Exova 2395 Speakman Dr. Mississauga Ontario Canada L5K 183 T:+1 (905) 822-4111 F:+1 (905) 823-1446 E: sales@exova.com W: www.exova.com



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Surface Flammability, Smoke and Toxic Gas Generation of "3M™ Polyurethane Adhesive Sealant 540"

A Report To: 3M Center

Industrial and Transportation Division

3M Center, Building 223-1N-14

St, Paul, MN 55144

USA

Phone: (651) 733-8456

Email: hcertain@mmm.com

Attention: Harry Certain

Submitted By: Fire Testing

Report No. 11-002-639(A1)

5 pages + appendix

Date: October 18, 2011

ACCREDITATION To ISO/IEC 17025 for a defined Scope of Testing by the International Accreditation Service

SPECIFICATIONS OF ORDER

Determine surface flammability in accordance with ASTM E 162, rate of smoke generation according to ASTM E 662 and toxic gas production in accordance with Bombardier SMP 800-C and Boeing BSS 7239, as per your Purchase Order No. USMMM8U18T and our Quote No. 11-006-08170 RV1 dated September 21, 2011.

<u>IDENTIFICATION</u> (Exova sample identification number 11-002-S0639-1)

Polyurethane sealant, identified as "3M™ Polyurethane Adhesive Sealant 540".

SAMPLE PREPARATION

The coating material was applied onto 6 mm thick fiberglass reinforced cement substrate using a $1/32 \times 1/32$ " square notched trowel and was allowed to dry 48 hours prior to testing.

TEST RESULTS

ASTM E 162-11a

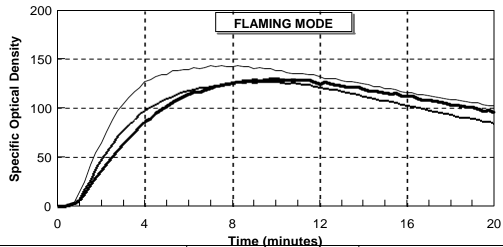
Surface Flammability of Materials Using a Radiant Heat Energy Source. (Is = Flame Spread Index).

	<u>Fs</u>	<u>Q</u>	<u>ls</u>	<u>Observations</u>
1:	2.1	2.5	5	Maximum flame front propagation to a distance of 7 inches.
2:	2.1	4.0	8	No flaming running or flaming dripping observed.
3:	1.9	2.7	5	
4:	2.1	2.7	<u>6</u>	
Rounded	Average	:	5	
Specified Maximum:			35	No flaming running or flaming dripping allowed

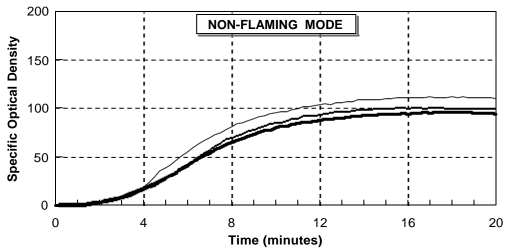
TEST RESULTS (continued)

ASTM E 662-09

Specific Optical Density of Smoke Generated by Solid Materials



Relative Room Humidity: 29%		ration: 20	,	Chamber Wall Temp: 35°C		np: 35°C
Flaming Mode Test		#1	#2	#3	Average	Specified Maxima
Specific Optical Density at 1.5 minu	22	27	42	30	100	
Specific Optical Density at 4.0 minu	86	98	126	104	200	
Maximum Specific Optical Density	130	128	143	133	-	
Maximum Corrected Optical Density	127	120	138	129	-	



Relative Room Humidity: 29%	Test Duration: 20 min.			Chamber Wall Temp: 35°C		
Non-Flaming Mode	Test	#1	#2	#3	Average	Specified Maxima
Specific Optical Density at 1.5 minu	2	1	1	1	100	
Specific Optical Density at 4.0 minu	18	17	19	18	200	
Maximum Specific Optical Density	96	102	112	103	-	
Maximum Corrected Optical Density	93	99	109	100	-	

TEST RESULTS (continued)

ASTM E 662 Observations

In the flaming mode, ignition was initially observed at the point of pilot flame impingement, increasing to full surface ignition within 25 seconds. Surface charring and visible smoke production were also observed. In the non-flaming mode, visible smoke production was observed within 30 seconds followed by surface charring.

