

TECHNICAL MANUAL

Part number : RAA 009 A Ind. B



· **REVISIONS**

INDEX	P/N REVISION	REVISION	SECTION	DATE
А		Creation	all	30/03/98
В	RAH 581 AA	Miscellaneous corrections	1, 3, 4	07/10/98



- 1. HYDRAULIC & PNEUMATIC PRINCIPLES
- 2. ELECTRIC & ELECTRONIC PRINCIPLES
- 3. MAINTENANCE PROCEDURES
- 4. OUTPUT FORMAT
- 5. TRAINING SLIDES

CONTENTS

1. GENERALITIES	. 2
2. MICROS 60 OT HYDRAULIC	
2.1.1. Transmission tubes list	. 4
2.1.2. Hydropneumatic connections	
2.2. Pack 2.2.1. Transmission tubes list	
2.2.1. Harshission tubes list	
2.3. Hydraulic cycle description	
2.3.1. Atmosphere circuit	
2.3.2. Diluent circuit	
2.3.3. Clean circuit	
2.3.4. Lyse circuit	
2.3.5. WBC/RBC counting circuit	
	. 0
3. MICROS 60 CT HYDRAULIC	15
3.1. With bottles	-
3.1.1. Transmission tubes list	
3.1.2. Hydropneumatic connections	
3.2. Pack 3.2.1. Transmission tubes list	
3.2.2. Hydropneumatic connections	
3.3. Hydraulic cycle description	
3.3.1. Atmosphere circuit	
3.3.2. Diluent circuit	
3.3.3. Clean circuit	
3.3.4. Lyse circuit	
3.3.5. WBC/RBC counting circuit	
	13
4. PNEUMATIC DIAGRAMS	26
4.1. Micros 60 CT bottle version	
4.2. Micros 60 CT pack version	
4.3. Micros 60 OT bottle version	
4.4. Micros 60 OT pack version	26

1. GENERALITIES

MICROS 60 instrument has been designed for simple mechanical operations.

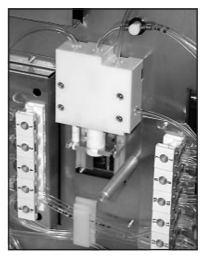
4 stepper motors provide movements to mechanical assemblies.

Pressure and vacuum are provided by the vacuum/waste syringe up and down movements (diag 1).

Liquid movements are achieved either by means of mechanical assembly movements (liquid



Diag.1



Diag.2

syringes diag 2) or pressure/vacuum and simultaneous action of specific valves.

• Dilution chambers (Diag .3)

WBC/HGB and RBC chambers are made of GRILAMID TR55 LY injected. The diode and the cell of the spectrophotometer are glued on the WBC/HGB chamber. Chamber positions can be modified in order to obtain the best sampling position possible.

Dilutions :

First dilution is carried out in the WBC/HGB chamber (with a bubbling phasis). The RBC blood sample is aspirated from this dilution. Lyse is sent from the drain nipple of the WBC/HGB chamber. Counts have a duration of 2×6 seconds.

(see procedures RAS 188 A and RAS 187 A for cycle hydraulic details)



Rinse :

To obtain the best rinse in the counting heads, diluent is sent from the liquid syringes. This is carried out before, between and after the two counts.



A window on the HGB/WBC chamber allows the needle to move down into the chamber and to inject reagents. As important light or variation of light can cause HGB result drifts, close instrument cover and door before running blood analyses.

Bubbling:

Insulators avoid polluted liquid overflows during bubbling phasis. they also allows an accurate adjustment of the bubbling volume.

MICROS 60 CT specifics :

- The piercing needle is equipped with two injectors to obtain a homogeneous diluent flow during needle rinsing phasis (see procedures RAS 188 A and RAS 187 A).

- Atmosphere is provided to sample tubes to allow a correct aspiration of blood.

2. MICROS 60 OT HYDRAULIC

Liquid circuits, hydropneumatic connections, as well as the transmission tubes used, are described in the following chart tables.

2.1. With bottles

DESIGNATION	PART NUMBER	DIAMETER	LENGTH	QUANTITY
SLEEVE HPS3	DBD005A	5-9		0.5
T CONNECTOR	EAB006B	2.3		3
T CONNECTOR	EAB032A	1.5		1
TUBE CAP	EAC017A	2.5		1
TYGON TUBE 0.051"	EAE006A	1.30	140	1
TYGON TUBE 0.051"	EAE006A	1.30	350	1
TYGON TUBE 0.060"	EAE007A	1.52	20	1
TYGON TUBE 0.060"	EAE007A	1.52	40	3
TYGON TUBE 0.060"	EAE007A	1.52	50	1
TYGON TUBE 0.060"	EAE007A	1.52	60	1
TYGON TUBE 0.060"	EAE007A	1.52	70	2
TYGON TUBE 0.060"	EAE007A	1.52	80	1
TYGON TUBE 0.060"	EAE007A	1.52	150	1
TYGON TUBE 0.060"	EAE007A	1.52	170	1
TYGON TUBE 0.060"	EAE007A	1.52	220	1
TYGON TUBE 0.060"	EAE007A	1.52	240	1
TYGON TUBE 0.060"	EAE007A	1.52	300	1
TYGON TUBE 0.060"	EAE007A	1.52	370	1
TYGON TUBE 0.060"	EAE007A	1.52	410	1
TYGON TUBE 0.060"	EAE007A	1.52	450	2
TYGON TUBE 0.060"	EAE007A	1.52	480	1
TYGON TUBE 0.081"	EAE008A	2.05	20	1
TYGON TUBE 0.081"	EAE008A	2.05	35	1
TYGON TUBE 0.081"	EAE008A	2.05	200	1
TYGON TUBE 0.081"	EAE008A	2.05	330	1
TYGON TUBE 0.081"	EAE008A	2.05	1080	1
TYGON TUBE 0.090"	EAE009A	2.28	20	2
TYGON TUBE 0.090"	EAE009A	2.28	50	1
TYGON TUBE 0.090"	EAE009A	2.28	60	2
TYGON TUBE 0.090"	EAE009A	2.28	120	1
TYGON TUBE 0.090"	EAE009A	2.28	140	1
TYGON TUBE 0.090"	EAE009A	2.28	150	1
TYGON TUBE 0.090"	EAE009A	2.28	190	1
BLUE TYGON TUBE 0.090"	EAE036A	2.28	1100	1
SLEEVE	GAL098A			30
TUBE SHIELD	GBC088A	4.4	30	1
TUBE SHIELD	GBC088A	4.4	60	1

2.1.1. Transmission tubes list

2.1.2. Hydropneumatic connections

CIRCUIT	FROM	SLEEVE	DIAM	LENGTH	то	SLEEVE
	(atmosphere)		2.28	190	Liquid valve 2_2	
AIR	Liquid valve 2_1	Y	2.28	50	Waste-chamb1	Y
	Waste-chamb2		2.28	20	Cap EAC017A	
	Diluent input		2.05	330	Liquid valve11_1	
	Liquid valve11_3	Y	1.52	220	Temp. sensor xba281a	Y
	Temp. sensor. xba281a	Y	1.52	40	Liquid syringes_3	Y
	Liquid valve11_2	Y	1.52	40	Liquid valve10_3	Y
	Liquid valve10_1	Y	1.52	70	Liquid valve7_3	Y
	Liquid valve7_1	Y	1.52	370	needle rinsing block2	
DILUENT	Needle rinsing block1		1.52	410	Liquid valve8_1	
	Liquid valve8_2	Y	1.52	300	Waste-chamb3	
	Liquid valve 7_2	Y	1.52	240		
	Té 2.3 2		1.52	50	Bac WBC/HGB 3	
	 Liquid valve10_2	Y	1.30	140	 Liquid syringes_1	Y
	Liquid syringes_2	Y	1.30	350	needle 1	
	Liquid syringes_2	Y			Needle 1	Y
	Clean bottle		2.28	1100 blue	 Liquid valve4_2	Y
CLEAN	Liguid valve4 1	Y	1.52	450	T connector 2.3 3	
	Lyse bottle		2.05	1080	liquid valve 1 1	
	Liquid valve1 3		1.52	150	Liquid syringes 4	
LYSE	Liquid valve1 2		1.52	480	WBC grounding connector	
	WBC grounding connector		1.52	20	T connector 1.5_1	
	WBC/HGB_2 chamber		1.52	170	RBC 3 chamber	
WBC/RBC	RBC 2 chamber		1.52	450	liquid valve 6 2	Y
COUNTING	Liquid valve 6 1	Y	1.52	60	Waste-chamb. 4	
	WBC/HGB_1 chamber		1.52	40	T connector 1.5 2	
	WBC/HGB_1 chamber		gbc088a	30	T connector 1.5 2	
	T connector 1.5_3		1.52	80	Insulator WBC_1	
	Insulator WBC_2		2.28	120	liquid valve12_2	Y
	Liquid valve12_1	Y	2.05	35	T connector 2.3_1	
	RBC_1 chamber		1.52	70	Insulator RBC_1	
	RBC_1 chamber		gbc088a	60	Insulator RBC_1	
DRAIN/BUBBLING	Insulator RBC_2		2.28	150	liquid valve13_2	Y
	 Liquid valve13_1	Y	2.05	20	T connector 2.3_2	
	T connector 2.3_3		2.05	200	T connector 2.3_2	
	T connector 2.3_3		2.28	20	Cell xba199a	Y
	 Cell xba199a	Y	2.28	60	Waste-chamb5	Y
	T connector 2.3_1		2.28	60	Liquid valve5_2	Y
	Liquid valve5 1	1	2.28	140	Waste ouput	



Read this table as follows in this example : The Liquid valve 7_2 corresponds to the ouput 2 of the valve number 7 (see attached pneumatic diagram).

2.2. Pack

2.2.1. Transmission tubes list

DESIGNATION	PART NUMBER	DIAMETER	LENGTH	QUANTITY
SLEEVE HPS3	DBD005A	5-9		0.5
T CONNECTOR	EAB006B	2.3		3
T CONNECTOR	EAB032A	1.5		1
TUBE CAP	EAC017A	2.5		1
TYGON TUBE 0.051"	EAE006A	1.30	140	1
TYGON TUBE 0.051"	EAE006A	1.30	350	1
TYGON TUBE 0.060"	EAE007A	1.52	20	1
TYGON TUBE 0.060"	EAE007A	1.52	40	3
TYGON TUBE 0.060"	EAE007A	1.52	50	1
TYGON TUBE 0.060"	EAE007A	1.52	60	1
TYGON TUBE 0.060"	EAE007A	1.52	70	2
TYGON TUBE 0.060"	EAE007A	1.52	80	1
TYGON TUBE 0.060"	EAE007A	1.52	150	1
TYGON TUBE 0.060"	EAE007A	1.52	170	1
TYGON TUBE 0.060"	EAE007A	1.52	220	1
TYGON TUBE 0.060"	EAE007A	1.52	240	1
TYGON TUBE 0.060"	EAE007A	1.52	300	1
TYGON TUBE 0.060"	EAE007A	1.52	370	1
TYGON TUBE 0.060"	EAE007A	1.52	410	1
TYGON TUBE 0.060"	EAE007A	1.52	450	2
TYGON TUBE 0.060"	EAE007A	1.52	480	1
TYGON TUBE 0.081"	EAE008A	2.05	20	1
TYGON TUBE 0.081"	EAE008A	2.05	35	1
TYGON TUBE 0.081"	EAE008A	2.05	200	1
TYGON TUBE 0.081"	EAE008A	2.05	590	1
TYGON TUBE 0.090"	EAE009A	2.28	20	2
TYGON TUBE 0.090"	EAE009A	2.28	50	1
TYGON TUBE 0.090"	EAE009A	2.28	60	2
TYGON TUBE 0.090"	EAE009A	2.28	120	1
TYGON TUBE 0.090"	EAE009A	2.28	150	1
TYGON TUBE 0.090"	EAE009A	2.28	190	1
TYGON TUBE 0.090"	EAE009A	2.28	510	1
TYGON TUBE 0.090"	EAE009A	2.28	550	1
TYGON TUBE 0.090"	EAE009A	2.28	1100	1
SLEEVE	GAL098A			31
TUBE SHIELD	GBC088A	4.4	30	1
TUBE SHIELD	GBC088A	4.4	60	1
METALLIC SHEATH	GBC170A	5.2	35	3

2.2.2. Hydropneumatic connections

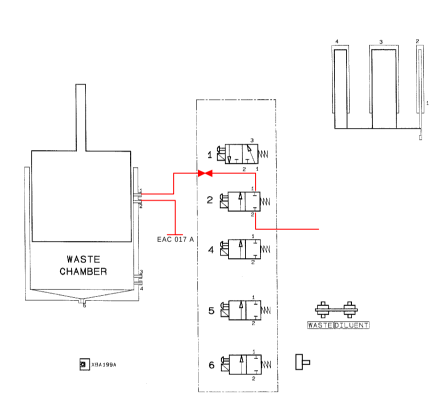
CIRCUIT	FROM	SLEEVE	DIAMETER	LENGTH	ТО	SLEEVE
	(atmosphere)		2.28	190	Liquid valve 2_2	
AIR	Liquid valve 2_1	Y	2.28	50	Waste-chamb1	Y
	Waste-chamb2		2.28	20	cap EAC017A	
	Pack_3 (Diluent)		2.28	550	Liquid valve 11_1	Y
	Pack_3 (Diluent)		gbc170a			
	Liquid valve 11_3	Y	1.52	220	temp. sensor xba281a	Y
	temp. sensor xba281a	Y	1.52	40	Liquid syringes _3	Y
	Liquid valve 11_2	Y	1.52	40	Liquid valve 10_3	Y
	Liquid valve 10_1	Y	1.52	70	Liquid valve 7_3	Y
	Liquid valve 7_1	Y	1.52	370	Needle rinsing block_2	
DILUENT	needle rinsing block1		1.52	410	Liquid valve 8_1	
	Liquid valve 8_2	Y	1.52	300	Waste-chamb3	
	Liquid valve 7_2	Y	1.52	240	T connector 2.3_1	
	T connector 2.3_2		1.52	50	Bac WBC/HGB_3	
	Liquid valve 10_2	Y	1.30	140	Liquid syringes _1	Y
	Liquid syringes _2	Y	1.30	350	NEEDLE_1	
	Liquid syringes _2	Y			NEEDLE_1	Y
	Pack-1 (Clean)		2.28	510	Liquid valve 4_2	Y
CLEAN	Pack-1 (Clean)		gbc170a			
	Liquid valve 4_1	Y	1.52	450	T connector 2.3_3	
	Pack_2 (Lyse)		2.05	590	Liquid valve 1_1	
	Pack_2 (Lyse)		gbc170a			
LYSE	Liquid valve 1_3		1.52	150	Liquid syringes _4	
	Liquid valve 1_2		1.52	480	WBC grounding connector	
	WBC grounding connector		1.52	20	T connector 1.5_1	
	WBC/HGB chamber _2		1.52	170	RBC chamber _3	
WBC/RBC	RBC chamber _2		1.52	450	Liquid valve 6_2	Y
counting	Liquid valve 6_1	Y	1.52	60	Waste-chamb4	
	WBC/HGB chamber _1		1.52	40	T connector 1.5_2	
	WBC/HGB chamber _1		gbc088a	30	T connector 1.5_2	
	T connector 1.5_3		1.52	80	Insulator WBC_1	
	Insulator WBC_2		2.28	120	Liquid valve 12_2	Y
	Liquid valve 12_1	Y	2.05	35	T connector 2.3_1	
	Bac RBC_1		1.52	70	Insulator RBC_1	
DRAIN /	Bac RBC_1		gbc088a	60	Insulator RBC_1	
BUBBLING	Insulator RBC_2		2.28	150	Liquid valve 13_2	Y
	Liquid valve 13_1	Y	2.05	20	T connector 2.3_2	
	T connector 2.3_3		2.05	200	T connector 2.3_2	
	T connector 2.3_3		2.28	20	Cellule xba199a	Y
	Cell xba199a	Y	2.28	60	Waste-chamb5	Y
	T connector 2.3_1		2.28	60	Liquid valve 5_2	Y
	Liquid valve 5_1		2.28	1100	Pack_4 (Waste)	

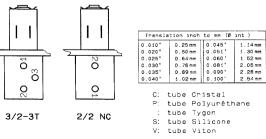
Read this table as follows in this example :

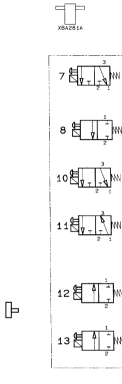
The Liquid valve 7_2 corresponds to ouput 2 of the valve number 7 (see attached pneumatic diagram).

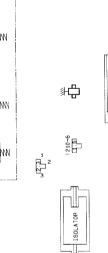
- 2.3. Hydraulic cycle description
 - 2.3.1. Atmosphere circuit
 2.3.2. Diluent circuit
 2.3.3. Clean circuit
 2.3.4. Lyse circuit
 2.3.5. WBC/RBC counting circuit
 2.3.6. Drain/bubbling circuit

ATMOSPHERE CIRCUIT







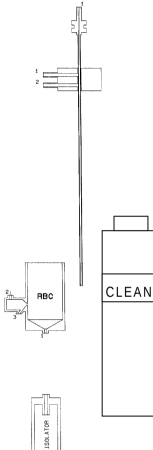


MM

WBC HGB

 $\geq \psi$

<u>"</u>ф



П,

LYSE



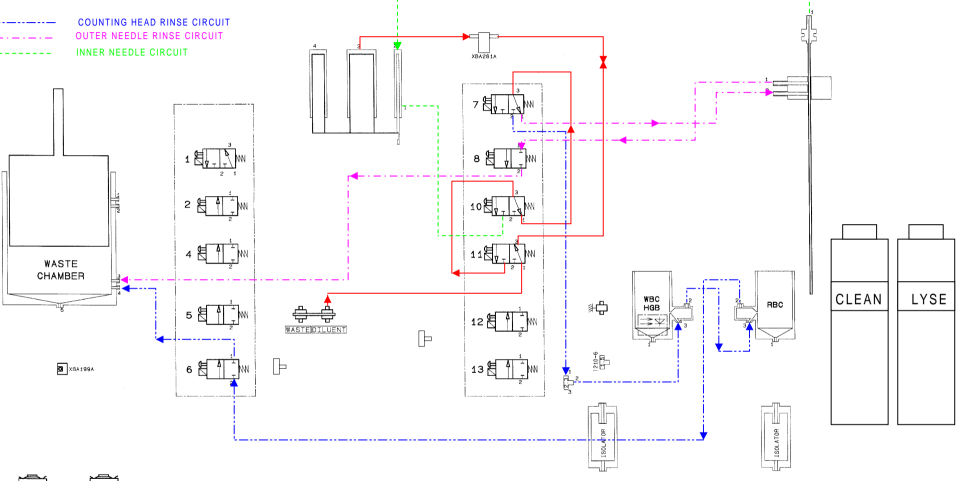
RAA 009

A

ind

B

DILUENT CIRCUIT

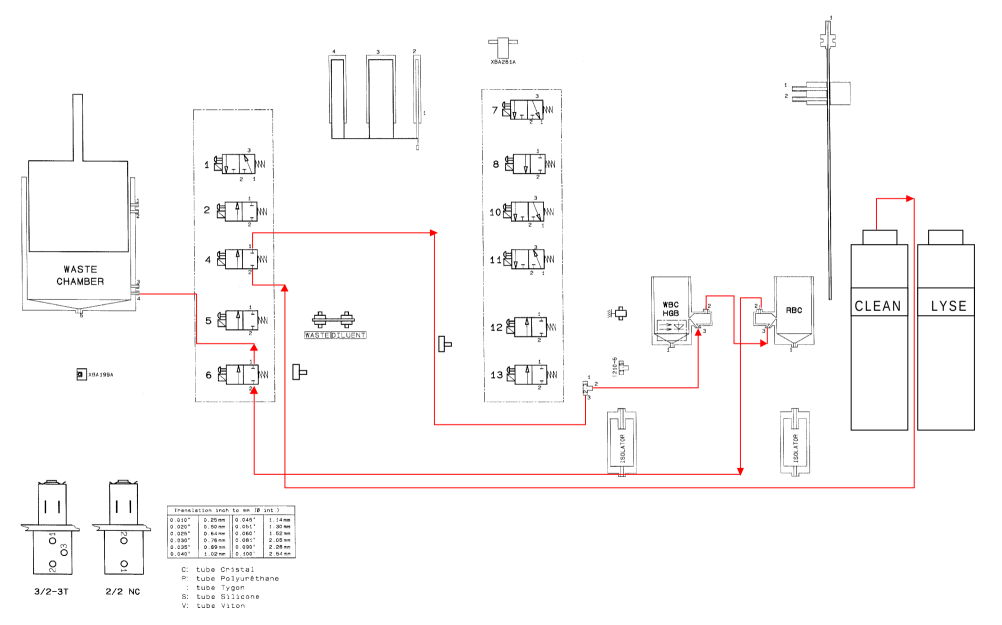


| | 11 Translation inch to mm (Ø int.) 0.010" 0.020" 0.025" 0.030" 0.035" 0.040" 0301 0 0 20 C: D 3/2-3T 2/2 NC S:

	cion inch	CC 1111 10	1116.7
010"	0.25 mm	0.045°	1.14 mm
020"	0.50 mm	0.051	1.30 mm
025"	0.64mm	0.060'	1.52 mm
030"	0.76 mm	0.081"	2.05 mm
035"	0.89 mm	0.090"	2.28 mm
040 *	1.02 mm	0.100"	2.54 mm
P: 1 : 1 S: 1		lyuréth gon licone	iane

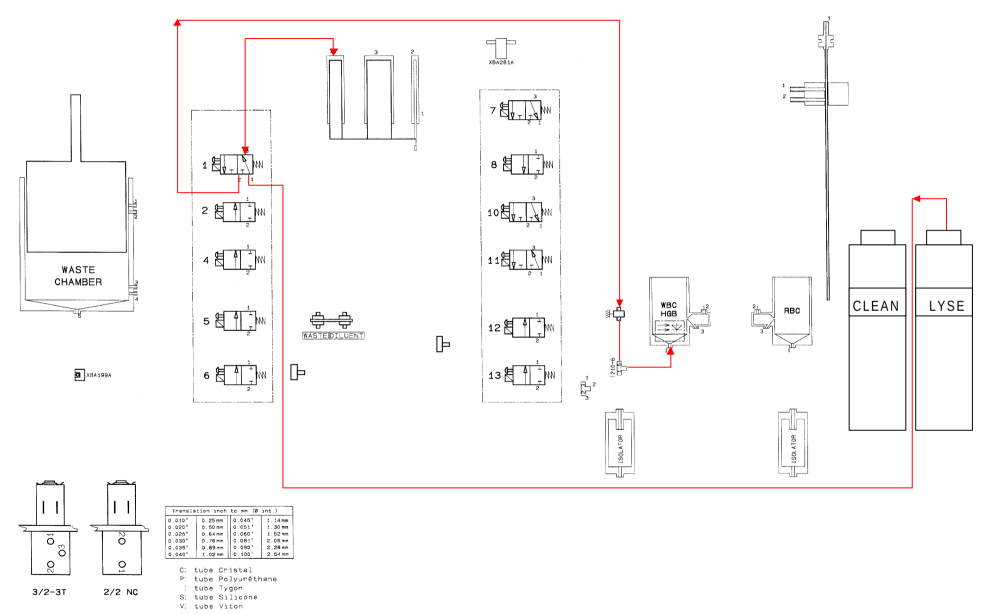
RAA 009 A ind B

CLEAN CIRCUIT

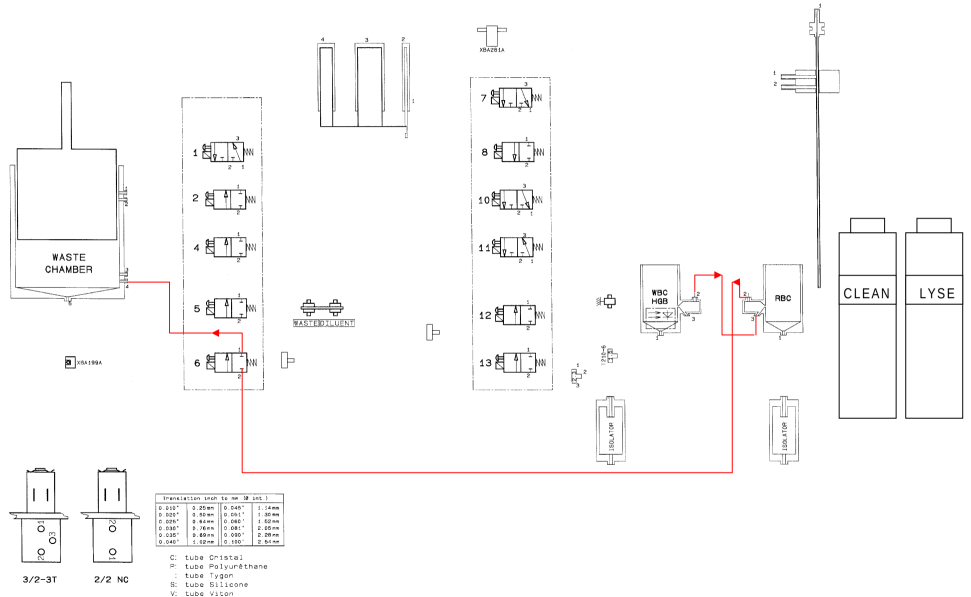


RAA 009 A ind B

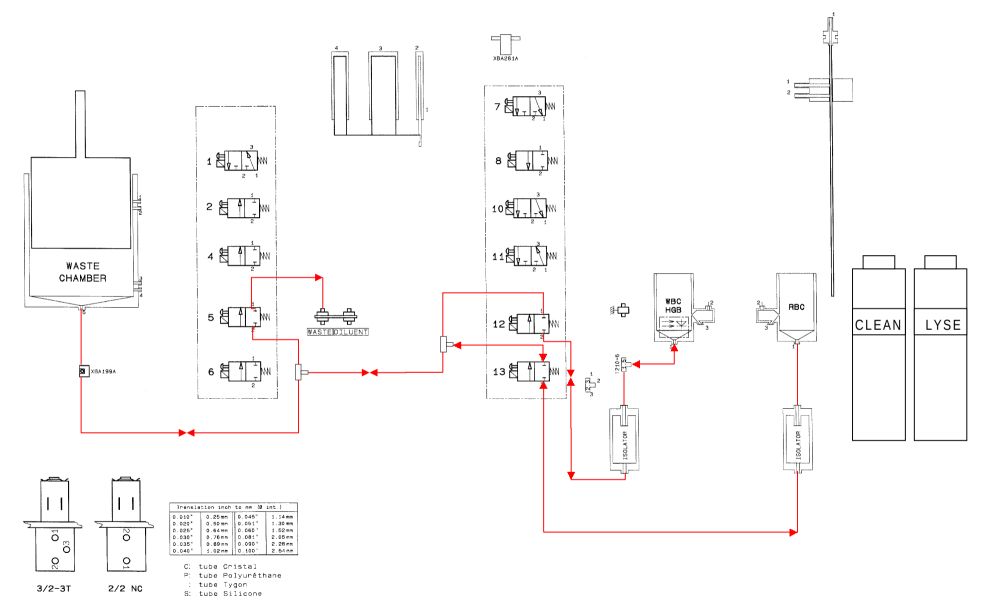
LYSE CIRCUIT



WBC / RBC COUNTING PHASIS



DRAIN / BUBBLING CIRCUIT



V: tube Viton

3/2-3T

2/2 NC

3. MICROS 60 CT HYDRAULIC

3.1. With bottles

3.1.1. Transmission tubes list

DESIGNATION	PART NUMBER	DIAMETER	LENGTH	QUANTITY
T CONNECTOR	EAB006B	2.3		4
STRAIGHT CONNECTOR	EAB015A	1.5/2.5		3
T CONNECTOR	EAB032A	1.5		2
TUBE CAP	EAC017A	2.5		1
TYGON TUBE 0.040"	EAE005A	1.02	110	1
TYGON TUBE 0.040"	EAE005A	1.02	335	1
TYGON TUBE 0.060"	EAE007A	1.52	15	3
TYGON TUBE 0.060"	EAE007A	1.52	20	2
TYGON TUBE 0.060"	EAE007A	1.52	40	4
TYGON TUBE 0.060"	EAE007A	1.52	50	2
TYGON TUBE 0.060"	EAE007A	1.52	60	1
TYGON TUBE 0.060"	EAE007A	1.52	70	2
TYGON TUBE 0.060"	EAE007A	1.52	80	1
TYGON TUBE 0.060"	EAE007A	1.52	100	1
TYGON TUBE 0.060"	EAE007A	1.52	150	1
TYGON TUBE 0.060"	EAE007A	1.52	170	1
TYGON TUBE 0.060"	EAE007A	1.52	220	1
TYGON TUBE 0.060"	EAE007A	1.52	240	1
TYGON TUBE 0.060"	EAE007A	1.52	420	2
TYGON TUBE 0.060"	EAE007A	1.52	450	2
TYGON TUBE 0.060"	EAE007A	1.52	480	1
TYGON TUBE 0.081"	EAE008A	2.05	20	1
TYGON TUBE 0.081"	EAE008A	2.05	35	1
TYGON TUBE 0.081"	EAE008A	2.05	200	1
TYGON TUBE 0.081"	EAE008A	2.05	330	1
TYGON TUBE 0.081"	EAE008A	2.05	650	1
TYGON TUBE 0.081"	EAE008A	2.05	1080	1
TYGON TUBE 0.090"	EAE009A	2.28	20	2
TYGON TUBE 0.090"	EAE009A	2.28	50	1
TYGON TUBE 0.090"	EAE009A	2.28	60	3
TYGON TUBE 0.090"	EAE009A	2.28	120	1
TYGON TUBE 0.090"	EAE009A	2.28	140	1
TYGON TUBE 0.090"	EAE009A	2.28	150	1
TYGON TUBE 0.090"	EAE009A	2.28	190	1
SILICONE TUBE	EAE025A	1.5/3.5	50	2
BLUE TYGON TUBE 0.090"	EAE036A	2.28	1100	1
SLEEVE	GAL098A			32
TUBE SHIELD	GBC088A	4.4	30	1
TUBE SHIELD	GBC088A	4.4	60	1

3.1.2. Hydropneumatic connections

CIRCUIT	FROM	SLEEVE	DIAMETER	LENGTH	то	SLEEVE
	(atmosphere)		2.28	190	Liquid valve 2_2	
	Liquid valve 2_1	Y	2.28	50	Waste-chamb1	Y
AIR	Waste-chamb2		2.28	20	cap EAC017A	
	(atmosphere)		2.28	60	Liquid valve 3_2	
	Liquid valve 3_1		1.52	40	T connector 2.3_1	
	Diluent input		2.05	330	Liquid valve 11_1	
	Liquid valve 11_3	Y	1.52	220	temp sensor. xba281a	Y
	temp sensor. xba281a	Y	1.52	40	liquid syringes_3	Y
	Liquid valve 11_2	Y	1.52	40	Liquid valve 10_3	Y
	Liquid valve 10_1	Y	1.52	70	Liquid valve 7_3	Y
	Liquid valve 7_1	Y	1.52	50	Liquid valve 9_3	Y
	Liquid valve 9_1	Y	1.52	420	Needle rinsing block2	
	Liquid valve 9_2	Y	1.52	420	T connector 1.5 2	
	T connector 1.5 1		S1.5/3.5	50	 Needle rinsing block3	
	T connector 1.5_3		S1.5/3.5	50	Needle rinsing block4	
DILUENT	Liquid valve 7_2	Y	1.52	240	T connector 2.3 1	
2.202	T connector 2.3 2		1.52	50	WBC/HGB chamber 3	
	Liquid valve 10_2	Y	1.52	15	Connector 1.5/2.5	
	Connector 1.5/2.5		1.02	110	Connector 1.5/2.5	
	Connector 1.5/2.5		1.52	15	liquid syringes_1	Y
	liquid syringes_2	Y	1.52	15	Connector 1.5/2.5	· ·
	Connector 1.5/2.5		1.02	335	needle_1	
	Connector 1.5/2.5	Y	1.02	000	needle 1	Y
	Needle rinsing block1		1.52	100	T connector 2.3_3	· ·
	T connector 2.3_2		1.52	20	Liquid valve 8_1	
	Liquid valve 8_2		2.05	650	Waste-chamb3	
	Clean bottle		2.28		Liquid valve 4_2	Y
CLEAN	Liquid valve 4_1	Y	1.52	450	T connector 2.3_3	· ·
	Lyse bottle		2.05	1080	Liquid valve 1_1	
	Liquid valve 1_3		1.52	150	liquid syringes_4	
LYSE	Liquid valve 1_3		1.52	480	WBC grounding connector	
	WBC grounding connector		1.52	20	T connector 1.5_1	
	WBC/HGB chamber _2		1.52	170	RBC chamber 3	
WBC/RBC	RBC chamber_2		1.52	450	Liquid valve 6_2	Y
counting	Liquid valve 6_1	Y	1.52	430 60	Waste-chamb. 4	1
	WBC/HGB Chamber 1		1.52	40	T connector 1.5 2	
	WBC/HGB Chamber 1		gbc088a	30	T connector 1.5_2	
	T connector 1.5_3		1.52		insulator WBC_1	
	insulator WBC_2		2.28	120	Liquid valve 12_2	Y
	Liquid valve 12_1	Y	2.05	35	T connector 2.3_1	1
	RBC_1 chamber		1.52	70	insulator RBC_1	
DRAIN /	RBC_1 chamber		gbc088a	60	insulator RBC_1	
DRAIN / BUBBLING	insulator RBC_2		2.28	150	Liquid valve 13_2	Y
DODDLING	Liquid valve 13 1	Y	2.20	20	T connector 2.3_2	
	T connector 2.3_3					
			2.05	200	T connector 2.3_2	V
	T connector 2.3_3	Y	2.28	20	Cell xba199a Waste-chamb5	Y Y
	Cell xba199a		2.28	60 60		Y Y
	T connector 2.3_1		2.28	60	Liquid valve 5_2	r
	Liquid valve 5_1		2.28	140	Waste ouput	

NOTE

Read this table as follows in this example :

The Liquid valve 7_2 corresponds to ouput 2 of the valve number 7 (see attached pneumatic diagram).

