The applicability of spectroscopy methods for estimating potentially toxic elements in soils: state-of-the-art and future trends

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Abstract: Potentially toxic elements (PTEs) in soils pose severe threats to the environment and human health. It is therefore imperative to have access to simple, rapid, portable and accurate methods for their detection in soils. In this regards, the review introduces recent progresses made in the development and applications of spectroscopic methods for \textit{in-situ} semi-quantitative and quantitative detection of PTEs in soil and critically compares them to standard analytical methods. The advantages and limitations of these methods are discussed together with recent advances in chemometrics and data mining techniques allowing to extract useful information based on spectral data. Furthermore, the factors influencing soil spectra and data analysis are discussed and recommendations on how to reduce or eliminate their influences are provided. Future research and development needs for spectroscopy techniques are emphasized, and an analytical framework based on technology integration and data fusion is proposed to improve the measurement accuracy of PTEs in soil.

Keywords: Spectroscopy, analytical techniques, contaminated soil, potentially toxic elements.
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1 Introduction

Soil contamination by PTEs is a global concern as they can cause serious health problems to human and ecosystems (1, 2, 3). While it is usually difficult to list or specify all PTEs found in soil as there are over 40 chemical elements (4), a list of 21 elements is commonly used for PTEs. The list includes antimony (Sb), arsenic (As), barium (Ba), cadmium (Cd), chromium (Cr), cobalt (Co), copper (Cu), gold (Au), lead (Pb), manganese (Mn), mercury (Hg), molybdenum (Mo), nickel (Ni), selenium (Se), silver (Ag), thallium (Tl), tin (Sn), tungsten (W), uranium (U), vanadium (V) and zinc (Zn) (5). Among these elements, 8 PTEs, including Pb, Cr, As, Zn, Cd, Cu, Hg, and Ni are considered as the most commonly elements found at contaminated sites (6).

Metal ions are non-biodegradable and can accumulate in soft tissues, hence, they are considered as posing a significant risk to health, and cause of many toxic and physiological diseases to both humans and livestock (7, 8). This is the case particularly for, but not limited to, As, Cd, Hg, and Pb, even they are present in relatively low concentrations (9–11). Thus, there is a need to gain a better understanding and assessment of the distribution and potential hazards posed by the presence of PTEs in soils, in order to ensure food safety and environmental and public health protection.

Various analytical techniques have been developed for the detection of PTEs, including electrochemical (12, 13), voltammetry (14), and optical (15) methods. Laboratory optical methods include inductively coupled plasma atomic emission spectroscopy (ICP-AES) (also referred to as inductively coupled plasma optical emission spectroscopy (ICP-OES)), inductively coupled plasma mass spectroscopy (ICP-MS)), atomic fluorescence spectrometry (AFS), X-ray fluorescence spectrometry (XRFS), laser-induced breakdown spectroscopy
(LIBS), neutron activation analysis (NAA), and atomic absorption spectrophotometry (AAS). These laboratory methods are sufficient for quantitative estimation with good detection limits, and have been widely used to detect different PTEs in soil samples, with high sensitivity, selectivity and accuracy (14, 15). However, all these techniques require relatively expensive instrumentation, the measurement procedure takes as relatively and needed long time and specialized personnel are necessary to carry out the measurements correctly (14).

Given the limitations and cost of the laboratory methods listed above, there have been tremendous efforts made to provide accurate rapid measurement techniques for \textit{in-situ} measurement (16). Cutting-edge technologies such as portable and powerful computers, software packages, fibre optics and spatial modelling have contributed significantly to assessment of the presence, distribution and potential hazards posed by the presence of PTEs in soils. Contextually, proximal soil sensing offers high sampling resolution combined with cheaper measurements procedures (16, 17). Proximal soil sensing refers to the use of all field methods to obtain signals from the soil when the probe is in contact with or close to (within 2 m) the soil (18). It includes electromagnetic induction (EMI), gamma radiometry (GR), and ground-penetrating radar (GPR), electrochemical sensors, and field spectroscopic methods such as visible and near-infrared spectroscopy (Vis-NIRS), mid-infrared spectroscopy (MIRS) and portable X-ray fluorescence spectroscopy (PXRFS) spectroscopy and portable LIBS. The suitability of these techniques for laboratory and/or the field measurement has been extensively verified (16, 19).

Since the 1990s, there has been a tremendous increase in the use of portable Vis-NIRS spectrophotometers for on-site determination of various analytes (17). These portable instruments are non-destructive, easy to use and offer rapid measurement as there is no sample preparation needed. These characteristics are all essential for cost-effective \textit{in situ} measurement (16, 20). Likewise, MIRS has been successfully used for predicting soil
parameters including PTEs concentrations (21), despite the complex nature of soil matrix and the sensitivity of MIRS to moisture content that can impact the accuracy of measurement. PXRFS has advantages over the traditional methods, in that it offers *in situ*, rapid detection and accurate quantification of a wide range of elements present in the soil, with little, or no, need for sample preparation (19).

In recent years, there has been a remarkable increase in the development of rapid sensor-based *in situ* applications of proximal soil sensing technologies to detect and monitor PTEs contaminants in soils. Among them, PXRFS has been extensively adopted (16, 22). In the last few years, portable LIBS has been developed to measure PTEs *in situ* (23–25). Others used vibrational spectroscopy methods including Vis-NIRS and MIRS to monitor and measure PTEs in soil also has been used (3, 16, 26, 27). All studies conducted to date agree that field-based, *in situ*, spectroscopic techniques offer rapid, non-destructive and cost-effective methods to determine on-site presence, level and spatial distribution of PTEs prior to undertaking laboratory-based chemical estimation that might be more accurate but would be much more expensive and time-consuming. Furthermore, on-site spectroscopic techniques can provide real-time monitoring and, consequently, helpful information for preliminary site assessment, and guide further sampling for a comprehensive risk assessment of contaminated sites. Nevertheless, spectral data analysis requires careful examination to avoid misleading interpretations. Multi-sensor and data fusion approaches have attracted much attention for estimating soil properties (28–30); however, this field is still in its infancy as far as accurate PTE detection and mapping in soils is concerned.

The aim of this paper is to provide a critical review of the potential of those spectroscopy techniques currently available for the analysis of PTEs in soils, with a particular emphasis on portable XRFS, LIBS, Vis-NIRS and MIRS methods for on-site measurement. In addition, a
discussion on the main factors or variables affecting the implementation of these methods is provided along with suggestions for improving performance and accuracy for in situ measurement.

2 Reasons for using the term “PTE” instead of heavy metals

The term “heavy metal” has never been precisely defined and has become the subject of a broad discussion (31–34), implying different things to different people (35). A wide variety of definitions can be found in the Web of Science and Google Search, including elements that form soaps upon reaction with fatty acids, have an atomic weight greater than Na (22.9), e.g., Al, Co, and Pb elements with a specific gravity of approximately 5.0 or higher, e.g., Pb, Ag, Cr, Co, Cu, Ni, and Hg; and elements that can be precipitated by hydrogen sulphide in acid solution, e.g., Ag, Au, Pb, Hg, Bi, and Cu (33).

Furthermore the term “Heavy metal” has never been defined by any authoritative body, such as the International Union of Pure and Applied Chemistry (IUPAC), nor is there any widely agreed criterion-based definition of a heavy metal, indeed the term “heavy metal” is considered imprecise at best, meaningless at worst and its use is strongly discouraged (36–37). It has been suggested by (34) that term “heavy metals” be replaced with, e.g., “potentially toxic element(s)/metal(s)” or “trace element(s)/metal(s)”, according to the context. Because not all trace metals are toxic, the term which has gained acceptance is “potentially toxic elements” (PTEs).

Indeed, the term PTEs is more inclusive and appropriate than “heavy metal” (33) and will be used in this paper. While laboratory atomic spectroscopic methods are frequently the choice for the analyses of PTEs (38–40), the focus of this review is on portable optical spectroscopy methods: XRFS and LIBS (both atomic spectroscopy), and Vis-NIRS, and MIRS (both molecular spectroscopy). The discussion will be steered towards the potential implementation
of these portable spectroscopy techniques for *in situ* measurement, highlighting the advantages and disadvantages of each technique. A new approach for multi-spectroscopy sensor data fusion is suggested, as the optimal solution for *in situ* detection and mapping of PTEs.

3 Laboratory/field spectroscopy methods for analysis of PTEs in soil

The laboratory methods for the analyses of PTEs listed above, are both expensive and time consuming, so the number of samples analysed will invariably be limited, which prevents the collection of a large number of samples from across the entire site to assess and accurately map field contamination. Such methods do not usually fulfil the requirement for rapid measurement and risk assessment of soil contamination. It is necessary to be able to detect and measure elements posing a danger to health quickly, cost-effectively and in *situ*. There is a need for portable, economical and rapid sensing technologies, and this can be achieved by spectroscopy techniques. This section focuses on portable spectroscopy methods that can be used in both field conditions and the laboratory; PXRFS, LIBS, Vis-NIRS and MIRS.

