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PIXE Analysis on Desorption of Elements from Coastal Sediments

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Coastal sedimental elements were desorbed with ultrapure water, artificial seawater, 0.2M (NH₄)₂C₂O₄, 1M CH₃COONH₄, 0.01M EDTA • 2Na, 1N HCl and 6N HCl, independently. Fluid volume to sediment weight ratio was 28ml/g for batch systems. Extraction by 10 minutes stirring per day were done for 8 to 16 days. 56 μ 1 of extracted fluid was dropped on Kimfoil to make a target. Beam bombardment was done with a cyclotron using a 4.5 MeV proton energy. For a typical sample, the maximum concentration of Fe seen in the samples was obtained by extraction with 6NHC1 (7.6mg/g). In comparison, the concentration of Fe observed after extraction with ultrapure water, artificial seawater, and 1M ammonium acetate were essentially zero. However, the amounts seen with 0.01M EDTA • 2Na and 0.2M ammonium oxalate were high as 18% and 9.5% of the 6N HCl value, respectively. About the extraction with EDTA having most excellent characteristics of chelate agent, alkali earth elements, Mg²⁺ and Ca²⁺ were detected in almost all samples because they form ionic bonding chelates. Desorption rates of transition metal elements which form coordinate bonding chelates were observed to be proportional to the stability constant except for Mn^{2+} and Cu^{2+} anomalies.

1. Introduction

At present it is internationally expected for Japan to achieve some responsibility and to play an important role for protection of the environment with a global scale. We have to respond to the international expection by displaying leadership in environmental problems, such as the application of charged particle beams to elemental analysis of environmental pollutants. The environments in the neighboring seas and the coasts around our country have seriously become worse, even though attempts are being made to recover the natural marine environment by improving the clarification capacity with an artificial seashore, partially constructed. It is important to find the adsorption-desorption kinetics of elements in coastal sediments, as a means to estimate the behavior of pollutants,

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especially the heavy metal elements. For the purpose of the assessment, it may be effective to make clear the behaviors of elements with coastal sediments by the use of the several kinds of the chemical extracting solutions.

This paper describes the result of attempts to obtain information on the adsorption/desorption kinetics of elements in coastal sediments. The extraction method uses independently the following seven kinds of extracting solution; ultrapure water, artificial seawater, 1M ammonium acetate, 0.2M ammonium oxalate, 0.01M EDTA • 2Na aqueous solutions, and 1N and 6N hydrochloric acids adopted by Fukui et al.¹²

It is very important to analyze the cause of turbidity and to discover the origin of the adsorbed matters^{3,0} on the coastal sediments for learning the adsorption/desorption characteristics of the elements^{1,5,6}.

The reason why the extraction procedures were attempted as the pretreatment for PIXE analysis will hereinafter be described. It may be expected that ultrapure water and artificial seawater desorb the seawater-soluble matters adsorbed on the surfaces or porous porosities of coastal sediments. Fukui²) reported that ammonium ions in aqueous solution could more readily desorb trace metals adsorbed by the ion-exchange mechanism than by the sodium ion abundant in seawater. In these studies, (1) both ammonium salts of acetate (1M) and oxalate (0.2M) were used in order to obtain information on desorption kinetics under readily exchangeable and suboxic conditions, respectively; (2) EDTA \cdot 2Na (0.01M) was used for examining the possible formation of chelate compounds with the heavy metals; and (3) HCl (1N and 6N) was used for investigating metal desorption destroying a part of mineral lattices in the solution.

Why PIXE? PIXE analysis has the following three outstanding characteristics for the assessment of the environmental pollution. (1) Multielements can be determined simultaneously without destroying the samples; (2) it needs only trace amounts; and (3) a valuable sample, involving great time, effort and expense in chemical extraction, can be analyzed by repeated procedure at various sensitivities. In brief, chemical extraction plus PIXE is predicted to be very useful for work in the marine environment.

2. Experimental

2.1 Sample and pretreatment

At five points in the Osaka Bay Shore, near South of Sensyu, Osaka Prefecture (Fig. 1), ten samples of deposits from the surface mud layer and the 20cm deep layer were collected³⁰ and used in this experiment. The samples were dehydrated by centrifuging (≤ 5800 rpm, 10min) for three days after the samples were collected, then dried for 64 hours in a thermostat kept at 75°C. The samples thus obtained were sieved into four stages with 16, 24 and 32 mesh screens as shown in Table 1. External characteristics of samples after dehydration, dry-



Fig. 1 Schematic map of the sampling stations

| Stag | e Mesh | Diameter (mm) | Diameter of sieving (μm) |
|------|-----------------|---------------|-------------------------------|
| 1 | >Mesh 16 | d>0.590 | 1000 |
| 2 | Mesh 16~Mesh 24 | 0.590>d>0.350 | 710 |
| 3 | Mesh 24~Mesh 32 | 0.350>d>0.290 | 500 |
| 4 | Mesh 32 $>$ | 0.290>d | |

Table 1 Characteristics of analytical sieve fractions

Table 2 External characteristics of samples

| | Collecting sites | Geographical position | Characteristics |
|-------|------------------------------------|----------------------------------|-----------------------------------|
| St. 1 | Surface layer* 20cm deep layer* | Dredging near Sano river | Mud, sand Shoulder and sand |
| St. 2 | 2 Surface layer 20cm deep layer | Aperture | Many shoulders Pebble |
| St. 3 | 8 Surface layer 20cm deep layer | Dredging near Kashii river | Bottom sediment Mud, sand |
| St. 4 | l Surface layer 20cm deep layer | Tarui, Aperture (Sennan City) | Bottom sediment Mud |
| St. 5 | 5 Surface layer 20cm deep layer | Okadaura near kashii river | Bottom sediment, shoulder Sand |

