APPENDIX R

KEMPER COUNTY IGCC PROJECT HAZARDOUS AIR POLLUTANT RISK SCREENING ANALYSES

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Prepared for: Southern Company Services, Inc.

Risk Screening Analysis of Hazardous Air Pollutant Emissions from the Proposed Kemper County IGCC Facility, 50% CO₂ Capture Case Kemper County, Mississippi

AECOM, Inc. May 2009 Document No.: 06204-041-0001a

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Risk Screening Analysis of Hazardous Air Pollutant Emissions from the Proposed Kemper County IGCC Facility, 50% CO₂ Capture Case Kemper County, Mississippi

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Contents

1.0	Introduction	.1
2.0	HAP Selection	.1
3.0	Evaluation Methods	.1
4.0	Modeling Approach	.2
5.0	Findings	.4
6.0	Summary	.4
7.0	References	.5

List of Tables

Table 1:	Estimates of HAP Emissions for Each of Two IGCC Stacks for Siemens Turbines and Ranking in Terms of Inhalation Risk
Table 2:	Estimate of Mercury Emissions for Each of Two IGCC Stacks for Siemens Turbines7
Table 3:	IGCC Source Parameters7
Table 4:	Modeled Project-Related Chronic Inhalation Risk Estimates for Arsenic, Cadmium, and Mercury
Table 5:	Modeled Project-Related Acute Inhalation Risk Estimates for Arsenic, Cadmium, and Mercury9
Table 6:	Maximum Modeled Long-Term Mercury Air Concentrations (µg/m ³)9
Table 7:	Comparison of Modeled Mercury Deposition from the Facility with Measured Deposition (g/m²/year)9

List of Figures

Figure 1:	Relative Location of Kemper IGCC Project Site, Meteorological Data Sources, and Ambient Mercury Monitors
Figure 2:	Meridian Key Field 5-year (1991-1995) Wind Rose11
Figure 3:	Project Receptors Used to Determine Maximum Annual and Short-Term Impacts – Near-Field Receptors
Figure 4:	Project Receptors Used to Determine Maximum Annual and Short-Term Impacts – Far-Field Receptors
Figure 5:	Kemper County Receptor Grid14
Figure 6:	Location of Maximum Air Concentration Impacts and Mercury Deposition15

1.0 Introduction

AECOM Environment has conducted a screening level evaluation of the potential effects of hazardous air pollutant (HAP) emissions from the proposed Kemper County Integrated Gasification Combined Cycle (IGCC) stacks.

2.0 HAP Selection

The evaluation considers the HAP emissions associated with continuous firing of syngas in the combustion turbines and natural gas in the associated duct burners. Although the combustion turbines are also capable of combusting natural gas, the fraction of time that natural gas alone would be fired is expected to be small. Two combustion turbine vendors are under consideration, Siemens and GE. HAP emissions were quantified for both turbines and the emission rates were found to comparable, with the Siemens turbines resulting in slightly higher potential emissions. Therefore, the Siemens turbines were addressed in this assessment. The HAPs emissions for a Siemens turbine are provided in Table 1 for all HAPs with the exception of mercury. Mercury emissions are provided in Table 2.

This assessment addressed two classes of health risk: carcinogenic risk and non-carcinogenic risk for longand short-term (acute and chronic) inhalation of airborne concentrations. For inhalation pathway risk, this screening-level assessment identifies two HAPs, which were selected based on relative emissions and longterm inhalation toxicity. To identify the HAPs to include in the assessment, the HAP emissions were ranked according to relative cancer and chronic non-cancer risk associated with their emissions. The U.S. EPA's Office of Air Quality Planning and Standards (OAQPS) has tabulated dose-response assessments that OAQPS uses for risk assessments of hazardous air pollutants and has made these available on the Technology Transfer Network at http://www.epa.gov/ttn/atw/toxsource/summary.html. For this ranking the Dose-Response Values compiled by U.S. EPA OAQPS (Ref: Prioritized Chronic Dose-Response Values, provided at http://www.epa.gov/ttn/atw/toxsource/table1.pdf) were applied. To estimate the relative contribution of HAP to inhalation cancer risk, each HAP emission rate listed in Table 1 was multiplied the by U.S. EPA's Unit Risk Estimates (URE). In a similar way, the relative percent contribution among HAPs was estimated by dividing the HAP emission rates by their respective chronic inhalation concentration levels (referred to here as Reference Air Concentrations or RACs). The result of this ranking, shown in Table 1, indicates that the two HAPs that contribute most to both cancer and non-cancer inhalation risks are arsenic and cadmium. Together these two HAPs contribute nearly 75% of the inhalation cancer risk associated with the IGCCs and nearly 50% of the chronic non-cancer inhalation risk. The implication is that if the modeled risk for these two HAPs is well below significance levels, it is assured that the combined risk associated with all of the HAPs will also be insignificant.

In addition to risks due to direct inhalation of HAPs, it is well-recognized that the routine ingestion of fish that contain elevated concentrations of mercury can pose non-carcinogenic chronic health risk. Some of the mercury in fish may result from deposition of airborne concentrations onto watersheds and subsequent bioaccumulation through the aquatic food chain. Although the presence of mercury in the ambient environment is a global issue, because mercury is among the HAPs that are known to be emitted from coal combustion, the local dispersion and deposition of this HAP from the IGCCs were also evaluated.

3.0 Evaluation Methods

To assess inhalation risk, AERMOD was applied to estimate the facility's contribution to ambient concentrations at the point of maximum off-site impact, as well as the average concentrations throughout Kemper County. Modeled long-term concentrations of arsenic, cadmium and mercury were then added to

U.S. EPA's model estimates of the pre-existing concentrations in Kemper County associated with other local and regional emission sources for these three HAPs as computed by the 1999 National-Scale Air Toxics Assessment (NATA, http://www.epa.gov/ttn/atw/nata1999/tables.html). In modeling long-term concentrations, as described in more detail below, an average concentration over a 5-year period was computed assuming that the IGCCs ran on syngas continuously at peak load. The level of incremental inhalation cancer and non-cancer risk, and combined risk (facility plus NATA), for each HAP were then evaluated. To evaluate the maximum individual lifetime cancer risk the estimated long-term average air concentration was multiplied by the URE (see Table 1). The result of this calculation is the increase in probability of a person contracting cancer after 70 years of exposure. A cancer risk value of less than 1x10⁻⁶ is generally recognized to be insignificant (U.S. EPA, 1991). For chronic non-cancer effects, the long-term average concentration was divided by the respective RAC to form what is referred to as a Hazard Quotient (HQ). A HQ value of less than 1.0 indicates that adverse non-cancer health effects are highly unlikely to occur (U.S. EPA, 2008). To assess potential acute health effects, the maximum modeled 1-hour average concentrations for the IGCCs were divided by minimum acute RAC found in EPA guidance (Acute Dose-Response Values for Screening Risk Assessments (6/12/2007) in http://www.epa.gov/ttn/atw/toxsource/table2.pdf.

In addition to the evaluation of inhalation health risks, the long-term deposition of mercury was also evaluated. Mercury is emitted in three forms, elemental mercury vapor, reactive gaseous mercury (RGM), and particulatebound mercury (PBM). As indicated in Table 2, it is estimated that 90% of the total mercury emitted from the IGCCs is in the form of elemental mercury, 10% RGM and un-guantified trace amounts of PBM. In terms of the contribution to wet or dry deposition, elemental mercury is essentially inert because it has an extremely low solubility and is non-reactive. Therefore, most of the elemental mercury remains airborne and is transported over long distances (U.S. EPA, 1997). PBM is deposited with the particles with which it is incorporated, but in this case the rate of emission is so low that the rate of PBM deposition cannot be quantified. In contrast, RGM is soluble and reactive such that it is subject to both dry and wet deposition. Dry deposition occurs when airborne RGM comes into contact with elements on the earth's surface such as vegetation and water bodies. Wet deposition occurs when precipitation falling through the plume captures soluble RGM vapor. As discussed further below, these deposition processes are simulated by U.S. EPA's AERMOD air quality model. The ambient concentration and deposition of mercury from the IGCC units at the point of maximum impact, and the average over Kemper County, are evaluated by comparing modeled results to measurements taken at two locations along the Gulf coast, and a third located in Oak Grove, MS, where there are long-term mercury monitoring networks (see Figure 1).

