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Test Program to Demonstrate the Stability of Hydrazine in Propellant Tanks

Final Report

Clifford M. Moran Roy A. Bjorklund

April 1983

Prepared for United Kingdom Treasury and Supply Delegation Washington, D C

through an agreement with

National Aeronautics and Space Administration

by

Jet Propulsion Laboratory California Institute of Technology Pasadena, California



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Reference to any specific commercial product, process, or service by trade name or manufacturer does not necessarily constitute an endorsement by the United States Government or the Jet Propulsion Laboratory, California Institute of Technology.

CONTENTS

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	Acknowledgements						
	Abstract						
	Acronyms						
T							
1.							
	A. COUPON TEST PROGRAM						
	B. BACKGROUND-RELATED TECHNICAL WORK						
	C. OBJECTIVES OF THE COUPON TEST PROGRAM						
	D. MATERIAL COUPON SOURCE						
11.	TEST PROCEDURES						
	A. TEST UNIT PREPARATION						
	B. STORAGE TESTING						
	C. POSTTEST ANALYSIS						
	l. Discussion						
	2. Procedure - Complete Analysis						
	3. Procedure - Limited Analysis						
	4. Gases of Decomposition						
	5. Residual Hydrazine						
гтт							
LLLe							
	A. DESCRIPTION OF TEST SPECIMENS						
	B. DETAILS AND SUMMARIES OF POSTTEST ANALYSES AND RESULTS						
	C. PROPELLANT CONTROLS						
	D. SURFACE ANALYSIS						
	1. Introduction						
	2. XPS Techniques						
	3. Results						
	4. Discussion						
	5. Scanning Electron Microscopy (SEM) Examination						
	6. Conclusions						
EV.	CONCLUSIONS						
	A. PROGRAM A: 6 MONTHS STORAGE						
	B. PROGRAM B: 24 MONTHS STORAGE						
Ĩ.	DATA TABLES						
/I.	REFERENCES						
ALLEI	NDIGES						
	A. PRETEST HYDRAZINE ANALYSIS						
	B. SPECIMEN LOGS						
	C. ANALYSIS PROCEDURE FOR CO ₂						
	D. PHOTOGRAPHS OF COUPONS						
	E. EXAMINATION OF 347 CRES WELD SPECIMENS						

Figures								
1-1.	Hydrazine Actuation System (HAS) Propellant Tank 1							
1-2.	Coupon Location on HAS Tank 1							
2-1.	Procedures for Capsule Filling 2							
2-2.	Typical Glass Capsule Test Unit 2.							
2-3.	"Lazy-Susan"-Type Storage Facility 2-							
2-4.	Procedure for Posttest Chemical Analysis 2-							
2-5.	Specimen/Capsule Test Opening Fixture 2							
3-1.	JPL XPS Laboratory 3-4							
C-1	Test Procedures for CO ₂ Analysis C-2							
D-1	Test Specimens, Hydrazine Decomposition Program "A" -							
	Secondary Containment System D-2							
D-2	Test Specimens, Hydrazine Decomposition Program "B" -							
	Primary Containment System D-7							
E-1	CRES 347 Weld Specimens, Surface Features at Weld E-S							
Е-2	CRES 347 Weld Specimens, Surface Features at Heat-							
	Affected Zone E-6							
Е-З	CRES 347 Weld Specimens, Cross Sections at Weld E-7							
Е-4	CRES 347 Weld Specimens, Cross Sections at Heat-							
	Affected Zone E-8							

Tables

1.	Listing of Coupon Test Numbers and Description	5-2
2.	Summary of Analyses and Results	5-3
3.	Details of Analyses and Results	5-6
4.	Summary of Analysis of Hydrazine Controls	5-9

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ABSTRACT

This document reports the analyses and results of a 24-month coupon test program to evaluate the decomposition of hydrazine by metallic components of propellant tanks. The propellant fuel evaluated was monopropellant-grade hydrazine (N_2H_4) , which is normally a colorless, fuming, corrosive, strongly reducing liquid. The degree of hydrazine decomposition was determined by means of chemical analyses of the liquid and evolved gases at the end of the test program. The experimental rates of hydrazine decomposition were determined to be within acceptable limits.

The propellant tank materials and material combinations were not degraded by a 2-year exposure to hydrazine propellant. This was verified using changeof-weight determinations and microscopic examination of the specimen surfaces before and after exposure, and by posttest chemical analyses of hydrazine liquid for residual metal content.

ACRONYMS

ACS	attitude control system
ARDE	ARDE, Inc., Mahawah, NJ
BAT	Bell Aerospace Division of Textron, Inc., Buffalo, NY
BE	binding energy
Caltech	California Institute of Technology
CRES	corrosion-resistant steel
CPR	coupon preparation requirement
EB	electron beam
EPR	ethylene propylene rubber
ETS	Edwards Test Station, JPL
FEP	fluorinated ethylene propylene (Teflon)
HAS	hydrazine actuation system
HAZ	heat-affected zone
JPL	Jet Propulsion Laboratory
NASA	National Aeronautics and Space Administration
OAST	NASA Office of Aeronautics and Space Technology
OSS	NASA Office of Space Sciences
PES	photoelectron spectra
SEM	scanning electron microscope
STP	standard temperature and pressure
TBD	to be defined/determined/done
TIG	tungsten inert gas
UKTSD	United Kingdom Treasury and Supply Delegation, Washington, D.C.
XPS	X-ray photoelectron spectroscopy

SECTION I

INTRODUCTION

A. COUPON TEST PROGRAM

The Coupon Test Program has been an investigation of the reactive compatibility of hydrazine with various metallic components of a propellant storage tank. The hydrazine/material compatibility research reported here was performed by the Jet Propulsion Laboratory (JPL), California Institute of Technology (Caltech), under Contract NAS7-198 with the National Aeronautics and Space Administration (NASA) for the United Kingdom Treasury and Supply Delegation (UKTSD) in accordance with the UKTSD Letter Agreement F-2479, dated July 5, 1979.

This coupon test program is an extension of the ongoing JPL/NASA long-term propellant/material compatibility program. The same procedures, test methods, and test facilities developed under the JPL/NASA program have been applied to this program.

This document is the final report for the Coupon Test Program. An interim report from Program A was prepared in October 1981 (Reference 1).

B. BACKGROUND-RELATED TECHNICAL WORK

JPL has collaborated with other agencies on a variety of research, development, test, and evaluation projects. The laboratory, with its Pasadena facility and Edwards Test Station (ETS) at Edwards Air Force Base, California, maintains an institutional capability and technical expertise in evaluating and testing Earth- and space-storable liquid propellants and materials for spacecraft propulsion system applications. Specifically, JPL has been investigating material compatibility involving Earth-storable propellants, including hydrazine, since 1962 under sponsorship of the NASA Offices of Aeronautics and Space Technology (OAST) and of Space Sciences (OSS). The details of the JPL material compatibility program and interim experimental results of the longterm storage testing are reported in References 2 and 3. The long-term exposure testing continues, and the accumulated time for some test specimens exceeds 12 years.

The results obtained have provided reliable data for designing and qualifying chemical propulsion systems and components for long-life spacecraft. The work performed has directly supported the early JPL planetary flight projects such as Ranger, Surveyor, and Mariner, and the Viking 1975 and Voyager 1977 (Jupiter-Saturn-Uranus).

The general technology areas involved are propellant chemistry, metallurgy, long-term (10-year) propellant/material compatibility, metal fracture/ toughness characteristics, and fracture mechanics design of pressurized systems. Typical Earth-storable propellants are hydrazine, refined-grade hydrazine (monopropellant grade), hydrazine-hydrazine nitrate, monomethylhydrazine, and nitrogen tetroxide. Spacecraft propulsion system materials include aluminum alloys, corrosion-resistant steels (CRES), titanium alloys, and elastometric materials, for example, AF-E-332.

C. OBJECTIVES OF THE COUPON TEST PROGRAM

The overall objective of the coupon test program was to verify the longterm compatibility of hydrazine actuation system (HAS) propellant tank materials and other material combinations with monopropellant-grade hydrazine. To accomplish this overall objective, the program was divided into two parts.

Program A was intended to evaluate short-term compatibility of the secondary propellant containment system shown in Figure 1-1. It should be noted that the secondary containment system will be exposed to hydrazine only if there is leakage from the primary containment system. The program objectives were:

- Determine rates of hydrazine decomposition at 43°C by means of pressure rise monitoring throughout the term of the test program.
- (2) Verify that pressure containment materials and material combinations are not degraded by 6-month exposure to hydrazine propellant, using weight determinations and microscopic examination of specimen surfaces, after exposure.

Program B was intended to evaluate long-term compatibility of the primary propellant containment system shown in Figure 1-1. The program objectives were:

- Determine rates of hydrazine decomposition at 43°C and 60°C by monitoring pressure rise throughout the term of the test program.
- (2) Determine degree of hydrazine decomposition by means of chemical analysis of liquid and evolved gases at the end of the test program.
- (3) Verify that primary containment materials and material combinations were not degraded by 2-year exposure to hydrazine propellant, using weight determinations and microscopic examination of specimen surfaces, after exposure, and also by posttest chemical analysis of hydrazine liquid for metal content.

D. MATERIAL COUPON SOURCE

The material coupons used in this program were provided by the UKTSD. They were obtained from sections cut out of a HAS tank fabricated by Bell Aerospace Division of Textron (BAT). A total of 82 coupons from 26 different locations are listed in Table 1.¹ The locations on the tank from which the coupons were cut are shown in Figure 1-2. Each coupon was processed, weighed, cleaned, and individually sealed in a plastic bag by BAT before delivery to JPL. The coupons remained sealed until they were removed and placed immediately into glass capsule test units prepared at JPL.

¹All tables are contained in Section V.







Figure 1-2. Coupon Location on HAS Tank

SECTION II

TEST PROCEDURES

A. TEST UNIT PREPARATION

Eighty-two hermetically sealed, glass-encapsulated test units were prepared by JPL with the materials specified and in accordance with the priorities established by the sponsor. All units were prepared in accordance with the procedures shown in Figure 2-1.

The test requirements for this program were specified in JPL Proposal 90-965, Revision 2, October 11, 1978 and are summarized below.

- Purified hydrazine (VL-75 grade) was supplied by JPL. Pretest analysis of this propellant is shown in Appendix A. This propellant met the BAT material specification (Reference 4).
- (2) All test coupons were supplied by UKTSD, together with appropriate documentation prepared by BAT. For Program A, 38 coupons were designated; for Program B, 44 coupons were designated. The distribution of coupons by test storage temperatures was also designated.
- (3) The test containers were Pyrex capsules, as described in Reference 2, and designed to have an internal volume, when sealed, of about 80 cm³. Figure 2-2 is a photograph of a typical test unit.

Test equipment, instrumentation, and techniques duplicated those employed in the JPL 10-year test program and reported in Reference 2. The pretest procedures are summarized below.

- (1) Strain gauges used to measure internal pressure buildup were mounted on the open capsules. A preliminary pressure calibration was used to check the sensitivities of the strain gauges.
- (2) Test specimens were installed into the clean capsules in "as received" condition from BAT except for the EPR-515 O-rings which were coated with Krytox lubricant by JPL.
- (3) Funnel necks were fused onto the capsules, with care taken not to overheat the strain gauges. Final pressure calibration of the gauges was then made.
- (4) Internal volumes of the test capsules were measured by the expanding volume technique using high-purity gaseous nitrogen at ambient temperature.
- (5) Capsules were then loaded with enough hydrazine so that the combined volume of propellant and specimen was 40 ±0.5 cm³, and the specimen was fully immersed. Three of the specimens were found to be oversized and additional quantities of hydrazine were added.



Figure 2-1. Procedures for Capsule Filling



Figure 2-2. Typical Glass Capsule Test Unit

- (6) The hydrazine was frozen in a dry-ice/alcohol slurry and the capsules were pumped down to a pressure of less than 1 mm Hg.
- (7) The capsule necks were sealed off by fusing the glass tube, and the completed test units were made ready for shipment to Edwards Test Station (ETS).

The hydrazine was pretest-analyzed for purity in accordance with the JPL standard methods (Reference 2). A special CO_2 analysis of hydrazine in one test unit, representative of a typical encapsulation, was performed to verify the exclusion of air during filling. (See Appendix A). No examination or analyses were performed on the coupons.

B. STORAGE TESTING

The experimental storage phase was conducted at ETS to determine the compatibility of the hydrazine propellant with the materials. A photograph of the "Lazy-Susan" type of storage facility is shown in Figure 2-3. The exposure tests on the 82 units were conducted for a period of up to 24 months (730 days) at temperatures of either $43 + 3^{\circ}$ C ($110 + 5^{\circ}$ F) or $60 + 1^{\circ}$ C ($140 + 2^{\circ}$ F). The temperature of 43° C was used in the JPL long-term program (Reference 2) as the "normal" temperature of a noncryogenic propellant in space. The higher temperature was chosen as being the highest temperature likely to be experienced by the propellant in service. During the 24-month exposure term, the following was accomplished:

- (1) Pressure readings were taken once per week for the first month.
- (2) Pressure readings were taken once per month for the next five months.
- (3) Pressure readings were taken bimonthly for the remaining eighteen months.
- (4) Test units were visually inspected after taking the pressure readings.
- (5) All results were recorded for the above observations.

Details of all test units are presented in specimen logs in Appendix B.

C. POSTTEST ANALYSIS

1. Discussion

At the completion of the storage tests, all test units were analyzed in accordance with the JPL standard methods (from Reference 2). The capsules were opened and the decomposition gases, hydrazine, and coupons removed. The test coupons were weighed and their surfaces were visually examined at 50x magnification. The decomposition gases and hydrazine were analysed using posttest procedures developed and used in JPL's original program. The procedure



Figure 2-3. "Lazy-Susan"-Type Storage Facility

(Figure 2-4) provides for an analysis of all components. The choice of the posttest analyses to be applied is dependent on the test-unit history and the level of information required to permit satisfactory assessment of results. Details of the posttest analysis procedure are discussed in the following subsections.

2. Procedure - Complete Analysis

The hydrazine was frozen by immersion in liquid nitrogen (LN₂), and the capsule placed in the opening fixture (Figure 2-5). The capsule tip was broken and the volume of noncondensable gases (N₂ and H₂) measured in a calibrated vacuum system. The hydrazine was thawed and refrozen at -30° C (-22°F) and the gas at that temperature, mainly NH₃, measured. The residual hydrazine was removed using a syringe and the purity determined by a gas-chromatographic technique which measures NH₃ and H₂O. Metal content in the residual hydrazine was analyzed by atomic absorption techniques. A turbidimetric method was used for low concentrations of chloride; higher concentrations were titrated. Fluoride was determined colorimetrically.

3. Procedure - Limited Analysis

This procedure measures only the noncondensable gases, hydrogen and nitrogen. After thawing, the residual hydrazine was removed from the capsule using a syringe and was analyzed by gas chromatography for NH_3 and H_2O . If the NH_3 content is low, there may be an error due to NH_3 evolution before analysis.

4. Gases of Decomposition

a. Composition. The contents of the posttest capsule were frozen in liquid nitrogen and then prepared for sampling as follows: The strain gauge was very carefully scraped off with a sharp razor blade, a small scratch was made on the neck of the capsule, and the capsule was then enclosed in the opening fixture (Figure 2-5). The system was pumped down for several hours until moisture was removed from the outside of the capsule. The fixture was then filled with dry helium to 0.5 atmosphere to aid in heat transfer, and then immersed into liquid nitrogen to a depth equal to one-half the length of the capsule. After an hour, the helium was pumped out. When a satisfactory vacuum had been attained (1.3 x 10^{-2} N/m²), the gas sampling system was isolated from the vacuum pump, and the neck of the capsule was broken by turning the handle on the fixture. By means of a Toepler pump, the released noncondensable gases were pumped off through a liquid-nitrogen trap. The volume of the collected gases was measured manometrically, and a sample was taken for mass spectrometric analysis.

The nitrogen-to-hydrogen ratio of noncondensable gas was determined in most of the test units containing more than 5 to 10 cc at standard temperature and pressure (STP). For most of these analyses, the hydrogen content was undetectable. With few exceptions, the hydrogen content of the remaining test units was no more than 4%, and these exceptions were welded or brazed specimens. An unexplainable exception was test unit 4019, which contained a Lee plug, and for which the hydrogen content of the noncondensable gas was 12.5%. The total amount of gas was also high, 4 to 5 times the quantity found with the other three Lee-plug test units.



I la internet

Figure 2-4. Procedure for Posttest Chemical Analysis



Figure 2-5. Specimen/Capsule Test Opening Fixture

The liquid nitrogen traps were replaced by traps at -30° C (-22° F). After repeated thawing and refreezing of the hydrazine, the remaining condensable gases (mainly NH₃) were pumped off, measured, and sampled. The fixture was opened and the hydrazine removed using a syringe.

b. <u>Calculated Final Capsule Pressure</u>. The mean volume of the test capsules was 82 cm³. With 40 g of hydrazine and a standard metal coupon, the ullage was about 40 cm³. The pressure calculations depend upon the volume of nitrogen plus hydrogen (assuming negligible solubility in the hydrazine) and the volume of ammonia in the vapor phase.

The contributions of nitrogen and hydrogen were calculated from the perfect gas law:

 $P_1 = \frac{NRT}{V}$

where

The calculation of pressure due to the ammonia is not so simple and straightforward. Ammonia is highly soluble in hydrazine, and may not necessarily be an ideal gas at the temperatures and pressures considered.

Fortunately, solubility data for ammonia in hydrazine are available (Reference 5). Although the data do not cover the temperatures of interest, viz., 43°C and 60°C, it was possible to extrapolate the data of the above-referenced report. It can be shown that for the ammonia dissolved in hydrazine, the following relationship can be used to determine a close approximation of the ammonia pressure:

$$P \cong \frac{N/m}{K}$$
(2)

where

P = partial pressure of ammonia, atm N = moles of ammonia in system m = moles of hydrazine in system K = equilibrium constant (0.0455 atm⁻¹at 43°C, 110°F), (0.0295 atm⁻¹ at 60°, 140°F) (1)

c. <u>Pressure Rise Rate</u>. It would be of interest, both practically and theoretically, to have curves expressing pressure as a function of time for each test capsule. Unfortunately, due to aging of the bonding material, some of the strain-gauge data have proven to be unreliable.

d. <u>Percentage of Hydrazine Decomposed</u>. The percentage of hydrazine decomposed is calculated from the total weight of the gaseous products of decomposition - viz, nitrogen and ammonia. Some hydrogen may arise from the attack of metals by acidic constituents, but the error, if any, is insignificant because of the low molecular weight of hydrogen.

