

Determining nanoform similarity *via* assessment of surface reactivity by abiotic and *in vitro* assays

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ABSTRACT

Grouping of substances is a method used to streamline hazard and risk assessment. Assessment of similarity provides the scientific evidence needed for formation of groups. This work reports on justification of grouping of nanoforms (NFs) *via* similarity of their surface reactivity. Four reactivity assays were used for concentration dependent detection of reactive oxygen species (ROS) generated by NFs: abiotic assays FRAS, EPR and DCFH2-DA, as well as the *in vitro* assay of NRF2/ARE responsive luciferase reporter activation in the HEK293 cell line. Representative materials (CuO, Mn₂O₃, BaSO₄, CeO₂ and ZnO) and three case studies of each several NFs of iron oxides, Diketopyrrolopyrroles (DPP)-based organic pigments and silicas were assessed. A novel similarity assessment algorithm was applied to quantify similarities between pairs of NFs, in a four-step workflow on concentration-response curves, individual concentration and response ranges, and finally the representative materials. We found this algorithm to be applicable to all abiotic and *in vitro* assays that were tested. Justification of grouping must include the increased potency of smaller particles *via* the scaling of effects with specific surface, and hence quantitative similarity analysis was performed on concentration-response in mass-metrics. CuO and BaSO₄ were the most and least reactive representative materials respectively, and all assays found BaSO₄/CuO *not* similar, as confirmed by their different NOAECs of *in vivo* studies. However, similarity outcomes from different reactivity assays were not always in agreement, highlighting the need to generate data by *one* assay for the representative materials *and* the candidate group of NFs. Despite low similarity scores *in vitro* some pairs of case study NFs can be accepted as sufficiently similar because the *in vivo* NOAECs are similar, highlighting the conservative assessment by the abiotic assays.

1. Introduction

Many substances are produced as nanomaterials, each of which can be generated in a variety of nanoforms (NFs) differing in size, crystallinity, shape or surface chemistry (Stone et al., 2020). The wide range of nanoforms available provides the possibility to generate a high diversity of commercial nano-enabled products. The different physicochemical characteristics of NFs may influence their toxicological profiles.

According to the REACH regulation (Helsinki, 2019), for each NF of a substance a set of minimum standard information has to be provided, and therefore the cost, duration, and effort of testing may hugely increase. However, this process can be streamlined through the use of alternative approaches such as grouping and read-across. The GRACIOUS Framework provides a logical and science evidenced approach to group similar NFs, allowing read-across of hazard information from source NFs (or non-NFs) with adequate hazard data to

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target NFs that lack such data (Stone et al., 2020). The GRACIOUS Framework supports the user to generate a grouping hypothesis that encompasses the relevant physicochemical characteristics, route of exposure and hazard endpoints. Integrated Approaches to Testing and Assessment (IATAs) are then used to gather the existing information needed to test the grouping hypothesis, and to guide the generation of new data to fill data gaps. The IATAs consist of decision trees, with each decision node posing a question that allows identification of the most relevant information needed to test the grouping hypothesis. Each decision node is supported by a tiered testing strategy (e.g. (Verdon et al. (2021))) to guide the gathering of evidence *via* the most appropriate, and if possible, standardised methods available. For human health studies, the tiered testing strategy includes simple *in vitro* models at tier 1, as well as a number of alternative more complex multi-cellular *in vitro* models at tier 2 and *in vivo* models at tier 3. The lower tier data provides the evidence to assess similarity of NF physicochemical characteristics (what they are), fate in the environment and toxicokinetics (where they go) and hazards (what they do – including surface reactivity). If sufficiently similar, the data can then be used to support read-across.

Application of grouping of NFs would therefore help to reduce the amount of experimental testing for hazard and risk assessment, thus reducing animal testing (Stone et al., 2020). As indicated above, grouping requires methods to assess the similarity of different NFs. A summary of different methods is available in the white paper of this issue (Jeliazkova et al., 2022). For the case studies investigated in this paper, various NFs were available for study, allowing an assessment of their similarity. As the white paper demonstrates, the use of scalar descriptors (a single value that represents a range of data or a concentration-response curve) is a convenient way to compare similarity. Here, we explore methods to assess similarity using the full concentration-response curve, in order to take into consideration that variations in the shape of the concentration-response curve can lead to loss of information when reduced to a scalar descriptor. To achieve this, reactivity concentration-response curves of the different NFs were evaluated *via* Bayes Factor (BF) calculations of pairwise (two NFs directly compared at a time) similarity assessments, in addition to assessing their similarity across the separate concentration (x-axis) and reactivity (y-axis) scales separately. The white paper (Jeliazkova et al., 2022) demonstrates that for scalar descriptors, the BF algorithm was consistent with simpler approaches, but statistically was more robust, and especially well-suited for the comparison of two-dimensional data such as concentration-response curves. The method of assessing similarities between concentration-response curves *via* BF calculations is originally presented in this issue by Tsiliki et al. (2021), however here we introduce a novel similarity assessment approach and integrate information from the similarity assessment between reactivity concentration-response curves and the comparisons of the varying ranges of the concentration and reactivity data available. This integrated information, which we denote by similarity score, quantifies how similar two NFs are and can then be compared to threshold values set by the representative materials to justify grouping.

In line with other scientific approaches and frameworks for grouping of NFs developed (Worth et al., 2017; Giusti et al., 2019; Arts et al., 2015), the GRACIOUS IATA for the inhalation (Braakhuis et al., 2021; Murphy et al., 2021) route of exposure identifies lung deposition, dissolution rate, *in vitro* inflammation, and surface reactivity as decisive properties to compare NFs. Surface reactivity is also a key parameter to compare the toxicity of NFs in the IATA for the oral route of exposure (Di Cristo et al., 2021). Moreover, the ECHA guidance recommends justifying a grouping of NFs *via* similarity of their surface reactivity (Hel-sinki, 2019), and the same guidance advises to justify grouping decisions “mainly using physicochemical parameters and/or *in vitro* screening methods”, consistent with tier 1 and tier 2 of the testing strategy of the “reactivity” decision node of the inhalation IATA (Braakhuis et al., 2021; Echa, 2019). Assessment of similarity is key to decide whether different NFs can be included in a group. To achieve this well-defined algorithm

to quantify the similarity between two (or more) NFs are required. Until now, there are no harmonized and standardised assays to assess either similarity or the reactivity of NFs. We considered the ROS generation as mode of action of toxicity of NFs; we chose several assays for assessing ROS generation (which could be integrated in a testing strategy) and tested the same NFs in the different assays. Then we developed a procedure to evaluate the similarity between NFs tested in the same assay.