* Surface layer: Collecting mud from surface to 2cm depth

★20cm deep layer: Collecting 2 ~ 3 cm bottom mud obtained by inserting a pipe to 23~24cm depth of bottom of the sea

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| | Dalla i Chaulden | Rate of sieve fraction (%) | | | | | | | | | |
|---------------------|-------------------|----------------------------|--------|--------|--------|--|--|--|--|--|--|
| Collecting sites | Peddle + Snoulder | Stg. 1 | Stg. 2 | Stg. 3 | Stg. 4 | | | | | | |
| St.1 Surface layer | 20. 58% | 24.03 | 8.57 | 13. 81 | 33.01 | | | | | | |
| 20cm deep layer | 25.05% | 28.45 | 6.97 | 14.29 | 25.23 | | | | | | |
| St.2 Surface layer | 37.90% | 28.28 | 6.44 | 8.38 | 18.99 | | | | | | |
| 20cm deep layer | 25. 51% | 38.50 | 5.65 | 8.94 | 21.44 | | | | | | |
| St.3 Surface layer | 0% | 0.82 | 1.01 | 3.00 | 95.17 | | | | | | |
| 20cm deep layer | 0% | 11.18 | 3.75 | 4.03 | 81.04 | | | | | | |
| St. 4 Surface layer | 5. 50% | 33.43 | 7.32 | 8.35 | 45.40 | | | | | | |
| 20cm deep layer | 17.98% | 24.48 | 9.60 | 9.84 | 38.10 | | | | | | |
| St.5 Surface layer | 0% | 6.87 | 1.75 | 5.00 | 86.38 | | | | | | |
| 20cm deep layer | 0% | 0.23 | 0.60 | 4.76 | 94. 41 | | | | | | |

Table 3 Particle size distribution of deposition

ing and sieving are shown in Table 2. Table 3 represents a distribution of the grain size of deposits, indicating the distribution of the whole containing the pebbles and the shells which have been removed before sieving.

2.2 Chemical extraction procedures

The stage 4 sample was most finely sieved. Each sample consisting of 1g at this stage was used in this extraction procedure. In the desorption test of the surface mud layer of the station 2 and 4, 0.5g of each sample was ground to powder in an agate mortar and was used for the extractions. Table 4 represents the constitution of extracting solution. A flow chart of the desorption test is shown in Fig. 2. To each sample of 1g or 0.5g, extracting solutions of 27.85 or 13.9ml were added in volumetric flasks of 100 or 50ml, respectively. In the desorption tests using ultrapure water and artificial seawater as extracting solution, samples were shaken occasionally for 10min per day during 15 and 16 days, respectively. In the desorption tests using 1M ammonium acetate, 0.2M ammonium oxalate, 0.01M EDTA • 2Na, and 1N and 6N hydrochloric acids, samples were extracted during eight and four days, respectively with the same shaking condition as that described above. It was considered that the extraction condition was pearly the same, although a shaking period differed somewhat with the kind of extracting solution. The liquids obtained by extraction with occasional shaking were filtered with a membrane filter (TM-2, pore size $0.45 \,\mu$ m, cellulose nitrate 47mm diameter) by the use of a Witt type filtering flask to which a stainless filter support screen was attached, in vacuo. The extracts from ultrapure water, artificial seawater, 1M ammonium acetate and 1N hydrochloric acid were filtered by the suction method mentioned above. In order to prevent the corrosi on by strong acid, the filtrations of the extracts from 0.2M (NH₄)₂C₂O₄, 0.01M EDTA • 2Na and 6N HCl were carried out by the use of a polycarbonate support screen instead of the stainless support screen. Each filtration time of 27ml of extract was about 3min, devided into two protions, where it took about 30s per filtration. Analytical data of the extracts from 1M ammonium acetate and 1N ydrochloric acid which were first filtered must be checked against the contamination arising from the stainless support mesh screen, which is readily corroded by acids.

| Extracting solution | Preparation method |
|---|---|
| Ultrapure water Artificial seawater 1M Ammonium acetate | Distilled water → ion exchange column → Millipore filter *Pure water → Lyman and Fleming method *Pure water → 1M CH ₃ COONH ₄ dissolved |
| 0.2M Ammonium oxalate 0.01M EDTA • 2Na 1N Hydrochloric acid 6N Hydrochloric acid | *Pure water → 0.2M (NH₄)₂C₂O₄ • H₂O dissolved *Pure water → 0.01M C₁₀H₁₄N₂O₈Na₂ • 2H₂O dissolved *Pure water → Diluted Ultrapure HCl (40%) to 3.59% Ultrapure water → Diluted ultrapure HCl (40%) to 19.93% |

Table 4 Preparation method of extracting solution

*Pure water was obtained by treating distilled water through ion exchange

Pretrentment of coastal sediment samples for desorption test Coastal mediment sampling: each 1g, grounded sample: 0.5g (Each surface layer or 20cm deep layer sample: total 10 samples

