Heavy metal fractionation studies in tidal sediment cores in the clam farms from Tan Thanh commune, Go Cong district, Tien Giang province, Vietnam

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This paper introduces the results from a study on the distribution of heavy metals in chemical fractions in tidal sediment cores at four sample stations inside the clam breeding plain, inside the harvested clam breeding plain, and on the frontier between the plains in the farms in Tan Thanh commune, Go Cong district, Tien Giang province, Vietnam. The partitioning of metals among the compartments of the sediment's solid phase was investigated indirectly by selective sequential extraction of substances that are water-soluble, exchangeable, bound to carbonates, bound to Mn oxides, bound to amorphous Fe oxides, bound to crystalline Fe oxides, associated with organics and residual. In case of investigated heavy metals (HM), the concentrations of Zn and Hg exceeded the National Technical Regulation on Sediment Quality QCVN 43/2012/BTNMT by 1.12 - 3.53 times and 26.58 - 171.96 times, respectively. The highest HMs concentration was found in the oxidable fraction (more than 60%). The data demonstrates the important role of organic matters in the oxidizable condition at the surficial sediment layer. Besides the oxidable fraction, high HMs concentrations were measured also in the residual fraction. The HMs content present in the solid residue also indicates the level of contamination in the river system: the greater the percentage of HMs present in the solid residue, the lesser the pollution in the environment because this solid residue involves components that can not be remobilized. The HMs concentrations depending on the depth of the sediment indicate that clam's digestive activity or the decomposition of tissue and shell of clams possibly affects the content of HMs.

Bài báo giới thiệu các kết quả nghiên cứu về sự phân bố kim loại nặng dưới các dạng liên kết khác nhau trong các mẫu môi trường của các bãi nuôi nghêu giống, bãi nuôi nghêu sau khi đã thu hoạch, và ranh giới giữa các bãi nuôi nghêu tại xã Tân Thanh, huyện Go Công Đông, tỉnh Tiền Giang, Việt Nam. Các dạng pha liên kết bao gồm: 1. pha hòa tan; 2. pha trao đổi; 3. liên kết với các chất 다만; 4. liên kết với Man gan ở xỉ; 5. liên kết với sắt ở xỉ và dòng hình; 6. liên kết với sắt ở xỉ dạng tĩnh thể; 7. liên kết với thành phần hữu cơ và; 8. phần bã rắn. Trong các kim loại nặng (KLN) được phân tích, hàm lượng Zn và Hg vượt quá tiêu chuẩn cho phép QCVN 43/2012/BTNMT lần lượt từ 1,12 - 3,53 và 26,58 - 171,96 lần. HBM lượng KLN tồn tại nhiều nhất dưới dạng liên kết với các thành phần có khả năng ở xỉ hòa tan với 60% chỉ ra rằng các thành phần hữu cơ và trong điều kiện ở xỉ hòa tan lơ lửng. Các thành phần có khả năng ở xỉ hòa tan các KLN hiện diện trong phần bã rắn nhiều hơn trong các pha khác. Sự có mặt của KLN trong phần bã rắn chỉ ra mức độ ảnh hưởng của hệ thống sông; càng nhiều phần trăm KLN có mặt trong phần bã rắn, càng ít ở nhiễm trong môi trường bã phân bã rắn này liên quan đến các thành phần không thể bị rụng tích lũy. Sự phụ thuộc của độ sâu của hàm lượng kim loại nặng dưới đáy nước và sự hình thành của hoạt động tiêu hóa và quá trình phân hóa của nghêu liên hâm lượng kim loại nặng.

1. Introduction

Vietnam ranks the third after China and India in the aquaculture industry and has been growing strongly since 2003. With a coastline of more than 3,200 km long with over 3,000 islands, a wealth of natural inland water bodies (lakes and rivers) and seasonally flooded grounds, since 2000, the fisheries sector is an important contributor to the economy of Vietnam and fisheries are identified as a key economic growth sector by the Vietnamese Government (MOFI, 2006). Seventy percent of coastal aquaculture production of Vietnam occurs in the southern area, in the Mekong River Delta (MOFI, 2006). With over 700 km of coastline, the coastal aquaculture of the Mekong River Delta has been well known for the farming of marine shrimps and mollusks. However, in the past few years, the phenomenon of widespread mass clam deaths had occurred in this Delta (Zing (2013), Nhan Dan (2011), Bao Moi (2013), Nong.
The aim of the present study is to investigate different chemical forms of HMs in sediments and to assess their association and bioavailability in the river.

