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JOHN F. KENNEDY SPACE CENTER UNIVERSITY OF CENTRAL FLORIDA

CHARACTERIZATION OF A TURBOMOLECULAR-PUMPED MAGNETIC SECTOR MASS SPECTROMETER

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ABSTRACT

A Perkin Elmer MGA-1200, turbomolecular-pumped, magnetic sector, multiple gas analyzer mass spectrometer with modified inlet for fast response was characterized for the analysis of hydrogen, helium, oxygen and argon in nitrogen and helium background gases. This instrument was specially modified for the Vanderberg AFB SLC-6 Hydrogen Disposal Test Program, as a part of the Hydrogen Sampling System (H2S2). Linearity, precision, drift, detection limits and accuracy among other analytical parameters for each of the background gas were studied to evaluate the performance of the instrument. The results demonstrate that H2S2 mass spectrometer is a stable instrument and can be utilized for the quantitative analytical determination of hydrogen, helium, oxygen and argon in nitrogen and helium background gases.

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I- INTRODUCTION

Hazardous gases are used extensively in various phases of the space shuttle launch at the Kennedy Space Center (KSC). Presence of hydrogen (H₂), even in small amounts in unwanted areas may generate severe hazardous conditions. In order to avoid the accumulation of these gases, nitrogen (N₂) and helium (He) are used as purge gases in various shuttle compartments and in cryogenic fuel lines respectively. This results in the importance of monitoring the presence of hydrogen, oxygen and other inorganic gases in the purged environments for a safe space shuttle launch.

Gas detection system used at KSC for the space shuttle launches involves the monitoring of the hazardous gases in various purged environments. Mass spectrometers and other analytical instruments, located in the Mobile Launch Pad (MLP) are used for the analysis of these purged gases. The spectrometers, in the past, had diode ionpumps for their high vacuum systems, and have proven to be highly reliable over numerous shuttle launches for the detection of hydrogen, oxygen, argon and helium in nitrogen purged areas. Helium purged areas could not be monitored by these systems since ion-pumps are not well suited to pumping with a nearly 100% helium background.

Recently, mass spectrometers with turbo-molecular vacuum pumps have become available which are ideal for monitoring the helium purged environments. These pumps have an advantage that they are not affected by the noble gases. Currently at KSC, turbomolecular-pumped mass spectrometers are being tested for use in monitoring helium and nitrogen purged areas and the results have been verv promising. The present investigation has focussed primarily on the evaluation of a modified version of a Perkin Elmer's magnetic sector, multiple gas analyzer mass spectrometer having a turbo-molecular pump installed for its vacuum system. Precisely, it involves the evaluation of the instrument for the analysis of hydrogen, oxygen, helium and argon in nitrogen and helium background gases.

II- MATERIALS AND METHODS

Α.

1- Mass Spectrometric System

The mass spectrometer used for this work was a Perkin Elmer's MGA-1200 Multiple Gas Analyzer (H2S2) with a mass range of 2-135 atomic mass unit. The spectrometer is a magnetic sector type which utilizes a turbo-molecular pump (50 L/sec capacity) for its vacuum system, The system was not further modified for this investigation. The MGA operates with a pressure of 50 Torr at its inlet leak and is equipped with Faraday detectors for intercepting the ion beams. The block diagram of the MGA system is presented in Figure 1. Figure 2A and 2B respectively demonstrate the flow diagram of the sample delivery system and the operation of the magnetic sector mass spectrometer.

2- Calibration Gases

Matheson Gas Products compressed gas cylinders containing mixtures of various gases of interest at different levels of concentrations in nitrogen and helium were used for this study. Pure nitrogen, air and helium gases were supplied by KSC.

B. Sample Delivery System

The sample delivery system used was a 8-port rotary gas valve for selecting the correct calibration gas. Standard gas mixtures with nitrogen and helium as the background gases were connected to the rotary valve having an electronically controlled switch. The switch facilitated rapid changes between various gas mixtures. The outlet of the rotary valve was connected to a pre-calibrated sample flow controller and the average of the flow controller was connected to the capillary of one of the inlet ports of the instrument via a digital calibrated gas flow meter (Sierra) and a T-connector. This T-connector is important

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for maintaining the sample inlet capillary at the atmospheric pressure. The total length of the tubing from the rotary outlet to the inlet capillary of the spectrometer was about six feet. Standard gas bottles were connected to the rotary valve with 2-4 feet long tubing. PVC tubing was used for all the gas transport lines. All the pressure regulators at the standard gas bottles were operated at an outlet pressure of about 15 lbs/in2. MGA required a minimum of 0.40 SLPM sample inlet flow for the capillary having inner diameter of 0.025 inches. The flow was optimized by monitoring the oxygen channel reading of the mass spectrometer while cycling a standard gas mixture through the system. Below this setting the "clean capillary" indicator light indicates that an insufficient amount of sample is reaching the inlet capillary. A higher flow rate of upto 0.8 SLPM of gas did not affect the performance of the instrument. Unless otherwise specified, a sample flow rate of 0.48-0.52 SLPM through the system was maintained for all the analytical measurements. The gas flow requirements may vary according to the diameter of the capillary connected to the sample inlet port of the instrument.