Reactive oxygen species (ROS) cause oxidative stress and cytotoxicity, and can be generated by NFs. Understanding NF reactivity is a key stage towards understanding the toxicology, because the generation of ROS can trigger sub-lethal (e.g. inflammation) and lethal (e.g. apoptosis) effects (Zhao and Riediker, 2014). NF based ROS production occurs *via* different mechanisms such as Fenton reaction (in the presence of divalent metal ions such as Fe²⁺), redox cycling and radical generation. Fenton-like reactions are reported to be the most common mechanism for metal NFs, leading to the generation of hydroxyl radicals (Bi and Westerhoff, 2019). Different assays are available to measure free radicals and ROS, however they all differ in the mechanism of detection, sensitivity and specificity. Since the exact mechanisms of ROS mediated effects of NFs are not well understood, several *in vitro* and abiotic reactivity assays are usually used (Zhao and Riediker, 2014; Angelé-Martínez et al., 2017a; Eom and Choi, 2009). The ferric reduction ability of serum (FRAS) assay utilizes human blood serum (HBS), to quantify the total antioxidant depletion induced by NFs as a measure of their oxidative potential. Moreover, this assay has shown potential to separate active from passive NFs (Pal et al., 2014), and to be specifically useful for grouping purposes because it can distinguish between different amounts of oxidative stress (Arts et al., 2016). Electron spin resonance (ESR) spectroscopy identifies qualitatively and quantitatively free radical species in abiotic and cellular environments (He et al., 2014). The ESR spin-trapping technique uses chemical species called spin traps, which react with short-lived free radicals to form relatively stable adducts having a half-life long enough for ESR measurement (Buettnner, 1987). Another commonly used assay assesses the oxidation of the non-fluorescent molecule 2'-7'-dichlorodihydrofluorescein diacetate (DCFH₂-DA), into a fluorescent form in the presence of ROS (Pal et al., 2014). DCFH₂-DA was first devised to detect ROS in the absence of cells (Brandt and Keston, 1965) and more recently it was suggested to be used as a tool to study cellular and abiotic ROS produced in response to nanomaterials (Wilson et al., 2002) and has been set up as a standard operating procedure for use with nanomaterials (Boyles et al., 2022). There are also several cellular assays which can be used to measure the impacts of ROS on cells. For example, the nuclear factor erythroid 2-related factor 2 (Nrf2)/ antioxidant response element (ARE) pathway is an important cellular defence system that is activated by various stresses (Niture et al., 2014). NRF2/ARE Responsive Luciferase Reporter HEK293 Cell Line can be used as an *in vitro* model for monitoring the activation of antioxidant response pathways triggered by treatment with NFs. The light induction in response to ROS and Nrf2 interaction with the ARE makes this an attractive model to study.

Here, reactivity of representative materials (CuO, Mn₂O₃, BaSO₄, CeO₂ and ZnO) plus three case studies (iron oxides, Diketopyrrolopyrroles (DPP)-based organic pigments and silica NFs) were tested *via* the abiotic assays FRAS, EPR and DCFH₂-DA, as well as the cellular *in vitro* assay, activation of NRF2/ARE Responsive Luciferase Reporter HEK293 Cell Line. In this work we address both the experimental data acquisition and the similarity evaluation. The similarity level of NFs in each reactivity assay and the consistency of similarity when compared with higher-tier (*in vivo*) results were evaluated.

2. Material and methods

2.1. Materials

ZnO NM110 and CeO₂ NM212 were kindly provided by the NM repository at the Joint Research Center (JRC) repository (JRC

Nanomaterials Repository, n.d.). BaSO₄ NM220 was provided by the NM repository at the Fraunhofer Institute (IME). Mn₂O₃ was purchased from Skyspring Nanomaterials. CuO was purchased from Plasma Chem. Fe₂O₃ nanoA (small rods, about 15 nm), Fe₂O₃ nanoB (rounded particles, about 30 nm), Fe₂O₃ larger (irregular particles, above 100 nm), FeOOH (small rods, about 15 nm) and organic pigments (DPP nano (43 nm), DPP nonano (233 nm), DPP premixed (230 nm)) were supplied by BASF Colors and Effects BASF Schweiz AG. Silica std. (standard silica, 30% in water), Silica Al (Al substituted into the silica surface, 25% in water), Silica silane (silane modified, 28% in water), Silica anis Al (Al substituted into the surface, aggregated nanoparticles, 7% in water) and Silica-anis-std (aggregated silica nanoparticles, 12% in water) were supplied by NOURYON.

The physicochemical properties and TEM images of materials are reported separately (Jeliazkova et al. 2021 data base publication) and are reproduced here in the supplementary information (Table S1 and Fig. S1). The reagents employed during each reactivity assay are reported in the supplementary information.

All NFs were tested in all reactivity assays.

2.2. Ferric reduction ability of serum (FRAS)

The SOP, which described a multi-dose protocol of the FRAS assay and was published in 2017 by Gandon et al. (Gandon et al., 2017), was used for reactivity testing of samples.

Briefly, samples were incubated with human blood serum for 3 h at 37 °C. Before incubation, bath sonication for 1 min was applied to prevent the formation of large agglomerates and allow the reagents to access the whole surface area. NF were separated from HBS via ultracentrifugation (AUC-Beckman XL centrifuge (Brea, CA, USA) at 14,000 G for 150 min). Subsequently, a 100 µL of NF-free HBS supernatant was incubated in the FRAS reagent that contains the Fe³⁺ complex. The total antioxidant depletion, as a measure of the oxidative potential of NF, was determined by assessing the UV-vis spectrum of the iron complex solution. Trolox, a water-soluble analog of vitamin E, was used as an antioxidant to calibrate the FRAS results. Different Trolox concentrations (from 0.001 to 0.1 mg/mL) were tested by the FRAS assay to obtain FRAS absorption signals that were linearly fitted. Finally, the oxidative damage induced by NF was calculated in Trolox equivalent units (TEUs).

Background FRAS signal level is up to 5000 nmol TEU/L and saturation of FRAS signals occurred at the level of about 250,000 nmol TEU/L, indicating that all antioxidants contained in the human serum are consumed during the incubation. 0.02 to 40 mg/mL dose range was applied to the representative test materials and case study materials. This range is a two-sided extension of the range of 0.15 to 10 mg/mL that was used in the extensive screening of 138 nanomaterials, each at an “adjusted” single concentration (Hsieh et al., 2013). We find that CuO and Mn₂O₃ reach saturation of the assay for all concentrations above 0.2 mg/mL.

2.3. Electron paramagnetic resonance spectroscopy (EPR)

EPR can be used to identify and quantify unpaired electron spins, e.g. to characterize the active sites of solid-state catalysts. Two methods have been established to assess the surface-induced reactivity of nanomaterials: Method 1 utilizes the nitron spin trap 5,5-Dimethyl-1-pyrroline-N-oxide (DMPO), one of the most established spin traps for nanosafety purposes (Hellack et al., 2017a). This method involves trapping reactive short-lived free radical intermediates (e.g. hydroxyl radical) via the creation of the spin adduct DMPO-OH with a characteristic 1:2:2:1 peak pattern and g-value. Method 2 employs the cyclic hydroxylamine spin probe 1-hydroxy-3-carboxy-pyrrolidine (CPH). CPH directly probes/interacts with short-lived reactive oxygen species (e.g. superoxide radical) on the material surface, forming the spin adduct CP• with characteristic 1:1:1 peak pattern and g-value. Both methods are standardised by ISO TS 18827 (ISO 2017) and iuta SOP: EPR

spectroscopy analysis using the spin probe CPH (by B. Hellack), respectively.

For EPR detection of superoxide anions, the spin trap 1-hydroxyl-2,2,6,6-tetramethyl-4-oxo-piperidine (Tempone-H) was prepared at 100 nM in 0.01 M EDTA, to be used at a final concentration of 1 mM. Test materials were prepared in a phosphate buffer at a starting concentration of 4 mg/mL, with a concentration-response measured between 0.002 and 4 mg/mL, dependent on material. Pyrogallol, at 32 mM, was used as a positive control. Measurements were taken 60 min after addition of Tempone-H, with samples maintained at 37 °C during this time. Using a Miniscope MS 200 (Magnetech, Berlin, Germany), the EPR spectrum was obtained with the following parameters: microwave frequency, 9.3–9.55 Hz; microwave power, 20 mW; modulation frequency, 100 kHz; modulation amplitude, 1500 mG; center field, 3350 G; sweep width, 55 G; sweep time, 30 s; number of passes, 1.

