




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The characteristics of mineralogy, morphology and sintering during co-combustion of Zhundong coal and oil shale

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The high-sodium coal from Zhundong region induces severe ash-related problems during combustion. Oil shale rich in silicon and aluminum has been adopted as an additive of Zhundong coal during co-combustion to investigate the characteristics of mineralogy, morphology and sintering at various temperatures and blending ratios in this study. The size distributions of the ash depositions have also been determined. The study results reveal that the fluxing minerals, such as sodium-based compounds, with sizes less than 10 μm are dominant in Zhundong coal. These fine particles may present in a molten or sintering state at high temperature. Some refractory minerals with size exceeding 10 μm , such as nepheline or anorthite, form during co-combustion. These irregular particles retain their shapes during combustion resulting from their high melting temperatures. The blending of oil shale can change the migration path of the sodium during combustion, indicating that sodium gradually migrates from the water-soluble phase to the aluminosilicate phase. The amount of ash particles less than 10 μm decreases from 84.31% to 40.29% with an increase in temperature. Furthermore, the refractory minerals with high melting point and high viscosity can result in high sintering temperature. The sintering temperature increases from 700 $^{\circ}\text{C}$ to 1055 $^{\circ}\text{C}$ when the blending ratio of oil shale reaches 20% and combustion temperature reaches 1050 $^{\circ}\text{C}$. In addition, 30–50% sodium-based compounds volatilize to the gaseous phase during combustion of Zhundong coal. When the proportion of oil shale reaches 10%, and combustion temperature is 850 $^{\circ}\text{C}$, sodium volatility will decrease to the lowest point of 14.49%, which can suppress ash sintering to a large extent.

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1. Introduction

The Zhundong region is situated in the east of the Junggar Basin, Xinjiang, China, where the coal resource is abundant, with prognostic reserves of 390 billion tons. This can meet the demand of China's coal consumption for 100 years based on the current annual average consumption.^{1,2} Zhundong coal ranges from lignite to sub-bituminous, which is obviously characterized by a low content of ash and sulfur, moderate calorific value and thus high quality.^{3–6} It has been widely considered as one of the most important coal resources in China, which possesses a wide research and application potential. However, Zhundong coal applied as a fuel material has been characterized by a high AAEMs level which is attributed to the local alkaline geographical environment caused by a marine sedimentary coalfield. Especially, the content of sodium oxide exceeds 2% in the ash, the highest even exceeds 10%, which is much higher than other steam coals.⁷ It is well known that the mineral constituents of coal exert a significant effect on thermal transformation efficiency of coal combustion

and safety of boilers operation. In some power plants, massive combustion of high sodium Zhundong coal has led to ash-related problems in the furnace wall and heat exchange surface of coal-fired boilers, such as fouling, slagging and sintering, which influence safe operation and service life of boilers, even results in boiler collapse.⁸ The volatilization of sodium-based compounds in Zhundong coal could be one of the major factors during combustion.⁹ Therefore, it is of theoretical value and practical significance to study the slagging and sintering mechanism and control technology of sodium element by means of investigating the characteristics of mineralogy, morphology and sintering of the ash in order to have Zhundong coal been applied widely.

Yang *et al.* adopted improved sequential extraction method to study the content of sodium in different modes. The result indicated that the sodium in Zhundong coal was mainly dominated by water-soluble sodium, which approximately accounted for 50–85% of the total sodium.¹⁰ The volatility of water soluble sodium at high temperature was considered to be noxious during Zhundong coal combustion in boilers of power plants. Meanwhile, Zhou *et al.* evaluated the mineral phases of Zhundong coal ash under oxy-fuel combustion, and revealed that the main mineral phases are NaCl, quartz, calcite, anhydrite, and andradite, which was consistent with the research

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result by Shen.^{11,12} Among these minerals, NaCl was considered as water-soluble sodium which played a dominated role in slagging and sintering performance. Xu *et al.* studied ash formation during combustion of Zhundong coal in a drop tube furnace and proposed that the ash of Zhundong coal had a large amount of particles with sizes less than 10 μm which typically rich in Ca, Fe, and Mg.² Li *et al.* investigated the formation of fine particulates from Zhundong coal during combustion and revealed that fine particles (<10 μm) rich in Na, S, and Cl had agglomerated and stuck to coarse ash particles (>10 μm), indicating sintering had occurred.¹³ In addition, sintering temperature has been widely considered as a criterion for the evaluation of coal ash slagging tendency.^{6,14} Llorente concluded that kaolin, lime, calcined dolomite, and ophite could reduce the sintering behaviour, whereas silica did not efficiently reduce the sintering tendency of biomass ash.¹⁵

A few strategies to alleviate ash-related problem caused by sodium during Zhundong combustion have been proposed in the previous research, such as, pretreatment,^{16–18} addition of less-fouling coal,^{19–23} different additives blending,^{24–26} as well as modifying boiler structure and optimizing operating conditions,²⁷ which have some limitations and shortcomings. Based on the current researches, co-combustion of silicon–aluminum-rich additives could inhibit sodium volatility, which is one of the effective modes to solve the ash-related problems.²⁸ When Zhundong coal was ashed with silicon-rich Heilongjiang coal, Wang *et al.* drew a conclusion that sodium volatility could be non-obvious if $\text{SiO}_2/\text{Na}_2\text{O}$ or $(\text{SiO}_2 + \text{Al}_2\text{O}_3)/\text{Na}_2\text{O}$ in the blending coal sample were more than 20 or 25, respectively.²⁹ Wei *et al.* blended Zhundong coal with SiO_2 , kaolin and fly ash respectively, compared the effects of three additives on capturing sodium and it turned out that the effect of SiO_2 was best at a most appropriate temperature of 1000 $^\circ\text{C}$.³⁰ Xu *et al.* investigated the efficiency of capturing sodium during co-combustion Zhundong coal with silicon–aluminum-rich kaolin, and the result showed that the more the kaolin content was, the better the capturing efficiency was, conversely, the higher the temperature was and the bigger the particle size was, then the lower the capturing efficiency is. Furthermore, 70% sodium could be captured with kaolin content ratio of 6%, particle size of 11 μm and combustion temperature of 1200 $^\circ\text{C}$.³¹

The oil shale, a new type fuel with rich reserve in Xinjiang Province, has been adopted as an additive of Zhundong coal during co-combustion, which is flammable sedimentary rock rich in silicon and aluminum elements.³² Those two elements are beneficial components for coal ash, which have an inhibitory effect on sodium volatility of Zhundong coal and change the path of sodium transformation during co-combustion, furthermore can enhance the sintering temperature. To date, the co-combustion of Zhundong coal and oil shale is rarely reported, some relevant mechanisms should be investigated for deeper. Thus, in this paper, the mineralogy and morphology of the blended fuel during co-combustion were studied through XRD and SEM-EDS. In addition, the variation of particle size and sintering temperature of the blended fuel would also be investigated. These above could provide valuable reference for clean and efficient utilization of Zhundong coal.