All the measurements were taken after a delay of about 15 seconds stabilization time. This was the time required by the mass spectrometer to respond accurately to the sample change at the rotary value at the flow rate used for the experimental work.

C. Calibration

For its characterization, the mass spectrometer was calibrated using analyzed gas mixtures from Matheson Gas Products. The instrument was calibrated with a zero gas and a standard gas mixture having known concentrations of the gases to be analyzed in their respective background gases. For the analysis in nitrogen background. The calibration standard gas mixture contained:

H₂ - 2.601%

He	-	7.006%
02	-	7.007%
Ar	-	1.001%

For the analysis in helium background, the calibration standard gas mixture contained:

H ₂	-	2.600%
02	-	5.003%
Ar	-	1.000%

D. Linearity and Precision Tests

The linearity and repeatability tests for the analyte gases in nitrogen and helium background gases were performed by analyzing standard gas mixtures containing varying concentrations of the analyte gases in their respective backgrounds. The experimental setup procedure for the determination of these parameters is shown in Figure 3. The output of each analyte gas channel was recorded in a digital form through an external digital voltmeter and also on a 6channel chart recorder as the zero gas and the standard gas mixtures were cycled through the rotary valve.

E. Drift Tests

Drifts in the output of the various gas channels of the instrument were monitored for 24 hours by cycling standard gas mixtures in nitrogen and helium background gases. The setup procedure for this study is presented in Figure 4. The standard gas mixtures used for this study are as follows:

1. Analyte gases in nitrogen background

H ₂	-	2.598%
He	-	5.006%
0 ₂	-	5.122%
Ar	-	1.000%

2. Analyte gases in helium background

H ₂	-	7.058%
O2	-	3.297%
Ar	-	0.999%

The instrument was calibrated for the appropriate background gas (N_2 or He) with the standard mixture before the start of the test. Appropriate zero gas kept flowing through the system for most of the time except for when the instrument inlet was switched to the other inlet port for analyzing the standard gas mixture. After recording the output of the various channels on a strip chart recorder and in a digital form at certain intervals of time, the inlet was switched back to the original zero gas inlet of the instrument.

F. Detection Limits

The output of H2(0-100% range), He(0-100% range), O2(0-25% range) and Ar(0-100% range) analyte channels of the instrument were recorded on a strip chart recorder preset in milli-volts ranges by using a fast scan speed (30 cm/min) for 30 seconds when appropriate zero gas was flowing through the system.

G. Response to a Gas Pulse

H2S2 is equipped with a most model for a fast response to detect a sudden sample change in the zero gas flowing through the system. The purpose of this test was to see how fast the instrument can detect 90% of the gas peak from the base value when a gas pulse of one zero gas was generated into another zero gas. The response of the instrument was recorded on a strip chart recorder. The setup procedure for this test is presented in Figure 5. Gas pulses of half a second to one second were generated from one zero gas into another zero gas through a fast-response Marotta electric valve. An auto timer to generate the timed pulses, and to control the strip chart recorder simultaneously was constructed in the laboratory.

III - RESULTS AND DISCUSSIONS

A. Linearity and Precision Tests

A1- Nitrogen Background Gas

Data obtained by analyzing a series of seven standard gas mixtures containing varying concentrations of analyte gases are presented in Table 1A. The experimental averages with background correction are the average of a set of eleven consecutive runs. The background levels of various analyte gases were obtained when pure nitrogen was cycling through the system. The relative errors in the analytical results for most of the standards for the analyte gases are in the range of 0.05-3.05% except for hydrogen on standard mixture bottle numbers 2 and 6. No obvious explanation can be offered for this deviation except to question the accuracy of the supplied standard bottle mixtures. The linear curves obtained for the various analyte gases are presented in Figures 6A-6E. The curves demonstrate a linear response of the mass spectrometer to the concentration range of the various analyte gases tested under the conditions of the experiment.

Repeatability test data are presented in a graphic form in Figures 7A-7C. The coefficient of variation for the eleven consecutive test runs is less than 2% for all the analyte gases.

