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Minggang Cai
Wenlu Zhao
Mohammed Khairy
Mian Chen

See next page for additional authors

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Net volatilization of PAHs from the North Pacific to the Arctic Ocean observed
by Passive sampling

Haowen Zheng a, b; Minggang Cai a, b; Wenlu Zhao; Mohammed Khairy d, e; Mian Chen a, b; Hengxiang Deng a, b; Rainer Lohmann d, *

a State Key Laboratory of Marine Environmental Science, Xiamen University, Xiamen 361102, China
b College of Ocean and Earth Science, Xiamen University, Xiamen 361102, China
c School of Environmental Science and Engineering, Zhejiang Gongshang University, Hangzhou 310018, China
d Graduate School of Oceanography, University of Rhode Island, Narragansett, Rhode Island 02882-1197, United States
e Department of Environmental Sciences, Faculty of Science, Alexandria University, 21511 Moharam Bek, Alexandria, Egypt

*Corresponding author: Rainer Lohmann
E-mail: rloehmann@uri.edu
The North Pacific-Arctic Oceans are important compartments for semi-volatile organic compounds' (SVOCs) global marine inventory, but whether they act as a "source-sink" remains controversial. To study the air-sea exchange and fate of SVOCs during their poleward long-range transport, low-altitude atmosphere and surface seawater were measured for polycyclic aromatic hydrocarbons (PAHs) by passive sampling from July to September in 2014. Gaseous PAH concentrations (0.67-13 ng m$^{-3}$) were dominated by phenanthrene (Phe) and fluorene (Flu), which displayed an inverse correlation with latitude, as well as a significant linear relationship with partial pressure and inverse temperature. Concentrations of PAHs in seawater (1.8-16 ng L$^{-1}$) showed regional characteristics, with higher levels near the East Asia and lower values in the Bering Strait. The potential impact from the East Asian monsoon was suggested for gaseous PAHs, which – similar to PAHs in surface seawater - were derived from combustion sources. In addition, the data implied net volatilization of PAHs from seawater into the air along the entire cruise; fluxes displayed a similar pattern to regional and monthly distribution of PAHs in seawater. Our results further emphasized that air-sea exchange is an important process for PAHs in the open marine environments.

Keywords

Polycyclic aromatic hydrocarbons; Low-density polyethylene; Air-sea exchange; Long-range transport; High-latitude marine environment.
1. Introduction

Polycyclic aromatic hydrocarbons (PAHs) are ubiquitous in the environments worldwide, and some of them are toxic, carcinogenic and mutagenic (Bozlaker et al., 2008; Okona-Mensah et al., 2005; Perera et al., 2005). Besides their petrogenic origin, PAHs are also generated during the incomplete combustion of organic materials, like fossil fuels and biomass (Yunker et al., 2002). As a class of semi-volatile organic compounds (SVOCs), they could arrive in the polar area from temperate regions with subsequent deposition and re-volatilization, representing an emerging concern in the Arctic like other conventional pollutants (Cai et al., 2016; Friedman and Selin, 2012; Laender et al., 2011).

Atmospheric transport has been considered the primary pathway for PAHs transported from the lower latitudes to the Arctic environments (Mulder et al., 2015; Dotel et al., 2020). Previous studies showed that PAHs found in the Arctic seawater and sediments mainly originated from natural underwater hydrocarbon seeps, while those in the air were from atmospherically derived sources (Harvey et al., 2014; Yunker et al., 2011; Foster et al., 2015). With the concern of climate change, the “polar sinks” for many conventional pollutants may become secondary sources via air-water exchange, that is, the declining sea ice coverage and rising temperature could lead to an accelerating release of PAHs from sea ice and seawater to the atmosphere (Hung et al., 2010; Ma et al., 2011, 2013; Galban-Malagon et al., 2013). Thus, studying PAHs in the atmosphere and related interfaces is helpful to better know the current state of the PAH emission and their fates in the Arctic (Friedman et al., 2014).

