Colloidal toxic trace metals in urban riverine and estuarine waters of Yantai City, southern coast of North Yellow Sea

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Highlights

- An optimized method for determining aquatic colloidal metals was provided.
- The real water samples were successfully tested and obtained excellent results.
- Colloidal metal heterogeneity in two urban rivers of coastal China was studied.
- Strong ligand, mixing and flocculation co-regulated colloidal dynamics in urban rivers.

Abstract

The environmental characteristics of colloidal toxic trace metals Cd, Cu and Pb in riverine and estuarine waters collected from two urban rivers of Yantai City in eastern China, the Guangdang and Xin’an Rivers, were investigated using a modified centrifugal ultrafiltration (CUF) method in conjunction with acid extraction and inductively coupled plasma mass spectrometry. The target metals in dissolved pool were divided into four CUF fractions, i.e. <1 kDa, 1–3 kDa, 3–10 kDa and 10 kDa–0.2 μm, and the results showed that colloidal Cd, Cu and Pb were dominated by 1–10 kDa (1–3 and 3–10 kDa), 1–3 kDa and 10 kDa–0.2 μm fractions, respectively. The coagulation/flocculation of low-molecular-weight (1–10 kDa) colloidal Cd and Cu in the estuaries was obvious and strong, while the enrichment of dissolved Pb in the 10 kDa–0.2 μm fraction may be mainly related to its biogeochemical interactions with Fe-oxides, which is easy to occur in macromolecular colloids. In addition, the actual molecular weight cutoffs (MWCOs) of the three used CUF units with nominal MWCOs of 1, 3 and 10 kDa were determined to be 4.9, 8.5 and 33.9 kDa, respectively, indicating that membrane calibration is essential for explaining the actual fraction of dissolved trace metals and verifying the integrity of ultrafiltration membrane. Overall, the results in this study provide a further understanding of the heterogeneity in biogeochemical features, migration and fate of toxic trace metals in aquatic ecosystems, especially that of the river-sea mixing zone.

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1. Introduction

Trace metals exist in various physical and chemical forms in natural water systems and play an important role in the biogeochemical cycle (Li et al., 2015). Based on size, metals are divided into three fractions, i.e. particulate, colloidal and truly dissolved phases (Lead and Wilkinson, 2006). Within these three forms, dissolved phase is usually defined as the portion that can pass through the 0.45 μm filter, but this operational definition ignores the colloidal phase in water system (Waeles et al., 2008).

The specific size range of the colloidal state has not been uniformly defined in previous studies, and the main pore diameters of the filters used for pre-filtering seawater were 0.2 (Guo et al., 2000), 0.45 (Zhou et al., 2016), and 1 μm (Zhang et al., 2007). Meanwhile, a variety of techniques have been successfully applied for the separation of colloidal materials, such as cross-flow ultrafiltration (Liu et al., 2013), centrifugal ultrafiltration (Xu et al., 2018) and field flow fractionation (Zhou and Guo, 2015). Due to the small amount of sample, easy operation and low cost, centrifugal ultrafiltration (CUF) is considered to be a high-efficiency colloidal substance separation technology. The CUF is also one of the few separation methods that do not require auxiliary reagents (Burba et al., 1998). But no matter which size standard or separation technique was used, the study found that a large part of the dissolved metal was present in the macromolecular organic complex state (i.e. the colloidal phase), which largely controls the operation of the marine ecosystem (Wen et al., 2011). Therefore, the determination of the metal forms in aquatic systems is critical for the understanding of the geochemical cycle and bioavailability of trace metals (Liu et al., 2013; Cornu et al., 2018). Among them, as an important transition zone between freshwater and seawater, the strong physico-chemical behaviors and hydrodynamic gradients of the estuary zone could have great impact on metal transfer and geochemical cycles (Waeles et al., 2008). Although previous studies have reported colloidal metal transfer as a function of salinity in the estuary area, studies of transfer behavior between various low-molecular-weight (LMW) colloidal fractions (1–10 kDa) are particularly inadequate (Waeles et al., 2008).

In this study, the distribution of Cu, Pb and Cd in four size fractions (i.e. <1 kDa, 1–3 kDa, 3–10 kDa and 10 kDa–0.2 μm) in estuaries of the Guangdong River and the Xin'an River in the northern part of the Shandong Peninsula (Eastern China) is reported. The two rivers are located in typical Chinese coastal urban systems (Zhang et al., 2016), and occupy an important area of the Shandong Peninsula Blue Economic Zone which is integrated in the national development strategy (Huang et al., 2013). The originality of this work is the combined use of CUF, acid extraction and ICP-MS to separate and measure trace elements in colloidal/dissolved fractions along an estuarine salinity gradient. Besides, the molecular weight cutoffs (MWCOs) of the ultrafiltration membranes used in this study were calibrated using standard macromolecules in combination with UV spectrophotometry and high performance liquid chromatography.