Bombardier SMP 800-C (Rev. 6 2009-08-31)

Toxic Gas Generation from Material Combustion

		Flaming	Non-Flaming	Specified
		<u>Mode</u>	<u>Mode</u>	<u>Maxima</u>
Carbon Monoxide (CO ppm				
	at 1.5 minutes	48	<10	-
	at 4.0 minutes	250	<10	-
	at maximum	1088	533	3500
Carbon Dioxide (CO2 ppm)				
	at 1.5 minutes	550	200	-
	at 4.0 minutes	2750	250	-
	at maximum	16500	1800	90000
Nitrogen Oxides (as NO2 p	1	3	100	
Sulfur Dioxide (SO2 ppm)	25	6	100	
Hydrogen Chloride (HCl pp	11	52	500	
Hydrogen Fluoride (HF ppn	4	4	100	
Hydrogen Bromide (HBr pp	<1	<1	100	
Hydrogen Cyanide (HCN p	4	3	100	
Original Weight (g)(includin	44.0	43.3	-	
Final Weight (g)	Not determinable	Not determinable	-	
Weight Loss (g)	-	-	-	
Weight Loss (%)		-	-	-
Time to Ignition (s)		5	Did not ignite	-
Burning Duration (s)	60	-	-	
3				

TEST RESULTS (continued) Boeing BSS 7239 (Rev.: A 1-18-88)

Toxic Gas Generation

				M-7 Technical
		Flaming	Non-Flaming	Specification
		<u>Mode</u>	<u>Mode</u>	<u>Maxima</u>
Carbon Monoxide (CO ppm)	at 1.5 minutes	s 25	<10	-
	at 4.0 minutes	165	<10	-
	at maximum	n 1040	503	3500
Nitrogen Oxides (as NO2 ppm)		<1	<1	100
Sulfur Dioxide (SO2 ppm)		10	<6	100
Hydrogen Chloride (HCl ppm)		<12	74	500
Hydrogen Fluoride (HF ppm)		19	19	200
Hydrogen Bromide (HBr ppm)		<3	<3	-
Hydrogen Cyanide (HCN ppm)		3	<1	150
Original Weight(g)(including subst	rate)	49.9	43.3	-
Final Weight (g)	<u> </u>	lot determinable	Not determinable	-
Weight Loss (g)		-	-	-
Weight Loss (%)		-	-	-
Time to Ignition (s)		5	Did not ignite	-
Burning Duration (s)		60	-	-

CONCLUSIONS AND COMMENTS

The polyurethane sealant identified in this report, when tested at applied onto 6 mm thick fiberglass reinforced cement substrate, meets The Federal Railroad Administration requirements as they pertain to surface flammability (ASTM E 162) and rate of smoke generation (ASTM E 662).

The polyurethane sealant also meets Bombardier requirements as they pertain to toxic gas production (Bombardier SMP800-C).

Boeing BSS 7239 is solely a test procedure and, as such, has no specific pass/fail criteria of its own. The M-7 Technical Specification criteria are cited for reference purposes only, and may or may not apply to this specific product. The polyurethane sealant identified in this report meets the M-7 Technical Specification requirements as they pertain to toxic gas generation (Boeing BSS 7239).

Note: This is an electronic copy of the report. Signatures are on file with the original report.

Mel Garces, Ian Smith, Fire Testing. Fire Testing.

Note: This report and service are covered under Exova Canada Inc. Standard Terms and Conditions of Contract which may be found on the Exova website (www.exova.com), or by calling 1-866-263-9268.

APPENDIX

(4 pages)

Summaries of Test Procedures

ASTM E 162-11a

Surface Flammability of Materials Using a Radiant Energy Source.

Four specimens, 6×18 inches in size, are pre-dried for 24 hours at 60° C and conditioned to equilibrium at $50 \pm 5\%$ relative humidity and $23 \pm 3^{\circ}$ C before testing.

Each specimen is mounted into a holder and inclined at 30° from the vertical in front of a 12×18 inch gas-fired radiant panel. The orientation of the specimen is such that ignition is forced near its upper edge by a pilot flame, and the flame front progresses downwards.

A factor derived from the rate of progress of the flame-front and the rate of heat liberation by the material under test is calculated as follows and then reported after rounding the average of the tests to the nearest multiple of 5:

 $Is = Fs \cdot Q$

Where: Is is the flame spread index

Fs is the flame spread factor

Q is the heat evolution factor

Transit authorities generally specify a maximum Is acceptance criterion of 35 for general applications, and 100 for light diffusers, windows and transparent plastic windscreens.

ASTM E 662-09

Standard Test Method for the Specific Optical Density of Smoke Generated by Solid Materials

This method of test covers a procedure for measuring the smoke generated by solid materials and assemblies in thickness up to and including 1 inch (25.4 mm). Measurement is made of the attenuation of a light beam by smoke (suspended solid or liquid particles) accumulating within a closed chamber due to nonflaming pyrolytic decomposition and flaming combustion. Results are expressed in terms of specific optical density (Ds), which is derived from a geometrical factor and the measured optical density (absorbance).

Specimens are dried for 24 hours at 60°C and conditioned to equilibrium at 50% RH and 23°C.

