TSTA LOOP OPERATION WITH 100 GRAMS-LEVEL OF TRITIUM

- FULL COMPONENTS MILESTONE RUN IN JUNE, 1988 -

December 1988

Hiroshi YOSHIDA, Hiroshi FUKUI, Shingo HIRATA*¹

Toshihiko YAMANISHI, Taisei NAITO*²

J. L. ANDERSON*³ J. R. BARTLIT*³ C. R. WALTHERS*³, R. H. SHERMAN*³

D.O. COFFIN*³ R.V. CARLSON*³ R.S. WILLMS*³ and J.E. NASISE*³

日 本 原 子 力 研 究 所 Japan Atomic Energy Research Institute

JAERI-Mレポートは、日本原子力研究所が不定期に公刊している研究報告書です。 入手の問合わせは、日本原子力研究所技術情報部情報資料課(〒319-11茨城県那珂郡東 海村)あて、お申しこしください。なお、このほかに財団法人原子力弘済会資料センター (〒319-11 茨城県那珂郡東海村日本原子力研究所内)で複写による実費頒布をおこなって おります。

JAERI-M reports are issued irregularly.

Inquiries about availability of the reports should be addressed to Information Division, Department of Technical Information, Japan Atomic Energy Research Institute, Tokaimura, Naka-gun, Ibaraki-ken 319-11, Japan.

© Japan Atomic Energy Research Institute, 1988

編集兼発行 日本原子力研究所

印 刷 ㈱原子力資料サービス

TSTA Loop Operation with 100 Grams-Level of Tritium - Full Components Milestone Run in June, 1988 -

Hiroshi YOSHIDA, Hiroshi FUKUI, Shingo HIRATA*1
Toshihiko YAMANISHI, Taisei NAITO*2, J.L. ANDERSON*3
J.R. BARTLIT*3, C.R. WALTHERS*3, R.H. SHERMAN*3
D.O. COFFIN*3, R.V. CARLSON*3, R.S. WILLMS*3
J.E. NASISE*3

Department of thermonuclear Fusion Research
Naka Fusion Rsearch Establishment
Japan Atomic energy Research Institute
Naka-machi, Naka-gun, Ibaraki-ken

(Received November 7, 1988)

A fully integrated loop operation test of Tritium systems Test Assembly (TSTA) with 107 grams of tritium was completed at Los Alamos National Laboratory (LANL) in June, 1988. In this test, a compound cryopump with a charcoal panel was incorporated into the main process loop for the first time.

The objectives were (i) to demonstrate the compound cryopump system with different flow rates and impurities, (ii) to demonstrate the regeneration of the compound cryopump system, (iii) to accumulate operating experience with other process systems such as the fuel cleanup system, the isotope separation system, the tritium supply and recovery system, etc. and (iv) to improve the data-base on TSTA safety systems such as the secondary containment system, tritium waste treatment system and tritium monitoring system.

This report briefly describes characteristics of the main subsystems observed during the milestone run.

Keywords: Fusion Reactor, Tritium Technology, Fusion Fuel Cycle,
Plasma Exhaust Gas, Compound Cryopump, Fuel Cleanup, Isotope
Separation, Secondary Containment, Tritium Monitoring

^{*1} Kawasaki Heavy Industries, Ltd.

^{*2} Mitsubishi Atomic Power Industries, Inc.

^{*3} Los Alamos National Laboratory

100 グラムレベルのトリチウムを用いたTSTAループ試験 - 1988 年 6 月のフル・コンポーネント試験-

日本原子力研究所那珂研究所核融合研究部 吉田 浩·福井 裕·平田 慎吾·山西 敏彦 内藤 大靖^{*2} J. L. ANDERSON ^{*3} J. R. BARTLIT ^{*3} C.R.WALTHERS ^{*3} R.H. SHERMAN ^{*3}D.O. COFFIN ^{*3} R. V. CARLSON ^{*3} R. S. WILLMS ^{*3}I. E. NASISE ^{*3}

(1988年11月7日受理)

本報は、1987年6月に発足した核融合燃料技術に関する日米協力(Annex IV)の下で行った 実規模D-Tガス循環試験(1988年6月実施分)の結果をまとめたものである。この試験は、 Annex IV 第1年目におけるTSTAの各種試験(ループ試験を6回,深冷蒸留塔,複合クライオポンプ等のサブシステム試験を6回)を総合したものであり、模擬トーラス、複合クライオポンプ及び同再生ポンプ系、D-Tガス精製系、NBI ガス循環系、深冷蒸留塔系等の主要システムを連結し運転することを目的とした。

那珂研究所:〒 311 - 01 茨城県那珂郡那珂町大字向山 801 - 1

^{*1} 川崎重工業㈱

^{*2} 三菱原子力工業 ㈱

^{*3} ロスアラモス国立研究所

JAERI-M 88-242

Contents

1. Introduction .	•••••	1
1.1 Layout of J	AERI/TSTA-LANL collaboration	1
1.2 Brief descr	ription of TSTA	2
1.2.1 Process	systems	2
1.2.2 Safety	systems	3
1.2.3 TSTA de	sign dose	4
1.2.4 Acciden	t considerations	4
2. Test plan		21
2.1 Objectives	•••••	21
2.2 Configurati	on	21
2.3 Procedures		21
3. Results of loo	p operation	25
3.1 Process sys	tems	25
3.1.1 Vacuum	system (VAC)	25
3.1.2 Tritium	storage and supply system (UTB)	25
3.1.3 Fuel cl	eanup system (FCU)	26
3.1.4 Isotope	separation system (ISS)	26
3.2 Safety syst	ems	27
3.2.1 Seconda	ry containment system (SEC)	27
3.2.2 Room ra	diation levels	28
3.2.3 Stack r	adiation levels	28
3.2.4 Tritium	waste treatment system (TWT)	29
4. Results of sys	tem shutdown	63
4.1 Process sys	tems	63
4.2 Safety syst	ems	63
4.2.1 Seconda	ry containment system (SEC)	63
4.2.2 Tritium	waste treatment system (TWT)	64
5. Conclusion	•••••	72
Acknowledgement		73
References		73

目 次

1.	J	予		論		1
	1.	1		J A	ERI/TSTA-LANL協力の概要	1
	1.	2	,	TS	TAの概要	2
		1.	2.	1	プロセスシステム	2
		1.	2.	2	安全システム	3
		1.	2.	3	被曝管理	4
		1.	2.	4	想定事故	4
2.	į	试	験	計画		21
	2.	1	Ē	試験	语 目的 ·······	21
	2.	2	É	試験	装置	21
	2.	3	Ē	試験	: 方法 ·······	21
3.	i	式	験	結果		25
	3.	1	-	プロ	セスシステム	25
ż		3.	1.	1	真空排気系	25
		3.	1.	2	トリチウムの貯蔵・供給系	25
		3.	1.	3	精 製 系	26
		3.	1.	4	同位体分離系	26
	3.	2		安全	システム	27
		3.	2.	1	第 2 次格納システム ····································	27
		3.	2.	2	室内トリチウム濃度	28
		3.	2.	3	スタックトリチウム濃度	28
		3.	2.	4	トリチウム廃ガス処理システム	29
4.		シ	ス	テム	。停止	63
	4.	1		プロ	1セスシステム	63
	4.	2		安全	システム	63
		4.	2.	1	第 2 次格納システム	63
,		4.	2.	2		64
5.		結		ā	À	72
謝		i	辞			73
参	老	文	献			73

1. INTRODUCTION

1.1 LAYOUT OF JAERI/TSTA-LANL COLLABORATION

The joint work of the TSTA Loop Operation in the first Annex IV year (June 11, 1987 - June 10, 1988) defined by the JAERI/TSTA-LANL collaboration program under Japan/US Fusion Cooperation Program was successfully completed, while producing many scientific and technological advances.

Table 1.1 summarizes the TSTA activities achieved in this period.

The loop tests in June and July, 1987 were the first loop operation of the TSTA process systems and safety systems with 100 grams-level of tritium. An emergency situation due to the loss of helium refrigeration of the cryogenic hydrogen isotope separation system, which was one of the most serious scenarios in the TSTA failure mode and effects analysis, happened during the June run. The test eventually had to be halted. There were, however, no offnormal environmental tritium releases or personnel exposures, and no resulting damage to TSTA systems which might have impacts on the Annex IV first year program. The second loop operation was successfully resumed, only one month after this unexpected event.

Hydrogen isotope separation experiments on a single column with H-D system (October, 1987) and D-T system (December, 1987), and with a two-column cascade configuration with H-D-T system (April, 1988) were performed to study isotope separation characteristics of full-scale cryogenic distillation columns.

Demonstration experiments on the fuel cleanup system, FCU (Front-end: impurity removal process consisted of catalytic oxidizer and cryogenic molecular sieve beds; Regeneration process: D-T recovery process of catalytic oxidizer, DTO freezer and hot metal beds) were carried out from January through March, 1988. The March run was the first complete functional test of the FCU under a full loop operating condition using impurities of 0.099%N $_2$ and 0.019%CH $_4$ in the D-T stream.

Compound cryopump experiments of TSTA-VAC system (composed of the torus mockup, two different compound cryopumps having D-T condensing chevrons and a He adsorbing charcoal panel, and a regeneration train) had been done with $\rm D_2$, $\rm H_2$, D-T, He, $\rm N_2$ and

their mixtures in the period from March, 1988 through July, 1988. The D-T experiments were done in both operating modes of non-loop flow and full loop flow with FCU, ISS and TPU (gas circulation system in the compound cryopump regeneration train). The loop operation of May-June, 1988 was the first test of all components of a fusion fuel cycle on an engineering scale. The pumping speed of D-T, D-T-He (10 - 25%), D-T-N $_2$ (1%), and the He/D-T

separation performance of both chevrons and a charcoal panel, and the regeneration back to the TSTA process loop were successfully demonstrated during a series of VAC system experiments.

This report briefly describes the results of this integrated operation of the full fuel processing loop test.

1.2 BRIEF DESCRIPTION OF TSTA [Ref. 1,2 and 3]

The detailed description of the above systems is given in previous reports "TSTA Loop Operation with 100 grams-level of Tritium - Milestone Run in June, 1987 - " and "TSTA Loop Operation with 100 grams-level of Tritium - Milestone Run in July, 1987 - ".[Ref. 1 and 2]

1.2.1 PROCESS SYSTEMS

Figure 1.1 shows the conceptual process loop which was used during this run. The current TSTA process loop, which is shown in Figure 1.2, is composed of the following subsystems.

(1) VAC

The Vacuum Facility (VAC) with a torus mock-up, two different designs of compound cryopumps, and a turbopump for regeneration of cryopumps (Figure 1.3).

(2) UTB

Tritium and other hydrogen isotopes storage /supply system with five large uranium beds (UTB-1, 2, 3, 4 and 5). Figure 1.4 shows the general arrangement of UTB.

(3) ISS

Hydrogen isotope separation system with four interlinked cryogenic distillation columns (I, H, D and T) and two hydrogen isotope equilibrators (EQ).

(4) TP1

Transfer pump system with a scroll pump (S) and metal bellows pumps (MBPA and MBPB).

(5) TP3

Transfer pump system with metal bellows pumps (MBPA and MBPB), a gas mixer (MX) and a hydrogen isotope equilibrator (EQ).

(6) FCU

Fuel cleanup system with catalytic reactors (CR1 and 2), cryogenic molecular sieve beds (MSB1, 2, 3 and 4), uranium beds (HMB4 and 5) and a tritiated water vapor freezer (DTOF). This system includes a NBI front end composed of MSB3 and 4.

(7) LIO

Load-in/Load-out system for tritium using \mathbf{T}_2 gas shipping containers.

1.2.2 SAFETY SYSTEMS

The major environmental and safety subsystems of the TSTA facility are as follows;

(1) SEC

The primary process equipment and process lines are doubly contained with secondary containment consisting of gloveboxes and plastic tubing, with nitrogen gas purge and control of their atmospheric pressures. Figure 1.5 shows the concept of the instrumentation of a typical glovebox. Purge gas (N_2) is supplied to the glovebox at a preset tritium level.

(2) TWT

The tritium waste treatment system (Figure 1.6) processes all tritium-bearing gaseous effluents generated in various subsystems in TSTA. This system is operated either in once-through or recirculation mode as necessary, and routes the effluents to the TSTA stack after detritiation and monitoring the tritium release level.

The capacities of the TWT compressors are 90 and 25 STP- m^3/h .

(3) TM

A number of tritium monitors (Stack, Duct, Room, Glovebox and Process) have been installed in the TSTA facility. Figure 1.7 shows the location of the room-tritium-monitors. The monitors perform the following functions; (i) quantitative determination of stack releases, (ii) monitoring tritium concentrations in room and room-exhaust air, secondary containment atmosphere, and process system lines, (iii) initiation of local alarms and computer-directed signals (in secondary containment and room or room exhaust ducts) and (iv) initiation of room air isolation and an evacuation alarm and computer-directed signals.

(4) ETC

The TSTA main cell contains approximately 3000 m³ of building atmosphere which would be contaminated with tritium in the event of an accidental release from a secondary containment. This system, the emergency tritium cleanup system (Figure 1.8), has the primary function to reduce the probability and amount of a tritium release to the environment from the TSTA main cell after such an accident.

The free air capacity of the primary compressor is $2500~\text{m}^3/\text{h}$ at 585~Torr (0.77 atm) and 293~K.

(5) VEN

The ventilation system is divided into two zones. The Zone I system provides heating and ventilation for areas (rooms for

non-tritium handling) from which tritium will be excluded and is maintained at a slight positive pressure (0.23 Torr) with respect to atmospheric pressure. The Zone II system (shown in Figure 1.9) for the main cell and other tritium handling rooms is maintained at a slight negative pressure (0.23 Torr) with respect to atmospheric pressure to minimize possible diffusion of tritium to Zone I or to the environment.

The ventilation capacity of the Zone II system is 15291 m3/h.

(6) MDAC

The TSTA is designed to be a computer controlled system. This system, master data acquisition and control, contains two computers (shown in Figure 1.10); the process computer and an interface, and the backup computer.

1.2.3 TSTA DESIGN DOSE

For TSTA to handle tritium the following subsystems must be on line: SEC, TM, TWT, ETC, VEN, MDAC, Power and Utilities. Most safety system operations are under total computer control. Table 1.2 shows the TSTA design dose. In addition, actions are taken to reduce radiation exposure from tritium to As Low As Reasonably Achievable (ALARA).

The value for occupational exposure is one-fifth of the guide level (5 rem/year) in DOE manual (DOE Order 5480.1A Chapter XI). The nonoccupational dose will be less than 1 mrem/year for the routine environmental tritium release rate of 200 Ci/year (selected as a design goal for TSTA). The dose for personnel in adjacent buildings will be less than 500 mrem/year.

1.2.4 ACCIDENT CONSIDERATIONS

(1) Failure Modes and Effects Analysis

TSTA subsystems have been analyzed with Failure Modes and Effects Analysis to evaluate the safety features of the design, identify critical failure modes, and recommend alternatives or precautions that will mitigate the effect of failures.

Table 1.3 gives a summary of the effects resulting from representative accidents which have been postulated for TSTA. Table 1.4 shows an expected tritium inventory in TSTA.

Analysis and discussions of the postulated accident and failures for TSTA are described in the TSTA Final Safety Analysis Report.[Ref.3]

(2) Accident Scenarios

(i) A single failure in any secondarily contained subsystem, i.e., a release into a glovebox, will not result in exposures to individuals or release to the environment, since the TWT will process the secondary containment atmosphere.

(ii) Double failures in systems which have secondary containment or a single failure in a single containment containing low levels of tritium may result in a tritium release into the experimental area. The extent of personnel exposures depends on the tritium concentration in the area, the tritium form (oxide or elemental), and the time required to exit from the area. The exposures for all credible accidents are within the TSTA design goal (25 rem) for accident situations.

Examples of accident scenarios are:

- glove rupture plus a release in that glovebox,
- rupture of ISS system.
- TWT low pressure receiver failure.

The release to the environment for the above type of accident is very small because the ETC is designed to process the contaminated room air. For a postulated release of 100

grams (10^6 Ci) of tritium into the facility only 10 Ci would be released from ETC to the environment (design).

