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# Determination of trace element concentrations in organic materials of "intermediate-thickness" *via* portable X-ray fluorescence spectrometry

Shubin Zhou, Dabed Qiuming Cheng, David C. Weindorf, De Biying Yang, Zhaoxian Yuang and Jie Yang \*\*David C. Weindorf, De Biying Yang, Thaoxian Yuang \*\*David C. Weindorf, De Biying Yang, De Biyin

This research explored the feasibility of using portable X-ray fluorescence (pXRF) spectrometry to quantify Fe, Cu, Zn, and As concentrations in dried and ground organic materials (fungi, vegetation, and animal tissues) with intermediate thickness. The experimental results indicated that pXRF determined concentrations of 39 organic samples (calibration set) with intermediate thickness decreased with increasing sample mass per unit area (also known as surface density or mass thickness) in a power function. The derived mass-correction model allows for correcting the matrix effect caused by analyzing samples with intermediate thickness and for predicting pXRF determined concentrations for infinitelythick samples. Then a simple linear regression (SLR) model was established based on the linear relationship between the pXRF and inductively coupled plasma mass spectroscopy (ICP-MS) determined concentrations of the organic samples. The SLR model allows for matrix effect correction caused by analyzing organic materials using pXRF soil mode and predicting the ICP-MS concentrations. The masscorrection and SLR calibration models were employed to predict the ICP-MS based trace element concentrations of 10 other organic samples of intermediate thickness (validation samples). The validation results were very good for Zn and As, and acceptable for Fe and Cu. The relative errors (REs) of the predicted Zn, As, Fe, and Cu concentrations were <18%, <22%, <34%, and <30%, respectively. Thus, pXRF shows strong potential for determining the trace element concentration in organic materials of intermediate thickness. The use of pXRF provides an appealing compromise between data quality and analytical time, and a potential alternative to laboratory analysis techniques such as ICP-MS.

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### Introduction

Portable X-ray fluorescence (pXRF) spectrometry is a technique capable of non-destructive, rapid, economical, and multi-elemental analysis. *In situ* pXRF analysis can provide elemental data in the field and assist in on-site decision making. *Ex situ* pXRF analysis provides a good compromise between data quality and analytical time. For *ex situ* pXRF analysis of organic materials (*e.g.*, vegetation and animal

tissues), drying is often recommended and should be done as rapidly as possible to minimize chemical and biological changes after sampling and washing.3 Drying and grinding can greatly improve pXRF data quality, with such processes and ex situ pXRF analysis possible in a field camp or laboratory. 4,5 Compared with other laboratory-based analytical methods such as inductively coupled plasma - optical emission spectroscopy (ICP-OES), ex situ pXRF analysis is more efficient since it saves the time required for sample digestion for wet chemistry analysis. Given the advantages of using pXRF in compositional analysis, pXRF has been widely used in the elemental assessment of various matrices, such as soils, 1,6-10 rocks, 11,12 water, 13,14 coal,15,16 dietary supplements,17 external markers (e.g., TiO2 and Cr<sub>2</sub>O<sub>3</sub>),<sup>17,18</sup> alloys,<sup>19,20</sup> nails,<sup>21,22</sup> ceramics,<sup>23,24</sup> compost,<sup>25,26</sup> fossils,27 marine litter,28 and vegetation.29,30 Physical (e.g., particle size, uniformity, and homogeneity) and chemical (differences in concentrations of interfering elements) effects, collectively known as matrix effects, are important considerations in pXRF analysis.

Considering the variable chemical composition of different types of samples, pXRF manufacturers developed different precalibration settings and analysis modes (e.g., geochem, soil, and

<sup>&</sup>quot;State Key Lab of Geological Processes and Mineral Resources, China University of Geosciences, Beijing 100083, China. E-mail: sinixyang@gmail.com

<sup>&</sup>lt;sup>b</sup>School of Natural Sciences, Faculty of Science and Engineering, Macquarie University, NSW, 2109, Australia

<sup>&</sup>lt;sup>c</sup>Bremick Preston Laboratory, Bremick Pty Ltd, Preston, NSW 2170, Australia

<sup>&</sup>lt;sup>d</sup>School of Earth Sciences and Engineering, Sun Yat-sen University, Zhuhai 519000,

<sup>&</sup>lt;sup>e</sup>Department of Earth and Atmospheric Sciences, Central Michigan University, Mount Pleasant, MI, 48859, USA

Daqing Oilfield Production Technology Institute, Daqing 163453, China

<sup>\*</sup>Institute of Resource and Environmental Engineering, Hebei Geo University, Shi Jiazhuang 050031, China

