Removal of Ions From Produced Water Using Powder River Basin Coal

Zaixing Huang  
University of Wyoming

Fang-Jing Liu  
University of Wyoming

Mingcheng Tang  
University of Wyoming

Yangyan Gao  
University of Wyoming

David Bagley  
University of Wyoming

Xin He  
University of Wyoming

Alexander Goroncy  
University of Wyoming

Maohong Fan  
mfan@uwyo.edu  
https://orcid.org/0000-0003-1334-7292

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Abstract

Although becoming less attractive as an energy source, coal has significant potential for other, more sustainable uses including water treatment. In this study, we present a simple approach to treat water that was produced during oil production and contained a total dissolved solids (TDS) content of over 150 g/L using Powder River Basin (PRB) coal. PRB coal used as packing material in a flow-through column effectively removed 60-80% of the cations and anions simultaneously. Additionally, 71-92% of the total organic carbon in the produced water was removed as was all of the total suspended solids. The removal mechanisms of both cations and anions were investigated. Cations were removed by ion exchange with protons from oxygen-containing functional groups such as carboxylic and phenolic hydroxyl groups. Anions, mainly Cl⁻, appeared to be removed through either the formation of resonance structures as a result of delocalization of electrons within coal molecules or through ion-π interactions. We propose that coal is a “pseudo-amphoteric” exchange material that can remove cations and anions simultaneously by exchanging ions with both ionized and non-ionized acids that are ubiquitous in coal structure or resonance effect.

Introduction

Global population growth and climate change have made sustainable water supply a challenge to all human beings. Hoekstra estimated that 2 billion people are living in areas with severe water scarcity (Hoekstra 2014). The same author updated the number to 4 billion in a later model (Mekonnen and Hoekstra 2016). Yet “produced water” is produced in large quantities concurrently with oil production, gas production, and other unconventional oil and gas production including coalbed methane and shale oil and gas extraction (Stoll et al. 2015; Veil et al. 2004; Vengosh et al. 2014). The water-to-oil ratio for oil production is about 3:1 (SPE). The oil and gas industries produce about 14.5 billion m³/yr (250 million barrels/day) of produced water, of which 40% is discharged into the environment (Fakhru’l-Razi et al. 2009). Produced water accounts for more than 80% of the liquid waste (Igunnu and Chen 2012) produced during oil and gas production.

Although a significant potential source of water, produced water contains organic and inorganic compounds. Specifically, dissolved and dispersed oils (BTEX, PAHs, and phenols), grease, heavy metals, radionuclides, treating chemicals, formation solids, salts, dissolved gases, scale products, waxes, microorganisms, and dissolved oxygen are commonly found in produced water. Furthermore, the treatment and disposal of produced water are costly. The expense for treating this potential resource is estimated at $40 billion per year (SPE) while the disposal cost on a unit basis could be as high as $4/barrel, depending on the quality of the produced water and the methods that are used (Duraisamy et al. 2013).