2.4. Dichlorodihydrofluorescein diacetate (DCFH2-DA)

Detection of ROS produced using the DCFH2-DA probe was conducted as follows. DCFH2-DA was chemically hydrolysed by incubation with 0.01 M NaOH, neutralized and diluted to form 10 µM DCFH2 in phosphate-buffered saline (PBS). During this reaction, test particles were prepared by suspension in phenol red-free minimum essential medium (MEM) with 2% FCS at a concentration of 40 mg/mL, followed by ultra-sonication in a water bath and serial dilutions to obtain a range of 5, 10, 20 and 40 mg/mL. Each treatment was then added, in triplicate to a 96-well plate at a volume of 25 µL, followed by addition of 225 µL 10 µM DCFH2 to each well. Final concentrations of 0.5, 1, 2 and 4 mg/mL were obtained, which were incubated at 37 °C for 90 min. After this time, samples were centrifuged at 3000 xg for 15 min, and 100 µL of each well was moved to a black 96-well plate to read fluorescence at ex/em wavelengths of 485/530 nm. To address the potential for interference of particles with the light detection, the same process as above was replicated using particles suspended in solutions of PBS alone (no DCFH2), or with 0.1 µM fluorescein diacetate (FDA). To account for background interference, signals generated with incubation in solutions of PBS alone were removed from signals generated in solutions of DCFH2.

2.5. In vitro assay: Nrf2-activation

The detailed SOP for the assessment of Nrf2-activation is published in Guisti et al. (2021) The main points are briefly reported hereafter for more clarity.

2.5.1. Preparation of NF dispersion

NF stock suspensions were prepared according to NanoToxClass dispersion protocol, which consists in cup horn sonication of NF in serum free cell medium at a concentration of 0.5 mg/mL (Bandelin Cuphorn UW 2200 for 23 min at 100% power). These suspensions were then diluted to the desired concentrations. For Nrf2 activation the dilution media is the complete cell medium but containing only 1% FBS; the final NF concentrations are to 0, 1.2, 3.6, 10.7, 32.1 µg/mL.

2.5.2. Measure of Nrf2-activation

A stably transfected cell line encoding a firefly luciferase reporter gene under the control of ARE element (NRF2/ARE Luciferase Reporter HEK293 Stable Cell Line) from Signosis was used. Cells were grown in DMEM cell culture medium (w/o phenol red and L-glutamine, high glucose, PAN Biotech GmbH) supplemented with 584 mg/L L-Glutamin, 0.1 mg/mL Penicillin/Streptomycin, 110 mg/L Sodium Pyruvate, 80 µg/mL Hygromycin B gold and 10% Fetal Bovine Serum (non-heat inactivated FBS Good from PAN Biotech). The cells were grown in T75 cell culture flasks in an incubator (37 °C, 5% CO₂ and 90% humidity) and sub-cultured regularly two times a week at ca. 70% confluence. Cells were seeded in 96-well white plates Greiner Bio-One P/N 655098 24 h before the treatment (10,000 cells in 0.1 mL pro well). After 24 h

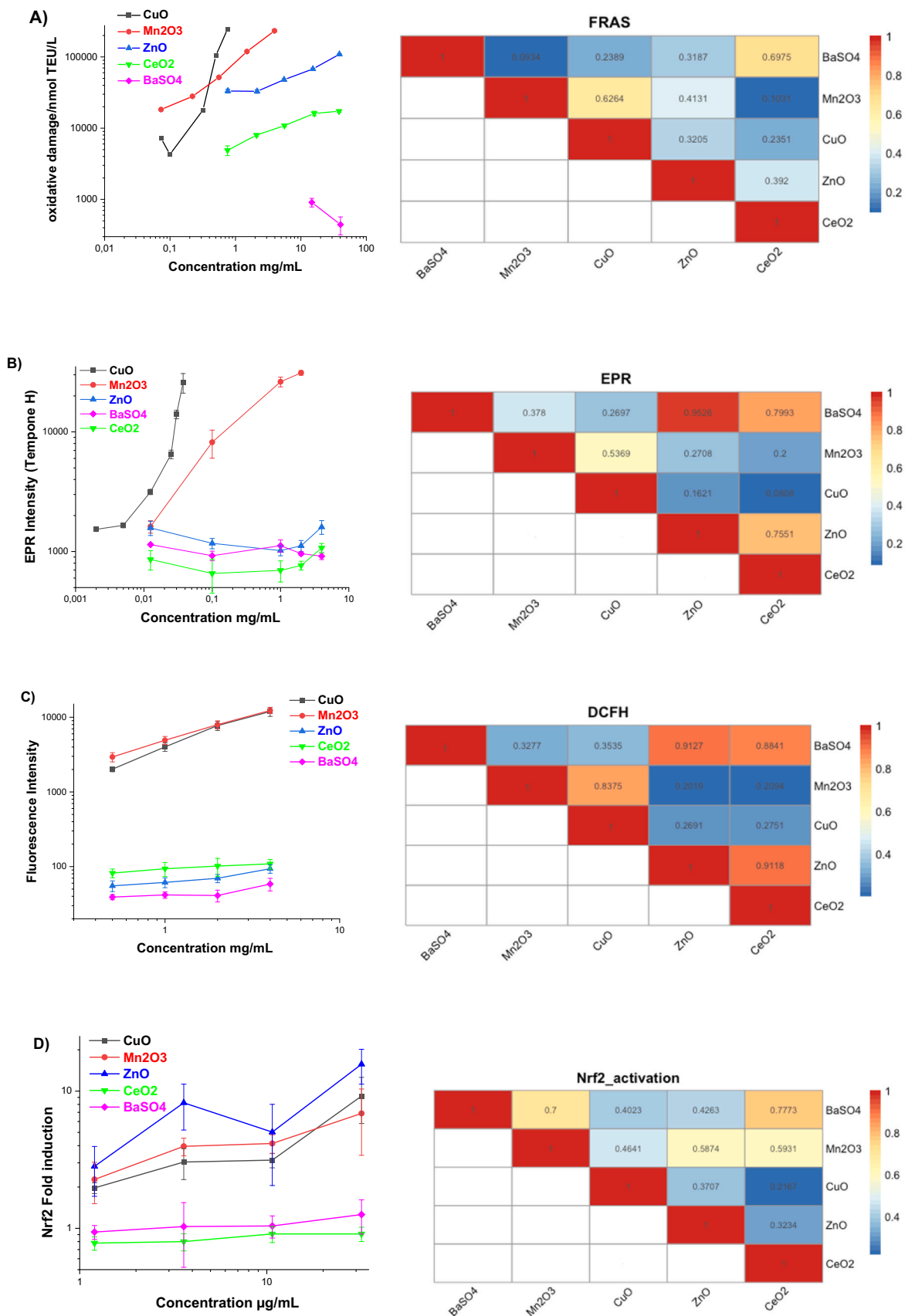


Fig. 1. Left side concentration-response curves of FRAS (A), EPR (Tempone H) (B), DCFH2-DA (C), Nrf2 activation (D) assays for representative test materials and on the right side their corresponding similarity plots for all possibly pairwise comparisons. Ideal similarity has a score of 1 (red), and the lowest similarity has a score of 0 (blue). (For interpretation of the references to colour in this figure legend, the reader is referred to the web version of this article.)

