



# GEORGIA

DEPARTMENT OF NATURAL RESOURCES

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## ENVIRONMENTAL PROTECTION DIVISION

**Quality Assurance Project Plan  
for the Georgia Ambient Air Monitoring Program  
Ethylene Oxide**

**Category II**

March 2020  
Revision 1.1

**Air Protection Branch  
Ambient Air Monitoring Program  
4244 International Parkway, Suite 120  
Atlanta, GA 30354**

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### *Acronyms and Abbreviations*

|                   |  |
|-------------------|--|
| AAMP              | Ambient Air Monitoring Program                 |
| APB               | Air Protection Branch                          |
| ASTM              | American Society for Testing and Materials     |
| ATMP              | Air Toxics Monitoring Program                  |
| °C                | Degrees Celsius                                |
| CAA               | Clean Air Act                                  |
| CFR               | Code of Federal Regulations                    |
| COC               | Chain of Custody                               |
| DQA               | Data Quality Assessment                        |
| DQI               | Data Quality Indicator                         |
| DQO               | Data Quality Objectives                        |
| EPA               | Environmental Protection Agency                |
| EPD               | Environmental Protection Division              |
| ESMB              | Extraction Solvent Method Blank                |
| GA EPD            | Georgia Environmental Protection Division      |
| GC                | Gas Chromatography                             |
| GC/MS             | Gas Chromatography/Mass Spectrometry           |
| HAPs              | Hazardous Air Pollutants                       |
| IB                | Instrument Blank                               |
| ICAL              | Initial Calibration                            |
| ICB               | Initial Calibration Blank                      |
| IO                | Inorganic                                      |
| IS                | Internal Standards                             |
| ISO               | International Organization for Standardization |
| K                 | Kelvin   |
| kPa               | Kilopascal                                     |
| LCS               | Laboratory Control Sample                      |
| LCSD              | Laboratory Control Sample Duplicate            |
| LIMS              | Laboratory Information Management System       |
| MB                | Method Blank                                   |
| MDL               | Method Detection Limit                         |
| µg                | Micrograms                                     |
| µg/m <sup>3</sup> | Micrograms per Cubic Meter                     |
| µg/mL             | Micrograms per Milliliter                      |
| MS                | Matrix Spike                                   |
| MSD               | Matrix Spike Duplicate                         |
| MQO               | Measurement Quality Objectives                 |
| MSA               | Metropolitan Statistical Area                  |
| NATA              | National Air Toxics Assessment                 |
| NATTS             | National Air Toxics Trends Stations            |
| NIST              | National Institute of Standards and Technology |
| OAQPS             | Office of Air Quality Planning and Standards   |
| PAMS              | Photochemical Assessment Monitoring Station    |

|        |   |
|--------|---|
| PPB    | Parts per Billion                             |
| PPBV   | Parts per Billion Volume                      |
| PQAO   | Primary Quality Assurance Organization        |
| QC     | Quality Control                               |
| QA     | Quality Assurance                             |
| QAPP   | Quality Assurance Project Plan                |
| r      | Correlation Coefficient                       |
| RPD    | Relative Percent Difference                   |
| RSD    | Relative Standard Deviation                   |
| RRF    | Relative Response Factor                      |
| RRT    | Relative Retention Time                       |
| RT     | Retention Time                                |
| SB     | Solvent Blank                                 |
| SLAMS  | State and Local Monitoring Stations           |
| SOP    | Standard Operating Procedure                  |
| TAD    | Technical Assistance Document                 |
| TM     | Trademark                                     |
| TO     | Toxic Organic                                 |
| UATS   | Urban Air Toxics Strategy                     |
| US EPA | United States Environmental Protection Agency |
| VOC    | Volatile Organic Compound                     |

## 1.0 Quality Assurance Project Plan Identification Approval

The attached Category II *Quality Assurance Project Plan for the Georgia Ambient Air Monitoring Program for Ethylene Oxide* is hereby recommended for approval and commits the Georgia Environmental Protection Division (GA EPD) to follow the elements described within.

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Designated Approving Official, US EPA Region 4 – Air and Radiation  
Division

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### **3.0 Distribution List**

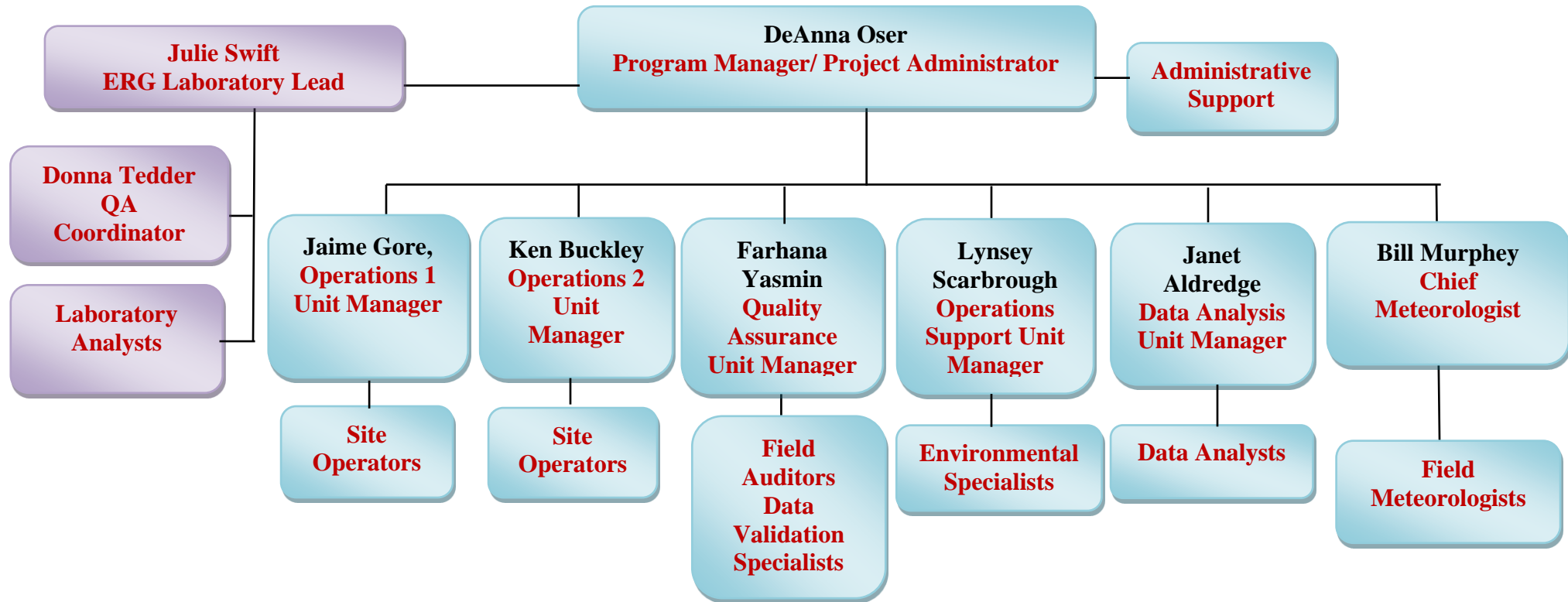
This section is not required for a Category II Quality Assurance Project Plan (QAPP).

### **4.0 Project/Task Organization**

The Georgia Ambient Air Monitoring Program (GA AAMP) and the Eastern Research Group Laboratory (ERG) have important roles in developing and implementing this ethylene oxide study. GA AAMP is responsible for taking this information and developing a study to meet the data quality requirements. ERG is the contract laboratory for the U.S. Environmental Protection Agency (EPA) for air toxics programs such as the National Air Toxics Trends (NATTS) sites. They are the laboratory utilized by EPA for previous ethylene oxide studies. Therefore, the laboratory quality assurance requirements are sufficient for the purposes of this study. For detailed information on the ERG Lab, see the ERG's *Support for the EPA National Monitoring Programs (UATMP, NATTS, CSATAM, PAMS, and NMOC Support) QAPP*, dated March 2019 (Laboratory Attachment of this document).

To make the best use of available resources and to meet timelines for collection and analysis of this study, the flow of information and samples must be optimally organized. The deployment and operation of the project is a shared responsibility among all the involved parties. This section describes the roles of all parties and establishes the lines of authority, communication and reporting, with the goal of facilitating a smoothly operated project. Figure 1 represents the division of function in the organization of the GA AAMP (blue blocks) and ERG Lab (purple blocks). The following information lists the specific responsibilities of each position.

Figure 1. GA AAMP Project Organizational Chart



## **4.1 Program Manager/Project Administrator**

Under supervision of the GA Air Protection Branch (APB) Chief, the Program Manager of GA AAMP is the Project Administrator for all the ambient air monitoring projects. He/she has the overall responsibilities for managing all aspects of the GA AAMP according to policy. Ultimately, the Program Manager/Project Administrator is responsible for establishing QA policy and for resolving QA issues identified through the QA program. The major responsibilities of the Program Manager/Project Administrator include, but are not limited to:

- Serving as a public relations contact for monitoring activities with this project
- Reviewing and maintaining budgets and milestones for GA AAMP
- Ensuring this study meets EPA quality assurance requirements
- Communicating with the ERG Laboratory Lead on issues related to routine sample analysis and related QA activities
- Reviewing and approving QAPPs and Standard Operating Procedures (SOPs) for the GA AAMP
- Managing GA AAMP's documents and records

## **4.2 Quality Assurance (QA) Unit**

### **4.2.1 Quality Assurance Unit Manager**

The QA Unit Manager is the delegated supervisor of the GA AAMP's QA Program for field and data handling activities. He/she has direct access to the Project Administrator (GA AAMP Manager) on all matters pertaining to quality assurance activities regarding field monitoring, sampling, measuring operations, and data handling procedures. His/her responsibilities are detailed below:

- Implementing GA AAMP's quality system in accordance with EPA's and GA EPD's QA policies within the project
- Reviewing and approving GA AAMP SOPs
- Managing data validation of air quality monitoring data
- Reviewing field audit reports
- Ensuring that reviews and audits are scheduled and completed
- Performing data verification of the data for this study

The QA Unit Manager has the authority to carry out these responsibilities and to bring to the attention of the Program Manager/Project Administrator any issues associated with these responsibilities.

### **4.2.2 Field Auditor**

The Field Auditor is responsible for:

- Scheduling and conducting field audits
- Assisting QA Unit Manager in developing and updating QAPPs
- Preparing and finalizing field audit reports

The Field Auditor has the authority to carry out these responsibilities and to bring to the attention of the QA Unit Manager any issues related to these responsibilities.

#### **4.2.3 Data Validation Specialist**

The Data Validation Specialist is responsible for:

- Preparing and updating SOPs for data review and validation activities
- Performing review to ensure that the ambient air monitoring data are validated in accordance with GA AAMP's data validation SOPs

### **4.3 Operations Units**

#### **4.3.1 Operations Unit Managers**

GA AAMP has two different Operations Units due to the heavy workload on field activities. The Operations Unit Managers are the delegated supervisors of the GA AAMP for the field monitoring and sampling operations, which include the QC activities that are implemented as part of routine data collection activities. Responsibilities of the Operations Unit Managers include:

- Supervising personnel in Operations Unit
- Establishing, operating, and maintaining all ambient air monitoring locations
- Developing the monitoring plan for this study
- Understanding GA AAMP QA policy and ensuring the Site Operators understand and follow the policy
- Assisting in resolution of technical problems

#### **4.3.2 Site Operators**

Under the supervision of the Operations Unit Managers, the Site Operators responsibilities include:

- Operating the air monitoring samplers following all the manufacturers' specifications, GA AAMP's SOPs, and this QAPP
- Maintaining a schedule of sample collection and shipments
- Verifying that all required QC activities are performed and that measurement quality standards are met as required in this QAPP
- Documenting and reporting all problems and corrective actions to the Operations Unit Managers

### **4.4 Operations Support Unit**

#### **4.4.1 Operations Support Unit Manager**

Under supervision of the GA AAMP Manager, the Operations Support Unit Manager is responsible for:

- Directing the activities of staff members responsible for overseeing the functions of GA AAMP Workshop (including inventory, testing of new equipment, maintenance and repair)

- Coordinating with ERG Lab for sample media pickup and sample delivery
- Updating and writing SOPs for new equipment added to the GA AAMP
- Budgeting for the Operations Units in managing purchasing and equipment procurement related to the field monitoring and sampling activities

#### **4.4.2 Environmental Specialist**

The Environmental Specialist in the Operations Support Unit assists Operations Support Unit Manager in his/her activities including:

- GA AAMP Workshop activities including testing of new equipment, maintenance and repair, and preventative maintenance activities
- Coordinating with ERG Lab for sample media pickup and sample delivery
- Updating and writing SOPs for new equipment added to the GA AAMP

#### **4.5 Data Analysis Unit**

##### **4.5.1 Data Analysis Unit Manager**

Under supervision of the GA AAMP Manager, the Data Analysis Unit Manager is responsible for:

- Supervising personnel in Data Analysis Unit
- Managing data analysis of this study
- Composing and updating GA AAMP's QAPPs
- Managing, reviewing and editing SOPs for the GA AAMP

##### **4.5.2 Data Analyst**

Under the supervision of the Data Analysis Manager, the Data Analyst's responsibilities include:

- Assisting in data analysis of this study
- Assisting in preparation of QAPPs for the GA AAMP
- Assisting in preparation of SOPs for the GA AAMP

#### **4.6 Meteorological Unit**

##### **4.6.1 Chief Meteorologist**

The Chief Meteorologist supervises the Meteorological Unit by:

- Supervising, training, and evaluating personnel in the Meteorological Unit
- Evaluating wind rose data in relation to monitoring locations

##### **4.6.2 Field Meteorologist**

The Field Meteorologist is responsible for:

- Evaluating wind rose data in relation to monitoring locations

## **4.7 Eastern Research Group Laboratory**

While GA AAMP handles all ambient air monitoring field activities, the ERG Lab handles the laboratory supplies, sample analysis, and laboratory QA/QC. The ERG Lab forwards the analytical data to GA AAMP for further data processing, review, and data validation. The ERG Lab is a contract laboratory and is utilized by US EPA for National Air Toxic Trends Site (NATTS) analysis, which includes the TO-15 analysis, operating under a QAPP approved by EPA Office of Air Quality Planning and Support (OAQPS). Therefore, the quality assurance activities of the ERG Lab are presumed sufficient. For more description of the ERG Lab, see *Support for the EPA National Monitoring Programs (UATMP, NATTS, CSATAM, PAMS, and NMOC Support) QAPP* (Laboratory Attachment of this document).

### **4.7.1 Laboratory Lead**

The Laboratory Lead has overall responsibility for managing all aspects of the ethylene oxide analyses for the ERG Lab. Ultimately, the Laboratory Lead is responsible for establishing the QA policy and for resolving QA issues identified through the Laboratory QA program. The laboratory operates under an EPA approved QAPP for TO-15 analysis for volatile organic compounds.

### **4.7.2 QA Coordinator**

The ERG Lab QA Coordinator has responsibility for ensuring that the ERG Lab follows the ERG Lab's QAPP, as approved by EPA.

## **5.0 Problem Definition/Background**

The National Air Toxics Assessment (NATA), which is updated approximately every three years, provides estimates of the risk of cancer and other serious health effects from inhaling air contaminated with toxic pollutants from large and small industrial sources, from on- and off-road mobile sources, and from natural sources such as fires. The latest available NATA report uses the 2014 National Emission Inventory (NEI), and in August of 2018, the NATA presented the updated estimated cancer risks at the census tract level. With this updated information, the NATA report identifies 18 areas of the U.S. that potentially have elevated long-term (chronic) cancer risks due to ethylene oxide emissions from stationary industrial sources. The Atlanta-Sandy Springs-Roswell Metropolitan Statistical Area (Atlanta MSA) was identified as one of these areas. The main use of ethylene oxide includes manufacture of ethylene glycol (antifreeze), solvents, detergents, adhesives and other products. Also, ethylene oxide is used as a fumigant and a sterilant for surgical equipment and plastic devices.

With the Atlanta MSA being one of the areas identified to have elevated risk in the NATA report, the Georgia Ambient Air Monitoring Program (GA AAMP) will conduct a study to characterize ethylene oxide concentrations in the ambient air. The GA AAMP will begin a study of ambient air levels of ethylene oxide concentrations as of September 2019. The plan for this study is that samples will be collected for approximately six months; however, if significant changes are seen in the data, the study may be extended for further characterization.

This ambient air monitoring study will yield data of sufficient quality that will allow a preliminary assessment of any potential ethylene oxide found at the monitoring sites. The preliminary assessment will be used to determine subsequent steps that may include considering longer-term monitoring where initial data are inclusive and additional information is needed to better characterize the ethylene oxide concentrations.

This QAPP describes the quality system developed, implemented and maintained by GA AAMP for the collection of air samples; the data quality assessment; the data validation; and the reporting of results to GA EPD's website (<https://epd.georgia.gov/ethylene-oxide-information>). The GA AAMP of the GA EPD acts as primary quality assurance organization (PQAO) in charge of monitoring ethylene oxide data.

## **6.0 Project/Task Description**

This QAPP was developed to ensure that GA AAMP has a quality program to characterize ethylene oxide concentrations in the ambient air. The plan for this study is that samples will be collected for approximately six months. The ethylene oxide monitoring study was developed to ensure consistent data quality is sufficient to characterize the ethylene oxide concentrations in the areas monitored. This study data will be posted to the GA EPD's website (<https://epd.georgia.gov/ethylene-oxide-information>).

The monitoring objectives for this study include the following specific aims:

- Characterizing ethylene oxide concentrations in the ambient air within approximately ¼ mile of two facilities (Sterigenics, Cobb County, Georgia and Becton Dickinson-Covington, Newton County, Georgia)
- Providing background concentrations for comparison at two previously established GA AAMP network sites, South DeKalb (13-089-0002) and the General Coffee monitoring station (13-069-0002)
- Providing quality data for risk characterization by other agencies
- Additional sample locations may be added as resources allow, following the methodologies outlined in this document, as applicable

Before the study begins in September, the GA AAMP began preliminary sampling for ethylene oxide at the South DeKalb (13-089-0002) National Air Toxics Trends Site (NATTS) in June of 2019 to gain an understanding of collection and analytical methods of the samples.

This study will utilize passive samplers for the measurement of ethylene oxide in the Atlanta area. For each day that samples are collected in the Covington and Cobb County areas, a sample will also be collected at the South DeKalb site utilizing the same passive sampling equipment. This comparison will provide information on the variability in the ethylene oxide concentrations in an urban area which is not influenced by the two facilities discussed above.

