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Trace metals partitioning between particulate and dissolved phases along a tropical mangrove estuary (Can Gio, Vietnam)

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h i g h l i g h t s

- Partitioning and distribution of 8 trace metals were studied in a mangrove estuary.
- TSS was the main trace metals carrier during their transit in the estuary.
- The monsoon induced increased metal inputs to the estuary.
- Most of dissolved metals exhibited a non-conservative behavior whatever the season.
- Mn, Cr and As were highly reactive, OM or DO playing a key role in their dynamics.

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Mangroves can be considered as biogeochemical reactors along (sub)tropical coastlines, acting both as sinks or sources for trace metals depending on environmental factors. In this study, we characterized the role of a mangrove estuary, developing downstream a densely populated megacity (Ho Chi Minh City, Vietnam), on the fate and partitioning of trace metals. Surface water and suspended particulate matter were collected at four sites along the estuarine salinity gradient during 24 h cycling in dry and rainy seasons. Salinity, pH, DO, TSS, POC, DOC, dissolved and particulate Fe, Mn, Cr, As, Cu, Ni, Co and Pb were measured. TSS was the main trace metals carrier during their transit in the estuary. However, TSS variations did not explain the whole variability of metals distribution. Mn, Cr and As were highly reactive metals while the other metals (Fe, Ni, Cu, Co and Pb) presented stable log K_D values along the estuary. Organic matter dynamic appeared to play a key role in metals fractioning. Its decomposition during water transit in the estuary induced metal desorption, especially for Cr and As. Conversely, dissolved Mn concentrations decreased along the estuary, which was suggested to result from Mn oxidative precipitation onto solid phase due to oxidation and pH changes. Extra sources as pore-water release, runoff from adjacent soils, or aquaculture effluents were suggested to be involved in trace metal dynamic in this estuary. In addition, the monsoon increased metal loads, notably dissolved and particulate Fe, Cr, Ni and Pb.

1. Introduction

Estuaries are key environment for the transfer of trace metals from land to open ocean (Fu et al., 2013; Wang et al., 2016). They are characterized by strong physico-chemical gradients, e.g. salinity, dissolved oxygen (DO), pH, redox condition (Elliott and McLusky, 2002), as well as gradients of organic matter quantity and quality

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([Abril et al., 2002](#)), which may affect trace metal speciation and distribution. Some elements may have a conservative behavior along the estuary, and their distribution is controlled by physical mixing of river and sea waters ([de Souza Machado et al., 2016](#)). Conversely, others may have a non-conservative additive or subtractive behavior, which means that due to biogeochemical processes a gain or a loss of dissolved metal concentrations in comparison to the theoretical dilution line will be observed. Metals also have affinities with suspended solid ([Lacerda et al., 1988](#)), which are by far the most important metal carriers from the rivers to the coastal area ([Yao et al., 2016](#)). Consequently, the distribution and partitioning of trace metals between the particulate and the dissolved phases may vary along the estuarine salinity gradient due mainly to changes in water chemistry (pH, DO, organic matter) and ionic strength ([Benoit et al., 1994](#); [de Souza Machado et al., 2016](#)) which can induce trace metal desorption by metal complexation with chloride and sulfate forming soluble inorganic complexes ([Acosta et al., 2011](#); [Greger et al., 1995](#)) or metal adsorption by cation exchange capacity ([Wang et al., 2016](#); [Yang and Wang, 2017](#)).

In developing countries, the capacity of wastewater treatment plants are not sufficient enough to treat the metal loads from urban, domestic or industrial activities, which are thus released into the rivers and then estuaries. As most of the tropical estuaries are colonized by mangrove forests ([Giri et al., 2011](#)), those inorganic contaminants can thus be deposited in these ecosystems. Mangrove sediments are known to act as a sink of trace metals due to their richness in organic matter, their clay content, and their dense root systems that can efficiently trap suspended matter and their associated metals from the water column ([Chu et al., 1998](#); [Harbison, 1986](#); [Marchand et al., 2006](#); [Natesan et al., 2014](#)). However, because of the reactivity of these sediments and the alternation between oxic and anoxic conditions, trace metals bearing phases may be dissolved ([Marchand et al., 2016](#); [Néel et al., 2014](#)) and metals can, thus, be exported to adjacent tidal creeks ([Holloway et al., 2016](#)). As mangroves have specific ecological, sociological and economical roles, notably for the local population and the fishing resources ([Lee et al., 2014](#)), their ecological status and the understanding of the metal fate in their water column is highly relevant.

Can Gio is the biggest mangrove forest in Vietnam, being a well-known sea-food producing area, and having its core zone registered as a UNESCO biosphere reserve. Can Gio is also an estuary located at the confluence of Sai Gon and Dong Nai Rivers, which drain a megacity of almost 10 million inhabitants, Ho Chi Minh City. In addition, this estuary is acting as a unique gate for water traffic connecting Ho Chi Minh City to the South China Sea. Despite the long list of ecosystem services provided by the Can Gio mangrove and the increasing anthropogenic pressure on this zone ([Kuenzer and Tuan, 2013](#)), there is a lack of published studies along this estuary, especially on trace metal dynamics. Some publications were interested in organic compounds ([Minh et al., 2007](#); [Oxmann et al., 2010](#)), and recently accumulation of trace metal in sediments fringing the mangrove was studied ([Costa-Boddeker et al., 2017](#)). The only available data concerning trace metals distribution was reported in surface water of the Sai Gon River, upstream the mangrove ([Strady et al., 2017a](#)).

Within this context, our main objectives were to determine: i) the spatio-temporal variations of salinity, pH, DO, total suspended solids (TSS), particulate organic carbon (POC) and dissolved organic carbon (DOC) along this mangrove estuary, from the downstream end of Ho Chi Minh City to the South China Sea; ii) the distribution and partitioning of metals between the particulate and dissolved phases along the salinity gradient; iii) the potential influence of the monsoon on the fate of trace metals. Our main hypothesis are: i) a decrease of trace metals concentrations due to the dilution with

marine waters and/or desorption from bearing phase resulting from increased oxygen content; ii) increased inputs during the monsoon caused by elevated runoff and soil leaching. To reach our goals, water and suspended matter samples were collected for organic carbon and trace metals analysis during 24 h cycling in order to take into account the whole tidal cycling at four sites along the estuary during two distinct seasons.

2. Materials and methods

2.1. Study area

The Can Gio mangrove is located in the south of Vietnam ($10^{\circ}22'14''\text{N}$ - $10^{\circ}40'09''\text{N}$; $106^{\circ}46'12''\text{E}$ - $107^{\circ}00'59''\text{E}$) and is part of the densely populated megacity of Ho Chi Minh City (HCMC; almost 10 million of inhabitants), 35 km downstream of the City urban center and industrial zones ([Strady et al., 2017a](#); [Vo, 2007](#)). This mangrove covers an area of 35,000 ha ([Fig. 1](#)) and is a UNESCO biosphere reserve which can be divided into three areas according to their anthropogenic pressure and level of conservation ([Nam et al., 2014](#)): (i) a core zone protected for long term conservation of biodiversity, in which human activities are prohibited, (ii) a buffer zone to protect the core zone with limited human activities and (iii) a transition zone with human activities like agriculture, aquaculture, fishing, salt-pan, trading and tourism. Can Gio mangrove is home to high biodiversity with more than 200 species of fauna (e.g. planktonic and benthic organism, fishes, reptiles and amphibians) and 20 species of flora, in which three mangrove species are dominant *Avicenia alba*, *Rhizophora apiculata*, *Phoenix paludosa* ([Kuenzer and Tuan, 2013](#)). Freshwater inputs originate from the Sai Gon and Dong Nai Rivers, emptying out the South



Fig. 1. Satellite image of Can Gio mangrove forest (black area) and sampling locations along the estuary. Site 1 characterizing the riverine and urban end-member; Site 2 characterizing aquaculture activities (mainly shrimps) and the buffer zone of mangrove forest; Site 3 characterizing the core of mangrove forest; site 4 characterizing the marine end-member.

