs was evaluated over around 50 min during both their synthesis (*i.e.*, material weighing and calcination) and the (MC)NM-based mortar formulation. The particle number and mass concentrations both in air and in the personal breathing zone (*i.e.*, PBZ, around 30 cm from the respiratory tract) of the workers was determined by using a portable condensation particle counter (CPC, model TSI 3007) and NanoTracer PNT 1000. To gain insights into the possible transport of particles through the air, measurements were performed at around 0.5–1 m (near field, NF) and 5–7 m from the source (far field, FF). Regarding the chemical composition and morphology of the particles potentially released during the monitoring campaign, a Tygon tube (length 1 m) at 30 cm from the mouth of the worker was set to monitor particle release near the breathing zone. Moreover, two high-flow peristaltic pumps (Casella, model APEX) containing a polycarbonate HEPA filter with a

diameter of 37 mm and pore size of 0.4  $\mu\text{m}$ , were fixed on the lab coat of the worker at 30 cm from their mouth to collect the particles in air during the monitoring campaign. Subsequently, the filters were observed by scanning

**Table 1** Exposure scenarios (ES), related activities and (MC)NMs investigated during the occupational monitoring campaign

Exposure scenario (ES)	Contributing exposure scenario (CES)	(MC)NMs
ES1 (material synthesis)	1. Background measurement 2. $\text{ZnAc}_2 \cdot 2\text{H}_2\text{O}$ weighing 3. $\text{SiO}_2$ weighing 4. $\text{SiO}_2$ addition 5. Local exhaust ventilation 6. 1st $\text{ZnAc}_2 \cdot 2\text{H}_2\text{O}$ addition 6. 2nd $\text{ZnAc}_2 \cdot 2\text{H}_2\text{O}$ addition	$\text{SiO}_2$
ES2_I (calcination)	7. Background measurement 8. Moving the dispersion 9. Muffle opening	$\text{SiO}_2$ , ZnO
ES2_II (calcination)	10. High temperature ramp	$\text{SiO}_2$ , ZnO
ES3 (mortar formulation)	11. Background measurement 12. Low temperature ramp 13. Mortar addition to the stirring tank 14. Additive addition to the stirring tank 15. Additive addition to the stirring tank + stirring 16. Test specimen unmolding	$\text{SiO}_2$ , ZnO



electronic microscopy analysis using a field emission scanning electron microscope (FESEM, Carl Zeiss Sigma NTS, Germany). Elemental analysis was performed by image analysis using FESEM coupled to an energy dispersive X-ray micro-analyser (EDS, mod. INCA).

Finally, according to the standard operating procedure published as part of the three-tiered methodology,<sup>27</sup> the criterion used to evaluate the results from tier 2 was the comparison of the particle concentration values obtained during the different activities monitored and the particle background concentrations using eqn (1), as follows:

$$C_{\text{net}} - C_{\text{bg}} > 3 \cdot S^2(\text{DBI}) \quad (1)$$

where  $C_{\text{net}}$  is the particle emission/exposure concentration,  $C_{\text{bg}}$  is the particle background concentration and  $S^2(\text{DBI})$  is the standard deviation of the particle background concentration. If  $C_{\text{net}} - C_{\text{bg}}$  is greater than  $3 \cdot S^2(\text{DBI})$  (*i.e.*, the standard deviation of the background concentration), it means that the recorded particle concentration is statistically significant with respect to the background values. Therefore, the workplace or process concentration must be further assessed for the release of airborne nano-objects in tier 3, making the implementation of mitigation measures necessary. Tier 3 requires repeating tier 2 measurements together with the simultaneous collection of particles for the off-line analysis of mass or fiber concentration, particle morphology and chemical composition. More details on this approach are reported in the ESI.†

## 2.5 Human health and environmental aspects in the final application phase (step 3)

**Dissolution investigation.** In this stage, the exposure assessment to hazardous substances from the mortars with embedded MCNMs was evaluated, covering the safety dimension within the application/use phase.

Firstly, a preliminary release study of Zn and Si from both individual and MCNM pristine powders was conducted. In detail, stock dispersions of (MC)NMs (*i.e.*,  $\text{SiO}_2$ , ZnO and  $\text{SiO}_2@\text{ZnO}$ ) at  $0.5 \text{ g L}^{-1}$  (20 mg in 40 mL ultrapure water, UPW) were sonicated with an ultrasound probe (UP-200S Hielscher Ultrasonics GmbH, Germany) in an ice bath for 4 min (power = 80%, frequency = 0.5 and sonication energy =  $43 \text{ J mL}^{-1}$  (ref. 35)). Afterwards, ultrafiltration of each sample was performed using Amicon Ultra 15 mL centrifugal 3k filters (Millipore) at 0.5, 1, 3, 24 and 48 h after sample preparation and acidified with 2%  $\text{HNO}_3$  prior to the ICP-OES analysis. The calibration curves and limits of detection (LOD) are reported in the ESI.†

Subsequently, the potential release of hazardous substances from the MCNM-based mortars was studied. In brief, the dissolution of Zn, Si and Ti was estimated from cement mortars with different percentages of MCNMs (*i.e.*, 1% or 5% of  $\text{SiO}_2@\text{ZnO}$ , and as references, 1% or 5% of  $\text{TiO}_2$  and 1% or 5% of silica-fume), according to the ISO

2812:2007 on Paints and varnishes – determination of resistance to liquids-Part 2: Water immersion method<sup>36</sup> and to EN 12457-3:2004 for leaching evaluation.<sup>37</sup> Before the release experiment, each mortar tested (weight:  $29 \pm 0.7 \text{ g}$ , size:  $4 \times 4 \text{ cm}$ ) was subjected to a thorough cleaning process with compressed air to get rid of potential impurities on its surface and weighed. After this pre-treatment, each mortar was totally immersed in 200 mL of UPW, placed on an upside-down glass petri dish to avoid contact with the bottom of the beaker and magnetically stirred during the test. Leachates were collected at different intervals (*i.e.*, 0.5, 3, 24, 48 and 72 h, which correspond to the typical time points used in (eco)toxicity assays) and filtered through a  $0.2 \mu\text{m}$  mesh filter. Afterwards, each sample was acidified with 2%  $\text{HNO}_3$ , and then analysed by ICP-OES to determine the concentration of Zn, Si and Ti.

