

Mineralogical and Chemical Composition of Inorganic Matter from Marketed Illinois Coals

Ilham Demir, Randall E. Hughes, Philip J. DeMaris, and Rodney R. Ruch

LIBRARY JAN 2 9 2002 IL GEOL SURVEY

Illinois Minerals 121 2001

George H. Ryan, Governor

Department of Natural Resources Brent Manning, Director

ILLINOIS STATE GEOLOGICAL SURVEY William W. Shilts, Chief Natural Resources Building 615 East Peabody Drive Champaign, IL 61820-6964



Mineralogical and Chemical Composition of Inorganic Matter from Marketed Illinois Coals

Ilham Demir, Randall E. Hughes, Philip J. DeMaris, and Rodney R. Ruch

Illinois Minerals 121 2001

George H. Ryan, Governor

Department of Natural Resources Brent Manning, Director

ILLINOIS STATE GEOLOGICAL SURVEY William W. Shilts, Chief Natural Resources Building 615 East Peabody Drive Champaign, IL 61820-6964 (217) 333-4747 LIBRARY JAN 2 9 2002 IL GEOL SURVEY

ACKNOWLEDGMENTS

We thank R. A. Cahill, Y. Zhang, and J. D. Steele for chemical analyses of the samples. We also thank W. R. Roy and R. A Cahill for critically reviewing the manuscript. This study was supported, in part, by grants from the Illinois Department of Commerce and Community Affairs through the Illinois Coal Development Board and the Illinois Clean Coal Institute.

Editorial Bo	bard
Jonathan H. Goo	dwin, Chair
Michael L. Barnhardt 3. Brandon Curry Heinz H. Damberger William R.	Anne L. Erdmann David R. Larson Donald G. Mikulio Roy



Printed by authority of the State of Illinois/2001/600

Printed with soybean ink on recycled paper

CON	TEN	ITS
-----	-----	-----

ACKNOWLEDGMENTS	ii
ABSTRACT	1
INTRODUCTION	1
RELATED RESEARCH Inorganic Matter in Coal Coal Combustion Residues Chemical and Mineralogical Analyses of Coal and CCRs	2 2 3 3
OBJECTIVES	5
EXPERIMENTAL PROCEDURES Samples and Sample Preparation Chemical Analysis of Coal and CCR Samples Low-temperature Ashing of Coal Samples Water Leaching of LTA Samples Mineralogical Analysis of LTA Samples Mineralogical Analysis of CCRs and Limestone	5 5 6 7 7 7 8
RESULTS AND DISCUSSION Mineralogical Composition of Marketed Coals Relationships between Minerals and Minor and Trace Elements Combustion Behavior of Minerals in Coal Emission of HAPs from Power Plants Partitioning of HAPs between Fly Ash and Bottom Ash Temporal Variations in the Characteristics of Marketed Coals	8 8 10 11 15 22 23
SUMMARY AND CONCLUSIONS	25
REFERENCES	26
FIGURES	
Hooneo	

1	US production and use of CCRs for the year 1998	3
2	Coal regions of Illinois	5
3	Average mineral content of 35 marketed coals	10
4	Major mineral composition of mineral matter from 35 marketed Illinois coals	10
5	Clay mineral composition of the mineral matter from 35 marketed Illinois coals	11
6	Relationships of $r^2 > 0.5$ between elements and mixed-layered illite/smectite in	
	marketed Illinois coals	11
7	Relationships of r ² > 0.5 between elements and mixed-layered illite/smectite	
	plus illite in marketed Illinois coals	16
8	Relationships of $r^2 > 0.5$ between elements and kaolinite in marketed Illinois coals	17
9	Relationship between quartz and Mg in marketed Illinois coals	17
10	Calcite-Mn and calcite-Zn relationships in marketed Illinois coals	18
11	Relationship between pyrite and Hg in marketed Illinois coals	18
12	Mineralogical composition of CCRs from three types of plants	
	(FBC, PC, CYC) in Illinois	19

TABLES

1	Amounts and descriptions of samples of coal and CCRs from three types of	
	power plants	6
2	Mineralogical composition of marketed Illinois coals and CCRs	9
3	Chemical composition of 35 marketed Illinois coals from preparation plants and of	
	feed coals and combustion residues from four selected power plants	12
4	Mineral-element correlations in marketed Illinois coals	15

5	Concentrations of AI and 15 elements of environmental concern in coal, CCRs, and limestone samples from the four power plants and related combustion	
	mobility and relative enrichment values	20
6	Mobilities of HAPs during combustion	21
7	Relative distribution of HAPs among combustion ashes	23
8	Temporal variations in the mineralogical and chemical composition of marketed coals	
	from four selected Illinois preparation plants	24

ABSTRACT

The mineralogical and chemical compositions of marketed Illinois coals and coal combustion residues (CCRs) were investigated to obtain a better understanding of the relationship of these materials to air pollution and ash formation at coal-fired power plants. The coal samples were collected from 35 coal preparation plants in Illinois. The samples of CCRs (fly and bottom ashes), along with samples of feed coals, were collected from a fluidized bed combustion (FBC) plant, two cyclone combustion (CYC) plants, and a pulverized coal combustion (PC) plant. Average (mean ± standard deviation) abundances of minerals in the marketed coals were kaolinite, $2.07 \pm 0.56\%$; quartz, 2.06 ± 0.78%; mixed-layered illite/smectite, 1.57 ± 0.70%; pyrite, 1.56 ± 0.50%; illite, 1.29 $\pm 0.46\%$; calcite, 0.84 $\pm 0.78\%$; chlorite, 0.18 $\pm 0.13\%$; marcasite, 0.13 $\pm 0.12\%$; plagioclase, $0.07 \pm 0.05\%$; and K-feldspar, $0.01 \pm 0.01\%$. Nineteen minor and trace elements positively correlated with some of the minerals. The correlations that involved elements of environmental concern were Mn-calcite, F-illite and F-illite/smectite, Th-illite and Th-illite/smectite, Th-kaolinite, Cr-illite/ smectite, and Hg-pyrite. A comparison of the mineralogical composition of coals with CCRs suggested that combustion at the power plants partially converted guartz to silica glass. The clay minerals and feldspars were converted largely to aluminosilicate glass and, to a much lesser extent, to mullite. Pyrite and marcasite were converted to hematite, magnetite, and amorphous Fe oxides. Calcite was converted to anhydrite, gypsum, lime, and portlandite; some of the lime and portlandite was converted back to calcite through reaction with atmospheric CO. The minerals derived from calcite formed 30% and 71% of the FBC fly ash and bottom ash, respectively. The CCRs from the CYC and PC plants contained mostly amorphous material (78-98%). Other minerals in the CCRs from the four power plants were magnetite (0-4.95%), guartz (0.008-8.44%), mullite (0-5.09%), hematite (0-3.03%), and calcite (0-1.95%).

The combustion behaviors of 15 elements (As, Be, Cd, Co, Cr, F, Hg, Mn, Ni, P, Pb, Sb, Se, Th, and U) of environmental concern in coal were estimated by comparing the analytical data from the feed coals and CCRs from the power plants. These elements were partly retained in the CCRs and partly mobilized and airborne upon combustion. Fluorine and Mn mobilities were greatest (>50% of the original amounts in the feed lost) at the FBC plant, and F, Hg, and Se mobilities were greatest at the CYC and PC plants. The majority of the other elements were less mobile (<25%). The mobilized portions of the elements may not have been entirely emitted into the atmosphere; they may have been partially condensed on hardware at the cool side of the power plants. The concentrations of the 15 trace elements were greater in the fly ashes than in the bottom ashes, with a few exceptions. Some of the elements could be commercially produced from the CCRs.

Samples collected at four selected coal preparation plants at a 4-year interval showed some variations in the composition of the coals over time. These results give a general idea about the limited reliability of using outdated coal-quality data from a given mine for assessing the environmental and economic impacts of burning the coal currently produced from the mine.

INTRODUCTION

The industrial use of coal is greatly affected by the amount and variety of its inorganic content. The inorganic matter is the main source of pollutants emitted to the atmosphere during coal combustion or leached from wastes from various coal utilization processes. The inorganic matter in the coal, however, may also be a source of useful by-products.

Data on the mineralogical composition of coal and CCRs have broad applications in studies relating to commercial uses of Illinois coals and of the residues from cleaning, combustion, and conversion of these coals. Interpreting combined mineralogical and chemical data helps identify the modes of occurrence of toxic elements in coal; such information is essential for designing coal cleaning or post-combustion control methods to meet the requirements of current or future environmental regulations. Improved understanding of the fate of minerals in different types of boilers also increases the ability to design improved methods to predict slagging, fouling, and ash properties from the mineralogical and chemical composition of feed coal. Ash fusion temperatures, slagging, and fouling are strongly dependent on the mineralogical composition of coal and, therefore, cannot be accurately

estimated from chemical analysis of coal or ash alone. The minerals in coals melt over a range of temperatures, which reflects the refractoriness of each type of mineral. The difference between the ash fusion temperature predicted from bulk chemistry and the actual or correct value increases as the partitioning of elements among different minerals increases. The CCR data can also help generate new ideas about increasing their commercial utilization.

RELATED RESEARCH

Inorganic Matter in Coal

The inorganic matter content of run-of-mine coal is highly variable and depends on geological conditions and the mining techniques used. Inorganic matter in coal is present mostly as discrete minerals with five different modes of occurrence: (1) microscopically disseminated inclusions within macerals; (2) layers or partings of variable thickness; (3) nodules, including lenticular and spherical concretions; (4) fissures, including cleat and other fracture and void fillings; and (5) megascopic rock fragments found within the coal bed because of faulting, slumping, or related disturbances and from intentional or accidental mining of the roof or floor strata (Harvey et al. 1983, Harvey and Ruch 1986). Physical coal-cleaning processes normally remove large, discrete grains and layers of inorganic matter more efficiently than they remove finely disseminated minerals. Mineralogical compositions of inorganic matter in channel and core samples of Illinois coals were investigated by Rao and Gluskoter (1973), Ward (1977), and Harvey et al. (1983). Minerals that commonly occurred in measurable quantities in these Illinois coal samples included illite, mixed-layered illite/smectite, kaolinite, pyrite, quartz, and calcite. Minerals with rare occurrences and trace abundances in the samples included chlorite, feldspars, zircon, sphalerite, dolomite (ankerite), siderite, barite, anhydrite, gypsum, rosenite, melanterite, coquimbite, jarosite, hematite, rutile, and apatite.

Inorganic minerals host the elements of environmental concern in coal. However, there are a few exceptions; for example, S can be part of the organic structure, and Cl can partly be adsorbed on the organic matter in exchangeable ionic form. Most elements of environmental concern in coal are categorized as trace elements (generally <200 mg/kg). Gluskoter et al. (1977) and Harvey et al. (1983) reported on the trace element contents of a large number of mostly channel and core samples of Illinois coals. Demir et al. (1998) used principal component analysis to determine the modes of occurrence of environmentally critical elements in channel samples of Illinois coals.

With respect to coal utilization, the National Committee for Geochemistry (National Research Council 1980) identified three categories of elements of environmental concern: (1) those of greatest concern—As, B, Cd, Pb, Hg, Mo, and Se; (2) those of moderate concern—V, Cr, Ni, Cu, Zn, and F; and (3) those of minor concern-Li, Na, Sr, Ba, Mn, Co, Ge, Cl, Br, Ra, Po, Rn, Th, and U. These categories were based upon known toxicity, levels of occurrence of each element in coal, and anticipated mobility upon combustion or upon disposal of ash. Sixteen elements in coal (As, Be, Cd, Cl, Co, Cr, F, Hq, Mn, Ni, P, Pb, Sb, Se, Th, U) are among the 189 hazardous air pollutants (HAPs) mentioned in the 1990 Clean Air Act amendments (U.S. Public Law 101-549, 1990). It should be pointed out that some of the elements in the National Research Council list were not considered HAPs by the amendments. Partitioning of HAPs and other elements among the CCRs and flue gas is highly variable (Swaine 1990, Clarke and Sloss 1992, Wesnor 1993, Davidson and Clarke 1996) because of the variation in the types and operational conditions of combustion units and in the chemical properties and modes of occurrences of the elements. Analytical errors and difficulties in obtaining representative samples also add to the variability of data on the retention or emission of potentially toxic elements from power plants. The HAP provisions of the 1990 amendments to the Clean Air Act focus on non-utility industries, including metals, petrochemicals, and papermaking (Moore 1994). A risk analysis by the U.S. Environmental Protection Agency (USEPA 1998) concluded that, for now, only Hg emission from the coal-fired electrical utilities is of potential health risk and thus requires further investigation; a final regulation on Hg will be issued by 2004.



Figure 1 US production and use of CCRs for the year 1998 (plotted using data from ACAA 1999). FGD, flue gas desulfurization.

Coal Combustion Residues

Utilities in the U.S. annually generate108 million short tons of CCRs, 58% of which is fly ash (fig. 1). Current and potential commercial uses of CCRs include treatment of soil and of mine and industrial wastes; admixture in cement, concrete, and grout; making bricks and other ceramic products; mineral filler; fill materials in civil engineering projects including road base/subbase; blasting grit; roofing granules; making wallboards; snow/ice control; and extraction of valuable materials (Clarke 1992, DeBarr et al. 1996, Dreher et al. 1996, Hughes et al. 1996, Wirtz and Bukowski 1996, Roy et al. 1997, ACAA 1999, Barbieri et al. 1999, Dhir and Jones 1999, Niewiadomski et al. 1999, Sloan et al. 1999). However, only about 29% of the total CCR generated in the U.S. is used for commercial purposes (fig. 1); the rest is disposed of on land or in abandoned mines.

Chemical and Mineralogical Analyses of Coal and CCRs

Methods for determining the chemical composition of coal and CCRs have been well established at the Illinois State Geological Survey (Gluskoter et al. 1977, Harvey et al. 1983, Demir et al. 1994). Methods for mineralogical analysis of these same materials, however, vary greatly in their usefulness for solving various problems associated with coal use. Application of an improved x-ray diffraction (XRD) method for mineralogical analysis of coal and CCR samples was an important task of this study. Therefore, the mineralogical analysis methods are discussed in considerable detail.

The methods of analyzing minerals in coal are based on XRD analysis, chemical analysis, or a combination of the two. Methods based solely on conventional XRD analysis contain large errors, as do those based solely on chemical analysis (called normative methods). Errors in determining the mineralogy of a coal based solely on the chemical analysis generally result from assigning the chemical elements to the wrong normative minerals. Another shortcoming of normative methods is that they are costly and generally lack procedures for estimating errors (Slaughter 1989). As a result, most investigators have favored methods that combine XRD, chemical analysis, and sometimes other procedures such as scanning electron microscopy. An interlaboratory study of quantitative coal mineral analysis by computer-controlled scanning electron microscopy indicated that the mineralogical composition of coal cannot be determined precisely using this technique alone (Galbreath et al. 1996). Beginning in the 1970s, investigators at the Illinois State Geological Survey began to apply combined XRD and chemical methods for analysis of the non-clay mineral fraction of coals. It is important to note that the methods described in the literature generally lack provisions for the analysis of some important minerals, particularly marcasite, mixed-layered illite/ smectite, and mixed-layered kaolinite/expandables.

