THE COMMON PITFALL OF USING ENRICHMENT FACTOR IN ASSESSING SOIL HEAVY METAL POLLUTION

(Kepincangan Penggunaan Faktor Pengkayaan dalam Menilai Pencemaran Logam Berat di dalam Tanah)

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Abstract

A soil pollution assessment of heavy metal Zn, Cu, Cr, Cd, Ni and Pb was carried out in Kuala Terengganu, Malaysia. A total of 245 surface soil samples were sampled and have been subjected to a total digestion and analyzed by inductive couple plasma-optical emission spectrometry (ICP-OES). The contamination of the soils was assessed on the basis of enrichment factors (EFs). Large discrepancies in results of enrichment factor associated with the choices of reference elements in the EFs’ calculation have been observed. Our study show that the choice of four commonly-used reference elements (Al, Fe, Ca and Mn) in enrichment calculation could easily alter the enrichment factor values of element of interest by more than 4-folds although the results were supposed to be similar regardless of which reference elements being used. This could lead to serious misinterpretation of soil quality assessment result.

Keywords: soil quality assessment, Malaysia soil, elemental analysis, acid digestion

Abstrak


Kata kunci: kajian kualiti tanah, tanah Malaysia, analisa unsur, penghadaman asid

Introduction

Heavy metals like Cd, Pb, Zn, Cu, Cr and Ni in soil have received wide attention due to their potential toxic impact on the environment if present in excessive concentrations. It is critically important to carry out soil quality monitoring program on a regular basis, particularly in urban watershed to support sustainable planning and development for better livelihood. There are three common approaches of assessing soil quality: (1) take measurements periodically over the time to monitor changes or trends in soil quality; (2) compare measured values to a standard or reference soil condition (i.e. Netherlands’ Dutch Target, Australia’s Soil Investigation Level,
USEPA Ecological Screening Levels, British Soil Guideline and Japanese Environmental Quality Standards). The former method is commonly used by stakeholders to identify areas where problems occur, to identify areas of special interest, or to compare fields under different management systems over time and latter generally adopted by law enforcement authorities for soil pollution control and mitigation. Finally, the third approach is based on calculation of enrichment factors (EFs) to evaluate anthropogenic influences on soil quality.

Enrichment factor is a proxy to assess the heavy metals contamination level in soil by estimating the differential of heavy metal concentrations against uncontaminated background or reference levels [1,2]. An enrichment factor can be defined as in equation 1:

$$EF_i = \frac{y/x}_{\text{sample}} / \frac{y/x}_{\text{background}}$$  \hspace{1cm} (2)

where, $y$ is concentration of target metal, $x$ is concentration of reference element or normalizer in soil and in background or baseline reference material. In soil quality study, the interpretation of enrichment factor is based on the deviation of the ratio between a metal of interest and a reference element that exhibit conservative and immobile behavior in environment in a sample to the equivalent ratio in a reference material from unity ($EFs = 1$). Enrichment factor equal to one is generally taken to mean that there has been no net gain or loss of a metal in the sampled soil relative to the respective material. Enrichment factor ratio should increase ($EFs > 1$) if there has been a net gain of metal concentration due to extra input (e.g. anthropogenic sources), and a net loss of metal concentration in the system (e.g. due to weathering, biological assimilation) should decrease the ratio value ($EFs < 1$).

Enrichment factor is practice widely in environmental studies. Nevertheless, the assumption of enrichment factor as pollution detection tool inherits many theoretical shortcomings [3,4]. The work done by Reimann and de Caritat [3,4] concluded that EFs calculation is strongly influenced by the natural variable composition of the reference material, biogeochemical processes and physico-chemical alteration of elements in crust materials. The authors regarded using EFs to interpret pollution event in any environmental studies as doubtful. The primary objective of this study is to determine the metal concentrations in soil of Kuala Terengganu. Fortuitously, this study will allow to test the reliability of enrichment factor as a pollution proxy for soil quality assessment.

Materials and Methods

Soil sampling

The soil quality assessment was carried out in Kuala Terengganu district, Malaysia (5.33°N, 103.1°E, Figure 1). Sampling of soil had been accomplished on 245 cells of 1 km x 1 km square grid using the topographic maps. About 1 kg of soil sample was collected from the first 10-15 cm of the upper soil horizon using a plastic spade at each grid. The collected samples were air-dried, ground, and passed through sieve of a 0.6 mm mesh size for subsequent acid digestion.