<sup>&</sup>lt;sup>h</sup>Institute of Earth Sciences, China University of Geosciences, Beijing 100083, China

mining).31 However, there is presently no available pXRF analvsis mode for an organic matrix. While soil mode can be used for the elemental assessment of organic samples, reductions in pXRF data accuracy may ensue<sup>2</sup> as soil mode was designed for quantifying elements in a soil matrix mainly composed of O, Si, and Al rather than an organic matrix rife with C, H, O, and N. Considering this, many researchers have established calibration models to correct for the matrix effect caused by the use of an alternate analytical mode. For example, Pearson et al. (2017) and Zhou et al. used soil mode to determine the elemental concentrations in water samples and corrected pXRF readings using simple linear regression (SLR) calibration models.14,32 Zhou et al. used mining mode to determine Fe and Si concentrations in an iron ore concentrate; and a SLR calibration model was developed to improve pXRF accuracy.33 Furthermore, SLR calibration models have been used for predicting the ICP-MS results for organic samples. For example, McGladdery et al. used soil mode to determine the elemental concentrations in vegetation samples.34 Validation statistics showed that pXRF could be used for determination of Zn and Cu even under field moist conditions, while dried and powdered sample analyses allowed for prediction of Pb, Fe, Cd, and Cu concentrations. Similarly, Zhou et al. used soil mode to determine the elemental concentrations in dried and ground leech samples.2 The validation results for 15 leech samples suggested that the established SLR calibration models substantially improved the accuracy for all studied elements ( $R^2 \ge 0.92$ ). However, very few studies have established whether different types of organic materials can be classified as a uniform matrix, and whether a singular calibration model can be applied universally to all

Organic materials feature high concentrations of light elements (C, H, O, and N), which are mostly transparent to X-rays.<sup>2</sup> Thus, the absorption of X-rays by organic matrices is smaller than that by other matrices (*e.g.*, rocks and soils) containing heavier elements. To some extent, X-ray absorption in organic materials is determined by trace metal concentrations.<sup>35</sup> To study the extent of sample self-absorption and the corresponding attenuation correction models, it is important to consider the mass thickness (surface density) of the sample pellets being analyzed (thin, intermediate-thickness, and infinitely-thick samples).<sup>35</sup>

types of organic samples.

For samples with very low sample mass per unit area (thin sample), the total mass-absorption coefficient of the sample is almost negligible, which means X-ray fluorescent intensity is linearly correlated to the concentration.<sup>35</sup> However, for thin samples in an organic matrix, the sample mass per unit area for heavier elements may be too low to form sample pellets.<sup>36</sup> For an infinitely thick sample, the fluorescent X-rays are fully absorbed within the sample; the thickness at which X-rays are fully absorbed is defined as the critical penetration depth (CPD). Further increase in thickness beyond the CPD will not cause variations in fluorescent X-ray intensity (concentration).<sup>35,37</sup> For samples with intermediate thickness, a change in mass thickness (sample mass per unit area) may cause variations in the emitted fluorescent X-ray intensity. Furthermore, different from soil and rock samples, organic materials generally have low

density. Potts et al. noted that the CPDs of X-rays increase with decreasing sample density.37 The CPDs of organic materials are generally a few centimeters, much higher than those of soil and rock materials.2,38,39 Therefore, pXRF analysis of organic samples with "intermediate-thickness" (e.g., several mm for light elements such as Mg and several cm for heavier elements such as Cu and Zn) is more common than for soil or rock samples. For XRF analysis of intermediate-thickness samples, Sitko detailed the quantitative X-ray fluorescence analysis of samples of less than "infinite thickness". 40 Studies conducted by the Sitko group also provided many approaches to correct matrix effects in X-ray fluorescence analysis of intermediatethickness samples, such as influence coefficient algorithms and the "two masses" (2M) method.41-43 This established the strong capability of XRF analysis of intermediate-thickness samples. For pXRF analysis of intermediate-thickness samples, some researchers have adopted thickness corrections for the analysis of vegetation and marine litter with intermediate thickness with the use of a "plastic mode" (an analytical mode with an integrated algorithm designed for plastic materials).28,44,45 In situ pXRF determined concentrations for F. ceranoides and F. vesiculosus were corrected for moisture content and thickness and the results suggest that corrected Zn concentrations were correlated with ICP-MS determined concentrations  $(R^2 = 0.664)$ . Gherase and Fleming and Fleming et al. investigated calibration methods to quantify elemental concentrations in nail clippings using pXRF.46,47 The introduction of a clipping mass-dependent factor to account for mass differences between samples was essential.22 Kagiliery et al. studied water influence on pXRF elemental analysis and proposed a statistical adjustment method enabling accurate prediction of elemental concentrations in water samples with a thickness of 4.19 mm. 13 The Cohen group first employed pXRF in biogeochemical mapping and quantitatively determined the X-ray penetration depth in cellulose filter papers (a proxy for wood).48 Ribeiro et al. investigated direct foliar analysis via pXRF; pXRF determined concentrations were found to decrease with increasing sample thickness.38 pXRF determined concentrations of Zn and Fe in dried, ground, and pressed leech sample pellets decreased with sample thickness in a power law relationship, which provides insights into accurate pXRF analysis for organic samples with intermediate-thickness.<sup>2</sup>