There is extensive literature relating to the determination by XRD of mineral matter in coal. The following developments were especially relevant to this present study:

- 1. Development of a procedure for removing the combustible material from coal at low temperatures, called low-temperature ashing (Gluskoter 1965).
- 2. Development of an accurate and precise method for the determination of pyritic sulfur in coals (ASTM 1995).
- 3. Development of methods for the quantitative determination of minerals by XRD (Snyder and Bish 1989).
- Development of NEWMOD[®] (Reynolds 1985) and its application to the quantitative analysis of clay minerals (Hughes and Warren 1989, Reynolds 1989, Hughes et al. 1994, Moore and Reynolds 1997).

The mineral concentrations in bulk coal samples often are too small to be accurately and precisely determined. Therefore, XRD of the low-temperature ashes (LTAs) of coals has been the principal method for mineralogical determinations for more than 30 years. The method yields a virtually certain identification of the minerals in a sample. In addition, variations in the relative intensity of XRD peaks for some minerals commonly can be used to estimate their chemical composition. The precision of different versions of XRD analysis varies widely, as do claims of their accuracy (Snyder and Bish 1989, Moore and Reynolds 1997). However, as a result of refinements during the 1980s, precise and accurate determination of non-clay minerals such as pyrite (and marcasite), quartz, dolomite, and calcite became possible (Harvey et al. 1983, Demir and Harvey 1990). A later study established analytical procedures that meet the requirements of most investigators for mineralogical analysis of inorganic matter in coal (Kruse et al. 1994). These procedures reduced or eliminated the following errors associated with historical XRD procedures:

- 1. Preferred orientation error. Two advances have been made that largely eliminate the error resulting from preferred orientation of mineral particles. The first is to grind the sample in a manner that creates a more random orientation of the crystallites. The McCrone micronizers® in our laboratory yield powders with crystallite sizes in the optimal 5-µm to 10-µm range. The second advance involves using side-loading powder holders and the capability to spin the samples during exposure to the x-ray beam to minimize preferred orientation effects. The typical practice of the past decade has been to use a random powder pack for non-clay mineral analysis and a bulk smear sample for analysis of clay minerals (Hughes et al. 1994). Historically, XRD analysis of clay minerals has been run on the segregated <2-µm fraction of a sample. This practice was developed partly because instruments at the time gave low peak intensities for most clay minerals and partly because clay minerals were thought to be essentially associated with the finest particle fractions, <2 µm by definition. In reality, the clay minerals in coal occur in a wide range of particle sizes. For example, illite/smectite crystallites commonly are <1 µm, while some kaolinite crystallites exceed 50 µm. As a result, errors in excess of an order of magnitude can occur from XRD analysis of samples prepared as <2-µm aggregates. This error has been eliminated by grinding the clay minerals to approximately the same size and by analyzing a bulk sample that includes all of the particles (Hughes and Warren 1989, Hughes et al. 1994).
- 2. Standard mineral error. The standard mineral error results from the measurement or calculation of a reference intensity for a "standard" mineral (Russell and Rimmer 1979, Hughes et al. 1994, Moore and Reynolds 1997). In the past, mineralogists analyzed standard samples with the highest purity possible and refined their procedure whenever a better standard was located. For clay minerals, Reynolds (1985) developed a computer program (NEWMOD®) that makes it possible to calculate a reference intensity ratio (RIR) for any clay mineral with a layer structure. This program makes it easy to calculate XRD peak ratios for the wide range of chemical variations observed in natural samples. Furthermore, NEWMOD® gives peak intensities relative to non-clay minerals. Therefore, the calculated intensities can be used to check the clay mineral

content, which is arrived at by proportioning the difference of 100% minus the non-clay mineral content. The net result of these improvements is to minimize the errors associated with the mismatch between reference standards and minerals in coal and thus eliminate the need for an almost infinite number of natural standard samples.

- 3. X-ray absorption error. The x-ray beam must pass through crystallite to be diffracted into the detector and recorded as a series of peaks. However, x-rays are absorbed in the crystallite in proportion to the atomic weight of its elements. Therefore, large grains of pyrite and marcasite will seem to be present at less than their true content because their diffracted x-rays are partially absorbed. In theory, highly precise and accurate analyses of these types of minerals require methods of sequential grinding to reduce errors to a minimum. In practice, it is possible to obtain adequate precision and accuracy with a single grinding in a McCrone[®] micronizer to produce 5-µm to 10-µm crystallites.
- 4. Error in matching XRD peak intensities with mineral contents. Attempts to calculate the content of all minerals directly from the XRD spectrum of a random bulk powder sample generally give erroneous results. This error was eliminated by applying a method that used chemical analysis of pyritic sulfur and XRD peak intensities from a random bulk pack to calculate the amount of pyrite, quartz, and calcite in LTA samples (Harvey et al. 1983, Demir and Harvey 1990). The bulk clay mineral fraction can then be calculated as 100% minus the non-clay mineral fraction. By extension, the amount of calcite (confirmed to be present in the sample by XRD) can be calculated also from CO₂ or CaO analyses by traditional chemical methods, providing that the concentration of other major Ca minerals in the samples is insignificant. For most coals, this calculation leaves only quartz among the major non-clay minerals in the sample to be determined solely with XRD.

OBJECTIVES

The objectives of this study were (1) to determine the mineralogical and chemical composition of the inorganic matter in marketed, conventionally cleaned Illinois coals from all operating mines and of CCRs from selected power plants; (2) to correlate the chemical composition with the mineralogical composition of the coals; (3) to investigate the fate of the inorganic constituents during combustion at the power plants; and (4) to determine the temporal variations in the mineralogical and chemical composition of marketed coals from selected mines.

EXPERIMENTAL PROCEDURES

Samples and Sample Preparation

Marketed coal samples were collected from each of 35 coal preparation plants that operated in the different regions of the Illinois coal field (fig. 2) between 1992 and 1996. In 1996, sets of fly ash, bottom ash, and feed coal samples were collected from a fluidized bed combustion (FBC) plant, two cyclone combustion (CYC) plants, and a pulverized coal combustion (PC) plant; a sample of the limestone used in the FBC plant was also collected. The coal sample from the FBC plant was a blend of coals from 17 different mines. Coal samples from the CYC and PC plants came from one of the 35 preparation plants that had been sampled four years earlier in 1992. In addition, three other preparation plants among the 35 were sampled again four years after the first



Figure 2 Coal regions of Illinois. Southeastern Illinois coal field is divided into Herrin (4H) and Springfield (4S) Coal seams. sampling. Results for these three samples and for the coal samples from the PC and CYC plants were compared with results from samples collected at the same plants four years earlier to study the temporal variations in the mineralogical and chemical composition of marketed Illinois coals.

Details of location, collection, processing, and storage of the marketed coal samples from the coal preparation plants are described in Demir et al. (1998). As for sampling at the power plants, the company staffs at the PC and CYC plants used automatic samplers to collect the samples as part of their routine sampling. For the FBC plant, we used a sampling shovel to collect 15 increments of the coal from the belt carrying coal to the boiler. To sample the FBC fly ash, the ash stream was directed to a sampling bag for about 30 sec through an opening at the particulate collection system. We sampled the FBC bottom ash and crushed limestone by using a sampling shovel to collect 15 to 20 increments of the materials from widely spaced locations of their stock piles. The power plant samples were weighed and described after a visual inspection (table 1). Representative splits of the coal and coarse-grained bottom ash samples were separated by riffling, splitting, and then grinding to -60 mesh (<250 µm) particle size in a Holmes mill. The fly ash samples were already -60 mesh in particle size and did not need to be ground. Small splits of the ground coal and bottom ash samples were prepared for chemical, mineralogical, and microscopic analyses by riffling and splitting.

Plant type	Sample type	Amount (kg)	Sample description
FBC	coal	11.3	<0.95-cm (3/8 in) particle size
	fly ash	8.2	very fine particle size, light gray color
	bottom ash	10.9	<0.64-cm (<1/4 in) particle size, mostly yellowish and grayish particles, and small number of black particles
	limestone	11.3	crushed and off-white color
CYC1	coal	4.1	<4.76-cm (-4 mesh) particle size
	fly ash	3.6	fine particle size, dark gray color
	bottom ash	5.4	<0.95-cm (<3/8 in) particle size, black color, and high moisture content because of quenching in water
CYC2	coal	4.5	<4.76-cm (-4 mesh) particle size
	fly ash	3.2	fine particle size, dark gray color
	bottom ash	5.4	<0.95-cm (<3/8 in) particle size, black color, and high moisture content because of quenching in water
PC	coal	5.4	<4.76-cm (-4 mesh) particle size
	fly ash	3.6	fine particle size, light gray color
	bottom ash	6.8	<0.95-cm (<3/8 in) particle size, dark gray to black color, and high moisture content because of quenching in water

 Table 1
 Amounts and descriptions of samples of coal and CCRs from three types (FBC, CYC, PC) of power plants.

Chemical Analysis of Coal and CCR Samples

The samples of coals and CCRs were analyzed using the following methods:

x-ray fluorescence spectrometry: Al, Ba, Ca, Fe, Ge, K, Mg, Mn, Mo, Na, P, Si, Sn, Sr, Ti

energy-dispersive x-ray: Ba, Ge, Mo, Sn, Sr, Zr

instrumental neutron activation: As, Br, Ce, Co, Cr, Cs, Dy, Eu, Ga, Hf, La, Lu, Mn, Rb, Sb, Sc, Se, Sm, Ta, Tb, Th, U, W, Yb, Zn

atomic absorption spectrometry: Cd, Cu, Li, Ni, Pb, Zn, Hg (by cold vapor atomic absorption)

optical emission spectrometry: Ag, B, Be, Ge, Mo, Pb, Sn, Tl, V

pyrohydrolysis-ion chromatography: F

inductively coupled plasma-mass spectrometry: Cd, Pb

ASTM methods: moisture, ash, volatile matter, C, H, N, O, total S and S forms, CI, calorific value

Note that some of the elements were determined using more than one method. In such cases, the average or the most reliable value was used. Chemical analyses of the samples were performed following strict quality assurance and quality control rules to ensure reproducibility and accuracy of the data. The quality assurance and control plan (ISGS 1994) included standard operating procedures for sample preparation and for the analytical work. Blank, standard, and replicate samples were analyzed to ensure data reliability.

Low-temperature Ashing of Coal Samples

Generating the LTA of a coal sample was the first step for analysis of coal minerals. The LTAs were generated at about 150°C in an activated oxygen-plasma atmosphere that had been created by passing oxygen into a radiofrequency field (Gluskoter 1965). The minerals in the LTA were essentially unaltered from their original state in the coal. In the standard procedure, 10-20 g of the sample were spread evenly to a depth of 3 mm over the bottoms of Pyrex boats (12×4 cm in size). The boats with samples were put in a vacuum desiccator for a minimum of 24 hr for drying. The dry samples were then placed in the asher, and the pressure in the asher was reduced to about 1 torr. The radiofrequency field was then introduced at 37.5 W per chamber, and oxygen gas was introduced at a rate of 20 cm³/min to produce the plasma. The ashing temperature was monitored occasionally with an infrared remote thermometer. The samples were stirred twice and weighed once each day. When the sample weight no longer changed, the ashing was complete.

Water Leaching of LTA Samples

Previous research indicated that the LTA samples contained some water-soluble, inorganic elements that reduced the accuracy of the mineralogical analysis (Kruse et al. 1994). To improve the accuracy of the mineralogical analyses, each LTA sample was washed with water to remove the water-soluble fraction. To accomplish the water extraction, each LTA sample was dispersed in demineralized water (1 g of LTA in 500 mL of water). After 12–16 hr, the LTA and water mixture was centrifuged, the clear supernatant was poured off, and the solid fraction was dried overnight at 70°C and weighed.

Mineralogical Analysis of LTA Samples

The water-extracted LTA samples were mineralogically analyzed by XRD. The data were then converted to whole-coal basis. For the XRD analysis, the LTA samples were micronized for 15 min with 10 mL of water buffered with a few drops of sodium dithionite in a McCrone[®] grinder. The samples were then transferred to 50-mL centrifuge tubes and centrifuged for 20 min at 2,000 rpm, which resulted in the separation of a clear supernatant. After the clear supernatant was discarded, the paste was mixed thoroughly, and some of it was smeared on glass XRD slides. The remaining paste was dried overnight at 70°C, mixed with a mortar and pestle, and then packed into an end-loading sample holder (a random bulk pack). The random bulk pack samples were x-rayed with a Scintag[®] diffractometer, and the peak areas were calculated with the DMS[®] software that is part of the instrument operating system. The smear slides were x-rayed after air-drying and after at least two days exposure to ethylene glycol, heating for 1 hr at 350°C, and then heating to 550°C, which produced the strong 14 Å XRD peak for detection of small amounts of chlorite. The results of XRD analysis of bulk packs and smear slides were fairly comparable. The bulk pack XRD analysis was the preferred method for determination of non-clay minerals. The XRD analysis of smear slides was

used for clay mineral analysis. The detection limit of calcite determination with XRD was 0.5 wt% or perhaps even greater. Therefore, the calcite content was calculated from the chemically determined CaO content of coal. This approach was judged to be reasonable because no other Ca minerals were detected in the samples by the XRD, and the amount of Ca associated with clay minerals and plagioclase was insignificant relative to that associated with calcite. The steps for calculating the mineralogical composition of the coal samples from the XRD spectra and chemical analyses were as follows:

- 1. Deconvolute XRD patterns and transfer data electronically to a computer spreadsheet.
- From the percent of insoluble LTA and chemically determined pyritic-S (S_{py}) content of coal, %(pyrite + marcasite) in LTA = %S_{py} in LTA × 1.871.
- 3. From bulk pack XRD peak areas, calculate the percentages of quartz, K-feldspar, plagioclase, calcite, pyrite, and marcasite on a 100% basis.
- 4. Decimal non-clay minerals (DNC) in the LTA = (%[pyrite + marcasite] from step 2) ÷ (%[pyrite + marcasite] from step 3).
- 5. Percentage of each non-clay mineral in the LTA = (its percentage in the non-clay fraction) × (DNC).
- 6. From smear slide XRD peak areas, calculate the percentages of mixed-layered illite/smectite, illite, kaolinite, and chlorite on a 100% clay basis.
- 7. Percentage of each clay mineral in the LTA = (percentage in the clay fraction) \times (1-DNC).
- 8. Recalculate the percentage of calcite in LTA using the %LTA and chemically determined CaO contents of coal, replace this chemical calcite content for the XRD calcite content from step 5, and adjust the percentages of other minerals from steps 5 and 7 accordingly.
- 9. Percentage of each mineral in whole $coal = (\%LTA \div 100) \times (percentage in the LTA).$

Mineralogical Analysis of CCRs and Limestone

The CCRs from the power plants contained large amounts of amorphous (glass) material and variable amounts of lime and anhydrite, which react with water. Therefore, a somewhat different procedure of mineralogical analysis was followed for the fly and bottom ashes as well as for the limestone sample: the water extraction step was omitted, 8 wt% dolomite was added to the sample as the internal standard, and the samples were micronized in propanol. Among a small number of available standards, the dolomite was the best because its peaks did not overlap with other mineral peaks, and 8 wt% dolomite produced enough XRD counts to ensure accuracy without diluting the other peaks too much.