2. Materials and methods

2.1. Coal and additive samples

A “run of mine” Zhundong coal (ZD) and Changji oil shale (CJ) were used in this current study, which were crushed, ground and screened to 75–100 μm . The proximate analysis, ultimate analysis and ash component analysis of Zhundong coal and oil shale were performed in Table 1. It could be obviously seen that Zhundong coal has higher moisture and lower ash content than oil shale. In addition, it has been widely considered that the base to acid ratio, denoted as B/A, $B/A = (\text{Na}_2\text{O} + \text{MgO} + \text{CaO} + \text{K}_2\text{O} + \text{Fe}_2\text{O}_3)/(\text{SiO}_2 + \text{Al}_2\text{O}_3 + \text{TiO}_2)$, is also an important index in slagging and sintering propensities according to previous researches. At the same time, Zhundong coal has a higher content of alkali metal and alkali earth metals (AAEMs), and lower content of SiO_2 and Al_2O_3 than oil shale, which results in higher B/A and slagging tendency. In contrast, oil shale has a higher content of SiO_2 and Al_2O_3 , and lower content of AAEMs, which leads to lower B/A and slagging propensity.

In addition, coal blends were prepared by physically mixing ZD and CJ samples with five blending ratios in the weight percentage: 100 : 0, 95 : 5, 90 : 10, 85 : 15, 80 : 20, which were termed as ZD, MF1, MF2, MF3, and MF4 for convenience. These samples were dried in an air blowing thermostatic drying oven at 105 $^\circ\text{C}$ for 24 h and stored in sealed bags for latter use.

2.2. Coal ashing

A programmable high-temperature tube furnace SRJK-2.5-13 was applied in the ashing experiment, with maximum temperature of 1350 $^\circ\text{C}$, furnace pot size of 2- ϕ 22 \times 180 mm, voltage of

Table 1 Properties of the Zhundong coal and Changji oil shale

	ZD	CJ
Proximate analysis (wt% as air dried base)		
M	12.93	2.92
A	5.48	78.72
V	25.27	13.18
FC	56.32	5.18
Ultimate analysis (wt% as air dried base)		
C	65.29	9.06
H	2.96	1.84
O	15.10	6.73
N	0.54	0.48
S	0.16	0.25
Ash composition (wt%)		
SiO_2	18.26	63.50
Al_2O_3	8.98	22.24
CaO	27.30	0.58
Fe_2O_3	7.28	5.24
K_2O	0.52	2.56
MgO	9.88	1.70
Na_2O	4.58	0.33
SO_3	20.34	0.08
P_2O_5	0.38	0.09
TiO_2	0.34	0.78
B/A ratio	2.54	0.12



0–220 V, power of 2.5 kW. In the experiment, ZD and CJ were put into the tube furnace for ashing, final combustion temperature was set to 550 °C for low temperature ashing. Because some minerals such as AAEMs would escape to gaseous phase at high temperature during coal combustion, which could cause relatively large errors in the analysis of mineral composition. At the same time, MF1, MF2, MF3, MF4 were conducted separately with final temperature of 850 °C, 950 °C, 1050 °C, respectively. All ash samples were heated in air atmosphere with the flow rate of 0.4 L min⁻¹ and remained constant for 2 hours under final temperature to make sure the mixed fuel sample were ashed completely. Furthermore, the heating rate was controlled to 10 °C min⁻¹ below 600 °C, and 5 °C min⁻¹ above 600 °C.

2.3. Chilling and drying

The ash samples obtained from co-combustion were rapidly immersed in liquid nitrogen for chilling which can be cooled down to less than 300 °C within 2 s in order to prevent a crystal phase change during the temperature drop. After these samples were cooled down to room temperature, they were dried in a thermostatic drying oven at 105 °C for 2 h. The ash samples after chilling and drying were subjected to subsequent determination and characterization.

2.4. XRD, SEM-EDS and particle size analysis

The Rigaku MiniFlex600 X-ray diffractometer (XRD) would be adopted to identify the mineral phase in the ash samples, with radiation source of Cu K α , tube voltage of 40 kV, tube current of 15 mA, scanning rate of 8° min⁻¹, scanning range of 20–60°, and also software jade 6.5 could be applied for comparison of absorption peak and ICDD Power Diffraction File (PDF) database to confirm the mineral composition of ash samples with different target temperature and blending ratio. In addition, different characteristic absorption peaks represent different mineral compositions of coal ashing, and the strength of characteristic absorption peak also represents the content of mineral composition.³³

The texture, morphological characteristic and spot elemental composition of the ashes particles in the deposit were observed using a Tescan Vaga 3 scanning electron microscopy (SEM) coupled with an Oxford X-ray energy dispersive spectroscopy (EDS). Each sample was scattered on a carbon tape carefully so that the particles could be observed individually.

The particle size distribution of each ash sample was also determined using a CILAS-1190 laser particle size analyzer in wet mode with a measurement range of 0.04–2500 μ m. Each measurement was repeated three times to ensure good repeatability of the results.

2.5. Sintering temperature determination

A designed pressure-drop device, as introduced elsewhere,¹⁴ was used to determine the sintering temperature of the ash samples prepared in the preceding section. Briefly, 0.4–0.8 g of ash was compacted into a pellet under a pressure of 0.8 MPa which was then heated in a tube furnace at a heating rate of 6.7 °C min⁻¹ in inert atmosphere (N₂) at a flow rate of 10 mL min⁻¹. The gauge

pressure prior to the pellet and the temperature of the pellet were continuously recorded. The sintering temperature of the ash samples was measured based on the first-order and second-order derivatives of the pressure-drop curves as a function of temperature in accordance with the research paper, which was an improved and accurate method compared to other sintering temperature measurement.

