Development of a novel integrated membrane system incorporated with an activated coke adsorption unit for advanced coal gasification wastewater treatment

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**HIGHLIGHTS**

- Activated coke (AC), a low-cost substitute to the more expensive activated carbon, is firstly used as the pretreatment of integrated membrane processes for the advanced coal gasification wastewater treatment.
- The effect of AC adsorption pretreatment on the UF, RO, and MD units is investigated.
- The mechanism that AC adsorption alleviates membrane fouling is discussed.
- TOC, FEEM, MWD, and GC–MS give a systematic characterization of water quality.
- The techniques that activated coke used as pretreatment of integrated membrane processes has been applied in a local coal gasification plant in China.

**ABSTRACT**

Lurgi–Ruhrgas coal gasification process is commonly used to produce methane in many regions. However, such a process produces wastewater containing toxic pollutants that are difficult to remove by the conventional integrated membrane system consisting of a coagulation unit, an ultrafiltration (UF) unit, a reverse osmosis (RO) unit, and a membrane distillation (MD) unit. In this regard, an activated coke was added prior to the UF unit in order to remove the organic matters and meet the wastewater discharge regulation. The activated coke is an abundant, low-cost substitute to the more commonly used expensive activated carbon. The effect of the adsorption unit on the removal efficiency and membrane fouling...
1. Introduction

Lurgi–Ruhrgas coal gasification process is commonly used to produce large quantity of syngas (methane as the primary component) in the regions rich in coal but poor in oil or gas (e.g., China) [1,2]. The wastewater discharged from the Lurgi–Ruhrgas gasifier usually contains large amount of toxic organic and inorganic pollutants, including phenolic compounds, polynuclear aromatic hydrocarbons, heterocyclic compounds, ammonium, sulfate, cyanide, and thiocyanate which are required to be removed to meet the wastewater discharge regulation [3,4]. However, most of the currently available treatment processes, such as anoxic–oxic (A/O) and anaerobic–anoxic–oxic (A²/O), were proven to be insufficient to reduce these contaminants to an acceptable level [5,6]. In this regard, the need for an effective and efficient process to treat the coal gasification wastewater is certainly warranted.

Among all the research efforts aiming for maximally reducing the contaminant contents in the coal gasification wastewater, the so-called “integrated membrane system (IMS)” combining various types of membrane processes and other separation processes has received considerable attention mainly because such an integrated configuration synergistically enhances the advantages of each individual process, and subsequently improves the contaminant removal efficiency from the coal gasification wastewater [7–12]. A typical IMS configuration for coal gasification wastewater treatment consists of a coagulation pre-treatment unit for colloids removal, an ultrafiltration (UF) unit to remove suspended solids, a reverse osmosis (RO) unit for metal ions and small organic matters removal, and a complimentary membrane distillation (MD) unit to further increase the global water recovery of the process. However, despite the great benefits derived from this process, the undesired fouling behavior commonly observed in the UF, RO, and MD unit processes causes a substantially loss in clean water productivity and quality, thus impeding the further advance of IMS in coal gasification wastewater treatment, where large quantity of wastewater is required to be purified and strict discharge regulation needs to be complied with [13–15]. With this concern, developing an appropriate strategy and IMS configuration not only delivering higher contaminant removal efficiency but also suppressing the undesirable membrane fouling phenomenon is of great importance for the IMS process.