RESULTS AND DISCUSSION

Mineralogical Composition of Marketed Coals

On average, the most abundant minerals in the 35 marketed Illinois coals were kaolinite ($\bar{x} = 2.07\%$) and quartz ($\bar{x} = 2.06\%$)(table 2, fig. 3). K-feldspar was the least abundant mineral; its average concentration was 0.01%. The clay mineral fraction generally was more abundant than the non-clay fraction in the 35 samples (fig. 4), and the clay mineral composition was slightly skewed towards kaolinite (fig. 5). The average clay content was 5.11% on whole coal basis or 52% on an LTA basis. This proportion of the clay fraction in the LTAs of the marketed Illinois coals is only slightly less than the 56% value for channel samples of Illinois coals reported in Harvey and Ruch (1986), which suggests that conventional cleaning does not significantly alter the ratio of clay to non-clay minerals compared with that of standard channel samples.

Table 2 Mineralogical composition of marketed Illinois coals and CCRs. All values are in weight percent on a dry weight basis.¹

Sample no.	Region	I/S	L	Kaol	Chl	Qtz	Pyr	Marc	Calc	K-f	Plag
C32773	1	1.67	0.79	1.50	0.07	2.24	1.42	0.24	0.71	0.05	0.07
C32774	1	0.43	0.53	0.93	0.05	1.26	2.35	0.26	0.27	<0.01	0.04
C32777	1	1.12	1.20	1.78	0.16	2.66	1.47	0.03	4.46	<0.01	0.08
C32778	1	1.79	1.23	2.31	0.23	1.82	0.91	<0.01	0.87	<0.01	0.06
C32782	1	0.61	1.10	1.69	0.06	3.62	2.00	0.06	1.27	<0.01	0.11
C32783	1	1.94	1.18	2.02	0.12	1.37	2.00	0.37	0.52	< 0.01	0.10
C32785	1	1.83	1.21	1.77	0.09	2.16	1.54	0.29	0.64	0.03	0.05
C32797	1	1.87	1.06	1.89	0.13	1.87	1.64	0.06	0.77	<0.01	0.06
C32814	1	0.69	0.47	0.66	0.02	1.19	1.71	0.01	0.87	<0.01	0.07
C32779	2	1.87	0.77	1.34	0.21	2.25	1.68	0.09	0.77	<0.01	<0.01
C32794	2	1.09	1.20	2.34	0.17	2.36	2.02	0.12	0.79	<0.01	0.04
C32798	2	2.30	1.17	2.00	0.10	2.70	1.78	0.16	1.62	<0.01	0.06
C32800	2	2.70	1.17	2.23	0.20	2.17	1.41	0.18	0.82	<0.01	0.05
C32813	2	3.11	2.50	2.19	0.32	3.07	2.42	0.12	0.96	<0.01	0.05
C32815	2	1.65	1.03	1.88	0.13	2.95	2.17	0.11	1.52	<0.01	<0.01
C32784	3	1.81	1.00	2.34	0.18	1.41	1.26	0.09	0.20	<0.01	0.05
C32795	3	0.59	1.29	1.81	0.15	1.20	0.31	<0.01	0.18	<0.01	0.03
C32796	3	2.95	1.42	3.22	0.44	4.06	0.73	0.02	0.61	<0.01	0.11
C32799	3	1.06	2.65	3.21	0.44	1.42	0.34	0.01	1.89	<0.01	0.18
C32801	3	1.08	1.44	2.44	0.28	1.26	1.17	0.07	0.41	<0.01	0.07
C32802	3	1.39	1.73	2.09	0.60	1.32	1.42	0.10	0.71	<0.01	0.05
C32803	3	1.69	1.34	2.45	0.07	1.53	1.31	0.09	0.50	<0.01	0.05
C32661	4H	1.41	1.22	1.84	0.11	1.80	1.78	<0.01	0.34	<0.01	<0.01
C32664	4 H	2.04	1.55	2.86	0.12	1.84	2.05	0.09	0.36	<0.01	0.06
C32665	4H	1.33	1.41	1.83	0.12	2.29	1.83	0.05	0.36	<0.01	0.03
C32771	4H	2.94	1.63	2.15	0.10	2.33	2.11	0.03	0.79	<0.01	0.05
C32776	4H	1.44	0.88	1.57	0.12	1.87	1.71	0.20	0.86	0.05	0.08
C32662	4S	1.48	1.20	1.64	0.19	1.15	1.01	0.06	0.23	<0.01	0.09
C32663	4S	1.31	1.18	2.80	0.19	1.77	1.30	0.53	0.29	<0.01	0.05
C32772	4S	1.10	1.19	2.19	0.18	2.24	1.50	0.19	0.80	0.02	0.09
C32775	4S	1.09	0.84	1.83	0.03	1.51	1.97	0.13	0.25	<0.01	0.14
C32780	4S	0.53	2.05	2.42	0.27	1.47	1.59	0.21	0.82	<0.01	0.06
C32781	4S	0.93	1.01	1.71	0.05	2.86	1.63	0.32	0.48	<0.01	0.07
C32793	4S	2.13	1.90	2.18	0.37	3.85	1.43	0.20	0.68	0.07	0.29
C35564	5	2.05	1.09	2.92	0.28	1.57	1.39	0.23	1.70	<0.01	0.06
Mean ²	all	1.57	1.29	2.07	0.18	2.06	1.56	0.13	0.84	0.01	0.07
SD ²	all	0.70	0.46	0.56	0.13	0.78	0.50	0.12	0.78	0.01	0.05

Power plant coals

Sample no.	Sample type	I/S	1	Kaol	Chl	Qtz	Pyr	Marc	Calc	K-f	Plag
C35300	FBC feed	1.78	2.07	2.70	0.16	1.70	0.89	0.30	0.86	<0.01	0.00
C35301	CYC1 feed	0.46	1.47	2.14	0.04	3.03	1.12	<0.01	1.00	<0.01	0.13
C35302	CYC2 feed	0.60	1.54	2.13	0.30	2.79	1.06	0.05	1.05	<0.01	0.03
C35303	PC feed	0.35	2.32	3.47	0.19	1.61	1.27	0.07	1.00	<0.01	0.02

Power plant ashes

Sample no.	Sample type	Mul	Qtz	Calc	Hemat	Magn	Anhyd	Gyps	Lime	Portl	Glass
C35304	FBC-fly ash	ND	8.44	ND	3.03	0.69	17	ND	2.30	11	57
C35308	FBC-bottom ash	ND	8.18	1.95	ND	ND	16	2.03	40	13	19
C35305	CYC1-fly ash	1.87	7.57	0.75	2.96	4.03	ND	5.02	ND	ND	78
C35309	CYC1-bottom ash	ND	1.29	ND	ND	0.44	ND	0.50	ND	ND	98
C35306	CYC2-fly ash	2.29	6.83	1.11	2.16	3.79	0.60	ND	ND	ND	83
C35310	CYC2-bottom ash	ND	1.27	ND	ND	0.48	ND	0.29	ND	ND	98
C35307	PC-fly ash	5.09	0.008	ND	2.07	4.74	0.71	ND	ND	ND	87
C35311	PC-bottom ash	ND	0.97	0.37	1.97	4.95	ND	ND	ND	ND	92
C35312	FBC - limestone3.4	% illite,	1.5% kad	olinite, 1	.5% chlori	te, 5% q	uartz, 86%	calcite, a	nd 4.0%	dolomite	Э

¹ I/S, mixed layered illite/smectite; I, illite; Kaol, kaolinite; Chl, chlorite; Qtz, quartz; Pyr, pyrite; Marc, marcasite; Calc, calcite; K-f, K-feldspar; Plag, plagioclase; Mul, mullite; Hemat, hematite; Magn, magnetite; Anhyd, anhydrite, Gyps, gypsum; Portl, portlandite; Glass, amorphous material (calculated by difference); SD, standard deviation; H, Herrin Coal; S, Springfield Coal; ND, not detected

² For values below detection limits, one-half of the detection limits were used in statistical calculations of means and standard deviations.



Figure 3 Average mineral content of 35 marketed coals. Numbers above bars are relative standard deviations ((standard deviation/mean) × 100). Keys: Kaol, kaolinite; I/S, illite/smectite; I, illite; ChI, chlorite; Qtz, quartz; Pyr, pyrite; Marc, marcasite; Calc, calcite; Plag, plagioclase; K-f, K-feldspar.



Figure 4 Major mineral composition of mineral matter from 35 marketed Illinois coals.

Relationships between Minerals and Minor and Trace Elements

By using the mineralogical and chemical data (tables 2 and 3), statistical correlations were sought among the minerals and minor and trace elements in the marketed Illinois coals. The statistical analysis yielded many significant correlations (table 4). The square of the correlation coefficient (r²) for some of the correlations was greater than 0.5 (figs. 6–10). Statistical correlations may or may not indicate that an element is located in the mineral with which it is positively correlated. For example, the correlation between quartz and Mg indicates that most of the quartz and the Mg-bearing mineral (illite/smectite) was incorporated into the coal by the same geological and geochemical conditions. The same interpretation can be made for the correlation between calcite and Zn. Calcite and the main Zn-bearing mineral (sphalerite) were incorporated into the coal by the



Figure 5 Clay mineral composition of the mineral matter from 35 marketed Illinois coals.



Figure 6 Relationships of $r^2 > 0.5$ between elements and mixed-layered illite/smectite in marketed Illinois coals.

same geochemical conditions, namely epigenetic precipitation; sphalerite was apparently present in the samples, but its concentration was too small to be detected by the XRD method. There is some evidence that F may be part of the illite and mixed-layered illite/smectite structure (Thomas et al. 1977). Manganese is known to replace Ca in the calcite structure, and, therefore, the Mn/calcite correlation was not surprising. Although the overall Hg-pyrite correlation is weak (fig. 11), there appear to be two populations, one above and one below the regression line, each with a different slope. This evidence suggests that Hg may be partitioned distinctly between pyrites of different origins (syngenetic vs. epigenetic). The modes of occurrence of most trace elements in coal are subject to multiple interpretations unless constricted by separating and directly analyzing individual minerals in coal.

Combustion Behavior of Minerals in Coal

Upon combustion, coal minerals (fig. 3) were converted to other minerals and amorphous phases that made up the CCRs (fig. 12). Among the minerals present in coal, only some of the quartz survived the combustion process; the rest of the quartz that was present in the coal was converted to silica glass. The clay minerals and feldspar were converted largely to aluminosilicate glass and,

