



Title	Fate of the Chlorine in Coal in the Heating Process
Author(s)	Tsubouchi, Naoto; Mochizuki, Yuuki; Wang, Yanhui; Ohtsuka, Yasuo
Citation	ISIJ International, 58(2), 227-235 <a href="https://doi.org/10.2355/isijinternational.ISIJINT-2017-302">https://doi.org/10.2355/isijinternational.ISIJINT-2017-302</a>
Issue Date	2018-02-15
Doc URL	<a href="http://hdl.handle.net/2115/79331">http://hdl.handle.net/2115/79331</a>
Rights	著作権は日本鉄鋼協会にある
Type	article
File Information	ISIJ Int. 58(2)_ 227-235 (2018).pdf



[Instructions for use](#)

# Fate of the Chlorine in Coal in the Heating Process

Naoto TSUBOUCHI,<sup>1)\*</sup> Yuuki MOCHIZUKI,<sup>1)</sup> Yanhui WANG<sup>2)</sup> and Yasuo OHTSUKA<sup>2)</sup>

1) Center for Advanced Research of Energy and Materials, Faculty of Engineering, Hokkaido University, Kita 13 Nishi 5, Kita-ku, Sapporo, 060-8628 Japan. 2) Research Center for Sustainable Materials Engineering, Institute of Multidisciplinary Research for Advanced Materials, Tohoku University, Katahira 2-1-1, Aoba-ku, Sendai, 980-8577 Japan.

(Received on May 29, 2017; accepted on November 7, 2017; originally published in *Tetsu-to-Hagané*, Vol. 103, 2017, No. 8, pp. 443–450)

Pyrolysis of 29 coals with carbon contents of 71–92 mass% on a dry, ash-free basis (daf) has been performed mainly in a temperature-programmed mode at 10°C/min up to 800°C with a flow-type fixed bed quartz reactor, and some factors controlling HCl formation have been examined. The rate profiles of HCl formation exhibit at least three distinct peaks at around 260–360, 470–510 and 580–630°C, and the lowest temperature peak is present for 8 coals alone, whereas the middle and highest temperature peaks are common with almost all of the coals. The HCl profile is also affected by the size of coal particles and the height of coal particles in the fixed bed. Yields of HCl and char-Cl at 800°C for 28 coals except an American bituminous coal are 44–95 and 4–54%, respectively, and tar-Cl is as low as  $\leq 7\%$  in all cases. The chlorine distribution is almost independent of the heating rate in the range of 2.5–400°C/min and has no distinct relationship with carbon or chlorine content in coal, but HCl tends to increase with increasing amount of (Na + 2Ca) in coal with a corresponding decrease in char-Cl. When an Indonesian sub-bituminous coal is injected into an O<sub>2</sub>-blown entrained bed gasifier under pressure, there is an almost 1:1 relationship between carbon and nitrogen conversions, whereas the sulfur and chlorine are enriched in the remaining char, and the degree of the enrichment is higher with chlorine. The method of evaluating coal-Cl forms quantitatively using model chlorine compounds is proposed.

KEY WORDS: coal; pyrolysis; gasification; chlorine distribution; HCl formation; chlorine enrichment.

## 1. Introduction

Chlorine contained in coal—denoted as coal-Cl—is transformed into hydrogen chloride (HCl) via coal conversion processes, such as pyrolysis, combustion and gasification. This is feared to pose harmful consequences for a number of technologies currently under development, including integrated gasification combined cycle (IGCC) and integrated gasification fuel cell (IGFC) systems. Moreover, it is believed that HCl arising from chlorine contained in coke may cause corrosion of furnace installations.<sup>1)</sup> These problems arising from chlorine—while relatively new compared to long-studied phenomena arising from nitrogen or sulfur in coal—pose unavoidable challenges for the development of coal-utilization technologies that seek to achieve zero-emission operation.

The chlorine content of coal typically measures 2 000  $\mu\text{g/g-dry}$  or below, and in many cases lies around 1 000  $\mu\text{g/g-dry}$ . However, some coal produced in the U.K. or the U.S. contains high concentrations of chlorine in the vicinity of 10 000  $\mu\text{g/g-dry}$ .<sup>2)</sup> By way of contrast, in Japan—where environmental regulations are stringent—the coal used for electric power production is said to have a chlorine content of 500  $\mu\text{g/g-dry}$  or below, and the chlorine content of the

caking coal imported by Japan for coke has been determined to be in the range of 100–1 500  $\mu\text{g/g-dry}$ .<sup>1)</sup>

When coal is heated during combustion or gasification, pyrolysis occurs first, and the volatile portion is released; it has been reported that coal-Cl separates in the form of HCl during this process.<sup>3–5)</sup> According to Herod *et al.*,<sup>3)</sup> when 6 types of U.K.-produced coal containing high concentrations of chlorine in the range of 4 000–9 000  $\mu\text{g/g-dry}$  were heated at a rate of 2°C/min to 300°C, then maintained at that temperature for 24 hours, some 40–60% of the chlorine was converted to HCl; this was confirmed via mass analysis. In addition, Fourier-transform infrared spectroscopy was used to demonstrate that HCl and NH<sub>3</sub> are produced in nearly the same temperature region, somewhere in the vicinity of 450°C.<sup>5)</sup> Thus, we see that reports of HCl formation during coal pyrolysis have appeared in various places to date; however, there have been essentially no systematic studies involving multiple types of coal.<sup>6)</sup> Elucidation of the nature of HCl formation during pyrolysis would be of significant interest in conjunction with the formation of HCl in subsequent combustion or gasification steps. Moreover, HCl reacts easily with chemical species, such as alkali metals and mercury, and thus affects the evolution of these substances,<sup>7,8)</sup> and for this reason is thought to be closely related to the behavior of trace quantities of harmful metals in coal, a focus of current interest. Previous studies<sup>2,9,10)</sup> have reported that (a) the oxidation of elemental mercury during

\* Corresponding author: E-mail: tsubon@tagen.tohoku.ac.jp  
DOI: <http://dx.doi.org/10.2355/isijinternational.ISIJINT-2017-302>

pulverized coal combustion proceeds via heterogeneous reactions involving HCl and unburnt carbon in an electrostatic precipitator (90–300°C), and (b) when attempting to eliminate mercury from the gas produced by coal gasification using activated coal, the yield at 80°C is significantly improved in the presence of HCl.

For these reasons, in this study we subject 29 varieties of coal, of significantly different carbon and chlorine content, to temperature-programmed pyrolysis. In addition to conducting a detailed study of the effect of types of coal and reaction conditions on the chlorine distribution and the temperature dependence of the rate of HCl formation, we focus on the fate of nitrogen, sulfur, and chlorine during pressurized O<sub>2</sub>-blown coal gasification. Our primary objective is to clarify differences in the behavior of the distribu-

tions of these substances at high temperature and pressure. We also apply curve-fitting techniques to profiles of HCl formation during pyrolysis in an attempt to identify and quantify sources of HCl by their chemical form.

