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Overall comparison and source identification of PAHs in the sediments of European Baltic and North Seas, Chinese Bohai and Yellow Seas

Pu Wang a,b,c, Wenyi Mi d, Zhiyong Xie c,* Jianhui Tang e, Christina Apel c, Hanna Joeriss c, Ralf Ebinghaus c, Qinghua Zhang b,a

a Hubei Key Laboratory of Environmental and Health Effects of Persistent Toxic Substances, Institute of Environment and Health, Jianghan University, Wuhan 430056, China
b State Key Laboratory of Environmental Chemistry and Ecotoxicology, Research Center for Eco-Environmental Sciences, Chinese Academy of Sciences, Beijing 100085, China
c Institute of Coastal Research, Helmholtz-Zentrum Geesthacht, Geesthacht 21502, Germany
d MINJIE Institute of Environmental Science and Health Research, Geesthacht 21502, Germany
e Yantai Institute of Coastal Zone Research, Chinese Academy of Sciences, Yantai 264003, China

Abstract

An international sampling campaign was carried out to comprehensively investigate the occurrence of polycyclic aromatic hydrocarbons (PAHs) in the marine sediments from the European Baltic and North Seas, Chinese Bohai and Yellow Seas. The concentrations of ∑18PAHs in the samples from these four seas were in the range of 0.91–5361 ng/g dry weight (dw), 0.46–227 ng/g dw, 25.0–308 ng/g dw and 4.3–659 ng/g dw, respectively. 4-rings PAHs, e.g., fluoranthene, pyrene and benzo(b)fluoranthene, were commonly the dominant compounds in all the samples. The PAH sources were identified via composition patterns, diagnostic ratios, principal component analysis (PCA) and positive matrix factorization (PMF). Coal combustion, vehicular emission, coke plant and petroleum residue were apportioned as the main sources in these marine sediments. However, through PMF modeling, different contributions of these sources were quantified to the deposited PAHs in the seas, suggesting distinct anthropogenic impacts on the adjacent marine system. It is noteworthy that biomass combustion may not be the main source of PAHs in the majority of sediments from these seas. This was evidenced by the ratios of naphthalene against its methylated derivatives (i.e. 1-, 2-methylnaphthalenes) other than the composition pattern in the samples, of which the approach is in prospect of developing in future studies.

GRAPHICAL ABSTRACT
1. Introduction

With the globally expanding economic and industrial developments in the 20th century, many developed and developing countries, such as the United Kingdom, USA, Japan and China, have experienced various environmental problems, including air, water, and soil pollution, even the food contamination, which raised great concerns about the environmental safety and local human health. Thereafter, energetic efforts have been made and significant improvements were achieved for the environment and human life in those developed countries. As two of the largest economy groups in the world, European Union (EU) and China have been experiencing different stages of the economic development in recent years. For example, Germany, the largest economy in EU, has established the mature industrial system with relatively less contamination, by contrast, many industries with severe pollution are still situated in China, which results in relatively higher emission of various pollutants in the environment including the surrounding marine system.

As typical bordering seas and semi-enclosed seas in the world (Fig. S1), the European Baltic and North Seas, and the Chinese Bohai and Yellow Seas have their societal significance with a broad range of important modes of uses, from oil and gas, nearshore aquaculture, offshore fishing, wind energy, shipping and other industrial activities to tourism and natural preservation. However, they also serve, or served, as dumping sites for onshore activities. A difference is the fact that the North European seas have many different national coasts while in case of the Chinese seas only a limited number of international partners share responsibility for the marginal seas. This possibly makes the pollution by anthropogenic activities of difference in these seas. Among the pollutants, persistent toxic substances (PTSs) received great attention due to their toxicity, persistence and bioaccumulation in the environment in the past several decades (Porta, 2015; Wang et al., 2017). Many PTSs are released from the terrestrial sources and may enter into the marine environment via long-range transport by water and/or air, leading to a potential exposure risk of the marine animals to various toxic chemicals (Wolska et al., 2012). Since most PTSs are hydrophobic and tend to bind strongly to particulate matter, which facilitates their deposition in water, sediment is therefore considered a primary reservoir of PTSs in aquatic systems (Maletić et al., 2019). Poly-cyclic aromatic hydrocarbons (PAHs) are such a kind of PTSs, which are known carcinogens and mutagens, and their sources, transport, and effects have been studied extensively (Kim et al., 2013). PAHs are natural components of coal and oil, and also formed during the combustion of fossil fuels and organic material, as well as biomass burning (Kim et al., 2013). They enter the marine environment through atmospheric deposition, road run-off, industrial discharges and oil spills (CEMP, 2009). PAHs sorbed on sediment are considered to be stabilized by physicochemical association with the sediment matrix, and undergo practically no further changes (Maletić et al., 2019). Therefore, its occurrence and distribution pattern in the marine sediment may provide important information for comparison of the impact from the anthropogenic activities on the adjacent marine system (Wolska et al., 2012; Maletić et al., 2019).