3.2. Pack

3.2.1. Transmission tubes list

DESIGNATION	PART NUMBER	DIAMETER	LENGTH	QUANTITY
T CONNECTOR	EAB006B	2.3		4
STRAIGHT CONNECTOR	EAB015A	1.5/2.5		3
T CONNECTOR	EAB032A	1.5		2
TUBE CAP	EAC017A	2.5		1
TYGON TUBE 0.040"	EAE005A	1.02	110	1
TYGON TUBE 0.040"	EAE005A	1.02	335	1
TYGON TUBE 0.060"	EAE007A	1.52	15	3
TYGON TUBE 0.060"	EAE007A	1.52	20	2
TYGON TUBE 0.060"	EAE007A	1.52	40	4
TYGON TUBE 0.060"	EAE007A	1.52	50	2
TYGON TUBE 0.060"	EAE007A	1.52	60	1
TYGON TUBE 0.060"	EAE007A	1.52	70	2
TYGON TUBE 0.060"	EAE007A	1.52	80	1
TYGON TUBE 0.060"	EAE007A	1.52	100	1
TYGON TUBE 0.060"	EAE007A	1.52	150	1
TYGON TUBE 0.060"	EAE007A	1.52	170	1
TYGON TUBE 0.060"	EAE007A	1.52	220	1
TYGON TUBE 0.060"	EAE007A	1.52	240	1
TYGON TUBE 0.060"	EAE007A	1.52	420	2
TYGON TUBE 0.060"	EAE007A	1.52	450	2
TYGON TUBE 0.060"	EAE007A	1.52	480	1
TYGON TUBE 0.081"	EAE008A	2.05	20	1
TYGON TUBE 0.081"	EAE008A	2.05	35	1
TYGON TUBE 0.081"	EAE008A	2.05	200	1
TYGON TUBE 0.081"	EAE008A	2.05	590	1
TYGON TUBE 0.081"	EAE008A	2.05	650	1
TYGON TUBE 0.090"	EAE009A	2.28	20	2
TYGON TUBE 0.090"	EAE009A	2.28	50	1
TYGON TUBE 0.090"	EAE009A	2.28	60	3
TYGON TUBE 0.090"	EAE009A	2.28	120	1
TYGON TUBE 0.090"	EAE009A	2.28	150	1
TYGON TUBE 0.090"	EAE009A	2.28	190	1
TYGON TUBE 0.090"	EAE009A	2.28	510	1
TYGON TUBE 0.090"	EAE009A	2.28	550	1
TYGON TUBE 0.090"	EAE009A	2.28	1100	1
SILICONE TUBE	EAE025A	1.5/3.5	50	2
SLEEVE	GAL098A			33
TUBE SHIELD	GBC088A	4.4	30	1
TUBE SHIELD	GBC088A	4.4	60	1
METALLIC SHEATH	GBC170A	5.2	35	3

3.2.2. Hydropneumatic connections

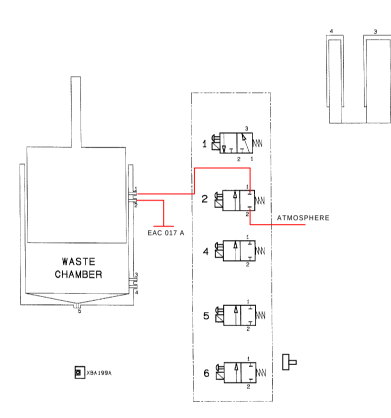
CIRCUIT	FROM	SLEEVE	DIAMETER	LENGTH	то	SLEEVE
	(atmosphere)		2.28	190	Liquid valve 2_2	
	Liquid valve 2_1	Y	2.28	50	Waste-chamb1	Y
AIR	Waste-chamb2		2.28	20	cap EAC017A	
	(atmosphere)		2.28	60	Liquid valve 3_2	
	Liquid valve 3_1		1.52	40	T Connector 2.3_1	
	Pack_3 (Diluent)		2.28	550	Liquid valve 11_1	Y
	Pack_3 (Diluent)		gbc170a			
	Liquid valve 11_3	Y	1.52	220	Temp. sensor xba281a	Y
	Temp. sensor xba281a	Y	1.52	40	Liquid syringes_3	Y
	Liquid valve 11_2	Y	1.52	40	Liquid valve 10_3	Y
	Liquid valve 10_1	Y	1.52	70	Liquid valve 7_3	Y
	Liquid valve 7_1	Y	1.52	50	Liquid valve 9_3	Y
	Liquid valve 9_1	Y	1.52	420	Needle rinsing block_2	
	Liquid valve 9_2	Ý	1.52	420	T Connector 1.5 2	
	T Connector 1.5 1		S1.5/3.5	50	Needle rinsing block_3	
	T Connector 1.5_3		S1.5/3.5	50	Needle rinsing block_4	
DILUENT	Liquid valve 7_2	Y	1.52	240	T Connector 2.3_1	
	T Connector 2.3_2		1.52	50	Bac WBC/HGB_3	
	Liquid valve 10_2	Y	1.52	15	connector1.5/2.5	
	connector1.5/2.5		1.02	110	connector1.5/2.5	
	connector 1.5/2.5		1.52	15	Liquid syringes_1	Y
	Liquid syringes_2	Y	1.52	15	connector 1.5/2.5	
	connector 1.5/2.5		1.02	335	Needle_1	
	connector 1.5/2.5	Y	1.02	335	Needle 1	Y
	Needle rinsing block_1	1	1.52	100	T Connector 2.3_3	I
	T Connector 2.3_2		1.52	20	Liquid valve 8_1	
	Liquid valve 8_2				Waste-chamb3	
	· –		2.05	650	_	Y
CLEAN	Pack_1 (Clean)		2.28	510	Liquid valve 4_2	ř
GLEAN	Pack_1 (Clean)	Y	gbc170a	450	T. Osansstan 0.0.0	
	Liquid valve 4_1	Ý	1.52	450	T Connector 2.3_3	
	Pack_2 (Lyse)		2.05	590	Liquid valve 1_1	
	Pack_2 (Lyse)		gbc170a	450	Linuid cominence A	
LYSE	Liquid valve 1_3		1.52	150	Liquid syringes_4	
	Liquid valve 1_2		1.52	480	WBCgrounding connec.	
	WBC grounding connect.		1.52	20	T Connector 1.5_1	
Comptage	Bac WBC/HGB_2		1.52	170	RBC chamber_3	
WBC/RBC	RBC chamber_2	N N	1.52	450	Liquid valve 6_2	Y
		Y	1.52	60	Waste-chamb4	
	Bac WBC/HGB_1		1.52	40	T Connector 1.5_2	
	Bac WBC/HGB_1		gbc088a	30	T Connector 1.5_2	
	T Connector 1.5_3		1.52	80	insulator WBC_1	
	insulator WBC_2		2.28	120	Liquid valve 12_2	Y
	Liquid valve 12_1	Y	2.05	35	T Connector 2.3_1	
	RBC chamber_1		1.52	70	insulator RBC_1	
			gbc088a	60	insulator RBC_1	
/ Bullage	insulator RBC_2		2.28	150	Liquid valve 13_2	Y
	Liquid valve 13_1	Y	2.05	20	T Connector 2.3_2	
	T Connector 2.3_3		2.05	200	T Connector 2.3_2	
	T Connector 2.3_3		2.28	20	CELL xba199a	Y
	CELL xba199a	Y	2.28	60	Waste-chamb5	Y
	T Connector 2.3_1		2.28	60	Liquid valve 5_2	Y
	Liquid valve 5_1		2.28	1100	Pack_4 (Waste)	

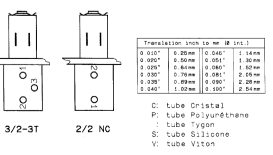
Read this table as follows in this example :

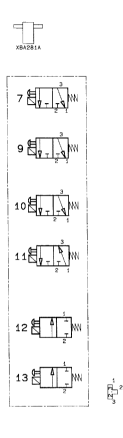
The Liquid valve 7_2 corresponds to ouput 2 of the valve number 7 (see attached pneumatic diagram)

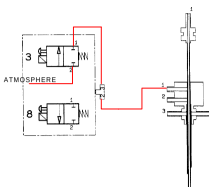
- 3.3. Hydraulic cycle description
 - 3.3.1. Atmosphere circuit
 3.3.2. Diluent circuit
 3.3.3. Clean circuit
 3.3.4. Lyse circuit
 3.3.5. WBC/RBC counting circuit
 - 3.3.6. Drain/bubbling circuit

ATMOSPHERE CIRCUIT





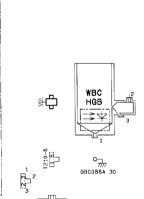




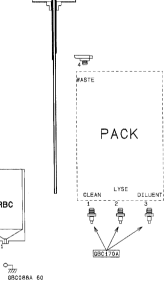
2

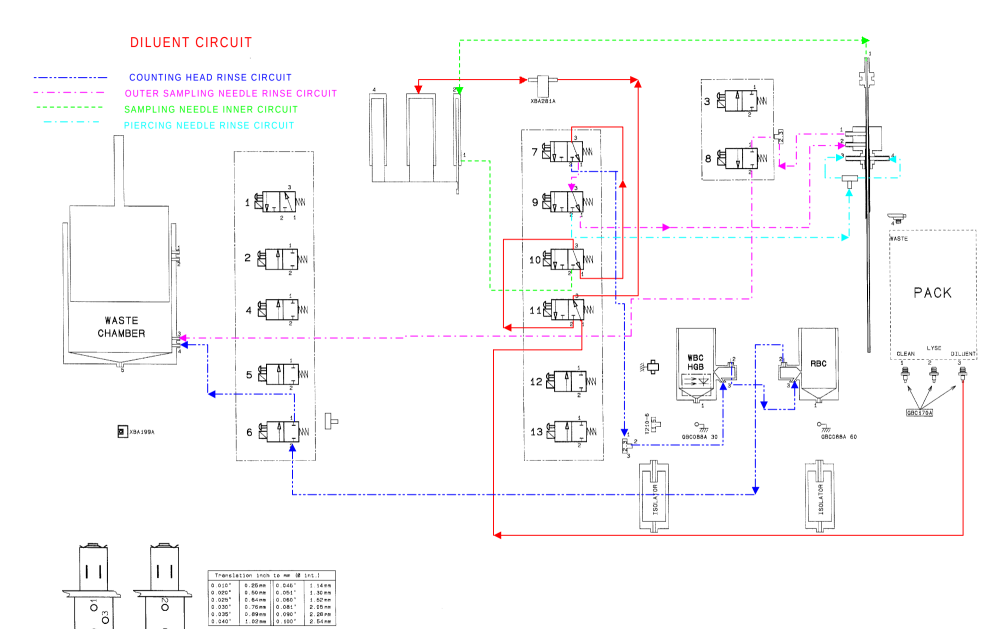
RBC

Ш ISOLATOR [



ISOLATOR





20

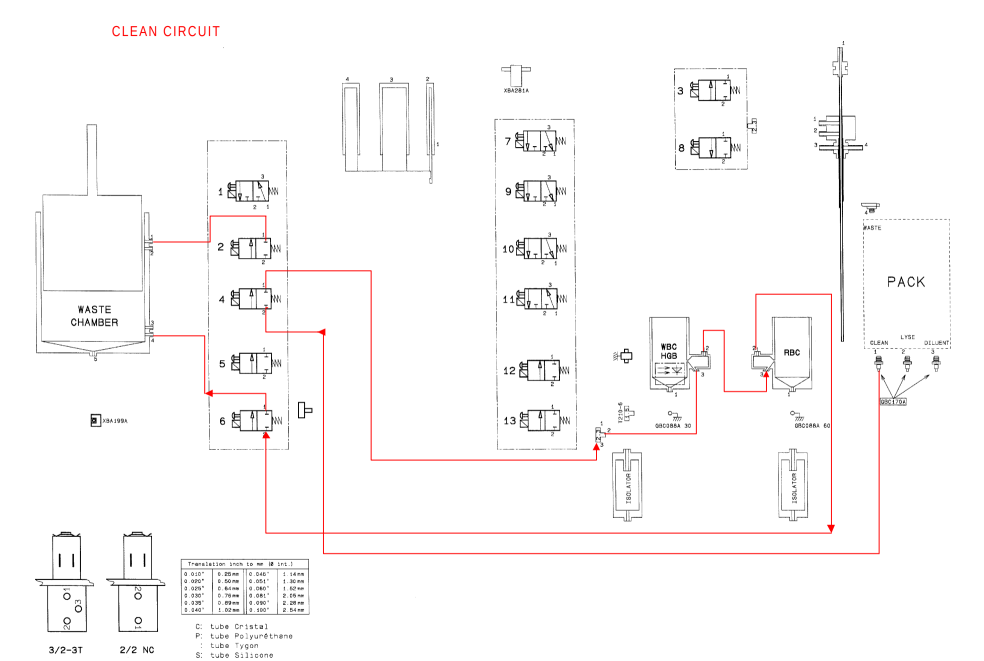
3/2-3T

0

2/2 NC

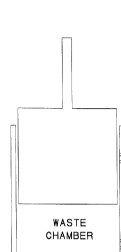
C: tube Cristal P: tube Polyuréthane : tube Tygon

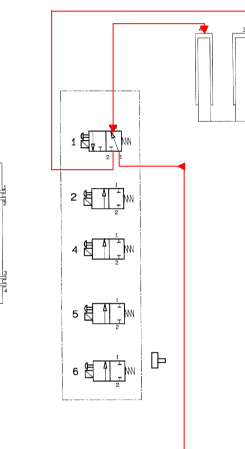
S: tube Silicone V: tube Viton

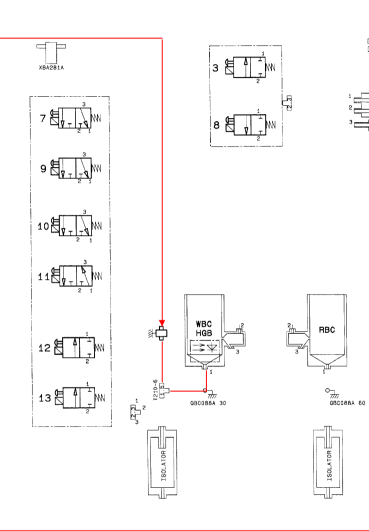


V: tube Viton

LYSE CIRCUIT







512

4 WASTE

PACK

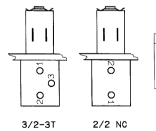
LYSE

۽ ا

GBC170A

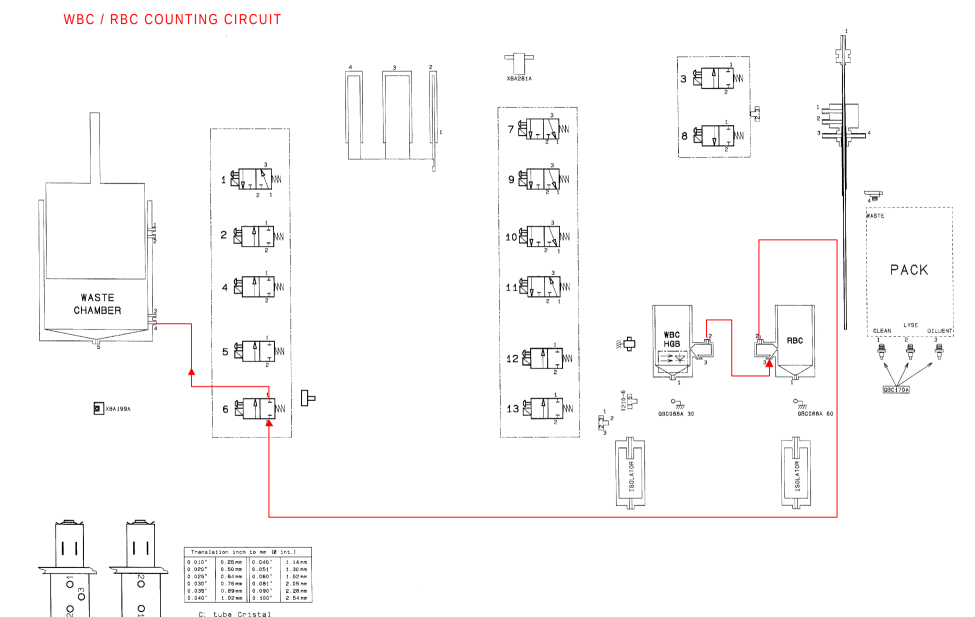
DILUENT

2



G XBA199A

Transla	tion inch	to mm (Ø	int.)
0.010"	0.25 mm	0.046"	1.14 mm
0.020	0.50 mm	0.051	1.30 mm
0.025*	0.64mm	0.060*	1.52 mm
0.030"	0.76 mm	0.081"	2.05 mm
0.035"	0.89mm	0.090*	2.28 mm
0.040"	1.02 mm	0.100"	2.54 mm
P: 1	tube Cr tube Po tube Ty tube Si	lyurét⊁ gon	ane

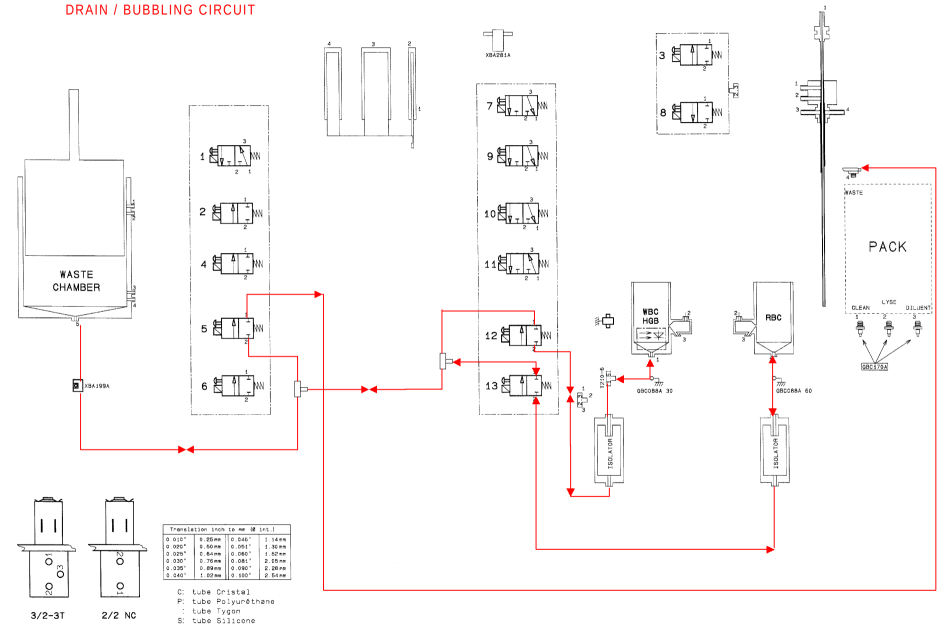


C: tube Cristal P: tube Polyuréthane : tube Tygon S: tube Silicone V: tube Viton

3/2-3T

2/2 NC

RAA 009 A ind B



Silicon
 V: tube Viton

4. PNEUMATIC DIAGRAMS

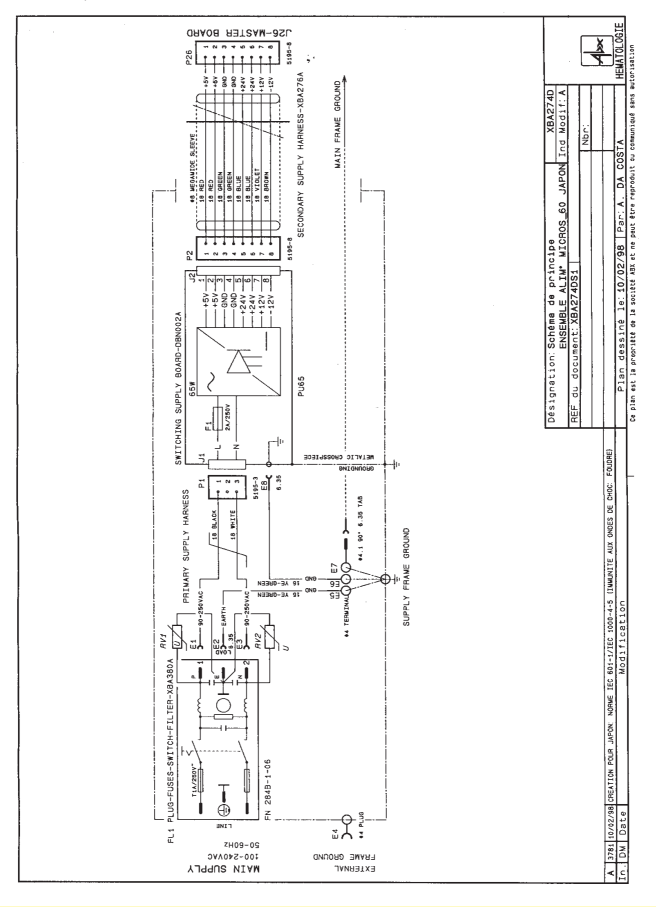
- 4.1. Micros 60 CT bottle version
- 4.2. Micros 60 CT pack version
- 4.3. Micros 60 OT bottle version
- 4.4. Micros 60 OT pack version

CONTENTS

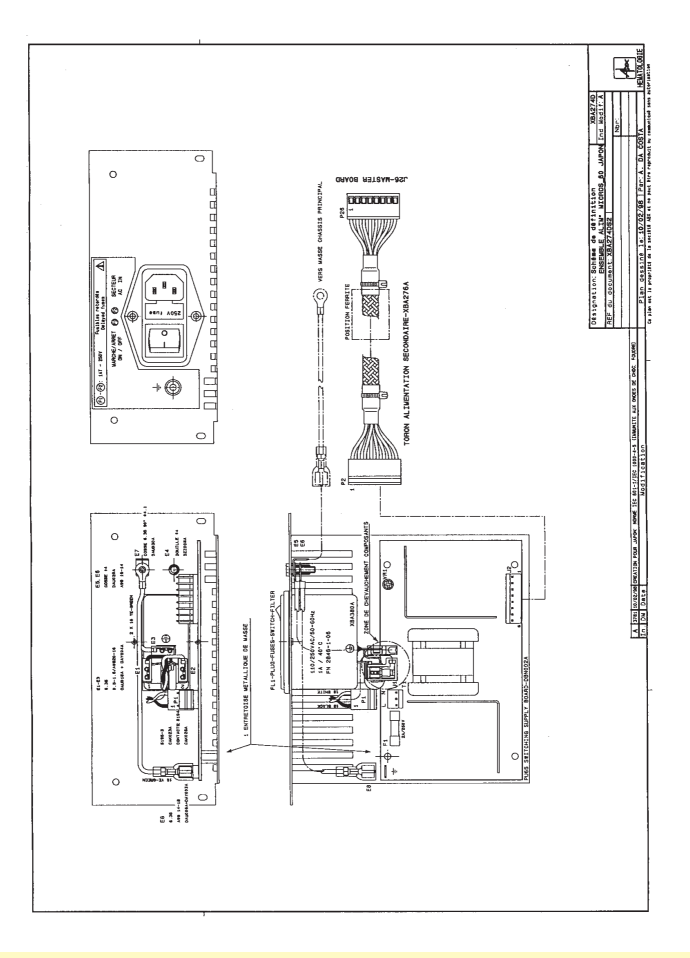
1. POWER SUPPLY ASSEMBLY	. 2
 1.1. Principle 1.2. Power supply fan 1.3. Secondary supply cable 1.4. Main supply filter 	. 4 . 5
2. MOTHER BOARD	. 7
2.1. Configuration	. 7
3. LCD BOARD	. 8
4. COAXIALS	. 9
4.1. RBC coaxial	
4.2. WBC coaxial	11
5. SENSORS	12
5.1. Drain detection sensor	
5.2. Carriage & needle sensors	
6. MISCELLANEOUS	
6.1. Needle carriage motor	
6.2. Barcode cable	
6.4. HGB Chamber assembly	
6.5. Chip card reader cable	
6.6. CT Piercing assembly cable	
6.7. CT twin valve cable	21
7. FLAT CABLES	22
8. ELECTRICAL SYNOPTICS	26

1. POWER SUPPLY ASSEMBLY

1.1. Principle

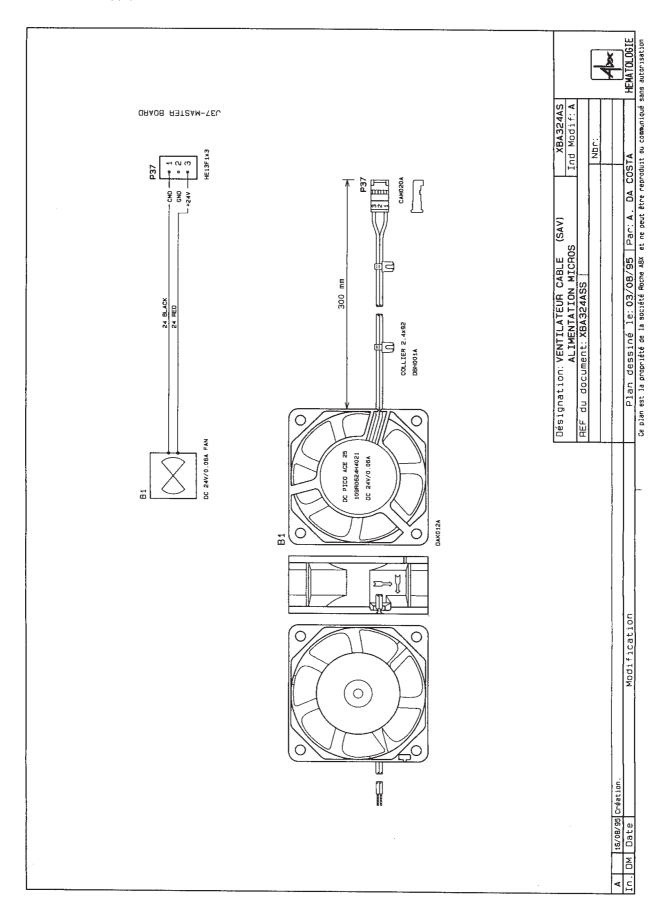


RAA 009 A Ind.A

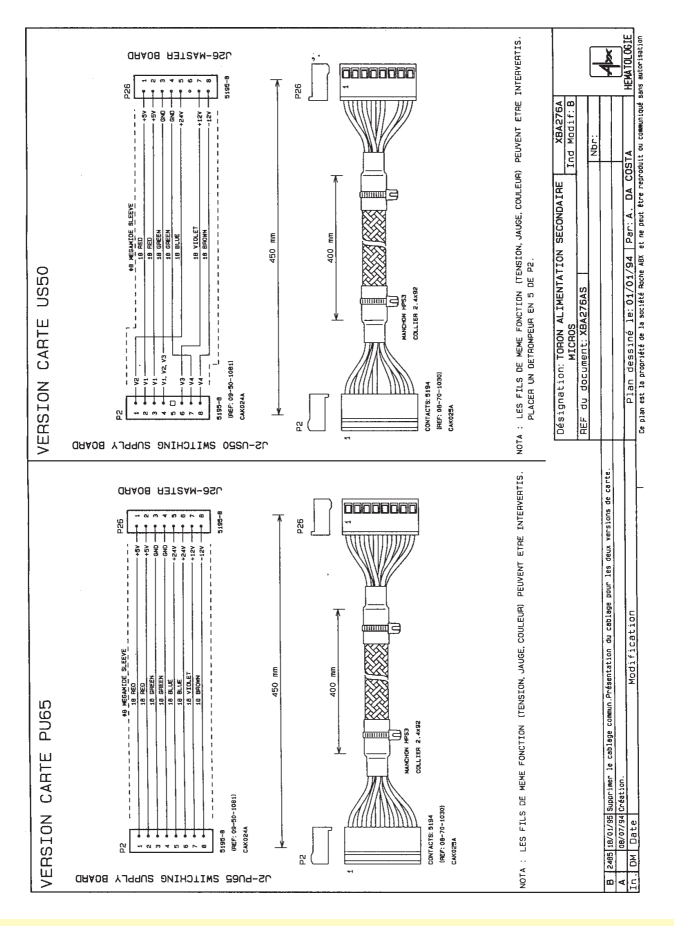


RAA 009 A Ind.A

1.2. Power supply fan

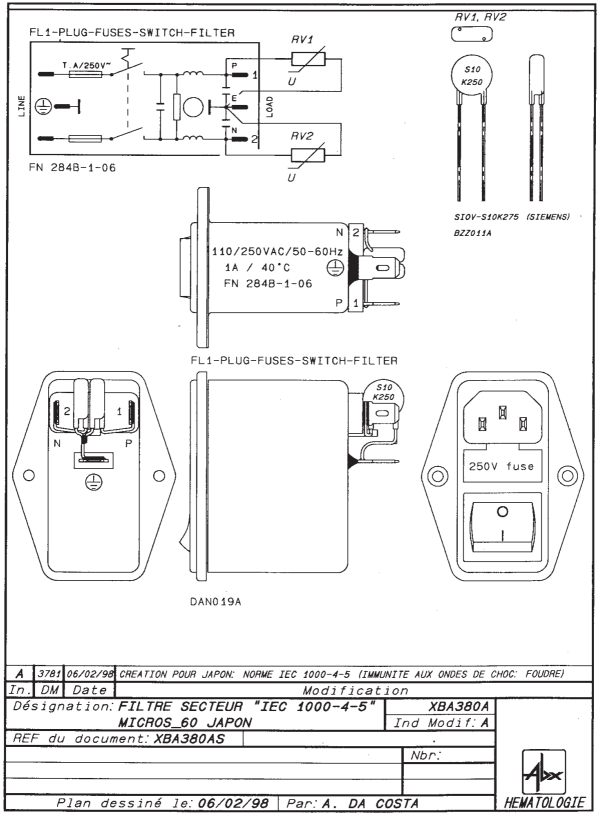


1.3. Secondary supply cable



RAA 009 A Ind.A

1.4. Main supply filter



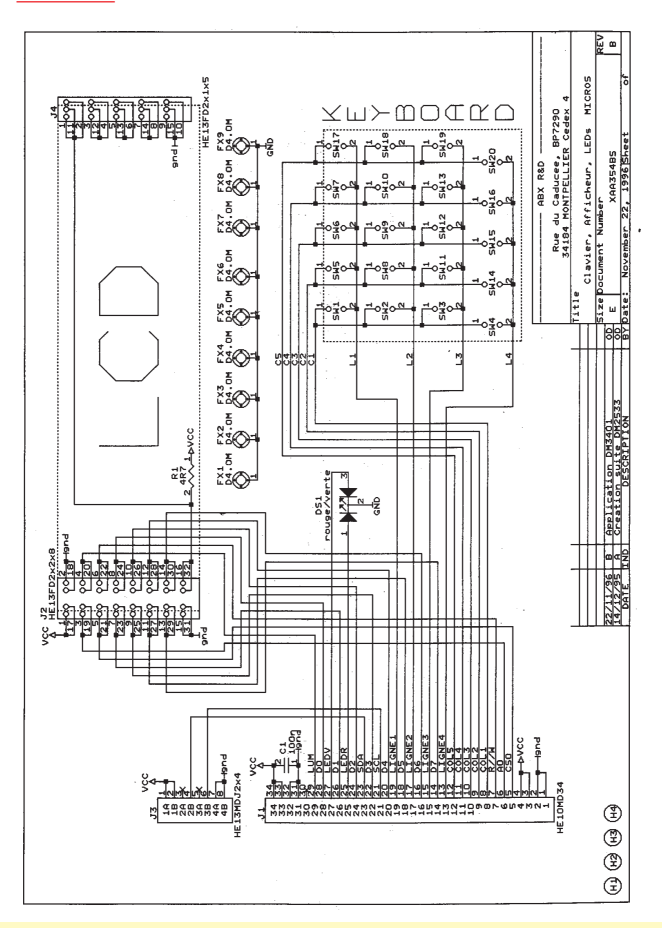
Ce plan est la propriété de la société ABX et ne peut être reproduit ou communiqué sans autorisation

2. MOTHER BOARD

2.1. Configuration

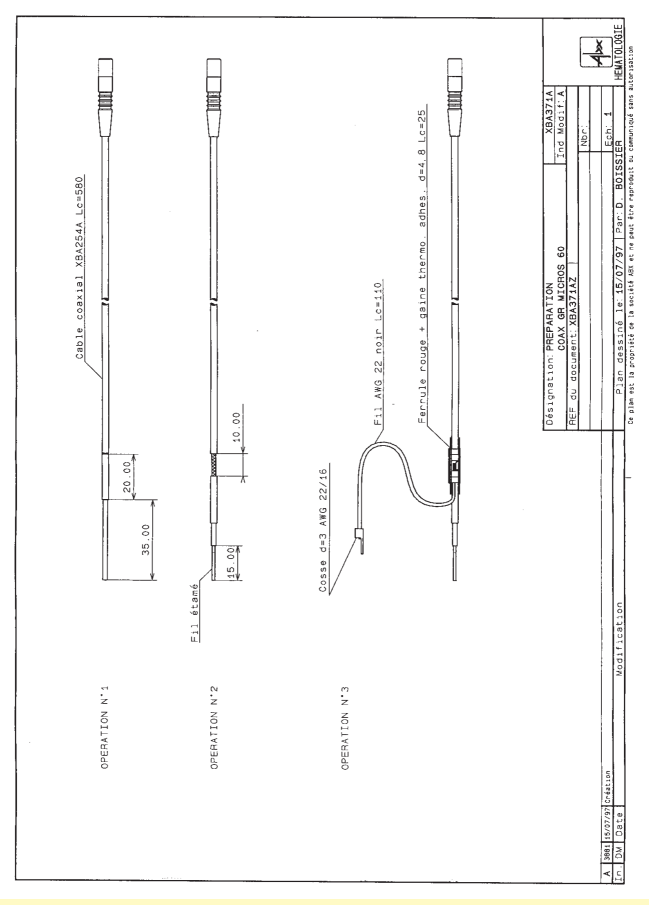
E1: EPROM	ELI EPROM address area option EPROM JUNPERS AREA 128Kx8 PCM NC	 		
i – –	1-3 don't francais	<u>E3: 60HCil MoDB signal option</u> on: MoDB = 0 (test mode, future use) Scals off: WODB = 1 (default value)		л Т
	.P.G.M. A.K7	 Edd, an: clock convected (default value) means off: clock no convected	FUNCTION	NCT
	248	E6: R5232 Contlawration		-]
	3-5 2-4 all tencals 3-5 2-4 all tencals 1-4 4-6 anglais 1-3 2-6 anglais	EZ2 on: off:	RTS/CTS are connected to DSR RTS/CTS are not connected to DSR (default value)	
			CONFIGURATION	ATION
		E3	 ,	
				`
			ABX Recherche & Developpement 128 Rue du Caducee, 34184 MONTPELLIER 1118 MI CROS MASTER Board 512 Document Number 512 Document Number 0 BY Date October 10, 1995 Sheet 1 of	i oppement 34184 NONTPELLIER ER Board ESBR SSBR SSBR

