NASA Contractor Report 165809

Development and Demonstration of Manufacturing Processes for Fabricating Graphite/LARC 160 Polyimide Structural Elements

R.K. Frost, J.S. Jones P.J. Dynes and D.H. Wykes

Rockwell International Downey, CA 90241

Contract NAS 1-15371 DECEMBER 1981

DEPARTMENT OF DEFENSE MASTICS TECHNICAL EVALUATION CREETER RENADCOM, DEVER, M. A. 67824

National Aeronautics and Space Administration

Langley Research Center Hampton, Virginia 23665 19960104 080

DTIC QUALITY INSPECTED 1

DISTRIBUTION STATEMENT A

Approved for public release; Distribution Unlimited



lestre edtre ad 67 100 10 1 --ENTER FORMAT DESIRED AND TERMINATE WITH END edsr e 7f with the base **(** ... end ***DTIC DOES NOT HAVE THIS ITEM*** (AD NUMBER: D434866 CORPORATE AUTHOR: ROCKWELL INTERNATIONAL DOWNEY CALIF* UNCLASSIFIED TITLE: DEVELOPMENT AND DEMONSTRATION OF (See MANUFACTURING PROCESSES FOR FABRICATING GRAPHITE/LARC-160 POLYIMIDE STRUCTURAL ELEMENTS, PERSONAL AUTHORS: FROST, R. K. ; DYNES, P. J. ; JONES, J. S. ; WYKES, D. H. REPORT DATE: DEC , 1981 PAGINATION: 498P 5 CONTRACT NUMBER: NAS1-15371 MONITOR ACRONYM: NASA MONITOR SERIES: CR-165809 REPORT CLASSIFICATION: UNCLASSIFIED LIMITATIONS (ALPHA): APPROVED FOR PUBLIC RELEASE; DISTRIBUTION UNLIMITED. AVAILABILITY: NATIONAL TECHNICAL INFORMATION SERVICE, SPRINGFIELD. VA. 22161. NASA-CR-165809. -33 - LIMITATION CODES: 1 24 -***** END OF DISPLAY LIST <<ENTER NEXT COMMAND>>

A Starting

The La Harling

Sec. 19. 19

		DTIC Unanr Justifi	TAB	
		By		
		Distrib	oution /	
		Ą	vailability Codes	
		Dist	Avail and/or	
	TABLE OF CONTENTS	DISC	Special	
		0-1		
	I I I I I I I I I I I I I I I I I I I		Page	
SUMMARY			1	
INTRODUC	TION		3	
PROCESS	DEVELOPMENT		7	
२ 1			7	
3 I I	Task (a) - QUALITY ASSURANCE		7	
3 1 2	Material Specification	•	7	
3 1 3	Process Specification		Ω .	
3 1 4	Nondestructive Inspection Techniques		8	
3 1 5	Chemical Characterization of LAPC-160		0	
3 1 5 1	LARC-160 Synthesis and Characterization Play	n	9	
3.1.5.2	Variables Study Materials		10	
3.1.5.3	Liquid Chromatography		11	
3.1.5.4	Ester Analysis		12	
3.1.5.5	Resin Intermediates		17	
3.1.5.6	Unreacted Amines		18	
3.1.5.7	Chemical Characterization Conclusions		19	
3.1.6	LARC-160 Variables Study		20	
3.1.6.1	Formulation and Process Variables		21	
3.1.6.2	Repeatability and Usability		22	
3.2	Task (b) - DEVELOP PROCESS		26	
3.2.1	Laminate Processing		26	
3.2.1.1	In Situ Cycle		26	
3.2.1.2	Two Stage Cycle		30	
3.2.1.3	Improved Two Stage Cycle		36	
3.2.2	"I" Stringer Processing		38	
3.2.2.1	"I" Stringer Tooling		39	
3.2.2.2	Layup and Debulking		39	
3.2.2.3	Vacuum Forming "I" Stringer Elements		39	
3.2.2.4	Imidizing "I" Stringer Elements		41	

1.0 2.0 3.0

TABLE OF CONTENTS (cont.)

3 2 2 5	Accombling of UTU Chainson Flowerts	Page
3.2.2.3	Assembling of "I" Stringer Elements	43
3.2.2.6	Cure Procedure	44
3.2.2.7	Postcure Procedure	45
3.2.3	Hat-Section Stringer Processing	45
3.2.3.2	Layup and Debulking	45
3.2.3.3	Imidizing (0) ₁₆ Cap Element	46
3.2.3.4	Shaping (O) ₁₆ Cap Element	46
3.2.3.5	Vacuum Forming Hat-Section Elements	46
3.2.3.6	Assembly for Cure	48
3.2.3.7	Cure Procedure	49
3.2.3.8	Postcure Procedure	49
3.2.4	Honeycomb Sandwich Processing	50
3.2.4.1	Prime Laminate Face Sheets	50
3.2.4.2	Prime Honeycomb Core	50
3.2.4.3	Assembly Sandwich Panel and Bond	51
3.2.5	Chopped Fiber Molding Processing	51
3.3	TASK (c) - FABRICATION AND TEST	53
3.3.1	Fabrication - Mechanical Properties Specimens	53
3.3.2	Testing - Mechanical Properties	54
3.3.2.1	Beam Test Description	54
3.3.2.2	Tensile Test Description	54
3.3.2.3	Compression Test Description	55
3.3.2.4	Flexural and Short Beam Shear Test Description	- 55
3.3.2.5	Tensile and Flexural Chopped Fiber Test Description	56
3.3.3	Test Results - Mechanical Properties	56
3.3.3.1	Tension	56
3.3.3.2	Compression	57
3.3.3.3	Flexural	58
3.3.3.4	Short Beam Shear	58
3.3.3.5	Tension and Flexural-Chopped Fiber Molding	59

TABLE OF CONTENTS (cont.)

Page

3.3.4	Fabrication - Structural Elements	59
3.3.4.1	Hat-Stringer Stiffened Skin Elements	60
3.3.4.2	"I" Stringer Stiffened Skin Elements	62
3.3.4.3	Honeycomb Sandwich Panel Elements	62
3.3.5	Prepration and Testing - Structural Elements	63
3.3.6	Test Results - Structural Elements	64
DEMONSTR	ATION COMPONENTS	67
4.1	Task (d) - LAMINATE FABRICATION	67
4.2	Task (e) - SKIN/STRINGER PANEL FABRICATION	67
4.3	Task (f) - HONEYCOMB PANEL FABRICATION	68
4.4	Task (g) - CHOPPED FIBER MOLDING FABRICATION	68
4.5	Task (h) - TECHNOLOGY DEMONSTRATOR SEGMENT	69
4.5.1	TDS Selection Rationale	70
4.5.2	TDS Front Spar Fabrication	71
4.5.2.1	Honeycomb Sandwich Panel Fabrication	72
4.5.2.2	"h' Frame Fabrication	72
4.5.2.3	"U" Closeout Ring Fabrication	72
4.5.2.4	Front Spar Panel Assembly	72
4.5.2.5	Front Spar Tee Fabrication	73
4.5.3	TDS Rear Spar Fabrication	73
4.5.4	Rib Modification for Load Introduction Plates	73
4.5.5	Alignment of Ribs to Covers	74
4.5.6	Assembly of the TDS	75
4.5.6.1	TDS Assembly - Stage 1	75
4.5.6.2	TDS Assembly - Stage 2	76
4.5.6.3	TDS Assembly - Stage 3	76
CONCLUSI	ONS	79
5.1	PROCESS DEVELOPMENT	79
5.2	DEMONSTRATION COMPONENTS	80

5.0

TABLE OF CONTENTS (cont.)

Page

6.0	REFERENCE	81
	Appendix A	343
	Appendix Al	344
	Appendix A2	347
	Appendix A3	349
	Appendix B	361
	Appendix B1	363
	Appendix B2	391
	Appendix B3	403
	Appendix C	413
	Tensile Coupon Curves	415
	Tension Beam Curves	445
14 1	Compression Beam Curves	455
	Appendix D	473
	Appendix E	479

FIGURE		PAGE
1	Idealized Polymerization Sequence for LARC-160 Polyimide Resin	83
2	Schematic Outline for the Synthesis of LARC-160 Polyimide Resin	84
3	Liquid Chromatographic Separation of LARC-160 Intermediate Ester Mixture	85
4	Liquid Chromatographic Separation of Synthetic Mixture of BTDA and NA Ester Compounds	86
5	Chemical Structures of Theoretical Isomer Products of BTDA Esterification	87
6	Chemical Structures of Theoretical Esterification Products of NA	88
7	Liquid Chromatographic Separation of LARC-160 Polyimide Resin	89
8	Relative BTA Concentration in LARC-160 Intermediate Ester Mixtures by HPLC Analysis	90
9	Relative BTA Concentration in LARC-160 Neat Resins and Prepregs by HPLC Analysis	91
10	Relative BTDE Monoethyl Ester Concentration in LARC-160 Intermediate Ester Mixtures by HPLC Analysis	92
11	Relative BTDE Monoethyl Ester Concentration in LARC-160 Neat Resins and Prepregs by HPLC Analysis	93
12	Relative BTDE Diethyl Ester Concentration in LARC-160 Intermediate Ester Mixtures by HPLC Analysis	94
13	Relative BTDE Diethyl Ester Concentration in LARC-160 Neat Resin and Prepregs by HPLC Analysis	95
14	Relative NE Monoethyl Ester in LARC-160 Intermediate Ester Mixtures by HPLC Analysis	96

FIGURE		PAGE
15	Relative NE Monoethyl Ester in LARC-160 Neat Resin and Prepregs by HPLC Analysis	97
16	Relative BTDE Triethyl Ester Concentration in LARC-160 Intermediate Ester Mixtures by HPLC Analysis	98
17	Chemical Structures of LARC-160 Resin Intermediates	99
18	Relative Methylene Dianiline Bis-Nadimide Resin Intermediate Concentration in LARC-160 Neat Resins and Prepregs	100
19	Liquid Chromatographic Separation of LARC-160 Polyimide "Variables Study", Batch 11, Extended Resin Cook Time Variation	101
20	Ion-Pair Liquid Chromatographic Separation of LARC-160 Polyimide Resin	102
21	Ion-Pair Liquid Chromatographic Separation of Jeffamine AP-22	103
22	Relative Methylene Dianiline Concentration in LARC-160 Neat Resins and Prepregs by HPLC Analysis	104
23	C-Scan Resin Variable No. 1	105
24	C-Scan Resin Variable No. 2	106
25	C-Scan Resin Variable No. 3	107
26	C-Scan Resin Variable No. 4	108
27	C-Scan Resin Variable No. 5	109
28	C-Scan Resin Variable No. 6	110
29	C-Scan Resin Variable No. 7	111
30	C-Scan Resin Variable No. 8	112
31	C-Scan Resin Variable No. 9	113

FIGURE		PAGE
32	C-Scan Resin Variable No. 10	114
33	C-Scan Resin Variable No. 11	115
34	C-Scan Resin Variable No. 12	116
35	C-Scan Resin Variable No. 13	117
36	C-Scan Resin Variable No. 14	118
37	C-Scan Resin Variable No. 15	119
38	C-Scan Resin Variable No. 16	120
39	Variation in Tg with Change in AP22 Concentration	121
40	TMA-Tg Values, Amine Variables	122
41	TMA-Tg Values NA/BTDA Variables	123
42	TMA-Tg Values Resin Processing Variables	124
43	TMA-Tg Values Anchamine & Tonox Amines	125
44	C-Scan of Laminate (Batch 23723) Imidizing Pressure	126
45	C-Scan of Laminate (Batch 23723) Imidizing Pressure 6.7 KN/m ² (2 in. Hg)	127
46	C-Scan of Laminate (Batch 23725) Imidizing Pressure 16.9 KN/m ² (5 in. Hg)	128
47	C-Scan of Laminate (Batch 23725) Imidizing Pressure 6.7 KN/m ² (2 in. Hg)	129
48	C-Scan of Laminate (Batch 23727) Imidizing Pressure 16.9 KN/m ² (5 in. Hg)	130
49	C-Scan of Laminate (Batch 23727) Imidizing Pressure 6.7 KN/m ² (2 in. Hg)	131
50	LARC-160 Preliminary Cure Cycle	132

FIGURE		PAGE
51	Flat Laminate Autoclave Tooling Concept	133
52	C-Scan Recordings of Panels EX41 and EX47 Minimum Cure Pressure Study	134
53	C-Scan Recordings of Panels EX48 and EX49 Minimum Cure Pressure Study	135
54	C-Scan Recordings of Panels, EX74, EX69, and EX70 Minimum Cure Temperature Study	136
55	C-Scan Recordings of Panels EX71 and EX72 Minimum Cure Temperature Study	137
56	TMA-Tg Characteristics of Specimen EX74	138
57	TMA-Tg Characteristics of Specimen EX69	139
58	TMA-Tg Characteristics of Specimen EX70	140
59	TMA-Tg Characteristics of Specimen EX71	14 1
60	TMA-Tg Characteristics of Specimen EX72	142
61	Typical Configuration for Debulking Celion/LARC-160 Prepreg	143
62	Typical Configuration for Imidizing Celion/LARC-160 Prepreg	144
63	LARC-160/Celion Imidizing Cycle Window and Sequence of Events	145
64	Typical Configuration for Curing Imidized Laminates	146
65	LARC-160/Celion Cure Cycle Window and Sequence of Events (Imidized Prepreg)	147
66	C-Scan of Laminate Imidized at 163 C, 60 Minutes	148
67	C-Scan of Laminate Imidized at 163 C, 90 Minutes	149
68	C-Scan of Laminate Imidized at 163 C, 120 Minutes	150

FIGURE		PAGE
69	C-Scan of Laminate Imidized at 163 C, 150 Minutes	151
70	C-Scan of Laminate Imidized at 163 C, 180 Minutes	152
71	C-Scan of Laminate Imidized at 177 C, 60 Minutes	153
72	C-Scan of Laminate Imidized at 177 C, 90 Minutes	154
73	C-Scan of Laminate Imidized at 177 C, 120 Minutes	155
74	C-Scan of Laminate Imidized at 177 C, 150 Minutes	156
75	C-Scan of Laminate Imidized at 177 C, 180 Minutes	157
76	C-Scan of Laminate Imidized at 191 C, 30 Minutes	158
77	C-Scan of Laminate Imidized at 191 C, 60 Minutes	159
78	C-Scan of Laminate Imidized at 191 C, 90 Minutes	160
79	C-Scan of Laminate Imidized at 191 C, 120 Minutes	161
80	C-Scan of Laminate Imidized at 191 C, 150 Minutes	162
81	C-Scan of Laminate Imidized at 199 C, 30 Minutes	163
82	C-Scan of Laminate Imidized at 199 C, 60 Minutes	164
83	C-Scan of Laminate Imidized at 199 C, 90 Minutes	165
84	C-Scan of Laminate Imidized at 199 C, 120 Minutes	166
85	C-Scan of Laminate Imidized at 199 C, 150 Minutes	167
86	C-Scan of Laminate Imidized at 218 C, 30 Minutes	168
87	C-Scan of Laminate Imidized at 218 C, 60 Minutes	169
88	C-Scan of Laminate Imidized at 218 C, 90 Minutes	170
89	C-Scan of Laminate Imidized at 218 C, 120 Minutes	171
90	C-Scan of Laminate Imidized at 218 C, 150 Minutes	172

xi

FIGURE		PAGE
91	Improved Two Stage Processing Cycle for Celion/LARC-160	173
92	Layup Sequence LARC-160/Celion "I" Beam - Balanced Cap	174
93	Fillet Radius Stock Tooling and Vacuum Bagging Arrange- ment - Imidizing Process	175
94	Layup and Tooling Process Hat-Stringer Assembly	176
95	LARC-160/Celion In-Situ Imidizing Cure Cycle Window and Sequence of Events	177
96	ASTM D647 Compression Mold for Celion/LARC-160 Molding Compound Flexure Specimen per ASTM D790	178
97	C-Scan Laminate EX 199 (0) _s for Aged Tensile Properties	179
98	C-Scan Laminate EX 200 (0, <u>+</u> 45,90) _s for Aged Tensile Properties	180
99	C-Scan Laminate EX 201 (90) ₄₀ for Aged Tensile and Compressive Properties	181
100	C-Scan Laminate EX202 (+45) $_{ m s}$ for Aged Tensile Properties	182
101	C-Scan Laminate EX 204 $(0)_{26}$ for Aged Short Beam Shear and Flexural Properties	183
102	C-Scan Laminate EX 220 (<u>+</u> 45) _S 32 Ply for Aged Compressive Properties	184
103	ASTM D647 Tensile Mold and Molded ASTM D651 Tension Coupon	185
104	Composite Tension and Compression Critical Beam Specimen Designs	186
105	Tension Coupon Specimen Designs	187
106	Compression Coupon Design and Test Set-up	188
107	Tensile Properties of LARC-160/Celion Laminates Postcured and 125 Hours -316C (600 F) Aged Conditions	189

FIGURE		PAGE
108	Compression Properties of LARC-160/Celion Laminates Postcured and 125 Hours -316 C (600 F) Aged Conditions	190
109	Flexural Strength of LARC-160/Celion 0 ⁰ Laminates	191
110	Short Beam Shear Strength of LARC-160/Celion Laminates	192
111	Typical C-Scans of "Hat" Elements	193
112	"Hat" Stringer Molded on Reverse Formed Tool - Showing Flat Condition	194
113	Influence of Tooling Mass on Cure Cycle and Heat Rise Rates - Flat Panels vs Hat Elements	195
114	Hat Stringers in Position on Skin - Fit-up Operation	196
115	Hat Stringer in Position on Skin with "1" Pressure Cauls Installed	197
116	Pressure Augmenter Plate in Position Over "1" Pressure Caul	198
117	"Hat" Stiffened Skin/Stringer-Bonding Complete	199
118	"Hat" Stiffened Skin/Stringer Showing Concave Skin Surface	200
119	C-Scan of Hat-to-Skin Bond, Typical Area - 1/2 Scale	201
120	C-Scan and Photomicrograph Correlation of "I" Stringer Cap Void Characteristics	202
121	"I" Stringers in Dry Fit Position on Skin Assembly	203
122	"I" Stringer in Bonding Position with "1" Pressure Cauls Installed	204
123	Pressure Augmenter Plate in Position Over "1" Pressure Cauls, "I" Stiffened Skin/Stringer Assembly	205
124	"I" Stringer Stiffened Skin Element in Vaccum Bag Bond- ing Fixture	206

FIGURES		PAGE
125	"I" Stiffened Skin/Stringer Panel Bonded Complete	207
126	C-Scan of 35X34 5 Harness Satin Weave Celion Fabric/ LARC-160 Laminate 0.33 cm Thick	208
127	"I" Stringer Stiffened Skin Panel Element EX111/EX113 Being Readied for -132 C (-270 F) Test	209
128	"I" Stiffened Skin/Stringer Panel, LN ₂ Manifolds and Baffle Plates in Place	210
129	Typical Test Set-up for -132 C (-270 F) Compression Element Test	211
130	Sandwich Element R.T. Test Set-up	212
131	Load/Strain Characteristics of "Hat" Stringer Stiffened Skin Element EX109/EX110 AT-132C (-270F)	213
132	Compression Load/Strain Characteristics of "Hat" Stringer Stiffened Skin Element EX195-4A Aged 125 Hours at 316 C (600 F) and Tested at -132 C (-270 F)	213
133	Compression Load/Strain Characteristics of Hat Stringer Stiffened Skin Element EX109/EX110A, Postcured Condition Tested at Room Temperature	215
134	Compression Load/Strain Characteristics of Hat Stringer Stiffened Skin Element EX109/EX110B, Postcured Condition Tested at Room Temperature	216
135	Load/Strain Characteristics of "Hat" Stringer Stiffened Skin Element EX195-2A Aged 125 Hours at 316 C (600 F) Tested at Room Temperature	217
136	Compression Load/Strain Characteristics of "Hat" Stringer Stiffened Skin Element EX195-1PC at 316 C (600 F)	218
137	Compression Load/Strain Characteristics of "Hat" Stringer Stiffened Skin Element EX195-3A Aged for 125 Hours at 316 C (600 F) and Tested at 316 C (600 F)	219
138	Load/Strain Characteristics of "I" - Stringer Stiffened Skin Element EX111/EX113 Tested at -132C (-270 [°] F)	220

FIGURES		PAGE
139	Compression Load/Strain Characteristics of "I" Stringer Stiffened Skin Element EX 194-4A Aged for 125 Hours at 316 C (600 F) and Tested at -132 C (-270 F)	221
140	Compression Load/Strain Characteristics of "I"-Stringer Stiffened Skin Element EX111/EX113, Postcured Condition Tested at Room Temperature	222
14 1	Compression Load/Strain Characteristics of "I" Stringer Stiffened Skin Element EX111/EX113, Postcured Condition, Tested at RT	223
142	Compression Load/Strain Characteristics of "I" Stringer Stiffened Skin Element EX194-2A, Aged for 125 Hours at 316 C (600 F), and Tested at Room Temperature	224
143	Load/Strain Characteristics of "I" Stringer Stiffened Skin Element EX194-1 Postcured Condition, Tested at 316 C (600 F)	225
144	Load/Strain Characteristics of "I" Stringer Stiffened Skin Element EX194-3A, Aged for 125 Hours at 316 C (600 F) and Tested at 316 C (600 F)	226
145	Load/Strain Characteristics of Sandwich Element EX241-1, Postcured Condition, Tested at -132 C (-270 F)	227
146	Load/Strain Characteristics of Sandwich Element EX241-3A Aged 125 Hours at 316 C (600 F), Tested at -132 C (-270 F)	228
147	Load/Strain Characteristics of Sandwich Panel Element EX150-1, Postcured Condition, Tested at Room Temperature	229
148	Load/Strain Characteristics of Sandwich Element EX241-2A Aged 125 Hours at 316 C (600 F) Tested at R.T.	230
149	Load/Strain Characteristics of Sandwich Element, EX150-2 Postcured Condition Tested at 316 C (600 F)	231
150	Load/Strain Characteristics of Sandwich Element EX241-4A Aged 125 Hours at 316 C (600 F), Tested at 316 C (600 F)	232
151	"Hat" Stringer Element EX109/EX110B Local Compression Failure, -132 ⁰ C (-270 ⁰ F) Test, Postcured	233

FIGURES		PAGE
152	"Hat" Element EX195-1PC Showing Skin Compression and Buckling Failures, 316 C (600 F) Test, Postcured	234
153	"Hat" Element EX195-1PC Showing Local Debonds and Flange Compression Modes, 316 [°] C (600 [°] F), Test, Postcured Condition	235
154	"Hat" Element EX195-3A Showing Local Flange Compression Modes, 316°C (600°F) Aged Condition	236
155	"Hat" Element EX195-3A, Local Skin Compression and Buckling Failure, 316 ^o C (600 ^o F), Test, Aged Condition	237
156	"I" Element EX194-1PC Showing Local Skin Compression Failure, 316 C (600 F) Test Postcured	238
157	Skin Compression Failure "I" Stringer Element EX194-4A -132 [°] C (-270 [°] F) Test, Aged Condition	239
158	Cap Compression Failure, "I" Element EX194-4A, -132 [°] C (-270 [°] F) Test, Aged Condition	240
159	Local Compressive Failure - Sandwich Element EX150-1, RT Test, Side 2, Postcured	24 1
160	Compressive Failure Mode Sandwich Element EX150-1, RT Test, Side 1, Postcured	242
161	Local Compressive Failure Sandwich Element EX150-2, 316°C (600°F) Test, Postcured	243
162A	Failure Modes of Sandwich Element EX 241-1, Postcured Condition, Tested at -132 C (-220 F)	244
1 62 B	Failure Modes of Sandwich Element EX 241-1, Postcured Condition, Tested at -132 C (-220 F)	245
163A	Failure Modes of Sandwich Element EX 241-3A Aged 125 Hours at 316 C (600 F), Tested at -132 C (-270 F)	246
163B	Failure Modes of Sandwich Element EX 241-3A Aged 125 Hours at 316 C (600 F). Tested at -132 C (-270F)	247

FIGURES		PAGE
1 6 4A	Failure Modes of Sandwich Element EX 241-2A, Aged 125 Hours at 316 C (600 F), Tested at RT	248
164B	Failure Modes of Sandwich Element EX 241-2A, Aged 125 Hours at 316 C (600 F), Tested at RT	249
165A	Failure Modes of Sandwich Element EX241-4A, Aged 125 Hours at 316 C (600 F) Tested at 316 C (600 F)	250
165B	Failure Modes of Sandwich Element EX 241-4A, Aged 125 Hours at 316 C (600 F) Tested at 316 C (600 F)	251
166	Comparison of Structural Efficiencies of Compression Panels	252
167	C-Scan Laminate CL 6C-11, (0,+45) _s , 6 Plies	253
168	C-Scan Laminate CL 12C-6, (0,+45) _s , 12 Plies	254
169	C-Scan Laminate CL 24C-7, (0 <u>+</u> 45) _s , 24 Plies	255
170	Three - 122 X 26.0 Cm Hat Stringer/Skin Panels Delivered to NASA-LaRC	256
171	End Views of Hat Stringer/Skin Panels Delivered to NASA- LaRC	257
172	C-Scans of Stringer to Skin Bond Joints, Panels EX249/ 248/245 and EX279/278/277	258
173	C-Scan of Stringer to Skin Bond Joints, Panel EX270/278/ 277	259
174	C-Scan Laminate CL8C-18, (0, <u>+</u> 45,90) _s 8 Plies	260
175	C-Scan Laminate Cl 12C-8 (0,90) _t , 12 Plies, Skin for Honeycomb Panels	261
176	C-Scan Laminate CL 12C-9, (0,90) _t , 12 Plies, Skin for Honeycomb Panels	262
177	C-Scan Honeycomb Panel 8A with Location of 25.4 X 25.4- cm Panels No. 1 and No. 2	263

FIGURE		PAGE
178	C-Scan Honeycomb Panel 9A with Location of 25.4 X 25.4-cm Panels No. 3 and No. 4	264
179	C-Scan Honeycomb Panel 9C with Location of 25.4 X 25.4-cm Panels No. 5 and No. 6	265
180	Molding Compound Process Demonstration Part Design	266
181	Celion/LARC 160 Molding Compound Complex Part Demonstration Mold	267
182	Complex Shaped Process Demonstration Part Showing Good Com- pound Flow and Mold Conformance-Imidized at 191 ^o C (375 ^o F), Removed From Mold Cold	268
183	Representative Demonstration Component	269
184	Completed Technology Demonstrator Segment Bonded Assembly	270
185	GR/PI Body Flap Concept and Demonstrator Segment	271
186	Bagging Assembly for Imidizing	272
187	Bagging Assembly for Autoclave Curing	273
188	Bagging Assembly for Honeycomb Sandwich Panel Bonding	274
189	C-Scan of Front Spar Sandwich Panel 18.5 x 13.5	275
190	Sketch of Three Section Tool for "h" Frame Member	276
191	Cured "h" Section Before Machining	277
192	"U" Closeout Ring Tool	278
193	Bonding Configuration for TDS Front Spar Panel	279
194	Elements of TDS Front Spar Panel	280
195	TDS Front Spar Panel Bonded Assembly	281
196	Typical Tool Configuration for TDS Front Spar Tee Members	282
197	Imidizing Configuration for Tee Member Element	283

FIGURE		PAGE	
198	Tee Member Bagging for Autoclave Cure	284	
199	Cured Tee Member Before Machining	285	
200	TDS Rear Spar Tool	286	
201	201 Load Introduction Pi Sections Bonded to the Machined TDS Rear Spar		
202	TDS Rib Areas Requiring Shim Bonding	288	
203	TDS in Rigged Condition	289	
204	Mechanical Fasteners Hold Ribs to Covers at Aft End of TDS	289	
205	Bagging Configuration for Bonding Ribs to Covers	290	
206	Honeycomb Inserted in TDS Cover Open Channel Closeout	291	
207	TDS With Breather Material in Place	292	
208	TDS Bagged for Bonding Cycle	293	
209	Harmonic Bond Testing	294	
210	Ultrasonic Pulse Echo Contact Testing	295	
211	Teflon Tubes Inserted in Aft Open Channel Closeouts	296	
212	Lower Leading Edge Cover With Breather Material in Place	297	
213	TDS With Lower Leading Edge Cover and Aft Spar Installed and all Breather Material in Place	298	
214	TDS Ready for Bagging	299	
215	TDS Bagged for Bonding Cycle	300	
216	TDS Bagged for Bonding Cycle	301	
217	TDS Upper Leading Edge Cover Before Machining	302	

FIGURE		PAGE
218	Partially Trimmed Upper Leading Edge Cover Clamped in Position on Bonded TDS Assembly	303
219	Final Trim of TDS Upper Leading Edge Cover	304

LIST OF TABLES

FABLE		PAGE
1	LARC-160 Intermediate Ester Batches	305
2	LARC-160 Neat Resin and Prepreg Batches	306
3	Experimental Conditions for HPLC Analysis of LARC-160 Resin	307
4	Experimental Conditions for Ion-Pair HPLC Analysis of LARC-160 Resin	307
5	Test Matrix Effect of Resin Formulation/Process Variables	308
6	Prepreg and Composite Physical, Short Beam Shear and Flexural Properties, LARC-160 Resin Stoichiometry and Process Variable Program	309
7	Minimum Molding Pressure and Temperature Investigations	312
8	Test Matrix - LARC-160 Imidizing and Cure Cycle Process Improvement Study	314
9	Imidizing and Cure Cycle Process Improvement Study Observations - Celion 3000/LARC-160 Composites	315
10	Test Matrix Mechanical Properties and Structural Elements	316
11	RT Bulk Compressive Properties of 352 Kg/m ² (22 PCF) 5052 Alloy 3.18 mm (0.125 in.) Cell Aluminum Honeycomb Core	317
12	Tensile Properties of LARC-160/Celion Unidirectional (0)5 Oriented Composite Postcured Condition - Test Beam	318
13	Tensile Properties of LARC-160/Celion Unidirectional (0)5 Oriented Composite, Aged 125 Hours at 316 C (600 F) - Beam Test	319
14	Compressive Properties of LARC-160/Celion Unidirectional (0)5 Oriented Composite Postcured Condition - Beam Test	320
15	Compressive Properties of LARC-160/Celion Unidirectional (0)5 Oriented Composite, Aged 125 Hours at 316 C (600 F) - Beam Test	321

LIST OF TABLES

TABLE		PAGE
16	Compressive Properties of LARC-160/Celion (0, +45, 90) _S Oriented Composite, Postcured Condition - Beam Test	322
17	Compressive Properties of LARC-160/Celion (0, +45, 90) _S Oriented Composite, Aged 125 Hours at 316 C (600 F) - Beam Test	323
18	Summary of LARC-160/Celion Tensile Properties	324
19	Tensile Properties of LARC-160/Celion (0)5 Oriented Postcured Composites (1) (2)	325
20	Tensile Properties of LARC-160/Celion (0) ₅ Oriented Composite, Aged 125 Hours at 316 C (600 F) (1) (2)	326
21	Tensile Properties of LARC-160/Celion (90) 40 Oriented Postcured Composites (1) (2)	327
22	Tensile Properties of LARC-160/Celion (90) 40 Oriented Composite Aged 125 Hours at 316 C (600 F) (1) (2)	328
23	Tensile Properties of LARC-160/Celion (<u>+</u> 45) _S Oriented Postcured Composites (1) (2)	329
24	Tensile Properties of LARC-160/Celion (+ 45) _S Oriented Composite, Aged 125 Hours at 316 C (600 F) (1) (2)	330
25	Tensile Properties of LARC-160/Celion (0, <u>+</u> 45, 90) _S Oriented Postcured Composite (1) (2)	331
26	Tensile Properties of LARC-160/Celion (, 145, 90) _S Oriented Composite, Aged 125 Hours at 316 C (600 F) (1) (2)	332
27	Summary of LARC-160/Celion Compression Properties	333
28	Compressive Properties of (90) ₄₀ and (<u>+</u> 45) _S Oriented Fiber LARC-160/Celion Composites	334
29	Flexural Properties of 0 ⁰ Oriented Fiber LARC-160/Celion Laminate	335
30	Short Beam Shear Properties of O ^O Oriented Fiber LARC-160/Celion Laminates	336

LIST OF TABLES

TABLE		PAGE
31	Tensile and Flexural Properties (Average) of Celion/LARC-160 Chopped Unidirectional Tape Molding Compounds	337
32	Structural Element Weight Losses After Aging at 316 C (600 F) for 125 Hours	338
33	Structural Element Potting Materials and Processes	339
34	Structural Element Target Loads	340
35	Results of Compression Tests on "Hat" and "I" Stiffened Skin and Sandwich Panel Structural Elements (1)	341
36	Psysical Properties of Celion/LARC-160 Composite Laminates Delivered to NASA-LaRC, Tasks (d) and (f)	342

1.0 SUMMARY

This final report describes the program effort performed by Rockwell International for the NASA/LaRC under Contract NAS1-15371. The objective of the program was to develop and demonstrate manufacturing technologies for the structural application of Celion graphite/LARC-160 polyimide composite material.

The program consisted of two parts: Process Development and Fabrication of Demonstration Components. Process development included establishing quality assurance of the basic composite material and processing, non-destructive inspection of fabricated components, developing processes for specific structural forms, and qualification of processes through mechanical testing. In the second part of the program, demonstration components were fabricated using the processes developed in part one. The demonstration components consisted of flat laminates, skin/stringer panels, honeycomb panels, chopped fiber compression moldings, and a Technology Demonstrator Segment (TDS) representative of the Space Shuttle aft body flap. The TDS, initially intended to be only a display article, was later directed in the program to a testable component. TDS test results will be reported in a separate final report.

2.0 INTRODUCTION

This program was conducted for the NASA Langley Research Center, Materials Division, Materials Application Branch under NASA contract NAS1-15371. Mr. Robert M. Baucom was the NASA Program Manager. Mr. Roger Frost of the Advanced Manufacturing Technology Department of Rockwell International, Downey, California was responsible for program management and technical direction. Acknowledgement is made for the technical assistance provided during the program by the following Rockwell personnel:

в.	D. Bhombal	C. L. Hamermesh	J. D. Leahy
I.	Bouton	P. J. Hodgetts	W. H. Morita
R.	J. Demonet	K. C. Hong	M. A. Morrow
Ρ.	J. Dynes	D. W. Houston	F. Roje
H.	E. Flanery	J. S. Jones	D. H. Wykes
s.	R, Graves	C. C. Kammerer	

The initial objective of this program was to develop and demonstrate manufacturing technologies for structural application of Celion graphite/ LARC-160 polyimide composite materials. Later this was expanded by contract modification to include mechanical ground testing of a Technology Demonstrator Segment (TDS), a three-bay test article representative of the Space Shuttle aft body flap.

This final report presents the accomplishments and results of the original contract requirements with regard to manufacturing technologies. Mechanical ground testing of the TDS will be documented in a subsequent report.

The manufacturing technologies phase of the program was divided into two parts, process development and the fabrication of demonstration components, each consisting of several tasks. The following briefly describes the objective.

Part 1. Process Development

Task (a) - Develop a quality assurance program including specification for Celion/LARC-160 polyimide materials, quality control of materials and processes, including studies of the effects of monomer and/or polymer variables and prepreg variables on the processibility of Celion/LARC-160 prepreg and on the mechanical properties of test specimens fabricated from the prepreg, and NDI of fabricated components.

Task (b) - Develop processes for fabricating laminates, hat and "I" stiffeners, honeycomb core panels, and chopped fiber moldings.

Task (c) - Fabricate specimens and conduct tests to qualify the processes for fabrication of demonstration components,

Part 2. Demonstration Components

4

Task (d) - Fabricate and NDI three (3) laminates 61x22-cm (24x48-in.) with 0° , $\pm 45^{\circ}$ lay up symmetrical about the neutral axis. Laminate thickness to be 0.08-cm, 0.15-cm, and 0.32-cm (0.030 in., 0.060 in., and 0.125 in.).

Task (e) - Fabricate and NDI three (3) secondarily bonded hat-stiffened skin-stringer panels 23-cm (9 in,) wide x 122-cm (48 in.) long with three lengthwise stiffeners.

Task (f) - Fabricate six (6) honeycomb core panels 25.4x24.5-cm (10x10in.) having 0.15-cm (0.060 in.) thick face sheets with 0°, 90° layup symmetrical about the neutral axis of the panel and 2.54-cm (1 in.) thick honeycomb core.

Task (g) - Fabricate six (6) chopped fiber moldings according to a specimen design mutually agreeable to Contractor and Contracting Officer's technical representative.

Task (h) - Fabricate a representative component of a Space Shuttle aft body flap that is mutually agreeable to the Contractor and Contracting Officer's technical representative.

Each of the above tasks is documented separately in the main body of this final report.

Use of commercial products or names of manufacturers in this report does not constitute official endorsement of such products or manufacturers, either expressed or implied, by the National Aeronautics and Space Administration.

3.0 PROCESS DEVELOPMENT

3.1 TASK (a) - QUALITY ASSURANCE

3.1.1 Selection of Prepreg Supply Source

On initiating the program, a survey of seven potential prepreg suppliers was conducted to determine their ability for providing Celion/LARC-160 prepreg materials to support the objectives of the contract. Each was evaluated by questionaire, Appendix A1. Following screening of returned questionaires, an on-site inspection of supplier prepreg facilities was made by a Rockwell survey team. The objectives of the survey were to determine the potential supplier's level of quality control procedures and record keeping, the type and adequacy of analytical test equipment, specific personnel capabilities, and their indicated cooperation and willingness to work with Rockwell to achieve reproducible prepreg material. Each was requested to provide samples of Celion/ LARC-160 prepreg tape for evaluation. Four of the seven suppliers responded to this request.

The prepreg samples were subjected to physical tests to determine percent of volatiles, resin solids content, fiber areal weight, and calculated thickness per ply. Visual examinations were performed to evaluate fiber collimation, gaps/edge waviness, tack and drape. Three suppliers, one primary and two backup, were selected to provide Celion/LARC-160 prepreg for the program.

3.1.2 Material Specification

All prepreg material used for this program was ordered against the requirements shown in Appendix A2 included as a flysheet attachment with each purchase order. Test conditions and calculations imposed by item 5 of the flysheet were defined by document LTR 2433-4462 which is presented in Appendix A3. In addition, prepreg acceptability for program production use was based on chemical

analysis of the intermediate ester and neat resin used to ensure prepreg batch-to-batch repeatability. Each prepreg batch was subjected to physical and mechanical testing before being used for program requirements.

From this program activity and also that conducted against NASA/LaRC contract NAS1-15183, Graphite/Polyimide Design and Fabrication, Rockwell material Specification MB0130-152, Graphite/ Polyimide Resin Prepreg - 600°F Applications was developed and is presented in Appendix B1. This specification includes both LARC-160 and PMR-15 resin impregnated graphite material systems and will be invoked when procuring these materials for production use.

3.1.3 Process Specification

Rockwell Process Specification MA0105-328, Fabrication of 600^oF Polyimide Graphite made from Norborene Terminated Methylene Di or Larger Analine Benzophenonetetracarboxylic Base Resin, is presented in Appendix B2. This specification presents the improved processing technique developed for both LARC-160 (Contract NAS1-15371) and PMR-15 (Contract NAS1-15183). While this specification describes processing for flat laminates, using perforated metal caul plates, the staging and curing procedures are directly applicable to complex structural shapes and were used in fabricating detail elements of the Technology Demonstrator Segment required by Task (h) of the program.

3.1.4 Nondestructive Inspection Techniques

Ultrasonic C-scan was the primary nondestructive inspection (NDI) technique utilized throughout this program to detect porosity, voids and debonds in laminates and bonded structures.

C-scan sensitivity is one of the most important variables that must be considered in the quality assessment of laminates and bonded structures. To optimize calibration sensitivity, comparative reference standards having internal defects of known type and size were fabricated and used for sensitivity settings. At the beginning of the NASA Graphite/Polyimide Design and Fabrication Contract (NAS1-15183), personnel at LaRC and Rockwell shared sample specimens of graphite/polyimide laminates having known defects. Ultrasonic C-scans and destructive correlative tests were made followed by the selection of an "A" sensitivity agreed to by both LaRC and Rockwell personnel. Therefore, "A" sensitivity has been used for ultrasonic C-scan inspection of products produced under this program with only minor exceptions as will be noted in appropriate sections of the test.

Solid laminate structures were C-scanned in the NDI laboratory by engineering personnel. Sandwich structures, however, were Cscanned in the production facility. The specification governing production C-scan procedure is presented in Appendix B3.

3.1.5 Chemical Characterization of LARC-160

The primary goal of this effort was to provide the methodology for specifying the chemical composition of LARC-160 polyimide resin materials. Once these analytical procedures were developed their sensitivity could then be assessed by analysis of a series of LARC-160 resin materials containing intentional composition and processing variations. This would thus provide the first step toward establishment of a chemical quality assurance specification for LARC-160.

One of the most successful techniques for characterizing composite matrix resins is high pressure liquid chromatography (HPLC). The objectives of the program were to describe the development of HPLC methods for characterizing LARC-160 resin and their application to the analysis of standard and Variables Study (discussed in 3.1.6) lots of commerical LARC-160 Resin materials.

3.1.5.1 LARC-160 Synthesis and Characterization Plan

A comprehensive characterization plan was devised for monitoring the chemical composition of LARC-160 at three stages during its manufacture. The monomeric LARC-160 ingredients and their idealized polymerization sequence are given in Figure 1. The three LARC-160 components are BTDE (diethyl ester of 3,3',

4,4' - benzophenonetetracarboxylic acid); the endcap NE (monoethyl ester of 5-norbornene-2,3-dicarboxylic acid); and Jeffamine AP-22, a liquid polymeric amine mixture containing approximately 85% methylene dianiline isomers with the remaining being mostly higher molecular weight tri-, tetra-and penta-functional amines. The commercial preparation of LARC-160 resin is outlined schematically in Figure 2. In the first step, BTDA and NA, the anhydride forms of BTDE and NE, respectively, are refluxed at 82-99°C (180-210°F) with alcohol for a total of approximately 60 minutes to give the BTDE/NE ester mixture. The alcohol used is an industrial mixture composed of 85.8% ethyl, 9.0% isopropyl, and 4.3% methyl alcohol with 0.9% methyl isobutyl ketone. The BTDE/NE ester mixture is a viscous liquid at room temperature containing several percent of unreacted alcohol. Neat resin is prepared in the next step by blending the liquid amine into the BTDE/NE ester mixture at 32°C (90°F). After mixing for five minutes, the material is stored in a freezer. Prepreg is produced in the final step by a proprietary hot-melt impregnation technique.

The chemical characterization plan called for HPLC analysis to be carried out subsequent to the BTDE/NE esters, neat resin, and prepreg stages of LARC-160 production. This approach was expected to help identify the introduction of chemical variations during manufacture and permit possible corrective measures to be taken during production.

3.1.5.2 Variables Study Materials

Many factors contribute to the consistent performance of an advanced composite matrix material. In the Variables Study, the most important LARC-160 formulation and processing variations were assessed by studying a series of modified resin materials prepared by the prepreg supplier. The formulation and process variations incorporated in the intermediate ester batches are described in Table 1. Several standard production batches from the same supplier which were analyzed are also identified. Only five of the intermediate ester Variables Study batches involved

modifications. Batches 7 through 10 contain \pm 5% variations in the standard amounts of BTDA and NA normally used. Batch 13 was given a six hour reflux compared to the normal 1.75 hour processing time.

The formulation and process variations contained in the neat resin and prepreg batches is given in Table 2. Batches 1 through 10 contain percentage variations on the formulated weights of amine and anhydrides normally used to prepreg LARC-160. An extended resin cook time of two hours at $79^{\circ}C$ ($175^{\circ}F$) was used in Batch 11, compared to a normal operation involving about 10 minutes at $60^{\circ}C$ ($140^{\circ}F$).

Two polymeric amine substitutes for Jeffamine AP-22 were evaluated in Batches 14 and 15. They are Anchamine DL, a product of Pacific Anchor Chemical Corp., and Tonox-22 from UniRoyal. Final Batch 16 represents a control material for the Variable Study, prepared with the standard LARC-160 formulation and under the same processing conditions as used for the other standard processed materials for Variables Study.

Resin Batches 1 and 2 in the Variables Study were rejected and replaced by Batches 1A and 2A, respectively. Original Batches 1 and 2 were rejected due to improper hot melt impregnation procedures. Although original Batches 1 and 2 of the Variables Study were not processed into prepreg, their chemical composition is included for comparative purposes. Included as a second part of the Variables Study was a test of the repeatability of the supplier in producing three standard batches of LARC-160.

3.1.5.3 Liquid Chromatography

The development of a liquid chromatographic separation technique for LARC-160 involved the selection of a type of column absorption material, an eluting solvent or solvent combination, and a mode of detecting the eluting components. The most common type of column packing material was chosen which is the so-called "reverse-phase" material consisting of octadecyl silane bonded 10µm silica spheres. Reverse-phase separations are nearly always carried out with an

aqueous/organic solvent gradient. The presence of both acidic BTDE and NE together with basic amine ingredients results in the pH being a critical factor for good resolution in the separation of LARC-160. Adequate column retention and resolution in a reverse-phase separation requires that component species be maintained in their nonionic forms by control of pH.

The ionization of carboxylic acid groups on BTDE and NE esters was suppressed by using a $pH = 3.00 \text{ KH}_2 PO_4$ buffer in the aqueous portion of the solvent gradient. This buffer serves not only to maintain BTDE and NE esters in their neutral forms but also enhances separation by a complex mechanism believed to be based on differences in the dissociation constants for various acidic groups present in BTDE and NE components. The basic amine components are present as protonated cations under the above acidic solvent conditions and are therefore poorly resolved.

A key feature of this technique was the adaption of low ultraviolet wavelength (200 NM) enabling the detection of the NE ester endcap. This in turn required the use of a solvent which was transparent at this wavelength. A water/acetonitrile solvent gradient was found to best fulfill all the requirements. Experimental details of the acidic reverse-phase techniques are given in Table 3.

The analysis of amine components in LARC-160 was made by an ion-pair HPLC method in which tetrabutyl ammonium phosphate is added to the aqueous solvent phase. The tetrabutyl ammonium cations are believed to ion-pair with anionic sample species, neutralizing their charge and improving retention and separation. This technique provides good separation of free amine ingredients in LARC-160, but BTDE and NE ester components are not as well resolved as with the buffered acidic solvent gradient. The experimental parameters used in the ion-pair method are given in Table 4.

3.1.5.4 Ester Analysis

BTDE/NE Ester Mixture

The initial step in LARC-160 synthesis is the co-esterification

of BTDA and NA. The products expected from this process are BTDE diester and NE monoester as shown in Figure 1. The HPLC separation of a BTDE/NE ester batch is shown in Figure 3. The complexity of the ester mixture is the result of several factors. First, the esterification is carried out using an alcohol mixture rather than pure ethyl alcohol. The major products detected are BTDE and NE ethyl esters; however, methyl esters are also present as BTDE monomethyl and BTDE methylethyl mixed esters. Isopropyl alcohol is apparently much less reactive in that none of its esters have been detected.

A second factor contributing to the complicated BTDE/NE ester composition is the manner in which the BTDE/NE ester mixture is prepared. In order to minimize the solvent content, only a small stoichmetric excess of alcohol is used for co-esterification. This results in a highly viscous reaction medium which tends to inhibit complete conversion of BTDA to BTDE diester and results in BTDE monoester and unreacted BTDA. When a large excess of alcohol is used in BTDA esterification, only diesters are produced. NA appears to be a more reactive anhydride than BTDA and is converted to monoester completely during esterification.

The identification of BTDE/NE ester components in Figure 3 was made by comparison with characteristic elution times of reference materials. A synthetic mixture of these BTDE and NE ester compounds was prepared by reacting the two anhydrides under conditions that favor the formation of a particular species. The HPLC separation of this mixture is shown in Figure 4. The isomeric makeup is that expected from the possible theoretical forms shown in Figures 5 and 6, except that three rather than four BTDE diester isomers are observed. The BTDE ortho diester isomer probably does not occur because of the much greater reactivity of the anhydride ring compared to the carboxylic acid groups toward esterification. Endo and exo isomer forms of NE esters were not separated in the present study. These reference ethyl ester compounds, together with others prepared from methyl alcohol and methyl/ethyl mixtures, made

possible the identification of the major BTDE/NE ester components.

BTDE/NE mixtures from the Variables Study, as well as several batches from larger scale standard production runs, were examined by HPLC. The relative concentration of each ester constituent was calculated as the peak area percentage of that component relative to the total area of all the peaks in the chromatogram. Although original Batches 1 and 2 of the Variables Study were not processed into prepreg, their chemical composition is included for comparative purposes.

Neat Resin and Prepreg

The characterization of BTDE and NE esters, ingredients in LARC-160 neat resin and prepreg, was made by the same HPLC method as that for analyzing BTDE/NE ester mixtures. The separation of a LARC-160 neat resin batch is shown in Figure 7. BTDE components and NE ester endcapper are separated, as are a number of resin intermediate compounds. The relative concentration of each BTDE and NE ester ingredient in the resin batches was expressed as its HPLC peak area percentage relative to the total ester peak area of the chromatogram.

The following is a discussion of the variation in the relative concentration of BTDE/NE ester components separated by HPLC as shown in Figure 3.

BTA

One form of the major anhydride ingredient which was detected in LARC-160 was BTA, the tetra-acid of BTDA. The identification of this component is complicated by the fact that if BTDA (anhydride) were present in LARC-160 resins, it would be hydrolyzed immediately to the tetra-acid form by the aqueous HPLC solvent gradient. It is therefore impossible to determine by HPLC whether the anhydride or tetra-acid were present in the resin. Infrared spectra of BTDE/NE ester mixtures, however, show neither of the strong anhydride absorption peaks at 1785 and 1860 cm⁻¹ characteristic of BTDA (anhydride). This implies that primarily only the tetra-acid occurs in

the LARC-160 materials. The levels of BTA in the intermediate ester and resin batches examined is shown in Figures 8 and 9. Except for Batches 15 and 16 of the Variables Study only a small amount of BTA is detected in the LARC-160 materials. One significant feature of these data is that the trend in BTA concentration among the samples is the same for intermediate ester, neat resin, and prepreg. This indicates that variations in BTA present at the intermediate ester stage of production are carried through both resin formulation and hot-melt prepregging.

The presence of BTA can affect later prepreg processing in several ways. First, BTA is more acidic than BTDA diester and is thus more likely to form insoluble salts with amine components added in the neat resin manufacturing operation. Salt formation has been reported (Ref. 1) in a similar PMR resin in which the tetra acid, 2,2-bis (3',4'-dicarboxylphen1) hexafluoropropane, and an aromatic amine were utilized. The presence of organic salts in PMR resin could introduce compositional variations and prevent proper prepregging. Furthermore, BTA may also be more reactive than BTDE toward amines in forming polyamide acid which can result in faster build-up of viscosity during processing. The high concentration of BTA in Batches 15 and 16 is believed to be the result of hydrolysis of BTDA by moisture in the air. These two batches were prepared late in the program using the original lot of BTDA which had been opened frequently, allowing exposure to moisture. It is not known for certain whether the excessive levels of BTA are responsible for the poor physical properties of laminates prepared from Batches 15 and 16 in the Variables Study. Replacement materials for these batches prepared with fresh BTDA showed only a small trace of BTA present.

BTDE Monoethyl Ester

Another unexpected form of the basic BTDA ingredient present is BTDE monoethyl ester. The peak area percentage of the second of the two isomeric forms (see Figure 5) is plotted in Figures 10 and 11. The concentration trend in this component is also very similar
between the intermediate ester, neat resin, and prepregs. The concentration variations are also similar to those observed for BTA. This is not unexpected because both BTA and BTDE monoester are the result of incomplete esterification and reaction conditions that promote BTA formation also favor BTDE monoester. This compound, which is also believed to be hydrolyzed, contains the ortho dicarboxylic acid group and could affect resin processing similar to that of BTA described above.

BTDE Diethyl Ester

The major BTA ingredient in the ester mixture is the diethyl ester. The variation in the second of three BTDE diethyl ester isomers separated by HPLC is plotted in Figures 12 and 13. The relative variations in Figure 12 and 13 correspond to different levels of BTA and BTDE monoester described earlier. For example, Batches 15, 16 and 16A contain a lower relative concentration of BTDE diethyl ester due to the greater amounts of BTA and BTDE monoester in these batches of material. The use of relative peak area analysis does not provide very meaningful data for the major component of a system. It works best for detecting relative variations in the concentration of minor ingredients.

NE Monoethyl Ester

The relative levels of NE ester endcap in the LARC-160 intermediate ester mixtures is plotted in Figure 14. In contrast to the variety of BTDA products present in the ester mixtures, the NE endcap occurs predominantly as the monoethyl ester. The relationship between NE ester peak area percentage and initial formulation changes is somewhat ambigious due to the larger variations in BTDA related ingredients. It is found, however, among Variables Study Batches 1 through 12, which are similar to BTDA related components, that correlations between NA formulation changes and peak area percentages can be made.

The relative levels of NE ester endcap in the LARC-160 resins are plotted in Figure 15. The irregularity in the data does not correlate with NA formulation changes and is believed to be related to resin cook variations. For example, the low level of NE ester in the extended resin cook time material (Batch 11) is a result of the preferential reaction of endcap with amine components to form resin intermediates. The greater reactivity of NE compared to BTDE esters have been discussed (Ref. 2 & 3). It is very difficult, therefore, to determine NE formulation errors by HPLC analysis.

BTDE Triethyl Ester

The extent of BTDE triethyl ester in the BTDE/NE ester batches examined is plotted in Figure 16. Excess ester reflux time in Variables Study batch 13 is indicated by an increased concentration of BTDE triester. The detrimental effects of the ortho ester group of this compound have been described for PMR-15 polyimide resin (Ref. 4). The amount of triester in the LARC-160 materials is, however, very small. BTDE triethyl esters could not be detected in LARC-160 neat resin or prepregs due to interference by ester/ amine reaction products which elute at the same time.

3.1.5.5 Resin Intermediates

Polymerization of LARC-160 monomers begins during neat resin preparation and prepregging, as evidenced by the appearance of new resin intermediate peaks in the HPLC separation of neat resin in Figure 7. According to the simplified curing sequence in Figure 1, BTDE ester ingredients first react with amines to form polyamide acids. These species are then endcapped by the NE ester to give the desired molecular weight product. Imidization and cross-linking then occur in subsequent steps at higher temperatures. Contrary to this reaction scheme, the resin intermediates identified in LARC-160 are predominantly NE ester/amine compounds.

The identification of three resin intermediate peaks in the HPLC separation of LARC-160 was made by comparison with elution times of model compounds whose structures are given in Figure 17. The variation in MDA bis-nadimide is plotted in Figure 18 for the LARC-160 resin materials. The effect of extended resin cook time and temperature, 2 hours at $79^{\circ}C$ ($175^{\circ}F$) vs 10 minutes at $60^{\circ}C$ ($140^{\circ}F$), is clearly indicated for Batch 11 of the Variables Study. The HPLC separation of this material is shown in Figure 19. A large increase in both MDA mono- and bis-nadimide results from excess resin cook time.

The presence of MDA bis-nadamide and bis-nadimide can affect both the processing and cured physical properties of LARC-160. Cross-linking of MDA bis-nadimide will result in a low molecular weight cross-link with different physical properties compared to the idealized structure shown in Figure 1. The remaining resin, however, will be deficient in endcapper, thus increasing its molecular weight between cross-links. The overall result is to produce a less homogeneous network than is predicted by the idealized polymerization sequence for LARC-160.

3.1.5.6 Unreacted Amines

The remaining ingredients to characterize in the LARC-160 resin system are the free or unreacted Jeffamine AP-22 amine compounds. An ion-pair HPLC technique was developed for this purpose. The separation of a standard production batch of LARC-160 resin by this method is shown in Figure 20. The identification of Jeffamine AP-22 components in the resin was made by comparison with the chromatogram of a pure Jeffamine AP-22 sample shown in Figure 21. Three major components are detected, including MDA, and two higher molecular weight MDA homologs. The relative MDA content was calculated as the peak area ratio of MDA to BTDE diester in the chromatograms. This ratio is plotted in Figure 22 for the LARC-160 batches. The results for Variables Study resin Batches 1-6 agree qualitatively with the Jeffamine AP-22 formulation changes incorporated in those materials. A low ratio is shown for Batch 11 because of its extended cooking time. The wide variation in the data for the standard batches is most likely due to the effect of B-staging rather than formulation changes.

3.1.5.7 Chemical Characterization Conclusions

This chemical characterization activity of the program demonstrates the applicability of HPLC techniques for characterizing the chemical composition of LARC-160 polyimide resin. Monomeric ester and amine ingredients, as well as a number of resin advancement products, are detected and identified by HPLC analysis. These techniques not only provide the basis for improved quality control procedures but also reveal information regarding the mechanism of LARC-160 polymerization. Some specific conclusions resulting from this study are:

- o Relative batch-to-batch chemical variations in BTDA in NA ingredients can be detected precisely, but absolute formulation deviations are difficult to determine because of the complexity of ester products and the effects of varying resin advancement composition.
- Resin B-staging or advancement caused by processing variations can be monitored from relative concentration of intermediate products formed.
- Complex conversion of BTDA to BTDE diester during esterification does not always occur, resulting in potentially detrimental BTDE monoesters and tetra acid (BTA) products.
- High levels of BTA can also result from hydrolysis of BTDA by exposure to air prior to formulation.
- NE ester endcap reacts preferentially with amine components, producing the bis-nadimide resin intermediate that modifies the ideal cured network structure.

- o Only a slight trace of BTDE triethyl ester is present in the resin materials examined.
- o. Free amine components present in neat resin and prepreg can be characterized by an ion-pair HPLC method.
- The chemical composition of neat resin and prepreg are quite similar, indicating that hot-melt impregnation has little affect on composition.

Based on the foregoing conclusions and observations made during the program, the following recommendations are given:

- Preparation of BTDE/NE intermediate esters should be modified to permit complete conversion of BTDA to BTDE diester.
- Pure methyl or ethyl alcohol should be considered for esterification. The use of a complex alcohol mixture introduces components whose behavior is unknown. The use of a pure alcohol would also greatly simplify the quality assurance of LARC-160.
- Care must be taken to prevent exposure of BTDA to moisture in the air which results in hydrolysis to the tetra-acid.
- The HPLC methodology needs to be made more reproducible.
 It is presently too difficult to obtain consistent data from one analysis to another.
- Improved quantification of the HPLC methodology is required.
 The use of relative peak area ratios is not satisfactory for analyzing all ingredients in LARC-160.

3.1.6 LARC-160 Variables Study

This portion of the program was incorporated by contract modification with the objective (1) of establishing the limits within which LARC-160 polyimide resin could vary with regard to formulation and processing without detriment to prepreg quality and (2) to demonstrate prepreg batch-to-batch repeatability.

3.1.6.1 Formulation and Process Variables

Table 5 presents the variables in formulation and processing assessed. All resin formulation and processing and prepreg production was conducted under laboratory conditions primarily to better control the variable requirements and also because of the 1.4 Kg (3 lb) prepreg batch requirement which would have been cost prohibitive if accomplished on a production line. The prepreg tape produced was 15.2-cm (6 in.) wide.

A total of sixteen prepreg batches were produced for this study. Batches 1 through 10 varied resin stoichiometry, batches 11 through 13 varied processing, and batches 14 and 15 substituted Anchamine-DL and Tonox 22 respectively for the Jeffamine AP-22. Batch 16 was utilized for chemical standardization since all other standard batches of prepreg used for this program were produced on a production line because of the quantity required.

Chemical analyses were conducted on each of the 16 batches of prepreg material produced (see 3.1.5). Analysis of the starting materials, NA, BTDA, and AP-22 was done only once since these materials were procured in large quantities. The intermediate ester, neat resin, and resin extracted from the prepreg of each batch were chemically analyzed as noted in Table 5.

Laminate panels 15.2x15.2-cm (6x6 in.), 14 ply unidirectional, were fabricated from each batch of prepreg material produced for the variable study using the two stage imidizing and cure process (see 3.2.1.2). The laminate panels were postcured free-standing for four hours at $316^{\circ}C$ ($600^{\circ}F$) in an air circulating oven.

Laminate Physical Properties

NDI C-scan tests resulted in 100% transmission through all resin stoichiometry and processing variable panels except EX207, resin variable number 5. Panel EX207 had a 10% excess concentration of Jeffamine AP22 and showed 40% transmission. The two panels with amine components, Anchamine-DL and Tonox 22, showed 70% and 0% transmission respectively. NDI C-scan recordings are shown in

Figures 23 through 38.

Target fiber volume of $60 \pm 2\%$ was achieved in five of fifteen panels fabricated. Low and high fiber volumes in the remaining panels is attributed to prepreg resin content inconsistencies within each laboratory scale tape roll. Detailed composite physical properties are presented in Table 6.

Laminate Mechanical Properties

Flexural and short beam shear (SBS) properties were determined on each postcured panel at room temperature and $316^{\circ}C$ ($600^{\circ}F$). High flexural strength and modulus and short beam shear strength was achieved in all specimens tested with the exception of panel EX219 which employed the Tonox 22. Detailed properties are presented in Table 6.

Laminate TMA-Tg Properties

Figure 39 shows a plot of Tg temperature determined for laminates as a function of Jeffamine AP-22 concentration. The NE and BTDE quantities were held constant at the standard formulation concentrations. Although the number of data points is limited, and only one determination per laminate sample was made, the plot indicates that Tg increases with increasing AP-22 concentration up to +10%. Data were too limited with respect to variable NE/BTDE concentrations to determine a trend in Tg temperature variations.

Significantly, panel EX 219 (Tonox 22) yielded a higher Tg than all other laminates. Individual TMA-Tg curves are presented in figures 40 through 43.

3.1.6.2 Repeatability and Usability

Three 4.5Kg (10 1b) batches (23723, 23725, and 23727) of 30.4cm (12 in.) wide unidirectional tape were produced under production conditions with separate batches of resin formulated for each 4.5 Kg (10 1b) of prepreg to demonstrate formulation repeatability and material usability over a six month period at $-18^{\circ}C$ ($0^{\circ}F$) storage and after ambient out-time exposure of seven days.

The resin batches were formulated within limits established from evaluating batches of prepreg 1 through 13 which varied in stoichiometry and processing. Formulation and processing limits for the three batches were as follows:

> AP-22 \pm 2.5% by weight NA \pm 2.5% by weight BTDA \pm 2.5% by weight Reflux time - not to exceed 90 minutes at 82 \pm 3^oC (180 \pm 5^oF)

> Cook time - not to exceed 115 minutes at $82 \pm 3^{\circ}C$ (180 + 5°F)

Prepreg material physical properties specified were:

Resin solids: $37 \pm 3\%$ Volatiles: $12 \pm 3\%$ Fiber Areal Weight: 134 ± 3 grams/m²

Samples of the neat resin and intermediate ester were provided with each batch of prepreg for HPLC analysis to determine repeatability and establish a standard for comparison of resin extracted from the prepreg during the storage period.

Although this phase of the variables study was to be completed in a six months period, equipment schedule priorities, equipment malfunctions, and unexplainable processing problems have not allowed a clear definition of usability after six months storage. Equipment priorities prevented laminates from being processed at the end of one month storage and laminates representing two months storage were not usable because of an autoclave malfunction. At the end of four month's storage, laminates were processed. However, these laminates were not deemed usable because of excessive resin bleed and fiber washing.

Chemical analysis of resin extracted from each of the three repeatability batches showed no marked difference to the samples of

neat resin supplied with each batch. A detailed examination of the prepreg supplier's records for the repeatability batches also showed no possible cause for the processing difficulties. Further chemical analysis by Rockwell did not disclose any extreme differences between the repeatability batches and other batches used successfully to date.

After eight months storage at $-10.8^{\circ}C$ ($0^{\circ}F$), two 15.2x15.2-cm (6.0x6.0 in.), 14 ply unidirectional laminates, laid up from each of the three Celion/LARC-160 repeatability batches were autoclave cured.

Prior to curing, the laminates were imidized: one from each prepreg batch at $190^{\circ}C$ ($375^{\circ}F$) and the remaining laminates at $207^{\circ}C$ ($405^{\circ}F$). Imidization staging at the noted temperatures was conducted under < 16.9KN/m² (5 in. Hg) vacuum bag pressure for one hour. The higher imidizing temperature was used, in an attempt to reduce resin flow during the cure process.

The laminates were autoclave cured for two hours at $316^{\circ}C$ $(600^{\circ}F)$ under full vacuum and augmenting pressure of $1379KN/m^2$ (200 psi). Total pressure was applied at the start of the cure and cure temperature was attained at the rate of $2.2^{\circ}C$ $(4^{\circ}F)/minute$. The cured laminates were forced cooled to $< 66^{\circ}C$ $(150^{\circ}F)$ before release of pressure. The cured laminates exhibited surface and edge fiber washing. However, this condition was to a lesser degree for those laminates imidized at the higher temperature. NDI C-scan, "A" sensitivity, of the laminates showed five of the six having minor voids to a maximum of 3% of laminate area.

Flexural properties of the laminates from one batch (23723) were $1476MN/m^2$ (214 ksi) at room temperature and $786MN/m^2$ (114 ksi) and $765MN/m^2$ (111 ksi) at $316^{\circ}C$ ($600^{\circ}F$) for laminates imidized at $190^{\circ}C$ ($375^{\circ}F$) and $207^{\circ}C$ ($405^{\circ}F$) compared to target values of > $1572MN/m^2$ (228 ksi) and > $938MN/m^2$ (136 ksi) respectively for the test temperature conditions. The remaining two batches (23725 and 23727) provided flexural strengths in the range of 1862 to $1924MN/m^2$ (270 to 279 ksi) at room temperature and 931 to $1282MN/m^2$

(135 to 186 ksi) at $316^{\circ}C$ ($600^{\circ}F$). Short beam shear values for the three batches were essentially equivalent: 110 to $122MN/m^2$ (16 to 17.7 ksi) at room temperature and 42 to 61 MN/m² (6 to 8.8 ksi) at $316^{\circ}C$ ($600^{\circ}F$) compared to target values of $> 103 MN/m^2$ (15 ksi) and $> 48 MN/m^2$ (7 ksi) respectively for the test temperature conditions.

