Robert J Giraud <Robert.J.Giraud@US A.dupont.com> 03/01/04 12:55 PM .

0900-1700-502-1990

To: Rich Leukroth/DC/USEPA/US@EPA cc: John Blouin/DC/USEPA/US@EPA, Greg Fritz/DC/USEPA/US@EPA, david.menotti@shawpittman.com, Stephen H Korzeniowski <Stephen.H.Korzeniowski@USA.dupont.com>, Robert C Buck <Robert.C.Buck@USA.dupont.com>, bill.beers@omnova.com Subject: revisions to final draft ECA appendices based on today's drafting committeecall

Colleagues,

Thank you for your comments on the appendices during this morning's call.

Here is revised Telomers Appendix A.4. Per the discussion (to prevent confusion with the new text for Part IX of the ECA cover document, the word "assembling" is changed to "receiving" on line 4 of page A.4-2. Similarly, I have revised the title of Section 4.1 (of Appendix A.4 for telomers) from "Assembly of Components" to "Handling of Components" for the same reason.

(See attached file: App A.4 telomers incin test draft 2-27-04-revised.pdf)

Here is revised Fluoropolymers Appendices A.2 through A.4. I have corrected errant mention of Composite 21 to Composite 1 in line 4 of page A.4-2 of fluoropolymers appendix A.4

(See attached file: App A.2 through A.4 Fluoropolymers draft 2-27-04-revised.pdf)

Since I had forgotten to Here is revised compiled Appendices B through G. introduce Appendix E.3 in the opening of Appendix C.2 and based on this morning's discussion, please see new section C.2.5.6 introducing Appendix Also, please see revised Appendix E.3 based on this morning's E.3. discussion. Please note that the parentheticals for "provided Phase II is performed" is to clarify that there will be no report for Phase II incineration testing if Phase II is not carried out (e.g., need for Technical Consultation and no agreement to proceed with Phase II testing) and to clarify that the test substance composites are germane only to Phase I have also corrected an oversight on Friday and II and not to Phase I. have bolded the docket number in the title section of the second page (Fluoropolymers) of Appendix G. Furthermore, I have added a cross-reference to Appendix C.2.5 in the last paragraph of Appendix C.1.3 (line 8 of page C.1- 4) with regard to reporting results for improved clarification.

(See attached file: Appendices B through G draft 2-27-04-revised.pdf)

If you have any questions or comments, please let me know.

I look forward to our 9 am to 10:30 am call on Tuesday March 2.

Best Regards,

Robert Giraud

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App A.4 telomers incin test draft 2-27-04-revise: App A.2 through A.4 Fluoropolymers draft 2-27-04-revise



Appendices B through G draft 2-27-04-revised

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## 1 APPENDIX A.2

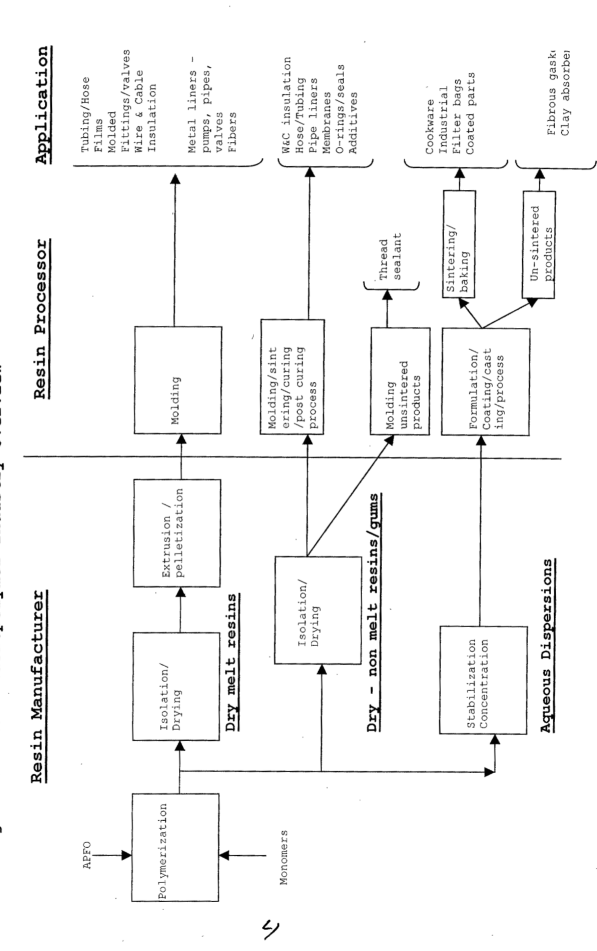
2 RATIONALE FOR SELECTING COMPOSITES TO BE TESTED 3 4 Review of Figure A.2-1 demonstrates that fluoropolymers industry products can be divided into 3 broad categories as 5 6 follows: 7 8 • Dry melt resins 9 • Dry non-melt resins and gums 10 • Aqueous dispersions 11 12 These three broad categories can in turn be divided into 13 four representative classes as follows: 14 15 • Dry melt resins 16 1. FEP, PFA, THV, ETFE, HTE 17 • Dry non-melt resins and gums 18 19 2. Dry non-melt resins 20 3. Fluoroelastomers (dry non-melt gums) 21 22 • Aqueous dispersions 23 4. PTFE, FEP, PFA, THV 24 25 Composite samples of each of these four representative 26 classes were selected as the test substance for this 27 testing program in order to represent the entire range of fluoropolymers involved. 28

A.2-1

INTERESTED PARTY REVIEW

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A.2-2

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1 APPENDIX A.3

## 2 COMPOSITION OF COMPOSITES TO BE TESTED

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6 7 The four composite test substances for this test program are presented below in Table A.3-1 with the fluoropolymer types, CAS numbers, and associated monomers for these fluoropolymers. Each fluoropolymer used in each relevant test substance composite will have been made using APFO.

8 9 10

Table	A 2_1	Test	Substance	Composites	her memo	
Table	A. 3-1.	. Test	Substance	COMPOSITES	DV TVDE	<u>×</u>

Test Substance	Fluoropolymer Type	CAS Number	Associated Monomers
Composite 1 -	PTFE	9002-84-0	TFE
Dry non-melt resin	Modified PTFE	26655-00-5	TFE, PPVE
Composite 2 -	FEP	25067-11-2	TFE, HFP
Dry melt resins	PFA	26655-00-5	TFE, PPVE
		31784-04-0	TFE, PEVE
	THV	25190-89-0	TFE, HFP, VDF
	ETFE	68258-85-5	TFE, E
	HTE	35560-16-8	TFE, HFP, E
Composite 3 - Fluoroelastomers	Fluoroelastomer Copolymers	9011-17-0	VDF, HFP
	Fluoroelastomer Terpolymers	25190-89-0	TFE, HFP, VDF
	Base resistant elastomers	54675-89-7, 27029-05-6	TFE, VDF, P TFE, P
	Perfluoroelastomers	26425-79-6	TFE, PMVE
	CTFE elastomers	9010-75-7	CTFE, VFD
	Low temperature elastomers	CBI	TFE, VDF
Composite 4 - Aqueous	PTFE	9002-84-0	TFE
Dispersions	FEP	25067-11-2	TFE, HFP
_	PFA	26655-00-5	TFE, PPVE
	THV	31784-04-0 25190-89-0	TFE, PEVE TFE, HFP, VDF

11

12 Confidential business information (CBI) regarding the

13 chemical identity of Low temperature elastomers has been

14 submitted to EPA under separate cover.

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1 APPENDIX A.4 2 PREPARATION OF COMPOSITES TO BE TESTED 3 4 4.1 Approach 5 6 A composite mixture of representative fluoropolymers, as 7 solids, will be prepared for each of the four test 8 substance composites. 9 10 The polymer samples will be first quality product polymer, 11 substantially free of inorganic constituents. Each sample will be from a representative grade for each applicable 12 13 fluoropolymer type from each applicable company. 14 15 A hypothetical example for Composite Z in Table A.4-1 below shows how the composites will be assembled. 16 In this 17 example with 4 types across 4 companies, there are 11 x's. 18 Hence, composite Z would be made up of 11 equal proportions 19 of the materials indicated with an x.

20 21

Table A.4-1. Example for Compositing Across Companies & Types

Test Substance	Fluoropolymer Type	Company A	Company B	Company C	Company D
Composite Z	Type 1		х	х	х
	Туре 2	X	x	x	х
	Туре З			x	
	Туре 4	X	X	х	

22

## 23 4.2 Preparation

24

25 Representative samples of each component from each 26 applicable company for each composite will be sent to the 27 compositing facility(ies) in packaging customarily used for 28 product sample packaging or in polyethylene, polypropylene, 29 or glass containers.

30

31 Each composite will be prepared under conditions designed 32 to prevent cross-contamination and designed to assure 33 solids temperatures less than or equal to 60 °C.

34

Following preparation of each composite, the composite will be placed in a polyethylene, polypropylene, or glass container.

38 4.2.1 Composite 1

39

40 Dry non-melt resins are available in powder form. Equal

A.4-1

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weights of the powder form of each of the two types of 1 2 components (following the approach in the example for 3 Composite Z in Section 4.1 above) will be mixed together in 4 dry form to yield Composite 1. 5 6 4.2.2 Composite 2 7 8 FEP, PFA, THV, ETFE, and HTE dry melt resins are available in powder form. Equal weights of the powder form of each 9 component (following the approach in the example for 10 11 Composite Z in Section 4.1 above) will be mixed together in 12 dry form to yield Composite 2. 13 14 4.2.3 Composite 3 15 Fluoroelastomers are available in slab, lump, or sheet 16 17 form. Composite 3 will be prepared following one of the 18 following approaches: 19 20 a) Equal weights of each component (following the approach 21 in example for Composite Z in Section 4.1) will be mixed 22 on a rubber mill to produce a homogenous slab of preset 23 thickness to yield Composite 3. 24 25 Or 26 27 b) Each component of Composite 3 will be cyrogenically 28 cooled (to make the elastomers brittle) and size-reduced 29 (e.g., ground) to produce powder. Equal weights of the 30 powder form of each component (following the approach in 31 the example for Composite Z in Section 4.1) will be mixed 32 together in dry form to yield Composite 3. 33 34 4.2.4 Composite 4 35 36 Aqueous dispersions of PTFE, FEP, PFA, and THV are 37 available as dispersions containing 20 to 60% fluoropolymer 38 solids by weight. Composite 4 will be prepared following one of the following approaches: 39 40 41 a) Equal weights (on a dry solids basis) of each component in aqueous dispersion form (following the approach in 42 43 example for Composite Z in Section 4.1) will be mixed 44 together in liquid form. Solids will be separated from the resulting liquid composite to yield low water content 45 (i.e., drip free) fine solids. 46

A.4-2

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1 2 Or 3 4 b) Solids will be separated from liquid for each component 5 of Composite 4 to yield low water content (i.e., drip free) fine solids for each component. Equal weights of 6 7 the solids form of each component (following the approach in the example for Composite Z in Section 4.1) will be 8 mixed together to yield Composite 4. 9 10 11 4.3 Verification 12 13 In order to assure that composite samples in this testing program have been made up of clearly identified materials, 14 the preparation of the composites will include formal Chain 15 16 of Custody procedures. A chain of custody form will be included with each component material going into the 17 18 composite to show the identity of the component material 19 and each transfer of custody from its point of origination 20 to preparation of the composite. 21 22 Once prepared, each composite will be accompanied by a new 23 chain of custody until it reaches the incineration testing 24 facility. 25 26 For documentation, the facility preparing a given composite 27 will generate a report to be submitted to EPA with the 28 final report for Phase II incineration testing.