Passive sampling is an effective monitoring technique for SVOCs, which is easy to operate, cost-effective and with high enrichment of the target compounds, and has been applied in various global monitoring projects (Lohmann et al., 2001; Harner et al., 2003; Meijer et al., 2003; Jaward et al., 2004; Khairy and Lohmann, 2014; Mcdonough et al., 2014; Zhao et al., 2018). This method integrates contaminant concentrations over time, representing time-weighted averages (Stuer-Lauridsen, 2005; Shaw and Mueller, 2009; Wania and Shunthirasingham, 2020). For PAHs, the freely-dissolved pollutants can be sampled with low-density polyethylenes (LDPEs) (Khairy and Lohmann, 2014). Concentrations of target compounds that do not reach equilibrium during the exposure period can be estimated relying
on the diffusive loss of performance reference compounds (PRCs) (Booij et al., 2002; Mayer et al., 2003; Khairy and Lohmann, 2012).

Although previous studies have reported the distribution and direction of PAHs air-water exchange in the northern Pacific and the Arctic, as well as fugacity model simulations, the seasonal and regional trends of PAHs remain uncertain (Ke et al., 2017; Ma et al., 2013). Hence the air-sea exchange process of PAHs from the North Pacific to the Arctic needs further research.

In this study, we performed spatially resolved air and water measurements during the sampling cruise, and the objectives were to (1) obtain the spatial and temporal distribution characteristics of atmospheric and dissolved PAHs in seawater from the North Pacific Ocean to the western Arctic Ocean; (2) assess the source of gaseous and freely dissolved PAHs at most sites; (3) derive the direction and magnitude of the air-sea flux exchange process of PAHs.

2. Materials and methods

2.1 Area description

During the Chinese sixth Arctic scientific expedition cruise from July to September in 2014, we collected 32 atmosphere and 16 surface seawater samples in the Japan Sea, Bering Sea and Chukchi Sea onboard the R/V Xuelong (Snow Dragon). The Bering Strait, connecting the Bering Sea with the Chukchi Sea, and linking the Asian and American continents, is the boundary of the Arctic and Pacific Oceans. The Bering Sea is a semi-enclosed, high-latitude sea that is almost divided equally between a deep basin (maximum depth 3500 m) and the continental shelves (<200 m). The eastern broad (>500 km) shelf of the Bering Sea contrasts with the narrow (<100 km) western shelf (Stabeno and Van Meurs, 1999). Water in the vast (~500 km wide from east to west and ~800 km long from north to south) and shallow (~50 m) Chukchi Sea (Arctic Ocean) is strongly forced by Pacific Ocean water entering through the Bering Strait (Woodgate et al., 2005), which delivers heat, freshwater, nutrients, and carbon to the Chukchi shelf and the Arctic Ocean.

In this study, there were 16 sampling stations (Table S1 in Supporting Information, SI), of which PS-01 to PS-07 were located in the Sea of Japan and northwest Pacific Ocean, while PS-08 to PS-10 in the Bering Sea and the Bering Straits, and PS-11 to PS-16 in the Arctic Ocean (mainly in Chukchi Sea).
2.2 Materials Preparation and Field sampling

2.2.1 Preparation for LDPEs

Both atmospheric and seawater samplers were made of LDPEs with PRCs, which is similar to the material used in semipermeable membrane devices. The inclusion of PRCs provided a means to identify the absorption of hydrophobic PAHs by estimating dissipation rates of PRCs (Booij et al., 2002; Lei et al., 2020). LDPEs (10 cm × 40 cm size, 50 μm thickness) were cleaned in dichloromethane and n-hexane for 24 h respectively. Deuterated PAHs (pyrene-d₁₀ and benzo[a]pyrene-d₁₂, namely Pyr-d₁₀ and BaP-d₁₂) were used as PRCs to infer the equilibrium concentration of compounds in the passive samplers as in previous work (Booij et al., 2002; McDonough et al., 2014). LDPEs were soaked in PRCs and continuously shaken for one month to achieve homogenous uptake prior to deployments.

2.2.2 Simultaneous sampling in atmosphere and surface seawater

Atmospheric and surface-seawater LDPEs were deployed and collected simultaneously on the underway R/V Xuelong in the Japan, Bering and Chukchi Seas (Fig. S1), and LDPEs were changed every three days. The sampling site coordinates were calculated as the middle position of the start and end of sampling positions.