The objective of this pilot study was to demonstrate the feasibility of using pXRF to quantitatively determine elemental concentrations in different organic materials. For pXRF analysis, we hypothesize that different types of organic materials can be classified as a uniform matrix and that any matrix effects can be corrected using a universal calibration model.

## Materials and methods

## Theory

Many XRF researchers have investigated thickness corrections to calculate concentrations using characteristic X-ray intensities. 40,41,43 In pXRF analysis, research conducted by Sharma *et al.* suggested that coherent normalization using the Monte Carlo code may be a particularly robust normalization procedure in

y = ax + b (4)

pXRF determination of Zn in nail clippings.49 Gherase and Fleming and Fleming et al. investigated calibration methods to quantify elemental concentrations in nail clippings. 46,47 Given the existence of built-in algorithms in pXRF analyzers allowing a direct output of compositional data rather than raw counts of fluorescent X-rays, many scientists focus on calibration methods using pXRF determined concentrations as inputs. For example, Ribeiro et al. suggested that the effect of thickness of ground plant material on pXRF determined concentrations was well fitted by using a polynomial inverse model.38 Zhou et al.2 normalized pXRF determined concentrations (measured pXRF concentrations for sample pellets with different thicknesses normalized by the concentrations for infinitely thick samples) of Zn and Pb in dried and ground leech pellets. They were found to decrease with sample thickness in a power law relationship as per eqn (1)

 $\frac{C_{\text{intermediate-thickness}}}{C_{\text{infinitely-thickness}}} = e \times L^f$  (1)

where L is the sample thickness,  $C_{\text{infinitely-thickness}}$  is the corresponding pXRF determined concentration for samples with intermediate thicknesses,  $C_{\text{infinitely-thickness}}$  is the pXRF determined concentration for infinitely thick samples, and e and f are constants (f < 0). If the density of leech samples is constant, then  $L \propto m$ , where m is sample mass per unit area (surface density).

Therefore, the relationship between standardized pXRF determined concentrations and sample mass per unit area should follow eqn (2).

$$\frac{\text{MC}}{\text{TC}} = i \times m^k \tag{2}$$

where MC is defined as the measured pXRF concentrations for samples with different values of mass per unit area (intermediate-thickness samples), TC is defined as the pXRF determined concentrations (for infinitely thick samples), m is sample mass per unit area (surface density) and i and k are constants which are element dependent, and can be calculated through experiments.

Then, eqn (2) can be converted to eqn (3)

$$TC = \frac{MC}{i \times m^k} \tag{3}$$

which allows for predicting the pXRF determined concentrations for infinitely thick samples using m and MC as inputs.

Besides the matrix effect caused by samples with intermediate-thickness which can be corrected as per eqn (3), the matrix effect caused by the use of soil mode for elemental analysis of organic samples may also cause less accurate results.<sup>2,34</sup> Previous research suggests that a SLR model can effectively correct for the matrix effect and improve the accuracy of the results.<sup>2,29,34</sup>

Here, a SLR model was used to approximate the relationship between concentrations determined using pXRF and ICP-MS, respectively. The linear regression model was described using eqn (4): where x is the concentration determined using ICP-MS and y is the corresponding pXRF determined concentration for infinitely thick samples (also referred to as TC in eqn (3)). The value of a represents the slope of the line and b is the y intercept. Therefore, the regression can be expressed using eqn (5):

[pXRF readings] = 
$$a$$
[ICP – MS determined concentrations]  
+  $b$  (5)

with constants a and b determined using pXRF concentrations and ICP-MS results for the 39 samples in the calibration dataset. This equation then allows for adjustment of pXRF data for the 10 validation dataset samples whereby the final results are comparable with the ICP-MS results. eqn (5) was then converted to eqn (6):

[calibrated concentrations] = 
$$\frac{1}{a}$$
([pXRF readings] – b) (6)

which is simplified to eqn (7) as:

[calibrated concentrations] = 
$$p[pXRF \text{ readings}] + d$$
 (7)

where p=1/a and d=-b/a. Notably, the calibrated concentrations are ICP-MS based concentrations. eqn (7) represents the calibration model using pXRF readings as inputs and predicting ICP-MS based concentrations. This model was then used to calibrate pXRF by analysing 10 samples (validation set) for performance validation.