2.6. Sodium content determination

The sodium content of the ash sample was determined using the ICAP 6000 Series inductively coupled plasma-optical emission spectrometer (ICP-OES) produced by Thermo Fisher Scientific. Each ash sample would be digested with a hydrofluoric acid–perchloric acid mixture to obtain the sample solution before the determination of the absorbency by ICP-OES, at a radio frequency power of 1.15 kW, plasma gas flow rate of 10.0 L min⁻¹, auxiliary gas flow rate of 0.75 L min⁻¹, atomization gas flow rate of 0.50 L min⁻¹, peristaltic pump flow rate of 1.5 mL min⁻¹, and integration time of 5 s (average of three tests), with an axial torch and a radial viewing mode.

2.7. Volatilization proportion of sodium

To better present the volatilization characteristic of sodium in the ash sample after co-combustion, the following parameters were introduced for calculation.

Sodium content, m_1 (mg g⁻¹), of mixed fuel after ashing (coal-based)

$$m_1 = m'_1 \times A \quad (1)$$

where m'_1 = sodium content of mixed fuel ash after ashing (ash-based), mg g⁻¹, and A = yield of ash produced from mixed fuel ashing.

Percentage of sodium residue in mixed fuel ash, α (%)

$$\alpha = \frac{m_1}{m_0} \times 100\% = \frac{m'_1 \times A}{m_0} \times 100\% \quad (2)$$

where m_0 = sodium content of mixed fuel before ashing (coal-based), mg g⁻¹.

Volatilization proportion of sodium, β (%), at combustion temperature T , which was calculated as follows:

$$\beta = 1 - \alpha = 1 - \frac{m_{\text{Na,MF}}(\omega, T)}{m_{\text{Na,ZD}} + m_{\text{Na,CJ}}} \times 100\% \quad (3)$$

where $m_{\text{Na,MF}}(\omega, T)$ = content of sodium in the ash of mixed fuel with an oil shale blending ratio of ω and final combustion temperature of T ; $m_{\text{Na,ZD}}$ = content of sodium in Zhundong coal; and $m_{\text{Na,CJ}}$ = content of sodium in Changji region oil shale.

3. Results and discussion

3.1. Ash mineralogy

Fig. 1 shows the XRD pattern of the ash samples of ZD, CJ and mixed fuel heated at different temperatures and blending ratios in air atmosphere. In Fig. 1(b)–(d), only the 20–60° XRD patterns



are shown here, because the ash samples of mixed fuel had weak absorption peaks at 10–20° and 60–90°. It can be evidently observed that the combustion temperature plays a significant role in the ash mineralogy transformation process, by contrast, the oil shale ratio has little influence.

Fig. 1(a) shows that the primary mineral phases in the ZD ash deposit at combustion temperature of 550 °C are mainly identified as anhydrite (CaSO₄), quartz (SiO₂), thenardite (Na₂SO₄), halite (NaCl). This coal also contains calcite (CaCO₃), larnite (Ca₂SiO₄), and periclase (MgO) in minor quantities. Among these minerals, the fluxing minerals including calcite, anhydrite, thenardite and halite with high viscosity and low melting point are dominant while the refractory minerals such as quartz, larnite and periclase are deficient, indicating severe sintering tendency and high deposition propensity.^{34,35} At the same time, the primary minerals in CJ ash deposit are identified as SiO₂ and Al₂O₃ in major quantities, which are consistent with the ash composition analysis results in Table 1. These minerals are refractory phases which suggest high melting point and low deposition propensity.

Fig. 1(b) shows that the mineral compositions in the ash deposition of mixed fuels with different blending ratio of oil shale at combustion temperature of 850 °C, are mainly quartz (SiO₂), anhydrite (CaSO₄), albite (NaAlSi₃O₈), anorthite

(CaAl₂Si₂O₈) and kyanite (Al₂O₃·SiO₂). It can be seen that all the absorption peaks of halite, thenardite, larnite and calcite disappears which will result from a series of physical and chemical changes. The entire amount of the halite and thenardite in ZD volatilizes into the gas phase which will adhere to the heat exchange surface and boiler wall. Meanwhile, the calcite disappears which suggests that it decomposes into CaO and CO₂. Furthermore, larnite decomposes into CaO and SiO₂. The generated CaO reacts with SiO₂ and Al₂O₃ in oil shale to form anorthite with a melting point of 1553 °C. Also, NaCl in ZD reacts with SiO₂ and Al₂O₃ in oil shale to form albite with a melting point of 1180 °C. Albite readily reacts with Fe₂O₃, CaO and some aluminosilicates to form a eutectic with a low ash fusion point of the mixed fuel. In addition, SiO₂ and Al₂O₃ in the mixed fuel react to form kyanite which will easily transform into mullite at high temperatures.

Fig. 1(c) shows that the mineral compositions in the ash deposition of mixed fuels with different blending ratio of oil shale at combustion temperature of 950 °C, are mainly quartz, anhydrite, anorthite, kyanite, gehlenite (Ca₂Al₂SiO₇) and akermanite (Ca₂MgSi₂O₇). It can be seen that the kyanite in the ash of the mixed fuel at the temperature of 950 °C is more than that at 850 °C, indicating that more kyanite is further generated with the increase in combustion temperature. Meanwhile, the