Surface-seawater LDPEs were put into a stainless-steel pipe, and continuously exposed to fresh marine surface seawater from the onboard seawater supply. We controlled the flow rate of the seawater such that the LDPEs were submerged constantly. The atmospheric sampling device was installed on the top deck, which was about 26 m above sea level. The device consisted of two stainless-steel bowls, connected in the middle by a stainless-steel center shaft. The LDPEs, thread on a metal wire, were fastened on the center shaft for sampling. All the LDPE samples were sealed with an aluminum foil bag and stored at -20°C.

2.2.3 Other auxiliary parameters

Auxiliary parameters such as temperature and salinity were provided by the Xuelong ship real-time monitoring system, and other parameters such as wind speed, wind direction and air temperature were provided by the automatic weather station on board Xuelong ship.

2.3 Pretreatment and analysis

LDPE samples were thawed out at room temperature, then cleaned with Milli-Q water and
any excess water or biofouling was removed with KimWipes. 200 mL of n-hexane was added to completely cover LDPEs after addition of 50 μL 100 ppb PAHs recovery indicator surrogates (acenaphthene-d_{10}, phenanthrene-d_{10}, chrysene-d_{12} and perylene-d_{12}, namely Acp-d_{10}, Phe-d_{10}, Chry-d_{12} and Pery-d_{12} respectively). After extraction for 24 h, the n-hexane was decanted and the extract was kept in a clean glass bottle. After a solvent exchange to dichloromethane, and repeating the above steps, extracts were combined. The extract was concentrated to 1mL in a 30°C water bath by a fully automatic sample concentrator, transferred to a volumetric tube and slowly purged to 100 μL with high-purity nitrogen. Then 50 ng of deuterated terphenyl and 35 ng of 2, 4, 6-Tribromobiphenyl were added as internal standards. The extract was sealed and frozen for storage before analysis. The weight of the extracted LDPE samples was recorded before and after treatment.

An Agilent 6890-5973 GC-MS equipped with a DB5-MS quartz capillary column (30 m × 0.25 mm i.d. × 0.25 μm, J&W Scientific Inc., Folsom, U.S.A.) was used to detect PAHs. The high purity helium was used as carrier gas with the flow rate of 1mL min^{-1}. The temperature programming of chromatography column started at 90 °C with three 3 minutes hold, then reached 110 °C at 5 °C min^{-1}(holding 2 minutes), increased by 8 °C min^{-1} until 200 °C (holding 3 minutes), and finally reached 315 °C at 5 °C min^{-1}fiv, keeping the final temperature for 5 minutes.

2.4 Quality assurance and quality control

Avoiding contamination was an important consideration in all steps associated with the extraction and analysis of LDPEs. All glassware used during pretreatment were cleaned, baked at 450°C for at least 4 hours, and thoroughly solvent-rinsed before use. The field blanks were regularly included in the sampling protocol, and included when samplers were changed during deployment and retrieval. Method detection limits (MDLs) were calculated as three times the standard deviation of the average field blank concentration. The MDL of the target compounds are listed in SI as well as instrument detection limits. All of the data were blank-corrected. When the concentration of the target compounds was less than the detection of limits, half of MDL was taken as its concentration value (Antweiler and Taylor, 2008).

22 PAHs were detected in this study. In addition to the 16 priority PAHs listed by U.S.
EPA, the other six compounds were Biphenyl (Biph), 1-Methylphenanthrene (1-MePhe), Retene (Ret), Benzo[e]pyrene (BeP), Perylene (Pery) and Benzo[j]fluoranthene (BjF) (Table S2 in SI). Due to the high environmental background concentration of naphthalene, it will not be discussed in this study. In order to improve the quality and credibility of the data, if the concentration of a certain target compound was greater than MDL but detected at less than 20% of the total sites, it was omitted from the discussion. A total of nine PAHs in atmospheric samples were regularly detected, which included phenanthrene (Phe), fluorene (Flu), Phe, anthracene (Ant), 1-MePhe, fluoranthene (FluA), pyrene (Pyr), chrysene (Chry), benzo[b]fluoranthene (BbF)/ benzo[k]fluoranthene (BkF) (because the peaks of BbF and BkF were hard to chromatographically separate, the total concentration of BbF/BkF are reported). While 11 PAHs in surface seawater samples were effectively detected, which included Flu, Phe, Ant, 1-MePhe, FluA, Pyr, Chry, Biph, benz[a]anthracene (BaA), BbF/BkF. Concentrations of PAHs in the samples were blank-corrected for the amounts detected in the field blanks, but not recovery corrected. The average recovery rates of the four surrogates that Acp-d_{10}, Phe-d_{10}, Chry-d_{12}, and Pery-d_{12} were 70 ± 12, 75 ± 15, 85 ± 13, and 74 ± 18% respectively.