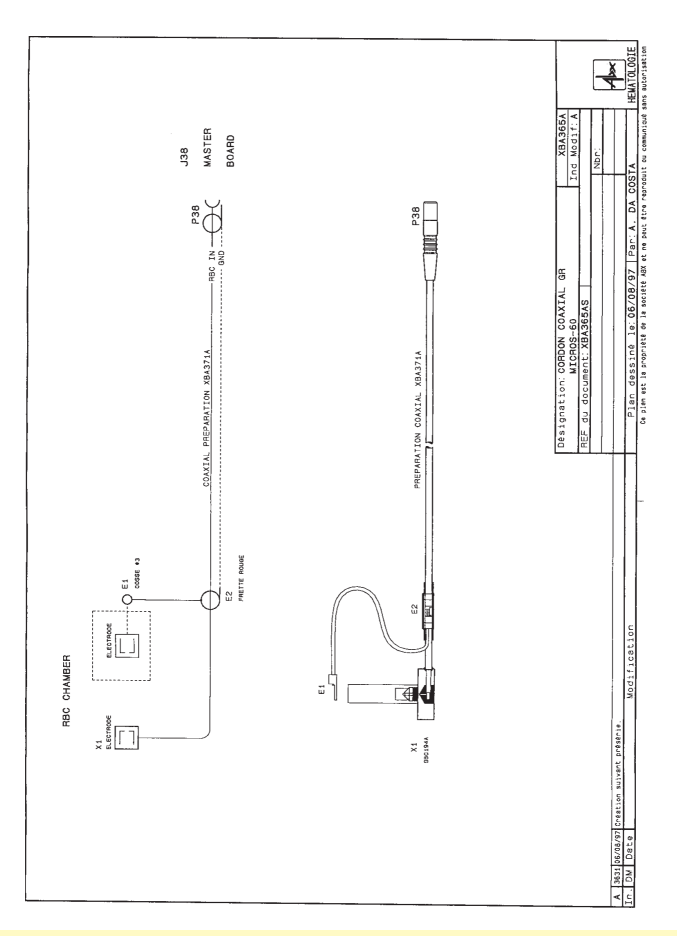
3. LCD BOARD



4. COAXIALS

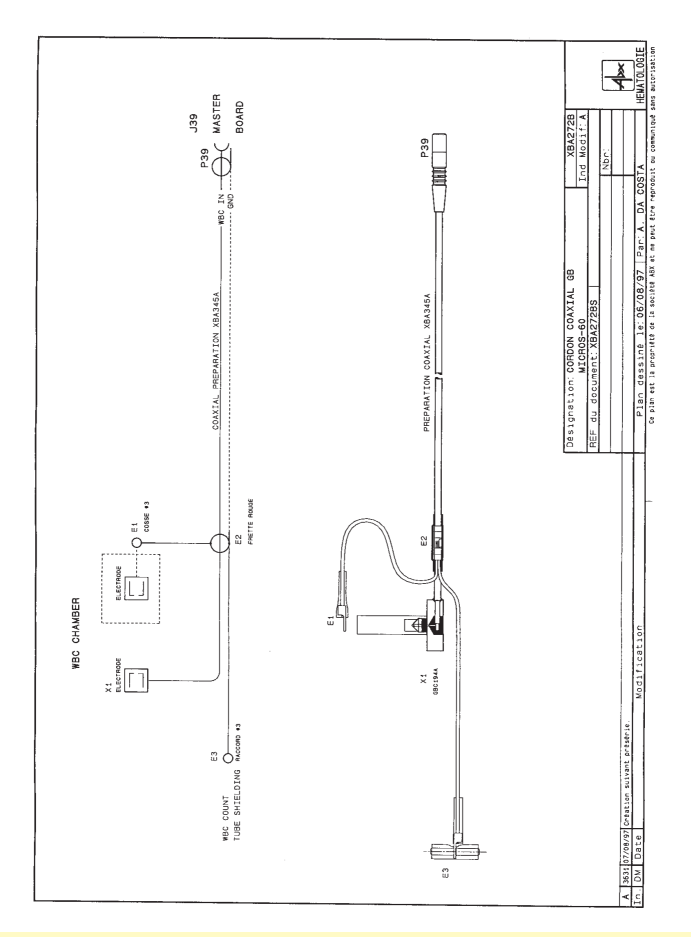
4.1. RBC coaxial





RAA 009 A Ind.A

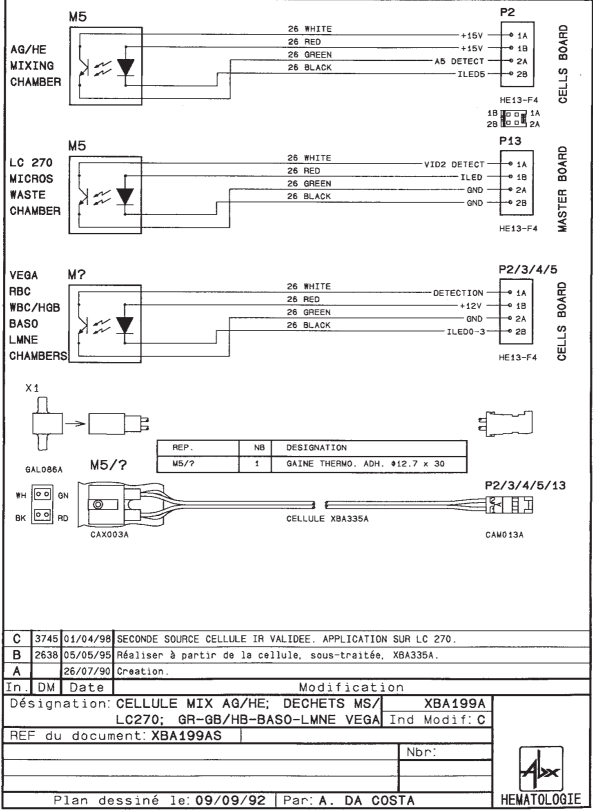
4.2. WBC coaxial



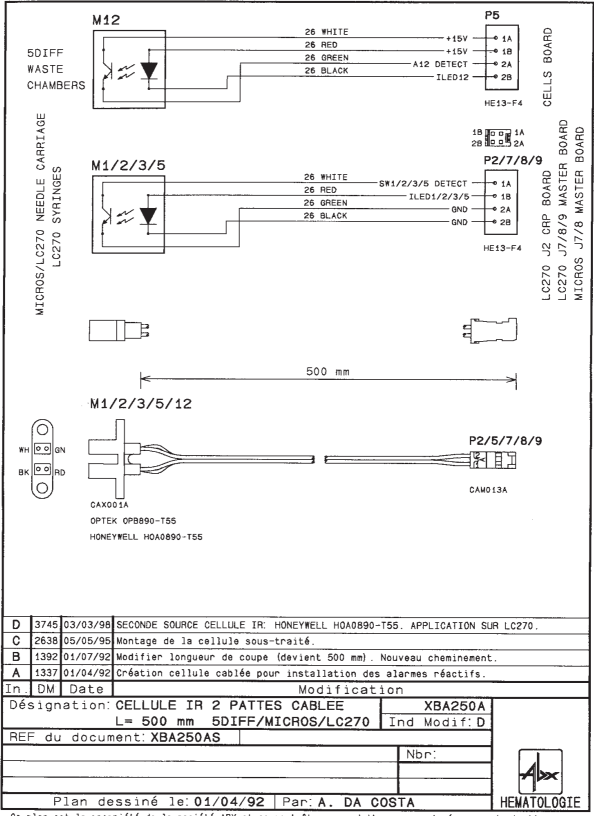
RAA 009 A Ind.A

5. SENSORS

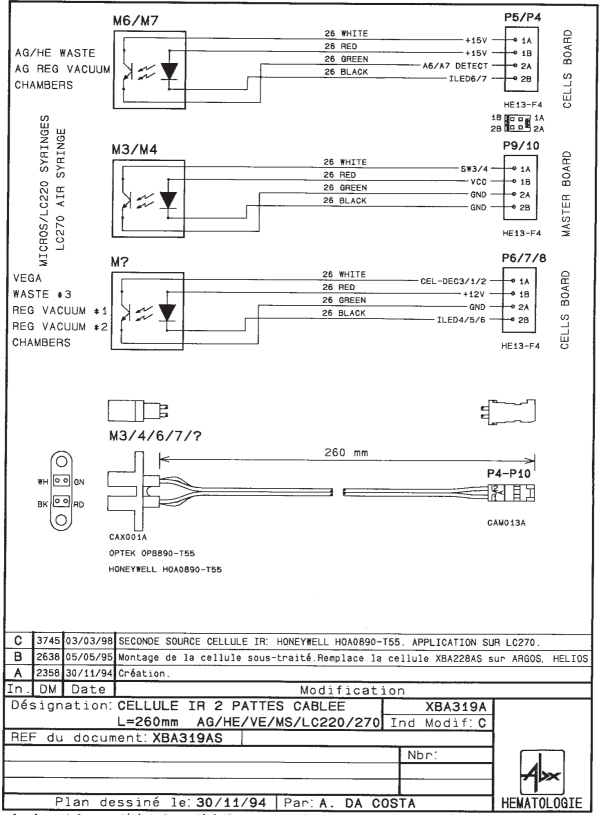
5.1. Drain detection sensor



5.2. Carriage & needle sensors

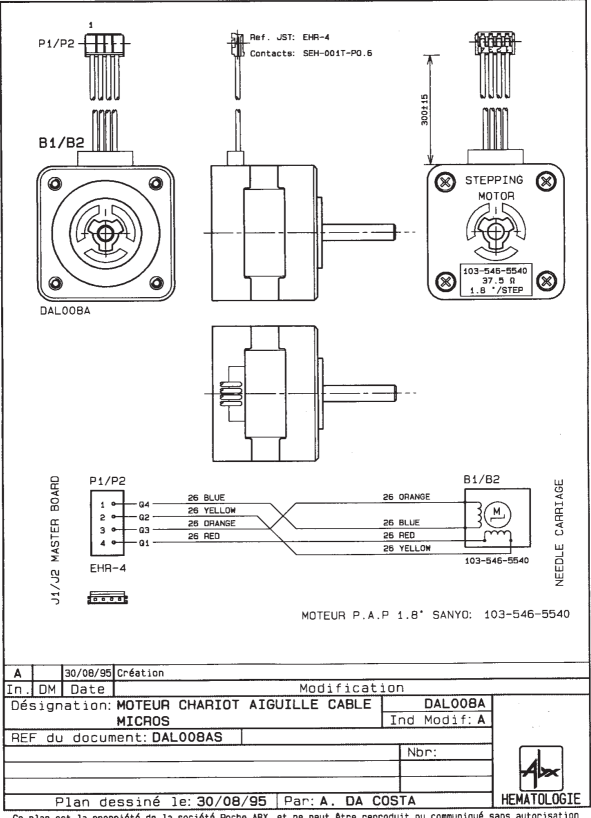


5.3. Syringe sensors

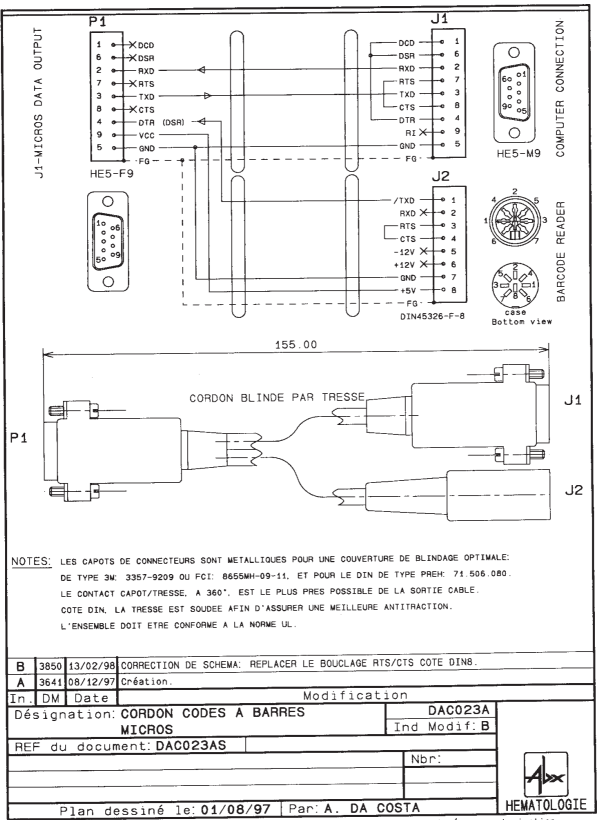


6. MISCELLANEOUS

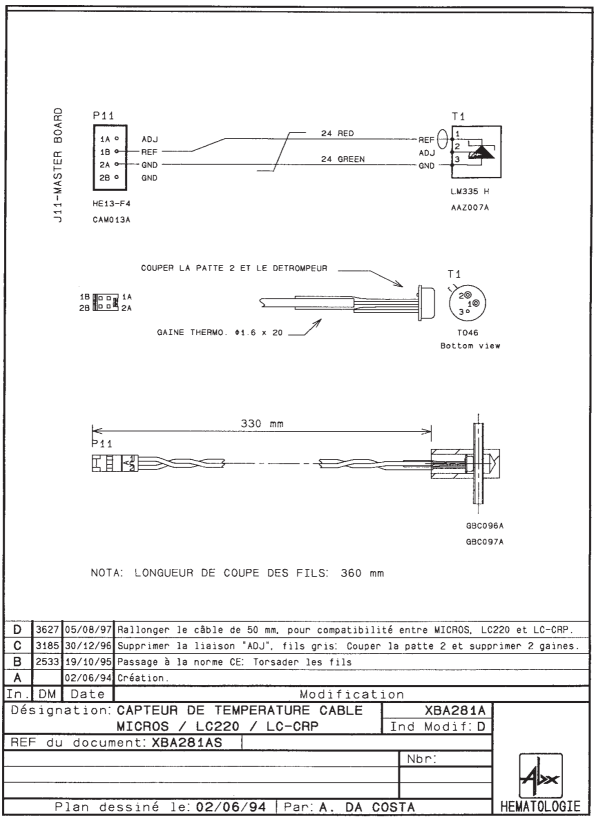
6.1. Needle carriage motor



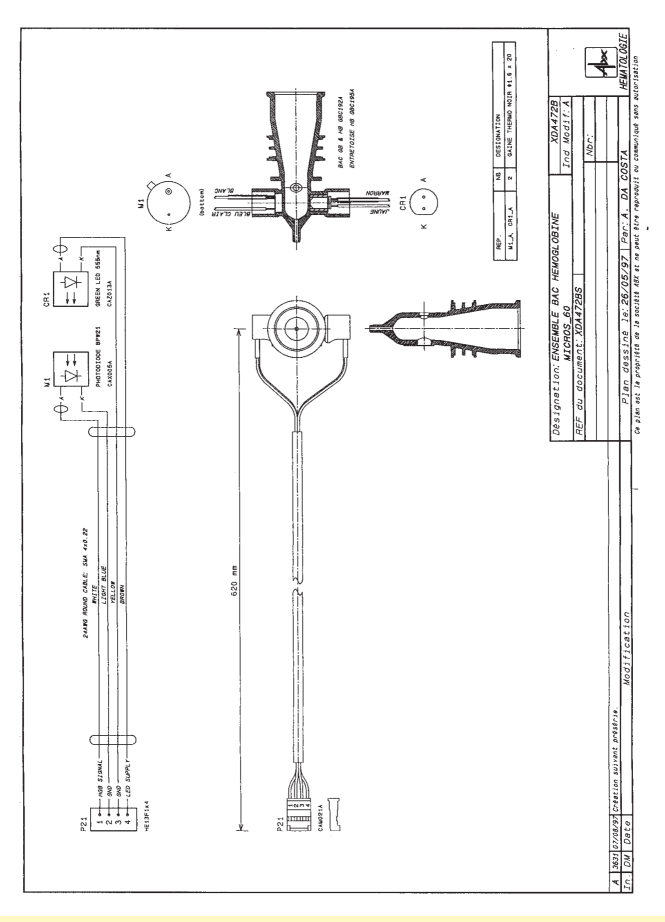
6.2. Barcode cable



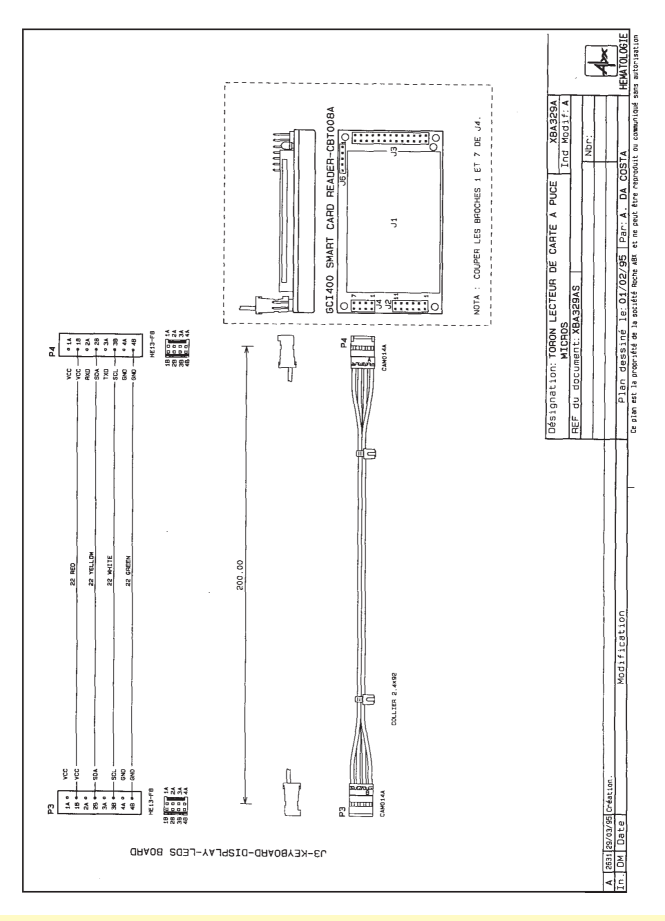
6.3. Temperature sensor



6.4. HGB Chamber assembly

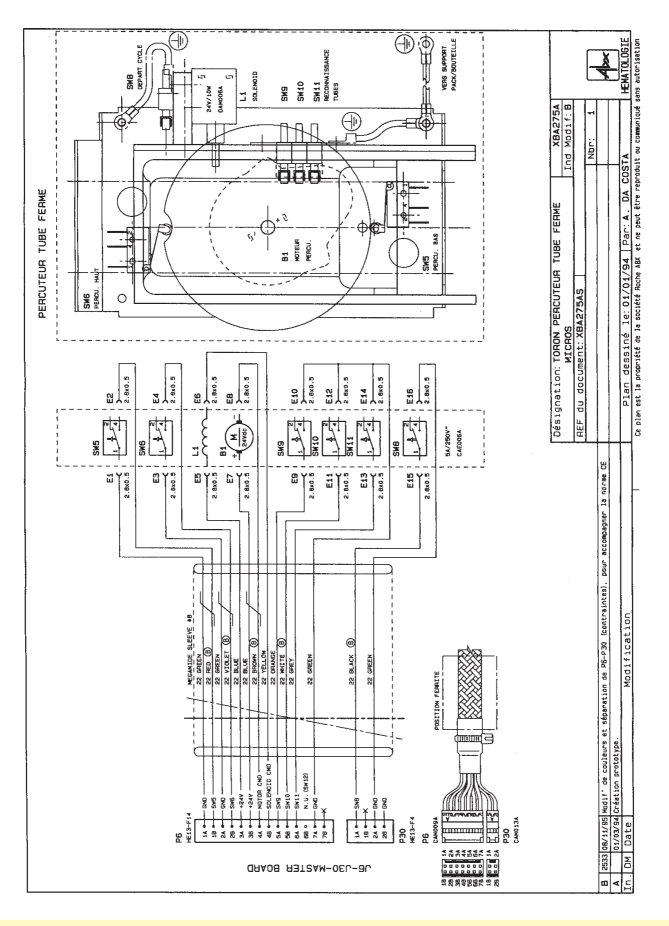


6.5. Chip card reader cable

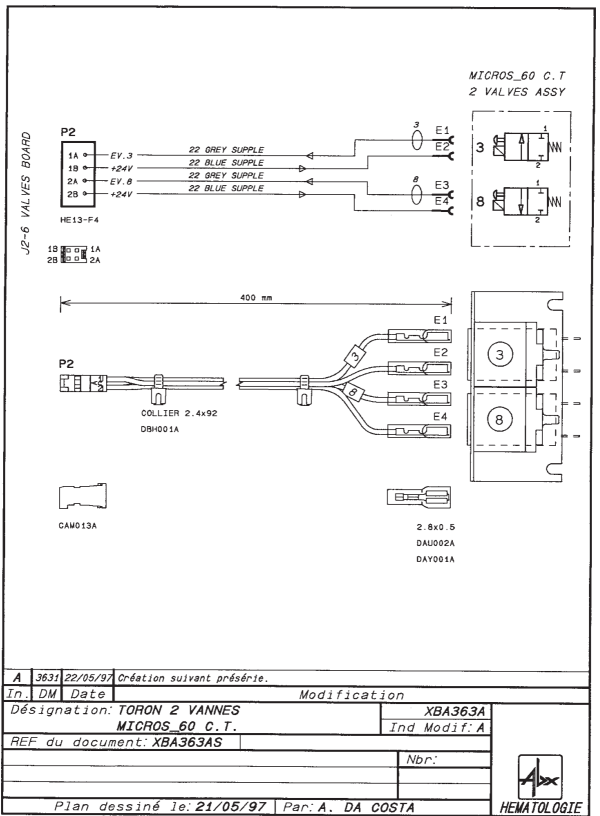


RAA 009 A Ind.A

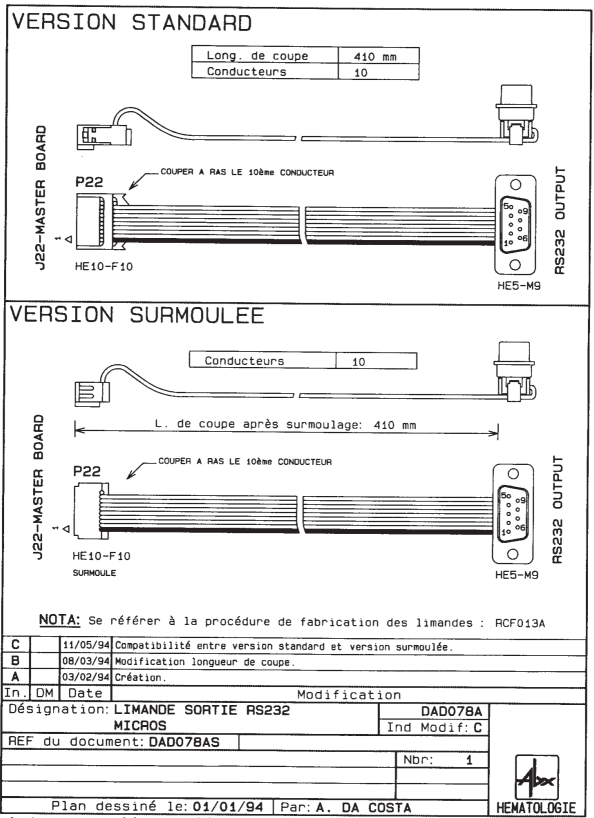
6.6. CT Piercing assembly cable

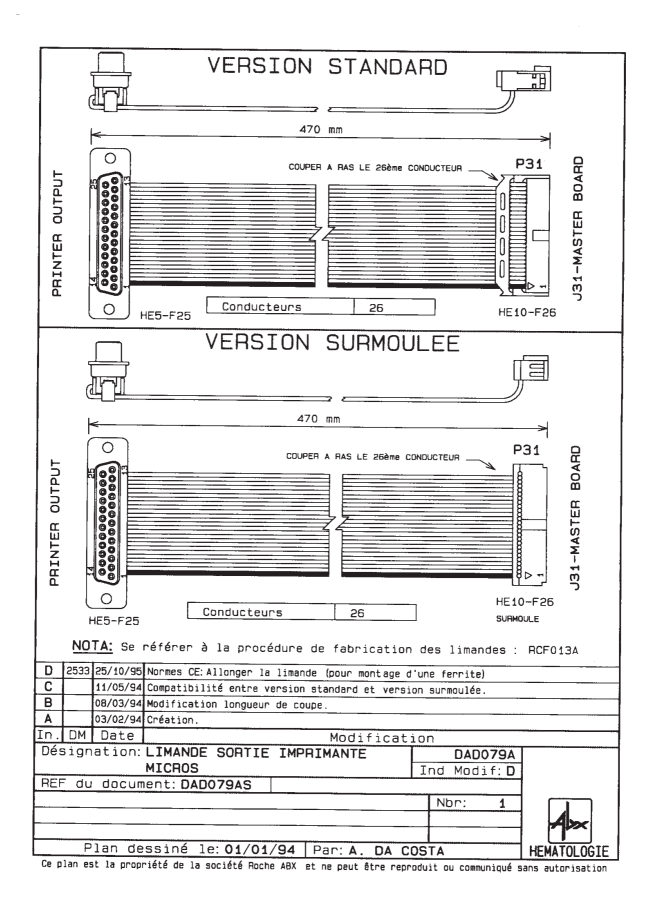


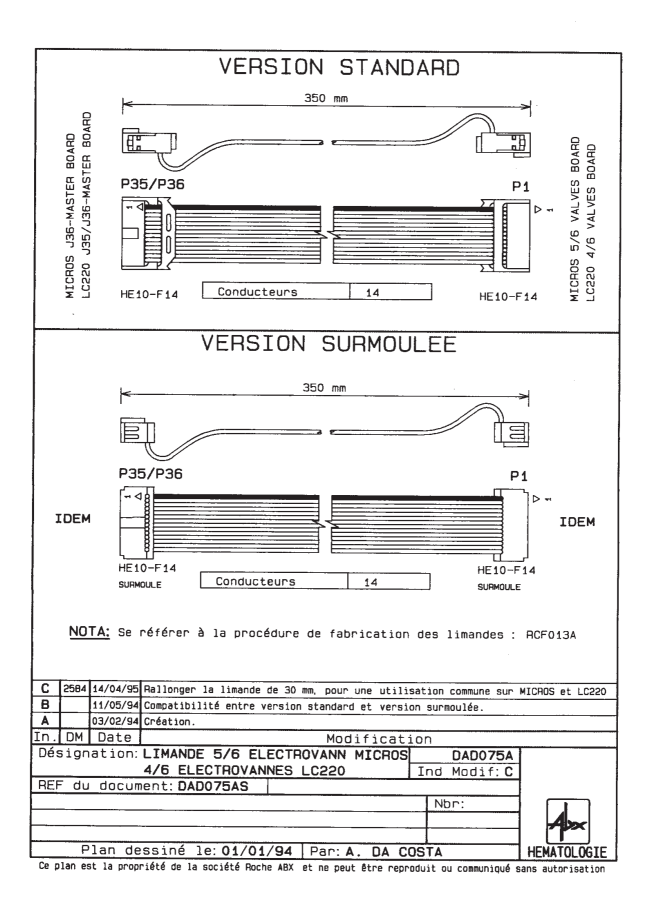
6.7. CT twin valve cable

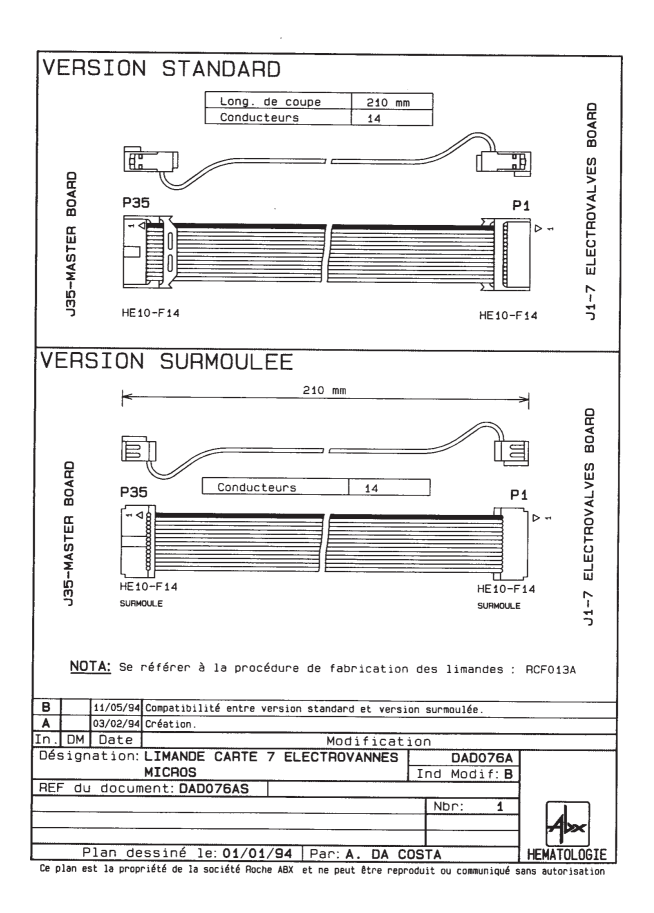


7. FLAT CABLES









8. ELECTRICAL SYNOPTICS

CONTENTS

1.	MAINTENANCE	. 2
	1.1. Introduction 1.2. Daily customer maintenance 1.3. Weekly customer maintenance	. 2
2.	Maintenance kits	
3.	PROCEDURES	4
	3.1. Procedure chart tables3.2. Required tools and products	5 7

1.1. Introduction



1.2. Daily customer maintenance

No special adjustments or maintenance has to be done on your equipment if the recommended startup and shutdown procedures are explicitly respected. See the **ABX MICROS 60** User Manual for the daily rinsing and cleaning of the system.

1.3. Weekly customer maintenance

An overall check for cleanliness of the system is recommended every week. All traces of blood or reagent have to be wiped off as soon as possible using a piece of cloth and distilled water.



Never use solvent or abrasive cleaning material to clean the system.

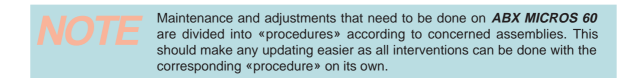
2. MAINTENANCE KITS

0	RING KIT : XEA 328 AS
FAA 036 A	1 Lyse dispenser O ring 1 Piercing needle O ring (MICROS CT only)
FAA 029 A	1 Diluent dispenser O ring
FAA 055 A	2 Sampling syringe O rings
FAA 049 A	2 Aperture O rings
FAA 046 A	2 Coaxial O rings
FAA 054 A	2 Sampling needle O rings (CT)
FAA 053 A	1 Sampling needle O ring (OT)
FAA 017 A	1 Vacuum/waste syringe O ring
KAM 022 A	6 Board holder clips (MICROS 45 only)
FAA 023 A	1 WBC/HGB chamber cap O ring
GBC 030 A	1 Air Syringe piston

Part number	SPARE PARTS KIT XEA 458 AS	Quantity
CAE 006 A	tube holder detection switch	1
EAE 005 A	Tygon tube 1.02mm (0,040")	2
EAE 007 A	Tygon tube 1,52mm (0,060")	2
EAE 008 A	Tygon tube 2,05mm (0,081")	2
EAE 009 A	Tygon tube 2,29mm (0,090")	2
FAK 001 A	Aperture 50µ	1
FAK 003 A	Aperture 80µ	1
FBR 002 A	Notched belt 290	1
FBR 003 A	Notched belt 380	1
GBC 052 A	C.T sampling needle	1
GBC 069 A	O.T sampling needle	1
GBC 189 A	CT Piercing needle	1
GBC 193 A	RBC chamber	1
XBA 199 A	Drain detection sensor	1
XBA 250 A	Carriage/needle sensor	1
XBA 272 B	WBC coaxial	1
XDA 472 B	WBC/HGB chamber	1
XDA 481 B	Liquid valve 2W NC without solenoid	1
XDA 483 B	Liquid valve 3W without solenoid	1
XBA 365 A	RBC coaxial	1
XBA 319 A	Liquid/air syringe sensor	1
XEA 328 AS	Maintenance kit	1

RAA 009 A Ind.A

3. PROCEDURES



CONCERNING PARTS :

- Hydraulic maintenance and adjustments.
- Pneumatic maintenance and adjustments.
- Electrical maintenance and adjustments.
- Power supply maintenance and adjustments.
- Electronic maintenance and adjustments.
- Printer maintenance and setup.

Each procedure has to be read entirely before beginning the intervention.



When cleaning instruments, disposable gloves should be worn.

3.1. Procedure chart tables

P / N	MICRO PROCEDURE	S 6 0 O T concerns
RAS 165 A	MICROS 60 OT INSTALLATION AND STARTUP	Unpacking - Working conditions - Visual checks - Reagent connection - Printer & instrument connections - Priming & Startup
RAS 168 A	SAMPLING NEEDLE MAINTENANCE MICROS OT	Needle replacement - O ring replacement
RAS 169 A	CHAMBER MAINTENANCE	RBC, WBC/HGB chamber cleaning - Aperture O ring replacement - Coaxial O ring replacement
RAS 170 A	LIQUID VALVE MAINTENANCE	Liquid valve assy replacement - Valve body replacement
RAS 171 A	POWER SUPPLY CHECK/REPLACEMENT	Voltage supply check - Power supply module replacement Fan operation check.
RAS 172 A	TECHNICIAN FUNCTIONS MICROS 60 OT	Version display Adjustments : HGB photometer calibration, Aperture voltage, pressure check, WBC gain , RBC & PLT gain, Sensor, Needle heigth and motion, bubbling Temperature sensor adjustment - Run mode - Reagent pack - Serial number - Cycle number - Burning
RAS 173 A	MECHANIC FUNCTIONS	Sensor replacement - Needle motion check - Carriage motion check - Liquid syringe motion check - Vac/Waste syringe motion check - Valve operation check - LCD contrast - Piercing mechanism check
RAS 174 A	DRAIN DETECTION ADJUSTMENT	Drain detection sensor adjustment
RAS 175 A	PCB VOLTAGE CHECKS	Voltage supply check/adj Aperture voltage check/adj RBC threshold check/adj WBC threshold check/adj PLT threshold check/adj HGB blank voltage check - Stepper motor voltage adjustment
RAS 177 A	LX300 PRINTER	Configuration - Control panel - Control LEds and keys - Description
RAS 178 A	LIQUID SYRINGE MAINTENANCE	Lyse dispenser O ring replacement - Diluent dispenser O ring replacement - Sampling needle dispenser O ring replact Lubrification
RAS 179 A	VACUUM/WASTE SYRINGE MAINTENANCE	O ring replacement
RAS 180 A	CHANGING THE INSTRUMENT LANGUAGE	Changing the instrument language
RAS 181 A	REAGENT PACK	Connector O ring replacement
RAS 182 A	BARCODE READER SETUP	Reader configuration
RAS 187 A	HYDRAULIC CYCLE CHECKUP MICROS 60 OT	Step by step control of the hydraulic cycle
RAS 191 A	OVERALL MAINTENANCE	Instrument maintenance step by step
RAS 192 A	DECONTAMINATION	Instrument decontamination
RAS 197 A	DRAIN & RINSE	Instrument rinse and drain for an extended shutdown

	MICRO	S 6 0 C T
P / N	PROCEDURE	C O N C E R N S
RAS 166 A	MICROS 60 CT INSTALLATION AND STARTUP	Unpacking - Working conditions - Visual checks - Reagent connection - Printer & instrument connections - Priming & Startup
RAS 167 A	SAMPLING NEEDLE MAINTENANCE MICROS CT	Needle O ring replacement - Sampling needle replacement Piercing needle replacement
RAS 169 A	CHAMBER MAINTENANCE	RBC, WBC/HGB chamber cleaning - Aperture O ring replacement - Coaxial O ring replacement
RAS 170 A	LIQUID VALVE MAINTENANCE	Liquid valve assy replacement - Valve body replacement
RAS 171 A	POWER SUPPLY CHECK/REPLACEMENT	Voltage supply check - Power supply module replacement Fan operation check.
RAS 176 A	TECHNICIAN FUNCTIONS MICROS 60 CT	Version display Adjustments : HGB photometer calibration, Aperture voltage, pressure check, WBC gain , RBC & PLT gain, Sensor, Needle heigth and motion, bubbling Temperature sensor adjustment - Run mode - Reagent pack - Serial number - Cycle number - Burning
RAS 173 A	MECHANIC FUNCTIONS	Sensor replacement - Needle motion check - Carriage motion check - Liquid syringe motion check - Vac/Waste syringe motion check - Valve operation check - LCD contrast - Piercing mechanism check
RAS 174 A	DRAIN DETECTION ADJUSTMENT	Drain detection sensor adjustment
RAS 175 A	PCB VOLTAGE CHECKS	Voltage supply check/adj Aperture voltage check/adj RBC threshold check/adj WBC threshold check/adj PLT threshold check/adj HGB blank voltage check - Stepper motor voltage adjustment
RAS 177 A	LX300 PRINTER	Configuration - Control panel - Control LEds and keys - Description
RAS 178 A	LIQUID SYRINGE MAINTENANCE	Lyse dispenser O ring replacement - Diluent dispenser O ring replacement - Sampling needle dispenser O ring replact Lubrification
RAS 179 A	VACUUM/WASTE SYRINGE MAINTENANCE	O ring replacement
RAS 180 A	CHANGING THE INSTRUMENT LANGUAGE	Changing the instrument language
RAS 181 A	REAGENT PACK	Connector O ring replacement
RAS 182 A	BARCODE READER SETUP	Reader configuration
RAS 188 A	HYDRAULIC CYCLE CHECKUP MICROS 60 CT	Step by step control of the hydraulic cycle
RAS 191 A	OVERALL MAINTENANCE	Instrument maintenance step by step
RAS 192 A	DECONTAMINATION	Instrument decontamination
RAS 197 A	DRAIN & RINSE	Instrument rinse and drain for an extended shutdown
RAS 198 A	PIERCING BLOCK	Description - Maintenance - Sampling position

3.2. Required tools and products

то	D L S	PROD	UCTS
DESIGNATION	PART NUMBER	DESIGNATION	PART NUMBER
HEXAGONAL KEYS		EMPTY SAMPLE TUBES	
DYNAMOMETRIC SCREW DRIVER A302	MAG 019 A	SILICONE GREASE	LAM 004 A
DYNAMOMETRIC SCREW DRIVER A301	MAG 020 A	GREASE FOR MECHANICAL ASSEMBLIES	XEA 381 AS
DYNAMOMETRIC SCREW DRIVER A300	MAG 013 A	SOFT TISSUE	
CLAMPS		LIQUID SOAP	
SCALPEL		DISTILLED WATER	
CUTTING PLIERS		MICROPIPETTE TIP	
PAIR OF SCISORS		FLAT PIECE OF STIFF PLASTIC	
VOLTMETER		LATEX WBC	LAD 001 AS
FLAT SCREWDRIVER		LATEX RBC	LAD 002 AS
BARFLEX		FELT PEN	
THERMOMETER		SYRINGE 5ML	
TORX KEYS			

INSTALLATION

• CONCERNS

- 1 Unpacking
- 2 Working conditions
- 3 Visual checks
- 4 Reagent connections
- 5 Printer and instrument connections
- 6 Priming and startup

· REQUIRED TOOLS

Hexagonal keys

· REQUIRED PRODUCTS

- MICROS 60 Reagents : Bottles or Pack.
- Waste container (for bottle model).