## 2. Experimental

### 2.1. Coal Samples

We used 29 varieties of coal as samples, ranging from lignite to anthracite. In most cases, upon obtaining coarse particles of a given sample, we air-dried the particles, then ground and sieved them to yield particles of size 150–250  $\mu\text{m}$ . Finally, we dried the particles for 1 h in N<sub>2</sub> at 108°C before using the sample in our experiments. **Table 1** lists results of elemental analysis and proximate analysis for

**Table 1.** Analyses of coals used.

Coal	Country <sup>a</sup>	Elemental analysis					Proximate analysis			
		(mass%-daf)					( $\mu\text{g/g-dry}$ )	(mass%-dry)		
		C	H	N	S	O	Cl	Ash	VM <sup>b</sup>	FC <sup>c</sup>
BFT	CAN	69.2	4.4	1.3	0.61	24.5	150	16.5	37.4	46.1
WMG	USA	69.9	4.9	1.1	0.49	23.6	250	5.4	45.0	49.6
BRU	IDN	71.1	5.1	1.6	0.61	21.6	230	3.1	45.6	51.3
BZP	USA	71.3	5.8	0.71	0.76	21.4	220	9.6	44.9	45.5
PSR	IDN	72.1	5.2	1.4	0.27	21.0	160	4.1	45.2	50.7
OBM	CAN	72.5	5.0	1.7	0.63	20.2	300	12.6	40.7	46.7
MBW	USA	73.4	5.2	1.6	0.43	19.4	420	4.3	38.0	57.7
WAN	USA	73.7	5.4	0.76	0.63	19.5	190	8.8	44.7	46.5
THY	JPN	74.8	6.0	1.3	0.25	17.6	440	11.3	42.6	46.1
BCN	USA	76.1	5.4	1.3	0.65	16.5	300	4.7	45.8	49.5
ILL	USA	76.9	5.3	1.3	5.5	11.0	810	14.3	40.1	45.6
BON	IDN	77.2	5.4	1.8	0.58	15.0	340	5.7	43.4	50.9
PRM	IDN	77.7	5.4	1.9	0.57	14.4	280	4.1	41.7	54.2
DTN	CHI	78.8	4.8	0.87	0.91	14.6	240	8.1	27.6	64.3
DRT	AUS	79.7	5.3	1.9	1.3	11.8	260	12.4	32.8	54.8
TGH	AUS	79.7	5.4	2.0	0.64	12.3	260	10.9	35.9	53.2
FSL	CHI	80.0	6.0	1.2	0.56	12.2	640	12.6	35.2	52.2
EML	SAF	80.1	5.3	1.9	1.1	11.6	230	12.9	29.7	57.4
LDL	AUS	81.1	5.4	2.1	0.57	10.8	470	8.3	33.9	57.8
LSN	USA	81.3	5.3	1.5	0.89	11.0	580	19.4	30.2	50.4
ENM	AUS	82.3	5.1	1.9	0.96	9.7	240	11.2	25.4	63.4
PIT	USA	82.4	5.4	1.6	2.4	8.2	840	9.0	37.8	53.2
MUR	AUS	83.7	5.5	1.8	0.75	8.2	370	12.0	28.0	60.0
UFT	USA	85.8	5.2	1.5	2.7	4.8	1 600	13.0	27.5	59.5
CNZ	CHI	88.4	4.1	1.7	0.44	5.4	380	15.0	9.4	75.6
YRB	AUS	89.5	3.7	2.0	0.78	4.0	1 300	9.8	7.3	82.9
JET	AUS	89.7	4.5	2.1	0.56	3.1	750	7.4	13.6	79.0
POC	USA	90.8	4.8	1.2	0.77	2.4	1 500	4.6	18.6	76.8
HGI	VIE	91.5	3.4	1.3	0.55	3.2	190	10.2	4.7	85.1

<sup>a</sup> CAN, Canada; USA, United States of America; IDN, Indonesia; JPN, Japan; CHI, China; AUS, Australia; SAF, South Africa; VIE, Vietnam.

<sup>b</sup> Volatile matter.

<sup>c</sup> Fixed carbon.

the coal samples used. The concentrations of carbon, chlorine and ash varied respectively over the ranges of 71–92 mass%-daf, 160–1 600  $\mu\text{g/g-dry}$  and 3.1–19 mass%-dry. For this study, we selected only coal samples within or near the van Krevelen coal band.

## 2.2. Pyrolysis Experiments and Analysis of Products

For pyrolysis, we used a flow-type fixed-bed quartz reactor.<sup>11,12</sup> In our experiments, a quartz cell filled with 2.5 g of coal was held in a quartz holder; after evacuation, ultra-high purity (>99.99995%) He was introduced into the system and a high-speed micro gas chromatograph (Agilent Technologies) was used to analyze the gas at the reactor outlet, with the system left untouched until the  $\text{N}_2$  and  $\text{H}_2\text{O}$  concentrations fell to 20 ppm or below. Then, the gas flow was switched to ultra-high purity (99.99995%)  $\text{N}_2$  and the heating of the sample was initiated. The temperature was controlled via K- or R-type thermocouples in contact with the bottom of the cell; the cell was heated to 800°C primarily at a rate of 10°C/min.

The HCl evolved by the pyrolysis process was analyzed online at 1-min intervals using an IR analyzer (Thermo Environmental Instruments).<sup>6,13–19</sup> The correlation coefficient for the calibration curves—constructed for HCl diluted by ultra-high purity  $\text{N}_2$  to 20, 50, 100, 200 ppm—was 0.99986. The gas resulting from  $\text{O}_2$  bomb combustion was captured in an aqueous solution of  $(\text{NH}_4)_2\text{CO}_3$  and an absorption spectrophotometer (Hitachi) was used to measure quantities of coal-Cl and of chlorine contained in tar (tar-Cl) and char (char-Cl).<sup>6,13,15,17–19</sup> The reproducibilities of our chlorine-content analyses were  $\pm 5$ ,  $\pm 7$  and  $\pm 4\%$  respectively for HCl, tar-Cl and char-Cl.

Using the results of these analyses, we express the yields of HCl, tar-Cl and char-Cl as fractions of the total content of chlorine present in the coal supplied. As discussed in more detail below, the total of these pyrolysis yields fell within the range of 90–108% for the 29 types of coal we considered, indicating that the chlorine balance was roughly appropriate. All gases at the reactor outlet were collected in a gas bag and investigated with a gas-detection tube in an effort to detect the presence of  $\text{Cl}_2$ . No  $\text{Cl}_2$  was detected (levels below 0.5 ppm) for any of the 29 types of coal we tested.

