UNCLASSIFIED

AD NUMBER

ADB010414

NEW LIMITATION CHANGE

TO

Approved for public release, distribution unlimited

FROM

Distribution authorized to U.S. Gov't. agencies only; Administrative/Operational Use; DEC 1975. Other requests shall be referred to Federal Aviation Administration, Supersonic Transport Office, 800 Independence Avenue, SW, Washington, DC 20590. Export Control.

AUTHORITY

faa ltr, 26 apr 1977

THIS PAGE IS UNCLASSIFIED



ADB010414

SST Technology Follow-On Program-Phase II

DEVELOPMENT AND EVALUATION OF FUEL TANK SEALANTS

Marlan Pollock

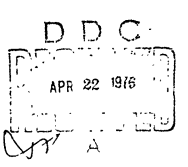


D6-60282 December 15, 1975

FINAL REPORT Task II

Approved for U.S. Government only This document is exempted from public availability because of restrictions imposed by the Export Control Act. Transmittal of this document outside the U.S. Government must have prior approval of the Supersonic Transport Office

Prepared for FEDERAL AVIATION ADMINISTRATION Supersonic Transport Office 800 Independence Avenue, S.W. Washington, D.C. 20590



The contents of this report reflect the views of 'The Boeing Company, which is responsible for the facts and the accuracy of the data presented herein. The contents do not necessarily reflect the official views or policy of the Department of Transportation. This report does not constitute a standard, specification, or regulation.

TECHNICAL REPORT STANDARD TITLE PAGE Government Accession No 3 Recipient's Catalog No FAA-SS 5 Report Date SST TECHNOLOGY FOLLOW-ON PROGRAM-PHASE II December 15, 1975 DEVELOPMENT AND EVALUATION OF FUEL TANK 6 Performing Organization Code SEALANTS Performing Organization Report No. Marlan/Pollock D6-69282 v Performing Organization Name and Address **Boeing Commercial Airplane Company** P.O. Box 3707 DOT_F Seattle, Washington 98124 12 Sponsoring Agency Name and Address **Federal Aviation Administration** Supersonic Transport Office 800 Independence Avenue, S.W. 14. Sponsoring Agency Code Washington, D.C. 20590 15 Supplementary Notes The most promising candidate for sealing fuel tanks of the U.S.A. supersonic transport is a fluorosilicone designated DC 77-028 and DC 94-529. The sealing system was exposed to various environments and the effects evaluated on appearance, physical properties, and sealing effectiveness. Test specimens were exposed to more than 36 000 hr of 219 to 227 C (426 to 441 F) fuel vapor in an accelerated cycle and more than 8000 flight cycles. A sealed tank simulating SST construction was exposed to more than 16 000 hr of 219 to 227 C (426 to 442 F) fuel vapor in an accelerated cycle and \approx 6500 torsional load cycles each at -46 C (-50 F), +232 C (+450 F), and room temperature. Confidence was developed that the DC 77-028/DC 94-529 fluorosilicone system would perform for 50 000 flight-hours as an integral fuel tank fillet sealant in a about Mach 2.7 commercial supersonic airplane. Further development is necessary, however, to achieve this level of confidence for faying surface and injection sealant systems. Progress was also made in developing a backup sealant based on fluorocarbon, and exposing and testing were conducted on a fluorosilicone-fluorocarbon hybrid developed by Dow Corning under an Air Force Materials Laboratory contract. 17 Key Words Approved for U.S. Government only. This document Sealant is exempted from public availability because of restrictions imposed by the Export Control Act. **Fuel Tank** Transmittal of this document outside the U.S. Supersonic Government must have prior approval of the **Temperature Resistant** Supersonic Transport Office. 19 Security Classif (of this report) 20 Security Classif (of this page) 21 No of Pages 22 Price Unclassified Unclassified 286 Form DOT F 1700 7 (8-69)

PREFACE

Among the agencies participating in this program by furnishing samples for test, doing developmental work, and exchanging information were Dow Corning, General Electric, Products Research and Chemical Corporation, the Air Force Materials Laboratory, and NASA-Ames. The Boeing Company is especially grateful to the Dow Corning Corporation for its continuous interest, diligence, and close cooperation.

The work was administered by the Supersonic Transport Office of the Department of Transportation. The technical monitors were C. R. Ritter and Charles Troha. The work was performed at the Renton Materials Technology Laboratory, the hazardous test cell at Boeing Field, Seattle, and the Materials Technology laboratories at Wichita.

PRECEDING PAGE BLANK-NOT FILMED

CONTENTS

à

Page

in in Hereinia

and and a

and the second second

And a second second

いいたいないまたがっていますがいていたいないない。 このはないない ないないない ないない しょうしょう

ŵ

•,

A CONTRACTOR

.

3

ħ,

0	INT	RODU	CTION	••••••	1
2.0	TES	т мет	HODS AN	D PROCEDURES	5
	2.1			nental Exposures	5
		2.1.1		cle	5
		2.1.2		ed Cycle	5
		2.1.3		Cycles	5
	2.2			est Specimens	5
		2.2.1		of Test Panels	5
		2.2.2		on of Primer	12
		2.2.3		on of Sealant	12
		2.2.4	-	ure	13
		_ ,		Fluorosilicone	13
			2.2.4.2	Fluorosilicone-Fluorocarbon Hybrid	13
	2.3	Test F			13
		2.3.1		and Mechanical Properties	13
		2.0.1		Tensile Strength and Elongation	13
				Hardness	14
				Weight Loss	14
				Volume Change	14
				Adhesion to Titanium (Tee Peel)	14
		2.3.2		I Testing	15
		D.O.D		Picture Frame Shear	15
				Flight Cycling Fillet Deflections	18
		2.3.3			18
		2.0.0	10000 1000	• • • • • • • • • • • • • • • • • • • •	10
3.0	RES	SULTS			21
0.0	3.1			ated and Flight Cycling	21
	3.2			area and 1 mino cycling	21
		3.2.1		rame	21
		3.2.2		lection	39
		3.2.3		ting	39
				0 0	•••
4.0	BAG	CKUPS	SEALANT		47
	4.1			nn	47
	4.2			bon-Fluorosilicone	47
ō.0	COI	NCLUS	IONS		48
6.0	REC	COMMI	ENDATION	IS	49
APF	'ENI			ION OF THE PROPOSED SST FUEL ALANT IN SMALL SIMULATED FUEL TANKS	51
APF	PEME			E, EASILY APPLIED FLUOROCARBON S FOR SUPERSONIC AIRCRAFT FUEL TANKS	213

v

~

- -

FIGURES

EN.

'n.

\$

NO.		Page
1	Typical Fillet, Faying Surface, and Injection Scaling	3
2	Standard Flight Cycle	6
3	Flight Cycle Apparatus	7
4	Flight Cycle Apparatus Without Insulation	8
5	Specimen Loading of Flight Cycle Chamber	9
6	Accelerated Exposure Cycle	10
7	Accelerated Cycle Exposure Chamber	11
8	Peel Strength Testing	16
9	Picture Frame Shear Specimen	17
10	Picture Frame Test Fixture	19
11	Flight Cycle Deflection Specimen	20
12	Accelerated Cycle Exposure of DC77-028 Lot 1222,	
	Tested at Room Temperature	22
13	Accelerated Cycle Exposure of DC77-028 Lot 1222,	
	Tested at -46° C (-50° F)	23
14	Accelerated Cycle Exposure of DC 77-028, Lot 1222,	
	Tested at 232° C (450° F)	24
15	Specimens From Wichita Tank, Lot 205180, Tested at	
	at Room Temperature	25
16	Specimens From Wichita Tank, Lot 205180, Tested at -46° C (-50° F)	26
17	Specimens From Wichita Tank, Lot 205180, Tested at 232º C (450º F)	27
18	Accelerated Cycle Exposure of DC 77-028, Lot 401117, Tested	
	at Room Temperature	28
19	Accelerated Cycle Exposure of DC 77-028, Lot 401117,	
	Tested at -46° C (-50° F)	29
20	Accelerated Cycle Exposure of DC 77-028, Lot 401117,	
	Tested at 232° C (450° F)	30
21	Flight Cycle Exposure of DC 77-028, Lot 401117, Tested at	
	Room Temperature	31
22	Flight Cycle Exposure of DC 77-028, Lot 401117, Tested	
	-46° C (-50° F)	32
23	Flight Cycle Exposure of DC 77-028, Lot 401117, Tested	
	at 232° C (450° F)	33
24	Flight Cycle Exposure of DC 77-028, Lot 206177, Tested at	
	Room Temperature	31
25	Flight Cycle Exposure of DC 77-028, Lot 206177, Tered at	
	-46° C (-50° F)	35
26	Flight Cycle Exposure of DC 77-028, Lot 206177, Tested	
	at 232° C (450° F)	36
27	Weight and Volume Loss for DC 77-028, Lot 401117	37
28	Fillet Seal of Picture Frame Panel After Exposure	38
29	Test Tank	40

NT

in a station of the state of th

States and Street

ં કરી કેલ્ડોલ્ડ જે ત્યાં છે.

の言いたちのと

and the second second second

> Contraction of the and the second

FIGURES (Concluded)

.[()	Scalant Condition After Exposure	/# 8
31	Repaired Aren	43
32	Lower Web-to-Chord Faying Scal Location	48
	Stiffener-to-Web Faying Seal Location	
	Injection Seal	-

TABLES

No.		Page
1	Exposure Times for the Test Tank	39
2	DC 77-108 Test Results	47

1.0 INTRODUCTION

A major technological requirement of the U.S.A. supersonic transport program was to ultimately provide a commercial supersonic aircraft which would be competitive from a maintenance standpoint with current subsonic aircraft. The need for an elastomeric material to function as an integral fuel tank sealant in the environment generated by a Mach 2 7 commercial supersonic aircraft was a major requirement which necessitated an extensive sealant research and development program by both The Boeing Company and its material suppliers.

Polysulfide-type sealants now used in subsonic jet aircraft have performed satisfactorily for more than 55 000 hr of flight time on individual aircraft. Fuel tank leakage caused solely by sealant degradation is almost nonexistent and, if mechanical damage to the sealant occurs, repairs are easily accomplished.

Integral fuel tank sealant requirements for the SST impose technical and environmental requirements far beyond the capabilities of the subsonic polysulfide sealants. The SST sealant material must be able to accommodate structural strains over a temperature range of -46° to $+232^{\circ}$ C (-50° to $+450^{\circ}$ F) under tank conditions of hot fuel and fuel vapor. It also must adhere to titanium alloy 6Al-4V and be inert to both the titanium alloy and the fluids encountered in manufacturing and flight operations. Finally, it must function satisfactorily for at least 50 000 flight-hours. A search for such a sealant was begun in 1961 through literature search and supplier contacts. Every available sealing system in use on supersonic airplanes was investigated to ascertain its suitability. When the SST fuel tank sealing program was initiated in 1965, it was obvious that only two elastomeric polymer systems could reasonably be considered as leading candidates on the basis of thermal stability and resistance to fuels. These were the fluorocarbons and the fluorosilicones.

Even though their thermal and fuel resistances are excellent, the fluorocarbons such as the Fluorels and Vitons have a tendency to cause stress corrosion of titanium and currently depend on being suspended in solution for proper application. Volatilization of the solvent precludes their effectiveness as a filleting sealant, since they must be applied in thin layers to prevent excessive porosity from outgassing. The fluorocarbons also have unacceptable low-temperature flexibility.

A MARK A REPORT

On the other hand, at low temperatures the fluorosilicones exhibit excellent properties which get progressively poorer as the temperature increases. Under conditions of high temperature and pressure, they tend to revert to a gummy mass. In 1967, Dow Corning introduced a fluorosilicone which was a significant improvement over the others. Originally, it was designated DC 94-516 which later was changed to DC 77-028. Today it is marketed as DC 94-529.

Dow Corning and Boeing worked closely together to develop the fluorosilicone system with the emphasis on improving reliability of adhesion and repairability and creating satisfactory faying surface and injection sealants. Primary sealing of the SST was to be by means of fillet seals, but in many situations it was necessary to use fillet sealing in

conjunction with other sealing methods. The faying surface sealant passed laboratory tests but was suspect because of visible channels through it and failure to seal the cover of a test tank. The problem with injection sealant was one of excessive thermal expansion, which caused it to extrude and to tear itself and the fillet seal covering it.

The basic fillet sealant, designated DC 77-028, was varied to obtain properties that would make it more suitable for faying surface and injection or prepack sealing by using different fillers and combinations of fillers. The faying surface sealant was designated DC 77-053, and the injection or prepack sealant was designated DC 77-066. Figure 1 shows how these sealants are used.

Another obstacle to determination of the suitability of a fuel tank sealant material for SST application was the lack of complete physical property requirements. On this basis, Boeing established the following fillet sealant target for 50 000-hr flight service life in the U.S.A. SST.

Technical Requirements

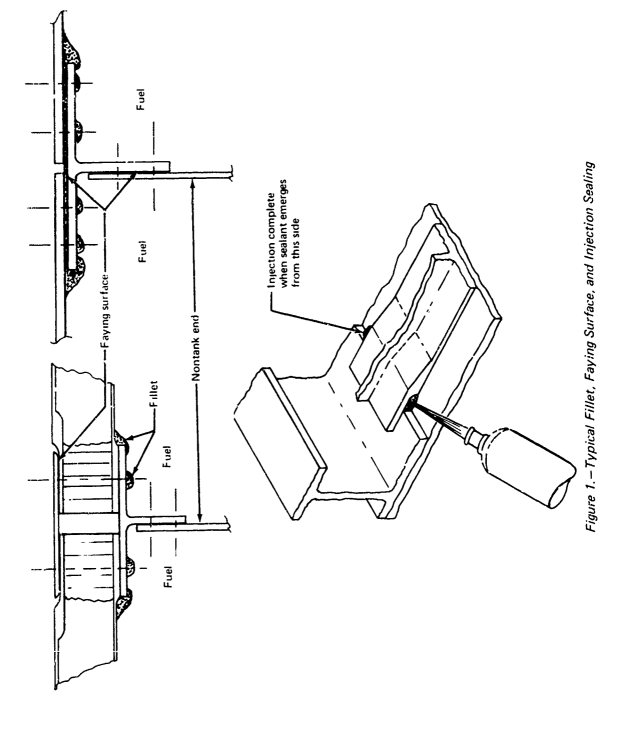
and a star with the second star and the second star and the second star and the second star and the second star

f i

ester about the sub-tree su

ومعالك المشاهدة والمتلك ومسالا مطارعتها وتتعاري

1.	Adhesion to 6Al 3V titanium (peel and tear resignance)	1.07 kg/cm (6 lb/in.), 85% cohesive failure, minimum
2.	Tensile elongation	15% minimum
3.	Shore A hardness	Less than 20 points change
4.	Weight change	Less than 20%
5.	Volume change	Less than 10% loss
6.	Deflection simulation [0.028-cm (0.011-in.) deflection]	Less than 30% tear after 100 cycles
7.	Leakage (picture frame)	No leaks
App	lication Requirements	
1.	Flow (slump)	1.25 cm (0.5 in. maximum in 2 hr
2.	Extrusion rate	15 g/min. minimum
3.	Cure	Less than 71° C (160° F)
4.	Working life	At least 0.5 l.r
5.	Repairability (peel test)	1.07 kg/cm (6 lb/in.), 85% cohesive failure, minimum



法認知

Ŕ.

A state of the second s

ł

- -

Work on the basic DC 77-028 fluorosilicone system was conducted through the SST prototype program and through a Phase I follow-on development program, under contract DOT-FA-SS-71-12, awarded by DOT after termination of the SST activities. The results of these efforts led to improvements in the sealant characteristics and test and evaluation criteria as reported in D6-60221, *Development and Evaluation of Fuel Tank Sealants*. Based on these findings, a Phase II technology follow-on program under contract DOT-FA-72WA-2893 was awarded to continue development of the system and demonstrate its capabilities for commercial supersonic aircraft applications. The primary objectives of the Phase II effort were:

- 1. Conduct 30 000 hr of simulated service testing at elevated temperatures.
- 2. Verify suitability for 50 000 flight-hours.
- 3. Develop improved adhesion.

er an stade for the state of the state of the base of the state of the

- 4. Develop repair techniques.
- 5. Develop faying surface/injection sealant capabilities.

Also as part of the Phase II program. a search was conducted for a backup to the fluorosilicone system. A contract was issued to Products Research and Chemical Corporation to develop a practicable sealant based on a fluorocarbon system pioneered by the Air Force Materials Laboratory. Both NASA and AFML were active in funding fuel tank sealant development programs, and these programs were followed closely by Boeing. Another backup candidate was a hybrid fluorocarbon-fluorosilicone developed by Dow Corning under an AFML contract. AFML supplied samples for test.

2.0 TEST METHODS AND PROCEDURES

The approach was to subject test specimens to various static and dynamic exposures and to measure changes in physical and mechanical properties. Responses of the sealant to the exposures were used as indications of useful life and deficiencies of the basic material or system. Functional tests, which measured sealing ability depending on all properties of the total system, were also conducted to evaluate the practical aspects of the sealant's use

2.1 CYCLING ENVIRONMENTAL EXPOSURES

Two principal cycling environments were used: flight and accelerated. The test tanks were exposed to cycles similar to the accelerated cycle.

2.1.1 FLIGHT CYCLE

The simulated flight cycle (fig 2) provides about 50 hr per week of elevated-temperature fuel vapor exposure. The equipment is shown schematically in figure 3 and photographically in figure 4. The loaded specimen rack is shown in figure 5.

2.1.2 ACCELERATED CYCLE

The accelerated cycle (fig. 6) exposes the specimens to a minimum of 140 hr per week of elevated-temperature fuel vapor. The exposure chamber is shown in figure 7 with the test specimens in place

2.1.3 TEST TANK CYCLES

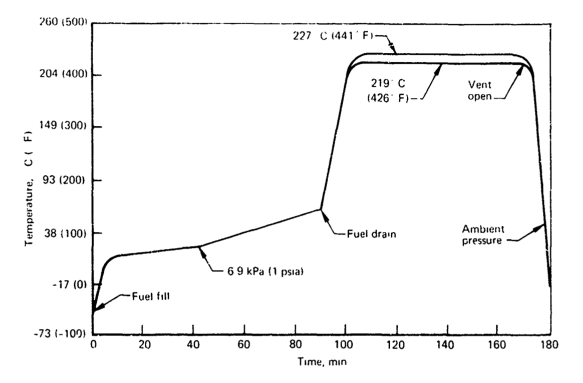
Environmental exposure cycling of the tank was the same as for the accelerated cycle except that ritrogen at atmospheric pressure was substituted for air at 6.9 kPa (1 psia). The tank was not designed to withstand the near-vacuum pressure.

2.2 PREPARATION OF TEST SPECIMENS

2.2.1 CLEANING OF TEST PANELS

All test panels and parts were cleaned by applying cleaner from clean polyethylene (or equivalent) squeeze bottles. The cleaner consisted of:

	Volume percent
Aromatic naphtha (TT-N-97, type I, grade B)	50
Ethyl acetate (TT-E-751) or isopropyl acetate (TT-1-721)	20



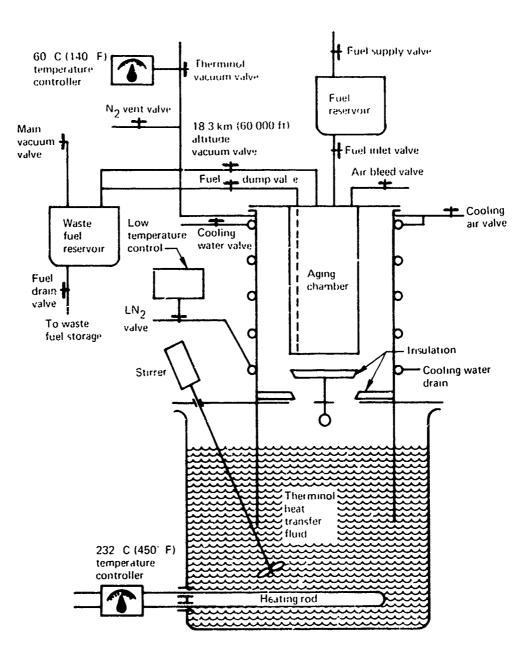
nen förstallaring som att den som det naktionen att att det som att att det som att det som att det som att det

ber alle and the set of a log of the state of the set o

فليغوض فالمحيطة والمعرفة فالمعافلا فالمترج وتعمدتها للتمنيفة ومهما الافاد أنطحه فلاطوه

ner sen ner sin her bestelsen is is in er state er state figtig til se er state bisket bitte bet etter state s

Figure 2.-Standard Flight Cycle



1.

ولاللاف فشارك أستانك

at the second solid

فتشتاه أدماك كموطحا والا

Figure 3.-Flight Cycle Apparatus

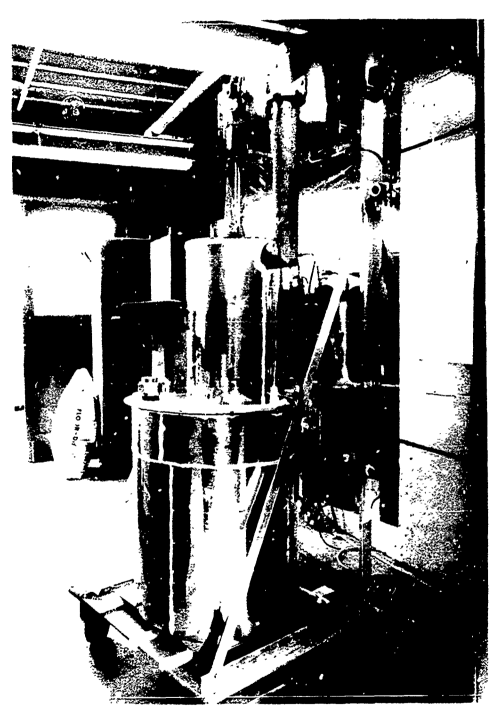
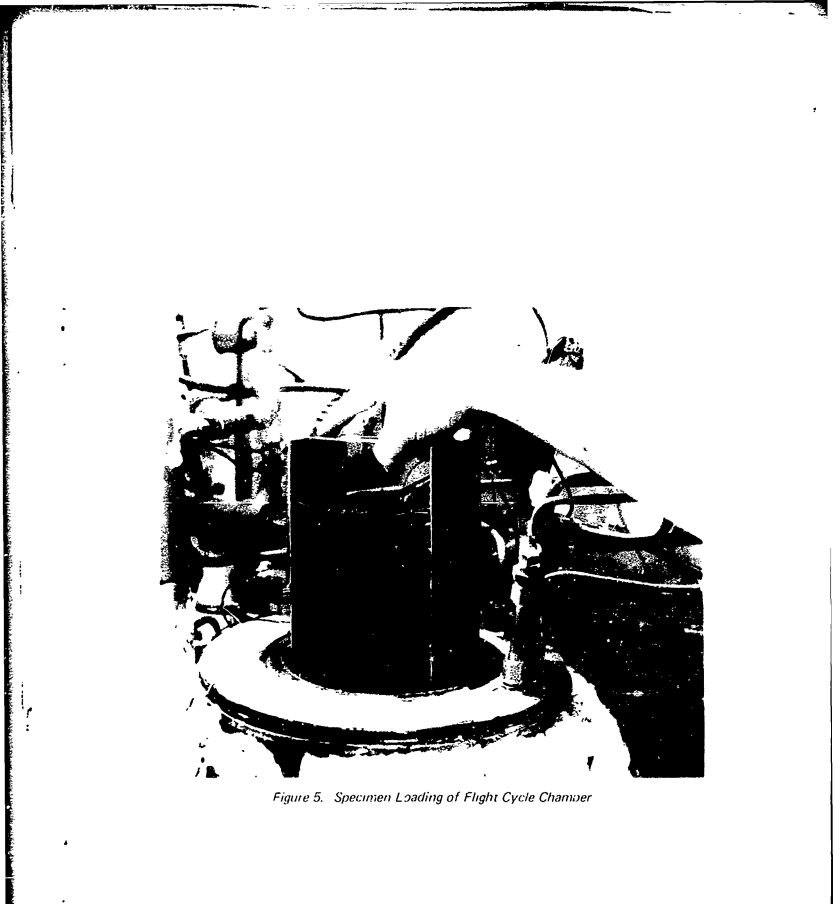


Figure 4 Flight Cycle Apparatus Without Insulation



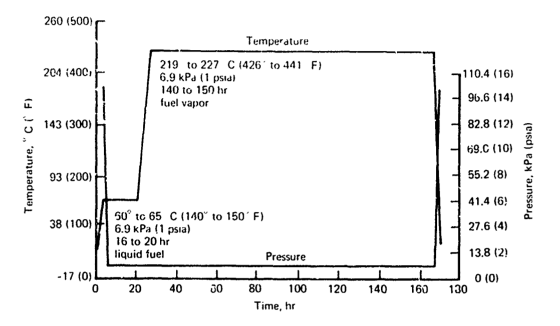


Figure 6. – Accelerated Exposure Cycle

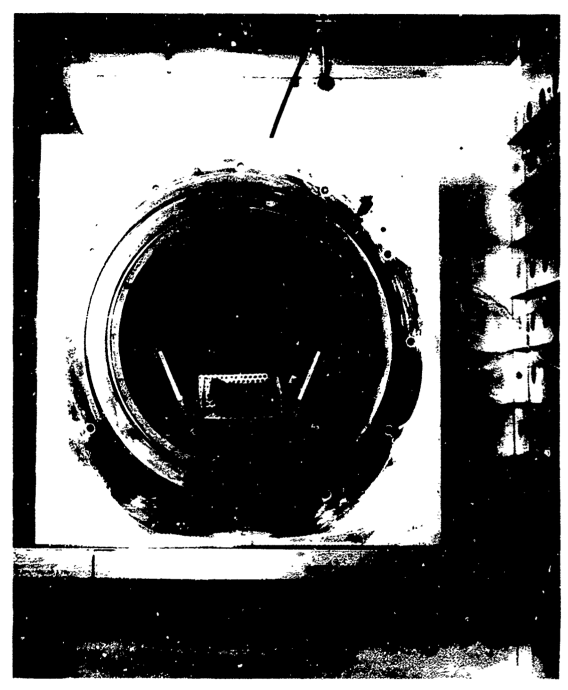


Figure 7. – Accelerated Cycle Exposure Chamber

Methyl ethyl ketone (377-M-261)	20

Isopropyl alcohol (MIL-F-5566) 10

The cleaner was applied directly from the bottle to the panel or part so that the entire surface was wetted. The surfaces to be cleaned were thoroughly scrubbed with clean gauze pads wetted with cleaner. The solvent was wiped off while wet with clean, dry gauze pads. This procedure was repeated as required to produce a clean surface as determined by no visibly detectable residue on the gauze. Each gauze and was used for one scrubbing or drying application only. Cleaned panels were not stacked, but were covered with paper toweling or equivalent until used. Test panels and parts were used within 8 hr after cleaning.

2.2.2 APPLICATION OF PRIMER

A coat of primer was applied to cleaned test panels before applying the sealant. The primer was applied in a continuous coat as uniformly as possible using a gauze pad. The primed panels were dried for 30 ± 10 min at $24^{\circ} \pm 1.5^{\circ}$ C ($75^{\circ} \pm 5^{\circ}$ F) and $50\% \pm 5\%$ relative humidity. Then the panels were immediately placed in an oven and the primer was cured at 232° C (450° F) for 25 ± 5 min. After curing, the primed surfaces were protected from contamination until sealant application. If sealant was not applied within 90 min after the primer was cured, the panels were cleaned and reprimed.

2.2.3 PREPARATION OF SEALANT

というないないのである。ためではないです

Prior to mixing, two-component sealant compounds were stored at $24^{\circ} \pm 1.5^{\circ}$ C $(75^{\circ} \pm 5^{\circ}$ F) for a sufficient time to allow the material to reach a state of equilibrium with that temperature. The activator was added immediately before weighing. Mixing instructions were as follows:

- 1. Weigh the correct amounts of base and activator onto a clean. flat stainless steel plate or pan immediately prior to mixing. The activator must not be allowed to contact the plate.
- 2. Handmix the sealant compound by folding and squeezing with a spatula for a minimum of 5 min until the sealant compound appears uniform.
- 3. Spread the sealant compound on a clean, flat stainless steel plate or pan so that the maximum depth is less than 1.21 cm (0.5 in.). Vacuum degas the sealant compound for 10 min at 1.73 kPa (0.25 psia) or less.

- 4. Remove the plunger and plug the nozzle end of a cartridge for the Semco 250 gun. Scoop up the sealant with a spatula, place in the open end of the cartridge, and drive down by sharply rapping the nozzle end of the cartridge on something solid. Repeat until the cartridge is filled.
- 5. Vacuum degas the filled cartridge for 5 min at 1.73 kPa (0.25 psia) or less. Use a plastic film as necessary as an extension of the cartridge to prevent overflow of the sealant. Place the plunger in the cartridge using care to minimize air entrapment.

When required, the sealant was put into refrigerated storage at or below -40° C $(-40^{\circ}$ F) immediately after being placed into the cartridges. Dry ice was not allowed to be used for refrigeration. The sealant was stored at or below -40° C $(-40^{\circ}$ F) for a minimum of 16 hr but not longer than 72 hr and conditioned at $4.4^{\circ} \pm 1.5^{\circ}$ C $(40^{\circ} \pm 5^{\circ}$ F) for 4 ± 2 hr immediately prior to thawing. It was thawed by vertically immersing the frozen cartridges in a $49^{\circ} \pm 0.6^{\circ}$ C $(120^{\circ} \pm 2^{\circ}$ F) water bath for 4 min ± 5 sec with the plugs installed and the upper end of the cartridge 2.54 cm (1 in) above the liquid level.

In BANK to the state of the second

2.2.4 SEALANT CURE

2.2.4.1 Fluorosilicone

Standard cure was 14 consecutive days at $24^{\circ} \pm 1.5^{\circ}$ C $(75^{\circ} \pm 5^{\circ}$ F) and $50\% \pm 5\%$ relative humidity. An accelerated cure consisting of 24 hr (minimum) at $71^{\circ} \pm 3^{\circ}$ C $(160^{\circ} \pm 10^{\circ}$ F) followed by 1 hr (minimum) at $149^{\circ} \pm 3^{\circ}$ C $(300^{\circ} \pm 10^{\circ}$ F) was sometimes used instead of the standard cure.

2.2.4.2 Fluorosilicone-Fluorocarbon Hybrid

The FCS-210, also known as DC 77-108, was cured 1 hr at 100° C $(212^{\circ}$ F) followed by 1 hr at 150° C $(302^{\circ}$ F).

2.2.5 PREPARATION OF SEALANT SLABS

The sealant was cast to a thickness of 0.32 ± 0.02 cm $(0.125 \pm 0.008$ in.) in a closed mold lined with Teflon. The mold was filled by extruding the sealant from a sealant gun with a Semco 440 nozzle. The nozzle was freed of air by a preliminary extrusion of 5 to 7 cm (2 to 3 in.) of sealant. During the casting operation, the tip of the nozzle was placed in an injection mold and was not removed until the mold was filled to excess.

When the standard cure was used, the sealant was kept in the closed mold until the cure was completed or removed at any time after 96 hr. When the accelerated cure was used, the sealant was allowed to remain in the mold at $24^{\circ} \pm 1.5^{\circ}$ C ($75^{\circ} \pm 5^{\circ}$ F) and $50\% \pm 5\%$ relative humidity for a minimum of 48 hr prior to the 71° C (160° F) exposure. The slabs were removed from the mold before completing the cure at 149° C (300° F).

2.3 TEST PROCEDURES

2.3.1 PHYSICAL AND MECHANICAL PROPERTIES

2.3.1.1 Tensile Strength and Elongation

Ultimate tensile strength and elongation were determined in accordance with ASTM D-412 using a jaw separation rate of 51 cm (20 in.)/min. Miniature specimens were used cut from slabs prepared as described in section 2.2.5. Five specimens were tested for each point except for those from lot 401117 where eight were tested per point.

2.3.1.2 Hardness

Shore A durometer hardness was determined in accordance with ASTM D-2240 taking the median value as the hardness for each specimen. The hardness is reported as the average of four specimens. Volume chang: specimens were used for hardness measurements.

2.3.1.3 Weight Loss

Specimens approximately 2.5 by 5 cm (1 by 2 in.) were cut from a slab of sealant prepared in accordance with section 2.2.5. Before and after the applicable environmental exposure period, the test specimens were conditioned for 24 hr in a dessicator and then weighed immediately.

Percentage weight loss was calculated as follows:

$$\frac{(W_1 - W_2) \times 100}{W_1}$$

where

and the second second second

 W_1 = weight of sample before aging

 W_2 = weight of sample after as $_{4g}$

Percentage weight loss for each determination is the average of four specimens.

2.3.1.4 Volume Change

Specimens approximately 2.5 by 5 cm (1 by 2 in.) were cut from a slab of sealant prepared in accordance with section 2.2.5. The volume change of environmentally aged specimens was determined in accordance with ASTM D-471. Volume change is reported as the average of four specimens.

2.3.1.5 Adhesion to Titanium (Tee Peel)

Details of Specimen Preparation.-The required number of 0.127- by 7.4- by 15.2-cm (0.05- by 2.9- by 6-in.) panels were prepared from annealed 6Al-4V titanium alloy. An equal number of 7.4- by 30.5-cm (2.9- by 12-in.) strips were prepared from 200-mesh stainless steel screen.

The panel surfaces and screen were cleaned as described in section 2.2.1 and primer was applied as in section 2.2.2.

The sealant was applied to approximately 12.7 cm (5 in.) at one end of the panel to a depth of $0.318 \pm 0.06 \text{ cm} (0.125 \pm 0.025 \text{ in.})$ and leveled, using a suitable jig. The screen was impregnated with sealant for approximately 12.7 cm (5 in.) on one end. The sealant-impregnated end of the screen was placed on the panel so that the loose,

unimpregnated end (aced the end of the panel free from sealant. The screen was smoothed down on the layer of scalant, taking care not to trap air under the screen. An additional 0.318 ± 0.06 cm $(0.125 \pm 0.025 \text{ in.})$ coating of sealant compound was applied over the impregnated screen. The scalant was cured in accordance with section 2.2.4.

Testing of Specimen . After exposure, two 2.5-cm (1-in) wide strips were prepared on each panel by cutting completely through the screen and sealant to the metal lengthwise along the panel and continuing completely along the unimpregnated screen. The loose + ud of each 2.5-cm (1-in) wide strip in turn was clamped in one jaw of a suitable recording tensile testing machine, and the adjacent end of the panel was fastened in the other jaw as shown in figure 8. Cuts through the sealant under the screen were made so that an initial separation of sealant from the metal panel was promoted. The screen was pulled at an angle of 180° from the panel and at a jaw separation rate of 5.1 cm/min (2 in./min).

Cuts in the sealant to the metal panel at the junction of separation were made at an angle of 45° toward the direction of separation at approximately 1.5-cm (0.6-in.) increments (approximately every 24 sec) on the left side of the panel as shown in figure 8 No cuts were required for 100% adhesive failure; however, any cobesive failures were trated as described. On the right side, except for the initial cut to promote separation, cuts were made only as necessary to prevent the sealant from peeling from the screen. All cuts extended completely across the strip being peeled and penetrated completely thro igh the sealant to the panel.

The percentage of cohesive paration was determined from the ratio of cohesive separation area to total cohesive and adhesive separation on both test areas. The cohesive strength was determined during cohesive tear. The average of the cohesive strength, as determined from an extensometer graph of the right-side pull, was recorded. Values recorded during cutting, or while load was being picked up after cutting, were not included in the average. Panels that were environmentally exposed were tested within 24 hr after removal from the exposure condition.

2.3.2 FUNCTIONAL TESTING

2.3.2.1 Picture Frame Shear

Details of Specimen Preparation.-The test panel was cleaned and primed as described. For a faying surface sealed panel, a layer of sealant was applied to one of the mating surfaces and the panel assembled while the sealant was still wet so that a continuous bead was extruded. The extruded sealant was then removed. Sealant was applied to fillet-sealed panels after assembly. A continuous fillet was applied around the periphery of the joint. Curing was in accordance with section 2.2.4.

Testing of Specimens-Leak testing was accomplished by placing a plexiglass box over the T-side of the panel (fig. 9), filling the box with water, and pulling a 13.8-kPa (2.0-psig) vacuum. Leakage was detected by observing the formation of bubbles. The test panel was then assembled in a picture frame fixture for application of structural loads. The frame was placed in an insulated conditioning chamber and loaded from the

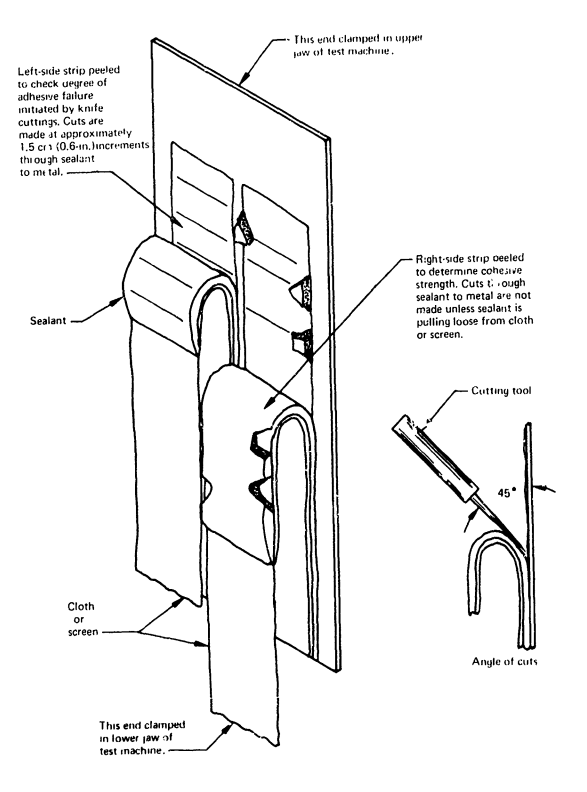
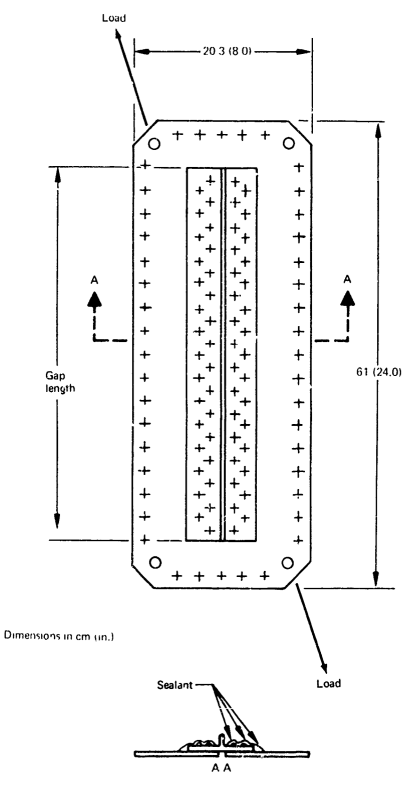


Figure 8.—Peel Strength Testing

:



ž

-

ł

こうちょう ちょうちょう シー・シー・シー・シート ちょうちょう ちょうちょう しょうちょう しょうしょう しょうしょう しょうしょう しょうしょう しょうしょう しょうしょう しょうしょう しょうしょう しょうしょう

Contraction of the

and the second state of the second second

NATION DE MILLINGS

1

ì

ヤモアンション

7

•

Figure 9.-Picture Frame Shear Specimen

corners. The test apparatus is shown in figure 10. A 22 680-kg (50 000-lb) tension load was applied to the frame 10 times at -46° C (-50° F), 10 times at room temperature, and 10 times at 232° C (450° F). A hydraulic cylinder was used for load application. The 22 680-kg (50 000-lb) load represents 100% of design limit load for the test panel.

After loading, the test panel was tested for leaks as described. The panel was then subjected to environmental aging. After various periods of aging, the panel was subjected to the loading and leak-testing conditions described.

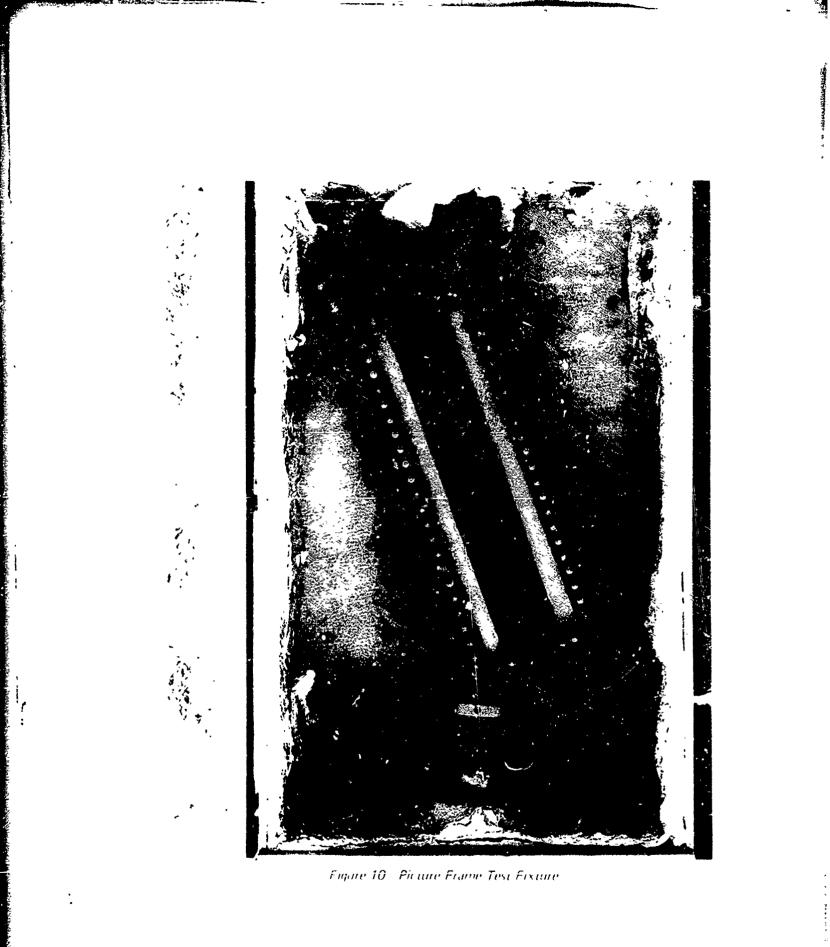
2.3.2.2 Flight Cycling Filler Deflections

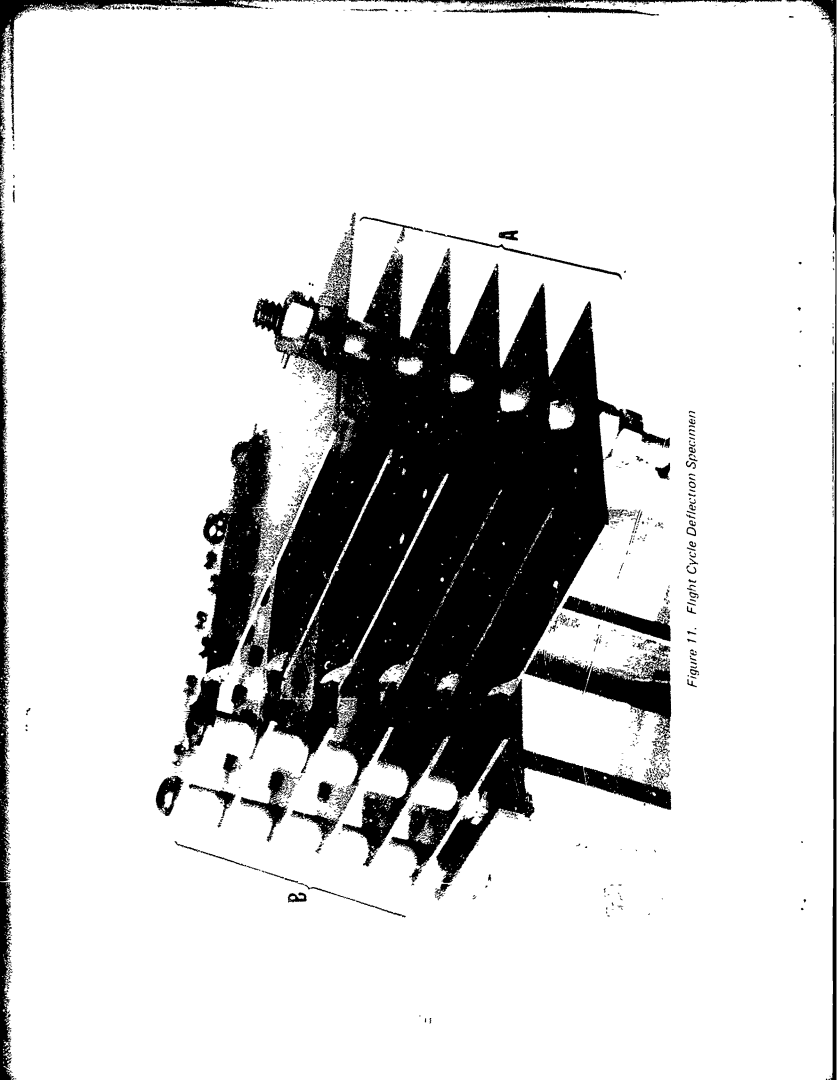
Specimens shown assembled in figure 11 were prepared from annealed 6AI-4V titanium alloy. They were cleaned and primed prior to fillet application according to sections 2.2.1 and 2.2.2. The titanium strips, B, were of constant length. Strips A overlapped by B were varied in length so that the gaps at the juncture of the A strips and B strips would vary with a constant end deflection of the strips B. Deflections were 0.008 cm (0.003 in.), 0.013 cm (0.005 in.), 0.020 cm (0.008 in.), 0.038 cm (0.015 in.), 0.051 cm (0.02 in.), and 0.076 cm (0.03 in.). The six specimens were mounted in a vertical row and fitted to the interior of the lid of the flight cycle apparatus. The end of each of the B strips was deflected at a rate of approximately 2 cycles per minute by means of a rod extending through the lid. The length of tear in each fillet throat was measured each time the flight cycle chamber was opened.

2.3.3 TANK TEST

and the first the second second and the second second in the second second second second second second second s

The tank test was performed by the Wichita Division of The Boeing Company. A complete description of tanks, tacilities, tests, and results is given in appendix A.





3.0 RESULTS

Four different lots of DC 77-028 were tested. Only one lot, 401117, was tested in both accelerated cycling and flight cycling. Specimens from lot 1222 were also in accelerated cycling. Specimens from lot 203117 were in flight cycling. The tank used material from lot 205180. All lots were manufactured at different times; the oldest was lot 1222 and the newest was lot 205180. Lot 401117 was a 226.79-kg (500-lb) production lot manufactured to demonstrate production capability and proof of ability to scale-up from laboratory size lots.

The only experimental sealant providing test results was the FCS-210 developed by Dow Corning with AFML funding.

3.1 EFFECTS OF ACCELERATED AND FLIGHT CYCLING

Figures 12 through 26 summarize changes in tensile strength, elongation, volume, weight, and adhesion after various periods of exposure to accelerated and flight cycling. Note that hours at the most severe condition, fuel vapor at 219° to 227° C (426° to 441° F), were used as a common base for comparison. For accelerated cycling, a factor of 1.2 may be used to calculate total exposure time to all fuel tank conditions. For flight cycling, a factor of 3 may be used. Each hour plotted is equivalent to one flight; therefore, 8217 flights were simulated. The plots also indicate the scatter in tensile and elongation data. Considering the inherent scatter, the difference between lots of material appears to be insignificant.

The sealant remained flexible and all but elongation at 232° C (450° F) and volume loss met target requirements after completing the exposures indicated. Elongation and volume loss were very close to the target requirements, and failure to meet them should not be construed as evidence that the sealant would not be serviceable.

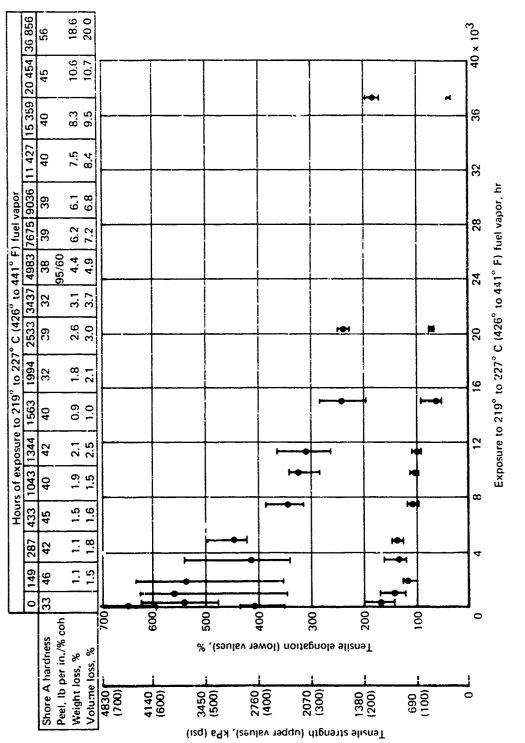
The lack of adhesion (peel test) data was due to invalid results, mainly adhesive failure between the sealant and embedded screen rather than between the sealant and titanium substrate.

It was hoped that a correlation of test results from accelerated cycling and flight cycling would be possible so that service life might be predictable using the shorter calendar time accelerated cycle. The indication is, however, that there is not a direct correlation and flight cycling is more severe than accelerated cycling. The comparison of weight and volume loss data in figure 27 dramatically depicts this.

3.2 FUNCTIONAL TESTING

3.2.1 PICTURE FRAME

The condition of the fillet-seeled picture frame test specimen is shown in figure 28 after 28 634 hr of 219° to 227° C (426° to 441° F) fuel vapor exposure in the accelerated cycle and 100 load cycles at each temperature. The sealant was tlexible and the panel

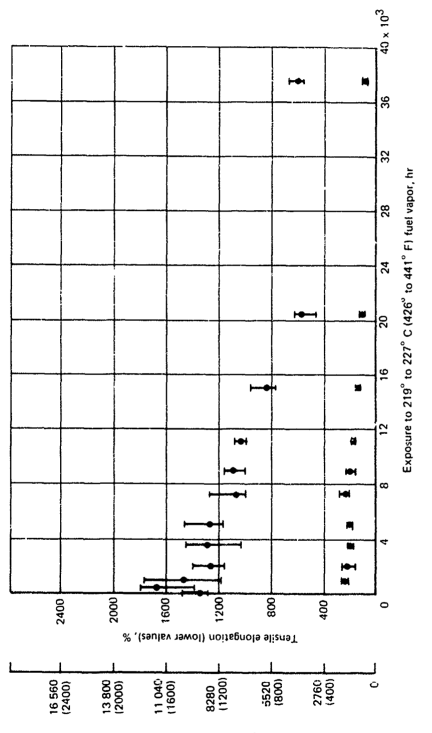


!



٩

.



THE ADDRESS OF

- ひかくがかく ひとう ふたんかい キャッシュー オット・ビン・シント かんしゅう ひかがた やまたれたい ひどうかん たかいちん ちゅう・ドン たんかん しゅう

ARK SKILL

i nati bishi kabara tali takukatin

Contraction of the second

ź

and a second and the second second and the second of the second second second second second second second second

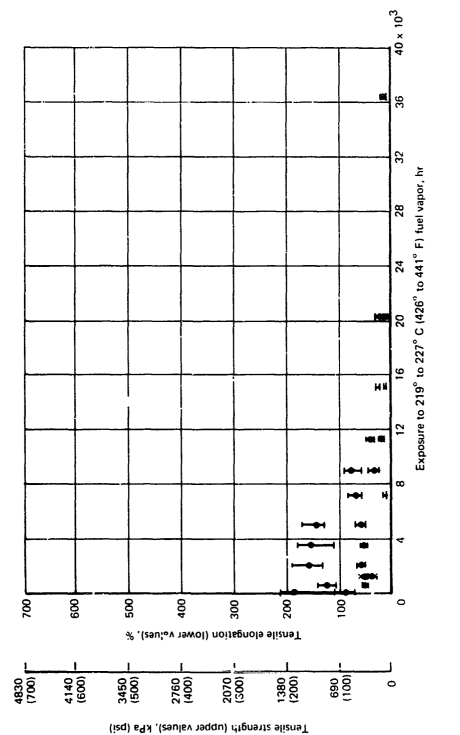
.