The results obtained for the detection limits of the four analyte gases when pure nitrogen zero gas was cycled through the system are presented in Table 1B. The detection limits for the four analyte gases in nitrogen background gas taken as twice the standard deviation are as follows:

H ₂ (0-100% range)	-	197	PPM	
O ₂ (0-25% range)	-	26	PPM	ORIGINAL PAGE IS OF POOR ONALITY
He (0-100% range)	-	30 5	PPM	
Ar (0-100% range)	-	27	PPM	

A2- Helium Background Gas

Table 2A demonstrates the data obtained by analyzing a series of standard gas mixtures in helium zero gas. The experimental averages with background correction are again the averages of a set of eleven consecutive runs. The background levels of various analyte gases were also obtained when pure helium was cycling through the system. The relative errors in the analytical results for various gas mixtures for the three analyte gases are in the range of 0.05-4.03% except for the two standard mixtures corresponding to bottle numbers 2 and 5. Again no explanation can be offered for this discrepancy except suggesting to check the accuracy of these standard mixtures by running fresh and reliable new standards. The linear curves obtained for the various analyte gases are presented in Figures 8A-8D. The curves demonstrate a linear response of the mass spectrometer to the concentration range of the various analyte gases tested under the prescribed conditions of the test.

The repeatability test data is presented in a graphic form in Figures 9A and 9B. The coefficient of variation of the eleven consecutive test runs is less than 1% for all the analyte gases in helium background gas.

The results obtained for the detection limits of the three analyte gases when pure helium was cycled through the system are presented in Table 2B. The detection limits for the three analyte gases taken as twice the standard deviation are as follows:

H ₂ (0-100% RANGE)	-	109	PPM
0 ₂ (0-25% RANGE)	-	22	PPM
Ar (0-100% RANGE)	-	15	PPM

B. Drift Study

B1- Nitrogen Background Gas

During the analysis of the zero gas over a 24-hour period, all of the analyte channels demonstrated a negative drift from their initial values and fell between 138 to 300 PPM except for argon which drifted about 70 PPM.

Table 3 demonstrates the drift in the analytical readout obtained for the standard gas mixture in nitrogen with background correction. The drift study plots of the various analyte gas channels are presented in Figures 10A-10C. Most of the channels demonstrate stability with small fluctuations, about 1% from the initial value, except for argon which deviated about 4% of the initial value in the positive direction.

B2- Helium Background Gas

For the zero helium gas analysis, all the channels drifted in the negative direction from the start of the experiment. The drift in all the analyte gas channels fell in the range of 50 to 220 PPM.

Table 4 demonstrates the drift in the analytical readout of the various analyte gas channels for the standard gas mixture in helium with background corrections. The data is presented in a graphic form in Figures 11A and 11B. Hydrogen channel demonstrates a negative drift from the initial value while oxygen and argon reflect an upward trend. Hydrogen and oxygen channels drifted about 2% and argon channel drifted about 6% from their initial readout.

B3- General

The results obtained on the drift study of the various analyte gas channels of the instrument reflect that for accurate and precise analytical results, periodic background corrections are needed during the analysis of gases in nitrogen background gas. It is especially important during the analysis of the analyte gases in helium background gas.

C. Response to a Gas Pulse

Tables 5A and 5B demonstrate the results obtained for the time required by the instrument to detect 90% peak of a zero gas from the baseline when pulses of half to one second durations were genearated into another zero gas flowing through the system. The results demonstrate that H2S2 can detect 90% of the peak from the baseline, from a distance of about six feet from its T-capillary connection. a half a second pulse of nitrogen zero gas into helium zero gas flowing at a rate of 3-6 SLPM in about 0.6 to 0.7 second (Table 5A). For pulses of similar durations of helium zero gas into nitrogen zero gas flowing at the same rate, the time was found to be about 0.4 second (Table 5B). Figure 12 demonstrates a typical strip chart recording of the response time for a pulse of half a second of nitrogen zero gas genearated into helium zero gas flowing at a rate of 6 SLPM.

IV - CONCLUSIONS

- 1 The linearity data demonstrate a linear response of the instrument in the concentration range tested for the analysis of oxygen, hydrogen, helium and argon in nitrogen background gas, and hydrogen, oxygen and argon in helium background gas.
- 2-Optimum sample flow required for the instrument is 0.48-0.52 SLPM for the sample inlet port capillary having inner diameter of 0.025 inch. Higher flow rates of upto 0.8 SLPM did not affect the analytical readout of the instrument. The flow rate should be optimized for the individual sample inlet port capillary.
- 3 The detection limits of the multiple gas analyzer mass spectrometer for the analysis of the analyte gases in nitrogen zero gas are: H₂(0-100% range)-197 PPM, O₂(0-25% range)-26 PPM, He(0-100% range)-305 PPM and Ar(0-100% range)-27 PPM. The detection limits in helium zero gas are: H₂(0-100% range)-109 PPM, O₂(0-25% range)-22 PPM and Ar(0-100% range)-15 PPM.
- 4 The drift study reflects that the analyte gas channel output are much stable in nitrogen zero gas as compared to helium zero gas. For accurate and precise analytical results, periodic background correction is needed during the analysis of the analyte gases in nitrogen background gas. However, it is especially important during the analysis in helium background gas.
- 5 Minimum response time required by the instrument to detect 90% of a peak of nitrogen zero gas from the baseline was found to be 0.6-0.7 second when publics of half a second duration of nitrogen zero gas were generaled into helium zero gas flowing at a rate of 3-6 SLPM. For pulses of helium zero gas of similar duration generated into nitrogen zero gas flowing at the same rate, the response time was found to be

about 0.4 second.

