

120 Industrial Way Longview, WA 98632

Date: 5/18/07

To: SWCAA

C/O Wess Safford 11815 NE 99<sup>th</sup> St. Suite 1294 Vancouver, WA 98682

Subject: VOC Emissions Final

Wess,

Enclosed you will find the final VOC emissions from drying of red alder for the Weyerhaeuser Hardwoods, Longview facility. The report is in accord with ADP 05-2597, 2.4 Emission Monitoring and Testing Requirements, section 36. The final report was completed by Michael R. Milota from the department of Wood Science and Engineering of Oregon State University, in Corvallis Oregon.

As per our agreement during the conference call on February 27<sup>th</sup>, 2007, the results of February 7<sup>th</sup>, 2007, would not be utilized to calculate future average emission factors. However, the emissions calculations from the second round of testing would be used for future average emission factors.

Testing was completed by May 15th, 2007 as scheduled.

Respectfully,

Rodney B. Smith

**Environmental Coordinator** 

Bob\_\_ Randy

Paul Wess Clint

**Natalia** 

Jerry\_ David

Laurie Brian

Kathy

Leo

Traci

Mary File

Vannessa John\_ Jackie

# VOC Emissions From the Drying of Red Alder Lumber

Report to

Weyerhaeuser Hardwoods Longview, WA

Report by

Michael R. Milota
Department of Wood Science and Engineering
Oregon State University
Corvallis, OR 97331

May 15, 2007

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# VOC emissions from the drying of red alder lumber

#### I. Results Summary

One charge each of 5/4 random width red alder lumber was dried from green in a small kiln at Oregon State University. The kiln dry- and wet-bulb temperatures based on a schedule provided by Weyerhaeuser Hardwoods. The maximum temperature was 170° (77°C). The air velocity was 500 feet per minute (2.5 m/s). The kiln was indirectly heated with steam. There was no humidification. Regulating the amount of air entering the kiln controlled venting and the humidity.

A JUM VE-7 total hydrocarbon analyzer was used to measure organic emissions following EPA Method 25A. The results are shown in Table 1.

**TABLE 1.** Summary of results.

	Initial MC	Time	VOCB		
	%	hr:min	lb/mbf		
Alder	102.1	126:14	0.17		

<sup>&</sup>lt;sup>A</sup> at time VOC was reported, actual final MC was 5.2% at 145.8 hrs

#### II. Lumber Source and Handling

Enough wood for the two charges was delivered by Weyerhaeuser Hardwoods on 4-19-07. The wood appeared fresh. The wood was wrapped in plastic and stored at -10°C in a freezer. Half the alder was randomly selected for the first charge. It was dried for six days starting on Thursday May 3.

<sup>&</sup>lt;sup>B</sup> as carbon reported for drying to 8% moisture content

#### III. Kiln Description and Operation

A schematic of the kiln is shown in Figure 1. The kiln box is approximately 4' by 4' by 4'. It is indirectly heated by steam. Four dry-bulb thermocouples and two wet-bulb thermocouples are located on the entering-air side of the load. The dry-bulb thermocouples are spaced in a grid. The two wet-bulb thermocouples are under a single sock at the center of the entering-air side of the load.

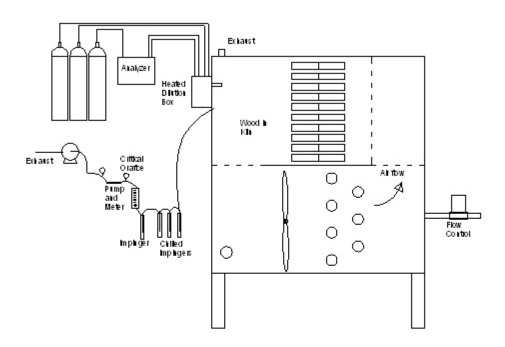


FIGURE 1. Schematic of kiln and sampling system.

#### **Humidity control**

A 200 L/min MKS mass flow meter controlled and measured the amount of air entering the kiln. It was factory calibrated and checked using a bubble meter. The amount of air entering the kiln is based on the wet-bulb temperature - if it is above setpoint, the airflow is increased and if it is below setpoint the airflow is decreased. This is analogous to venting for a commercial kiln. A minimum of 12 L/min entered the kiln at all times, more than removed by the analyzer (1.6 L/min). Putting air into the kiln at a rate of 100 L/min causes the pressure in the kiln to be 60 to 130 Pa above ambient, depending on location in the kiln (high-pressure or low-pressure side). Thus, any fugitive leakage should be out of the kiln. Two additional flow meters can be manually set to provide additional airflow. These were not used in this study. The steam spray line is disabled, so no water vapor is added to the kiln atmosphere. The impinger train in Figure 1 was not used in this work.

#### **Temperature control**

Temperature in the kiln is controlled by indirect steam heating. When the average of the four dry-bulb thermocouples is below setpoint, the steam pressure in the coil is increased. When it is above setpoint, steam flow to the coil is reduced.

#### **Schedules**

The drying schedule used (Figure 2) was based on drying conditions supplied by the mill. The values in Figure 2 are based on the entering-air temperature. This represents the highest temperature the wood would experience in a commercial kiln.

#### **Charge Sequence**

The lumber was thawed, unwrapped, and 2" were trimmed from each end of each board to give 44" samples. These were then weighed, placed in the kiln and dried according to one of the schedules in Figure 2. The actual temperatures are shown in Figure 3. Sampling for hydrocarbon was done as described in section IV. At the end of drying the wood was weighed, oven dried, and reweighed so initial and final moisture contents could be determined by ASTM D4442 (oven-dry method).