## 3. Results and discussion

### 3.1 Physicochemical characterization

Among the intrinsic properties that can influence the toxicity and modulate the intensity of adverse effects exerted by the (MC)NMs,<sup>38</sup> chemical composition, constituent particle size, morphology, and crystallinity were considered as the most relevant and were investigated in detail. Regarding the chemical composition, the ICP-OES analysis estimated that  $\text{SiO}_2$  accounted for 80% of the total MCNM, while ZnO for the remaining 20%. The particle size and shape of the (MC)NMs were determined by TEM (Fig. 3), showing highly agglomerated constituent  $\text{SiO}_2$  particles, with an average particle size of around 20 nm (Fig. 3a). ZnO NM (Fig. 3b), synthesized as benchmark material, showed elliptic/roundish shape particles with an average particle size of  $<200 \text{ nm}$ . Fig. 3c displays the TEM image of MCNM, which is constituted of agglomerated particles similar to that observed for  $\text{SiO}_2$  NM but with the presence of a non-homogeneous coating.

The crystalline structure of MCNM was determined *via* XRD analysis, showing that the  $\text{SiO}_2$  core was amorphous, whereas the ZnO shell revealed a crystalline phase, as observed for the individual components (Fig. S4†).

To gather further information on the coating, energy dispersive X-ray (EDX) mapping, reactivity analysis *via* MeOH-TPRx and scanning transmission electron microscopy (STEM) analysis were also performed. The EDX mapping of the single elements, *i.e.*, Si (Fig. S5c†), Zn (Fig. S5d†), O (Fig. S5f†), and the overlayed mapping of Si–Zn (Fig. S5e†) confirmed that MCNM was an Si core–Zn shell structure with a non-homogeneous coverage. Finally, the reactivity analysis *via* MeOH-TPRx provided insight into the primary reactivity of the surface reactive sites of (MC)NMs.  $\text{SiO}_2$  (Fig. S6a†) exhibited acidic reactivity, as indicated by the formation of dimethyl ether (DME, green trace). ZnO (Fig. S6b†) displayed redox character, as evidenced by the formation of formaldehyde. Lastly, the presence of ZnO on the surface of  $\text{SiO}_2$  resulted in a system that has both acidic and redox



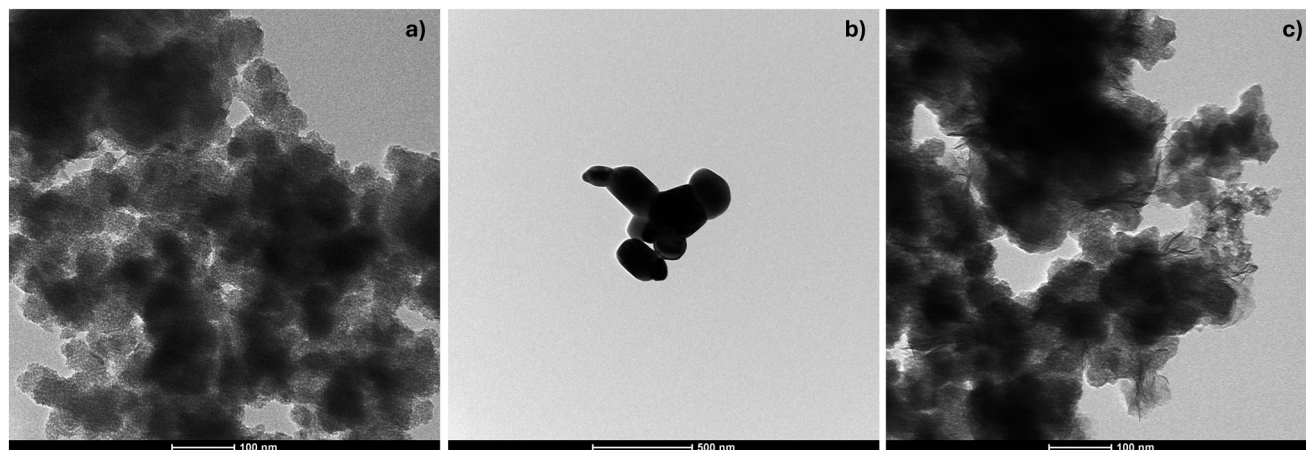


Fig. 3 TEM images of a)  $\text{SiO}_2$  NM, b)  $\text{ZnO}$  NM, and c)  $\text{SiO}_2@\text{ZnO}$  MCNM.

properties (DME and HCHO formation, respectively) (Fig. S6c†). This reactive profile indicated that silica (acidic) is not totally covered by ZnO. Therefore,  $\text{SiO}_2@\text{ZnO}$  exhibited reactive properties similar to that of its individual components ZnO and  $\text{SiO}_2$ , without any distinct effects arising from the interaction between  $\text{SiO}_2$  and ZnO being clearly observed. Finally, the non-homogeneous coverage of  $\text{SiO}_2$  by ZnO was also corroborated by STEM (Fig. S7†).

### 3.2 Hazard properties evaluation

According to the SSbD framework, the hazard assessment of chemicals/materials to synthesize the  $\text{SiO}_2@\text{ZnO}$  MCNM was performed considering the information gathered from the physico-chemical characterization and the criteria defined in Table 3 of the SSbD framework<sup>11</sup> (*i.e.*, criterion H1 = most harmful substances, criterion H2 = substances of concern,

and criterion H3 = other hazard properties) based on the hazard classes and categories determined through CLP and REACH.

The  $\text{SiO}_2$  NM powder (CAS no. 7631-86-9), which was used to improve the stability, compatibility, and dispersibility of the cementitious material with respect to its bulk phase,<sup>39</sup> was classified as non-hazardous according to the current Globally Harmonized System of Classification and Labelling of Chemicals (GHS). However, given that there is still no final consensus on the risk assessment of NMs,<sup>40,41</sup> this classification may be updated by the European Chemicals Agency (ECHA) if new insights on their toxicity are provided.

Alternatively, the hazard identification of  $\text{ZnAc}_2 \cdot 2\text{H}_2\text{O}$  (CAS no. 557-34-6) showed both health and environmental hazards, *i.e.*, acute toxicity (oral, category 4, H302), serious eye damage (category 1, H318) and long-term (chronic) aquatic hazard (category 2, H411). The workflow to assess the