6 - The present study on the characterization of the Perkin Elmer's MGA-1200, turbomolecular-pumped, magnetic sector, multiple gas analyzer mass spectrometer (H2S2) demonstrates that the instrument can be utilized for the quantitative analytical determination of hydrogen, oxygen, helium and argon in nitrogen and helium background gases. TABLES

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		B	c l	D 1	E	F	G
		H2 10%	H2 100%		N2	O2 25%	Ar
; †	BOTTIF2						
3							
Ă	STREAM	0.9850	0.9850	0.7998	96.7132	1.0000	0.5020
$\overline{\mathbf{x}}$		0.3030					
-	EXPAVERAGE	0.8301	0 8327	0.7936	97.3364	1.0107	0.5017
	DAT AVENUC						
	STDDEV	0.0038	0.0065	0.0163	0.0446	0.0015	0.0070
-	3.000	0.0000					
10	COE OF VARIAT	0 4217	0 7244	2.0529	0.0458	0.1600	1.4040
11							
12	REI FRADA	9.6354	9.3678	0.7705	0.6443	1.0632	0.0543
13	<u> </u>						
14					••••		•••••
15	tt						
16	1	H2 10%	H2 100%	He	N2	02 25%	Ar
17	BOTTLES						
18							
19	STDCONC	2 5040	2.6040	11.3990	72,3840	12.0110	1.0020
20							
21	EXPAVERAGE	2 54 34	2,6445	12.0292	72.2145	12.0439	1.0223
22							
23	STD DEV	0.0061	0.0082	0.0133	0.0169	0.0225	0 0 1 9 5
24							
25	COF OF VARIAT	0.2309	0.3101	0,1104	0.0235	0.1881	1,9034
26	1						
27	RELEBROR	1.5117	1.5570	0.2412	0.1927	0.2736	2.0232
28							<u> </u>
29							
30	1					·	
31		H2 10%	H2 100%	He	N2	02 25%	Ar
32	BOTTLE 4						
33							ļ
34	STD COAC	2.5980	2.5980	5.0060	86.2740	5.1220	1.0000
35							<u> </u>
36	EXPAVERAGE	2.6620	2.5555	4.9545	86.0273	5.1305	1.0202
37							
38	STD CEV	0 0027	0.0059	0.0059	0.0509	0.0035	0.0165
39						1	
40	CE CE VARA	0.1022	0.2579	0,1388	0.0707	0 1860	1 5255
41	· · · · · · · · · · · · · · · · · · ·						
4	PELEPRCA	2.4634	1 2.5964	1 0279	0 2950	0.1551	2 0182