(3) Emergency Evacuation Procedures

Evacuation procedures for a TSTA emergency have been established in accordance with the TSTA Emergency Plan. Any tritium release to the room atmosphere is quickly picked up by the room monitor. Local alarms will be given at each of three levels of tritium concentrations measured.

- Low $(20x10^{-6} \text{ Ci/m}^3)$; amber light
- Mid $(100x10^{-6} \text{ Ci/m}^3)$; red light, steady sonalert (mutable)
- High(10000x10⁻⁶ Ci/m³); flashing red light, pulsing sonalert

The evacuation alarm is automatically triggered by the high level.

The stay time in the contaminated room to keep the committed dose below 50 mrem is:

- (i) without protective clothing; stay time (min) = 6000/tritium level (10⁻⁶ Ci/m³),
- (ii) with bubble suits and supplied air; stay time (min) = 1200000/tritium level (10^{-6} Ci/m³).

Re-entry at a concentration over 10^{-3} Ci/m³ requires full protective clothing (bubble suits) and may only be made if accompanied by a second person who is also suited up. Self-contained supplied air suits also be used for re-entry with permission of TSTA management and HSE-1 (branch of the LANL health physics group).

Table 1.1 Schedule of TSTA Operations

MAJOR EXPERIMENTS & OPERATIONS AT TSTA, FIRST YEAR OF ANNEX IV

	Date	Operation	Description
1.	June 22-26, 1987	loop test of FCU/ISS	with continuous impurity flow with continuous impurity flow
2.	July 23-27, 1987	loop test of FCU/ISS	• •
3.	September 1987	maintenance on ISS	lower ISS vacuum jacket after tritium use
4.	October 4-9, 1987	non-loop ISS (UTB,TPU)	single column experiments of ISS
5.	October 1987	FCU-related testing	begin assembly/tests (w/o tritium) of high temperature metal beds
6.	December 6-11, 1987	non-loop ISS (UTB,TPU)	single column experiments of ISS
7.	January 1988	non-loop FCU	non-tritium test of FCU operation
8.	February 7-9, 1988	loop test of FCU/ISS	preparatory for first full integration of FCU/ISS with continuous impurity flow
9.	February 21-24, 1988	loop test of FCU/ISS	same as item 8 above
10.	February 28	loop test of FCU/ISS	first full integration of FCU/ISS
	March 4, 1988		with continuous impurity flow
11.	March- May 1988	non-loop VAC tests	without tritium
	April 4-8, 1988	non-loop ISS (UTB,TPU)	ISS experiments, 2 column
	May 31-June 6, 1988	loop test of VAC/FCU/ISS	first integrated operation of full fuel processing loop

Table 1.2 TSTA Design Dose Commitments

Condition Design Objectives Radiation Workers Public Public bNormal Operation (mrem/yr) 1000 170 ^CAccident (rem)

^aDesign dose commitment is the exposure which, under the given conditions, shall not be exceeded in design. These dose commitments are less than specified in DOE 5480.1A.

bExamples:

- (1) Normal component operation including shutdown. repair, maintenance, and checkout
- (2) Operational occurrences such as leaks, loss of power. and component malfunctions likely to occur once or more during the life of the facility

25

5

CExamples:

- (1) Very low probability events such as "most intense predicted" natural phenomena
- (2) Major component failure events which are not likely to occur during the life of the facility

Table 1.3 Summary of Postulated Accidents

		Delescable	e Inventory	Mitigation	Release to Stack	Boundary ² Site Whole Body	Whole Body Worker Dose Rate ³	
Failu		Ci	Form	Method	Cf	Dose (mrem)	(mrem/min)	Freq
1 6110	<u></u>							
1. a)	Rupture or	100	DT	A	100	1 x 10 ⁻⁶	1 x 10 ⁻³	E
	large leak			В	1 x 10 ⁻³	7 x 10 ⁻⁷	1 x 10 ⁻³	E
	from torus							
	plus loss							
	of SEC			_		7 x 10 ⁻²		-
ь)	Same as la,	100	DTO/HTO/T _Z O	A	100 1 x 10 ⁻³	7 x 10 ⁻⁷	67 67	E E
	accompanied			В	1 x 10 -	/ x 10 '	0/	E
	by a fire	5.8 x 10 ⁴	DT	A	5.8 x 10 ⁴	7 x 10 ⁻⁴	0.8	Ε
2. 2)	Rupture of	5.8 X 10	וע	В	5.8 x 10 ⁻¹	4 x 10 ⁻⁴	0.8	E
	cryopump plus loss of SEC			•	3.0 X 10	, , , ,	2.0	-
ы	Same as 2a,	5.8 x 10 ⁴	HT0/T ₂ 0	A	5.8 x 10 ⁴	38	3.9 x 10 ⁴	E
٠,	accompanied	0.0 x 10		8	5.8 x 10 ⁻¹	4×10^{-4}	3.9 x 10 ⁴	E
	by a fire							
3. a)	Rupture of	9.7 x 10 ⁵	DT.T2	С	1	7 x 10 ⁻⁴	0	E
	distillation		. 2					
	columns to							
	vacuum jacket				_	_		
b)	Same as 3a	9.7 x 10 ⁵	DT.T2	A	9.7 x 10 ⁵	1 x 10 ⁻²	13	E
	followed by a		•	В	9.7	6 x 10 ⁻³	13	E
	breach of the							
	vacuum jacket	_			E	,	5	_
c)	Same as 3b,	9.7 x 10 ⁵	D ₂ 0,DT0,T ₂ 0	A	9.7 x 10 ⁵	6 x 10 ²	6.5 x 10 ⁵ 6.5 x 10 ⁵	E
	accompanied by			В	9.7	6 x 10 ⁻³	6.5 X 10"	Ē
	a fire, release	•						
	at 50 m	3000	DT,T2	С	3 x 10 ⁻²	2 x 10 ⁻⁵	O	E
4. 4) Leakage of	•	21.12	·				
	transfer lines from NBI and					•		
	IMS to FCU int	0						
	secondary							
	containment.					_K	4.0 x 10 ⁻²	E
ь) Same as 4a.	3000	DT,T ₂	A	3000	4 x 10 ⁻⁵	4.0 x 10 ⁻²	E
	followed by		•	В	3 x 10 ⁻²	2 x 10 ⁻⁵	4.0 x 10 °	t
	breach of							
	secondary							
	containment					_	2.0 x 10 ³	E
	;) Same as 4b,	3000	DTO,D ₂ 0,T ₂ 0	A	3000	2 2 x 10 ⁻⁵	2.0 x 10 ⁻³	E
	accompanied			В	3 x 10 ⁻²	2 1 10		
	by a fire	5		612	NA	4.7 x 10 ³	NA ·	E
5.	Aircraft cras	9.7 x10°	HTO	NA	BA .			
	into facility				NA	0.23	KA	E
6.	Earthquake	1.45 x 10 ⁶		NA NA	NA NA	11 × 10 ³	NA	Ē
	totał		нто	RA.	· - ·			
	destruction							

Mitigation Method

A - Ventilate Experimental Room

^{8 -} Process Room Air with Emergency Cleanup System. Release form is tritiated water vapor.

C - Process Contaminated Air with Tritium Waste Treatment System

MA - Not Applicable

 $^{^2}$ This is maximum dose comitment and includes the skin intake. The dose is determined from

Fig. B-1. The dose to the skin itself is discussed in Sec. 6.1. Site boundary is 400 m from TSTA.

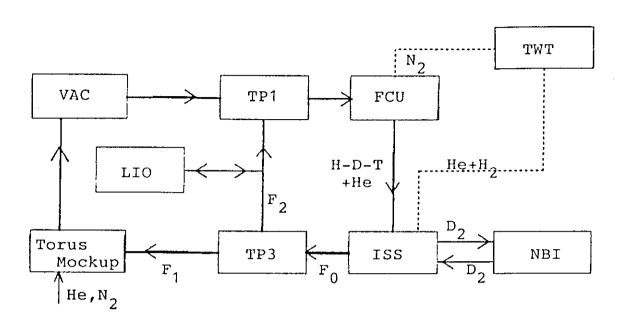
The dose from any tritium which escapes before the room is isolated has been neglected.

 $[\]frac{1}{3}$ Worker dose per minute of exposure. It is expected that personnel will exit from the room in less than 30 seconds. The calculations also assume uniform mixing in the room.

Table 1.4 Expected Tritium Inventory

SUBSYSTEM	COMPONENT	INVENTORY 1,2	FORM
VAC	a. Torus b. Cryopump	.01 g 6 g	DT DT
FCU	HMB's, MSB's DTOF	10 g to 30 g	T ₂ ,DT,DTO C(D,T) ₄ ,N(D,T) ₃
ISS	a. Total	100 g	T ₂ ,DT
UTB	a. TSTA Shutdown b. TSTA Normal	150 g 0.25 g	T ₂ ,DT
TWT	a. Low Pressure Receiver	0.1 g	DT,T_2,HTO,C_xT_y,H_2
	b. Molecular Sieve Drier	4-6 g	нто
XCS		0.1 g	т ₂ , нто
ETC	a. Normal Operation	small or none	UTO T O DTO
	<pre>b. After Spill of X g</pre>	X g	HTO,T ₂ 0,DT0

- 1. The inventory in the piping of TSTA is estimated at 1 g.
- 2. In some cases the amount of tritium is at its maximum in the system before regeneration or removal for disposal.