DExtraction with @Extraction with @Extraction ction DExtraction with 0.01W EDTA • 2Na GExtraction with with DExtraction with ultrapure wate artificial : IN NEL AC 0.2N (NEL),0x IN HCL **GN HC1** 1 1 1 I I 65 -oles mles Shaking for 10min king for 10mi Shaking for 10min for 10mi 10min 100 fo king for 1 Omi per day during da per day during 8 days per day during 15 days 16 ő days days 75 8 days dava 1 1 um filtratio The left way as the The with filtering fla left left left left left Filter: TH-2 stainle 5' M polycas ate stainless mesh polycari nate stainless eesh polycarbonate support scree moort scree -L 1 I ł I d 56 µl on the The The 6av The . left The left The : left The left Kimfoil membrane left left dried w nier infra .! ţ 1 20 samples pies 28 = oles 20 28 for target for target for target for target

Fig. 2 Pretreatment of coastal sediment samples for desorption test

2.3 PIXE analysis

PIXE target samples were analyzed by the PIXE analysis system in the cyclotron room at the Faculty of Science, Osaka University. A block diagram of the PIXE analysis system is shown in Fig. 3. Beam bombardment was done using 5MeV protons of the cyclotron, degraded to about 4.5MeV with a Kapton foil beam smoother $(2.5mg/cm^2)$. The beam cross section diameter at the target

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position was defined to 8mm by double graphite collimators. Beam current was about 1nA and cumulative charge was about 600nC. Characteristic X-rays from samples were measured with a Si(Li) detector (effective area: 28.3mm², FWHM: 160eV at 5.9keV) with a pierced Lumilar "funny" filter in front of the Be window. The angle between the sample plane and the beam direction was 45°. The distance between the sample plane and the Be window of Si(Li) detector was 5cm, and the angle between the detector head and the beam direction was 135°.



Fig. 3 Block diagram of PIXE+FAST system



Fig. 4 Sensitivity curves of the PIXE system

As references, standard samples for PIXE of Micromatter Co. were used. Sensitivity curves of the system are shown in Fig. 4. Spectrum analysis was done by using a personal computer based commercial MCA software, and the quantitative determination of elements was undertaken by calculating X-ray self absorption corrections.

2.4 NAA analysis

Part of the samples made as described in 2.2, two sets of samples of St.1 surface mud layer and 20cm deep layer were assayed by NAA to assure analytical results and increase the numbers of determined elements. Each sample portion on target Kimfoil membranes was scissored out and was seeled into a small These bags were put into a polypropylene capsule together polvethylene bag. with NBS standard river sediment and bovine lever references. This capsule was neutron irradiated in the F ring (thermal neutron flux: 1.4×10^{12} n · cm⁻² · sec⁻¹) of the reactor of Rikkyo University for 6 hours. A gamma-ray measurement was done tow times, for 4,000 and 40,000s with the γ -ray emmitting nuclide analysis system consisted of a shield box (10cm thick lead, 5mm thick copper lining, 5mm thick opaque acrylic lining, volume: 320×320×420mm³) and a coaxial ntype pure Ge detector (relative efficiency: 30%) of Research Institute for Advanced Science and Technology, University of Osaka Prefecture. Neutron flux was confirmed with an NBS river sediment standard sample, and found to be equal to the nominal value. Spectral analysis was done by using a personal computer based commercial MCA software for nuclide analysis.

3. Analytical results and discussion

The PIXE analytical results of St.1 surface and 20cm deep layer samples are listed in Table 5 as a typical example for main elements, together with the results of extracting liquid blanks. 21 elements, Mg, Al, Si, P, S, Cl, K, Ca, Sc, Ti, V, Cr, Mn, Fe, Ni, Cu, Zn, Br, Sr, Hg and Pb were determined. Seven elements, Si, S, Cl, K, Ca, Fe and Zn, were determined for all kinds of extracted samples. The NAA results corresponding to them are listed in Table 6. From the results, the following eight facts were found.

1) Hydroxides of Ca, Mg, and K were released from silicates in sediments by hydrolysis⁹. They were extracted with ultrapure water in the order, Ca>K>Mg (Ca:47-726 μ g/g, K:17-134, Mg:not detected), and with artificial seawater in the order, Mg>Ca>K (Mg:350-2090 μ g/g, Ca: 94-1087, K:4-540).

2) By 0.2M $(NH_4)_2Ox$, K⁺ was extracted much $(K:99-417 \mu g/g)$ in the form of $K_2C_2O_4$, while Ca^{2+} was scarcely detected because it precipitated as CaC_2O_4 . S was extracted fairly much $(S:142-2360 \mu g/g)$ in the forms of SO_4^{2-}/S^{2-} through some reactions with CaSO₄ and FeS. Ti⁴⁺ ion was extracted much $(Ti:205-342 \mu g/g)$ in the form of $[Tio(C_2O_4)_2(H_2O)]^{2-}$. V⁵⁺ is desorbed as VO_2^+ by the exchange with

Table 5-1 Analytical results of PIXE

Content (unit: $\mu g/g$ sediment or $\mu g/27.85$ ml extracting blank)