The organic matters present in the coal gasification wastewater are believed to be the main foulants that (i) foul the UF membrane thus causing the flux decline in the UF process, (ii) wet the RO membrane surface and consequently lower the salt rejection, and (iii) accumulate on the MD membrane surface thus encouraging the crystallization and scaling of the metal ions on the membrane surface, and consequently lead to a lowered salt rejection. As such, the addition of a pre-treatment step to eliminate these organic foulants thus minimizing the fouling propensity in each individual membrane unit process has become a research focal point and several separation techniques have been considered as the potential pre-treatment candidates. Among them, the use of activated carbon to adsorb contaminant compound (mainly organic matters) in the wastewater shows great process flexibility and potential to be incorporated in the IMS process as a pre-treatment to safeguard the subsequent membrane processes. However, several issues associated with the activated carbon substantially limit the feasibility of using such a material in the wastewater treatment, including: (i) the difficulty to regenerate the activated carbon, and (ii) the high cost of these adsorbents [16–19]. In this regard, it is essential to search for a low-cost, more efficient and easily accessible adsorbent. Activated coke, a low-cost adsorbent, is often used as a substitute adsorbent for activated carbon, due to (i) its excellent SO₅ and NOX adsorption ability in the gas phase, and (ii) its exceptional adsorption capacity to remove organic pollutants from the liquid phase. Furthermore, activated coke is produced from naturally occurring carbonaceous materials such as lignite, petroleum coke, wood and other biomass [16,18,20,21]. Therefore the abundant resource of activated coke in a coal gasification plant guarantees a secured and low-cost supply of such absorbent in the IMS process. However, to the best knowledge of the authors, no studies have yet been reported on the incorporation of an activated coke adsorption unit in an IMS process for the coal gasification wastewater treatment.

In this study, the addition of an adsorption unit with activated coke as a pre-treatment step prior to the UF unit process in a typical IMS configuration was studied. Real coal gasification wastewater provided by a local coal gasification plant instead of synthetic model wastewater was used to demonstrate the feasibility of using such a system in an industrial-relevant environment. The effect of the activated coke adsorption pre-treatment on (i) contaminant removal, and (ii) membrane fouling mitigation was systematically examined by comparing the performance of two different IMS configurations (with and without activated coke adsorption unit process). The effluent properties of each unit process were characterized by total organic carbon (TOC), fluorescence excitation-emission matrix (FEEM), molecular weight distribution (MWD), and gas chromatograph/mass spectroscopy (GC–MS) analysis. In addition, the surface morphology of the activated coke as well as the fouling layer formed on the membrane surface was characterized by the scanning electron microscopy (SEM).

2. Materials and methods

2.1. Materials

The wastewater influent used in the experiments was provided by Yima gasification plant (Henan, China). The existing treatment processes in the plant included a phenolic compounds extraction unit, an ammonia stripping unit, and a sequencing batch reactor (SBR) treatment. After being treated by a series of chemical and biological treatments, the wastewater was still deep brown in color with a bad odor. The chemical and physical properties of the wastewater are showed in Table 1.

The activated coke from lignite was obtained from the same gasification plant, with particle sizes ranging from 0.038 mm to 0.15 mm, and the Brunauer–Emmett–Teller (BET) surface area of
Table 1
Water quality parameters of the raw wastewater.

| Parameter | Value   | Parameter | Value   |
|-----------|---------|-----------|---------|
| pH        | 7.1–7.4 | Ca²⁺ (mg/L)| 7.14–7.57 |
| Conductivity (μS/cm) | 1452–1777 | Mg²⁺ (mg/L)| 6.11–6.56 |
| TDS (mg/L) | 725–888 | Al³⁺ (mg/L)| 1.78–2.04 |
| Turbidity (NTU) | 44.5–73.7 | Cl⁻ (mg/L) | 48.4–50.8 |
| CODₐₐ (mg/L) | 342–503 | SO₄²⁻ (mg/L)| 463–486 |
| TOC (mg/L) | 138–187 | NO₃⁻ (mg/L)| 11.5–12.8 |
| NH₃-N (mg/L) | 8.60–13.2 | SiO₂ (mg/L)| 18.3–18.5 |

350 m²/g (measured by the N₂ adsorption isotherms using an ASAP 2000 Micromeritics instrument) (Micromeritics Instrument Corporation, USA). The activated coke was washed with deionized water for several times to remove the contaminants attached on the surface until the washed water became visibly clear. The activated coke was then dried in an oven at 105 °C for 48 h and stored in an air-tight glass bottle for further use.