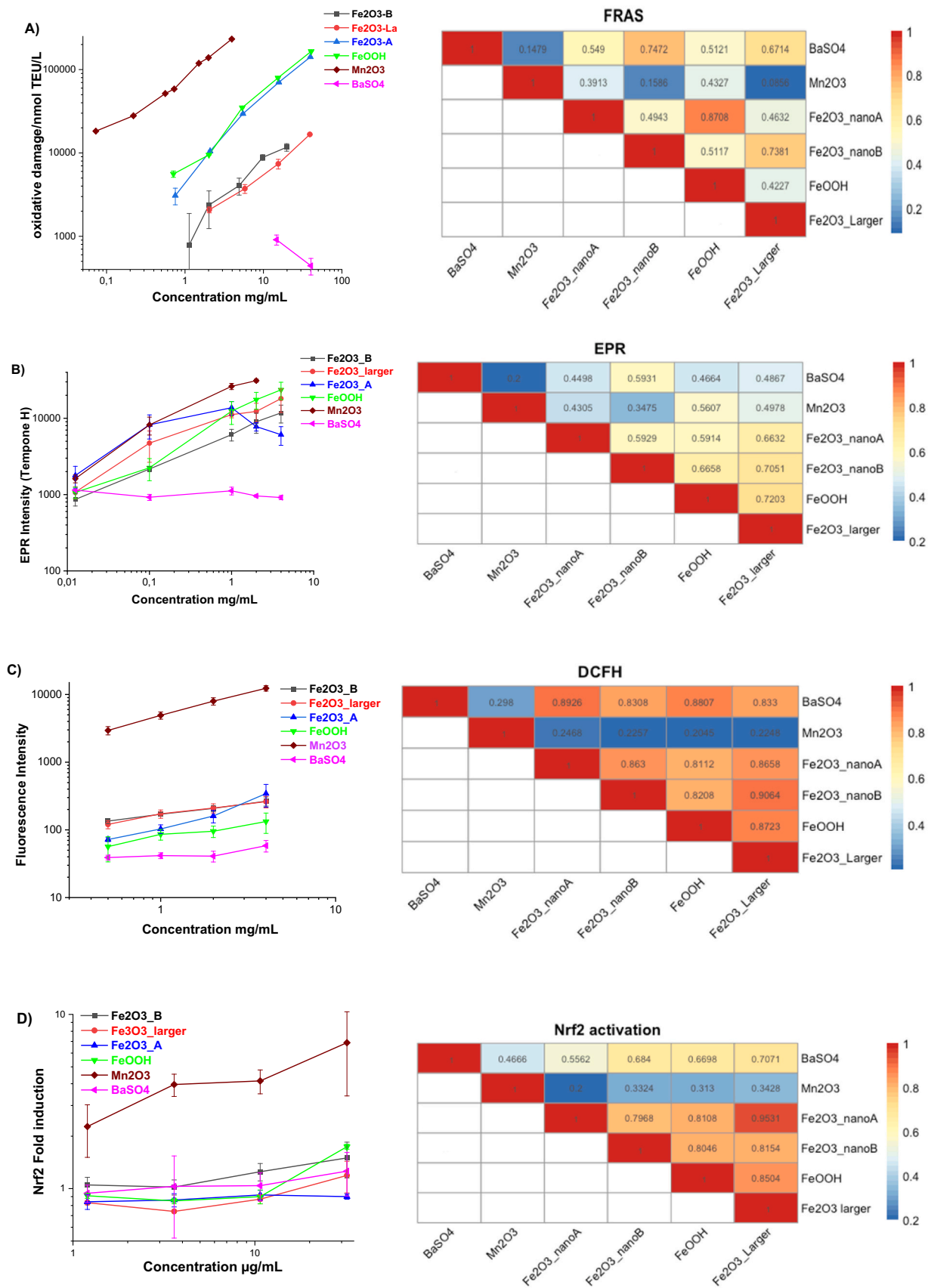


Fig. 2. Left side concentration-response curves of FRAS (A), EPR (Tempone H) (B), DCFH2-DA (C), Nrf2 activation (D) assays for Fe-based materials and additional Mn₂O₃ and BaSO₄; on the right side their corresponding similarity plots for all possibly pairwise comparisons. Ideal similarity has a score of 1 (red), and lowest similarity has a score of 0 (blue). (For interpretation of the references to colour in this figure legend, the reader is referred to the web version of this article.)

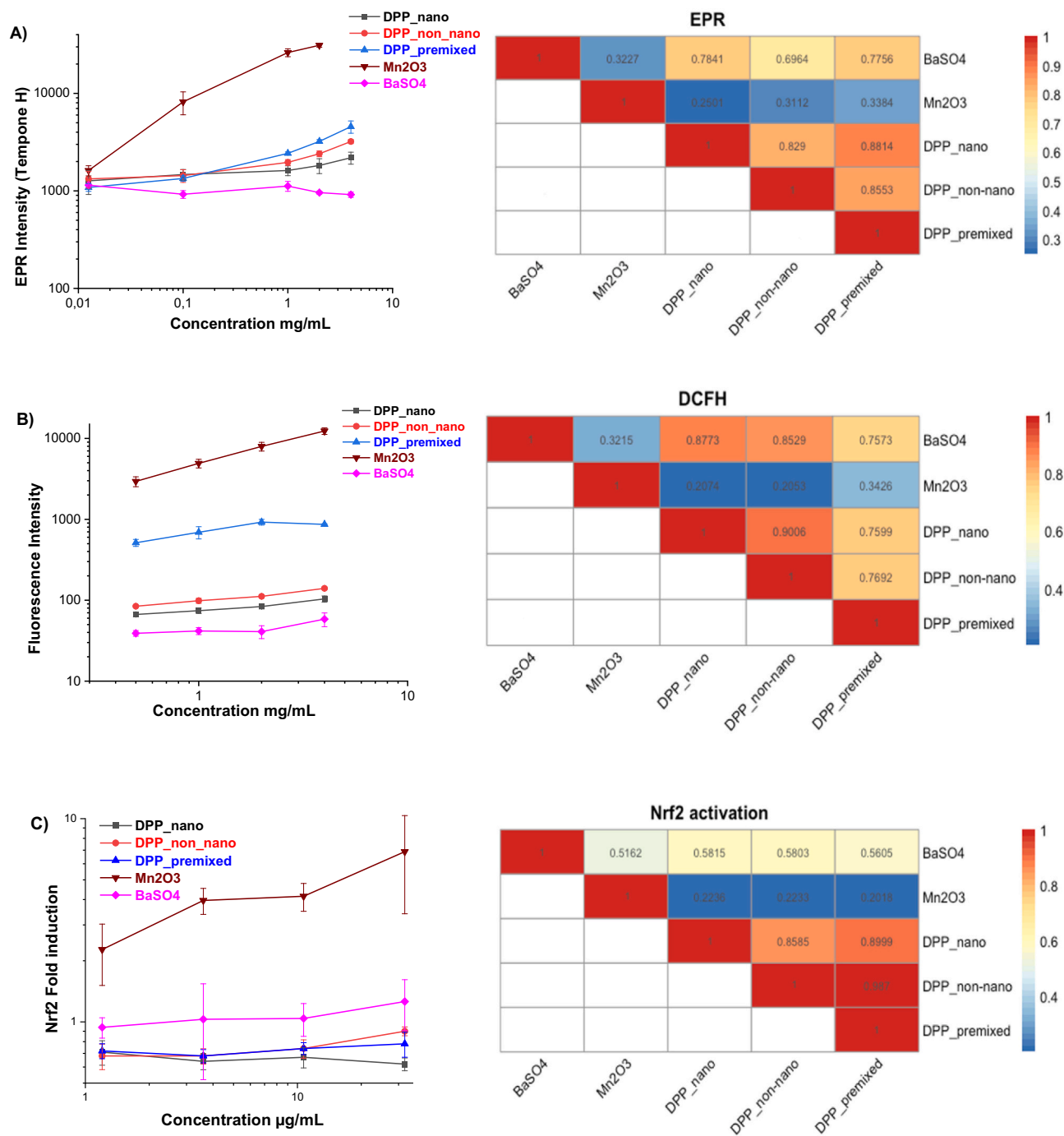


Fig. 3. Left side concentration-response curves of EPR (Tempone H) (A), DCFH2-DA (B), Nrf2 activation (C) assays for DPP pigments and additional Mn₂O₃ and BaSO₄; on the right side their corresponding similarity plots for all possibly pairwise comparisons. Ideal similarity has a score of 1 (red), and lowest similarity has a score of 0 (blue). (For interpretation of the references to colour in this figure legend, the reader is referred to the web version of this article.)

incubation at 37 °C, 5% CO₂, the cell culture medium is carefully removed and replaced with 0.2 mL treatment cell medium containing the desired final NF concentration (i.e. 0, 0.7, 2.1, 6.3, 18.9 µg/cm² corresponding to 0, 1.2, 3.6, 10.7, 32.1 µg/mL). Cells were treated for 48 h.

