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## A refined dose metric for nanotoxicology based on surface site reactivity for oxidative potential of engineered nanomaterials†

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The increasing production of engineered nanomaterials (ENMs) raises significant concerns about human and environmental exposure, making it essential to understand the mechanisms of their interaction with biological systems to manage the associated risks. To address this, we propose categorizing ENM reactivity using in chemico methodologies. Surface analysis through methanol chemisorption and temperature-programmed surface reaction allows for the determination of reactive surface sites, providing accurate estimates of effective ENM doses in toxicity studies. Additionally, antioxidant consumption assays (dithiothreitol, cysteine, and glutathione) and reactive oxygen species (ROS) generation assays (RNO and DCFH2-DA) are employed to rank the oxidative potential of ENM surface sites in a cell-free environment. Our study confirms the classification of ZnO NM-110, ZnO NM-111, CuO, and carbon black as highly oxidant ENMs, while TiO2 NM-101 and NM-105 exhibit low oxidative potential due to their acidic surface sites. In contrast, CeO2 NM-211 and NM-212 demonstrate redox surface sites. SiO<sub>2</sub> nanomaterials (NM-200 and NM-201) are shown to be inert, with low oxidation rates and minimal reactive surface density, despite their high surface area. Quantifying reactive surface sites offers a refined dose metric for assessing ENM toxicity, advancing safe-by-design nanomaterial development.

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### Introduction

Engineered nanomaterials (ENMs) are increasingly utilized in various sectors, including catalysis, agriculture, cosmetics, electronics, medicine, or food.1,2 The massive production of these materials calls for an accurate evaluation of their impact on both human health and the environment.3 Understanding their modes of action enables the adoption of Safe and Sustainable by Design (SSbD) approaches to develop nano-

growing importance of New Approach Methodologies (NAMs) in

the evaluation of substances, including ENMs. 9,10 NAMs,

particularly those designed for ENM screening, for instance by

enabled products characterized by low risks throughout their

life cycle: these approaches make it possible to design ENMs or nano-enabled products with safety considerations integrated

early in the product development stage. However, the physico-

chemical properties that make them useful also complicate defining their dosage. Unlike bulk chemicals, mass and mole exposure do not always correlate with the ENM exposure dose; different particle sizes dramatically change the fraction of ENM that is exposed. In addition, even for a given ENM, the surface area may have very different populations of reactive sites depending on exposed planes, defects and other variables.4-7 For this reason, traditional approaches to chemical safety assessment, which rely on established toxicological protocols, may not fully address the unique challenges presented by ENMs. Furthermore, nano-derived effects as well as the complexity and the speed of new innovations for ENMs require methodological approaches that can be run with less timeconsuming procedures.8 In addition, the field of toxicology is currently undergoing a significant transformation, moving towards animal-free methods. This context underscores the

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measuring ENM reactivity, are valuable tools in the Safe by Design (SbD) framework.

This paper provides a phenomenological description of the reactivity of ENMs by linking physicochemical characteristics to their oxidative potential, a parameter that has been proposed to rank the risk of ENMs. 11-16 often quantified by FRAS, 17,18 CPH, and DMPO probes.19 These methods, however, are blind to the characteristics of the surface that determine the observed oxidative damage. From a surface-chemistry point of view, it becomes important, in fact, to determine the fraction of the whole surface that is really active, and the nature of this reactive surface. To this goal, we propose a novel, in chemico methodology to identify the nature, number, and reactivity of surface sites on an ENM using probe molecules. Gas-phase methanol chemisorption and TPSR are used to quantify the reactive site surface density and their reactive profile. The intrinsic oxidative potential is measured via oxidation rates for (1) ROS generation and (2) antioxidant consumption. We employed antioxidant consumption assays (dithiothreitol, cysteine, and glutathione) and reactive oxygen species (ROS) generation assays (detected by N,N-dimethyl-4-nitrosoaniline, RNO, and 2,7-dichlorodihydrofluorescein diacetate DCFH2-DA).18 Reactive surface site quantification is used to normalize oxidative potential data, applying the concept of Oxidative Turnover Frequency (OxTOF). This normalization enables a refined and relevant definition of effective dose in terms of reactive surface sites. Therefore, the paper emphasizes that integrating surface reactivity and material chemistry concepts into toxicology protocols may allow an accurate screening and comprehensive assessment of ENM risks.7 The study combines the analysis of the reactive surface sites with the data of reactivity to rank a set of 14 common nanomaterials based on the estimations of their oxidative potential normalized to the number of active sites.

## 2 Experimental

#### 2.1 Nanomaterials

Fourteen common powdered ENMs were analyzed without any pretreatment:  $TiO_2$  NM-101 (JRCNM01001a),  $TiO_2$  NM-105 (JRCNM01005a),  $CeO_2$  NM-211 (JRCNM02101a),  $CeO_2$  NM-212 (JRCNM02102a), ZnO NM-110 (JRCNM62101a), ZnO NM-111 (JRCNM01101a), SiO<sub>2</sub> NM-200 (JRCNM02000a), SiO<sub>2</sub> NM-201 (JRCNM02001a), MWCNT NM-400 and MWCNT NM-401 were supplied by Joint Research Centre (JRC), while CuO (ref. number: 544868),  $Mn_2O_3$  (ref. number: 933791), carbon black (PRINTEX® 90, CB) and  $Fe_2O_3$  (ref. number: 544884) were obtained from Sigma-Aldrich (SA).  $TiO_2$  NM-105 is the same as Aeroxide P25 (Degussa/Evonik), which comprises 73–85% anatase, 14–17% rutile, and 0–13% amorphous titania.  $^{20-22}$  The main physicochemical properties of the evaluated NMs are summarized in Table S1.†

# 2.2 Quantification and characterization of reactive surface sites

Reactive sites on the surface of metal oxide nanomaterials were probed by chemisorption of methanol followed by its

Temperature Programmed Surface Reaction (TPSR)23,24 to identify the number and characteristics of reactive surface sites.25 The protocol has been adapted from our previous studies.<sup>26,27</sup> Briefly, 100-250 mg of NM powder (nanoparticle aggregate size from 25 to 100 µm) were mixed with 0.5 g SiC (black 180, Navarro SiC S.A.) and introduced into a fixed-bed reactor (0.4 cm diameter) to follow the procedure described elsewhere.26 Methanol chemisorption at 100 °C (50 °C in the case of CuO and ZnO NM-110 due to their high reactivity) results in the release of H<sub>2</sub>O molecules due to the formation of methoxy species (-OCH<sub>3</sub>) bonding to the reactive surface sites. During linear heating, different reaction products are released according to the reactivity of the surface sites: some unreacted methanol can be released in low-activity sites, acidic sites yield dimethyl ether (CH<sub>3</sub>OCH<sub>3</sub>), redox sites generate formaldehyde (HCHO), and carbon dioxide (CO<sub>2</sub>) can be produced on basic sites (typically above 300 °C) or highly reactive redox sites (below 300 °C). This catalytic clue about the primary reactivity of a nanomaterial is independent of the stability of a dispersion, pretreatments, bio-transformations or ion release. All the evaluated nanomaterials were previously screened for thermal stability in an STA 6000 simultaneous thermal analyzer (PerkinElmer). The thermal stability of the nanotubes up to 450 °C was confirmed by the supplier.28

#### 2.3 Consumption of antioxidants

To estimate the ability of ENMs to consume antioxidants, we measured the rates at which terminal thiol groups are oxidized in three biologically relevant molecules: (i) dithiothreitol (DTT), (ii) cysteine (Cys) and (iii) glutathione (GSH). DTT is a synthetic di-sulfide used as an unspecific reducing agent. Glutathione is a natural antioxidant characterized by the lowest redox potential that can be found in a cell.<sup>13</sup> This low redox potential (*i.e.*, high thermodynamic tendency to reduce oxidants) also enables, in principle, the direct oxidation of the S-terminus by species other than ROS (*e.g.*, metal ions).<sup>13,29</sup> Cysteine is an amino acid, and the oxidation of its terminal –S is relevant for the oxidative damage of proteins.<sup>30</sup> The details of these acellular procedures are provided in the ESI.†

#### 2.4 Generation of reactive oxygen species

We estimated the rates of ROS generation by two acellular approaches. We used *N*,*N*-dimethyl-4-nitrosoaniline (RNO) for the spectroscopic determination of the production of hydroxyl radicals (*i.e.*, OH<sup>+</sup>),<sup>31</sup> and 2',7'-dichlorodihydrofluorescein diacetate (DCFH<sub>2</sub>-DA) to detect ROS other than \*OH.<sup>18</sup> The specifics of the procedure are detailed in the ESI.† The methodology for DCFH<sub>2</sub>-DA assay is based on the SOP developed by the GRACIOUS EU project (GA760840).

