Occurrences and Behaviors of Naphthenic Acids in a Petroleum Refinery Wastewater Treatment Plant

Beili Wang,† Yi Wan,*,† Yingxin Gao,‡ Guomao Zheng,† Min Yang,‡ Song Wu,§ and Jianying Hu†

†Laboratory for Earth Surface Processes, College of Urban and Environmental Sciences, Peking University, Beijing 100871, People’s Republic of China
‡Chinese Academy of Sciences, State Key Laboratory of Environmental Aquatic Chemistry, Ecoenvironmental Science Research Center, Beijing 100085, People’s Republic of China
§Petrochina Ji Dong Oilfield Company, Tangshan, 063200 Hebei, People’s Republic of China

ABSTRACT: Naphthenic acids (NAs) are one class of compounds in wastewaters from petroleum industries that are known to cause toxic effects, and their removal from oilfield wastewater is an important challenge for remediation of large volumes of petrochemical effluents. The present study investigated occurrences and behaviors of total NAs and aromatic NAs in a refinery wastewater treatment plant, located in north China, which combined physicochemical and biological processes. Concentrations of total NAs were semiquantified to be 113–392 μg/L in wastewater from all the treatment units, and the percentages of aromatic NAs in total NAs was estimated to be 2.1–8.8%. The mass reduction for total NAs and aromatic NAs was 15 ± 16% and 7.5 ± 24% after the physicochemical treatment, respectively. Great mass reduction (total NAs: 65 ± 11%, aromatic NAs: 86 ± 5%) was observed in the biological treatment units, and antiestrogenic activities observed in wastewater from physicochemical treatment units disappeared in the effluent of the activated sludge system. The distributions of mass fractions of NAs demonstrated that biodegradation via activated sludge was the major mechanism for removing alicyclic NAs, aromatic NAs, and related toxicities in the plant, and the polycyclic NA congener classes were relatively recalcitrant to biodegradation, which is a complete contrast to the preferential adsorption of NAs with higher cyclicity (low Z value). Removal efficiencies of total NAs were 73 ± 17% in summer, which were higher than those in winter (53 ± 15%), and the seasonal variation was possibly due to the relatively high microbial biotransformation activities in the activated sludge system in summer (indexed by O₃−NAs/NAs). The results of the investigations indicated that biotransformation of NA mixtures by the activated sludge system were largely affected by temperature, and employing an efficient adsorbent together with biodegradation processes would help cost-effectively remove NAs in petroleum effluents.

INTRODUCTION

Due to the increasing importance of petroleum for industrial development in the last decades, petrochemical effluents have had a significant impact on pollution of surface water systems. The amount of wastewater generated from the refining of petroleum is 0.5–1.6-fold greater than the volume of petroleum that is refined. Petroleum refinery wastewater (PRWW) comprises different amounts of hazardous pollutants, of which naphthenic acids (NAs) are a class of compounds that have been reported to be toxic and persistent, especially in the oil sands process-affected water (OSPW) in northeastern Alberta, Canada. Degradation of NAs in wastewater from the petroleum industry is an important challenge for remediation of the large volumes of petrochemical effluents.

Removal of NAs in wastewater treatment plants (WWTP) plays a crucial role in pollution control, and the treatment process of the PRWW generally consists of mechanical and physicochemical technologies including oil–water separation and coagulation, followed by biological treatment within the integrated activated sludge treatment plant. Laboratory studies have evaluated various technologies and processes for removing NAs in OSPW, such as biological treatment, membrane filtration, ozonation, and chemical coagulation, but much less attention has been given to treatment of PRWW. While nanofiltration could remove >95% NAs in wastewater and ozonation treatment achieved approximately 50% and 75% NA degradation in lab, biological treatment is the most efficient and cost-effective method in refineries. Recently, occurrences of NAs in activated sludge wastewater from six petroleum refineries in the United States has been investigated, but the fates of NAs remain unknown and there is still
a lack of understanding as to which processes in the WWTP are affecting the removal of NAs. It is well-known that use of mass balance analysis to understand behaviors of pollutants in WWTP can be useful for identification of mechanisms that are best suited for removal of pollutants and that assessment of efficiencies of different treatment units of a WWTP for removal of NAs could provide information needed to improve performance of these plants. However, little is known about the fate and major mechanisms for the removal of NAs in full-scale WWTPs.

