

# **Exhibit 18**

**STUDY PROFILE for MRID 49888501**

Field Volatility of Dicamba Formulation MON 119096  
Following a Pre-Emerge Application Under Field  
Conditions in the Southeastern USA

**TEST GUIDELINE**

OPPTS 835.8100

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**DATE**

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## **Field Volatility of MON 119096 Herbicide**

**Report:** MRID 49888501: Field Volatility of Dicamba Formulation MON 119096 Following a Pre-Emerge Application Under Field Conditions in the Southeastern USA. 2016. Jacobson B., Urbanczyk-Wochniak E., Mueth M.G., Riter L.S., Sall E.D., Honegger J., South S., Carver L.

**Document No.:** MRID 49888501

**Guideline:** OCSPP 835.8100

**Statements:** This study was conducted according to the USEPA FIFRA Good Laboratory Practice (GLP) Standards (40 CFR Part 160), with the following exceptions:

### Field Phase:

- 1) Test site information such as estimation of slope, depth to water table elevation, and GPS coordinates.
- 2) Study weather data.
- 3) Pesticide and crop histories for the test plots.
- 4) Soil information provided by the USDA Natural Resource Conservation Service (NRCS).
- 5) Test plot preparation prior to application.

Data Confidentiality, GLP Compliance, Quality Assurance, and Authenticity Certification statements are provided.

**Classification:** This study is [provide classification and very concise statement of any deficiencies that impacted the classification]. [If multiple classification terminologies are needed for multilateral reviews, list or tabulate them.]

**PC Code:** [xxxxxxx]

**Reviewer:** [Provide final reviewer(s)'s name and title.]

**Signature:**

**Date:** [Type date of signature.]

### **Executive Summary:**

Field volatilization of MON 119096 Herbicide, a dicamba diglycolamine salt formulation containing 350 g dicamba acid equivalent (a.e.)/L applied in a spray solution containing 0.25% of the non-ionic surfactant Induce®, was examined from a bare plot at one site in Tift County, Georgia. The site where the study was conducted was near Chula, Georgia. The experiment at this site (Site 1/Plot 1) was conducted for approximately three days. The nominal application rate in the plot was 1 lb a.e./A (1.121 kg/ha).

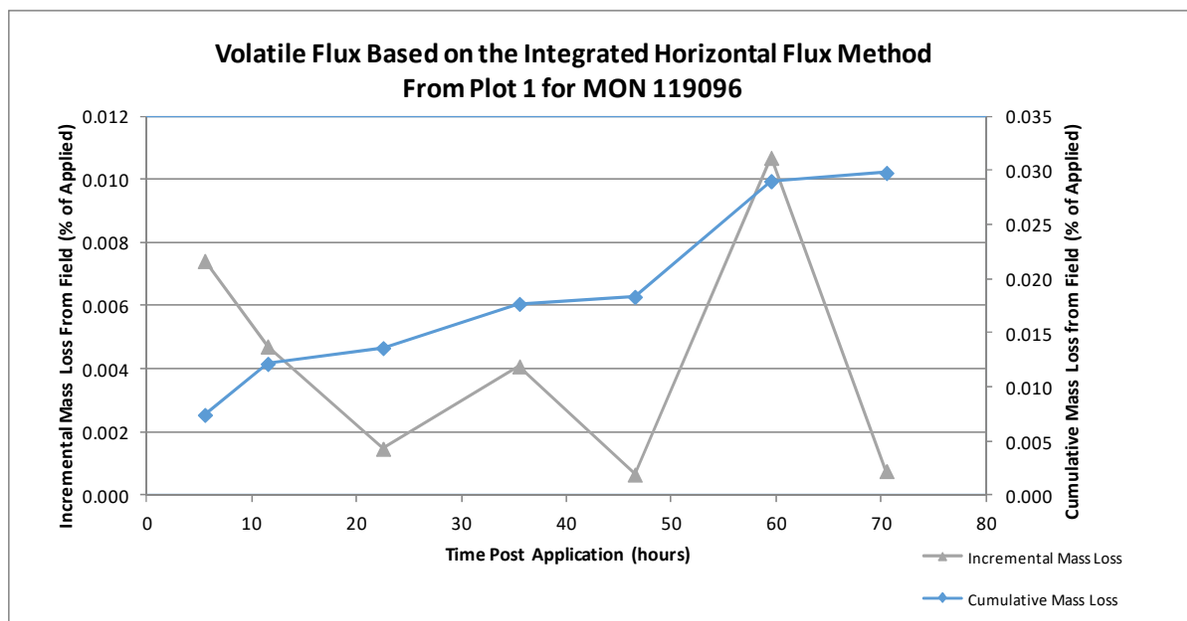
Under field conditions at Site 1/Plot 1 based on calculations using the Integrated Horizontal Flux method, dicamba in the MON 119096 formulation had a peak volatile flux rate of 0.000420  $\mu\text{g}/\text{m}^2\cdot\text{s}$  (0.000466  $\mu\text{g}/\text{m}^2\cdot\text{s}$ )<sup>1</sup> of the applied accounting for 0.007% (0.008 %) of the applied observed 0 – 5.5 hours post-application. By the end of the study, a total of 0.03% of dicamba

<sup>1</sup> Values for flux and percent of applied calculated by the registrant in the submitted report (MRID 49888501) are shown in parentheses, if different.

volatilized and was lost from the field after application. Other secondary peak volatile flux rates of 0.000244  $\mu\text{g}/\text{m}^2\cdot\text{s}$ , 0.000042  $\mu\text{g}/\text{m}^2\cdot\text{s}$ , 0.000098  $\mu\text{g}/\text{m}^2\cdot\text{s}$ , 0.000018  $\mu\text{g}/\text{m}^2\cdot\text{s}$ , 0.000256  $\mu\text{g}/\text{m}^2\cdot\text{s}$ , and 0.000022  $\mu\text{g}/\text{m}^2\cdot\text{s}$  (0.000271  $\mu\text{g}/\text{m}^2\cdot\text{s}$ , 0.000046  $\mu\text{g}/\text{m}^2\cdot\text{s}$ , 0.000088  $\mu\text{g}/\text{m}^2\cdot\text{s}$ , 0.000021  $\mu\text{g}/\text{m}^2\cdot\text{s}$ , 0.000276  $\mu\text{g}/\text{m}^2\cdot\text{s}$ , and 0.000023  $\mu\text{g}/\text{m}^2\cdot\text{s}$ ), occurred at 5.5 to 11.5, 11.5 to 22.5, 22.5 to 35.5, 35.5 to 46.5, 46.5 to 59.5, and 59.5 to 70.5 hours post-application, respectively. Higher flux rates were observed during the first day after application and during the daytime on Days 2 and 3. See Figure 1 for a graph of calculated incremental and cumulative mass loss from the treated area based on the Integrated Horizontal Flux Method.