2. Materials and methods

2.1 Sampling site

The Tan Thanh commune of Go Cong Dong district, Tien Giang province is located in the Mekong Delta region of Vietnam where famous clam farms are situated. The area selected for this study is between the Den Do hamlet and Tan Thanh resort (Figure 1) with a water area of about 2,000 hectares (ha) used for clam aquaculture. The elevation of this tidal area is from 0.6 to -6.0 m. The core sediments were collected using a gravity type sediment corer of a 1m length of polycarbonate tube. Our research focuses on the 0.4 m above of core-length. The sampling stations include: P8 (10°17'20"N, 106°46'58"E) and P10 (10°16'53"N, 106°46'44"E) inside the baby clam breeding plain; P9 (10°17'06"N, 106°46'52"E) inside the harvested clam breeding plain; P11 (10°16'38"N, 106°46'28"E) on the frontier between the plains (Figure 1). All samples were collected in April, during the dry season.

Figure 1 - Location of the sample stations in the research region - Tan Thanh, Go Cong Dong, Tien Giang

2.2 Sampling and analysis

This study focuses on the surficial sediment (0 - 60 cm) in a tidal wetland in the estuary Tieu, Tan Thanh, Go Cong Dong, Tien Giang, which is the sediment layer reflecting the historical pollution (Salomons and Forstner, 1984; Forstner, 1989). This layer is also the living environment of clams, which can grow, digest and die at the depth of 20 cm. For this reason, the sediment cores were cut after collection in three slices: from 0 to 20 cm, 20 to 40 cm, and 40 to 60 cm of depth. The samples were stored in freeze room at 4°C. At position Tan Thanh P11 (TT - P11), the core has just two slices (from 0 to 40 cm) because the third one was damaged after cutting.

To measure the total HMs concentration in every slice, the sediment sample was digested following the method of Jarvis et al. (1992) by using acid mixture HF/HClO₄ and then...
HNO₃ before being analysed by Inductively Coupled Plasma Spectrometer (ICP-MS, Ultramas-700) in the laboratory of the Institute of Geological Sciences (Vietnam Academy of Science and Technology).

Table 1. Selective sequential extraction (SSE) procedure

| No | Fraction                  | Extractant                             | Vol. | Time | Temp. | pH  |
|----|---------------------------|----------------------------------------|------|------|-------|-----|
| F1 | Water soluble             | deionized water                        | 10ml | 30 min | 20°C | 7   |
| F2 | Exchangeable              | 1M Mg(NO₃)₂                            | 10ml | 2 h  | 20°C | 3   |
| F3 | Bound to carbonate        | 1M NaCOOCH₃                            | 10ml | 5 h  | 20°C | 4   |
| F4 | Bound to Mn - O           | 0.1M NH₄OH.HCl                         | 10ml | 30 min | 20°C | 3.5 |
| F5 | Bound to amorphous Fe-O   | 0.2M (NH₄)₂C₂O₄.H₂O + 0.2M             | 10ml | 4 h  | 20°C | 3   |
| F6 | Bound to crystalline Fe-O | 0.2M (NH₄)₂C₂O₄.H₂O + 0.2M             | 10ml | 30 min | 80°C | 2   |
| F7 | Oxidizable                | 0.02M HNO₃ + 30% H₂O₂                  | 3ml  | 5 h  | 85°C | 2   |
| F8 | Residues                  | H₂O pure                               | 10ml | 30 min | 85°C | 2  |