INTERVENTION TIME

- 30 minutes

• FREQUENCY

• SPECIFIC KIT OR CONSUMABLES

- PACK installation kit : XEA 314 A
- or
- Bottle installation kit : XEA 332 A



RAS 165 A Ind.B

INSTALLATION

PROCEDURE

1 - Unpacking



The instrument is enveloped in a special, protective foam before being placed in a cardboard box. Cut the four angles of the box to unpack the system.

Remove the cardboard box containing the **ABX MICROS 60** installation kit from its location (see Diag.1).

Diag.1

2 - Working conditions

Environment

ABX MICROS 60-*OT* should be operated in an indoor location only. Operation at an altitude over 2000 meters is not recommended. Instrument is designed to be safe for transient voltages according to INSTALLATION CATEGORY II and POLLUTION DEGREE 2.

Location

ABX MICROS 60-*OT* should be placed on a clean and level table or work station. Please note that **ABX MICROS 60-***OT*, printer and reagents weigh approximately 30 kilograms (66 lbs). Avoid exposure to sunlight. Proper ventilation requires that a space of at least 20 cm (8 inches) must be left behind the apparatus.

• Grounding

Proper grounding is required. Check that the wall ground (earth) plug is correctly connected to the laboratory grounding electricity installation. If there is no ground then use a ground stake. Current electricity norms must be applied.

• Humidity and temperature conditions

ABX MICROS 60-*OT* can function between 18 to 32° C (65 to 90° F), with relative humidity, meaning less than 80% with no condensation. If it is kept at a temperature less than 10° C (50° F), the instrument should be allowed to sit for an hour at the correct room temperature before use.

INSTALLATION

3 - Visual checks

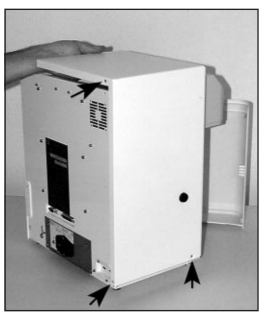


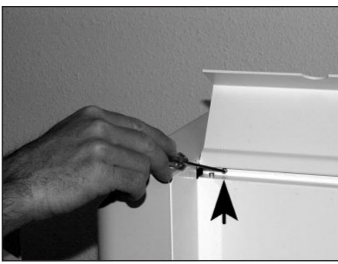
Diag.2

Using the key contained into the installation kit, turn the locker as shown on the Diag. 2 to open the pneumatic protection door.

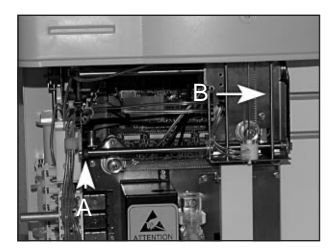
Unscrew the 3 cover fixation screws (Diag. 3) and loosen the 2 tightening screws under the reagent flap (Diag 4).

Remove the cover : pull it backward and lift it up to the rear of the instrument .





Diag.4



Diag.5

Diag.3

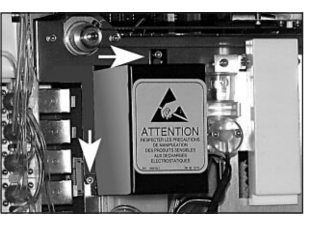
Push the black plastic carriage locking clip (A) as far as possible to the left and place the sample needle carriage (B) as far forward as possible to the right-hand side, as shown in Diag. 5.

Check that the needle is not bent and make sure it is in its upper position.



INSTALLATION

Unscrew slightly the 2 screws of the WBC/HGB chamber protection cover (diag 6). Remove the cover and check that both chambers (RBC/PLT, WBC/HGB) are fixed properly in their clips and the electrode blocks are attached firmly to the chambers (Diag. 7).

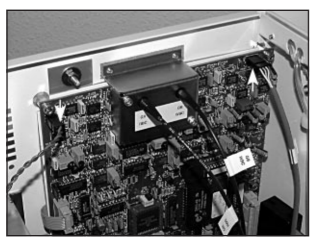


Diag.6



Diag.7

Re-install the HGB/WBC chamber cover.



Check that the connectors on the printed circuit board are securely in place (Diag. 8).

Re-install the instrument cover.

Diag.8



Diag.9

Remove the fuse holder from its location on the rear panel pressing on the holder lock (Diag.9) and check the fuse characteristics : they should be 1 Ampere, 250 Volts Slow-Blow.



4 - Reagent connections

• Bottle connections



Lyse and cleaning reagents are placed inside the reagent compartment as shown in the Diag. 10. Install the reagent straws and the bottle stoppers. Connect the blue tube to the MINICLEAN bottle and the white tube to the MINILYSE bottle. Close the compartment cover.

Install the male connectors included in the installation kit at the liquid input and output located at the bottom of the instru-

Connect the diluent container (see CAU-

TION above) using the diluent straw and

a 3x6 cristal tube (1 meter maximum) on the diluent input located at the bottom of the instrument rear panel (Diag.11). Connect the waste container using the cristal tube 3 x 6 on the waste output, and place the waste container below the instrument level (under the bench).

ment rear panel (Diag.11).

Diag.10

IMPORTANT

When the ABX MICROS 60-*oT* is set up with the 16 or 18 parameters mode, it is mandatory to use specific MINILYSE LMG and MINIDIL LMG reagents.

CAUTION

The Diluent container will be located on the bench at the same level than the instrument.

Waste connection



Diag.11





Always follow the recommended procedures for waste disposal. Never connect the instrument wastes directly to the laboratory drain pipes. For each waste container, follow the neutralization procedure as described in the user manal.



INSTALLATION

• Reagent pack connection

Remove the reagent output protections, as well as the waste input protection (Diag.12 & 13)









Install the pack directly into the compartment of the instrument as shown on the Diag. 14, 15 &

16. Push the pack down in order to plug correctly the pack on the male connectors.



Diag.14



The free male connector (see Diags. 14, 15 & 16) must be plugged on the pack upper valve in order to receive the waste liquids.

Diag.16

CAUTION

In order to avoid leak problems it is recommended not to unplug several times the same reagent pack.



Diag.15

Page 6/13

INSTALLATION

5 - Printer and instrument connections

• Instrument connection.

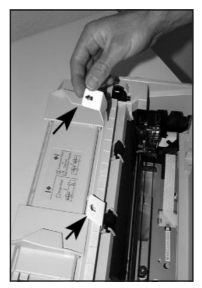
If the instrument has to be connected to a laboratory computer, use the plug RS232.

Connect the power cable to the plug located on the rear left-hand side of the device (Diag. 17).



Diag.17

- Setting up the printer
- Remove all the package protections
- Install the paper feed-knob



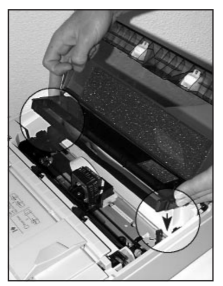


Diag.19

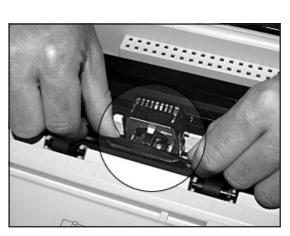
Diag.18

Open the ink ribbon access door at the top of the printer and install the ribbon as shown in diag 20 & 21 : Slide the printer head to the middle of the printer.

Insert the ribbon cartridge into the printer Guide the ribbon between the print head and ribbon guide. Slide the printer head from side to side to make sure it moves smoothly.



Diag.20



Diag.21

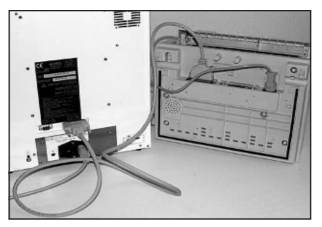
INSTALLATION

- Install the paper supports (Diag 22) for single sheets paper use only.



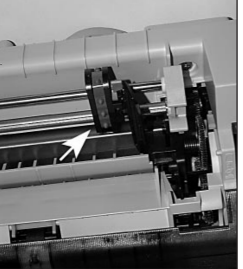


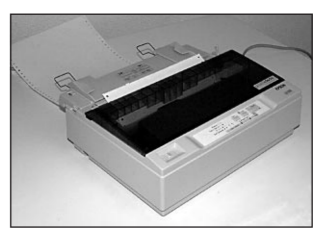
- The printer is connected to **ABX MICROS 60**-*OT* with the cable delivered with the instrument. Lock the connector in place by tightening the 2 screws on each end of the connector to the **ABX MICROS 60**-*OT*. Attach the other end of the cable to the printer and lock the printer connector in place by the means of the 2 clips located on the connector itself (Diag. 23).





- For continuous paper, introduce it in the slot at the back of the printer and use the sprocket covers to load paper, or else feed the paper frontward when using single sheets (see printer user manual).





Diag.25

Diag.24





• Printer command keys

Roman				Font	LF/FF	BING 1	Pause
Sans Serif	-0	•				201	Contraction of the second
Draft					S.S. manual S	and the second	
Draft Condensed	• •	• 1	2	L []Pa	per Park	Ռո	CONCERCION OF
	12/13	Micro	o Adjust	1	1	3sec	

Diag.26

LED PAUSE : The orange LED PAUSE lights when the printer stops printing. During each power ON, this LED blinks for few seconds and 4 audible beeps occur. When the printer runs out of paper, the LED blinks and 3 audible beeps occur. This LED lights also when the paper is in its tear off position. When a problem occurs, this LED lights ON and 5 audible beeps occur.

LEDS FONT 1 and FONT 2 : These 2 green LEDS indicate the selected font. Refer to the printer user's manual to select the font.

Key FONT : During normal operation, the FONT key allows the font selection. For each pressure on this key, the selection is modified. Refer to the printer user's manual to select the font. When this key is pressed during the printer power ON, the printer setup menu is entered.

Key LF/FF : During normal operation, a quick pressure on this key allows a ligne feed of the paper. Keep the pressure on this key to feed a whole page. This key can be used to load or eject the paper.

When this key is pressed during the printer power ON, the printing test starts. Key PAUSE : When this key is pressed during the printing, the printout stops. Press again on this key to restart the printout.

PAPER PARK : If Z folded paper is used, the paper can be driven to its parking position when pressing simultaneously on the keys LF/FF and FONT.

MICRO ADJUST : This function allows to adjust the loading paper position. See the user's manual for details.

• Printer Configuration

Switch on the printer when pressing the . The configuration should be the following :

0000000000	**************************************	<pre>(************************************</pre>	0000000000
$\overline{}$			- U

Diag.29

INSTALLATION

Each parameter can be modified by the corresponding parameter chart. Each chart is accessible using the keys <PAUSE>, and <LF/FF> according to the control LED combinations (Diag 30).

	Links		Table B	
1	Lights 2	PAUSE	Setting	Go to submenu
BLINKS	\$12, 123 all, 105 all the test feet that the	OFF	Character spacing	Table C
BLINKS	ON	OFF	Shape of zero	Table D
OFF	BLINKS	OFF	Skip-over-perforation	Table E
ON	BLINKS	OFF	Character table	Table F
BLINKS	BLINKS	OFF	Auto line feed	Table G
BLINKS	OFF	ON 1	Page length	Table H
BLINKS	; ON	ON :	Auto tear off	Table I
OFF	BLINKS	I DN I	Tractor	Table J
ON	: BLINKS	ON :	Interface	Table K
BLINKS	BLINKS	I ON I	Bit rate	Table L
OFF	OFF	BLINKS	Parity	Table M
BLINKS	: OFF	BLINKS	Data length	Table N
ON	OFF	BLINKS	ETX/ACK	Table D
ON	; ON	BLINKS	Software	: Table P
BLINKS	ON ·	BLINKS	Auto CR	Table Q

Diag.30

6 - Instrument startup

Reagent priming

When the **ABX MICROS 60-***ot* is first installed, it contains no reagents. All the reagents have to be primed now. Turn ON instrument by pressing the ON/OFF switch located on the rear panel. When the instrument turns on, the display shows :

PLEASE WAIT FOR 3 MIN ESCAPE : ESC

This time is required at the startup for the instrument initialization and stabilization, specifically for the HGB diode to reach its operationnal temperature. Press ESC several times in order to abort the cycle : the LED of the front panel turns from red to green and the display shows the following :

STARTUP NOT INITIATED PRESS A KEY TO CONTINUE...

This message appears when the instrument is setup with the manual startup cycle to prevent any analysis cycle before running a startup cycle. Press any key, the main menu is displayed :

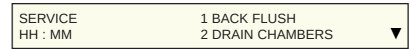
MAIN MENU	1 RESULTS	
HH : MM	2 QC	7



• Bottles and containers set :

Install the reagent bottle and carry out a PRIME cycle to clear the reagent line of air bubbles. This procedure should be done whenever a new bottle of reagent is installed.

From the MAIN MENU, move the cursor to the function $\begin{pmatrix} 4 \\ \end{pmatrix}$ SERVICE and press ENTER. The service menu is displayed :



Move the cursor to function 3 PRIME REAGENTS and press ENTER. The PRIME menu is displayed :

PRIME	1 ALL REAGENTS	
HH : MM	2 DILUENT	▼

Select either the function $\begin{pmatrix} 1 \\ \end{pmatrix}$ prime ALL REAGENTS or move the cursor next to the required reagent and press ENTER.

The priming cycle starts while the following menu is displayed :

ALL REAGENTS	WAIT FOR 2 MN 3 S

IMPORTANT

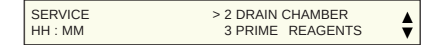
Before analyzing samples, visually inspect reagent lines and pumps for air bubbles. Repeat priming if air bubbles are still present. Call the *ABX representative service department* if priming does not eliminate air bubbles.

Never initiate two Lyse prime cycles back-to-back. This causes excessive foaming in the waste chamber. Run a blank cycle between each Lyse prime cycle.

- Run a STARTUP cycle.

Reagent pack

From the MAIN MENU, move the cursor to the function $\begin{pmatrix} 4 \\ \end{pmatrix}$ SERVICE and press ENTER. The service menu is displayed :



Move the cursor to $\begin{pmatrix} 3 \\ \end{pmatrix}$ PRIME REAGENTS and press the ENTER key. Select the function

CHANGE PACK and follow the instructions given by the LCD in order to install the pack.

REAGENT PACK	> 1 CHANGE PACK	
HH : MM	2 CBC LEFT < 150>	▼

Once the new PACK is installed a priming cycle will be automatically carried out and the following menu is displayed.

PRIME

WAIT FOR 2 MIN 3 S

MPORTANT

Before analyzing samples, visually inspect reagent lines and pumps for air bubbles. Repeat priming if air bubbles are still present. Call the *ABX representative service department* if priming does not eliminate air bubbles.

From the REAGENT PACK menu, the function $\binom{2}{}$ "CBC LEFT" displays the number of analysis cycles left to run with the same pack.

It is also possible to run a priming cycle at any time using the selection $\binom{3}{}$ "PRIME REAGENTS" of the SERVICE menu.

IMPORTANT

It is recommended not to remove the pack several times before the reagents are totally used in order to avoid leak problems.

- Run a STARTUP cycle.

INSTALLATION

Once the instrument ready for the analyses, remove the adhesive protection from the front panel (diag 31)



Diag.31

• CONCERNS

- 1 Unpacking
- 2 Working conditions
- 3 Visual checks
- 4 Reagent connections
- 5 Printer and instrument connections
- 6 Priming and startup

· REQUIRED TOOLS

Hexagonal keys

· REQUIRED PRODUCTS

Reagents : Bottles or Pack. Waste container (for bottle model).

INTERVENTION TIME

30 minutes

• FREQUENCY

· SPECIFIC KIT OR CONSUMABLES

PACK installation kit : XEA 317 A or Bottle installation kit : XEA 335 A or



RAS 166 A Ind.B

PROCEDURE

1 - Unpacking



The instrument is enveloped in a special, protective foam before being placed in a cardboard box. Cut the four angles of the box to unpack the system.

Remove the cardboard box containing the instrument installation kit from its location (see Diag.1).

Diag.1

2 - Working conditions

Environment

ABX MICROS 60-*CT* should be operated in an indoor location only. Operation at an altitude over 2000 meters is not recommended. Instrument is designed to be safe for transient voltages according to INSTALLATION CATEGORY II and POLLUTION DEGREE 2.

Location

ABX MICROS 60-*CT* should be placed on a clean and level table or work station. Please note that **ABX MICROS 60-***CT*, printer and reagents weigh approximately 30 kilograms (66 lbs). Avoid exposure to sunlight. Proper ventilation requires that a space of at least 20 cm (8 inches) must be left behind the apparatus.

Grounding

Proper grounding is required. Check that the wall ground (earth) plug is correctly connected to the laboratory grounding electricity installation. If there is no ground then use a ground stake. Current electricity norms must be applied.

• Humidity and temperature conditions

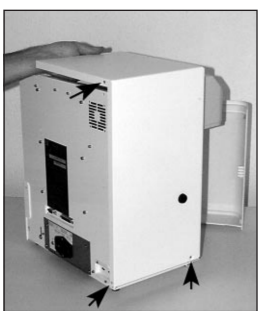
ABX MICROS 60-*CT* can function between 18 to 32° C (65 to 90° F), with relative humidity, meaning less than 80% with no condensation. If it is kept at a temperature less than 10° C (50° F), the instrument should be allowed to sit for an hour at the correct room temperature before use.

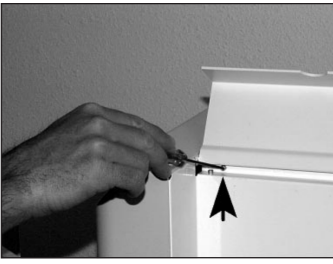
INSTALLATION

3 - Visual checks



Diag.2





Using the key contained into the installation kit, turn the locker as shown on the Diag. 2 to open the

Unscrew the 3 cover fixation screws (Diag. 3) and loosen the 2 tightening screws under the reagent

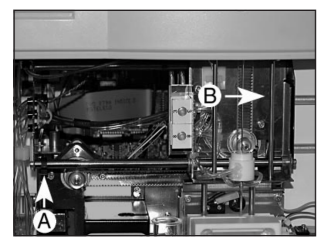
Remove the cover : pull it backward and lift it up to

pneumatic protection door.

the rear of the instrument.

flap (Diag 4).

Diag.4



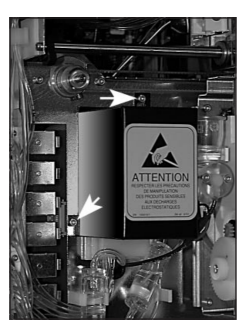
Diag.5

Diag.3

Push the black plastic carriage locking clip (A) as far as possible to the left and place the sample needle carriage (B) as far forward as possible to the right-hand side, as shown in Diag. 5.

Check that the needle is not bent and make sure it is in its upper position.

Unscrew slightly the 2 screws of the WBC/HGB chamber protection cover (diag 6). Remove the cover and check that both chambers (RBC/PLT, WBC/HGB) are fixed properly in their clips and the electrode blocks are attached firmly to the chambers (Diag. 7).

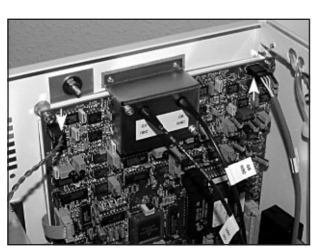


Diag.6

Re-install the HGB/WBC chamber cover.



Diag.7



Check that the connectors on the printed circuit board are securely in place (Diag. 8).

Re-install the instrument cover.

Diag.8



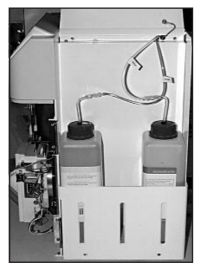
Diag.9

Remove the fuse holder from its location on the rear panel pressing on the holder lock (Diag.9) and check the fuse characteristics : they should be 1 Ampere, 250 Volts Slow-Blow.



4 - Reagent connections

• Bottle connections



Lyse and cleaning reagents are placed inside the reagent compartment as shown in the Diag. 10. Install the reagent straws and the bottle stoppers. Connect the blue tube to the MINICLEAN bottle and the white tube to the MINILYSE bottle. Close the compartment cover.

Install the male connectors included in the installation kit at the liquid input and output located at the bottom of the instru-

Connect the diluent container (see CAU-

TION above) using the diluent straw and

a 3x6 cristal tube (1 meter maximum) on the diluent input located at the bottom of the instrument rear panel (Diag.11). Connect the waste container using the cristal tube 3 x 6 on the waste output, and place the waste container below the instrument level (under the bench).

ment rear panel (Diag.11).

Diag.10

IMPORTANT

When the ABX MICROS 60-*cT* is set up with the 16 or 18 parameters mode, it is mandatory to use specific MINILYSE LMG and MINIDIL LMG reagents.

CAUTION

The Diluent container will be located on the bench at the same level than the instrument.

Waste connection



Diag.11





Always follow the recommended procedures for waste disposal. Never connect the instrument wastes directly to the laboratory drain pipes. For each waste container, follow the neutralization procedure as described in the user manal.



• Reagent pack connection

Remove the reagent output protections, as well as the waste input protection (Diag.12 & 13)









Install the pack directly into the compartment of the instrument as shown on the Diag. 14, 15 & 16. Push the pack down in order to plug correctly the pack on the male connectors.



Diag.14



Diag.16

CAUTION

In order to avoid leak problems it is recommended not to unplug several times the same reagent pack.



Diag.15

The free male connector (see Diags. 14, 15 & 16) must be plugged on the pack upper valve in order to receive the waste liquids.

5 - Printer and instrument connections

• Instrument connection.

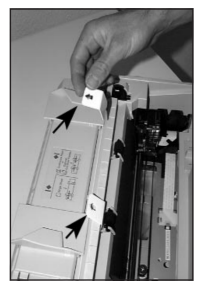
If the instrument has to be connected to a laboratory computer, use the plug RS232.

Connect the power cable to the plug located on the rear left-hand side of the device (Diag. 17).



Diag.17

- Setting up the printer
- Remove all the package protections
- Install the paper feed-knob



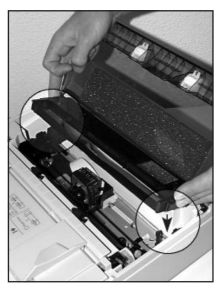


Diag.19

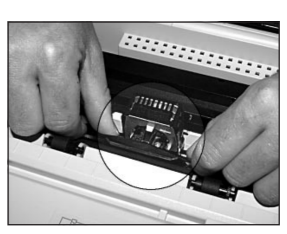
Diag.18

Open the ink ribbon access door at the top of the printer and install the ribbon as shown in diag 20 & 21 : Slide the printer head to the middle of the printer.

Insert the ribbon cartridge into the printer Guide the ribbon between the print head and ribbon guide. Slide the printer head from side to side to make sure it moves smoothly.



Diag.20



Diag.21

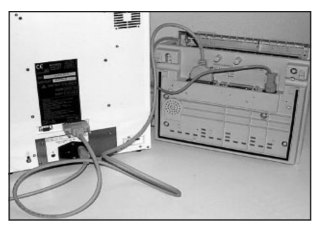
INSTALLATION

- Install the paper supports (Diag 22) for single sheets paper use only.





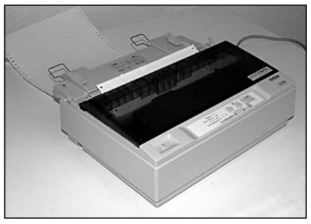
- The printer is connected to **ABX MICROS 60-***CT* with the cable delivered with the instrument. Lock the connector in place by tightening the 2 screws on each end of the connector to the **ABX MICROS 60-***CT*. Attach the other end of the cable to the printer and lock the printer connector in place by the means of the 2 clips located on the connector itself (Diag. 23).





- For continuous paper, introduce it in the slot at the back of the printer and use the sprocket covers to load paper, or else feed the paper frontward when using single sheets (see printer user manual).





Diag.25

Diag.24



• Printer command keys

Roman		•		Font	LE/FF	- BRE 1	Pause
Sans Serif		•		1		21	- BEERS
Draft	•				S.S. Margaret		
Draft Condensed	• •	• 1	2	L CIPe	per Park	ŀ'n	CONTRACTOR OF
	12/12	Micro	o Adjust	1	1	3sec	

Diag.26

LED PAUSE : The orange LED PAUSE lights when the printer stops printing. During each power ON, this LED blinks for few seconds and 4 audible beeps occur. When the printer runs out of paper, the LED blinks and 3 audible beeps occur. This LED lights also when the paper is in its tear off position. When a problem occurs, this LED lights ON and 5 audible beeps occur.

LEDS FONT 1 and FONT 2 : These 2 green LEDS indicate the selected font. Refer to the printer user's manual to select the font.

Key FONT : During normal operation, the FONT key allows the font selection. For each pressure on this key, the selection is modified. Refer to the printer user's manual to select the font. When this key is pressed during the printer power ON, the printer setup menu is entered.

Key LF/FF : During normal operation, a quick pressure on this key allows a ligne feed of the paper. Keep the pressure on this key to feed a whole page. This key can be used to load or eject the paper.

When this key is pressed during the printer power ON, the printing test starts. Key PAUSE : When this key is pressed during the printing, the printout stops. Press again on this key to restart the printout.

PAPER PARK : If Z folded paper is used, the paper can be driven to its parking position when pressing simultaneously on the keys LF/FF and FONT.

MICRO ADJUST : This function allows to adjust the loading paper position. See the user's manual for details.

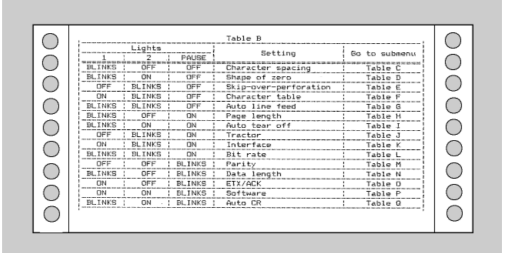
• Printer Configuration

Switch on the printer when pressing the . The configuration should be the following :

000000000	**************************************	**************************************	000000000000000000000000000000000000000
-----------	--	--	---

Diag.29

Each parameter can be modified by the corresponding parameter chart. Each chart is accessible using the keys <PAUSE>, and <LF/FF> according to the control LED combinations (Diag 30).



Diag.30

6 - Instrument startup

Reagent priming

When the **ABX MICROS 60-***ct* is first installed, it contains no reagents. All the reagents have to be primed now. Turn ON instrument by pressing the ON/OFF switch located on the rear panel. When the instrument turns on, the display shows :

PLEASE WAIT FOR 3 MIN ESCAPE : ESC

This time is required at the startup for the instrument initialization and stabilization, specifically for the HGB diode to reach its operational temperature. Press the ESC key in order to abort the cycle : the LED of the front panel turns from red to green and the display shows :

STARTUP NOT INITIATED PRESS A KEY TO CONTINUE...

This message appears when the instrument is setup with the manual startup cycle to prevent any analysis cycle before running a startup cycle. Press any key, the main menu is displayed :



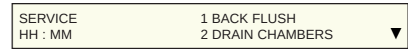
I C K



Bottles and containers set :

Install the reagent bottle and carry out a PRIME cycle to clear the reagent line of air bubbles. This procedure should be done whenever a new bottle of reagent is installed.

From the MAIN MENU, move the cursor to the function $\begin{pmatrix} 4 \\ \end{pmatrix}$ SERVICE and press ENTER. The service menu is displayed :



Move the cursor to function 3 PRIME REAGENTS and press ENTER. The PRIME menu is displayed :

PRIME	1 ALL REAGENTS	
HH : MM	2 DILUENT	/

Select either the function $\begin{pmatrix} 1 \\ 1 \end{pmatrix}$ prime ALL REAGENTS or move the cursor next to the required reagent and press ENTER.

The priming cycle starts while the following menu is displayed :

ALL REAGENTS	WAIT FOR 2 MN 3 S

IMPORTANT

Before analyzing samples, visually inspect reagent lines and pumps for air bubbles. Repeat priming if air bubbles are still present. Call the *ABX representative service department* if priming does not eliminate air bubbles.

Never initiate two Lyse prime cycles back-to-back. This causes excessive foaming in the waste chamber. Run a blank cycle between each Lyse prime cycle.

- Run a STARTUP cycle.

Reagent pack

1

From the MAIN MENU, move the cursor to the function $\begin{pmatrix} 4 \\ \end{pmatrix}$ SERVICE and press ENTER. The service menu is displayed :



Move the cursor to $\begin{pmatrix} 3 \\ \end{pmatrix}$ PRIME REAGENTS and press the ENTER key. Select the function

CHANGE PACK and follow the instructions given by the LCD in order to install the pack.

REAGENT PACK	> 1 CHANGE PACK	
HH : MM	2 CBC LEFT < 150>	▼

Once the new PACK is installed a priming cycle will be automatically carried out and the following menu is displayed.

PRIME

WAIT FOR 2 MIN 3 S

IMPORTANT

Before analyzing samples, visually inspect reagent lines and pumps for air bubbles. Repeat priming if air bubbles are still present. Call the *ABX representative service department* if priming does not eliminate air bubbles.

From the REAGENT PACK menu, the function $\binom{2}{}$ "CBC LEFT" displays the number of analysis cycles left to run with the same pack.

It is also possible to run a priming cycle at any time using the selection $\binom{3}{}$ "PRIME REAGENTS" of the SERVICE menu.

IMPORTANT

It is recommended not to remove the pack several times before the reagents are totally used in order to avoid leak problems.

- Run a STARTUP cycle.

INSTALLATION

Once the instrument ready for the analyses, remove the adhesive protection from the front panel (diag 31)



Diag.31

SAMPLING NEEDLE MAINTENANCE

· CONCERNS

- Needle O ring replacement
- Sampling needle replacement
- Piercing needle replacement

· REQUIRED TOOLS

- Hexagonal keys
- Dynamometric screw driver A302 : MAG 019 A

· REQUIRED PRODUCTS

- Silicone grease : LAM 004 A

INTERVENTION TIME

- 15min

FREQUENCY

- See frequency chart table for cleaning.
- O ring replacement : 1/year
- Needle replacement : On request only

SPECIFIC KIT OR CONSUMABLES

- O ring kit : XEA 328 AS
- Sampling needle : GBC 052 A
- Piercing needle : GBC 189 A

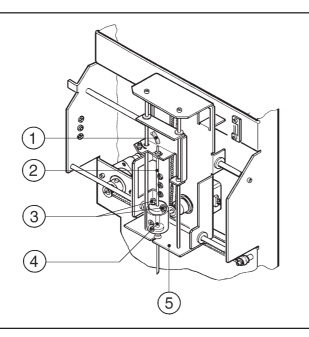




18/06/98

SAMPLING NEEDLE MAINTENANCE

PROCEDURE



Diag.1

3

A - Needle or O ring replacement

- Remove the rinsing block/needle assy from the carriage taking care not to bend the needle.

- Lift up the O ring holder (1) and re-

place the O rings 2 by new ones

previously greased. Wipe all excess of grease away.

- If necessary clean the inner surface of the rinsing block with a little piece of paper.

Diag.2

(1)

4

Proceed the same way to replace the needle if necessary

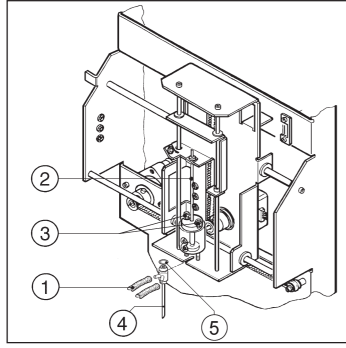
2

- Reassemble in the reverse order. Use a dynamometric screw driver to tighten the screws (3) (Diag.1) to **700 mN.m (99.4 Ozf.in)**.

SAMPLING NEEDLE MAINTENANCE

B - Piercing needle replacement

- Disconnect the tube 1 from the piercing needle 4 (Diag.3).



- Lift the needle 2 in the upper po-

- Loosen the 2 screws (3) just enough to enable the rinsing block to be lifted up of about 5 mm.

- Pull the piercing needle 4

(foreward) and replace it by a new one if necessary.

- Replace the piercing needle O ring

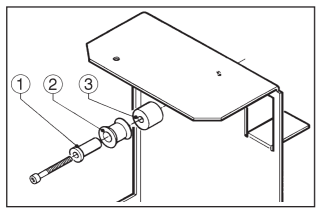
⁵ (FAA 036 A) by a new one.

- Reassemble in the reverse order. Apply the same torque : **700mN.m** (**99.4 Ozf.in**) to tighten the two screws (3).

Diag.3



It is recommended to check the correct motion of the needle. Proceed as follows : Enter the «SERVICE» menu and then the sub menu «MECHANIC» and perform a «NEEDLE U/D» cycle.



Diag.4

Blockage problems may occur on some instruments during the needle or carriage motions giving some motor error messages.

Before replacing the concerned motor, it is necessary to check the correct rotation of the free puley located at the end of the notched belt.

Remove the axle screw of the puley and clean its 2 parts and the washer. Reinstall the puley assy, the rounded edge facing the puley. Tighten the screw with a torque of **400mN.m (56.8 Ozf.in)**.

Check that the puley turns freely after the tightening. Add a drop of oil (LAM 007 A) if necessary.

