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Alternative digestion strategy for Ti, Zr and Hf oxides: eliminating hydrofluoric acid†

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Group IV metal oxides have a broad impact on the environment and human health due to their diverse applications in industry, consumer products and biomedicine. However, their chemical inertness poses significant challenges for accurate quantification in biological matrices, which is essential for assessing biodistribution, toxicity, and regulatory compliance. Traditional digestion methods often rely on hydrofluoric acid (HF), a hazardous reagent requiring specialized handling and infrastructure. Here, we present an alternative, HF-free microwave assisted digestion protocol for group IV metal oxides in biological contexts, utilizing sulfuric acid/water/hydrogen peroxide mixtures to achieve complete solubilization across nano- to microscale particles. The method's efficacy was evaluated on various commercially available TiO₂, ZrO₂, and HfO₂ powders. Optimization of digestion parameters, including acid-to-peroxide ratios, temperature, and reaction time, led to recoveries exceeding 90% for all tested materials. Notably, higher temperatures and extended digestion times were required for larger particles and higher atomic number oxides, reflecting the increased metal–oxygen bond dissociation energies. The method's applicability was further demonstrated through successful quantification of spiked nanoparticles in human cancer cells and bovine liver tissue, with detection limits down to ~1 ppb and achieving recoveries within 80–100%, maintaining sample stability over four weeks. Comparative analysis with HF-based digestion revealed comparable sensitivity and detection limits using inductively coupled plasma optical emission spectrometry (ICP-OES), with the HF-free method offering a safer and more accessible alternative for routine laboratory analysis. This validated protocol facilitates accurate quantification of group IV metal oxides in complex biological matrices, supporting preclinical and clinical studies while mitigating the risks associated with HF usage.

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Introduction

Group IV metal (Ti, Zr, Hf)-containing materials are utilized in various biomedical and consumer-good applications, including cancer therapy, dental and orthopaedic implants, pigments, catalysts and coatings.^{1–10} This is primarily due to their biocompatibility and desirable mechanical, electrical or (physico)chemical properties. For instance, Ti and Ti alloys (*e.g.*, Ti₆Al₄V) are employed in long-lasting, load-bearing medical

implants due to their elastic modulus, which closely resembles that of bone, and their naturally occurring oxide surface. Zirconium oxide (ZrO₂) serves as a dental ceramic implant material owing to its white color, high biocompatibility, low thermal conductivity, toughness, and low bacterial attraction.^{11,12} Although these materials are typically used as bulk implants, it is now well documented (*e.g.*, refs. 13–15) that wear particles are released and accumulate in surrounding tissues, highlighting an increasing need for biodistribution analysis and a better understanding of their material fate. In nanoparticulate form, group IV metal oxides and metal–organic frameworks (MOFs) show promise as anticancer agents due to their photocatalytic, radiocatalytic, or X-ray absorption properties.¹⁶ For example, nano-TiO₂ exhibits significant photocatalytic activity and can be used in photodynamic cancer therapies.¹⁷ Similarly, Zr- and Hf-carrying nanomaterials demonstrate promising X-ray radiation-based anticancer effects.^{18–20} Some of these materials are under clinical investigation,^{21,22} or have received regulatory approval for pharmaceutical applications (*e.g.*, by the FDA).²³ All in all, the various

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use cases in research suggest group IV metals will have increasing impact in the future.

For clinical translation, it is imperative to gain insights into the stability, biotransformation, uptake behavior, and bio-distribution of these materials. Additionally, metal analysis in tissues surrounding medical implants is necessary to understand tissue reactions and the release of metal ions or wear particles.²⁴ Therefore, there is a growing need to quantify these materials in different complex matrices such as culture media, blood, cells and tissues. This is also reflected in demands from regulatory authorities such as the FDA,²⁵ that emphasize bio-distribution and uptake studies, especially for non-biodegradable materials.