Main flow rate : $F_0 = 15 - 19 \text{ mol-DT/h}$

VAC flow rate : $F_1 = 2 - 10 \%$ of F_0

Circulating

flow rate : $F_2=90 - 98 \%$ of F_0

Fig. 1.1 TSTA Conceptual Block Diagram

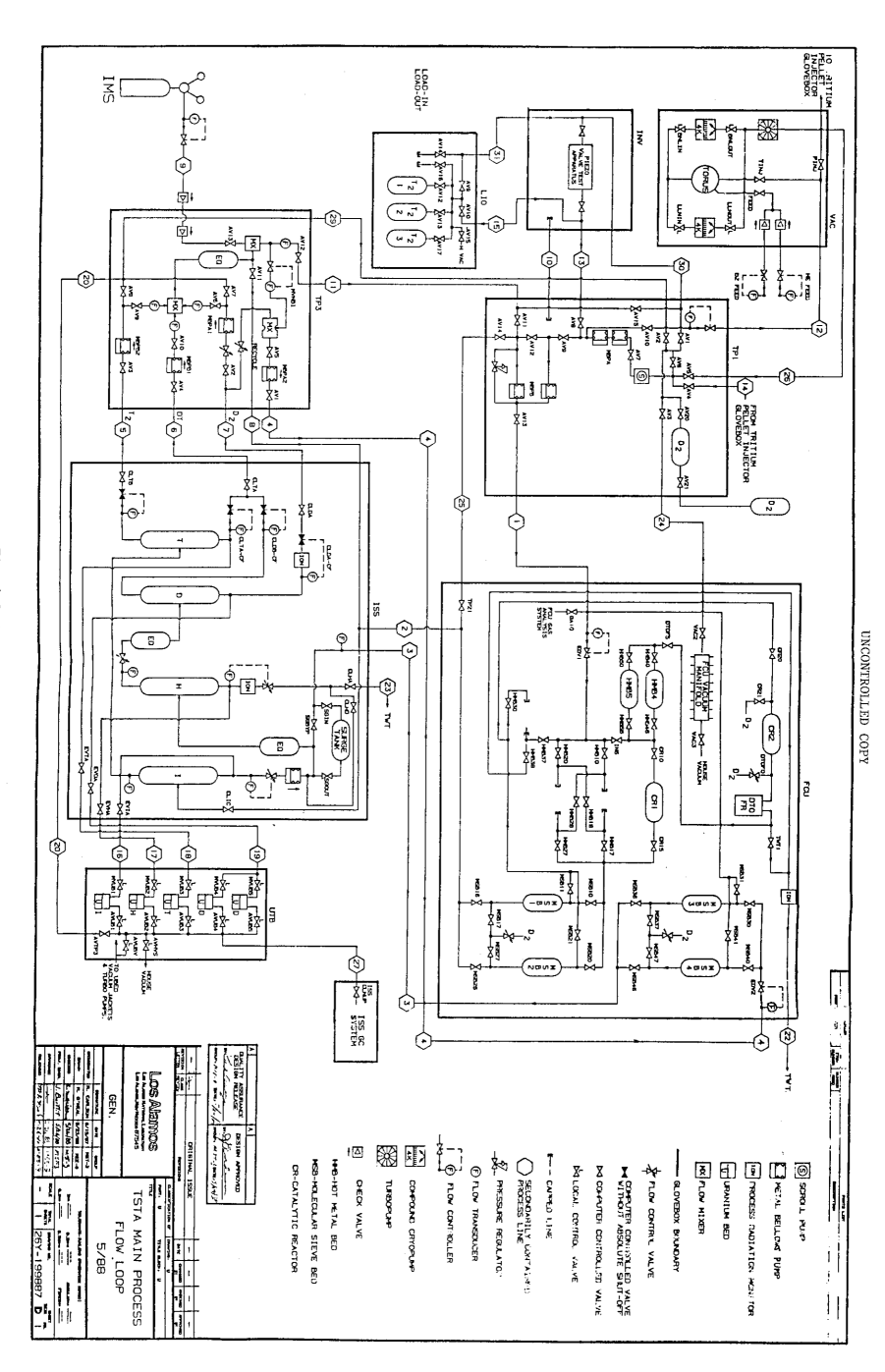


Fig. 1.2 TSTA Main Process Flow Loop

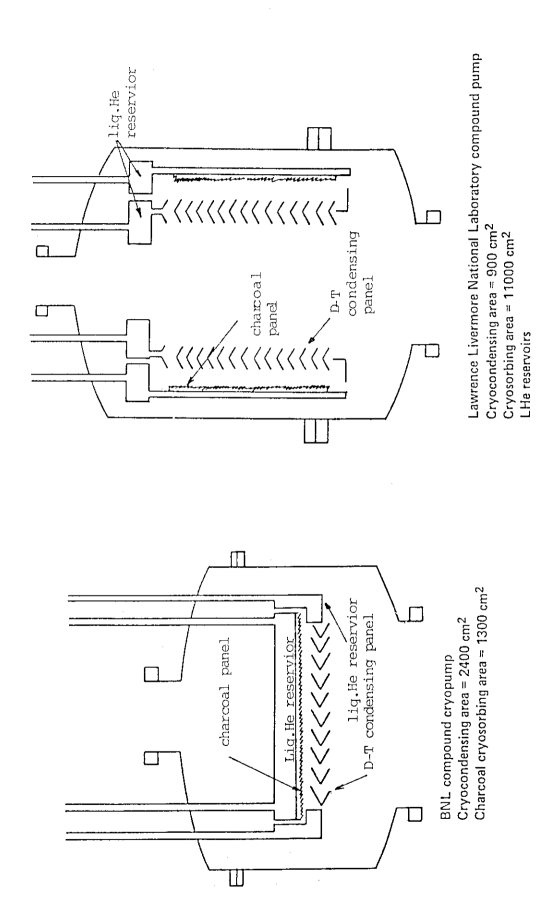


Fig. 1.3 BNL and LLNL Compound Cryopumps

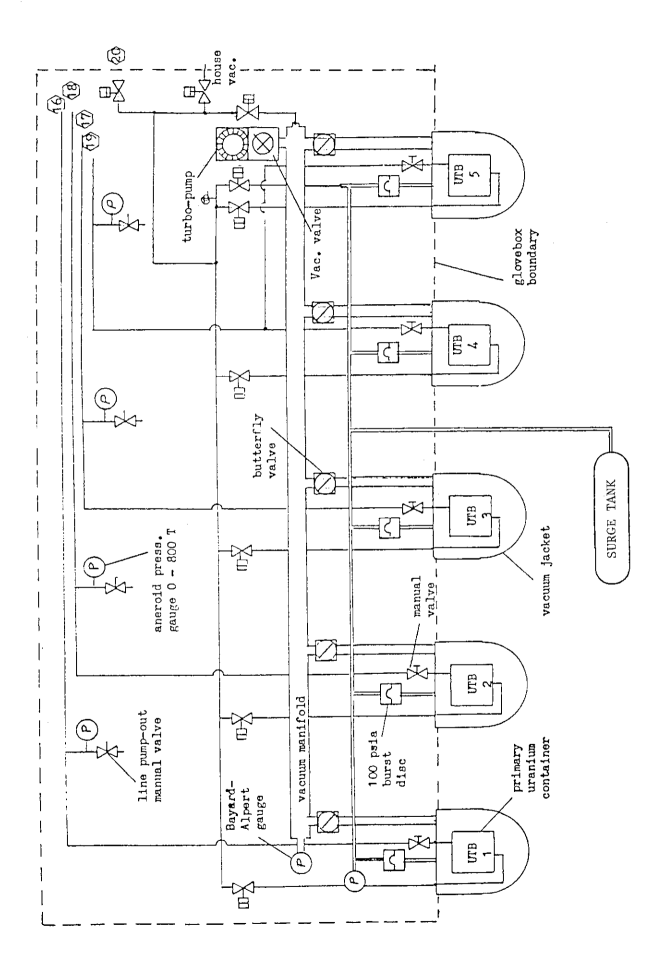


Fig. 1.4 Schematic Configuration of UTB system

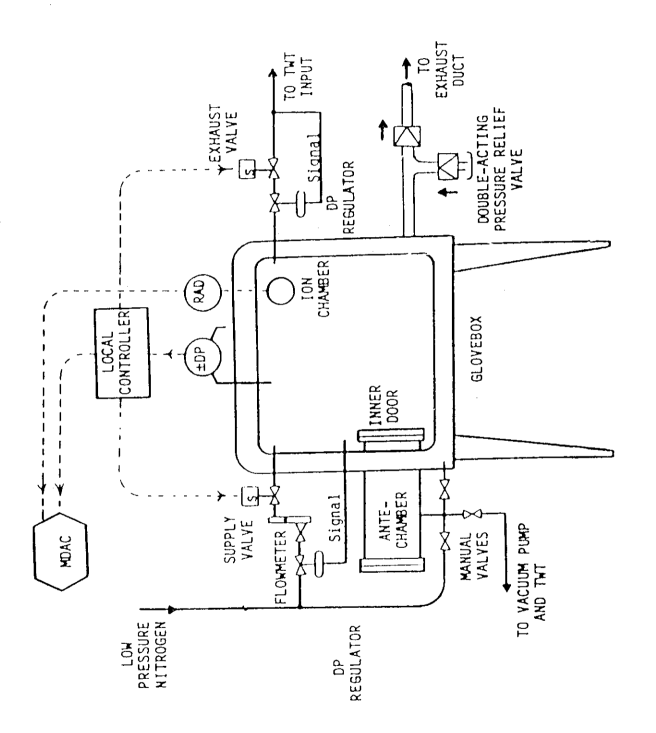


Fig. 1.5 Instrumentation of a Typical Glovebox

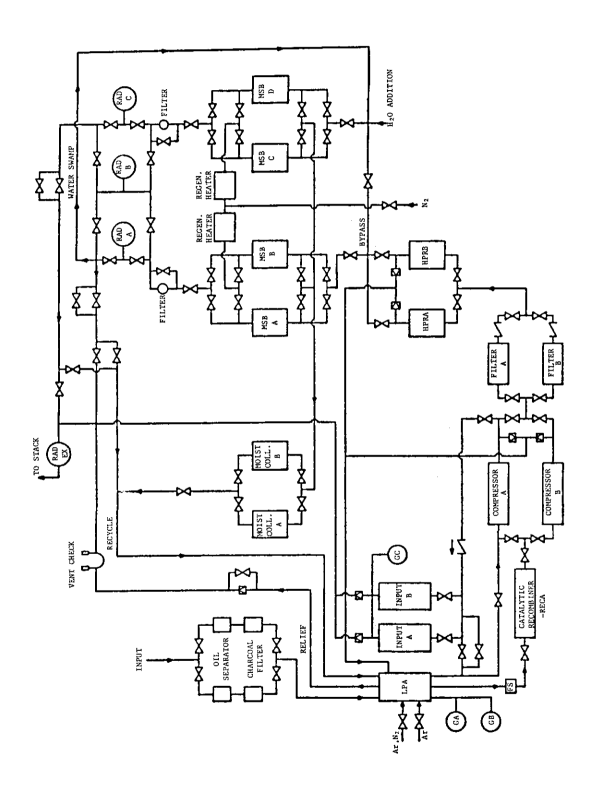


Fig. 1.6 TWT Primary Components and Flow Paths

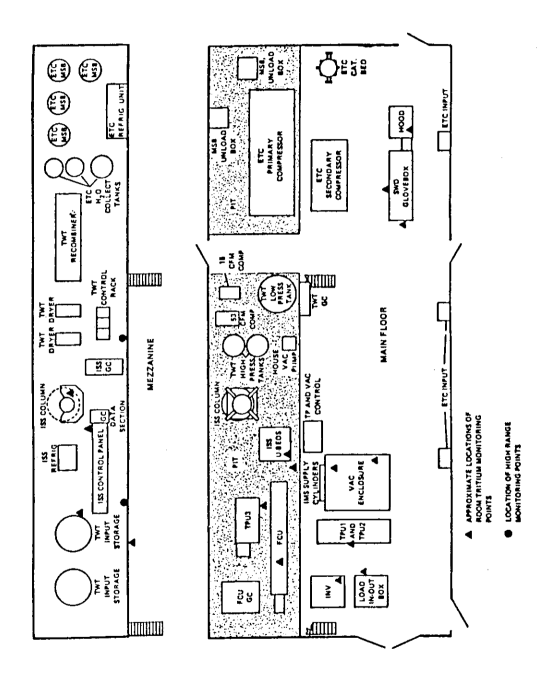


Fig. 1.7 Tritium Monitoring Points

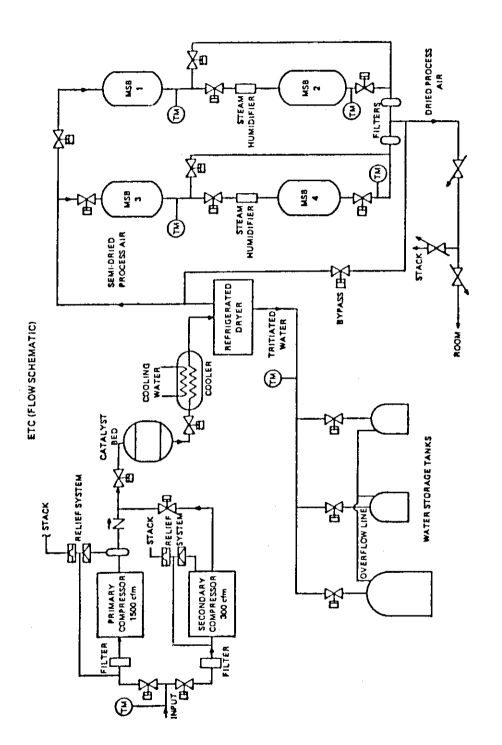


Fig. 1.8 ETC Primary Components and Flow Paths

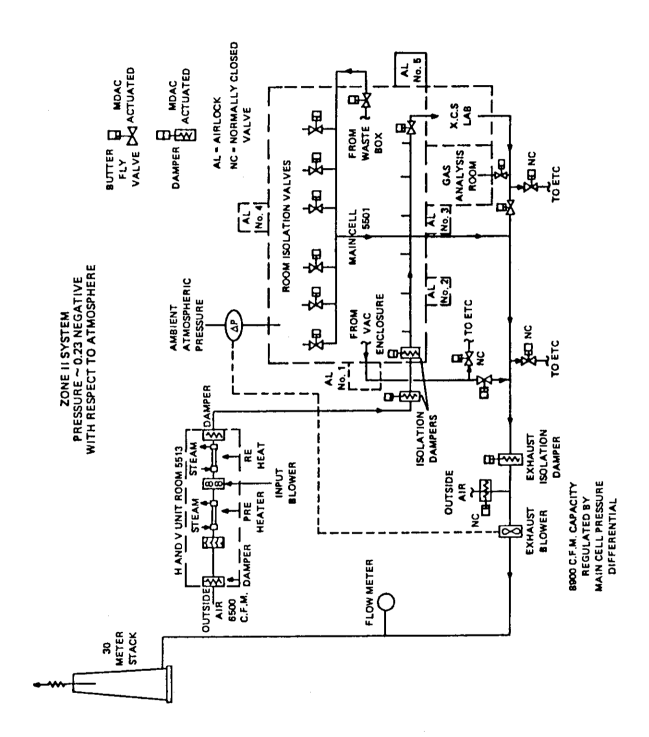


Fig. 1.9 Ventilation System of TSTA Facility (Zone II)

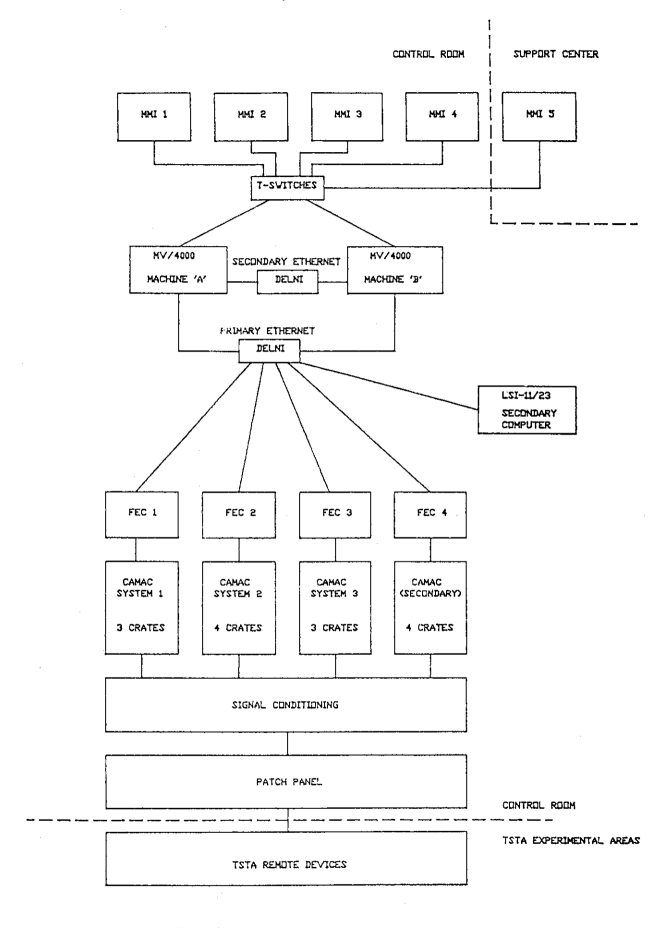


Fig. 1.10 General Block Diagram of MDAC

2. TEST PLAN

2.1 OBJECTIVES

The primary objectives of this milestone run were:

- to integrate the VAC system into the TSTA main process loop.
- to measure pumping speeds of a compound cryopump under different flow rates and impurity compositions (N₂, He).
- to measure D-T/He separation performance of a compound cryopump and demonstrate He and D-T regeneration from cryopanels to the process main loop,
- to improve the data-base and operating experience on both process systems (UTB, ISS, FCU, etc.) and safety systems (SEC, TM, TWT, etc.).

2.2 CONFIGURATION

The TSTA process loop is configured as shown in Figure 1.2. The new subsystem VAC for this test is enclosed within a secondary containment $(2.4x2.7x3.6\ \text{mH})$ backed up by ETC.

The system is able to operate in any of the following modes:

- Local,
- Computer manual,
- Computer macro,
- Full computer control.

The hydrogen isotope separation system however, was operated in the local and computer manual mode during this run.

Process systems used were: VAC, LIO, UTB, IMS, FCU, ISS, TP1, TP3 and GAN. All safety systems were required or available during the run, these were SEC, TM, TWT, VEN, ETC, MDAC, and the emergency generator and uninterruptible power supply. The following utility systems were required; the house vacuum system, breathing air, helium, nitrogen (liquid and gaseous) and hydrogen-argon. Liquid helium was supplied to the VAC system. The supplied breathing air system was available for use in the unlikely event of a tritium release into the main cell.

2.3 PROCEDURES

(1) Start up operation

Major steps for the start up of the process systems were:

- low level tritium leak checking of the main process

system,

- evacuation of the ISS vacuum jacket, pre-cooling of the ISS thermal shield with liquid nitrogen and cooling down of the columns with the cryogenic He refrigerator,
- evacuation of the vacuum jackets of FCU process components, and cooling down of cryogenic molecular sieve beds (no-heating the catalytic reactor because there was no impurity except N_2 in the main process loop).
- supply hydrogen isotopes to the FCU to pre-saturate the cryogenic molecular sieves (MSB1),
- evacuation of the vacuum jackets and cooling down of NBI return process component (cryogenic molecular sieve bed, MSB4),
- evacuation of the UTB vacuum jackets and pre-heating

Tritium loading and He stripping (2)

The loading of tritium into the process loop was performed from the UTB (except UTB-2) and from the tritium gas shipping containers attached to the LIO. Approximately 84.4 g-mole of H-D-T mixture(18.1g-H, 230.1g-D, 107.0g-T) was supplied from the UTB and tritium gas shipping containers in this run. The stripping of He and excessive H₂ was performed by using the ISS

column H during the full loop operation.

Loop operation (3)

Full loop operation through the flow paths FCU-ISS-TP3-TP1-(DT main flow) and ISS-TP3-FCU(NBI)-ISS (NBI D2 flow) was FCU

started after the completion of the tritium loading from the UTB and tritium gas shipping containers. Additional loading of tritium from an another gas shipping container was carried out after achieving a steady state conditions in the loop. tests of a compound cryopump were started when steady stat was reached.

The major tasks performed during loop operation were:

- to measure the characteristics of compound cryopumps of VAC (Table 2.1 shows the experimental condition of VAC), to measure H-D-T separation characteristics under
- steady state distillation,
- withdrawal of He, H_2 -HD, D_2 , DT and pure T_2 from the

ISS under continuous distillation conditions.

(4) Shutdown of the process loop

The ordinary shutdown procedures of the process loop are:

JAERI-M 88-242

- stop process gas circulation through flow paths FCU-ISS-TP3-TP1-FCU and ISS-TP3-FCU(NBI)-ISS,
- offload the H-D-T mixture in the process loop to the UTB,
- warm up the ISS by stopping He refrigerant and liquid nitrogen supply, and continue offloading of H-D-T mixture to the UTB,
- warm up cryogenic molecular sieve beds in the FCU and NBI return process by stopping liquid nitrogen supply.
- if blanketing effect occurs over the uranium beds in the UTB, circulate process gas through the flow path FCU-ISS-UTB-TP3-TP1,
- evacuate process residual gas(including impurities) in the process loop to empty tritium gas shipping containers by using the TP1,
- turn off all electrical heaters of process components and heaters in the ISS.

Table 2.1 Set Values for VAC Operation in the Integrated Run

	Ę	Our and the control of the	THE PART WATER OF THE PART OF		Ę		, F	
DATE	KON	EXFERIMENTS	TOTAL THROOGHFOI		ドーレエ		r-ne	ດງ
			TORR.L/S	SCCM	V-DC	COUNTS	SCCM	DIV.
2 June, Thu		* D-T + N_2 : 1 % $_2$ * Regen. to FCU	1x10min 3 " 5x30min 3x10min	78.95 236.85 394.75 236.85	0.801 2.191 3.581 2.191	53.5 59 59 59		
3 June, Fri	2	* Pure D-T * Regen. to FCU	1x10min 3 " 5x30min 3x10min	78.95 236.84 394.74 236.84	0.801 2.191 3.581 2.191	53 59 59 59		
3 June, Fri	E	* D-T/He: 2%He 5%He * Regen. 10%He to FCU 20%He	5×10min 5 " 5 "	386.86 375.01 355.28 315.80	3.511 3.407 3.233 2.886	•	7.90 19.74 39.48 78.95	
4 June, Sat	₹"	* D-T/He : 10%He 10%He * Regen. 10%He to LIO 10%He	1xlomin 3 " 5x60min 3xlomin	276.33 71.06 213.17 355.28 213.17 71.06	0.731 1.982 3.233 1.982 0.731	53 53 58 58 58 58	7.90 23.69 39.48 23.69	5.44 27.43 27.43 27.43
5 June, Sun 6 June, Mon	5 SHIJTD	5 * D-T/He : 10%He * Regen. to LIO SHJIDOWN LOOP OPERATION	1.5x120min	106.58	1.044	54.5	11.84	8.17
7-9 June, Tue-Thu	Thu	* Ion gauges(on TORUS, BNL, LINL)Calibration with D-T * D-T will be supplyed from UTB * Complementary runs will be performed with UTB and TP1.	S, ENL, LINL)Calibration with D-T ed from UTB will be performed with UTB and T	ation wit	h D-T and TP1.			