2. Material and methods

2.1. Reagents

The mixed standard solutions for three target elements (Cd, Cu and Pb, 1000 mg L⁻¹) were purchased from the General Research Institute for Nonferrous Metals (China). All used ultra-pure deionized water (DIW) of 18.2 MΩ-cm in resistivity was provided by a Pall Cascada™ lab water purification system (Pall Corporation, USA). Different concentrations of nitric acid (HNO₃, Sinopharm Chemical Reagent Co., Ltd., China) and hydrochloric acid (HCl, SinoPharm Chemical Reagent Co., Ltd., China) solution were prepared by diluting the concentrated HNO₃ and concentrated HCl which were re-distilled through a sub-boiling distillation apparatus (BSB-939-IR, Berghof, Germany). The internal-fluorinated HDPE bottles (Nanjing Shuishan Plastic Products Co., Ltd., China) for storing the cleaning reagents (NaOH and HCl), ultrafiltered and pre-filtered water samples were first cleaned with 10% Decon 90™ detergent (Decon Laboratories Ltd., UK) and then sonicated sequentially in 1.5 mol L⁻¹ HNO₃, 1.2 mol L⁻¹ HCl and DIW for 3 h, modified from the cleaning process in Li et al. (2015). All the used experimental labwares were also pre-cleaned following the same cleaning process, which were rinsed 3–5 times with DIW between each washing cycle and allowed to dry in a Class 100 clean bench. Finally, each dried bottle was double-packed and stored at −25 °C for sampling. All experiments were performed in the Class 100 clean bench in the Yantai Institute of Coastal Zone Research, Chinese Academy of Sciences.

2.2. Sample collection

Surface samples (1 m deep), including riverine and estuary waters of two rivers, were collected in Yantai City, Eastern China, in January 2019 (Fig. 1). Six sampling sites, namely 1, 2, 3, 4, 5 and 6, were on the verge of a rapidly developing aquaculture zone of China (Zhang et al., 2016). Surface water samples were collected using trace-metal-clean pole-sampling techniques that was equipped with a Teflon pole and attached to internal-fluorinated HDPE bottles at the end. After the water sample was collected, the sampling bottle was double bagged in polyethylene bags, and then stored in a sample box at −4 °C (Lu et al., 2019). The salinities of the 6 sampling sites (No. 1–6) were 29.6, 26.7, 0.5, 28.6, 24.0 and 0.3, respectively, measured by using the SX713 Model salinometer from Shanghai San-Xin Instrumentation, Inc (Shanghai, China). Water samples were taken back to the laboratory and filtered (within 2 h) by using an enclosed filtration system, which was actually a pre-cleaned polysulfone vacuum filtration device (DS0320-5045, Nalgene™). Thermo Fisher Scientific Inc., USA equipped with a perforated screw cap and a diaphragm vacuum pump (Tianjin Jingteng Experimental Equipment Co., Ltd., China). The unfiltered water samples were pumped along the C-flex™ tubing (Cole-Parmer Inc., USA) while vacuum filtration was performed. Finally, pre-filtered samples were collected in another pre-cleaned 1 L internal-fluorinated HDPE bottles and then stored in freezer at 4 °C before further processing. The Supor™-200 0.2 μm membrane filters (Pall Corporation, USA) used in this work were pre-cleaned with 2% HNO₃ and then rinsed with DIW before use.