4.0 Modeling Approach

U.S. EPA's AERMOD model was applied to estimate air concentrations of arsenic, cadmium and mercury, and deposition of mercury emitted from the IGCC units. Five years of representative meteorological data (1991-1995) were applied, including National Weather Service (NWS) surface and precipitation data from Meridian Key Field in Meridian, MS and upper air data from Jackson International Airport in Jackson, MS. The relative location of the meteorological measurement sites with respect to the project site are provided in Figure 1. Processed, AERMOD-ready meteorological data was provided by the Mississippi Department of Environmental Quality (MDEQ) on their website (http://www.mississippi.gov/frameset.jsp?URL=http%3A%2F%2F %2Fwww.deq.state.ms.us%2FMDEQ.nsf%2Fpage%2FMain_Home%3FOpenDocument). The 5-year wind rose shown in Figure 2 indicates prevailing winds are primarily southerly, but northerly winds are also frequent. In accordance with U. S. EPA guidance, a sensitivity analysis was conducted using both the surface site characteristics (roughness, albedo and Bowen ratio) from the project site and the NWS station site, and the site characteristics that resulted in the greater modeled concentrations were applied. On this basis, the surface characteristics of the NWS site at Meridian were used. Two sets of receptors were used in the analysis. The first set, which was developed for the air quality modeling in support of the project permitting effort, was applied to model the concentration and deposition at the maximum point of impact. This receptor grid consisted of "tight" receptors spaced 50 meters apart along the fence line and extending out 500 meters

from the fence-line. Beyond 500 meters, a spacing of 100 meters was used up to one-kilometer from the facility. Between one- and five-kilometers, a spacing of 500 meters was used. Between five- and ten-kilometers, a spacing of 1,000 meters was used. Between ten- and twenty-kilometers, a spacing of 2,000 meters was used. For this grid, the near field receptors are shown in Figure 3 and the far field receptors are shown in Figure 4. A second set of receptors comprised of a uniform, 1 km spaced receptor grid across Kemper County, was used to model the Kemper County-wide average for comparison to NATA county-level results. Receptor elevations were developed using Digital Elevation Model (DEM) data from USGS. The Kemper County grid is shown in Figure 5.

Table 3 provides the source parameters used to apply AERMOD. For the IGCC dispersion modeling and RGM deposition analysis, the general modeling procedures and options specified in the current versions of in the AERMOD User's Guide (U.S. EPA 2004), AERMOD Implementation Guide (U.S. EPA 2009), and the Guideline on Air Quality Models (GAQM; U.S. EPA 2005) were followed. To model ambient concentrations of arsenic, cadmium and elemental mercury, it was conservatively assumed that no deposition occurred. Deposition was accounted for in modeling the concentration of RGM. Modeling was conducted in a manner consistent with U.S. EPA guidance and standard practices, including the use of regulatory default options, as appropriate. The building downwash analysis was performed using the most recent version of EPA's Building Profile Input Program (BPIP) (Version 04274) with the plume rise model enhancements (PRIME) building downwash algorithms, consistent with the air quality modeling.

Aerial imagery (2004) was examined to determine the land-use characteristics around the project site for use as input into the model for mercury deposition calculations. The area in the vicinity of the project was found to be heavily forested. As a result, category 4 (forest) was used for the gas dry deposition land-use category for all 36 100 radials in the mercury deposition modeling as described in the Addendum to the AERMOD User's Guide (U.S EPA, 2006).

The transport and mobility of a pollutant are determined by the physical properties of the specific

pollutant. For deposition modeling, AERMOD requires several pollutant-specific parameters:

(1) diffusivity in air; (2) diffusivity in water; (3) leaf cuticular resistance to lipid uptake; and (4) the Henry's Law constant. The following values of these parameters recommended by U.S. EPA were applied:

- diffusivity in air: 6.0E-2 cm²/sec⁽¹⁾
- diffusivity in water: 5.25E-6 cm²/sec⁽²⁾
- Cuticular resistance: 1.0E7 sec/m⁽¹⁾
- Henry's Law constant: 6.0E-6 pa-m³/mol⁽¹⁾
- Reactivity factor: 1.0⁽¹⁾
- ⁽¹⁾ Source is Wesley, M.L., Doskey, P.V., and Shannon, J.D. Deposition Parameterizations for the Industrial Source Complex (ISC3) Model, Argonne National Laboratory, June, 2002, Table 7 (p.27).
- ⁽²⁾ Source is U.S. EPA's Human Health Risk Assessment Protocol for Hazardous Waste Combustion Facilities [Publication No. EPA 530-R-05-006], September 2005. Information from companion HHRAP Database – Value for mercuric chloride. <u>http://www.epa.gov/osw/hazard/tsd/td/combust/riskvol.htm#volume2</u> - scroll down to "Download HHRAP Companion Database (ACCESS)"

5.0 Findings

Figure 6 provides the location of the maximum modeled air concentrations for the modeled HAPs and the location of the highest RGM deposition relative to the facility. The maximum air concentration impacts, along with the maximum dry deposition, were found to be along the southeast fence-line of the power block portion of the facility. The maximum wet and total RGM deposition was found to occur along, and just past, the northeast fence-line of the power block, directly to the north of the IGCC stacks.

Table 4 provides inhalation risk estimates associated with arsenic, cadmium and mercury emitted from the facility and the combined county-wide risk of these HAPs, incorporating the 1999 NATA Kemper County modeling results. The maximum cancer risk, which is associated with arsenic, is less than 2×10^{-7} at the point of maximum, and the county-wide-average risk, including the proposed IGCC units is only about 1×10^{-8} . These levels are below the EPA's recommended significance levels of 1×10^{-6} . The hazard quotients for each of the three HAPs are also insignificant with the highest incremental HQ, which occurs for cadmium, of less than 3×10^{-3} compared to a significance level of 1.0. The corresponding Kemper County HQ for cadmium is about 2×10^{-4} . The results of acute inhalation assessment, provided in Table 5, indicate that arsenic is the HAP with the largest acute hazard quotient with a value of about 5×10^{-3} . This analysis demonstrates that the proposed IGCC units will not cause or contribute significantly to acute or chronic health effects and cancer risk.

The analyses for mercury are provided in Tables 6 and 7. Table 6 compares maximum modeled long-term air concentrations of elemental mercury and RGM with available measurements from two measurement sites located to the south near the Gulf Coast. The long-term concentration of elemental mercury is two orders of magnitude less than the measured ambient concentrations, and the maximum modeled RGM concentration is about a third of the ambient concentrations. Thus, the combined maximum long-term modeled concentration of elemental mercury and RGM is only one hundredth of the measured ambient concentrations. This finding is consistent with NATA which specifies a U.S. background mercury concentration associated with global transport of 1.5 x 10-3 μ g/m³. Table 7 provides a comparison of the maximum modeled RGM deposition from the facility with values developed from measurements from the OLF study and the NADP Mercury Deposition Network. The maximum long-term wet deposition is one to two orders of magnitude less than measurements. The modeled dry deposition at the location of maximum deposition is comparable to the dry deposition estimates from the OLF study. The result is that the total modeled mercury deposition is only about one eighth of total ambient deposition. The deposition analysis therefore demonstrates that the project will not contribute substantially to the ambient concentration or deposition of mercury in the vicinity.

6.0 Summary

An assessment of the HAP emissions from the proposed IGCC units was conducted using the conservative assumption that the IGCC units will operate continuously at 100% load on syngas for five years. Modeling of concentrations and RGM deposition was conducted with AERMOD, a U.S. EPA designed model that provides conservative estimates of air quality impacts. The results of the assessment indicate that no HAP emitted from the project will result in, or contribute to, human health risk due to inhalation. In addition, the ambient concentrations of mercury resulting from the IGCC units were modeled to be orders of magnitude less than ambient concentrations, and the modeled deposition of mercury was less than has been measured at nearby study areas.

7.0 References

U.S. Environmental Protection Agency (EPA), 1997. Mercury Stud report to Congress, Volume III: Fate and Transport of Mercury in the Environment, EPA-452/R-97-005, December.

U.S. EPA, 2009. AERMOD Implementation Guide - Revised, March 19, 2009

U.S. EPA, 2006. Addendum: Users Guide for the AMS/EPA Regulatory Model – AERMOD

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U.S. EPA, 2005. Guideline on Air Quality Models (as incorporated in Appendix W of 40 CFR Part 51; EPA, 2005).

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U.S.EPA, 2008. 1996 National-Scale Air Toxics Assessment: Summary of Results [http://www.epa.gov/nata/risksum.html].

U.S.EPA, 1991. Role of the Baseline Risk Assessment in Superfund Remedy Selection Decisions. OSWER Directive #9355.0-30. April. [http://www.epa.gov/oswer/riskassessment/pdf/baseline.pdf].