3. RESULTS OF LOOP OPERATION

Loop operation began at 9:23 AM on June 1, 1988. The following is the results of main subsystems during loop operation.

3.1 PROCESS SYSTEMS

3.1.1 VACUUM SYSTEM (VAC)

Torus pumping tests with the compound cryopump (BNL) were performed with D-T, N_2 , and He. The pump employed a charcoal panel as the helium sorbing media. The nominal pumping surface areas are 2400 cm 2 for DT condensing chevrons and 1300 cm 2 for the He sorbing panel.

The principal results were the following:

- the BNL compound cryopump was successfully integrated into the TSTA process loop and demonstrated with a D-T flow rate of 1 7% of D-T main stream,
- impurity effects on the pumping speed with a D-T stream was investigated with 1% N_2 and 1-25% He, demonstrating little effect,
- He/DT separation performance on both panels of BNL pump was measured: DT gas regenerated from the condensing chevron revealed no He contained, He gas from the charcoal panel showed the level of DT of 10-25% DT

 $(8x10^{-4} - 4x10^{-3} \text{ g-mol/h}).$

The detailed results and discussion of this system will be described in a separate report.

3.1.2 TRITIUM STORAGE AND SUPPLY SYSTEM (UTB)

Approximately 84.4 g-mole of H-D-T mixture (18.1 g-H, 230.1 g-D, 107.0 g-T) was loaded to the process system from four uranium beds (UTB-1, -3, -4, and -5) in the UTB and from tritium gas shipping containers. Hydrogen isotopes from UTB were loaded directly to ISS during cooling down with the cryogenic He refrigerator, and tritium gas from gas shipping containers was supplied through the FCU.

Figures 3.1 and 3.2 show temperatures of the uranium beds (each contains 6 kg-U). Figure 3.3 shows the pressure of column I in the ISS during offloading of the UTB. The required time periods to heat the UTB from 473 K to 700 K were approximately 4 hrs, and two days to cool to ambient temperature.

3.1.3 FUEL CLEANUP SYSTEM (FCU)

 $\ensuremath{\text{No}}$ impurity except $\ensuremath{\text{N}}_2$ was added into the main loop in this run, so the catalytic reactors of FCU were not heated (Figure

Figure 3.5 shows the temperatures of molecular sieve beds MSB1 in the FCU and MSB4 in the FCU (NBI). Nitrogen gas used for the VAC experiments was removed by this molecular sieve bed (MSB1) during the regeneration of the BNL cryopump.

Figures 3.6(a) and 3.6(b) show the pressures in the main

process line of FCU during this run.

3.1.4 ISOTOPE SEPARATION SYSTEM (ISS)

This system (Figure 3.7) was operated with the FCU and NBI for 140 hrs. Withdrawals of He, H, and HD from the top of column

H, D_2 from the top of column D, and T_2 from the bottom of column

T were performed under full loop operation.

Figures 3.8(a) and 3.8(b) show typical distillation conditions of four interlinked columns during the removal of He to TWT and steady state, respectively. The concentration profile was calculated under the condition of steady state shown in Figure 3.8(b), using 5 cm of HETP (Height Equivalent to a Theoretical Plate). Figures 3.9(a) and 3.9(b) show the profile of column I in vapor and liquid phase, respectively.

The mixture of He, Ho and HD was withdrawn four times from

the top of ISS column H to the TWT. The following is the log of these withdrawals.

First time : Time 13:10 - 17:30 Flow rate 280.0 scc/min Total 72.8 I-STP

No tritium was detected by GC analyses.

Second time: Time 19:52 - 21:56 (6/3/88)Flow rate 282.0 scc/min Total 35.0 1-STP

No tritium was detected by GC analyses.

Third time: Time 19:10 - 21:50 Flow rate 280.0 scc/min (6/4/88)Total 44.8 1-STP

No tritium was detected by GC analyses.

Time 20:19 - 20:50; 20:50 - 21:56 Flow rate 280.0 ; 309.0 scc/min Forth time: Time (6/5/88); 309.0 scc/min Total 29.1 1-STP

No tritium was detected by GC analyses.

During these withdrawals, no increase of tritium level the TWT-LPR was observed except the forth withdrawal (maximum level: 1 Ci/m^3) in which period ISS was not in a good steady state conditions.

Approximately 60 samples of gas analysis were performed with the on-line gas chromatograph system to certify a steady state condition of each column. Tables 3.1(a)-(c) show the results of the analysis.

Figures 3.10(a), 3.10(b), 3.11(a) and 3.11(b) show the pressures and the liquid levels of ISS, respectively.

Figure 3.12 shows the inlet flow rate of column I and compositions at the top of column H. No tritium was detected by gas chromatograph at the top of column H during the entire period of this run.

Figure 3.13 shows the temperatures of columns during cooldown. The required time periods of cooling down to 20K from ambient temperature were approximately 9.5hrs for columns I and T (the packed section of both columns is equipped with He looping coils), 11hrs for column D and 26hrs for column H. It took much time to cool down column H because the concentration of He in column H was high.

Figures 3.14(a) and 3.14(b) show the temperatures of columns during warm-up. The required time to warm up from 20 K to ambient temperature was approximately 50 hrs.

3.2 SAFETY SYSTEMS

3.2.1 SECONDARY CONTAINMENT SYSTEM (SEC)

Notable increments of the radiation levels in gloveboxes were observed at ISS (GB1 and GB2) and LIO. The following are the characteristics of these tritium releases in each glovebox.

(1) Radiation levels of ISS gloveboxes

Figure 3.15 shows the radiation levels of two ISS gloveboxes and UTB glovebox.

The GB1 radiation level (on 31 May) started to increase

(from 1.0×10^{-3} Ci/m 3 to 2.5×10^{-2} Ci/m 3) with the increase of ISS column I pressure during loading tritium from two uranium beds (UTB1 and 3). In the previous loop experiments [Refs.1 and 2], this typical increase has been identified as due to a leak through a rupture disc assembly on the top of column I. The radiation levels decreased gradually with the progress of isotopic separation between light elements and tritium in the column I (decrease of tritium concentration in the top of this column). Purge of this glovebox had automatically been performed

at a level higher than $1x10^{-3}$ Ci/m³. The second peak (maximum

 4.0×10^{-2} Ci/m³) shown in this Figure, was also caused by the increase of tritium at the top of column I due to the disturbance of distillation conditions of the four columns.

Figure 3.16 shows a relationship among radiation levels of ISS gloveboxes and the pressure of ISS. A relationship between

radiation levels and pressure of the column H can be seen from this Figure.

Most peaks on the ISS-GB2 were caused by ISS gas analysis (approximately 60 samples were withdrawn from columns to ISS.GAN during this run). The increase of the GB2 radiation level with an increase of the ISS pressure occurred from unidentified small leaks in the plumbing of the ISS gas analysis system (sampling manifold for two gas chromatographs). This glovebox was always purged with nitrogen during this run, since the tightness of isolating valve sheets between the sampling manifold and the ISS columns has deteriorated.

(2) Radiation levels of FCU glovebox

There were no leaks in the glovebox of FCU during this run.

(3) Radiation levels of LIO and INV gloveboxes

As shown in Figure 3.17, five peaks (maximum; $8x10^{-3}$ Ci/m³) on the level of LIO-GB were observed. This normal release occurred when a tritium gas shipping container was attached to the LIO manifold. The source of tritium might be some residual gas in a short connecting tube on the gas cylinder. Successive leaks did not occur during loading and offloading.

(4) Radiation levels of TPU glovebox

There appeared no notable radiation peak in TPU-GB1 and GB2 (Figure 3.18). They contain the TPI and TP3 systems composed of two sets of double-headed metal bellows pumps, a scroll pump (in TP1) and a hydrogen isotope equilibrator (in TP3).

3.2.2 ROOM RADIATION LEVELS

No offnormal radiation levels were detected with tritium monitors placed at several key places in the TSTA experimental room (main cell) during this run. Figures 3.19, 3.20 and 3.21 show the radiation levels in main cell. The sharp peaks were due to the daily check of detectors.

3.2.3 STACK RADIATION LEVELS

No offnormal release were detected during this run (Figure 3.22).

The total environmental release measured with the stack monitor (bubbler system)[Ref.1] was evaluated as follows:

5/30/88 - 6/5/88 HTO: 973.0 mCi

HT : 62.5 mCi

3.2.4 TRITIUM WASTE TREATMENT SYSTEM (TWT)

The tritium waste treatment system operated the entire period of this run without any problems. Figure 3.22 shows the radiation levels at the exit of TWT and the stack. During this run, two peaks were detected on the radiation level at the exit of TWT. No off-normal release to the environment was observed, because the TWT system went into recycle mode from stack mode, automatically. The evacuation of the main loop caused the first peak on the radiation level of TWT shown in Figure 3.22. The main remaining gas was hydrocarbon.

Figure 3.23 shows the radiation level in the low pressure receiver (LPR) of TWT. Although there are many tritium sources contributing to the LPR radiation level, most peaks observed in Figure 3.23 were due to exhaust gas from the ISS gas analysis system (ISS.GAN).

Figure 3.24 shows the temperatures of the TWT recombiner during this run. The temperature of about 760 K was reached under recycle mode and about 720 K was reached under stack mode.

It is important to keep the pressure in the LPR between set points because the LPR receives the exhaust gases from process systems and safety systems. During this run, the pressure in the LPR was successfully kept between 0.27 atm and 0.54 atm (Figure 3.25).

It is possible to estimate the total amount of tritium transferred to the TWT. The following is the breakdown of tritium exhausted from the ISS-GAN based on the composition, pressures and number of samples analyzed with the ISS gas

analysis system, and holdup of its sampling manifold (40 cm³):

Gas composition : Table 3.1(a)-(c)
Gas pressures : 600 torr for sampled gas

Gas pressures : 600 torr for sampled gas 300 torr for flushing

Number of sampling : 60 times

Number of flushing: 2 times in each sampling

Total tritium amount: 1871 Ci

The integrated amount of tritium was determined from the system flow rate (0.425 m 3 /min) and the reading of the LPR radiation monitor every minute. The total amount of tritium was 5.8×10^3 Ci, which includes a background contribution (1x10 3 Ci) of the LPR tritium monitor (background radiation level; 2.7×10^{-4} Ci/m 3).

Table 3.1(a) Results of GC Analysis for This Run (I) (Experiments have been performed in June '88.)

day/time	position	Не	^H 2	comp HD	osition HT	(%) D ₂	DT	т ₂
1/16:38	H-T	11.75	52.08	35.64	_	0.53	_	_
1/17:15	H-T	8.88	48.23	42.13	-	0.76	-	-
1/17:53	H-T	5.02	53.45	37.61	-	3.92	_	-
1/19:13	I-F	0.12	1.42	12.39	5.60	32.92	38.35	9.20
1/20:54	H-T	2.61	64.62	32.36	-	0.41	-	-
1/21:29	H-T	1.29	72.90	25.51	-	0.31	-	-
1/22:07	I-F	-	0.26	6.89	2.03	48.83	36.58	5.42
2/15:16	I-F	-	-	-	-	38.71	48.47	12.72
2/16:29	I-T	-	-	2.90	0.24	96.85	-	-
2/17:14	I-S(S1)		0.82	11.61	2.35	54.74	28.59	1.88
2/18:10	I-S(A2S)	-	-	0.72	0.12	69.85	29.11	0.11
2/18:16	I-S(S1A)	-	-	-	-	52.14	44.53	3.33
2/18:51	I-B	-	_	-		19.15	61.72	19.13
2/19:07	T-T	-	· -	_	-	26.96	60.45	12.60
2/19:16	T-S(S3)		-	-	-	7.91	44.80	47.29
2/19:41	T-S(S3A)	-	-	-	-	0.24	6.73	93.02
2/20:03	T-S(S3B)	-	-	-	-	-	0.18	99.82
2/20:08	T-B	-	-	-	-	-	-	99.99

Table 3.1(b) Results of GC Analysis for This Run (II) (Experiments have been performed in June '88.)

day/time	position	Не 	^H 2	compo	sition HT	(%) D ₂	DT	^T 2
2/20:34	H-T	6.10	38.49	53.91	-	1.45	-	_
2/21:04	H-S(S2)		0.42	10.21	2.63	80.79	5.93	_
2/21:08	H-S(S2A)	1.12	5.71	22.96	7.08	31.24	28.15	3.74
2/21:31	H-B	-	-	-	_	91.93	7.89	0.19
2/21:58	D-F	-	_	0.09	-	91.46	8.30	0.16
2/22:05	D-T		-	0.12		99.83	-	_
2/22:34	D-S(S4)	-	0.07	3.82	-	79.95	15.50	0.66
2/23:06	D-B	-	-	-	-	18.63	55.75	25.62
2/23:25	H-F	0.07	0.19	2.24	-	89.49	7.79	0.22
3/00:03	T-T	-	-	_		46.13	51.30	2.57
3/00:35	T-T		-	-	-	50.59	48.10	1.31
3/01:12	T-T	-	-	-	-	39.03	57.69	3.29
3/01:36	T-T	-	<u>.</u> .	_	_	32.34	62.61	5.06
3/02:09	$\mathbf{T} - \mathbf{T}$	-	-	_	-	25.17	67.58	7.24
3/02:36	T-T	_	_	_	_	23.12	63.07	7.81
3/03:09	T-T	-	-	~	-	23.30	67.81	6.89
3/03:42	T-T	_	-	-	-	30.57	63.69	5.74
3/04:15	T-T	-	-	-	_	35.84	59.78	4.38
								

Table 3.1(c) Results of GC Analysis for This Run (III) (Experiments have been performed in June '88.)

day/time	position	Не	^Н 2	compo HD	sition HT		DT 	^Т 2
3/17:37	H-T	2.10	97.33	0.57	_	-	-	-
3/18:10	H-T	6.68	93.32	-	-	_	-	-
3/20:57	H-T	4.35	95.23	0.41	-	-	-	_
3/21:30	$\mathbf{T} - \mathbf{T}$	-	-	-	-	27.85	65.85	6.30
3/22:00	T-T	-	-	_	-	23.39	70.05	6.56
3/22:30	T-T	-	-	-	-	25.08	67.69	7.23
3/22:56	T-T	-	-	-	-	29.54	65.42	5.04
3/23:50	T-T	. -	-	-	-	35.79	56.77	7.44
4/08:00	I-F			•	-	54.07	39.44	6.49
4/09:11	I-F	-	-	-	_	52.44	40.70	6.86
4/10:12	H-T	3.42	34.57	59.92		2.10	-	-
4/19:00	H-T	15.03	35.92	48.01	-	1.05	_	-
4/20:34	H-T	4.49	37.69	56.68	_	1.13	-	-
4/21:14	$H-\mathbf{T}$	3.15	30.93	64.62		1.30	-	_
4/22:01	H-T	2.19	25.03	70.81	-	1.97	_	-
5/08:39	I-F	-	-	-	-	44.95	45.06	9.99
5/18:34	H-T	2.43	9.17	83.11	-	5.28	· •	_
5/20:20	H-T	2.79	12.72	77.66	-	6.84	-	-
5/21:02	H-T	4.57	7.47	85.57	-	2.38	-	-
5/21:42	H-T	4.22	3.98	89.35	-	2.45	-	-