	nless indi	cated Utili	in lasima			0000																
$ \begin{array}{cccccccccccccccccccccccccccccccccccc$	o no. F	Region' %	Moisture	%Ash	Btu/Ib	%Volatile	%Fixed C	%C	H%	N%	0%	%Total S	%Organic S	%Pyritic S	%Sulfate S	Ag	A	As	<u>ه</u>	Ba	Be	
$ \begin{array}{c ccccccccccccccccccccccccccccccccccc$	2773	-	14.02	8.72	12.810	41.22	50.06	71.47	5.04	1.21	9.41	4.14	3.03	0.97	0.14	v	8,997	1.3	135	4	1.0	
$ \begin{array}{c ccccccccccccccccccccccccccccccccccc$	774	-	13.45	7.07	13,270	40.87	52.06	74.58	5.38	1.31	8.09	3.58	1.56	1.46	0.56	$\overline{\nabla}$	5,345	20	116	17	4.0	
$ \begin{array}{c ccccccccccccccccccccccccccccccccccc$	111	-	14.94	14.52	12,450	38.60	46.88	69.44	5.04	1.22	6.63	3.14	2.05	1.01	0.08	v	8,574	5.1	119	66	1.2	
	1778	-	17.36	9.80	12,710	36.68	53.53	71.95	5.16	1.42 1	0.08	1.60	0.98	0.54	0.07	V	10,955	10	202	20	2.2	
	782		14.99	11.62	12,500	38.98	49.40	69.45	5.17	1.25	8.61	3.90	2.45	1.25	0.20	ī	8,574	2.4	106	38	<1.0	
	2783	-	15.59	9.74	12,690	40.50	49.76	70.02	5.36	1.26	9.21	4.42	2.96	1.34	0.12	ī	9,421	2.2	155	33	1.4	
	2785		13.98	9.75	12,740	41.43	48.81	70.22	5.34	1.30	9.22	4.17	3.00	1.05	0.12	v	9,738	2.3	178	43	1.5	
$ \begin{array}{c c c c c c c c c c c c c c c c c c c $	2797		15.73	10.29	12,730	40.27	49.45	71.11	5.06	1.23	8.68	3.62	2.45	0.99	0.18	V	10,215	4	147	4	- 12	
$ \begin{array}{c ccccccccccccccccccccccccccccccccccc$	2814		11.39	6.00	13,840	45.68	48.31	76.20	5.49	1.28	7.35	3.68	2.41	1.09	0.18	<u>v</u> .	3,811	6.4	83	ee e	2.0	
$ \begin{array}{cccccccccccccccccccccccccccccccccccc$	2779	5	12.73	9.63	12,750	39.39	50.98	71.19	5.13	1.24	8.62	4.20	3.07	1.04	0.09	v.	9,685	1.	114	40	0.1>	
$ \begin{array}{c ccccccccccccccccccccccccccccccccccc$	2794	0	11.13	10.52	12,690	38.49	51.00	71.22	4.92	1.31	8.37	3.67	2.29	1.24	0.14	īv	10,585	2.3	128	36		
$ \begin{array}{ccccc} 2 & 1130 & 1200 & 364 & 3019 & 7018 & 437 & 130 & 335 & 234 & 134 & 105 & 022 & 61 & 1305 & 2 & 127 & 64 & 105 \\ 31 & 3 & 3 & 931 & 113 & 1200 & 364 & 3019 & 7018 & 437 & 139 & 133 & 237 & 234 & 139 & 102 & 012 & 61 & 1305 & 2 & 127 & 64 & 105 \\ 31 & 3 & 3 & 931 & 112 & 3200 & 334 & 323 & 323 & 324 & 123 & 127 & 101 & 012 & 61 & 10273 & 124$	2798	5	12.38	13.16	12,290	38.31	48.53	68.84	4.97	1.27	8.28	3.48	2.13	0.20	0.15	v	12,543	2.2	128	82	<1.0	
$ \begin{array}{c c c c c c c c c c c c c c c c c c c $	2800	0	11.93	11.39	12,600	38.42	50.19	70.45	4.94	1.20	8.39	3.63	2.49	0.92	0.22	ī	11,696	2	137	49	<1.0	
	2813	5	10.80	14.70	12,090	36.49	48.80	67.69	4.72	1.26	8.16	3.47	1.97	1.45	0.05	ī	15,507	2.4	127	69	1.4	
778 3 100 5/1 3/3 0/2 0/3 1/2 0/3 1/2 0/3 1/2 0/3 1/2 0/3 1/2 0/3 1/2 0/3	2815	2	12.65	12.03	12,420	37.16	50.82	69.42	4.93	1.41	8.48	3.73	2.19	1.41	0.12	ī	11,855	ო	126	4	<1.0	
770 81 7 73 0.55 0.75 <td>2784</td> <td>e</td> <td>10.00</td> <td>8.13</td> <td>13,330</td> <td>35.62</td> <td>56.24</td> <td>74.81</td> <td>5.22</td> <td>1.57</td> <td>8.48</td> <td>1.79</td> <td>0.77</td> <td>0.74</td> <td>0.13</td> <td>v</td> <td>10,056</td> <td>18</td> <td>74</td> <td>32</td> <td>1.8</td> <td></td>	2784	e	10.00	8.13	13,330	35.62	56.24	74.81	5.22	1.57	8.48	1.79	0.77	0.74	0.13	v	10,056	18	74	32	1.8	
779 3 91 1610 72.96 519 711 705 73 711 712 72 711	2795	e	10.06	5.76	13,780	35.25	58.99	78.02	5.11	1.63	8.75	0.73	0.55	0.17	0.02	īv	7,992	3.6	57	26	1.0	
9 3 9 7 14 7 8 9 7 14 7 8 10 0 7 14 7 8 10 9 1 14 7 8 10 9 11 10 11 10 1	9620	e	9.81	16.10	12.120	32.00	51.90	69.01	4.64	1.49	7.71	1.05	0.53	0.42	0.10	īv	21,541	9.8	49	81	1.0	
	6620	3	9.67	11.42	12.730	34.40	54.18	72.42	4.82	1.94	8.62	0.76	0.52	0.21	0.03	īv	15,983	17	58	91	1.0	
3 66 93 93 933 51 14 75 31 21 73 14 75 31 17 36 17 37 37 31 30 36 17 37 37 31 37 31 37 31<	2801		10.38	8.36	13.280	35,93	55.71	75.08	5.19	1.56	7.82	1.98	1.11	0.70	0.17	ī	10.479	10	64	38	1.0	
$ \begin{array}{cccccccccccccccccccccccccccccccccccc$	2802) m	8.56	9.36	13.090	38,34	52.30	73.74	5.11	1.41	7.25	3.12	2.01	0.88	0.23	ī	10.056	3.1	78	36	<1.0	
#i 7i0 Bi1 132-0 552 741 750 214 710 014 <1240 014 <1240 014 <1240 014 <1240 014 <1240 014 <1240 014 <1240 014 <1240 014 <1240 014 <1240 014 <1240 014 <1240 014 <1240 014 <1240 014 <1240 014 <1240 014 <1240 014 <1240 014 <1240 014 <1240 014 <1240 014 <1240 014 <1240 014 <1240 014 <1240 014 <1240 014 <1240 014 <1240 014 <1240 014 <1240 014	2803		10.08	9.19	13.150	36.62	54.19	74.10	5.11	1.46	7.59	2.54	1.56	0.79	0.19	īv	10.638	4.1	78	45	1.2	
866 41 500 987 3336 7390 987 1314 47 50 47 131 47 51 41 50 51 41 50 51 41 50 51 41 50 51 141 50 141 44 50 51 141 50 51 141 50 51 141 50 51 141 50 51 141 50 51 141 50 51 141 50 51 141 50 51 50 51 50 51 50 51 50 51 50 51 5	2661	4H	7 60	8 17	13 240	36.60	55 23	74 99	5 11	1 48	7.36	2 89	1 76	0 04	0 14	7	9.262	34	47	33	16	
660 670 671 717 500 717 500 717 500 717 500 717 710 711 711 711 711 71 711 711 711 711 711 711 710 711	2664	4H	5 80	9.87	13 150	36.67	53 45	73.80	0000	1.41	7.06	2.87	1.64	1.18	0.05	; .	12.490	5.4	41	36	1.4	
	0665	H	6.80	0.20	13 150	36.31	54 30	74 17	5.05	1 44	7 23	2 73	161	1.05	0.07	7	11 114	44	90	68	1	
min min <td>1770</td> <td>Ŧ</td> <td>0.00</td> <td>10.57</td> <td>12 620</td> <td>35.06</td> <td>50.37</td> <td>71.65</td> <td>4.75</td> <td>134</td> <td>6 75</td> <td>0 03</td> <td>1 59</td> <td>00-1</td> <td>0.10</td> <td>7 7</td> <td>13 972</td> <td></td> <td>8</td> <td>6 FC</td> <td> i -</td> <td></td>	1770	Ŧ	0.00	10.57	12 620	35.06	50.37	71.65	4.75	134	6 75	0 03	1 59	00-1	0.10	7 7	13 972		8	6 FC	 i -	
5 5 7 1 7 1 1 1 1 1 1 1 1 1	776	F F	10.15	0.07	13 180	37.92	50 B1	72 08	514	1 44	8.05	0 1 C	1 90	113	010	7 7	9.579	2.7	76	6	- 12	
35 45 6.70 896 13.100 35.36 7.55 7.55 7.56 7.16 7.17 7.8 7.10 9.16 7.10 9.16 7.10 9.16 7.10 9.16 7.11 9.16 7.11 9.16 7.11 9.16 7.11 9.16 7.11 9.16 7.11 9.16 7.11 9.16 7.11 9.16 7.11 9.16 7.11 9.16 7.11 9.16 7.11 9.16 7.11 9.16 7.11 9.17 7.11 9.16 0.11 0.	660	F V	00- 00-	7.00	13 530	34 88	58.10	77 00	л - с - с	1 59	7.75	151	0.85	0 59	0.08	, .	9 209	14	40	2 G	14	
775 55 57.0 57) (0.40	0.0				70.75	1 10		1 1 1		0.0	0.00 F		7 7	0100	1 0	P 4	5 9	- -	
717 55 77.0 53.0 73.0 53.0 73.0 53.0 73.0 53.0 73.0 53.0 73.0 53.0 73.0 53.0 73.0 53.0 73.0 53.0 73.0 53.0 73.0 53.0 73.0 53.0 73.0 53.0 73.0 53.0 73	2003	0 v V	0./0	0.00	13,100	00.09	00.00	10.03	2.00	1.03	6 50	0 00 0	1.00	10.1		7	0 044	ţ a	<u></u>	9 ç		
700 45 6.0 9.0 17.0 3.3.6 1.4 7.17 3.3.6 1.4 7.17 3.3.6 1.4 7.17 3.3.6 1.4 7.17 3.3.6 1.10 0.12 1.15 0.3.6 3.1 64 1.66 0.71 1.51 0.3 2.1 1.51 0.3 2.1 <td>2112</td> <td>0 v</td> <td>20.0</td> <td>7.00</td> <td>10/2/01</td> <td>44.00 00 10</td> <td>20.20</td> <td>77.00</td> <td>0.00 F 11</td> <td>00.1</td> <td>0.00 27 2</td> <td>00000</td> <td>1 70</td> <td>116</td> <td>0.00</td> <td>7 7</td> <td>9,044</td> <td>0 5</td> <td>2 0</td> <td>38</td> <td>ο α - τ</td> <td></td>	2112	0 v	20.0	7.00	10/2/01	44.00 00 10	20.20	77.00	0.00 F 11	00.1	0.00 27 2	00000	1 70	116	0.00	7 7	9,044	0 5	2 0	38	ο α - τ	
7.9 5. 0.0 5 0.0 <th0.0< th=""> <th0.0< th=""></th0.0<></th0.0<>	0020	0 v	0/.7	10.1	13,700	00.00 27 75	20.40	EU.11	0	1.40	21.0	2 20	0.10	1.10	0.04	7	0,002	1 0 7 -	8 8	57	0. 5	
38 5 113 114 12,070 33.25 2.335 7.337 4.36 1.32 0.33 2.5 0.33 2.5 0.33 2.5 0.33 2.5 0.33 2.5 0.33 2.5 0.33 2.5 0.33 0.33 2.5 0.33 <t< td=""><td>100</td><td>ç Ç</td><td>0.02</td><td>9.01</td><td>10,120</td><td>01.10</td><td>26.00</td><td>75 60</td><td>0.00</td><td>1 0 0</td><td>1 20</td><td></td><td>0.7 00 F</td><td></td><td>0.6</td><td>7 5</td><td>200,0</td><td>- 0</td><td>5 8</td><td>58</td><td></td><td></td></t<>	100	ç Ç	0.02	9.01	10,120	01.10	26.00	75 60	0.00	1 0 0	1 20		0.7 00 F		0.6	7 5	200,0	- 0	5 8	58		
35 11.3 14.4 12.30 0.03	10/2	ξ 2 ¢	4.30	3.11	10,100	04.44	10.00	20.02	20.00	00.1	1,00	2.02	00.1	0	2.0	7 7	15 100	2. 50			2 0	
all 10.40 10.20 12.952 37760 52.70 4.90 1.40 7.80 2.90 1.70 1.00 0.10 0.5 10812 7.4 92 48 1.3 werplant samples 3.40 12.50 476 2.90 0.20 1.00 0.10 0.5 10812 7.4 92 48 1.3 werplant samples 3.40 1.00 0.50 0.20 0.50 0.50 0.10 0.5 10812 7.4 92 48 1.3 werplant samples 5 3.40 1.40 %0 %10tals %0ratils %7 92 48 1.3 94 92 48 1.3 92 48 1.3 93 1.4 82 69 1.1 1.4 1.4 1.3 1.4 1.3 1.3 1.4 1.4 1.3 1.4 1.3 1.1 1.3 1.1 1.4 1.3 1.1 1.4 1.3 1.1 1.4 1.3 1.4 1.3 1.4 1.3 1.4 1.3 1.4 1.3	26130	f u	8 00 8	14.14	12,400	43.75	56.25	10.01	4'.00	р †	DC: -	2 30	0.94	1.32	0.13	7 5	16.407	54	146	154	- C	
ari ali 10.40 10.20 12.952 37.60 52.70 72.70 4.90 1.40 7.80 2.90 1.10 1.00 0.80 0.30 0.10 0.5 10812 7.4 92 48 1.3 werplant samples retraint sample no. type %Moisture %Ash Blu/lb %Volatile %FixedC %C %H %N %O %TotalS %OrganicS %PyriticS %SulfateS Ag Al As B Ba Be Sample % The %Moisture %Ash Blu/lb %Volatile %FixedC %C %H %N %O %TotalS %OrganicS %PyriticS %SulfateS Ag Al As B Ba Be Sample % TotalS	5	>	22.2			2	0.00					1	2	10.1	2	;	5	5	2	2) i	
rer plant samples Note that the sample no. type %Moisture %Ash Bu/lb %Volatile %FixedC %L %H %N %OrganicS %PyriticS %Sulfate S As B B Be 5300 BC-coal 12.45 10.96 %Volatile %FixedC %L %H %N %OrganicS %PyriticS %Sulfate S Ag As B Be Be 5300 FBC-coal 12.45 10.96 40.81 48.24 - - 3395 22.48 0.67 0.80 60.7 20.8 8.09 55 1 7 2 190 38 1 3305 5 1 3305 5 1 3305 5 1 3305 5 1 3305 6 1 320 130 3 1 300 3 1 3505 5 3 1 350 30 1 350 30	an²	all all	10.40 3.40	10.20 2.50	12,952 476	37.60 2.90	52.70 3.00	72.70 2.70	4.90 0.90	1.40 0.20	7.80 1.10	2.90 1.00	1.70 0.80	1.00 0.30	0.10 0.10	0.5	10,812 3,240	7.4 8.1	92 45	48 25	1.3 0.7	
Sample Sample<	ver plant	samples																				
Sample Sample Solution Molatile %FixedC %C %H %N %CoranicS %CorganicS %EvriticS %SulfateS Ag As B Be 5300 FBC-cool 12.45 10.96 40.81 48.24 - - 4.21 2.94 0.69 0.57 <0.2																						
5300 FBC-coal 12.45 10.96 40.81 48.24 - - 4.21 2.94 0.69 0.57 <0.2	o no.	Sample ype %	Moisture	%Ash	Btu/lb	%Volatile	%Fixed C	%C	H%	N%	0%	%Total S	%Organic S	%Pyritic S	%Sulfate S	Ag	A	As	В	Ba	Be	
	5300 FF 5301 C) 5302 C) 5302 C) 5302 FF 5304 FF 5304 FF 5306 C) 5306 FF 5306 FF 5309 C) 5311 PC	3C-coal 3C-coal 7C1-coal 7C2-coal C-coal 7C1-fly as 7C2-fly as 7C2-bottom 3C-bottom 7C2-bottom	12.45 12.26 13.286 13.72 13.72 13.7 13.7 13.7 12.94 13.7 12.94 13.7 12.94 13.7 12.94 13.7 12.94 13.7 12.94 13.7 12.94 13.7 12.94 13.7 12.94 13.7 12.94 13.7 12.95 13.7 12.95 13.7 12.95 13.7 12.95 13.7 12.72 13.72 11.7.72 11	10.96 10.63 10.64 10.64		40.81 39.05 50.19 38.77	48.24 50.31 50.60					4.21 3.95 4.02 4.00	2.94 2.48 2.52 2.52	0.69 0.67 0.79 0.79	0.57 0.85 0.69	002 002 003 003 003 003 003 003 003 003	9,897 8,045 8,045 8,203 8,150 33,925 70,179 72,296 10,076 80,076 69,067	2.6 2.6 2.2 2.2 5.1 1.8 6.1 3.4 1.1 3.4 1.1 2.1 2.1 3.4 3.3 3.3	195 198 198 198 198 429 429 429 100 27 27 359 385	39 35 37 37 37 37 387 357 334 357 357 357 357	1.7 1.7 1.7 1.7 1.7 1.7 1.7 1.7 1.7 1.7	

Table 3 Chemical composition of 35 marketed Illinois coals from preparation plants and of feed coals and combustion residues from four selected power plants. The values are in milligrams per kilogram