Beside alicyclic “classical” NAs, aromatic NAs were recently identified by comprehensive two-dimensional gas chromatography–mass spectrometry (GC × GC-MS) of the methyl esters in OSPW.26–28 Although aromatic NAs make up a small percentage of NA mixtures (<10% in crude oils), they are the most environmentally significant components of the overall toxicity and recalcitrance of NAs found in wastewaters from the petroleum industry.29,30 For example, the aromatic NAs with chemical formulas similar to estrone- and estradiol-like compounds have been demonstrated to cause estrogenicity in fish larvae.31,32 Several biodegradation studies reported the microorganisms and mechanisms involved in degradation of individual aromatic NA compounds,30,33 however, there is very little understanding of aromatic NAs in water and solid fractions during physicochemical treatment and activated sludge treatment processes.

The present study investigated occurrences of total NAs (including both alicyclic and aromatic NAs) in aqueous and solid matrices in a petroleum refinery wastewater treatment plant, north China, and performed a mass balance of total NAs and aromatic NAs based on field measurements. The toxicities of wastewater samples from different treatment units were also measured by the yeast two-hybrid assay. The obtained field results were applied to assess the physicochemical and biological processes that may be effective for the removal of total NAs and aromatic NAs in wastewater. The seasonal variations of total NAs and their oxidized products (oxy-NAs) were further explored for clarification of the possible factors affecting the removal efficiencies.

### MATERIALS AND METHODS

#### Sample Collection.

The petroleum refinery wastewater treatment plant investigated in the present study is designed to treat 20000 m³/d wastewater generated by oil production platforms in Hebei Province, north China. The scheme of the refinery wastewater treatment plant and sampling locations are shown in Supporting Information (SI) Figure S1. The wastewater is first treated by physicochemical treatment units including gravity setting, coagulation, walnut shell filtration, and flotation. The hydraulic retention times were 3.5 h in each physicochemical treatment unit. The primary sludge from the gravity setting and coagulation chambers is pumped into the physicochemical treatment unit. The primary sludge from the secondary clarifier step, the effluent of the activated sludge reactor is discharged to the environment.

Occurrences and mass balance of NAs in the treatment plant were explored by analyzing NAs in wastewater, suspended solids, and sludge samples collected from different treatment units in the summer time (May 15, 2013 and May 15, 2014). The refinery processed light crude oil during the sampling periods. Seasonal variations of NAs in the plant were further studied by analyzing NAs in wastewater collected in winter (October 30, 2013). The characteristics of collected wastewater, suspended solids, and sludge samples are given in SI Table S1. For each sampling location, wastewater samples were collected every 3 h, and four samples were mixed as a 12 h composite water sample. The composite samples were collected twice and analyzed separately for each occasion (May 15–16, 2013, October 30–31, 2013, and May 15–16, 2014). The suspended solids were collected by filtering the water samples, and the glass microfiber filters (Whatman GF/C 1.2 μm, Maidstone, UK) used for filtering four composite water samples collected in each treatment unit in summer were extracted as one sample for analysis. The dewatered sludge and excess sludge were collected at the outlets of physicochemical treatment chambers and the activated sludge system, respectively, and sludge samples collected in summers were mixed and extracted as one sample for analysis. All water samples were collected in 500 mL amber glass bottles, which were washed by methanol and purified water before use. Water samples were extracted within 4 h in the local laboratory after being filtered. Sludge samples were stored at –20 °C until analysis. During the sampling period, the wastewater for treatment was around 13000 m³/d, and the average nitrogen concentrations in the secondary effluent were 0.02 mg/L for NO₂-N and 0.009 for NO₃-N. The other treatment parameters in the treatment plant (e.g., H₂SO₄, Sulfur Compounds (SCs), Suspend Solids (SS), and petroleum content) were listed in SI Table S2.