Even though the occurrence of PAHs in the marine environment was constantly studied in many coastal or oceanic areas (Maletić et al., 2019) including the European seas (CEMP, 2009) and Chinese seas (Zhao et al., 2020), less was done to comprehensively compare their distribution pattern and source composition in the marine sediment from different seas, which may limit the understanding of environmental fate of PAHs and anthropogenic effects on marine system on a global scale.

In the present study, an international sampling campaign was conducted on these four seas to comprehensively investigate occurrence of PAHs in the surface sediments. The potential sources were identified based on the composition patterns, diagnostic ratios, principal component analysis (PCA) and positive matrix factorization (PMF), and the quantitative comparison between the different areas may shed light on the fate of PAHs in the marine environment and anthropogenic impact from the surrounding countries.

2. Materials and methods

2.1. Sampling areas

The Baltic Sea is a relatively shallow inland sea in north-east Europe, bounded by the coastlines of Denmark, Estonia, Finland, Germany, Latvia, Lithuania, Poland, Russian Federation and Sweden (Fig. 1).
Fig. S1). Its average depth is 57 m. As a semi-enclosed sea, the Baltic Sea received much runoff from coastal and river areas with high concentrations of population and industry, and has been recognized as an ecologically vulnerable system (Johannesson and Andre, 2006). The shallow sounds between Sweden and Denmark provide a limited water exchange with the North Sea. The North Sea is located in the northeastern arm of the Atlantic Ocean, and between the British Isles and the mainland of northwestern Europe (Fig. 1; Fig. S1). It serves as a prominent shipping zone among European countries and between Europe and the Middle East. A third aspect of economic importance has been the extensive reserves of petroleum and natural gas discovered beneath the seafloor. Few parts of the North Sea are N90 m in depth.

The Bohai and Yellow Seas are semi-enclosed marginal seas in the northwest Pacific Ocean, and some of the most extensive shelf seas in the world (Fig. S1). The Bohai Sea is surrounded by the Bohai Economic Rim (BER), which is the largest economic engine in North China. The Bohai Sea has an area of approximately 77,000 km² and an average depth of 18 m (Song et al., 2014). It received many fluvial sediments especially from the Yellow River, and nearly 30% of the Yellow River-derived sediment has been ultimately transported out of the Bohai Sea into the Yellow Sea (Yang and Liu, 2007). The Yellow Sea is surrounded by the west coast of the Korean Peninsula and the east coast of China. It is connected to the Bohai Sea through the Bohai Strait, and its northern extension is referred to Korea Bay. The Yellow Sea has an area of approximatively 380,000 km² and an average depth of 44 m (Song et al., 2014).

2.2. Sample collection

A total of 101 surface sediment samples were collected from the four different seas, including 24 from the Baltic Sea, 27 from the North Sea (including 6 from the river Elbe Estuary, Germany) (Fig. 1), 11 from the Bohai Sea and 39 from the Yellow Sea (Fig. 2). The samples in the German Bight of the North Sea and Germany Baltic Sea coastline were collected on the research vessel Ludwig Prandtl in June and September 2017, respectively. While on the sampling campaign in the Chinese Bohai and Yellow Seas, the surface sediment samples were collected on the Chinese research vessel Dongfanghong 2 in June and July 2016, where a gridded sampling strategy was carried out. Stainless-steel box corers were used and the top 10 cm of sediment was taken on both the sampling campaigns. The samples were put afterward into the aluminun bowls (pre-cleaned with acetone and baked out at 250 °C) and stored at −20 °C until sample pretreatment.