Q.	6 0 0 0 0 0 0 0 0 0 0 0 0 0 0 0 0 0 0 0	۵ 4
4		94
2		33 4 97 1,3
Mg		4'30 1'30
З	$\begin{array}{c} 0.11\\ 0.02\\ 0.03\\ 0.02\\ 0.03\\ 0.02\\ 0.03\\ 0.02\\ 0.03\\$	0.1
	<u>៰៵៵៷៰៵៹៵៸៰៸៷៰៸౼៹៹៰៰៰៰៰៰៰៸៰៰៰៸៰៰៰៸៰៰៰៸៰៰</u>	0.0
		± 0
La		51.(4 27.(
×	1,577 1,577 1,577 1,577 1,577 1,577 1,577 1,543 1,543 1,743 1,743 1,743 1,743 1,743 1,743 1,743 1,577 1,577 1,577 1,577 1,573 1,743 1,577 1,543 1,743 1,743 1,743 1,577 1,543 1,744 1,744 1,1444 1,1444 1,1444 1,1444 1,1444 1,1444 1,1444 1,1444 1,14444 1,14444 1,144444 1,144444 1,144444444	1,04 1 1,743
	222 222 222 222 222 222 222 222	5.6
βΪ		v Q
Ŧ	0.38 0.38 0.39 0.49 0.44 0.44 0.44 0.44 0.45 0.45 0.44 0.44	4.4 0.4
Ge	ကို ႏို ႏ ႏ ၈ က က ကို ရာ ကို	07 80
Ga	<pre>% 4 4 4 4 4 4 4 4 4 4 4 4 4 4 4 4 4 4 4</pre>	0.5 1.5
	000400000000000000000000000000000000000	00
Ъ.	$\begin{array}{c} 11\\ 12\\ 23\\ 23\\ 23\\ 23\\ 23\\ 23\\ 23\\ 23\\ 23\\ 2$	214,23 7,10
ш	252 253 253 253 253 253 253 253	ני א
		00
Ē		0.2
Ŋ	0.64 0.64 0.64 0.64 0.64 0.66 0.76 0.76 0.76 0.76 0.78 0.78 0.78 0.78 0.78 0.78 0.78 0.78 0.78 0.78 0.76 0.68 0.74 0.76 0.76 0.76 0.76 0.77 0.076 0.076 0.076 0.076 0.076 0.076 0.076 0.076 0.076 0.076 0.076 0.0777 0.0777 0.0777 0.0777 0.0777 0.0777 0.0777 0.0777 0.0777 0.0777 0.0777 0.077	22
5	2343 2343 2350 2350 2350 2350 2350 2350 2350 235	39.1 15.0
s	8878888890002000	0.1
^O		μΟ
ò	55525555555555555555555555555555555555	136
8	2002 4 5 2 2 5 5 4 4 5 2 5 5 4 4 5 5 5 5 5	3.5
σ	100 100 1150 1150 1150 1150 1150 1150 1	, ,
8	6.1 6.1 6.1 6.1 6.1 6.1 6.1 6.1	55 9.4
8	0.25 0.25).47).55
		37
S	$\begin{array}{cccccccccccccccccccccccccccccccccccc$	39,2, 356,9(
ы	$\begin{array}{c} 0.5\\ 0.5\\ 0.5\\ 0.5\\ 0.5\\ 0.5\\ 0.5\\ 0.5\\$	<1 0.6
no.	777 777 777 777 777 778 779 880 779 880 880 779 880 880 779 880 880 779 880 779 779 777 779 880 779 880 779 777 779 800 800 779 800 770 800 770 800 770 800 770 800 700 800 700 800 700 800 700 800 700 800 700 800 700 800 700 800 700 800 700 800 700 800 700 800 700 800 700 800 700 800 700 800 700 800 700 800 700 800 8	311 312 #inu ood
Lab		C35

0		-																				
Lab no.	Na	Ĩ	٩	Pb	Вb	Sb	Sc	Se	Si	Sm	Sn	Sr	Та	- qu	L L	H I	n	>	M	ЧY	Zn	Zr
C30773	1 039	ŧ	44	ų	1	0 0	1 0	1	20.988	0.65	<u>د</u> ہ	28 (1 0	. 12	5	39 <1	-	7 16	0.35	0.29	32 9	00
C3077A	148	ų L	87 2	124	. LC	0	0	0	11 312	0	i rć	14	0107	1.25 (2 780			. α	000	0.61	20.20	ç Ç
C30777	445	2 1	44	36.0	, t	- - i -		- - 1 (c	27 20g	 0.45) {{	- 92	11	24 C	,		-, -		100	000	447	2 6
C32778	1 187	31	87	13.5	2 6	00	2.6	, LC	23,886	0.97) r		12	15		20	: c	- 6	0.35	5 0 43	105	1 %
C32782	1 113	5 ~	87	2. 42	i 8	10	9 -	6	27,438	0.49	<5.5	25.0	1.0	. 200	2	39	; -		0.35	5 0.26	42.2	3 8
C32783	1,484		5 4	<5.5	12	0.1	2.0	1.3	20,520	0.75	5	58	.11	0.13	.3	39 <1	. <u> </u>	112	7 0.3	0.44	54.1	22
C32785	1,261	17.5	131	<5.3	11	0.4	2.0	3.9	23,091	0.73	\$	30	0.12 0	0.13	.3 5	39 <1	÷.	8	9 0.4	0.37	59.9	22
C32797	1,113	16	87	8.5	11	0.6	2.2	1.4	23,232	0.72	\$	44	0.11 0	0.13	.5	99 <1	÷.	9 27	7 0.45	5 0.39	105	25
C32814	74	12	<44>	23.0	5.1	1.9	1.1	1.3	9,956	-	ŝ	17 0	0.05 0).29 (.61 2	40 <1	8.	0 16	3 0.25	5 0.76	67.9	÷
C32779	1,039	=	44	7.0	10	0.4	2.0	2.3	22,203	0.8	ŝ	27 0	0.1	0.13	.3 5	39 <1	ю.	1 4	1 0.55	5 0.41	77.9	22
C32794	371	14	4	14.5	10	0.3	2.1	1.9	23,559	0.8	ŝ	20	0.1	.14	.4 5	99 <1	¢.	0 23	3 0.3	0.43	120	24
C32798	1,039	18	87	12.0	13	0.3	2.5	3.2	28,701	0.93	ŝ	24 (0.12 0	0.16	.7 6	59 <1	c,i	7 46	5 0.45	5 0.47	78	28
C32800	964	15.5	87	8.0	14	0.4	2.4	2.8	26,457	0.89	ιŞ	22	0.12 0	.13	.6	59 <1	¢.	90 30	0.4	0.42	68.5	26
C32813	890	24	305	12.5	19	0.7	3.1	5.4	35,572	1.2	\$	27 (0.14 0	0.2	1.1	39 <1	ю	7 65	5 0.5	0.65	141	34
C32815	371	10	4	12.3	11	0.6	2.4	2.1	25,849	0.91	\$	20	0.13 0	0.13	.7 6	59 <1	÷.	9 27	7 0.25	5 0.45	101	24
C32784	890	15	4	39.5	13	1.1	2.6	1.4	18,510	1.2	ŝ	30	0.1	0.16	.4 5	99 <1	÷.	0 18	3 0.4	0.47	28.8	31
C32795	1,039	18	4	16.0	8.8	1.1	1.9	1.1	15,005	0.82	\$	27 0	0.1	. 60.0	.2	80 <1	0	6 15	5 0.25	5 0.33	39.9	20
C32796	1,261	23.5	87	31.3	24	1.0	3.8	2.0	44,079	1.8	ŝ	30	0.21 0	0.23	1,1	99 <1	÷.	0	9 0.3	0.74	318	51
C32799	519	14.5	436	63.7	15	3.6	2.8	1.3	26,363	1.4	ŝ	156 0	0.12 0	. 19	.6	59 <1	0	7 28	3 0.75	5 0.47	121	28
C32801	816	14	44	22.0	1	0.6	2.4	1.5	20,146	0.97	Ŝ	22	0.11 0	0.14 1	.5 6	59 <1	÷.	0 2	1 0.75	5 0.42	34.5	27
C32802	593	7	44	7.0	11	0.3	2.2	1.5	20.801	0.78	ιŜ	29	111 0	.13	.4 5	39 <1	-	33	0.35	5 0.38	32.1	24
C32803	816	13	44	12.5	: =	0.4	2.6	15	20,661	0.93	ŝ	27	1	15	9	12		5	1 0 4	0.43	25.4	22
C32661	519	13.5	44	19.0	10	0.5	94	200	18 791	0.85	592	i 6		. 13		30	- 2	Э С	5 0 4	0.47	68.2	22
C32664	245	2. 	5 5	30.5	14	0.0 7	i (*) (r - 1	23 746		, r , r	15		218	о г с		: ~	3 8 0 0	100	0 53	38.7	i g
CRORES	207	: -	54	18.5	r (*	0.0 0	- r	 	22 764	0 98	20.07	1 %		14	, u		iu	16		0.45	23.5	200
C32771	445	: C	44	215	2 4	0.3	0.0	4	30.570	1.3	; .c		14 (16	0.0	12 00	i a		0.5	0.55	104	3 8
C32776	668	2 0	44	10.0	2 =	0.6	s i c	1 7	20.053	0.79) -C	14	11	19	. 4 . 6	30	i	1 00	0.45	5 0 44	6	3 K
C32662	593	17	175	22.5	: =	1.0	2.5	1.3	16.173	1.7	ιų	82	1.0	0.24	. 6.	39		9 21	7 0.4	0.61	28.4	22
C32663	252	13	144	96.0	9.7	1.4	2.4	1.5	21,549	0.88 <	10	46 (0.1	0.13	.4 6	59 1	0	6	3 0.45	5 0.44	61.8	39
C32772	371	13	87	27.0	10	0.7	2.4	1.3	22,343	0.95	۲ŷ	43 (0.13 0	.14	.3	99 <1	v	, ç	9 0.45	0.39	39.8	32
C32775	148	17.5	44	51.5	11	1.2	2.3	3.6	16,360	0.9	\$ ℃	44	0.1 0	0.14 1	.3 4	80 <1	Ö	8	9 0.35	5 0.43	109	24
C32780	445	7.5	4	11.7	10	1.4	2.0	2.3	20,941	0.7	ŝ	19 0	0.11 0	1.0	.2	99 <1	c,	0	1 0.5	0.33	35.7	27
C32781	223	1	44	46.0	11	1.4	1.7	2.5	22,343	0.74	\$2	28	0.1	1.1	·2 5	39 <1	i2	0-9-	1 0.35	5 0.34	62.4	27
C32793	1,039	22	175	35.5	18	1.2	2.9	1.1	33,655	1.3	ŝ	40 0	0.16 0	. 17	8.8	39 <1	0	8	2 0.5	0.56	91.3	49
C35564	1,335	25	87	35.4	17.6	2.1	3.7	2.2	31,084	1.3	ŝ	35 (0.2	0.17	.4 8	99 <1	O	9 26	0.1	0.62	252	12
Mean	716	14.4	87	25.8	12.0	0.9	2.4	1.9	23.314	1.0	2.6	33	0.1	0.2	.5	11 0	5 2.	3	1 0.4	0.5	89	27
SD	397	5.5	83	25.9	3.7	0.7	0.6	0.9	6,605	0.3	0.5	24 (0.0	0.0	.4 1	53 -	,	9 16	s 0.1	0.1	87	8
C35300	1.113	18	131	4.3	19	0.4	2.5	5.5	26,597	1.0	\$	34	0.1	0.2	8.	39 <1	÷.	9 41	0.3	0.5	22	10
C35301	1,039	8	44	3.3	14	0.7	1.8	2.7	24,961	0.7	\$	25 (0.1 0	.1	·2	80 <1	2	0 32	4 <0.3	0.5	34	б
C35302	964	17	4	3.2	13	0.8	1.9	2.0	25,475	0.6	ŝ	25 (0.1 0	1.1	.3 5	39 <1	,	8 37	7 <0.5	0.3	39	6
C35303	1,039	1	4	3.2	12	0.6	1.9	2.9	25,475	0.7	ŝ	26 (.1 0	.1	.4	80 <1	0	3 41	0.4	0.5	34	б
C35304	3,858	99	567	14.1	54	1.3	7.8	14.4	96,572	3.8	Ŝ	300	0.4 0).5 5	6.0 1,9	78 2	7.	1 69	9 3.8	1.7	193	10
C35305	14,615	173	742	89.7	132	13.6	19.4	43.3	217,404	6.3	1	264	ς.	.0	1.6 6,1	14 10	21.	7 420	4.0	3.9	962	111
C35306	16,247	308	8/3	170	178	49.9	23.1	24.3	189 451	8.5	10	371	5	.12	.8 6,1	14 4	<u>1</u> 00	1 560	5.55	6 4.35	1,286	131
C35307	11,425	1	524	48.9	104 i	7.9	18.0	7.4	243,347	6.7	ں م	264		0.8	2.6 5,4 2	55 4	16.	8 428	3.4	00 I 00 I	306	105
0225308	1,113	14	480	8.4 8.0	15	0.8	2.4	⊽ [.]	61,32/	1./	. 0	506	2.2	2.2	6. 9	59 . <1	4	÷ ۲	3 0.9	0. /	2	ND ND
C35309	6,380	98	524	€0. 1	99 F		16.3		244,235	1.0	18	255	0.0	. I . I	.3 4,0	16 2	Ξ.	1 197	9.V	а. 1 0	97	4
015350	10,/0	20	349	<4.8	5 5	0.1	10.1		246,852	5.5	14	152			.1 4,1	36		247	D. I	מ מ		5 2
C35311	5,935	ςΩ γ	349	10.2	88	N C	14.4 1	N L	217,638	4.8 •	26	194 0	8.0	0.6	.4 3,0 2	96	N C	53	0. I V	7.7	55 20	00
215050	919	9.4	292	5.03	=	0.2	G. [<0.0>	21,409	-	₽	929	1.0	2.0	0.	[V]	N	2	t <1./	C.U	89	
¹ Regions	H Herrir	Coal: S	Spring	ufield Coal																		
² For these	soulev e	ndinater	4 with "	" sinn on	f juit of t	illen ed	OC MOL	i head ir	n tha ctatic	tical co	montati	h of and	otormin	o the me	an and ct	ondard c		soulen				
	ה עמותכס	ווחורמובי		signi, un	6-11411 OL	ווב עמות	es wei	nasn :	ו וווב אמווא	llual co	IIpuiain	טווא ויט מ	erennin	e li le li le	สโเสเนอเ	aliuaiu	leviation	Values.				

Table 3. Continued Chemical composition of 35 marketed Illinois coals from preparation plants and of feed coals and combustion residues from four selected power plants. The values are in milligrams per kilogram (unless indicated otherwise) on a moisture-free basis.