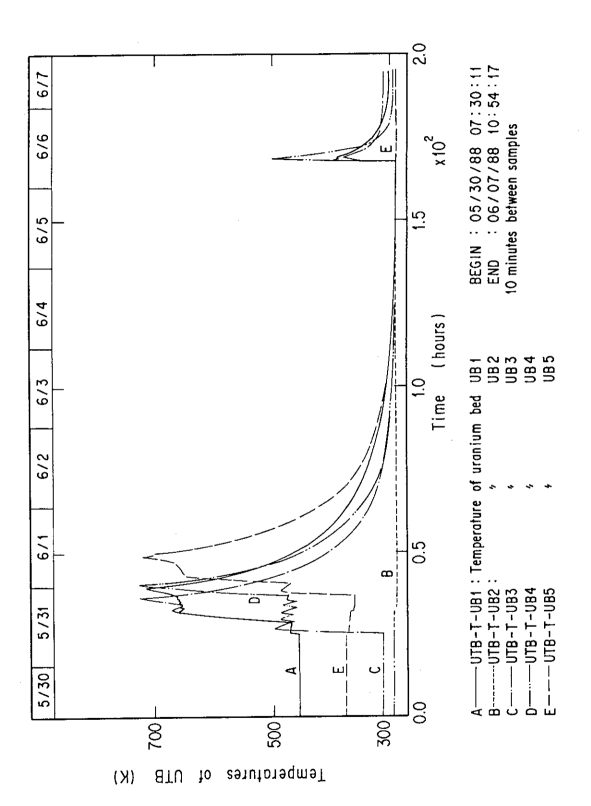


Fig. 3.1 Temperatures of UTB (1) at Tritium Loading to the TSTA Loop

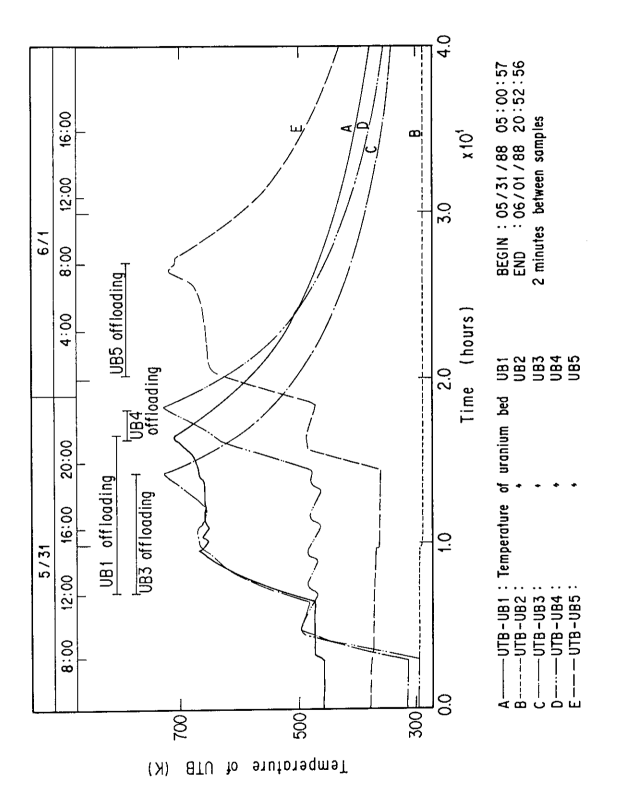


Fig. 3.2 Temperatures of UTB (2) at Tritium Loading to the TSTA Loop

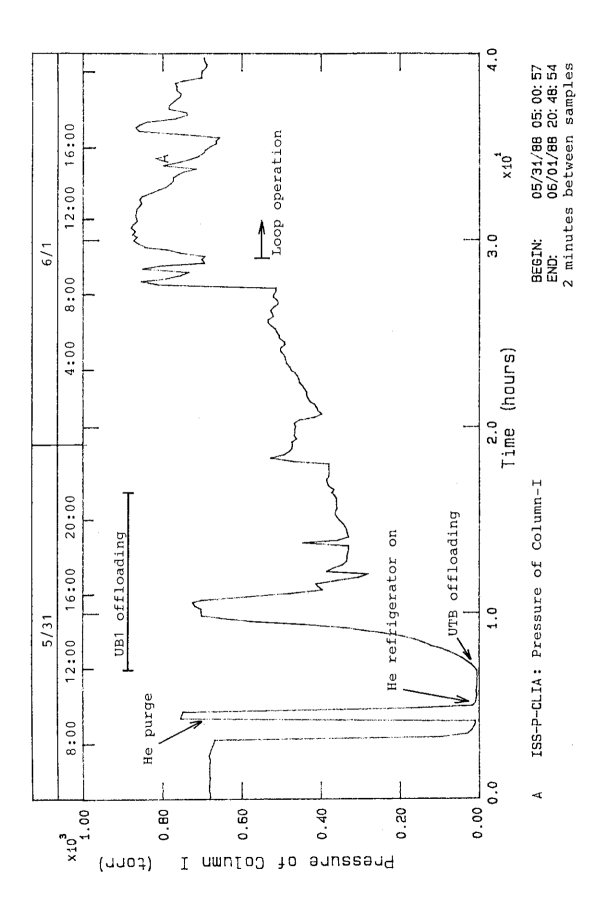


Fig. 3.3 Pressure of Column I during UTB Offloading

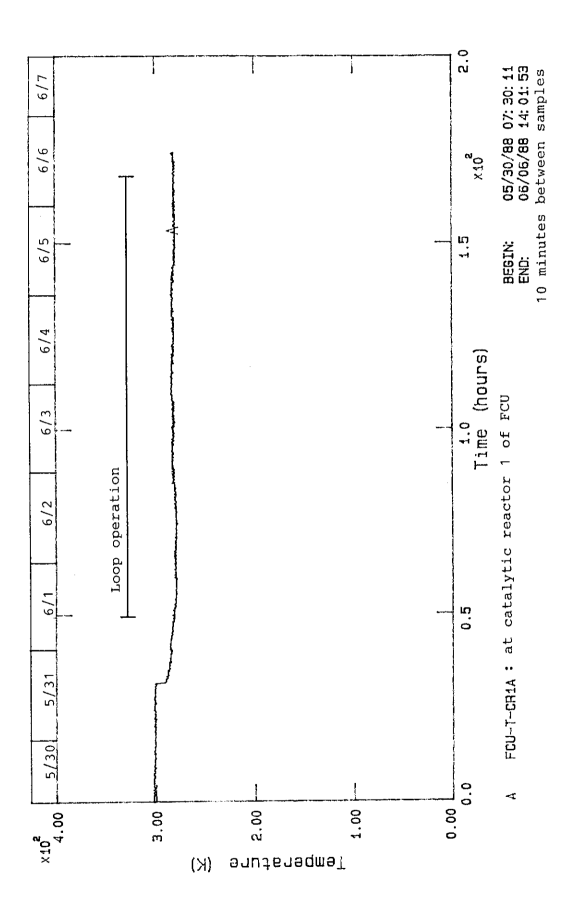


Fig. 3.4 Temperatures of Catalytic Reactor 1 in FCU

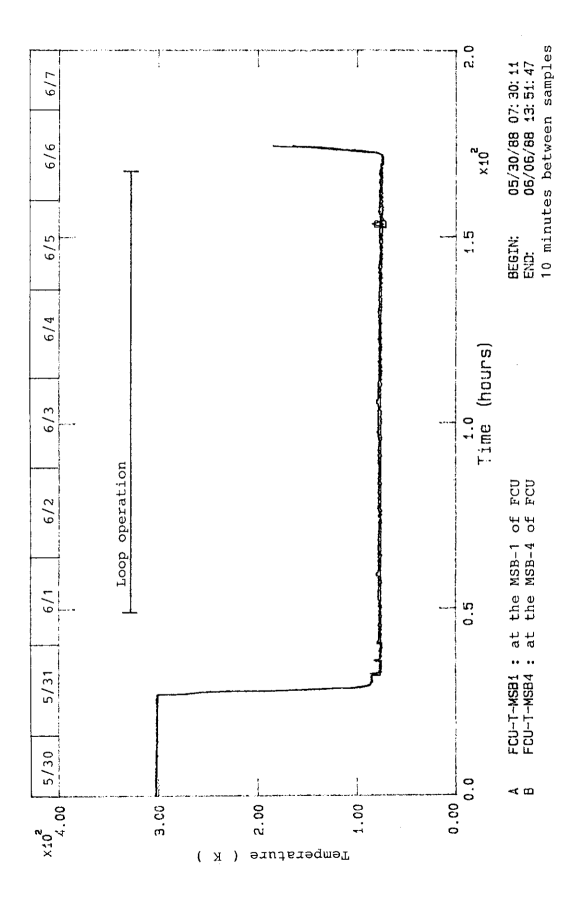


Fig. 3.5 Temperatures of Molecular Sieve Beds in FCU and NBI

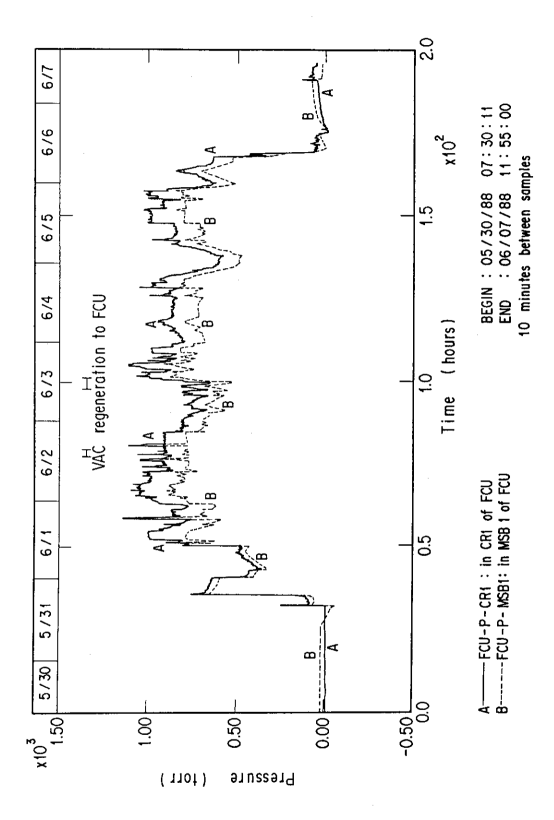


Fig. 3.6(a) Process Pressures in FCU

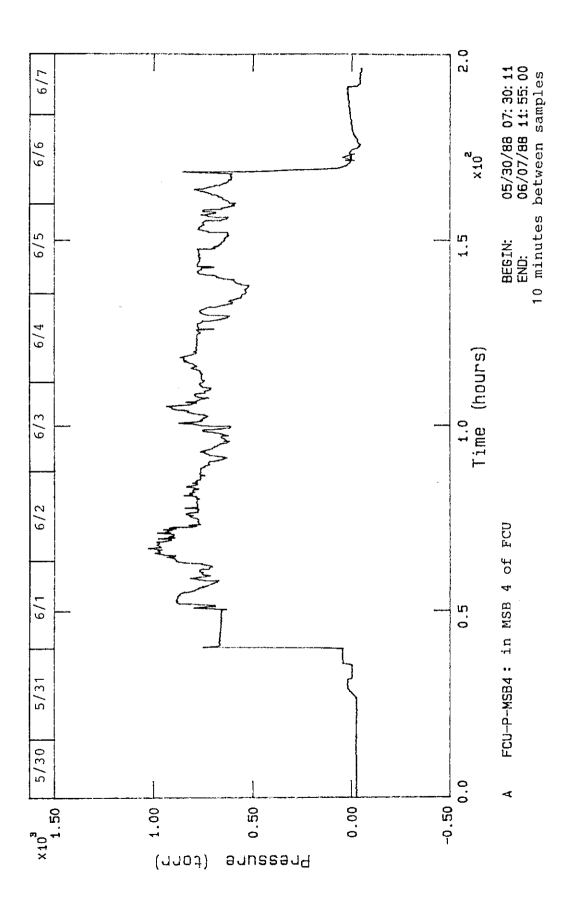
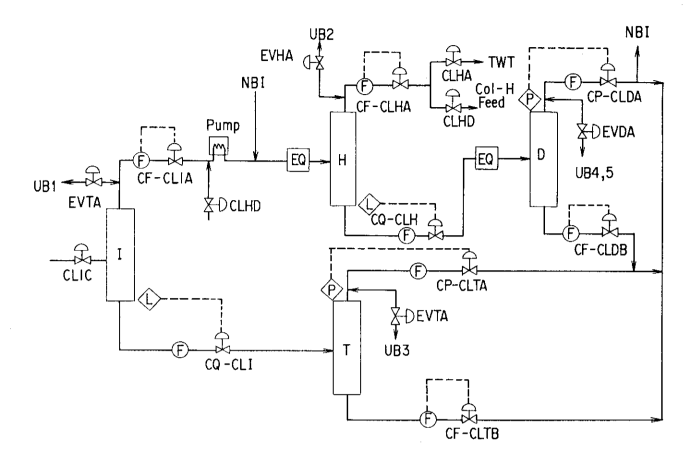


Fig. 3.6(b) Process Pressures in NBI



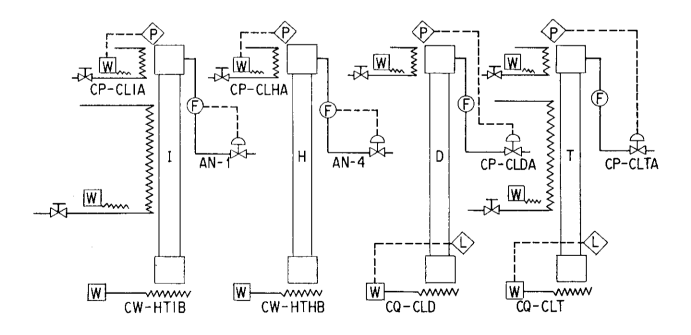


Fig. 3.7 Configuration of ISS Control System

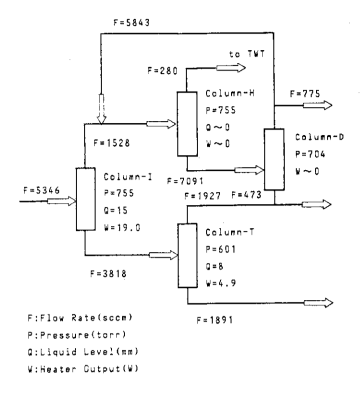


Fig. 3.8(a) Flow Balances in ISS (1)

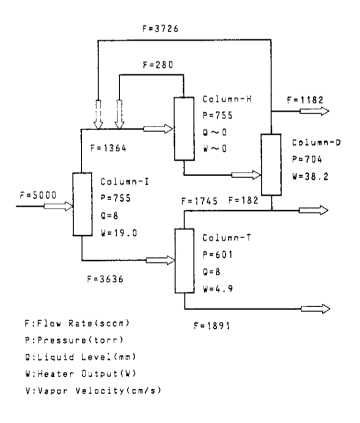


Fig. 3.8(b) Flow Balances in ISS (2)

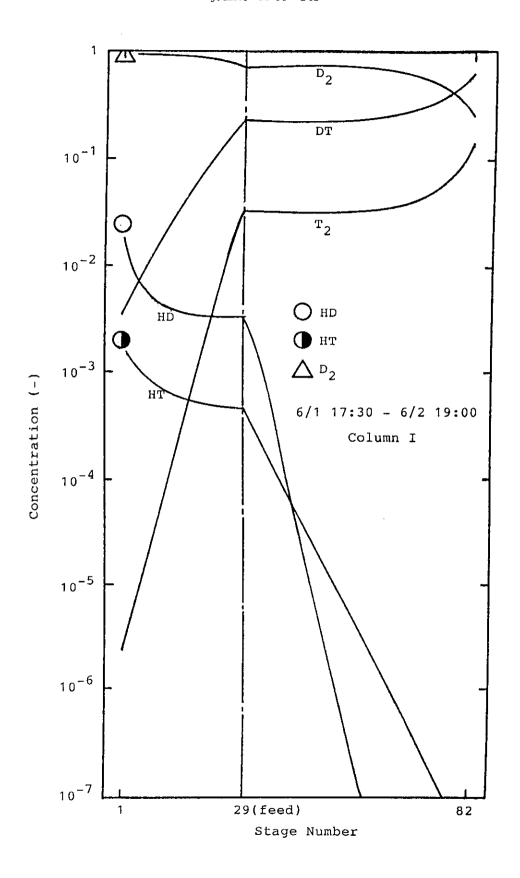


Fig. 3.9(a) Concentration Profile of ISS Column I (in Vapor Phase)