For the uncertainty analysis, it is difficult to estimate the overall uncertainty of air-water exchange by passive sampling, because it involves multiple mathematical functions and values that have both normal and lognormal errors associated with them. Additionally, the uncertainties in $H^0$ are not well characterized, causing the uncertainty associated with the temperature- and salinity corrections were not included.

2.5 Calculations and data analysis

2.5.1 Calculations of PAH concentrations

For each compound, the fraction of equilibrium (f) achieved for each compound was determined by fitting the equilibrium of the PRCs using temperature-corrected air-water partitioning coefficient in LDPEs ($K_{PE}$) values to a model curve derived from equation (1) (Liu et al., 2016),

$$f = 1 - e^{-\frac{R_s t}{K_{PE} M_{PE}}}$$

Where $R_s$ (L day$^{-1}$) is the sampling rate defined as the volume of water or air that comes into contact with the sampler per day; $t$ is deployment time (days); $M_{PE}$ is the PE weight (kg);
the values of $R_s$ can be estimated using nonlinear least squares methods, by considering $f$ as a continuous function of $K_{PE}$, with $R_s$ as an adjustable parameter using Excel Solver to obtain the best fit (Booij and Smedes, 2010). The average gaseous sampling rate of PAHs was $1700 \pm 150$ m$^3$ day$^{-1}$, among which the Pacific Ocean and the Bering Sea was $620 \pm 410$ m$^3$ day$^{-1}$, however the Arctic Ocean was $3100 \pm 1300$ m$^3$ day$^{-1}$, resulted from different wind speed and directions described below. The aqueous sampling rate of PAHs was $140 \pm 54$ L day$^{-1}$.

Gaseous and freely dissolved concentrations, $C_{A/W}$ of compounds were calculated from the equation (2),

$$C_{A/W} = \frac{c_{PE} - c_{Blank}}{K_{PE} (1 - e^{-R_s \cdot K_{PE} M_{PE}})} \tag{2}$$

Where $c_{PE}$ is the PE-normalized concentrations. For more details, see the Supporting Information.

2.5.2 Brief description of calculation and two-film model

The Whitman two-film resistance model was used for the air-sea exchange flux ($F_{aw}$) calculation in the modified version (Liss and Slater, 1974).

$$F_{aw} = v_{a/w} (C_w - \frac{C_a}{K_{aw}}) \tag{3}$$

where $F_{aw}$ is calculated with the mass transfer velocity ($v_{a/w}$), the concentrations in seawater and atmosphere, $C_w$ and $C_a$, and air-water partitioning coefficient corrected by ambient temperature, $K_{aw}$ (Liu et al., 2016). Calculation details of PAHs air-sea exchange fluxes at different sites are listed in SI from Table S6 to S12.

2.6 Air mass back trajectories

The NOAA’s HYSPLIT model was used to track the origins of air masses (Draxler and Hess, 1998). Air mass back trajectories were set as 12 h steps that traced back the air masses for 5 days, using the sampling height as arrival height.
3. Results and discussion

3.1 PAHs in the atmosphere

3.1.1 Concentrations and distributions

Total gaseous concentrations of $\sum_9$PAH ranged from 0.67 to 13 ng m$^{-3}$ with the mean value of 3.7 ± 3.0 ng m$^{-3}$, close to previous data of 0.93-93 ng m$^{-3}$ for $\Sigma_{15}$PAH reported over the North Pacific and Arctic Ocean in 2003, as well as 0.91-7.4 ng m$^{-3}$ with a mean of 3.3 ± 1.7 ng m$^{-3}$ for $\Sigma_{18}$PAH reported in 2010 (Ding et al., 2007, Ma et al., 2013). Observation showed the maximum concentrations occurred at PS-01 located in the Bohai Sea, and lowest at PS-16 in the Arctic Ocean (Fig. 1, Table S3).