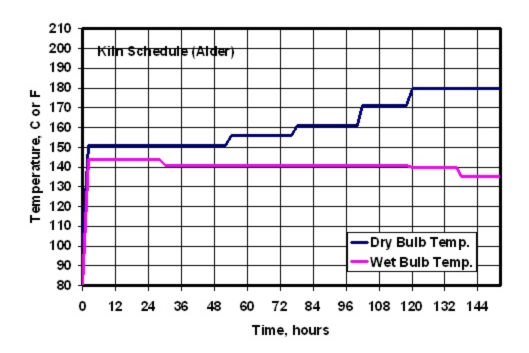


FIGURE 2. Drying schedule.

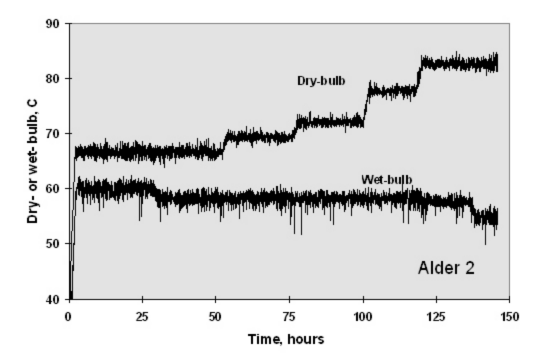


FIGURE 3. Dry- and wet-bulb temperatures.

#### IV. Sampling Systems and Methodologies

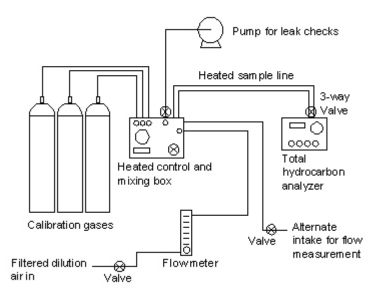
Sampling for total hydrocarbon is done directly from the kiln as shown in Figure 1. The concentration obtained from the hydrocarbon analyzer and the amount of air entering the kiln allow the total hydrocarbon emissions to be calculated. No impingers were used for the work reported here.

Figures 4a and 4b show the hydrocarbon sampling system. Unlike stack testing, all necessary equipment is permanently mounted on the kiln and flows are controlled with valves. The sample is withdrawn from the kiln under the assumption that the gas in the kiln is well-mixed and that the composition in the kiln near the exhaust is the same as the composition of the exhaust. The THC sample was drawn from the kiln directly into a heated dilution/filter box mounted on the side of the kiln. The box was heated to 125°C. Heated dilution gas can be added to the hydrocarbon sample gas to lower the gas moisture content to the detector. Dilution air was used when the gas moisture content in the kiln was greater than 15% so that the air moisture content to the detector remained less than 15%. The sample line from the box to the analyzer was heated to 135°C. The valve at the back of the analyzer was heated to 145°C.

The fuel gas was hydrogen. The span gas was EPA Protocol 905 ppm propane in air, the mid-gas was certified 412 ppm propane. The zero gas was 0.1 ppm air. Detailed sampling procedures are in Appendix 1 and a summary is presented below.

Leak checks were conducted before and after the charge was dried. Valves are closed and all components from just behind the probe tip to the valve at the back of the analyzer are placed under a 18-20 inHg vacuum. Less than one inHg pressure change during two minutes is acceptable and this was met.

Total flow and sample flow to the analyzer were checked using an NIST-traceable flow meter. Total flow is measured with the dilution gas off. Sample flow is measured with it on. This was done at the beginning and end of each sampling interval during which the sample gas was diluted. The meter was attached to the system near the probe tip within the heated box. The valves were repositioned so that the sample came from the flow meter rather than the kiln. Readings of flow were made with the dilution gas both off and on. The flow readings were verified by observing the change in the analyzer reading for span gas with the dilution gas off and on. The dilution ratio calculated based on the analyzer readings was within 4% of that determined by the flow meter.



**FIGURE 4A.** Schematic of heated filter box with air dilution system, heated sample line, and analyzer. Sample enters heated box from back of drawing (box is attached to kiln).



**FIGURE 4B.** Photo of VOC sampling system showing heated sample box (with white insulation), toggle valves and flow meter for calibration gases (upper left), on/off valve for calibration gas (3 at upper center right), heated sample line to analyzer (green tube, lower left), valve for sample (2 at center), toggle valve to vacuum pump (near calibration gas valves), and vent/flowmeter valve (4 at upper right).

Calibration of the zero and span of the detector (JUM VE-7) was done at the beginning of each run (about every four to eight hours). The calibration gas was introduced by setting the valves so the calibration gas entered the system near the probe tip at ambient pressure. The calibration was checked at the end of each run with no adjustments made to the zero or span during the run. The span drift was always less than three percent of full scale for a run and generally less than one percent. The zero drift was minimal during entire drying cycles.

#### V. Data Reduction and Treatment

The "FlowCalc" worksheet in the Excel file "Kiln, ....XLS" in Appendix 2 shows the calculations for each 3-minute interval during the charges. Column A is a reading number. Columns B and C are the clock and charge times, respectively. Columns D and E are the average dry- and wet-bulb temperatures. Column F is the vapor pressure of water at the wet-bulb temperature. The absolute humidity is shown in column G and the molal humidity in column H. These are calculated based on the dry-bulb temperature, wet-bulb temperature, vapor pressure.