A.4-3

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#### APPENDIX B.1 1

#### GUIDELINE FOR THERMOGRAVIMETRIC ANALYSIS 2

3 4

ASTM E 1868-02 "Standard Test Method for Loss-On-Drying by 5 Thermogravimetry" will be used as the guideline for conducting the analysis described in Appendix C.2.3 with the following modifications for this testing program:

7 8

6

Section	Modification
2.1	• Standard practices at the University of Dayton Research Institute (UDRI) may be used as references throughout the standard in place of the ASTM standards noted in this section.
4.1	• The loss-on-drying value specified in the second through fifth sentences of this section will not be recorded.
7.1.3	<ul> <li>The programming rate of the furnace will be set at 10 to 25°C/min, rather than 5°C/min. Pursuant to section 11.6, the temperature program rate will be documented in the report.</li> <li>The isothermal temperature within the range of 25 to</li> </ul>
7.1.4	<ul> <li>1000°C will be maintained ±3°C, rather than ±2°C.</li> <li>The specimen atmosphere control system will be capable of supplying dry air in addition to "inert dry gas (usually purified grade nitrogen)".</li> </ul>
7.1.7	• The temperature program rate will be set at 10 to 25°C/min, rather than 5°C/min. Pursuant to section 11.6, the temperature program rate will be documented in the report.
	<ul> <li>The temperature program rate will be controlled to within the range of ±3°C/min, rather than ±0.1°C/min.</li> </ul>
	• Within the range of 25 to 1000°C, the isothermal temperature will be maintained within $\pm 3$ °C, rather than $\pm 2$ °C.
11.4	<ul> <li>The mass of the test specimen noted in the first sentence of this section will be 0.005 to 5 mg, rather than 10±1 mg (i.e., 9 to 11 mg).</li> </ul>
11.6	• The test specimen heating rate will be set at 10 to 25°C/min, rather than 5°C/min noted in the first sentence of this section. Pursuant to section 11.6, the temperature program rate will be documented in the report.
11.9	• Termination criteria will follow Test Method A as outlined in section 11.10.1.

B.1-1

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11.10.1	<ul> <li>The "fixed period of test time" mentioned in this section will be set at 5 min.</li> </ul>
11.10.1.1	<ul> <li>Loss-on-drying values will not be recorded.</li> </ul>
12.1	• The loss-on-drying value will not be calculated.
13.1.1	<ul> <li>The "identification and description of the material being tested" will be consistent with the information known to the analyst.</li> </ul>
13.1.5	<ul> <li>The loss-on-drying value will not be included in the report.</li> </ul>
14.2	<ul> <li>This section is not applicable because the Test Method A termination criteria will be used.</li> </ul>

9

## 10

## 11 Reference

ASTM E 1868-02 "Standard Test Method for Loss-On-Drying by Thermogravimetry", ASTM International. For referenced ASTM standards, visit the ASTM website, www.astm.org, or contact ASTM Customer Service at <u>service@astm.org</u>. For Annual Book of ASTM Standards volume information, refer to the standard's Document Summary page on the ASTM website.

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1 APENDIX C.1

## 2 **PFOA TRANSPORT TESTING**

C.1.1 Significance

Testing will be performed to verify that potential PFOA
emissions from the combustion testing described in Appendix
C.2 can be quantitatively transported from the high
temperature reactor into the exhaust gas sampling apparatus
(aqueous solution bubblers).

12 Acceptable PFOA transport will be demonstrated if the 13 transport efficiency (as computed in one or more of the 14 formulas below) is greater than or equal to 70%.

16 C.1.2 Experimental Plan

17 18 C.

3 4

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18 C.1.2.1 Base Plan 19

20 Transport of PFOA across the laboratory-scale thermal 21 reactor system described in Appendix C.2.4 and into the 22 exhaust gas bubblers described in Appendix D.1 will be 23 quantitatively determined as an indication of transport 24 from the high temperature reactor into the bubblers. 25

A PFOA standard of known purity greater than or equal to 97% will be gasified at 150 to 250 °C (based on thermogravimetric analysis of PFOA) with transfer line and reactor temperatures 0 to 100 °C higher than the gasification temperature.

32 Three replicate transport efficiency test runs will be 33 conducted. A minimum of one blank run will be conducted 34 prior to each transport efficiency test run. 35

36 The sample size of the PFOA standard to be gasified will be 37 less than 5 mg. The reactor exhaust gas will be collected 38 into bubbler aqueous solution as described in Appendix D.1 39 (including an HPLC water rinse of the flexible tubing [used 40 to connect the thermal reactor system and the bubbler 41 assembly] into the aqueous solution composite), which will 42 be analyzed for PFOA as described in Appendix D.2. In 43 order to provide a second way of demonstrating quantitative 44 transport, this aqueous solution composite will also be 45 analyzed for total fluorine as described in Appendix D.3. 46 (Testing for total fluorine is included due to possibility 47 of thermal degradation of PFOA under transport test

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1 conditions.) Therefore, for this transport testing the 2 amount of PFOA fed to the thermal reactor system will be sufficiently high to assure that the total fluorine input 3 4 to the thermal reactor system will be greater than 140% of 5 the mass corresponding to the limit of quantitation (LOO) 6 for total fluorine in the aqueous solution composite. (The 7 LOQ for total fluorine in aqueous solution is much higher ` 8 than the LOQ for PFOA in aqueous solution.) 9 10 The amount of PFOA and total fluorine in the thermal 11 reactor system exhaust gas will be determined via analysis 12 of the aqueous solution composite as noted above. 13 14 The amount of PFOA fed to the thermal reactor system will 15 be known based on measurement prior to gasification and 16 will be verified by weighing the pyroprobe insert cartridge 17 before and after each test run. The amount of fluorine ( input to the system will be calculated from the amount of 18 19 PFOA fed, the known purity of the PFOA, and the known 20 fluorine fraction of the PFOA standard. 21 22 PFOA transport efficiency (TE) as a percentage will be 23 computed as follows: 24 25 % PFOA TE = mass of PFOA in aqueous solution composite \* 100 (1)26 mass of PFOA fed to thermal reactor system 27 28 Total fluorine (TF) transport efficiency as a percentage 29 will be computed as follows: 30 31 Total F TE = mass of total F in aqueous solution composite \* 100 (2) 32 mass of total F fed to thermal reactor system 33 34 C.1.2.2 Contingent Testing 35 36 If the transport efficiencies for both PFOA (equation 1) 37 and total fluorine (equation 2) are less than or equal to 38 70%, then additional work will be performed. This 39 additional work will be performed in a step-wise fashion to determine if consideration of one or more of the following 40 procedural revisions enables achievement of 70% transport 41 efficiency as follows: 42 43 44 Step 1. The flexible tubing between the thermal reactor 45 system and the bubbler assembly from the experiment described in Section C.1.2.1 would be 46 quantitatively rinsed with methanol. This methanol 47 48 rinsate would be analyzed for PFOA (as described in

C.1-2

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Appendix D.2) and/or for total fluorine (as 1 described in Appendix D.3). Revised transport 2 efficiency (TE) as a percentage for PFOA (equation 3 3) and/or total fluorine (equation 4) would be 4 computed by including the mass of analyte in the 5 methanol rinse in the numerator as follows: 6 7 8 massproa out % PFOA TE = ----- \* 100 (3)9 10 mass<sub>PFOA</sub> in 11 mass of PFOA in bubbler where  $mass_{PFOA}$  out = 12 aqueous solution composite 13 + mass of PFOA in methanol 14 rinsė 15 16 mass of PFOA fed to thermal and mass<sub>PFOA</sub> in = 17 reactor system 18 19 20 masstotal F out % Total F TE = ----- \* 100 (4)21 22 mass<sub>total F</sub> in 23 where  $mass_{total F}$  out = mass of total F in 24 bubbler aqueous 25 solution composite 26 + mass of total F in 27 methanol rinse 28 29 and mass<sub>total F</sub> in = calculated mass of 30 total F in PFOA fed to 31 thermal reactor system 32 33 Step 2 (if necessary) The experiment described in Section 34 C.1.2.1 would be repeated with 35 reagent(s) (e.g. NaOH) added to the 36 bubbler aqueous solution to determine 37 if reagent addition enhances analyte 38 absorption, thereby improving transport 39 efficiency. Transport efficiency would 40 be calculated using equation (1) and/or 41 (2) above. The impact of reagent 42 addition on LOQ for PFOA analysis 43 described in Appendix D.2 would be 44 determined. 45 46 C.1.3 Reporting of Results 47 48 Following completion of PFOA transport testing as described 49 in this appendix and prior to beginning incineration 50 testing described in Appendix C.2, a letter report will be 51

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1 submitted to EPA with the transport efficiency result(s) 2 and indication of what contingent testing, if any, was 3 performed.

5 If Appendix C.2 incineration testing is performed, the 6 detailed results of Appendix C.1 transport testing will be 7 included in the test report for Appendix C.2 incineration 8 testing described in Appendix C.2.5. If Appendix C.2 9 incineration testing is not performed, the detailed results 10 of Appendix C.1 transport testing will be provided in a 11 test report for Appendix C.1 transport testing.