China Sea via the Long Tau and Soai Rap Rivers. The hydrology is affected by a semi-diurnal tidal regime with a tidal amplitude up to 4 m, and by typical tropical monsoon climate, with a rainy season lasting from May to October and a dry season extending from November to April. The annual average temperature and precipitation are 25.8 °C and 1400 mm, respectively. During the rainy season, highest precipitation can reach up 400 mm in September while it is usually less than 80 mm per month during the dry season. The main economic activities of local people are: aquaculture, in which shrimp farming dominate with approximately 3160 ha, fisheries, agriculture and salt production (Tuan and Kuenzer, 2012). The industrial activities are excluded of the three zones (core, buffer and transition zones) of the Can Gio mangroves but are developed to 35 km upstream the mangrove close to HCMCity urban center and along the Dong Nai river (mainly plastic and rubber production, mechanical engineering, electrical engineering, packaging, textile and dyes industry, oil activities and cement production).

2.2. Field sampling and measurements

Samples were collected during 24 h tidal cycles, with a sampling every 2 h, at four sites along the Long Tau River according to their environmental characteristics and their locations into the different zones of the mangrove described in the section above: site 1 (10°39'55"Ne106°47'30"E) characterizing the riverine and urban end-member; site 2 (10°34'19"Ne106°50'11"E) located in the buffer zone of the mangrove forest characterizing aquaculture activities (mainly shrimps); site 3 (10°31'04"Ne106°53'13"E) located in the core zone of the mangrove forest; site 4 (10°29'32"Ne106°56'55"E) characterizing the marine end-member (Fig. 1). The sampling campaigns were carried out during dry season (January and February 2015) and rainy season (September and October 2015).

The physicochemical parameters such as salinity, pH and temperature (°C) were logged continuously *in-situ* using a multi-probe (Yellow Spring Instrument® meters YSI 6920). DO was monitored with a HOB0 Dissolved Oxygen data logger (HOB0 U26-001). pH probes were pre-calibrated using buffer solutions: 4, 7 and 10 (NIST scale) at the same day prior of sampling. Probes were immersed 50 cm below the air-water interface and data were recorded every at 5 min.

At each sampling site and time, surface waters (50 cm below the air-water interface) were collected in duplicates using a bucket, immediately transferred into acid pre-cleaned polypropylene bottles (1 L) - previously rinsed with surface water. Then, a first aliquot of surface water was filtered through 0.2 µm PTFE Omnipore™ Membrane Filters for trace metal analysis. The filtrate was transferred into pre-cleaned 50 mL polypropylene, immediately acidified to pH < 2 using concentrated suprapur® HNO₃ (HNO₃/sample ¼ 1/1000 (v/v)) for dissolved trace metals M_D, while the filters were kept in pre-cleaned plastic petri dishes for suspended particulate trace metals analysis (M_P). Both samples were maintained in cool box during sampling campaign and stored at 4 °C in the laboratory for dissolved metal and at -18 °C for particulate metal (Strady et al., 2017b). Filters were then freeze dried, weighed and preserved at room temperature until analysis. A second aliquot of surface water was filtered in triplicates through pre-combusted (500 °C) and pre-weighted glass fiber filters (Whatman® GF/F 0.7 µm): the filter was stored at -18 °C for particulate organic carbon (POC) and total suspended solid (TSS) determinations while the filtrate was transferred into sterile 15 mL polypropylene, acidified using concentrated suprapur® HCl and stored at 4 °C in fridge for dissolved organic carbon (DOC) analysis.

2.3. Samples analysis

2.3.1. Dissolved metals concentrations (M_D)

Dissolved Mn, Fe, Cr, Co, Ni, Cu, As and Pb concentrations were directly measured by Thermo Scientific iCAPQ ICP-MS using internal standard calibration (AETE-ISO platform, OSU-OREME/Université de Montpellier), with a Kinetic Energy Discrimination e Argon Gas Dilution module (KED e AGD mode) (Kutscher et al., 2014). Accuracy and precision were controlled using certificate reference material: riverine water (SLRS6) for salinity lower than 1 and estuarine water (SLEW-3) for salinity higher than 1 (Table 1a).

2.3.2. Particulate metal concentrations (M_P)

Particulate Mn, Fe, Cr, Co, Ni, Cu, As and Pb concentrations were analyzed according to a total extractable metal digestion adapted from the USEPA 3051a method (USEPA, 2007). Filters were put on PTFE vessel in which 6 mL of concentrated HNO₃ and 2 mL of

Table 1
Quality control of analytical methods applied for dissolved and particulate metal concentrations analysis: a) accuracy, precision and detection limit using estuarine water SLEW-3; b) BCR-277R for wet digestion method.

a) Dissolved metal concentration analysis					
Element	Detection limit (mg L ⁻¹)	Certificated values (mg L ⁻¹)	Measured values (mg L ⁻¹)	Recovery (%)	Relative standard deviation (%)
Cr	0.022	0.183 ± 0.019	0.0182 ± 0.020	99	11
Mn	0.013	1.61 ± 0.22	1.469 ± 0.016	91	1
Fe	0.031	0.568 ± 0.059	0.661 ± 0.073	116	11
Co	0.0092	0.042 ± 0.010	0.0464 ± 0.0025	110	5
Ni	0.011	1.23 ± 0.07	1.230 ± 0.014	100	1
Cu	0.011	1.55 ± 0.12	1.491 ± 0.041	96	3
As	0.0062	1.36 ± 0.09	1.569 ± 0.013	115	1
Pb	0.003	0.009 ± 0.0014	0.0115 ± 0.0017	130	15
b) Particulate metal concentration analysis					
Element	Certificated values (mg kg ⁻¹)	Measured values (mg kg ⁻¹ , n ¼ 9)	Recovery (%)	Relative standard deviation (%)	Analytical method
As	18.3 ± 1.8	17.56 ± 0.87	96	5.0	ICP-MS
Pb	NA	28 ± 1.3	e	4.8	ICP-MS
Co	22.5 ± 1.4	22.9 ± 0.7	102	2.8	ICP-MS
Cr	188 ± 14	171.3 ± 5.4	91	3.2	ICP-MS
Cu	63 ± 7	60.5 ± 1.8	96	2.9	ICP-MS
Ni	130 ± 8	128.9 ± 3.2	99	2.5	ICP-MS
Mn	NA	835 ± 29	e	3.5	FAAS
Fe	NA	51855 ± 3146	e	6.1	FAAS

concentrated HCl were added. The vessels were placed into ultrasonic bath during 15 min and were then heated at 110 °C during 12 h in the electrical oven. After cooling, the digestions were put again in ultrasonic bath during 15 min, and temperature was then increased up to 160 °C during 4 h. The samples were cooled and filtrated to reject residue, then diluted to 25 mL with deionized water and stored in pre-cleaned PP tubes at 4 °C until analysis. All reagents were of analysis grade (Merck) and were purified using a sub-boiling quartz distillation equipment prior digestion.

Concentrations of Cr, Co, Ni, Cu, As and Pb were measured by ICP eMS (Agilent 7700x) using spiked ^{103}Rh and ^{197}Au as internal standard while Fe and Mn concentrations were determined by Flame Atomic Absorption Spectrophotometer (Shimadzu AA-6650). The analytical precision and accuracy were insured by analyzing certificate reference material estuarine sediments (BCR-277R) (Table 1b).

2.3.3. Dissolved organic carbon (DOC) and particulate organic carbon (POC) analysis

The DOC analysis were performed on a Shimadzu® TOC-L series analyzer employing a 680 °C combustion catalytic oxidation method. The analyzer was combined with a solid sample module (SSM-5000A) heated up to 900 °C for POC measurement (Leopold et al., 2013). For both, 40% glucose standard was used for calibrations. Repeated measurements of the standard at different concentrations indicated a measurement deviation <2%.

2.4. Data analysis

In order to better understand the interaction of trace metals between dissolved and particulate phases along the salinity gradient, the partitioning coefficient (K_D) was calculated. The K_D provide a quantitative value for the partitioning of metal concentration between solution and particle. It is defined as the ratio of particulate metal concentration (M_P) to dissolved metal concentration (M_D) in the water column (Turner et al., 1993).

$$K_D = M_P / M_D$$

where M_P is the particulate metal concentration; M_D is the dissolved metal concentration.

The Pearson correlation coefficient and the one-way ANOVA were performed using statistical package software (SPSS; version 23) to identify major relationships between metal concentrations and physico-chemical parameters as well as interrelations between metals together in order to identify the main factors controlling metal partitioning.