#### Flow calculations

The volumetric dry gas flow rate in column I is the flowmeter reading adjusted for the meter calibrations and the molar humidity of the entering gas. This is in standard (at 0°C) liters per minute. In column J this has been converted to a mass flow rate in kg/min and in column K is the same information is expressed as a molal flow rate. These values are for the dry gas vented from the kiln.

#### **Moisture calculations**

The water removal rate in g/min (column L) is calculated from the humidity (column G) and the gas flow (column J). The and the total water (column M) is an integration of column L over time.

The moisture content of the wood at each time interval in the event (column N) was determined by reducing the MC of the wood from the previous value by accounting for the amount of water leaving the kiln during the interval. This amount has been adjusted by adjusting the wet-bulb temperature to make the ending moisture content match.

#### **Total hydrocarbon calculations**

The original total hydrocarbon analyzer reading is shown in column O. In column P this has been corrected to compensate for the range setting switch on the analyzer and scaling between the analyzer reading and the computer reading. Also in column P, the THA data between sampling runs has been adjusted to the average of the data during the 12-minute period before the analyzer testing and calibration time. The dilution THA (column Q) is the corrected THA reading divided by the dilution ratio (from column Y). In column R we have the opportunity to compensate for the effect of moisture on the JUM detector. This was not done so column R equals column Q. Finally in column S, the hydrocarbon concentration is converted to a dry gas basis concentration using the molar humidity (column H).

In column T, the hydrocarbon flow rate in  $g_{carbon}$ /min is calculated in a manner analogous to the water flow rate using the dry gas flow rate and the hydrocarbon concentration. Column U is the integral of column T over time, the cumulative hydrocarbon release up to that point in the schedule. Column V is the cumulative unit emissions, that is, column U divided by the oven-dry weight of the wood in the kiln.

Column X indicates the hydrocarbon sampling run and column Y is the dilution ratio during that run. The next two columns, Z and AA, are the cumulative dry gas and water during the kiln cycle. These are used obtain the average gas moisture contents. The uncorrected wood moisture content is shown in column AC. This is the MC in column N before adjustment of the wet-bulb to make the beginning and ending MCs match the ovendry test. The kiln air and analyzer air moisture contents (based on volume) are shown in columns AD and AE.

At the end of the FlowCalc spreadsheet are summaries by run of the flow data for the total hydrocarbon run intervals. Further down are summaries by impinger interval. These are the tables that appear in the body of the report. The other pages in the files "Kiln, ....XLS" are graphs of the data in the FlowCalc page.

Moisture content and board weight data are in the files named "Weights, ....XLS."

#### VI. Sampling Results

The hydrocarbon emissions are summarized graphically here. All emission data is presented in detail in electronic form in Appendix 2. A summary for each sampling interval is in Table 2.

Figure 5 shows total hydrocarbon concentration (left scale) and dry gas vent rate (right scale) versus time. The vent rate is low for the first few hours as the kiln comes up to temperature and the wet-bulb depression is small. The venting then increases followed by decreases in a sawtooth pattern for the remainder of the schedule. The increases in vent rate at each sawtooth correspond the changes in the drying schedule.

The total hydrocarbon concentration is very dependent on the venting. Early in the schedule the high vent rate results in a low hydrocarbon concentration and vice versa. The spike early in the schedule occur while the kiln is warming and before the vents open. When the venting increases, the total hydrocarbon concentration decreases. Note that total hydrocarbon concentration is not indicative of the amount of hydrocarbon emissions unless one also considers the vent rate. These two factors combined determine the emissions.

Figure 6 shows the cumulative hydrocarbon emissions and the rate of emissions versus time. The cumulative emissions is the emissions up to any point in time in the schedule. The rate of emissions is how much is coming out per unit time. The maximum emission rates occur early in each schedule. The rate of emissions is very low at the end of each schedule.

Figure 7 shows the wood moisture content versus time. The estimated moisture content is obtained from the humidity, vent rate, initial sample weight, final sample weight, and oven-dry sample weight. The initial moisture content was 102.1% on a dry basis by ASTM D4442. The final moisture content was 5.3%. The endpoints for the estimated moisture content lines match the oven-dry method. The adjustment was made by adjusting the wet-bulb temperature.

Figure 8 shows the cumulative hydrocarbon emissions versus moisture content. The hydrocarbon emissions for drying to any moisture content can be read from this graph. In agreement with past studies, there is a fairly linear relationship between the emissions and the decrease in moisture content at lower moisture contents.

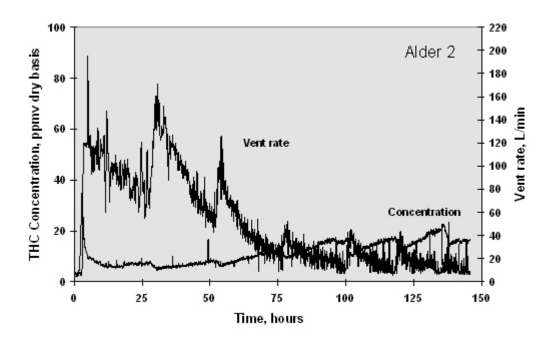


FIGURE 5. Hydrocarbon concentration and vent rate versus time.

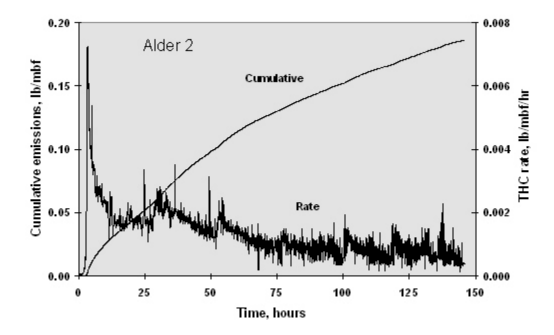


FIGURE 6. Cumulative and rate of emissions versus time (as carbon).