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APPENDIX C.2 1 2 INCINERATION TESTING 3 C.2.1 ELEMENTAL ANALYSIS 4 5 6 C.2.1.1 Introduction 7 8 Elemental analysis as described in Section C.2.1 will be 9 performed for each test substance composite to aid in preparation for combustion testing described in Section 10 11 C.2.4. 12 13 As Kissa (1998) points out, technique strongly affects analytical results for fluorinated organic compounds such 14 15 as fluorinated surfactants and fluorinated polymers due to the carbon-fluorine bond: 16 17 18 Fluorine in organic compounds is usually determined by converting organic fluorine to an 19 inorganic fluoride. Various combustion methods 20 21 are routinely used for this purpose. However, the carbon-fluorine bond is exceptionally strong, 22 23 and extremely vigorous conditions are needed for a guantitative mineralization. Conventional 24 combustion conditions used for the determination 25 of carbon and hydrogen in nonfluorinated organic 26 27 compounds are not adequate for a quantitative 28 analysis of fluorinated surfactants. 29 30 Therefore, total fluorine analysis will be performed using "extremely vigorous conditions" as described in Section 31 32 C.2.1.2, and the commercially available conventional technique used for empirical determination of carbon and 33 34 hydrogen content (described in Section C.2.1.3) will provide estimated values. 35 36 37 C.2.1.2 Total Fluorine 38 39 Each test substance composite will be characterized via analysis of total fluorine content. 40 41 Based on manufacturing process knowledge, the levels of 42 total fluorine in the components of test substance 43 composites are orders of magnitude higher than the 44 potential trace level of inorganic fluoride in these 45 materials. Therefore, for this test program, the total 46 organic fluorine value for each test substance composite 47

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1 will be considered to be the same as the total fluorine 2 value. 3 Total fluorine content will be measured via the Wickbold 4 5 Torch method; see Appendix D.3. 6 7 C.2.1.3 Carbon and Hydrogen 8 9 In order to provide information for stoichiometric 10 calculations in Section C.2.2, the carbon and hydrogen content of each test substance composite is needed. Based 11 12 on manufacturing process knowledge of the polymers in this 13 program, levels of sulfur, and nitrogen are expected to be 14 less than 0.1% and to thereby have negligible effect on stoichiometric calculations. 15 16 17 C.2.1.3.1 Theoretical Determination 18 19 Where the elemental composition of a test substance composite is known from the identity of the components in a 20 given composite, the carbon and hydrogen content of the 21 22 test substance composite can be calculated. 23 For example, where each of the components of a test 24 substance composite are polytetrafluoroethylene (PTFE), the 25 carbon and hydrogen can be determined knowing the molecular 26 27 formula for PTFE is  $(C_2F_4)_n$  as follows: 28

	number	atomic weight	weight %
carbon (C)	2	12	24
hydrogen (H)	0	1	0
fluorine (F)	4	19	76
total			100

29 30

31

C.2.1.3.2 Empirical Determination

Where compositional information on carbon and hydrogen content is not known from the identity of the components in a given composite, each such test substance composite will be analyzed for carbon and hydrogen.

36

37 As noted in Section C.2.1.1, empirical determination of 38 carbon in test substance composites via commercially 39 available conventional techniques is expected to 40 underestimate the carbon content of the test substance 41 composites due to the strength of the carbon-fluorine bond. 42 Similarly, empirical determination of hydrogen in test

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1 substance composites via commercially available 2 conventional techniques is expected to overestimate the 3 hydrogen content of the test substance composites. 4 5 The carbon content of the test substance composite can be 6 measured by determining the carbon dioxide  $(CO_2)$  generated 7 by the oxidation of the sample. This oxidation may be 8 accomplished by high temperature combustion, catalytic 9 combustion, or wet chemical oxidation. The  $CO_2$  is measured 10 directly by an infrared detector or a thermal conductivity detector, via absorption into a suitable solution (e.g., 11 potassium hydroxide) and gravimetric determination, or by 12 13 conversion to methane for measurement via a flame 14ionization detector. 15 16 The hydrogen content of the sample can be determined by 17 difference with knowledge of the fluorine content and carbon content of the sample where the moisture content and 18 19 chlorine content of the sample are negligible or known. 20 Alternatively, the hydrogen content of the sample is 21 measured by determining the water generated by high 22 temperature combustion of the sample. Measurement of water 23 in the combustion gas for this analysis may be accomplished 24 by techniques such as use of an infrared detector or 25 absorption on a dessicant with gravimetric determination. 26 With empirical hydrogen determination, it is important to 27 correct for the water in the combustion gas attributable to 28 the moisture content in the sample to obtain the hydrogen 29 content of the sample; see Section C.2.1.4. 30 31 Manufacturing process knowledge of the polymers will be 32 used to review the elemental analysis results and to form 33 the basis for interpreting non-detects. For example, if the hydrogen analytical result for a perfluorinated polymer 34 35 is less than a quantitation limit of 0.1%, then the 36 analytical result will be replaced with 0. 37 38 C.2.1.4 Moisture 39 40 Where preparation (as described in Appendix A.4) for a 41 given test substance composite has involved dewatering, the moisture (or solids) content of each such test substance 42 43 composite will be determined in order to provide a dry basis for calculations as needed. 44 45 Moisture is determined by measuring the loss of weight of 46 47 the sample when heated under controlled conditions. A

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representative sample is weighed and placed in a crucible 1 2 (or dish) and evaporated to dryness in an air or nitrogen 3 atmosphere at a defined temperature setpoint (e.g., 103 °C to 105 °C) in the range of 100 °C to 125 °C. The moisture 4 5 value is calculated as the loss in weight (difference between the starting weight of sample and the final weight 6 7 of sample) divided by the starting weight of sample. 8 Similarly, a solids value can be calculated as the final 9 weight of sample divided by the starting weight of sample. 10 C.2.2 COMBUSTION STOICHIOMETRY 11 12 13 Combustion stoichiometry calculations as described in Section C.2.2 will be performed to aid in preparation for 1415 combustion testing described in Section C.2.4 16 17 First, the weight percent values from Section C.2.1 are converted to molar quantities on a dry basis. 18 19 20 Second, based on Chapter 3 of Combustion Fundamentals for Waste Incineration (American Society of Mechanical 21 Engineers, 1974), the reaction products for these molar 22 quantities are calculated assuming complete combustion with 23 24 the following rules: 25 a) All carbon (C) in feed converts to carbon dioxide  $(CO_2)$ 26 27  $C + O_2 \rightarrow CO_2$ 28 b) All sulfur (S) in feed converts to sulfur dioxide (SO<sub>2</sub>) 29  $S + O_2 \rightarrow SO_2$ 30 31 32 c) The halogens (Cl, F) in feed convert to hydrogen halides 33  $H_2 + Cl_2 \rightarrow 2HCl$  $H_2 + F_2 \rightarrow 2HF$ 34 35 d) Hydrogen (H) present in feed in excess of that 36 required to yield products in item c) above will be 37 38 converted to water  $2H_2 + O_2 \rightarrow 2H_2O$ 39 40 41 e) Nitrogen (N) from feed or air is emitted as molecular 42 nitrogen  $N_2 \rightarrow N_2$ 43 44 Third, with these rules, the balanced chemical reaction for 45 combustion of a compound can be written. 46

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1 For example, the resulting reaction equation for a 2 hydrocarbon like methane (CH<sub>4</sub>) is 3  $CH_4 + 2 O_2 \rightarrow CO_2 + 2H_2O$ 4 5 Note that the term feed in the preceding rules (a through e) includes both material being combusted and the fuel 6 7 source of hydrogen such as methane or methanol. Additionally, stoichiometric calculations as described 8 9 above presume that the compounds undergoing combustion are essentially free of inorganic constituents. 10 11 These calculations provide the theoretical amount of oxygen 12 needed for the overall combustion reaction for the feed 13 14 based on the available information used in the 15 calculations. The initial estimate for the amount of oxygen to be used in combustion testing will be determined 16 from this theoretical amount with adjustments for target 17 18 oxygen level in thermal reactor system exhaust gas. The 19 actual amount of oxygen to be used in combustion testing 20 will be based oxygen monitoring described in Section C.2.4. 21 22 These stoichiometric calculations will also be used as needed to initially estimate and adjust experimental 23 24 conditions for combustion testing in Section C.2.4. 25 26 C.2.3 THERMOGRAVIMETRIC ANALYSIS 27 28 Thermogravimetric analysis (TGA) will be conducted to 29 determine the temperature range required for gasification 30 of each test substance composite. TGA will be conducted in 31 flowing air from room temperature to 1000°C as described in 32 Appendix B.1. 33 34 The TGA weight-loss profile for each test substance composite 35 will be evaluated to determine the temperature at which the 36 weight loss reaches a final asymptote across the temperature range investigated. This temperature corresponds to the 37 38 point at which no further gasification (under test 39 conditions) occurs for the material and will be considered 40 the temperature for complete gasification of the material. 41 42 C.2.4 Combustion Testing 43 C.2.4.1 Test Objective 44 45 46 The objective of the testing program described in Appendix 47 C.2 is to assess the potential for waste incineration of

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each test substance composite to emit PFOA, based on quantitative determination of potential exhaust gas levels of PFOA from laboratory-scale combustion testing under conditions representative of typical municipal waste combustor operations in the U.S.

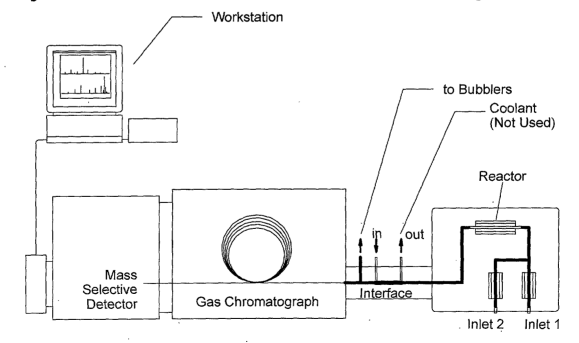
7 C.2.4.2 Experimental Apparatus

9 Combustion testing will make use of the Advanced Thermal 10 Reactor System (ATRS) at the University of Dayton Research 11 Institute (UDRI). The ATRS is a laboratory-scale, non-12 flame, batch-charged, continuous flow thermal reactor 13 system. The use of this non-flame thermal reactor system 14 gives a conservative representation of full-scale waste 15 incineration prior to air pollution controls.