## 2.3. Elemental Analysis of Char after Gasification under Pressure

To investigate the behavior of coal-Cl under gasification following pyrolysis, we conducted an elemental analysis of a sample of high-pressure-gasified char (from BON raw coal) supplied by Babcock Hitachi, and then calculated the carbon conversion and the chlorine conversion. For comparison, we also determined the nitrogen conversion and sulfur conversion. Gasification was conducted at pressures of 0.9–2.6 MPa using an entrained bed gasification system with  $\text{O}_2/\text{coal}$  ratios in the range 0.66–0.83 kg/kg.<sup>15</sup> Gasification temperatures varied slightly depending on the experimental conditions; the average temperature at an  $\text{O}_2/\text{coal}$  ratio of 0.83 kg/kg and 2.6 MPa was 1 400°C.

## 2.4. Mineral Composition Analysis

To measure the mineral composition of our coal samples, high-temperature ash obtained by combustion at 815°C in

air was fully dissolved in a blended aqueous solution of HF and aqua regia, and metal ions were measured using an inductively coupled plasma emission spectrometer (ICP-ES) (Perkin Elmer).<sup>20</sup> In our ICP-ES analysis we measured 8 elements: Na, Mg, Al, Si, K, Ca, Fe and Ti. We assumed the chemical form of these metals would be oxides (with the exception of Ca in high-sulfur-content coal, which forms sulfate). Calculated values of ash contents fell in the range of 94–110% of the experimental ash-content values determined by proximate analysis (Table 1).

## 2.5. Solid-State Nuclear Magnetic Resonance Measurements

In previous research, the presence of NaCl hydrate has been noted among the chemical forms of coal-Cl,<sup>21,22</sup> and it has even been reported that the majority of coal-Cl forms hydrated NaCl.<sup>3,4,23</sup> Thus, to test for the presence of NaCl, we used a Fourier-transform nuclear magnetic resonance spectrometer (Chemagnetics) to conduct solid-state  $^{23}\text{Na}$ -NMR measurements of YRB coal.<sup>24</sup> We used the method of single-pulse excitation with magic-angle spinning (SPEMAS) with a rotation rate of 15 kHz and a  $^{23}\text{Na}$  resonance frequency of 79.4 MHz at 25°C and with the number of iterations in the range of 130 000–170 000. As an external reference, we used 1 N NaCl aqueous solution and set the chemical shift of the  $^{23}\text{Na}$  obtained to 0 ppm.

## 3. Results and Discussion

### 3.1. Temperature Dependence of HCl Formation

Figure 1 shows the temperature dependence of the rate of HCl formation in the process of temperature-programmed pyrolysis of WMG, DRT and HGI coals. For WMG coal, HCl evolution starts near 200°C; the formation rate exhibits a slight peak at 260°C and a major peak at 490°C, as well as a shoulder that appears in the range of 580–630°C. In contrast, for DRT coal the formation of HCl occurs at higher temperatures than for WMG coal; here the formation rate attains its maximum at 470°C, with a peak also visible near 580°C. For HGI coal, we observe just a single peak at 580°C.

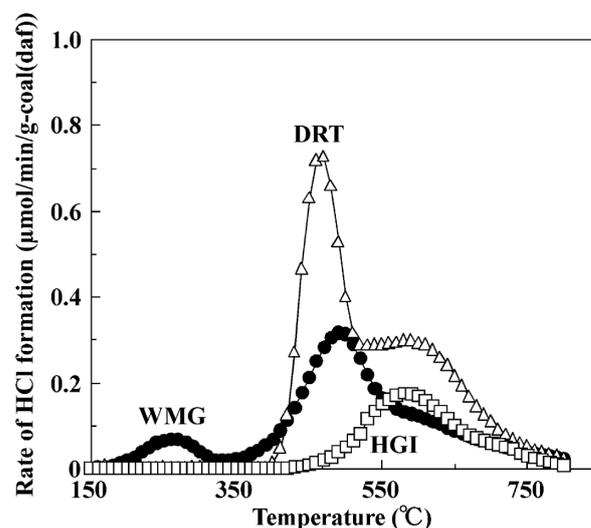


Fig. 1. Rate profiles for HCl formation in the temperature-programmed pyrolysis of WMG, DRT and HGI coals.

**Figure 2** shows results for JET and MBW coals. As is clear from comparison to Fig. 1, MBW coal exhibits an HCl formation profile similar to that seen for DRT coal. In contrast, for JET coal we see two peaks in the vicinity of 360°C and 510°C, as well as a shoulder in the range of 580–630°C.

**Table 2** collects the peak temperatures of HCl formation for all of the coals studied. From this table the following findings are evident:

- (1) A low-temperature HCl peak in the range of 260–360°C is present only for BFT, WMG, LDL, PIT, UFT, YRB, JET and POC coals.
- (2) An HCl peak in the range of 470–510°C is observed for all types of coal with the exception of CNZ, YRB and HGI coals.
- (3) The high-temperature HCl peak at 580–630°C is common to all types of coal.

Thus, we see that peaks in HCl formation may be broadly divided into three temperature regions: 260–360°C, 470–510°C and 580–630°C; the presence of peaks in the various regions depends on the type of coal.

### 3.2. Chlorine Distribution and Coal Type

Table 2 also lists the HCl yield at 800°C for all coals used. For all types of coal, Cl<sub>2</sub> was not detected (concentration, <0.5 ppm). Previous studies of pyrolysis also did not report formation of Cl<sub>2</sub>.<sup>3,4,23</sup> As we see from Table 2, the HCl yield depends strongly on the type of coal; yield values vary over the range of 23–95%, with discrepancies as large as over fourfold depending on the type of coal. We may summarize the results by saying that, with the exception of POC coal, more than half of the coal-Cl is converted to the gas phase. Yields of tar-Cl are smaller, in the range of 0–7%. This suggests that tar-Cl is converted to HCl via secondary reactions in the pyrolysis process. Yields for char-Cl vary from 4 to 72%, differing significantly for different coal types, as is true for the case of HCl. Total yields (HCl + tar-Cl + char-Cl) were reasonable, in the range of 90–108%. For char-Cl, multiple scenarios may be considered: perhaps the coal-Cl originally present in the coal remains unchanged, or with some partial change of form, in char; or perhaps a portion of the HCl evolved reacts with char

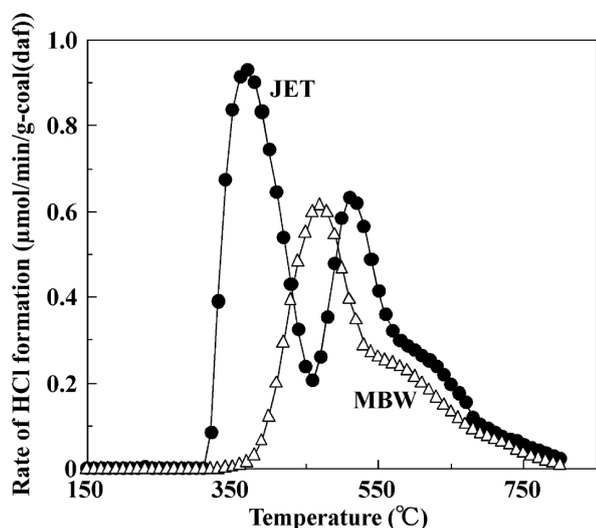
(minerals or carbon substances) and is incorporated into the solid phase. For example, calcium contained in coal exists in various forms—including limestone, dolomite, gypsum and ion-exchange Ca—while iron exists in the forms of pyrite, carbonate, ferric oxide and ion-exchange Fe and undergoes reduction reactions during pyrolysis. The reactions of HCl with CaCO<sub>3</sub> and FeO are expressed by Eqs. (1) and (2), respectively.