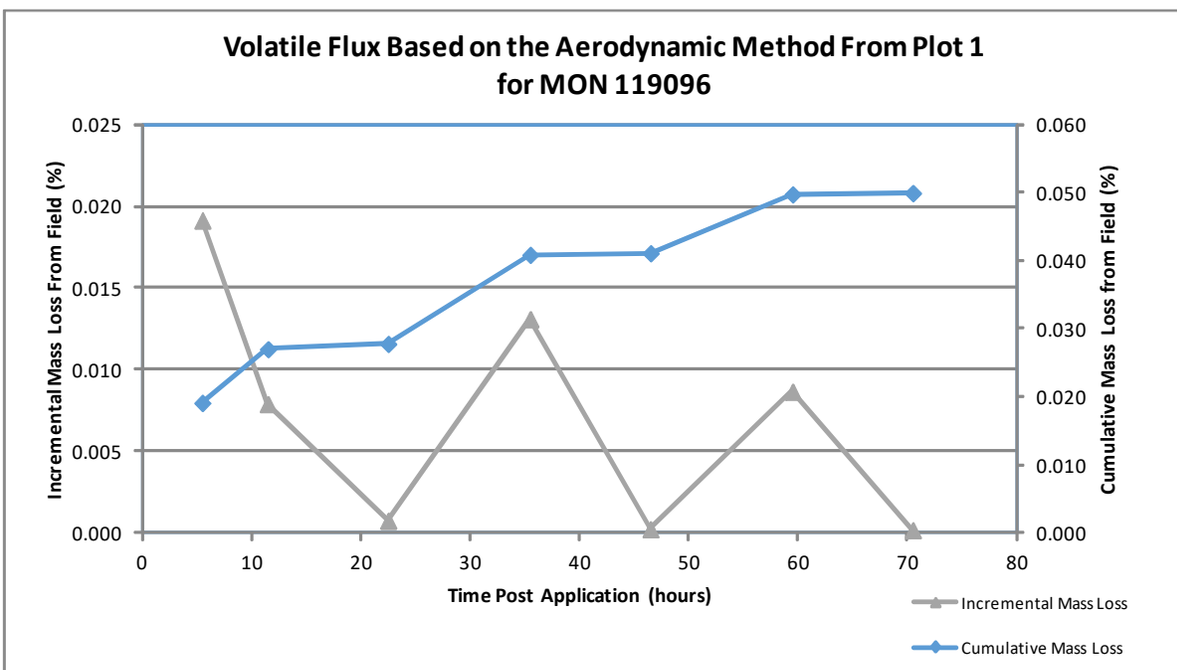
Under field conditions at Site 1/Plot 1 based on calculations using the Aerodynamic Flux method, dicamba in the MON 119096 formulation had a peak volatile flux rate of 0.00108  $\mu\text{g}/\text{m}^2\cdot\text{s}$  (0.00102  $\mu\text{g}/\text{m}^2\cdot\text{s}$ )<sup>2</sup> of the applied accounting for 0.019% (0.018 %) of the applied observed 0 to 5.5 hours post-application. By the end of the study, a total of 0.05% of dicamba volatilized and was lost from the field after application. Other secondary peak volatile flux rates of 0.000409  $\mu\text{g}/\text{m}^2\cdot\text{s}$ , 0.000021  $\mu\text{g}/\text{m}^2\cdot\text{s}$ , 0.000314  $\mu\text{g}/\text{m}^2\cdot\text{s}$ , 0.000007  $\mu\text{g}/\text{m}^2\cdot\text{s}$ , 0.000207  $\mu\text{g}/\text{m}^2\cdot\text{s}$ , and 0.000005  $\mu\text{g}/\text{m}^2\cdot\text{s}$  (0.000382  $\mu\text{g}/\text{m}^2\cdot\text{s}$ , 0.000019  $\mu\text{g}/\text{m}^2\cdot\text{s}$ , 0.000294  $\mu\text{g}/\text{m}^2\cdot\text{s}$ , 0.000006  $\mu\text{g}/\text{m}^2\cdot\text{s}$ , 0.000193  $\mu\text{g}/\text{m}^2\cdot\text{s}$ , and 0.000005  $\mu\text{g}/\text{m}^2\cdot\text{s}$ ), occurred at 5.5 to 11.5, 11.5 to 22.5, 22.5 to 35.5, 35.5 to 46.5, 46.5 to 59.5, and 59.5 to 70.5 hours post-application, respectively. Higher flux rates were observed during the first day after application and during the daytime sampling on Days 2 and 3. See Figure 2 for a graph of calculated incremental and cumulative mass loss from the treated area based on the Aerodynamic Flux Method.

Figure 1. Dicamba Mass Loss Based on the Integrated Horizontal Flux Method



<sup>2</sup> Values for flux and percent of applied calculated by the registrant in the submitted report MRID 49888501 are shown in parentheses, if different.

Figure 2. Dicamba Mass Loss Based on the Aerodynamic Flux Method



## I. Materials and Methods

### A. Materials

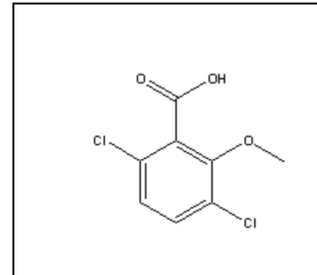
#### 1. Test Material

Product Name: MON 119096 Herbicide

Formulation Type: Liquid

CAS #: 104040-79-1

Storage stability: Stability of the active ingredient dicamba in the test substance MON 119096 has previously been demonstrated for 14 days at 55°C and for a related formulation one year under heated warehouse conditions.



#### 2. Storage Conditions

The test substance was shipped by Monsanto Company under ambient conditions on April 28, 2015 and received by Southeastern Ag Research, Inc. (SEAgRes) in Chula, Georgia on April 30, 2015. Upon receipt, the test substance was logged in and stored in a pesticide storage area at ambient temperatures ranging from approximately 66 to 73 °F prior to the application.

## B. Study Design

### 1. Site Description

The test site was located near Chula, Georgia in Tift County. A 3.4 acre (1.37 ha) plot was located in a larger field of bare soil. The plot dimensions were 384 ft x 384 ft (117m x 117m). The soil type of the plot was classified as a Tifton sandy loam, a very deep, well drained, moderately slowly permeable soil on interfluves that formed in loamy marine sediments. The taxonomic class of the Tifton series is – fine-loamy, kaolinitic, thermic Plinthic Kandiudults (Order – Ultisols, Suborder – Udults). The crops on this site for the previous three years were cotton (2012), cotton (2013), and peanut (2014). No herbicides containing dicamba had been used on the site for the last three years, although a number of other cotton and peanut herbicides, fungicides, and insecticides were used. The elevation of the site was approximately 370 feet above sea level, and the slope in the test area was estimated to be approximately 2%.

### 2. Application Details

Application rate(s): The target application rate was 1.0 lb a.e./A (1.121 kg/ha). Based on application monitoring samples in the treated area, the measured application rate was 113 ±26% of the target application rate. All nozzles were calibrated and pass times were within 2% of the target pass time.

Irrigation and Water Seal(s): No irrigation water was used.