3. Results and discussion

3.1. Spatial and seasonal variations of physico-chemical parameters

Can Gio is a tropical mangrove-dominated estuary characterized by semi-diurnal tidal regime and subject to a monsoon season. All the studied parameters varied from the upstream site, at the edge of Ho Chi Minh City, to the downstream site, at the mouth of the estuary (Table 2, Fig. 2). In addition, some seasonal variations were observed. Salinity, DO and DOC were affected by seasonal change (ANOVA, $p < .01$), whereas pH, TSS and POC were not (ANOVA, $p > .05$). From the upstream site to the sea, salinity varied from 0 to 25 during the rainy season and from 5 to 25 during the dry season. The very low salinity (<2) observed at the upstream site during the rainy season, whatever the tides, was attributed to intense rainfall increasing the freshwater inflow and limiting the saline intrusion. The pH values varied from 6.8 to 7.8 during the dry season and from

6.4 to 7.5 during the rainy season, and were positively correlated to salinity ($r = 0.96$ and 0.97 during the dry and the rainy season, respectively). DO ranged from 3.5 to 7.0 $\text{mgO}_2 \text{ L}^{-1}$ and from 1.2 to 5.3 $\text{mgO}_2 \text{ L}^{-1}$ during the dry and the rainy seasons respectively, also increasing with salinity ($r = 0.86$ and 0.94 for the dry and the rainy season, respectively). Both pH and DO reflected the mixing of fresh and sea waters (Fig. 2). The acidic pH values measured at the upstream site during the monsoon are likely related to the contribution of the Sai Gon River, which is characterized by low pH, down to 5.7 (Strady et al., 2017a), possibly due to the leaching of the surrounding acidic sulfate soils (Nguyen et al., 2011), but also to the decay processes of organic inputs. The latter may also be responsible for the drastic decrease of DO values measured at this site, down to 1.2 $\text{mgO}_2 \text{ L}^{-1}$. These organic inputs may originate from domestic and urban discharges of Ho Chi Minh City urban center and also from the industrial areas located along the Sai Gon and Dong Nai Rivers.

The POC concentrations decreased along the salinity gradient but not linearly, from 3.8% to 1.3%, and from 5.0% to 1.7% during dry and rainy seasons, respectively. Those patterns suggest that POC distributions resulted both from the dilution by seawater and from the decomposition processes occurring during its transit in the estuary (Fig. 2). Decreasing POC concentrations have also been observed in most estuaries like in temperate European estuaries (Abril et al., 2002; Etcheber et al., 2007) or tropical estuaries like in the Mandovi Estuary (Shynu et al., 2015). At the upstream site, DOC concentrations ranged from 1.22 mgC L^{-1} to 5.89 mgC L^{-1} during the rainy season and could be related to intense inputs from the city and industrial zones (Strady et al., 2017a). Along the salinity gradient, DOC concentrations varied from 1.64 mgC L^{-1} to 3.54 mgC L^{-1} and were stable during the rainy season. However, during the dry season, DOC values slightly increased in the downstream part of the estuary, which might be the result of intense carbon degradation processes in well oxygenated waters (Ni et al., 2008), and possible inputs from the adjacent mangrove forest (Dittmar and Lara, 2001).

The TSS concentrations varied from 16 to 361 mg L^{-1} and 23 to 283 mg L^{-1} during the dry and the rainy season, respectively, and did not present any conclusive distribution along the salinity gradient (Fig. 2). They were in the same range than those measured upstream the mangrove by Strady et al. (2017a), whatever the seasons. However, higher values were measured at site 2, located within the outlet of small tributaries, which are suggested to be related to disturbance of bottom sediment due to confluences of watersheds (Cang et al., 2007) and the increased erosion of the river banks caused by the sinuous flow (Nam et al., 2014).

To summarize, the spatial and seasonal variations of the physico-chemical properties reflected well the mixing of the fresh and the sea waters in the estuary. The characteristics of the upstream site during the monsoon are those of the Saigon River: salinity almost null, acidic pH values, low DO concentrations, and high POC and DOC contents. We suggest that this site can be considered as the river end-member during this season. During the dry season, the characteristics of site 1 were estuarine ones, with a mix between fresh and sea waters, and thus cannot allow us to determine inputs in the estuary at this season. Concerning the downstream site, it exhibited similar salinity, pH and DO ranges during both seasons and was thus considered as the seawater end member of the studied system.

3.2. Trace metal distributions at the upstream site during the monsoon

The characteristics of the upstream site during the monsoon being those of the river end-member, and not those of estuarine

Table 2

Bi-hourly water level (m), salinity, pH, DO (mgO₂ L⁻¹), DOC (mgC L⁻¹), POC (%) and TSS (mg L⁻¹) measured at the four sampling sites (site 1, site 2, site 3, site 4) during the dry and the wet seasons.