Two sets of $15.2 \times 15.2 - \text{cm}$ (6x6 in.) 14 ply unidirectional laminates were fabricated from each of the three repeatability batches after the material had been stored for eleven months. The laminates were imidized at 218C (425F) for 30 minutes at two pressures 16.9 KN/m² (5 in. Hg) and 6.7 KN/m² (2 in. Hg). This change is processing, over that used for the prior specimens, was in keeping with the improved processing developed for the program.

Fiber washing was evident after imidizing to approximately the same extent as seen on prior laminates made from these material batches. However, it is felt this would have been controlled by the use of edge dams.

The laminates were autoclave cured for three hours at $329^{\circ}C$ (625°F) under full vacuum and augmenting pressure of 1379 KN/m² (200 psi). The laminates were not postcured. All laminates showed excellent C-scan quality as shown in Figures 44 through 49.

Mechanical properties of these laminates were not determined because of variable thickness caused by the fiber washing. However, specimens were taken from each laminate for determination of fiber volume and Tg. Results of these determinations are as follows:

Batch	Imidizing Pressure	Fiber Volume (%)	Tg
23723	16.9 KN/m ²	66	343 ⁰ C
	6.7 KN/m^2	63	349 ⁰ C
23725	16.9 KN/m^2	65	340 ⁰ C
	6.7 KN/m^2	64	341 ⁰ C

Batch	Imidizing Pressure	Fiber Volume (%) <u>Tg</u>
23727	16.9 KN/m ²	70	343 ⁰ C
	6.7 KN/m ²	67	339 ⁰ C
	Target	60 <u>+</u> 2	340 ⁰ C

Further effort on this particular study was impossible because of fabrication of the Technology Demonstrator Segment, Task (h), which was of higher priority. However, batch repeatability was demonstrated by the repetitive chemical analysis conducted in attempting to determine a possible cause for the "processing" problem.

With regard to usability after storage, experience with production materials used for this program has indicated no problem with materials that have been kept at $-10.8^{\circ}C$ ($0^{\circ}F$) for over six months.

3.2 TASK (b) - DEVELOP PROCESSES

Processing of Celion/LARC-160 was developed for fabrication of laminates, hat and "I" stiffeners, honeycomb sandwich panels, and chopped fiber moldings. Primary effort was devoted to developing process procedures for laminate structures which provided the basis for process development of other structure variations.

3.2.1 Laminate Processing

Three laminate processing procedures were evaluated during this program: (1) in situ cycle, imidizing and cure, (2) two stage cycle, imidizing and cure, and (3) improved two stage cycle, imidizing and cure.

3.2.1.1 In Situ Cycle

The objective of the in situ cycle was to accomplish imidization and cure in one continuous operation. The in situ cycle is shown schematically in Figure 50 and is described as follows:

- o Apply 6.7 KN/m² (2 in. Hg) vacuum and maintain throughout cycle.
- o Raise temperature from RT to $163^{\circ}C$ (325°F) at 1.7 to 2.8°C (3 to 5°F)/minute.
- o Hold for 1 hour at $163^{\circ}C$ (325°F).
- o Raise part temperature from $163^{\circ}C$ (325°F) to 329°C (625°F) at 1.7 to 2.8°C (3 to 5°F)/minute.
- o Apply 1378 KN/M^2 (200 psi) pressure at 274°C (525°F).
- o Cure at $329^{\circ}C$ ($625^{\circ}F$) for 2 hours.
- o Force cool to $< 149^{\circ}$ C (300°F) prior to pressure release.
- o Postcure free-standing in an air-circulating oven. Raise temperature from RT to $316^{\circ}C$ ($600^{\circ}F$) at $5^{\circ}C$ ($9^{\circ}F$)/ minute. Hold at $316^{\circ}C$ ($600^{\circ}F$) for 4 hours. Force cool to RT.

The tooling/layup configuration for the in situ cycle is shown in Figure 51. A limitation to this cycle was the inability to control resin flow resulting in higher than target fiber volume (60 + 2% laminates.

Concurrent with the development of the in situ cycle, studies were conducted to evaluate minimum pressure/minimum temperature cure conditions.

Minimum Pressure Cure Study

Celion/ LARC-160 unidirectional, 32 ply laminates 12.7-cm (5 in.) wide by 11.5-cm (4.50 in.) long were prepared at cure pressure levels of 1.38 MN/m^2 (200 psi), 1.03 MN/m^2 (150 psi), .69 MN/m^2 (100 psi) and .34 MN/m^2 (50 psi). The in situ cycle time/temperature profile and pressure application point of 274°C (525°F) was employed in autoclave molding the laminates and final cure was accomplished at 329°C (625°F) for two hours using standard tooling. Panels were postcured at 316°C (600°F) for four hours. As indicated by NDI C-scan, "A" sensitivity, tests and actual void volume measurements, .69 MN/m^2 (100 psi) was the lowest molding pressure that produced < 1% target void volume laminates. Panels

EX41 1.38 KN/m^2 (200 psi), EX47 1.03 KN/m^2 (150 psi) and EX48 .69 KN/m^2 (100 psi) NDI C-scan, "A" sensitivity, recordings showed near 100% sound laminates with only two discrepant areas <3.2-mm (1/8 in.) diameter showing on EX48. NDI C-scan recordings of postcured laminates are shown in Figures 52 and 53. Therefore, the minimum pressure limit for processing Celion/ LARC-160 prepreg appears to be somewhere between .69 MN/m^2 (100 psi) and .34 MN/m^2 (50 psi) since the 34 MN/m^2 (50 psi) pressure molded laminate EX49 showed total porosity in C-scan test and has a 6.4% void volume.

Significantly, panel EX 48 cured at .69 MN/m^2 (100 psi) had an optimum fiber volume of 60.2%, compared with panels cured at 1.38 MN/m^2 (200 psi) and 1.03 MN/m^2 (150 psi) which had fiber volumes of 69.9% and 67.7% respectively.

Composite physical, flexural and short beam shear properties of panels autoclave molded at 1.38 MN/m^2 (200 psi), 1.03 MN/m^2 (150 psi), .69 MN/m² (100 psi), and .34 MN/m² (50 psi) are compared in Table 7. Data indicated no definite trend related to flexural strength or elastic modulus properties in comparing cure pressure levels. Short beam shear strength, 316°C (600°F) properties, however, in all specimens cured at 1.03 MN/m^2 (150 psi), .69 MN/m² (100 psi), and .34 MN/m² (50 psi) fell slightly below the 48.2 MN/m^2 (7 psi) target level.

Minimum Cure Temperature Study

A study was conducted to determine the practical minimum temperature/time cycle for curing Celion/LARC-160 prepreg. This study was undertaken in an effort to take advantage of the possible use of reusable silicone rubber vacuum bags and/or premolded pressure caul sheets. These bags and/or caul sheets would be particularly advantageous in fabrication of hat and "I" beam stringers, sine wave ribs, and other complex shaped parts. Reusability is of prime importance for cost effectiveness. In addition, lower curing temperatures will reduce the chances of seal and bag leakage.

Fabrication of unidirectional, 32 ply 11.5x12.7-cm (4.5x5.0 in.) laminates was accomplished using the two stage Celion/LARC-160 prepreg cure cycle and tooling with 1.38 MN/m^2 (200 psi) pressure applied at 274°C (525°F). Cure cycles employed to bracket temperature/time limits are described as follows: 316°C (600°F) 2 hours; 302°C (575°F) 3 hours; 288°C (550°F) 4 hours; and 274°C (525°F) 5 hours. Postcure of all laminates was accomplished at 316°C (600°F for four hours.

NDI C-scan, "A" sensitivity, tests on laminates EX74, EX69, EX70, EX71, EX72 cured at respective temperatures of $329^{\circ}C$ ($625^{\circ}F$), $316^{\circ}C$ ($600^{\circ}F$), and $302^{\circ}C$ ($557^{\circ}F$) and $274^{\circ}C$ ($525^{\circ}F$) had 99.5% sound laminate areas after final postcure at $316^{\circ}C$ ($600^{\circ}F$) for four hours. C-scan recordings of postcured laminates are presented in Figures 54 and 55.

As cured laminate TMA-Tg values gradually decreased with decreasing cure temperature. For example laminate EX74 cured at $329^{\circ}C$ ($625^{\circ}F$) had a Tg of $346^{\circ}C$ ($655^{\circ}F$) and EX72 cured at $274^{\circ}C$ ($525^{\circ}F$) had a Tg of $257^{\circ}C$ ($495^{\circ}F$), All laminates, however, after being postcured at $316^{\circ}C$ ($600^{\circ}F$) for four hours had essentially equivalent Tg values, with panel EX72 having the highest value of $360^{\circ}C$ ($680^{\circ}F$). TMA-Tg recordings are presented for comparison in Figures 56 through 60.

Composite physical and mechanical properties are presented in Table 7. Composite fiber volume decreased with decreasing cure temperature, from 68.4% for panel EX74, cured at $329^{\circ}C$ ($625^{\circ}F$) to 62.1%, for panel EX72, cured at $274^{\circ}C$ ($525^{\circ}F$). No definitive trend was noted in normalized room temperature flexural strengths and none of the panels achieved minimum target values. All panels except EX70, cured at $302^{\circ}C$ ($575^{\circ}F$) achieved target $316^{\circ}C$ ($600^{\circ}F$) strength. Room temperature and $316^{\circ}C$ ($600^{\circ}F$) short beam shear target strengths were achieved on all panels.

3.2.1.2 Two Stage Cycle

The two stage cycle was developed to improve resin flow control not obtainable by the in situ process. The following fully describes the two stage cycle including layup and debulking procedures.

Prepreg Tape Layup Procedure

Stack prepreg tape in the required ply orientation and number of plies, paper backing surface up, on a smooth tooling surface such as a glass plate. Supporting layup materials such as parting films, bleeders, porous teflon coated 104 fiber glass (TX1040 or 3TLL), caul plate, breathers and vacuum bag are shown in the layup sequence in Figure 61.

During layup, it is preferable to slice the edges of the tape using a straight edge in order to remove irregularities. If the prepreg tape edges are cut clean and uniform, this operation can be omitted. Normally, the Celion/LARC-160 tape will have adequate tack to adhere to itself, however, if tack is not adequate, a hot iron may be employed to aid layup. This is accomplished by ironing directly over the paper backing; either locally tacking or flat ironing to adhere the plies. The iron heat setting should be in the range of $176^{\circ}C$ ($350^{\circ}F$). Great care shall be exercised in insuring either absolute butt joints or a slight overlap of 0.76-mm (0.03 in) in the layup of tape elements. Tow splices should be flagged on the tape roll by the supplier, however, extreme care shall be taken to visually inspect each tape element. Any sections having tow splices must be removed.

Debulking Procedure

Debulking of stacked prepreg has been found to be advantageous in fabricating both flat and complex shaped laminates for the following reasons:

 Preconsolidation debulks the prepreg close to final laminate thickness; therefore, when augmented pressure is applied during final cure, less resin and fiber movement is required to achieve ultimate thickness.

- During the debulking operation, supporting materials such as TX1040 and bleeders are adhered to the laminate stack, forming a well consolidated unit that is easily handled during subsequent operations.
- Prestacked and debulked laminates are easily stored, under refrigeration, in sealed bags.
- o The preconsolidated preforms, with bleeder materials inplace, are easily handled in vacuum forming operations during the fabrication of hat, "I", "Pi" and other complex shape elements.
- o Where dry prepreg is used in the layup, the debulking operation not only consolidates and adheres the materials but also rejuvenates the resin tack, making the total stack pliable.

Two approaches to the flat laminate debulking operations have been developed and successfully qualified for use in Tasks (a), (b), (c), (d), (e), and (f).

Dry Preg Debulking

The following operations are to be performed after completing prepreg tape layup.

- o Select a flat 4.6 to 6.34-mm (0.18 to 0.25 in.) thick aluminum caul plate the same size as the stacked layup and apply Frekote 33 mold release to the caul surface which will face the layup.
- o Preheat the caul plate to $127 + 6.6^{\circ}C$ (260 + $12^{\circ}F$).
- On removal from the oven (within 2 3 minutes), immediately assemble the caul plate to the stacked layup surface, seal in a vacuum bag and apply vacuum to 67 KN/m² (20 in. Hg).
 - Note: The layup area shall have been previously prepared for rapid vacuum bag application so that minimal heat loss is incurred from the hot caul

plate. This operation softens the LARC-160 resin and causes it to flow into the TX1040 and bleeder materials, creating a consolidated prepreg/bleeder preform. The dry prepreg stack will become mildly tacky and pliable after this operation.

 Allow the assembly to stabilize to room temperature before removing the vacuum bag.

Tacky Prepreg Debulking

The same procedures shall be employed as for dry prepreg except that heating of the caul plate is optional. If heat is used, the procedure described for dry prepreg shall be followed. The layup shall be debulked at room temperature under vacuum bag pressure, $< 67 \text{ KN/m}^2$ (20 in. Hg) for a minimum of four hours.

Imidizing Procedure

Principal concerns with the LARC-160 polyimide resin/graphite materials are ensuring (1) efficient, uniform removal of solvent and condensation reaction volatiles from large and complex surface areas, and (2) resin flow control in the composite prior to application of augmented pressure during the cure cycle.

Prepreg volatile removal techniques were developed using the tooling shown in Figure 62. The concept of uniform removal of volatiles is based on use of a perforated layup tool surface. Perforations in the surface act as individual, unrestrained vacuum ports serving local surface areas of about 6-cm² (1 in.²). Vacuum channel separator strips are located in-line between the vacuum ports to support the caul layup surface and to provide an unrestrained venting system to a central manifold that, in turn, leads to the main vacuum source.

The laminate preform is imidized on porous tooling shown in Figure 62. The laminate is contained in a Celgard 4500 or 4510 polypropylene microporous membrane which allows removal of volatiles through the bottom perforated caul plate while preventing resin loss into surrounding breathers. Volatiles are reduced to < 3% by this procedure. The imidizing cycle is shown in Figure 63.

A perforated top pressure caul without vacuum channels may be used rather than a perforated bottom plate, if desired, although this concept has not been proved for larger, thicker, laminates. If the top perforated caul is used, efficient methods for venting volatiles to the vacuum source must be employed.

Detailed Imidizing Procedure

- Apply one 7781 fiberglass breather ply and one layer
 of Celgard 4500 or 4510 to the surface of the perforated
 layup plate as shown in Figure 62. Celgard may be spliced
 with an electronic heat sealer or with a thin band of
 Kapton tape 6.35-mm (0.25 in.) wide.
- o Transfer the debulked preform and the integral bleeders from the layup tool to the prepared perforated layup plate.
- Assemble all bagging components per Figure 62 and install thermocouples in the trim edge of the part.
- o Install the assembly in an air-circulating oven and perform the imidizing cycle per Figure 63. Thermocouples placed within the part trim, not oven temperature, shall be used in controlling the imidizing cycle. Thermocouple data shall be autographically recorded.

Cure Procedure

Imidized flat laminates are autoclave cured on tooling shown in Figure 64. Since the laminate volatile content has been reduced to < 3% during the imidizing procedure it can now be treated analogous to an epoxy laminate in the cure process. Nonperforated cauls are employed with the bleeder arrangement shown in Figure 64. The autoclave cure cycle is shown in Figure 65.

Process development, performed for Task (b) and verified in Tasks (a), (b), (c), (d), (e), and (f), has shown that considerable latitude exists in cure levels. For example, ultimate cure temperature and pressure can be accomplished as low as $274^{\circ}C$ ($525^{\circ}F$) and 689 KN/m² (100 psi) respectively. Minimum cure temperature has been set at $287^{\circ}C$ ($550^{\circ}F$) and pressure 1378 KN/m² (200 psi).

Autoclave pressurization rates evaluated have been in the 5 to 7 minute range from 0 to 1378 KN/m^2 (0 to 200 psi). The pressure application window appears to be optimum in the temperature range of 274 to 287°C (525 to 550°F). Resin hot melt flow is in the range of 254 to 265°C (490 to 510°F), however, the apparent viscosity is very low. If pressure is initiated before 274°C (525°F) excessive resin losses have been realized, resulting in high fiber volume laminates.

Laminates are postcured at $316^{\circ}C$ ($600^{\circ}F$) for four hours in an air circulating oven in free-standing position. Postcuring studies during Task (b) proved that the composite Tg is significantly increased and mechanical properties at $316^{\circ}C$ ($600^{\circ}F$) are improved. Regardless of Tg or mechanical properties values attained after initial cure, postcured of all laminates has been specified to insure retention of laminate quality after exposure to the operating service temperatures.

Detailed Autoclave Curing Procedure

- Remove the Celgard membrane from the bottom surface of the laminate, being careful not to damage or disturb the imidized layup. Leave the 2.3 to 4.8-mm (0.09 to 0.18 in.) top pressure caul in place; do not remove bleeder.
- Transfer the laminate to the steel curing tool (nonperforated surface). The steel tool surface shall be prepared using Frekote 33 parting agent. A Kapton glide sheet shall be employed to separate the laminate from the tool surface. Tape all components in place with Kapton pres-

sure sensitive tape.

Install type 162 fiberglass breathers (or equivalent)
 over the aluminum pressure caul and onto the tool surface. Adequate breather material shall be placed be tween the part and vacuum source to insure efficient re moval of any residual volatiles.

o Install thermocouples into the edge of the part. These thermocouples shall be used in monitoring the recording time/temperature data during cure and shall be used to control the cure cycle.

 Install a 0.051-mm (0.002 in.) thick Kapton film bag over the breather and tool surface and seal around the periphery of the tool using a high-temperature sealant. Insure that no bridges exist or sharp protrusions bear against the vacuum bag. Make vacuum bag "ear" seals as required to insure adequate bag slack to prevent bridging.

o Install the steel clamping ring and secure with bolts around the periphery of the tool. The complete tooling arrangment is shown in Figure 64.

o Install the tooling in an autoclave and apply full vacuum to the tooling system. Apply 689KN/m² (100 psi) to the autoclave and inspect the system for leaks. If a vacuum leak greater than 33.6 KN/m² (10 in. Hg) occurs within five minutes, the source shall be located and repaired.

Perform the cure cycle within the time temperature profile of Figure 65. The heat up rate may be less than the 3.9°C (7°F)/min. as shown depending on autoclave capability and/or tooling mass. No cool down rate is specified as this also depends on autoclave/tooling characteristics. Record all events such as application of various levels of vacuum, pressure and autographically record temperature from each thermocouple on parts and

autoclave. The part thermocouples shall be used in controlling the cure cycle. All part temperatures shall be in the range of 274 to $287^{\circ}C$ (525 to $550^{\circ}F$) when 1378 KN/m² (200 psi) autoclave pressure is applied.

o Remove bleeder materials and clean up parts.

o Submit for NDI C-scan test.

Postcure Procedure

Postcure in an air-circulating oven by raising the oven and part temperature from room temperature to $316^{\circ}C$ ($600^{\circ}F$) at an average heat rise rate of 1.6 to $8.3^{\circ}C$ (3 to $15^{\circ}F$)/minute and hold at $316^{\circ}C$ ($600^{\circ}F$) for four hours.

3.2.1.3 Improved Two Stage Cycle

0

Process studies, initiated under NASA/LaRC Contract NAS1-15183, were further developed under this program to simplify the imidizing and autoclave cure cycles. Prepreg tape layup and debulking procedures are as defined in 3.2.1.2. The objective of this effort were as follows:

o Reduce the number of steps in the imidizing cycle.

Increase the prepreg preform imidizing temperature and/or time at temperature, thereby increasing the LARC-160 resin viscosity when hot melt occurs during the cure cycle. This would allow application of 1378 KN/m² (200 psi) pressure at the initiation of cure at room temperature and, in turn, eliminate the chance of error of pressure application in the temperature range of 274 to 287°C (525 to 550°F) with the existing cure cycle. Allowing the resin to seek its normal flow point while under constant pressure, also eliminates problems related to temperature non-uniformity due to varying part thickness or tooling mass.

Eliminate the intermediate 163[°]C (325[°]F) steps in the autoclave cure cycle by raising the temperature from

room temperature to final cure temperature within the established heat rise rate band.

Twenty-five panels, approximately $15.2 \times 15-2$ -cm (6x6 in.) were fabricated and imidized as noted in Table 8. All panels were autoclave cured under the same bag with 1378 KN/m² (200 psi) pressure applied at the start of the cure. Actual heat rise rate for the cure cycle averaged 1.3° C (2.3° F)/minute. Panels were submitted for NDI C-scan testing and specimens removed for physical properties tests.

The following observations were made on panels cured using the improved cycle:

High resin flow was noted on all panels imidized at 162 and 177°C (325 and 350°F) as indicated by saturation of surrounding fiberglass breather layers. Panels imidized at 191 and 199°C (375 and 390°C) showed excellent compaction characteristics and good resin beading at laminate edges. The panels imidized at 218°C (425°F) had minimal evidence of resin flow. The panels imidized for 30 and 60 minutes had overall excellent cosmetic appearance while those imidized for longer periods at 218°C (425°F) showed surface roughness discrepancies indicating inadequate resin flow.

NDI C-scan testing results showed all panels imidized at 162°C (325°F) for time periods of 60, 90, 120, 150, and 180 minutes had void area discrepancies ranging between 40 and 80%. Void areas increased with lesser time at imidizing temperature. Panels imidized at 177°C (350°F) showed ultra sound penetration improvement starting at 150 minutes with approximately 7% void area showing. Panels imidized at 191°C (375°F) starting at 60 minute thru 150 minutes showed 100% ultra sound transmission except for some cellulose acetate fiber splice voids. All panels imidized at 199 and 218°C (390 and 425°F) had 100% ultra sound transmission.

As previously noted, panels imidizing at 218°F (425°F) for time periods exceeding 60 minutes showed surface roughness irregularities due to resin flow reduction.

NDI C-scan recordings of each panel evaluated are presented in Figures 66 through 90.

Physical properties testing results verified imidizing time/temperature relation as observed in panel cosmetic appearance and NDI C-scan tests. All panels imidized at 163 and $177^{\circ}C$ (325 and $350^{\circ}F$) had high fiber and void volumes. Panels imidized at 199 and $202^{\circ}C$ (395 and $390^{\circ}F$) for time periods between 30 and 150 minutes achieved target fiber volume of $60 \pm 2\%$ and void volume $\leq 2\%$. Those panels imidized at $218^{\circ}C$ ($425^{\circ}F$) for 30 minutes also achieved target requirements. Panels imidized at $218^{\circ}C$ ($425^{\circ}F$) for longer time periods had low fiber volumes; void volumes were $\leq 2\%$. Detailed physical properties are presented in Table 9.

Results of this process improvement study indicated the best imidizing time/temperature bands to be in the range of $191^{\circ}C$ (375°F) for 60 to 150 minutes, $199^{\circ}C$ (390°F) for 30 to 150 minutes and $218^{\circ}C$ (425°F) for 30 to 60 minutes.

The improved two stage cycle, employed in fabricating elements for the Technology Demonstrator Segment (TDS), Task (h), is presented schematically in Figure 91. The cure cycle permits temperature options based on tooling requirements e.g. formed silicone rubber blankets utilized on complex contoured details. Postcure time at temperature varies from four hours at the lower temperature to two hours when cured at $329^{\circ}C$ ($625^{\circ}F$).

3.2.2 "I" Stringer Processing

0

The following process description is specific for the "I" stringer design developed for this program. It is, however, applicable to the fabrication of any "I" stringer element with minor modifications.

Stock for the individual components of the "I" stringer ("C" channels, caps and radius fillets) are laid up and debulked as flat laminates per 3.2.1.2 and then vacuum formed and imidized. Vacuum forming of "C" channel and bottom surface top cap flat pre-forms enables easy, wrinkle-free shaping of components.

Preformed imidized radius fillets, that fill the interstices between "C" channels and caps to near net shape, and cap elements are easily handled and located in position during assembly for autoclave cure.

3.2.2.1 "I" Stringer Tooling

The following tools were required for the fabrication of the "I" stringer designed for this program:

- "C" channel mandrels, 6061-T6 or equivalent aluminum alloy.
- Cap forming silicone rubber pressure caul stabilized on an aluminum bar.
- Imidizing tool consisting of a caul plate and perforated layup plate.
- o Preforming and imidizing tool for fillet stock.

Tools are shown in Figures 92 and 93.

3.2.2.2 Layup and Debulking

Layup flat stock for "C" channels per design requirements. Follow the layup and debulking procedures described in 3.2.1.2. During the layup procedure, the TX1040 and bleeder materials applied to "C" channels must be laid on a 45° bias to the rectangular flat preform. This is required to prevent wrinkles during preforming and imidizing operations.

3.2.2.3 Vacuum Forming "I" Stringer Elements

The "I" stringer elements; "C" channels, bottom surface top cap, and fillets are vacuum formed prior to imidizing.

Bottom Surface Top Cap

- Apply Frekote 33 parting agent to the aluminum "C" channel mandrels surfaces and oven dry for 15 minutes at 176°C (350°F).
- o Assemble two "C" channels together with attachment bolts.
- Cut a 3.5-cm (1.48 in.) wide, 96.5-cm (38.0 in.) long
 strip of debulked (0)₇ ply prepreg stock. Remove TX1040
 and bleeder ply.
- o Assemble the $(0^{\circ})_7$ layup to the "I" beam mold top cap surface, between two edge dams as shown in Figure 92.
- Place the silicone rubber surfaced aluminum pressure caul in place over the layup.
- o Apply a breather and nylon film vacuum bag, seal and place in an autoclave. Apply $> 84 \text{ KN/m}^2$ (>25 in. Hg) vacuum and 689 KN/m² (100 psi) pressure. Raise the temperature to 65°C (150°F) and hold for 15 minutes. Force cool to room temperature.
- o Remove the tooling from the bag.
- o Cut the preformed laminate along the center line of the cap cleavage to separate the two mandrels. The cap (0⁰)₇ preforms will adhere to each "C" mandrel after this operation. These remain in place for vacuum forming the "C" channels. Remove TX1040 from laminate.

"C" Channels

- Prepare a flat plate of suitable size for holding each
 "C" channel mandrel in a vacuum bag. Vacuum bag materials and seals shall be prepared previously so that a rapid seal can be made in the subsequent vacuum forming oper-ation.
- o Cut a 12.7-cm (5 in.) wide, 96.5-cm (38.0 in.) long strip of debulked (+ 45[°]), 4 ply prepreg stock. Remove

the TX1040 faying with the laminate surface facing the tool.

 Place the mandrel on the flat plate. Transfer the debulked laminate with integral 45[°] bias TX1040 and one ply 120 bleeder to the mandrel and secure in place on the ends with a small piece of tape.

- o Drape the vacuum bag over the flat-debulked laminate on the mandrel, seal and draw vacuum. This operation will form the laminate with TX1040 and bleeder in place, over the "C" channel, wrinkle-free.
- o Place the assembly in an air-circulating oven and raise the mandrel and part temperature to 65 to $93^{\circ}C$ (150 to $200^{\circ}F$) and hold for 15 minutes.
- Allow the assembly to return to room temperature before releasing vacuum. The laminate preform, with integral 45° bias bleeders and porous TX1040 separators, will remain secure to the mandrel and is now ready for imidizing operations.

3.2.2.4 Imidizing "I" Stringer Elements

Imidizing of all "I" stringer elements was accomplished as described in 3.2.1.2. The following defines preparations prior to imidization of formed elements, "C" channels and fillets.

"C" Channels

o Drape a layer of Celgard 4500 or 4510 microporous membrane over the preform bleeder surface and secure in place, wrinkle-free, with pressure sensitive tape on the backside of the mandrel. Celgard is used to contain the resin and release volatiles during imidization.

 Place the two "C" channel mandrels on a flat plate suitable for applying a vacuum bag.

 Install thermocouples under the breather material over the part, outside trim lines. Data from the thermo-

couples shall be autographically recorded and used as the basis for controlling the imidizing cycle.

 Drape one ply of type 120 or 7781 fiberglass or Mochburg paper breather material over the Celgard film surface.

o Install a nylon film vacuum bag, drape in place over the preform bleeder surface and seal around the periphery of the flat aluminum plate. Insure that an efficient breather system such as multi-plies of type 162 fiberglass breather are connected between the parts and vacuum source. The "C" channel imidizing arrangement is shown in Figure 92.

 Place the bagged assembly in an air circulating oven and imidize per Figure 63. Monitor and record thermocouple and other cycle events such as vacuum data.

Fillets

The fillet imidizing tooling and molding concept is shown in Figure 93. The tooling is designed to augment vacuum bag pressure through a pressure augmenter plate to a maximum of 710 KN/m^2 (103 psi) when 101 KN/m^2 (30 in. Hg) vacuum is applied. This feature produces well defined and consolidated imidized fillet preforms.

- o For each fillet preform, cut a 6.35-mm (0.25 in.) wide strip from the debulked fillet stock, remove TX1040 and bleeder, and place the strip in the fillet preform tool cavity.
- Install dams and pressure mandrels, thermocouples,
 pressure augmenter plate, breathers, vacuum bag, and
 seal per Figure 93.
- Place the assembly in an air circulating oven and imidize per Figure 63. Thermocouple data shall be used for controlling the imidizing cycle and shall be

recorded autographically.

3.2.2.5 Assembling of "I" Stringer Elements

- Remove Celgard, TX1040, and bleeder materials from outside surfaces of imidized "C" channel elements. Care shall be exercised to prevent damage to imidized preforms.
- Join the two "C" channel mandrels together with undersize diameter fasteners to allow for mandrel movement while under pressure during cure.
- Install two imidized Q⁰ fillet elements in cleavage, top and bottom, between the "C" channels and secure in place at part ends, outside the part trim area, with a small piece of Kapton tape.
- o Install tooling dams on top cap edges.
- Place the assembly on a flat steel tool for autoclave curing at 329°C (625°F), 1378 KN/m² (200 psi). The tool surface shall be prepared by coating with Frekote 33 parting agent. Cover tool surface with Kapton film glide sheet.
- Cut a 3.5-cm (1.48 in.) wide, 96.5-cm (38.0 in.) long strip of imidized 0°, 7 ply laminate stock and remove TX1040 and bleeder materials.
- o Place the laminate in the top cap recess over the $(\pm 45^{\circ})_{s}$, 4 ply flanges of the "C" channels and 0° fillet elements.
- Install the top pressure caul. The pressure caul shall
 be prepared by coating with Frekote 33 parting agent.
- Install type 162 fiberglass breathers (or equivalent)
 over the "I" beam tooling and onto the tool surface.
 Adequate breather material shall be placed between the
 part and vacuum source to insure efficient removal of
 volatiles and protection of the bag.

- Install thermocouples into the edge of the part. These thermocouples shall be used in monitoring and recording time/temperature data during cure and shall be used to control the cure cycle.
- o Install a 0.051-mm (0.002 in.) thick Kapton film bag over the breather and tool surface and seal around the periphery of the tool using a high-temperature sealant. Insure that no bridges exist or sharp protrusions bear against the vacuum bag. Make vacuum bag "ear" seals as required to insure adequate bag slack to prevent bridging.
- Install the steel clamping ring and secure with bolts around the periphery of the tool.

3.2.2.6 Cure Procedure

- Install the tooling in an autoclave and apply full vacuum to the tooling system. Apply 689 KN/m² (100 psi) to the autoclave and inspect the system for leaks. If a vacuum leak greater than 33.7 KN/m² (10 in. Hg) occurs within five minutes, the source shall be located and repaired.
- Perform the cure cycle within the time temperature profile of Figure 65. Record all events such as application of various levels of vacuum, pressure and autographically record temperature from each thermocouple on parts and autoclave. The part thermocouples shall be used in controlling the cure cycle. All part temperatures shall be in the range of 274 to $287^{\circ}C$ (525 to $550^{\circ}F$) when 1378 (KN/m²) (200 psi) autoclave pressure is applied.

 Remove bleeder materials, clean up parts, and submit to Quality Engineering for NDI C-scan test.

3.2.2.7 Postcure Procedure

Postcure the "I" stringer in an air-circulating oven by raising the oven and part temperature from room temperature to 316°C (600°F) at an average heat rise rate of 1.6 to 8.3°C (3 to 25°F)/minute and hold at temperature for four hours. The "I" stringer shall be supported on a flat base, free standing, during postcure.

3.2.3 Hat-Section Stringer Processing

Processing procedures are defined for the specific hatsection stringer designed for this program but are applicable to any hat-section element with minor modifications.

Stock for the individual components of the hat-section stringer are laid up and debulked as flat laminates per 3.2.1.2 except for modifications noted herein. Only the $(0^{\circ})_{16}$ unidirectional cap reinforcement is imidized in accordance with 3.2.1.2. Imidization of the web/flange components is accomplished in situ during the autoclave cure cycle.

3.2.3.2 Layup and Debulking

Prepreg tape having a nominal 0.145-mm (5.7 mil) cured ply thickness and 152 ± 4 grams/m² areal fiber weight was used for this hat-section stringer design.

Flat laminate stock, $(0^{\circ})_{16}$ was laid up and debulked per 3.2.1.2 for the hat-section cap.

Two flat laminates, $\pm 45^{\circ}$ two ply, for inner and outer web/ flange elements of the hat-section were laid up and debulked. However, the laminate stock for the inner element was debulked without bleeder material. In the layup of the laminate stock for both inner and outer elements, the TX1040 only and TX1040 with bleeder were applied to the laminate surface, as determined by the assembly, on a 45° bias to the rectangular laminate shape to prevent wrinkling during vacuum forming.

3.2.3.3 Imidizing (0[°])₁₆ Cap Element

The $(0^{\circ})_{16}$ cap stock was imidized per 3.2.1.2 except that 172 KN/m² (25 psi) was applied after the $115^{\circ}C$ (240°F) cycle to increase compaction and to reduce material movement during the cure process and thereby eliminated wrinkles in the cap area.

3.2.3.4 Shaping (0⁰)₁₆ Cap Element

Trim a 2.79-cm (1.1 in.) wide strip from the imidized $(0^{\circ})_{16}$ laminate stock parallel to the fibers using a sharp knife and straight edge. Remove TX1040 and bleeder.

Place the strip on the top of the mandrel and bevel the edges to match the angle of the tool using a sanding block.

3.2.3.5 Vacuum Forming Hat-Section Elements

o Apply Frekote 33 parting agent to the hat-section mandrel surfaces and oven dry for 15 minutes at $176^{\circ}C$ (350°F).

 Cut a 15.2-cm (6.0 in.) wide strip from the debulked two ply +45° flat laminate (inner element) to desired length for the inner layer of the hat-section.

- o Prepare a flat plate of suitable size for holding the mandrel in a vacuum bag. Vacuum bag materials and seals shall be prepared previously so that a rapid seal can be made in the subsequent vacuum forming operation.
 - o If the prepreg is dry and nontacky, heat the hatsection mandrel to $65 \pm 6^{\circ}C$ (150 $\pm 10^{\circ}F$) to promote improved drape for vacuum forming.
 - Place the mandrel on the prepared flat plate. Transfer
 the debulked flat laminate to the mandrel and secure
 in place with tape at each end.

o Drape a layer of nylon film and Mochburg paper breather over the surface of the flat laminate.

46

- o Drape the vacuum bag over the layup, seal and draw vacuum. Insure that the vacuum bag conforms to the radius areas by rubbing with a teflon paddle. This operation will form the flat debulked laminate with TX1040 in place over the mandrel, wrinkle-free. The laminate preform with integral 45° bias TX1040 separators will remain secured to mandrel.
- Remove nylon bag, Mochburg breather, and bias ply of TX1040.
- Place the shaped $(0^{\circ})_{16}$ cap element prepared over the <u>+</u> 45[°] two ply layup on the tool cap. Tack in-place on each end with a small piece of tape.
- Cut a 15.2-cm (6.0 in.) wide strip from the debulked two ply <u>+</u> 45^o flat laminate (outer element) to the desired length for the outer layer of the hat-section. Remove the single ply of TX1040. The bias oriented TX1040 and 120 fiberglass bleeder are to remain in place.
- o Place the laminate on the $(0^{\circ})_{16}$ cap element to have the graphite surfaces in contact. Secure each end to mandrel with tape.
- Repeat the above vacuum forming operation. In order to prevent the outer plies from tacking to the flange of the inner plies, insert a strip of polyethelene, or F.E.P. film between the two preforms along each flange. During the vacuum forming operation rub and force the bag into the radius areas with a teflon paddle.
- Remove the vacuum bag and then carefully remove the two parting film strips from between each flange of the "hat".

o Tack the two flanges into final position.

0

Drape a parting film such as mylon or F.E.P. over the vacuum formed hat assembly. Install the molded silicone

rubber caul over the parting film. Seal in a nylon film vacuum bag, place in an autoclave, apply vacuum and pressurize to 689 KN/m^2 (100 psi). Hold under pressure for approximately 15 minutes. This operation is performed to insure proper seating of prepreg preforms, bleeder materials and rubber caul.

• Remove bag, rubber tooling and parting film and inspect for part conformance to the tooling.

3.2.3.6 Assembly for Cure

- Apply Frekote 33 parting agent to the rubber caul. Air dry for 15 minutes minimum.
- o Install the rubber caul over the $\pm 45^{\circ}$ bias 120 fiberglass bleeder surface of the preformed hat.
- Place the mandrel on a flat steel tool suitable for curing parts at 1378 KN/m² (200 psi), 287^oC (550^oF). Install shims under the curved base of the tool to prevent bending the tool when autoclave pressure is applied.
- Install 162 fiberglass breather material over the external surface of the rubber caul. Apply material as required to prevent bridging and any sharp protrusions from coming in contact with the bag. Adequate breather material shall be placed between the part and vacuum source to insure efficient removal of volatiles.
- Install thermocouples into the edge of the part under the rubber caul. These thermocouples shall be used in monitoring and recording time/temperature data during cure and shall be used to control the cure cycle.
- Install a 0.051-mm (0.002 in.) thick Kapton film bag over the breather and tool surfaces and seal around the periphery of the tool using a high-temperature sealant. Insure that no bridges exist or sharp protrusions bear

against the vacuum bag. Make vacuum bag "ear" seals as required to insure adequate bag slack to prevent bridging.

 Install the steel clamping ring and secure with bolts around the periphery of the tool. The tooling arrangement is shown in Figure 94.

3.2.3.7 Cure Procedure

- Install the tooling in an autoclave and apply full vacuum. Apply 689 KN/m² (100 psi) to the autoclave and inspect the system for leaks. If a vacuum leak greater than 33.7 KN/m² (10 in. Hg) occurs within five minutes, the source shall be located and repaired.
- o Perform the in situ imidizing the cure cycle within the time temperature profile of Figure 95. Record all events such as application of various levels of vacuum, pressure and autographically record temperature from each thermocouple on parts and autoclave. The part thermocouples shall be used in controlling the cure cycle. All part temperatures shall be in the range of 274 to $287^{\circ}C$ (525 to $550^{\circ}F$) when 1378 KN/m^2 (200 psi) auto-clave pressure is applied. The ultimate cure temperature temperature shall not exceed $293^{\circ}C$ (560°F).
- o Force cool the part to $< 65^{\circ}$ C ($< 150^{\circ}$ F) prior to pressure release.
- o Remove the part from the tooling. Care shall be exercised to prevent tearing the rubber caul during removal from the surface of the bleeder material on the part.
- o Remove bleeder materials, clean up parts, and submit to Quality Engineering for NDI C-scan test.

3.2.3.8 Postcure Procedure

Postcure the hat-section in an air-circulating oven by raising the oven and part temperature from room temperature to $316^{\circ}C$ (600°F) at an average heat rise rate of 1.6 to $8.3^{\circ}C$ (3 to $15^{\circ}F$)/ minute and hold at temperature for four hours. The hat-section shall be supported on a flat base, free standing, during postcure.

3.2.4 <u>Honeycomb Sandwich Processing</u>

Laminate face sheets are processed in accordance with procedures defined in 3.2.1.3 except for postcuring which is accomplished after bonding the sandwich structure.

3.2.4.1 Prime Laminate Face Sheets

- Secure face sheets to a flat surface or in a holding frame.
- Abrade faying surfaces with Scotch Brite, Type A, pads and water. A water break free test will be performed and each skin force dried at 121°C (250°F) for 30 minutes in an air circulating oven.
- o Prime skin faying surfaces by spraying two box coats of 35% solids BR34 aluminum powder filled polyimide resin. (A box coat consists of two spray coats, with the second spray coat applied 90° to the first to ensure even coverage). The primer will be allowed to air dry for 45 minutes minimum, staged in an air circulating oven by raising the temperature from room temperature to 51.7°C (125°F) and then to 204°C (400°F) in 13.9°C (25°F) increments every 15 minutes. This procedure is required to prevent blistering the primer.
- Skins will be stored in clean kraft paper until ready for use.

3.2.4.2 Prime Honeycomb Core

- o Vapor and spray clean in trichlorethylene and oven dry at $121^{\circ}C$ (250°F) for 30 minutes.
- Apply 35% solids BR34 primer to the core cell faying edges in four box coats. Air dry 45 minutes minimum and stage primer in an air circulating oven per 3.2.4.1.

• Primed core elements will be stored in clean kraft paper until ready for use.

3.2.4.3 Assemble Sandwich Panel and Bond

- Apply adhesive film, FM34B-18, .44 Kg/m² (0.09 psf)
 to the face sheets, assemble with honeycomb, and vacuum
 bag for bonding.
- o Apply 84.4 to 94.5 KN/m^2 (25 to 27 in. Hg) vacuum and 2.76 KN/m^2 (40 psi) autoclave pressure.
- o Raise temperature from room temperature to $177^{\circ}C$ (350°F) at 2.8°C (5°F)/minute.
- o Cure two hours at $188^{\circ}C$ (370°F).
- Postcure sandwich panels free standingin an air-circulating oven by raising the temperature at 5-7°C (9-12°F)/minute to 316°C (600°F) and holding at 316°C (600°F) for four hours. Force cool panels to room temperature.

3.2.5 Chopped Fiber Molding Processing

Unidirectional prepreg tape 15.2-cm (6 in.) wide was procured for this process development to the following physical properties requirements:

- o Resin Solids: 38 ± 3%
 o Volatiles: 12 ± 3%
 o Fiber Areal 67 ± 3 grams/m²
 weight:
 o Calculated thickness, 60% fiber volume:
 - 0.0064-cm (2.5 mils)/ply

The tape material was chopped to produce random size pieces 1.25 to 2.54-cm (0.5 to 1.0 in.) long, in the filament direction, by 0.25 to 3.0-cm (0.1 to 1.2 in.) wide.
Development of chopped fiber molding processing was accomplished using an ASTM D790 flexure specimen molded in an ASTM D647 compression mold. The flexure mold and typical molded coupons are shown in Figure 96.

The following two stage process was developed for chopped fiber molding:

- Spread chopped unidirectional prepreg uniformly over a teflon sheet positioned in a shallow pan.
- o Imidize material in an air-circulating oven by raising the temperature from room temperature to $190^{\circ}C$ (375°F) and staging at temperature for one hour.
- Load mold with net weight imidized material to obtain a target fiber volume of 60%.
- o Place mold in a press preheated to $316^{\circ}C$ ($600^{\circ}F$) and close to contact position.
- o Apply 13790 KN/m^2 (2000 psi) pressure when part temperature reaches 204°C (400°F).
- o Cure one hour at $316^{\circ}C$ (600°F).

Postcure chopped fiber molding free standing for four hours at $316^{\circ}C$ (600°F).

3.3 TASK (c) - FABRICATION AND TEST

3.3.1 Fabrication - Mechanical Properties Specimens

Initial laminate panels to be used for mechanical properties testing were laid up and autoclave cured using the single stage in situ imidizing and cure process described in 3.2.1.1. These panels were used to obtain all postcured condition mechanical properties specified in the test matrix, Table 10. Resin flow control proved to be a problem using this cure cycle, sometimes resulting in high composite fiber volumes in the range of 64 to 68%. All laminates for specimen fabrication had essentially zero void content as determined analytically, and by NDI C-scan test.

Process optimization studies performed in Task (b) led to a two stage processing requiring an imidizing cycle where volatiles are removed to < 2% from the stacked prepreg prior to the autoclave cure. Resin flow control was maintained during imidization by low vacuum levels and a Celgard 4500 or 4510 microporous polypropylene film which allows volatile matter to escape through a perforated tooling plate while the membrane contains the low viscosity resin. Excess resin was absorbed into bleeder materials calculated to yield a laminate with a target 60 + 2\% fiber volume.

The autoclave cure was accomplished between two flat tooling plates. Since the major portion of volatile matter was removed in the imidizing cycle, the laminates were treated similarly to epoxy materials. Final laminate cure was accomplished at $287^{\circ}C$ ($500^{\circ}F$) for three hours or $326^{\circ}C$ ($625^{\circ}F$) for two hours.

This two stage process was employed in the fabrication of all laminates for mechanical properties specimens that were aged for 125 hours at $316^{\circ}C$ ($600^{\circ}F$). All panels had essentially zero void with fiber volumes in the 61 to 63% range. NDI C-scan recordings confirmed high quality and are shown in Figure 97 through 102. The detailed description of this two stage processing is presented in 3.2.1.2. Flexural and tensile specimens were molded from chopped fiber using the compression molds shown in Figure 96 and 103. The

molding process used is described in 3.2.5.

3.3.2 Testing - Mechanical Properties

Testing was performed in accordance with the matrix, Table 10. Three specimens for each test mode and temperature were tested in the postcured and aged, 125 hours 316°C (600°F), conditions.

3.3.2.1 Beam Test Description

Tension and compression critical beams were employed to determine (0)_t tension (F_{tu} , E_t , ε_{ult} , μ %) and (0[°])_t and (0[°], + 45[°], 90[°])_s compression (F_{cn} , E_c , $\varepsilon_{ult}\mu$ %) properties. The beam designs are presented in Figure 104.

Analytical studies (Ref. 5) performed by Mr. Mark Shuart, NASA-LaRC, proved that the 352 Kg/m^3 (22 pcf) honeycomb core significantly affects the measured strength and elastic modulus properties of composite specimens. A computer program was developed by NASA-LaRC to assess the actual effect this core has on the laminate properties and to establish property adjustment factors. Bulk core properties for both aluminum, 352 Kg/m^3 (22 pcf), and 301 CRES, 639 Kg/m^3 (40 pcf), core materials were developed by Rockwell International and transmitted to NASA-LaRC for use in the computer program in developing the composite property adjustment factors. These data are shown in Table 11. Specific adjustment factors are given in the individual mechanical property data Tables 12 through 17.

During test, individual specimens were stabilized at each test temperature for 10^{+10}_{-0} minutes prior to application of stress at a head travel of 1.27-mm (0.05 in.)/minute. Data were obtained by autographic recording of axial strain gages installed on the composite specimens at the beam midpoint.

3.3.2.2 Tensile Test Description

Tensile coupons were employed to determine $(0^{\circ})t$, $(90^{\circ})t$, $(+45^{\circ})s$ and $(0^{\circ}, \pm 45^{\circ})s$ and $(0^{\circ}, \pm 45^{\circ}, 90^{\circ})s$ tension properties. Specific properties determined were F_{tu} , E_t , ε_{ult} , $\mu(\pi)$ and ν .

The $(0^{\circ})_t$, $(90^{\circ})_t$ and $(+45^{\circ})_s$ coupons employed a straight sided design and the $(0^{\circ}, \pm 45^{\circ}, 90^{\circ})$ s coupons were necked down in the test section. Specimen design is shown in Figure 105.

During test, specimens were loaded at a head travel of 1.27-mm (0.05 in.)/minute after stabilizing for 10^{+10}_{-0} minutes at temperature. Data were obtained using biaxial strain gages mounted back-to-back on two of three specimens in each test group.

Load/strain data were obtained incrementally in testing postcured condition specimens by digital readout using a data logger. Stress/strain data plots were made using a Hewlett Packard 9820 computer system. The remaining single specimen in each group was instrumented with clip-on hang down extensometers.

In testing the 125 hour 316[°]C aged coupon specimens, load/ strain data was obtained autographically from biaxial strain gages at a constant head travel of 1.27-mm (0.05 in.)/minute.

3.3.2.3 Compression Test Description

Compression coupons were employed to determine (90)_t and (+45)_s compression properties, F_{cu} , E_c and $\varepsilon_{ult} \mu$ (%). The compression specimen is a 7.62-cm long x 2.54-cm wide (3.000X1.000 in.) coupon. Specific tolerances and test fixture design are presented in Figure 106. During test, specimens were loaded, after stabilizing at each test temperature for 10^{+10}_{-0} minutes, at a constant head travel of 1.27-mm (0.05 in.)/minute. Load/strain data were obtained autographically using a hang down deflectometer.

3.3.2.4 Flexural and Short Beam Shear Test Description

Specimens were machined from 0° , 26 ply, nominal 0.063-mm (2.5 mils)/ply, 1.65-mm (0.0565 in.) thick test panels. Specimen configurations were in accordance with ASTM D790 (flexural) and ASTM D2344 (short beam shear). Respective span to thickness ratios for each test are 32:1 and 4:1. Strain measurements were made auto-graphically during eact test using an isolated deflectometer positioned at the specimen midpoint. Elastic modulus properties were derived from load/strain curves obtained in the flexural tests

and the load/strain curves obtained in the SBS tests were used to give a positive indication of when actual specimen failure occurred. Specimens were loaded at a head travel of 1.27-mm (0.05 in.)/minute after being stabilized at the test temperature for 10^{+10}_{-0} minutes.

3.3.2.5 Tensile and Flexural Chopped Fiber Test Description

Specimen configuration as molded were in accordance with ASTM D651 (tensile) and ASTM D790 (flexural). The tensile and compression specimens were tested at room temperature and $316^{\circ}C$ ($600^{\circ}F$). For the $316^{\circ}C$ testing, the specimens were stabilized at temperatures for 10^{+5}_{-0} minutes.

3.3.3 Test Results - Mechanical Properties

3.3.3.1 Tension

Data obtained during testing are summarized in Table 18. The effects of test temperature and postcuring versus aging (125 hour, $316^{\circ}C$ ($600^{\circ}F$) conditioning) on composite mechanical properties are presented graphically in Figure 107.

Testing problems were experienced in some cases with beam specimens at $204^{\circ}C$ and $316^{\circ}C$ ($400^{\circ}F$ and $600^{\circ}F$) when either composite facing-to-core or steel facing-to-core bond failures occurred. Test data were tabulated at the composite stress level reached when bond failure occurred and are therefore not averaged.

Tension test results of $(0^{\circ})_{t}$ beam specimens adjusted per paragraph 3.3.2.1 show that the postcured specimens have higher strength than the aged under all conditions except room temperature. All tensile strengths were quite high starting at 2068 MN/m² (300 ksi) at -168°C (-270°F) and steadily decreasing to 1648 MN/m² (239 ksi) at 316°C (600°F). The $(0^{\circ}, \pm 45^{\circ}, 90^{\circ})_{s}$ and $(\pm 45^{\circ})_{s}$ tensile coupons in the postcured condition also maintained higher strength than the aged counterparts although the spread was very close. There was virtually no decrease in the postcured $(0^{\circ}, \pm 45^{\circ}, 90^{\circ})_{s}$ tensile strength between -168°C (-270°F) and 316°C (600°F). The resin critical $(90^{\circ})_{t}$ aged tensile coupon specimens

showed good strength retention in comparison with the postcured units, having slightly higher $-168^{\circ}C$ ($-270^{\circ}F$) and room temperature strengths and slightly lower $204^{\circ}C$ ($400^{\circ}F$) and $316^{\circ}C$ ($600^{\circ}F$) strengths.

Elastic modulus properties of the fiber critical $(0^{\circ})_{t}$ specimens were not significantly affected regardless of test temperature while the $(0^{\circ}, \pm 45^{\circ}, 90^{\circ})_{s}$, $316^{\circ}C (600^{\circ}F)$ test specimens showed some modulus loss. For the resin critical $(\pm 45^{\circ})_{s}$ and $(90^{\circ})_{t}$ coupons, a gradual decrease in elastic modulus properties was noted between $-168^{\circ}C (-270^{\circ}F)$ and $316^{\circ}C (600^{\circ}F)$.

Detailed tensile properties and failure modes from beam tests are presented in Tables 12 and 13 and properties from coupon testing are presented in Tables 19 through 26. Stress/strain curves from beam and coupon testing are presented in Appendix C.

3.3.3.2 Compression

Data obtained during test are summarized in Table 27. Effects of test temperature and postcuring versus aging (125 hours, $316^{\circ}C$ (600°F) conditioning) on composite mechanical properties are presented graphically in Figure 108.

Testing problems occurred in compression test of $204^{\circ}C$ ($400^{\circ}F$), (0° , <u>+</u> 45° , 90°)_s beams as discussed in the tensile results 3.3.3.1.

Analysis of compression test results indicated somewhat different trends in strength properties than found in tension, with lower ultimate strengths in the fiber critical orientations, $(0^{\circ})_t$ and $(0^{\circ}, \pm 45^{\circ}, 90^{\circ})_s$, and higher strengths in the resin critical orientations, $(90^{\circ})_t$ and $(+45^{\circ})_s$. For the $(0^{\circ})_t$ beam tests the aged, room temperature strength was higher than the postcured specimens as found in tension tests. A greater loss from room temperature compression strength was noted at $316^{\circ}C (600^{\circ}F)$ than in the tension tests, a 37% reduction for postcured and 50% reduction for aged condition. Beam specimens with $(0^{\circ}, \pm 45^{\circ}, 90^{\circ})_s$ fiber orientation, postcured condition, showed only 33% strength loss from room temperature to $316^{\circ}C (600^{\circ}F)$ while the aged spec-

imens had a 20% loss indicating a postcure effect. The resin dependent $(90^{\circ})_{t}$ and $(\pm 45^{\circ})_{s}$ coupon specimen strengths were almost identical in both postcured and aged conditions at each test temperature except for the postcured $316^{\circ}C$ $(600^{\circ}F)$ tested $(\pm 45^{\circ})_{s}$ specimens, indicating the influence of the resin. Strength losses from room temperature to $316^{\circ}C$ $(600^{\circ}F)$ ranged from 54% in the postcured $(\pm 45^{\circ})_{s}$ specimens, while the aged specimens lost only 20%, again indicating a postcure effect on the resin.

Elastic modulus properties of the $(\pm 45^{\circ})_{\rm s}$ specimens at each test temperature were increased after aging at 316° C $(600^{\circ}$ F), while the $(90^{\circ})_{\rm t}$ specimens showed no significant difference. $(0^{\circ})_{\rm t}$ and $(0^{\circ}, \pm 45^{\circ}, 90^{\circ})_{\rm s}$ specimens showed no significant change in elastic modulus properties, regardless of test temperature or aged condition. Detailed compression data and failure modes from beam tests are presented in Tables 14 through 17. Table 28 presents the data for coupon testing. Stress/strain curves are presented in Appendix C.

3.3.3.3 Flexural

Results of flexural strength tests on postcured and aged specimens show a drop in strength from room temperature to $316^{\circ}C$ $(600^{\circ}F)$ of 51% and 34% respectively. Specimens tested at $-168^{\circ}C$ $(-270^{\circ}F)$ yielded respective strength increases from room temperature of 18% and 6.1%. The aged specimens demonstrated higher strengths at all test temperatures except $-168^{\circ}C$ $(-270^{\circ}F)$. Elastic modulus properties were not significantly affected regardless of aged conditions or test temperature. Failure modes of $-168^{\circ}C$ $(-270^{\circ}F)$ and room temperature tested specimens were by outer fiber tension and by compression in specimens tested at $204^{\circ}C$ $(400^{\circ}F)$ and $316^{\circ}C$ $(600^{\circ}F)$. Detailed data are presented in Table 29 and the relative performance of postcured and aged specimens is shown graphically in Figure 109.

3.3.3.4 Short Beam Shear

Results of short beam shear tests on postcured and aged

specimens show exceptionally good strength retention at all test temperatures. The strengths of postcured specimens was slightly higher at $-168^{\circ}C$ ($-270^{\circ}F$) and room temperature than the aged specimens, equivalent at $204^{\circ}C$ ($400^{\circ}F$) and slightly lower at $316^{\circ}C$ ($600^{\circ}F$). All failure modes were by interlaminar shear. The relative performance of postcured and aged specimens is presented graphically in Figure 110 and tabulated data in Table 30.

3.3.3.5 Tension and Flexural-Chopped Fiber Molding

Four batches of chopped fiber molding material were evaluated. These varied in areal weight of the starting prepreg tape and chopped fiber length. The material batches and variations are noted below in the order of evaluation.

Batch A - Areal weight: 66.8 gm/m^2 Fiber length: 1.27 to 2.54-cm (0.5 to 1 in.)

Batch B - Areal weight: 153.7 gm/m^2 Fiber length: 1.27-cm (0.5 in.)

Batch C - Areal weight: 60.4 gm/m² Fiber length: 1.27-cm (0.5 in.)

Batch D - Areal weight: 67 gm/m² Fiber length: 2.54-cm (1 in.)

Flexural specimens were molded from all batches. Tension specimens were molded from batches A and B. Average properties are presented in Table 31.

3.3.4 Fabrication - Structural Elements

Hat-section and "I" stiffened panel design requirements, design assumptions for optimization of panel configuration, analysis, and supportive calculations for the designs are presented in Appendix D.

3.3.4.1 Hat-Stringer Stiffened Skin Elements

The detailed process description for fabricating hat-section elements is presented in 3.2.3. Difficulties were encountered during fabrication of some later 193-cm (76 in.) long hat elements in the form of $(\pm 45^{\circ})$ layers locally wrinkling along the upper cap corners. Wrinkling was caused by insufficient compaction of the 30.5X193-cm (12.0X76.0 in.) 16 plies thick unidirectional cap preform during the imidizing cycle. However, NDI C-scan test results showed that essentially void free parts were attained.

The wrinkling problem was resolved by modifying the unidirectional cap imidizing procedure by applying 84KN/m² (25 in. Hg) vacuum plus 172KN/m² (25 psi) autoclave pressure at the end of the 114°C (240°F) cycle. Resultant preforms were reduced in bulk thickness from 3.30 to 3.56-mm (0.13 to 0.14 in.) to 2.79 to 1.41-mm (0.11 to 0.095 in.) which decreased material movement during final compaction in the cure process. The Celgard contained the resin during pressurization and no excessive losses were noted. Additional debulking of the lay up was accomplished after final lay up using a molded silicone rubber caul at room temperature under 689 KN/m² (100 psi). The resultant preform on the aluminum mandrel closely matched the shape of the rubber caul, producing smooth, wrinkle-free surfaces. Hat elements were autoclave molded using the in situ cure process described in 3.2.1.1. Excellent NDI C-scan test results were obtained on all hats in the cured and postcured conditions, as typically shown in Figure 111.

Concave warpage of the hat stringers occurs along the element flange and inside cap length with a maximum flatness deviation of approximately 5.08-mm (0.20 in.) at the midpoint as shown in the photograph, Figure 112. This condition was partially removed after bonding the three hat elements (EX191, EX193, EX195) to skin (EX197) and was almost completely removed when the 193-cm (76 in.) long skin/stringer assembly was cut into five 30.48-cm (12 in.) long test sections, Specimen No's EX195-1, -2, -3, -4, -5. This lengthwise concave warped conditions is caused by imbalance in the hat design, where the major quantity of fiber contained in the cap section places the part's neutral axis off center. This condition would have posed a problem in the test of the stringer stiffened skin elements that were to be delivered to NASA-LaRC in Task (e) of the program, since nonuniform cap-skin loading would result from the concave condition.

To resolve this problem, the 127-cm (50 in.) long hat mandrel tool was modified by reverse rolling it concave to the cap surface, 6.35-mm (0.25 in.) at the midpoint. This modification, 1.27-mm (0.05 in.) more than actually observed in the elements as they are removed from the tool, was made on the assumption that upon removal from the reverse cured tool after curing, the hat element would approach a flat condition. Any minor longitudinal warping, concave or convex, would be eliminated when the hat is bonded to the skin.

This approach was verified in molding hat elements EX249 and EX250 to be used in fabricating hat-stiffened skin/stringer elements in Task (e). These parts were fabricated, per specific procedures defined in 3.2.3 on the reverse formed tooling. Resultant hat elements were flat and linear with no warpage. The photograph in Figure 112 shows the flatness of the hat element.

Due to the hat tooling mass, heat rise rates during the in situ imidizing cure cycle are extremely low. For example in the final critical temperature range between 257 to $271^{\circ}C$ (500 to $525^{\circ}F$) the average heat rise rate was only 0.51 to $0.55^{\circ}C$ (0.92 to $1.0^{\circ}F$)/minute. Figure 113 gives actual heat rise rate ranges observed during two hat element autoclave curing cycles. This indicates that the LARC-160 system is apparently not affected by long dwell periods close to the hot melt resin flow point, demonstrating that a large processing window exists. Heating rate comparisons are plotted in Figure 113 for typical flat panels, which show 340 minutes total cure time versus 590 minutes for hat elements, not counting cool-down.

The vacuum bag oven cure employing the pressure augmentation process was used to bond the three hat elements to skin using FM34B-18, 439 grams/m² (0.09 psf), adhesive. Tooling was improved for fabrication of the subsequent 193-cm (76 in.) long article, EX195, by employing inverted "T"-bars to distribute bonding pressure to hat flange/bond areas. This innovation improved handling and assembly of tooling elements. The tooling concept is shown in photographs, Figures 114 through 118. A section of the NDI C-scan recording of the hat-toskin bond is shown in Figure 119.

3.3.4.2 "I" Stringer Stiffened Skin Elements

The detailed process description for fabricating "I" section elements is presented in 3.2.2. NDI C-scan tests showed considerable void areas in the caps. To determine the void characteristics, 120X photomicrographs were taken of discreet cap areas where both 100% and 0% sound penetrations were recorded. From these it was determined that the void shapes were irregular micron sized pits distributed throughout the cap thickness. The cap side showing 100% sound penetration showed no voids in the photomicrographs. Actual respective void volumes detemined analytically were 8.31% and 1.13% and fiber volumes 58-62%. NDI C-scans and photomicrographs of the "I"-stringer cap are shown in Figure 120.

"I"-stringers were bonded to the skin assembly with FM34B-18 adhesive film on a 104 glass cloth carrier using the vacuum bag pressure augmentation process. Since the maximum pressure augmentation area-to-bond area available in this part design is only 1.5:1, an autoclave was required in the bonding operation. A minimum 3:1 pressure augmentation area to bond is required for one atmosphere oven bonding operations. The bonding sequence is shown in Figures 121 through 125.

3.3.4.3 Honeycomb Sandwich Panel Elements

Fabrication of sandwich panel EX150, 63.5x71.1x4.85-cm (25.0x28.0x1.91 in.) was accomplished in accordance with processes described in 3.2.4. The skins were comprised of unbalanced (0⁰₂,

 \pm 45°, 0°), 5 ply, nominal 0.144-mm (5.7 mil)/ply unidirectional tape. NDI C-scan recordings indicated a good skin to core bond was attained.

Celion IK, 34x35, 5 harness satin weave graphite fabric/ LARC-160, 20 ply doubler stock was fabricated using the two stage processing, imidization and autoclave curing. Panel size was 30.5x60.9x0.33-cm (12x24x0.130 in.). NDI C-scan showed 100% transmission; a recording is shown in Figure 126. Doubler stock was machined to a tapered configuration and bonded to the ends of each sandwich element with FM34B-18 adhesive film.

3.3.5 Preparation and Testing - Structural Elements

All structural elements were tested on a 1957KN (440,000 lb.) capacity Tinius Olsen universal testing machine. Test preparation and testing was accomplished as follows:

Specimen ends were ground flat and parallel to within
+ 0.127-mm (+ 0.005 in.).

One half of all structural elements were aged 125 hours at 316°C (600°F). Initial weights, and percent weight loss after aging are shown in Table 32.

Hat-and "I"-stiffened skin/stringer panels were stabilized at each end by potting in place to approximately 1.27-mm (0.50 in.) thick to match equivalent thickness precision ground tool steel load plates. Potting materials were selected based on test temperature and processed as shown in Table 33. Sandwich panel ends were stabilized by the tapered doublers described in paragraph 3.3.4.3. Doublers were clamped between parallel bars resting on the test bed during test to prevent specimen ends from spreading.

 Bi-axial gages were positioned and bonded at the midpoint of the center stringer cap and skin, in a backto-back pattern and in rosette, 0°, 45°, 90°, on one webof the center stringer. Sandwich panels were instrumented

with back-to-back strain gages in rosette, 0° , 45° , 90° , at the midpoint.

 Hat and "I"-stiffened skin/stringer panel edges were clamped to provide fixity; sandwich panel edges were not clamped.

Specimens were positioned in the test machine on a special spherical seat fixture designed to ensure optimum axial alignment. A pre-load of 2244 to 8896 N (500 to 2000 lb.) was applied and the specimen was aligned by adjusting the spherical seat to match back-to-back skin and hat cap axial strain gage deflections to a tolerance of 50µ. Typical specimens are shown in position on the test machine in Figures 127 through 130.

 After stabilizing at test temperature, a compressive load was applied incrementally up to the individual specimen calculated design ultimate. Strain measurements were taken at each loading increment. Specific target ultimate loads are shown in Table 34.

3.3.6 Test Results - Structural Elements

Data are summarized in Table 35 and load/strain curves are presented in Figures 131 through 150. Figures 151 through 165 show failures modes of elements that failed during test.

All of the room temperature elements met the design ultimate load requirement of 525KN/m (3000 lb./in). Specimen EX109/EX110A hat stringer failed while being held at the predicted ultimate load, showing good correlation between theory and design practice. A small degree of strength degradation was noted in the hat stiffened skin/stringer element EX109/EX110B during $-168^{\circ}C$ ($-270^{\circ}F$) testing where failure occurred at 117 KN (26,250 lb.), 3.4% below room temperature design ultimate. This specimen had previously been tested to design ultimate of 120.8 KN (27,150 lb.) at room temperature and then fatigue tested to 265,000 cycles, 5% to 67% of design ultimate (compression/compression). Premature failure may have been caused

by the combination of previous static and fatigue testing and resin embrittlement at -168 °C (-270 °F).

The strain gage data showed, for the most part, linear compression properties for the section designs tested. This was most apparent in the tests on the two honeycomb panels, which represent a balanced section and the six "I"-stringer panels which are unbalanced. The unbalanced hat-stiffened skin/stringer elements showed fairly linear strain increase until just before failure where it was found that probable local instabilities caused fairly large excursions in the gage readings in some test cases. The nonlinearity would be aggravated if the section designs were more unbalanced.