Nevertheless, due to the need for water and the desire to limit environmental damage due to discharge, treatment of produced water for potable and irrigation uses has become an option (Qi et al. 2021). Conventionally, produced water is disposed by direct discharge or reinjection into disposal wells
(Duraisamy et al. 2013; Jiménez et al. 2018), thus providing no beneficial use of the water. Physical treatment processes include adsorption, sand filters, hydrocyclones, evaporation, dissolved air precipitation, C-TOUR, freeze-thaw evaporation, devaporation, electrodialysis/electrodialysis reversal, gas flotation, and macro-porous polymer extraction. Chemical treatment processes include chemical precipitation, chemical oxidation, electrochemical processes, photocatalytic treatment, in situ chemical oxidation (ISCO), room temperature ionic liquid, and demulsifier (Dickhout et al. 2017; Duraisamy et al. 2013; Fakhru'l-Razi et al. 2009; Iggunu and Chen 2012; Li et al. 2010; Munirasu et al. 2016). Membrane filtration, using microfiltration, ultrafiltration, reverse osmosis, nanofiltration, bentonite clay, and zeolite membrane technologies (Dickhout et al. 2017; Munirasu et al. 2016; Nasiri et al. 2017), has been extensively examined with emerging produced water treatment technologies including membrane distillation (Woo et al. 2017), forward osmosis (Bell et al. 2017), and the two combined (Kim et al. 2017). However, membrane fouling and high capital and operations costs remain a big hurdle for these membrane filtration technologies (Duraisamy et al. 2013; Stoll et al. 2015). Hackney and Wiesner estimated the cost for treating average quality produced water to remove most of solids, organic, and inorganic components with unit processes to be $8.06/m³ adjusted to the inflation rate of 2019 (Hackney and Wiesner 1996) and increasing up to $35.00/m³ for produced water having high concentrations of organic constituents, total suspended solids (TSS), and total dissolved solids (TDS), with TDS removal dominating the cost.

Another fossil fuel, coal, may have potential for cost-effectively treating produced water from other fossil fuel production processes. Coal accounts for over 88% of the world’s fossil fuels (Wang et al. 2019), is widely available and found in about 100 countries all over the world (Andruleit et al. 2016) and is predicted to outlast other hydrocarbon resources by hundreds of years. Furthermore, the conventional uses of coal for power generation, steel making, and chemical feedstock production (Falbe et al. 1982) are becoming less attractive because these uses produce pollutants and emit more CO₂ than any other energy sources (Huang et al. 2017). Coal is now being examined as an inexpensive raw material for manufacturing activated carbon (Rosa et al. 2017), carbon fiber (Li et al. 2017; Liu et al. 2018), silicon metal (WCA), and sorbent (virus removal from water) (Oza and Chaudhuri 1975), for example. Other studies have examined using coal fly ash for synthesizing zeolites (Otal et al. 2005; Querol et al. 2002) or ceramic microfiltration membrane (Jedidi et al. 2009) to be used for produced water treatment. Song et al. reported fabrication of a microfiltration carbon membrane with anthracite to remove oil from oily wastewater (Song et al. 2006). The preparation required carbonization of the coal at an elevated temperature for up to 900 °C. Sulfonated or ammoniated coal were prepared with chemical reactions to add additional functional groups to enhance the ion exchange capacity of coal (Nachod 2012).

These uses require processing of coal in some manner prior to use. Yet coal is a complex material that may have inherent properties suitable for treating produced water directly, without extensive processing of the coal. The objectives of this work were (a) to test the hypothesis that native subbituminous coal from Wyoming’s Powder River Basin can effectively treat produced water by removing extremely high concentrations of TDS as well as organic carbon, and suspended solids simultaneously, and (b) to
evaluate the ion exchange and sorption mechanisms by which cations and anions are removed. To the best of our knowledge, this is the first work to investigate the treatment of produced water with native coal directly for simultaneous removal of suspended solids, organic, and inorganic components.

Materials And Methods

2.1 Coal and Produced Water

Powder River Basin coal from Wyodak was provided by Black Hills Corporation. Coal was milled by Wyoming Analytical Lab (Laramie, WY, USA). The fraction of coal with particle sizes between 40–60 mesh was used for column preparation. The raw coal contained 16.00% moisture, 8.18% ash on a moisture-free basis, and 48.73% volatile matter on a dry and ash-free basis. The elemental composition of the coal was 78.87% C, 3.72% H, 1.01% N, and 15.93% O on a dry and ash-free basis (Liu et al. 2018). The pore properties of the coal were BET specific surface area (N2 at 77 K) 2.598 m2/g, DFT pore diameter 0.844 nm, average pore diameter 13.64 nm, and total pore volume 0.018 cm3/g (Xu et al. 2020; Zhang et al. 2021).