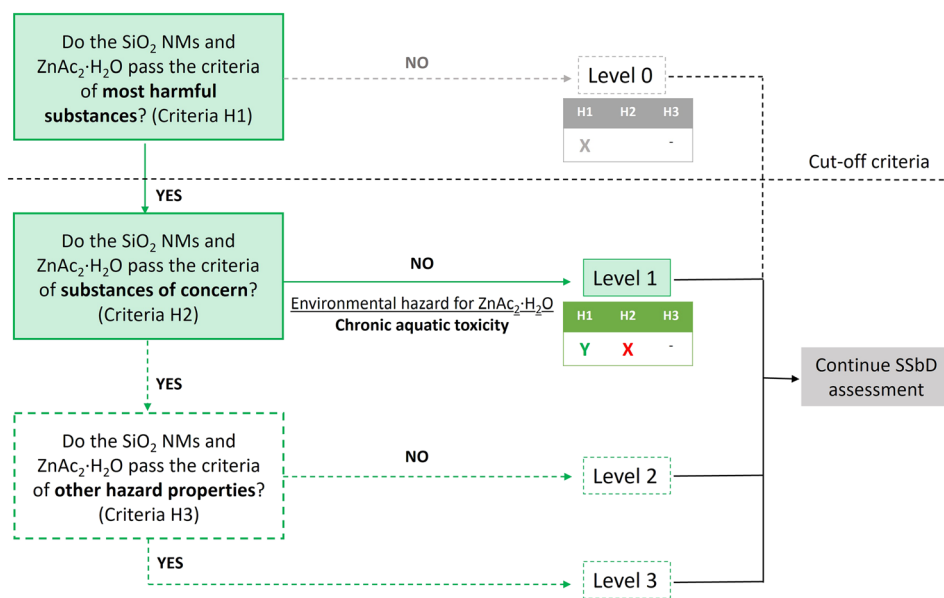


Fig. 4 Workflow to assess hazard of  $\text{SiO}_2@\text{ZnO}$  MCNM based on step 1 of the SSbD framework. The pathway followed is represented by solid lines and colored rectangles.





hazard of the chemicals used for the synthesis of  $\text{SiO}_2@\text{ZnO}$  MCNM is displayed in Fig. 4.

Therefore, based on the three criteria defined in Table 3 of the SSbD framework,<sup>11</sup>  $\text{SiO}_2$  NMs and  $\text{ZnAc}_2\cdot 2\text{H}_2\text{O}$  pass the H1 but not H2 criteria, which refers to substances of concern described in the Chemical Strategy for Sustainability (CSS) and not already included in the H1 criteria. In addition to these criteria, the framework suggests the use of an evaluation system, which includes 4 levels for step 1, moving from level 0 (chemicals or materials that do not pass hazard criterion H1 (e.g., considered most harmful substances) to level 3 (chemicals or materials that pass all the safety criteria in step 1). Accordingly, the NMs and chemicals used for the synthesis of  $\text{SiO}_2@\text{ZnO}$  MCNM belong to level 1, corresponding to chemicals or materials that pass hazard criterion H1 but not criterion H2 (i.e.,  $\text{ZnAc}_2\cdot 2\text{H}_2\text{O}$  has the potential to induce chronic adverse effects to aquatic life).

In addition to the preliminary evaluation of the hazard properties, the cytotoxicity of the (MC)NMs was assessed and the results are displayed in Fig. 5. Fig. 5a shows the viability of THP-1 cells exposed to  $\text{SiO}_2@\text{ZnO}$ ,  $\text{SiO}_2$  or ZnO NMs for 24 h. Viability is expressed as a percentage of control over a range of exposure concentrations from 0 (control) to  $100\text{ }\mu\text{g mL}^{-1}$ . All three materials induced a concentration-dependent decrease in viability, with the ZnO material being the most potent and  $\text{SiO}_2$  being the least potent. Considering that MCNM is a combination of both ZnO and  $\text{SiO}_2$ , and that the curve for MCNM was between the curves for the individual materials, the interaction is additive rather than synergistic or antagonistic. The production of the pro-inflammatory cytokine IL-1 $\beta$  by the NM- and MCNM-treated THP-1 cells (Fig. 5b) also exhibited a concentration-dependent effect. For this particular indicator, there was no significant difference between any of the materials assessed, suggesting that none of the components dominated the pro-inflammatory activity of MCNM and there was no mixture effect.

The *in vitro* assessment of the MCNM hazard indicated that MCNM was more reflective of the cytotoxicity and pro-inflammatory responses induced by ZnO NM rather than  $\text{SiO}_2$  compared at the same mass exposure concentration. This suggested the responses are driven by the ZnO NM component. There is no evidence of an enhanced hazard response due to the  $\text{SiO}_2$  NM; however, a slight increase in response compared to ZnO alone means some interaction between the NM components may impact the bioactivity of ZnO when in MCNM form. Considering the worst-case scenario, it can be assumed in the early innovation stages that the hazard of MCNM is comparable to that of ZnO. Further work in a wider array of human and environmental models for both acute and longer-term exposure will be required in later innovation stages for regulatory risk assessment.

### 3.3 MCNM production and processing phase: human health and safety aspects

The survey to identify the potential sources/points of (MC) NM emission is included in the ESI.<sup>†</sup> According to the results obtained, the OSH assessment during the material synthesis (weighing and calcination) and the (MC)NM-based mortar formulation was carried out using the monitoring procedure described in the Materials and methods section. The background levels of particle number measured over at least 15 min for the three ES considered are summarized in Table S3<sup>†</sup> and the corresponding SEM-EDX images are displayed in Fig. S8–S13.<sup>†</sup> The results in Table S3<sup>†</sup> show that the background levels were in the range of  $3000$  to  $6000\text{ }\#\text{ cm}^{-3}$  for ES1,  $4500$  to  $9500\text{ }\#\text{ cm}^{-3}$  for ES2 and  $2450$  to  $4500\text{ }\#\text{ cm}^{-3}$  for ES3. The SEM-EDX images of the background measured for the three exposure scenarios considered revealed the presence of agglomerated particles in the nm– $\mu\text{m}$  range and their main composition was C, O and Si, but Ca, Al, K and Fe were also detected. Zn was only identified in the mortar formulation laboratory (ES3).