2.6. Similarity analysis to compare concentration-response curves

Pairwise similarity analysis was performed in a 3-step manner employing three different criteria, namely assessing the similarities between shapes of reactivity concentration-response curves, similarities

between the concentration factor ranges, and similarities between the reactivity factor ranges. The three criteria were quantified with a scalar metric each and aggregated to a unique value, denoted by similarity score, which takes values in the range between 0 and 1, with values close to 1 denoting high similarity. Lastly, representative test materials were used to set the biological relevant range of the assay. Similarities between concentration-response curves for each pair of NF were assessed using BF calculations which can be interpreted as indexes of preference for one model over another, suggesting by how much a data sample should update our belief in one model over a competing one (Faulkenberry, 2018). Given a pair of concentration-response curves, two models

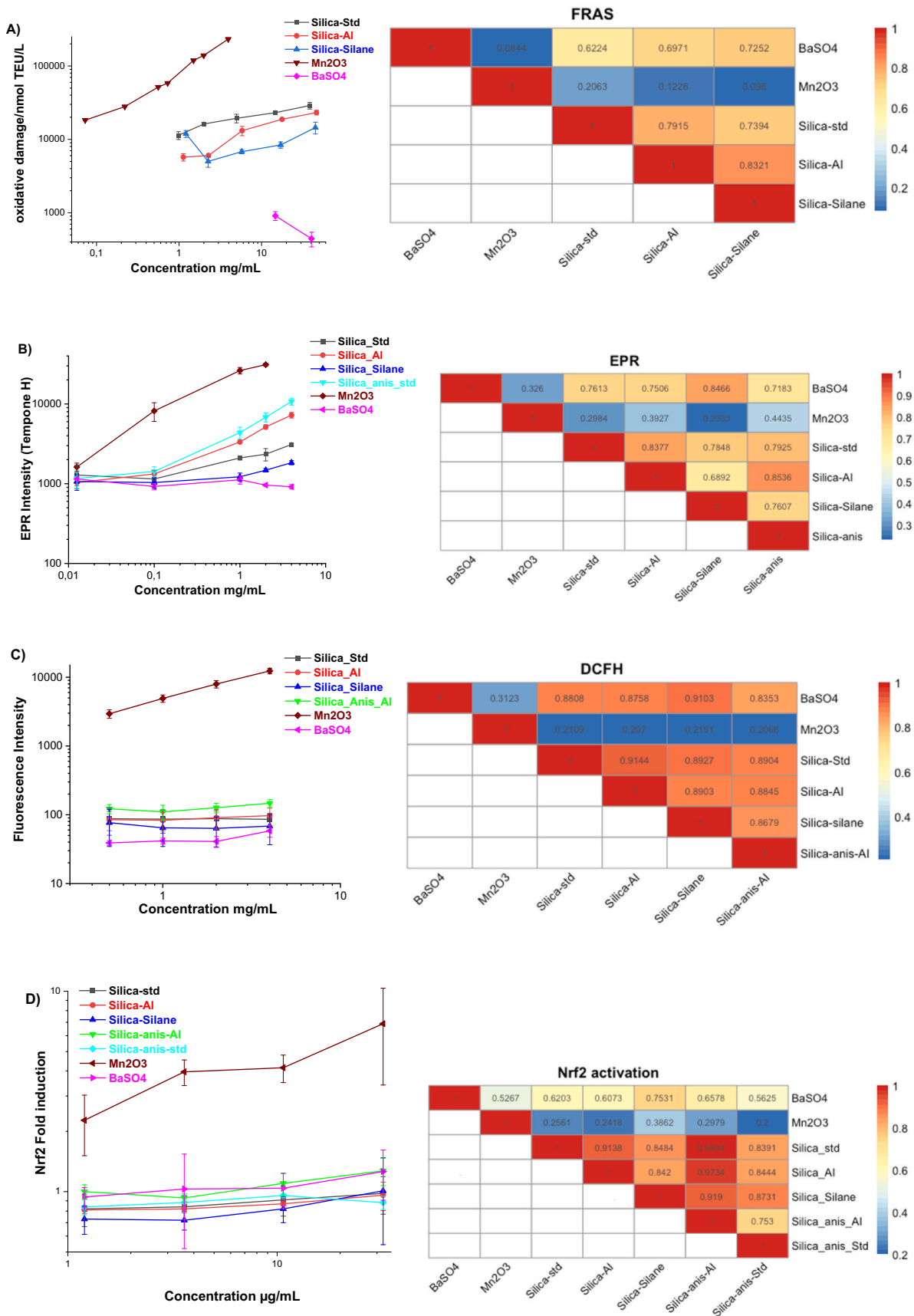


Fig. 4. Left side concentration-response curves of FRAS (A), EPR (Tempone H) (B), DCFH2-DA (C), Nrf2 activation (D) assays for silica-based materials and additional Mn₂O₃ and BaSO₄; on the right side their corresponding similarity plots for all possibly pairwise comparisons. Ideal similarity has a score of 1 (red), and lowest similarity has a score of 0 (blue). (For interpretation of the references to colour in this figure legend, the reader is referred to the web version of this article.)

Table 1

Pairwise similarity scores for representative test materials in FRAS, EPR (Tempone H), DCFH and Nrf2 activation assays. Ideal similarity has a score of 1, and lowest similarity has a score of 0. Keeping in mind that reactivity is only one of several relevant properties, the results can be compared to the available *in vivo* inhalation NOAEC values for each pair of NFs (Wohlleben et al., 2019).

Pair of materials	Similarity scores by reactivity assays				NOAEC (mg m ⁻³)
	FRAS	EPR	DCFH	Nrf2	
BaSO ₄ / CuO	0.2389	0.2697	0.3535	0.4023	50 / 0.6
BaSO ₄ / ZnO	0.3187	0.9526	0.9127	0.4263	50 / 0.5
BaSO ₄ / CeO ₂	0.6975	0.7993	0.8841	0.7773	50 / <0.5
CuO / ZnO	0.3205	0.1621	0.2691	0.3707	0.6 / 0.5
CuO / CeO ₂	0.2351	0.0808	0.2751	0.2167	0.6 / <0.5
ZnO / CeO ₂	0.3920	0.7551	0.9118	0.3234	0.5 / <0.5

are compared, specifically the first model (M₁) assumes the concentration-response curves of the two NFs are identical as opposed to the second model (M₂) which assumes that curves are coming from different distributions (Tsiliki et al., 2021). Positive B₁₂ values suggest that M₁ is preferable compared to M₂, and the two NFs can be assumed to be similar given the data.

Similarities between ranges of the concentration-response factors, in this case concentration-reactivity data, were also quantified per factor using the Manhattan distance metric in both cases. This was found to be an important adjustment to the BF calculations in order to cope with large differences in the concentration ranges measured.