During the study, approximately three qualitative samples will be taken at the South DeKalb site utilizing the passive sampling system as well as the ATEC system that was used for the initial

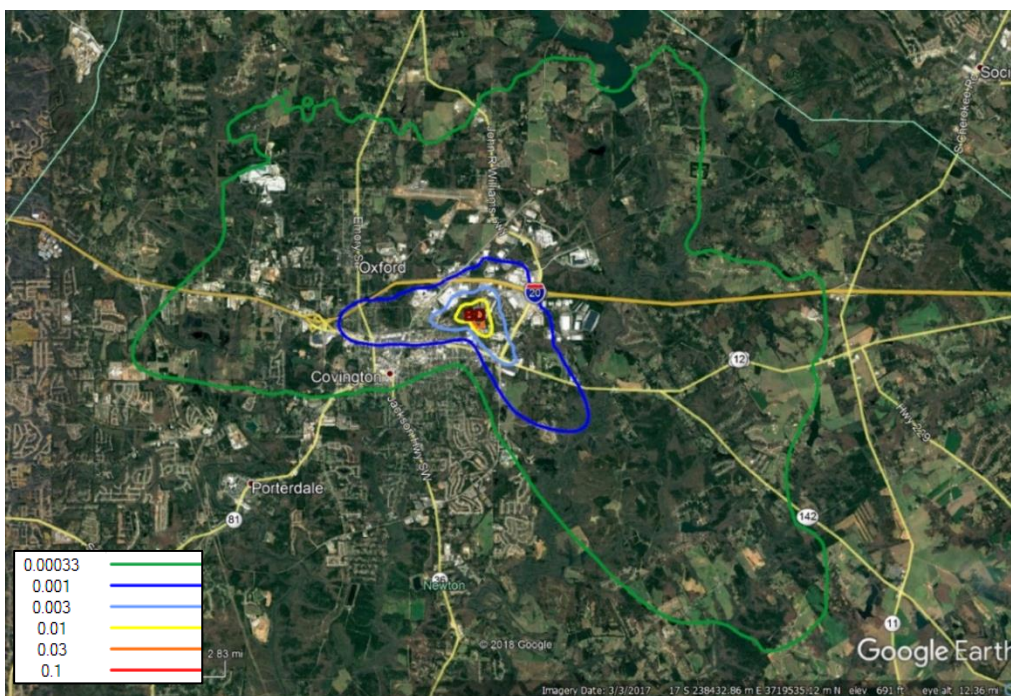


measurements prior to the commencement of this study. In addition, due to its proximity to the South DeKalb site and Interstate 285, the GA AAMP will also collect approximately three ethylene oxide samples with the VOCs canister collection at the Near Road-285 (NR-285) site (13-089-0003) for qualitative comparison to the data collected at the South DeKalb site. The following figure shows the proximity of the two sites. This comparison may provide insight on the contribution of mobile sources to the ethylene oxide concentration measured at the South DeKalb site.



**Figure 2. Location of South DeKalb and NR-285 Sites**

To determine the ambient monitoring sites near Becton-Dickinson in Covington, GA (Figure 3 and Figure 4) and Sterigenics in Cobb County, GA (Figure 6 and Figure 7), the GA AAMP considered the modeled emission data which was generated by the Planning and Support Program of GA EPD. Dispersion models of the ethylene oxide emissions data from these two facilities had been conducted to determine concentrations of ethylene oxide around each facility. These models are shown in the following figures. The modeled values are shown in micrograms per cubic meter ( $\mu\text{g}/\text{m}^3$ ). Based on previous ethylene oxide monitoring conducted by EPA, the determination was made to characterize the ethylene oxide concentrations within approximately  $\frac{1}{4}$  mile of the facilities and to qualitatively determine the gradient (change in concentration) within approximately 1 mile of each facility. Therefore, each model was overlaid with  $\frac{1}{4}$  mile,  $\frac{1}{2}$  mile and 1 mile radius measurements around each facility (Figure 5 and Figure 8).

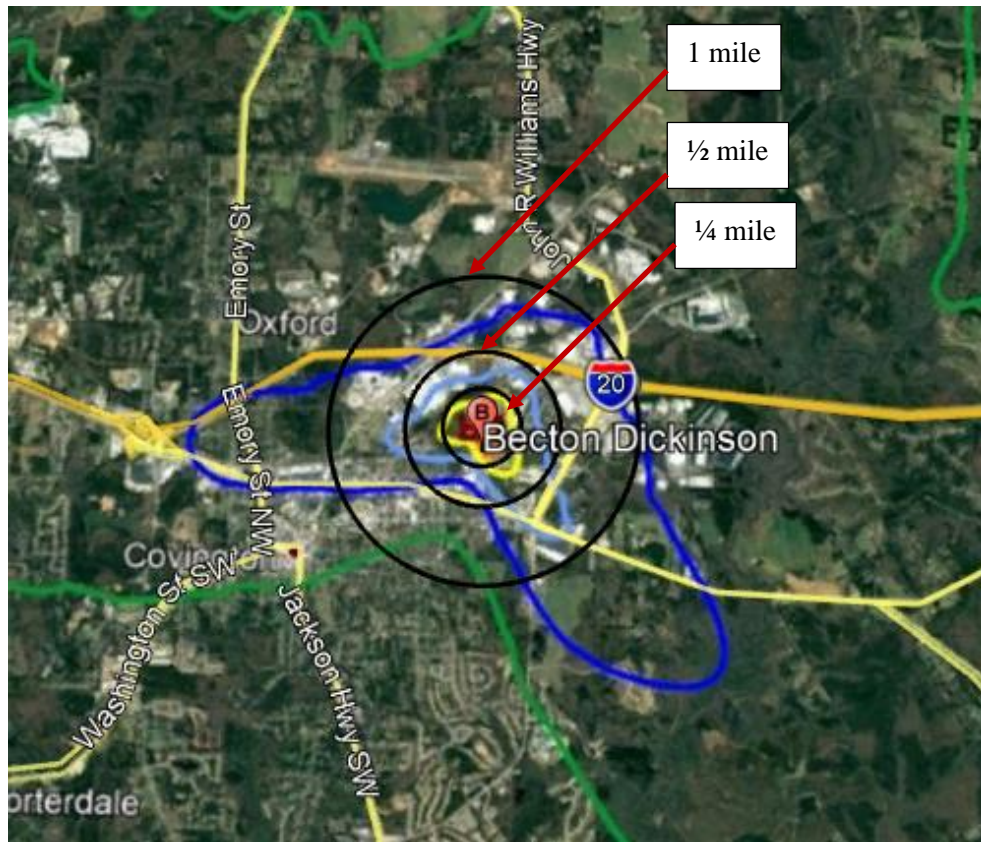


**Figure 3. Contours of 5-year Annual Average Ground-level Concentrations (in  $\mu\text{g}/\text{m}^3$ ) of Becton Dickinson (Covington) Modeled Overlaid on a Google Earth Map**



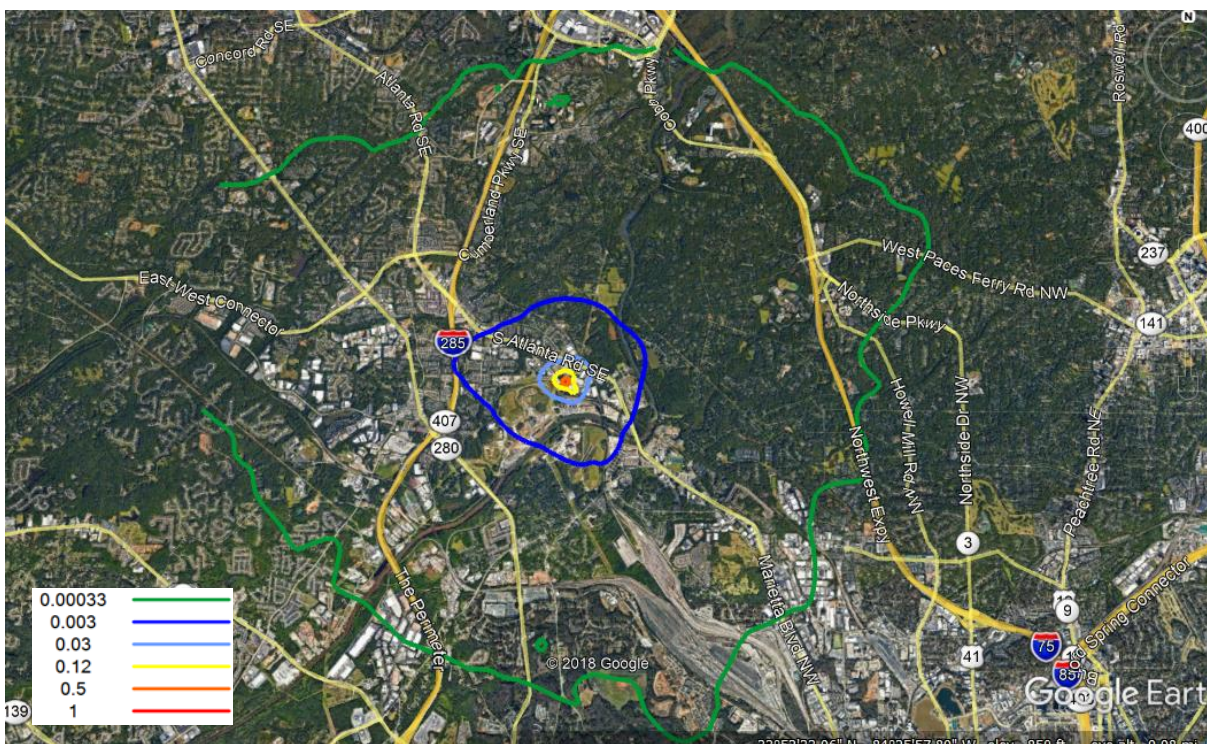
**Figure 4. A Close-up of Figure 3 (Becton Dickinson)**



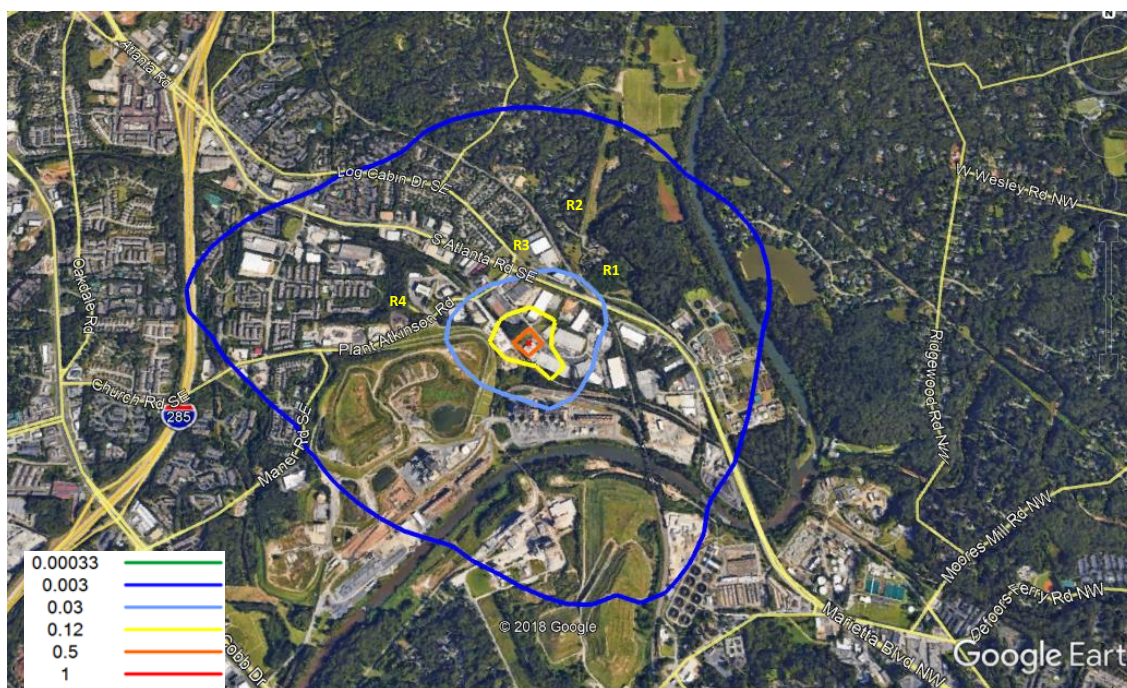


**Figure 5. Model Overlaid with Distances from Becton Dickinson**



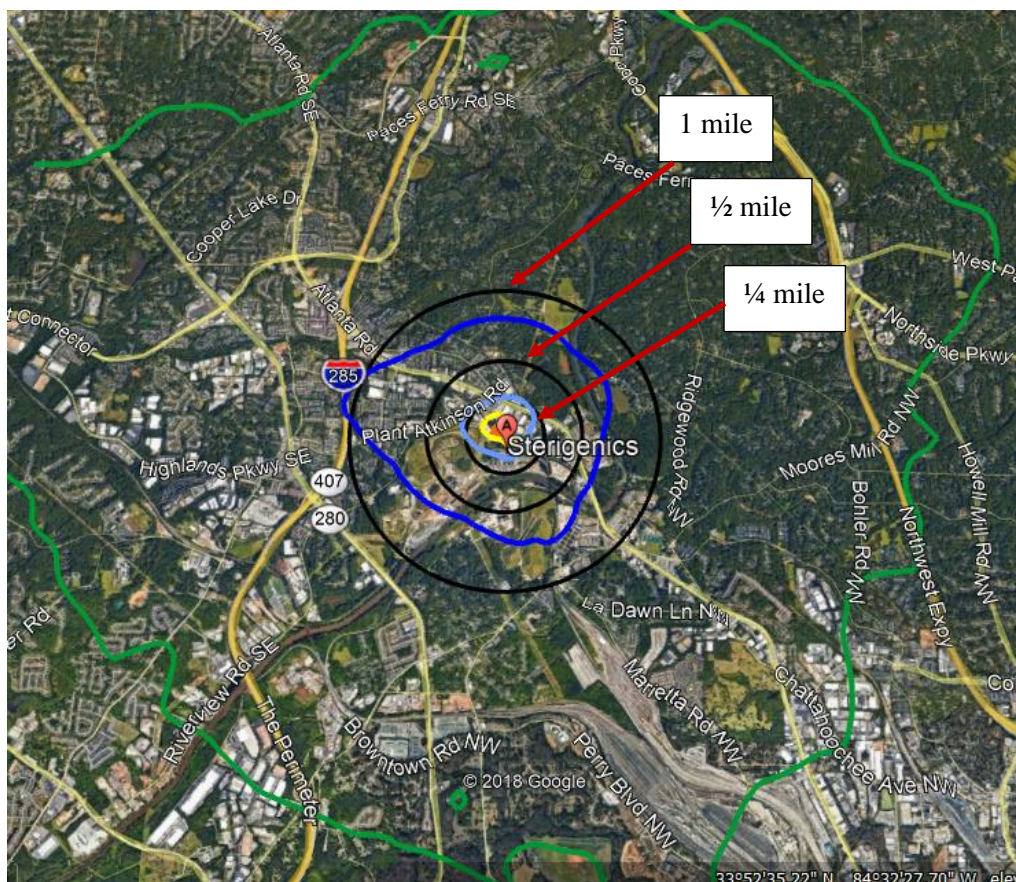


**Figure 6. Contours of 5-year Annual Average Ground-level Concentrations (in  $\mu\text{g}/\text{m}^3$ ) from Sterigenics (Cobb County) Modeled with the Current Emission Scenario Overlaid on a Google Earth Map**



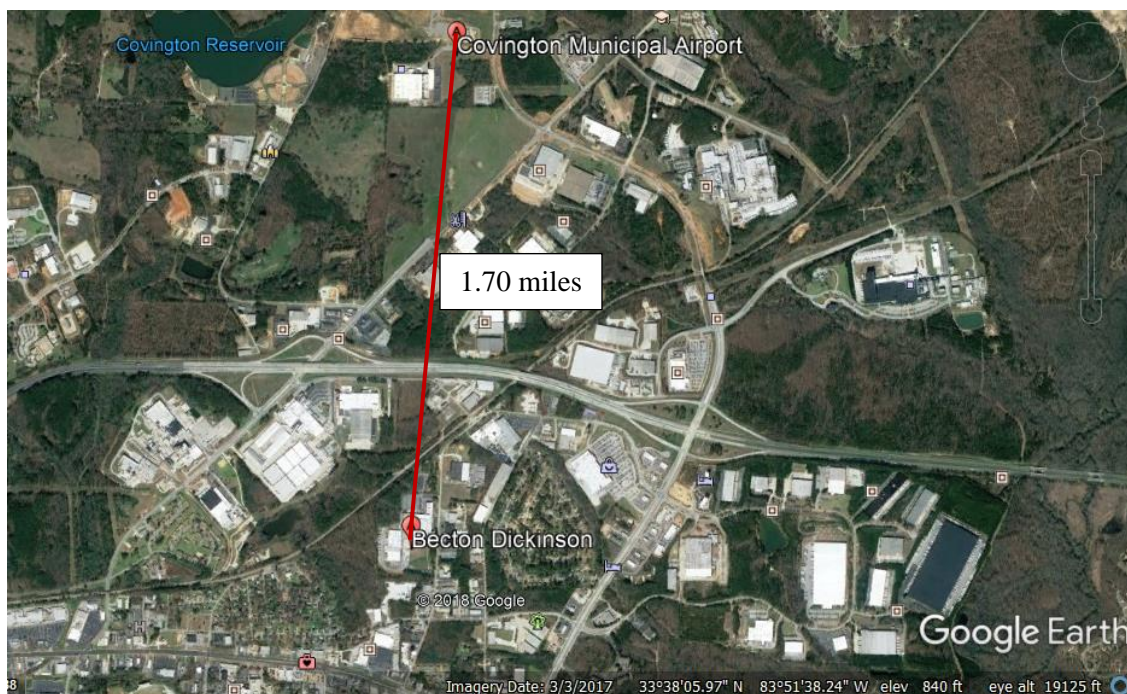
**Figure 7. A Close-up Look of Figure 6 (Sterigenics)**





**Figure 8. Model Overlaid with Distances from Sterigenics**

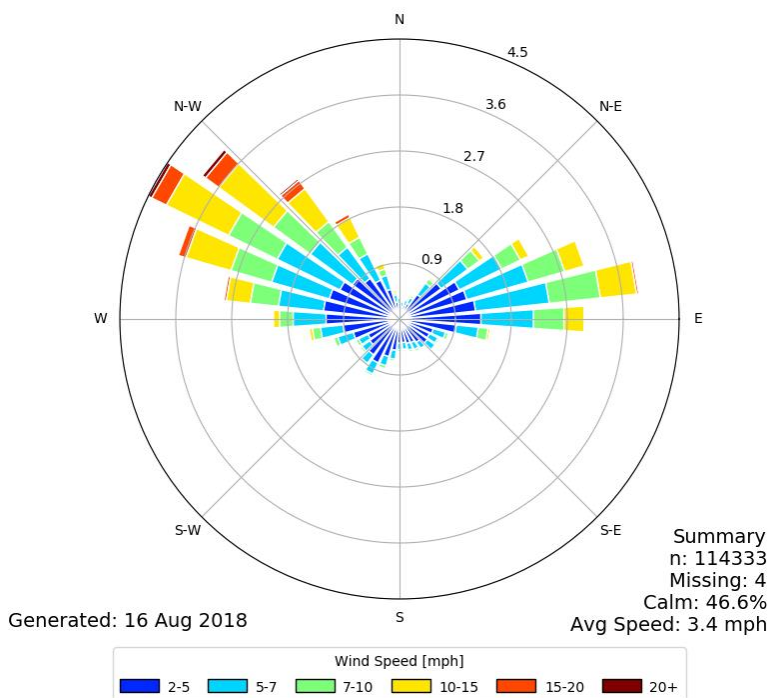
Wind rose data from airports near each facility was assessed by the GA AAMP, and primary and secondary wind patterns were determined. The available wind data from the Covington Municipal Airport was used for the Becton Dickinson facility, and the available wind data from the Dobbins Air Reserve Base was used for the Sterigenics facility. Distances from the nearby airports to the facility are shown in Figure 9 and Figure 11 below. Wind rose data from each airport is shown in Figure 10 and Figure 12.