2.3. CUF

A fixed rotor centrifuge (Velocity 14R, Dynamica Corporation, Australia) was used with three kinds of CUF units (Macrosep™, Pall Laboratory, USA) with respective cutoffs of 1 kDa (MAP001C38), 3 kDa (MAP003C36) and 10 kDa (MAP010C38). Supplementary Material Fig. S1 shows a schematic diagram of the experiment process. Before CUF, the unit was thoroughly cleaned with a method modified from Guo et al. (2000) and Xu and Guo (2017), i.e. sequentially washed 5–10 times with 0.05 mol L⁻¹ NaOH, 0.02 mol L⁻¹ HCl and DIW to remove possible contamination. The DIW blank was collected to check that the leaching of metals is not contributing to the concentration in samples. Then, the CUF unit was loaded with 15 mL of samples (<0.2 μm), and centrifuged (4200×g, 50–70 min) to less than 0.5 mL at 4 °C constant temperature to separate the concentrates (>1 kDa, >3 kDa and >10 kDa) for the determination of concentration of colloidal metals (Cd, Cu, Pb).
and Pb) with different molecular weights. Then, after CUF, 3% HNO₃ was added 3 times with 1 mL each time to the concentrate for extraction (≥1 h each time). Finally, three extracts were mixed and volume adjusted to 5 mL with 3% HNO₃ (v/v). As for the ultrafiltrate, 0.2 mL was diluted to 10 mL with 3% HNO₃ (v/v). <0.2 μm total dissolved fraction was prepared by diluting 0.2 mL of prefiltered water sample to 10 mL with 3% HNO₃. All treated samples were stable at 25 °C for ≥48 h and then measured using ICP-MS. Colloidal trace metal concentrations in different size ranges were calculated based on the difference in concentration between the initial sample and the ultrafiltrate samples (Xu et al., 2018). Based on this method, fractions can be defined as <1 kDa, 1–3 kDa, 3–10 kDa and 10 kDa–0.2 μm. In this way, a concentration factor (cf) of 3 was achieved for the colloids and of 0.02 for the ultrafiltrates (a 50 times dilution). After the whole experiment was completed, each CUF unit was recirculated, and then poured into sterile water and stored at 25 °C in the dark.

2.4. Membrane calibration

To check the deviation of the MWCO of the ultrafiltration membrane used in this experiment from the manufacturer's cutoff ratings, we used standard macromolecules to calibrate the CUF unit, including vitamin B₁₂ (VB₁₂, Aladdin Industrial Corporation, Shanghai, China), with a nominal molecular weight (NMW) of 1.3 kDa, and standard fluorescent tagged dextrans (FTDs, TdB Consultancy AB, Uppsala, Sweden) with a NMW of 4.5, 9.5, 19.5 and 40 kDa, respectively.

Briefly, the standard macromolecules were added to the prefiltered seawater and thoroughly mixed (Supplementary Material Fig. S1), followed by CUF treatment. Then, the concentrations of the standard macromolecule in ultrafiltrate, concentrate and prefiltered water sample to 10 mL with 3% HNO₃. All treated samples were stable at 25 °C for ≥48 h and then measured using ICP-MS. Colloidal trace metal concentrations in different size ranges were calculated based on the difference in concentration between the initial sample and the ultrafiltrate samples (Xu et al., 2018). Based on this method, fractions can be defined as <1 kDa, 1–3 kDa, 3–10 kDa and 10 kDa–0.2 μm. In this way, a concentration factor (cf) of 3 was achieved for the colloids and of 0.02 for the ultrafiltrates (a 50 times dilution). After the whole experiment was completed, each CUF unit was recirculated, and then poured into sterile water and stored at 25 °C in the dark.

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Briefly, the standard macromolecules were added to the prefiltered seawater and thoroughly mixed (Supplementary Material Fig. S1), followed by CUF treatment. Then, the concentrations of the standard macromolecule in ultrafiltrate, concentrate and prefiltered sample reached extremely high levels. In order to prevent sample overflow during centrifugation, an initial sample volume of 15 mL in the ultrafiltration unit was chosen. As two factors that directly affect the ultrafiltration efficiency, optimization of centrifugation time and centrifugal force is necessary. When the centrifugal force is constant, the ultrafiltration efficiency can be increased by prolonging the ultrafiltration time, however excessive time is likely to cause membrane rupture and shortens its service life.

3. Results and discussion

3.1. Centrifugal force and centrifugation time

Ultrafiltration efficiency is affected by many factors, such as sample viscosity and initial volume, membrane pore size, centrifugation angle, centrifugal force and centrifugation time (Elshobaby et al., 2018; Loginov et al., 2017). Xu and Guo (2017) found that when the volume of the concentrate on the ultrafiltration membrane was <500 μL after CUF, the concentration ratio of the colloidal substance and the desalting efficiency of the seawater sample reached extremely high levels. In order to prevent sample overflow during centrifugation, an initial sample volume of 15 mL in the ultrafiltration unit was chosen. As two factors that directly affect the ultrafiltration efficiency, optimization of centrifugation time and centrifugal force is necessary. When the centrifugal force is constant, the ultrafiltration efficiency can be increased by prolonging the ultrafiltration time, however excessive time is likely to cause membrane rupture and shortens its service life.

