Adsorption and fractionation of Pt, Pd and Rh onto inorganic microparticles and the effects of macromolecular organic compounds in seawater

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A R T I C L E   I N F O

Article history:
Received 30 October 2018
Received in revised form 5 September 2019
Accepted 5 September 2019
Available online 10 September 2019

Keywords:
Adsorption
Fractionation
Inorganic microparticles
Macromolecular organic compounds
Platinum group elements
Seawater

A B S T R A C T

Adsorption and fractionation of Pt, Pd and Rh (defined here as platinum group elements, PGEs) onto the representative inorganic microparticles, including Fe2O3, MnO2, CaCO3, SiO2, Al2O3 and kaolinite in seawater were investigated. The effects of macromolecular organic compounds (MOCs) as the representatives of organic matter, including humic acids (HA), bovine serum albumin (BSA) and carrageenan, on the adsorption were also studied considering that organic matter is ubiquitous in seawater and indispensable to marine biogeochemical cycles. In the absence of MOCs, the representative mineral particles Fe2O3 and MnO2 had the strongest interaction with PGEs. The adsorption of PGEs onto the representative biogenic particles SiO2 and CaCO3 and lithogenic particles Al2O3 and kaolinite was similar or weaker than onto the mineral particles. MOCs inhibited the interaction between PGEs and the particles except for Pt and Pd onto the biogenic particles in artificial seawater. This impediment may be closely related to the interaction between particles, MOCs and elements. The partition coefficient (log Kd) of Pt was similar (~4.0) in the presence of MOCs, indicating that the complexation between Pt and MOCs was less important than hydrolysis or adsorption onto the acid oxide particle surface. Rh tended to fractionate onto the mineral and lithogenic particles in the presence of HA and carrageenan, while Pd was more likely to fractionate onto the biogenic particles. However, BSA enhanced the fractionation tendency of Pd onto the mineral particles. The results indicate that the adsorption behavior of Pd onto inorganic particles was significantly affected by the composition or the type of MOCs. Hence, the interaction between PGEs and inorganic particles may be greatly affected by the macromolecular organic matter in the ocean.

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1. Introduction

There has been a huge increase in three anthropogenically derived platinum group elements (PGEs) in natural environments, namely Pt, Pd and Rh, because of their wide application in anticancer drugs, chemical catalysis, and especially automotive catalytic converters (Koek et al., 2010; Mudd, 2012; Pawlak et al., 2014; Speder et al., 2014). Owing to the potential hazards of these pollutants to humans (Merget and Rosner, 2001; Wiseman and Zereini, 2009; Sorensen et al., 2016; Zimmermann et al., 2017), researchers have investigated the biogeochemical behaviors of PGEs in various environmental matrixes (Reith et al., 2014). Furthermore, a migration path of these elements in the environment, vehicle exhaust—soil—aquatic system, was preliminarily defined (Mitra and Sen, 2017). Tiny particulate PGEs have been released from vehicles into the atmosphere in the abrasion and deterioration of catalysts and subsequently enter the soil with atmospheric deposition and rainfall (Colombo et al., 2008; Wiseman and Zereini, 2009; Pawlak et al., 2014; Clement et al., 2015; Ruchter and Sures, 2015). Later, PGEs in their elemental states are transformed
into oxidation states with the compounds present in soil (Ek et al., 2004; Sebek et al., 2011; Sucha et al., 2016). Finally, PGEs enter the aquatic environments such as rivers, estuaries and seawater with runoff and rainfall (Pawlak et al., 2014; Ruchter and Sures, 2015; Mitra and Sen., 2017). Moreover, PGEs have the characteristics of forming complexes easily with donor ligands such as suspended particulate matter (SPM) and dissolved organic matter (DOM) present in the natural aquatic system (Baker and Spencer, 2004; Turner et al., 2006; Turner and Wu, 2007). Hence, biogeochemical behaviors of PGEs in aquatic systems could dramatically affect the status of these elements in the environment. However, there is a lack of the information about the interactions between PGEs and SPM or DOM, because the detection of PGEs is restricted at typical environmental levels (Rao and Reddi, 2000; Suzuki et al., 2014; Formies et al., 2016; Mashio et al., 2016).

Some researchers have explored the role of SPM or sediment in the scavenging of PGEs in aquatic systems. Turner et al. (2006) reported that the adsorption of PGEs onto suspended estuarine sediment was dominated by the hydrous metal oxides of the par-

mineral fraction was represented by Fe₂O₃ and MnO₂ (Yang et al., 2013; Lin et al., 2014; Lin et al., 2015; Yang et al., 2015). Comparative experiments were conducted in the presence of diverse DOM for the investigation of the role of organic matter in regulating this adsorption in seawater. The organic matter used in this study was some model MOCs, namely humic acid (HA), bovine serum albumin (BSA) and carrageenan (Sigma-Aldrich® Product Number C1867), which represented the known major substances of natural DOM including humic matter, protein and acid polysaccharides, respectiv-

elatively (Gueguen et al., 2006; Yang et al., 2013; Ilina et al., 2014; Lin et al., 2014; Lin et al., 2015; Yang et al., 2015). This study focused on the role of the chemical components of particles and DOM in the fractionation of PGEs in seawater; the effects of living organisms and the precipitation of PGEs were not taken into account; in addition, the electrical potential of the solutions was simply regarded as qualitative evidence and not taken into deep discussion.