The CPDs for light elements are quite low (mm level) and a thickness of 1 cm seems to be thick enough. <sup>38,39,50</sup> Thus, heavy elements were targeted in the present research. This is because the variation between pXRF determined concentrations for heavier elements (with higher CPDs) and mass thickness (sample mass per unit area) should be easier to observe.

# Sample preparation

The 49 samples used for this pilot study consist of fungi, vegetation, and animal tissues (Table 1). All samples were ovendried at 60 °C for 72 h to remove moisture. Next, the dried samples were ground to pass through a 250  $\mu$ m sieve. The grinding time was adjusted according to the toughness of the samples. For example, the leech samples were ground for  $\sim$ 6 min ensuring that over 95% leech sample particles passed through a 250  $\mu$ m sieve. For mushroom samples, the grinding time was adjusted to  $\sim$ 3 min.

#### Measurement methods

A Niton XL3t 950 (Thermo Fisher Scientific) pXRF spectrometer was used for elemental analysis following the methodology reported by Weindorf and Chakraborty. The instrument was equipped with a silicon drift detector (SDD) and an X-ray tube with a maximum voltage/current of 50 kV/40  $\mu$ A and an Ag anode target excitation source. The diameter of the detection window was 8 mm. The dried and ground organic samples were

Table 1 Description of the 49 analysed organic samples

Fungi $(n = 17)$	Vegetation $(n = 16)$	Animal tissues $(n = 16)$
Pholiota nameko $(n = 2)^a$ Lepista nuda $(n = 10)^b$ Lentinula edodes $(n = 2)^a$ Suillus luteus $(n = 1)^a$ Pleurotus citrinopileatus sing $(n = 1)^a$ Panellus serotius $(n = 1)^a$	Leaf of Salix matsudana Koidz $(n = 1)^b$ Leaf of Rhamnus davurica Pall $(n = 1)^b$ Leaf of Sorbaria sorbifolia $(n = 1)^b$ Straw of Oryza sativa L. (rice) $(n = 3)^b$ Leaf of Cirsium japonicum $(n = 1)^b$ Purple potato $(n = 1)^c$ Cocoa powder $(n = 1)^c$ Matcha powder $(n = 1)^c$ Spices (ginger; citrus peel; white pepper) $(n = 3)^a$ Ginseng $(n = 1)^c$ Carrot $(n = 1)^c$ Spinach $(n = 1)^c$	Leech $(n = 3)^d$ Oyster $(n = 2)^c$ Shrimp $(n = 2)^c$ Codfish $(n = 1)^c$ Salmon $(n = 1)^c$ Pork liver $(n = 2)^c$ Goose liver $(n = 1)^c$ Chicken liver $(n = 3)^c$ Lamb liver $(n = 1)^c$

<sup>&</sup>lt;sup>a</sup> Purchased from a market in Weihe County, Harbin City, China. <sup>b</sup> Collected from the Bailing Cu-Zn deposit area in A'cheng District, Harbin City, China. <sup>c</sup> Purchased online at Taobao China. <sup>d</sup> Purchased from the Anguo Traditional Chinese Medicine (TCM) Market, Anguo City, China.

placed in a sample cup with a diameter of 23 mm and a height of 25 mm. The sample cup was placed in a test stand with a thin polyethylene film protecting the aperture of the pXRF spectrometer from the powered organic samples. Soil mode was used in this study, as established in previous studies used to determine element concentrations in various organic samples.<sup>2,34</sup> Fe, Cu, Zn, and As were determined under the main range scanning condition (50 keV and 40 µA) with a dwell time of 30 s; low and high energy ranges were disengaged. Each sample was scanned to produce 20 replicates. "System selfcheck" was performed to monitor spectral interference.51 "System self-check" is a factory calibration check used for confirming that the instrument is operating within resolution and stability tolerance (e.g., no shifts in energy line positions, regions of interest, or shift in gain control due to temperature changes).52 For quality assurance (monitoring signal drift), analysis of a certified reference material (CRM) (CRM Soil 180-649 supplied by Thermo Fisher Scientific) was conducted within every four samples runs. Elemental recovery percentages (PXRF determined/CRM reported) for As, Zn, Ni, Fe, Mn, Cr, K, and Ba were: 81-115%, 83-95%, 104-226%, 95-97%, 70-84%, 94-132%, 70-75%, and 72-88%, respectively. Notably, the certified As concentration in CRM 180-649 is 10.5 mg kg<sup>-1</sup>; pXRF readings near the limit of detection (LOD) are often less accurate.51