However, non-validated methods are often used in the biomedical community (e.g. aqua regia for Hf^{26,27}) or actual concentrations are not even determined.²⁸ Validated methods for determining group IV oxides are often either hazardous or labour-intensive and therefore often not performed or implementable in standard labs. Due to passivation, TiO₂ as well as ZrO₂ and HfO₂ do not dissolve in cold mineral acids with the exception of hydrofluoric acid (HF).²⁹ Consequently, most common digestion protocols involve HF in combination with other acids, such as nitric acid (HNO₃) to break down the oxides and stabilize the analyte ions as fluoride complexes. The use of HF necessitates stringent safety measures and trained personnel as it poses significant health risks upon skin contact and requires immediate medical action upon an incident. Due to these risks and associated costs, alternative digestion methods have been explored, particularly for TiO₂, one of the most studied nanoscale inorganic materials. Most commonly, fusion reactions have been shown to break down TiO₂. For example, SiO₂ has been determined in a TiO₂ matrix using alkali fusion with KOH and boric acid, followed by dissolution in dilute HCl.³⁰ Many other methods rely on fusion with sulfates. For instance, TiO₂ has been determined in sunscreens using a three-step sequential microwave digestion in a 1 : 1 mixture of HNO₃/HCl followed by fusion in a crucible with KHSO₄ and subsequent dissolution in H₂SO₄.³¹ Furthermore, ammonium persulfate fusion in a crucible followed by soaking in 2% HNO₃ and subsequent hot plate boiling was employed to determine TiO₂ in water and wastewater.³² Titanium dioxide has also been analyzed in chewing gum by heating a sulfuric acid and catalyst mixture to 400 °C for 2 hours in a Kjeldahl unit, followed by dilution to 10% H₂SO₄ for measurement.³³

The success of such sulfate-based alternative methods is not surprising, since TiO₂ in concentrated H₂SO₄ is known to form Ti(IV)sulfate (Ti(SO₄)₂), which is colorless and dissociates in water. Titanium(IV) also forms different complexes in sulfuric solutions, e.g. [Ti(OH)₂]²⁺, [Ti(OH)₃]⁺, [Ti(OH)₂(HSO₄)]_{aq} and [Ti(OH₃(HSO₄))]_{aq}.²⁹ Similarly, Zr and Hf form sulfates in sulfuric solutions, including (Zr(SO₄)₂·4H₂O, Hf(SO₄)₂·4H₂O, Zr(OH)₂·SO₄ and Hf(OH)₂·SO₄·H₂O.³⁴ Therefore, generating sulfate complexes presents a promising route for solubilizing group IV oxides without using HF. For instance, Ma *et al.* used H₂SO₄ to extract Zr and Hf with recoveries of 89.1 and 81.2%, respectively, from a zirconosilicate.³⁵ Furthermore, Watkins *et al.* demonstrated in 2018 that using H₂SO₄/water mixtures and heating to

110 °C for 8 hours effectively digested nano-TiO₂ in various matrices including water, fish tissue, periphyton and sediment.³⁶ Based on these results, we evaluated this approach in a previous study, where we achieved recoveries between 96 and 107% for Ti, Ti/Zr and Hf-containing metal-organic frameworks.³⁷

Encouraged by these results, we further evaluated, developed and validated this HF-free digestion method also for the oxides, which were expected to be more difficult to solubilize. In this study, we investigated and validated the use of a sulfuric acid-hydrogen peroxide-based microwave digestion to quantitatively dissolve group IV oxides in nano- and microparticulate form as a cost effective, quick and readily available alternative to HF and fusion-based methods. We assessed the digestion performance of TiO₂, ZrO₂ and HfO₂ powders with different, clinically and industrially relevant primary particle sizes from several nm up to microns. After optimizing digestion parameters such as temperature, digestion time and acid/peroxide ratios with the parent powders we finally optimized and demonstrated the applicability of the validated method to quantify these materials in cancer cells and tissues.

Materials and methods

Particles, chemicals and reagents

For our experiments, we chose commercially available materials where applicable to allow for accessibility for future comparative experiments. Most of the metal oxide powders were available from Sigma Aldrich. The respective product and CAS numbers can be found in Table S1 in the ESI.†

All reagents were of analytical grade unless stated otherwise. Concentrated HNO₃ (69%), H₂SO₄ (97%), H₂O₂ (30%) as well as HF (40%) were obtained from Sigma Aldrich. Ultrapure water (18.2 MΩ) was obtained through an ELGA Chorus purification device. NIST-traceable certified reference materials (CRM) for instrument calibration were obtained from inorganic ventures.

Digestion procedures

In total, we evaluated five different digestion procedures and compared them to a combination of HNO₃, H₂O₂ and HF as the “gold standard” method of reference for the selected materials.

Method I: 1.5 mL HNO₃ (65%) and 1 mL H₂O₂ (30%) were added to the weighted particle powder in a quartz glass tube. The samples were digested in 10 mL quartz tubes closed by a PTFE cap with pressure-exchange opening in a pressurized microwave (TurboWAVE, MLS GmbH) at 120 bar pressure and 200 °C for 10 min, after an initial ramping phase from room temperature to 200 °C of 12 min.

Method II: 1 mL ultrapure H₂O, 1.5 mL H₂SO₄ (97%) and 1 mL H₂O₂ (30%) were added to the weighted particle powder in the quartz digestion tubes. The samples were digested in the pressurized microwave at 120 bar pressure and 200 °C for 10 min, after an initial ramping phase from room temperature to 200 °C of 12 min.