The petroleum produced water was provided by Hunt Oil Co. (Knox City, TX, USA). It was collected on February 22, 2018 from the feed before entering the wastewater treatment plant. Key parameters of the produced water included: pH 6.83, TSS 233 mg/L, TDS 152,320 mg/L, conductivity 238 mS/cm, total organic carbon (TOC) 42 mg/L, and alkalinity 80 mg/L.

2.2 Experimental Design and Operation

All experiments were conducted by passing produced water through a column packed with coal. A 500-mm long, 25-mm ID Kontes chromatography column equipped with a PTFE stopcock plug (Kimble Chase, Rockwood, TN, USA) was packed with ground coal (30 g). A pinch of cotton was used as a strainer right before the stopcock plug to strain the coal particles. The coal was washed with 100-mL of deionized (DI) water that was passed through the packed coal by gravity. The final height of packed coal was approximately 65 mm.

Filtration experiments were conducted to examine the removal of suspended solids, organic, and inorganic compounds from produced water. An aliquot of 30 mL produced water was filtered (by gravity unless otherwise stated) through the column packed with coal. The filtrate was collected and designated as the sample to be analyzed for that cycle. Then the column was washed with 30 mL of 1% HCl (wt/vol, 37% TraceMetal Grade, Fisher Scientific, Pittsburgh, PA, USA), followed by a DI water rinse until the pH of the wash was circumneutral. After washing, another 30-mL aliquot of produced water was filtered through the same coal column. This filtration-washing cycle was repeated 10 times.

To investigate the impact of functional groups on the water treatment, the coal was extracted with solvents including NMP (N-Methyl-2-pyrrolidone), (THF Tetrahydrofuran), and methanol before being placed in the column. The ground coal was soaked in solvent at a ratio of 1 to 3 (coal wt/solvent wt) at ambient temperature for one week. The coal-solvent mixture was then filtered through a filter paper.
(Whatman qualitative, Grade 1, GE, Pittsburg, PA) and the coal was washed with that solvent until the filtrate contained no color. The coal was recovered and dried at 80 °C. The dried, extracted coal was then used in the filtration apparatus as described for the sequential filtration experiments.

### 2.3 Analytical Measurements

Conductivity, TDS and pH were measured by an H280G pH, conductivity and DO meter (Hach, Longmont, CO, USA). The alkalinity measurement was done with an alkalinity test kit AL-AP MG-L (Hach, Longmont, CO, USA). TOC was measured for both raw and treated produced water. TOC was analyzed with a Shimadzu TOC analyzer (TOC V CSN, Shimadzu Corporation, Kyoto, Japan). The original water sample was also analyzed with a Varian Cary Eclipse Fluorescence Spectrophotometer (Agilent, Walnut Creek, California, USA) for 3D-EEM under emission 3D mode. The scanning setup is as follows: emission wavelengths (Em) 290–590 nm with 2 nm increments; excitation wavelengths (Ex) 225–450 nm with 2.5 nm increments, slit bandwidths 5 nm for both emission and excitation; and scan rate 1200 nm/min (Wang et al. 2019). The spectra were processed using MatLab (Natick, Massachusetts, USA).

The concentrations of major inorganic elements, including Ca, Cl, Fe, K, Mg, and Na of the raw and treated produced water were determined by a Perkin Elmer Nexlon 300 ICP/MS (Waltham, MA, USA) under standard mode (Huang et al. 2020; Lewińska-Preis et al. 2021). The certified ICP/MS standards were purchase from Inorganic Ventures (Christiansburg, VA, USA). A Re solution (50 ppb) was used as the internal standard. The water samples were diluted to meet the requirements of the device prior to analysis.