In terms of structural efficiency, the hat-stiffened panel yielded the lightest weight design. It should also be noted that the $(0^{\circ}, \pm 45^{\circ}, 90^{\circ})_{s}$ skin configuration of the hat-and "I"-stiffened panels was dictated as a design requirement, and thus these sections did not represent optimal designs. A measure of the structural efficiency of these configurations may be obtained by plotting the design parameters as shown in Figure 166. The relative weights per unit area of the three configurations are as follows:

Hat-Stiffened panel:	4.5Kgm/m ²	(0.939	1b/ft ²)
"I"-stiffended panel:	5.1Kgm/m ²	(1.04	1b/ft ²)
Honeycomb panel:	5.6Kgm/m ²	(1.14	1b/ft ²)

4.0 DEMONSTRATION COMPONENTS

4.1 TASK (d) - LAMINATE FABRICATION

Three laminates, 64x127-cm (25x50 in.), were fabricated using 0.127-mm (5 mil), 30.5-cm (12 inch) wide prepreg tape. Laminate ply orientation was (0^o, \pm 45^o) symmetrical about the neutral axis. The laminates, indentified as CL6C-11, CL12C-6, and CL24C-7, were 0.75, 1.5, and 3.0-mm (0.030, 0.060, and 0.120 in.) thick having 6, 12, and 24 plies of prepreg tape respectively.

Two stage processing, imidization and cure, of the laminates was accomplished as described in 2.3.1.2. After post curing for four hours at 316°C (600°F), the laminate panels were C-scanned and trimmed to finished dimensions, 60x120-cm (24x48 in.). C-scans are shown in Figures 167, 168, and 169. Laminate physical properties are presented in Table 36.

TASK (e) - SKIN/STRINGER PANEL FABRICATION

The secondarily bonded hat-section stringer configuration, rather than the "I" stringer was selected on the basis of the test data (presented in 3.3.6) which showed that both met load requirements, with the hat configuration having a weight savings advantage.

Prepreg tape, 0.145-mm (5.7 mil) thick and 30.5-cm (12 in.) wide, was used for fabricating the laminate skins and hat-section elements for the three demonstration articles required by this task. Laminate and hat-section processing is described in 3.2.1.2 and 3.2.3. Assembly bonding of the three hat-section to the laminate skin was accomplished as described in 3.3.4.

The completed skin/stringer panels, measuring 26x122-cm (10.2 x48 in.) are shown in Figure 170. An end view of one panel is presented in Figure 171. NDI C-scan recordings of the FM34B-18 adhesive stringer to skin bond areas are shown in Figures 172 and 173. C-scans typical of the laminate skins and hat-sections are shown in Figures 174 and 111.

4.2

TASK (f) - HONEYCOMB PANEL FABRICATION

Six honeycomb sandwich panels 25.4x24.4-cm (10x10 in.) were required by this task. These panels consisted of 0.15-cm (0.060 in.) thick, (0^o, 90^o)_t face sheets bonded to 2.54-cm (1 in.) thick glass/polyimide (HRH 327) honeycomb cure having a .48-cm (3/16 in.) cell size and 64.04 Kg/m³ (4.0 1b/ft³) density. The face sheets were bonded to BR34 primed core with FM34 adhesive, 0.44 Kg/m² (0.09 psf). Face sheet to core bonding was accomplished as defined in 3.2.4.

Two 12-ply laminates, designated as CL12C-8 and CL12C-9, 61x91-cm (24x36 in.) were laid up using 0.127-mm (5 mil) prepreg tape. The layup was such that the $(0^{\circ}, 90^{\circ})_{t}$ fiber orientation was symmetrical about the neutral axis of the honeycomb core. Laminate cure was as defined in 3.2.2. Postcure of the laminates was accomplished in two stages: two hours at $316^{\circ}C$ ($600^{\circ}F$) free standing and two hours at $316^{\circ}C$ ($600^{\circ}F$) in the bonded condition which also postcured the adhesive bond line. Physical properties of the postcured laminates are shown in Table 35. Specimens for physical properties determination were taken from laminate trim areas and received the full four hour exposure.

C-scans of laminates CL12C-8 and CL12C-9, showing the location of the three face sheets 28x56-cm (llx22-in.) cut from each are presented in Figures 175 and 176. Face sheet pairs 8C9C, 8A8B, and 9A9B were bonded to honeycomb core. C-scans of the three sandwich panels, identified as 8A, 9A, and 9C, and showing the location of each 25.4x25.4-cm (l0x10 in.) panel, designated #1 through #6, are shown in Figures 177, 178, and 179.

4.4 TASK (g) - CHOPPED FIBER MOLDING FABRICATION

Six chopped fiber molding were fabricated using the longer fiber length/lower areal fiber weight molding material (Batch D) noted in 3.3.3.5. The moldings were made using the NASA/BAC matched metal molds utilized in performing a similar task for Contract NAS1-15009 (Graphite/PMR-15 Composite Materials). The molding procedure is

68

4.3

is defined in 3.2.5. Part configuration is shown in Figure 180. Figures 181 and 182 show the charged matched metal mold and a finished part.

4.5

TASK (h) - TECHNOLOGY DEMONSTRATOR SEGMENT

The initial intent of this task was to fabricate a representtive demonstration component of the Space Shuttle aft body flap. It was intended that the component design would incorporate all processes and structural configurations developed in task (a) through (g) to demonstrate manufacturing feasibility of graphite/ LARC-160 to full-scale structures. An example of such a demonstration component, presented in the program proposal, is shown in Figure 183.

The requirement of this task "to fabricate a representative demonstration component" was later changed "to fabricate a representative structural test component". This structural test component has been given the designation of Technology Demonstrator Segment (TDS). The completed TSD, ready for installation of instrumentation for ground testing, is shown in Figure 184.

In changing from a demonstration to a structural test component, the complexity of the task changed correspondingly. All aspects of design, tooling, NDI, fabrication, and assembly became more critical. To implement this change, the scope of Contract NAS1-15843 (Develop, Demonstrate, and Verify Large Area Composite Structural Bond with Polyimide Adhesive) was amended to fabricate the cover panels, ribs, and leading edge covers of the TDS. Fabrication of the remaining TDS components and final assembly operation was accomplished under this program, Contract NAS1-15371.

All fabricated elements of the TDS, i.e., solid laminate structures, laminate skins, and bonded honeycomb panels, were nondestructively inspected and the results recorded and filed. NDI records for the TDS are not included in this report because of

volume considerations and also because many of the recordings, being very large, would lose definition when reduced for report inclusion.

4.5.1 TDS Selection Rationale

Increase in orbiter inert weight during maturation of the design adversely affects the deliverable and recoverable payload weight capability. The decreased deliverable payload could be restored with additional Shuttle system propulsion, but the decreased recoverable weight cannot be restored in this manner. However, reduction of the basic orbiter weight could result in the restoration of both the deliverable and recoverable payload weight. Significant weight savings are predicted for application of advanced composites in orbiter structural components.

Early Shuttle orbiter studies showed that the use of advanced composites on the vertical tail, elevon, and aft body flap would achieve significant weight savings, particularly if high-temperature graphite/polyimide (Gr/PI) were employed. In 1976, the NASA selected the orbiter body flap as a demonstration component for the Composites for Advanced Space Transportation Systems (CASTS) program. Since that time, orbiter composite-structures IR&D studies have emphasized the body flap. In FY 1976, a preliminary design concept for a body flap was identified. Adhesive bonding of joints was used throughout, thus eliminating stress-concentration and fatigue problems associated with mechanical fasteners. In FY 1977 through 1980, the design data base for Gr/PI structure was expanded through an extensive test program of body flap related subelements.

The body flap was chosen because it is a large, relatively simple, and easily retrofittable structure. It is subjected to extreme acoustic (165dB OASPL), aerodynamic (13.8 KN/m²), and thermal (1482°C) environment. The flight environment would thus thoroughly test the advanced structural concepts and demonstrate feasibility of application to other Orbiter structures.

Structural weight reduction and increased performance can be realized by taking advantage of the large strength-to-weight and stiffness-to-weight ratios of advanced composites. Savings of up to 145Kg (320 lb) of the total body flap structure/TPS weight can be realized by application of 316°C (600°F) structural allowable Gr/PI. In comparison to the baseline aluminum structure (177°C structural allowable), Gr/PI has reduced TPS requirements; and the TPS tiles can be directly bonded to the Gr/PI substructure because of the thermal compatability and stiffness of the components.

The design of the TDS simulates a section of the orbiter aft body flap incorporating three ribs and extending from the forward cover panels to the rear spar as shown in Figure 185. This section is 137x152-cm (54x60 in), 43-cm (17 in) at the front spar, and 18-cm (7 in) at the rear spar.

Specific objectives of the TDS were as follows:

- o Verify advanced composite design/analysis techniques.
- Develop and verify manufacturing techniques for large, complex Gr/PI structure.
- Demonstrate the integrity of Gr/PI all bonded structure to sustain orbiter aerodynamic, thermodynamic, and acoustic environments.

4.5.2 TDS Front Spar Fabrication

The TDS front spar panel (Appendix E, SS79-00253) is a bonded structure consisting of a honeycomb sandwich panel having a 13.33-cm (5.25 in) diameter access opening edged with a "U" shaped ring laminate, solid laminate "h" shapes framing the honeycomb panel, and laminate doublers backing the "h" section.

4.5.2.1 Honeycomb Sandwich Panel Fabrication

Laminate fabrication and sandwich adhesive bonding processes were performed as described in 3.2.1.2 and 3.2.4 except for minor modifications in the laminate bagging assembly procedures. Bagging assembly for imidizing and curing of the 4-ply $(0^{\circ}, \pm 45^{\circ}, 90^{\circ})$ laminates for the honeycomb sandwich are shown in Figures 186 and 187. Assembly bagging of laminate skins to honeycomb core for adhesive bonding is presented in Figure 188. A C-scan recording of one honeycomb sandwich panel prior to machining is shown in Figure 189.

4.5.2.2

"h" Frame Fabrication

A three section tool, shown in Figure 190, was used to fabricate the "h" frame members for the front spar panel. The "h" frame was comprised of the three laminate layup shapes: flat, "Z" and "U". Each laminate layup was imidized on the respective tool section. After imidization "Z" and "U" laminate layups and tool sections were assembled, an imidized unidirectional fillet section was placed in the space between "Z" and "U" laminates, and the flat laminate and flat tool plate were positioned to complete the layup/tooling assembly. This was bagged and autoclave cured. Laminate processing was in accordance with 3.2.1.2. A completed section, before machining, is shown in Figure 191.

4.5.2.3

"U" Closeout Ring Fabrication

The "U" closeout ring was fabricated in four 100° arc segments from woven Thornell 300/LARC-160. The segments were fabricated using the tool shown in Figure 192. Laminate processing was in accordance with 3.2.1.3.

4.5.2.4 Front Spar Panel Assembly

A two piece aluminum picture frame tool was developed for assembly bonding of the front spar panel. Figure 193 presents a detail of the tool showning its application for bonding the "h" frames and doublers. FM34B-18 adhesive was used for bonding. Bond processing was in accordance with 3.2.4. Elements of the front spar assembly and the completed assembly are shown in Figure 199 and 195.

4.5.2.5 Front Spar Tee Fabrication

The three configurations of the front spar Tee members, SS79-00253-003, -004 and -005 are presented in Appdendix E. A typical tool for Tee fabrication is shown in Figure 196. The Tee members consists of three sections (two opposed "L" laminates and a flat laminate) and a unidirectional fillet to fill the space at the laminate junction.

The Gr/PI laminates and fillet section were imidized on the appropriate tooling. Typical imidization setup for an "L" laminate is shown in Figure 197. Following imidization, the tooling was assembled with the Gr/PI fillets in place. The assembly was bagged as shown in Figure 198 and autoclave cured. Proccessing was in accordance with 3.2.1.3. A typical Tee is shown in Figure 199.

4.5.3 TDS Rear Spar Fabrication

The TDS rear spar, SS79-00253-006 is presented in Appendix E. It was fabricated on steel tooling, shown in Figure 200, dimensionally corrected for differences in thermal coefficient of expansion between the Gr/PI laminate and the tool. Laminate layup, imidization and autoclave cure was conducted in accordance with 3.2.1.3. After C-scan inspection, load introduction Pi sections were bonded in three places using FM34B-18 adhesive on the spar OML in line with the TDS ribs. The completed aft spar is shown in Figure 201.

Rib Modification for Load Introduction Plates

Installation of the load introduction plates on the TDS for subsequent mechanical testing required flat and parallel rib interface surfaces. To meet this requirement, it was necessary to shim the interface surfaces of the rib Pi caps (SS79-00251-002) and the front spar Tee (SS79-00253-003) to achieve co-planar and parallel surfaces. Shims were fabricated from graphite fabric/LARC-160 and bonded to the TDS rib assembly in the areas shown in Figure 202 using FM34B-18 polyimide adhesive. Only minimum machining was required to establish the co-planar and parallel condition after bonding.

4.5.4

After shim bonding, the location of holes for attaching the load introduction plates to the rib was established using a trim/ drill template. These locations in the rib were potted to provide a solid area for bolt clamp-up pressure in the rib honeycomb core. The potted rib and load introduction plates were match drilled to the trim/drill template using diamond core drills.

4.5.5 Alignment of Ribs to Covers

TDS cover panels (SS79-00250-008) were net trimmed and the location of the three rib assemblies was carefully laid out on the IML skin of the one panel. The three ribs were rigged into position and securely spring-clamped to the cover forward and aft closeout channels (SS79-00250-005 and -006) after establishing the optimum inplane condition for the aft spar attach caps (SS79-00251-003) and the open rib Pi caps (SS79-0025-002) aft of the Tee (SS79-00253-003). At optimum rigging, the open rib Pi caps were inplane while the center rib aft spar attach was out of plane by approximately 0.25-mm (0.010 in.).

In the clamped position, tooling holes were drilled in the rib Pi cap base and the cover IML skin/inner aft closeout channel leg, two places each rib, and slightly aft of the Tee in the rib Pi cap base and the cover IML skin, also two places each rib. Mechanical fasteners were installed in the aft tooling holes and the spring clamps replaced with C-clamps.

The clamped partial assembly was placed on the second cover panel. The two cover panels were squared using large 90° tooling knees and perpendicularity of the three ribs to the cover panels confirmed. The ribs were clamped to the cover panel, tooling holes drilled and mechanical fasteners installed as previously described. Figures 203 and 204 show the TDS in the completed rigged condition and the mechanical fasteners at the aft end holding ribs to covers.

Tees (SS79-00249-004 and -005) were positioned with respect to the rib Tees (SS79-00253-003), clamped in place and tooling holes drilled for precise location.

4.5.6 Assembly of the TDS

Assembly of the TDS was accomplished in three stages as follows:

- Stage 1 Bonding of the ribs and front spar Tees to upper and lower cover panels.
- Stage 2 Bonding of the lower leading edge cover panel and aft spar to the above.
- Stage 3 Mechanically attaching the upper leading edge cover and front spar panels to complete the assembly.

4.5.6.1 TDS Assembly - Stage 1

All bond faying surfaces of the upper and lower cover panels, ribs, and front spar Tees were cleaned to obtain a water break surface and primed as defined in 3.2.4. FM34B-18 adhesive film, .44 Kg/m² (0.09 psf), was applied to the prepared surfaces. Covers, rib, and Tees were assembled to the rigged positon established in 4.5.5.

This assembly was prepared for autoclave bonding as shown in Figure 205. The open channel closeouts, at the forward and aft end of the cover panels, were filled with honeycomb core, as shown in Figure 206, to prevent buckling under bonding pressure. All glass breather plies were taped in place to prevent movement and facilitate the bagging operation. The assembly was enclosed in an envelope bag which was appropriately tailored for the two cavities formed between ribs and covers. Thermocouples were placed to monitor the cure. Figure 207 shows the assembly with breather in place ready for bag installation. A view of the envelope bag through one of the two cavities is shown in Figure 208. Five vacuum valve stems were installed in the envelope bag: four active and one static. The bagged assembly was autoclave cured under full vacuum and 172 KN/m² (25 psi) augmenting pressure at 191°C ($375^{\circ}F$) for two hours.

After cure all bagging materials were removed and the bond areas were visually and nondestructively inspected. NDI utilized harmonic bond testing, Figure 209, and ultrasonic pulse echo contact testing with a delay, Figure 210. Manual recordings were made of any discrepant areas.

The bonded assembly was postcured for two hours at (316C) $600^{\circ}F$.

4.5.6.2 TDS Assembly - Stage 2

The bond faying surfaces of the ribs, cover panels, lower leading edge cover and aft spar assembly were prepared for bonding. FM34B-18 adhesive film was applied to the prepared surfaces and the lower leading edge cover and aft spar assembly positioned. The open channel closeouts of the lower leading edge cover and mating, forward, open channel of the lower cover were filled with honeycomb core. A teflon tube was inserted in the aft open channel closeouts of the upper and lower covers as shown in Figure 211. These tubes extended through the bay and provided equalized pressure during cure.

Bagging, cure, and NDI were as noted in 4.5.6.1. Figures 212 through 216 show the assembly with breather plies in place and after bagging. After cure and NDI, the bonded assembly was postcured.

4.5.6.3 TDS Assembly - Stage 3

Attachment holes in the front spar panels were diamond core drilled using an applied trim/drill tool. The drilled panels were positioned to the front spar Tees for match drilling of holes in these attaching members. All drilling operations were performed with proper backup to prevent fiber breakout on drill exit.

The upper leading edge cover shown in Figure 217 was trimmed to net dimensions on the ends and aft edge. This was positioned on the assembly and forward end trimmed to match the rib trim as shown in Figures 218 and 219 respectively.

Attachment locations were laid out on the upper leading edge cover at the aft edge and ribs were diamond core drilled. Potted inserts were installed for the rib attachments. The cover was clamped in the assembled condition and holes match drilled through the ribs and upper Tee elements of the front spar.

The completed TDS has been delivered for installation of strain gages, thermocouples, and deflection transducer mounting pads in preparation for the ground test phase of this program.

5.0 CONCLUSIONS

The objectives of this program, to develop processes and fabricate demonstration components, have been accomplished. Principal accomplishments and their importance to potential application of graphite/LARC-160 material to Aerospace structures are presented in the areas of process development and component fabrication.

PROCESS DEVELOPMENT

5.1

Quality Assurance of the material system has been furthered by implementation of specifications for material procurement and fabrication processing. Also, nondestructive inspection techniques have been advanced by the cooperative efforts of Langley Research Center and Rockwell International in the establishing of "A" standards for C-scan inspection.

The chemical characterization activity demonstrated the applicability of high pressure liquid chromatography (HPLC) techniques for characterizing the chemical composition of LARC-160 polyimide resin and its mechanism of polymerization. However, this investigation also determined that the HPLC methodology needs further refinement to obtain better reproducibility and improved quantification to satisfactorily analyze all components of the LARC-160 system. At present, it can be said that HPLC provides an indication of material acceptability which must be fully substantiated by appropriate mechanical testing.

The Resin Variables Study indicated the basic range within which changes in both resin formulation and processing could occur without degradation. However, it must be noted that the variation matrix presented in the text could be greatly expanded to include more subtle variations which could not be evaluated within the scope of this program.

Specific processing was developed for fabrication of flat laminates, stiffened panels, honeycomb sandwich panels, and chopped fiber moldings. These processes were demonstrated by fabricating demonstration components which were delivered to LaRC.

Developed processes were qualified by fabrication and mechanical testing. The test data presented in this report provides an adequate starting point for further effort directed to the establishment of design standards necessary for effectual application of the LARC-160 system.

DEMONSTRATION COMPONENTS

The fabrication of demonstration components, laminates, stringer stiffened panels, honeycomb sandwich panels, and chopped fiber moldings, Task (d) through (g), demonstrated the applicability of developed processes to scaled-up structure. Fabrication of the Technology Demonstrator Segment (TDS), a full-size segment of the Space Shuttle aft body flap, demonstrates the applicability of the processes to a manufacturing environment. The TDS is one of the largest all bonded Gr/PI structures fabricated to date.

The results of this program demonstrates that Gr/LARC-160 is a viable material system for structural application. To achieve its potential, however, further development effort must be expended.

5.2

6.0 REFERENCE

- Gibbs, H. H., and J. R. Ness, "Development of Quality Control Techniques for NR150 Polyimide Adhesive and Binder Materials", SAMPE Journal, January/February (1979).
- Lauver, R. W., W. B. Alston, and R. D. Vannucci, "Stability of PMR-Polyimide Monomer Solutions," 34th Annual Conference of SPI Reinforced Plastics/Composites Institute, New Orleans, Louisiana, January 29-February 2, 1979.
- Young P. R., and G. F. Sykes, Polymer Preprints, "Characterization and Aging Effects of LARC-160," ACS/CSJ Chemical Congress, Honolulu, Hawaii, April 1-6, 1979.
- 4. Lauver, R. W., and R. D. Vannucci "Characterization of PMR Polyimides-Correlation of Ester Impurities with Composite Properties," Proceedings 24th National SAMPE Symposium and Exhibition, San Francisco, California, May 8-10, 1979, p. 522.
- Shuart, M. J. and Herakovich, C. A. "An Evaluation of the Sandwich Beam in Four point Bending a Compressive Test Method for Composites" NASA Technical Memorandum 78783, 1978.



Figure 1. Idealized Polymerization Sequence for LARC-160 Polyimide Resin







-ELUTION TIME ----





Figure 4. Liquid Chromatographic Separation of Synthetic Mixture of BTDA and NA Ester Compounds



Chemical Structures of Theoretical Isomer Products of BTDA Esterification Figure 5.















90

`




•







BTDA DIETHYL ESTER PEAK AREA PERCENTAGE



Relative NE Monoethyl Ester in LARC-160 Intermediate Ester Mixtures by HPLC Analysis Figure 14.



Relative NE Monoethyl Ester in LARC-160 Neat Resin and Prepregs by HPLC Analysis Figure 15.









p,p METHYLENE DIANILINE MONONADIMIDE



p,p METHYLENE DIANILINE BIS-NADIMIDE

Figure 17. Chemical Structures of LARC-160 Resin Intermediates









DETECTOR RESPONSE AT 254NM









	PANEL EX217	FORM	JLATION IABLES	PRO VARI	CESS ABLES	
	PREPREG RUN & (BATCH)	CONC. AP-22	CONC. ANHYDRIDES	COOK TIME	REFLUX TIME	
1. A.	1 (22990)	+2%	STD	STD	STD	

Figure 23. C-Scan Resin Variable No. 1



Figure 24. C-Scan Resin Variable No. 2

J		Ba Galer	'ı, aaabaaa				ily Blincum	ส่งและเปลี่	d III	鯯		朝御		li l	anta	andanti				1
1.100 f																				
ų.																				
1																				
																5				
																				ì
	1																			
						2.7														ļ
																	1			
																				1
																				ĥ
	160 C	8 1. dets	C. 10. 10.05000	1.0.15.00.0.0.0	 11010101012-00	(12)1 24034113		LES. (1) (#11.1/2	E 82 8 2 4 2 4 5 7 5	551. 614.14	1.15-11-11-10		111111111	19111111	1111111111111	8m104849	1000000	i ji liniti	igda in	ļ.

PANEL	FORM	ULATION	PROCESS	
EX205	VAR	IABLES	VARIABLES	
PREPREG RUN	CONC.	CONC.	COOK	REFLUX
ε (BATCH)	AP-22	ANHYDRIDES	TIME	TIME
3 (22945)	+5%	STD	STD	STD

Figure 25. C-Scan Resin Variable No. 3



PANEL	FORM	ULATION	PRO	ABLES	
EX206	VAR	IABLES	VARI		
PREPREG RUN	CONC.	CONC.	COOK	REFLUX	
& (BATCH)	AP-22	ANHYDRIDES	TIME	TIME	
4 (22946)	-5%	STD	STD	STD	

Figure 26. C-Scan Resin Variable No. 4



EX20/	VAR	IABLES	VART	ABLES
PREPREG RUN & (BATCH)	CONC. AP-22	CONC. ANHYDRIDES	COOK TIME	REFLUX TIME
5 (22947)	+10%	STD	STD	STD

Figure 27. C-Scan Resin Variable No. 5



PANEL	FORM	ULATION	PROCESS	
EX208	VAR	IABLES	VARIABLES	
PREPREG RUN	CONC.	CONC.	COOK	REFLUX
& (BATCH)	AP-22	ANHYDRIDES	TIME	TIME
6 (22948)	-10%	STD	STD	STD

Figure 28. C-Scan Resin Variable No. 6



PANEL	FORM	ULATION	PRO	CESS	
EX209	VAR	IABLES	VAR I	ABLES	
PREPREG RUN	CONC.	CONC.	COOK	REFLUX	
& (BATCH)	AP-22	ANHYDRIDES	TIME	TIME	
7 (22949)	STD	NA (+5%) BTDA (STD)	STD	STD	

Figure 29. C-Scan Resin Variable No. 7



PANEL	FORM	ULATION	PROCESS		
EX210	VAR	IABLES	VARIABLES		
PREPREG RUN	CONC.	CONC.	COOK	REFLUX	
ε (BATCH)	AP-22	ANHYDRIDES	TIME	TIME	
. <u>8</u> (22950)	STD	NA (-5%) BTDA (STD)	STD	STD	

Figure 30. C-Scan Resin Variable No. 8



PANEL	FORM	ULATION	PRO	CESS
EX211	VAR	IABLES	VAR L	Ables
PREPREG RUN	CONC.	CONC.	COOK	REFLUX
& (BATCH)	AP-22	ANHYDRIDES	TIME	TIME
9 (22951)	STD	NA (STD) BTDA (+5%)	STD	STD

Figure 31. C-Scan Resin Variable No. 9



PANEL	FORM	ULATION	PRO	CESS	
EX212	VAR	IABLES	VARI	ABLES	
PREPREG RUN	CONC.	CONC.	COOK	REFLUX	
ε (BATCH)	AP-22	ANHYDRIDES	TIME	TIME	
10 (22952)	STD	NA (STD) BTDA (-5%)	STD	STD	

Figure 32. C-Scan Resin Variable No. 10

PANEL	FORMU	ULATION	PRO	CESS	
EX213	VAR	IABLES	VARI	ABLES	
PREPREG RUN	CONC.	CONC.	COOK	REFLUX	
& (BATCH)	AP-22	ANHYDRIDES	TIME	TIME	
11 (22953)	STD	ŜTD	2HRS @ 79C	STD	

Figure 33. C-Scan Resin Variable No. 11



Figure 34. C-Scan Resin Variable No. 12



PANEL	FORM	ULATION	PRO	CESS
EX215	VAR	IABLES	VAR1	Ables
PREPREG RUN	CONC. CONC.		COOK	REFLUX
ε (BATCH)	AP-22 ANHYDRIDES		TIME	TIME
13 (22955)	STD	STD	STD	6 HRS

Figure 35. C-Scan Resin Variable No. 13



PANEL	FORMULATION		PROCESS	
EX216	VARIABLES		VARIABLES	
PREPREG RUN	CONC.	CONC.	COOK	REFLUX
ε (BATCH)	AP-22	ANHYDRIDES	TIME	TIME
14 (23107)	STD ANCH- AMINE DL	STD	STD	STD

Figure 36. C-Scan Resin Variable No. 14





PANEL	FORMULATION		PROCESS	
EX219	VARIABLES		VARIABLES	
PREPREG RUN	CONC	CONC	COOK	REFLUX
& (BATCH)	AP-22	ANHYDRIDES	TIME	TIME
16 (24266)	STD	STD	STD	STD

Figure 38. C-Scan Resin Variable No. 16



Figure 39. Variation in Tg With Change in AP22 Concentration



Figure 40. TMA-Tg Values, Amine Variables



Figure 41. TMA-Tg Values NA-BTDA Variables





ŧ



Figure 43. TMA-Tg Values Anchamine & Tonax Amines






Figure 46. C-Scan of Laminate (Batch 23725) Imidizing Pressure 16.9 KN/m² (5 in. Hg)



Figure 47. C-Scan of Laminate)Batch 23725) Imidizing Pressure 6.7 KN/m² (2 in. Hg)



Figure 48. C-Scan of Laminate (Batch 23727) Imidizing Pressure 16.9 KN/m² (5 in. Hg)



Figure 49. C-Scan of Laminate (Batch 23727) Imidizing Pressure 6.7 KN/m² (2 in. Hg)



Figure 50. LARC 160 Preliminary Cure Cycle







COMPOSITE DESCRIPTION: EX 41 NO. OF PLIES/ORIENTATION: 32/0° THICKNESS MM (MILS): 2.03-1.79 (80-70.4) PANEL SIZE CM (INCH): 10.8x12.7 (4.25x5.0)

PROCESS VARIABLE: 163 C (325 F) 60 MINUTES (CONTROL). STANDARD CURE AND POSTCURE CYCLE

EX47



COMPOSITE DESCRIPTION: EX 47

NO. OF PLIES/ORIENTATION: 32/0°

THICKNESS MM (MILS): 2.03-1.79 (80-70.4)

PANEL SIZE CM (INCH): 10.8x12.7 (4.25x5.0)

PROCESS VARIABLE: CURE PRESSURE EVALUATION STUDY-1.0 N/M2 (150 PSI). STANDARD CURE TEMPERATURE AND POSTCURE

Figure 52. C Scan Recordings of Panels EX41 and EX47 Minimum Cure Pressure Study





EXTENSIVE MICRO AND MACRO POROSITY

Figure 53. C Scan Recordings of Panels EX48 and EX49 Minimum Cure Pressure Study

EX74 EX71 A. II/form

COMPOSITE DESCRIPTION: EX 74

NO. OF PLIES/ORIENTATION: 32/0°

THICKNESS MM (MILS): 2.1-1.9 (83.2-73.6)

PANEL SIZE CM (INCH): 10.8x12.7 (4.25x5.0)

PROCESS VARIABLE: CURE TEMPERATURE EVALUATION STUDY-329 C (625 F). 1.3 N/M² (200 PSI). STANDARD POSTCURE





COMPOSITE DESCRIPTION: EX 69 NO. OF PLIES/ORIENTATION: 32/0° THICKNESS MM (MILS): 2.1-1.87 (76.4-73.6) PANEL SIZE CM (INCH): 10.8x12.7 (4.25x5.0) PROCESS VARIABLE: CURE TEMPERATURE EVALUATION STUDY—316 C (600 F) 13 N/M² (200 PSI). STANDARD POSTCURE

EX70



COMPOSITE DESCRIPTION: EX 70

NO. OF PLIES/ORIENTATION: 32/0°

THICKNESS MM (MILS): 2.3-2.2 (89.6-86.4)

PANEL SIZE CM (INCH): 10.8x12.7 (4.25x5.0)

PROCESS VARIABLE: CURE TEMPERATURE EVALUATION STUDY-302 C (575 F), 1.3 N/M2 (200 PSI). STANDARD POSTCURE

Figure 54. C Scan Recordings of Panels EX74, EX69, and EX70 Minimum Cure Temperature Study



EX72



COMPOSITE DESCRIPTION: EX 71

NO. OF PLIES/ORIENTATION: 32/0°

THICKNESS MM (MILS): 2.1-2.0 (83.2-80)

PANEL SIZE: CM (INCH): 10.8x12.7 (4.25x5.0)

PROCESS VARIABLE: CURE TEMPERATURE EVALUATION STUDY-288 C (550 F), 1.3 N/M² (200 PSI). STANDARD POSTCURE

COMPOSITE DESCRIPTION: EX 72

NO. OF PLIES/ORIENTATION: 32/0°

THICKNESS MM (MILS): 2.2-2.1 (86.4-83.2)

PANEL SIZE CM (INCH): 10.8x12.7 (76.4x5.0)

PROCESS VARIABLE: CURE TEMPERATURE EVALUATION STUDY-274 C (525 F) 1.3 N/M² (200 PSI). STANDARD POSTCURE



'											
	SAMPLE:	SAMPLE HEIGHT		X-AXIS SCA	LE	50 <u>810</u>	RUN	NO.			
		LOADING ON TRA	X	Y-AXIS SCA	LE	W	DAT	E //2	- 2 -	28	
		PROBE: EXPANSION/	PENETRATION	Y-AXIS SEN	ѕιтитγ		OPE	RATOR_	Ŵ	0	
	ORIGIN:	HEATING RATE	oc MIN.			IN. PROBE DISPL. IN. OF CHART	. 1 54				
	COMPOSITE DESCRIPTION: EX 74	4							· · · ·		
	PREPREG MFG/BATCH: USP/2W426	52					1	2 EX 74	r pc		
	NO. OF PLIES/ORIENTATION: 3	2/0°									
	THICKNESS MM (MILS): 2.1-1.4	8 (83.2-73.6)				· · ·			· · · ·		
	PANEL SIZE MM (MILS): 15.2x	15.2 (6x6)							2 4	01	
LN3	PROCESS VARIABLE: EFFECT OF THICKNESS ON MECHANI TIES. TARGET THICKNE	LAMINATE CAL PROPER- SS: 20 MM					X		<u> </u>		
CEME	(80 MILS). STANDARD POSTCURE CYCLES	CURE AND				3360					
AJ										• •	•
ISID											
BE											
РКО						346	0	· · · ·	• • •		
		4									
						· · · · · · · · · · · · · · · · · · ·		· · · · · · · · · · · · · · · · · · ·			
		· · · · · · · · · · · · · · · · · · ·		•	 		:			 	
								• •			
0	50 100	150 20	00 25(30	0	350	40	00	45(500
	-	T, °C (CORRECTED	FOR CHROME	EL ALUMEL .	THERMO	COUPLES)					
	F18	gure 56. TMA-T	g Character	tistics of	Specim	en EX 74					

			~					
SAMPLE: LARC		as malicate	K-AXIS SCAL	Е 50	0 0 1 0 1 0	RUN NO		
	LOADING ON TRA	Y Sg.	Y-AXIS SCAL	E .00 2	¥	DATE	11-30-7	00
	PROBE: EXPANSION	PENETRATION	Y-AXIS SENS	ΤΙΛΙΤΥ		OPERAT	OR A.O	
ORIGIN:	HEATING RATE	S MIN.		IN. PROBI	E DISPL.			
COMPOSITE DESCRIPTION: EX 6	6				· · · · ·			
PREPREG MFG/BATCH: USP/2W42	62 2 /0°				· · ·	3	EX 69 45Cu	red 600F
THICKNESS MM (MILS): 2.1-1.	2/0 87 (76.4-73.6)			· · · · · · · ·		(2)	Ex69 Afte	1 PC 68 F
PANEL SIZE CM (INCH): 10.8x	12.7 (4.25×5.0) Frature					/2		
EVALUATION STUDY -31 13 N/M ² (200 PSI).	6 C (600 F) STANDARD			· · · · · · · · · · · · · · · · · · ·				
F DISH					A		TA (1) 3	35°C
8044					358 2		(2) 3.	580
				3380	·			
	2				· · ·			
0 50 100	150 20 T,°C (CORRECTED	DO 250 FOR CHROME) 300 L ALUMEL TH fatice of 6	35(ERMOCOUP) LES) /	400	450	200
4	י-ניזיוי יור שיווא.	-8 VIIALALLEI	TALLCO OF .	ה וומוודהאלנ	202			

CORIGIN: LOADING ON TRAY Y-AXIS SCALE IN DATE ORIGIN: HEATING RATE Y-AXIS SENSITIVITY OPERATOR ORIGIN: HEATING RATE IN OPERATOR ORIGIN: HEATING RATE IN OPERATOR ORIGIN: HEATING RATE IN OPERATOR OPERATOR US IN OPERATOR DREPREG MEG/BATCH: US IN OPERATOR DREPREG MEG/BATCH: US IN IN PAREL SIZE CM (INLLS): I.0. BAI2.7 (4.25x5.0) PAREL SIZE CM (INCH): I.0. BAI2.7 (4.25x5.0) PAREL EXAMINATION IN IN IN PAREL SIZE CM (INCH): I.0. BAI2.7 (4.25x5.0) PAREL SIZE CM (INCH): I.0. BAI2.7 (4.25x5.0) PROCESS VARIAGE: US IN I.3. AVAZ IN IN I.3. AVAZ IN IN I.3. AVAZ		
PROBE: EXPANSION/FENETRATION PROBE: EXPANSION/FENETRATION PROBE: EXPANSION/FENETRATION PREPRES METERS IN MILES PREPRES METERS IN MILES PREPRES METERS IN MILES PROFESS WM (MILES): 2.3-2.2 (89.6-86.4) PANEL SIZE CM (INCH): 10.8x12.7 (4.25x5.0) PROFESS VM (MILES): 2.3-2.2 (89.6-86.4) PROFESS VM (MILES): 2.3-2.2 (89.6	-AXIS SCALE DATE 22	-6-78
ORIGIN: HEATING RATE Image: Composite decident in the	-AXIS SENSITIVITY	0
COMPOSITE DESCRIPTION: EX 70 PREPREG MEG/BATCH: USP/244262 NO. OF PLIES/ORIENTATION: 32/0° THICKNESS MM (MLLS): 2.3-2.2 (89.6-86.4) PAREL SIZE CM (INCH): 10.8x12.7 (4.2555.0) PAREL SIZE CM (INCH): 10.8x12.7 (4.2555.0) PROEES VARIABLE: CURE TERMERATURE EVALUATION STUDY-302 C (575 F), 1.3 N/M2 (200 PS1). STANDARD PRODEE DISPLACEMENT	IN. PROBE DISPL	
PREPREG MEG/BATCH: USP/2W4262 NO. OF PLIES/ORIENTATION: 32/0° THICKNESS MM (MILS): 2.3-2.2 (89.6-86.4) PANEL SIZE CM (INCH): 10.8x12.7 (4.25x5.0) PANEL SI		
NO. OF PLES/ORTENTATION: 32/0° THICKNESS MM (MILS): 2.3-2.2 (89.6-86.4) THICKNESS MM (MILS): 2.3-2.2 (89.6-86.4) PANEL SIZE CM (INCH): 10.8x12.7 (4.25x5.0) PROCESS VARIABLE: CURE TEMPERATURE PROCESS VARIABLE: CURE TE		
THICKNESS WM (MILS): 2.3-2.2 (89.6-86.4) PANEL SIZE CM (INCH): 10.8x12.7 (4.25x5.0) PROCESS VARIABLE: CURE TEMPERATURE PROCESS VARIABLE: CURE TEMPERATURE EVALUATION STUDY-302 C (575 F). 1.3 N/M2 (200 PSI). STANDARD POSTCURE POSTCURE PROCESS VARIABLE: CURE TEMPERATURE PROCESS VARIABLE: CURE PROCESS VARIABLE: CURE PROCESS VARIABLE: CURE TEMPERATURE PROCESS VARIABLE: CURE TEMPERATURE PROCESS VARIABLE: CURE TEMPERATURE POSTCURE PROF PROF PROF PROF PROF PROF PROF POSTCURE	1 EX 70 45	72 mils
PANEL SIZE CM (INCH): 10.8x12.7 (4.25x5.0) PROCESS VARIABLE: CURE TEMPERATURE EVALUATION STUDY = 302 C (575 F). 1.3 N/M2 (200 PS1): STANDARD POSTCURE POSTCURE PROBE DISPLACEMENT 3535 C 353 C		
PROCESS VARIABLE: CURE TEMPERATURE EVALUATION STUDY-302 C (575 F). 1.3 N/M2 (200 PS1). STANDARD POSTCURE		
	12 64 70 46 4	73 mi/s
	3530	
	3176	
0 50 100 150 200 250 300 350 400	300 350 400 4	50 500
T, ^o C (CORRECTED FOR CHROMEL ALUMEL THERMOCOUPLES) Figure 58. TMA-Tg Characteristics of Specimen EX70	. ALUMEL THERMOCOUPLES) ristics of Specimen EX70	

.

500 83 21:15 78 EX 71 AS CURED 804:15 С. О 450 6-445 600F ป : ١ Ż **OPERATOR** Ex 71 RUN NO. 400 DATE ø 3550 ≩⊻ T, °C (CORRECTED FOR CHROMEL ALUMEL. THERMOCOUPLES) ° Å IN. PROBE DISPL. 350 20 Y-AXIS SENSITIVITY 21295 X-AXIS SCALE. Y-AXIS SCALE. 300 250 NIN. PROBE: EXPANSION/PENETRATION 200 LOADING ON TRAY PANEL SIZE: CM (INCH): 10.8x12.7 (4.25x5.0) SAMPLE HEIGHT HEATING RATE. d) PROCESS VARIABLE: CURE TEMPERATURE EVALUATION STUDY-288 C (550 F), 1.3 N/M2 (200 PSI). STANDARD POSTCURE THICKNESS MM (MILS): 2.1-2.0 (83.2-80) 150 NO. OF PLIES/ORIENTATION: 32/0° PREPREG MFG/BATCH: USP/2W4262 COMPOSITE DESCRIPTION: EX 71 100 20 SAMPLE: **ORIGIN:** PROBE DISPLACEMENT

TMA-Tg Characteristics of Specimen EX71 Figure 59.

500 78 mils 7-78 EX72 AS CURED SH \$25F 76 0 . چ PC 4hr 450 12-е. ; 4 EX 72 OPER ATOR. RUN NO. 400 DATE ন্দ Ľ 111 3605 -. 11 ≩l≚ S S IN. PROBE DISPL. IN. OF CHART T, °C (CORRECTED FOR CHROMEL ALUMEL THERMOCOUPLES) 350 20 Y-AXIS SENSITIVITY î X-AXIS SCALE Y-AXIS SCALE. 300 ____ 2570-250 WIN. PROBE: EXPANSION/PENETRATION 200 LOADING ON TRAY SAMPLE HEIGHT HEATING RATE PANEL SIZE CM (INCH): 10.8x12.7 (76.4x5.0) THICKNESS MM (MILS): 2.2-2.1 (86.4-83.2) Τ. PROCESS VARIABLE: CURE TEMPERATURE EVALUATION STUDY-274 C (525 F) 1.3 N/M2 (200 PSI). STANDARD POSTCURE 150 NO. OF PLIES/ORIENTATION: 32/0° COMPOSITE DESCRIPTION: EX 72 PREPREG MFG/RATO PREPREG MFG/BATCH: USP/2W4262 100 50 SAMPLE: **ORIGIN:** TIT TΤ T 0

TMA-Tg Characteristics of Specimen EX 72 Figure 60.



Typical Configuration for Debulking Celion/LARC-160 Prepreg Figure 61.



















Figure 66. C-Scan of Laminate Imidized at 163 C, 60 Minutes



Figure 67. C-Scan of Laminate Imidized at 163 C, 90 Minutes



Figure 68. C-Scan of Laminate Imidized at 163 C, 120 Minutes



Figure 69. C-Scan of Laminate Imidized at 163 C, 150 Minutes



Figure 70. C-Scan of Laminate Imidized at 163 C, 180 Minutes



Figure 71. C-Scan of Laminate Imidized at 177 C, 60 Minutes



Figure 72. C-Scan of Laminate Imidized at 177 C, 90 Minutes

6/2/80 350/120 "A"

Figure 73. C-Scan of Laminate Imidized at 177 C, 120 Minutes



Figure 74. C-Scan of Laminate Imidized at 177 C, 150 Minutes



Figure 75. C-Scan of Laminate Imidized at 177 C, 180 Minutes



Figure 76. C-Scan of Laminate Imidized at 191 C, 30 Minutes



Figure 77. C-Scan of Laminate Imidized at 191 C, 60 Minutes



Figure 78. C-Scan of Laminate Imidized at 191 C, 90 Minutes

3715/120 "A" 6/2/80

Figure 79. C-Scan of Laminate Imidized at 191 C, 120 Minutes

375/150 A" 6/2/80

Figure 80. C-Scan of Laminate Imidized at 191 C, 150 Minutes
390/30 A" 6/2/80

Figure 81. C-Scan of Laminate Imidized at 199 C, 30 Minutes



Figure 82. C-Scan of Laminate Imidized at 199 C, 60 Minutes



Figure 83. C-Scan of Laminate Imidized at 199 C, 90 Minutes

390/120 A" 6/5/80

Figure 84. C-Scan of Laminate Imidized at 199 C, 120 Minutes



Figure 85. C-Scan of Laminate Imidized at 199 C, 150 Minutes



Figure 86. C-Scan of Laminate Imidized at 218 C, 30 Minutes



Figure 87. C-Scan of Laminate Imidized at 218 C, 60 Minutes



Figure 88. C-Scan of Laminate Imidized at 218 C, 90 Minutes



Figure 89. C-Scan of Laminate Imidized at 218 C, 120 Minutes

425/150 "A" 6/2/80 199

Figure 90. C-Scan of Laminate Imidized at 218 C, 150 Minutes







Figure 92. Layup Sequence LARC-160/Celion "I" Beam - Balanced Cap

VACUUM		PRESSURE		AUGMENTED	
PRESSURE		TO AUGMENTER		PRESSURE	
LEVEL		PLATE		TO FILLET	
KN/m ²	I NCHES HG	KN/m ²	PSI	KN/m ²	PSI
6.75	2	6.75	0.98	48	6.9
84.0	25	84.0	12.2	586	85
101	30	101	14.7	710	103

MOLDING PRESSURE CRITERIA (1)(2)(3)

```
(1)
PRESSURE AUGMENTER PLATE AREA RATIO
TO MANDRELS = 7:1
```

(2) PROCESS PER 1.1 AND 1.2.

:

(3) Fillet mold is 96.5 CM (38.0 INCHES) LONG



Figure 93. Fillet Radius Stock Tooling and Vacuum Bagging Arrangement - Imidizing Process



Figure 94. Layup and Tooling Process Hat-Stringer Assembly











Figure 97. C-Scan Laminate EX 199 (0) for Aged Tensile Properties







Figure 99. C-Scan Laminate EX 201 (90)₄₀ for Aged Tensile and Compressive Properties



Figure 100. C-Scan Laminate EX202 (± 45) for Aged Tensile Properties



Figure 101. C-Scan Laminate EX 204 (0)₂₆ for Aged Short Beam Shear and Flexural Properties



Figure 102. C-Scan Laminate EX 220 (+45) 32 Ply for Aged Compressive Properties





FLEX SANDWICH BEAM TEST -ULTIMATE COMPRESSION STRESS ON FACE SHEET



$$F_{cu} = \frac{4P}{W \times t \left(15 + \frac{T+t}{2}\right)}$$

WHERE

P = ULTIMATE FAILING LOAD

W = BEAM WIDTH

t = COMPRESSION FACE SHEET THICKNESS

T = TENSION FACE SHEET THICKNESS

FOIL PER AMS 5510H.

† FOR -270 F, 75 F & 400 F TESTS, USE 0.12 PSF FM 400 ADHESIVE. FOR 600 F TESTS, USE 0.135 PSF ADHESIVE (FM34).

FLEX SANDWICH BEAM TEST -ULTIMATE TENSILE STRESS ON FACE SHEET



LOAD PADS - 1.0 IN. WIDE REACTION PADS - 1.5 IN. WIDE

CALCULATION OF ULTIMATE TENSILE STRESS (Ftu) ON COMPOSITE FACE SHEET

$$F_{tu} = \frac{4P}{W \times t \left(1.5 + \frac{T+t}{2}\right)}$$

WHERE

P = ULTIMATE FAILING LOAD

W = BEAM WIDTH

- t = TENSILE FACE SHEET THICKNESS
- T = COMPRESSION FACE SHEET THICKNESS

* FOR -270 F, 75 F & 400 F TESTS USE ALUMINUM HONEYCOMB CORE, 1/8-22 PCF, 5052 ALLOY.

FOR 600 F TESTS, USE 321 CRES, ANN., 0.005 IN. FOIL PER AMS 5510 H

+ FOR -270 F, 75 F & 400 F TESTS, USE 0.12 PSF FM 400 ADHESIVE. FOR 600 F TESTS, USE 0.135 PSF ADHESIVE (FM34)

Figure 104. Composite Tension and Compression Critical Beam Specimen Designs



			NOMINAL	GAUGE		TAB AD	HESIVE	
LAMINATE	SPEC IMEN		THICKNESS "T"	SECTION		TEST TE	() (C)	
OR I ENTATION	DESIGN	NO. PLIES	(MILS)	CM (INCH)	-132	RT	204	316
(0)t	۵	5	0.032/(12.5)	1.27 (0.500)	FM400	FM400	FM400	FM34
(90) t	60	32	0.20/(80.0)	2.54 (1.000)	FM400	FM400	FM400	FM34
(+45) _S	В	t	0.0254/(10.0)	2.54 (1.000)	FM400	FM400	FM400	FM34
(0, <u>+</u> 45,90) _S	A	ω	0.051/(20.0)	1.588 (0.625)	FM400	FM400	FM400	FM34

Figure 105. Tension Coupon Specimen Designs





Figure 107. Tensile Properties of LARC-160/Celion Laminates Postcured and 125 Hours -316 C (600 F) Aged Conditions



Σ[™]\MM 223ЯT2 3TAMITJU 3∂AЯ3VA





Figure 110. Short Beam Shear Strength of LARC-160/Celion Laminates



Figure 111. Typical C-Scans of "Hat" Elements



Figure 112. "Hat" Stringer Molded on Reverse Formed Tool-Showing Flat Condition







A800122 C-9



Hat Stringer in Position on Skin With "1" Pressure Cauls Installed Figure 115.




A800122 C-11 C







A800122 C-10 C



Figure 118. "Hat" Stiffened Skin/Stringer Showing Concave Skin Surface





EX190 NON VOIDED CAP 120X

Figure 120. C-Scan and Photomicrograph Correlation of "I" Stringer Cap Void Characteristics





Figure 121. "I" Stringers in Dry Fit Position on Skin Assembly



Figure 122. "I" Stringer in Bonding Position With "1" Pressure Cauls Installed



Figure 123. Pressure Augmenter Plate in Position Over "L" Pressure Cauls, "I" Stiffened Skin/Stringer Assembly



Figure 124. "I" Stringer Stiffened Skin Element in Vacuum Bag Bonding Fixture

A800122 C-5

A800125 C-2



A800125 C-1



Figure 125. "I" Stiffened Skin/Stringer Panel Bonded Complete



Figure 126. C-Scan of 35X34 5 Harness Satin Weave Celion Fabric/ LARC-160 Laminate 0.33 cm Thick



Figure 127. "I" Stringer Stiffened Skin Panel Element EX111/EX113 Being Readied for -132 C (-270 F) Test



Figure 128. "I" Stiffened Skin/Stringer Panel, LN2 Manifolds and Baffle Plates in Place









GAGES 1 AND 2 ON LOWER SKIN DIRECTLY OPPOSITE 3 AND 4







GAGES 1 AND 2 ON LOWER SKIN DIRECTLY OPPOSITE 3 AND 4



Figure 132. Compression Load/Strain Characteristics of "Hat" Stringer Stiffened Skin Element EX195-4A Aged 125 Hours at 316 C (600 F) and Tested at-132 C (-270 F)



STRAIN GAGES 1 AND 2 ON LOWER SKIN DIRECTLY OPPOSITE 3 AND 4



Figure 133. Compression Load/Strain Characteristics of Hat Stringer Stiffened Skin Element E109/EX110A, Postcured Condition Tested at Room Temperature



Figure 134. Compression Load/Strain Characteristics of Hat Stringer Stiffened Skin Element EX109/EX110B, Postcured Condition Tested at Room Temperature



GAGES 1 AND 2 ON LOWER SKIN DIRECTLY OPPOSITE 3 AND 4



Figure 135. Load/Strain Characteristics of "Hat" Stringer Stiffened Skin Element EX195-2A Aged 125 Hours at 316 C (600 F) Tested at Room Temperature

• •







Figure 136. Compression Load/Strain Characteristics of "Hat" Stringer Stiffened Skin Element EX195-1PC at 316 C (600 F)





3 AND 4 INSTALLED ON LOWER . SKIN OPPOSITE 1 AND 2







STRAIN GAGES 3 AND 4 INSTALLED ON LOWER SKIN OPPOSITE 1 AND 2



Figure 139. Compression Load/Strain Characteristics of "I" Stringer Stiffened Skin Element EX 194-4A Aged tor 125 Hours at 316 C (600 F) and Tested at -132 C (-270 F)



STRAIN GAGES 3 AND 4 INSTALLED ON LOWER SKIN OPPOSITE 1 AND 2



Figure 140. Compression Load/Strain Characteristics of "I"-Stringer Stiffened Skin Element EX111/EX113, Postcured Condition Tested at Room Temperature



3 AND 4 INSTALLED ON LOWER SKIN OPPOSITE 1 AND 2







STRAIN GAGES 3 AND 4 INSTALLED ON LOWER SKIN OPPOSITE 1 AND 2



STRAIN µ (%)

Figure 142. Compression Load/Strain Characteristics of "I" Stringer Stiffened Skin Element EX194-2A, Aged for 125 Hours at 316 C (600 F), and Tested at Room Temperature



GAGES 3 AND 4 INSTALLED ON LOWER SKIN OPPOSITE 1 AND 2



STRAIN µ (%)

Figure 143. Load/Strain Characteristics of "I" Stringer Stiffened Skin Element EX194-1 Postcured Condition, Tested at 316 C (600 F).



STRAIN GAGES 3 AND 4 INSTALLED ON LOWER SKIN OPPOSITE 1 AND 2











Figure 146. Load/Strain Characteristics of Sandwich Element EX241-3A Aged 125 Hours at 316 C (600 F), Tested at -132 C (-270 F)



Figure 147. Load/Strain Characteristics of Sandwich Panel Element EX150-1, Postcured Condition, Tested at Room Temperature

LOAD K NEWTONS



Load/Strain Characteristics of Sandwich Element EX241-2A Aged 125 Hours at 316 C (600 F) Tested at R.T. Figure 148.



Figure 149. Load/Strain Characteristics of Sandwich Element, EX150-2 Postcured Condition, Tested at 316 C (600 F)











Failures, 316 C (600 F) Test, Postcured




Figure 153. "Hat" Element EX195-IPC Showing Local Debonds and Flange Compression Modes, 316°C (600°F) Test, Postcured Condition.



"Hat" Element EX195-3A Showing Local Flange Compression Modes, 316°C (600°F) Aged Condition.

Figure 154.

A800331 G-11 C



Figure 155. "Hat" Element EX195-3A, Local Skin Compression and Buckling Failure, 316°C (600°F) Test, Aged Condition.



Figure 156. "I" Element EX194-IPC Showing Local Skin Compression Failure, 316 C (600 F) Test Postcured



Figure 157. Skin Compression Failure "I" Stringer Element EX194-4A -132°C (-270°F) Test, Aged Condition.





-



Figure 159. Local Compressive Failure-Sandwich Element EX150-1, RT Test, Side 2, Postcured.

A800331 G-3C



Figure 160. Compressive Failure Mode Sandwich Element EX150-1, RT Test, Side 1, Postcured.



Figure 161. Local Compressive Failure Sandwich Element EX150-2, 316°C (600°F) Test, Postcured.

A800331 G-8 C



Figure 162A. Failure Modes of Sandwich Element EX 241-1, Postcured Condition, Tested at - 132 C (-220 F) FAILING LOAD: 160 KN (36,000 LB)

SIDE 1

¢



Failure Modes of Sandwich Element EX 241-1, Postcured Condition, Tested at -132 C (-220 F) Figure 162B.



Figure 163A. Failure Modes of Sandwich Element EX 241-3A Aged 125 Hours at 316 C (600 F), Tested at -132 C (-270 F)

FAILING LOAD: 156 KN (35,100 LB)

SIDE 1

A800808 G-4



Figure 163B. Failure Modes of Sandwich Element EX 241-3A Aged 125 Hours at 316 C (600 F), Tested at -132 C (-270 F)

SIDE 2



Figure 164A. Failure Modes of Sandwich Element EX 241-2A, Aged 125 Hours at 316 C (600 F), Tested at RT

FAILING LOAD: 136 KN (30,650 LB)

A800808 G-6



Figure 164B. Failure Modes of Sandwich Element EX 241-2A, Aged 125 Hours at 316 C (600 F), Tested at RT



Figure 165A. Failure Modes of Sandwich Element EX 241-4A, Aged 125 Hours at 316 C (600 F) Tested at 316 C (600 F)

FAILING LOAD: 120 KN (27,000 LB)

SIDE 1



Figure 165B. Failure Modes of Sandwich Element EX 241-4A, Aged 125 Hours at 316 C (600 F) Tested at 316 C (600 F)

A800808 G-9



REF. "ANALYTICAL AND EXPERIMENTAL STUDY OF STRUCTURALLY EFFICIENT COMPOSITE HAT-STIFFENED PANELS LOADED IN AXIAL COMPRESSION"

JERRY G. WILLIAMS & MARTIN M. MIKULAS, JR. NASA-LARC AIAA PAPER #75-754, 1975

Figure 166. Comparison of Structural Efficiencies of Compression Panels.



Figure 167. C-Scan Laminate CL 6C-11, $(0, \pm 45)_s$, 6 Plies



Figure 168. C-Scan Laminate CL 12C-6, (0,<u>+</u>45)_s, 12 Plies



Figure 169. C-Scan Laminate CL 24C-7, (0+45), 24 Plies



Figure 170. Three - 122 X 26.0 Cm Hat Stringer/Skin Panels Delivered to NASA-LaRC



Figure 171. End Views of Hat Stringer/Skin Panels Delivered to NASA-LaRC



EX279/278/27 and EX249/248/245 Panels Joints, Bond Skin t t Stringer of Figure 172. C-Scans







Figure 174. C-scan Laminate CL8C-18, $(0, \pm 45, 90)_S$, 8 Plies



Figure 175. C-Scan Laminate Cl 12C-8, (0,90)_t, 12 Plies, Skin for Honeycomb Panels



Figure 176. C-Scan Laminate CL 12C-9, (0,90)_t, 12 Plies, Skin for Honeycomb Panels



Figure 177. C-Scan Honeycomb Panel 8A with Location of 25.4 X 25.4-cm Panels No. 1 and No. 2



Figure 178. C-Scan Honeycomb Panel 9A with Location of 25.4 X 25.4-cm Panels No. 3 and No. 4



Figure 179. C-Scan Honeycomb Panel 9C with Location of 25.4 X 25.4-cm Panels No. 5 and No. 6







Figure 181. Celion/LARC 160 Molding Compound Complex Part Demonstration Mold





Figure 182. Complex Shaped Process Demonstration Part Showing Good Compound Flow and Mold Conformance—Imidized at 191°C (375°F), Removed From Mold Cold



Figure 183. Representative Demonstration Component





A810826 F-3








Figure 186. Bagging Assembly for Imidizing



FOR CERTAIN LAYUPS, THE 120 CLOTH, STAINLESS STEEL CAUL PLATE AND 162 CLOTH WERE NOT USED

Figure 187. Bagging Assembly for Autoclave Curing



Figure 188. Bagging Assembly for Honeycomb Sandwich Panel Bonding







Figure 190. Sketch of Three Section Tool for "h" Frame Member



Figure 191. Cured "h" Section Before Machining

A801003 C-14



Figure 192. "U" Closeout Ring Tool







Figure 194. Elements of TDS Front Spar Panel



Figure 195. TDS Front Spar Panel Bonded Assembly







Figure 197. Imidizing Configuration for Tee Member Element

AUTOCLAVE CURE FOR TEE MEMBER (END VIEW)



Figure 198. Tee Member Bagging for Autoclave Cure

A801023 G-11



Figure 199. Cured Tee Member Before Machining



Figure 200. TDS Rear Spar Tool



Figure 201. Load Introduction Pi Sections Bonded to the Machined TDS Rear Spar



Figure 202. TDS Rib Areas Requiring Shim Bonding



Figure 203. TDS in Rigged Condition



Figure 204. Mechanical Fasteners Hold Ribs to Covers at Aft End of TDS



Figure 205. Bagging Configuration for Bonding Ribs to Covers









A810807 A-6



Figure 208. TDS Bagged for Bonding Cycle

A810811 G-3



Figure 209. Harmonic Bond Testing

A810727 G-6



Figure 210. Ultrasonic Pulse Echo Contact Testing





A810807 A-9



Figure 212. Lower Leading Edge Cover With Breather Material in Place

A810807 A-38





A810807 A-33



Figure 214. TDS Ready for Bagging

A810807 A-34



A810811 G-1



Figure 216. TDS Bagged for Bonding Cycle.

A810811 G-2





A810803 A-19



Figure 218. Partially Trimmed Upper Leading Edge Cover Clamped in Position on Bonded TDS Assembly



Figure 219. Final Trim of TDS Upper Leading Edge Cover



Table 1. LARC-160 Intermediate Ester Batches

Batch Code

Standard Production Batches

Α	22367 544 F	Run 2	Standard Compositio	n &	Processing
В	22361 544 1	run 4	Standard Compositio	n &	Processing
С	22408 545 F	Run 1	Standard Compositio	n &	Processing
D	22746 545 F	Run 1	Standard Compositio	n &	Processing
E	22746 545 1	Run 2	Standard Compositio	n &	Processing
F	22746 545 F	Run 3	Standard Compositio	n &	Processing

Variables Study Batches

22943	Standard Composition & Processing
22944	Standard Composition & Processing
22945	Standard Composition & Processing
22946	Standard Composition & Processing
22947	Standard Composition & Processing
22948	Standard Composition & Processing
22990	Standard Composition & Processing
22991	Standard Composition & Processing
22949	+5% NA, Standard Processing
22950	+5% BTDA, Standard Processing
22951	-5% NA, Standard Processing
22952	-5% BTDA, Standard Processing
22953	Standard Composition & Processing
22954	Standard Composition & Processing
22955	Standard Composition, 6 hr Reflux
23107	Standard Composition & Processing
23236	Standard Composition & Processing
23357	Standard Composition & Processing
	22943 22944 22945 22946 22947 22948 22990 22991 22949 22950 22951 22952 22953 22953 22954 22955 23107 23236 23357

Repeatability Batches

G	23724	Standard Composition & Processing
Н	23726	Standard Composition & Processing
Ι	23728	Standard Composition & Processing

.

Table 2. LARC-160 Neat Resin and Prepreg Batches

Batch Code	Neat Resin	Prepreg
А	544 22408	Not Available
В	544 22513	Not Available
C	544 22368 Cut 3	22368 Roll 1
D	544 22361 Cut 4	22361 Roll 1
Ε	544 22367 Cut 2	22367 Roll 1
F	Not Available	22322 Roll 1
G	544 22332 Cut 1	22332 Cut 1
H	Not Available	22999-2
I	Not Available	22999-3
J	Not Available	22999-4
K	Not Available	22999-11
L	Not Available	23091 Roll 4

Variables Study Batches

	Neat Resin	Prepreg	
1	22943	22943	+2% AP-22. Standard Processing
2	22944	22944	-2% AP-22, Standard Processing
3	22945	22945	+5% AP-22, Standard Processing
4	22946	22946	-5% AP-22, Standard Processing
5	22947	22947	+10% AP-22, Standard Processing
6	22948	22948	-10% AP-22, Standard Processing
1A	22990	22990	+2% AP-22, Standard Processing
2A	22991	22991	-2% AP-22, Standard Processing
7	22949	22949	+5% NA, Standard Processing
8	22950	22950	+5% BTDA, Standard Processing
9	22951	22951	-5% NA, Standard Processing
10	22952	22952	-5% BTDA, Standard Processing
11	22953	22953	Standard Composition, Extended Resin Cook
12	22954	22954	Standard Comp., Intermediate Resin Cook
13	22955	22955	Standard Composition, 6 hr Ester Reflux
14	22956	22956	Anchamine DL, Std. Comp. & Proc.
15	23236	23236	Tonox-22, Standard Composition & Proc.
16	23357	23427	Standard Composition & Processing

Repeatability Batches

М	23724	23723
Ν	23726	23725
0	23728	23727
Table 3. Experimental Conditions for HPLC Analysis of LARC-160 Resin

Column: Spectra-Physics Sperisorb ODS

Solvents: Baker HPLC water with 0.01 M KH_2PO_4 at pH=3 WITH HCL Burdick & Jackson Acetonitrile

Gradient: 10-50% acetonitrile/water, 30 min linear gradient with hold at 50% 10 min and equilibrate at initial for 10 min.

Detection: 200 nm, 0.4 aufs

Flow: 1 ml/min

Sample: 10 μ l of 1.5 mg/ml solution in THF

Table 4. Experimental Conditions for Ion-Pair HPLC Analysis of LARC-160 Resin

Column: Whatman Partisil 10, ODS-2

Solvents: Baker HPLC Water with Waters PIC A Ion Pair Reagent Burdick & Jackson UV Grade Tetrahydrofuran (THF)

Gradient: 15-50% THF in water with PIC A, linear 15 min gradient with 15 min hold at 50% THF and equilibrate at initial 15 min

Detection: 254 nm, 0.4 aufs

Flow: 1 ml/min

Sample: 10 μ of 1.5 mg/ml solution in THF

Test Matrix Effect of Resin Formulation/Process Variables Table 5.