At 200–800°C, the change in standard Gibbs free energy is calculated to be in the range of –10 to –7.3 kcal/mol for Eq. (1) and in the range of –8.7 to –0.72 kcal/mol for Eq. (2). Thus, the formation of CaCl<sub>2</sub> or FeCl<sub>2</sub> is thermodynamically favored. The results of mineral composition analysis (section 2.4) indicate that, among all alkali metals and alkaline-earth metals present in our 29 coal samples, the

**Table 2.** Results for HCl formation from all coals examined.

Coal	Peak temperature observed in HCl profile (°C)			HCl yield at 800°C (%)
	260–360	470–510	580–630	
BFT	○	○	○	91
WMG	○	○	○	86
BRU		○	○	47
BZP		○	○	81
PSR		○	○	75
OBM		○	○	71
MBW		○	○	86
WAN		○	○	70
THY		○	○	95
BCN		○	○	91
ILL		○	○	44
BON		○	○	45
PRM		○	○	55
DTN		○	○	56
DRT		○	○	80
TGH		○	○	69
FSL		○	○	69
EML		○	○	77
LDL	○	○	○	66
LSN		○	○	63
ENM		○	○	60
PIT	○	○	○	57
MUR		○	○	56
UFT	○	○	○	74
CNZ			○	61
YRB	○		○	89
JET	○	○	○	82
POC	○	○	○	23
HGI			○	69



**Fig. 2.** Rate profiles for HCl formation in the temperature-programmed pyrolysis of JET and MBW coals.

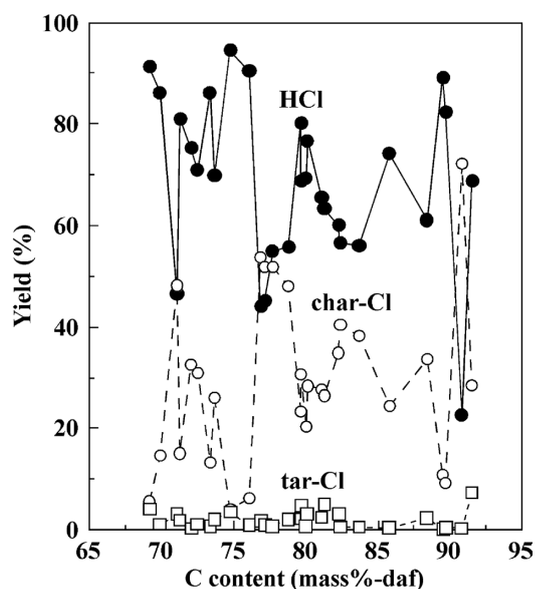
abundance of Ca is greatest (0.011–1.4 mass%-dry) for most types of coal, while the Fe content (0.23–2.2 mass%-dry) is also fairly significant. Reactions between HCl and surface OH groups in silicate or aluminosilicate may also occur. For non-minerals, we may also consider scenarios in which reactions of surface active sites of carbon in char with HCl proceed and the resulting chlorine is observed as char-Cl.

### 3.3. Effects of Coal and Mineral Compositions on Chlorine Distribution

**Figure 3** shows the effect of the carbon content of coal (C%) on the yield of HCl, tar-Cl and char-Cl. We see no clear relationship between HCl yield and C%; for example, although the C% values of THY and ILL coals are nearly equal (75 and 77 mass%-daf, respectively), their HCl yields differ by nearly 50%. We find similar behavior for tar-Cl and char-Cl. We also investigated the relationship between chlorine distribution and chlorine content (Cl%), but again found no clear correlation. We conclude that the yields of HCl, tar-Cl and char-Cl produced under the pyrolysis conditions of this study do not depend on the C% or Cl% of the feed coal.

As noted above, the majority of coal-Cl is thought to exist in the form of NaCl hydrate,<sup>3,4,23</sup> and Cl-XANES measurements indicating the presence of NaCl·nH<sub>2</sub>O, CaCl<sub>2</sub>·6H<sub>2</sub>O and organic hydrochlorides have been reported.<sup>21,22</sup> Thus, assuming that Na and Ca are present in the form of NaCl and CaCl<sub>2</sub>, we investigated the relationship between chlorine distribution and the quantity (Na+2Ca) in coal. The results are shown in **Fig. 4**. Although exceptions exist, the general trend is that, as the value of (Na+2Ca) increases, HCl increases and char-Cl decreases. We also attempted to plot the data with only Na or Ca on the horizontal axis, but found no clear trend of the sort visible in Fig. 4. Thus, it seems that the fate of coal-Cl during pyrolysis is best characterized in terms of the quantity (Na + 2Ca) in coal.

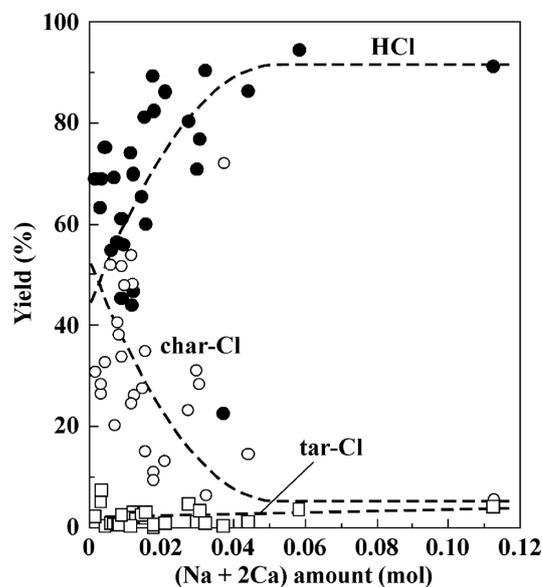
Several possible explanations for the fluctuations in the data of Fig. 4 may be considered. One explanation derives from the fact, noted above, that calcium in coal exists in