2.2. Experimental

2.2.1. Process 1: Coagulation-UF-RO-MD

The Process 1 consisted of a coagulation unit, followed by a UF unit, a RO unit, and finally a MD unit, a schematic representation of the entire IMS process is provided in Fig. 1.

In the coagulation unit, polyaluminum chloride (PAC, flocculant) and polyacrylamide (PAM, coagulant) were added with a dosage of 250 mg/L and 1.0 mg/L, respectively. The coagulation process was performed on an agitation apparatus (MY3000-6G, Meijiu Environmental Equipment Corporation, China). PAC was firstly added in the wastewater (1 L), after 1 min of vigorous mixing at 250 rpm, PAM was then added in the wastewater. The wastewater samples were then subjected to a three-step agitation: a 1 min rapid mixing at 150 rpm, followed by a 10 min slow mixing at 60 rpm and a 5 min slow mixing at 30 rpm. The samples were then allowed for precipitation for 30 min, the supernatant was collected at the end of the 30 min precipitation and used as the feed for the UF process.

A polysulfone (PSf) hollow fiber UF membrane module (GHF-3540, Gohigher Environmental Tech, China) with a molecular weight cut-off (MWCO) of 20–30 kDa was used in the UF unit process in a cross-flow configuration at 0.50 bar. The retentate stream of the UF process was recirculated back to the feed tank, whilst the permeate stream was used as the feed of the subsequent RO process.

In the RO unit, a laboratory-scale cross-flow filtration setup (CEPA CF II, GE Osmonics, USA) consisting of a feed tank, a pump, a pressure gauge and a membrane filtration cell was used, and the flat-sheet RO membrane (BW30-4040) was provided by FilmTec Corporation, USA. The new membrane was pre-compacted for 24 h by filtering with deionized water until a steady-state flux was obtained. The feed solution was stored in a 4 L reservoir and pressurized to 10 bar before pumping into the RO membrane unit, whilst the retentate stream was recirculated back to the feed reservoir. The weight of the permeate was continuously monitored with a digital balance (ML3002, Mettler-Toledo Corporation, Switzerland) to observe the flux decline behavior. Water recovery was limited to 75% to avoid the precipitation of soluble salts on the membrane surface. The RO brine was then sent to the MD unit to be desalinated and concentrated.

Microporous polyvinylidene fluoride (PVDF) hollow fiber membrane (provided by Tianjin Polytechnic University, Tianjin, China) was used in the MD unit process. The feed and permeate temperatures were set at 323 K and 298 K, respectively, and the flow rates of the feed and permeate streams were maintained at 0.5 m/s and 0.1 m/s. A bore side feed configuration was used in this study. During the first 24 h of the experiment, both permeate and retentate streams were recirculated back to the feed tank, then the permeate was collected in a separate reservoir where the weight and the conductivity of the permeate were measured by a digital balance (ML3002, Mettler-Toledo Corporation, Switzerland) and a conductivity meter (Thermo’s Orion 145A+, USA), respectively.

2.2.2. Process 2: Coagulation-activated coke adsorption-UF-RO-MD

The set-up for Process 2 was similar to that of Process 1, with the additional activated coke adsorption unit prior to the UF unit process (Fig. 2). In the activated coke adsorption unit, 75 g activated coke was added into a feed tank containing 5 L of coagulant effluent with an Electric mixer (JL-1, Ronghua Equipment Corporation, China). After 2 h of adsorption (preliminary experiments revealed that the adsorption equilibrium was reached within 2 h), the water sample was settled for 12 h and then the supernatant was provided as the feed of the UF unit process.