5. Residual Hydrazine

The residual hydrazine was removed from the capsule and analyzed as follows:

a. Impurities: NH_3 and H_2O . The NH₃ and H_2O contents of the hydrazine were analyzed by gas chromatography using a 0.0065-m-diam x 2-m-long (1/4-in.-diam x 6-ft-long) column filled with powdered Teflon coated with 15 wt% tri-ethanolamine. The inlet and column temperatures were held at $90^{\circ}C$ (194°F) and the helium flow set at 100 cm³/min. This column separates NH₃, H₂O, and N₂H₄, in that order.

b. Contaminants: Metals, Halogen and CO₂.

(1) <u>Metals</u>. The capsule was cut open and the metal sample rinsed with water, while any adhering material was rubbed loose with a rubber policeman. Any residue in the capsule was also rinsed out. All washings and residue were acidified with 5% HNO3, diluted to a known volume with water and analyzed for the appropriate metals by atomic absorption.

(2) <u>Halogens</u>. An aliquot of the acidified washings was checked for chloride by turbidimetry. If the chloride content was high, titration was used. The fluoride ion content was determined by a spectrophotometric method based on the bleaching of a zirconium alizarin color complex by fluoride ion. The absorbance was measured at 525 nm.

(3) <u>Carbon dioxide</u>. Hydrazine reacts with carbon dioxide to form the salt, hydrazinium carbazate. The equation for this reaction is

 $CO_2 + 2N_2H_4 \rightarrow (N_2H_5) (NH_2NHCOO).$

The method of analysis involved the addition of a sample of hydrazine to an excess of sulfamic acid. The sulfamic acid liberates CO_2 from the hydrazinium carbazate. Sulfamic acid was selected for use in the analysis because hydrazinium sulfamate is soluble in water and sulfamic acid is nonvolatile.

The liberated CO_2 was swept out of solution with helium gas, through a trap containing concentrated sulfuric acid to remove the water, and then through a special trap containing small glass beads where the CO_2 present was frozen out at the temperature of liquid nitrogen.

The trap containing the frozen CO_2 was provided with a special four-way stopcock that permits the CO_2 to be isolated in its loop. This trap is attached to a special sample introduction system on a custom-built chromatograph that permits the collected CO_2 to be quantitatively transferred through a chromatographic column for separation and assay. (Refer to Appendix C for details of this method.)

SECTION III

TEST RESULTS

A. DESCRIPTION OF TEST SPECIMENS

To determine the hydrazine/tank material compatibility, tests were performed on 26 types of metallic and nonmetallic coupons obtained from a fabricated tank, as shown in Figure 1-2. A detailed description of these coupons by CPR numbers and the results of the posttest visual examination are discussed in this section and are compiled in Table 2. Photographs of all test coupons in the posttest condition are presented in Appendix D. Additional data on 347 CRES weld specimens from the JPL/NASA long-term program are presented in Appendix E. A summary of posttest visual examination of coupons is presented below.

a. <u>CPR 1</u>. Specimens 001 to 008 are Type-304L corrosion-resistant steel (CRES) coupons from the ARDE, Inc., diaphragm. All specimens were observed to have a matte finish with no visible corrosion on their surfaces.

b. <u>CPR 2</u>. Specimens 014 to 017 are Type-304L CRES coupons from the ACS tank liner/diaphragm assemblies, which were girth-welded. All four specimens were shiny in appearance with no surface corrosion apparent. Specimen 014 had a very thin film along the heat-affected zone (HAZ) of the liner. Specimens 015 and 016 also had thin films along the HAZ of the diaphragm.

c. <u>CPR 3</u>. Specimens 023 to 026 are coupons from burst-disc assemblies made of Type-347 CRES preformed sheet stock, which was electron-beam (EB) welded to a ring made of Type-304L CRES. Each specimen was shiny and bright with no evidence of corrosion on its surface. Each diaphragm had identification numbers and symbols inscribed on it, in addition to the preformed scoring marks. All specimens were observed to have very small parallel scratches on the surfaces.

d. <u>CPR 4</u>. Specimens 034 to 037 are commercial Lee plugs made of Type-304L CRES. All four specimens had shiny and bright surfaces with no corrosion evident.

e. <u>CPR 5.</u> Specimens 045 to 048 are coupon sections from the Type-304L CRES liner/diaphragm liquid outlet housing, which was tungsten-inert-gas (TIG) welded. All specimens appeared bright and shiny with no signs of surface corrosion apparent.

f. <u>CPR 6</u>. Specimens 056 to 059 are coupon sections from the Type-304L CRES EB weld joint No. 407. All specimens' metallic surfaces were shiny and bright with no signs of visible corrosion.

g. <u>CPR 7</u>. Specimens 067 to 070 are coupon sections from the Type-304L CRES EB weld joint No. 406. All coupons' metallic surfaces were shiny and bright with no signs of visible corrosion.

h. <u>CPR 8</u>. Specimens 078 to 081 are coupon sections of Types 308L and 304L CRES from the stiffening ring/diaphragm spot-welded subassemblies. All were shiny and bright with no evidence of surface corrosion.

i. <u>CPR 9</u>. Specimens 300 to 307 are coupon sections of Type-304L CRES from the ACS tank liner. All samples were shiny, with a matte finish. No corrosion was noted on these coupons.

j. <u>CPR 10</u>. Specimens 100 to 103 are coupon sections from the tank shell composed of Type-A286 CRES. The metal was still shiny with a minor amount of gray discoloration. No corrosion on the coupon was evident at 45x magnification.

k. <u>CPR 11</u>. Specimens 109 and 110 are coupon sections from TIG-welded joint No. 411 of Type-A286 CRES, which was a part of the ACS tank girth weld. The metal was shiny except for a gray discoloration in the HAZ along each side of the weld. No corrosion was evident.

1. CPR 12. Specimens 112 and 113 are coupon sections of the TIG-welded joint No. 417 composed of Type-A286 CRES, which was welded to Type-304L CRES. The specimen metal surfaces were shiny and bright with no evidence of corrosion.

m. <u>CPR 13</u>. Specimens 115 and 116 are sections of Type-308L CRES filler wire used in the diaphragm stiffening ring. The wire was shiny except for dark-gray bands about 1/2 in. from each end. No other corrosion was evident.

n. <u>CPR 14</u>. Specimens 118 to 121 are coupon sections of Type-304L CRES from the liner-diaphragm assembly with a gold-nickel brazed Type-308L CRES filler wire. All coupons were shiny and bright with no evidence of corrosion.

o. <u>CPR 15.</u> Specimens 127 and 128 are pieces of the Mylar plastic sensor disc from the vapor-detection assembly. Both specimens dissolved in the hydrazine propellant after three hours of testing. (The hydrazine became light yellow in color and contained 0.016 mg of iron.)

p. <u>CPR 16.</u> Specimens 130 and 131 are coupon sections of Type-304L CRES from the liquid outlet housing to the tank-half EB-welded joint No. 404. Specimen 130 was shiny except for some etching on the bottom of the coupon and a gray discoloration in the HAZ along each side of the weld. Specimen 131's surface was shiny and bright except for an irregular surface on the top. No corrosion was seen on either coupon.

q. <u>CPR 17</u>. Specimens 133 and 134 are fluorinated-ethylenepropylene-(Teflon-) coated samarium-cobalt magnets from the vapor-detection assembly. The Specimen 133 coating was intact, but uneven. No corrosion was evident. The posttest weight was significantly lower than the pretest weight. A reason for the loss was not readily apparent. However, it was observed that the magnet did influence the action of the scale balance; an accurate weight was obtained only after neutralizing the magnetic influence. The specimen-134 Teflon coating was irregular but apparently intact. There were dark spots on the magnet surface, but it was not possible to identify them as corrosion spots. r. <u>CPR 18</u>. Specimens 200 and 201 are coupon sections of Type-17-4 PH CRES, H 1050 temper, and electropolished, from the piston in the flow-equalizer valve. The surface of the specimens was bright with no evidence of corrosion.

s. <u>CPR 19</u>. Specimens 210 and 211 are electropolished and chrome-plated sections of Type-17-4 PH CRES, H 1050 temper, from the shaft end of the flow-equalizer valve. The specimens were shiny and bright with no corrosion evident.

t. <u>CPR 20</u>. Specimens 220 and 221 are springs of electropolished Type-17-4 PH CRES, CH 900 temper, from the flow-equalizer valve. Specimen 220 is bright with a light-gray tarnish. No other corrosion was evident. Specimen 221 was shiny with no corrosion evident.

u. <u>CPR 21</u>. Specimens 230 and 231 are coupon sections of Type-17-4 PH CRES, TIG welded and electroplated, from the shaft end assembly of the flow-equalizer valve. The metal was shiny and bright with some etching near the coupon identifying number. No corrosion was seen.

y. <u>CPR 22</u>. Specimens 240 and 241 are bourdon tubes of Inconel 902 (Ni Span C) from the pressure-switch assembly. The metal surfaces have an even gray oxidized coating with no corrosion evident.

w. <u>CPR 23</u>. Specimens 250 and 251 are coupon sections and pieces of Type-347 CRES bar stock from the propellant distribution manifold fitting. Specimen 250 was shiny with a very light-gray mottling perceptible on the surfaces. No corrosion was apparent. Specimen 251, with circular machining marks, was shiny and bright. No corrosion was seen.

x. <u>CPR 24</u>. Specimens 260 and 261 are of Type-347 CRES tubing. A light-gray tarnish was evident on the surface, but no corrosion can be seen.

y. <u>CPR 25</u>. Specimens 270 and 271 are coupon sections of Type-347 CRES tubing Astroarc welded (Weld No. 78) to Type-347 CRES tubing. The tubing was shiny except for a gray discoloration in the HAZ on each side of the weld. No corrosion was evident.

z. <u>CPR 26</u>. Specimens 280 and 281 are O-rings made of Parker seal compound EPR 515. Microscopic examination of the O-rings revealed no crazing or cracking; the surface appeared smooth and unbroken. There was no Krytox 240AC coating on the specimens before cleaning.

B. DETAILS AND SUMMARIES OF POSTTEST ANALYSES AND RESULTS

The posttest analyses and results are summarized in Table 2. The duration of the test units in storage, test temperatures in degrees Celsius, and capsule posttest pressures at test temperature in N/cm² are given. The specimen material, configuration, and weight change in milligrams are listed. The percentage decomposition of hydrazine and the gas evolution rate in cm³ x 10^{-3} day⁻¹. cm⁻² are also given.

Data on the individual test units, test specimens, and the hydrazine propellant is given in Table 3. The BAT number is identified with the test unit number. The initial weight in grams and the change in weight is given for each specimen. The analysis of the hydrazine is given in milligrams for dissolved iron (Fe) and by percent for water (H₂0) and ammonia (NH₃). The hydrazine decomposition into the noncondensable gases nitrogen (N₂) and hydrogen (H₂) is given as total volume (cm³) at standard temperature and pressure (STP); the gas evolution rate, both uncorrected and control corrected, is in cm³ x 10^{-3} .

C. PROPELLANT CONTROLS

Table 4 presents data on the hydrazine propellant, unit number, days on test, temperature in degrees Celsius, capsule pressure at test temperature, and the H₂O and NH₃ analyses, by percent, along with information on hydrazine decomposition. The noncondensable gases (N₂ and H₂) are listed as total volume cm³ at STP and the rate as cm³ x 10^{-3} .

D. SURFACE ANALYSIS

1. Introduction

As noted in Table 2, in a few pairs (or groups) of specimens, one of the test units shows a significantly higher gas evolution than the others in that group. For example, specimen BA 008 had twice the gas evolution rate of any of the others in the CPR 1 group. In the CPR 4 group, one of the Lee plugs greatly enhanced the decomposition of hydrazine. Other examples can be seen in Table 2.

Two sets of specimens were chosen for very scrupulous examination by X-ray photoelectron spectroscopy (XPS) and the scanning electron microscope (SEM). Both sets are from the primary containment system (Program B) and both are of 304L CRES. The specimens examined were BA 005 and BA 008 of the CPR 1 group, and BA 305 and BA 307 of the CPR 9 group. In each case an untreated specimen, i.e., one not exposed to hydrazine, was used as a control in the analysis.

2. XPS Techniques

The six specified stainless-steel samples were submitted for surface analysis by the XPS technique; Figure 3-1 shows the JPL XPS laboratory. The samples were cut from the original strips using metal shears. Then, immediately prior to insertion into the XPS spectrometer, each sample was cleaned ultrasonically in absolute ethanol for 10 minutes and dried with flowing nitrogen. The following samples were analyzed:

<u>Set I</u>	Set II
CPR 1 (CONTROL)	CPR 9 (CONTROL)
BA 005	BA 305
BA 008	BA 307

3-4



Figure 3-1. JPL XPS Laboratory

Analyses were performed with a modified Hewlett-Packard 5950A electron spectrometer under computer control. Photoemitted electrons, characteristic of the surface composition of the sample, were formed by interaction with 1486.6-eV incident photons from an aluminum K α radiation source. For the instrumentation employed, the measured photoemission represents an average signal over an area of approximately 1 mm x 5 mm. Although the photoemitted electrons from a given atomic core level may sometimes originate as much as 100 Å below the sample surface, the immediate surface region of the sample will actually contribute significantly more to the measured signal due to an exponential attenuation (with depth) of photoelectrons by the solid.

For each sample, a wide energy-range (100 to 1300 eV) scan was run to determine the major contributors to the total photoelectron spectrum, followed by careful measurement of a 20 to 40 eV binding energy-range characteristic of each of eight elements of major interest. Each such run for a given sample took a total of 15.2 hours of instrument time to obtain adequate statistics. Photoelectron spectra (PES) of the following elements were obtained:

Energy	Measured Binding
Levels	Energy Range, eV
1s	275 to 295
1s	390 to 410
1s	523 to 543
$2p_3/2$, $2p_1/2$	560 to 600
$2p_{3/2}, 2p_{1/2}$	625 to 665
$2p_{3/2}, 2p_{1/2}$	695 to 735
$2p_{3/2}, 2p_{1/2}$	845 to 895
$2p_{3/2}, 2p_{1/2}$	1010 to 1050
	Energy Levels 1s 1s 1s 2p3/2, 2p1/2 2p3/2, 2p1/2 2p3/2, 2p1/2 2p3/2, 2p1/2 2p3/2, 2p1/2 2p3/2, 2p1/2 2p3/2, 2p1/2

The XPS BE (binding energy) spectra can be used to identify different elements on a surface and to distinguish between the same element in different chemical environments. For example, oxidation of a metal causes an effective unbalanced positive change localized on the metal atom and the remaining electrons are therefore held more tightly; i.e., their binding energy increases.

3. Results

For the various spectral regions examined in this investigation, some general observations are presented below.

a. <u>Carbon 1s Region</u>. Carbon is the major constituent of the surface of all samples, amounting to 40 +4 atomic percent (at.%) for five samples and to a significantly higher 68.8 at.% for CPR 1 (which also gave anomalous results in almost all the other measurements). The primary peak in the carbon spectra comes from aliphatic carbon compounds, while there are also substantial intensities in the regions expected for carbon-nitrogen compounds ($\approx 286 \text{ eV}$) and carbon-oxygen compounds ($\approx 288 \text{ eV}$). A quantitative measure of the relative amounts of these three carbon species may be obtained from the application of computer curve-fitting routines to the individual spectra.

b. Oxygen 1s Region. The apparent surface concentration of oxygen is approximately the same $(46 \pm 3 \text{ at.}\%)$ for all samples, with the exception of CPR 1(24.7\%). In the case of oxygen, a variety of metal oxides and hydroxides contribute to the measured signals and, in the absence of adequate reference data, it is very difficult to make any specific assignments to the various peaks and shoulders observed. It should be noted that the oxygen 1s binding energy for CPR 1 is shifted to a higher energy than that observed for the other five samples, suggesting a significantly different distribution of metal oxides/hydroxides for this particular sample, when compared to the others.

c. <u>Nitrogen 1s Region</u>. The apparent surface concentration of nitrogen is observed to be approximately the same $(2.8 \pm 0.5 \text{ at.}\%)$ for all six samples. The measured binding energy of the nitrogen peak is, for all samples, consistent with that to be expected for protonated amines and amino polymers.

d. <u>Chromium 2p Region</u>. In both Sets I and II, the apparent surface concentration of chromium for the samples exposed of the control specimens, CPR 9 (4.3 at.%) or CPR 1 (0.8 at.%, anomalously low). The predominant species present in all samples are metallic chromium (relatively small) and chromium oxide or hydroxide (relatively large). The ratio of metallic to oxidized chromium is approximately the same for all samples.

e. <u>Manganese 2p Region</u>. In both Sets I and II, the apparent surface concentration of manganese for the samples exposed to hydrazine $(1.0 \pm 0.2 \text{ at.\%})$ is found to be greater than that for either of the control specimens CPR 9 (0.5 at.\%) or CPR 1 (0.1 at.\%, again low). The predominant species present in all samples is oxidized manganese as the monoxide; hydroxides may be present as well, due to the close similarity of binding energies.

f. Iron 2p Region. The dominant surface species present in all samples are metallic iron and ferric oxide; iron hydroxides are also possible since oxides and hydroxides exhibit similar binding energies. In both Sets I and II, the ratio of metallic iron to oxidized iron is much greater for the samples subjected to hydrazine than for the control specimens. In Set II, the control specimen CPR 9 has a greater total iron surface concentration (3.9 at.%) than that for the samples contacted by hydrazine (3.1 +0.8 at.%). In Set I just the opposite occurs; the total surface iron for CPR 1 (1.0 at.%) appears to be anomalously low.

g. <u>Nickel 2p Region.</u> In all cases, the primary peak in the nickel spectrum is due to metallic nickel; a much less intense peak at higher binding energy is due to the monoxide. In both Sets I and II, the apparent surface concentration of nickel was approximately the same (0.3 at.%) for the samples exposed to hydrazine. The nickel concentration for control sample CPR 9 was somewhat lower (0.1 at.%), while for CPR 1 it was exceptionally low (0.2 at.%).

h. <u>Zinc 2p Region</u>. In all cases, the surface zinc is present in oxidized form, probably as the simple oxide. In Set I, the apparent surface concentration of zinc is approximately equal (0.2 at.%) for all three samples. In Set II, the control specimen, CPR 9, shows 2 to 3 times more zinc (0.6 at.%) than the samples subjected to hydrazine.

3-7

4. Discussion

Both the carbon and nitrogen binding energy spectra give strong evidence of surface contamination by carbon-nitrogen compounds such as amino polymers. It should be noted that although these samples were apparently stored in polyethylene bags in pretreatment steps, the <u>final</u> pretest step consisted of heat-sealing in nylon, a process which could readily provide the observed amino polymers on the surface.