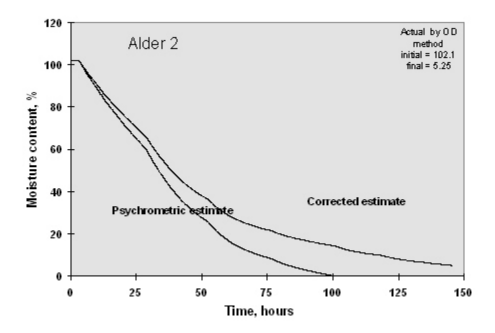


FIGURE 7. Moisture content versus time.

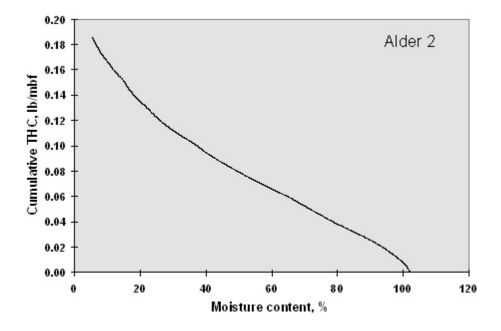


FIGURE 8. Cumulative emissions (as carbon) versus moisture content.

Table 2 shows the VOC results by run for the charge. A run is an interval between analyzer calibrations, about six hours of data. The interval time periods shown in the table include the times between sampling and mass calculations are adjusted to account for these. Sampling occurred for approximately 98% of the drying time. Samples of field sampling sheets, including dilution system and heated component data are given in Appendix 3 with full PDF format versions in Appendix 2.

**TABLE 2.** Summary of sample runs for analysis of total hydrocarbon..

Sample	Time	Cumu	lative	Dry Flow	THC	THC	conc	THC mass	THC rate	Av	erage N	1 C
Run		Dry Gas	Water	Rate @68	asC	wet	dry	as C	as C	Wood	Air	A nal.
	hrs	kg	kg	l/min	g	ppmv	ppmv	lbg/mbf	lb/hr/mbf	%	%	%
1	2.26	1.30	0.07	7.9	0.01	4.0	4.5	0.000	0.0001	102.1	7.7	7.7
2	4.48	35.08	4.98	108.2	0.54	12.2	15.1	0.017	0.0037	99.2	18.6	18.6
3	2.54	21.71	3.05	118.4	0.21	6.4	7.9	0.007	0.0026	93.5	18.5	11.7
4	3.17	27.87	3.96	121.5	0.22	5.2	6.5	0.007	0.0022	88.9	18.6	11.9
5	2.85	20.96	2.96	101.5	0.16	4.9	6.1	0.005	0.0017	84.4	18.5	11.8
6	2.72	19.36	2.73	98.5	0.15	4.9	6.1	0.004	0.0016	80.7	18.5	11.8
7	3.31	22.67	3.23	94.8	0.20	5.6	7.0	0.006	0.0018	76.7	18.7	11.9
8	3.26	20.64	2.97	87.5	0.18	5.8	7.2	0.006	0.0017	72.7	18.8	12.0
9	5.16	37.44	5.24	100.3	0.32	5.6	7.0	0.010	0.0019	67.6	18.4	11.7
10	2.58	27.99	3.54	149.9	0.18	4.3	5.3	0.006	0.0022	61.6	16.9	10.7
11	3.76	36.26	4.61	133.4	0.26	4.7	5.7	0.008	0.0021	56.2	17.0	10.9
12	3.49	29.36	3.74	116.4	0.22	5.0	6.1	0.007	0.0020	50.8	17.0	10.9
13	8.60	53.43	6.81	85.8	0.45	5.7	6.9	0.014	0.0016	43.7	17.0	10.9
14	3.76	16.87	2.13	62.1	0.18	6.9	8.4	0.005	0.0014	38.1	16.9	10.8
15	7.92	46.56	5.88	81.2	0.42	6.0	7.3	0.013	0.0016	32.7	16.9	10.8
16	3.26	12.74	1.62	54.0	0.14	7.3	8.9	0.004	0.0013	28.0	17.0	10.9
17	9.19	24.26	3.07	36.5	0.31	8.5	10.3	0.009	0.0010	24.8	17.0	10.9
18	2.99	6.04	0.76	27.9	0.09	9.7	11.8	0.003	0.0009	22.4	16.9	10.9
19	8.56	20.83	2.59	33.6	0.27	8.8	10.7	0.008	0.0010	20.2	16.7	10.7
20	3.76	6.77	0.85	24.9	0.11	10.4	12.6	0.003	0.0009	18.0	16.8	10.9
21	8.01	12.00	1.49	20.7	0.22	12.4	15.1	0.007	0.0009	16.5	16.7	10.8
22	2.94	3.75	0.47	17.6	0.07	13.0	15.8	0.002	0.0008	15.2	16.9	10.9
23	8.92	17.68	2.17	27.4	0.27	10.6	12.8	0.008	0.0009	13.5	16.5	10.7
24	4.08	5.84	0.72	19.8	0.10	11.6	14.1	0.003	0.0008	11.6	16.5	10.8
25	8.24	9.73	1.20	16.3	0.18	12.7	15.4	0.006	0.0007	10.4	16.5	10.7
26	3.40	6.42	0.74	26.1	0.10	10.4	12.5	0.003	0.0009	9.0	15.7	10.2
27	2.99	3.95	0.46	18.3	0.07	12.1	14.5	0.002	0.0007	8.3	15.8	10.3
Sum	126.2	547.5	72.1		5.6			0.173				
Average				66.3		8.0	9.7		0.0014			

#### VII. Quality Assurance

#### Leak checks

Leak checks were performed on the VOC system before and after drying each species by pulling a 17 inHG vacuum and sealing the system for 2 minutes with no change in pressure.