| Sample | | | | | | | | | E | lemer | its | | | | | | | | | | |
|---|------|-----|------|-----|------|-------|-------|------|------|-------|-----|-----|------|------|------|-----|----|-----|----|-----|------|
| | Mg | Al | Si | Р | S | C1 | K | Ca | Ti | V | Cr | Mn | Fe | Ni | Cu | Zn | Hg | As | Pb | Br | Sr |
| ultrapure water blank | NÐ | ND | 405 | ND | ND | 27.5 | ND | 21.5 | 310 | ND | ND | ND | 22 | ND | ND | 17 | ND | ND | ND | ND | ND |
| artificial seawater blank | ND | ND | 333 | ND | ND | 318 | ND | 32.5 | 460 | ND | ND | ND | 26 | ND | ND | 11 | NÐ | ND | ND | ND | ND |
| 0.2M (NH₄)₂Ox blank | ND | ND | 633 | ND | ND | ND | 2.5 | 34.5 | 468 | 1.5 | ND | ND | 23 | ND | ND | 9.3 | ND | ND | ND | ND | ND |
| 1M NH₄Ac blank | ND | ND | 343 | ND | ND | ND | 3.5 | 18.8 | 222 | ND | ND | ND | 19 | ND | ND | 7.3 | ND | ND | ND | ND | ND |
| 0.01M EDTA · 2Na blank | ND | ND | 300 | ND | ND | 39.3 | 5.0 | 9.75 | 173 | 14 | ND | ND | 17 | ND | ND | 5.3 | ND | ND | ND | ND | ND |
| IN HCI blank | NÐ | ND | 633 | ND | ND | 201 | 33. 8 | 32.8 | 152 | 16 | ND | ND | 21 | ND | ND | 18 | ND | 1.3 | ND | 82 | ND |
| 6N HC1 blank | ND | ND | 64.8 | ND | ND | 7.0 | 7.25 | 67.8 | 131 | 8.5 | ND | ND | 17 | ND | ND | 17 | ND | ND | ND | 500 | ND |
| St.1 sf ultrapure water | ND | ND | 107 | ND | 71 | 3400 | 110 | 400 | ND | ND | ND | ND | 23 | ND | ND | ND | ND | ND | ND | ND | ND |
| St.1 sf artificial seawater | 680 | ND | ND | ND | 81 | 2140 | 97.5 | 463 | 18.0 | ND | ND | 2.8 | 51 | 0.5 | ND | 1.5 | ND | ND | ND | 40 | ND |
| St.1 sf 0.2M (NH ₄) ₂ Ox | ND | ND | 550 | ND | 2360 | 2030 | 308 | 5.3 | 710 | 9.8 | 7 | 48 | 740 | ND | ND | 29 | ND | ND | ND | ND | ND |
| St.1 sf 1M NH ₄ Ac | 310 | ND | ND | ND | 88 | 2680 | 125 | 3230 | 11.8 | ND | ND | 29 | 16 | ND | ND | ND | ND | ND | ND | ND | ND |
| St. 1 sf 0.01M EDTA · 2Na | 29 | ND | 613 | ND | 343 | 2470 | 202 | 2600 | 715 | ND | ND | 79 | 1390 | ND | 8.5 | 48 | ND | ND | ND | ND | ND |
| St. 1 sf IN HCl | ND | 190 | 147 | ND | 465 | 10100 | 68.8 | 938 | ND | ND | ND | ND | 3850 | ND | ND | ND | ND | 2.5 | ND | ND | ND |
| St. 1 sf 6N HC1 | ND | 780 | 64 | 35 | 273 | 17000 | 475 | 1990 | 490 | ND | ND | 95 | 7600 | ND | ND | ND | 22 | ND | 79 | 320 | 13 |
| St.1 20cm ultrapure water | ND | ND | 137 | ND | 10 | 3080 | 28 | 283 | 16 | 8 | ND | ND | 19 | ND | ND | ND | NÐ | ND | ND | 17 | - ND |
| St.1 20cm artificial seawater | 350 | 22 | 31 | 1.3 | 33 | 953 | 13 | 127 | 16 | ND | ND | ND | 17 | 1.3 | . 25 | 1.0 | ND | ND | ND | 13 | NÐ |
| St. 1 20cm 0. 2M (NH ₄) ₂ 0x | 780 | NÐ | 483 | ND | 370 | 2500 | 293 | 57 | 673 | 55 | ND | 1.8 | 36 | . 25 | ND | 17 | ND | ND | ND | ND | ND |
| St.1 20cm 1M NH ₄ Ac | 150 | ND | 73 | ND | 273 | 2050 | 335 | 4530 | 6 | ND | ND | ND | 19 | 2.8 | ND | ND | ND | ND | ND | ND | 8.0 |
| St. 1 20cm 0.01M EDTA • 2Na | 110 | ND | 620 | ND | 26 | 1800 | 98 | 5600 | 630 | ND | ND | 16 | 595 | . 75 | 3.3 | 19 | ND | ND | ND | 7.5 | 25 |
| St.1 20cm 1N HC1 | ND | 103 | 173 | ND | 598 | 17800 | 390 | 3200 | 21 | ND | ND | 38 | 5730 | ND | ND | ND | ND | ND | ND | ND | ND |
| St. 1 20cm 6N HC1 | 1750 | 308 | 148 | ND | 400 | 18900 | 388 | 3800 | 558 | ND | ND | 61 | 6950 | ND | 1.0 | 49 | 60 | ND | ND | 343 | 33 |
| | Mg | Al | Si | Р | S | Cl | K | Ca | Ti | V | Cr | Mn | Fe | Ni | Cu | Zn | Hg | As | Pb | Br | Sr |

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Table 5-2 Analytical results of PIXE