1 - FAG 011 A : Autolub. axle 2 - GBC 146 A : Free pulley 3 - GBC 147 A : Pulley holder

SAMPLING NEEDLE MAINTENANCE

· CONCERNS

- Needle replacement

- O ring replacement

· REQUIRED TOOLS

- Hexagonal keys : 2,5
- Dynamometric screw driver A300 : MAG 013 A

· REQUIRED PRODUCTS

- Silicone grease : LAM 004 A

INTERVENTION TIME

- 15 min

• FREQUENCY

- Needle replacement : On request only
- O ring replacements : 1/year

· SPECIFIC KIT OR CONSUMABLES

- O ring kit : XEA 328 AS
- Spare parts kit : XEA 458 AS

RAS 168 A Ind.A



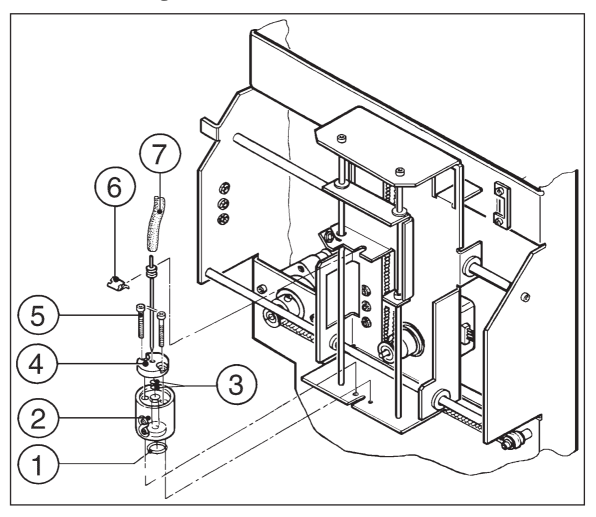
SAMPLING NEEDLE MAINTENANCE

PROCEDURE

- Disconnect the tube 7 from the top of the needle. Manually lift up the sampling needle (Diagram 1).

- Unscrew the 2 screws (5) in order to freed the needle rinsing block (2) from the carriage

frame. Remove the clip \bigcirc .



Diag.1

- Remove the rinsing block/needle assy from the carriage taking care not to bend the needle.

- If necessary clean the inner surface of the rinsing block by means of a little piece of soft paper. Spread a little amount of grease in between the rinsing block 2 and its support.

- Lift up the O ring holder (4) and replace the O rings (3) by new ones previously greased. Wipe all excess of grease away.

Proceed the same way to replace the needle if necessary

SAMPLING NEEDLE MAINTENANCE

- Reassemble in the reverse order. Use a dynamometric screw driver to tighten the screws (5) (Diag.1) to 100 mN.m.



It is recommended to check the correct motion of the needle. Proceed as follows : Enter the «SERVICE» menu and then the sub menu «MECHANIC» and perform a «NEEDLE UP/DOWN» cycle.

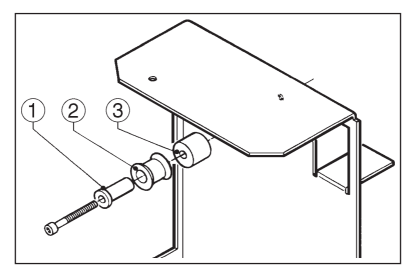
Blockage problems may occur on some instruments during the needle or carriage motions giving some motor error messages.

Before replacing the concerned motor, it is necessary to check the correct rotation of the free pulley located at the end of the notched belt.

Remove the axle screw of the pulley and clean its 2 parts and the washer.Reinstall the pulley assy, the rounded edge facing the pulley. Tighten the screw with a torque of **400mN.m (99.4 Ozf.in)**.

Check that the pulley turns freely after the tightening. Add a drop of oil (LAM 007 A) if necessary.

- 1) FAG 011 A : Autolub. axle
- 2 GBC 146 A : Free pulley
- 3 GBC 147 A : Pulley holder



Diag.2

CHAMBER MAINTENANCE



• CONCERNS

- RBC & WBC/HGB cleaning
- Aperture O ring replacement
- Coaxial O ring replacement

· REQUIRED TOOLS

- Hexagonal keys : 2,5
- Soft paper
- Dynamometric screw driver : MAG 013 A
- Cutting pliers

· REQUIRED PRODUCTS

- Liquid soap
- Distilled water
- A scalpel
- A Micropipette tip

INTERVENTION TIME

- 30 min

• FREQUENCY

- RBC & WBC cleaning : 2 (type 1 & 2) or 3/year (type 3).
- Aperture O ring replacement : 1/year
- Electrode O ring replacement : 1/year

• SPECIFIC KIT OR CONSUMABLES

- O ring kit : XEA 328 AS

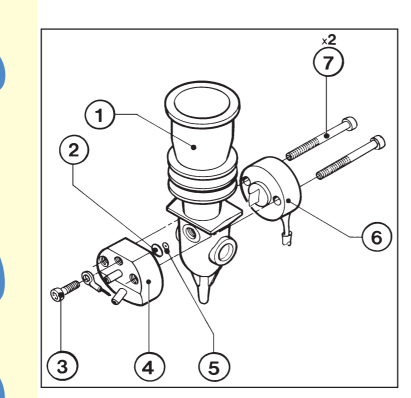


RAS 169 A Ind.A

CHAMBER MAINTENANCE

• PROCEDURE

1 - RBC chamber cleaning :



- Run a drain chamber cycle (SERVICE menu, DRAIN CHAMBERS Sub menu).

- Record the tube positions before dismantling the chambers.

- Disconnect the chamber tubes. - Unclip the RBC chamber.

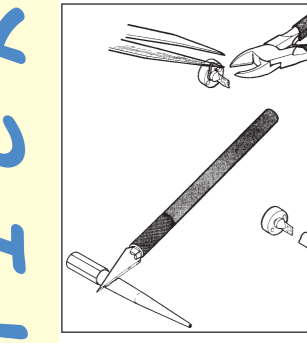
- Dismantle the electrode

6 loosening the 2 fixation

screws $\overbrace{7}$ and the terminal

holding screw (3) (Diagram 1).

Diag.1



• Electrode O ring replacement : - Use a previously cut micropipette tip to replace the electrode O ring as shown on Diag.2.

• Aperture O ring replacement :

- Install the chamber over a piece of white paper or cloth.

- Carefully remove the counting

head (4) and plunge the aper-

ture 5 in distilled water.

Replace the O ring 2 by a new one.

Diag.2

- Clean the chamber and the counting head with liquid soap, do not introduce any sharp instruments inside so as to avoid damaging the inside of the chamber and the aperture.

CHAMBER MAINTENANCE

CAUTION

Do not manipulate the aperture using hard instruments. Clean the aperture with a piece of soft paper or preferably, in between 2 fingers.

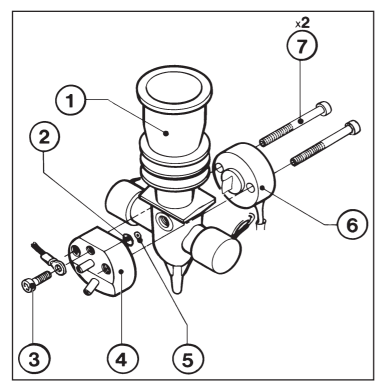
- Rinse thoroughly with distilled water
- Dry the exterior of the chamber with a soft paper.

CAUTION

Do not apply too much pressure on the electrode fixation screws, as it can break the aperture (tightening torque = 100mN.m / 14.2 Ozf.in).
It is recommended to reconnect the tubes on the counting head before reassembling the "electrode/chamber/counting head" assy in order to avoid applying constraint on the chamber.

- Position the chamber in its fixation clips.
- Reconnect the tubes

2 - WBC/HGB chamber cleaning :



Run a drain chamber cycle (SERVICE menu, DRAIN CHAMBERS Sub menu).
Loosen the cover screws of the WBC/HGB chamber and remove the cover.

- Record the tube positions before dismantling the chambers.

- Disconnect the chamber tubes.

- Unclip the RBC chamber.

- Dismantle the electrode

b) loosening the 2 fixation

screws (7) and the termi-

nal holding screw 3.

- Proceed as described in **1** - *RBC chamber* to clean the chamber and to replace the lectrode and aperture O rings.

0



Diag.3



The spectrophotometer can not be dismantled from the chamber. If this one has been dammaged it is necessary to replace the whole chamber assy. When cleaning the spectrophotometer, make sure to thoroughly rinse it in order to obtain a correct HGB blank measure.

LIQUID VALVE MAINTENANCE

• CONCERNS

Liquid valve assy replacement Valve body replacement



· REQUIRED TOOLS

Hexagonal keys Pair of pliers

S

· REQUIRED PRODUCTS

Soft paper

INTERVENTION TIME

15 min

• FREQUENCY

On request only

· SPECIFIC KIT OR CONSUMABLES

- 6 Valve assembly (MICROS 60 CT) : XDA 579 CS
- 6 Valve assembly (**MICROS 60** *OT*) : XDA 578 CS
- 5 Valve assembly : XDA 580 CS
- 2 ways NC liquid valve without solenoïd : XDA 481 B 3 ways liquid valve without solenoïd : XDA 483 B
- Solenoïd 24V 4W : EAZ 004 A

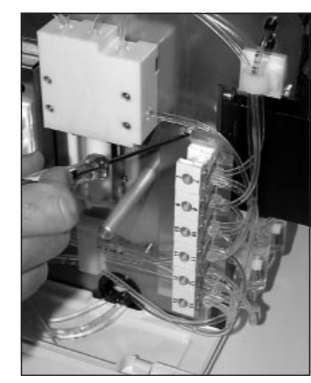
RAS 170 A Ind.A

LIQUID VALVE MAINTENANCE



- Several reasons can unable the correct operations of the valves :

- Leaks on the valve bodies
- Defective Solenoïds
- Liquid discharge on valves
- Corrosion traces on the axis, ect...
- Run a «DRAIN CHAMBERS» cycle.
- Switch off the instrument.
- Note the tube positions on the valve assembly that requires to be dismantled.
- Unscrew the valve assembly fixation screws (Diag.1).





Diag.1

Diag.2

- Disconnect the supplying flat cable from the valve assembly (diag.2).

NOTE

When leaks occur on valves it is recommended to replace the entire valve assembly by a new one.

- If only one valve has been damaged it is possible to dismantle the valve body on its own as shown on the diagram 3 : use a pair of pliers to disconnect the valve holder clip and remove the body.



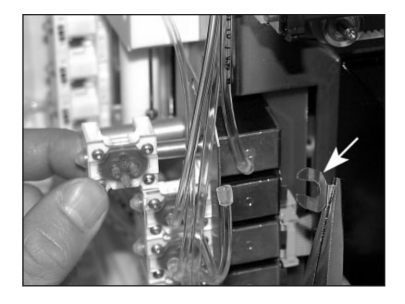
とこ



When replacing one valve only, it is important to check if the «neighboured» valves have not been damaged too.

LIQUID VALVE MAINTENANCE

When replacing one valve only, it is important to check if the «neighboured» valves have not been damaged too.



Diag.3



The solenoïd can not be dismantled unless unsoldering it. If this one is suspected solder a new one or replace the entire valve assembly.

- Reconnect the tubes on the nipples (with the sleeves).

- Re-install in the reverse order. Switch on the instrument.

- Control the watertightness of the valves and check for the correct operations :

Go to «SERVICE» menu, then to «MECHANIC» sub menu (5) and require a «VALVES» (6) test (see Mechanic functions : RAS 173 A).

- Check the calibration too.

• REMARKS

On MICROS 60 - CT, when piercing several times the same tube cap, some pieces of cork may

be dragged along towards the WBC/HGB chamber and then the liquid valve 2. This may damaged the operation of the valve.





POWER SUPPLY CHECK OR REPLACEMENT

• CONCERNS

- Voltage supply check
- Power supply module replacement
- Fan operation check

· REQUIRED TOOLS

- Hexagonal keys
- Flat screw driver
- Volmeter

· REQUIRED PRODUCTS

- None

· INTERVENTION TIME

- 15 min

• FREQUENCY

- 1/year

· SPECIFIC KIT OR CONSUMABLES

- None

RAS 171 A Ind.A



10/04/98

POWER SUPPLY CHECK OR REPLACEMENT

• PROCEDURE

1 - Supply voltage check

CAUTION

The supply voltage check has to be done with the power supply module connected only.

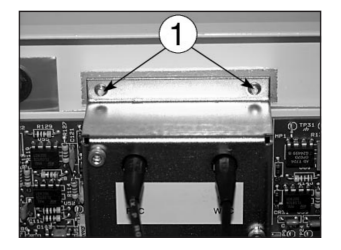
- The supply voltage check is done on the mother board (XAA 355 A) according to the following chart table :

* Ground on TP31, TP 30 or TP 29

TEST POINTS	VOLTAGES
TP 20	-12V <u>+</u> 0,5V
TP 22	24V + 1.5V - 0V
TP 23	5V + 0,3V - 0V
TP21	12V <u>+</u> 0,5V

CAUTION

The two screws on the top of the coaxial cover holding the board on the MICROS frame are the grounding connection of the mother board (diag 1). It is mandatory to check the correct tightening of these screws to obtain correct voltage values.

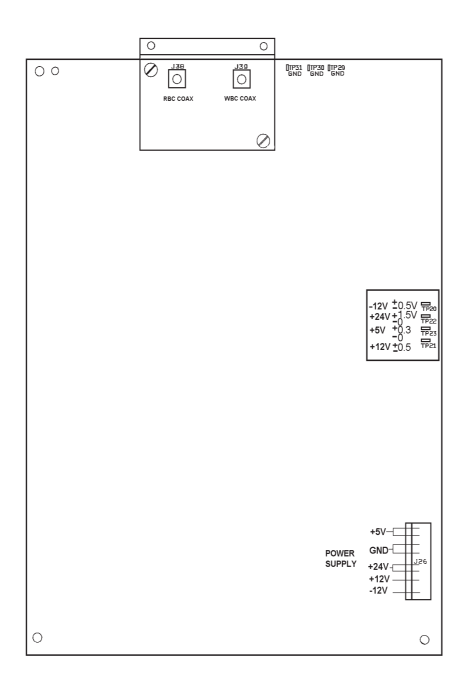


Diag.1

R



The test points are located as shown on the diagram below :



Diag.2



If the voltages values are not correct or among the ranges no adjustment can be carried out either on the board or on the power supply module. Replace the power supply module as described below :

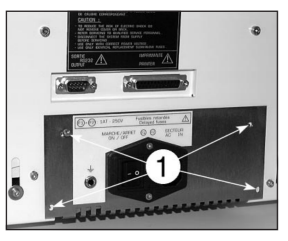
POWER SUPPLY CHECK OR REPLACEMENT

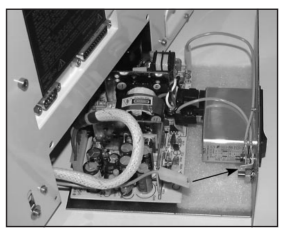


2 - Power supply module replacement

- Switch off the instrument.
- Disconnect the main supply voltage cable and the printer cable.
- Disconnect the power supply cable from the mother board, connector J26 (see Diag.2).

- Unscrew the 4 screws (1) (Diag.3) and start to move out the power supply module. Disconnect the grounding wire (Yellow/green wire see diag 4) from the rear panel of the module. Route the cable (from J26) down to make the removing of the module easier.





Diag.3

Diag.4

WARNING !

The power supply module internal fuse is not to be replaced even when this one has blown down.

- Replace the power supply module by a new one and reinstall in the reverse order.

- Switch on the instrument and check the voltages on the mother board as described in the previous paragraph.

• REMARKS

The four leds in front of the test points are lit to indicate a voltage presence but whatever its value!!!

Check the operation of the fan as following : When the fan has stopped, move the cursor to the 4 SERVICE menu and press ENTER : the fan should start.

TECHNICIAN FUNCTIO

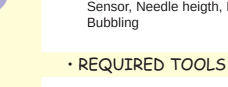
CONCERNS

1 - Version display. 2 - Adjustments

HGB photometer calibration, Aperture voltage Vacuum check, WBC gain, RBC & PLT gain Sensor, Needle heigth, Needle motion **Bubbling**

3 - Temperature sensor adjustment 4 - Run mode

- 5 Reagent pack
- 6 Serial number
- 7 Cycle number
- 8 Burn-in



- Flat screw driver
- thermometer
- Barflex
- Hexagonal keys
- Voltmeter

· REQUIRED PRODUCTS

- WBC latex : LAD 001 AS
- RBC and PLT latex : LAD 002 AS
- Soft paper
- Flat piece of stiff plastic

INTERVENTION TIME

- 60 min

FREQUENCY

N H C R O

- See maintenance chart table.

· SPECIFIC KIT OR CONSUMABLES

- Needle position tool : GBC 218 A

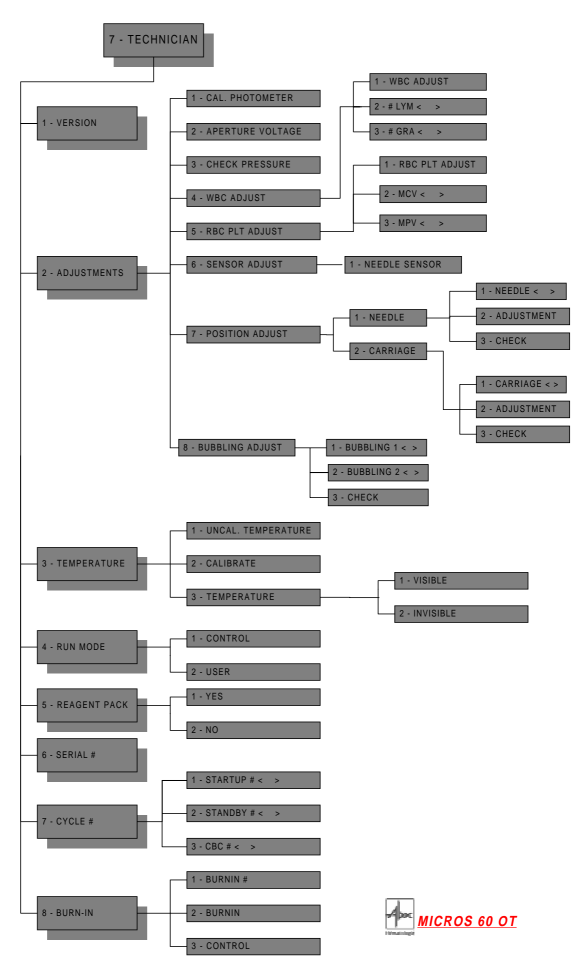


16/06/98



• PROCEDURE

TECHNICIAN FUNCTION



TECHNICIAN FUNCTION

Once entering the «SERVICE» sub menu, move to «TECHNICIAN FUNCTIONS» and press . A specific password (421) is required to enter the sub menus.

Move the cursor by means of result and result and choose the required menus pressing

the key.

I - The version number is displayed.

II - Adjustments :

1 - HGB photometer calibration

- Dismantle the WBC/HGB chamber cover.
- Check the general cleanliness of the WBC chamber/spectrophotometer assy.
- Re-install the chamber cover.

If the WBC chamber has been dismantled previously make sure no liquid has flown in between the spectrophotomer and the chamber. Clean the inner surfaces of the spectrophotometer as well as the chamber. Reassemble the assy and tighten the two screws to the following torque : 400mN.m (see RAS 169 A : Chamber maintenance)

- Run the CAL PHOTOMETER function (function 1 of the «ADJUSTMENTS» menu) : diluent is delivered to the WBC/HGB chamber twice.

An HGB channel is displayed on the LCD screen :

VALUE XXX

IMPORTANT

1 - The HGB photometer calibration must be done 20min at least after the instrument has been switched on.

2 - This adjustment must be done with the WBC chamber cover installed!!!

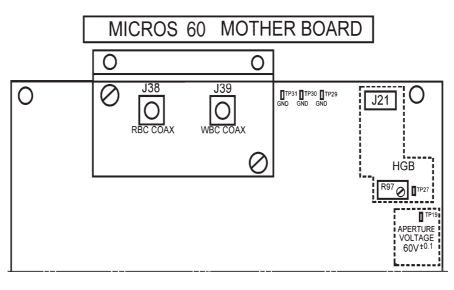
- By means of **R97** (See diagram 1) adjust the HGB channel according to the room temperature using the chart table given on next page.



After 40 seconds approximately, the function is automatically exited.

- Run the CAL PHOTOMETER function again to verify the adjustment.

TECHNICIAN FUNCTION



Diag.1

ROOM TPT (°C)				
	Mini.	Nominal	Maxi.	
15	240	245	250	
16	240	245	250	
17	239	244	249	
18	238	243	248	
19	237	242	247	
20	236	241	246	
21	235	240	245	
22	234	239	244	
23	234	239	244	
24	233	238	243	
25	232	237	242	
26	231	236	241	
27	230	235	240	
28	229	234	239	
29	228	233	238	
30	228	233	238	
31	227	232	237	
32	226	231	236	
33	225	230	235	
34	224	229	234	
35	223	228	233	

RUS

TECHNICIAN FUNCTION

2 - Aperture voltage

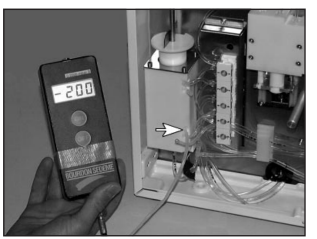
- Once entering the «APERTURE VOLTAGE» menu, connect a voltmeter between the ground (TP30) and TP19.

- Check that the value is 60V +2.8V -1.5V (Diagram 1). The aperture voltage is not adjustable.

- Press any key to escape.

3 - Vacuum check

Enter the 2 - ADJUSTMENTS / 3 - CHECK PRESSURE menu.



- Disconnect the tube from the vacuum/waste

syringe coming from the valve (s) (see diagram 2).

- Follow the instructions given on the LCD screen :

«PLEASE PLUG BARFLEX ON AIR SYRINGE» (On the free nipple).

«CHECK PRESSURE : -200mB ± 10mB» (The piston has raised in order to create a vacuum in the syringe body).

- Check the stability of the vacuum during 30 secondes : The vacuum drop down must be ≤ 2 mbar.

Diag.2

- If the results are not correct check the O ring and the tubing watertightness.

 $\ensuremath{\text{ eplus}}$ (disconnect the Barflex and replug the tube instead).

4 - WBC adjust

- Put the WBC latex to mix on a Vortex during 1min or shake thoroughly

IMPORTANT

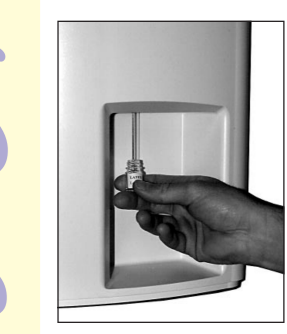
As the WBC gain is a factory adjustment it is mandatory not to readjust it without taking the following precautions :

Carry out previously, an autoconcentrated cleaning to make sure of the cleanliness of the WBC counting circuit.

If necessary clean the WBC chamber aperture as described in RAS 169 A. Make sure the Latex has been thoroughly mixed before.

- Run a blank cycle to check the cleanliness of the instrument.
- Enter the 4 WBC ADJUST sub menu and then 1 WBC ADJUST





- Present the vial of Latex to the open probe as shown on diagram 3 and press the sampling bar located behind the sampling needle : an analysis cycle begins.

During the cycle measuring phasis (around 1 minute) the **Lymphocyte** and the **Granulocyte** volumes are displayed on the screen every 3 seconds as shown below :

Diag.3

LYM <57 +/- 1>	GRA <180 +/- 2>	
57	180	

- Wait for several results to be displayed and check the stability of both values.

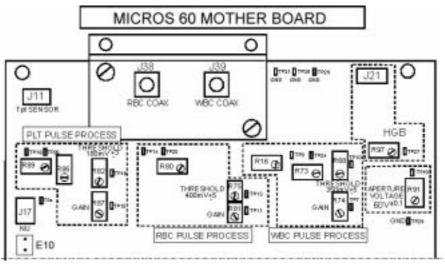
IMPORTAN7

It is mandatory not to operate the gain adjustment as long as the lympho and granulo values are not stable.

After one minute the last volume values displayed on the screen are printed out as well as the WBC, PLT and RBC histograms and the CBC's results. Check that these printed values correspond to the following target values :

-	LYM	= 57 ± 1
-	GRA	= 180 ± 2

- If not rerun a «WBC ADJUST» cycle after having previously mixed the latex vial again. By means of **R74** (see diagram 4) adjust the volumes to the target values during the measuring phasis.



Diag.4



Both sub menus 2 - # LYM < > and 3 - # GRA < > allow the technician to change the Latex target values if the latex run on the instrument different from the Latex recommended above.

5 - RBC PLT adjust

- Put the RBC and PLT latex to mix on a Vortex during 1min or shake thoroughly

IMPORTANT

As the RBC/PLT gain is a factory adjustment it is mandatory not to readjust it without taking the following precautions :

Carry out previously an autoconcentrated cleaning to make sure of the cleanliness of the RBC/PLT counting circuit.

If necessary clean the RBC/PLT chamber aperture as described in the procedure RAS 169 A.

Make sure the Latex has been thoroughly mixed before.

- Run a blank cycle to check the cleanliness of the instrument.
- Enter the «5 RBC PLT ADJUST» sub menu.

- Present the vial of Latex to the open probe as shown on diagram 3 and press the sampling bar located behind the sampling needle : the needle directly delivers the latex sample in the RBC chamber dilution and a measuring phasis begins.

- During the cycle measuring phasis (around 1 minute) the **Platelet** and the **Red Blood cell** volumes are displayed on the screen every 3 seconds as shown below :



- Wait for several results to be displayed and check the stability of both values.

IMPORTANT

It is mandatory not to operate the gain adjustment as long as the platelet and RBC values are not stable.

After one minute the last volume values displayed on the screen are printed out as well as the PLT and RBC histograms and the CBC's results.

- Check that these printed values correspond to the following target values :

- RBC = 74 ± 1 - PLT = 59 ± 1



From the latex **lot # 980311** included, balls having a different size, a drift of MPV peak has been noticed, i.e. a modification of the PLT gain target value : it becomes **64** instead of **59**. The program default value will be modified in the next MICROS version.

Both sub menus 2 - MCV < > and 3 - MPV < > allow the technician to change the Latex target values. If the lot $\# \ge 980311$, modify the target values and proceed the same way to adjust the PLT gain.

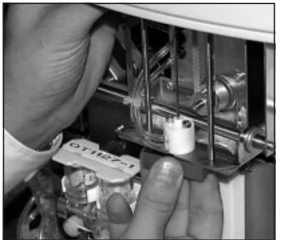
- If not, rerun a «RBC PLT ADJUST» cycle after having previously mixed the latex vial again.

- Adjust the volumes to the target values during the measuring phasis by means of the following potentiometers (see Diagram 4) :

- RBC gain with R81
- PLT gain with R87

6 - Needle sensor adjust

If the needle detector has been replaced by a new one or dismantled for any reason, it is mandatory to re-position it at the right heigth. Proceed as following :



- Install the piece of plastic (diagram 5) underneath the needle rinsing block.

- Once entering the menu «6 - SENSOR ADJUST», enter the sub menu «1 - NEEDLE SENSOR».

- Push the sampling needle downward until it stops against the piece of plastic and press any key in order to raise the needle back in its upper position.

The current number of steps, the mini and maxi values are displayed as well as the way to move the sensor (shown by an arrow) if the current value is out of ranges (see below).

Diag.5



- For a current number of steps out of ranges, unloosen the 2 cell fixation screws (diagram 6) and gently move the sensor

- upward if the current value is too low
- downward if the current value is too high.

- Tighten the screws and rerun a «NEEDLE SENSOR» cycle. Check that the current value is correct.

Diag.6



10 steps correspond to around 1 mm. The target number of steps is 70 \pm 5

7 - Position adjustment

• Needle heigth adjustment

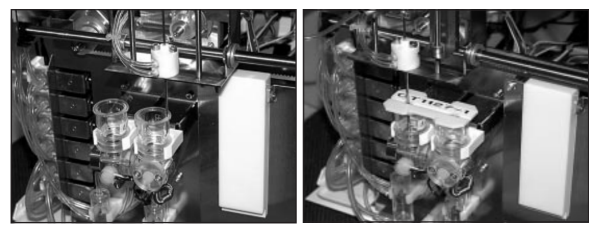
When replacing a needle, it is mandatory to adjust the heigth of the needle in the chambers. Proceed as follows :

- Remove the WBC/HGB chamber cover.

- Enter the menu 7 - POSITION ADJUST / 1 - NEEDLE / 2 - ADJUSTMENT.

- Manually pull down the needle until it comes into contact with the edge of the RBC chamber (Diag. 7).

- Press a key to continue : the needle moves up to the initial position ; the value is stored.



Diag.7

Diag.8

Needle motion adjustment

The needle position in the WBC chamber can be adjusted as follows :

- Enter the menu 7 POSITION ADJUST / 2 CARRIAGE / 2 ADJUSTMENT.
- Position the tool (P/N : GBC 218 A) over the RBC and WBC chambers (Diag 8).
- Manually lower the needle into the WBC chamber.

- Press a key to continue : the needle moves up, and the carriage comes back to the initial position ; the value is stored.

- Carry out a 3 CHECK cycle : the needle comes down to the WBC chamber.
- Check that the needle is centered in the hole.

If not, enter the 2 - CARRIAGE / 1 - CARRIAGE < > menu ; the display shows the current number of steps carriage motion.

If the needle goes too far on the right, add 1 step to the current value for 0.1mm. If the needle is too much on the left, decrease the current value of 1 step for 0.1mm.

CARRIAGE ? :	EXIT : ESC
CURRENT : 893	SAVE : ENTER
OORALEAT . 000	

Carry out a 2 - CARRIAGE / 3 - CHECK cycle again to control the needle position.

8 - Bubbling adjustment

An overflow protection tank is installed on the drain circuit of each chamber. This one prevents from polluted liquid overflow during bubbling phasis (Diag .9).



Diag.9

Two bubbling phasis are adjustable :

- "BUBBLING 1" is the first dilution (WBC/HGB chamber) bubbling value.

- "BUBBLING 2" is the second dilution (WBC/HGB chamber + lyse) value and RBC chamber bubbling value.

Both values correspond to a number of steps carried out by the waste/vacuum syringe. Default values are BUBBLING 1 : 175 BUBBLING 2 : 120

CAUTION

These values are factory adjusted (and may be different from the default values shown above) and should be modified only when hematologic erroneous results are given by the instument : If values are too important, liquid overflows can occur or if bubbling is too low homogeneity of the dilution can be decreased. Ranges : 150 < BUBBLING 1 < 200 80 < BUBBLING 2 < 140

To modify the bubbling values, enter the menu :

2 - ADJUSTMENTS / 8 - BUBBLING ADJUST. / BUBBLING 1 < > 2 - ADJUSTMENTS / 8 - BUBBLING ADJUST. / BUBBLING 2 < > and type in new step value.

Carry out a 3 - CHECK to control the adjustment.

III - Temperature

When entering the «temperature adjustment» menu the following sub menus are displayed :

1 - Uncal. temperature

When pressing the \sum_{Enter} key the sensor temperature value **uncalibrated** is displayed. This value should be close to the diluent temperature.

2 - Calibrate :

The temperature must be calibrated according to the diluent temperature :

- Plunge a thermometer directly into the diluent container and leave it for a while until stabilization.



For a pack equipped instrument, the thermometer must be plunged in the WBC/HGB chamber and the temperature must be note as soon as possible.

- Run 2 diluent primes («SERVICE» menu, «PRIME» sub menu, «DILUENT» selection).

- Enter the «CALIBRATE» menu. Note the temperature of the diluent and type in the value (if it is different from the previous on the instrument).

- Press on to save the new value.

3 - Temperature :

- 1 Visible : Press The value displayed is the calibrated temperature.
- 2 Invisible : Press to cancel the temperature display. (Temperature invisible by default)

IV - Run mode

The instrument must be configurated in the «USER» mode (configurated by default in the «USER « mode), the «CONTROL» mode intends for a factory use.

V - Reagent pack

This function is used to update the instrument from a bottle mode to a pack mode.

VI - Serial

Displays the instrument serial number.

VII - Cycle

Displays the : Startup number since the first use of the instrument. Stand by number since the first use of the instrument. CBC number since the first use of the instrument.

IMPORTANT

The startup, stand by and CBC numbers are adjustable in this menu but it is mandatory to keep the initial values (useful for maintenance schedules).

VIII - Burn-in

This function which allows the burn-in of the instrument is intended for a factory use.

RAS 172 A Ind.A

MECHANIC FUNCTION

CONCERNS

- Sensor replacements
- Needle motion check
- Carriage motion check
- Liquid syringe motion check
- Vacuum/waste syringe motion check
- Valve operation check
- LCD contrast adjustment
- Piercing mechanism check (MICROS 60 CT)

• REQUIRED TOOLS

- Hexagonal keys
- Felt-pen

· REQUIRED PRODUCTS

- None

INTERVENTION TIME



FREQUENCY

- See maintenance chart table.

· SPECIFIC KIT OR CONSUMABLES

- Vacuum/waste syringe sensor : **XBA 319 AS**
- Liquid syringe sensor :
- **XBA 319 AS** XBA 250 A - Carriage and needle sensors :





RAS 173 A Ind A

MECHANIC FUNCTION

• PROCEDURE

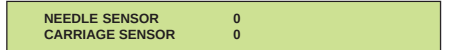
The MECHANIC function arborescence is as follows :

CHANIC	1 - Sensors
	2 - Needle U/D
	3 - Carriage L/R
	4 - Liquid syringe
	5 - Pres. syringe
	6 - Valves
	7 - Chg. Contrast
	8 - Park
	9 - Piercing

From the main menu, enter the 4 - SERVICE menu and move to 5 - MECHANIC sub menu.

1 - Sensors







should be displayed on the LCD screen.

Manually raise the needle support in the upper position (see diag 1)

The «0» should switch to «1». This indicates the correct operation of the needle sensor (diagram 1).

If nothing happened try to move the needle up and down again to get the commutation from <0> to <1>.

If the test is still wrong, check the correct connection of the sensor on the connector J7 of the mother board.

• Replacing the needle sensor

Switch off the instrument.

Use a felt-pen to mark the sensor position.

Unscrew the 2 sensor fixation screws and remove the cell holder .

Disconnect the wire from the J7 connector (See diagram 2).

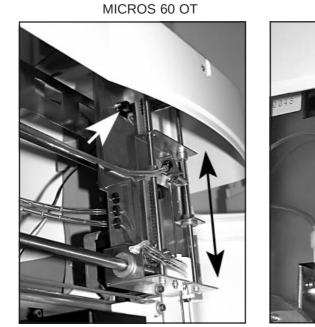
Replace the previous sensor by a new one and reassemble in the reverse order.



The new sensor must be installed exactly on the same position.

Proceed as described in **RAS 172 A (MICROS 60 OT)** or **RAS 176 A (MICROS 60 CT)** «Sensor adjustment» to check the correct position of the sensor.

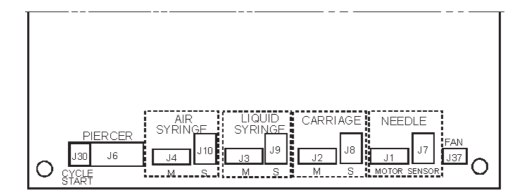






MCROS 60 CT

Diag.1



Diag.2

• Replacing the carriage sensor

Use the 1 - SENSOR test to check the carriage motion detection :

Move rightward the needle carriage in order to perform the sensor detection (diagram 3). To replace it, note its position with a felt-pen, unscrew the cell fixation screws and remove the cell holder.

Disconnect the wire from the connector J8 (See diagram 2)

Replace the previous sensor by a new one and reassemble in the reverse order.



The new sensor must be installed exactly on the same position.

Proceed as described in **RAS 172 A (MICROS 60 OT)** or **RAS 176 A (MICROS 60 CT)** «Sensor adjustment» to check the correct position of the sensor.

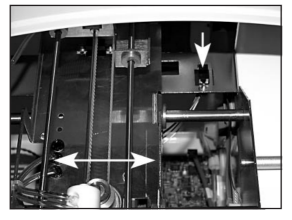


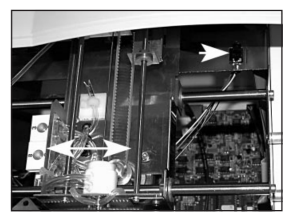


MECHANIC FUNCTION

MICROS 60 OT

MICROS 60 CT





Diag.3

2 - Liquid syringes motion check

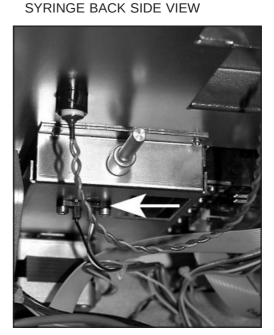
Run a 4 - LIQUID SYRINGE cycle : The syringes (diag 4) are moved upward and downward once.