#### Calibration

Data for the calibration gases are given in Appendix 4 The calibration sheet for the flow meter is also included is also included in Appendix 4 as is the thermocouple calibration check.

#### **Anomalies**

There were no anomalies during the work that would significantly affect the data.

	Appendix 1. Detai	led Sampling	Procedures
OSU, Wood Scien	nce and Engineering	14 Wey	erhaeuser Hardwoods, May, 2007

#### NSTRUCTIONS FOR CHECKS OF EMISSIONS KILN

**Purpose:** Ensure kiln is operating correctly

**Clock time:** Record from computer

Run time: Record from computer. Check the box if the computer screen being refreshed

and time is advancing.

**Box temperature:** Read from metal electrical box under desk, left controller. The top and

bottom numbers should be similar on the box should be similar, about 126 C...

**Valve temperature:** Read from metal electrical box under desk, right controller. The top and bottom numbers should be similar on the box should be similar, about 154 C..

**Dry-bulb temperature:** Read from computer screen. Compare to graph to be sure it's correct. If it's not within a degree or two of the chart, check again in a few minutes. During startup (the first 3 or so hours), it may not be able to track. If it's too high, the heat valve should be closed, too low and the heat valve should be open. If it does not appear to be working correctly, call Mike or Mark.

**Wet-bulb temperature:** Read from computer screen. Compare to graph to be sure it's correct.

If it is too low, it means that the kiln atmosphere is too dry. Check the flow meters. If Flow1 is about 10 L/min (its lower limit), make sure that Flow2 and Flow3 are turned off

If it's too high, then either the kiln atmosphere is too humid or the sock is not being wetted. If Flow 1 is near 200 L/min (its upper limit) add venting by opening Flow2 and/or Flow 3. The maximum for Flow2 is 50 L/min, if it reads over this value for several readings, reduce it to about 45 L/min. Don't change Flow3 often, rather set it and leave it for several hours if possible. Keep the Flow 3 reading constant by small adjustments. As Flow1 decreases or Flow2 turned down, there is more pressure behind Flow3 and the flow increased. Check for water in the wet-bulb reservoir (push the float down and make sure it's getting water).

Check both Wet-bulb1 and Wet-bulb2 and make sure they are reading about the same. If they differ by more than 2 C, call Mike or Mark.

If both wet-bulbs are reading the same as the dry-bulb, check the wet-bulb water.

If these procedures do not correct the wet-bulb temperature within 30 minutes, call Mike or Mark.

**Line temperature:** Read from gray box on wall above analyzer. It should read about 275°F.

**Chiller temperature:** Read the chiller temperature. It should be about -1°C.

**Flow 1:** Read from computer. The value of Flow1 changes depending on the wet-bulb. If Flow 1 is 10 L/min and the wet-bulb is too low, there's probably nothing we can do. If it's 200 L/min and the wet-bulb is too high, Flow2 and/or Flow3 can be opened. Flow2 and Flow3 should be adjusted so that Flow1 stays below 175 to 200 L/min.

**Flow 2:** Read from computer. The value of Flow2 is set by you. It will vary a little - as flow 1 goes down, flow 2 will go up. Do not set it to < 40 L/min if you think Flow1 is going to decrease or it will go off scale and not be read by the computer

**Flow 3:** Read from meter. The value of Flow3 is set by you. It will vary a little - as flow 1 goes down, flow 2 will go up. Be sure to clearly record this value and when you change it

**Dilution flow:** Read dilution flow meter. It should read the same setting as the red flag. Do not adjust. If significantly different, investigate.

**F/M Flow:** Read from rotometer. This should be about 400 to 500 cc/min.

**Line vacuum:** Read from the vacuum gauge. This should be about 20"Hg.

# INSTRUCTIONS - FIELD DATA SHEET FOR TOTAL HYDROCARBON ANALYZER PRE-SAMPLE PROCEDURE

#### BACKGROUND INFORMATION

Get the dry- and wet-bulb temperatures from the kiln schedule or off the computer. Use the highest expected values for the run.

Read absolute humidity off the psychrometric chart or table.

Calculate or read from tables -

Percent moisture = 100 / [ 1 + 1 / 1.61\*AbHum ]

Target Dilution Ratio (TDR) = 15 / Percent Moisture

Event = the name of the drying cycle.

Run = the number of the 3-hour interval.

Operator, that's you.

Date and time are now, as you start the data collection process.

#### **AMBIENT DATA**

Call 9-754-0081 and get altimeter setting.

Read the laboratory temperature from the thermometer.

#### **ANALYZER CALIBRATION**

Set valves so that 1, 2 = off; 3=on; 4=vent. This allows gas to flow out of the vents from the calibration tanks and shuts off all other sources. Only calibration gas should go through the detector.