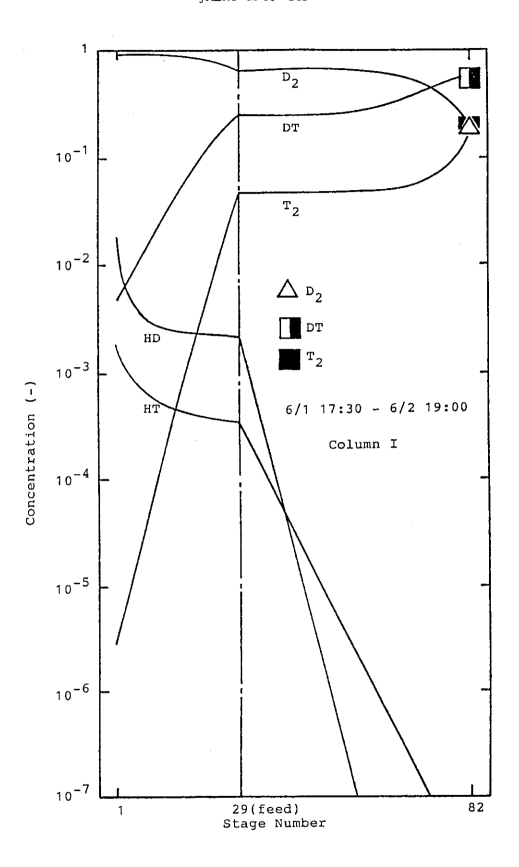


Fig. 3.9(b) Concentration Profile of ISS Column I (in Liquid Phase)