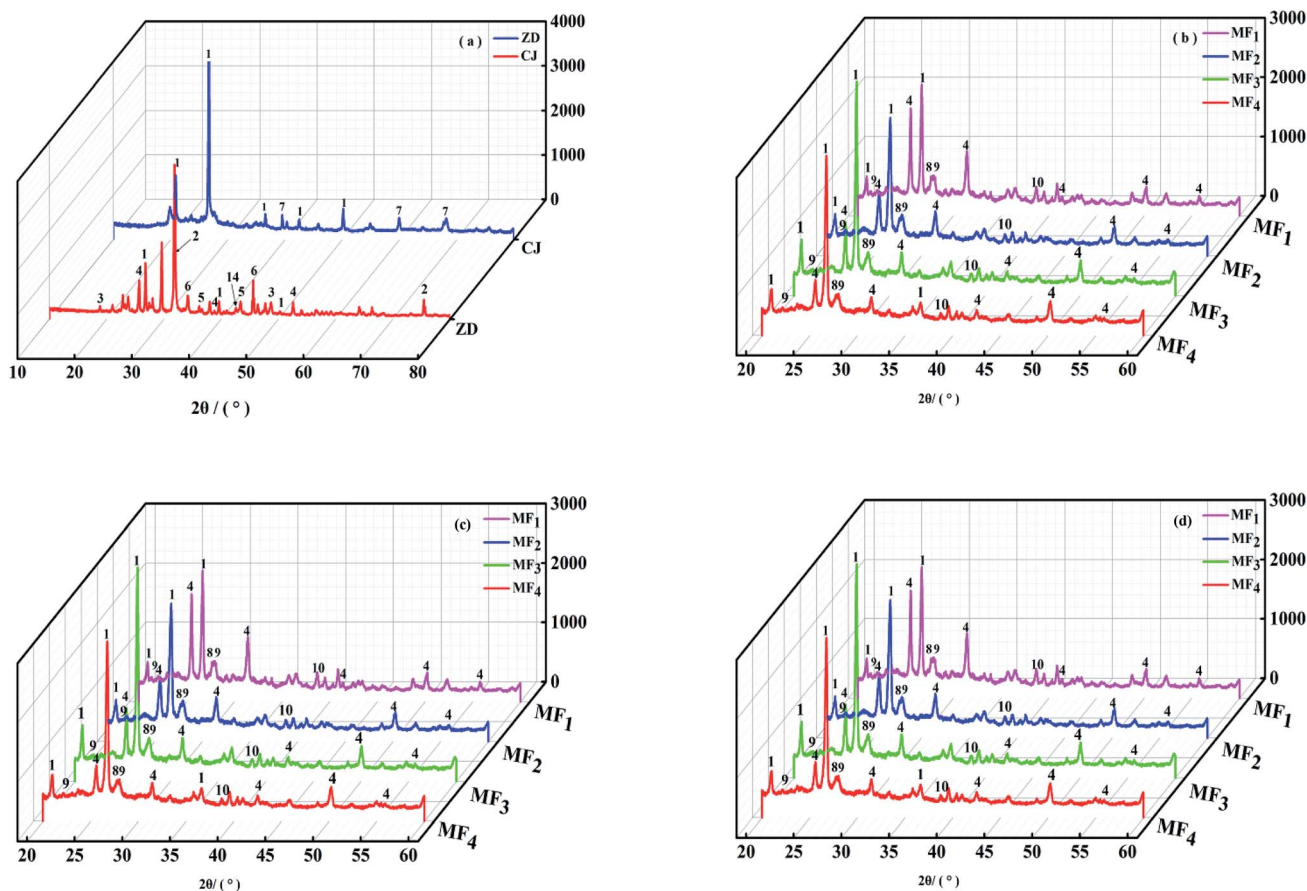


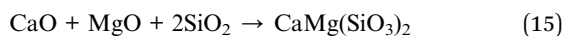
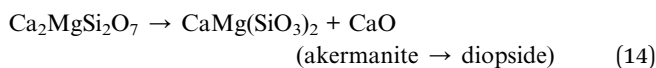
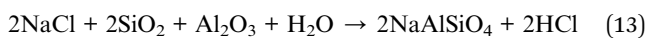
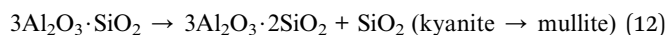
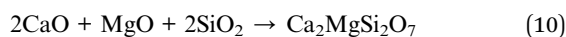
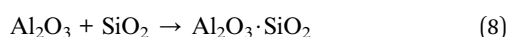
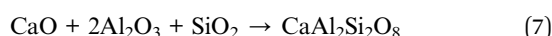
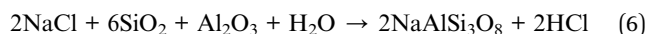
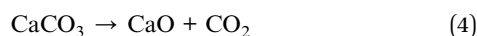
Fig. 1 The XRD pattern of ash samples of ZD, CJ and mixed fuel obtained at different blending ratios and temperatures. 1. Quartz; 2. halite; 3. thenardite; 4. anhydrite; 5. calcite; 6. larnite; 7. Al₂O₃; 8. anorthite; 9. albite; 10. kyanite; 11. anorthite; 12. nepheline; 13. mullite; 14. periclase; 15. gehlenite; 16. akermanite; 17. diopside.



anorthite in the ash of the mixed fuel at temperature of 950 °C is less than that at 850 °C, resulting from that the anorthite reacts with CaO to form gehlenite with a melting point of 1593 °C. In addition, the CaO generated from former reaction reacts with periclase in ZD coal and quartz in oil shale to form akermanite with a melting point of 1450 °C.

Fig. 1(d) shows that the mineral compositions in the ash deposition of mixed fuels with different blending ratio of oil shale at combustion temperature of 1050 °C, are mainly quartz, gehlenite, anhydrite, akermanite, mullite (3Al₂O₃·2SiO₂), nepheline (NaAlSiO₄) and diopside (CaMg(SiO₃)₂). It can be seen that all the absorption peaks of the anorthite in the ash of the mixed fuel disappears, owing to that all of the anorthite transforms into gehlenite with the temperature increase. Also, the kyanite transforms into mullite with a melting point of 1850 °C, which is consistent with previous research results.³⁶ In addition, albite disappears and decomposes into nepheline with a melting point of 1550 °C and quartz. Nepheline can also be produced by the reaction of sodium-containing compounds in ZD coal with SiO₂ and Al₂O₃ in oil shale directly at this temperature. Akermanite decomposes into diopside with a melting point of 1391 °C and CaO. And diopside can also be generated by the reaction of CaO, MgO, and SiO₂ in the ash of the mixed fuel directly at this temperature.

These reactions above are given as follows:



It can be distinctly known that all the ash deposition of mixed fuel heated at various temperatures with various blending ratios of oil shale contain a certain amount of quartz and anhydrite. This is because oil shale contains a large amount of SiO₂ and the following reactions also produce it. Meanwhile, anhydrite is hard to volatilize which will decompose above 1700 °C in an oxidizing atmosphere. After blending oil shale in Zhundong coal, a series of physical and chemical changes have

occurred in the ash of the mixed fuel during co-combustion, which can change the migration path of the sodium. The XRD patterns show that sodium gradually migrated from the water-soluble phase to the aluminosilicate phase, which has been deposited in the ash and inhabited to volatilize by the Si and Al in oil shale.

3.2. Ash morphology and chemistry

Fig. 2 presents typical ash depositions SEM images of ZD coal heated at 550 °C and mixed fuel containing 90% ZD coal and 10% oil shale heated at 850 °C, 950 °C, 1050 °C and EDS results at selected spots of interest in order to investigate the morphology and chemistry. In addition, the particle size distributions of ash depositions are shown in Fig. 3.