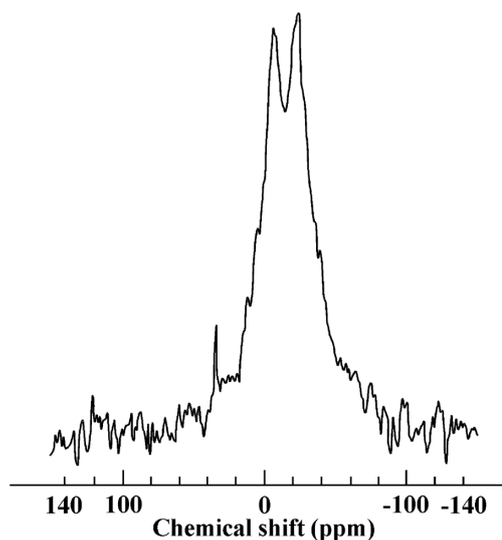
**Fig. 3.** Effect of carbon content in coal on chlorine distribution at 800°C.

the form of carbonate and hydrosulfate. The results of our mineral composition analysis (section 2.4) revealed that atomic Ca/Cl ratio was greater than 1 for almost all of the 29 coal samples used; thus, chlorides are relatively scarce, and limestone, dolomite and gypsum are the principal components. In contrast, atomic Na/Cl ratio was less than 1 in many cases, and thus chlorides may mainly constitute Na-containing compounds. The coal-Cl concentration has been reported to increase with increasing quantities of Na in coal.<sup>2)</sup> For these reasons, we performed solid-state <sup>23</sup>Na-NMR measurements of YRB coal, and the results are shown in **Fig. 5**. As expected, an Na<sup>+</sup>Cl<sup>-</sup> chemical shift appears at -5 ppm, confirming the presence of NaCl. Another possible explanation for the fluctuations in the data of Fig. 4 is the presence of Cl species other than inorganic chlorides, such as organic hydrochlorides.<sup>21,22)</sup>

In the pyrolysis of 15 types of caking coal for coke (10°C/min, 900°C), char-Cl was reported to increase as the Ca content of the coal increased.<sup>1)</sup> However, we observed no such trend in this study. A possible reason for the increase



**Fig. 4.** Relationship between the amount of (Na+2Ca) in coal and yield of HCl, tar-Cl or char-Cl at 800°C.



**Fig. 5.** Solid-state <sup>23</sup>Na-NMR spectrum for YRB coal.

in char-Cl is the formation of  $\text{CaCl}_2$  due to the progress of secondary reactions between HCl once-evolved and  $\text{CaCO}_3$  (Eq. (1)) or CaO. However, if the coal contains sulfur in abundance, secondary reactions between  $\text{H}_2\text{S}$  and  $\text{CaCO}_3$  or CaO occur with greater frequency, creating conditions unfavorable for the formation of  $\text{CaCl}_2$ . The S content of the caking coal for coke remains unclear,<sup>1)</sup> but is probably considerably smaller than the values used in this study for the quantity of S in coal (Table 1). Thermodynamic considerations suggest that the driving force for Eq. (1) is larger at a higher temperature; thus, even under the conditions of our experiments this could take place at high temperatures. In either case, there can be no doubt that the chemical form of coal-Cl is one important factor governing the HCl formation profile and the chlorine distribution.

### 3.4. Effect of Coal Sample State on HCl Formation

As is well known, the ash content and mineral composition of coal may vary depending on particle size. Since coal-Cl consists primarily of inorganic chlorides, the content of coal-Cl may also be affected by the coal particle size. For this reason, we used the method described in section 2.1 to prepare JET coal particles of size 55–75  $\mu\text{m}$  and subjected these to elemental analysis. The results indicated a Cl content of 930  $\mu\text{g/g-dry}$ , which is somewhat increased from the value of 750  $\mu\text{g/g-dry}$  observed for particles of size 150–250  $\mu\text{m}$ . This particle-size dependence merits a detailed future investigation.

Figure 6 shows the effect of particle size on the temperature dependence of the HCl formation rate. The HCl peak observed near 360°C for the original sample sizes (150–250  $\mu\text{m}$ ) is reduced by a factor of more than 4 for particle sizes of 55–75  $\mu\text{m}$ ; however, above 450°C the two rates do not differ significantly. The reason for this is presently unknown; however, it means that, if HCl in the vicinity of 360°C arises from a specific chloride compound, its content varies with coal particle size. Clarifying this point is a topic for future work; however, the results of Fig. 6 already suffice to indicate at the very least that particle size must be held constant for experiments testing HCl formation in pyrolysis.

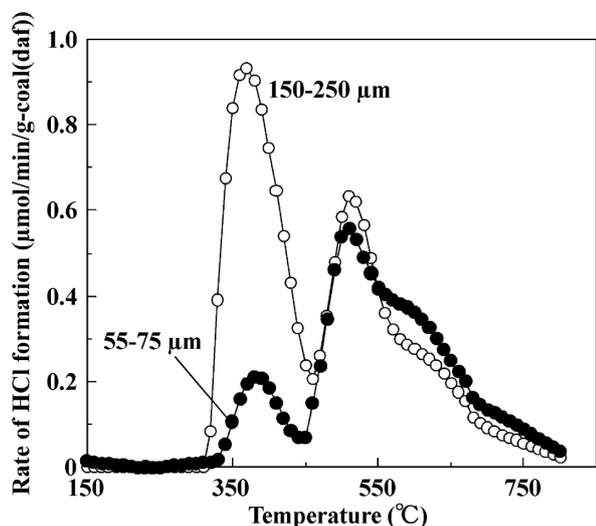


Fig. 6. Effect of the size of coal particles on HCl formation in the temperature-programmed pyrolysis of JET coal.

Figure 7 shows the effect of sample layer height on HCl formation from DRT coal. Here we have fixed the particle size in the range of 150–250  $\mu\text{m}$  and reduced the layer height to 1.5 mm from its original value of 8 mm. In other words, we have reduced the sample volume to decrease the layer height to approximately 1/5 of its original value. Comparing results for the two sample heights, we see that the rate of HCl formation does not differ much at 500°C or below, but is greater for the higher-height layer above this temperature, particularly in the vicinity of 580°C. Thicker sample layers facilitate more secondary reactions among initial products, so the results of Fig. 7 suggest that secondary reactions promoting HCl formation proceed near 580°C.

### 3.5. HCl Formation and Coal-Sample Heating Rate

To investigate the impact of heating rate on HCl formation, we measured the temperature dependence of the HCl yield in pyrolysis of DRT coal with heating rates reduced from the original rate of 10°C/min to 2.5°C/min and 5°C/min. The results are shown in Fig. 8. As the heating rate

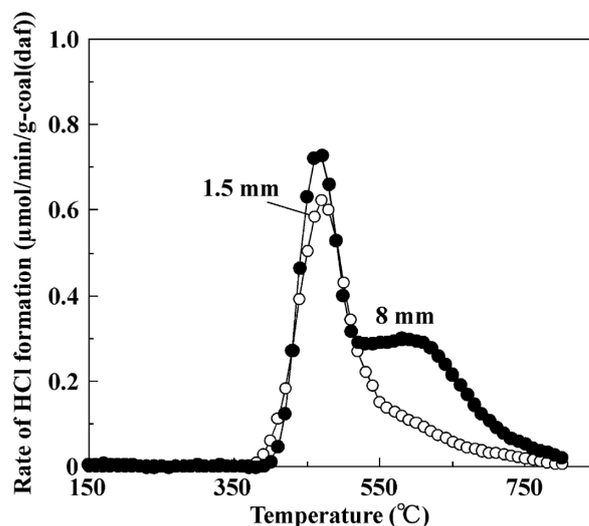


Fig. 7. Effect of the height of coal particles in the fixed bed on HCl formation in the temperature-programmed pyrolysis of DRT coal.