Content (unit: μ g/g sediment or μ g/27.85ml extracting blank)

| Sample | Elements | | | | | | | | | | | | | | | | | | | | |
|---|---|---------------------------------------|--|--|---|--|---|--|---|---|--|---|---|------------------------------------|---|---|---|--|-----------------------------------|---|---|
| | Mg Al Si P S Cl K Ca Ti V Cr Mn Fe Ni Cu Zn Hg As Pb Br S | | | | | | | | | | | | | | Sr | | | | | | |
| St. 2 sf ultrapure water St. 2 sf artificial seawater St. 2 sf 0.2M (NH ₄) ₂ Ox St. 2 sf 1M NH ₄ Ac St. 2 sf 0.01M EDTA · 2Na St. 2 sf 1N HCl St. 2 sf 6N HCl | ND 568 14.8 470 60.5 ND 1930 | ND ND ND ND 249 420 | 115 ND 403 83 573 ND 135 | ND ND ND ND 88.8 ND | 433 2270 345 233 470 3 535 418 | 4330 9580 3230 3730 3900 10100 17800 | 70.3 200 420 94.5 348 43.3 60.3 | 748 315 49.3 5130 4530 1650 2630 | ND ND 685 21 658 ND 580 | ND ND ND ND ND ND ND | ND ND ND ND ND ND ND | ND ND 12.3 21 30.5 ND 19.8 | 6 1 683 ND 733 3180 5900 | ND ND 1.5 .25 ND ND | ND ND 1.3 2.0 ND ND | 1.8 ND 33 ,75 3.3 ND ND | ND ND ND ND ND ND 6.8 | ND ND ND ND ND ND | ND ND ND ND ND 16 | 18.5 ND 24.3 ND 19.8 ND 285 | ND ND ND 11 ND ND |
| St. 2 20cm ultrapure water St. 2 20cm artificial seawater St. 2 20cm 0. 2M $(NH_4)_2Ox$ St. 2 20cm 1M NH_4Ac St. 2 20cm 0.01M EDTA · 2Na St. 2 20cm 1N HC1 St. 2 20cm 6N HC1 | ND 1430 ND 153 ND ND 1490 | ND ND ND ND 168 265 | ND 538 73.5 448 14.3 166 | ND ND ND ND ND ND | 1970 3300 142 213 69.5 530 1850 | 2850 31800 1000 1930 1790 8500 13600 | 17.3 380 140 433 118 313 161 | 258 898 12.5 5100 5350 1830 2680 | ND ND 703 12.3 433 ND 525 | ND ND 34 2.5 ND ND ND | ND ND ND 2.5 ND ND | ND 2.5 ND 10 12.5 40 28.3 | 16.3 3.5 298 17.8 625 3550 5480 | ND ND ND ND ND ND | ND ND ND 2.0 ND ND | ND 1.0 22 ND 18 ND 20 | ND ND ND ND ND 94 | ND ND . 75 ND ND ND ND | ND ND ND 4.5 25 ND | ND 418 ND 3.3 1 ND 308 | ND 110 ND 24 ND 19 |
| | Mg | Al | Si | Р | S | Cl | K | Ca | Ti | V | Cr | Mn | Fe | Ni | Cu | Zn | Hg | As | Pb | Br . | Sr |
| St.3 sf ultrapure water St.3 sf artificial seawater St.3 sf 0.2M (NH ₄) ₂ Ox St.3 sf 1M NH ₄ Ac St.3 sf 0.01M EDTA · 2Na St.3 sf 1N HC1 St.3 sf 6N HC1 | ND 2090 588 200 229 ND ND | ND ND 143 ND 528 863 | 91 155 568 ND 520 75.5 142 | ND ND ND ND ND ND | 161 2730 328 247 4.0 184 73.3 | 3800 32500 2650 1680 1930 8350 7250 | 81. 8 540 300 247 179 93. 3 68. 0 | 101 1120 49.8 715 710 390 510 | ND ND 808 15 618 ND 793 | ND ND ND ND ND ND | ND ND ND ND ND ND ND | ND ND 7.3 11.3 23.8 68.8 33.0 | ND 758 19 1910 4080 4730 | ND ND ND 1.8 ND ND | ND ND ND ND ND ND 8.0 | ND ND 15 ND 24 ND 42 | ND ND ND ND ND ND | ND ND ND ND ND ND | ND ND ND ND 24 ND | ND 450 ND 7.5 6.8 ND 270 | ND 156 ND ND ND ND ND |
| St. 3 20cm ultrapure water St. 3 20cm artificial seawater St. 3 20cm 0. 2M $(NH_4)_2Ox$ St. 3 20cm 1M NH_4Ac St. 3 20cm 0. 01M EDTA · 2Na St. 3 20cm 1N HC1 St. 3 20cm 6N HC1 | ND 1230 515 170 410 305 2160 | ND ND ND 62.5 -232 325 | 78.3 5450 540 81.0 690 250 149 | ND ND ND ND ND 72 ND | 255 3380 222 146 128 273 300 | 1830 32800 2220 1690 2040 6950 13300 | 118 283 253 226 208 89.8 24 | 72 855 53.5 543 733 298 360 | ND ND 775 8.75 728 ND 583 | ND ND ND ND ND 9.8 ND | ND ND ND ND ND ND ND | ND ND 21.3 ND 26.5 61.5 42.3 | 15.3 13.5 142 19.5 2410 3300 5400 | ND ND ND ND ND ND | ND ND ND 4.5 ND ND | ND 21 7.3 41 ND 34 | ND ND ND ND ND ND | ND ND ND ND ND ND | ND ND ND ND ND A2 | ND ND 11 11 4.5 ND 298 | ND ND 8.5 5.5 ND ND |
| | Mg | Al | Si | Р | S | Cl | K | Ca | Ti | V | Cr | Mn | Fe | Ni | Cu | Zn | Hg | As | Pb | Br | Sr |