In this study, nine different PAHs were regularly detected in most gaseous samples, including three-ring PAHs like Flu, Phe, 1-MePhe, Ant, and four-ring PAHs like FluA, Pyr, and Chry. Higher molecular weight PAH concentrations in the atmosphere were typically below their limits of detection. Across the entire sampling cruise, Phe displayed the highest average contribution of 52% to $\sum_9$PAH, followed by Flu, which contributed another 44%. Hence, the cumulative contributions of other congeners were less than 5%. Our results mirror previous results, in that Phe was also the dominant compound among the gas phase PAHs, contributing 50% to $\Sigma_{15}$PAH over the North Pacific and Arctic Ocean (Ding et al., 2007).

3.1.2 Decreasing distribution with latitude

Overall, a significant negative correlation was observed between gaseous $\sum_9$PAH and latitude (the coefficients of determination, $R^2$ was 0.61, the P-value < 0.0010). The partial pressure of seven PAH congeners were calculated by a modified Clausius-Clapeyron equation (Venier et al., 2012), these results were obviously consistent with the trends in concentration with temperature. The mobility of PAHs was usually reduced by decreasing temperature, which contributed to decreasing volatility of PAHs, especially for higher molecular weight PAHs. As shown in Fig. 2, the partial pressure of FluA, Pyr and Chry decreased faster than for the low molecular weight PAHs such as Flu and Phe (All the $R^2$ and P-values are given in SI). Higher molecular weight PAH congeners were more sensitive to temperature change, in line with expectations for gaseous PAH concentrations with latitude (Fig. 2).

On the other hand, PAHs are more susceptible to photochemical-degradation in the
atmosphere. The concentration ratios of Flu to Phe were positively related to latitude ($R^2=0.58$, $P<0.0010$) from the North Pacific to the Arctic (Figure 1); and the ratio FluA/(FluA+Pyr) was greater than 0.5 at all stations (Fig. 4). The total photochemical residence time ($\tau_{\text{total}}$) of Flu and FluA were derived as 21 and 20 hours, and 9 and 5 hours for Phe and Pyr, respectively (Keyte et al., 2014); the observed low concentrations of Ant in the atmosphere were likely caused by its 2 hours of $\tau_{\text{total}}$. Furthermore, the atmospheric half-life of Flu is higher than Phe, which means higher removal rates occurred in Phe, and Flu presents a greater transmission potential than Phe for long-range atmospheric transport (LRAT) (Halsall et al., 2001).

### 3.1.3 Influence of air mass

Diagnostic ratios of gaseous PAHs were calculated, but need to be interpreted with caution, because the ratios might be affected by potential post emission processes. 1-MePhe/Phe in the atmosphere were less than 0.5, implying that the source of the PAHs could be attributed to combustion processes (Yunker et al., 2002; Deka et al, 2016; Wu et al., 2019). This was further supported by the isomeric ratios of FluA/(FluA+Pyr) that ranged from 0.42-0.82; a ratio greater than 0.5 indicates the main sources of grass, wood or coal combustions, and between 0.4 to 0.5 usually suggests petrogenic combustion (Qu et al., 2019; Qi et al., 2020).

At the mid-latitude sites, the air mass mainly originated from the coast of China, Korea, Japan and Russia, and close-by terrestrial regions as indicated by the air mass back trajectories (Fig. S7 and S8 in SI). With the trajectories of air mass that was used to study the origin of atmospheric PAHs in Japan and other neighboring areas, high concentrations of PAHs and emissions have been already observed in East Asian areas (Ohura et al., 2004; Primbs et al., 2007). Continuously decreasing concentrations of PAHs occurred from the Bering Sea to the Arctic, but increased again between PS-09 and PS-11, where back trajectories indicated the air had passed through Alaska and Siberia, especially on Kamchatka peninsula, consistent with the impact of terrestrial source (Fig. S8 in SI). According to the air mass back trajectories, the source of the atmosphere PAHs in July was significantly affected by the southwest monsoon, which is also an important pathway for long-range transport.
3.2 PAHs in the seawater

3.2.1 PAH profiles in seawater

The concentrations of total freely dissolved $\sum_{11}$PAH in the surface seawater ranged from 1.8 to 16 ng L$^{-1}$ with the mean value of 7.7 $\pm$ 4.6 ng L$^{-1}$. The maximum concentrations occurred at PS-03 on the northeast coast of Japan, followed by station PS-02 and PS-04, and the minimum was observed at PS-10 in the Bering Strait (Fig. 3, Table S4). A total of 11 different PAHs were regularly detected in the freely-dissolved phase in the seawater samples, including Biph, Flu, Phe, 1-MePhe, Ant, FluA, Pyr, and Chry, BaA, BbF/BkF. The concentrations of higher molecular weight PAHs in seawater were typically below the limits of detection, similar to the atmosphere.