Wesley, M.L., Doskey, P.V., and Shannon, J.D. Deposition Parameterizations for the Industrial Source Complex (ISC3) Model, Argonne National Laboratory, June, 2002.

	Maximum Short-Term Emissions ^(1,3) (Ib/hour)	Cancer Unit Risk Estimate ⁽⁴⁾ (1/µg/m ³)	Percent of Cancer Risk	Reference Air Concentration ⁽⁴⁾ (µg/m3)	Percent of Chronic Hazard Index
Volatile Organic Compounds					
Acetaldehyde	2.29E-02	2.20E-06	0.1%	9.00	0.1%
Acrolein	2.22E-03	NA		0.02	5.4%
Benzene	2.16E-02	7.80E-06	0.2%	30.00	0.0%
Ethylbenzene	9.03E-03	NA		1000.00	0.0%
Formaldehyde	1.01E-01	5.50E-09	0.0%	9.80	0.5%
Toluene	2.69E-02	NA		5000.00	0.0%
Xylene	2.58E-02	NA		100.00	0.0%
Polycyclic Organic Matter (POM)					
Polycyclic Aromatic Hydrocarbons					
(PAH)	1.90E-04	1.10E-03	0.2%	200.00	0.0%
2-Methylnapthalene	1.20E-03	NA		NA	
Acenapthylene	8.50E-05	NA		NA	
Benzo(a)anthracene	7.50E-06	1.10E-04	0.0%	NA	
Benzo(e)pyrene	1.80E-05	NA		NA	
Benzo(g,h,i)perylene	3.10E-05	NA		NA	
Napthalene	1.60E-03	3.40E-05	0.1%	3.00	0.0%
Metals					
Antimony	1.27E-02	NA		NA	
Arsenic	9.79E-03	4.30E-03	45.7%	0.03	15.8%
Beryllium	3.00E-03	2.40E-03	7.8%	0.02	7.3%
Cadmium	1.37E-02	1.80E-03	26.8%	0.02	33.1%
Chromium VI (2)	1.49E-03	1.20E-02	19.4%	0.10	0.7%
Cobalt	2.64E-03	NA		0.10	1.3%
Lead	1.30E-02	NA		0.15	4.2%
Manganese	1.40E-02	NA		0.05	13.6%
Mercury (total)	3.61E-03	NA		0.30	0.6%
Elemental Mercury	3.25E-03	NA		NA	
Reactive Gaseous Mercury	3.61E-04	NA		NA	
Particulate-bound Mercury	Trace	NA		NA	
Nickel	1.83E-02	NA		0.09	9.8%
Phosphorous	1.11E-02	NA		0.07	7.7%
Selenium	1.40E-02	NA		NA	
Inorganic Compounds					
Carbon Disulfide	1.47E-01	NA		700.00	0.0%

Table 1: Estimates of HAP Emissions for Each of Two IGCC Stacks for Siemens Turbines and **Ranking in Terms of Inhalation Risk**

(1) Emission rates based on emission factors from "A Study of Toxic Emissions from a Coal-Fired Gasification Plant", Radian Corporation, December 1995.

(2) 12% of total chromium emissions characterized as hexavalent for coal combustion per U.S. EPA 2005 National Emissions Inventory Data & Documentation (http://www.epa.gov/ttn/chief/net/2005inventory.html). Estimates are based on full load operating scenarios with Duct Burner firing.

(3)

(4) Source: Prioritized Chronic Dose-Response Values: http://www.epa.gov/ttn/atw/toxsource/table1.pdf.

	Average Short- Term Syngas (Ib/hour)	Average Annual (Ib/year)
Total Mercury	0.0036	31.64
Elemental (90% of Total)	0.0033	28.47
RGM (10% of Total)	0.00036	3.16
Particle-Bound (Only trace amounts)	Trace	Trace

Table 2: Estimate of Mercury Emissions for Each of Two IGCC Stacks for Siemens Turbines

Table 3: IGCC Source Parameters

Parameter	Valu	le
	CC1A	CC2A
UTM Coordinate East (m) ⁽¹⁾	335223.3	335177.6
UTM Coordinate North (m) ⁽¹⁾	3614049.2	3614050.0
Stack Base Elevation (ft)	474	474
Stack Height (ft)	325	325
Stack Diameter (ft)	24	24
Exit Temperature (°F)	270.6	270.6
Exit Velocity (ft/s)	51.4	51.4

⁽¹⁾ Coordinates for UTM Zone 16 referenced to Datum NAD27.

Table 4:Modeled Project-Related Chronic Inhalation Risk Estimates for Arsenic, Cadmium, andMercury

	Maximum Proje	ect Related R	kisk Estimates	Kemper County-Wide Average Risk Estimates				
Hazardous Air Pollutant	Maximum Modeled Long-Term Concentration (μg/m³)	Chronic Hazard Quotient	Individual Inhalation Cancer Risk	Long-Term Modeled Concentration from Project ⁽¹⁾ (µg/m ³)	NATA Modeled Kemper County Long- Term Concentration (µg/m ³) ⁽²⁾	Total Modeled Long-Term Kemper County Concentration (μg/m ³)	Chronic Hazard Quotient	Individual Inhalation Cancer Risk
Arsenic	3.7E-05	1.2E-03	1.6E-07	2.3E-06	7.8E-07	3.1E-06	1.0E-04	1.3E-08
Cadmium	5.1E-05	2.6E-03	9.3E-08	3.2E-06	7.2E-07	3.9E-06	1.9E-04	7.0E-09
Mercury - Total	1.4E-05	4.5E-05	not applicable	8.3E-07	8.9E-07	1.7E-06	5.7E-06	not applicable

(1) Values are the average of all receptors (total 2085) in an evenly spaced grid covering all of Kemper County in 1000m increments.

(2) 1999 National Air Toxics Assessment (<u>http://www.epa.gov/ttn/atw/nata1999/tables.html</u>) NATA's National Ambient Background for mercury is 1.50E-03 μg/m³.

Table 5: Modeled Project-Related Acute Inhalation Risk Estimates for Arsenic, Cadmium, and Mercury

Pollutant	Maximum Modeled Concentration (µg/m ³)	Acute Dose Response Value ⁽¹⁾ (µg/m ³)	Acute Hazard Quotient
Arsenic	0.00093	0.19	4.90E-03
Cadmium	0.00130	900	1.45E-06
Mercury - Total 0.00034		1.8	1.90E-04

(1) Minimum value provided in Table 2: Acute Dose-Response Values for Screening Risk Assessments (<u>http://www.epa.gov/ttn/atw/toxsource/summary.html</u>).

Table 6: Maximum Modeled Long-Term Mercury Air Concentrations (µg/m³)

Hazardous Air Pollutant	Maximum Modeled Long- Term Concentration	Grand Bay Lab Ambient Conc. ⁽¹⁾	OLF Mercury Study Average Concentration (2005-2008) ⁽²⁾
Mercury - Elemental	1.22E-05	1.40E-03	1.35E-03
Mercury – RGM	1.35E-06	4.02E-06	3.68E-06
Elemental + RGM	1.36E-05	1.41E-03	1.35E-03

(1) http://www.arl.noaa.gov/documents/reports/Luke_NADP_v2.pdf.

(2) Data provided by Southern Company.

Table 7: Comparison of Modeled Mercury Deposition from the Facility with Measured Deposition (g/m²/year)

Pollutant	Max. Annual Modeled Impact	NADP Mercury Deposition Network ⁽¹⁾	OLF 2005-08 Average Low/High Estimates ⁽²⁾
Mercury - Wet Deposition	3.21E-07	1.68E-05	1.47E-05
Mercury - Dry Deposition	1.71E-06	not available	1.22E-06/2.45E-06
Mercury - Total Deposition	1.90E-06	not available	15.9E-06/17.2E-06

(1) Average concentration at MS22 monitor (Oak Grove, MS) from 2001-2007.

(2) Data for OLF provided by Southern Company. Wet deposition is directly measured and dry deposition has been estimated based on measured concentrations and estimated range of deposition velocity applicable to the surface type in the OLF study area.