2. Materials and methods

2.1. Reagents and seawater

The microparticles and MOCs selected for the adsorption experiment are representatives of the major known particulate and DOM components in seawater (Yang et al., 2013; Lin et al., 2015). The characteristics of MOCs including the main constituents, functional groups and molecular weights are shown in Table 1; the details of microparticles including the surface area and zeta potential are shown in Table 2. The ultrapure deionized water (DIW, >18 MΩ·cm) used in this study was prepared with a Pall® Cascad™ water purification system.

The mixed stock solution of 1 mg L⁻¹ each of Pt, Pd and Rh was prepared in 2 mol L⁻¹ HCl in a 1 L high-density polyethylene bottle (Thermo®) with a standard of 1000 mg L⁻¹ individual elements of Pt, Pd and Rh supplied by General Research Institute for Nonferrous Metals, China. The chemical forms of Pt, Pd and Rh in the stock solution were Cl-forms and the oxidation states were +2, +2 and +3, respectively. The labwares used in this study were cleaned strictly according to Li et al. (2015) and Liu et al. (2019). They were washed with Decon® 90 (2% in DIW), then infused in 10% HNO₃ and 10% HCl for 48 h, followed by rinsing with DIW for more than five times, and finally dried in a Class 100 clean bench.

For comparison, two solution systems were applied, one prepared with artificial seawater (ASW) which did not contain organic

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| Table 1 | Characteristics of macromolecular organic components used in this research. |
|---------|--------------------------------------------------------------------------------|
| Macromolecular organic matter | Represented component of DOM | Main functional group | Molecular weight | Provider |
| Humic acid sodium salt | Humic matter | -COO⁻ | >10 kDa | Sigma-Aldrich China, Inc. |
| Bovine serum albumin | Protein | Amino acid residue | -66.4 kDa | Shanghai Luoshen Biological Technology Co. Ltd |
| Mixture of κ- and λ-carrageenans | Acid polysaccharides | -SO₃, -OH | >20 kDa | Sigma-Aldrich China, Inc. |

| Table 2 | Characteristics of inorganic microparticles used in this research. |
|---------|----------------------------------------------------------------------|
| Particle | Median grain size (µm) | Specific surface area (m² g⁻¹) | Zeta potential in ASW (mV) | Zeta potential in UVSW (mV) | Provider |
| Fe₂O₃ | 7.1 | 23.8 | -6.82 | 2.16 | Sinopharm Chemical Reagent Co., Ltd |
| MnO₂ | 11.6 | 18.9 | -5.47 | -3.53 | Sinopharm Chemical Reagent Co., Ltd |
| CaCO₃ | 11.1 | 2.56 | 14.50 | 10.80 | Sinopharm Chemical Reagent Co., Ltd |
| SiO₂ | 2.6 | 2.12 | -15.60 | -8.10 | Sinopharm Chemical Reagent Co., Ltd |
| Al₂O₃ | 8.0 | 21.3 | -6.78 | -4.64 | Sinopharm Chemical Reagent Co., Ltd |
| Kaolinite | 9.5 | 35.7 | -5.78 | -3.45 | Sinopharm Chemical Reagent Co., Ltd |
mixture and nutrients, and the other prepared with ultraviolet irradiated natural seawater (UVSW) from which DOM was removed. ASW was prepared according to the method D1141-98, Reapproved 2013 (American Society for Testing and Materials, ASTM). UVSW was prepared by irradiating natural seawater under ultraviolet light (500 W) for over 48 h to remove most of the active DOM, and the residual background concentration of dissolved organic carbon in UVSW was less than 0.2 mg L\(^{-1}\). The natural seawater was collected from the North Yellow Sea (123.912° E, 38.965° N), filtered through a 0.2 μm hydrophilic polyethersulfone membrane (Pall®) and stored in 4 L polypropylene (PP) bottle (Thermo®) at 4 °C for further use.

2.2. Adsorption studies

The adsorption experiment was operated in 100 ml Teflon beakers with lid. Teflon was reported to have low surface adsorption of PGEs ions (Cubelo-García et al., 2008). The adsorption procedures are schematized in Fig. 1. The experimental preparations were performed in the clean bench. The adsorption was conducted at room temperature (20 ± 2 °C) in the dark under two conditions: single kind of particle system and particle-MOC system. The following is the brief account of the experimental procedure. 48.5 ml of ASW or UVSW were added into 100 ml Teflon beakers, and then each was spiked in 1 ml non-complexing Tris-HCl buffer (pH ~8.5). The buffer could maintain the pH of the experimental solution at ~8.2 and prevent the irreversible formation of interfering colloids, caused by pH drift, which could significantly influence the adsorption of PGEs onto particles (Roberts et al., 2009; Yang et al., 2013; Li et al., 2015).