The untreated and solvent-extracted coal were characterized by a Fourier transform infrared (FTIR) spectrometer (Nicolet iS50, Thermo Scientific, USA). The attenuated total reflectance (ATR) technique was used to acquire data without complicated sample preparations. The samples were scanned between 4000 and 400 cm$^{-1}$ at a resolution of 4 cm$^{-1}$. The data of each spectrum were collected with 128 scans. Coal samples after filtration and after acid wash were studied using a Scanning Electron Microscopy with Energy Dispersive X-Ray Spectroscopy (SEM-EDS, Model #51-XMX0005, FEI, Quanta FEG MK2; Oxford Instruments America, USA) to examine the adsorption of the inorganic ions. The coal samples were dried at 80 °C, spread evenly on a double-sided carbon tape which was then mounted on an SEM stub. The scanning was operated in HV mode with an accelerating voltage of 20 kV.

### Results And Discussion

#### 3.1 Removal of Inorganic Ions, TOC and TSS

The primary inorganic components of the untreated produced water include cations: Na (49.2 g/L), Ca (10.0 g/L), Mg (1.70 g/L), K (0.31 g/L), and Fe (0.56 g/L), and an anion: Cl (90.8 g/L). Na and Cl comprised about 92% by weight of the total inorganic ions determined by ICP-MS. These inorganic ions were effectively removed by filtration through the coal column (Fig. 1, **Supplementary Table 1**). Ten filtration-regeneration cycles were conducted and showed 60–80% removal of individual ions with an
average removal of 73%. Interestingly, the percent overall removal of all ions was similar, ranging between 64% and 76%, suggesting there was no specific removal selectivity of these ions. An average of about 0.12 g of TDS was removed per g coal for each filtration-regeneration cycle, although there is residual water in the pores of the packed column after washing. There may be dilution effects from the residual water that need further investigation.

The TOC concentration of the original produced water was 42 mg/L. During filtration through the coal column, 71–92% of the TOC was removed. The removal efficiency of the initial cycle (Cycle-1) was the lowest (71%), whereas the mean value of the remaining 9 cycles was 90%. The removal capacity of organic components was significantly improved after coal regenerations. The TSS was decreased from 233 mg/L in the raw produced water to a negligible level (below detection) in the filtrate.

Figure 2 shows SEM and EDS images of the coal samples directly after filtration and also after regeneration with the acid wash. The EDS analysis (Fig. 2b) clearly shows adsorption/deposition of major inorganic ions on the surface of coal particles. After regeneration with the acid wash, most of the cations were removed (Fig. 2d), thus confirming how the treatment cycles worked. Ions were retained in the coal through filtration and subsequently removed by acid wash, thereby regenerating the coal medium for further filtration cycles. After regeneration with the acid wash, Cl dominated the surface of coal (Fig. 2d) because diluted HCl solution was used as the washing agent.

### 3.2 Results of $^{13}$C NMR analysis

The raw coal, washed coal before filtration (Before filtration), and coal after filtration (After Filtration) were subjected to NMR analysis to identify functional groups and investigate the changes of these functional groups. As shown in Fig. 3, many oxygen-containing functional groups, including alcohol, carboxyl, carbonyl, phenol, ester, and ether were identified in all samples with different intensity (Kim et al. 2013). Other non-oxygen-containing functional groups including aliphatic and aromatic C-H groups such as methyl, methylene, and methyne were also identified. This is in line with other studies suggesting low rank coal contains hydroxyl-, methoxy-, and/or methyl-substituted benzene rings, carboxylic acids, and aliphatic linkers line –CH₂CH₂− group (Liu et al. 2013; Liu et al. 2019).

In general, the intensity of the three samples decreased in the order Before filtration > raw coal > After filtration. Before filtration coal was raw coal that had been washed in preparation for use in the column. The washing process removed ions within the coal. In contrast, the After filtration sample contained ions that were removed from the produced water and were bound/adsorbed onto the coal. Because the electron distribution of $^{13}$C can be affected by factors such as binding partners, the binding of ions can significantly reduce the response in NMR analysis. This may explain the intensity differences among coal samples of before filtration, raw and after filtration. In particular, the differences in intensity of the phenolic and carboxylic functional groups were prominent, suggesting their involvement in the removal of ions from the produced water.