3.1 X-ray fluorescence spectroscopy (XRFS)

Several studies and practical applications have demonstrated that XRFS is a suitable tool for industrial and environmental applications. XRFS is a well-known laboratory technique (41), which has been widely used since the 1950s for elemental analysis (19). The energy as well as intensity associated with characteristic-fluoresced radiation from the elements in a given sample are used to recognize and determine their presence and concentrations (42). When a single atom is excited by an external energy source, it emits photons of a characteristic energy or wavelength. The elements present in a given sample may be identified and quantified by counting the number of photons of each energy emitted (43).
Fluorescence is a form of luminescence that occurs when the emitted light has a longer wavelength (lower energy) than the absorbed radiation (19). Elemental identification in a given sample is based on the relationship between emission wavelength and atomic number, while the elemental concentration can be estimated from the characteristic line intensities (number of photons of each wavelength) (44, 45). This has proved to be a highly sensitive technique for PTEs analysis (46–48) with accurate multi-element analysis of rock samples (47), and soils matrices (49–53). With recent technological improvements, PXRFS has now become available, for rapid and cost-effective on-site measurement of soil contaminants (16, 54).

Using portable analytical methods, such as PXRFS, for assessing contamination has many advantages, including; increased efficiency, cost-effectiveness, convenience and ease of use, real-time information for on-site decision-making, minimal sample preparation required, multi-element measurement capability, potential for relatively low detection limits and better representation of contaminant distribution. (55–58). Such non-destructive testing also has no negative impact on the environment, PXRFS has been accepted by the environmental community as valuable technique for analysing environmental samples with several advantages over other optical analytical methods such as AAS, ICP-AES and ICP-MS (59). PXRFS has been used for rapid field screening of As contaminants with a strong correlation between the obtained results, and those from of AAS laboratory analysis (60). In addition, PXRFS has been successfully used for both in situ and ex situ (laboratory) analysis for the quantification of Pb, Zn, As, Cd, and Cu in paddy soils near a multi-contaminated mining area (61). The obtained results for both in situ and ex situ sampling were close to those obtained by Korean standard laboratory tests (coefficient of determination: \( R^2 \) close to 1.0) for Pb and Zn (61). Radu and Diamond (62) compared two PXRFS instruments with AAS analysis of aqua regia digested soil samples for estimating As, Zn, Pb, and Cu in old mining
areas of Ireland. The authors reported excellent to very strong correlations ($R^2$ of 0.99 for As, 0.99 for Pb, 0.96 for Cu, and 0.84 for Zn) between the two methods. They recommended PXRFS due to its speed because there was no need for sample digestion. Kilbride et al. (63) compared ICP-OES assessment of \textit{aqua regia} extraction method with two types of field PXRFS systems (X-ray tube and dual isotope) for determination of As, Cd, Cu, Fe, Mn, Pb and Zn in selected industrial areas of the UK. These Authors reported a high correlation between X-ray tube measurements and ICP-OES results for Fe and Pb better than the dual isotope, whereas higher correlations for Cd, Cu, Mn, and Zn were obtained with the dual isotope instrument measurements.

There are general limitations faced by users of PXRFS such as the need for official approval for field measurement, the need for a field expertise (19) and the health risks associated with X-ray radiation. The complex relationship between the intensity of the characteristic X-ray emission of an element and its concentration within a material is a limitation on the measurement which requires a correction to be applied to the measurements (64). However, internal calibration by the manufacture such as fundamental parameter (FP) overcomes this limitation and PXRFS produces robust data that accurately reflects material composition (45). Interference caused by physical-matrix effects (\textit{e.g.,} particle size, and surface conditions), chemical-matrix effects causing elemental interference (\textit{e.g.,} As and Pb), and moisture content ($>20\%$) should be considered and minimized (19). Variation in soil samples is a challenge for PXRFS measurement, inhomogeneity resulting from coarse textures, veins, phenocrysts, localized mineralization and irregularities of the surface reduce the accuracy of measurement (65, 66). Similarly, spectral interference between elements such As and Pb associated with a shared spectral peak reduces the estimation accuracy (67, 68). Moisture content reduces the PXRFS measurement and needs to be corrected to dry conditions. For example, Weindorf et al. (69) compared the accuracy of PXRF measurement for soils
samples under four different moisture conditions: dry, moist, frozen *in situ*, refrozen in lab. They reported significantly improved results using linear regression equations (higher $R^2$, lower RMSE, and slopes closer to 1) by applying a moisture correction factor of elemental data to dry.

In order to optimise the PXRF measurement and output, Binstock et al. (70) developed a new protocol for measurements. This consisted of drying and sieving collected samples to <50 μm particle sizes. Results for Pb showed no statistical difference between paired ICP-AES and PXRFs. Figueroa-Cisterna et al. (71) used a combination of *ex situ* PXRFs with different multivariate statistical techniques (*e.g.*, cluster analysis (HCA), principle component analysis (PCA), factor discriminant analysis (FA), and linear discriminant analysis (LDA)) to develop calibration models for *in situ* measurement of K, Ca, Cu, Fe, Mn, Pb, Ti, Zn, Rb, and Sr in Boris Angelo mining area of Central Chile. They concluded that the proposed *ex situ* calibration approach could be applied directly without pre-treatment of samples *in situ*. Rouillon and Taylor (2016) evaluated the analytical capabilities of PXRFs for the measurement of contaminated soil samples, achieving excellent correlation between the ICP-AES measurements and the PXRFs results for most elements (*e.g.* $R^2 = 0.999$ for Pb and Mn, $R^2 > 0.995$ for Zn, Cd, and Cu). Both ICP-AES and ICP-MS gave poor measurement accuracy for Ti and Cr, compared to PXRFs. They concluded that PXRFs is a suitable alternative tool to ICP-AES in the measurement of Cd, Cr, Cu, Fe, Mn, Pb, Sr, Ti, and Zn in metal-contaminated soils, and provides a high efficiency solution for high sampling density, cost-effectively for complete geochemical investigations.

XRFS when coupled with multivariate analysis or machine learning becomes a potentially powerful tool for rapid and accurate estimation of PTEs and other soil quality parameters. This is normally called recalibration of the instrument and has been used by several
researchers to improve the results of XRFS measurement (72–74). For instance, Moros et al. (73) evaluated the feasibility of using partial least squares regression (PLSR) for modelling of XRFS spectra of estuarine sediments to predict the concentrations of Al, As, Cd, Co, Cr, Cu, Fe, Mg, Mn, Ni, Pb, Sn, V, and Zn from 116 samples. They reported an average prediction error of $\pm 37\%$ for the 14 elements and residual predictive deviation (RPD) values ranged from 1.1 to 3.9. Towett et al. (72) developed simple linear models for estimating total element concentrations of Al, Cr, Cu, Fe, Ga, K, Mn, Ni, Ti, V, and Zn in soil samples collected from across sub-Saharan Africa using an S2 PICOFOX™ spectrometer (Bruker AXS Microanalysis GmbH, Germany). The results revealed that XRFS accurately determined the studied total element concentrations ($R^2 > 0.80$). Kaniu et al. (74), used XRFS (with isotope source) spectrometer coupled with PLSR method and artificial neural network (ANN) for non-destructive analysis of soil nutrients needed for soil quality indicator assessment (SQIs). They reported that PLSR was more accurate for analysing C, N, Na, P and Zn ($R^2 > 0.9$) with low standard error of prediction (SEP) of 0.05%, 0.01%, 0.01%, and 1.98 µg g$^{-1}$, respectively, while for Mg, Cu and Fe the ANN analysis was better suited with $R^2 > 0.9$ and SEP of 0.08%, 4.02 µg g$^{-1}$, and 0.88 µg g$^{-1}$, respectively. However, these methods have not received sufficient attention in PXRFS measurement. Machine learning can handle the overlapping spectral futures, nonlinearity in the spectral response and matrix effects, generally improving the model prediction accuracy of PTEs, a statement that needs be approved by further research. A summary of case studies that used XRFS for the estimation of PTEs in soil matrices from different regions is presented in Table 1. The majority of reported results are based on specific calibrations provided by the manufacturers of the PXRF devices, whereas only two studies reported recalibration results using spectra analysis with PLSR (75) and cubist regression method (30).
For PXRFS measurements in Table 1, we see that median $R^2$ values, ranging between 0.91 and 0.98 can be observed in Table 1 for Zn, As, Pb, Cu, Cd, Mn, and Fe, whereas less accurate estimations were found for Cr and Ni (median $R^2 = 0.87$ and 0.52, respectively). The measurement accuracy of Fe and Pb is among the highest (median $R^2 = 0.98$). Minimum values of $R^2$ for in situ measurement As and Zn were 0.70 and 0.81, respectively (56). However, the in situ results were rather poor (median $R^2$ of 13% and 20% for Zn and As, respectively), suggesting the former measurement mode can be only used for screening and not for quantitative estimations. Further studies should confirm whether or not the performance of PXRFS can be improved when fresh soil samples are used. Table 1 also shows that accuracy of PXRFS measurements of processed samples (air-drying and sieving) are better than those reported for in situ measurements using fresh soil samples without any sample pre-processing (56).