2.3. Analytical methods

2.3.1. Wastewater quality measurement

China national standard methods were adopted to measure and determine the chemical and physical properties of the coal gasification wastewater. The pH and turbidity were determined using a pH meter (FE20-FiveEasy, Mettler-Toledo Corporation, Switzerland) and a turbidimeter (Hach 2100, Hach Corporation, USA). The total organic carbon (TOC) was monitored using a TOC analyzer (Torch, Tekmar Dohrmann Corporation, USA) and the chemical oxygen demand (COD) was measured by the titrimetric method using dichromate as the oxidant in acidic solution at 150 °C for 2 h (Hach heating system, Hach Corporation, USA). The concentration of NH₃-N was obtained through Nessler’s reaction using a 722S spectrophotometer (Jingke Equipment Corporation, China). The iron content was measured using the inductively coupled plasma atomic emission spectroscopy (ICP-AES) (Prodigy, Leeman Corporation, USA) and the ion chromatograph (ICS–1000, Dionex Corporation, USA). All the chemicals used in the experiments were with analytic grade.

2.3.2. FEEM analysis

The FEEM spectra of the wastewater samples were recorded with a fluorometer (F-7000, Hitachi, Japan). A 450–W Xenon lamp was used as the excitation source. FEEM were collected every 5 nm over an excitation (Ex) range of 200–400 nm, with an emission (Em) range of 280–550 nm by 5 nm. The spectra were obtained by subtracting the Milli-Q water blank spectra, recorded under the same conditions, to eliminate water Raman scatter peaks. The software Origin 8.5 (OriginLab Inc., USA) was used to process the data. The X-axis indicates the emission spectra from 300 nm to 550 nm while the Y-axis represents the excitation from 200 nm to 400 nm, and the contour line as the third dimension, is given to express the fluorescence intensity.

2.3.3. Molecular weight distribution (MWD) analysis

MWD was measured using a laboratory-scale membrane separation apparatus (Minimate TFF system, Pall Corporation, USA). The wastewater was first filtrated through a 0.45 μm cellulose acetate membrane, and then followed by the filtration through a series of ultrafiltration membranes with different molecular weight cut-off (MWCO) from 650 Da to 30,000 Da (Minimate TFF capsule, Pall Corporation, USA). The membranes were washed with deionized water for at least 24 h prior to use.

2.3.4. GC–MS analysis

GC–MS was used for organic compound analysis. The samples were pretreated by liquid–liquid extraction using CH₂Cl₂.
(chromatogram pure grade, Fisher Corporation, USA). Then, 1 μL pretreated sample was analyzed using a 6890N/5973 GC–MS system (Agilent Corporation, USA), coupled with a DB-35MS capillary column with the inner diameter of 0.25 mm and the length of 30 m. The temperature of the gasification compartment was maintained at 280 °C. The temperature was first set at 40 °C for 3 min and then increased to 280 °C with an increment of 3 °C/min. The electron energy and the electron double voltage were set at 70 eV and 1200 V, respectively, and the molecular weight scan varied from 50 Da to 800 Da.

2.3.5. SEM analysis

The surface morphology of the activated coke and the fouling cake layer formed on the membrane surface was studied using scanning electron microscopy (S-3000N, Hitachi Company, Japan).

3. Results and discussion

3.1. Pollutants removal efficiency in each unit of two different processes

Fig. 3 presents the TOC and turbidity results in each unit of the two different processes. In terms of the TOC, it was found that the coagulation unit substantially reduced the TOC level from 152 mg/L to 87.4 mg/L, whilst the subsequent UF process only slightly reduced it to 75.3 mg/L in the case of Process 1 where no adsorption unit was added. On the other hand, in the case of Process 2, a sharp drop in TOC from 87.4 mg/L (after coagulation) to 12.3 mg/L (after adsorption) was observed due to the addition of the activated coke adsorption unit. Similarly, the subsequent UF unit in Process 2 only managed to further reduce TOC from 12.3 mg/L to 11.8 mg/L. Such a high TOC removal efficiency of the activated coke adsorption (up to 85.9%), along with the low TOC removal efficiency of UF process suggested that a large number of organic matters and colloids that cannot be effectively removed by UF membranes due to the relatively smaller particle size than the pore size of UF membrane, can be instead, sufficiently removed by the activated coke adsorption. The similar level of TOC after the RO unit in both processes indicated that the RO process alone is also capable of removing the majority of the organic matters, similar to that of the adsorption unit. However, it should be noted that the organic matters present in the wastewater stream is the main foulant contributing to the undesired fouling behavior as aforementioned. As such, the addition of an activated coke adsorption unit prior to the UF unit not only functioned as an additional step to better remove contaminants, but also served as a safeguard to prevent the subsequent membrane processes (UF, RO, and MD) from undesired fouling. In terms of the effluent turbidity, similar trend in comparison with TOC was observed with the additional adsorption unit, suggesting the excellent adsorption efficiency of the activated coke.