Site	Label	Dry season							Wet season						
		Water level m	Sal	pH	DO mgO ₂ L ⁻¹	DOC mgC L ⁻¹	POC %	TSS mg L ⁻¹	Water level m	Sal	pH	DO mgO ₂ L ⁻¹	DOC mgC L ⁻¹	POC %	TSS mg L ⁻¹
S1	h0	12.8	8.84	6.87	3.89	2.8	3.76	16.6	11	0.55	6.57	1.84	2.36	5.05	44.53
S1	h2	13.6	7.4	6.81	3.47	3.25	3.84	17.23	10.6	0.2	6.55	1.21	3.88	5.04	45.66
S1	h4	14.6	7.92	6.93	4.07	3.18	3.4	18.96	10.8	0.3	6.55	1.32	5.89	4.35	58.46
S1	h6	16.2	10.68	7.09	4.62	2.94	2.6	29.03	11.3	0.64	6.59	1.8	NA	3.94	62.23
S1	h8	15.1	12.56	7.17	4.15	2.92	2.95	21.53	11.9	1.92	6.68	2.33	3.42	2.92	136.76
S1	h10	14.2	11.01	7.1	4.1	3.33	2.41	33.46	12.2	3.19	6.75	2.57	2.52	4.96	22.8
S1	h12	13.1	10.05	7.09	3.52	2.92	2.73	25.1	12	0.91	6.62	1.72	2.04	4.73	34.00
S1	h14	12.9	9.77	7.04	3.6	3.59	2.39	60.26	11.8	0.75	6.6	1.59	3.18	4.00	42.53
S1	h16	12.4	8.15	6.97	4.17	3.35	2.7	43.06	11.9	0.69	6.6	1.64	2.42	4.04	42.96
S1	h18	13.6	8.44	7	4.65	2.54	2.98	22.96	12.1	0.84	6.61	1.77	1.22	4.36	32.9
S1	h20	15.3	10.19	6.99	4.04	3.12	3.12	27.53	11.8	0.83	6.64	2.26	2.44	4.45	37.7
S1	h22	15.3	11.43	6.84	4.65	3.88	2.52	42.56	11.5	0.85	6.66	2.3	1.97	4.74	33.36
S1	h24	13.5	9.17	6.87	4.08	3.83	3.61	15.96	11	0.57	6.63	2.12	1.41	4.26	47.23
S2	h0	11.7	21.33	7.32	4.98	4.41	1.98	198.26	10.8	17.76	7.59	4.6	2.35	2.78	53.00
S2	h2	10.8	21.71	7.43	4.93	2.47	1.94	139.43	10.6	16.7	7.37	4.28	1.96	2.31	75.96
S2	h4	9.3	21.32	7.4	4.89	4.32	1.62	168.66	8	15.06	7.42	4.27	1.94	1.93	236.26
S2	h6	9.4	20.22	7.39	4.96	2.3	2.02	151.16	7.3	11.78	7.24	3.88	1.99	1.99	178.33
S2	h8	10.5	20.17	7.38	4.87	2.62	1.28	70	8.2	6.49	7.01	3.16	2.12	2.64	50.76
S2	h10	11.1	21.56	7.42	4.84	2.39	1.31	61.53	10.8	8.78	7.28	3.93	2.3	2.01	178.63
S2	h12	11.1	22.67	7.45	4.8	4.11	NA	147.9	13.3	16.28	7.46	4.07	1.64	1.67	266.8
S2	h14	9.7	21.69	7.36	4.87	4.02	NA	108.96	13.6	17.07	7.49	4.12	1.86	2.03	82.56
S2	h16	8.2	20.9	7.38	4.85	4.72	NA	206.6	8.6	15.97	7.39	4.05	1.89	2.29	190.7
S2	h18	7.5	18.8	7.33	4.85	3.62	NA	361.16	7.6	14.88	7.38	3.94	2.82	2.11	283.06
S2	h20	8.1	13.89	7.19	4.52	2.8	NA	93.96	7.3	10.31	7.17	3.51	3.54	2.22	80.23
S2	h22	10.3	17.55	7.3	4.75	3.19	NA	55.13	9.8	10.92	7.26	3.98	2.38	2.62	67.36
S2	h24	10.8	21.17	7.38	4.83	3.13	NA	281.33	10.5	15.82	7.43	3.93	2.03	2.25	170.23
S3	h0	NA	NA	NA	NA	NA	NA	NA	11.8	15.86	7.43	3.58	2.09	2.65	30.76
S3	h2	12.6	NA	NA	4.93	3.77	1.66	77.8	11.2	14.71	7.35	3.25	2.07	2.47	51.16
S3	h4	12.7	NA	NA	5.13	2.22	1.7	167	11.4	13.47	7.29	3.11	2.01	2.51	48.26
S3	h6	12.8	NA	NA	5.36	2.3	1.49	73.8	12.3	13.47	7.31	3.34	2.24	2.92	33.3
S3	h8	11.7	NA	NA	5.55	2.54	1.51	44.66	12.9	17.05	7.45	3.61	1.78	2.46	38.8
S3	h10	NA	NA	NA	NA	NA	NA	NA	13.3	18.93	7.56	3.8	2.41	2.92	43.83
S3	h12	10.3	NA	NA	4.99	2.77	1.57	107.46	13.1	20.18	7.66	4.73	2.04	2.9	36.53
S3	h14	10.6	NA	NA	4.91	2.36	1.77	52.46	12.9	16.48	7.48	4.17	1.82	3.51	31.5
S3	h16	12.4	NA	NA	4.98	3.94	1.29	53.9	12.7	16.68	7.47	3.79	2.39	2.65	33.23
S3	h18	12.7	NA	NA	5.11	2.82	1.53	54.96	12.6	16.98	7.48	3.58	2	2.86	36.86
S3	h20	11.3	NA	NA	5.17	3.1	1.48	39.7	13.1	16.9	7.48	3.72	2.08	2.92	24.56
S3	h24	10.6	NA	NA	5.22	2.79	1.5	66.8	12.9	17.21	7.5	3.88	2.14	2.78	24.5
S4	h0	12.6	26.06	7.75	6.82	3.66	1.66	73.56	12.6	16.52	7.5	4.2	2.21	3.06	23.23
S4	h2	12.3	26.1	7.81	7.01	4.48	1.58	45.43	12.3	18.93	7.39	3.86	2.2	2.22	41.96
S4	h4	12.1	25.91	7.81	6.85	3.64	1.26	42.3	13.5	23.01	7.59	4.52	1.66	1.85	62.36
S4	h6	12	24.17	7.73	6.49	3.64	1.58	40.66	14	25.61	7.7	5.27	3.19	1.87	50.06
S4	h8	12.6	25.24	7.76	6.65	4.32	2.09	34.83	12.3	22.92	7.59	4.97	2.32	1.82	60.73
S4	h10	13	25.78	7.79	6.75	4.16	2.9	34.83	12.4	21.43	7.48	4.29	2.24	1.83	89.66
S4	h12	12.5	25.56	7.77	6.61	4.84	2.37	28.66	14.1	20.09	7.44	4.09	1.89	1.84	42.8
S4	h14	11.3	23.63	7.69	6.24	3.28	1.77	78.73	14.5	23.09	7.57	4.51	1.91	2.02	51.06
S4	h16	11	22.78	7.64	5.97	3.92	2.33	121.86	13.8	24.09	7.63	4.63	2.71	2.18	48.43
S4	h18	11.4	20.98	7.58	5.84	4.44	2.38	50.63	13.1	23.74	7.59	4.83	2.35	2.09	46.46
S4	h20	11.9	23.59	7.64	5.99	4.46	1.97	93.93	11.7	21.92	7.52	4.53	3.46	2	120.93
S4	h22	12.8	25.97	7.75	6.57	3.79	1.85	98.8	11.4	19.44	7.41	4.37	2.78	2.09	102.76
S4	h24	13.1	25.82	7.76	6.6	3.8	1.89	54.33	12	18.01	7.38	3.95	3.14	2.42	39.73

waters, we choose to discuss them separately from the other data in order to evidence riverine trace metals inputs in the Can Gio Estuary. The dissolved concentrations measured during this period were (mg L⁻¹): Mn_D, 3.4e37; Fe_D, 2.4e18; Cu_D, 0.2e15; Pb_D, 0.15e2.1; Ni_D, 0.3e1.37; As_D, 0.09e1.4; Cr_D, 0.053e0.26 and Co_D, 0.01e0.5 (Fig. 4; supplementary data 1) while the particulate concentrations were (mg kg⁻¹): Fe_p, 42,959e55,238; Mn_p, 306e822; Cr_p, 73e131; Ni_p, 57e93; Cu_p, 43e76; Pb_p, 18e27; Co_p, 10e19 and As_p, 9e14 (Fig. 4; supplementary data 1). Both dissolved and particulate concentrations were in the same range as previously measured in the Sai Gon River (Nguyen et al., 2011; Strady et al., 2017a) and were in the low range of World River average concentrations (Gaillardet et al., 2014; Viers et al., 2009). We noticed that the higher dissolved and particulate concentrations were measured during the ebb tide, namely during riverine water

flowing, attesting their inputs from urban and anthropogenic sources (Strady et al., 2017a). Elevated Cr_p, Cu_p and Ni_p concentrations coincided with high POC values, as previously observed by Strady et al. (2017a) suggesting that organic matter might be an important factor affecting those metal distributions. The decrease and variation of Cr_p, Cu_p and Ni_p concentrations, at the riverine part and the beginning of estuary (Fig. 4c, e and 4f), may be thus related to the organic matter degradation.

3.3. Relationship between TSS and total trace metal concentrations

To assess the control of TSS concentrations on metal transfer in the Can Gio Estuary, the total metal concentrations in a volume of water (i.e. the particulate concentrations, expressed in mg L⁻¹, plus dissolved concentrations) were calculated (Strady et al., 2017b;

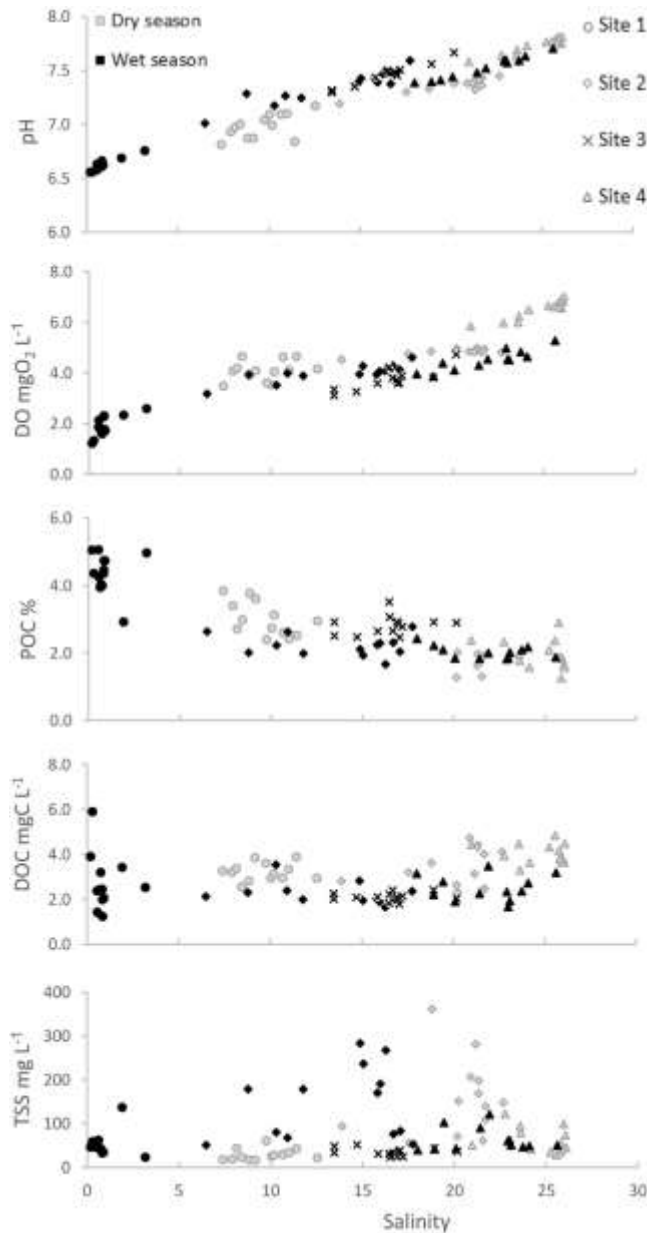


Fig. 2. Spatio-temporal variations in pH, DO, POC, DOC and TSS along the salinity gradient during the dry and the wet seasons.