Tarp Applications: Tarps were not used.

Application Equipment: A tractor-mounted boom sprayer was used for the broadcast application of Dicamba Formulation MON 119096 to the treated plot. The spray boom was fitted with sixteen Turbo Teejet Induction (TTI 04) nozzles and 50 mesh screens. The nozzles were spaced 18 inches apart, providing a 24-foot swath width. The boom height was set approximately 18 inches above the soil. The sprayer was equipped with a 80 gallon tank, and a roller pump (PTO driven) was used to pressurize the system and provide agitation (recirculation).

Equipment Calibration Procedures:

The sprayer output prior to application was calibrated using water and the spray from each nozzle was collected for 20 seconds in three separate runs. The boom pressure was set at approximately 27-28 psi for the calibration trial and test substance application. The calibrated sprayer output for the application was calculated to be 13.5 GPA. The target sprayer speed was 9.98 feet/sec or 6.80 mph.

Application Regime: The application rates and methods used in the study are summarized in **Table 1**.

**Table 1. Summary of application methods and rates for MON 119096**

Field	Application Method	Time of Application (Date and Start Time)	Amount Dicamba Applied (lbs)	Area Treated (acres)	Calculated Application Rate (lb ae/acre)	Reported Application Rate (lb ae/acre)
1	Broadcast Spray	5/5/2015 08:09	3.4	3.4	1.0	0.999* 1.13**

\*Based on sprayer pass timing

\*\*Based on analysis of deposition in the spray area.

Application Scheduling: Critical events of the study in relation to the application period are provided in **Table 2**.

**Table 2. Summary of MON 119096 application and monitoring schedule**

Field/Plot	Treated Acres	Application Period	Initial Air/Flux Monitoring Period	Water Sealing Period	Tarp Covering Period
Field 1 /Plot 1	3.4	05/05/15 between 08:05 – 8:30	05/05/15 between 8:30 – 14:00	Not Applicable	Not Applicable

### 3. Soil Properties

Soil properties measured before the study are provided in **Table 3**.

**Table 3. Summary of soil properties for fields/plots**

Field	Sampling Depth	USDA Soil Textural Classification	USGS Soil Series	WRB Soil Taxonomic Classification	Bulk Density (g/cm <sup>3</sup> )	Soil Composition
1	0– 6 inches	Sand	Tifton loamy sand	Fine-loamy, kaolinitic, thermic Plinthic Kandiodults (Order – Ultisols, Suborder – Udults)	1.45	% Organic Carbon = 0.31% % Sand = 90% % Silt = 6% % Clay = 4%

Figure 3 presents a plot of the soil surface temperature measured 1 mm below the soil surface during the time of the study. Figure 4 presents a plot of the volumetric water content (VWC) as fraction of total soil volume at a depth of 2 inches. Note that the VWC measurement is a relative measurement since the probe was not calibrated to the specific soil and the test site.

Figure 3. Soil Temperature at the Soil Surface (1 mm Depth) after Application

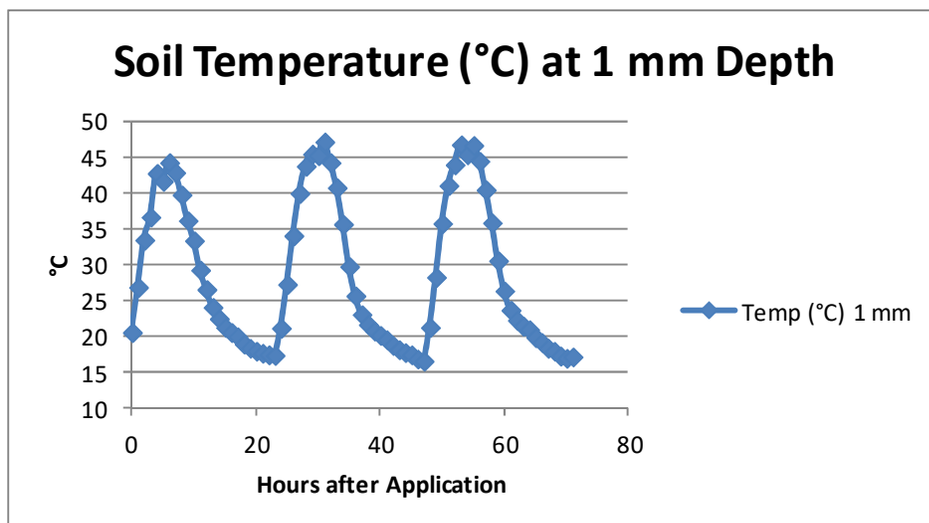
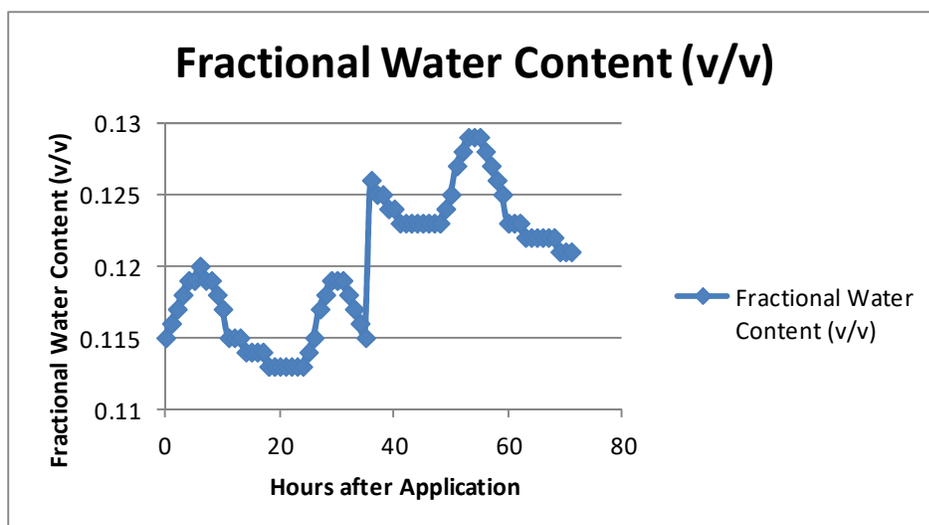


Figure 4. Soil Volumetric Water Content after Application



#### 4. Meteorological Sampling

Air temperature, precipitation, relative humidity, solar radiation, wind speed, wind direction, soil temperature (1 mm and 2 and 6 inches deep) and reference evapotranspiration, were recorded at the test site using an on-site weather station. A summary of hourly and daily climatological data including air temperature (max/min/avg), relative humidity (max/min/avg), ETo, wind speed (max/min/avg), average wind direction, solar radiation, soil temperature (max/min/avg), volumetric water content, and precipitation for the duration of the field volatility sampling is presented in Appendix 1 of the report.