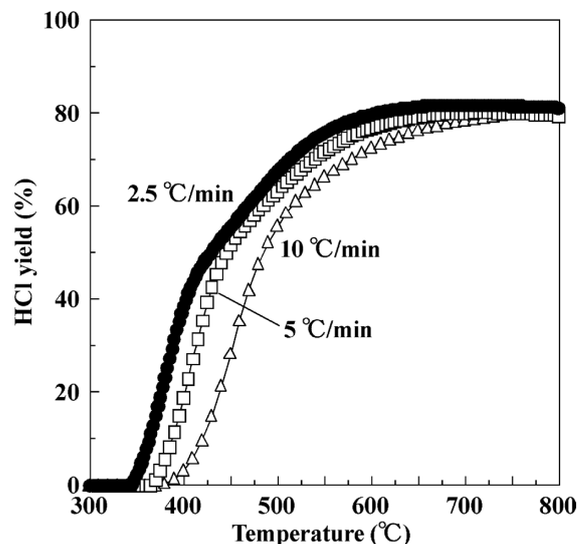


Fig. 8. Temperature change in HCl yield in the temperature-programmed pyrolysis of DRT coal at 2.5–10°C/min.

decreases, the HCl yield in the low-temperature region increases; however, at 800°C the yield is essentially independent of heating rate.

Based on the results of these temperature-programmed pyrolysis tests, we calculated the apparent activation energy of HCl formation,<sup>25)</sup> obtaining a value of 20 kcal/mol for HCl yields in the range of 10–40% and 25–30 kcal/mol for HCl yields in the range of 50–70%. Herod *et al.*<sup>3)</sup> heated at 2°C/min and performed pyrolysis at 300°C, concluding that the rate-controlling step of HCl formation corresponded to diffusion within micro-pores; thus, it is possible that the rate-determining step of HCl formation from bituminous coal involves diffusion in pores at low temperature region but crosses over to chemical reactions at higher temperatures of 450°C or above.

**Figure 9** shows the chlorine distributions resulting from pyrolysis of CNZ, YRB and JET coals at a heating rate of 400°C/min. When the heating rate is increased from the original value of 10°C/min to this higher rate, the HCl yields for all types of coal decrease slightly. On the other hand, the char-Cl yield at the higher heating rate increased 6% for CNZ coal and 13% for JET coal, but did not increase at all for YRB coal. Tar-Cl was not detected for any of the three coal types. As shown in Figs. 1 and 2, HCl formation increases with increasing temperature; however, the HCl yield at 800°C does not depend much on heating rate within the range of 2.5–400°C/min. When coal is injected into a pulverized coal combustion furnace or a gasification furnace, it is heated rapidly at a heating rate of 10<sup>4</sup>–10<sup>5</sup>°C/s to induce pyrolysis; future work is likely needed to study the fate of coal-Cl under these types of heating conditions.

### 3.6. Behavior of Chlorine in High-Temperature and -Pressure Gasification

As noted in section 3.2 above, although exceptions exist, in general at least 50% of coal-Cl is evolved in the form of HCl in pyrolysis up to temperatures of 800°C, with the majority of the non-HCl portion remaining in char. If all of this char-Cl were converted to HCl in the processes of combustion or gasification, the majority of coal-Cl would be converted to the gas phase. According to previous

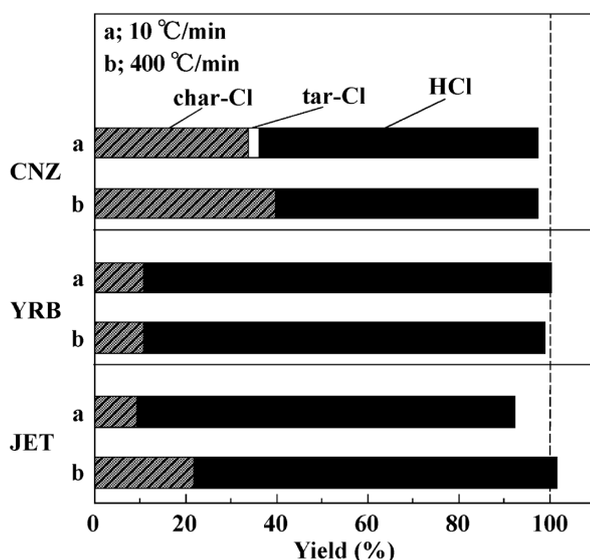


Fig. 9. Effect of heating rate on chlorine distribution at 800°C.

research,<sup>26)</sup> in an entrained bed gasification test of THY coal using Coal Partial Combustor (CPC) techniques (1 600°C, 2.0 MPa), approximately 90% of coal-Cl was converted to gas phase, although chlorine remaining in slag or char was also observed. In the pyrolysis of the THY coal used in this study, 95% of coal-Cl was released as HCl (Table 2); applying this value to gasification suggests that chlorine remaining in slag or char is produced by secondary reactions of HCl.

Thus, to investigate the behavior of coal-Cl in high-temperature, high-pressure gasification processes, we conducted elemental analysis of char obtained by the pressurized O<sub>2</sub>-blown coal gasification (section 2.3) and studied the relationship between the carbon and chlorine conversions. The results are shown in **Fig. 10**. For comparison, the figure also shows conversion for nitrogen and sulfur. The values plotted here, which are the quantities of carbon, chlorine, nitrogen and sulfur relative to the corresponding quantity of each substance in the char after pyrolysis under ambient pressure, represent conversion for the process of char gasification after volatile matters have been released. As can be seen from Fig. 10, whereas nitrogen in char gasifies to approximately the same extent as carbon, sulfur and chlorine are more resistant than carbon to gasification. The conversion for chlorine is particularly low: whereas some 80% of carbon is converted, 70–80% of char-Cl remains in char. Another way of saying this is that, compared to nitrogen and sulfur, chlorine is an easily condensed heteroelement. Results of this type strongly suggest secondary reactions between HCl once-evolved and alkali-metal compounds or carbon active sites, and corroborate reports that some portion of coal-Cl transitions to slag or char.<sup>26)</sup>

### 3.7. Quantifying Differences in the Chemical Forms of the Substances Responsible for HCl Formation

In this section we perform a curve-fitting analysis of profiles plotting HCl formation rate versus temperature during temperature-programmed pyrolysis (Fig. 1, *etc.*), attempting to quantify the HCl sources for various different forms.