PIXE Analysis on Desorption of elements from coastal sediments

Table 5-3 Analytical results of PIXE

Content (unit: $\mu g/g$ sediment or $\mu g/27.85$ ml extracting blank)

| Sample | | | | | | | · | | | Eleme | nts | | | | | | | | | | |
|--|---|---------------------------------------|--|--|--|--|--|--|---|--|--|--|---|---|------------------------------------|--|--|--|--|--|--|
| · · · | Mg | Al | Si | Р | S | C1 | K | Ca | Ťi | V | Сг | Mn | Fe | Ni | Cu | Zn | Hg | As | Pb | Br | Sr |
| St. 4 sf ultrapure water St. 4 sf artificial seawater St. 4 sf 0. 2M $(NH_4)_2 \Omega x$ St. 4 sf 1M NH_4Ac St. 4 sf 0. 01M EDTA \cdot 2Na St. 4 sf 1N HCl St. 4 sf 6N HCl | ND 618 515 338 152 575 ND | ND 68.8 ND ND 240 755 | 116 3 34 418 83 610 202 20 | ND ND ND ND ND ND | 120 358 303 465 99, 5 458 2530 | 3480 3150 3400 2680 2130 9880 18500 | 64 25.5 268 59 228 83 408 | 608 470 278 2800 2160 1140 1630 | ND ND 703 ND 620 ND 445 | ND ND ND ND ND ND | NÐ ND ND ND ND ND | ND ND ND 28.5 79.3 74.3 | 7.3 20.3 61.8 11.0 1540 3850 7330 | ND ND ND 2.3 ND ND | ND ND ND ND ND ND | ND 2.3 17 4.8 30 26 48 | ND ND ND 9.8 ND ND | ND ND ND ND ND S. 3 | ND ND ND 3.5 ND ND | ND ND 17.5 11 2.3 ND 360 | ND ND 5 ND 5 ND ND 14 ND |
| St. 4 20cm ultrapure water St. 4 20cm artificial seawater St. 4 20cm 0. 2M (NH ₄) ₂ Ox St. 4 20cm 1M NH ₄ Ac St. 4 20cm 0.01M EDTA • 2Na St. 4 20cm 1N HC1 St. 4 20cm 6N HC1 | NÐ ND 373 325 223 ND 1840 | ND 30 ND ND 338 236 | 158 27.3 525 56.5 643 320 179 | ND ND ND ND ND ND ND | 2880 413 318 253 111 980 222 | 4200 3200 2400 1990 1580 16800 14300 | 134 147 318 55 137 76.5 268 | 228 165 53.3 2800 2380 1720 1100 | 8.3 ND 738 3.0 658 52.8 610 | ND ND ND ND ND ND ND | ND ND ND ND ND ND | ND 25.3 ND 32.5 79.8 45.3 | 20 17.8 320 14.8 1580 6700 5080 | ND ND 1.3 ND 1.3 2.5 ND | ND ND ND ND ND ND | ND ND 15 ND ND 39 5.8 | ND ND ND ND ND ND | ND ND ND ND ND ND | ND ND ND ND ND ND | ND 45 ND 6. 5 76 320 | ND 14 ND ND 6.8 ND |
| | Mg | Al | Si | Р | S | C1 | K | Ca | Ti | V | Cr | Mn | Fe | Ni | Cu | Zn | Hg | As | Pb | Br | Sr |
| St.5 sf ultrapure water St.5 sf artificial seawater St.5 sf $0.2M (NH_4)_2 0x$ St.5 sf $1M NH_4 Ac$ St.5 sf $0.01M EDTA \cdot 2Na$ St.5 sf $1N HC1$ St.5 sf $6N HC1$ | ND 2040 710 236 393 ND 1780 | ND ND ND 72.3 325 1680 | 91 ND 485 47.8 590 73.5 117 | ND ND ND ND ND ND ND | 503 1570 275 121 12 410 473 | 3830 6930 2550 2440 2700 7700 9250 | 102 500 323 214 283 59.8 214 | 533 853 50. 5 893 1330 483 925 | ND ND 810 8.5 748 ND 558 | ND ND ND ND ND ND | ND ND ND ND ND ND | ND 5.3 ND 2.8 20.3 ND 13.5 | 40 6.75 174 16.3 2650 3430 5950 | ND 1.3 ND ND ND ND ND | ND ND ND 4.3 2.3 ND | NÐ ND 22 3. 0 27 23 43 | ND ND ND ND ND ND ND | ND ND ND ND ND ND | ND ND ND 9.5 ND ND | ND 149 17 13 6.5 ND 330 | ND ND ND ND ND ND |
| St. 5 20cm ultrapure water St. 5 20cm artificial seawater St. 5 20cm 0.2M $(NH_4)_20x$ St. 5 20cm 1M NH ₄ Ac St. 5 20cm 0.01M EDTA \cdot 2Na St. 5 20cm 1N HC1 St. 5 20cm 6N HC1 | ND ND 217 167 98.3 ND ND | ND ND ND 171 211 197 | 51.3 44.5 490 37.8 448 308 385 | ND ND ND ND ND ND | 338 18.0 184 145 368 57.3 498 | 2160 57.0 1400 1330 2230 5100 4400 | 96 4.0 102 95.3 2.0 109 66 | 68.8 31.8 20.5 132 280 109 140 | ND 15.5 695 10.3 715 22.3 855 | ND ND 43.8 ND ND ND ND ND | ND ND ND ND ND ND ND | ND ND 2.75 ND ND ND 7.8 | 27.5 16.8 570 14.3 1660 2110 2750 | ND ND ND ND ND ND ND | ND ND ND 4.5 ND 13 | ND ND 18 3.0 33 8.0 31 | ND ND ND ND ND ND | ND ND ND ND ND ND ND | NÐ ND ND ND ND ND ND | ND ND .5 4.0 11.3 48.8 218 | ND ND ND ND ND ND ND |
| | | Al | Si | Р | S | Cl | К | Ca | Ti | V | Cr | Mn | Fe | Ni | Cu | Zn | Hg | As | Pb | Br | Sr |