	PANEL NO.	EX217	EX218	EX205	EX206	EX207	EX208	EX209	EX210	EX211	EX212	EX213	EX214	EX215	EX216	EX219	
	LAM	~	, ,	>	>	>	>	~	>	>	>	>	~	~	>	>	
	PREPREG EXTRACT	~	>	>	>	>	>	>	>	>	>		>	>	>	~	>
ES	RESIN	~	>	>	>	>	>	~	>	>	>	>	~	~	>	>	>
ANAL YS	INT. ESTER	~	>	>	>	>	>	~	>	>	>	>	~	1	>	>	>
	AP-22	~															· /
	BTDA	>															/
	NA	>															>
ARIABLES	REFLUX TIME	STD	-				STD	STD			STD	STD	STD	6 HR	STD	STD	STD
PROCESS V	C00K TIME	STD					STD	STD A			STD	2 HRS AT 70C	2 HRS AT 60 C	STD	STD	STD	STD
VARIABLES	CONC. ANHYDRIDES	STD		-			STD	NA (+5%) BTDA (STD)	NA (-5%) BTDA (STD)	NA (STD) BTDA (+5%)	NA (STD) BTDA (-5%)	STD	STD	STD	STD	STD	STD
FORMULATION	CONC. AP-22	+2%	-2%	+5%	+5%	+10%	-10%	STD			STD	STD	STD	STD	ANCHAMINE DL STD	TONOX 22 STD	STD
PREPREG	BAICH 1.35 KG (3 LB) EA	-	2	m	4	ß	6	7	œ	б	10		12	13	14	15	91

Table	6.	Prepreg and	Composite Physical,	Short Beam Shear and	Flexural Properties,
		LARC-160	Resin Stoichiometry	and Process Variable	Program

						1		r			
Properties Laminate No.	Target(2) Property	1	X217	I	X218	E	:x205	. I	X206		8X207
Resin/Process Variable											
 Resin Run No. Concentration AP22 Concentration Anhydrides Cook time Reflux time 			1 +2% STD STD STD		2 - 2% STD STD STD		3 +5% STD STD STD		4 -5% STD STD STD		5 +191 570 570 570
Processing Parameters ⁽¹⁾											
 Type of bleeder/No. Plies(6) 	-	120	/2T & 1B	120	/2T & 1B	120	/1Т & В	120	/1T & B	12	0/1T & B
Prepreg Physical Properties							· · ·		·····		
 Prepreg batch Fiber areal weight (grams/m²) Calc. Thick./ply, 60% fiber vol, mm (mils) Resin solids content (%) as is Resin solids content staged (%)(5) Volatile content as in (%) Volatile content staged (%)(5) 	134+4 0.131-0.124 (5.2-4.9) 38.0 ± 3 32-36 9-14 <2	22 12 0. 41 32 15 1.	991 7 122 (4.8) .2 .2 .6 13	23 11 0. 43 31 13 1.	107 8 112 (4.4) .0 .7 .8 11	22 13 0. 42 28 14 3.	945 1 127 (5.0) .1 .4 .3 04	22 12 0. 40 30 15 2.	946 7 122 (4.8) .4 .5 .2 65	229 120 0.1 42 34 14 1.5	247 5 22 (4.8) 8 4 5 7
Composite Physical Properties(1)											
 Specific gravity (grams/cc) Resin weight content (%) Fiber volume (%) Void volume (%) Thickness nm (mils) Thickness (ASTM 2583) Weight lass in postcure (%) 	1.573-1.591 31.05-34.71 58-62 <2 >70 <1	1. 34 59 -0 1.42-1.6 0.102-0. 72- 0.	59 .0 .3 .18 0 (56-63) 114 (4.0-4.5) 75 23	1. 29 63 0. 1.45-1.6 0.104-0. 72- 0.	595 .1 .9 97 5 (57-65) 117 (4.1-4.6) 74 23	1. 30 62 0. 1.16-1.3 0.084-0. 72- 0.	59 .8 .2 74 7 (46-54) 097 (3.3-3.8) 76 27	1. 28 64 0. 1.45-1.5 0.104-0. 74- 0.	60 .2 .9 92 5 (57-61) 112 (4.1-4.4) 76 24	1. 31. 60 2. 1.63-1.7 9.117-0. 64- 0.2	66 8 1 3 6 (64-69) 24 (4.6-4.9 58 24
9. weight loss after 125 nrs at 316 C (600 F) (%)	<3	1.	48	1.	40	1.	58	1.4	46	1.3	57
10. TMA-1g C, (F) postcured 4 hrs at 316 C (600 F)	>340 (644)	356	(673)	340	(644)	347	(357)	337	(639)	36	(693)
Cured Postcured 4 hrs at 316 C (600 F)	>95 >95	10 10	0	10 10	0	10))	10 10	0	100 100	1
Composite Mechanical Properties(4)	MN/m ² (Ksi)	MN/m ²	(Ksi)	MN/m ²	(Ksi)	'MN/m ²	(Ksi)	MN/m ²	(Ksi)	MN/m ²	(Ksi)
1. Flexural strength RT		1840	(253) (278) (<u>269)</u> (267)	1778	(268) (254) (252) (258)	1812	(254) (267) (267) (263)	1833	(273) (250) (274) (266)	1530	(222) (228) (216) (222)
Avg normalized strength, 60% F/V	>1571 (>228)	1861	(270)	1669	(242)	1750	(254)	1695	(246)	1530	(222)
316 C (600 F)	Avg	1033	(161) (130) (159) (150)	916	(132) (132) (134) (133)	1096	(160) (149) <u>(169)</u> (159)	978	(155) (138) (132) (142)	937	(131) (136) (142) (136)
Avg normalized strength, 60% F/V	>937 (>136)	1046	(152)	860	(125)	1057	(153)	900	131	(937)	(136)
2. Flexural modulus RT	CN/m ² (Msi) Avg	GN/m ²	(Ms1) (19.7) (17.3) (18.5) (18.5)	GN/m ²	(Msi) (19.3) (19.3) (19.7) (19.4)	GN/m ²	(Ms1) (19.6) (19.9) (18.7) (19.4)	GN/m ²	(Msi) (21.0) (18.4) (19.4) (19.6)	GN/m ²	(Ms1) (18.0) (25.2) (17.3) (20.2)
Avg normalized modulus, 60% F/V	>124 (>18)	129	(18.7)	126	(18,2)	129	(18.7)	124	18.0	140	(20.2)
316 C (600 F)	Avg	132	(20.5) (16.5) (20.3) (19.1)	916	$(17.4) \\ (16.9) \\ (18.5) \\ (17.6)$	130	(19.2) (18.1) (19.2) (18.8)	121	(19.1) (18.1) (15.2) (17.5)	119	(17.8)(17.4)(16.9)(17.4)
Avg normalized modulus, 60% F/V	>124 (>18)	133	(19.3)	114	(16.5)	125	(18.1)	111	(16.1)	119	(17.4)
 Short beam shear strength RT 	MN/m ² (Ks1) >103 (>15) Avg	MN/m ²	(Ks1) (17.1) (17.9) (17.9) (17.6)	MN/m ²	(Ks1) (16.2) (16.2) (16.6) (16.3)	MN/m ²	(Ksi) (16.4) (17.1) (18.1) (17.2)	MN/m ²	(Ksi) (16.4) (18.4) <u>(17.6)</u> (17.5)	MN/m ²	(Ks1) (11.8) (12.5) (13.1) (12.5)
316 C (600 F)	>48 (>7) Avg	55	(8.9) (8.2) (6.9) (8.0)	43	(6.5) (6.4) (5.9) (6.3)	53	(8.4) (7.3) (7.4) (7.7)	41	(6.1) (6.2) $-\frac{(5.8)}{(6.0)}$	59	(8.8) (8.6) (8.5) (8.6)

,

Table 6. Prepreg and Composite Physical, Short Beam Shear and Flexural Properties, LARC-160 Resin Stoichiometry and Process Variable Program (Cont)

Properties	T	1		<u> </u>				[
Laminate No.	Target(2) Property	E	208	E	x209	EX	210	EX	211	EX2.	12
Resin/Process Variable								1			
 Resin Run No. Concentration AP22 Concentration anhydrides Cook time Reflux time 		6 -1 ST ST ST	0% D D	7 S1 NA (+5%) S1 S1	TD , BTDA(STD) TD TD	8 S NA(-5%), B S S	TD TDA (STD) TD TD	9 S' NA(STD), S' S'	TD BTDA(+5%) TD TD	10 STI XA(STR), 1 STI STI) 3TDA (~5.%))
Processing Parameters(1)		1									
1. Type of bleeder/no. plies(6)		120/2	T & 1B	181/1	LT & 1B	120/1	B only	120/21	Г & 18	120/2T	6 1B
Prepreg Physical Properties											
 Prepreg batch Fiber areal veight (grams/m²) Calc thick/ply, 60% fiber vol, mm (mils) Resin solids content (%) as is Resin solids content staged (%) (5) Volatile content as is (%) Volatile content staged (%) 	134+4 0.131-0.124 (5.2-4.9) 38.0 +3 32-36 9-14 <2	2294 132 0.12 36.9 30.7 15.1 2.92	8 4 (4.9)	2294 122 0.11 40.0 30.3 14.1 2.09	49 17 (4.6) 3 1	2295 122 0.11 42.0 35.7 14.0 2.79) 7 (4.6)	2295: 116 0.111 43.0 30.8 14.3 1.58	1 2 (4.4)	22952 120 0.112 (42.1 28.3 14.8 1.34	4.4)
Composite Physical Properties(1)											
 Specific gravity (grams/cc) Resin weight content (X) Fiber volume (X) Void volume (X) Thickness nm (mils) Thickness/ply, mm (mils) Thickness/ply, mm (mils) Weight loss in posture (X) Weight loss after 125 hr at 316°C (600°F) (X) The C. (F) postcured 4 hr at 316°C (600°F) Cured 	1.573-1.591 31.05-34.71 58-62 <2 >70 <1 <3 >340 (644) >85	1.62 25.0 68.6 0.68 1.45-1.62 0.104-0.11 1.48 0.24 1.48 339	(57-64) 6 (4.1-4.6) (642)	1.60 28.6 64.2 0.91 1.55-1.75 0.117-0.12 1.48 0.22 1.48 364	(61-69) (61-69) (64-4-9) (687)	1.57 33.3 59.2 1.24 1.37-1.55 0.099-0.11 1.5 0.25 1.51 340	(54-61) 2 (3.9-4.4) (644)	1.59 30.7 67.2 2.02 1.32-1.55 0.094-4.4 1.31 0.23 1.31 340 ((52-61) (3.7-4.35) (644)	1.5 31. 61. 1.2 1.47-1.57 (5 0.104-0.112 1.1 0.2 1.1 353	8 5 6 8-62) (4.1-4.4) 7 3 7 (667)
Postcured 4 hr at 316°C (600°F)	>95	100		100		99 100		100		100	
Composite Mechanical Properties(4)			Γ								
1. Flexural strength RT	MN/m ² (Ksi)	MN/m ²	(Ks1) (305) (321) (274) (200)	MN/m ²	(Ks1) (232) (245) (233) (227)	MN/m ²	(Ksi) (333) (248) (270)	MN/m ²	(Ksi) (296) (319) (283)	MN/m ²	(Ksi) (248) (262) (260)
Avg normalized strength, 60% F/V	>1571 (>228)	1808	(262)	1526	(221)	1930	(283)	1839	(299)	1728	(251)
316°C (600°F)	Avg	1027	(144) (155) <u>(148)</u> (149)	1013	(144) (157) (140) (147)	1102	(167) (155) (159) (160)	1109	(180) (143 (160) (161)	985	(148) (134) (147) (143)
Avg normalized strength, 60% F/V	>937 (>136)	898	(130)	942	(137)	1113	(162)	987	(143)	956	(139)
2. Flexural modulus RT	GN/m ² (Msi)	GN/m ²	(Msi) (20.0) (18.5) (19.1) (10.2)	GN/m ²	(Msi) (18.2) (17.9) (18.0)	GN/m ²	(Ms1) (20.4) (20.0) (22.5)	MN/m ²	(Msi) (19.1) (20.3) (19.1)	GN/m ²	(Ms1) (18.4) (22.5) (18.7)
Avg normalized modulus, 60% F/V	>124 (>18)	115	(19.2)	115	(16.7)	145	(21.0)	134	(19.5)	137	(19.9)
316°C (600°F)	Aug	121	$(16.8) \\ (17.6) \\ (18.4) \\ (17.6) \\ ($	119	$(17.4) \\ (16.7) \\ (17.5) \\ (17.2) \\ ($	127	(19.4) (17.9) (18.3) (18.5)	115	(20.4) (19.4) (18.4) (10.4)		(18.2) (15.7) (17.8) (17.0)
Avg normalized modulus, 60% F/V	>124 (>18)	106	(15, 3)	110	(17.2)	129	(18.7)	119	(19.4)	115	(17.2)
3. Short beam shear strength	MN/m ² (Ksf)	MN/m ²	(Ka1)	MN/m ²	(Ket)	MN/m ²	(Kei)	MN/m ²	(Ket)	MN /m ²	(Ket)
RT	>103 (>15)		(17.6)		(13.9)		(18.0)	. at / at	(17.5)	4 MT / IL -	(16.9)
		120	(17.8) (16.9) (17.4)		(16.3) (15.5) (15.2)		(17.2) (16.0) (17.1)		(16.3) (15.5) (16.4)		(16.6) (16.5) (16.3)
316°C (600°F)	>48 (>7)	2.00	(8.1)	103	(8.6)	110	(7.0)	11.7	(8.4)	113	(8.5)
	Avg	54	(7.6) (7.9) (7.9)	 58	(8.3) (7.9) (8.3)	54	(8.3) (8.5) (7.9)	58	(8.3) (8.5) (8.4)		(8.1) (8.4) (8.3)

(1) The inidizing 2 stage cure cycle and tooling specified was employed in fabrication of 14 ply undirectionsl laminates 17.78 x 13.59 Cm (7.7 x 5.5 inches). Laminates were postcured at 316 C (600 F) for 4 hours, freestanding in an air circulating oven. Prepreg and composite physical properties were calculated per Appendix A.
 (2) Target property values are based on Celion fiber minimum properties of 2618 NN/m², (360 ksi) tensile strength and 234 CN/m², (34 Msi) tensile modulus using the rule of mixtures, 602 composite fiber volume. Target 316°C (600 F?) strength properties are based on a 60 percent retention of room values.
 (3) NDI ultrasonic through transmission tests were performed using the NASA-LARC established "A" sensitivity standards.
 (4) Specimens were tested after stabilizing at 316°C (600°F?) for 10 minutes.
 (5) Volattles and resin solids content determined on portion of stacked laminate.
 (6) T = number of bleeder plies on top surface of laminate; B = number of bleeder plies on bottom surface.

Properties	Target ⁽²⁾					F	vo1:=				/ . 1 <i>r</i> .
Resin/Process Variable	Toperty					E	A215	<u>_</u>	A210	F.7	
 Resin run no. Concentration AP22 Concentration anhydrides Cook time Reflux time 		11 STD. STD. Exte STD.	nded	12 STD. STD. Inte STD.	rmediate	13 STD. STD. STD. 6 hr		14 STD. STD. STD. STD.	Ancamine	15 STD. STD. STD. STD.	Tonax
Processing Parameters ⁽¹⁾											
1. Type of bleeder/No. Plies ⁽⁶⁾	-	181/	1T & 1B	120/	2T & 1B	120/	2T & 1B	120/	27 & 18	120/1	IT & IB
Prepreg Physical Properties											
 Prepreg batch Fiber areal weight (grams/m²) Calc. Thick./ply, 60% fiber vol, mm (mils) Resin solids content (%) as is Resin solids content staged (%) Volatile content as is (%) Volatile content staged (%) 	134+4 0.131-0.124 (5.2-4.9) 38.0 ± 3 32-36 9-14 <2	2295 115 0.10 46.1 37.9 12.7 1.62	3 9 (4.3)	2295 120 0.11 40.5 33.9 12.9 1.36	4 (4.5)	22955 121 0,111 40.0 30.3 14.8 1.64	5 7 (4.6)	22994 126 0.119 43.0 31.9 12.0 1.39	9 (4.7)	23236 133 0.127 38.9 11.7	5.0)
Composite Physical Properties(1)					•				· · · · · · · · · · · · · · · · · · ·		
 Specific gravity (grams/cc) Resin weight content (%) Fiber volume (%) Void volume (%) Thickness mm (mils) Thickness/Ply, mm (mils) Barcol hardness (ASTM 2583) Weight loss in postcure (%) Weight loss in postcure (%) Weight loss after 125 hrs at 316 C (600 F) (%) TKA-Tg C, (F) Postcured 4 hrs at 316 C (600 F) C. Scon ultra sound transitson (%) 	1.573-1.591 31.05-34.71 58-62 <2 >70 <1 <3 >340 (644)	1.56 36.0 56.1 1.04 1.62-1.78 0.117-0.1 69-73 0.27 1.73 337 (639)	(64-70) 27 (4.6-5.0) 3	1.59 30.5 62.4 0.82 1.49-1.80 0.106-0.1 69-73 0.21 1.45 340 (644)	(59-71) 29 (4.2-5.1)	1.59 31.6 61.6 0.31 1.39-1.65 0.099-0.14 73-76 0.26 1.34 340 (644)	(55-65) 7 (3.9-4.6)	1.59 29.5 63.3 1.19 1.52-1.72 0.109-0.12 72-74 0.25 1.37 355 (671)	(60-68) 24 (4.3-4.9)	1.489 32.7 56.6 6.5 1.72-1.98 0.124-0.14 64-68 0.23 2.12 374 (705)	(68-78) 2 (4.9-5.6)
Cured Postcured 4 brs at 316 C (600 F)	>95 >95	100		100		100		60 70		0	
Composite Mechanical Properties ⁽⁴⁾		MN/m ²	(Ks1)	MN/m ²	(Ksi)	MN/m ²	(Ksi)	MN/m ²	(Ksi)	MN/m ²	(Ksi)
1. Flexural strength RT	MN/m ² (Ksi)	1557	(219) (243) (221) (227)	1612	(237) (225) (240) (234)	1743	(261) (239) (262) (254)	1626	(240) (229) (240) (236)	1233	(188) (180) (169) (179)
Avg normalized strength, 60% F/V	>1571 (>228)	1665	(242)	1550	(225)	1698	(246)	1541	(224)	1307	(190)
316 C (600F)	·		(120)		(146)		(156)		(147)		(121)
	Avg	896	(119) (151) (130)	1027	(155) (145) (149)	1047	(144) (156) (152)	985	(138) (145) (143)	820	(110) (125) (119)
Avg normalized strength, 60% F/V	>937 (>136)	958	(139)	987	(143)	1020	(148)	934	(134)	869	(126)
2. Flexural modulus RT	GN/m ² (Msi) Avg	GN/m ²	(Ms1) (17.7) (18.5) <u>(18.3)</u> (18.2)	GN/m ²	(Ms1) (20.0) (19.6) (20.3) (20.0)	GN/m ²	(Ms1) (20,0) (19.2) <u>(18.2)</u> 19.1	GN/m ²	(Msi) (18.2) (18.7) <u>(18.7)</u> (18.5)	GN/m ²	(Mst) (15.6) (16.0) (15.1) (15.6
Avg normalized modulus, 60% F/V	>124 (>18)	134	(19.5)	132	(19.2)	128	(18.6)	121	(17.5)	114	(16.5)
316 C (600 F)	Avg	112	(15.7) (15.5) (17.7) (16.3)	127	(18.3) (19.2) (17.8) (18.4)	127	(19.5) (17.5) (18.2) (18.4)	119	(17.5) (16.7) (17.5) (17.2)	114	(16.7) (15.7) (17.2) (16.5)
Avg normalized modulus, 60% F/V	>124 (>18)	120	(17.4)	122	(17.7)	123	(17.9)	112	.(16.3)	120	(17.4)
3. Short beam shear strength RT	MN/m ² (Ksi) >103 (>15) Avg	MN/m ²	(Ksi) (17.6) (16.8) (16.3) (16.9)	MN/m ²	(Ks1) (18.1) (16.9) <u>(18.0)</u> (17.7)	MN/m ²	(Ks1) (16.3) (17.4) (17.4) (17.0)	MN/m ²	(Ksi) (14.2) (12.8) (13.8) (13.6)	MN/m ²	(Ksi) (7.4) (10.3) (7.7) (8.5)
316 C (600 F)	>48 (>7) Avg	48	(5.9) (5.9) (9.3) (7.0)	67	(9.1) (9.4) <u>(10.6)</u> (9.7)	64	(11.1) (8.6) (8.2) (9.3)	47	(7.6) (6.7) (6.5) (6.9)	33	(5.1) (4.5) (4.9) (4.8)

Table 6". Prepreg and Composite Physical, Short Beam Shear and Flexural Properties, LARC-160 Resin Stoichiometry and Process Variable Program (Cont)

(1) The imidizing 2 stage cure cycle and tooling specified was employed in fabrication of 14 ply unidirectional laminates 17.78 x 13.59 cm (7.0 x 5.5 inches). Laminates were postcured at 316°C (600°F) for 4 hours, freestanding in an air circulating oven. Prepreg and composite physical properties were calculated per Appendix A.

(2) Target property values are based on Celion fiber minimum properties of 2618 MN/m², (380 Ksi) tensile strength and 234 GN/m², (34 Ksi) tensile modulus using the rule of mixtures, 60% composite fiber volume. Target 316 C (600 F) strength properties are based on a 60 percent retention of room values.

(3) NDI ultrasonic through transmission tests were performed using the NASA-LaRC established "A" sensitivity standards.

 $\ensuremath{^{(4)}}$ Specimens were tested after stabilizing at 316 C (600 F) for 10 minutes.

(5) $_{Volatiles}$ and resin solids content determined on portion of stacked laminate.

(6) $T \approx$ number of bleeder plies on top surface of laminate; $\beta \approx$ number of bleeder plies on bottom surface.

	<u> </u>		•	<u> </u>	• •				•		+			+		+		+
	74		2-73.6)) (2.6-2.4)			(Ksi)	(231) (224) (232) (229)	(201)	(154) (147) (151) (151)	(132)	(Ms1)	(18.7) (20.1) (20.0) (19.6)	(17.2)	(19.0) (18.6) (19.2) (18.9)	(16.6)	(Ks1)	(17.3) (16.4) (16.6) (16.8)	(8.17) (8.18) (8.18) (8.15) (8.16)
	EX	329 (625) 1378 (200)	1.613 25.8 68.4 -0.40 2.11-19 (83 75-78 75-78 75-78	356 (673) 100 100		MN/m ²	1578	1385	1040	606	GN/m ²	132	119	130	511	MN/m ²	116	56.2
2	49		76.4-73.6) 1.(2.4-2.3			(Ks1)	(195) (207 (206) (203)	(201)	(102) (99) (109) (103)	(102)	(Ms1)	(19.1) (19.2) (20.2) (19.5)	(19.3)	$(17.1) \\ (18.1) \\ (17.4) \\ (17.5) \\ (17.5) $	(17.3)	(Ksi)	(12.4) (10.7) (10.7) (11.3)	(6.24) (5.92) (6.79) (6.32)
1041 1991	EX	329 (625) 345 (50)	1.49 1.49 34.0 6.16 6.37 2.0-1.87 (2.0-1.87 (73-78 73-78 0.10 685)	361 (682) 0 0		MN/m ²	. 1399	1385	710	703	GN/m ²	134	133	121	119	MN/m ²	779	43.5
	48		3.2-76.8) 9 (2.6-2.4)			(Ksi)	(211) (221) (222) (222)	(221)	(136) (124) (133)	(133)	(Nsi)	$(17.1) \\ (17.6) \\ (18.5) \\ (17.7) \\ (17.7)$	(17.6)	$(16.9) \\ (16.2) \\ (18.4) \\ (17.2) \\ (17.2)$	(17.1)	(Ks1)	$(15.3) \\ (14.9) \\ (15.2) \\ ($	(6.98) (6.88) (6.78) (6.88)
, , , , , , , , , , , , , , , , , , , ,	EX	329 (625) 689 (100)	1.54 1.54 60.2 0.51 2.1-1.9 (8: 2.1-1.9 (8: 74-76 0.062 339 (642)	346 (655) 99.5 99.5		MN/m ²	1530	1523	916	916	GN/m ²	122	121	119.	118	MN/m ²	105	47.4
->	47		9 (2.5-2.2)			(Ksi)	(237) (240) (232) (236)	(209)	(143) (158) (148) (149)	(132)	(Msi)	(18.8) (18.4) (18.5) (18.5)	(16.5)	(18.7) (19.8) (19.9) (19.5)	(17.2)	(Ks1)	(14.4) (13.9) (14.5) (14.5)	(7.17) (6.89) (6.94) (6.94)
	EX	329 (625) 1034 (150)	1.60 26.0 67.7 0.33 0.14 (80 2.0-1.8 (80 2.0-1.8 (80 0.14 0.14 34.5 (653)	342 (648) 100 100		MN/m ²	1626	1440	1027	910	GN/m ²	128	114	134	119	MN/m ²	100	47.8
	41		-70) (2.5-2.2)			(Ksi)	(214) (224) (220) (219)	(188)	(146) (145) (142) (142)	(122)	(Nsi)	(19.3) (19.2) (19.2) (19.2)	(16.4)	(18.8) (18.4) (18.5) (18.6)	(16.0)	(Ks1)	$(15.9) \\ (17.7) \\ (16.2) \\ (16.6)$	(6.93) (7.12) (7.31) (7.12)
0	EX	329 (625) 1378 (200)	1.618 24.4 62.7 -0.27 2.0-1.8 (80 2.0559 .0535-0559 .0320 (608)	331 (628) 100 100		MN/m ²	Avg 1509	1295	Avg 978	841	GN/m ²	Avg 132	113	Avg 128	110	MN/m ²	Avg 114	Avg 49.1
	et(5) berty		9 3.2-76.8) 9 (2.6-2.4)			(Ksi)		(>228)		(>136)	(isi)		(>18)		(>18)	(Ksi)	(>15)	(/<)
	Targ		1.561-1.57 35.0-31.3 58-62 58-62 22 22 21-1.9 (8 2060060 270 270	>340 >95 >95		MN/m ²		>1570		>942	GN/m ²		>124		>124	NN/m ²	>103	>48
	Properties Panel No.	Processing Variable(1) 1. Maximum cure temperature, C (F) 2. Maximum molding pressure, N/m ² (psi)	Composite Physical Properties(2) 1. Specific gravity (grams/cc) 2. Resin weight content (2) 3. Fiber volume (3) 4. Void Volume (3) 5. Thickness mm (mils) 6. Thickness/ply, mm (mils) 7. Barcol hardness (ASTM D5583) 8. Weight loss in postcure (2) 9. TMA-TB C, (F) cured	Postcured 4 hrs at 316 C (600 F) 10. C Scan ultra sound transmission (%)(3) Cured Postcured 4 hrs at 316 C (600 F)	Composite Mechanical Properties(4)	l. Flexural strength	RT	Avg normalized strength, 60% F/V	316 C (600 F)	Avg normalized strength, 60% F/V	2. Flexural modulus	RT	Avg normalized modulus, 60% F/V	316 C (600 F)	Avg normalized modulus, 60% F/V	3. Short beam shear strength	KI	316 C (600 F)

Minimum Molding Pressure and Temperature Investigations Table 7.

(1) luminates, 32 ply, unidirectional were autoclave molded using the two stage cycle. (2) Prepreg and composite physical properties were calculated per Appendix A. (3) NDI ultrasonfc through transmission tests were performed using the XMSN-LaKK established "A" sumsitivity stundards. (4) Specimens were tested after stabilisting at 316 C (600 F) for 10 minutes. (5) Fragrep roperty values are based on Celion fiber miniawa properties of 2018 $\rm KX/m^2$, (13 MSI) tensile modulus. (5) Fragrep roperty values are based on Celion fiber miniawa properties of 2018 $\rm KX/m^2$, (13 MSI) tensile modulus. (5) Fragrep roperty values are based on Celion fiber miniawa properties of 2018 $\rm KX/m^2$, (13 MSI) tensile modulus. (6) Fragrep roperty values are based on Celion fiber miniawa properties of 2018 $\rm KX/m^2$, (13 MSI) tensile modulus.

Properties Panal No.	FY	60	FX	70	FX	71	EX	72
Panel No.								
 Maximum cure temperature, C (F) Maximum molding pressure, N/m² (psi) 	316 (600) 1378 (200)		302 (575) 1378 (200)		288 (550) 1378 (200)		274 (525) 1378 (200)	
Composite Physical Properties(2)								
 Specific gravity (grams/cc) Resin weight content (%) Fiber volume (%) Void Volume (%) Thickness mm (mils) Thickness (ASTM D2583) Weight loss in postcure (%) TMA-Tg C, (F) cured Postcured 4 hrs at 316 C (600 F) C Scan ultra sound transmission (%)(3) 	1.612 26.2 67.9 -0.34 2.1-1.87 (7 .06090584 76-82 0.36 335 (635) 358 (676)	26.4-73.6) 6 (2.4-2.3)	1.594 28.5 65.1 -0.07 2.3-2.2 (89 0.7110685 75-80 0.43 317 (603) 353 (667)	.6-86.4) (2.8-2.7)	1.591 29.4 64.2 -0.16 2.1-2.0 (8: .0660063 76-81 0.39 305 (581) 355 (671)	3.2-80) 5 (2.6-2.5)	1.582 31.3 62.1 -0.20 2.2-2.1 (86 75-77 0.39 257 (495) 360 (680)	5.4-83.2) 5 (2.7-2.6)
Cured Postcured 4 hrs at 316 C (600 F)	100 100		100 100		100 100		99.5 99.5	
Composite Mechanical Properties(4)								
1. Flexural strength	MN/m ²	(Ksi)	MN/m ²	(Ksi)	MN/m ²	(Ksi)	MN/m ²	(Ksi)
_ RT	1702	(230) (254) <u>(256)</u> (247)	1516	(216) (218) (226) (220)	1530	(227) (216) (224) (222)	1440	(197) (217) (214) (209)
Avg normalized strength, 60% F/V	1502	(218)	1399	(203)	1426	(207)	1392	(202)
316 C (600 F)	1158	(173) (167) (164) (168) (1(6)	951	(136) (138) (141) (138) (197) (197)	1054	(154) (156) (151) (153) (1(2))	1054	(156) (145) (157) (153)
Avg normalized strength, 60% F/V	1020	(148)	8/5	(127)	985	(143)	1020	(140)
2. Flexural modulus RT	GN/m ²	(Ms1) (18.7) (20.2) (21.1) (20.0)	GN/m ²	(Msi) (17.2) (17.7) (17.8) (17.6)	GN/m ²	(Msi) (19.0) (18.1) (17.7) (18.3)	GN/m ²	(Ms1) (18.2) (18.5) (18.5) (18.4)
Avg normalized modulus, 60% F/V	122	(17.7)	112	(16.2)	118	(1/.1)	123	(17.8)
516 C (600 F)	139	(19.2) (20.3) (21.1) (20.2) (17.8)	117	(17.0) (17.1) (17.0) (17.0) (15.7)	127	(18.2) (18.8) (18.4) (18.5) (17.3)	125	(17.7) (18.4) (18.6) (18.2) (17.6)
Avg normalized modulus, 60% F/V	123	(17.8)	100	(15.7)		(17.5)	. 2	(17.0)
3. Short beam shear strength RT	MN/m ²	(Ks1) (16.6) (16.7) (15.4) (16.2)	MN/m ²	(Ks1) (16.9) (15.9) <u>(16.0</u> (16.3)	MN/m ²	(Ksi) (16.2) (16.9) <u>(17.5)</u> (16.9)	MN/m ² 99.9	(Ks1) (15.5) (14.4) (14.4) (14.5)
316 C (600 F)	57.2	(8.52) (8.36) (8.24) (8.37)	57.8	(8.77) (8.23) (8.17) (8.39)	51.3	(7.73) (7.84) (6.75) (7.44)	. 53.7	(8.04 (7.76) <u>(7.60)</u> (7.80)

Table 7. Minimum Molding Pressure and Temperature Investigations (Cont)

 $\ensuremath{^{(1)}}_{\text{Laminates, 32 ply, unidirectional were autoclave molded using the two stage cycle.}$

(2) Prepreg and composite physical properties were calculated per Appendix A.

(3) NDI ultrasonic through transmission tests were performed using the NASA-LaRC established "A" sensitivity standards. (4) Specimens were tested after stabilizing at 316 C (600 F) for 10 minutes.

(5) Target property values are based on Celion fiber minimum properties of 2618 MN/m², (34 MSI) tensile modulus, using the rule of mixtures, 60% composite fiber volume. Target 316 C (600 F) strength properties are based on a 60 percent retention of room temperature values.

Imidizing Temperature C (F)		Imid	izing Time	(Mins)	
163 (325 F)	60	90	120	150	180
177 (350 F)	60.	90	120	150	180
191 (375 F)	30	60	90	120	150
199 (390 F)	30	60	90	120	150
218 (425 F)	30	60	90	120	150

Table 8. Test Matrix - LARC 160 Imidizing and Cure Cycle Process Improvement Study

Total: 25-6 X 6 - 26 ply 0° onidirectional laminates.

Imidizing and Cure Cycle Process Improvement Study Observations -- Celion 3000/LARC 160 Composites⁽¹⁾ Table 9.

						_		—					·						-									
		Remarks	Corrugated depressions top and bottom laminate surfaces	parallel to fibers. Fiber wash and resin flash excessive	on panel sides	Surface depressions have	decreased, fiber wash and resin flash somewhat decreased	Corrugated depressions top and	bottom surfaces parallel to fibers. Fiber wash and resin	LIASH EXCESSIVE ON PAHEL SIGES Laminate surfaces smooth: same	fiber wash and resin flashing	011 STGES	Very minor top and bottom surface depressions and edge fiber washing	Top and bottom surfaces very	smooth and uniform, minor fiber			Top and bottom surfaces very	smooth and uniform, minor fiber washing	D			Top and bottom surfaces very	smooth and uniform, minor fiber	0	Top and bottom surfaces smooth	but appear resin rich due to lack of flow, minor fiber wash and resin flash	
	C-Scan Ultrasound Transmission	(%)	40	40	40 J	60	50 🕈	50	50	75	95	95 J	95	95 (2 splices)	95 (l splice)	95 (l splice)	100	(00T	100	T00	100	100 1	100	100	100	100	95	
tions	Barcol	(Cured)	72-75	72-75	72-76	72-75	68-73	72-74	72-75	73-76	74-78	73-77	72-77	74-78	74-77	75-79	72-77	12-27	74-76	73-75	73-76	70-76	74-77	72-76	74-76	73-76	72-77	
Observat	1 ess	(mils)	53-64	20-25	51-61	45-57	45-60	48-65	48-60	53-59	48-61	50-65	59-62	60-64	60-63	59-63	62-65	60-63	63-65	62-64	64-65	63-66	61-64	61-64	62-63	63-66	63-65	
Cure Cycle	Pane Thickn	шш	1.34-1.63	1.27-1.91	1.30-1.55	1.14-1.45	1.14-1.52	1.22-1.65	1.22-1.52	1.35-1.50	1.22-1.55	1.27-1.65	1.50-1.57	1.52-1.63	1.52-1.60	1.50-1.60	1.57-1.65	1,50-1,60	1.60-1.65	1.57-1.63	1.63-1.65	1.60-1.68	1.55-1.63	1.55-1.63	1.57-1.60	1.60-1.68	1.60-1.65	
	Void	(%)	4.61	4.20	3.64	3.20	2,21	4.59	4.42	3.74	1.91	2.01	0.58	0.19	-0.04	-0.04	-0.25	-0.19	-0.10	-0.76	0.18	-0.015	0.13	0.43	-0.011	-0.042	0.35	
	Fiber Vol	(%)	64.4	64.91	64.98	64.8	64.9	62.4	65.2	64.2	64.4	63.8	63.0	61.6	59.8	59.7	58.1	61.0	58.1	59.8	58.3	58.2	59.9	59.0	58.9	56.8	57.2	
	Resin	(%)	26.3	25.8	26.4	26.8	27.4	25.7	28.2	27.1	28.0	28.5	30.0	31.5	33.3	33.6	35.0	32.3	34.9	33.7	34.6	34.8	33.1	33.8	34.1	35.1	35.5	
	.	g/cc	1.530	1.531	1.545	1.550	1.563	1.536	1.521	1.540	1.565	1.561	1.576	1.575	1.570	1.574	1.565	1.577	1.563	1.579	I.560	1.562	1.568	1.560	1.565	1.550	1.553	
	Resin Flow Characteristics	in Cure	high	high	high	high	high	high	high	high	high	high	Low	med	med	med	med	med	med	Том	low	law	low	low	low	low	low	
vations	Resin	(%)	34.7	33.0	35.0	34.0	32.6	35.4	35.2	34.4	34.6	34.2	34.9	34.6	35.0	33.8	34.9	35.0	35.1	33.3	34.2	36.7	34.7	33.2	35.2	34.0	35.2	
) Observ	νοΊ	(%)	1.64	1.52	1.48	1.68	1.43	1.56	1.15	1.45	1.06	1.14	0.73	1.22	1.24	1.17	0.92	1.21	0.87	1.10	0.71	0.66	1.29	1.35	1.37	1.15	1.16	
; Cycle ⁽²	Imidize Time	(nin)	60	06	120	150	180	60	06	120	150	180	30	60	6	120	150	30	60	90	120	150	30	60	96	120	150	
Imidizing	Imidize Temp C	(F)	163 (325)					177	(005)				191 (375)		••			199	(065)				218	(425)				3

⁽¹¹⁾Celion 3000 fiber (epoxy resin sized) employed in making 30.4 cm (12.0 inches) wide, nominal 67 ± 3 grams/m² areal fiber weight prepreg. Prepreg. The resin solids: 38 $\pm 3\%$; volatiles: 12 $\pm 3\%$. (2) One ply 120 fiberglass top and 1 ply bottom surfaces used in imidizing and cure cycles to absorb excess resin to target 60 ± 2 composite fiber colume.

		T INTERIAMINAR SHFAR	able 10.	Test Matri	Lx Mechani	cal Prope	rties and	Structura	L Elements	I.REAM STIFFENED	HONEVCOME PANEL
		INTERLAMINAR SHEAR	FLEXURE	LONGITUDINAL TENSION	BEAM FLEXURE TENSION	BEAM FLEXURE Compression	COMPRESSION	HONEYCOMB FLATWISE TENSION	HAT STIFFENED Skin Stringer Panel	I-BEAM STIFFENED SKIN-STRINGER PANEL	HONEYCOMB PANEL
NOITATNAIRO RABIA	(A) D° ARUTARAMAT 1231	0.625	4 4 4 1 0.50 0.06 - 32:1	10,000 100 100 100 100 100 100 100 100 1							
		Fisu	() F _{fu} ,E _f	(1) F _{lu} . Et ∈ tu ^p	() F _{tu} Et ∉ _{tu}	(j) Feu Ec écu	() F _{cu} E _c [€] cu	<u>ک</u>	(2) 3) 4) Fc · ^c u	2 3 4 Fc · ⁶ cu	2 3 4
	-168 (-270)	9	9	۵	9	9	1	1	2	2	2
	24 (75)	6	ø	6	9	9	1	1	2	2	2
2	202 (400)	6	g	5	9	0	1	I	I	1	ŀ
L	316 (600)	Q	6	ø	ی	ø	ł	I	2	2	2
	-168 (-270)	1	1	9	1	1		1			
۱	24 (75)	1	1	ø	1	I	9	1			
۰	202 (400)	1	1	۵	1	l	6	1			
I	316 (600)	1	1	6	L	1	9	I			
	-168 (-270)	I	1	9	1	ł	Q	١			
	24 (75)	1	1	9	I	I	9	1			
- 	202 (400)	1	1	6	ł	I	9	1			
L	316 (600)	1	1	9	i	I	9	1			
	-168 (-270)	1	1	9	1	10	1	9			
 ; ;	24 (75)	1	1	۵	1	۵	ł	9			
n, c+∓.n	202 (400)	1	1	Q	I	9	1	9			
	316 (600)	1	I	59 .	1	ę	1	9			
	Θ	3 SPECIMENS POST-CU 1 SPECIMEN POST-CUR	JRED, 3 SPECIMENS AGEI (ED, 1 SPECIMEN AGED A	D AT 316 C (600F) FOR 1: It 316 C (600F) FOR 125	25 HOURS 3 FIBER Hours 4 desig	ORIENTATION IN LAMIN. N LOAD > 3000 LB/IN.	ATE AS REQUIRED BY S At room temperatur	PECIFIC DESIGN (REF 2ND IE) & 3RD QUARTERLY REP	ORTS)	

														_	
			JIMENSIONS: SAME AS		Flatwise	(1310) 1 (1520) (1540)	2 (1560) 2 (1660)	(1200) 3 (1290)	10,011 (1453)						
		16 CM 0 IN.)	POTTED WITH EA934 N	pon	Perpendicular	(6.10) (6.03) (7.67)	45.47 (6.6)			0.72 0.738 0.751	(0.736)				
re	The second secon		CM (N)	Core Rib	Specimen ID	10 11 12				10 11 12					
Honeycomb CO			CM 0 IN.)	load Direction to	Parallel	(16.73) (17.64) (19.89)	124.64 (18.09)			1.26 1.15 1.21	(1.21)				
unutun		0.38 CM (0.15 1	3.81 (1.5	Test I	Specimen ID	r 8 6				~ ≈ 6					
TA LIEU V.		см IN.)	ON-POTTED	bbon	Perpendícular	(7.56) (7.20) (6.59)	(8.95) (6.20)	50.30 (7.3)		0.69	08.0 (0.79)		(154.9) (146.7)	1.04 (150.8)	0.71 0.72 0.72
	r A		T S W W W	to Core Ri	Specimen ID	4 M 19	νø			0.044	n v				
2. LO TER (1			M.) (4.01	oad Direction t	Parallel	(13.6) (10.8) (11.20)	(11.88) (9.82)	78.96 (11.46)		1,15 1,15	(1, 20)		(254.0) (258.1)	1.76 (256.1)	1.08 1.12 1.10
			3.81 C (1.50	Test I	Specimen ID	4 31 1	φN			н - т - т - т	л ю П				
					Compressive Property	E _{cu} NN/m ² (Ksi)		(AVC		о д	(<u>Av</u> c	Fcu	E _{cu} MN/m ² (Psi)	(AV(µс (AVG
				Honevcomb	Core Density					352 Kg/m ³ (22 PCF) AL H/C			c	96 Kg/m ³	(6 PCF) AL H/C

÷.

-

RT Bulk Compressive Properties of 352 Kg/m² (22 PCF) 5052 Alloy 3.18 mm (0.125 fn.) Cell Aliminim Honevromh Core Table 11.

<pre>D) 5 Oriented Composite,</pre>	
n Unidirectional (n-Beam Test
of LARC-160/Celior	Postcured Condition
Tensile Properties	
Table 12.	

		Failure Mode	Tension, 2 inches both sides of G Tension on G Tension on G	Tension on G Tension, 1.5 inch off G Tension on G	Tension in center area Tension in center area Tension in center area	Steel face to core failure Tension on Q and steel face to core Steel face to core failure
	- + ; ;	с итг н (%)	$\frac{1.22 *}{1.16 *}$ $\frac{1.04 *}{1.14 *}$	$1.12 \\ 1.22 \\ 1.11 \\ 1.15 \\ $	$\frac{1.15}{1.02}\\\frac{1.07}{1.28}$	>0.816 0.970 >0.800 0.970
	ted ⁽²⁾	(Msi)	(26.50) (26.24) (27.44) (26.72)	(25.58) (26.04) (24.27) (25.30)	$\begin{array}{c} (23.61) \\ (25.95) \\ (22.31) \\ (23.17) \end{array}$	$\begin{array}{c} (25.45) \\ (25.44) \\ (25.44) \\ \hline (21.31) \\ \hline (24.07) \end{array}$
	Adjus	GN/m ²	184	<u>174</u>	160	<u>166</u>
Et	est	(țsW)	(28.28) (28.00) (29.29) (28.52)	(27.30) (27.80) (25.90) (27.00)	(25.20) (27.70) (23.8) (24.7)	(26.96) (26.95) (22.57) (25.49)
	Ĩ	GN/m ²	<u> 161</u>	<u>186</u>	<u>170</u>	<u>176</u>
	ted ⁽²⁾	(Ks1)	(321) (296) (282) (300)	(278) (304) (263) (282)	(274) (251) (243) (256)	>(206) (239) > <u>(163)</u> (239)
	Adjus	MN/m ²	2065	1941	<u>1763</u>	<u>1647</u>
-	st t	(Ksi)	(347) (319) (304) (323)	(300) (329) (284) (304)	(296) (271) (262) (276)	>(219) (255) >(174) (255)
Ft. Tes	Te;	MN/m ²	2228	2097	1076	<u>1757</u>
(3)	Test ⁽³⁾ Temp. C (F)		-168 (-270) Avg	RT Avg	204 (400) Avg	(316) (600) Avg
(4) Specimen Number		EX107T-4 -5 -6	EX107T-1 -2 -3	EX107T-7 -8 -9	EX107T-10 -11 -12	

(1) Tension critical beams per the design described. Aluminum honeycomb 5052 alloy core 1/8 cell, 352 g/m³ (22 pcf) density was employed in -168 C, RT and 204 C (-270 F, RT and 400 F) tests. CRES core, 301 alloy, 1/8 cell, 0.127 mm (0.005 inch) foil, 40 pcf density was employed in 316 C (600 F) tests.
(2) Adjusted properties were calculated from equations derived by Mr. Mark Shuart, NASA/LaRC using a computer program that considers the effect of bulk core properties on the strength and elastic modulus properties of the laminate. Adjustment factors

for -168 C, RT and 204 C(-270 F, RT and 400 F) test temperature: $F_{tu} = \frac{F_{tu}}{1.0792}$; $E_t = 0.937 \times E_t$ test value. For

FOT -100 V, $F_{tu} = \frac{F_{tu} \text{ test}}{1.0651}$; $E_t = 0.944 \times E_t$ test value. 316 C (600 F) tests: $F_{tu} = \frac{1.0651}{1.0651}$; $E_{t} = 0.944 \times E_t$ test value.

(3) Tests were performed at a load rate of 0.127 Cm/minute (0.05 inch/minute) after stabilizing at the test temperature for 10 minutes.

*Projected from point of strain gage failure.

	Properties:
•	Physical H
,	Composite

rties:
l Propei
Physica.
mposite

	C-Scan Transmission	100	
	ange	(mils)	(12-13)
kness	Actual R	mm	0.305-0.330
Thickr	ulated	(mils)	(10.4)
	Calc	шш	0.264
	Void	(%)	-0.65
•	Fiber Vol (%)		68.8
- -	Resin Content (%)		25.6
	Densitv	(grams/cc)	1,618

 $^{(5)}$ The insitu imidizing-cure cycle specified was employed in laminate fabrication.

Tensile Properties of LARC-160/Celion Unidirectional (0)5 Oriented Composite, Aged 125 Hours at 316 C (600 F)-Beam Test Table 13.

Failure Mode		Tensile failure on \mathfrak{C} Tensile failure on \mathfrak{C} Tensile failure on \mathfrak{C}	Tensile failure 1.0 inch off E Tensile failure on E Tensile failure on E	Tensile failure on C Tensile failure on C Bond failure specimen to core	Bond failure steel face/core Bond failure steel face/core Bond failure steel face/core			
	-	ε μτ ε μ	12.5 9.6 <u>12.3</u> 11.5	$13.3 \\ 13.3 \\ 14.1 \\ 13.6 \\ $	9.4 9.4 9.4	>4.5 >5.5 >6.4		
	ted(2)	(Msi)	$\begin{array}{c} (24.6) \\ (27.0) \\ (23.1) \\ (24.4) \end{array}$	$\begin{array}{c} (23.9) \\ (23.7) \\ (23.5) \\ (23.4) \\ (23.4) \end{array}$	(25.8) (27.8) (26.8)	$\begin{array}{c} (25.2) \\ (24.8) \\ (24.8) \\ (24.5) \\ (24.8) \\ (24.8) \end{array}$		
	su į bA	GN/m ²	172	161	185	<u>171</u>		
14	Test	st	st	(Wsi)	(26.3) (28.9) (24.7) (26.6)	$\begin{array}{c} (25.5) \\ (25.3) \\ (25.3) \\ (24.0) \\ (24.9) \end{array}$	(27.6) (29.7) (28.7)	$\begin{array}{c} (26.7) \\ (26.3) \\ (25.9) \\ (26.3) \\ (26.3) \end{array}$
		GN/m ²	184	172	*	181		
	ted (2)	(Ksi)	$\begin{array}{c} (260) \\ (227) \\ (227) \\ (251) \\ (251) \end{array}$	(294) (315) (307) (305)	$\begin{array}{c} (243) \\ (231) \\ (231) \\ (238) \\ (237) \end{array}$	(>120) (>135) (>156)		
'n	Adjusi	MN/m ²	1727	2104	1633	1		
Ľ,		(Ksi)	(281) (245 (286) (271)	$\begin{array}{c} (317) \\ (340) \\ (331) \\ (329) \end{array}$	(259) (249) (>257) (254)	(>128) (>144) (>166)		
	Te	MN/m ²	1867	2269	1750			
. (3)	Test ⁽³⁾ Temp. C (F)		-168 (-270) Avg	RT Avg	204 (400) Avg	(316) (600) Avg		
	Specimen(4) Number		EX199T-1 -2 -3	EN199T-4 -5 -6	EX199T-10 -11 -12	EN199T- 7 -8 -9		

 $(1)_{Tension}$ critical beams per the design described. Aluminum honeycomb 5052 alloy core, 1/8 cell, 352 g/m³ (22 pcf) density was employed in ~168 C, RT and 204 C (-270 F, RT and 400 F) tests. CRES core, 301 alloy, 1/8 cell, 0.127 mm (0.005 inch) foil, 40 pcf density was employed in 316 C (600 F) tests.

(2) Adjusted properties were calculated from equations derived by Mr. Mark Shuart, NASA /LaRC using a computer program that considers the effect of bulk core properties on the strength and elastic modulus properties of the laminate. Adjustment factors for -168 C, RT and 204 C (-270 F, RT and 400 F) test temperature:

 $F_{tu} = \frac{F}{1.0792}$; $E_t = 0.937 \times E_t$ test value. For 316 C (600 F) tests:

 $F_{tu} = \frac{F_{tu} \text{ test}}{1.0651}$; $F_{t} = 0.944 \text{ x} F_{t}$ test value.

(3) Tests were performed at a load rate of 0.127 Cn/minute (0.05 inch/minute) after stabilizing at the test temperature for 10 minutes. (4) Composite Physical Properties:

Wt Loss	Wt Loss After 125 Hours at 316 C (600 F) (5)				
	C-Scan Transmission	1101 estimation 11	66		
	Range	(mils)	(11-12.5)		
Thickness	Actual F	шш	0.279-0.318		
	lated	(mils)	(11.2)		
	Calcu	ШШ	0.284		
	Void	(%)	-0.22		
	Fiber		63.4		
	Resin Content (۵)				
	1000	(grams/cc)	1.601		

 $^{(5)}{\rm The}~2$ stage cure cycle specified was employed in laminate fabrication. *Strain gage failure. Compressive Properties of LARC-160/Celion Unidirectional (0)5 Oriented Composite Postcured Condition-Beam Test Table 14.

		Failure Mode	Compression on G Compression 1.0 inch off G Compression 2.0 inch both sides G	Compression overloading hole Compression on G Compression on G	Composite to core bond failure Compression on G No test damaged specimen	Compression on Q. Compression 1.5 inch off Q. Compression 1.75 inch off G						
		ε ult μ (%)	$1.42 \\ 1.34 \\ 1.51 \\ 1.42 \\ $	1.11 0.976 1.05 1.05	>0.472 0.960 0.960	0.610 0.660 0.660 0.643						
	ed (2)	(Msi)	(23.20) (23.12) (22.78) (23.03)	(20.93) (21.52) (20.80) (20.98)	(21.18) (20.42) (204)	$(19.71) \\ (21.00) \\ (20.95) \\ 20.55$						
	Adjust	GN/m2	<u>159</u>	<u>145</u>		141						
E	st	est	(Wsi)	(24.76) (24.67) (24.31) (24.58	(22.34) (22.97) (22.20) (22.50)	(22.6) (21.8) (21.8) (21.8)	$\begin{array}{c} (20.88) \\ (22.24) \\ (22.20) \\ (21.77) \\ (21.77) \end{array}$					
	Te	GN/m ²	164	<u>155</u>	150	150						
	ed(2)	(Ksi)	(273) (254) (262) (263)	(207) (193) (198) (199)	>(95.7) (170) (170) (170)	(114) (133) (136) (128)						
n	Test Adjust	Test Adjust	Test Adjust	Adjust	: Adjust	t Adjust	MN/m ²	<u>1812</u>	1373		(880)	
ъ Б								(Ksi)	(295) (274) (283) (284)	(223) (208) (214) (215)	>(102) (183) - (183)	$(121) \\ (142) \\ (145) \\ (136) \\ (136)$
				MN/m ²	<u> 1957</u>	1481	<u></u> 1261	937				
Tact (3)	Temp	E)	-168 (270) Avg	RT Avg	204 (400) Avg	316 (600) Avg						
	Creatman(4)	Number	EX107C-4 -5 -6	EX107C-1 -2 -3	EX10C-7 -8 -9	EX107C-10 -11 -12						

(1) Compression critical beams per the design described. Aluminum honeycomb 5052 alloy core 1/8 cell, 352 g/m³ (22 psf) density was employed in -168 C, RT & 204 C (-270 F, RT & 400 F) tests, CRES Core, 301 Alloy, 1/8 cell, 0.0127 mm (0.005 inch) foil 40 pcf was employed in 316 C (600 F) tests.

(2) Adjusted properties were calculated from equations derived by Mr. Mark Shuart, NASA/LaRC using a computer program that considers the effect of bulk core properties on the strength & elastic modulus properties of the laminate. Adjustment factors for -168 C, RT &

 $F_{cu} = \frac{F_{cu} \text{ test}}{1.0651}$; $E_c = 0.944 \text{ x } E_c$ test value.

(3)Tests were performed at a load rate of 0.127 cm/minutes (0.05 inch/minute) after stabilizing at the test temperature for 10 minutes. (4)Composite Physical Properties:

	23) 100		
	nge	(mi)	(12-2
hickness	Actual Rar	LE	0.305-0.330
T	Calculated	(mils)	(10.4)
		шu	0.264
	Void	(%)	-0.65
	68.8		
Resin Content (%)			25.6
	1.618		

 $^{(5)}$ The insitu imidizing-cure cycle specified was employed in laminate fabrication.

Table 15. Compressive Properties of LARC-160/Celion Unidirectional (0)5 Oriented Composite, Aged 125 Hours at 316 C (600 F)--Beam Test

		Failure Mode	Compression 1.5 inch off E Compression on E Compression on E		Compression on E Compression 1.25 inch off E Compression over loading hole		Compression 1.5 inch off & Compression 1.0 inch off & Compression 0.5 inch off &		Compression 1.0 inch off C Compression on E	Compression 1.0 mich 914
	*	ε ult μ (%)	 11.7 13.9	12.8	10.5 14.4 13.7	12.9	4.60	4.7	5.40	3.94
	ted(2)	(Msi)	(21.8) (22.1)	(22.0	(19.8) (19.4) (20.2)	(19.8)	(26.3)	(24.1)	(22.8)	$\frac{(22.9)}{(22.9)}$
	Adjust	GN/m ²	1	151		136	t	166		157
ш	st	(Wsi)	(23.6)	(23.5)	(21.1) (20.7) (21.6)	(21.1)	(28.1)	(25.8)	(24.2)	$\frac{(24.3)}{(24.3)}$
	Te	GN/m ²	*	162		145	*	177	*	167
	ced(2)	(Ksi)	(221) (263) (242)	(242)	(184) (245) (235)	(222)	(119) (114)	(104)	(119)	(83.7) (106)
-	Adiust	MN/m ²		1667		1529		774		727
Ц,		(Ksi)	(257) (284) (261)	(267)	(198) (264)	(072)	(128) (123)	$\frac{(112)}{(121)}$	(127) (122)	(113) (113)
	Ē	MN/m ²		1842		1653		834		777
	Test(3) Temp C (F)		-168 (-270)	Avg	КТ	Avg	204 (400)	Avg	316 (600)	Avg
	Specimen(4) Number		EX199C-1 -2) 	EX199C-4 -5	9	EX199C-10 -11	-12	EX199C-7 _8) 6

 $(1)_{Compression}$ critical beams per the design described. Aluminum honeycomb 5052 alloy core, 1/8 cell, 352 g/m³ (22 psf) density was employed in -168 C, RT and 204 C (-270 F, RT and 400 F) tests, CRES core, 301 alloy 1/8 cell, 0.127 mm (0.005 inch) foil 40 pcf was employed in 316 C (600 F) tests.

(2) Adjusted properties were calculated from equations derived by Mr. Mark Shuart, NASA/LaRC using a computer program that considers the effect of bulk core properties on the strength and elastic modulus properties of the laminate. Adjustment factors for -168 C, RT and 204 C (-270 F, RT and 400 F) test temperature:

 $F_{cu} = \frac{f_{cu} \text{ test}}{1.0792}$; $E_c = 0.937 \text{ x } E_c$ test value. For 316 C (600 F) tests:

 $F_{cu} = \frac{f_{cu} \text{ test}}{1.0651}; E_c = 0.944 \text{ x } E_c \text{ test value}.$

(3)_{Tests} were performed at a load rate of 0.127 cm/minutes (0.05 inch/minute) after stabilizing at the test temperature for 10 minutes. (4) Composite Physical Properties:

Wt Loss After 125	Wt Loss After 125 Hours at 316 C (600 F) (%)					
	C-Scan Transmission	(%)	66			
	Range	(mils)	(11-12.5)			
Thickness	Actual	шш	0.279-0.318			
	lated	(mils)	(11.2)			
	Calcu	Ē	0.284			
	Void	10A	-0.22			
	Fiber	Vol (%)	63.4			
	Res In	Content (%)	29.9			
		Density (grams/cc)	1.601			

 $^{(5)}{\rm The}$ 2 stage cure cycle specified was employed in laminate fabrication. *Strain gage failure

Compressive Properties of LARC-160/Celion (0, ±45, 90)S Oriented Composite, Postcured Condition-Beam Test Table 16.

		Failure Mode	Compression on & Compression overloading hole Compression in center	Compression overloading hole Compression on C Compression on C	Composite-to-core bond failure Composite-to-core bond failure Composite-to-core bond failure	Compression on C Composite-to-core bond failure [Tested in tension-tensile failure] Composite-to-core bond failure Compression on C	
	ε ult μ	(%)	1.73 1.14 1.44	$1.33 \\ 1.16 \\ 1.11 \\ 1.21 \\ 1.21$	>0.950 >1.10 >1.16	0.984 >0.906 [0.890] >1.00 1.10	
	ed ⁽²⁾	(msi)	(9.13) (8.83) (8.98)	$\begin{array}{c} (8.88) \\ (9.54) \\ (9.54) \\ \hline (8.31) \\ \hline (8.91) \end{array}$	(6.85) (7.61) (7.04) (7.17)	(7.22) (7.68) [9.04] (8.86) <u>9.06</u> (8.14)	
	Adjust	GN/m ²	61.87	61.39	49.38	[62.3]	
Е Е	st	(msi)	(10.43) (10.10) (10.27)	$\begin{array}{c} (10.15) \\ (10.90) \\ (9.50) \\ (10.18) \end{array}$	(7.83) (8.70) (8.04) (8.04) (8.19)	(7.99) (8.50) [10.00] (9.8) (9.05)	
	Te	GN/m ²	70.73	<u>70.16</u>	56.43	[68.9] 62.0	
	sted ⁽²⁾	(ksi)	(98.4) (116) (99.2) (105)	(97.5) (95.0) (79.9) (90.8)	>(57.20) >(68.75) >(72.08) >(66.01)	(59.1) >(62.0) [85.2] >(71.6) (82.0) 70.6	
cu	Adju	MN/m ²	720	626	455	[587] 486	
	st	(ksi)	(117) (138) (118) (124)	(116) (113) (95) (108)	>(68.1) >(81.8) >(85.73) (78.5)	(66.1) >(69.4) [95.3] >(80.1) (91.8) 79.0	
	Ĩ	MN/m ²	<u>857</u>	744	541	[657] 544	
Tost(3)	Test(3) Temp C (F)		-168 (-270) Ave	RT Ave	204 (400)	316 (600) Ave	U
	T (4)(5) Specimen Number		EX106C-4 -5 -6	EX106C-1 -2 -3	EX106C-7 -8 -9	EX106C-10 -11 -12 -12 -13	

(1) Compression critical beams per the design described. Aluminum honeycomb 5052 alloy core, 1/8 cell, 352 g/m³ (22 pcf) density was employed in -168 C, RT and 204 C (-270 F, RT and 400 F) tests. CRES core, 301 alloy, 1/8 cell, 0.127 mm (0.005 inch) foil, 40 pcf employed in 316 C (600 F) tests.

(2)Adjusted properties were calculated from equations derived by Mr. Mark Shuart, NASA/LaRC using a computer program that considers the effect of bulk core properties on the strength and elastic modulus properties of the laminate. Adjustment factors for -168 C,

RT and 204 C (-270 F, RT, and 400 F) test temperatures: $F_{cu} = \frac{F_{cu} \text{ test}}{1.1893}$, $E_{c} = 0.875 \text{ x} E_{c}$ test value. For 316 C (600 F) tests: $F_{cu} = \frac{F_{cu} \text{ test}}{1.1192}$; $E_{c} = 0.904 \text{ x} E_{c}$ test value.

(3)Tests were performed at a load rate of 0.127 cm/minute (0.05 inch/minute) after stabilizing at the test temperature for

10 minutes.
(4) Composite Physical Properties:

	TMA-Tg	(c)	362	
	C-SCAN Transmission	(%)	100	
	ange	(mils)	(18-20)	
nickness	Actual Ré	unu	0.457-0.503	
Ţ	lated	(mils)	(18.1)	
	Calcu	шш	0.461	
	Void	(%)	0.47	
	Fiber	(%)	62.9	
	Resin	30.2		
		1.588		

 $^{(5)}$ The insitu imidizing-cure cycle specified was employed in laminate fabrication.

¢

		Failure Mode	Compression over loading hole Compression over loading hole Compression 1.5 inch off &		Compression on & Compression outside loading hole Compression on &		Compression over loading hole Compression on E Compression over loading hole		Compression 1.25 inch off & Compression over loading hole Compression on &	
		ε ult μ (%)	9.44 9.80 9.62		10.00 14.00 13.90	C0.21	9.80 13.10	11.45	11.40	10.6
	ted(2)	(Msi)			(9.19) (7.44) (7.13)	(76.1)	(7.07) (7.11)	(2.09)	(8.02)	(7.84)
с Е	Adjust	GN/m ²				54.6	1	48.9	I	54.0
	st	(Wsi)	(9.60) (9.10) (9.35)		(10.50) (8.50) (8.48)	(9.16)	(8.08) (8.13)	(8.11)	(8.88)	(8.68)
	Te	GN/m ²	* 79			63.1	*	55.9	*	59.8
	ted(2)	(Ksi)	(103) (66.5) (69.4) (70.6)	(2.2.1)	(75.7) (85.9) (85.7)	(82.4)	(65.4) (55.8) (73.6)	(67.9)	(58.7) (72.2)	(64.2)
	Adjus	MN/m ²	0,12			568		447		442
H	Rt	(Ksi)	(122) (79.1) (82.5)	(1.40)	(90.1) (102) (102)	(0.86)	(77.8) (66.4) (87.5)	(77.2)	(65.7) (80.8)	(71.8)
	1 	MN/r1 ²		TCO		675		532		495
	Test(3)	E)	-168 (-270)	AVB	RT	Avg	204 (400)	Avg	316 (600)	Avg
		Specimen(4)(5) Number	EX200C-1 -2 -3		EX200C-4 -5 -6		EX200C-10 -11 -12	4	EX200C-7 -8	61

(1) Compression critical beams per the design described. Aluminum honeycomb 5052 alloy core, 1/8 cell, 352 g/m³ (22 pcf) density was employed in -168 C, RT and 204 C (-270 F, RT and 400 F) tests. CRES core, 301 alloy, 1/8 cell, 0.127 mm (0.005 inch) foil, 40 pcf employed in 316 C (600 F) tests.

(2) Adjusted properties were calculated from equations derived by Mr. Mark Shuart, NASA/LaRC using a computer program that considers the effect of bulk core properties on the strength and elastic modulus properties of the laminate. Adjustment factors for -168 C, RT and 204 C (-270 F, RT, and 400 F) test temperatures:

 $F_{cu} = \frac{F_{cutest}}{1.1893}$, $E_{c} = 0.875 \times E_{c}$ test value. For 316 C (600 F) tests:

 $F_{cu} = \frac{F_{cutest}}{1.1192}; E_{c} = 0.304 \times E_{c} \text{ test value.}$

(3) Tests were performed at a load rate of 0.127 cm/minute (0.05 inch/minute) after stabilizing at the test temperature for 10 minutes. (4) Composite Physical Properties:

					ŢŢ.	ickness				llt Ince ∆ftar	
	Resin	Fiber	Void	Calcu	ulated	Actual	Range	C-Scan Transmission	TMA-Te	125 Hrs at 316 C (600 F)	
~	Content (%)	Vol (%)	107 (%)	ЩЩ	(mils)	ш	(mils)	(%)	(C)	(%)	
+ -	31.2	62.0	-0.21	0.465	(18.32)	0.48-0.54	(19-21.5)	100	365	3.2	-

 $^{\rm (5)}{\rm The}$ 2 stage cure cycle specified was employed in laminate fabrication. *Strain gage failure.

Table 18. Summary of LARC-160/Celion Tensile Properties

						Test Temp	erature			
Panel	Fiher			Postci	ıred		Aged 12	5 Hours	at 316 C	(600 F)
No. & Spec.	Orientation/ Specimen	Tensile Property(1)	-168 C (-270 F)	RT	204 C (400 F)	316 C (600 F)	-168 C (-270 F)	RT	204 C (400 F)	316 C (600 F)
EX107	(0) ^t /	$F_{tu} MN/m^2$	2065	1941	1076	1647	1727	2104	1633	1
(P.C.)	Beam	(Ksi)	(300)	(282)	(276)	(239)	(251)	(305)	(237)	ł
EX199		E _t GN/m 2	184	174	160	166	172	161	185	171
(Aged)		(Ws1)	(26.72)	(25.30)	(23.7)	(24.07)	(24.4)	(23.4)	(26.8)	24.8
		e ult µ (%)	1.14	1.15	1.28	0.970	1.51	1.36	0.94	ł
		v	0.370	0.275	0.310	0.310	1	1	ł	1
EX98	(90)+/	$F_{tu} MN/m^2$	35.6	23.0	15.8	18.1	47.3	35.1	12.1	14.0
(.p.g.)	Coupon	(Ks1)	(5.17)	(3.34)	(2.30)	(2.63)	(6.87)	(01.2)	(1.75)	(2.03)
EX201		Et GN/m ²	11.02	9.20	8.20	5.23	TBD	TBD	TBD	TBD
(Aged)		(Msi)	1.60	1.60	(1.19)	0.759	TBD	TBD	TBD	TBD
_		ε ULT μ (%)	0.33	0.33	0.20	0.37	TBD	TBD	TBD	TBD
-		ν	0.068	0.068	0.041	0.031	TBD	TBD	TBD	TBD
EX105	(+45) _S /	$F_{tu} MN/m^2$	201	201	149	141	15.7	134	132	103
(P.C.)	Coupon	(Ksi)	29.1	29.1	(21.6)	(20.5)	(22.8)	(19.4)	(1.91)	(14.9)
EX202		$E_{t} GN/m^{2}$	27.69	27.69	20.26	17.71	TBD	TBD	TBD	TBD
(Aged)		(Msi)	(4.02)	(4.02)	(2.94)	(2.57)	TBD	TBD	TBD	TBD
		€ ULT µ (%)	0.74	0.74	ł	1	TBD	TBD	TBD	TBD
		A	0.75	0.75	0.84	0.92	TBD	TBD	TBD	TBD
EX106	(0, 14 5,90) _S	$F_{tu} MN/m^2$	517	569	556	560	480	446	434	164
(P.C.)	Coupon	(Ksi)	(6.47)	(82.5)	(80.7)	(81.3)	(9.6)	(64.7)	(63.3)	(71.3)
EX200		$E_{t} GN/m^{2}$	56.03	53.28	55.14	41.96	TBD	TBD	49.61	TBD
(Aged)		(Wsi)	(8.13)	(7.73)	(8.00)	(60.9)	TBD	TBD	(7.2)	TBD
		€ ULT µ (%)	0.96	1.10	1.02	0.89	TBD	TBD	TBD	TBD
		A	0.320	0.295	0.325	0.300	TBD	TBD	TBD	TBD

(1) u Poissons ratio values reported for tension beam specimens were calculated from coupon specimen data.