One meteorological station was established near the plot so that a good representation of the wind pattern of the test plot was achieved (Figure 3 of the report). The station monitored air temperature (using air temperature sensors and radiation shields), and wind speed and wind direction (using sonic anemometers) located at approximately 0.33, 0.55, 0.90 and 1.5 m above the soil surface. Each parameter was measured every second, and the data were summarized every minute and every hour. The hourly wind speed and direction data for each monitoring height is summarized in Appendix 2 of the report. The hourly air temperature data for each monitoring height is summarized in Appendix 3 of the report. The temperature, wind speed, and wind direction at each sampling height summarized on a per minute basis is provided in the Raw Met Data worksheet in the Excel spreadsheets embedded in Attachment 2 of this document.

Details of the sensor heights and the meteorological parameters for which data were collected are illustrated in Table 4. The location of the meteorological equipment for the plot is shown in Attachment 3.

**Table 4. Summary of meteorological parameters measured in the field**

Field	Minimum Fetch* (m)	Parameter	Monitoring heights (m)	Averaging Period
1	64.8 (58.6)**	Wind speed/Wind direction	Height 0.33, Height 0.55, Height 0.90, Height 1.5	1 minute
		Ambient air temperature	Height 0.33, Height 0.55, Height 0.90, Height 1.5	1 minute

\* Value indicated is the minimum average fetch for a sampling period. Values on a per minute basis are calculated in the spreadsheets provided in Attachment 2.

\*\*Value in parenthesis is from registrant submitted report MRID 49888501. In this report mean fetch for a sampling period is calculated using the resultant wind direction over the sampling period rather than calculating a fetch value for each minute of the sampling period and then taking the mean of those values.

## 5. Air Sampling

The on-field air profile monitoring station in the plot consisted of air sampling pumps mounted on a sampling mast located at the approximate center of the plot. Five sample locations (heights) were established on the sample mast at approximately 0.15, 0.33, 0.55, 0.90 and 1.5 m above the soil surface. All air samples for a specific time increment were used for the regression analysis for that increment unless samples were disqualified by breakage or other defined cause. For each sampling time increment, a mass rate of chemical transfer per unit surface area (flux rate) was calculated. Two pre-application (PA) samples were collected at the 0.15 m height near the air monitoring location (center of plot) in the treated plot on May 4, 2015 prior to the start of the application on the following day. Samples were collected from 8:04 until 14:20.

## 6. Sample Handling and Storage Stability

Upon collection, the air sampling tubes were immediately placed in an ice chest with ice packs until they were transported to field cooperator's facility freezer. Transit stability samples were prepared/spiked by Monsanto Company, and then shipped frozen to the field site then stored and shipped in the same manner as the actual field samples. A set of air sampling tubes to be exposed in the field were prepared/spiked by Monsanto Company, and then shipped frozen to the field site. At the field site, these samples were removed from frozen storage, allowed to reach ambient temperature then exposed to an air flow then placed in frozen storage along with the

other field samples.

All samples were later shipped overnight via Federal Express in insulated containers with dry ice to Monsanto Company for analysis. The samples were received frozen and were maintained frozen except when a sample was being prepared for extraction. Samples were then stored in a continuously monitored freezer set at approximately -20 °C until analysis. All PUF samples in this study were extracted and analyzed within a period of 16 days or less after collection or spiking.

Stability of dicamba on PUF sample collectors has been demonstrated in a concurrent stability study. Samples were prepared by fortifying polyurethane foam (PUF) air sampling traps separately with dicamba at a concentration of 0.003 µg /PUF. Samples were removed from frozen storage and analyzed at intervals of 0, 7, 15, 55 and 78 days. The estimated degradation of dicamba was calculated using a statistical regression model. The statistical analysis of the change in measured concentration over time showed no statistically significant change in dicamba over the course of 78 days. Dicamba is therefore considered stable on PUF sample collectors under frozen storage conditions for at least a period of 78 days.

## 7. Analytical Methodology

- **Sampling Procedure and Trapping Material:** Dicamba was monitored using glass vapor collection tubes containing polyurethane foam (PUF) sorbent (SKC Inc. Catalog No. 226-92) and SKC® (Model Number 224-52) personal air sampling pumps. The vapor collection tubes were attached to a pump using Tygon® tubing. The flow for each sample was calibrated for each sample period using a Bios International Dry-Cal model DC-L12K. The flow was set to approximately 3 L per minute.
- **Extraction method:** The polyurethane foam sorbent (PUFs) was transferred from the air sampling glass tube to a 50 mL conical tube. Dicamba was extracted from PUFs using methanol containing stable-labeled internal standard. The sample tubes were capped and agitated on a high-speed shaker for extraction. An aliquot of the supernatant was filtered, evaporated, reconstituted in 25% methanol in water and then dicamba was quantitated using LC-MS/MS with electrospray ionization in negative ion mode.
- **Method validation (Including LOD and LOQ):** The method used for data collection was ME-1902. This procedure was validated in study by assaying 18 untreated control samples, 18 replicates each of the control samples fortified at 0.0003 (LOQ), 0.003 and 0.060 µg/PUF. Capability of dilution was demonstrated using 18 replicate samples fortified at 0.6 µg/PUF. Validation assessments demonstrated accuracy within 70-120% and precision ≤20% RSD at all fortification levels tested. The specificity of the method was demonstrated through the analysis of control matrices. The limit of quantitation (LOQ) of dicamba was 0.0003 µg/PUF determined during method development and confirmed through in-study method validation. The LOD was determined to be  $6.5 \times 10^{-5}$  µg/PUF through statistical determinations on validation data.
- **Instrument performance:** Calibration curves were linear with 1/x weighting. The working range of the method was from 0.0003 to 0.075 µg/PUF. Extension of the upper range of the

method to 0.75 µg/PUF can be achieved using lower injection volumes, therefore overall the method covered sample concentrations from 0.0003 to 0.75 µg/PUF.

## 8. Quality Control for Air Sampling

- Lab Recovery:** Most laboratory spike recoveries are within the acceptable range (90-110%) with overall recoveries between 90.2 to 102.7 percent. The exceptions include two laboratory spike samples with percent recovery of 78.5 and 82.8 at the fortification level of 0.0003 µg/PUF and two laboratory spike sample with percent recovery of 84.2% and 85% at fortification level of 0.6 µg/PUF.
- Field blanks:** There was no dicamba measured in the samples collected before application.
- Field Recovery:** Most field spike recoveries are within the acceptable range with overall recoveries between 97.7 and 110 percent. The exceptions are one sample extracted after six hours sampling time with percent recovery of 85.0 at the fortification level of 0.003 µg/PUF. One sample extracted after six hours sampling time with percent recovery of 83.3% at the fortification level of 0.6 µg/PUF, and three samples extracted after twelve hours sampling time with percent recovery of 82.0, 85.0 and 114 at the fortification level of 0.6 µg/PUF. Mean recoveries for six hour samples fortified at 0.003 µg/PUF, 0.03 µg/PUF and 0.6 µg/PUF were 95.7%, 103%, and 98.2%, respectively, and for the twelve hour samples were 102%, 103% and 93.6%, respectively, (n=3 for each fortification level).
- Travel Recovery:** Stability of dicamba on PUFs during transit of samples from the field was determined by sending three samples fortified at 0.003 µg/PUF and three samples fortified at 0.03 µg/PUF. The overall recoveries ranged between 94.3 to 106.0%.
- Breakthrough:** Samples that were fortified at 0.6 µg/PUF and exposed in the field for 12 hours had an average recovery of 93.6%. Samples fortified at 0.030 µg/PUF and exposed for 12 hours had an average recovery of 103%. The highest dicamba amount measured on a PUF was less than one third of the highest fortification indicating that dicamba loss due to breakthrough did not occur at the dicamba air concentrations present in this study.