If the sensor (diag 5) detection is not correct a message «ERROR : SYRINGE MOTOR» is displayed.

Three parts are involved :

- The motor ———> Check the connection on the board (connector J3)
 The syringes ———> Check the motion up/down
 The sensor ———> Check the connection on the board or replace it.

- SYRINGE FRONT SIDE VIEW



Diag.5

• Replacing the sensor :

Diag.4

- Switch off the instrument.
- Unscrew the 2 screws (Diag.5) and disconnect the wire from the connector J9 (Diag.2)
- Replace the previous sensor by a new one and re-install in the reverse order.
 - Switch on the instrument.
 - Run a 4 LIQUID SYRINGE cycle again to control the correct detection.



RAS 173 A Ind A



3 - Vacuum/waste syringe motion check

Run a 5 - PRESSURE SYRINGE cycle : The syringe piston moves upward and downward once.

If the sensor (Diag.6) detection is not correct a message «ERROR : PRESSURE SYRINGE MOTOR» is displayed.

Three parts are involved :

- The motor ———> Check the connection on the board (connector J4)
- The syringe ———> Check the motion up/down
- The sensor _____> Check the connection on the board or replace it.



Diag.6

- Replacing the sensor :
- Switch off the instrument.
- Unscrew the 2 cell fixation screws (Diag.6) and disconnect the wire from the connector J10 (Diag.2)
- Replace the previous sensor by a new one and re-install in the reverse order.
- Switch on the instrument.
- Run a 5 PRESSURE SYRINGE cycle again to control the correct detection.

4 - Valve operation check

Run a 6 - VALVES cycle. The number of the valves from 1 to 13 is displayed and each valve is activated once.

Check their correct operations. If a valve is suspected proceed as described in the procedure : RAS 170 A.



MECHANIC FUNCTION

5 - LCD contrast adjustment

Enter the sub menu «7 - CHG. CONTRAST». Use the arrows to modify the contrast and press



to validate the adjustment.

If the LCD screen happens to be unreadable because of a bad contrast adjustment it is possible to get back a correct contrast pressing at the

DEL

same time both keys : 🄇

6 - Piercing mechanism check (MICROS 60 CT)

Enter the sub menu 9 - PIERCING. Follow the instructions : «PLEASE CLOSE TUBE HOLDER DOOR» : a piercing operation is simulated.

This function gives the sample tube holder position and the heigth of the needle in its lower position. See RAS 172 A (MICROS 60 *OT*) or RAS 176 A (MICROS 60 *CT*)

NOTE

If the sample tube holder has been removed the following message is displayed : «ERROR : NO SAMPLE TUBE HOLDER». If the sample tube holder has been turned in between two piercing positions (it means that the tube is not in front of the piercing needle) the following message is displayed : «ERROR : TUBE HOLDER POSITION».

アンマン

DRAIN DETECTION

· CONCERNS

0

- Drain detection sensor adjustment/replacement

· REQUIRED TOOLS

- Voltmeter
- Flat screw driver
- 5ml syringe

· REQUIRED PRODUCTS

- None

INTERVENTION TIME

- 15 min

• FREQUENCY

- On request or once a year

· SPECIFIC KIT OR CONSUMABLES

- Drain detection sensor : XBA 199 A.



RAS 174 A Ind.B

DRAIN DETECTION

PROCEDURE

The Vacuum/Waste syringe and the chambers drains are controled by an infrared sensor located below the syringe.

Controls carried out by the cell during a cycle are as follows :

- Control of the correct operation of the cell

At the first chamber drain, air must be detected in the cell within defined timeout, and followed by liquids.

If this switch "air-liquid" has been successfull, the adjustment of the cell (see below) is validated. If not, the cycle is stopped and the following message is triggered : *"sensor error or diluent empty"* (check the connection or the adjustment of the cell).

- Control of the chamber drains

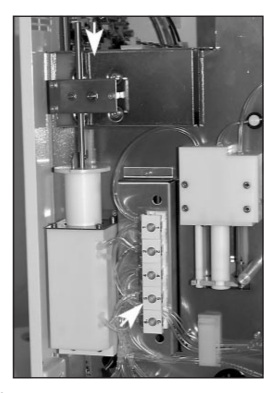
A first measure controls that liquids circulate in the cell during a defined timeout. The second measure checks that air has replaced liquid. If so, the drain phasis is validated.

If not, the following message is displayed : «.....» (the instrument carries on the current cycle) It means that the sensor always detects liquids. (Check the watertightness of the syringe.)

• Drain detection sensor adjustment

- Raise the piston up and press the valve <2>.

- Manually perform a syringe drain pressing the valve <5> and pulling down the syringe piston (Diag.1).



- Make sure the cell is perfectly drained (no bubble).

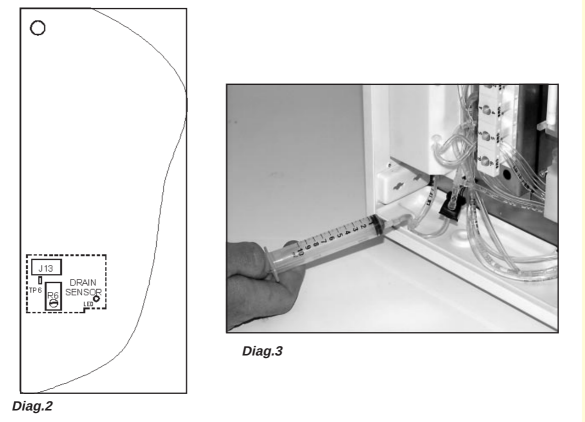
- Connect a voltmeter between the ground **TP30** and **TP6** (See Diag.2) and adjust the voltage to **4,5V** ± **0,3V** by means of **R6**.

Diag.1

DRAIN DETECTION

Fill the syringe up with diluent and connect it on the detection cell as shown on diagram 3.

MICROS 60 MOTHER BOARD



- Push diluent through the sensor and check that the voltage falls down below **1Volt** on the voltmeter.



Check the commutation from 4,5Volts to \geq 0Volt once again pushing and drawning alternately liquid and air through the sensor by means of the syringe.

- If nothing happens switch off the instrument and disconnect the connector from J13.
- Replace the drain detection sensor by a new one.
- Switch on the instrument
- Carry out the new sensor adjustment as described above.



However it is posible to control the correct operation of the sensor by means of the LED located next to **R6**. Indeed the LED should be lit when liquid circulates through the sensor and switched off with air.



Account of the shold check/adjustment Put threshold check/adjustment

· REQUIRED TOOLS

- Voltmeter
- Flat screw driver

S

REQUIRED PRODUCTS
 - None

INTERVENTION TIME

- 20min

• FREQUENCY

- See maintenance chart table.

· SPECIFIC KIT OR CONSUMABLES

- None

RAS 175 A Ind.B

PCB VOLTAGE CHECKS/ADJUSTMENTS

• PROCEDURE

1 - RBC, PLT, WBC threshold checks/adjustments

- Ground on TP31.
- Adjust the thresholds according to the below chart table :

THRESHOLDS	TEST POINTS	VOLTAGE	POTENTIOMETERS
WBC	TP 10	350 mV <u>+</u> 7	R68
RBC	TP13	400 mV <u>+</u> 7	R75
PLT	TP16	180 mV <u>+</u> 3	R82



(See Diag.1)

2 - Aperture voltage check

See RAS 172 A : MICROS 60 OT Technician function RAS 176 A : MICROS 60 CT Technician function

3 - Voltage supply check See RAS 171 A



4 - HGB blank voltage check

See RAS 172 A : MICROS 60 *OT* Technician function RAS 176 A : MICROS 60 *CT* Technician function

5 - Stepper motor voltage checks/adjustments

- Ground on TP31.

- Adjust the motor voltages according to the below chart table :

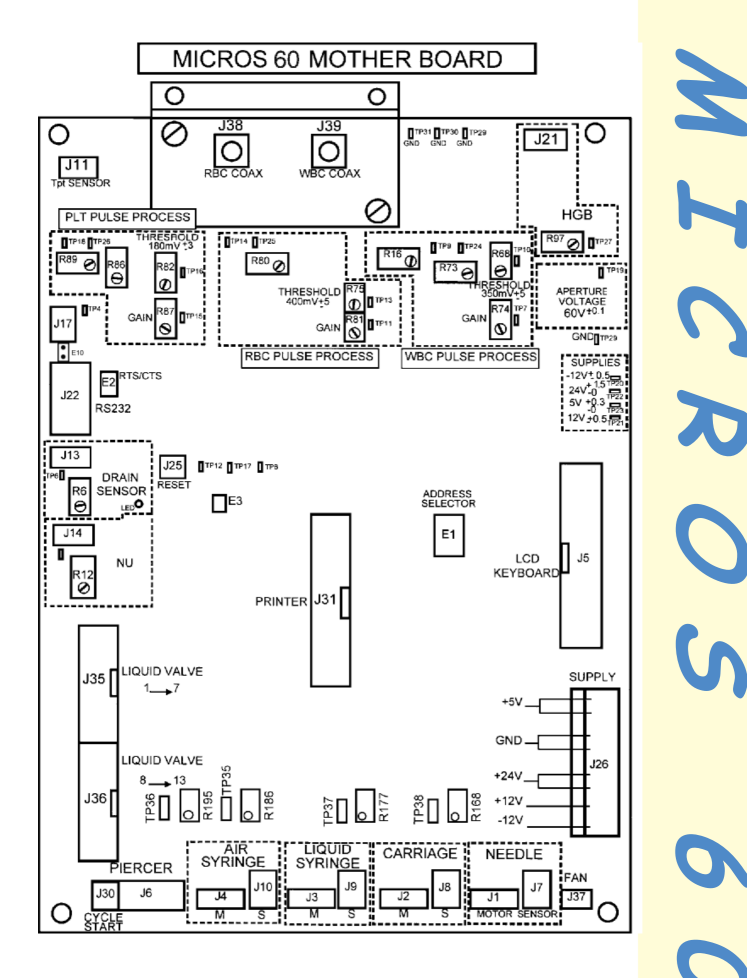
MOTORS	TEST POINTS	VOLTAGES	POTENTIOMETERS
Air syringe	TP36	2.50 V <u>+</u> 0.05 V	R195
Liquid syringe	TP35	2.00 V <u>+</u> 0.05 V	R186
Horizontal carriage	TP37	1.50 V <u>+</u> 0.05 V	R177
Vertical carriage	TP38	1.00 V <u>+</u> 0.05 V	R168



(See Diag.1)

Page 2/4

PCB VOLTAGE CHECKS/ADJUSTMENTS



• CONCERNS

- 1 Version display.
- 2 Adjustments :

HGB photometer calibration, Aperture voltage Vacuum check, WBC gain, RBC & PLT gain Sensor, Needle heigth, Needle motion Bubbling

• REQUIRED TOOLS

- Flat screw driver
- thermometer
- Barflex
- Hexagonal keys
- Voltmeter

· REQUIRED PRODUCTS

- WBC latex : LAD 001 AS
- RBC and PLT latex : LAD 002 AS
- Soft paper
- Flat piece of stiff plastic

INTERVENTION TIME

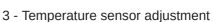
- 60 min

• FREQUENCY

- See maintenance chart table.

· SPECIFIC KIT OR CONSUMABLES

- None



- 4 Run mode
- 5 Reagent pack
- 6 Serial number
- 7 Cycle number
- 8 Burn-in



16/06/98

RAS 176 A Ind.A

>

>

1 - NEEDLE < > 2 - ADJUSTMENT

1 - CARRIAGE < >

2 - ADJUSTMENT

3 - CHECK

3 - CHECK

1 - NEEDLE ADJUSTMENT

2 - NEEDLE 1 < >

3 - NEEDLE 2 < >

4 - NEEDLE 3 < >

6 - NEEDLE 5 < >

7 - NEEDLE 6 < >

MICROS 60 CT

< >

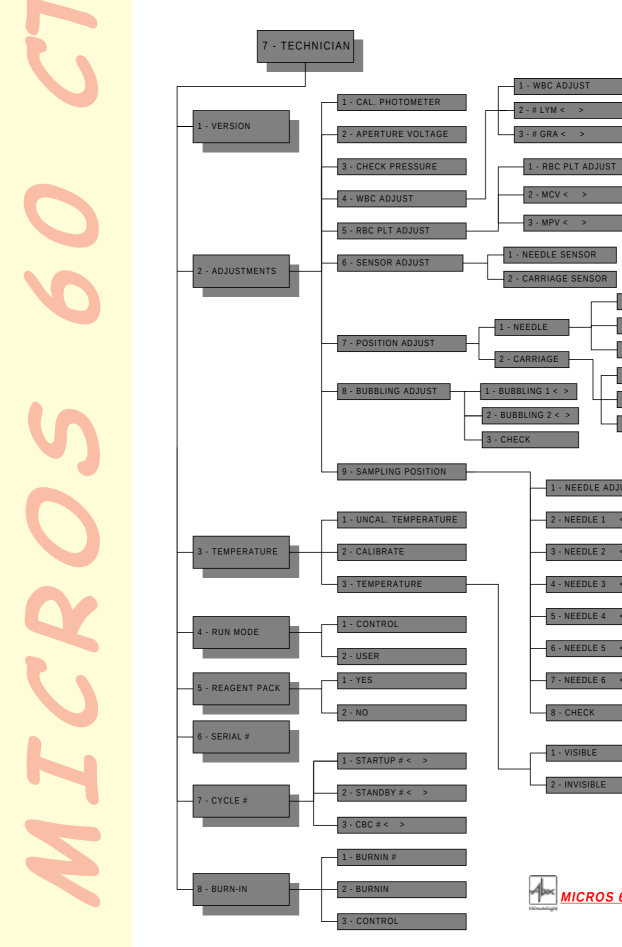
5 - NEEDLE 4

8 - CHECK

1 - VISIBLE

2 - INVISIBLE

PROCEDURE



Once entering the «SERVICE» sub menu, move to «TECHNICIAN FUNCTIONS» and press

. A specific password (421) is required to enter the sub menus.

Move the cursor by means of value and value and choose the required menus pressing

the key.

- I The version number is displayed.
- II Adjustments :

1 - HGB photometer calibration

- Dismantle the WBC/HGB chamber cover.
- Check the general cleanliness of the WBC chamber/spectrophotometer assy.
- Re-install the chamber cover.

If the WBC/HGB chamber has been dismantled previously make sure no liquid has flown in between the spectrophotomer and the chamber. Clean the inner surfaces of the spectrophotometer as well as the chamber. Reassemble the assy and tighten the two screws to the following torque : 400mN.m (see RAS 169 A : Chamber maintenance)

- Run the CAL PHOTOMETER function (selection 1 of the «ADJUSTMENTS» menu) : diluent is delivered to the WBC/HGB chamber twice.

An HGB channel is displayed on the LCD screen :

VALUE XXX

IMPORTANT

1 - The HGB photometer calibration must be done 20min at least after the instrument has been switched on.

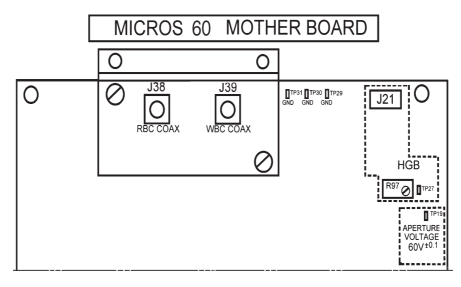
2 - This adjustment must be done with the WBC chamber cover installed!!!

- By means of **R97** (See diagram 1) adjust the HGB channel according to the room temperature using the chart table given on next page.



After 40 seconds approximately, the function is automatically exited.

- Run the CAL PHOTOMETER function again to verify the adjustment.





ROOM TPT (°C)	CHANNEL		
	Mini.	Nominal	Maxi.
15	240	245	250
16	240	245	250
17	239	244	249
18	238	243	248
19	237	242	247
20	236	241	246
21	235	240	245
22	234	239	244
23	234	239	244
24	233	238	243
25	232	237	242
26	231	236	241
27	230	235	240
28	229	234	239
29	228	233	238
30	228	233	238
31	227	232	237
32	226	231	236
33	225	230	235
34	224	229	234
35	223	228	233

0
S
Q
S
H
Z

RAS 176 A Ind.A

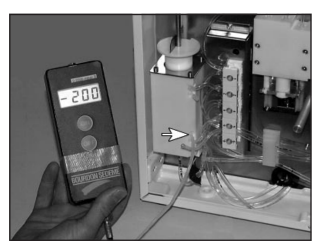


2 - Aperture voltage

- Once entering the «APERTURE VOLTAGE» menu, connect a voltmeter between the ground (TP30) and TP19.

- Check that the value is 60V +2.8V -1.5V (Diagram 1). The aperture voltage is not adjustable.
- Press any key to escape.

3 - Vacuum check



- Disconnect the tube from the vacuum/ waste syringe coming from the valve (see diagram 2).

- Follow the instructions given on the LCD screen :

«PLEASE PLUG BARFLEX ON AIR SYRINGE» (On the free nipple).

«CHECK PRESSURE : -200mB ± 10mB» (The piston has raised in order to create a vacuum in the syringe body).

- Check the stability of the vacuum during 30 secondes : The vacuum drop down must be ≤ 2 mbar.

Diag.2

- If the results are not correct check the O ring and the tubing watertightness.

 $\ensuremath{\text{ eplus}}$ (disconnect the Barflex and replug the tube instead).

4 - WBC adjust

- Put the WBC latex to mix on a Vortex during 1min or shake thoroughly

IMPORTANT

As the WBC gain is a factory adjustment it is mandatory not to readjust it without taking the following precautions :

Carry out previously an autoconcentrated cleaning to make sure of the cleanliness of the WBC counting circuit.

If necessary clean the WBC chamber aperture as described in RAS 169 A. Make sure the Latex has been thoroughly mixed before.

- Run a blank cycle to check the cleanliness of the instrument.
- Enter the «WBC ADJUST» sub menu.

TECHNICIAN FUNCTION

- Enter the «WBC ADJUST» sub menu and close the door of the piercing mechanism (See Diag 3) : A CBC's cycle starts.

During the cycle measuring phasis (around 1 minute) the **Lymphocyte** and the **Granulocyte** volumes are displayed on the screen every 3 seconds as shown below :

Diag.3

LYM <57 +/- 1>	GRA <180 +/- 2>	
57	180	

- Wait for several results to be displayed and check the stability of both values.

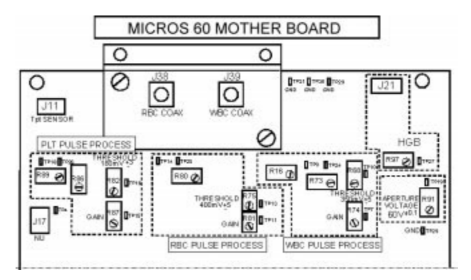
IMPORTANT

It is mandatory not to operate the gain adjustment as long as the lympho and granulo values are not stable.

After one minute the last volume values displayed on the screen are printed out as well as the WBC, PLT and RBC histograms and the CBC's results. Check that these printed values correspond to the following target values :

- LYM = 57 ± 1 - GRA = 180 ± 2

- If not rerun a «WBC ADJUST» cycle after having previously mixed the latex vial again. By means of **R74** (see diagram 4) adjust the volumes to the target values during the measuring phasis.



Diag.4

RAS 176 A Ind.A





Both sub menus 2 - # LYM < > and 3 - # GRA < > allow the technician to change the Latex target values if the latex run on the instrument is different from the one recommended above.

5 - RBC PLT adjust

- Put the RBC and PLT latex to mix on a Vortex during 1min or shake thoroughly

IMPORTANT

As the RBC/PLT gain is a factory adjustment it is mandatory not to readjust it without taking the following precautions :

Carry out previously an autoconcentrated cleaning to make sure of the cleanliness of the RBC/PLT counting circuit.

If necessary clean the RBC/PLT chamber aperture as described in the procedure RAS 169 A.

Make sure the Latex has been thoroughly mixed before.

- Run a blank cycle to check the cleanliness of the instrument.
- Enter the «RBC PLT ADJUST» sub menu.

- Present the vial of Latex to the open probe as shown on diagram 3 and press the sampling bar located behind the sampling needle : the needle directly delivers the latex sample in the RBC chamber dilution (for a usual analysis cycle the sample is first delivers to the mixing chamber) and a measuring phasis begins.

- During the cycle measuring phasis (around 1 minute) the **Platelet** and the **Red Blood cell** volumes are displayed on the screen every 3 seconds as shown below :

- Wait for several results to be displayed and check the stability of both values.

IMPORTANT

It is mandatory not to operate the gain adjustment as long as the platelet and RBC values are not stable.



From the latex **lot # 980311** included, balls having a different size, a drift of MPV peak has been noticed, i.e. a modification of the PLT gain target value : it becomes **64** instead of **59**.

The program default value will be modified in the next MICROS version. Both sub menus 2 - MCV < > and 3 - MPV < > allow the technician to change the Latex target values. If the lot $\# \ge 980311$, modify the target values and proceed the same way to adjust the PLT gain.

- If not, rerun a «RBC PLT ADJUST» cycle after having previously mixed the latex vial again.

- Adjust the volumes to the target values during the measuring phasis by means of the following potentiometers (see Diagram 4) :

- RBC gain with R81
- PLT gain with R87

6 - Sensor adjust

Needle sensor

If the needle sensor (diag 6) has been replaced by a new one or dismantled for any reason, it is mandatory to re-position it at the right heigth. Proceed as following :



- Install the piece of plastic (diagram 5) underneath the needle rinsing block.

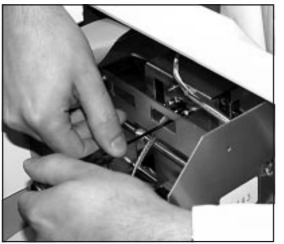
- Once entering the menu «SENSOR ADJUST», enter the sub menu «NEEDLE SENSOR».

- Push the sampling needle downward until it stops against the piece of plastic and press any key in order to raise the needle back in its upper position.

The current number of steps, the mini and maxi values are displayed as well as the way to remove the sensor (shown by an arrow) if the current value is out of ranges (see below).

Diag.5





- For a current number of steps out of ranges, unloosen the 2 cell fixation screws (diagram 6) and gently move the sensor

upward if the current value is too low
downward if the current value is too high.

- Tighten the screws and rerun a «NEEDLE SENSOR» cycle. Check that the current value is correct.

Diag.6



10 steps correspond to around 1 mm. The target number of steps is 70 ± 5

Carriage sensor

This function allows the adjustment of the carriage sensor (diagram 8) position.

- Proceed as described below :



Diag.7

- Install the piece of plastic against the left side of the tube holder compartment (see Diagram 7).

- Once entering the menu «SENSOR ADJUST» (Selection 6), enter the sub menu «CARRIAGE SENSOR».

Move the carriage on the left until the piercing needle stops against the piece of plastic. Press any key : the carriage comes back in its initial position.

The current number of steps, the mini and maxi values are displayed as well as the way to remove the sensor if the current value is out of ranges (see below).

CURRENT : 340	MIN : 345	MAX : 355	►
---------------	-----------	-----------	---



- For a current number of steps out of ranges, unloosen the 2 screws (see diagram 8) and gently move the sensor

- towards the right if the current value is too low

- towards the left if the current value is too high.

Diag.8



10 steps correspond to around 1 mm. The target number of steps is 350 ± 5

- Tighten the screws (see diagram 8) and rerun a «CARRIAGE» cycle. Check that the current value is correct.

7 - Position adjustment

Needle heigth adjustment

When replacing a needle, it is mandatory to adjust the heigth of the needle in the chambers. Proceed as follows :

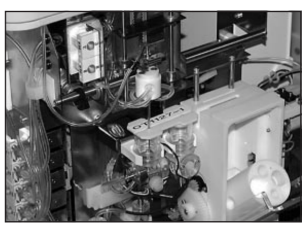
- Remove the WBC/HGB chamber cover.

- Enter the menu 7 - POSITION ADJUST / 1 - NEEDLE / 2 - ADJUSTMENT.

- Manually pull down the needle until it comes into contact with the edge of the RBC chamber (Diag. 9).

- Press a key to continue : the needle moves up to the initial position ; the value is stored.





Diag.10

Diag.9

Needle motion adjustment

The needle position in the WBC chamber can be adjusted as follows :

- Enter the menu 7 POSITION ADJUST / 2 CARRIAGE / 2 ADJUSTMENT.
- Position the tool (P/N : GBC 218 A) over the RBC and WBC chambers (Diag 10).
- Manually lower the needle into the WBC chamber.

- Press a key to continue : the needle moves up, and the carriage comes back to the initial position ; the value is stored.

- Carry out a 3 CHECK cycle : the needle comes down to the WBC chamber.
- Check that the needle is centered in the hole.

If not, enter the 2 - CARRIAGE / 1 - CARRIAGE < > menu ; the display shows the current number of steps carriage motion.

If the needle goes too far on the right, add 1 step to the current value for 0.1mm. If the needle is too much on the left, decrease the current value of 1 step for 0.1mm.

CARRIAGE ? : CURRENT : 893 EXIT : ESC SAVE : ENTER

Carry out a 2 - CARRIAGE / 3 - CHECK cycle again to control the needle position.

8 - Bubbling adjustment

An overflow protection tank is installed on the drain circuit of each chamber. This one prevents from polluted liquid overflow during bubbling phasis (Diag .11).



Diag.11

Two bubbling phasis are adjustable :

- "BUBBLING 1" is the first dilution (WBC/HGB chamber) bubbling value.
- "BUBBLING 2" is the second dilution (WBC/HGB chamber + LYSE) value and RBC chamber bubbling value.

Both values correspond to a number of steps carried out by the waste/vacuum syringe. Default values are BUBBLING 1 : 175 BUBBLING 2 : 120

CAUTION

These values are factory adjusted (and may be different from the default values shown above) and should be modified only when hematologic erroneous results are given by the instument : If values are too important, liquid overflows can occur or if bubbling is too low homogeneity of the dilution can be decreased. Ranges : 150 < BUBBLING 1 < 200 80 < BUBBLING 2 < 140

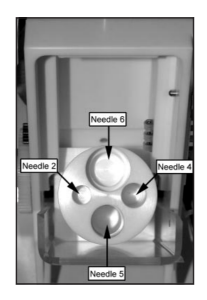
To modify the bubbling values, enter the menu :

2 - ADJUSTMENTS / 8 - BUBBLING ADJUST. / BUBBLING 1 < > 2 - ADJUSTMENTS / 8 - BUBBLING ADJUST. / BUBBLING 2 < > and type in new step value.

Carry out a 3 - CHECK to control the adjustment.

TECHNICIAN FUNCTION

8 - Needle sampling position



- Move the tube holder to one of the four sampling positions.

- Move the cursor to the function 9 - SAMPLING POSITION / 1 - NEEDLE ADJUSTMENT.

- Close the sample holder door. The sampler assy moves to the upper position (except in the *Needle 6* position).

- Manually push the sampling needle to the bottom of the required sampling position and

press \bigoplus_{Enter} : The **MICROS 60** *CT* adjust automatically the needle depth whatever the tube holder position can be. The sampler assy comes back to the initial position and the sampler door is opened.

Diag.13

WARNING !



OEM instruments equipped with specific sample tube holders (see RAS 198 A procedure) must be used with the specific tubes or blood controls they were intended for.

- Turn the tube holder to another needle position and carries out the same procedure to adjust the needle depth.

It is possible to enter directly the required number of steps for each sampling position. Proceed as following :

Enter the sub menu that corresponds to the number of the needle : the current number of steps is displayed. Enter the new value.

Increase the number of steps to move the needle deeper or decrease the value to raise the needle.

Confirm the new value with ______. The minimum and maximum step values are as follow :

NEEDLE	NUMBER OF STEPS		
	MINI.	DEFAULT	MAXI.
1	1	788	1100
2	1	661	1100
3	1	612	1100
4	1	948	1100
5	1	1003	1100
6	1	845	1100

* Check

The CHECK function allows to check the piercing operation on each sampling position. From the SAMPLING POSITION menu, move the cursor to the function <9> CHECK and press ENTER.

Select the required position on the tube holder and close the sample door. The piercing cycle is carried out and the number of steps for this position is displayed :



III - Temperature adjustment

When entering the «temperature adjustment» menu the following sub menus are displayed :

1 - Uncal. temperature

When pressing the \sum_{Enter} key the sensor temperature value **uncalibrated** is displayed. This value should be close to the diluent temperature.

2 - Calibrate :

The temperature must be calibrated according to the diluent temperature :

- Plunge a thermometer directly into the diluent container and leave it for a while until stabilization.

For a pack equipped instrument, the thermometer must be plunged in the WBC/HGB chamber and the temperature must be note as soon as possible.

- Run 2 diluent primes («SERVICE» menu, «PRIME» sub menu, «DILUENT» selection).

- Enter the «CALIBRATE» menu. Note the temperature of the diluent and type in the value (if it is different from the previous on the instrument).

- Press of to save the new value.

3 - Temperature :

1 - Visible : Press to validate the temperature display. The value displayed is the calibrated temperature.

2 - Invisible : Press to cancel the temperature display. (Temperature invisible by default)

IV - Run mode

The instrument must be configurated in the «USER» mode (configurated by default in the «USER « mode), the «CONTROL» mode intends for a factory use.

V - Reagent pack

This function is used to update the instrument from a bottle mode to a pack mode.

VI - Serial

Displays the instrument serial number.

VII - Cycle #

Displays the :

Startup number since the first use of the instrument. Stand by number since the first use of the instrument. CBC number since the first use of the instrument.

IMPORTANT

The startup, stand by and CBC numbers are adjustable in this menu but it is mandatory to keep the initial values (useful for maintenance schedules).

VIII - Burn-in

CAUTION

This function which allows the burn-in of the instrument is intended for a factory use.

LX300 PRINTER SETUP

· CONCERNS

1 - Configuration

- 2 Control panel
- 3 Control LEDS and keys
- 4 Printer description

· REQUIRED TOOLS

- None

S

· REQUIRED PRODUCTS

- None

· INTERVENTION TIME

- 15 minutes

• FREQUENCY

- On request

· SPECIFIC KIT OR CONSUMABLES

- None



RAS 177 A Ind.A

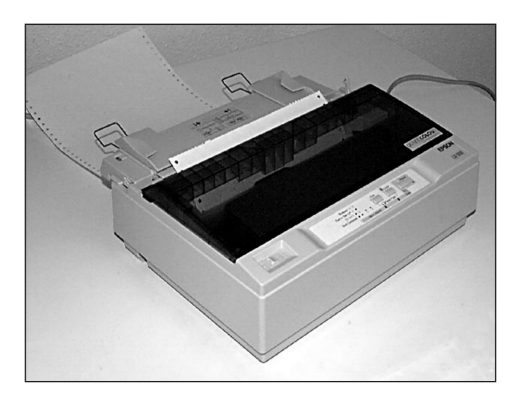
LX300 PRINTER SETUP

• PROCEDURE

0

a

On the MICROS 60, it is necessary to select the printer format RESERVED 1 of the "PRINTER" menu (function 4) accessible through the "OPTIONS" menu (function 5 of the main menu) then "RESULTS" (function 1).



Diag.1

LX300 PRINTER SETUP

1 - Printer configuration :

The printer configuration is printed out when pressing the key when the printer is switched ON. The configuration used for the MICROS 60 is the factory configuration :

******	******
Character spacing	s ₁ 2 ^{>} cpi
Shape of zero	O
Skip-over-perforation	Off
Character table	PC 437
Auto line feed	Off
Page length	12 inches
Auto tear off	Off
Tractor	Single
Interface	Auto selection (10 sec.)
Bit rate	9600 bps
Parity	None
Data length	8 bit
ETX/ACK	Off
Software	ESC/P
Auto CR	Off

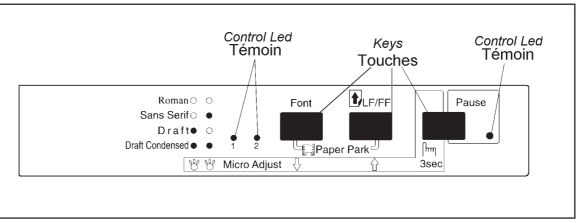
Diag.2

Each parameter can be modified by the corresponding parameter chart. Each chart is accessible using the keys <PAUSE>, and <LF/FF> according to the control LED combinaisons :

Lights				,,,,,,,,,,,,,,,,,,,,,,,,,,,,,,,,,,,,,,
1	2	PAUSE	Setting	Go to submenu
BLINKS	OFF	OFF ;	Character spacing	¦ Table C
BLINKS	ON	OFF ;	Shape of zero	; Table D
OFF	BLINKS	OFF ;	Skip-over-perforation	Table E
ON	BLINKS	OFF ;	Character table	; Table F
BLINKS	; BLINKS	OFF ;	Auto line feed	; Table G
BLINKS	; OFF	ON ;	Page length	; Table H
BLINKS) ON	ON ;	Auto tear off	; Table 1
OFF	BLINKS	ON ¦	Tractor	; Table J
ON	BLINKS	ON ;	Interface	; Table K
BLINKS	BLINKS	ON ;	Bit rate	¦ Table L
OFF	; OFF	BLINKS ;	Farity	¦ Table M
BLINKS	OFF	BLINKS ;	Data length	; Table N
ON	OFF	BLINKS ;	ETX/ACK	; Table O
ON	I ON	BLINKS ;	Software	; Table P
BLINKS	ON ·	BLINKS ;	Auto CR	¦ Table Q



2 - Control pannel : (See Diag.4)



Diag.4

The control pannel keys allow the user to set up the main functions of the printer : paper advance, paper ejection, and font selection. Control LEDS indicate the printer status.

3 - Control LEDS and keys :

LED PAUSE : The orange LED PAUSE lights when the printer stops printing. During each power ON, this LED blinks for few seconds and 4 audible beeps occur. When the printer runs out of paper, the

LED blinks and 3 audible beeps occur. This LED lights also when the paper is in its tear off position.

When a problem occurs, this LED lights ON and 5 audible beeps occur.

LEDS FONT 1 and **FONT 2**: These 2 green LEDS indicate the selected font. Refer to the printer user's manual to select the font.

Key FONT : During normal operation, the FONT key allows the font selection. For each pressure on this key, the selection is modified. Refer to the printer user's manual to select the font. When this key is pressed during the printer power ON, the printer setup menu is entered.

Key LF/FF : During normal operation, a quick pressure on this key allows a ligne feed of the paper. Keep the pressure on this key to feed a whole page. This key can be used to load or eject the paper.

When this key is pressed during the printer power ON, the printing test starts.

Key PAUSE : When this key is pressed during the printing, the printout stops. Press again on this key to restart the printout.

PAPER PARK : If Z folded paper is used, the paper can be driven to its parking position when pressing simultaneously on the keys LF/FF and FONT.

MICRO ADJUST : This function allows to adjust the loading paper position. See the user's manual for details.