Method III: 1 mL ultrapure H₂O, 1.5 mL H₂SO₄ and 1 mL H₂O₂ were added to the weighted particle powder in the quartz



digestion tubes. The samples were digested in the pressurized microwave at 120 bar pressure and 250 °C for 30 min, after an initial ramping phase from room temperature to 250 °C of 12 min.

Method IV: The particle powders were digested in 1.5 mL H₂SO₄ and 1 mL H₂O₂ using the otherwise same procedure as in method III. Since the digestion matrix is an acid piranha solution requiring special precautions,³³ we first placed all quartz digestion tubes containing the samples into a cold water bath, then added H₂SO₄ and finally, slowly, H₂O₂. Both reagents did not immediately mix and react, and a short agitation using a vortex was necessary. After careful agitation, the samples were immediately placed back into the cold water bath, as in all cases an exothermic reaction could be observed. The reactions observed were never violent, *i.e.* we did not observe heating to >> 100 °C or very extensive foaming/effervescence or even deflagration.

HF reference method: For comparison, the HF-digestion was performed by adding 2 mL HNO₃, 1 mL H₂O₂ and lastly 0.3 mL HF (40%) in a protected HF-facility to pre-weighted particle powders. Samples were digested in Teflon tubes at 250 °C and 120 bar for 10 min after a temperature ramping time of 15 min using an ultraCLAVE (MLS GmbH) microwave.

After any digestion method, the samples were transferred to conical 50 mL polypropylene test tubes and filled to the mark with ultrapure water and measured without further dilution (unless stated otherwise) using an inductively-coupled plasma optical emission spectrometer (ICP-OES). All measured concentrations and recovery calculations can be found in the ESI, in Tables S3–S7.†

Instrumentation

For all measurements, an Agilent 5110 ICP-OES was used. Sample introduction was performed *via* an SPS4 autosampler connected to a glass concentric nebulizer followed by a glass cyclonic double pass spray chamber. Operating parameters of the instrument are summarized in Table S2 (ESI).† Calibration curves for all elements of interest were made from single element standards (CRMs) obtained from inorganic ventures (Suisse TP product numbers: CGT11 (Ti), CGZR1 (Zr) and CGHF1 (Hf)) and were prepared in an acid solution matched exactly to the respective samples. Specifically, the calibration solution matrix for the HF-free digestion solutions consisted of 2.91% v/v H₂SO₄, corresponding to 1.5 mL H₂SO₄ (97%) per 50 mL total volume, whereas that for the HF digestion solutions contained 2% v/v HNO₃. Calibrations were additionally verified by measuring a certified multi-element standard (CCS-5, Inorganic Ventures). Recoveries for all elements were usually >96% in this case. Data were initially evaluated using the software ICP-expert (v7.4.1.10449, Agilent Technologies) to calculate mass concentrations from intensities before they were exported for further analysis.

Method development and spiking experiments

Initial method development was carried out with pure powders. To distribute a defined particle mass to the digestion tubes, first

a 1 mg mL⁻¹ aqueous dispersion of each powder was prepared by accurately weighing the powder into a small 2 mL glass vial (VWR). After weighing, ultrapure water was added so that the final mass concentration reached 1 mg mL⁻¹. From this dispersion, defined aliquots were then distributed to the digestion tubes after reaching a homogenous sample dispersion using vortex agitation and ultrasonication and it was proceeded as described above. Similarly, for cell spiking experiments, 10 µL of the homogenous nanoparticle dispersion was added to 100.000 HT1080 cancer cells, corresponding to <0.1 ng metal mass per cell or <10 wt%. Cells were previously taken from a routinely cultured cell flask. Cells were cultivated in minimum essential medium eagle (MEM, Sigma-Aldrich) supplemented with 10% fetal calf serum, 1% L-glutamine, 1% penicillin–streptomycin, 1% nonessential amino acids, and 1 mm sodium pyruvate. To detach cells from a flask, HT1080 cells (ATCC CCL121TM) were trypsinized and counted using a hemocytometer. One million cells per mL solution were recovered and 100.000 cells were added to each quartz digestion tube before nanomaterial dispersions were spiked. Standard culturing practices that were used are described elsewhere.³

For tissue spiking, the CRM BCR 185R (bovine liver, powder) was weighed in approx. 20 mg aliquots to the quartz tubes and mixed with 200 µL of the homogenous particle dispersion, corresponding to <1 wt% nanomaterial uptake; clinically relevant cell or tissue uptake scenarios are considered to be below 10 wt% of metal uptake.³⁹ For instance, for intratumoral nanoparticle injections of NBTXR3 (HfO₂) a total oxide mass of 1.8 wt% should be injected, corresponding to 1.5 wt% of Hf.⁴⁰