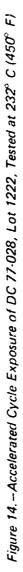
Figure 13.-Accelerated Cycle Exposure of DC 77-028 Lot 1222, Tested at -46° C (-50' F)

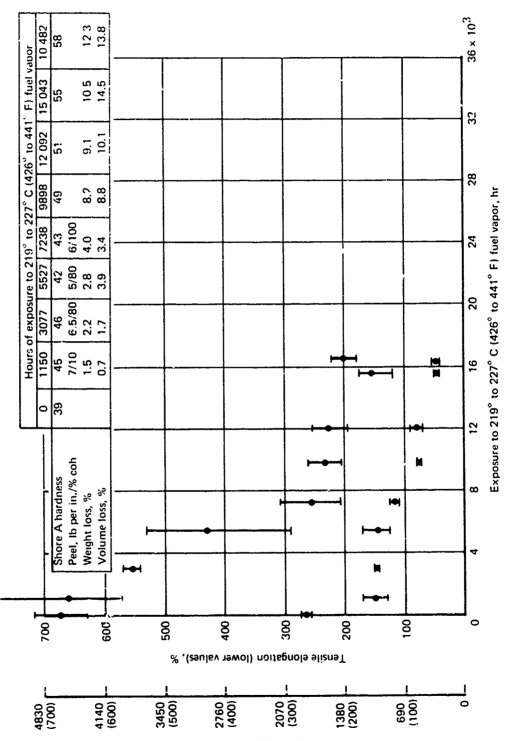
- - C.

. ч.

Tensile strength (upper values), dPa (psi)







÷.

149.1

Alexandre In State

10.2

ALTER AN ALTERNA

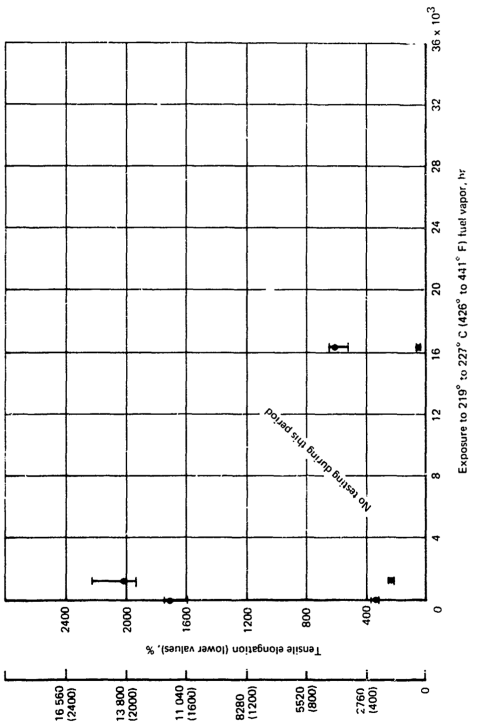
;

î

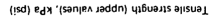
۰.

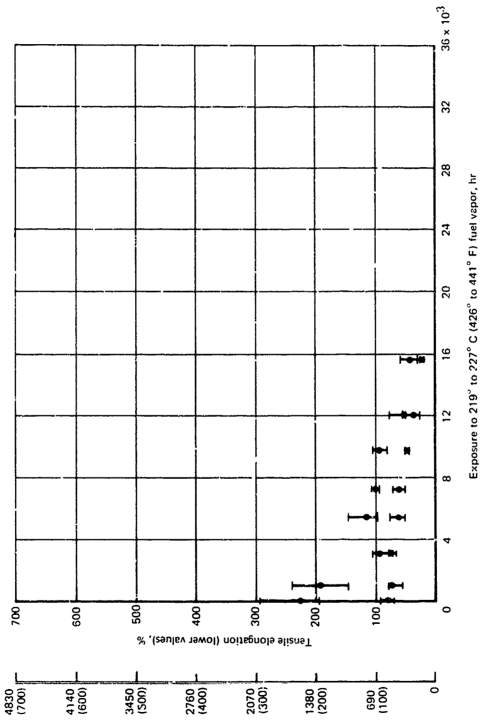
Figure 15.-Specimens From Wichita Tank, Lot 205180, Tested at Room Temperature

Tensile strength (upper values), kPa (psi)





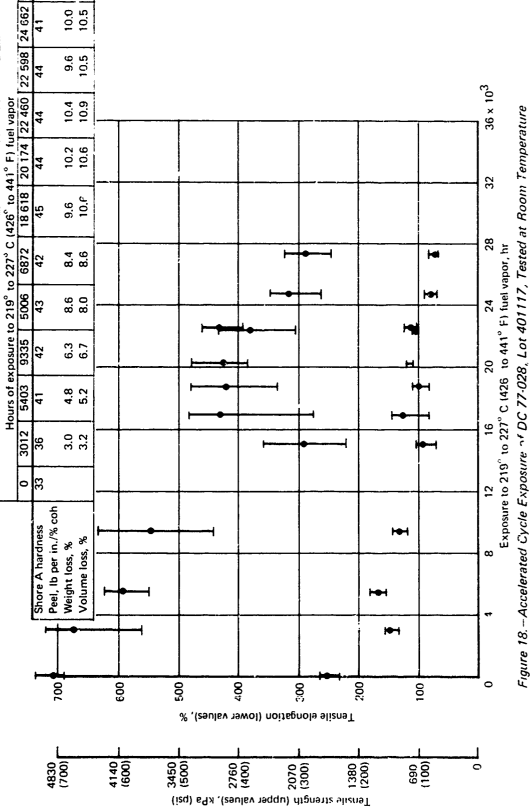




a state of the sta



Tensile strength (upper values), kPa (psi)



and the set of the set

الملكمة التلو فالقحم والألا أوكاله الاردا ولا

1

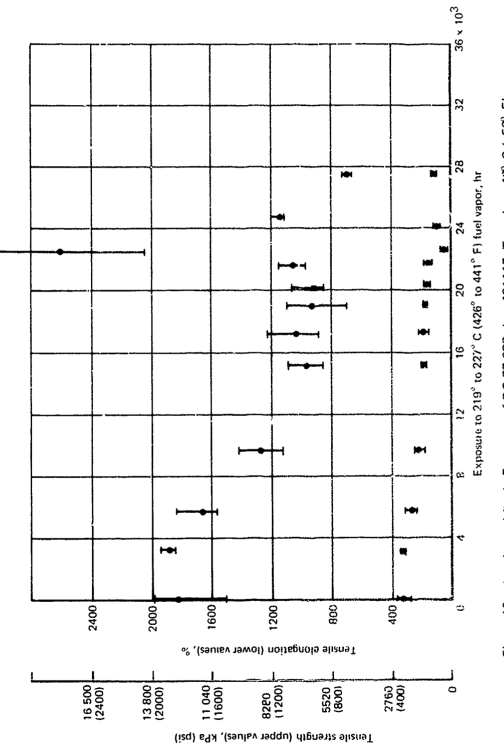
i

1. .

51

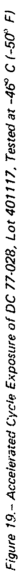
13.3 14 1

٢



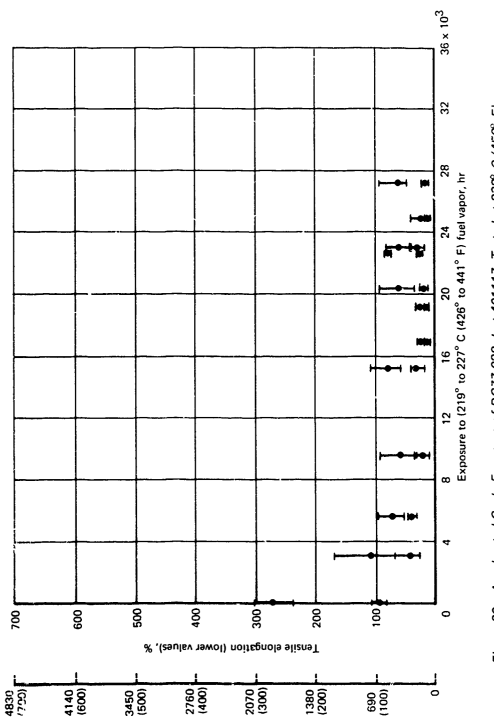
TINGS OF STREET

•



いたのかはおいればいました。そのなるなどのないので、「たいである」というないないです。

A STATE AND A STATE AN



第二、日本になるというないので、たちになっていたいである

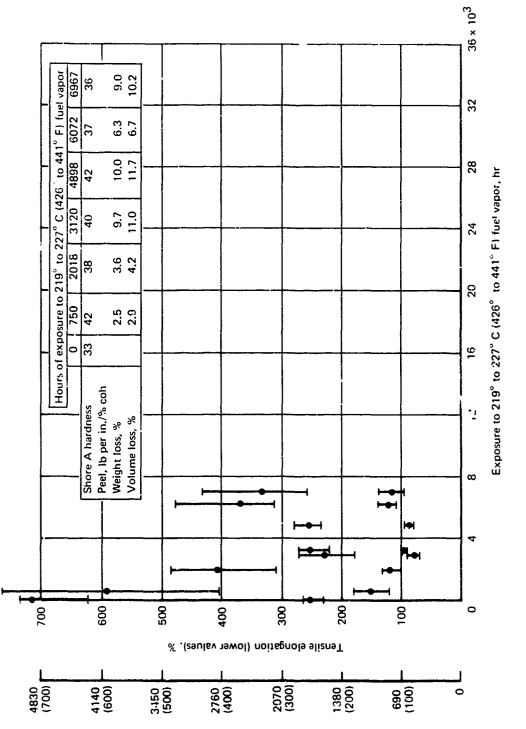
Statute Barth Samet

۽ *



30

Tensile strength (upper values), kPa (psi)



•

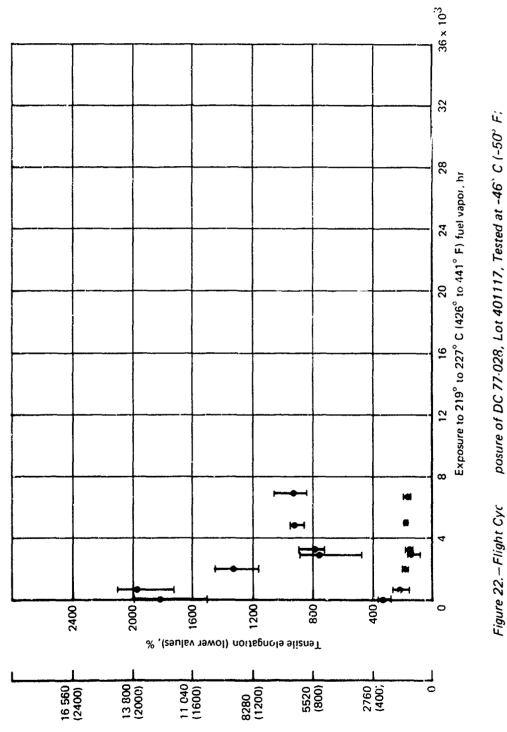
G



ちょうか ふまう いっき ちまう あんちかんなまっ 男子ないいろう ス・・・・チー・・バ

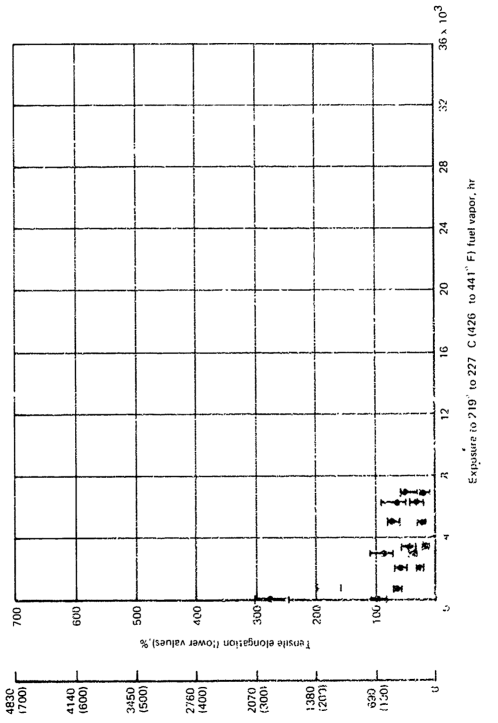
the state of the

Tensile strength (upper values), kPa (psi)



È,

Tensile strength (upper values), kPa (psi)



anno a suite de la constante d Carlo de la constante de la cons

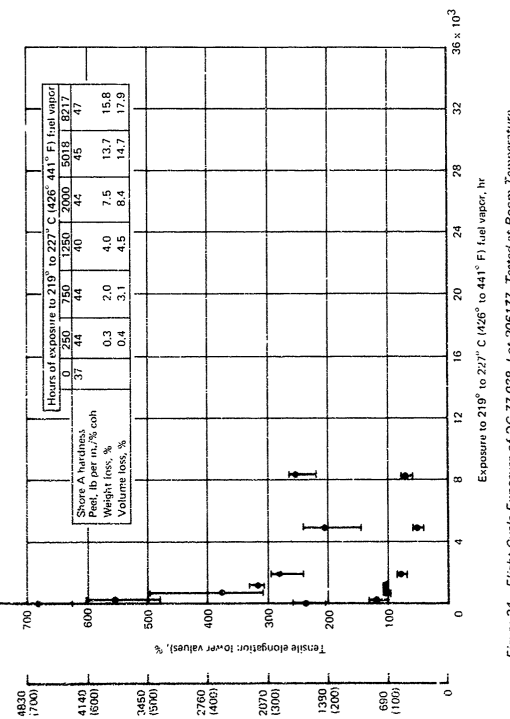
A State of the second second

تكتلونهم الأو

:



(isd) 644 (solues and (upper values). KPa (psi)



į

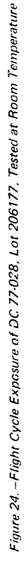
١, .

τ.

Stand Market was

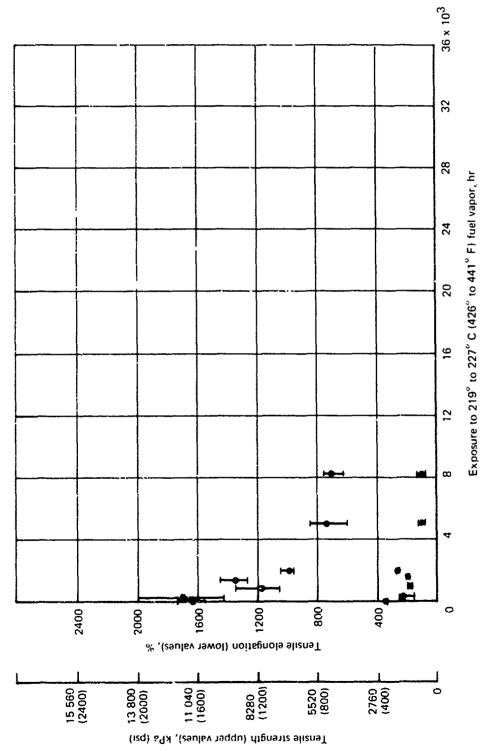
See 2 .

5



34

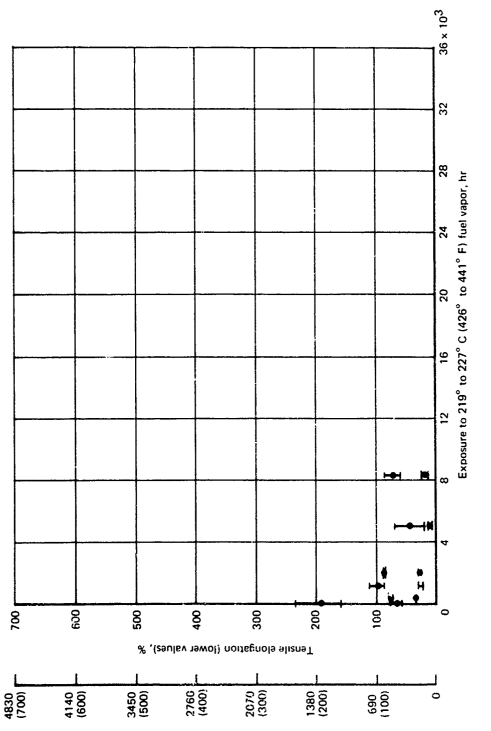
Tensile strength (upper values), kPa (psi)



į

ſ

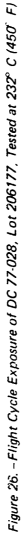


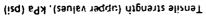


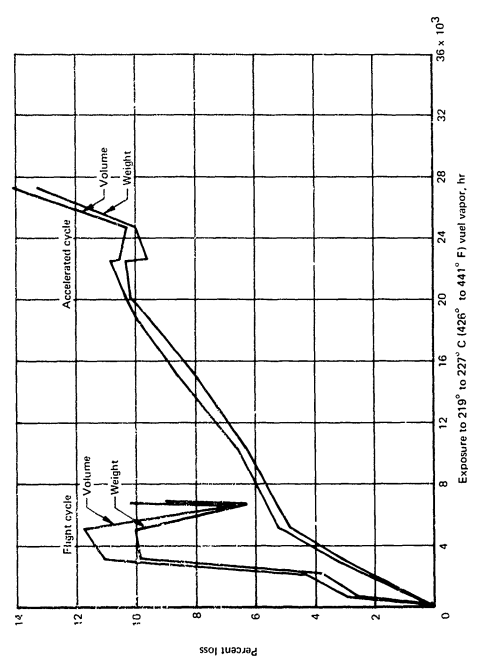
and a second second

م حد م محمد م

1







وتعفيه فالمستكرج فكالمستلك

ž

Figure 27.-Weight and Volume Loss for DC 77-028, Lot 401117

1. All - March States and and



Figure 28. - Fillet Seal of Picture Frame Panel After Exposure

remained essentially leak-free. The faying surface sealed panel was leaking slightly. Based on this and the fact that other tests showed that the faying surface scalant is not effective, we must assume that the test is unreliable.

3.2.2 FILLET DEFLECTION

After 6967 hr of 219° to 227° C (426° to 441° F) fuel vapor exposure and 836 040 load cycles in the flight cycle, only small cracks were apparent at the juncture of each fillet and the titanium overlap. By visual inspection, these cracks did not show evidence of growing after they were noted initially.

3.2.3 TANK TESTING

1

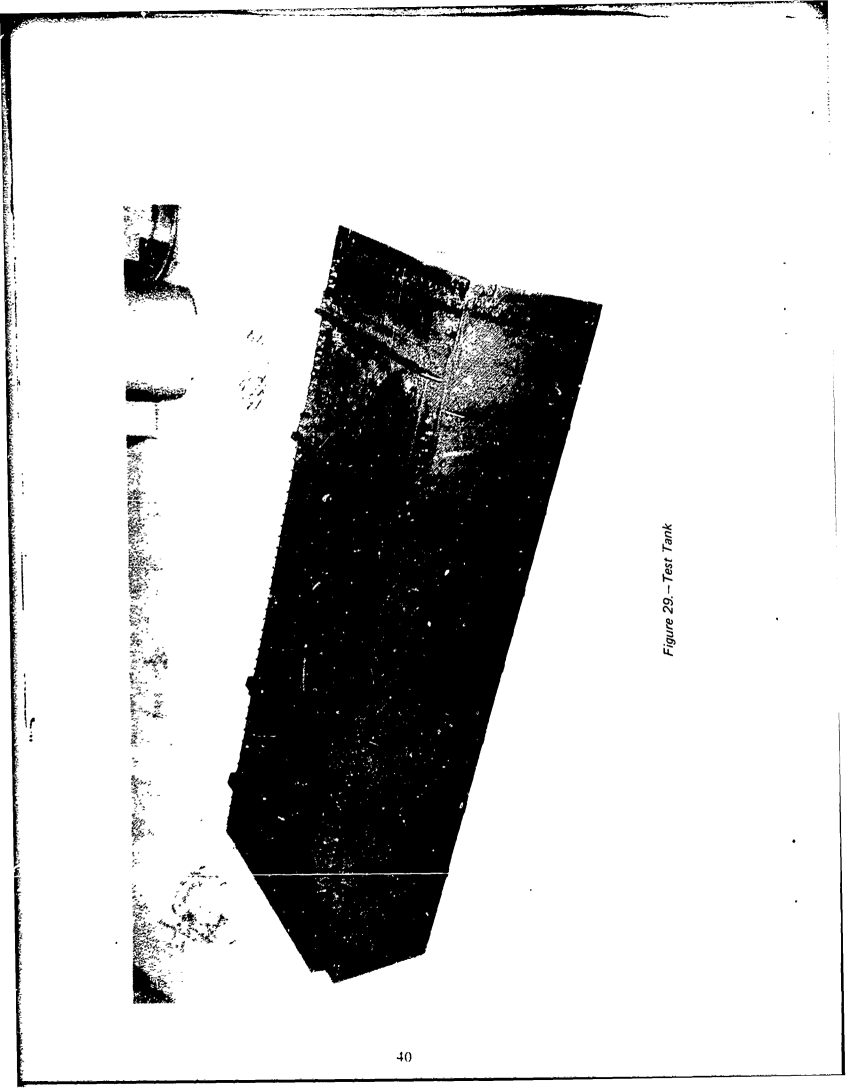
The tank test is considered to be the most informative in regard to sealant performance in an airplane structure. The tank is shown in figure 29 after 6000 load cycles at room temperature, 4500 cycles at 232° C (450° F), 4330 cycles at -46° C (-50° F), and exposure to fuel and fuel vapor for the temperature and hours indicated in table 1. The tank was inspected and tested for leaks once every 6 months. The leaks varied each time in number and location. Detection of leaks was never possible without pressurization. For the final test, the tank was left to stand for about 1 hr with water in it and it did not leak. Approximately 13.8-kPa (2-psi) pressure was then applied, which caused two spots to leak very slightly. After a short period of time, oozing was noted at one other location. Pressure of 34.5 kPa (5 psi) was then applied and the leaks increased in amount and number. When the pressure was removed, the tank continued to leak. It was then drained.

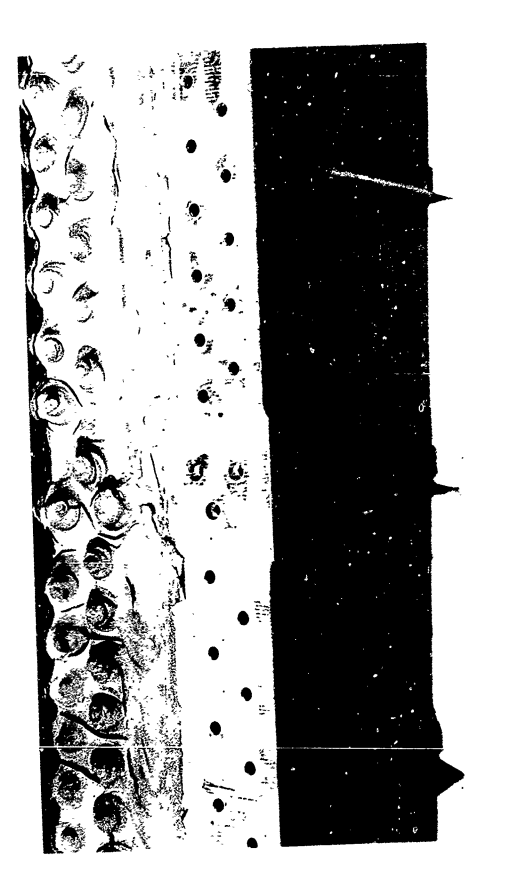
Temperature, °C (°F)	Hours	
Liq	luid	
To 59 (139)	676	
60 (140) to 65 (150)	2 133.5	
Over 65 (150)	113.2	
Below 60 (140)	521.5	
Vap	or	
60 (140) to 204 (400)	1 794.8	
205 (401) to 218 (425)	470	
213 (415) to 227 (441)	78.5	
219 (426) to 227 (441)	16 481.5	
Over 227 (441)*	65	

Table 1.-Exposure Times for the Test Tank

*Included in hours at 219" to 227° C (426° to 441° F); highest was 238° C (460° F).

It had been noted previously that cracks in the sealant, such as those shown in figure 30, were more prevalent when the sealant fillet was thick, so a small section was





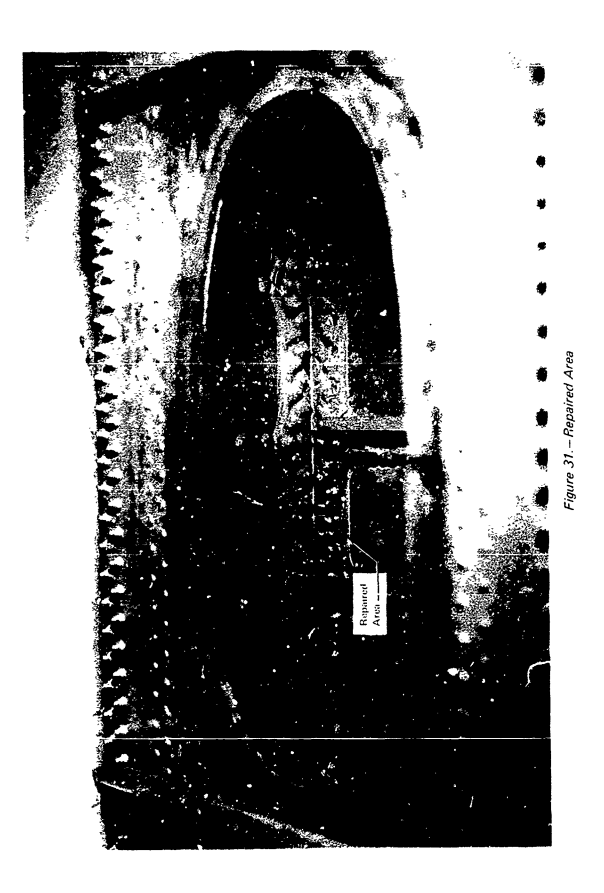
ţ

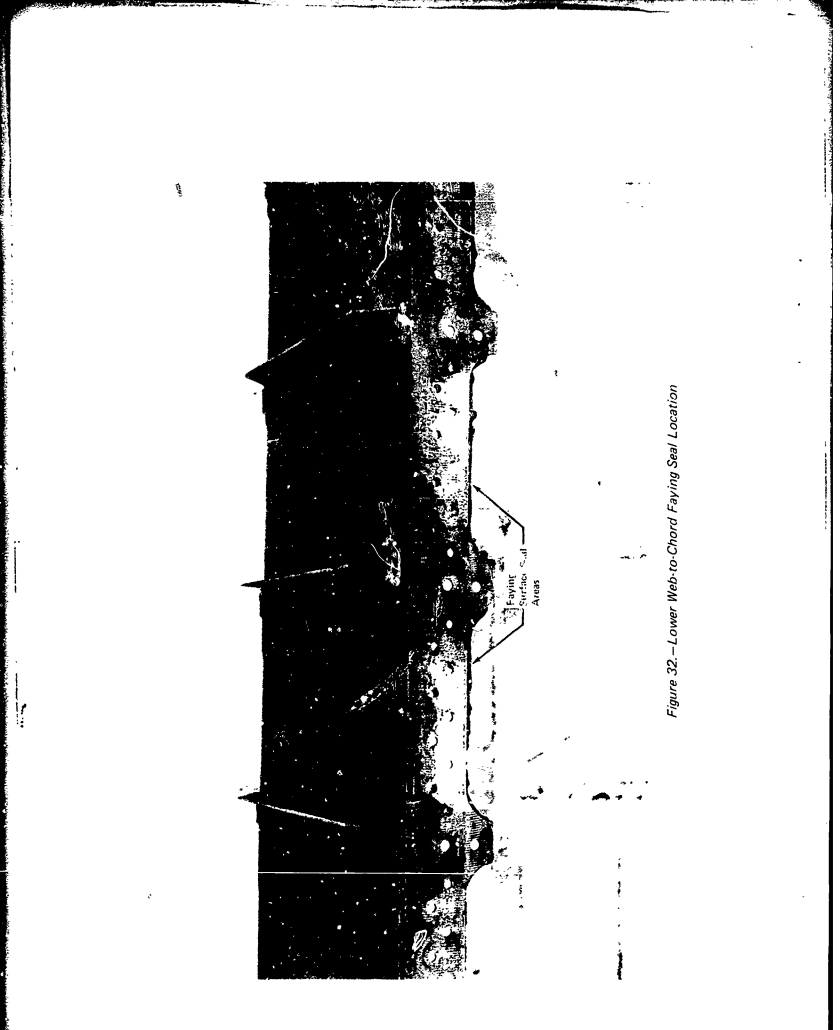
Figure 30.--Sealant Condition After Exposure

repaired using small fillets. The section was repaired after 7238 hr of 219° to 227° C (426° to 441° F) fuel vapor exposure in February 1973 and had no splits and no leaks. Adhesion was retained to the old sealent as shown in figure 31.

The top and bottom skins were removed. Adhesion of the sealant to the fasteners and structure was surprisingly good, and the sealant fillets failed cohesively when the structural components were separated, as can be seen in the pictures. Areas where the faying surface isolation seals had been were evident (figs. 32 and 33), but there was little sealant to be seen. What was there was easily scored with the blunt edge of a fingernail. Condition of the Teflon rings in the self-sealing fasteners (which were unsuccessful from the beginning) varied from no change through a crumbly powder to nonexistent. The sealant on the top skin had separated from the seal plane, had numerous large splits, and was absent in some places.

Each inspection revealed leaking corners which were usually repaired before returning the tank to environmental exposure. The injection sealant was inadequate in that it expanded, gradually extruded out of the groove, and ruptured the adjacent fillet. Modification of the DC 77-028 fillet sealant to produce DC 77-066 injection sealant was demonstrated to be partially effective. At one time, corner repairs were made using DC 77-028; the corners started leaking badly immediately when exposed to temperature. The injection seal is shown in figure 34.





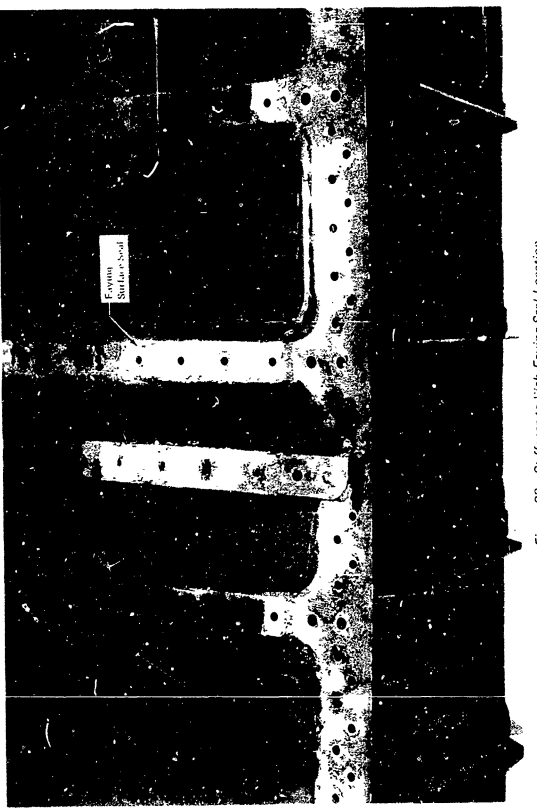
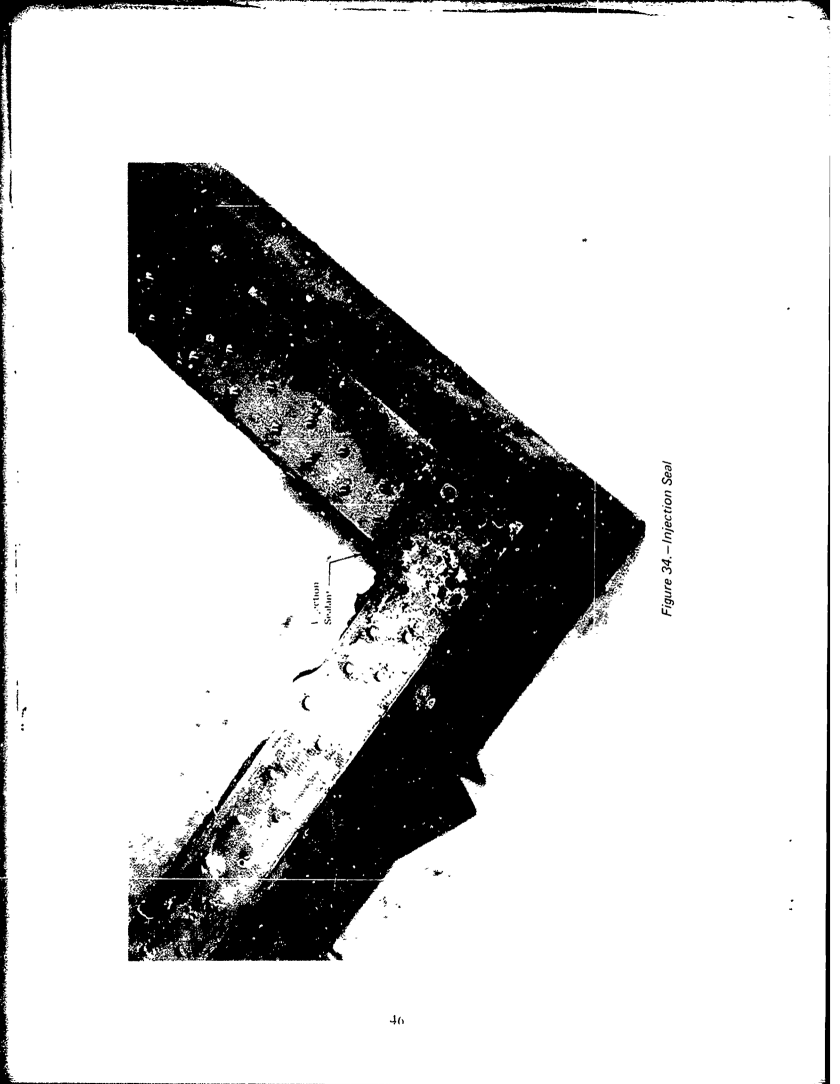


Figure 33.--Stiffener-to-Web Faying Seal Location



4.0 BACKUP SEALANT

4.1 HYDROFLUOROCARBON

The Products Research and Chemical Corporation (PRC) was given a contract to develop a sealant based on fluorocarbons and having a high solids content. Existing sealants based on fluorocarbons are normally supplied with a maximum of 40% solids by volume, requiring them to be applied in thin cross sections and resulting in shrinkage on drying and curing. PRC was asked to develop one having a solids content of 80% by volume; one having 70% was produced, which is considered a significant advancement. Solvent content was further reduced to less than 10% by volume by pre-extruding into a tape which was tacky enough to adhere to titanium. After a heat cure, adhesion was retained; however, when ar attempt was made to cure a sheet of sufficient thickness to provide test specimens, bubbling took place and destroyed the sheet. The progress made was encouraging and further investigation and development are warranted, specifically in adhesion, thermal stability, and stress corrosion. A complete report of the development is attached as appendix P.

4.2 HYBRID FLUOROCARBON-FLUOROSILICONE

The Air Force Materials Laboratory funded the development of hybrid fluorocarbon-fluorosilicones by the Dow Corning Corporation. One of the most promising of these, DC 77-108 (FCS 210), was exposed to accelerated cycling and tested periodically. Test results are shown in table 2.

Property	Hours of 219	Hours of 219 ' to 227 C (426 to 441' F) fuel vapor exposure				
rioperty	0	2520	9938	15 826		
Tensile strength, kPa (psi) -46° C (~50 [°] F) Room temperature 232° C (450° F)	24 019 (3486) 7 406 (1074) 875 (127)	17 749 (2576) 7 917 (1149) 1 523 (221)	19 120 (2775) 7 237 (1059) 1 054 (153)	8342 (1209) 4885 (708) 849 (123)		
Elongation, % -46 C (-50" F) Room temperature 232" C (450" F)	8 585 64	1 7 640 111	7 ∠10 44	8 190 26		
Shore A durometer (hardness) Weight loss, % Volume loss, %	34	45 7.0 6.8	76 7.4 7.5	46 7.4 7.3		

Table 2. - DC 77-108 Test Results

5.0 CONCLUSIONS

- 1. Confidence was develope.' that the Dow Corning DC 77-028 (DC 94-529) flucrosilicone sealant system would perform for 50 000 flight-hours as an integral fuel tank fillet sealant in a commercial supersonic airplane. This confidence was established by the completion of the following evaluations:
 - a. More than 36 000-hr accelerated life cycle test in 219° to 227°C (426° to 441°F) fuel vapor
 - b. More than 8000 flight cycles test (24 000 hr) in 219° to 227° C (426° to 441° F) fuel vapor under ±96.6-kPa (±14-psi) pressure variance
 - c. More than 16 000 hr of test as sealant in a simulated SST fuel tank under accelerated life cycle conditions in 219° to 227° C (426° to 441° F) fuel vapor
 - d. Torsional load cycle tests of the sealed tank (item c) of 4330 at -46° C (-50° F), 6000 at room temperature, and 4500 at 232° C (459° F)
 - e. Demonstration of fillet sealant repairability during tank tests

and a state of the state of the

- f. Demonstration of lot to-lot reproducibility of production size lots of DC 77-028
- 2. Neither the faying surface sealant (DC 77-053) nor the injection sealant (DC 77-066) derivatives of the DC 77-028 fluorosilicone system were satisfactory; however, the DC 77-066 was superior to DC 77-028 as an injection sealant.
- 3. The backup fluorocarbon and hybrid fluorocarbon-fluorosilicone sealant systems show promise but do not have the low-temperature ductility capabilities demonstrated by the fluorosilicone sealant.
- 4. Flight cycle testing provides more critical evaluation of basic sealant properties for a given total test time than accelerated life cycle testing. Testing of the sealant in its intended application in a simulated fuel tank is most informative from a practical standpoint of sealant application, behavior, and repairability.

6.0 RECOMMENDATIONS

Further development of a faying surface and injection sealant should be pursued.

and the second secon

ちんてい ちかい どうちん かいちん ひろう ストライン

したいまましたとうといういろうないのである

Flight testing of the DC 77-028 system in a supersonic aircraft is the next logical step in investigation of its functionality and should be initiated.

Attempts should be made to develop fuel tank sealants from polymers other than fluorosilicones, especially if aircraft designed to fly in excess of Mach 2.7 are contemplated. These sealants should be tested in the same way as DC 77-028 for comparison purposes.

APPENDIX A



CODE IDENT. NO. 81205

NUMBER	<u>D3-8297</u>	REV LTR _F
INITIAL	RELEASE DATE	APR I 3 1970
	EVALUATION	OF THE PROPOSED SST FUEL TANK
	SEALANT IN	SMALL SIMULATED FUEL TANKS

FOR LIMITATIONS IMPOSED ON THE USE OF THE INFORMATION CONTAINED IN THIS DOCUMENT AND ON THE DISTRIBUTION OF THIS DOCUMENT, SEE LIMITATIONS SHEET.

MODEL	2707	_CONTRACT	····

			1	10 1	
PREPARED BY		duli	lid	ditor	3-31-70
SUPERVISED BY		Phar	1 -	- ()	on 17.1-70
	/	PI	B	hi a	·
APPROVED BY	$\frac{\zeta}{1}$	$4 \sqrt{4}$	L'a	emer	3 7 7 7
APPROVED BY	(]	aler G	·	King.	2 4/3/10
				δ	· · ·
	\Box				

NO. D3-8297 PAGE 1 OF 114 BUEING

etti≪3D

TOM.

1 7 11

E-3039 R4

LIMITATIONS

INFORMATION CONTAINED HEREIN IS PROPRIETARY TO THE BOEING COMPANY AND SHALL NOT, WITHOUT PERMISSION OF THE BOEING COMPANY, BE REPRODUCED, DISCLOSED, OR USED BY AKYONE OTHER THAN AGENCIES OF THE GOVERNMENT ACTING PURSUANT TO CONTRACTUAL OR STATUTORY AUTHORIZATION.

THIS DOCUMENT IS CONTROLLED BY MATERIALS TECHNOLOGY UNIT ORGANIZATION 3-7584

ALL REVISIONS TO THIS DOCUMENT SHALL BE APPROVED BY THE ABOVE NOTED ORGANIZATION PRIOR TO RELEASE.

BOEINO	NO. D3-8297
SECT	PAGE 2

REV LTR: A

E-3043 R1

ABSTRACT

In order to determine the fuel tank sealing capabilities of the sealant (Dow Corning 77-028) proposed for the fuel tanks of the Model 2707 supersonic transport prototypes, two small titanium test tanks were initially fabricated and sealed at Boeing-Wichita. These two tanks were subjected to long term simulated fuel tank environments and were periodically subjected to cyclic structural loading at three temperatures, followed by pressure testing for leakage. A third test tank, which more closely simulated the SST fuel tank structure, was subsequently fabricated and sealed by CAG. This tank was exposed to similar environmental conditioning and structural loading.

Testing of tank numbers 1 and 2 is described in Section 2. Testing of tank no. 2 was terminated after 524 hours of fuel vapor exposure at 415 - 441°F (4 weekly environmental conditioning cycles). Testing of tank no. 1 continued to 3862 hours of fuel vapor exposure (31 environmental conditioning cycles).

Testing of tank number 3 is described in Section 3. This tank was exposed to a total of 16,482 hours of fuel vapor at $426 - 441^{\circ}F$ or a total of 22,269 hours at all conditions (liquid fuel, ambient to 205°F, and vapor fuel, ambient to 460°F). Total number of weekly environmental conditioning cycles was 129. In addition the tank was periodically load cycled for an accumulated 6000 cycles at ambient temperatures, 4500 cycles at 426 - 441°F, and 4330 cycles at -40 to -50°F.

Early in the environmental conditioning of all three tanks the sealant exhibited numerous cracks and areas of loss of adhesion resulting in significant leakage. This document describes in detail the environmental conditioning and structural loading used, and the condition of the sealant and the observed leakage of the tanks after each increment of specified testing. However, all conclusions concerning the significance of the testing and the anticipated performance of the sealant will be presented elsewhere by the Commercial Airplane Group, and is therefore beyond the scope of Wichita Division responsibility.

BOEING	NO.	D3-8297
SECT	PAGE	3

REVLTR: E

E-3033 R1

TABLE OF CONTENTS

			Page
	TITL	E PAGE	1
	LINI	TATIONS	2
	ABST	RACT	З
	TABL	E OF CONTENTS	4
	LIST	OF ILLUSTRATIONS	6
	FORE	WORD	8
1.	INTR	ODUCTION	9
2.	TEST	ING OF TANK NUMBERS 1 AND 2	10
	2.1	Fabrication and Sealing	10
	2.2	Preparation and Testing Of Physical Property Specimens	17
	2.3	Environmental Conditioning	25
	2.4	Dynamic Load Cycling	29
		2.4.1 Description Of Dynamic Test Device 2.4.2 Determination Of Torsional Load Levels 2.4.3 General Test Procedure For Dynamic	29 30
		Cycling and Leak Testing Of Tanks	33
	2.5	Evaluation Of Tanks After Four Cycles Of Environmental Conditioning	34
	2.6	Repairs and Continuation Of Testing Of Tank No. 1	38
		 2.6.1 Repairs Of Tank 2.6.2 Dynamic Load Cycling After Repairs 2.6.3 Evaluation Of Tank No. 1 After 12 Cycles Of Environmental Conditioning 	38 39 41
	2.7	Evaluation Of Sealant Repair Procedures In Tank No. 1	46

BOEINO	NO.	D3-8297
SECT	PAGE	4

A +

.

٠

.

REVLTR: A

E-3033 R1

- 94

7

l

1.1.1

مون المواتين المحالية. المانية بالمعالية المحالية المحالية المحالية المحالية المحالية المحالية المحالية المحالي محالية المحالية المحالية من محالية المحالية المحالية المحالية المحالية المحالية المحالية المحالية المحالية المح

a substanting a state with the set of a state of a

1.2.1.1.L.

sa di baba na Minda ndanya ning sulat na na na na na

Ê !

ţ

and a second second

1.4

TABLE OF CONTENTS (Continued)

,		INDEE OF CONTENTS (CONTINUED)	Page	
3.	TEST	ING OF TANK NO. 3	50	
	3.1	General Description	50	
	3.2	Initial Preparation and Testing Of Tank	51	
	3.3	Dynamic Load Cycling	52	
		3.3.1 General Test Procedure	52	
	3.4	3.3.2 Initial Load Cycling Prior To Any Environmental Conditioning Preparation and Testing Of Physical Property	52	
		Specimens For Tank No. 3	56	
	3.5	Environmental Conditioning Of Tank No. 3	57	
	3.6	Dynamic Load Cycling Of Tank No. 3 After 22 Cyles Of Environmental Conditioning	57.2	
	3.7	Dynamic Load Cycling Of Tank No. 3 After 39 Environmental Conditioning Cycles (August 1972) .	57.4	
	3.8	Repair Of Tank No. 3 After 46 Environmental Conditioning Cycles (November 1972)	57.6	
	3.9	Dynamic Load Cycling Of Tank No. 3 After 54 Environmental Conditioning Cycles (February 1973)	57.7	
	3.10	Dynamic Load Cycling Of Tank No. 3 After 74 Environmental Conditioning Cycles (August 1973) .	57.8	
	3.11	Dynamic Load Cycling Of Tank No. 3 After 92 Environmental Conditioning Cycles (January 1974)	57.9	
	3.12	Dynamic Load Cycling Of Tank No. 3 After 117 Environmental Conditioning Cycles (August 1974)	57.10	Ε
•	3.13	Dynamic Load Cycling Of Tank No. 3 After 129 Environmental Conditioning Cycles (December 1974) RENCES	57.11 58	Ε
4.		RENCES	58 84	
	AFFL	NDIN A - Leakaye lests - rank numbers i and 2	54	
	APPE	NDIX B - Leakage Tests - Tank Number 3	98	
	APPE	NDIX C - Measurement Of Web Shear Wrinkles		
		Gf Tank Number 3	105	
	REVI	SIONS	113	
	CHAN	GE RECORD PAGE	114	
		BOEING NO. D3-82	97	
RE.V	LTR:E	SECT PAGE 5		

REVLTR:E

an - were date in the location of the second strategy of the second states of the second states in the second s

and and some as the second stranger with a set of the second stranger of the second stranger with the second st

the second s

St. North

FIGURES	LIST OF ILLUSTRATIONS	Page
1	Typical Cross-Sectional View Of Fillet Sealed Tank No. 1	59
2	Partial Assembly and Sealing Of Tank No. 1	60
3	Partial Assembly and Sealing Of Tank No. 1 - Upper End View	61
4	Partial Assembly and Sealing Of Tank No. 1 - Lower End View	62
5	Partial Assembly and Sealing Of Tank No. 2	63
6	Completed Test Tanks	64
7	Peel Strength Testing	65
8	Test Specimens Ready For Installation In One Test Tank	66
9	Flow Chart - Environmental Conditioning Of Test Tanks	67
10	Environmental Conditioning Test Equipment	68
11	Environmental Chamber With Side Heaters Installed .	69
12	Dynamic Test Device (With Insulating Chamber For Hot Air Circulation)	70
13	Dynamic Test Device (With Circulating Cold Fuel)	71
14	Cracks In Fillets At Base Of Fasteners	72
15	Cracked Fillets At Center Rib (Tank No. 1)	73
16	Cracked Fillets At Center Rib (Tank No. 2)	74
17	Fillet At Center Rib (Tank No. 2) Showing Crack and Loss Of Adhesion	75
18	Area of Tank No. 3 Repaired With Small Fillets in February 1974	75.1 E

BOEING	NO.	D3-8297
SECT	PAGE	6

REVLTR: E

E-3013 R1

States & March

i

ALL ALL ALL ALL

たちのうちょうとうないできたのできたのできたのできたのでありますの

Ly, Grobert & S. S. S. S.

ł

TABLES	LIST OF ILLUSTRATIONS (Continued)	Page
1	Physical Property Specimen Test Results For Tank Numbers 1 and 2	76
2	Environmental Conditioning Of Tank No. 1	77-78
3	Environmental Conditioning Of Tank No. 2	79
4	Leakage Of Tank Numbers 1 and 2 After Four Cycles Of Environmental Conditioning	80
5	Summary Of Leakage Of Tank No. 1 In Primary Test Areas	81
6	Physical Property Specimen Test Results For Tank No. 3	82
7	Environmental Conditioning Of Tank No. 3	83
8	Corner Leakage History of Tank No. 3	83.10 E

BOEING	NO.	D3-8297
SECT	PAGE	7

,

FOREWORD

This report was prepared by the Materials Technology Unit of The Boeing Company, Wichita Division. Work was initiated under Engineering Work Authorization Number 36529221, titled "Environmental and Functional Testing Of SST Fuel Tank Sealant", dated August 4, 1969 (written by the Commercial Airplane Group, Structures Technology-Materials-SST).

On March 25, 1971 this program was cancelled when the government denied additional funds for SST prototype construction.

Work was subsequently resumed in August 1971 under Commercial Group IDWA No. 630600 (EWA No. 964020), and Wichita EWI No. 964020 funded by the Department of Transportation.

Funding was continued in April 1972 under Commercial Airplane Group IDWA No. 630600 (EWA No. 03866), and Wichita EWI No. 989001.

In December 1973 the final portion of the program was authorized under Commercial Airplane Group IDWA No. B30600 and Wichita EWI No. 989001-1.

The program was completed in March 1975.

BOEING	NO.	D3-8297
SECT	PAGE	8

REVLTR: E

E-3033 R1

58

♥ E

1. INTRODUCTION

The most promising sealant available to date for sealing the fuel tanks of the Model 2707 Supersonic Transport is Dow Corning 77-028, a two-part experimental fluorosilicone, which was developed specifically for this purpose. This sealant must function for at least 50,000 hours in the presence of liquid hydrocarbon fuel up to 150°F and fuel vapor and nitrogen up to 441°F.

The purpose of the work conducted herein is to verify the sealing capabilities of this sealant by environmental and structural lord testing of small sealed titanium test tanks.

To accomplish the above objective two small titanium tanks were initially fabricated and sealed. These tanks simulate typical skin and stiffener type integral fuel tank structure. One tank was fillet sealed only and the other was fillet sealed and faying surface sealed. These tanks were exposed to long term fuel tank environments, and were periodically subjected to structural dynamic torsional loading, followed by pressure testing, to determine the capability of the sealant to retain fuel at the environmental temperature extremes of the SST fuel system.

A third tank was subsequently built by CAG and tested by Wichita. This tank more closely simulated the SST fuel tank structure than did the first two tanks.

59

BOEINO	NO.	D3-8297
SECT	PAGE	9

REVLTR:

E-3033 R1

Laboratory test specimens were prepared and installed in each tank. These specimens were environmentally conditioned in the tanks and were withdrawn periodically and evaluated to measure the change in physical properties.

2. TESTING OF TANK NUMBERS 1 AND 2

2.1 Fabrication and Sealing

The test tank design used herein was originally developed 1. by Boeing-Wichita under a contract with the Air Force Materials Laboratory (AFML). Reference 1 is the fina? report under this contract. The detail parts for two test tanks, which were originally fabricated for this contract but not used, were returned to Boeing-Wichita by AFML and used to build the two test tanks described herein. Optimum design of the test tanks required that they be large enough to simulate actual fuel tank structure and small enough to be economical to evaluate. The tank size chosen was approximately 9 by 14 by 50 inches. The tank design is a semi-monocoque box beam of 6A1-4V titanium sheet skins and webs, and extruded titanium chords and stiffeners. The end plates and test jig attach fittings are 4130 steel. Reference 2 is the Boeing drawing for the tank details. This type and size of tank is believed to

BOEINO	NO.	D3-8297
SECT	PAGE	10

REVLTR: A

E-1011 R1

be the most economical and the least complex construction with which significant and reliable results may be obtained.

- Two tanks were fabricated and sealed. Tank Number 1 was fillet sealed only. Tank Number 2 was completely faying surface sealed as well as fillet sealed.
- 3. Tank detail parts were vapor degreased, followed by nitrichydrofluoric acid etching. Steel parts were descaled in nitric acid, sandblasted, and spray painted with Sermetal W (BMS 14-4) per BAC 5840 for corrosion protection.
- 4. The tanks were assembled using titanium bolts (BACB30HY6) and collars (BACC30AE6W) from Voi-Shan manufacturing Company, except for the end plates and access doors where cadmium plated steel bolts were used.
- 5. In general the sealing was conducted using BAC 5504 procedures as a guide. Dow Corning 77-028 sealant was hand-mixed as described in Section 2.2, Item 2. Some sealant was used immediately after mixing, and some was frozen at -40°F and used within 4 days. The first lot of sealant received (Lot No. 411147, dated 19 November 1969) wruld not cure properly, even at 300°F. This lot was replaced by Lot No. 1114, manufactured 3 December 1969, which was used for all sealing of both tanks and for the preparation of all Section 3. laboratory test specimens.