3.2 Laser-induced breakdown spectroscopy (LIBS)

LIBS is one of the most promising tools for elemental analysis (76, 77). LIBS is an atomic emission spectroscopy technique whereby a high energy laser (typically $10^8$-$10^{10}$ W/cm$^2$), usually pulsed, is focused onto a sample surface as a means of generating a plasma with a temperature high enough ($\sim$8000 K) for the ablated material to dissociate and a volume of excited atoms and/or ions expands rapidly into a state of so-called phase relaxation when the atomic and molecular emission lines characteristic of the elements present can be observed (78, 79). A charge-coupled detector (CCD) or intensified CCD is subsequently used to collect, spectrally resolve, and detect the light emitted. The detector is gated, deferring detection until the state of phase relaxation is reached - to optimise the signal-to-noise ratio of the acquired spectrums. The individual peaks that characterise the spectrums represent the elements that comprise the sample and consist of three principal parameters: spectral shape,
wavelength, and intensity. These parameters vary according to the emitting elements but will be modified by the surrounding environment.

LIBS is a reliable technique that can be employed in a range of challenging contexts to directly sample several different material types (76, 77). The fact that the laser plasma is formed by specifically directed laser light gives it a range of important properties; e.g. the ability to focus the pulse on a very specific region, thereby creating a specifically defined area for analysis. Such an ability is very useful for a number of different applications, including spatial mapping of elements (micro-analysis). In addition, it is possible not only to transmit the laser pulse via an open-path (line-of-sight), but to use fibre-optics to generate a plasma located on a remote target. As such, it is highly suitable for elemental analysis. Another significant advantage is that LIBS can be merged with other analysis techniques to improve detection competences. It is also possible to improve the excitation capabilities of laser plasma by using auxiliary power sources that remotely add energy into the plasma; for example, a CO$_2$ laser (23).

By generating the LIBS plasma on a sample that is located within a microwave cavity, it is also possible to introduce microwave energy (77). Having the option to improve the emissions that are produced by the elements that form the LIBS plasma represents a significant development because it has the potential to augment detection competences and, through doing so, expand its applicability. Recent signal enhancement schemes such as the double-pulse method (a second laser pulse is timed to “hit” the plasma released by the first pulse, increasing the heating effect and amplifying the signal) and orthogonal pulse method (a second laser is oriented to be orthogonal to the first, and by timing the initial and orthogonal laser pulses even greater signal amplification can be obtained), however, this achieved signal enhancement with double-pulse LIBS in either configuration, they are element and matrix dependent (77).
The LIBS plasma technique is not without its limitations. For example it is not always possible to control the various operational parameters to optimise the analytical capabilities of LIBS. The ablated mass and nature of the plasma that is generated vary according to the operational parameters that are employed including the distance between the lens and the sample, the power density of the laser pulse, and the pressure and composition of the environment in which the plasma is formed (77, 80). The analyte signals are directly influenced by changes in these parameters in addition to the relative intensities of the emission signals. This undermines the effectiveness of the quantitative analyses and ability to use element ratios to identify materials. It is possible to generate LIBS plasma in intense conditions; however, the nature of the conditions has a direct and significant impact on the character of the spectrum changes. In some cases, there may be a requirement for alternate analytical lines and it can be challenging to calibrate for quantitative analysis. The analysis of bulk materials, especially when they are inhomogeneous, can also be challenging due to the minute mass involved in a single LIBS measurement. Effective LIBS analyses that rely on ablation to generate the atomised material followed by the subsequent excitation of the atoms depends on both the sampling, the laser parameters and the physical and chemical properties of the studied sample (77, 81). Effects in LIBS play a significant role in quantitative analysis using LIBS include the grain size, moisture content and the chemical matrix effect the on the LIBS emission lines (77, 82). This effect is often more severe in LIBS than in ICP methods because of the high density of the materials contained within the small laser plasma (77). Therefore, methods to amend matrix effects in LIBS are very important for quantitative analysis.

LIBS has been employed successfully for the qualitative and quantitative analysis of various elements in samples of diverse origin and nature, including soils (25, 83–85). Good calibration results were obtained for the determination of PTE concentrations with detection
limits in the 10 mg g⁻¹ range. The analysis of PTEs in soils has been investigated using LIBS in combination with established laboratory-based analytical technique such as ICP-OES, ICP-MS, GF-AAS, and AAS (86–88). Generally, LIBS-based analysis was reliable and results reasonably correlated with the other traditional analytic methods, encouraging authors to conclude that LIBS is a rapid and useful tool for screening and estimation of PTE contamination (86, 87, 89, 90). For example, the accuracy of LIBS for determining total contents of PTEs (e.g. Cr, Cu, Fe, Mn, Ni, Pb and Zn) in 20 reference soil samples has been compared with the results obtained by ICP-AES and AAS methods (86). Results showed that mean values of the relative standard deviation (RSD %) of metal concentrations measured by LIBS were generally much higher, with the exception of Cr, than the corresponding values obtained by ICP, and are comparable with, or lower than the RSD % values obtained in other studies. Authors report that the results obtained indicate that PTE concentrations measured by LIBS oscillate much more around the mean value, and often overestimate compared to the corresponding concentrations measured by ICP (86).

Due to the large amount and complexity of the information provided by a LIBS spectrum, chemometric approaches are now commonly used for processing the data (91–93) which open a new avenue for semi-quantitative and quantitative analysis beyond the traditional univariate approach. Several studies have addressed the use of chemometrics for PTEs based on LIBS spectral data (24, 89, 94). For example, the estimation of Al, Ba, Ca, Fe Mg, Na, and Si in situ has been investigated using 59 measured spectra of weathered terrestrial samples using a portable LIBS instrument in the ultraviolet (UV) and infrared (IR) spectral ranges coupled with a PLSR method (89). The LIBS results were compared with energy depressive X-ray fluorescence spectrometer (ED-XRF) and Raman spectroscopy. The results of LIBS-PLSR models for the tested elements ranged between r values of 0.77 (for Al) and 0.92 (for Mg) in the UV spectral range, and r values of 0.48 (for Fe) and 0.74 (for Si) for the IR
spectral range (89). Authors concluded that the obtained results were in agreement with the range of composition measured by XRFS. LIBS data coupled with an ANN was used for determination of Cu in 59 spectra from a heterogenic set of reference soil samples and their respective Cu concentration (24). The ANN provided remarkable prediction accuracy for Cu, although portable instrumentation was employed. The authors reported a limit of detection (LOD) of 2.3 mg dm\(^{-3}\) of Cu and a mean squared error (MSE) of 0.5 mg dm\(^{-3}\) for the predictions (24). The performance of LIBS spectral data for determination of Cd under air and Argon (Ar) conditions were analysed and compared (94). PLSR and least-squares support vector machine (LS-SVM) models of Cd under both conditions were constructed. Results showed that the LS-SVM model for an Ar environment provided the best performance of prediction with \(R^2\) of 0.98, and root mean square error of prediction (RMSEP) of 0.034 mg kg\(^{-1}\). The authors concluded that LIBS combined with LS-SVM for an Ar environment condition could be a useful tool for the accurate prediction of Cd for environmental monitoring (94).

Table 1 summarizes some of the available studies on the application LIBS for predicting PTEs in the soils. Compared with XRFS, few papers are available reporting results obtained by LIBS. The use of multivariate statistical analyses is still limited to the commonly used PLSR, with a very limited application of machine learning methods (94). Spectral data preprocessing is not regularly applied, although standard normal variate (SNV) and wavelet methods are used to correct the spectral information, and reduce noise in order to improve the prediction performance (89). The highest correlation coefficients were found for the total concentrations of Zn (0.99), Pb (0.99), and Cu (0.98) (25). LIBS can also be used for mobile, and rapid semi-quantitative analysis and screening of PTEs in soils using the traditional calibration curve method (25).
3.3 Visible and near-infrared spectroscopy (Vis-NIRS)

Visible and near-infrared (Vis-NIR) spectroscopy has proved to be a useful tool for soil analysis (95). Once a soil medium is subjected to a light source, part of the light is absorbed, part is scattered and part diffusely reflected from the soil. The final shape of the soil spectra is a reflection of both the light scattering and absorption phenomena that differs due to the sample physical and chemical characteristics, respectively. The working principle of the absorption Vis-NIR spectroscopy stems from activating chemical bonds by irradiating mineral mixtures thereby creating resonance vibration, either by bending or stretching of the molecular bonds. These types of vibrations result in light absorption, in different levels, having a particular power quantum related to the difference between two energy levels. Since the energy quantum is actually straightforwardly associated with frequency, the resulting absorption features of the spectrum curve can be used for analytical purposes (95). In the visible (Vis) range (400–780 nm), absorption bands related to soil colour are due to electron excitations, which assists the measurement of soil organic matter content (SOM) and moisture content (MC). Within the NIR range (780 – 2500 nm), overtones of OH and overtones and combinations of C-H + C-H, C-H + C-C, OH+ minerals, and N-H of fundamental vibrations (e.g., C–H, N–H, O–H, C–O, Si–O) that occur in the MIR spectral range (2500–25000 nm or wave number of 4000–400 cm⁻¹) (96, 97). These are the major spectral features essential for the detection and quantification of key soil properties having direct spectral responses e.g., MC, SOM, clay minerals, and total nitrogen (98). These properties can be accurately predicted with Vis-NIRS with appreciable accuracy, compared to properties with indirect spectral responses (e.g., P, K, Mg, Ca, Na, CEC, pH) that are potentially predictable through covariation with properties having direct spectral responses in the Vis-NIR spectral range (95). The NIR overtones and combinations are of broad bands that overlap but 10-100 times weaker compared with the MIR spectrum where fundamentals are more
resolved, permitting the structure of a sample to be better elucidated (95, 99, 100). The higher energy of NIR radiation and the relatively low absorptivity of water makes NIR spectroscopy as a better technique when wet soils and sediments samples are analysed, compared with MIR spectroscopy (99, 100). Due to the overlaps among overtones and combinations due to their broad bands, advanced chemometrics and machine learning tools are used to extract quantitative and qualitative information. Overall, Vis-NIRS is widely adopted to determine SOM, minerals, texture, nutrients, and PTEs in soils (95, 100).