# 2.5 Ranking of ENMs reactivity and investigation of reactivity-toxicity correlations

We used *K*-means and hierarchical dendrogram unsupervised learning algorithms to detect similarity-based descriptors and patterns of reactivity.<sup>32–34</sup> The main goal of this approach was to group the ENMs based on distance-based metrics or similarity

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according to their oxidative potential. Reactivity data based on ENM mass, surface and reactive surface sites exposed to the probe molecule were compared in a pair plot. The oxidative potential per reactive surface site obtained for the different probe molecules was compared utilizing Pearson's correlation coefficient.

We used Spearman's coefficient to compare the acellular oxidative potential of ENMs with outcomes of in vitro studies by LDH and WST-1 assays in lung cell lines A549 and dTHP-1.35 As a toxicological metric, we employed the highest concentration that showed no adverse effects after 24 h exposure; gravimetric concentrations of ENMs (µg mL<sup>-1</sup>) were converted to reactive site concentrations ( $\mu$ mol L<sup>-1</sup>) based on the data obtained with a methanol probe (one methanol molecule titrates one reactive surface site).

#### Results 3

#### Number and nature of reactive surface sites of ENMs

For each ENM, we evaluated (i) specific surface area  $(m^2 g^{-1})$ , (ii) specific number of reactive surface sites (mmol g<sup>-1</sup>, one methanol molecule titrates one reactive surface site), and (iii) reactive site surface density (sites per nm<sup>2</sup>) (Table 1). The ENMs with the largest specific surface area were CB (317 m<sup>2</sup> g<sup>-1</sup>), MWCNT NM-400 (240 m<sup>2</sup> g<sup>-1</sup>) and TiO<sub>2</sub> NM-101 (225 m<sup>2</sup> g<sup>-1</sup>), whereas the two ZnO, Mn<sub>2</sub>O<sub>3</sub> and CuO exhibited the lowest values: 9 m<sup>2</sup> g<sup>-1</sup> (NM-110), 15 (NM-111), 17 m<sup>2</sup> g<sup>-1</sup> and 12 m<sup>2</sup> g<sup>-1</sup>, respectively. Regarding the number of reactive surface sites, TiO2 NM-101 had the highest specific number (2.8 mmol g<sup>-1</sup>), followed by MWCNT NM-400, TiO<sub>2</sub> NM-105 and MWCNT NM-401, with values between 1.7 and 1.1 mmol  $g^{-1}$ , whereas zinc oxides, ceria, silica, CuO and Mn2O3 exhibited a lower specific number of sites  $(0.1-0.4 \text{ mmol g}^{-1})$ . These were specific values (per unit mass); however, the analysis of the reactive site surface density, with a variety of values, revealed that the physical specific surface area did not directly correlate with the

number of reactive surface sites, as determined by methanol chemisorption. The ENM with the highest reactive site surface density was CuO (22 sites per nm2), followed by ZnO NM-110, Fe<sub>2</sub>O<sub>3</sub> and TiO<sub>2</sub> NM-105, while the lowest values were those of silica and MWCNT, all below 5 sites per nm<sup>2</sup>.

The nature of surface reactivity of the ENMs was tested by MeOH-TPSR (Fig. 1) to identify the type of reactive surface sites and classify them accordingly. TiO<sub>2</sub> nanomaterials presented a main desorption band related to dimethyl ether formation (Fig. 1A and D), i.e., to acid reactivity, in line with the TPSR results obtained in a previous study with anatase titania DT51 from CristalACTiVTM.26 In addition, low signals of formaldehyde and carbon dioxide were registered, indicating the presence of some redox and basic sites in titania samples. Ceria, however, exhibited mainly redox reactivity (Fig. 1C and F); NM-211 showed the maximum methanol conversion to formaldehyde at 259 °C, whereas NM-212 formed formaldehyde at two types of sites, more (207 °C) and less (283 °C) active than those of NM-211, with intermediate carbon dioxide generation at around 265 °C. CO<sub>2</sub> reports two kinds of reactive surface sites: (a) highly oxidising sites readily convert surface methoxy species to CO2 with a maximum at temperatures ranging from 170 °C or lower to ca. 280 °C; (b) basic sites bind methoxy species so strongly that these may only desorb at high temperature (above 300 °C), being readily converted to CO2, as described in the literature.25 In the case of ZnO, CuO, Mn<sub>2</sub>O<sub>3</sub> and Fe<sub>2</sub>O<sub>3</sub> carbon dioxide was the only reaction product. Fig. 1B shows that CuO was a highly oxidizing material (CO<sub>2</sub> maximum formation at 221 °C), and Mn<sub>2</sub>O<sub>3</sub> was the next highly oxidizing material (CO<sub>2</sub> maximum at 274 °C in Fig. 1G). The results for ZnO and Fe<sub>2</sub>O<sub>3</sub> are shown in Fig. 1E and H, where it can be observed that CO<sub>2</sub> formed at significantly higher temperatures, indicating an essentially basic character of the reactive sites. The temperature of maximum CO2 formation can be used to rank the reactivity of these four ENMs from a kinetic point of view as follows: CuO  $(221 \, {}^{\circ}\text{C}) > \text{Mn}_2\text{O}_3 (274 \, {}^{\circ}\text{C}) > \text{Fe}_2\text{O}_3 (314 \, {}^{\circ}\text{C}) \ge \text{ZnO} (315 \, {}^{\circ}\text{C}); \text{ here,}$ 

Table 1 Surface descriptors for the tested ENMs. Specific surface area was obtained by using N<sub>2</sub> adsorption isotherms and the specific number of reactive surface sites by methanol chemisorption. Their ratio is the reactive site surface density. The number of reactive surface sites on carbon black could not be measured due to agglomeration issues in the reactor

Nanomaterial	Specific surface area $(m^2 g^{-1})$	Specific number of reactive sites (mmol $g^{-1}$ )	Reactive site surface density (reactive site per nm <sup>2</sup> )	
TiO <sub>2</sub> NM-101	225	2.8	7	
TiO <sub>2</sub> NM-105	58	1.1	13	
ZnO NM-110	9	0.2	17	
ZnO NM-111	15	0.1	5	
SiO <sub>2</sub> NM-200	182	0.3	1	
SiO <sub>2</sub> NM-201	140	0.5	2	
CeO <sub>2</sub> NM-211	76	0.4	6	
CeO <sub>2</sub> NM-212	25	0.4	10	
MWCNT NM-400	$240^a$	1.7	4	
MWCNT NM-401	$140^a$	1.1	4	
CuO	12	0.4	22	
Carbon black	317	NA	NA	
Fe <sub>2</sub> O <sub>3</sub>	41	1.0	15	
$Mn_2O_3$	17	0.3	11	

<sup>&</sup>lt;sup>a</sup> Data obtained from the supplier.

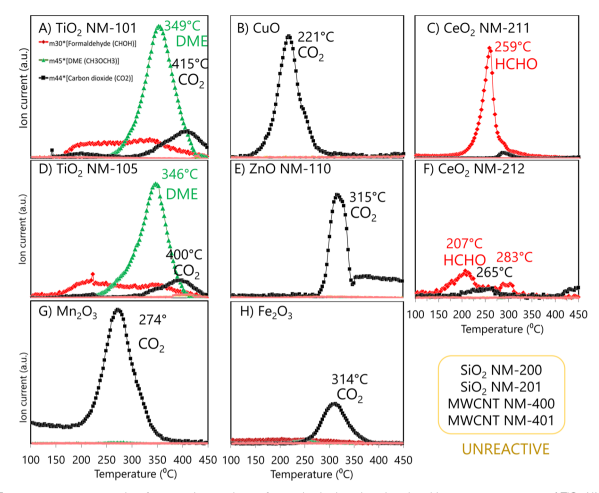


Fig. 1 Temperature-programmed surface reaction products of pre-adsorbed methanol analysed by mass spectroscopy of  $TiO_2$  NM-101 (A), CuO (B), CeO<sub>2</sub> NM-211 (C),  $TiO_2$  NM-105 (D), ZnO NM-110 (E), CeO<sub>2</sub> NM-212 (F), Mn<sub>2</sub>O<sub>3</sub> (G), and  $Fe_2O_3$  (H). Formaldehyde (red diamond) is formed at redox surface sites, dimethyl ether (green triangle) at acid surface sites, and carbon dioxide (black square) at basic or highly reactive redox surface sites. No detectable formation of species with mass 60 (methyl formate) or mass 75 (dimethoxymethane) was observed, indicating the absence of bifunctional reactive surface site activity.

lower temperatures are representative of methanol over-oxidation due to highly reactive redox sites, while methanol combustion is the  $\rm CO_2$  production mechanism at high temperatures, indicative of strong affinity between basic sites and the methoxy group. Methanol-TPSR data for highly inert  $\rm SiO_2$  (NM-200 and NM-201) and MWCNT (NM-400 and NM-401) are summarized in Fig. S1.† Only  $\rm SiO_2$  NM-201 and MWCNT NM-400 showed some reactivity, with a slight band related to  $\rm CO_2$  formation that suggested a basic character of their low reactivity sites.