![Fig. 1. Study area and sampling sites in the rivers and estuaries of Yantai City, China. The black dots indicate sampling sites.](image-url)
life (Sun et al., 2015). Therefore, the water sample (site 1) was used to investigate the effect of the centrifugal force (1000, 2000, 3000 and 4000 g) and the centrifugation time (20, 30, 40, 50, 60 and 70 min) on the extraction efficiency. It should be noted that the volume of the concentrate was approximate because it could not be accurately measured. 10 µL, 100 µL, 1000 µL and 5 mL pipettes were used for volume transfer and estimation during the experiment. By measuring the remaining volume of the concentrate, optimal conditions for each MWCO CUF units could be determined (Supplementary Material Fig. S3a-c).

As can be seen from Supplementary Material Fig. S3a-c, the concentrates in the three CUF units reached a minimum volume under test conditions of NO. 24 (4000 g, 70 min), with volumes of ~450 (1 kDa), ~300 (3 kDa) and ~310 µL (10 kDa), respectively. However, the experimental conditions satisfying the experimental requirements of the concentrate volume of <500 µL were quite numerous. For example, a 3 kDa ultrafiltration centrifuge unit could concentrate the sample to <500 µL in NO. 18 (3000 g, 70 min), 22 (4000 g, 50 min), 23 (4000 g, 60 min) and 24 (4000 g, 70 min), 10 kDa could reach the requirements in both NO. 22 (4000 g, 50 min), 23 (4000 g, 60 min) and 24 (4000 g, 70 min). Therefore, in order to meet both the centrifugal effect and the shortest experimental time, the 1 kDa unit was used under the NO. 24 condition (4000 g, 70 min) and the 3 and 10 units were used under the condition of NO. 22 (4000 g, 50 min).

As matrix effects can alter the centrifugation efficiency from one sample to another, the optimal ultrafiltration parameters calibrated for sample 1 were tested on the five remaining samples (Supplementary Material Fig. S3d). As sample 3 (riverine water) was the only sample that was subjected to centrifugation with a 10 kDa unit, its centrifugal force of 4200 g and centrifugation time of 70 (1 kDa), 50 (3 kDa) and 50 min (10 kDa), respectively.

### 3.2. Extraction method

In order to investigate the ability of ultrapure water and different concentrations of HNO₃ to extract colloidal Cd, Cu and Pb trapped on the membrane, sample 1 was selected and centrifuged ultrafiltration according to the method described in Section 2.3. 6 different HNO₃ concentrations (0.0, 0.5, 1.0, 1.5, 2.0 and 3.0%, v/v) were tested to assess the efficiency of Cd, Cu and Pb extraction from concentrated colloids prior to the 3% HNO₃ 5 mL dilution step. In addition, 0.2 mL of ultrafiltrate was aspirated and the same extract was used to dilute it to 10 mL. All samples were measured using ICP-MS and the results of recovery (or extraction rate) are shown in Fig. 2.

The results showed that the recoveries of colloidal Cd, Cu and Pb in the DIW extracts of the three ultrafiltration centrifuge units were lower, only 20.7–70.9%. However, when the concentration of HNO₃ was higher than 1.5%, the recoveries of Cd, Cu and Pb measured in different concentrations of HNO₃ extract were close and stable, both higher than 90.3%. This indicates that the method blanks were low and the target metals could be efficiently extracted by a higher concentration (~1.5%) of HNO₃ solution. Since the dilution of the standard solution and the working curve was 3% HNO₃ in the ICP-MS measurement, 3% HNO₃ was selected as the acidified extract.

### 3.3. Actual molecular weight cutoff (MWCO) of ultrafiltration membrane

It should be noted that the manufacturer-rated MWCOs of CUF units may be different from the actual MWCOs (Guo et al., 2000; Xu and Guo, 2017). Therefore, after the calibration process described in Section 2.4, the actual MWCOs of the used CUF units and their retentions (R) were evaluated. Retention characteristics of different standard macromolecules by CUF units are shown in Supplementary Material Fig. S4.

As depicted in Supplementary Material Fig. S4, for the 1 kDa unit, R of VB₁₂ was about 1.3%, which indicated that the manufacturer-rated MWCO of CUF was much lower than the actual MWCO of the ultrafiltration membrane. As for 3 kDa unit, R was ~83.1% for the 4.5 kDa FTD although it reached 93.4% for the 9.5 kDa FTD. The 10 kDa unit was no exception; R was ~77.6% for the 19.5 kDa FTD although it reached 93.3% for the 40 kDa FTD. The corresponding molecular weight based on the 90% R was taken as the actual MWCO of the ultrafiltration membrane (Guo and Santschi, 2007), and its value was 4.9, 8.5 and 33.9 kDa for the manufacturer-rated 1, 3, and 10 kDa membranes, respectively, which were comparable to the corresponding values of 2.5, 7.4 and 32 kDa gained through the same method by Xu and Guo (2017).