For a single kind of microparticle system, 2.5 ± 0.05 mg of particles was added into each of the 100 ml Teflon beakers. The final concentration of particles was 50 mg L\(^{-1}\), which was close to the general SPM concentration (1–100 mg L\(^{-1}\)) in natural seawater (Yang et al., 2013). Then, 0.5 ml of PGEs stock solution was added to make the work concentration of each Pt, Pd, and Rh at 10 μg L\(^{-1}\). The final experimental solution volume was 50 ml. Notably, 10 μg L\(^{-1}\) of PGEs was dramatically higher than the natural concentration of these elements in the ocean. Nevertheless, this concentration was comparable to that of some other trace elements in seawater and lower enough compared with the concentration of ligands that could potentially bind with PGEs (Turner et al., 2006; Turner and Wu, 2007). Hence, the experimental condition could, at least to some extent, reflect the adsorption situation of PGEs in seawater (Turner et al., 2006; Turner and Wu, 2007). Furthermore, this concentration is necessary to allow for an interference-free detection by inductively coupled plasma mass spectrometry (ICP-MS) (Turner et al., 2006). Based on the results of pre-experiments (Fig. S1), the solutions were shaken for about 12 h in the dark for adsorption equilibrium. After the adsorption, the particles and the work solution were separated through filtration with a 0.45 μm pre-weighted hydrophilic polyethersulfone membrane (Pall®). The filtration device was pretreated with 10 ml of work solution three times to reduce the retention of PGEs (Turner et al., 2006; Turner and Wu, 2007). 5 ml of filtrates were pipetted into 15 ml pre-weighted polypropylene (PP, FCOLONTM) centrifuge tubes, diluted to 10 ml by 2% HCl and stored at 4 °C. The particles retained on the membranes were dried in the clean bench for 24 h and weighted to calculate the mass balance of the particles. The weighted membranes were soaked in 10 ml of 2.5 M HCl in 15 ml PP centrifuge tubes at room temperature to be digested for one week. After the digestion, the tubes were centrifuged at 2500 rpm for 5 min. And then, 5 ml of supernatants were pipetted to 15 ml PP centrifuge tubes and stored at 4 °C.

The role of MOCs in the fractionation of PGEs onto inorganic microparticles was investigated with the binary experiments in which the microparticle and MOCs were simultaneously added into the adsorption solutions and stirred for 30 min before spiking PGEs, guaranteeing the sufficient interaction between MOCs and microparticles (Yang et al., 2013). The final concentration of MOCs was 2 mg L\(^{-1}\), which was in the range of natural DOC concentrations of 1.09–4.22 mg L\(^{-1}\) in the Bohai and Yellow Seas (Ran et al., 2013; Liu et al., 2015). Other procedures were the same as the single kind of particle system. The control experiments were conducted simultaneously in which the same concentrations of PGEs as that of the single kind of particle and particle-MOC experiments were spiked but no microparticles and MOCs were added.

The pH throughout the adsorption experiments was monitored with a Meteler Toledo® pH meter, because Cubelo-García et al. (2008) reported that significant changes in pH may lead to undesirable colloid formation, particle dissolution, and precipitation of PGEs in the solution, which interfere in the adsorption dramatically, and the sample must be discarded if the pH of the experimental system drifted more than ~0.3 pH units. Three repeated operations were conducted simultaneously for each adsorption process.

2.3. Sample analysis

The concentration of Pt, Pd and Rh in filtrates and digestions was measured using an ICP-MS instrument. The precision was ± 3% and the detection limit was 0.01 mg L\(^{-1}\) for Pt, 0.005 mg L\(^{-1}\) for Pd, and 0.001 mg L\(^{-1}\) for Rh. All additions and subtractions of the mass balance were performed with two replicates. The pH throughout the adsorption experiments was monitored with a Meteler Toledo® pH meter.
analyzed with ICP-MS (ELAN DRC II, PerkinElmer®). The filtrates were diluted 25-fold in 2% HNO3 to maintain the concentration of dissolved salt being <0.1% w/v to minimize signal interference from the saline matrix. The digestions were diluted to twice their original volumes with DIW. The instrument was calibrated using 2% HNO3-based multi-element standards which were prepared with UVSW and ASW, and \(^{103}\text{Rh}\) was added as the internal standard to make a concentration of 10 mg L\(^{-1}\). The concentrations of Pt, Pd and Rh were determined with the signal of \(^{103}\text{Rh}\), the average signals of \(^{194}\text{Pt}\) and \(^{199}\text{Pt}\), and \(^{106}\text{Pd}\) and \(^{108}\text{Pd}\), respectively (Djigova et al., 2003). A standard and a blank were inserted in every analysis of the 12 samples to monitor the instrument errors.

The zeta potential of the solutions was determined with a Malvern Panalytical® Nano-WS90 analyzer. The surface morphology of the particles was obtained with a Hitachi® S-4800 high-resolution scanning electron microscope (SEM). The work conditions of the microscope were 3 kV accelerating voltage, 2,000–100,000 x magnification, 3800–4200 μm working distance and 6,000–10,000 nA emission current. In this study, the images with a magnification of 50,000 x were selected for the analysis, which clearly reflected the surface conditions of the particles.

2.4. Recovery, partition coefficient, and fractionation factor

The mass recovery was used to measure the adsorption of PGEs onto the container walls and the experimental operation on the adsorption results (Yang et al., 2013). The total recovery was defined as R to evaluate the loss of PGEs during the experiment:

\[
R = \frac{M_\text{ads} + M_\text{p}}{M_\text{i}} \times 100\%
\]

where R represents the ratio of the total mass recovery of PGEs before and after the adsorption; \(M_\text{ads}\) represents the mass of PGEs in the liquid phase after adsorption; \(M_\text{p}\) represents the mass of PGEs in the particulate phase after adsorption; \(M_\text{i}\) represents the initial mass of PGEs added to the solution. When \(R = 100\%\), there is no loss of PGEs in the experiment; when \(R < 100\%\), there is some loss; the greater the difference between \(R\) and 100% is, the greater loss occurs in the experiment.

The partition coefficient (Kd) of PGEs between seawater and particulate phases was calculated with a traditional formulation which has been used in different experimental systems (Honeyman and Santschi, 1989; Yang et al., 2013):

\[
K_d = \frac{C_p}{C_d C_T}
\]

where \(C_p\) represents the concentrations of PGEs in the particulate phase; \(C_d\) is the concentrations of PGEs in the dissolved phase; \(C_T\) is the total concentrations of microparticles in the solution. The Kd values (in L kg\(^{-1}\)) are reported in terms of log Kd.