As shown in Fig. 2(a), at combustion temperature of 550 °C, it is evidently observed that an irregular ash particle in area 1 mostly consist of Si and O, indicating the presence of quartz. Meanwhile, the spherical ash particle in area 2 has experienced fully molten at high temperature whose elemental compositions are mainly Na, S and O, suggesting that sodium sulfate exists. It is because of the low melting temperature of sodium sulfate that the spherical ash particle in area 2 is in the fusional state. In addition, the bonding ash particles aggregate in area 3 which is mainly composed of Na and Cl, showing the present of halite. These fine particles have stuck with each other, and the particle sintering occur which is observed in the SEM image resulting from the low sintering temperature of halite. An irregular ash particle has been found in area 4 rich in Ca, S and O, which shows that calcium sulfate exists in ZD coal indubitably. In the ash deposition of ZD coal, discrete fine particles with sizes less than 10 μm play a dominate role which account for 84.31% of total ash particles according to the ash particle size distribution of ZD coal as shown in Fig. 3. This is in agreement with the literature research studied by Li. The fine particles rich in AAEMs (especially Na and Ca) are fluxing minerals with size less than 10 μm, which have low sintering temperature and high viscosity. This can easily stimulate slagging tendency and enhance sintering propensity.

As shown in Fig. 2(b), at combustion temperature of 850 °C, an oblate spherical ash particle in area 1 is observed whose elemental compositions are mainly Si and O, suggesting the presence of quartz. Meanwhile, the coarse irregular ash particle exists in area 2 which is mainly composed of Na, Si and Al according to the EDS result, indicating the presence of sodium aluminosilicate which is identified as albite further referring to the weight fraction of elements and result of XRD above. Some discrete fine particles less than 10 μm aggregate on its surface so that particles sintering occur under this circumstance. In addition, the elemental compositions of the irregular ash particle mainly consist of Ca, S and O (area 3) and Ca, Si, Al and O (area 4), indicating the existence of calcium sulfate and aluminosilicate. These irregular ash particles retain their shapes during combustion which resulted from their high melting temperatures.

As shown in Fig. 2(c), at combustion temperature of 950 °C, it is evidently observed that the ash deposition consists of both



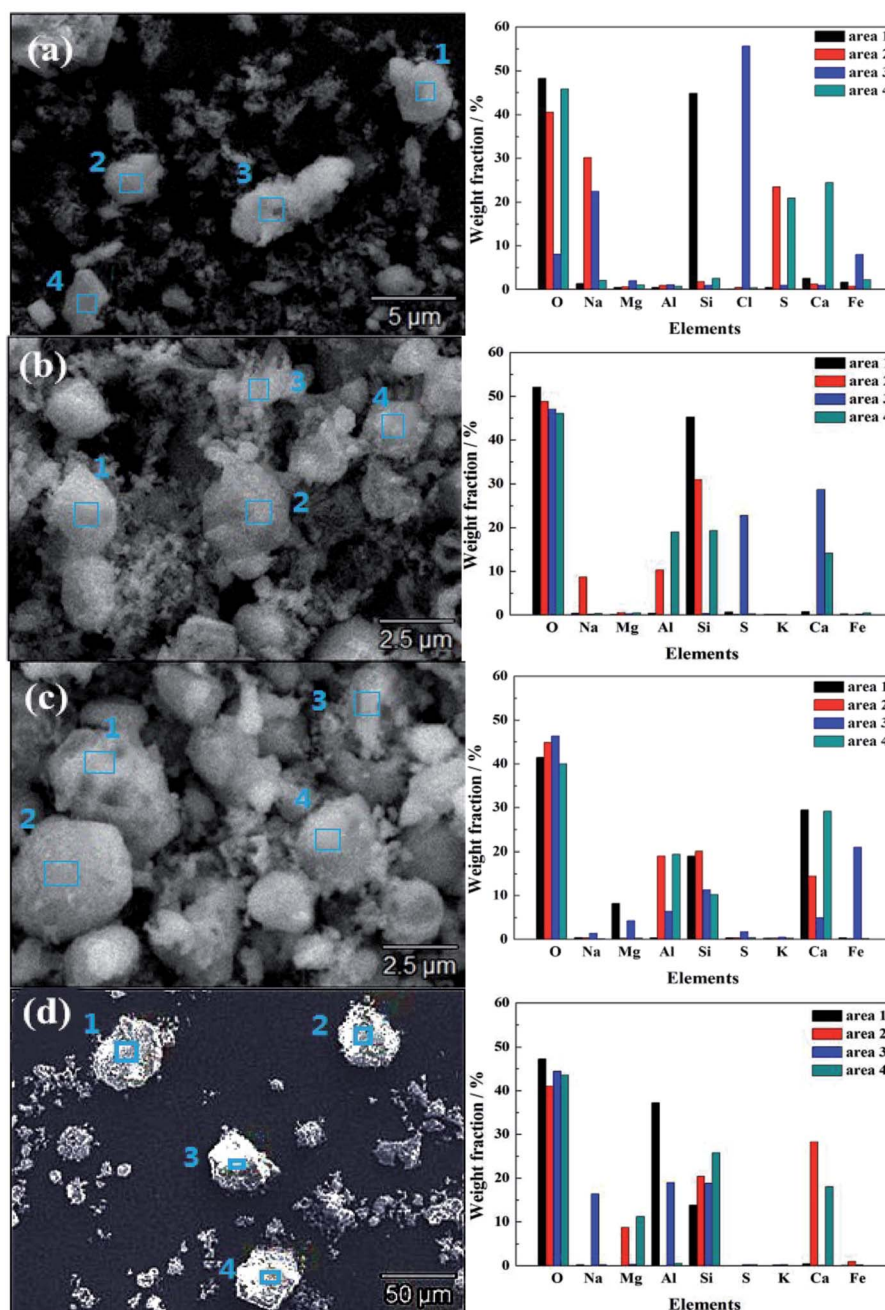


Fig. 2 Typical ash SEM images of ZD coal at temperature of (a) 550 °C and mixed fuel at temperatures of (b) 850 °C, (c) 950 °C, (d) 1050 °C, and EDS results of elemental analysis at selected spots.

fine ash particles and large coarse ash particles. Large irregular particles rich in silicate or aluminosilicate with high melting point are dominated in the ash deposition. The large irregular ash particle in area 1 whose elemental compositions mainly consist of Ca, Mg, Si and O, confirming the present of akermanite with size more than 10 μm. Meanwhile, the larger irregular spherical ash particle in area 2 is observed which is mostly composed of Ca, Si, Al and O, indicating the of calcium aluminosilicate which is identified as anorthite further referring to the weight fraction of elements and result of XRD above. In addition, a molten agglomeration form in area 3, which

contains majority of Fe, Ca, Mg, Si and O, resulting from that the oxides of iron, calcium, magnesium and silicate lead to form a low-temperature eutectic. The irregular spherical ash particle in area 4, is identical in the elemental composition with that in area 2, but different in weight fraction of each element, which is determined as gehlenite further according to the weight fraction of elements and result of XRD above. Furthermore, some discrete fine particles have adhered to gehlenite, resulting in a decrease in the amount of discrete fine ash particles, as shown in Fig. 3.