Yet, since the curve in Supplementary Material Fig. S4 was a fitting curve, the results obtained from the figure may be different from the true value. For example, the retention of a 1 kDa ultrafiltration membrane for a 4.5 kDa FTD was 97.1% (see Supplementary Material Fig. S4), which indicated that the actual MWCO of 1 kDa ultrafiltration membrane should be less than 4.5 kDa, but the value from the graph was 4.9 kDa. However, it was undeniable that the ultrafiltration performance of the ultrafiltration membrane could still be better evaluated by the above necessary calibration experiments. Besides, the results of the calibration indicated that the rated MWCOs significantly overestimated the colloidal trace element concentration in ultrafiltrate.

The results of this study further confirmed the conclusion made in Xu and Guo (2017) that the actual MWCO for the 1, 3, and 10 kDa CUF units were estimated to be >2, >7, and >32 kDa, respectively, and the MWCO was not necessarily the same even with the different batches of CUF units produced by the same manufacturer.

### 3.4. Mass balance

The mass balance directly determines the quality of the analytical test results (Liu and Lead, 2006; Liu et al., 2007). In this study, sample 1 was selected as the study object and mass balance was performed by adding known concentrations of metal standard solutions to the filtered seawater sample (<0.2 µm). Then, by using the CUF method described in Section 2.3, the samples were separated according to different molecular weight ranges (<1 kDa, 1–3 kDa, 3–10 kDa and 10 kDa–0.2 µm), and then 3% HNO₃ was used for extraction and dilution. Spiked total dissolved fraction (<0.2 µm) was prepared using the same dilution method (a 50 times dilution). After being respectively processed by CUFs with

| Element | Linear ranges (nmol L⁻¹) | Linear correlations (R²) | Detection limit (nmol L⁻¹) | Quantification limit (nmol L⁻¹) |
|---------|--------------------------|--------------------------|---------------------------|-----------------------------|
| Cd      | 0.2–89.0                 | 0.9999                   | 0.022                     | 0.073                       |
| Cu      | 1.6–786.8                | 0.9999                   | 0.019                     | 0.131                       |
| Pb      | 0.2–120.7                | 0.9999                   | 0.005                     | 0.015                       |
1, 3 and 10 kDa MWCOs and the other necessary pre-processing, the concentrations of the three target metals were obtained by 3 parallel determinations of ICP-MS under the optimal conditions. The recoveries (mass balance) of three metals were obtained and calculated by the following form (Liu et al., 2013):

\[
\text{Recovery} = \frac{C_c V_c + C_u V_u}{C_t V_t} \times 100\%
\]

C: metal concentration; V: volume; c, u and t: concentrate, ultrafiltrate and total dissolved phase (<0.2 μm). If the recovery is <100%, the target element is lost during CUF; if the recovery is >100%, it indicates that the target element is contaminated during CUF. The corresponding results are shown in Fig. 3.

The results showed that the recoveries of the three CUF units at different spike levels were 85.3–100.8%, which proved that the developed method could be used to determine the three colloidal metals in the actual water samples. At the same time, a thorough pre-cleaning process is indispensable and necessary for improving the accuracy of the experiment and the efficiency of the entire CUF system (Guo and Santschi, 2007).

3.5. Contrast with other methods

Although the separation and determination of colloidal trace metals in seawater has become no longer complicated, the diversity of available colloids separation techniques and their operational characteristics leads to a lack of standardized methods within the community (Wen et al., 1996; Liu et al., 2013; Hargreaves et al., 2017). The method described in this study was compared with other approaches in literature (Table 2). Wen et al. (1996) proposed a method for the collection of colloidal material for trace metal phase speciation studies in marine environments. The pre-cleaning time of the cross-flow ultrafiltration (CFUF) system was >1.5 h, and the cf was 5–10. Liu et al.
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3.6. Method application

In this study, the surface waters of 6 sites in Yantai City (sites 1–6 in Fig. 1) were collected, and the concentrations of studied metals were analyzed by the proposed method to evaluate the applicability. The results of the colloidal trace metal concentrations and percentages for each molecular weight are shown in Figs. 4 and 5, respectively. For the sake of comparison, concentrations and percentages of studied metals in collected samples and other aquatic systems are shown in Table 3.