#### Sample Preparation.

The chemicals and reagents used in the present study are provided in the SI. The methods for semiquantification of NAs and oxy-NAs in wastewater and sediment samples have been reported previously.34,35 Briefly, approximately 250 mL of wastewaters spiked with 0.1 μg of surrogate standards (12-oxochenoodeoxycholic acid and 1-pyrenebutyric acid) were extracted with MAX cartridge (Oasis MAX, 6 mL, 150 mg, Waters, USA) which was preconditioned by use of 6 mL of methanol and 6 mL of purified water. The cartridges were rinsed with 6 mL of 5% ammonia and then dried under a flow of nitrogen. The MAX cartridge was precleaned with 6 mL of methanol which was discarded and then eluted with 12 mL of ethyl acetate saturated with hydrochloric acid (2 M HCl:ethyl acetate = 1:10, v/v). The eluate was washed three times with pure water and reconstituted with 100 μL of methanol for analysis by an ultrahigh-pressure liquid chromatograph (UPLC) coupled to a quadrupole time-of-flight mass spectrometer (QTOF-MS) with electrospray ionization in negative ionization mode (ESI−).

The sludge (about 0.5 g) and suspend solid (about 0.1 g) samples were first freeze-dried and then were spiked with 5 μg of surrogate standards (1-pyrenebutyric acid and 12-oxochenoodeoxycholic acid) before Soxhlet extraction with a total of 200 mL of hexane/MTBE/methanol (1:1:1, v/v) mixture solution for 24 h. The extracts were concentrated to approximately 5 mL by rotary evaporation and then dried under a gentle stream of nitrogen gas to approximately 3 mL to...
Table 1. Concentrations of NAs in Wastewater Collected from Different Treatment Units in a Petroleum Refinery Wastewater Treatment Plant, North China (μg/L)\textsuperscript{a}

<table>
<thead>
<tr>
<th>NAs</th>
<th>PW</th>
<th>GE</th>
<th>CE</th>
<th>WE</th>
<th>FE</th>
<th>AE</th>
<th>SE</th>
</tr>
</thead>
<tbody>
<tr>
<td>total</td>
<td>392 ± 113</td>
<td>289 ± 26</td>
<td>270 ± 36</td>
<td>263 ± 19</td>
<td>349 ± 17</td>
<td>123 ± 57</td>
<td>113 ± 82</td>
</tr>
<tr>
<td>Z = 0</td>
<td>51 ± 10</td>
<td>30 ± 18</td>
<td>30 ± 7.1</td>
<td>28 ± 6.6</td>
<td>32 ± 14</td>
<td>5.1 ± 4</td>
<td>6.3 ± 5.2</td>
</tr>
<tr>
<td>Z = −2</td>
<td>109 ± 21</td>
<td>91 ± 18</td>
<td>86 ± 5.7</td>
<td>84 ± 10</td>
<td>111 ± 18</td>
<td>20 ± 28</td>
<td>5.8 ± 1.9</td>
</tr>
<tr>
<td>Z = −4</td>
<td>109 ± 31</td>
<td>91 ± 10</td>
<td>84 ± 13</td>
<td>82 ± 7.5</td>
<td>109 ± 5.8</td>
<td>25 ± 26</td>
<td>15 ± 7.1</td>
</tr>
<tr>
<td>Z = −6</td>
<td>52 ± 31</td>
<td>35 ± 13</td>
<td>33 ± 11</td>
<td>33 ± 8.3</td>
<td>50 ± 4.8</td>
<td>22 ± 8.1</td>
<td>24 ± 19</td>
</tr>
<tr>
<td>Z = −8</td>
<td>31 ± 23</td>
<td>19 ± 5.8</td>
<td>16 ± 5.1</td>
<td>16 ± 3.3</td>
<td>21 ± 3.5</td>
<td>17 ± 9.2</td>
<td>20 ± 17</td>
</tr>
<tr>
<td>Z = −10</td>
<td>19 ± 13</td>
<td>11 ± 3.6</td>
<td>10 ± 3.2</td>
<td>10 ± 2.3</td>
<td>13 ± 3.0</td>
<td>12 ± 6.1</td>
<td>14 ± 11</td>
</tr>
<tr>
<td>Z = −12</td>
<td>13 ± 11</td>
<td>7.5 ± 3.2</td>
<td>6.6 ± 2.8</td>
<td>6.5 ± 1.9</td>
<td>8.1 ± 2.6</td>
<td>12 ± 8.7</td>
<td>16 ± 13</td>
</tr>
<tr>
<td>Z = −14</td>
<td>8.9 ± 8</td>
<td>4.6 ± 2.2</td>
<td>4.1 ± 2</td>
<td>4 ± 1.4</td>
<td>4.7 ± 2.1</td>
<td>10 ± 8.6</td>
<td>13 ± 13</td>
</tr>
</tbody>
</table>