Method validation

The accuracy of the method was determined from best estimates for a true value obtained from own spiking experiments, as no certified reference materials for group IV oxides in biological material were available. Specifically, the accuracy was evaluated by relating the measured metal mass, $m_{\text{meas.}}$, to an expected metal mass (*i.e.*, best estimate for the true value), $m_{\text{exp.}}$, as follows:

$$\text{Recovery (\%)} = \frac{m_{\text{meas.}}}{m_{\text{exp.}}} = \frac{c_{\text{meas.}} \times V_{\text{s}}}{(m_{\text{MO}} \times (1 - \text{SR}) \times f_{\text{M}}) \times (V_{\text{s}})^{-1}} \quad (1)$$

$m_{\text{meas.}}$ = metal mass in the sample. $m_{\text{exp.}}$ = expected metal mass. $c_{\text{meas.}}$ = metal concentration in the sample. V_{s} = sample volume. m_{MO} = mass of the metal oxide powder. SR = water and organic surface residue mass fraction determined from thermogravimetric analysis. f_{M} = metal mass fraction of the respective metal oxide with 0.848 (M = Hf), 0.740 (M = Zr) and 0.599 (M = Ti).

Furthermore, we determined the method precision according to the following condition:⁴¹

$$\Delta x(\%) = \frac{|x_1 - x_2|}{\bar{x}} \times 100 \leq \sqrt{2} \times 2 \times s_{\text{v,rel}} \quad (2)$$

$\Delta x(\%)$ = relative difference of measured values to the mean. $x_{1,2}$ = measured values. \bar{x} = average of measured values. $\sqrt{2}$ = root of replicate number ($n = 2$). $s_{\text{v,rel}}$ = relative standard deviation – set to a limit of 2%.



In our case for method II–IV, this condition was fulfilled with $\overline{\Delta x} = 2.8\% \leq \sqrt{2} \times 2 \times 2\% = 5.65$, hence digestions were performed in duplicates. All experimentally determined as well as calculated values for each experiment can be found in Tables S3–S7 in the ESI.† Furthermore, sample stability was determined by measuring the same samples after a specific time with a fresh calibration.

Thermogravimetric analysis

Water and organic surface content was assessed using thermogravimetric analysis (TGA) and was performed with a NETZSCH TG 209 F1 instrument (NETZSCH-Gerätebau GmbH, Selb, Germany) and heating of the weighted sample at room temperature to 700 °C, with a heating rate of 10 °C min⁻¹ under nitrogen flow. The surface residue mass fraction (SR) was quantified from the weight loss at 700 °C.

X-ray diffraction

X-ray powder diffraction was performed on a PANalytical X'Pert Powder Diffractometer (Malvern Panalytical, UK) equipped with a copper X-ray source. Particle powders were placed on a low background sample holder and diffraction patterns were recorded from the rotating powder sample. Rietveld refinement for phase and grain size analysis was performed using Profex⁴² (Version 4.3.5).

Results and discussion

Particle characterization and reference digestion

Prior to determining element/oxide recoveries, all nano- and microparticles were characterized to confirm or estimate the theoretical element composition, their sizes and potential organic surface residues. X-ray powder diffraction (XRD) patterns indicated that all particles were crystalline (ESI, Fig. S1†). Based on XRD Rietveld refinement calculations, nanoparticles had grain sizes around 9–25 nm, while microparticles ranged from 26 to 107 nm (Table 1). Phase analysis identified typical metal oxide specific phases. While most particles were present in a major single phase, nano HfO₂ and nano TiO₂ displayed multiple phases. The analysis also confirmed the respective phases for micro anatase TiO₂ and