#### 3.2 Consumption of thiol-based antioxidants

The oxidative potential evaluated as Cys and GSH consumption after 24 h and as the DTT oxidation rate after 1 h are summarized in Fig. 2 and sorted by the k-means clustering algorithm as high, moderate, or low.  ${\rm SiO_2~NM-200~was}$  substantially unreactive in all assays, but site normalization indicated a moderate reactivity of its few reactive surface sites for Cys and DTT.

The most active ENM for Cys and GSH consumption was CuO, both per mass and per surface area, whereas active site normalization highlighted the oxidative potential of the zinc oxides, in particular NM-111. Carbon black and  ${\rm TiO_2}$  NM-105 consumed quite a large amount of these thiols per unit mass; however, they were in the low reactivity cluster when normalized by surface area or reactive site. Ceria exhibited low thiol consumption in GSH, but moderate Cys consumption per site. MWCNT NM-400 exhibited moderate reactivity for mass-normalized data, but low reactivity according to surface area normalization.

According to DTT consumption assay, carbon black, CuO, and MWCNT NM-400 were classified as highly oxidative according to the mass-based oxidation rate. However, for data normalized per area or reactive surface site, carbon black and MWCNT NM-400 fell into the moderate and low reactivity group, whereas  $\rm Mn_2O_3$ ,  $\rm Fe_2O_3$  and  $\rm CeO_2$  NM-212 were upgraded to highly reactive according to site normalization. Normalization per reactive surface site estimated the low reactivity of TiO\_2 NM-101 and TiO\_2 NM-105, which were, however, clustered as moderately reactive by mass normalization.

#### 3.3 Generation of ROS

ROS production by nanomaterials was analyzed using DCFH<sub>2</sub> assay and \*OH trapping with RNO. The results of DCFH<sub>2</sub> assay

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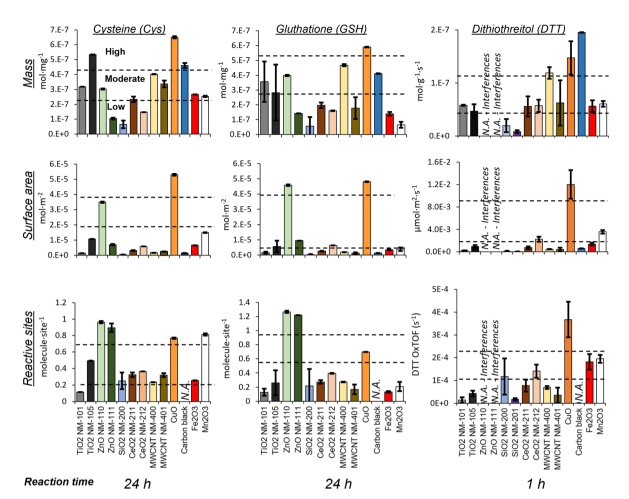


Fig. 2 Oxidative potential of the tested ENMs evaluated based on Cys (left) and GSH (center) 24 h consumption and the DTT 1 h oxidation rate (right) normalized by mass (top), surface area (middle) and reactive site (bottom). Averaged values (n = 3) with error bars indicating the standard deviation. Statistical clustering by the k-means algorithm according to reactivity (high-moderate-low) is indicated by horizontal dashed lines.

were expressed as the concentration of standard fluorescein diacetate (FDA) to obtain normalized data. DCFH2 depletion (Fig. 3a-l) was observed for carbon black, MWCNT NM-400, ZnO NM-111, CeO<sub>2</sub> NM-212, TiO<sub>2</sub> NM-105, and TiO<sub>2</sub> NM-101, which displayed positive and linear slopes, informing that ROS induction occurred in a dose-dependent manner. The significantly higher slope of NM-105 compared to NM-101 is indicative of higher rates of reactive oxygen species production by NM-105. Unlike CeO2 NM-212 and ZnO NM-111, CeO2 NM-211 and ZnO NM-110 show no ROS production. Four additional nanomaterials did not show any difference compared to the negative control for ROS: both silicon oxides (NM-200 and NM-201), CuO, and Fe<sub>2</sub>O<sub>3</sub>. Carbon-based ENMs and CuO were only tested between 0 and 12.5 µg mL<sup>-1</sup> following the interference test results (Table S4†). Thus, DCFH2 reaction was unsuitable for detecting ROS generated by CuO.

The results for RNO depletion by \*OH trapping are shown in Fig. 3m-o. Five nanomaterials appeared very active in producing \*OH: ZnO NM-110, ZnO NM-111, carbon black, CuO and MWCNT NM-400. These data highlight the elevated oxidative potential of zinc oxide reactive surface sites. Low reactivity was observed for TiO<sub>2</sub> NM-105, Fe<sub>2</sub>O<sub>3</sub> and Mn<sub>2</sub>O<sub>3</sub>,

while TiO<sub>2</sub> NM-101, both CeO<sub>2</sub> and MWCNT NM-401 performed similarly to the negative control. A large standard deviation for MWCNT NM-401 data was caused by non-stable dispersions in water, probably due to its high hydrophobicity.

#### Discussion 4

#### ENM's reactive surface sites

The reactive site surface density of the tested ENMs in this work ranged between 0.8 and 22 sites per nm2 (Table 1), consistent with literature data. 23,26,36-39 Methanol chemisorption was also performed at 50 °C for highly reactive CuO and ZnO NM-110 materials to avoid the formation of methoxy multilayers due to strong reactive interactions. The reactive site surface densities obtained at 100 °C were higher than those corresponding to a monolayer, which is in line with the observed CO<sub>2</sub> formation in MeOH-TPSR via decomposition of carbonates (high temperatures) and formates (low temperatures).40-42 The carbonates form at basic sites, while formates originate at redox sites. SiO2 NM-200 and NM-201 presented a low specific number of reactive sites, also resulting in low reactive sites surface density; these results explained the low reactivity observed in all assays

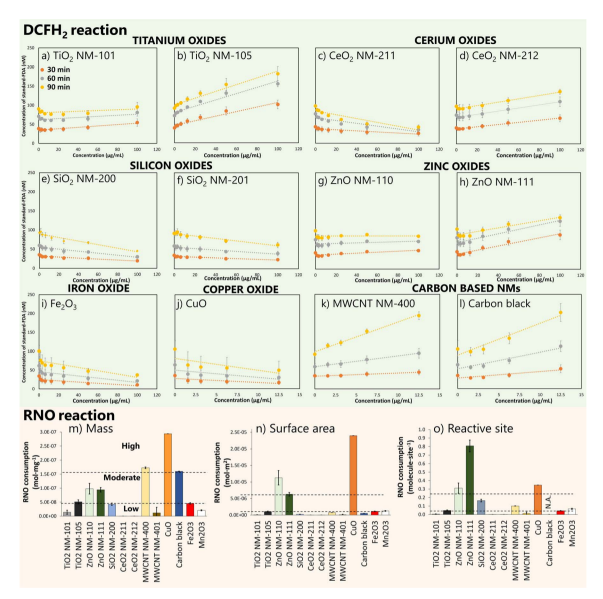


Fig. 3 ROS production estimation based on DCFH<sub>2</sub> assay (a)–(l) and \*OH trapping with RNO (m)–(o). Depleted DCFH<sub>2</sub> at different ENM concentrations measured by using a standard-FDA calibration curve for  $TiO_2$  NM-101 (a),  $TiO_2$  NM-105 (b),  $CeO_2$  NM-211 (c),  $CeO_2$  NM-212 (d),  $SiO_2$  NM-200 (e),  $SiO_2$  NM-201 (f),  $ZiO_3$  NM-110 (g),  $ZiO_3$  NM-111 (h),  $ZiO_3$  (i),  $ZiO_3$  (i),  $ZiO_3$  NM-400 (k) and carbon black (l). Carbon based NMs and CuO were only tested between 0 and 12.5  $ziO_3$  m mL<sup>-1</sup> according to the results of the interference test. RNO depletion is normalized per mass (m), per surface area (n) and per reactive site (o). Averaged values ( $ziO_3$  with error bars indicating the standard deviation.

and the absence of MeOH-TPSR products. We conclude that, although very large, the surface of these silica ENMs was mostly unreactive. The same behavior was observed for carbon nanotubes (MWCNT NM-400 and MWCNT NM-401). Despite their high specific surface area, they were actually two of the engineered nanomaterials with the lowest reactive site surface density, only slightly higher than that of silica oxides. Therefore, due to their low specific number of reactive sites and the absence of products in TPSR-MeOH, it is suggested that the reactivity observed for NM-400 in the liquid-phase reactions must be produced by reactive contaminants (Table S1†). 43,44

The chemisorption capacity of ZnO NM-111 seemed to be limited by the triethoxycaprylsilane coating, so that only about 5 reactive sites per nm<sup>2</sup> were available to interact with methanol,

whereas the uncoated ZnO NM-110 showed a smaller area but a very high surface density of reactive sites. Their reactive surface sites were similarly high consumers of GSH and Cys antioxidants in both nanomaterials. The site-normalized reactivity highlighted the oxidative capacity of NM-110 and NM-111 in Cys and GSH consumption tests.