## 9. Application Verification

Verification of the application rate was accomplished in two ways. First, the spray application rate was determined by measuring the time required for the tractor to drive the 117 m swath

length (pass time) for each pass and the calibrated volume output. The calculated spray application rate for each pass was consistent with the target dicamba application rate of 1.0 lb a.e./A with pass times and thus application rate varying by less than 2%. For the second method to verify the application of the test substance, ten (10) application monitoring devices/samples, pre-labeled pans each containing five (5) Whatman #3 filter papers each 12.5 cm in diameter (total area of 614 cm<sup>2</sup>) were placed diagonally across the plot area. As soon as possible after application the five filter papers in each pan were placed into a pre-labeled 500 mL wide-mouth, plastic bottle. Each bottle was sealed with a cap and taped. As soon as possible after collection (placed into bottle), the application monitoring samples were stored (while in the field) in a running, portable chest type freezer.

Recoveries achieved on extraction and analysis of application monitoring devices were in the range 64 to 157%, averaging  $113 \pm 26\%$  (n=10). Recoveries achieved on extraction and analysis of laboratory fortification of application monitoring filter papers ranged from 102 to 104%.

## II. Results and Discussion

### A. Empirical Flux Determination Method Description and Applicability

For estimation of flux after application in this study, both the Aerodynamic Method and the Integrated Horizontal Flux Method were used.

#### Aerodynamic Method

The aerodynamic method, also referred to as the “flux-gradient” method, was one of the two techniques employed for estimating flux rates from the field treated with MON 119096 for this field study given the available data. For the aerodynamic method, a mast was erected in the middle of the treated field, and samples for air concentration measurement were collected at five different heights, 0.15, 0.30, 0.55, 0.90, and 1.5 meters above the soil surface. Likewise, temperature and wind speed data were collected at 0.30, 0.55, 0.90, and 1.5 meters above the soil surface at a weather mast positioned just outside of the spray plot. Log-linear regressions were performed relating the natural logarithm of the sample height to the concentration, temperature, and wind speed. These relationships were then utilized to calculate input parameters for the equation to estimate flux.

The methods to estimate flux and related equations are presented in Majewski et al., 1990. The equation for estimating flux using the aerodynamic method in particular is the Thornthwaite-Holzman Equation, which is shown in the following expression:

$$\text{Equation AD-1} \quad P = \frac{k^2 (\Delta \bar{c})(\Delta \bar{u})}{\phi_m \phi_p \left[ \ln \left( \frac{z_2}{z_1} \right) \right]^2}$$

where P is the flux in units of  $\mu\text{g}/\text{m}^2\cdot\text{s}$ , k is the von Karman's constant (dimensionless  $\sim 0.4$ ),  $\Delta \bar{c}$  is the vertical gradient pesticide residue concentration in air (regressed) in units of  $\mu\text{g}/\text{m}^3$  between heights  $z_{\text{top}}$  and  $z_{\text{bottom}}$  in units of meters,  $\Delta \bar{u}$  is the vertical gradient wind speed (regressed) in units of m/s between heights  $z_{\text{top}}$  and  $z_{\text{bottom}}$ , and  $\phi_m$  and  $\phi_p$  are atmospheric momentum and vapor stability correction terms, respectively. The heights  $z_{\text{top}}$  and  $z_{\text{bottom}}$  are chosen to fall within the range of the sampler heights deployed in the field test, and, as indicated above, both the air concentration and wind speed values at these two heights are obtained from linear regression equations relating concentration and wind speed to the natural log of vertical height.

Following the conditions expected in the neutrally stable internal boundary layer characterized by an absence of convective (buoyant) mixing but mechanical mixing due to wind shear and frictional drag, a log-linear regression is performed relating the natural logarithm of the sample height to the concentration, temperature, and wind speed. The adjusted values of the concentration, temperature, and wind speed from this regression are incorporated into Equation AD-1 to arrive at Equation AD-2 which is ultimately used to compute the flux.

$$\text{Equation AD-2} \quad \text{Flux} = \frac{-(0.42)^2 (c_{z_{\text{top}}} - c_{z_{\text{bottom}}}) (u_{z_{\text{top}}} - u_{z_{\text{bottom}}})}{\phi_m \phi_p \ln \left( \frac{z_{\text{top}}}{z_{\text{bottom}}} \right)^2}$$

$\phi_m$  and  $\phi_p$  are internal boundary layer (IBL) stability correction terms determined according to the value of the Richardson number,  $R_i$ , which is calculated as shown in Equation AD-3:

$$\text{Equation AD-3} \quad R_i = \frac{(9.8)(z_{\text{top}} - z_{\text{bottom}})(T_{z_{\text{top}}} - T_{z_{\text{bottom}}})}{\left[ \left( \frac{T_{z_{\text{top}}} + T_{z_{\text{bottom}}}}{2} \right) + 273.16 \right]^3 * (u_{z_{\text{top}}} - u_{z_{\text{bottom}}})^2}$$

where  $T_{z_{\text{top}}}$  and  $T_{z_{\text{bottom}}}$  are the regressed temperatures at the top and bottom of the vertical profile in units of  $^{\circ}\text{C}$ .

if  $R_i > 0$  (for Stagnant/Stable IBL)

$$\phi_m = (1 + 16R_i)^{0.33} \text{ and } \phi_p = 0.885(1 + 34R_i)^{0.4}$$

if  $R_i < 0$  (for Convective/Unstable IBL)

$$\phi_m = (1 - 16R_i)^{-0.33} \text{ and } \phi_p = 0.885(1 - 22R_i)^{-0.4}$$

### **Special Comments regarding EPA and Registrant's calculations for the flux estimation by the AD Method**

Important special comments relating to the Registrant's use of the AD Method in the present submission are:

1. The original articulation of Equation AD-3 in this template document specified the denominator as the sum of the two bracketed terms, when in fact it should be specified as the product. This has been corrected in the text. The calculation in the EPA spreadsheet embedded with this document did correctly multiply the two terms.
2. While the equations above indicate that the exponent for calculating  $\phi_m$  is -0.33 or +0.33 (depending on the value of  $R_i$ ), the Excel spreadsheet for the aerodynamic method provided with this template uses the value +/- 0.30483 for this exponent. For the flux values provided by the Registrant in Table 5, the exponent value of +/- 0.33 has been used.
3. The flux values provided in Table 5 as Registrant values use 0.40 as the von Karman's constant (see, for example, Majewski, et. al., 1990) while the Aerodynamic Method calculations in the example spreadsheet use a value of 0.42.
4. The Registrant has corrected two cell assignment errors that are present in the example Excel spreadsheet for the aerodynamic method.
  - i. In the Conc. Summary worksheet, the regression formulas in Columns I, J, and K, Row 3, reference the sampler heights specified by the User in Columns C through F, Row 2. However, the reference to the Row 2 Heights is lost when Row 3 of Columns I, J, and K, is copied down to other rows (sampling periods). This is because the reference to the Row 2 Heights is not specified as an absolute reference.