ı İ

324

Specimen (4)	Test (3) Temperature	F _{tu}	(5)	Et			εÜtu
Number	(F)	MN/m ²	(Ksi)	GN/m ²	(MSI)	νt	(%)
EX107-2-1	-168	1626	(236)		(24.5)		0.96A
-2-2	(-270)	1605	(233)		(22.5)	0.390	1.08 ¥
-2-3		1743	(253)		(21.6)	0.350	<u>1.16</u> *
				Avg 158	(22.2)	0.370	
EX107-1-1		1709	(248)		(23.0)	_	1.05 A
-1-2	RT	9 50	(137)		(18.4)	0.260	0.75 *
-1-3		1337	(194)		(21.4)	0.290	0.89 *
				Avg 145	(21.1)	0.275	
EX107-3-1	204	1633	(232)		(21.7)	_	1.17A
- 3- 2	(400)	1578	(229)		(21.6)	0.330	0.98*
- 3- 3		1357	(197)		(24.4)	0.290	<u>0.81</u> *
				Avg 156	(22.6)	0.310	
EX107-4-1	316	1357	(197)		(21.3)		0.90A
-4-2	(600)	1240	(180)		_		
-4-3		1433	(208)	· · · ·	(24.0)	<u>0.310</u>	0.87*
				Avg 156	(22.7)	0.310	

Table 19. Tensile Properties of LARC-160/Celion (0)₅ Oriented Postcured Composites(1)(2)

(1) Coupon tensile specimen design, straight sides, 2.54 cm (1.00 inch) wide. Laminate consisted of 5 ply 0° oriented, nominal 0.064 cm (2.5 mils) per ply.

(2) Composite Physical Properties:

	Resin	Fiber	Void		T	hickness		C-Scap
Density	Content	Vol	Vol	Calcu	ulated	Actual R	ange	Transmission
(grams/cc)	(%)	(%)	(%)	mm	(mils)	mm	(mils)	(%)
1.618	25.6	68.8	-0.65	0.264	(10.4)	0.305-0.330	(12-13)	100

(3) Load was applied at 1.27 mm/minute (0.05 inch) after specimens had stabilized at test temperature for 10 minutes. Strain gaged specimens were loaded incrementally to allow (4) for data aquisition. -1 specimens were tested with a 5.08 cm (2.0 inch) gage length extensometer. Specimens

-2 and 3 tested at -168 C and 316 C (-270 F and 600 F) employed type WK-00-125AD-350 gages; -2 and 3 specimens tested at 204 C (400 F), type OK-00-125A-A-350 (LEN) gages; -2 and 3

(5) specimens tested at RT, type CEA-00-125UT-350 gages. Ftu data points were not averaged since strain gaged specimens were not tested under a constant loading condition.

* Projected from last strain gage reading

A Actual

Specimen ⁽⁴⁾	Test (3) Temperature C	F	tu	E	t		€ U]† µ
Number	(F)	MN/m ²	(Ksi)	GN/m ²	(MSI)	νt	(%)
EX199TC-1	-168		207		22.1		
-2	(-270)		182				
-3			-*			IRD	IBD
	AVG	1340	195				
EX199TC-4			192		22.8		
-5	RT		-**			-	TOD
-6			199			IRD	, IRD
	AVG	1347	196				
EX199TC-7	204		211		24.1		
-11	(400)		_**			TPD	TRD
-12			201			100	100
	AVG	1419	206				
EX199TC-8	316		_**				
-9	(600)		-**			TRD	TRD
-10			-**		20.8		
	AVG			l 		 	

Table 20. Tensile Properties of LARC-160/Celion (0), Oriented Composite, Aged 125 Hours at 316 C (600 F) (1) (2)

(1) Coupon tensile specimen design, straight sides, 2.54 cm (1.00 inch) wide. Laminate consisted of 5 ply 0° oriented, nominal 0.064 cm (2.5 mils) per ply.

(2) Composite Physical Properties:

					Thi	ckness			Wt Loss After
Density (aroms/cc)	Resin Content (%)	Fiber Vol (%)	Void Vol (%)	Calcu	lated	Actual	Range (mils)	C-Scan Transmission (%)	125 Hours at 316 C (600 F) %
	(%)	(%)	(%)	0.094	(11.0)	0.070.0.219	(11.10.5)	(%)	(000 F) %
1.601	29.9	63.4	-0.22	0.284	(11.2)	0.2/9-0.318	(11-12.5)	100	3.4

(3) Load was applied at 1.27 mm/minute (0.05 inch) after specimens had stabilized at test temperature for 10 minutes. Strain gaged specimen data aquisition was obtained autographically on two X, Y, Y recorders.

(4) -1 specimens were tested with a 5.08 cm (2.0 inch) gage length extensometer. Specimens -2 and 3 tested at -168 C and 316 C (-270 F and 600 F) employed type WK-00-125AD-350 gages; -2 and 3 specimens tested at 204 C (400 F), type 0K-00-125A-A-350 (LEN) gages; -2 and 3 specimens tested at RT, type CEA-00-125UT-350 gages.

* Damaged specimen

**Graphite/polyimide tab caused slipping in grips, damaged specimen.

Specimen ⁽⁴⁾	Test (3) Temperature C	F	(5)		Et		εUltμ
Number	(F)	MN/m ²	(Ksi)	GN/m ²	(MSI)	ν _t	(%)
EX98-2-1	-168	24.05	(3.49)	-	(1.63)	_	0.21 A
-2-2	(-270)	47.20	(6.85)		(1.57)	0.068	0.45*
-2-3			-				
		L		Avg 11.02	(1.60)	0.068	
EX98-1-1		24.87	(3.61)		(1.26)	_	0.28 A
-1-2	RT	-	-		-		
-1-3		21.15	(3.07)		<u>(1.41)</u>	0.051	<u>0.22*</u>
				Avg 9.20	(1.34)	0.051	
EX98-3-1	204	15.16	(2.20)		(1.11)		0.20 A
-3-2	(400)	15.23	(2.21)		(1.28)	0.032	0.17*
-3-3		17.16	(2.49)		(1.18)	0.049	0.22*
				Avg 8.20	(1.19)	0.041	
EX98-4-1	316	24.87	(3.61)		(0.883)	_	0.43A
-4-2	(600)	14.74	(2.14)		(0.740)	0.021	0.33*
-4-3		14.81	(2.15)		(0.653)	0.041	0.35*
				Avg 5.23	(0.759)	0. 0 31	

Table 21. Tensile Properties of LARC-160/Celion (90)40 Oriented Postcured Composites(1) (2)

(1) Coupon tensile specimen design straight sides, 2.54 cm (1.0 inch) wide. Laminate consisted of 40 ply 90° oriented, nominal 0.064 cm (2.5 mils) per ply.

(2) Composite Physical Properties:

Density	Resin	Fiber	Void Vol	Calc	<u>Thi</u> ulated	ckness Actual	Range	C-Scan Transmission	TMA-Tg
(grams/cc)	(%)	(%)	(%)	mm	(mils)	mm	(mils)	(%)	(C)
1.598	28.4	65.4	-0.29	2.21	86.7	1.80-2.05	75-81	100	340

(3) Load was applied at 1.27 mm/minute (0.05 inch) after specimens had stabilized at test temperature for 10 minutes. Strain gaged specimens were loaded incrementally to allow for ,,, data aquisition.

- (4) data aquisition.
 -1 specimens were tested with a 5.08 cm (2.0 inch) gage length extensometer. Specimens
 -2 and 3 tested at -168 C and 316 C (-270 F and 600 F) employed type WK-00-125AD-350 gages;
 -2 and 3 specimens tested at 204 C (400 F), type 0K-00-125a-a-350 (LEN) gages; -2 and 3
- (5) specimens tested at RT, type CEA-00-125UT-350 gages.
 (5) Ftu data points were not averaged since strain gaged specimens were not tested under a constant loading condition.
- * Projected from last strain gage reading
- A Actual

(4)	Test (3) Temperature	Ft	u	E	t		c III t u
Number	(F)	MN/m ²	(Ksi)	GN/m ²	(MS1)	νt	(%)
EX201TC-1	-168		6.90			-	
-2	(-270)		6.85	TBD	TBD	TBD	TBD
-3			*				
	Avg	47.3	6.87				
EX201TC-4			4.80	9.64	1.4		
-5	RT		5.40	TBD	TBD	TBD	TBD
-6			5.10				
	Avg	35.1	5.10				
EX201TC-7	204		1.90	7.57	1.1		
-10	(400)		1.60	TBD	TBD	TBD	TBD
	Avg	12.1	1.75				
EX201-8	316		1.80				
-9	(600)		2.30	TBD	TBD	TBD	TBD
-11			2.00				
	Avg	14.0	2.03				

Table 22. Tensile Properties of LARC-160/Celion (90)40 Oriented Composite Aged 125 Hours at 316 C (600 F) (1) (2)

(1) Coupon tensile specimen design straight sides, 2.54 cm (1.0 inch) wide. Laminate consisted of 40 ply 90° oriented, nominal 0.064 cm (2.5 mils) per ply.

(2) Composite Physical Properties:

					Th	ickness				Wt/ Loss After
Density	Resin Content	Fiber Vol	Void Vol (%)	Calc	ulated	Actual	Range	C-Scan Transmission (%)	TMA-Tg	125 Hours at 316C (6005) %
(grams/cc)	30.0	(%)	(%) -0.45	mm 2.27	(89.2)	mm 2.48-2.57	(98-101)	100	359	0.86

(3) Load was applied at 1.27 mm/minute (0.05 inch) after specimens had stabilized at test temperature for 10 minutes. Strain gaged specimen data aquisition was obtained autographically using X, Y, Y recorders.

(4) -1 specimens were tested with a 5.08 cm (2.0 inch) gage length extensometer. Specimens -2 and 3 tested at -168 C and 316 C (-270 F and 600 F) employed type WK-00-125AD-350 gages; -2 and 3 specimens tested at 204 C (400 F), type OK-00-125a-a-350 (LEN) gages; -2 and 3 specimens tested at RT, type CEA-00-125UT-350 gages.

* Damaged specimen.

(4) Specimen	Test (3) Temperature C	F,	(5)	1	² t		εUltμ
Number	(F)	MN/m ²	(Ksi)	GN/m ²	(MSI)	ν _t	(%)
EX105-2-1	-168	200	(29.1)		(3.58)	—	_
-2-2	(-270)	209	(30.4)		(4.30)	0.72	0.74*
-2-3		192	(27.9)		(4.17)	0.77	0.70*
			Avg	27.69	(4.02)	0.75	
EX105-1-1		176	(25.5)		(3.10)		_
-1-2	RT	163	(23.7)		(3.48)	0.77	-
-1-3		168	(24.7)		(3.22)	0.74	
	·		Avg	22.25	(3.23)	0.76	
EX105-3-1	204	164	(23.8)		(2.53)		
-3-2	(400)	149	(21.6)		(3.15)	0.75	—
-3-3		135	(19.6)		(3.15)	0.93	
			Avş	20.26	(2.94)	0.84	
EX105-4-1	316	150	(21.8)		(2.15)		
-4-2	(600)	130	(18.9)		(2.22)	0.93	—
-4-3		143	(20.7)		(3.33)	0.91	
[Avg	, 17.71	(2.57)	0.92	

Table 23. Tensile Properties of LARC-160/Celion (+45)_S Oriented Postcured Composites(1) (2)

(1) Coupon tensile specimen design, straight sides, 2.54 cm (1.0 inch) wide. Laminate consisted of 4 ply (+45)_S oriented, nominal 0.064 cm (2.5 mils) per ply.

(2) Composite Physical Properties:

Density	Resin Content	Fiber Vol	Void Vol	Calcu	Thio lated	ckness Actual Ra	nge	C-Scan Transmission
(grams/cc)	(%)	(%)	(%)	mm	(mils)	mm	(mils)	(%)
1.592	29.0	64.6	-0.10	0.223	8.88	0.229-0.254	9-10	100

(3) Load was applied at 1.27 mm/minute (0.05 inches) after specimens had stabilized at test temperature for 10 minutes. Strain gaged specimens were loaded incrementally to allow for

(4) data aquisition. -1 specimens were tested with a 5.08 cm (2.0 inch) gage length extensometer. Specimens -2 and 3 tested at -168 C and 316 C (-270 F and 600 F) employed type WK-00-125AD-350 gages; -2 and 3 specimens tested at 204 C (400 F), type OK-00-125A-A-350 (LEN) gages; -2 and 3

(5) specimens tested at RT, type CEA-00-125UT-350 gages. Ftu data points were not averaged since strain gaged specimens were not tested under a constant loading condition.

*Projected from last strain gage reading.

Spacimen ⁽⁴⁾	Test (3) Temperature	F	tu	E	t		e III t u
Number	(F)	MN/m ²	(Ksi)	GN/m ²	(MSI)	νt	(%)
EX202TC-1	-168		(23.1)	8.27	1.2		
-2	(-270)		(21.3)	TBD	TBD	TBD	TBD
-3			(24.0)				
	Avg	157	(22.8)		·		
EX202TC-4			(17.6)	14.5	2.1		
-5	RT		(18.7)	TBD	TBD	TBD	TBD
-6			(21.9)				
	Avg	134	(19.4)				
EX202TC-7	204		(19.2)	16.5	2.4		
-11	(400)		-*	TBD	TBD	TBD	TBD
-12			(19.0)				
	Avg	132	(19.1)				
EX202-8	316		(14.0)				
-9	(600)		(15.0)	TBD	TBD	TBD	TBD
-10			(15.7)	12.4	1.8		
	Avg	103	(14.9)				

Table 24. Tensile Properties of LARC-160/Celion $(\pm 45)_S$ Oriented Composite, Aged 125 Hours at 316 C (600 F) (1) (2)

 $^{(1)}$ Coupon tensile specimen design, straight sides, 2.54 cm (1.0 inch) wide. Laminate consisted of 4 ply (+ 45) $_{\rm S}$ oriented, nominal 0.064 cm (2.5 mils) per ply.

(2) Composite Physical Properties:

					Th	ickness			Wt. Loss After
Density	Resin	Fiber Vol	Void Vol	Calcu	lated	Actual Ra	nge	C-Scan Transmission	125 Hours at 316 C
(grams/cc)	(%)	(%)	(%)	mm	(mils)	mm	(mils)	(%)	(600 F) %
1.580	33.6	59.26	-0.10	0.238	(9.36)	0.203-0.254	(8-10)	100	5.2

(3) Load was applied at 1.27 mm/minute (0.05 inches) after specimens had stabilized at test temperature for 10 minutes. Strain gaged specimen data acquisition was obtained autographically using two X, Y, Y recorders

(4)
 -1 specimens were tested with a 5.08 cm (2.0 inch) gage length extensometer. Specimens -2 and 3 tested at -168 C and 316 C (-270 F and 600 F) employed type WK-00-125AD-350 gages; -2 and 3 specimens tested at 204 C (400 F), type 0K-00-125A-A-350 (LEN) gages; -2 and 3 specimens tested at RT, type CEA-00-125UT-350 gages.

* Damaged specimen

Specimen (4)	Test (3) Temperature C	F	(5)	F	t		εUltμ
Number	(F)	MN/m ²	(Ksi)	GN/m ²	(MSI)	Υ _t	(%)
EX106-2-1	-168	581	(84.3)		(8.67)		0.99A
-2-2	(-270)	393	(57.0)		(7.55)	0.310	0.79*
-2-3		576	(83.6)		(8.18)	0.330	1.11*
			Avg	56.03	(8.13)	0.320	
EX106-1-1		619	(89.9)		(7.55)	_	1.22*A
-1-2	RT	. 558	(81.0)		(7.76)	0.280	1.08*
-1-3		528	(76.7)		(7.89)	0.310	1.01*
			Avg	53.28	(7.73)	0.295	
EX106-3-1	204	634	(92.0)		(7.79)	_	1.20 A
-3-2	(400)	544	(79.0)		(8.04)	0.310	0.99*
- 3- 3		490	(71.1)		(8.18)	0.340	0.86*
			Avg	55.14	(8.00)	0.325	
EX106-4-1	316	604	(87.6)		(7.82)		0.90A
-4-2	(600)	573	(83.1)		(5.20)	C.320	0.96*
-4-3		504	(73.2)		(5.25)	0.280	<u>0.83*</u>
			Avg	41.96	6.09	0.300	

Table 25. Tensile Properties of LARC-160/Celion (0, +45, 90)s Oriented Postcured Composite(1) (2)

(1) Coupon tensile specimen design necked down test section. Laminate consisted of 8 ply (0, +45, 90)_S oriented, nominal 0.064 cm (2.5 mils) per ply.

(2) Composite Physical Properties:

Density	Resin Content	Fiber Vol	Void Vol	Calcu	Tlated	nickness Actual Ra	ange	C-Scan Transmission	TMA-Te
(grams/cc)	(%)	(%)	(%)	mm	(mils)	mm	(mils)	(%)	(C)
1.588	30.2	62.9	0.47	0.461	(18.1)	0.457-0.503	(18-20)	100	365

(3) Load was applied at 1.27 mm/minute (0.05 inch) after specimens had stabilized at test temperature for 10 minutes. Strain gaged specimens were loaded incrementally to allow for (4)^{data} aquisition. -1 specimens were tested with a 5.08 cm (2.0 inch) gage length extensometer. Specimens

-2 and 3 tested at -168 C and 316 C (-270 F and 600 F) employed type WK-00-125AD-350 gages;

-2 and 3 specimens tested at 204 C (400 F), type OK-00-125A-A-350 (LEN) gages; -2 and 3

(5) specimens tested at RT, type CEA-00-125UT-350 gages. Ftu data points were not averaged since strain gaged specimens were not tested under a constant loading condition.

* Projected from last strain gage reading

A Actual

Table 26. Tensile Properties of LARC-160/Celion (0, 145, 90)S Oriented Composite, Aged 125 Hours at 316 C (600 F) (1) (2)

	·						
(4)	(3) Test Temperature	F	tu	E	t		
Specimen '' Number	C (F)	MN/m ²	(Ksi)	GN/m ²	(MSI)	γ_{t}	ε01tμ (%)
EX200TC-1	-168		(71.2)	53.1	7.7		
-2	(-270)		(70.8)	TBD	TBD	TBD	TBD
- 3			(66.8)				
	Avg	480	(69.6)				
EX200TC-4			63.1	47.5	6.9		
-5	RT		70.9	TBD	TBD	TBD	TBD
-6			60.0				
	Avg	446	64.7				
EX200TC-7	204		63.3	49.6	7.2		
	(400)			TBD	TBD	TBD	TBD
	Avg	434	63.3		7.2		
EX200TC-8	316		70.9				
-9	(600)		68.1	TBD	TBD	TBD	TBD
-10			74.8	46.1	6.7		
	Avg	491	71.3				

⁽¹⁾Coupon tensile specimen design necked down test section. Laminate consisted of 8 ply $(0, \pm 45, 90)_S$ oriented, nominal 0.064 cm (2.5 mils) per ply.

(2) Composite Physical Properties:

	Pasin	Fibor	Void		Thi	ckness		(. See		Wt. Loss After
Density (grams/cc)	Content (%)	Vo1 (%)	Vo1 (%)	Calc mm	ulated (mils)	Actual mm	Range (mils)	Transmission (%)	TMA-Tg (C)	at 316 C (600 F) %
1.594	31.2	62.0	-0.21	0.465	(18.32)	0.48-0.54	(19-21.5)	100	365	3.2

⁽³⁾Load was applied at 1.27 mm/minute (0.05 inch) after specimens had stabilized at test temperature for 10 minutes. Strain gaged specimen data acquisition was obtained autographically on 2 X, Y, Y recorders.

(4) -1 specimens were tested with a 5.08 cm (2.0 inch) gage length extensometer. Specimens -2 and 3 tested at -168 C and 316 C (-270 F and 600 F) employed type 0K-00-125AD-350 gages; -2 and 3 specimens tested at 204 C (400 F), type)K-00-125A-A-350 (LEN) gages; -2 and 3 specimens tested at RT, type CEA-00-125UT-350 gages.

Table 27. Summary of LARC-160/Celion Compression Properties

316 C (600 F) 54.0 (7.84) (106) 157 (22.9) 0.47 (15.0) 7.19 (1.04) 1.83 125 (18.2) 12.7 (1.85) (64.2) (600 F) 1.06 727 103 442 316 C (177.7) 7.37 (1.07) 1.86 204 C (400 F) 138 (20.0) 14.1 (2.05) (64.9) 48.8 (7.09) 1.15 (24.1) 0.47 (112) 166 774 122 I 447 Aged 125 Hours at (22.8) 8.98 (1.30) 1.43 (222) 136 (19.8) 1.29 (22,8) 17.2 (2.50) 1.47 (7.92) 1.26 (82.4) 54.6 1529 RT 568 157 157 -168 C (-27°F) Test Temperature (242) 151 (22.0) (23.7) 11.3 (1.65) 1.64 208 (30.2) (3.02) 1.60 (79.6) 56.4 (8.18) 20.8 1.28 0.96 1667 548 163 (13.4) 5.79 (0.841) 2.80 (20.55) 0.643 (600 F) 316 C 486 (70.6) 56.0 (8.14) 1.07 63.5 (9.22) 10.1 (1.47) 2,27 (128) 141 92.3 880 >(66.01) 49.38 (7.17) >1.10 204 C (400 F) (201) 7.02 (1.02) 2.44 130 (18.9) 107 (1.55) 4.17 (170) 141 (20.4) 0.960 1171 138 >455 Postcured (199) 155 (20.98) 1.05 182 (26.4) 15.8 (2.29) 3.00 626 (90.8) 61.39 (8.91) 1.21 (25.4) 9.3 (1.35) 2.17 RI 1373 175 -168.C (-27°F) (263) 159 (23.03) 1.42 (33.5) 11.8 (1.71) 2.03 242 (35.1) 19.9 (2.90) 1.50 720 (105) 61.87 (8.98) 18.2 **1.**44 231 Compression Property ε ULT μ(%) E_{CU} MN/m² (Ksi) ε ULT μ(%) E_{CU} MN/m² (Ksi) ε ULT μ(%) ε ULT μ(%) ECU MN/m² E_C GN/m² (Msi) E_{CU} MN/m² E_C GN/m² (Msi) E_C GN/m² (Msi) Ec GN/m² (Ksi) (Ksi) (Msi) Orientation/ (0,±45,90)_S/ Specimen Fiber (±45)_S/ Coupon (90) t/ Coupon (0)t/ Beam Beam EX106 (P.C.) (P.C.) (P.C.) (P.C.) (Aged) (Aged) (Aged) (Paged) Panel EX199 EX105 EX200 EX201 EX107 No. EX98 220

Compressive Properties of $(90)_{AA}$ and $(+45)_{c}$ Oriented Fiber LARC-160/Celion Composites Table 28.

									2		, 									.	
Panel			-168°C	(-270	°F) (2)				RT				204°C	(400°F	(2)			316°C	(600°F	(2)	
No. / Orientation/ C	ondition	ч	n	ພິ	+s.	, ult (μ)	ч	ц	о Э	ω	ult (µ)	F	n	ы		ε ult (μ)	H	cu		F _C	: ult (µ)
(plies)	(1)	4N/m ²	(Ks1) (:N/m ²	(Ws1)	(%)	MN/m ²	(Ks1)	3N/m ²	(Wsi)	1 (%)	۸N/m ²	(Ksi)	GN/m ²	(Wsi)	(%)	MN/m ²	(Ksi)	GN/m ²	(Wsi)	(%)
89N.3			(31.2)		1.72)	1.86		(23.1)		1.35)	1.96		20.7		1.00	2.46		13.1		0.826	3.00
/(06)	O	_ <u> </u>	(35.6)	~	1.76)	2.13	_	(27.3)	~	1.39)	2.32		20.1		1.01	2.48		13.7		0.837	2.50
(07)			(33.8)		1.64)	2.11		(25.7)		1.32)	2.23		19.4	1	1.04	2.37		<u>13.3</u>		0.860	<u>2.90</u>
	Avg	231 ((33.5)	11.8	1.71	2.03	175	(25.4)	9.3 (1.35)	2.17	138	20.1	7.02	1.02	2.44	92.3	13.4	5.79	0.841	2.80
EX201			(25.2)		1.64)	1.68		(14.9)	Ĭ	1.29)	1.19		1			1		(14.2)		(1.06)	1.61
/ (06)	0	<u> </u>	(21.5)	<u> </u>	1.64)	1.66	_	(18.9)	<u> </u>	1.30)	1.55		1		1	1		(14.6)		(10.1)	1.89
(40)			(24.4)		1.67)	1.59		(19.8)		1.30)	1.55		(1-1)		1.07	1.36		(16.1)		(<u>1.06</u>)	2.01
	Avg	163 ((23.7)	11.3 (1.65)	1.64	157	(22.8)	8.98 (1.30)	1.43	122 ((17.7)	7.37	1.07	1.86	103	15.0	7.19	(1.04)	1.83
FX91			37.7	_	2.93	1.51		25.3		2.23	2.91		17.8		1.47	3.49		8.49		1.66	1.89
(+45) _S /	Θ		37.0		2.84	1.76		25.9		2.46	2.30		19.0		1.40	4.50		9.35		1.11	2.58
(32)			30.5		2.94	1.22		27.9		2.18	4.06		19.8		<u>1.78</u>	4.53		9.82		1.64	2.37
	Avg	242	35.1	19.9	2.90	1.50	182	26.4	15.8	2.29	3.00	130	18.9	107	1.55	4.17	63.5	9.22	10.1	1.47	2.28
EX220	 	Ť	(28.8)		3.18)	1.17		(22.1)		2.56)	1.41		18.6)		(2.07)	ł		(19.7)		(1.77)	
(+45)S/	0	~	(32.8)	~	2.80)	2.45		(23.6)	<u> </u>	2.52)	1.54		(20.3)		(2.07)	1		(16.7)		(1.73)	1
(32)	1		(1.62)		3.08)	1.18		(22.7)		2.48)	;		21.2)		(2.01)	1		(18.1)		(2.05)	+
	Avg	308	30.2	20.8 (3.02)	1.60	157	(22.8)	17.2	2.50	1.47	138	20.0	14.1	(2.05)		125	(18.2)	12.7	1.85	
Composite	Physical	Proper	ties (4		Tar	set proper	rties		EX96			E	X91			EX201			EX220		
1. Specific g	ravity (gı	ams/cc	;		1.561-	1.579		1.595			1.556	~			1.604			1.589			
2. Resin weig	ht content	(%)			35.0-3	1.3		28.4			34.7				30.0		_	32.5			<u>.</u> ,
3. Fiber volu	me (%)				58-62			65.4			58.1				63.4			59.4			
4. Void Volum	e (%)				<2			-0.25	-		0.29				0.45			0.90			
5. Thickness	mm (mils)							1.90-	2.05 ()	75-81)	2.00-	.2.13 (79-84)		2.48-2	27 (98-TC	(1)	1.98-2	.08 (78	-82)	
6. Thickness/	ply, mm (n	(ils)			0.0660	-0.0609 (2.6-2.4	1) 0.047	-0.051	(1.87-2.	0) 0.063	1-0.067	(2.47-	-2.63)	.062(64 (2.45-	2.52)	.062	065 (2.	43-2.56	
7. Barcol har	dness (AST	'M D258	(6)		>70			75-78			26-75	_			72-78			73-78			
8. Weight los	s in postc	"ure (%	C		<1			0.15			0.31				ł			00.73			_
9. Weight los	s after 12	5 hrs	at 316°	C (%)	7										0.86						
10. TMA-Tg C,	(F) cured				> 330 (626)		330 (626)		352 ((999)			ł			ł			
Postcui	red 4 hour	s at 3	16°C (6	00°F)	>340 ((779		340 ((779)		332 ((020)			ł			349 (6	(09		
Aged 1.	25 hours a	t 316°	C (600°	F)	>340 (644)					}				359 (6	(8)					
11. C-scan ult	ra sound t	ransmi	ssion ((E) (X																	
Cured					>95			100			100				100			100			
Postcul	red 4 hour	s at 3	.T6°C (6	00°F)	>95			100			100				100			100			

(1)Condition (1): Postcured 4 hours at 316°C (600°F); (2) aged 125 hours at 316°C, (600°F) for 125 hours.

(2) production (0.05 function) at test temperature for 10 minutes at a load rate of 1.27 mm (0.05 funch)/minute in the test fixture shown in the Third Quarterly Report. (3) fixture shown in the Third Quarterly Report. (3) MDI ultra sonic through transmission tests were performed using the NASA-LARC established "A" sensitivity standards.

 $\langle 4 \rangle$ Instructure cycle used for EX91 & E98 laminates; two stage cycle used for EX201 and E220 laminates.

Table 29. Flexural Properties of 0° Oriented Fiber LARC-160/Celion Laminates

	Ff	-108 C	(-270°F) E		Ff	ч _ л		J.J.	Here and the second sec	204 ° C		ų	Ff		(600°F)	1 E
on (1)	MN/m^2	(ksi)	GN/m ²	(msi)	MN/m^2	(ksi)	GN/m ²	(msi)	MN/m^2	(ksi)	GN/m ²	(ms1)	MN/m^2	(ksi)	GN/m ²	(msi)
Avg	- - 2033	(296) (293) (295) (295)	<u>126</u>	$(18.9) \\ (17.8) \\ (18.1) \\ (18.3) \\ (18.3)$	<u>1674</u>	(238) (248) (241) (241) (243)	<u>123</u>	$(17.3) \\ (18.3) \\ (18.0) \\ (18.0) \\ (17.9) \\ (17.9)$	1144	(143) (152) (203) (166)	<u>138</u>	$(19.7) \\ (21.4) \\ (21.4) \\ (19.1) \\ (20.1) \\ ($	<u>.</u> 992	(161) (137) (134) (144)	132	(18.4) (19.6) (19.6) (19.2)
Avg	1902	(304) (272) (255) (276)	<u>145</u>	(19.6) (21.6) (21.1) (21.1) (21.1)	<u>1785</u>	(254) (255) (267) (259)	<u>129</u>	$(18.0) \\ (19.1) \\ (19.0) \\ (18.7) \\ (18.7)$	1509	(210) (220) (227) (219)	132	(19.0) (20.1) (18.3) (19.1)	<u>1171</u>	$\begin{array}{c} (161) \\ (174) \\ (176) \\ (170) \\ (170) \end{array}$	<u>128</u>	$(17.7) \\ (18.7) \\ (18.7) \\ (19.4) \\ (18.6) \\ (18.6)$
nosi te	Physica	1 Prone	rties (4)] [aroot P				FX22				FX204		
224224		24211			' 	419cc -		3								T
cific g	ravity	(grams/	(cc)		1.56	1-1.579			l.581			н 	.608			
in weigh	ht cont	ent (%)			35.0	-31.3			30.4			́́л	0.4			
er volu	те (%)				58-6	2			62.2				3.7			
d volum	e (%)				× 2				0.87			ī 	0.83			
ckness 1	mm (mil	s)			1.71	6-1.583	(67.6-	62.4)	1.47-1	.62 (58	-64)		.37-1.54	t (54–6)	1)	
ckness/j	ply, mm	(mils)			0.06	60-0.06(39 (2.6 -	-2.4)	0.056-	0.062 (:	2.23-2.4	(9) 0	.059-0.0)53 (2.(38-2.35	~
col har	dness (.	ASTM D2	583)		> 70				72-78			-	3-74			
íght los:	s in po	stcure	(%)		< 1				0.22					ı		
lght los:	s after	125 hr	at 316'	C (°F)						ı			.03			
A-Tg°C,	(°F) c	ured			> 33	0 (626)				I				I		
Postcu: Aged 1	red 4 h 25 hour	ours at s at 314	316°C - 6°C (60((600°F))°F)	× 34 × 34	0 (644) 3 (644)			353 (6	- -		์ 	62 (689)	1		
scan ult	ra soun	d transı	mission	(%) (3)												
Cured					> 95				100			ה 	00			
Postcu	red 4 h	ours at	316°C	(600°F)	> 95				100			́н	00			

(2) Specimens were tested after stabilizing at test temperature for 10 minutes at a load rate of 1.27 mm (0.05 inch). (3) NDI ultra sonic through transmission tests were performed using the NASA-LaRC established "A" sensitivity standards.

 $^{(4)}$ The two stage cycle was employed in laminate fabrication.

Table 30. Short Beam Shear Properties of 0° Oriented Fiber LARC-160/Celion Laminates

(600 °F)	n	(ksi)	9.3 5.9	8.8 (9.2)	(8.9) (8.5)	$\frac{(8.1)}{(8.5)}$
316°C (Fs	MN/m ²		63.4		58.5
400°F)	n	(ksi)	(12.6) (12.8)	$\frac{(12.6)}{(12.6)}$	(12.8) (12.9)	$\frac{(12.0)}{(12.6)}$
204°C (ы	MN/m ²		86.8		86.8
		(ksi)	(17.1) (17.6)	(17.5) (17.4)	(17.9) (18.3)	$\frac{(17.7)}{(18.0)}$
RT	Fsu	MN/m ²		120		124
-270'F)	n	(ksi)	(0.61) (0.61)	(20.9) (19.9)	(21.1) (23.9)	<u>(22.9)</u> (22.6)
-168°C (с Ч	MN/m ²		137		156
		Condition (1)	Ô	Avg	Θ	Avg
	Panel No./	No. of Plies	EX204/ 26		EX225/ 26	

	Composite Physical Properties ⁽⁴⁾	Target Properties	EX225	EX204
-	Specific gravity (grams/cc)	1.561-1.579	1.581	1.608
2.	Resin weight content (%)	35.0-31.3	30.4	30.4
ъ.	Fiber volume (%)	58-62	62.2	63.2
4.	Void volume (Z)	¢1	0.87	-0.83
<u>ۍ</u>	Thickness mm (mils)	1.716-1.583 (67.6-624)	1.47-1.62 (58-64)	1.37-1.54 (54-61)
6 .	Thickness/ply, mm (mils)	0.0660-0.0609 (2.6-2.4)	0.059-0.062 (2.23-2.46)	0.059-0.053 (2.08-2.35)
7.	Barcol hardness (ASTM D2583)	>70	72-78	73-74
8.	Weight loss in postcure (%)	<1	0.22	ł
9.	Weight loss after 135 hours at 316°C (%)		I	1.03
10.	TMA-Tg C, (F) cured	>330 (626)	I	ł
	Postcured 4 hours at 316°C (600°F)	>340 (644)	353 (667)	
	Aged 125 hours at 316°C (600°F)	>340 (644)	1	365 (689)
11.	C-scan ultra sound transmission $(z)^{(3)}$			
	Cured	>95	100	100
	Postcured 4 hours at 316°C (600°F)	>95	100	100

(1) Condition ①: Postcured 4 hours at 316°C (600°F); ② aged 125 hours at 316°C (600°F)

(2) Specimens were tested after stabilizing at test temperature for 10 minutes at a load rate of 1.27 mm (0.05 inch)/minute.
(3) NDI ultrasonic through transmission tests were performed using the NASA-LaRC established "A" sensitivity standards.

4

١

 $^{(4)}_{\mathrm{The}}$ two state cycle specified was employed in laminate fabrication.

Table 31. Tensile and Flexural Properties (Average) of Celion/LARC-160 Chopped Unidirectional Tape Molding Compounds

			Tensile Pr	operties (1)	F	lexural Pr	operties (2)
			22 C (75 F)	316 C (600 F)	22 C	(75 F)	316 C	(600 F)
	1	2 	F _{tu} (3)	F _{tu} (3)	F _{fu} (3)	E f	F _{fu} (3)	н Н
Batch	Meight gm/m ²	riber Length cm(in.)	MN/m ² (ksi)	MN/m ² (ksi)	MN/m ² (ksi)	GN/m ² (Msi)	MN/m ² (ksi)	GN/m ² (Msi)
А	66.8	1.27 to 2.54 (0.5 to 1.0)	194 (28.1)	184 (26.7)	586 (85.1)	74.4 (10.8)	497 (72.2)	72.3 (10.5)
ф	153.7	1.27 (0.5)	116 (16.9)	89.6 (13.0)	245 (35.6)	37.9 (5.5)	196 (28.5)	35.1 (5.1)
U	60.4	1.27 (0.5)		7	484 (70.3)	73.7 (10.7)	278 (40.3)	55.8 (8.1)
Q	67	2.54 (1.0)			855 (124)	82.7 (12)	537 (77.9)	73 (10.6)

Parts were force (1) Individual tensile coupons per ASTM D 651 were net molded at 13.78 MN/m² (2,000 psi) 316 ± 5 C (600 ±10 F) for 1 hour. Pressure was applied when part reached 204 C (400 F). cooled under pressure to 66 C (150 F).

(600 ±10 F). Pressure was applied when part reached 204 C (400 F). Parts were force cooled under (2) Individual flexure coupons per ASTM D 790 were net molded at 13.78 MN/m² (2,000 psi) 316 ±5.5 C 66°C (150 F). pressure to

(3) Load was applied at 0.127 cm (0.05 inch)/minute after stabilizing at 316 C (600 F) for 10 ±5 minutes.

Table 32. Structural Element Weight Losses After Aging at 316 C (600 F) for 125 Hours.

Configuration	Specimen No.	Initial Wt. (grams)	Wt. Loss %
"HAT"	EX195-2A	349.1	1.43
	EX195-3A	348.0	1.39
	EX195-4A	343.8	1.45
"I"	• EX194-2A	399.6	1.25
	EX194-3A	398.2	1.36
	EX194-4A	395.1	1.27
Sandwich	EX241-2A	*	0.75
	EX241-3A	—	0.72
	EX241-4A	-	0.72

*Not comparable-doublers are bonded to panel ends.

Table 33. Structural Element Potting Materials and Processes.

Test Temperature	Potting Material		Process ⁽¹⁾
-168 C (-270 F)	Filled epoxy paste, EA934, Hysol Corporation	1.	Mix, pot & cure at R.T, 4 hours.
		2.	Post cure at 121 C (250 F) for 2 hours.
RT	Filled epoxy paste, EA911-11, Hysol Corporation	1.	Mix, pot and cure at R.T., 8 hours minimum
316 C (600 F)	Aluminum filled polyimide resin, BR 34B-18, American Cyanamid Corporation	1.	Modify base material by adding 35% - 0.8 mm (1/32 in) fiberglass milled fibers. Mix on paint shaker for 30 minutes minimum.
		2.	Pot specimen ends with Compound approximately 10.16 mm (0.40 inch) deep.
		3.	Place in oven and raise temperature R.T. to 177 C (350 F) at < 1.1 C (2 F)/min.
		4.	Raise temperature 177 C (350 F) to 316 C (600 F) at < 1.7 C (3 F)/min.
		5.	Post cure at 316 C (600 F) for 2 hours.

(1) Alignment of all specimens during cure was maintained within machined tolerance by clamping to right angle fixtures.

	Design Ult. Load, KN/cm (lbs/inch) Test Temperature C (F)							
Specimen	-168	RT	316 C					
Design	(-270 F)		(600 F)					
"Hat" Stringer	528	528	319					
	(3016)	(3016)	(1819)					
"I" Stringer	542	542	325					
	(3016)	(3096)	(1858)					
Sandwich	805	805	483					
	(4600)	(4600)	(2760)					

Table 34. Structural Element Target Loads

			Test Temperature		Ultimate Load		
Element Configuration	Condition(2)	No.	с	(F)	KN	(LBS)	Remarks
		EX109/EX110A	24	(75)	120.8	(27,150)	Achieved design ultimate, compressive failure of hat caps with transfer thru webs occurred during strain gage readout at 27,150 lbs. Skin and bond failures were secondary.
		EX109/EX110B	24	(75)	120.8	(27,150)	Achieved design ultimate - no failure. Specimen was then fatigue tested 5% to 67% of design ultimate, compression/compression load to 265,000 cycles - no failure.
ATVAT VIATVA		EX109/EX110B	-168	(~270)	116.8	(26,250)	Skin compression failure - load dropped to 19,500 lbs. Retest of EX109/EX110B specimen tested at RT. Failed at 97% of design ultimate.
STRAIN GAGES 1 AND 2 INSTALLED ON LOWER SKIN DIRECTLY OPPOSITE 3 AND 4		EX195-1 PC	316	(600)	87.84	(19,750)	Skin compression failure, bottom 2 corners followed by buckling through bottom-center. Minor bond failure, center stringer under skin buckle. 73% of RT design ultimate.
		EX195-2A	24	(75)	120.8	(27,150)	Achieved design ultimate - no failure.
	(2)	EX195-4A	-168	(-270)	120.8	(27,150)	Achieved design ultimate - no failure.
		EX195-3A	316	(600)	124.3	(27,950)	Skin compression failure inboard of 1 bortom corner followed by diagonal skin buckling toward center of panel. Minor debond under center stringer. Failed at 103% of RT design ultimate.
		EX111/EX113	24	(75)	125.4	(28,187)	Achieved design ultimate - no failure.
		EX111/EX113	-168	(~270)	125.4	(28,187)	Achieved design ultimate - no failure. Retest of EXIII/EXII3 tested at RT.
7/3/7/	1	EX111/EX113	24	(75)	125.4	(28,200)	Achieved design ultimate - no failure. Retest of EX111/EX113 tested at RT and -270 F.
<u>DENERSE</u>		EX194-1 PC	316	(600)	125.4	(28,187)	Achieved design ultimate - minor skin compression failure in 1 corner. Did not cause drop in load. No debonds.
STRAIN GAGES 3 AND 4 Installed on lower		EX194-2A	24	(75)	125.4	(28,187)	Achieved design ultimate - no failure.
SKIN UPPUSITE I AND 2	2	EX194-4A	-168	(-270)	125.7	(28,250)	Achieved design ultimate. Compression failure of skin starting at 1 upper corner extends inboard 1 inch. Two stringer caps and webs also failed in compression with caps splitting axially. No debonds.
		EX194-3A	316	(600)	125.4	(28,187)	Achieved design ultimate - no failure.
	_	EX150-1	24	(75)	125.7	(28,260)	Achieved required 0.53 MN/m (3000 lbs/inch) compression load. Skin compression failures occurred along edges of bottom doublers. No debonds.
STRAIN GAGES 4, 5, AND 6 INSTALLED ON LOWER SKIN OPPOSITE 1, 2, AND 3	1	EX241-1PC	-168	(-270)	160,1	(36,000)	Achieved 133% RT design ultimate. Compression failure both skins next to doubler.
		EX150-2	316	(600)	97.86	(22,000)	Skin compression 1 side only, top corner, 1.12 inch above doubler at edge, extends 3.0 inches inboard. Achieved 81% of RT requirement of 0.53 MN/m (3000 lbs/inch). No debonds.
		EX241-2A	24	(75)	136.3	(30,650)	Achieved 113% RT design ultimate. Compression failure both skins next to doubler; also core shear failure after skin failure due to instability.
	2	EX241-3A	-168	(-270)	156.1	(35,100)	Achieved130% RT design ultimate. Compression failure both skins 1.0 inch above doubler and next to doubler.
		EX241-4A	316	(600)	120.1	(27,000)	Achieved 100% RT design ultimate. Compression failure 1 skin next to doubler and 1 skin 1.0 inch above doubler.

Table 35. Results of Compression Tests on "Hat" and "I" Stiffened Skin and Sandwich Panel Structural Elements(1)

Condition 1: Postcured 4 hours at 316 C (600 F); 2 aged 125 hours at 316 C (600 F)

Physical Properties of Celion/LARC-160 Composite Laminates Table 36.

CL12C-9/ 12-(0,90)	-	1.560	35.2	57.1	0.65	344 (651)	100	100
CL12C-8/ 12-(0,90)		1.567	31.7	60.5	1.32	341 (646)	98	98
CL24C-7/ 24-(<u>+</u> 45,90)S		1.587	30.3	62.5	0.51	344 (651)	97	97
cL12C-6/ 12-(<u>+</u> 45,90) _S				_	_	348 (658)	66	66
cL6C-11/. 6-(<u>+</u> 45,90) _S		1.565	34.5	57.91	0.56	361 (682)	98	98
Target Property		1.561-1.579	35.0-31.3	58-62	<2	>340 (644)	>95	>95
Properties Panel No./No. Plies- Orientation	Composite Physical Properties(1)	1. Specific gravity (grams/cc)	2. Resin weight content (%)	3. Fiber volume (%)	4. Void Volume (%)	5. TMA-Tg C, (F) Postcured 4 hrs at 316 C (600 F)	6. C-Scan ultra sound transmission (%)	Postcured 4 hrs at 316 C (600 F)

Delivered to NASA-LaRC, Tasks (d) and (f)(1)

(1) Prepreg physical properties are as follows: Fiber areal weight: 127 grams/m²; calculated thickness/ply: 0.122 mm (4.80 mils); resin solids content: 37.4%; volatile content: 12.5%

1
APPENDIX A

This appendix contains three documents which were used during the course of the program to assure prepreg quality. Appendix Al presents the questionaire which was utilized as a preliminary screening device for potential prepreg suppliers. The flysheet attachment, Appendix A2, was submitted with each purchase order for prepreg materials. Appendix A3 presents test procedures and defines calculations which were imposed by the flysheet requirements.

APPENDIX A1

QUESTIONS

1. Have you ever produced graphite/LARC-160 prepregs?

What graphite filaments were utilized?

- a. HTS
- b. HTS-II
- c. Celion
- 3. Is your process proprietary?
- 4. As a producer of LARC-160 prepregs, how do you obtain the LARC-160 varnish?
 - a. Buy from supplier
 - b. Produce yourself
- 5. If you produce the LARC-160 varnish yourself, which method do you use to maintain accurate ratio control of BTDE, AP22 and NE?
 - a. By dispensing the initial amounts of BTDS, AP22 and NA in solid form for each batch
 - b. By the use of commercial alcohol solutions of BTDE and NE
 - c. By the use of alcohol solutions of BTDE and NE which you have produced yourself
 - d. Other
- 6. Do you use methyl or ethyl ester of BTDE and NE for your LARC-160 varnish?
 - a. Methyl ester
 - b, Ethyl ester
- 7. If you use the commercial LARC-160 varnish, what solids concentration do you buy?
 - a, 99% solids
 - b. 85% solids
 - c. 66% solids
 - d. Other explain why

- 8. In the production of your LARC-160 prepregs, do you use a hot melt process or a solvent process?
 - a. 99% solids (hot melt)
 - b. 85% solids
 - c. 66% solids
 - d. Others explain why
- 9. How do you assure high purity of the LARC-160 varnish for prepregging?
 - a. Use high purity ingredients
 - b. Store the LARC-160 alcohol solutions at a low temperature such as $40^{\rm O}F$ and for less than one month
 - c. Use the varnish in solids form (99%) only
 - d. Other
- 10. If you buy ingredients for preparation of the LARC-160 varnish, do you obtain certification on the assay of these ingredients from the suppliers?
 - a. Methyl and/or ethyl alcohol
 - b. BTDA, NA and AP22
 - c. BTDE and NE
- 11. What facilities do you have for producing LARC-160/Graphite Fiber Prepregs?
 - a. Hot melt coater
 - b. Solvent coating facility
 - c. Continuous fabric or tape coating facility
- 12. What type of LARC-160 prepregs can you produce?
 - a. Continuous unidirectional tape up to 12 inches wide
 - b. Continuous unidirectional tape up to 24 inches wide
 - c. Up to 50 inch wide fabric
 - d. Up to 60 inch wide fabric

- 13. What in-process quality control tests do you conduct on your LARC-160 prepregs?
 - a. Resin content
 - b. Volatiles

•

- c. Resin flow
- d. Other explain why
- 14. What quality control tests do you conduct on your finished prepreg?
 - a. Resin content
 - b. Volatiles content
 - c. Fiber content
 - d. Interlaminar shear test
 - e. Compression test
 - f. Gel time
 - g. Resin flow
 - h. Flex and flex modulus
 - i. Other explain why
- 15. What documentation do you provide on your prepreg materials?
 - a. Purchase orders and certification of all basic raw materials
 - b. Date of prepration and batch number of the monomer reactants
 - c. Specifications for prepregging and test methods
 - d. Date of prepregging and batch number of prepreg
- 16. Can you produce LARC-160 molding compounds?
 - a. Chopped roving
 - b. Chopped fabric
 - c. Other explain
- 17. Can you produce LARC-160 adhesives?

APPENDIX A2

CELION/LARC-160 PREPREG REQUIREMENTS

- 1. Graphite/polyimide prepreg tape, continuous, 6 to 12 inches wide, slit net. Supply to the following target requirements:
 - a. Resin type: LARC-160
 - b. Fiber type: Celion 6000, NR150-B2G sized
 - c. Resin solids (%): 37 + 3 3 = 3
 - d. Volatile content (%): 12 + 3 at 600° F, 30 minutes
 - e. Fiber areal weight $(gram/m^2)$: 153 + 3
 - f. Graphite fiber tensile strength (ksi: 400 min,)
- 2. Supplier to perform all resin mixing operations. Detailed mixing procedures to be supplied with material shipment,
- 3. Prepreg materials and required resin samples shall be packed with dry ice for shipment. All containers shall be plainly labeled to indicate dry ice shipping conditions.
- 4. Supply all available resin, resin precursor, and fiber physical properties certified by Celanese and batch numbers. Supplier will provide duplicate sample quantities of resin precursor materials, intermediate esters, and neat resin. One set of samples will be sent with each prepreg batch. The second set of samples will be sent directly to:

Science Center 1049 Camino Dos Rios Thousand Oaks, CA 91360

Attention: Paul J. Dynes, A12

Samples sent to the Science Center shall be labeled to note contents (e.g. analytical samples) and storage requirements.

Frequency of sample submission and sample size are as follows:

a. Precursor materials - 300 gm each of BTDA, NA, and AP22 and 1 liter of Fotocol or equivalent will be submitted with the resin batch. Where the precursors are from a lot used in formulating prior resin batches, these samples will not be required. However, the precursor lot number and prior resin batch number shall be noted.

- b. Intermediate esters and neat resin 10 gm each of the intermediate ester and neat resin will be submitted for each resin batch formulated. These samples will be shipped under temperature controlled conditions commensurate with those used for shipping the prepreg tape representative of the resin batch.
- 5. Test conditions and calculations for prepreg target requirements are contained in LTR 2433-4462 which is in supplier's possession.
- 6. Two (2) copies of the following will be furnished to Rockwell for each lot or batch of material processed:
 - a. Graphite Yarn Q.C. Summary (Celanese)
 - b. Tensile Strength and Young's Modulus of Graphite Fibers (Celanese)
 - c. Mix Order and Specifications
 - d. Collimated Tape Traceability
- 7. The prepreg tape batch will not be accepted by Rockwell unless samples of the intermediate ester and neat resin specified in 4(b) have been received by both the Space Systems Group (Downey) and the Science Center prior to prepreg shipment or unless these samples accompany the prepreg shipment to the Space Systems Group and evidence of sample submittal to the Science Center is documented.

Documentation items noted in 6 will be submitted no later than fifteen (15) days after prepreg shipment.

APPENDIX A3

Space Systems Group



Laboratory Test Report

LTR 2433-4462

GRAPHITE/POLYIMIDE PREPREG AND COMPOSITE PHYSICAL PROPERTY TESTING PROCEDURES

March 1978

Authored by

J. S. Jones Responsible Test Engineer

Approved by

ntyp

J. A. Leing

S. Kritzer, Śupervisor Non-Metallic Materials & Processes Laboratories and Test J. H. Diaz, Manager Materials & Processes Laboratories Laboratories and Test



PREPREG AND COMPOSITE CONSTITUENT CALCULATIONS * Ι FOR POLYIMIDE RESIN/GRAPHITE FIBER MATERIALS

The following procedures define calculations to be employed in establishing graphite/polyimide prepreg and composite constituents. Example calculations for a typical PMR15 I/HTS II prepreg and composite, 60% fiber volume are given.

1.0 Resin and Fiber Densities

1.1 Establish resin $\ell r = 1.32$ grams/cc Vendors Certification

1.2 Establish Fiber $\rho f = 1.65 \text{ grams/cc}$

2.0 Establish Theoretical Density of Composite, O Void Assumed

 $c = (VF_f \times cf) + (Vr_f \times cr)$

Where: (c = density of composite VF_f = desired fiber volume fraction (f = density of fiber, grams/cc Vr_f = desired resin volume fraction (r = density of resin, gram/cc Example: $lc = (.60 \times 1.65) + (.40 \times 1.32)$ (.990 + .528)lc = 1.518 grams/cc

3.0 Establish Resin Solids (%) and Fiber Weight (%) for a Desired 60% Composite Fiber Volume, O Void Content

R.S. (%) =
$$(Vr_f \times (r) + (Vf_f \times (f)) \times 100)$$

Where: R.S. (%) = resin solids weight (%)
 Vr_f = desired resin volume fraction
 Vf_f = desired fiber volume fraction
 $(r = actual resin density (g/cc))$
 $(f = actual fiber density (g/cc))$
 FW (%) = fiber weight (%)

Example: R.S. (%) = $\frac{(.40 \times 1.32)}{(.40 \times 1.32) + (.60 \times 1.65)} = \frac{.528}{1.518}$.3478 $.3478 \times 100 = 34.78\%$

$$FW(\%) = 100 - 34.78 = 65.22\%$$

* These procedures have been programmed on a Hewlett Packard computer, Model 9820A, to facilitate rapid data acquisition.





4.0 Establish Fiber Weight Percent of Prepreg

4.1 <u>Test Procedure</u>

- Prepare nominal 3-inch square specimens of prepreg. For handling convenience the specimen may be cut (before weighing) into several narrow strips. Remove release paper before analyzing.
- 2) Determine the area of each nominal 3 x 3 inch specimen to an accuracy of 0.01 square inch. Record as "AF"(Use in step 7).
- 3) Obtain a clean dry extraction thimble and weigh to the nearest 0.1 mg and record as W_1 .
 - Note: may be purchased from Van Waters & Rogers Company as fritted glass extraction thimbles, medium E.C. 35 x 90 mm, Catalog No. 27743-120.
- 4) Place specimen in thimble (step 2) and weigh to the nearest 0.1 mg. Record as W_2 .
- 5) Place thimble and specimen in beaker and add enough solvent to cover the extraction thimble, and let set at room temperature with intermittent agitation for 30 minutes. The solvent used shall be selected on the basis of being able to dissolve the resin completely under the conditions of the test. Normally methyl ethyl ketone is suitable.
- 6) Remove extraction thimbles with specimen from beaker of solvent and drain. Discard used solvent and rinse beaker with fresh solvent;
- 7) Place extraction thimbles inside beaker and cover with fresh solvent. Repeat steps 5 and 6 until solvent is visually clean and fibers stand apart.
- 8) Following the last extraction, remove extraction thimble with specimen from beaker of solvent and place in a rubber crucible holder on a vacuum filter flask capable of maintaining a vacuum of at least five inches of mercury. Drain free of solvent and rinse once more with fresh solvent.
- 9) Dry for 30 minutes at 300 320F in a mechanical convection oven or, alternatively, for 15 minutes at 130 150F in a vacuum oven.
- 10) Remove from oven and cool to room temperature in a desiccator.
- 11) Weigh each extraction thimble and specimen to nearest 0.1 mg.
- 12) Record test specimen weight as W₂.



4.2 Calculations and Report of Results

Calculate fiber content as follows:

Weight % fiber =
$$\frac{W_3 - W_1}{W_2 - W_1} \times 100$$

- Where: W_1 = weight of extraction thimble, grams W_2 = weight of extraction thimble plus specimen, in grams W_3 = weight of extraction thimble plus test sample after extraction, in grams
- Example: 33.20352 32.61985 = .58367 = .575233.63455 - 32.61985 = 1.01420

.5752 x 100 ₩ 57.52%

5.0 Establish Total Volatile Weight Percent of Prepreg

5.1 <u>Test Procedure</u>

- 1) Obtain clean, dry aluminum weighing dish (expendable).
- Prepare specimens of a size not to exceed four square inches. The sampling plan shall permit assessment of distributional uniformity within the prepreg tape.
 - Note: release paper must be removed prior to analyzing. Any resin adhering to the release paper will be lost to the test.
- 3) Weigh each aluminum dish to the nearest 0.1 mg and record as W_2 .
- 4) Place a test specimen in each dish and weigh to the nearest 0.1 mg. Record as W_1 .
- 5) Remove volatiles by heating the dish and the test specimens in a pre-heated air-circulating oven at 600F for 30 minutes.
- 6) Remove dish and test specimens from oven and cool to room temperature in a desiccator.
- 7) Weigh each dish and test specimens to the nearest 0.1 mg and record as W_3 .



5.2 Calculations and Report of Results

1) Calculate the volatiles content according to the following equation:

Volatiles content, percent by weight = $\frac{W_1 - W_3}{W_1 - W_2} \times 100$

- Where: W_1 = weight of dish plus test specimen in grams, before volatiles removal.
 - W_2 = weight of dish in grams
 - W₃ = weight of dish plus test specimen after volatiles removal
- 2) Report results in percent by weight to the nearest .01 percent.

Example: 10.4770 - 10.3461 = .1309 = .086510.4770 - 8.9642 = 1.5128

 $.0865 \times 100 = 8.65\%$

6.0 Establish Prepreg Tape Resin Solids Weight Percent

R.S. (%) = $\left(1.0 - \frac{FW_f}{1.0 - VW_f}\right)$ x 100 Where: R.S. (%) = resin solids in prepreg FW_f = fiber weight fraction of prepreg per step 4.0 VW_f = volatile weight fraction of prepreg per step 5.0 Example: R.S. (%) = $1.0 - \frac{.5752}{1.0 - .0865} = \frac{.5752}{.9135} = .6296$ 1.0 - .6296 = .3704

 $.3704 \times 100 = 37.04\%$



7.0 Establish Fiber Areal Density

Ad
$$(g/m^2) = \frac{Wf}{Af} \times 10.76$$

Where: Ad = fiber areal density (g/m^2) Wf = total weight of fiber in prepreg sample per 4.0 above (grams) Af = fiber area per step 4.0 (ft²) 10.76 = conversion factor, g/ft^2 to g/m^2 Example: Ad = 0.872 = 13.95 grams/ft² .0625

conversion to grams/meter² = 13.95 x 10.76 = 150.1 gram/m²

8.0 Establish Predicted Thickness per Ply of Cured Composite, 0 Void Assumed, 60 percent Fiber Volume

$$T_p = \frac{A d^4}{W f_f \times (c \times (25.4))}$$

Where: Tp = thickness per ply of composite (mils) Ad = fiber areal density, grams/meter² (per step 7.0) Wf_f = fiber weight, fraction (per step 3.0) C = theoretical density of composite (per step 2.0) 25.4= conversion factor, inch to m m

Example: $Tp = \frac{150.1}{.6522 \times 1.518 \times (25.4)} = \frac{150.1}{25.147} = 5.97$ mils/ply

9.0 Establish Prepreg Fiber Areal Density Based on Desired Cured Ply Thickness of Composite with a 60% Fiber Volume

Ad = Tp (Wff) ((c) (25.4)

Example: 5.967 (.6522) (1.518) (25.4) = $150.05 \text{ grams/meter}^2$



10.0 Establish Specific Gravity of Composite

10.1 Apparatus

- Analytical Balance a balance with a precision within 0.1 mg, accuracy within 0.05 percent relative (that is, 0.05 percent of the weight of the specimen in air), and equipped with a stationary support for the immersion vessel above the balance pan ("pan straddle").
- 2) Wire a corrosion-resistant wire, Awg No. 36 or finer, for suspending the specimen.
- 3) ^Immersion Vessel a beaker or other wide-mouthed vessel for holding the water and immersed specimen.
- 4) Thermometer a thermometer with an accuracy of ±0.1C (±0.18F) is required.

10.2 Materials

1) Water - the water shall be substantially air-free, distilled or demineralized water.

10.3 <u>Test Procedure</u>

- Cut the required number of test specimens of any convenient size, weighing from 1 to 2 grams each. Edges must be smooth, square, and surfaces clean for accurate density determination.
- 2) Weigh the specimen in air to the nearest 0.1 mg or 0.05 percent relative, which ever is greater.
- 3) Attach to the balance a piece of fine wire sufficiently long to reach from the hook above the pan to the support for the immersion vessel. Attach the specimen to the wire such that it is suspended about 1 inch above the vessel support.
- 4) Mount the immersion vessel on the support, and completely immerse the suspended specimen in water (10.2) at a temperature of $23 \pm 2C$. The vessel must not touch wire or specimen. Remove any bubbles adhering to the specimen and wire. Usually these bubbles can be removed by rubbing them with another wire. Weigh the suspended specimen to the required precision. Record this weight as b (the weight of the specimen, and the partially immersed wire in liquid). Weigh rapidly in order to minimize absorption of water by the specimen.
- 5) Weigh the wire in water with immersion to the same depth as used in the previous step. Record this weight as w (weight of the wire in liquid).





10.4 Calculations

1) Calculate the specific gravity of the composite as follows:

Sp gr (g/cc) =
$$\frac{a}{a + w - b}$$

Where: a = apparent weight of specimen, without wire in air b = apparent weight of specimen, completely immersed and of the wire partially immersed in liquid w = apparent weight of partially immersed wire. (w = density of water at water temperature

11.0 <u>Establish Resin Content of a Graphite/Polyimide Composite Using Acid</u> <u>Digestion Techniques</u>

11.1 <u>Test Procedure</u>

- 1) Use the same composite specimen employed in determining specific gravity per 10.0 above.
- 2) Obtain a clean dry extraction thimble or clean the thimble in a beaker containing HNO_3 for a minimum of 1 hour at 200 ± 10F. Wash with distilled water, dry in oven at 250 ± 10F, desiccate and cool.

Note: may be purchased from Van Waters & Rogers Co. as fritted glass extraction thimbles, medium E.C. 35 X 90 mm, Catalog No. 27743-120.

- 3) Weigh each extraction thimble to the nearest 0.1 mg and record as " W_1 ".
- 4) Dry specimen used in density determination (step 10.0) and place in clean extraction thimble and weigh to the nearest 0.1 mg. Record as " W_2 ".
- 5) Place thimble and specimen in a beaker fitted with a raised platform and a magnetic stirring bar, and add concentrated H_2SO_4 until the specimen is covered. Bring slowly to a boil and hold for 30 minutes.
- 6) Remove thimble and decant spent H_2SO_4 . Replace thimble in same beaker and repeat step 6.
- Continue boiling until fibers are completely separated and the resin is completely decomposed. This is determined by visual examination and may require additional H₂SO₄.
 - Note: Complete digestion is indicated when the test specimen changes its appearance from a unitized mass to loose, soft fibers which have a tendency to sink to the bottom of the thimble.





- 8) After digestion, place on a magnetic stirrer, stir slowly and allow to cool below 300F.
- 9) While stirring, <u>carefully</u> add 10% H_2O_2 to the hot solution.

Caution: Allow the H_2O_2 to run down the side of the beaker. Add very slowly.

- 10) Continue adding H_2O_2 until the acid solution turns a transparent clear color. If a clear color is not obtained, the test is invalid and should be repeated.
- 11) Allow acid to digest three more minutes.
- 12) Remove extraction thimble from the acid, drain, place in a rubber crucible holder on a vacuum filter flask and wash fibers with distilled water until free of acid.
- 13) Remove thimble containing fibers and dry in an oven maintained at 300 \pm 10F for a minimum of 30 minutes. Cool in a desiccator and weight to the nearest 0.1 mg. Record weight as "W₃".
- 11.2 <u>Calculations and Report of Results</u>

Calculate resin content according to the following equation:

Resin content weight, $\% = \frac{W_1 - W_3}{W_1 - W_2} \times 100$

Where: W_1 = weight of extraction thimble plus test specimen before acid digestion in grams W_2 = weight of extraction thimble in grams W_3 = weight of extraction thimble plus specimen after acid digestion in grams.

Example: 33.61985 - 33.27205 = .3478033.61985 - 32.61985

.3478 x 100 - 34.78%



13.0 Establish Void Volume Percent of Composite

 $V_{V} = 1 - \left(c \left(Wf_{f}/(f) + (Wr_{f}/(r) \times 100\right)\right)$ Where: $V_{V} = \text{void volume percent, composite}$ $\left(c = \text{actual density of composite (gram/cc)}_{\text{determined per step 10.0}}\right)$ $\left(r = \text{density of resin (gram/cc) per step 1.1}_{\text{Wf}_{f}}\right)$ $\left(f = \text{density of fiber (gram/cc) per step 1.2}_{\text{Wr}_{f}}\right)$ $\left(r = \text{resin solids weight fraction per step 11.0 or 12.0}\right)$ Example: $V_{V} = 1.518 \left(\frac{.6522}{1.65}\right) + \left(\frac{.3478}{1.32}\right)$ $\left(.3953\right) + \left(.2635\right)$ $1.518 \left(.6587\right) = 1.00$ 1 - 1.00 = 0.00% void

14.0 Establish Fiber Volume Percent of Composite

 $V_{f} = Wf_{f} \times (fc) \times 100$ Where: V_{f} = fiber volume percent of composite Wf_{f} = fiber weight fraction per step 13.0 ff = density of fiber per step 1.2 fc = density of composite (actual per step 10.0) Example: $V_{f} = \left(\frac{.6522}{1.65}\right) \times (1.518) = .6000$

.6000 x 100 - 60.00%



12.0 Establish Resin Content and Coke Number (char yield) of Cured Polyimide Laminates by Thermogravimetric Analysis (TG).