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Table 6 Analytical results of NAA

Content (unit: μ g/g sediment or μ g/27.85ml extracting blank)

| Sample | | | | | | | | | El | ement | S | | | | | | | | | | | |
|---|---|---|--|--|---|--|---|---|---|--|---|---|--------------------------------------|--|--|--|--|---|---|--|---|--|
| | Na | К | Sc | Cr | Mn | Fe | Co | Zn | Br | Ag | Cd | Sn | Cs | Ba | La | Sm | Eu | Lu | Hf | Ta | Pa | Yb |
| ultrapure water blank artificial seawater blank 0.2M (NH ₄) ₂ Ox blank 1M NH ₄ Ac blank 0.01M EDTA • 2Na blank 1N HCl blank 6N HCL blank | 39 140000 77 47 3400 64 50 | ND 1150 ND ND ND ND ND | .06 ND .01 ND ND ND | ND ND ND ND 6.6 ND | ND ND ND ND ND ND ND | ND ND ND ND ND ND | ND ND ND ND ND 42 ND | ND ND ND ND ND ND ND | 49 13200 34 40 8300 1300 8330 | ND ND ND ND ND ND ND | ND ND ND ND ND ND ND | NÐ NÐ ND ND ND ND | ND ND ND ND ND ND | ND ND ND ND ND ND ND | ND ND ND 5.5 ND ND | ND ND ND 2.9 ND ND | ND ND ND ND ND ND 24 | ND 120 ND ND ND ND 160 | ND ND ND ND ND ND | ND ND ND ND ND ND | ND ND ND ND ND ND | . 68 ND ND ND ND ND ND |
| St. 1 sf ultrapure water St. 1 sf artificial seawater St. 1 sf 0.2M (NH ₄) ₂ Ox St. 1 sf 1M NH ₄ Ac St. 1 sf 0.01M EDTA · 2Na St. 1 sf 1N HC1 St. 1 sf 6N HC1 | 3200 220000 5300 3400 11000 3400 3400 | ND 6000 ND ND ND 490 ND | ND ND .11 .01 ND .22 .33 | ND ND 1.1 1.9 22 21 12 | ND ND 4.7 ND 156 165 | ND ND 140 2870 10300 12100 | ND 11 1.7 1.4 2.9 5.7 7.0 | ND ND 17 57 91 3090 | 340 17100 356 323 326 1510 8300 | ND 6.2 ND .71 10 ND ND | ND ND 2.8 ND ND ND ND | ND 550 ND 360 300 ND ND | ND ND . 30 ND ND 1. 1 | ND 5990 ND 5650 164 ND ND | ND ND ND 1.4 4.5 5.5 | ND ND . 12 ND 1. 1 2. 4 2. 9 | ND ND . 02 . 13 . 27 . 22 | ND ND ND ND ND ND ND | ND 4.9 ND 1.2 ND ND 4.3 | ND 5.3 .84 ND 4.1 3.4 ND | ND ND ND . 52 ND . 35 | ND ND ND . 69 . 23 ND |
| St. 1 20cm ultrapure water St. 1 20cm artificial seawater St. 1 20cm 0.2M $(NH_4)_2Ox$ St. 1 20cm 1M NH_4Ac St. 1 20cm 0.01M EDTA \cdot 2Na St. 1 20cm 1N HC1 St. 1 20cm 6N HC1 | 2500 124000 2900 2600 9600 3200 2700 | ND 2900 ND ND ND ND ND | ND ND . 13 . 07 . 17 . 48 . 44 | 2.8 5.6 5.0 ND ND ND 7.8 | ND ND 5.1 ND ND 143 167 | ND ND 3400 ND 810 9600 11000 | 8.6 5.2 24 ND 1.2 7.9 8.1 | ND ND 25 19 32 110 84 | 260 20900 368 384 243 1580 8100 | ND ND . 68 2. 2 ND ND ND | ND ND ND ND ND ND | ND ND 230 450 160 ND ND | ND ND ND ND ND 1.8 | 5800 ND 4500 3500 ND ND 7300 | ND ND 4.4 ND 1.2 5.2 4.8 | ND ND . 13 . 87 2. 6 2. 5 | ND 11 ND ND . 07 . 16 | ND ND 38 ND 170 ND . 11 | ND ND ND 86 ND ND | ND ND 1.4 ND ND 7.0 7.5 | ND ND ND ND ND . 50 8.9 | ND ND .54 ND ND 1.2 ND |
| | Na | К | Sc | Cr | Mn | Fe | Co | Zn | Br | Ag | Cd | Sn | Cs | Ba | La | Sm | Eu | Lu | Hf | Ta | Pa | Yb |

NH₄⁺ and forms complex anion, $[VO_2(C_2O_4)_2]^{3-}$ with oxalate ion $(V:8-53 \mu g/g)$. Mn²⁺ is desorbed by the exchange with NH₄⁺, but Mn was not detected or detected with low concentrations (Mn:2-48 $\mu g/g$) because the stability of the nonionic complex $[Mn(C_2O_4)(H_2O)_4]^{9}$ is low. As for Fe, the portion adsorbed as Fe³⁺ is desorbed as very stable complex anion $[Fe(C_2O_4)_3]^{3-}$. Fe was detected at a fairly high concentration (Fe:13-735 $\mu g/g$). As for Zn, desorbed Zn²⁺ by the exchange with NH₄⁺ forms the stable complex anion $[Zn(C_2O_4)_2]^{2-}$. Zn was detected in almost all samples (Zn:6-24 $\mu g/g$).