micro rutile TiO₂ as specified by the distributor. Transmission electron microscopy (TEM) corroborated the nanometer-scale sizes, ranging from 6 to 34 nm (d_{TEM}), suggesting a single crystalline domain per nanoparticle. Hafnium dioxide was the smallest nanoparticle, followed by TiO₂ and ZrO₂. The sizes of all microparticles were estimated using scanning electron microscopy (SEM) images and were well above the XRD grain sizes, implying multiple crystalline domains per microparticle. HfO₂, ZrO₂ and anatase TiO₂ exhibited average particle sizes between 150 and 280 nm (d_{SEM}). Rutile TiO₂ was the biggest microparticle with an average size of 860 nm. All nano- and microparticles were spherical to elliptical in shape (ESI, Fig. S2†). Given that nanomaterials possess a higher surface area than micromaterials, they are more prone to adsorbing moisture or other volatile organic substances on their surface. To estimate the amount of such surface adsorbed volatile substances, and with this also the inorganic metal oxide particle weight, thermogravimetric analysis (TGA) was performed (ESI, Fig. S3†). Microparticles exhibited minimal mass loss (<0.2%) after heating to 700 °C, as expected. In contrast, nanoparticles showed notable volatile organic surface residues ranging from 0.7 to 2.6 wt%. Interestingly, the amount of surface residues was highest for the smallest nanoparticle and decreased with particle size, which is in line with the higher surface area of smaller nanoparticles. These surface residues were accounted for in calculating the metal oxide purity correction factor (1-SR, eqn (1)). Subsequently, metal recovery investigations were performed using ICP-OES with the state-of-the-art HF digestion method as a reference for developing an HF-free digestion method. As expected, the HF method efficiently solubilized the selected materials, yielding recoveries between 93 and 106%. For nano HfO₂, we noted an increased recovery of slightly over 100%, which might be attributed to minor contaminations. However, all recoveries were compliant to FDA criteria; according to the FDA, an acceptable recovery range for an element in drug testing is between 80% and 120% of the declared content to ensure method accuracy and reliability.⁴³

Furthermore, we noted impurities of Zr in Hf oxide and *vice versa*. The metal impurity of Zr in HfO₂ materials was quantified to be <0.2% (micro) and <1.2% (nano) (Zr/Hf). The impurity of Hf in ZrO₂ materials, however, was around ~2% (Hf/Zr) (ESI,

Table 1 Material properties (crystal size and phase, particle size and surface residues) of the particles used in this study and calculated oxide recovery rate (eqn (1), ESI, Table S3) after HF digestion. Corresponding XRD spectra, EM images and size distributions can be found in the (ESI)

Material		Grain size value \pm ESD (nm)	Main phase	dTEM/SEM average \pm SD (nm)	Surface residue, SR mass loss (weight %)	Recovery HF reference method ($n = 2$) (%)
HfO ₂	Micro	68.1 \pm 0.4	Monoclinic (>99%) ⁴⁶	150.4 \pm 66	~0.1	95.4 \pm 3.7
ZrO ₂		26.4 \pm 0.2	Monoclinic (>98%) ⁴⁷	276.4 \pm 103	~0.2	96.4 \pm 1.2
TiO ₂		39.1 \pm 0.2	Anatase (>99%) ⁴⁸	218.5 \pm 76	~0.1	97.1 \pm 0.4
TiO ₂		127.1 \pm 1.2	Rutile (>96%) ⁴⁹	859 \pm 370	~0	96.8 \pm 1.2
HfO ₂	Nano	8.9 \pm 1.1	Monoclinic (>83%) ⁴⁶	5.8 \pm 1.5	~2.6	105.5 \pm 2.0
		12.4 \pm 3.5	Orthorhombic (>15%) ⁵⁰			
ZrO ₂		25.4 \pm 0.2	Monoclinic (>99%) ⁴⁷	31 \pm 7.5	~0.7	93.5 \pm 1.0
TiO ₂		14.8 \pm 0.1	Anatase (>93%) ⁴⁸	16.5 \pm 6.3	~2.0	95.5 \pm 0.4
		18.7 \pm 0.8	Rutile (>6%) ⁴⁹			



Table S3†). These impurities were excluded from the recovery calculations. Zirconium (Zr) and hafnium (Hf) commonly coexist in natural minerals⁴⁴ making them mutual impurities in each other's materials. Major Zr sources include Zircon (ZrSiO_4) and baddeleyite (natural form of ZrO_2) in which Zr and Hf coexist.⁴⁵ Zircon is the primary source of all Hf and contains up to 1.2 wt% Hf.⁴⁵ Since they are chemically nearly identical, separating these elements remains challenging⁴⁵ explaining the observed presence of impurities.

Digestion method optimization using pure powders

Initially, materials were digested using a commonly employed HF-free method comprising only HNO_3 and H_2O_2 (method I, Fig. 1) and compared to the reference HF digestion. This approach resulted in unacceptably low recoveries (<30%) for all microparticles. Similarly, nano ZrO_2 exhibited a low recovery (~20%) with this simple nitric acid/peroxide digestion. Consequently, this method also did not fulfill our precision criterion (see eqn (2) and ESI Table S4†), which is expected for such low recoveries. Interestingly, the recovery for nano TiO_2 and nano HfO_2 was found to be around 80%. Compared to microparticles, nanoparticles have a higher accessible surface area increasing digestion probability. Additionally, sufficiently small and colloiddally stable nanoparticles are likely to achieve good atomization efficiency in the ICP plasma without complete digestion. For instance, slurries with particles below ~5 μm often achieve good atomization in the plasma and recoveries comparable to digested solutions.^{51,52} Such size effects could

explain the high recoveries of TiO_2 and HfO_2 nanoparticles, which are smaller than nano ZrO_2 and all microparticles, using the simple nitric acid route.