Mineral-element pair		r ²	Mineral-element pair		۲²
Illite/smectite	Cs Mg Ti Ta Sc	0.62 0.53 0.52 0.49 0.48	Illite + Illite/smectite	Mg F Ti Cs Rb	0.66 0.63 0.63 0.60 0.59
	Rb Hf Ga Ba	0.47 0.46 0.42 0.41 0.36		Th Ta Hf Ba Ga Ce La	0.56 0.55 0.49 0.47 0.45 0.37 0.34
Kaolinite	Hf Sc Ba Ti	0.58 0.55 0.53 0.53	Chlorite	Ta Ti Hf	0.34 0.33 0.31
	Li Ta Th Ce	0.51 0.51 0.46 0.38	Quartz	Mg Ti Ta	0.53 0.32 0.31
	La Cs	0.36 0.32	Calcite	Mn Zn	0.94 0.54
Marcasite	B Br	0.48 0.37			

Table 4 Mineral-element correlations in marketed Illinois coals at $\alpha = 0.001$ levels of significance, which correspond to $r^2 > 0.30$ for the number of samples used.

to a lesser extent, to mullite. Pyrite and marcasite were converted to hematite, magnetite, and amorphous Fe oxides. Some of the calcite reacted with sulfur released from pyrite, marcasite, and organic matter and formed anhydrite, which later reacted with moisture and partially hydrated to gypsum. The rest of the calcite was converted to CaO (lime) and Ca(OH)₂ (portlandite); some of these minerals subsequently changed back to calcite through reaction with atmospheric CO₂. At the FBC plant, large amounts of lime and portlandite were generated (mostly in the bottom ash) because of the limestone addition; subsequent reaction with atmospheric CO₂ most likely took place only on the exterior surfaces of these relatively coarse Ca minerals from the FBC ashes. The mineral conversion reactions apparently varied from one type of combustion unit to another, which resulted in distinct differences in the mineralogical as well as chemical compositions of the CCRs from the four coal-fired power plants. These differences have important implications for the effect of plant type on the HAPs emissions and on the suitability of the CCRs for use in cement, ceramic, and other commercial applications.

Emission of HAPs from Power Plants

If a HAP element is entirely retained in CCR, its emission into the atmosphere at coal-fired power plants is prevented. However, a portion of each HAP generally is mobilized and becomes airborne. The mobilized portions of the HAPs may be partially adsorbed on surfaces of hardware on the cool side (between the boiler and the stack) of a power plant and partially emitted into the atmosphere. Thus, the mobility values of HAPs would overestimate their atmospheric emissions, except in the case of highly volatile HAPs (F, Hg, Se). Therefore, it is difficult to obtain mass balance for trace elements from power plant samples, and the mobility values reported here should be used cautiously.



Figure 7 Relationships of $r^2 > 0.5$ between elements and mixed-layered illite/ smectite plus illite in marketed Illinois coals.



Figure 8 Relationships of $r^2 > 0.5$ between elements and kaolinite in marketed Illinois coals.

Figure 9 Relationship between quartz and Mg in marketed Illinois coals.



The mobilities of 15 HAP elements from the four power plants were determined by using mass balance calculations (table 5). The mass balance of an element was calculated by comparing its mass in the feed (w_F) and in the CCRs (w_{CCRs}):

Mass balance (%) = $(w_{CCBs} / w_{F}) \times 100$.

The calculations of w_{cCRs} and w_{F} took into consideration the mass ratios of fly ash to bottom ash as well as the measured concentrations of elements and high-temperature ash in the samples. The fly ash to bottom ash ratios were 80/20 for the FBC plant, 25/75 for the CYC plants, and 75/25 for the PC plant. The feed for the FBC plant included not only coal but also the limestone added to the bed at a ratio of 75% coal to 25% limestone.

The mass balances of the 15 elements were normalized to that of AI (by dividing them by AI mass balance and multiplying the result by 100) to minimize the effect of sampling and analytical errors on the mass balance computations. Aluminum was used for normalization because it is a refractory element with a relatively large concentration in coal and is expected to be retained almost completely in the combustion ashes; no more than 5% (most of the time <1%) of AI was expected to escape the particulate collection systems with ultra-fine, airborne fly ash particles. Aluminum mass balances of approximately 100% for the CYC and PC units (Table 5) indicated that the mass balance calculations performed for these two types of plants were highly accurate. The AI mass balance for the FBC plant (76%) was somewhat small but still reasonable. The AI mass balance deficiency for the FBC plant could have resulted from variations in the composition of the feeds that would make it difficult to obtain a truly representative sample during a few hours of operation. The limestone and the coal used in the FBC furnace were blends of products from many different quarries and mines in Illinois. Therefore, future studies should consider collecting at least several



Figure 10 Calcite-Mn and calcite-Zn relationships in marketed Illinois coals.







Figure 12 Mineralogical composition (from table 2) of CCRs from three types of plants (FBC, PC, CYC) in Illinois. The cyclone plant values were the averages of two plants. Key: Mul, mullite; Calc, calcite; Qtz, quartz; Magn, magnetite; Hemt, hematite; Anhyd, anhydrite; Portl, portlandite; Gyps, gypsum; Glass, amorphous phase.

Table 5 Concentrations of AI and 15 elements of environmental concern in coal, CCRs, and limestone samples from the four power plants and related combustion mobility and relative enrichment (RE) values. The concentration values are in milligrams per kilogram unless indicated otherwise. All values are on a dry weight basis.

ltem	Plant and sample type	AI (%)	As	Be	Cd	°	ç	щ	Hg	Mn	ïŻ	٩	Pb	Sb	Se	Ч	
C35300 ¹ C35304 C35308 C35308 C35312 Mass balt Mobility (* RE for fly RE for bo	FBC-coal FBC-fly ash FBC-bottom ash FBC-limestone ance (%) %) ash ttom ash	1.00 3.39 1.09 0.63 24 0.9 0.3	1.8 6.1 3.3 81 19 0.6 0.6	1.7 6.0 96 0.9 0.9	1.00 2.48 0.63 0.55 0.55 27 0.6 0.6	4.6 12.3 4.3 3.5 3.5 76 24 0.6 0.6	25 75 35 96 0.8 0.8	137 53 48 5 15 85 0.1 0.1	0.10 0.25 0.01 <0.01 82 0.7 0.7	77 620 310 1,394 42 58 0.3 0.2	18 66 14 9.4 0 1.0 0.2	131 567 480 262 104 0 0.8	4.34 14.1 4.8 5.03 84 16 0.7 0.2	0.4 1.3 0.8 0.2 0 0.8 1.5	5.5 14.4 <1.4 <1.6 <0.5 87 13 0.8 0.0	1.8 5.6 1.9 94 0.8 0.8 0.3	1.9 7.1 2.8 94 0.8 0.8
C35301 C35305 C35309 Mass bala Mobility ('	CYC1-coal CYC1-fly ash CYC1-bottom ash ance (%) 1	0.80 7.02 8.01 102 0	2.6 50.6 1.2 54 46	4.0 14.0 5.5 20 80	0.19 11.5 0.48 170 0	4.0 24.2 54 54 46	17 227 112 85 15	63 15 11 89	0.08 0.73 0.02 25 75	77 310 542 65 35	8 173 70 123 0	44 742 524 135 0	3.34 89.7 2.65 75 25	0.7 13.6 1.1 62 38	2.7 43.3 2.1 47 53	1.2 14.6 11.3 104 0	2.0 21.7 11.1 71 29
RE for fly RE for bo	ash ttom ash	0.9	2.0	0.4	6.2 0.3	0.6	1.4 0.7	0.0	1.0 0.0	0.4 0.7	2.3 0.9	1.8 1 i2	2.8 0.1	2.0 0.2	1.7 0.1	1.3 1.0	1.1 0.6
C35302 C35306 C35310 Mass bal Mobility (RE for fly RE for bo	CYC2-coal CYC2-fly ash CYC2-bottom ash ance (%) %) ash ttom ash	0.82 7.23 7.97 107 0 0.9	2.4 2.4 0.05 160 6.5 0.0	1.9 15.0 62 38 0.8 0.6	0.20 16.2 0.49 232 8.7 2.6	3.5 3.5 45.8 17.2 73 27 1.4 0.5	18 208 125 85 15 1.2 0.7	61 200 5 91 0.4	0.08 0.41 0.005 14 76 0.5 0.0	77 310 542 66 34 0.4	17 262 62 69 31 1.7 0.4	44 873 349 115 0 2.1 0.8	3.22 170 2.4 144 0 5.7 0.1	0.8 48 1.0 167 6.4 0.1	2.0 22.2 1.8 36 64 0.1 0.1	1.3 15.6 11.1 99 1 1.3 0.9	1.08 19.2 13.2 86 1.1 1.1 0.8
C35303 C353307 C35311 Mass bal Mobility (RE for fly RE for bo	PC-coal PC-fly ash PC-bottom ash ance (%) %) ash ttom ash	0.82 8.47 6.91 105 1.1 1.1	2.2 34.0 3.0 121 0 1.6 0.1	9.0 9.0 91 9 1.0	0.42 4.89 0.47 92 8 0.1	3.5 20.2 22.2 60 40 0.6 0.6	175 175 99 1 1.1 0.9	63 67 5 8 92 0.1	0.08 0.10 >.01 90 0.1 0.0	77 468 62 38 0.6	11 77 85 73 73 27 0.7 0.7	44 524 262 106 0 1.3 0.6	3.2 48.9 10.2 124 1.6 0.3	0.6 7.9 2.2 110 0 1.4	2.9 7.4 2.2 21 79 0.3 0.3	1.4 12.6 10.4 87 13 1.0 0.8	2.3 16.8 12.5 0.8 0.8 0.8

¹ Lab number.

Table 6	Mobilities	of HAPs	during	combustion.
---------	------------	---------	--------	-------------

	Mobility	(%)	
Plant type	Low (<25%)	Moderate (25-50%)	High (>50%)
FBC	As, Be, Co, Cr, Hg, Ni, P, Pb, Sb, Se, Th, U	Cd	F, Mn
CYC ¹	As, Cd², Cr, Ni, P², Pb, Sb², Th, U	Be, Co, Mn	F, Hg, Se
PC	As², Be, Cd, Cr, P, Pb, Sb², Th	Co, Mn, Ni, U	F, Hg, Se

¹ Average of two CYC plants. The two plants varied significantly with respect to the mobilities of some elements (see table 5).

² Negative mobilities resulting from excess mass balance suggest the possibility of contamination of combustion ashes by erosion or corrosion of hardware (see text for more on the definition of mobility).

sets of samples over a period of several months of operation from the FBC plant, and the average mass balance data on these samples should be used to smooth out the variance.

The mobility of HAP elements from the four plants was calculated as follows:

Mobility (%) = 100 - mass balance.

According to this mobility relationship, if the amount of an element recovered from the CCRs accounted for 100% of the amount in the feed, then the element was assumed to be immobile. If the mass balance of an element was less than 100%, then the difference was considered to indicate the percentage of the element that entered and remained in the gas phase or condensed on the ultra-fine, airborne fly ash particles or on the power plant hardware.

For convenience, the mobility values were divided into low, moderate, and high categories (table 6). Negative mobility values caused by the excess mass balance (101–167%) in some cases (table 5) probably resulted mainly from contamination of the CCRs by the erosion or corrosion of hardware in the combustion process. In particular, the excess mass balances observed for Cd for the CYC1 unit and for As, Cd, Pb, and Sb for the CYC2 unit (table 5) were too large to explain as sample variability or analytical error. Analysis of the CYC2 fly ash was repeated three times, and no significant variation was found between the three analyses. Although the negative mobility values were arbitrarily assumed to be in the low-mobility category, we recommend that periodic sampling of the CYC2 plant and analysis of the samples be carried out in the future to better understand the combustion behavior of As, Cd, Pb, and Sb at this unit.

For the FBC plant, only the mobilities of F (85%) and Mn (58%) were large (table 6). The mobilities at the FBC plant were surprisingly low for the normally volatile elements Hg (18%) and Se (13%). Overall, the HAPs mobilities were lower at the FBC plant than at the other plants (table 5). Apparently, a relatively low combustion temperature (~900°C) or the chemical environment created by the addition of limestone resulted in generally lower HAPs mobilities. The FBC fly and bottom ashes naturally contained large amounts of anhydrite and lime-portlandite (table 3). Several investigators (Clarke and Sloss 1992; Meij 1993, 1994a; Gullet and Ragnunathan 1994; Bool and Helble 1995; Querol et al. 1995; DeVito and Bhagwat 1997) reported that Ca-rich materials such as lime and limestone can capture substantial amounts of As, Hg, Sb, and Se during combustion. Suarez-Fernandez et al. (1996), however, did not find any major differences between the combustion behavior of trace elements in a laboratory-scale FBC unit with or without the addition of limestone. This result suggests that low temperatures may influence the mobility of certain elements more than does limestone addition.

For the CYC and PC plants, mobilities of the volatile elements F (89-92%), Hg (75-90%), and Se (53-79%) were high (table 6), as expected. The mobility values of some elements for the CYC and PC plants were inconsistent with the volatility data summarized in figure 7 of Clarke and Sloss (1992). According to Clarke and Sloss (1992), As, Cd, Pb, and Sb are moderately volatile. However, our data indicated that all four elements had low mobilities at the PC plant, and Cd and Pb had low mobilities at the two CYC plants. Also, the Mn mobility was supposed to be low, which is inconsistent with the moderate Mn mobility calculated for the CYC and PC plants.

Although Mn is not a volatile element at normal coal combustion temperatures, its mobility from the FBC plant was surprisingly large. Querol et al. (1995) reported that Mn has an affinity for Fe oxide in the CCRs. The Mn content of the feed coal from the CYC and PC plants was less than the Mn content of the feed (a blend of coal and limestone) from the FBC plant (table 5). Furthermore, CCRs from the CYC and PC plants contained more Fe oxide minerals (magnetite + hematite) than did the CCR from the FBC plant (table 3). These differences could be the reason Mn mobility was considerably greater at the FBC plant than at the CYC and PC plants.

Studies reviewed by Clarke and Sloss (1992) and Davidson and Clarke (1996) indicated that among the 15 HAPs investigated, substantial portions of only F, Hg, and Se were mobilized in the gas phase and emitted into the atmosphere during coal combustion. The emission of other elements generally resulted from their enrichment in the submicron-size, fly ash particles that passed through the particulate control systems.