Wang et al., 2016) and expressed in mg L^{-1} (Fig. 3). All total metal concentrations increased with increasing TSS concentrations along the estuary ($r > 0.95$, $p < .01$ for all metals during the dry season, and $r > 0.9$ for Mn, Fe, Cr, Ni and Pb; $r \geq 0.80$ for Co; $r \geq 0.75$ for Cu and $r \geq 0.46$ for As; $p < .01$ during the rainy season) supporting the fact that the total metal concentrations transfer in the Can Gio Estuary is controlled by the change of TSS concentrations whatever the season, like in the Mekong Delta, Vietnam (Strady et al., 2017b), in the Huanghe River Estuary, China (Wang et al., 2016) or in the Changjiang River Estuary, China (Yang et al., 2014). However, for a given TSS concentration, total concentrations in Fe, Cr, Ni and Pb were higher during the monsoon than the dry season (Fig. 3) suggesting enhanced metal transport from land to sea during the monsoon. The particulate phase was the dominant one for Fe, Cr and Co, representing more than 99% of the total for Fe, 75e99% for Cr and 70e98% for Co, while the percentage of Ni_p , Cu_p , As_p , Pb_p and

Mn_p varied from 22 to 91%, 28e95%, 13e94%, 24e99% and 29e99%, respectively. This result suggests different partitioning between particulate and dissolved metal phases during their transit along the Can Gio mangrove Estuary.

3.4. Trace metal partitioning in the estuarine zone

3.4.1. Iron

Along the salinity gradient, Fe_p varied from 35,276 to 55,302 mg kg^{-1} during the monsoon, and from 13,917 to 41,328 mg kg^{-1} during the dry season with minimum values reached at the seaside of the estuary (Fig. 4a). These lower values during the dry season could be attributed to the dilution with the sea water (i.e. Fe_p concentrations of 15,000 mg kg^{-1} in coastal zones of the Pacific Ocean (Hatje et al., 2001)). During the monsoon, dilution was not observed, probably due to intense leaching of Fe from the surrounding mangrove soils, Fe being a rich element of mangrove soils (Marchand et al., 2006; Miola et al., 2016; Neel et al., 2014). The Fe_p varied between 1.01 and 3.3 mg L^{-1} during the dry season and 1.6e5.0 mg L^{-1} during the rainy season. Along the salinity gradient, Fe_p showed a non-conservative behavior with additive concentrations at mid salinity (site 2 and 3) (Fig. 4a, supplementary data 1 and 2). During the dry season at site 2, these higher concentrations might be attributed first to the contribution of the pore-water inputs from mangrove sediments, as observed in mangrove estuaries like in Australia (Santos et al., 2011). These authors reported extremely high dissolved iron concentration up to 374 mg L^{-1} in pore-water. They also evidenced that iron in surface water originated from mangrove soils by pore-water discharge, using radio-isotopes (Radon, ^{222}Rn). In fact, mangrove soils can be rich in dissolved iron as described in the Pai Matos mangrove system, Brazil (Otero et al., 2009) or in the French Guiana mangrove (Marchand et al., 2006). This richness result from the alternation of different redox processes, notably iron oxide reduction and iron sulphide oxidation, leading both to the release of dissolved iron in pore-waters, which can thus be exported to adjacent ecosystems through pore-water seepage (Deborde et al., 2015; Sanders et al., 2015). Secondly, we assumed that irregular inputs from the shrimp farms might also be a source of dissolved iron concentration to the water column as observed in the Estuaries of East-Hainan, China (Fu et al., 2013). The Fe_p concentrations were influenced by pH changes, associated to the cultivation cycles (Azevedo et al., 2009), reaching approximately 2 mg L^{-1} in ponds on acid sulfate soils (Jayasinghe et al., 2010). Unfortunately, we were not able to determine the Fe_p concentrations at site 3 during the dry season, which was only surrounded by mangrove forest. During the rainy season, scattered Fe_p were also measured at site 2 and at site 3 at both ebb and flood tides, which may be related to pore-water releases from mangrove sediments, aquaculture ponds or runoff from the surrounding watershed. Along the Can Gio

mangrove Estuary, $\log K_d^{\text{Fe}}$ was stable whatever the season, ranging from 6.1 to 7.6 (Fig. 4a). The absences of significant correlations between $\log K_d^{\text{Fe}}$ and salinity, pH, DO, DOC, POC or TSS do not allow us to better characterize Fe partitioning along this mangrove estuary.

3.4.2. Manganese

Mn_p ranged from 402 to 1055 mg kg^{-1} during the dry season and from 397 to 922 mg kg^{-1} during the monsoon along the salinity gradient (Fig. 4b), showing an absence of seasonal effect. These values were in the same range than those measured at the upstream site during the rainy season, and in tropical estuaries like the Tan-shui Estuary, Northern Taiwan (Fang and Lin, 2002). The high Mn_p fluctuations and its absence of conclusive distribution along the

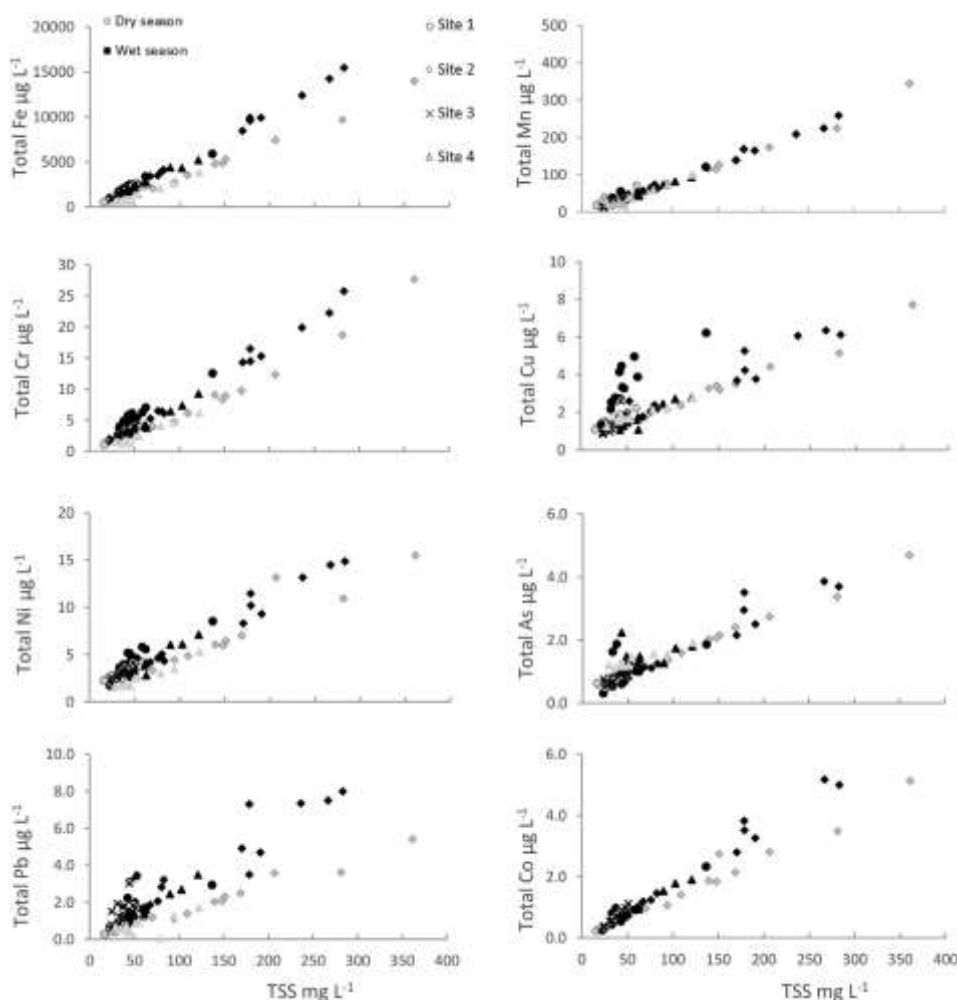


Fig. 3. Relationship between the total metal concentrations (i.e. dissolved plus particulate metal concentrations, expressed in mg L^{-1}) of Fe, Cr, Ni, Pb, Mn, Cu, As, Co and TSS concentration during the dry and the wet seasons.