Subsequently, nitric acid was replaced with an aqueous 60% H_2SO_4 mixture (digestion method II). This change significantly improved nanoparticle recoveries to acceptable levels (90–100%). Consequently, also the precision was improved considerably to ~1.6 $\Delta x(\%)$. Microparticle recoveries also improved, following an atomic number (Z) dependent trend. Notably, micro Anatase TiO_2 was recovered at nearly 100% while the rutile form achieved around 70%. Given that rutile TiO_2 had a much larger particle size, the differing recoveries between TiO_2 phases likely resulted from size and grain size differences in the micropowders.

Compared to TiO_2 , micro ZrO_2 and micro HfO_2 had lower recoveries (~50% and ~20%, respectively). This behavior suggests an atomic number-dependent recovery rate trend, consistent with the Z -dependent increase in metal–oxygen bond dissociation enthalpy for group IV transition metals and oxygen.⁵³

As microparticle recoveries improved but remained suboptimal, we implemented digestion method III, increasing the target temperature to 250 °C and maintaining it for 30 minutes before cooling down. This adjustment tremendously increased microparticle recovery rates; nanoparticle recovery remained between 90 and 100%, with a slight improvement observed for nano ZrO_2 . Additionally, recovery rates for micro rutile TiO_2 and micro ZrO_2 exceeded 90%. Only micro HfO_2 showed comparably low recovery rates of just below 80%.

To further improve micro HfO_2 digestion, in digestion method IV, we omitted additional water in the digestion matrix, resulting in a more aggressive acid piranha solution (1.5 mL 97% H_2SO_4 + 1 mL 30% H_2O_2) and increased peroxomonosulfuric acid (Caro's Acid) formation. Handling this solution generally requires special caution.³⁸ However, in this setting, we never observed excessive heat or gas development, so we consider working to be sufficiently safe. Under these final conditions, all micro- and nanoparticle recovery rates were found to be well above 90%.

Application of the optimized method for nanoparticle cell uptake quantification

In nanomedical applications, quantifying nanomaterial uptake into cells is crucial for understanding nanomaterial–cell interactions. To demonstrate the applicability of the optimized HF-free digestion method for cell uptake studies, we used digestion method IV to digest nanoparticles spiked to cancer cells (Fig. 2) and investigated the stability of the digested samples over four weeks. Cells were spiked with nanoparticle concentrations up to 10 wt%, representing the higher end of reported cell uptake.³⁹ All nanoparticles showed recoveries between 80% and 100% with a tendency of higher recoveries for TiO_2 , followed by ZrO_2 and HfO_2 . All samples remained stable over the course of 4 weeks, indicating that the metals remained stable within the liquid phase, allowing for (re-)evaluation within at least one month.

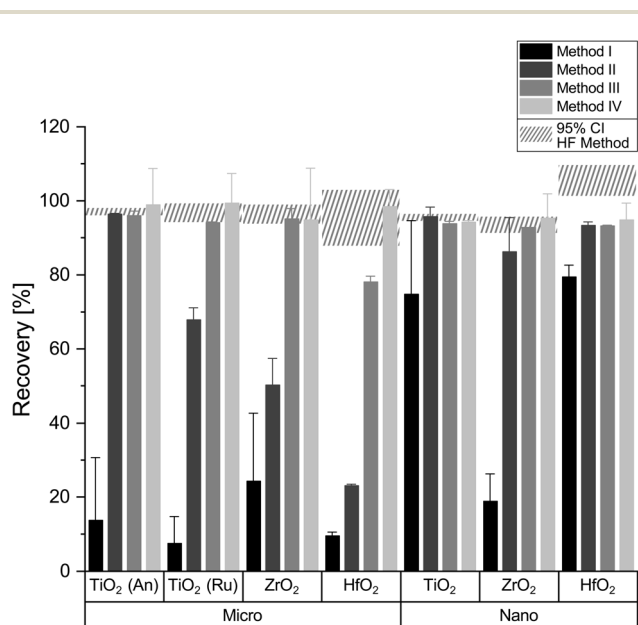


Fig. 1 Metal oxide recoveries for micro- and nanopowders and four different digestion methods determined from measuring the single elements (Ti, Zr, Hf) and calculated using eqn (1); method I: standard HNO_3 , 10 min@200 °C, method II: H_2SO_4 , 10 min@200 °C; method III: H_2SO_4 , 30 min@250 °C, method 4: piranha, 30 min@250 °C; An: anatase, Ru: rutile. Error bars signify $2\sigma_{\text{rel}}$ (~95% confidence interval (CI)) and the hatched area the 95% CI of the HF reference method for direct comparison.