The fractionation factor (F) was defined as the ratio of Kd values to evaluate the fractionation behavior of those elements by a specific microparticle (Yang et al., 2013; Lin et al., 2015):

\[
F_{A/B} = \frac{K_{dA}}{K_{dB}}
\]

where \(K_{dA}\) and \(K_{dB}\) are the partition coefficients of PGE A and B, respectively; \(F_{A/B}\) is the fractionation factor between A and B. When \(F_{A/B} = 1\), there is no fractionation in that particular particle during adsorption. When the \(F_{A/B} > 1\) or <1, the particle preferentially absorbs PGE A over B or B over A.

3. Results

3.1. The recovery of Pt, Pd and Rh

The R value of each PGE was measured for the evaluation of its loss in the experiments, and the results under different experiment conditions are shown in Fig. 2. Overall, R ranged from 80 to 100% and varied with different elements, which means that experimental design and operation satisfied the objectives of this research.

The R values of Pt in all conditions were higher than 90%, indicating that no significant loss of Pt occurred during the adsorption experiment. For Pd, a different feature of R was found with the addition of MOCs in ASW (Fig. 2). In the absence of MOCs, most of the R values of Pd were ~95%. But a slight increase about 2–3% was observed with the addition of MOCs. It may be due to the impeding effects of MOCs on the adsorption of Pd onto the container walls (Cobelo-Garcia et al., 2008). The organic complexation formed between MOCs and Pd had a stronger solvability than the inorganic complexes of Pd (Wood, 1996; Colombo et al., 2008).

For Rh, a wider range of R value about 80–95% was observed and varied with the experimental conditions (Fig. 2). It indicated that relatively lower R values were generally obtained in the presence of MOCs, which might be caused by the Rh adsorption onto container walls, the precipitation of Rh complexes (Cobelo-Garcia et al., 2008; Colombo et al., 2008) and other unknown factors during the experimental operation.

3.2. Partition coefficient of Pt, Pd and Rh

The partition coefficient of Pt, Pd and Rh could reflect an intuitive adsorption behavior of PGEs onto the particles. In this study, the log Kd varied with the types of organic matter, element and particles, and the results are shown in Table S1 and Fig. 3.

The strongest adsorption occurred in the interaction between PGEs and MnO\(_2\) in UVSW, with the log Kd value being greater than 5.7, followed by FeO\(_2\) being approximately 5.6. The lowest adsorption occurred on the Al\(_2\)O\(_3\) particles and the log Kd was about 3.89 ± 0.02 (Fig. 3 and Table S1). The interactions between the studied PGEs and SiO\(_2\), CaCO\(_3\) and kaolinite particles were weaker than mineral particles and the log Kd values ranged from 4.11 to 4.57. Moreover, the fractionation of PGEs onto different particles in ASW and UVSW were diverse.

For Pt, the log Kd on each particle in ASW and UVSW was similar. However, a slight decrease in the coefficient value by ~0.04–0.15 was observed in ASW compared to in UVSW except for SiO\(_2\) with ~0.07 increase in log Kd. For Pd, a larger change of log Kd by 0.06–0.33 appeared. In UVSW, the adsorption of Pd onto Fe/Mn oxides was weaker than in ASW (log Kd decreased by ~0.32), but its adsorption onto biogenic particles were stronger (log Kd increased by ~0.22); besides, the adsorption of Pd onto lithogenic particles were comparable in ASW and UVSW (log Kd changed by ~0.05). For Rh, an obvious difference in its adsorption onto Fe/Mn oxides in ASW and UVSW was observed and the variation of log Kd was about 0.38–0.40; however, its adsorption onto lithogenic and biogenic particles differed slightly in the two seawater conditions expect for Al\(_2\)O\(_3\), with the log Kd increased ~0.16 in UVSW. From the above results, it could be obtained that the variation in adsorption reactivity of PGEs onto particles might be related to the residual dissolved organic matter in UVSW, which was the main difference between UVSW and ASW.

In the particle-MOC binary systems, in general, MOCs weakened the adsorption of PGEs onto particles (Fig. 3), and the interactions among them varied greatly. All comparisons in the following were made with the corresponding particle system.

For Rh, the strong adsorption still occurred on the Fe/Mn oxides.
The log $K_d$ of these binary systems decreased by about 0.47–1.00 and tended to have very similar values of 4.79 ± 0.06 except for the ones with carrageenan in ASW. For the adsorption of Rh onto biogenic particles, MOCs hindered its interaction with CaCO$_3$ in ASW, and log $K_d$ decreased correspondingly by 0.38–0.4; however, they strengthened the adsorption in UVSW and the maximum change of log $K_d$, about 0.10–0.25, occurred in the systems containing HA. Moreover, the promoting effects of carrageenan and BSA on the distribution of Rh onto CaCO$_3$ particles were similar (Fig. 3). For the interaction between PGEs and SiO$_2$, HA promoted the adsorption, BSA hindered the adsorption and carrageenan had no effects on the adsorption in ASW which showed the promoting effects and the log $K_d$ increased by 0.10–0.14. Among them, the strongest inhibition effect was on mineral particles (Fig. 3), with the decrease in log $K_d$ by 0.40–1.40. For Pt, the weakening effects of MOCs on the interactions between particles and PGEs were also observed, and the biggest decrease in log $K_d$ by 0.47–1.03 was also on the mineral particles (Fig. 3 and Table S1). The log $K_d$ of lithogenic and biogenic particles was 3.91–4.38 in the presence of MOCs and the fluctuation was no more than 0.2–0.4 (Table S1). The log $K_d$ of the adsorption of Pt onto Fe/Mn oxides in ASW was higher than that in UVSW, and slight promoting effects on the adsorption of Pt onto SiO$_2$ in ASW were observed (Fig. 3).