Thus, as an example, the formula in Column J, Row 3 of the Conc. Summary worksheet should be “=INTERCEPT(C3:F3, LN(C\$2:F\$2))”, wherein the addition of the “\$” designation in the Row 2 references ensures that the reference to sampler heights always points to Row 2 when the formula is copied down to subsequent sampling periods. This correction has been applied to Columns I, J, and K in the Conc. Summary worksheet.
  - ii. Similar referencing corrections have been applied to the wind and temperature regression formulas in Columns I, J, and K of the Met Data Summary worksheet to ensure that the appropriate row is specified when accessing the sampler height data.
5. The minimum fetch requirement that the fetch is 100 times the highest height of the air sampler for this method to be valid was not satisfied for the samplers positioned at 0.90 and 1.5 meter above the soil surface for any of the sampling times; however, this treated plot was bare soil, and the separation of the plot from obstacles such as trees that would impact air flow was indeed sufficient to establish excellent linearity for the regression of temperature and wind speed with the natural log of sampler height. Also, flux and total

mass loss values are in very good agreement between the aerodynamic flux method and the Integrated Horizontal Flux Method. This agreement indicates that the fetch was of sufficient length for the aerodynamic method to be valid for this plot.

### Integrated Horizontal Flux Method

The integrated horizontal flux method, also referred to as the “mass balance” method, was used as the second technique employed for estimating flux rates from the field treated with MON 119096 for this field study given the available data. For the integrated horizontal flux method, a mast was erected in the middle of the treated field, and samples for air concentration measurement were collected at five different heights, 0.15, 0.30, 0.55, 0.90, and 1.5 meters above the soil surface. Likewise, temperature and wind speed data were collected at 0.30, 0.55, 0.90, and 1.5 meters above the soil surface at a weather mast positioned just outside of the spray plot. A log-linear regression was performed relating the natural logarithm of the sample height to the air concentration and wind speed following the log linear relationships for the atmospheric boundary layer. Subsequent wind speeds and air concentrations required as inputs to the equation used to estimate flux are then determined from these regression equations.

The methods to estimate flux and related equations are presented in Majewski et al., 1990. The equation for estimating flux using the integrated horizontal flux (IHF) method in particular is given by the following expression:

$$\text{Equation IHF-1} \quad P = \frac{1}{x} \int_{Z_0}^{Z_p} \bar{c} \bar{u} dz$$

where P is the volatile flux in units of  $\mu\text{g}/\text{m}^2 \cdot \text{s}$ ,  $\bar{c}$  is the average pesticide residue concentration in units of  $\mu\text{g}/\text{m}^3$  at height Z in units of meters,  $\bar{u}$  is the average wind speed in units of m/s at height Z, x is the fetch of the air trajectory blowing across the field in units of meters for the sampling period in question,  $Z_0$  is the aerodynamic surface roughness length in units of meters,  $Z_p$  is the height of the plume top (the height at which air concentration falls to zero) in units of meters, and dz is the depth of an incremental layer in units of meters.

The text and the embedded spreadsheet that accompany this template document describe a numerical evaluation of the integral in Equation IHF-1 by application of the trapezoidal integration technique. In this case Equation IHF-1 is modified to the summation shown in Equation IHF-2, below (Yates, 1996):

$$\text{Equation IHF-2} \quad P = \frac{1}{x} \sum_{Z_0}^{Z_p} (A * \text{Ln}(z) + B) * (C * \text{Ln}(z) + D) dz$$

where A is the slope of the wind speed regression line by  $\ln(z)$ , B is the intercept of the wind speed regression line by  $\ln(z)$ , C is the slope of the concentration regression by  $\ln(z)$ , D is the intercept of the concentration regression by  $\ln(z)$ , and z is the height above ground level. The embedded spreadsheet (and the Registrant) interprets  $Z_0$  as the aerodynamic surface roughness, i.e., the point at which the wind speed profile extrapolates to zero velocity. In turn, the material supplied with this template document calculates  $Z_p$ , the height of the concentration plume, using Equation IHF-3, below.

$$\text{Equation IHF-3} \quad Z_p = \exp\left[\frac{(0.1 - D)}{C}\right]$$

However, the Registrant is not familiar with the derivation behind this equation for evaluating  $Z_p$  and uses instead the alternative (but similar) equations IHF-6 and IHF-7, as described in Comment 2 of the following Special Comments section.

### **Special Comments regarding EPA and Registrant's methods and calculations**

Important special comments relating to the Registrant's use of the IHF Method in the present submission are:

1. The linear equations relating the wind speed and air concentration to the natural log of the height, Z, afford an analytical solution to the integral specified in Equation IHF-1. The Registrant utilizes this analytical solution to evaluate Equation IHF-1, rather than the numerical integration through application of the Trapezoidal Rule, represented in Equation IHF-2 and used in the provided embedded IHF spreadsheet.

Thus, if

$$\text{Equation IHF-4} \quad P = \frac{1}{x} \int_{Z_0}^{Z_p} (C \ln(z) + D)(A \ln(z) + B) dz$$

Then

*Equation IHF-5*

$$P = \left[ \frac{A \cdot C}{x} (z(\ln(z))^2 - 2z \ln(z) + 2z) + \frac{(A \cdot D + C \cdot B)}{x} (z \ln(z) - z) + \frac{(B \cdot D)}{x} z \right]_{Z_0}^{Z_p}$$

In which the symbols have the same meanings as those described for Equation IHF-2, above.