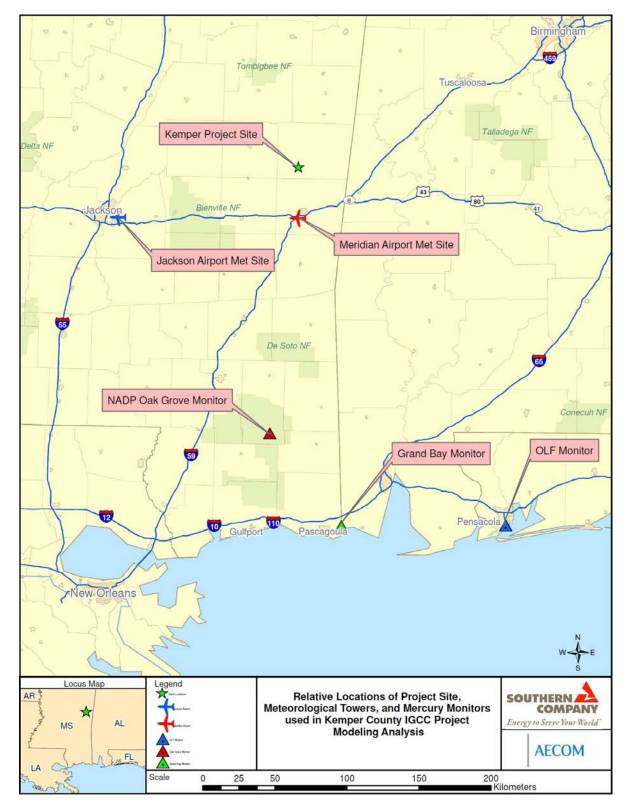


Figure 1: Relative Location of Kemper IGCC Project Site, Meteorological Data Sources, and Ambient Mercury Monitors

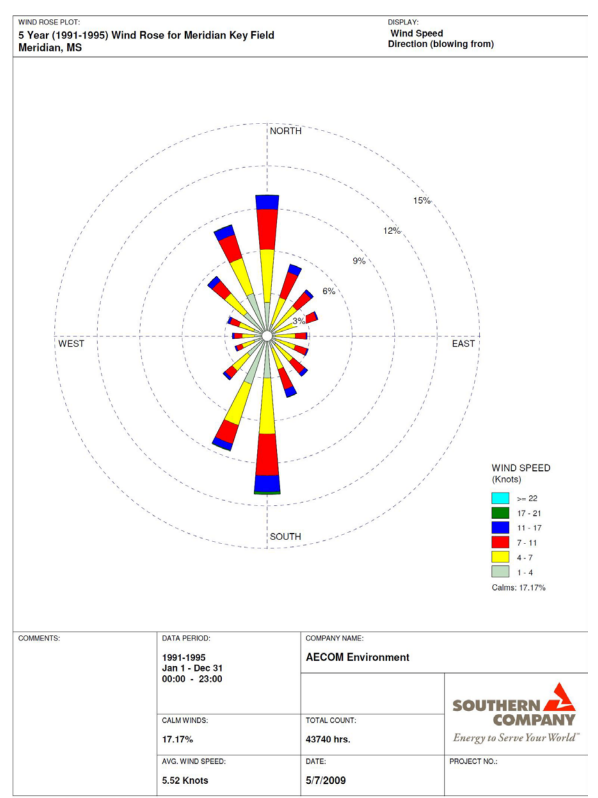


Figure 2: Meridian Key Field 5-year (1991-1995) Wind Rose

WRPLOT View - Lakes Environmental Software

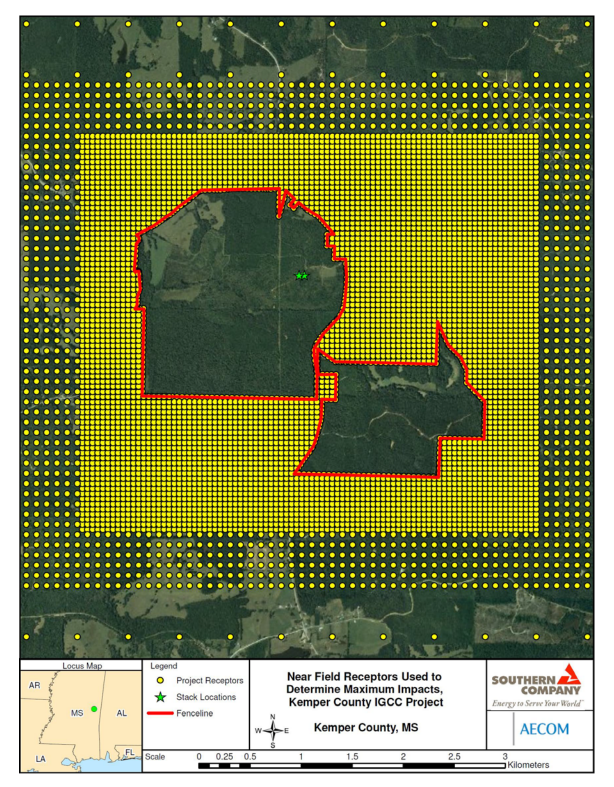
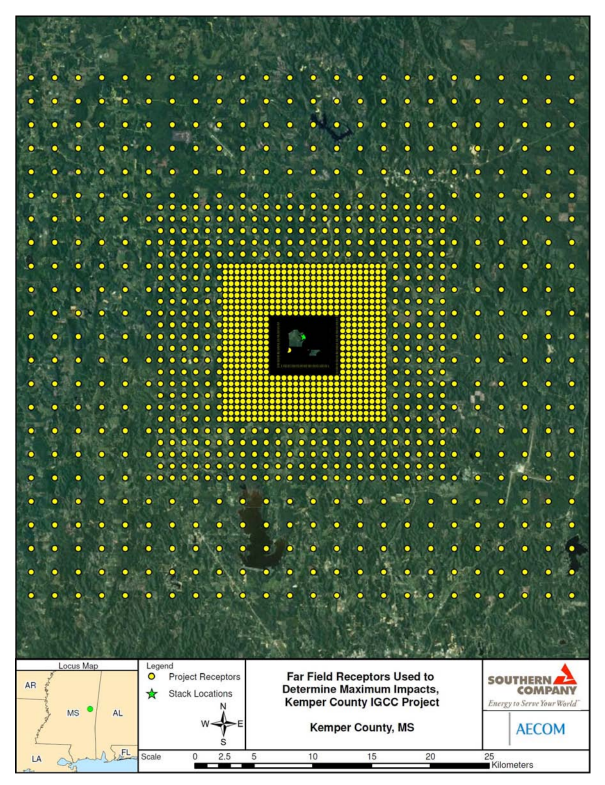
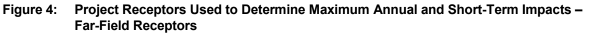


Figure 3: Project Receptors Used to Determine Maximum Annual and Short-Term Impacts – Near-Field Receptors