In general, MOCs changed to a certain extent the adsorption of PGEs onto inorganic particles, and this effect was closely related to the type of particles and organic matter. This suggested that MOCs had a profound influence on the partition of Rh, Pd and Pt between the particles and the seawater.

### 3.3. Fractionation factors between Pt, Pd and Rh

Fractionation factors ($F$) between Pt, Pd and Rh which were derived from their $K_d$ values and expressed by the pairs of Rh–Pd ($F_{Rh/Pd}$), Rh–Pt ($F_{Rh/Pt}$) and Pd–Pt ($F_{Pd/Pt}$) are shown in Fig. 4 and
Table S2. This parameter reflects the adsorption inclination of particles to elements in different environments. The more the F value deviated from 1, the more obvious fractionation of the PGEs onto the tested particles was.

In a single kind of particle system, F of PGEs adsorption onto MnO₂ had the largest value and was obviously higher in UVSW than in ASW except for Pt-Pd. It indicated that the fractionation between Pt and Pd was not significantly affected by the solution matrix. In addition, the difference in F values of PGEs adsorption onto other particles were not significant. In general, the fractionation of PGEs onto Fe₂O₃ and MnO₂ in ASW and UVSW showed a big difference (Table S2).

In the particle-MOC binary systems, MOCs dramatically changed the fractionation between PGEs, and a significant difference between the two seawater conditions was also observed. For Fe₂O₃, F increased in the presence of HA, carrageenan and BSA in ASW; but in UVSW, the effects of MOCs varied: HA and carrageenan enhanced (F increased by 10^3–10^4) while BSA inhibited (F decreased by 8) the fractionation between Rh and Pd. The enhancing effect in ASW was significantly stronger than in UVSW. For MnO₂, the effect of MOCs on the fractionation between PGEs in UVSW was similar to that of Fe₂O₃; in ASW, HA and BSA slightly enhanced the fractionation between PGEs (F increased by 0.15–2.5) but carrageenan weakened it (F decreased by about 1).

For biogenic particles, the effect of MOCs in ASW and UVSW also varied significantly. MOCs enhanced the fractionation between PGEs in the process of their adsorption onto CaCO₃ in ASW; however, this effect only occurred in the presence of BSA in UVSW. The influence of MOCs on the fractionation between PGEs in the process of their adsorption onto SiO₂ was more complicated: in ASW, HA and carrageenan made a slight enhancement on the fractionation but BSA had no effects; in UVSW, conversely, BSA had an obvious enhancement but carrageenan and HA had almost no effects.

For lithogenic particles, MOCs had a similar effect on the fractionation between PGEs in ASW and UVSW with HA and carrageenan promoting the fractionation but BSA having no obvious effects. The fractionation between the pairs of Rh-Pd and Pd-Pt was promoted more in UVSW than in ASW.

The results revealed that, on the whole, MOCs impacted especially on the fractionation of PGEs in adsorbing onto Fe₂O₃ and MnO₂. The highest F about 43.65 was observed in Fe₂O₃-HA system and followed by that in MnO₂-HA system about 25.11 (Fig. 4). It indicated a very important role of HA in regulating PGEs adsorption onto mineral particles in seawater. For other MOCs, the enhancement of BSA for the adsorption of PGEs onto the particles was higher than that of carrageenan. Hence, the ability of MOCs to affect the adsorption of PGEs onto mineral particles was HA > BSA > carrageenan. But for lithogenic particles, the order changed to be HA > BSA ≈ carrageenan. For biogenic particles, there were different behaviors in the presence of MOCs. The adsorption of PGEs onto CaCO₃ was more influenced by MOCs compared with their adsorption onto SiO₂. The ability of MOCs to affect the adsorption of PGEs onto CaCO₃ and SiO₂ was
4. Discussion

4.1. Adsorption characteristics of Pt, Pd and Rh onto the microparticles

Based on the results of adsorption coefficient and fractionation factor, it can be found that the adsorption behaviors of PGEs onto particles varied significantly. The mineral particles showed the maximum adsorption propensity (with the log $K_d$ being $4.54$–$5.75$) and fractionation for PGEs, more than other particles used in this study (Figs. 3 and 4). $\text{Fe}_2\text{O}_3$ and $\text{MnO}_2$ exhibited similar behavior of preferring to adsorb Rh and it was followed by Pt and Pd. It indicated that the metal oxides/hydroxides particles had a higher affinity to PGEs ions in seawater. This result is consistent with the contents of PGEs in ferromanganese crusts/nodules in the ocean, which had a high level of PGEs (Sun et al., 2006; Guan et al., 2017). The oceanic ferromanganese crusts/nodules show strong enrichment in Pt and Rh and depletion in Pd (Guan et al., 2017), which was consistent with the strong fractionation of Rh-Pd and Rh-Pt that was generated in the process of their adsorption onto Fe/Mn oxides in this research. However, the fractionation between Pt-Pd was not as obvious as that in the natural oceanic ferromanganese crusts/nodules (Guan et al., 2017), which may be caused by the difference in residual dissolved organic matter between UVSW and ASW. To identify this speculation, the zeta potential of each kind of particle in UVSW and ASW was investigated, and the results are shown in Table 2. The absolute value of Fe/Mn oxides in UVSW was lower than in ASW, which means that these particles in UVSW had a stronger affinity with the negatively charged PGEs complexes. Hence, Fe/Mn oxides play a more important role in the scavenging of Pt, Pd and Rh in the ocean than other inorganic particles.