ensure the complete removal of hexane and MTBE. The residues were dissolved in 30 mL of water and then purified by the same method of solid-phase extraction for water sample preparation (Oasis MAX cartridges, 6 mL, 150 mg, Waters, Milford, MA). The elute of cartridge was washed with pure water to reduce the pH levels toward neutral and reconstituted with 500 μL of methanol for UPLC-ESI–QTOF-MS analysis.

Since UPLC-QTOF-MS analysis cannot differentiate aromatic NAs from alicyclic acids, aromatic NAs were isolated and semiquantified using a previously developed method.\textsuperscript{28} Aliquot of elute was derivatized with BF₃-methanol, and separation of aromatic NAs from alicyclic NAs was then achieved by using 6 mL Discovery Ag-Ion SPE cartridges (750 mg sorbent; Sigma–Aldrich, Dorset, UK). The isolated aromatic NA esters were analyzed by gas chromatography–mass spectrometer (GC-MS). Details of sample preparations for aromatic NAs and parameters for instrumental analysis of UPLC-QTOF-MS and GC-MS were provided in the SI.

Extracts of water and suspend solids in wastewater samples collected from different treatment units were combined and redissolved in dimethyl sulfoxide (DMSO) to assess the estrogenic and antitestrogenic activities using the yeast two-hybrid assay.\textsuperscript{46} The details of the bioanalysis were provided in the SI.

Quality Assurance and Quality Control (QA/QC). All equipment were rinsed with acetone and hexane to avoid sample contamination, and laboratory blanks were conducted for every batch of samples. In this study, total NAs in field blank samples were determined to be <10 μg in water and <1.3 μg in sludge samples. Efficiencies of the sample preparation procedure were assessed by analyzing sludge and wastewater samples collected from each treatment units spiked with standard solutions of model NAs and oxy-NAs. The absolute recoveries of model NA and oxy-NA compounds in triplicate spiked samples were 80 ± 17%, 84 ± 16%, 80 ± 15%, 83 ± 19%, 87 ± 17%, 89 ± 17%, and 89 ± 18% in petroleum wastewater, gravity settling effluent, coagulation effluent, walnut shell effluent, flotation effluent, A/O effluent, and secondary effluent, respectively, and 71 ± 19% for sludge samples. Currently, concentrations of NA mixtures were generally semiquantified based on integration of the hump peak of each NA congener assuming that the responses for individual compounds in the hump peaks were similar,\textsuperscript{12,37–41} since the separation method and standards for all the individual NAs were not available. While the commercial NAs and oil field NAs may contain NA congeners with different structures and response factors,\textsuperscript{42} semiquantification of the analytes was achieved using an internal standard method with calibration against standard solutions of commercial NAs (Acros) reported previously.\textsuperscript{3,25,34} NAs and oxy-NAs were semiquantified in sample extracts relative to 1-pyrenebutyric acid and 12-o xochoenodeoxycholic acid, respectively. Surrogate standards were spiked to samples prior to extraction to compensate for the loss of target compounds during the extraction process and correct the variation of instrument response and matrix effect, although the response variability would still occur in the analysis. Recoveries of 1-pyrenebutyric acid and 12-o xochoenodeoxycholic acid were 76 ± 12% and 70 ± 11% in all water samples and 74 ± 14% and 71 ± 17% in all suspended solid samples. The method detection limits (MDL) were based on the peak-to-peak noise of the baseline near the analyte peak obtained by analyzing field samples and on a minimum value of 3 for signal-to-noise for model NA and oxy-NA compounds. The MDLs of the model compounds ranged from 0.7 to 15 ng/L in wastewater samples and from 0.1 to 1.0 ng g\textsuperscript{−1} dw in suspended solid samples. Mass ions of 119 and 105 were selected for semiquantification of aromatic NAs relative to the surrogate (1-pyrenebutyric acid),\textsuperscript{26} and percentages of aromatic NAs in total NAs were estimated assuming that the responses for NAs and aromatic NAs were similar to that of 1-pyrenebutyric acid in UPLC-QTOF-MS and GC-MS analysis, respectively. The data analysis including the calculation of mass flow, solid–water partition coefficient (K\textsubscript{ow}, L/kg), and contribution of sorption and degradation to the removal of NAs in the plant was provided in the SI.