salinity gradient indicated that the Mn_p distribution in the Can Gio Estuary was not dominated by a simple mixing of the two end members, and might be influenced by biogeochemical processes. The Mn_D varied between 15.2 and 0.93 mg L^{-1} and 12.5 to 0.98 mg L^{-1} during the dry and the rainy season, respectively (Fig. 4b, supplementary data 1 and 2). Additionally, it decreased along the salinity gradient exhibiting a non-conservative subtractive behavior whatever the season. Both in laboratory experiments (Hatje et al., 2003) and in the field (Fang and Lin, 2002), it was observed that DO increase can result in Mn_D adsorption onto particles surface by the oxidation of dissolved Mn (II) to insoluble Mn (III) and Mn (IV) (hydr)-oxides. Log K^D increased from 4.0 to 5.8 along the salinity gradient whatever the seasons (Fig. 4b) exhibiting good correlation with both DO and pH (Fig. 5a and b), suggesting that Mn partitioning is probably affected by the change of oxygenation in the estuary. During the dry season, higher Mn_D were measured at site 2, similarly to Fe_D , and could also originate from mangrove sediment porewaters (Holloway et al., 2016; Sanders et al., 2015) or shrimp ponds effluents (Inoue and Asano, 2013). During the rainy season, Mn_D presented scattered values at flood tide that might be related to runoff inputs of the adjacent watershed. Consequently, Mn dynamic along the estuary was strongly influenced by geochemical processes, mainly Mn precipitation resulting from oxygenation and pH changes with water mixing.

3.4.3. Chromium

The Cr_p ranged from 25 to 79 mg kg^{-1} during the dry season and from 57 to 94 mg kg^{-1} during the rainy season (Supplementary data 1 and 2), showing seasonal variations and decreasing concentrations along the salinity gradient (Fig. 4c). The Cr_p in the estuarine zone were slightly lower than in the riverine part (upstream site during the rainy season), while the Cr_p at the seaside of the estuary were close to the Cr_p reported in the coastal area of South China Sea (Cenci and Martin, 2004). Therefore, the Cr_p concentrations decrease along the salinity gradient might be attributed to both dilution with seawater containing low Cr_p and to its desorption processes during organic matter degradation in the estuary, Cr having a strong affinity for organic matter (Masscheleyn et al., 1992). We observed that later process may play an important role on Cr_p distribution during the rainy season (i.e. a positive correlation observed between Cr_p and POC, $r^2=0.71$) and a minor role during the dry season ($r^2=0.53$), affecting also Cr partitioning (Fig. 5c). The Cr_D showed distinct behaviors along the salinity gradient between seasons: i) conservative distribution with gradually Cr_D increase seaward during the dry season, and ii) non-conservative with a gain of Cr_D at mid salinity during the rainy season (Fig. 4c, supplementary data 1 and 2). Scattered and higher Cr_D at site 2 and site 3 observed during the monsoon might be originated from the same sources as Fe_D (positive correlation

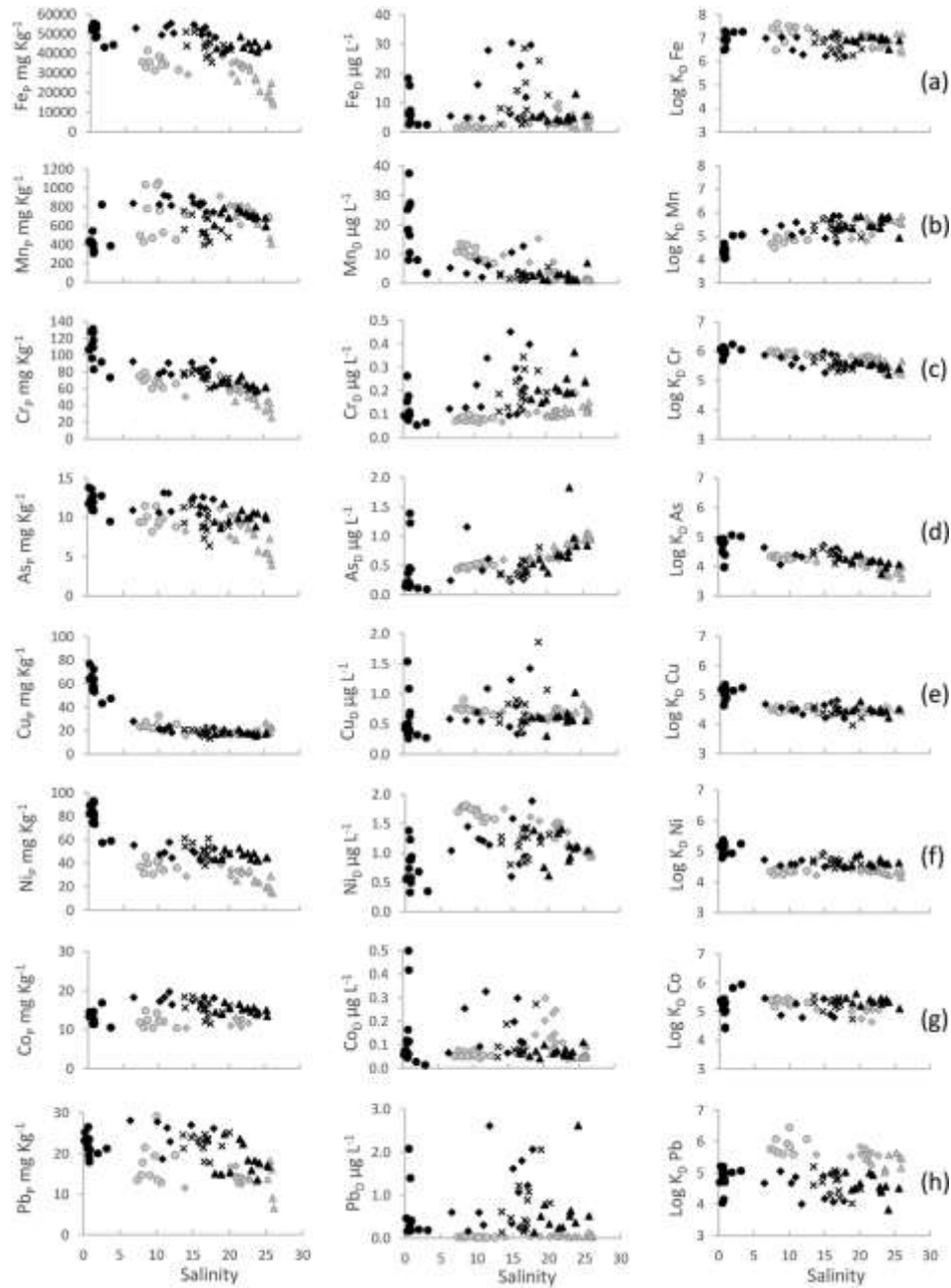


Fig. 4. Distribution of dissolved, particulate, log K_d of metals in the Can Gio mangrove estuary during the dry (gray dots) and the wet seasons (black dots), with circles, squares, multiplication signs and triangles representing the site 1, site 2, site 3 and site 4 respectively: Fe (a), Mn (b), Cr (c), As (d), Cu (e), Ni (f), Co (g) and Pb (h).

between Cr_D and Fe_D ; Fig. 5f), i.e. mangrove sediment pore-water releases (Szymczycha et al., 2016), runoff and/or shrimp pond effluents inputs. Along the estuary, $\log K_D^D$ decreased from 6.2 to 5.2 and presented a low variation range. The combination of Cr_D increase, $\log K_D^D$ decrease from the upstream site to the seaside and the positive correlation between $\log K_D^D$ and POC (Figs. 4c and 5d) confirm the Cr desorption from particulate organic matter during the water transit in the estuary. We suggest that the oxidation level may also influence Cr partitioning as evidenced by the negative correlation observed between $\log K_D^{Cr}$ and $\log DO$ (Fig. 5e). Thus, Cr partitioning in the Can Gio mangrove Estuary was controlled by physical mixing and also biochemical processes, notably OM decomposition and increased oxygenation along the estuary.