### 3.3 Effects of Solvent Extraction on Coal and Solvents
To investigate the mechanisms of ion removal by coal filtration, PRB coal was solvent-extracted with tetrahydrofuran (THF), methanol, or N-methyl-2-pyrrolidone (NMP). The total ion removal capacity of the coal (as determined from a filtration cycle without subsequent acid washing) was significantly decreased by solvent extraction, to 26%, 23%, and 16% removal for THF, methanol, and NMP extractions, respectively.

Solid coal, both before and after extraction, and liquid solvent after extraction were examined by FTIR analysis. Although there are difficulties in using FTIR to analyze heterogeneous materials like coal with respect to sample preparation, band assignments, and baseline correction (Solomon and Carangelo 1982), a number of functional groups and changes due to solvent extraction were identified in this study. Figure 4 shows the FTIR spectra of coal and liquid from solvent extraction. The peaks were identified according to the literature (Sigma-Aldrich; Xie 2015). The coal structure was modified by solvent extraction to different extents (Fig. 4a). Specifically, coal extracted with NMP exhibited the greatest structural changes with respect to functional groups, followed by methanol and THF. This is consistent with the ion removal capacity where NMP extraction showed the greatest decrease in capacity, followed by methanol and then THF which showed the smallest decrease in ion removal capacity.

The liquid solvents after extraction contained compounds with oxygen- and nitrogen-containing functional groups (Fig. 4b), consistent with the changes in the solid coal functional groups. The intensity of N-H stretching for solvent-extracted coal was significantly reduced, whereas peaks of N-H stretching existed in all the liquid extracts. The intensity of the N-H peaks of the coal followed the same tendency of ion removal capacity, with less ion removal capacity corresponding to decreased intensity of the N-H peaks. This was also true for other functional groups such as carboxyl and phenolic-hydroxyl groups, suggesting that these functional groups may be involved in ion removal. These results are most evident in the NMP treatment where the intensity of these peaks in the extracted coal were significantly reduced while the NMP after extraction had a strong presence of these peaks. NMP has been shown to facilitate the extraction of hydroxyl-containing moieties from bituminous coal (Sun et al. 2014). These results indicate that the extraction of these functional groups significantly impaired the ion removal capacity of the coal, largely related the removal capacity to the number of these oxygen- and nitrogen-containing moieties.

3.4 Composition of coal extracts

Volatile and small-molecular compounds in the extracts were characterized by GC/MS to investigate the impact of extractable components on the ion removal. As shown in Fig. 5, the group components detected in the extracts mainly included alkanes, alkenes, arenes, alcohols, phenols, ketones, carboxylic acids, and esters. The relative abundances of oxygen-containing compounds (i.e., alcohols, phenols, ketones, carboxylic acids, and esters) compared to all extracted compounds were 47.2%, 86.4%, and 82.0% for THF, methanol, and NMP extracts, respectively. Oxygen-containing functional groups, especially carboxyl and hydroxyl, in these compounds may interact with metal cations to form cation-bridging linkages (Liu et al. 2016; Mathews and Chaffee 2012), which are beneficial for ion removal. Extraction of
these oxygen-containing compounds appears to have significantly reduced the ion removal capacity of the coal.

The detected compounds in the methanol and NMP extracts were dominated by esters with 76.6% and 52.9% relative abundance, respectively (Fig. 5), while the hydroxyl-containing compounds (like carboxylic acids, alcohols, and phenols) had a relatively low abundance. As polar solvents, methanol and NMP have proven to be effective for extracting hydroxyl-containing compounds from coals (Liu et al. 2016; Sun et al. 2014). Because GC/MS is only sensitive for volatile and less polar compounds, the GC/MS detectable compounds account for only a fraction of the compounds in the extracts. According to FTIR analysis, NMP extract may also contain many hydroxyl-containing compounds, especially carboxylic acids with low carbon numbers, which were not detected by GC/MS.