The anomalous results for control CPR 1 were confirmed by a second run which gave results identical to the original run. This specimen was more reduced in oxygen and metals concentration, but more heavily contaminated by carbon than any of the other five samples. Therefore, more meaningful comparison may probably be made using CPR 9 as a control for both sets of samples subjected to hydrazine.

The overall picture is that, upon exposure to hydrazine, oxidized iron is removed, leaving a surface richer in the protective chromium oxide. A more thoughtful analysis may be aided by taking into account the information presented in Reference 6, and the references contained therein.

5. Scanning Electron Microscopy (SEM) Examination

These same specimens were then examined by SEM to look for differences in surface morphology. The surfaces of specimens BA 005 and BA 008 were identical in appearance when examined at 50x and 500x magnifications. When compared to the CPR 1 control, the specimens exposed to hydrazine appeared to have very minor surface pitting. The surfaces of specimens BA 305 and BA 307 also were identical and no differences could be seen when compared to the CPR 9 control.

In addition, the specimen BA 036, Lee plug, was thoroughly examined by SEM because of the greater decomposition rate of its test unit compared to the three other replicates. A direct comparison to specimen control BA 038 indicated no differences.

6. Conclusions

These highly detailed and sensitive analyses failed to indicate the causes for differences observed in the rates of hydrazine decomposition between pairs of nominally identical specimens. Some of the more subtle differences in surface character were unfortunately masked by the presence of significant contamination by carbon-nitrogen compounds from the sealed nylon storage bags. The thickness of this carbon-nitrogen layer is such (<100Å) that it would have no effect on either the rate of hydrazine decomposition or the effect of corrosion of the coupons. SEM examination of the above-mentioned six coupons and Lee plug BA 036 again revealed no cause for the difference in decomposition rates.

SECTION IV

CONCLUSIONS

With few exceptions, mainly attributable to catalysis, possible contamination, or inherent sample-to-sample variation, the rate of hydrazine decomposition in these tests was very low -- producing less than 1.0 cc of gas per year per cm² of specimen area.

The degree of corrosion of the metal coupons was virtually unmeasurable in all instances. The elastomer EPR-515 did not appear to degrade, and the Mylar film dissolved as expected.

A. PROGRAM A: 6 MONTHS STORAGE

The rates of hydrazine decomposition were low in most test units -- less than 1.0 cm^3 gas per year per cm² of specimen area. The following were a few exceptions to the low rates:

- (1) 304L liner-diaphragm with Au-Ni brazed 308L filler wire (BA 118-121), 1.0 to 3.7 cm³/yr/cm²
 - (a) Possible catalysis by Au-Ni braze
 - (b) Rate based on total coupon area.
- (2) FEP-coated Sm-Co magnets (BA 133-134), 3.7 to $13.2 \text{ cm}^3/\text{yr/cm}^2$
 - (a) FEP coating intact
 - (b) Possible permeation and catalysis.
- (3) ERP-515 O-rings
 - (a) Catalysis by carbon black used in compounding elastomer
 - (b) Very rapid decomposition, but area rate not meaningful.
- (4) Mylar film (BA 127-128), $3.5 \text{ cm}^3/\text{yr/cm}^2$
 - (a) Film dissolved
 - (b) Area rate not meaningful.

The corrosion of metallic coupons was minimal and only very light tarnish was seen on a few specimens. The weight changes of coupons were negligible, and dissolved iron in the propellant was almost unmeasurable. Two nonmetals were included in Program A:

- (1) Mylar film (BA 127-128), which dissolved as expected.
- (2) EPR-515 O-rings (BA 280-281), which appeared unchanged after exposure to hydrazine.

B. PROGRAM B: 24 MONTHS STORAGE

The rates of hydrazine decomposition were low in most test units -- less than 1.0 cm^3 gas per year per cm^2 of specimen area. There were a few exceptions to low rates, but the results were not consistent:

- (1) 304L liner/diaphragm girth weld (BA 017), 1.3 cm³/yr/cm²
 - (a) Only one of four specimens produced an anomalously large volume of gas
 - (b) Possible contamination or sample-to-sample variation.
- (2) 304L Lee plug (BA 036), $5.2 \text{ cm}^3/\text{yr/cm}^2$
 - (a) Only one of four specimens showed a high rate of decomposition
 - (b) Possible contamination or sample-to-sample variation.

(3) 304L EB weld No. 407 (BA 058-BA 059), $2.3-3.3 \text{ cm}^3/\text{yr/cm}^2$

- (a) Two specimens at 43° C showed low rate of gas formation
- (b) Possible effect of 60^oC storage
- (c) Possible contamination or sample-to-sample variation.
- (4) 304L EB weld No. 406 (BA 070), $1.8 \text{ cm}^3/\text{yr/cm}^2$
 - (a) Only one of two at 60°C was high
 - (b) Possible contamination or sample-to-sample variation.

None of the specimens appeared to corrode, and only a very light tarnish was seen on a few specimens. Weight changes of the coupons were negligible, and the dissolved iron in the propellant was almost unmeasurable.

In general, the results of this study agree very well with the JPL/NASA long-term compatibility program (References 2 and 3). The Type-304L CRES chosen for the primary containment of hydrazine in the propellant tank appears to be entirely suitable for use in systems requiring at least a 2-year service life. In the secondary containment side of the tank, the A286 CRES used in the outer pressure vessel is compatible with hydrazine for 6 to 12 months exposure. The Au-Ni braze material, FEP-coated Sm-Co magnets, and the EPR 515 O-rings have been shown to cause hydrazine decomposition that could result in an undesirable gas pressure buildup which must be accomodated in the system design. However, these materials would be in contact with hydrazine only if a leak occurred in the primary containment system.

SECTION V

DATA TABLES

The tables are, generally, self-explanatory. The following comments are given to expand on certain topics.

The strain gauge data shown in Table 2 indicate that this is not a reliable method of determining capsule pressures of less than one atmosphere. The strain gauges are normally calibrated at positive pressure only, and an extrapolation is made to zero pressure. Attempts to calibrate a capsule-mounted strain gauge at subatmospheric pressure produced results that indicated random shifting of the calibration line. At pressures greater than one atmosphere, the strain gauge data agree very well with the actual pressures found in the capsules. Posttest recalibration of several capsules indicates that while the zero point may shift, the sensitivity is maintained during handling and testing.

The decomposition of hydrazine in the control capsules (Table 4) presumably occurs through homogeneous (bulk) catalysis; glass should not act as an active surface for hydrazine decomposition. Purified hydrazine contains very little dissolved iron (a known catalyst) and, therefore, the rate of decomposition is predictably slow. With the introduction of a metallic specimen, there is the possibility of an active surface and heterogeneous (surface) catalysis. If metal is dissolved from the surface of the specimen, it is possible for both reaction mechanisms to occur. Obviously, from an inspection of some of the results (Tables 2 and 3), there are metallic surfaces which are not catalytically active towards hydrazine, especially Type-304L CRES Alloy.

	Test Numbers					
CPR	BAT	Test Unit	Material Compatibility Test Specimen Description			
1	001-008	4001-4008	304L Arde diaphragm			
2	014-017	4009-4012	304L/304L liner/diaphragm girth weld			
3	023-026	4013-4016	347 burst disc			
4	034-037	4017-4020	304L Lee plug			
5	045-048	4021-4024	304L/304L liner/diaphragm outlet housing TIG weld			
6	056-059	4025-4028	304L/304L EB weld #407			
7	067-070	4029-4032	304L/304L EB weld #406			
8	078-081	4033-4036	308L/304L ring/diaphragm spot welded			
9	300-307	4037-4044	304L liner			
10	100-103	4045-4048	A286 tank shell			
11	109-110	4049-4050	A286/A286 girth TIG weld #411			
12	112-113	4051-4052	304L/A286 polar TIG weld #417			
13	115-116	4053-4054	308L stiffening ring			
14	118-121	4055-4058	308L/304L wire/diaphragm, Au-Ni braze			
15	127-128	4059-4060	Mylar sensor disc			
16	130-131	4061-4062	304L/304L EB weld #404, liquid outlet housing to tank half			
17	133-134	4063-4064	Samarium-cobalt magnet, FEP coated			
18	200-201	4065-4066	17-4 PH, H 1050 temper, electropolished			
19	210-211	4067-4068	17-4 PH, H 1050 temper, electropolished, chrome plated			
20	220-221	4069-4070	17-4 PH, CH900 temper, spring			
21	230-231	4071-4072	17-4 PH/17-4 PH, TIG weld, shaft end			
22	240-241	4073-4074	Inconel 902, Ni span C, Bourdon tube			
23	250-251	4075-4076	347 manifold fitting			
24	260-261	4077-4078	347 tube, annealed			
25	270-271	4079-4080	347/347 Astro-arc weld #78			
26	280-281	4081-4082	EPR 515, Parker seal, O-ring, Krytox coated			

Table 1. Listing of Coupon Test Numbers and Description

		Days on Test	Test Temp., oC	Capsule Pressure at Test Temp., N/cm ²	Specimen			Propellant		
BAT No.	Test Unit				(Strain Gage Reading)	Material	Configuration	Weight Change mg	Decompo- sition, %	Gas Evolution cc x 10 ⁻³ · day ⁻¹ · cm ⁻²
BA001	4001	807	43	0.77	(0)	304L CRES	ACS Diaphragm	-0.4	0.02	0.01
BA002	4002	807	43	0.72	(0)		"	0.0	0.02	-0.01
BA003	4003	807	43	0.99	(0)	**	. 75	-0.2	0.03	0.06
BA004	4004	807	43	0.65	(0)	"	"	-0.1	0.01	-0.01
BA005	4005	765	60	1.70	(0)	39	**	+2.5	0.02	-0.10
BA006	4006	765	60	2.85	(0)	**	"	-0.2	0.05	0.10
BA007	4007	765	60	2.12	(0)	"	"	0.0	0.04	-0.03
BA008	4008	765	60	3.10	(0)	**	39	-0.3	0.06	0.19
BA014	4009	807	43	0.89	(0)	"	Liner/diaphragm,	-0.8	0.02	0.02
BA015	4010	807	43	1.41	(0)	**	Girth weld	-1.2	0.02	0.10
BA016	4011	807	60	3.25	(0)	**	**	-0.9	0.06	0.10
BA017	4012	765	60	33.39	(35.1)	,,	"	-0.5	0.97	3.52
BA023	4013	378	43	2.07	(5.0)	CRES 347	Burst Disc	-0.4	0.12	0.74
BA024	4014	378	43	2.48	(0)	**	"	-0.4	0.16	0.95
BA025	4015	378	60	9.81	(0)	"	"	-0.4	0.41	3.91
BA026	4016	378	60	8.24	(0)	"	"	-0.1	0.35	3.14
BA034	4017	835	43	1.05	(0)	304L CRES	Lee Plug	-0.4	0.03	0.61
BA035	4018	835	43	1.95	(0)	**	77 .	-0.3	0.04	2.40
BA036	4019	835	60	10.98	(5.5)	39	"	+15.5	0.26	14.10
BA037	4020	835	60	3.77	(0)	"	"	+7.1	0.04	2.62
BA045	4021	835	43	1.12	(8.0)	**	TIG weld	+0.1	0.03	0.29
BA046	4022	835	43	1.30	(0)	"	**	-0.3	0.04	0.33
BA047	4023	835	60	2.80	(0)	"	"	-0.2	0.06	0.28
BA048	4024	835	60	7.44	(2.5)	**	"	0.0	0.14	2.90
BA056	4025	771	43	2.73	(0)	"	EB weld #407	-1.0	0.09	0.44
BA057	4026	771	43	6.81	(2.1)	"	"	-1.0	0.25	1.18
BA058	4027	729	60	49.35	(38.6)	**	,,	-1.1	1.40	9.00
BA059	4028	715	60	35.34	(29.6)	**	22	-1.2	1.09	6.22

Table 2. Summary of Analyses and Results
							Specimen		P	ropellant
BAT No.	Test Unit	Days on Test	Test Temp., °C	Capsule Pressure at Test Temp., N/cm ²	(Strain Gage Reading)	Material	Configuration	Weight Change mg	Decompo- sition, %	Gas Evolution cc x 10 ⁻³ · day ⁻¹ · cm ⁻²
BA067	4029	771	43	1.27	(0)	304L CRES	EB weld #406	-0.7	0.02	0.14
BA068	4030	771	43	2.97	(3.4)	"	"	-0.8	0.10	0.49
BA069	4031	757	60	8.10	(0)	39	**	+142.2 ^a	0.28	1.28
BA070	4032	715	60	25.15	(16.5)	"	**	-0.8	0.77	4.80
BA078	4033	799	43	1.05	(0)	308L/304L CRES	Ring/diaphragm,	-0.4	0.03	0.05
BA079	4034	799	43	0.85	(0)	77	Spot weld	0.0	0.02	0.02
BA080	4035	785	60	6.17	(0)	"	"	0.0	0.16	0.57
BA081	4036	785	60	9.37	(1.8)	"	"	0.0	0.26	1.07
BA300	4037	807	43	0.85	(0)	304L CRES	Liner	-0.9	0.02	0.03
BA301	4038	765	43	b	(14.0)	"	**	+0.5		_
BA302	4039	807	43	0.70	(0)	**	**	-0.6	0.01	< 0.01
BA303	4040	807	43	0.74	(0)	"	**	-1.2	0.02	< 0.01
BA304	4041	765	60	с	(0)	**	"	-0.7	_	-
BA305	4042	765	60	3.81	(0)	"	**	+0.6	0.08	0.41
BA306	4043	765	60	3.59	(0)	3 9		0.0	0.08	0.31
BA307	4044	765	60	2.50	(0)	"	"	-0.5	0.04	0.08
BA100	4045	184	43	0.69	(0)	A286	Tank Shell	-0.7	< 0.05	0.13
BA101	4046	245	43	2.02	(0)	"	"	-0.3	0.15	0.68
BA102	4047	308	43	1.21	(0)	"	"	-0.9	0.10	0.14
BA103	4048	365	43	1.02	(0)	**	"	+0.2	0.04	0.14
BA109	4049	245	43	1.43	(2,1)	**	TIG weld #411	-0.7	0.10	0.31
BA110	4050	365	43	1.74	(1.0)	**	**	-0.3	0.07	0.41
BA112	4051	245	43	3.04	(2.8)	A286/304L	TIG weld #417	-0.9	0.03	3.67
BA113	4052	365	43	3.57	(1.5)	57	**	-0.4	0.13	2.60
BA115	4053	245	43	1.31	(1.7)	308L	Wire	-0.7	0.01	2.07
BA116	4054	365	43	0.99	(0)	**	**	+0.1	0.08	0.14
BA118	4055	220	43	7.48	(11.6)	308L/304L	Liner/diaphragm,	-1.1	0.36	3.46d
BA119	4056	281	43	11.39	(10.0)	••	Au-Ni Braze	-0.3	0.40	5.05d
BA120	4057	344	43	10.53	(10.7)	"	"	-0.8	0.29	3.63d

Table 2. Summary of Analyses and Results (continuation 1)

							Specimen		Propellant	
BAT No.	Test Unit	Days on Test	Test Temp., °C	Capsule Pressure at Test Temp., N/cm ²	(Strain Gage Reading)	Material	Configuration	Weight Change mg	Decompo- sition, %	Gas Evolution cc x 10 ⁻³ . day ⁻¹ . cm ⁻²
BA121	4058	401	43	29.04	(29.1)	308L/304L	Au-Ni Braze	-0.2	0.70	9.98d
BA127	4059	365	43	2.70	(2.4)	Mylar Film	(Specimen Dissolved)	-	0.16	9.15 ^e
BA128	4060	365	43	2.44	(1.4)	**	**		0.09	9.43e
BA130	4061	308	43	1.61	(0)	304L	EB weld #404	-1.1	0.01	0.53
BA131	4062	340	43	0.97	(0)	**	"	-1.3	0.08	0.05
BA133	4063	308	43	4.67	(0)	Samarium-Cobalt	Magnet, FEP-	-28.7	0.11	36.17
BA134	4064	365	43	2.25	(0)	**	Coated	+0.8	0.11	10.46
SE200	4065	308	43	2.10	(0)	17-4 PH, H1050	Valve, Electro-	-0.5	0.02	0.90
SE201	4066	365	43	0.92	(0)	**	Polished	+0.1	0.04	0.08
SE210	4067	365	43	0.88	(3.4)	,,	Valve, E.P.,	0.0	0.05	0.04
SE211	4068	184	43	0.58	(0)	**	Chrome plated	-1.1	< 0.05	0.03
SE220	4069	184	43	0.81	(6.9)	17-4 PH, CH900	Spring	-2.4	< 0.05	0.26
SE221	4070	36 5	43	1.05	(5.5)	**	9 9	+0.6	0.05	0.16
SE230	4071	308	43	0.56	(0)	17-4 PH	Valve, TIG weld	-0.4		0.01
SE231	4072	365	43	0.76	(0)	**	99	-0.4	0.04	0.01
SC240	4073	308	43	1.29	(0)	Inconel 902	Bourdon tube	-0.3	0.01	0.11
SC241	4074	365	43	1.71	(0)	**	99	-0.8	0.08	0.10
BA250	4075	245	43	1.04	(1.7)	347	Bar Stock	-1.2	0.06	0.18
BA251	4076	365	43	1.52	(0)	**	"	-0.2	0.06	0.35
BA260	4077	245	43	0.76	(4.5)	**	Tube, annealed	-0.8	< 0.01	0.16
BA261	4078	365	43	0.81	(3.4)	**	29	-0.2	0.04	0.04
BA270	4079	245	43	1.97	(3.8)	**	Astro arc weld #78	-0.1	0.16	0.68
BA271	4080	365	43	7.34	(2.1)	**	"	-0.4	0.26	3.41
BA280	4081	37	43	27.34	(27.5)	EPR 515	"O" Ring,	+2.1	0.93	416.3 ^f
BA281	4082	65	43	44.07	(44.0)	"	Krytox Coated	+3.5	1.62	370.0 ^f

Table 2. Summary of Analyses and Results (continuation 2)

^aProbable error in pretest weighing.

^dRate of decomposition is proportional to the area of exposed gold-nickel braze.

^eMylar film dissolved in propellant; area rate is not relevant.

^cCapsule broke in breaker fixture; gas lost.

^bCapsule tip had microscopic leak; gas data meaningless.

^fDecomposition catalyzed by carbon black used in compounding; area rate values are meaningless.