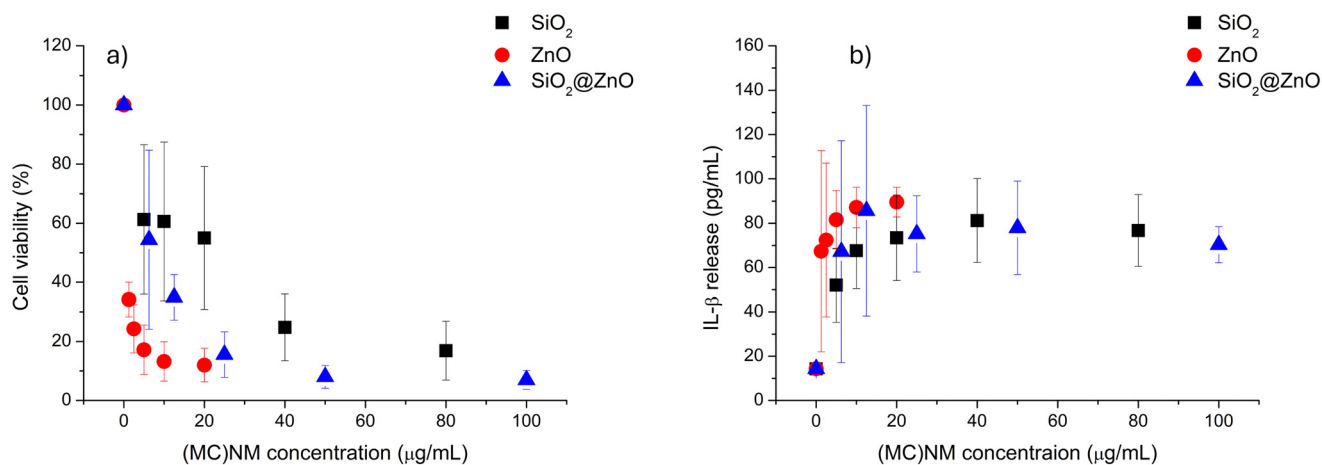
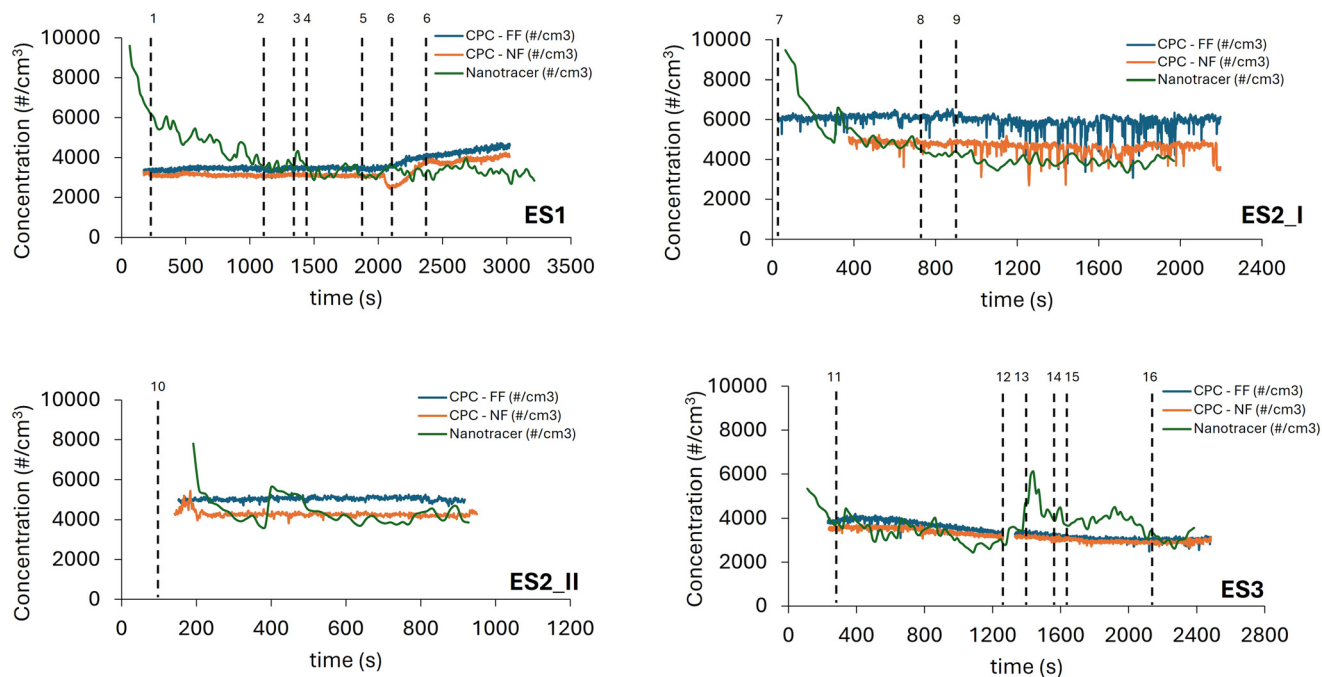


Fig. 5 Cytotoxicity assessment by treating THP-1 cells with  $\text{SiO}_2@\text{ZnO}$  MCNM (blue triangle),  $\text{SiO}_2$  NM (black square) or ZnO NM (red circle) for 24 hours. a) Viability of THP-1 cells assessed by the Alamar blue assay. b) Production of IL-1 $\beta$  by THP-1 cells.





**Fig. 6** Particle number concentration for different exposure scenarios considered (ES1, ES2\_1, ES2\_2, and ES3). Dot lines refer to a specific contributing exposure scenario (CES) listed in Table 1. CPC-FF = condensation particle counter-far field (5–7 m from the source) and CPC-NF (0.5–1 m from the source) = condensation particle counter-near field.

The results of the monitoring campaign through portable air monitoring devices for the three ES, showing particle concentration (number of particles per  $\text{cm}^3$ ) vs. time, are displayed in Fig. 6.

Regarding the measurements for ES1, corresponding to weighing  $\text{ZnAc}_2 \cdot 2\text{H}_2\text{O}$  and  $\text{SiO}_2$  powders, Fig. 6 shows that the particle concentration was relatively stable during the whole activity monitored and did not exceed  $5000 \text{ # cm}^{-3}$ . The only significant increment in the values recorded was observed when  $\text{ZnAc}_2 \cdot 2\text{H}_2\text{O}$  was added. As observed for ES1, the monitoring campaign for ES2 related to the calcination step again revealed that the particle concentration was relatively stable over time, even if all the values started at least  $1000 \text{ # cm}^{-3}$  higher than that in ES1. Concerning the mortar formulation step (ES3), the results in Fig. 6 highlight that while the CPC values were approximately the same during all the activities monitored, the data collected by the NanoTracer presented a peak reaching  $6000 \text{ # cm}^{-3}$ . This is related to the addition of mortar to the stirring tank. According to the NanoGEM Standard Operation Procedures (SOP) for assessing exposure to nanomaterials,<sup>27</sup> a peak is statistically significant as follows:

$$(\text{Peak Value} - \overline{\text{BG}}) > (3 \times S^2(\text{BG})) \quad (2)$$

where  $\overline{\text{BG}}$  is the mean value of the background and  $S^2(\text{BG})$  is the standard deviation of the background.

Therefore, given that  $(\text{Peak Value} - \overline{\text{BG}}) = 1682 \text{ # cm}^{-3}$  and  $(3 \times S^2(\text{BG})) = 1354 \text{ # cm}^{-3}$ , the peak recorded by NanoTracer was considered statistically significant.

The corresponding SEM-EDX images obtained during the monitoring campaign are displayed in Fig. S14–S19,<sup>†</sup> highlighting the presence of some agglomerated particles in the micrometer size range, mainly composed of C, O, Si, while a very low signal corresponding to Zn emerged, which was not present in the background samples.