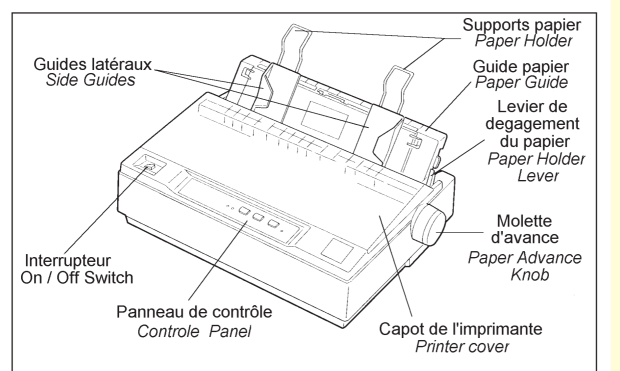


Q

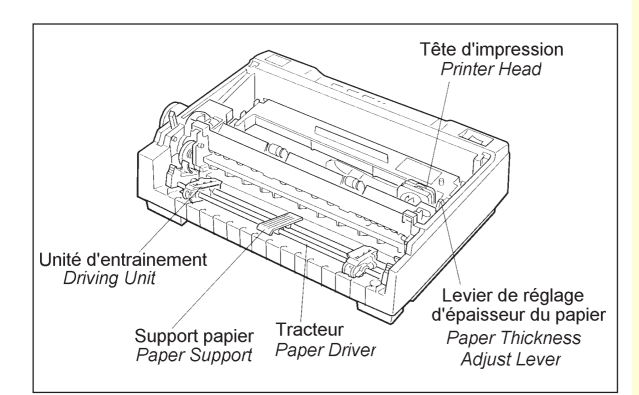


LX300 PRINTER SETUP

4 - Printer description :



Diag.5



Diag.6

T

LIQUID SYRINGES

CONCERNS

Lyse dispenser O ring replacement Diluent dispenser O ring replacement Sampling needle dispenser O ring replacement Lubrication of the liquid syringes

· REQUIRED TOOLS

- Hexagonal keys
- Dynamometric screw driver : A302 : MAG 019 A A301 : MAG 020 A

· REQUIRED PRODUCTS

- Silicone grease : LAM 004 A
- Soft paper
- Grease for mechanical assemblies : XEA 381 AS

INTERVENTION TIME

- 30 min

 FREQUENCY - Once a year or on request

· SPECIFIC KIT OR CONSUMABLES

- O ring kit : XEA 328 AS

RAS 178 A Ind.A

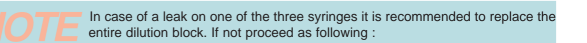


16/06/98



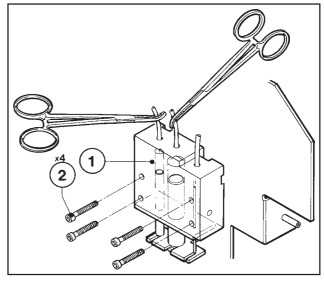
PROCEDURE

0



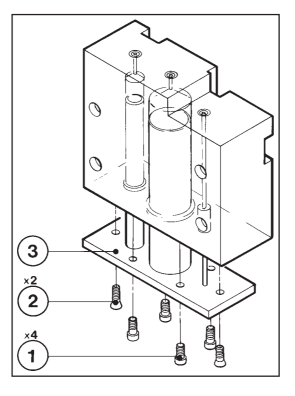
- Pull the piston assy in the upper position and clamp the diluent and lyse tubing as described on the diagram 1.

- Disconnect the diluent/lyse/ sampling tubings from the 3 syringes and the tube on the sampling syringe side.





- Unscrew the 4 fixation screws 2 , and remove the dilution block 1 (Diagram 1).



Diag.2

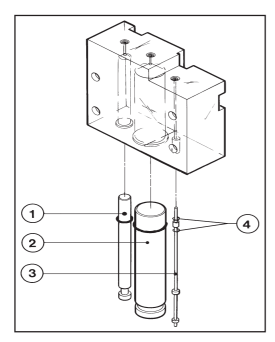
- Unscrew the 6 screws 1 and 2 in

order to remove the body cover 3 (Diagram 2).

- Pull out the pistons 1, 2 and 3 from the body with their respective o ring still around (Diagram 3).

- Replace the lyse and diluent O rings by new ones. Check the cleanliness of the piston and of the syringe bodies. If necessary clean with a soft paper.

LIQUID SYRINGES



Diag.3

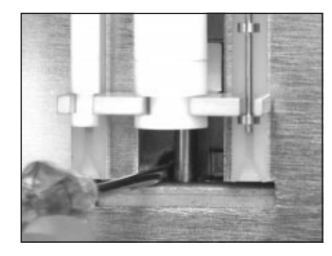
- Spread a little amount of silicone grease between two fingers and apply a very thin film of grease on the 2 new sampling syringe O rings. Replace the

old ones (4).

- Reinstall the dilution block assy in the reverse order.

Use the A302 dynamometric screw driver to tighten the screws 1 to **700mN.m** (Diagram 2).

Use the A301 dynamometric screw driver to tighten the screws 2 to **400mN.m** (Diagram 2).



Lubrication

- Disconnect the diluent and waste inputs located at the rear of the instrument (or remove the reagent pack)

- Move the liquid syringe by hand in order to have an access to the motor gearings.

- Spread a little amount of grease on the gearings and on the piston axis (Diag.4).

- Move by hand the syringe assembly to spread the grease on all parts of the gearings and piston axis.

- Re-install the instrument cover, reconnect the waste and diluent tubes, reconnect the power cable.

- Switch the instrument on and run several priming cycles.

CAUTION

Place some absorbant paper at the instrument rear connections (diluent and waste) as some liquids may come out when the syringe is pushed.

VACUUM/WASTE SYRINGE MAINTENANCE

CONCERNS

- O ring replacement



· REQUIRED TOOLS

· REQUIRED PRODUCTS

- Silicone grease : LAM 004 A

INTERVENTION TIME

- Hexagonal keys
- Dynamometric screw driver A302 : MAG 019 A

- Grease for mechanical assemblies : XEA 381 AS.

- 20 min

FREQUENCY

- See maintenance chart table

· SPECIFIC KIT OR CONSUMABLES

- O ring kit : XEA 328 AS

RAS 179 A Ind.A

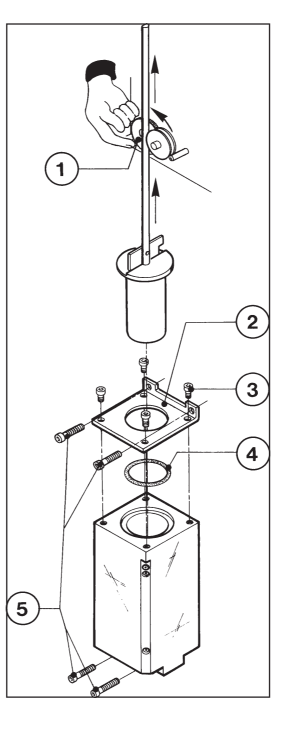
VACUUM/WASTE SYRINGE MAINTENANCE

• PROCEDURE

- Manually pull the syringe piston up in order to freed it from the syringe body.

The syringe has to be linked to the atmosphere, pressing the valve 2, to pull the piston out from the syringe body. Turn the cylindrical gearing 1 by hand to help the raising of the piston (See Diagram 1).

いろ



Diag.1

VACUUM/WASTE SYRINGE MAINTENANCE

- Unscrew the fixation screws (5) in order to remove the syringe body.
- Unscrew the O ring tightening screws (3) and remove the O ring (4).
- Spread a little amount of silicone grease between two fingers and apply a very thin film of grease on a new O ring.
- Reinstall in the reverse order. Apply the following torque to the screws (5): 700 mN.m.



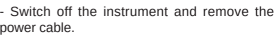
Push the piston back inside the syringe body before tightening the o ring fixation screws (3). Use the dynamometric screw driver to tighten the screws (3) to 400 mN.m

- Check the watertightness of the syringe running a «CHECK PRESSURE» cycle (see procedure : RAS 172 A for OT or RAS 176 A for CT).

- Run cycles and check for correct operations.
- Lubrication



Diag.2



- Disconnect the diluent and waste inputs located at the rear of the instrument.

- Using a small and flat screwdriver, spread a little amount of grease on the gearings of the air syringe reductor plate (Diag.2).



- Spread a little amount of grease on the coggs of the piston axis (Diag.3).

Diag.3

VACUUM/WASTE SYRINGE MAINTENANCE





- Move by hand the piston axis up and down in order to spread the grease all around the gearings and along the axis (Diag.4).

Diag.4

Z

Q

INSTRUMENT LANGUAGE

01/04/98

• CONCERNS Hématologie Changing the instrument language · REQUIRED TOOLS Pair of pliers · REQUIRED PRODUCTS None INTERVENTION TIME R 5 minutes • FREQUENCY On request only · SPECIFIC KIT OR CONSUMABLES

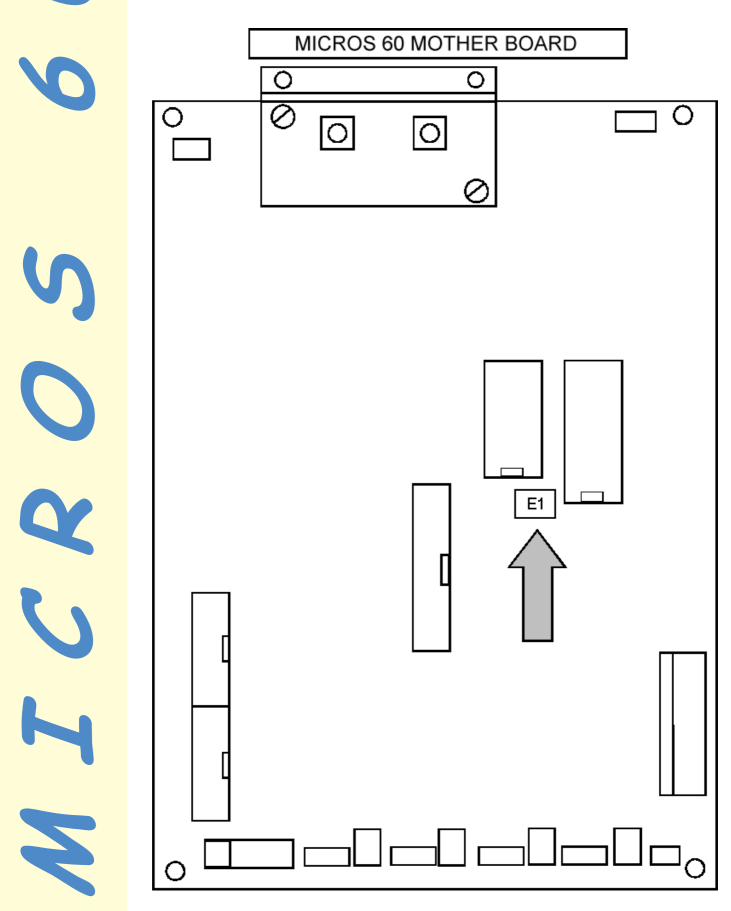
None

5

RAS 180 A Ind.A

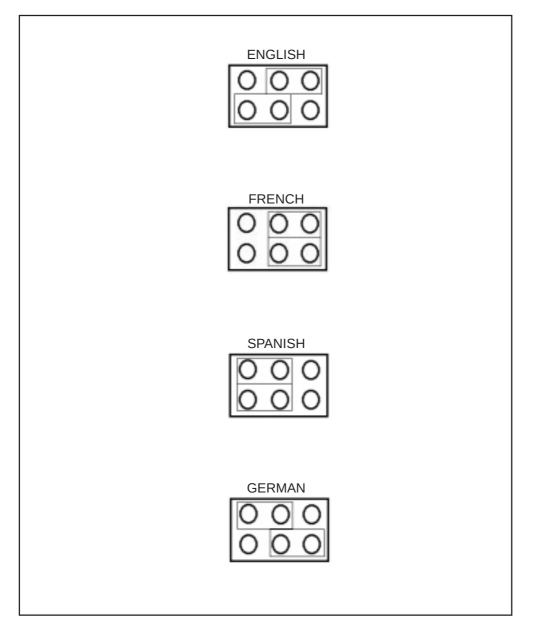
INSTRUMENT LANGUAGE

• PROCEDURE





- Replug the jumpers on E1 according to the wished language :



JUMPER CONFIGURATION ON E1

- Switch on the instrument.

REAGENT PACK

• CONCERNS

- Replacement of the waste connector O ring
- Replacement of the diluent/clean/lyse O rings

· REQUIRED TOOLS

- Pair of pliers
- Torx keys

· REQUIRED PRODUCTS

- None

INTERVENTION TIME

- 15 min

• FREQUENCY

- O ring replacements : 1/year

· SPECIFIC KIT OR CONSUMABLES

- O rings : FAA 036 A

5





RAS 181 A Ind.A

REAGENT PACK

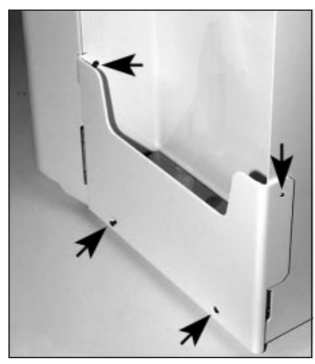
• PROCEDURE

- Disconnect the pack if this one is still connected.

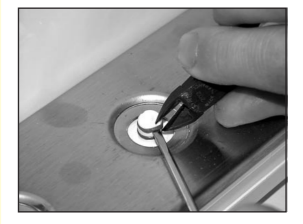


C

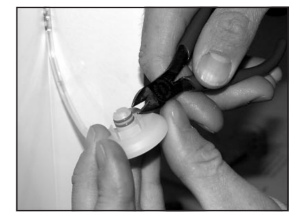
- Dismantle the reagent pack front panel unscrewing the four torx screws as shown on the diagram 1.













- Use a pair of pliers to cut the O rings of the reagent connectors (diag 2) .
- Replace the O rings by new ones.
- Replace as well the waste connector O ring (diag 3).

Hématologie

20/04/98

• CONCERNS Barcode reader installation & configuration · REQUIRED TOOLS None. · REQUIRED PRODUCTS None. INTERVENTION TIME P 10 minutes. • FREQUENCY On request. · SPECIFIC KIT OR CONSUMABLES Installation kit : XBA 379 AS





PROCEDURE

Installation :

- Switch off the instrument.
- Open the instrument cover.
- Install the jumper (included in the XBA 379 AS kit) on E10 as shown on the diagram 3.

WARNING !

Once the jumper installed on E10, the instrument data ouput receives 5 volts to supply the barcode reader. This voltage should cause damages on computer connection if this one is directly connected on the MICROS 60 data ouput. It is then mandatory to connect the computer connection only on the cable (DAC 023 AS shown on the diagram 2) intended for it.

- Connect the BARCODE reader in the DIN plug of the RS adaptor wiring (diag 1 and 2).



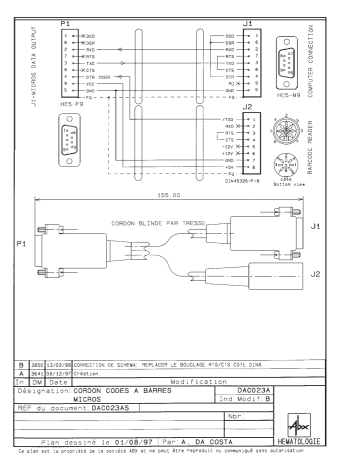
Diag.1

Setup :

- Switch on the instrument and within the 15 first seconds, read from the top to the bottom the 3 barcode labels located on the top left of the page 4 of this procedure. The audible beep occurs after each reading.

- After the 3 labels (the audible signal beeps 5 times) read all the labels from top to bottom and from left to right.

- When the last label is read, the signal beeps 5 times in order to indicate the end of the setup. Check on the test labels located on the last page that the reading is correct.



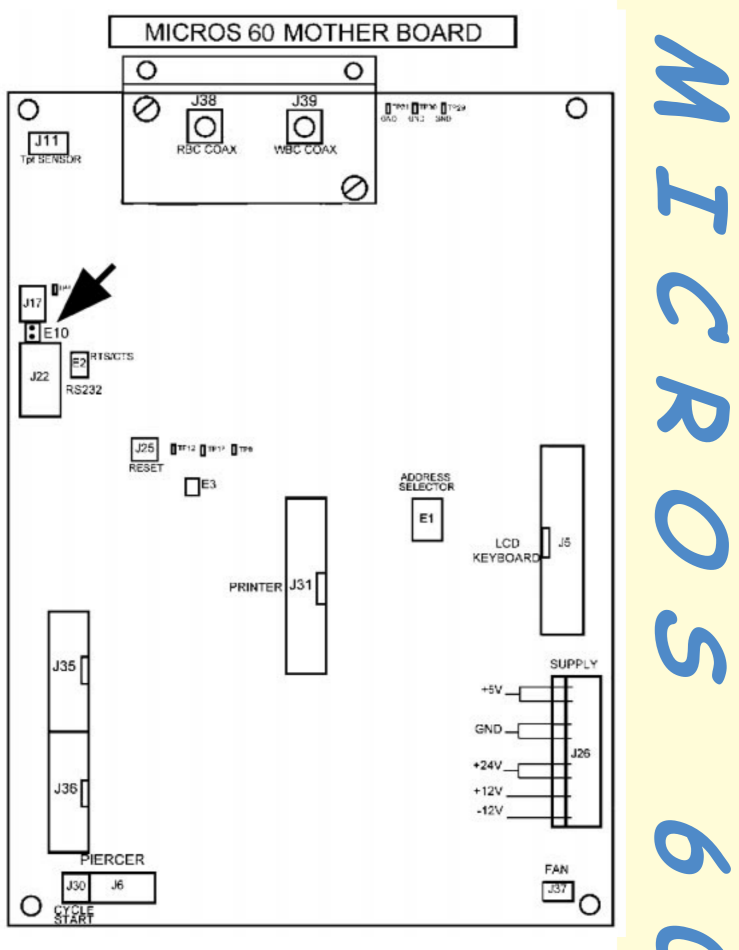


- In case of difficulties to read 2 consecutive 0, move away the barcode

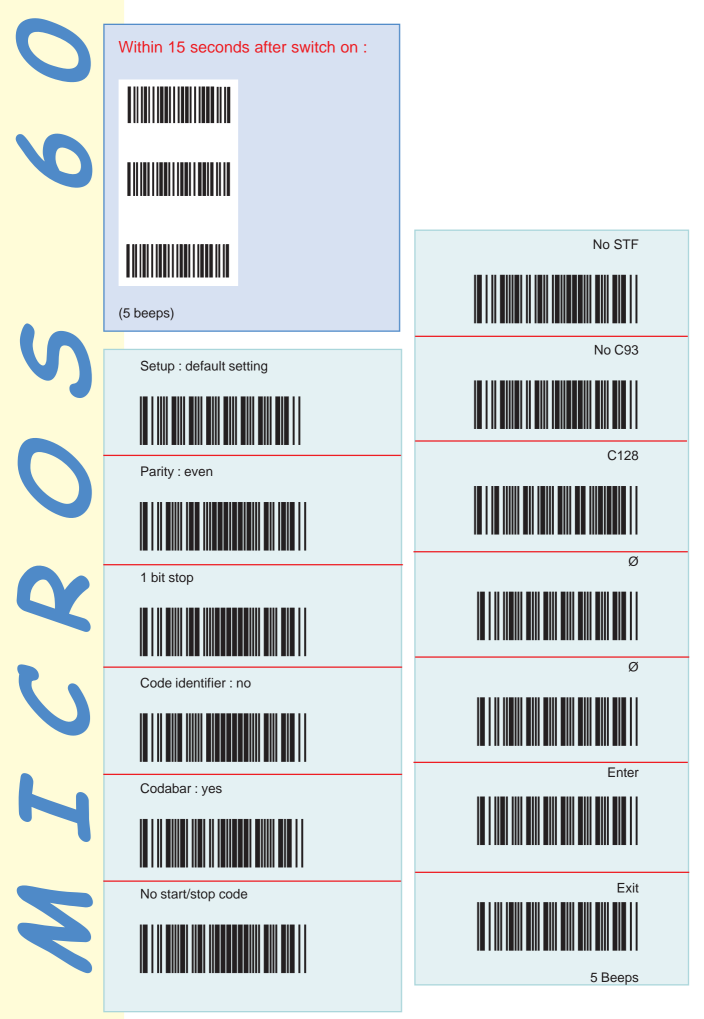
- reader from the page after the first reading and then read the second 0.
- To carry out the *read test after setup* the instrument must be configurated in a US mode in order to obtain the barcode identification in the ID field (menu 5 - SETUP / 3 - SPECIAL / 7- ID MODE).
- The barcode reader can be configurated according to the type of barcode label in use (from the menu 5 - SETUP / 6 - BARCODE) and allows to enable the checksum or not.

The barcode setup can be printed out by the function 5 - SETUP / 3 - SPE-CIAL / 5 - PRINT CONFIG.

RAS 182 A Ind A









READ TEST AFTER SETUP



• CONCERNS

Step by step control of the hydraulic cycle.



· REQUIRED TOOLS

None

· REQUIRED PRODUCTS

Blood samples

INTERVENTION TIME

15 minutes

• FREQUENCY

On request

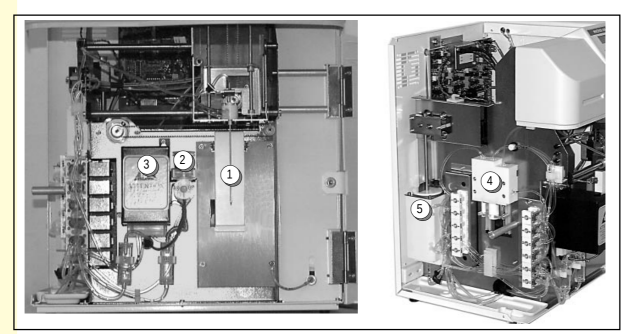
N H C N H C

· SPECIFIC KIT OR CONSUMABLES

None

RAS 187 A Ind.A

PROCEDURE

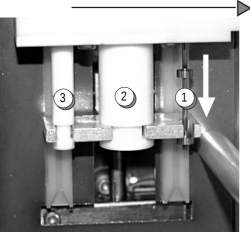


Diag.1

- 1 Cycle start condition
- Needle 1) in the sampling position (diag 1).
- RBC chamber 2 filled with 2.5ml of diluent.
- WBC/HGB chamber 3 filled with 2.5ml of diluent.
- Liquid syringes 4 in standby position.
- Vacuum/waste syringe 5 in the lower position.

2 - Sampling

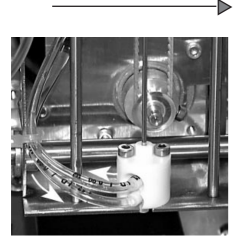
• Aspiration of 10µl of blood sample



Diag.2

<u>Causes</u> The liquid syringes assembly moves down (diag 2) and pulls down the micro sampling syringe 1.

3 - Outer sampling needle rinse



Diag.3

<u>Causes</u>

- The sampling needle moves up.

- The liquid syringes 2 (diag 2) send diluent for rinse

through the rinsing block

- Polluted diluent is aspirated (from the lower to the upper tube) by means of waste/Vacuum syringe raise.



Þ

 \triangleright

Þ

Diag.4

4 - WBC/HGB chamber rinse & HGB blank measure

- Sampling carriage transfer over the WBC/HGB chamber.
- Counting head rinse
- WBC/HGB chamber drain -
- Needle motion downward of a few steps (Diag 5)
- Diluent is delivered from the rinsing block

<u>Causes</u> Diluent is delivered by means of the liquid syringes raise

Aspiration by means of the Vacuum/waste syringe raise (diag 4)

The liquid syringes move up and a flow of diluent is delivered to the chamber via the outer needle.



Diag.5

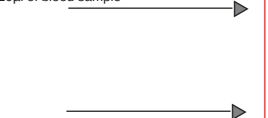
Page 3/6

- WBC/HGB chamber drain (second time)
- Diluent is delivered from the outer needle (second time)
- HGB blank measure (beep triggered)
- RBC and WBC/HGB chamber drains.

5 - Dilutions

Bubbling

- Sampling needle moves down to the WBC/HGB chamber
- Injection of 1.7ml of diluent into the WBC/HGB chamber
- + Injection of 10µl of blood sample



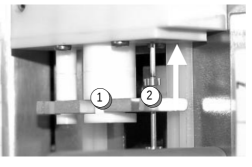
Causes

Raise of the liquide syringes (diag 6) :

- delivers 0.5ml of diluent from the outer sampling needle

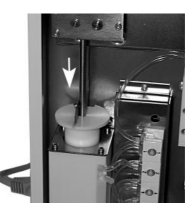
- injects 1.2ml of diluent + blood sample from the inner sampling needle

Bubbling by means of vacuum/ waste syringe downward motion (Diag 7).



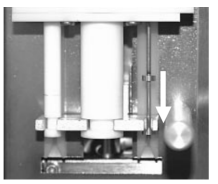
Diag.6

- Sampling needle moves up
- Outer needle short aspiration (dries the needle)
- Sampling needle moves back in the chamber
- Aspiration of 28.3µl of diluted blood (dilution 1/170)



Diag.7

<u>Causes</u> The liquid syringes move down and pull down the micro sampling syringe (diag. 8)



Diag.8

- Sampling needle moves up
- Injection of 0.4 ml of diluent into the WBC/HGB chamber

<u>Causes</u> Raise of the liquide syringes (diag.6) : - delivers 0.4ml of diluent from the outer sampling needle

A L C R O S

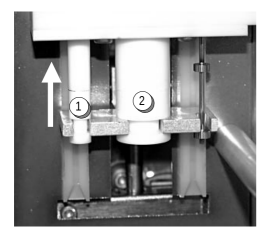
- Outer needle aspiration
- Carriage motion over the RBC chamber
- Sampling needle moves down into the RBC chamber
- Lyse injection into the WBC/HGB chamber

<u>Causes</u>

The liquid syringes raise (diag 8)

and the syringe ① delivers 0.52ml of lyse via the WBC chamber bottom

FINAL DILUTION in the WBC CHAMBER : 1.7ml + 0.4ml diluent + 0.52ml of lyse = 1/260





Diag.8

• Injection of 2.5 ml of diluent into the RBC chamber

+ Injection of 28.3 µl of diluted blood

<u>Causes</u>

Diag.9

The liquid syringes raise (diag 8) : - delivers 0.5ml diluent via the outer sampling needle

- injects diluted blood from the inner needle + 2ml of diluent

FINAL DILUTION in the RBC CHAMBER : 28.3µl of diluted blood at 1/170 + 2.5ml diluent

_1	28.33	1
170	2500	15000

- Bubbling (diag 10)
- Sampling needle moves up
- Carriage motion over the WBC/HGB chamber
- Counting head rinse



Diag.10

6 - Counts

- First counts (beep triggered)
- · Counting head rinse
- Second counts (beep triggered)

A third count (C3) is carried out if the difference between first (C1) and second count (C2) is not within acceptable limits :

<u>- WBC</u> :

if C1 or C2 > 3000 C3 is carried out if difference between C1 and C2 > 7%

if Max C1 or C2 \leq 3000 C3 is carried out if difference between C1 and C2 > 9%

<u>- RBC</u> :

if C1 or C2 > 16000 C3 is carried out if difference between C1 and C2 > 5%

if Max C1 or C2 \leq 16000 C3 is carried out if difference between C1 and C2 > 8%

<u>- PLT</u> :

if C1 or C2 > 400 C3 is carried out if difference between C1 and C2 > 15%

if Max C1 or C2 \leq 400 C3 is carried out if difference between C1 and C2 > 20%

- · Counting head rinse
- WBC chamber drain
- Diluent injection into the WBC chamber from the outer needle
- RBC chamber drain
- Carriage motion over the RBC chamber
- Diluent injection into the RBC chamber from the outer needle (diag 11)



Diag.11

- Carriage & needle motions back to the initial positions
- Results display and printed out

Page 6/6

• CONCERNS

Step by step control of the hydraulic cycle.



· REQUIRED TOOLS

None

· REQUIRED PRODUCTS

Blood samples

INTERVENTION TIME

15 minutes

H

• FREQUENCY

On request

· SPECIFIC KIT OR CONSUMABLES

None

RAS 188 A Ind.A

PROCEDURE



Diag.1

- 1 Cycle start condition
- Needle (1) in the sampling position (diag 1 & 2).
- RBC chamber 2 filled with 2.5ml of diluent.
- WBC/HGB chamber (3) filled with 2.5ml of diluent.
- Liquid syringes 4 in standby position.
- Vacuum/waste syringe 5 in the lower position.

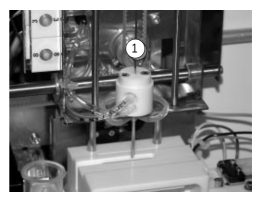
2 - Cap piercing

- Carriage motion over the WBC chamber.
- WBC/HGB chamber drain

• Diluent injection into the WBC/HGB chamber through the channels (A) and (B) (see diag.4)

- HGB blank measure (beep triggered)
- Carriage return over the piercing device
- Atmosphere is provided inside the tube

Cap piercing



Diag.2

<u>Causes</u> Aspiration by means of the Vacuum/waste syringe raise.

Diluent is delivered by means of the liquid syringes raise

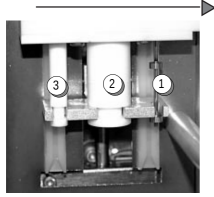
<u>Causes</u> Liquid valve #3 is activated

Sampling holder rises in the upper position. The needle pierces the tube cap

RAS 188 A Ind.A

3 - Sampling

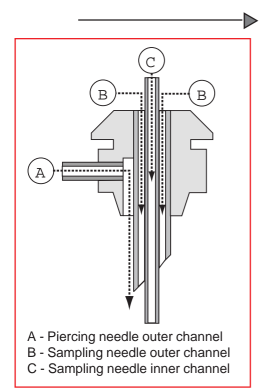
- Sampling needle (diag. 2) moves down to the lower position (inside the tube)
- Aspiration of 10µl of blood sample



<u>Causes</u> The liquid syringes assembly moves down (diag 3) and pulls down the micro sampling syringe 1).

Diag.3

- The sampling needle comes back in the upper position
- RBC chamber drain
- · Counting head rinse
- Carriage motion over the WBC/HGB chamber
- Sampling holder door opens
- WBC/HGB chamber drain
- 4 needle rinses



<u>Causes</u>

- The liquid syringe (2) (diag 3) sends diluent for rinse through the outer piercing needle (A) (diag 4) and inner piercing needle (B). The polluted diluent is sent to the WBC/HGB chamber.



Diag.5



The piercing needle inner rinse is equivalent to the sampling needle outer rinse.

- WBC/HGB chamber drain
- Second needle rinses
- WBC/HGB chamber drain
- The sampling needle moves down into the WBC/HGB chamber





Bubbling

Injection of 1.7ml of diluent into the WBC/HGB chamber
 + Injection of 10µl of blood sample

<u>Causes</u>

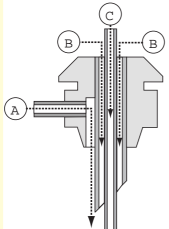
 \triangleright

Raise of the liquide syringes (diag 7) :

- delivers 0.5ml of diluent from the outer sampling needle (channel B diag.6)

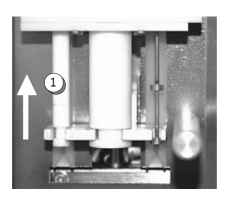
- injects 1.2ml of diluent + blood sample from the inner sampling needle (channel C)

Bubbling by means of vacuum/ waste syringe downward motion (Diag 8).