In terms of concentrations (Fig. 4), the results showed that the <1 kDa fraction of Cd, Cu and Pb in the Guangdang River (sites 1, 2 and 3) were 0.12–0.24, 1.13–30.93 and 1.52–1.65 nmol L⁻¹, respectively, the 1–3 kDa fraction of Cd, Cu and Pb were 0.007–0.01, 0.24–2.03 and 0.06–0.34 nmol L⁻¹, respectively, the 3–10 kDa fraction of Cd, Cu and Pb were 0.0006–0.01, 0.03–2.32 and 0.02–0.31 nmol L⁻¹, respectively, and the 10 kDa–0.2 µm fraction of Cd, Cu and Pb were 0.01–0.02, 0.08–2.54 and 0.16–0.56 nmol L⁻¹, respectively; in the Xin’an River (sites 4, 5 and 6), the concentrations of Cd, Cu and Pb were 0.14–0.23, 0.81–22.09 and 1.21–2.53 nmol L⁻¹ in the <1 kDa fraction, 0.001–0.02, 0.12–2.37 and 0.02–0.05 nmol L⁻¹ in the 1–3 kDa fraction, 0.003–0.01, 0.06–1.68 and 0.01–0.02 nmol L⁻¹ in the 3–10 kDa fraction, and 0.01–0.02, 0.06–2.30 and 0.20–0.26 nmol L⁻¹ in the 10 kDa–0.2 µm fraction. Obviously, the concentrations of the three metals in each fraction significantly increased along the river, which could be attributed to a release from sediments during the estuarine mixing (Santos-Echeandia et al., 2008). However, under similar sampling conditions, the concentrations of the three metals in the two rivers in each fraction were not exactly the same, especially Pb, showing the most obvious difference (Fig. 4e and f). Meanwhile, the geographical proximity of the two rivers did not
produce the expected similar results. On the contrary, in the LMW colloids (1–10 kDa), Pb showed significant differences in the two rivers, which was mainly attributable to the physical and chemical characteristics. We compared the results with other published data (Table 3) and found that the concentrations of colloidal Cd in the two studied rivers were close to that of the Venice Lagoon (0.0001–0.04 nmol L\(^{-1}\)) (Martin and Dai, 1995), but significantly lower than that of the Penzé estuary (0.07–0.30 nmol L\(^{-1}\)) (Waeles et al., 2008); the concentrations of colloidal Cu were similar to that of the Venice Lagoon (1.89–8.34 nmol L\(^{-1}\)), but higher than that of the Penzé estuary (0.07–0.30 nmol L\(^{-1}\)); colloidal Pb, due to the obvious difference in colloidal concentration between the two rivers, was at a lower colloidal concentration level in the Xin'an River, which was close to the Penzé estuary (0.03–0.27 nmol L\(^{-1}\)), and the concentrations in the Guangdang River were higher than that of the Venice Lagoon (0.002–0.97 nmol L\(^{-1}\)).

As for the percentages (Fig. 5), truly dissolved Cd, Cu and Pb (<1 kDa) in the Guangdang River were 83–87%, 76–84% and 58–87% of their respective total dissolved pools (<0.2 \(\mu m\)), respectively, the 1–3 kDa fraction of three metals were 3–5%, 4–16% and 3–12%, respectively, the 3–10 kDa fractions were 0.4–4%, 2–6% and 1–12%, respectively, and the 10 kDa–0.2 \(\mu m\) fractions were ~7%, 5–7% and 8–14%, respectively; in the Xin'an River, the corresponding values of Cd in the four fractions (i.e. <1 kDa, 1–3 kDa, 3–10 kDa and 10 kDa–0.2 \(\mu m\)) were 80–90%, 1–8%, 2–4% and 7–8%, respectively, the percentages of Cu in the four fractions were 77–83%,
Fig. 5. Percentage distribution of Cd, Cu and Pb in the dissolved fractions of the samples. (a), (c) and (e) represent the corresponding values of Cd, Cu and Pb in Guangdang River, respectively, and (b), (d) and (f) represent the corresponding values of Cd, Cu and Pb in Xin'an River, respectively.

| Location           | Prefilter (µm) | Membrane’s cutoff (kDa) | cf | C_T (nmol L⁻¹) | C_C (nmol L⁻¹) | C_C/(C_T + C_C) (%) | Reference          |
|--------------------|----------------|-------------------------|----|----------------|----------------|---------------------|--------------------|
| Guangdang River    | 0.2            | 1, 3 and 10             | >30| 0.12–1.13–1.52–| 0.02–0.35–0.24– | 13–16–13–         | This study         |
| Xin'an River       | 0.2            | 1, 3 and 10             | >30| 0.14–0.81–1.21–| 0.02–0.24–0.23– | 10–17–10–         | This study         |
| Venice Lagoon, Italy | 0.4          | 10⁷                    | ~1.1| 0.007–3.15–0.02– | 0.001–1.18–0.002–| 34–41–58          | Martin and Dai (1995) |
| Penze estuary, France | 0.2 and 0.45 | 5, 10, 30, 50 and 300  | 13 | 0.01–0.13–0.003–| 0.07–2.14–0.03– | 82–94–91          | Waeles et al. (2008) |