				Anal	ysis of l	Propellan	it ^{a,b}		Rate	······		
BAT No.	Test Unit	Specimo Initial g	en Weight Change g	Fe	e ppm	н ₂ 0 %	NH3 %	Total cc STP	Uncorrected cc x $10^{-3} \cdot day^{-1}$	Corrected for Control cc x 10 ⁻³ • day ⁻¹	Specimen Surface Area cm ²	Area Rate cc x 10 ⁻³ • day ⁻¹ • cm ⁻²
BA001	4001	1.8307	-0.0004	<0.02	< 0.5	0.63	0.02	0.67	0.83	0.16	19.4773	0.01
BA002	4002	1.8808	0.0000	< 0.02	<0.5	0.66	0.02	0.45	0.56	-0.11	19.4926	-0.01
BA003	4003	1.8674	-0.0002	< 0.02	<0.5	0.66	0.02	1.51	1.87	1.20	19.4926	0.06
BA004	4004	1.8745	-0.0001	< 0.02	< 0.5	0.68	0.01	0.39	0.48	-0.19	19.4773	-0.01
BA005	4005	1.8774	+0.0025	< 0.02	< 0.5	0.68	0.01	1.59	2.08	-1.92	19.4621	-0.10
BA006	4006	1.8189	-0.0002	< 0.02	< 0.5	0.65	0.03	4.60	6.01	2.01	19.4926	0.10
BA007	4007	1.8308	0.0000	<0.02	< 0.5	0.84	0.03	2.58	3.37	-0.63	19.4926	-0.03
BA008	4008	1.8121	-0.0003	< 0.02	<0.5	0.65	0.04	5.88	7.69	3.69	19.4773	0.19
BA014	4009	17.9183	-0.0008	c		0.58	0.02	1.15	1.43	0.76	36.55	0.02
BA015	4010	19.7705	-0.0012	<u></u>		2.95	0.01	3.52	4.36	3.69	37.59	0.10
BA016	4011	20.5209	-0.0009			0.56	0.04	6.57	8.14	4.14	39.57	. 0.10
BA017	4012	19.8750	-0.0005	0.125	3.1	0.86	0.60	110.57	144.54	140.54	39.92	3.52
BA023	4013	10.5977	-0.0004	0.02	0.5	0.63	0.10	6.17	16.32	15.65	21.1697	0.74
BA024	4014	9.8477	-0.0004	0.02	0.5	0.50	0.14	7.69	20.34	19.67	20.7925	0.95
BA025	4015	10.4620	-0.0004	0.02	0.5	0.62	0.32	32.86	86.93	82.93	21.2021	3.91
BA026	4016	10.5614	-0.0001	0.02	0.5	0.70	0.27	28.31	74.89	70.89	22.5703	3.14
BA034	4017	0.6948	-0.0004	< 0.02	<0.5	0.52	0.02	1.95	2.34	1.67	2.7429	0.61
BA035	4018	0.6995	-0.0003	< 0.02	<0.5	8.21	0.02	5.99	7.17	6.50	2.7040	2.40
BA036	4019	0.6989	+0.0155	< 0.02	<0.5	0.76	0.16	35.60	42.63	38.63	2.7392	14.10
BA037	4020	0.7015	+0.0071	<0.02	< 0.5	0.71	0.02	9.27	11.10	7.10	2.7068	2.62
BA045	4021	2.0661	+0.0001	< 0.02	<0.5	0.57	0.02	2.18	2.61	1.94	6.61	0.29
BA046	4022	2.1141	-0.0003	< 0.02	< 0.5	2.33	0.03	2.50	2.99	2.32	7.04	0.33
BA047	4023	2.1348	-0.0002	< 0.02	≼0.5	0.93	0.04	5.01	6.00	2.00	7.03	0.28
BA048	4024	2.0952	0.0000	< 0.02	<0.5	0.90	0.08	19.80	23.71	19.71	6.80	2.90
BA056	4025	11.9468	-0.0010	< 0.02	<0.5	1.03	0.06	7.86	10.19	9.52	21.75	0.44
BA057	4026	12.4409	-0.0010	< 0.02	< 0.5	1.21	0.18	20.57	26.68	26.01	22.06	1.18
BA058	4027	12.3262	-0.0011	0.050	1.3	0.70	0.89	155.32	201.45	197.45	21.93	9.00
BA059	4028	12.3092	-0.0012	0.125	3.1	0.86	0.74	108.45	140.66	136.66	21.96	6.22
BA067	4029	10.5426	-0.0007	_		0.81	0.01	2.76	3.58	2.91	21.42	0.14

Noncondensable Gas $(N_2 + H_2)$

5-6

										Noncondensable Ga	s (N ₂ + H ₂)	
				Anal	ysis of I	ropellar	nt ^{a,b}		Rate	······································	<u>, p</u>	
BAT No.	Test Unit	Initial g	change g	Fe	e ppm	н ₂ 0 %	NH3 %	Total cc STP	Uncorrected cc x 10 ⁻³ • day ⁻¹	Corrected for Control cc x 10 ⁻³ • day ⁻¹	Specimen Surface Area cm ²	Area Rate cc × 10 ⁻³ • day ⁻¹ • cm ⁻²
BA068	4030	11.2825	-0.0008			0.98	0.07	8.50	11.02	10.35	21.05	0.49
BA069	4031	11.6180	+0.1422d	_		0.86	0.20	24.00	31.70	27.70	21.59	1,28
BA070	4032	11.5931	-0.0008	< 0.02	< 0.5	0.92	0.52	76.87	107.51	103.51	21.56	4.80
BA078	4033	5.5458	-0.0004	<0.02	< 0.5	1.18	0.02	1.66	2.08	1.41	26.8885	0.05
BA079	4034	5.6452	0.0000	<0.02	< 0.5	1.26	0.02	0.99	1.24	0.57	28.3715	0.02
BA080	4035	5.6520	0.0000	< 0.02	< 0.5	0.52	0.11	15.23	19.40	15.40	27.0450	0.57
BA081	4036	5.5455	0.0000	< 0.02	< 0.5	0.55	0.18	26.42	33.66	29.66	27.7521	1.07
BA300	4037	5.2560	-0.0009	<0.02	< 0.5	0.80	0.02	1.06	1.31	0.64	20.0694	0.03
BA301	4038	4.9612	+0.0005	< 0.02	< 0.5	0.65	0.04	<u> </u>	_	-	20.0753	_
BA302	4039	4.3396	-0.0006			0.84	0.01	0.61	0.76	0.09	20.0754	<0.01
BA303	4040	4.8056	-0.0012	_		1.56	0.02	0.55	0.68	0.01	20.1092	<0.01
BA304	4041	5.1021	-0.0007	< 0.02	< 0.5	0.83	0.04	f	-		20.2661	_
BA305	4042	5.2821	+0.0006	0.02	0.5	0.65	0.02	9.37	12.25	8.25	20.3587	0.41
BA306	4043	5.0690	0.0000	< 0.02	< 0.5	0.62	0.05	7.91	10.34	6.34	20.2475	0.31
BA307	4044	5.0142	-0.0005	< 0.02	<0.5	0.70	0.03	4.36	5.70	1.70	20.0859	0.08
BA100	4045	8.3294	-0.0007	-		-	< 0.05	0.63	3.42	2.75	21.2098	0.13
BA101	4046	8.3596	-0.0003	_		-	0.14	3.67	14.98	14.31	21.1961	0.68
BA102	4047	8.3758	-0.0009	_		-	0.10	1.10	2.90	2.90	21.1961	0.14
BA103	4048	8.3975	+0.0002	_			0.04	1.33	3.64	2.97	21.1825	0.14
BA109	4049	15.0951	-0.0007	_			0.10	1.97	8.04	7.37	23.9895	0.31
BA110	4050	14.6483	-0.0003	_		_	0.06	3.95	10.82	10.15	24.7093	0.41
BA112	4051	4.6890	-0.0009				< 0.05	9.32	38.04	37.37	10.1851	3.67
BA113	4052	4.7486	-0.0004	_			0.10	9.85	26.99	26.32	10.1216	2.60
BA115	4053	2.7318	-0.0007			_	<0.05	3.15	12.86	12.19	5.8837	2.07
BA116	4054	2.7818	+0.0001	_		_	0.08	0.55	1.51	0.84	5.9570	0.14
BA118	4055	6.0872	-0.0011			—	0.30	21.89	99.50	98.83	28.58	3.46 ^g
BA119	4056	5.9381	-0.0003	_		_	0.29	38.47	136.90	136.23	27.00	5.05 ^g
BA120	4057	5.8159	-0.0008	_		_	0.18	34.54	100.41	99.74	27.45	3.63 ^g
BA121	4058	5.8271	-0.0002	0.020	0.5		0.08	110.08	274.51	273.84	27.45	9,98 ^g

Table 3. Details of Analyses and Results (continuation 1)

Table 3. Details of Analyses and Results (continuation 2)

											2 2'	
			TTT * 1 .	Analy	ysis of I	Propellar	it ^{a,b}		Rate			
BAT No.	Test Unit	Initial g	change g	Fe	ppm	н ₂ 0 %	NH3 %	Total cc STP	Uncorrected cc x 10 ⁻³ • day ⁻¹	Corrected for Control cc x 10 ⁻³ • day ⁻¹	Specimen Surface Area cm ²	Area Rate cc × 10 ⁻³ • day ⁻¹ • cm ⁻²
BA127	4059	0.0090	<u></u>	0.016	0.4	<u> </u>	0.14	6.19	16.96	16.29	1.7806	9.15 ^h
BA128	4060	0.0089		_		_	0.07	6.31	17.29	16.62	1.7620	9.43 ^h
BA130	4061	21.7411	-0.0011				0.11	2.43	7.89	7.22	13.7489	0.53
BA131	4062	21.4048	-0.0013	_			0.08	0.48	1.41	0.74	13.5632	0.05
BA133	4063	0.6402	-0.0287	_		_	0.06	15.58	50.58	49.91	1.38	36.17
BA134	4064	0.6172	+0.0008	—		-	0.09	5.51	15.10	14.43	1.38	10.46
SE200	4065	12.1607	-0.0005	_			<0.05	6.41	20.81	20.14	22.4651	0.90
SE201	4066	11.7659	+0.0001	_ `		-	0.04	0.90	2.47	1.80	22.2418	0.08
SE210	4067	10.6931	0.0000	<u> </u>			0.05	0.55	1.51	0.84	22.07	0.04
SE211	4068	10.9716	-0.0011			-	< 0.05	0.24	1.30	0.63	22.15	0.03
SE220	4069	6.8595	-0.0024	_			< 0.05	1.05	5.71	5.04	19.5243	0.26
SE221	4070	6.8879	+0.0006				0.04	1.36	3.73	3.06	19.5243	0.16
SE230	4071	12.1497	-0.0004			-	<0.05	0.24	0.78	0.11	22.07	0.01
SE231	4072	11.9301	-0;0004	-			0.04	0.35	0.96	0.29	22.22	0.01
SC240	4073	6.7289	-0.0003	· _		_	< 0.05	3.13	10.16	9.49	88.232	0.11
SC241	4074	6.7560	-0.0008	-		-	0.07	3.46	9.48	8.81	88.232	0.10
BA250	4075	12.7079	-0.0012	_			0.06	1.12	4.57	3.90	22.2552	0.18
BA251	4076	12.7510	-0.0002	-		-	0.05	3.10	8.49	7.82	22.1961	0.35
BA260	4077	7.1138	-0.0008			·	< 0.05	1.00	4.08	3.41	21.21	0.16
BA261	4078	7.0385	-0.0002			-	0.04	0.52	1.43	0.75	20.95	0.04
BA270	4079	6.7365	-0.0001	-			0.15	3.24	13.22	12.55	18.56	0.68
BA271	4080	6.6650	-0.0004	0.023	0.6	-	0.18	23.40	64.11	63.44	18.58	3.41
BA280	4081	0.5132	+0.0021	0.010	0.3	0.56	0.61	98.60	2664.9	2664.2	6.40	416.3 ⁱ
BA281	4082	0.5141	+0.0035	0.010	0.3	0.55	0.40	153.96	2368.6	2367.9	6.40	370.0 ⁱ

Noncondensable Gas $(N_2 + H_2)$

a Based on actual weight of propellant in capsule.

b Halide level undetectable, i.e., < 0.02 mg.

c – Not measured; data not available.

d Probable error in pretest weighing.

e Capsule tip had microscopic leak; gas data meaningless.

f Capsule broke in breaker fixture; gas lost.

g Rate of decomposition is proportional to the area of exposed gold-nickel braze.

h Mylar film dissolved in propellant; area rate is not relevant.

i Decomposition was catalyzed by carbon black used in compounding; area rate values are meaningless.

							P	ropellant ^a		
					Ana	lysis ^b		No	n-Condensable Gas (N ₂ +	- H ₂)
Test Unit	Days on Test	Test Temp., °C	Capsule Pressure at Test Temp., N/cm ²	(Strain Gage Reading)	н ₂ 0, %	NH ₃ , %	Decomposition, %	Total cc STP	$cc \times 10^{-3} \cdot day^{-1}$	Average
4100	220	43	0.81	(6.0)	_c	< 0.05	< 0.05	1.03	4.68 ^f	
4101	365	43	0.75	(0)	0.74	0.04	0.04	0.26	0.71	
4102	549	43	_	(0)	0.72	0.02	_	_d		
4106	729	43	_	(0)	0.70	0.01		<0.25 ^e	_	
4108	729	43	0.67	(0)	0.71	0.01	0.01	0.45	0.62	0.67
4103	220	60	<1.0	(0)	_	< 0.05	< 0.01	2.08	9.45 ^f	
4104	401	60	2.21	(0)	0.56	0.10	0.11	1.26	3.14	
4105	547	60	2.00	(0)	0.72	0.02	0.03	2.45	4.48	
4107	729	60	1.71	(0)	1.55	0.01	0.02	2.82	3.87	
4109	729	60	2.17	(0)	0.65	0.01	0.02	3.25	4.46	4.00

Table 4. Summary of Analysis of Hydrazine Controls

^aBased on 40.0 cc hydrazine.

^bMetals and halides undetectable.

^c– Not measured; data not available.

^dCapsule broke in test fixture; gas lost.

^eInsufficient quantity of gas to measure.

^fValue not included in average.

SECTION VI

REFERENCES

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- Toth, L., et. al., Propellant Material Compatibility Program and Results, Technical Memorandum 33-779, Jet Propulsion Laboratory, Pasadena, California, August 15, 1976.
- 3. Moran, C. M., and Bjorklund, R. A., <u>Propellant/Material Compatibility</u> <u>Program and Results, Ten-Year Milestone</u>, JPL Publication 82-62, Jet <u>Propulsion Laboratory</u>, Pasadena, California, July 15, 1982.
- 4. Material Specification, <u>Anhydrous Hydrazine for the HAS System</u>, Report 8803-947047, Revision A, <u>Bell Aerospace Textron</u>, <u>Buffalo</u>, <u>New York</u>, undated.
- 5. Chang, E. T., et al., <u>Solubilities of NH₃, CO, CO₂, and SF₆ in Liquid Propellants</u>, The Aerospace Corp., El Segundo, Calif., Air Force Report SAMSO-TR-71-17, November 1970.
- 6. Adams, R. O., "Review of the Stainless Steel Surface," Journal of Vacuum Science Technology, Vol. 1, No. 12, 1983.

APPENDIX A

PRETEST ANALYSIS OF HYDRAZINE

The pretest analysis of Drum H8367 indicated a very high purity which even meets the requirements of the current MIL-P 26536C, Amendment 2, High Purity Grade. The drum was from JPL's supply of hydrazine at ETS and was chosen because of the low CO₂ content. The full pretest analysis is presented in the report form contained in this appendix.

The handling of the propellant during the filling operation, and the process of removing it after completion of the storage period, can influence the CO_2 content through inadvertent exposure to air. A special test capsule was designated as a CO_2 control capsule. It was filled as part of a regular series of capsules being processed. After freezing and thawing, this control capsule was opened and the propellant removed by the standard procedure. The CO_2 content was determined to be 15 ppm, slightly higher than the 9 ppm in the original hydrazine, but considerably lower than the 30 ppm limit in the above-mentioned military specification. Although this determination was not normally part of the posttest procedure, two other test units were analyzed for CO_2 : 4081 contained 5 ppm CO_2 ; 4082 contained 19 ppm CO_2 .

The data from these three capsules indicated that the procedures employed in filling the capsule were adequate to maintain the desired low CO_2 content. The greatest risk for exposure to air occurs during the opening of the capsule and removal and transfer of the hydrazine to the analysis vial.

REPARED BY)	(DATE)	(REPORT NO.)	
ſoth	4 Dec. 1979	79X10201	
Assay By Toulow/Morror	(DATE)	(PROJECT)	
TAY TOT / HOT AN	10 Dec. 1979	N2 ^H 4Compacibility	
SSAY-HYDRAZINE JPL Dru	um H8367		T
Constituent or Property	,	Results	Specification
Hydrazine assay, % by w	eight Note 1	99.4	98.5% min.
Density at 298 K (77°F)	g/cm^3	1.004	
Particulate, mg/cm ³		0.0007	1 milligram/1 lite:
Water plus soluble impu	rities.	0.62%	1.0% max.
% by weight	- -		
Major impurities. % by	weight		
Ammonia (NH _a)		< 0.1%	0.4% max.
Aniline (C.H.NH.)		None detected, n.d.	0.5% max.
Toluene (C.H.CH.)		n.d.	
Carbon Dioxide (CO)	0.0009	50.0 ppm max.
		nd (0.1%	
Other		n.u. \0.1%	
Sulfated Ach % by weigh	·h+	(0.0005	
Atomic Absorption A	malveie	0.0003	
of ach	ularysis		
Diash Diash and motals	CNH (nom)		
JISSOIVEd metals, Mg/	^g ^N 2 ⁿ 4 ^(ppm)	0.12	
Aluminum			
NICKEI		0.15	
Manganese		< 0.03	
Cobalt		ζ 0.03	
Chromium		< 0.01	
Copper		< 0.03	
Zinc		0.03	
Silicon		< 0.1	
Magnesium		0.05	
Sodium		0.05	
Calcium		0.3	
Barium		< 0.1	
Boron		< 10	
Other Potassium		0.03	

JPL 0999-5 (REV 11-68)

			PAGE OF
REPARED BY)	(DATE)	(REPORT NO.)	
Toth Assay By	(DATE)	(PROJECT)	a Mary and a second
Taylor/Moran	10 Dec 197	N ₂ H ₂ Compatibility	
SSAY-HYDRAZINE JPL Dr	um H8367		
Constituent or Proper	ty	Results	Specification
Dissolved anions, Mg	/g		
Fluoride		n.d. <5 ppm	•
Chloride		n.d. <1 ppm	5.0 ppm max.
Sulfate		n.d. <5 ppm	
Nitrate		n.d. <5 ppm	
Nonvolatile residue.	mg/cm ³	< 0.005	
Identification/Histor Specification Storage Container	y	Notes 2, 3	Note 1
Storage container		JPL Drum H836/	
Test Sample (250 ml size)		2 bottles	
NOTES OR REFERENCES	ni antoni Pelanta manana manangi 1993 til til paga sa na men katikan diteri		
 Hydrazine must cor Revision A. Chemi CO₂ requirement of 	nform with Bell A Ical composition 50 ppm maximum	erospace specification 8 requirements listed in S is critical.	803-947047 ection 3.2
2. W/A 4078-1 and shi	lpper E6076; hydr	azine received 7 Decembe	r 1979.
3. Purified or refine "Voyager 1977", ar Martin Marietta Co during CY 1973-197	ed grade hydrazin nd "Mars Viking I orp., Denver, Col 75 period.	e used on NASA-JPL fligh ander 1975". Hydrazine a orado, their specificati	t projects manufactured by on STM NO20,

al Hiller | |

A-3

APPENDIX B

DETAILED LOGS OF ALL TEST UNITS

Log I is a listing of specimens grouped by specimen (BAT) numbers and test unit (JPL) numbers in ascending order. The "Material Description" column also lists the CPR number and the material scheduled for storage. The "Test Duration" column under "Cell" gives the dates of storage at test temperature. The "Refrigeration" column lists the dates for posttest storage in the freezer before analysis. The "Analysis Document" column lists the JPL internal memoranda reporting results of analysis. The "Remark" column lists test temperature (43°C unless otherwise noted), and other information.