Soil digestion and chemical analysis

All soil digestion and element analysis was performed in analytical chemistry laboratory of Universiti Malaysia Terengganu (UMT). All apparatus used in this experiment were acid-cleaned with 10% HNO$_3$. The soil digestion procedure for heavy metal analysis was carried out using USEPA method 3052. In general, an acid mixture of HNO$_3$:HCl:HF (9:4:1) ml ratio was added to 0.5 g of soil sample inside a Teflon vessel. Later, digestions of soils were performed in a microwave-heated oven (Ethos Plus Milestone, USA) at 210 °C for 20 minutes. After digestion, small amount of saturated H$_3$BO$_3$ acid was added into solution to neutralize remaining active HF in solution. The digested solution then makes up to 25 ml of final volume with ultrapure deionized water. A complete digested soil sample will normally yield a clear solution. Quality control and quality assurance also carried out by using two standard reference materials SRM-2709 and SRM-2711 obtained from the National Institute of Standards and Technology (NIST, USA). The SRMs were subjected to similar sample treatment procedures as described above. All samples and SRMs were analyzed for Al, Fe, Mn, Ca, Zn, Cu, Cr, Cd, Ni and Pb by inductive couple plasma optical emission spectrometer (Varian Vista Pro, Australia). SRMs recovery result shows that the analytical accuracy for all measured metals in SRM (n = 12) was within 5-15% error for all metals, expect for Cr which was 10-25% error.
Enrichment factor calculation and geochemical mapping

The calculation of a normalized enrichment factor for a given metal for this study is represented by equation 1. Where, $EF_i$ is the enrichment factor of target metal $i$, $(y/x)_{sample}$ is the target metal to reference (conservative) element’s ratio in the samples of interest; while $(y/x)_{background}$ is the target metal to reference metal ratio in world soil baseline reference taken from Ure and Berrow [5].

Results and Discussion

Heavy metals concentration in Kuala Terengganu soil

The concentration range of metals in Kuala Terengganu soils and their mean values are given in Table 1. The mean concentration values of all metals, except Cd in surface soil of Kuala Terengganu were lower than those reported in worldwide average values [5].

Level of Al (0.45%), Ca (0.14%), Mn (82.9 mg/kg) in the soil samples studied was found much lower (10 fold) than those reported by Ure and Berrow [5] for world average values 6.65%, 1.96% and 760 mg/kg, respectively. The studied soil samples were relatively poor in Fe, Cu, Cr, Zn and Ni. The mean values of these metals were lower than those reported by Zarcinas et al. [7] for Malaysian agricultural soils (Table 2) and for world average value [5]. Level of Pb (23.6 mg/kg) detected in the soil samples were similar to those found in Zarcinas et al. [7] (26.4 mg/kg) and worldwide average value (29.2 mg/kg). Conversely, the content of Cd in studied soil ranged between 0.38 and 6.78 mg/kg with the median concentration of 1.28 mg/kg and was found to be two times greater than the mean concentration of world average (0.62 mg/kg).

The Baging, Rhutapai, Rusila and Rudua soil series associated to BRIS (Beach Ridges Interspersed with Swales) were reported as dominant soils in these areas [6]. The low heavy metal concentrations in Kuala Terengganu soil indicates the sampled soils are mainly originated from highly weathered BRIS soils. The decreasing order of concentration on the different metals Fe>Al>Ca>Mn>Zn>Pb>Cr>Cu>Ni>Cd reflects their abundance in BRIS soil are associated with low cation exchange capacity, low organic matter content, and excessively-drained sandy soil properties promote leaching losses.
Table 1. The range, median, mean of selected metals in surface soil of Kuala Terengganu (in mg kg\(^{-1}\), n=245)

<table>
<thead>
<tr>
<th>Metal</th>
<th>Range</th>
<th>Median</th>
<th>Mean</th>
<th>Worldwide Average [5]</th>
</tr>
</thead>
<tbody>
<tr>
<td>Al*</td>
<td>0.14-1.44</td>
<td>0.42</td>
<td>0.45</td>
<td>6.65</td>
</tr>
<tr>
<td>Fe*</td>
<td>0.22-7.70</td>
<td>1.54</td>
<td>1.57</td>
<td>3.20</td>
</tr>
<tr>
<td>Mn</td>
<td>2.53-356</td>
<td>51.3</td>
<td>82.9</td>
<td>760</td>
</tr>
<tr>
<td>Ca*</td>
<td>0.01-1.17</td>
<td>0.08</td>
<td>0.14</td>
<td>1.96</td>
</tr>
<tr>
<td>Zn</td>
<td>4.61-204</td>
<td>31.4</td>
<td>38.3</td>
<td>59.8</td>
</tr>
<tr>
<td>Cu</td>
<td>0.82-148</td>
<td>9.39</td>
<td>10.9</td>
<td>25.8</td>
</tr>
<tr>
<td>Cr</td>
<td>3.05-50.1</td>
<td>21.6</td>
<td>20.5</td>
<td>84.0</td>
</tr>
<tr>
<td>Cd</td>
<td>0.38-6.78</td>
<td>1.21</td>
<td>1.28</td>
<td>0.62</td>
</tr>
<tr>
<td>Ni</td>
<td>1.91-16.7</td>
<td>7.18</td>
<td>7.02</td>
<td>33.7</td>
</tr>
<tr>
<td>Pb</td>
<td>2.54-160</td>
<td>21.9</td>
<td>23.6</td>
<td>29.2</td>
</tr>
</tbody>
</table>