LINEARITY AND PRECISION STUDY: STATISTICAL DATA OF GASES IN N2 BKG GAS

LINEARITY AND PRECISION STUDY: STATISTICAL DATA OF GASES IN N2 BKG GAS

						5	6
	A .	8	C	D	E		
3				115		02 25%	Ar
4		H2 10%	H2 100%	110			
5	BOTTLES			+			
6			2 6010	7 0050	82,3850	7.0070	1.0010
	SIUCURE	2.6010	2.5010	- /.0000			
	EVR AVERACE	2 5472	2 6491	7 0009	82.1827	7.0255	1.0207
	EAPATEMOE	2.04/2	2.0431				
	STODEY	0.0055	0.0083	0 0083	0 0 2 0 5	0 0117	0 0184
2	310000						
12	OF OF VARIAT	0.2075	0.3138	0.1187	0.0250	0.1660	1.8039
54							
55	RELERADA	1.7755	1 8489	0 0727	0 2455	0 2634	1.9705
56							
57							
58						00.05 %	
59		H2 10%	H2 100%	He	N2	02 25%	<u> </u>
60	BOTTLES			 			
61					02.0000		
62	STD CONC	8.0000	8.0000		92.0000		
63				+	93.0427	+	
64	EXPAVERAGE	84567	8 4582		33.042.		
65	cm cov		0.0126	+	0.0429		
60	SIDUEV	0.0107	Q.0125				
60/	CE CE VARIA	0 1270	0.1478		0.0461		
60	por or earling	1 0.1210					
70	RELETING	5 7341	5,7273	-	1.1334		
71	1	1	1				
72							· · · · · · · · · · · · · · · · · · ·
73							
74		H2 10%	H2 100%	Ha	N2	02 25%	Ar
75	BOTTLE 7						+
76							1 0000
77	STD CONC	5,1450	5,1450	5.0040	85.54.30	3.2940	10000
78						2 3 2 5 0	0 9946
79	FXP AVERAG	E 4.9921	4 9852	4.9000	85.4691	3.3230_	
80					0.0270	0.0061	0.0167
61	STD CEV	0.0087	0.0060	0.0089			
82					0.0316	0.1842	1.6741
6 3	COF CF VARIA	0.1734	0.1209	0.1825	0.0310		
84				2.0791	0.0934	0.9411	0 5 3 6 4
83	HEL EFFICIE	29/20	3 0480	2.0783		_	
1							
8.4			-				
80		H2 10%	H2 100%	He	N2	02 25%	Ar
100	BOTTLE						
91							
19	STOCCER	10 00 20	10 0030	10 00 10	75 5370	3 3000	1 0.010
9	3						
9.	ERPACEA	1E 98792	9.900	9 30.7	75 5645	3 3 3 3 3 3 3 3 3 3 3 3 3 3 3 3 3 3 3 3	0 3351
9	5						
9	6 STDCFV	0 0 1 3 1	0.0352	0.00 9	0 0 1 97	0.0051	00181
9	7						1 0 3 1 6
9	B COF CF VAR	AT 0 1327	0.3561	0.0274	0.0260	0.1839	1 1 8 3 1 3
9	9						1 2716
10	0 RELEFIED	A 1 2872	1.1791	+01/3	1 0.1618	1 0 9031	

Table 1B

	A [В	С	D	<u> </u>	F
1			02 25%	He	H2 100%	Ar
2						
3	n=30		CONC, PPM	CONC, PPM	CONC, PPM	CONC, PPM
4						
5			31.25	200.00	150.00	40.00
6			25.00	400.00	50.00	30.00
7			25.00	140.00	20.00	20.00
8			18.75	700.00	50.00	50.00
9			43.75	340.00	10.00	30.00
10			25.00	300.00	170.00	50.00
11			25.00	200.00	120.00	62.00
12			25.00	240.00	10.00	20.00
13			50.00	240.00	400.00	42.00
14			6.25	100.00	280.00	40.00
15			37.50	260.00	50.00	8.00
16			43.75	240.00	120.00	48.00
17			18.75	300.00	50.00	40.00
18			18.75	600.00	100.00	20.00
19			12.50	220.00	50.00	36.00
20			43.75	500.00	250.00	20.00
21			12.50	700.00	170.00	34.00
22			6.25	400.00	200.00	24.00
23			31.25	240.00	150.00	30.00
24			.27.50	200.00	120.00	30.00
25			31.25	500.00	160.00	40.00
26			50.00	- 400.00	150.00	30.00
27			50.00	360.00	100.00	32.00
28			50.00	480.00	70.00	52.00
29			31.25	440.00	200.00	54.00
30			31.25	240.00	400.00	62.00
3 1			18.75	400.00	150.00	26.00
32			18.75	360.00	50.00	40.00
33			25.00	200.00	150.00	40.00
34			25.00	300.00	150.00	20.00
35						
36	EXP AVERACE	(BKG)	28.63	340.00	136.67	35.67
37						
38	STDCEV	(S.D.)	12.77	152.50	98.73	13.27
39						
40	DETECT LIMIT	(2 S D.)	25.54	305 00	197.46	26.54

BACKGROUND CONCENTRATIONS IN N2 FOR DETECTION LIMIT DETERMINATION

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Та	p	le	2A
Ia	b	le	2A

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LINEARITY AND PRECISION STUDY: STATISTICAL DATA OF GASES IN HE BKG GAS