## RESULTS AND DISCUSSION

### Occurrence

Table 1 shows the concentrations of total NAs (including both alicyclic and aromatic NAs) in wastewater samples collected from various treatment units in a petroleum refinery wastewater plant. Concentration of total NAs was 392 ± 113 μg/L in the crude petroleum wastewater, and the percentage of aromatic NAs in total NAs was estimated to be 60%, suggesting that most NAs (>90%) detected in the treatment plant were alicyclic NAs. The relative composition of NA mixtures compared to the sum of all ions observed in the negative-ion ESI full scan mass spectra was 70 ± 12% in the crude petroleum wastewater, suggesting that NA mixtures were dominant contaminants in the wastewater. Concentrations of total NAs in the crude petroleum wastewater in the present study were much lower than those in petroleum wastewaters (4.5–16.6 mg/L) from six United States refineries,\textsuperscript{42} of which concentrations of NAs were semiquantified with the same method, and the possible reason could be the different type of...
crude oils. The dominant ions in the crude oil refinery wastewater in the present study were for \( Z = -2 \) and \( Z = -4 \) series, accounting for 55.6% of the total concentrations, followed by \( Z = -6 \) (13.3%), \( Z = 0 \) (13.0%), \( Z = -8 \) (7.9%), \( Z = -10 \) (4.8%), \( Z = -12 \) (3.3%), and \( Z = -14 \) (2.3%), which all centered around \( n = 10 - 20 \) (Figure 1a). The profile with a high abundance of one-three ring NAs with 10-20 carbons in crude oil wastewater were similar to that of crude oil total extractable NA extract in six United States refineries, with the dominant ions of \( Z = -2 \), \( Z = -4 \), and \( Z = -6 \) and \( n = 10 - 20 \), except for that the proportions of high molecular weight NAs (\( n = 25 - 35 \)) were relatively high in the present study.

Concentrations of total NAs in physicochemical treatment units were 289 ± 26 \( \mu g/L \), 270 ± 36 \( \mu g/L \), 263 ± 19 \( \mu g/L \), and 349 ± 17 \( \mu g/L \) in the effluent of gravity settling, coagulation, walnut shell, and flotation, respectively. About 26.3% of total NAs were removed after the treatment by gravity settling, and concentrations of total NAs only decreased by 2.2-6.6% due to the treatment of coagulation and walnut shell units, suggesting that adsorptions in the gravity setting unit would slightly help the initial removal of total NAs in the wastewater. The profiles of total NAs were very constant in wastewater from the physicochemical treatment units, with the dominant ions of \( Z = -2 \), \( Z = -4 \), and \( Z = -6 \) and \( n = 10 - 20 \) (Figure 1b, c, d, and e). The profiles are differentiated from that of crude oil wastewater by the extreme low percentage of NAs with high carbon numbers (\( n = 25 - 35 \)), which have been removed by the gravity settling unit. The removal efficiencies of different NA congeners by gravity setting increased linearly from 16.4 to 47.9% with the Z value decreasing from \(-2 \) to \(-14 \) (most alicyclic NAs, >90%), indicating that high cyclization NAs could be relatively easily removed by adsorption during the gravity setting processes.

Concentrations of total NAs in effluent of the activated sludge system were 123 ± 57 \( \mu g/L \), and the removal efficiencies for the total NAs by the A/O process were 65 ± 16%, suggesting that the A/O process treatment could effectively remove most NAs in the treatment plant. The removal efficiencies of different NA congeners by the A/O process treatment increased linearly with an increase of the Z value. This indicated that low cyclization NAs are relatively easily removed in the activated sludge system, resulting in the relatively high proportions (8.1-13.8%) of high cyclization NAs (\( Z = -14 - 8 \)) in the effluent of the activated sludge system (Figure 1f).