2.3. Sample extraction, cleanup and instrumental analysis

The details about sample extraction, cleanup procedures and instrumental analysis were given in the Supplementary materials. Briefly, the frozen sediment samples were freeze-dried, homogenized and then extracted by Ultrasonic assisted extraction. The extract was cleaned up through a silica gel column and finally concentrated for instrumental analysis. PAH analysis was performed on a gas chromatograph/mass spectrometer (GC/MS) (Agilent 6890N GC/5973N MSD, USA), and 16 US EPA priority PAHs and 2 methylnaphthalenes (1- and 2-MN) were measured (Table S1).

2.4. Quality assurance/quality control (QA/QC)

16 deuterated internal standards (d-PAHs) were spiked in the samples for qualification and quantification, and 13C-labeled PCB-208 was added for recovery calculation. The average recoveries were 87 ± 27% in all the samples. The procedural (laboratory) blank was processed for each batch of 10 samples. Method detection limits (MDLs) were calculated as average blank plus three times standard deviation (SD) of the targets in the procedural blanks, and the values were in the range of 0.003–0.04 ng/g (Table S1).
3. Results and discussion

3.1. The overall concentrations and distribution

The sum concentrations of 18 PAHs (Σ18PAHs, including 1- and 2- MN) were in the range of 0.91–5361 ng/g dry weight (dw) with a mean of 549 ± 1124 ng/g dw in the sediments from the Baltic Sea, while they were in the range of 0.46–227 ng/g dw (mean 30.2 ± 57.2 ng/g dw), 25.0–308 ng/g dw (mean 184 ± 109 ng/g dw) and 4.3–659 ng/g dw (mean 138 ± 133 ng/g dw) in the North Sea, Bohai Sea and Yellow Sea, respectively (see Table 1). Relatively higher level was observed in the Baltic Sea, which may be attributed to the fact that most of the sampling sites were in close proximity to land and/or in bays or estuaries, and revealed the effect of proximity from source. Even so, they were generally comparable to those reported in the prev-ious studies (Baumard et al., 1999; Ricking and Schulz, 2002; SRU, 2004) (Table S2). The two highest values were obtained in the sampling sites L9–44 (209 ng/g dw) and L9–45 (5361 ng/g dw), both of which were located in the Flensburg Fjord, the northern end of Germany (Fig. 1). The main industries in Flensburg are shipbuilding, metal processing and paper making, which may lead to elevated emission of PAHs into the water system. On the other hand, the Flensburg Fjord is semi-enclosed with low water exchange rate against the Baltic Sea, probably increasing the deposition of PAHs into sediment. Higher concentrations were also found in the sites L9–6 (998 ng/g dw), and L9–20/21 (471/ 480 ng/g dw), which were near the port cities of Stralsund and Wismar, respectively. Furthermore, relatively higher concentration was ob- served in the site SH-01 (648 ng/g dw), which was located at the Peine River Estuary, Germany.

lower than those from the other three seas, but comparable to the mon- itoring data from the convention for the Protection of the Marine Envi-ronment of the North-East Atlantic (OSPAR) over the period of 2010–2015 (OSPAR, 2017). Whereas, they were lower than those in a special report by the German Advisory Council on the Environment in