2. Regarding  $Z_p$ , the upper limit of the indicated integration –

The Registrant has not seen reference to an equation for calculating the height of the concentration plume that includes the subtraction of D from 0.1 in the numerator of the bracketed fraction seen in Equation IHF-3. Instead, the Registrant calculates the plume height as that value of Z at which the air concentration falls to zero. This is found by specifying a zero concentration in the regressed linear equation relating air concentration to the natural log of the height, Z, as seen in Equation IHF-6 below. Solving Equation IHF-6 for  $Z_p$  yields Equation IHF-7. It is this value for  $Z_p$  that the Registrant uses for the upper limit of the integral of Equation IHF-1. These  $Z_p$  are also entered as  $Z_{\max}$  in the IHF Flux Analysis worksheet, Column N.

$$\text{Equation IHF-6} \quad 0 = C \cdot \ln(Z_p) + D$$

$$\text{Equation IHF-7} \quad Z_p = \exp\left[\frac{-D}{C}\right]$$

3. Regarding  $Z_0$ , the lower limit of the indicated integration –

As specified in the example calculations provided with this template, the Registrant uses the Aerodynamic Surface Roughness, termed  $Z_0$ , as the lower limit for the integral of Equation IHF-1.  $Z_0$  is the height at which the wind speed profile extrapolates to zero. In the present case this is found by specifying a zero wind speed in the regressed linear equation relating wind speed to the natural log of the height, Z, as seen in Equation IHF-8 below. Solving Equation IHF-8 for  $Z_0$  yields Equation IHF-9. It is this value for  $Z_0$  (for each sampling period) that the Registrant uses for the lower limit of the integral of Equation IHF-1. These  $Z_0$  are also the values entered as the Aerodynamic Surface Roughness in the IHF Flux Analysis worksheet, column M.

$$\text{Equation IHF-8} \quad 0 = A \cdot \ln(Z_0) + B$$

$$\text{Equation IHF-9} \quad Z_0 = \exp\left[\frac{-B}{A}\right]$$

4. As in the aerodynamic method spreadsheet, the regression equations in the Conc. Summary and the Met Data Summary worksheets contain non-absolute cell assignment errors when referencing the user specified sampling heights. See Comment 4 in the Aerodynamic Method Section above. These errors have been corrected by specifying the sampling height references as absolute row addresses.
5. The minimum fetch requirement of 20 meters for this method to be valid was satisfied at all times. The treated field was a bare soil field with an estimated slope of about 2% . The maximum surface roughness was therefore very much less than 0.1 meters, thus this method validity criterion was met.
6. Regarding the value of x, the average fetch distance for a given sampling period –

The Registrant and the example IHF spreadsheet embedded in this template document approach the calculation of  $x$  in slightly different manners.

For any given sampling period (Event), the Registrant calculates the average fetch by first calculating the wind speed weighted average wind angle over the course of the Event time window. This average velocity weighted wind angle is then used in a single calculation to calculate the average fetch. The embedded IHF spreadsheet, on the other hand, calculates average fetch by first calculating the fetch associated with the minute by minute wind angle data compiled in the Raw Met Data worksheet and then averages those results over the time period of the sampling event. In more concise terms, for each Event the Registrant calculates fetch from the wind speed weighted average of the raw wind direction data, while the embedded spreadsheet calculates fetch as the average of the raw fetch data.

The distinctions described above lead to minor differences in the fetch values used in IHF-5 when calculating flux by the IHF method.

## **B. Temporal Flux Profile**

The flux as determined by the registrant as reported in MRID 49888501 and as determined by using the Excel spreadsheet provided by EPA in the Study Profile Template for the Field Volatility study (<http://www.epa.gov/pesticide-registration/study-profile-templates#fate>) for each sampling period after the application is provided in **Table 5**. Note that the Excel spreadsheet used for the flux calculation has been reviewed as described in Section A above to ensure that cell assignments are correct for each calculation.

**Table 5. Field volatilization flux rates of MON 119096 obtained in Georgia study, 1 lb a.e./A application rate**

Sampling Period	Date/ Time	Sampling Duration (hours)	Flux Estimate			
			EPA Spreadsheet ( $\mu\text{gm}^{-2}\text{sec}^{-1}$ )	Registrant Report ( $\mu\text{gm}^{-2}\text{sec}^{-1}$ )	Empirical Flux Determination Method	Notes
1	05/05/15 8:30 – 14:00	5.5	0.001083 0.000420	0.001017 0.000466	AD IHF	
2	05/05/15 14:00 – 20:00	6.0	0.000409 0.000244	0.000382 0.000271	AD IHF	
3	05/05/15 20:00 – 05/06/15 7:00	11	0.000021 0.000042	0.000019 0.000046	AD IHF	
4	05/06/15 7:00 – 20:00	13	0.000314 0.000098	0.000294 0.000088	AD IHF	
5	05/06/15 20:00 – 05/07/15 7:00	11	0.000007 0.000018	0.000006 0.000021	AD IHF	
6	05/07/15 7:00 – 20:00	13	0.000207 0.000256	0.000193 0.000276	AD IHF	
7	05/07/15 20:00 – 05/08/15 7:00	11	0.000005 0.000022	0.000005 0.000023	AD IHF	

\*Methods legend: AD = Aerodynamic Method, IHF = Integrated Horizontal Flux.

### III. Study Deficiencies and Reviewer's Comments

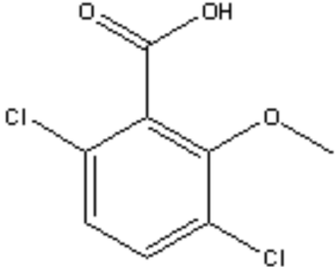
Left blank

### IV. References

Johnson, B., Barry, T., and Wofford P. 1999. Workbook for Gaussian Modeling Analysis of Air Concentrations Measurements. State of California Environmental Protection Agency, Department of Pesticide Regulation. Sacramento, CA.

- Majewski, M.S., Glotfelty, D.E., Kyaw Tha Paw U., Seiber, JN. 1990. A field comparison of several methods for measuring pesticide evaporation rates from soil. *Environmental Science and Technology*, 24:1490-1497.
- Wilson, J.D., and Shum. W.K.N. 1992. A re-examination of the integrated horizontal flux method for estimating volatilisation from circular plots. *Agriculture Forest Meteor.* Vol 57:281-295.
- Yates, S.R., F.F. Ernst, J. Gan, F. Gao, and Yates, M.V. 1996. Methyl Bromide Emissions from a Covered Field: II. Volatilization," *Journal of Environmental Quality*, 25: 192-202.

## Attachment 1: Chemical Names and Structures

Code Name/ Synonym	Chemical Name	Chemical Structure
<b>PARENT</b>		
<b>MON 119096/ Dicamba as the diglycolamine salt</b>	<b>IUPAC:</b> 3,6-dichloro- <i>o</i> - anisic acid  <b>CAS:</b> 3,6-dichloro-2- methoxybenzoic acid  <b>CAS No.:</b> 104040-79-1  <b>Formula:</b> C <sub>8</sub> H <sub>6</sub> Cl <sub>2</sub> O <sub>3</sub>  <b>MW:</b> 221.04 g/mol  <b>SMILES String:</b>	  <chem>CLc1ccc(CL)c(OC)c1C(=O)(O)</chem>

**Attachment 2: Statistics Spreadsheets and Graphs**

The electronic spreadsheet files are inserted below for calculations using the Aerodynamic Method and the Integrated Flux Method for determination of the emission rate of dicamba after application in the form of the diglycolamine salt in the formulation MON 119096.