**Figure 9. Location of Covington Municipal Airport (Wind Rose Data) in Relation to Becton Dickinson**

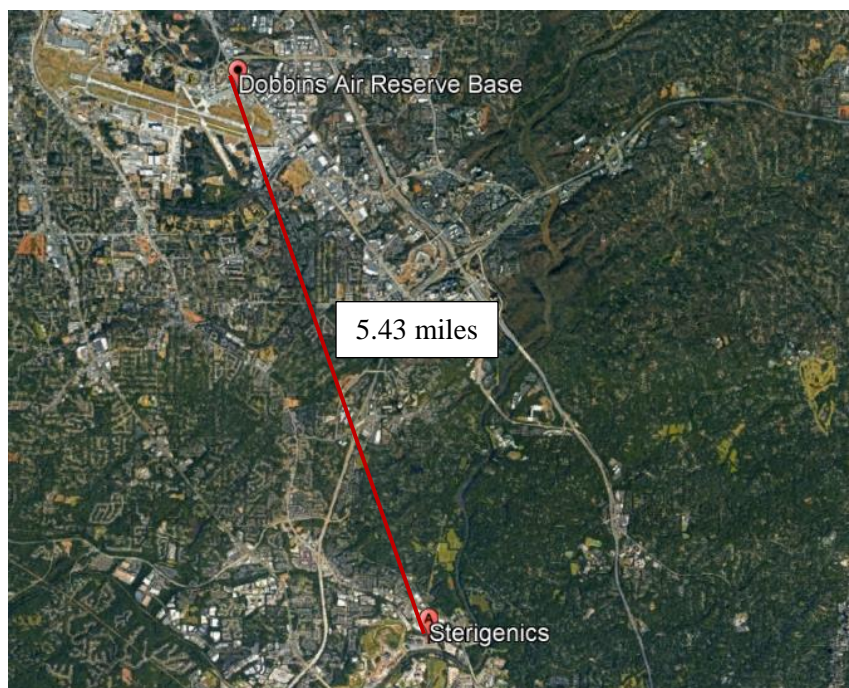


[CVC] Covington  
Windrose Plot [All Year]  
Period of Record: 12 Dec 2013 - 16 Aug 2018

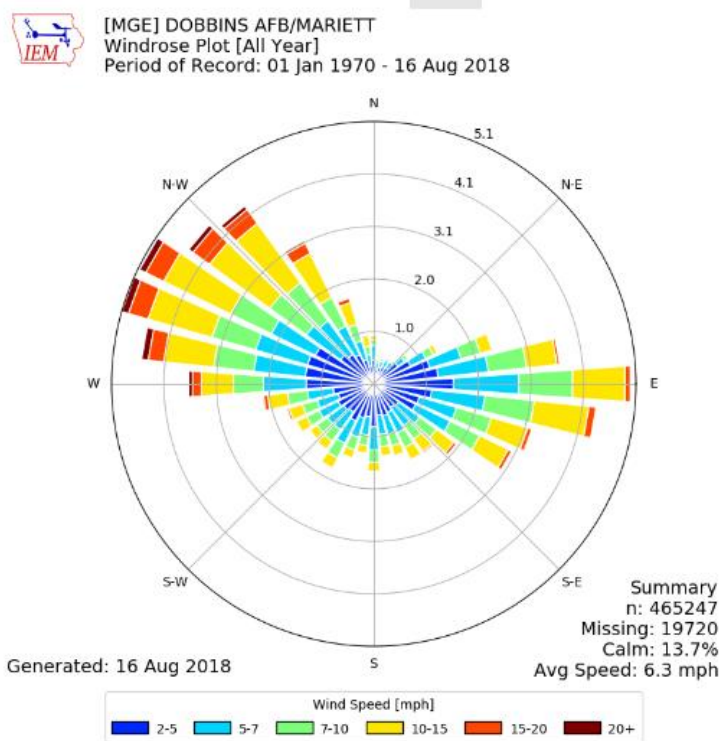


**Figure 10. Annual Wind Rose Data from Covington Municipal Airport, 2013-2018**





**Figure 11. Location of Dobbins Air Reserve Base (Wind Rose Data) in Relation to Sterigenics**

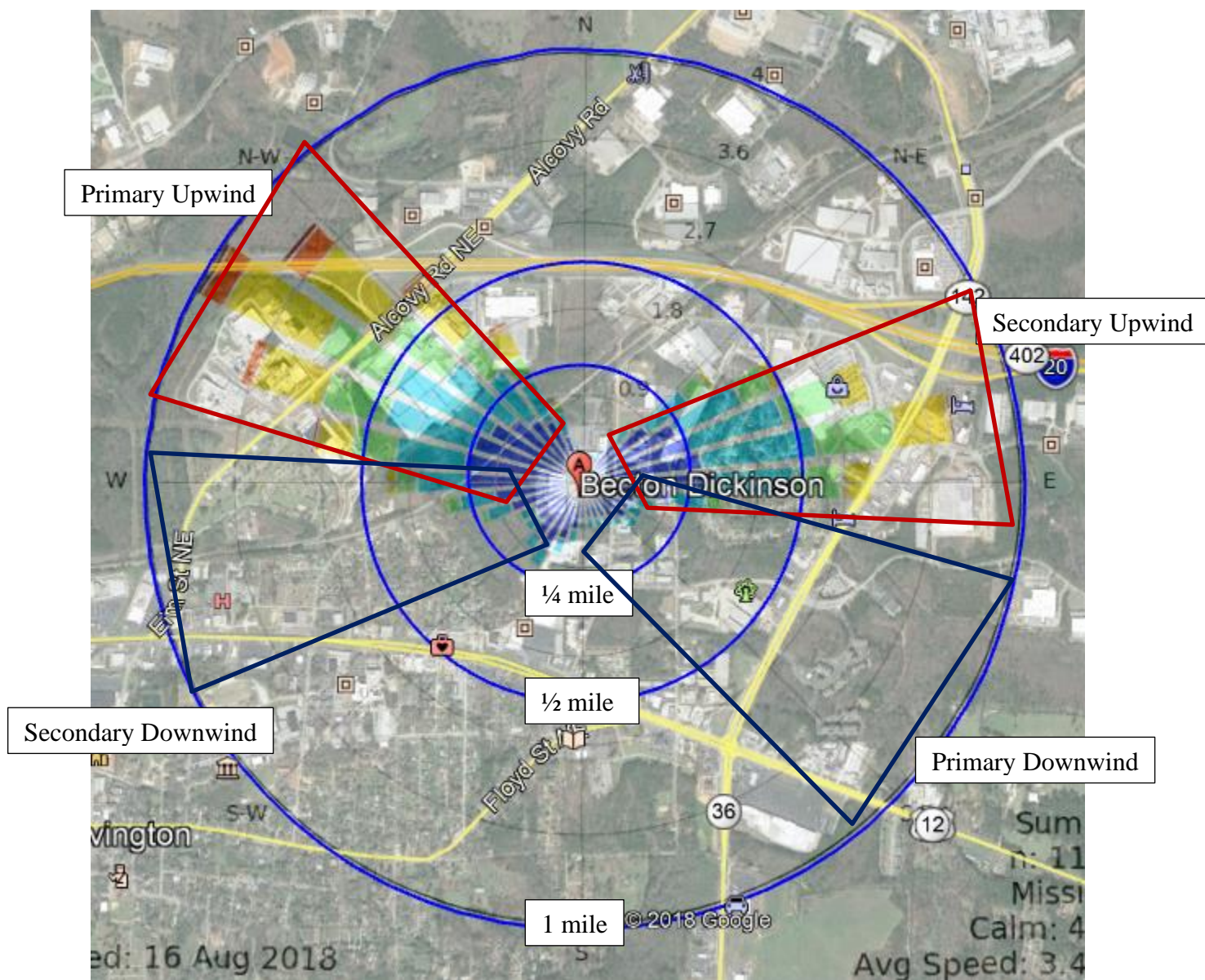


**Figure 12. Annual Wind Rose Data at Dobbins Air Reserve Base, January 1970-August 2018**

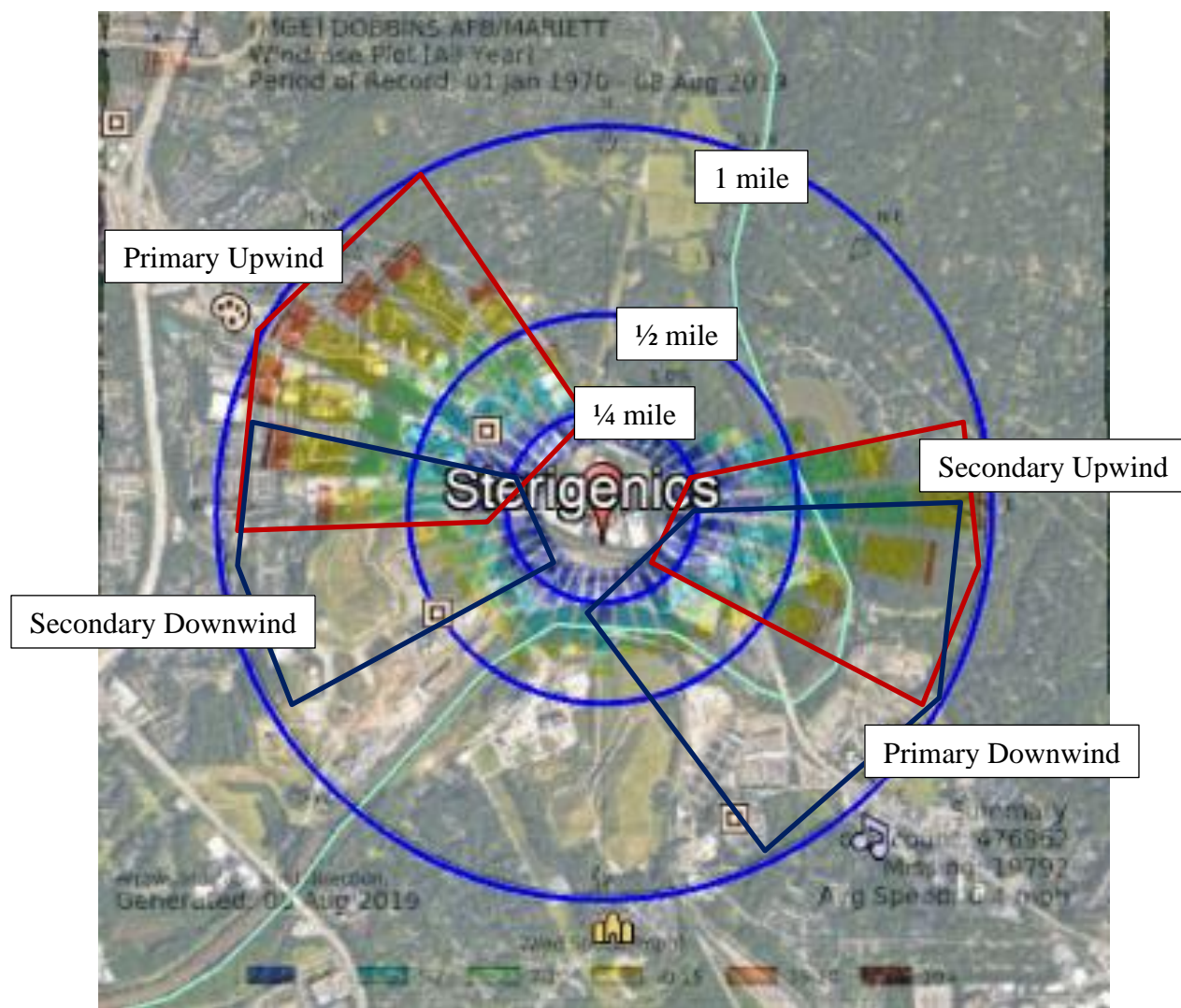
The wind roses were overlaid on Google Earth maps to help pinpoint the appropriate locations to place monitors around each facility. The GA AAMP plans to collect samples within four locations around each facility for each sampling event: primary upwind direction, primary downwind direction, secondary upwind direction and secondary downwind direction (indicated with red polygons for upwind and blue polygons for downwind in Figure 13 and Figure 14). Samples will be taken within approximately  $\frac{1}{4}$  mile of each facility in the four quadrants every six days. Samples will also be taken using the same passive sampling equipment at South DeKalb site for each sampling event. The measurements at the South DeKalb site will provide information for the relative comparison of the three locations (near Becton Dickinson, near Sterigenics, and South DeKalb). To help determine concentrations of spatial relativity to increased distance from the site of emissions, qualitative comparisons will also be made at distances of approximately  $\frac{1}{4}$  mile,  $\frac{1}{2}$  mile, and 1 mile radius of each facility. This will be accomplished by comparing a sample taken at approximately  $\frac{1}{4}$  mile to a sample taken at either approximately  $\frac{1}{2}$  mile or 1 mile. Refer to Table 1 for more information.

The GA AAMP will take reasonable precautions in placement of the passive samplers to ensure Site Operator safety. The samplers will be placed in the best places to characterize emissions in the air surrounding each facility, at heights up to 10 meters, within the breathing zone, and with an open fetch for unobstructed air flow across the samplers.





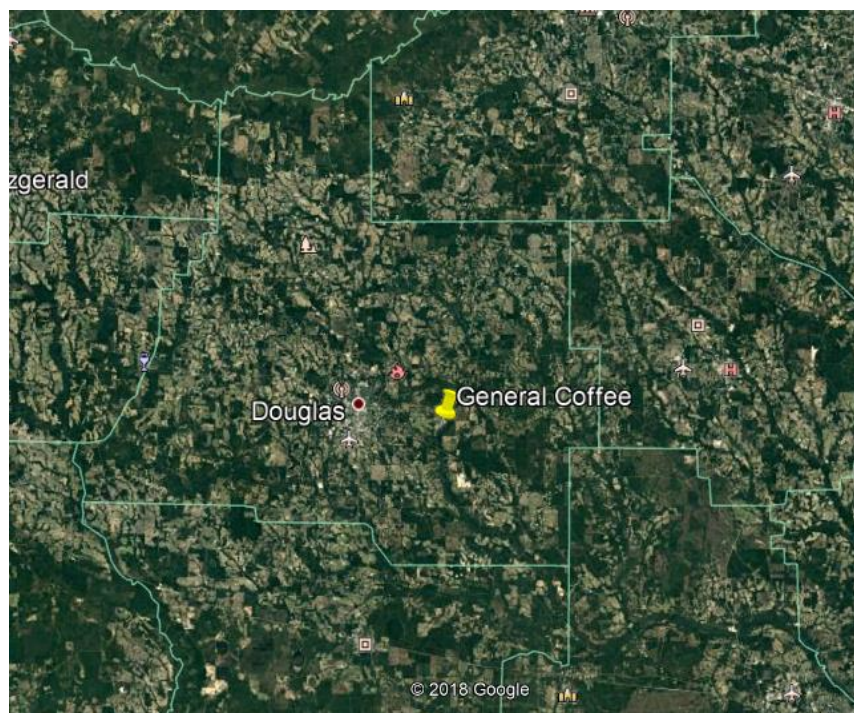
**Figure 13. Primary and Secondary Wind Directions and Distances from Becton Dickinson**



**Figure 14. Primary and Secondary Wind Directions and Distances from Sterigenics**

The GA AAMP also plans to collect samples at a rural, “background” site to compare to the samples collected near the facilities. This will help determine how much ethylene oxide is in the ambient air, with no influence from urban area activities. This background site is located at the General Coffee site (13-069-0002) (Figure 15) in Coffee County. Samples will be collected on a one in 12- day schedule at the General Coffee site.





**Figure 15. General Coffee Site**

Due to the difficulty in laboratory analysis, the ethylene oxide samples will be analyzed by the EPA contract laboratory, ERG Lab, for consistency in measurements as compared to previous EPA studies.

The GA AAMP is collecting ethylene oxide data at the South DeKalb, NR-285, and General Coffee sites to make comparisons to the data collected near each facility. In addition, comparison analyses are planned between EPA Region 4 Laboratory at Laboratory Services and Applied Science Division in Athens, GA and the EPA contract laboratory, ERG Lab, for a laboratory comparison as available.

To summarize, the GA AAMP is sampling ethylene oxide as follows (also see Table 1 below):

- Every 6 days, samples will be collected at each of four sites around each identified facility (Becton Dickinson and Sterigenics) at approximately the  $\frac{1}{4}$  mile radius mark to capture primary and secondary upwind and downwind concentrations (see above figures for primary and secondary upwind and downwind quadrants).
- Once a month, a collocated sample should be collected at one of four sites around each identified facility (Becton Dickinson and Sterigenics) at approximately the  $\frac{1}{4}$  mile radius mark to capture primary and secondary upwind and downwind concentrations (see above figures for primary and secondary upwind and downwind quadrants). The same site(s) should be used for collocation throughout the study for consistency.
- Once a month, samples should be also collected at approximately  $\frac{1}{2}$  mile or 1 mile radius from each facility, in one of the four quadrants, to assess spatial variation. Comparisons will be made between the samples collected within approximately  $\frac{1}{4}$  mile mark and the  $\frac{1}{2}$

mile mark or between the samples collected within approximately ¼ mile mark and the 1 mile mark to determine gradient of ethylene oxide concentration at the specified distance from each facility.

- Every 6 days, samples will be collected at the South DeKalb site for comparison.
- Every 12 days, samples will be collected at the background General Coffee site for comparison.
- Approximately 3 samples per study will be collected at the NR-285 site for a qualitative comparison.
- Approximately 3 samples per study will be collected concurrently using the passive canister sampler and the ATEC canister sampler at the South DeKalb site for a qualitative comparison.
- Approximately 350 samples will be collected as part of this ethylene oxide study.
- Additional sample locations may be added as resources allow, following the methodologies outlined in this document, as applicable.

A unique code will be assigned to identify and differentiate each of the monitoring sites during this study.