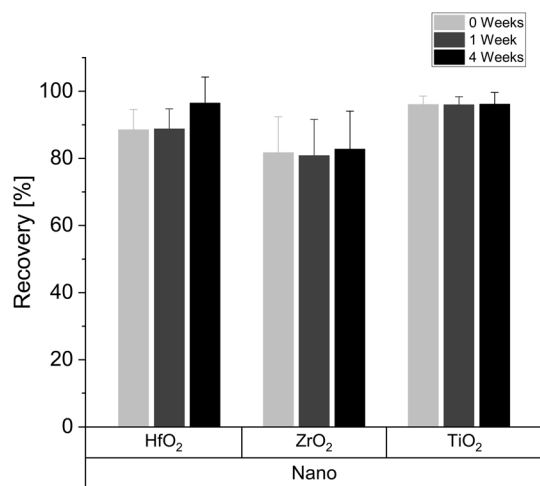


Fig. 2 Metal oxide recoveries for nanopowders spiked to cancer cells quantified directly after digestion (0 weeks) using digestion method IV as well as one and four weeks after digestion. Recoveries were determined from single element (Ti, Zr, Hf) quantifications and calculations using eqn (1). Corresponding nanomaterials were spiked to 100.000 cells per sample resembling a nanomaterial cell-uptake scenario between 1 and 10 wt%. Error bars signify $2s_{rel}$ (~95% confidence interval).

Application and further method optimization in spiked bovine liver tissue

For *in vivo* toxicity or efficacy studies, metal recovery in organs or target tissues from inserted or injected engineered nano- and micron-sized materials is essential. To simulate such scenarios, the optimized digestion method (IV) was applied to bovine liver tissue (BCR185r) samples spiked with nano- and micro-metal oxide powders.

Metal recoveries were excellent for micro- and nano-TiO₂ as well as for all nanopowders (>90%, Fig. 3a). However, the material-dependent decrease in recovery observed during the initial method optimization reappeared (e.g., micro: TiO₂

Table 2 Detection limit (DL), quantification limit (QL) and calibration slope (sensitivity) for each element (Ti, Zr, Hf) for the HF-free and the HF reference digestion method as quantified using ICP-OES

Limits of detection	HF-free method		HF method		
	Method (ppb)	Instrument (ppb)	Method (ppb)	Instrument (ppb)	
Ti	DL	0.69	0.21	0.37	0.34
	LOQ	2.10	0.63	1.12	1.03
	Slope	96.16 Int. ppb ⁻¹		96.03 Int. ppb ⁻¹	
Zr	DL	2.23	0.71	0.86	1.03
	LOQ	6.76	2.14	2.62	3.13
	Slope	88.08 Int. ppb ⁻¹		89.05 Int. ppb ⁻¹	
Hf	DL	10.52	1.90	1.95	1.69
	QL	31.87	5.76	5.89	5.13
	Slope	5.27 Int. ppb ⁻¹		5.35 Int. ppb ⁻¹	

(~100%) > ZrO₂ (~85%) > HfO₂ (~70%)/nano: TiO₂ (~100%) = ZrO₂ (~100%) > HfO₂ (~90%). Extending the reaction time from 30 to 60 minutes increased metal recoveries to >90% for ZrO₂ and HfO₂ micropowders as well as HfO₂ nanoparticles in liver tissue. Thus, more complex sample matrices require longer reaction times. Furthermore, these results suggest that the method might be also applicable to other types of aqueous and/or organic matter containing samples, such as for example wastewater or sewage sludge. Overall, HF-free digestion method IV provided excellent group IV metal recoveries also in *in vivo* scenarios for both nano- and microparticles. Additionally, digested samples displayed a very good long term sample stability post digestion of at least four weeks (Fig. 3b).

Sensitivity and detection limits of HF-free vs. HF digestion method

Finally, ICP-OES detection limits and sensitivity were compared between the HF-free and HF reference digestion method. The