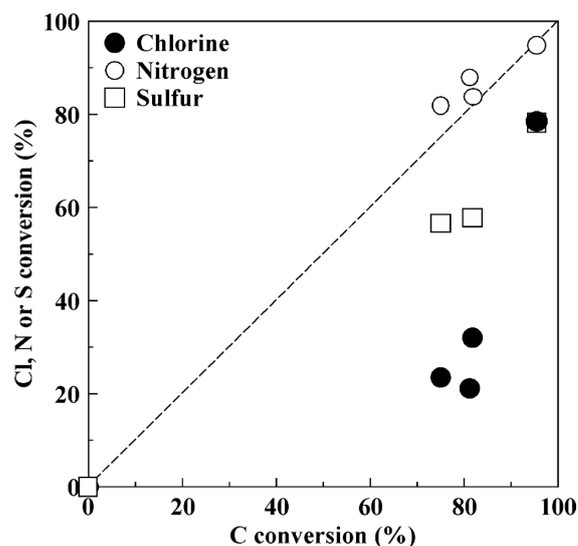


Fig. 10. Relationship between carbon conversion and chlorine, nitrogen or sulfur conversion during O<sub>2</sub>-blown entrained bed gasification under pressure.

For curve-fitting in the low-temperature region of 450°C or below, we used temperature and peak-width (full width at half maximum) data for previously-reported HCl formation peaks for chlorine model compounds.<sup>6,13</sup> As fitting parameters for HCl peaks in the range of 470–510 and 580–630°C, we computed the mean values of the peak temperature and peak width for all types of coal that exhibited the respective peaks (Table 2). The resulting values were: for the range of 470–510°C, a peak temperature of  $490 \pm 30^\circ\text{C}$  and a peak width of  $60 \pm 10^\circ\text{C}$ ; for the range of 580–630°C, a peak temperature of  $590 \pm 10^\circ\text{C}$  and a peak width of  $200 \pm 10^\circ\text{C}$ . The reproducibility of the present curve-fitting method was  $\pm 3\%$ .

Figures 11 and 12 show curve-fitting results for YRB and JET coals, which exhibited characteristic HCl peaks in the low-temperature region. For both of these coals, the particle size was 150–250  $\mu\text{m}$ . For YRB coal, we estimate values of 25% for organic hydrochlorides, 35% for  $\text{CaCl}_2 \cdot 6\text{H}_2\text{O}$ , 31% for  $\text{NaCl} \cdot n\text{H}_2\text{O}$  and 9% for the remainder (labeled in the figure as Cl-containing carbon species: C(Cl) or C(HCl)). In our previous studies,<sup>6,13</sup> HCl formation rates for YRB coal at temperatures of 210–280°C decreased significantly with water washing before pyrolysis; however, a peak at 360°C

remained, and ultimately 28% of coal-Cl eluted in water. If the curve-fitting results of Fig. 11 are applicable in their current form, this indicates the partial elimination of organic hydrochlorides and  $\text{CaCl}_2$  hydrate. The peak in the range of 580–630°C may be attributed to Cl-containing carbon species formed by secondary reactions between carbon active sites in char and HCl evolved. If 100 ppm HCl/ $\text{N}_2$  gas is exposed over lignite char at 500°C, the HCl reacts easily with carbon active sites, transforming into organic C–Cl bonds and/or carbon species containing HCl; in subsequent temperature-programmed desorption runs, an HCl-formation peak was reported in the range of 550–620°C.<sup>6</sup> As shown in Fig. 12, for JET coal with particle sizes of 150–250  $\mu\text{m}$ , estimated values were 53% for  $\text{NaCl} \cdot n\text{H}_2\text{O}$  and 47% for the remainder [total for 470–510°C (unidentified chlorine compounds) and 580–630°C]. For JET coal with particle sizes of 55–75  $\mu\text{m}$  we estimated a value of 13% for NaCl hydrates. After water washing of the latter sample, the peak at 360°C disappeared entirely<sup>6</sup> and approximately 15% of coal-Cl was removed,<sup>6</sup> indicating that all NaCl elutes in water. If this argument is correct, it implies that the elimination extent for NaCl hydrate should depend on the type of coal; this may signify differences in the structure or distribution of pores in which NaCl is present.<sup>2)</sup>

It remains unclear why the HCl peak at 260–360°C appears only for BFT, WMG, LDL, PIT, UFT, YRB, JET and POC coals (Table 2). As noted above, if the source of this HCl is attributed to organic hydrochlorides and inorganic chlorides  $\text{CaCl}_2 \cdot 6\text{H}_2\text{O}$  and  $\text{NaCl} \cdot n\text{H}_2\text{O}$ , our findings perhaps suggest that these chloride species are present only in the 8 types of coal listed above. Alternatively, it may be the case that, although organic hydrochlorides and inorganic chlorides are also present in other types of coal, all HCl molecules evolved in this temperature region are recaptured by mineral matters and/or carbon active sites in those types of coal.

Needless to say, the methods used in this section to quantify separately the sources of the HCl evolved in the pyrolysis are insufficient. Future studies are needed to corroborate the accuracy of the present curve-fitting analysis through a combination of measurements of Ca or Na ions in the filtrate remaining after water washing, investigations of changes in solid-state  $^{23}\text{Na}$ -NMR spectra before and after washing and chemical state analysis of coal-Cl using solid-state  $^{35}\text{Cl}$ -NMR.

#### 4. Conclusions

Using 29 types of coal with carbon content in the range of 71–92 mass%-daf and chlorine content in the range of 160–1 600  $\mu\text{g/g-dry}$ , we studied the temperature dependence of the rate of HCl formation during temperature-programmed pyrolysis and the effects of coal type and reaction conditions on chlorine distribution, and obtained the following conclusions.

(1) HCl formation peaks may be roughly separated into three temperature ranges: 260–360°C, 470–510°C and 580–630°C. Although instances of HCl formation at low temperatures were rare, HCl formation at intermediate temperatures was observed for essentially all types of coal.

(2) The yields of HCl and char-Cl at 800°C fell respec-

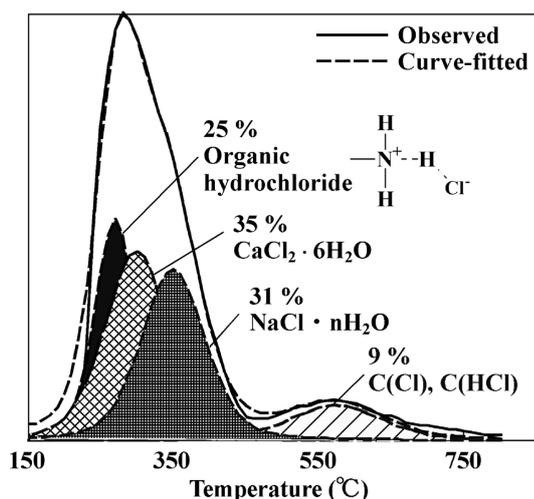


Fig. 11. Deconvolution of the profile for HCl formation in the temperature-programmed pyrolysis of YRB coal.