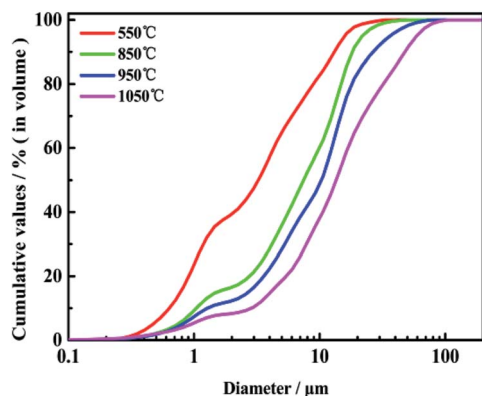


Fig. 3 Size distributions of ZD coal at temperature of 550 °C and mixed fuel at temperatures of 850 °C, 950 °C, 1050 °C.

As shown in Fig. 2(d), at combustion temperature of 1050 °C, it has obviously been found that an irregular coarse ash particle in area 1 whose elemental compositions are mainly Si, Al and O, which is probably large-size mullite with some fine ash particle stuck on it. Meanwhile, an oblate spherical ash particle mainly consisted of Ca, Si, Al and O has been observed in area 2, which is considered as calcium aluminosilicate with large size and high melting point. The large angular ash particle in area 3 is mostly composed of Na, Si, Al and O which is identified as sodium aluminosilicate, confirming the formation of nepheline according to the result of XRD above. Furthermore, another large angular ash particle has been found in area 4, with large amount of Mg, Ca, Si and O, which is further considered as diopside. In addition, it has been shown that the dispersion of ash particle considerably increases and the amount of large ash particle with size more than 10 μm also increase, which accounts for 60% approximately, resulting in the mitigation of sintering.

It has been firmly considered that the combustion temperature has a significant influence on the morphology and size distribution of ash deposition, resulting from the action of combustion on the mineral reaction. With an increase in combustion temperature, the amount of fine ash particle decreases and large ash particle increases. The amount of ash particle less than 10 μm decreases from 84.31% at temperature of 550 °C to 40.29% at temperature of 1050 °C according to the curve of ash particle size distribution, as shown in Fig. 3. Some large ash particles have been found in SEM image resulting from the formation of some minerals rich in silicate and aluminum, such as nepheline, anorthite, mullite, akermanite *etc.*, which are refractory minerals with low viscosity and high melting point, leading to mitigate slagging tendency and suppress sintering propensity.

3.3. Sintering temperature of the ash

The sintering of coal ash refers to the welding and bonding of adjacent powder particles under the effect of excessive surface free energy, which is one of the most important factors of bed material agglomeration, pipe fouling, heat-exchange surface

slagging.^{37–39} Therefore, the sintering temperatures of ZD coal and mixed fuel ash are measured by a designed pressure-drop device to evaluate the effect of combustion temperature and oil shale blending. It has been firmly convinced that the higher sintering temperature, the lower sintering propensity and slagging tendency.

Fig. 4 shows that the sintering temperature of the ash samples of the mixed fuel heated at various temperatures. It can be clearly known that the sintering temperature increases first slightly and then decreases rapidly in general with an increase in the blending ratio of oil shale, regardless of any combustion temperatures. The sintering temperature of coal ash is related to mineral components in the ash, the reactions among them and the formation of a low-temperature eutectic.⁴⁰ The more content of basic oxides such as Na₂O, CaO, MgO indicates low sintering temperature and high slagging tendency in combustion condition. On the contrary, the more content of acidic oxides such as SiO₂ and Al₂O₃ indicates high sintering temperature and low slagging tendency in combustion condition.^{41,42}

The Zhundong coal ash with low sintering temperature of 700–730 °C which is due to the high alkali and alkaline earth metals content which are fluxing minerals according to the results of ash component analysis and XRD. The acidic cations with high ionic potential (Na⁺, Ca²⁺, Mg²⁺, Fe³⁺) are easily combined with oxygen to form complex ions or polymers, while the alkaline cations (Si⁴⁺, Al³⁺) are oxygen donors, which can terminate the agglomeration of the polymers and reduce their viscosity, showing the fluxing effect. In addition, these crystalline fluxing minerals will act as the amorphous minerals under the action of the surface tension showing the flowing state and stimulating ash sintering. When the oil shale blending ratio is in the range of 0–5%, the sintering temperature decreases gradually. It is for the reason that the proportion of mineral component in the ash deposition is in the low-temperature eutectic region of the ternary phase diagram which is consistent with the research results of Li.^{13,43} When the oil shale blending ratio increases further, the sintering temperature increases rapidly, which is because the ash depositions of the mixed fuel contain higher contents of refractory minerals but lower contents of fluxing minerals.

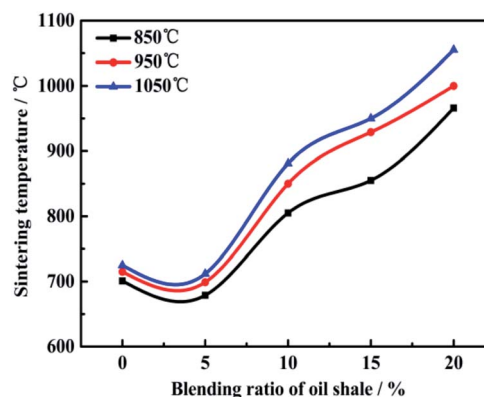


Fig. 4 Sintering temperature of ZD and mixed fuel ash heated at various temperatures.



It can be obviously observed that the sintering temperatures of the ash depositions increase with an increase in temperature, which is attributed to the differences in mineral components of the ash depositions. According to the results of XRD above, more kinds and contents of refractory minerals with high melting point and high viscosity generate, which can result in high sintering temperature. As a result, the sintering temperature increases from 700 °C to 1055 °C, which can suppress ash sintering to a large extent.

3.4. Sodium volatility of the ash

Fig. 5 shows the variation trend of sodium volatilization rates at different blending ratios (0–20%) and temperatures (850–1050 °C). It has been founded that the oil shale blending ratio significantly influenced the sodium volatilization and such effect varied with the combustion temperature.