The affinity of biogenic particles with PGEs was weaker than that of the mineral particles (Figs. 3 and 4). And the adsorption behaviors of Pt, Pd and Rh onto $\text{SiO}_2$ and $\text{CaCO}_3$ particles were similar, with the log $K_d$ being around $4.11$–$4.47$ (Table S1). However, some subtle differences could still be noticed. In ASW, the...
difference in log $K_d$ between PGEs was so small that it could be neglected; while in UVSW, the corresponding log $K_d$ values were higher than in ASW and slightly decreased in the following order: Pd > Rh > Pt, indicating that the residual dissolved organic matter in UVSW may have some effects on the promotion of the adsorption. There was no obvious fractionation between PGEs in the process of their adsorption onto biogenic particles (Fig. 4). As shown in Table 2, the absolute value of zeta potential in SiO$_2$ and CaCO$_3$ particle systems (around 10 mV) was higher than that in Fe$_2$O$_3$ and MnO$_2$ particle systems (around 2–3 mV), indicating that the interaction between PGEs and biogenic particles could be more stable. Moreover, the difference in zeta potential between SiO$_2$ and CaCO$_3$ particle systems was not obvious (Table 2), which may demonstrate the inconspicuous differences in the adsorption and fractionation behaviors of PGEs observed between the two biogenic particles. Other studies also found that SiO$_2$ exhibited similar adsorption behaviors to Rh and Pt ions without being affected by organic matter (Wood, 1996; Takahashi et al., 1999). If this is the case, in the ocean, the fractionation between Pt, Pd and Rh in the process of their adsorption onto biogenic particles would depend largely on the abundance of diatom frustules and the predominant phytoplankton species (Yang et al., 2013). But in reality, the surface of biogenic particles in the ocean is attached by some organic matter such as humic substances, proteins and carbohydrates. These attachments may change the surface properties of the particles, affecting the adsorption of Pt, Pd and Rh onto these particles. In this study, the difference in the results of the single kind of particle experiments and particle-MOC experiments provided a clue to this speculation (Figs. 3 and 4).

Due to their large surface area and ubiquity in marine environments, lithogenic particles may be an important carrier for PGEs. Unlike that of biogenic particles, Al$_2$O$_3$ and kaolinite did not cause obvious fractionation of PGEs in the adsorption process (Fig. 4). From Table 2, it could be found that kaolinite used in this study has the largest specific surface area. For the two kinds of lithogenic particles used in this study, $K_d$ of Rh, Pd and Pt adsorption onto kaolinite was obviously higher than that onto Al$_2$O$_3$ (Fig. 3). However, there was no obvious difference in zeta potential between kaolinite and Al$_2$O$_3$ in both UVSW and ASW (Table 2). Therefore, specific surface area might be an important factor in the adsorption of PGEs onto lithogenic particles in seawater.

4.2. The roles of macromolecular organic matter

4.2.1. Humic acid

HA, a mixture which has many types of functional groups, is usually regarded as an important component of colloidal organic matter (Xu and Guo, 2017). It can greatly affect the geochemical behavior of trace metals in seawater by providing electron donating groups to their ions (Wood, 1996). In this study, an obvious decrease in the adsorption of PGEs onto Fe/Mn oxides was observed in the presence of HA (Fig. 3 and Table S1). Pd had the greatest decrease, followed by Pt and Rh. This may be related to the adsorption of HA onto the surface of particles and the speciation dynamics of HA and PGEs in seawater (Parfitt et al., 1977; Wood, 1996; Lyubomirova and Djingova, 2011).

As shown in Fig. 5, compared to a single kind of particle system, something obviously attached to the surface of Fe$_2$O$_3$ particles in the Fe$_2$O$_3$-HA binary system, which should be HA considering there was no other matter. From the image it could be found that HA was bulky and occupied a large portion of the surface area of the particles. Hence, humic acid, mainly with aromatic ring structures (Wood, 1996; Thimsen and Keil, 1998; Vermeer and Koopal, 1998; Kleber et al., 2007), blocked the available surface of Fe/Mn oxide particles (Takahashi et al., 1999; Tombacz et al., 2004; Qin et al., 2015), and weakened the adsorption of Rh, Pd and Pt onto the particles. In addition, irrespective of particles, there were complex interactions between PGEs and HA in seawater (Liu et al., 2019), which implies that HA competes with particles for adsorbing the ions of PGEs in seawater.

The zeta potential change of particles may be another factor of...
influencing the adsorption. The solutions containing Fe₂O₃, CaCO₃ and Al₂O₃ were selected for the investigation of the relationship between potential change and adsorption in this study. These particles have some special adsorption characteristics such as the adsorption of PGEs onto Fe₂O₃ exceeds MnO₂, the adsorption of PGEs onto CaCO₃ varies significantly, and the adsorption occurring onto Al₂O₃ changes slightly. Therefore, to test the potential change of those particles will be helpful for the understanding of the mechanism of the interactions between particles, organic matter and PGEs. As shown in Fig. 6, the zeta potential for Fe₂O₃ changed from 2.16 mV in the single kind of particle system to −10.8 mV in the particle-MOC binary systems in UVSW. Moreover, it was reported that the potential of inorganic PGEs complexes in natural seawater was negative (Colombo et al., 2008), resulting in the decline of their adsorption onto particles in Fe₂O₃-MOC binary systems (Wood, 1996; Takahashi et al., 1999).