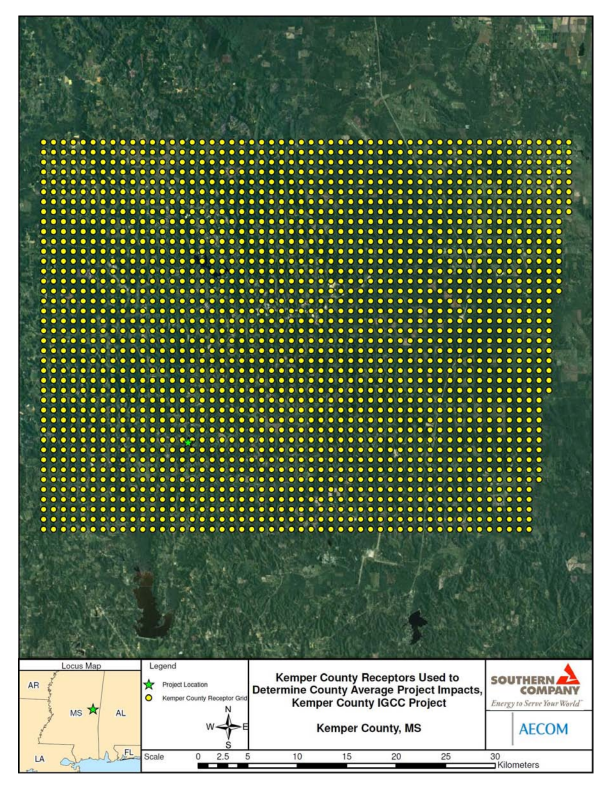


Figure 5: Kemper County Receptor Grid

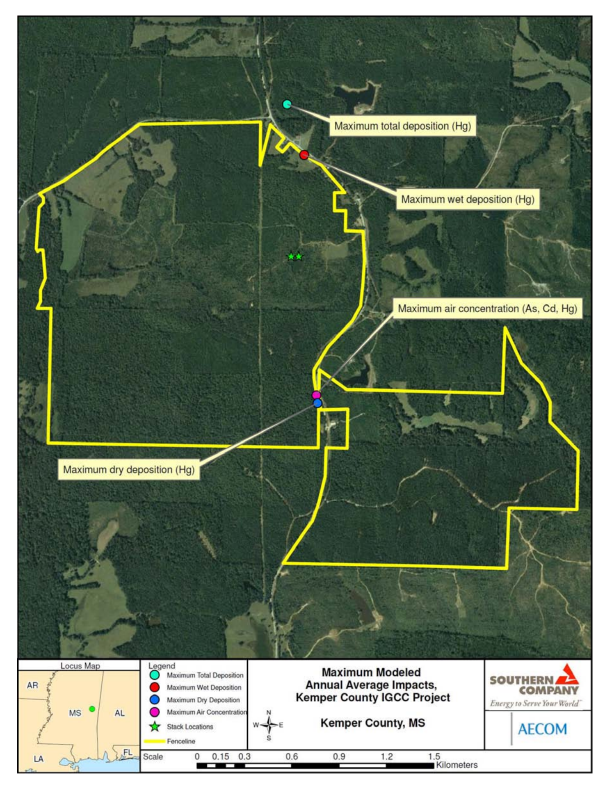


Figure 6: Location of Maximum Air Concentration Impacts and Mercury Deposition

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Contents

1.0	Introduction	.1
2.0	HAP Selection	.1
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List of Tables

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Table 7:	Comparison of Modeled Mercury Deposition from the Facility with Measured Deposition (g/m²/year)9

List of Figures

Figure 1:	Relative Location of Kemper IGCC Project Site, Meteorological Data Sources, and Ambient Mercury Monitors
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Figure 4:	Project Receptors Used to Determine Maximum Annual and Short-Term Impacts – Far-Field Receptors
Figure 5:	Kemper County Receptor Grid14
Figure 6:	Location of Maximum Air Concentration Impacts and Mercury Deposition15

1.0 Introduction

AECOM Environment has conducted a screening level evaluation of the potential effects of hazardous air pollutant (HAP) emissions from the proposed Kemper County Integrated Gasification Combined Cycle (IGCC) stacks.

2.0 HAP Selection

The evaluation considers the HAP emissions associated with continuous firing of syngas in the combustion turbines and natural gas in the associated duct burners. Although the combustion turbines are also capable of combusting natural gas, the fraction of time that natural gas alone would be fired is expected to be small. Siemens turbines were addressed in this assessment. The HAPs emissions for a Siemens turbine are provided in Table 1 for all HAPs with the exception of mercury. Mercury emissions are provided in Table 2.

This assessment addressed two classes of health risk: carcinogenic risk and non-carcinogenic risk for longand short-term (acute and chronic) inhalation of airborne concentrations. For inhalation pathway risk, this screening-level assessment identifies two HAPs, which were selected based on relative emissions and longterm inhalation toxicity. To identify the HAPs to include in the assessment, the HAP emissions were ranked according to relative cancer and chronic non-cancer risk associated with their emissions. The U.S. EPA's Office of Air Quality Planning and Standards (OAQPS) has tabulated dose-response assessments that OAQPS uses for risk assessments of hazardous air pollutants and has made these available on the Technology Transfer Network at http://www.epa.gov/ttn/atw/toxsource/summary.html. For this ranking the Dose-Response Values compiled by U.S. EPA OAQPS (Ref: Prioritized Chronic Dose-Response Values, provided at http://www.epa.gov/ttn/atw/toxsource/table1.pdf) were applied. To estimate the relative contribution of HAP to inhalation cancer risk, each HAP emission rate listed in Table 1 was multiplied the by U.S. EPA's Unit Risk Estimates (URE). In a similar way, the relative percent contribution among HAPs was estimated by dividing the HAP emission rates by their respective chronic inhalation concentration levels (referred to here as Reference Air Concentrations or RACs). The result of this ranking, shown in Table 1, indicates that the two HAPs that contribute most to both cancer and non-cancer inhalation risks are arsenic and cadmium. Together these two HAPs contribute nearly 75% of the inhalation cancer risk associated with the IGCCs and nearly 50% of the chronic non-cancer inhalation risk. The implication is that if the modeled risk for these two HAPs is well below significance levels, it is assured that the combined risk associated with all of the HAPs will also be insignificant.

In addition to risks due to direct inhalation of HAPs, it is well-recognized that the routine ingestion of fish that contain elevated concentrations of mercury can pose non-carcinogenic chronic health risk. Some of the mercury in fish may result from deposition of airborne concentrations onto watersheds and subsequent bioaccumulation through the aquatic food chain. Although the presence of mercury in the ambient environment is a global issue, because mercury is among the HAPs that are known to be emitted from coal combustion, the local dispersion and deposition of this HAP from the IGCCs were also evaluated.

3.0 Evaluation Methods

To assess inhalation risk, AERMOD was applied to estimate the facility's contribution to ambient concentrations at the point of maximum off-site impact, as well as the average concentrations throughout Kemper County. Modeled long-term concentrations of arsenic, cadmium and mercury were then added to U.S. EPA's model estimates of the pre-existing concentrations in Kemper County associated with other local and regional emission sources for these three HAPs as computed by the 1999 National-Scale Air Toxics Assessment (NATA, http://www.epa.gov/ttn/atw/nata1999/tables.html). In modeling long-term concentrations, as

described in more detail below, an average concentration over a 5-year period was computed assuming that the IGCCs ran on syngas continuously at peak load. The level of incremental inhalation cancer and noncancer risk, and combined risk (facility plus NATA), for each HAP were then evaluated. To evaluate the maximum individual lifetime cancer risk the estimated long-term average air concentration was multiplied by the URE (see Table 1). The result of this calculation is the increase in probability of a person contracting cancer after 70 years of exposure. A cancer risk value of less than 1x10⁻⁶ is generally recognized to be insignificant (U.S. EPA, 1991). For chronic non-cancer effects, the long-term average concentration was divided by the respective RAC to form what is referred to as a Hazard Quotient (HQ). A HQ value of less than 1.0 indicates that adverse non-cancer health effects are highly unlikely to occur (U.S. EPA, 2008). To assess potential acute health effects, the maximum modeled 1-hour average concentrations for the IGCCs were divided by minimum acute RAC found in EPA guidance (Acute Dose-Response Values for Screening Risk Assessments (6/12/2007) in http://www.epa.gov/ttn/atw/toxsource/table2.pdf.

In addition to the evaluation of inhalation health risks, the long-term deposition of mercury was also evaluated. Mercury is emitted in three forms, elemental mercury vapor, reactive gaseous mercury (RGM), and particulatebound mercury (PBM). As indicated in Table 2, it is estimated that 90% of the total mercury emitted from the IGCCs is in the form of elemental mercury, 10% RGM and un-guantified trace amounts of PBM. In terms of the contribution to wet or dry deposition, elemental mercury is essentially inert because it has an extremely low solubility and is non-reactive. Therefore, most of the elemental mercury remains airborne and is transported over long distances (U.S. EPA, 1997). PBM is deposited with the particles with which it is incorporated, but in this case the rate of emission is so low that the rate of PBM deposition cannot be quantified. In contrast, RGM is soluble and reactive such that it is subject to both dry and wet deposition. Dry deposition occurs when airborne RGM comes into contact with elements on the earth's surface such as vegetation and water bodies. Wet deposition occurs when precipitation falling through the plume captures soluble RGM vapor. As discussed further below, these deposition processes are simulated by U.S. EPA's AERMOD air quality model. The ambient concentration and deposition of mercury from the IGCC units at the point of maximum impact, and the average over Kemper County, are evaluated by comparing modeled results to measurements taken at two locations along the Gulf coast, and a third located in Oak Grove, MS, where there are long-term mercury monitoring networks (see Figure 1).