3.4.4. Arsenic

The As_P varied from 6.4 to 13.8 $mg\ kg^{-1}$ during the rainy season, and from 3.9 to 11.5 $mg\ kg^{-1}$ during the dry season. Values were stable along the salinity gradient during the rainy season and presented a drop of concentrations at site 4 during the dry season (Fig. 4d). The concentrations at the seaside during the dry season were in the same range than those measured in the coastal zone of South China Sea (Cenci and Martin, 2004) and in the Bohai Sea (Wang et al., 2016). The As_P distribution in the Can Gio mangrove Estuary reflected thus the water mixing between the river and the sea end-members but also biogeochemical processes. Along the estuary, As_P exhibited a strong correlation to Fe_P ($r > 0.8$, for both seasons, Fig. 5g) implying that Fe_P was the main carrier phase of As_P (Strady et al., 2017a) and an important factor controlling As_P .

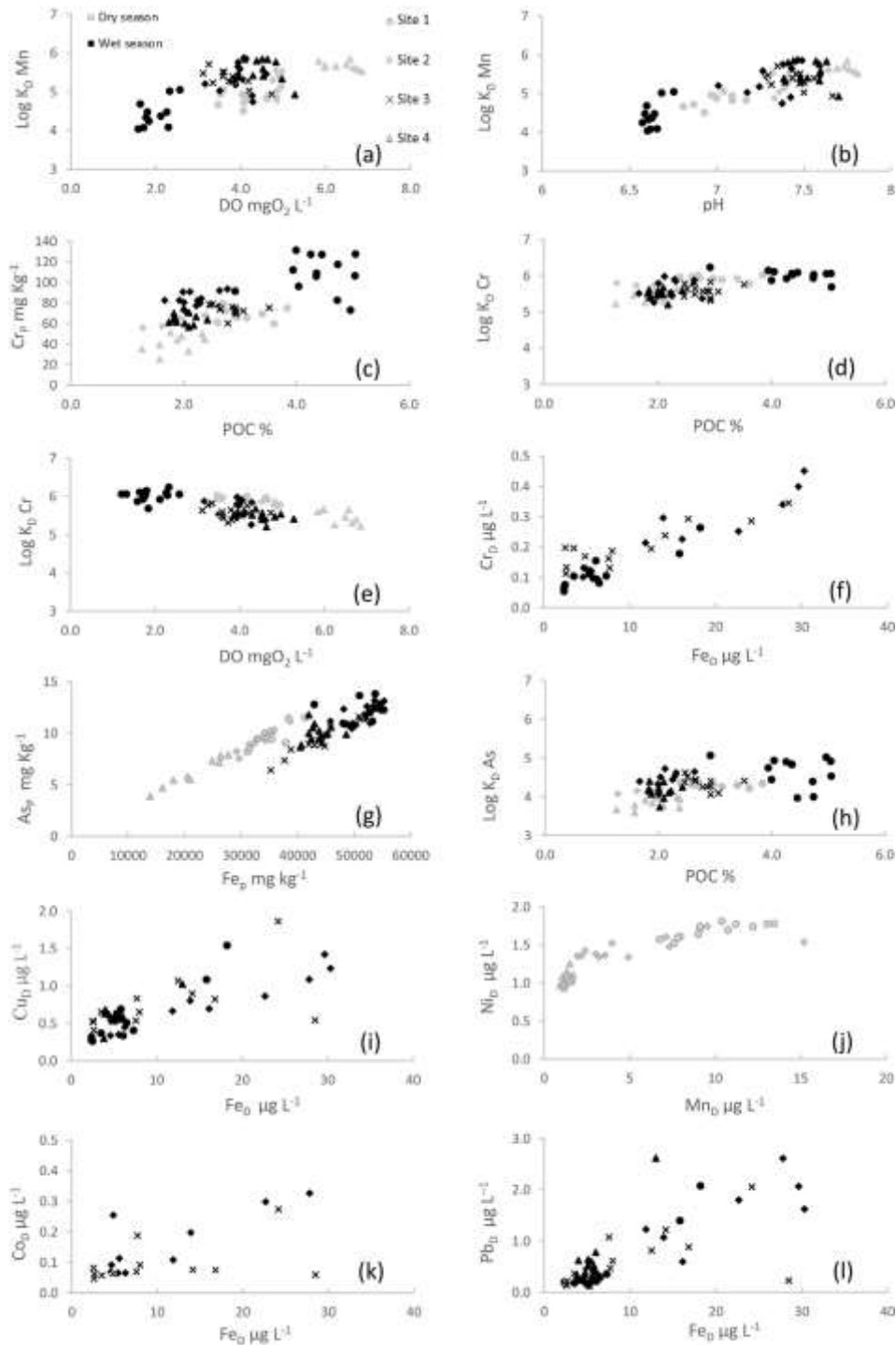


Fig. 5. Interrelations between dissolved metal concentrations and relationships between log K_d of metal with physico-chemical parameters during the dry and the wet seasons: (a) $\text{Log } K_d^{\text{Mn}}$ and DO; (b) $\text{Log } K_d^{\text{Mn}}$ and pH; (c) Cr_D and POC; (d) $\text{Log } K_d^{\text{Cr}}$ and POC; (e) $\text{Log } K_d^{\text{Cr}}$ and DO; (f) Cr_D and Fe_D ; (g) As_D and Fe_D ; (h) $\text{Log } K_d^{\text{As}}$ and POC; (i) Cu_D and Fe_D ; (j) Ni_D and Mn_D ; (k) Co_D and Fe_D ; (l) Pb_D and Fe_D .

distribution in the estuary. Both metals probably had a terrigenous origin (Cances et al., 2005; Oursel et al., 2014). Actually in Vietnam, many soils are naturally enriched in As, which is a public health problem for the local populations (Gustafsson and Tin, 1994; Nguyen et al., 2016; Phuong et al., 2010). Along the estuary, As_D concentrations increased from 0.5 to 0.95 during the dry season and from 0.15 to 0.65 mg L^{-1} during the rainy season, which might be caused by dilution with sea water i.e. 1.5 mg L^{-1} of As_D in the

Bohai Sea (Wang et al., 2016) and probably by As_D release during degradation of organoarsenic compounds from particulate phase (i.e. negative correlations were observed between As_D and POC, $r = -0.64$). This hypothesis is supported by the $\text{Log } K_d^{\text{As}}$ decrease along the salinity gradient, especially from mid salinity (Fig. 4d) and by positively correlations between $\text{Log } K_d^{\text{As}}$ and POC (Fig. 5h). In previous studies, some authors noticed that dissolved organic carbon plays an important role of As partitioning between dissolved and

particulate phase because it can strongly interact with As species (Bauer and Blodau, 2006; Liu and Cai, 2010; Yanan et al., 2017). In the Can Gio mangrove estuary, however, we observed the correlation coefficients between As_D concentrations and DOC being low during the dry season ($r < 0.4$) and very low during the rainy season ($p > .05$). As a consequence, we suggest that the As-DOC complexation may occur as a minor processes and possibly causing $\log K^As_D$ decrease along the estuary. The As_D distributions presented conservative to non-conservative behavior with slightly subtractive concentrations at mid salinity during both seasons (Fig. 4d), which might be caused by the As_D adsorption onto particle phase containing rich (hydr)oxide Fe_p (Lenoble et al., 2013). We also noticed low $\log K^As_D$ values at the upstream site during the ebb tide, which could be related to the Sai Gon River inputs containing high As_D . In conclusion, As partitioning in the Can Gio Estuary was affected by both physical and biogeochemical processes, mainly organic matter decomposition.