The ratios of 1-MePhe/Phe and FluA/(FluA+Pyr) indicated the main sources of combustions included many types (Yunker et al., 2002; Qu et al., 2019; Qi et al., 2020). The results in seawater were consistent to previous studies in the North Pacific and Arctic Ocean (Ding et al., 2007; Ma et al., 2013).

While Phe also dominated the dissolved $\sum_{11}$PAH, its average contribution was only 36%. Hence more PAHs were regularly present beyond Phe, including Flu, which contributed 25%, FluA (11%), Pyr (10%), in addition 1-MePhe and Biph were 8% and 5%, respectively. Other congeners like Ant, BaA, BbF/BkF were only detected in a few of the samples beyond the Bering Sea (for details see SI), whose contributions were less than 4%. The combined influences of sources like rivers, runoff, currents as well as biogeochemical process cause more higher molecular weight PAHs to be present in the surface seawater (Sambrotto, 1984; McDonough et al., 2014). We speculate that differences in biodegradation and photochemical-degradation of PAHs increased the proportion of higher molecular weight PAHs in seawater (González-Gaya et al., 2019). Bacterial degradation rate constants ($K_D$) showed a significant inverse correlation with $K_{ow}$, which means the capacity to biodegrade higher molecular weight and hydrophobic PAHs (i.e., FluA, Pyr, Chry, BbF/BkF) were slower than for lighter weight PAHs (Table S5) (Tucca et al., 2020). Besides, higher molecular weight PAHs in the air sorb to aerosol according their higher octanol-air ratios, contributing to wet and dry deposition entering seawater.
3.2.2 Changes of freely-dissolved PAH distribution with region

Freely dissolved $\Sigma_{11}$PAH concentrations varied with sampling locations as well, but displayed obvious regional trends instead of correlation with latitude. They ranged from 12 to 16 ng L$^{-1}$ in the Japan Sea to the North Pacific Ocean, located in a mid-latitude temperate zone. Concentration were lower in the nearby Bering Strait ranging only from 1.8 to 4.8 ng L$^{-1}$, as well as sites in the Arctic Ocean (2.7-6.5 ng L$^{-1}$). In general, dissolved $\Sigma_{11}$PAH concentrations closer to shore were relatively higher than those in the open ocean (Fig. 3).

Lower concentrations of seawater $\Sigma_{11}$PAH were present at station PS-08, PS-09 and PS-10 in the Bering Strait, which was known as one of the most productive waters in the Arctic Ocean. Likely removal of PAHs by high amounts of biogenic particles significantly decreased the concentration of PAHs in the ocean, where the plankton biomass was higher, consistent with the relevance of the biological pump (González-Gaya et al., 2019). Another reason for low concentrations of $\Sigma_{11}$PAH in the Arctic Ocean are the oceans’ long residence time, as well as the weak sea ice melting in summer. The ocean current exchange is relatively slow, causing the residence time of surface water to be about 10 years. Melting sea ice has been recognized as an important factor controlling the distribution of PAHs in the Arctic, where only part of the sea ice melts even in summer (MacDonald et al., 2000). The lower salinity supported that continental runoff or melting ice rather than ocean currents were the pathways of PAHs into the surface water, and the atmospheric deposition was also a key source (see below).