In the ATRS, the test sample is gasified and transported to a high temperature reactor. In the high temperature reactor, the sample vapors are subjected to controlled conditions for residence time and temperature. As described in Sections C.2.4.5 and C.2.4.6, combustion products will be monitored or collected for quantitative analysis.

A schematic of the ATRS as configured for this test program is shown in Figure C.2-1.

### 27 Figure C.2-1. Schematic of ATRS for this Test Program



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The ATRS consists of a reactor assembly and in-line gas 1 chromatograph/detector system connected via an interface. 2 The reactor assembly consists of a thermally insulated 3 enclosure housing the sample introduction, reactor, and 4 5 transfer line systems. 6 Sample introduction for solid materials (Inlet 1) employs a 7 pyroprobe, a device designed to gasify samples by heating 8 9 them at a fixed rate. The main gas flow will also be fed 10 via Inlet 1, and Inlet 2 will be used to feed supplemental 11 flow. 12 13 During combustion tests, the transfer line between the 14 pyroprobe and the reactor is heated and maintained above 200 °C. The reactor is housed within its own small tube 15 16 furnace and may be independently heated to as high as 1100 (Actual conditions for this test program are presented 17 °C. in Section C.2.4.3.) The transfer line from the reactor to 18 19 the interface is heat traced to greater than 200 °C to 20 prevent cool regions where reactor products could otherwise 21 be lost through condensation. 22 23 The interface routes the combustion exhaust gas to the in-24 line gas chromatograph (GC) and mass selective detector 25 (MSD) or to sample collection for off-line analysis. For 26 combustion testing in this test program, the interface will 27 also be maintained above 200 °C. Exhaust gas monitoring for 28 this program is described in Section C.2.4.5. 29 30 C.2.4.3 Combustion Test Experimental Conditions 31 Each test substance composite will be subjected to 32 33 laboratory-scale incineration using the experimental 34 apparatus described in Section C.2.4.2. 35 36 C.2.4.3.1 Combustion Air 37 38 Synthetic air (mixture of 21% oxygen and 79% nitrogen) will be used in place of compressed air to prevent potential 39 40 interference in the experimental system due to background 41 levels of  $CO_2$  in compressed air. 42 43 C.2.4.3.2 Fuel 44 45 Methanol will be used, as needed, as a supplemental fuel to

ensure the presence of sufficient hydrogen to convert

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fluorine to hydrogen fluoride (HF) and chlorine to hydrogen 1 2 chloride (HCl). 3 As noted in Municipal Solid Waste in the United States: 4 2000 Facts and Figures (EPA, 2002), paper and paper 5 products (made from wood) make up the largest component of 6 municipal solid waste (MSW). The sum of paper and paper 7 products with wood in MSW makes up over 30% of MSW. 8 9 During the 19<sup>th</sup> century, methanol was produced from wood and 10 was known as wood alcohol. Therefore, methanol can be used 11 in this experimental program as a surrogate for the paper 12 and wood fraction of MSW. 13 14 C.2.4.3.3 Operating Conditions 15 16 The target operating conditions for the high temperature 17 reactor during the combustion tests for each test substance 18 composite identified in Appendix A.3 are presented in Table 19 20 C.2-1. 21 TABLE C.2-1. COMBUSTION TEST TARGET OPERATING CONDITIONS 22 1000 °C Temperature Residence Time 2 sec  $O_2$  concentration in exhaust gas 10% 15% H<sub>2</sub>O concentration in exhaust gas 3 Number of replicate runs 23 These conditions are conservatively representative of 24 typical furnace operating conditions of municipal waste 25 combustors (MWCs) and of typical secondary chamber 26 operating temperatures for medical waste incinerators in 27 the U.S. See Appendix D.4 for supporting information. 28 29 Temperature and residence time values in Table C.2-2 will 30 be fixed setpoints for these experiments. The temperature 31 of the high temperature reactor will be controlled within 32 +10 °C to assure isothermal operation. 33 34 The amount of each test substance composite fed to the ATRS 35 in this testing program will be a measured amount less than 36 5 mg. The actual amount fed, gasification rate (determined 37 from TGA), air supply, and fuel supply will be adjusted to 38 assure that the oxygen level in the exhaust will be greater 39 than or equal to the concentration in Table C.2-1 40 throughout each test to be representative of typical MWC 41 conditions. The fuel supply and air supply will also be 42

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adjusted as needed to approach the target  $H_2O$  concentration 1 2 in exhaust gas in Table C.2-1. 3 The pyroprobe section final temperature (at end of 4 temperature ramp-up) will be 750 °C or as needed to assure 5 this section is 50 to 100 °C above the highest temperature 6 7 for complete gasification across the test substance 8 composites as determined from the TGA results; see Section C.2.3. This is necessary to assure complete gasification 9 10 of the sample of test substance composite and a common set 11 of experimental conditions across the test materials during 12 combustion testing. 13 C.2.4.3.4 Blanks 14 15 A minimum of one thermal blank will be run prior to each 16 17 set of three combustion test runs for a given test substance composite. Each thermal blank run will be at the 18 corresponding combustion test conditions with all feeds 19 except for the test substance. 20 21 22 C.2.4.4 Process Monitoring 23 24 ATRS process parameters in Table C.2-2 will be monitored for each combustion test at key points during the test as 25 noted in the table. Each combustion test will be a minimum 26 of 5 minutes in duration. If the duration of a combustion 27 test is greater than 15 minutes, each parameter in Table 28 29 C.2-2 will be recorded at least once every 15 minutes. 30 TABLE C.2-2. COMBUSTION TEST MONITORING 31 Parameter Key Time for Recording Before & after gasification Temperature-Reactor Temperature-Transfer line Before & after gasification After gasification Temperature-Inlet 1 Temperature-Inlet 2 Before & after gasification Gas flow rate-Inlet 1 Before & after gasification Before & after gasification Gas flow rate-Inlet 2 Before & after combustion test Total Gas Flow rate Before & after combustion test

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Temperature-Inlet 1 will be recorded at the end of the 33

34 temperature ramp-up for gasification to monitor the

35 pyroprobe final temperature.

Pressure-Reactor

Make-up Gas (He) Flow rate

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Before & after gasification

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The flow rate of the exhaust gas routed to the bubblers 1 2 (see Section C.2.4.5.2) will be determined based on the 3 flow measurements listed in Table C.2-2. Δ 5 The amount of material fed to the system will be verified 6 by weighing the pyroprobe insert cartridge before and after 7 each experiment. 8 9 Exhaust gas monitoring is described in Section C.2.4.5. 10 11 Exhaust Gas Monitoring C.2.4.5 12 13 Combustion exhaust gas will be continuously monitored for 14 oxygen during each combustion test via in-line MSD or via 15 an oxygen monitor.  $CO_2$  in exhaust gas will be monitored via 16 in-line GC, in-line MSD, or a continuous monitor; or 17 exhaust gas will be collected in Tedlar® bags for off-line 18 analysis of CO<sub>2</sub>. Carbon monoxide (CO) in exhaust gas will 19 be monitored via in-line GC or a continuous monitor; or exhaust gas will be collected in Tedlar® bags for off-line 20 21 analysis of CO. Tedlar® bag samples may be collected at 22 the exit of the bubblers described in Section C.2.4.6. 23 24 C.2.4.6 Exhaust Gas Sampling 25 26 Gas samples for off-line analysis will be collected as described in Appendix D.1, revised as necessary pursuant to 27 Appendix C.1.2.2 if applicable. 28 29 A minimum of 60 mL of bubbler aqueous solution composite is 30 expected from each combustion test. Of this, a minimum of 31 45 mL will be directed to PFOA analysis, and the remainder 32 will be directed to fluoride ion analysis. 33 34 35 C.2.4.7 Exhaust Gas Analysis 36 37 C.2.4.7.1 Fluoride Ion 38 39 A portion of the composite bubbler aqueous solution sample 40 from each combustion test collected as described in Section C.2.4.6 will be analyzed for fluoride ion via ion 41 42 chromatography using EPA Method 300.0. 43 44 C.2.4.7.2 PFOA 45 A portion of the composite bubbler aqueous solution sample 46 from each combustion test collected as described in Section 47

C.2.4.6 will be analyzed for PFOA via LC/MS/MS as described 1 2 in Appendix D.2. 3 4 As described in Appendix D.2, composite bubbler aqueous 5 solution sample results less than method detection limit 6 (MDL) will be reported as not detected (ND), results 7 between MDL and the limit of quantitation (LOQ) will be reported as not quantifiable (NQ), and numerical values 8 9 will not be reported. 10 11 Due to background levels of PFOA, the analytical laboratory will only report numerical values for PFOA concentration in 12 13 the aqueous solution greater than or equal to the LOQ. This is required to assure that the reported concentration 14 15 value is attributable to the aqueous solution sample rather than to background. 16 17 18 C.2.5 Reporting of Results 19 20 C.2.5.1 Elemental Analysis Results 21 22 The results of elemental analysis for each test substance 23 composite (as noted in Section C.2.1) will be reported. 24 The laboratory reports will be included in an appendix to 25 the final report for incineration testing (test report). 26 27 C.2.5.2 Combustion Stoichiometry Results 28 29 Combustion stoichiometry (as noted in Section C.2.2) 30 calculations for each test substance composite will be 31 included in an appendix to the test report. 32 33 C.2.5.3 TGA Results 34 35 The temperature for complete gasification and the TGA 36 graphical results for each test substance composite (as 37 noted in Section C.2.3) will be included in an appendix to 38 the test report. 39 40 C.2.5.4 Combustion Test Results 41 42 C.2.5.4.1 Process Monitoring 43 Process monitoring data (as noted in Section C.2.4.4) 44 45 recorded for each combustion test will be reported in 46 tabular form. 47

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1 C.2.5.4.2 Exhaust Gas Monitoring 2 3 Exhaust gas  $O_2$ , CO and  $CO_2$  monitoring results will be 4 reported as the integrated or average value for each 5 combustion test. CO will be reported in terms of parts per 6 million by volume (ppmv).  $O_2$  and  $CO_2$  will be reported in 7 terms of percent by volume (%). 8 9 C.2.5.4.3 Exhaust Gas Analytical Results 10 11 Results of analyses noted in Section C.2.4.7 will be 12 reported for each replicate of each combustion test. 13 14 The analytical result for each analyte in Section C.2.4.7 will be reported in terms of concentration (mass per 15 16 volume) in the bubbler aqueous solution. For each analyte, 17 this value will be used with the associated exhaust gas 18 volume to compute an exhaust gas concentration and with the 19 associated test substance mass to compute mass of analyte 20 per mass of test substance composite. 21 22 C.2.5.4.3.1 Fluoride 23 24 Fluoride ion in the exhaust gas will be reported on the 25 basis of mass of fluoride ion per mass of test substance 26 composite. The corresponding hydrogen fluoride value for 27 each will also be computed and reported for reference. 28 29 C.2.5.4.3.2 PFOA 30 31 PFOA results for the bubbler aqueous solution samples will 32 be reported as described in Section C.2.4.7.2. PFOA 33 results for associated blanks will also be reported. 34 35 If present in the bubbler aqueous solution at a 36 concentration above the matrix-specific LOQ, PFOA in the 37 exhaust gas will be reported on the basis of mass of PFOA 38 per mass of test substance composite. 39 40 C.2.5.5 Release Assessment 41 42 In the event that PFOA is reported for the exhaust gas bubbler aqueous solution at a concentration at or above the 43 LOQ (as defined in Appendix D.2) for two or more of the 44 three runs for a given test substance composite, a release 45 46 assessment report for the full-scale waste incineration of products represented by the test substance composite will 47

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be prepared following the outline in Appendix E.2 and will 1 2 be included in the test report.