Log II is a listing of specimens by ascending capsule number. The "Capsule" column also includes the total internal volume of each capsule. The "Material Description" column lists the date and time of capsule filling. The capsules were then kept in a freezer until the date shown in the "Cell" column, i.e., the beginning of the storage at test temperature. The "Remarks" column lists the volume of hydrazine placed in each capsule.

		n marangan kana pananan kana pananan kana kana k	Report Number79X07500)
			Project <u>Hydrazine Con</u>	patibility
			Classification Unclass	fied
		JET PROF CALIFORNIA I	PULSION LABORATORY	
		su	IMMARY	
	HYD	RAZINE MATER	RIAL COMPATIBILITY	
		TEST SPECT	MEN/CAPSULES	
			1 1 1070	
Prepar	red by <u>L. K. I</u>	στη	Date JULY 1979	
	.		Dote	
		API	PENDIX OR REVISION	
Date	Pages Affected	Appendix OR Revision	Remarks	Changed by
28 May 1980	14	A	Terminate test units: 4081 (SE 280); 4082 (SE 281)	L. Toth
1			1	

NUM	BER	TEST MONTH	TEST I	URATION	ANALYSIS		REV
SPECIMEN	CAPSULE	DESCRIPTION	CELL	REFRIG- ERATION	DOCUMENT, IOM, etc.	REMARKS	ISIO
BAT No.		CPR No.	IN/OUT	IN/OUT	DATE		N
4001	7903	304L CRES 2 DIAPHRAGM	4 28 Jan 80	14 APR 82 13 MAY 82	344AT-82- 204		
BA 001		/	14-Apr 82- 207	30	9 Sept 82		
4002	7924	304L CRES 2 DIAPHRAGM	4 28 Jan 80	14 APR 82 11 MAY 82	344AT-82-204		
BA 002	-	1	8.01	28	9 58 81 - 00-		
4003	7929	304L CRES 2 DIAPHRAGM	28 JAN 80	14 APR 82	344AT-82-204	Hydr	DATE)
BA 003		1	11-24-1892- Cari	1C MHY 02 29	1 4 sept. 02	azin	<u>v 19</u> 7
4004	7947	3041 CRES 29	28 Jan 80	14 APR 82	344AT-84-204		
RAAAA		DIAPHRAGM	H-Desc Ele	14 MAY 82 31	9 Sep + 82		NEPORT
ANNE	7921	3041 CRES 24	28 Jan 80	3 MAR 82	344AT-82-117	60°C	500 1 M.
1005		DIAPHRAGM	03 Mar 82	30 MAR 82.	13 May 82	lant	ater L Pi
BA005		1	765	28		Gran Gran Gran Gran Gran Gran Gran Gran	copc
4006	7902	304 L CRES 29 DIAPHRAGM	28 Jan 80 03 Mar 82	3 MAR 82 31 MAR 82	344AT-82-117	60°C	Comp osal
BA006			765	29	13 may or		atibi 90-96
4007	7932	304L CRES 24 DIAPHRAGM	38 Jan 80 03 Mar 82	3 MAR 82. 18 MAR 82.	344AT-82-117	60 °C	líty 5 rei
BA 007		1	765	16	15 May 200	X X	2
4008	7936	304L CRES 29 DIAPHRAGM	28 Jan 80 03 Mar 82	3 MAR 82	344AT-82-117	60°C	
BADOS		1	765	15	13 May 82		
	************	*Specificati	on: Bell Aer	ospace Texti	ron; Report No.	8803-947047, rev. A	

в-3

NUM	IBER	TEST	TEST I	URATION	ANALYSIS			
SPECIMEN	CAPSULE	MATERIAL DESCRIPTION BAT	CELL	REFRIG- ERATION	DOCUMENT, IOM, etc.	REMARKS	m	
BAT No.	ļ	CPR No.	IN/OUT	IN/OUT	DATE			
4009	7937	304 L CRES 24 LINER	28 Jan 80	14 APR 82	344AT-82-204			
00000			14-and 82	21	8 Sept. 82			
BAOTT		2	- SON	14 000 82	- WAT 07 704		-	
4010	7955	LINER	20000000	4 MAY 82	34441-82-207 8 Sept 82			
0			7 - Day Ch	21				
BAOIS	7010	2 ZARICOGE 24	28 Jan FO	11 400 02	24447-82-204	60°C		
4011	1712	LINER		10 MAY 82	8 Sept 82	42.0 00 N2H2	ydra	
R. M.			14-and 82	27			ızin	
BA 016	7020	2 2011 0000 20	28 Tan 80	2 000 82	34445-87-117	60°0	-le	
4012	1730	LINER	03 mar 82	26 MAR 82	13 May 82		Mon	
BADIT		.2.	765	24			opro	
4013	79202	347 CRES 24	18 MAR 80	1 APR \$1	744-AT-81-061	Special sapsule 1.25" de	Le L	191
1010		DISC	1 11/2 81	7 192 81	22/14/81		lant	LI
BA 023		3	5/8 (241)	6 DAYS			G	rop
4014	79204	347 CRES 24	18 Mar 80	1AP2 81	344-47-81-061	special Rapsule 1.25" dis	ade	osa
		DISC	- oc - tis	7,400.81	2242881		7	1 90
BA OJE		3	218 0405	GDAYS				96-(
4015	79205	347 CRES 24	18 Mar 80	129951	344-47-81-04	Special copsule 1.25" dia	P	5
		DISC	378645	1.4442.81	224/02 81	60°C		ev.
BA 025		3		SIGNIS				2
4016	79206	347 CRES 24	18 Mar 80	1 400-21	344-147-81-56	Special capsule 1, 25" dia	-	
		DISC	APIC ST	LAGE CI	22/18/2 81	00 °C	ľ	
BA 026		3		3.24.12				
	•	*Specificatio	n: Bell Aer	ospace Texti	on: Report No.	8803-947047, rev. A	-	

NUM	BER	TEST MATEDIAT MONTHS	TEST D	URATION	ANALYSIS		TITL	REV	PRE
SPECIMEN	CAPSULE	DESCRIPTION	CELL	REFRIG- ERATION	DOCUMENT, IOM, etc.	REMARKS	1 "	ISIO	70
BAT No.		CPR No.	IN/OUT	IN/OUT	DATE		_		77
4017	7926	3046 CRES 24	28 Jan 80	12 MAY 82	344AT-82-205				
10//		PLUG	12 12 12 12 2	25 MAY 82	8 Sept 82				
BA 034		4	835	14					
4018	7933	3041 CRES 24	28 Jan 80	12 MAY 82	344AT-82-205			1 1	
/		PLUG	12 MAY82	25 MAY 82	8 Sept 82				
BA 035		4	835 Jay	14	•				
4019	7925	3041 CRES 24	28 Jan 80	12 MAY 82	344AT-82.205	60°C	Hy		ر م
,		PLUG	12MA182	30 JUNE 82	8 Sept 82		lra		Ē
8			6.21	50			211	1.4	11
DA 036		4	835		2 11 15 0 2 2 05		_je	1	79
4010	7935	3046 CRES 24	28 Jan 80	12 MAY 82	344AT-82-205	60°C	M		
		PLUG	12 44 84	30 JUNE 82	9 seption		no	l g	9X
BA 037		4	835	50			oro	ECT	75
1091	TOAN	3001 CPES 24	28 Jan 80	12 MAY 82	344AT-82-205		Pel	4 3	8 §
4021	1740	WELD TIG	12 MAY82	27 14-182	8 Sept 82		lar	PL	
	• •			1.5			Ħ	Pr	1
BA OQS		5	835	and the second			_ £	opo	
6012	7944	3041 CRES 24	28 Jan 80	12 MAY 82	344A7-82-205		de	Sal	
/ •		WELD TIG	1214482	27 MAY 82	8 Sept 82		1	l 9	
BA 046		5	835	16				t1191	
4023	7911	3041 CRES 24	28 Jan 80	12 MAY 82	344AT-82-205	60°C		5 1	
1000	/ ///	WELD TIG	12012482	29 JUNE 82	8 Sept 82			rev	
BA NAT		5	825	49				. 2	
ANTA	7978	304LCEES 24	28 Jan 80	12 MAY 82	344AT-82-205	60°C	1		
4024	1120	WELD TIG	12 MAY 82	1.101 1 82.	8 Sept 82				
				51					
BA 048	<u> </u>	5	l						
		*Specificatio	n: Bell Aer	ospace Texti	on; Report No.	8803-947047, rev. A	1		

NUMI SPECIMEN	BER CAPSULE	TEST MATERIAL MONTH DESCRIPTION	CELL	REFRIG-	ANALYSIS DOCUMENT, IOM. etc.	- 	REMARKS		TITLE	REVISI	IPREPARI
RAT Na		BAT	TN /OUT	TN/OUT	DATE					N N	
4125	7971	3046 CRES 24	4 Mar 80	14 APP 82	344AT-82-204				1		23
TUZJ		WELD EB # 40 7	Hayse 85	II MAY 82	8 Sept 82						
BA 056		6	271	28							
4026	7960	3046 CRES 24	4 Mar 80	14 APR 82	344AT-82-204						ŀ
/ 000		WELD EB #407	H-apr 82	12 MAY 82	8 Sept 82						
BA 057		6	771	61					-		
4027	7922	WELD EB	4 Mar 80 03 Mar 82	3 MAR 82 26 MAR 82	344AT-82-117 8 Sept 82	60°C			lydra	ATE)	July
BA 058		# \$ 8 7 6	129	24	1				zine		1979
4028	7950	3042 CRES 24	18 Mar 80	3 MAR 82	344AT-82-17	60°C			M		
1000	,,	WELD, EB	O3 Mar 82	30 MAIN BA	13 May 82				ono	Į Į	79X
RA 0.59		# 407	115	27.					pro	ECT)	075
1120	70A1	304L CRES 29	4 Mar 80	14 APR 82	344A7-82-204				Tel .	53	8 6
FUAI	/ / 7 0	WELD EB	11-1-0382	14 MAY 82	9 Sept 82				lan	L	
BA 067	4. 	# fo 6 7	771	31			-		t Gr	rial Prop	
1170	7965	304LCRES 24	\$ Mar 80	14 APR 82	344AT-82-204				ade	Sa Co	
7030	//00	WELD EB	14 0 50 8 20	12 MAY82	8 Sept 82				*	I 9	÷.,
RA ALS		7	1 721	. 29						0-9	
1021	79.58	3041 CRES 24	18 Mar 80	14 APR 82	344AT. 82-204	6000		· · ·	1	111	ļ
7051	1100	WELD EB	110 . 1.2	10 MAY82	8 Sept 82					rev	1
BA ALA		#406	745.00	27						2	
1029	7930	3046 CRES 24	18 May 80	3 MAR 82	344AT-82-117	60°C			1 .		
TUJA	1757	WELD EB	03 dias 22	25 MAR 82	13 May 42						
BA 070		# 406 7	nik.	23							
		*		· · · · · · · · · · · · · · · · · · ·	4						1

NUM	BER	TEST	TEST D	URATION	ANALYSIS			E	
SPECIMEN	CAPSULE	MATERIAL MONTHS DESCRIPTION	CELL	REFRIG- ERATION	DOCUMENT, IOM, etc.	REMARKS		VISIO	2
BAT No.		CPR No.	IN/OUT	IN/OUT	DATE			Z	N a
4033	7969	3081/3091 CRES 24 RING-DIAPH	4 Mar 80 12 MAY 82	12 MAY 82 26 MAY 82	344AT-82-205 8 Sect 82				R.
BA 1178		general de la companya de la compa	799	15					
4034	7978	3081/3041 CRES 24 RING-DIAPH.	4 Mar 80	12 MAY 82	344AT-82-205				
BA 079	1. 1. 1. 1. 1. 1. 1. 1. 1. 1. 1. 1. 1. 1. 1. 1. 1	8	799	26 MAY 8C 15	8 Sept 82				
4035	7966	3081/3041 CRES 29 RING-DIAPH.	18 Mar 80	12 MAY 82 10 JUNE 82	344AT-82-205 8 Sep + 82	60°C	Hydraz	(DATE)	July
BA 080		8	785	30			ine		197
4036	7970	3081/3041 CRES 24 RING-DIAPH.	18 Man 80	12 MAY 82 29 JUNE 82	344AT-82-205 8 Sept 82	60°C	, Monc	(PRO	9 791
BA 081		8	785	49			prop	JECT)	(0750
4037	7914	304L CRES 14 LINER	28 Jan 80 14-19-19-82	14 APR 82 13 MAY 82	344AT-82-204 8 Sept 82	Replace 84 085	ellant	Mater JPL I	ðġ
BA 300		9	807	30			្ន	rop	
4038	7945	309L CRES 29 LINER	28 Jan 80 03 Mar 82	3 MAR 82. 25 MAR 82.	344A7-82-117 13 May 82	Replace BA 086	ade*	Compa osal 9	
BA 301		9	165	23				0-9	
4039	7954	304L CRES 24 LINER	28 Jan 80	14 APR 82. 5 MAY 82	344AT-82-204 8 Sept 82	Replaces BA 087		ility 65 rev.	
BA 302		9	\$ \$27	22				N	
4040	7956	304LCRES 24 LINER	28 Jan 80	14 APR 82 5 MAY 82	344AT-82-204 8 Sept.92	Replaces BA 088			
BA 303		9	<u> </u>	22					
		*Specification	on: Bell Aer	ospace Textr	on; Report No.	8803-947047, rev. A			1.11

В-7

NUM	BER	Test	TEST D	URATION	ANATYCTC	T		
SPECIMEN	CAPSULE	MATERIAL MONTHS DESCRIPTION	CELL	REFRIG- ERATION	DOCUMENT, IOM, etc.	REMARKS	I'LE	EVISIO
BAT No.		CPR No.	IN/OUT	IN/OUT	DATE			NUN
4041	7918	304L CRES 24 LINER	28 Jan 80 03 Mar 82	3 MAR 82 30 MAR 82	344AT-82-117 13 May 82	Replaces BA 089 60°C		
BA 304		9	765	27				
4042	7927	304L CRES 24 LINER	28 Jan 80 03 Mar 82	3 MAR 82 19 MAR 82	344AT-82-117 13 May 82	Replaces BA 090 60°C		
BA 305		9	765	17				
4043	7943	304L CRES 24 LINER	28 Jan 80 03 Mar 82	3 MAR 82 19 MAR 82	344AT-82-117 13 May 82	Replaces BA 091 60°C	Hydraz	(DATE) July (DATE)
BA 306		9	765	17			ine	197
4044	7930	304L CRES 24 LINER	285an80 03mar 82	3 MAR 82 17 MAR 82	344AT-82-117	Replaces 84 092 60°C	, Mone	9 793 (PRO
84 307		9	105	15	13 may 20		opro	VORT N
4045	7982	A286 CRES 6 TANK SHELL	4 Mar FO 4 SEPT 80	4 SEPT SO 15 OCT 80	344 AT- 80-160 7 NOV 80		bellant	yo.) 00 Matej JPL I
BA 100		10	184 Dauga	HIDAYS			Gr	rial rop
4046	7920	A 286 CRES 6 TANK SHELL	4 Mar 80 4 Nov 80	4 NOU 80 19 NOU 80	344 AT - 20-173 505 - 80		ade*	Compa osal 9
BA IDI		10	243 0413	15 DAYS				t1b
4047	7957	A286 CRES 6 TANK SHELL	4Mar 80 6 JAN 81	6 JAN 81 20 JAN 81	344AT-81-019 13 FEB 81			111ty 55 rev
BA 102		10	308 2445	14 2440				2
4048	7959	A186 CRES 6 TANK SHELL 6	4 Mar 80 4 MAR 81	4 MAR 81 1 Mary 81	344AT-81-086 5 MAY 81			
Dillo				100			100	

в-8

NUM	BER	TE	ST	TEST I	DURATION	ANALYSIS				KE	
SPECIMEN	CAPSULE	DESCRIPTION	14/145	CELL	REFRIG- ERATION	DOCUMENT, IOM, etc.	REMARI	KS		1ST0	Z
BAT No.		CPR No.		IN/OUT	IN/OUT	DATE					H
4049	7963	A286 CRES WELD TIG # BII	6	4 Mar 80 4 NOU 80	4 NOU 80	344-AT-20-173 SDEC 80					H.
BA 109		11.		245 Days	16 DAYS						
4050	7981	A 286 CRES WELD TIG	6	4 Mar 80 4 Mar 81	4 MAR81 23 APR 81	344AT-81-086 15 MAY 81					
BA 110		# 4/1		365 04-5	50						
4051	7974	A 286 CRES WELD TIG # 417	6	4 Mar 80 4 MOU 80	4 NOV 80 20 NOV 80	344-AT- 80-173 5 DEC 80			Hydraz	(DATE)	July
BA 112		12		245 8000	16 DAYS				Ine		197
4052	7977	A286 CRES WELD TIG	6	4 Mar 80 4 Mar 81	4 mon 81 22 April	344 AT-81-086 15 MA-1 81			, Mono	(PRO	X64 6
BA 113		# 417		3652000	49				prop	ECT)	0750
4053	7934	308L CRES FILLER WIRE	6	4 Mar 80 4 NOU 80	4 NON 80 14 NON 80	344-AT-90-173 SDEC 80			oel lant	Mater JPL 1	ŏ
BA 115		13		245 Days	15 DAYS				Gr	rop	
4054	7951	3081 CRES FILLER WIRE	6	4 May 80 4 May 81	4 mar 81 24 Ago 81	3414-19-81-086 15 may 51			ade*	Compa osal 9	and the subscription
BA 116		13		365Darg	51		· · · · · · · · · · · · · · · · · · ·			0-9	
4055	7915	308L/304L CRES LINER-DIAPH,	6	28 Jan 80 4 SEPT 80	45297 80 1705780	344 AT-80-160 71100 80	42 CC N2 H4			111ty 55 rev	
BA 118		14		220 Days	43 DAYS					. 2	-
4056	7913	3081/304L CRES LINER- DIAPH,	6	28 Jan 80 4 NOU 80	4 NON 80 18 NON 80	3,44, 67 - 40 -M	42 cc N2H&				
RA 110		14		2x1 days	NUDAIS						