#### 4.2 Nanomaterial ranking

To compare, rank and categorize the ENMs by reactivity, we adopted the K-means clustering algorithm (Fig. 4). CuO stands out as the ENM with the highest oxidative potential, as it was consistently classified as highly reactive across all antioxidant probe molecules, including \*OH radicals observed in RNO depletion. However, in the DCFH<sub>2</sub> probe reaction, the

Reactive profile

Main reactivity

Acid

Basic

Redox

Unreactive

Not available

MeOH

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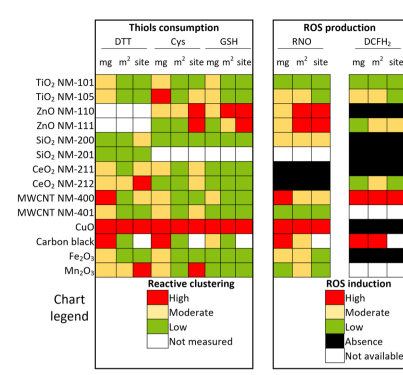


Fig. 4 Heat map for the 14 ENMs evaluated based on their intrinsic oxidative capacity to react with the thiol group in DTT, Cys, and GSH, or their production of ROS that are trapped by RNO and DCFH<sub>2</sub>, as well as based on their reactive profile obtained via methanol temperature programmed surface reaction. Clustering was performed by the k-means algorithm.

interference caused by this ENM prevented the detection of ROS. Following CuO, the subsequent nanomaterials in the reactivity scale were the two ZnO (NM-111 and NM-110) and carbon black. CuO and uncoated ZnO (NM-110) showed great similarities: they exhibited a high oxidizing character, form CO<sub>2</sub> in MeOH-TPSR, but did not react against DCFH2; in addition, their consumption of RNO suggest that they form \*HO.

Reported ESR and FRAS (ferric reducing ability of the serum) assay data for ZnO NM-110 17,18 confirm the absence of ROS production. Moreover, a high depletion of cyclic hydroxylamine spin probe 1-hydroxy-3-carboxy-pyrrolidine (CPH) and nitrone spin trap 5,5-dimethyl-1-pyrroline-N-oxide (DMPO) molecules, sensitive to different ROS, was reported for this material compared to the blank, and surface normalized data provided further evidence of the oxidative capacity. 19,35 The FRAS assay revealed high oxidation levels caused by ZnO NM-110, in line with its reactivity per site in GSH, Cys, and RNO reactions. Similar to this uncoated ZnO, silane coated ZnO (NM-111), one of the most oxidizing ENMs in the series, tested positive in FRAS experiments, 35 as well as CPH and DMPO probes. 19 The different behavior of NM-110 vs. NM-111 in the DCFH<sub>2</sub> probe reaction suggests that both nanomaterials have different oxidation mechanisms, being NM-111 a nanomaterial with higher ROS induction capacity.

Lastly, carbon black was utilized as a positive control in DCFH<sub>2</sub> assay due to its oxidative capacity evidenced by the high ROS production, high DTT depletion and high sensitivity for FRAS assay reported in the literature. 15,45-48 This high reactivity of CB agreed with the results obtained in this study by mass normalization, where a high consumption of GSH, Cys, DTT,

and RNO, and especially a high slope in the DCFH2 reaction were obtained, according to its capacity to induce ROS.

These data clustered SiO<sub>2</sub> NM-200, Fe<sub>2</sub>O<sub>3</sub>, CeO<sub>2</sub> NM-211, CeO2 NM-212, TiO2 NM-101 and TiO2 NM-105 in a lower reactivity group. It is worth noting the similarity between the two titanias reflected by MeOH-TPSR, mainly when compared to the higher oxidizing capacity of ceria NM-211 vs. NM-212. The different oxidative potential for CeO<sub>2</sub> ENMs is supported by XPS results in the literature, which detected a higher percentage of Ce(III) on the surface of NM-211 (22%) vs. NM-212 (14%).49 Concerning SiO<sub>2</sub> NM-200, it exhibited low reactivity in all reactions and metrics, except for DTT consumption per active site. The limited number of reactive sites on the surface leads to an artificially high value of molecule consumption per site, even though the nanomaterial is, in fact, classified as unreactive. There is no substantial consumption of any molecule, no measurable ROS formation, and no products observed in MeOH-TPSR whatsoever. This evidence underlines the relevance of the binomial constituted by (a) the number of reactive surface sites and (b) the reactivity of such sites.

The results of oxidative potential per reactive surface site obtained with the different probe molecules were compared utilizing Pearson's correlation coefficient (Fig. S2†). The comparison of probe molecules enabled an exploration of the underlying reaction mechanisms governing their consumption. The significant Pearson correlation coefficient between GSH, Cys, DTT and RNO (0.91-0.80) suggests a shared oxidation mechanism. As outlined in previous studies,50-52 this mechanism primarily involves the oxidation of thiol (-SH) groups, which are critical functional components in GSH, Cys, and DTT.

During this process, thiols undergo a single-electron oxidation, resulting in the formation of sulfur-centered radicals that later combine to form disulfides. Other authors, such as Jiang and colleagues, attributed the oxidation of DTT to the presence of transition metals, like Cu, which induce Fenton-like reactions, producing H<sub>2</sub>O<sub>2</sub> and ultimately generating \*OH radicals.<sup>53</sup> Consequently, for ENMs like CuO, the consistent data showing oxidation of -SH groups and generation of \*OH radicals that deplete RNO are in perfect agreement. Additionally, the surface of CuO particles is predominantly reduced to either Cu<sub>2</sub>O or metallic copper, as demonstrated by Yiwen Wang et al.54 For ZnO and TiO2, thiol adsorption has been reported in the literature; however, the acidic surface of TiO2 may not oxidize DTT, while the highly reactive surface sites of ZnO result in significant depletion of -SH antioxidants. In contrast, CeO<sub>2</sub>, with its oxidative surface sites, exhibits only slight oxidation of antioxidants, likely due to the absence of -SH adsorption on its surface. 55 However, DCFH2, which theoretically should provide results similar to RNO, does not correlate with the obtained dataset for ROS generation (RNO) and antioxidant consumption (DTT, Cys, and GSH). This discrepancy is primarily attributed to interference from carbon-based and CuO ENMs, while showing no reactivity in the presence of Zn2+, as evidenced by the low reactivity of ZnO NM-110.

# 4.3 Correlation between surface reactivity and *in vitro* toxicity data: a new dosimetry approach

The results for in chemico reactivity described in this work and literature cytotoxicity data suggest that enhanced reactivity in the tested ENMs correlates with increased cytotoxicity in pulmonary cells post 24 h exposure. Notably, CuO, ZnO NM-110, and ZnO NM-111, identified as highly oxidizing, markedly reduced cell viability in primary pulmonary cell lines. Specifically, CuO emerged as the most cytotoxic ENM, significantly impacting A549 cell viability at concentrations of 5-10 μg mL<sup>-1</sup> across MTT, NRU, CFA, and IL-8 assays, <sup>56-59</sup> while zinc oxide NM-110 exhibited significant effects on cell viability at  $6.4-50~\mu g~mL^{-1}$  for PMA treated THP-1,60,61 128  $\mu g~mL^{-1}$  for Calu-3  $^{62}$  and 48–75  $\mu g\ mL^{-1}$  for A549,  $^{63-65}$  and even lower concentrations are required for ZnO NM-111: 37 μg mL<sup>-1</sup> in A549 65 and 32–128 μg mL<sup>-1</sup> in Calu-3.62 Exposure to CuO, ZnO NM-110 and ZnO NM-111 may result in different modes of action, including those derived from ion release into the medium, as they are known to dissolve. For example, in a recent in vivo study in rats, ZnO NM-110 was compared to highly soluble ZnSO<sub>4</sub>, concluding that ZnO nanoforms most likely exhibit their effects by zinc ions, since the exposure to zinc sulfate had similar effects. However, they are highly reactive not only in the consumption of probe molecules during dissolution, but also in methanol-TPSR, which is performed with the powdered material, suggesting that surface reactivity may provide very relevant complementary information for toxicity evaluation.66 Indeed, it is remarkable that Cronholm et al. observed significant cell and DNA damage caused by CuO nanoparticles but no adverse effects from highly soluble CuCl<sub>2</sub> salts.<sup>67</sup> In agreement with this, Wang et al.

concluded that dissolved Cu<sup>2+</sup> ions contributed to less than half of the total effects caused by CuO NPs, including ROS generation and DNA damage.<sup>68</sup> Thus, we identify surface reactivity as one of the key factors driving the electronic imbalance within cells, ultimately resulting in cellular damage.