The GA AAMP will place the collocated samplers at the site with the expected highest concentration within reason and considering the safety of the Site Operators.

The measurement goal of the ethylene oxide study is to estimate the 24-hour average passive canister sampling concentrations in units of micrograms per cubic meter ( $\mu\text{g}/\text{m}^3$ ). The GA AAMP ethylene oxide monitoring project will follow EPA Compendium Method TO-15, as applicable, for collecting volatile organic compounds. The sampling instruments, sampling media, sampling schedules and monitoring purposes used by GA AAMP to collect air samples for the analyses of ethylene oxide are shown in the following table. Ethylene oxide will be collected at the locations around Becton Dickinson and Sterigenics, the South DeKalb site (13-089-0002), the NR-285 site (13-089-0003), and the General Coffee site (13-069-0002).

| Site Location  | Sampling Instruments                     | Sampling Media                   | Monitor Type           | Sampling Schedule                                | Monitor Purpose                                |
|--|--|----------------------------------|------------------------|--|--|
| Within approximately ¼ -mile radius of Becton-Dickinson      | Entech CS1200E Passive Canister Samplers | 6-Liter stainless steel canister | Primary and collocated | Primary-Every 6 days;<br>Collocated-Once a month | Characterization of air surrounding facilities |
| Within approximately ½ and 1-mile radius of Becton-Dickinson | Entech CS1200E Passive Canister Samplers | 6-Liter stainless steel canister | Primary                | Approximately one spatial sample per month       | Qualitative spatial comparison                 |

|   |  |                                  |                        |   |  |
|---|--|----------------------------------|------------------------|---|--|
| Within approximately ¼ mile radius of Sterigenics       | Entech CS1200E Passive Canister Samplers | 6-Liter stainless steel canister | Primary and collocated | Primary-Every 6 days; Collocated-Once a month | Characterization of air surrounding facilities |
| Within approximately ½ and 1-mile radius of Sterigenics | Entech CS1200E Passive Canister Samplers | 6-Liter stainless steel canister | Primary                | Approximately one spatial sample per month    | Qualitative spatial comparison                 |
| South DeKalb  | Entech CS1200E Passive Canister Sampler  | 6-Liter stainless steel canister | Primary and Collocated | Every 6 days; Collocated - once per month     | Comparison/background                          |
| South DeKalb  | ATEC 2200 Sampler                        | 6-Liter stainless steel canister | Primary                | Approximately 3 per study                     | Qualitative comparison                         |
| NR-285  | Xonteck Model 910 Air Sampler            | 6-Liter stainless steel canister | Primary                | Approximately 3 per study                     | Qualitative comparison                         |
| General Coffee  | Xonteck Model 911 Air Sampler            | 6-Liter stainless steel canister | Primary                | Every 12 days                                 | Rural background                               |

**Table 1. Sampling Details for Collecting Ethylene Oxide Data**

The GA AAMP may place samplers in other locations around additional facilities deemed necessary for the collection of ethylene oxide data. Those samplers will follow this QAPP, and samplers around any additional facilities will follow the monitoring objectives and procedures defined in this QAPP. For all sites used for this data study that are not established GA AAMP network sites, placement of samplers in relation to any affected source is contingent on the availability of ambient monitoring locations. All other aspects of the sample (from collection to verification) will follow the procedures outlined in this QAPP.

The work required to collect, document and report the ethylene oxide data includes:

- Appropriate placement of the sampler
- Ensuring accurate and reliable monitors records of data collected
- Developing SOPs for equipment checks, operation, and maintenance
- Establishing assessment criteria
- Validating the data produced in accordance with criteria herein

## 6.1 Field Activities

The Site Operators will perform field activities to include:

- Performing routine site operations and maintenance activities that include verifying sampler status, and recording pertinent field data and measurements
- Performing leak checks
- Collecting ethylene oxide samples and sending to ERG Lab for analysis

The Field Auditor will perform on-site assessments of the ethylene oxide collection, at least once during the study at each of the five target locations (Cobb County, Covington, South DeKalb, NR-285, General Coffee).

## 6.2 Laboratory Activities

The GA AAMP sends the ethylene oxide samples to the ERG Lab for analysis. The ERG Lab delivers an electronic data package to GA AAMP for validation and upload to the GA AAMP website (<https://epd.georgia.gov/ethylene-oxide-information>). Any issues observed with the laboratory data are discussed with the ERG Lab. The ERG Lab maintains copies of their SOPs and are available to the GA AAMP staff as needed. Copies of the ERG Lab SOPs are available upon request and the ERG Lab's *Support for the EPA National Monitoring Programs (UATMP, NATTS, CSATAM, PAMS, and NMOC Support) QAPP*, dated March 2019 is Laboratory Attachment of this document.

## 6.3 Project Assessment Techniques

The evaluation process used to measure the performance or effectiveness of the system is called an assessment. This includes the audit, performance evaluation, inspection, peer review, or surveillance.

An audit of the Site Operator's sample collection will be conducted at each of the five locations (Cobb County, Covington, South DeKalb, NR-285, and General Coffee) during the study. This audit will review equipment, adherence to the SOP, field documentation, and chain of custody records to ensure compliance with the QAPP. The results of the audits (and any identified corrective actions) are summarized in a report to the QA Unit Manager.

## 6.4 Ethylene Oxide Project Records

The GA AAMP will maintain procedures for preparation, review, approval, use, revision and maintenance of documents and records. The categories and types of records and documents that are applicable to GA AAMP are shown in Table 2. More detail is shown in Section 9.0.

**Table 2. Critical Documents and Records**

| Categories                  | Record/Document Types                                       |
|-----------------------------|---|
| Management and Organization | Organizational Chart of GA AAMP<br>Personnel qualifications |

|                               |   |
|-------------------------------|---|
| Network & Site Information    | Network description<br>Site characterization file<br>Site maps/pictures   |
| Environmental Data Operations | Quality Assurance Project Plans (QAPPs)<br>Standard operating procedures (SOPs)<br>Field and laboratory logbooks<br>Sample handling/custody records<br>Inspection/maintenance records |
| Raw Data                      | Any original data from the laboratory   |
| Data Reporting                | Data/summary reports  |
| Data Management               | Data Validation Folders   |
| Quality Assurance             | Field audits of site operations   |

## 6.5 Project Schedule

The schedule for field and laboratory analysis activities are summarized in Table 3. As the project progresses, feedback from local stakeholders may initiate changes to the project. The dates of these activities may change due to unforeseen circumstances. However, this is the general timeline that the GA AAMP will follow for this project.

**Table 3. Schedule of Monitoring Activities**

| Activity                               | Date           | Comments  |
|--|----------------|---|
| Monitoring plan development            | August 2019    | Monitoring plan vetted through official channels.   |
| QAPP development                       | August 2019    | Input taken and incorporated into official document.  |
| Sampling devices procured              | August 2019    | ERG Laboratory in place for receiving samples.  |
| Sampling devices prepared              | September 2019 | Sampling equipment zero checked at ERG Laboratory   |
| Identification of the monitoring sites | September 2019 | List of candidate sites selected considering NATA results, wind rose data, and information from local stakeholders. |
| Sampler siting/testing                 | September 2019 | Establishment of sites and preliminary testing of samplers.   |
| Field / laboratory training            | September 2019 | Field and laboratory training activities.   |
| Sampling begins                        | September 2019 | Sampler testing completed and media shipped to monitoring locations.  |
| Laboratory analysis begins             | September 2019 | Samples received and analysis begins.   |

|                              |                   |   |
|------------------------------|-------------------|---|
| Field audit assessment       | 1 audit per study | Once per location per study                                 |
| Data evaluation phase begins | April 2020*       | Data set evaluated to determine if more sampling is needed. |

\*In April 2020, evaluate to see if further measurements are needed.

## 7.0 Quality Objectives and Criteria for Measurement Data

This short-term study will be conducted under the quality program of the GA AAMP EPA-approved *Environmental Protection Division Air Protection Branch Quality Management Plan*, dated August 2015 where applicable.

### 7.1 Data Quality Objective (DQO)

The GA AAMP did not go through a formal data quality objective (DQO) process for the ethylene oxide monitoring project; however, the GA AAMP agreed upon measurement quality objectives for this project with the stakeholders. Measurement quality objectives for the various data quality indicators were developed based on the requirements of EPA Compendium Method TO-15.

### 7.2 Measurement Quality Objectives (MQOs) for Ethylene Oxide

Measurement quality objectives (MQOs), or acceptance criteria, are designed to evaluate and control various phases (sampling, preparation, analysis) of the measurement process. These MQOs are defined in terms of the following data quality indicators (DQIs):

- Precision - "Precision is a measure of agreement between two replicate measurements of the same property, under prescribed similar conditions. This agreement is calculated as either the range or as the standard deviation," (US EPA QA/G-5, Appendix D). This is the random component of error.
- Bias - "Bias is the systematic or persistent distortion of a measurement process that causes errors in one direction," (US EPA QA/G-5, Appendix D). Bias is determined by estimating the positive and negative deviation from the true value as a percentage of the true value.
- Comparability - "Comparability is the qualitative term that expresses the confidence that two data sets can contribute to a common analysis and interpolation. Comparability must be carefully evaluated to establish whether two data sets can be considered equivalent in regard to the measurement of a specific variable or groups of variables," (US EPA QA/G-5, Appendix D).
- Representativeness - "Representativeness is a measure of the degree to which data accurately and precisely represent a characteristic of a population parameter at a sampling point or for a process condition or environmental condition. Representativeness is a qualitative term that should be evaluated to determine whether in situ or other measurements are made and physical samples collected in such a manner that the resulting data appropriately reflect the media and phenomenon measured or studied," (US EPA QA/G-5, Appendix D).



- **Completeness** - Completeness is a metric quantifying the amount of valid data obtained from a measurement system compared to the amount that was expected to be obtained under correct, normal conditions. Completeness can be expressed as a ratio or a percentage. Data completeness requirements are included in the reference methods (40 CFR Part 50).
- **Sensitivity** – Sensitivity is determined by method detection limits (MDLs) for each measurement method for each pollutant (40 CFR 53.20, Table B-1 and manufacturer’s guidance).

The DQIs of representativeness, completeness, precision, bias, and sensitivity must meet specific MQOs, or acceptance criteria. The MQOs for each of the DQIs are as follows:

- **Representativeness:** Sampling must occur at one in 6-day frequency, from midnight to midnight local standard time, over  $24 \pm 1$  hours
- **Completeness:** At least 75% of all data available in a given quarter must be reported
- **Precision:** The percent difference must be no more than 25% for results  $\geq 5 \times \text{MDL}$
- **Bias:** Measurement error must be  $<3 \times \text{MDL}$
- **Sensitivity:** MDL as required by EPA as part of national contract (see ERG Laboratory’s QAPP attached)

For the GA AAMP ethylene oxide monitoring project to follow these MQOs, the data produced will be considered of sufficient quantity and quality for the decision making to commence. The following data validation table outlines the acceptance criteria to meet these MQOs. GA AAMP uses the acceptance criteria provided in EPA supplied guidance *Technical Assistance Document for the National Air Toxics Trends Stations Program, Revision 3*, dated October 2016 as a guide, and unless otherwise noted, the references shown in the table refer to this document. The MQOs are used by GA AAMP to control and assess measurement uncertainties.

**Table 4. Data Validation Table**

**VOCs via EPA Compendium Method TO-15**

| Parameter   | Description and Required Frequency   | Acceptance Criteria  | Reference                            | Category         |
|---|--|--|--------------------------------------|------------------|
| <i>Field Readiness Checks and Collection Activities</i> |  |  |                                      |                  |
| Canister Viability                                      | All canisters  | Canister must be used within 30 days from final evacuation         | Section 4.2.6.2<br>TO-15 Section 1.3 | Operational      |
| Canister Starting Pressure Determination                | Each canister prior to collection of a field sample or preparation of a calibration standard or laboratory QC sample | Vacuum $\geq -28$ inHg   | Section 4.2.5.2.1                    | Critical         |
| Sample Setup Leak Check                                 | Each canister prior to collection - draw vacuum on canister connection   | Leak rate must be $\leq 1$ inHg over 5 minutes                     | Section 4.2.5.2.1                    | Critical         |
| Sampling Frequency                                      | One sample every six days according to the EPA National Monitoring Schedule  | Sample must be valid to be included in $\geq 75\%$                 | Section 4.2.5.3                      | Critical and MQO |
| Sampling Period   | All field-collected samples  | 1380-1500 minutes ( $24 \pm 1$ hr) starting and ending at midnight | Section 4.2.5.3                      | Critical and MQO |
| Field-collected Sample Final Pressure                   | All field-collected samples  | Must be determined with a pressure gauge                           | Section 4.2.5.2.4                    | Operational      |
| Trip Blanks   | Once a month on primary field-collected samples  | Measurement $<3 \times \text{MDL}$                                 | Section 4.3.8.2.2                    | Operational      |
| <i>Sample Receipt</i>                                   |  |  |                                      |                  |

|   |  |   |  |  |
|---|--|---|--|--|
| Chain-of-custody  | All field-collected samples including field QC samples   | Each canister must be uniquely identified and accompanied by a valid and legible COC with complete sample documentation     | Sections 3.3.1.3.7 and 4.2.5.2.4                   | Critical   |
| Sample Holding Time                                       | All field-collected samples, laboratory QC samples, and standards  | Analysis within 30 days of end of collection (field-collected samples) or preparation (QC samples or standards)             | Section 4.2.1 TO-15 Sections 1.3, 2.3, and 9.2.8.1 | Operational  |
| Canister Receipt Pressure Check                           | All field-collected samples upon receipt at the laboratory – measured with calibrated pressure gauge or transducer | Pressure change of $\leq 3$ inHg from the final pressure at retrieval   | Section 4.2.8                                      | Critical for subambient sample collection, operational for pressurized sample collection |
| <b>GC/MS Analysis</b>                                     |  |   |  |  |
| Refer to ERG Lab's attached QAPP                          |  |   |  |  |
| <b>Laboratory Readiness and Proficiency</b>               |  |   |  |  |
| Refer to ERG Lab's attached QAPP                          |  |   |  |  |
| <b>Canister and Sampling Unit Testing and Maintenance</b> |  |   |  |  |
| Refer to ERG Lab's attached QAPP                          |  |   |  |  |
| <b>Site Specifications and Maintenance</b>                |  |   |  |  |
| Sample Inlet Filter                                       | Particulate filter maintenance<br>Beginning of study   | Change filter when canister pressure shows necessary<br><br>Clean or replace the 2- $\mu$ m sintered stainless steel filter | Section 4.2.3.3 TO-15 Section 7.1.1.5              | Operational  |
| <b>Data Reporting</b>                                     |  |   |  |  |
| Data Completeness   | Valid samples compared to scheduled samples<br>For duration of study   | $\geq 75\%$ of scheduled samples  | Section 3.2  | MQO  |

### 7.3 Intended Use of Data

This data will be used to:

- Characterize ambient levels of ethylene oxide
- Establish background concentration of ethylene oxide
- Provide ethylene oxide data for risk characterization by other agencies

The quality of the data must be evaluated and controlled to ensure that it is maintained within the established acceptance criteria. Measurement quality objectives (MQOs) are designed to evaluate and control various phases (sampling, preparation, analysis) of the measurement process.

### 7.4 Measurement Scale

Each sampler operated by GA AAMP is assigned a scale of representativeness based on 40CFR58, Appendix D. The ethylene oxide monitors represent a middle scale to neighborhood scale. These

representativeness definitions are found in GA AAMP's *Annual Ambient Air Monitoring Plan* at <https://airgeorgia.org/>.

## 8.0 Personnel Training and Development Program

This section is not required for a Category II QAPP.

## 9.0 Documentation and Records

GA AAMP, as a PQAQ performing environmental data operations and management activities, has established and maintained procedures for the timely preparation, review, approval, issuance, use, control, revision and maintenance of documents and records. These procedures are elaborated in this section as a documentation and records management policy to address at least the following elements:

- A list of files considered the official records and their media type (e.g., paper, electronic)
- Schedule for retention and disposition of records
- Storage and retrieval system of records
- Person(s) responsible at each level of storage and retrieval for records
- Assignment of appropriate levels of security

A document, from a records management perspective, is a volume that contains information that describes, defines, specifies, reports, certifies, or provides data or results pertaining to environmental programs. Table 5 lists the categories and types of records and documents that are applicable for document control in the GA AAMP. Information on key documents in each category is included in this section. With the exception of Field Logbooks which are kept with the operator, all paper records are stored in the GA AAMP central office. In addition to paper records, all the applicable documentation referred to in this section is saved as an electronic record with a format of MS Word, MS Excel, or PDF on the local network on the GA AAMP server. Retention of both paper and electronic records is explained in Section 9.3 below. The paper and electronic records are stored in a logical order for ease of access. For details of the ERG Lab's record management process, refer to the ERG Lab's QAPP attached.

**Table 5. Types of Information Retained Through Document Control**

| Categories                  | Record/Document Types           | Electronic Copy | Paper Copy |
|-----------------------------|---------------------------------|-----------------|------------|
| Management and Organization | Organizational Chart of GA AAMP | X               | X          |
|                             | Personnel qualifications        | X               | X          |
|                             | Support contracts               | X               | X          |
| Network & Site Information  | Network description             | X               | X          |
|                             | Site description for study      | X               | X          |
|                             | Site characterization file      | X               | X          |
|                             | Site maps/pictures              | X               | X          |

|                               |   |   |   |
|-------------------------------|---|---|---|
| Environmental Data Operations | Quality Assurance Project Plans (QAPPs) | X | X |
|                               | Standard operating procedures (SOPs)    | X | X |
|                               | Field logbooks                          |   | X |
|                               | Sample handling/custody records         | X | X |
|                               | Inspection/maintenance records          | X | X |
|                               | NIST traceable records                  | X | X |
| Raw Data                      | Any original data                       | X | X |
| Data Reporting                | Data/summary reports                    | X | X |
| Data Management               | Data Validation Folders                 | X | X |
| Quality Assurance             | Field Audits of Site Operations         | X | X |
|                               | NIST traceable records                  | X | X |

The GA AAMP has permission from the property owners to place its ethylene oxide ambient air samplers.

For the GA AAMP SOPs and QAPPs, the original copies are considered controlled copies and are maintained by the Program Manager. GA AAMP SOPs and QAPPs are available in ‘read only’ format on the local network drive and through online database records for operations. The current GA AAMP SOPs are retained in a folder for the GA AAMP at S:\Ambient\SOPs for Operations Unit. The current GA AAMP QAPPs are stored at S:\Ambient\QAPPs. The GA AAMP’s historical SOPs and QAPPs are removed as they are replaced.

The GA AAMP’s raw data records on the local network server are backed up every 24 hours. In addition, the local network server files are kept as a redundant system to ensure proper storage of GA AAMP raw data records.