FIVLTR: A

_	BOEINO	NO.	D3-8297
ſ	SECT	PAG	. 11

E-3013 R1

- 6. All structure to be sealed was solvent cleaned with BNS 11-7 cleaner prior to applying primer. Initially some methylethylketone was used instead of BMS 11-7. After solvent cleaning, a thin coat of Dow Corning 77-037 primer (Reference No. E-23-233, dated December 1969) was applied with a gauze pad moistened with primer. The primer was allowed to dry a minimum of 45 minutes at room temperature prior to applying sealant.
- 7. Freshly mixed (or freshly thawed) sealant was applied using a Semco sealant gun, cartridge and formed nozzle to apply a fillet in one application to the dimensions specified in BAC 5504. Faying surface seals were applied by injecting the sealant on the structure and spreading it out to a thin continuous coating with a spatula or fairing tool.
- 8. Both access doors were installed with faying surface seals. Tank No. 1 (fillet sealed only) was sealed as shown typically in Figure 1. Block-off faying surface seals were installed (as shown typically in Figure 5) to aid in isolating any fillet leaks which may be evident in the future. Figures 2, 3 and 4 show details of Tank No. 1 during one stage in the assembly and

DEINCE NO. 03-8297 PAGE 12

REVLTR: A

E-3033 R1

sealing. The total weight of the sealant fillets in this tank was 2.79 pounds, based on weighing the sealant applied.

- 9. Tank No. 2 was committely faying surface sealed and fillet sealed. Figure 5 shows details of this tank prior to installation of the spar web.
- 10. The sealant in each tank received several conditions of cure depending upon location, since the tanks were progressively fabricated and sealed in several steps. Interim cures were generally 1 to 2 hours at 160°F. Upon completion of each tank a minimum cure of 2 hours at 160°F plus 1 hour at 300°F was used. The thermocouples and laboratory test specimens were then installed in each tank (see figure 6) and the access doors installed using steel bolts in nutplates through faying surface seals. The tanks then received one additional cure of 2 hours at 160°F and 1 hour at 300°F.

BGEINO	NO.	D3-	8297
SECT	PAGE		3

FEVLTR A

E-3033 M1

- 11. Completed tanks were leak tested by applying 5 10 psi air pressure internally and immersing tanks in water or applying bubble (soap) solution externally. Both tanks exhibited several (up to 9) fastener leaks which were repaired by applying additional sealant over the fasteners on the tank interior. The completed tanks are shown in Figure 6 prior to installation of the access doors.
- 12. After the sealing of the tanks had been completed, Dow Corning became concerned that sealant lot No.. 1114, used to seal both tanks, way not be representative of their best sealant. It was requested that we check the thermal stability of this lot by exposing standard tensile strength and elongation specimens to dry heat at 450°F for 7 days. This was done, along with several other tests, with results as follows:

Tensile Strength and Elongation **a**. Ultimate Tensile Strength, psi Average % Exposure Range (5 Specimens/Condition) Elongation Average 62 680 (1) Control (No Exposure) 284 503 129 (2) Dry Heat, 450°F, 7 Days 232 (3) Immersed in Fuel 2 Days 160 140°F; Plus Dry Heat. 225 482 450°F, 7 Days (4) Same as (3) Except Exposed in a Closed Quart 220 347 18 Jar During Dry Heat DOEINO NO. 03-8297 PAGE 14 SECT

64

PEVLTR: A

E-2011 R1

- b. Reversion resistance was determined by injecting sealant in aluminum and steel tubes 6 to 24 inches long, and with inside diameters of .11 to .25 inch. The sealant generally reverted to an uncured fluid after 3 to 7 days exposure to dry heat at 450°F. The sealant appears unsuitable for enclosed injections.
- c. A control peel panel pulled 19 lbs./inch at room temperature with 100 percent cohesive separation.
- d. Original application time of this lot, as measured
 by "snap-time" was 2 4 hours at room temperature when
 the material was first received in December 1969.
 In January 1970 this had increased to about 48 hours.
 This indicates a considerable change during storage.
- e. The sealant cured adequately in a faying surface after 3 hours at 300°F, or 2 hours at 160°F plus 1 hour at 300°F. Adhesion to the primer was very poor. The sealant exhibited extensive "mud-cracking" or channeling, apparently caused by tearing due to thermal expansion and contraction.

BOEING	NO.	D3-8297
SECT	PAGE	15

PEVLIR: A

E-3033 R1

- f. Fillet cure properties were as follows:
 - (1) At room temperature the durometer hardness
 was only 20 after 10 days. Sealant was tackfree in approximately 48 hours.
 - (2) At 160°F the sealant was tack-free in 15
 minutes and cured to a hardness of 27 in
 3-1/2 hours.
 - (3) At 300°F the sealant curve to a hardness of
 37 in 70 minutes.

The sealant appeared to be unsuitable for faying surfice or injection sealing. However, the thermal stability appeared satisfactory and it was decided to go ahead and initiate environmental conditioning of the two test tanks.

BOEINO	NO. 03-8297
SF (T	PAGE 16

REVLTR: A

E-2033 RT

- [•]2.2 Preparation and Testing Of Physical Property Specimens
 - Twenty-one sets of the following specimens were prepared for environmental aging in the two test tanks.
 - a. Ultimate Tensile Strength and Elongation per ASTM-D-412 Four ASTM-D-412 Die C (half-scale: 0.5 by 2.0 inc., with 0.125 in. neck) specimens were prepared for each set from slabs prepared as described in paragraph 3, below. Specimens were tested before and after periods of environmental conditioning at a jaw separation rate of 10 inches per minute. Ultimate tensile strength and elongation was reported as the average of four specimens.
 - b. Volume Change per ASTM-D-471

Four specimens approximately 1 by 2 inches were prepared from slabs prepared for each set as described in paragraph 3, below. Volume change, reported as the average of four specimens, was determined after periods of environmental conditioning.

c. Hardness per ASTM-D-2240

Type A (Shore A) hardness was determined using the above volume change specimens before and after periods of environmental conditioning. Tests were in accordance with ASTM-D-2240 using a one kilogram weight and a five second reading. The hardness reported was the average of four specimens.

BOEINO	NO.	03-8297
SECT	PAGE	17

ſ

REVLTR: C

E-3033 R1

d. <u>Weight Loss</u>

Weight loss was determined using the above volume change specimens. Before and after periods of environmental conditioning, the specimens were conditioned for 24 hours in a dessicator and then weighed immediately. Percentage weight loss was calculated as follows:

$$(W_1 - W_2) \times 100$$

 W_1

where W_1 = Weight of sample before aging. W_2 = Weight of sample after aging.

Percent weight loss was reported as the average of four specimens.

e. Peel Strength

(1) One peel strength panel was prepared for each set using 0.05 by 2.9 by 6 inch panels from 6A1-4V titanium alloy. An equal number of 2.9 by 12 inch strips of 200 mesh stainless steel screen were also prepared. Panels were vapor degreased, followed by nitric-hydrofluoric acid etching. The screen was vapor degreased.

BOEINLS	NO. D3-8297
SECT.	PAGE 18

REVLTR: A

E-8033 R1

- e. <u>Peel Strength</u> (Continued)
 - (2) Immediately prior to preparing specimens they were solvent cleaned with BMS 11-7 cleaner per BAC 5504. Panels and screen were then primed with a thin coat of Dow Corning 77-037 primer. Primer was allowed to dry a minimum of 45 minutes prior to applying sealant.
 - (3) Sealant, mixed per paragraph 2 below, was applied to approximately five inches at one end of each panel to a depth of 0.125 ±0.025 inches using a suitable jig. The primed screen was impregnated for five inches on one end and placed on each panel in such a manner that the loose unimpregnated end faced the end of the panel free from sealant. The screen was smoothed down on the sealant carefully to avoid trapping air under the screen. An additional 0.125 ±0.025 inch thick layer of sealant was applied over the unimpregnated screen. The specimens were cured per paragraph 4, below.
 - (4) Before and after periods of environmental conditioning, two 1.0 inch wide strips were prepared on each panel by cutting completely through the screen and sealant to the metal length-wise along the panel and continuing completely along the unimpregnated

BOEING NO. **D3-8297** SECT. PAGE 19

REVLTR. A

E-3033 R1

- e. <u>Peel Strength</u> (Continued)
 - (4) (Continued)

screen. The loose end of each 1 inch wide strip in turn was clamped in one jaw of a suitable recording tensile testing machine and the acjacent end of the panel was fastened in the other jaw as shown in Figure 7. Cuts through the sealant under the screen were made so that an initial separation of sealant from the metal panel was promoted. The screen was pulled at an angle of 180 degrees from the panel and at the rate of 2 inches per minute in jaw separation.

(5) Cuts in the sealant to the metal panel at the junction of separation were made at an angle of 45 degrees towards the direction of separation at approximately 0.4 inch increments (approximately every 24 seconds) on the left side of the panel as shown in Figure 7. No cuts were required for 100 percent adhesive failure; however, any cohesive failure was treated as above. On the right side, except for the initial cut to promote separation, cuts were made only as necessary to prevent the sealant from paeling from the screen. All cuts extended completely across the strip being peeled and penetrated completely through the sealant to the panel.

BALEINA NO. 03-8297 SECT PAGE 20

REVLTR: A

E-2033 R1

- e. <u>Peel Strength</u> (Continued)
 - (6) The percent cohesive separation was determined from the ratio of cohesive separation area to total cohesive and adhesive separation on both test areas. The cohesive strength was determined during cohesive tear. The average of the cohesive strength, as determined from an extensiometer graph of the right side pull, was recorded. Values recorded during cutting or while load is being picked up after cutting were not included in the average. Panels which were environmentally exposed were tested within 24 hours after removal from the exposure condition.
- 2. The sealant was mixed in accordance with the following procedure:
 - a. Weigh onto a clean flat stainless steel plate or pan the correct amounts of base and activator immediately prior to mixing. Do not allow activator to contact the plate.
 - b. Hand mix by folding and squeezing the sealant compound with a spatula. Hix for a minimum of 5 minutes and until sealant compounds appears uniform.

BOEINO	NO. D3-8297
SECT	PAGL 21

FEVLTR. A

8-3033 R1

7	1
1	8

- c. Spread the sealant compound on a clean flat stainless steel plate or pan so that the maximum depth is less than 1/2 inch. De-gas the sealant compound for 10 minutes with a vacuum of 0.24 psia or better.
- d. Remove the plunger and plug the nozzle end of a cartridge for the Semco No. 250 gun. Scoop up scalant with a spatula, place it in the open end of the cartridge and drive it down by sharply rapping the nozzle end of the cartridge on something solid. Repeat until the cartridge is filled.
- e. De-gas the filled cartridge for five minutes with a vacuum of 0.25 psia or better. A plastic film may be used as an extension of the cartridge to prevent overflow of the sealant. Place the plunger in the cartridge using care to minimize air entrapment.
- f. Sealant was used within 2 hours after mixing.

BOEINO	NO. D3-8297
SECT	PAGE 22

REVLTR: A

E-3033 R1

and the second states and the second states

- 3. Sealant slabs were prepared as follows:
 - a. Sealant mixed per paragraph 2, was injected into
 a teflon lined closed mold to a thickness of 0.125
 +0.008 inches.
 - b. The mold was filled by extruding the sealant from a sealant gun with a Semco No. 440 nozzle. The nozzle was freed of air by a preliminary extrusion of 2 to 3 inches of sealant. During the casting operation, the tip of the nozzle was placed in an injection hole and was not removed until the mold was filled to excess.
 - c. Sealant was cured 48 hours minimum at room temperature plus 2 hours at 160°F. The sealant slabs were then removed from the mold and cured 1 hour at 300°F.
 - 4. Sealant specimens were cured a minimum of 2 hours at 160°F plus 1 hour at 300°F. Subsequent room temperature cure prior to environmental conditioning was at least 30 days.

BOEINO	NO.	D3-8297
SECT	PAGE	23

REVLTR: A

E-3033 R1

- 5. Ten sets of the above specimens were positioned inside of each test tank (see Tank No. 2 in Figure 6). These specimens were environmentally conditioned along with each tank. Specimens were removed periodically and tested as described above in paragraph 1. One set of specimens was a control (unexposed) set and was held at ambient laboratory conditions. This set was tested with the first set of environmentally conditioned specimens. Test specimens ready for installation in one tank are shown in Figure 8.
- Physical property specimen test results are reported in Table 1.

BOEINO	NO.	DJ-8297
SECT	PAGE	24

REVLTR: A

8-1011 RT

al hand a state of the state of

and a strange of the second strange

2.3 Environmental Conditioning

- 1. The environmental conditioning of the test tanks was accomplished by placing each tani in separate electrically heated aluminum chambers which were inerted with nitrogen. Figure 9 is a flow chart and Figure 10 is a photograph of the environmental conditioning test equipment. Figure 11 shows one chamber during installation of the side bayonet type strip heaters. Similar heaters were also installed on the bottom of the chambers. The chambers were insulated on the exterior with firebrick covered with fiberglas batting.
- 2. Thermocouples were placed on three heaters (two side and one bottom) for each chamber. These six thermocouple temperatures were continuously recorded on one six-point recorder. This recorder was equipped with an over-temperature control to preclude over-heating of the heaters.

BOEINO	NO.	D3-8297
SECT	PAGE	25

REVLTR: A

E-3033 A1

3. Thermocouples were installed in the tank interiors in the locations indicated in Figure 6. Thermorouples were also inserted in drilled holes in one exterior corner of each heavy steel end plate at the ends of each tank. Three thermocouple temperatures for each tank (top plate, tank interior, and bottom plate) were continuously recorded on the second six-point recorder. This recorder was also equipped with an over-temperature control to preclude over-heating of the tanks. For each chamber, the side and bottom heater temperatures were separately controlled by two Thermac Temperature Controller and Power Regulator Units (one for side heaters, one for bottom heaters). Each side heater Thermac Unit controlled side heater temperatures by means of a thermocouple in each tank interior. Each bottom heater Thermac Unit controlled bottom heater temperatures by means of a thermocouple in each tank bottom plate.

NO. 03-8297 SE.CT PAGE 26

A

REVLTR: A

E-3033 R1

and a complete state and a set of the state of the state

- 4. Initial heat up of one environmental chamber was conducted with an old previously tested titanium tank installed in the chamber to check tank temperature variations. This chamber, in its original configuration, had side heaters only. Temperature variations in any horizontal plane around the test tank were less than 10°F which was considered satisfactory. However, the temperature variations in a vertical direction was over 100°F (tank bottom was only 200°F when the tank top was 312°F). It was therefore considered Becssary to install heaters on the bottoms of the chambers.
- 5. Initially it had been planned to expose the tanks to liquid fuel for 1 hour at 140 - 150°F each working day (5 times a week). However, excessively long times were required to heat up the tanks for the elevated temperature fuel vapor exposure (approximately 2 hours) and to stabilize at the desired temperature range (approximately 8 hours, subsequently reduced to about 3 hours). It was therefore agreed that one weekly liquid fuel exposure of approximately 16 hours is satisfactory. This will allow a weekly elevated temperature fuel wapor exposure of approximately 130 hours.

D3-8297 DEINO NO. 27 PAGE

A

REVLTR A

8-3033 R1

- 6. During the elevated temperature environmental conditioning nitrogen and liquid fuel were added to each tank at the following rates:
 - a. Nitrogen was added at a rate of one tank volume every three hours. With a tank volume of approximately 27 gallons, this was a nitrogen purge rate of approximately 570 cubic centimeters per minute.
 - b. Liquid fuel was added at a rate of 0.1 percent tank
 volume every 90 minutes. This was calculated to be
 approximately 1.5 cubic centimeters per minute.
 - c. Nitrogen and fuel additions were metered using shielded type compact flowmeters (size 12 and 11 respectively) from Roger Tilmont Industries, Incorporated.
- 7. Fuel used for the first eight environmenta? conditioning cycles was MIL-J-5161 Grade JI with 18.2 percent aromatics. This fuel was purchased from Crystal Refining Company, Carson City, Michigan. Subsequently all fuel was ASTM D1455 Jet A with 14.8 to 15.3 percent aromatics, and purchased from Standard Oil of California, Salt Lake City, Utah.
- 8. The environmental conditioning for each cycle is shown in Tables 2 and 3 for Tank Numbers 1 and 2 respectively.

BOEINO	NO.	D3-8297
SEC1	PAGE	28

A

REVITR: A

E-3033 R1

2.4 Dynamic Load Cycling

2.4.1 Description Of Dynamic Test Device

- 1. The Dynamic Test Device was originally developed and used under Boeing-Air Force Materials Laboratory research contract reported in Reference 1. An improved version of this test device (Figure 12) was subsequently built under a Boeing research program. Torsional loads are applied by mounting a test tank horizontally as a cantilever beam, with one end firmly attached to the steel frame and the other end subjected to dynamic loading using a hydraulic ram. Cyclic loading is accomplished by a timer which operates a valve causing the ram to cycle.
- 2. To evaluate a tank at elevated temperatures, an insulated chamber attaches to the test jig, completely enclosing the test tank. The chamber is a stainless steel cylinder, consisting of two half-sections with fiberglass insulation, as shown in Figure 13. Heating of the tank is accomplished by blowing hot air between the tank and the insulated chamber.
- 3. To evaluate a tank at low temperatures, cold JP-4 fuel is inculated through the tank. The fuel is cooled to the desired temperature by pumping it through a coll immersed in a mixture of trichloroethylene and dry ice (see Figure 12).

BOEINO	NO.	D3-8297
SEC 1	PAGE	29

REVETR

ţ

E-3033 R1

• A

2.4.2 Determination Of Torsional Load Levels

- 1. Commercial Airplane Group stress analysis on the test tanks indicated that they were shear end fatigue resistant up to 159,000 inch-pounds torsional loading. Some tests were conducted to determine the range of joint deflections that would be obtained during cyclic loading. The tank used for this purpose was one used previously in an Air Force Material Laboratory Contract. This tank had been dissembled, stripped, cleaned up, and reassembled. This tank is identical to the two tanks being tested in this program, with the following exceptions:
 - a. It had been used previously at torsional loads up to 270,000 inch-pounds.
 - b. It was assembled with 0.25 inch steel nuts and bolts, torqued to 60-100 inch-pounds.
 - c. The access door is 0.10 inch thick 2024-T3 aluminum with two reinforced 4 inch diameter holes to permit measuring joint deflections by hand.
 - d. The center rib web contained no cut-outs.

BOEINO	NO. D 3-8297
SECT	PAGE 30

2

REVLTR: A

E-1011 Rt

2.4.2 (Continued)

- 2. It was very difficult to measure the joint gaps due to lack of visibility and the restricted access through the holes in the access doors. Joint gaps were measured manually by inserting small feeler gauges in the joints. Many of the extrusions used in the tank had curved edges making exact measurement impossible. The smallest gauge that could be handled under these circumstances was 3 mils. Initially (prior to loading) the joint gaps were generally 0 - 3 mils, with the following exceptions:
 - a. Center rib attachments to the spar web and the upper and lower skins: 3 13* mil gaps.
 - b. Spar web to spar chord adjacent to lower skin:
 3 11* mil gaps.

c. Lower skin at longitudinal stiffener: 13 - >18* mil gaps.
d. Upper skin at longitudinal stiffener: 3 - 5 mil gaps.

*Larger joint gaps were due to, (1) dimensional variations in titanium extrusions purchased back in 1963, (2) poor fit of center rib, and (3) possibly some permanent deformation of skins and web when previously used.

BOEIND	NO. 03-8297
SECT	PAGE 31

RLVLTR A

E-3033 R1

2.4.2 (Continued)

3. Torsional loading of this tank at 160,000 inch-pounds produced little measurable increases in joint gaps. Two places on the spar web increased--one 2 mils and one 4 mils. Torsional loading at 200,000 inch-pounds produced several increases in joint gaps:

a. Spar Web: 1 place 4 mil increase.

b. Upper Skin at

Longitudinal Stiffener: 1 place 2 mil increase.

1 place 5 mil increase.

c. Lower Skin at

Longitudinal Stiffener: Several places where gaps increased 4 - 10 mils.

4. Loading at this level did not appear to produce permanent visible deformation of the structure. However, when loaded to 240,000 inch-pounds torque, some slight permanent deformation of lower skin was observed. Therefore, the load levels chosen for the two test tanks was 160,000 inch-pounds (many cycles) and 200,000 inch-pounds (fewer cycles).

BOSINO	NO.	D3-8297
SECT	PAGE	32

REVLTR: A

E-1011 R1

2.4.3 General Test Procedure For Dynamic Cycling and Leak Testing Of Tanks

> Environmental conditioning of each test tank per Section
> 2.3 was interrupted periodically to conduct cyclic torsional load testing at a high temperature, low temperature, and at room temperature.

Torsional loading was conducted as follows:

Tank Test Temperature	500 Torsional Load Cycles At	10 Torsional Load Cycles At		
Room Temperature	160,000 inch-pounds	200,000 inch-pounds		
Low Temperature (-45 to -50°F)	360,000 inch-pounds	200,000 inch-pounds		
High Temperature (426 to 441°F)	128,000 inch-pounds	160,000 inch-pounds		

Load cycling rate was generally 8 - 12 cycles per minute. Each cycle included torsional loading in both directions.

2. After dynamic cycling at each temperature, each tank was leak tested by applying 10 psig pressure internally, applying aqueous soap solution to the tank externally, and recording all leakage as indicated by soap bubbles at the external leak location. Leakage is recorded in Appendix A.

BO SINS	NO.	D3-8297
SEC !	FAGE	33

REVLTR: A

#-3673 R1

2.5 Evaluation Of Tanks After Four Cycles Of Environmental Conditioning

The primary test areas of each tank were considered 1. to be the upper and lower skins and the spar. These areas, including the center rib, more closely simulated typical integral fuel tank structure and therefore the performance of the sealant in these areas is of primary significance. Areas of the tank that are out of the primary test area are the tank ends and the access door. The ends of the tank are composed of heavy steel end plates and jig attach fittings. These areas do not contain typical structure and may be subjected to unrealistic loads and deflections. The access door was faying surface sealed and attached by a single row of removable steel bolts. Door joint deflections and faying seal width is not typical of fuel tank structure. Therefore the tank ends (including areas under the jig attach fitting fingers), and the access door are not part of the primary test areas, and the performance of the sealant in these areas is of secondary importance only. These areas are designated secondary test areas.

MOEINO	NO.	D3-8297
SEC1	PAGE	34

REVLTR: A

#-1011 #1

84

~ A

2.5 (Continued)

- 2. Upon completion of four cycles of environmental conditioning (524 hours exposure to fuel vapor at 415-441°F, per Tables 2 and 3) both tanks were removed from the environmental chambers and leak tested. Leakage is summarized in Table 4. Specific leak locations are shown in Appendix A. There was considerable leakage in the secondary test areas (tank ends and access door). The reasons for these leaks appeared to be due to, (1) extrusion of sealant from the injections, and (2) loss of adhesion of the sealant to the Sermetal W painted steel parts in these areas. Considerable work was expended to clean up these leaking areas on the exterior of the tanks and to seal these leaks externally with the test sealant. Since the tanks were not opened at this point the cause of the fastener leaks in the primary test areas was not evident.
- 3. Both tanks were installed (separately) on the Dynamic Test Device and load cycled per Section 2.4.3 at an elevated temperature of 428-439°F. Prior to heating, one liter of test fuel was added to Tank No. 1, and three liters were added to Tank No. 2. The tanks were continuously purged with nitrogen and were not pressurized.

On completion of dynamic cycling, the tanks were again leak tested. Leakage is shown in Table 4.

85

REVLTR: A

BOEINO	And the Party of t		the second second second
SECT	PAGE	3	5

E-3013 R1

W. Lower

2.3 (Continued)

6.4 4 4 66 C + 10 0

4. The tank access doors were removed, and examination of the interior revealed the following:

a. Fastener Leaks

The fastener leaks were generally due to, (1) loss of sealant adhesion to titanium adjacent to fasteners (see Figure 14), or (2) small cracks in the sealant. Numerous fasteners not leaking also exhibited these conditions. The causes of the 22 fasteners leaking in Tank No. 1 were generally much less evident than for Tank No. 2.

b. Joint Leaks

Although only Tank No. 2 contained joint leakage, both tanks exhibited cracking and/or loss of aúhesion of joint fillets at corners (where fillets make a 9G degree turn). At the center rib Tank No. 1 and No. 2 had two and five cracked corner fillets respectively (see Figures 15, 16, and 17). Both tanks also had cracked and loose fillets at the corners of the end plates.

BOSINO	NO,	D3-8297
SECT	PAGE	30

REV LTR: A

- 2.5 (Continued)
 - 5. Apparent causes of sealant failures were:
 - a. Poor sealant adhesion to titanium.
 - Low adhesive and cohesive strength of sealant at elevated temperatures (peel strength is only 2.5 pounds per inch at room temperature per Section 2.2).
 - c. Thermal expansion problems (expansion of injection seals under fillets in corners may be the cause of corner fillet failures).
 - d. Undersize applications of sealant over fasteners, or entrapped air bubbles in sealant may have caused a few fastener leaks.
 - 6. Joint fillets not in corners were generally satisfactory in both tanks. The fifteen joint leaks in Tank No. 2 were generally adjacent to leaking fasteners or corners where joint fillets were cracked or loose. Tank No. 2 was completely faying surface sealed, but exhibited 15 joint leaks and 51 fastener leaks. Tank No. 1, faying sealed at center rib only, exhibited no joint leaks and only 22 fastener leaks. This was a paradox not readily understood. It appeared that the faying surface seals in Tank No. 2 were completely ineffective.
 - It was decided at this point to, (1) repair discrepant sealant in Tank No. 1 and continue testing, and (2) to terminate testing of Tank No. 2 because of excessive leakage.

SECT PAGE 37

REVLTR:

E-3053 M1

2.6 Repairs and Continuation Of Testing Of Tank No. 1 2.6.1 Repairs Of Tank

> 1. Four pounds of Dow-Corning 77-028 sealant, Lot No. 104105 and one pint of 77-037 primer, Lot No. E23-276, were received on April 24, 1970, to repair Tank No. 1. Preliminary testing showed that the new sealant would not adhere to cured unexposed sealant, and when applied to sealant previously exposed to fuel the new sealant would not cure at the interface. Subsequent testing showed that if the fuel exposed sealant was dried to remove the fuel (overnight at about 425°F) and the exposed surface then cut away and fresh sealant applied, marginal compatibility (2 pounds per inch peel strength) could be obtained. The tank was therefore exposed to dry heat for 18 hours at 425°F. Some additional small cracks in fillets and areas of poor adhesion were discovered during repairs. The sealant was very easy to damage. Cutting away discrepant sealant frequently caused adjacent sealant to crack or come loose.

BOEINO	NO.	D3-8297
SECT	PAGE	38

REVLTR: A

- 1033 R1

2.6.1 (Continued)

2. All fillets showing cracks and/or poor adhesion were removed and replaced with new sealant. Cleaning of the metal was accomplished with aluminum wool, Scotchbrite, and gauze dampened with BMS 11-7 cleaner. Leakage before and after repairs is summarized in Table 5 (Leakage Test No. 2 and 3). In general, the repairs eliminated very few leaks. Sixteen old fastener leaks were eliminated, but 20 new ones developed. Additional leakage was probably due to, (1) damage of sealant during repairs, and (2) to the need to bake the aged sealant to remove the fuel in order to make the aged sealant repairable. Leakage at the tank ends was not reduced appreciably, but load cycling and additional environmental conditioning was still considered feasible.

2.6.2 Dynamic Load Cycling After Repairs

1. The tank was installed on the dynamic test device and load cycled per Section 2.4.3 at -47 to -49°F and at room temperature. A detailed leak test could not be conducted readily at this point since the tank was still on the dynamic test device and was wet with fuel which prevented the soap solution from working properly. However, leakage recorded is shown in Table 5 (Leakage Test No. 4).

89

BOEINO	NO. D3-8297
SECT	PAGE 39

REVLTR: A

E-3033 M1

2.6.2 (Continued)

- 2. The tank was subsequently heated to 429-441°F and load cycled for 20 cycles at 160,000 inch-pounds torque plus 30 cycles at 128,000 inch-pounds. This hot load cycling was conducted to see if the new fillet repairs would crack with no prior environmental conditioning. Leakage is summarized in Table 5 (Leakage Test No. 5). Ten joint leaks and 80 fastener leaks were present in the primary test areas. Approximately 54 of the fastener leaks were new. Seven of the ten joint leaks were probably the source of seven joint leaks, and may be the source of the other three.
- 3. Examination of the tank interior after the above described load cycling can be summarized as follows:
 - a. The new fillets installed during repairs showed no major cracking, but there were numerous areas of loss of adhesion to the structure and the old sealant. This was about 20 percent of the repaired areas. There were some small cracks in new and old sealant in fillets at ends of tank in corners.
 - b. Old sealant fillets in tank ends show additional loss of adhesion.
 - c. Cause of the many additional fastemer leaks was not apparent.

d. Thefillets in general continue to look good.AJLCTPAGE 40

E-1013 (11

REV LTR:

90

, A

2.6.2 (Continued)

- 4. After the above load cycling the access door was again leaking badly. During the room temperature cycling it was observed that the access door was moving (sliding) excessively at the faying surface due to the oversize fastemer holes in the door (which were necessary to make the door fit). A new door was fabricated and installed prior to continuation of environmental conditioning per Section 2.6.3. A small amount of a low expansion sealant (Dow Corning 77-066, Lot Z-40-15) was used in the prepack channels at each end of the access door; however, this was not considered in be a good evaluation of this material.
- 2.6.3 Evaluation Of Tank No. 1 After 12 Cycles Of Environmental Conditioning
 - Tank No. 1 was exposed to eight additional cycles of environmental conditioning per Table 2 (Cycles 5 through 12). Accumulated exposures after 12 cycles were as follows:
 a. Liquid Fuel Exposure

 Ambient
 139°F
 204.6 Hours

 140°F
 150°F
 173.0 Hours

 152°F
 175°F
 <u>30.5</u> Hours

 Total Liquid Fuel Exposure
 408.1 Hours At Ambient

 Total Liquid Fuel Exposure
 408.1 Hours At Ambient

BØEINO	NO. D3-8297
SECT	PAGE 41

REVLTR: A

E-1011 R1

2.6.3 (Continued)

b. Fuel Vapor Exposure

A	mbient	-	4 00°F	118.0	Hours			
	401°F	-	425°F	51.3	Hours			
	415°F	-	441°F	624.5	Hours			
	426°F	•	441°F	939.1	Hours			
	443°F	-	457° F	25.2	Hours			
Total	Fue 1	¥apor	Exposure	1758.1	Hours		Ambient 457°F	
				1563.6	Hours	At	415 - 441°F	

- 2. The tank was then removed from environmental conditioning and leak tested. Leakage is shown in Table 5 (Leakage Test No. 6).
- 3. The tank was then load cycled per Section 2.4.3 at all three temperatures and again leak tested (see Table 5, Leakage Test No. 7). There are 24 joint lengths, separated from each other by faying surface seals, in the primary test areas (eight each on top skin, lower skin, and spar). It is difficult to summarize the leakage since many leaks "come and go" (as seen by comparing the total leaks obtained during Leakage Test No. 6 with those obtained during (Leakage Test No. 7).

BORINO	NO.	93-8297
SECT	PAGE	42

REVLTR: A

E-3033 A1

92

Ą

2.5.3 (Continued)

4. The tank access door was removed and the sealant examined. Appendix A (Page A-11) shows the locations and types of joint fillet defects found, which can be summarized as follows:

Symbol	Type Of Defect	Of Defects
(15)	Small crack or loss of adhesion. Crack generally penetrates to sub- strate with some accompanying loss of adhesion.	40
14	Crack in repair fillet.	1
13	Lucs of adhesion of repair fillet to titanium.	2
(12)	Loss of adhesion of repair fillet to original fillet.	2

BLEIND	NO. D3-8297
SECT	PAGE 43

REVLTR: A

E-3033 R1

/ A

Number

2.6.3 (Continued)

- 5. In an effort to determine where the leaks originated, an interior leakage analysis test was conducted by placing a transparent access door on the tank, applying 5 psig negative (vacuum) pressure to the tank interior and observing leaks on the interior bubbling through a 1 to 2 inch layer of water. It was desired to determine the following:
 - (1) Whether the exterior leaks were primarily caused by leaking fasteners or joint fillets.
 - (2) Whether joint fillets were leaking in straight fillet areas or only at corners (where cracks and loss of adhesion had made previous repairs necessary).

Interior leakage obtained by this method is shown in Appendix A (Page A-12) and the numbers of leaks obtained can be summarized as follows:

	Relative Rate Of Leakage				
Origin Of Leak	One Bubble Indication (1)	Small Continuous Bubbling	Large Vigorous Bubbling		
Sealant over fastener	41 Leaks	9 Leaks	1 Leak		
Joint fillet at or near corner	8 Leaks	4 Leaks	2 Leæks		
Joint fillet in straight-seam areas	28 Leaks	7 Leaks	0 Leaks		
Uncertain (Joint fillet and sealant over fastener are continuous)	0 Leaks	3 Leaks	0 Leaks		

1 One bubble forms on sealant but does not break and reform. These can be classified as possible or suspected leaks only.

INO. D3-8297 PAGE 44 SECT

v A

REVLTR: A

E-1011 A1

a bear do a birth and a bir

alanda mula adamating ana ata

2.5.3 (Continued)

- 6. An attempt was made to correlate the exterior leakage (Table 5), the interior sealant defects (Item 4 above) and the interior leakage (Item 5 above). In general, there is <u>very little correlation</u>; however, the following items are considered significant:
 - a. On the tank exterior 11 to 16 percent of the fastemers
 and 25 to 33 percent of the 24 joint areas are leaking.
 - b. On the tank interior 10 fastener leaks were confirmed, ith 41 more suspected. Thirteen joint fillet leaks were confirmed, with 36 more suspected.
 - c. Of the 13 confirmed joint fillet leaks, 7 leaks were in straight seam areas (not at or near corners). None of these 7 leaks were at the 32 defects mentioned in Item d., below.
 - d. Forty defects (small cracks and/or spots showing loss of adhesion) were observed in the joint fillets. Thirty-two of these defects were in straight seam areas (not at or near corners).
- 7. At this point it appeared that the sealant was cracking and losing adhesion because of the low physical properties resulting from inadequate resistance to the environmental conditioning and structural loading. Continuation of the testing of this tank in Section 2.7 was primarily to evaluate sealant repair procedures.

MOEINO NO. 03-8297 PAGE 45 SECT

REV LTR:

E-1011 R1

Strate Antis and an and the state

2.7 Evaluation Of Sealant Repair Procedures In Tank No. 1

- Defective and/or leaking joint fillets were removed from 14 locations (crosshatched areas in Appendix A, Page A-12). In order to evaluate the repairability of the fuel aged discrepant sealant, these areas were repaired in August 1970 using sealant received from Commercial Airplane Group. The following repair procedure was used:
 - Fuel aged sealant was dried 16 hours at 420°F to remove fuel.
 - b. Dried sealant was then scrubbed with cheesecloth and SMS 11-7 solvent to clean and remove gloss.
 - c. Structure was cleaned with Scotchbrite and cheesecloth wet with BMS 11-7.
 - d. A thin coat of primer was applied to structure only and dried 1 to 2 hours.
 - e. A thin brush coat of freshly thawed sealant was applied to structure and old sealant with acid brushes.
 - f. Sealant fillets were applied within one hour.
 - g. Sealant was cured overnight at room temperature, 2 hours at 160°F and 1 hour at 300°F.

NO. 03-8297 SECT PAGE 46

REVLTR: A

E-3033 R1

2.7 (Continued)

- 2. It was not possible to clean the old sealant without causing some loss of adhesion. No ettempt was made to repair the leaking fasteners. The ends of the tank (out of the primary test areas) were still leaking considerably and not considered repairable. (Fuel leakage during the 16 hour liquid fuel exposure portions of environmental conditioning cycles 8 through 12 ranged from 1200 to 10,300 milliliters.)
- 3. The tank was exposed to 8 additional cycles of environmental conditioning per Table 2 (Cycles 13 through 20). Total accumulated exposures after 20 cycles were as follows:

a. Liquid Fuel Exposure

Ambient	-	140°F	236.6 Hours
140°F	-	150°F	290.5 Hours
152 ° F	-	175°F	<u>54.0</u> Hours

Total Liquid Fuel Exposure 581.1 Hours At Ambient To 175°F

b. Fuel Vapor Exposure

Ambient	-	400°F	220.6	Hours	
401°F	-	425°F	67.i	Hours	
415 °F	•	441°F	637.0	Hours	
426 ° F	-	441*F	1983.1	Hours	
443°F	-	457°F	42.2	Hours	
Total Fuel Va	por E	kposure			At Ambient To 457°F At 415°F To 441°F

The fuel vapor exposure after the repairs reported in Item 1, above was 1056.5 hours at 415°F to 441°F MO. D3-8297

PAGE

REVLTR:

E-1011 R1

2.7 (Continued)

e al 75 al 2718 et diversitation dia di thate di tanàna dia amin'ny dia

a Bischild Bill and Aller States and

4. The tank was removed from environmental conditioning and leak checked by putting approximately 2 inches of water in the tank, installing an acrylic door, applying 5 psi of vacuum and observing bubbles. Results are summarized in Appendix A (Page A-13). There were 2 leaks in one of the 6 repaired areas of the upper panel. In the lower panel there was 1 leak in one of the 3 repaired areas, one new fillet leak and 4 new fastener leaks. The spar had 2 of 5 repaired areas leaking and 1 new fastener leak.

A number of cracks in the thin faired portion of the fillets was observed. Under these cracks are bare spots where the sealant separated adhesively. A portion of the cracked fillets were removed in 4 locations. Also, one leaking repair in the spar was removed, revealing a section where there was no adhesion.

These were all repaired according to the procedure described in Item 1 above, except that 400 grit aluminum oxide sandpaper was used to abrade the surface. Following repair and an overnight bake in air at 180°F, three of the repairs were removed to check adhesion. Good adhesion had been maintained. These three repairs were then repaired again.

BOEINO	NO. D3-8297
SECT	PAGE 48

REVILTR: A

2-3033 R1

98

A

v 2.7 (Continued)

- 5. The tank was closed up and placed in the environmental chamber where it was filled with fuel and exposed at room temperature for 14 days. It was refilled periodically as fuel leaked out.
- 6. The tank was then placed on the dynamic test device and load cycled as follows:
 - a. 500 Cycles at 426 to 441°F and 128,000 inch-pounds torque.
 - b. 510 cycles at room temperature (500 cycles at 160,000 inch-pounds torque and 10 cycles at 200,000 inchpounds torque).
- 7. The tank was then environmentally conditioned in accordance with Table 2, Cycles 21 through 31. Environmental conditioning was the same as used previously (Section 2.3) except the liquid exposure for each cycle was reduced to one hour due to the excessive leakage of this tank.
- 8. Environmental conditioning of Tank No. 1 was terminated upon completion of Cycle 31 due to program cancellation in March 1971. In August 1971 the program was reactivated and Tank No. 1 was leak tested and examined after 31 environmental conditioning cycles (see Table 2). The leakage observed is shown in Appendix A (page A-13). With the exception of the new leak at the 1/4" crack in the spar fillet,

BOESNA	NO. D3-8297
SECT	PAGE 49

REVLTR: B

E-3033 #1

99

Å

B

all new leaks were small and nearly undetectable. Only two of these new leaks were in any of the repairs made after 20 environmental conditioning cycles and they were in the same repaired section. There were 10 new leaks in repairs made after 12 environmental conditioning cycles, and 12 new leaks in original sealant. Note however, that no new leaks occurred in seven of the fourteen repairs made after 12 environmental conditioning cycles. These repairs had accumulated a total of 2298 hours fuel vapor exposure, while the repairs after 20 environmental conditioning cycles had accumulated 1242 hours fuel vapor exposure. The only difference in the repair procedures was that in the latter case the surfaces to be repaired were abraded using 400 grit emery paper rather than the Scotchbrite used in the former case. The test results do not indicate that the severe abrading was any more effective.

9. Testing of Tank No. 1 is considered complete. No additional testing of this tank is planned.

D3-8297 NO. 49.1 SECT PAGE

REVLTR: B

E-3033 #1

100

B

3. TESTING OF TANK NO. 3

- 3.1 General Description
 - 1. This tank was designed, fabricated, and sealed by Commercial Airplane Group per Reference 4, and was intended to simulate the SST fuel tank structure more closely than the first two tanks. The tank was built to much closer tolerances than was possible for the first two tanks (which were built in 1954 with badly warped extrusions that could not be properly machined). The tank is 10 by 18 inches in cross section and 50 inches long. Skin and spar webs are chemically milled and the top skin contains a full size elliptical shaped fuel tank access door. The tank is of conventional skin and stiffener type construction with joggled internal skin stiffeners, and exterior skin and spar stiffeners. It has no intermediate ribs. The tank was primarily fillet sealed with Dow Corning 77-028 (same sealant used in previous tanks), but also contained a few primary faying surface and injection seals on the lower skin.

BOEINO	NO.	D3-8297
SECT	PAGE	50

2-1015 R1

and the second state of the second state of the second states

1

an de statut de la constantia de la constan

3.2 Initial Preparation and Testing Of Tank

- Tank No. 3 was leak tested initially by applying 5 psig internal pressure and immersing the tank in water. One fastener was leaking (see Appendix B, Page B-1).
- 2. Holes in the tank end plates (for installation on the dynamic test device) were located, drilled and tapped (sixteen 1/2 inch diameter holes in each end, drilled about half way through the 2 inch thick end plates).
- 3. Six thermocouples were installed in the tank (two each on upper and lower skin stiffeners, and one in the center of each end plate).
- 4. The tank was placed on the dynamic test device and initial torsional loads were applied to measure the extent and depth of web shear wrinkling at five locations. Test data is shown in Appendix C. Initial web buckling was obtained at approximately 50,000 inch-pounds of torque. Results showed that at 216,000 inch-pounds of torsion (typical flight load), significant web wrinkles <u>do</u> extend into the pad-up areas of the chem-milled webs and even into the chords and stiffeners.

BOEINO	NO. D3-8297
SECT	PAGE 51

REVLTR: A

E-3033 R1

102

▼ A

3.3 Dynamic Load Cycling

3.3.1 General Test Procedure

in water.

Dynamic load cycling of Tank No. 3 was conducted per Section 2.4.3 with the following exceptions:

a. Torsional load levels were as follows:

<u>Tank Test Temperature</u>	500 Torsional Load Cycles At
Room Temperature	216,000 inch-pounds
Low Temperature (-45 to -50°F)	216,000 inch-pounds
High Temperature (426 to 441°F)	172,800 inch-pounds
b. Leak testing was conducted at 5 psig wit	h the tank immersed

3.3.2 Initial Load Cycling Prior To Any Environmental Conditioning

1. The tank was load cycled per Section 3.3.1 at room temperature, at the high temperature, and at the low temperature. The low temperature cycling was terminated at 330 cycles when the ends began leaking fuel and it was discovered that the bolts in the end plates were loose. Both rows of bolts in the upper skin and both spars were very loose (2 to 3 turns). Only a few of the bolts in the lower skin were loose and these only slightly. Four bolts in the upper skin had broken at the threads. All bolts were subsequently B replaced with CRES bolts, B30LM3HU, lock-wired in place.

B . JE INO	NO.	D3	-829	7
SECT	PAG	E	52	

REVLTR: B

E-2033 #1

3.3.2 (Continued)

yyddy thry ann nighteissin e cyfyte . Alsystemae gyn mewnedy i falsen eniffest etter anter anter a the trebe bet the state in the st

- 2. After the above load cycling (but prior to installing new bolts) the tank was leak tested. Leakage is shown in Appendix B. Page B-2. There were two joint leaks in the lower skin and two corners were leaking at the ram end. The fastener leak was the same one leaking prior to any load cycling. Based on the fuel leakage from the corners during dynamic cycling it is probable that more than two corners would be leaking if the tank could be leak tested under load.
- 3. The tank was then opened and the interior examined. Numerous cracks in the fillets were found in the lower skin as shown in Appendix B, Page B-3. Eight cracks were in joint fillets and two were in sealant over fasteners in the primary test areas of the tank (excluding tank ends). Cracks were also found in four of the eight corners of the tank (four of these cracks appeared to be due to separation of repair sealant). Vacuum leak testing at 3 psig with a layer of water C on the bottom skin revealed the source of two exterior leaks to be fillet seal cracks (see Appendix B, Page B-3). Portions of the fillets containing cracks at the two leak locations were carefully removed to preserve the cracked section. <u>Adhesion</u> was good except at one fastener.

NO. D3-8297 PAGE 53

REVLTR: C

E-2033 R1

3.3.2 (Continued)

- 4. The tank was then mounted on the dynamic test device. Members of the Stress Group were in attendance to observe the wrinkles formed during loading and to assess the effect on structural life of the tank. It was concluded that the tank would easily last for the program life. The wrinkles were very prominent. Three to 4 thousandths of an inch gap could be measured between the skin and stiffeners on the outside of the tank in one or two places; however, deflections inside were less than .0015 inch (in the areas where there were no fillets and measurement was therefore possible).
 5. The tank was load cycled at room temperature for an additi-
- onal 1500 cycles in an effort to see if additional fillet cracks would be obtained or if existing cracks would grow.
- 6. The two areas where cracked fillets were removed (Itom 3) were repaired, and the tank was load cycled per Section 3.3.1 at the elevated temperature (426 to 441°F). Small test specimens were attached to the tank during the loading. They consisted of heavy fillets on 6A14V titanium. One was for control, one had been "puddled" extensively, and one was permanently deformed to put the fillet in tension. Following the exposure with the tank, these specimens were thermally shocked a number of times by heating to 440°F and immediately placing in contact with a -40°F surface. In no instance did we produce new cracks or extend the old ones.

NO. 03-8297 PAGE SECT

REVLTR: B

E-1011 A1

105

r A

8

B

3.3.2 (Continued)

A. T. P. L. J. S. Stat. & Styles

A South States

- 7. The tank was then leak tested (see Appendix B, Page B-4). One new leak was found on the lower skin. The tank was opened and the fillets examined (see Appendix B, Page B-5). Five small fillet cracks were found on one spar fillet. These were very small and <u>may</u> have been overlooked previously. Three small cracks were found on fillets of the lower skin (1 seam fillet crack and 2 fastener fillet cracks).
- 8. Leaks were repaired in four areas of the tank identified by (B) in Appendix B, Page B-6. These repairs stopped all but one leak (which had not been repaired because the interior source could not be identified).
- 9. The length of all existing cracks was measured and recorded (see Appendix B, Page B-3).
- 10. Physical property test specimens were installed in the tank, the original access door installed, and environmental conditioning was initiated on January 13, 1971.

BASINO	NO.	03	- 8297	
SECT	PAG	E	55	

REVILTR: A

E-3033 R1

- 3.4 Preparation and Testing Of Physical Property Specimens For Tank No. 3
 - Physical property test specimens were prepared from slabs of cured sealant provided by Commercial Airplane Group.
 Slabs were cast from Batch No. 205180 on October 8, 1970.
 Ten sets of specimens were prepared. Each set contained:
 - a. 15 Tensile Strength and Elongation Specimens.
 - b. 3 Volume Change Specimens.

Five peel panels (prepared by Commercial Airplane Group) and nine sets of the above specimens were placed in Tank No. 3. One set was a Control (Unexposed) Set.

- 2. In general, test specimens were prepared and tested in accordance with Section 2.2. Tensile Strength and Elongation Specimens were tested at three temperatures (5 specimens per condition): 75 ±5°F, 450 ±5°F and -50 ±5°F.
- 3. Test specimens were removed from the tank and tested per B Section 2.2, paragraph 1. Results obtained are reported B in Table 6.

BOEINO	NO.	D3-	8297
SECT	PAGE	5	6

A

В

REVLTR: B

1

E-3033 R1

3.5 Environmental Conditioning Of Tank No. 3

- 1. Environmental conditioning of Tank No. 3 was generally accomplished as described in Section 2.3. The specific environments for each environmental conditioning cycle is shown in Table 7. Eight cycles were completed when this program was cancelled in March 1971. The tank remained closed up and essentially empty (in the Figure 10 Environmental Chamber) until work resumed in August 1971.
- 2. The tank was removed from the environmental chamber and leak tested at 5 psi while immersed in water. Leakage is shown in Appendix B, page E-7. Six of eight corners were leaking badly. Five small seam leaks were found on the lower skin. Examination of the interior revealed numerous new cracks. The six leaking corners contained large cracks. When the tank was dried out (at 310°F for 16 hours) in preparation for repairs, additional cracks appeared and many increased in length. Locations of interior cracks are shown in Appendix B, page 3-8. A 3 psi vacuum leak test with water on the interior lower skin revealed 7 interior leaks as shown in Appendix B, page B-9. These leaks were small. The numerous cracks in the sealant fillets were the main item of concern. Those in the corners were gapping and up to 1.5 inches long. New cracks were found in fillets at ends of lower skin stiffeners. It appeared that the thicker the fillet the greater the chances of it cracking. Eight

NO. D3-8297 SECT PAGE 57

R

REVLTR: B

E-2033 R1

cycles of environmental conditioning did not cause the fillet cracks found initially (prior to environmental conditioning) to grow or cause leaks.

- 3. The 6 leaking corners were repaired and the tank was again leak tested. See Appendix B, page B-10. Two of the six corners were still leaking. One new small seam leak was found in the upper skin.
- 4. The tank was installed on the dynamic test device and load cycled at all three temperatures per Section 3.3.1. The tank was removed from the dynamic test device and leak tested. Leakage is shown in Appendix B, page B-11. Seven new leaks were observed, but were not large enough to require repairing. Examination of the tank interior showed no significant changes.
- Environmental conditioning of Tank No. 3 was resumed on September 7 with Cycle No. 9, and was terminated on December 23, 1971 with Cycle No. 22. See Table 7.

BOEINO	NG.	D3-8297
SECT	PAGE	57.1

REVLTR: B

ころいたからないないないないないです。 やま いっしゅうゆう

K-1015 R1

109

B

- 3.6 Dynamic Load Cycling of Tank No. 3 After 22 Cycles of Environmental Conditioning
 - 1. After 22 cycles of environmental conditioning (see Table 7) Tank No. 3 was leak tested by immersion in water as previously described. Leakage is shown in Appendix B, page B-12. There were 9 new leaks, including 2 new corner leake. One of the 2 new corner leaks was a previously repaired corner, the other was a leak in the original sealant. Except for the corners all leaks were very small. The access door was leaking badly also.
 - 2. The tank was opened and the interior examined. The leakage of the access door was due to 100 percent compression set of the fluorosilicone door seal. All defects in the tank interior are recorded in Appendix B, page B-13. There were approximately 28 new defects in the tank since the last examination (after 8 environmental conditioning cycles, page B-8).
 - 3. The tank was again placed on the dynamic test device and load cycled at all three temperatures per Section 3.3.1. The tank was removed and leak tested immersed in water. Leakage is shown in Appendix B, page B-14. Six new leaks were obtained, for a total of approximately 24 leaks. A few leaks traveled to adjacent areas or disappeared. All leaks, except for the 6 leaking corners, were very small.

NC. D3-8297 PAGE 57.2

C

E-3033 R1

- 4. The six leaking corners were repaired and the tank leak tested. Leakage after repairs is shown in Appendix B, page B-15. While leak testing, an additional corner (F-2) was found to be leaking badly. This appeared to be due to a crack approximately 1/4 inch long in the sealant that apparently opened up when the tank was dried out prior to repairs. The sealant was chamfered back and the crack well filled with fresh sealant.
- 5. Table 8 shows the corner leakage history of tank No. 3.
- 6. The door was installed using a gasket made of fluorosilicone rubber sheet. The tank was placed in the environmental chamber and environmental conditioning was resumed on 23 February 1972.