The samples were subjected to ICP-MS analysis at ALS Minerals.53 To this end, the samples were digested using a HNO<sub>3</sub>/HCl acid mixture for 8 h at 20 °C. The digestion vessels were then transferred to a EG 20A-PLUS hotplate produced by LabTech (85 °C for 15 min and then 115 °C for 2 h). After digestion, when the vessel cooled to room temperature, HCl acid was added. Lastly, the prepared samples underwent ICP-MS analysis. A PerkinElmer Elan 6000 instrument was used for ICP-MS analysis. The LODs of the ICP-MS method for Fe, Cu, Zn, and As are 1 mg kg<sup>-1</sup>, 0.01 mg kg<sup>-1</sup>, 0.1 mg kg<sup>-1</sup>, and 0.005 mg kg<sup>-1</sup>, respectively. ICP-MS measurement accuracy was confirmed by measuring certified reference materials (GL-03 leaves from Eucalyptus tree supplied by ALS Global Brisbane) at regular intervals. The certified values for Fe, Cu, Zn, and As in reference material GL-03 are 103 mg kg<sup>-1</sup>, 3.37 mg kg<sup>-1</sup>,

14.7 mg kg<sup>-1</sup>, and 0.293 mg kg<sup>-1</sup>, respectively. The recovery percentages (ICP-MS determined/CRM reported) for Fe, Cu, Zn, and As were 102.9-106.8%, 97.3-107.1%, 98.0-106.1%, and 100.3-106.5%, respectively. Furthermore, replicate ICP-MS measurements were conducted on random samples to observe the standard deviations of the ICP-MS measurements. Standard deviations (and relative standard deviations) of replicate measurements can reflect the uncertainty of ICP-MS analysis. The average relative standard deviations of replicate ICP-MS measurements were 4.00%, 2.02%, 1.61%, and 2.76% for Fe, Cu, Zn, and As, respectively.

#### Procedures and pXRF analysis

To determine the constants i and k in eqn (2), experiments were conducted as follows. The dried and ground organic samples were progressively added to sample cups and underwent pXRF analysis. As an example, for a mushroom sample (YHM-3-F), 0.256 g of the sample was added to a sample cup, and then the sample cup was lightly tapped on a table by hand for  $\sim$ 40 seconds ( $\sim$ 30 times). This process ensures homogeneity of the sample in the sample cup. The table was covered with a piece of plain paper to prevent potential contamination to the film sealing the sample cup. Then the prepared sample was subjected to pXRF analysis. The sample mass in the sample cup was increased to 0.549 g, and subjected to pXRF analysis again. The sample mass in the sample cup was progressively increased to  $\sim$ 3.1-3.5 g, since pXRF determined concentrations of the target elements varied little when the sample mass in the sample cup reached ~3 g. The pXRF determined concentrations for the sample with higher mass (all >3.1 g) in the sample cup (with a diameter of 2.3 cm) were regarded as "infinitely thick" samples. The sample mass per unit area was calculated as per eqn (8):

$$m = \frac{4M}{\pi D^2} \tag{8}$$

where m is sample mass per unit area; M is sample mass in the sample cup; D is the diameter of the sample cup.

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#### Calibration and performance verification

The samples were randomly split into an  $\sim$ 80% calibration set (39 samples) and a  $\sim$ 20% validation set (10 samples) a priori. The constants i and k in eqn (2) and (3) were calculated through experiments of samples in the calibration set, which allows for correction of the matrix effect caused in the analysis of intermediate-thickness samples.

Constants p and d can be calculated from constants a and b (slopes and intercepts of the SLR equations between ICP-MS and pXRF determined concentrations for infinitely thick samples in eqn (4)) as per eqn (7), which corrects the error caused by analyzing organic materials using soil mode. Therefore, successful calibration of the pXRF determined concentrations of organic materials with intermediate thickness should combine eqn (3), (7) and (9) as follows:

[calibrated pXRF reading] = 
$$p\left[\frac{MC}{i \times m^k}\right] + d$$
 (9)

where MC is the measured pXRF concentration for samples with different values of mass per unit area in mg kg<sup>-1</sup> (intermediate-thickness samples), m is the sample mass per unit area in g cm<sup>-2</sup> defined as per eqn (8), [calibrated pXRF reading] denotes the predicted ICP-MS based concentrations, and other terms (p, d, i and k) are constants aforementioned in egn (2) and (7).