<table>
<thead>
<tr>
<th>Compound</th>
<th>Baltic Sea (n = 24)</th>
<th>North Sea (n = 27)</th>
<th>Bohai Sea (n = 11)</th>
<th>Yellow Sea (n = 39)</th>
</tr>
</thead>
<tbody>
<tr>
<td>NaP</td>
<td>0.02–16.4</td>
<td>1.21 ± 3.30</td>
<td>0.36</td>
<td>0.009–176</td>
</tr>
<tr>
<td>2-MN</td>
<td>0.01–10.2</td>
<td>0.98 ± 2.91</td>
<td>0.34</td>
<td>0.00–4.3</td>
</tr>
<tr>
<td>1-MN</td>
<td>0.006–60.0</td>
<td>0.68 ± 1.41</td>
<td>0.25</td>
<td>0.00–1.79</td>
</tr>
<tr>
<td>Acy</td>
<td>0.01–20.1</td>
<td>2.79 ± 5.52</td>
<td>0.46</td>
<td>0.006–1.72</td>
</tr>
<tr>
<td>Ace</td>
<td>0.01–21.0</td>
<td>2.79 ± 5.52</td>
<td>0.46</td>
<td>0.006–1.72</td>
</tr>
<tr>
<td>Flu</td>
<td>0.04–97.7</td>
<td>3.49 ± 9.94</td>
<td>0.68</td>
<td>0.007–2.09</td>
</tr>
<tr>
<td>Phe</td>
<td>0.03–497.0</td>
<td>30.3 ± 100.1</td>
<td>6.40</td>
<td>0.005–100.9</td>
</tr>
<tr>
<td>Ant</td>
<td>0.009–28.1</td>
<td>3.94 ± 6.82</td>
<td>1.16</td>
<td>0.006–33.9</td>
</tr>
<tr>
<td>Fla</td>
<td>0.07–1663.0</td>
<td>117 ± 336</td>
<td>23.4</td>
<td>0.004–44.0</td>
</tr>
<tr>
<td>Pyr</td>
<td>0.11–970.0</td>
<td>92.2 ± 204.8</td>
<td>21.5</td>
<td>0.004–37.5</td>
</tr>
<tr>
<td>BaA</td>
<td>0.06–324.0</td>
<td>35.2 ± 69.9</td>
<td>9.30</td>
<td>0.002–15.7</td>
</tr>
<tr>
<td>Chr</td>
<td>0.07–472.0</td>
<td>50.6 ± 102.1</td>
<td>12.9</td>
<td>0.03–24.4</td>
</tr>
<tr>
<td>BbF</td>
<td>0.10–432.0</td>
<td>57.8 ± 105.3</td>
<td>15.2</td>
<td>0.04–18.9</td>
</tr>
<tr>
<td>BkF</td>
<td>0.08–297.0</td>
<td>44.2 ± 74.0</td>
<td>11.0</td>
<td>0.02–20.9</td>
</tr>
<tr>
<td>BaP</td>
<td>0.05–279.0</td>
<td>40.1 ± 71.9</td>
<td>11.3</td>
<td>0.01–17.5</td>
</tr>
<tr>
<td>InP</td>
<td>0.10–120.0</td>
<td>28.9 ± 34.3</td>
<td>15.1</td>
<td>0.04–15.0</td>
</tr>
<tr>
<td>DBA</td>
<td>0.02–269.0</td>
<td>57.6 ± 71.2</td>
<td>2.76</td>
<td>0.004–22.7</td>
</tr>
<tr>
<td>BaghiP</td>
<td>0.08–248.0</td>
<td>32.8 ± 55.7</td>
<td>12.0</td>
<td>0.01–5.83</td>
</tr>
<tr>
<td>Σ3+PAHs</td>
<td>0.91–5361.0</td>
<td>549 ± 1124.0</td>
<td>135.0</td>
<td>0.06–227.0</td>
</tr>
<tr>
<td>TOC (%)</td>
<td>0.05–613.0</td>
<td>21.6 ± 7.74</td>
<td>2.13</td>
<td>0.01–6.00</td>
</tr>
<tr>
<td>Water temperature (°C)</td>
<td>17.2–19.8</td>
<td>18.3 ± 0.8</td>
<td>18.1</td>
<td>13.5–21.1</td>
</tr>
<tr>
<td>Depth (m)</td>
<td>2.0–25.0</td>
<td>12.1 ± 7.8</td>
<td>11.0</td>
<td>1.8–40.8</td>
</tr>
<tr>
<td>Salinity (%)</td>
<td>1.1–18.0</td>
<td>12.3 ± 4.6</td>
<td>12.1</td>
<td>0.6–32.3</td>
</tr>
<tr>
<td>pH value</td>
<td>7.88–8.76</td>
<td>8.25 ± 0.18</td>
<td>8.23</td>
<td>7.75–8.25</td>
</tr>
</tbody>
</table>

Table 1
The concentrations of PAHs (ng/g dw) and other parameters in the sediments from the Baltic Sea, North Sea, Bohai Sea and Yellow Sea.