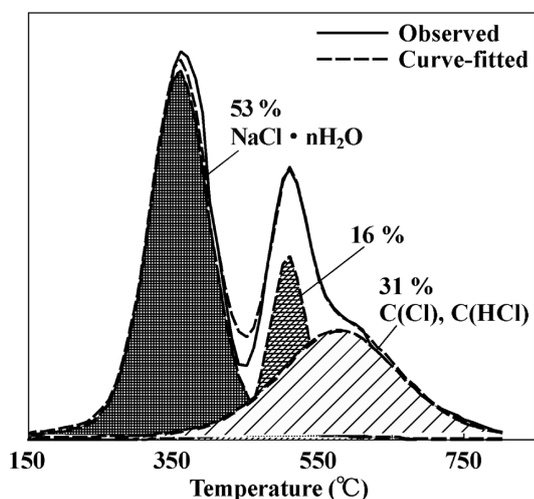


Fig. 12. Deconvolution of the profile for HCl formation in the temperature-programmed pyrolysis of JET coal.

tively in the ranges of 23–95% and 4–72%. On the other hand, for tar-Cl the yield never exceeded 7%. Thus, 90% or more of coal-Cl was transformed into HCl and char-Cl.

(3) As the quantity (Na + 2Ca) in coal increased, the HCl yield increased, while the char-Cl yield decreased.

(4) Although profiles plotting HCl formation rate versus temperature were affected by the size of coal particles and the height of coal particles in the fixed bed, the HCl yield at 800°C was essentially independent of heating rate over the range of 2.5–400°C/min.

(5) When an Indonesian sub-bituminous coal was injected into an O<sub>2</sub>-blown entrained bed gasifier under pressure, nitrogen and carbon contained in char gasified to the same extent, but sulfur and chlorine did not gasify, instead remaining in the char. In particular, we found that chlorine was the most easily condensed in char.

(6) We also showed that curve-fitting methods based on results for model chlorine compounds may be successfully applied to profiles of HCl formation during temperature-programmed pyrolysis to quantify different chemical forms of the chlorine present in coal.

#### Acknowledgment

This work was supported in part by the 24th Iron and Steel Institute of Japan (ISIJ) Research Promotion Grant. The authors gratefully acknowledge the assistance of Ms. Megumi Nishio in carrying out experiments.

#### REFERENCES

- 1) S. Nomura, K. Kato and Y. Maeno: *J. Jpn. Inst. Energy*, **82** (2003), 866.
- 2) R. M. Davidson: Chlorine and Other Halogens in Coal, IEAPER/28, IEA Coal Research, London, (1996).
- 3) A. A. Herod, N. J. Hodges, E. Pritchard and C. A. Smith: *Fuel*, **62** (1983), 1331.
- 4) G. Fynes, A. A. Herod, N. J. Hodges, B. J. Stokes and W. R. Ladner: *Fuel*, **67** (1988), 822.
- 5) D. Shao, E. J. Hutchinson, H. Cao, W. P. Pan and C. L. Chou: *Energy Fuel.*, **8** (1994), 399.
- 6) N. Tsubouchi, S. Ohtsuka, Y. Nakazato and Y. Ohtsuka: *Energy Fuel.*, **19** (2005), 554.
- 7) S. C. Mitchell: Hot Gas Cleanup of Sulfur, Nitrogen, Minor and Trace Elements, CCC/12, IEA Coal Research, London, (1998).
- 8) S. Benson: Fuel Cells - Use with Coal and Other Solid Fuels, CCC/47, IEA Coal Research, London, (2001).
- 9) N. Fujiwara: *Chem. Eng. Jpn.*, **70** (2006), 339.
- 10) M. Ozaki, S. Nagano, R. Ochiai, M. A. Uddin, E. Sasaoka and S. Wu: Proc. 44th Conf. of Japan Institute of Energy for Coal Science, The Japan Institute of Energy, Tokyo, (2007), 60.
- 11) N. Tsubouchi, Y. Ohshima, C. Xu and Y. Ohtsuka: *Energy Fuel.*, **15** (2001), 158.
- 12) N. Tsubouchi and Y. Ohtsuka: *Fuel*, **81** (2002), 1423.
- 13) N. Tsubouchi, S. Ohtsuka, H. Hashimoto and Y. Ohtsuka: *Energy Fuel.*, **18** (2004), 1605.
- 14) N. Tsubouchi, E. Kasai, K. Kawamoto, H. Noda, Y. Nakazato and Y. Ohtsuka: *Tetsu-To-Hagané*, **91** (2005), 751.
- 15) M. Takeda, A. Ueda, H. Hashimoto, T. Yamada, N. Suzuki, M. Sato, N. Tsubouchi, Y. Nakazato and Y. Ohtsuka: *Fuel*, **85** (2006), 235.
- 16) N. Tsubouchi, H. Hashimoto, N. Ohtaka and Y. Ohtsuka: *J. Hazard. Mater.*, **183** (2010), 116.
- 17) N. Tsubouchi, T. Saito, N. Ohtaka and Y. Ohtsuka: *Energy Fuel.*, **27** (2013), 87.
- 18) N. Tsubouchi, T. Saito, N. Ohtaka, Y. Nakazato and Y. Ohtsuka: *Energy Fuel.*, **27** (2013), 5076.
- 19) N. Tsubouchi, N. Ohtaka and Y. Ohtsuka: *Energy Fuel.*, **30** (2016), 2320.
- 20) N. Tsubouchi and Y. Ohtsuka: *Fuel*, **81** (2002), 2335.
- 21) F. E. Huggins and G. P. Huffman: Chlorine in Coal, Coal Science and Technology, Vol. 17, Elsevier, Amsterdam, (1991), 43.
- 22) F. E. Huggins and G. P. Huffman: *Fuel*, **74** (1995), 556.
- 23) N. J. Hodges, W. R. Ladner and T. G. Martin: *J. Inst. Energy*, **56** (1983), 158.
- 24) K. Sakanishi, E. Akashi, T. Nakazato, H. Tao, H. Kawashima, I. Saito and T. Takarada: *Fuel*, **83** (2004), 739.
- 25) T. Ozawa: *Bull. Chem. Soc. Jpn.*, **38** (1965), 1881.
- 26) K. Fujii: Behavior of Heteroatoms and Heavy Metals in Coal, Research Report on BRAIN-C Project (NEDO-C-9840), NEDO, Tokyo, (1999), 197.