3.2. Mechanism of the organic pollutants removal in the activated coke adsorption unit

The results presented in Fig. 3 clearly indicate that the activated coke can sufficiently adsorb a large amount of organic matters in the wastewater. To better reveal the removal mechanism of the adsorption unit, a variety of characterization techniques were applied, and the results are presented in this section.
3.2.1. **FEEM spectra analysis**

Fig. 4 shows the fluorescence spectra of the raw wastewater, coagulation effluent and adsorption effluent. Region I, II, III, and IV represents aromatic protein, fulvic acid-like organics, humic acid-like organics, and soluble microbial by-product-like materials, respectively [22]. The raw wastewater contained all the above mentioned pollutants. After coagulation, the fluorescence intensities decreased but it was still difficult for the coagulation unit to efficiently remove the pollutants. This was also consistent with some previous studies, which reported that many humic and fulvic acid substances derived from raw wastewater and some soluble microbial product produced during biodegradation could not be effectively removed by coagulation [23–25]. However, significant variations were observed after the AC adsorption unit. Two peaks associated with humic acid and fulvic acid compounds disappeared, indicating that AC effectively absorbed the refractory organics. The adsorption effluent only contained aromatic protein and soluble microbial by-product-like materials with decreased concentration.

3.2.2. **MWD analysis**

The molecular weight distribution of the raw wastewater, coagulation, and adsorption effluents are shown in Fig. 5. It was clear that the majority compounds presented in the raw wastewater had a molecular weight either higher than 30,000 Da or less than 10,000 Da. Furthermore, it appeared that the coagulation unit displayed exceptional removal capability for compounds with molecular weights higher than 30,000 Da, and mediocre efficiency for removing compounds with molecular weights less than 10,000 Da. After adsorption, contaminants with molecular weight more than 3000 Da have been removed effectively. Combined with the FEEM analysis, it can be concluded that the contaminants mainly included refractory humic acid and fulvic acid compounds. These compounds imposed significant fouling propensity on membrane system reported by numerous studies [26–31]. The fouling study was also performed in this work and the results will be presented in the following sections to reveal the effect of the additional adsorption unit on the fouling behavior of the subsequent membrane units.
3.2.3. GC–MS analysis

Tables 2 and 3 show the organic compositions of the coagulation and activated coke adsorption effluent, respectively. MWD results from Fig. 5 illustrated the presence of a large portion of refractory organic compounds with molecular weight less than 30,000 Da in the coagulation effluent. As shown in Table 2, these compounds were mainly ketones, heterocyclic compounds, alcohols, esters and benzenes, which could lead to membrane fouling on the RO and MD units. As a result of the activated coke adsorption, majority of the heterocyclic compounds were removed and only little ketones, alcohols, esters, and benzenes were still detected in the effluent (Table 3). These residual contaminants were mainly from aromatic protein and soluble microbial by-product-like materials with a molecular weight less than 650 Da.

3.3. Effect of adsorption pretreatment on the fouling behavior of the IMS

3.3.1. Effect of adsorption pretreatment on the fouling behavior in the UF unit

Fig. 6 shows the normalized flux ($J$/$J_0$) and trans-membrane pressure (TMP) profiles as a function of time in the UF unit of two different processes. The results clearly indicated that in the case of no additional adsorption unit, the permeate flux sharply dropped by 24.0% and the TMP increased by 66.0% (from 0.50 bar to 0.83 bar), suggesting the occurrence of severe fouling. On the other hand, when the adsorption unit was introduced, the permeate flux remained almost constant and the TMP only increased by 12.0% (from 0.50 bar to 0.56 bar). These results indicated that the activated coke adsorption unit effectively removed contaminants responsible for fouling; consequently, the fouling behavior in the subsequent UF process was significantly mitigated.