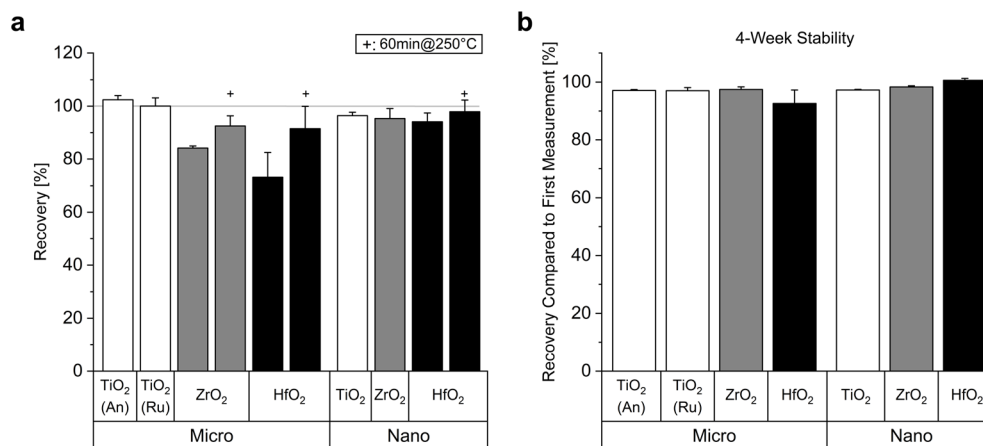


Fig. 3 (a) Metal oxide recoveries for micro- and nanopowders mixed with bovine liver tissue simulating an *in vivo* scenario after digestion using method IV or an adapted method IV with a longer microwave reaction time at 250 °C (60 instead of 30 minutes, indicated by the "+" sign). Recoveries were determined from single element (Ti, Zr, Hf) quantifications and calculations using eqn (1). (b) Four-weeks stability of digested samples plotted as recovery compared to the first measurement in (a) performed directly after digestion; An: anatase, Ru: rutile. Error bars signify $2s_{rel}$ (~95% confidence interval).



detection and quantification limits (DL and QL, respectively) were calculated per FDA guidelines: $DL = \frac{3.3\sigma}{\text{slope}}$ and $QL = \frac{10\sigma}{\text{slope}}$, where σ is the standard deviation of the blank response (here: emission intensity).⁴³ Sensitivities, indicated by calibration slopes (intensity/ppb), were comparable for both methods, following atomic number order (Ti: ~ 96 Int. ppb⁻¹ > Zr: ~ 88 Int. ppb⁻¹ > Hf: ~ 5 Int ppb⁻¹, Table 2). Consequently, detection and quantification limits followed this trend but were generally slightly lower for the HF method. While the instrument and method DLs and QLs of the HF method were very similar to each other, they showed a discrepancy to those of the HF-free method. This is because the standard deviation of the procedure blanks was generally higher for the HF-free method compared to the HF method. This behavior might be attributed to matrix differences (2.9% H₂SO₄ compared to 2% HNO₃) which can affect viscosity and, consequently, nebulization and transport efficacies in both methods. This is supported by the observation that the DL and QL can be lowered by roughly a factor 2–3 by doubling the readout time (ESI, Table S8†). Further improvements in DLs and QLs can be achieved by implementing tube pre-cleaning procedures to prevent analyte carryover between measurement campaigns and by increasing rinsing time between individual measurements. Nevertheless, given the comparable quantification sensitivities and the similar ranges of detection and quantification limits, we conclude that the HF-free method is a viable alternative to the reference HF digestion method which can be used in every standardly equipped analytical laboratory.

Conclusion

The reliable detection of group IV metal oxides is playing an increasingly important role in the biomedical as well as environmental fields. To digest group IV metal oxide materials, HF- or fusion-based methods are standardly used to date. Here, we have shown that sulfuric acid and hydrogen peroxide can be used as easily available, comparatively safe alternative HF-free digestion method for Ti, Zr and Hf metal oxides, both as pure powders and in aqueous environments with organic matter (cells, tissues). While nanomaterials were easier to digest, micron-sized powders required higher reaction temperatures, longer reaction times, and higher concentrations of peroxomonosulfuric acid for successful digestion. Digestion efficacy was also atomic number dependent. The proposed methods can be used to quantify the oxide elements in cells, organs and tissues from average particle sizes <6 nm up to 860 nm. ICP-OES revealed good linearity between 0.01 and 5 mg L⁻¹ (corresponding to 0.005–2.5 ng cell⁻¹, considering 100 000 digested cells per sample, or, 0.25–12.5 mg g⁻¹ of dried tissue, considering 20 mg digested tissue per sample) and common metal-dependent detection limits, comparable to those of an HF-containing reference method. The proposed HF-free method has fewer safety and infrastructural demands compared to HF digestions and allows cheaper and easier access to pre-clinical and clinical toxicity as well as efficacy studies with group IV metal oxide materials.

Data availability

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

Author contributions

LRHG: investigation, methodology, validation, formal analysis, data curation, writing – original draft, writing – review & editing; MR: formal analysis, writing – review & editing; IKH: resources, funding acquisition, writing – review & editing; AG: conceptualization, supervision, validation, writing – original draft, writing – review & editing.

Conflicts of interest

There are no conflicts to declare.

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