Concentrations and profiles of NAs detected in the suspended solids and sludge samples in the treatment plant are given in SI Table S3, Figure S2, and Figure S3. The total concentrations of NAs were 860-2772 mg/kg in suspended solids from all the treatment units and 1594-2489 mg/kg in sludge samples from gravity settling and A/O treatment units, of which the low variation of the concentrations suggested the constant deposit of NAs in the suspended solids and sludge (SI Table S3). The profiles with dominant ions of \( Z = -8 - 2 \) were found for all the solid and sludge samples, but the carbon...
numbers centered around 20–33 in sludge were different from those in suspended solids, in which NAs with high molecular weight and carbon number (33–40) were generally detected with relatively high abundance (SI Figures S2 and S3). The solid–water partition coefficients \(K_p\) \(L/kg\) for different classes of NA congeners were further calculated, and \(K_p\) of total NAs was in the range of 2194–12715 in all the treatment units (Table 2). It is interesting to note that \(K_p\) decreased with the Z value of NAs (most alicyclic NAs, >90%) in all the physicochemical treatment units, but this phenomenon disappeared in the A/O process unit. The results indicate that high cyclization NAs were relatively easy to remove by adsorption, which is consistent with the positive correlation between the removal efficiencies and cyclization of NAs in the gravity settling unit, and the sorption behaviors were greatly affected in the A/O process.

The total NA concentrations (113 ± 82 \(\mu\)g/L) for secondary effluent were similar to those in effluent of the activated sludge system, and the removal efficiencies, calculated by comparing concentrations in the raw sewage and secondary effluent, were 73 ± 17% for total NAs. A recent study reported occurrences of NAs in oil refinery activated sludge wastewater treatment systems in six United States refineries, but the different type of crude oil hampered a direct comparison of NAs for the same refinery.\(^{25}\) The present study found relatively high removal efficiencies of NAs in a refinery wastewater treatment plant, and the removal efficiencies of different NA congeners decreased linearly with the increased cyclization of NAs, which is similar to the structure-persistence relationship of NAs in the A/O process and different from the adsorption behaviors of NA congeners, possibly suggesting the relatively high contribution of the activated sludge system to the removal of NAs in the treatment plant.

### Mass Flow and Mass Balance

Mass flows and mass balance of total NAs were determined to assess their potential removal mechanisms in a petroleum refinery wastewater treatment plant (Figure 2 and SI Table S4). In petroleum wastewater, the combined aqueous and solid phase mass flows of total NAs were about 7725 ± 1471 g/d, and mass flows entering the wastewater treatment plant were 5097 g/d and 2628 g/d in the water-phase and sorbed fraction, respectively. The proportion of total NAs sorbed on particles was 34% of that in the water-phase, suggesting that NAs were discharged to the environment mainly in aqueous phase although these compounds have high values of \(K_p\) (Table 2). In the physicochemical treatment units, mass change percentages ranged from 5.3 to 10.6% in coagulating, walnut shell, and flotation treatment units. The low mass change percentage and variability in the treatment units indicated that adsorption was of minor importance in the removal of total NAs in the plant. Total NAs mass flow entering the activated sludge system was about 4536 ± 150 g/d, and the great mass reduction percentage was observed in both the water-phase (64.8%) and sorbed fraction (65.7%) in the treatment unit. The high mass reduction in the activated sludge system suggested that the A/O process greatly contributed to the removal of NAs. For filtered secondary effluents, the mass flow was about 1016 ± 735 g/d, and the sorbed amount was ignored due to the low concentrations of suspended solids in the effluent. The total NAs mass in the dewatered and excess sludge was 498 and 9.6 g/d, respectively.