4. Proposed Mechanisms Of Ion Removal

The structure of a coal macromolecule may be visualized as a condensed aromatic carbon-atom lattice surrounded by a typical “fringe” formed by functional side groups. The left panel of Fig. 6 is a hypothetical model of coal structure (Malumbazo 2011). It is a heterogeneous mixture composed of a macromolecule network with varying degrees of cross-linking (Smith et al. 2013). Modified lignin, as well as cellulose and melanoidin-type materials, are considered to be the ‘backbone’ of this macromolecule network. The cross-linkage of lower rank coal, including subbituminous coal, is dominated by alkyl and aryl ether groups with oxygen functional groups. The chemical heterogeneity of coal decreases from low rank coal to high rank while the aromaticity increases, suggesting that lower rank coals (lignite and subbituminous) have more complex chemical structures than high-rank coals (bituminous coal and anthracite) because the low rank coal contains several distinct classes of constituents (Hofrichter and Fakoussa 2001; Wang et al. 2015). Carboxyl and hydroxyl groups, among others, are the main oxygen-containing functional groups in coal structure which are present in low-rank coal (Xie 2015). Phenolic hydroxyl groups are the main form, but some alcoholic hydroxyl groups also exist.

The removal of cations from produced water by coal filtration is proposed to be through ion exchange (Fig. 6a). The carboxyl and hydroxyl groups may act as ion exchanger sites by exchanging protons (H⁺) for other cations (Na⁺, K⁺, Ca²⁺, Mg²⁺, etc.) allowing these cations to bind to the negatively charged hydroxyl and carboxyl functional groups. This hypothesis is supported by the pH measurements of the raw produced water before filtration (6.83) and the treated water (as low as 2.28), indicating that protons were released by the coal, presumably due to cation exchange. Additionally, NMR analysis of before and after filtration coal samples also showed significant intensity reduction of carboxyl and hydroxyl groups, further supporting the cation exchange hypothesis. The solvent extraction results provide additional support for the hypothesis. Solvent extraction removed carboxyl and hydroxyl functional groups, as illustrated by the FTIR analyses where the functional groups were removed from the coal and present in the extracts, resulting in significantly impaired ion removal capacity.
Proposed mechanisms for Cl\(^-\) removal are less clear. One possible removal mechanism is adsorption of Cl\(^-\) due to the delocalization of electrons within molecules to form resonance structures (Fig. 6b). This resonance effect or mesomeric effect or electron-donating effect occurs between a lone pair of electrons and a pi bond or two pi bonds next to each other (Dewick 2006). For example, the lone electron pair in the oxygen of the phenolic hydroxyl group may be donated to form a double bond and leaving a positive charge. The donation stabilizes the structure of the non-ionized acid such as phenols or derivatives. The donating effect passes the electrons on to the pi bonds along the ring to produce a negative charge of the para- or ortho-carbon in the same aryl ring. These charged molecules could then bond with both negative and positive ions in the produced water to remove them. The resonance effect could be positive or negative. In the positive resonance effect, -OH, -SH, -OR, and -SR can increase the electron density of the stabilizing ring while -NO\(_2\), -S = O, and -C = O could decrease the electron density of the stabilizing ring in the negative resonance effect (Dewick 2006). The removal of these functional groups, as occurred with solvent extraction, would reduce the ion removal capacity of coal.