Exposure to CuO, ZnO NM-110 and ZnO NM-111 may result in different modes of action. In contrast, exposure to nonsoluble ENMs with moderate or low oxidative potential causes less pronounced effects in pulmonary cells. TiO2 NM-101 produces no significant alteration in BEAS-2B and A549 cell viability.69-71 Similarly, TiO2 NM-105 shows no notable impact on A549 and THP-1 cell lines. 72,73 Cerium oxide NM-212 does not disrupt LDH, cytokine release, or Alamar blue assay markers in A549 and PMA-treated THP-1 cells.73,74 SiO2 NM-200 cytotoxicy has been tested for BEAS-2B,70 Calu-3 62 and PMA treated THP-1,60 but significant effects were on; v obtained with the MTS test on THP-1 at a concentration of 50 μg mL<sup>-1</sup>. An MTT assay in A549 and BEAS-2B indicates no impact on cell viability after 24 h exposure to 200 µg mL<sup>-1</sup> Fe<sub>2</sub>O<sub>3</sub>.75 Conversely, carbonaceous materials exhibit varied effects. Exposure of A549 cells to 50 μg mL<sup>-1</sup> carbon black induces ROS and IL-8 release without affecting cell viability, whereas no cytotoxicity is observed in Calu-3 at this concentration. MWCNT NM-400 at 100 μg mL<sup>-1</sup> in A549 prompts LDH release and reduces cell viability, and at 80  $\mu g \text{ mL}^{-1}$  in PMA-treated THP-1 it triggers IL-8 release. MWCNT NM-400 at 100  $\mu g \ mL^{-1}$  in A549 induced LDH release and decreased cell viability, 77 and at 80 μg mL<sup>-1</sup> in PMA treated THP-1 it triggers IL-8 release.78 The toxicity of MWCNT NM-400 produced by surface reactivity could be explained due to their impurities;44 however, it is not expected to initiate reactions leading to toxicity effects. 43,77,79 Additionally, based on previous studies, these impurities are not released in the medium.79 Finally, when MWCNT NM-401 is exposed to PMA treated THP-1 and A549, the main effect observed is IL-8 release at 40  $\mu g$  mL<sup>-1</sup> for both cell lines.78

Understanding the interplay between toxicity and reactivity poses a complex challenge, particularly without complementary assays addressing biotransformation, bioaccumulation, corona protein formation, and related factors. Moreover, the variety of reactivity and toxicity testing methods and conditions employed in different studies hampers a comprehensive analysis. Taking this into account, available toxicity data of TiO<sub>2</sub> NM-101, TiO<sub>2</sub> NM-105, ZnO NM-110, ZnO NM-111, SiO<sub>2</sub> NM-200, SiO<sub>2</sub> NM-201, CeO<sub>2</sub> NM-211, CeO<sub>2</sub> NM-212, CuO and Fe<sub>2</sub>O<sub>3</sub> evaluated by LDH and WST-1 assays with A549 and dTHP-1 cell lines, collected in Table S3,† were selected owing to their relative uniformity (same two assays and two cell lines), same exposure time (24 h) and the extensive range of tested ENMs (11 out of the 14 studied ENMs).80 These data are expressed as no observed adverse effect concentration (NOAEC) of reactive sites concentration ( $\mu$ mol L<sup>-1</sup>), our new proposed dose metric, to calculate Spearman's coefficients summarized in Table 2, aiming to evaluate possible correlations between oxidative reactivity against the five specific probe molecules used in this work and the reported cell viability. DCFH<sub>2</sub> depletion rates were measured at 50 μg mL<sup>-1</sup>

Table 2 Spearman correlation between oxidative potential per reactive surface site measured by DCFH2, RNO, DTT, Cys and GSH probe reactions in this work and bibliographic in vitro toxicity data assessed by LDH and WST-1 tests in A549 and dTHP-1 cell lines after 24 h exposure from Table S3. Raw toxicological data extracted from Alcolea-Rodriguez et al.80

		RNO	GSH	Cys	DTT	$DCFH_2$
A549 A549 dTHP-1 dTHP-1	LDH WST-1 LDH WST1	-0.55 -0.57 -0.55 -0.57	-0.54 $-0.58$ $-0.54$ $-0.59$	-0.56 $-0.59$ $-0.56$ $-0.59$	-0.53 -0.53 -0.53 -0.53	-0.16 $-0.21$ $-0.16$ $-0.21$
	n	9	9	9	8	9

of ENMs, and normalized by mass, area and sites (Table S2†). Oxidative reactivity in cell-free environments and in vitro toxicity values exhibit negative Spearman's coefficients, indicating an inverse proportionality. This means that ENMs with higher reactivity require lower concentrations to induce cell damage. DTT, GSH, Cys and RNO reactions showed similar moderate Spearman correlation factors for in vitro adverse effects; the least predictive marker is the oxidation rate of DCFH<sub>2</sub>, which demonstrates a lower prognostic efficacy for A549 and dTHP-1 cell viability.

This new dosimetry shares the limitations of other normalizations when applied to soluble nanomaterials: ion secretion results in mass loss (leading to a reduction in nanomaterial concentration) and surface area alteration (with the exposed surface area remaining unknown). Consequently, if ions originating from the ENM are dissolved in the reaction media, in addition to interference problems, it is not possible to ascertain the state of reactive sites. Therefore, for cytotoxic ENMs such as CuO and ZnO (NM-110 and N-111) that are partially soluble, it is challenging to determine their true exposure levels based on reactive surface site concentrations. Other nanomaterials not exhibiting solubility in the literature, such as titania, Fe<sub>2</sub>O<sub>3</sub>, and MWCNTs,79 exhibit significant disparities in their actual exposure of reactive sites for a similar concentration, e.g., 100 µg mL<sup>-1</sup> of TiO<sub>2</sub> NM-101 presents a high number of reactive sites (280 µmol sites per L) compared to its analog NM-105 (110 µmol sites per L) or to other nanomaterials like Fe<sub>2</sub>O<sub>3</sub> (110 μmol sites per L). This phenomenon highlights the relevance of using reactive site concentration as an alternative to gravimetric concentration, as comparisons in terms of mass that do not account for the exposure of each nanomaterial. However, when using volumetric surface area (cm<sup>2</sup> mL<sup>-1</sup>), the Spearman correlation factors for GSH, Cys, and DTT depletion increase to approximately 0.80 (data not shown) compared to cell-based parameters, indicating a stronger correlation. This trend arises because the most reactive ENMs, ZnO and CuO, have low BET surface areas. We expect that, since not all physical areas correspond to chemical areas, expanding the study to include a broader cohort of more spherical ENMs will likely reduce the Spearman correlation. The current high correlation is due to the limited number of cytotoxic ENMs in this study.

#### Conclusions and outlook 5

This study reinforces the significance of reactivity by introducing new probe-molecule methods to evaluate reactivity that produce results comparable with established assays, such as DCFH<sub>2</sub>. Our measurements aimed to establish that (1) methods such as methanol-TPSR, antioxidant consumption, and ROS generation measured by RNO and DCFH2 can elucidate the surface reactivity of ENMs; (2) ENMs with identical compositions can exhibit varying oxidative capacities; (3) the number and nature of reactive surface sites significantly influence their reactivity and, consequently, their toxicity; and (4) a site-based dose metric is proposed for establishing the correct effective dose of non-soluble ENMs.