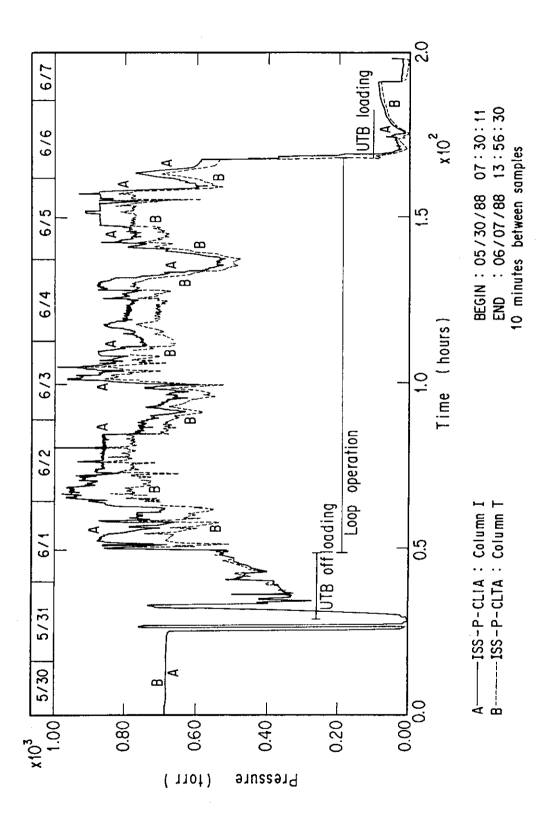


Fig. 3.10(a) Pressures of ISS Columns

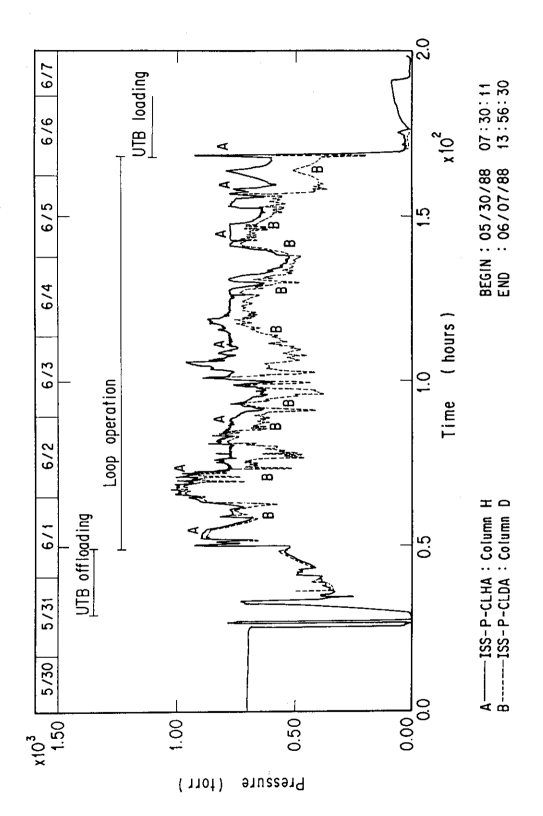


Fig. 3.10(b) Pressures of ISS Columns

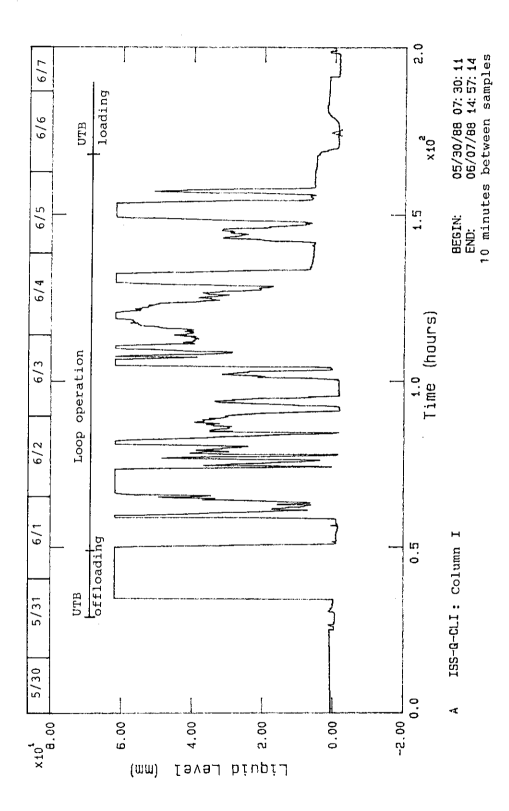


Fig. 3.11(a) Liquid Levels of ISS Columns

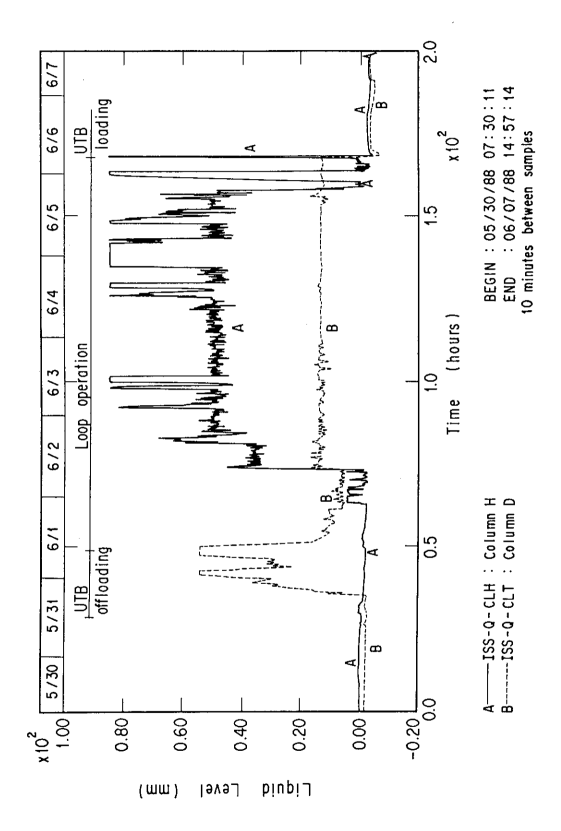


Fig. 3.11(b) Liquid Levels of ISS Columns

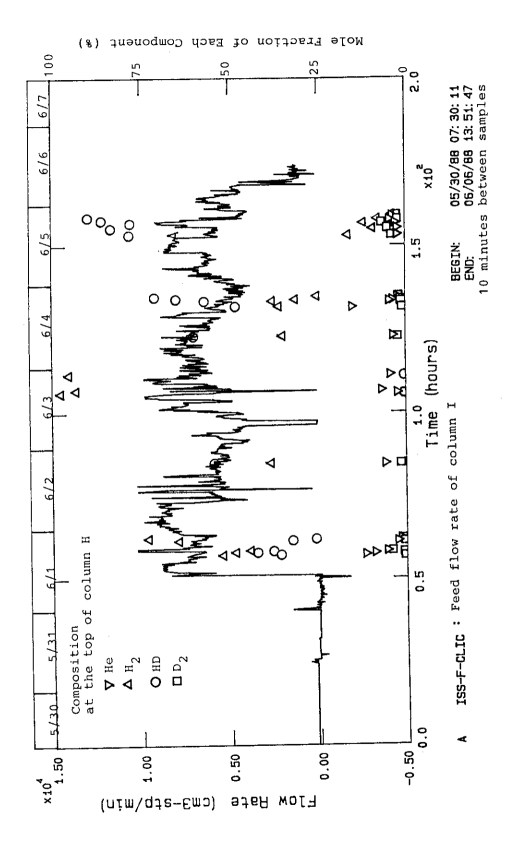


Fig. 3.12 Feed Flow Rate for ISS

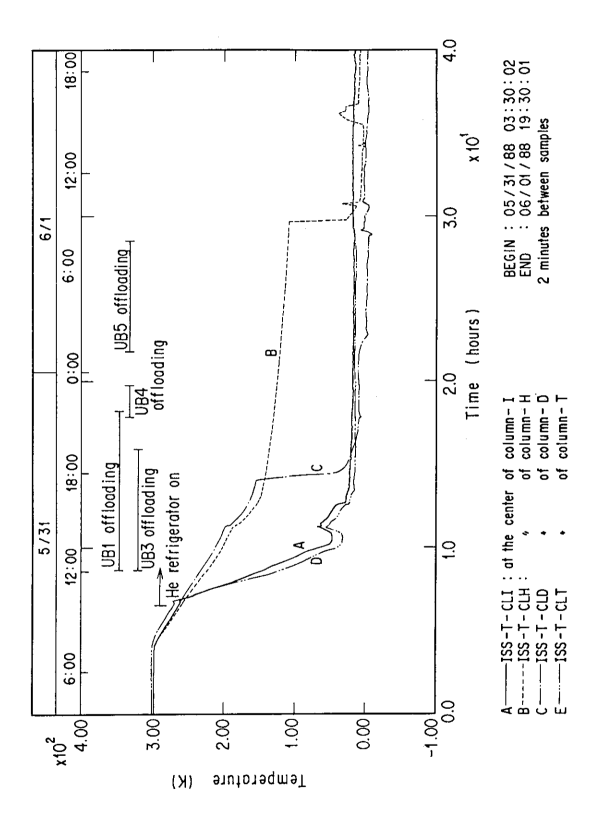


Fig. 3.13 Temperatures of ISS Columns during cool down

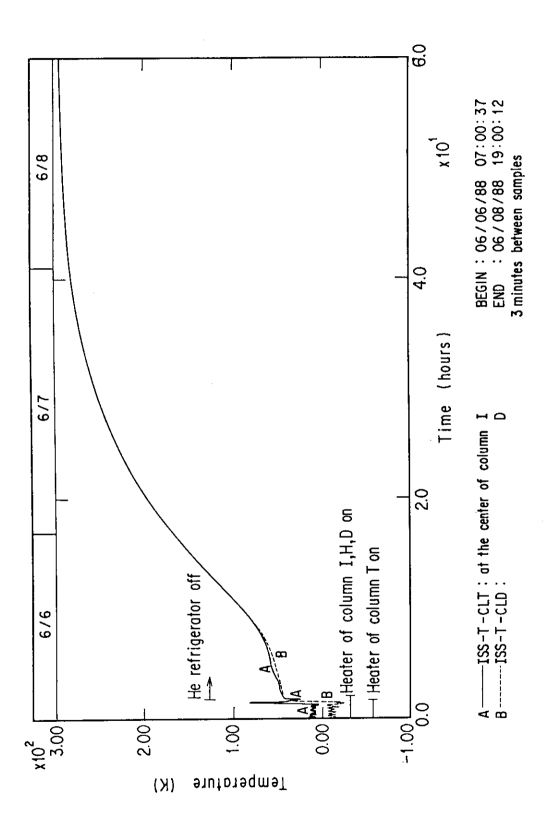


Fig. 3.14(a) Temperatures of ISS Columns during warm up

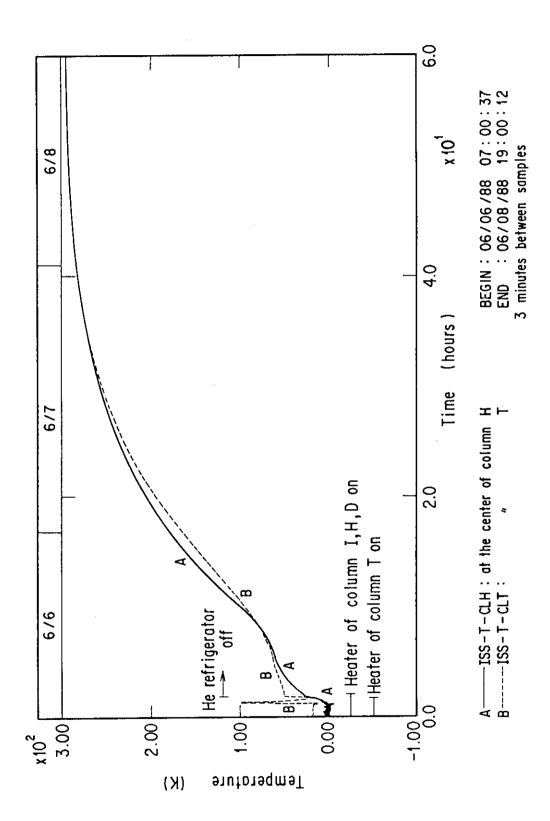


Fig. 3.14(b) Temperatures of ISS Columns during warm up

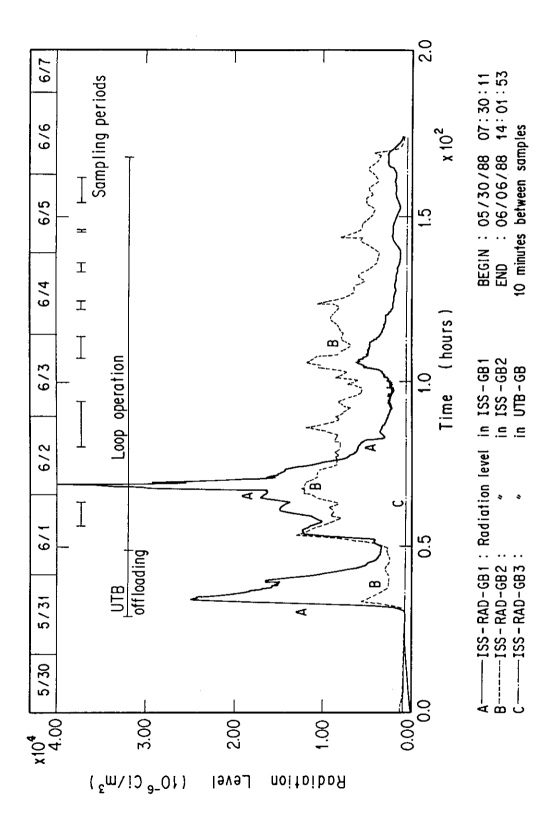


Fig. 3.15 Radiation levels in ISS Gloveboxes

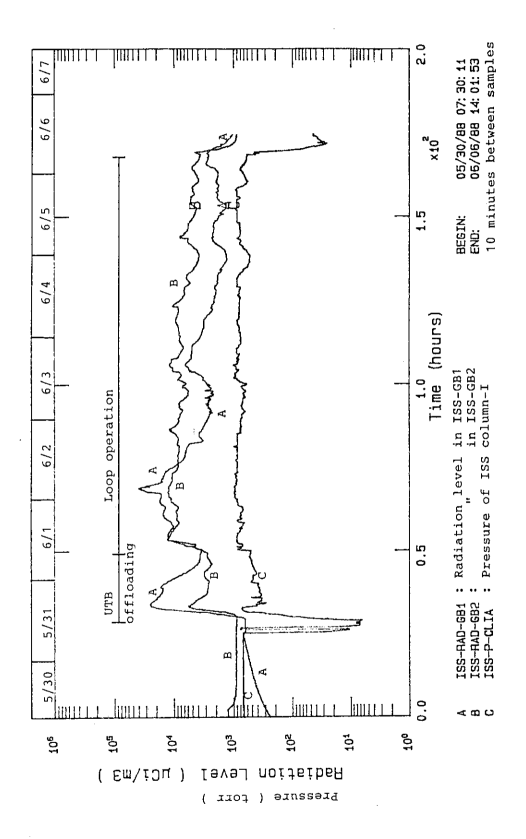


Fig. 3.16 Relationship between Radiation Level and Pressure in ISS

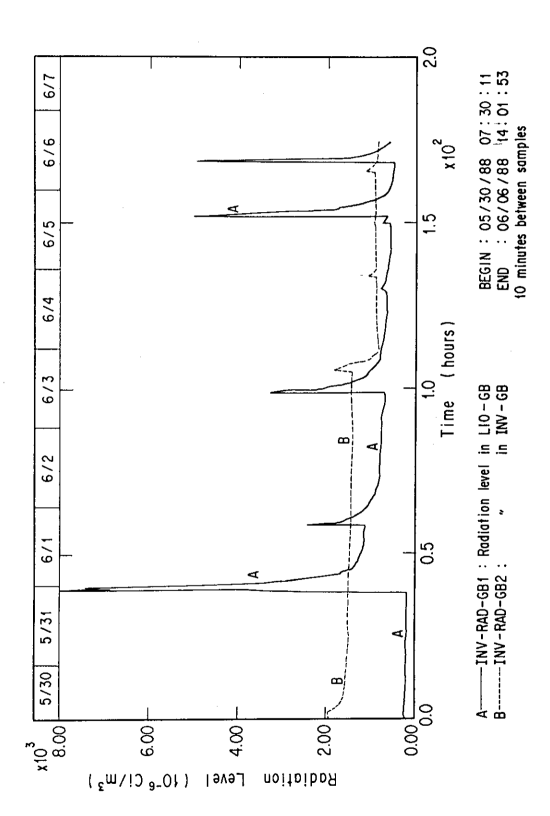


Fig. 3.17 Radiation Levels in LIO and INV Gloveboxes

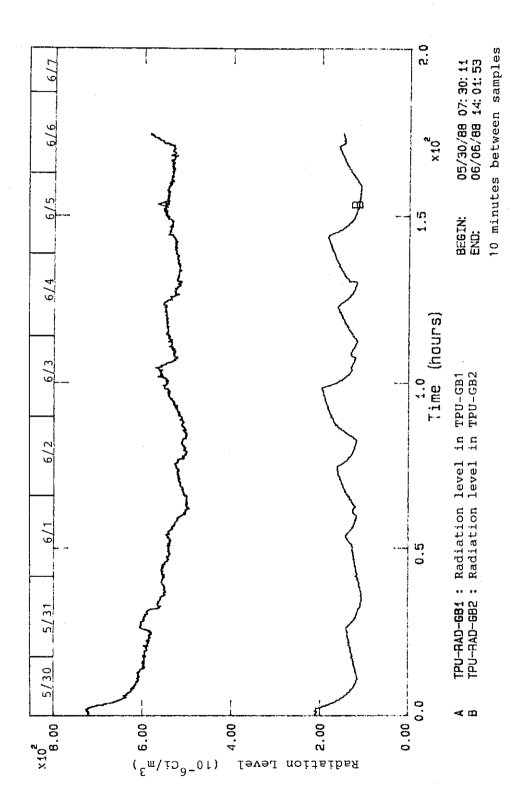


Fig. 3.18 Radiation Levels in TPU Gloveboxes

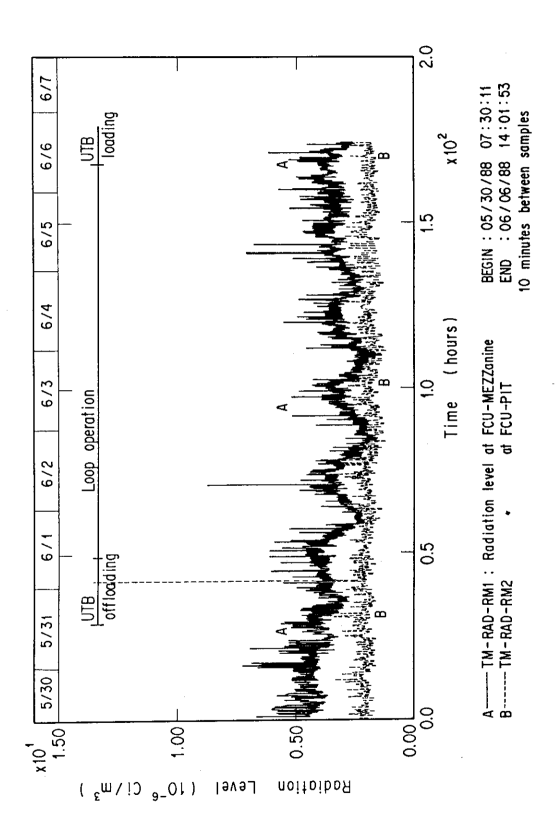


Fig. 3.19 Radiation Levels in Main Cell (1)

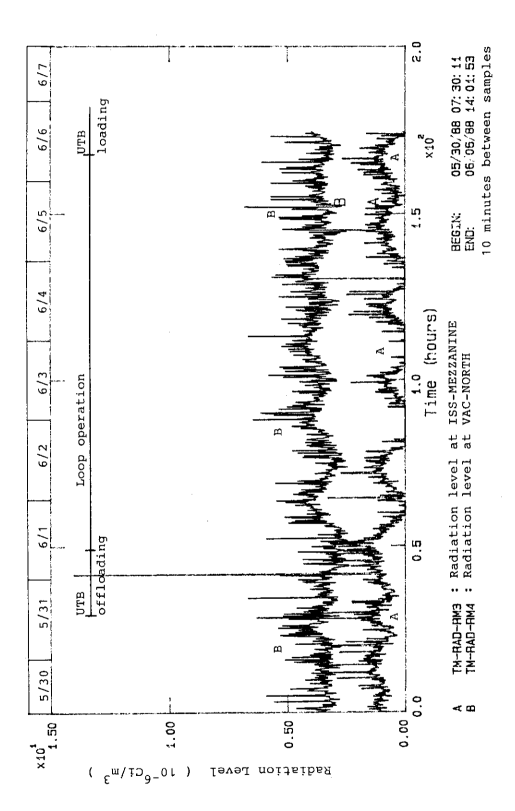


Fig. 3.20 Radiation Levels in Main Cell (2)

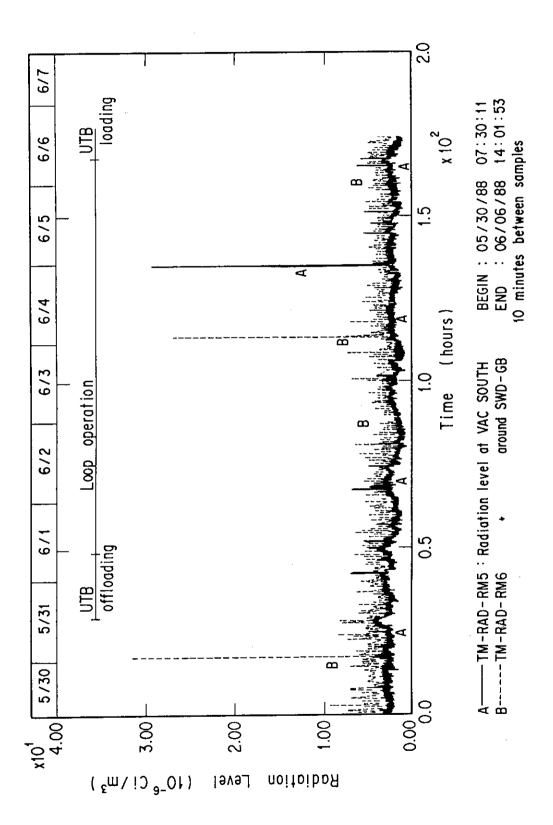


Fig. 3.21 Radiation Levels in Main Cell (3)

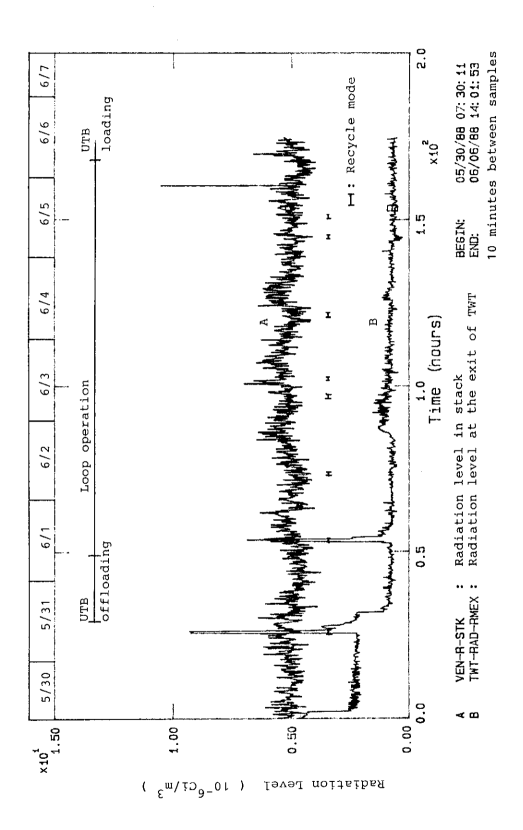


Fig. 3.22 Radiation Levels in the TWT-Outlet

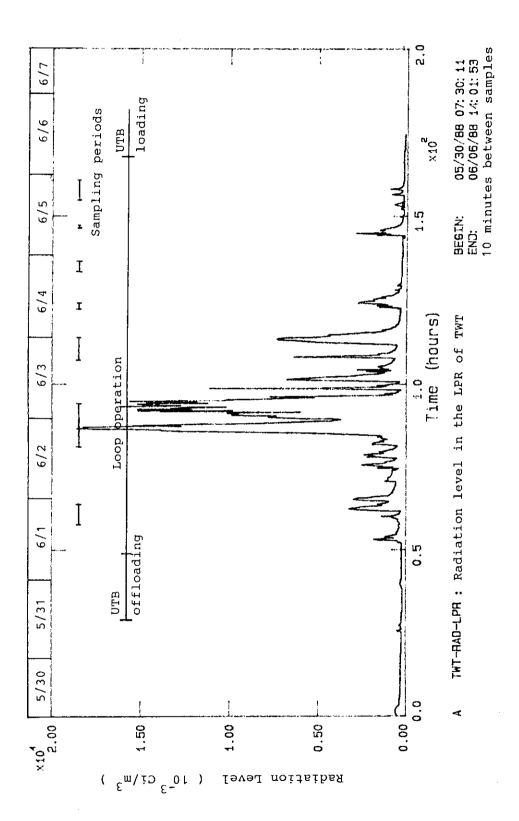


Fig. 3.23 Radiation Level in the TWT-Inlet (LPR)

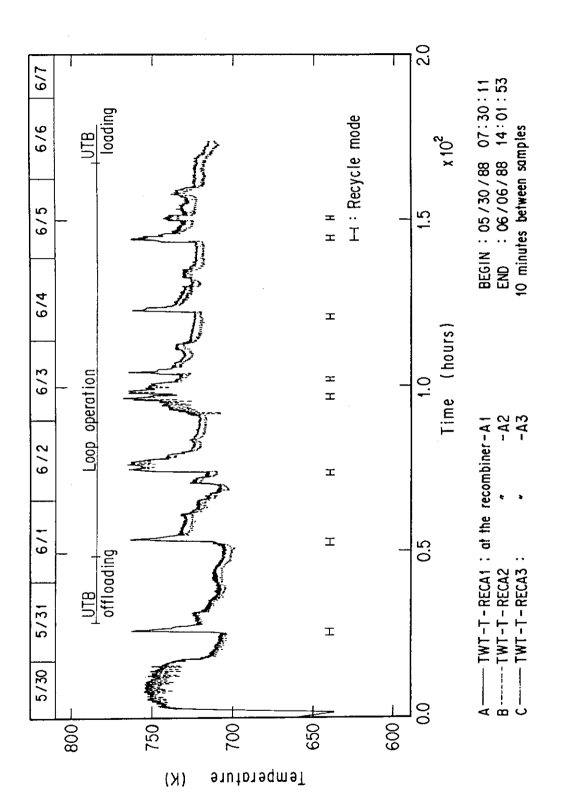


Fig. 3.24 Temperatures at the TWT Recombiner

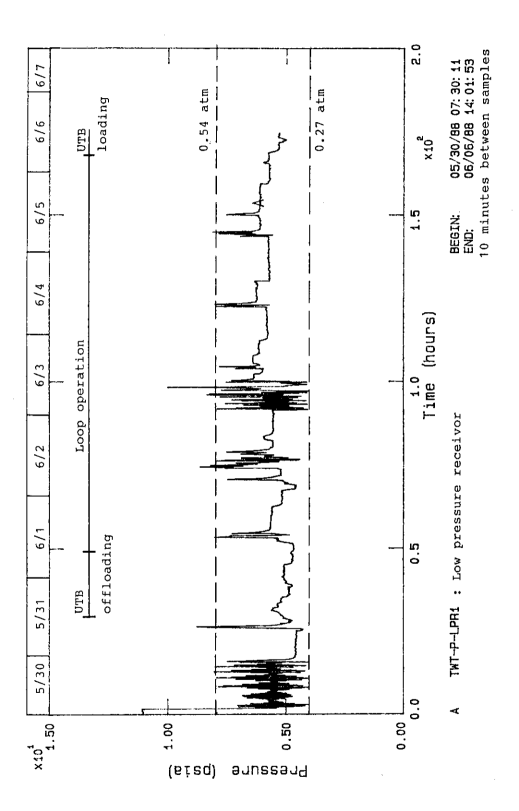


Fig. 3.25 Pressure in the TWT-Inlet (LPR)

4. RESULTS OF SYSTEM SHUTDOWN

4.1 PROCESS SYSTEMS

The following Figures show variations of the major process parameters on the ISS, FCU and UTB during the shutdown operations (Loop operation was stopped: 6/6/88~8:12, H-D-T dumping to UTB: 6/6/88~8:12).

Figure 4.1, outputs of the reboiler heaters in each column, shows that the heaters were maintained for approximately 1 hour because of enhancement of liquid boiled up. They were turned off when no liquid was observed in each reboiler.

Figure 4.2 shows that pressures of all columns immediately dropped by opening valves to UTB (Figures 3.10(a) and 3.10(b)), but increased to 500 Torr (column T) and to 800 Torr (columns I, H and D). These complex pressure changes are assumed to be depending on the amount of liquid in each column. The pressures were again increased after the helium refrigerator was turned off. The pressure of column T took time to increase after the He refrigerator was turned off, because the packed section of column H was cooled by He. Reduction of process loop pressures to a level lower than 400 torr was achieved in about 1 hour.

Figure 4.3 shows pressures in the FCU and NBI. The pressures of the FCU and NBI behaved like that of ISS columns, because gas was circulated in the process loop during shutdown.

During loading to UTB, blanketing effect due to the impurities was not observed in any of the uranium beds. The pressure was under 50 torr (Figure 4.2). The major components of

remaining gas were $^3{\rm He}$ and $^4{\rm He}$ which were shown in Table 4.1 (N $_2$ gas was already evacuated to TWT).

Figure 4.4 shows the rapid temperature rise due to the exothermic reaction between H-D-T mixture and uranium beds, that occurred immediately after opening the valves to UTB. The temperature peak levels on UTB2 and UTB4 connected to column H and D respectively were higher than that of the others. This phenomenon might have resulted from the difference in column inventory of hydrogen isotopes. The temperatures of UB1 and UB3 were relatively higher than that of others at a steady state after dumping of H-D-T mixture from the process loop, reflecting the effect of the tritium decay heat from uranium tritide.

4.2 SAFETY SYSTEMS

4.2.1 SECONDARY CONTAINMENT SYSTEM (SEC)

Figure 4.5 shows the radiation levels of the ISS-GB during process loop shutdown operation. The broad peak observed in the ISS-GB2 was caused by leaks from the sampling system of ISS due to pressure increase (up to 900 torr) of column I at shutdown operation (Figure 4.2).

No increase of radiation levels in other GBs was observed.

4.2.2 TRITIUM WASTE TREATMENT SYSTEM (TWT)

Figure 4.6 shows the radiation levels in the LPR and at the exit of TWT. The radiation level in the LPR increased slightly on account of the increase of that in the ISS-GB2. The TWT kept the radiation level low ($< 1 \times 10^{-3}$ Ci/m³) at the exit.