4.0 Modeling Approach

U.S. EPA's AERMOD model was applied to estimate air concentrations of arsenic, cadmium and mercury, and deposition of mercury emitted from the IGCC units. Five years of representative meteorological data (1991-1995) were applied, including National Weather Service (NWS) surface and precipitation data from Meridian Key Field in Meridian, MS and upper air data from Jackson International Airport in Jackson, MS. The relative location of the meteorological measurement sites with respect to the project site are provided in Figure 1. Processed, AERMOD-ready meteorological data was provided by the Mississippi Department of Environmental Quality (MDEQ) on their website (http://www.mississippi.gov/frameset.jsp?URL=http%3A%2F%2F %2Fwww.deq.state.ms.us%2FMDEQ.nsf%2Fpage%2FMain_Home%3FOpenDocument). The 5-year wind rose shown in Figure 2 indicates prevailing winds are primarily southerly, but northerly winds are also frequent. In accordance with U. S. EPA guidance, a sensitivity analysis was conducted using both the surface site characteristics (roughness, albedo and Bowen ratio) from the project site and the NWS station site, and the site characteristics that resulted in the greater modeled concentrations were applied. On this basis, the surface characteristics of the NWS site at Meridian were used. Two sets of receptors were used in the analysis. The first set, which was developed for the air quality modeling in support of the project permitting effort, was applied to model the concentration and deposition at the maximum point of impact. This receptor grid consisted of "tight" receptors spaced 50 meters apart along the fence line and extending out 500 meters from the fence-line. Beyond 500 meters, a spacing of 100 meters was used up to one-kilometer from the facility. Between one- and five-kilometers, a spacing of 500 meters was used. Between five- and tenkilometers, a spacing of 1,000 meters was used. Between ten- and twenty-kilometers, a spacing of 2,000

meters was used. For this grid, the near field receptors are shown in Figure 3 and the far field receptors are shown in Figure 4. A second set of receptors comprised of a uniform, 1 km spaced receptor grid across Kemper County, was used to model the Kemper County-wide average for comparison to NATA county-level results. Receptor elevations were developed using Digital Elevation Model (DEM) data from USGS. The Kemper County grid is shown in Figure 5.

Table 3 provides the source parameters used to apply AERMOD. For the IGCC dispersion modeling and RGM deposition analysis, the general modeling procedures and options specified in the current versions of in the AERMOD User's Guide (U.S. EPA 2004), AERMOD Implementation Guide (U.S. EPA 2009), and the Guideline on Air Quality Models (GAQM; U.S. EPA 2005) were followed. To model ambient concentrations of arsenic, cadmium and elemental mercury, it was conservatively assumed that no deposition occurred. Deposition was accounted for in modeling the concentration of RGM. Modeling was conducted in a manner consistent with U.S. EPA guidance and standard practices, including the use of regulatory default options, as appropriate. The building downwash analysis was performed using the most recent version of EPA's Building Profile Input Program (BPIP) (Version 04274) with the plume rise model enhancements (PRIME) building downwash algorithms, consistent with the air quality modeling.

Aerial imagery (2004) was examined to determine the land-use characteristics around the project site for use as input into the model for mercury deposition calculations. The area in the vicinity of the project was found to be heavily forested. As a result, category 4 (forest) was used for the gas dry deposition land-use category for all 36 10° radials in the mercury deposition modeling as described in the Addendum to the AERMOD User's Guide (U.S EPA, 2006).

The transport and mobility of a pollutant are determined by the physical properties of the specific pollutant. For deposition modeling, AERMOD requires several pollutant-specific parameters: (1) diffusivity in air; (2) diffusivity in water; (3) leaf cuticular resistance to lipid uptake; and (4) the Henry's Law constant. The following values of these parameters recommended by U.S. EPA were applied:

- diffusivity in air: 6.0E-2 cm²/sec⁽¹⁾
- diffusivity in water: 5.25E-6 cm²/sec⁽²⁾
- Cuticular resistance: 1.0E7 sec/m⁽¹⁾
- Henry's Law constant: 6.0E-6 pa-m³/mol⁽¹⁾
- Reactivity factor: 1.0⁽¹⁾

¹Source is Wesley, M.L., Doskey, P.V., and Shannon, J.D. Deposition Parameterizations for the Industrial Source Complex (ISC3) Model, Argonne National Laboratory, June, 2002, Table 7 (p.27).

² Source is U.S. EPA's *Human Health Risk* Assessment Protocol for Hazardous Waste Combustion Facilities – [Publication No. EPA 530-R-05-006], September 2005. *Information from companion HHRAP Database – Value for mercuric chloride*. <u>http://www.epa.gov/osw/hazard/tsd/td/combust/riskvol.htm#volume2</u> - scroll down to "<u>Download</u> <u>HHRAP Companion Database (ACCESS)</u>"

5.0 Findings

Figure 6 provides the location of the maximum modeled air concentrations for the modeled HAPs and the location of the highest RGM deposition relative to the facility. The maximum air concentration impacts, along with the maximum dry deposition, were found to be along the southeast fence-line of the power block portion of the facility. The maximum wet and total RGM deposition was found to occur along, and just past, the northeast fence-line of the power block, directly to the north of the IGCC stacks.

Table 4 provides inhalation risk estimates associated with arsenic, cadmium and mercury emitted from the facility and the combined county-wide risk of these HAPs, incorporating the 1999 NATA Kemper County modeling results. The maximum cancer risk, which is associated with arsenic, is 2.0×10^{-7} at the point of maximum, and the county-wide-average risk, including the proposed IGCC units is only about 1.7×10^{-8} . These levels are below the EPA's recommended significance level of 1×10^{-6} . The hazard quotients for each of the three HAPs are also insignificant with the highest incremental HQ, which occurs for cadmium, of 3.2×10^{-3} compared to a significance level of 1.0. The corresponding Kemper County HQ for cadmium is about 2.6×10^{-4} . The results of acute inhalation assessment, provided in Table 5, indicate that arsenic is the HAP with the largest acute hazard quotient with a value of about 5.6×10^{-3} . This analysis demonstrates that the proposed IGCC units will not cause or contribute significantly to acute or chronic health effects and cancer risk.

The analyses for mercury are provided in Tables 6 and 7. Table 6 compares maximum modeled long-term air concentrations of elemental mercury and RGM with available measurements from two measurement sites located to the south near the Gulf Coast. The long-term concentration of elemental mercury is two orders of magnitude less than the measured ambient concentrations, and the maximum modeled RGM concentration is less than half of the ambient concentrations. Thus, the combined maximum long-term modeled concentration of elemental mercury and RGM is only about one hundredth of the measured ambient concentrations. This finding is consistent with NATA which specifies a U.S. background mercury concentration associated with global transport of $1.5 \times 10^{-3} \,\mu g/m^3$. Table 7 provides a comparison of the maximum modeled RGM deposition from the facility with values developed from measurements from the OLF study and the NADP Mercury Deposition Network. The maximum long-term wet deposition is one to two orders of magnitude less than measurements from the OLF study. The result is that the total modeled mercury deposition is only about one seventh of total ambient deposition. The deposition analysis therefore demonstrates that the project will not contribute substantially to the ambient concentration or deposition of mercury in the vicinity.

6.0 Summary

An assessment of the HAP emissions from the proposed IGCC units was conducted using the conservative assumption that the IGCC units will operate continuously at 100% load on syngas for five years. Modeling of concentrations and RGM deposition was conducted with AERMOD, a U.S. EPA designed model that provides conservative estimates of air quality impacts. The results of the assessment indicate that no HAP emitted from the project will result in, or contribute to, human health risk due to inhalation. In addition, the ambient concentrations of mercury resulting from the IGCC units were modeled to be orders of magnitude less than ambient concentrations, and the modeled deposition of mercury was less than has been measured at nearby study areas.

7.0 References

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Wesley, M.L., Doskey, P.V., and Shannon, J.D. Deposition Parameterizations for the Industrial Source Complex (ISC3) Model, Argonne National Laboratory, June, 2002.