Open the zero gas tank valve

zero toggle switch up (on), others down (off) set flow to 3.5 L/min using regulator on tank wait for a stable reading (about 30 to 60 seconds) use the zero dial (pot) on THA to get a zero reading read the analyzer read computer note pot setting

#### Open span gas tank valve

span toggle switch up (on), others down (off) set flow to 3.5 L/min using regulator on tank set analyzer to range 3 wait for a stable reading (about 30 to 60 seconds) use the span dial (pot) on THA to get a reading of 905 ppm

close valve on zero gas tank

read the analyzer, record, for example, 9.05 or 900 read computer (should read about 905) note pot setting

#### Open mid gas tank valve

mid toggle switch up (on), others down (off) set flow to 3.5 L/min using regulator on tank wait for a stable reading (about 30 to 60 seconds) read analyzer (do not adjust pot settings), record, for example, 4.12 or 412 read computer (should about 412) check for within tolerance turn off mid gas all toggle switches off

#### SET DILUTION FLOW BEFORE RUN

Set valves so that 1, 2, 3 = off; 4=meter. This allows gas to flow only from the meter to the detector.

Use the Gilibrator to take 4 readings of the total flow rate (TFR). This is the total flow drawn by the analyzer and should be about 2.6 L/min

Make sure the average does not include any "bad" readings Record the average, L/min = cc/min / 1000

Write the Event, Run, and "Pre-TFR" on the Gilibrator printout.

#### Calculate the next two values -

Target dilution flow rate (TDFR) is the TFR x (1 - DR) Target sample flow rate (TSFR) is the TFR x DR Check that the sum of these is the Total Flow Rate

#### Set dilution flow

Set red pointer to desired dilution flow (on meter with valve 1) Slowly open lower valve on dilution flow meter (1=on; 2, 3=off; 4=meter) Use upper valve on dilution flow meter to adjust flow Do not adjust this meter after this point Read the meter that you just set and record the value

Use the Gilibrator to take 4 readings of the sample flow rate (SFR). This is the flow through the analyzer after dilution is set. It will vary, depending on the dilution setting.

Make sure the average does not include any "bad" readings Record the average, L/min = cc/min / 1000

Write "Pre-SFR" on the Gilibrator printout.

#### **CHECK DILUTION FLOW BEFORE RUN**

Set valves so that 1, 3 = on; 2=off; 4=vent. This allows gas to flow out of the vent from the calibration tank and shuts off all other sources. Calibration gas and dilution air will go through the detector.

Open span gas tank valve

span toggle switch up (on), others down (off)
set flow to 3.5 L/min using regulator on tank
set analyzer to range 3
wait for a stable reading (about 30 to 60 seconds)
record
turn off all calibration gas tank valves
all toggle switches off

Calculate the dilution ratio based on gas flow by dividing the Sample Flow Rate by the Total Flow Rate.

Calculate the dilution ratio based on span gas by dividing the Diluted span by the undiluted span.

If the Dilution ratios do not agree within 5% - DO NOT PROCEED\*\*\*\*. Use  $100*(DR_{Span} - DR_{Flow})/DR_{Flow}$  to calculate the % difference.

\*\*\*\* check calculations, check that values for ppm and flows make sense, remeasure everything. If it still does not agree, call Mike or Mark

#### **START RUN**

Set valve so that 1, 2, 5 = on; 3, 4=off; all calibration tank valves off

Record the start time. Use the computer clock for all times or set your watch to the computer time.

Make sure analyzer is on appropriate range, usually range 3, to keep THC reading on computer between 60 and 750.

Monitor system, as needed. Record system condition at least hourly.

End time should be no more than 3 hours from start time.

#### **POST-SAMPLE PROCEDURE**

#### AT END OF RUN

Record your name as the operator.

Event = the drying cycle. Run = the 3-hour interval.

Operator, that's you. Date and time are now, as you start the data collection process.

#### AMBIENT DATA

Call 9-754-0081 and get temperature and altimeter

Local pressure = (Altimeter - 0.23) x 3.3867

Read the laboratory temperature from the thermometer.

Fill out appropriate information on Pre-sample side of data sheet for next run. This will save time in between runs.

#### **END TIME**

Record computer time.

DO NOT adjust dilution gas yet.

#### **CHECK DILUTION FLOW AFTER RUN**

Set valves so that 1, 3 = on; 2=off; 4=vent. This allows gas to flow out of the vent from the calibration tank and shuts off all other sources. Calibration gas and dilution air will go through the detector.

Open span gas tank valve

span toggle switch up (on), others down (off) set flow to 3.5 L/min using regulator on tank wait for a stable reading (about 30 -60 seconds) record all toggle switches off

**Sample flow rate.** Set valves so that 1=on; 2, 3 = off; 4=meter. This allows gas to flow only from the meter and the dilution to the detector.

Use the Gilibrator to take 5 readings of the sample flow rate (SFR). This is the flow through the analyzer with dilution on.

Make sure the average does not include any "bad" readings

Record the average, L/min = cc/min / 1000

Write "Post-SFR" on the Gilibrator printout.

Read dilution flow meter
To calculate the L/min, divide scfh by 2.12
Turn off dilution flow meter using valve 1

**Total flow rate**. Set valves so that 1, 2, 3 = off; 4=meter. This allows gas to flow only from the meter to the detector.

Use the Gilibrator to take 5 readings of the total flow rate (TFR). This is the total flow drawn by the analyzer and should be about 2.6 L/min Make sure the average does not include any "bad" readings Record the average, L/min = cc/min / 1000 Write "Post-TFR" on the Gilibrator printout.

#### CHECK CALIBRATION OF ANALYZER

Set valves so that 1, 2 = off; 3=on; 4=vent. This allows gas to flow out of the vents from the calibration tanks and shuts off all other sources. Only calibration gas should go through the detector.