As shown in Fig. 5, during the combustion of “run of mine” Zhundong coal, 30–50% sodium-based compounds volatilize to gaseous phase, and condense on the heat exchange surface and furnace wall, then react with SO_2 , Fe_2O_3 in the flue gas to generate various kinds of sulfates with low-melting-point and high-viscosity,⁴⁴ such as sodium sulfate (Na_2SO_4), compound sodium sulfate ($\text{Na}_3\text{Fe}(\text{SO}_4)_3$) and sodium pyrosulfate ($\text{Na}_2\text{S}_2\text{O}_7$), which will form a compacted and cohesive sediment layer and result in high-temperature slagging, as shown in Fig. 6.⁴⁵ On the other hand, the fused boding layer would capture fly ash particles in the flue gas. These particles with low-sintering-temperature become fused and softened together through welding and bonding of the ambient particles, which led to high-temperature sintering. Therefore, it is widely assumed that the slagging caused by high-sodium coal results from the joint effects of high-temperature cohesive ash deposition and high-temperature sintering.

When blending Zhundong coal with a certain proportion of oil shale, the volatility of sodium-based compounds obviously reduces. It commendably shows that oil shale could inhibit sodium volatility with a significant effect during co-combustion. The sodium in Zhundong coal is captured by silicon and aluminum in oil shale, and then react to form the high-melting-

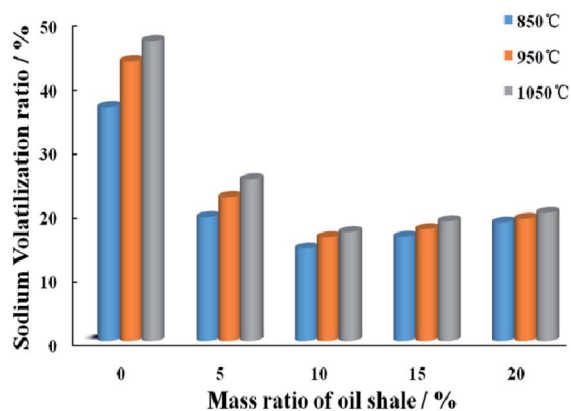


Fig. 5 Variation trend of sodium volatilization rates at different blending ratios and temperatures.



Fig. 6 Situation of slagging and deposition on the heat-transfer surface during Zhundong coal combustion.

point aluminosilicates, which can be exhausted to outside of the boiler by slag discharge system. In addition, with the proportion of oil shale increases, volatility rate of sodium-based compounds rapidly decreases first and then increases slowly under different temperatures. When the proportion reaches 5–10%, oil shale reactivity is strong, and reacts rapidly with sodium in the ash, then volatility rate of sodium reduces obviously. When the proportion reaches 10%, combustion temperature is 850 °C, sodium volatility will decrease to its lowest point of 14.49%. When the proportion further increases, the total content of sodium in the blending coal sample decreases, then the volatility rate will slightly increase.

At the same time, temperature has a profound effect on the release and transformation of sodium during Zhundong coal combustion.⁴⁶ As the temperature increases from 850 °C to 1050 °C, volatility of sodium in different blending coals all increase slightly, which partly results from the formation of mullite. This reaction will lead to provide less SiO_2 and Al_2O_3 for sodium to form sodium aluminosilicates, which will reduce the chance of the capture sodium reaction. Furthermore, with an increase in temperature, the increase of release rate of sodium is always absolutely synchronous with that of oil shale's reaction of capturing sodium element. Since the release rate is higher than capturing rate, the volatility rate of sodium increased slightly. In addition, as the temperature increases, the separation of moisture and volatile in coal become faster, and water-soluble sodium arrives at the surface of coal ash particle with moisture through interspace, thus new channels from in the interior structure of coal, which further accelerates sodium volatility.

4. Conclusions

(1) The combustion temperature plays a significant role in the ash mineralogy transformation process, conversely, the oil shale ratio has little influence. The primary minerals in the Zhundong coal are mainly identified as anhydrite, quartz, the nardite and halite. After blending oil shale, some



aluminosilicates formed, such as albite and nepheline, with an increase in temperature. The blending of oil shale can change the migration path of the sodium during combustion, indicating that sodium gradually migrated from the water-soluble phase to the aluminosilicate phase.

(2) The fluxing minerals, such as sodium-based and calcium-based compounds, with size less than 10 μm are dominated in Zhundong coal. These fine particles may present in molten or sintering state at high temperature. After blending of oil shale, some refractory minerals with size exceeds 10 μm , such as nepheline, anorthite, form during co-combustion. These irregular particles retain their shapes during combustion resulting from their high melting temperatures. The amount of ash particle less than 10 μm decreases from 84.31% to 40.29% with an increase in temperature.

(3) With an increase in the blending ratio of oil shale, the sintering temperature increases first and then decreases. When the blending ratio is in 0–5%, the sintering temperature decreases, because the proportion of mineral component is in the low-temperature eutectic region. When the blending ratio increase further, more kinds and contents of refractory minerals with high melting point and high viscosity generate, which can result in high sintering temperature. The sintering temperature increases from 700 $^{\circ}\text{C}$ to 1055 $^{\circ}\text{C}$, which can suppress ash sintering to a large extent.

(4) During combustion of Zhundong coal, 30–50% sodium-based compounds volatilize to gaseous phase. After blending Zhundong coal with some oil shale, with the proportion increases, volatility rate of sodium-based compounds decreases first and then increases. The effect of oil shale on inhibiting sodium volatility at low temperature is better than at high temperature. When the proportion reaches 10%, combustion temperature is 850 $^{\circ}\text{C}$, sodium volatility will decrease to its lowest point of 14.49%.

Conflicts of interest

There are no conflicts to declare.