The GA AAMP’s raw data records that are housed on the local network are only available to the GA AAMP staff. The raw data is validated as discussed in Section 20.0 and posted to the GA EPD’s website (<https://epd.georgia.gov/ethylene-oxide-information>). Historical QA documents are retained in hardcopy in GA AAMP files and/or electronic ‘read only’ access. Any of GA AAMP’s hard copy site information (maps, photos, etc.) is housed in the central files.

## 9.1 Routine Data Activities

GA AAMP maintains records in appropriate files that allow for the efficient archival and retrieval of records. Ambient air quality information is included in this system. Table 5 includes the documents and records that are filed according to the statute of limitations discussed in Section 9.3.

## 9.2 Documentation Control

The details of the documents and records listed in Table 5 will be discussed in the appropriate sections of this document. All raw data required for calculations is collected electronically or on

data forms that are included in the field and analytical methods. All hardcopy information shall be filled out in indelible ink. Corrections shall be made by inserting one line through the incorrect entry, initialing and dating this correction, and placing the correct entry alongside the incorrect entry, if this can be accomplished legibly, or by providing the information on a new line if the above is not possible.

### **9.2.1 Logbooks**

Each Site Operator is responsible for obtaining appropriate field logbooks uniquely numbered and associated with the individual and/or a specific program. These logbooks will be used to record information about the site and laboratory operations, as well as document routine operations.

Completion of data entry forms, associated with all routine environmental data operations, are required even when the field logbooks contain all appropriate and associated information required for the routine operation being performed.

- Field Logbooks - Logbooks are used for each monitoring site, specific program, audit, or individual. Each notebook should be hardbound and paginated. After use in the field, field logbooks are retained in Site Operator's office.
- Laboratory Logbooks – Logbooks are used for sample custody, sample preparation and instrumental analysis. Each notebook should be paginated. An electronic database (Laboratory Information Management System or LIMS) exists in which the ERG Lab retains all data records pertaining to sample tracking, preparation, and analysis, as well as general comments and notations and other pertinent information required for support of the GA AAMP's data integrity activities. Refer to ERG's Laboratory Attachment for more details.

### **9.2.2 Chain-of-Custody Forms**

For any samples that are taken to the ERG Lab for analysis, a Chain-of Custody (COC) form is created. Custody records document the "chain of custody": the date and person responsible for the various sample handling steps associated with each sample and the information that acknowledges that sample integrity remained intact. Custody records also provide a reviewable trail for quality assurance purposes and can be used as evidence in legal proceedings. The GA AAMP and ERG Lab track and document the whereabouts of each sample at each stage throughout the data collection operation using the Field Data Sheet and the COC form as shown in the applicable SOPs listed in Table 7. Entries on the COC form are made by hand. The information is then entered into the sample tracking system, where an electronic record is kept. More information about COC forms is detailed in Section 12.0.

## **9.3 Data Archiving and Retrieval**

The storage and retrieval of the air quality monitoring data are conducted through the archiving system of GA EPD. All the information listed in Table 5 will be retained in house for at least five years from the date of collection. However, if any litigation, claim, negotiation, audit, or other action involving the records has been started before the expiration of the five-year period, the

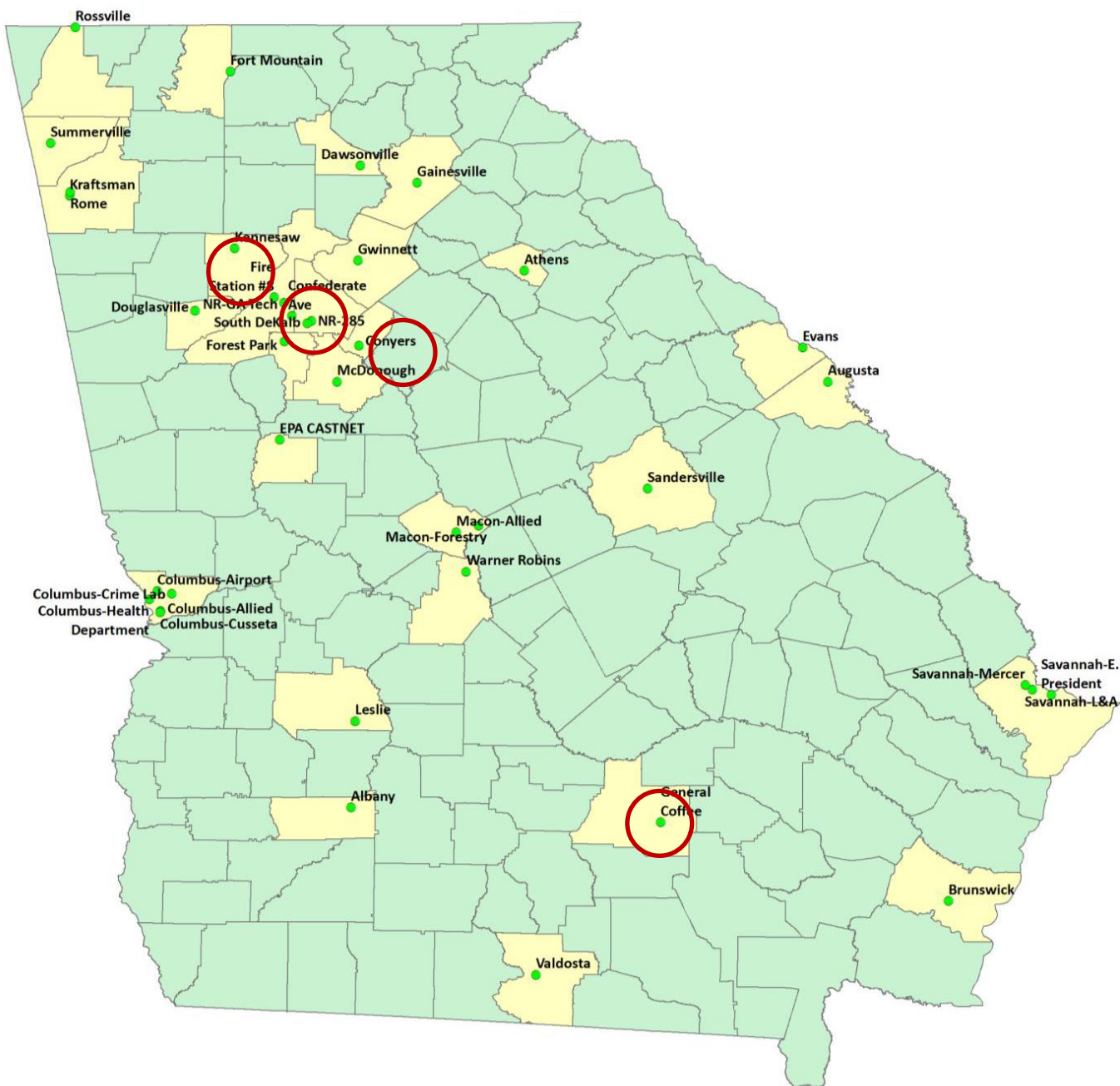
records will be retained until completion of the action and resolution of all issues which arise from it, or until the end of the five year-period, whichever is later.

## **10.0 Network Description**

For a detailed description of the GA AAMP's ethylene oxide study sampling design, refer to Section 6.0. Figure 16 shows the areas that the GA AAMP will monitor for ethylene oxide (indicated by red circles).

Details regarding the South DeKalb, NR-285, and General Coffee sites can be found in GA AAMP's *Annual Ambient Air Monitoring Plan* at <https://airgeorgia.org/>.

The GA AAMP may place samplers in other locations around additional facilities deemed necessary for the collection of ethylene oxide data. Those samplers will follow this QAPP, and samplers around any additional facilities will follow the monitoring objectives and procedures defined in this QAPP. For all sites used for this data study that are not established GA AAMP network sites, placement of samplers in relation to any affected source is contingent on the availability of ambient monitoring locations. All other aspects of the sample (from collection to verification) will follow the procedures outlined in this QAPP.



**Figure 16. Location of Ethylene Oxide Monitoring Sites**

### 10.1 Monitoring Objective

The GA AAMP's ethylene oxide sites are representative of a middle to neighborhood scale and collect data with a source-oriented monitoring objective.

## 10.2 Sampling Frequency

For a detailed description of the GA AAMP's ethylene oxide study sampling frequency, refer to Section 6.0. Latitude and longitude coordinates will be disclosed after the study is complete. Samples will be collected from midnight to midnight. Sampling frequencies are shown in Table 6.

**Table 6. Sampling Frequency of Ethylene Oxide Monitors**

| Site Location  | Sampling Instruments                     | Sampling Media                   | Monitor Type           | Sampling Schedule                                | Monitor Purpose                                |
|--|--|----------------------------------|------------------------|--|--|
| Within approximately ¼ -mile radius of Becton-Dickinson      | Entech CS1200E Passive Canister Samplers | 6-Liter stainless steel canister | Primary and collocated | Primary-Every 6 days;<br>Collocated-Once a month | Characterization of air surrounding facilities |
| Within approximately ½ and 1-mile radius of Becton-Dickinson | Entech CS1200E Passive Canister Samplers | 6-Liter stainless steel canister | Primary                | Approximately one spatial sample per month       | Qualitative spatial comparison                 |
| Within approximately ¼ mile radius of Sterigenics            | Entech CS1200E Passive Canister Samplers | 6-Liter stainless steel canister | Primary and collocated | Primary-Every 6 days;<br>Collocated-Once a month | Characterization of air surrounding facilities |
| Within approximately ½ and 1-mile radius of Sterigenics      | Entech CS1200E Passive Canister Samplers | 6-Liter stainless steel canister | Primary                | Approximately one spatial sample per month       | Qualitative spatial comparison                 |
| South DeKalb   | Entech CS1200E Passive Canister Sampler  | 6-Liter stainless steel canister | Primary and collocated | Every 6 days;<br>Collocated – once a month       | Comparison/background                          |
| South DeKalb   | ATEC 2200 Sampler                        | 6-Liter stainless steel canister | Primary                | Approximately 3 per study                        | Qualitative comparison                         |
| NR-285   | Xonteck Model 910 Air Sampler            | 6-Liter stainless steel canister | Primary                | Approximately 3 per study                        | Qualitative comparison                         |
| General Coffee   | Xonteck Model 911 Air Sampler            | 6-Liter stainless steel canister | Primary                | Every 12 days                                    | Rural background                               |



### 10.3 Site Selection

GA AAMP considered the following aspects when establishing the ethylene oxide air monitoring sites:

- Understanding the monitoring objective(s)
- Identifying the spatial scale most appropriate for the monitoring objective(s)
- Identifying the general locations where the monitoring site(s) should be placed according to wind direction
- Identifying specific monitoring sites

The sites will be chosen as GA AAMP's ethylene oxide sites due to the following factors:

- Modeled ethylene oxide emissions data showing highest concentrations
- Transport of pollutants downwind of facilities
- Characterize air upwind of facilities

## 11.0 Sampling Method Requirements

### 11.1 Field Collection Description

Ethylene oxide samples are collected in 6 Liter stainless steel canisters. The Site Operators receive certified "clean" canisters from the ERG Lab. These canisters are evacuated to at least -28 inches of mercury gauge pressure when connected to the samplers. When not attached to the sampler, the canister is capped using a brass or stainless steel cap. Unique sample identification (ID) numbers are printed on tags attached to the canister (Figure 22, in next Section). Each canister also has a unique ID permanently written on the canister. For this short term study, the passive ethylene oxide monitor kits were sent to ERG Lab for a zero leak check at the beginning of the study. These results were evaluated and no contamination was found. The ERG Lab data is available upon request.

Prior to sampling, each canister must pass the leak check procedure. Primary samples will be collected on a one in 6-day schedule. Collocated samples should be collected once a month. Refer to Section 6.0 for more details regarding sampling schedules. The sample will run for 24 hours  $\pm$  1 hour. The Entech passive sampler is complete when it reaches subambient pressure, typically 2 to 4 inches mercury (inHg) (all other samplers i.e. the ATEC and Xonteck samplers should have an ending pressure of  $\geq +5$  psig). The filled canister is then removed from the VOC sampler and subsequently delivered to the ERG Lab for analysis. For more information regarding the ERG Lab, see Laboratory Attachment of this document.

### 11.2 Sampling Methodology

The methods described herein provide for measurement of the relative concentration of ethylene oxide in ambient air for a 24-hour sampling period. The method described in this section is based on *Compendium Method for the Determination of Toxic Organic Compounds in Air*, United States Environmental Protection Agency, Section TO-15, January 1999. The samplers located near the facilities and at the South DeKalb site will be a CS1200E Passive Sampler from Entech Instruments, which will connect directly to a 6-liter stainless steel canister (Figure 17). A TM1200

Canister Sampling Timer treated with silica for non-reactivity will be used to automatically start and stop the sampling at a 24-hour period. See GA AAMP's *Standard Operating Procedure Entech CS1200E Passive Sampler Kit* for more details.



**Figure 17. Entech CS1200E Passive VOCs Sampler Set-Up**

In addition, at the South DeKalb site, the ethylene oxide sampler will be the ATEC 2200 with a 6-liter stainless steel canister (Figure 18). See GA AAMP's *Standard Operating Procedure for Operation of a Volatile Organic Compound (VOC) Canister Sampler for a National Air Toxics Trends Station (NATTS)* for more details.



**Figure 18. ATEC 2200 VOCs Sampler**

The ethylene oxide sampler at the NR-285 site will be the Xonteck Model 910 Sampler with a 6-liter stainless steel canister (Figure 19). See GA AAMP's *Standard Operating Procedure for the Xonteck Model 910 VOCs Canister Sampler* for more details.



**Figure 19. Xonteck Model 910 VOCs Sampler**

At the General Coffee site, the ethylene oxide sampler will be the Xonteck Model 911 Sampler with a 6-liter stainless steel canister (Figure 20). See GA AAMP's *Standard Operating Procedure for Xonteck Model 911 VOCs Canister Sampler* for more details.



**Figure 20. Xonteck Model 911 VOCs Sampler**

### 11.3 Standard Operating Procedures

In order to perform the sampling, analysis, and QC activities consistently, GA AAMP has prepared and updated standard operating procedures (SOPs) for each routine or repetitive task as a part of the QAPP. The SOPs prepared and updated by GA AAMP for the ethylene oxide monitoring study are summarized in Table 7. At the time of writing this QAPP, some SOPs were still being updated.

The GA AAMP and ERG Lab's SOPs detail the instrument operation requirements. Table 7 shows a list of GA AAMP's SOPs that apply to the VOCs samplers. For ERG Lab's SOPs, see Section 8.0 and Appendix D of the ERG Laboratory Attachment of this document.

**Table 7. GA AAMP's SOPs for Ethylene Oxide Collection**

| <b>SOP</b>  | <b>Revision</b> | <b>Date</b>    |
|---|-----------------|----------------|
| Standard Operating Procedure for Operation of a Volatile Organic Compound (VOC) Canister Sampler for a National Air Toxics Trends Station (NATTS) | 1               | August 2017    |
| Standard Operating Procedure for the Xonteck Model 910 VOCs Canister Sampler  | 0               | September 2019 |
| Standard Operating Procedure for Xonteck Model 911 VOCs Canister Sampler  | 0               | September 2019 |
| Standard Operating Procedure Entech CS1200E Passive Sampler Kit   | 0               | September 2019 |
| Standard Operating Procedure for Data Validation of Integrated Data   | 2               | September 2018 |

#### **11.4 Sample Probe/Sample Train**

For the VOCs samplers at the ethylene oxide monitoring sites, the GA AAMP uses the Entech Passive VOCs samplers, ATEC samplers, and the Xonteck 910/911 VOCs samplers, which are free standing samplers and do not have a sampler probe/train that requires maintenance. In addition to the leak checks described in these documents and in Section 14.0 of this QAPP, GA AAMP will clean the exterior of the VOCs sampler at least once during this study, or as needed. Details are shown in the applicable Operations' SOPs listed in Table 7.

#### **11.5 Sampler Leak Check**

The GA AAMP performs a leak check before each sample is collected. Details are explained in the applicable Operations' SOPs listed in Table 7. Per the SOPs noted above and Table 4 contained in this QAPP, the passive ethylene oxide samplers underwent a leak check performed by the ERG Laboratory prior to beginning this study.

#### **11.6 Maintenance of Sampler Probe/Sampler Train**

Preventative maintenance is performed on the ethylene oxide samplers by GA AAMP as described in the applicable SOPs listed in Table 7. Per the SOPs noted above and Table 4 contained herein this QAPP, the following maintenance is performed as stipulated. The GA AAMP replaces the sample inlet filter as indicated by pressure issues. The sample probes and inlets will be cleaned as needed, in addition to the sample line replacement.

## **11.7 Modifications to Samplers**


In the event of needed corrective action, the Site Operator notifies the Operations Unit Manager. The QA Unit Manager and Program Manager should also be notified. Details are described in the applicable SOP listed in Table 7.

## **12.0 Sample Numbering and Custody**

Unique sample IDs are generated by the ERG Lab and labeled appropriately on the sampling media (see Section 11.0 for details of how sample IDs are addressed). The GA AAMP utilizes these sample IDs to match the laboratory data to the field data, as applicable. GA AAMP may employ custody seals on the samples, and the samples are either in secured GA EPD buildings, ERG buildings, secured at the sampling location, or in the possession of GA EPD or ERG personnel.

A critical activity within any data collection phase involving physical samples is the handling of sample media prior to sampling; transporting sample media to the field, handling samples in the field at the time of collection; storage of samples (in the field or other locations); transport of samples from the field site; and the analysis of the samples. Custody records document the “chain of custody”: the date and person responsible for the various sample handling steps associated with each sample and the information that acknowledges that sample integrity remained intact. Custody records also provide a reviewable trail for quality assurance purposes and can be used as evidence in legal proceedings. The GA AAMP and ERG Lab track and document the whereabouts of each sample at each stage throughout the data collection operation using the Field Data Sheet, Chain-of-Custody (COC) Form, and ERG Tracking Tag as shown in the applicable SOPs listed in Table 7. Entries on the COC form are made by hand. The information is then entered into the ERG sampling tracking system (LIMS), where an electronic record is kept. More details are shown in the SOPs in Table 7 and the ERG’s Laboratory Attachment of this document. Examples of the COC Form, Sample Tracking Tag, and Logbook are shown below.