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4 C.2.5.6 Test Report Outline

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6 The outline for the test report is presented in Appendix 7 E.3. All reporting discussed in Sections C.2.5.1 through 8 C.2.5.5 will be included in this test report, as 9

applicable.

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## 1 APPENDIX D.1

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## 2 EXHAUST GAS SAMPLING VIA BUBBLERS

Gas samples for off-line analysis will be collected from a vent line off the interface of the thermal reactor system described in Appendix C.2.4. Flexible (silicone or equivalent) tubing will connect the vent line and a set of bubblers.

10 Gas absorption via these bubblers will provide aqueous solution (of documented content) to analyze for prescribed 11 12 parameters. Two to four bubblers (low pressure drop impingers) in series will be used. Each bubbler will 13 contain a predetermined amount of aqueous solution, and the 14 total amount of solution at the beginning of each test run 15 will be a minimum of 55 mL. The temperature of the gas 16 17 exiting the last bubbler will be monitored.

19 An additional bubbler (which is empty) will be added to the 20 front end of this series of bubblers to serve as a knock-21 out pot if calculations or preliminary measurements 22 indicate that greater than 10 mL of water will be produced 23 during the testing for a given material. 24

Upon completion of sample collection, the amount in each bubbler will be weighed and recorded, and the contents of the bubblers will be composited for subsequent analysis. Additionally, the flexible tubing will be rinsed with 5 mL of HPLC water to collect potential condensate in the tubing; this rinsate will be combined with the bubbler composite prior to analysis.

33 Bubbler aqueous solution composites will be conveyed to 34 analytical laboratory(ies) in polyethylene, polypropylene, 35 or glass container(s).

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1 APPENDIX D.2 2 PFOA ANALYSIS BY LC/MS/MS 3 4 D.2.1 Introduction 5 Samples to be analyzed for PFOA in this study will be 6 subjected to Liquid Chromatography with Tandem Mass 7 Spectrometry (LC/MS/MS) in accordance with "Method of 8 9 Analysis for the Determination of Ammonium Perfluorooctanoate (APFO) in Water Revision 1" (Exygen 10 method) revised per the section-by-section comments listed 11 in Section D.2.4 below. These revisions are necessary to 12 adapt a method originally developed for liter quantity 13 water samples to samples related to testing described in 14 Appendix C. 15 16 The testing programs described in Appendix C are expected 17 to generate samples of aqueous solution, methanol (e.g., as 18 used for extraction or rinsing), and corresponding blanks. 19 The expected sample size for aqueous solution samples (from 20 exhaust gas bubbler sample collection) available for 21 analysis via this method is approximately 50 mL. 22 23 24 D.2.2 Method Summary 25 PFOA is extracted from water using a disposable  $C_{18}$  solid 26 phase extraction (SPE) cartridge. PFOA is eluted from the 27 cartridge with methanol. Quantification of PFOA is 28 accomplished by electrospray liquid chromatography/tandem 29 mass spectrometry (LC/MS/MS) analysis. 30 31 32 D.2.3 Reporting 33 The target limit of quantitation (LOQ) for this study with 34 this method is 50 ng/L based on prior work with water 35 samples where an 8-fold concentration via extraction using 36  $C_{18}$  SPE cartridge has been demonstrated. The actual LOQ 37 will be matrix dependent; for samples (e.g., methanol 38 rinsate) where the 8-fold concentration cannot be 39 performed, the target LOQ for this study is 400 ng/L. 40 41 Sections 4.5.4 and 5 of the Exygen method explain reporting 42 for field samples such as bubbler aqueous solution 43 composites, which are distinct from blanks and spikes, as 44 45 follows: 46 Field samples in which either no peaks or peaks 47

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less than the MDL are detected at the corresponding analyte retention time will be reported as ND (not detected). Samples in which peaks are detected at the corresponding analyte retention time that are less than the LOQ and greater than or equal to the MDL will be reported as NQ (not quantifiable).

8 Therefore, sample results less than method detection limit 9 (MDL) will be reported as ND, and sample results between 10 MDL and the limit of quantitation (LOQ) will be reported as 11 NQ. Numerical values will not be reported for such 12 samples. Only concentrations above the LOQ, where the 13 reported concentration is attributable to the sample rather 14 than to background, are reported with numerical values.

Additionally, if the PFOA anion is found in a sample at a concentration above the LOQ for the matrix but is less than 5 times the concentration found in the associated blank, the result will be flagged and treated as ND.

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21 D.2.4 Study-Specific Comments on the Method

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Section	Comment
1	<ul> <li>The concentration of PFOA found will be reported directly and the mathematical conversion for reporting as APFO mentioned in the 4<sup>th</sup> sentence of the 2<sup>nd</sup> paragraph will not be performed.</li> <li>Since the 8-fold concentration described in the 2<sup>nd</sup> sentence 4<sup>th</sup> paragraph (which forms the basis for the LOQ in the 3<sup>rd</sup> paragraph and the MDL in the 4<sup>th</sup> paragraph) is dependent on having a minimum of 40 mL of aqueous sample amenable to extraction using the C<sub>18</sub> SPE cartridge described in section 4.4 of the method, the LOQ and MDL in the method will be a factor of 8 higher than reported where less than 40 mL of sample is available or where the sample is not amenable to extraction using the C<sub>18</sub> SPE cartridge described in section 4.4 of</li> </ul>
2 2 Mate	the method (e.g., methanol).
3.3 Note at top of page 8	<ul> <li>The note stating "Equivalent materials may be substituted for those specified in this method if they can be shown to produce satisfactory results" will not be used in the analysis for this testing program.</li> </ul>

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3.3	• The following text will be used in place of
Notes,	Note 1 with respect to the PFOA analysis
Note 1	conducted for this testing program:
3.3	In order to avoid contamination, the use of disposable labware (tubes, pipets, etc.) is required.
Notes, Note 4	• The following text will be used in place of Note 4 with respect to the PFOA analysis conducted for this testing program:
	Solvents (e.g., methanol) used for this analysis must be checked for the presence of contaminants by LC/MS/MS before use.
3.5 opening text prior to 3.5.1	• Where the available amount of sample is expected to be much less than 1 liter, insufficient sample is available to prepare the fortified matrix spikes described in the opening text of section 3.5. In this case, the analytical standards discussed in this opening text will be limited to two purposes since the third purpose (matrix spike) stated in the method cannot be done.
4.3, item b	<ul> <li>Where the available amount of sample is expected to be less than 80 mL (= 2 * 40), the replicate extraction noted in the first sentence of this item cannot be performed.</li> <li>If the sample is not an aqueous sample amenable to extraction using the C<sub>18</sub> SPE cartridge described in section 4.4 of the method, then section 4.4 is skipped such that the sample is analyzed directly. (Note: For such samples, the LOQ and MDL will be 8 times higher than the values quoted in the method.)</li> </ul>
4.3, item c	<ul> <li>As noted in comment on section 3.5 opening text above, fortified matrix spikes will not be prepared when the available amount of sample is much less than 1 liter.</li> <li>Where the available amount of sample is expected to be less than 80 mL (= 2 * 40), the conditional repeat fortification and extraction described in the third sentence of this item cannot be performed.</li> </ul>
4.4	• Extraction using the $C_{18}$ SPE cartridge requires a suitable aqueous sample. This extraction and the corresponding 8-fold concentration

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	pointed out in the NOTE at the end of this section cannot be performed on non-aqueous (e.g., methanol) samples.
4.4, item 1	• In order to measure out the 40 mL mentioned in this item, it is necessary to have at least 45 mL of sample to enable pipet transfer.
4.5.4, item g	• A storage stability study for PFOA in water performed independently of the development of the method indicates that PFOA may be stored in glass, polystyrene, polypropylene, or polyethylene containers without measurable degradation for up to 68 days prior to extraction. Therefore, the total holding time between sample collection and analysis for aqueous PFOA samples in this study may exceed the 14 day limit noted in the first sentence of this item provided that the sample is not held for greater than 68 days unless additional storage stability testing justifies a longer hold time.
4.6, item 3	• As noted in comment on section 3.5 opening text above, fortified matrix spikes will not be prepared when the available amount of sample is much less than 1 liter. In this case, acceptance criteria for matrix spike recoveries will not be considered.
5, item c	• The calculation in equation 3 in this section will not be performed since it is not necessary to convert the PFOA analytical results to APFO for this study.