On the whole, when HA was added, a slight decrease in the adsorption of PGEs onto CaCO₃ occurred but a slight increase in their adsorption onto SiO₂ occurred in ASW, while the corresponding variations were relatively weaker in UVSW, especially for Rh and Pt. This may be related to the difference in the affinity between HA and particles and the dominated adsorption factors between PGEs and particles. As shown in Fig. 5, an obvious HA attachment on CaCO₃ particles could be found. However, although a certain portion of the surface area of CaCO₃ particles was occupied by HA, their surface was still dominated by the unoccupied portions. Hence, the adsorption of PGEs onto CaCO₃ in this study could be mainly determined by the hydrophobicity and surface area as that reported in the research of Takahashi et al. (1999), even if the zeta potential of this particle changed from 10.8 mV to −8.27 mV in the presence of HA (Fig. 6). In addition, the relatively low adsorption affinity of PGEs onto biogenic particles may be another factor (Fig. 4 and Table S2), which means that the amount of PGEs in this study could be sufficient to support the adsorption by both HA and biogenic particles.

The adsorption characteristics of PGEs onto the lithogenic particles showed a similar contradictory trend with that of the biogenic particles but varied between elements. For Rh and Pt, HA promoted their distribution onto Al₂O₃ particles, however, it showed an inhibitory effect for Pd, which may be caused by the interaction between HA and the particles being reported in literature (e.g. Fein et al., 1999; Tombácz et al., 2004; Feng et al., 2005; Wang and Xing, 2005; Janot et al., 2012). Shown by the SEM image in Fig. 5, there was no obvious HA attachment observed on the surface of Al₂O₃ particles. However, HA could produce thick coronas as adsorbed layers, which were covered with a large number of different polar groups, around Al₂O₃ particles in seawater (Ghosh et al., 2010). PGEs in solutions might react firstly with the polar groups on the thick corona around the surface of Al₂O₃ particle. Therefore, the inhibitory effect may be related to the positive or negative electrical characteristics of HA corona. The zeta potential of the solutions with Al₂O₃ particles was negative, and it changed from −4.64 to −12.3 mV when HA was added (Fig. 6), indicating the increase of repellant between the particle surface and PGEs. For kaolinite, the inhibition effect of HA on the adsorption of PGEs could be deduced. The interaction between the HA modified kaolinite particle surface and PGEs ions was speculated to be more like a chelating ion-exchange rather than a simple inorganic ion exchange, which was similar to that reported in Jülide and Apak (2006). Therefore, the adsorption of PGEs onto kaolinite particles decreased in the presence of HA, may be due to its broad combination with the −COO or −C=O groups in humic acid (Wood, 1996). However, the interactions among PGEs, tested particles and HA involve complicated specific molecular mechanisms which are beyond the scope of this research, and further research is needed for the understanding of the processes.

4.2.2. Bovine serum albumin

Like HA, BSA also changed the adsorption characteristics of Pt, Pd and Rh onto different particles (Figs. 3 and 4). For some particles, their morphology could be significantly altered by BSA adhering onto their surface (Liang and Wang, 2010; Rezwan et al., 2004; Song et al., 2012). These BSA-coated particles weakened the direct interaction between PGEs and the particles in solutions (Winsiewska et al., 2015; Givens et al., 2017). In this research, BSA showed an inhibition effect on the adsorption of Rh and Pt onto mineral particle Fe₂O₃ and MnO₂, which was similar to that of HA (Fig. 3 and Table S1). The SEM images show a similar adsorption feature of HA and BSA interacting with Fe₂O₃ particles (Fig. 5), and the log Kd and zeta potential values of the Fe₂O₃-BSA system were comparable to those of the Fe₂O₃-HA system (Figs. 3 and 6), which provided a clue that the adsorption behavior of Rh and Pt onto Fe₂O₃ particles in the presence of HA and BSA may be similar.

For the tested biogenic and lithogenic particles, in the presence of BSA, the log Kd values of PGEs onto all of them were approximately around 4 except for Pd onto CaCO₃ and kaolinite (Fig. 3). Shown by the SEM images, the adsorption feature of BSA onto CaCO₃ and Al₂O₃ particles was similar (Fig. 5). Moreover, compared to their corresponding values of the single kind of particle systems, the zeta potential of BSA-CaCO₃ and Al₂O₃ binary systems changed smaller than that of HA-CaCO₃ and Al₂O₃ binary systems (Fig. 6). It indicated that the adsorption of Rh and Pt onto these two kinds of particles may be dominated by the hydrophobicity/hydrophilicity and/or available surface area of the particles even in the presence of BSA. The adsorption behavior of Pd onto Fe/Mn oxides, CaCO₃ and kaolinite particles was different from Pt and Rh in the presence of BSA, which was possibly controlled by the distinction in their ability of interacting with BSA reflected by their partition and fractionation features (Figs. 3 and 4). In BSA-particle binary system, the inhibition effect of BSA on the adsorption of Pd onto the particles was smaller than that of HA, which was consistent with the results of the previous research that the interaction between BSA and Pd was weaker than that between HA and Pd (Liu et al., 2019).

4.2.3. Carrageenan

Like that of HA and BSA, carrageenan inhibited the adsorption of Rh and Pt onto mineral particles Fe₂O₃ and MnO₂ (Fig. 2 and Table S2). Compared to the corresponding system free of MOCs, the change of zeta potential of the Fe₂O₃ particle system in the presence of carrageenan was smaller than that of HA and BSA (Fig. 6). And carrageenan did not adhere to the Fe₂O₃ particles as significantly as that of HA and BSA, reflected by the SEM image (Fig. 5). These provided a clue that the direct interaction between
carrageenan micelles and PGEs would compete with and weaken the adsorption of PGEs onto Fe₂O₃.