The reactivity screening conducted in this study with 14 ENMs using 6 probe reactions underscores the oxidizing potential of nanomaterials such as CuO, ZnO NM-110, ZnO NM-111, and carbon black, whereas nanomaterials such as MWCNT NM-400, TiO<sub>2</sub> NM-105, CeO<sub>2</sub> NM-212, SiO<sub>2</sub> NM-200, TiO<sub>2</sub> NM-101, CeO<sub>2</sub> NM-211, MWCNT NM-401, Mn<sub>2</sub>O<sub>3</sub>, and Fe<sub>2</sub>O<sub>3</sub> are categorized into groups with moderate to low oxidative reactivity. Additionally, distinct oxidative mechanisms were proposed for each ENM. For ZnO, its reactivity and toxicity have been reported in the literature as dependent on the release of Zn<sup>2+</sup> ions. In contrast, CuO's surface reactivity emerged as the primary mechanism of oxidation and toxicity, as Cu salts, in comparison, failed to induce similar toxicity or reactivity effects according to the literature. These findings reflect the complex mechanisms ocurring at the nanolevel, and highlight the multifactorial mode of action of ENMs in biological systems.

The proposed methodology offers a deep understanding of the reactivity of ENMs. It has versatile applications, among which a novel approach is suggested for normalizing toxicity data. This method considers the reactive surface sites rather than solely relying on the mass or the total surface area of ENMs. Such an approach could significantly enhance the field of nanotoxicology in the future, providing a more accurate risk assessment of ENMs.

## Data availability

The data supporting this article have been included as part of the ESI.†

### Conflicts of interest

The authors declare no competing financial interest.

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#### References

- 1 I. Khan, K. Saeed and I. Khan, Nanoparticles: properties, applications and toxicities, *Arabian J. Chem.*, 2019, 12(7), 908–931, DOI: 10.1016/j.arabjc.2017.05.011.
- 2 C. Kinnear, T. L. Moore, L. Rodriguez-Lorenzo, B. Rothen-Rutishauser and A. Petri-Fink, Form Follows Function: Nanoparticle Shape and Its Implications for Nanomedicine, *Chem. Rev.*, 2017, 117(17), 11476–11521, DOI: 10.1021/acs.chemrev.7b00194.
- 3 M. A. Bañares, A. Haase, L. Tran, et al., CompNanoTox2015: novel perspectives from a European conference on computational nanotoxicology on predictive nanotoxicology, *Nanotoxicology*, 2017, **11**(7), 839–845, DOI: **10.1080/17435390.2017.1371351.**
- 4 F. Polo-Garzon, Z. Bao, X. Zhang, W. Huang and Z. Wu, Surface Reconstructions of Metal Oxides and the Consequences on Catalytic Chemistry, *ACS Catal.*, 2019, 9(6), 5692–5707, DOI: 10.1021/acscatal.9b01097.
- 5 Z. Wu, A. K. P. Mann, M. Li and S. H. Overbury, Spectroscopic investigation of surface dependent acid base property of ceria nanoshapes, *J. Phys. Chem. C*, 2015, **119**(13), 7340–7350, DOI: **10.1021/ACS.JPCC.5B00859**.
- 6 Z. Wu, M. Li, J. Howe, H. M. Meyer and S. H. Overbury, Probing defect sites on CeO<sub>2</sub> nanocrystals with well-defined surface planes by Raman spectroscopy and O<sub>2</sub> adsorption, *Langmuir*, 2010, 26(21), 16595–16606, DOI: 10.1021/ LA101723W.
- 7 M. A. Bañares, V. Alcolea-Rodriguez and R. Portela, A catalytic perspective to nanomaterials reactivity-based toxicity; implications for single- and multiple-component nanomaterials (nanocomposites), *NanoImpact*, 2025, 37, 100542, DOI: 10.1016/J.IMPACT.2025.100542.
- 8 T. Stobernack, N. Dommershausen, V. Alcolea Rodríguez, *et al.*, Advancing Nanomaterial Toxicology Screening Through Efficient and Cost Effective Quantitative Proteomics, *Small Methods*, 2024, 8(12), 2400420, DOI: 10.1002/smtd.202400420.
- 9 S. Schmeisser, A. Miccoli, M. von Bergen, *et al.*, New approach methodologies in human regulatory toxicology not if, but how and when!, *Environ. Int.*, 2023, **178**, 108082, DOI: **10.1016/j.envint.2023.108082**.
- 10 S. H. Doak, M. J. D. Clift, A. Costa, et al., The Road to Achieving the European Commission's Chemicals Strategy for Nanomaterial Sustainability—A PATROLS Perspective on New Approach Methodologies, Small, 2022, 18(17), DOI: 10.1002/smll.202200231.
- 11 S. Halappanavar, J. D. Ede, J. A. Shatkin and H. F. Krug, A systematic process for identifying key events for advancing the development of nanomaterial relevant adverse outcome pathways, *NanoImpact*, 2019, **15**, DOI: **10.1016/j.impact.2019.100178**.
- 12 S. Halappanavar, S. Van Den Brule, P. Nymark, *et al.*, Adverse outcome pathways as a tool for the design of testing strategies to support the safety assessment of emerging

- advanced materials at the nanoscale, *Part. Fibre Toxicol.*, 2020, 17(1), DOI: 10.1186/s12989-020-00344-4.
- 13 F. C. Simeone and A. L. Costa, Assessment of cytotoxicity of metal oxide nanoparticles on the basis of fundamental physical-chemical parameters: a robust approach to grouping, *Environ. Sci.: Nano*, 2019, 6(10), 3102–3112, DOI: 10.1039/C9EN00785G.
- 14 D. Ag Seleci, G. Tsiliki, K. Werle, et al., Determining nanoform similarity via assessment of surface reactivity by abiotic and in vitro assays, NanoImpact, 2022, 26, 100390, DOI: 10.1016/j.impact.2022.100390.
- 15 M. Boyles, F. Murphy, W. Mueller, *et al.*, Development of a standard operating procedure for the DCFH<sub>2</sub>-DA acellular assessment of reactive oxygen species produced by nanomaterials, *Toxicol. Mech. Methods*, 2022, 32(6), 439–452, DOI: 10.1080/15376516.2022.2029656.
- 16 M. D. Driessen, S. Mues, A. Vennemann, *et al.*, Proteomic analysis of protein carbonylation: a useful tool to unravel nanoparticle toxicity mechanisms, *Part. Fibre Toxicol.*, 2015, 12(1), 1–18, DOI: 10.1186/s12989-015-0108-2.
- 17 D. Ag Seleci, G. Tsiliki, K. Werle, *et al.*, Determining nanoform similarity via assessment of surface reactivity by abiotic and in vitro assays, *NanoImpact*, 2022, **26**, 100390, DOI: **10.1016/j.impact.2022.100390**.
- 18 M. Boyles, F. Murphy, W. Mueller, *et al.*, Development of a standard operating procedure for the DCFH<sub>2</sub>-DA acellular assessment of reactive oxygen species produced by nanomaterials, *Toxicol. Mech. Methods*, 2022, 32(6), 439–452, DOI: 10.1080/15376516.2022.2029656.
- 19 M. D. Driessen, S. Mues, A. Vennemann, *et al.*, Proteomic analysis of protein carbonylation: a useful tool to unravel nanoparticle toxicity mechanisms, *Part. Fibre Toxicol.*, 2015, **12**(1), 1–18, DOI: **10.1186/s12989-015-0108-2**.
- 20 C. Mungsuk, S. Yommee, S. Supothina and P. Chuaybamroong, Solar photocatalytic degradation of carbendazim in water using TiO<sub>2</sub> particle- and sol-gel dipcoating filters, *Results Eng.*, 2023, 19, 101348, DOI: 10.1016/ j.rineng.2023.101348.
- 21 B. Ohtani, O. O. Prieto-Mahaney, D. Li and R. Abe, What is Degussa (Evonic) P25? Crystalline composition analysis, reconstruction from isolated pure particles and photocatalytic activity test, *J. Photochem. Photobiol.*, *A*, 2010, 216(2–3), 179–182, DOI: 10.1016/j.jphotochem.2010.07.024.
- 22 Joint Meeting of the Chemicals Committee and the Working Party on Chemicals, Pesticides and Biotechnology, Titanium Dioxide: Summary of the Dossier, Series on the Safety of Manufactured Nanomaterials, No. 73, 16th Meeting of the Working Party on Manufactured Nanomaterials, 2016.
- 23 J. M. Tatibouët, Methanol oxidation as a catalytic surface probe, *Appl. Catal.*, *A*, 1997, **148**(2), 213–252, DOI: **10.1016**/**S0926-860X**(96)00236-0.
- 24 Y. Wu, F. Gao, H. Wang, L. Kovarik, B. Sudduth and Y. Wang, Probing Acid–Base Properties of Anatase TiO<sub>2</sub> Nanoparticles with Dominant {001} and {101} Facets Using Methanol Chemisorption and Surface Reactions, *J. Phys. Chem. C*, 2021, 125(7), 3988–4000, DOI: 10.1021/acs.ipcc.0c11107.