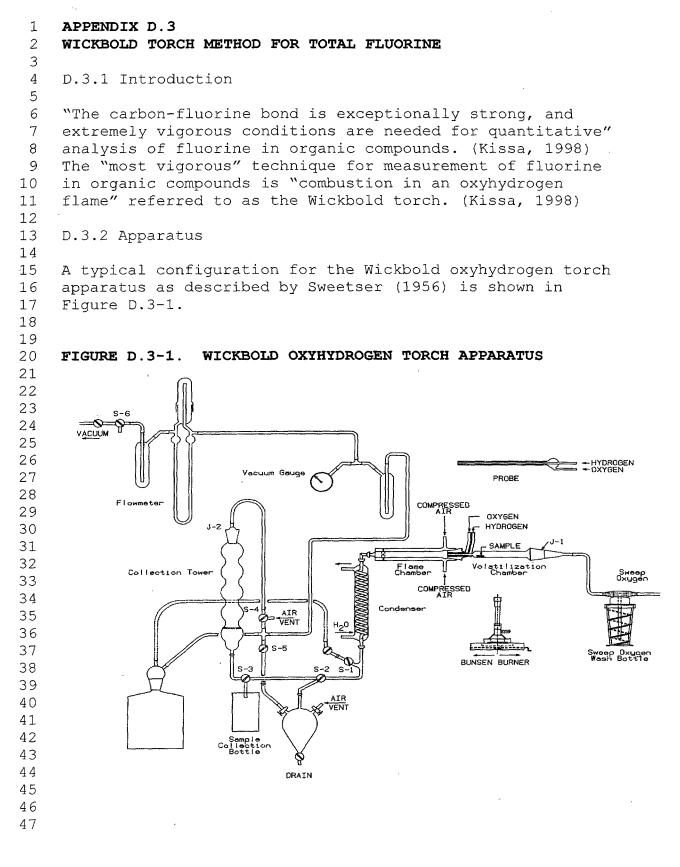
D.2.5 Reference

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Flaherty, J. and K. Risha, "Method of Analysis for the
Determination of Ammonium Perfluorooctanoate (APFO) in
Water Revision 1", Exygen Method No. 01M-008-046 Revision
January 2003. (EPA Docket ID OPPT-2003-0012-0040)

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D.3.3 Method Description 1 2 3 The sample size for the standard sample boat is up to 20 mg 4 for a solid or up to 5 mL for a liquid. 5 6 With the oxyhydrogen torch in operation, the sample is 7 pyrolyzed or vaporized with a Bunsen burner moving on a rail below the volatilization chamber. The vapors and 8 pyrolysis products are swept through the oxygen-hydrogen 9 flame chamber operating at up to approximately 2000 °C to 10 mineralize the fluorine in the sample to fluoride ion. 11 The resulting fluoride ion is absorbed in the collection tower 12 containing water or an alkaline solution. 13 14 The absorbed fluoride ion is measured via fluoride ion-15 selective electrode or ion chromatography. 16 17 The reported limit of quantitation for total fluorine via 18 19 the Wickbold Torch method is 0.5 ppm (0.5 mg/kg). The accuracy of this method for determination of total fluorine 20 21 in fluorinated polymers is exemplified by total fluorine values of 75.35% to 75.84% for PTFE with known total 22 fluorine content of 76.0%. (Sweetser, 1956) 23 24 25 D.3.4 Safety Considerations 26 Use of hydrogen presents a potential fire and explosion 27 hazard. Use of oxygen presents a potential fire hazard. 28 Safe operation of the oxyhydrogen torch is assured by the 29 use of specialized equipment with shielding and elaborate 30 safety devices by well-trained personnel at a qualified 31 32 laboratory. 33 34 D.3.5 References 35 Kissa, E. "Analysis of Anionic Fluorinated Surfactants", 36 Chapter 8 in Anionic Surfactants: Analytical Chemistry -37 2nd Edition, Revised and Expanded, edited by John Cross. 38 Marcel Dekker Surfactant Science Series, volume 73, 1998. 39 40 Sweetser, P. B. "Decomposition of Organic Fluorine 41 Compounds by Wickbold Oxyhydrogen Flame Combustion Method", 42 Analytical Chemistry, vol. 28, pp. 1766-1768, 1956. 43

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## 1 APENDIX D.4

### 2 WASTE INCINERATION AND OPERATION CONDITIONS

4 Polymers of the sort being investigated in this testing 5 program may be present at trace to low concentrations in 6 the feedstreams to municipal waste combustors and/or 7 medical waste incinerators in the U.S.

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## D.4.1 Types of Incinerators

- 11 D.4.2.1 Municipal Waste Combustors
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According to the Integrated Waste Services Association (IWSA), there are a total of 98 waste-to-energy facilities operating municipal waste combustors (MWCs) in the U.S. as of 2002. (IWSA 2002) Table D.4-1 summarizes the number and annual capacity of these units by type of technology employed.

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## 20 Table D.4-1. MWCs in 2002

Туре	Number of	Annual Capacity	Fraction		
	Facilities	(million Ton/year)	of Waste		
Mass Burn	68	22.5	76.5%		
Refused Derived	18	6.4	21.8%		
Fuel (RDF)					
Modular	12	0.5	1.7%		
Total	98	29.4	100.0%		

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22 D.4.1.2 Hospital/Medical/Infectious Waste Incinerators 23

24 Although earlier reports indicated approximately 2400 medical waste incinerators in the U.S. in the 1990s burning 25 approximately 846 thousand tons of hospital and 26 medical/infectious waste (EPA 1997), the current EPA Office 27 28 of Air Quality, Planning, and Standards (OAQPS) inventory 29 indicates that there are 116 hospital/medical/infectious 30 waste incinerators (HMIWIS) in the U.S. as of July 28, 31 2003. (EPA 2003)

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33 This represents a greater than 90% reduction in the number of operating HMIWIs in the U.S. Many medical waste 34 35 incinerators were closed rather than upgraded to meet new emission standards, as hospitals improved their programs to 36 37 segregate infectious ("red bag") waste burned in HMIWIs 38 from non-infectious ("black bag") waste handled as municipal solid waste after it leaves the hospital. 39 Consequently, the amount of segregated infectious waste 40

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1 burned in HMIWIs is expected to be less than 0.3 million 2 tons per year.

4 EPA notes that over 97% of medical waste incinerators are 5 controlled air modular units (EPA 2000a). Recent 6 communication with EPA OAQPS indicates that virtually all 7 existing HMIWIS are controlled air modular (two-chamber) 8 units.

## 10 D.4.2 Incinerator Operating Conditions

12 Many incinerators for municipal solid waste are designed to operate in the combustion zone at 1800 °F (982 °C) to 2000 13 14 °F (1093 °C) to ensure good combustion. (EPA 1995) EPA's new source performance standards (NSPS) and emission 15 16 guidelines for both municipal waste combustors (MWCs) and 17 hospital/medical/infectious waste incinerators (HMIWIs) are 1.8 based on the use of "good combustion practices" (GCP). (EPA 1997, EPA 2000b, EPA 2000c, Van Remmen 1998) 19 20

Referring to MWCs, Donnelly notes, "Design of modern efficient combustors is such that there is adequate turbulence in the flue gas to ensure good mixing, a hightemperature zone (greater than 1000 °C) to complete burnout, and long enough residence time at high temperature (1-2 sec) for complete burnout." (Donnelly 2000) The term "flue gas" here refers to the gas above the grate.

With respect to HMIWIS, Van Remmen states "any unit which presently [prior to compliance date] has a [secondary chamber] residence time less than two seconds at 1000 °C does not meet the requirement for good combustion under the new regulations." (Van Remmen 1998)

35 Similarly, most MWCs operate with a 2 second gas residence 36 time in the high temperature zone in order to assure 37 compliance with emission standards on carbon monoxide (CO) 38 and dioxins.

40 D.4.2.1 MWC Operating Conditions

42 D.4.2.1.1 Mass Burn MWC

44 Review of the IWSA Directory (IWSA 2002) indicates that 45 almost all of these mass burn units are mass burn water 46 wall furnaces. Nearly all mass burn water wall furnaces

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1 have reciprocating grates or roller grates to move the 2 waste through the combustion chamber. (EPA 1996a) 3 4 Studies on the Millbury, Massachusetts mass burn water wall 5 MWC produced gas temperature versus residence time results. 6 (Scavuzzo, Strempek, and Strach 1990) Calculations based 7 on Figure 6 of this paper indicate a time-averaged temperature of 2238  $^{\circ}\text{F}$  (1226  $^{\circ}\text{C})$  over a 2 second. The 8 9 corresponding gas temeperature at the 2 second level from 10 this figure is 1750 °F (954 °C). 11 12 A report on the Warren County, New Jersey mass burn water wall MWC indicates that the design gas temperature between 13 14 the grate and secondary air inject was greater than 2000 °F 15 (1093 °C) over a gas residence time of an additional 2.2 16 seconds. (Schuetzenduebel and Nobles 1990) This report 17 also shows that this MWC was designed for 2 seconds 18 residence time above 1800 °F (982 °C) between the 19 introduction of secondary air and the exit of the furnace 20 section. (Schuetzenduebel and Nobles 1990) The temperature 21 profile (Figure 21) in the temperature correlation test 22 report (Schutzenduebel 1989) for this MWC shows the full load gas temperature at the secondary air injection point 23 24 is 2650 °F, and the gas temperature at the 2-second point is 25 1850 °F. Therefore, testing indicates an average temperature of 2250  $^{\circ}\text{F}$  (1232  $^{\circ}\text{C})$  over this 2 second gas 26 residence time for the Warren County unit. A related 27 28 report for the Warren County MWC by the design firm 29 indicates that the exhaust gas oxygen concentration is 30 nominally 10% (dry basis). (Blount Energy Resource Corp. 31 1989) 32 Information from these 2 MWCs demonstrates that the average 33 / 34 gas temperature across a 2 second residence time for mass 35 burn MWCs is conservatively expected to be greater than 1100 °C. 36 37 38 Test report data from a typical mass burn MWC (Fairfax, 39 Virginia) indicates typical average furnace exit gas 40 concentrations are 10.8% oxygen (dry basis) and 18.4% 41 moisture (water). (Clean Air Engineering, 1997) 42 43 As indicated in Table D.4.1, mass burn units account for 44 over 76% of the municipal solid waste incinerated in the 45 U.S.

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#### 2 D.4.2.1.2 RDF MWC

Furnace temperatures as well as flue gas oxygen and moisture
(H<sub>2</sub>O) levels for the Mid-Connecticut RDF combustor during
performance tests while operating under good combustion
conditions across a range of steam loads (Finklestein and
Klicius 1994) are summarized in Table D.4-2.

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### Table D.4-2. RDF MWC - Mid-Connecticut

Steam load test number	low PT-13	low PT-14	inter- mediate PT-10	inter- mediate PT-02	normal FT-09	normal	normal PT-11	high PT-12
Furnace temperature (°C)	965	1004	1012	1022	1033	1015	1026	1049
flue gas $O_2$ (%)	10.1	9.6	9.2	9.1	7.6	7.5	7.9	6.4
flue gas moisture	12.4	11.1	12.3	15.4	15.1	16.3	14.1	16.2

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12 The average operating conditions for this RDF unit across 13 the range of steam loads are 1016  $^{\circ}$ C, 8.4% O<sub>2</sub> (dry basis), 14 and 14.1% moisture. 15

Examination of the report and MWC temperature monitoring practices indicates that these temperatures are effectively combustion zone exit temperatures. Therefore, in order to determine the average MWC combustion zone temperature across a 2 second gas residence time, it is necessary to understand the time-temperature profile of the MWC.