The final similarity score reported is a weighted average distance metric, which for each pair of NFs, combines the BF value with quantification of the distance between the ranges of the response reactivity values d_R and the distance between the ranges of their concentration d_D . The specific weights shown below were selected after a recurring adaptation procedure to adequately distinguish the active and passive NFs from the representative test materials. Other factors may be possible, but their validity need to be demonstrated on suitable representative test materials.

$$\text{similarity score} = (0.3 \cdot BF) + (0.5 \cdot d_R) + (0.2 \cdot d_D)$$

3. Results and discussion

3.1. Results description

Concentration dependent abiotic and cell-based *in vitro* reactivity assays were applied on representative test materials and case study materials. The following sections first present the concentration-dependent reactivity in mass dose metrics (as required by the IATA to compare potency) (Braakhuis et al., 2021), and then in surface dose metrics (to check for *qualitative* similarity of the specific reactivity).

3.1.1. Results in mass dose metrics

CuO and Mn₂O₃ induced a concentration dependent reactivity in all reactivity assays (Fig. 1A-D). The FRAS, EPR (DMPO) and Nrf2 activation assays showed concentration dependent reactivity of ZnO NF (Fig. 1A, D and S2 A). The FRAS assay distinguished very reactive materials (CuO, Mn₂O₃), from the intermediate reactive material ZnO, and from rather passive materials (Fig. 1A). ZnO presented the highest reactivity in the Nrf2 activation assay (Fig. 1 D). The DCFH2-DA assay and measurement of two spin traps (CPH and Tempone H) by EPR found only low reactivity of ZnO (Fig. 1 B, C and S2A); this is not in keeping with the known *in vivo* toxicity of ZnO which induces inflammation (in rats) and ecotoxicity at low concentrations (Di Cristo et al., 2021; Boyles et al., 2022). However, the exact mechanism of these different cases is not discussed here. Only FRAS assay presented concentration dependent responses by CeO₂ (Fig. 1A). BaSO₄ did not produce any concentration dependent responses in any assay (Fig. 1 A-D).

All Fe-based materials induced a concentration dependent increase

in reactivity according to FRAS assay and EPR (Tempone H) (Fig. 2 A, B). However, Fe₂O₃-nanoA showed an unexplained decrease in reactivity at high concentrations (from 1.0 mg/mL) in EPR assay by using Tempone H spin trap. Except Fe₂O₃-Larger, other Fe-based NFs had concentration dependent reactivity in EPR (DMPO) assay (Fig. S3 A). DCFH2-DA and Nrf2 activation assays did not generate any concentration dependent reactivity for Fe-based materials (Fig. 2 C, D).

On pigments, the FRAS assay may have suffered from optical interference and flocculation of pigment samples in serum. Colorimetric assays, including the FRAS assay, are not recommended for NM with high absorption coefficient (i.e. where traces of NM generate a large interference with colorimetric readout). All DPP based pigments demonstrated slightly increased reactivity in a concentration dependent manner as assessed by the EPR (Tempone H) assay (Fig. 3A), however no concentration response was observed by DCFH2-DA, Nrf2 activation and EPR (DMPO) assays (Fig. 3 B, C and S4 A).

Due to low particle concentrations of the pristine products Silica-anis-Al (7%) and Silica-anis-std (12%), the FRAS Assay could not be applied to these NFs. Silica-std, Silica-Al and Silica-silane showed concentration dependent responses in the FRAS assay (Fig. 4A). The EPR (Tempone H) assay also generated concentration dependent reactivity for Silica-std, Silica-Al, Silica-silane and Silica-anis-std (Fig. 4B). DCFH2-DA, Nrf2 activation and EPR (DMPO) assays did not demonstrate any concentration dependent responses for all silica-based NFs (Fig. 4 C, D and S5 A).

3.1.2. Results in surface dose metrics

The reactivity of particles is in fact often referred to as “surface reactivity”, because the reaction is thought to occur at the particle surface (Hellack et al., 2017b). If two NFs induce different reactivity in mass-dose metrics, they may still induce similar reactivity after rescaling to surface-dose metrics (Bahl et al., 2020). This would then indicate a *qualitative* similarity of the reactivity. Examples of qualitative similarity were observed on the Fe-based materials, where all Fe₂O₃ NFs collapse onto one concentration-response curve in the surface-dose representation of FRAS reactivity (Fig. S8A), whereas the chemically different FeOOH stands out (Fig. S8A). Another example is given by the NF and the non-nano-form of DPP pigment, which collapse onto one concentration-response curve in the surface-dose representation of both EPR and DCFH reactivity (Fig. S7A, S7B). In contrast, the different representative test materials maintain a very different concentration response relationship also in surface-dose representation (Fig. S6). It has been argued that surface dose allows the best understanding of inhalation toxicity (Schmid and Stoeger, 2016; Oberdörster and Kuhlbusch, 2018). Notwithstanding that systematic understanding -or even based on it- the scaling of effects with specific surface area is one of the reasons for regulators to demand a separate assessment of NFs of a substance. A justification of grouping several NFs with respect to their inhalation hazard must consider that even at the same surface-specific reactivity, particles with a higher specific surface area induce more oxidative damage. In line with the IATA (Braakhuis et al., 2021), the following section is devoted to quantitative similarity of mass-dose potency.

3.2. Results of quantitative similarity assessments

Previously, single reactivity descriptors were used for grouping purposes in the nanoGRAVUR framework (Wohlleben et al., 2019). For two reasons we are unable to do this here for each assay using concentration response curves: the range in effect was so vastly different between many of the particles, which makes the EC50 comparison unrealistic, as the method provides a value based on the range restrictions of each individual material. Secondly, some of the treatments have such a low effect (negative for the BaSO₄) that a reliable EC50 is unattainable. Therefore, we use a similarity algorithm (BF calculations) for further discussion that facilitates the discussion based on the complete concentration dependency. This enables a direct comparison of the

Table 2

Pairwise similarity scores for Fe₂O₃_nanoA and Fe₂O₃_Larger in FRAS, EPR (Tempone H), DCFH and Nrf2 activation assays, and available NOAEC values for each pair of NFs (Niture et al., 2014).

Samples	Similarity scores				NOAEC (mg m ⁻³)
	FRAS	EPR	DCFH	Nrf2	
Fe ₂ O ₃ _nanoA / Fe ₂ O ₃ _Larger	0.4632	0.6632	0.8658	0.9531	30 / 30

similarity assessment based on different assays.

3.2.1. Representative test materials

Pairwise similarity was analyzed by Bayes factor algorithm using reactivity data from four different assays. The pairwise comparisons result in a triangular similarity matrix (Fig. 1) that allows pairwise comparison of NFs by reading along rows and down columns. Colors and similarity score numbers are used in the triangular similarity matrices to indicate the degree of similarity between two NF, with warm colors (yellow/red colors and numbers close to 1.0) indicating a high degree of similarity, and cool colors (blue colors and numbers close to 0.0) at the opposite end of the spectrum representing the NFs that are not similar.