To evaluate the effects of calibrations, the relative errors (REs) [calibrated pXRF reading] were calculated. The RE was defined as per eqn (10) as:

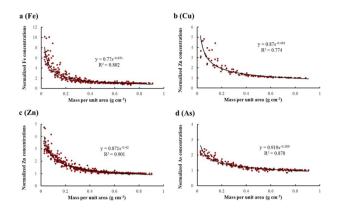


Fig. 1 Relationships between the normalized portable X-ray fluorescence (pXRF) concentrations (measured pXRF concentrations for samples with different values of sample mass per unit area normalized by the concentrations of the infinitely thick samples) of Fe (a), Cu (b), Zn (c), and As (d) and the sample mass per unit area (surface density).

correlation, especially considering that Fe concentrations for nearly 1/3 of samples were <200 mg kg<sup>-1</sup> (~twofold of the Fe LOD),51

Fig. 1b shows the relationship between the standardized pXRF determined concentrations for Cu and sample mass per unit area. The calibration set only contains 10 samples due to the inherent low Cu concentrations in the samples. The values of constants i and k were 0.87 and -0.491 for Cu, respectively (Fig. 1b). The  $R^2$  value for the Cu relationship was 0.774. Notably, Cu concentrations for four samples were <60 mg kg<sup>-1</sup>

$$RE = \frac{|[calibrated pXRF reading] - [ICP-MS determined concentration]|}{[ICP-MS determined concentration]}$$
(10)

# Results and discussion

# Standardized pXRF determined concentrations versus sample mass per unit area (surface density)

Standardized pXRF determined Fe concentrations decreased with mass per unit area in a power law relationship (Fig. 1a). The standardized pXRF Fe concentrations in samples with low sample mass in the sample cup (mass per unit area) were significantly higher than those for "infinitely-thick" samples. For example, the measured Fe concentration for lamb liver was 1197.3 mg kg<sup>-1</sup> with a sample mass of 0.119 g in the sample cup (0.0286 g cm<sup>-2</sup>), while the measured Fe concentration for the sample with infinite-thickness was 118.3 mg kg<sup>-1</sup>. The significant overestimation of Fe concentrations in samples with low sample mass indicates that the algorithm in pXRF analyzers incorrectly calculated the Fe concentrations and the original pXRF determined concentrations for intermediate-thick samples are not reliable. The values of constants i and k in eqn (5) were 0.770 and -0.651 for Fe, respectively (Fig. 1a). The coefficient of determination  $(R^2)$  was 0.802, indicating a good

and close to the LOD for Cu ( $\sim 20 \text{ mg kg}^{-1}$ ). The  $R^2$  values increased to 0.900 after removing the four samples.

Fig. 1c shows the relationship between the standardized pXRF determined concentrations for Zn and sample mass per unit area. The values of constants i and k were 0.871 and -0.420for Zn, respectively (Fig. 1c). Zinc concentrations were detected in all 39 samples in the calibration set and the LOD for Zn was <12 mg kg $^{-1}$ . The correlation was better for Zn with an  $R^2$  value of 0.901, suggesting that the proposed regression model (eqn (5)) fitted quite well; the calibration model (eqn (8)) should have strong potential to correct the effect caused by sample thickness. The strong correlation for Zn was likely due to the robust sensitivity for Zn detection via pXRF and the relatively high Zn concentration in the samples used for calibration.54

The standardized pXRF As concentrations also decreased with increasing sample mass per unit area with an  $R^2$  value of 0.878 (Fig. 1d). The values of constants i and k were 0.919 and -0.289 for As, respectively. The law's exponent (constant k) for As was the highest among those of all studied elements, which

means the overestimation of As concentrations by pXRF in samples with low sample mass was less significant.

If the normalized concentration (MC/TC) is set at 1.01, then xrepresents the mass per unit area (surface density) which caused only a 1% overestimation of pXRF determined concentrations in the infinitely thick samples. This can be roughly regarded as the minimum mass per unit area of the infinitely thick samples under pXRF analysis (in which the X-rays are fully absorbed). According to the functions given in Fig. 1, the minimum mass per unit area causing full X-ray absorption was  $0.659 \mathrm{\ g\ cm^{-2}}, 0.738 \mathrm{\ g\ cm^{-2}}, 0.703 \mathrm{\ g\ cm^{-2}}, \mathrm{and} \ 0.721 \mathrm{\ g\ cm^{-2}}$  for Fe, Cu, Zn, and As, respectively. Theoretically, the critical penetration depths of X-rays (also stands for sample mass per unit area if density is a constant) should increase with increasing atomic number.37,55 However, the calculated minimum sample mass per unit area which fully absorbs X-rays is greater for Cu than for Zn. This is due to the difference between the sample set for Cu and Zn: the set for Cu only contains 10 samples, while the set for Zn contains all 39 samples. If the same set was applied for Zn, the calculated minimum sample mass per unit area (surface density) would change to 0.730 g cm<sup>-2</sup>, very close to that of Cu (0.738 g cm<sup>-2</sup>), which might be explained by random errors or the limited sample size.