NUM	BER	TEST	TEST I	URATION	ANALYSIS			KE	Ţ
SPECIMEN	CAPSULE	MATERIAL POWINS DESCRIPTION	CELL	REFRIG- ERATION	DOCUMENT, IOM, etc.	REMARKS	m	IOTST/	
BAT No.		CPR No.	IN/OUT	IN/OUT	DATE				F
4057	7941	308L/304L CRES 6	28JAn 80	6 JAN 81	344-17-81-99				P
,		LINER- UMPA	6 JAN 8/	23 244 01	13 FEB 81				
BA 120		14	344 DAVS	IDDAYS					
4058	7952	3081/3041 CRES 6	28 Jan 80	4 Aren 81	344AT-81-086		1		
1050	////	LINER-DIAPH.	4 Mar 8	21Aprol	15 MAY 81				
BA 121		14	401 days	48					
6159	701 A	MYLAR FILM 6	4 Mar 80	4 mar 81	344AT-81-086		Hy		Ŀ
1001	1764	TYPE G DISC	4 mars	25Mar 81	1511481		dras	Ē	<u>ATn</u>
BA 127		15	365 Dausa	2			51ne		121
ANLO	7010	MYLAR FILM /	4 May 80	4 Ameril	344AT-81-086	•			ľ
7000	1168	TYPEG	4 mar 81	30Apr 81	15 MAY81		lonc	PRO	19
RA 128		DISC	365 Sam	57			pro		
10/1	7079	304L CRES 6	4 Mar 80	6 24481	344-41-81-09		Pel	L 3	Ę
7061	1.112	WELD EB	6 5 1 81	21 34481	13 FEB 81		lan	PL	
80.00		# 404	308 DHYS				t G	r1a Pro	
DA 130	7000	16 3011 CRES 1	70 Mar. Ca	UMU 81	244-17-81-081	a capsulo 7979 broken.	rad	1 C	
4061	1770	WELD EB	4 Mar 81	21 April	1544481	& copsule 7980 broken	10° *	日日	
		# 404		L.K.		c. specimen recleaned a, &	F.	90-	
BA 131		16 SAMADIUMA	Subday	1 14181	244-47-81-019		-	965	
40.63	7975	COBALT 6	6 JAN 81	21 JAN 8	17 FPR SI			re	
		MAGNET	308 0445		131.0001			۷.	
BA 133		17		1512445			4	2	
4064	7976	SAMARIUM 6 COBALT	4 Mar 80	29 Am x	344AT-81-086				
		MAGNET	-7 marti		IS MAY 81	1			
BA 134		17	3650 am	1.0				1 .	

NUM	BER	MATERIAL MONTHS	TEST D	URATION	ANALYSIS	DEMADUC		REVI	PREP
PECIMEN	CAPSULE	DESCRIPTION	CELL	REFRIG- ERATION	IOM, etc.	KEMARKS	1	SIO	HAR
BAT No.		CPE No.	IN/OUT	IN/OUT	DATE			Z	
4065	7948	17-4 PH CRES 6	4 Mar 80	6 JAN 81 22 JAN 81	344/17-581-019]		R.
		ELECTRO POLISH	6 JAN 81 228 DOVC		13 7 1 1 5 81				
SE 200		18		17 0445	and the set	·	4		
4066	7973	HINSN	4 Mar 80	Garen 81	544151-81-086				
		ELECTRO POLISH	7 one of	1 May of	130-001-01				
SE 201		18	365 dauge	50				ليسا	
4067	7961	17-4 PH CRES 6	4 Mar 80	ymon 81	34447-81-086		Hyd	DAT	J
		ELECTRO POLISH	4 mars	20Mar 81	15 14 181		ras		ilv
5E 110	-	CHROME PLATE	2150	1 2 2			in		19
AALA	7017	17-4 PH CRES 1	& Mar 80	4 SEPT80	344-17-80-160		-t [®]		79
7060	1761	H1050	4 SEPT 80	1600 80	7 100,80		Mon	(PP)	(RE
		ELECTRO POLISH	184 Daux				opr	DIEC	X07
SE 211				420445		Charles Consult on let A.8-75 dia	- Pa	3	500
4069	7986	17-7PH CRED 6	4 Mar 50	4 SEPT 80	7 14 . 41 . 201 40	special report of 15 end	11	JPI	- 2
		SPRING	4 500 00	1,	11(00 80		1 T	PH	
SE 220		20	189 acrys	43 DAYS			ទ្ឋ	cop	
1 À 70	7999	17- TPH CRES 6	4 Mar 80	4 Anan 81	344AT-81-086	special copsule meck 0.825 dia	ade	Co Co	
7070	1112	CH 910	4 mar 81	20APA 81	12 WAY EI		*	19 19	
SE 231		SPRING	2450000	1-47					
1091	7010	17-4 PH/17-4 PH CRES 6	4 Mar 80	6 JAN 81	344 1.81-019		1	65 11	
40.11	1762	WELD TIG	6 JAN 81	2224USI	3FEB81		1	rev	
s e 230		21	308 DAXS	16DAYS				. 2	
1171	7001	17-4 PH/17-4PH CRES6	4 Mar 80	yman 81	344 AT-81-086		1		
TUIL	//7/	WELD TIG	y dren sol	1 May 81	15 13 181				
SE 231		21	365 Dauge	S SO				1	
	. 	*Specificatio	n. Boll Aer	ognaca Tavti	on: Report No.	8803-947047, rev. A			

LASS	NUM SPECTMEN	BER CAPSULE	MATERIAL MONTHS	TEST I	BEFRIC-	ANALYSIS DOCUMENT,	REMARKS	ITLE	LEVI:
FICA	or bornait		DESCRIPTION		ERATION	IOM, etc.			501S
TIO	BAT No.		CPR No.	IN/OUT	IN/OUT	DATE			N
	4073	7993	INCONEL 902 6	4 May 80	CJANSI	54447-81-019	Special Repairle mack 0,856 dia		1
			TUBE	CJAN 81	12 MACK	1 3 PC13 01			
	56240		22	308 DAVS	10 DAYS				
	4074	7994	INCONEL 902 6	4 Mar 80	4 mar 81	344 15-81-086	special capsule neck 0.856 dia		
	/ • / /		NI SPANC TURE	4 Aren 81	26 Mar 81	15 MAR 81			
	SC 241		29	3650m	22				
	4075	7984	347 CRES 6	4 Mar 80	440080	344 AT. 80-173		Hy	J
	1010		V64	4 No 80	20 NON SU	SIDECXO		iras	uly E)
	PA 950		22	24sdays	LIG DAYS			Ine	19
ŀ	A N71	7002	347 CRES 6	9 Mm 80	4 man 81	34447-81-086			979
	4010	1105	J64	your st	2020-81	15MAY 81		lone	79) (PRC
				2151	47			opro	X07
	BA 251	2000	23 3A7 CPCC /	1 Mr. 80	11 1 1 1 1 1 1 1 1	344 15.50.173		pe	500
. 4	4077	1995	ANNEALED 6	4 NOV 80	YNOU XO	SDEC 80		11a1	fat.
			TUBE	245 Daux				Ĩ	Pro
-	BA 260		24	A 44 80	14 0445	4		irac	al (
4	4078	7996	347 CRES 6	4 man 81	yren 81	344 M-81-080		le*	ia1
			TUBE		LANG OF				90-
Ļ	BA 261	ļ	24	365Day				4	-96.
	4079	7987	397/347 6	4 Mar 80	19 NOU 80	1344 AT 80.173			5 14 13
_			WELD ASTROARC					1	/ 2V.
r c	BA 270		25	2012 grings	15DAYS	-		1	N
-989	4080	7989	347/347 6	4 Mar 80	4mar 81	344AT-81-086			
í R	,		WELD ASTRO ARC	1 y mar 81	22/4128	1.5174181			
EV 1	BA 271		# 78	365 Daug					
1			*Specificatio	n: Bell Aer	ospace Text	ron; Report No.	8803-947047, rev. A		
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NUMI	BER	TEST MATERIAL MONTHS	TEST D	URATION	ANALYSIS	DEMADIZO		S	PREP
SPECIMEN	CAPSULE	DESCRIPTION BAT	CELL	REFRIG- ERATION	IOM, etc.	KEMAKAS) ISION	ARED
BAT No.		CPR No.	IN/OUT	IN/OUT	DATE				2,≧
4081	7988	EPR 515 6 SEAL PAPKEP	4 MAN 80 IN ADD 80	10 APR 80	344AT-80-076	Terminate - TPI permet 90: 96 F- 0			H H
SE 280		KPYTOX 240AC	37 days	6 0445	111091100	(A)			
4082	799N	EPR 515 6	4 Mar 80	8 MAY 80	344 AT-80-091	Terminate -	-1		
7000	1110	SEAL PARKER	8 May 80	13 MAY 80	28 MAY 1980	JPL report 90:965-9			
SE 281		KKYTOX 240AC 26	65 days	5 DAYS	l	A			
			-				Hydı	DATE	DATE
							razi		ייים 1
							ne,		979
							Mon	Î.	7 (19
							opr	OJEC	POR
	L						pe]		50 NO
							lan	late IPL	-
							tG	ria] Proj	
							cade	L Co	
							*	mpa: 1 90	
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								5 r	
								ev.	
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-	CAPSIII	MATERIAL Mon	ST NTHS	TEST I	DURATION	ANALYSIS	REMARKS			NAME I
ES	T CAPSUL	DESCRIPTION BAT		CELL	ERATION	IOM, etc.			NOT C	STON
100	<u>.</u> 7949	N2 Hq only	24 2	IN/OUT 28 Jan 80 4 SEPT 80	1N/OUT 4513 PT 80 13 OCT 80	7400 80	control			
		40 xx	3	120 days	39 0445					
10,	7908	N2H2 only	14	4 Mar 80 -1 mar 81	4 mar 81 30 Apr 81	344AT-81-086 15MAY 81	control			
. •		40 m	3	65day						-
10	? 7916	N2H4 only	24	4 Mar 80 45ept 81	4 SEP 81 8 FLB 82	244AT 82-117 13 May 82	control	Hydrazi		DATE)
		40 KR		549		5		ne,		
-10	3 7904	NaH& Only.	24	28 JAN 80 4 SEPT 80 120 Dans	4 SEPT 80 14 OCT 80	JALAL ROLLEO	Control 60°C	Monop		(PROJE
		40 LL			40 DAYS					3
10	4 7931	N2 Hq only	24	28 Jan 80 4 Mar 81	4 man 81 25 Man 81	34417-81-081 15 MAN 81	60°C	ellant	JPL Pi	
		40 xx	4	101 days				Gra	ropo	5
10	5 7942	N2 H& only	24	28 Jan 80 28 July 81	28 JUL 81 10 FEB 82	344AT-82-117 13 May 82	controf 60°C	lde*	Compationsal 90	n
		40 KK		541	190				-96	
10	6 7985	Nr. H& only	24	4 Mar 80 3 Mar 82	3 MAR 82 31 MAR 82	344AT-82-117 13 May 82	Control		5 rev.	
		40 xx		11.7	27				N	
10	7 7910	N2H& only	24	4Mar 80 3 mar 82	3 MAR 82 24 MAR 82	344AT-82-117 13 May 82	Control 60°C			
		40 KK	<u> </u>	71 a	22		<u> </u>			
		*Specificat	tion:	Bell Aer	ospace Text	ron; Report No.	8803-947047, rev. A			_

NUM	BER	TES	7	TEST D	URATION	ANALYSIS			RE	PA
TEST	CAPSULE	MATERIAL 770, DESCRIPTION	MTH	CELL	REFRIG- ERATION	DOCUMENT, IOM, etc.	REMARKS	m	VISIO	TO
BAT NO.	·	CPR No.		IN/OUT	IN/OUT	DATE				YEY
4108	7919	N2H& only	24	4 MAR80 03 mar 82		344AT-82-117	control			
		40.cx		729		157 00.9				
4109	7913	N2H4 only	24	& MARSO	3 MAR 82 24 MAR 82	344AT-82-117 13 May 82	control			
		40 xc		729	22		60 C			
4110	7907	Empty.	24	10 Mar 80			open Junnel	Hydrazir	DATE)	DATE) July 19
4111	7917	Empty	2¢	10 Mar 80			calibration unit open funnel 60°C	ie, Monopro	(PROJECT	(REPORT
41/2	7905	Sealed Internal presson 733 mm hg	24 ne	18 Mar 80			Reference unit	pellant Gr	^{r)} Material JPL Prop	NO.)
4113	7997	Sealed Internal presson 733 mm hg	24 L	18 Mar 80			Reference unit 60°C	ade*	Compatib posal 90-9	
4114	NONE No gauge	N2 H4 only 40 cc	-	28 Feb 80 8 Opr 80 40 days	28 Feb 80 8 Apr 80 80 days	394AT-80-081 May 1980	Special for CO2 analysis check Terminated		ility 65 rev. 2	
4115	79201	Empty	24	10 Mar 80	- F		calibration unit open funnel			

LASSIFICA	NUM FST	BER CAPSULE	MATERIAL DESCRIPTION	TEST D	URATION REFRIG- ERATION	ANALYSIS DOCUMENT, IOM, etc.	REMARKS	TITLE	REVISIO	To
TION BA	<u>т No.</u> 116	79203	Empty It	IN/OUT 18 Mar, Fi	IN/OUT	DATE	calibration unit open funnel			TH
							60°C	-		
								Hydrazine,	DATE)	July 1979
								Monoprop	(PROJECT)	79X0750
								ellant Gra	Material JPL Propo	0
	·				-			ade*	Compatibs	
Jaf									11ty 55 rev. 2	
999-5 (REV										
11-68)		L	1*Specificatio	Don: Bell Aer	l ospace Textr	on; Report No.	8803-947047, rev. A	1		

в-16

Report Number 79×07501 Project N2 H& COMPATIBILITY Classification <u>UNCL</u>

JPL 0988 MAR Et

JET PROPULSION LABORATORY CALIFORNIA INSTITUTE OF TECHNOLOGY

SUMMARY

TEST CAPSULE SPECIMEN NUMBERS DETAIL INFORMATION

Prepared by J. TOTH Dote July 1979 C. MORAN Date July 1999

	1. A.	AP	PENDIX OR REVISION	
Date	Pages Affected	Appendix OR Revision	Remarks	Changed by
		-		

Classification

NU	MBER		TEST D	URATION	ANALYSIS	S	TOP HOAP	1
SPECIMEN	CAPSULE	MATERIAL DESCRIPTION	CELL	REFRIG- ERATION	DOCUMENT, IOM, etc.	REMARKS	1401	m
BA	Vol. Cm3	Vol. cm3	IN/OUT	IN/OUT	DATE	$N_2H \neq V_{0L} = 40.$	0-Speck	فبهرة
NONE	7901	Capsule -	broken		f. of	Broken	5.10	
		NO Capsu.	20 To Pa	all and	miller	184.0	6.80	
4006	7902		20 Jan 80 8:30 A M	12 N		60°C.	6.00	
006	88.8	0.2311		21°C		39.7689		-
4001	7903		28 Jan 80 8 ; 30 A M	21 Jan 80 3 PM			5,40 Hydraz	
001	83.9	0.2320		21°C		39.768	ine	•
4103	7904	NeHq 40 cc	28 Jan 80 8:30 A M 4 SEQT 80	18 Jan 80 1 PM	Evacuated to Imm may	60°C Contro P	3,10 Mono	
NONE	86.3	0	220days	22.0	-77° C	40,00	prop	
4/12	7905	NOTHING PRESSURE: JPL I ATMOSPHERE//	/8MAR 80 8:30 AM	11 Mar 80 10; 30 AM			3, 82 2, 70	JPL P
NONE	84.8	0 733 mm hg		23°C		Blank se	alof 9	rop
NONE	7906	Experimental				Ref no sig no	ading *	osal 90-9
4110	7907	NotHING	10 MAR 80 8:30 AM				4.14 2.90	165 rev
		Open meete		2		Blank cat	ibration	N
4101	7908	N2 H4 40RR	4 MAR 80 8:30 AM	27 Feb-80 9:30 AM		control	6.36	
NONE	85.6	0		2300	24	40.00		

в-18

2]	NUM	BER	<u> </u>	TEST T	TRATION		1		2	1	
ASSIFIC	SPECIMEN	CAPSULE	MATERIAL DESCRIPTION	CELL	SEALED REFRIG-	DOCUMENT, IOM. etc.	REMARKS		EVISI	REPARI	V
ÄTIO	BA	Vol cm?	Val am3	IN/OUT	ERAIION IN/OUT	DATE	NeH& Nof		N N	0 T	
z	NONE	7909	Experimental				Ref. unit for sg			H,	
										30	
,	4107	7910	N2H4 40RC	4 Mar 80 8130 AM	27 Feb 80 9:30 AM		60°C . 5.5 Control 4.80	A 2		PAN	
	NONE	85.8	0	· · ·	23°C		40.00				
¢	4023	7911		28 Jan 80 8:30 AM	225an80 10 A M		60°C 7.13	Hydras	(DATE)	(DATE) July	
	047	87.2	0.2698		20°C		39.7302	ine		197	
7	4011	7912		28 Jan 80 8:30 AM	22 Jan 80 10 AM		60°C 7.54 Total Not. 42.0 cc	, Mono	(PRO	(REP) 9 79X	
	016	87.9	3.3606		20°C		36,6394	prof	JECT)	075(
/	4109	7913	N2H4 40KK	4 Mar 80 8:30 AM	27Fet80 9;30 AM		60°C 9.0 Control 8.0	ellant	Mater JPL F	ē.	
	NONE	86.0	0		23°C		40.00	Gr	rop		
7	4037	7914		28 Jan 80 8:30 AM	18 Jan 80 1 PM		4.11	ade*	Compa osal 9		2
	300	87.2	0.6657		2200		39.3343		0-9		
4	4055	7915		28 Jan 80 8:30 AM	18 Jan 80 11 A M		4.12 Total.vol. 42.0 cc	2	llfty 55 rev		PAGE .
	118	87.1	0.7679		2100		39.2321		2		R
	4102	7916	N2H4 40KK	4 Mar 80 8:30 AM	27 Feb 80 10 AM		Control 8.90				0F
NEV 11	NONE	84.8	0		2300		40.00	j			
			*Specificatio	n: Bell Aer	ospace Textr	on; Report Nó.	8803-947047, rev. A				Ĺ