Table 4.1 The Result of Residual Gas Analysis of Process Loop after Shutdown of Run

6/ 7/88 Residual after Circ Mass Spec Analysis 3/ 8/88			225.00	torr	23.0 C	.050	liters	
		yy		Corr		anl Y	Summary	HDT mas
		Obs, I						7157 YES
H2	2.014	503, *	884		י ער	:	•	ш
He-3	2.016 3.016	9.1550	1.739	917	! Ha-3	49.447	, ! 341	74.993 D 25.007 T
HD	3.027	/11000	.715	* * * * * *	i ne o	1111111	. 047	25 AA7 T
Ho~4	4 003	2.6160		777	! Hp-4	12.270	!	10,007
			.795	* LL1	, nc , ! WT	121210	<u>'</u> !	ຄວກ
02	4.028	.0480 .0450	.729	002	! 137	.094	. 28.794	CD4
DT	5.030	0450	774	002	! DT	094	125	N?
12	6.032	••••	1,000		; DT ; T2	••••	;	
CH4	15.031	4,1790	.272	.028	CH4			
	17.027		.272		; NH3			T gas !
CH3D	17.038	8.8560	.272	. 058	: CH3D	3.155	, , -, }	
H20	18,011		.272		H20		 D	·
		17.3790	.272	.111	CH3T	6.016	1	;
	18.044		.272		; CH202		Grams H	;
	19.017		.272		HD0		Grags D	1
		17.5090	.272	.107	CHD2	5.898	Grams T	1
		2.5530	1.080	.062	Ne-20	3.328	Curies =	}
HTO	20.019		.272	,	: HTO		¦	
020	20.023		.272		1 020		as water	;
CH2T2	20.048		.272		: CH2T2		; H	+
CD4	20.056	15.1800	.272	.092	: CD4	4.984	; D	1
DTO	21.025		.272		: DTO		: T	
CD3T	21.058	11.0990	.272	.066	CD3T	3,556	Grams H	;
Ne-22	21.991		1.073		Ne-22		Grams D	!
120	22.027		.272		: 120		Grams T	1
CHT3	22.056		.272		ETH3		Curies =	;
		4.8470	.272					;
		3.4740						ne :
		3.5710	.272	.020		1.070		!
CO	27.995	.5330 .0440	.230		; C0		; D	;
N2	28.005	.5330	.230	.002	1 N2 1 O2	.125	1	i
82	31.990	.0440	. 281			.012	: D T Grams H Grams D	1
Ar	37.762	32.4760	.210	.108				
CO2	43.990		.193		1 CO2		Grams I	
								3.5 !
				1.853		100.000	•	•

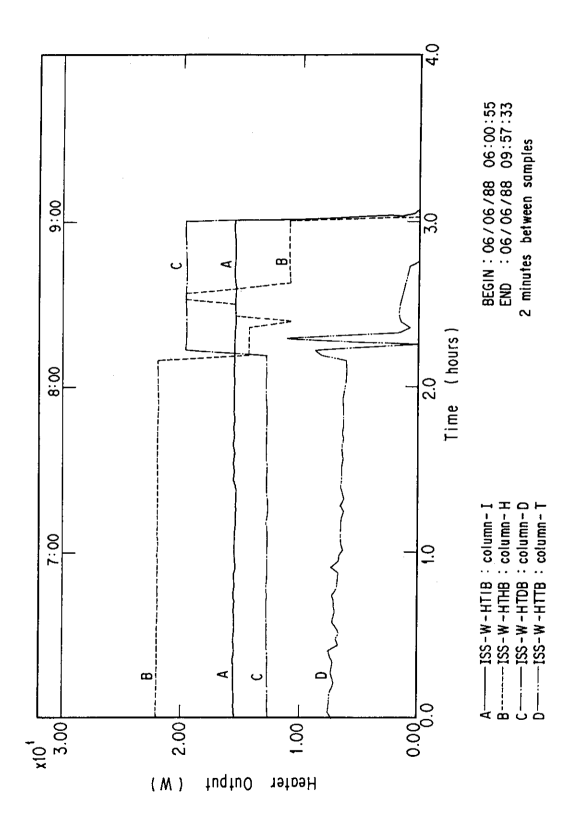


Fig. 4.1 Heater Outputs of ISS

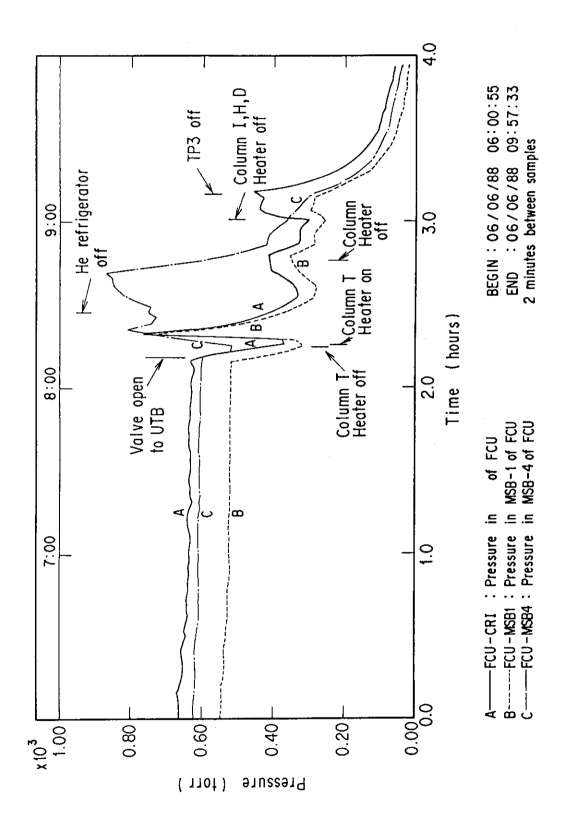


Fig. 4.2 Pressures of ISS Columns during Shutdown

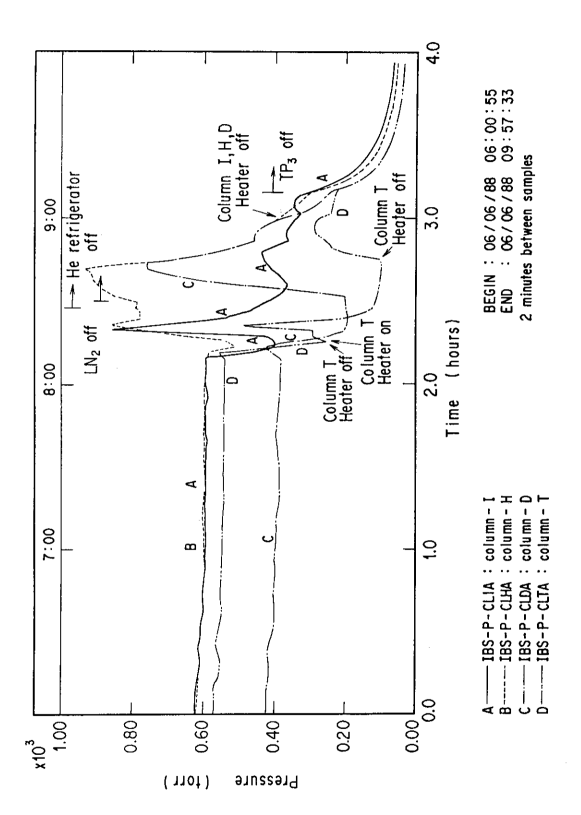


Fig. 4.3 Process Pressures in FCU during Shutdown

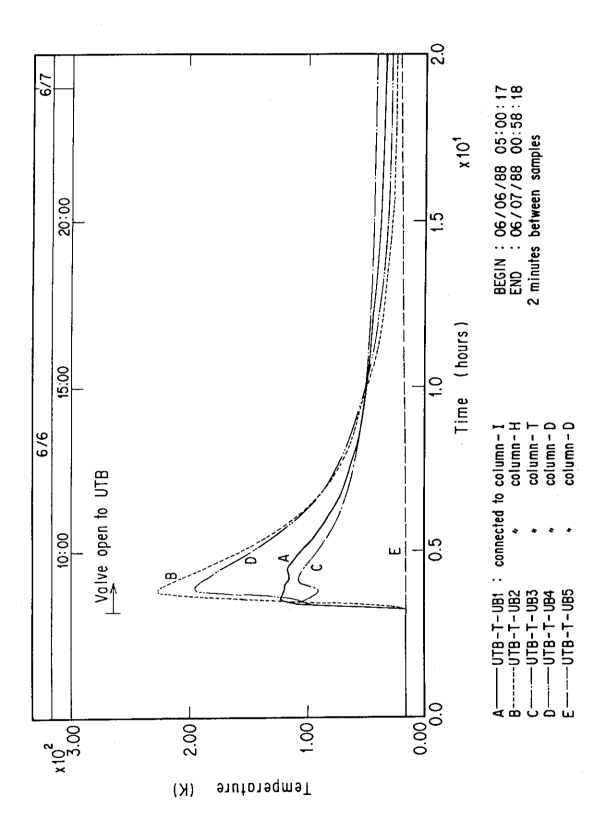


Fig. 4.4 Temperatures of UTB during Shutdown

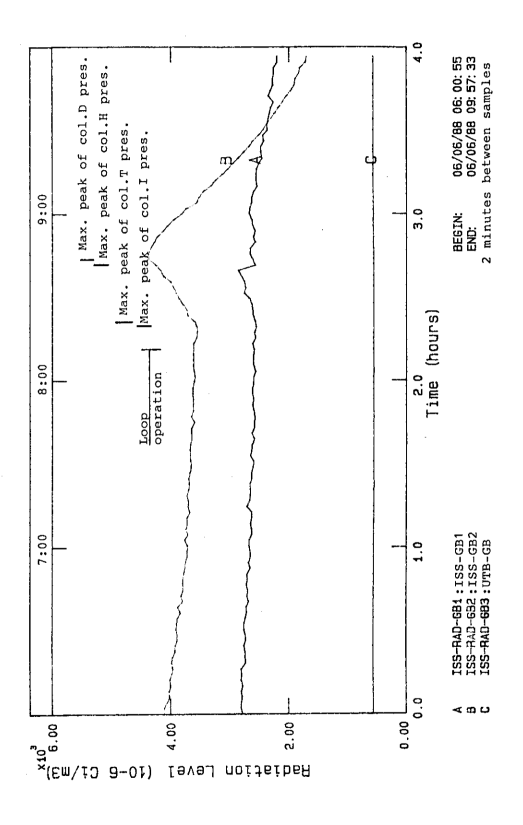


Fig. 4.5 Radiation Levels in ISS Gloveboxes during Shutdown

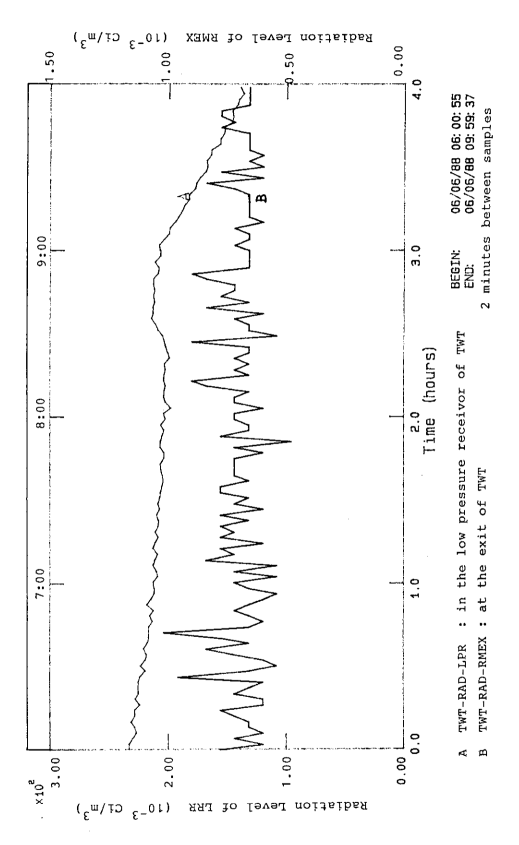


Fig. 4.6 Radiation Levels in the TWT-Inlet and Outlet during Shutdown

5. CONCLUSION

This milestone run was performed with the VAC system to demonstrate the TSTA full integrated fuel loop. Primary objectives were achieved without any accidental release of tritium to the experimental room or the environment.

Achievements of process systems are:

- full loop operation with the VAC, FCU, NBI and ISS was performed with approximately 84.4g-mole of H-D-T mixture(tritium; 107.0g, D/T ratio; about 2.15) for five days,
- the compound cryopump (BNL) was successfully integrated into the main loop. Pumping performance, as well as regeneration characteristics of VAC system, was measured using D-T, D-T with $\rm N_2$ and D-T with He (The

details will be reported in a separate paper.),

 the ISS was operated for five days under cryogenic distillation conditions with the VAC, FCU and NBI return process. Withdrawal of He, H₂ and HD mixture

from the top of column H was successfully carried out to remove He and ${\rm H}_2$ from the process loop,

- removal of ${\rm N}_2$ injected into the loop and the VAC system was successfully done by the FCU.

Issues of the safety systems are:

- there were several small released tritium into the gloveboxes, but no tritium release into the experimental room occurred,
- the amount of tritium exhausted to the TWT was estimated to be 4800 Ci. The amount of tritium from the ISS gas analysis was approximately 1900 Ci.
- there was no off-normal tritium release from the TSTA process loop into the environment during this run. The total release measured with the stack monitor (bubbler system) was:

5/30/88 - 6/5/88 HTO: 973.0 mCi HT: 62.5 mCi.

ACKNOWLEDGEMENT

The authors wish to express their sincere thanks to Dr. M. Enoeda, who has joined TSTA activity after this milestone run as a member of Annex IV 2nd year program, for all his efforts in the simulation study of ISS.

REFERENCES

- 1. H. Yoshida, S. Hirata, and T. Naito, T. Yamanishi :JAERI Research Team, and J.L. Anderson, J.R. Bartlit, R.V.Carlson, D.O. Coffin, R.H. Sherman and R.S. Willms : TSTA Research Team, "TSTA Loop Operation with 100 grams-level of Tritium Milestone Run in June, 1987 -," JAERI-M 88-204 , Japan Atomic Energy Research Institute (1988).
- 2. H. Yoshida, S. Hirata, and T. Naito, T. Yamanishi : JAERI Research Team, and J.L. Anderson, J.R. Bartlit, R.V.Carlson, D.O. Coffin, R.H. Sherman and R.S. Willms : TSTA Research Team, "TSTA Loop Operation with 100 grams-level of Tritium Milestone Run in July, 1987 -," JAERI-M 88-205, Japan Atomic Energy Research Institute (1988).
- 3. TSTA design team, "TRITIUM SYSTEMS TEST ASSEMBLY; FINAL SAFETY ANALYSIS REPORT," SAR-82-1F, Los Alamos National Laboratory (1982).

ACKNOWLEDGEMENT

The authors wish to express their sincere thanks to Dr. M. Enoeda, who has joined TSTA activity after this milestone run as a member of Annex IV 2nd year program, for all his efforts in the simulation study of ISS.

REFERENCES

- 1. H. Yoshida, S. Hirata, and T. Naito, T. Yamanishi :JAERI Research Team, and J.L. Anderson, J.R. Bartlit, R.V.Carlson, D.O. Coffin, R.H. Sherman and R.S. Willms : TSTA Research Team, "TSTA Loop Operation with 100 grams-level of Tritium Milestone Run in June, 1987 -," JAERI-M 88-204 , Japan Atomic Energy Research Institute (1988).
- 2. H. Yoshida, S. Hirata, and T. Naito, T. Yamanishi :JAERI Research Team, and J.L. Anderson, J.R. Bartlit, R.V.Carlson, D.O. Coffin, R.H. Sherman and R.S. Willms: TSTA Research Team, "TSTA Loop Operation with 100 grams-level of Tritium Milestone Run in July, 1987 -," JAERI-M 88-205, Japan Atomic Energy Research Institute (1988).
- 3. TSTA design team, "TRITIUM SYSTEMS TEST ASSEMBLY; FINAL SAFETY ANALYSIS REPORT," SAR-82-1F, Los Alamos National Laboratory (1982).