*values reported in weight percent, wt.%

Soil pollution assessment of heavy metals using enrichment factor
To allow quantitative comparisons between degrees of metals enrichment in the soil pollution assessment, Sutherland [8] and Loska et al [9] had distinguished five contamination categories of the enrichment factor. An EF < 2 is deficiency of minimal enrichment, suggestive of no or minimal pollution; EF 2 - 5 is moderate enrichment; EF 5 - 20 is significant enrichment; EF 20 - 40 is very high enrichment and EF > 40 is extremely high enrichment, indicating an extreme pollution signal. Several elements are utilized as normalizer in enrichment factor, in particular element that have been associated with rock-forming minerals (see summary in Table 2).

Table 2. Selection and justification of reference element for enrichment factor calculation in relation to anthropogenic impact

<table>
<thead>
<tr>
<th>Normalizer</th>
<th>Justification</th>
<th>Reference</th>
</tr>
</thead>
<tbody>
<tr>
<td>Al</td>
<td>Al has no significant anthropogenic input compared to the large of geogenic element; Ion potential of Al is close to target elements, reduce variation in comparison</td>
<td>[10]</td>
</tr>
<tr>
<td>Mn</td>
<td>Mn is one of largest components of soil</td>
<td>[11]</td>
</tr>
<tr>
<td>Fe</td>
<td>Fe is abundant in soils; considered to be free from anthropogenic contribution</td>
<td>[1, 12]</td>
</tr>
<tr>
<td>Ca</td>
<td>Ca is one of the main components of the Earth’s Crust, concentration in soil is connected with the matrix, lower mining interference factor (MIF) compared to Al and Zr</td>
<td>[9]</td>
</tr>
<tr>
<td>Zr</td>
<td>Zr has no significant anthropogenic source and resides primarily in the weathering-resistant heavy mineral Zircon</td>
<td>[13]</td>
</tr>
<tr>
<td>Sc</td>
<td>Sc is stable to supergene processes associated with oxidation and chemical weathering</td>
<td>[14]</td>
</tr>
<tr>
<td>Sr</td>
<td>Sr is one of the main components of the Earth’s Crust, concentration in soil is connected with the matrix</td>
<td>[15]</td>
</tr>
<tr>
<td>Ti</td>
<td>Ti is a common rock-forming element and very resistant to weathering in the soil.</td>
<td>[16]</td>
</tr>
</tbody>
</table>

The scatter plot in Figure 2 illustrates a comparison study of selected elements EF against different normalizers (Fe, Al, Ca and Mn) and the obtained results showed that EF values of metal were different corresponding to the
normalizers used. For better interpretation, four different colour lines representing the contamination categories scale suggested by Sutherland [8] and Loska et al. [9] were plotted (Figure 2). For example, if a sample fall within green and yellow line, then the sample was classified as being moderate enriched (EF = 2-5). If it falls between the yellow and orange line, then the sample was classified as significant enrichment (EF = 5-20). If a sample surpassed the orange line but was within the red line, then the sample was classified as being very highly enriched (EF = 20-40). Samples with the enrichment factor greater than 40 that were found to have surpassed the red line were classified as samples with severe pollution. If samples fall below green line (EF < 2), then samples were classified as depletion to minimal enrichment suggestive of no or minimal pollution.

Figure 2. Comparison of enrichment factor for target metals using different normaliser; green, yellow, orange and red colour lines representing contamination categories scale proposed by Loska et al. [9] and Sutherland [8]
Theoretically, the selection of reference element used in EFs calculation should hardly be affected by weathering processes and should show little variability of occurrence [3]. If different reference elements are selected to calculate EFs, they should all show similar pollution signals regardless of which normalizers were being used. In this study, however, all four sets of EFs calculation (refer Figure 2) bear no relationship to each other.

Figure 2 show EFs values for all metals derived from all four normalizers differ in the range from 1 to 4-folds. EFs derived from Ca and Mn normalizer generally had overestimated amount compared to Al and Fe. We found that 93% and 6% of the Cd samples were classified as extremely high polluted and very high polluted respectively when Mn was used as reference element. These outcomes are very different to the other reference elements, in particular to Fe normalizer. For Fe reference-EFs, 95% of Cd samples were classified as moderate enriched or lower. Al and Ca normalizer also produced a wide range of EFs for Cd samples and these values ranged between Fe and Mn references derived EFs. These discrepancies reflect that EFs do not provide a simple technique to differentiate particular suite of metals was whether of anthropogenic origin or vice versa.