B-20		BA 4111 4041 304 4108 NoNE 4046	Vol. cm ³ 7917 7918 87.0 7919 84.8	Volam ³ Nothing Open mech 0.6492 N2 H4 40 cc 0	IN/OUT 10 MAR 80 8:30 AM 28 Jan 80 8:30 AM 4 Mar 80 8;30 AM	-IN/0000 18 Jan 80 1 PM 22°C 27 Feb-80	DATE	N2H4 Nop 60°C Blank 60°C 39.3508	8.36 7.50 4.91 8.2		TH/MORAN
B-20	3	4111 4041 304 4108 None 4046	7917 7918 87.0 7919 84.8	NotHING Open mech 0.6492 N2 H4 40 xx 0	10 MAR 80 8: 30 AM 28 Jan 80 8: 30 AM 4 Mar 80 8; 30 AM	18 Jan 80 1 PM 22°C 27 Feb-80		60°C Blank 60°C 39.3508	8.36 7.50 4.91 8.2		MORAN
B-20	3	4041 304 4108 None 4046	7918 87.0 7919 84.8	Open mech 0.6492 N2 H4 40 xx 0	28 Jan 80 8: 30 A M 4 Mar 80 8; 30 AM	18 Jan 80 1 PM 22°C 27 Feb-80		Blank 60°C: 39.3508	4.91 8.2		YORAN
B-20	3	4041 <u>304</u> 4108 <u>None</u> 4046	7918 87.0 7919 84.8	0.649 <u>2</u> N ₂ H4 40 xx	28 Jan 80 8: 30 A M 4 Mar 80 8; 30 AM	18 Jan 80 1 PM 22°C 27 Feb-80		60°C· 39.3508	4.91 8.2		OAN
B-20	1	304 4108 None 4046	87.0 7919 84.8	0,6492 N2 H4 40 xx 0	4 Mar 80 8; 30 AM	22°C 27Feb-80		39.3508	8.2	_	
₽-20	1	4108 _{None} 4046	7919 84.8	N2 H4 40 RR	4 Mar 80 8; 30 AM	27 Feb-80		A	8.2		~ ~
R-20	5	None 4046	84.8	0	1	10 111		Control	-	Hydraz	July DATE)
B-20	5	4046				2300		40.00		ine	1979
- <u>></u> 0	11		7920		4 Mar 80 8:30 AM	27 Fet 80 10 30 AM			5.3	Mono	(PROJ
1 C C C C C C C C C C C C C C C C C C C	L	101	86.8	1.0618		23°C		38.938		prop	0750 ECT)
	2	4005	7921		28 Jan 80 8:30 A M	21 Jan 80 12 N		60°C.	7.0	JPL P ellant	Mater
		005	88.6	0.2367		21°C		39.7633		ropo	
	14	4027	7922		18 Mar 80 9:00 AM	11 Mar80 1:30		60°C	8.10	osal 9 ade*	Compation
		058	85.3	1.5605		2300	29	38.4395		0-96	
	4	4056	7923		28 Jan 80 8:31 Am	21JAm 80 3 PM		Total vol. 42.0	5.98 CC	5 rev.	14tv
	JPLO	119	88.7	0.7494		2.1°C		39.2506		2	
	H)) (1966	4002	7924		28 Jan 80 8:30 ЛМ	21 Jan 80 3 P M			5.35		
	EV 11	002	85.1	0.2371		21°C		39.7629			
	-68)		-	*Specificati	lon: Bell Aer	ospace Textro	on; Report No.	8803-947047, rev. A			

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LASSIFICA	NUM SPECIMEN	BER CAPSULE	MATERIAL DESCRIPTION	TEST I CELL	URATION SEALED REFRIG- ERATION	ANALYSIS DOCUMENT, IOM, etc.	REMARKS		11TLE	REVISIO	
To	BA	Val an?	Volan?	IN/OUT	-IN/OUT-	DATE	NaHqVol			N Y PY	
2 4	4019	7925		28 Jan 80 8:30 AM	21Jan80 11AM		60°C	5.85		H/V	
	036	90.5	0.083		2100		39,917			101	
19	4017	7926		28 Jan 80 8530 AM	2150m80 4PM			5.55		OAN	
	034	90.2	0.084		2100		39.916			-	
3	4042	7927		28 Jan 80 8 30 AM	21 JAN 80 11 A M		60°C	5.1	Hydraz	(date) July (date)	
l	305	87.4	0.6706		2100		39,3294		ine	197	
14	4024	7928		28 Jan 80 8:30 AM	22 Jan 80 10 A M		60°C'	6,95	, Mono	9 79X	
	048	83.1	0.2653		20.0		39. 7347		prop	075(
2	4003	7929		28 Jan 80 8:30 AM	21 Jan 80 3 P M			5.9	ellant	Mater	
	003	84.2	0.2345		2100		39.7655		Gr	ial	
3	4044	7930		28 Jan 80 8: 70 AM	21 JAN 80 3 PM	-	60°C.	9.25	ade*	Compa	A
	307	90.2	0.6360		2100		39.364			o tib	
1	4104	7931	N2H4 40RR	28 Jan 80 8:30 AM	18 Jan 80 1 PM		60°C.	9.5		lifty	PAGE.
Jac	NONE	88.0	0		2200		40.00			з	A
1) 🛉 🖓 60	4007	7932		28 Jan 80 8:30 H 14	21 Jay 80 2 PM		60°C.	6.75			۱, ا
REV 11	007	86.2	0.2324		2100		39.7676				
-68)			*Specificatio	on: Bell Aer	ospace Textro	on; Report No.	8803-947047, rev. A				I

ASSIF	SPECIMEN	CAPSULE	MATERIAL	CELL	SEALED REFRIG-	ANALYSIS DOCUMENT,	REMARK	5	TLE	EVIS
CATIC	BA	Vol an?	Volam 3	TN/OUT	ERATION	DATE	NaHaval			ION
ž 14	4018	79 33		2850m 80 8.30 AM	21 Jan 80 4 PM			5.50		
	035	91.9	0.086		2100		39.914			100
9	4053	7934		4 Mar 80 8:30 AM	27Fet.80 11 AM			5.90		MCI.
	115	86.1	0.2815		23°C		39.7185			
14	4020	7935		28 Jan 80 8:30 AM	21 Jan 80 4 PM		60°C.	5.85	Hydraz	DATE)
	037	87.6	0.087		2100		39.913		sine	161
2	4008	7936		28 JAM 80 8:30 117	21 Jan 80 2 PM		60°C	6.75	, Mono	(PROJ
	008	85.4	0.2301		2100		39.7699		prop	ECT)
14	4009	7937		28 Jan 80 8:30 AM	21 Jan 80 11 AM			6.0	ellant	Mater JPL P
	014	86.2	2,2708		2100		37. 7292		Gre	fal
19	4012	7938		28 Jan 80 8; 30 Ary	21 Jan 80 11 AM		60°C	7.9	ide*	Compations Compations (Compation) Compation (Compation) Compation)
	017	88.3	2.5323		2100	······	37.4677			0-96-0
14	4032	7939		18 Mar 80 9:00 AM	11 Mar 80 1: 30 PM		60°C.	7.85		lity 5 rev.
2	.070	86.2	1,4709		23°C	24	38,5291			2
H 10000	402.1	7940		28 Jan 80 8:30 AM	29 JAN 80 10 AM			5.65		
EV 11	045	88.3	0.2620		20°C		39.738			
-68)			*Specificat	tion: Bell Aer	ospace Textro	on; Report No.	8803-947047, rev.	A		

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LASSIFIC.	NUM	BER CAPSULE	MATERIAL DESCRIPTION	CELL	URATION SEALED REFRIG- ERATION	ANALYSIS DOCUMENT, IOM, etc.	REMARKS		ITLE	PREPAREI	T
ATIO	BA	Vol an3	Val an?	IN/OUT	EN/OUT	DATE	N2HANA	-		NIC BY	
4	4057	79\$1		28 Jan 80 81 30 AM	18 Jan 80 1 PM			3.92		N/H	
	120	83.8	0.7366		22°C		39.2634			10,	
- /	4105	79 42	N2H& 40KK	28 Jan 60 8:30 A M	18 Jan 80 1 P M		60°C. Control	8.08		RAN	
	NONE	88.3	0		2200		40.00				
3	4043	79 43		28 Jan 80 8: 20 AM	21 Jan 80 12 N		60°C	4.64	Hydraz	(DATE) July (DATE)	
	306	88.1	0,6430		2100		39,357		ine	197	
18	4022	7944		28 Jan 80 8:30 AM	22 Jan 80 10 AM			6,68	, Mono	9 79X	
	046	84.2	0,2676		20°C		39.7324		prop	ОРТ N 075(IECT)	
3	4038	7945		28 JAMEO 8:30 AM	18 Jan 80 1 PM			\$.0	ellant	Nater	
	301	82.7	0,6299		22°C		39.3701		Gr	fal	
19	4029	7946		4 Mar 80 8:30 A M	27 Fet 80 2:30 PM			6,40	ade*	Compa	6
·	067	84.7	1.3348		2300	24	38,6652			0-96	
٦	4004	7947		28 JAN 80 8:30 AM	21 JAM80 3 PM			5,88		1 1 1 1 1 1 1 1 1	PAGE
JPLO	004	86.8	0.2363		2100		39.7637			5	6
PL 2000	4065	79 48		4 Mar 80 8:30 AM	27 Fel 80 1 PM			6,90			or I
EV 11	SE 200	87.4	1.5647		23°C	24	38,4353				
-68)			*Specificatio	on: Bell Aer	ospace Textr	on; Report No.	8803-947047, rev. A	4			

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LASSIFIC	NUM SPECIMEN	BER CAPSULE	MATERIAL DESCRIPTION	TEST D CELL	SEALED REFRIG-	ANALYSIS DOCUMENT, IOM. etc.	REMARKS	REVISI	PREPARE
ATIO	BA	Val and	Vol cm 3	IN/OUT	ERAITON IN/OUT	DATE	N2H4 vol		
2 /	4100	7949	N2H4 40cc	28 Jan 80 8:30 AM	18Jan 80 1 PM		3, /2		
	NONE	88.8	0		22°C		40.00		1
14	4028	79 50		18 Mar 80 9:00 AM	11 Mar 80 2 PM		Broken, replaced 7.4 60°C	1 CH	5
	059	85.7	1.5595		2300	24	38.4405		
9	4054	7951		4 Mar 80 8:30 AM	27 Feb 80 11 AM		5.0	July DATE) Hydraz	(DATE)
	116	85.7	0,3517		23.0		39,6483	1979	
		7952	Spare	18 Mar 80 9:00 AM			7, 4	9 79X (PROJ	(REPC
						24		0750 ргор	ATI
4	4058	7953		28 Jan 80 8:30 AM	18 Jan 80 1 PM		3.92	Mater JPL P ellant	L <u>2</u>
	121	87.7	0.7380		22°C	· .	39.262	fal	
3	4039	7954		28 Jan 80 8:30 AM	21Jan 80 2PM		3.8	Compa osal 9	7
	302	86.8	0,5506		2100		39,4494	0-96	
14	4010	7955		18 Jan 60 8:30 AM	21 Jan 80 11 A M		6.08	lity 5 rev.	PAGE
JPL 0	015	88.3	2.5024		2100		37.4976	2	
but-666	4040	7956		28 Jan 80 8: 30 AM	18 Jan 80 1 P M		3,84		۰ ۶
EV 11	303	85.3	0.6097	L	22°C	s	39.3903		
-68)			*Specificatio	n: Bell Aer	ospace Textr	on; Report No.	8803-947047, rev. A		

NUM SPECIMEN	BER CAPSULE	MATERIAL	TEST I CELL	URATION SEALED REFRIG-	ANALYSIS DOCUMENT,	REMARKS	TITLE	IPREPA REVĮ S
BA	Val am?	Vol am ²	IN/OUT	ERATION	IOM, etc. DATE	Na H& wol		TON TON
4047	7957		4 Mar 80 8:30 AM	27 Feb 80 10 AM		Broken, replaced 5.0		TH-
102	86.3	1.0645		23'0		38,9355		Mo
4031	7958		18 Mar 80 9:00 AM	11 Mar 80 2 PM		60°C 9,30	2	CAN
069	84.1	1.4920		2300	24	38,508		
4048	7959		4 Mar 80 8:30 AM	27 Feb 80 10:30 AM		5-0	Hydraz	DATE) July DATE)
103	85.4	1.0666		2300	-	38.9334	ine	1979
4026	7960		4 Mar 80 8:30 AM	27 Fet 80 2:05 PM		6.0	, Mono	9 79X
057	83.5	1.5745		2300	24	38,4255	prop	DRT N 075(JECT)
4067	7961		4 Mar 80 8; 30 AM	27 Feb 80 1 PM		6.9	ellant	o.))0 Mater JPL P
SE210	80.9	1.3813		2300	24	38.6187	Gra	fal
4071	7962		4 Mar 80 8:30 AM	27 Feb 80 1 PM		5.6	1de*	Compa Isal 9
SE 230	88.7	1.5657		2300	· .	38.4343		tib1
4049	7963		4 Mar 80 8:30 AM	27 Feb 80 3 PM		5,1		lity 5 rev.
109	86.0	1.8789		2300		38.1211		2
4059	7964		4 Mar 80 8; 30 AM	27 Feb 80 11 AM	t.	6.0		
127	85.3	0.0056		2300	24	39,9944		
		*Specificati	on: Bell Aer	ospace Textro	on; Report No.	8803-947047, rev. A		

LASSI	NUM SPECIMEN	CAPSIII F	MATERIAL	TEST I	SEALED	ANALYSIS DOCUMENT.	REMARKS	5	TTLE	REVI
FICA			DESCRIPTION	CELL	ERATION	IOM, etc.				IOIS
TION	BA	Val am	Volem	IN/OUT	IN/OUT	DATE	No Ha Nal		-	20
14	4030	7965		4 Mar 80 8:30 AM	27 Febro			6.20		
	068	86.1	1,4278		2300	24	38,5722			741
14	4035	7966		18 Man 80 9:00 AM	11 Mar 80		60°C'	8.20		
	080	84.6	0.7187		2300	24	39.28/3			2
10	4068	7967		4 Mar 80 8:30 AM	ITFER 80 1. PM			4,36	Hydra	July DATE)
	SE211	81.2	1,4163		23°C		38,5837		zine	197
8	4060	7968		4 Mar 80 8:30 AM	27 Fet 80 11 AM			5.9	, Mone	9 79) (PRO
	128	85.5	0.0061		2300		39,9939		oprop	JECT)
14	4033	7969		4 Mar 30 8:30 AM	27 Fel 80 245 PM			6.9	oel lant	Matej JPL J
	078	82.3	0.7104		2300	24	39.2896		E Gr	rial
14	4036	7970		18 Mar 80 9:00 AM	11 Mar 80 1:30 PM		60°C'	9.0	ade*	Compa osal 9
' '	081	86.9	0.7048		2300	24	39,2952			t1b:
14	4025	7971		4 Mar 80 8:30 AM	27 Feb 80			6.2		llity 55 rev
JPL (056	87.0	1.5/12		2300	24	38,4888	1.		. 2
1) (16664	4061	7972		4 Mar 80 8:30 AM	27Feb 80 10.30 AM			9.50		
7EV 11	130	84.6	2.7659		2300		37,2341			
-68)			*Specificat:	ion: Bell Aer	ospace Textro	on; Report No.	8803-947047, rev.	A		

в-26
0			r	1					1		
LASSIFIC	NUM SPECIMEN	BER CAPSULE	MATERIAL DESCRIPTION	TEST D CELL	SEALED REFRIG- ERATION	ANALYSIS DOCUMENT, IOM, etc.	REMARKS		ITLE	REVISIO	
ATIO	BA	Vol cm	Vol cm3	IN/OUT	IN/OUT	DATE	NºH& vol	<u></u>		V O	D BY
z /0	4066	7973		4 Mar 80 8:30 AM	27 Fet 80 1 PM			4.44		/ H	
	SE201	84.1	1.5179		2300		38.4821			20	
15	4051	7974		4 Mar 80 8:30 A M	27 Feb 80 2:15 PM			5.6		RAN)
	112	83.3	0.5820		23.0		39,418				
6	4063	7975		4 Mar 80 8:30 AM	275ef 80 10; 30 AM			5,9	Hydraz	July (DATE)	(DATE)
	133	86.7	0.1157		23°C		39.8843		ine	197	
6	4064	7976		4 Mar 80 8:30 A M	27 Fet 80 11 AM			5.4	, Mono	9 79X	(REP
	134	87.5	0.1191		2300		39,8809		prop	<u>16С1)</u>	ORTN
15	4052	7977		4 Mar 80 8:30 AM	27 Feb 80 2:45 PM			4.24	ellant	Mater JPL P	io.)
	113	83.9	0,5916		2300		39.4084		Gr	fal	
14	4034	7978		4 Mar 80 8:30 AM	27 FLA-80 2; 95 PM			6.2	ade*	Compa Dsal 9	1
1	079	84.4	0.7174		2300	24	39,2826	· · · · · · · · · · · · · · · · · · ·		6411	Ø
6	(4062)	7979	Capsule br Specimen	more more	during	final	calibration mimber 79	(4,96) 80	Reg	265-	PAGE .
1ªL C	(131)		(2,7234) N	10 caps	ulo rep	acement	(37.2766)		Tel	Em	L
0999-V	4062	7980	capsule bro handling or	ben af transfer	ter find from E	l calibra 75 for d.	tion during elivery to Pa	7.0 aden	TU	'X	2 of
11 ∧ ∰	/3/		2.7234 Spe	apsule	repeaced	7998 34 ment 34	37.2766				
-68)			*Specificatio	n: Bell Aer	ospace Textr	on; Report No.	8803-947047, rev. A				