Partitioning of HAPs between Fly Ash and Bottom Ash

The atmospheric emission of elements from coal-fired power plants is directly related to the partitioning of elements among CCRs. For example, the enrichment of an element in the bottom ash would reduce its atmospheric emission. However, if an element is preferentially enriched in submicron-size, airborne, fly ash particles, its emission would be enhanced. The partitioning of elements in CCRs affects not only their atmospheric emissions but also the commercial value of the CCRs. To compare the distribution of HAP elements in fly and bottom ashes relative to their concentrations in coal, a relative enrichment factor (RE) was calculated for each HAP using the formula of Meij (1992):

$$RE = (C_{CCR}/C_F) \times (%ash_F/100).$$

where C_{CCR} and C_{F} are the concentrations of an element in the CCR (fly ash or bottom ash) and feed, respectively, and %ash_F is the percentage of ash in the feed. The feed refers to coal or, in the case of the FBC unit, a mixture of coal and limestone. Elements that are neither enriched nor depleted in the CCR should ideally have relative enrichment values of 1. Elements with relative enrichment values of significantly greater or less than 1 are enriched or depleted, respectively, in the CCR. Based on the literature (Meij 1992) and for convenience, the relative enrichment values in this study were divided into "normal," "enriched," and "depleted" categories (table 7).

In most cases, the relative enrichment factor of the HAPs was greater for the fly ash than for the bottom ash samples from the four plants (table 5). The only reverse cases were Sb at the FBC plant, Mn at the CYC plants, and Co and Ni at the PC plant. The greater relative enrichment values for fly ash than for bottom ash probably resulted from the combination of two factors: (1) large portions of most HAP elements were volatilized during combustion and then, upon cooling, condensed on the fly ash particles, and (2) some elements were enriched in finely disseminated coal minerals that ended up in the fly ash in greater proportions than they did in the coarse coal minerals.

Some elements with large relative enrichment values could be recovered from the respective CCR and sold. For example, Se was highly enriched in the CYC1 fly ash (RE = 1.7). The average concentrations of Se in the earth's crust, shale, and soils are 0.05, 0.5, and 0.4 mg/kg, respectively (Clarke and Sloss 1992). As a result, the Se concentration in the CYC1 fly ash was 866, 87, and 108 times greater than the earth's crust, shale, and soil averages, respectively. Therefore, the

Table 7 Relative distribution of HAPs among combustion ashes.

Plant and		Distribution	
ash type	Normal (RE = 0.7-1.3)	Depleted (RE < 0.7)	Enriched (RE > 1.3)
FBC-fly FBC-bottom	Be, Cr, Hg, Ni, P, Pb, Sb, Se, Th, U P	As, Cd, Co, F, Mn As, Be, Cd, Co, Cr, F, Hg, Mn, Ni, Pb, Se, Th, U	none Sb
CYC ¹ -fly CYC-bottom	Be, Co, Hg, U Cr, Mn, Ni, P, Th, U	F, Mn As, Be, Co, F, Hg, Pb, Sb, Se	As, Cr, Cd, Ni, P, Pb, Sb, Se, Th Cd
PC-fly PC-bottom	Be, Cd, Cr, Ni, P, Th, U Be, Co, Cr, Ni, Th	Co, F, Hg, Mn, Se As, Cd, F, Hg, Mn, P, Pb, Sb, Se, U	As, Pb, Sb none

¹Average of two CYC plants.

CYC1 fly ash can be considered to be a potential source for commercial Se production. Rose et al. (1979) reported that Se is produced industrially as a by-product of Cu refining; for an Arizona Cu ore deposit, for example, Se concentration was ~8 mg/kg. This value is much greater than the crustal average of 0.05 mg/kg Se, but similar in magnitude to the range of 7.4 to 43.3 mg/kg for Se found in the fly ashes from the three Illinois plants (table 3).

In the past, relative enrichment values in the combustion products were used to assign the inorganic elements in coal to differing volatility classes (Clarke and Sloss 1992, Meij 1992, Davidson and Clarke 1996). However, such a task is often complicated by substantial variations in the combustion behavior of elements depending on the characteristics of coal, the type and operating conditions of power plants, and sampling and analytical errors. For example, some elements were enriched in the fly ashes from CYC or PC plants but not in the fly ash from the FBC plant (table 7). The only obvious trend common to all four power plants was that most HAPs were depleted or normally distributed in the bottom ashes (table 7). Although depletion of Al, a non-HAP conservative element, is not expected in the CCRs, its depletion in the FBC bottom ash (RE = 0.3; table 5) is not surprising. The likely reason for the observed AI depletion is that the abundant and AI-rich clay minerals from the FBC feed coal ended up in the fly ash in greater proportions than the Aldeficient calcite that was the principal component of the limestone. This study did not examine the enrichment of the trace elements in particle-size fractions of the fly ash samples. Previous studies (Tumati and DeVito 1991, 1993; Dale et al. 1992; DeVito and Jackson 1994; Helble 1994; Meij 1994b; Querol et al. 1995; Cereda et al. 1995; Suarez-Fernandez et al. 1996) indicate that there is generally a negative correlation between ash particle size and the concentrations of As, Co, Cr, Hg, Mn, Ni, Sb, and Se.

Temporal Variations in the Characteristics of Marketed Coals

Marketed coals from four of the Illinois preparation plants were sampled twice with a 4-year interval between the two sampling times. The mineralogical and chemical characteristics of these asshipped coals changed somewhat over the 4 years (table 8); some of these changes could partly be the result of the simple sample variation that can occur any time. There was no apparent systematic change that was common to all four coals. For example, the ash content increased for two of the coals and decreased for the other two. The temporal change in coal quality for some individual mines is not surprising because over time the production in a mine moves to different reserve blocks, commonly with somewhat differing coal qualities. And, in some cases, coal cleaning efficiency is altered over time to meet the specifications of coal buyers. For example, if the customer of a mine changes from a PC boiler without a scrubber to a PC boiler with a scrubber, a FBC unit, or gasification plant, the run-of-mine coal does not have to be cleaned as much for sulfur (pyrite) removal. The average coal quality for the entire basin, however, would not change significantly over a 4-year period. The coal C35300 from the FBC unit (Tables 3 and 5) was a blend of production from 17 different and mostly high-sulfur coal mines; thus, this sample probably represented a typical marketed coal from high-sulfur Illinois coal mines.

 Table 8
 Temporal variations in the mineralogical and chemical composition of marketed coals from four selected Illinois preparation plants.

 Sample 1
 from each plant was collected in 1992 (Demir et al. 1994), and sample 2 from each plant was collected in 1996. All values are on a dry weight basis, except equilibrium moisture.

	Pla	nt A	Pla	int B	Pla	nt C	Plar	nt D
Composition	Sample 1	Sample 2	Sample 1	Sample 2	Sample 1	Sample 2	Sample 1	Sample 2
	(C32773)	(C35559)	(C32774)	(C35562)	(C32782)	(C35303) ¹	(C32814)	(C35561)
Mineralogical (wt%) Illite/smectite Illite Kaolinite Chlorite Quartz Pyrite Marcasite Calcite K-feldspar Plagioclase	1.67 0.79 1.50 0.07 2.24 1.42 0.24 0.24 0.71 0.05 0.07	1.94 1.72 2.85 0.14 2.04 1.52 0.55 0.91 <0.01 <0.01	0.43 0.53 0.93 0.05 1.26 2.35 0.26 0.27 <0.01 0.04	3.21 1.96 1.60 0.20 0.79 2.87 0.56 0.21 <0.01 0.11	0.61 1.10 1.69 0.06 3.62 2.00 0.06 1.27 <0.01 0.11	0.35 2.32 3.47 0.19 1.61 1.27 0.07 1.00 <0.01 0.02	0.69 0.47 0.66 0.02 1.19 1.71 0.01 0.87 <0.01 0.07	0.63 0.87 1.33 0.05 0.35 1.31 0.60 0.45 <0.01 <0.01
Ash	8.7	11.2	7.1	10.7	11.6	10.6	6.0	4.9
Volatile M	41.2	40.9	40.9	41.1	40.0	38.8	45.7	42.6
Fixed C	50.1	48.0	52.1	48.2	49.4	50.6	48.3	52.6
Pyritic S	0.97	1.20	1.46	1.87	1.25	0.79	1.09	1.11
Organic S	3.03	3.27	1.56	1.89	2.45	2.52	2.41	1.43
Sulfate S	0.14	0.19	0.56	0.19	0.20	0.69	0.18	0.51
Eq. moisture	13.9	12.1	13.4	12.2	14.3	14.7	11.3	13.9
Al	0.90	1.02	0.53	1.21	0.86	0.82	0.38	0.34
Si	2.10	2.53	1.13	1.94	2.74	2.55	1.00	0.67
Fe	1.19	1.41	2.11	2.25	1.50	1.55	1.26	1.67
Ca	0.29	0.36	0.11	0.09	0.51	0.40	0.35	0.18
K	0.16	0.20	0.091	0.27	0.16	0.15	0.075	0.06
Na	0.10	0.10	0.015	0.022	0.11	0.10	0.007	0.007
Chemical (mg/kg) As B Br Ba Cd Cc Cc Cc Cc Cc Cc Cc Cc Cu Eu F Ga Ge Hf Hg La Li Lu Mg Mn Mo Ni P Rb Pb Sb Sc Ss Sm Sr Ta Tb Tt Tt U V W Yb Zn	$\begin{array}{c} 1.3\\ 135\\ 6.5\\ 41\\ 1.0\\ <0.3\\ 6.3\\ 2.3\\ 12\\ 0.88\\ 7.4\\ 0.16\\ 90\\ 3.2\\ <5\\ 0.36\\ 0.13\\ 3.5\\ 3.9\\ 0.1\\ 482\\ 39\\ <6\\ 11\\ 44\\ 11\\ <6\\ 0.2\\ 1.9\\ 0.65\\ <5\\ 28\\ 0.1\\ 0.12\\ 1.2\\ <1\\ 539\\ 1.7\\ 16\\ 0.35\\ 0.29\\ 33\\ 20\end{array}$	1.6 163 8.3 44 <1 1.0 8.0 3.4 3.4 1.1 8.2 0.21 141 2.9 <5 0.5 0.07 4.3 4.3 0.09 603 42 6.5 11 87 14 5.4 0.3 2.0 2.7 1.0 <5 33 0.11 1.4 <1 599 1.3 36 0.50 0.40 99 11	$\begin{array}{c} 20\\ 116\\ 3.9\\ 17\\ 4.0\\ <0.3\\ 4.1\\ 3.1\\ 5.8\\ 0.75\\ 13.6\\ 0.32\\ 68\\ 4.0\\ 27\\ 0.32\\ 0.22\\ 2.2\\ 5.8\\ 0.11\\ 241\\ 18\\ <5\\ 15\\ 87\\ 5\\ 102\\ 1.2\\ 1.9\\ 1.9\\ 1.9\\ 1.9\\ 1.9\\ 1.9\\ 1.9\\ 1.9$	$\begin{array}{c} 26\\ 98\\ 5\\ 61\\ 6.5\\ 0.7\\ 13.6\\ 7.5\\ 14\\ 1.8\\ 18.2\\ 0.52\\ 133\\ 5.3\\ 61\\ 0.6\\ 0.15\\ 7.1\\ 14.1\\ 0.17\\ 663\\ 24\\ <5\\ 22\\ 87\\ 19\\ 169\\ 2.3\\ 3.4\\ 1.6\\ 2.4\\ <5\\ 33\\ 0.1\\ 0.44\\ 2.1\\ <1\\ 599\\ 1.0\\ 26\\ 0.3\\ 1.1\\ 287\\ 21\\ \end{array}$	2.4 106 10 38 <1 0.4 5.1 1.6 14 0.68 6.2 0.1 78 2.1 6 0.4 0.07 3.1 2.8 0.07 543 55 11 7 87 8.2 <6 0.5 1.6 1.9 0.49 <5.5 25 0.1 0.07 1.1 1 539 1.3 23 0.26 42 23	2.2 181 13.2 37 1.0 0.4 7.1 3.5 17 1.0 5.0 0.10 63 2.4 7 0.6 0.08 2.9 3.3 0.10 543 43 8.0 11 44 12 3.2 0.6 1.9 2.9 0.70 <5 26 0.10 1.4 <1 480 2.3 41 0.4 0.5 34 9	$\begin{array}{c} 6.4\\ 83\\ 2\\ 33\\ 2.0\\ 1.0\\ 5.0\\ 2.5\\ 5.7\\ 0.92\\ 8.8\\ 0.3\\ 63\\ 2.2\\ 16\\ 0.17\\ 0.07\\ 1.7\\ 0.9\\ 0.12\\ 241\\ 30\\ 15\\ 12\\ 30\\ 5.1\\ 23\\ 1.9\\ 1.1\\ 1.3\\ 1.0\\ <5\\ 17\\ 0.05\\ 0.29\\ 0.6\\ <1\\ 240\\ 8.0\\ 16\\ 0.25\\ 0.76\\ 68\\ 11\end{array}$	$\begin{array}{c} 46\\ 161\\ 3.6\\ 11\\ 3.0\\ 0.7\\ 2.7\\ 11\\ 5.7\\ 0.80\\ 20.5\\ 0.35\\ 50\\ 2.7\\ 125\\ 0.2\\ 0.13\\ 1.2\\ 1.2\\ 0.14\\ 121\\ 20\\ <3\\ 20\\ 87\\ 4.3\\ 184\\ 5.1\\ 1.5\\ 1.9\\ 1.3\\ <5\\ 10\\ <0.1\\ 0.29\\ 0.6\\ <1\\ 240\\ 0.8\\ 16\\ 0.10\\ 0.73\\ 179\\ 9\\ 9\end{array}$

¹ Two other newly collected coals (tables 3 and 5, CYC 1 and CYC 2 feed coals) came from the same mine as the sample C35303.