	Maximum Short-Term Emissions ^(1,3) (Ib/hour)	Cancer Unit Risk Estimate ⁽⁴⁾ (1/µg/m ³)	Percent of Cancer Risk	Reference Air Concentration ⁽⁴⁾ (µg/m3)	Percent of Chronic Hazard Index
Volatile Organic Compounds					
Acetaldehyde	1.79E-02	2.20E-06	0.0%	9.00	0.1%
Acrolein	1.58E-03	NA		0.02	4.0%
Benzene	1.91E-02	7.80E-06	0.2%	30.00	0.0%
Ethylbenzene	6.43E-03	NA		1000.00	0.0%
Formaldehyde	8.61E-02	5.50E-09	0.0%	9.80	0.4%
Toluene	1.92E-02	NA		5000.00	0.0%
Xylene	1.83E-02	NA		100.00	0.0%
Polycyclic Organic Matter (POM)					
Polycyclic Aromatic	1 225 04	1 105 02	0.29/	200.00	0.00/
Hydrocarbons (PAH) 2-Methylnapthalene	1.33E-04 1.14E-03	1.10E-03 NA	0.2%	200.00 NA	0.0%
Acenapthylene	8.25E-05	NA		NA NA	
Benzo(a)anthracene	7.30E-06	1.10E-04	0.0%	NA	
Benzo(e)pyrene	1.75E-05	NA		NA	
Benzo(g,h,i)perylene	3.02E-05	NA		NA	
Napthalene	1.45E-03	3.40E-05	0.1%	3.00	0.0%
Metals	1.102.00	0.102.00	0.170	0.00	0.070
Antimony	1.24E-02	NA		NA	
Arsenic	9.52E-03	4.30E-03	45.7%	0.03	16.0%
Beryllium	2.92E-03	2.40E-03	7.8%	0.02	7.4%
Cadmium	1.33E-02	1.80E-03	26.8%	0.02	33.6%
Chromium VI (2)	1.45E-03	1.20E-02	19.4%	0.10	0.7%
Cobalt	2.57E-03	NA		0.10	1.3%
Lead	1.27E-02	NA		0.15	4.3%
Manganese	1.36E-02	NA		0.05	13.8%
Mercury (total)	3.67E-03	NA		0.30	0.6%
Elemental Mercury	3.31E-03	NA		NA	
Reactive Gaseous Mercury	3.67E-04	NA		NA	
Particulate-bound Mercury	Trace	NA		NA	
Nickel	1.78E-02	NA		0.09	10.0%
Phosphorous	1.08E-02	NA		0.07	7.8%
Selenium	1.36E-02	NA		NA	
Inorganic Compounds					
Carbon Disulfide	1.43E-01	NA		700.00	0.0%

Table 1: Estimates of HAP Emissions for Each of Two IGCC Stacks for Siemens Turbines and **Ranking in Terms of Inhalation Risk**

(1) Emission rates based on emission factors from "A Study of Toxic Emissions from a Coal-Fired Gasification Plant", Radian Corporation, December 1995.

(2) 12% of total chromium emissions characterized as hexavalent for coal combustion per U.S. EPA 2005 National Emissions Inventory Data & Documentation (<u>http://www.epa.gov/ttn/chief/net/2005inventory.html</u>). Estimates are based on full load operating scenarios with Duct Burner firing.

(3)

(4) Source: Prioritized Chronic Dose-Response Values: http://www.epa.gov/ttn/atw/toxsource/table1.pdf.

	Average Short- Term Syngas (lb/hour)	Average Annual (lb/year)
Total Mercury	3.67E-03	32.18
Elemental (90% of Total)	3.31E-03	28.96
RGM (10% of Total)	3.67E-04	3.22
Particle-Bound (Only trace amounts)	Trace	Trace

Table 2: Estimate of Mercury Emissions for Each of Two IGCC Stacks for Siemens Turbines

Table 3: IGCC Source Parameters

Parameter	Value		
	CC1A	CC2A	
UTM Coordinate East (m) (1)	335175.01	335223.77	
UTM Coordinate North (m) (1)	3614077.76	3614076.92	
Stack Base Elevation (ft)	474	474	
Stack Height (ft)	325	325	
Stack Diameter (ft)	24	24	
Exit Temperature (°F)	213.0	213.0	
Exit Velocity (ft/s)	46	46	

⁽¹⁾ Coordinates for UTM Zone 16 referenced to Datum NAD27.

Table 4: Modeled Project-Related Chronic Inhalation Risk Estimates for Arsenic, Cadmium, and Mercury

	Maximum Project Related Risk Estimates		Kemper County-Wide Average Risk Estimates					
Hazardous Air Pollutant	Maximum Modeled Long-Term Concentration (μg/m³)	Chronic Hazard Quotient	Individual Inhalation Cancer Risk	Average ⁽¹⁾ Long-Term Modeled Concentration from Project (µg/m ³)	NATA ⁽²⁾ Modeled Kemper County Long- Term Concentration (µg/m ³)	Total Modeled Long-Term Kemper County Concentration (μg/m ³)	Chronic Hazard Quotient	Individual Inhalation Cancer Risk
Arsenic	4.62E-05	1.54E-03	1.99E-07	3.24E-06	7.83E-07	4.02E-06	1.34E-04	1.73E-08
Cadmium	6.46E-05	3.23E-03	1.16E-07	4.53E-06	7.22E-07	5.25E-06	2.63E-04	9.45E-09
Mercury - Total	1.78E-05	5.93E-05	not applicable	1.24E-06	8.91E-07	2.13E-06	7.10E-06	not applicable

(1) Values are the average of all receptors (total 2085) in an evenly spaced grid covering all of Kemper County in 1000m increments.

(2) 1999 National Air Toxics Assessment (<u>http://www.epa.gov/ttn/atw/nata1999/tables.html</u>) NATA's National Ambient Background for mercury is 1.50E-03 μg/m³.

Table 5: Modeled Project-Related Acute Inhalation Risk Estimates for Arsenic, Cadmium, and Mercury

Pollutant	Maximum Modeled Concentration (μg/m³)	Acute Dose Response Value ⁽¹⁾ (μg/m ³)	Acute Hazard Quotient
Arsenic	0.00106	0.19	5.58E-03
Cadmium	0.00148	900.0	1.65E-06
Mercury - Elemental	0.00041	1.8	2.27E-04

(1) Minimum value provided in Table 2: Acute Dose-Response Values for Screening Risk Assessments (<u>http://www.epa.gov/ttn/atw/toxsource/summary.html</u>).

Table 6: Maximum Modeled Long-Term Mercury Air Concentrations (µg/m³)

Hazardous Air Pollutant	Maximum Modeled Long- Term Concentration	Grand Bay Lab Ambient Conc. ⁽¹⁾	OLF Mercury Study Average Concentration (2005-2008) ⁽²⁾
Mercury - Elemental	1.60E-05	1.40E-03	1.35E-03
Mercury – RGM	1.78E-06	4.02E-06	3.68E-06
Elemental + RGM	1.78E-05	1.41E-03	1.35E-03

(1) http://www.arl.noaa.gov/documents/reports/Luke_NADP_v2.pdf.

(2) Data provided by Southern Company.

Table 7: Comparison of Modeled Mercury Deposition from the Facility with Measured Deposition (g/m²/year)

Pollutant	Max. Annual Modeled Impact	NADP Mercury Deposition Network ⁽¹⁾	OLF 2005-08 Average Low/High Estimates ⁽²⁾
Mercury - Wet Deposition	3.44E-07	1.68E-05	1.47E-05
Mercury - Dry Deposition	2.24E-06	not available	1.22E-06/2.45E-06
Mercury - Total Deposition	2.46E-06	not available	15.9E-06/17.2E-06

(1) Average concentration at MS22 monitor (Oak Grove, MS) from 2001-2007.

(2) Data for OLF provided by Southern Company. Wet deposition is directly measured and dry deposition has been estimated based on measured concentrations and estimated range of deposition velocity applicable to the surface type in the OLF study area.

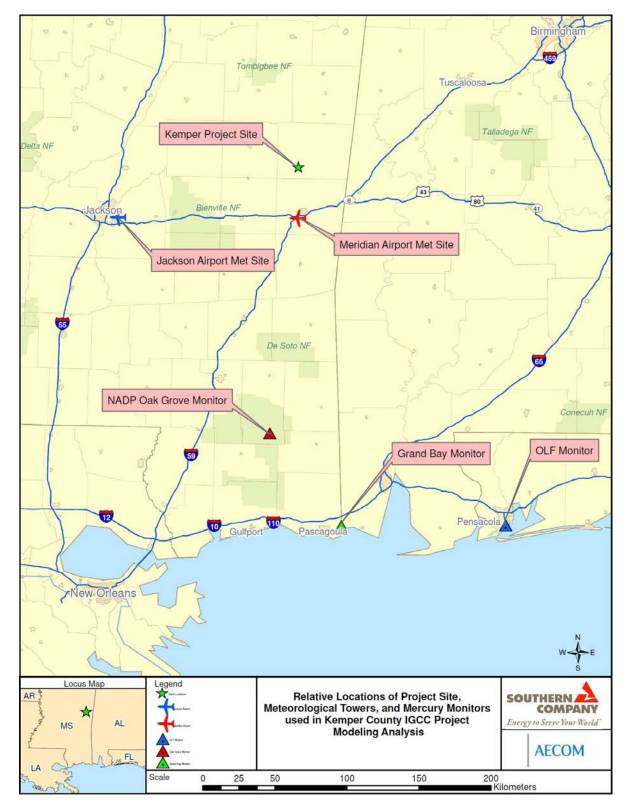


Figure 1: Relative Location of Kemper IGCC Project Site, Meteorological Data Sources, and Ambient Mercury Monitors