Subsequently, the results from the monitoring campaign were assessed by adopting the approach reported by Asbach *et al.*, 2014 (ref. 27) as the criteria to follow, where the values of the background (Table S3<sup>†</sup>) and data from tier 2 were compared (Tables S4–S6<sup>†</sup>). Based on this comparison, no significant exposure to particles (*i.e.*, almost all agglomerates in the  $\mu\text{m}$  size range) in any of the three exposure scenarios was determined. A further comparison was also performed between the results included in this work and some recommended limit values (recommended benchmark level – RBL or nano reference values – NRV) proposed by various international bodies, as reported in the literature.<sup>42</sup> For example, according to the Institute for Occupational Safety and Health of the German Social Accident Insurance (IFA), considering an average airborne concentration during an 8 h workday, the RBL values are  $20\,000 \text{ # cm}^{-3}$  (from 1 to 100 nm) for bio persistent granular NMs with a density of  $>6000 \text{ kg m}^{-3}$  and  $40\,000 \text{ # cm}^{-3}$  for particles with a density of  $<6000 \text{ kg m}^{-3}$ . Following the RBL values proposed by IFA, the Social and Economic Council of the Netherlands (SER) proposed the same values, referred to as NRV. According to the results obtained, it can be concluded that none of the exposure scenarios investigated generate a particle concentration higher than the RBL/NRV values. However,

considering the continuous improvements in the risk assessment of advanced materials, including NMs, a further SSbD measure to mitigate the potential exposure to  $\text{SiO}_2$  NMs in powder form through inhalation would be its replacement with colloidal  $\text{SiO}_2$ , but ensuring that the same functionality of the  $\text{SiO}_2@\text{ZnO}$  MCNM is achieved.

### 3.4 MCNM final application phase: human health and environmental aspects

To assess the potential exposure to the (MC)NMs under investigation in the final application phase, the release of potentially hazardous metals (*i.e.*, Zn, Si, and Ti) from both the pristine materials, used as the reference from which the release could be maximum, and the embedded mortars was evaluated. Considering the application of mortars on the surface of buildings that are subject to different weathering agents such as raining events, which can wash out undesirable and harmful substances, their potential release was evaluated considering the worst-case scenario. This test involved the total immersion of a mortar brick in UPW over time, as described by the standard leaching test methods ISO 2812:2018 for mortar immersion conditions and EN 12457-3:2004 for leaching evaluation.<sup>43</sup> The results obtained from these experiments are reported below.

The dissolution of the (MC)NM powder dispersed in UPW was investigated at room temperature by ICP-OES at 0.5, 1, 3, 24 and 48 h after preparation of the dispersion and the values are displayed in Fig. 7. The data are expressed as % of undissolved Zn and Si normalized by the molecular weight for each (MC)NM, considering the 4:1 ratio of  $\text{SiO}_2:\text{ZnO}$  for the MCNM, as initially determined by ICP-OES. The results showed that Si dissolution in UPW, from both individual and MCNM, was negligible over time (around 2%), while Zn dissolution revealed some differences between ZnO and  $\text{SiO}_2@\text{ZnO}$  MCNM only at 48 h, reaching around 9% and 6%, respectively. This slight difference could be ascribed to their

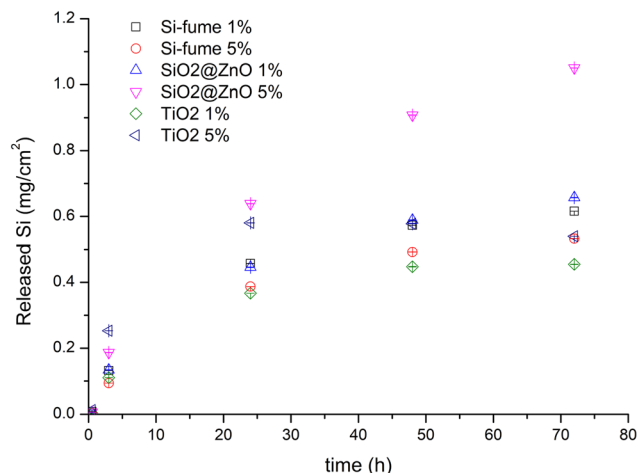


Fig. 8 Si released from the different mortars tested at 0.5, 3, 24, 48 and 72 h.

particle size and surface chemistry, which are generally recognized to be the main physico-chemical parameters affecting the colloidal behaviour of NMs.<sup>44,45</sup> Specifically, as the particle size decreases, the surface area and number of reactive surface sites increase, which often results in different behavior in NMs with respect to their bulk state. In the case of the ZnO NMs investigated in this study, their particle size was <200 nm with an elliptic/roundish shape, while in the case of the core-shell structure, ZnO was present as a thin coating on the surface of  $\text{SiO}_2$  NMs.

Moving from the synthesis to the use phase, the potential release of hazardous substances from (MC)NMs embedded in mortars was investigated. According to the experimental design followed in this work, no release of Zn and Ti from the mortars immersed in UPW after 72 h was detected, while the Si release data (expressed as mg of Si per  $\text{cm}^2$  of mortar) are displayed in Fig. 8. The results of this experiment allowed us to discriminate whether the source of Si was the mortar itself or the MCNMs

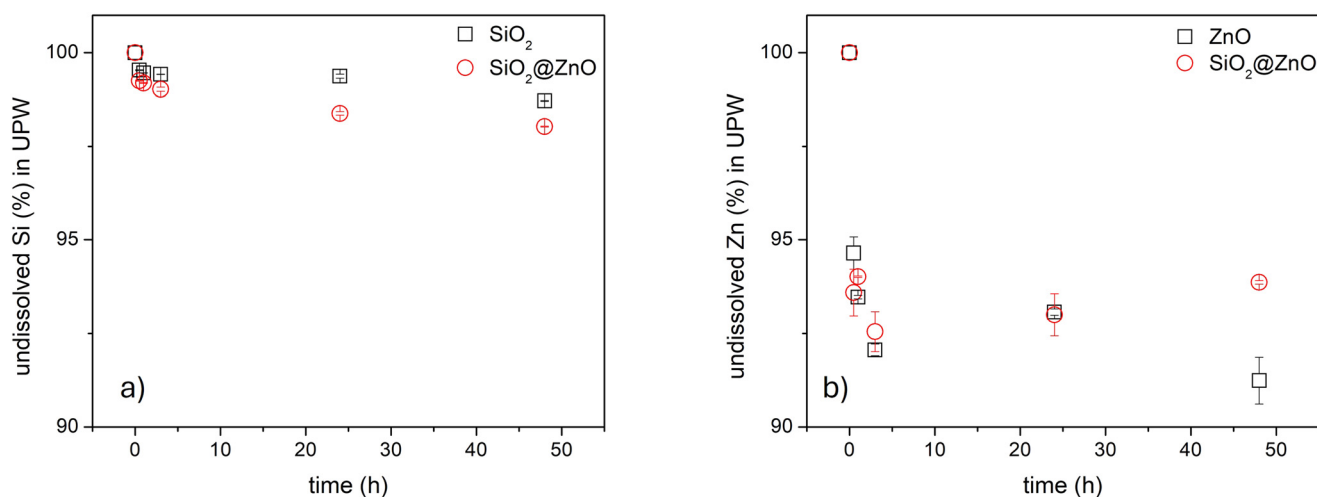


Fig. 7 Dissolution of Si and Zn from both individual and MCNM. a) Si% from  $\text{SiO}_2$  NM and  $\text{SiO}_2@\text{ZnO}$  MCNM. b) Zn% from ZnO NM and  $\text{SiO}_2@\text{ZnO}$  MCNM.