To assess the contribution of sorption and degradation to the removal of total NAs in the plant, the mass balance of NAs in the wastewater treatment plant was expressed in chemical mass fractions (%) detected in (i) secondary effluent, (ii) dewatered and excess sludge, and (iii) total lost, relative to the calculated

![Figure 2](image-url)  
**Figure 2.** Mass flows (g/d) of total NAs and percentages of aromatic NAs in total NAs (%) in a petroleum refinery wastewater treatment plant, north China.
initial loading (100%) (Figure 3). The calculated fraction of mass losses due to degradation for total NAs accounted for 74.4% of initial loadings, while the contribution of sorption and output of sludge was much less (6.6%). The mass proportion of total lost for different NA congeners increased linearly from 36 to 90% with increasing of Z values, which is consistent with the fact that NAs with increased cyclicity are more recalcitrant than lower cyclization NA during the biodegradation processes.5,10,13,14 (Figure 3). The results demonstrated that biodegradation via activated sludge treatment system was the major removal mechanism of NAs in the plant.

Mass flows and mass balance of aromatic NAs were further determined since the aromatic acids might represent a toxic and/or recalcitrant fraction of NA mixtures. As shown in Figure 2, mass flows of aromatic NAs entering the wastewater treatment plant were 306 g/d and 29 g/d in the water-phase and sorbed fraction, respectively. In the physiochemical treatment units, the percentage of aromatic NAs in total NAs were constant in wastewater (6.0–8.8%), and mass flows of aromatic NAs were in the range of 246–309 g/d (Figure 2). The mass flow of aromatic NAs entering the activated sludge system was about 306 g/d, and great mass reduction (92%) was observed in water-phase. Similar high mass reduction in the A/O treatment unit (97%) was also found for individual aromatic NAs (C11H16O2, selected ions: 133, 105, 7726), of which mass flows were shown in SI Figure S4. In the mass balance analysis, the fraction of mass losses due to degradation for aromatic NAs accounted for 87.4% of initial loadings, and the contribution of sorption and output of sludge was only 0.78%. The results demonstrated that aromatic NAs, similar to total NAs, can be greatly removed by biodegradation in the A/O treatment unit. This conclusion is consistent with the results of microbial biodegradation of individual aromatic alkanoic NAs.30,33 Furthermore, estrogenic activities in wastewater from different treatment units were determined, since the aromatic NA fraction was reported to have an estrogenic effect.31,32 Estrogenic activities were not observed in the sample extracts possibly due to the low concentrations of NA mixtures, but antiestrogenic activities were found in wastewater from physiochemical treatment units with ER antagonist potency ranging from 750 to 2100 ng Tamoxifen/L (Figure 4 and SI Figure S5). It should be noted that the antiestrogenic activities disappeared in the A/O process effluent and secondary effluent, suggesting the effective removal of the toxic compounds by the activated sludge system (Figure 4). The results further confirmed that biodegradation was the major mechanism for removing alicyclic NAs, aromatic NAs, and related toxicities in the plant.

It is interesting to note that the NAs with low Z values (most alicyclic NAs, >90%) were relatively recalcitrant to biodegradation (Figure 3), which is a complete contrast to the preferential adsorption of NAs with less negative Z. Previous studies have tested various types of adsorption materials in a laboratory to help remove the persistent fraction of NAs.21–24 Thus, employing an efficient adsorbent together with biodegradation processes within the integrated activated sludge treatment plant would help to cost-effectively and efficiently remove most NAs including toxic and recalcitrant fractions in wastewater to meet the no-discharge policy for the compounds.

**Seasonal Differences.** Concerning the possible seasonal difference in the removal efficiencies of NAs, seasonal variations in concentrations of total NAs in the PRWW were investigated. Concentrations of total NAs were 337 ± 17, 312 ± 30, 293 ± 9.4, 234 ± 7.0, 248 ± 30, 219 ± 16, and 159 ± 58 μg/L in petroleum wastewater, gravity settling effluent, coagulation effluent, walnut shell effluent, flotation effluent, A/O process effluent, and secondary effluent, respectively, in winter. While the concentration and profile of NAs were similar in the two seasons, the removal efficiencies of total concentrations of NAs in winter (53 ± 15%) were relatively low compared with those in summer (73 ± 17%) (Figure 5a, p = 0.228, t test), and the negative average removal efficiencies for Z = −12 and −14 in summer were observed possibly due to the wide variations among four times of sampling. The removal efficiencies of NAs in the physiochemical treatment units, calculated by comparing concentrations in petroleum wastewater and flotation effluent, were both relatively low in summer and winter (<30%), which is consistent with the low contribution of physiochemical treatment to the removal of NAs in the plant; and the removal efficiencies of the activated sludge system in winter (11 ± 17%) was significantly lower than those in summer (65 ± 16%) (p < 0.05, t test), suggesting that the activated sludge units are more effective in decreasing the NA
concentrations in the summer time and temperature would affect the removal efficiencies during the treatment.