BOEING	NO.	D3-8297
SECT	PAGE	57.3

REVLTR: D

E-3033 #1

- 3.7 Dynamic Load Cycling of Tank No. 3 After 39 Environmental Conditioning Cycles (August 1972)
 - 1. After completion of 39 environmental conditioning cycles (see Table 7) Tank No. 3 was removed from the environmental chamber and leak tested. Leakage is shown in Appendix B, page B-16. Prior to repairing leaks the tank was dried out 16 hours at 400°F. The tank interior was then examined and fillet cracks are shown in Appendix B, page B-17. A few new cracks were evident. It was subsequently decided to repair leaks <u>after</u> load cycling the tank since no primer was available at this point.
 - 2. The tank was placed on the dynamic test device and load cycled at all three temperatures per Section 3.3.1. The tank was removed and again leak tested. Leakage, shown in Appendix B, page E 18, can be summarized as follows as compared with leakage after 22 environmental conditioning cycles:

Leak Location	After 22 Cycles	After <u>39 Cycles</u>
Corners (see Table 8)	4	4
Seams	15	7 (1)
Fasteners	3	0

(1) 4 new seam leaks; 12 seam leaks disappeared.

There was no significant increase in interior fillet cracks after the load cycling (see page 8-17).

BOEING	NO.	D3-8297
SECT	PAGE	57.4

REVLTR: D

E-1033 R1

- 3. Of the four leaking corners only two (R-1 and R-2) exhibited interior defects which appeared to be the cause of the leakage. These two corners were leaking significantly and were therefore repaired as follows.
 - All defective seglant in corner area was removed,
 including sealant in the two injections adjacent
 to each corner.
 - Metal was scrubbed with aluminum oxide paper
 and cleaned with BMS 11-7 cleaner.
 - c. A thin coat of a new two-part primer (Dow Corning 77-123) was applied to the metal. This primer replaced the previously used Dow Corning 77-037 which is no longer available.
 - d. The primer was dried 90 minutes at room temperature, and freshly mixed 77-028 sealant applied. Sealant was cured immediately at 300°F for 1 hour minimum.

The 1wo repaired corners were leak tested and showed no leakage.

- Test specimens removed from the tank were tested as before and results are reported in Table 6.
- 5. Remaining test specimens and new thermocouples were installed in the tank and environmental conditioning was resumed on 6 September 1972. The fluorosilicone rubber gasket on the door was in good condition and was not replaced.

BOEING	NO.	D3-8297
SECT	PAGE	57.5

REVLTR: D

E-3033 R1

T

3.8 Repair of Tank No. 3 After 46 Environmental Conditioning Cycles (November 1972)

- 1. After completion of the liquid fuel exposure portion of environmental conditioning cycle no. 47, approximately 11.5 gallons of fuel had leaked out of the tank into the environmental chamber. Prior leakage had been gradually increasing from 3/4 gallon (cycle no. 42) to 5-1/2 gallons (cycles 45 and 46). Due to the excessive leakage the tank was removed from the environmental chamber and leak tested with 5 psig air pressure. All 8 corners of the tank ware leaking (see Appendix B, page B-19).
- 2. Since most of the leakage would be from those corners that ware at the bottom of the tank during environmental conditioning, it was decided to repair only these four corners (R-1, R-2, R-3, and R-4). Corners R-1 and R-2 were previously repaired in September 1972 (see Table 8), and it was apparent that the new sealant in the injections adjacent to these corners had expanded and failed the fillets in these corners. These two corners were repaired with new fillets and the adjacent injections were cleaned out and left empty. All 4 corners were repaired using the procedure used in September (see Section 3.7, Item 3) except sealant was cured overnight at ambient temperatures prior to curing at 300°F.
- 3. After the repairs the bottom four corners were leak tested and only corner R-4 was still leaking. This was a small leak and was not repaired. Environmental conditioning was resumed on November 20.

REVLTR: D

BOEING	NO.	D3-8297
SECT	PAGE	57.6

E-3033 R1

- **3.9** Dynamic Load Cycling of Tank No. 3 After 54 Environmental Conditioning Cycles (February 1973)
 - After completion of 54 environmental conditioning cycles (see Table 7) tank no. 3 was removed from the environmental conditioning chamber and installed on the dynamic test device. The tank was load cycled at all three temperatures per Section 3.3.1. The tank was removed and leak tested. Leakage is shown in Appendix B page B-20. There were 26 seam leaks not including one seam on the upper skin which was leaking slightly over almost its entire length of 48 inches. All eight corners were leaking, primarily at the two injections in each corner.
 - 2. The tank was opened and examined. There was no significant change in interior fillet cracks over that previously reported. Interior leakage as indicated by air at 3 psi vacuum bubbling through a water layer is shown in Appendix B, page B-21.
 - 3. The four corners which would be at the bottom of the tank during environmental conditioning (R-1, R-2, R-3, and R-4) were repaired. All sealant in these corners, including the adjacent injections, was removed. The fillets between corners were also removed. Structure was cleaned as previously described and primed with a new sample of Dow Corning 77-123 one-part primer. The primer was air dried 90 minutes at ambient temperature and frozen Dow Corning 77-028 (Lot 40111/, mfg. Aug. 1971) was applied. The sealant was cured at ambient temperatures overnight and then o.on cured 1 3 hours at 300°F. The corners were leak tested and were leak-free.
 - Laboratory test specimens were removed, tested, and results added to Table 6. Environmental conditioning was resumed on February 19.

BOEING	NO. D3-8297	
SECT	PAGE 57.7	

REVLTR: D

Ł

E-3033 R1

3.10 Dynamic Load Cycling of Tank No. 3 After 74 Environmental Conditioning Cycles (August 1973)

- 1. During the liquid exposure portion of environmental conditioning cycle no. 71 the tank leaked 16 gallons of fuel. The tank was removed from the environmental conditioning chamber and leak tested. One corner at the tank bottom (R-2) appeared to be causing the leakage. This corner was repaired with new sealant using previously described procedures. The door seal was badly cracked and may have been leaking. This seal was replaced with a .125 inch thick fluorosilicone (BMS 1-53) rubber gasket. Environmental conditioning was resumed on June 22.
- 2. After completion of 74 environmental conditioning cycles (see Table 7) the tank was removed from the environmental conditioning chamber, installed on the dynamic test device and load cycled at all three temperatures per Section 3.3.1. The tank was removed and leak tested. Leakage is shown in Appendix B, page B-22. There were fewer seam and corner leaks that when the tank was tested after 54 environmental conditioning cycles (see 3.9). There were 18 seam leaks and 3 corners leaking. Seam leaks at 5 psi pressure were generally very small. Two of the three corner leaks were at the end of the tank which was up during environmental conditioning. The remaining corner leak was very small. Therefore it was decided not to attempt to repair any of the leaks.
- 3. The tank was opened and examined. There was no significant change in interior appearance. Interior leakage as indicated by air at 3 psi vacuum bubbling through a water layer is shown in Appendix B page B-23. Only three leaks were visible.
- 4. Laboratory test specimens were removed from the tank and tested as before. Results were added to Table 6. Environmental conditioning was subsequently resumed the third week in August.

REVLTR D

BOEING	NO.	D3-8	29	7
SECT	PAG	E 5	7.	8

116

- *3.11 Dynamic Load Cycling of Tank No. 3 After 92 Environmental Conditioning Cycles (January 1974)
 - After completion of 92 environmental conditioning cycles the tank was removed from the environmental conditioning chamber and leak tested in water. Leakage is shown in Appendix B, page B-24. There was considerably more leakage than when the tank was tested after 74 environmental conditioning cycles. There were approximately 33 seam leaks and 6 corners were leaking. Nine of the 16 injections at the tank corners were leaking. Based on previous experience no attempt was made to repair any leaks.
 - The tank was opened and examined. Most cracks appeared to be gapped open wider than before. Interior leakage is shown on Appendix B, page B-25.
 - 3. In an effort to see if a properly repaired area would again exhibit cracks, the area shown as 1 in Appendix B, page B-25, was stripped and resealed using minimum size fillets. This area is also shown in Figure 18.
 - 4. Laboratory test specimens were removed from the tank and tested & before. Results were added to Table 6.
 - 5. The tank was placed on the dynamic test device and load cycled at all three temperatures per Section 3.3.1.
 - 6. The four corners at the end of the tank which would be down ouring environmental conditioning (R-1, R-2, R-3, and R-4) were teak tested and were essentially the same as shown in Appendix B, page B-24; therefore, no repairs were made. The tank was returned to the environmental chamber for additional environmental conditioning.

BOEING	NO.	D3-8297
SECT	PAGE	57.9

REVLTR: D

2-3033 R1

- 3.12 Dynamic Load Cycling of Tank No. , After 117 Environmental Conditioning Cycles (August 1974)
 - After completion of 117 environmental conditioning cycles the tank was removed from the environmental conditioning chamber and leak tested in water. Leakage is shown in Appendix B, page B-26. There was considerably more leakage than when the tank was tested after 92 environmental conditioning cycles. Two seams were leaking slightly along their entire length. No attempt was made to repair any leaks.
 - The tank was opened and examined. Interior leakage is shown on Appendix B, page B-27.
 - Laboratory test specimens were removed from the tank and tested as before. Results were added to Table 6.
 - 4. The tank was placed on the dynamic test device and load cycled at all three temperatures per Section 3.3.1.
 - 5. The four corners at the end of the tank which would be down during environmental conditioning (R-1, R-2, R-3, and R-4) were leak tested and leakage was not judged severe enough to repair; therefore, no repairs were made. The tank was returned to the environmental chamber for additional environmental conditioning.

BOEING NO. D3-8297 57.10 SECT PAGE

E-1031 81

118

• E

- 3.13 Dynamic Load Cycling of Tank No. 3 After 129 Environmental Conditioning Cycles (December 1974)
 - During the liquid portion of the third environmental conditioning cycle (no. 120), 11.5 gallons of fuel leaked into the environmental chamber. The tank was removed and examined. Two lower corners (F-1 and F-2) were leaking extensively. These corners were repaired with new sealant. The tank was then returned to environmental conditioning.
 - 2. After completion of 129 environmental conditioning cycles the tank was removed from the environmental conditioning chamber and load cycled on the dynamic test device at all three temperatures per 3.3.1. The tank was then leak tested in water. Leakage is shown in Appendix B, page B-28. There was considerably more leakage than when the tank was tested after 117 environmental conditioning cycles. Six of the eight longitudinal seams were leaking over much of their entire length. Fifteen of the 16 injections at the tank corners were leaking. All eight corners were leaking. The tank was opened and examined. Interior leakage is shown on Appendix B, page B-29.
 - 3. Laboratory test specimens were removed from the tank and tested as before. Results were added to Table 6.
 - 4. Testing of this tank was considered completed at this point.

BOEING	NO.	D3-8297
SE.CT	PAGE	57.11

REVILTR: E

E-3033 R1

΄ Ε

4. **REFERENCES**

Spring at the second standard and

Sh Pixel

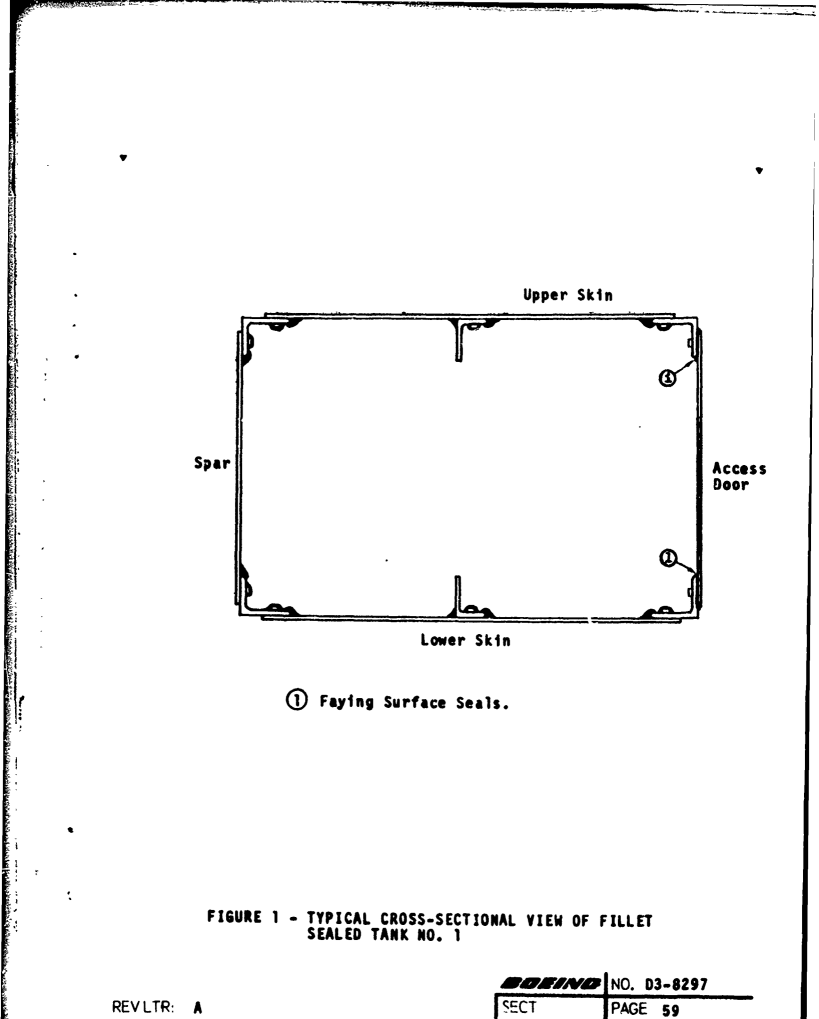
- ML-TDR-64-207, High Temperature Elastomeric Integral Fuel Tank Sealant Evaluations.
- 2. Boeing Drawing 35-22615, Tank Assembly-Sealant Test.
- 3. BAC 5504, Integral Fuel Tank Structure Sealing.
- 4. Bosing Drawing 65A14621, Tank Assembly Sealant Test.

BOEING	NO.	D3-8	297	
SECT	PAGE	58		

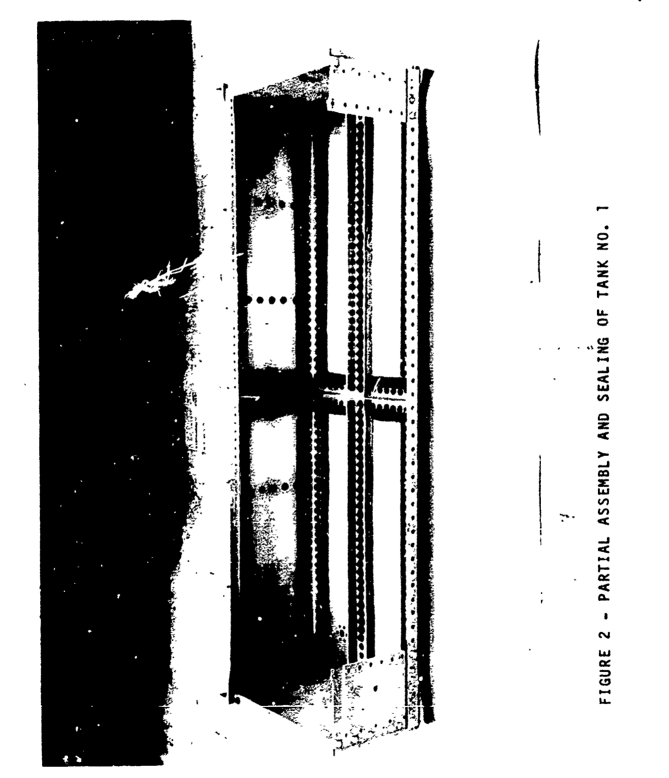
A

REVLTR: A

E-3033 R1



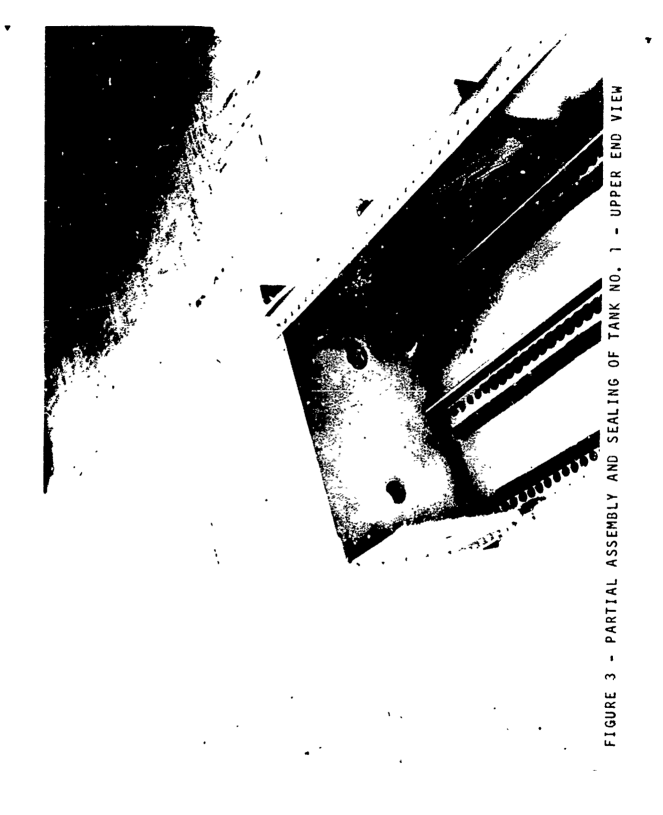
E-3033 R1

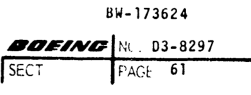


BW-173623

DEING NO. 03-8297 Sec.1 PAGE 60

E-3033 R1

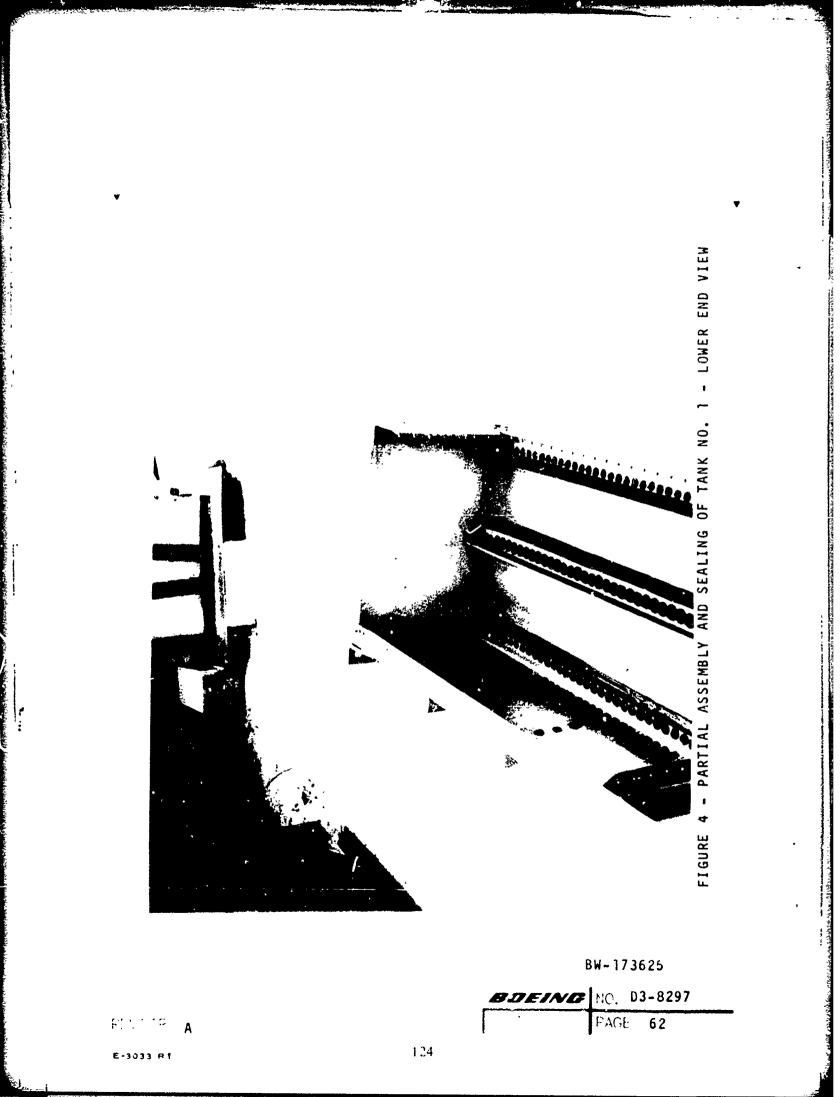


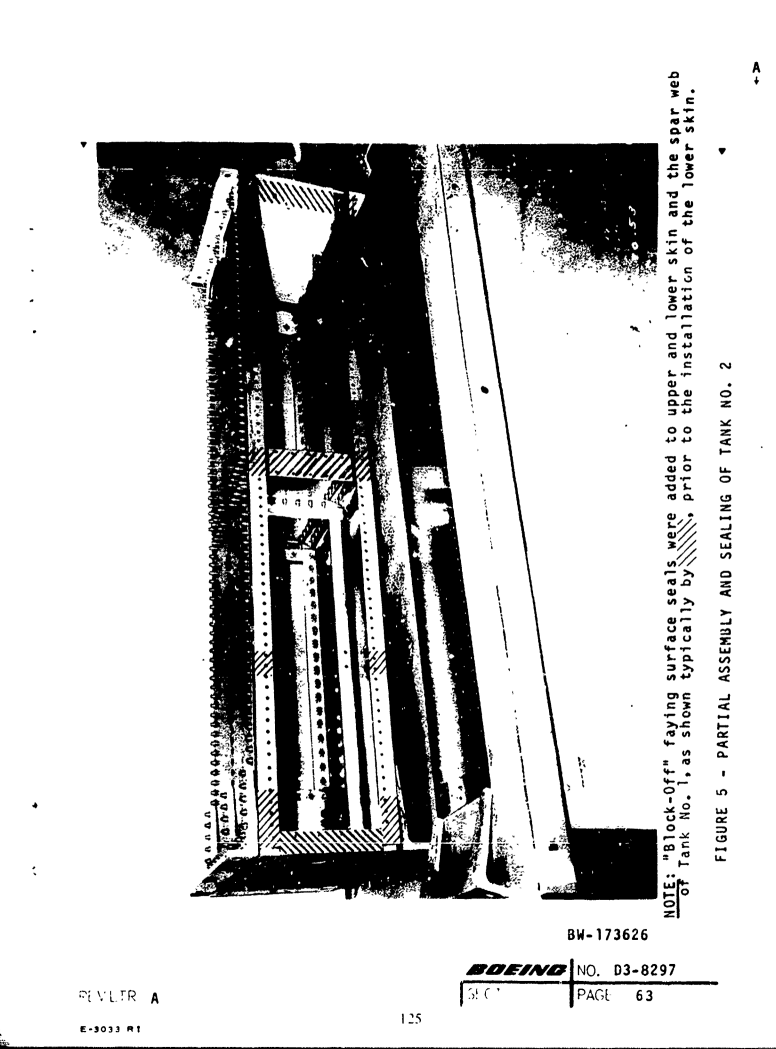


PEVLTR A

E-3033 R*

:

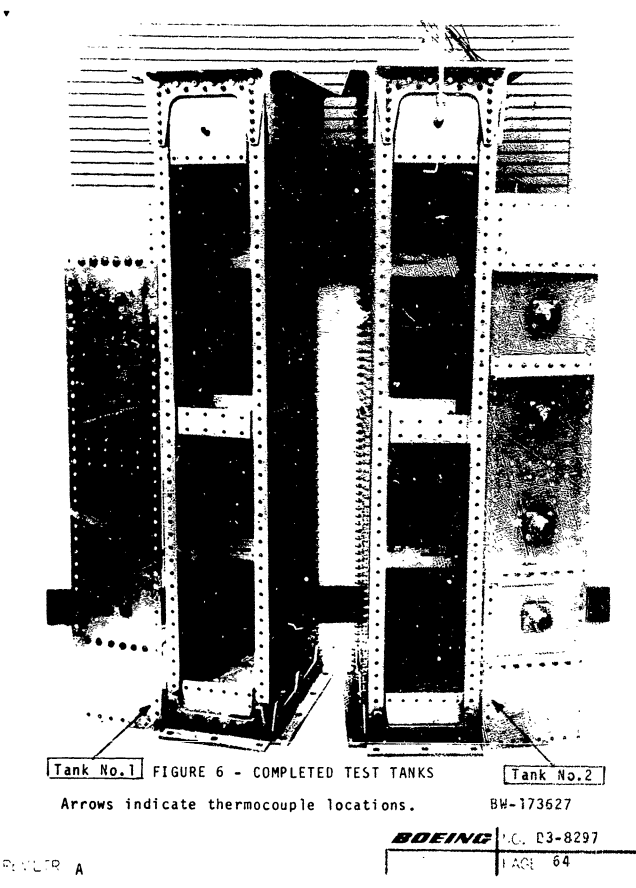




ŗ

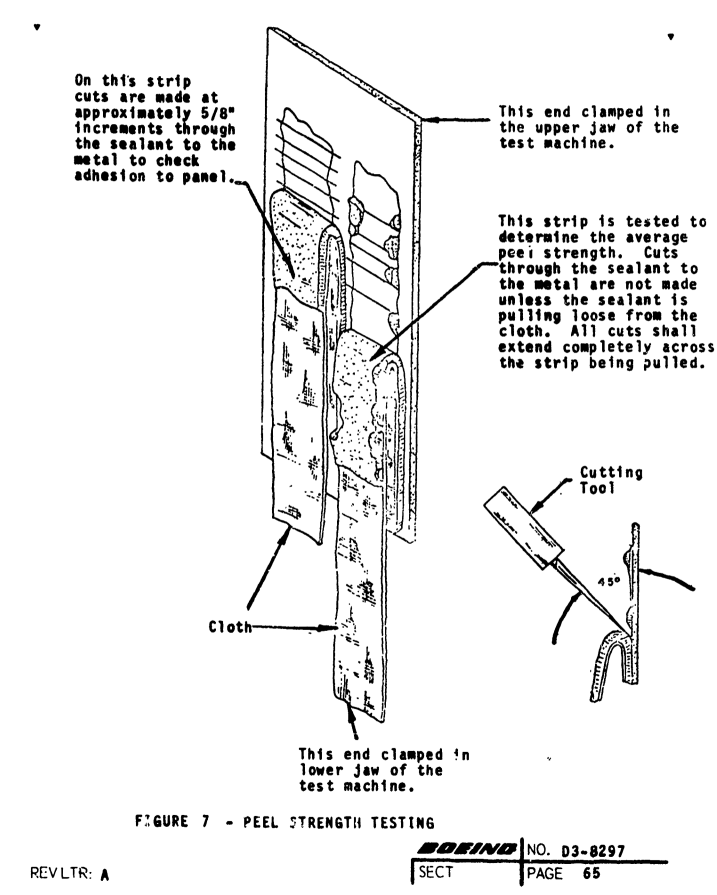
i d

K,



E-3033 R1

ľ

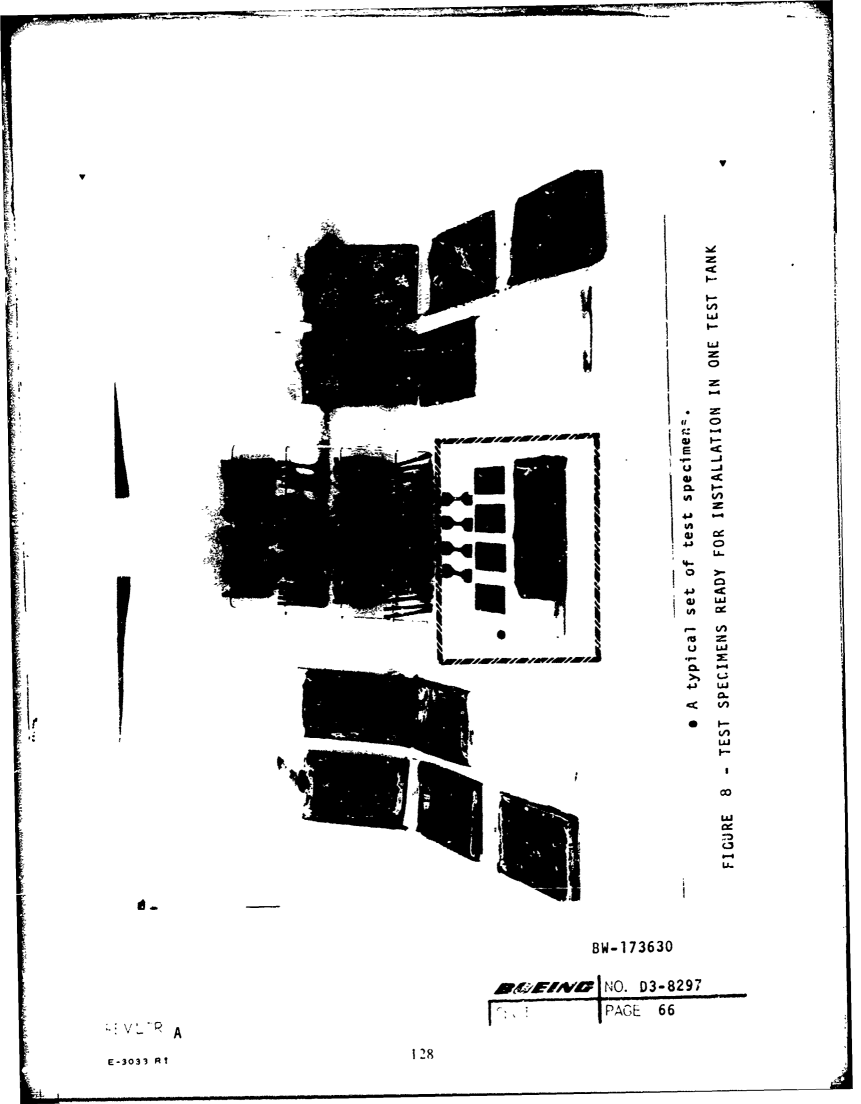


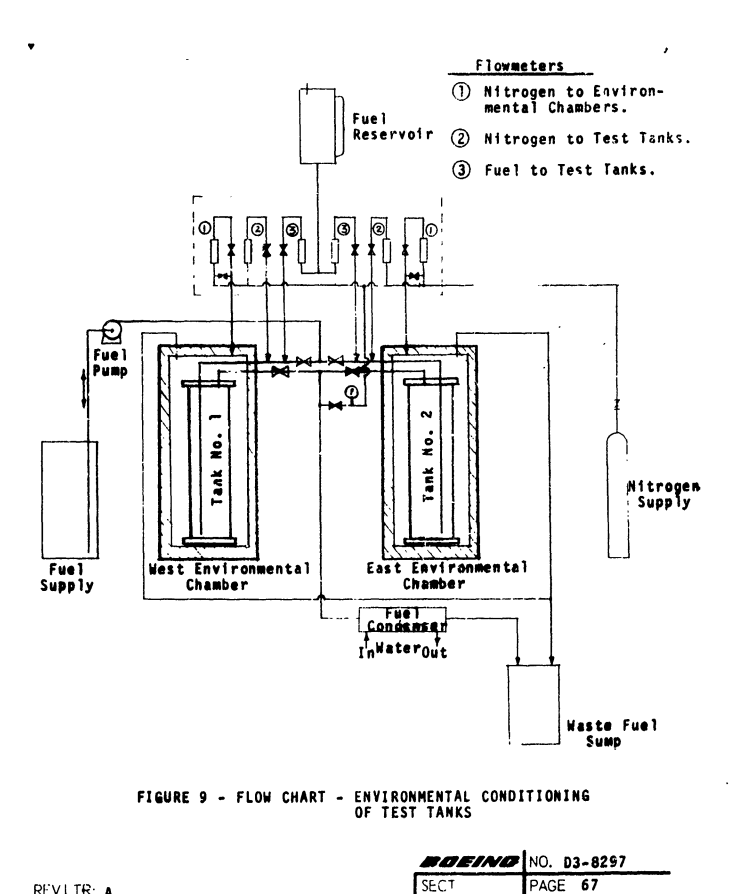
ውስት የሚያስት በማስት የሚያስት የሚያስት

そうちんとう シンチャック シスジックト

E-3033 R1

•



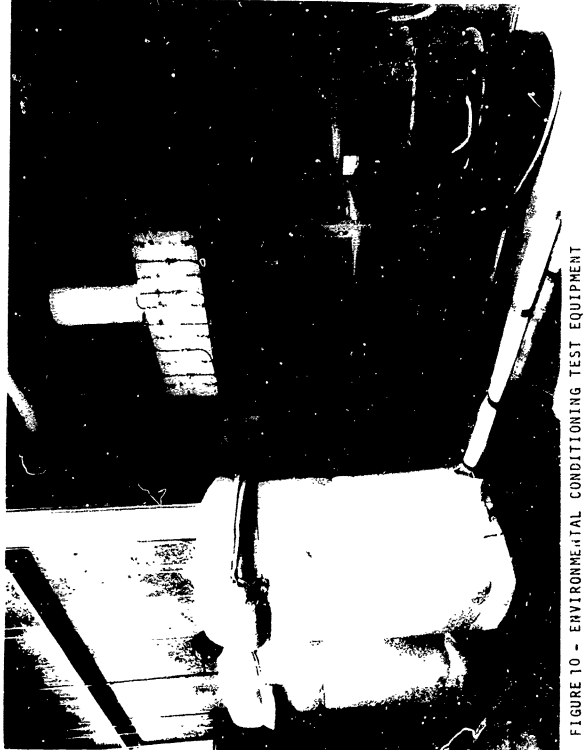


ちょう じょう たんしょう

REVLTR: A

1

E-1011 #1



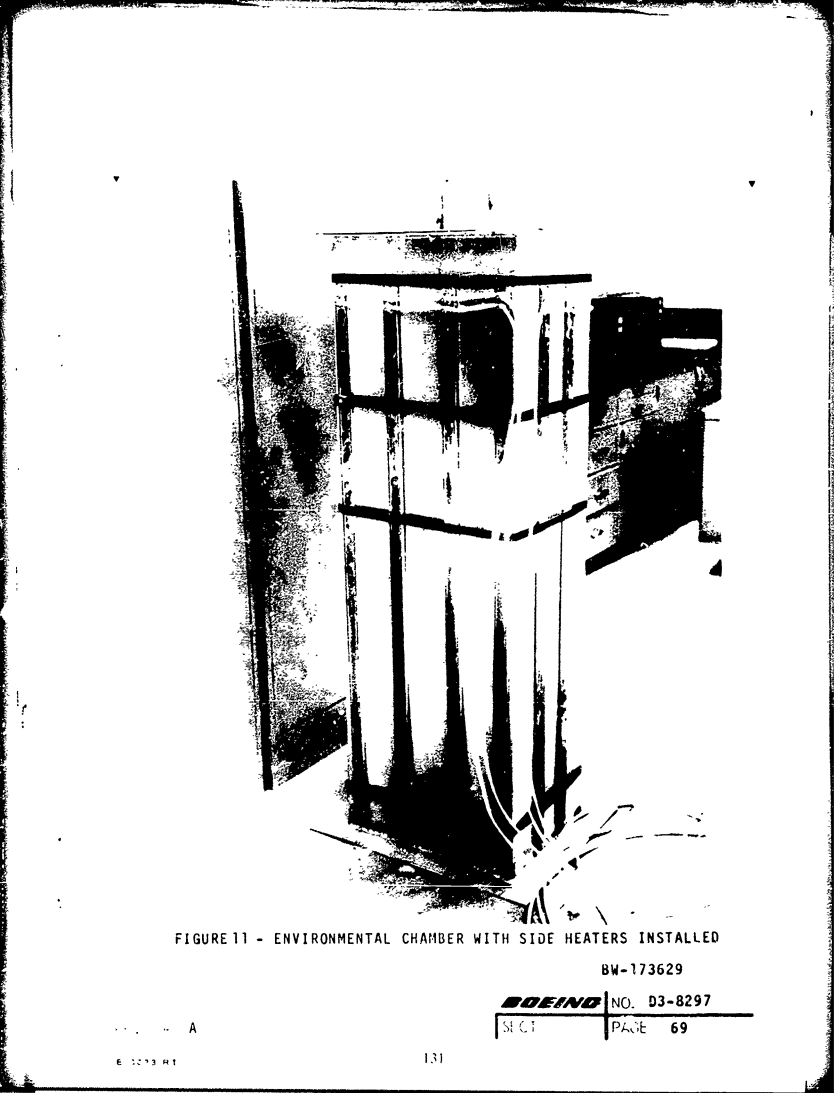
-W-1/3028

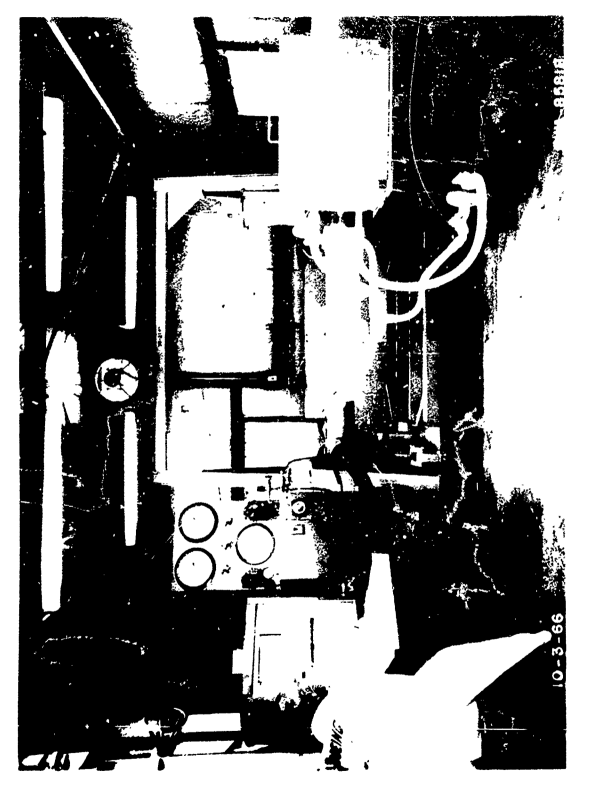
. 93-3297 58

BOEING

ί. · Υ ~ A

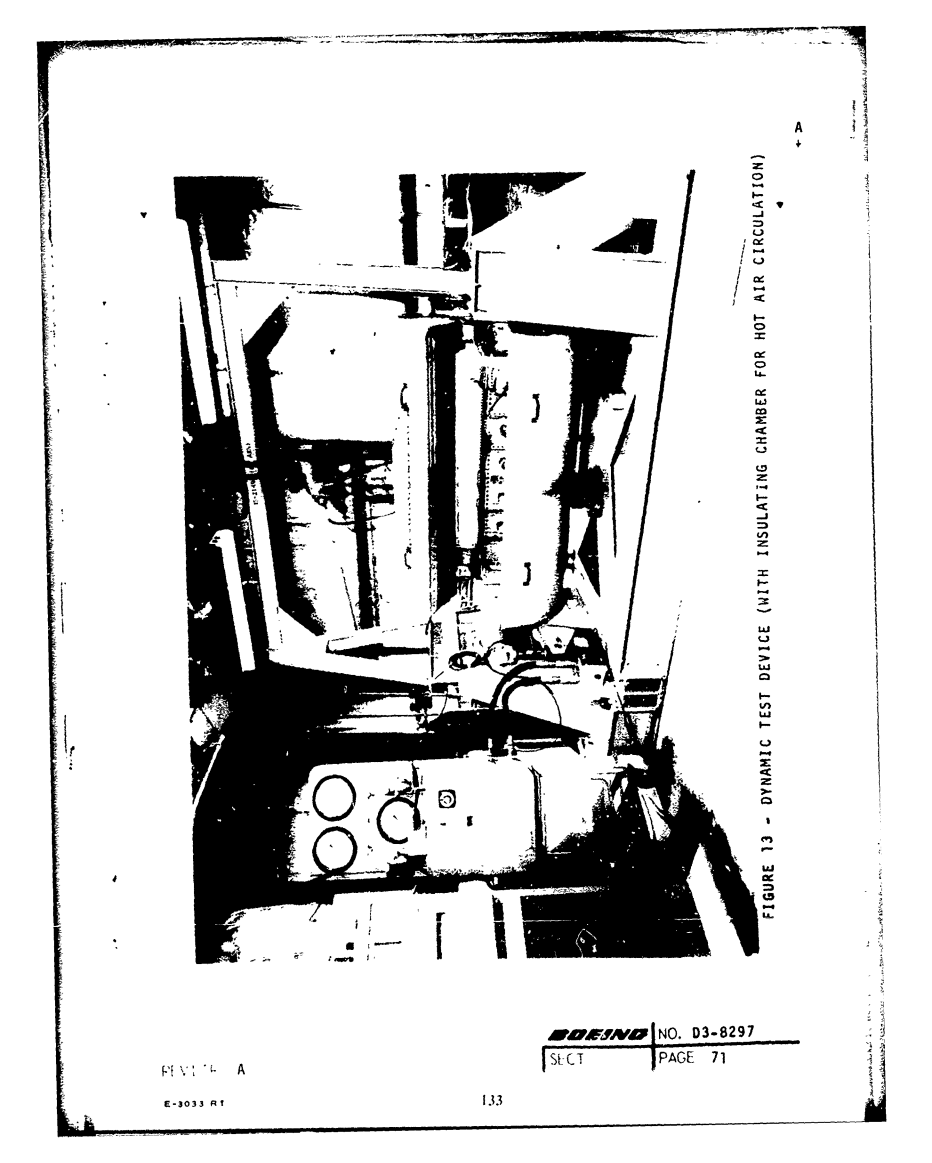
E-3033 R1

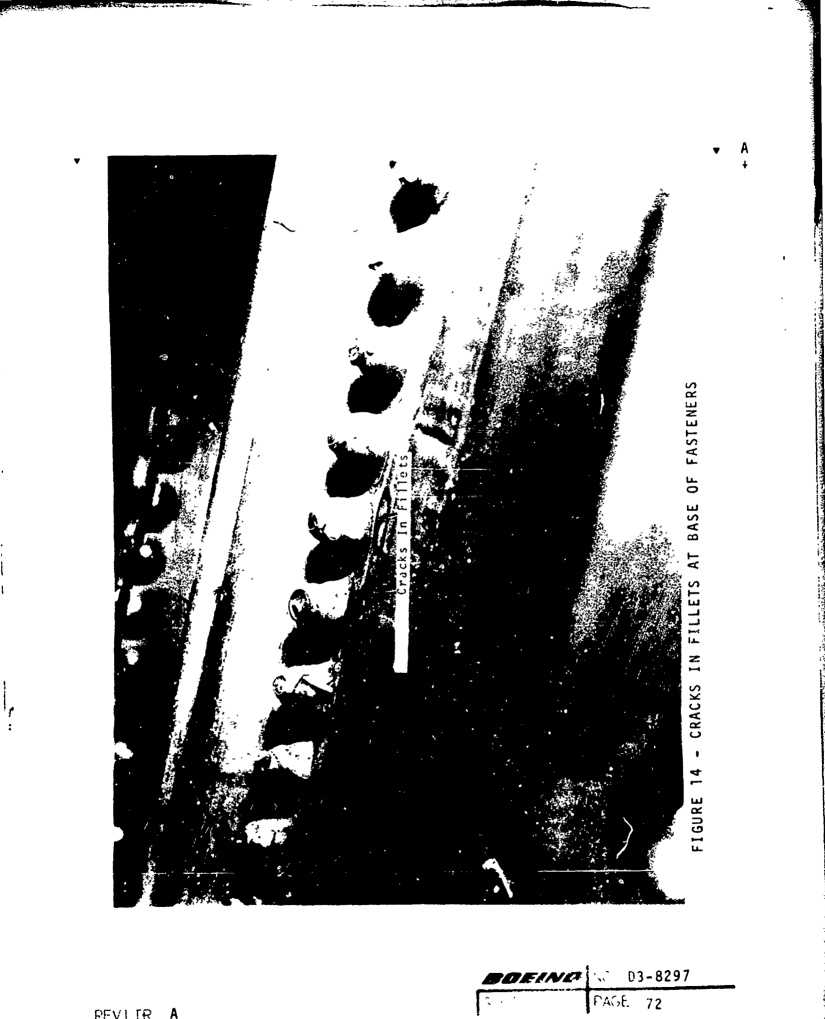




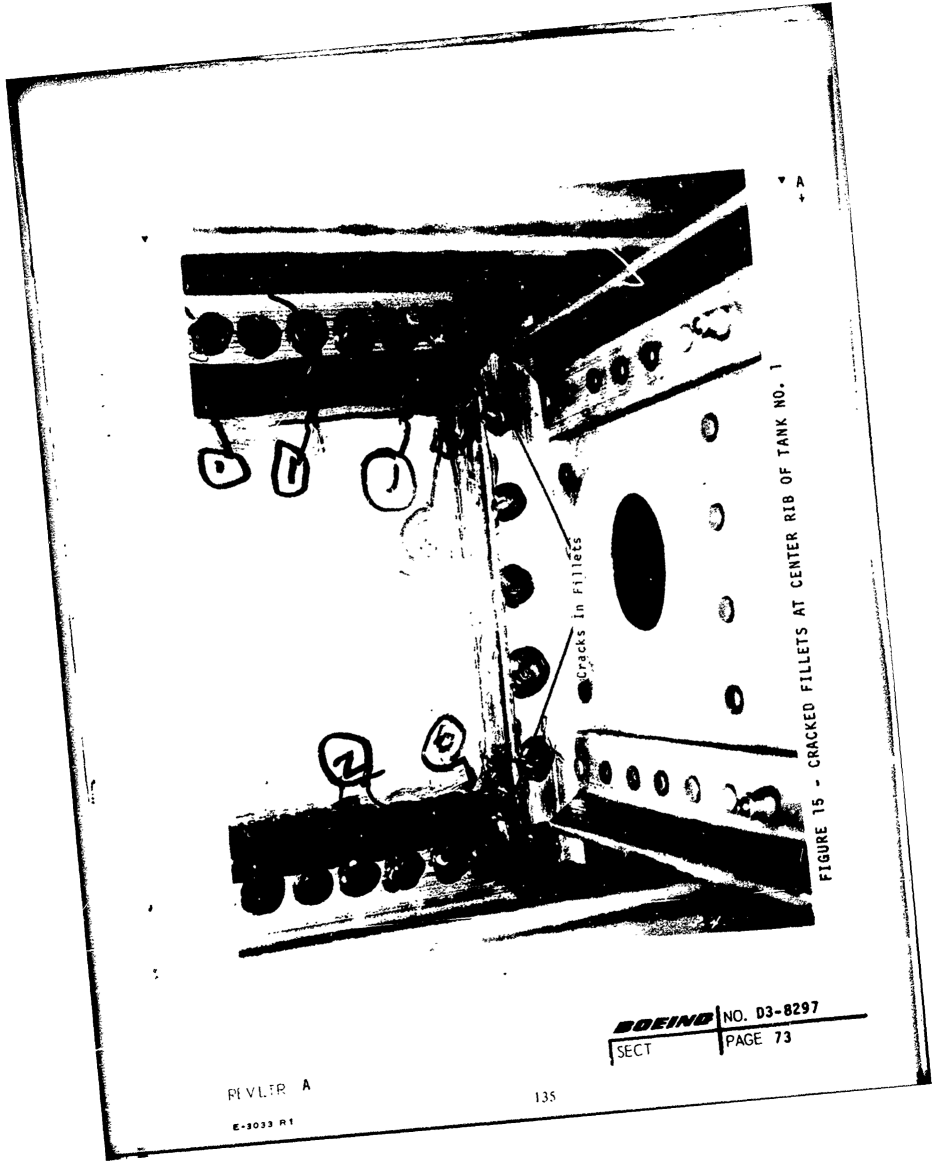
DYMAMIC TEST DEVICE (WITH CIRCULATING COLD FUEL) 1 FIGURE 12 A

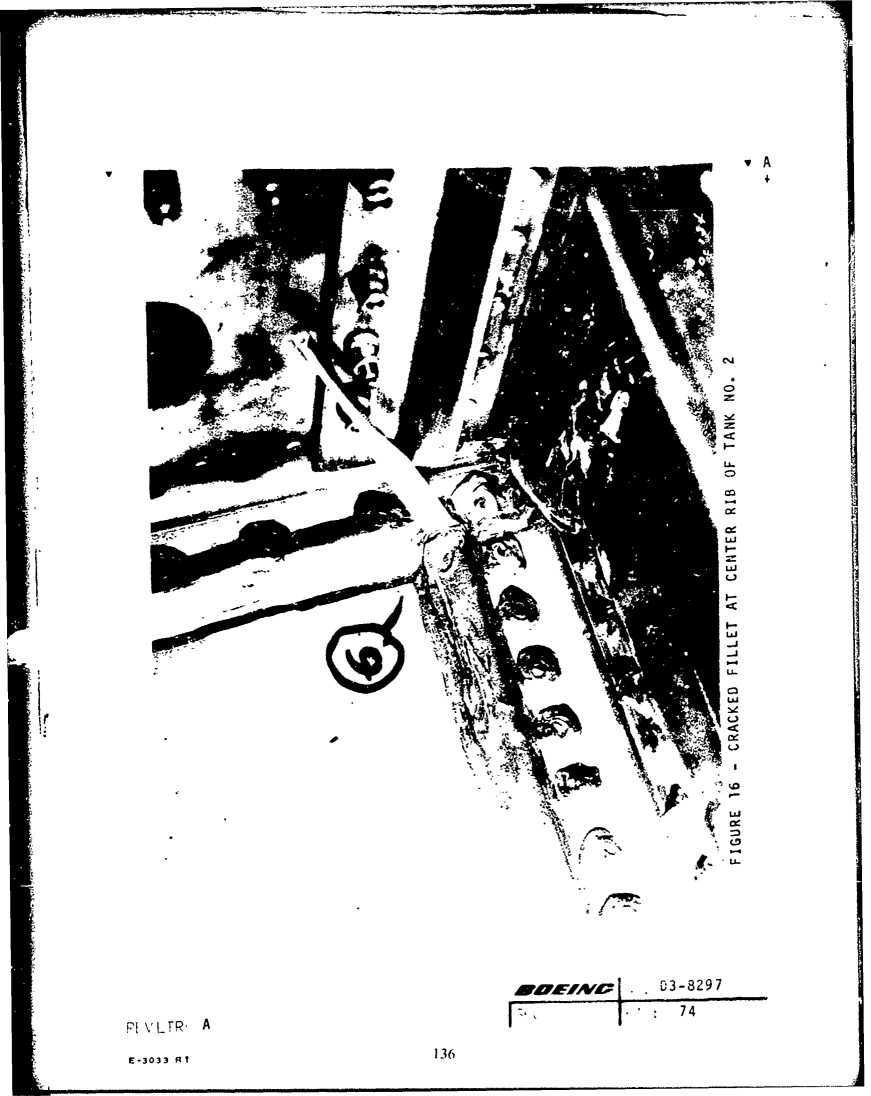
E-3033 R1

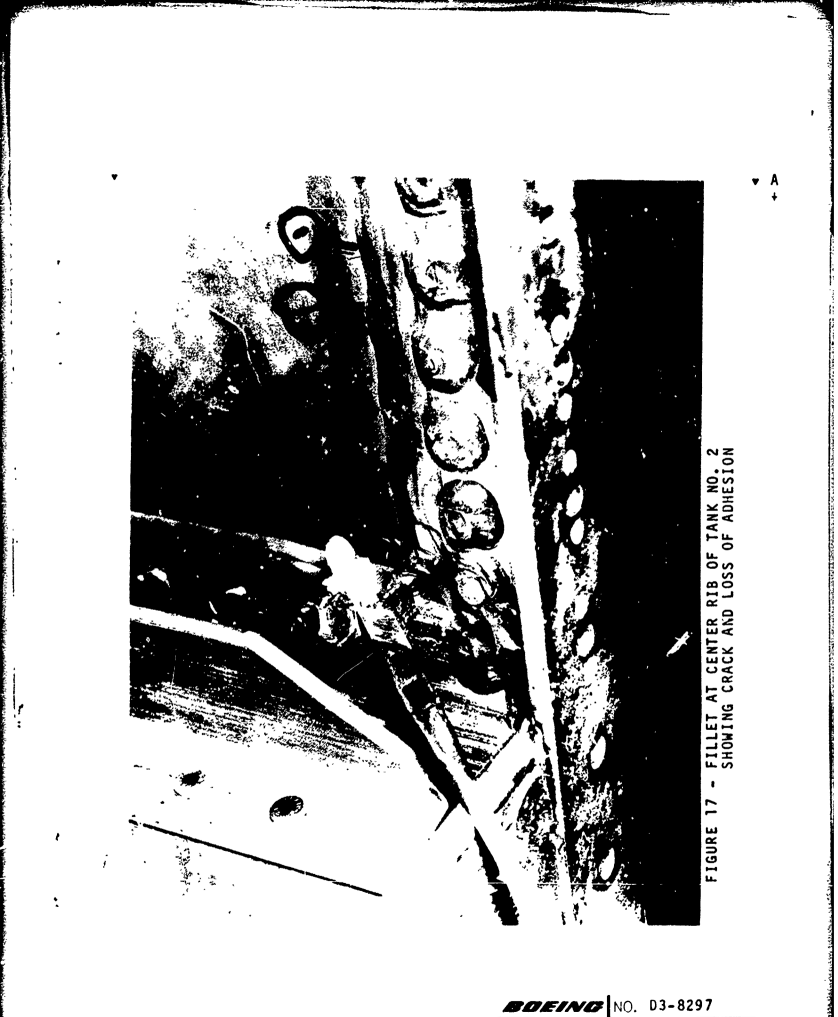




r	v	<u> </u>	17		
ۥ3	10:	33	R	1	



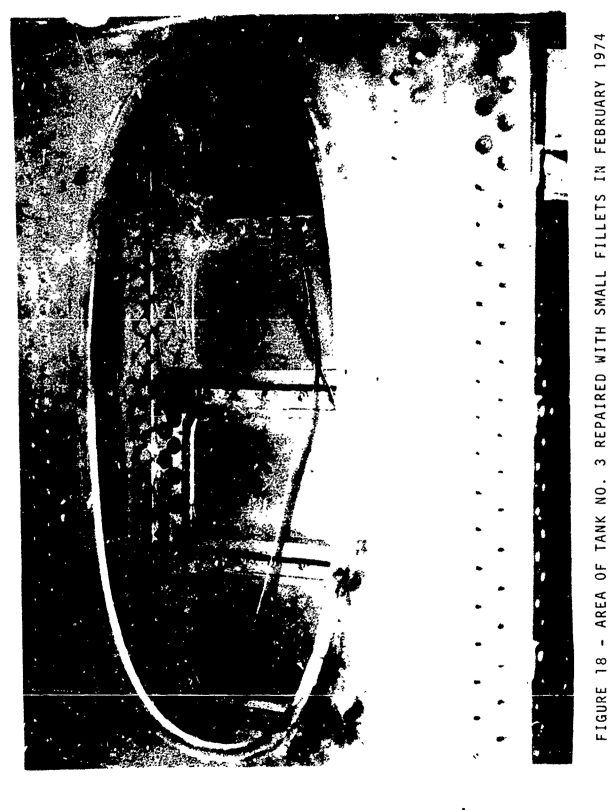




FEVER A	FE	V	L	Ţ	R	A
---------	----	---	---	---	---	---

SECT PAGE 75

E-3033 R1



₩. \/[⁻₩ Ε

BOEING	NO.	D3-8297
SI.CT	PAGE	75.1

3 RESULTS FOR TANK NUMBERS I AND TABLE PHYSICAL PROPERTY TEST

The second second

and and a set of the state of t

Strate States in

1

Completed[] 400 2.5 +0.6 -0.3 (25) Tank 476 (139) 9 0 M 117 8 90 111 1 111 No. 20 CVCLes ୭ Cycles <u>n</u> -1.3 -2.1 L) 2.+ 98 Tank No. 310 91 (25) 205 8 3 111 5 1 1 Conditioning 6 12 Cvcles 5-2 Tank -1-2 +0.2 No. 354 (130) 86 (15) 50 50 1 1 546 111 Tank 500 2.5 **6.**[+ Environmental 414 (145) 113 (55) 236 (105) Hours Hours Hours 78 10 200 60 No. Selay 524 1564 2620 665 675 +4.5 +2.9 2 S Tank No. 454 (118) 2 170 95 (20) 69 (15) **3**2 4 4 IJ Cycles Cycles Cycles 24 Hours. Specimens Exposure) (No Env. Con trol 603 (351) 277 101) 196 00 + 22 13 100 111 111 * : 111 20 ¢ Circulating Air Oven 440 ±10°F for Hours in fuel vapor at 4415-441°F 01+ ₽ +| \$1 \$ +1 \$ +1 \$÷ Υ. +| \$÷1 Test Test 9 07 75 75 75 35 75 75 Avg. (Range) (Range) (Range) pst Åvg. (Range) Physical Property Specimens per Condition) Fuel Soaked Dried Out2 Lb./In. Avg. Separation Elongation, % Avg. (Range) pst Avg. (Range) pst Avg. (Range) Avg. Hardness, pts Fire 1-Soaked Avg. Avg. Initial Dried-Out Strength. Cohestve × 24 Strength. Change, Change . Durometer Ultimate Tensile × **Height** Volume Peel ୦ Э 3

REVLTR: C

D3-8297 DEIND NO. 76

SECT

E-3033 R1

139

PAGE

Conditioning

4 cycles of Environmental

first

2 during the

Thise specimens were in Tank No.