3.4.5. Copper

During both seasons, Cu_p varied between 13 and 33 $mg\ kg^{-1}$ (Fig. 4e, supplementary data 1 and 2) and presented stable values in the estuarine part (5e25 salinity), as observed in the East Hainan, China (Fu et al., 2013) and in the Pearl River Estuary, South China (Zhang et al., 2013). Adrop of Cu_p was observed between the riverine part (e.g. 43e77 $mg\ kg^{-1}$, upstream site during the rainy season) and the beginning of the estuary. This loss of Cu_p at the fresh water - estuary interface could be related to intense organic matter degradation processes leading to desorption reaction due to the decomposition of organo-copper complexes; the urban water was characterized by both high POC and Cu_p . The higher $\log K^{Cu_p}$ at the upstream site during the monsoon (Fig. 4e) confirmed those hypothesis. The Cu_p varied from 0.5 to 0.9 and 0.3e1.8 $mg\ L^{-1}$ during the dry season and the monsoon, respectively, with a high variation range and no clear evolution along the salinity gradient. However, punctual elevated Cu_p at site 2 and site 3 were measured during the monsoon (Fig. 4e) and were positively correlated to Fe_D ($r = 0.73$, Fig. 5i) possibly suggesting the same sources as for Fe : inputs from adjacent ecosystems. However, these punctual Cu_p inputs did not seem to affect the Cu partitioning, as evidenced by the stable $\log K^{Cu_p}$ along the estuary (Fig. 4e). These results suggest the biogeochemical processes might be minor factors influencing the Cu partitioning in this environment.

3.4.6. Nickel

The Ni_p distribution along the salinity gradient differed between seasons. During the rainy season, Ni_p varied from 61 to 38 $mg\ kg^{-1}$, with stable concentrations in the estuarine part, which were approximately 2-fold lower than in the riverine part (Fig. 4f, supplementary data 1). At the river end-member, high Ni_p were suggested to be related to the high POC concentrations. Like for Cu, the decrease of Ni_p at the river-estuary interface may result from Ni_p desorption due to organic matter degradation. The Ni_D varied from 0.3 to 1.8 $mg\ L^{-1}$ along the estuary with scattered concentrations. The absences of correlations between Ni_D and physico-chemical parameters or TSS or other metals do not allow us to identify the origin of scattered Ni_D concentrations. $\log K^{Ni_D}$ was stable along the salinity gradient, except a slight decrease at the river-estuary interface (Fig. 4f), inferring that Ni was poorly reactive in the Can Gio mangrove Estuary during the rainy season. During the dry season, Ni_p slightly decreased along the salinity gradient and dramatically dropped to minimum values at the mouth of the estuary, from 45.6 $mg\ kg^{-1}$ to 14.4 $mg\ kg^{-1}$ (Fig. 4f, supplementary data 1). This phenomena might be induced by the marine dilution, the coastal area of South China Sea containing low Ni_p , 12 $mg\ kg^{-1}$ (Cenci and Martin, 2004). The decrease of Ni_p caused by sea water

dilution was previously observed in the Tanshui Estuary, Northern Taiwan (Fang and Lin, 2002) and in the East Hainan, China (Fu et al., 2013). The gradual Ni_D decrease during the dry season (1.81 $mg\ L^{-1}$ to 0.93 $mg\ L^{-1}$) from site 1 to site 4 could be also related to sea water dilution, Ni_D were lower than 0.3 $mg\ L^{-1}$ in the coastal area of South China Sea (Cenci and Martin, 2004). This decrease can also be related to co-precipitation of Ni and Mn onto solid phase (i.e. a positive correlation between Ni_D and Mn_D , $r = 0.88$, Fig. 5j), like previously observed in the Port Curtis Estuary, Australia (Angel et al., 2010) and in Yangtze River Estuary (Wen et al., 2013). During both seasons, the stability of $\log K^{Ni_D}$ suggests that Ni was poorly reactive in the estuary. The physical mixing was thus the main factor controlling the Ni partitioning in the Can Gio mangrove Estuary.

3.4.7. Cobalt and lead

Because of their similar distributions in dissolved, particulate concentrations and partitions along the studied estuary, Co and Pb concentrations are presented and discussed together. The Pb_p presented scattered values close to 21 $mg\ kg^{-1}$ and 15 $mg\ kg^{-1}$ during the monsoon and the dry season respectively, while Co_p were varied around 13 $mg\ kg^{-1}$ for both seasons (Fig. 4g and h, supplementary data 1 and 2). Those scattered concentrations were often observed in tropical estuaries, like in the Wenchang/Wenjiao River Estuary, East-Hainan, China (Fu et al., 2013) and in the Changjiang Estuary, Eastern China (Wang and Liu, 2003). The physico-chemical parameters had a limited effect to Co_p and Pb_p distributions (e.g. no significant correlation with pH, DO or POC), implying that biogeochemical processes have a restricted control on the Co and Pb distribution in the estuary. The Co presented baseline concentration close to 0.05 $mg\ L^{-1}$ during both seasons whereas the Pb_D were closed to 0.3 $mg\ L^{-1}$ and 0.03 $mg\ L^{-1}$ and the dry season, respectively, exhibiting a seasonal effect. The higher measured Pb_D during the rainy season might be attributed to intense leaching from mangrove soil (Defew et al., 2005) and/or to Pb enrichment by atmospheric deposition (Hien et al., 1997, 1999). Along the salinity gradient, both Co_D and Pb_D showed non-conservative additive behaviors (Fig. 4g and h), as previously observed in the Mekong delta (Cenci and Martin, 2004) and in the Changjiang Estuary, Eastern China (Wang and Liu, 2003). Scattered Co_D and Pb_D concentrations were observed at site 2 and site 3 during both flood and ebb tides in the rainy season that suggested similar sources between those metals and Fe_D (significant correlations with Fe_D : $r = 0.56$ for Co_D and $r = 0.77$ for Pb_D , Fig. 5k and l). In the Southern Baltic Sea, a significant pore-water discharge along the coastal area was reported to be a source of dissolved Co and Pb (Szymczycha et al., 2016). $\log K^{Co_D}$ and $\log K^{Pb_D}$ were stable and scattered along the estuary during both seasons but not correlated to any physico-chemical parameters, suggesting a poor reactivity (Fig. 4g and h). The scattered $\log K^{Co_D}$ and $\log K^{Pb_D}$ evidenced thus that the contribution of extra sources played an important role in Co and Pb distributions in the Can Gio Estuary.

4. Conclusions

The Can Gio mangrove Estuary is a dynamic environment at the edge between the biggest City in Vietnam and the South China Sea. The water delivered to the estuary during the rainy season was acidic, almost anoxic, and rich in organic matter and in trace metals, evidencing strong anthropogenic pressure on the ecosystem. We suggest that during the monsoon season, heavy rainfall induced increased runoff and soil leaching, resulting in enhanced trace metal inputs to the estuary, both in particulate and dissolved phases. However, as soon as the trace metals enter the estuary, their distribution and partitioning changed due to the physical mixing

with seawater and/or biogeochemical processes. The first parameter controlling metal dynamics and transport from Ho Chi Minh City to the South China Sea was the concentration of suspended solids, which can vary as a function of tide and season. Strong correlations were measured between total trace metal concentrations and TSS, whatever the element, but they did not explain the whole variability of trace metal dynamics along the estuary. Organic matter played also a key role on these dynamics. First, we suggest that the elevated inputs of Cu_p , Ni_p and Cr_p were related to the high POC concentrations at the upstream site during the monsoon season. Then, OM decomposition along the estuary resulted in metal release in the dissolved phase, with increasing concentrations with the salinity gradient, specifically for As and Cr. Conversely, concentrations of other dissolved elements decreased along the salinity gradient, either due to water mass mixing, or geochemical processes, which was the case for Mn, Fe, Co and Pb which were poorly reactive during their transit, as evidenced by the absence of any specific correlation between their $\log K_D$ and the physico-chemical parameters studied. For the latter metals, water mass mixing was suggested to be the main factor driving their distribution along this mangrove Estuary. Regarding the potential impacts of extra sources on metal partitioning like: i) pore-waters flushed out from mangrove soils at low tide, ii) runoff from adjacent soils, or iii) shrimp pond effluents, a further detailed investigation should be carried out at sites without freshwater discharge to get a better assessment of their role in trace metals dynamic along the estuary. Furthermore, trace metal geochemistry should be studied in the whole ecosystem (e.g. mangrove soils, trees, animals) and not only in the water column, to understand their potentially bioavailability and ecologically risks.

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Appendix A. Supplementary data

Supplementary data related to this article can be found at <https://doi.org/10.1016/j.chemosphere.2017.12.189>.

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