Figure 21. Example of Chain-of-Custody Form

|   |  |  |                                       |
|---|--|--|---------------------------------------|
|  |  | <div style="border: 1px solid black; padding: 2px;">ERG Lab ID #</div> |                                       |
| <small>601 Keystone Park Drive, Suite 700, Morrisville, NC 27560</small>          |  |  |                                       |
| <b>AIR TOXICS SAMPLE CHAIN OF CUSTODY</b>   |  |  |                                       |
| <b>Lab<br/>Pre-Sampling</b>   | Site Code: _____   |  | Canister Number: _____                |
|   | City/State: _____  |  | Lab Initial Can. Press. ("Hg): _____  |
|   | AQS Code: _____  |  | Cleaning Batch #: _____               |
|   | Collection Date: _____   |  | Date Can. Cleaned: _____              |
|   | Options:<br>SNMOC (Y/N): _____                                     |  | Duplicate Event (Y/N): _____          |
|   | TOXICS (Y/N): _____  |  | Duplicate Can #: _____                |
|   | METHANE (Y/N): _____   |  |                                       |
|   | Relinquished by: _____   |  | Date: _____                           |
| <b>Field<br/>Setup</b>  | Received by: _____   |  | Date: _____                           |
|   | Operator: _____  |  | MFC Setting: _____                    |
|   | System #: _____  |  | Elapsed Timer Reset (Y/N): _____      |
|   | Setup Date: _____  |  | Canister Valve Opened (Y/N): _____    |
|   | Field Initial Can. Press.: _____ <b>psig psia "Hg (Circle one)</b> |  |                                       |
| <b>Field<br/>Recovery</b>   | Recovery Date: _____   |  | Sample Duration (3 or 24 hr): _____   |
|   | Operator: _____  |  | Elapsed Time: _____                   |
|   | Field Final Can. Press.: _____ <b>psig psia "Hg (Circle one)</b>   |  |                                       |
|   | Status: <b>VALID</b> <b>VOID</b> (Circle one)                      |  | Canister Valve Closed (Y/N): _____    |
|   | Relinquished by: _____   |  | Date: _____                           |
| <b>Lab<br/>Recovery</b>   | Received by: _____   |  | Date: _____                           |
|   | Lab Final Can. Press.: _____ <b>psig "Hg (Circle one)</b>          |  | Converted to psia: _____              |
|   | Status: <b>VALID</b> <b>VOID</b> (Circle one)                      |  | Gauge: <b>1</b> <b>2</b> (Circle one) |
|   | If void, why: _____  |  |                                       |
| <b>Samples stored in Air Tox Lab (Room 130)</b>                                   |  |  |                                       |
| Comments: _____<br>_____<br>_____<br>_____<br>_____                               |  |  |                                       |

White: Sample Traveler

Canary: Lab Copy

Pink: Field Copy



|                      |                  |
|----------------------|------------------|
| Analysis: _____      |                  |
| Sample ID: _____     |                  |
| Laboratory ID: _____ |                  |
| Date Sampled: _____  |                  |
| Canister #: _____    | Press/Vac: _____ |
| Site: _____          | Dup/Rep: _____   |
| Comment: _____       |                  |

**Figure 22. ERG's Sample Tracking Tag**

|         |                                  |                        |
|---------|----------------------------------|------------------------|
| 8/29/19 | Entech passive Sampler: NP-26534 |                        |
|         | Sample Date: 8/30/19             |                        |
|         | Setup Date: 8/29/19              | Pickup Date: 9/3/19    |
|         | Sample ID: AK11563               |                        |
|         | Begin Vacuum: -30 inHg           | End Press/Vac: -3 inHg |
|         | Leak check: PASSED               |                        |
|         | Comment:                         |                        |
|         | Operator: <i>[Signature]</i>     |                        |

**Figure 23. Example of the GA AAMP Logbook Entry**

## 12.1 Pre-Sampling Custody

The pre-sampling custody is the sample handling stage that includes sample media purchasing, logging in, labeling, identification, pre-sampling weighing, transportation, and installation on

sampler. For GA AAMP's SOPs, see the applicable SOPs listed in Table 7 for more details. For the ERG Lab, see Laboratory Attachment, Section 9.1 for more details.

### **12.1.2 Sample Preparation**

Sample preparation is an essential portion of the ethylene oxide project. Cleaning, evacuation, testing, verification and storage of canisters are functions that are required for sample preparation.

Sample set-up of the ethylene oxide samplers take place any day after the previous sample has been recovered. Canisters for air collection for VOCs analyses must be used within 30 days after certified clean. Detailed sample set-up procedures are available from the corresponding GA AAMP's SOPs. For a description of ERG Lab's sample preparation, see Laboratory Attachment Section 10.0 of this document.

### **12.1.3 Sample Volume**

The volume of air to be sampled is specified by the manufacturer and is in the method specifications. Samples are expected to be 24 hours; therefore, the Site Operators must set the flow rates to collect a sufficient sample to obtain the minimum sample volume. In some cases, a shorter sample period may occur due to power outages. A valid sample run should not be less than 23 hours or greater than 25 hours. If the sample period is less than 23 hours or greater than 25 hours, the sample will be nulled and the Operations Unit Manager notified. The Entech passive sampler is complete when it reaches subambient pressure, typically -2 to -4 inHg (all other samplers i.e. the ATEC and Xonteck samplers should have an ending pressure of  $\geq +5$  psig).

## **12.2 Post Sampling Custody**

Post sampling procedures include: sample removal, field record keeping and transportation of samples, how to protect the sample from contamination, temperature preservation requirements, and the permissible holding times to ensure against degradation of sample integrity. See the applicable GA AAMP's SOP in Table 7, and for the ERG Lab, see Laboratory Attachment, Section 9.1 for more details.

### **12.2.1 Sample Contamination Prevention**

To prevent contamination during transport to the laboratory, the VOCs stainless steel canisters should be capped and handled to ensure that the valve to canister connection remains intact and the interior surface is not compromised.

### **12.2.2 Temperature Preservation Requirements**

During transport from the ERG Lab to the sample location, VOCs canisters have no specific requirements for temperature control per TO-15 Compendium Sections 1.3, 2.3, and 9.2.8.1.



### 12.2.3 Permissible Holding Times

The *Technical Assistance Document for the National Air Toxics Trends Station Program, Revision 3*, dated October 2016 states the permissible holding times for the VOCs samples. The VOC Canister analysis should be within 30 days of end of collection or preparation according to TO-15 Compendium Sections 1.3, 2.3, and 9.2.8.1.

### 12.3 Delivery to ERG Lab

Once the ethylene oxide samples are collected and prepared for delivery, the Site Operators send the samples to the ERG Lab via UPS, following protocol in applicable SOPs. When the samples are received at the ERG Lab, the chain-of custody form is filled in to record the sample receipt by Laboratory personnel. The ERG Lab analyst maintains records of sample preparation, analysis, and data input and management. See the applicable ERG Lab SOPs and Section 9.0 of the Laboratory Attachment for details.

### 12.4 Make-up Samples

Due to the number of sites involved in this study, there will not be make-up ethylene oxide samples taken. The frequency and duration of the sampling should ensure sufficient ethylene oxide data is available.

## 13.0 Analytical Methods

The method stated here provides for chromatographic analyses at the ERG Lab for samples collected at the GA AAMP ethylene oxide sites. The basic method used by ERG Lab is based on the Toxic Organic Compendia (TO-15) listed in Section 11.0. The sample media used to collect samples at ethylene oxide sites is a canister as shown in Table 1. In addition, the trip blank and laboratory blank must also be prepared. See Section 12.1.2 and the applicable ERG Lab's SOPs for more detail. The instruments used for laboratory analysis of the samples collected at the GA AAMP's ethylene oxide sites are listed in Table 8.

**Table 8. Instruments Used in the ERG Lab**

| Parameter | Instrument  | Method       |
|-----------|---|--------------|
| VOCs      | Agilent HP 8890/5977B with Entech 7200A interface<br>Agilent HP 6890/5973 with Entech 7200A interface | GC/MS, TO-15 |

### 13.1 Sample Contamination Prevention

The analytical support component of the ethylene oxide sites has rigid requirements for preventing sample contamination. To minimize contamination, the sample media clean-up and sample

preparation rooms are separate from the instrumentation rooms. In addition, heating and ventilation systems are checked by certified technicians. Hoods are also checked quarterly.

For the VOCs analytical method, the best prevention of contamination is not opening the canister in the laboratory. All post sampling Entech passive canisters that enter the ERG Lab should have subambient pressure of -2 to -4 inHg (all other samplers i.e. the ATEC and Xonteck samplers should have an ending pressure of  $\geq +5$  psig). Care must be taken when the canisters are under vacuum and stored in the laboratory. If there is a slight leak in the canister cap or valve, then laboratory air can enter into the canister and contaminate the run.

### **13.2 Temperature Preservation Requirements**

There are no temperature requirements.

### **13.3 Permissible Holding Times**

The permissible holding times for the ethylene oxide samples are detailed in the TO Compendia and the SOPs shown in Table 7.

## **14.0 Quality Control Requirements**

Quality Control (QC) is a means of periodic evaluation of the acceptability of the data. That is, does the data meet certain criterion. This section contains descriptions of the various QC checks which GA AAMP performs in conjunction with collecting ethylene oxide data. For a description of ERG Lab's quality control requirements, see Laboratory Attachment, Section 11.0.

### **14.1 Instrument Checks**

For this short-term study, the passive ethylene oxide monitor kits were sent to ERG Lab for collection and analysis of a zero sample, as well as a leak check, at the beginning of the study. The certification data is stored on the GA AAMP's local network for reference by anyone in the GA AAMP. The ATEC sampler was zero checked prior to this study as part of the NATTS Network requirements. For any samplers that were not zero checked, if high values are suspected due to a bias in the data, a comparison between that sampler and the passive sampler may be done for qualitative purposes. Each sampler will be uniquely identified. For a description of ERG Lab's calibration requirements, see Laboratory Attachment, Section 13.0.

The initial canister pressure must be checked prior to sample collection by measurement of the canister vacuum with a pressure gauge or pressure transducer. If a built-in gauge on the sampling unit cannot be calibrated, a standalone gauge will be employed for this measurement. This initial pressure will be documented on the sample collection form. Canisters must show  $\geq -28$  inHg.

Once vacuum is verified, the canister is connected to the sampling unit and a leak check is performed. A leak check may be performed by quickly opening and closing the valve of the canister to generate a vacuum in the sampling unit. The vacuum/pressure gauge in the sampling

unit will be observed for a minimum of 5 minutes to ensure that the vacuum does not change by more 1 inHg.

## 14.2 Precision Checks

One of GA AAMP's ethylene oxide samplers at each facility will be collocated with an additional sampler that will allow GA AAMP to make precision determinations. Collocated samplers operate monthly. There are two types of precision that will be determined for ethylene oxide data: collocated precision and replicate precision.

### 14.2.1 Precision Determination

Collocated precision evaluates the results of two monitors sampling side by side. The monitors separately operate at the same time and undergo the same sample collection, handling, and analysis procedures. In order to determine the precision, one compares results from the primary sampler concentration to the collocated sampler concentration by using the Relative Percent Difference noted below:

$$\text{Equation 14.2.1: Relative Percent Difference (RPD)} = \left[ \frac{ABS(Value_1 - Value_2)}{\frac{(Value_1 + Value_2)}{2}} \right] \times 100\%$$

The replicate precision is a measure of the reproducibility of the laboratory analyses. A replicate evaluation is performed on each batch by the ERG Lab with results sent to GA AAMP. A replicate is simply a re-analysis of the same canister of sample and then comparing the results of the replicate analysis to the first analysis. The ERG Lab will perform replicate analysis on 10% of samples. The percent RPD calculation for determining replicate precision is the same as the collocated calculation. Refer to the ERG's Laboratory Attachment for more details.

### 14.2.2 Precision Acceptance Criteria

Precision acceptance criteria are found in Section 7.2 of this QAPP.

### 14.2.3 Corrective Actions

Any non-conformances from the criteria specified in Section 14.2 above would be determined on a case-specific basis. In general, data validity for posting results on the GA EPD website purposes is a collective team effort and appropriate actions will be considered based on the circumstances. See the GA AAMP's *Standard Operating Procedure for Data Validation of Integrated Data* for further details. For a description of ERG Lab's corrective actions, see Laboratory Attachment, Section 16.3 of this document.

## 14.3 Quality Assurance Audits

An in-house technical systems audit (TSA) will be performed on the GA AAMP's ethylene oxide sampling equipment once per location per study. This will include a review of the Site Operators'

implementing SOPs, sampler maintenance, QC checks, and use of field logbooks and chain of custody forms. Audits are performed by the independent QA Unit in the GA AAMP. A summary report will be prepared by the Field Auditor. Please see the appropriate SOP shown in Table 7 for further details.

#### **14.4 Trip Blanks**

Trip blanks are collected for primary ethylene oxide samples once per month at the Sterigenics, (Cobb County) and Becton Dickinson (Covington) monitoring locations. Please see the GA AAMP's VOCs SOPs for details of the trip blanks. Trip blank acceptance criteria are found in Section 7.0 of this QAPP. Any non-conformances from the criteria specified in Section 7.0 would be determined on a case-specific basis. In general, data validity is a collective team effort and appropriate actions will be considered based on the circumstances. See the GA AAMP's *Standard Operating Procedure for Data Validation of Integrated Data* for further details.

### **15.0 Instrument/Equipment Testing, Inspection, and Maintenance Requirements**

#### **15.1 Maintenance**

The GA AAMP sends each passive ethylene oxide sampler to the ERG Lab for maintenance and leak check. This was conducted prior to beginning this study. For details of ERG's maintenance and leak check procedures, see ERG's Laboratory Attachment, Section 12.0. See the applicable Operations SOPs for maintenance of other ethylene oxide samplers.

#### **15.2 Instrument Check-In**

##### **15.2.1 Receipt from Maintenance**

When GA AAMP receives a VOCs monitor after it has undergone its maintenance, GA AAMP inspects the monitor for any damage during shipment. GA AAMP also turns on the unit and evaluates for proper operation.

##### **15.2.2 Zero Bias Check**

Please see the *Standard Operation Procedure for Operation of a Volatile Organic Compound (VOC) Canister Sampler for a National Air Toxics Trends Station* for further details of how to determine bias using ultra pure zero air or nitrogen.

#### **15.3 New Equipment**

When GA AAMP receives a new VOCs sampler, the same procedures will be used for instrument check-in as outlined in Section 15.2.

## **15.4 Spare Parts Inventory**

The GA AAMP maintains appropriate spare parts for the VOCs samplers. Primarily, GA AAMP has at least two backup monitors which are rotated through the maintenance program so that the GA AAMP has ample supply in case of failure of a critical part in a sampler. In addition, spare stainless steel valves and sample lines are available as needed.

## **15.5 Site Maintenance**

### **15.5.1 Cleaning of the Sample Inlets**

For the ethylene oxide samplers, leak checks will be performed before sampling. The particulate filter should be replaced as indicated by the final pressure on the canister. Pressure/vacuum indicates a blockage. Vacuum pressure gauges are calibrated initially before use. Documentation of these checks is stored on the GA AAMP local network drive.

### **15.5.2 Quality Assurance Audits**

An audit of the Site Operator's sample collection will be conducted at each of the five locations (Cobb County, Covington, South DeKalb, NR-285, and General Coffee) during the study. This audit will review equipment, adherence to the SOP, field documentation, and chain of custody records to ensure compliance with the GA AAMP's QAPP. The results of the audits (and any identified corrective actions) are summarized in a report to the QA Unit Manager.

## **16.0 Instrument Checks Frequency**

For the Entech CS 1200E Passive VOCs Canister Samplers at the sites near each facility and at the South DeKalb site, the ERG Lab performed a canister leak check and blank check on each canister prior to beginning this study. The initial canister pressure/vacuum is checked prior to each sampling. The initial pressure will be documented on the sample collection COC form. Canisters must show  $\geq -28$  inches Hg vacuum to conduct sampling. Once vacuum is verified, the canister is connected to the sampling unit and a leak check is performed. A leak check is performed in the field by quickly opening and closing the valve of the canister to generate a vacuum in the sampling unit. The vacuum/pressure gauge in the sampling unit will be observed for a minimum of 5 minutes to ensure that the vacuum does not change by more than 1 inHg. The vacuum/pressure gauges are calibrated initially before use, and on an as needed basis. Particulate filters are disposable and replaced if the sampling flow rate or final canister pressure/vacuum indicates a blockage or buildup of particulates.

For the Xontek Model 910 (NR-285), Xontek Model 911 (General Coffee) and ATEC 2200 (South DeKalb) VOCs Samplers, the GA AAMP uses a NIST traceable flow measurement device, a thermometer (if separate from flow meter), and barometer (if separate from flow meter). The calibration standards were sent to the supplier for NIST traceable certification prior to the study. An Excel spreadsheet is maintained by the GA AAMP to ensure that these standards are re-certified in a timely manner.

For a description of ERG Lab's calibration requirements, see Laboratory Attachment, Section 13.0 of this document.

## **17.0 Inspection, Acceptance, Requirements for Supplies and Consumables**

This section is not required for a Category II QAPP.

## **18.0 Non-Direct Measurements**

GA AAMP relies on the data that is generated through field and laboratory operations. However, other significant data is obtained from sources outside the GA AAMP or from historical records. This section addresses data not obtained by direct measurement from the GA AAMP. Possible databases and types of data and information that might be used include:

- Chemical and Physical Properties Data
- Sampler Manufacturers' Operational Literature
- Geographic Location Data
- External Monitoring Databases
- Population Data from the US Census Bureau
- Traffic Data from Georgia Department of Transportation
- Wind Roses and other atmospheric data from other meteorological stations
- Emission Inventory from EPA

Any use of outside data will be quality controlled to the extent possible following the QA procedure outlined in this document and in applicable EPA guidance documents.

### **18.1 Chemical and Physical Properties Data**

Physical and chemical properties data and conversion constants are often required in the processing of raw data into reporting units. This type of information that has not already been specified in the monitoring regulations will be obtained from the following nationally and internationally recognized sources. Other data sources may be used with approval of the Program Manager.

- National Institute of Standards and Technology (NIST)
- ISO, IUPAC, ANSI, and other widely-recognized national and international standards organizations
- EPA
- The current edition of certain standard handbooks may be used without prior approval of the QA Unit Manager

### **18.2 Sampler Operation and Manufacturers' Literature**

Another important source of information needed for sampler operation is manufacturers' literature. Operations manuals and users' manuals frequently provide numerical information and equations pertaining to specific equipment. The GA AAMP's personnel are cautioned that such information is sometimes in error, and appropriate cross-checks will be made to verify the reasonableness of information contained in manuals. Whenever possible, the Site Operators will compare physical

and chemical constants in the operations' manuals to those given in the sources listed above. If discrepancies are found, the applicable Operations Manager should be the one to determine the correct value by contacting the manufacturer. The following types of errors are commonly found in such manuals:

- Insufficient precision
- Outdated values for physical constants
- Typographical errors
- Incorrectly specified units
- Inconsistent values within a manual
- Use of different reference conditions than those called for in EPA guidance

### **18.3 Geographic Location**

Another type of data that will commonly be used in conjunction with the GA AAMP ethylene oxide project is geographic information. The GA AAMP locates the site using global positioning system (GPS) that meets the requirements in Appendix A of EPA's National Geospatial Data Policy (August 2005). Google Earth is used as the primary means for locating and siting sampling locations.