One Panel per Condition

All specimens immersed in liquid fuel at room temperature for one day prior to testing.

TABLE 2

ļ

.

ENVIRONMENTAL CONDITIONING CYCLES FOR TANK NO. 1

ienta l	Liquid F	uel Exp	osure	Hours		Fuel Va	Por	sure	2	
E Y		140		em D.	Ambient	401	4]5	4 26	Over 1	Temp. ()
-	39°F	150°F	15] - F to + ° F	Hrs.	400°F	425°F	441°F	441°F	to + °F	Hrs.
	169.0				16.0	18.0	133.0		457	.4
	6.0	16.0			5°U		138.5			
	1.6	20.0					138.5			
	5.0	10.5	152	6.0			114.0			
	16.5	2.0			18.5	5.0		165.5	443 445	4.0
	1.0	21.N	152	2.0	8.0	3.0	10.Û	124.8	444	6.5
		15.5	175	5.0	3.0	1.0	5.0	113.0		
		17.0	160	4.0	5.5	1.5	11.0	128.5		
		15.5	160	5.5	28.5	1.3		0.06	443	~.
	2.0	13.5	165	и.U	4.5	1.0	19.0	146.5	443	5.0
	1.5	20.0			6,5	1.5	21.0	90.8	450	-, -
	2.0	22.0			25.5	19.0	34.5	80.0	445	6.5
	5.0	16.5			3.5	1.8	9.5	105.8		Í
	16.5	5.0	160	2.0	52.5	3.0		89.5		1
	1.0	17.0	157	3.0	£3°3	3.8		115.3	4 46	1.0
	2.0	17.5	i 75	z•0	3.3	1.3	ż.0	140.0	448	·2.
	1.0	18.5	175	3.0	3.0	.8	1.0	140.8	443	
				d in	ur s	reported at	415-441°F	P C L	426-441	
5	using fuel	el from	Standard	110	of California	nia.				

140

REVLTR: A

E-1033 R1

D3-8297 **DEIND** NO. PAGE SECT 77

(Continued on next page)

(Continued) C YCLES 2 TABLE CONDITIONING

1

t

•

ature A ŧ ure ۰. at ambient temp columns are due 0.5 4.0 50.5 8. C 0.3 temper Hrs. em p --U. ₩. • 4 This was a id day exposure at ambien 5 Most of the hours in thise columns a unplanned cool downs raused by over controls turning heaters off. Over 442-457 1211 443 446 445 4.5 5 t t Hour 44]°F œ. 0 304.5 93.01 92.0 0. -S. m 0 5 S. 5. 5. S **4**26 to Exposure 3862, ٠ 138 140. 173. 140. 00 94. 158. 370. 101 187. 237 441°F 415 63.7 2 Vapor 2 FOR TANK 401 to 425°F 5.5 117.7 3.3 0.3 5.0 1.3 1.8 ي. بد 2.5 ŝ ŝ • 0 S 0 Fuel 2 9 -0 v Q ~ 30,0 140°F 200°F 37.0 ō S ŝ 5. 0. S 17.0 0 Ambient to 400°F 65. °. 2.5 18 32 45. 5. 30 20 37 ŝ (J 824.1 **N** S 4 4.0 80 5 . 0. -un . 140 F rer . 52. v ٠ 29, 42. 69. **b** 19 ω 35 13.0 0.5 С Hrs. 53.(Temp. hours from 5 Hour S°FI S°FI LL. • • l' l' Hours in this column are included in h reported at 415-441°F or 426-441°F. Startstusing short liquid fuel cycle here on due to excessive leakage. Over Exposure 21 175 151-17 175 to excessive leakage. **ENVIRONMENTAL** 50°F 0**4**0 7.5 • 17.5 290.5 5 18 Fuel Ambient Liguid to 139•F • 3**36.**0 584.1 3.0 •• 1.0 2.5 1.0 • S. s. 1.0 ۰. .. ഹ <u>__</u> ~ _ --------nvironmental onditioning No. TOTALS (m Cycle 22 19 23 18 20 24 25 26 28 29 30 3 2] 27 $\overline{\mathbb{C}}$ ୦ ROEINO NO. D3-8297 SECT PAGE 75 **REVLTR:** A

34

「ない」日本なるかです。

E-\$033 R 1

141

	ENVIR	DIMENTAL	C OND I T I OI	VING OF	ENVIRONMENTAL CONDITIONING OF TANK NO. 2		
Environmental	Liguid	Fuel Ex	Liquid Fuel Exposure Hours	urs	Fuel Vapor	Exposu	re Hours
Conditioning	Ambient	0+1	Over T	em p .	Ambient 401 415	401	415
LYCIE NO.	to 139°F	to 150°F	150°F to +°F	Hrs.	400°F	to 425°F	to 441°F
-	169.0				16.0	18,0	133.0
2	6.0	16.0			2.0		138.5
e	1.6	20.0					138.5
4	2.0	9*5	155	10.0			114.0
Total	178.6	45.5	150-155°F	10.0	18.0	18.0	524.0
		234.] Hours	ours		56	560 Hours	

TABLE 3

ţ

1

REVLTR: A

BOEINO	NO.	D3-9297
SECT	PAGE	79

K-3032 RT

142

A +

•	TABL LEAKAGE OF TANK	NUMBERS 1			•
	AFTER FOUR CYCLES OF ENV	IRONMENIA	Number 0		
Tank No.	Location Of Leakage*	Envir	Cycles Of onmental tioning	After 4 C Environm Conditio Pl 510 Hot L	ental ning
				Per Sect	10n 2.4.3
		Joint Leaks	Fastener Leaks	Joint Leaks	Fastener Leaks
1	<u>Primary Test Areas</u>				
	Upper Skin	0	8	0	11 ()
	Lower Skin	0	2	0	7
	Spar	0	1	0	4
	Secondary Test Areas				
	Access Door	8	0	Many	Many
	Ends Of Tank	=13	2	11	29
2	Primary Test Areas				
	Upper Skin	0	2	14	20 (2)
	Lower Skin	0	3	0	5
	Spar	0	1	1	=26
	Secondary Test Areas				
	Access Door	6	12	Many	Many
	Ends Of Tank	10	3	11	28
	-8 leaks in one area.	2 •!	j leaks in	one area.	-L
-	*See Appandix A	for speci	fic locatio	ns of leak	s. 8297
REVL	TR: A		SECT	PAGE 8	
E-305	3 A 1	143			

1

44

נות אלאר לאות הלאות האלא האלי של אר שלה לאות אולה אלי היות היות אלי יישר אלי היות אלי אלא לא אלי אלי אלי אלי א מיני ביימי היות היות היות היו ביווי היווי היו מים מימי אות יות מית אלי מים ביוו ביר אלי אלי אלי אלי אלי אלי אלי

10.4.4.10

1

TAB	LE	5
-----	----	---

A

SUMMARY OF LEAKAGE OF TANK NO. 1 IN PRIMARY TEST AREAS

ſ

	Number				us Cumul Leak Loc		-
			Leak	age Tes	t No.*		
Type Of Leak	î	2	3	4	5	6	7
Fastener Leaks	11	22	26	12	80	89	63
Joint Leaks (Total)	0	0	0	4	10	3	4
Joint Areas Leaking ①	0	0	6.	4	7	8	6

Leakage	Environmental Conditioning	Total Fuel Vapor Exposure	Cycle	Dynamic L es Complet st Tempera	ed	
Test No.	Cycles Completed	At 415 - 441°F	440°F	-50 F	Ambient	1
ı	4	524 Hours	0	0	0	
2	4	524 Hours	510	0	0	
3 (2)	4	524 Hours	510	C	0	
4	4	524 Hours	510	510	510	
5	4	524 Hours	560	510	510	
6	12	1564 Hours	560	510	510	В
7	12	1564 Hours	1070	1020	1020	В

- (1) There are 24 joint areas, separated by faying surface seals, in the primary test area (8 each on top skin, bottom skin, and spar).
- (2) Same exposures as Leakage Test No. 2, except after repair of loose and/or cracked fillets.
- (3) Eight leaks in four joint areas, plus continuous joint leakage along entire lengths of four additional joint areas.
- (4) Eight leaks in five joint areas, plus continuous joint leakage along entire lengths of two additional joint areas.

ROEINO	NO. D3-8297
SECT	PAGE 81

REVLTR: i	3
-----------	---

E-2033 Rt

5

and a second second

and a second second

. .

,

ntrakania 24 J

ي م د واستان ويور الي م

S. S. Balanta

Environ	onmental Cycles Cycles Cycles	(90) (4) (67(198)556(33)429(243)254(107) 104) (5) 197(90) 54(44)113(51) 101(20) 159) (5) 2058(308)	$ \begin{array}{c ccccccccccccccccccccccccccccccccccc$	7 10% 80% 6.5 80% 100 6	-0.7% -1.7% -3.9 -3.4%	-1.5% -2.2% -4.0 ⁻	(2) ③ 45 46 42 43	F.EOnly 3 Specimens Per ConditiF.ENot true cohesive separation was adjacent to wire screenSpecimens.7At 438°F.
rol nens		667 197 2058	150 150 231	7	':	• 1	<u>ن</u> ق	Specimens.
Test	emp.	75 450 -50	75 450 -50	75 75	75	75	75	Per c
hysical	Prope rty	Tensile Strength, psi Avg. (Range)	Ultimate Elongation. % Avg. (Range)	Peel Strength, lb./in. Avg. % Cohesive Separatio	Volume Change, % Avg.	Weight Change, % Avg. Fuel Soaked	J rometer, Haidness PTS Avg.	 Temperature Toler Per Table 7. Average and Range Only 2 Spectmens
	rnys i ca i	Property	enstle S psi	Property Strengt Avg. (* Avg. (Tensile Strengt Tensile Strengt psi Avg. (Ultimate Elonga Voltimate Strength, Peel Strength, Cohesive Sep	Property Property psi Avg. (% Avg. (% Avg. (% Avg. (1b./in ohesive Sep me Change,	Tensile Strength. Tensile Strength. Property (Ra Ultimate Elongati Peel Strength. Ra Peel Strength. Volume Change. % Weight Change. %	Tensile Strength. Tensile Strength. Instrength, kaye. (Ra Ultimate Elongati & Avg. (Ra Ra Peel Strength, kaye. Volume Change, k Weight Change, k Height Change, k Proneter, Haldne, PTS, P

ì

- ALINA BALKIN

· · Number

ł.

The second

. . . .

- ----

601.00

SF-CI

82

EAG

TABLE 7 Environmental conditioning of tank number 3

كندحاله عاوتك سنادها

1

		Fuel Liquid	EX	posure h	Hours		Fuel	l Vapor	Exposure	e Hours		
	Environmenta	Ambient	140	Over Temperati	0ver mperatures	Below	140	104	415	426	0ver Temperatures	tures
	Conditioning Cycle No.	139°F	150°F	151°F To +°F	Hours	Θ	- 6	425°F	44	441°F	441°F To +°F	Hours
•	-	4.0	15.0				19.5	5.5		100.0		
	2	18.5					3.3	2.5		115.8		
	E	0.4	16.5			16.0	32.0	7.5		92.5		
	+	5.0	12.5			36.0	35 . 0	14.5		227.0		
•	5	2.5	12.0	155	8.0	2.0	4.5	7.:		156.0		
	ç	3.0	12.5	156	6.0	58.0	32.0	ó.(75.0		
•	7 (3)	5.5	15.0			35.0	53.5	14.5		288.5 _a	460 445	1.0
•	80	4.0	18.5			7.0	35.5	9.5		95.5		
•												
+ 	Aost of	of the hours temperature	t con t	these c rols tu	columns turning o	are due to off heaters	•	unplanned cool	ol downs	caused	bу	
				1. ()		el ebim	, et a t a	coated with	th polvo	oolvohenvlouinoxalir ^e	noxaltre	
- 82	A NEW O	installed p	rior to	inig se di ni	ning th	prior to beginning this cycle.						

3 These hours are included in hours reported under 426 - 441°F.
4 Lower half of tank may have been in the range of 401 to 425°F for 50 hours.

(Table Continued on next page)

and the second second

1

REVLTR· B

8297 83 SECT PAGE

146

E-3033 R1

A +

Environmental Cycle Ko. No. 10 11 12 13 13 13 13 15 15 15 15 18 18 18 18 20 20	Fuel Li Fuel Li 70°F 139°F 139°F 5.5 3.0 5.0 4.0 4.0 4.0 8.0 8.0 2.5 8.0 10.0	quid Exp 140 150°F 150°F 19.5 19.5 19.5 18.5 18.5 18.5 18.5 18.5 13.5 13.5 23.5 23.5 11.0	posure.Ho Tempere.Ho 151 cf + F + F 	e, Hours Over to Hours Aours	Below 140°F () 87.0 87.0 15.0	Fue 140 140 140 400°F 400°F 33.0 33.0 8.0 6.5 6.5 6.5 6.5 6.5 6.5 6.5 6.5 6.5 8.5 8.5 7.5 8.5 8.5 8.5 8.5 8.5 8.5 8.5 8.5 8.5 8	l Vapor 401 10 441°F 3.5 3.5 3.5 2.0 2.0 2.5 2.5 2.5 2.5 2.5 2.5 2.5 2.5 2.5 6.5	Exposure 415 441°F 441°F	. Hours 426 70 162.0 162.0 161.0 138.0 138.0 138.0 137.5 135.5 135.5 135.5 135.0 120.0	
22	10.0	10.0				6.5	3.0		167.0	
TOTAL SUST	1 3 0	217 0	151 156		256.0	0 0 1 4				

`**!**

and a second second

.'

ţ

ي جانبي ماريد المارين المارية ا ماريخ

. . .

÷ :

1.141.141.1

		iture	Hours							3.5										
		Over Temperature	441°F to↓°F							446										Ъу
	Hours	4 26 to	441°F	192.0	108.0	139.5	115.0	257.5	120.0	96.5	105.0	164.5	141.5	130.0	141.0	141.5	196.0	118.5	112.5	caused h
	Exposure,	415 to	441°F							63.5							15.0			l downs
0.3	Vapor Ex	401 to	425°F	7.5	5.0	3.0	2.0	4.5	4.0	43.0	2.0	2.0	1.5	1.5	1.5	2.0	3.5	7.0	2.0	nned cool
ING OF TANK NO.	Fuel '	140 to	400°F	49.0	58.0	5.5	7.0	29.0	39.0	65.0	14.0	2.0	4.0	9.5	5.0	5.0	55.5	50.5	4.5	e to unplanned off.
O SNINOI		Below 140°F	[]	42.0																a O
CONDITIONING	e,Hours	ture	Hours										3.0 19.0	14.0		10.0	12.0	13.0	10.01	columns are du turning heaters
ENVIRONMENTAL	Exposure, l	Over Tempera	151 to ↓ °F										205 175	175		160	170	170	160	e
ENVIRON	Liquid Exp	140 to	150°F	8°.5	18.5	17.0	18.0	15.5	16.0	5.0	17.0	18.0		6.0	18.5	8.0	5.5	10.0	8,0	s in cont
	Fuel Lig	Anbient To	139°F	11.0	3.0	3.0	3.0	5.0	3.0	17.0	2.0	5.0	1.0	1.0	2.0	2.0	2.0	1.0	2.0	Most of the hours over-temperature
	Environmental			23	24	25	26	27	28	29	30	31	32	33	34	ທ ຕ	36		38	(-)
	REVI	TR:	D										SECT			-	GE	- 8: 8:	29 3.1	

determinent of the other

a a ser a construction de la construcción d

J. L. P. J. Methods of L. S. M. S. Mathematical Social Social

and the subscription of the

i e

ţ.

3 These hours are included in hours reported under 426 - 441°F.

(Continued) TABLE 7

and the second

and and the state of the second

والمعالية المحالية الم

والمتعادية والمتعادية والمتعادية والمتعادية

ŧ

Hours 14 I Temperature \bigcirc 41 leakage Over 441°F to+°F 448 445 liquid Most of the hours in these columns are due to unplanned cool downs caused by over-temperature controls turning heaters off. 441°F 93.5 137.0 87.0 171.0 135.0 123.0 85.0 131.5 ഗ ഗ 131.5 163.5 121.0 133.0 144. **4**26 to 138. Hours to excessive 8. 441°F Fuel Vapor Exposure, 415 t t Section duc to 425°F 5.5 1.5 5.0 S 1.5 0.3 1.5 2.5 4.0 ഹ 0 0 ഗ ഹ . M 2. . M 401 n M sse exposure CONDITIONING OF TANK NO. 19.5 400°F 8.0 30.0 5.0 8.0 12.5 28.5 42.0 3.0 10.5 33.5 26.5 ŝ ഹ 7 (Continued) 10. . 6 t t 140 ન vapor . Below 140°F 29.5 23.0 1.5 (no E Hours TABLE 18.2 Fuel Liquid Exposure, Hours **Cemperature** Over с то то ENVIRONMENTAL 190 51 + 150°F 17.5 17.0 0 24.0 23.5 8.5 10.0 ഗ 22.0 ഗ 11.0 S 18.5 ഗ 12.5 19. 19. 12. 3. 8 t t 140 Ambient TO 139°F 5.0 1.5 1.0 11.0 3.0 1.5 11.5 14.0 3.0 11.0 13.0 S S ຽ . . Environmental onditioning Cycle 40 39 43 64 41 41 4 S 46 50 44 47 48 S 52 53 No. 7 BOEING 1)3-8297 NO. SEC 7 PAGE REV LTR: 83.3 D

E-3033 R1

Sun, Some

مى مە^{مىر}ىدىكىر<mark>مەمىرىد</mark>. مەمەر مەمەر مەمەر مەمەر

; .

.

· when he was to start a

and a second a substantial

ちちしているいろうちちというという いいち いっちいちいちょうちょうちょう あんにちに

2

1

and the second second

149

- 441°F

These hours are included in hours reported under 426

 \bigcirc

	·	r	T	1	1	<u> </u>	_	1		r	1	<u>.</u>	.	1			r	T	-1
		ture	Hours																
•		Over Temperature	441°F to+°F																
	Hours	426 to	441°F	87.5	137.5	156.5	133.0	133.5	132.5	131.5	132.0	132.5	141.0	133.5	130.0	131.5	133.0	132.5	caused by
	Exposure, H	415 to	441°F																duwns cal
NO. 3	Vapor Exi	401 to	425°F	3.5	3.0	2.5	3.5	3.0	2.0	1.5	2.5	2.0	1.5	1.5	2.0	2.5	1.0	2.0	ed cool
OF TANK N	Fuel V	140 to	400°F	38.5	3.5	10.0	9.0	0.6	10.0	9.5	0.6	0.6	9.5	9.5	10.5	9.0	10.0	8.5	unplanned
CONDITIONING O		Below	140°F																e due to
CONDIT	ours	ture	Hours																mns are
MENTAL	Exposure, II	Over Tempera	l51 to ↓ °F																se colu
ENV IRONMENTAL		140 to	150°F	14.5	10.5	14.0	15.0	18.5	13.0	·17.5	14.0	12.5	15.0	15.5	15.0	19.0	15.0	16.5	in these colu
	Fuel Liquid	Ambient To	139°F	8.5	14.0	0°0'	8.0	6.0	10.5	7.0	9.5	11.0	0.0	8.5	8.5	6.0	0.0	<u>8</u> .5	e hours
	Environmental Conditioning	Cycle No.		54	55	56	57	53	59	60	61	62	63	64	65	66	67	6.9	() Most of
רי	VLT	R: D										-	C1	5810	E	NO. PAG	.)3 E	829 83	
17.1. 									15	50		I						-	

TABLE 7 (Continued)

And a second second

مى بىلىغۇر بىلىغۇرىيىتى بىلىغۇرىيى بىلىغۇرىيى بىلىغۇرىيى بىلىغۇرىيى بىلىغۇرىيى بىلىغۇرىيى بىلىغۇرىيى بىلىغۇرىي بىلىغۇر بىلىغۇرىيى بىلىغۇرىيى بىلىغۇرىيى بىلىغۇرىيى بىلىغۇرىيى بىلىغۇرىيى بىلىغۇرىيى بىلىغۇرىيى بىلىغۇرىيى بىلىغ

1

ŧ 1 1 over-temperature controls turning heaters off. \odot

These hours are included in hours reported under 426 - 441°F.

1.040.4

•		Over Temperature	441°F Hours tot°F (3)	>							_								
	Hours		441°F 4	158.5	115.0	120.0	181.0	81.0	114.0	32.5	135. `	111.5	127.0	114.0	98.5	137.0	62.5	131.5	caused by
	Exposure, H	415 to	441°F																downs cau
NO. 3	Vapor Exp	401 to	425°F	2.5	2.5	2.5	3.0	2.5	2.0	5.0	2.5	2.0	3.0	2.5	3.0	2.0	2.5	2.0	cool
×	Fuel V	140 to	400°F	6.0	14.0	8.5	6.5	7.0	22.0	56.5	C.7	6.5	14.0	6.5	15.0	6.5	6.5	6.0	unplanned
TABLE 7 (Continued) CONDITIONING OF TAN		Below	140%										15.5		21.5		71.5		e due to
TABLE CONDIT	iours .	r ature	Hours																mns are
MENTAL	Exposure, liours	Over Tempera	151 to ↓ °F																se columns
ENVIRONMENTAL	Liquid Exp	140 to	150°F	14.0	16.5	14.5	23.5	21.0	40.5	7.5	22.5	40.5	22.0	18.5	22.5	18.0	18.5	19.0	in these
	Fuel Lig	Ambient To	139°F	0.6	3.0	10.5	2.0	2.0	2.0	16. ^م	2.5	2.0	ī.5	7.0	2.0	3°2	3.0	3.5	the hours
	Environmental Conditioning	Cycle No.		69	70	71	72	73	74	.°5	76	77	78	79	80	81	82	83	(1) Most of t
	EVLT	R: D	-		-					51	-	SE			G	NO. PAG		- 82 83	

Standard and Carlos A. Standard and the Standard Street

2

1

A PARTICULAR DE LA CARACTERIZA DE LA C

A STATE OF SUPPORT STATE

2

のためにないのない

151

ta Ali Maguzan Mula - undu ana undu a un

ļ

These hours are included in hours reported under 426 - 441°F.

over-temperature controls turning heaters off.

 \odot

س		ture	Hours																
		Over Temperature	441°F to4°F																
	Hours	426 to	441.05	137.5	136.0	138.5	137.0	137.5	136.0	137.5	145.5	138.5	110.0	128.5	132.0	138.5	137.5	129.0	caused by
	Exposure, H	415 to	441°F																downs ca
NO. 3	Vapor Exp	401 to	425°F	1.5	2.5	2.0	2.0	2.0	2.5	2.5	2.0	3.0	3.5	2.5	3.0	3.5	3.0	3.5	cool
×	Fuel V	14C to	400°F	5.5	7.0	6.0	6.5	6.5	6.5	6.5	16.5	6.0	6.5	10.5	6.5	7.0	8.0	6.5	unplanned
<pre>' (Continued) IONING OF TAN</pre>		Below	140°F																mns are due to
TABLE ' (Cont CONDITIONING	ours	ture	Hours																mns are
MENTAL	Exposure, H	Over Tempera	151 to + °F					1											se colui trait
ENV I RONMENTAL		140 to	150°F	19.5	18.5	19.5	19.0	18.0	19.0	19.5	18.5	14.5	12.5	16.5	19.5	18.0	21.0	19.5	s in these colu
	Fuel Liquid	Ambient To	139°F	3.0	3.0	3.0	3.0	3.0	3.5	3.0	3.0	8.0	11.5	3.5	3.5	4.5	3.5	4.0	
	Environmental Conditioning	Cycle No.		84	85	я6	87	88	89	90	16	92	93	94	ý 5	96	97	86	Most of the hour
R.	EVLT)				15	2		Sit	CI.	:// 3	187	NO. PAG		829	

ļ

to more and a state of the

ļ

ł

ì

i

•

ころの あいまたい あまま にない うちになる ひまいがっ

These hours are included in hours reported under 426 - 441°F. over-temperature controls turning heaters off.

 \odot

ω.		r ature	Hours																
		Over Temperature	441°F tot°F																
	Hours	426 +0	441°F	138.0	63.0	97.5	114.5	1 .	131.0	132.5	131.5	132.5	133.5	116.0			132.0	131.5	caused by
	Exposure, 1	415 to	441°F																downs ca
NO. 3	Vapor Ex	401 to	425°F	4.5	3.0	2.2	3.0	3.5	3.5	4.0	3.0	3.0	4.0	6.5	3.5	•	3.5	3.5	ed cool
X	Fuel 1	140 to	400°F	37.5	54.5	14.5	5.5	6.0	6.5	5.5	5.5	6.5	5.5	14.5	. 5.0	5.5	6.5	5.0	to unplanned off.
TABLE 7 (Continued) CONDITIONING OF TAN		Below																	۱۵۷
TABLE CONDIT	Hours	r ature	Hours																umns are du ing heaters
MENTAL	Exposure, l	Ove mper	151 to + °F																se colu s turn:
ENVIRONMENTAL	ש	140 to	1.50°F	18.5	19.5	0.01	21.5	16.5	19.5	20.5	19.0	19.5	19.0	20.5	21.5	20.0	19.0	18.0	s in these controls
	Fuel Liqui	Ambient To	139°F	3.5	4.0	3.0	3.5	3.0	3.5	3.0	4.0	3.0	4.0	4.5	4.0	-4.5	4.5	·5 . 0	hours ature
	Environmental Conditioning	Cycle No.		66	100	101	102	103	104	105	106	107	108	109	011	111	112	113	(1) Most of the hours over-temperature
	EVLT		•							- •		SE		-	<i>Æ</i> 7	NO. PAG	_	- 82	
r`(K; L							153	3		1	~··		l	. 40	•		

4

1

يند بدو تام المالية المالية من المالية المالية المالية المالية المالية . 2014 من المالية .

(3) These hours are included in hours reported under 426 - 441°F.

w

والمحمومة والمحمولة والمعادية والمحمولة والمحمولة والمحمومة والمحمومة والمحمولة والمحمولة والمحمولة والمحمولة

Hours Temperature \odot Over 441°F to+°F 426 to 441°F 131.0 139.5 130.5 44.5 130.0 131.5 27.0 129.0 131.0 131.0 129.5 132.0 121.5 Hours Fuel Vapor Exposure, 415 to 441°F 401 to 425°F 4.0 3.5 4.5 3.5 3.5 4.5 4.0 3**.**5 3.5 4.0 7.0 4.0 3.5 ო 1 CONDITIONING OF TANK NO. 140 to 400°F TABLE 7 (Continued) 5.0 6.0 5.0 8.5 ວ. ວ 5.0 5.0 5.0 4.5 4.5 12.5 4.5 5.5 Below 140°F 61 નિ Hours Fuel Liquid Exposure, Hours Temperature Over ч Ч С Ч ENVIRONMENTAL 151 -> 150°F 18.5 17.5 18.5 18.5 16.5 18.5 19.0 17.0 17.5 18.5 18.0 18.5 18.0 19.0 t t 140 Ambient To 139°F 4.0 5.5 4.5 4.0 6.0 4.5 4.5 6.5 4.0 4.5 4.0 4.5 4.5 4.0 Snvironmental onditioning Cycle No. 118 114 116 115 119 117 120 122 123 124 125 126 121 127 BOEING NO. D3-8297

Most of the hours in these columns are due to unplanned cool downs caused by over-temperature controls turning heaters off.

131.5

4.5

4.5

19.0

4.5

128

83.8

PAGE

These hours are included in hcurs reported under 426 - 441°F

 \bigcirc

F

ļ

FF-VLTR-Ε

154

SECT

يىة		ture	Hours)	65.0												
-	,	Over Temperature	441°F to+°F		442-460												-
	Hours	426	441°F	132.5	16,481.8		HRS.							 			caused by
	Exposure, H	415	441°F		78.5		FUEL VAPOI										downs cau
NO. 3	Vapor Exi	401 to	425°F	4.5	470.0	-	19,346.6										ed cool downs - 441°F.
Ä	Fuel V	140 to	400°F	4.5	1794.8				F.XPOSURE								olann 426
TABLE , (Continued) CONDITIONING OF TAN		Below			521.5				TOTAL HOURS								
_	ure, Hours	Over Temperature	I to Hours °F		151- 205°F 113.2		FUEL LIG. HRS.		22.269.3 TO				 			 	columns turning n hours
ENVIRONMENTAL	uid Exposure,	140 to	150°F [151	19.0	2133.5 20		2,922'.7 m						 			 	s in these controls included in
	Fuel Liquid	Ambient To	139°F	5.0	675.0												hours ture are i
	Environmental Conditioning	Cycle No.		129	TOTALS (1-129)		SURTOTALS		GRAND TOTAL							 	 Most of the hours over-temperature These hours are :
R.E	EVLTF			ł	•••••••			{		55	4.	SEC	 115		NO. PAGE	7	

a v v a div a a anishanan

ちゅうろう くしせんじい ひろくろし しゅいいしい とくせいれん

NAMES A CONSTRUCTION MADE IN CONSTRUCTION

2

わたいてもうちょうちゃう あっかかかいろう ちゅうちん ひしゅう アンドレート

لتوزعه فلامتناقذه وعدوما فالمانا فللتوجيد ومعرار المتواسيكم

Date	Test Interval Completed			Cor	ners	Leaki	ng (1)	
		R-1	R-2	R-3	R-4	F-1	F-2	F-3	F-4
3-71	After 8 Env. Cond. Cycles		x	x	x		x	x	x
8-71	After Repairs of Corners		x					X	
9-71	After Load Cycling at 3 Temperatures		x		x			x	
1-72	After 22 Env. Cond. Cycles		x		x	x		x	x
1-72	After Load Cycling at 3 Temperatures	х	x		x	x		x	x
2-72	After Repairs		x				x	x	x
8-72	After 39 Env. Cond. Cycles Plus Loading at 3 Temperatures	x	x			x		x	
9-72	After Repairs of R-1 & R-2					X		x	
11-72	After 46 Env.Cond.Cycles	X	X	X	X	X	x	x	X
11-72	After Repairs of R-1,R-2,R-3,R-4				X	X	X	X	X
2-73	After 54 Env. Cond. Cycles	X	X	X	X	X	X	X	X
2-73	After Repairs of R-1,R-2,R-3,R-4			1		X	X	X	X
8-73	After 74 Env.Cond.Cycles		X	1	1			X	X
2-74	After 92 Env.Cond.Cycles			X	X	X	X	X	X
7-74	After 117 Env.Cond.Cycles		X		x	X	X	X	X
9-74	After 120 Env.Cond.Cycles 2		x		x			x	X
1-74	After 129 Env.Cond.Cycles	x	X	X	X	X	X	X	X
Corne	age B-15 for locations of co rs F-1 and F-2 were repaired LTR: E	rners.	•	B D SECT	EIN	G NO	. D3- GE 8	8297	

TABLE 8

CORNER LEAKAGE HISTORY OF TANK NO. 3

E-3033 RT

and the second second

Section 2.

n China and an east in a picture state.

Souther Statistics and the states of the sta

ning and a start of the start o

فستنفاق فسيلاق بالاختر وتجاورا أفلأخ والتحمل مراجع والموجع

in the second

156

E ↓ !

APPENDIX A

LEAKAGE TESTS - TANK NUMBERS 1 AND 2

	NO.	D3-8297
SECT	PAGE	84

าร ค.ศ.ศ.ศ.ศ. 2. เป็นสาย 2. เป็นสา 2. เป็นสาย 2.

ł

مەر ئىلگۈم چۈم ئەر ئىلغانلىق ئەر ئىلغىگەندۇ. يەڭ ئەر ئەلگەنگەنگەنگەنگەنگە كەر كەركەنگەنگە ئەلگەنگەنگەر ئەلگەن مەر ئەم ئەر مەر ئەم ئەلغەن ئەر ئەلگە ئەر ئەلغەن يەڭ ئەلگەن ئەلگەن ئەلگەن ئەل ئەر ئەم ئەر ئەلگەن ئەلگەن ئەر ئەلم

. NCM X.

ちょうちん ちょうちょう

ちょうちょうかい おおう かんかい いったいかい たいちょう ちんちょう ちょうちょう ひょうちょう

197

ariandiana akas

• A •

REVLTR:

E-1011 R1

ļ

ġ.,

1.

1

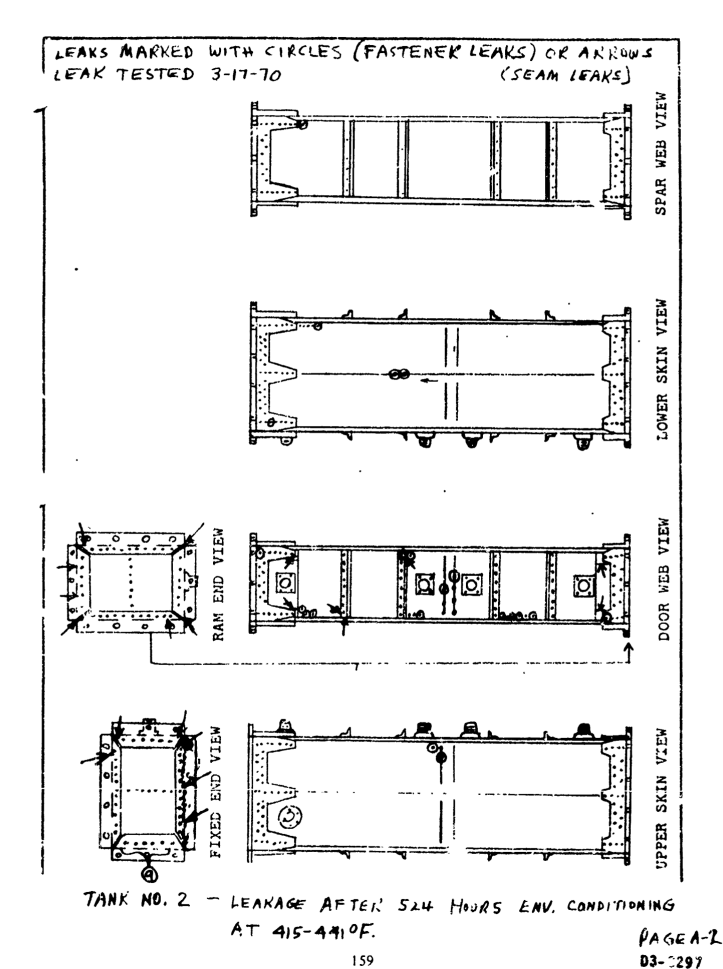
ŝ

LEAKAGE TEST DATA SPECTS

LEGEND FOR LEAKAGE SYMBOLS : FILLET OVER FASTENER () SEALANTA SHOWS LOSS OF ADHESION TO TITANIUM. 2 SMALL CRACK IN FILLET OVER FASTENER 3 INADEQUATE FILLET APPLICATION CAUSE OF LEAK UNKNOWN 5 LOSS OF ADHESION, BUT NO LEAK 6 SEALANT FILLET CRACKED \mathbf{O} SEALANT FILLET SHOWS LOSS OF ADHESION TO T) TANIUM 8 SEALANT FILLET SHOWS LOSS OF ADHESION SERNETAL W TO **(1)** EVIDENCE OF SEALANT REVERSION $(\mathbf{0})$ FILLET CRACKED OR LOOSENED BY EXPANSION OF INJECTION UNDERNEATH . (\mathbf{I}) GENERAL OR CONTINUOUS JOINT LEAKAGE LOSS OF ADHESION OF REPAIR FILLET TO ORIGINAL FILLET (12)(13) LOSS OF ADHESION OF REPAIR FILLET TO STRUCTURE (14) CRACK IN REFAIR FILLET SMALL CRACE OR LOSS OF ADHESION IN JOINT FILLET, GENERALLY (15) T" INCH LONG, CRACK GENERALLY PENETRATES TO SUBSTRATE WITH SOME ACCOMPANYING LOSS OF ADHESION .

7 CRACK

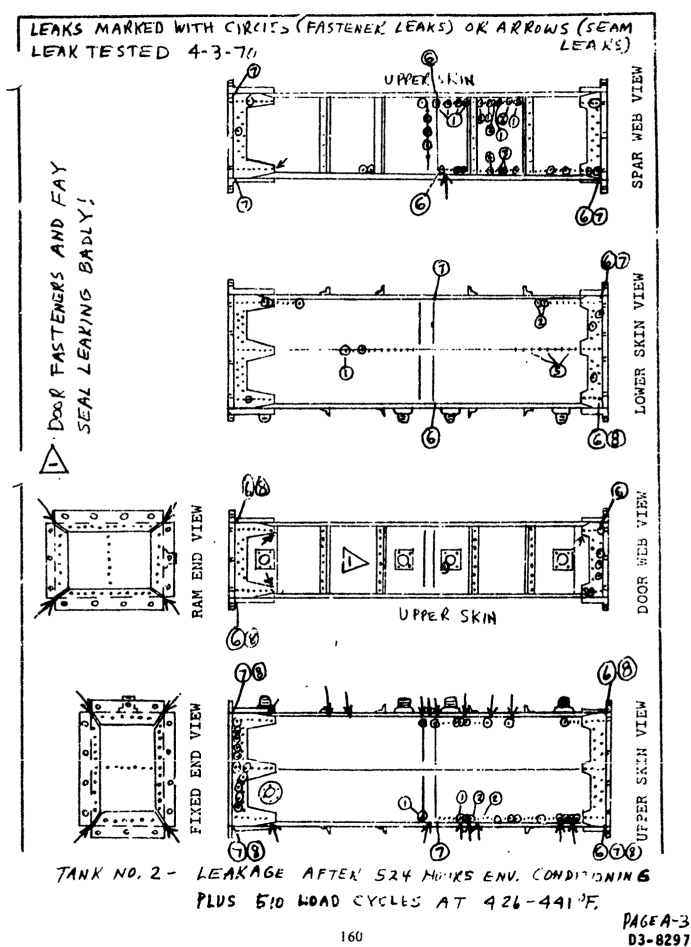
LOSS OF ADHESION

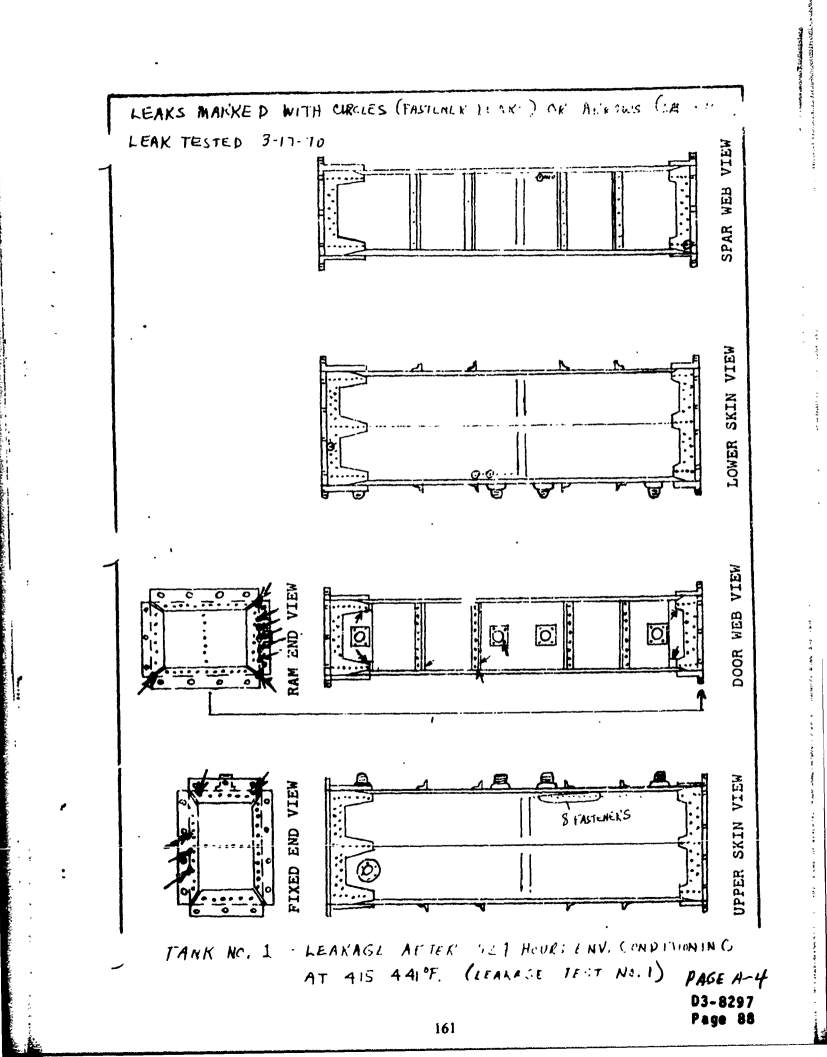


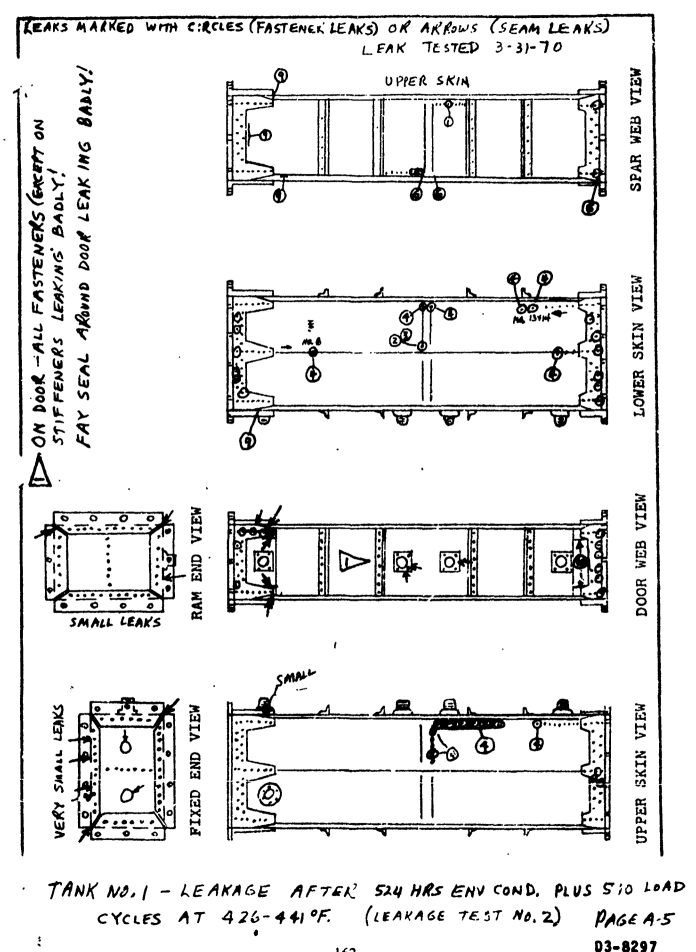
Ř

5

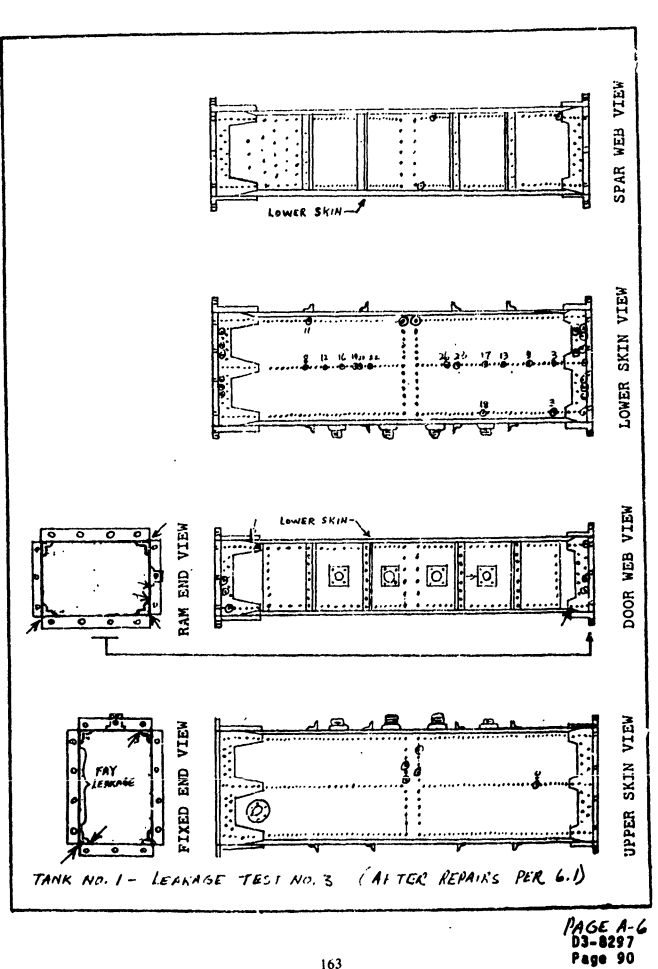
159



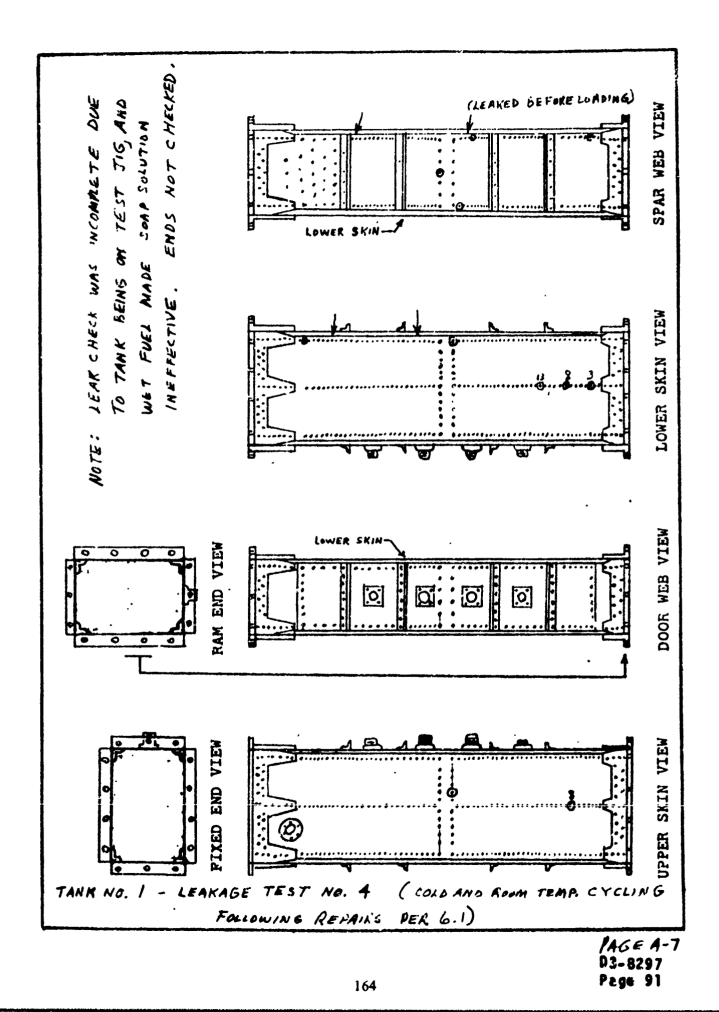


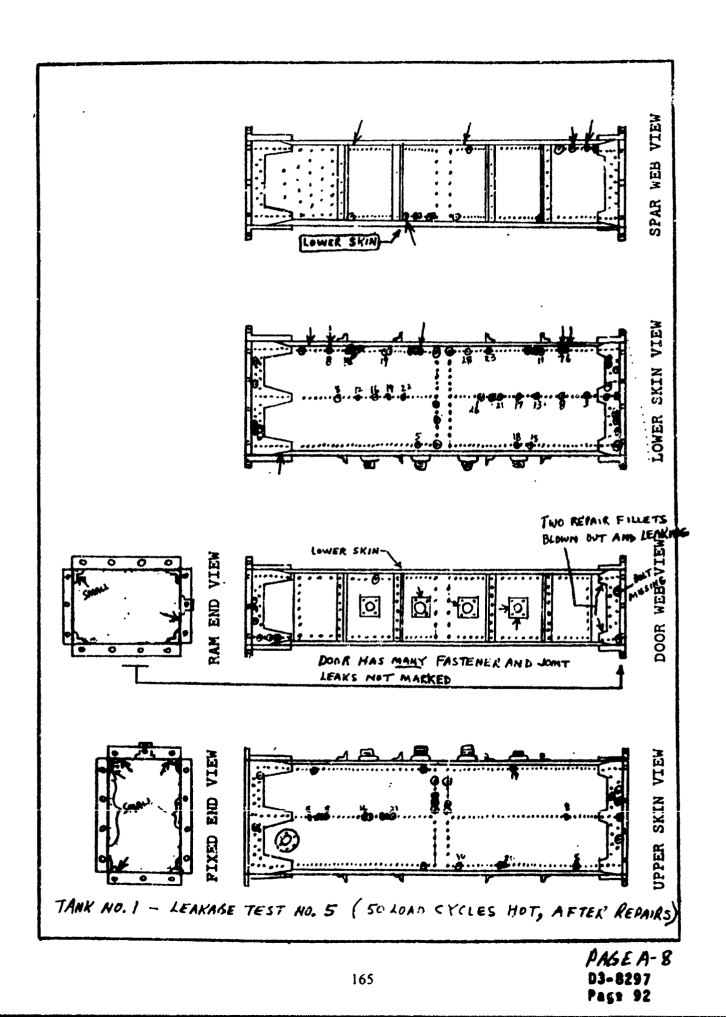


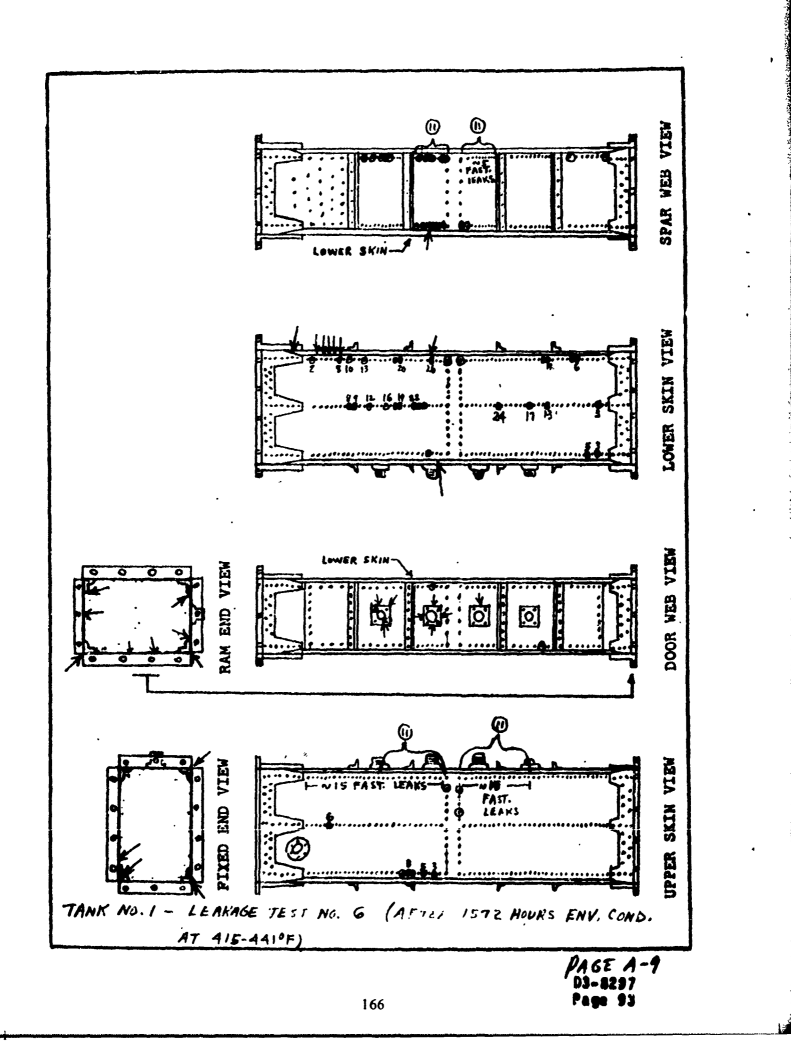
¹⁶²

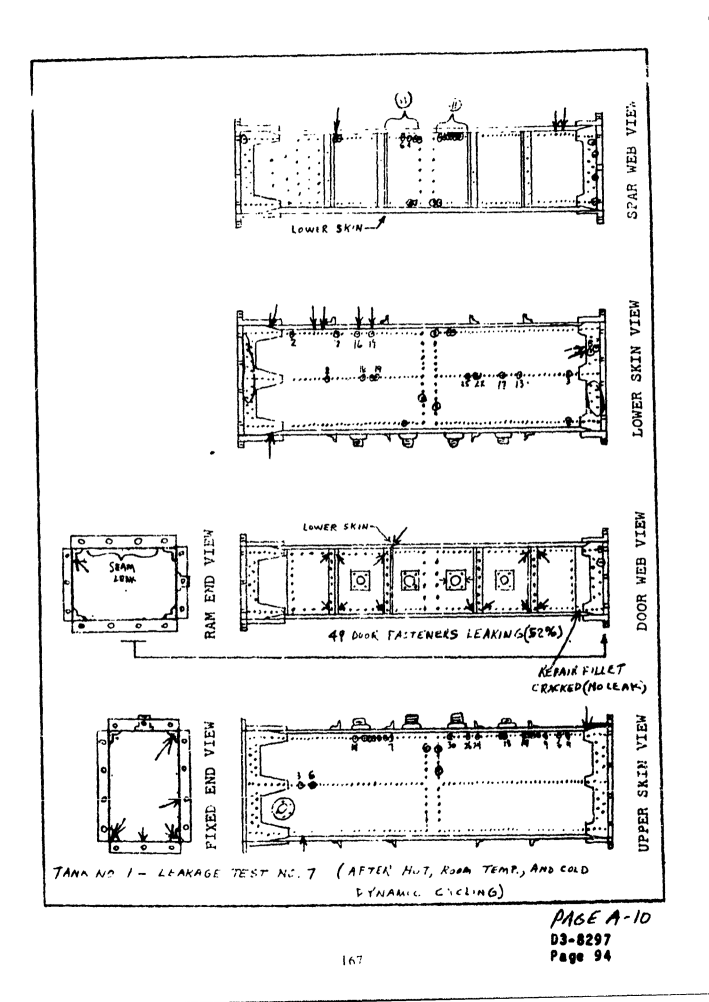


:

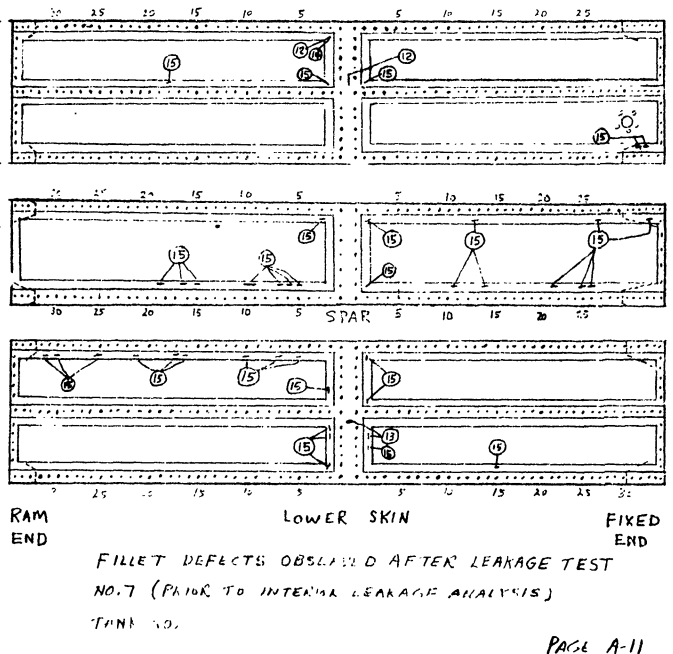




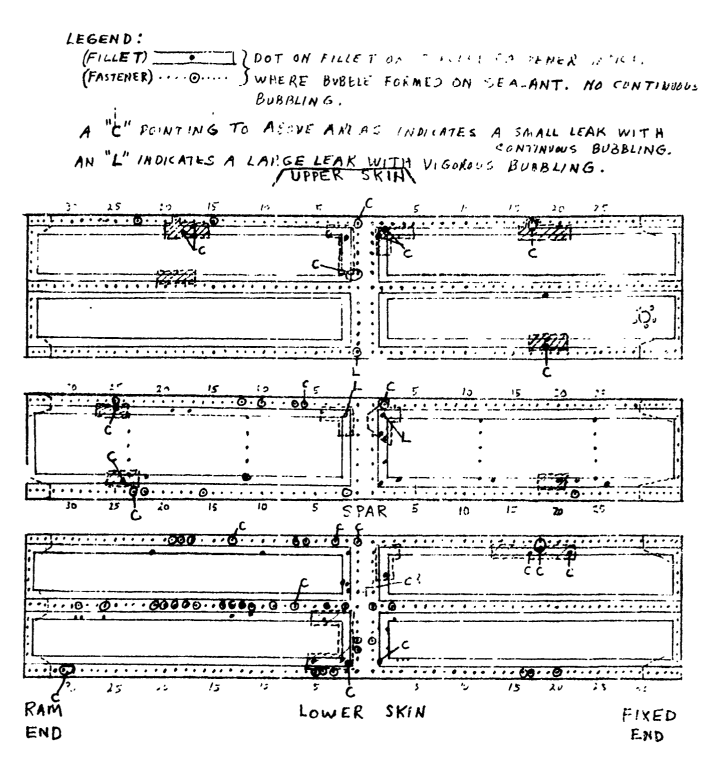




UPPER SKIN

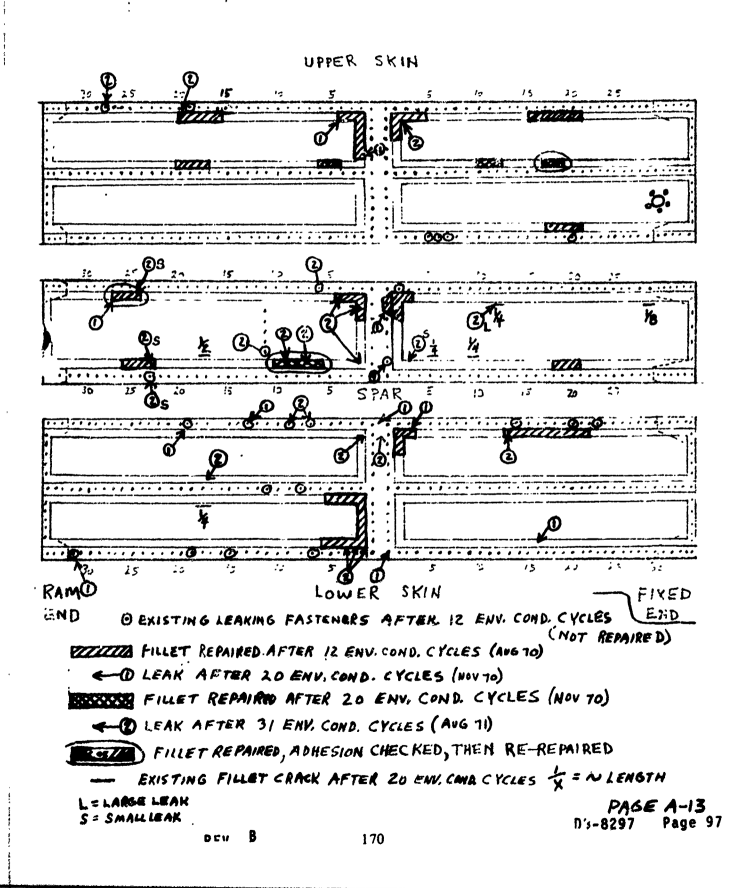


D3-8297 Page 95



TANK NO, 1 INTERIOR LEARAGE ANALYSIS CONFUCTED AFTER 12 ENVIRONMENTAL CONDITIONING CYCLES, ANALYSIS CONDUCTED USING 5 PSIG NEGATIVE (VACUUM) PRESSURE IN JANK WITH WATER LAYER ON INTERIOR TO VISUALLY INDICATE LEARAGE BY BUBBLING. PAGE A-12 03-8297

169



B ·

APPENDIX B

LEAKAGE TESTS - TANK NUMBER 3

MOEINS	NO. D3-8297	
	PAGE 98	-94-

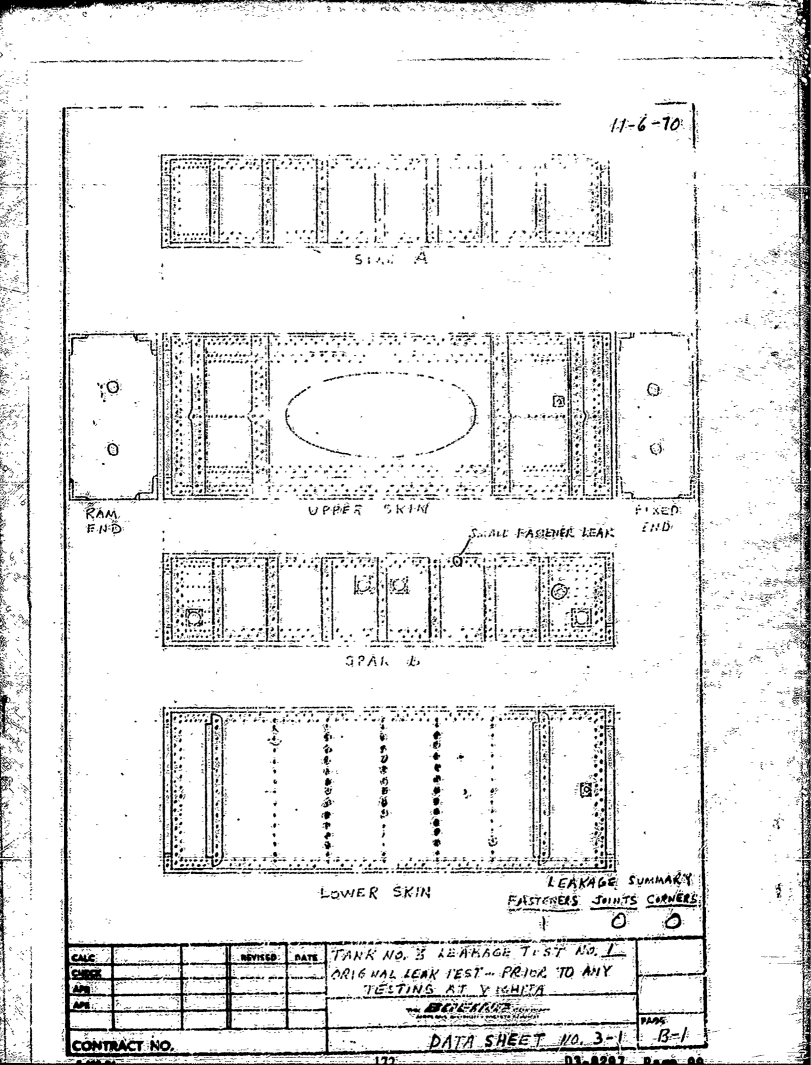
L TELEVIL

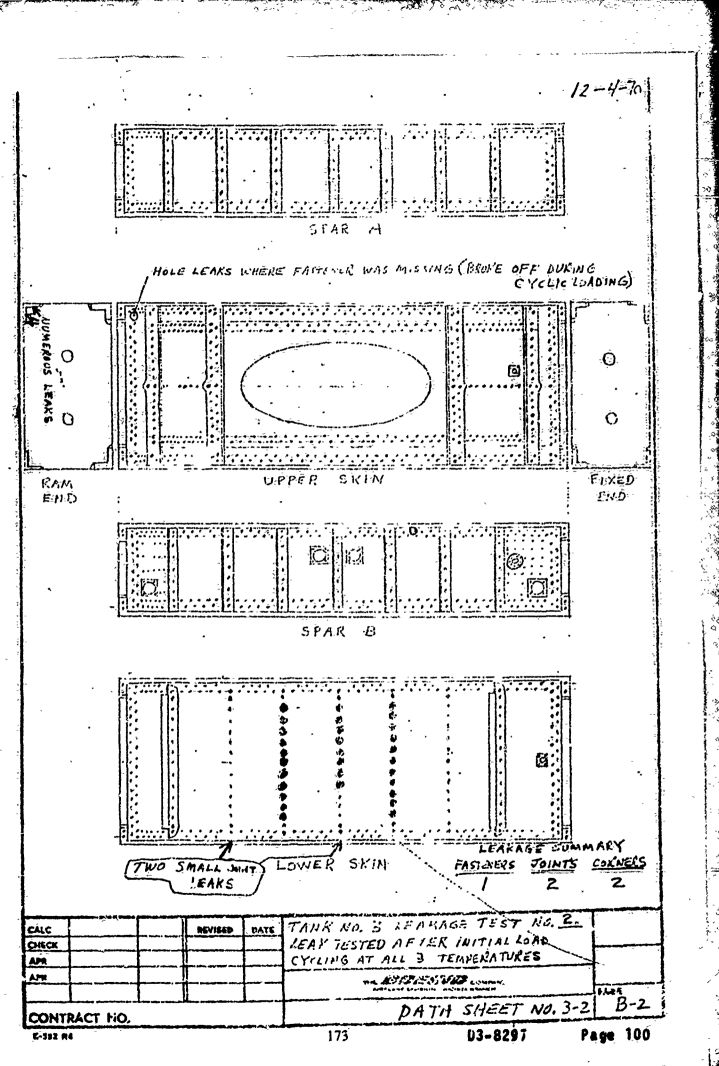
REVLTR: A

E-3033 R1

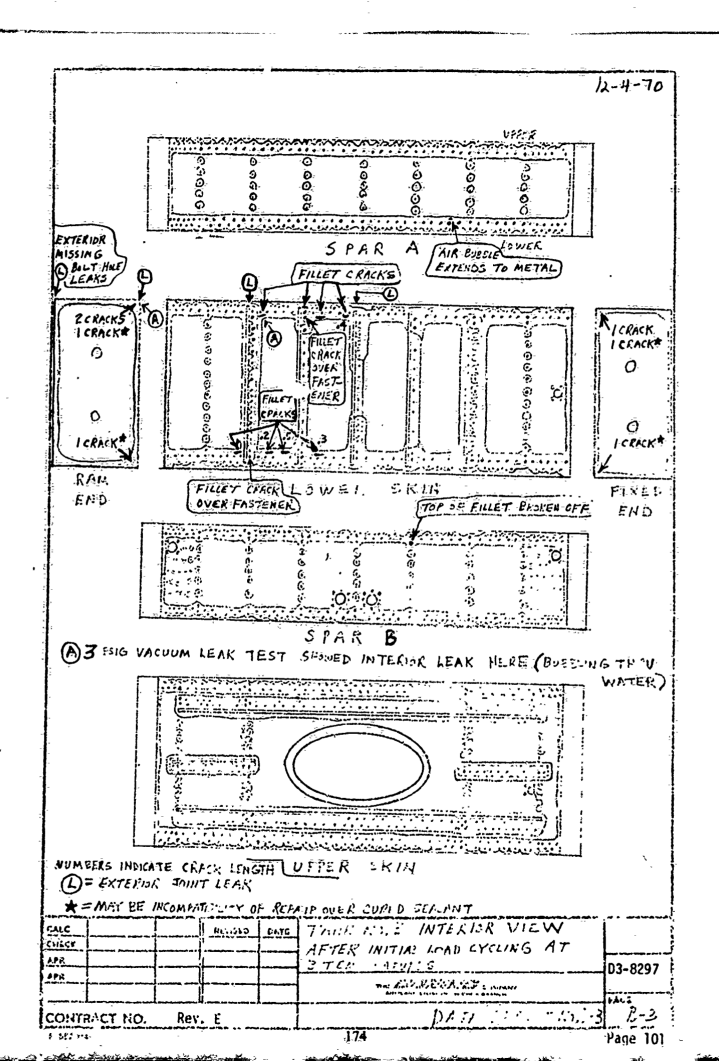
3

*



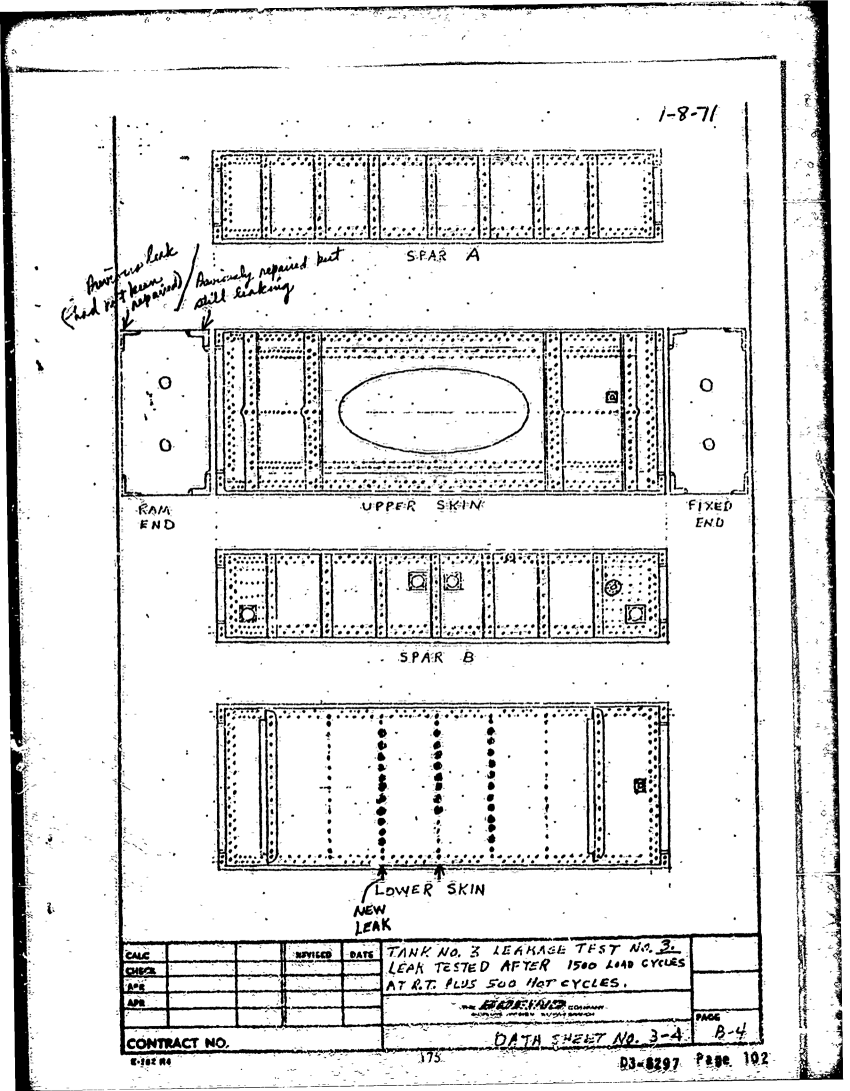


<u>______</u>

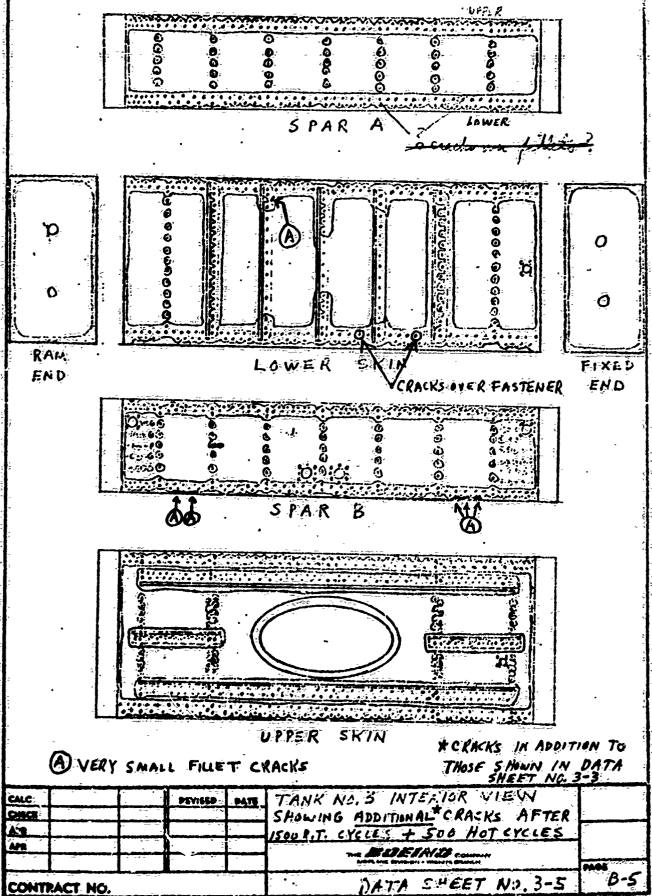


1. S.

67.0



En D. C



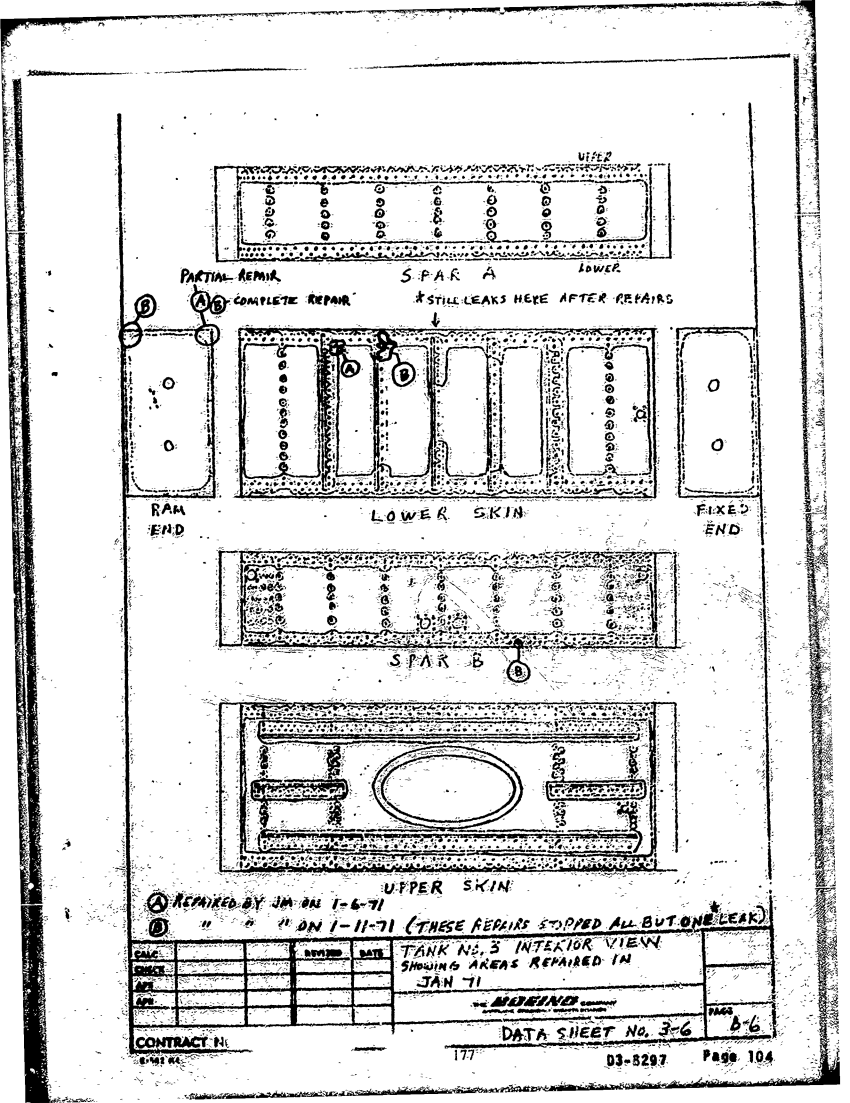
176

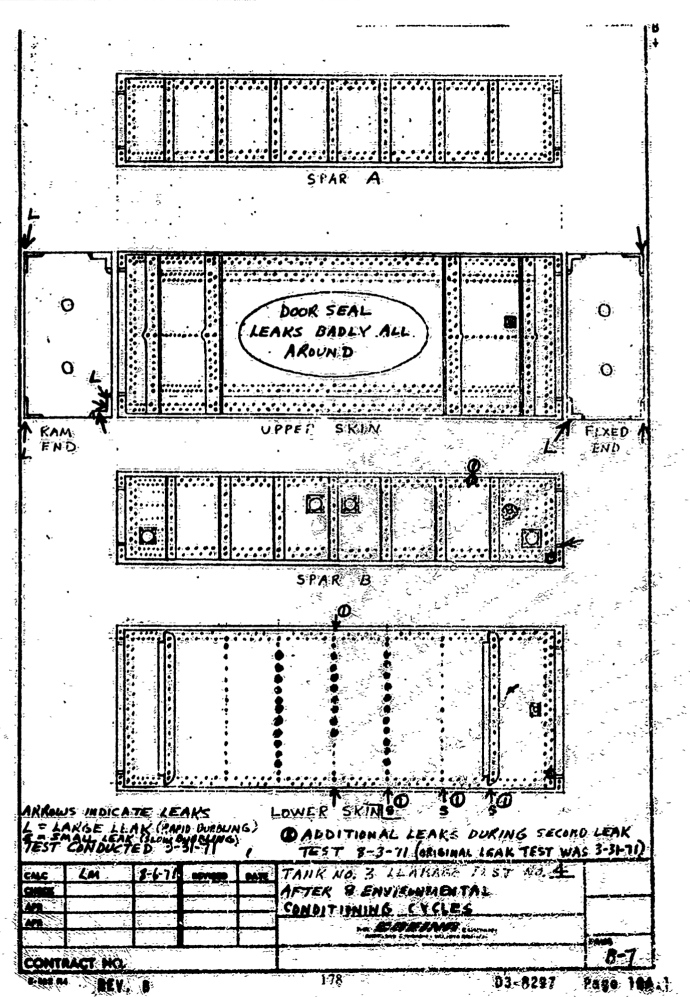
D3-8297 Page 103

いたというというかい

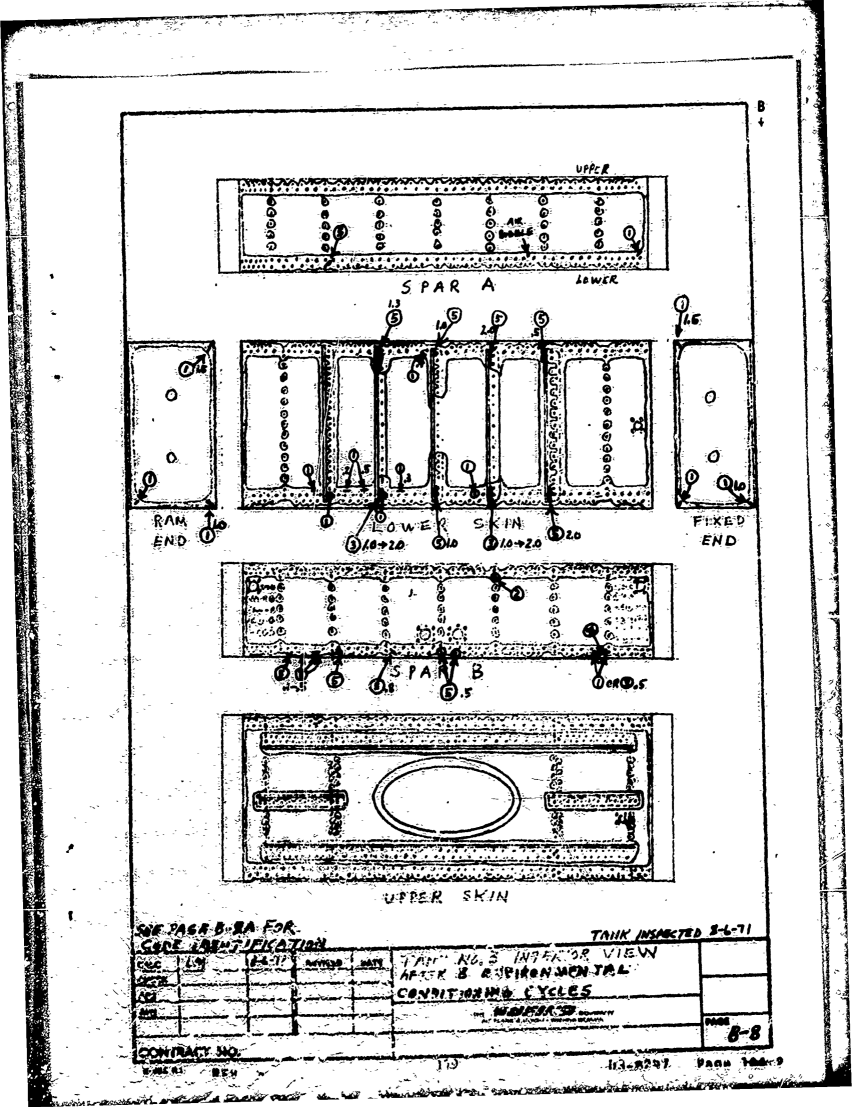
「「「「「「「」」」

C-362 84





「「「「「「「「」」」」



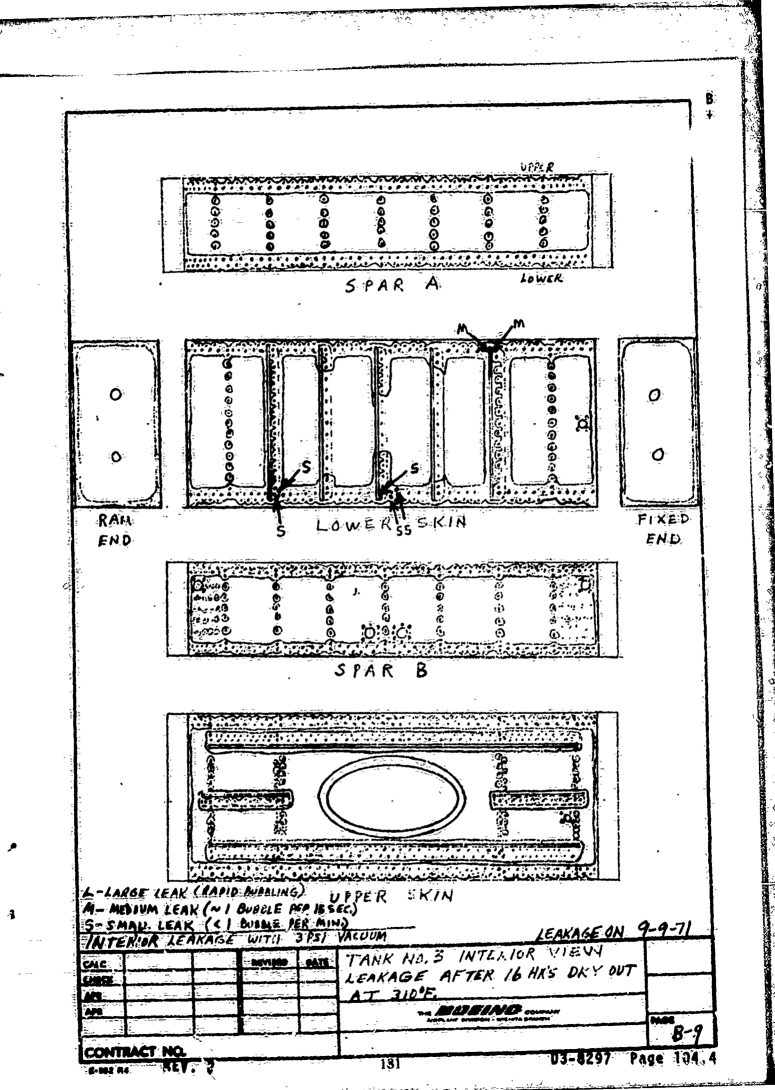
- CODE FOR FAGE B-8
- Der Indigates FILLET CRACK (LENGTH AS NOTED IN INCHES)
- D + FILLET OVER FASTENER PARTIALLY, TORN OFF
- OUT (IGHES 310) ON AUG 5 TO PREPARE FOR REPAIRS.
- A Loss OF ADHESION
- (BETWEEN GAPRIL AND GAUGUST 1991) OR WHEN TANK WAS DRIED. OUT (16 HRS @ 310°F) ON AUG 5, 1991 TO PREPARE FOR REPAIRS,

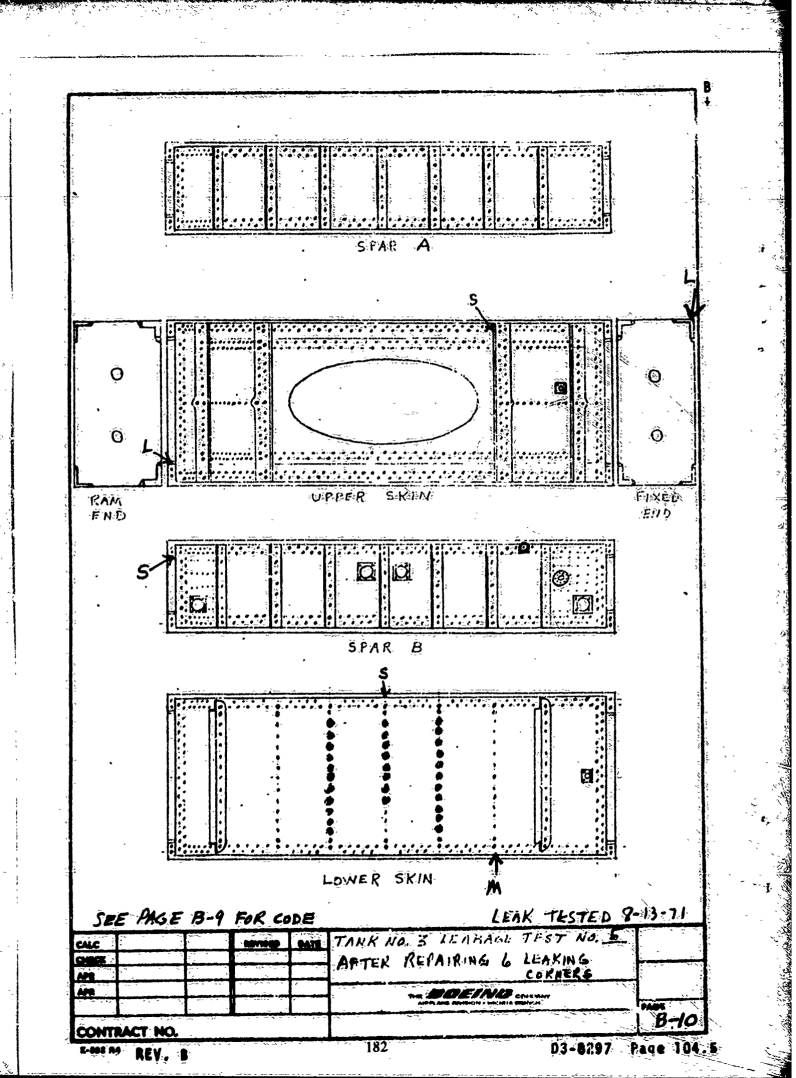
180

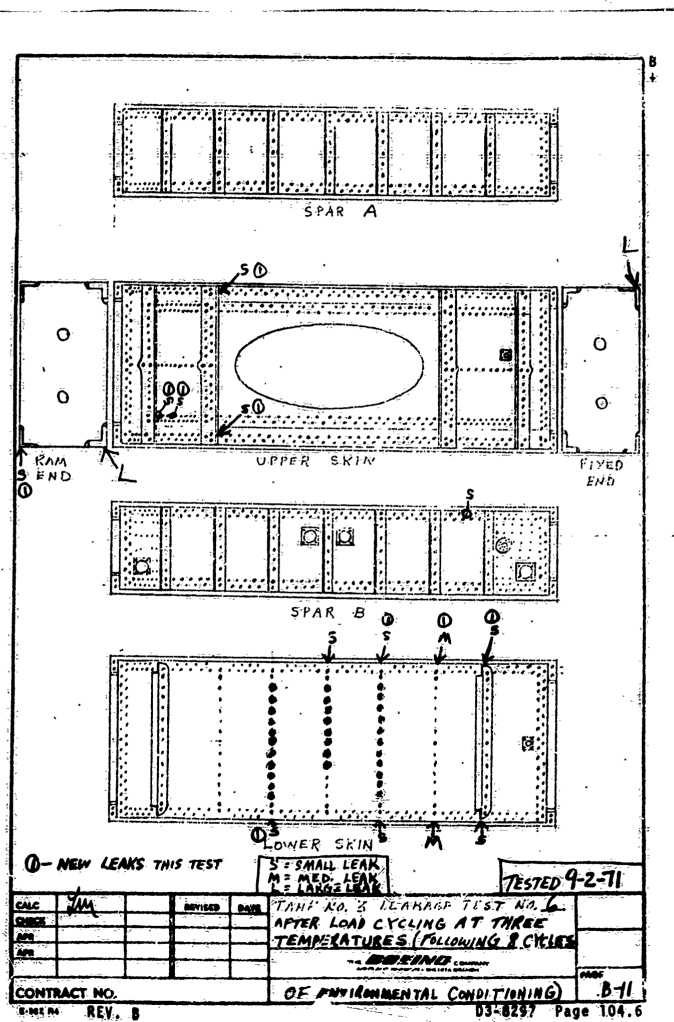
PAGE B

e 104. 7

03-8297



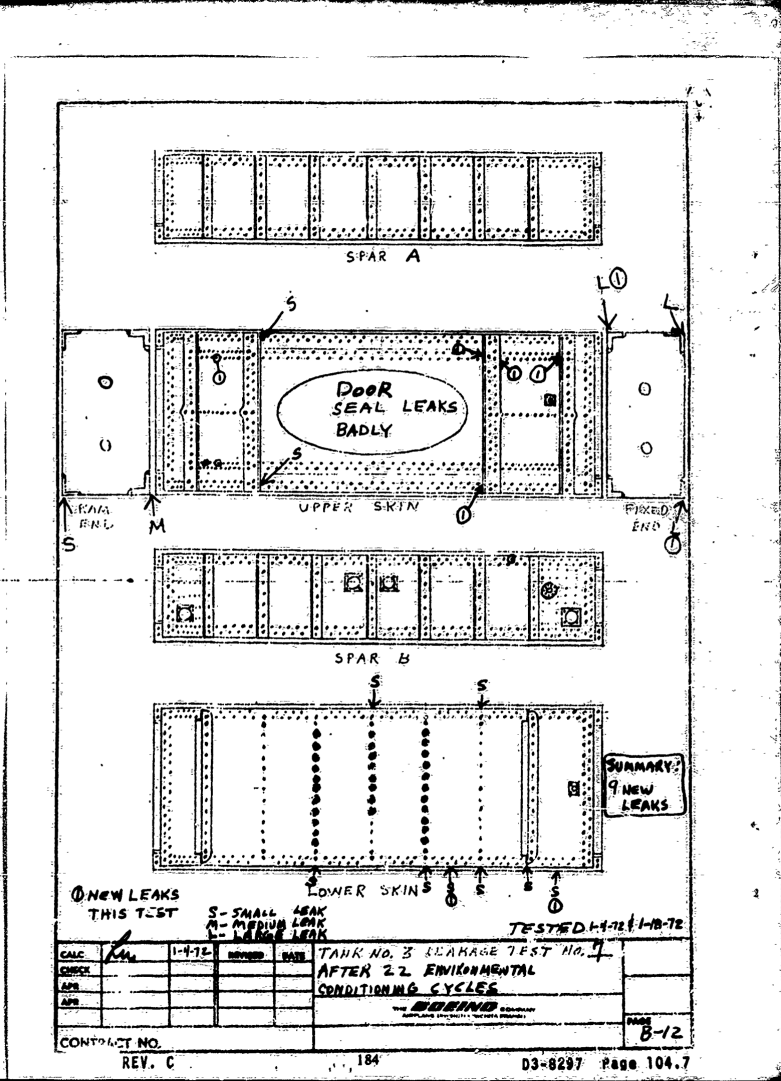


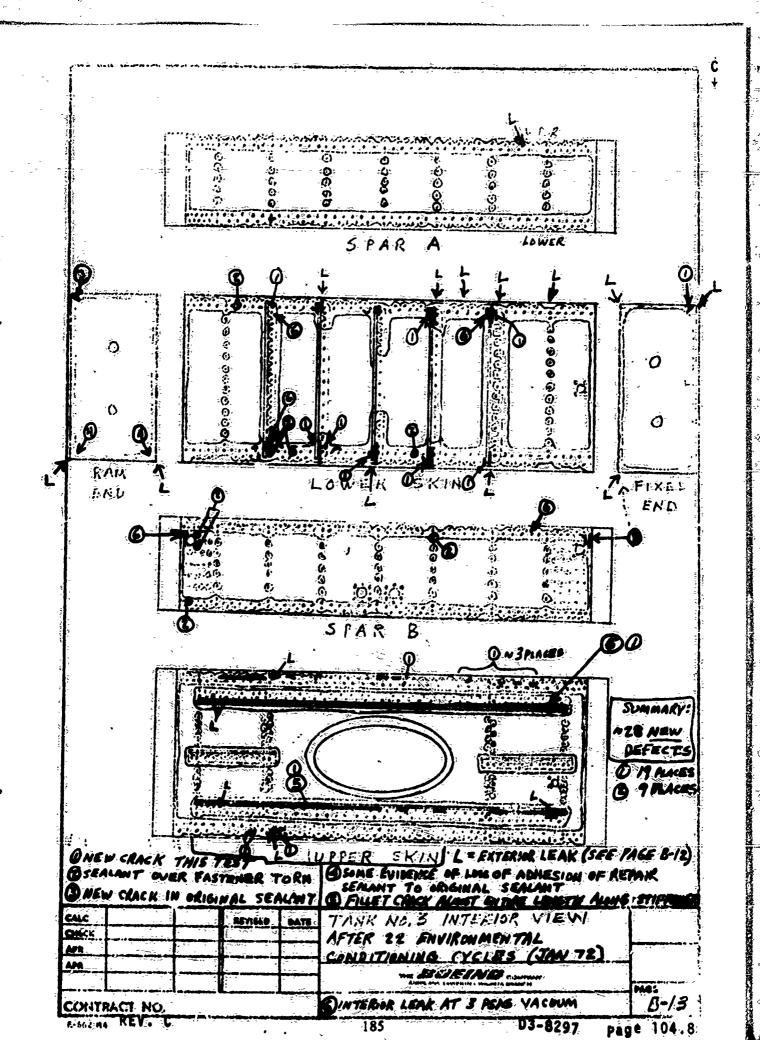


C 7 7

ς - ³

.....

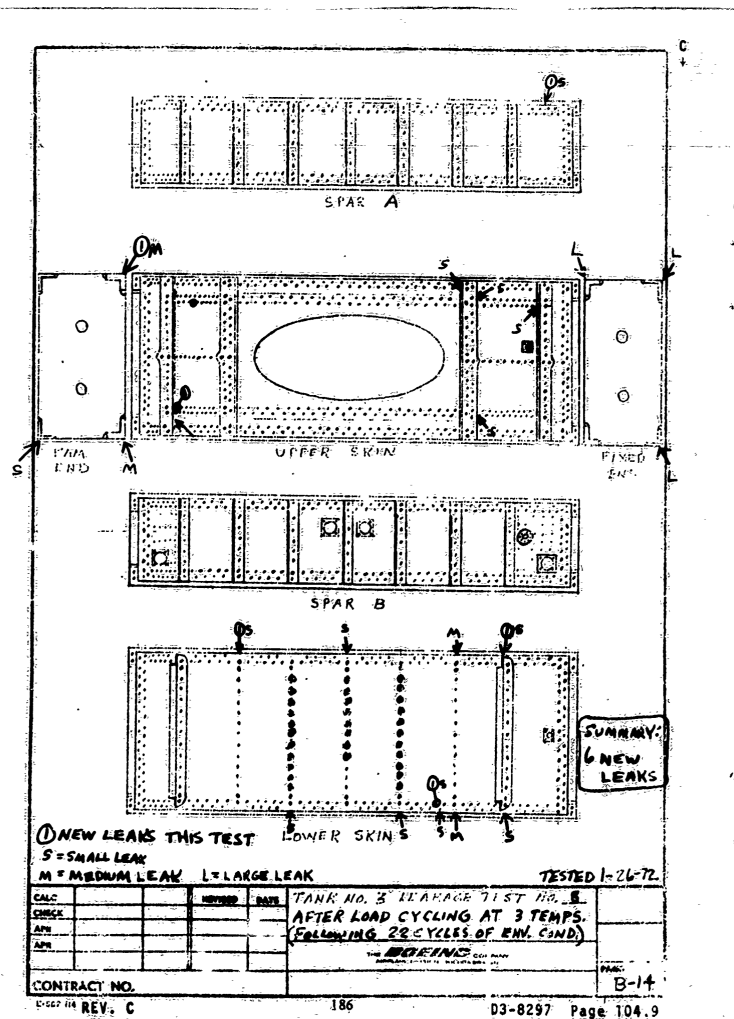




`--',

and the second

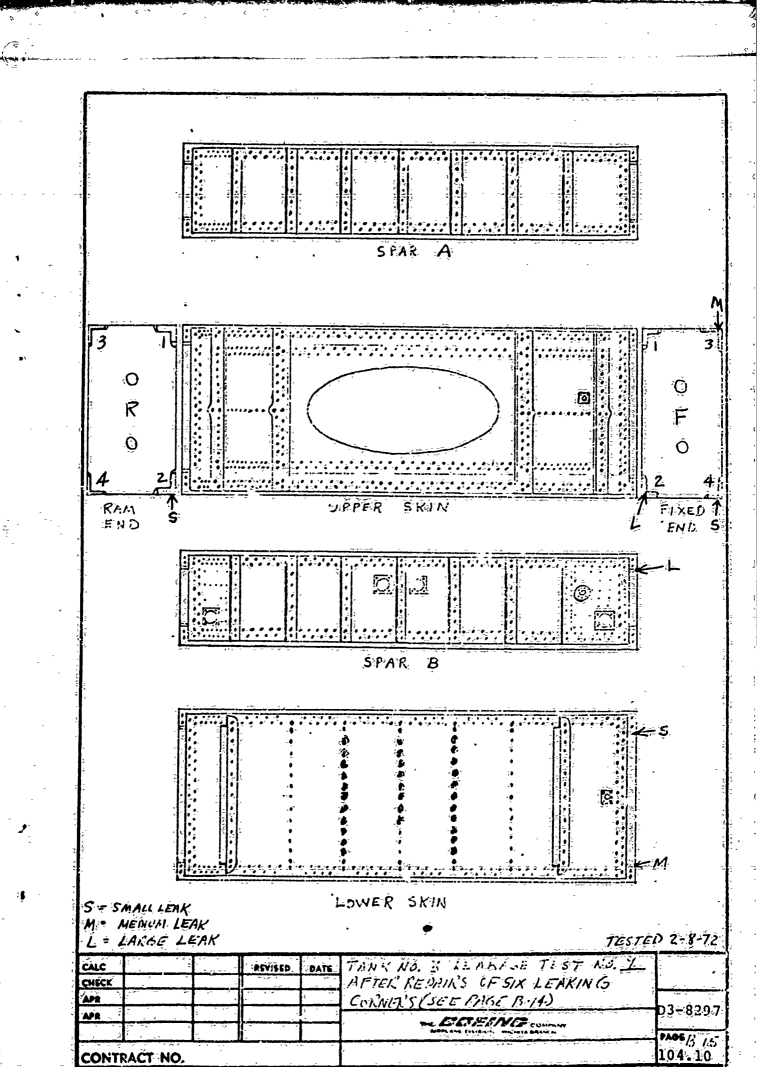
127



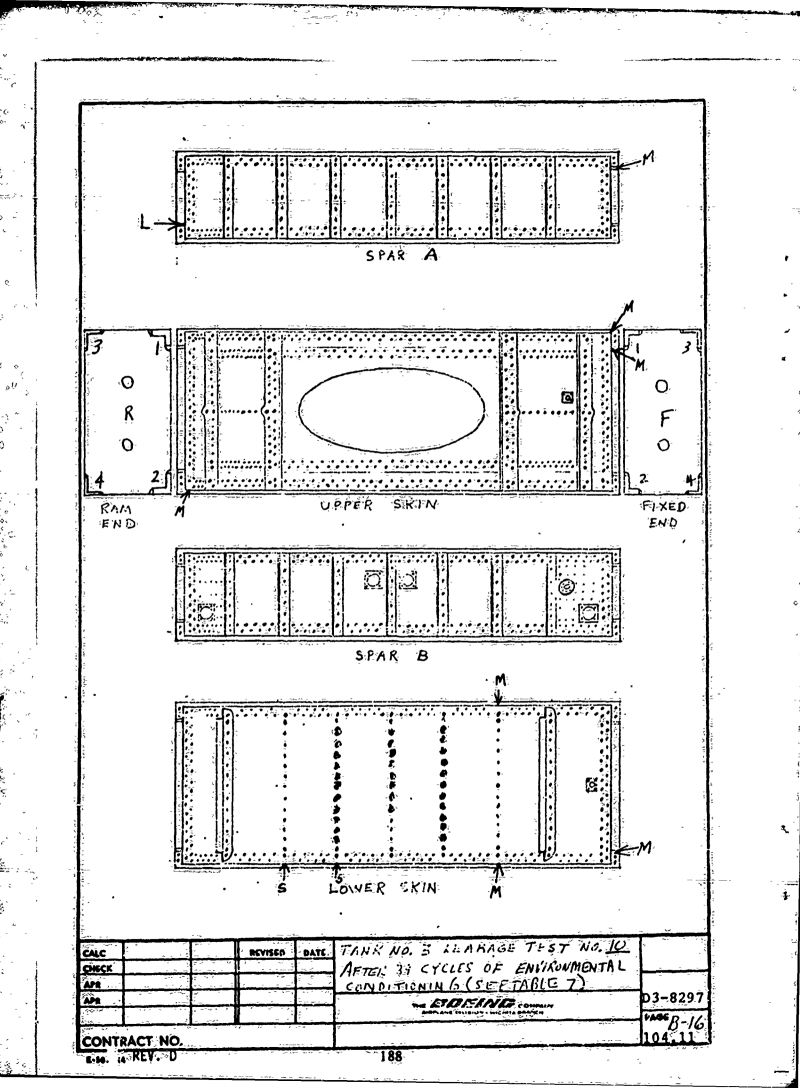
and the second sec

× .