3.3.2. Effect of adsorption pretreatment on the fouling behavior in the RO unit

Fig. 7 shows the normalized flux ($J$/$J_0$) as a function of time in the RO units of two different processes. It can be seen that the normalized flux in both cases decreased over time. More specifically, a more pronounced flux decline was observed in the IMS process without the adsorption pretreatment. At the end of operation cycle, the normalized fluxes of Processes 1 and 2 decreased by 28.0% and 11.4%, respectively. The fouling of the RO membrane was caused by the accumulation of organic and inorganic matters on the membrane surface that cannot be effectively removed by the preceding UF process. As such, the result demonstrated in Fig. 7 also indicated that the activated coke adsorption showed great influence not only on the following UF process but also on the RO unit further downstream.

3.3.3. Effect of adsorption pretreatment on the fouling behavior in the MD unit

Fig. 8 shows the normalized flux and the conductivity profile as a function of time in the MD units of two different processes. Similar to the observations in the UF and RO processes, a much more moderate flux decline was observed in the MD process when the activated coke adsorption unit was included, suggesting the suppression of the fouling behavior due to the removal of foulants in the adsorption unit.

In terms of the conductivity profiles in two processes, it was found that the conductivity of the permeate streams increased sharply in the initial stage of the MD process, suggesting that some metal ions have penetrated into the distillate. Furthermore, the maximal conductivity for Processes 1 and 2 were found to be 140.1 mS/cm and 122.3 mS/cm, respectively. Afterward, a gradual decrease in conductivity was observed for both processes. Through
Table 2
GC–MS analysis of the coagulation effluent.

| Number | Organic compounds in coagulation effluent | Formula | Relative content (%) |
|--------|------------------------------------------|---------|----------------------|
| 1      | 2-Methyl-1,3-cyclopentanedione             | C₆H₁₂O₂ | 2.1                  |
| 2      | 5-Methyl-isothiazole                      | C₅H₁₀N  | 18.8                 |
| 3      | Dibutyl phthalate                         | C₈H₁₀O₄ | 10.3                 |
| 4      | 2,2,6,6-Tetramethyl-4-piperidone          | C₈H₁₂NO | 9.9                  |
| 5      | 1,2-Benzenedicarboxylic acid bis(2-methylpropyl) ester | C₅H₁₀O₄ | 6.1                  |
| 6      | 4-Butyl-benzenamine                       | C₆H₁₃N  | 4.8                  |
| 7      | Benzothiazole                             | C₅H₁₀N  | 4.2                  |
| 8      | 2,3-Dimethyl-3-buten-2-ol                 | C₈H₁₆O  | 4                    |
| 9      | 2-Methylpiperidine                        | C₅H₁₁N  | 3.7                  |
| 10     | 4-Hydroxy-3-methoxy-benzonitrile          | C₈H₁₃NO₂| 3.2                  |
| 11     | 2-Aminopyridine                           | C₆H₁₂O  | 2.7                  |
| 12     | 5,6,7,8-Tetrahydro-2-naphthalenol         | C₁₀H₈N  | 2.1                  |
| 13     | Alpha,alpha-dimethyl-benzemethanol        | C₁₆H₂₂O₄| 13.8                 |
| 14     | 5-Acetyl-2-methylpyridine                 | C₈H₁₈O  | 17                   |
| 15     | 1,2-Benzenedicarboxylic acid disoctyl ester | C₁₆H₂₂O₄| 1.7                  |
| 16     | Tetramethyl-thiourea                      | C₅H₁₂N₂S| 1.3                  |
| 17     | Butylated hydroxytoluene                  | C₁₀H₁₄O | 1                    |