- 25 J. M. Jehng, I. E. Wachs, G. S. Patience and Y. M. Dai, Experimental methods in chemical engineering: temperature programmed surface reaction spectroscopy— TPSR, Can. J. Chem. Eng., 2021, 99(2), 423-434, DOI: 10.1002/cjce.23913.
- 26 V. Alcolea-Rodriguez, R. Portela, V. Calvino-Casilda and M. A. Bañares, In chemico methodology for engineered nanomaterial categorization according to number, nature and oxidative potential of reactive surface sites, Environ. Sci.: Nano, 2024, 11(9), 3744-3760, DOI: 10.1039/ D3EN00810J.
- 27 A. Brunelli, A. Serrano-Lotina, M. A. Bañares, et al., Safe-bydesign assessment of an SiO<sub>2</sub>@ZnO multi-component nanomaterial used in construction, Environ. Sci.: Nano, 2024, DOI: 10.1039/D4EN00352G.
- 28 K. Rasmussen, J. Mast, P. J. Temmerman, E. Verleysen, N. Waegeneers and F. Steen, Multi-Walled Carbon Nanotubes, NM-400, NM-401, NM-402, NM-403: Characterisation and Physico-Chemical Properties, in NMseries of Representative Manufactured Nanomaterials, 2014, DOI: 10.2788/10753.
- 29 B. Hellack, C. Nickel, C. Albrecht, et al., Analytical methods to assess the oxidative potential of nanoparticles: a review, Environ. Sci.: Nano, 2017, DOI: 10.1039/c7en00346c.
- 30 D. Garrido Ruiz, A. Sandoval-Perez, A. V. Rangarajan, E. L. Gunderson and M. P. Jacobson, Cysteine Oxidation in Proteins: Structure, Biophysics, and Simulation, Biochemistry, 2022, 61(20), 2165-2176, DOI: 10.1021/ acs.biochem.2c00349.
- 31 I. Kraljić and C. N. Trumbore, p-Nitrosodimethylaniline as an OH Radical Scavenger in Radiation Chemistry, J. Am. Chem. Soc., 1965, 87(12), 2547-2550, DOI: 10.1021/ JA01090A004/ASSET/JA01090A004.FP.PNG\_V03.
- 32 G. Mancardi, A. Mikolajczyk, V. K. Annapoorani, et al., A computational view on nanomaterial intrinsic and extrinsic features for nanosafety and sustainability, Mater. Today, 2023, 344-370, DOI: 10.1016/ 67, j.mattod.2023.05.029.
- 33 N. Jeliazkova, E. Bleeker, R. Cross, et al., How can we justify grouping of nanoforms for hazard assessment? Concepts and tools to quantify similarity, NanoImpact, 2022, 25, 100366, DOI: 10.1016/j.impact.2021.100366.
- 34 K. Paunovska, D. Loughrey, C. D. Sago, R. Langer and J. E. Dahlman, Using Large Datasets to Understand Nanotechnology, Adv. Mater., 2019, 31(43), 1902798, DOI: 10.1002/adma.201902798.
- 35 A. Bahl, B. Hellack, M. Wiemann, et al., Nanomaterial categorization by surface reactivity: a case study comparing 35 materials with four different test methods, NanoImpact, 2020, 19, 100234, DOI: 10.1016/j.impact.2020.100234.
- 36 L. E. Briand, Investigation of the Nature and Number of Surface Active Sites of Supported and Bulk Metal Oxide Catalysts through Methanol Chemisorption, Metal Oxides, 375-412, DOI: CRC Press, 2005, pp. 9781420028126-15.

- 37 I. E. Wachs, Number of surface sites and turnover frequencies for oxide catalysts, J. Catal., 2022, 405, 462-472, DOI: 10.1016/j.jcat.2021.12.032.
- 38 J. M. Jehng, I. E. Wachs and M. Ford, Temperature-Programmed (TP) Techniques, in Springer Handbook of Advanced Catalyst Characterization, Springer Handbooks, 2023, ch. 45, vol. 4, pp. 1005-1029, DOI: 10.1007/978-3-031-07125-6 45.
- 39 L. E. Briand, A. M. Hirt and I. E. Wachs, Quantitative determination of the number of surface active sites and the turnover frequencies for methanol oxidation over metal oxide catalysts: application to bulk metal molybdates and pure metal oxide catalysts, J. Catal., 2001, 202(2), 268-278, DOI: 10.1006/jcat.2001.3289.
- 40 A. Davydov, Molecular Spectroscopy of Oxide Catalyst Surfaces, Wiley, 2003, DOI: 10.1002/0470867981.
- 41 S. E. Collins, L. E. Briand, L. A. Gambaro, M. A. Baltanás and A. L. Bonivardi, Adsorption and Decomposition of Methanol on Gallium Oxide Polymorphs, J. Phys. Chem. C, 2008, 112(38), 14988-15000, DOI: 10.1021/jp801252d.
- 42 P. Bazin, S. Thomas, O. Marie and M. Daturi, New insights into the methanol oxidation mechanism over Au/CeO2 catalyst through complementary kinetic and FTIR operando SSITKA approaches, Catal. Today, 2012, 182(1), 3-11, DOI: 10.1016/j.cattod.2011.10.001.
- 43 C. Darne, A. Desforges, N. Berrada, et al., A non-damaging purification method: decoupling the toxicity of multiwalled carbon nanotubes and their associated metal impurities, Environ. Sci.: Nano, 2019, 6(6), 1852-1865, DOI: 10.1039/C8EN01276H.
- 44 J. H. E. Arts, M. A. Irfan, A. M. Keene, et al., Case studies putting the decision-making framework for the grouping and testing of nanomaterials (DF4nanoGrouping) into practice, Regul. Toxicol. Pharmacol., 2016, 76, 234-261, DOI: 10.1016/j.yrtph.2015.11.020.
- 45 E. Koike and T. Kobayashi, Chemical and biological oxidative effects of carbon black nanoparticles, Chemosphere, 2006, **65**(6), 946-951, DOI: 10.1016/ j.chemosphere.2006.03.078.
- 46 A. Bahl, B. Hellack, M. Wiemann, et al., Nanomaterial categorization by surface reactivity: a case study comparing 35 materials with four different test methods, NanoImpact, 2020, 19, 100234, DOI: 10.1016/j.impact.2020.100234.
- 47 J. J. Sauvain, M. J. Rossi and M. Riediker, Comparison of three acellular tests for assessing the oxidation potential of nanomaterials, Aerosol Sci. Technol., 2013, 47(2), 218-227, DOI: 10.1080/02786826.2012.742951.
- 48 F. Murphy, N. R. Jacobsen, E. Di Ianni, et al., Grouping MWCNTs based on their similar potential to cause pulmonary hazard after inhalation: a case-study, Part. Fibre Toxicol., 2022, 19(1), 1-23, DOI: 10.1186/s12989-022-00487-6.
- 49 J. Keller, W. Wohlleben, L. Ma-Hock, et al., Time course of lung retention and toxicity of inhaled particles: short-term exposure to nano-ceria, Arch. Toxicol., 2014, 88(11), 2033-2059, DOI: 10.1007/s00204-014-1349-9.
- 50 A. L. Kramer, S. Dorn, A. Perez, et al., Assessing the oxidative potential of PAHs in ambient PM2.5 using the DTT