The potential of NIR reflectance spectroscopy coupled with PLSR was investigated by Kooistra et al. (101) for the assessment of soil contaminated by Cd and Zn in river floodplains in the Netherlands (101). The authors reported a coefficient of determination ($R^2$) of 0.88 and 0.90 and root mean square error in cross validation (RMSECV) of 0.68 and 80.97 mg kg$^{-1}$, for Cd and Zn, respectively. Also, the prediction of PET (As, Cd, Cu, Fe, Hg, Pb, S, Sb, and Zn) concentrations in soil samples was performed based on Vis-NIRS coupled with an ANN and stepwise multiple linear regression analysis (SMLR) (7). The authors reported high prediction accuracy results for six out of the nine elements, with $R^2$ values of 0.84, 0.72, 0.96, 0.95, 0.87 and 0.93 for As, Fe, Hg, Pb, S, and Sb, respectively, for SMLR. No successful correlations were obtained for Cd (0.51), Cu (0.43), and Zn (0.24). Siebielec et al. (102) based on full Vis-NIR spectra (400-2500 nm) of soil samples collected from a wide range of pH, organic carbon and soil textures, reported good prediction using PLSR models with $R^2$ of 0.97, 0.94, 0.80, 0.99 and 0.96 for Fe, Cd, Cu, Ni and Zn, respectively. Wu et al. (103) used a Vis-NIR spectrometer coupled with univariate regression and principal component regression (PCR) analyses for predicting Hg concentration in agricultural soils from the Nanjing region in China. The authors used some pre-treatments such as Kubelka–Munk transformations, absorbance and derivative to optimize the prediction of Hg concentration, achieving the best results by using the PCR regression with the Kubelka–
Munk transformation with $R^2$ of 0.69, and RMSEP of 0.15 mg kg$^{-1}$. They found a positive correlation between Hg concentration and the absorption of goethite and clay minerals, and they suggested that Hg sorption by clay-size mineral aggregates in soils was the mechanism explaining the successful prediction of Hg having spectral featureless (103). The capability of Vis-NIRS (400–2,500 nm) coupled with PLSR was evaluated for the measurement of AS in forest soil (104). PLSR was also used to estimate AS concentration in soil samples (105), reporting $R^2$ of 0.69–0.71, RMSEP of 1.64–1.61 and RPD of 1.70–1.80 for the best calibration model. Authors pointed out the impact of soil organic matter (OM) content on the model prediction accuracy. Estimating and mapping the distribution of PTEs in soils using Vis-NIRS combined with SMLR and enter-MLR (EMLR) was conducted by Choe et al. (106). The best results for predicting AS and Cu were achieved with EMLR models with $R^2$ values of 0.60 and 0.81, respectively. The authors provided a distribution map (contours) of As and Cu concentrations resulting from the EMLR models, which exhibited identical spatial patterns to a map produced based on the reference measured values of As and Cu (106).

In order to better predict low concentrations of As, Cu, Pb, and Zn metals in agricultural soils, a comparison was made between PLSR and genetic Algorithm-PLSR (GA-PLSR) has been made (107). Results showed that the GA-PLSR model outperformed the PLSR models with $R^2$ of cross-validation ($R^2_{cv}$) of 0.60–0.69 for the four metals analysed while the $R^2_{cv}$ of PLSR ranged between 0.49 and 0.58 (107). The authors attributed the Vis-NIRS measurement of studied PTE (having no direct spectral responses) to be due to co-variation with OM, as both heavy metals and OM have the same spectral features (e.g., same significant wavelengths) (107). The feasibility of using Vis-NIRS to predict PTEs (Cr, Zn, Cu, Ni, and Pb) in several soil types using PLSR was reported for a collection of samples from Bulgaria (27). The results revealed that the best prediction obtained was with the Cu model. However, the prediction of Zn, Pb and Ni models were less accurate, and the results
for Cr in both cross-validation and prediction were the worst (27). The authors found that the values of both RMSECV and RMSEP increased when the number of samples in the calibration set (cross-validation) was decreased. They concluded that Vis-NIRS is a useful tool to predict PTEs in soil samples, on the condition that a large number of samples was used in the calibration. A case study to monitor the suitability of Vis-NIRS for predicting concentration of PTEs based on PLSR and support vector machine regression (SVMR), using samples from mining dumpsites in the Czech Republic was reported in (108). The best prediction of As was obtained with the SVMR model with the first derivative pre-treatment ($R^2 = 0.89$, RMSEP = 1.89 mg kg$^{-1}$, RPD = 2.63), where acceptable prediction was obtained for Cd and Cu ($R^2 = 0.66$ and 0.81, RMSEP = 0.0.8 and 4.08 mg kg$^{-1}$, RPD = 2.0 and 2.5, respectively).

Vis-NIRS has been reported as a rapid, non-destructive and cost-effective tool for estimating the concentration of PTEs in soil (3, 16). Nevertheless, the actual correlations obtained between Vis-NIR spectra and the PTEs are not necessary based on the direct response of these element in the Vis-NIR spectrum (3, 16). The majority of the reported studies have proposed that the calibrations obtained were based on indirect correlations, between the measured metals and other soil properties, having direct spectral features in the NIR spectral range such as clay, organic carbon, and iron oxides (3, 16, 109). Therefore, caution is necessary before applying the contaminate-based model to a different sample matrix.

Wu et al. (109) explored the impact associated with surrogate calibrations. The authors compared soil samples naturally rich in Cd and Zn with the spiked soil samples with Cu and Zn, concluding that the Vis-NIR spectrum is sensitive to sample handling as well as the orientation of the sample relative to the Vis-NIR instrument. They could not find distinguishable effect in respect of the presence of PTEs on the Vis-NIR spectrum (109), which supports the hypothesis of “surrogate” correlation for PTEs, and suggests that even in
highly contaminated soils PTEs may not be identifiable over the OM absorption bands in the Vis-NIRS region (109). However, there are several cases of successful application of NIR spectroscopy to measure PTEs in soils, which necessitates further research to explain the physico-chemical relations between the PTEs and other soil properties (e.g., SOM, and clay minerals), having a direct spectral response in the NIR spectral range.

Previous research has shown that sample preparation (drying, grinding, sieving) before scanning with NIR spectroscopy affected the measurement accuracy of soil fertility parameters (95). Likewise for PTEs, according to published literature sample preparation has a clear effect on measurements. The effect of sample preparation such as homogenization, drying, grinding, and sieving on estimating the accuracy of cyanide concentration using a hand held NIR instrument and PLSR was investigated by Sut et al. (110). The authors found that the root mean square error of measured cyanide concentrations was larger for unprepared fresh soil samples (RMSE > 2400 mg kg\(^{-1}\)), compared to prepared samples in the laboratory (RMSE > 1750 mg kg\(^{-1}\)). They concluded that the direct in situ measurements yielded high a LOD, which, in laboratory, can be improved by sample preparation, with the sample preparation actually resulting in reducing the influences of light scattering, soil moisture and particle size on accuracy of measurement. They also found that the prediction of cyanide concentration in processed soil samples (dried and sieved) (R\(^2\) = 0.41) was improved after grinding (R\(^2\) = 0.86).

More case studies concerning the use of NIR spectroscopy for the estimation of PTEs in soil matrices are provided in Table 1 for different countries over the world. The full Vis-NIR spectral range of 350-2500 nm was used in the majority of the reported studies. Spectral data pre-processing was used intensively to improve the prediction results, among which the first derivative (FD) and continuum removal (CR) methods were the most used. The PLSR linear multivariate analysis was the most common method used for modelling. In general,
comparable prediction accuracies were found for the chalcophile elements (As, Cu, Pb and Zn), compared to the siderophile elements (Co, Fe, Mn and Ni). The highest prediction performance was for the total concentrations of Pb ($R^2 = 0.94$) and Ni ($R^2 = 0.93$) (Table 1). The successful prediction of these elements can be attributed to the relationship between these elements and carbonate (111). Fe minerals are the main parameter determining the prediction accuracy (7, 103, 112), in addition to clay minerals and SOM (102, 113), all having direct spectral responses in the NIR range (95).