### **18.4 External Monitoring Databases**

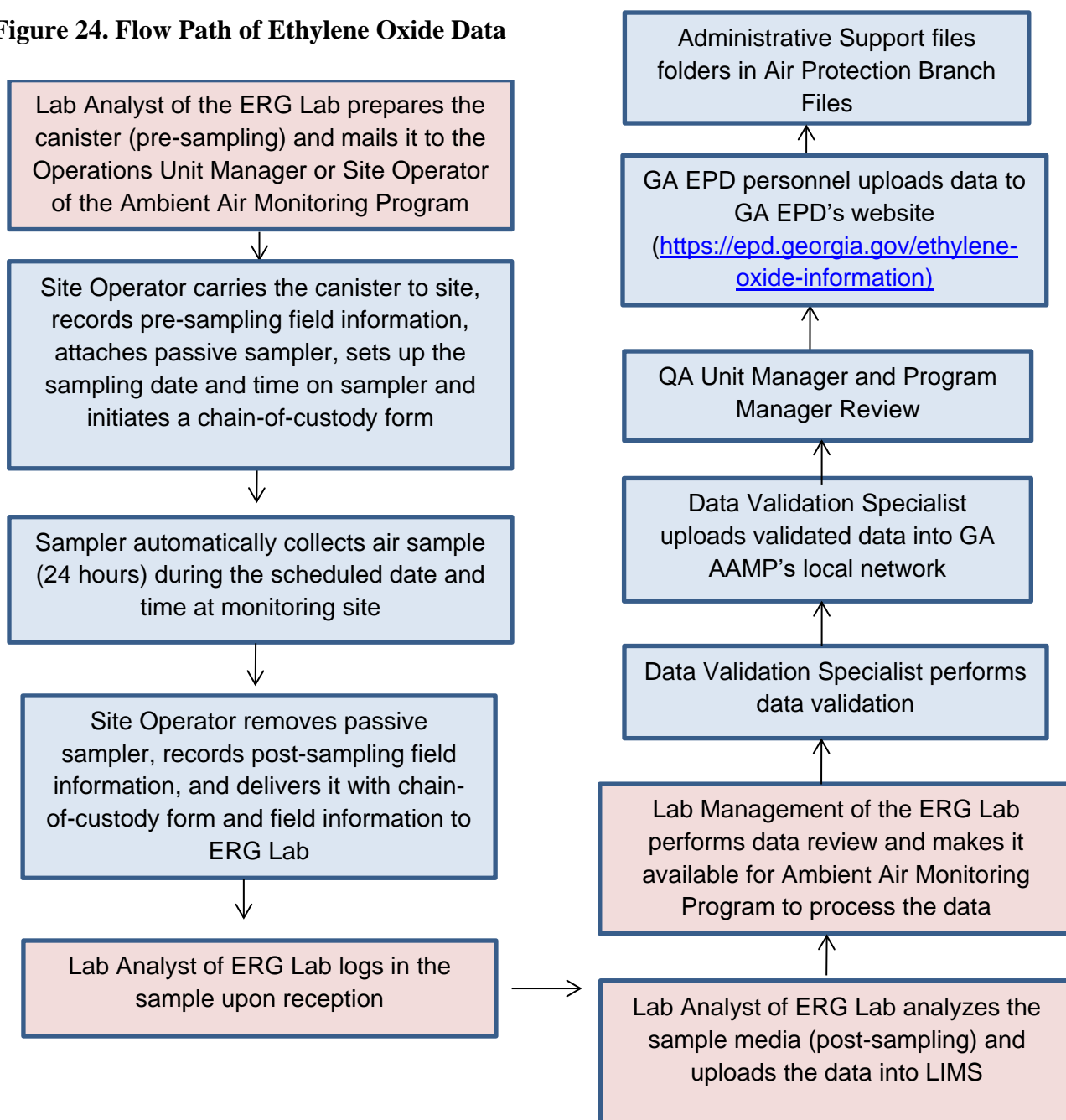
Data from the GA AAMP/GA EPD website may be used in published reports with appropriate caution. Care must be taken in reviewing and using any data that contain flags or data qualifiers. If data is flagged, such data shall not be utilized unless it is clear that the data still meets critical QA/QC requirements. It is impossible to assure that a database such as the GA AAMP/GA EPD website is completely free from errors including outliers and biases, so caution and skepticism is called for in comparing GA AAMP data from other reporting agencies. Users should review available QA/QC information to assure that the external data are comparable with GA AAMP measurements and that the original data generator had an acceptable QA program in place.

## **19.0 Data Management**

This section identifies the procedures that are followed to acquire, transmit, transform, reduce, analyze, store, and retrieve ambient air monitoring data by the field and office personnel of GA AAMP. The details of the processes and procedures in the ERG Lab are described in the ERG Lab's *Support for the EPA National Monitoring Programs (UATMP, NATTS, CSATAM, PAMS, and NMOC Support) QAPP*, and the ERG Laboratory Attachment of this document.

The following chart shows the flow of ambient air data collection process for the data. The collection and management of the data involves two operational entities: GA AAMP (blue blocks) and the ERG Lab (pink blocks). The GA AAMP performs the field activities and the ERG Lab conducts the analytical operations. For more description of ERG Lab's sample and data flow, see Laboratory Attachment, Section 15.0. In addition, please refer the applicable GA AAMP SOPs listed in Table 7 for more detail.

**Figure 24. Flow Path of Ethylene Oxide Data**





## **19.1 Data Collection and Recording**

The GA AAMP uses EPA-approved ambient air samplers for collection of ethylene oxide data. The canisters are collected manually and sent to the ERG Lab for analysis. The analysis results are saved in the ERG's Laboratory Information Management System (LIMS) and sent to the GA AAMP where the data is shared on the GA AAMP's local network. The leak check data is collected by the Site Operator and recorded on the local shared network for the Data Validation Specialist to review the data. The audit information is collected by the Field Auditor and recorded on the local shared network for the Site Operator and Data Validation Specialist to review.

## **19.2 Data Transmittal**

For the GA AAMP ethylene oxide data, all sampling media is sent back to the ERG Lab for analysis. Once the laboratory analysis is complete, the data is sent to GA AAMP office via email in a 'read only' portable document format (pdf) and an Excel file.

## **19.3 Data Review and Reduction (Validation)**

For ethylene oxide data, the ERG Lab analyzes the samples and summarizes the data as well as the corresponding QA/QC information in the ERG LIMS system and sends a copy to the GA AAMP. These files are 'read only' to ensure the data are not modified or deleted. The Data Validation Specialist reviews the laboratory data from the ERG Lab and the corresponding information on the chain-of-custody form and field data sheet. The holding time and delivery storage requirements for samples as listed in the SOPs shown in Table 7 must be followed; otherwise, the data will be invalidated. After completion of data review, the Data Validation Specialist prepares the final data associated with any applicable flags or null data codes into reportable data format and prepares a hard copy folder of the relevant information. For more detail, refer to the GA AAMP *Standard Operating Procedure for Data Validation of Integrated Data*.

## **19.4 Data Storage and Retrieval**

The storage and retrieval of the air quality monitoring data are conducted through the archiving system of GA EPD. The raw data is stored in the GA AAMP's local network (electronic data), and central file room (paper copy) for a period of at least five years, unless any litigation, claim, negotiation, audit, or other action involving the records has been started before the expiration of the five-year period. If this happens, the records will be retained until completion of the action and resolution of all issues that arise from it, or until the end of the regular five-year period, whichever is later.

The GA AAMP's raw data records that are housed on local network are only available to the GA AAMP staff. The raw data is then validated as discussed in the next Sections and posted to the GA EPD's website (<https://epd.georgia.gov/ethylene-oxide-information>).

## **20.0 Assessment and Response Actions**

Assessments are used to measure the performance and effectiveness of the quality system. These assessments and evaluations ensure the implementation of this QAPP, and that the ethylene oxide data is being collected for its intended use.

An in-house technical systems audit (TSA) will be performed on the GA AAMP's ethylene oxide sampling equipment. This will include a review of the Site Operators' implementing SOPs, sampler maintenance, QC checks, and use of field logbooks and chain of custody forms. These audits are performed by Field Auditor of the independent QA Unit in the GA AAMP. A summary report will be prepared by the Field Auditor.

The field assessments are performed as described in Section 14.0. The data validation will be performed as described in Sections 22.0 and 23.0. Detailed procedures of the quality assessment items can be found in the corresponding GA AAMP's SOPs (Table 7).

The laboratory assessments are performed as described in the ERG Lab's QAPP. For details of the ERG Lab assessments, see the ERG Lab QAPP attached. As EPA contract laboratory, the ERG Lab is subject to oversight by the EPA contract auditing group.

Although the GA AAMP produces quality data, the ethylene oxide data does not have to be certified by the GA AAMP Program Manager/Project Administrator, as the samplers are not SLAMS samplers.

## **21.0 Reports to Management**

With each set of ethylene oxide samples, a report summarizing the information will be sent to the GA AAMP and GA EPD management. The report will include a summary of sampling and analysis. Communication is an integral part of operating the GA AAMP ethylene oxide sites, and the status of the sites is directly communicated with the Site Operators, Operations Unit Manager, QA Unit Manager, and Program Manager as necessary. In addition, each of the Unit Managers meets with the Program Manager at least on a monthly basis to discuss pertinent issues.

## **22.0 Data Validation and Usability**

In order for the ethylene oxide data to be usable, the data undergoes validation procedures to determine that the data has met quality specifications. Validation, performed by Site Operators and Data Validation Specialists, can be defined as confirmation, through provision of objective evidence, that the particular requirements for a specific intended use are fulfilled. Site Operators and Data Validation Specialists evaluate the data to establish and confirm that the data was collected according to this QAPP and the SOP requirements. The Data Validation Specialist estimates the potential effect that any deviation from the QAPP and SOP may have on the usability of the associated data item, its contribution to the quality of the reduced and analyzed data, and its effect on decisions.

For GA AAMP, data validation is a process of reviewing and reducing raw data, with the use of objective evidence, to confirm requirements have been fulfilled and the intended use of the processed data for posting on the GA EPD's website (<https://epd.georgia.gov/ethylene-oxide-information>). The data validation process is based on sound documentation and checks. It is a systematic approach to produce reportable data that is accurate and complete. The GA AAMP performs data validation as data is received from the ERG Lab. It involves the data handling personnel of all units in GA AAMP as shown in the organization chart (Figure 1 in Section 4). Refer to the GA AAMP's *Standard Operating Procedure for Data Validation of Integrated Data* for more information.

## 22.1 Sampling Design

The GA AAMP chose the ethylene oxide monitoring sites according to emission models, wind rose data, proximity to the facilities, and proximity to the interstate or rural area as described in Section 6.0 and Section 10.0.

## 22.2 Sample Collection Procedures

The ethylene oxide sample collection procedures for the GA AAMP are outlined in Section 12.0 of this QAPP. The field audits discussed in Section 14.0 verify that the applicable SOPs listed in Table 7 are being followed when collecting samples. Potentially unacceptable data points are routinely identified through the application of error flags/codes. Each flag/code is associated with a unique error shown in Table 9. These error flags/codes are routinely reviewed as part of the data validation process. This activity assists in identifying suspect data points that could invalidate the resulting averaging periods. Any deviation from the established sampling criteria must be documented in the appropriate logbook and on the field data sheet. Accurate and complete documentation of any sample collection deviations will assist in any subsequent investigations or evaluations. Investigations and evaluations may be necessary to determine whether the data obtained from a particular site may qualify as a baseline or indicator for other sites.

**Table 9. Data Codes**

| <u>Null Codes</u> | <u>Description</u>                   |
|-------------------|--------------------------------------|
| AA                | Sample Pressure out of Limits        |
| AB                | Technician Unavailable               |
| AC                | Construction/Repairs in Area         |
| AD                | Shelter Storm Damage                 |
| AE                | Shelter Temperature Outside Limits   |
| AF                | Scheduled but not Collected          |
| AG                | Sample Time out of Limits            |
| AH                | Sample Flow Rate out of Limits       |
| AI                | Insufficient Data (cannot calculate) |
| AJ                | Filter Damage                        |
| AK                | Filter Leak                          |
| AL                | Voided by Operator                   |

|                        |  |
|------------------------|--|
| AM                     | Miscellaneous Void   |
| AN                     | Machine Malfunction  |
| AO                     | Bad Weather  |
| AP                     | Vandalism  |
| AQ                     | Collection Error   |
| AR                     | Laboratory Error   |
| AS                     | Poor Quality Assurance Results   |
| AT                     | Calibration  |
| AU                     | Monitoring Waived  |
| AV                     | Power Failure  |
| AW                     | Wildlife Damage  |
| AX                     | Precision Check  |
| AY                     | Q C Control Points (zero/span)   |
| AZ                     | Q C Audit  |
| BA                     | Maintenance/Routine Repairs  |
| BB                     | Unable to Reach Site   |
| BC                     | Multi-point Calibration  |
| BD                     | Auto Calibration   |
| BE                     | Building/Site Repair   |
| BG                     | Missing ozone data not likely to exceed level of standard              |
| BH                     | Interference/co-elution/misidentification                              |
| BI                     | Lost or damaged in transit   |
| BJ                     | Operator Error   |
| BK                     | Site computer/data logger down   |
| BM                     | Accuracy check   |
| BN                     | Sample Value Exceeds Media Limit                                       |
| BR                     | Sample Value Below Acceptable Range                                    |
| CS                     | Laboratory Calibration Standard  |
| DA                     | Aberrant Data (Corrupt Files, Aberrant Chromatography, Spikes, Shifts) |
| DL                     | Detection Limit Analyses   |
| FI                     | Filter Inspection Flag   |
| MB                     | Method Blank (Analytical)  |
| MC                     | Module End Cap Missing   |
| SA                     | Storm Approaching  |
| SC                     | Sampler Contamination  |
| ST                     | Calibration Verification Standard                                      |
| TC                     | Component Check & Retention Time Standard                              |
| TS                     | Holding Time Or Transport Temperature Is Out Of Specs.                 |
| XX                     | Experimental Data  |
| <u>Qualifier Codes</u> | <u>Description</u>   |
| 1                      | Deviation from a CFR/Critical Criteria Requirement                     |
| 2                      | Operational Deviation  |

|                    |  |
|--------------------|--|
| 3                  | Field Issue  |
| 4                  | Laboratory Issue   |
| 5                  | Outlier  |
| 6                  | QAPP Issue   |
| 7                  | Below Lowest Calibration Level   |
| 9                  | Negative value detected - zero reported                                |
| 1V                 | Data reviewed and validated  |
| CB                 | Values have been Blank Corrected                                       |
| CC                 | Clean Canister Residue   |
| CL                 | Surrogate Recoveries Outside Control Limits                            |
| DI                 | Sample was diluted for analysis  |
| EH                 | Estimated; Exceeds Upper Range   |
| FB                 | Field Blank Value Above Acceptable Limit                               |
| FX                 | Filter Integrity Issue   |
| HT                 | Sample pick-up hold time exceeded                                      |
| LB                 | Laboratory blank value above acceptable limit                          |
| LJ                 | Identification Of Analyte Is Acceptable; Reported Value Is An Estimate |
| LK                 | Analyte Identified; Reported Value May Be Biased High                  |
| LL                 | Analyte Identified; Reported Value May Be Biased Low                   |
| MD                 | Value less than MDL  |
| MS                 | Value reported is 1/2 MDL substituted.                                 |
| MX                 | Matrix Effect  |
| ND                 | No Value Detected  |
| NS                 | Influenced by nearby source  |
| QX                 | Does not meet QC criteria  |
| SQ                 | Values Between SQL and MDL   |
| SS                 | Value substituted from secondary monitor                               |
| SX                 | Does Not Meet Siting Criteria  |
| TB                 | Trip Blank Value Above Acceptable Limit                                |
| TT                 | Transport Temperature is Out of Specs.                                 |
| V                  | Validated Value  |
| VB                 | Value below normal; no reason to invalidate                            |
| W                  | Flow Rate Average out of Spec.   |
| X                  | Filter Temperature Difference out of Spec.                             |
| Y                  | Elapsed Sample Time out of Spec.                                       |
| <u>Inform Code</u> | <u>Description</u>   |
| IA                 | African Dust   |
| IB                 | Asian Dust   |
| IC                 | Chem. Spills & Industrial Accidents                                    |
| ID                 | Cleanup After a Major Disaster   |
| IE                 | Demolition   |
| IF                 | Fire – Canadian  |

|    |                               |
|----|-------------------------------|
| IG | Fire - Mexico/Central America |
| IH | Fireworks                     |
| II | High Pollen Count             |
| IJ | High Winds                    |
| IK | Infrequent Large Gatherings   |
| IL | Other                         |
| IM | Prescribed Fire               |
| IN | Seismic Activity              |
| IO | Stratospheric Ozone Intrusion |
| IP | Structural Fire               |
| IQ | Terrorist Act                 |
| IR | Unique Traffic Disruption     |
| IS | Volcanic Eruptions            |
| IT | Wildfire-U. S.                |
| J  | Construction                  |

Null codes are used when the data is not usable and needs to be invalidated.

Quality Assurance (“QA”) qualifier codes are input when there is an issue that may affect the data due to a procedural malfunction, or general quality assurance.

Informational qualifiers (“INFORM”) are only for informational purposes.

### **22.3 Sample Handling**

Pertinent deviations from established sample-handling protocols for each sample physically retrieved for monitoring sites and equipment must be recorded on the sample custody sheet assigned to each filter for collection and recorded in the applicable electronic database for all pollutants.

### **22.4 Analytical Procedures**

The ethylene oxide data is validated and verified utilizing both manual and electronic methods. Specific criteria are utilized at the ERG Lab with blanks, duplicates, replicates, and collocated samples to determine acceptable data, the minimum acceptable values, and other criteria that are indicative of valid qualifying data. The ERG Lab can flag suspect data utilizing the list provided in Table 9.

### **22.5 Instrument Check Procedures**

Refer to Section 16.0 for details regarding checking the sampling instruments. More information can be found in applicable Operations’ and Data Validation SOPs found in Table 7.

## **22.6 Quality Control Procedures**

Section 14.0 specifies the QC checks that are to be performed during sample collection, handling, and analysis. These include analyses of standards, blanks, spikes, and replicates, which provide indications of the quality of data being produced by specified components of the measurement process. For each specified QC procedure, the acceptance criteria and corrective action (and changes) should be specified. Data Validation Specialists should document the corrective actions that were taken, which samples were affected, and the potential effect of the actions on the validity of the data. More information regarding QC checks and corrective actions can be found in Section 14.0, as well as the applicable Operations' and Data Validation SOPs found in Table 7.