2004 (SRU, 2004), and those from the river Elbe Estuary, Germany (260–906 ng/g dw) (Otto et al., 2013) (Table S2). The two highest con- centrations were obtained from the sites L6–04 (227 ng/g dw) and L6–12 (187 ng/g dw), which were situated along the river Elbe and close to the Hamburg harbor and the Kiel Canal Estuary, respectively (Fig. 1). Moreover, higher concentrations were found in the sites L6–23 (96.3 ng/g dw) and L6–30 (74.3 ng/g dw), which were close to the PAHs concentrations in the North Sea, Bohai Sea.

In the Chinese seas, the PAH concentrations in the sediments from the Bohai Sea were generally higher than those from the Yellow Sea, and those in the north Yellow Sea were higher than in the south part (Fig. 2). Furthermore, PAHs showed an increasing tendency along the distance from the coastline toward the open sea, which was consistent with the previous observations (Lin et al., 2011; Li et al., 2015; Lin et al., 2017), as well as some other organic pollutants in this area (Apel et al., 2018; Mi et al., 2019). The fluvial sediments are transported, deposited and redistributed, forming coarse-grained sand, fine-grained mud and sand-mud mixture deposits on the continental shelf under in- tende tidal, wave and coastal conditions in the east China seas. (Yang and Liu, 2007; Qiao et al., 2017). However, all of the coastal currents generally flow southwards and warm currents enter northwards, resulting in the mud deposit areas located off the Yellow River and in the central Bohai Sea, around the end of the Shandong Peninsula, and in the central South Yellow Sea (Qiao et al., 2017). The higher PAH levels were commonly found in the mud deposit areas, which may reflect hy- drodynamic effect on spatial distribution of the deposited PAHs (Qiao et al., 2017). The PAH levels in the Chinese seas were generally compa- rable to those reported in the same areas (Lin et al., 2011; Li et al., 2015; Lin et al., 2017; Xu et al., 2019) (Table S2). The highest concentration (659 ng/g dw) was observed in the site ΩN843 in the north Yellow Sea, followed by the nearby site ΩN844 (424 ng/g dw), both of which were in the mud deposit area close to the Shandong peninsula.

This ob- servation was consistent with the previous studies on the organic UV stabilizers and UV filters (Apel et al., 2018), as well as phthalate esters (Mi et al., 2019) in the same area.

3.2. PAH profiles in sediment

Regarding the PAH profiles, 4-rings PAHs, e.g., fluoranthene (Fia), pyrene (Py), and benzo(b)fluoranthene (BbF) were commonly dominant compounds in the samples from all the seas, accounting for 14.1%, 11.8% and 12.1% of Σ18PAHs on average, respectively (Fig S2). This profile was generally consistent with many previous studies in these seas (Li et al., 2015; Lin et al., 2017; OSPAR, 2017).

Specifically, in the European seas, indeno(1,2,3-cd)pyrene (InP) made relatively higher contributions with a percentage of 9.8% on average, especially in the site L6–24 (22%), which was close to the island of tourist attraction in the North Sea. For the in- dividual site, relatively higher percentage of Fia. was observed in the sites L9–45 (31%) and L9–21 (25%) in the Baltic Sea. Moreover, although the levels of naphthalene (NaP)
and its alkylated derivatives (1- and 2-MNs) were relatively low in the sites L6–17/19/21/36 in the North Sea (b133 ng/g dw), they contributed evidently high with a sum percentage of about 20% \( \sum \text{PAHs} \) in each site, indicating a source characteristic of petrogenic origin (Yang et al., 2017).

In the Bohai and Yellow Seas, chrysene (Chr) generally accounted for higher percentages (mean 10.5%). This was different from the results by Li et al. (Li et al., 2015) and Lin et al. (Lin et al., 2017), where phenanthrene (Phe) was found as one of dominant compounds other than 4- rings PAHs. Furthermore, NaP showed relatively higher contributions (over 6.5%) in the sites C\H20/21/23, C\H35/36 and \( \text{C} \)\H41, as well as 1- and 2-MNs. These sites were generally close to the coastline of Jiangsu province, where highly developed manufacturing and shipping industries were located. The higher contribution of NaP and its methylated derivatives may therefore relate to the industrial emission in this area (ATSDR, 2005).