T	A	В	С	D	E	F
1		H2 10%	H2 100%	He	02 25%	Ar
2	BOTTLE 2					
3						
4	STDCONC	3.0140	3.0140	92.9830	3.3003	1.0000
5						
6	EXP AVERAGE	3.2092	3.2464	92.3700	2.8966	0.9789
7						
8	STDDEV	0.0076	0.0150	0.0307	0.0038	0.0066
9	005 05 14 5 4 7		0.4005	0.0222	0.1206	0.5721
10		0.2381	0.4625	0.0332	0.1290	0.0721
11		6 4769	7 7095	0.6593	12 2325	2 1091
12	HELENNUN	0.4/30	7.7095	0.0333	12.2025	
1 4						
15						
1 6		H2 10%	H2 100%	He	O2 25%	Ar
17	BOTTLE 3					
18						
19	STDCONC	9.9980	9.9980	90.0020		
20]					
21	EXP AVERAGE	9.8432	9.9827	91.0164		
22						
23	STD DEV	0.0144	0.0200	0.0329		
24						
25	COF OF VARIAT	0.1461	0.2008	0.0362		
26		1.5405	0 15 29	1 1 2 7 0		
21	RELERHOR	1.5485	0.1528	1.12/0	+	+
20			+		+	
120	1	<u> </u>				
31						1
3 2	+	H2 10%	H2 100%	He	02 25%	Ar
3 3	BOTTLE 4	1				
34						
3 5	STDCONC	2.5980	2.5980	87.4120	9.9900	0.9990
36					1	
37	EXP AVERAGE	2.5230	2.5718	88.1964	9.7530	1.0079
38						
39	STD DEV	0.0105	0.0214	1.0214	0.0086	0.0060
40				4.5500		0 5050
4 1	UUF UF VARIAT	0.3956	0.8306	4.5596	0.0887	0.5960
4 2		2	1.0074	0.8973	2 3728	0.8918
4 3	I ACLEMMUM	2.0000	1.0078	0.0973	2.3720	1 0.0010
<u> </u>		<u>!</u>		<u></u>		<u> </u>

Table 2A Continue

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	A	В	С	D	E	F
45						
46						
47		H2 10%	H2 100%	He	02 25%	Ar
48	BOTTLES		·····			
49	CTD COALD	2 0 0 2 0	0.0000	05 1097	1.0000	0.0000
50	SIDUNC	2.0020	2.0020	95.1967	1.9900	0.0003
52	EXPAVEBAGE	1 8584	1 8836	95 1609	1 9425	0 7899
53	DV AVD VOL		1.0030		1.5425	0.7000
54	STDDEV	0.0060	0.0050	0.0365	0.0034	0.0053
55						
56	COF OF VARIAT	0.3231	0.2678	0.0383	0.1727	0.6686
57						
58	RELERROR	7.1746	5.9123	0.0397	2.8264	1.2984
59			······································			
60			<u></u>		• • • • • • • • • • • • • • • • • • •	
61						
62		H2_10%	H2 100%	He	02 25%	Ar
63	BOTTLE 6		· · · · · · · · · · · · · · · · · · ·			
64						
65	STDCONC	2.6000	2.6000	91.3970	5.0030	1.0000
66		0.5140	0.5470	01.1545	1 0000	0.0050
	EXP AVEHAGE	2.5118	2.54/3	91.1045	4.8380	0.9950
60	STODEV	0.0049	0.0195	0.0419	0.0168	0.0077
70	510024	0.0043	0.0165	0.0418	0.0100	0.0077
71	COF OF VARIAT	0.1949	0.7258	0.0459	0.3476	0.7744
72						
73	RELEAROR	3.3916	2.0280	0.2543	3.2853	0.4364
74						
75					1	• • • • • • • • • •
76						
77		H2 10%	H2 100%	He	02 25%	Ar
78	BOTTLE 7					
1/9		7.0500	7.0500	00.0400		0.0000
80	- SIDUNU	7.0580	1.0580	88.6460	3.2970	0.9990
01	EVPANEDACE	7 2422	7 4245	97 9055	3 1025	0.0945
81	LAF AVERAUE	1.3423	1.4345	07.0933	3.1925	0.3045
8 4	STODEV	0.0091	0.0216	0 0372	0.0030	0.0074
8 5			0.0210	0.0072	1 0.0000	
86	COF OF VARIAT	0.1239	0.2908	0 0424	0.0927	0.7483
87	1		1			
88	RELEBBOR	4 0277	5 3 3 5 0	0.8467	3 1695	1 4459

LINEARITY AND PRECISION STUDY: STATISTICAL DATA OF GASES IN He BKG GAS

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Table 2A Continue

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LINEARITY AND PRECISION STUDY: STATISTICAL DATA OF GASES IN He BKG GAS

	A	B	С	D	E.	F
89						
90						
91						
92		H2 10%	H2 100%	He	02 25%	Ar
93	BOTTLE 8					
94						
95	STDCONC	5.0040	5.0040	90.6950	3.3010	1.0000
96						
97	EXP AVERAGE	5.0011	5.0782	90.2473	3.1961	0.9871
98						
99	STD DEV	0.0073	0.0060	0.0297	0.0034	0.0074
100						
101	COF OF VARIAT	0.1462	0.1187	0.0329	0.1070	0.7547
102						
103	RELERROR	0.0581	1.4825	0.4937	3.1767	1.2909