Test Procedure

- Prepare a cured sample of polyimide resin. The resin shall be free of graphite fiber by prior extraction if taken from a prepreg sample, with appropriate solvent followed by filtration and then solvent evaporation. TG sample shall be taken from specific gravity sample.
- 2) Determine the coke number (char yield) by carrying out a TG run to 800C in N_2 on the cured neat resin. The balance and control/readout system shall be operated per the applicable instrument manual. Initial sample weight shall be from 10 to 20 mg.

3) The coke number is calculated as follows:

coke number (Cn) = $\frac{W_r}{W_i}$

Where: W_i = initial sample weight W_r = weight remaining at 800C W_v = volatile weight loss to 300C (if present)

4) A sample of cured laminate weighing 20 to 30 mg is then run by TG to 800C in N_2 . Calculation of resin content is as follows:

Wt % resin =
$$\left(\frac{W_{r} - W_{v}}{(W_{i} - W_{v})(C_{n})}\right)^{10^2}$$

Where: W_r = total weight loss W_v = volatile weight loss to 300C (if present) W_i = initial weight C_n = coke number

- Note: 1) Twenty to 30 mg initial sample weight is only a guide: Samples of greater weight may be required if graphite content and coke number are high.
 - 2) This method is based on the assumption that graphite present in the laminate loses no weight to 800C in N₂.



APPENDIX B

Two specifications, material (Appendix B1) and process (Appendix B2), were prepared to comply with the requirements of Task (a). The ultrasonic inspection procedure for adhesive bonded assemblies (Appendix B3), in effect since October 1974, was utilized where production facilities were necessary for NDI of sandwich structure.