3.3 Air-sea exchange of PAHs with two-film model

3.3.1 General net volatilization

The air-sea exchange fluxes of the seven PAHs was generally dominated by net volatilization especially from the North Pacific to the Arctic (Fig. 5, Table S12). Generally, fluxes of the more volatile PAHs especially for Flu and Phe were greater and nearly accounted for >70% of all (Table S12 in SI), based on their dominant contribution to gaseous and free-dissolved PAHs. The maximum values of volatilization and deposition fluxes of $\Sigma_{11}$PAHs were observed at site PS-04 (11 μg m$^{-2}$ day$^{-1}$) and PS-01 (-2.5 μg m$^{-2}$ day$^{-1}$) respectively. Flux distributions indicated that the deposition was only observed near land from Shanghai to the Bohai Sea, while net volatilizations were more widespread. For instance, Flu dominated the
flux at site PS-04 with the value of 4.2 μg m$^{-2}$ day$^{-1}$, and Phe at site PS-01 was -5.2 μg m$^{-2}$ day$^{-1}$, due to the greatest concentrations of dissolved Flu at PS-04 and gaseous Phe at PS-01 respectively. The effect of volatilization from coastal waters with high PAH levels has reported in other PAH-impacted regions such as the Atlantic, Narragansett Bay (USA) and the southeast Mediterranean (Nizzetto et al., 2008; Castro-Jiménez et al., 2012; Lohmann et al., 2011). The fluxes were mainly contributed by three-ring PAHs, while the fluxes of four-ring PAHs showed less variation in the North Pacific and the Arctic.

3.3.2 Flux trend closely coupling with dissolved phase

The derived air-water exchange fluxes correlated significantly to regional and monthly changes, similar to the freely-dissolved PAH distributions, which might imply the one of driver by concentration gradients between seawater and air (Fig. 5). Volatilizations dominated in the air-sea exchange because the seawater concentrations corrected by $K_{aw}$ were one to three orders of magnitudes higher than equivalent concentrations in the atmosphere.

As for individual congeners, there were no clear trends for three-ring PAHs, but diverse ones for higher-ring PAHs, especially FluA and Pyr declined with increasing latitude (Fig. S5 in SI). These results verified that higher molecular weight PAHs were more sensitive to changes with temperature. PAHs are moderately volatile and hydrophobic, which facilitates their partitioning from air and water into organic phases (Nizzetto et al., 2010). Obvious outputs of biogenic particles occurred in the Bering Strait and caused a significant decrease in seawater concentration as well as fluxes. In the Arctic, low PAHs fluxes were driven basically by low atmospheric and seawater concentrations. However, the net volatilizations only had a minor impact on atmospheric concentration. According to the long-range transport and diffusion advection of PAHs, the atmospheric concentrations tend to decrease as the latitude increases, so the transports depend largely on temperature changes. In our study, strong correlations were observed for air-sea exchange fluxes with concentrations rather than temperature.

3.4 Implication for the source-sink pattern of PAHs

3.4.1 Source pattern of PAHs from the Pacific to the Arctic

For further analysis, principal component analysis (PCA) was used to elucidate linear combinations of PAHs, to distinguish between the samples to assess different sources (Nemr et
Two compounds were obtained from PCA, namely the first principal component (PC1) and second principal component (PC2), contributed 74% to the total variance (Fig. 4). Correlation analysis showed high relevance of PC1 for four-ring PAHs, such as FluA and Pyr (r=0.98), FluA and Chry (r=0.90), Pyr and Chry (r=0.94); and PC2 represented mainly three-ring PAHs such as Flu and Phe (r=0.82), Flu and Ant (r=0.56), Phe and 1-MPhe (r=0.67). The scores on PCA showed that all the sites in the Bering Sea and Arctic were concentrated together in the negative axis of PC1 and PC2, while sites in the North Pacific were more dispersed.

The diagnostic ratios and PCA analysis provided qualitative information about the spatial patterns of PAH sources. Based on the analysis mentioned earlier, we inferred that sources of seawater and atmospheric PAHs in the Bering Sea and Arctic were mainly from combustion processes. In the Arctic, the diagnostic ratios of PAH concentrations in seawater were similar to that in atmosphere, which might provide the evidence that PAHs reached high latitudes by long-range atmosphere transport (LRAT). Hence the atmospheric concentrations contributed only low levels in the high latitudes, and combined with sea ice melting, runoff and other biogeochemical process to affect the ratio characteristics. On the other hand, both of the seawater and atmospheric PAHs in the Pacific Ocean were also mainly derived from combustion processes. The difference is that the North Pacific is known as an important cruise routes area, which potentially contributes additional sources factors, complicating the seawater PAHs. Therefore, petroleum combustion emitted by ships is an inevitable source for PAHs in addition to runoff, ocean currents and atmospheric deposition.