3.4 Mid-infrared spectroscopy (MIRS)

MIRS is a rapid and cost-effective techniques developed over the last 40 years for soil analysis (114). Fundamental vibrations of chemical bonds e.g., C–H, N–H, O–H, C–O, Si–O take place in the MIR electromagnetic range (2500–25000 nm or wave number of 4000–400 cm$^{-1}$) (97), and can provide extensive and wealth information about the chemical and physical properties of the sample components (115). The MIRS spectra can reveal essential information related to both organic and some inorganic compounds. However, the complexity of soil matrices, and the relatively low concentrations of PTEs usually do not show absorption features in MIR spectrum (16), but they are bound to spectrally active soil components (e.g., OM and clay minerals) and as a result modify the resultant spectral features (21). However, compared to the Vis-NIRS, the MIRS offers more informative spectrum with well delineated absorption bands (97, 116). By using MIRS, it has been demonstrated that particular markers exist in soil MIR spectrum for minerals and organometallic complexes (117).

Many reports on the use of MIRS for the measurement of key soil physical and chemical properties can be found in the literature. For example, MIRS has been successfully used to estimate soil total carbon, organic, and inorganic carbon, providing more accurate estimations than Vis-NIRS (118, 119). Recent research has revealed that the MIR can be used to measure
PTEs in soil (97, 102). In their study, Siebielec et al. (102) used 70 soil samples collected from the mining area of Tarnowskie Gory (Upper Silesia, Poland) for predicting Fe, Cd, Cu, Ni, Zn, and Pb concentrations based on the combined spectral range of NIRS and MIR spectra (2500 to 25000 nm) coupled with the leave-one-out PLSR cross-validation procedure. They achieved accurate calibrations between Fe, Cd, Cu, Ni, and Zn and MIR spectra with $R^2$ of 0.97, 0.94, 0.80, 0.99, and 0.96 for those metals, respectively, but Pb had a low correlation with $R^2 = 0.66$, and the calibrations using NIRS were less accurate than those based on MIR.

By testing 4130 European agricultural soils, Soriano-Disla et al. (97) found that MIRS was a powerful tool for predicting the concentrations of PTEs in soil. Given that the bioavailability of PTEs in soil is basically ruled through the interactions between the trace elements species and existing anionic functional groups, the MIRS method can be used successfully for estimation PTEs bioavailability. The feasibility of using MIR to determine Cr using tannery sludge in tropical soils has been tested (120). The authors reported changes in soil spectral signal due to metal adsorption to soil components, demonstrating differences between relatively contaminated-free and highly contaminated soils. The authors pointed out that MIRS outperformed Vis-NIRS for measuring Cr, suggesting that the OM content played a more substantial role of the determination of spectral characteristics than mineralogical compositions, especially those that have high OC content. Very recently, the feasibility of Vis-NIRS and MIRS handheld (In particular the 4300 Handheld Fourier Transform Infrared (FTIR) Spectrophotometer) for quantification of PTEs (Al, As, Ca, Cu, Fe, K, Mn, Na, Ni, Pb, and Zn) in 203 soil samples collected from Saxony, Germany, were investigated (21). The authors used PLSR methods with different spectral data pre-processing, namely, first and second derivatives, multi scatter correction (MSC), continuum removal (CR) and standard normal variate (SNV). The authors reported a high performance for the model of MIR-spectra with multi-scatter correction (MSC), and SNV-pre-processing with very good to good
prediction for Al, Fe, K and Ni models \( (R^2 = 0.70-0.88) \). It was concluded that MIR spectroscopy is a reliable and promising alternative approach for the estimation of metal contents in soil. Table 1 summarizes the most important studies on applying the MIRS for predicting PTEs in soil. Compared with Vis-NIRS, few articles are available reporting predictions of PTE concentrations in soils by MIRS. A wide range of pre-processing methods (e.g., CR, MSC, and SNV) were used to correct the spectral information, reduce noise and scattering effects, to improve the prediction performance. In general, PLSR was the common calibration method. Better predictions were found for the total concentrations of siderophile elements, compared to the chalcophile elements (see Table 1). The highest correlation coefficients were found for the total concentrations of Ni \( (R^2 = 0.98) \), Zn \( (0.96) \), and As \( (0.92) \) \(^{(21)}\). The successful prediction of these elements can be attributed to the fact that the spectral features in the MIR-region are mainly caused by fundamental molecular vibrations \(^{(21)}\).

(Table 1)

4. Data and model fusion methods

Soil studies are usually considered as a multifactorial problem and it is worth exploring and measuring the same samples simultaneously using several analytical tools (platforms) processing with on or a set of models. Although research on individual proximal sensors has demonstrated potential, no individual sensor (or technique) is able to completely characterize soil complexity. Soils are typically characterized by high variability on all scales that can be monitored using different sensor systems. Each sensor offers an exclusive perspective on one aspect of a soil system that will change both temporally and spatially. Consequently, requiring a single sensor to perform more than one function in these circumstances is almost unfeasible \(^{(136)}\). Data fusion is the combination of data from different sources to produce more rigorous data than could be obtained from a single system or sensor \(^{(16)}\). Sensor fusion
can be a sensible option to integrate diverse variations across scales and different soil properties. Multi-model fusion or ensemble methods are a useful tool for improving predictions, and the data fusion approach has been applied extensively in agriculture in recent years (137–139). Such an approach has not been yet tested for PTE analysis but sensor data and multi-model fusion methods have the potential to improve performance when estimating PTEs in soil using various complementary sensors and a set of rigorous models.

The optimal goal of the fusion technique is to gain high quality and reliable information, providing better and greater in-depth understanding (140). Fusion techniques are expected to deduce inferences that are potentially more accurate than if they were achieved by a single sensor or model. Fusion of data from multiple sensors and/or sets of models provides several advantages over data from a single sensor such as robustness, accuracy, extended feature coverage and harmonizing information on particular phenomena (30, 140, 141). The fusion technique can be categorized into two major types: (1) proximal sensor data fusion in which data from different sensors are applied jointly into a single multivariate or machine learning method; (2) multi-model fusion, in which models are combined to provide the optimal prediction. (see Fig.1).

Measuring with multiple sensors such as portable/handheld Vis-NIRS, MIRS, and PXRFS is versatile and provides a larger amount of data. This progress in data acquisition offers the possibility of a better understanding of soil contamination and improved analysis. Vis-NIRS, MIRS, and PXRFS have been used independently for determination of PTEs in soil sample data, but model fusion is still new in environmental studies particularly, for PTE determination, and has attracted no or little attention in the literature. Wang et al. (29) successfully combined Vis-NIRS and PXRFS for the quantification of soil nitrogen and carbon. Likewise, Chakraborty et al. (142) combined Vis-NIRS spectra and PXRFS to
develop an optimised model for the rapid prediction of petroleum hydrocarbons in soils. Towett et al. (143) combined MIRS and PXRFS to predict soil properties and distributions of available Al, Cu, Fe, Zn and Mn from sub-Saharan Africa soils using RF regression. The authors reported a very good prediction for Al ($R^2 = 0.86$), poor prediction for Mn ($R^2 = 0.56$), and very poor predictions for Cu, Fe, and Zn ($R^2 = 0.43, 0.40, \text{ and } 0.04$, respectively). The authors found that the prediction of some elements was strongly improved by adding XRFS data to the MIR data.

Multi-model fusion can improve the overall model prediction accuracy. Combining different model outcomes in this way is termed as model ensemble or averaging (144). This combination of models reduces the variances of predictions found with individual models, hence, the combined predictions typically outperform those from individual models. The majority rule (simple averaging) is the common model fusion technique, which integrates all models by using an equally weighted averaging technique that integrates several ensemble methods (141). For instance, Chakraborty et al. (142) applied a model fusion technique by using Vis-NIR based penalized spline regression (PSR) method followed by PXRFS elemental based RF regression, which resulted in improved prediction accuracy of total petroleum hydrocarbon (TPH) estimation in contaminated soil. The PSR was used to fit the training set (containing Vis-NIR spectra only) and RF was used to fit the residual of the PXRF elemental data. The authors found that the combined modelling approach produced better results compared to individual model outputs. In another study, O’Rourke et al. (30) investigated the combined approach of Vis-NIRS, MIRS, and PXRF data to improve the accuracy in estimating PTEs in soil. These authors again found integration between Vis-NIRS, MIRS, and PXRFS resulted in better determination of (As, Cd, Co, Cu, Hg, Mn, Ni, and Zn) distributions in soil than could be obtained using any one spectral method. The authors further argued that the multi-sensor data fusion approach can be an alternative to
traditional wet chemistry analyses, and is potentially suited for large scale routine soil monitoring.

(Figure 1)

5. Selecting of optimal optical technique for PTE analysis

In order to select the most suitable optical technique for the determination of PTEs in soil, information such as analysis time, cost per sample, portability and ease of field application, robustness, repeatability, prediction accuracy and the final application of results should be available. One example is, when time is not a significant issue and accuracy is of prime importance, here laboratory techniques are the ideal option, as their accuracy is higher than portable field devices. However, in this case pre-processing of soil samples could also enhance the prediction accuracy and enable more reliable results to be obtained.