Alternatively, the ion removal of any particular ion may not be attributed to any single functional group. Other possible mechanisms to produce charges in coal include changes of electron density by binding alkali metal ions (which are dominant species in produced water) with the functional groups in aromatic structures and electrostatic induction (Xie 2015). In recent decades, ion-π interactions have been recognized and found to be widely exist as a form of general noncovalent bonding (Dougherty 1996; Ma and Dougherty 1997; Schottel et al. 2008). Ion-π interactions happen not only in aromatic systems, but are also well documented in other simple π systems such as ethylene and acetylene. Studies show that highly solvated cations can be sequestered by such binding force in aromatic-containing structures (Ma and Dougherty 1997), while anion-π interactions happen in electron deficient aromatic systems (Schottel et al. 2008). The anion-π interaction combines effects of electrostatic and anion-induced polarization, with the former correlated to permanent quadruple moment, Q\(_{zz}\) and the latter to molecular polarizability (Quiñonero et al. 2004; Schottel et al. 2008). Aromatic molecules with lower absolute values of Q\(_{zz}\) could bind to both anion and cation which might account for some of the ion removal in our system (Schottel et al. 2008). The ion-π interaction binding energy is estimated to be 20–50 kJ/mol which is energetically favorable and is comparable to the binding energy of hydrogen bonds. In the ion-π theory, the inductive effect, rather than the resonance effect, facilitates binding (Ma and Dougherty 1997; Schottel et al. 2008). Although the mechanisms of anion removal remain hypothetical, based on the results presented herein, coal may be considered a “pseudo-amphoteric” exchanger that has the capability of removing both cations and anions simultaneously and effectively.

**Conclusions**

In this study, we have demonstrated that TOC, TSS, and especially TDS can be removed simultaneously from produced water using Power River Basin subbituminous coal as the treatment material. The material is stable with repetitive filtration-regeneration cycles. More importantly, the coal was shown to have a surprisingly high ion removal capability for produced water containing extremely high dissolved
solids (> 150,000 mg/L or 15%). We propose that the PRB coal is a “pseudo-amphoteric” ion exchange material that could simultaneously remove both positive and negative ions. The analyses strongly support the hypothesis of ion exchange as the mechanism for cation removal whereas the mechanism for anion removal is proposed to be by adsorption through resonance structures induced by delocalization of electrons within coal molecules or through ion-π interactions with aromatic clusters. Further work is needed to confirm the anion removal mechanisms. These results may have important ramifications in the reclamation of produced water from energy production sector.

Declarations

Availability of data and materials

The data supporting the findings of this study are available within the article and its Supplementary Information files.

The authors declare no competing interests

Authors’ contributions

Zaixing Huang: Conceptualization, data curation, formal analysis, investigation, methodology, visualization; Writing-original draft/reviewing & editing

Fangjing Liu, Mingchen Tang, Yangyan Gao, Xin He, Alexander Goroncy: Data curation, formal analysis, reviewing & editing

David Bagley: Formal analysis, supervision, validation; Writing-original draft/reviewing & editing

Maohong Fan: Conceptualization, supervision, resources; Writing-original draft/reviewing & editing

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Figures
Figure 1

Inorganic ions removed by filtering 30-mL of produced water through 30-g of coal. Cycles 2-10 were conducted with washed (regenerated) coal as described in the text. The raw produced water had TDS of 152 g/L.
**Figure 2**

SEM & EDS images of the coal samples. (a) SEM of coal directly after filtration; (b) EDS of coal directly after filtration; (c) SEM of coal after regeneration with acid wash; (d) EDS of coal after regeneration with acid wash.
Figure 3

13C NMR analysis of raw coal, washed coal before filtration of produced water and coal after filtration of produced water.
Figure 4

FTIR spectra of solvent-extracted coal: (a) Solid Coal (Original was coal prior to solvent extraction, NMP, Methanol and THF were extracted coal); (b) Liquid solvents after extraction.
Figure 5

GC/MS detectable constituents (by group component) in the coal extracts
Figure 6

Proposed mechanism of inorganic ions removal. (a) Cation exchange; (b) Resonance effect. The cations (monovalent and divalent) were removed by ion exchange with H-bearing moieties (-OH, COOH, etc.). Both cations and anions are proposed to be removed by non-ionized charges induced by resonance effect.

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