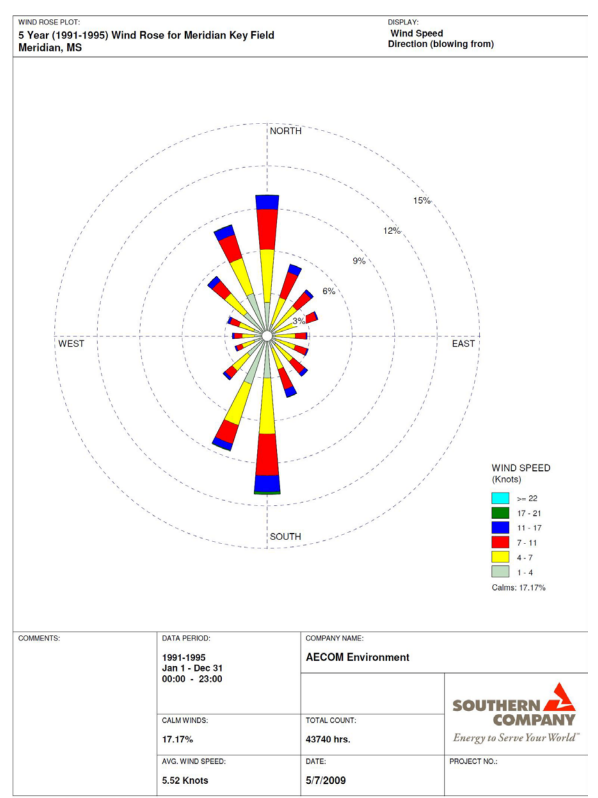


Figure 2: Meridian Key Field 5-year (1991-1995) Wind Rose

WRPLOT View - Lakes Environmental Software

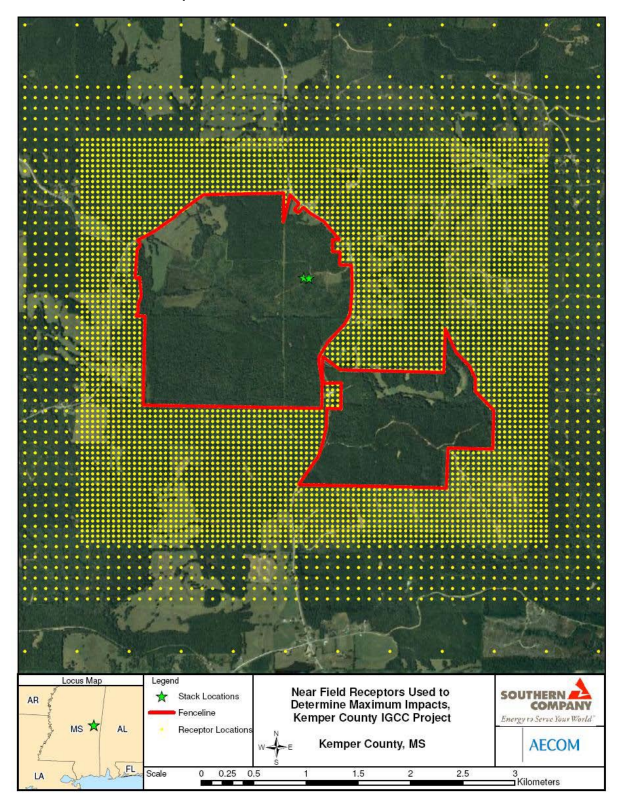


Figure 3: Project Receptors Used to Determine Maximum Annual and Short-Term Impacts – Near-Field Receptors

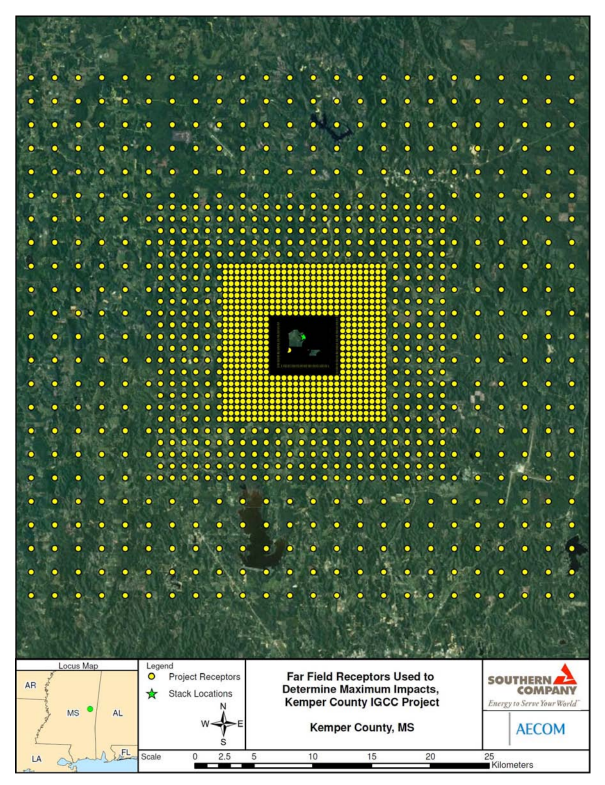


Figure 4: Project Receptors Used to Determine Maximum Annual and Short-Term Impacts – Far-Field Receptors

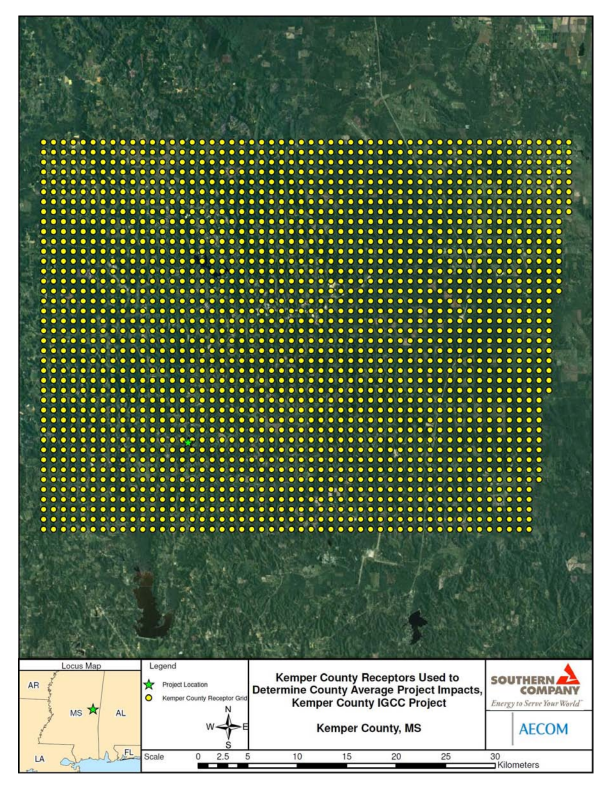


Figure 5: Kemper County Receptor Grid

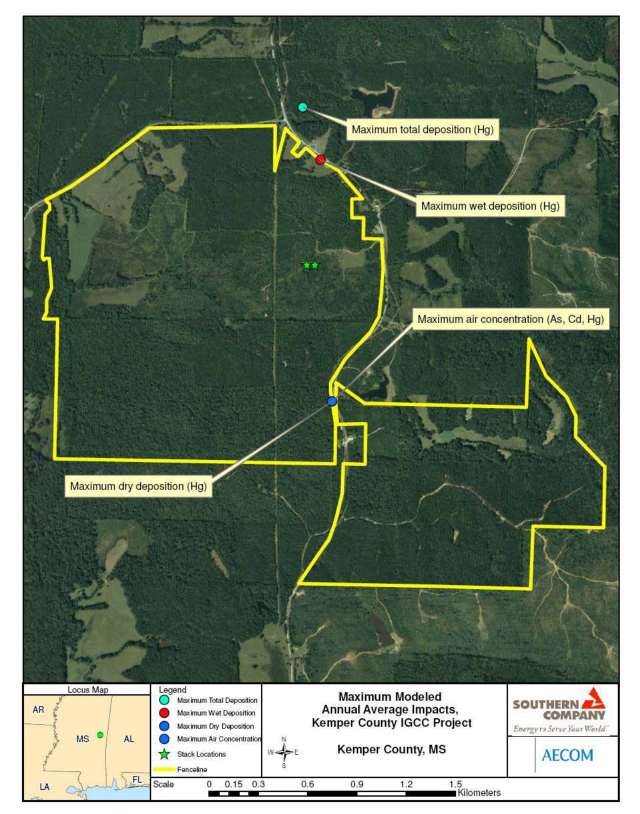


Figure 6: Location of Maximum Air Concentration Impacts and Mercury Deposition