The partitioning of metals among the compartments of the sediment solid phase was investigated indirectly by selective sequential extractions (SSE), as described in Table 1. The SSE was performed with 1 g of ground sediment in 50 ml polypropylene centrifugation tubes to minimize losses of material. All extractions were performed in duplicate on three replications of each treatment. In each extraction series, a standard sample was introduced to follow the reproducibility of the procedure. After each extraction step, the tubes were centrifuged at 5200 rpm for 20 min. The supernatants were then filtered through 0.45 μm membranes (Sartorius), whereas the residues were washed with 10 ml of ultra-pure water, centrifuged again, and then the supernatants were pooled. The leachates (extract and rinsing) were stored in polypropylene bottles or glass vials at 4°C until chemical analysis. The residues were dried at 40°C to remove all of the water prior to the next extraction step. Blanks without a sediment sample were used on each extraction step to determine the purity and quality of the procedure. The chemical forms were labelled according to the targeted geochemical compartments during each extraction step: F1 - water soluble (WAT), F2 - exchangeable (EXCH), F3 - bound to carbonates (CAR), F4 - bound to Mn oxides (MNOX), F5 - bound to amorphous Fe oxides (FEOX1), F6 - bound to crystalline Fe oxides (FEOX2), F7 - associated with organics (OM) and F8 - residual (RES). Each element in the different fractions was expressed as micrograms extracted per gram of soil and as a percentage of the total amount of metal extracted after the seven steps. The elements were analysed by ICP-MS.

3. Results and discussion

3.1 Total HMs contents

The results of total HMs content of every slice (TT-Px-1: 0 - 20 cm, TT-Px-2: 20 - 40 cm, and TT-Px-3: 40 - 60 cm) at 4 sample stations (P8 and P10: inside the baby clam breeding plain; P9: inside the harvested clam breeding plain; P11: on the frontier between the plains) are given in Table 2. These results are compared with the National Technical Regulation on Sediment Quality of Vietnam (QCVN 43:2012 BTNMT). This standard is the same with the standard given by Canadian sediment quality guidelines for the protection of aquatic life (CCME, 2003).

Table 2. HMs distribution in slices of sediment cores in the clam plains

| Name         | Cr   | Zn   | As   | Cd   | Hg   | Pb   |
|--------------|------|------|------|------|------|------|
| TT-P8-1      | 70.65| 448.76| 15.24| 2.15 | 40.63| 62.93|
| TT-P8-2      | 62.39| 620.40| 35.99| 5.95 | 34.87| 77.18|
| TT-P8-3      | 144.75| 958.71| 93.71| 2.00 | 17.21| 103.76|
| TT-P9-1      | 57.55| 463.08| 17.64| 4.74 | 53.03| 117.29|
| TT-P9-2      | 66.10| 439.20| 13.27| 3.05 | 120.37| 61.45|
| TT-P9-3      | 116.83| 305.79| 39.12| 3.56 | 71.06| 93.52|
| TT-P10-1     | 67.52| 374.75| 13.70| 1.80 | 86.76| 60.74|
| TT-P10-2     | 79.20| 534.62| 29.26| 2.52 | 61.49| 83.83|
| TT-P10-3     | 87.91| 568.31| 19.46| 3.18 | 40.07| 47.37|
| TT-P11-1     | 62.30| 381.22| 16.70| 4.25 | 42.79| 47.67|
| TT-P11-2     | 63.23| 423.68| 27.26| 2.79 | 36.82| 43.22|
| QCVN 43:2012 BTNMT | 160.00| 271.00| 41.60| 4.20 | 0.70 | 112.00|
We have seen that contents of Cr, As, Cd and Pb were under the standard QCVN 43:2012/ BTNMT, except at the depth 40 - 60 cm at TT-P8 where As concentration was exceeded 2.25 times. The concentrations of Zn were also found 1.12 - 3.53 times higher than the standard. They increased with depth at TT-P8, TT-P10, TT-P-11 and decreased with depth at TT-P9. Another HM, Hg, its concentration in the samples was 24.58 - 171.96 times higher than the standard and decreased with depth at TT-P8, TT-P10 and TT-P-11. The increase or decrease in Zn and Hg concentrations may be influenced by the activities of clams living in sediment.