1. Validation spreadsheet for studies following the Aerodynamic Method:



field-volatility-835-81  
00-MRID\_49888501\_

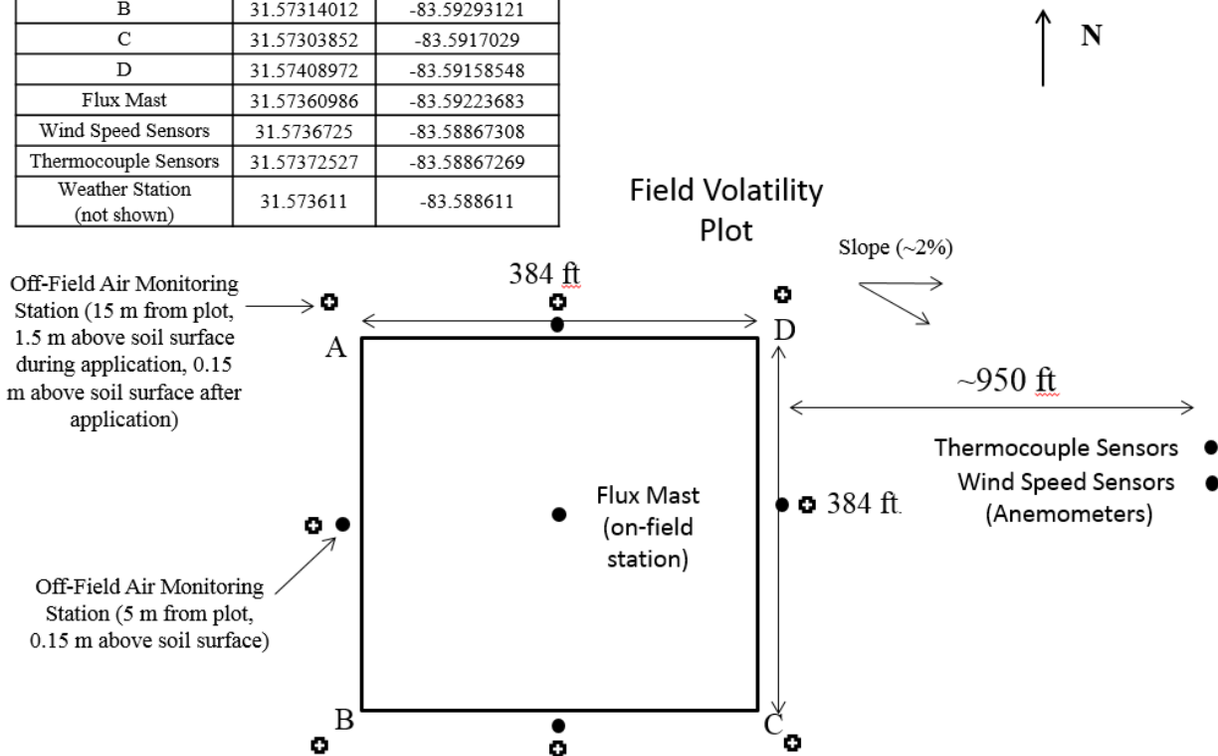
2. Validation spreadsheet for studies following the Integrated Horizontal Flux Method:



field-volatility-835-81  
00-MRID\_49888501\_

### Attachment 3: Field Volatility Study Design and Plot Maps

Point	Latitude (°N)	Longitude (°W)
A	31.5741914	-83.59281253
B	31.57314012	-83.59293121
C	31.57303852	-83.5917029
D	31.57408972	-83.59158548
Flux Mast	31.57360986	-83.59223683
Wind Speed Sensors	31.5736725	-83.58867308
Thermocouple Sensors	31.57372527	-83.58867269
Weather Station (not shown)	31.573611	-83.588611



Note: This study profile focuses on results of flux determination after application using methods employing dicamba concentrations measured from the center flux mast. The purpose of the off-field air monitoring stations is described in the report (MRID 49888501).

## Attachment 4: Calculations and Index of Variables Used in Flux Determination Methods

### Aerodynamic Method

$$\text{Equation } x_1 \quad \text{Flux} = \frac{-(0.42)^2 (c_{z_{top}} - c_{z_{bottom}})(u_{z_{top}} - u_{z_{bottom}})}{\phi_m \phi_p \ln\left(\frac{z_{top}}{z_{bottom}}\right)^2}$$

Flux ( $\mu\text{g}/\text{m}^2\text{s}$ ): volatile flux of pesticide from release source surface

$c_{z_{top}}$  ( $\mu\text{g}/\text{m}^3$ ): concentration at the top sampler adjusted according to the regression of concentration vs. ln (height)

$c_{z_{bottom}}$  ( $\mu\text{g}/\text{m}^3$ ): concentration at the bottom sampler adjusted according to the regression of concentration vs. ln (height)

$u_{z_{top}}$  (m/s): wind speed at the top sampler adjust according to the regression of wind speed vs. ln (height)

$u_{z_{bottom}}$  (m/s): wind speed at the top sampler adjust according to the regression of wind speed vs. ln (height)

$\phi_m$  and  $\phi_p$  (dimensionless): Internal Boundary Layer (IBL) stability correction terms determined according to the following conditions based on the calculation of the Richardson number,  $R_i$ :

$$\text{Equation } x_2 \quad R_i = \frac{(9.8)(z_{top} - z_{bottom})(T_{z_{top}} - T_{z_{bottom}})}{\left[\left(\frac{T_{z_{top}} + T_{z_{bottom}}}{2}\right) + 273.16\right] * (u_{z_{top}} - u_{z_{bottom}})^2}$$

where:

$T_{z_{top}}$ : Temperature at the top sampler adjusted according to the regression of temperature vs. ln (height)

$T_{z_{bottom}}$ : Temperature at the bottom sampler adjusted according to the regression of temperature vs. ln (height)

$R_i$  (dimensionless): Richardson Number

if  $R_i > 0$  (for Stagnant/Stable IBL)

$$\phi_m = (1 + 16R_i)^{0.33} \text{ and } \phi_p = 0.885(1 + 34R_i)^{0.4}$$

if  $R_i < 0$  (for Convective/Unstable IBL)

$$\phi_m = (1 - 16R_i)^{-0.33} \text{ and } \phi_p = 0.885(1 - 22R_i)^{-0.4}$$

### Integrated Horizontal Flux Method

$$\text{Equation } x_3 \quad P = \frac{1}{x} \sum_{z_0}^{z_p} (A * \ln(z) + B) * (C * \ln(z) + D) dz$$

P ( $\mu\text{g}/\text{m}^2\text{s}$ ): volatile flux of pesticide from release source surface

z (m): height above ground level

A ( $\text{s}^{-1}$ ): slope of the wind speed regression line by  $\ln(z)$

B (m/s): intercept of the wind speed regression line by  $\ln(z)$

C ( $\mu\text{g}/\text{m}^4$ ): slope of the concentration regression by  $\ln(z)$

D ( $\mu\text{g}/\text{m}^3$ ): intercept of the concentration regression by  $\ln(z)$

$Z_0$  (m): aerodynamic surface roughness length of release source surface

$Z_p$  (m): volatile plume top height; calculated from the following equation:

$$\text{Equation } x_4 \quad Z_p = \exp\left[\frac{(0.1 - D)}{C}\right]$$