APPENDIX B1



FORM M 131-H-1 REV 5-73

Newsen Avison Letter Most 1 CONTENTS Paragraph Page 1.0 SCOPE 2 1.1 Description 2 1.2 Classification 2 1.2.1 Type 2 1.2.2 Classification 2 1.2.2.3 Grade 2 1.3 Form 2 2.0 APPLICABLE DOCUMENTS 2 2.0 APPLICABLE DOCUMENTS 2 3.1 Workmanship 3 5.1.1 Uniformity 3 5.2.2 Resin Properties 3 5.2.1 Infrared Spectography 3 5.2.2 High Preseure Liquid Chronatography 3 5.3.1 Specific Gravity 4 5.3.2 Mechanical Properties 4 5.3.4 Thermal Properties 5 5.4.1 Piber Content 5 5.4.2 Flinsh - 5 5.4.3			CODE IDE	NT. NO.												
HB0130-152 A I CONTENTS Paragraph Page 1.0 SOPE 2 1.1 Description 2 1.2 Classification 2 1.2.1 Type 2 1.2.2 Classification 2 1.2.3 Grade 2 1.3 Form 2 2.0 APPLICABLE DOCUMENTS 2 3.0 REQUIENMENTS 3 3.1.1 Uniformity 3 3.2.2 Besin Properties 3 3.2.2 REQUIENMENTS 3 3.2.2 Besin Properties 3 3.2.2 Besin Properties 3 3.3.1 Special Properties 4 3.3.1 Special Properties 4 3.3.1 Fiber Properties 5 3.4.4 Thermal Oxidation Resistance 4 3.4.4 Theregreg Froperties 5 3.4.4 Theregreg Splices 5		NUMBER			P	EVISI	ON LE	TTE				PAG	E		_	
DATES Paragraph Paragraph 1.0 SOFE 2 1.2 Classification 2 1.2 Classification 2 1.2.1 Type 2 1.2.2 Classification 2 1.3 Forde 2 2.0 AFPLICABLE DOCUMENTS 2 3.1 Workmanship 3 3.1.1 Uniformity 3 3.2.2 Resin Properties 3 3.2.2 High Pressure Liquid Chromatography 3 3.2.2 High Pressure Liquid Chromatography 3 3.2.2 High Pressure Liquid Chromatography 3 3.3.1 Specific Gravity 3 3.3.2 High Properties 4 3.3.1 Specific Gravity 5 3.4.1 Piber Cooperties 5 3.5.2 Hicharical Properties 5 3.5.4 Fiber Splices 6 3.5.4 Fiber Splices 5 3.5.4 Fiber Splices 6 3.5.5 High Properties 6 <		MB0130	-152	A										1 		
Paragraph Page 1.0 SCOPE 2 1.2 Classification 2 1.2.1 Type 2 1.2.2 Class 2 1.2.3 Grade 2 1.3 Form 2 2.0 APFLICABLE DOCUMENTS 2 2.0 APFLICABLE DOCUMENTS 3 3.1 Workmanship 3 3.1.1 Workmanship 3 3.1.2 Defects 3 3.2.1 Infrared Spectography 3 3.2.2 Rein Properties 3 3.2.1 Infrared Spectography 3 3.2.2 High Pressure Liquid Chronatography 3 3.5.3 Fiber Properties 4 3.5.4 Thermal Oxidation Resistance 4 3.5.3 Fiber Vetting 5 3.4.1 Piber Content 5 3.4.2 Fiber Vetting 5 3.4.3 Alignment 5 3.4.4					CONTE	NTS										
Paragreph Page 1.0 SCOPE																
1.0 SCOPE 2 1.1 Description 2 1.2.1 Type 2 1.2.2 Classification 2 1.2.3 Grade 2 1.2.4 Class 2 1.2.5 Grade 2 1.5 Form 2 2.0 APPLICABLE DOCUMENTS 2 2.0 APPLICABLE DOCUMENTS 2 2.0 ReQUIREMENTS 3 3.1 Workmanship 3 3.1.1 Uniformity 3 3.1.2 Defects 3 3.2.1 Infrared Spectography 3 3.2.2.1 Infrared Spectography 3 3.2.2.2 High Pressure Liquid Chromatography 3 3.5 Fiber Properties 3 3.5.1 Specific Gravity 4 3.5.2 Mechanical Properties 5 3.5.3 Fiber Properties 5 3.5.4 Thermal Oridation Resistance 4 3.4.4 Thermal Oridation Resistance 5 3.4.5	Paragr	aph														Page
1.0 SCOPE 2 1.1 Description 2 1.2 Classification 2 1.2.1 Type 2 1.2.2 Class 2 1.2.3 Grade 2 1.2.5 Grade 2 1.3 Form 2 2.0 APPLICABLE DOCUMENTS 3 3.1 Workmanship 3 3.1 Uniformity 3 3.1 Defects 3 3.1 Defects 3 3.2.1 Infrared Spectography 3 3.2.2 High Pressure Liquid Chromatography 3 3.3.1 Specific Gravity 4 3.3.2 Rechaical Properties 4 3.3.4 Thermal Oxidation Resistance 4 3.4.1 Fiber Proferties 5 3.4.2 Fiber Wring																
1.1 Description. 2 1.2.1 Type 2 1.2.2 Classification 2 1.3 Form 2 1.3 Form 2 2.0 APPLICABLE DOCUMENTS 2 2.0 REQUIREMENTS 3 3.1 Workmanship 3 3.1.1 Uniformity 3 3.1.2 Defects 3 3.2.2 Resin Properties 3 3.2.2 High Pressure Liguid Chromatography 3 3.3.5 Fiber Properties 4 3.4.7 Thermal Oridation Resistance 4 3.5.3 Fiber Properties 5 3.5.4 Thermal Oridation Resistance 4 3.5.3 Fiber Properties 5 3.4.4 Thermal Oridation Resistance 5 3.4.4 Fremes Opporties 5 3.4.5.1 Friber Splices 5 3.4.6 Splices 5 3.4.7 Alignment 5 3.4.8 Splices 6 3.4.7	1.0		SCOPE	• • •	• •	• •	• •	•	•	• •	•	••	• •	• •	٠	. 2
1.2.1 Type 2 1.2.2 Class 2 1.3 Form 2 2.0 APPLICABLE DOCUMENTS 2 3.0 REQUIREMENTS 3 3.1 Workmanship 3 3.1.1 Uniformity 3 3.2 Defects 3 3.2.2 Hein Properties 3 3.2.2 High Pressure Liquid Chromatography 3 3.2.2 High Pressure Liquid Chromatography 3 3.3.5 Fiber Properties 4 3.5.1 Specific Gravity 4 3.5.2 Mechanical Properties 4 3.5.3 Finish 5 3.5.4 Thermal Oxidation Resistance 4 3.5.1 Specific Gravity 5 3.5.2 Mechanical Properties 5 3.4.1 Piber Content 5 3.4.2 Fiber Wetting 5 3.4.3 Alignment 5 3.4.4.1 Prepreg Splices 5 3.4.5.1 Prepreg Splices 6	1.1		Description	• • •	R •	• •	•••	•	• •	• •	•	• •	• •	••	•	2
12.2 Class 2 1.2.3 Grade 2 1.3 Form 2 2.0 APPLICABLE DOCUMENTS 2 3.0 REQUIREMENTS 2 3.1 Workmanship 3 3.1.1 Uniformity 3 3.1.2 Defects 3 3.2.4 Infrared Spectography 3 3.2.2 High Pressure Liquid Chronatography 3 3.3.1 Specific Gravity 4 3.3.1 Specific Gravity 4 3.3.1 Specific Gravity 4 3.3.2 Mechanical Properties 4 3.3.3 Flinsh 4 3.3.4 Thermal Oxidation Resistance 4 3.4.4 Prepreg Properties 5 3.4.1 Fiber Content 5 3.4.2 Hick Property Splices 6 3.4.4 Gaps 5 3.4.5 Splices 6 3.4.6 Width 6 3.4.7 Bregreg Splices 6 3.4.6 Width <td>1.2.1</td> <td></td> <td>Tassiica.</td> <td>• 10</td> <td>• •</td> <td>• •</td> <td>• •</td> <td>• •</td> <td>•</td> <td>• •</td> <td>•</td> <td>••</td> <td>• •</td> <td>• •</td> <td>•</td> <td>2</td>	1.2.1		Tassiica.	• 10	• •	• •	• •	• •	•	• •	•	••	• •	• •	•	2
1.2.3 Grade 2 1.3 Form 2 2.0 APPLICABLE DOCUMENTS 2 3.0 REQUIREMENTS 3 3.1 Workmanship 3 3.1.1 Uniformity 3 3.1.2 Defects 3 3.2.1 Infrared Spectography 3 3.2.2 Resin Properties 3 3.2.1 Infrared Spectography 3 3.2.2 High Pressure Liquid Chronatography 3 3.2.1 Specific Gravity 3 3.2.2 High Properties 4 3.3.3 Finish 4 3.4.1 Fiber Properties 4 3.5.2 Hechanical Properties 5 3.4.3 Frierg Properties 5 3.4.4 Prepreg Properties 5 3.4.2 Fiber Weiting 5 3.4.3 Alignment 5 3.4.4 Gaps 5 3.4.5 Splices 5 3.4.6 With 6 3.4.7 Edges	1.2.2	•		• • •	• •	••	• •	•	•	• •	•	••	•	••	•	2
1.3 Form 2 2.0 APPLICABLE DOCUMENTS 2 3.0 REQUIREMENTS 3 3.1 Workmanship 3 3.1.1 Uniformity 3 3.1.2 Defects 3 3.2.3 Infrared Spectography 3 3.2.4 Infrared Spectography 3 3.2.2 High Pressure Liquid Chronatography 3 3.3.1 Specific Gravity 3 3.3.2 Mechanical Properties 4 3.3.3 Flineh 4 3.4.4 Thereral Oxidation Resistance 4 3.5.3 Flineh 5 3.4.4 Thereral Oxidation Resistance 5 3.4.4 Thereral Oxidation Resistance 5 3.4.5 Splices 5 3.4.6 Splices 5 3.4.7 Prepreg Properties 5 3.4.8 Splices 5 3.4.5 Splices 5 3.4.5 Splices 6 3.4.6 Width 6 3.4.7 <td>1.2.3</td> <td></td> <td>Grade</td> <td>•••</td> <td>•••</td> <td>•••</td> <td>•••</td> <td>•</td> <td></td> <td>•••</td> <td></td> <td>•••</td> <td></td> <td></td> <td>:</td> <td>2</td>	1.2.3		Grade	•••	•••	•••	•••	•		•••		•••			:	2
2.0 APPLICABLE DOCUMENTS 2 3.0 MEQUIREMENTS 3 3.1 Workmanship 3 3.1.1 Uniformity 3 3.1.2 Defects 3 3.2.2 Resin Properties 3 3.2.2 High Pressure Liquid Chromatography 3 3.2.2 High Pressure Liquid Chromatography 3 3.3.1 Specific Gravity 3 3.5.2 Mechanical Properties 4 3.5.3 Finish 4 3.5.4 Thermal Oxidation Resistance 4 3.5.4 Prepreg Properties 5 3.4.2 Fiber Wetting 5 3.4.3 Alignment 5 3.4.4 Gaps 5 3.4.5 Splices 6 3.4.6 Width 6 3.4.7 Fdeges 6 3.4.8 Length 6 3.4.9 Arcal Weight 6 3.4.9 Arcal Weight 6 3.4.9 Arcal Weight 6 3.5.1	1.3		Form		•••	•••				•••		••	•		•	2
2.0 APPLICABLE DOCUMENTS 2 3.0 REQUIREMENTS 3 3.1 Workmanship 3 3.1.1 Uniformity 3 3.1.2 Defects 3 3.2.1 Infrared Spectography 3 3.2.2 High Pressure Liquid Chromatography 3 3.2.2 High Pressure Liquid Chromatography 3 3.3.1 Specific Gravity 3 3.5.2 Mechanical Properties 4 3.5.3 Finish 4 3.5.4 There Properties 4 3.5.5 Finish 4 3.5.4 Thermal Oxidation Resistance 4 3.4.1 Fiber Gontent 5 3.4.2 Fiber Wetting 5 3.4.3 Alignment 5 3.4.4 Gaps 5 3.4.5 Splices 5 3.4.6 Width 6 3.4.7 Edges 6 3.4.6 Width 6 3.4.6 Length 6 3.5.1 Cured Ply														•		
3.0 REQUIREMENTS 3 3.1 Workmanship 3 3.1.1 Uniformity 3 3.1.2 Defects 3 3.1.2 Defects 3 3.1.2 Defects 3 3.2 Resin Properties 3 3.2.1 Infrared Spectography 3 3.2.2 High Pressure Liquid Chromatography 3 3.3.1 Specific Gravity 4 3.5.2 Mechanical Properties 4 3.5.3 Finish 4 3.5.4 Thermal Oxidation Resistance 4 3.5.3 Filer Properties 5 3.4.1 Fiber Content 5 3.4.2 Fiber Wetting 5 3.4.3 Alignment 5 3.4.4 Gaps 5 3.4.5 Splices 5 3.4.6 Width 6 3.4.7 Edges 6 3.4.6 Wight 6 3.4.6 Length 6 3.5.1 Cured Ply Thickness 6	2.0		APPLICABLE 1	DOCUME	NTS	• •	••	•	• •	••	•	••	•	••	•	2
3.1 Workmanship	3.0		REQUIREMENT	5	• •			•			•		•		•	3
3.1.1 Uniformity 3 3.1.2 Defects 3 3.2 Hegin Properties 3 3.2.1 Infrared Spectography 3 3.2.2 High Pressure Liquid Chromatography 3 3.3 Piber Properties 4 3.5.1 Specific Gravity 4 3.5.2 Mechanical Properties 4 3.5.3 Finish - 4 3.5.4 Thermal Oxidation Resistance 4 3.4.1 Fiber Content 5 3.4.2 Fiber Wetting 5 3.4.3 Alignment 5 3.4.4 Gaps 5 3.4.5 Splices 6 3.4.6 Width 6 3.4.7 Edges 6 3.4.8 Length 6 3.4.9 Areal Weight 6 3.4.9 Areal Weight 6 3.4.9 Areal Weight 6 3.4.9 Areal Properties 7 4.0 Qualification 8 3.5.2 P	3.1		Workmanship		• •	• •	• •	•	• •	• •	•	• •	•	• •	•	3
3.1.2 Defects 3 3.2.1 Infrared Spectography 3 3.2.2 High Pressure Liquid Chromatography 3 3.3 Fiber Properties 4 3.3.1 Specific Gravity 4 3.3.2 Mechanical Properties 4 3.3.1 Specific Gravity 4 3.3.2 Mechanical Properties 4 3.3.4 Thermal Oxidation Resistance 4 3.4.1 Fiber Content 5 3.4.2 Fiber Wetting 5 3.4.3 Alignment 5 3.4.4 Gaps 5 3.4.5 Splices 5 3.4.6 Splices 5 3.4.7 Edges 6 3.4.8 Length 6 3.4.9 Areal Weight 6 3.4.10 Storage Life 6 3.4.9 Areal Weight 6 3.4.10 Storage Life 6 3.4.11 Cured Ply Thickness 6 3.5.2 Physical Properties 7 4.0<	3.1.1		Uniformity		• •	••	• •	•	• •	• •	•	••	•	• •	•	3
3.2Resin Properties33.2.1Infrared Spectography33.2.2High Pressure Liquid Chromatography33.3Fiber Properties43.3.1Specific Gravity43.3.2Mechanical Properties43.3.3Finish	3.1.2		Defects	• • •	• •	• •	• •	•	• •	• •	٠	• •	•	• •	•	3
3.2.1 Infrared Spectography	3.2		Resin Proper	rties	• •	••	• •	•	• •	• •	•	• •	• •	• •	•	3
3.2.2 High Pressure Liquid Chromatography	3.2.1		Infrared Sp	ectogra	aphy	• •	• •	•	• •	• •	٠	• •	• •	• •	•	3
3.3 Fiber Properties 4 3.3.1 Specific Gravity 4 3.3.2 Mechanical Properties 4 3.3.3 Finish 4 3.3.4 Thermal Oridation Resistance 4 3.4 Pripreg Properties 5 3.4.1 Fiber Content 5 3.4.2 Fiber Wetting 5 3.4.3 Alignment 5 3.4.4 Gaps 5 3.4.5 Splices 5 3.4.4 Gaps 5 3.4.5 Splices 6 3.4.6 Width 6 3.4.7 Edges 6 3.4.8 Length 6 3.4.9 Areal Weight 6 3.4.10 Storage Life 6 3.5.1 Cured Ply Thickness 6 3.5.2 Physical Properties 7 4.0 QUALITY ASSURANCE 8 4.1.1 Qualification Sample 8	3.2.2		High Pressu:	re Liqu	uid C	hrom	ato	gra]	phy	• •	٠	• •	•	• •	•	3
3.3.1 Specific Gravity 4 3.3.2 Mechanical Properties 4 3.3.3 Finish 4 3.4 Thermal Oxidation Resistance 4 3.4 Prepreg Properties 5 3.4.1 Fiber Content 5 3.4.2 Fiber Wetting 5 3.4.3 Alignment 5 3.4.4 Gaps 5 3.4.5 Splices 5 3.4.6 Width 6 3.4.7 Edges 6 3.4.8 Length 6 3.4.7 Edges 6 3.4.8 Length 6 3.4.9 Areal Weight 6 3.4.10 Storage Life 6 3.5.1 Gured Ply Thickness 6 3.5.2 Physical Properties 7 4.0 QUALITY ASSURANCE 8 4.1 Qualification Sample 8 4.1.11 Qualification Sample 8	3.3		Fiber Prope:	rties	• •	• •	• •	•	• •	• •	•	••	•	• •	•	4
5.3.2 Mechanical Properties 4 3.3.3 Finish	3.3.1		Specific Gr	avity	• •	• •	• •	•	• •	• •	•	• •	•	• •	•	4
3.3.3FinishFinish43.3.4Thermal Oxidation Resistance43.4Prepreg Properties53.4.1Fiber Content53.4.2Fiber Wetting53.4.3Alignment53.4.4Gaps53.4.5Splices53.4.5.1Prepreg Splices63.4.6Width63.4.7Edges63.4.8Length63.4.9Areal Weight63.5.1Cured Ply Thickness63.5.2Physical Properties63.5.3Mechanical Properties74.0QUALITY ASSURANCE84.1.1Qualification84.1.1.1Qualification Sample8	3.3.2		Mechanical 1	Proper	ties	••	• •	•	• •	• •	•	• •	•	• •	٠	4
5.3.4Thermal Oxidation Resistance43.4Prepreg Properties53.4.1Fiber Content53.4.2Fiber Wetting53.4.3Alignment53.4.4Gaps53.4.5Splices53.4.5.1Prepreg Splices63.4.6Width63.4.7Edges63.4.8Length63.4.9Areal Weight63.4.10Storage Life63.5.2Physical Properties63.5.3Mechanical Properties63.5.4Gualification74.0QUALITY ASSURANCE84.1.1Qualification Sample8	3.3.3	5	Finish	• • •	••.	• •	• •	•	••	• •	•	• •	•	•••	. •	4
3.4Prepreg Properties53.4.1Fiber Content53.4.2Fiber Wetting53.4.3Alignment53.4.4Gaps53.4.5Splices53.4.5.1Prepreg Splices63.4.5.2Fiber Splices63.4.6Width63.4.7Edges63.4.8Length63.4.9Areal Weight63.4.10Storage Life63.5.1Cured Ply Thickness63.5.2Physical Properties63.5.3Mechanical Properties74.0QUALITY ASSURANCE84.1.1Requirements84.1.1.1Qualification Sample8	3.3.4	•	Thermal Oxid	lation	Kesi	stan	ce	•	• •	•. •	•	• •	•	• •	•	4
3.4.1 Fiber Uontent 5 3.4.2 Fiber Wetting 5 3.4.3 Alignment 5 3.4.4 Gaps 5 3.4.5 Splices 5 3.4.5.1 Prepreg Splices 6 3.4.6 Width 6 3.4.7 Edges 6 3.4.8 Length 6 3.4.9 Areal Weight 6 3.4.10 Storage Life 6 3.5.1 Cured Ply Thickness 6 3.5.2 Physical Properties 7 4.0 QUALITY ASSURANCE 8 4.1.1 Requirements 8 4.1.1 Qualification Sample 8	5.4		Prepreg Pro	pertie	5 %	• •	• •	•	••	• •	•	• •	•	• •	•	2
3.4.2 Fiber Wetting 5 3.4.3 Alignment 5 3.4.4 Gaps 5 3.4.5 Splices 5 3.4.5 Splices 5 3.4.5 Prepreg Splices 6 3.4.5.1 Prepreg Splices 6 3.4.5.2 Fiber Splices 6 3.4.6 Width 6 3.4.7 Edges 6 3.4.8 Length 6 3.4.9 Areal Weight 6 3.4.10 Storage Life 6 3.4.10 Storage Life 6 3.5 Laminate Properties 6 3.5.1 Cured Ply Thickness 6 3.5.2 Physical Properties 7 4.0 QUALITY ASSURANCE 8 4.1 Qualification 8 4.1.1 Requirements 8 4.1.1 Qualification Sample 8	5.4.1		Fiber Conte	nt • •	• •	• •	• •	•	• •	• •	•	• •	•	•••	•	2
3.4.4 Gaps 5 3.4.4 Gaps 5 3.4.5 Splices 5 3.4.5.1 Prepreg Splices 6 3.4.5.2 Fiber Splices 6 3.4.6 Width 6 3.4.7 Edges 6 3.4.8 Length 6 3.4.9 Areal Weight 6 3.4.10 Storage Life 6 3.5.1 Cured Ply Thickness 6 3.5.2 Physical Properties 6 3.5.3 Mechanical Properties 7 4.0 QUALITY ASSURANCE 8 4.1.1 Requirements 8 4.1.11 Qualification Sample 8	3.4.2		Fiber wetti	ng • •	• •	•••	• •	•	••	• •	•	• •	•	• •	•	2
3.4.4GapsSplices53.4.5Splices53.4.5.1Prepreg Splices63.4.5.2Fiber Splices63.4.6Width63.4.7Edges63.4.8Length63.4.9Areal Weight63.4.10Storage Life63.5Laminate Properties63.5.1Cured Ply Thickness63.5.3Mechanical Properties74.0QUALITY ASSURANCE84.1.1Requirements84.1.1.1Qualification Sample8	3.4.3)	Alignment .	• • •	• •	• •	• •	•	• •	• •	•	••	•	••	•	25
3.4.5.1Splices63.4.5.2Fiber Splices63.4.6Width63.4.7Edges63.4.8Length63.4.9Areal Weight63.4.10Storage Life63.5.1Cured Ply Thickness63.5.2Physical Properties63.5.3Mechanical Properties74.0QUALITY ASSURANCE84.1.1Qualification84.1.1Qualification Sample8	7•4• 4	•	Gaps • • •	• • •	• •	• •	• •	•	••	• •	•	• •	•	••	•	5
3.4.5.1Frepreg Splices63.4.5.2Fiber Splices63.4.6Width63.4.7Edges63.4.8Length63.4.9Areal Weight63.4.10Storage Life63.5Laminate Properties63.5.1Cured Ply Thickness63.5.2Physical Properties63.5.3Mechanical Properties74.0QUALITY ASSURANCE84.1Qualification84.1.1Requirements84.1.1.1Qualification Sample8	2+4+2) · •	Depres • •	• • •	• •	• •	• •	•	• •	• •	•	• •	•	• •		5
3.4.6Width63.4.6Width63.4.7Edges63.4.8Length63.4.9Areal Weight63.4.10Storage Life63.5Laminate Properties63.5.1Cured Ply Thickness63.5.2Physical Properties74.0QUALITY ASSURANCE84.1Qualification84.1.1Requirements84.1.1Qualification Sample8	. 2•4•2)•1 5.2	Fiber Splic	LCes .	•••	• •	•.•	•	•••	•••	:	••	•	••	:	6
5.4.6With	3 4 6		Width		•••	•••	•••	•		•••					Ī	6
3.4.8Length63.4.9Areal Weight63.4.10Storage Life63.5Laminate Properties63.5.1Cured Ply Thickness63.5.2Physical Properties74.0QUALITY ASSURANCE84.1Qualification84.1.1Requirements84.1.1.1Qualification Sample8	J•4•0 % A 7) 7		• • •	••	• •	• •	. •	• •	• •	•	••	•	••	•	6
3.4.9Areal Weight63.4.10Storage Life63.5Laminate Properties63.5.1Cured Ply Thickness63.5.2Physical Properties74.0QUALITY ASSURANCE84.1Qualification84.1.1Requirements84.1.1.1Qualification Sample8	3.4.8	2	Length	•••	•••	•••	•••	•			:					6 6
3.4.10Storage Life63.5Laminate Properties63.5.1Cured Ply Thickness63.5.2Physical Properties63.5.3Mechanical Properties74.0QUALITY ASSURANCE84.1Qualification84.1.1Requirements84.1.1Qualification Sample8	3.4.9	, 1	Areal Weigh	•••				•								6
3.5Laminate Properties63.5.1Cured Ply Thickness63.5.2Physical Properties63.5.3Mechanical Properties74.0QUALITY ASSURANCE84.1Qualification84.1.1Requirements84.1.1.1Qualification Sample8	3.4.1	, O	Storage Life													6
3.5.1Cured Ply Thickness63.5.2Physical Properties63.5.3Mechanical Properties74.0QUALITY ASSURANCE84.1Qualification84.1.1Requirements84.1.1Qualification Sample8	3.5		Laminate Pr	operti	es .							•••		• •	•	6
3.5.2Physical Properties63.5.3Mechanical Properties74.0QUALITY ASSURANCE84.1Qualification84.1.1Requirements84.1.1.1Qualification Sample8	3.5.1		Cured Plv T	nickne	88 .								•			6
3.5.3Mechanical Properties74.0QUALITY ASSURANCE84.1Qualification84.1.1Requirements84.1.1.1Qualification Sample8	3.5.2	2	Physical Pr	operti	es .			•			•	• •	•	• •	•	6
4.0QUALITY ASSURANCE84.1Qualification84.1.1Requirements84.1.1.1Qualification Sample8	3.5.3	5	Mechanical	Proper	ties	• •	• •	•	• •	• •	•	•••	•	• •	•	7
4.1Qualification84.1.1Requirements84.1.1.1Qualification Sample8	A.0		OULT TOV ASS	TRANCE	. -								-		•	8
4.1.1Requirements	4•V A_1		Qualificati	n							-				•	8
4.1.1.1 Qualification Sample	4.1.1	L .	Requirement	3				•		• •	•				•	8
	4.1.1	.1	Qualificati	on Sam	ple								•		•	8
			~~~~~~~~~~~		r - *			-	•		-	- •		2	-	
										,						

M131-H-28 REV. 4-73

CODE IDENT. NO.

	MB0130-	152				Τ		1	T	FAGE		ii	1
<b>_</b>		<b>1</b> <i>JL</i>	<u>1 ~ 1</u>				1		1	L		**	I
			CON	ITENI	'S (Co	nt'd	)						
Para	agraph												Page
4.1.1	•2	Certified T	est 1	Repo:	rt .			•••		••	• •	• •	. 8
4.1.1	•3	Test Facili	ties	• •			• •	• •		• •	• •	• •	8
4.1.1	•4	Demonstrati	on T	ests	• •	• • •	• •	• •	• •	• •	• •	• •	8
4.1.2		Changes	• •	• •	• •	• • •	• •	•••	• •	• •	• •	• •	8
4.2		Acceptance	• •	• •	• • •	• • •	•••	•••	• •	• •	• •	• •	8
4.2.1		Definitions	• •	• •	• • •	• •	• •	•••	• •	• •	• •	• •	8
4.2.2		Testing	• •	• •	• • •	• •	• •	• •	• •	• •	•••	••	0
4.5		Certificati	on •		•••	• • •	• •	•••	••	•••	• •	•••	
4.4		Responsibil	ity :	for.	Inspe	tior	anc	Tes	ting	• •	• •	• •	
4.4.1		Ingrection	Page	•••	• • •	• •	• •	••	• •	• •	• •	• •	
4.4.6		TUSPECTION	neco:	rus	• • •	•••	•••	•••	• •	• •	• •	• •	
4.5.1		iest method Standerd Co	• • ndi +	i one	• • •	•••	••	••	• •	•••	• •	•••	
A.5.2		Regin		-0113	• • •	• •	•••	• •	• •	• •	• •	• •	2
4.5.2	.1	Infrared Sn		•••	•••	•••	•••	•••	• •	• •	•••	• •	-
4.5.2	.2	High Pressu	re L	iguid	1 Chro	mato	pret	hv .	•••	•••	• • •	•••	זֹנ
4.5.2	.3	Record Rete	ntio	n .									10
4.5.3	• >	Fiber											10
4.5.3	•1	Specific Gr	avit	y .					•••		•••	•••	1
4.5.3	•2	Mechanical	Prop	erti	es of	Fibe	ers .					• •	12
4.5.3	•3	Thermal Oxi	dati	on Re	esista	ince	• •	• •	• •		• •		15
4.5.4	-	Prepreg	• •	• •			• •	•••	• •	• •	• •	• •	10
4.5.4	•1	Fiber Conte	nt.	• •	• • •	• •	•••	• •	• •	•	• •	• •	16
4.5.4	•2	Areal Weigh	t.	• •	• •		• •	•••		• •			1'
4.5.4	•3	Volatiles C	onte	nt .	• • •			•••	• •	• •	• •	• •	l.
4.5.4	•4	Prepreg Res	in Co	ontei	nt • ·	• •	•••	• •	• •	• •	• •	• •	18
4.7.4	• 5	Gel Time .	••	• •	• • •	• • •	•••	• •	• •	• •	• •	• •	18
4.7.7	•	Curea Lamin Desession	ate	•••	•••	• • •	••	•••	••	••	• •	• •	1
4.0.0	•⊥ ?	Cured Plum	. OI ( hiolo		i rami	nate	• •	• •	••	••	• •	• •	27
4.5.5	• Z _ R	Specific Cr	avi t	uess v	• • •	•••	•••	•••	•••	• •	• •	• •	24
4.5.5	• )	Fiber Conte	av _ v. n+	<b>,</b> •	•••			•••	• •	•••	•••	••	20
4.5.5	•5	Fiber Conte	$nt_{-}$	Volu	ne Per	cent		•••				•••	20
4.5.5	•6	Glass Trans	itio	n Ter	npera	ture				• •		• •	2
4.5.5	•7	Short-Beam	Shea	r .	• • •				• •	• •	• •	• •	24
4.5.5	•8	Longitudina	1 F1	exura	al Tes	st .		•••	• •	••	• •	••	24
5.0		PREPARATION	FOR	DEI.	IVERY								25
5.1		Packaging .		• •						•••	•••	•••	2
5.2		Identificat	ion									• •	25
5.3		Packing	• •	• •		• •			• •	• •		· ·	25
5.4		Marking of	Ship	ment	• • •		• •	• •	• •	• •	• •	• •	25
			2										
0.0		NOTE	• •	• •	• • •	• •	• •	••	••	• •	• •	• •	25
0+T		intended Us	е.	• •	• • •	• • •	•••	•••	• •	• •	••	• •	25

		CODE IC	DENT. NO	L.							
	NUMBER				REVISIO	NLETTER	1	]	PAGE		
	MB0130-	-152	A					1		2	
unidi react tempe	1.0 SCOPE 1.1 <u>Descripti</u> rectional tape ing polyimide ratures up to	on. This and woven resins sui 316C (600F)	specif graph table ).	ficat nite for	ion es fiber fabric	tablis fabric ation	hes t impr of st	he re egnat	equireme ted with ares for	nts fo conde use a	or ensation- it
class	1.2 Classific es, types and	grades:	e prei	.mpre	gnated	mater:	ials	shall	l be of	the fo	llowing
as fo	1.2.1 The typ llows:	e shall spe	ecify	the	chemic	al nat	ure o	f the	e matrix	resir	n system
	Type I	Nonhalog identifi	enated ed <b>a</b> s	l sys LaRC	tem wi 160	th dia	mine	to te	etra-ami	ne miz	tures,
	Type II	Nonhaloge	enated	l sys	tem, i	dentif	ied a	s PMF	1-15		
	1.2.2 The cla	ss shall s	pecify	gra	phite	fiber	stren	gth a	ind modu	lus pr	operties
	Class 1 -	Graphite pr strength of (33 msi).	repreg f 2.76	g mad GN/	e from n ² (40	high ( ) ksi)	and	gth f modul	lus 230	aving GN/m ²	a minimu
	Class 2 -	Graphite pr strength of (50 msi).	repreg f 2.20	g mad ) GN/1	e from n ² (32)	high 1 O ksi)	nodul and	us fi modul	ber hav lus 345	ing a GN/m ²	minimum
	Class 3 -	Graphite p 483 GN/m ²	repreg (70 ms	g mad si).	e from	fiber	havi	ng a	modulus	in ex	cess of
:	1.2.3 The gra	de shall s	pecify	the	graph	ite fo	<b>r-</b> m				
	Grade U	Unidired	ctions	l ta	pe						
	Grade B	Broad Go	oods,	Wove:	n fabr	ic					
the r	1.3 <u>Form</u> - Th equired widths	e graphite on a core	tape which	prei sha	npregna 11 not	ated ma be de:	ateri forme	al is d by	to be the mat	furnis erial	hed in weight.
a par betwee	2.0 APPLICABL t of this spec en these docum	E DOCUMENTS ification f ents and th	5. Th to the nis sp	e la e exte ecif:	test in ent spe ication	ssues ( ecified n, this	of th 1 her 3 spe	e fol ein. cific	lowing In cas ation s	docume e of c hall p	ents form onflict prevail.
	Test Metho	d				1	le tho	ds of	Testin	g	
	FED-STD-40	6			Plas	tics:	Meth	ods c	f Testi:	ng	
	MIL-B-117				Bags Pack	, Sleev aging	/es,	and T	ubing -	Inter	ior
	MIL-G-8341	0			Grap] Shoo	nite F	iber i	Resin	Inpreg	rated	Tape and
					onee	t, 10r	Hard	Layu	p		

M131-H-28 REV. 4-73

CODE IDENT. NO.

	NUMBER				REVIS	ION LE	TER			PAGE		]
	MB0130-	-152	A								3	
	ASTM D 638	3			Ter	sile	Prop	perti	es o	f Plast	ics	
	ASTM D 790	)			Fle	xural	l Pro	opert:	ies	of Plas	tics	
	ASTM D 792	2			Spe by	cific Displ	: Gra Lacen	vity ment	and	l Densit	y of	Plastics
	ASTM D 234	14			Apj Rei Mei	aren nfor hod	t Hoi ced H	rizon Plast:	tal ics	Shear S by Shor	trena t-Bea	gth of am Shear
•	3.0 REQUIREM	ents										
	3.1 <u>Workmans</u>	nip										
and c are do accep	3.1.1 Uniform lean and free etrimental to table only to	nity. The main from foreign fabrication the limits	ater n ma , ap give	rial teri opean en in	shall als, a ance, 1 3.4 a	be un ind sl or pe ind 3	nifor nall erfor .5.	m in not 1 mance	qua have e.	lity and charac These do	d cor teria efeci	ndition, stics which ts shall b
is fla in tha	3.1.2 Defects agged, and repat roll.	g. Material placement ya	con rdag	ntair ge is	ning de saddeo	fect:	s sha the 1	all be coll :	e al for	lowed i every d	f eac efect	ch defect t occurin

3.2 <u>Resin Properties</u>. A one-pint sample shall be taken from each batch of neat resin used to perform the prepreging of the material to this specification and shall be sent with the prepreg order represented. The resin sample shall be identified with cook batch, filming batch and other processing identification as applicable.

3.2.1 Infrared Spectrography. An infrared spectrogram shall be made on a sample of the neat resin from each batch of prepreg per 4.5.2.2 and a copy of this spectrogram shall be transmitted as supporting data to the prepreg certification.

3.2.2 <u>High Pressure Liquid Chromatography</u>. Liquid Chromatographic analysis shall be made on neat resin sample from each batch of prepreg per 4.5.2.3 and a copy of all chromatograms shall be transmitted as supporting data to the prepreg certification.

3.3 Fiber Properties. Fiber properties shall be determined by the fiber manufacturer and the information transmitted to the prepreger by certification for ultimate transmission to Rockwell International as specified herein. A 50 gram sample from each batch of fiber used in the prepreging operation shall be supplied to Rockwell International with the prepreg order represented. Fibers used shall exhibit the properties shown in Table I. The prepreg supplier shall certify in writing that the fibers meet the values of Table I. The actual test values and fiber lots may be obtained from the fiber manufacturer.

0004S

M131-H-28 REV. 4-73

NUMBER       MB0130-152       A         3.3.1       Specific Gravity. The symptotic symptot	REVISION LETT pecific gravit ansmitted as s Fiber room tem shall be measu all be polyimid l prior to shi al, and quanti ance. Fiber we for 168 hours ht Loss Proper	y of each lo upporting de perature ten red per 4.5 de. Epoxy s pment. Fibe ty expressed eight loss s per 4.5.3.3 ties of Grag	PAGE 4 ot of fiber sh ata to the pre- nsile strength .3.2. sizing may be er certificati d as weight ad shall not exce 3. phite Fibers	ion siditi
ABOI 100-152       A         3.3.1       Specific Gravity. The service         measured per 4.5.3.1 and shall be trapertification.         3.3.2       Mechanical Properties. If         gensile modulus and ultimate strain a         3.3.3       Finish. Fiber sizing shall be traperties. If         3.3.4       Thermal Oxidation Resistan         3.3.4       Thermal Oxidation Resistan         Deercent when exposed to 600°F in air         Table 1       Strength and Weight	pecific gravit ansmitted as s Fiber room tem shall be measu all be polyimi l prior to shi al, and quanti ance. Fiber we for 168 hours ht Loss Proper	y of each lo upporting d perature ten red per 4.5 de. Epoxy s pment. Fibe ty expressed eight loss s per 4.5.3.3 ties of Graj	4 ot of fiber sh ata to the pro- nsile strength .3.2. sizing may be er certification d as weight ad shall not exce 3. phite Fibers	hall preg
3.3.1 <u>Specific Gravity</u> . The speasured per 4.5.3.1 and shall be transmission. 3.3.2 <u>Mechanical Properties</u> . 1 Sensile modulus and ultimate strain a 3.3.3 <u>Finish</u> . Fiber sizing sha substituted with engineering approval include a statement of sizing materia 3.3.4 <u>Thermal Oxidation Resistan</u> bercent when exposed to 600°F in air Table 1 Strength and Weigh	pecific gravit ansmitted as s Fiber room tem shall be measu all be polyimi l prior to shi al, and quanti ance. Fiber we for 168 hours ht Loss Proper	y of each lo upporting de perature ter red per 4.5 de. Epoxy s pment. Fibe ty expressed eight loss s per 4.5.3.3 ties of Gray	ot of fiber sl ata to the pro- nsile strength .3.2. sizing may be er certification d as weight act shall not excess. phite Fibers	nall preg i, ion s lditi
Table 1 Strength and Weig	ht Loss Proper	ties of Graj	phite Fibers	
-	 	Graphite	e Fiber	
Properties	Class I	Class	II Clas	s II
Modulus, GPa (msi), min.	230 (33)	320 (50)	520 (7	'O)
Tensile Strength, MPa (ksi), min.	2760 (400)	2200 (320	) <b>1860 (</b>	270)
Density, g/cc (lb/cu in), <u>+</u> .04	1.77 (.0639)	1.90 (.06	586) 2.07 (	•074
Weight Loss After 125 Hours at 316C (600°F), % W./W ₂ , max.	1.5	1.5	1.5	
Weight Per Unit Length of Tow, Kg/m (lb/in) X 10 ⁻⁶ , <u>+</u> 10%	780 (44)	760 (43)	800 (4	.5)
efinitions GPa = Gigapascals msi = Million pounds per squar MPa = Megapascals KSI = Thousand pounds per squar g/cc = grams per cubic centime lb/cu in = pounds per cubic ir % W ₁ /W ₂ = Weight lost over weight loss in p kg/m = kilograms per meter lb/in = pounds per inch	re inch eter ach r original weig percent	t times a	hundred gives	

3.4 <u>Prepreg Properties</u>. The prepreg material furnished to this specification shall be of quality workmanship. It shall be essentially free from crimped fibers, gelled resin particles, twists, fiber balling, unwetted fibers and dry or boardy areas. Individual tows shall be parallel to the tape or sheet centerline. Indications of impurities, dry areas, areas of nonuniformity, incomplete impregnation, gelled resin, hard spots, or localized color difference in impregnated cloth shall be marked by tape as nonconforming area. The physical properties shall be as shown in Table 2.

Table 2 Physical Properties of the Uncured Prepreg

Property	Requirement	Test Method Paragraph
Volatiles, Weight percent	12 <u>+</u> 3	4.5.4.3
Resin Solids, Weight percent	38 <u>+</u> 3	4.5.4.4
Gel Time at 204°C (400°F) Minutes	0.5 to 2.0	4•5•4•5 ·

3.4.1 <u>Fiber Content</u>. The prepreg as specified is intended to produce a cured laminate with a 60 + 2 percent fiber content by volume per 4.5.4.1.

3.4.2 Fiber Wetting. The filaments shall be completely wetted by the resin; no cured resin particles are permitted when determined visually using magnification as necessary.

3.4.3 <u>Alignment</u>. The filament bundles in the tape shall be parallel to the longitudinal direction of the prepreg within an angle of one degree when examined visually using appropriate aids to measure angular alignment.

3.4.4 Gaps. Any gaps within or between tows in unidirectional tape shall comply with the following when determined visually using adequate scales.

(a) No gap shall exceed 0.010 inch in width.

4171 M 70 DEV 4 72

- (b) The length of any gap shall not exceed 4 inches.
- (c) Gaps in line with each other and no more than one inch apart shall be considered as one gap, regardless of number.
- (d) Gaps with excessive width or length will be considered defective and must be replaced as described in 3.1.2.

3.4.5 <u>Splices</u>. Splices shall be determined visually using magnification as necessary.

0004S

CODE IDENT. NO.

NUMBER		P	PAGE	]					
MB0130-152	A							6	

3.4.5.1 <u>Prepreg Splices</u>. Prepreg splices are permitted on any roll of tape where processing is continuous without change in fiber or resin batch. Such splice must be marked by tape as a nonconforming area per 3.1.2.

3.4.5.2 <u>Fiber Splices</u>. Splices of filament bundles shall be kept as short as possible. Adhesive used for bonding filement bundles shall be compatable with the resin system and shall be identified and transmitted as supporting data to the prepreg certification. Where possible, fiber splices shall be flagged during prepreg production.

3.4.6 Width. The prepreg width shall be as specified on the purchase order. Width tolerance for unidirectional tape shall be + 0.125 inch.

3.4.7 Edges. Maximum acceptable waviness of any 24-inch length of tape shall be 0.030 inch from the edge when measured with appropriate straight edge.

3.4.8 Length. The length of each individual roll of prepreg shall be provided together with sequence in production and batch identification as supporting data to the prepreg certification. The maximum length of prepreg on any single roll shall be as specified in the purchase order.

3.4.9 <u>Areal Weight</u>. Variation from the nominal areal weight specified on the purchase order shall not exceed + 5 percent.

3.4.10 <u>Storage Life</u>. The prepreg material shall have a storage life of a least nine months when stored in the original package at temperatures of 255°K (0°F) or below. Storage life is defined as the length of time, starting with the date of manufacture, during which the material continues to meet all the requirements of this specification.

3.5 Laminate Properties. The following requirements shall apply when laminates are cured to the procedure specified in 4.5.5.1.

3.5.1 <u>Cured Ply Thickness</u>. The average thickness per ply of the laminate shall be as specified in the purchase order.

3.5.2 <u>Physical Properties</u>. The physical properties of the cured laminate shall conform to the requirements specified in Table 3.

0004S

M131. H. 28 REV 4.72

CODE IDENT. NO.

Property	Requirement	
Specific Gravity	1.57 <u>+</u> .03	
Resin Solids Content, Weight percent	31.0 <u>+</u> 3	
Fiber Content, Volume percent	60 <u>+</u> 2	
Glass Transition Temperature, •C ( *F) minimum (3 specimens)	316 (600)	

	TEST		Graphite Fiber	r
Property	TEMP. C(*F)	Class I	Class II	Class III
Flexural Strength (ultimate) MPa (ksi) Mpa (ksi)	RT 316(600)	1571 (228) 942 (126)	1515 (220) 757 (110)	1394 (202.2) 900 (130.6)
Flexural Modulus GPa (msi) GPa (msi)	RT 316(600)	124 ( 18) 124 ( 18)	117 (17.0) 103 (15.0)	118 (17.2) 108 (15.6)
Short Beam Shear MPa (ksi) MPa (ksi)	RT  316(600)	103 (15) 48 (7)	96 (14.0) 41 ( 6.0)	95 (13.8) 50 ( 7.2)

Note: All values are minimum for the average of the specimens tested with no individual value less then 80 percent of the value listed here. Where the number of specimens is not otherwise specified, five replicates shall be fabricated and tested.

•

0004S

M131.H 28 REV 4.73

CODE IDENT. NO.

NUMBER		 R	EVISIO	IN LET	TER		PAGE			
MB0130-152	A							8		

4.0 QUALITY ASSURANCE

4.1 Qualification

4.1.1 <u>Requirements</u>. Material submitted for qualification to this specification shall be tested to all requirements of this specification. In addition, each prepreg manufacturing facility must be qualified individually. The adequacy of the manufacturing facility may be verified, as deemed necessary, by company representatives, by a survey of such facilities. All requests for qualification shall be directed to the company's Material Department which will request data and samples when desired for qualification purposes.

4.1.1.1 Qualification Samples. The qualification samples shall consist of one representative production sample roll (at least 1.5 kg (3.5 lbs.)) of the particular type from each of a minimum of three separate resin mixes. Each type must be qualified individually.

4.1.1.2 <u>Certified Test Report</u>. The qualification sample submitted for approval shall be accompanied by a certified test report in duplicate which shows that the sample meets the prepreg physical, chemical and composite property requirements of this document. The supplier qualification report shall contain the following:

- (1) Supplier product designation.
- (2) Prepreg type in accordance with this document.
- (3) Test results including individual test values

4.1.1.3 <u>Test Facilities</u>. All suppliers shall have test facilities, or access to test facilities required to test, in accordance with this document, including chemical characterization requirements. The adequacy of test facilities may be verified, as deemed necessary, by a survey by company representatives of such facilities.

4.1.1.4 <u>Demonstration Tests</u>. Qualification testing shall consist of a demonstration of the conformance of the sample supplied in accordance with 4.1.1.1 to all requirements of this document.

4.1.2 <u>Changes.</u> If there is any change in formulation of the material originally qualified to this specification, a new manufacturer's designation shall be assigned and the material shall be resubmitted for qualification.

4.2 Acceptance

4.2.1 <u>Definitions</u>. For purposes of sampling, inspection and maintenance of records the following definitions shall apply.

a. Fiber Lot - Fiber that is produced in one single continuous operation and is separately identified by the supplier.

0004S

M131-H-28 REV. 4-73

$\sim$	IDENT	

			CODE IDE	NT. NO	).								
	NUMBER					REVISI	ON LE	TTER			PAGE		
	L	MB0130-152		A							9		
the pr (ref. lamine ASTM I	<ul> <li>Re</li> <li>Re</li> <li>Pi</li> <li>Pi</li> <li>Si</li> <li>1c</li> <li>us</li> <li>4.2.2</li> <li>repreg</li> <li>3.3),</li> <li>te (re</li> <li>Flexure</li> <li>790.</li> </ul>	esin Batch - S lended in one oll - Any sub- repreg Lot - S roduct an one ingle fiber 10 ot, certifica- sed and placer <u>Testing</u> . Acc will meet sp prepreg (ref ef. 3.5).	The qua mixer -sectio That qu appara ot. Wh tion da ment wi ceptanc ecified . 3.4) t room	ntii and n o: anti tus en : ta s thin rec and tem	ty of separ f a pr ity of from it is shall n the hall b quirem the f	resin representation representation recess docum produ- produ- produ- produ- produ- produ- produ- produ- produ- produ- produ- prosentation restantat	a con y ide g lo preg resin sar, set isar, for ting ad 3.	mpoun enti t. whiin bai y to rela form the tes	nded fied tch is tch a util ated supp neat ts sp (600*	in ( as a projection percond p	one operat such by the oduced as preferrably more than cents of f r certific sin (ref. fied for t when test	ion an e supp a cont y from one f iber lo ation 3.2), he cur ed per	d lier. inuous one iber ots that fiber ed
I F Shipme report number design	nterla ber AS 1.3 <u>Ce</u> ent sta 5 shall 5, batc nation	minar snear a M D 2344. Antification. Ating conformation include this th number, rol and date of n	A cer A cer ance to s speci ll numb manufac	tif: the fice er e ture	t room ied re e requ ation and fo	port ireme numbe otage	from from ents er, in	n the of type eacl	and e sup this and n rol	oplie spec clas	er shall a cification ss, purcha manufactur	wnen to ccompa: • This se ordo er's	ested ny each s er
4	•4 <u>R</u> e	sponsibility	for In	spec	ction	and 1	'est	ing					
4 inspec Intern Rockwe set fo compli	.4.1 etion a pationa oll res orth in ance w	Supplier. The and testing spand use his or serves the right this specification that the the the the the the specification of the	he supp pecifie wn faci ght to ication ation r	lien d he liti peri whe equi	r is r erein ies or form o ere su iremen	espor and m those those ich an its.	nsib nay nes nes re do	le f (wit) f a ( s an; eeme	or th h the comme y of d nec	e pe app crcia the cessa	erformance proval of 1 al laborat inspectionary to ens	of al Rockwe ory. n and ure	l ll testing
. 4 and te comple	•4.2 sts fo te and	Inspection Re or conformance available to	ecords. e to th o Rockw	Tr e re ell	ne sup equire Inter	oplien ements natio	's : of nal	insp thi upon	ectio s <b>spe</b> n req	n re cifi uest	ecords of a liction sha	examina 11 be 3	ation kept
4	•5 <u>Te</u>	st Methods											
4 temper and re	•5.1 ature lative	Standard Cond tests shall h humidity of	ditions be cond 70 per	ucte cent	Jnless ed at t maxi	othe a ter mum.	rwis pera	se sj ature	ecif of	ied 25 t	herein, a to 27°C (7	11 room 5 to 79	⊓ 9°F),
4 conduc aceton a film applic	•5.2.1 ted or e or c will able i	Infrared S a sample of ther appropri- be formed af infrared proce	pectrog neat r iate so ter sol edures.	ram. esir lver vent	The obta nt. D tevap	infr ined eposi orati	ared per t tl	l spe 3.2 ne re Obt	ectro Ex esin tain	grag trac on a the	bhic analy t the res salt pla spectra u	sis sha in with te such sing	all be n C.P. n that
													00045

M131-H-28 REV. 4-73

CODE IDENT. NO. NUMBER **REVISION LETTER** PAGE MB0130-152 A 10 4.5.2.2 High Pressure Liquid Chromatography. This or an equivalent procedure may be used. From a sample of neat resin obtained per 3.2 prepare the sample solutions and inject into the instrument with the liquid chromatograph containing the following columns and settings. Test 1. Experimental Conditions for High Pressure Liquid Chromatographic (HPLC) Analysis of 3, 3', 4, 4'- benzophenonetetracarboxylic acid dimethylester (BTDE), 5 - norboreze - 2, 3 - dicarboxylic acid monomethylester (NE) and Reactions Products in Type I and Type II Polymide Resins. Column: Spectra-Physics Sperisorb ODS Solvents: Baker HPLC water with 0.01 M dihydrogenorthophosphate (KH PO ) at pH-3 Burdisk A & Jackson Acetonitrile Gradient: 10 to 50 percent acetonitrile/water, 15 minutes linear gradient with hold at 50 percent 10 minutes and equilibrate at initial for 10 minutes. Direction: 200 nanometers (nm). 0.4 aufs Flow: 1 milliliter (ml)/minute Sample: 10 micrograms (ul) of 1/5 milligrams per milliliter (mg/ml) of solution Test 2. Experimental Conditions for Ion-Pair Liquid Chromatographic Analysis of Amine Components in LaRC-160 and PMR-15 Polymide Resins. Column: Whatman Partisil 10. ODS-2 Solvents: Baker HPLC water with Waters PIC A Ion Pair Reagent Burdick & Jackson ultraviolet (UV) Grade Tetrahydrofurane (THF) Gradient: 15 to 50 percent THF in water with PIC A, linear 15 min gradient with 15 min hold at 50 percent THF and equilibrate at initial 15 minutes. SP-770, 254 nm, 0.4 aufs Detector: Flow: 1 ml/minute Sample: 10 ul of 1/5 mg/ml solution in THF 4.5.2.3 Record Retention. Copies of all test data and chromatograms shall be retained for a period of three years and shall be made available to Rockwell upon request. 4.5.3 Fiber The propreg supplies may use certification properties supplied by the fiber manufacturer. If such certification is not available, the following procedures shall apply. 0004S

14171 L 79 DEV 4.73

CODE IDENT NO

	CODE IDENT. NO.																
	NUMB	ER			REVISION LETTER								PAGE				
		MBOI	.30 <b>-</b> 152		A									11			
· · ·					<u>.</u>	- <b>1</b>	<b>.</b>				A						
	4.5.3	.1 <u>S</u>	ecific (	ravity	-												
	(1)	Equi	ment														
		(a) (b) (c) (d) (e)	Pycnomet Oven, væ Balance, Inert gæ Vacuum s	ter, mo acuum , analy as (hel: source	del tica ium	930 al, 1 pre	, Be mini ferr	ckma mum ed)	n or sens bott	equ eitiv led	vity vity with	0.00 n rea	Ol gram gulator	1 *•			
	(2)	Obtai	n a repi	resenta	tive	e sai	mple	of	appr	oxin	atel	ly f:	ive (5)	grams	·		
	(3)	Condi 275 <u>-</u>	tion the 5°F for	e sampl r 20 to	e in 30	n a ' min'	vacu utes	um o	ven	oper	ratir	ng a	t full	vacuum	and		
	(4)	Cool	in a des	siccato	r ai	nd w	eigh	to	the	near	rest	0.0	Ol gram	1.			
	(5)	Calil	orate the	e equip	men	t ju	st p	rior	to	use	•		••				
	(6)	Adjus great	st gas pi ter than	ressure 2 psi.	reį Tu	gula urn `	tor vacu	on t um s	he h uppl	elin y or	um te 1.	ank '	to a pr	ressure	no no		
	(7)	With count wheel the s	purge an ter-clock until s side of t	nd coup kwise d startin the cas	lin ire g nu e al	g va ctio umbe bove	lves n to r is the	ope the rea	n, 1 ext ched suri	rota reme (10 .ng h	te bo e pos ocate nandu	oth i sitio ed on wheel	handwhe on. Tu n a pla 1).	els in Irn the Ite aff	n a mea: fixed	suring to	
	(8)	Inser and i	rt previo insert sa	ously c ample c	ond: up :	itio: into	ned pos	and itic	weig n ar	;hed Id se	spec ecure	cimen e it	n into firmly	the sa	mple	cup	
	(9)	Open	purge an	nd çoup	lin	g va	lves	•									
	(10)	Open close	vacuum v valve.	valve a	nd e	allo	w 10	sec	onds	fo	r sys	3 tem	to eva	icuate,	the	n	
	(11)	Open then	gas (hei close ga	lium) v as valv	alvo e.	e an	d al	low	5 ве	con	is fo	or p	ressure	e <b>e</b> quil	libri	um,	
	(12)	Open close	vent va e vent an	lve for nd purg	• 5 a ;e va	seco alve	nds s•	to a	1100	fo:	r pre	essu	re <del>c</del> qui	libriu	am, tl	nen	
	(13)	Wait rota	for 10 a ted a cou	seconds uple of	the time	en l mes)	00se •	n th	e cc	upli	ing v	alv	e (this	shoul	d be		
	(14)	Wait until after this	for 10 a the rearches the who process	seconds ference eel ini	the whe tia	en, eel : lly :	turn stop stop	bot s. s.	h ha Appl Keep	indwi y a the	neels mini e poi	s clo imum inte:	ockwise amount r on th	e simul ; of pr ne scal	taneo ressur .e dur	ously re ring	
	(15)	Wait mark	for 10 s with the	seconds e measu	and rind	d ad g ha:	just ndwh	; poi eel.	nter	by	ima <i>į</i>	ze a	lignmer	nt to t	he ze	ero	
	(16)	Open cent:	coupling	g valve	• ]	Read	spe	cime	n vo	lume	e òn	cou	nter di	.rectly	'in d	ubic	
																00045	

M101 H 00 DEV 4.73

.

()	NUMBER MBC 17) Repe spec 18) Calc Densi 5.3.2 <u>Me</u> ecimen Pr	p130-152 at steps 7 thro imen. ulations: ty = weight of volume rea chanical Proper	A dried	6 unti speci	l co	nsis	tent	readin	PAGE	12 >btain	ed on	each
4.5	MBC 17) Repe spec 18) Calc Densi 5.3.2 <u>Me</u> ecimen Pr	<pre>pl30-152 pat steps 7 thro imen. ulations: ty = weight of volume rea chanical Proper</pre>	A   ugh 1 dried ding	6 unti speci	1 co	nsis	tent	readin	ngs are o	12 >btain	ed on	each
() () 4•!	17) Repe spec 18) Calc Densi 5.3.2 <u>Me</u> ecimen Pr	at steps 7 thro imen. ulations: ty = weight of volume rea chanical Proper	ugh 1 dried	6 unti speci	.l co	nsis	tent	readin	ngs are o	obtain	ed on	each
4.5	18) Calc Densi 5.3.2 <u>Me</u> ecimen Pr	ulations: ty = weight of volume rea chanical Proper	dried ding	speci								
4.5	Densi 5.3.2 <u>Me</u> ecimen Pr	$ty = \frac{\text{weight of}}{\text{volume rea}}$ chanical Proper	dried ding	speci								
4.5	5.3.2 <u>Me</u> ecimen Pr	chanical Proper		obtain	men, ed,	gra cubi	ms c ce:	ntimete	ers			
1	ecimen Pr		ties	of Fib	er							
<u>Spe</u>		eparation										
Ma	terials R	equired:										
	3 piece 4 piece Air cir 4 - 3-i Mochber	s - Stainless S s - Stainless S culating oven, s nch Boston No. g bleeder cloth	teel teel maxim 4 cla No.	plates plate um tem mps. CW-185	22 X 22 X pera	12 1/ 12 1 ture	16-in X 1/4 200	nch (s) 4-inch *C (328	arp edge (to take 3°F)	s rou: 150°	nded) C) (2	38°F))
()	l) Coat	l stainless st	eel p	late o	n on	e si	de w:	ith				
	(a)	2 heavy coats	of Fr	ekote ;	#33 :	relea	ase s	spray a	ind			
	(b)	2 coats of FM-	122 r	elease	spra	ay.						
(2	2) Coat 1 (a	2nd and 3rd st ) and (b).	ainle	ss ste	el pi	late	s on	one si	de only.	Sam	e as	step
(3	3) Dres diam	s drum and work eter rotating c	tabl ylind	e as f er.)	0110	NS:	(Dru	um is ]	.2-inch 1	ong,	9-inc	h
	(a)	Cut a 32-inch	piece	of re	leas	e pa	per.					
	(b)	Tape one end o used) and wrap	f the aróu	paper nd as	to tigh	the a	drum as po	(30-ir ossible	ch circu	mfere	nce d	rum
	(c)	Tape the two en	nds o	f relea	ase j	pape	r to	gether.				
	(d)	Cut a 32-inch	piece	of bl	eede	r paj	per.					
	(e)	Tape the bleed drum as tightl;	er paj y as j	per to possib	the le.	rele	ease	paper	and wrap	aroui	nd th	9
	(f)	Cut a 40-inch i paper.	lengti	h of r	eleas	se pa	aper	and 40	-inch le	ngth d	of blo	eder
	(g)	Place the blee stainless stee	ler p l tab	aper on le•	n toj	o of	the	releas	e paper	and ta	ape to	the
	(h)	Mark off the to	op of	the b	leede	er på	aper	into l	0-inch w	ide se	ection	15.
	(i)	Cut 2-inch and table's edge.	l-in	ch masl	king	tape	e and	l faste	n loosel	y to 1	the	00045

	CODE IDENT, NO.										
NU	MBER REVISION LETTER PAGE										
	MB0130-152 A 13										
(4)	Tape a strip of 2-inch masking tape (sticky side up) across the full width of the drum.										
(5)	Attach one end of the sample to 2 inch masking tape $1/2$ -inch from the left edge of bleeder cloth.										
(6)	Rotate drum at desired speed until one wrap of sample is achieved.										
(7)	Cut sample and attach end to face up masking tape. Identify sample.										
(8)	Place next sample $1/2$ -inch to the right of previous sample and repeat steps 5 thru 7.										
(9)	Once around the drum is generally enough material for 1 (one) strand tensile samples.										
(10)	) Impregnate samples with pre-mixed X506 resin.										
(11)	) Place bleeder cloth over sample and hold ten minutes.										
(12)	) Press very lightly with roller to bleed off excess resin.										
(13)	) Remove the bleeder cloth.										
(14)	) Cut strand at 2-inch masking tape and unwind.										
(15)	) Place each impregnated strand on prepared work table and identify.										
(16)	) Repeat Steps 14 and 15 for each strand.										
(17)	) Place 20-inch sample across 1/16 inch plate.										
(18)	Repeat step 17 for remaining strands, maintaining approximately 1/2-inch spacing between samples.										
(19)	Sandwich sample plate with remaining Stainless Steel plates (coated side to the samples). Note: Do not slide plates against each other because sample could be damaged.										
(20)	) Place the four Boston No. 4 clamps equidistant around the perimeter of the sandwiched assembly.										
(21)	Place the sample assembly into a preheated air circulated oven set @ 150°C (302F). The assembly should be horizontal and supported to permit air circulation on all sides.										
(22)	Hold in oven for two hours.										
(23)	) Remove sample from oven and allow to cool to room temperature.										
(24)	Place in freezer for approximately 15 minutes to ease separation.										
(25)	) Carefully break open assembly and cut cured samples to an 18-inch length and identify. 00045										

M131-H-28 REV 4-73

÷.

MB0130-152       A       A         Specimen Test       A         Apparatus:       Tension testing Machine - a testing machine having a constant-rate-of-crosshead movement with a stationary a         Grips -       grips for holding the test specimen between members should be of the self-aligning type	rAGE 14 nd a movable membra the fixed and mova with a ng 3000 pounds of	er. able										
Specimen Test         Apparatus:         Tension testing Machine - a testing machine having a constant-rate-of-crosshead movement with a stationary a         Grips -       grips for holding the test specimen between members should be of the self-aligning type	nd a movable membra the fixed and mova with a ng 3000 pounds of	er. able										
<u>Specimen Test</u> Apparatus: Tension testing Machine - a testing machine having a constant-rate-of-crosshead movement with a stationary a Grips - grips for holding the test specimen between members should be of the self-aligning type	nd a movable membe the fixed and move with a ng 3000 pounds of	er. able										
Apparatus: Tension testing Machine - a testing machine having a constant-rate-of-crosshead movement with a stationary a Grips - grips for holding the test specimen between members should be of the self-aligning type	nd a movable membe the fixed and move with a ng 3000 pounds of	er. able										
Tension testing Machine - a testing machine having a constant-rate-of-crosshead movement with a stationary a Grips - grips for holding the test specimen between members should be of the self-aligning type	nd a movable member the fixed and move with a ng 3000 pounds of	er. able										
Grips - grips for holding the test specimen between members should be of the self-aligning type	the fixed and move with a ng 3000 pounds of	able										
pneumatic-hydraulic action capable of applyi pressure to the jaws.												
Jaws: 2 by 1-inch rubber faced jaws are recommende indicating their center.	d; with edge mark:	ings										
Specimen: Cured impregnated tows ten inches in length.												
Testing Procedure:												
(1) Set speed of testing machine at 1.27 millimeters	(0.05 in.)/minute	•										
(2) Set the load scale range as follows:												
3K = 200 pounds 6k = 500 pounds												
(3) Set the jaws at an effective gage length of 127 m apart.	illimeters (5 inc)	nes)										
(4) Adjust recorder speed to lay curve at approximate	ly 45°.											
(5) Sandwich one end of the specimen between aluminum and place in upper jaws, aligning it with centeri	oxide cloth (320 ng marks on jaws.	gri										
(6) Repeat Step 5 for lower jaws.												
(7) Apply approximately 2000 pounds of pressure to ja specimens should break within the 5-inch gauge le occurs apply more pressure, if specimens continua reduce pressure.	ws. Majority of ngth, if slippage lly break at jaws											
(8) Manually remove any positive or negative load app jaws.	lied to specimen b	bу										
(9) Attach 2-inch extensometer to specimen and connec drive so that the abcissa of the stress-strain cu strain, directly.	t to Instron char rve is fractional	t										
(10) Start test.												
(11) Test all specimens with extensometer for modulus breaking load to calculate strength.	determination and	use										
		~										

378

44191 L 98 DEV 4.73

CODE IDENT. NO.

· · · · · · · · · · · · · · · · · · ·	NUMB	REVISION LETTER     PAGE       MB0130-152     A     15
	Calcu	lations
	Cross	-sectional area, square inches = $.172 \times 10^{-6} \times \text{Denier/Density}$
	Denie Densi	r = original yarn denier of specimen ty = fiber density grams per cubic centimeter $(g/cm^3)$
	Modul	us, psi = (Load)/(Area X strain)
	Stren	gth, psi = (Load)/(Area)
	Strai	n, in/in = (Deflection)/(Gauge length)
	4.5.3	-3 Thermal Oxidation Resistance
	(1)	Weigh a clean dry 2-inch diameter $1/2$ -inch high wall aluminum weighing dish. Record the weight as $W_1$ .
	(2)	Put on polyethylene gloves.
	(3)	Roll approximately 3 grams of clean dry fiber into a roll so that it fits snugly into the weighing dish. Bend the wall of the dish in slightly to hold the coil of fiber in position.
	(4)	Weigh the dish and fiber. Record the weight as $W_2$ .
	(5)	Place the weighing dish containing the fiber in a hot block oven heated to $316C (600 + 10^{\circ}F)$ or equivalent. There should be only a slight air flow.
	(6)	Leave for 130 + 10 minutes.
	(7)	Remove from the oven, cool in a desiccator and weigh. Record weight as $W_3$ .
		Note. This should be the start - weight where any combined moisture, sizing or other non fiber contaminates are driven off leaving a pyrolytically clean fiber.
	(8)	Place the weighing dish with the specimen back in the oven heated to $316^{\circ}C$ (600°F) and heat for 168 <u>+</u> 1 hours.
	(9)	Remove the weighing dish with the specimen from the oven. Place in a desiccator to cool. Weigh and record the weight as $W_4$ .
•		
		0004s
14121 U 30 DEV 4		·

		CO	DE IDE	NT. NO.										
	NUM	NUMBER				P	EVISIO	N LET	TER		PAGE			
·····		MB0130-152		A								16		
	(10)	Calculations: Percent Weight L	098	= ( W	W3 -	• W ₄	) X (	100						
		W _l = weight of	wei	.ghi	ng d	lish								
		<ul> <li>W₂ = weight of weighing dish and non-pyrolytically cleaned fiber (includes moisture and sizing)</li> <li>W₃ = weight of weighing dish and pyrolytically cleaned fiber (moistu and sizing free)</li> <li>W₄ = weigh of weighing dish and fiber thermally oxidized for 168 hour at 600°F in air.</li> </ul>												<b>n</b> .
														isture
														hours
	(11)	Test shall be co recorded. All s	nduc peci	ted men	on Is sh	thro	ee sj mee1	eci: th	mens e re	wit quir	h th emer	ne average nts specifi	and mi ed in	nimum 3.3.
	4.5.4	Prepreg												
	4.5.4	.1 Fiber Content	• T	his	or	an e	equiv	ale	nt p	roce	dure	e may be us	ed.	-
	(1)	Prepare nominal convenience the narrow strips.	3-in spec Remo	ch ime ve	squa n ma rele	re s y be ase	speci e cu pape	.men t (b er b	s of efor efor	prej e we: e an:	preg ighi alyz	g. For han ng) into s ing.	dling everal	
	(2)	Determine the ar 0.01 square inch used in determin	ea o • R ing	f e eco the	ach rd a fib	nomi s fi er a	inal iber areal	3 X are: we:	3-in a (A ₁ ight	nch ( f).	spec Thi	imen to an s value wi	accur 11 als	acy of o be
	(3)	Obtain a clean d milligram and re	ry e: cord	xtr as	acti W _l .	on 1	thimb	le a	and w	veig	h to	the neare	st 0.1	
		NOTE: May be pu glass ext Catalog N	rcha ract o. 2	sed ion 774	fro thi 3-12	m Va mble O.	in Wa es, D	tera	s&] um E	Roge: •C•	rs C 35 X	onpany as 90 millim	fritte eters,	đ
	(4)	Place specimen in milligram. Reco	n th: rd a:	imb s W	le ( 2•	s tej	2)	and	wei	gh to	o th	e nearest	0.1	
	(5)	Place thimble and extraction thimble agitation for 30 basis of being all of the test. No:	d spo le, a min ble rmal	ecia and utes to ly n	men let s. diss meth	in t set The olve yl e	eake tat solv the thyl	r an roon ent res ket	nd ac n ter used sin c tone	ld en npera l sha compl is a	noug atur all Lete suit	h solvent e with int be selecte ly under t able.	to cov ermitt d on t he con	er the ent he dtions
	(6)	Remove extraction drain. Discard w	n thi used	imb so	les lven	with t an	n spe nd ri	cime nse	en fi beal	com 1 cer 1	eak vith	er of solv fresh sol	ent and vent.	đ
	(7)	Place extraction Repeat steps 5 an apart.	thin nd 6	nblo un	es i: til :	nsid solv	le be rent	aken is v	r and Visua	l cov ally	ver cle	with fresh an and fib	solve ers sta	nt. and
														00045

41191 LL 90 DEV 4.73
·····	CODE IDENT. NO.	
NUM	BER REVISION LETTER PAGE	
<u> </u>	MB0130-152 A 17	<u> </u>
(8)	Following the last extraction, remove extraction thimble with s from beaker of solvent and place in a rubber crucible holder on filter flask capable of maintaining a vacuum of at least five is mercury. Drain free of solvent and rinse once more with fresh	pecim a va nches solve
(9)	Dry for 30 minutes at 150 to 160C (300 to 320F) in a mechanical convection oven or, alternatively, for 15 minutes at 55 to 65C (130 to 150F) in a vacuum oven.	•
(10)	Remove from oven and cool to room temperature in a desiccator.	
(11)	Weigh each extraction thimble and specimen to nearest 0.1 milli,	gram.
(12)	Record test specimen weight as W3.	
(13)	Calculations and Report of Results	
	Calculate fiber content as follows:	
	Weight percent fiber = $\frac{(W_3 - W_1) \times 100}{W_2 - W_1}$	
	Where: $W_1$ = weight of extraction thimble, grams	
	$W_2$ = weight of extraction thimble plus specimen, in g	rams
	$W_3$ = weight of extraction thimble plus test sample af extraction, in grams	ter
4.5.4	4.2 Areal Weight	
	$A (g/M^2) = \frac{W_{f} X 1550}{A_{f}}$	
Where	: A = fiber areal weight (grams per square meter)	
•	$W_f$ = total weight in grams (g) of fiber in the prepreg sample, $W_3 - W_1$ .	,
	$A_{f} = 1$ fiber area in square inches (in ² ) from 4.5.4.1(2).	
	1550 = conversion factor of grams per square inch $(g/in^2)$ to grams square meter $(g/M^2)$	ams p
	3 Valatilas Content	
4.5.4.	volatiles content	
<b>4.5.4</b>	Obtain samples of the prepreg weighing 2.0 to 4.0 grams.	

M131-H-28 REV. 4-73

CODE IDENT. NO. **REVISION LETTER** NUMBER PAGE MB0130-152 18 A (3) Place the sample and dish in an air circulating oven pre-heated to 232°C (450°F) and hold at constant temperature for 30 minutes. (4) Remove dish from the oven and weigh. (5)Calculate volatile content as follows:  $W_1$  = weight of dish  $W_2$  = weight of dish plus prepreg sample before heating  $W_3$  = weight of dish plus prepreg sample after heating Volatile Weight percent =  $\frac{W_2 - W_1}{W_2 - W_1}$ 4.5.4.4 Prepreg Resin Content - Dry Analysis Method. Calculate dry resin content using data obtained in the preceeding paragraphs. Percent Resin Content, Wet = 100 - weight percent Fiber - weight percent Volaties Percent Resin Content, Dry = Weight percent Wet Resin X 100 weight percent fiber + weight percent Wet Resin 4.5.4.5 Gel Time Apparatus: (1)Fisher-Johns melting point apparatus or equivalent to read +1°C (1.8°F) of the specified temperature. (2)Thickness No. 2 cover glasses (18 millimeters) or equivalent. (3) Timer (4) Wooden picks Procedure: (1)Preset the melting point apparatus to 204°C (400°F). (2) Insert a 1/4 inch by 1/4 sample between two cover glasses and place in the melting point apparatus. Start the timer and probe the specimen with a wooden pick. (3) (4) Resin gel is evidenced when no resin movement is seen when moderate pressure is applied to the specimen. At this point stop the timer and report the gel time to the nearest 0.1 minute. 0004S

	C	ODE IDENT. NO	0.						_
	NUMBER			REVISIO	N LETT	ER	T	PAGE	]
<u> </u>	MB0130-152	A						19	L
produc checko full v approj	4.5.5 <u>Cured Laminat</u> 4.5.5.1 <u>Preparation</u> be the specimens require the bag and seal vacuum. Correct any priate process as des	of Cured nired by system t degradat scribed b	Lam the o be ion. elow	inate. tests i used t Relea :	Lay n thi o cur se th	up the is spec re thes ne vacu	lami ifica e spe um an	inates neces ation. Afte ecimens by a nd commence	sary to r assembly, pplying the
2	Staging Procedure								
	Types I & II Mate:	rial						·	
	<ol> <li>Make lay-1</li> <li>Install 14</li> <li>Apply 2 tr (1-3°C) (1</li> <li>Hold at tr (5) Reduce th vacuum or (6) Remove lay</li> </ol>	up for st ay-up ass b 15 inch 2 to 5°F) emperatur e tempera removing minate fr	agin embl; es o per e fo ture fro om 1	g as sh y in ai f mercu minute r 30 mi to 65° m oven. ay-up a	own i r cir ry va to 2 nutes C (15 nd ir	n Figu culati acuum a 218 <u>+</u> 3 minim 50°F) o nspect	re l. ng ov nd ra °C (4 oum. °r low	ven. hise the tem 425 <u>+</u> 5°F). wer before r	perature at eleasing
	i								
	• :-								
SEALANT	IL KAPTON FILM	VATE		- 2 PLIE /16 INC 	S 162 H PEI LY [®] TH SEPA LY [®] TH PTON DROUS	2 GLASS RFORATE IN MOO RATOR - UN MOO FILM O TEFLON	FABI ED CAI HBERG TEFI HBERG DR N GLA	VIC JL PLATE CLOTH LON COATED G CLOTH SS FABRIC	LASS CLOTH
	• NOTE- THE AMOUNT	OF MOCHB	ERG	cloth a	ND VA	A MUDA	PPLIE	Ð	
	<b>DURING STAGIN</b>	G SHALL B	e ad	JUSTED	as ri	EQUIREI	) TO	ACHIEVE	
	A CURED LAMIN	ATE FIBER	VOL	ume of	601 1	ased u	PON		
	CERTIFIED PRE	REG VOLA	TILE	AND RE	SIN C	ONTENT	•		
	FIGURE 1. La	yup Assem	bly	for Sta	ging	Туре І	: & I:	I Naterials	00045



M111.H.78 REV 4.73

**Rockwell International Corporation** 



	MINEK			REVISION	LETTER			PAGE		7	
	MB0130-152	A						raut.	22		
Postcurin	g Procedure										
<u> </u>	ypes I & II Materi	al									
	These mater excess of 3 is found ne shall be ut	rials nor 516°C (60 ecessary 511ized.	mally OF) af to pos	exhibi Ster cu stcure	t tra: ring j lamina	nsiti per p ates,	on t rece the	emper ding foll	ature proce owing	s (Tg) dure. proce	in If dure
	<ol> <li>Install lam</li> <li>Increase te</li> <li>Hold at 316</li> <li>Reduce temp oven.</li> </ol>	minate in emperatur 5 <u>+</u> 9°C ( perature	air c e to 3 600 <u>+</u> to 65•	circula 516 + 9 5°F) f C (150	ting ( *C (60 or 16 *F) o:	oven 00 <u>+</u> hour r low	in f 5°F) s er b	ree s efore	tandi: remo	ng con ving f	diti rom
4 measured represent ultrasoni readings	•5.5.2 <u>Cured Ply</u> to the nearest O.C ative locations us c gage. The thick and dividing by th	Thicknes 25 milli sing a do mess per ne number	meter ubleba ply s of pl	ne thic (0.001 11 3-m Shall b ies.	kness inch illime e comj	of t ) in eter puted	he c at 1 (1/8 by	ured east R) m avera	lamina five icroma ging	ate shi eter o: the fi	all r ve
4 accordanc	•5•5•3 Specific G e with Federal Tes	ravity. It Method	The s No. 4	pecifi 06, Me	c gray thod 5	<b>vity</b> 5011.	shal	l be	deter	nined :	in
4 ligestion Ca	.5.5.4 Fiber Cont method shall be u ution: Appropriat	ent. Ei sed. e safety	ther t	he acions	d dige	estio be o	n or bser	the ved u	hydra: sing (	zine either	
4 ligestion <u>Ca</u>	.5.5.4 Fiber Cont method shall be u ution: Appropriat method.	ent. Ei sed. e safety	ther t preca	he acio Autions	d dige must	estio be o	n or bser	the ved u	hydra: sing (	zine either	
4 Higestion <u>Ca</u> <u>Acid</u>	.5.5.4 Fiber Cont method shall be u ution: Appropriat method. Digestion	ent. Ei sed. e safety	ther t preca	he acio	dige must	estio be o	n or bser	the ved u	hydra: sing (	zine either	
4 Higestion <u>Ca</u> <u>Acid</u> (1)	.5.5.4 Fiber Cont method shall be u ution: Appropriat method. Digestion The composite sp	ent. Ei sed. e safety	ther t preca	he acid	d dige must	be o	n or bser	the ved u deter	hydra: sing ( minat	zine either ions p	er
4 ligestion <u>Ca</u> <u>Acid</u> (1)	.5.5.4 Fiber Cont method shall be u ution: Appropriat method. Digestion The composite sp 4.5.5.3 may be u	ent. Ei sed. e safety ecimens	ther t preca used f	he acions Nutions for spe	d dige must	estio be o grav	n or bser ity	the ved u deter	hydra: sing ( minat:	zine either ions po	er
4 ligestion <u>Ca</u> (1) (2)	.5.5.4 Fiber Cont method shall be u ution: Appropriat method. Digestion The composite sp 4.5.5.3 may be u Obtain a clean d containing nitri 10F). Wash with	eecimens sed. ry extra c acid f	ther t preca used f ction or a m ed wat	the acid tutions for spe thimble inimum ser, dr	d dige must cific e or c of or y in c	grav grav clean he ho	n or bser ity the ur <b>a</b> at 1	the ved u deter thim t 149 21 <u>+</u>	hydra: sing ( minat: ble in ± 5.1	zine either ions po n a ben 5°C (36 (250 <u>+</u>	er aker 00 <u>+</u> 10F
4 Aigestion <u>Ca</u> (1) (2)	<ul> <li>.5.5.4 Fiber Cont method shall be u ution: Appropriat method.</li> <li>Digestion</li> <li>The composite sp 4.5.5.3 may be u Obtain a clean d containing nitri lOF). Wash with desiccate and co Note: May be pu extractio 27743-120</li> </ul>	ecimens sed. ry extra c acid f distill ol. rchased n thimble	ther t preca used f ction or a m ed wat from V es, me	be acid utions for spe thimble inimum er, dr an Wate dium E	d dige must cific e or c of or y in c ers &	grav clean he ho oven Roge 5 X 9	n or bser ity the ur a at 1 rs C 0 mi	the ved u deter thim t 149 21 <u>+</u> o. as llime	hydra: sing ( minat: ble in 55°C ( frit ters,	zine either ions po n a bes 5°C (30 (250 <u>+</u> ted gla Catalo	er aker 10F ass og N
4 ligestion <u>Ca</u> (1) (2) (3)	.5.5.4 Fiber Cont method shall be u ution: Appropriat method. Digestion The composite sp 4.5.5.3 may be u Obtain a clean d containing nitri 10F). Wash with desiccate and co Note: May be pu extractio 27743-120 Weigh each extra	ecimens sed. ry extra c acid f distill ol. rchased n thimbl ction th	ther t preca used f ction or a m ed wat from V es, me imble	he acid outions or spe thimble inimum ser, dr dan Wat dium E to the	d dige must cific e or o of or y in o ers & .C. 35 neare	grav clean Roge X 9 est 0	n or bser ity the ur a at 1 rs C 0 mi .1 m	the ved u deter thim t 149 21 <u>+</u> 0. as llime illig	hydra: sing ( minat: ble in ± 5. 55°C frit ters, ram an	zine either ions per 5°C (30 (250 <u>+</u> ted gla Catalo	er 00 <u>+</u> 10F ass og N
4 ligestion <u>Ca</u> (1) (2) (3) (4)	.5.5.4 Fiber Cont method shall be u ution: Appropriat method. Digestion The composite sp 4.5.5.3 may be u Obtain a clean d containing nitri lOF). Wash with desiccate and co Note: May be pu extractio 27743-120 Weigh each extra "W1". Dry the specimen each of three in milligroup Bocc	eecimens sed. ry extra c acid f distill ol. rchased n thimbl ction th s used i: a clean	ther t preca used f ction or a m ed wat from V es, me imble n the extra	be acid butions for spe thimble inimum er, dr dium E to the specif: ction	d dige must cific e or o of or y in o ers & .C. 35 neare ic gra thimbl	grav clean he ho oven Roge 5 X 9 est 0 avity le an	n or bser ity the ur a at 1. rs C O mi .1 m det	the ved u deter thim t 149 21 <u>+</u> o. as llime illig ermin igh t	hydra: sing ( minat: ble in ± 5. 55°C frit ters, ram an ation o the	zine either ions po 5°C (30 (250 <u>+</u> ted gla Catalo nd reco and pl neares	er 10F ass bog N bord lace
4 Aigestion <u>Ca</u> (1) (2) (3) (4) (5)	.5.5.4 Fiber Cont method shall be u ution: Appropriat method. Digestion The composite sp 4.5.5.3 may be u Obtain a clean d containing nitri lOF). Wash with desiccate and co Note: May be pu extraction 27743-120 Weigh each extra "W1". Dry the speciment each of three in milligram. Reco Place thimble an a magnetic stirr	eecimens sed. ry extra c acid f distill ol. rchased n thimbl ction th s used i: a clean ord as W ₂ d speciming bar,	ther t preca used f ction or a m ed wat from V es, me imble n the extra en in and a	be acid utions for spe thimble inimum er, dr an Wate dium E to the specif: ction a beak dd con	d dige must cific e or o of or y in o ers & .C. 35 neare ic gra thimbl er fit	grav clean he ho oven Roge 5 X 9 est 0 hvity le an tted	n or bser ity the at l .1 m det d we with	the ved u deter thim t 149 21 <u>+</u> o. as llime illig ermin igh t a ra uric	hydra: sing ( minat: ble in ± 5. 55°C ( frit ters, ram a) ation o the ised ] acid (	zine either ions p n a bea 5°C (30 (250 <u>+</u> ted gla Catalo nd reco and pl neares platfon mtil 1	er aker 10F ass bg N bord lace st O rm a
4 Aigestion <u>Ca</u> (1) (2) (3) (4) (5) (6)	<ul> <li>.5.5.4 Fiber Cont method shall be u ution: Appropriat method.</li> <li>Digestion</li> <li>The composite sp 4.5.5.3 may be u Obtain a clean d containing nitri lOF). Wash with desiccate and co Note: May be pu extraction 27743-120</li> <li>Weigh each extra "W1".</li> <li>Dry the speciment each of three in milligram. Reco Place thimble and a magnetic stirr specimen is cove Remove thimble a beaker and repeat</li> </ul>	eecimens sed. ry extra c acid f distill ol. rchased n thimble ction th s used in a clean ord as W ₂ d speciming bar, red. Br nd decan t Step 5	ther t preca used f ction or a m ed wat from V es, me imble n the extra en in and a ing sl t spen	he acid utions for spe thimble inimum er, dr an Wate dium E to the specif: ction a beak dd con owly to t sulf	d dige must cific e or o of or y in o ers & .C. 35 neare ic gra thimbl er fit centra o a bo uric a	grav clean he ho oven Roge 5 X 9 est 0 hvity le an tted ited oil a: acid.	n or bser ity the at l .1 m det d we with nd h Re	the ved u deter thim t 149 21 <u>+</u> o. as llime illig ermin igh t a ra uric old f place	hydra: sing ( minat: ble in <u>+</u> 5. 55°C frit ters, ram an ation o the ised 1 or 30 thim	zine either ions pun n a best 5°C (30 (250 <u>+</u> ted gla Catalo nd reco and pl neares platfor minute ole in	er aker 10F ass bg N lace st O rm a the ss sam
4 ligestion <u>Ca</u> (1) (2) (3) (4) (5) (6)	.5.5.4 Fiber Cont method shall be u ution: Appropriat method. Digestion The composite sp 4.5.5.3 may be u Obtain a clean d containing nitri lOF). Wash with desiccate and co Note: May be pu extractio 27743-120 Weigh each extra "W1". Dry the specimen each of three in milligram. Reco Place thimble an a magnetic stirr specimen is cove Remove thimble a beaker and repea	ecimens sed. ry extra c acid f distill ol. rchased n thimbl ction th s used i: a clean ord as W ₂ d specim ing bar, red. Br nd decan t Step 5	ther t preca used f ction or a m ed wat from V es, me imble n the extra en in and a ing sl t spen	he acid autions for spe thimble inimum fer, dr an Wate dium E to the specif: a beak dd cond owly to t sulf	d dige must cific e or o of or y in o ers & .C. 35 neare ic gra thimbl er fit centra o a bo iric a	grav clean he ho oven Roge 5 X 9 est 0 hvity le an tted hited oil a: acid.	n or bser ity the ur a at 1 .1 m det d we with sulf nd h Re	the ved u deter thim t 149 21 <u>+</u> o. as llime illig ermin igh t a ra uric old f place	hydra: sing ( minat: ble in ± 5. 55°C frit ters, ram an ation o the ised ] acid to or 30 thim]	zine either ions p 5°C (30 (250 + ted gla Catalo nd reco and pl neares platfor minute ole in	er aker 00 <u>+</u> 10F ass bg N bord Lace st 0 rm a the es. sam
4 Higestion <u>Ca</u> (1) (2) (3) (4) (5) (6)	.5.5.4 Fiber Cont method shall be u ution: Appropriat method. Digestion The composite sp 4.5.5.3 may be u Obtain a clean d containing nitri lOF). Wash with desiccate and co Note: May be pu extractio 27743-120 Weigh each extra "W1". Dry the specimen each of three in milligram. Recc Place thimble an a magnetic stirr specimen is cove Remove thimble a beaker and repea	ecimens sed. ry extra c acid f distill ol. rchased n thimbl ction th s used i: a clean rd as W ₂ d specim ing bar, red. Br nd decan t Step 5	ther t preca used f ction or a m ed wat from V es, me imble n the extra en in and a ing sl t spen	he acid utions for spe thimble inimum er, dr an Wat dium E to the specifiction a beak dd cond owly to t sulfu	d dige must cific e or o of or y in o ers & .C. 35 neare ic gra thimbl er fit centra o a bo uric a	grav grav clean he ho oven Roge 5 X 9 est 0 avity le an tted acid.	n or bser ity the at l .l m: .l m: .l m: det d we with sulf nd h Re	the ved u deter thim t 149 21 <u>+</u> o. as llime illig ermin igh t a ra uric old f place	hydra: sing ( minat: ble in 55°C ( frit ters, ram an ation o the ised ] acid ( or 30 thim)	zine either ions p n a be 5°C (36 (250 <u>+</u> ted gla Catalo nd reco and pl neares platfor minute ole in	er aker 10F ass bog N bord lace st O rm a the ss. sam

CODE IDENT. NO.

	NUMB	R .	V		T	R	EVISIO	ON LET	TER			PAGE	_	
		MB0130-	152		A	T							23	
							•							
(	(7)	Continue complete require <u>Note</u> : C i	e boil ely de addii Comple its ap	ing un ecompos cional ete dig opearan	til fi ed. T sulfur estion ce fro	bers his i ic ac is i m a u	are s de id. ndic niti	comp term ated zed :	lete ined whe mass	ly so by n the to 1	epar visu e te loos	rated an mal exam est spec se, soft	d the inati imen fibe	resin is on and may changes ers which
_		h	ave a	tende	ncy to	sink	to	the	bott	om o:	f tł	ne thimb	le.	
(	(8)	After di cool bel	.gesti .ow 30	on, pl OF.	ace on	a ma	gnet	ic s	tirr	er, i	stir	slowly	and	allow to
(	.9)	While st the hot Caution:	irrin solui All	ig, <u>car</u> ion. low the	efully hvdro	add gen p	10 p erox	erce	nt b to r	y vo: un de	lume own	hydrog the sid	en pe e of	roxide to the
			bea	iker.	Add ve	ry sl	owly						•	
(	(10)	Continue transpar invalid	e addi ent c and s	ng hyd lear c bhould	rogen ; olor. be rep	perox If a eated	ide cle	unti ar c	l th olor	e ac: is i	id a not	olution obtaine	turn d, th	s a e test is
(	(11)	Allow ac	id to	diges	t thre	e mor	e mi	nute	s.					
(	(12)	Remove e crucible	extrac hold	tion t er on	himble a vacu	from um fi	the lter	aci fla	d, d sk a	rain nd wa	, pl ash	ace in fibers	a rub with	ber distilled
(	(13)	Remove 1 10F for	til i thimbl a mir	ree of .e cont nimum o	acid a aining f 30 m	as sn fibe inute	own TSA S.	by n nd d Cool	eutr ry i in	ai 11 n an a de:	naic ove sicc	ation o en maint ator an	i ph ained d wei	at 300 + gh to the
(	(14)	nearest Calculat	0.1 m te fil	illigr er con	am. R tent a	ecord ccord	wei ing	ght to t	as" hef	W3". Ollo	wine	g equati	on:	
		Fit	er co	ontent,	weigh	t per	cent	; = V V	3 - 1 -	W ₂ W ₂	,	100		
		Where:	W ₁	= we ac	ight o id dig	f ext estio	ract n in	ion gra	thim ms	ble j	plus	s test s	pecim	en before
			W ₂ -	• we	ight o	f ext	ract	ion	thim	ble _: :	in e	rams		
			₩3 *	• we di	ight o gestio	f ext n in	ract gram	ion 18•	thim	ble j	plus	s specim	en af	ter acid
H	lydraz	ine Dige	estion	1										
(	(1) (2)	A cured one-half the near Place th approxim hydrazim Cautions	lamin f incl rest ( ne sam nately ne hyde	nate wi n) and 0.1 mil nple in y 40 mi 1rate.	th a n a nomi ligram a 400 llilte	omina nal w . Re mill rs of invol	l si eigh cord ilit rea	ze o it of wei ers gent hvd	f 2. .5 ght wide gra razi	5 X : to . as W mou de au ne sl	1.75 8 gi 1. th H nhyd	5 centim ram shal Erlenmey frous hy be per	eters 1 be er fl drazi	(one by weighed to ask & add ne or d in a
· .		<u> </u>	fur esc pre sha con the	e hood caping. event a all be ndensed work	with The ony con config liqui area.	an ai opera tact ured ds or	rflo tor of t to p oth	shal shal he h recl ter c	equa l we ydra ude onta	te t ar a zine drop mina	o pi dequ to ping nt i	revent a late rub the ski g of rus into the	ny fu ber g n. T t, du flas	mes loves to he hood st, k or on
														00045

CODE IDENT. NO.

	NUMBE	MB0130	-152					PAGE	24	
			-1)2	~ <u></u>			_ <u></u>		24	
	(3)	Heat un	til the	mate:	rial is co	moletelv	degraded a	as indicat	ed bv	the
		separat	ion of	fiber	s free of	anv polv	imide resi	n. Do not	allow	the
		mixture	to go	to dry	vness.	ung porg				•
	(4)	Obtain	a clean	drv d	coarse gra	de fritt	ed ølass f	ilter and	weigh.	to the
		nearest	0.1 mi	lligra	am. Recor	d weight	as Wo.			
	(5)	Filter	the dig	rested	material	through	the weighe	d. coarse	grade.	fri tte
		glass f	ilter.	,				-,	,	
	(6)	Rinse t	he fibe	ers wi	th ethanol	until t	he washing	s are cold	rless.	
	(7)	Oven dr	<b>y</b> at 90	) to 10	00°C (194	to 212°F	).			
	(8)	Cool in	a desi	.ccato	r and weig	h the fi	lter and co	ontents to	the no	earest
		0.1 mil	ligram.	Reco	ord weight	as Wz.				
	(9)	Calcula	te fibe	r cont	tent accor	ding to	Step 14 of	acid meth	od.	
	4.5.5.	5 Fibe	r Conte	nt, Vo	olume Perc	ent. Ca	lculate the	e fiber vo	lume a	ccordin _é
to the	e foll	owing f	ormula:							
		Fiber v	olume,	percer	$nt = \frac{W^{f}}{\Sigma f}$	f XΣc	X 100			
		Where:	$w^{f/f}$	= 1	fiber weig	ht fract	ion per 4.	5.5.4.		
			<b>۲</b> ۴		longity of	fiham n				
				- (	TEURICY OT	TIDEL b	er 4•9•9•1	•		
					v					
	4.5.5.	6 Glas	Σc s Trans	= d	lensity of Temperatu	composi	te per 4.5. glass trai	.5.3. nsition te	mperati	ure of
the cu Analy: Measu: load.	4.5.5. ured c zer or rement Test	6 <u>Clas</u> omposit equiva s will s shall	Σc s Trans e shall lent. be made be con	= d ition be de The he on a ducted	density of <u>Temperatu</u> stermined sating rat 100 mil d d on lamin	composi <u>tre</u> . The using a e shall iameter ates tha	te per 4.55 glass tran DuPont 941 be 5 + 0.5 expansion p t have been	.5.3. TMA model C per min probe unde n cured pe	mperati /900 Th ute. r a 5-4	ure of hermal gram 5.1.
the cu Analy: Measu: load.	4.5.5. cured c zer or rement Test	6 <u>Clas</u> omposit equiva s will s shall	Σc s Trans e shall lent. be made be con	= d ition be de The he on a ducted	density of <u>Temperatu</u> stermined sating rat 100 mil d d on lamin	composi ure. The using a e shall iameter ates tha	te per 4.55 glass tran DuPont 941 be 5 + 0.55 expansion p t have been	.5.3. msition te TMA model C per min probe unde n cured pe	mperata /900 Ti ute. r a 5-4 r 4.5.5	ure of hermal gram 5.1.
the cu Analy: Measu: load.	4.5.5. cured c zer or rement Test 4.5.5.	6 <u>Glas</u> omposit equiva s will s shall 7 Shor	Σc s Trans e shall lent. be made be con t-Beam	= d ition be de The he ducted Shear	density of <u>Temperatu</u> etermined eating rat 100 mil d d on lamin . The int	composi re. The using a e shall iameter ates tha erlamina	te per 4.5. glass tran DuPont 941 be 5 + 0.5 expansion p t have been r shear str	.5.3. msition te TMA model °C per min probe unde n cured pe rength sha	mperata /900 Tl ute. r a 5-4 r 4.5.5	ure of hermal gram 5.1.
the cr Analy: Measu: load. deter	4.5.5. ured c zer or rement Test 4.5.5. mined	6 <u>Glas</u> omposit equiva s will s shall 7 <u>Shor</u> per MIL	Σc s Trans e shall lent. be made be con t-Beam -G-8341	= d ition be de The he on a ducted Shear	density of <u>Temperatu</u> etermined eating rat 100 mil d d on lamin . The int th a minim	composi re. The using a e shall iameter ates tha erlamina um half-	te per 4.5. glass tran DuPont 941 be 5 + 0.5 expansion p t have been r shear str hour soak a	.5.3. TMA model C per min probe unde n cured pe rength sha at each te	mperata /900 Ti ute. r a 5-4 r 4.5.5 ll be st tem	ure of mermal gram 5.1. peratur
the cr Analy: Measu: load. determ prior	4.5.5. ured c zer or rement Test 4.5.5. mined to lo	6 <u>Glas</u> omposit equiva s will s shall 7 <u>Shor</u> per MIL ading.	Σc s Trans e shall lent. be be made be con t-Beam -G-8341	= c ition be de The he on a ducted <u>Shear</u> 0, wit	density of <u>Temperatu</u> etermined eating rat 100 mil d d on lamin . The int th a minim	composi re. The using a e shall iameter ates tha erlamina um half-	te per 4.5. glass tran DuPont 941 be 5 + 0.5 expansion p t have been r shear str hour soak a	.5.3. msition te TMA model 'C per min probe unde n cured pe rength sha at each te	mperata /900 Ti ute. r a 5-4 r 4.5.5 ll be est tem]	ure of hermal gram 5.1. perature
the ci Analy: Measu: load. deter prior	4.5.5. ured c zer or rement Test 4.5.5. mined to lo	6 <u>Glas</u> omposit equiva s will s shall 7 <u>Shor</u> per MIL ading.	Σc s Trans e shall lent. be be made be con t-Beam -G-8341	= c ition be de The he on a ducted <u>Shear</u> 0, wit	density of <u>Temperatu</u> etermined eating rat 100 mil d d on lamin . The int th a minim	composi re. The using a e shall iameter ates tha erlamina um half-	te per 4.5. glass tran DuPont 941 be 5 + 0.5 expansion p t have been r shear str hour soak a	.5.3. msition te TMA model °C per min probe unde n cured pe rength sha at each te	mperata /900 Tl ute. r a 5-4 r 4.5. ll be st tem]	ure of nermal gram 5.1. perature
the cu Analys Measu: load. detern prior be det tempe:	4.5.5. ured c zer or rement Test 4.5.5. to lo 4.5.5. termin rature	6 <u>Glas</u> omposit equiva s will s shall 7 <u>Shor</u> per MIL ading. 8 <u>Long</u> ed per 1 prior	Σc s Trans e shall lent. b be made be con t-Beam -G-8341 itudina MIL-G-8 to load	= d ition be de The he on a ducted Shear 0, with 1 Fley 3410, ing.	density of <u>Temperatu</u> etermined eating rat 100 mil d d on lamin . The int th a minim <u>rural Test</u> with a mi	composi re. The using a e shall iameter ates tha erlamina um half- . The f nimum ha	te per 4.5. glass tran DuPont 941 be 5 + 0.5 expansion j t have been r shear str hour soak a lerural str lf-hour soa	.5.3. msition te TMA model °C per min probe unde n cured pe rength sha at each te rength and ak at each	mperate /900 Tl ute. r a 5-4 r 4.5.5 ll be st temp modulu	ure of hermal gram 5.1. perature us shal:
the ci Analys Measu: load. detern prior be det tempe:	4.5.5. ured c zer or rement Test 4.5.5. to lo 4.5.5. termine rature	6 <u>Glas</u> omposit equiva s will s shall 7 <u>Shor</u> per MIL ading. 8 <u>Long</u> ed per 1 prior	Σc s Trans e shall lent. b be made be con t-Beam -G-8341 itudina MIL-G-8 to load	= c ition be de The he on a ducted Shear 0, with 1 Fley 3410, ing.	density of <u>Temperatu</u> etermined eating rat 100 mil d d on lamin . The int th a minim <u>wural Test</u> with a mi	composi re. The using a e shall iameter ates tha erlamina um half- . The f nimum ha	te per 4.5. glass tran DuPont 941 be 5 + 0.5 expansion j t have been r shear str hour soak a lerural str lf-hour soa	.5.3. msition te TMA model °C per min probe unde n cured pe rength sha at each te rength and ak at each	mperate /900 Tl ute. r a 5-4 r 4.5.5 ll be st temp modulu	ure of hermal gram 5.1. perature us shal:
the ci Analys Measu: load. detern prior be det tempe:	4.5.5. ured c zer or rement Test 4.5.5. to lo 4.5.5. termine rature	6 <u>Glas</u> omposit equiva s will s shall 7 <u>Shor</u> per MIL ading. 8 <u>Long</u> ed per 1 prior	Σc s Trans e shall lent. b be made be con t-Beam -G-8341 itudina MIL-G-8 to load	= d ition be de The he on a ducted Shear 0, with 1 Fley 3410, ing.	density of <u>Temperatu</u> etermined eating rat 100 mil d d on lamin . The int th a minim <u>rural Test</u> with a mi	composi re. The using a e shall iameter ates tha erlamina um half- . The f nimum ha	te per 4.5. glass tran DuPont 941 be 5 + 0.5 expansion j t have been r shear str hour soak a lexural str lf-hour soa	.5.3. msition te TMA model °C per min probe unde n cured pe rength sha at each te rength and ak at each	mperate /900 Tl ute. r a 5-4 r 4.5.5 ll be st temp modulu	ure of hermal gram 5.1. perature us shal:
the ci Analys Measu: load. detern prior be de tempe:	4.5.5. ured c zer or rement Test 4.5.5. to lo 4.5.5. termine rature	6 <u>Glas</u> omposit equiva s will s shall 7 <u>Shor</u> per MIL ading. 8 <u>Long</u> ed per 1 prior	Σc s Trans e shall lent. be be made be con t-Beam -G-8341 itudina MIL-G-8 to load	= d ition be de The he on a ducted Shear 0, with 1 Fley 3410, ing.	density of <u>Temperatu</u> etermined eating rat 100 mil d d on lamin . The int th a minim <u>rural Test</u> with a mi	composi re. The using a e shall iameter ates tha erlamina um half- . The f nimum ha	te per 4.5. glass tran DuPont 941 be 5 + 0.5 expansion j t have been r shear str hour soak a lexural str lf-hour soa	.5.3. msition te TMA model C per min probe unde n cured pe rength sha at each te rength and ak at each	mperata /900 Tl ute. r a 5-4 r 4.5.5 ll be st temp modulu	ure of hermal gram 5.1. perature us shall
the ci Analy: Measu: load. determ prior be de tempe:	4.5.5. ured c zer or rement Test 4.5.5. to lo 4.5.5. termine rature	6 <u>Glas</u> omposit equiva s will s shall 7 <u>Shor</u> per MIL ading. 8 <u>Long</u> ed per 1 prior	Σc s Trans e shall lent. be made be con t-Beam -G-8341 itudina MIL-G-8 to load	= d ition be de The he on a ducted Shear. 0, with 1 Fley 3410, ing.	density of <u>Temperatu</u> etermined eating rat 100 mil d d on lamin . The int th a minim <u>rural Test</u> with a mi	composi re. The using a e shall iameter ates tha erlamina um half- . The f nimum ha	te per 4.5. glass tran DuPont 941 be 5 + 0.5 expansion j t have been r shear str hour soak a lexural str lf-hour soa	.5.3. msition te TMA model °C per min probe unde n cured pe rength sha at each te rength and ak at each	mperati /900 Tl ute. r a 5-4 r 4.5.5 ll be st tem modulu	ure of hermal gram 5.1. peratur us shal:
the ci Analy: Measu: load. determ prior be de tempe:	4.5.5. ured c zer or rement Test 4.5.5. to lo 4.5.5. termine rature	6 <u>Glas</u> omposit equiva s will s shall 7 <u>Shor</u> per MIL ading. 8 <u>Long</u> ed per 1 prior	Σc s Trans e shall lent. b be made be con t-Beam -G-8341 itudina MIL-G-8 to load	= d ition be de The he on a ducted Shear 0, with 1 Fley 3410, ing.	density of <u>Temperatu</u> etermined eating rat 100 mil d d on lamin . The int th a minim <u>wural Test</u> with a mi	composi re. The using a e shall iameter ates tha erlamina um half- . The f nimum ha	te per 4.5. glass tran DuPont 941 be 5 + 0.5 expansion j t have been r shear str hour soak a lexural str lf-hour soa	.5.3. msition te TMA model °C per min probe unde n cured pe rength sha at each te rength and ak at each	mperati /900 Tl ute. r a 5-4 r 4.5.5 ll be st temp modulu	ure of hermal gram 5.1. peraturn us shal:
the ci Analy: Measu: load. determ prior be de tempe:	4.5.5. ured c zer or rement Test 4.5.5. to lo 4.5.5. termine rature	6 <u>Glas</u> omposit equiva s will s shall 7 <u>Shor</u> per MIL ading. 8 <u>Long</u> ed per 1 prior	Σc s Trans e shall lent. b be made be con t-Beam -G-8341 itudina MIL-G-8 to load	= d ition be de The he on a ducted Shear 0, with 1 Fley 3410, ing.	density of <u>Temperatu</u> etermined eating rat 100 mil d d on lamin . The int th a minim <u>rural Test</u> with a mi	composi re. The using a e shall iameter ates tha erlamina um half- . The f nimum ha	te per 4.5. glass tran DuPont 941 be 5 + 0.5 expansion j t have been r shear str hour soak a lexural str lf-hour soa	.5.3. msition te TMA model °C per min probe unde n cured pe rength sha at each te rength and ak at each	mperati /900 Tl ute. r a 5-4 r 4.5.5 ll be st tem modulu	ure of hermal gram 5.1. peraturn us shal:
the ci Analy: Measu: load. determ prior be de tempe:	4.5.5. ured c zer or rement Test 4.5.5. to lo 4.5.5. termine rature	6 <u>Glas</u> omposit equiva s will 5 shall 7 <u>Shor</u> per MIL ading. 8 <u>Long</u> ed per 1 prior	Σc s Trans e shall lent. be be con t-Beam -G-8341 itudina MIL-G-8 to load	= d ition be de The he on a ducted Shear 0, with 1 Fley 3410, ing.	density of <u>Temperatu</u> etermined eating rat 100 mil d d on lamin . The int th a minim <u>rural Test</u> with a mi	composi re. The using a e shall iameter ates tha erlamina um half- . The f nimum ha	te per 4.5. glass tran DuPont 941 be 5 + 0.5 expansion j t have been r shear str hour soak a lexural str lf-hour soa	.5.3. msition te TMA model °C per min probe unde n cured pe rength sha at each te rength and ak at each	mperatu /900 T) ute. r a 5-4 r 4.5. ll be st tem modulu	ure of hermal gram 5.1. peraturn us shal:
the ci Analy: Measu: load. detern prior be de: tempe:	4.5.5. ured c zer or rement Test 4.5.5. to lo 4.5.5. termin rature	6 <u>Glas</u> omposit equiva s will 7 <u>Shor</u> per MIL ading. 8 <u>Long</u> ed per 1 prior	Σc s Trans e shall lent. b be made be con t-Beam -G-8341 itudina MIL-G-8 to load	= d ition be de The he on a ducted Shear 0, with 1 Fley 3410, ing.	density of <u>Temperatu</u> etermined eating rat 100 mil d d on lamin . The int th a minim <u>rural Test</u> with a mi	composi re. The using a e shall iameter ates tha erlamina um half- . The f nimum ha	te per 4.5. glass tran DuPont 941 be 5 + 0.5 expansion j t have been r shear str hour soak a lerural str lf-hour soa	.5.3. msition te TMA model °C per min probe unde n cured pe rength sha at each te rength and ak at each	mperati /900 Tl ute. r a 5-4 r 4.5.5 ll be st tem] modulu	ure of hermal gram 5.1. peraturn us shal:
the ci Analy: Measu: load. detern prior be de: tempe:	4.5.5. ured c zer or rement Test 4.5.5. to lo 4.5.5. termin rature	6 <u>Glas</u> omposit equiva s will 7 <u>Shor</u> per MIL ading. 8 <u>Long</u> ed per 1 prior	Σc s Trans e shall lent. be made be con t-Beam -G-8341 itudina MIL-G-8 to load	= d ition be de The he on a ducted Shear 0, with 1 Fley 3410, ing.	density of <u>Temperatu</u> etermined eating rat 100 mil d d on lamin . The int th a minim <u>rural Test</u> with a mi	composi re. The using a e shall iameter ates tha erlamina um half- . The f nimum ha	te per 4.5. glass tran DuPont 941 be 5 + 0.5 expansion j t have been r shear str hour soak a lerural str lf-hour soa	.5.3. msition te TMA model °C per min probe unde n cured pe rength sha at each te rength and ak at each	mperata /900 Tl ute. r a 5-4 r 4.5.5 ll be st tem] modulu	ure of hermal gram 5.1. perature us shal:
the ci Analy: Measu: load. detern prior be de: tempe:	4.5.5. ured c zer or rement Test 4.5.5. to lo 4.5.5. termine rature	6 <u>Glas</u> omposit equiva s will 7 <u>Shor</u> per MIL ading. 8 <u>Long</u> ed per 1 prior	Σc s Trans e shall lent. be be con t-Beam -G-8341 itudina MIL-G-8 to load	= d ition be de The he on a ducted Shear 0, with 1 Fley 3410, ing.	density of <u>Temperatu</u> etermined eating rat 100 mil d d on lamin . The int th a minim <u>rural Test</u> with a mi	composi re. The using a e shall iameter ates tha erlamina um half- . The f nimum ha	te per 4.5. glass tran DuPont 941 be 5 + 0.5 expansion j t have been r shear str hour soak a lerural str lf-hour soa	.5.3. msition te TMA model °C per min probe unde n cured pe rength sha at each te rength and ak at each	mperatu /900 T) ute. r a 5-4 r 4-5-5 ll be st tem modulu	ure of hermal gram 5.1. perature us shal:
the cr Analy: Measu: load. determ prior be de: tempe:	4.5.5. ured c zer or rement Test 4.5.5. to lo 4.5.5. termine rature	6 <u>Glas</u> omposit equiva s will 7 <u>Shor</u> per MIL ading. 8 <u>Long</u> ed per 1 prior	Σc s Trans e shall lent. be be con t-Beam -G-8341 itudina MIL-G-8 to load	= d ition be de The he on a ducted Shear 0, with 1 Flex 3410, ing.	density of <u>Temperatu</u> etermined eating rat 100 mil d d on lamin . The int th a minim <u>rural Test</u> with a mi	composi re. The using a e shall iameter ates tha erlamina um half- . The f nimum ha	te per 4.5. glass tran DuPont 941 be 5 + 0.5 expansion j t have been r shear str hour soak a lerural str lf-hour soa	.5.3. msition te TMA model °C per min probe unde n cured pe rength sha at each te rength and ak at each	mperatu /900 T) ute. r a 5-4 r 4.5. ll be st tem] modulu	ure of hermal gram 5.1. perature us shal:
the cr Analy: Measu: load. determ prior be de: tempe:	4.5.5. vered c verent Test 4.5.5. to lo 4.5.5. termine termine	6 <u>Glas</u> omposit equiva s will s shall 7 <u>Shor</u> per MIL ading. 8 <u>Long</u> ed per 1 prior	Σc s Trans e shall lent. be be con t-Beam -G-8341 itudina MIL-G-8 to load	= d ition be de The he on a ducted Shear 0, with 1 Flex 3410, ing.	density of <u>Temperatu</u> etermined eating rat 100 mil d d on lamin . The int th a minim <u>rural Test</u> with a mi	composi re. The using a e shall iameter ates tha erlamina um half- . The f nimum ha	te per 4.5. glass tran DuPont 941 be 5 + 0.5 expansion j t have been r shear str hour soak a lexural str lf-hour soa	.5.3. msition te TMA model °C per min probe unde n cured pe rength sha at each te rength and ak at each	mperatu /900 T) ute. r a 5-4 r 4.5. ll be st tem] modulu test	ure of hermal gram 5.1. perature us shal:
the cr Analy: Measu: load. deter prior be de: tempe:	4.5.5. ured c zer or rement Test 4.5.5. to lo 4.5.5. termine termine	6 <u>Glas</u> omposit equiva s will s shall 7 <u>Shor</u> per MIL ading. 8 <u>Long</u> ed per 1 prior	Σc s Trans e shall lent. be be con t-Beam -G-8341 itudina MIL-G-8 to load	= d ition be de The he on a ducted Shear 0, with 1 Flex 3410, ing.	density of <u>Temperatu</u> etermined eating rat 100 mil d d on lamin . The int th a minim <u>rural Test</u> with a mi	composi re. The using a e shall iameter ates tha erlamina um half- . The f nimum ha	te per 4.5. glass tran DuPont 941 be 5 + 0.5 expansion j t have been r shear str hour soak a lexural str lf-hour soa	.5.3. msition te TMA model °C per min probe unde n cured pe rength sha at each te rength and ak at each	mperatu /900 T) ute. r a 5-4 r 4.5. ll be st temp modulu test	ure of hermal gram 5.1. perature us shall
the cr Analy: Measu: load. deter prior be de: tempe:	4.5.5. ured c zer or rement Test 4.5.5. to lo 4.5.5. termine termine	6 <u>Glas</u> omposit equiva s will s shall 7 <u>Shor</u> per MIL ading. 8 <u>Long</u> ed per 1 prior	Σc s Trans e shall lent. be be con t-Beam -G-8341 itudina MIL-G-8 to load	= d ition be de The he on a ducted Shear 0, with 1 Flex 3410, ing.	density of <u>Temperatu</u> etermined eating rat 100 mil d d on lamin . The int th a minim <u>rural Test</u> with a mi	composi re. The using a e shall iameter ates tha erlamina um half- . The f nimum ha	te per 4.5. glass tran DuPont 941 be 5 + 0.5 expansion j t have been r shear str hour soak a lexural str lf-hour soa	.5.3. msition te TMA model °C per min probe unde n cured pe rength sha at each te rength and ak at each	mperatu /900 T) ute. r a 5-4 r 4.5. ll be st temp modulu test	ure of hermal gram 5.1. perature us shall
the cr Analy: Measu: load. deter prior be de: tempe:	4.5.5. ured c zer or rement Test 4.5.5. to lo 4.5.5. termine termine	6 <u>Glas</u> omposit equiva s will s shall 7 <u>Shor</u> per MIL ading. 8 <u>Long</u> ed per 1 prior	Σc s Trans e shall lent. be be con t-Beam -G-8341 itudina MIL-G-8 to load	= d ition be de The he on a ducted Shear 0, with 1 Flex 3410, ing.	density of <u>Temperatu</u> etermined eating rat 100 mil d d on lamin . The int th a minim <u>rural Test</u> with a mi	composi re. The using a e shall iameter ates tha erlamina um half- . The f nimum ha	te per 4.5. glass tran DuPont 941 be 5 + 0.5 expansion j t have been r shear str hour soak a lerural str lf-hour soa	.5.3. msition te TMA model °C per min probe unde n cured pe rength sha at each te rength and ak at each	mperatu /900 T) ute. r a 5-4 r 4.5. ll be st tem] modulu test	ure of hermal gram 5.1. perature us shall
the cr Analy: Measu: load. determ prior be de: tempe:	4.5.5. ured c zer or rement Test 4.5.5. to lo 4.5.5. termine rature	6 <u>Glas</u> omposit equiva s will s shall 7 <u>Shor</u> per MIL ading. 8 <u>Long</u> ed per 1 prior	Σc s Trans e shall lent. be be con t-Beam -G-8341 itudina MIL-G-8 to load	= d ition be de The he on a ducted Shear 0, with 1 Flex 3410, ing.	density of <u>Temperatu</u> etermined eating rat 100 mil d d on lamin . The int th a minim <u>rural Test</u> with a mi	composi re. The using a e shall iameter ates tha erlamina um half- . The f nimum ha	te per 4.5. glass tran DuPont 941 be 5 + 0.5 expansion j t have been r shear str hour soak a lerural str lf-hour soa	.5.3. msition te TMA model °C per min probe unde n cured pe rength sha at each te rength and ak at each	mperatu /900 T) ute. r a 5-4 r 4.5. ll be est temp modulu test	ure of hermal gram 5.1. perature us shal:

388

	NUMBER				REVISI	ON LET	TER	rr		PAGE		]
	MB0130-	152	A			I					25	
inches shall stick: evacus the fo	5.0 PREPARATI 5.1 <u>Packaging</u> 5 in diameter. be used on on ing to each ot ated, moisture 5.2 <u>Identific</u> collowing data:	ON FOR DE Prepreg A nonadh e side of her. Each -proof pla <u>ation</u> . Ea	g mate herent the m n roll astic ach ro	Y erial t pap nater l or bag. oll o	shall er or M ial to rolls c f prepr	be i lylai prev of pi reg s	olle ser ent epre hall	d on barat the g sh be	a r or o laye all perm	eel r f a c rs of be he anent	ot les contras `mater sat-ses :ly man	es than sting co rial fro aled in rked wi
Declar	Lal: Graphite	/Polyimide	e Kesi	in Pr	epreg -	. 600	) F A	ppli	cati	ons		
Morrie	ell internatio	nar Specif	LICATI	lon:		- 152	тy	pe:		0	1888 -	
manu18	No .	and Produ	1CT 10	ienti	11Catio	n: _	с¥-					
Roll					94 محتا	ue C	na ao+-	.nuia		=: N	ot Ad	unatod.
Stone	····	Nov 01	 סי			ar 1 	eet:	uro:	88:	, ^N	et Adj	usted:
DIOLAF	se iemperature	, Max	<b>r</b>		Sneii I	lie:	_9_	mont.	ns a			·
r												
follow Materi Purche	ving informati al: Graphite	f Shipment on: /Polyimide	t. Eac Resi	ch shi in Pre	ipping epreg -	cont 600	aine •F A	r sha	all 1 catio	oe ma ons	rked w	rith the
follow Materi Purcha Manufa	wing informati al: Graphite ase Order No.:	f Shipment on: /Polyimide	e Resi	in Pro	ipping epreg -	cont 600	aine •F A	r sha	all   catio	oe ma ons	rked w	vith the
follow Materi Purcha Manufa Storag STORE	A Marking o ying informati al: Graphite ase Order No.: actur's Name a ge Temperature AT 0°F	f Shipment on: /Polyimide 	t. Eac e Resi : Iden F	h shi in Pro	ipping epreg - cation Shelf L	cont 600 ife:	aine •F A 	r sha	all   catio	oe ma ons t O°F	rked w	rith the
follov Materi Purcha Manufa Storag Storag	Arking o ving informati al: Graphite ase Order No.: actur's Name a ge Temperature AT 0°F	f Shipment on: /Polyimide nd Product , Max0*	e Resi ; Iden ;	in Pro	ipping epreg - cation Shelf L	cont 600 ife:	aine •F A 	pplic	all   catio	ons t O°F	rked w	ith the
follow Materi Purcha Manufa Storag Storag STORE 6 when m aerosy and st	Arking o ving informati al: Graphite ase Order No.: actur's Name a ge Temperature AT O'F 5.0 NOTE 5.0 NOTE 5.1 Intended bolded using i bace, and simi crength-to-wei	f Shipment on: /Polyimide nd Product , Max. <u>O</u> <u>Use</u> . Mate ndicated l larly rela ght ratios	erial amina ated p	procenting requi	ipping preg - cation Shelf L ired in method ry stru ired.	cont 600 ife: acc s, i ctur	aine •F A  orda s su al c	nce titab:	all    ns a with le fo	this this whe	speci e in a re hig	fication fication fication fication fication fication

~

APPENDIX B2

•

PREPARED BY	CODE IDENT, NO.: 03953	NUMBER
R. L. Long		MAU105-328
APPROVALS	Space Division	Process
		DATE
	Downey, California 90241	1-13-82
	SPECIFICATION	SUPERSEDES SPEC. DATED:
		BEV LTB
		A PAGE 1 of 11
TITLE		Total Pages 12
т	RETCATION OF LODG 160 BOLVIMIDE CRADUTE	
1	DETERTION OF LARC 100 FOLITMIDE/GRAFAITE	COMPOSITES
		· · · · · ·
EO		
- E.U.		

.....

CODE IDENT. NO.

<u>ARAGRAPH</u> 1.0 S 2.0 H 3.0 H 4.0 G 5.0. H 6.0. M	MA0105-328          TABLE OF CONTENT         SCOPE         APPLICABLE DOCUMENT         REQUIREMENTS         QUALITY ASSURANCE         PREPARATION FOR DEL         NOTES	IS AND MA	TERIAL				i	PAGE 2 2 2 7 11 11
ARAGRAPH 1.0 \$ 2.0 # 3.0 H 4.0 0 5.0. H 6.0. M	TABLE OF CONTENT SCOPE APPLICABLE DOCUMENT REQUIREMENTS QUALITY ASSURANCE PREPARATION FOR DEL NOTES	<u>FS</u> TS AND MA	TERIAL					PAGE 2 2 7 11 11
ARAGRAPH 1.0 S 2.0 A 3.0 H 4.0 G 5.0. H 6.0. M	TABLE OF CONTENT SCOPE APPLICABLE DOCUMENT REQUIREMENTS QUALITY ASSURANCE PREPARATION FOR DEL NOTES	<u>FS</u> FS AND MAT	TERIAL					PAGE 2 2 7 11 11
ARAGRAPH 1.0 \$ 2.0 # 3.0 H 4.0 0 5.0. H 6.0. M	TABLE OF CONTENT SCOPE APPLICABLE DOCUMENT REQUIREMENTS QUALITY ASSURANCE PREPARATION FOR DEL NOTES	<u>FS</u> TS AND MAT	TERIAL					PAGE 2 2 7 11 11
ARAGRAPH 1.0 S 2.0 H 3.0 H 4.0 G 5.0. H 6.0. M	TABLE OF CONTENT SCOPE APPLICABLE DOCUMENT REQUIREMENTS QUALITY ASSURANCE PREPARATION FOR DEL NOTES	<u>FS</u> TS AND MAT	TERIAL					2 2 2 7 11 11
ARAGRAPH 1.0 \$ 2.0 # 3.0 H 4.0 0 5.0. H 6.0. M	TABLE OF CONTENT SCOPE APPLICABLE DOCUMENT REQUIREMENTS QUALITY ASSURANCE PREPARATION FOR DEL NOTES	<u>FS</u> TS AND MA LIVERY	TERIAL					2 2 2 7 11 11
1.0 S 2.0 H 3.0 H 4.0 G 5.0. H 6.0. M	SCOPE APPLICABLE DOCUMENT REQUIREMENTS QUALITY ASSURANCE PREPARATION FOR DEL NOTES	IS AND MA	TERIAL					2 2 7 11 11
1.0 S 2.0 H 3.0 H 4.0 G 5.0. J 6.0. M	SCOPEAPPLICABLE DOCUMENT REQUIREMENTS QUALITY ASSURANCE PREPARATION FOR DEL NOTES	IS AND MA	TERIAL					2 2 7 11 11
2.0 H 3.0 H 4.0 G 5.0. H 6.0. M	APPLICABLE DOCUMENT REQUIREMENTS QUALITY ASSURANCE PREPARATION FOR DEL NOTES	IS AND MA	TERIAL					2 2 7 11 11
3.0 H 4.0 ( 5.0. H 6.0. N	REQUIREMENTS QUALITY ASSURANCE PREPARATION FOR DEL NOTES	LIVERY .			· · · · · · · · · · · · · · · · · · ·			2 7 11 11
4.0 ( 5.0. ) 6.0. N	QUALITY ASSURANCE PREPARATION FOR DEL NOTES	LIVERY .			· · · · · · · · · · · · · · · · · · ·		•••••	7 11 11
5.0. I	PREPARATION FOR DEL NOTES	LIVERY .			• • • • • • • • •		•••••	11 11
6.0, N	NOTES			•••.	•••••			11
•		•						
• • •					,			
• • •					۲.			
• • •					۲			
•					۲			
•					۲			
•								
•								
						·		00055

CODE ID	ENT. N	0.								
NUMBER			R	EVISIO	ON"LET	PAGE				
MA0105-328	A								2	

1.0 SCOPE

1.1 <u>Description</u>. This specification establishes the materials to be used and the procedures to be followed for fabricating composites from LaRC 160 and PMR-15 polyimide resin systems reinforced with graphite fibers. This specification is intended for use in fabricating parts capable of structural application in a 316C (600F) environment.

2.0 APPLICABLE DOCUMENT AND MATERIALS

The latest issue of the following documents form a part of this specification to the extent specified herein. In case of conflict between these documents and this specification, this specification shall prevail.

2.1 Documents

FED-STD-406	Plastics: Methods of Testing
MA0110-306	Environmentally Controlled Areas
MT0302-001	Test Method Standards for Advanced Composite Materials
MT0501-510	Inspection, Ultrasonic
MT0302-502	Process Control for Fabrication of Reinforced Plastic Parts

2.2 <u>Materials</u>

MB0130-152, Graphite/Polyimide Resin Prepreg. - 600°F Applications

3.0 REQUIREMENTS

3.1 Safety Requirements

3.1.1 Supervision shall inform all personnel working to this specification of the hazards involved and the necessary precautions required when handling the chemicals used in the procedures of this specification.

3.1.2 Skin contact with the liquid chemicals and solutions required by this specification shall be prevented by wearing protective equipment (gloves, aprons, etc.).

3.1.3 Face shields or safety goggles shall be worn during the handling of solvents and liquid chemicals.

3.1.4 Solvents and chemical solutions required by this specification shall be handled only in areas approved by Industrial Safety. Operating department supervision, with Facilities and Industrial Engineering, shall ensure that local exhaust ventilation and/or general air circulation is adequate to prevent employee exposure to vapors or mists in concentrations above the Threshold Limit Values. (Threshold Limit Values can be obtained from Industrial Safety.)

0005S-2

CODE IDENT. NO.

NUMBER	NUMBER		R	EVISIO	DN LET	PAGE				
	MA0105-328	A							3	

3.1.5 Information or assistance on any aspects of safety associated with this specification shall be obtained from Industrial Safety.

3.2 <u>Material Storage and Handling</u>. Adhesive and preimpregnated material shall be stored in accordance with the storage-life requirements of the applicable material specifications and monitored per MT0302-001.

3.2.1 Materials that have exceeded the storage life shall be withheld from use pending retest to the acceptance requirements of the applicable specifications. Acceptable materials shall be returned to storage.

3.2.2 When not in use, the material shall be stored in a heat-sealed, moisture proof plastic bag. Allow refrigerated material to warm to at least  $16^{\circ}C$  ( $60^{\circ}F$ ), but less than  $32^{\circ}C$  ( $90^{\circ}F$ ), prior to opening container.

3.2.3 The total cumulative time that MB0130-152 prepreg material may be between zero and 24°C (75°F) during storage before the staging operation is started shall not exceed 168 hours.

3.3 Layup

3.3.1 Layup shall be conducted in an area classified per MAOllO-306 as GHA, Condition C, Level III. The air supplied to this area shall be filtered through an industrial grade (or better) air filter.

3.3.2 The part layup shall consist of the required number and orientation of plies of prepreg as shown on the Engineering Drawing. Material with different manufacturer's prepreg designations shall not be intermixed in the same part unless so directed in engineering documentation.

3.3.3 Each ply shall be continuous in the length direction; orientation of the prepreg shall not deviate more than two degrees from the specified direction.

3.3.4 Gaps between laterally adjacent tapes shall be no wider than 0.030 inch; tape edges in adjacent plies of the same orientation shall be staggered at least 1.0 inch.

3.3.5 Layups containing details to be co-cured into the part shall have the details prepared for processing in accordance with Engineering Drawing.

3.3.6 Heat may be applied locally to assist this operation using a hand held pressing iron or heat gun with temperatures up to 93C (200F).

3.3.7 Debulking under pressure without application of heat is permitted. If heat is used during debulking, any process control panel that is required by engineering drawings shall be debulked concurrently with the part using a proportionate number of plies and bleeders at each step. No more than two heat debulking cycles performed at  $93 \pm 6^{\circ}C$  ( $200 \pm 10^{\circ}F$ ) for  $20 \pm 5$  minutes, shall be applied prior to the final curing cycle.

0005S-3

	CODE IDI	ENT. NO	).						
	NUMBER		R	EVISIO	V LETT	TER		PAGE	
	MA0105-328	A		T				PAGE	
					L				L
3.4 configur	Assembly. Lay up the pration shown on the Engin	reimµ eerir	pregnat ng Draw	ed co ing.	mpos	site	materi	al to the	
3.5	Laminate Cure Procedure								
(1)	Assemble lavup for ouri	<b>n</b> a	a hours	4 m E	1.	1			
(2)	Install in autoplaye an	ug as	s shown	in P	igur	re I.			
(2)	Apply minipum upply		ach va	cuum	and	inst	rument	ation lines.	
(3)	autoclave pressure of 1 rate less than 2.5 cent any leaks.	68.5 378KF imete	ers (1 )	neter Psi) inch)	s (2 •Shu of	27 ir it of merc	iches) `f vacu cury pe	of mercury an um and verify r minute. Co	i leak rrect
(4)	Apply vacuum in the ran to be determined from pr Quality Assurance receiv	ge of repre ving	2 to 2 eg evalu	l5 in Latio	ches n te	s of sts	mercur; conduct	y. The actua ted as part of	l level
(5)	Raise the air tempe $(425+5^{\circ}F)$ and hold at t	eratu hat t	re at {	3-4°C	(5 for	to 7 60 <u>+</u> 2	°F) per ? minut	r minute to 21 es	L8 <u>+</u> 3°C
	tooling is used. Actual minimum laminate tempera	and d l con ature	well tinditions	.me m s sha )C (4	ay n 11 b 10ºF	eed e pr	to be : edeteri	increased if h nined to achie	eavy ve a
(6)	Reapply full vacuum and	main	ntain au	itocl	ave	pres	sure a	t 1378KPa (20)	) psig).
(7)	Increase temperature to	329	+ 3°C (	625	+ 5°	F) a	t a rat	ce of 3-4°C (5	5-7°F)
( )	per minute.								
(8)	Hold at temperature for	3 ho	ours.						
(9)	Reduce the temperature 1	to 65	°C (150	)°F)	or l	ower	before	e releasing va	acuum or
<b>ABO</b> Fi				APTO ES 162 (16" CA 2 PLI 7 3TL 7 2 1 1 2 1 2 1 2 1 2 1 2 1 2 1 2 1 2 1 2	N FIDER LUL PLIES L SEPJ PLIES 1/16 - 2	GLAS ATE- VATIO 0 GLA ARATO 120 CAUJ RFOR 9LIES 3TLL 	S OR BOA NS OPTIC USS CLOTH GLASS CL 120 GLASS CL 120 GLAS SEPARATO LIES 120 ( /16" CAL - PERFOR	ATCLOTH ONAL OMIT THESE TO CURE ONE LAMIN PER TOOL ONE LAMIN PER TOOL	NATE.
	FIGURE 1. Layu	TOOL	sembly	for S	Stagi	HEAN	PLY 162 G YY KAPIC Laminat	es	
									00058-4

CODE IDENT, NO.

NUMBER	REVISION LETTER							PAGE
 MA0105-328	A							5

3.6 <u>Postcure Procedure</u>. These materials normally exhibit glass transition temperature (Tg) in excess of 316°C (600F) after curing per preceding procedure. If it is found necessary to postcure laminates, the following procedure shall be utilized.

- (1) Install laminate in air circulating oven in free standing condition.
- (2) Increase temperature at a rate of 4-5°C (8-10°F) per minute to  $316 \pm 3°C$  (600 + 5°F)
- (3) Hold  $\overline{at}$  316 + 3°C (600 + 5°F) for 16 hours.
- (4) Reduce temperature to  $65^{\circ}$ C (150°F) or lower before removing from oven.

3.7 <u>Structure Requirements</u>. Cured composites shall conform to the requirements in Table I, and the dimensional requirements of the applicble Engineering Drawings, and shall meet the requirements in 3.7.1 through 3.7.6.

3.7.1 <u>Fractures</u>. A fracture is defined as a visible break in the fiber reinforcement. There shall be no defects of this nature.

3.7.2 <u>Blisters and Unbonded Areas</u>. A blister is defined as a local increase in thickness usually caused by the formation of trapped gas generating pockets between plies during cure. These gas pockets may not increase the thickness locally but may appear as unbonded areas rather than as blisters. There shall be no defects of this nature.

3.7.3 <u>Delaminations</u>. A delamination is defined as an area where plies at the trimmed edge of the laminate have become separated for any reason, i.e., machining with dull tools such as drilling, sawing, etc. There shall be no defects of this nature.

3.7.4 <u>Bridged Plies</u>. Bridged plies are defined as areas in which the plies have insufficient or no contact with the inside radius of the mold surface or with the preceding plies. There shall be no defects of this type.

3.7.5 <u>Wrinkles</u>. A wrinkle is defined as a raised fold in one or more layers of the laminate. Defects of this nature are not allowed unless identified on the Engineering Drawing.

3.7.6 <u>Foreign Objects</u>. There shall be no inclusions of materials not specifically called for in the applicable material specifications, process specifications, or Engineering Drawings.

3.8 <u>Ultrasonic Inspection</u>. An autographic recording shall be made of the ultrasonic inspection of all cured parts and shall be kept as a permanent record. This inspection shall be conducted per 4.3.1.

0005S-5

CODE IDENT. NO.

NUMBER		REVISION LETTER	PAGE	
MA0105-328	A			6

3.9 <u>Specific Gravity</u>. The specific gravity of the laminates shall be between 1.56 and 1.58 when determined per 4.3.2.

3.10 <u>Glass Transition Temperature</u>. The glass transition temperature of the cured assembly shall exceed  $316C(600^{\circ}F)$  when measured per 4.3.3.

3.11 <u>Fiber Content</u>. The fiber content shall be between 58 and 62 volume percent when determined per 4.3.5.

3.12 Void Content. The void content shall be no greater than 2 volume percent when determined per 4.3.6.

3.13 <u>Mechanical Properties</u>. The mechanical properties shall meet the requirements of Table 1.

PROPERTY	TEST   TEMPERATURE   C (F)	     test value	TEST METHOD PARAGRAPH
fu   Flexural Strength, F   X	21 (70)     316 (600)	  1515MPa (220 KSI)     757MPa (110 KSI)	4.3.7
f Flexural Modulus, E X	21 (70) 316 (600)	117GPa (17 MSI) 	
isu Short-Beam Shear, F X	21 (70)     316 (600)	96MPa (14 KSI) 	4.3.8

Table I. Mechanical Properties

## NOTES:

 All values shown are minimum requirements for the average of specimens tested. No individual value shall be less than 80 percent of the minimum average requirement. Where the number of specimens is not otherwise specified, five (5) replicates shall be fabricated and tested.

(2) Elevated temperature tests shall be preceded by a 30  $\pm$ 2 minute soak at test temperature.

00058-6

CODE IDENT. NO.

NUMBER		F	PAGE					
MA0105-328	A							7

4.0 QUALITY ASSURANCE

4.1 Process Qualification.

4.1.1 The fabrication process shall be qualified for each combination of materials used prior to fabrication of production parts. Qualification shall consist of documented evidence the process is capable of producing a part that will meet all the requirements of this specification. Any change in procedure or materials will require requalification.

4.2 Product Acceptance.

4.2.1 Product acceptance shall be based upon evidence of compliance with the requirements of this specification.

4.2.2 When required by engineering drawing, a minimum of four specimens shall be tested for each of the following properties to the requirements in Table I:

### Room Temperature Test

Flexural strength Flexural modulus Short-beam shear

After One-Half Hour at 316C (600F),

Flexural strength Flexural modulus

4.2.3 The specific gravity shall be determined from the trim of the part where trim is available or from the process control coupon, and these values shall meet the requirements of 3.9.

4.2.4 When a question arises as to an excessive variation of resin content on the part, resin content determinations on the part trim or a process control coupon shall be made at the request of Quality Control and shall meet the requirements of 3.11, 3.12, and 3.13.

<u>NOTE</u>: A process control coupon is a laminate fabricated with the same materials subjected to the same cure cycle as the part represented. The coupon shall have a size sufficient to supply all the specimens required by this specification.

4.3 Test Methods.

4.3.1 <u>Ultransonic Inspection</u>. The cured laminates shall be ultrasonically inspected per MT0501-510.

0005S-7

CODE IDENT. NO.

NUMBER	REVISION LETTER							PAGE		
MA0105-328	A									8

4.3.2 <u>Specific Gravity</u>. The specific gravity shall be determined in accordance with Federal Test Method No. 406, Method 5011.

4.3.3 <u>Glass Transition Temperature</u>. The glass transition temperature of the cured composite shall be determined using a DuPont 941 TMA mode/900 Thermal Analyzer, or equivalent. The heating rate shall be  $5 \pm 0.5^{\circ}C$  ( $9 \pm .9^{\circ}F$ ) per minute. Measurements will be made on a 100-mil-diameter expansion probe under a 5-gram load.

4.3.4 <u>Fiber Content</u>. The fiber content of the cured laminate shall be determined by either the acid digestion or hydrazine digestion method as follows:

### Acid Digestion

CAUTION: All operations involving acid digestion shall be performed in a fume hood having adequate airflow and other safeguards to prevent fumes from escaping. The operator shall wear adequate protective clothing to prevent contact between the acid or other reactants with the skin.

- 1. The composite specimens used for specific gravity determinations described in 4.3.2 shall be used.
- 2. Obtain a clean, dry extraction thimble or clean the thimble in a beaker containing nitric acid for a minimum of one hour at  $149 \pm 6^{\circ}$ C (300  $\pm 10^{\circ}$ F). Wash with distilled water, dry in oven at  $121 \pm 6^{\circ}$ C (250  $\pm 10^{\circ}$ F), desiccate and cool.
  - <u>NOTE</u>: May be purchased from Van Waters & Rogers co. as fritted glass extraction thimbles, medium E.C. 35X90 millimeters.
- 3. Weigh each extraction thimble to the nearest 0.1 milligram and record as "W1"
- 4. Dry the specimens used in the specific gravity determination and place each of three in a clean extraction thimble and weigh to the nearest 0.1 milligram. Record as " $W_2$ "
- 5. Place thimble and specimen in a beaker fitted with a raised platform and magnetic stirring bar, and add concentrated sulfuric acid until the specimen is covered. Bring slowly to a boil and hold for 30 minutes.
- 6. Remove thimble and decant spent sulfuric acid. Replace thimble in same beaker and repeat Step 5.
- 7. Continue boiling until fibers are completely separated and the resin is completely decomposed. This is determined by visual examination and may require additional sulfuric acid.
  - <u>NOTE</u>: Complete digestion is indicated when the test specimen changes its appearance from a united mass to loose, soft fibers which have a tendency to sink to the bottom of the thimble.

0005S-8

	CODE IDENT. NO.
NUMBEI	REVISION LETTER PAGE
 L	MA0105-520 A 9
8.	After digestion, place on a magnetic stirrer, stir slowly and allow to cool below 149°C (300°F).
9.	While stirring, <u>carefully</u> add 10 percent hydrogen peroxide to the hot solution.
	<u>CAUTION</u> : Allow hydrogen peroxide to run down the side of the beaker. Add very slowly and incrementally.
10.	Continue adding hydrogen peroxide until the acid solution turns a transparent, clear color. If a clear color is not obtained, the test is invalid and should be repeated.
11.	Allow acid to digest three more minutes.
12.	Remove extraction thimble from the acid, drain, place in a rubber crucible holder on a vacuum filter flask, and wash fibers with distilled water until free of acid as indicated by neutrality indication of pH paper. Denatured alcohol may be used to expedite drying after acid removed.
13.	Remove thimble containing fibers and dry in an oven maintained at 149 $\pm$ 6°C (300 $\pm$ 10°F) for a minimum of 30 minutes. Cool in a desiccator and weigh to the nearest 0.1 milligram. Record weight as "W ₃ ".
14.	Calculate the fiber content according to the following equation:
• •	Fiber content, weight percent = $\frac{W_3 - W_2}{W_1 - W_2}$ X 100
	Where $W_1$ = weight of extraction thimble plus test specimen before acid digestion in grams
	$W_2$ = weight of extraction thimble in grams
	W ₃ = weight of extraction thimble plus specimen after acid digestion in grams
Hydr	azine Digestion
	<u>CAUTION</u> : All operations involving hydrazine shall be performed in a fume hood with an airflow edequate to prevent any fumes escaping. It shall be configured to prevent dropping of any solid such as rust or dust or dripping of any condensed liquid contaminate onto the work area. The work area shall be free of all debris and dust. The operator shall wear adequate rubber gloves and other protective clothing to prevent any contact of the hydrazine to the skin.
1.	A cured laminate with a nominal size of 1 by $1/2$ inch and a nominal weight of 0.5 to 0.8 gram shall be weighed to the nearest 0.1 milligram. Record weight as "W _l ".
	00058-9

٠

		CODE IDENT. N	0.					
NUMBE	R		•	REVISIO	NLETTER		PAGE	
	MA0105-328	A					10	
2.	Place the s add approxim or hydrazin	ample in a mately 50 m e hydrate.	400-mi millili	llilii ters c	ters wi of reag	de mou ent gra	th Erlenmyer ade anhydrous	flask and hydrazine
3.	Heat until separation mixture to	the materia of fibers f go to dryne	al is c Tree of ess.	omplet any p	cely de oolyimi	graded de res:	as indicated in. Do not a	i by the llow the
· 4.	Obtain a cl the nearest	ean, dry co 0.1 millig	gram.	rade f Record	fritted d weigh	glass t as "	filter and w W2".	weigh to
5.	Filter the of fritted gla	digested ma ss filter.	iterial	throu	igh the	weight	ed, coarse-gr	rade,
6.	Rinse the f	ibers with	ethano	l unti	il the	washin	gs are color]	less.
7.	Oven dry at	90 to 1000	°C (194	to 21	L2°F).			
8.	Cool in a d 0.1 milligra	esiccator a am. Record	and wei I weigh	gh the t as "	e filte 'W ₃ .	r and a	contents to t	the nearest
9.	Calculate f	iber conter	nt as s	hown i	in Step	14 of	Acid Method:	:
4.3.5 <u>F</u> the following	iber Content, g formula:	Volume Per	cent.	Calcu	ulate t	he fib	er volume acc	cording to
Fiber vo	lume, percent	(V%) = (%V)	$\frac{\sqrt{6}}{f}$ x	e ^{p x}	100			
When	re (W _{f)f}	= fiber	weight	fract	ion pe	r 4.3.	4	
	٩f	= specif on the	fic gra e batch	vity of ma	of fibe aterial	r per used	vendor certif	fication
	ρ _c	= specif	ic gra	vity c	of comp	osite	per 4.3.2	
4.3.6 <u>V</u> following for	oid Content. mula:	Calculate	the vo	id vol	lume pe	rcent	according to	the
Percent	Void Volume =	$\frac{100}{\rho_{c}} - [\frac{(v)}{v}]$	$\frac{VT_{r}}{\rho_{r}} +$	(Wt%)	) <u>f</u> ]			
Where:	(Wt %) _r =	resin weig	ght per	cent p	ber 4.3	• 4		
	(W _f ) _f =	fiber weig	ght fra	ction	per 4.	3.4		
ſ	) _c =	specific g	gravity	of co	omposit	e per	4.3.2	
ĥ	°r =	specific g the batch	gravity of mat	of re erial	esin pe used	r·vend	or certificat	cion on
ſ	f =	specific g batch of p	gravity nateria	of fi l used	iber pe 1	r vendo	or certificat	ion on the

M131-H-28 REV. 4-73

401

00055-10

CODE IDENT. NO.

MA0105-328 A 11	NUMBER	REVISION L	ETTER	PAGE	
	MA0105-328	A		11	

4.3.7 <u>Flexure Test</u>. The flexure strength and modulus shall be determined in accordance with MT0302-001 with a minimum half-hour soak at each test temperature prior to loading.

4.3.8 <u>Short-Beam Shear</u>. The interlaminar shear strength of the laminate shall be determined in accordance with MT0302-001, with a minimum half-hour soak at each test temperature prior to loading.

4.4 <u>Records</u>. Records of the entire fabrication process and the control thereof shall be maintained for the purpose of demonstrating process control.

5.0 PREPARATION FOR DELIVERY

This section is not applicable to this specification.

6.0 NOTES

This section is not applicable to this specification.

00058-11

APPENDIX B3

PREPARED C. C.	вү Kammerer			мрр NO. 501MT51	L <b>OM</b> 03			
APPROVAL	If began	· ·	F	rev. ltr. New	PAGE 1 of 6			
E./e	Santo J.	MATERIAL		DATE ](	0-11-74			
		PROCESSING	EES	SUPERSEDE	S MPP DATED			
		PROCEDURE	E GO E	AUTHORIZI MT050	NG MPS 1-510			
QUAI	LITY PROCESS PRO	CEDURE TITLE						
	ULTRASON	IC INSPECTION OF ADHESIVE	BONDED AS	SEMBLIES	6			
	REVISION RECORD							
REV. NO. CHG LTR	D		APPROVAL/DATE					
	. •							
	EO 839733							

FORM 3916-5 REV 3-73

MPP NUMBER				REV	ISION	LET	TER			T	PAGE	)
501MT	51 <b>0M</b> 03										. Ade	2
1.	SCOPE											
1.1	This procedure descr ultrasonic through-t bonded honeycomb str and MT0501-508.	ibes th ransmis ructures	ne met sion s, to	hods insg the	s to bect req	be ion uire	used of v ement	l wher roids is of	n pe in MTO	rfo adh 501	rming ésive -510	
2.0	APPLICABLE DOCUMENTS	/MATERI	ALS									
2.1	501MT510M08	501MT510M08 Ultrasonic Pre-Inspection Preparation										
3.0	GENERAL NOTES/SAFETY	REQUI	EMENT	S								
3.1	Personnel Qualification and Certification - All personnel per- forming inspection operations to this procedure will require both "certification" and "qualification" as follows:											
3.1.1	Certification: All to Space Division Tr	inspect aining	cors w Cours	ill e Na	be . l	curi 82A:	rent] I/J.	Ly "ce	erti	fie	d"	
3.1.2	Qualification: All inspectors will, in addition to certification, be qualified in the ultrasonic inspection process (TCN 192AI) to be used.											
3.2	All test equipment shall be used with safety precautions suggested by suppliers.							ted				
3•3	Industrial Safety approved ventilation must be available and operating in all facility areas where chemicals in this procedure are used.							re				
3.4	Personnel must wear goggles during use o	imperv: of chem:	ious p icals.	last	tic	glo	ves a	and p	rote	cti	ve	
3.5	Reference Standards: Ultrasonic inspection shall be performed upon production parts only when standards with known size defects and cross sections are available which represent the part configuration							upon d tion.				
3.6	Equipment Maintenance in operating conditi Inspection supervisi	e: Ult on, and on.	trason 1 shal	ic e l be	equi e th	pmen e ro	nt sl espoi	nall I nsibi:	be m lity	ain of	taine	đ
3.7	Reference Standard M shall be maintained Repairs will be made material such as pol	Laintens with ec , when ysulfic	ance: lge se requi le sea	Ad) als red lant	nesi fre , wi ts.	ve 1 e o: th j	bonde f hol perma	ed re: les an anent	fere nd f sea	nce Tac	stan tures g	dards •
3.8	Electronic Instrumen be allowed to warm u (not less than 15 min instrument.	tation p prior utes) r	All to i recomm	ele nspe ende	ectr ecti ed b	oni on : y t)	c in: for a he ma	strume a per: anufac	enta iod ctur	tic of er	on sha time of th	11 e

FORM 3916-S-1 REV. 5-73

•

			REV	ISION LE	TTER			PAGE	
	2							2	
	)	L	JJ		Li		<u> </u>	<u>_</u>	
3.9	Equipment Calibrati brated in accordanc Engineering functio inspection process.	on: Ultra e with pronal group	asonic ocedure respor	instru s esta sible	men bli: for	tation shed b the u	shall y the ltraso	be cali- Quality nic	
3.10	The following defin the procedure:	itions are	e inclu	nded to	9 <b>8</b> 5	sist i	n unde	rstanding	
	Squirter: A clear ducer, a nozzle, a	plastic as water inp	ssembly ut fitt	housi ing, s	ing a ind a	an ult a nozz	rasoni le ins	c trans- ert.	
	<u>Nozzle:</u> An adjusta optimum laminar flo	ble plast w of wate	ic tube r throu	e which ugh the	n ca e no	n be a zzle i	djuste nsert.	d for	
	<u>Nozzle Insert</u> : An orifice control plastic insert in the nozzle end to provide water stream diameters of $1/8$ , $5/32$ , $3/16$ , $7/32$ , 1/4, $5/16$ and $3/8$ inch.								
	Attenuation: The l The amplitude measu (db).	oss of pr are of att	opagati enuatio	ion of on will	ult L be	rasoun expre	d in a ssed i	material. n decibels	
	<u>Water Noise</u> : Dissolved air and turbulent water flow will create ultrasonic reflective noise responses (water noise) on the ultra sonic transceiver video display cathode ray tube (CRT).								
	<u>Video:</u> Ultrasonic signal responses displayed visually on the f of the transceiver (CRT).							on the face	
	Gated Video: A por C-scan recording to	tion of t show the	he ove: presei	nall v: nce or	ideo <b>a</b> bs	that ence o	is sel of defe	ected for	
	Reference Standards and fabrication of in controlled size	: A test the hardw defects.	panel are ite	that d em to l	lupl be i	icates nspect	confi ed wit	guration h built <del>-</del>	
4.0	ACCEPT/REJECT CRITE	ERIA							
4.1	The bonded honeycon determined by the p drawing and/or as p	nb structu requiremen referenced	res ac ts of by the	cept/re the app e plan	ejec plic ning	t crit able e docum	eria s engines mentati	shall be ering ion.	
5.0	PROCEDURE								
5.1	Summary - The follo	owing step	s shal	l be p	erfo	ormed i	in sequ	lence.	
	Step 1. Enter 1	required p	arts d	ata on	ins	pectio	on form	16 <b>.</b>	
	Step 2. Prepare	e the part	for u	ltraso	nic	inspec	ction.		
	Step 3. Position fixture	on referen es.	ice sta	ndard	and	part i	into ho	olding	
1									

FORM 3916-S-1 REV. 5-73

405

MPP NUMBER			REVISION LETTER
501MT510M	)3		4
	Step 4	•	Set up for reference standard scanning.
	Step 5	•	Set up for production part scanning.
	Step 6		Identify indications.
	Step 7	•	Evaluate defects.
	Step 8	•	Complete inspection records.
	Step 9	•	Clean and dry production part.
5.2	Detail	ed Ste	eps
	Step 1	•	Enter required parts data on inspection forms.
	l	A	Check the Manufacturing Order to see that all previous operations have been completed.
	1	В	Review other documentation, such as drawings, photo- graphs, etc., to verify location of closeouts and metal inserts.
	Step 2		Prepare the part for ultrasonic inspection.
	2	2A	Assign a control number to the technique control record.
	2	2B	Mask and/or seal parts as necessary to prevent water entering areas that may be necessary to keep dry. Use method presented in 501MT510M08.
	2	2C	Attach any tooling that might be necessary to support the part or standard in the inspection system.
	Step 3	3	Position reference standard and part into holding
	3	<b>3</b> A	Select the holding fixture for the configuration to be inspected. For example, flat parts can be clamped vertical between squirters.
	Step 4	ŀ	Set up for reference standard scanning.
	4	łA	Turn instrumentation to ON position, and allow to warm up for a minimum of 15 minutes.
	٦ţ	₽B	Position squirter to scan position and adjust water controls for abundant flow to force air bubbles out of the system.

.

FORM 3916-S-1 REV. 5-73

	REVISION LETTER PAGE
501MT510M03	5
40	Reduce water flow until a smooth stream is obtained, free of spirals.
4D	Observe CRT of ultrasonic detector for reflective amplitude response from surface reflection, back surface and/or reflector surfaces.
4E	Move transducers to a reference standard void and observe signal response. Little or no response will be indicated on the CRT.
4F	If amplitude of signal response over the void is great- er than 10% of total signal, increase reject until 10% or less is obtained.
4G	Return transducers to a well bonded area and observe signal response. If signal response does not return to saturation, increase gain until saturation is achieved, then repeat steps 4E, 4F, and 4G.
4H	If the saturation and 10% of saturation signal differ- ential is not obtainable, different transducer frequency and squirter nozzle insert sizes shall be used until separation is obtained.
41	Adjust gate controls until gated video includes only the through-transmission signal response.
4.1	Adjust bridge controls to set scan limits for C-scan of the reference standard.
4K	Scan approximately one inch of the reference standard with the void included.
4L	Measure the width dimensions of the C-scan recorded void. If it is not the dimension of the standard defect, increase or decrease the receiver gain and repeat the one-inch scan.
Step 5.	Set up for production part scanning.
54	Move bridge controls for a C-scan of the part. If possible, part scanning shall include the standard simultaneously with the part. If not possible, a one-inch scan of the standard shall be completed at the end of the part C-scan.
5в	With scan speed reduced to zero, push scan switch to ON and advance speed control until operation speed is obtained (not to exceed 20 inch/sec.).

FORM 3916-S-1 REV. 5-73

MPP NUMBER				F	EVISIO	N LETT	TER			PAGE
501MT510M03										6
	Step 6.	Identif	y indica	tions.	•					
	<b>6</b> A	All C-s by comp closeou	can recon arison to ts or ot	rded d o drav her an	liscon vings reas t	iti <b>n</b> u illu hat	iti stri wou	es sl ating ld ca	nall b g loca ause s	e identified tion of ignal loss.
	6в	Mark an changes	eas on the in cross	he rec s sect	cordin tion t	ig thi hat	at a are	are ( not	neces	by internal sarily voids.
	Step 7.	Evaluat	e defect	<u>s</u> .						
	7A	Evaluat with th ing dra	ion of di me applica wing requ	iscont able p uireme	tinuit proces ents f	ies s sp or a	wil eci cce	l be ficat ptand	in ac tions ce cri	cordance or Engineer- teria.
	7B	Discont evaluat	inuities ed with a	that manua	may b l A-sc	e pr an t	ese: ech	nt sl nique	hall b es.	e further
	7C	Discont verifie on the	inuities d by A-s C-scan re	large can te ecord:	er tha echniq ing.	n al lues	low sha	able 11 be	that e labe	have been led defects
	7D	Defects uated t	in sand to provide	wich a e info	struct ormati	ion r	sh ela	all   tive	be fur to lo	ther eval- ocation.
	7E	Specifi the cro sonic t	ic techni bss-secti lest syst	ques : on sha em, ai	for lo all be nd/or	cati wit coin	on h p ta	of d ulse ppin	efects echo g.	within contact,
	7F	Mark lo	cation o	f defe	ect u <u>r</u>	pon p	art	sur	face.	
	Step 8.	Complet	e inspec	tion	record	<u>ls</u> .				
	8a	Accept records	parts fr and wor	ee fro k ordo	om def er.	fects	s an	d st	amp in	nspection
	δB	Squawk with wo	defectiv ork order	e deb proc	onds d edures	or de	lam	inat	ions :	in accordance
	8C	Complet informa frequen	te ultras ation rel ncies, wa	onic ative ter n	techni to tr ozzle	ique ransd inse	rec luce ert	ord r an size	with d d rece , etc.	the required eiver
	Step 9.	Clean a	assemblie	s wit	h clea	an wa	ter	and	wipe	dry.
	9A	All mas been us 501MT5	sking and sed shall LOMO8.	/or so be re	ealing	g mat 1 in	eri acc	al t orda	hat ma nce w:	ay have ith
	9 <b>B</b>	Oven di accords	rying whe ance with	n nec 501M	essary T51 <b>0M</b> (	y or 08.	reg	uire	d, sha	all be in

FORM 3916-S-1 REV. 5-73

FAOL			1000				
		·		SUMENI NUMBEI	R	REV	SE
- 1771 C	ODE IDENT.	NO. 03953	5	01MT510M03		A	01
			DOG	UMENT TYPE	l_	ł	
1	Space Di	ivision					
TA F	Rockwell Inte	ernational		UALITY			
4/23/5	1						
		JENT.		6.0.40039		1	
	AMENDA	viciti			PAGE	Ť	
Adhe	sive Bonded	l Assemblie		23/75	0F		
			ERS				
How and			60290	yange			
part of Qua	lity Proces	ssing Proce	dure 501	LMT510M03, N	lew, dat	ted	
•							
ion referen	ce standard	l and part	into hol	lding fixtur	<u>e</u> .		
t the holdi	na fixture	for the co	onfigura	tion to be i	nspect	ed.	
xample, fla	t parts car	n be clampe	ed vertic	cal between	squirt	ers	•
C	AUTION						
2							
i improvised	l clamping o	devices are	e used in	n lieu of ap	proved		
res, restra	unts must i	pe-brovrae	i to pro	UTDIC INAGVE	rtent		
ge.							
ion referen	nce standard	d and part	into ho	lding fixtu	<u>e</u> .		
ige.	nce standard	d and part	<u>into ho</u>	lding fixtur	<u>e</u> . Inspect	ed.	
ige. ion referen t the holdi xample, fla	nce standard Ing fixture at parts can	d and part for the co n be clamp	into ho onfigura ed verti	lding fixtur tion to be f cal between	<u>e</u> . Inspect squirt	ed, ers	•
ige. ion referen t the holdi xample, fla	nce standard ing fixture at parts can	d and part for the co n be clampo	<u>into ho</u> onfigura ed verti	lding fixtur tion to be f cal between	inspect squirt	ed, ers	•
ion referen t the holdi xample, fla	nce standard Ing fixture At parts can	d and part for the co n be clampo	into ho onfigura ed verti	lding fixtur tion to be d cal between	<u>se</u> . Inspect squirt	ed, ers	•
ige. ion referen t the holdi xample, fla	nce standard Ing fixture At parts can	d and part for the co n be clampo	<u>into ho</u> onfigura ed verti	lding fixtur tion to be f cal between	inspect squirt	ed, ers	•
ige. ion referen t the holdi xample, fla	nce standard Ing fixture at parts can	d and part for the co n be clampo	<u>into ho</u> onfigura ed verti	lding fixtur tion to be f cal between	inspect squirt	ed, ers	•
ion referen t the holdi xample, fla	nce standard Ing fixture at parts can	d and part for the co n be clampo	<u>into ho</u> onfigura ed verti	lding fixtur tion to be f cal between	inspect squirt	ed, ers	•
ige. ion referen t the holdi xample, fla	nce standard ing fixture at parts can	d and part for the co n be clampo	into ho onfigura ed verti	lding fixtur tion to be f cal between	inspect squirt	ed, ers	•
ion referen t the holdi xample, fla	nce standard Ing fixture at parts can	d and part for the co n be clampo	<u>into ho</u> onfigura ed verti	lding fixtur tion to be f cal between	inspect squirt	ed, ers	•
ige. ion referen t the holdi xample, fla	nce standard Ing fixture At parts can	d and part for the co n be clampo	into ho onfigura ed verti	lding fixtur tion to be f cal between	inspect squirt	ed, ers	•
ige. ion referen t the holdi xample, fla	nce standard Ing fixture at parts can	d and part for the co n be clampo	into ho onfigura ed verti	lding fixtur tion to be f cal between	inspect squirt	ed, ers	-
nge. aion referen at the holdi ample, fla	nce standard ing fixture at parts can	d and part for the co n be clampo	<u>into ho</u> onfigura ed verti	lding fixtur tion to be f cal between	inspect squirt	ed, ers	•
i <u>on referen</u> t the holdi xample, fla	nce standard Ing fixture at parts can	d and part for the con h be clampo	<u>into ho</u> onfigura ed verti	lding fixtur tion to be f cal between	inspect	ed, ers	-
nge. aion referen at the holdi ample, fla	nce standard ing fixture at parts can	<u>d and part</u> for the co n be clampo	<u>into ho</u> onfigura ed verti	lding fixtur tion to be f cal between	inspect squirt	ed, ers	•
ion referen t the holdi example, fla	nce standard ing fixture at parts can	<u>d and part</u> for the con n be clampo	<u>into ho</u> onfigura ed verti	lding fixtur tion to be f cal between OFFICIA	inspect squirt	ed, ers	•
ion referen t the holdi example, fla	nce standard	<u>d and part</u> for the co n be clampo	into ho onfigura ed verti	lding fixtur tion to be f cal between OFFICIA RELEAS	inspect squirt	ed, ers	-
ion referen t the holdi example, fla	nce standard	d and part for the con h be clampo	<u>into ho</u> onfigura ed verti	lding fixtur tion to be f cal between OFFICIA RELEA APR 29	inspect squirt	ed, ers	•
ion referen t the holdi example, fla	nce standard	<u>d and part</u> for the co n be clampo	into ho onfigura ed verti	Iding fixtur tion to be f cal between OFFICIA RELEA APR 29 44.10	LLLY SED 1975 2 Pm	ed, ers	-
ion referen	nce standard	d and part for the con h be clampo	<u>into ho</u> onfigura ed verti	Iding fixtur tion to be f cal between OFFICIA RELEA APR 29 4/12 ENGINE	LLLY SED 1975 ERING	ed, ers	-
ion referen t the holdi example, fla	nce standard	d and part for the con be clampo	<u>into ho</u> onfigura ed verti	Iding fixtur tion to be f cal between OFFICIA RELEAS APR 29 4,10 ENGINE	LLY SED 1975 ERING	ed, ers	-
	Jultr Adhe part of Qua ion referen t the holdi xample, fla	Space D Rockwell Inter AMENDA Ultrasonic Insp Adhesive Bonded part of Quality Process ion reference standard t the holding fixture xample, flat parts can <u>CAUTION</u> improvised clamping of res. restraints must h	Space Division Rockwell International AMENDMENT Ultrasonic Inspection of Adhesive Bonded Assemblie part of Quality Processing Proce ion reference standard and part t the holding fixture for the co xample, flat parts can be clampe <u>CAUTION</u> improvised clamping devices are res. restraints must be provided	Space Division Rockwell International AMENDMENT Ultrasonic Inspection of Adhesive Bonded Assemblies Boart of Quality Processing Procedure 501 ion reference standard and part into hold t the holding fixture for the configuration caution improvised clamping devices are used in res. restraints must be provided to provid	Space Division Rockwell International AMENDMENT Ultrasonic Inspection of Adhesive Bonded Assemblies Part of Quality Processing Procedure 501MT510M03, N ion reference standard and part into holding fixture t the holding fixture for the configuration to be in example, flat parts can be clamped vertical between <u>CAUTION</u> improvised clamping devices are used in lieu of appress. restraints must be provided to prohibit inadve	Space Division Rockwell International AMENDMENT Ultrasonic Inspection of Adhesive Bonded Assemblies part of Quality Processing Procedure 501MT510M03, New, da ion reference standard and part into holding fixture. t the holding fixture for the configuration to be inspect xample, flat parts can be clamped vertical between squirt <u>CAUTION</u> improvised clamping devices are used in lieu of approved res. restraints must be provided to prohibit inadvertent	Space Division Rockwell International AMENDMENT Ultrasonic Inspection of Adhesive Bonded Assemblies part of Quality Processing Procedure 501MT510M03, New, dated ion reference standard and part into holding fixture. t the holding fixture for the configuration to be inspected, xample, flat parts can be clamped vertical between squirters. <u>CAUTION</u> improvised clamping devices are used in lieu of approved res. restraints must be provided to prohibit inadvertent

•		3568 s	HUTTLE		·
PREPARED BY	044-220 AD50	CODE IDENT, NO. 03953	DOCUMENT NUMBER	REV	SEC
L. Owen	2-5146		501MT510M03	A	02
F. Sandy	11.02. hut	Space Division	DOCUMENT TYPE		
1000	, really	Rockwell International	QUALITY ANTHORITY		
The second	+				
CALL ENG		AMENDMENT	GO 40039 DATE		
MFG	4	l'Itrasonic Inspection of	PAGE	<u>ر</u> ر	L
- Truck		Adhesive Bonded Assembling	SUBJECT		
7 26 77 PROJ		382 E	CHANGE		
]	This amendment f 501MT510M03, Rev 3.1.1 C 3.1.2 Q (	Forms a part of Quality Process vision N5W, dated October 11, 19 Certification: Space Divis Course No. 182 AI/AJ. Pualification: All inspectors v pualified to Space Division On-1 OJQ) Course No. 888 AO/AP.	Procedure 974. sion Training sill be currently The-Job Qualification		
<u>_</u> <u>w</u>	MAS				
	3.1.1 · C C	ertification: Space Divis ourse No. 182 AI/J.	ion Training		
-	3.1.2 Q c i	ualification: All inspectors w ertification, be qualified in t nspection process (TCN 192 AI)	ill, in addition to he ultrasonic to be used.		
•					
	·.		OFFICIALLY RELEASED		
		<b>.</b>	AUG - 5 1977		
			8/32 Km ENGINEERING		
FORM 3943-E REV 2	2-74		RELEASED AUG - 5 1977 8:32 AM ENGINEERING		_

		\$12	SHU	ITTLE
PREPARLO BY C. G. K. m. MILA	D/344-230	CODE IDENT. NO. 03953	DOCUMENT NUMBER	HEV SEQ
	6afety	Rockwell International Space Systems Group	50 IMT5 10M03 DOCUMENT TYPE Quality	
ME	5/80	E85 1	GO 40039	
Mfg.		AMENDMENT Wight	May 5, 1980	OF
Proj.Eng. 6/9/80		OF ADHESIVE BONDED ASSEMBLIES	BYBJECT B 5 http://www.change	
	This Amendment 501MT510M03, Re	forms a part of Quality Process Pr vision New, dated October 11, 1974	rocedure •	
Page 6	<u>98</u>			
<u>15</u>	All parts subje be dried as soo after ultrasoni	cted to water squirter inspection n as practical (not to exceed 16 1 c inspection	shall hours)	
	<u>9C</u>	. •	, · ·	
<u>15</u>	Oven drying whe accordance with	en necessary or required, shall be n 501MT510M08.	in	
Page 6	<u>9B</u>			
WAS	Oven drying whe accordance with	en necessary or required, shall be n 501MT510M08.	in	
	• •			u 0861
· · .		· · · · · · · · · · · · · · · · · · ·		N 18 1
			Э	
FORM 3945-E REV 1-79				

	- · · · · · · · · · · · · · · · · · · ·	, <i>e -</i>	SHUTTL	
PREPARED BY		63 5 10 5 10 5 5 5 5 5 5 5 5 5 5 5 5 5 5 5	DOCUMENT NUMBER	REV SEO
J. Mamon D/344.	-230 2-3108	CODE IDENT. NO. 03953	501MT510M03	A 04
APPR	OVALS /1/	O Di ista	DOCUMENT TYPE	
Tol	NES Es	Space Division	Quality	•
A Stand	Fig. Eng.	Rockwell International	AUTHORITY	
y Sullim	-	OUALITY PROCEDURE		
746P	·		GO 40111	· · · · · · · · · · · · · · · · · · ·
mila		AMENDMENT		PAGE
Proj. Eng.		ULTRASONIC INSPECTION	8-10-81	OF
& gRaluts		0F 68	SUBJECT	· .
Safety		ADHESIVE BONDED ASSEMBLIES	CHANGE	
Page 6 9B IS	This amendment for revision new, dat All parts subject	orms a part of Quality Process Proted October 11, 1974. ted to water squirter inspection s	cedure 501MT510M0 shall be dried as	soon as
· · · .	practical (not to	o exceed 72 hours) after ultrason	ic inspection.	
		•	•	•
WAS	All parts subject	ted to water squirter inspection s	shall be dried as :	soon as
	practical (not th	b exceed to hours, after diffason.	ie inspection.	
		• • • • • • • • • • • • • • • • • • •		
			5	·
			81 SEP - J PH 1= 4	OFFICIALLY
1		· · · ·	10	. •
		·	<u></u>	
FORM 3945-E REV	2-74			

ł

# APPENDIX C

This appendix contains the stress/strain curves for testing of beam and coupon specimens in tension and compression. TENSILE COUPON CURVES

STRESS MN/m²



416

Figure C-1. LARC-160/Celion (0)₅ Stress/Strain Curves at RT






STRESS MN/m²







STRESS MN/m²



Figure C-7. LARC-160/Celion (0)5 Stress/Strain Curves at 316 C (600 F)

















Figure C-14. LARC-160/Celion (<u>+</u>45)_S Stress/Strain Curves at RT.



Figure C-15. LARC-160/Celion (<u>+</u>45)_S Stress/Strain Curves at RT.







Figure C-18. LARC-160/Celion (<u>+</u>45)S Stress/Strain Curves at 204 C (400 F)



Figure C-19. LARC-160/Celion (+45)S Stress/Strain Curves at 204 C (400 F)











**ZIBESS (KSI)** 



,

STRESS MN/m² 620 413 207 0 Figure C-26. LARC-160/Celion (0,  $\pm 45$ , 90)_S Stress/Strain Curves at 204 C (400 F) 0.80 i SPECIMEN EX 106-3-2 ORIENTATION (0, ±45,90) S (79.0) (8.04) 0.31 544 55.40 + 04.0 STRAIN ULT STRENGTH MN/m² (KSI) ELASTIC MODULUS GN/m² (MSI) POISSON'S RATIO AXIAL 0 TRANSVERSE STRAIN 2 -0.40 90.06 0 60.0 30.0

**STRESS (KSI)** 









TENSION BEAM CURVES

;

STRESS  $MN/m^2$ 



STRESS MN/m²





**STRESS (KSI)** 

STRESS MN/m²





Figure C-33. Tensile Stress/Strain Characteristics of (0) $_5$  Parallel LARC-160/Celion Laminates at 316 C (600 F) - Beam Test

STRESS  $MN/m^2$ 



Figure C-34. Tensile Stress/Strain Characteristics of (0)5, Parallel LARC-160/Celion Laminates Aged for 125 Hours at 316 C (600 F), Beam Test at - 132 C (-270 F)

STRESS MN/m² 2067 **T** 2412 1723 1378 1034 689 345 1.6 EX199T-4 1.4 1.2 EX199T-6 0.-- EX199T-5 STRAIN µ (%) 0.80 0.60 .... 0.40 0.20 0 350 E 300 250 200 150 001 20 0

Figure C-35. Tensile Stress/Strain Characteristics of (0)5 Parallel Oriented LARC-160/Celion Laminates Aged 125 Hours at 316 C (600 F), Beam Test at RT

**ZIKESS (KSI)** 







Figure C-37. Tensile Stress/Strain Characteristics of (0)₅ Parallel Oriented LARC-160/Celion Laminates Aged for 125 Hours at 316 C (600 F), Beam Test at 316 C (600 F)

453

14.
# COMPRESSION BEAM CURVES

STRESS MN/m2



Figure C-38. Compression Stress/Strain Characteristics of (0)5 Parallel LARC-160/Celion Laminates at--132 C(-270 F)--Beam Test

STRESS(KSI)



STRESS MN/m²



Figure C-40. Compressive Stress/Strain Characteristics of (0)5 Parallel LARC-160/Celion Laminates at 204 C(400 F) Beam Test





STRAIN MN/m2



Figure C-42. Compression Stress/Strain Characteristics of (0)5, Parallel LARC-160/Celion Laminates aged 125 Hours at 316 C (600 F), Beam Test at - 132 C (-270 F)







STRESS MN/m²

**ZIBESS (KSI)** 

STRESS  $MN/m^2$ 



ZIBESS (KSI)

STRESS MN/m²



STRESS  $MN/m^2$ 



**ZIBESS (KSI)** 

465

Figure C-47. Compression Stress/Strain Characteristics of (0, +45,90) S LARC-160/Celion Laminates at-132(-270 F)-Beam Test





STRESS MN/m²



**ZIBESS (KSI)** 

467

Figure C-49. Compression Stress/Strain Characteristics of (0,<u>1</u>45,90)_S LARC-160/Celion Laminates at 316 C (600 F)—Beam Test





Figure C-50. Compression Stress Strain Characteristics of (0, 445, 90)S LARC-160/Celion Laminated Aged for 125 Hours at 316 C (600 F), Beam Test at -132 C (-270 F)



STRAIN µ(%)

Figure C-51. Compression Stress/Strain Characteristics of (0, <u>+</u>45, 90)S LARC-160/Celion Laminates Aged for 125 Hours at 316 C (600 F) Beam Test at RT

**ZIBESS (KSI)** 



Figure C-52. Compression Stress/Strain Characteristics of (0, <u>+</u>45, 90)S LARC-160/Celion Laminates Aged for 125 Hours at 316 C (600 F), Beam Test at 316 C (600 F)



**ZIBESS (KSI)** 

# APPENDIX D

This appendix presents design requirements for hat-section and "I" stiffened panels, design assumptions for optimization of panel configuration, analysis, and supportive calculations.

#### PANEL DESIGN REQUIREMENTS

Test panels were designed in accordance with requirements specified by NASA/LaRC.

- (1) Panel Design Load: .53 MN/m (3000 1b/in.) nominal, .49 to .56 MN/m (2800 1b to 3200 1b/in.)
- (2) Panel Skin: 8-ply isotropic layup  $(0, \pm 45, 90)_s$
- (3) Stringers: 3 minimum
- (4) Panel Dimensions and Fabrication Concepts:
  - (a) 22.86 cm (9 in.) wide X 30.48 cm (12 in.) long panels
    (16) Hat stiffened 2 each, bonded, bolted, cocured
    (16) "I" stiffened 2 each, bonded, bolted, cocured
  - (b) 22.86 cm (9 in.) wide X 121.9 cm (48 in.) long of best configurations fabricated and tested above.
- (5) Panel failure mode: 121.9 cm (48 in.) long panels Euler buckling
   30.48 cm (12 in. long) panels local crippling or buckling
- (6) Minimum Weight Design
- (7) Test at -121C (-250F), room temperature, 204C (400F), and 316C (600F) after postcure and 125 hours exposure at 316C (600F)

#### DESIGN_ASSUMPTIONS

In addition to the above design requirements, the following assumptions were made:

- Panel components are to be made of symmetric layups in order to avoid material warpage, which could severly degrade panel strength.
- (2) Tape with the following cured and postcure properties will be used;

$$E_{x} = 148.1 \text{ GN/m}^{2} (21.5 \text{ Msi}) \text{ compression}$$

$$E_{y} = 9.85 \text{ GN/m}^{2} (1.43) \text{ Msi} \text{ compression}$$

$$G_{xy} = 5792 \text{ MN/m}^{2} (840 \text{ Ksi})$$

$$V_{xy} = .25$$

$$V_{yx} = .05$$
Specific gravity, 1.57 g/cc  
Fiber volume, 60%

- (3) Long panel tests will be made with simply supported ends and with non-local load transfer into the panel to as great a degree as possible.
- (4) <u>+</u> 45 plies will be used in the web and the top of the hat or "I" section will be reinforced with 0^o plies. Past investigations (Agarwal and Davis, 1974, "Minimum-Weight Designs for Hat Stiffened Composite Panels Under Uniaxial Compression," NASA TND-7779) in-dicate that this results in the most efficient stringer-stiffener panel design. This assumption was verified during the design process.

### ANALYSIS

Panel analysis was made utilizing the AC-3 computer program and other routines written for the HP9825 for the determination of section properties following the guidelines of the Rockwell Structures Manual.

The design and optimization methodology was as follows:

- (1) Select number of stiffeners.
- (2) Select tape thickness.
- (3) Select web layup This is influenced by the assumption that symmetric laminates are to be used in panel subelements. Thus, the web of the "I" sections, which is formed from two "C" channels, must be increased or decreased by 4 plies at a time. Similarly, the legs of the hat section must each be increased or decreased by 2 plies at a time.
- (4) Select hat or "I" section width This choice may be modified at a later time, especially in the case of the "I" section, in order to prevent local crippling.
- (5) Select hat height.
- (6) Sufficient  $0^{\circ}$  ply reinforcement is added to the flange of the "I" section or top of the hat section to give the required Polar moment of inertial, I, for Euler buckling of the 122 cm (48 in.) long panels. The total load on the column will be 120.096 N (27,000 lb.) nominal 112,090 N (25,200 lb.) -128,102 N (28,800 lb.), thus, the required I is 2.7 X  $10^{-7}$ m⁴ (.6635 in⁴) nominal (2.5 X  $10^{-7}$ m⁴) -2.9 X  $10^{-7}$ m⁴ (.7077 in⁴).

This is based on an effective modulus of elasticity  $E_{eff} = 65.5 \text{ GN/m}^2$  (9.5 Msi), which is equal to the experimentally determined compression modulus of the 8-ply "isotropic" laminate.

In order to calculate I, the areas of the various panel subelements are first converted to effective areas:

$$A_{eff_i} = A_i \frac{E_i}{E_{eff}}$$

Section properties are then calculated in accordance with procedures described in the Rockwell Structures Manual.

- (7) Parametric plots are generated displaying weight vs. hat/ "I" section height for various tape thicknesses and configurations. The least weight configurations are then analyzed for local crippling.
  - (a) The b/t for the panel subelements is generally large enough such that they may be molded as simply supported long plates in axial compression.
  - (b) The formula for the local crippling of simply supported laminated plates is given in the Advanced Composite Design Guide:

^Nx, cr = 
$$\frac{2\pi_2}{b^2}$$
 (  $\sqrt{D_{11}D_{22}} + D_{12} + D_{66}$  ), 1bs/in

or

$$\sigma_{\rm cr} = \frac{2\pi^2}{b^2 t} \qquad (\qquad \sqrt{D_{11} D_{22}} + D_{12} + 2 D_{66}), \ 1bs/in^2$$

where D₁₁, D₁₂, D₂₂, and D₆₆ are the flexural rigidities of the laminate. These are calculated using the AC-3 Point Stress Analysis computer program, which is also described in the Advanced Composites Design Guide.

(c) The flange of the "I" section is molded as a plate with 3 sides simply supported and one side free. The corresponding cripping stress is given by:

$$\sigma = \frac{t^2}{b^2} Gxy$$

(8) The local crippling analysis places cut-off points on the parametric curves of weight vs hat/"I" height.

Least weight practicable designs are chosen from the curves and one is chosen subject to manufacturing constraints. The most notable fabrication problem is found in the mechanically fastened configurations in which it is necessary to provide a wide lap for tool clearance and sufficient fastener edge distance.

## SUPPORTIVE CALCULATIONS FOR PANEL DESIGNS

The following provides supportive correlation with previous research and calculations pertinent to panel section properties.

(a) Local Buckling

Shown below are crippling stress levels for subelements of the hat and "I" sections. Also shown are the predicted stress levels in the subelements at the Euler limit load of the long column and column loads necessary to precipitate local crippling.



Euler Buckling Load, L = 1.22 m (48 in.) P = 112,588 N (25,312 1b.)

Predicted:

	Crippling Stress	Stress at Euler Limit	Column Load for Local Failure
Α.	163.8 MN/m ²	154.1 MN/m ²	120,763 N
	(23,760 psi)	(22,350 psi)	(27,150 1Ъ.)
в.	731.4 MN/m ²	$345.7 \text{ MN/m}^2$	238,235 N
	(106,077 psi)	(50,131 psi)	(53,560 1Ъ.)
с.	53.2 MN/m ²	48.2 MN/m ²	124,322 N
	(7,720 psi)	(6,990 psi)	(27,950 1b.)
D.	163.8 MN/m ²	154.1 MN/m ²	120,763 N
	(23,760 psi)	(22,350 psi)	(27,150 1b.)



Euler Buckling Load L = 1.22 m (48 in.) P = 123,975 N (27,872 1b.)

Predicted:

Crippling Stress	<u>Stress at Euler Limit</u>	Column Load for Local Failure
175.2 MN/m ²	$173.3 \text{ MN/m}^2$	125,376 N
(25,412 psi)	(25,128 psi)	(28,187 1b.)
433.6 MN/m ²	392.1 MN/m ²	137,078 N
(62,880 psi)	(56,869 psi)	(30,818 1b.)
212.8 MN/m ² (30,863 psi)	54.7 $MN/m^2$ (7,938 psi)	480,012 N (108,366 1b.)

## APPENDIX E

This appendix contains the Engineering Design drawings which define the TDS. The drawing list is as follows:

SS79-00249 2 Sheets	Technology Demonstration Segment - Graphite/Polyimide, Shuttle Body Flap
SS79-00250	Cover — Body Flap Segment
SS79-00251 2 Sheets	Rib-Stability Technology Demonstration Segment, Graphite/Polyimide
SS79-00252	Leading Edge Panels - Body Flap Segment
SS79-00253 3 Sheets	Spar Webs - Body Flap Segment







C SA MARTEN

6 - WANTER

7

.

Ć

1907 AND DIA HOLE POTTING PER DEPARTMENT O98 LTR 2423-4454 TOOLING HOLES AS REGID

----

8

· ----

TO INSTALL FASTENERS PER MADIOI- 301 CLASS

ŧ

4

- (9) DRILL 9/16 DIAMETER HOLE AND INSTA SL6149-3-770 TYPE R INSERTS M POTTING COMPOLIND PER DEPARTME D98 LTR-2423-4454
- A HE SUL WITH MBORD-002 FASA A HE SUS SYSTEM CORE PER D/095 JTR 2425-4430
- (3. UL TRASONIC INSPECT PER MT050-OIC
- & INSPECT LAMINATE MATERIALS PER LTR
- SPECIFIC PROCESS PER LIONE LTR
- 4 68074112/POLYIMIDE TAPE 0.0020 TH
- 3 LAMINATE ORIENTATION TOLERANCE: :
- 2. DIMENSIONAL LENGTH TOLERANCE: 10. ANGULAR TOLERANCE: 10'30'
- L. DIMENSIONS ARE IN INCHES

NOTES: UNLESS OTHERWISE NOTED

ŧ

4

				MODE FOR TEL	T FITTLE HAR	Η
						Η
· ·						1
						1
· · ·			• . 1			1
· · · ·						
						ł
						G
					1	
			······		·	
				1	1	
				,	÷	
			<u></u>		* 	
						F,
		1 1			1 	
				1 :	ł 	i
				1		
				1	1	
			······································	1	•	
		+	······	· ·	1 1	+-
				1		E
			······	· • · · · · · · · · · · · · · · · · · ·		i i
			!		1	<b>-</b>
				+	* *	¹
		<u> </u>			1	
		F/7 12:53	02:00:3 MASHE	2	3	
		160 MD144	00+1004 NUT	<u> </u>	<b>i</b>	÷ _
		181 DADHEN	220804 STREM			
		115 14042-5	002-244 SCREW	!	¢	
ADIOH-BOK CLASS 1		135 DID1210	02-00-01 STREW	+	Smulle LOC	
E INSERTS WITH		15	INSERT	1		÷
	1 1 11	1 5579-0	253 006 12 SPNE	<u>.</u>	1	
Male E Per	112	15579-00	1-00% STRAF	1225x643	• •	
		1 15574-30	1 006 TEE		104	
Rials per utr 2933-0062		1 15577-00		, 	165 Tu 00011 -001	
ES PER PROCESS		15574-0	29400, TEE		11570-00209-000	
R D/096 LTR 2425 - 445	1	1 357-00	258-00:	1	•	
	16	1		!	1 4	
22 0 0025 TH CK		2 557900	253-0021WEE	ŗ	<b>1</b>	
DLERANCE 22		11 5579-00	25200 I PANEL	i	¢	
PLERANCE: 10.03	1 13	1 1 15579-00	25H0051R1B ASSEM	BY	•	
		5579-00	250-008 20VER			
Ð			002 BONDASSY	7 <u>7</u> X8		
		1001 tss79-0	0249ISEGMENT			B
	HIEDEN RE DOWN DOWN BODY RECORD			AUTOM.	BUR ALLE South	37
	·		PARIS US:	4.6.		-1
ţ			5			
		MOLE NOTE: THEET	the in use	SAHTE/POL	nnol, "	A
	- 156.79 06.2879	- ACI 1998年 AND + 308-128 - ACI 1998年 328 + 422-42 - 331 1998年 228 + 428-2 - 331 1998年 228 + 428-2	11-7-7- 70-7		NT AT	
	Breghtern .	101 1401 140 - 101 - 10 70 1401 140 - 101 - 10 140 140 240 - 104 - 10		J 1 03953	5579-00245	

. . . .


























					- <u>T</u> -			<b>。</b>							
·								<u> </u>		#11			1		
:										bi score	· · · · · · · · · · · · · · · · · · ·		1		
											•				
											;		1 <b>H</b>		
														1	
a 											i				
· , .4															
											-				
										. <u></u>			G	1	
		•	•		i	1	<u>;</u>		<u> </u>	·	i +				
			•	+	1	1			<u> </u>		1 		{		
			, 	· · · ·	+	+	<del>.</del>		1		1			ſ	
			 		i	1			1	4	1 1				
					1		<u> </u>		1	1	•				
		•		<u> </u>		<u> </u>	<u>;                                    </u>		÷	<u> </u>					
						1	:		1	1	•	<u>-</u>	F		
	• 1 · · ·				t	1	t		<u> </u>	-	1		1		
		1	1		1							Ţ			
					1		<u>t</u>		1	<u> </u>	1	1	L		
							<u>i</u> t			1	t t		$\square$		
				+	+	<u> </u>	<u>;</u> ;		1	<u> </u> 			1		
					T		1		1				1		
				1	T		1			1			E		
		,					<u> </u>		<u> </u>		<u> </u>		{		
					+	<b> </b>	<u>:</u>		1	· · · · · · · · · · · · · · · · · · ·	•			•	
		1			I		I		<u>+</u>						
· .		1			T	ļ						1	+		
					+	<u> </u>			<u> </u>	<u> </u>	<u>.</u>				
	$\left  - \right $				+			·····		<u> </u>	·	_		•	
				1	1	Ē	1		1	<u> </u>	· · · · · · · · · · · · · · · · · · ·		D		
					T							1			
	$\vdash$		-+		+		<u> </u>		<b> </b>	 	¥ [			· .	
FIMIDE	$\vdash$		-+		+		<u> </u>		<u> </u>						
	H			1	+				İ.		• · · · · · · · · · · · · · · · · · · ·	+-	Π		
	$\Box$		$\square$	Ţ	T										÷
01-010	┝┼	;					[	<u> </u>	 	ļ	ŧ				
18 1TR 2433-4462	$\vdash$	-	+	-+-	+	<b> </b> -	<del> </del>		<u> </u>	<u> </u>	<b>.</b>	-+	c		
079-3334 SPECIAL					1					<u> </u>	ţ	-+			
16-50-50	ЦĪ		$\overline{+}$	-	$\Gamma$					ļ	È				
15 THICK	$\vdash$	-+	-+-	+	1.0	<b> </b>	FMA	4	ADJECT	ADH THE FUSA	-				
DI30-152A	$\vdash$			+	IAR					IN TOPTED OF	I DE STALE MERINE	CONSIGNAL OF			
E: 20.03					Ľ					1	<u> </u>		Ц		
	$\square$	1		1	T			· · · · · · · · · · · · · · · · · · ·	L						
	$\vdash$		+	-+-	<u> </u>		· · · · · ·			ļ	<u>.</u>				
	┝┥			┿	1.			003	CORE	SEE NOTES	(5)				
				1	12			002	SKINS	SEE NOTES	32(6)(7)(8)				
		1		1	1001		5379-00	251	RIB		k8X9)		B		
		2000											Η		
· · · · · ·			1		-				METS LIST						
	E		-					OR P	1.3007						
	$\pm$		+				,	17	MSt.	TECHNOL	ADILITY	R ]	A	-	
2001 3 157-00269 TES	<u></u>		E		-		1994 - 1994 - 1995 1994 - 1994 - 1995 1994 - 1994 - 1995		Inc.n	- SEGMENT	, GHANNITE/POLYIN	ALDE -			
	1									J 03953	15579-0025				
3	<u> </u>				,			2		1101171	1	112			
					·										





ŧ 

ï





		RIB ASSEMBLY FOR TECHNOLOGY DEMONSTRATES SEGMENT
-		J 03953 5579-00251
	2	A State of the second second

-004 FRANT NUS -004 FRANT NUS CLARK NUT SEE MOTES -000 ATTACA ATTACA -000 ATTA

-

HI MAN

in.

15 PARTS LIST (3(6)(7)(8)(1) (3(4)(6)(8)

K3X4X6X7X8

クニニ ===

.

2

MANATE BRIENTATION TOLERANCE: 12"				1	-004
GULAR TOLERANCE : : 0'20'		ł		1	-603
MENSIONS ARE IN INCRES				2	
SI WILESS OTHERWISE SPECIFIED				1	3079-40851-401
				<b>a</b> 5	
	C.				

3 ¥.

 $\mathcal{F}$ 

3

-

6 DI NOTE:

13 3

22

Χ.

10

.

4

1. A. C. S.

- ( LANSMATE BRIENTATION TELERNICE : 12" 2. DEALASIANAL LENGTH TOLERANCE : 10,03
- C GRAPHITE POLTMARE TAPE Q. MES' THICK CELINH 6000/LANC BD MER MBISD- MERA
- SPECIFIC PROCESSES PER BYONE-EVES- ++E+ (5) HEXCEL CORP. CORE HRW-387-% -.75-3.0
- (T) REPECT LANGENTE MATERIALS PER LTR 2435-4462 G FASAREATE GA/PI LAMINATES PER PROCESS SEL MADINE 253, CURE CIELE PER 30077-338, 6
- SYSTEM PER DIDIS LTR 2423-4454 ( ULTRASONE WEFELT PER MTOFOI-ON
- ( POTTING COMPOUND PER D/048 LTR 2423-4454 T BOND WITH MEDIED-OLE FMS+ ADDESIVE
- TIBERITE IN BRAPHITE / POLYMAIDE CLOTH . DOTS THICK 3 LAYERS
- .

4

3

T ī 1 1 1 ł ÷ + 1 1 1 . ŧ. 1 f ł ł ł 1 1 í. i i T 1 1 Ð ... ŧ i ī 1 T C ŧ 1 .i 1 . ŧ 5579-80E8-805 ASSEMBLY 1 2 15579-00E53-003 TEL

2

2000

1 4

-----

H

G

F

E

63

-----

270

25

Ŧ

5579-00251-003

I MAN









								-						ł
								T BERNEYING	MARKET I DE RECORDE		80 160			
-									PARTS LIST					Į.
		+												
		П			1				DE P					
					1				the state of the	LEAD A	- EDGC - AD 55	GMENT	3	1
	-001	#	579-00250				12 3	222	10					{
						ha				11039	5. 557	9-0025Z	. 1	
		<u> </u>			-					T 19935	ante la state	5. 1958au		
4				3	• .		[		2	Sec.			194 E	
-	1	-								20	in the		1	
Additional of the	50 mg 7								·····		and the second second			

		+		i E				1 <b>^</b> :			
				1	_		PARTS LIST		. <del> </del>		
-	HERE BEIT	000				CONTRACT INCOME	I OF RECOVERS		ES C'E SPPER		
	<b>GHT CT T</b>	 	ا <del>ہے جم</del>	1077	1000	1.3579-002 5 E		SEE NOT		$\square$	B
ILESS OTHERWISE SPECIFIED		!	1	11	<u> </u>	- 00Z	INK CORE	SEE NOTE	1 <u>5</u> , <del>1 <u>5</u></del>		12
INS ARE IN INCHES				12	ļ	-003	SKIN	SEE NOTES	5) 4 Y SY D D		5
R TOLERANCE : 10'50		1	1	11	!	-004	SKIN	SEE NOTES	5, E ( ) ( ) ( )		8
LE ORIENTATION TOLERANCE : 1 2"			1	1 Z		-005	I CHANNE:	SEE NETES	<u> DDDD</u>		25
6000/LARC IGO PER MBDISO- ISZA		ļ	1	1	Ī	1	1	1			
DPP. CORE HRH-327-345-35 TO 36 PCF			- <u>+</u> -	Ť	i	i	1		·		2
PROCESSES PER DO90-2423-4454		i	-+	- <u>-</u>	<u>.</u>	<u>.</u>			· · · · · · · · · · · · · · · · · · ·	$\vdash$	$\Box$
TE GRIPH LAMINATES PER MAT'L SPEC			-+		1	1 KM.34	ADHESIVE	SUMONE :	CH STYLE FERTILLAS CARE	MR.	
LAMMATE MATERIALS PER LTR 213-4462				+	<u> </u>	ļ		1 AT TYPE ANT	5		
23 - 4454 W WSBCCT BER MIDEOL-DID				+	!	<u> </u>	<u></u>	<u> </u>	۶ ۲		
TH MBOIZO-062 FM34 E System Per D/098			_	1	<u> </u>	1			·		
_			1	Î	1		1		۲ 		
		1	1	Ť	1	;	1	1	3		
	1	1 1	$-\frac{1}{1}$	÷-	1	<u>.</u>			\$	<u> </u>	
		<u> </u>				, t		1			H
	<u> </u>							<u> </u>	\$ <del>;</del>	$\vdash$	1
	<u>+</u>	<u>'  </u>	÷					1	د •	$\vdash$	
-		1	1		!	!	i	1	· •		
	1			1	;	1			s		D
	1			1	ł	1		1	r		
	1 1	1		<u>-</u>		+		i ī	<u>.</u>		
					÷	·		1 	, ,,,,,,,,,,,,_		
				<u>.</u>		·		: 	1 		-
					!	<u>.</u>			s •	<u> </u>	
		<u>.</u>		.!	·	<u>.</u>		I	۱ ۱		

----13 н ł İ. G 1 1 ł ÷ , : 1 i. ī i . ł + ł -1 1 • ÷ 1 ł. F ī . Ī 1 , 1 -! ÷ ; ; ; : 1 i , 1 i . . . 1 1 4 ł ŧ -f + ł 1 1 Ī ŧ. 1 1 + 1 1 : 1 • 1 ; . 1 i 1 1 ÷ 1 ł ÷ • 1 İ E . 1 1 1 1 ÷ ÷ Ī 1 8 1 1

:

Ì p

ŧ,

1

Т

3

4

۰.

2













	•	٠	 ·

.







Ð

. .





				5579-00253 3	1
+	<u>بمبد</u>	43	42	41	40
				1	

.___



---

41 1





1. Report No.	2. Government Accessi	on No.	3. Reci	pient's Catalog No.		
A Title and Subtitle				vet. Date		
Development and Demonstrat	tion of Manufactu	ring Proc	esses Dec	ember 1981		
for Fabricating Graphite/	LARC-160 Polyimid	e Structu	ral 6 Perfo	rming Organization Code		
Elements	• •		0	inning organization code		
7. Author(s)	· · · · · · · · · · · · · · · · · · ·		8. Perfo	orming Organization Report No.		
R.K.Frost J.S.	Jones					
P.J.Dynes D.H.V	Wykes		10 18/0-1			
9. Performing Organization Name and Addres		/	10. 10.			
Rockwell International						
12214 Lakewood Blvd.			11. Cont	ract or Grant No.		
Downey, Ca. 90241			NAS	1-15371		
			13. Type	e of Report and Period Covered		
12. Sponsoring Agency Name and Address			Con	tractor Report		
National Aeronautics and	Space Administrat	ion	14. Spon	soring Agency Code		
Washington, D.C. 20546						
15. Supplementary Notes	<u></u>		<b>I</b>			
Langley Technical Monitor	: Robert M. Bauc	om				
Final Report						
16. Abstract						
The program consisted of tw stration Components. Proce the basic composite materia components, developing proc processes through mechanica components were fabricated components consisted of fla fiber compression moldings, of the Space Shuttle aft bo be subjected to mechanical 400 cycles between 5-100% o between -107°C and 316°C.	o parts: Process ss development in l and processing, esses for specifi l testing. In th using the process t laminates, skin and a Technology dy flap. The TDS loading at room t f limit load at 2 TDS test results	Develop cluded es nondest c structu e second es develo /stringe Demonst is a fu emperatu 60°C; and will be	ment and Fabri stablishing qu ructive inspect iral forms, and part of the p oped in part of r panels, hone rator Segment 11-size testab re and 260°C; d thermal cycl reported in a	cation of Demon- ality assurance of tion of fabricated d qualification of orogram, demonstration one. The demonstration ycomb panels, chopped (TDS) representative ole component and will simulated fatigue, ing, 125 cycles separate final report.		
17. Key Words (Suggested by Author(s))		18. Distributi	on Statement			
LARC-160 Polyimide Resin, Gr	raphite/Polyimide					
Celion/LARC-160, Polyimide H	Processing	171	offica _ 11-1:-	nited		
Polyimide Characterization		Uncias	stited - outfi	nrea		
Polyimide Fabrication						
Polyimide Adhesive Bonding			Subjec	t lategory 24		
19. Security Classif. (of this report)	20. Security Classif. (of this	page)	21. No. of Pages	22. Price		
Unclassified	Unclassified		498			
For calo by th	ha National Tachainal Infor	mation Convio	o Springfield Virgin	- 221C1		

For sale by the National Technical Information Service, Springfield, Virginia 22161