### 3.2 HMs in Selective Sequential Extraction

#### 3.2.1 Cobalt (Co)

The Co concentrations in the fractions at 4 sample stations are shown in Table 3.

| Table 3. Co concentrations in the chemical fractions (per cent) |
|---------------------------------------------------------------|
| Sample          | TT-P8-1 | TT-P8-2 | TT-P8-3 | TT-P9-1 | TT-P9-2 | TT-P9-3 | TT-P10-1 | TT-P10-2 | TT-P10-3 | TT-P11-1 | TT-P11-2 |
| F1              | BDL     | BDL     | BDL     | 0.01    | 0.03    | BDL     | BDL      | BDL      | BDL      | BDL      | BDL      |
| F2              | 0.02    | 0.02    | 0.05    | 0.03    | 0.01    | 0.05    | 0.02     | 0.02     | 0.02     | 0.05     | 0.02     |
| F3              | 3.17    | 1.44    | 0.79    | 5.79    | 5.86    | 2.73    | 5.91     | 3.96     | 1.75     | 3.59     | 2.47     |
| F4              | 0.08    | 0.05    | 0.03    | 0.16    | 0.16    | 0.10    | 0.19     | 0.12     | 0.06     | 0.14     | 0.08     |
| F5              | 2.84    | 1.55    | 0.81    | 3.88    | 3.81    | 2.33    | 4.01     | 3.19     | 1.42     | 3.41     | 2.37     |
| F6              | 0.61    | 0.61    | 0.35    | 0.86    | 0.58    | 0.65    | 0.79     | 0.78     | 0.41     | 0.75     | 0.69     |
| F7              | 87.4    | 91.87   | 91.63   | 83.09   | 83.16   | 89.65   | 82.56    | 87.04    | 91.19    | 85.55    | 88.65    |
| F8              | 5.88    | 4.45    | 6.39    | 6.17    | 6.39    | 4.54    | 6.49     | 4.89     | 5.15     | 6.52     | 5.72     |

The average distribution of Co in different chemical fractions by SSE procedure shows the following distribution (Table 3): F1 < F2 < F4 < F6 < F5 < F3 < F8 < F7. The highest concentration was found in F7 (oxidizable fraction). The data more than 82% (82.5 - 91.9%) shows an important role of organic matters in the oxidizable condition at the surficial sediment layer. At this concentration, Co was a potential resource for the coastal estuary environment.

We found that the Co bound to carbonate (F3) decrease by the depth. The highest Co concentration/amount in the sediment layer 0 - 20 cm may due to the presence of clams. The Co amount accumulated in the carbonate shells after decomposition will be eventually released back into the environment. This Co amount has also the most ability to easily exchange with the environment and penetrate into the ecosystem.

In the F5 fraction, the Co concentration decreases also by the depth. This is in agreement with the results of Klinkhammer (1980) and Kumar and Edward (2009). The distribution of Co depends on the reduction-oxidation in the iron oxide phase. But in F7, the Co concentration increases by depth, which is related to the reduction of oxygen content by the depth of the sediment layers.

In the residue (F8), the distribution of Co is not clearly different at various depths. The amount of metal in this residue is difficult to separate from sediment, so the ability to enter the ecosystem is very low.

#### 3.2.2 Zinc (Zn)

The Zn extractant in the fraction in Table 4 shows that the main distribution is in F7 and F8 with a rate of more than 97% of the total Zn content of all fractions. Accordingly, the distribution of Zn in the fractions increase as following: F1 < F2 < F4 < F3 < F6 < F5 < F8 < F7.

As we have known, Zn from the external environment enters the estuary area by some sources including wastewater (Boxall et al., 2000). The high content of Zn-containing organic matter present in this wastewater would lead to an increase in Zn concentration in the sediment. In addition, the Zn content present in the solid residue also indicates the level of Zn contamination in the river system: the greater the percentage of HMs present in the solid residue, the lesser the pollution in the environment because this solid residue involves components that cannot be remobilized.

In the fraction F7, the Zn distribution increases by the depth at position TT-P8, TT-P10 and TT-P11, and decrease at the position TT-P9 where clams were harvested over two months before sampling date.
### 3.2.3 Arsenic (As)

Table 5 shows the arsenic distribution in fractions following the depth at the sample position. The increasing arsenic distribution in fractions is F2 < F1 < F4 < F6 < F5 < F7. The As concentrations in F8 are varying at different depths.