ð

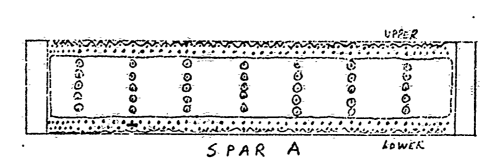


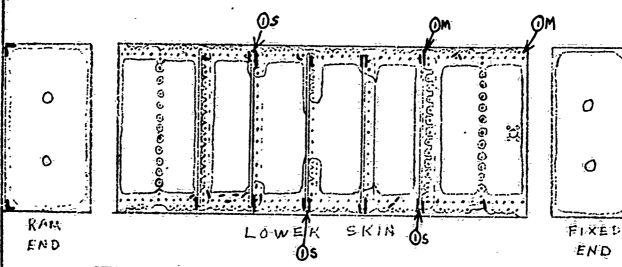
T

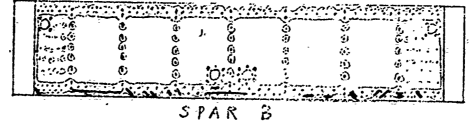


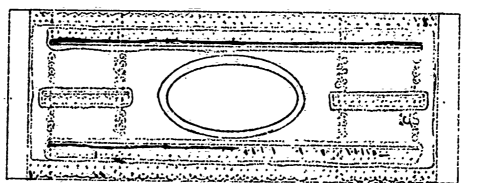
ૡૡૣૡૡૢ૱

ais féa









UPPER CRIN 5 = SMALL LEAK M = MEDIUM LEAK D->INTERIOR LEAK-WATER BUBBLING THRU 3 PSI VACUUM = LARGE LEAK

,

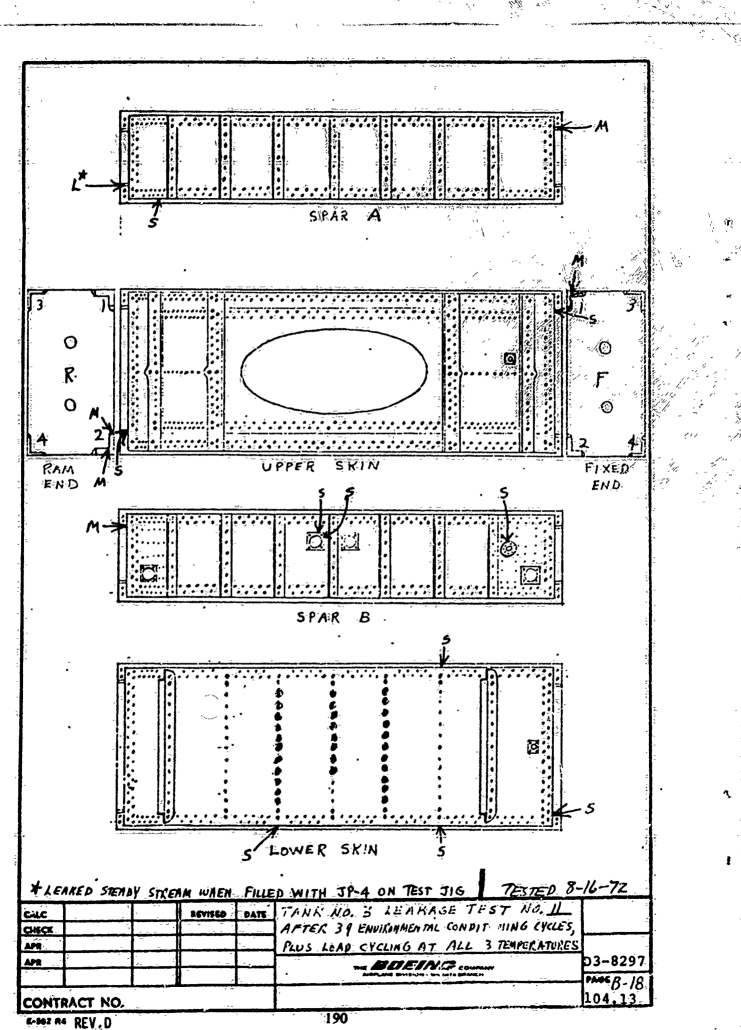
TRITI · JOK

TESTEP	0-1.12
11N	

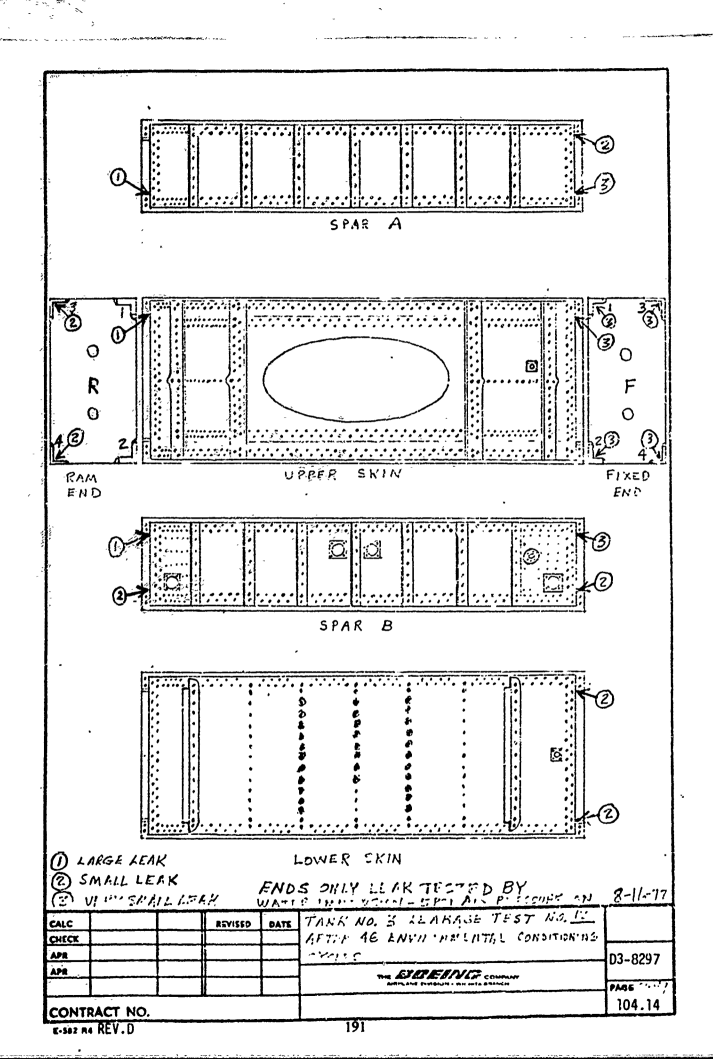
CALC <u>CHECK</u> <u>A82</u> A82		ASVISTÒ	 TANK NO INTERIAR THENTAL CONDITION - AFTER 39 CYCLES OF ENVILONMENTAL CONDITION - ING, PLUS IBHOURS DRYOUT AT 400 °F. The ALL OF STREET COMMONS	D3-8297
CONT	RACT NO.			104.12

6-132 44 REV. D

189



il.

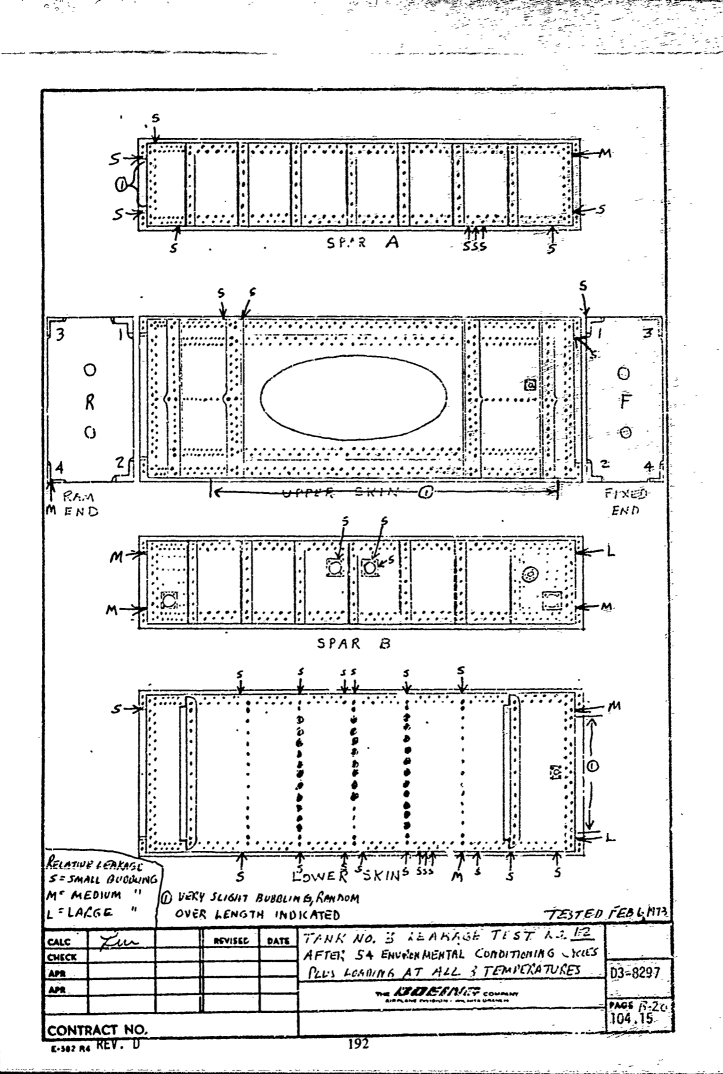


11-20

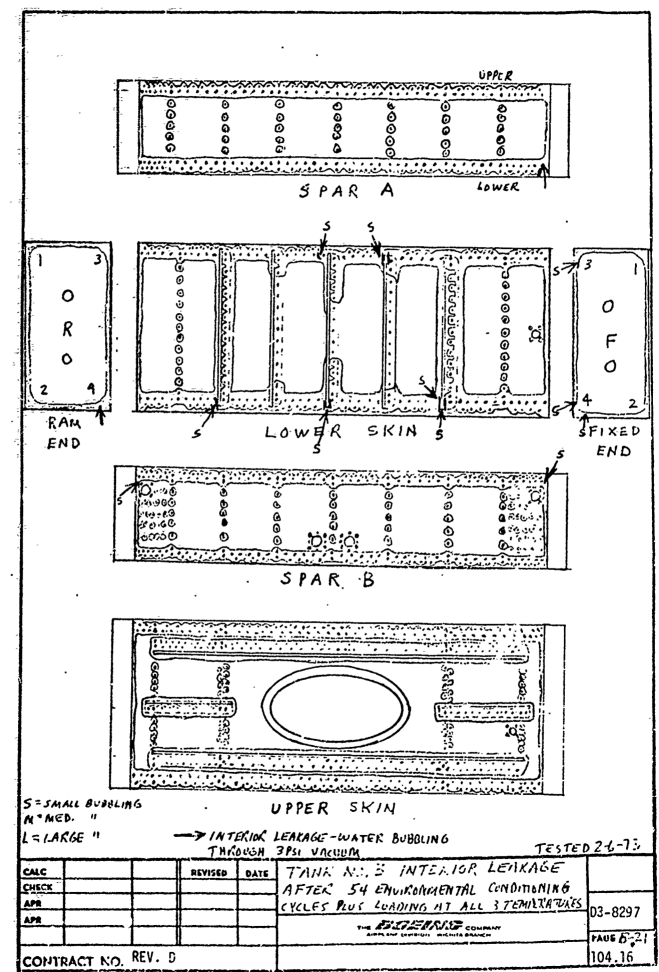
20

1480

王に、和川子にてたいました and the product of the second for きょうやく ţ 54 and the a



1 .



1 first

7

و بقبجة آ

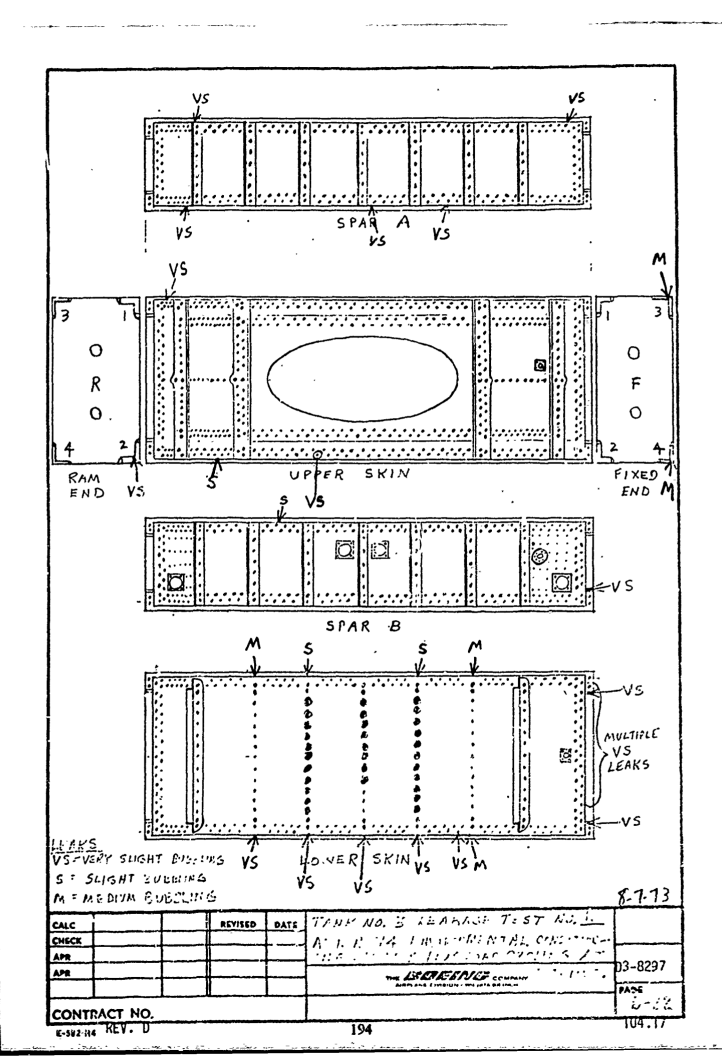
どういろんかい したんたい

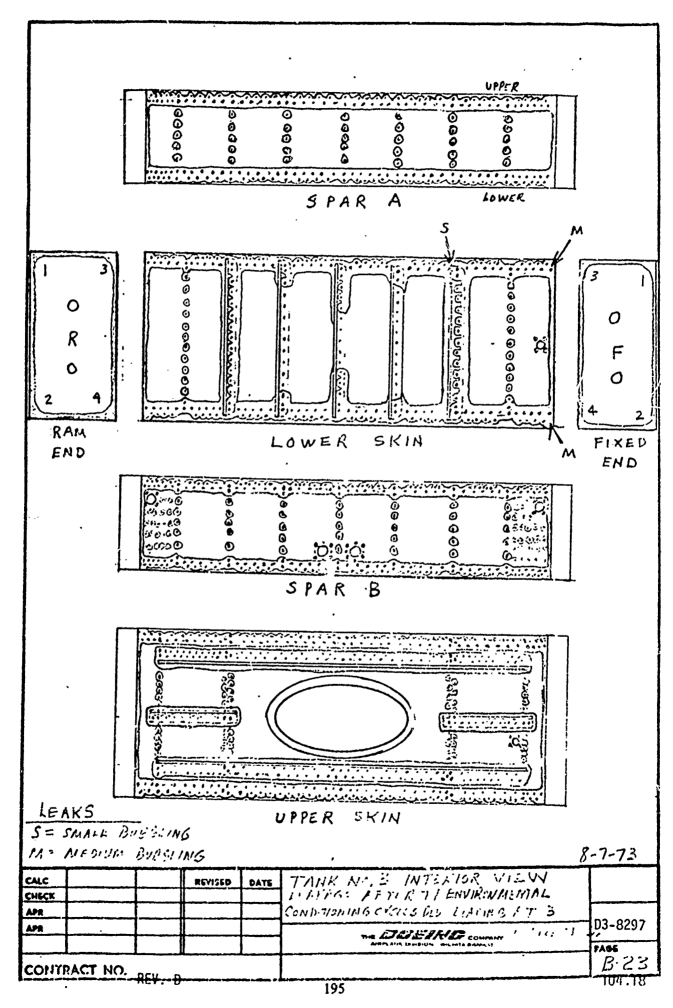
Active States

1

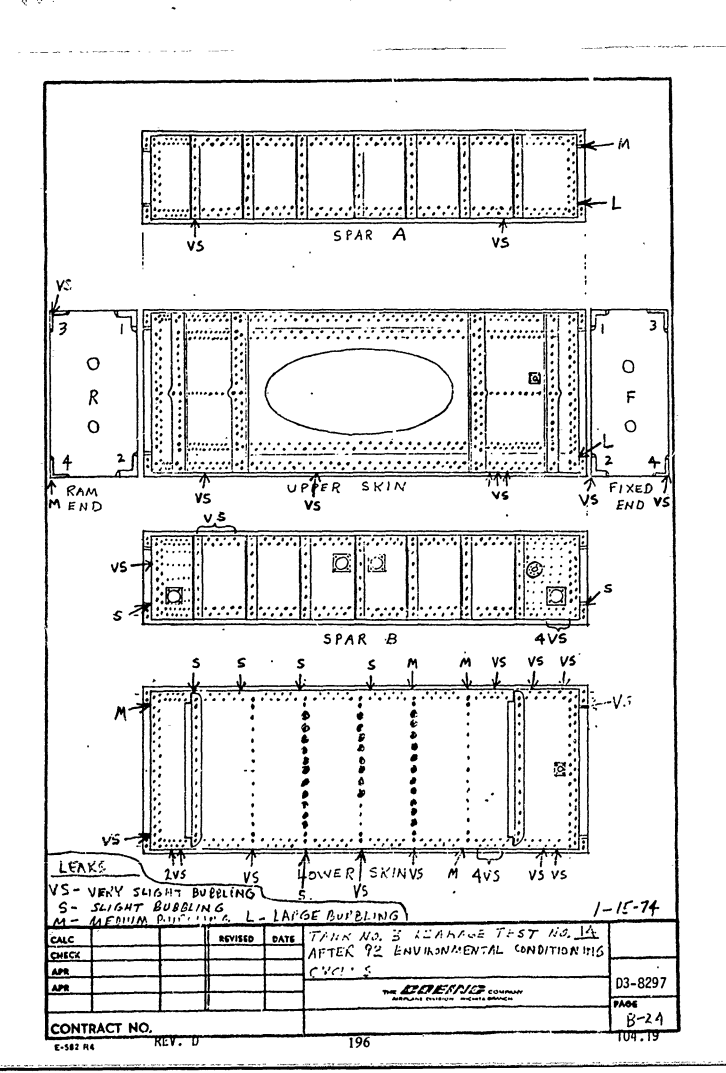
いるいないないというでもないないないない

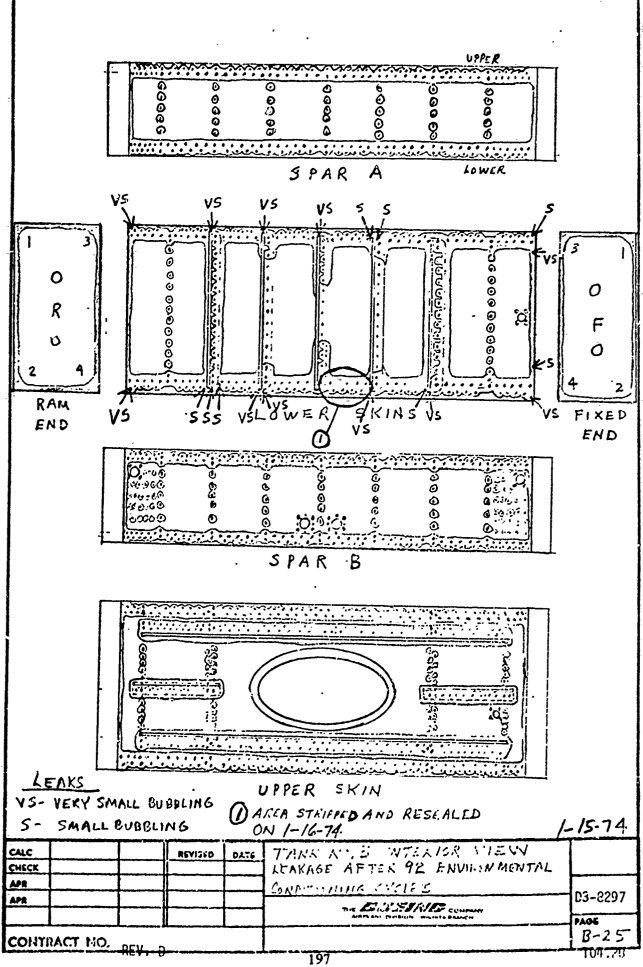
193

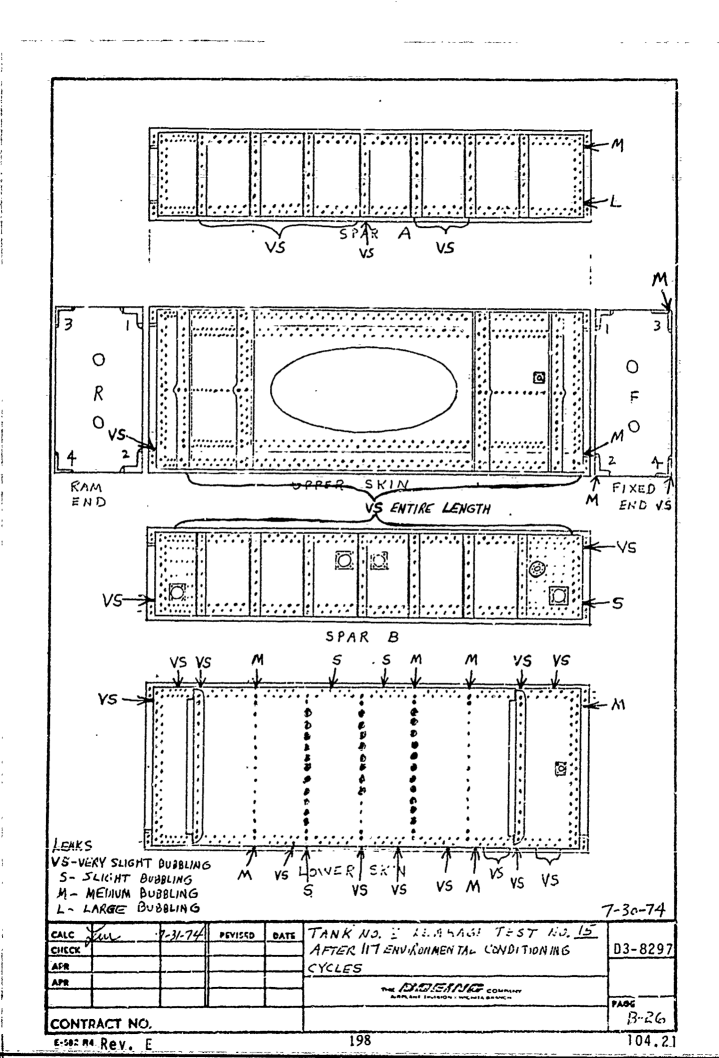




n n

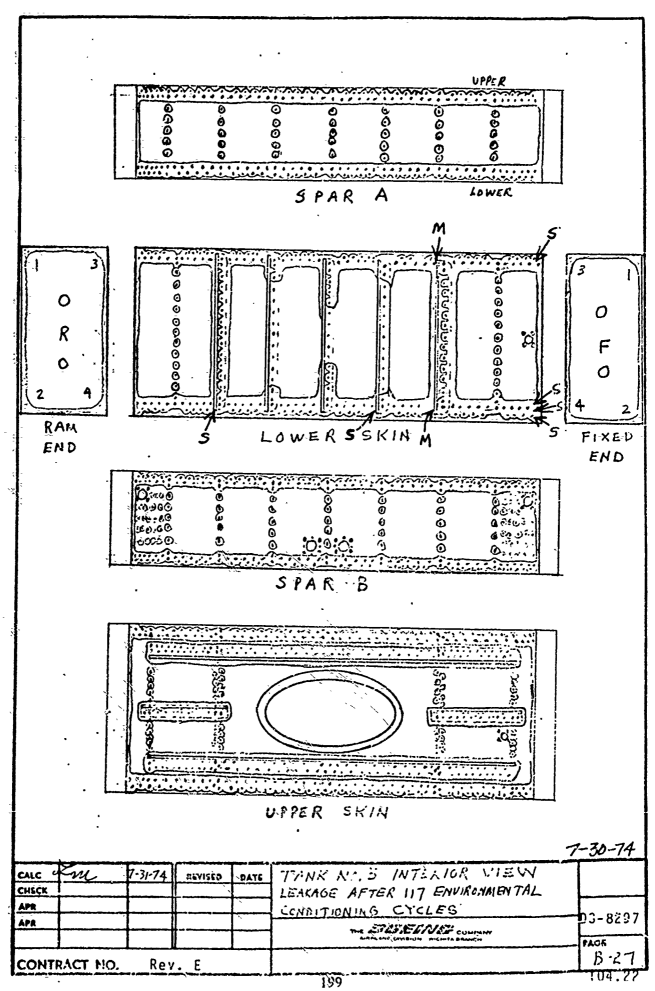


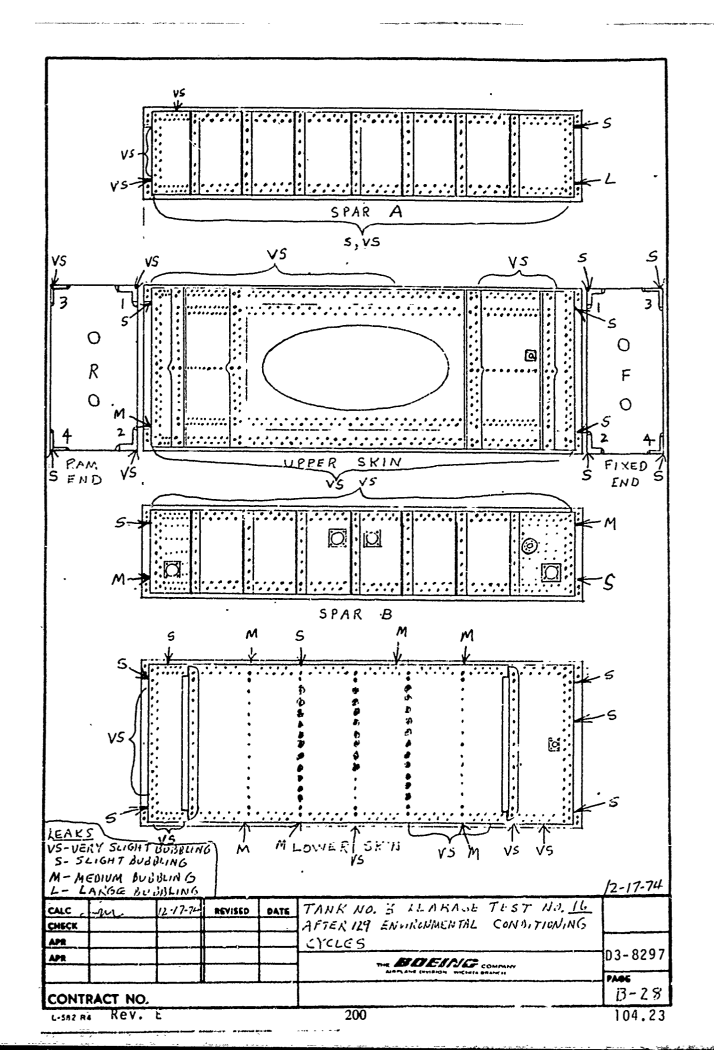




19 V

276.00





٠,

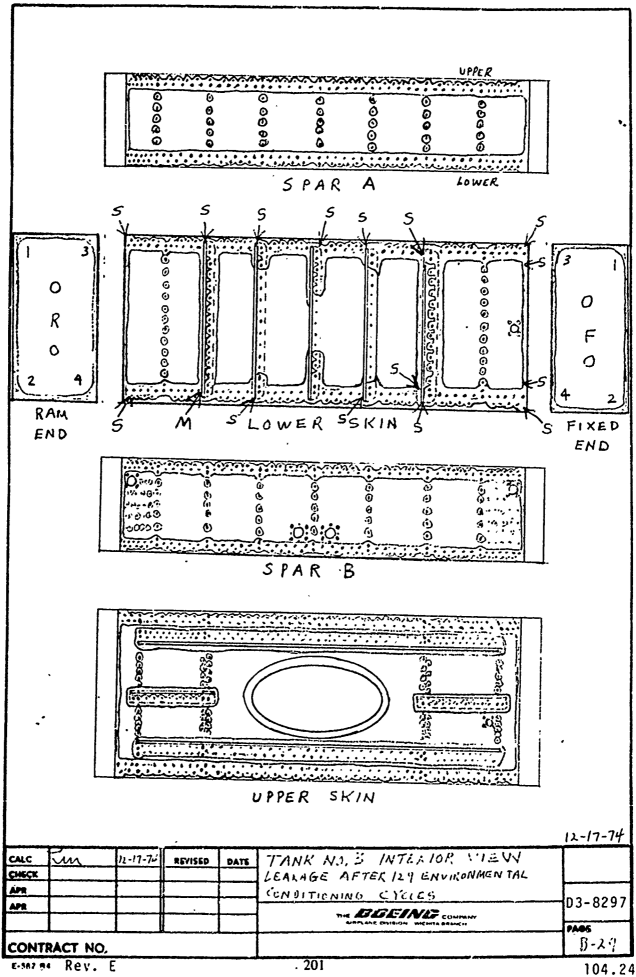
<u>ن</u>

· · ·

Circles And Circles X 2 7 1

.

 v_{s_i}



APPENDIX C

MEASUREMENT OF WEB SHEAR WRINKLES OF TANK NUMBER 3

POEINO	NO. D3-8297
SECT	PAGE 105

1

REVLTR: A

E-3033 R1

and shared a strength of the s

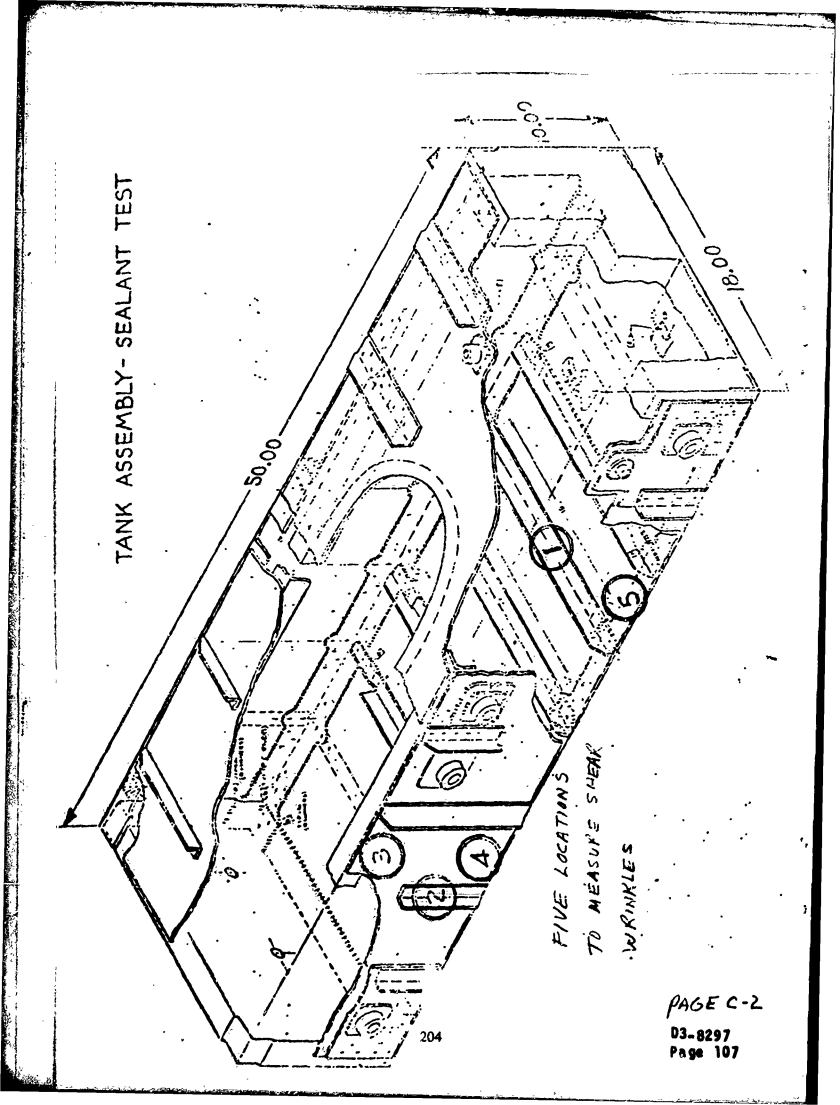
202

MEASUREMENT OF WEB SHEAR WRINKLES OF TANK NO. 3

Reference: Coordination Sheet No. SSI-Wing-661, Schuler to Beckmann, dated 4 November 1970, Subject: "Pad-Ups of Fuel Tank Shear Webs.

- Task A. Determine torque to obtain initial web buckling (estimated to be 52,000 inch-pounds).
 - 1. At 40,000 inch-pounds no wrinkling.
 - 2. At 50,000 inch-pounds very slight buckling.
 - 3. At 60,000 inch-pounds definite buckling.
- <u>Task B.</u> Determine extent and depth of web shear wrinkles at the five areas specified in the above reference. Measure at 100,000 and 216,000 inch-pounds torque in both directions.
 - This data is recorded in the attached 5 data sheets. Collecting and recording the desired information was difficult and, as can be seen, is confusing. It is suggested that Lyle Middleton (telephone 8-435-2337) be contacted if necessary for clarification.
 - 2. In general, the results show that at 216,000 inch-pounds torque, significant web wrinkles do extend into the pad-up areas and even into the chords and stiffeners.

Page C-1 D3-8297 Page 106



DEPTH OF WRINKLES ALONG LINES A THROUCH E WERE MEASURED WITH A 74 IN'H LONG STRAIGHT EDGE AT THE FOLLOWING LOADS: TORQUE WRINKLE DEPTH IN JOINT DEFLECTIONS LOAD, MILES AT LINE AT LINE INLB! A B'C D E C D O'C3 C)	
WERE MEASURED WITH A 74 IN'H LONG STRAIGHT EDGE AT THE FOLLOWING LOADS: TORQUE WRINKLE DEPTH IN JOINT DEFLECTIONS LOAD, MILS AT LINE AT LINE IN-LB! A B'C D E C D O' C3 C3 C3 C3 C3 C3 C3 C3 -100,000 C3. C3 C3 C3 C3 C3 -216,000 L 28 14 C3 7 C3 C3 -216,000 L 28 14 C3 7 C3 C3 +100,000 ~5 C3 C3 C3 C3 C3 +100,000 ~5 C3 C3 C3 C3 +100,000 L 28* 12* 2 16 C3 C3 -15 COUNTERCIDENTS TORQUE + 15 CLUCKWISE TORQUE L 15 EARGE C 19 LESS THAN WRINKLE MEASUREMENTS TAKEN FROM TANK EXTERIOR	* T	HESE VALUES TAKEN A	LONG LINES BAND	
WERE MEASURED WITH A $7\frac{1}{4}$ IN H LONG STRAIGHT EDGE AT THE FOLLOWING LUADS: TORQUE WRINKLE DEPTH IN JOINT DEFLECTIONS LOAD, <u>MILS AT LINE</u> AT LINE <u>IN-LB</u> : <u>A</u> <u>B</u> <u>C</u> <u>D</u> <u>E</u> <u>C</u> <u>D</u> <u>O'</u> <u>C</u> <u>C</u> <u>C</u> <u>C</u> <u>D</u> <u>C</u> <u>C</u> <u>C</u> <u>D</u> <u>C</u> <u>D</u> <u>C</u> <u>C</u> <u>D</u> <u>C</u> <u>D</u> <u>C</u> <u>C</u> <u>D</u> <u>C</u> <u>C</u> <u>D</u> <u>C</u> <u>C</u> <u>D</u> <u>C</u> <u>D</u> <u>C</u> <u>C</u> <u>D</u> <u>C</u> <u>D</u> <u>C</u> <u>C</u> <u>D</u> <u>C</u> <u>D</u> <u>C</u> <u>D</u> <u>C</u> <u>C</u> <u>D</u> <u>C</u> <u>C</u> <u>D</u> <u>C</u> <u>D</u> <u>C</u> <u>D</u> <u>C</u> <u>C</u> <u>D</u> <u>C</u> <u>C</u> <u>D</u> <u>C</u> <u>D</u> <u>C</u> <u>C</u> <u>C</u> <u>D</u> <u>C</u> <u>C</u> <u>C</u> <u>D</u> <u>C</u> <u>C</u> <u>D</u> <u>C</u> <u>C</u> <u>C</u> <u>C</u> <u>D</u> <u>C</u> <u>C</u> <u>C</u> <u>C</u> <u>D</u> <u>C</u> <u>C</u> <u>C</u> <u>D</u> <u>C</u> <u>C</u> <u>C</u> <u>D</u> <u>C</u> <u>C</u> <u>C</u> <u>C</u> <u>C</u> <u>C</u> <u>D</u> <u>C</u> <u>C</u> <u>C</u> <u>C</u> <u>C</u> <u>C</u> <u>C</u> <u>C</u> <u>D</u> <u>C</u> <u>C</u> <u>C</u> <u>C</u> <u>C</u> <u>C</u> <u>C</u> <u>C</u> <u>C</u> <u>C</u>	WRINKLE ME	ASUREMENTS TAKEN FRO	M TANK EXTERIOR	EAS
WERE MEASURED WITH A $7\frac{1}{4}$ IN H LONG STRAIGHT EDGE AT THE FOLLOWING LUADS: TORAVE WRINKLE DEPTH IN JOINT DEFLECTIONS LOAD, <u>MILS AT LINE</u> AT LINE <u>IN-LB</u> : <u>A</u> <u>B</u> <u>C</u> <u>D</u> <u>E</u> <u>C</u> <u>D</u> <u>O</u> <u>C</u> <u>C</u> <u>C</u> <u>D</u> <u>O</u> <u>C</u> <u>C</u> <u>C</u> <u>C</u> <u>C</u> <u>O</u> <u>C</u> <u>C</u> <u>C</u> <u>C</u> <u>O</u> <u>C</u> <u>C</u> <u>C</u> <u>C</u> <u>O</u> <u>C</u> <u>C</u> <u>C</u> <u>C</u> <u>C</u> <u>C</u> <u>D</u> <u>C</u> <u>C</u> <u>C</u> <u>D</u> <u>C</u> <u>D</u> <u>C</u> <u>D</u> <u>C</u> <u>C</u> <u>D</u> <u>C</u> <u>D</u> <u>C</u> <u>C</u> <u>D</u> <u>C</u> <u>C</u> <u>D</u> <u>C</u> <u>C</u> <u>D</u> <u>C</u> <u>C</u> <u>D</u> <u>C</u> <u>C</u> <u>D</u> <u>C</u> <u>D</u> <u>C</u> <u>C</u> <u>D</u> <u>C</u> <u>C</u> <u>D</u> <u>C</u> <u>D</u> <u>C</u> <u>D</u> <u>C</u> <u>C</u> <u>D</u> <u>C</u> <u>C</u> <u>D</u> <u>C</u> <u>C</u> <u>D</u> <u>C</u> <u>C</u> <u>D</u> <u>C</u> <u>C</u> <u>D</u> <u>C</u> <u>C</u> <u>D</u> <u>C</u> <u>D</u> <u>C</u> <u>C</u> <u>D</u> <u>C</u> <u>D</u> <u>C</u> <u>C</u> <u>D</u> <u>C</u> <u>C</u> <u>D</u> <u>C</u> <u>C</u> <u>C</u> <u>D</u> <u>C</u> <u>C</u> <u>D</u> <u>C</u> <u>C</u> <u>D</u> <u>C</u> <u>C</u> <u>C</u> <u>D</u> <u>C</u> <u>C</u> <u>C</u> <u>D</u> <u>C</u> <u>C</u> <u>C</u> <u>C</u> <u>D</u> <u>C</u>		S COUNTERCIDERWISE TORA	IVE	
WERE MEASURED WITH A 74 IN H LONG STRAIGHT EDGE AT THE FOLLOWING LUADS: TORQUE WRINKLE DEPTH IN JOINT DEFLECTIONS LOAD, MILS AT LINE AT LINE IN-LB: A B'C D E C D O' C3 C3 C3 C3 C3 C3 C3 C3 I-100,000 C3. C3 C3 C3 C3 C3 C3 C3	-216,	000 L 28 14 <3 7	 	
WERE MEASURED WITH A 74 IN H LONG STRAIGHT EDGE AT THE FOLLOWING LUADS: TORQUE WRINKLE DEPTH IN JOINT DEFLECTIONS		ត្រូតត្រួត	$ \begin{array}{c} \mathbf{C} \mathbf{D} \\ \hline 3 \overline{3} \\ 3 \overline{3} \end{array} $	
	STR	NIGHT EDGE AT THE FI	JOINT DEFLECTIONS	

1 -と し

States and a state of the states of the

50

AND REAL BORN

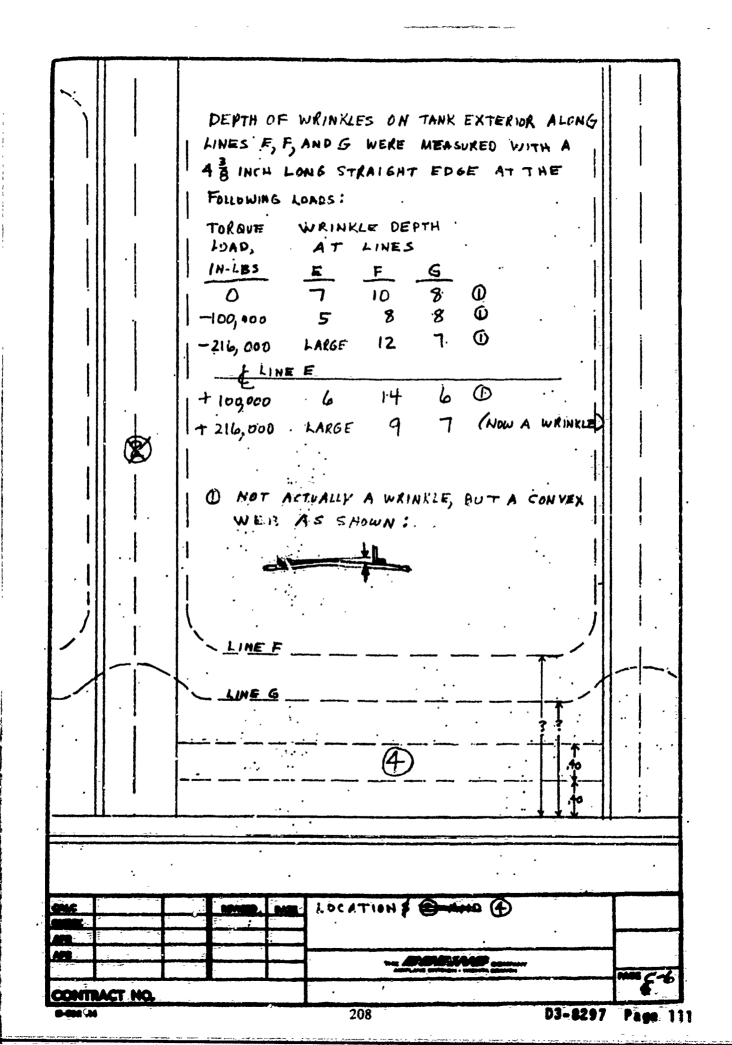
the second state of the second state of the second state of the

Î

.

10 Y Z DEPTH OF WRINKLES ON "ANK EXTERIOR ALONG LINES A, B, AND C WERE MEASURED WITH A 74 INCH LONG STRAIGHT EDGE AT THE FOLLOWING LOADS: JOINT DE FLECTION TORQUE LOAD WRINKLE DEPTH LINES AT STIFFE IN-LBS AT 展 C C 3 <3 <3 3 - 100,000 -**く**ろ **<**3 < 3 LARGE 8 8 -216,000 <3 <3 **<**3 +100,000 -くう ----~ 5 15 **<3**_ +216,000 LARGE 46 0 LOCATION \$ 2 AND CNC S JAMO CONTRACT NO. 206 03-8297 Pane 1

3 LINE A LINE B DEPTH OF WRINKLES ON TANK EXTERIOR ALUNG LINES A AND B WERE MEASURED WITH A A 3 INCH LONG. STRAGHT EDGE AT THE FULLOWING LOADS: TORQUE LOAD, WRINKLE DEPTH IN IN-LBS MILS AT LINE _ **B** A < 3 < 3 O 1 ¢ < 3 くら -100,000 6 10 -216,000 < 3 ٢3 +100,000 <3 + 216,000 5 CALC LOCATION (3) M · · · . NTRACT NO. 207 74.97 1.00



5 6 Z

DEPTH OF WRINKLES ON TANK EXTERIOR ALONG LINES H, J, AND K WERE MEASURED. WITH A 4 TINCH LONG STRAIGHT EDGE AT THE FOLLOWING LOANS: WRINKLE DEPTH IN TORQUE Ø LOAD, " MILS AT LINE IN-LBS <u>H</u><u>J</u><u>K</u> 3 3 3 3 3 <3 <3 0 **3**. 1 -100,000 L 5.4. -216,000 + 100,000 13* 9* 0 + 216,000 LINE H I NOT A WRINKLE, BUT CONVEX WEB AS SHOWN : - 32-4-45-* THESE VALUES TAKEN FROM OTHER SIDE OF STITFENER AT LEFT. LINE J LINE K 6 LOCATION & CHARD (VIEWED FROM TANK EXTERIOR) 6 rt No 209

-7

	REVISIONS		
LTR	DESCRIPTION	DATE	APPROVED
۸	To add additional data to the documentcovers work conducted from the beginning of the program August 1969 until cancellation March 25, 1971.	4-8-7 \	2. middleton RBallier
B	To add additional data to the documentcovers w conducted from August 1971 to December 23, 1971.	r ^k 1-5-7:	X middleton
C	To add work conducted from December 23, 1971 to January 31, 1972.	1-31-7	2. Middleton
D	To add work conducted from February 1, 1972 to January 1974.	2-18-74 (2. Widdliton
E	To add work conducted from January 1974 thru March 1975.	4-16-7	L'millitan
F	To add data to Table 6 (inadvertently omitted fr Revision E).	om 4 4-24-75 2	Middliton Pa Billion
+302	RI BOEING	NO. D3	-8297
	210	PAGE	

ς,

ł

0

)

			CHAN	ge re	CORD F	PAGE.		RE	V R F
SCOT		PAGES		REV			PAGES		REV
SECT	REVISED	ADDED	DELETED	LTR	SECT	REVISED	ADDED	CELETED	LT
		nent has b					104.18 104.19 104.20		D
	completely revised		•	A		1 3 5 6 7	57.11 75.1 83.8 83.9 83.10	3.1	
	1,8,49, 52,54,56, 57,81,82, 83,96,97, 113, 114	3.1 49.1 57.1 83.1 104.1 104.2 104.3		В		8 57.10 83.6 83.7 101 113 114	104.21 104.22 104.23 104.24		E
		104.3 104.4 104.5 104.6				1 82 113 114			F
	1,3.1,8, 17,53,76, 82, 113 & 114	57.2 104.7 104.8 104.9		с					
		57.3 57.4 57.5 57.6 57.7 57.8 57.9 57.10 83.2 83.3 83.4 83.5 83.6 83.7 104.10 104.11 104.12 104.13 104.14 104.15 104.16 104.17		D					
EVLTI	R: F			2	11	BOEN SECT	NO. D PAGE	93-8297 114	

1. Constant of the second s

ç

a second how the second s

APPENDIX B

Υ.

4

Cr.

LONG-LIFE, EASILY APPLIED FLUOROCARBON SEALANTS FOR SUPERSONIC AIRCRAFT FUEL TANKS

Final Report

Ьу

R. Gilliland P. Mallard R. Schubert L. Morris

Prepared under Contract No. Y555369-5835N

for

The Boeing Company Commercial Airplane Group Seattle, Washington

Products Research & Chemical Corporation

Burbank, California

PRECEDING PAGE BLATS.NOT FILMED

her weed

TABLE OF CONTENTS

Dane

		rage
١.	INTRODUCTION	1
11.	SUMMARY OF WORK	3
	CONCLUSIONS AND RECOMMENDATIONS	4
١٧.	PHASE I - IMPROVEMENT OF APPLICATION PROPERTIES	5
	A. Reduction of Viscosity of Current Type Materials	5
۷.	PHASE IIA - IMPROVING THE LIFE OF FLUOROCARBON SEALANTS	17
	A. Elimination of Metallic Impurities	17
	B. Antioxidants .	17
	C. Metallic Soaps	17
	D. Acid Scavengers	17
	E. Free Radical Scavengers	17
	F. Investigation of Cure Mechanisms on Stability	18
	G. Pre-extruded Sealant	20
۷١.	RAW MATERIALS	29

VII. PHOTOGRAPHS, Plates No. 1, 2 and 3

I. INTRODUCTION

The historical development of aircraft fuel sealants begins with the change from bladder-type fuel cells to integral wing tanks. The need for an easily applied sealing compound to resist the fuel, flexing and the temperatures encountered was met first by polymeric alkyd resins. These proved to be limited in both stability and application properties. Materials based upon polysulfide polymers initially used dispersions in water which could be made to fuse together by means of "chemical sufteners." The high shrinkage of such materials and the development of liquid, 100% solids, mercaptan-terminated, polysulfide polymers offered a new formulating tool. The standard lead peroxide-cured polysulfides, however, still suffered from attack by fuel. With the invention of the dichromate-cured polysulfides by Products Research & Chemical Corporation, and later the manganese dioxide cured materials, a truly fuel-resistant material was available for aircraft sealing. These materials are, in fact, the basis for sealing nearly all subsonic commercial and military aircraft in the worid today.