3.4.2 Net volatilization prospect under climate changing

Our results illustrated that the majority of PAHs displayed a net volatilization trend from East Asia to the Arctic, which was on the contrary of the previous studies (Zhong et al., 2012; Ma et al., 2013). The significant differences in the fluxes were partly attributed to the different concentrations of gaseous and dissolved PAHs between these two studies whose samples were from different years. The time lag to previous studies might have been sufficient to cause an increase in PAH concentration in surface seawater for the ongoing-emitted PAHs year by year. In addition, East Asia is a possible continental source area and has been estimated to contribute greatly to the global emission inventory of PAHs (Shen et al., 2013). Furthermore, sea surface temperatures (SSTs) in August 2014 were as much as 4 °C warmer than the 1982-2010 August
mean in the Bering Strait and the sixth smallest Arctic sea ice extent by the satellite record (1979-2014) occurred in the summer of 2014 (Chen et al., 2016), which might partially explain the different results reported here. Lastly, sampling of PAHs by passive sampling only captures freely dissolved or gas-phase compounds, which should lead to air-water exchange fluxes not impacted by inadvertent capture of PAH bound to colloids or small particles.

Some persistent organic pollutants, whose primary emissions have been reduced, are suggested having iterative processes (including deposition, volatilization and re-deposition) in the changing Arctic (Cai et al., 2012). Since the ice will prevent the escape of PAHs in winter, we can assume that atmospheric PAHs will deposit in snow and ice by dry/wet deposition processes and the Arctic turned into a sink again. In other words, as a result of global warming with sea ice retreat, whether the polar region is a sink or secondary source depends on the seasonal variety. The air-sea exchange fluxes in our study displayed significantly lower in September than in July (Fig. 5), which might be driven by decreasing temperature and weakened monsoon. PAH concentrations and fluxes are thus likely affected by temporal changes, but more evidence is needed (Fig. S6 in SI). Combining this with our result that fluxes were consistent with surface seawater PAHs concentrations, then seawater might dominate the air-sea exchange when not covered by sea ice. But whether the hypothesis is reasonable or not need to be further studied, and which might determine the role of the Arctic as “sink” or “source” of PAHs.

4 Conclusions

This study focused on the air-sea exchange process of PAHs, relying on LDPEs passive sampling from the North Pacific to Arctic. The gaseous concentrations of $\sum_9$PAH ranged from 0.67 to 13 ng m$^{-3}$ and displayed a significant decreasing trend with latitude, which was dominated by long-range transport and photochemical-degradation of PAHs. Different distributions were explained by the air mass back trajectories, coupled with the influence of air-sea exchange and dry/wet deposition processes. The concentrations of total freely dissolved $\sum_{11}$PAH ranged from 1.8 to 16 ng L$^{-1}$ whereas higher molecular weight PAHs showed more presence. The changes in PAH profiles can likely be attributed to their relative capacity to undergo biodegradation and dry/wet deposition. In addition, regional distributions were shown
in freely dissolved PAHs, showing high levels in the North Pacific. The lower levels were impacted by biogenic particle removals in the Bering Sea, and were attributed to slow renewal of seawater and melting ice in the Arctic. Molecular ratios of the PAHs in atmosphere and surface seawater indicated combustion sources, and the source indication might provide some evidence for LRAT of high-latitude marine environmental PAHs. Overall, the air-sea fluxes of PAHs presented mostly net volatilizations with special regional and monthly changes, controlled by seawater concentrations. The air-sea exchange process showed only a relatively minor importance of air concentrations, instead the fluxes and dry/wet depositions from atmosphere to seawater might were more important especially in the Arctic.

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Fig. 1. The gaseous concentration distribution of Flu and Phe along the cruise transect (the figure above), and the ratios of Flu to Phe as a function to latitude (the figure below). (The domination of Flu and Phe presented in Fig. S3)
Fig. 2. The partial pressure of PAH congeners as a function of water temperature (left), and the ratios of the sum of three-ring PAHs to four-ring PAHs as a function of air temperature (right).

Fig. 3. The PAH concentrations in surface seawater along the cruise transect.
Fig. 4. Results of PAH diagnostic ratios of MePhe/Phe and FluA/(FluA+Pyr) (left), and PCA (right) for samples in seawater (dots in blue) and atmosphere (dots in red).
Fig. 5. The distribution of PAH air-sea exchange fluxes, and their monthly changes.