Span gas tank valve should be open

span toggle switch up (on), others down (off)

set flow to 3.5 L/min using regulator on tank

set analyzer to range 4

wait for a stable reading (about 30 -60 seconds)

read analyzer (do not adjust pot settings), record, for example, 1.50 as 1500

read computer (should read about 152 due to range 4 setting)

note pot setting

check for within tolerance - between 1483 and 1573

#### Open mid gas tank valve

mid toggle switch up (on), others down (off)

set flow to 3.5 L/min using regulator on tank

set analyzer to range 3

wait for a stable reading (about 30 -60 seconds)

read analyzer (do not adjust pot settings), record, for example, 8.50 as 850

read computer (should read same as analyzer)

check for within tolerance

#### Open the zero gas tank valve

zero toggle switch up (on), others down (off)

set flow to 3.5 L/min using regulator on tank

wait for a stable reading (about 30 -60 seconds)

read analyzer (do not adjust pot settings)

read computer note pot setting

Calculate the dilution ratio based on gas flow by dividing the Sample Flow Rate by the Total Flow Rate.

Calculate the dilution ratio based on gas flow by dividing the Sample Flow Rate by the Total Flow Rate.

Calculate % difference as 100 \* {Absolute Value (DR<sub>Span</sub>-DR<sub>Flow</sub>)} / DR<sub>Flow</sub>

Record the time now as the end time for check.

Tear off the four sets of Gilibrator readings (Pre-TFR, Pre-SFR, Post-SFR, Post-TFR) and staple to paper with other records.

Start Pre-Sample procedure for next run.

## Appendix 2. Data in Electronic Form

		Appendix 3. Samp	oles of Field Dat	a <b>Sheet</b> s
OSU, Wood Science and Engineering 24 Weyerhaeuser Hardwoods, May, 2007	OSU, Wood Scie	nce and Engineering	24 Weyerha	euser Hardwoods, May, 2007

Weylo Alder 2

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			, , , , , , , , , , , , , , , , , , ,	Line 3	Vac.	inHg			22	22				//					1		\		
				rine z	Vac.	inHg			22	22			42	75	24]	/					\		
			7	rine 1 Line 2	Vac.	inHg	25	25	22	22	74	, 72				72	42	hz	24		3		28.5
					Line 3	ml/min	/	/	500	264													
					Line 2	ml/min			420	460			954	450	450			/	1				
					Line 1	ml/min	624	064	263	490	06h	490			/	490	Cb)~	9	06	1	190		र्पुरा
				FIOWS	Dilution	SCFM			(3	1.3	1.3	1.3	(,3	1.3	(.3	1-3	(, 3	(.)	<b>(1)</b>	(1,3	(,3		1,3
					Flow 3	SCFH																	
					Flow 2	L/min																	
					Flow 1	L/min		හ	561	દાા	911	951	SL	<u>-5</u> b	20	<b>}</b> 8	(ه ۶	(ه٤	18	041	771		(1)
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			İ		Line	ပံ			38)	581		-	135	135	135	581	581	52)	5		135		135
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te	20-2-		1	empe	Dry- bulb	ာ့	25	530	5		<b>189</b> 7	4. صاح)	Ge. B	674 61.8	670 617	60.1	1.00	Je. if	61.3	1.73	ا مام		Je. 2 60 4
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				ב ב		#	-	۵	4	3	4	77	¿S	$\sim$	S	9	6	(	8	0)	4	#	12
				un Y	time	hrs	0.01	1:00	4:59	6:36	4.59	11:49	15:4	13:36	14:44	15:35	16:42	7):4	21.33	95,56	36.02		37:08
Charge:		Page:	-	200 200 200 200 200 200 200 200 200 200	time		8.09	4:14		40:51	17:37	14:24	12:57	M:17	28:77	25:43	05:00	05:1	5:41	14,03	01:02	21-17	_
$\overline{}$		<u> </u>						Щ.														<u> </u>	6

### FIELD DATA SHEET FOR TOTAL HYDROCARBON ANALYZER - BEFORE

BACKGR	OUND INFORMA	TION	·				
Event (kilr	n charge): <u>Weyco</u>	Alder2	Time now: 17:32				
Run (sam	ple):		Dry-bulb temperature:				
Operator:	$\sim$ $\sim$ $\sim$ $\sim$ $\sim$	rev	Wet-bulb temp	eratu	re:		
Date:	5-2-07		Target Dilution		\$		
AMBIENT	DATA						
Laborator	y temperature:_2	1.6°C					
ANALYZE	R CALIBRATION	L		[1, 2	= off; 3=on; 4=vent]		
	Analyzer, ppm	Analyzer, ppm Computer		9	Pot settings		
zero	O (0)	0	does not apply		468		
span	611 (611)	611	does not app	ly	489		
mid	305 (300)	305	280 to 322		none		
SET DILU	TION FLOW BEF	1 - 1/	· · · · · · · · · · · · · · · · · · ·				
Total flow	rate (TFR):	1.716	L/min	[1,	2, 3 = off; 4=meter]		
Target dilu	ition flow rate (TD	FR)	_ L/min		[TFR x (1 - DR)]		
san	nple flow rate (TS	FR)	_ L/min				
Set and re	ead dilution meter:	_1.3	scfh	[	scfh = L/min * 2.12 ]		
Sample flo	ow rate (SFR):	1.093			2, 3 = off; 4=meter]		
CHECK D	ILUTION FLOW	BEFORE RUN	T	[1,	3=on; 2=off; 4=vent ]		
	Analyzer	DR <sub>Span</sub> [ Span <sub>Diluted</sub> /Span ]	DR <sub>Flow</sub> [SFR/TFR]	100*	Difference, % (DR <sub>Span</sub> - DR <sub>Flow</sub> )/DR <sub>Flow</sub>		
Span <sub>Dilt</sub>	uted 389	0.6367	0.6369		0.03		
START TI	ME: 17:35	T	1, 2, 5 = on: 3.	4 = 0	off: tank valves off 1		