# PXRF determined concentrations *versus* ICP-MS determined concentrations for "infinitely thick" samples

As shown in Fig. 2, the pXRF determined concentrations for Fe, Cu, Zn, and As were all strongly correlated with the ICP-MS determined concentrations. The  $R^2$  values were 0.922, 0.960, 0.940, and 0.925 for Fe, Cu, Zn, and As, respectively, which were

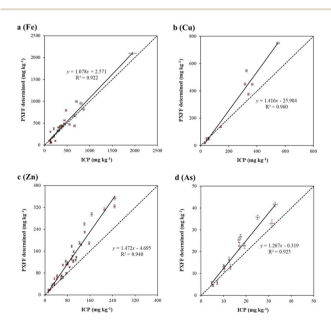


Fig. 2 Plots showing portable X-ray fluorescence (pXRF) vs. inductively coupled plasma mass spectroscopy (ICP-MS) determined Fe (a), Cu (b), Zn (c), and As (d) concentrations (mean  $\pm$  SD) for 39 organic samples. The solid line represents the best-fitting line of ordinary least squares regression and the dotted line represents the 1:1 line.

slightly higher than the  $R^2$  values reported by McGladdery et al. (Fe: 0.74; Zn: 0.93; Cu: 0.82)<sup>34</sup> and the  $R^2$  values reported by Turner et al. (Cu: 0.862; Zn: 0.959; As: 0.976).44 The particle size (<0.25 mm) in the present study was much smaller than those in the study carried out by McGladdery et al. (<2 mm), causing a higher degree of homogeneity and a higher R<sup>2</sup> value.<sup>34,56</sup> However, the  $R^2$  values for all studied elements were slightly lower than those reported by Zhou et al.,2 in which the elemental concentrations of dried and ground leech samples were determined using pXRF (Fe: 0.99; Cu: 1.00; Zn: 0.94; As: 0.95). This is likely because metal concentrations in leech samples can be high (e.g.  $C_{\text{Fe}} > 3500 \text{ mg kg}^{-1}$ ); highconcentration samples may have elevated the  $R^2$  values.<sup>2</sup> The slopes (95% confidence limits) for Fe, Cu, Zn, and As were 1.078  $\pm$  0.121, 1.416  $\pm$  0.235, 1.472  $\pm$  0.123, and 1.267  $\pm$  0.209, respectively. In this case, systemic errors were observed for Cu and Zn (and possibly As), whose slopes were different from 1. Overestimation of pXRF determined concentrations was likely due to the use of soil mode in the elemental determination of organic materials.

Overall, the correlation between ICP-MS determined concentrations and pXRF determined concentrations is quite solid, especially considering that the samples were of different species. This further demonstrates that different types of organic materials (fungi, vegetation, and animal tissues) may be regarded as a universal matrix in pXRF analysis; calibrations based on the same SLR models are then applicable for organic materials. Development of an organic mode suitable for analysis of organic materials is highly recommended to equipment manufacturers and further testing on a broader array of organic matrices is suggested to further refine the proposed universal organic matrix for pXRF analysis.

#### Validation of calibration models

Given the strong correlation between standardized pXRF determined concentrations and sample mass per unit area, as well as the correlation between pXRF and ICP-MS determined concentrations, eqn (9) was used to predict the elemental concentrations determined using ICP-MS in 10 samples of the validation set. The constants i and k can be obtained from the power-law functions given in Fig. 1. The constants p and p were calculated from constants p and p are slopes and intercepts of the linear regression equations in Fig. 2. Then, the calibration models for predicting elemental concentrations determined by ICP-MS in organic materials with intermediate thickness were obtained using eqn (11)–(14) as follows:

[calibrated Fe reading] = 
$$0.928 \times \left[ \frac{\text{MC}}{0.77 \times m^{-0.651}} \right] - 2.385$$
 (11)

[calibrated Cu reading] = 
$$0.706 \times \left[ \frac{\text{MC}}{0.87 \times m^{-0.491}} \right] + 18.294$$
 (12)

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[calibrated Zn reading] = 
$$0.679 \times \left[ \frac{MC}{0.871 \times m^{-0.42}} \right] + 3.19$$
 (13)

[calibrated As reading] = 
$$0.789 \times \left[\frac{\text{MC}}{0.919 \times m^{-0.289}}\right] + 0.252$$

where MC is the measured pXRF concentration for intermediate-thickness organic samples in mg kg<sup>-1</sup>, and m is the sample mass per unit area in g cm<sup>-2</sup>.