Three specimens, 3" square, are exposed to each mode of combustion. Prior to test initiation, the chamber wall temperature is established in the range of 33 to 37° C. The % light transmittance during the course of the combustion is recorded. These data are used to express the quantity of smoke in the form of Specific Optical Density based on the following formula, which assumes the applicability of Bouguer's law:

$$Ds = (V/AL) \cdot log(100/T) = G \cdot log(100/T) = 132 \cdot log(100/T)$$

Where: Ds = Specific Optical Density

T = % Transmittance

V = Chamber Volume (18 ft³)

A = Exposed Area of the Sample (0.0456 ft²)

L = Length of Light Path in Chamber (3.0 ft)

G = Geometric Factor

Among the parameters normally reported are:

Ds

1.5 - specific optical density after 1.5 minutes

Ds

4.0 - specific optical density after 4.0 minutes

Dm - maximum specific optical density at any time during the

20 minute test

Dm

(corr) - Dm corrected for incidental deposits on the optical surfaces

Transit authorities generally specify a maximum Ds 1.5 of 100 and a maximum Ds 4.0 of 200 in either flaming or non-flaming test mode.

Bombardier SMP 800-C (Rev. 6 2009-08-31)

Toxic Gas Sampling and Analytical Procedures

Toxic Gas Generation

Gases produced for analysis are generated in a specified, calibrated smoke chamber during standard rate of smoke generation testing (typically ASTM E 662), in both flaming combustion and non-flaming pyrolytic decomposition test modes.

Carbon Monoxide (CO) and Carbon Dioxide (CO2)

CO and CO2 are monitored continuously during the 20 minute test using a non-dispersive infrared (NDIR) analyzer. Data are reported in ppm by volume at 1.5 and 4.0 minutes and at maximum concentration.

Acid Gas Sampling

HCN, HF, HCl, HBr, NOx and SO2 are sampled by drawing 6 litres of the chamber atmosphere through two midget impingers, each containing 10 ml of 0.25N NaOH, at a rate of 375 ml per minute. The 16-minute sampling period is commenced at the 4 minute mark. All determinations are performed in both the flaming and non-flaming modes and all data are reported in parts per million (ppm) by volume in air.

Analysis of Impingers for Hydrogen Cyanide (HCN)

Cyanide in the NaOH impinger, as NaCN, is converted to CNCI by reaction with chloramine-T at pH greater than 8 without hydrolyzing to CNO⁻. After the reaction is complete, CNCI forms a red-blue colour on addition of a pyridine-barbituric acid reagent. Cyanide is quantified by spectrometric measurement of the increase in colour 578 nm.

Reference: In-house SOP 00-13-SP-1216 based on ASTM Method D 2036-91

Analysis of Impingers for Hydrogen Fluoride (HF)

Fluoride, as NaF, in the NaOH impinger is determined using SPADNS colorimetry.

Reference: In-house SOP 01-13-SP-1295

Analysis of Impingers for Hydrogen Chloride (HCI) and Hydrogen Bromide (HBr)

Alkali halides (chloride and bromide) formed in the NaOH solution are measured using ion chromatography and conductivity detection.

Reference: In-house SOP 02-13-SP-1402

Analysis of Impingers for Nitrogen Oxides (NOX)

Nitrite and nitrate formed in the alkaline solution are determined using ion chromatography and conductivity detection. The nitrite and nitrite results are combined and the total expressed as nitrogen dioxide (NO2). Reference: In-house SOP 02-13-SP-1402

Analysis of Impingers for Sulfur Dioxide (SO2)

SO2 is trapped in the NaOH impinger as sulfite and sulfate (SO3⁻² and SO4⁻²). Hydrogen peroxide is added to convert SO3⁻² to SO4⁻². Resulting sulfate is determined using ion chromatography and conductivity detection.

Reference: In-house SOP 02-13-SP-1402

Boeing BSS 7239 (Rev.: A 1-18-88)

Toxic Gas Sampling and Analytical Procedures

Toxic Gas Generation

Gases produced for analysis are generated in a specified, calibrated smoke chamber during standard rate of smoke generation testing (ASTM E 662), in both flaming combustion and non-flaming pyrolytic decomposition test modes.

Carbon Monoxide (CO)

CO is monitored continuously during the 20 minute test using a non-dispersive infrared (NDIR) analyzer. Data are reported in ppm by volume at 1.5 and 4.0 minutes and at maximum concentrations.

Acid Gas Sampling

HCN, HF, HCl, HBr, NOx and SO2 are sampled by drawing 1 litre of the chamber atmosphere through two midget impingers, each containing 10 ml of 0.25N NaOH, at a rate of 400 ml per minute. The 2½ minute sampling period is commenced at the 4 minute mark. Determinations are performed in both the flaming and non-flaming modes and data are reported in parts per million (ppm) by volume in air.

Analysis of Impingers for Hydrogen Cyanide (HCN)

Cyanide in the NaOH impinger, as NaCN, is converted to CNCI by reaction with chloramine-T at pH greater than 8 without hydrolyzing to CNO⁻. After the reaction is complete, CNCI forms a red-blue colour on addition of a pyridine-barbituric acid reagent. Cyanide is quantified by spectrometric measurement of the increase in colour 578 nm.

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Reference: In-house SOP 02-13-SP-1402