#### Table 5. As concentrations in the chemical fractions (per cent)

| Sample name | TT-P8-1 | TT-P8-2 | TT-P8-3 | TT-P9-1 | TT-P9-2 | TT-P9-3 | TT-P10-1 | TT-P10-2 | TT-P10-3 | TT-P11-1 | TT-P11-2 |
|-------------|---------|---------|---------|---------|---------|---------|---------|---------|---------|---------|---------|
| F1          | 0.03    | 0.01    | 0.02    | 0.01    | 0.00    | 0.00    | 0.00    | 0.02    | 0.02    | 0.02    | 0.01    |
| F2          | 0.01    | 0.01    | BDL     | 0.01    | 0.01    | 0.01    | 0.01    | 0.01    | 0.02    | BDL     | BDL     |
| F3          | 0.32    | 0.14    | 0.06    | 0.02    | 0.02    | 0.02    | 0.02    | 0.02    | 0.02    | 0.02    | 0.02    |
| F4          | 0.07    | 0.03    | BDL     | 0.02    | 0.02    | 0.02    | 0.02    | 0.02    | 0.02    | BDL     | BDL     |
| F5          | 7.36    | 2.53    | 0.74    | 0.05    | 0.05    | 0.05    | 0.05    | 0.05    | 0.05    | 0.05    | 0.05    |
| F6          | 1.23    | 0.58    | 0.37    | 1.04    | 1.04    | 1.04    | 1.04    | 1.04    | 1.04    | 1.04    | 1.04    |
| F7          | 90.97   | 89.37   | 84.51   | 88.29   | 88.56   | 94.61   | 85.60   | 93.54   | 86.93   | 90.39   | 96.24   |
| F8          | BDL     | 7.33    | 14.28   | 2.72    | BDL     | 2.53    | BDL     | 8.01    | BDL     | BDL     | BDL     |

Note: TT-PX-1: depth 0 - 20 cm, TT-PX-2: depth 20 - 40 cm, TT-PX-3: depth 40 - 60 cm; BDL: below detection limit

As shown in Table 5 for the arsenic content of F5 and F6, As also formed a relatively stable affinity for iron oxides. In particular, the association with amorphous Fe ranges from 0.74 to 12.2% depending on location and depth of sampling. Bounding with crystalline Fe only accounts for a very small proportion, 0.37 - 1.51%. In F5, As tends to decrease by the depth at TT-P8, TT-P10 and TT-P11 positions, especially drastically reduced between the first two layers. At TT-P9, however, the As content was unstable, increasing in the second layer and decreasing sharply in the third layer. Similarly, the As content in F6 also tends to decrease with depth at TT-P8, TT-P10, and TT-P11 positions, and at TT-P9, As increases in the second layer then decreases rapidly in the third layer. Particularly in F8, at depth 0 - 20 cm at position TT-P8 and in the 20 - 40 cm depth layer at position TT-P9, at 0 - 20 cm and 20 - 40 cm layers of TT-P10 and 0 - 20 cm and 20 - 40 cm layers at position TT-P11 we do not see the presence of As.

### 3.2.4 Cadmium (Cd)

The Cd distribution in the fractions is F1 < F4 < F2 < F6 = F5 = F3 < F8 < F7 (Table 6). Cd is mainly present in fraction F7, more than 93.6% (93.6 - 98.6%) of total Cd content in sediment. This demonstrates the close association of Cd with the organic compounds present in the sediment. However, in F8 this content is only from 1.18 to 5.56%, the highest at the depth 0 - 20 cm at TT-P8 and TT-P10 positions. This amount of Cd is very difficult to separate from sediment in order to get released into the surrounding environment. With F7, Cd increases with depth at TT-P8 and TT-P10 positions, gradually decreases at TT-P9 but is almost unchanged at TT-P11. But in F8, the Cd decreases at TT-P8 and TT-P10, increases at a depth of 20 - 40 cm at TT-P9 and is almost unchanged at TT-P11.
3.2.5 Mercury (Hg)

As it can be observed in Table 7, there is no presence of Hg in the exchangeable fraction F2 and the Mn oxides F4.

The distribution of Hg increases in the fractions as following: F1<F6 <F3 <F5 <F8 <F7. Like other HMs, Hg is predominantly present in the fraction F7, which accounts for more than 70% of the total. The amount of Hg in F8 also accounts for a relative proportion of 10.6 - 28.4%.