- consumption assay, *Environmental Pollution*, 2021, **285**(May), 117411, DOI: 10.1016/j.envpol.2021.117411.
- 51 J. J. Sauvain, S. Deslarzes and M. Riediker, Nanoparticle reactivity toward dithiothreitol, *Nanotoxicology*, 2008, 2(3), 121–129, DOI: 10.1080/17435390802245716.
- 52 J. G. Charrier and C. Anastasio, On dithiothreitol (DTT) as a measure of oxidative potential for ambient particles: evidence for the importance of soluble transition metals, *Atmos. Chem. Phys.*, 2012, 12(19), 9321–9333, DOI: 10.5194/acp-12-9321-2012.
- 53 H. Jiang, C. M. Sabbir Ahmed, A. Canchola, J. Y. Chen and Y. H. Lin, Use of dithiothreitol assay to evaluate the oxidative potential of atmospheric aerosols, *Atmosphere*, 2019, **10**(10), 1–21, DOI: **10**.3390/atmos10100571.
- 54 Y. Wang, J. Im, J. W. Soares, D. M. Steeves and J. E. Whitten, Thiol Adsorption on and Reduction of Copper Oxide Particles and Surfaces, *Langmuir*, 2016, 32(16), 3848–3857, DOI: 10.1021/acs.langmuir.6b00651.
- 55 O. C. Grimm, R. M. D. S. Somaratne, Y. Wang, S. Kim and J. E. Whitten, Thiol adsorption on metal oxide nanoparticles, *Phys. Chem. Chem. Phys.*, 2021, 23(14), 8309–8317, DOI: 10.1039/D1CP00506E.
- 56 E. Moschini, M. Gualtieri, M. Colombo, U. Fascio, M. Camatini and P. Mantecca, The modality of cell-particle interactions drives the toxicity of nanosized CuO and TiO<sub>2</sub> in human alveolar epithelial cells, *Toxicol. Lett.*, 2013, 222(2), 102–116, DOI: 10.1016/j.toxlet.2013.07.019.
- 57 A. Semisch, J. Ohle, B. Witt and A. Hartwig, Cytotoxicity and genotoxicity of nano and microparticulate copper oxide: role of solubility and intracellular bioavailability, *Part. Fibre Toxicol.*, 2014, **11**(1), 10, DOI: **10.1186/1743-8977-11-10**.
- 58 I. Perelshtein, A. Lipovsky, N. Perkas, A. Gedanken, E. Moschini and P. Mantecca, The influence of the crystalline nature of nano-metal oxides on their antibacterial and toxicity properties, *Nano Res.*, 2015, 8(2), 695–707, DOI: 10.1007/s12274-014-0553-5.
- 59 E. Moschini, G. Colombo, G. Chirico, G. Capitani, I. Dalle-Donne and P. Mantecca, Biological mechanism of cell oxidative stress and death during short-term exposure to nano CuO, *Sci. Rep.*, 2023, 13(1), 2326, DOI: 10.1038/s41598-023-28958-6.
- 60 T. Brzicova, E. Javorkova, K. Vrbova, et al., Molecular responses in THP-1 macrophage-like cells exposed to diverse nanoparticles, Nanomaterials, 2019, 9(5), 1–19, DOI: 10.3390/nano9050687.
- 61 R. Safar, Z. Doumandji, T. Saidou, et al., Cytotoxicity and global transcriptional responses induced by zinc oxide nanoparticles NM 110 in PMA-differentiated THP-1 cells, *Toxicol. Lett.*, 2019, 308, 65–73, DOI: 10.1016/j.toxlet.2018.11.003.
- 62 L. Farcal, F. T. Andón, L. Di Cristo, *et al.*, Comprehensive in vitro toxicity testing of a panel of representative oxide nanomaterials: first steps towards an intelligent testing strategy, *PLoS One*, 2015, **10**(5), 1–34, DOI: **10.1371**/ **journal.pone.0127174**.
- 63 A. Precupas, D. Gheorghe, A. Botea-Petcu, et al., Thermodynamic Parameters at Bio-Nano Interface and

- Nanomaterial Toxicity: A Case Study on BSA Interaction with ZnO, SiO<sub>2</sub>, and TiO<sub>2</sub>, *Chem. Res. Toxicol.*, 2020, 33(8), 2054–2071, DOI: 10.1021/acs.chemrestox.9b00468.
- 64 N. E. Yamani, A. R. Collins, E. Rundén-Pran, *et al.*, In vitro genotoxicity testing of four reference metal nanomaterials, titanium dioxide, zinc oxide, cerium oxide and silver: towards reliable hazard assessment, *Mutagenesis*, 2017, 32(1), 117–126, DOI: 10.1093/mutage/gew060.
- 65 Y. N. El, E. Mariussen, M. Gromelski, *et al.*, Hazard identification of nanomaterials: in silico unraveling of descriptors for cytotoxicity and genotoxicity, *Nano Today*, 2022, **46**, 101581, DOI: **10.1016/j.nantod.2022.101581**.
- 66 L. M. Gilbertson, E. M. Albalghiti, Z. S. Fishman, et al., Shape-Dependent Surface Reactivity and Antimicrobial Activity of Nano-Cupric Oxide, Environ. Sci. Technol., 2016, 50(7), 3975–3984, DOI: 10.1021/ACS.EST.5B05734/ASSET/ IMAGES/LARGE/ES-2015-05734J 0005.JPEG.
- 67 P. Cronholm, H. L. Karlsson, J. Hedberg, et al., Intracellular Uptake and Toxicity of Ag and CuO Nanoparticles: A Comparison between Nanoparticles and Their Corresponding Metal Ions, Small, 2013, 9(7), 970–982, DOI: 10.1002/SMLL.201201069.
- 68 Z. Wang, N. Li, J. Zhao, J. C. White, P. Qu and B. Xing, CuO nanoparticle interaction with human epithelial cells: cellular uptake, location, export, and genotoxicity, *Chem. Res. Toxicol.*, 2012, 25(7), 1512–1521, DOI: 10.1021/TX3002093/SUPPL\_FILE/TX3002093\_SI\_001.PDF.
- 69 W. Thongkam, K. Gerloff, D. van Berlo, C. Albrecht and R. P. F. Schins, Oxidant generation, DNA damage and cytotoxicity by a panel of engineered nanomaterials in three different human epithelial cell lines, *Mutagenesis*, 2017, 32(1), 105–115, DOI: 10.1093/mutage/gew056.
- 70 T. Cervena, A. Rossnerova, T. Zavodna, *et al.*, Testing strategies of the in vitro micronucleus assay for the genotoxicity assessment of nanomaterials in beas-2b cells, *Nanomaterials*, 2021, 11(8), 1–20, DOI: 10.3390/nano11081929.
- 71 A. Zijno, D. Cavallo, G. Di Felice, *et al.*, Use of a common European approach for nanomaterials' testing to support regulation: a case study on titanium and silicon dioxide representative nanomaterials, *J. Appl. Toxicol.*, 2020, **40**(11), 1511–1525, DOI: **10.1002/jat.4002**.
- 72 A. Lankoff, W. J. Sandberg, A. Wegierek-Ciuk, *et al.*, The effect of agglomeration state of silver and titanium dioxide nanoparticles on cellular response of HepG2, A549 and THP-1 cells, *Toxicol. Lett.*, 2012, **208**(3), 197–213, DOI: **10.1016/j.toxlet.2011.11.006**.
- 73 R. Lehner, I. Zanoni, A. Banuscher, A. L. Costa and B. Rothen-Rutishauser, Fate of engineered nanomaterials at the human epithelial lung tissue barrier in vitro after single and repeated exposures, *Frontiers in Toxicology*, 2022, 4, 1–13, DOI: 10.3389/ftox.2022.918633.
- 74 F. Cappellini, S. Di Bucchianico, V. Karri, et al., Dry Generation of CeO<sub>2</sub> Nanoparticles and Deposition onto a Co-Culture of A549 and THP-1 Cells in Air-Liquid Interface—Dosimetry Considerations and Comparison to

- Submerged Exposure, Nanomaterials, 2020, 10(4), 618, DOI: 10.3390/nano10040618.
- 75 X. Lai, Y. Wei, H. Zhao, et al., The effect of Fe<sub>2</sub>O<sub>3</sub> and ZnO nanoparticles on cytotoxicity and glucose metabolism in lung epithelial cells, J. Appl. Toxicol., 2015, 35(6), 651-664, DOI: 10.1002/jat.3128.
- 76 K. Lindner, M. Ströbele, S. Schlick, et al., Biological effects of carbon black nanoparticles are changed by surface coating with polycyclic aromatic hydrocarbons, Part. Fibre Toxicol., 2017, 14(1), 1-17, DOI: 10.1186/s12989-017-0189-1.
- 77 R. D. Bengalli, G. Zerbi, A. Lucotti, T. Catelani and P. Mantecca, Carbon nanotubes: structural defects as stressors inducing lung cell toxicity, Chem.-Biol. Interact., 2023, 382, 110613, DOI: 10.1016/j.cbi.2023.110613.
- 78 E. Di Ianni, J. S. Erdem, P. Møller, et al., In vitro-in vivo pulmonary inflammogenicity correlations of genotoxicity of MWCNT, Part. Fibre Toxicol., 2021, 18(1), 1-16, DOI: 10.1186/s12989-021-00413-2.
- 79 M. J. Burgum, V. Alcolea-Rodríguez, H. Saarelainen, et al., The dispersion method does not affect the in vitro genotoxicity of multi-walled carbon nanotubes despite inducing surface alterations, NanoImpact, 2025, 37, 100539, DOI: 10.1016/J.IMPACT.2024.100539.
- 80 V. Alcolea-Rodriguez, V. I. Dumit, R. Ledwith, R. Portela, M. A. Bañares and A. Haase, Differentially Induced Autophagy by Engineered Nanomaterial Treatment Has an Impact on Cellular Homeostasis and Cytotoxicity, Nano 11793-11799, Lett., 2024, 24, DOI: 10.1021/ acs.nanolett.4c01573