Oxy-NAs, considered as a potential marker for the degradation of NAs in aquatic samples, were observed during aerobic microbial biodegradation of commercial NAs. Concentration ratios between oxy-NAs and NAs has been successfully applied to the assessment of weathering characteristics of NAs in the oil spilled area. Thus, the concentration ratios (O₃-NAs/NAs) were used to assess the biodegradation of NAs in the petroleum refinery wastewater plant. In winter, the concentration ratios were similar in different treatment units with values ranging from 0.26 to 0.36 (Figure 5b). However, the concentration ratios in summer remained constant in physicochemical treatment units (0.15–0.19) and shift to 0.71–0.82 after the active sludge system (Figure 5b), suggesting the high microbial biotransformation activities of the A/O processes in summer, which is consistent with the high removal efficiencies in the treatment unit. The results confirmed that biodegradation via activated sludge was the major process contributing to the removal of most NAs including toxic and recalcitrant fractions, and the temperature would greatly influence the treatment efficiencies.

Overall, the present study reported for the first time the mass balance of total NAs and aromatic NAs in a petroleum refinery wastewater plant. The high cyclization NAs were relatively easily subject to adsorption, which slightly helps the initial removal of NAs in the wastewater. The distribution of mass fractions of NAs in the plant indicated that biodegradation was the major removal mechanism of NAs (including aliphatic NAs, aromatic NAs, and related toxicities) in the activated sludge system, and the removal efficiencies by the processes was largely affected by temperature. The results of the investigations indicated that employing an efficient adsorbent together with optimized biodegradation processes would help cost-effectively and efficiently remove NA mixtures in wastewater to meet the no-discharge policy for the pollutants.

**ASSOCIATED CONTENT**

Supporting Information
Text, figures, and tables addressing (1) chemicals and reagents and instrument analysis; (2) analysis of aromatic NAs; (3) assay of agonist and antagonist; (4) calculations of mass flow, mass proportions, and solid–water partition coefficients; (5) characteristics of collected wastewater, suspended solids, and sludge samples; (6) treatment parameters in the petroleum refinery wastewater treatment plant; (7) concentrations of NAs (mg/kg) in suspended solids and sludge samples; (8) mass flux (g/d) of NAs in the petroleum refinery wastewater treatment plant; (9) sample locations in the plant; (10) profiles of NAs in suspended solids, activated sludge, and dewatered sludge; (11) mass flows (g/d) of individual aromatic NAs (C₈H₁₀O₂); and (12) dose–response curves of antiestrogen activities in sample extracts. This material is available free of charge via the Internet at http://pubs.acs.org.

**AUTHOR INFORMATION**

*Corresponding Author*
Phone/Fax: 86-10-62759126. E-mail: wany@urban.pku.edu.cn. Corresponding author address: College of Urban and Environmental Sciences, Peking University, Beijing 100871, China.

**Notes**

The authors declare no competing financial interest.

**ACKNOWLEDGMENTS**

The research is supported by National Natural Science Foundation of China (21177003, 21422701) and State High Technology Research and Development (863) Project of China (2012AA063401).

**REFERENCES**


(30) Johnson, R. J.; Smith, B. E.; Sutton, P. A.; McGenity, T. J.; Rowland, S. J.; Whitby, C. Microbial biodegradation of aromatic naphthenic acids is affected by the degree of alkyl side chain branching. ISME J. 2011, 5, 486–496.


(41) Clemente, J. S.; Prasad, N. G. N.; Mackinnon, M. D.; Fedorak, P. M. A statistical comparison of naphthenic acids characterized by gas