Total TOC (mg/L) 74.8

Table 3
GC–MS analysis of the AC adsorption effluent.

| Number | Organic compounds in AC adsorption effluent | Formula | Relative content (%) |
|--------|-------------------------------------------|---------|----------------------|
| 1      | 1,2-Benzenedicarboxylic acid disoctyl ester | C₁₆H₂₄O₄| 50.8                 |
| 2      | Butylated hydroxytoluene                   | C₁₀H₁₄O | 14.5                 |
| 3      | Dibutyl phthalate                          | C₁₀H₁₆O₄| 13.8                 |
| 4      | 2,2,6,6-Tetramethyl-4-piperidinone         | C₈H₁₂NO | 8.7                  |
| 5      | 1,2-Benzenedicarboxylic acid, bis(2-methylpropyl) ester | C₁₆H₂₂O₄| 7                    |
| 6      | Alpha,alpha-dimethyl-benzemethanol         | C₁₀H₁₆O | 4.2                  |
| 7      | Tetramethyl-thiourea                       | C₅H₁₂N₂S| 1                    |

Total TOC (mg/L) 12.3

Fig. 8. Normalized flux (J/J₀) and conductivity as a function of time in the MD units of two different processes (a) Process 1 and (b) Process 2.

the analysis of MD distillate, only Ca²⁺, SO₄²⁻, NO₃⁻, and Cl⁻ were presented and the total concentration of these inorganic ions were 10.6 mg/L for Process 1 and 4.38 mg/L for Process 2, which shall only lead to a very low conductivity. Therefore the increase of the distillate conductivity in the initial stage of the both processes (as shown in Fig. 8) could be a result of penetration of the volatile organic compounds from the feed to the distillate [32]. Once the diffusion of volatile organic compounds was complete, the conductivity of the distillate stream began to decrease over the remaining period of operation time. The slightly lower maximal conductivity and final conductivity at the end of the MD operation in the case of Process 2 suggested that a portion of the volatile organic compounds were removed in the activated coke adsorption unit.

3.3.4. Surface morphology of the activated coke and membranes
Fig. 9(a and b) shows the SEM images of the activated coke before and after adsorption. The surface of raw activated coke appeared to be uneven and porous, indicating large available surface area for adsorption. After the adsorption, much smoother surface was observed, suggesting that most of the surface of the activated coke was occupied by the adsorbed organic and inorganic compounds from wastewater.

Fig. 9(c and d) and (e and f) shows the surface morphology of the RO and MD membranes after the experiments in two processes. For both the RO and MD membrane in Process 1 where no adsorption unit was presented, a thick cake layer was formed on the membrane surface, suggesting severe fouling which was confirmed by the larger flux decline in these two cases. On the other hand, when the adsorption pretreatment was applied, the cake layers appeared to be much thinner for both RO and MD membranes. Furthermore, the membrane fouling behavior was significantly alleviated, which was in good agreement with the results of flux decline in RO and MD units (the permeate fluxes of the RO and MD units increased by...
21.9% and 23.1%, respectively, when the adsorption unit was added in comparison with those of Process 1).

4. Conclusions

A novel integrated membrane system incorporated with an activated coke adsorption unit was proposed and examined with real coal gasification wastewater provided by a local coal gasification plant in China as influent. The effect of the additional adsorption unit on the contaminant removal and membrane fouling mitigation was systematically studied. The use of abundant and low-cost activated coke (provided by the same coal gasification plant) as adsorbent effectively removed majority of the organic matters presented in the wastewater which otherwise can be hardly removed by the conventional membrane systems. The FEEM, MWD, and GC–MS analysis further revealed that humic acid, fulvic acid, and heterocyclic compounds were mostly removed by the additional adsorption unit, most of which are commonly found as main contributors to the undesired membrane fouling phenomenon. As a result of such an efficient removal of these potential foulants, the integrated membrane system incorporated with adsorption unit displayed much better fouling resistance for
all the membrane units integrated in the process (UF, RO, and MD).

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