Since waste combustion in this and most other RDF units in 23 24 the U.S. involves burning on the grate (EPA 1996a) similar to the operation of mass burn MWCs, the time-temperature 25 profile in an RDF unit is expected to be similar to that 26 described in Section D.4.2.1.1 above. Based on this 27 similarity and the temperatures in Table D.4-2, the average 28 gas temperature across a 2 second residence time for RDF 29 30 units is conservatively expected to be greater than 1100 °C. 31

32 As indicated in Table D.4.1, RDF units account for 33 approximately 22% of the municipal solid waste incinerated 34 in the U.S.

36 D.4.2.1.1 Modular MWC

Modular MWCs are generally small dual-chamber units, accounting for less than a total of 2% of the municipal solid waste incinerated in the U.S. in 2002. Modular MWCs are generally equipped with auxiliary fuel burners in the

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1 secondary chamber. (EPA 1996a) EPA notes that the 2 secondary chamber exit temperature of modular MWCs is 3 maintained at typically 980 to 1200 °C. (EPA 1996a) 4 5 A typical modular MWC in Polk County, Minnesota is operated 6 with a gas residence time of 2 seconds, in the secondary 7 chamber, a secondary chamber exit temperature in the range of 1800 °F (982 °C) to 2000 °F (1093 °C), flue gas oxygen 8 concentrations in the range of 10% to 13% (dry basis), and 9 flue gas moisture in the range of 10% to 15% (Pace 10 11 Analytical 2003). 12 13 Since the secondary chamber exit temperature is expected to 14 be the minimum gas-phase temperature for the chamber, the secondary chamber average gas temperatures for modular MWCs 15 are expected to be 1000  $^{\circ}\mathrm{C}$  or greater. 16 17 18 As indicated in section D.4.1, such modular units are 19 generally small MWCs and account for less than a total of 20 2% of the municipal solid waste incinerated in the U.S. 21 22 D.4.2.1.4 MWC Summary 23 Considering the relative quantities of municipal waste 24 25 burned annually in each type of MWC and the data in this section, typical operating conditions for the high 26 27 temperature zone of most MWCs are >1000 °C average 28 temperature across 2 second residence time with exit gas 29 concentrations of 10%  $O_2$  (dry basis) and >15% moisture. 30 31 D.4.2.2 HMIWI Operating Conditions 32 33 The range of temperatures for the secondary chamber of controlled air medical waste incinerators has been reported 34 as 980 to 1200 °C. (Theodore 1990) EPA notes that auxiliary 35 36 fuel (e.g., natural gas) is burned in the secondary chamber. 37 of medical waste incinerators to sustain temperatures in the range of 985 to 1095 °C and that combustion air at 150 38 to 250 % of the stoichiometric requirement is usually added 39 40 to the secondary chamber. (EPA 2000a, EPA 1994a). 41 42 In its model plant description background document, EPA 43 notes that the average moisture content in HMIWI flue gas 44 was about 10 % based on available data, and EPA states 45 "limited data show that older [HMIWI] units typically have 46 residence times that range from essentially 0 seconds up to 47 about 1 second." (EPA 1994b) However, as noted above, a

more recent report indicates that HMIWIS still in operation 1 have secondary chamber temperatures greater than or equal 2 to 1000  $^{\circ}\mathrm{C}$  with a gas residence time of 2 seconds. (Van 3 4 Remmen 1998) For example, EPA studied the incinerator at Weeks Hospital in New Hampshire as a typical HMIWI with a 5 design residence time of 2 seconds in the secondary 6 7 chamber. (EPA 1996b) During this testing, the average exit 8 secondary chamber exit temperature was 1024 °C, and the flue 9 gas oxygen concentration was 13.5%. (dry basis) (EPA 1996b) 10 Review of test reports for all HMIWIS in the EPA docket for 11 the HMIWI NSPS and EG rulemakings that are listed in EPA's 12 current HMIWI inventory (EPA 2003) does not refute Van 13 14 Remmen's statement above on residence time and temperature and indicates HMIWI flue gas oxygen concentrations for 15 these units in the range of 10 to 15% (dry basis) and stack 16 17 moisture concentrations as high as 30% (after wet (Environmental Laboratories Inc. 1993, EPA 18 scrubbing). 1996, HDR Engineering 1994a, HDR Engineering 1994b, METCO 19 Environmental 1992, Technical Services, Inc. 1993, 20 21 Technical Services, Inc. 1994a, Technical Services, Inc. Apparently, the older HMIWIs referred to in EPA's 22 1994b) model plant description background document either have 23 been shut down or upgraded to operate with secondary 24 chamber exit temperatures higher than 1000 °C at a gas 25 26 residence time of 2 seconds. 27 28 Secondary chamber temperature of HMIWIs is monitored near the secondary chamber outlet. (EPA 1994) Hence, when the 29 30 auxiliary burner (located on the end opposite from the outlet) is in use, the average gas temperature in an HMIWI 31 secondary chamber is greater than the outlet temperatures 32 noted above. Therefore, secondary chamber average gas 33 temperatures for HMIWIs are expected to be 1000 °C or 34 35 greater with a gas residence time of 2 seconds. 36 In summary, typical operating conditions for the secondary 37 chamber of operating HMIWIs in the U.S. are 1000 °C average 38 temperature across 2 second residence time with exit gas 39 concentrations of 13%  $O_2$  (dry basis) and >10% moisture. 40 -41 42 D.4.3 Pollution Control Equipment 43 Over 99% of large MWC capacity operates with a spray dryer 44 absorber/scrubber. (IWSA 2003) Approximately 80% of large 45 MWC capacity operates using carbon injection as part of the 46 pollution control system. (IWSA 2003) Due to requirements 47

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in the NSPS (EPA 2000b) and EG (EPA 200c) for small MWCs, small MWCs planning continued operation are generally upgrading or have upgraded their pollution control equipment to add spray dryer absorbers or other acid gas control and carbon injection.

7 Review of EPA's HMIWI inventory (EPA 2003) indicates that 8 essentially all HMIWIs have some form of wet or dry 9 scrubbing for acid gas control.

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# 11 D.4.4 Summary

13 Approximately 30 million tons per year of municipal solid waste was combusted in the United States annually in waste-14 to-energy municipal waste combustors in 2003. 15 Approximately 0.3 million tons per year of segregated 16 medical waste was combusted annually in the United States 17 in hospital/medical/infectious waste incinerators in 2003. 18 Considering the relative amounts of waste combusted 19 annually, typical operating conditions for waste 20 incineration in the U.S. across these two classes of units 21 are as follows: 22

Average Temperature	>1000 °C
Residence Time	>2 sec
$O_2$ concentration in exhaust gas	$\overline{10\%}$ (dry basis)
H <sub>2</sub> O concentration in exhaust gas	15%

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23

EPA emission regulations currently in place or in place by 26 2005 require that operating municipal waste combustors and 27 hospital/medical/infectious waste incinerators have or will 28 have air pollution control equipment such as wet or dry 29 scrubbing for acid gas control. 30

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1		NDIX E.1
2	OUTL:	INE FOR INTERIM PROGRESS REPORTING
3		
4 5 7	Title	e: Enforceable Consent Agreement for the Laboratory- Scale Incineration Testing of Fluorotelomer Based Polymers - Interim Report
8 9	OPPT	Docket ID No: OPPT-2004-0001
10		
11 12 13	Date	of Interim Report: [ date ]
14 15 16	This	Report covers the period from [date] to [date]
17 18 19 20 21	1)	List or description of significant ECA Test Program milestones during this period:
22 23 24 25 26 27	2)	Description of Difficulties: (If none indicate N/A)
28 29 30 31 32 33 34 35 36	3)	Actions taken in response to difficulties: If none indicate N/A)
37 38 39 40 41 42 43	4)	Other information relevant to the progress of the testing program: (If none indicate N/A)

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1 2 2		NDIX E.1 (continued) INE FOR INTERIM PROGRESS REPORTING
3 4		
4 5 6 7 8	Title	e: Enforceable Consent Agreement for the Laboratory- Scale Incineration Testing of Fluoropolymers - Interim Report
9 10	OPPT	Docket ID No: OPPT-2003-0071
11 12 13	Date	of Interim Report: [ date ]
14 15 16 17	This	Report covers the period from [date] to [date]
18 19 20 21	1)	List or description of significant ECA Test Program milestones during this period:
22 23 24 25 26 27 28	2)	Description of Difficulties: (If none indicate N/A)
29 30 31 32 33 34 35 36 37	3)	Actions taken in response to difficulties: If none indicate N/A)
38 39 40 41 42	4)	Other information relevant to the progress of the testing program: (If none indicate N/A)

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### 1 APPENDIX E.2

### 2 OUTLINE FOR RELEASE ASSESSMENT REPORT

4 As described in Appendix C.2.5.5 of this ECA, if PFOA is reported for the exhaust gas bubbler aqueous solution at a 5 concentration at or above the LOQ (as defined in Appendix 6 7 D.2) for two or more of the three runs for a given test 8 substance composite, then the potential for release from full-scale municipal and/or medical waste incineration, as 9 applicable, (including application of air pollution 10 controls) of products represented by the test substance 11 composite in the United States will be assessed to put the 12 data into perspective. At a minimum, the report will 13 follow the general outline described below and will state 14 assumptions, document the basis for the assumptions made, 15 quantitatively estimate the variability of calculated 16 estimates (based on the variability of the parameters in 17 the evaluation), and qualitatively discuss the uncertainty 18 of calculated estimates. 19

- 21 1.0 Introduction
- 22 23

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• Statement of objective for combustion testing of test substance composites.

• Applicability of the laboratory-scale combustion testing to full-scale municipal waste combustors. (MWCs) and/or medical waste incinerators (as applicable) in the United States.

#### 31 2.0 Summary of study results

• A listing of exhaust gas analytical results reported for each applicable test substance composite.

• A listing of test substance composite analytical results reported for each applicable test substance composite.