PCA was employed to further explore the PAHs distribution characteristic among the different seas. As exhibited in Fig. S3, 85.5% of the PAHs variation could be explained by PC1 (70.4%) and PC2 (15.1%) (Eigenvale N1). PC1 was characterized by the US EPA priority PAHs with NaP, while PC2 was featured by NaP and its methylated derivatives, i.e. 1- and 2-MNs. Regarding the score plot, L9–45, –44, and \( \text{C} \)\B43 were distinctively separated from the others. As mentioned above, both the sites L9–44 and –45 were located in the Flensburg Fjord, where much complex sources exist and may lead to different contamination characteristics from the others. Whereas, the site \( \text{C} \)\B43 was in the mud deposit area close to the Shandong peninsula, which probably induced the distinct distribution profile of PAHs in the adjacent marine system.

3.3. Source identification

PAHs encompass both parent (non-alkylated) compounds and alkylated homologues. Although PAHs can be produced through natural processes, they also arise from anthropogenic sources. Various combustion processes are the major source of PAHs, commonly comprising the heavier, parent (non-alkylated) PAHs, whereas they can also be of petrogenic origin (crude oils or refinery products) with mainly alkylated, 2–3 rings PAHs formed as a result of diagenetic processes.

The 16 US EPA priority PAHs were grouped into 2–3 rings low-molecular weight (LMW) and 4–6 rings high-molecular weight (HMW) PAHs, and the LMW/HMW ratio was generally employed for primary evaluation of PAH sources. In the present study, this ratio ranged from 0.08–0.81 in all the samples, suggesting a common source of pyrolytic origin. It was generally in agreement with previous studies in these seas (Lin et al., 2011; Jiao et al., 2012; Tobiszewski and Namieśnik, 2012; Louvado et al., 2015; Wang et al., 2015). One-way ANOVA analysis (Tukey HSD) indicated that significant higher ratios were obtained from the North Sea compared with those from the Baltic and Yellow Sea (p ≤ 0.001) (Fig. S5), which was in well agreement with the increasing tendency of PAH and TOC levels along the distance from the coastline toward the open sea, and confirmed the hydrodynamic effect on spatial distribution of the deposited PAHs.

Furthermore, diagnostic ratios of marker species were calculated. The majority of \text{Fla} (\text{Fla} + \text{Pyr}) and \text{InP} (\text{InP} + \text{benzo(g,h,i)-perylene (BghiP)}) ratios were N0.5 (Fig. 3), indicating pyrogenic source of bio- mass and coal combustion in most of the sampling sites (Guo et al., 2006). Some of these ratios were b0.5, but still higher than the threshold values of petrogenic origin, declaring the contribution from petroleum combustion. On the other hand, the lower benzo(a)anthracene (BaA) / (BaA + Chr) and anthracene (Ant)/(Ant+Phe) ratios ostensibly implied that PAHs in some sites especially the Chinese seas could be of petrogenic origin (Guo et al., 2006). Previous studies indicated that Ant and BaA have higher potential of photolysis than Phe and Chr, which made diagnostic ratios of BaA/(BaA + Chr) and Ant/(Ant+Phe) questionable during long range transport of PAHs (Yan et al., 2005; Deng et al., 2013; Lin et al., 2017). Despite that, river discharge and the coastal currents flowing southwards contributed much to the sediment in the two Chinese seas, and the sampling sites in the European seas were mainly along the coastlines. Therefore, the ratios of BaA/ (BaA + Chr) and Ant/(Ant+Phe) could to large extent distinguish PAH sources in the different seas. As exhibited in Fig. 3, the ratios from the Chinese seas were generally clustered together, while those from the Baltic and North Seas were more scattered, suggesting some distinct source characteristics between them. This phenomenon could also be analyzed by PCA analysis when using these ratios as variables (Fig. S6), where the sampling sites in the Baltic Sea were characterized by BaA/(BaA + Chr), Ant/(Ant+Phe) and InP/(InP + GghiP), while those in the North Sea were featured by \( \sum \text{LMW} / \sum \text{HMW} \) other than them. This further indicated discrepancy in the PAH sources between the two European seas, probably due to the effect of proximity from source, while similar sources in the two Chinese seas because of hydrodynamic effect.