embedded in the mortar, showing that the mortar sample with a higher percentage of MCNM (5%  $\text{SiO}_2\text{@ZnO}$ ) was the only material that released a different Si amount than the others, especially at 48 and 72 h. Therefore, the difference between the maximum amount of Si released from the 5%  $\text{SiO}_2\text{@ZnO}$  mortar and that released by the mortars without Si-based NMs allowed the determination of the amount of Si ascribable to the MCNM. For example, at 72 h, this amount ranged from around 0.4 to 0.6  $\text{mg cm}^{-2}$ , suggesting the maximum Si release of 4% with respect to the maximum amount potentially released by the MCNM.

### 3.5 SbD assessment summary

The combination of experimental data and literature information reported in this work allowed us to assess the safety of the  $\text{SiO}_2\text{@ZnO}$  MCNM. The results from the hazard assessment (step 1) are comprised of information from its basic physicochemical characteristics, preliminary evaluation of hazard properties and rapid cytotoxicity assays. The results of this first step showed the following: a) the MCNM was composed of an  $\text{SiO}_2$  NP core, with non-homogeneous coverage of ZnO, with a 4 : 1  $\text{SiO}_2$  : ZnO ratio; b) the hazard evaluation of the components used for the MCNM synthesis reached level 1 (according to the SSbD framework evaluation system), given that  $\text{ZnAc}_2\cdot\text{H}_2\text{O}$  can induce chronic environmental toxicity; and c) the *in vitro* assessment of the MCNM suggested that its hazard can be comparable to that of the ZnO NPs tested. The results from the air monitoring campaign (step 2) did not highlight any exposure of workers to particles for the three exposure scenarios considered, disclosing that the particle concentration recorded was always below the recommended RBL/NRV thresholds proposed by the IFA German Institute and by the SER Dutch advisory body. Lastly, the release experiments from the mortars with embedded MCNM (step 3) showed only the slight release of Si from the mortar loaded with the highest MCNM concentration. Therefore, considering the worst-case scenario, these SbD data suggest that this  $\text{SiO}_2\text{@ZnO}$  MCNM can be considered similar to <200 nm round-shape ZnO NPs for its cytotoxicity response but it does not pose significant concerns resulting from the occupational exposure assessment as well as from the final application phase. These outcomes, not only including the new experimental data but also the approach to assess the safety dimension of MCNM, can be very useful to different stakeholders, including value chain actors in the building sector such as governments, private investors, construction companies and real estate agents. Moreover, researchers, policy makers and the European Commission can take advantage of this case study to guide future efforts toward the sustainability assessment of advanced materials.

## 4. General remarks and conclusions

This work falls within the context of implementing the European Commission Chemicals Strategy for Sustainability and focused on the safety dimension aspects of the SSbD

framework proposed by the JRC. The stepwise approach followed can be considered one of the activities envisaged by the European Commission to support the testing and potential refinement of the framework, which was developed for chemicals and materials in general, and not specifically for advanced materials. The MCNM studied here is novel and innovative, and hence the existing data required for applying the framework are scarce. For example, the evaluation of hazard properties in step 1 of the framework requires data, which are not yet available from CLP and REACH registrations. Therefore, the framework encourages the use of New Approach Methodologies (NAMs), such as *in vitro* and *in silico* assays, as relevant means to support the innovation process in the early stages because even if they are not fully validated for regulatory purposes, NAMs can provide valuable information that is relatively economical to obtain.<sup>46</sup> Hence, faster and more affordable *in vitro* assays or *in silico* tools (in this case specifically for MCNMs) can be valuable to gather this information in the early innovation stages. Regarding these tools, the selection of the most suitable benchmark with which to compare the *in vitro* results of MCNM is crucial but a challenge at the same time because it depends greatly on the physicochemical characteristics of the MCNM and how it may interact with the surroundings.

The following tools/procedures related to experimental activities were identified to operationalize the safety aspects of the SSbD framework to evaluate the safety of the  $\text{SiO}_2\text{@ZnO}$  MCNM in the early innovation stage including basic physicochemical characterization, hazard classification, rapid *in vitro* assays, the three-tiered methodology developed by the nanoGEM research project and the release investigation of potential hazardous substances from MCNM-based mortars. It is worth noting that except for the procedure addressing step 3 of the SSbD framework, chosen specifically for this case study, the tools identified in this study are generally applicable. As a result, they can be effectively utilized for implementing the safety aspect of the SSbD framework for MCNMs.

Concerning the re-design phase of this MCNM, a valid alternative to decrease the exposure of workers during the synthesis of MCNM, acting in the early-stage design will be the replacement of the  $\text{SiO}_2$  powder with colloidal  $\text{SiO}_2$ , ensuring the same material functionality. Therefore, this approach can have a positive impact in the early stage of the design phase of chemicals/materials, steering innovation towards the green industrial transition, beyond current regulatory compliance.

In line with the framework, further assessments are ongoing to include the estimation of the environmental, social and economic impacts to address the other steps not covered in this work. For example, even if recycling of building materials is constantly increasing, it is worth noting that the end-of-life phase of MCNM-based mortars was not considered in this work, although it may be a critical step contributing to a more comprehensive safety and sustainability assessment. The ongoing research activity





within SUNSHINE is now focusing on integrating the information generated herein with the screening level approach already developed by project partners,<sup>16</sup> with the aim to include all these information in the SUNSHINE SSbD e-infrastructure. This methodology will provide an interactive tool to meet the expectations of stakeholders along the value chain of this MCNM, which can also be extended to other advanced materials. The digital integration of information through the e-infrastructure will help producers, both large companies and small and medium enterprises, to clearly identify criticalities along the entire MCNM life cycle in which to take action by eliminating hazardous materials, replacing them with less hazardous ones and disclosing the environmental, economic and social impacts compared to a benchmark. Overall, we believe that this approach is consistent with the Green Deal ambition towards a zero-pollution toxic-free environment to adequately protect citizens and the environment.

## Disclaimer

The content expressed in this paper is solely the opinion of the authors and does not necessarily reflect the opinion of their institutions.

## Data availability

The authors confirm that the data supporting the findings of this study are available within the article and its ESI.†

## Conflicts of interest

There are no conflicts to declare.