1 $O2 25\%$ $H2 100\%$ Ar 2
2 3 n=30 CONC,PPM CONC,PPM CONC,PPM 4
- - CONC,PPM CONC,PPM CONC,PPM 4 - - - - 5 35.00 70.00 35.00 6 30.00 90.00 8.00 7 10.00 90.00 20.00 8 35.00 20.00 10.00 9 20.00 70.00 20.00 10 12.50 190.00 15.00 11 25.00 110.00 18.00 12 42.50 140.00 18.00 13 25.00 130.00 10.00 14 2.50 60.00 15.00 15 12.50 70.00 5.00 16 2.50 100.00 5.00 17 30.00 100.00 30.00
4 35.00 70.00 35.00 5 30.00 90.00 8.00 7 10.00 90.00 20.00 8 35.00 20.00 10.00 9 20.00 70.00 20.00 10 12.50 190.00 15.00 12 42.50 140.00 18.00 13 25.00 130.00 10.00 14 2.50 60.00 15.00 15 12.50 70.00 5.00 16 2.50 100.00 5.00 17 30.00 100.00 30.00
5 35.00 70.00 35.00 6 30.00 90.00 8.00 7 10.00 90.00 20.00 8 35.00 20.00 10.00 9 20.00 70.00 20.00 10 12.50 190.00 15.00 11 25.00 110.00 18.00 12 42.50 140.00 18.00 13 25.00 130.00 10.00 14 2.50 60.00 15.00 15 12.50 70.00 5.00 16 2.50 100.00 5.00 17 30.00 100.00 30.00
6 30.00 90.00 8.00 7 10.00 90.00 20.00 8 35.00 20.00 10.00 9 20.00 70.00 20.00 10 12.50 190.00 15.00 11 25.00 110.00 18.00 12 42.50 140.00 18.00 13 25.00 130.00 10.00 14 2.50 60.00 15.00 15 12.50 70.00 5.00 16 2.50 100.00 30.00
7 10.00 90.00 20.00 8 35.00 20.00 10.00 9 20.00 70.00 20.00 10 12.50 190.00 15.00 11 25.00 110.00 18.00 12 42.50 140.00 18.00 13 25.00 130.00 10.00 14 2.50 60.00 15.00 15 12.50 70.00 5.00 16 2.50 100.00 30.00
8 35.00 20.00 10.00 9 20.00 70.00 20.00 10 12.50 190.00 15.00 11 25.00 110.00 15.00 12 42.50 140.00 18.00 13 25.00 130.00 10.00 14 2.50 60.00 15.00 15 12.50 70.00 5.00 16 2.50 100.00 30.00
9 20.00 70.00 20.00 10 12.50 190.00 15.00 11 25.00 110.00 15.00 12 42.50 140.00 18.00 13 25.00 130.00 10.00 14 2.50 60.00 15.00 15 12.50 70.00 5.00 16 2.50 100.00 30.00
$\begin{array}{c ccccccccccccccccccccccccccccccccccc$
$\begin{array}{c ccccccccccccccccccccccccccccccccccc$
12 42.50 140.00 18.00 13 25.00 130.00 10.00 14 2.50 60.00 15.00 15 12.50 70.00 5.00 16 2.50 100.00 5.00 17 30.00 100.00 30.00
13 25.00 130.00 10.00 14 2.50 60.00 15.00 15 12.50 70.00 5.00 16 2.50 100.00 5.00 17 30.00 100.00 30.00
14 2.50 60.00 15.00 15 12.50 70.00 5.00 16 2.50 100.00 5.00 17 30.00 100.00 30.00
15 12.50 70.00 5.00 16 2.50 100.00 5.00 17 30.00 100.00 30.00
16 2.50 100.00 5.00 17 30.00 100.00 30.00
<u>17</u> <u>30.00</u> <u>100.00</u> <u>30.00</u>
18 12.50 170.00 32.00
19 17.50 60.00 19.00
20 45.00 170.00 16.00
21 22.50 20.00 20.00
22 10.00 160.00 29.00
23 22.50 130.00 25.00
24 20.00 60.00 20.00
3 3 2 6 EXDAUEDACE (BKC) 20 76 111 67 17 80
20.75 111.07 17.00
38 STOCE/ (SD) 10.02 54.34 7.29
40 DETECTIONIT (25D) 21 86 108 68 14 58