With recent advances in proximal soil sensing and data mining techniques, portable field equipment is now available that can carry out measurements in situ for rapid detection and screening with good to acceptable accuracy. However, it is important to note that there are many challenges associated with field spectroscopy, among others, surface roughness, texture, moisture content, plant residues and gravels are the parameters with the most negative effects. Compared to laboratory spectroscopy, field spectroscopy requires extra attention to eliminate the effects of external (environmental) variables. Recent advances in pre-processing algorithms to eliminate these external factors, such as external parameter orthogonalization (EPO) and direct standardization (DS) are recommended by many studies (145, 146). Moreover, data mining techniques such as machine learning methods when combined with EPO and/or DS can deal with both linear and non-linear sources in the data set and improve prediction accuracy (95, 147, 148). It can be hypothesised that the
combination of multi-sensors using data fusion approach together with EPO and DS is the ultimate solution for optimal prediction accuracy for in situ measurement of PTEs. Future work is needed to confirm this assumption, although developments in agriculture has already demonstrated its general validity.

For decision-making, important factors such as analysis run time, cost per sample, operational skills required and portability need to be considered. The portable vis-NIRS, MIRS, LIBS, and PXRFS are cost-effective, time-saving, and provide sufficient accuracy in screening and monitoring PTE contamination levels in soil when compared to laboratory analytical methods (16, 21). Portable field measuring techniques enable the collection of a high number of samples that can cover the entire contaminated area in a relatively short period, which is crucial for effective hazard assessment and remediation action planning (149–151). High-resolution sampling offers a better understanding of the spatial distribution of the area being studied by using geostatistical spatial interpolation and the Geographic Information System as a mapping tool. This offers accurate elemental maps and derivation of interpretive hazard maps for rapid and accurate remediation actions (151). In this context, Suh et al. (150) developed an inexpensive, rapid and accurate mapping system for soil subjected to Cu contamination using in situ PXRFS data corrected by reference Cu values obtained from ICP-AES analysis at a mine site in South Korea. The authors reported accurate prediction of Cu concentrations similar to those of ICP-AES with an excellent $R^2$ values of 0.99. They concluded that the method is effective for mapping soil contamination if the PXRFS databases and laboratory measurement (e.g., ICP-AES) are integrated (150). One example of potential use of portable sensors is fusion between PXRFS and portable vis-NIRS data for accurate and rapid prediction of PTEs (75), which can be integrated with satellite and unmanned aerial vehicle-based hyperspectral imagining systems for large scale investigations (152, 153).
Table 2 shows the main factors to be considered in the decision-making process when selecting analytical techniques for PTEs analysis. These factors can be linked to cost (device and sample), detection (targeted analytes and LOD), the measurement scale, and the operator’s skills. For example, the LIBS provides a wide range of analysis (more than 20 elements) at acceptable LOD (1-10 mg g\(^{-1}\)). However, the initial cost for the device ($35,000-50,000), the cost per sample ($15-30), and the fact this a non-established handheld/portable device for \textit{in situ} measurements, limit the use of such a method for field measurement.

On the other hand, portable spectroscopy systems such as vis-NIRS, MIRS, and PXRF are of much lower instrument cost ($30,000-40,000) with per sample cost ($10-20) (Table 2). The targeted analytes of these portable techniques cover the majority of PTEs (26 elements), with PXRF able to provide real-time measurement in the field with good accuracy (\(R^2 = 0.80\)) (56). Regarding operational skills, vis-NIRS, MIRS and PXRF spectroscopy all require low to medium skills, whereas laboratory methods such as ICP-MS, OES, and AES require medium to high skills, and NAA requires a high level of skill. This highlights an advantage for portable techniques in that they do not need highly expert personnel for \textit{in situ} measurement and to provide hazard maps in less than one day (150).

\textbf{(Table 2)}

\section*{6. Conclusions}

There are several well-established standard laboratory techniques for determining the concentration and distribution of potentially toxic elements (PTEs) in soils. The methods offer a diverse sensitivity range, are highly accurate down to trace level detection. However, there are drawbacks such as sample extraction and analytical time required, cost, and relatively high level of technical expertise needed. Thus there is a growing
demand for in situ and real-time detection/measurement tool. This has contributed to the development of new portable/handheld devices such as visible and near infrared (Vis-NIR), mid infrared (MIR), laser-induced breakdown spectroscopy (LIBS), and X-ray fluorescence (XRF) spectrometry, which offer rapid measurement and allow for the high density sampling necessary for quantification and mapping of PTEs to support decisions for e.g., land reclamation. However, there are still limitations related to soil matrix properties such as variation in moisture content and soil texture that can reduce the prediction accuracy. To overcome these issues, advanced pre-processing algorithms such the external parameter orthogonalization (EPO) and direct standardization (DS) coupled with machine learning techniques such as support vector machine (SVM) or the random forest (RF) can be used. An artificial neural network (ANN) that can cope with nonlinearities in the spectral responses of the soil can eliminate the influence of these variables and improve prediction accuracy. Moreover, data fusion technique that merge data from different sensors, or predictions of different models, can improve estimation accuracy. However, much research is still needed in the area of portability and real-time prediction based on spectroscopic techniques that can speed up the field measurement, and maybe revolutionise the way we conduct risk assessment or site investigation.

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Table 1 Summary of case studies for estimation of potentially toxic elements (PTEs) concentrations in soils using visible and near-infrared spectroscopy (Vis-NIRS), mid-infrared (MIRS), and X-ray fluorescence spectroscopy (XRFS), and laser-induced breakdown spectroscopy (LIBS) in soil matrices from different regions over the world.