[ 60 < computer reading < 750 ]

ANALYZER RANGE: \_\_\_\_\_

Operator: _	). Kebu		E	vent (kiln ch	narge): <u>Weyco Alder2</u>	
Time now: _	20:35	-	F	Run (sample	): <u> </u>	
AMBIENT D	DATA	***************************************			, , , , , , , , , , , , , , , , , , ,	
Laboratory t	temperature: ¯	<u>S·I</u> °C				
END TIME:	70:35	<del>-</del> .				
CHECK DIL	UTION FLOW	AFTER RUN		[1,	3=on; 2=off; 4=vent]	
	. :	Analyzer			Computer	
Sr	oan <sub>Diluted</sub>	383		3A3		
Sample flow	rate (SFR) :	1.102	_ L/min	[1= on, 2, 3 = off, 4=mete		
Read dilutio	n meter:	_ scfh	_ L/min		[ L/min = scfh*0.472 ]	
Total flow ra (attach print	ate (TFR): out with all fou	r sets of data)	_ L/min	[-1	, 2, 3 = off; 4=meter]	
Dilution ratio		0.6385	5		[SFR/TFR]	
CHECK OF	ANALYZER C	ALIBRATION		ſ 1.	2=off; 3=on, 4=vent ]	
- VIII - VIII	Analyzer	Computer	With	in range	Pot settings	
span	601	601	593	to 629	489	
mid	299	300	287	to 313	none	
zero	8	9	-18	to +18	468	
Dilution ratio	-,	0-637			[ Span <sub>Diluted</sub> / Span ]	
Dilution ratio	o difference:	0.1879	.% [10	0*(Abs(DR	$_{Span}$ - $DR_{Flow}))/DR_{Flow}$	
End time for	r check:	20:41	-			
Comments:		·				

Appendix 4. Calibration Data

# Airgas

# **CERTIFICATE OF ANALYSIS Grade of Product: EPA Protocol**

Airgas Speciality Gases

12722 S. Wentworth Avenue

Chicago, IL 60628 1-773-785-3000

Part Number:

E02AI99E15A0453

Reference Number: 54-124086894-1 FAX: 1-773-785-1928

http://www.airgas.com

Cylinder Number:

XC031356B

Cylinder Volume:

146 Cu.Ft.

Laboratory:

ASG - Chicago - IL

Cylinder Pressure:

2015 PSIG

Valve Outlet:

Analysis Date:

Feb 09, 2007

590

Expiration Date: Feb 09, 2010

Certification performed in accordance with "EPA Traceability Protocol (Sept. 1997)" using the assay procedures listed. Analytical Methodology does not require correction for analytical interferences. This cylinder has a total analytical uncertainty as stated below with a confidence level of 95%. There are no significant impurities which affect the use of this calibration mixture. All concentrations are on a volume/volume basis unless otherwise noted.

Do Not Use This Cylinder below 150 psig.i.e. 1 Mega Pascal

ANALYTICAL RESULTS									
Component	Requested Concentration		Protocol – Total Relative Method Uncertainity						
PROPANE Air	300.000 PPM Balance	299.9 PPM G	-1 +/- 1% NIST Traceable						

CALIBRATION STANDARDS									
Type Lot ID Cylinder No Concentration Expiration Date									
NTRM	51919	SG9101963ALB	483.6PPM PROPANE/	Jul 01, 2009					
			ANALYTICAL EQUIPMENT	C					
Instrument/Make/Model			Analytical Principle	Last Multipoint Calibration					
VARIAN (	CP3800		FID	Feb 02, 2007					

Triad Data Available Upon Request

Notes:

QA Approval



## **Certificate of Analysis: EPA Protocol Gas Mixture**

Airgas Specialty Gases

12722 S. Wentworth Avenue

Chicago, IL 60628

1-733-785-3000 Fax: 1-733-785-1928

Cylinder Number: Cylinder Pressure: Certification Date:

CC44350

Reference Number: 54-124076439-1

10/4/2009

Expiration Date: 2000.6 PSIG 10/4/2006 Laboratory:

ASG - Chicago - IL

#### **Certified Concentrations**

Component	Concentration	Accuracy	Analytical Principle	Procedure
PROPANE	611.3 PPM	+/- 1%	FID	G1
Air	Balance			

Certification performed in accordance with "EPA Traceability Protocol (Sept. 4997)" using the assay procedures listed. Analytical Methodology does not require correction for analytical interferences.

Notes:

Do not use cylinder below 150 psig.

Approval Signature

Reference Standard Information

<u>Type</u>

**Balance Gas** 

R 310893

S 392797

Component Cyl.Number

Concentration

NTRM 51919

PROPANE

SG9101963ALB

483-6 PPM

**Analytical Results** 

1st Component

**PROPANE** 

1st Analysis Date:

10/04/2006

S 391575 Z 0

Conc 609.7 PPM

R 310807 \$ 393458 Z 0

Z 0 R 316077 Conc 612.6 PPM Conc 611.6 PPM

AVG: 611.3 PPM