BACKGROUND CONCENTRATIONS IN He FOR DETECTION LIMIT DETERMINATION

Table 2B

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5.1550

F G E A 8 С D 1 2 N2 02 25% Ar H2 10% H2 100% Н 3 4 STDOONC 5.0060 86.2740 5.1220 1.0000 2.5980 2.5980 5 6 7 7-Jul-88 avc and 3/CD and 3 C CONC: 8 TIME 0.9930 9 PM 5.30 2.4900 5.1275 2.4700 4.9200 10 7.00 2.5080 0.9840 5.1450 2.4900 4,9300 11 7.30 5.1400 0.9790 2.4980 2.4700 4.9300 12 8.00 2.5070 5.1450 0.9820 2.4700 4.9400 13 5.1500 8.30 2.5160 2.4800 4.9200 0.9840 14 8-Jul-88 0.9950 15 AM 8.00 5.1450 2 4910 2.4600 4.9200 16 9.00 5.1500 1.0120 2.4930 2.4600 4.9200 17 5.1450 1.0230 10.15 2.4920 2.4600 4.9100 18 11.15 1.0310 2.4920 2.4700 4.9000 5.1550 19 PM 12.00 1.0340 5.1375 2.4970 2.4500 4,9100 20 1.00 5.1500 1.0360 2.4990 4.9300 2.4600 21 2.10 1.0360 5.1525 2.4970 4.9100 2.4800 22 3.15 2.4990 5.1550 1.0360 4,9300 2.4600 23 4.00 5.1550 1.0320 2.4960 2.4800 4.9100 24 25 26 EVPAVERAGE 2.4982 5.1466 1 0113 2.4686 4,9200 27 28 STD DEV 0.0078 0.0236 0.0075 0.0110 0.0111 29 3 0 MAX READOUT 2.5160 5.1550 1.0360 2.4900 4.9400 31 3 2 MIN READOUT 2.4900 2.4500 4.9000 5.1275 0.9790 33 3 4 INI READOUT 5.1275 0.9930 2.4900 2.4700 4.9200 35 3 6 FIN READOUT 1.0320

DRIFT STUDY: READOUT OF THE STD GAS MIXTURE IN N2 WITH BKG CORRECTIONS

Table 3

4,9100

2.4960

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DRIFT STUDY: READOUT OF STD GAS MIXTURE IN He WITH BKG CORRECTIONS

	Α	8	С	D	E	F	G
1							
2		H2 10%	H2 100%	He	N2	02 25%	Ar
3							
4	STDCONC	7.0580	7.0580	88.6460	0.0000	3.2970	0.9990
5							
6							
7	JULY 5,88		CONC	<u> </u>		anc	2/20
8	TIME	<u> </u>					
9	AM 3.45	7.0880	7.0500			3.1900	1.0360
10	10.15	7.0640	7.0200			3.1775	1.0470
11	11.00	7.0940	7.0500			3.1850	1.0060
12	11.30	7.1170	7.0700			3.1975	0.9973
13	PM 12.20	7.0670	7.0100			3.2100	0.9660
14	1.00	7.0590	7.0100			3.2025	0.9580
15	1.30	7.0670	7.0200			3.2075	0.9590
16	2.30	7.0610	7.0100		ļ	3.2125	0 9720
17	3.00	7.0420	6.9900			3.2125	0.9720
18	3.30	7.0270	6.9900			3.2150	0.9350
19	4.00	7.0260	7.0000			3.2150	0.9.30
20	4.30	7.0250	6.9700	L		3.2175	0.9510
21	JULY 6,88			ļ			
22	AM 8.00	7.0030	6.9600		·	3.2450	1.0870
23	8.30	7.0010	6.9500			3.2475	1.0970
24	9.00	6.9940	6.9500			3.2500	1.1010
25	10 00	6.9970	6.9200			3.2500	1.10 0
1	<u>+</u>	······································		<u></u>			
20	END VERVER					2 2147	1 0102
20	EVP AVEFAGE	7.0458	6.9981	· • • • • •	-	3.2147	1.0103
120	STDODY	0.0373				0.0220	0.0600
11	SIDUEV	0.0373	0.0409		+	0.0229	0.0000
1 1 2	MAX READOUT	7 1170	7 0500			3 2500	1 1070
133		7.1170	7.0500			3.2300	
3 4	MINREADOUT	6 9940	6 9200		+	3 1775	0 9350
3 5			0.3250	+			
36	IN READOUT	7 0880	7 0500		+	3 1900	1.0360
37	1		1.0300				T
38	FIN READOUT	6 9970	6 9200		- 	3.2503	1 1070

Table 5A

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RESPONSE TIME, N2 GAS PULSE IN He (90% PEAK DETECTION FROM BASELINE)

Table 5B

RESPONSE TIME, He GAS PULSE IN N2 (90% PEAK DETECTION FROM BASELINE)



ILLUSTRATIONS





Figure 1

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Typical MGA-1200 Inldt System

Figure 2A



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Idealized Operation of a Hagnetic Sector Mass Spectrometer

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Figure 2B





Figure 3

164



DRIFT TEST SETUP

Figure 4

165



Figure 5

RESPONSE TIME SETUP

166

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LINEAR CURVE H2/N2 10%



LINEAR CURVE H2/N2 100%





LINEAR CURVE He/N2





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Figure 6E







171





Figure 8B











DRIFT STUDY : STD GAS MIXTURE IN N2 WITH BKG CORRECTION

DRIFT STUDY : STD GAS MIXTURE IN N2 WITH BKG CORRECTION



Figure 10B















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Figure 12

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