SUMMARY AND CONCLUSIONS

The mineralogical and chemical composition of (1) marketed Illinois coals from 35 mines and (2) feed coal, fly ash, and bottom ash from three different types of power generating units (PC, CYC, and FBC) burning Illinois coals were investigated. This investigation was the first time the nature and combustion behavior of mineral matter in marketed Illinois coals were determined in such a systematic and detailed manner. The general findings were as follows:

- The mean clay mineral content of the marketed Illinois coals was about 52% of the total mineral matter present in these coals. The mean values for abundances of the individual minerals in the coals (with standard deviations shown as ± values) were kaolinite, 2.07 ± 0.56%; quartz, 2.06 ± 0.78%; mixed-layered illite/smectite, 1.57 ± 0.70%; pyrite, 1.56 ± 0.50%; illite, 1.29 ± 0.46%; calcite, 0.84 ± 78%; chlorite, 0.18 ± 0.13%; marcasite, 0.13 ± 0.12%; plagioclase, 0.07 ± 0.05%; and K-feldspar, 0.01 ± 0.01%. Quartz partially survived the combustion process and was partially converted to silica glass at the power plants. The clay minerals and feldspars were converted to aluminosilicate glass and to mullite. Pyrite and marcasite were converted to hematite, magnetite, and amorphous Fe oxides. Calcite was converted to lime, portlandite, anhydrite, and gypsum.
- Concentrations of some minor and trace elements increased with increased concentrations of certain minerals in the 35 coals. Among the 15 HAP elements (As, Be, Cd, Co, Cr, F, Hg, Mn, Ni, P, Pb, Sb, Se, Th, U) in the coals, the concentrations of Cr, F, and Th were strongly and positively correlated with the content of mixed-layered illite/smectite or the combination of mixed-layered illite/smectite and illite. Strong positive correlations existed also between Th and kaolinite contents and between Mn and calcite contents. There were weak positive correlations between Co and chlorite, Hg and marcasite, and Hg and pyrite. These relationships between minerals and elements are important considerations for designing methods for pre- and post-combustion control of HAP emissions from coal-fired power plants.
- Upon combustion, a portion of each HAP element is mobilized and becomes airborne. The mobilized portions of an element could partially be adsorbed on surfaces of hardware on the cool side of a power plant instead of being emitted into the atmosphere. In that case, the atmospheric emission of HAPs, except highly volatile ones (F, Hg, Se), could be less than the emission values reported here. Therefore, in this study, the term "mobility" instead of "emission" was used to express the portion of an element that was not retained in CCRs. The mobilities of 15 HAPs from the FBC plant were generally less than the mobilities from the other plants. Relatively large mobilities (more than 50% of the original amount in coal) were observed for F and Mn at the FBC plant and for F, Hg, and Se at the CYC and PC plants.
- Combustion mobilities of trace elements are controlled by their volatility and affinity for various coal combustion phases. The elements investigated in this study occurred in greater concentrations in the fly ash than in the bottom ash in most cases, which is generally consistent with the literature data. The relative enrichment of the elements in the fly ash most likely resulted from the combination of (1) the partial volatilization of the elements from the bottom ash and their subsequent condensation on the fly ash particles upon cooling and (2) preferential enrichment of some elements in finely disseminated coal minerals that ended up largely in the fly ash. Some CCRs contained high concentrations of certain elements that could be recovered and sold.
- There were differences in the mineralogical and chemical compositions of two sets of marketed coal samples taken from four Illinois preparation plants at a 4-year interval. Some of the differences could result from expected sample variations independent of when the samples were collected. However, it is likely that the observed differences largely resulted from changes in characteristics of in-place coal as mining moved to different reserve blocks over time and from changes in coal cleaning in response to market demands. Therefore, historical coal quality data on clean coal products from a given mine cannot be used to accurately determine the quality of the current production from the same mine.

REFERENCES

- ACAA, 1999, Coal combustion product production and use for 1998: Alexandria, Virginia, American Coal Ash Association.
- ASTM, 1995, Annual book of ASTM standards, section 5, volume 0.505—Gaseous fuels; coal and coke: Philadelphia, Pennsylvania, ASTM.
- Barbieri, L., I. Lancelotti, T. Manfredini, I. Queralt, J. M. Rincon, and M. Romero, 1999, Design, obtainment and properties of glasses and glass-ceramics from coal fly ash: Fuel, v. 78, p. 271–278.
- Bool, L.E., and J.J. Helble, 1995, A laboratory study of the partitioning of trace elements during pulverized coal combustion: Energy and Fuels, v. 9, p. 880–887.
- Cereda, E.M.,G.M. Braga, M. Pedretti, G.W. Grime, and A. Baldacci, 1995, Nuclear microscopy for the study of coal combustion related phenomena: Nuclear Instruments and Methods in Physics Research B, v. B99, no. 1/4, p. 414–418.
- Clarke, L.B., 1992, Application for coal-use residues: London, International Energy Agency (IEA) Coal Research IEACR/50.
- Clarke, L.B., and L.L. Sloss, 1992, Trace elements—Emission from coal combustion and gasification: London, IEA Coal Research IEACR/49.
- Dale, L.S., S.A. Lawrensic, and J.F. Chapman, 1992, Mineralogical residence of trace elements in coal—Environmental implications of combustion in power plants, CSIRO investigation report CET/IRO58: North Ryde, New South Wales, Australia, CSIRO, 79 p.
- Davidson, R.M., and L.B. Clarke, 1996, Trace elements in coal: London, IEA Coal Research, IEAPER/21.
- DeBarr, J.A., D.M. Rapp, M. Rostam-Abadi, J.M. Lytle, and M.L. Rood, 1996, Valuable products from utility fly ash: Cartersville, Illinois, Illinois Clean Coal Institute Final Report.
- Demir, I., and R.D. Harvey, 1990, Abundance and origin of major minerals in the Herrin Coal,
 Illinois Basin, *in* R. W. Bryers and K. S. Vorres, eds., Mineral matter and ash deposition from coal: New York, Engineering Foundation, p. 13–22.
- Demir, I., R.D. Richard, R.R. Ruch, H.H. Damberger, C. Chaven, J.D. Steele, and W.T. Frankie, 1994, Characterization of available (marketed) Illinois coals from Illinois mines: Illinois State Geological Survey Open File Series 1994-2.
- Demir, I., R.R. Ruch, H.H. Damberger, R.D. Harvey, J.D. Steele, and K.K. Ho, 1998, Environmentally critical elements in channel and cleaned samples of Illinois coals: Fuel, v. 77, p. 95–107.
- DeVito, M., and S.B. Bhagwat, 1997, Correlate coal/scrubber parameters with Hg removal and Hg species in flue gas: Paper presented at the 15th Annual Contractors' Meeting of Illinois Clean Coal Institute, July 29–30, 1997, Champaign, Illinois.
- DeVito, M.S., and B.L. Jackson, 1994, Trace element partitioning and emissions in coal-fired utility systems: Paper presented at the 87th Annual Meeting and Exhibition of Air and Waste Management Association, June 19–24, 1994, Cincinnati, Ohio.
- Dhir, R.K., and M.R. Jones, 1999, Development of chlorite-resisting concrete using fly ash: Fuel, v. 78, p. 137–142.
- Dreher, G.B., W.R. Roy, and J.D. Steele, 1996. Laboratory studies on the codisposal of fluidizedbed combustion residue and coal slurry solid: Illinois State Geological Survey Environmental Geology 150.
- Galbreath, K., C. Zygarlicke, G. Casuccio, T. Moore, P. Gottlieb, N. Agron-Olshina, G. Huffman, A. Shah, N. Yang, J. Vleeskens, and G. Hamburg, 1996, Collaborative study of quantitative coal mineral analysis using computer-controlled scanning electron microscopy: Fuel, v. 75, p. 424–430.
- Gluskoter, H.J., 1965, Electronic low temperature ashing of bituminous coal: Fuel, v. 44, p. 285–291.

- Gluskoter, H.J., R.R. Ruch, W.G. Miller, R.A. Cahill, G.B. Dreher, and J.K. Khun, 1977, Trace elements in coal: Illinois State Geological Survey Circular 499, 154 p.
- Gullet, B.K., and K. Ragnunathan, 1994, Reduction of coal-based metal emissions by furnace sorbent injection: Energy and Fuels, v. 8, p. 1068–1076.
- Harvey, R.D., R.A. Cahill, C.-L. Chou, and J.D. Steele, 1983, Mineral matter and trace elements in the Herrin and Springfield Coals, Illinois Basin Coal Field: Illinois State Geological Survey Contract/Grant Report 1983-4.
- Harvey, R.D., and R.R. Ruch, 1986, Mineral matter in Illinois and other coals, *in* K. S. Vorres, ed., Mineral matter and ash in coal, American Chemical Society Symposium Series 301: Washington, D.C., American Chemical Society, p. 10–40.
- Helble, J.J., 1994, Trace element behavior during coal combustion: Results of a laboratory study: Fuel Processing Technology, v. 39, p. 159–172.
- Hughes, R.E., P.J. DeMaris, G.B. Dreher, D.M. Moore, and M. Rostam-Abadi, 1996, Brick manufacture with fly ash from Illinois coals: Carterville, Illinois, Illinois Clean Coal Institute Final Technical Report.
- Hughes, R.E., D.M. Moore, and H.D. Glass, 1994, Qualitative and quantitative analysis of clay minerals in soils, *in* J .E. Amonette and L.W. Zelazny, eds., Quantitative methods in soil mineralogy: Madison, Wisconsin, Soil Science Society of America Miscellaneous Publication, p. 330–359.
- Hughes, R.E., and R.L. Warren, 1989, Evaluation of the economic usefulness of earth materials by x-ray diffraction, *in* R.E. Hughes and J.C. Bradbury, eds., Proceedings of the 23rd Forum on the Geology of Industrial Minerals: Illinois State Geological Survey Industrial Minerals Notes 102, p. 47–57.
- ISGS, 1994, QA/QC plan for the analytical geochemistry section: Illinois State Geological Survey internal document.
- Kruse, C.W., C. Chaven, R.E. Hughes, R.R. Ruch, and J.M. Lytle, 1994. Illinois Basin coal sample program: Carterville, Illinois, Illinois Clean Coal Institute Final Technical Report.
- Meij, R., 1992, A mass balance study of trace elements in a coal-fired power plant with a wet fgd facility, *in* G. Vourvopolis, ed., Elemental analysis of coal and its by-products, River Edge, New Jersey, World Scientific Publishing Co., p. 299–318.
- Meij, R., 1993, The fate of trace elements at coal-fired power plants, *in* W. Chow and L. Lewin, eds., Proceedings of 2nd International Conference on Managing Hazardous Air Pollutants: Palo Alto, California, Electric Power Research Institute EPRITR-104295, p. V83–V105.
- Meij, R., 1994a, Distribution of trace species in power plant streams: A European perspective, *in* Proceedings of 56th Annual Meeting of American Power Conference, April 25–27, 1994, Chicago, Illinois, v. 56-1, p. 458–463.
- Meij, R., 1994b, Trace element behavior in coal-fired power plants. Fuel Processing Technology, v. 39, p. 199–217.
- Moore, D.M., and R.C. Reynolds Jr., 1997, X-ray diffraction and the identification and analysis of clay minerals: New York, Oxford University Press.
- Moore, T., 1994, Hazardous air pollutants: measuring in micrograms: EPRI Journal, January/ February, p. 7–15.
- National Research Council, 1980, Trace element geochemistry of coal resource development related to environmental quality and health: Washington, D.C., National Academy of Science, 153 p.
- Niewiadomski, M., J. Hupka, R. Bokotko, and J.D. Miller, 1999, Recovery of coke fines from fly ash by air sparged hydrocyclone flotation: Fuel, v. 78, p. 161–168.
- Querol, X., J.L. Fernandez-Turiel, and A. Lopez-Soler, 1995, Trace elements in coal and their behavior during combustion in a large power station: Fuel, v. 74, p. 331–343.

- Rao, C.P., and H.J. Gluskoter, 1973, Occurrence and distribution of minerals in Illinois coals: Illinois State Geological Survey Circular 476.
- Reynolds, R.C., Jr., 1985, NEWMOD[®], a computer program for the calculation of one-dimensional diffraction patterns of mixed-layered clays: Hanover, New Hampshire, published by R.C. Reynolds, Jr.
- Reynolds, R.C., Jr., 1989, Principles and techniques of quantitative analysis of clay minerals by xray powder diffraction, *in* D. R. Pevear and F. A. Mumpton, eds., Quantitative mineral analysis of clays: Evergreen, Colorado, Clay Minerals Society, p. 4–36.
- Rose, A.W., H.E. Hawkes, and J.S. Webb, 1979, Geochemistry in mineral exploration (2nd ed.): New York, Academic Press.
- Roy, W.R., G.B. Dreher, J.D. Steele, R.G.Darmody, D.Tungate, W.E. Giles, and S.C. Phifer, 1997, Direct revegetation of coal slurry after amendment with FBC residues: Carterville, Illinois, Illinois Clean Coal Institute Final Technical Report.
- Russell, S.J., and S.M. Rimmer, 1979, Analysis of mineral matter in coal, coal gasification ash, and coal liquefaction residues by scanning electron microscopy and x-ray diffraction, *in* C. Kan Jr., ed., Analytical methods for coal and coal products, v. 3: New York, Academic Press, p. 133–162.
- Slaughter, M., 1989, Quantitative determination of clay and other minerals in rocks, *in* D. L. Bish and J. E. Post, eds., Quantitative mineral analysis of clays, Washington, D.C., Mineralogical Society of America, p. 183–216.
- Sloan, J.J., R.H. Dowdy, M.S. Dolan, and G.W. Rehm, 1999, Plant and soil response to fieldapplied flue gas desulfurization residue: Fuel, v. 78, p. 169–174.
- Snyder, R.L., and D.L. Bish, 1989, Quantitative analysis, *in* D.L. Bish and J.E. Post, eds., Modern powder diffraction, Reviews in mineralogy, v. 20: Washington, D.C., Mineralogical Society of America, p. 101–144.
- Suarez-Fernandez, G., X. Querol, J.L. Fernandez-Turiel, A.B. Benito-Fuertes, and M.R. Martinez-Tarazona, 1996, The behavior of elements in fluidized bed combustion: American Chemical Society Fuel Chemistry Preprints, v. 41, p.796–800.
- Swaine, D.J., 1990, Trace elements in coal: London, Butterworth, 276 p.
- Thomas, J. Jr., H.D. Glass, W.A. White, and R.M. Trandel, 1977, Fluoride content of clay minerals and argillaceous earth materials: Clays and Clay Minerals, v. 25, p. 278–284.
- Tumati, P.R., and M.S. DeVito, 1991, Retention of condensed/solid phase trace elements in an electrostatic precipitator, *in* Managing hazardous air pollutants—State of the art: Palo Alto, California, Electric Power Research Institute.
- Tumati, P.R., and M.S. DeVito, 1993, Trace element emissions from coal combustion—A comparison of baghouse and especially collection efficiency, *in* Proceedings of 3rd International Conference on Effects of Coal Quality on Power Plants: Palo Alto, California, Electic Power Research Institute EPRI TR-1-2280, p. 335–348.
- U.S. Public Law 101-549, 1990, Clean Air Act Amendments, Title 3, 104 Statute 2531–2535.
- USEPA, 1998, Study of hazardous air pollutant emissions from electric utility steam generating units, Final report to Congress: v. 1 and 2, EPA-453/R-98-004b.
- Ward, C.R., 1977, Mineral matter in the Springfield-Harrisburg (No. 5) Coal Member in the Illinois Basin: Illinois State Geological Survey Circular 498, 35 p.
- Wesnor, J.D., 1993, ABB's investigation into the utility air toxic problems, Paper presented at EPRI/EPA SO₂ Conference, Boston, Massachusetts.
- Wirtz, G.P., and J.M. Bukowski, 1996, Extruded honeycomb construction materials from fly ash: Carterville, Illinois, Illinois Clean Coal Institute Final Report.