### Table 7. Hg concentrations in the chemical fractions (percent)

| Sample name | TT-P8-1 | TT-P8-2 | TT-P8-3 | TT-P9-1 | TT-P9-2 | TT-P9-3 | TT-P10-1 | TT-P10-2 | TT-P10-3 | TT-P11-1 | TT-P11-2 |
|-------------|---------|---------|---------|---------|---------|---------|---------|---------|---------|---------|---------|
| F1          | BDL     | BDL     | BDL     | BDL     | BDL     | BDL     | BDL     | 0.02    | 0.01    | 0.01    | 0.01    |
| F2          | 0.02    | 0.01    | 0.01    | 0.04    | 0.02    | 0.02    | 0.02    | 0.05    | 0.03    | 0.02    | 0.04    |
| F3          | 0.20    | 0.11    | 0.10    | 0.12    | 0.16    | 0.30    | 0.39    | 0.30    | 0.21    | 0.13    | 0.27    |
| F4          | 0.02    | 0.01    | 0.01    | 0.01    | 0.01    | 0.03    | 0.01    | 0.02    | 0.01    | 0.01    | BDL     |
| F5          | 0.17    | 0.06    | 0.05    | 0.07    | 0.27    | 0.11    | 0.23    | 0.15    | 0.15    | 0.10    | 0.19    |
| F6          | 0.17    | 0.06    | 0.03    | 0.07    | 0.10    | 0.12    | 0.19    | 0.17    | 0.15    | 0.07    | 0.18    |
| F7          | 95.24   | 97.57   | 98.64   | 98.44   | 97.13   | 97.45   | 93.57   | 96.91   | 97.87   | 98.49   | 98.23   |
| F8          | 4.18    | 2.18    | 1.17    | 1.26    | 2.30    | 1.96    | 5.56    | 2.38    | 1.57    | 1.17    | 1.07    |

Note: TT-PX-1: depth 0 - 20 cm, TT-PX-2: depth 20 - 40 cm, TT-PX-3: depth 40 - 60 cm; BDL: below detection limit

3.2.6 Lead (Pb)

According to data shown in Table 8, Pb is completely absent in the water-soluble fraction F1 and fraction F2 and present with a negligible content in association with Mn oxides (F4). The distribution of Pb in fractions is as follows: F3 <F6 = F5 <F8 <F7. Similar to other metals, Pb is also found primarily in the oxidative fraction F7. In F8, Pb accounted for 8.61 - 20.0%. At depths with F7, Pb increases at 20 - 40 cm layer at TT-P8 and TT-P10, while it decreases at the same layer at TT-P11 and TT-P9.

### Table 8. Pb concentrations in the chemical fractions (percent)

| Sample name | TT-P8-1 | TT-P8-2 | TT-P8-3 | TT-P9-1 | TT-P9-2 | TT-P9-3 | TT-P10-1 | TT-P10-2 | TT-P10-3 | TT-P11-1 | TT-P11-2 |
|-------------|---------|---------|---------|---------|---------|---------|---------|---------|---------|---------|---------|
| F1          | BDL     | BDL     | BDL     | BDL     | BDL     | BDL     | BDL     | BDL     | BDL     | BDL     | BDL     |
| F2          | BDL     | BDL     | BDL     | BDL     | BDL     | BDL     | BDL     | BDL     | BDL     | BDL     | BDL     |
| F3          | 0.33    | 0.37    | 0.49    | 0.15    | 0.28    | 0.79    | 0.23    | 0.49    | 0.88    | 0.13    | 0.71    |
| F4          | BDL     | 0.01    | BDL     | BDL     | 0.01    | BDL     | BDL     | BDL     | 0.01    | 0.01    | 0.01    |
| F5          | 1.06    | 0.73    | 0.46    | 0.64    | 1.18    | 0.48    | 0.94    | 0.56    | 0.89    | 1.35    | 1.18    |
| F6          | 0.98    | 0.79    | 0.86    | 0.67    | 0.93    | 1.34    | 1.30    | 0.91    | 1.21    | 1.28    | 1.20    |
| F7          | 83.14   | 86.68   | 83.10   | 89.93   | 82.50   | 88.13   | 82.74   | 88.68   | 77.03   | 83.14   | 81.24   |
| F8          | 14.48   | 11.43   | 15.08   | 8.61    | 15.10   | 9.25    | 14.78   | 9.36    | 20.03   | 14.1    | 15.66   |