*10 kDa was used as the boundary between truly dissolved and colloidal phases in that study.
7–11%, 5–6% and 6–8%, respectively, and the percentages of Pb in the four fractions were 84–90%, 1–2%, ~1% and 8–14%, respectively. For colloidal Cd (Fig. 5a and b) and colloidal Pb (Fig. 5e and f), the percentages of 10 kDa–0.2 μm fraction were higher than the 1–3 kDa fraction and the 3–10 kDa fraction, indicating that the colloidal material in this size range may dominate the forms of Cd and Pb in the colloidal phase. As for Cu, the 1–3 kDa fraction was clearly the dominant one (Fig. 5c and d). However, the studies by Dai and Martin (1995) in the Ob estuary and Waeles et al. (2008) in the Penzé estuary have shown that most of the colloidal Cu was present in the 10 kDa–0.2 μm fraction of the organic matter, and colloidal Cd and Pb may be more likely to combine with the LMW colloids (1–10 kDa), which are the opposite to the results we got. These obvious differences should thus be attributed to the properties of the colloidal material in the study areas. From the comparative data in Table 3, it can be seen that the percentages of colloidal Cd and Cu of their respective total dissolved concentrations in the study area were lower than that of the Venice Lagoon (34% Cd and 42% Cu) (Martin and Dai, 1995) and the Penzé estuary (82% Cd and 94% Cu) (Waeles et al., 2008). Only the percentages of colloidal Pb in the estuary area were close to 42%, but still lower than the Venice Lagoon (58%) (Martin and Dai, 1995) and the Penzé estuary (91%) (Waeles et al., 2008).

3.7. Size distribution variations of colloidal metals along the salinity

3.7.1. Cd

Previous studies have shown that changes in the concentration of chloride ions (Cl−) have a significant effect on the dynamic of some metals in the aquatic system (Byrne and Yao, 2000; Colombo et al., 2008). Fig. 4a and b show the distinct characteristics of various Cd concentrations increasing with salinity gradient, indicating that all of the four fractions were enriched during the river-sea mixing process (Waeles et al., 2008). However, as shown in Fig. 5a and b, 10 kDa–0.2 μm Cd was always at a relatively stable level (~8%), whereas the 1–10 kDa Cd was opposite (2.9–11.3%). For example, in the Xin’an River, 1–3 kDa Cd was constantly increasing with salinity gradient, and the percentage of variation varied widely (from 0.9% to 7.8%) and larger than that of the other two colloidal fractions (3–10 kDa and 10 kDa–0.2 μm), indicating that this fraction may dominate the migration of Cd in this region. In the Guangdang river, 3–10 kDa fraction may have similar characteristics (0.48–3.7%). However, in the Guangdang River, both 1–3 kDa and 3–10 kDa fraction showed significant downward trend accompanied by a change in salinity from 26.7 to 29.6 (Fig. 5a), indicating that the coagulation/floculation of colloidal organic matter in this region was considerable, which was generally thought to occur in macro-molecular colloidal matter (Tang et al., 2001). Typically, 1–10 kDa organics are primarily considered to be humic/fulvic compounds or peptides (newly released, such as polypeptides) (Kawakami et al., 2006; Lead and Wilkinson, 2006). In the Xin’an River (Fig. 5b), the apparent enrichment of 1–10 kDa Cd could be attributed to the change of Cd from particle-desorbed phase to stable organic-Cd complex (Kozelka and Bruiland, 1998; Wells et al., 1998), and the contribution of organic matter to the formation of colloidal Cd was particularly important in winter (Waeles et al., 2005). According to Gerringa et al. (1996), the source of these organics (1–10 kDa) may be derived from the degradation of particulate organic matter or the particle desorption.