Pairwise similarity scores (numbers) of representative test materials in FRAS, EPR (TemponeH), DCFH2-DA and Nrf2 activation assays were summarized in Table 1 and Table S2. CuO and Mn₂O₃ demonstrated high reactivity in all abiotic reactivity assays, which resulted in similar/very similar similarity as indicated with warm colors (Fig. 1). On the other hand, BaSO₄ and CeO₂ were non-reactive in all assays and presented very similar similarity with orange colour (Fig. 1 A–D). However, BaSO₄ and CeO₂ are only similar in reactivity while the dissolution data and *in vivo* NOAECs (Table 1) of BaSO₄ and CeO₂ are different (Keller et al., 2021a) (Wohlleben et al., 2019). EPR assays using different spin traps (CPH, DMPO and Tempone H) resulted in different similarity scores compared to the data with TemponeH (as well as DCFH2-DA and Nrf2) (Fig. 1B and S2 B, C). All abiotic assays detected CuO and BaSO₄ as the most and least reactive materials respectively and clearly BaSO₄/CuO were not found to be similar for all reactivity assays. Since CuO with a NOAEC of 0.6 mg/m³ also differed substantially in its *in vivo* response from BaSO₄ with a NOAEC of 50 mg/m³ (Table 1), their choice as representative materials was confirmed.

It should be noted here that BF calculations assume log-normally distributed data and sampling from the log-normal distributions of concentration-reactivity data for each of the groups of the materials studied. Parameters of the distributions are estimated from the data. *i.e.* the group of NFs considered each time, and for that reason BF values vary even though referring to the same pair of NFs.

3.2.2. Case studies

Concentration dependent reactivity curves of case study materials were compared with very reactive Mn₂O₃ (Arts et al., 2015; Arts et al., 2016) very reactive CuO (Gosens et al., 2016), and non-reactive BaSO₄ (Buesen et al., 2014; Landsiedel et al., 2014). These three materials served as representative test materials, and their dose response was included a) in the graphical presentation for each NM class and each assay (Figs. 2 to 4), and b) in the quantitative similarity analysis (Figs. 2 to 4), where they represent the biologically relevant range, as recommended by the white paper (Jeliazkova et al., 2022). Similarity assessment is relevant for NFs with reactivity values in the biologically relevant range. Differences between case study NFs that are very small compared to this range would still allow grouping. For this reason, quantitative similarity assessment of same-substance NFs must always include at least two other substances that represent the biologically relevant range (Jeliazkova et al., 2022).

3.2.2.1. Fe-based materials. Pairwise similarity scores of Fe-based materials, Mn₂O₃ and BaSO₄ in FRAS, EPR (Tempone H), DCFH2-DA and

Nrf2 activation assays were compared in Table S3. All Fe-based materials were similar to each other and demonstrated low reactivity, as did BaSO₄ in DCFH2-DA and Nrf2 activation assays (Fig. 2 C and D). Depending on spin traps, different pairwise similarity scores were presented by EPR assays (Fig. 2B and S3). It could not be confirmed that EPR, in general, is a suitable tool for analysing Fe-based materials; different similarity outcomes were found for different spin traps, indicating the importance of understanding the specificity of spin-traps, and possible, but undetermined, sample interferences were observed in EPR assays, including the interrupted concentration-response curve observed after 1 mg/mL measurements of Fe₂O₃_nanoA.

The iron oxides were intermediate in reactivity, and different compared to both negative and positive controls in FRAS assay. Only FRAS assay distinguished two groups of very similar (nano)forms. In the first group, Fe₂O₃_nanoA and FeOOH and in the second one Fe₂O₃_nanoB and Fe₂O₃_larger showed very similar mass-dose reactivity with a score of 0.87 and 0.74 respectively (Fig. 2A). The representation of FRAS reactivity in surface-dose metrics (Fig. S8A), the response of FeOOH equals the positive control in the FRAS assay (Fig. S8A) and is an order of magnitude different from the Fe₂O₃ NFs.

The present results allow us to adjust the decision criteria of the tiered testing strategy such that the grouping decisions made with tier 1 abiotic method are not in conflict with tier 2 *in vitro* tests and tier 3 *in vivo* testing, considered as gold standard (Verdon et al., 2021; Braakhuys et al., 2021). We maintain that reactivity alone is not predictive of inhalation effects but also the reactivity assessment must not create conflicts between tiers. Similarity scores listed in Table 2 have to be accepted as sufficiently similar as Fe₂O₃_nanoA and Fe₂O₃_larger have a similar *in vivo* NOAEC (tested by short-term inhalation screening on rats, Table 1) (Wohlleben et al., 2019). In another perspective, the difference in reactivity is a false positive result of the integrated approach to testing and assessment (IATA) that should not prevent grouping, if also the other decision nodes of the IATA indicate sufficient similarity (Murphy et al., 2021). On the other hand, FeOOH is well known catalyst for Fenton like reactions (Li et al., 2015), which is a relevant ROS production mechanism in biological media. Accordingly, detection of higher reactivity in FeOOH was expected and confirmed by the FRAS assay (Fig. 2A). Especially in surface dose metrics, the response of FeOOH equals the positive control in the FRAS assay (Fig. S8A) and is an order of magnitude different from the Fe₂O₃ NFs.

3.2.2.2. DPP-based organic pigments. Pairwise similarity scores of DPP-based materials, and Mn₂O₃ and BaSO₄ in EPR (Tempone H), DCFH2-DA and Nrf2 activation assays were compared in Table S4. Except EPR (DMPO), all reactivity assays scored very high pairwise similarity (orange colour) of three pigments (Fig. 3 and S4). Moreover, they had low reactivity similar to BaSO₄, where the *in vivo* data is available (DPP nano and DPP non-nano), the similarity of low reactivity matches the similarity of low *in vivo* toxicity with inhalation NOAEC at >30 mg/m³ (Table 3) (Wohlleben et al., 2019; Hofmann et al., 2016). One notes that the similarity of the NF and the non-NF of DPP pigment is even higher in surface-dose metrics, where their curves collapse onto one concentration-response curve (Fig. S7A, S7B). The comparison to the NOAEC, however, which is a mass-based value, must equally adhere to mass-based similarity assessment.

3.2.2.3. Silica-based materials. Comparing between silica NFs, all abiotic and *in vitro* reactivity assays indicated a high similarity within all samples (orange and red colour) (Fig. 4). Pairwise similarity scores of silica materials, and Mn₂O₃ and BaSO₄ in FRAS, EPR (Tempone H), DCFH2-DA and Nrf2 activation assays were compared in Table S5. All assays demonstrated low reactivity of all silica samples which is very similar to BaSO₄.

Table 3

Pairwise similarity scores for DPP_nano and DPP_non-nano in EPR (Tempone H), DCFH and Nrf2 activation assays, and available NOAEC values for each pair of NFs (Niture et al., 2014).

Samples	Similarity scores			NOAEC (mg m ⁻³)
	EPR	DCFH	Nrf2	
DPP_nano / DPP_non-nano	0.6632	0.8658	0.9531	>30 / >30