B-27

NUM SPECIMEN	CAPSULE	MATERIAL	TEST D	SEALED REFRICE	ANALYSIS DOCUMENT,	REMARKS		REVI
DΛ	1.0	DESCRIPTION	0000	ERATION	IOM, etc.	11 11		
4050	7981	Voram ²	IN/OUT 4 Mar 80 8;30 A M	17 Fel. 80 3 PM	DATE	N2H& NDI	4.20	TH
110	87.3	1.8275		23°C		38,1725		Mo
4045	7982		4 Mar 80 8: 30 AM	27 Fel 20 10 AM	· · · · · · · · · · · · · · · · · · ·		4,70	RAN
100	87,2	1.0567		2300		38,9433		
4076	7983		4 Mar 80 8:30 AM	27 Feb 80 3:15 PM			4.12	July DATE) Hydraz
251	85.5	1:6193		2300		38.3807		1979 ine
4075	7984		\$ Mar 80 8; 30 AM	27Fell 80 3; 15 PM			4,8) 79X(PROJ
250	87.6	1.6191		2300		38,3809)750 Ест)
4106	7985	N2H4 40 KR	4 Mar 80 8:30 AM	27 Fet 80 9:30 AM		Control	4.70	Mater JPL P ellant
None	85.0	0		2300		40.00		1a1 rop
4069	7986	Special dia ,875	4 Mar 80 8:30 AM	27 Feb 80 1 PM			3.9	Compa osal 9 ade*
SE220	90.5	0,9018		2300		39.0982		0-96
4079	7987		4 Mar 80 8;30 AM	27Feb 80 3:15 PM			3. 88	lity 5 rev.
270	85.8	0.8548		2.3°C		39.1452		2
4081	7988	EPR 515 Oring Krytox 240 Ar.	4 Mar 80 8; 30 AM 10 AU.2 80	27 Fel 80 3 2:30 1M	44 AT-80-076 May 80	WA 4102, 4103	3,92	
280	87,8	0,4024	37 dys	23.0	· .	39.5976		
· · · · · · · · · · · · · · · · · · ·		*Specificatio	n: Bell Aer	ospace Textr	on; Report No.	8803-947047, rev. A		

B-28

01			P				T		<u>– – – – – – – – – – – – – – – – – – – </u>
LASSIFIC	NUM SPECIMEN	CAPSULE	MATERIAL DESCRIPTION	CELL	SEALED REFRIG- ERATION	ANALYSIS DOCUMENT, IOM, etc.	REMARKS		T PREPARE T VEVISIO
ATIO	BA	Vol an	Vol an'	IN/OUT	EN/OUT-	DATE	N2H& vol		
/5	4080	7989		4 Mar 80 8:30 AM	27 Feb 80 3 PM			3,70	4/1
	271	85.2	0,8496		2300		39.1504		Yo I
13	4082	7990	EPR 515 Oring Kry tox 240AC	4 Mar 80 8:30 AM	27 Fet 80 2:10 PM		W/A \$102, 4103	4.2	RAN
	281	86.6	0.4118	65 dys 10:3	AM 2500	- 	39,5882		
10	4072	7991		4 Mar 80 8; 30 AM	27 Feb 80 1 PM			4.5	(DATE) July (DATE)
	SE 231	90.3	1.5383		23.0		38.4617		197 197
	4070	7992	Special dia, 875	4 Mar 80 8:30 AM	27 Feb 80 2:10 PM			3.8	9 (PRO
	SE221	83.8	0.9064	•	2300		39.0936		DAT N 075(
12	4073	7993	Special dia.856	4 Mar 80 8:30 AM	27 Fil 80 21 10 PM			3,76	Nater JPL P
	50240	87,2	0.8975		2300		39.1025	-	Ial Gr.
12	4074	7994	Special dia.856	4 Mar 80 8:30 AM	27 Feb 80 1 PM			3.8	Compa Compa osal 9
		85.4	0.8975		2300	nan Maria	39,1025		2 11b1 0-96
15	4077	7995		4 Mar 80 8:30 AM	27 Fet 80 3:15 PM			3.5	PAGE . Llity 55 rev.
JPL C	260	87.5	0,9202		2300		39.0799		2
2000 X	4078	7996		4 Mar 80 8:30 AM	27 Fel:80 3 PM			3.6	
11 VEV	261	86.7	0.8960		23°C		39.104		
-68)			*Specificatio	n: Bell Aer	ospace Textre	on; Report No.	8803-947047, rev. A		

NUM	BER	WATEDTAT	TEST D	URATION	ANALYSIS			REV	PRE
SPECIMEN	CAPSULE	DESCRIPTION	CELL	REFRIG-	DOCUMENT, IOM, etc.	REMARKS		ISI	PARE
BA	Vol cm3	Volam?	IN/OUT	ENATION EN/OUT	DATE	NºHa Val		NN	
4113	7997	NoTHING PRESSURE: JPL I ATMOSPHERE	18 Mar 80 9:00 AM	11 Mar 80 10:30 AM		60°C 3,80			4 E /
	85.1	0		2.300		Blank sealed			2
4062	7998		19 Mar 80 SATURDAN 1:00 PM	25 Mar 80 2:30 PM	-	See 7980 3.5	•		D.A.
131	85.9	2.7234		21°C		37.2766			\leq
NONE	7999	Nothing	11 Mar 80 10:30 AM	11 Mar 80 10:30 AM		Calib 3.76 open neck	Hydraz	DATE)	(DATE)
	86.8			2300			ine		107
							Mo	1	
							nopr	POJEC	EPORT
					· · · · · · · · · · · · · · · · · · ·		opel		5 N.O.
							lant	later PL P	
							Gr	1al rope	
4114	N O NUMBER	40.0 CC N2H4	28 Fet 80 8 apr 80	Sealed 3 placed in	49 AT-80-081 May 80	W/A 4103	ade*	Companosal 9	
SPECIAL		CUZ annary sis	40 dys	<i>γj</i> =0		40.00 CC		96-0	
								11t	
								y ev.	
								2	
	1	*Specifi	Poll Acr	L Tertr	on: Report No	8803-947047, rev. A	1		

в-30

NUM	BER		TEST DURATION		ANALYSIS	Special copsulsa	TITL	REV
SPECIMEN	CAPSULE	MATERIAL DESCRIPTION	CELL	REFRIG- ERATION	DOCUMENT, IOM, etc.	1.15 IN CHREMARKS 0, D.	m	TO-
BA	Vol em3	Vol cm ³	IN/OUT	IN/OUT	DATE	N2H& VOL = 40.0-spec	em3	
4115	79201	NoTHING	10 Mar 80 8:30 AM					1
		Open mech			· · · ·	Blank calibration		NO V
4013	79202		18 Mar 80 9:00 AM					RAI
023	103.6	1.3765				38.6235		<u> </u>
4116	79203	NOTHING	18 Mar 80 9:00 AM			60°C	Hydra	(DATE) July (DATE)
		Open nech				Blank calibration	zíne	197
4014	79204		18 Mar 80 9:00 AM				, Moj	19 (19)
024	105.1	1.2676				38.7324	nopro	POJECT
4015	79205		18 Mar 80 9:00 AM			60°C	JPL pellar	Mate
025	102.2	1.3479				38.6521	Prop lt Gr	rial
4016	79206		18 Mar 80 9:00 AM			60°C	osal : ade*	Comp
026	106.5	1.3607				38.6393	90-96	tibi
							5 rev	lity
							· · · · · · · · · · · · · · · · · · ·	
	L	tSpecificati	Bell Acr	Cenace Tevta	L	8803-947047. rev. A		

B-31

APPENDIX C

CO₂ ANALYSIS

A. DETERMINATION OF CARBON DIOXIDE ABSORBED BY HYDRAZINE

The general laboratory test setup for CO_2 analysis is shown schematically in Fig. C-1. The sulfamic acid solution is prepared by dissolving 150 g of reagent grade material in 1.0 liter of distilled water. To reduce the CO_2 content of the sulfamic acid, high-purity helium, passed through Ascarite, is bubbled through the sulfamic acid solution via the glass frit, which provides a fine gas dispersion and efficient purging. The helium gas is passed through the sulfamic acid delivery tube for about 16 hours at 50-60 cm³/min. The exit end of the helium gas from the sulfamic acid bottle is protected against air and CO_2 with an Ascarite tube. This Ascarite tube is replaced with a new one after the helium purge. With the precautions outlined, the blank CO_2 is under 2.0 ppm.

The apparatus is standardized by means of a $NaHCO_3$ solution prepared by dissolving 0.381 g of dried $NaHCO_3$ in 1.0 liter of distilled water. The solution is stored in glass, and air exposure is minimized. This solution provides 0.20 mg CO₂ per milliliter. Its CO₂ content is 200 ppm by weight.

The column is 6.0-mm-diam tubing, 3.66 m long (0.24-in.-diam, 12 ft long), filled with 60 to 80 mesh F & M Polypack No. 5. This packing gives good separation of CO_2 at ambient temperature. The peaks are sharp, permitting direct reading of the heights and eliminating the need for peak area measurements. The column is bent into a number of 0.7-m (2-ft) sections arranged close together and contained in a glass jacket. The filament-type thermal conductivity detector unit is kept at ambient temperature in a glass dewar to minimize temperature fluctuations. A 1.0-mV recorder records the detector output. Helium flow is 60 cm³/min.

The first step in the analysis is the determination of the blank: the CO_2 picked up from the reagents and the system. The flow of the high-purity helium purge gas, after passage through Ascarite, is adjusted to 50 cm³/min by means of a flowmeter in the system. A 60-ml sulfamic acid solution is run into the unit via the stopcock. The stirrer is adjusted to give vigorous constant stirring. Once set, the helium flow and stirring are kept fixed through the whole run.

After addition of the sulfamic acid, the helium gas is passed through the traps for 30 min to purge the system of air. The CO_2 trap is then immersed in liquid nitrogen to the top level of the glass beads. The flow of helium is continued for 20 min, after which time the stopcock on the CO_2 trap is turned to isolate the loop on the trap.

The trap, immersed in liquid nitrogen, is transferred to the gas chromatograph sampling system. The stopcock on the CO_2 trap is turned so as to evacuate the noncondensable gases in the trap and then turned to isolate the loop containing the frozen CO_2 . LIBERATION AND TRAPPING OF CO2



Figure C-1. Test Procedures for CO_2 Analysis

C-2

The next step is to flow the helium gas through the branched leg of the sampling system. During this operation the liquid nitrogen is removed, and the CO_2 trap thawed with warm water. After a few minutes, the stopcock is turned so as to flush the CO_2 with helium into the chromatographic column for separation and assay. The blank run is repeated until consistent, low values are obtained.

The standardization run is made in the same manner as the blanks, except that after a 5-min preliminary purge with helium, 0.50 ml of standard NaHCO₃ solution is injected into the vigorously stirred sulfamic acid via the septum on the sulfamic acid unit. The released CO_2 is frozen out during the 20-min duration in the CO_2 trap immersed in liquid nitrogen. The trapped CO_2 from the standard solution is transferred to the gas chromatographic sampling system. This yields a peak height for a standard of 100 ppm CO_2 .

The CO_2 in the hydrazine is similarly determined. A 1.0-ml sample is injected into the sulfamic acid solution via the septum, and the released CO_2 is swept out of the solution for a period of 20 min. The hydrazine injections should be made rapidly with a minimum exposure to air. The sulfamic acid solution is sufficient to neutralize 1.0 ml of hydrazine and should therefore be discarded after each hydrazine analysis. If another sample is to be run, the sulfamic acid unit is refilled, and the blank and the standard determinations are made as before.

B. CALCULATION FOR CARBON DIOXIDE CONTENT

The formula for determining parts per million of carbon dioxide is

<u>peak sample - peak blank</u> x 100 peak standard - peak blank

For hydrazine, where the density can be taken as 1.0, a density correction term is not applied. The error due to this omission is about 1%, well within the +10% precision for CO₂ determination when the values are under 20 ppm.

C. CONCLUSION

The method described provides meaningful results for the determination of CO₂ in hydrazine or its methyl-substituted derivatives.

APPENDIX D

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TEST COUPON PHOTOGRAPHS IN THE POSTTEST CONDITION

Figure D-1 shows test specimens from Program A, the secondary containment system; Figure D-2 shows test specimens from Program B, the primary containment system.



Figure D-1. Test Specimens, Hydrazine Decomposition Program "A" - Secondary Containment System



Figure D-1. Test Specimens, Hydrazine Decomposition Program "A" -Secondary Containment System (Continued)



Figure D-1. Test Specimens, Hydrazine Decomposition Program "A" - Secondary Containment System (Continued)



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Figure D-1. Test Specimens, Hydrazine Decomposition Program "A" - Secondary Containment System (Concluded)



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Figure D-2. Test Specimens, Hydrazine Decomposition Program "B" -Primary Containment System (Continued)



Figure D-2. Test Specimens, Hydrazine Decomposition Program "B" - Primary Containment System (Continued)



Figure D-2. Test Specimens, Hydrazine Decomposition Program "B" - Primary Containment System (Continued)







Figure D-2. Test Specimens, Hydrazine Decomposition Program "B" - Primary Containment System (Concluded)

APPENDIX E

347 CRES WELDS

A. INTRODUCTION

Some concern was expressed by the sponsor that the 347 CRES diaphragm in the burst disc assemblies (BA 023 to BA 026) might be subject to stress corrosion cracking in a hydrazine environment. All four specimens were removed from storage at 12.5 months rather than the scheduled 24 months. A cursory examination at JPL was inconclusive. However, a very thorough examination by BAT revealed that any crack formation in the diaphragms was due to the manufacturing and coining (crimping) process.

In the NASA long-term material compatibility test program, a few welded 347 CRES specimens were still in storage. Two were removed - one of which had been stored in the stressed configuration. These were thoroughly tested to determine the differences induced by storage while stressed versus storage while relaxed. Those results are not directly relatable to the 347 CRES diaphragms, but it was believed that the information gained would be useful to this program.

B. SAMPLES

All samples were Type-347 CRES which had been heliarc welded. For all samples, the area analyzed was from the "top" surface immediately adjacent to the weld. The samples analyzed were those described below:

JPL 0981 - welded, no $N_{2}H_{4}$ (control)

JPL 2005 - welded, in N_2H_4 - 4054 days

JPL 1977 - welded, stressed in N_2H_4 - 4054 days

For samples 2005 and 1977, the areas chosen had been at the liquid/gas interface during the testing. A dark discoloration of interest was particularly obvious in this area.

C. XPS EXAMINATION

All three samples had been rinsed with distilled water upon removal from the hydrazine. Prior to analysis with XPS technique, the samples were further cleaned in an ultrasonic cleaner by serial rinses in trichloroethylene, acetone, and absolute ethanol. Samples were then blown dry with dry nitrogen gas.

The XPS spectrometer (modified HP5950A) averages over a region approximately 1 mm x 5 mm in area and 50-100 Å in depth. Because of the exponential attenuation of the photoelectrons, the immediate atomic surface (approximately 30 Å) was weighted more heavily than the rest.

Except for Na, Zn, and N, all the observed elements are expected for 347 CRES. (The approximate theoretical composition of 347 CRES is 0.08% C, 2% Mn, 0.05% P, 0.03% S, 1% Si, 18% Cr, 10% Ni, 0.1% Ta, remaining % Fe.) Oxygen is present in the form of various metal oxides and hydroxides as discussed below.

1. Chromium Region

JPL 0981. Cr_2O_3 , CrO_3 , and some reduced Cr^{+3} species at lower binding energy (BE) than CrO_3 , but higher than chromium metal, were observed. chromium hydroxides may also be present.

<u>JPL 2005 and JPL 1977</u>. No differences were observed between these two samples. Spectra were consistent with Cr_2O_3 and chromium hydroxides. The atomic percent of Cr observed increased in the order 0981 <2005 \cong 1977.

2. Fe Region

The atomic percent of iron observed decreased in the order $0981 > 2005 \cong 1977$. For all three samples, effectively Fe₂O₃ and iron hydroxides were observed. There were some differences in the high BE side of the oxide peak. A small amount of FeO was observed on the control (0981) as well as the possibility of low-level FeO or iron sulfides.

3. Mn Region

The atomic percent of manganese observed decreased in the order 0981 >2005 \cong 1977. The spectra for all three samples may be assigned to MnO and manganese hydroxides.

4. Zn Region

The atomic percent of zinc observed decreases in the order 0981 >2005 \cong 1977. ZnO was present in all three samples.

5. Ni Region

The atomic percent of nickel observed was approximately the same for all three samples, with perhaps slightly more for 2005 as compared to 1977. The Ni was present as Ni $_20_3$ and nickel hydroxides. The lower BE on 0981 was probably due to Fe $_20_3$.

6. Carbon Region

Approximately the same amount of carbon was observed on all three samples. Primarily aliphatic carbon was present although substantial intensities in the C-O and C-N regions were observed, showing some differences in detail between the samples.

7. Oxygen Region

Approximately the same amount of oxygen was observed on all samples. The primary peak was due to metal oxides and hydroxides. The lower BE on 0981 was probably due to Fe_2O_3 .

8. Nitrogen Region

The nitrogen intensity increases significantly in the order 0981 <2005 \cong 1977. The primary differences appear on the low binding energy side of 2005 and 1977, where a shoulder characteristic of reduced nitrogen species such as amines and ammonia transition metal complexes was observed. The high binding energy peak was consistent with a variety of species including protonated amines and amide polymers as well as hydrazine salts and nitrites.

D. CHEMICAL ANALYSIS

Chemical analysis of propellant and analysis of the decomposition gases indicated no significant differences between stressed and unstressed conditions (Table E-1). Both specimens were exposed to hydrazine for 4054 days.

	Prope1		
Specimen	Decomposition (wt%/yr)	Dissolved Fe (mg)	N_2H_2 gas, cc x10 ⁻³ ·day ⁻¹ ·cm ⁻²
JPL 1977, stressed	0.174	0.21	1.48
JPL 2003, unstressed	0.154	0.21	1.15

Table E-1. Summary of Posttest Hydrazine Analysis

E. SEM EXAMINATION OF SPECIMENS

SEM photomicrographs of the surfaces of the specimens indicated that some pitting has occurred at the liquid-vapor interface. As seen in the accompanying photographs, (Figures E-1 to E-4) the distribution of pit sizes varies between the stressed and unstressed specimens. These same samples were sectioned and etched. Photomicrographs of the cross sections revealed no intergranular or intragranular corrosion. The markings seen on the photographs are the result of over-etching.

F. CONCLUSIONS

- 1. The pattern of surface corrosion was similar for each of the specimen examined.
- 2. No intergranular or intragranular corrosion was observed in stressed specimen.

- 3. Fe and Mn dissolved more readily than does Cr, leaving a corroded surface rich in chromium.
- 4. The XPS examination indicated no difference in the chemical nature of stressed and unstressed specimens.
- 5. Only minor differences were observed in metal content or decomposition of propellant between stressed and unstressed configurations.





UNSTRESSED WELD

CONTROL WELD

Figure E-1. CRES 347 Weld Specimens, Surface Features at Weld



STRESSED HAZ



CONTROL HAZ

Figure E-2. CRES 347 Weld Specimens, Surface Features at Heat-Affected Zone

Е-6



STRESSED WELD-TENSION



UNSTRESSED WELD





STRESSED HAZ-TENSION

UNSTRESSED HAZ

Figure E-4. CRES 347 Weld Specimens, Cross Sections at Heat-Affected Zone

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