The color contour maps of Pb concentration and EFs in Figure 3 were created using the inverse distance weighting (IDW) spatial interpolation technique in ESRI ArcMap 9.2. In Figure 3, the Pb hotspots in Kuala Terengganu soils are marked in red colors. The Pb concentration for Kuala Terengganu soils were also compared to Dutch environmental pollutant reference values, also known as “Dutch target” used in environmental remediation, investigation and cleanup purposes. For Kuala Terengganu, three out of 245 samples had exceeded the Dutch recommendation value of 85 mg/kg. The spatial variations for each metal for Kuala Terengganu soil in this study can be found in Poh [17].

![Figure 3. Comparison of the calculated EFs for Pb in Kuala Terengganu soils](image-url)
Three striking features of the EFs pattern of Pb are evident in Figure 3. First, the EFs for Pb show great spatial variation compared to Pb distribution pattern in Kuala Terengganu soil. Second, this variation is multidimensional; that is three different set of reference element show mainly independent of EFs variation. For example, when EFs was calculated relative to the Pb/Al ratio, the pattern of high Pb enrichment was recorded at the same areas in the Pb concentration map. However, when the EFs Pb/Fe ratio distribution was compared to Pb concentration map, it turns out, several high enrichment values near the central part of the map disappeared. Furthermore, when EFs Pb/Ca ratio distribution was compared, the high anomaly near to river mouth was no longer present in the map. Meanwhile, an unexpected extremely high EFs had appeared scattered in the central part of the map. Third, these EFs patterns have a pervasive missed-interpretation of pollution signal. In this case, EFs Pb/Al ratio distribution showed 50% of Kuala Terengganu soils were significantly enriched by Pb but EFs Pb/Fe ratio distribution, on the other hand illustrated that 99% of areas were still less contaminated by Pb. The results could overestimate, underestimate, or simply incorrectly estimate the environmental pollution.

The potential sources of errors for EFs technique mentioned above can be explained in a number of ways. First, a huge natural variation in concentration of the different elements in reference materials (i.e. continental crust value, world soil average) exist and using single mean value from a skewed distribution dataset (i.e. Ure and Berrow [5]) does not permit a reasonable EFs assessment. Ure and Berrow [5] reported Pb concentration in world soil is ranges from 1 – 888 mg/kg with average value of 29.2 mg/kg, obviously the given mean value does not reflect the true average Pb concentration for world average soil. This is one of a very good example to show that using the mean world soil value as a proxy of “natural background values” in enrichment factor inherent a serious flaw to estimate pollution signal in environment.

Second, an inhomogeneous distribution of reference elements in the study area could also lead to misinterpretation in EFs. Overestimation of EFs occurs when the chosen reference elements (normalizer) showed a natural spatial variation that was as large as, or larger than, that of the target metal of interest [4]. In this study, we used quartile coefficient of dispersion (cqv) statistical analysis to compare the dispersion of the maximum concentration differences between Pb and other reference elements. The calculated cqv for Ca, Fe and Al against Pb was 2.2, 1.1 and 0.7 respectively, indicated that the natural spatial variation of Ca was found 2 times greater than Pb. We found that the large dispersion of Ca concentration has attributed greater EFs value of Pb in Kuala Terengganu and the Pb/Ca ratio EFs was found greater than Pb/Al and Pb/Fe ratio EFs by order of two magnitudes (Figure 3).

The large discrepancies in EFs value could be due to two other factors: (i) the differential solubility of refractory minerals in acid digestion protocols as well as grain size fractionation effect resulted different concentration of metal elements in Earth materials and; (ii) the significance of biogeochemical and weathering processes that control element concentration in Earth surface are not considered in EFs assessment on pollution studies. Reimann and de Caritat [3,4] and Desaules [18] have provide a detailed explanation and references in conceptual shortcomings of EFs in environmental studies.

**Conclusion**

Our work demonstrated that application of enrichment factor in the assessment of soil pollution level in Kuala Terengganu inherent a major weakness. One unanticipated finding was that the choice of four commonly-used reference elements (Al, Fe, Ca and Mn) for enrichment calculation could easily alter the enrichment factor values of Pb by more than 4-folds although the results were supposed to be similar regardless of which reference elements being used. This bias can be attributed to the absent of a reliable “natural background value” or references. Another shortcoming of EFs approach in assessing the level of anthropogenic sources to environment is EFs did not take into account the natural variation of element concentration in environmental samples in relation to biogeochemical and localized lithogenic processes as mentioned in Reimann and de Caritat [3,4] and Desaules [18]. Therefore, application of enrichment factor as a way to fingerprint the anthropogenic source of heavy metals in soils requires extra caution in future study.

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References