3.3. Sensitivity of assays for the reactivity induced by specific substances

By means of several commonly used assays for the assessment of NM reactivity, we have identified a number of important factors to consider. One consideration is that specific assays can be observed as being sensitive (or insensitive) to specific material classes. This was observed here on numerous occasions, and was evident for representative test materials and case study substances. For example, CuO and Mn₂O₃ consistently induced dose dependent reactivity across all reactivity assays, while the reactivity of CeO₂ and ZnO was particularly confounded: Only FRAS assay demonstrated a dose-dependent response to CeO₂, and when ZnO was assessed, FRAS, EPR (using DMPO) and Nrf2 activation assays demonstrated clear dose-dependent reactivity, while by DCFH₂-DA and EPR using either CPH or Tempone H, no such concentration-response was observed; these general responses to all substances tested can be seen in Table 4 as a simple portrayal of whether a dose response was observed or not. These inconsistencies raise an issue of how much understanding there needs to be in specific assay parameters and the interpretation of simple reactivity endpoints. Should we consider the low reactivity of ZnO in certain assays as a false-negative affect as ZnO is known to be hazardous *in vivo*? Probably not, it just means that the mode-of-action of ZnO is better represented by certain assays than others. These findings were reflected with the use of statistical analysis and quantification of similarity. The robust three-parameter assessment model used for the statistical analysis confirmed the similarity and high reactivity of substances, such as CuO and Mn₂O₃ in all assays, and furthermore identified BaSO₄ and CeO₂ as being non-reactive in all assays. The specificity of abiotic assays to certain modes-of-action is especially versatile (Lakshmi Prasanna and Vijayaraghavan, 2015; Angelé-Martínez et al., 2017b), e.g. via different EPR probe molecules, of which CPH and DMPO spin probes (respectively, spin traps) are included in the ISO standard (ISO, 2017), and have been used to group nanomaterials by surface-induced oxidative damage (Bahl et al., 2020; Wohlleben et al., 2019). Also the simplified assays such as the FRAN (Bi and Westerhoff, 2019) or FRAP (Thaipong et al., 2006; Benzie and Strain, 1996) versions of the FRAS assay, using individual probes instead of entire human serum, are less sensitive but more specific. The lack of “realism” may thus be seen as an advantage for targeted investigations (Hellack et al., 2017b), but was less of a focus here.

4. Conclusions

Assessment of reactivity is only one of the decision nodes on a complete IATA to justify grouping of NFs, while proving similarity in each of these decision nodes is required to justify a grouping decision; conversely other considerations, such as ranking of effects will help to inform source and target for read-across decisions. Although to fully define what contributions similarity in NF reactivity can play in grouping decisions are beyond scope of the current study, this work provides useful insights in how similarity in reactivity assessment can be assessed, using representative test materials and numerous NF case studies.

This study demonstrates that a similarity assessment of NFs can be compiled via use of well-defined reactivity assays, as long as the limitations of such an assessment are understood. Here, the BF calculations were applied to compare dose-dependent reactivity over different dose

ranges from four different reactivity assays. The strength in this analysis partially comes from the robust nature in which multiple concentration-response parameters are considered within one model, including the assessment of three distinct opportunities to address similarity: the shape of the reactivity concentration-response curve, the concentration factor ranges, and the reactivity factor ranges. We found the algorithm used to be applicable to all abiotic and cell-based *in vitro* assays that were tested. This similarity assessment can serve as decision criterion in an IATA, where reactivity is one of several criteria on a data matrix of NFs and control materials. However, in this comparison, the same analytical method should be used for all NFs and control materials.

We observed several examples of *qualitative* similarity, where materials of different shape and size (but same composition) collapse onto one concentration-response curve in the surface-dose representation reactivity. However, the scaling of effects with specific surface area is one of the reasons for regulators to demand a separate assessment of NFs of a substance, and the justification of grouping must respect that even at same surface-specific reactivity, particles with a higher specific surface area induce more oxidative damage. The *quantitative* similarity analysis should thus be performed on concentration-response data provided as a mass-metric representation.

We have used comparisons of *in vivo* NOAEC data here to provide a biological relevance to these reactivity measurements, and in doing so have again found correlations and disparities. With similarity scores for BaSO₄ and CuO, for example, being low in all assays and likewise were considerably different in their *in vivo* NOAEC. However, this was not always the case, as reflected in various examples of the representative test materials in Table 1, but also in assessment of case study substances such as the Fe-based materials; differences between Fe-based materials were observed in the FRAS assay which were not portrayed in relation to *in vivo* results. If we set the acceptable limit to a reactivity similarity score e.g. above 0.6, the Fe-based materials would not be justified for grouping by tier 1 reactivity methods (Table 2), and the GRACIOUS framework would require to exclude certain NFs from the candidate group, or to use another tier 1 assay (with justification), or to escalate to higher tier testing, where in the specific case tier 3 (*in vivo*, Table 2) confirms grouping.

These observations lead us to conclude that although our use of *in vivo* NOAEC values can provide some level of assurance that the similarity confirmations made have merit and justify an implication of a potential hazard, it should be stressed that any resulting *in vivo* NOAEC may be a culmination of many contributing factors, while our similarity assessment is implicit to one, that being reactivity (in this case specifically ROS generation). This illustrates the earlier discussion of how a reactivity provides just one decision node of a complex IATA, with other consideration being important. For example, we observed BaSO₄ and CeO₂ as statistically similar and both non-reactive in all assays, however, they differ considerably in their *in vivo* NOAEC. There must be a reason to this, and another decision node of the IATA prevents the grouping of these two materials, since they are not similar in their dissolution rate (Keller et al., 2021b; Keller et al., 2020).

In case study assessments we have also directly included the biologically relevant range of reactivity through use of high and low reactivity representative test materials (Mn₂O₃ and BaSO₄, respectively). In general, this allowed for a strong agreement in how the data from each of the different reactivity assays was interpreted, with Fe-based substances being consistently found (in FRAS, DCFH₂-DA and Nrf2 activation) similar to BaSO₄ and dissimilar to Mn₂O₃; only EPR was in disagreement. For the other case studies (pigments and silica particles) there was also a good level of correlation found across the different assays, when using this range of high to low reactivity. Table 1 highlights that the least similar pairs of NFs reach similarity scores around 0.2, whereas Tables 2 and 3 highlight that pairs of NFs which actually have similar *in vivo* NOAEC values -and thus should be accepted as being similar- score between 0.46 and 0.95 in their similarity of reactivity, depending on the chosen assay.

Table 4

Comparison of concentration-response observations according to different reactivity assays; x = concentration-response observed, o = no concentration-response observed, xo = some but not all case study NFs provided a concentration-response, – = not measured in this assay.

	FRAS	DCFH ₂ -DA	EPR-Tempone-H	Nrf2 activation
Representative test materials				
CuO	x	x	x	x
Mn2O3	x	x	x	x
ZnO	x	o	o	x
CeO ₂	x	o	o	o
BaSO ₄	o	o	o	o
Case studies				
Fe-based	x	xo	x	o
Pigments	–	o	x	o
Silica	x	o	x	o

When *excluding* from the similarity analysis the least similar pair of representative materials, which represent the biologically relevant range, it was possible to tease out sensitive details *within* individual case studies. For example, when considering individual Fe-based substances the extent of similarity of two Fe particles (Fe₂O₃-nanoA and Fe₂O₃-Larger) was shown to differ considerably across different assays, with a high level of similarity shown in the DCFH₂-DA and Nrf2 assays, slightly lower level of similarity in EPR, and what can be considered as closer to dissimilar in the FRAS assay. With these observations in mind we would suggest that there are different purposes of conducting such analysis under both these conditions: i) assessment of similarity for regulatory purposes must include the representative materials of high and low levels of reactivity to align findings to known benchmarking values; ii) for mechanistic studies, or to identify trends that can guide Safer-by-Design optimisations, one may decide to assess NFs independently from these benchmark values to allow more sensitive assessment.

The Bayesian similarity algorithm could also be used for *in vivo* dose response to quantify similarity of the tier 3 results. The reactivity similarity assessment calibration would then be more robust, and our methodology could be transferred for use with other data.

In summary, this work demonstrates that the grouping of candidate NFs with regard to the similarity of their surface reactivity can be justified or rejected by well-established, partially ISO-standardised assays and by a novel but transparent, easily reproduced algorithm. The data matrix must include materials that represent high and low reactivity—typically two NFs of other substances—and must be filled by only one assay for all candidate NFs and the representative materials.

Declaration of Competing Interest

The authors declare the following financial interests/personal relationships which may be considered as potential competing interests:

DAS, DAE, KW, KS, WW are employees of BASF SE, a company producing and marketing nanomaterials.

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Appendix A. Supplementary data

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