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## References

- 1 Y.-W. Huang, M. Cambre and H.-J. Lee, The Toxicity of Nanoparticles Depends on Multiple Molecular and Physicochemical Mechanisms, *Int. J. Mol. Sci.*, 2017, **18**, 2702.
- 2 K. Luyts, D. Napierska, B. Nemery and P. H. M. Hoet, How physico-chemical characteristics of nanoparticles cause their toxicity: complex and unresolved interrelations, *Environ. Sci.: Processes Impacts*, 2013, **15**, 23–38.
- 3 A. Sukhanova, S. Bozrova, P. Sokolov, M. Berestovoy, A. Karaulov and I. Nabiev, Dependence of Nanoparticle Toxicity on Their Physical and Chemical Properties, *Nanoscale Res. Lett.*, 2018, **13**, 44.
- 4 N. M. Tran, A. N. Nguyen, J. Bae, J. Kim, D. Kim and H. Yoo, Recent strategies for constructing hierarchical multicomponent nanoparticles/metal–organic framework hybrids and their applications, *Nanoscale Adv.*, 2023, **5**, 3589–3605.
- 5 D. Li, F. Guo and L. Qi, Gold Nanoarrow-Based Core–Shell and Yolk–Shell Nanoparticles for Surface-Enhanced Raman Scattering, *ACS Appl. Nano Mater.*, 2022, **5**, 126–132.
- 6 Y. Xiang, Y. Huang, B. Xiao, X. Wu and G. Zhang, Magnetic yolk-shell structure of ZnFe<sub>2</sub>O<sub>4</sub> nanoparticles for enhanced visible light photo-Fenton degradation towards antibiotics and mechanism study, *Appl. Surf. Sci.*, 2020, **513**, 145820.
- 7 G. F. Huseien, Potential Applications of Core-Shell Nanoparticles in Construction Industry Revisited, *Appl. Nano*, 2023, **4**, 75–114.
- 8 A. Speziale, J. F. González-Sánchez, B. Taşçı, A. Pastor, L. Sánchez, C. Fernández-Acevedo, T. Oroz-Mateo, C. Salazar, I. Navarro-Blasco, J. M. Fernández and J. I. Alvarez, Development of Multifunctional Coatings for Protecting Stones and Lime Mortars of the Architectural Heritage, *Int. J. Archit. Herit.*, 2020, **14**, 1008–1029.
- 9 F. Zhang, Z. Wang, W. J. G. M. Peijnenburg and M. G. Vijver, Review and Prospects on the Ecotoxicity of Mixtures of Nanoparticles and Hybrid Nanomaterials, *Environ. Sci. Technol.*, 2022, **56**, 15238–15250.
- 10 D. Pozzi, G. Caracciolo, L. Digiacomo, V. Colapicchioni, S. Palchetti, A. L. Capriotti, C. Cavaliere, R. Zenezini Chiozzi, A. Puglisi and A. Laganà, The biomolecular corona of nanoparticles in circulating biological media, *Nanoscale*, 2015, **7**, 13958–13966.
- 11 C. Caldeira, R. Farcas, A. I. Garmendia, L. Mancini, D. Tosches, A. Amelio, K. Rasmussen, H. Rauscher, S. J. Riego and S. Sala, *Safe and sustainable by design chemicals and materials - Framework for the definition of criteria and evaluation procedure for chemicals and materials*, EUR 31100 EN, Publications Office of the European Union, Luxembourg, 2022, DOI: [10.2760/487955](https://doi.org/10.2760/487955), JRC128591, ISBN 978-92-76-53264-4, <https://publications.jrc.ec.europa.eu/repository/handle/JRC128591>.
- 12 European Commission, *Commission recommendation of 8.12.2022 establishing a European assessment framework for 'safe and sustainable by design' chemicals and materials*, Brussels, 2022, 8.12.2022 C (2022) 8854 final.
- 13 A. Kraegeloh, B. Suarez-Merino, T. Sluijters and C. Micheletti, Implementation of Safe-by-Design for Nanomaterial Development and Safe Innovation: Why We Need a Comprehensive Approach, *Nanomaterials*, 2018, **8**, 239.
- 14 L. Yan, F. Zhao, J. Wang, Y. Zu, Z. Gu and Y. Zhao, A Safe-by-Design Strategy towards Safer Nanomaterials in Nanomedicines, *Adv. Mater.*, 2019, **31**, 1805391.
- 15 A. Sudheshwar, C. Apel, K. Kümmerer, Z. Wang, L. G. Soeteman-Hernández, E. Valsami-Jones, C. Som and B. Nowack, Learning from Safe-by-Design for Safe-and-Sustainable-by-Design: Mapping the current landscape of Safe-by-Design reviews, case studies, and frameworks, *Environ. Int.*, 2024, **183**, 108305.