| Elements | Country | Samples No | Spectral range (nm) | Pre-processing | Range (mg kg$^{-1}$) | Regression technique | $R^2$ | RMSE | RPD | Reference |
|----------|---------|------------|---------------------|----------------|---------------------|---------------------|------|------|-----|-----------|
| Cd       | Netherlands | 69       | 400-2500           | FD+ NM         | –                  | PLSR                | 0.88 | 0.72 | -   | (113)     |
| Zn       |          |           |                     |                |                     |                     |      |      |     |           |
| As       | Spain    | 214       | 350-2500           | AB+ FD         | 7–442              | ANN, MLR            | 0.85, 0.83 | SEP=0.007-0.007 | 3.28, 3.83 | (7)       |
| Cd       |          |           |                     |                | 0.05–14.8          |                     | 0.49, 0.51 | 0.085-0.07 | 0.96, 1.23 |
| Cu       |          |           |                     |                | 17.5–521           |                     | 0.44, 0.54 | 0.09, 0.085 | 0.84, 1.23 |
| Hg       |          |           |                     |                | 0.01–13.9          |                     | 0.92, 0.95 | 0.033, 0.027 | 4.30, 5.77 |
| Sb       |          |           |                     |                | 196–3362           |                     | 0.92, 0.92 | 0.028, 0.033 | 4.55, 4.66 |
| Pb       |          |           |                     |                | 17.5–331.5         |                     | 0.94, 0.94 | 0.028, 0.031 | 5.89, 5.30 |
| Zn       |          |           |                     |                | 94.0–3887.0        |                     | 0.22, 0.23 | 0.08, 0.08 | 0.51, 0.59 |
| Fe       | Poland   | 70        | 400-2498           | MSC            | 0-160.0            | PLSR                | 0.87 | RMSD=9.2 |   | (102)     |
| Cd       |          |           |                     |                | 0-38.0             |                     | 0.54 | 5.13 | -   |           |
| Cu       |          |           |                     |                | 3.0-78.0           |                     | 0.61 | 10.3  | -   |           |
| Pb       |          |           |                     |                | 0.0-6800           |                     | 0.45 | 839.0 | -   |           |
| Ni       |          |           |                     |                | 3.0-77.0           |                     | 0.84 | 6.21  | -   |           |
| Zn       |          |           |                     |                | 0-4500             |                     | 0.67 | 526.0 | -   |           |
| Hg       | China    | 105       | 380-2500           | AB, K-M        | 0.04–1.26          | PCR                 | 0.71 | 0.15 | -   | (121)     |
| As       | China    | 120       | 900-2500           | MSC            | 5.57–47.07         | PLSR                | 0.65 | 1.65 | -   | (103)     |
| Cr       |          |           |                     |                |                    |                     | 0.76 | 5.60  | -   |           |
| Element | Location | Sample Size | Concentration Range | Method | Coefficient | Slope | Intercept | R² | p-value |
|---------|----------|-------------|---------------------|--------|--------------|-------|-----------|----|---------|
| Cu      | China    | 61          | 9.60–73.40          | PLSR   | 0.62         | 7.61  | -         |    |         |
| Hg      |          |             | 0.04–1.26           |        | 0.59         | 0.117 | -         |    |         |
| Ni      |          |             | 10.60–57.50         |        | 0.86         | 2.21  | -         |    |         |
| Pb      |          |             | 22.30–208.00        |        | 0.66         | 4.97  | -         |    |         |
| Zn      |          |             | 43.90–625.00        |        | 0.62         | 13.6  | -         |    |         |
| As      | China    | 61          | 1.13–13.30          | SMLR, EMLR | 0.72 | 0.86 | 1.90 | (112)  |
| Cd      | China    | 61          | 0.22–0.54           |        | 0.20         | 0.05  | 1.23      |    |         |
| Co      | China    | 61          | 11.5–19.7           |        | 0.80         | 0.93  | 218.0     |    |         |
| Cr      | China    | 61          | 60.8–104.0          |        | 0.85         | 4.20  | 2.50      |    |         |
| Cu      | China    | 61          | 26.5–55.3           |        | 0.67         | 3.74  | 1.74      |    |         |
| Ni      | China    | 61          | 27.3–50.6           |        | 0.81         | 2.32  | 2.35      |    |         |
| Pb      | China    | 61          | 21.4–42.8           |        | 0.55         | 3.58  | 1.49      |    |         |
| Zn      | China    | 61          | 68.8–120.0          |        | 0.56         | 8.20  | 1.45      |    |         |
| As      | Spain    | 49          | 292.2–826.9         | SMLR, EMLR | 0.84 | 0.88 | 2.36, 2.56 | (122) |
| Pb      | Spain    | 49          | 1412.7–18,811.3     |        | 0.53, 0.61   | 0.38, 0.37 | 1.36, 1.39 |
| Zn      | Spain    | 49          | 200.9–728.1         |        | 0.54, 0.59   | 0.22, 0.23 | 1.39, 1.38 |
| As      | South Korea | 22   | 52.4–1493.8         | SMLR, EMLR | 0.37 | 0.59 | SE=. 0.27, 0.25 | -,- | (106) |
| Cu      | South Korea | 22   | 21.9–252.6S         |        | 0.24, 0.81   | 0.21, 0.12 | -,- |
| Pb      | South Korea | 22   | 56.8–152.5          |        | -,-          | -,-   | -,- |
| As      | China     | 33          | 19.33–403.77        | SMLR, EMLR | 0.61 | 0.24 | -,- | (123) |
| Cu      | China     | 33          | 31.83–190.51        |        | 0.39         | 0.20  | -,-       |    |         |
| Fe      | China     | 33          | 28.14–64.20         |        | 0.43         | 0.12  | -,-       |    |         |
| Cu      | Germany   | 109         | 0.22–0.54           |        | 0.20         | 0.18  | -,-       |    |         |
| Pb      | Germany   | 109         | 11.5–19.7           |        | 0.80         | 0.93  | 218.0     |    |         |
| Element | Value | Range |
|---------|-------|-------|
| Zn      | 40-1322 | 0.81  | 117.5 | -   |
| Fe      | -     | 0.84  | 4960  | -   |
| Mn      | -     | 0.70  | 363.3 | -   |
| Pb      | USA   | 24    | 350-2500 | 8-1823 | 0.74  | -   |
| Zn      | 0-605 | 0.76  | -     | -   |
| Cu      | 40-1322 | 0.81  | -     | -   |
| Cd      | -     | 0.42  | -     | -   |
| Mn      | -     | 0.80  | -     | -   |
| Cu      | 30    | 350-2500 | AB+ FD | 1.670–332.18 | UR | 0.79 | 17.85 | -   |
| Pb      | 2.480–50.920 | 0.65 | 0.91 | -   |
| Zn      | 73.76–269.53 | 0.61 | 22.51 | -   |
| Zn      | 43.90–625.00 | 0.62 | 13.6 | -   |
| Ni      | 120   | 380-2500 | PLSR, MARS | 0.75, 0.93 | 2.94, 229 | 1.99, 2.56 |
| Cr      | –     | 0.71, 0.83 | 5.54, 4.29 | 1.49, 2.54 |
| Cu      | –     | 0.55, 0.77 | 6.49, 4.62 | 1.28, 1.46 |
| As      | –     | 0.40, 0.53 | 1.14, 1.00 | 1.01 |
| Zn      | –     | 0.48, 0.68 | 14.78, 11.41 | 1.37, 1.78 |
| Pb      | –     | 0.38, 0.60 | 5.92, 4.67 | 1.25, 1.59 |
| Cd      | –     | 0.69, 0.80 | 0.04, 0.04 | 2.45, 2.45 |
| As      | 122   | 350-2500 | FD-UVN | 3.800–16.600 | PLSR | 0.45 | 1.60 | 1.13 | (127) |
| Cd      | Non   | 0.081–1.441 | 0.30 | 0.16 | 0.97 |
| Cr      | Non   | 30.990–113.90 | 0.98 | 19.46 | 1.01 |
| Cu      | Non   | 9.100–55.500 | 0.92 | 5.45 | 1.54 |
| Hg      | FD-UVN | 0.030–0.330 | 0.83 | 0.048 | 1.23 |
| Pb      | UVN   | 11.120–89.680 | 0.68 | 5.94 | 1.11 |
| Element | Location | Observed Concentration Range | MSC | Mean | PLSR | Standard Error | n | Reference |
|---------|----------|-----------------------------|-----|------|------|----------------|----|-----------|
| Fe      | Poland   | 2500-25000                  | 0.1–160 | PLSR | 0.97 | 4.10           | (102) |
| Cd      |          | 0.1–35.0                    | PLSR | 0.94 | 1.82 |
| Cu      |          | 0.1–78.0                    | PLSR | 0.80 | 7.52 |
| Pb      |          | 0.1–6800                    | PLSR | 0.66 | 662  |
| Ni      |          | 0.5–75.0                    | PLSR | 0.98 | 1.88 |
| Zn      |          | 0.5–4500                    | PLSR | 0.96 | 191  |
| Cu      | Australia| 2500-25000                  | 0.01–7.2 | Cubist | 0.14 | 0.65 | 1 | (128) |
| Fe      |          | 2.6–500                     | PLSR | 0.49 | 35.30 | 1.3 |
| Mn      |          | 0.67–152                    | PLSR | 0.15 | 18.66 | 1 |
| Zn      |          | 0.05–11                     | PLSR | 0.15 | 0.90  | 1 |
| As      | China    | 8000-25000                  | 3.73–16.51 | PLSR | 0.92 | 1.75 | (129) |
| Cu      |          | 8.70–196.7                  | PLSR | 0.80 | 1.14 |
| Zn      |          | 24.2–577.3                  | PLSR | 0.95 | 1.56 |
| Pb      |          | 7.10–767.0                  | PLSR | 0.56 | 1.08 |
| Cr      |          | 25.9–79.0                   | PLSR | 0.81 | 1.36 |
| Al      | 34 European countries | 2500–20000 | 329–65090 | De-trend | 0.83 | 0.13 | 2.4 | (130) |
| Fe      |          | 380–133926                  | De-trend | 0.79 | 0.16 | 2.2 |
| Ni      |          | 0.05–2475                   | De-trend | 0.77 | 0.24 | 2.1 |
| Zn      |          | 1.50–1396                   | De-trend | 0.60 | 0.19 | 1.6 |
| Cu      |          | 0.30–395                    | De-trend | 0.55 | 0.26 | 1.5 |
| Mn      |          | 1.60–14969                  | De-trend | 0.54 | 0.27 | 1.5 |
| Pb      |          | 1.60–1309                   | De-trend | 0.42 | 0.24 | 1.3 |
| Se      |          | 0.08–6.8                    | De-trend | 0.40 | 0.19 | 1.3 |
| Cd      |          | 0.005–7.5                   | De-trend | 0.40 | 0.29 | 1.3 |
| Element | Country | Sample Size | Range | Mean | R² | SD | Linear Regression | SNV + FD | 4000–130,000 | 0.85 | 6672.48 | (21) |
|---------|---------|-------------|-------|------|----|----|-------------------|---------|---------------|-------|----------|------|
| Hg      | Brazil  | 100         | 2500–16,666 | Absorbance 14.3–65 | PLSR 0.91 | 16.1 | 3.4 | (131) |
| Ag      | Brazil  | 100         | 2500–16,666 | Absorbance 14.3–65 | PLSR 0.91 | 16.1 | 3.4 | (131) |
| Ba      | Brazil  | 100         | 2500–16,666 | Absorbance 14.3–65 | PLSR 0.91 | 16.1 | 3.4 | (131) |
| Co      | Germany | 100         | 2500–15,000 | SNV +FD 4000–130,000 | 0.85 | 6672.48 | (21) |
| As      | Germany | 100         | 2500–15,000 | SNV +FD 4000–130,000 | 0.85 | 6672.48 | (21) |
| Cu      | Germany | 100         | 2500–15,000 | SNV +FD 4000–130,000 | 0.85 | 6672.48 | (21) |
| Fe      | Germany | 100         | 2500–15,000 | SNV +FD 4000–130,000 | 0.85 | 6672.48 | (21) |
| Mn      | Germany | 100         | 2500–15,000 | SNV +FD 4000–130,000 | 0.85 | 6672.48 | (21) |
| Ni      | Germany | 100         | 2500–15,000 | SNV +FD 4000–130,000 | 0.85 | 6672.48 | (21) |
| Zn      | Germany | 100         | 2500–15,000 | SNV +FD 4000–130,000 | 0.85 | 6672.48 | (21) |