The United States Air Force in the mid-1950's recognized, however, that the development of supersonic aircraft would require new polymer developments to produce materials beyond the temperature capabilities of the polysulfide rubbers. Products Research, at that time, under WADD 58-59, showed that fluorocarbon elastomers as well as liquid fluorosilicones had significantly improved high-temperature fuel resistance. However, the development of the fluorocarbons has been hampered by their poor low-temperature properties, their unavailability in high solids liquid forms, and finally by concern over stress corrosion effects on titanium alloys at high temperatures.

As a result of these difficulties, the commercial development of hightemperature aircraft sealants has taken place with the fluorosilicones. These products are not without their problems, however: Even stabilized trifluoropropyl polysiloxanes are subject to reversion when confined, so that long-time retention of properties, especially required for commercial supersonic craft, is not available.

As a result of these limitations of the fluorosilicones, the Air Force has funded programs in which the fluorocarbons are chemically copolymerized with the fluorosilicones to give products of improved reversion resistance while retaining adequate low-temperature properties. The Boeing Company has extensively tested such materials and finds them significantly better than the unmodified fluorosilicones for their requirements. They still show thermal deterioration, however. Likewise, the introduction of major fluorocarbon segments into any polymer gives concern with stress corrosion of titanium. This has led to use of the fuel-resistant cyanosiloxanes in some specialty applications.

In the past fifteen years, work has continued with efforts to produce more tractable sealants based on fluorocarbon backbones as well as polymers with heterocyclic nitrogen, etc. Wright Pattrrson Air Force Base has formulated a number of sealants based upon low molecular weight, vulcanizable fluorocarbons. Early materials showed varying degrees of stress corrosion with titanium. With the introduction of the fluorocarbon, ECD-487, a DuPont product, a material was available which was shown by the Boeing Company to have improved low-temperature properties along with retention of hightemperature properties and absence of stress corrosion. When this polymer was made into a sealant along the lines of the Air Force work, high shrinkage and bubbling was encountered.

ł

The purpose of this project was to find means of producing practical sealants from available low-temperature fluorocarbons which would be expected to perform well in the supersonic transport designed by the Boeing Company.

II. SUMMARY OF WORK

All goals for the fluorocarbon sealant and tape were met with the exception of slightly lower volume percent solids in the case of the sealant.

The sealant developed under this program is greatly superior in application properties to all known solvent-based fluorocarbon materials, has a volume shrinkage less than one-half that of the solvent-applied types, is safe to use from a flammability standpoint and is much easier to apply and tool. The discovery of a water miscible, "reverse phase," technology operable with fluorocarbon dispersions offers wide opportunities for room temperature fusions of otherwise difficult nonfusible, cracking latices.

The tape developed under the program, based upon ECD-487, has a much wider range of temperature usefulness than the standard Viton B materials.

Two different approaches for producing a practical fluorocarbon sealant were followed. Phase I had as a goal the formulation of an 80 volume percent solids material based on ECD-487 which had easy application properties and a useful life at 550°F--twice that of current materials. The second approach employed ECD-487 in a soft, self-vulcanizing tape form at high solids.

Work undertaken to improve sealant application properties and increase volume percent solids by means of pigment reduction and introduction of low molecular weight modifiers was of minor value. Solvent dispersion studies, especially through use of Viton latex sealants, give materials which closely approach the target goal of 80% solids by volume and greatly improved application properties. A volume percent solids of 70%, with weight percent

solids of 83%, was obtained. Low-temperature properties are intermediate between Viton B and ECD-487. Further research to improve stability of the sealants and adhesion to titanium is desirable.

Pre-extruded sealants were developed with not more than 10% by volume of volatiles. These sealants have excellent low-temperature properties, being prepared from ECD-487, and are stable when stored at moderately low temperatures. Methods have been developed for adhering these sealants to titanium. Adhesion to titanium was retained after considerable dry heat exposure.

A number of materials were investigated to improve the life of fluorocarbon sealants. One material, Vanstay 8050, was found to be effective in improving the life of solvent-dispersed and pre-extruded sealant.

III. CONCLUSIONS AND RECOMMENDATIONS

A quantum improvement in application properties of fluorocarbon-based sealants has been achieved by the discovery of means for producing room temperature fusible fluorocarbon latices. Combined with the availability of better low-temperature polymers, serious consideration should be given to such materials in supersonic aircraft sealing applications. Retention of good elastomeric properties at 500°F or lower for long periods was observed.

The pre-extruded tape formulations based upon ECD-487 show generally good temperature range of performance and have merit for configurations where high shrinkage or reversion in confined situations is of concern--such as faying surfaces.

Further work in the development of low temperature fusing fluorocarbon latices would be merited in the areas of evaluation of other fluorocarbon latex polymers, improved primer developments for both sealants and tapes, and exploration of the effects of pH, stabilizers and curing mechanisms on the long-term, high temperature retention of physical properties.

111. PHASE 1 - IMPROVEMENT OF APPLICATION PROPERTIES

The major limiting factor in the current study of fluorocarbon sealants is their difficult application which results in slow extrusion rates, high shrinkage and entrapped bubbles. The current Viton ECD-487 sealant has only 63.5% solids by weight, or 43.3% solids by volume. The goal of this phase is to prepare a one- or two-component thermosetting sealant which has the basic heat-, fuel- and low-temperature-resistance of the current Viton ECD-487 formulation, but has a volume percent solids of at least 80% and a room temperature viscosity of no more than 35,000 poise.

A. Reduction of Viscosity of Current Type Materials

1. Pigment Reduction

Viton ECD-487, the primary polymer of interest, is inherently quite strong compared with other commonly used fuel tank elastomers. The use of pigmentation by carbon black could be dispensed with if its elimination would increase solids content. The effect of elimination and increasing carbon black content is shown below.

	2211-22	2211-23	2211-36
Viton ECD-487	100.0	100.0	100.0
Maglite Y	15.0	15.0	15.0
Thermax		15.0	60.0
FS-1265	8.0	8.0	8.0
EDA DIBK Ketimine	5.5	5.5	5.5
MEK	78.0	81.5	94.5
Extrusion rate @ 90 psi (1/8" diameter nozzle)	38 gms/min.	26 gms/min.	38 gms/min.
Solids Content			
By weight	62.3%	63.9%	66.6%
By volume	42.0%	43.6%	46.8%
Tensile Strength, psi/ Elongation, %			
Cured 24 hrs. @ 350°F	240/1500	270/1200	250/380
Aged 7 days @ 500°F	1420/1000	1400/950	950/600
Aged 35 days @ 500°F	900/150	1300/120	1360/20

As a nonreinforcing grade of carbon black, Thermax, a medium thermal type, was used, its variation produced little effect on viscosity. Increasing carbon black loading gave a marginal increase in volume percent solids over the basic formula, 2211-23, but lowered elongation and heat resistance. The elimination of carbon black decreased volume percent solids. Note, however, the great improvement in elongation retention with reduction of carbon black.

2. Solvent-Dispersion Studies

Frequently, with high polymer solutions, it is possible to prepare colloidal dispersions which exhibit lower viscosities than true solutions of the polymer. This is based on the principle that molecular aggregates can present less interaction than molecularly dispersed molecules because of their lower surface area. A good example of this principle is found in the preparation of rubber cements where hydrocarbon solutions of rubber exhibit high tack and "web-like" character. Introduction of small amounts of the nonsolvent ethyl alcohol produce coiling of the rubber chains and a reduction in the viscosity and webbing occurs. This pheromena was first investigated by choosing solvents or solvent blends which give appropriate solubility parameter (d) and hydrogen bonding index (Y) for solution of Viton ECD-487. Solutions were prepared of 15 grams ECD-287 per 200 cc of solvent and viscosity determined with the Brookfield Viscometer, spindle #3 at 10 rpm. Results are shown below:

Solvents	<u></u>	<u> </u>	<u>Viscosity(poise</u>)
MEK	9.30	540	3.6
Ethyl acetate	9.10	5.20	3.4
Tetrahydrofuran	9.10	5.50	3.9
90% MEK, 10% toluene	9.25	5.18	4.1
80% MEK, 20% toluene	9,22	4.97	4.0
70% MEK, 30% toluene	9.15	4,83	5.5
90% MEK, 10% methanol	9.80	5.70	5.3
70% MEK, 30% methanol	10.85	6.33	4.2

Solvents	<u> </u>	$\underline{\gamma}$	<u>Visccsity(poise</u>)
50% MEK, 50% methanol	11.90	6.95	5.8
80% ethyl acetate, 20% methanol	10.18	6.10	3.8
80% tetrahydrofuran, 20% methanol	10,18	6.10	4.0
40% MEK, 30% methanol, 30% toluene	10.74	5.70 [°]	4.5
60% MEK, 20% methanol, 20% toluene	10.28	5.60	4.0
20% MEK, 40% methanol, 40% toluene	11.20	5.48	8.6

Lowest viscosity was obtained with the pure solvents. Blending a nonsolvent with the solvent generally increased viscosity. By choice of the appropriate solubility parameter and hydrogen-bonding coefficients, solutions were obtained with as little as 20% solvent, with 80% nonsolvent, but at a considerable sacrifice in viscosity. Later work with water and amides gave better results.

3. Introduction of Low Molecular Weight Modifiers

The preferred materia! for modifying ECD-487 to reduce viscosity would be a low molecular weight version of ECD-487. The DuPont Company had sampled Wright-Patterson AFB with such a material, but was unable to provide samples for this project. Other low viscosity Vitons available are grades LM and LD-011. These materials do not have the desirable low-temperature flexibility exhibited by ECD-487 or its low molecular weight version. However, it was felt that small quantities could be incorporated into the sealant without materially affecting low-temperature properties. In addition, DuPont supplied samples of LD-487-LV, a lower Mooney version of ECD-487, and VTX-3518, also lower Mooney, but intermediate in low-temperature properties between ECD-487 and Viton A. The following results were obtained:

	2211-17	2211-44	2211-51	2211-56
ECD-487	50	~-		
Viton LM	50			
LD-487LV		100		
LD-011		~~	100	
VTX-3518				100
Maglite D	3			
Calcium hydroxide	6			
Maglite Y		15	15	15

	2211-17	2211-44	2211-51	2211-56
Thermax	20	15	20	15
FS-1265		8	8	8
Hydroquinone	1	*-		
N,N-Dimethyl dodecylamine	2			
MEK	37	70		62.5
EDA-DIBK ketimine		5.5		5.5
Shell H-3		~ •	5	
Methyl Isobutyl Ketone			13,6	
Solids Content, by weight	77.5%	67.0%	91.5%	69.7%
Extrusion Rate @ 90 psi (1/8: diameter nozzle)	6.2 gms/min.	22.4 gms/min.	10.0 gms/min.	17.2 gms/min.
Tensile Strength, psi/ Elong∉tion, %				
Cured 24 hrs. @ 350°F Aged [.] 7 days @ 500°F	250/1450 	280/850 1000/530	 brittle	800/1000 960/600

The LD-487LV and VT-X-3518 gave only marginal improvement in solids content. With LD-011 Sealant Grade Fluoroelastomer, a 91.5% solids sealant was obtained, but the sealant bubbled badly and became brittle after seven days' oven-aging at 500°F. The ECD-487/LM blend was intermediate between the LD-011 and ECD-487 in solids content. A very low state of cure was obtained with the hydroquinone/ amine curing system used.

4. Viton Latex Sealants

Viton B is also available in a latex form designated Viton L-31 Aqueous Fluoroelastomer Dispersion. This material is supplied at 65% solids at a viscosity of 150 centipoise. This is an extremely low viscosity compared with solvent dispersions of the same solids content of about three million centipoise. Much higher solids content and lower shrinkage are possible if the material can be thickened to a sealant viscosity. While Viton B does not have the desirable low-temperature properties of ECD-487, this work was undertaken to devise means of preparing a latex sealant. The technology developed could be readily transferred to an ECD-487 type latex when such becomes available.

In preliminary experiments, merely thickening the latex to a paste consistency resulted in a material which cracked on drying. Blends of the L-31 latex with a polycyanosiloxane fluid improved handling properties. The addition of acid acceptors such as magnesium oxide, zinc oxide, litharge and Dyphos destabilized the latex. A 50-50 blend of L-31 latex and Viton LM dissolved at 80% in methyl isobutyl ketone was prepared and cured with Maglite Y and hexamethylene diamine. The cured product was not well knit and showed cracks on bending. Dimethyl formamide was then used as the solvent for Viton LM and this material was easily handled and applied as a sealant and dried at room temperature to a good elastomer. Again, the use of conventional acid acceptors destabilized the system. A surface-treated calcium carbonate, OMYA-BLH, which is neutral in pH, was found to be a satisfactory acid acceptor in the Viton L-31 latex system. The incorporation of Viton LM was eliminated because of its relatively poor heat and corrosion resistance. Modifiers such as fluorosilicone oil or polycyanosilosane fluid were required to obviate cracking of the film. The following formulations were prepared:

- -	2211-41	2211-42
Viton L-31 Latex	154.0	154.Q
Dimethyl acetamide	15.0	15.0
Polycyanosiloxane fluid		8.0
FS-1265 fluorosilicone oil	8.0	****
OMYA-BLH	20,0	20.0
Distilled water } dispersion	19.6	19.6
Witco 900	0.4	0.4

Degassed to paste consistency (vacuum removal of water to solids-indicated)

Curing agent - 20% aqueous solution hexamethylene diamine - 3.5 parts

Weight % solids		82.4%	83.0%
Volume % solids		70.0%	70.5%
Aging at 500°F:	42 days	flexible	flexible
	56 days	flexible	flexible
	84 days	weak and short	brittle

Films were cured three days at 75° F, one day at 140° F, one day at 200° F and one day at 350° F prior to 500° F exposure. Best heat resistance was obtained with the fluorosilicone modifier.

An attempt was made to produce a formulation containing ECD-487 as an external phase with internal latex. This resulted in coagulation and a stable sealant could not be produced. Emulsions of ECD-487 in water and methyl isobutyl ketone were prepared at 16%, 21% and 30% solids. Attempts to prepare a late~ sealant from these emulsions resulted in coagulation. Attempts to produce a latex from the emulsion by selective absorption of the methyl isobutyl ketone were unsuccessful.

The formulations below show the effect of decreasing calcium carbonate content and of substituting alumina for calcium carbonate. Increasing calcium carbonate from 20 to 30 parts caused cracking of the cured film.

	<u>2211-53</u>	2211-69	2211-72
Viton L~31 Latex Dimethyl acetamide FS-1265 OMYA-BLH	154.0 15.0 8.0 20.0	154.0 15.0 8.0 10.0	154.0 15.0 8.0
Tabular alumina, 325 mesh Distilled water Tamol 731 Witco 900		2.4 0.4 0.1	20.0 9.0 0.8 0.2

Degassed to paste consistency (vacuum removal of water to solids indicated) Curing agent - 20% aqueous solution hexamethylene diamine - 3.0 parts

Weight % solids	83.8%	83.2%	82 .6%
Storage Stability: @ 75°F	30 days		
@ 32°F	10 months		
Aging of cured films:			
Tensile Strength, psi/Elongation	. %		
14 days @ 500°F	590/310	560/360	450/470
21 days @ 500°F	410/310	650/300	
35 Jays @ 500°F	330/200	290/240	310/410
49 days @ 500°F	390/160	245/220	210/320
70 days @ 500°F	flexible	flexible	flexible

Reduction of calcium carbonate or use of alumina improved elongation after heat exposure, but caused corrosion of titanium to occur more rapidly With alumina filler, a rather thermoplastic material was obtained, and it is apparent that alumina is not a satisfactory acid acceptor in Viton latex.

Samples of experimental low-temperature latexes VT-X-3627 and VT-X-3627B were obtained from DuPont. These polymers were said to be intermediate in lowtemperature flexibility between ECD-487 and Viton B. The VT-X-3627B was a minor modification designed for potentially improved stability. These latexes were compounded into sealants in a formulation similar to 2211-53 (shown above) except that M-Pyrol (N-methyl, 2 pyrolidone) was used in place of dimethyl acetamide since subsequent work on L-31 latex indicated that the former imparted improved application and stability properties.

Use of the centrifuge was investigated to concentrate the later. Excessive localized hardening occurred unless the centrifuge was run at much slower speeds for longer periods. Concentration of the later by decanting excess water after the later was allowed to settle for several weeks was a more feasible method. Additional wetting agent was added in this case to compensate for that lost in the decanting process.

Preliminary VT-X-3627 and VT-X-3627B formulations were less stable than the Viton L-31 formulations. The former coagulated in a few days at room temperature. Storage at 0°F also resulted in coagulation. A temperature of approximately 32°F was found to be the optimum for storage.

Alkaline materials generally used as acid acceptors in Viton caused coagulation of the VT-Y-3627 and VT-X-3627B as well as the Viton L-31 latex.

A preliminary formula was prepared without concentrating the latex. Aging tests were conducted at 500°F, comparing two levels of curing agent.

221	-7!	5

•	IL W JARY COLON	104.0	
	M-Pyrol	15.0	
	FS-1265	8.0 .	
	OMYA-BLH	20.0	
	OMYA-BLH Distilled Water	4.8	
dispersion -	Tamol 731	0.8	
	Tamol 731 Witco 900	0.2	
	Degassed to paste		m removal of water lids indicated)
	Curing Agent - 20% he>	& aqueous solution wamethylene diamine	
	Weight % solids	82 . 2%	
Aging of Cured Films		Curing	Agent
		three parts	four parts
Tensile Strength, psi/ Elongation, %	/		
Initial - cured 1 day	@ 150°F plus		
l day @ 250		65/150	110/270
Initial - cured additi	ional I dav		
@ 350°F		90/500	150/400
Aged 7 days @ 500°F		280/520	330/370
" 30 days @ 500°F		260/340	320/260
" 60 days @ 500°F		220/260	260/120
•			

164.0

A

VT-X-3627 Latex

Initial tensile strength and elongation were quite low until curec °50°F. The lower tensile strength in the sample cured with three parts curing agent was due to greater porosity. The lower amine concentration did improve aging, as noted by the superior elongation after 60 days' exposure at 500°F.

In order to determine the practicality of sealing aircraft with Viton latex sealants, corner sections were constructed of titanium, primed and fillets of Viton latex extruded into the seams. Initial efforts, utilizing sealants such as 2211-75 above, resulted in the formation of large cracks at the corner as the latex dried. The formula was modified by increasing the content of FS-1265 fluorosilicone oil from eight to twelve parts. With this modification, no cracking occurred. (See Plate No. 1)

In an effort to improve stability of the VT-X-3627 and VT-X-3627B sealants, calcium carbonate was omitted from the latex. A ball milled dispersion of calcium carbonate together with hexamethylene diamine was used as the curing agent. Formulations and test results are shown below:

	2211-130	2211-131	2211-132	2211-133
	Part B	Part B	Part B	Part 8
VT-X-3627 Latex	164.0		164.0	
VT-X-3627B Latex		164.0		164.0
M-Pyrol	15.0	15.0	15.0	15.0
Witco 975	1.5	1.5	*****	۲۵ امی جغه همه های
FS - 1265	12.0	12.0	12.0 .	12.0
	<u>Part A</u>	Part A		
OMYA-BLH	20.0	20.0	20.0	20.0
Witco 900	0.2	0.2	0.2	0.2
Tamol 731	0.4	0.4	0,8	0.8
M-Pyroi	2.0	2.0	U,C 	
Distilled water	4.8	4.8	4.8	4.8
Hexamethylene diamine	0.8	0.8		
nexamethyrene dramme	0.0	0.0		
			Part A	<u>Part A</u>
20% aqueous solution				
hexamethylene diamine			4.0	4.0
·				
Dégassed Part B to paste	consistency (v	acuum removal	of water to s	solids indicated)
	-			
Weight % solids, Part B	79.3%	77.7%	82.4%	80.5%
Storage Stability @ 32°F	94 days	94 days	72 days	108 days
Aging of Cured Films Tensile Strength, psi/ Elongation, %				
18 days @ 500°F	560/280	430/200	560/330	350/300
25 days @ 500°F	480/230	370/140	350/270	300/230
38 days @ 500°F	460/250	290/100	365/180	220/160
52 days @ 500°F	380/70	250/40	280/120	200/100
	<i>J J J J J J J J J J</i>		200/ 140	

In both cases, VT-X-3627 was higher in solids content and aged better than VT-X-3627B at $500^{\circ}F$.

25

1

Incorporation of the calcium carbonate into the latex, as in 2211-132 and 2211-133, gave higher solids content and better aging--probably due to better dispersion of the pigment. Storage stability was not noticeably improved by omission of the pigment; however, sealants prepared at lower solids content and lower viscosity without pigment have been found to be more stable than those noted above. The suitability of such low solids sealants for use is questionable because of excessive shrinkage and flow. All VT-X-3627 and VT-X-3627B sealants have shown progressive increase in viscosity even on storage at 32°F, so storage times shown above can only be considered approximations. The desirability of a Viton latex with improved stability as well as good low-temperature properties is evident.

Improved ease of manufacture of the latex sealants was found by decanting the settled latex and adding a degassing agent, Foammaster S. A vinyl stabilizer, Vanstay 8050, was added to one of the formulations shown below in an attempt to improve high-temperature resistance by decreasing the tendency to dehydro-fluorination. Aging tests were conducted at 550°F to accelerate the test.

	2211-174	<u>2211-177</u>	2211-179
VT-X-36278 VT-X-3627	127.0	124.0	124.0
M-Pyrol	15.0	15.0	15,0
FS-1265	12.0	12.0	1z 0
Witco 975	1.5	1.5	1.5
Foammaster S	1 drop	1 drop	1 drop
OMYA-BLH Distilled water	20.0 4.8	20.0 4.8	20.0 4.8
Tamol 731 dispersion	0.8	0.8	0.8
Witco 900	0.2	0.2	0.2
Vanstay 8050			2.0

Degassed to paste consistency (vacuum removal of water to solids indicated) Curing agent - 3 parts 20% aqueous solution hexamethylene diamine Weight % solids 83.7% 82.5% 82.8%

	2211-174	2211-177	2211-179
Tensile Strength, psi/ Elongation, %			
Initial – cured 1 day @ 158°F plus 4 hrs. @ 250°F, plus 20 hrs. @ 350°F	70/410	120/600	60/440
7 days @ 550°F 14 days @ 550°F	260/300 170/120	240/300 150/120	290/240 260/100
Weight Loss: 7 days @ 550°F 14 days @ 550°F	-28.3% -39.2%	-27.4% -37.8%	-29.3% -44.3%

No improvement in aging was found with Vanstay 8050. Stability in storage at 32°F was not as good as with formulations mixed earlier in the project. This was probably due to changes occurring in the latex on standing over a period of several months.

Corners were sealed with these sealants initially as well as one hour after mixing with the curing agent. No cracking occurred in any case. Application life is considered to be at least one hour.

Adhesion studies were conducted on titanium, MIL-T-9046, Type 3, Composition C-6 A1/4V. Preliminary tests included titanate, several silane-type primers and MIL-C-23377 epoxy-polyamide primer. These were tested with and without a secondary coating of brush-type solvent-based Viton Ming 2211-43 cured with hexamethylene diamine. Best results were obtained with a silane-type PRC Primer #6, or with MIL-C-23377 epoxy-polyamide primer with a secondary coating, 2211-43, prior to application of the latex sealant. Curing at 350°F was generally required to obtain adhesion. Exposure to 500°F oven-aging resulted in loss of adhesion. Tests were conducted at 400° and 450°F to determine how long adhesion was retained at these temperatures. Due to the limited quantity of latex sealant available, peel adhesion tests were not conducted. The sealant was cut to the metal and manually pulled to determine

whether adhesive or cohesive failure resulted. Cohesive failure was considered to be satisfactory adhesion. The following results were obtained:

	2211-177	2211-179
PRC Primer #6 + 2211-43		
9 days @ 400°F	cohes i ve	cohes i ve
12 days @ 400°F	cohes î ve	adhes i ve
14 days @ 400°F	cohes i ve	
2 days @ 450°F	cohes i ve	cohes i ve
5 days @ 450°F	adhes i ve	adhes i ve
MIL-C-23377 + 2211-43		
2 days @ 450°F	cohes i ve	cohes i ve
5 days @ 450°F	adhes i ve	adh e s i ve

The formula for 2211-43 is shown below:

	2211-43
ECD-487	100
Maglite Y	15
FS-1265	8
Methyl isobutyl ketone	80
MEK	300
	503

Curing agent - 20% hexamethylene diamine in MEK - 8.0 parts

Two additional primers, Chemlok 607 and a polyimide polymer (Araldite P-13N, diluted one to one with dimethyl formamide) were tested with 2211-177. The secondary primer, developed under the work on tape sealants, was 2211-186.

	2211-186
ECD-487	100
Maglite Y	15
Thermax	15
Durez 12686	3
FS-1265	8
Diak No. 1	2
MEK	288
	431

The following results were obtained:

Cured 24 hrs. @ 158°F +	<u>Chemiok 607</u>	<u>Araldite P-13N</u>
4 hrs. @ 250°F + 20 hrs. @ 350°F	cohes i ve	partly cohesive
l day @ 450°F 2 days @ 450°F 5 days @ 450°F	cohesive partly cohesive partly cohesive	cohes i ve cohes i ve adhes i ve
· 14 days @ 450°F	partly cohesive	adhesive

IV. PHASE IIA - IMPROVING THE LIFE OF FLUOROCARBON SEALANTS

A. Elimination of Metallic Impurities

It has been shown that metallic catalysts such as iron and zinc greatly destabilize vinyl resins, producing catastrophic dehalogenation at elevated temperature. Use of procedures which introduce a minimum of iron and incorporation of organic soluble metal chelating agents to remove such impurities is indicated.

B. Antioxidants

The unsaturation resulting from loss of halogen as well as any hydrogen originally present is subject to reaction with oxygen of the air. Phenolic antioxidants and organic and metallic phosphites have been used to reduce this attack.

C. Metallic Soaps

Combinations of calcium, cadmium and barium soaps have been found useful in adding to the stability of vinyl chloride polymers.

D. Acid Scavengers

Organic materials which remove the acids of decomposition can greatly extend the life of halogenated polymers since these acids are autocatalytic in their effect on decomposition rates. Materials such as olefins and epoxies are known to provide stability until chemically exhausted.

E. Free Radical Scavengers

Very effective, free radical scavengers are a series of sulfur-bearing compounds along with substituted phenolics. These materials prevent the peroxide formation and decomposition occurring at unsaturated sites.

F. Investigation of Cure Mechanisms on Stability

Mechanisms for curing "Viton" involve removal of halogen acid by various procedures. This halogen acid, frequently present as a dissociable salt at high temperatures (or the fluoride ion), can produce problems. The effect of different cures and degrees of cure on stability were investigated.

Since it is beyond the scope of this project to evaluate all of the available materials of these types, representative materials from each category were chosen:

Formula	<u>Material</u>	Туре
2211-145	Nopchelate OS	Chelating Agent
2211-140	Santovar A Mark X Dyphos	Antioxidant " "
2211-143 2211-144	Vanstay 4030 (barium-cadmium) Vanstay 8050 (calcium-zinc)	Metallic Soap
2211-141 2211-139	Epon 1001 Tris-beta chloroethyl phosphine	Acid Scavenger
2211-137 2211-138	Catalin CAO-6 Stabilite White	Free Radical Scavenger
2211-136		None

The basic formula, 2211-23, was used for comparison. The ketimine, which is very slow-curing, was replaced by 1.6 parts 85% solution of hexamethylene diamine. The sealants were thinned to a flowable viscosity with MEK, curing agent added, and flowed onto glass plates at a dry thickness of 15 mils. After curing, weight loss and tensile properties were determined after exposure at 550°F.

Formula	Additive	Days @ 550°F	Weight Loss	Tensile Strength, psi/ Elongation, %
2211-136	None	0 4	-22 . 7% ·	570/1050 610/380
		7	-37.3%	860/230
		10	-52.0%	1240/85
2211-137	Catalin CAO-6	0		500/1200
	2 phr	4	-26.4%	670/500
		7	-39.3%	900/200
		10	-52.0%	1100/30
2211-138	Stabilite White	0		600/1200
	2 phr	4	-23.1%	640/450
	- F	7	-34.4%	770/120
		10	-46.7%	1130/20
		ĨŬ	-40.//	1130/20
2211-139	Tris-beta-chloroethyl	0.		270/950
•	phosphine, 2 phr	4	-20.4%	640/450
		7	-32 .2%	770/120
		10	-46.3%	1130/20
2211-140	Santovar A	0		560/1070
	2 ph:	4	-24.1%	700/300
	- •	7	-37.1%	770/120
		10	-50.6%	1510/30
2211-141	Faca 1001	0		E20/920
2211-141	Epon 1001	0	a 1 1.07	530/820
•	2 phr	4	-31.4%	990/135
		7	-50.3%	1420/20
	•	10	-63.4%	brittle
2211-142	Dyphos, 8 phr	0		1120/700
		4	-46.8%	420/120
		7		brittle
2211-143	Vanstay 4030, 2 phr	e		800/870
		4	-26.0%	830/175
		7	-40.4%	1140/50
		10	-54.1%	brittle
2211-144	Vanstay 8050, 2 phr	0		550/820
2211-144	vanstay 0050, 2 phr	0	20.0%	
		4	-20.0%	670/375
		7	-27.3%	860/240
		10	-36.4%	1240/130
2211-145	Nopchelate OS	0		520/840
	l phr	4	-25.3%	600/460
		7	-41.7%	760/155
		10	-56.3%	1260/15
2211-146	Mark X, 2 phr	0		540/870
······································	······································	4	-21.3%	590/355
		7	-36.7%	780/120
•		τó	-48.8%	1510/20
		• -		· · · · · · · · · · · · · · · · · · ·

ø

In most cases, the additives used were ineffective or were deleterious to aging of the Viton sealant. One material, Vanstay 8050, an organic mixture of calcium and zinc compounds supplied by R. T. Vanderbilt Co., showed improved aging. Weight loss after ten days at 550°F was considerably lower than for any of the other materials tested and elongation was greater. Additional aging tests indicated that 2 phr (parts per 100 of rubber) and 1.6 phr hexamethylene diamine were optimum quantities. The low film thickness of 15 mils in this test was conducive to rapid aging. A thicker fillet of sealant would be expected to retain flexibility for a longer period at 550°F.

G. Pre-extruded Sealant

Some of the properties of Viton fluorocarbon lend themselves nicely to the use of pre-extruded forms. This is not true of the liquid polymers which are too weak and uncured to be self-supporting. Such preextruded forms are widely used in construction in cases where the movement and design do not cause working out of these necessarily thermoplastic materials. This limitation is not present with Viton, however, since it actually cures to a thermoset condition. Further, the slow curing time of the Viton ECD-487 is a distinct advantage since it may be supplied already catalyzed, in a cold condition, and will slowly vulcanize in place. The high strength and viscosity of the polymer now become an advantage and it is not necessary to provide a fluid, easy-working sealant. While application and tooling methods are different, the criteria that make a suitable extrusion are well-defined. Solids contents of 80% or more become relatively simple.

1. Formulation and Tape Criteria

Information gathered in Part A of Phase I can be directly transferred here, but higher solids materials are inherent in this approach since the viscosity limitations disappear and other criteria become significant.

a. Cone Penetrometer

In place of viscosity, the cone penetrometer requirements are used as a control. These values range in the case of construction sealants from 70 mm to 140 mm, the lower values (higher viscosity) being used for the sealants requiring some resistance to deformation. Since Viton cures, this upper limit is more for application ease than for providing deformation resistance (ASTM D-5, 100 gms - 5 seconds).

b. Extrusion of Samples

Each material and shape requires a design allowance in the die to produce the proper cross-section. A laboratory extruder with the simple slot and circular cross-section dies was used, with notice being taken of the relation of formula characteristics to extruded shape and size. A Brabender laboratory extruder, Model 101, equipped with variable speed roll feeder and oil-heated barrel with Thermotron #1001 Controller, was used. The extrusion process is shown in Plate 2.

2. Goal

An easy-to-use, self-sticking (to suitable primer or brush-type Viton) extrusion with all the fuel and heat resistance of the liquid sealant containing not more than 10 volume percent of volatiles.

Preliminary tape formulations were prepared as shown below:

	2211-181	2211-182	2211-183
ECD-487	100.C	100.0	100.0
Maglite Y	15.0		
Calcium oxide		15.0	15.0
Thermax	15.0	15.0	15.0
FS-1265	8.0	8.0	8.0
Vanstay 8050	2.0	2.0	2.0
HMDA	1.6		1.6
Diak No. 3		4.0	
MEK	8.0	8.0	8.0

	2211-181	2211-182	2211-183
Extrusion properties	poor	good	poor
NVM of extruded tape, %	96.2	96.5	
Penetrometer, 1/10 mm	27.0	31.0	

Formulations cured with HMDA (hexamethylene diamine) extruded poorly, apparently because of incipient cure. Use of a blocked amine such as Diak No. 3 is necessary for good extrusion. Diak No. 3 is N,N¹-dicinnamylidine - 1,6 hexane - diamine. Diak No. 1 (hexamethylene diamine carbamate) was next investigated. J

	2211-184	2211-185	2211-186	2211-188
ECD-487	100.0	100.0	100.0	100.0
Maglite Y		15.0	15.0	15.0
Calcium oxide	15.0			
Thermax	15.0	15.0	15.0	15.0
FS-1265	8.0	8.0	8.0	8.0
Vanstay 8050	2.0	2.0		2.0
Durez 12686			3.0	
Diak No. 1	2.5	2.5	2.0	2.0
MEK	8.0	8.0	8.0	8.0
Extrusion properties	good	good	poor	poor
NVM of extruded tape, %	. 96.0	95.7	96.7	96.5
Penetrometer, 1/10 mm	31.0	31.0	28.0	29.0

Formulations prepared with 2.5 parts Diak No. 1 extruded well, but with 2.0 parts, extrusion was poor. The plasticizing effect of the higher amine content is beneficial to extrusion. Either calcium oxide or Maglite Y can be used as acid acceptor with 2.5 parts Diak No. 1 to obtain satisfactory extrusion. Durez 12686 was added to 2211-186 in an effort to improve tackiness of the tape for adhesion to metal. A portion of 2211-186 was thinned to brush consistency for use as a secondary primer for adhesion to titanium.

The best results on oven-aging at 550°F were obtained with 2211-181 and 2211-186. However, both of these formulations extruded poorly. Results are shown in the following table. 236

Aging of Tape Formulations (Tensile Strength (psi)/Elongation(%))

	2211-181	2211-182	2211-184	2211-185	2211-186	2211-188
Cured:						
24 hrs. @ 158°F	220/1100	105/600	350/900	320/980	130/700	250/900
+4 hrs.@250°F	300/910	325/1100	570/540	720/620	250/700	470/650
+20 hrs.@350°F	560/880	630/860	690/500	730/500	360/730	610/590
Aged :						
7 days @ 550°F	700/300	430/230	600/110	650/210	640/320	550/260
14 days @ 550°F	1220/75	brittle	710/20	810/60	990/80	900/60
Weight Loss						
Aged :						
7 days @ 550°F	-25.0%	-32.2%	-30.4%	-26.1%	-27.3%	-36.5%
14 days @ 550°F	-43.1%	-53.1%	-49.9%	-44.2%	-44.2%	-52.2%

Experiments were conducted in adhering the tape to titanium. A double primer system was used, the second coat consisting of a brush-type Viton sealant. The tape was washed with MEK and rolled down into the primer when the latter was barely dry as shown in Plate 3. As the tape is inherently not very tacky, it must be applied while a little solvent still remains in tape and primer. The panels were dried four days at room temperature prior to exposure in the 158°F oven. The following results were obtained;

Tape First Primer <u>Second Primer</u>	2211-181 PRC Primer #6 2211-43	2211-181 MIL-C-23377 2211-43	2211-182 PRC Primer #6 2211-43	2211-182 MIL-C-23377 2211-43
24 hrs.@ 158°F				
+ 4 hrs.@ 250°F	adhes i ve	cohes i ve	adhes i ve	cohes i ve
+ 20 hrs.@ 350°F	adhes i ve	cohesive	adhes i ve	cohes i ve

Tape First Primer Second Primer	2211-181 PRC Primer #6 2211-43	2211-181 MIL-C-23377 2211-43	2211-182 PPC Primer #6 2211-43	2211-182 MIL-C-23377 2211-43
1 day @ 450°F	adhes ive	cohes i ve	cohes i ve	cohes i ve
6 days @ 450°F	adhes i ve	cohes i ve	cohes i ve	cohes ive
8 days @ 450°F	adhes ive	adhes i ve	cohes i ve	cohes i ve
14 days @ 450°F	adhes i ve	adhes i ve	cohes i ve	cohes i ve
19 days @ 450°F	adhes i ve	adhes i ve	adhes ive	adhes i ve

The best results were obtained with the Diak No. 3 cured formulations 2211-182. PRC Primer #6 and MIL-C-23377 epoxy-polyamide primer were equal in retention of adhesion at 450°F, but the latter had better initial adhesion. The following formulations were prepared and extruded:

	2211-190	2211-191	2211-192	2211-193
ECD-487	100	90	100.0	100.0
Viton LM		10		
Maglite Y	15			
Maglite D		15	15.0	15.0
Thermax	15	15	15.0	15.0
FS-1265	8	8	8.0	8.0
Vanstay 8050	2	2	2.0	
Diak No. 1		2		
Diak No. 3	. 4		3.2	3.2
Durez 12686	**			3.0
MEK	8	8	12.0	8.0
Extrusion properties	good	poor	fair	poor
N/M of extruded tapes, %	96.0%	96.2	94.0	96.17
Penetrometer, 1/10 mm	32	32	35	31

The plasticizing effect of the higher amine content is evident in the above results. The formula with four parts Diak No. 3, 2211-190, had the best extrusion properties. Additional MEK (in 2211-192) was not as effective as the higher Diak No. 3 content. The use of low molecular weight Viton LM (in 2211-191) did not improve extrusion. Aging test results are shown on the following page.

٩.

Ŧ

	Tensile Strength(psi)/Elongation(%)				
	2211-190	2211-191	2211-192	2211-193	
Cured 24 hrs. @ 158°F + 4 hrs. @ 250°F	180/1200	680/400	380/1100	240/1050	
Cured 24 hrs. @ 158°F + 20 hrs. @ 350°F	670/8 20	920/300	640/920	900/700	
Aged 7 days @ 550°F	660/300	870/30	500/230	540/250	
Aged 14 days @ 550°F	1300/20	Brittle	1000/50	1120/30	
		Weight	Loss		
Aged 7 days @ 550°F	-28.9%	-39.6%	-22.7%	-24.1%	
Aged 14 days @ 550°F	-47.5%	-62.5%	-41.6%	-42.6%	

The formulation containing ten parts Viton LM, 2211-191, exhibited poor aging. Best aging results were obtained with formulations containing the lower Diak No. 3 content.

Adhesion studies were conducted using a pretreatment of the titanium surface with Pasa-Jell 107-prior to priming.

	PRC Pri 2211-186		Chemlok 2211-186		Araldit diluted 5 2211-186	0% w/DMF
	2211-190	2211-192	2211-190	2211-192	2211-190	2211-192
Cured 24 hrs.@ 158°F + 4 hrs.@ 250°F	cohes i ve	adhes i ve	cohes i ve	cohes i ve	adhes i ve	adhes i ve
Cured 24 hrs.@ 158°F + 20 hrs.@ 350°F	cohes i ve	cohes ive	cohes i ve	cohes i ve	adhes i ve	adhes i ve
1 day @ 450°F	cohes i ve	cohes i ve	cohes i ve	cohes i ve	cohes i ve	cohes i ve
2 days @ 450°F	adhes i ve	adh es i ve	adh as ive	cohes i ve	cohes i ve	cohes i ve
14 days @ 450°F	* *		~ ~	cohesive	cohes i ve	cohes i ve
21 days @ 450°F			**	adhes i ve	cohesive	cohes i ve
28 days @ 450°F					cohes i ve	cohes i ve
35 da ys @ 450°F					cohes i ve	cohes i ve
63 days @ 450°F		239		40	cohes i ve	cohes i ve

The polyimide resin primer, Araldite P-13N, was superior in retention of adhesion at 450° F. However, it requires a higher temperature to secure initial adhesion, and good initial adhesion was not obtained with this primer system until the panels were exposed to 450° F.

3

ų

ž

A small quantity of LD-48/-LV, a lower Mooney version of ECD-487, was obtained. This was evaluated in comparison with ECD-487 in a formulation using 3.6 parts Diak No. 3. This represents a compromise between the superior aging with 3.2 parts and the better extrusion properties with 4.0 parts Diak No. 3. In addition, formulations were prepared from each polymer with FS-1265 fluorosilicone oil omitted.

	2211-197	2211-198	2211-199	2211-200
ECD-487	100.0			100.0
LD-487-LV Maglite Y	 15.0	100.0 15.0	100.0 15.0	15.0
Thermax FS-1265	15.0 8.0	15.0 8.0	15.0 	15.0
Vanstay 8050 Diak No. 3	2.0 3.6	2.0 3.6	2.0 3.6	2.0 3.6
MEK	10.0	8.0	10.0	12.0
Extrusion properties	good	good	fair	fair
NVM of extruded tape	95.0%	95.9%	94.8%	93.9%
Penetrometer, 1/10 mm	29	30	30	34

Tensile Strength(psi)/Elongation(%)

Cured 24 hrs, @ 158°F + 4 hrs, @ 250°F	380/1100	340/1000	460/1100	420/1200	
Cured 24 hrs.@ 158°F + 20 hrs.@ 350°F	710/860	600/840	680/1050	650/1100	
Aged 7 days @ 550°F	550/3 70	690/270	680/230	430/250	
Aged 14 days @ 550°F	1090/40	1600/20	°50/15	620/35	

	Weight Loss					
•	2211-197	2211-198	2211-199	2211-200		
Aged 7 days @ 550°F	-20.6%	- 25.5%	-24.6%	-22.8%		
Aged 14 days @ 550°F	-40.8%	-43.4%	-45.4%	-42.2%		

2-6-6-8-4

Solids content of the LD-487-LV tape sealant was only slightly higher than that prepared from ECD-487, and there was no noticeable difference in application properties. The heat resistance properties of the LD-487-LV were not as good as those of ECD-487, as noted by the higher weight loss and lower elongation of the former. The omission of fluorosilicone oil was harmful to extrusion properties and made no appreciable difference in heat resistance.

Adhesion studies were conducted with and without the Pasa-Jell 107 treatment of titanium, using Araldite P-13N diluted 50% with DMF and comparing secondary primers with and without FS-1265 fluorosilicone oil. A portion of 2211-200 was thinned to brush consistency for use as a secondary primer. Test results are shown below:

> Araidite P-13N 2211-186 Primer No Pasa-Jeli 107

	<u>2211-197</u>	2211-198	2211-199	2211-200		
Cured 24 hrs.@ 158°F + 4 hrs.@ 250°F + 20 hrs. @ 350°F	adhes ive	adhes i ve	adhes i ve	adhesive		
l day @ 450°F 6 days @ 450°F 8 days @ 450°F	cohes ive cohes ive adhes ive	cohes i ve cohes i ve adhas i ve	cohes i ve adhes i ve 	cohes ive adhes ive		
	Araldite P-13N 2211-200 Primer No Pasa-Jell 107					
	2211-197	2211-198	2211-199	2211-200		
Cured 24 hrs. @ 158°F + 4 hrs. @ 250°F + 20 hrs. @ 350°F	adhes i ve	adhes i ve	adhes i ve	adhes i ve		

ī,

	2211-197	2211-198	2211-199	2211-200
l day @ 450°F 2 days @ 450°F 6 days @ 450°F 8 days @ 450°F 10 days @ 450°F 30 days @ 450°F	cohesive cohesive cohesive cohesive 40% con. 40% coh.	cohes ive cohes ive cohes ive adhes ive	adhes ive 50% coh. adhes ive 	cohes ive cohes ive adhes ive
	Araldite P-13N 2211-186 Primer Pasa-Jell 107			
	2211-197	2211-198	<u>2211-199</u>	2211-200
Cured 24 hrs. @ 158°F + 4 hrs. @ 250°F + 20 hrs.				
@ 350°F	adhes i ve	adķes i ve	adhes i ve	adhes i ve
1 day @ 450°F	cohes i ve	cohesive	cohes i ve	cohesive
6 days @ 450°F	cohes i ve	cohesive	cohes i ve	cohes i ve
8 days @ 450°F	cohes ive	cohes ive	cohes i ve	adhes i ve
10 days @ 450°F	cohes i ve	cohes i ve	30% coh.	
21 days @ 450°F	cohes ive	cohesive	30% coh.	
23 days @ 450°F	cohes ive	cohes i ve	adhes i ve	
30 days @ 450°F	cohes i ve	cohes i ve		
	Araldite P-13N 2211-200 Primer Pasa-Jell 107			
	2211-197	2211-198	2211-199	2211-200
Cured 24 hrs. @ 158°F + 4 hrs_ @ 250°F + 20 hrs.				11
@ 350°F	adhes i ve	adhes i ve	adhes i ve	adhes i ve
1 day @ 450°F	cohes i ve	cohes ive	cohes ive	cohesive
6 days @ 450°F	cohes ive	cohes i ve	adhes i ve	adhes i ve
8 days 🐼 450°F	cohes i ve	cohes i ve	~ -	
10 days @ 450°F	cohes i ve	30% coh.		
14 days @ 450°F	cohes ive	adhes i ve	~ *	-*
17 days @ 450°F	50% coh.		44 4 9	:
23 days @ 450°F	50% coh.			
27 days @ 450°F	adhes i ve			

Test results above confirm the improvement in adhesion with Pasa-Jell 107 treatment and 'Araldite P-13N plus 2211-186 Primer. This secondary primer, containing FS-1265 Fluorosilicone 0il, was superior to 2211-200 Primer which contained no FS-1265. ų

The tape sealants, 2211-197 and 2211-198, which contained FS-1265, also retained adhesion for a longer period at 450°F than 2211-199 and 2211-200, which contained no FS-1265.

V. RAW MATERIALS

وتوافقتها وأحمدهم كماناه مشاولته فالأقيم ومعاومتهم ومجروعه ومالوال والمقاولات والمرافقا تسويهم والمراجع

ģ

ſ

چ ۲

f

è

Viton ECD-487	E. I. DuPont de Nemours & Co.
Viton LD-487-LV	11
Viton LM	u
Viton LD-011	i i
Viton VT-X-3518	88
Viton VT-X-3627 Latex	· u
Viton VT-X-3627B Latex	11
Viton L-31 Latex	88
Diak No. 1	11
Diak No. 3	
Maglite D	Merck & Co.
Maglite Y .	Merck & Co.
Thermax .	R. T. Vanderbilt Co.
FS-1265, fluorosilicone oil	Dow Corning
EDA-DIBK Ketimine (ethylene diamine diisobutyl ketone)	Synthesized at Products Research & Chemical Corporation
MEK (methyl ethyl ketone)	Shell Chemical Co.
MIK (methyl isobutyl ketone)	11 12 11
Calcium hydroxide	Baker Chemical Co.
Calcium oxide	38 88 88
Hydroquinone	Van Waters & Rogers
N,N-dimethyl dodecylamine	Union Carbide
Shell H-3	Shell Chemical Co.
Dimethyl acetamide	Van Waters & Rogers

Polycyanos i loxane

OMYA-BLH

Witco 900

Witco 975

Tamo1 731

Tabular alumina, 325 mesh

M-Pyrol (N-methyl, 2-pyrolidone)

Foammaster S

Vanstay 4030

Vanstay 8050

Durez 12686

Nopchelate OS

Santovar A

Mark X

Dyphos

Epon 1001

Tris-betachloroethyl phosphine

Catalin CAO-6

Stabilite White

HMDA (hexamethylene diamine)

Products Research & Chemical Corporation

٦

z

٩.

Pleuss-Stauffer

Witco Chemical

Rohm and Haas

Alcoa

GAF Corp.

Nopco Chemical Division, Diamond Shamrock

R. T. Vanderbilt Co.

Hooker Chemical Corp.

Nopco Chemical Division, Diamond Shamrock

Monsanto Chemical Co.

Argus Chemical Co.

National Lead Co.

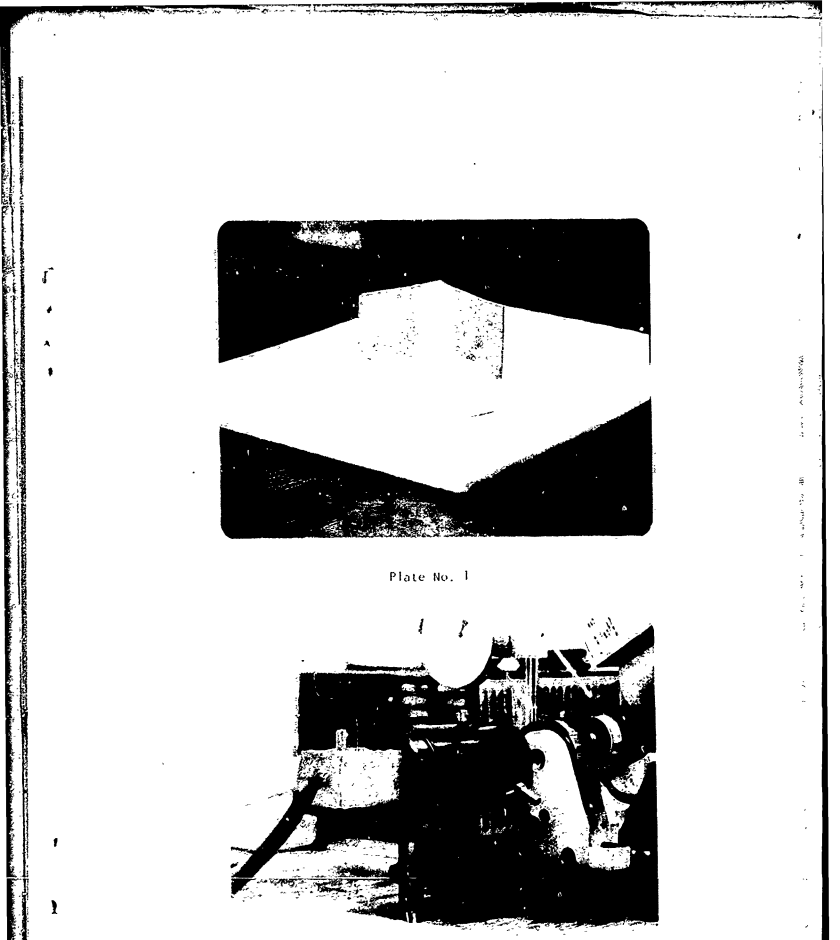
Shell Chemical Co.

Monsanto Chemical Co.

Catalin Corp. of America

R. T. Vanderbilt Co.

E. I. DuPont de Nemours & Co.





~~~~

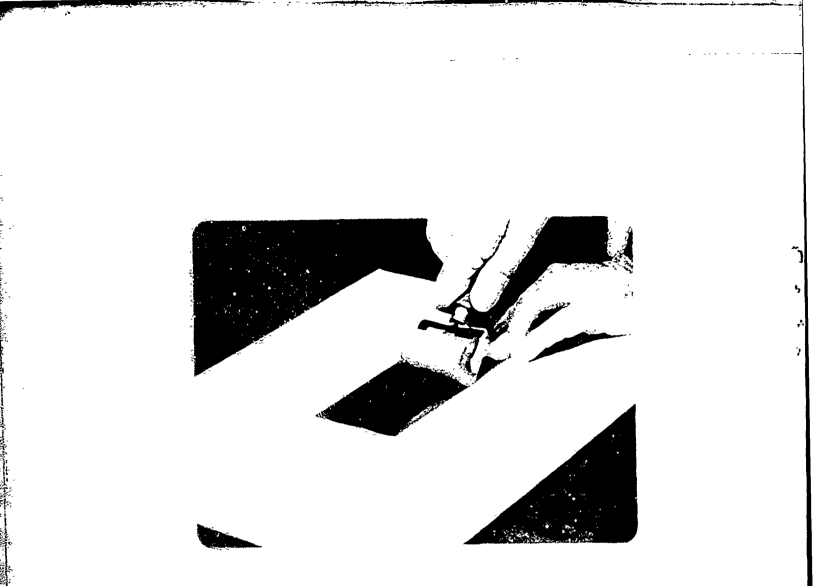


Plate No. 3