NaP and its methylated derivatives are not only the natural constituents of mineral oil, but also produced commercially. They are also un- intentionally discharges during the burning of wood or tobacco, and
accidental spills (ATSDR, 2005). Whereas, little was reported about environmental fate of the methylated derivatives, i.e., 1- and 2-MNs (ATSDR, 2005). In this case, the composition of NaP and its methylated derivatives in the samples may provide information for the source identification. As Conde et al. (Conde et al., 2005) indicated, PAHs emission from combustion of agricultural and sylvicultural debris (i.e., biomass) were dominated by NaP, 2-MN, 1-MN, Phe and acenaphthylene (Acy), and the ratios normalized to NaP were about 0.4, 0.3, 0.25 and 0.2, respectively. Jenkins et al. (Jenkins et al., 1996) also found a signature of NaP and Phe during various biomass combustion, and almost all the ratios of 2-MN/NaP were lower than 0.5. While for the PAHs of petrogenic origin, the ratios of 1-MN/2-MN in the Bakken crude oils, diesel and gas-oline samples were lower than 0.75 as Yang et al. (Yang et al., 2017) investigated, Soliman et al. (Soliman et al., 2014) also obtained that the ratios of 2-MN/NaP and 1-MN/NaP was 0.58 and 0.36 on average in the Qatar coastal sediments, and 0.62 for 1-MN/2-MN as expected. In the present study, there were significant correlations between the con-centrations of 1-/2-MNs and NaP in all the seas (p < 0.001), and the ratios of 2-MN/NaP and 1-MN/NaP (slope of the regression line) were about 0.86 and 0.50, respectively (Fig. 4). The ratio of 1-MN/2-MN was expectedly 0.58, comparable to those of petrogenic source (Soliman et al., 2014; Yang et al., 2017), but obviously lower than 0.75 for biomass burning (Conde et al., 2005). Considering that Phe was not the domi-nant PAH compound in the sediments, it suggested that biomass com-bustion should not be the main source of PAHs in the majority of sampling sites, while petroleum residue may be ubiquitous in the European and Chinese seas. Regarding the site L9–45, the ratios of 2-MN/NaP and 1-MN/NaP were 1.32 and 1.18, resulting in a 1-MN/2-MN ratio of 0.89, distinctly higher than that from the nearby site L9–44 (0.65). Since L9–45 was characterized by the parent PAHs without NaP (Fig. S3), and Fla. and Pyr were the dominant compounds with lower concentrations of 5–6 rings PAHs, it suggested a source characteristic of coal combustion in this site (Levendis et al., 1998; Li et al., 2016). However, the site L–44 was featured by NaP and 1-, 2-MNs via PCA analysis, and benzo[k]fluoranthene (BkF) and benzo[a]pyrene (BaP) accounted for a higher proportion, suggesting a contribution from vehicle emission (Li et al., 2016). This was also in agreement with the distribution characteristic in the site CNB43 from the Yellow Sea.

PMF was commonly utilized to estimate contribution of the specific PAH source categories (Lin et al., 2011; Tobiszewski and Namieśnik, 2012; Li et al., 2016; Maletic et al., 2019). Details about PMF modeling process could be found elsewhere (Lin et al., 2011; Norris et al., 2014; Li et al., 2016). In this study, an empirical uncertainty of 20% was adopted as an estimate of the confidence level based on the results from regularly analyzing the standard reference material (Mai et al., 2003; Lin et al., 2011). Taking the background of the PAH release sources in these seas into consideration, different factor (3, 4 and 5) and seed numbers (20, 50 and 100) were tested at first, and the solution with 4 factors and 50 seeds was selected to yield the most stable and interpret-able results. Four typical PAH source profiles were finally selected refer-ring to acknowledged references (Khalili et al., 1995; Levendis et al., 1998; Marr et al., 1999; Larsen and Baker, 2003; Liu et al., 2009; Li et al., 2016) to verify the source contribution from PMF modeling, including 1) vehicular emissions, 2) coal combustion, 3) petroleum resi-due, and 4) coke plant. In order to compare with the previous studies, PMF modeling was performed only for 16 US EPA priority PAHs. As shown in Fig. 5, each factor had a similar contribution in the samples from the Baltic Sea, and 0.62 for 1-MN/2-MN as expected. In the present study, there were significant correlations between the concentration of 1-/2-MNs and NaP in all the seas (p < 0.001), and the ratios of 2-MN/NaP and 1-MN/NaP (slope of the regression line) were about 0.86 and 0.50, respectively (Fig. 4). The ratio of 1-MN/2-MN was expectedly 0.58, comparable to those of petrogenic source (Soliman et al., 2014; Yang et al., 2017), but obviously lower than 0.75 for biomass burning (Conde et al., 2005). Considering that Phe was not the domi-nant PAH compound in the sediments, it suggested that biomass com-bustion should not be the main source of PAHs in the majority of sampling sites, while petroleum residue may be ubiquitous in the European and Chinese seas. Regarding the site L9–45, the ratios of 2-MN/NaP and 1-MN/NaP were 1.32 and 1.18, resulting in a 1-MN/2-MN ratio of 0.89, distinctly higher than that from the nearby site L9–44 (0.65). Since L9–45 was characterized by the parent PAHs without NaP (Fig. S3), and Fla. and Pyr were the dominant compounds with lower concentrations of 5–6 rings PAHs, it suggested a source characteristic of coal combustion in this site (Levendis et al., 1998; Li et al., 2016). However, the site L–44 was featured by NaP and 1-, 2-MNs via PCA analysis, and benzo[k]fluoranthene (BkF) and benzo[a]pyrene (BaP) accounted for a higher proportion, suggesting a contribution from vehicle emission (Li et al., 2016). This was also in agreement with the distribution characteristic in the site CNB43 from the Yellow Sea.
Fig. 4. The relationships between NaP and 1–2-MNs (A), 1-MN and 2-MN (B) in the sediments from the different seas.