3) By 1M NH₄Ac with an ion-exchange function of NH₄⁺ ion, Mg²⁺, K⁺ and Ca²⁺ were extracted from clay minerals in the order, Ca>Mg>K (Ca:113-5110 μ g/g, Mg:150-470, K:51-429).

4) Desorption behavior with 0.01M EDTA • 2Na is as follows. EDTA makes ionic chelates with alkali earth metal elements. Mg^{2+} and Ca^{2+} were detected in almost all samples. Among them, the desorbed amounts and rates of Ca with EDTA are especially high. This is because when $edta^{4-}$ ions as doner ions arranged around a metal ion, the size of Ca^{2+} ion (1.18Å, $e^2/r=3.4$) is suitable for making a chelate compound having maximum stability (stability•constant, $\log K_1=$ 10.6). The size of Mg^{2+} ion is smaller than Ca^{2+} ion ($e^2/r=4.9$), nevertheless the desorbed amounts and rates of Mg are smaller than Ca^{2+} by more than one order. This is because chelate compounds having lower stability constants ($\log K_1 =$ 8.7, etc.)⁹ than those of Ca were formed.

On the other hand, among transition metal elements, the detected elements in this experiment, Ti, V, Mn, Fe, Cu, and Zn, form coordinate bonding chelates with EDTA. Except V among them, generally the desorption rates of these elements were proportional to the stability constants of chelates. About the sediment samples collected at St.1 surface and St.1 20cm depth, desorption amounts and rates with 0.01M EDTA \cdot 2Na and the stability constants of chelates are shown in Table 7. According to this table, the order of the desorption rate of transition metal elements are

 $Mn^{2+}>Ti^{4+}>Zn^{2+}>Fe^{3+}>Cu^{2+}$ (St.1 surface)

 $Ti^{+}>Zn^{2+}>Fe^{3+}$ (St.1 20cm depth).

The desorption tendency was in accordance with the order of the stability constants, $Cu^{2+}>Ti^{4+}>Zn^{2+} \ge Fe^{3+}>Mn^{2+}$, except for Mn^{2+} and Cu^{2+} ions. These Mn^{2+} and Cu^{2+} anomalies show that a greater part of Mn is in inorganic forms and a greater part of Cu is in organic forms as reported before by Matsuda et al³⁰. The existing modes of chemical forms of these elements coincide with the report by Sugimura et al⁵⁰.

5) In 6N HCl extracting liquid, Al and Fe which are thought to be mineral components were detected with the highest concentrations among all extracting liquids tested. For a typical sample, the maximum concentration of Fe seen in the samples was obtained by extraction with 6N HCl(7.6 mg/g). In comparison, the

| Element Sample | Mg²+ | Ca²+ | Ti⁴+ | Mn²+ | Fe²+ | Cu²+ | Zn²+ |
|---|-------|-------|-------|-------|--------|-------|---------------|
| *Conc. (μg/g) St. 1 sf | 27700 | 12700 | 3990 | 488 | 39900 | 959 | 608 |
| St. 1 20cm | ≦8300 | 33800 | 5320 | < 480 | 63200 | ≦200 | 378 |
| St. 1 Desorption $(\mu g/g)$ | 29 | 2590 | 542 | 79 | 1373 | 8.5 | 42 . 7 |
| surface Desorption (%) | 0.105 | 2.04 | 13. 6 | 16. 2 | 3. 44 | 0.886 | 7.02 |
| St. 1Desorption (μg/g)20cmDesorption (%) | 110 | 5590 | 457 | 16 | 578 | 3.3 | 13.7 |
| | ≧1.3 | 16.5 | 8. 59 | >3.3 | 0. 915 | ≧1.7 | 3.62 |
| Stability constant, logK ₁ of metal-EDTA chelates ⁹⁾ | 8.7 | 10.6 | 17.5 | 14.5 | 15.2 | 18.8 | 15.9 |

Table 7 Desorption with 0.01M EDTA and the stability constants of chelates

 $* \leq$ and < show the values of the estimated limits because the elements were not detected.

Fe concentration observed after extraction with ultrapure water, artificial seawater, and 1M ammonium acetate were essentially zero. However, the amounts seen with 0.01M EDTA \cdot 2Na and 0.2M ammonium oxalate were as high as 18% and 9.5% of the 6N HCl value, respectively. Hg and Pb were extracted which were not desorbed by any other extracting liquids. In 1N HCl, mineral components were generally less desorbed than 6N HCl, as expected.

6) As for St.1 surface and 20cm deep layers, elements detected by both PIXE

and NAA methods had general tendencies that were similar to each other. However, the NAA values were greater than the PIXE values. It is thought that the main reason for these discrepancies is the evaporation loss of volatile matter in vacuum and during proton bombardments in vacuum.

7) With most elements, excepting a few elements (especially as seen in St.1 case Ca with each extracting liquid), desorption amounts tended to be greater in surface layer than in the 20cm deep layer.

8) From the NAA results, light rare earth elements, La, Sm and Eu, were extracted with chelate agent EDTA, 1N HCl and 6N HCl regularly, and the order of the desorpted concentrations were EDTA<IN HCl<6N HCl.

In summary, these techniques, chemical extraction plus PIXE, are shown to be very useful to understand adsorption/desorption behavior in environmental conditions. The accumulation of the experimental data obtained by continuation of the study should clarify the effects of seashore through the formation of a fundamental base of the study of marine science.

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