### 40 3.0 Discussion

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 Description of the combustion section of the applicable waste incineration process(es) being evaluated (MWC and/or medical waste incinerator) including the rationale for selecting test target temperature(s) and description of typical

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1		operational parameters. Cross-reference to or
2		submission of relevant parts of Appendix D.4 of this
3		ECA can satisfy this provision.
4		
5		<ul> <li>Decorrintion of the next-combustion oin pollution</li> </ul>
		<ul> <li>Description of the post-combustion air pollution</li> </ul>
6		control equipment (e.g., lime scrubbing, carbon
7		adsorption) employed by typical operating full-scale
8		waste incineration process(es) as applicable.
9		
10	4.0	Extrapolation of laboratory test results to the
11		typical waste incineration process(es), as applicable,
12		described in Section 3.0 (above) for each test
13		substance composite to be evaluated.
14		-
15		<ul> <li>The relevance of the subject test substance</li> </ul>
16		composite to MWCs and/or medical waste incinerators.
17		
18		<ul> <li>The estimated concentration of the subject test</li> </ul>
19		substance composite to the applicable type(s) of
20		waste incinerator. Available information on
21		hydrogen fluoride concentration in waste incinerator
22		exhaust can provide the basis for an upper bound on
23		this estimated concentration.
24		
25		• A description of the extrapolation.
		• A description of the exclaporation.
26		
27	•	<ul> <li>A description of any assumptions used.</li> </ul>
· · · · · ·		
28		
29		<ul> <li>Any unique qualitative or quantitative descriptors</li> </ul>
30		of the test, the testing equipment, and the results
31		deemed necessary for informative review of the test
32		and test results.
33		
	E 0	Constitution Booleania
34	5.0	Sensitivity Analysis
35		
36		<ul> <li>Assessment of the impact of variability</li> </ul>
37		(quantitative) and uncertainty (qualitative) in each
38		parameter on the evaluation results.
39		-
	<i>c</i> ^	Conclusions
40.	6.0	CONCLUSIONS
41		
42	7.0	References

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1	APPENDIX E.3
2	OUTLINE OF TEST REPORT <sup>*,**</sup>
3	
4	
5	Table of Contents
6	
7	Executive Summary
8	Excederve buildery
9	Introduction
10	Incroduceton
	I Dhace I DECA Twomenent Testing
11	I. Phase I PFOA Transport Testing
12	
13	1. Experimental Apparatus
14	2. Description of Test Conditions (including deviations
15	from protocol)
16	3. Documentation of PFOA Standard
17	4. Analytical Results
18	4.1 PFOA
19	4.2 Total Fluorine
20	5. Transport Efficiency
21	5.1 PFOA
22	5.2 Total Fluorine
23	6. Discussion of Results
24	7. Conclusions
25	
26	II. Phase II Incineration Testing (provided Phase II is performed)
27	
28	1. Documentation of Test Substance Composites
29	2. Elemental Analysis Results
30	3. Combustion Stoichiometry Results
31	4. TGA Results
32	5. Combustion Testing
33	5.1 Experimental Apparatus
34	5.2 Description of Test Conditions (including deviations
35	from protocol)
36	5.3 Combustion Testing Results
37	5.3.1 Process Monitoring
38	5.3.2 Exhaust Gas Monitoring
39	5.3.2 Exhaust Gas Sampling and Analysis
40	5.3.2.1 PFOA
41	5.3.2.2 Fluoride
42	5.4 Discussion of Results
42 43	5.5 Conclusions
43 44	J.J CONCLUSIONS
44 45	III Appardicas
45 46	III. Appendices
47	• Quality Assurance Report(s)

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1		
2		• Report(s) from Compositing Facility(ies) (provided Phase II
3		is performed)
4		
5		<ul> <li>Reports from Analytical Laboratories</li> </ul>
6		
7		<ul> <li>Release Assessment per Appendix E.2, if applicable</li> </ul>
8		
9		
10		
11	*	Test Report will include this information (as applicable)
12		but not necessarily in this format.
13		
14	* *	References to literature in this report will include full
15		citations.

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# 1 APPENDIX F

# 2 ECA INCINERATION TESTING QUALITY ASSURANCE PROJECT PLAN (QAPP):

- 3 REQUIRED CONTENT
- 4

EP	A QA/R-5 QAPP Guidance Element	Required Content of QAPP(s) for ECA Incineration Testing
A:	PROJECT MANAGEMENT	
	Al Title and Approval Sheet	to be included in QAPP
	A2 Table of Contents	to be included in QAPP
	A3 Distribution List	to be included in QAPP
	A4 Project/Task Organization	to be included in QAPP
	A5 Problem Definition/ Background	to be satisfied by cross- reference to ECA (Parts I, IV) and Appendix A, C.1, or C.2, as applicable
	A6 Project/Task Description	see element A5
	A7 Quality Objectives and Criteria	to be satisfied by cross- reference to Appendix A, C.1, or C.2 (as applicable) and to Appendix D.2 and/or D.3, as applicable
	A8 Special Training/	for facilities subject to
	Certifications	GLP (40 CFR Part 792) under this ECA, QAPP shall state that this element is satisfied by compliance with applicable GLP requirements; for compositing facilities, to be to be satisfied by providing a statement of the qualifications for each such facility
	A9 Documentation and Records	to be satisfied by cross- reference to ECA Part XIV and Appendix E
в:	DATA GENERATION AND ACQUISITION	
	B1 Sampling Process Design (Experimental Design)	see element A5
	B2 Sampling Methods	to be satisfied by cross- reference to Appendix C.1 or C.2 (as applicable) and to Appendix D.1
	B3 Sample Handling and Custody	to be included in QAPP consistent with Appendix A
	B4 Analytical Methods	to be satisfied by cross-

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	reference to analytical
	method descriptions in
	Appendices C.2, D.2, and
	D.3, as applicable
B5 Quality Control	to be satisfied by cross-
	reference to QC provisions
	(e.g., blanks) in
	Appendices A, C.1, C.2,
	D.2, and D.3, as applicable
B6 Instrument/Equipment	to be included in QAPP in
Testing, Inspection, and	summary form for chemical
Maintenance	analysis equipment for the
	analytical methods for
	element B4 above
B7 Instrument/Equipment	to be included in QAPP in
Calibration and Frequency	summary form for chemical
	analysis equipment for the
	analytical methods for
	element B4 above
B8 Inspection/Acceptance of	for facilities subject to
Supplies and Consumables	GLP (40 CFR Part 792) under
	this ECA, QAPP shall state
	that this element is
	\ (
	satisfied by compliance
	with applicable GLP
	requirements; not
	applicable to other
	facilities
B9 Non-direct Measurements	to be satisfied by cross-
	reference to Appendix C.2.2
B10 Data Management	for facilities subject to
	GLP (40 CFR Part 792) under
	this ECA, QAPP shall state
	that this element is
	satisfied by compliance
	with applicable GLP
	requirements; for
	compositing facilities, to
	be to be satisfied by
	cross-reference to Appendix
	A.4
C: ASSESSMENT AND OVERSIGHT	
C1 Assessments and Response	for facilities subject to
Actions	GLP (40 CFR Part 792) under
	this ECA, QAPP shall state
	that this element is
	satisfied by compliance
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	with applicable GLP
	requirements; for other
	facilities, to be included
	in QAPP
C2 Reports to Management	for facilities subject to
	GLP (40 CFR Part 792) under
	this ECA, QAPP shall state
	that this element is
	satisfied by compliance
	with applicable GLP
	requirements; for other
	facilities, to be included
	in QAPP
D: DATA VALIDATION AND USABILITY	2
D1 Data Review, Verification,	to be satisfied by cross-
and Validation	reference to Appendix A,
	C.1, or C.2 (as applicable)
	and to Appendix D.2 and/or
	D.3, as applicable
D2 Verification and	for facilities subject to
Validation Methods	GLP (40 CFR Part 792) under
	this ECA, QAPP shall state
	that this element is
	satisfied by compliance
	with applicable GLP
	requirements; for other
	facilities, to be included
	in QAPP consistent with
	Appendices A, C.1, C.2,
	D.2, D.3 as applicable
D3 Reconciliation with User	to be satisfied by cross-
Requirements	reference to Appendices
redattoweree	C.2.5.5 and $E.2$ , as
	applicable
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1	APPENDIX G
2	COPY OF EPA ORDER
3	
4	
5	UNITED STATES
6	ENVIRONMENTAL PROTECTION AGENCY
7	
8	TESTING CONSENT ORDER FOR THE LABORATORY-SCALE INCINERATION
9	TESTING OF FLUOROTELOMER BASED POLYMERS
10	
11 .	Docket No. OPPT-2004-0001
12	
13	The second se
14	Under the authority of section 4 of the Toxic Substances
15	Control Act (TSCA), 15 U.S.C. 2603, the United States
16	Environmental Protection Agency (EPA) issues this testing
17	consent order (Order) to take effect on the date of publication of the notice in the Federal Register announcing the issuance of
18 19	this Order. This Order incorporates the enforceable consent
20	agreement (ECA) for the laboratory-scale incineration testing of
20	fluorotelomer based polymer test substance composites listed in
22	Appendix A of the ECA.
23	Appendix A of the left.
24	·
25	
26	
27	
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29	
30	
31	Date Stephen L. Johnson,
32	Assistant Administrator
33	For Prevention, Pesticides,
34	And Toxic Substances
35	
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1	APPENDIX G (continued)
2	COPY OF EPA ORDER
3	
4	
5 6	UNITED STATES ENVIRONMENTAL PROTECTION AGENCY
7	ENVIRONMENTAL PROTECTION AGENCY
8	TESTING CONSENT ORDER FOR THE LABORATORY-SCALE INCINERATION
9	TESTING OF FLUOROPOLYMERS
10	
11	Docket No. OPPT-2003-0071
12	
13	
14	Under the authority of section 4 of the Toxic Substances
15	Control Act (TSCA), 15 U.S.C. 2603, the United States
16 17	Environmental Protection Agency (EPA) issues this testing
18	consent order (Order) to take effect on the date of publication of the notice in the Federal Register announcing the issuance of
19	this Order. This Order incorporates the enforceable consent
20	agreement (ECA) for the laboratory-scale incineration testing of
21	fluoropolymer test substance composites listed in Appendix A of
22	the ECA.
23	
24	
25	
26	
27	·
28 29	
29 30	
31	Date Stephen L. Johnson,
32	Assistant Administrator
33	For Prevention, Pesticides,
34	And Toxic Substances

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