The concentrations found using eqn (11)–(14) were then compared with those determined using ICP-MS; relative errors were calculated. The relationship between the sample mass per unit area and relative errors is shown in Fig. 3.

Better performance was observed for Zn (Fig. 3c). PXRF was effective in the analysis of all 10 samples of the validation set. The REs of the predicted Zn concentrations were all <18% in 105 measurements, with an average of 8.9% (Fig. 3c). PXRF only detected As in three samples. The REs of the predicted As concentrations were <22% in 35 measurements (<20% for 95% measurements), with an average RE of 9.1% (Fig. 3d). However, the validation results for Fe and Cu were even worse. For Fe, pXRF was effective for seven samples. The REs of the predicted Fe concentrations were <34% in 61 measurements, with an average RE of 11.5%. The less robust validation results for Fe were expected, since pXRF is more sensitive for heavier elements due to their higher excitation energy.7 The fluorescence yield characterizing the elements is poor for light elements due to the limited number of electrons contained therein. As such, only high concentrations of light elements (e.g. Na, Mg, and Al) are detectable via pXRF. Furthermore, as shown in Fig. 1a, Fe concentrations may vary dramatically with sample mass per unit area in thin samples; a tiny change of sample mass may cause huge variations in the Fe concentration

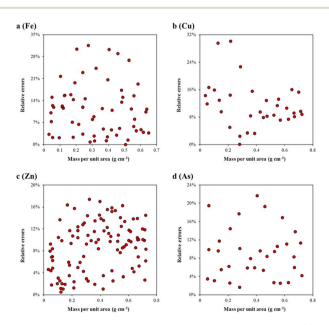


Fig. 3 Relationships between the relative errors of calibrated Fe (a), Cu (b), Zn (c), and As (d) concentrations and sample mass per unit area.

determined by pXRF, which coincided with the results obtained by Fleming *et al.*<sup>57</sup> Another source of errors may arise during experiments, since the powdered sample may not be uniformly distributed in the sample cup; geometrical placement is important in XRF analysis.<sup>47</sup> For Cu, pXRF was only effective for three samples. The REs of the predicted Cu concentrations were generally <30% in 33 measurements, with an average RE of 11.5%. Such results may be explained by the limited number of samples of the calibration set.

# Conclusions

This research introduced a calibration model allowing pXRF to quantify Fe, Cu, Zn, and As concentrations in organic materials (fungi, vegetation, and animal tissues) with intermediate thickness. The calibration model was divided into two parts:

The pXRF determined concentrations of Fe, Cu, Zn, and As in intermediate-thickness samples were found to decrease with increasing sample mass per unit area (surface density) in the sample cup in a power-law relationship. The coefficients of determination ( $R^2$ ) of the power-law regression models were 0.802, 0.774, 0.901, and 0.878 for Fe, Cu, Zn, and Zn, respectively. The derived mass-correction model allows correction for the matrix effect caused by analyzing intermediate-thickness samples.

The pXRF determined concentrations for infinitely-thick samples were linearly correlated with the ICP-MS determined concentrations. The  $R^2$  values of the SLR model were 0.922, 0.960, 0.940, and 0.925 for Fe, Cu, Zn, and Zn, respectively. The SLR calibration model allows pXRF determined concentrations for infinitely-thick samples to predict ICP-MS based concentrations.

The mass-correction model and the SLR calibration model were employed to calibrate the matrix effect caused by analyzing the intermediate-thickness organic samples (10 samples of the validation set). The corrected pXRF determined concentrations were compared with the ICP-MS determined concentrations. The validation results were very good for Zn and As, and acceptable for Fe and Cu. The REs of the predicted Zn, As, Fe, and Cu concentrations were <18%, <22%, <34%, and <30%, respectively. The validation results suggest that pXRF was capable of quantifying trace metal concentrations in organic materials with intermediate thickness. In future analysis, the sample mass per unit area (surface density) should be considered as the factor controlling pXRF determined concentrations rather than the sample thickness, since sample density may not be uniform in different organic matrices.

# Conflicts of interest

There are no conflicts to declare.

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