3.7.2. Cu

Obviously, compared with the estuary area, 1–3 kDa colloidal Cu in both rivers showed a decreasing trend (Fig. 5c and d) in the lower salinity zone (from ~0.3 to ~27), while the addition occurred in the 3–10 kDa fraction (Guangdang River) and 10 kDa–0.2 μm fraction (Xin’an River). This indicates that the flocculation process of LMW colloid in this area promotes the migration of the colloidal Cu to the high-molecular-weight form. However, in the Guangdang estuary, Cu in three colloidal fractions (1–3 kDa, 3–10 kDa and 10 kDa–0.2 μm) showed a removal behavior, which may be attributed to the reduction of strong Cu-ligands in this area (Tang et al., 2001). In addition, colloidal flocculation may be the main cause of Cu removal in studies of several estuaries (Apte et al., 1990; Waeles et al., 2008), but this is generally thought to occur in macromolecular colloids (10 kDa–0.2 μm), which are mainly considered to be newly released intra-cellular biomolecules (extracellular polysaccharides, peptidoglycans and proteins) or refractory humic aggregates (Kaplan et al., 1987; Lee et al., 1996; Hung et al., 2001; Lead and Wilkinson, 2006). In the Xin’an River, the removal behavior of Cu did not occur. On the contrary, the addition of three colloidal fractions indicated that the metal addition was stronger than colloidal coagulation/floculation in this region, which could be attributed to the delayed flocculation of the colloid by the strong Cu-ligands added from the sediment (Skrabal et al., 1997; Santos-Echeandia et al., 2008). In addition, as the study by Baeyens et al. (1998) in the Scheldt estuary, the contribution of suspended particles desorption to Cu addition should not be ignored.

3.7.3. Pb

In the Xin’an River, the percentages of colloidal Pb were always at a low level (≤15.0% of the total dissolved pool), and the percentage change of each colloidal fraction (from salinity ~0 to salinity ~30) was not as obvious as that of the Guangdang River (Fig. 4e, f and 9e, f). Although the variations of colloidal Pb in the Xin’an River are not dramatic (compared with the Guangdang River), in the lower salinity zone (from 0.3 to 24.0), the addition could be found in the 1–10 kDa fractions of the two rivers (Fig. 4e, f and Fig. 5e, f). However, except for the 3–10 kDa colloidal Pb, the other colloidal fraction of Pb showed similar removal in both estuaries (Fig. 4e, f and 5e, f). In general, Pb (<0.45 μm) had a significant correlation with Fe (<0.45 μm) in the estuary area (R² = 0.33, p < 0.01, n = 72, Waeles et al., 2007), and Wen et al. (1999) found that high-molecular-weight (>10 kDa) colloidal Pb was often accompanied by Fe-oxides. Therefore, Luoma (1989) proposed that the main carrier of colloidal Pb may be the colloid-Fe-oxides or colloid-Fe-sulphides complex, and the flocculation of Fe caused by the intense river-sea mixing in the estuary area may directly lead to the limitation of Pb in the estuary area (Benoit et al., 1994).

4. Conclusions

In this study, a separation and determination method of aquatic colloidal Cd, Cu and Pb was established by CUF, acid extraction and ICP-MS. By precisely adjusting the centrifugation time and centrifugal force, the colloidal metal of different molecular weights in 6 samples could be effectively separated within 70 min. In addition, combined with ICP-MS, lower LOD (0.005–0.131 nmol L⁻¹) and excellent recoveries (85.3–100.8%) were obtained. Membrane calibration experiments were also performed and the results showed that the difference between manufacturer-rated and actual MWCO would overestimate the colloidal metal concentration of each fraction in the water sample. After using the modified method for the analysis of the water samples of Yantai City, Eastern China, it could be found that Cd, Cu and Pb in the truly dissolved phase accounted for 58–90% of the total dissolved pool. In addition, Cd and Pb were more likely to bind to the 10 kDa–0.2 μm colloids, while Cu was related to the 1–3 kDa LMW colloids. By studying the size distribution variations of the trace metals along the salinity gradient, the data
explain the dynamic behaviors of the three metals in the lower salinity zone (from -0.3 to -27) and estuary area (~30). Briefly, the addition of dissolved Cd (in the low salt zone) involves the formation of CI-complexes and desorption or degradation of particulate matters, and this addition is more pronounced in winter and is strongly correlated with the 1–10 kDa colloids. The removal behaviors of 1–10 kDa Cd in the Guangdang estuary could be attributed to the coagulation/flocculation of the colloid. As for Cu, the removal behaviors in the Guangdang estuary may be the reduction of strong Cu-ligands and the coagulation/flocculation of colloids, and the intriguing additions in the Xin'an estuary may be the reason for the strong Cu-ligands introduced into the water by sediments. As a metal with strong correlation with Fe-oxides, removal behaviors of Pb at the two estuaries are significantly controlled by the coagulation/flocculation of Fe-oxides. In summary, due to the high heterogeneity of colloidal metals, detailed descriptions of various factors are needed to better understand the dynamic migration processes and biogeochemical cycling of the elements in aquatic systems, especially along the river-sea continuum.

Declaration of Competing Interest

The authors declare that they have no known competing financial interests or personal relationships that could have appeared to influence the work reported in this paper.

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Appendix A. Supplementary data

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