The results of source apportionment were generally consistent with the previous reports in the Chinese seas (Lin et al., 2011; Li et al., 2016) and the European seas (SRU, 2004; Lubecki and Kowalewska, 2010; OSPAR, 2010, 2017), where coal combustion, vehicular emission, petroleum residue and coke plant were commonly apportioned as the PAH sources. Whereas, limited data was available for their quantitative contributions in the marine sediments, especially in the European seas. In this case, the quantitative comparison between the different areas in the present study may shed light on the effect of proximity from source and hydrodynamic effect on spatial distribution of pollutants in the marine sediments.

4. Conclusions

In the present study, PAHs were intensively investigated in the sediments from four different seas including the European Baltic Sea and North Sea, Chinese Bohai Sea and Yellow Sea. Higher levels of \( \sum_{16}^{18} \text{PAHs} \) were detected in the Baltic Sea, while distinctly lower levels were measured in the North Sea. Combining with various approaches for source identification, coal combustion, petroleum residue, vehicular emission and coke plant were appointed as the main sources of PAHs in the surface sediments. Particularly, the ratios of NaP and its methylated derivatives (i.e. 1- and 2-MNs) evidenced that biomass combustion may not be the main source in the majority of sediments from these seas. This approach showed its great significance for source apportionment, and is in prospect of developing in the future study. Furthermore, different source contributions were quantified in the different seas, which may to some extent reflect the regional emission intensity of various PAH sources in the past decades.

One drawback is that, due to the limited tonnage of the research vessel Ludwig Prandtl, the majority of surface sediments from the European seas especially the Baltic Sea were obtained near the coastline, of which the results may reveal the effect of proximity from source. While the samples from the Chinese part were collected in the open sea because of the larger tonnage of Dongfanghong 2, of which the spatial distribution closely related to the hydrodynamic effect in this area. This may undervalue the quantitative comparison of source contributions in the different seas. More studies are warranted to further confirm the source contributions from the anthropogenic activities around these seas.
CRediT authorship contribution statement

Pu Wang: Writing - original draft, Methodology, Software, Formal analysis, Investigation, Data curation, Writing - review & editing, Visualization, Project administration, Funding acquisition.
Wenyi Mi: Methodology, Validation, Investigation.
Zhong Jie: Conceptualization, Methodology, Investigation, Resources, Data curation, Writing - review & editing, Visualization, Supervision, Project administration, Funding acquisition.
Jianhui Yang: Resources, Writing - review & editing, Funding acquisition.
Christina Apel: Resources.
Hanna Joens: Resources.
Ralf Ebinghaus: Writing - review & editing, Funding acquisition.
Zhengbo Qiang: Writing - review & editing, Funding acquisition.

Declaration of competing interest

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

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Appendix A: Supplementary data

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References