## **22.7 Data Reduction and Processing Procedures**

As mentioned in the above sections, internal technical systems audits will be performed to ensure the data reduction and processing activities mentioned in the QAPP are being followed. Data will be reviewed and final concentrations will be validated by the Data Validation Specialist. The data will also be reviewed to ensure that associated flags or any other data qualifiers have been appropriately associated with the data and that appropriate corrective actions were taken. Upon completion of adjustments and/or corrective actions, the Data Validation Specialist uploads the final monitoring data, along with any applicable null codes, to the GA AAMP's local shared drive. Also, he/she notifies the Data Analysis Unit Manager, Operations Unit Manager, Site Operator, and Quality Assurance Unit Manager with the results of validation. The final values uploaded to the local shared drive should match the independent spreadsheet. Then the final ethylene oxide data will be uploaded to GA EPD's website (<https://epd.georgia.gov/ethylene-oxide-information>) by GA EPD personnel.

## **23.0 Validation and Verification Methods**

For GA AAMP, data validation is a process of reviewing and reducing raw data, with the use of objective evidence to confirm requirements have been fulfilled. Data verification is the process of independently (QA) checking the processed data, and verifying, with objective evidence, the validity and intended use of the processed data for upload to GA EPD's website (<https://epd.georgia.gov/ethylene-oxide-information>). The data validation and verification process is based on sound documentation and valid Quality Control (QC) and Quality Assurance (QA) checks. It is a systematic approach to produce reportable data that is accurate and complete. GA AAMP performs data validation as the data is received from the ERG Lab. It involves the data handling personnel of all units in GA AAMP as shown in the organization chart (Figure 1 in Section 4). Refer to GA AAMP's *Standard Operating Procedure for Data Validation of Integrated Data* for more information.

The following outline shows steps involved in the data review. Data validation and verification are discussed in more detail below the outline.

Level 0 (Raw data review):

- Site Operator evaluates samples as they are collected and notes any anomalies observed with sample collection.

Level 1 (Data analyzed):

- Laboratory Analyst processes samples and notes any anomalies as samples are processed.

Level 2 (Data Validation):

- Data Validation Specialist reviews data from ERG Lab, field data sheets, COCs, etc., ensuring MQOs are met. Applies null data codes or qualifier codes, and prepares file for upload.

Level 3 (Data Verification):

- Quality Assurance Unit Manager and Program Manager review and sign to approve data for upload.

### **23.1 Data Validation**

The ERG Lab analyzes the ethylene oxide samples and posts the data in a spreadsheet in their LIMS system. Once the laboratory analysis is complete, the data is sent to GA AAMP office via email in a 'read only' portable document format (pdf) and an Excel file. The Data Validation Specialist reviews the data, as well as the corresponding QA/QC information and the corresponding information on the chain-of-custody form and field data sheet. The MQOs for the ethylene oxide samples as listed in Table 4 must be followed, otherwise the data will be flagged or invalidated appropriately, according to Table 9. After completion of data review, a data folder is then generated by the Data Validation Specialist as data is received from the ERG Lab for the next steps of data validation. Data will be reviewed to ensure that associated flags or any other data qualifiers have been appropriately associated with the data and that appropriate corrective actions were taken. Upon completion of adjustments and/or corrective actions, the Data Validation Specialist uploads the final monitoring data, along with any applicable null codes, to the GA AAMP's local shared drive and notifies the QA Unit Manager that the data is ready for his/her review. The final values uploaded to the local shared drive should match the independent spreadsheet. Also, the Data Validation Specialist notifies the Operations Unit Manager and Site Operator with the results of validation.

### **23.2 Data Verification and Upload**

The QA Unit Manager receives the folder prepared by the Data Validation Specialist and verifies the information therein. He/she ensures proper qualifying data codes or null data codes have been applied, and ensures data is acceptable and complete. The QA Unit Manager makes appropriate notation of review, and comments if any corrections need to be made by the Data Validation Specialist. The QA Unit Manager submits the data to the Program Manager for final approval, and the data is then forwarded through GA EPD management for posting on the GA EPD website (<https://epd.georgia.gov/ethylene-oxide-information>).



## **24.0 Reconciliation with User Requirements**

A preliminary data review will be performed to uncover potential limitations to using the data, to reveal outliers, and generally to explore the basic structure of the data. The next step is to calculate basic summary statistics, generate graphical presentations of the data, and review these summary statistics and graphs to determine if representativeness, comparability, completeness, precision, bias, and sensitivity, were met. Representativeness can be assessed with site location information and is based on potential sources and select weather station information. Comparability is based on method measure of the level of confidence with which one data set can be compared to another. Completeness is measured by the amount of valid sample data obtained compared to what was expected. Precision is determined from replicate collocated analyses. Sensitivity is demonstrated through MDLs.

If the sampling design and statistical tests conducted during the final reporting process show results that meet acceptance criteria, it can be assumed that the network design and the uncertainty of the data are acceptable. Further use of the data will include characterizing concentrations in potentially affected nearby neighborhoods based on method sensitivity.

To determine if the GA AAMP will continue sampling ethylene oxide data, a qualitative analysis of the data will be assessed. In addition, the GA AAMP will ensure that the MQOs for data completeness and percent difference are met.

## Revision History

| Versions of <i>Quality Assurance Project Plan for the Georgia Ambient Air Monitoring Program Ethylene Oxide</i> |                |   |
|---|----------------|---|
| Revision 0  | September 2019 | Original version  |
| Revision 1  | March 2020     | Edited language regarding placement of monitors, and collocated and spatial frequencies; added statement regarding placement of potential future monitors; specified trip blank locations; added GA AAMP Additional Sampling Sites Attachment |
| Revision 1.1  | June 2020      | Updated spatial description and wind direction in the GA AAMP Additional Sampling Sites Attachment  |

## References

Eastern Research Group. 2019. *Support for the EPA National Monitoring Programs (UATMP, NATTS, CSATAM, PAMS, and NMOC Support) QAPP*. Morrisville, Georgia. March 2019.

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Environmental Protection Agency. Code of Federal Regulations. <https://www.ecfr.gov/>

Environmental Protection Agency. 2018. *National Air Toxics Trends Station Work Plan Template*. (Revised March 2018).

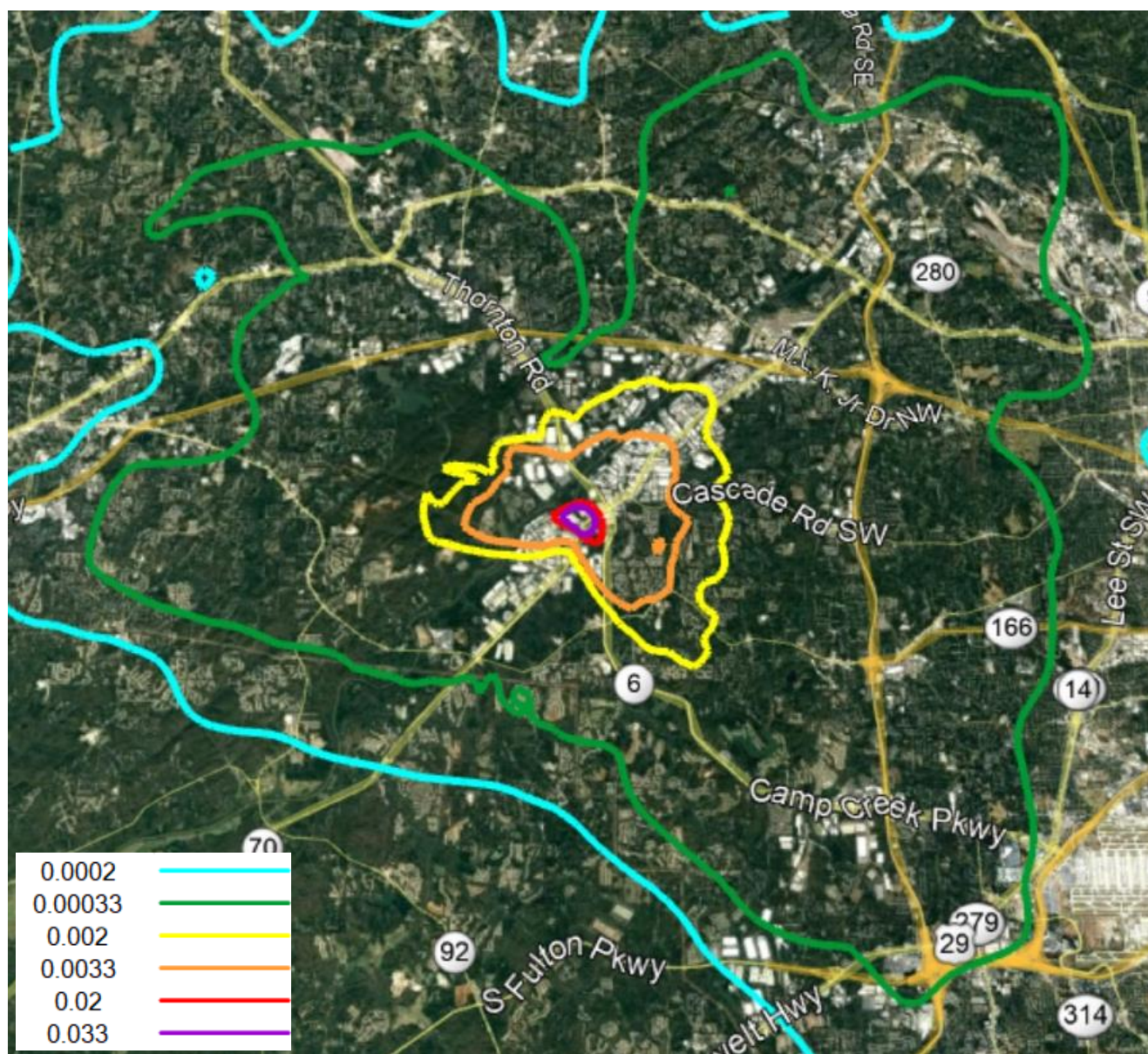
Environmental Protection Agency. 2018. *Quality Assurance Project Plan for Field Sampling Plan for Ambient Air Ethylene Oxide Monitoring Near Sterigenics Facility, Willowbrook, IL*. (November 2018).

[https://cfpub.epa.gov/ncea/iris2/chemicalLanding.cfm?substance\\_nmbr=1025](https://cfpub.epa.gov/ncea/iris2/chemicalLanding.cfm?substance_nmbr=1025)  
<https://www.epa.gov/national-air-toxics-assessment/2014-national-air-toxics-assessment>  
<https://www.epa.gov/air-emissions-inventories/2014-national-emissions-inventory-nei-data>

### **GA AAMP Additional Sampling Sites Attachment**

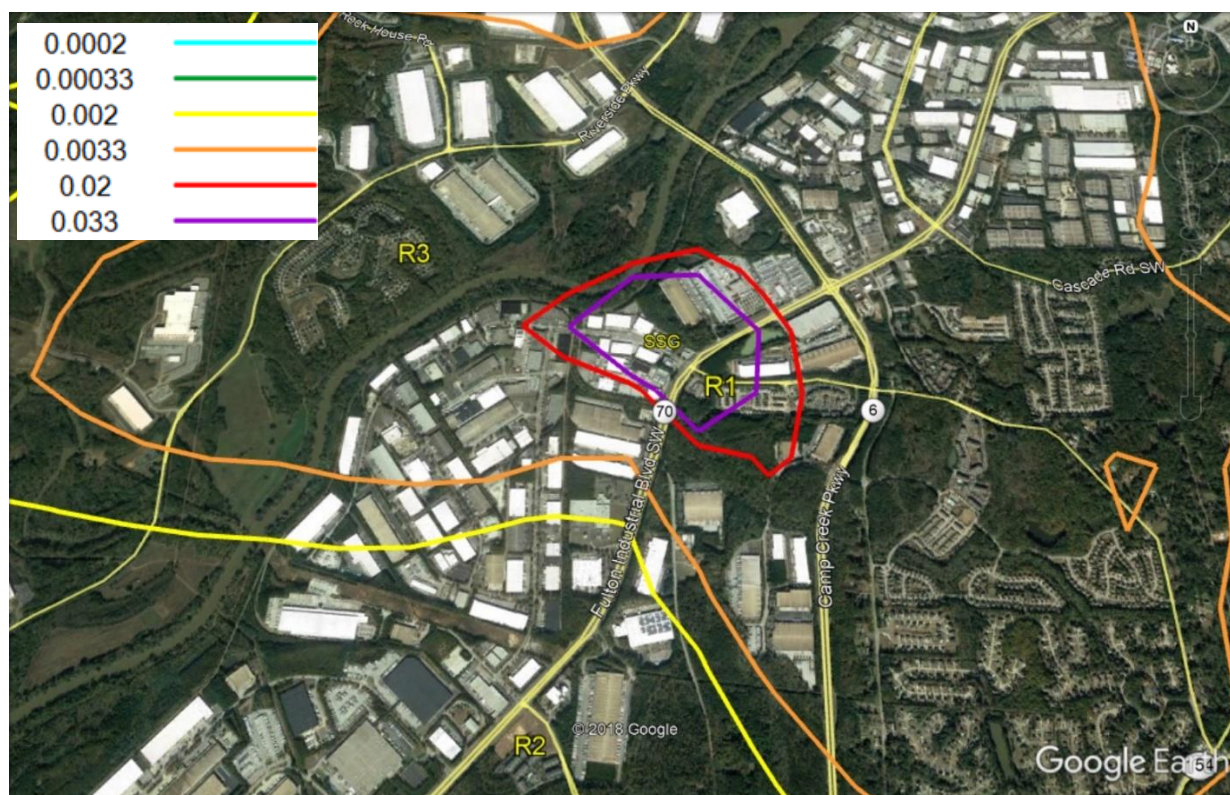
Sterilization Services of Georgia (SSG) in Fulton County is not a source of ethylene oxide that was identified by the 2018 NATA; however, the Planning and Support Program of GA EPD conducted modeling and SSG was modeled to have emissions above Georgia's Acceptable Ambient Concentrations (AACs). Computer models were used to predict the concentrations of toxic air pollutants (TAPs) being analyzed using facility information provided by the source and other information developed by GA EPD staff. The modeling results were compared to the 15-min, 24-hour, and annual AACs. GA EPD's 15-min and 24-hour AACs are derived from Occupational Safety and Health Administration (OSHA) permissible exposure limits (PEL), OSHA Total Weight Average (TWA) PEL. GA EPD's annual AACs are derived from U.S. EPA's risk values which are found in EPA's Integrated Risk Information System (IRIS) Risk Based Air Concentration (RBAC) database. The impacts of facility-wide ethylene oxide emissions were evaluated according to the Georgia Air Toxics Guideline.

To determine the ambient monitoring sites near SSG in Fulton County, GA, the GA AAMP considered the modeled emission data which was generated by the Planning and Support Program of GA EPD. Dispersion models of the ethylene oxide emissions data from the facility were conducted to determine concentrations of ethylene oxide around the facility. This model is shown in the following figures. The modeled values are shown in micrograms per cubic meter ( $\mu\text{g}/\text{m}^3$ ) and identified nearby residential areas with ethylene oxide concentrations above the annual AAC. Due to the limited availability of acceptable locations which can be considered ambient air in the industrialized area surrounding SSG, the GA AAMP identified two acceptable sampling locations within approximately  $\frac{1}{4}$  mile and  $\frac{1}{2}$  mile of the facility for the purposes of the ethylene oxide study. The sampling sites are both located in the primary downwind direction from SSG, and can also be used for spatial comparisons. These locations were selected due to areas of highest modeled concentrations and the identification of vulnerable populations in the vicinity of the facility. Both sampling locations are within or adjacent to a residential community, and no other acceptable monitoring sites were identified during the investigation.



**Figure 25. Contours of Modeled Annual Averaged Ground-level Concentrations Across the 5-Year Period (in  $\mu\text{g}/\text{m}^3$ ) for the Current Scenario Overlaid on a Google Earth Map**





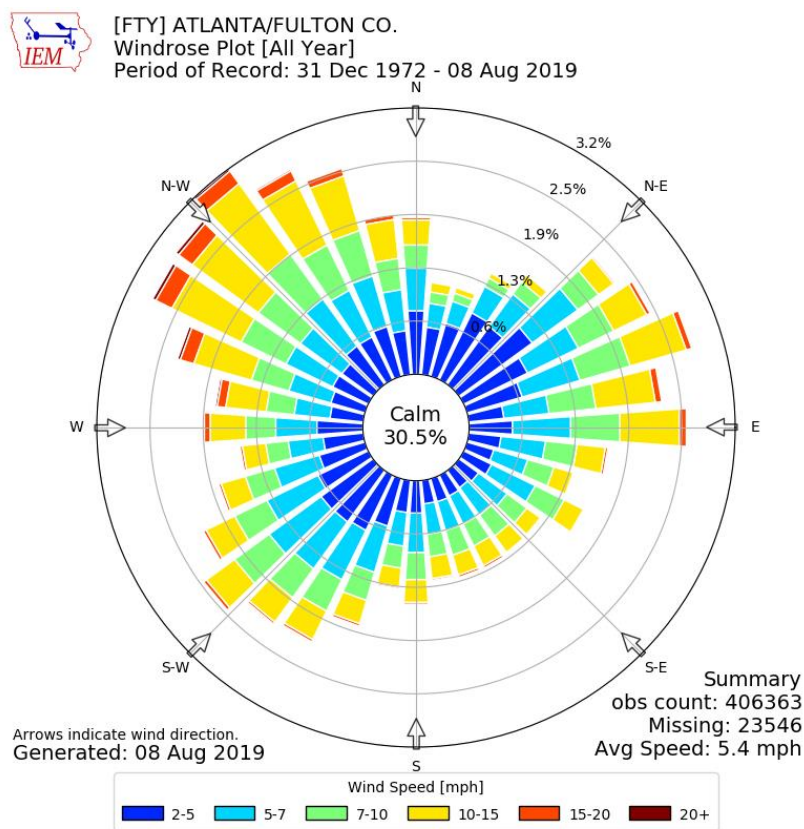
**Figure 26. A Close-up of Figure 25**

Wind rose data from the Atlanta Fulton County Airport near SSG was assessed by the GA AAMP, and primary and secondary wind patterns were determined. Distance from the Atlanta Fulton County Airport to SSG is shown in Figure 27 below. Wind rose data from the Atlanta Fulton County Airport is shown in Figure 28.





**Figure 27. Distance from SSG to Atlanta Fulton County Airport**



**Figure 28. Annual Wind Rose Data at Atlanta Fulton County Airport, December 31, 1970-August 8, 2019**

The GA AAMP began sampling near SSG on January 16, 2020. The sampling will follow the sampling schedule of once every six days as discussed for the other monitoring sites in Section 6.0 of this document. A collocated sample should be collected each month from the site nearest the residential community. Other than the number of sites used to characterize the ethylene oxide concentration around the facility, the quality assurance procedures described in this QAPP will be followed for the sampling in Fulton County. For the duration of the project, the sampling timeframe in Fulton County will be concurrent with the sampling timeframe for the Cobb County and Covington sites described in Section 6.0.

### **Laboratory Attachment**

ERG Laboratory QAPP available upon request.