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References

- J. Li, X. Zhuang, X. Querol, O. Font, N. Moreno and J. Zhou, *Fuel*, 2012, **95**, 446–456.
- J. Xu, D. Yu, B. Fan, X. Zeng, W. Lv and J. Chen, *Energy Fuels*, 2014, **28**, 678–684.
- J. H. Lv, X. Y. Wei, Y. H. Wang, T. M. Wang, J. Liu, D. D. Zhang and Z. M. Zong, *RSC Adv.*, 2016, **6**, 11952–11958.
- J. Li, M. Zhu, Z. Zhang, Z. Kai, G. Shen and D. Zhang, *Fuel Process. Technol.*, 2016, **144**, 155–163.
- J. Zhou, X. Zhuang, A. Alastuey, X. Querol and J. Li, *Int. J. Coal Geol.*, 2010, **82**, 51–67.
- J. Li, M. Zhu, Z. Zhang and D. Zhang, *Fuel*, 2016, **179**, 71–78.
- X. Wang, Z. Xu, B. Wei, L. Zhang, H. Tan, T. Yang, H. Mikulčić and N. Duić, *Appl. Therm. Eng.*, 2015, **80**, 150–159.
- C. Du, L. Liu and P. Qiu, *RSC Adv.*, 2017, **7**, 10397–10406.
- B. Gibson, S. Carter, A. Fisher, S. Lancaster, J. Marshall and I. Whiteside, *J. Anal. At. Spectrom.*, 2015, **30**, 1814–1869.
- Y. Yang, Y. Wu, H. Zhang, M. Zhang, Q. Liu, H. Yang and J. Lu, *Fuel*, 2016, **181**, 951–957.
- J. Shen, H. Hu, M. Xu, H. Liu, K. Xu, X. Zhang, H. Yao and I. Naruse, *Fuel*, 2017, **207**, 365–372.
- B. Zhou, H. Zhou, J. Wang and K. Cen, *Fuel*, 2015, **150**, 526–537.
- G. Li, S. Li, Q. Huang and Q. Yao, *Fuel*, 2015, **143**, 430–437.
- L. Chao, C. F. You and D. K. Zhang, *Energy*, 2014, **69**, 562–570.
- M. J. F. Llorente, P. D. Arocas, L. G. Nebot and J. E. C. García, *Fuel*, 2008, **87**, 2651–2658.
- Z. Huang, L. Yan, L. Dan, Z. Zhou, Z. Wang, J. Zhou and K. Cen, *Energy Fuels*, 2013, **27**, 2049–2056.
- H. B. Vuthaluru and D. Zhang, *Fuel Process. Technol.*, 1999, **60**, 145–156.
- S. Arvelakis, H. Gehrman, M. Beckmann and E. G. Koukios, *Biomass Bioenergy*, 2002, **22**, 55–69.
- J. Barroso, J. Ballester, L. M. Ferrer and S. Jiménez, *Fuel Process. Technol.*, 2006, **87**, 737–752.
- S. Su, J. H. Pohl and D. Holcombe, *Fuel*, 2003, **82**, 1653–1667.
- A. Rushdi, A. Sharma and R. Gupta, *Fuel*, 2004, **83**, 495–506.
- S. S. Lokare, J. D. Dunaway, D. Moulton, D. Rogers, D. R. Tree and L. L. Baxter, *Energy Fuels*, 2006, **20**, 1008–1014.
- B. H. Lee, S. I. Kim, S. M. Kim, D. H. Oh, S. Gupta and C. H. Jeon, *Korean J. Chem. Eng.*, 2016, **33**, 147–153.
- M. Öhman, D. Boström, A. Nordin and H. Hedman, *Energy Fuels*, 2004, **18**, 1370–1376.
- F. Wigley, J. Williamson and G. Riley, *Fuel Process. Technol.*, 2007, **88**, 1010–1016.
- T. Madhiyanon, P. Sathitruangsak, S. Sungworagarn, S. Fukuda and S. Tia, *Fuel Process. Technol.*, 2013, **115**, 182–191.
- R. W. Bryers, *Prog. Energy Combust. Sci.*, 1996, **22**, 29–120.
- H. Schürmann, S. Unterberger, K. R. Hein, P. B. Monkhouse and U. Gottwald, *Faraday Discuss.*, 2001, **119**, 433.
- J. R. Wang, F. Chen, B. Zhao, X. Li and L. Qin, *Fuel Process. Technol.*, 2016, **155**, 209–215.
- B. Wei, X. Wang, H. Tan, L. Zhang, Y. Wang and Z. Wang, *Fuel*, 2016, **181**, 1224–1229.
- L. Xu, L. Jie, K. Yong, Y. Miao, R. Wei and T. Wang, *Energy Fuels*, 2014, **28**, 5640–5648.
- Z. J. Liu and R. Liu, *Earth Sci. Front.*, 2005, **12**, 315–323.
- N. Jing, Q. Wang, L. Cheng, Z. Luo, K. Cen and D. Zhang, *Fuel*, 2013, **104**, 647–655.
- I. Panagiotidis, K. Vafiadis, A. Tournlidakis and A. Tomboulides, *Appl. Therm. Eng.*, 2015, **74**, 156–164.
- H. Zhou, B. Zhou, H. Zhang and L. Li, *Energy Fuels*, 2014, **28**, 7701–7711.



- 36 T. Sahraoui, H. Belhouchet, M. Heraiz, N. Brihi and A. Guermat, *Ceram. Int.*, 2016, **42**, 12185–12193.
- 37 A. Y. AlOtoom, G. W. Bryant, L. K. Elliott, B. J. Skrifvars, M. Hupa and T. F. Wall, *Energy Fuels*, 1999, **14**, 41.
- 38 A. Y. Alotoom, L. K. Elliott, F. T. Wall and B. Moghtaderi, *Energy Fuels*, 2000, **14**, 97–109.
- 39 F. Ishom, T. Harada, T. Aoyagi, K. Sakanishi, Y. Korai and I. Mochida, *Fuel*, 2002, **81**, 1445–1451.
- 40 Y. Lu, Y. Wang, Y. Xu, Y. Li, W. Hao and Y. Zhang, *Appl. Therm. Eng.*, 2017, **121**, 224–233.
- 41 H. Naganuma, N. Ikeda, T. Ito, H. Satake, M. Matsuura, Y. Ueki, R. Yoshiie and I. Naruse, *Fuel Abstr. Curr. Titles*, 2013, **105**, 77–81.
- 42 Q. H. Wang, N. J. Jing, Z. Y. Luo, L. I. Xiao-Min and T. Jie, *J. China Coal Soc.*, 2010, **35**, 1015–1020.
- 43 F. Li, J. R. Qiu and C. G. Zheng, *J. Huazhong Univ. Sci. Technol., Med. Sci.*, 1996, **24**, 577–584.
- 44 P. Radhakrishnan, V. Zakkay and A. Agnone, *Combust. Sci. Technol.*, 1986, **50**, 271–281.
- 45 S. Zhang, C. Chen, D. Shi, J. Li, J. Wang, X. Guo, A. Dong and S. Xiong, *Proc. Chin. Soc. Electr. Eng.*, 2013, **33**, 1–12.
- 46 G. Li, C. A. Wang, Y. Yan, X. Jin, Y. Liu and D. Che, *J. Energy Inst.*, 2015, **29**, 48–56.