**PXRFS**

| Element | Country | Sample Size | Range | Linear Regression | 0.97 | (63) |
|---------|---------|-------------|-------|-------------------|-------|------|
| Pb      | UK      | 81          | 5-40398 | linear regression | 0.97 | (63) |
| Cd      |         |             | 0-447  |                    | 0.46 |      |
| As      |         |             | 2.0-5646 |                  | 0.87 |      |
| Ni      |         |             | 1.0-84.0 |                 | 0.34 |      |
| Zn      |         |             | 3.0-25389 |                | 0.88 |      |
| Mn      |         |             | 6.0-38267 |                | 0.74 |      |
| Element | Location | Number | Range | Method | Slope | Reference |
|---------|----------|--------|-------|--------|-------|-----------|
| Fe      |          |        | 482-91566 |       | 0.97  |           |
| Cu      |          |        | 3.0-5140 |       | 0.88  |           |
| Pb      | USA      | 76     | 152-3590 | linear | 0.98  | 4.1       | (70) |
| Pb      | Ireland  | 17     | 80-30000 | linear | 0.99  |           | (62) |
| AS      |          |        | <LOD-1230 |       | 0.99  |           |
| Cu      |          |        | 10-910  |       | 0.95  |           |
| Zn      |          |        | 260-9000 |       | 0.84  |           |
| Cd      | South Korea | 30 | linear |       | 0.96  |           | (61) |
| Pb      |          |        |         |       | 0.97  |           |
| Cu      |          |        |         |       | 0.94  |           |
| Zn      |          |        |         |       | 0.76  |           |
| As      |          |        |         |       | 0.97  |           |
| Ti      | Australia | 75   |         |       | 0.95  |           | (22) |
| Cr      |          | 72     |         |       | 0.87  |           |
| Mn      |          | 75     |         |       | 0.99  |           |
| Fe      |          | 75     |         |       | 0.98  |           |
| Ni      |          | 27     |         |       | 0.51  |           |
| Cu      |          | 75     |         |       | 0.99  |           |
| Zn      |          | 75     |         |       | 0.99  |           |
| As      |          | 45     |         |       | 0.97  |           |
| Sr      |          | 75     |         |       | 0.97  |           |
| Cd      |          | 23     |         |       | 0.99  |           |
| Pb      |          | 75     |         |       | 0.99  |           |
|   |   |   |   |   |   |   |
|---|---|---|---|---|---|---|
| Al | Ireland | 322 | 0.6-9.7 | Cubist | 0.88 | 0.6 | 2.11 | (30) |
| As |   |   | 0.7-51.6 |   | 0.55 | 5.22 | 0.58 |
| Ba |   |   | 44.3-1178.9 |   | 0.90 | 40.35 | 1.66 |
| Cd |   |   | 0.01-5.0 |   | 0.86 | 0.23 | 0.57 |
| Co |   |   | 0.7-58.7 |   | 0.80 | 2.86 | 0.84 |
| Cr |   |   | 5.5-221.7 |   | 0.87 | 8.52 | 1.73 |
| Cu |   |   | 1.8-104.0 |   | 0.93 | 3.77 | 1.71 |
| Fe |   |   | 0.2-5.4 |   | 0.98 | 0.14 | 4.96 |
| Hg |   |   | 0.01-1.0 |   | 0.59 | 0.05 | 0.40 |
| Mg |   |   | 0.1-1.5 |   | 0.61 | 0.14 | 0.71 |
| Mn |   |   | 16.0-10287.0 |   | 0.90 | 316.41 | 0.90 |
| Mo |   |   | 0.1-14.4 |   | 0.82 | 0.52 | 0.57 |
| Ni |   |   | 1.7-176.0 |   | 0.84 | 5.84 | 1.27 |
| Pb |   |   | 5.2-123.0 |   | 0.82 | 7.21 | 1.02 |
| As | UK | 75 |   |   | 0.70 |   | (56) |
| Zn |   |   |   |   | 0.81 |   |   |
| Cr | China | 111 |   | Wavelet | 0.99 |   | (132) |
| Zn |   |   |   |   | 0.99 |   |   |
| Pb |   |   |   |   | 0.99 |   |   |
| As |   |   |   |   | 0.98 |   |   |
| Zn | China | 301 | 59.0-4194 | PLSR | 0.99 | 28.9 | 9.35 | (75) |
| Ni |   |   | 2.0-114 |   | 0.37 | 33.0 | 0.36 |
| Cu |   |   | 8.0-297 |   | 0.95 | 12.37 | 2.88 |
| Pb |   |   | 15.0-159 |   | 0.92 | 12.0 | 2.53 |
|   |   |   |   |   |
|---|---|---|---|---|
| As | Italy | 294 | 1.0-53.0 | 0.73 |
| Cu | Italy | 294 | 8-485 | 0.95 |
| Ni | | | Linear regression | |
| Zn | | | 0.97 | |

**LIBS**

| Cr | Italy | 20 | 400-700 nm | RSD % |
|----|-------|----|------------|-------|
| Cu | | | | (134) |
| Fe | | | | |
| Mn | | | | |
| Ni | | | | |
| Pb | | | | |
| Zn | | | | |

| Cu | Brazil | 59 | ANN | 0.96 |
|----|--------|----|-----|------|
| Cr | Italy | 6 | | 0.92 |
| Al | Spain | 19 | without | 0.77 |
| Ba | | | PLSR | 0.30 |
| Ca | | | | 1.57 |
| Fe | | | 0.80 | 0.10 |
| Mg | | | SNV | 2.63 |
| Si | | | SNV | 1.96 |
| Cd | China | 50 | Wavelet | 211.9–232.9 nm |
| | | | PLSR | 0.97 |
| | | | LS-SVM | 0.034 |

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| Element | Country | Sample Size | LOD (mg kg⁻¹) |
|---------|---------|-------------|---------------|
| Cu      | China   | 25          | 0.98 6.28     |
| Pb      |         |             | 0.99 8.64     |
| Zn      |         |             | 0.99 8.55     |

Stepwise-MLR; MARS, multivariate adaptive regression splines; UR, univariate regression; AB, absorbance; CR, continuum removal; FD, first derivative; SEP, standard error of prediction; UVN, unit vector normalization; K-M, Kubelka–Munk; MSC, Multiplicative scatter correction; SNV, standard normal variate spectra.
Table 2. Advantages and limitations of most common laboratory and field optical and spectroscopic techniques for determining potential toxic elements (PTEs) in soil samples.

| Price range (1000$) | Targeted analytes | Sample matrix | Cost per sample ($) | Limit of detection (mg kg\(^{-1}\)) | Accuracy range | Advantages | Limitations | References |
|---------------------|-------------------|---------------|---------------------|------------------------------------|----------------|------------|-------------|------------|
| **PXRF**            |                   |               |                     |                                    |                |            |             |            |
| 25-40               | 26                | Solid         | 10-20               | 1-10%                             | Fast, non-destructive, accuracy and precision increase with higher element concentration, capable of analysing a large range of elements from F to U, and well suited for the analysis of highly homogeneous materials. | Sensitive to matrix effect, relatively expensive to run, and analysis is slower. | (19, 50–52, 55, 56, 154) |
| **LIBS**            |                   |               |                     |                                    |                |            |             |            |
| 35-50               | 10-15             | Solid         | 15-20               | 1-10%                             | Fast, destructive, accuracy and precision increase with higher element concentration, capable of analyzing a large range of elements, and well suited for | Sensitive to matrix and water content effect, relatively expensive to run. | (24, 77, 82, 89, 94) |
| Price range (1000$) | Targeted analytes | Sample matrix | Cost per sample ($) | Limit of detection | Accuracy range | Advantages | Limitations | References |
|---------------------|-------------------|---------------|---------------------|-------------------|---------------|------------|-------------|-----------|
| the analysis of highly homogeneous materials. | | | | | | | | |

**Vis-NIRS**

| 20-40 | - | All matrices | 10-20 | - | - | Fast, non-destructive, portable, and suitable for on-line measurement, accuracy and precision increase in more homogeneous materials. | Sensitive to matrix effect, needs to be calibrated with reference samples, effects of moisture, texture are moderate on accuracy. | (16, 27, 29, 106, 107) |

**MIRS**

| 30-40 | - | All matrices | 10-20 | - | - | Very fast, non-destructive, portable, allows in situ measurement, accuracy and precision increase in more homogeneous materials | Sensitive to matrix effect and soil moisture, needs to be calibrated with reference samples, and effects of moisture, texture are high on accuracy and larger than in the Vis-NIRS | (21, 30, 120, 143, 155) |

Vis-NIRS = visible and near-infrared spectroscopy; LIBS = laser-induced breakdown spectroscopy; MIRS, mid-infrared spectroscopy; PXRF = portable x-ray fluorescence spectroscopy; LOD = limit of detection.