Note: TT-PX-1: depth 0 - 20 cm, TT-PX-2: depth 20 - 40 cm, TT-PX-3: depth 40 - 60 cm; BDL: below detection limit

3.2.7 Chromium (Cr)

Cr concentration is distributed in the fractions of the sediment sample in the following order: F1 <F3 <F5 <F6 <F8 <F7 and is completely absent in phase F2 and F4 with all samples analyzed. More than 62% of Cr is concentrated in fraction F7. This is the amount of Cr in the sediment sample that is likely to be involved in oxidation reactions. Cr content in F8 with a high ratio of 20.9 - 38.4% is hardly separable from sediment materials to participate in the environment. The rate of Cr present in the fraction F5 and
F6 is quite low, accounting for less than 2% of the total Cr content in sediment samples.

Table 9. Cr concentrations in the chemical fractions (percent)

| Sample name | TT-P8-1 | TT-P8-2 | TT-P8-3 | TT-P9-1 | TT-P9-2 | TT-P9-3 | TT-P10-1 | TT-P10-2 | TT-P10-3 | TT-P11-1 | TT-P11-2 |
|-------------|---------|---------|---------|---------|---------|---------|---------|---------|---------|---------|---------|
| F1          | BDL     | BDL     | BDL     | BDL     | BDL     | BDL     | BDL     | BDL     | BDL     | BDL     | BDL     |
| F2          | BDL     | BDL     | BDL     | BDL     | BDL     | BDL     | BDL     | BDL     | BDL     | BDL     | BDL     |
| F3          | 0.59    | 0.56    | 0.29    | 0.76    | 0.63    | 0.39    | 0.62    | 0.65    | 0.51    | 0.74    | 0.72    |
| F4          | BDL     | BDL     | BDL     | BDL     | BDL     | BDL     | BDL     | BDL     | BDL     | BDL     | BDL     |
| F5          | 1.40    | 1.62    | 0.74    | 1.90    | 1.57    | 0.96    | 1.55    | 1.23    | 1.25    | 1.57    | 1.48    |
| F6          | 1.43    | 1.77    | 1.17    | 1.94    | 1.62    | 1.06    | 1.64    | 1.46    | 1.36    | 1.694   | 1.77    |
| F7          | 58.09   | 62.98   | 64.16   | 68.21   | 70.01   | 62.05   | 63.46   | 75.78   | 69.15   | 71.32   | 75.11   |
| F8          | 38.49   | 33.07   | 33.64   | 27.19   | 26.16   | 35.53   | 32.72   | 20.88   | 27.73   | 24.67   | 20.92   |

Note: TT-PX: depth 0 - 20 cm, TT-PX: depth 20 - 40 cm, TT-PX: depth 40 - 60 cm; BDL: below detection limit

4. Conclusions

The concentration of total HMs in sediment cores in comparison with the standard QCVN 43:2012/BTNMT shows a pollution with Zn and Hg whose concentrations are 1.12 - 3.53 times and 24.6 - 171 times higher than the standard. The average distribution pattern of HMs in different SSE fractions increases in the order of F1 < F2 < F4 < F3 < F6 < F5 < F8 < F7. The highest concentration of heavy metals was found in the oxidable and residual phases. The data more than 60% shows that most of HMs seem to be available for the biological activities in the oxidable condition at the surficial sediment layer. The heavy metal concentrations depending on the depth of sediment show that clam's digestive activity or the decomposition of tissue and shell of clams possibly affects the content of heavy metals.

Acknowledgments. I would like to express my sincere thanks to The National Foundation for Science and Technology Development (NAFOSTED), affiliated with the Ministry of Science and Technology (MOST) for their precious financial support; without this, my research (project code: 1059.99-2010.17) cannot be completed. My special thanks are devoted to Dr. Nguyen Van Thuyen from the University of Natural Sciences, Vietnam for his endless support in the field trip. Last but not least, I would like to thank my colleagues working at the Institute of Geological Science (IGS) of the Vietnam Academy of Science and Technology (VAST), who often helped and gave me valuable support at critical junctures in terms of experiments during the making to this research.

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