The biogenic particles had a higher adsorption constant of PGEs than the lithogenic particles, and the log Kd values of PGEs in either of CaCO₃ and SiO₂ systems with and without carrageenan were close (Fig. 3), which indicated that carrageenan did not obviously affect the adsorption of PGEs onto biogenic particles. This may be the case because polysaccharides may not adsorb easily onto the particles with acidic surfaces such as SiO₂ (Liu et al., 2000; Laskowski et al., 2007). Besides, the SEM images show that only a very small amount of carrageenan attached to the surface of CaCO₃ and Al₂O₃. Therefore, for the biogenic and lithogenic particles used in the experiment, if there were no direct interactions between PGEs and carrageenan, the addition of carrageenan may have little influence on the adsorption of PGEs onto them, and the adsorption behavior of PGEs onto these particles with and without carrageenan should be similar. However, PGEs showed various adsorption behaviors onto biogenic and lithogenic particles according to the experimental results (Figs. 3 and 4). The degree of difference in log Kd values of Pt, Pd and Rh between the single kind of biogenic and lithogenic particle system and its corresponding carrageenan-particle binary system followed the order of Pt < Rh < Pd.

The results of zeta potential measurement showed that the values of the CaCO₃ and Al₂O₃ single kind of particle systems changed greatly from 10.8 to −9.01 mV and −4.64 to −30.1 mV respectively when carrageenan was added, indicating an increase in affinity to cations caused by carrageenan. A strong interaction between Pd and Rh ions and carrageenan was found in Liu et al. (2019). Therefore, the formation of PGEs-carrageenan complexes in carrageenan-biogenic and -lithogenic particle binary systems may influence the adsorption of PGEs onto particles greatly.

4.3. Mechanism of MOCs affecting PGEs adsorption onto inorganic particles in seawater

As discussed in Sections 4.1 and 4.2, MOCs significantly influenced the adsorption of Pt, Pd and Rh onto inorganic particles in seawater by the interactions among MOCs, particles and PGEs. As shown in Fig. 7, a possible mechanism of MOCs affecting the adsorption of PGEs onto inorganic particles in seawater could be obtained. In the following, three key processes of interactions which occurred in the particle-MOC binary systems were proposed based on the experimental results of this research and previous reports. First, PGEs interacted directly with particles like those of the single kind of particle system, in which the interaction was mainly determined by the physicochemical properties of PGEs and particles. Second, the interaction occurred between PGEs and MOCs in seawater (Wood, 1996; Sures and Zimmermann, 2007). PGEs could form stable complexes with MOCs and inorganic and small molecule organic ligands (Wood, 1996; Cobelo-García, 2013), which inhibited the adsorption of PGEs onto particles. Third, the competitive adsorption occurred between PGEs-MOCs and PGEs-particles. MOCs and the colloids they formed could adsorb PGEs in seawater (Zakharova, 1987; Liu et al., 2019), which led to a reduction in the distribution of PGEs onto the inorganic particles through the competitive adsorption. In addition, the presence of MOCs in the solution could change the surface properties of the particles (Fein et al., 1999; Tombacz et al., 2004; Feng et al., 2005; Janot et al., 2012; Sanderman et al., 2014). By adsorbing different kinds of MOCs, the surface potential and the area of the particles were changed (Figs. 5–7), which could directly affect the adsorption of PGEs. Besides, the attachment increased the macromolecular structure on the particle surface (Fig. 5), causing a reduction in macromolecules in the dissolved phase (Tombacz et al., 2004; Feng et al., 2005; Janot et al., 2012). On the whole, MOCs inhibited the adsorption of Pt, Pd and Rh onto inorganic particles in seawater in most instances and changed the fractionation of them.

5. Conclusions

In this research, controlled laboratory experiments were conducted to examine the adsorption characteristics of Pt, Pd and Rh ions onto inorganic particles. The results show that the chemical composition of particles played an important role in scavenging of Pt, Pd and Rh in seawater in a single kind of particle system. The representative mineral particles Fe₂O₃ and MnO₂ had the highest
degree of affinity with PGEs, followed by biogenic particles SiO2 and CaCO3 and lithogenic particle Al2O3 had the lowest adsorption capacity. The adsorption behavior of Pt, Pd and Rh in artificial seawater and UV irradiated natural seawater was different to an extent, which was possibly caused by the residual organic matter in the latter.

Macromolecular organic compounds (MOCs), including HA, BSA and carrageenan, changed the adsorption characteristics of Pt, Pd and Rh onto inorganic particles greatly. For Pt, MOCs inhibited its adsorption onto inorganic particles. For Pd, the effects of MOCs varied significantly with the particles, indicating complex interactions among Pd, MOCs and particles. For Rh, the effects of MOCs were similar to that of Pt in the presence of MOCs, except for Fe2O3 and MnO2 which indicated a more important role in scavenging of Rh. The presence of MOCs did not obviously change the adsorption feature of Pt onto biogenic particles; however, MOCs inhibited its adsorption onto mineral and lithogenic particles and the influence varied significantly. In summary, the results of this research suggest that the presence and type of macromolecular organic compounds are crucial in regulating the interactions between Pt, Pd and Rh ions and inorganic microparticles.

Conflicts of interest

We declare that we have no conflicts of interest to this work.

Acknowledgement

We would like to thank Yuxi Lu and Tianci Gao for their field and laboratory assistance, helpful advice and in-depth discussion. This study was funded by the National Natural Science Foundation of China (41370066).

Appendix A. Supplementary data

Supplementary data to this article can be found online at https://doi.org/10.1016/j.envpol.2019.113192.

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