- 16 L. Pizzol, A. Livieri, B. Salieri, L. Farcas, L. G. Soeteman-Hernández, H. Rauscher, A. Zabeo, M. Blois, A. L. Costa, W. Peijnenburg, S. Stoycheva, N. Hunt, M. J. López-Tendero, C. Salgado, J. J. Reinoso, J. F. Fernández and D. Hristozov, Screening level approach to support companies in making safe and sustainable by design decisions at the early stages of innovation, *Clean. Environ. Syst.*, 2023, **10**, 100132.
- 17 European Committee for Standardization, EN 196-1:2016 - Methods of Testing Cement - Part1: Determination of Strength, 2016.
- 18 European Committee for Standardization, EN 1015-11:2019 - Methods of test for mortar for masonry - Part 11: Determination of flexural and compressive strength of hardened mortar, 2019.
- 19 K. Rasmussen, H. Rauscher, A. Mech, J. Riego Sintes, D. Gilliland, M. González, P. Kearns, K. Moss, M. Visser, M. Groenewold and E. A. J. Bleeker, Physico-chemical properties of manufactured nanomaterials - Characterisation and relevant methods. An outlook based on the OECD Testing Programme, *Regul. Toxicol. Pharmacol.*, 2018, **92**, 8–28.
- 20 H. Bresch, V.-D. Hodoroaba, A. Schmidt, K. Rasmussen and H. Rauscher, Counting Small Particles in Electron Microscopy Images - Proposal for Rules and Their Application in Practice, *Nanomaterials*, 2022, **12**, 2238.
- 21 European Commission, Regulation (EC) No 1272/2008 of the European Parliament and of the Council of 16 December 2008 on classification, labelling and packaging of substances and mixtures, amending and repealing Directives 67/548/EEC and 1999/45/EC, and amending Regulation (EC) No 1907/2006, *Official Journal of the European Union L 353*, 31.12.2008, 2008, pp. 1–1355.
- 22 European Commission, Regulation (EC) No 1907/2006 of the European Parliament and of the Council of 18 December 2006 concerning the Registration, Evaluation, Authorisation and Restriction of Chemicals (REACH), establishing a European Chemicals Agency, amending Directive 1999/45/EC and repealing Council Regulation (EEC) No 793/93 and Commission Regulation (EC) No 1488/94 as well as Council Directive 76/769/EEC and Commission Directives 91/155/EEC, 93/67/EEC, 93/105/EC and 2000/21/EC, *Official Journal of the European Union, L 396*, 30.12.2006, 2006, pp. 1–849.
- 23 N. Ruijter, L. G. Soeteman-Hernández, M. Carrière, M. Boyles, P. McLean, J. Catalán, A. Katsumiti, J. Cabellos, C. Delpivo, A. Sánchez Jiménez, A. Candalija, I. Rodríguez-Llopis, S. Vázquez-Campos, F. R. Cassee and H. Braakhuis, The State of the Art and Challenges of In Vitro Methods for Human Hazard Assessment of Nanomaterials in the Context of Safe-by-Design, *Nanomaterials*, 2023, **13**, 472.
- 24 H. M. Braakhuis, A. G. Oomen and F. R. Cassee, The State of the Art and Challenges of In Vitro Methods for Human Hazard Assessment of Nanomaterials in the Context of Safe-by-Design, *Toxicol. Appl. Pharmacol.*, 2016, **299**, 3–7.
- 25 B. Sun, X. Wang, Z. Ji, R. Li and T. Xia, NLRP3 Inflammasome Activation Induced by Engineered Nanomaterials, *Small*, 2013, **9**, 1595–1607.
- 26 <https://www.patrols-h2020.eu/publications/sops/index.php>.
- 27 C. Asbach, T. A. J. Kuhlbusch, B. Stahlmecke, H. Kaminski, H. J. Kiesling, M. Voetz, D. Dahmann, U. Götz, N. Dziurawitz and S. Plitzko, Measurement and monitoring strategy for assessing workplace exposure to airborne nanomaterials, in *Safety of Nanomaterials along Their Lifecycle: Release, Exposure, and Human Hazards*, 2014, pp. 233–246.
- 28 M. Methner, L. Hodson and C. Geraci, Nanoparticle emission assessment technique (NEAT) for the identification and measurement of potential inhalation exposure to engineered nanomaterials—part A, *J. Occup. Environ. Hyg.*, 2010, **7**, 127–132.
- 29 OECD, Strategies, techniques and sampling protocols for determining the concentrations of manufactured nanomaterials in air at the workplace, Series on the Safety of Manufactured Nanomaterials N° 82, ENV/JM/MONO, 2017, vol. 30.
- 30 OECD, Series on the safety of manufactured nanomaterials number 11 - Emission assessment for identification of sources and release of airborne manufactured nanomaterials in the workplace: compilation of existing guidance, ENV/JM/MONO, 2009, vol. 16.
- 31 BAuA, BG RCI, IFA, IUTA, TUD and VCI, *Tiered Approach to an Exposure Measurement and Assessment of Nanoscale Aerosols Released from Engineered Nanomaterials in Workplace Operations*, 2011.
- 32 European Committee for Standardization, EN17058:2018 - Workplace exposure - Assessment of exposure by inhalation of nano-objects and their aggregates and agglomerates, 2018.
- 33 European Chemicals Agency, *An illustrative example of the exposure scenarios to be annexed to the safety data sheet. Part 1, Introductory note*, European Chemicals Agency, 2017.
- 34 European Chemicals Agency, *Guidance on information requirements and chemical safety assessment - Chapter 14: occupational exposure assessment, version 3.0 - August 2016*, European Chemicals Agency, 2016.
- 35 G. M. DeLoid, J. M. Cohen, G. Pyrgiotakis and P. Demokritou, Preparation, characterization, and in vitro dosimetry of dispersed, engineered nanomaterials, *Nat. Protoc.*, 2017, **12**, 355–371.
- 36 International Organization for Standardization, ISO 2812-1:2017(en), Paints and varnishes - Determination of resistance to liquids — Part 1: Immersion in liquids other than water, 2017.
- 37 UNI EN 12457-3, 2004, Characterization of waste - Leaching - Compliance test for leaching of granular waste materials and sludges - Part 3: Two stage batch test at a liquid to solid ratio of 2 l/kg and 8 l/kg for materials with high solid content and with particle size below.
- 38 OECD, Physical-chemical decision framework to inform decisions for risk assessment of manufactured nanomaterials, ENV/JM/MONO, 2019, vol. 12.
- 39 S. Gupta, Smart Nanoconcretes and Cement-Based Materials, in *Smart Nanoconcretes and Cement-Based Materials*, ed. M. S. Liew, P. Nguyen-Tri, T. A. Nguyen and S. Kakooei, Elsevier, 2020, pp. 601–617.
- 40 W. Brand, P. C. E. van Kesteren, R. J. B. Peters and A. G. Oomen, Issues currently complicating the risk assessment of



- synthetic amorphous silica (SAS) nanoparticles after oral exposure, *Nanotoxicology*, 2021, **15**, 905–933.
- 41 P. C. E. van Kesteren, F. Cubadda, H. Bouwmeester, J. C. H. van Eijkeren, S. Dekkers, W. H. de Jong and A. G. Oomen, Novel insights into the risk assessment of the nanomaterial synthetic amorphous silica, additive E551, in food, *Nanotoxicology*, 2015, **9**, 442–452.
  - 42 M. Visser, I. Gosens, D. Bard, P. van Broekhuizen, G. Janer, E. Kuempel, M. Riediker, U. Vogel and S. Dekkers, Towards health-based nano reference values (HNRVs) for occupational exposure: Recommendations from an expert panel, *NanoImpact*, 2022, **26**, 100396.
  - 43 A. Brunelli, L. Calgaro, E. Semenzin, V. Cazzagon, E. Giubilato, A. Marcomini and E. Badetti, Leaching of nanoparticles from nano-enabled products for the protection of cultural heritage surfaces: a review, *Environ. Sci. Eur.*, 2021, **33**, 48.
  - 44 E. A. Meulenkamp, Size Dependence of the Dissolution of ZnO Nanoparticles, *J. Phys. Chem. B*, 1998, **102**, 7764–7769.
  - 45 A. Albanese, P. S. Tang and W. C. W. Chan, The Effect of Nanoparticle Size, Shape, and Surface Chemistry on Biological Systems, *Annu. Rev. Biomed. Eng.*, 2012, **14**, 1–16.
  - 46 E. Abbate, I. Garmendia Aguirre, G. Bracalente, L. Mancini, D. Tosches, K. Rasmussen, M. Bennett, H. Rauscher and S. Sala, *Sustainable by Design chemicals and materials - Methodological Guidance*, Publications Office of the European Union, Luxembourg, 2024, DOI: [10.2760/28450](https://doi.org/10.2760/28450), JRC138035.