· Sampling needle moves up

• Outer needle short aspiration (dries the needle)





Diag.6

Diag.7

- Sampling needle moves back in the chamber
- Aspiration of 28.3µl of diluted blood (dilution at 1/170)

• Sampling needle moves up

- Injection of 0.4 ml of diluent into the WBC/HGB chamber
- Carriage motion over the RBC chamber

FINAL DILUTION in the WBC CHAMBER :

- Sampling needle moves down to the RBC chamber
- Lyse injection into the WBC chamber + Bubbling

1.7ml + 0.4ml diluent + 0.52ml of lyse = 1/260

<u>Causes</u> Raise of the liquide syringes (diag.7) : - delivers 0.4ml of diluent from the outer sampling needle (channel B)

Diag.8

<u>Causes</u> The liquid syringes raise (diag.7) and the syringe 1 delivers 0.52ml of lyse via the WBC chamber bottom

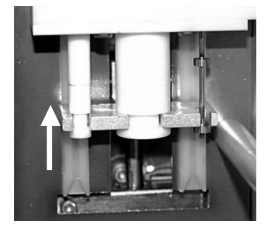
 \triangleright

• Injection of 2.5 ml of diluent into the RBC chamber + Injection 28.3 µl of diluted blood

<u>Causes</u>

·D>

The liquid syringes raise (diag.9) : - delivers 0.5ml diluent via the outer sampling needle (channel B diag. 6) - injects diluted blood from the inner needle + 2ml of diluent (channel C)



Diag.9



Diag.10

FINAL DILUTION in the RBC CHAMBER : 28.3µl of diluted blood at 1/170 + 2.5ml diluent

$$\frac{1}{170} * \frac{28.33}{2500} = \frac{1}{15000}$$

- Bubbling (diag 11)
- Sampling needle moves up
- Carriage motion over the WBC/HGB chamber
- Counting head rinse



Diag.11

6 - Counts

- First counts (beep triggered)
- · Counting head rinse
- Second counts (beep triggered)

A third count (C3) is carried out if the difference between first (C1) and second count (C2) is not within acceptable limits :

<u>- WBC</u> :

if C1 or C2 > 3000 C3 is carried out if difference between C1 and C2 > 7%

if Max C1 or C2 \leq 3000 C3 is carried out if difference between C1 and C2 > 9%

<u>- RBC</u> :

if C1 or C2 > 16000 C3 is carried out if difference between C1 and C2 > 5%

if Max C1 or C2 \leq 16000 C3 is carried out if difference between C1 and C2 > 8%

<u>- PLT</u> :

if C1 or C2 > 400 C3 is carried out if difference between C1 and C2 > 15%

if Max C1 or C2 \leq 400 C3 is carried out if difference between C1 and C2 > 20%

- · Counting head rinse
- WBC chamber drain
- Diluent injection into the WBC chamber from the outer sampling needle (channel B)
- RBC chamber drain
- Carriage motion over the RBC chamber
- Diluent injection into the RBC chamber from the outer sampling needle (channel B)
- Carriage & needle motions back to the initial positions
- · Results display and printed out

· CONCERNS

0

Instrument maintenance step by step



· REQUIRED TOOLS

Hexagonal keys Dynamometric screw driver A302, A301, A300 Clamps Scalpel Cutting pliers Pair of scisors Voltmeter Flat screw driver Barflex Thermometer Torx keys

· REQUIRED PRODUCTS

Empty sample tubes Silicone grease Soft tissue Liquid soap Distilled water Micropipette tip Flat piece of stiff plastic Latex WBC, RBC Felt pen syringe 5ml Fresh blood samples Calibrator Commercial control

INTERVENTION TIME

2 h 30

• FREQUENCY

The yearly maintenance frequencies vary with the instrument output. According to the cycle numbers, 3 categories are created : < 6000 cycles/year -> 1 overall maintenances/year 6000 to 15000 cycles/year -> 2 overall maintenances/year > 15000 cycles/year -> 3 overall maintenances/year

· SPECIFIC KIT OR CONSUMABLES

Spare parts kit : XEA 458 AS Needle position tool : GBC 218 A

RAS 191 A Ind.B

	A - INSTRUMENT CHECKUP	Procedures
0	 1 - Reagent check ✓ Type of reagent used ✓ Expiration dates ✓ Levels ✓ Pack : Number of cycles left 	
	 2 - Operation check ✓ Blank cycle control ✓ QC control ✓ Fresh blood sample run ✓ Calibration coefficient checkup ✓ Leak control and general cleanliness of the instrument 	
	B - CLEANING AND MAINTENANCE	
J	 1 - Chamber maintenance ✓ RBC & WBC/HGB chamber cleaning ✓ Aperture check ✓ Coaxial O ring replacement ✓ Aperture O ring replacement ✓ Aperture tightening torque : 100mN.m 	RAS 169 A
2	 2 - Liquid syringes ✓ Cleaning ✓ O ring replacement (Lyse, diluent, Micro syringe) Torque values : 	RAS 178 A
Y	 torx screws : 400mN.m Hexagonal screws : 700mN.m Diag.1 ✓ Lubrication of the gearings and piston axis 	
	 3 - Sampling needle ✓ Cleaning ✓ Sampling needle O ring replacement CT : O ring holder tightening torque : 700mN.m OT : O ring holder tightening torque : 100mN.m 	RAS 167 A (CT) RAS 168 A (OT)
	 ✓ Free pulley cleaning and lubrication Pulley tightening torque : 400 mN.m ✓ Piercing needle check/cleaning (CT) 	
	 4 - Air syringe ✓ Cleaning ✓ O ring replacement O ring tightening torque : 400mN.m Syringe holding torque : 700mN.m ✓ Lubrication of the gearings and piston axis 	RAS 179 A
2	 5 - Piercing block maintenance ✓ Check and clean 	RAS 198 A (CT)
	6 - Reagent pack (Option) ✓ Replacement of the connector O rings	RAS 181 A

C - MECHANICAL OPERATION CHECK 1 - Liquid syringes ✓ Operation check Menu 4 - SERVICE / 5 - MECHANIC / 4 - LIQ. SYRINGES	Procedures	2
 2 - Air syringe ✓ Operation check Menu 4 - SERVICE / 5 - MECHANIC / 5 - PRESSURE SYR. 	RAS 173 A	
 3 - Liquid valves ✓ Operation check Menu 4 - SERVICE / 5 - MECHANIC / 6 - VALVES 		H
 4 - Sampling needle and carriage ✓ Operation check Menu 4 - SERVICE / 5 - MECHANIC / 2 - NEEDLE U/D Menu 4 - SERVICE / 5 - MECHANIC / 3 - CARRIAGE L/R 		0
D - MECHANICAL ADJUSTMENTS		
1 - Needle heigth ✓ Menu 7 - TECHNICIAN / 2 - ADJUSTMENTS / 7 - POSITION ADJST. / 1 - NEEDLE / 2 - ADJUSTMENT	RAS 172 A (OT) RAS 176 A (CT)	N
\checkmark Needle on the edge of the RBC chamber, press a key.		
 2 - Needle motion ✓ Tool GBC 218 A over the RBC and WBC chamber. ✓ Menu 7 - TECHNICIAN / 2 - ADJUSTMENTS / 7 - POSITION ADJST. / 2 - CARRIAGE / 2 - ADJUSTMENT ✓ Lower the needle in the WBC chamber , press a key. 	RAS 172 A (OT) RAS 176 A (CT)	0
E - HYDRAULIC ADJUSTMENTS		
 ✓ PRIME / ALL REAGENTS ✓ Check for leaks 		1
 1 - Check vacuum ✓ Barflex connected instead of the tube coming from the valve 8. ✓ Menu SERVICE / 7 - TECHNICIAN / 2 - ADJUSTMENTS / 3 - CHECK PRESSURE Value : -200mbar ± 10mbar ✓ Check vacuum drop down during 30 secondes ≤ 2 mbar 	RAS 172 A (OT) RAS 176 A (CT)	S
2 - Bubbling ✓ Menu 7 - TECHNICIAN / 2 - ADJUSTMENTS / 8 - BUBBLING ADJUST. / BUBBLING 1 < > (default value : 175)	RAS 172 A (OT) RAS 176 A (CT)	
✓ Menu 7 - TECHNICIAN / 2 - ADJUSTMENTS / 8 - BUBBLING ADJUST. / BUBBLING 2 < > (default value : 120)		0
 3 - Drain detection ✓ Voltmeter between TP30 and TP6 ✓ Drained sensor : adjust the voltage to 4,5V ± 0,3V by means of R6 ✓ Fill the sensor with diluent : voltage falls down below 1Volt 	RAS 174 A	0

Page 3/7

					-				<u> </u>		
\mathbf{O}										Procedures	
	✓ Me	ermome enu 7 - 1	FECHNICIAN	1/3-TEI	liluent contaiı MPERATURE ure displayed	E / 2 - C		f pack)		RAS 172 A (OT) RAS 176 A (CT)	
	Tempera	IMPORTANT Temperature adjustment must be done 20 minutes at least after the instrument has been switched on.									
	<mark>5 - Photom</mark> ✓ Me			1/2 - AD	JUSTMENTS	5 / 1 - C/	AL PHOTOM	1ETER		RAS 172 A (OT)	
S	-		nel displayed the room ter		creen, adjuste :	ed by m	eans of R97	,		RAS 176 A (CT)	
	ROOM		CHANNEL	-	ROOM		CHANNEI	_			
	TPT (°C)	Mini.	Nominal	Maxi.	TPT (°C)	Mini.	Nominal	Maxi.			
	18	238	243	248	24	233	238	243			
	19	237	242	247	25	232	237	242			
	20	236	241	246	26	231	236	241			
	21	235	240	245	27	230	235	240			
	22	234	239	244	28	229	234	239			
	23	234	239	244	29	228	233	238			
Q	✓ Bla	tex thor ank cycl	oughly mixed e to check th	e cleanlir							
S	 ✓ Menu 7 - TECHNICIAN /2 - ADJUST./4 - WBC ADJUST /1 - WBC ADJUST ✓ Run an analysis on Latex. Adjust with R74 to obtain - LYM = 57 ± 1 - GRA = 180 ± 2 						RAS 172 A (OT) RAS 176 A (CT)				
H	 • RBC ✓ Latex thoroughly mixed ✓ Blank cycle to check the cleanliness ✓ Menu 7 - TECHNICIAN / 2 - ADJUST. / 5 - RBC PLT ADJUST / 1 - RBC PLT AD. ✓ Run an analysis on Latex. ✓ Adjust : 										
		- the F			obtain RBC =					RAS 172 A (OT) RAS 176 A (CT)	
Z											



F - VOLTAGE CHECKS

1 - Power supply

✓ Ground on TP31, TP30 or TP29

TEST POINTS	VOLTAGE
TP 20	-12V <u>+</u> 0,5V
TP 22	24V + 1.5V - 0V
TP 23	5V + 0,3V - 0V
TP21	12V <u>+</u> 0,5V

2 - RBC, PLT, WBC threshold

✓ Ground on TP31

THRESHOLDS	TEST POINTS	VOLTAGE	POTENTIOMETERS
WBC	TP 10	350 mV <u>+</u> 7	R68
RBC	TP13	400 mV <u>+</u> 7	R75
PLT	TP16	180 mV <u>+</u> 3	R82

3 - Stepper motor voltage

✓ Ground on TP31

MOTORS	TEST POINTS	VOLTAGE	POTENTIO.
Air syringe	TP36	2.50 V ± 0.05 V	R195
Liquid syringe	e TP35	2.00 V <u>+</u> 0.05 V	R186
Horizontal carria	age TP37	1.50 V <u>+</u> 0.05 V	R177
Vertical carriag	je TP38	1.00 V <u>+</u> 0.05 V	R168

4 - Aperture voltage

- ✓ Ground on TP30
- ✓ Menu 4 SERVICE / 7 TECHNICIAN / 2 ADJUSTMENTS /
- 2 APERTURE VOLTAGE
- ✓ Test point on TP 19 : check to have 60V +2.8V -1.5V



Procedures

RAS 171 A

RAS 175 A

RAS 175 A

RAS 172 A OT RAS 176 A CT

G - BLOOD SAMPLE RUN

1 - Preliminary

✓ Run a STARTUP cycle

2 - Repeatability

- ✓ Fresh and normal blood sample
- ✓ Run 20 consecutives analyses

CAUTION

For MICROS 60 CT, it is mandatory to remove the cap from the sample tube to prevent from piercing several times the same cap.

 \checkmark Control to have variation coefficients within the following acceptable limits :

PARAMETERS	% CV	TEST LEVEL
- WBC :	< 2,5%	at 10.10 ⁹ /I
- RBC :	< 2%	at 5.10 ¹² /I
- HGB :	< 1,5%	at 15 g/dl
- HCT :	< 2%	at 45 %
- MCV :	< 1%	at 90 fl
- PLT :	< 5%	at 300.10 ⁹ /I
- LYM :	< 5%	at 40%(16/18 param.)
- MON :	< 10%	at 10% (16/18 param.)
- GRA :	< 5%	at 50% (16/18 param.)

<u>CV calculated by means of the below</u> formula :

$$\overline{X} = \frac{\sum Xi}{n}$$
 $SD = \sqrt{\frac{\sum (\overline{X} - Xi)^2}{n-1}}$

 \overline{X} : Mean

Xi : measure value

n : Measure number

SD : Standard deviation

$$CV(\%) = \frac{SD}{\overline{X}} \times 100$$

H - CALIBRATION

✓ Calibration passed :

Check that the calibration coefficients remain within the following ranges :

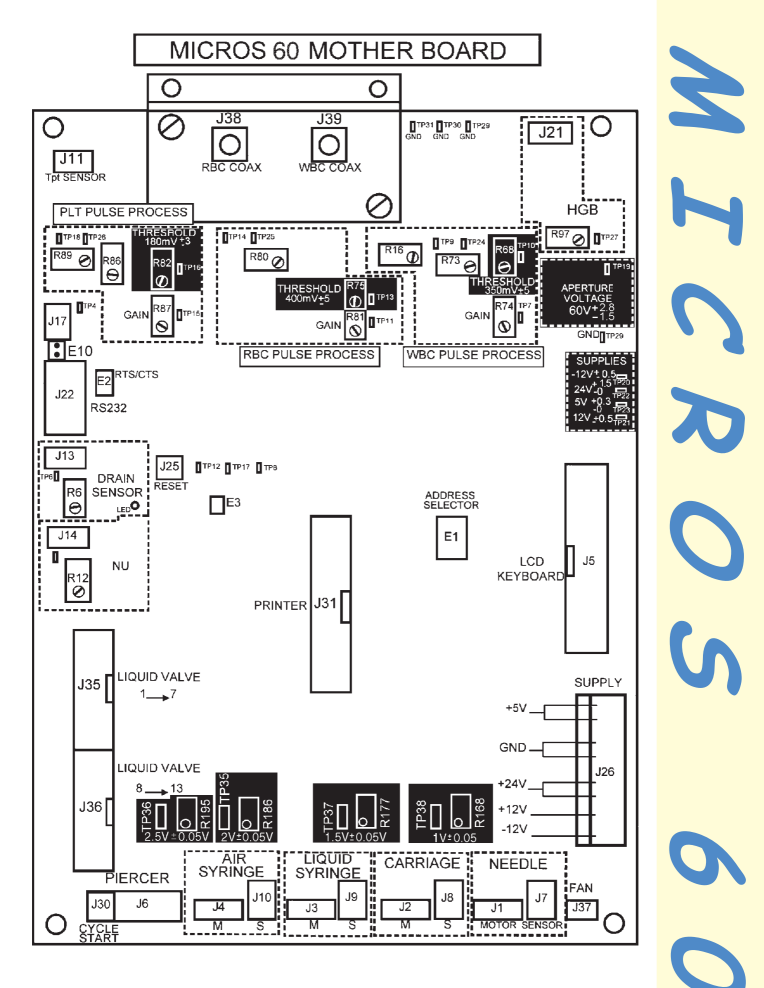
LIMITS	WBC	RBC	HGB	НСТ	PLT	MPV	RDW	PDW
MINIMUM	0.80	0.65	1.10	0.83	0.86	0.75	0.75	0.75
MEAN	1.00	0.81	1.38	1.04	1.07	0.94	1.00	1.00
MAXIMUM	1.20	0.97	1.66	1.25	1.28	1.13	1.25	1.25

I - QUALITY CONTROL

✓ Run a blood control

✓ Option : QC smart card use

OVERALL MAINTENANCE



DECONTAMINATION

· CONCERNS

Instrument decontamination before maintenance operation in the following cases

- Instrument move out of the biologic risks area
- Maintenance intervention on contaminate suspected assemblies

· REQUIRED TOOLS

Hexagonal keys Clamps Flat screw driver Torx keys

· REQUIRED PRODUCTS

Fungicidal, bactericidal, virus killing detergent spray, non corrosive for metals, Non plastic altering. Bleach solution 12°Cl Deionize water Protection gloves Absorbant paper Distilled water

INTERVENTION TIME

1h35min

• FREQUENCY

On request

· SPECIFIC KIT OR CONSUMABLES

Drain and rinse kit XEA 349 AS for Pack equipped instrument

RAS 192 A Ind.A



S

X

DECONTAMINATION

PROCEDURE

WARNING !

- Disposal gloves and white coat must be worn by the operator.
- Local or national regulations must be applied in all the operations .

1 - Preliminary (20min)

- Switch on the instrument
- Run a STARTUP cycle, then a SERVICE / AUTOCLEAN
- Switch off the instrument and remove the supplying cable
- Open the instrument cover

- Spray the bactericidal cleaner on all assemblies that may provide biologic risks and wait for 10 minutes (assemblies in contact with the operator such as instrument cover, tube holder, keyboard, start key, sampling needle neighboured assemblies...

2 - Manual decontamination (20 min)

- Remove the WBC/HGB chamber cover
- Dilute the 12°cl bleach to 1 part of bleach for 4 of deionize water (1/5).
- Instrument environment must be cleaned and decontaminated.
- No sponge, nor cloth must be used. Only absorbant paper, thrown after use, in contamination bins, can be employed. For small or weak assemblies use accurate drier papers.
 All assemblies that is suspected to have contact with biologic product must be disinfected
- with the diluted bleach (the stainless steel must bleached below 30°Celsius).
- Blood stains or salt marks must be cleaned with spray detergent first.
- Concerned assemblies
 - Outer surfaces of the instrument (perpex, covers, LCD, reagent locations....)
 - Keyboards
 - Waste connector plug
 - Liquid valve push
 - Needle neighboured assemblies
 - Tube holder assy.
 - overflow trays

Reinstall all the assemblies and setup the instrument in its initial configuration.

3 - Analysis circuit decontamination (30 min)

• BOTTLE VERSION

- Prepair 1 bottle containing 1/2 litre of bleach diluted to 1 part of bleach for 9 parts of deionize water (1/10).

- Prepair 1 bottle containing 1/2 litre of distilled water.
- Switch on the instrument
- Replace the reagent bottles by the diluted bleach bottle.
- Run a SERVICE / PRIME / ALL REAGENTS cycle.
- Fill a sample tube with diluted bleach to 1 part of bleach for 4 of deionize water (1/5).
- Enter the TECHNICIAN / BURN-IN function, Type in 15 cycles and leave the instrument operating until it stops (On MICROS 60 OT run 15 manual cycles).

RAS 192 A Ind.A

DECONTAMINATION

• PACK VERSION

- Prepair 1 bottle containing 1/2 litre of bleach diluted to 1 part of bleach for 9 parts of deionize water (1/10).

- Prepair 1 bottle containing 1/2 litre of distilled water.
- Prepair one empty bottle of 1 litre for waste.
- Switch on the instrument
- Replace the reagent pack by the Drain & Rinse kit (XEA 349 AS).

- Plunge the straws into the diluted bleach bottle and the waste tube into the empty waste bottle.

- Run a SERVICE / PACK / PRIME cycle.
- Fill a sample tube with diluted bleach to 1 part of bleach for 4 of deionize water (1/5).

- Enter the TECHNICIAN / BURN-IN function, Type in 15 cycles and leave the instrument operating until it stops (On MICROS 60 OT run 15 manual cycles).

4 - Drain and rinse (30 min)

- Remove the 3 reagent straws from the bottle containing the diluted bleach
- Wrap the straws in absorbant paper.
- Run two prime cycles : the bleach is drained.
- Replace the diluted bleach by the distilled water bottle and re-plunge the straws in distilled water.
- Run six PRIME / ALL REAGENTS cycles (Rinse).
- Remove the 3 reagent straws from the distilled water (Wrap the straws in absorbant paper).
- Run two PRIME / ALL REAGENTS cycles : the distilled water is drained.
- Run a STAND BY cycle.
- Re-install the reagent bottles and the straws (or re-install the Pack instead of the Drain & Rinse kit).
- Switch off the instrument.
- Close the instrument cover.

DRAIN & RINSE

· CONCERNS

Instrument Rinse and drain before - an extended shutdown - an instrument removing

· REQUIRED TOOLS

None

S

· REQUIRED PRODUCTS

Distilled water

INTERVENTION TIME

35min

• FREQUENCY

On request

· SPECIFIC KIT OR CONSUMABLES

Drain & rinse kit : **XEA 349 AS** for Pack equipped instrument **ADVIA** Pack optional : Reagent output protections FFZ 015 A

RAS 197 A Ind.B



DRAIN & RINSE

PROCEDURE

WARNING !

Disposal gloves and white coat must be worn by the operator.
Local or national regulations must be applied in all the operations .

1 - Preliminary (5min)

- Switch on the instrument
- Run a STARTUP cycle, then a SERVICE / AUTOCLEAN

2 - Drain and rinse (30 min)

• BOTTLE VERSION

- Prepair one bottle containing 1/2 litre of distilled water
- Remove the 3 reagent straws MINIDIL, MINILYSE, MINICLEAN from the bottles.
- Wrap the straws in absorbant paper.
- Run two PRIME / ALL REAGENTS cycles : the reagents are drained.
- Replace the reagent bottles by the distilled water bottle and plunge the straws into distilled water.
- Run 6 PRIME / ALL REAGENTS cycles (Rinse).
- Remove the 3 reagent straws from the distilled water (Wrap the straws in absorbant paper).
- Run two PRIME / ALL REAGENTS cycles : the distilled water is drained.
- Run a STAND BY cycle.
- Check that the diluent syringe piston is in park position (upper position).
- Remove the distilled water and install the installation kit box instead.
- Install the black plastic carriage locking clip in order to block the needle carriage (see INS-
- TALLATION procedures :

MICROS 60 CT : RAS 166 A

MICROS 60 OT : RAS 165 A)

- Clean the reagent stains from the instrument.
- Put an adhesive tape on the tube holder door (MICROS 60 CT) to prevent from opening it.
- Switch the instrument off.

DRAIN & RINSE

• PACK VERSION

- Prepair one bottle containing 1/2 litre of distilled water.
- Prepair one empty bottle of 1/2 litre for waste.
- Install the Drain & Rinse kit (XEA 349 AS) instead of the Reagent pack.
- Plunge the waste tube into the empty waste bottle.
- Wrap the 3 reagent straws MINIDIL, MINILYSE, MINICLEAN in absorbant paper.
- Run two SERVICE / PACK / PRIME cycles : the reagents are drained.
- Plunge the straws into distilled water.
- Run 6 SERVICE / PACK / PRIME cycles (Rinse).

- Remove the 3 reagent straws from the distilled water (Wrap the straws in absorbant paper).

- Run two SERVICE / PACK / PRIME cycles : the distilled water is drained.
- Run a STAND BY cycle.
- Check that the diluent syringe piston is in park position (upper position).
- Remove the Drain & rinse kit.
- Switch off the instrument.
- Install the black plastic carriage locking clip in order to block the needle carriage (see INSTALLATION procedures :

MICROS 60 CT : RAS 166 A MICROS 60 OT : RAS 165 A)

- Clean the reagent stains from the instrument.

- Put an adhesive tape on the tube holder door (MICROS 60 CT) to prevent from opening it.

- Switch the instrument off.



 ADVIA : Install the reagent ouput protections (see the warning sheet form : RAL 035 A and join it to the installation kit box)

- Install the installation kit box on the pack location.

• CONCERNS

- Sampling position and available tubes
- Piercing block description
- Maintenance

· REQUIRED TOOLS

- Torx keys

· REQUIRED PRODUCTS

- See decontamination procedure : RAS 192 A

INTERVENTION TIME

- 30 min

• FREQUENCY

- On request

· SPECIFIC KIT OR CONSUMABLES

- None



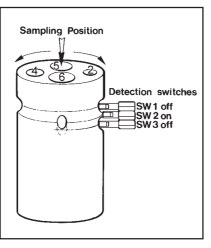
RAS 198 A Ind.A

PROCEDURE

1 - Sampling position



Diag.1



According to the sample tube models (Vacutainers, Microtainers ...) used and to their position into the sample tube holder, the heigth of the needle in its lower position can be modified.

The 3 switches (diag 1) associated to the tube holder are able to detect the sampling position according to the following principle :

Two states 0/1 are possible for the 3 switches (see diag 2) :

- Switched OFF : 1
- Switched ON : 0

The binary codes obtained from the states of the switches gives the positions of the tube holder.

For each position of the sample tube holder (from 1 to 6) corresponds a position of the needle (from 1 to 6).

Diag.2

SWITCH 1	SWITCH 2	SWITCH 3	Sampling position	Needle
0	0	0	Bad position of the tube holder	
1	0	0	position 1	Needle 1
0	1	0	position 2	Needle 2
1	1	0	position 3	Needle 3
0	0	1	position 4	Needle 4
1	0	1	position 5	Needle 5
0	1	1	position 6	Needle 6
1	1	1	No tube holder	



The code «0 0 0» means that the tube holder has been turned in between two sampling positions. The code «1 1 1» means that the tube holder has been removed.

The following chart tables give the tube positions available according to the models of sample holder used on the instrument

Diag.3

MICROS 60 STANDARD TUBE HOLDER



Diag.4

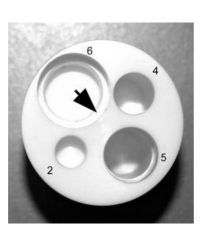
							Diag.					
Barrillet standard (GBC061AS) <i>Standard</i> <i>Tube holder</i>	Marque <i>Trademark</i>	Additif <i>Additive</i>	Volume <i>Volume</i>	Vide <i>Vacuum</i>	Volume mort (5) Dead volume	Ref. <i>P/N</i>	Modèle <i>Model</i>	Photo N° Photo #	Réglage (1) Adjustment	Perçage tube Tube piercing	Bouchon (2) <i>Stopper</i>	
POSITION 6 CONTROLE	R&D system	na	2ml	na	250µl	na	na	8	standard	sans bouchon without stopper		
	Becton D.	EDTA-K3	5ml	nc	390µl	368452	Vacutainer	9	standard	avec bouchon with stopper	gomme rubber	
	Becton D.	EDTA-K3	5ml	2ml	400µl	367651	Vacutainer	11	standard	avec bouchon with stopper	hemogard	
	Becton D.	EDTA-K3	5ml	3ml	410µl	367856	Vacutainer	10	standard	avec bouchon with stopper	hemogard	
	Becton D.	EDTA-K3	5ml	3ml	400µl	367652	Vacutainer	12	standard	avec bouchon with stopper	hemogard	
	Becton D.	EDTA-K3	5ml	4.5ml	410µl	367654	Vacutainer	7	standard	avec bouchon with stopper	hemogard	
	Terumo	EDTA-K2	5ml	3ml	430µl	VP-053SDK	Venoject II	15	standard	avec bouchon with stopper	Ultraseal	
POSITION 5 STANDARD	Terumo	EDTA-K3	5ml	5ml	460µl	VT-050STK	Venoject	17	standard	avec bouchon with stopper	gomme rubber	
	Terumo	EDTA-K3	5ml	3ml	460µl (4)	VT-053STK	Venoject	na	standard	avec bouchon with stopper	gomme rubber	
	ABX	EDTA-K3	5ml	4ml	480µl	ABX-3004002	na	14	standard	avec bouchon with stopper	gomme (non recommande	
	Greiner	EDTA-K3	5ml	2ml	370µl (4)	454087	Vacuette	na	standard	avec bouchon with stopper	hemogard	
	Greiner	EDTA-K3	5ml	3ml	370µl (4)	454086	Vacuette	na	standard	avec bouchon with stopper	hemogard	
	Greiner	EDTA-K3	5ml	4ml	370µl	454036	Vacuette	13	standard	avec bouchon with stopper	hemogard	
	LDM	EDTA-KE	5ml	4.5ml	480µl	nc	nc	2	standard	avec bouchon with stopper	hemogard	
POSITION 4	Becton D.	EDTA-K3	3ml	na	30µI	6385	Vacutainer	6	standard	sans bouchon (3) without stopper	gomme (non recommande rubber (not recommended	
	Terumo	EDTA-K3	3ml	3ml	30µI	VT-030STK	Venoject	16	standard	avec bouchon with stopper	gomme (non recommande rubber (not recommended	
	Sarstedt	nc	0.5ml	na	30µI	901091	nc	4	standard	sans bouchon without stopper	bouchon imperdable	
POSITION 2	ABX	nc	0.5ml	na	30µl	ABX-3001001	nc	5	standard	sans bouchon without stopper	bouchon imperdable unlostable stopper	

Table 1

CAUTION

The needle sampling position must not be modified without refering to the procedure RAS 176 A.

ADVIA 60 STANDARD TUBE HOLDER



Diag.5



Diag.6

Barrillet standard (GBC216AS) Standard Tube holder	Marque Trademark	Additif <i>Additive</i>	Volume <i>Volume</i>	Vide <i>Vacuum</i>	Volume mort (5) Dead volume	Ref. <i>P/N</i>	Modèle <i>Model</i>	Photo N° <i>Photo #</i>	Réglage (1) Adjustment	Perçage tube Tube piercing	Bouchon (2) Stopper
POSITION 6	STRECK	nc	nc	nc	nc	nc	nc	18	standard	avec bouchon with stopper	nc
CONTROLE	R&D system	na	2ml	na	250µl	na	na	8	standard	sans bouchon without stopper	
	Becton D.	EDTA-K3	5ml	nc	390µI	368452	Vacutainer	9	standard	avec bouchon with stopper	gomme rubber
	Becton D.	EDTA-K3	5ml	2ml	400µl	367651	Vacutainer	11	standard	avec bouchon with stopper	hemogard
	Becton D.	EDTA-K3	5ml	3ml	410µl	367856	Vacutainer	10	standard	avec bouchon with stopper	hemogard
	Becton D.	EDTA-K3	5ml	3ml	400µl	367652	Vacutainer	12	standard	avec bouchon with stopper	hemogard
	Becton D.	EDTA-K3	5ml	4.5ml	410µI	367654	Vacutainer	7	standard	avec bouchon with stopper	hemogard
	Terumo	EDTA-K2	5ml	3ml	430µI	VP-053SDK	Venoject II	15	standard	avec bouchon with stopper	Ultraseal
POSITION 5	Terumo	EDTA-K3	5ml	5ml	460µI	VT-050STK	Venoject	17	standard	avec bouchon with stopper	gomme rubber
	Terumo	EDTA-K3	5ml	3ml	460µl (4)	VT-053STK	Venoject	na	standard	avec bouchon with stopper	gomme rubber
	ABX	EDTA-K3	5ml	4ml	480µl	ABX-3004002	na	14	standard	avec bouchon with stopper	gomme (non recommand rubber (not recommended
	Greiner	EDTA-K3	5ml	2ml	370µI (4)	454087	Vacuette	na	standard	avec bouchon with stopper	hemogard
	Greiner	EDTA-K3	5ml	3ml	370µI (4)	454086	Vacuette	na	standard	avec bouchon with stopper	hemogard
	Greiner	EDTA-K3	5ml	4ml	370µI	454036	Vacuette	13	standard	avec bouchon with stopper	hemogard
	LDM	EDTA-KE	5ml	4.5ml	480µl	nc	nc	2	standard	avec bouchon with stopper	hemogard
POSITION 4	Becton D.	EDTA-K3	3ml	na	30µI	6385	Vacutainer	6	standard	sans bouchon (3) without stopper	gomme (non recommand rubber (not recommended
	Terumo	EDTA-K3	3ml	3ml	30µl	VT-030STK	Venoject	16	standard	avec bouchon with stopper	gomme (non recommand rubber (not recommended
	Sarstedt	nc	0.5ml	na	30µl	901091	nc	4	standard	sans bouchon without stopper	bouchon imperdable unlostable stopper
POSITION 2	ABX	nc	0.5ml	na	30µl	ABX-3001001	nc	5	standard	sans bouchon	bouchon imperdable unlostable stopper

Table 2

OPTIONAL TUBE HOLDER



Diag.8



Diag.9

Groove

Barrillet optionnel (GBC217AS) <i>Optional</i> <i>Tube holder</i>	Marque Trademark	Additif <i>Additive</i>	Volume <i>Volume</i>	Vide <i>Vacuum</i>	Volume mort (5) Dead volume	Ref. <i>P/N</i>	Modèle <i>Model</i>	Photo N° Photo #	Réglage (1) <i>Adjustment</i>	Perçage tube Tube piercing	Bouchon (2) Stopper
POSITION 6 CONTROLE	R&D system	na	2ml	na	250µl	na	na	8	standard	sans bouchon without stopper	na
	Becton D.	EDTA-K3	5ml	nc	390µl	368452	Vacutainer	9	standard	avec bouchon with stopper	gomme rubber
	Becton D.	EDTA-K3	5ml	2ml	400µl	367651	Vacutainer	11	standard	avec bouchon with stopper	hemogard
	Becton D.	EDTA-K3	5ml	3ml	410µl	367856	Vacutainer	10	standard	avec bouchon with stopper	hemogard
	Becton D.	EDTA-K3	5ml	3ml	400µl	367652	Vacutainer	12	standard	avec bouchon with stopper	hemogard
	Becton D.	EDTA-K3	5ml	4.5ml	410µl	367654	Vacutainer	7	standard	avec bouchon with stopper	hemogard
	Terumo	EDTA-K2	5ml	3ml	430µl	VP-053SDK	Venoject II	15	standard	avec bouchon with stopper	Ultraseal
POSITION 5 STANDARD	Terumo	EDTA-K3	5ml	5ml	460µl	VT-050STK	Venoject	17	standard	avec bouchon with stopper	gomme rubber
	Terumo	EDTA-K3	5ml	3ml	460µl (4)	VT-053STK	Venoject	na	standard	avec bouchon with stopper	gomme rubber
	ABX	EDTA-K3	5ml	4ml	480µl	ABX-3004002	nc	14	standard	avec bouchon with stopper	gomme (non recommandé) rubber (not recommended)
	Greiner	EDTA-K3	5ml	2ml	370µl (4)	454087	Vacuette	na	standard	avec bouchon with stopper	hemogard
	Greiner	EDTA-K3	5ml	3ml	370µl (4)	454086	Vacuette	na	standard	avec bouchon with stopper	hemogard
	Greiner	EDTA-K3	5ml	4ml	370µl	454036	Vacuette	13	standard	avec bouchon with stopper	hemogard
	LDM	EDTA-KE	5ml	4.5ml	480µl	nc	nc	2	standard	avec bouchon with stopper	hemogard
POSITION 3	Becton D.	nc	0.5ml	na	30µI	365975	Microtainer	3	avec tube with tube	sans bouchon without stopper	Microgard (Equipé adapteur: autre réglage aiguille) (Equipped with adaptor: other adjustment)
POSITION 1	Becton D.	nc	0.5ml	na	30µI	365973	Microtainer	1	standard	sans bouchon without stopper	na

Table 3



na - not applicable.

nc - not communicated.

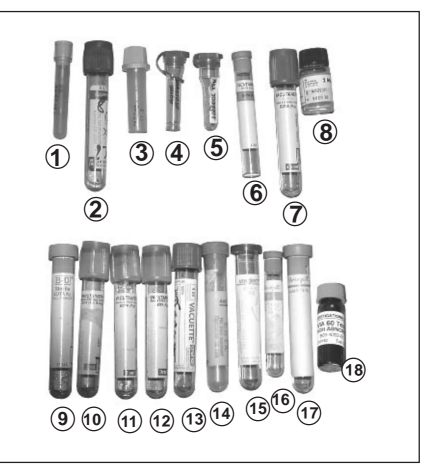
(1) - The standard adjustment positions the needle at the bottom of the tube holder, when mentioned "with tube", the presence of the tube is mandatory for the adjustment.

(2) - More information about sample tube is available in the user manual, section 3 "SPECIFICA-TIONS", point 3.4.2.

(3) - The thickness of the tube stopper blocks the tube into the piercing mechanism.

(4) - These volumes have been calculated; not measured.

(5) - The "dead volume" is determined after the manual adjustment of the sampling depth and increased by 20%, except in the pediatric tubes where a 30μ I volume has been fixed arbitrarily for security.



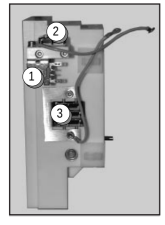




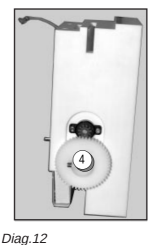
The needle heigth adjustments is explained in the procedure RAS 176 A.

3 - Description

Piercing block right hand side view



Piercing block left hand side view

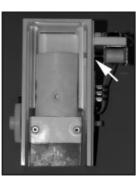


Door lock electro magnet

- 2 Door detection switch
- 3 Sampling position switches
- 4 Gearings

Diag.11

2 - Maintenance



- Control the correct operation of the door lock electro magnet (diag 13).

Diag.13



Diag.14



- The sample tube holder can be pulled out from its location as shown on diagram 15.

- To clean the piercing block it is easier to dismantle the door

- Re-install the reverse order.

front panel as shown on the diag 14.

Diag.15

CAUTION

Decontamination procedure (RAS 192 A) must be followed to clean the sample tube holder and its location.

CONTENTS

1. ARGOS FORMAT PRINCIPLES	2
1.1. Introduction 1.2. Results characteristics	2 2
1.2.1. Key	
1.2.2. Result format	3
1.2.3. End of communication key	4
2. STANDARD FORMAT	5
2.1 Message Structure	5
2.2. Details about the structure	
2.3. Identifier list and their formats	
2.3.1. Hematologyc numeric parameters	
2.3.1.1. Format description	
2.3.1.2. Identifier list	
2.3.2. Pathology	
2.3.3. Histograms and matrix	
2.3.3.1. Format description	
2.3.3.2. Identifier list	
2.3.4. Patient result identification	
2.3.4.2. Identifier list	
2.3.4.3. Analysis type (\$80)	
2.4. Heading title	
2.5. Other identifiers	9
	J
3. PIN ASSIGNMENTS 1	0

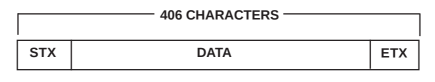
3 RS ouput formats are available on MICROS 60 :

- Format 1 : ARGOS / HELIOS type
- Format 2 : ABX internal format (factory use)
- Standard format
- TR off : Transmission off

1. ARGOS FORMAT PRINCIPLES

1.1. Introduction

The ARGOS format is a fixed format (406 characters for one result) including a STX and a ETX. These characters are splitted into fields representing a transmitted item.





The fields have a fixed length separated by the **/OD** character.

1.2. Results characteristics

1.2.1. Key

Total ASCII characters emitted : 406

- (-) : Space \$20

- (]) : Carriage return \$0D

- CRC : exclusive "OR" of all the transmitted bytes except ETX and STX, then an inclusive "OR" with a \$40 value.

zzzzz : numeric field completed by zeros on the left.
 ex : 04.55 (decimal separation with a period).
 When the analyser does not transmit parameters, the field (zzzzz) is put in place of (--.-).

- Y : Alphanumeric character from \$20 to \$7F.

- # : Space (\$20) if automatic sampling. Star (\$2A) if manual sampling.

1.2.2. Result format

Line 1: STX (\$02) Start of text 1 Line 3: z2 Analyser No 2 + 1 Line 4: YYYYYYYYYYYYYYYYYYYYYY Hoefficiation 30 + 1 Line 6: YYYYYYYYYYYYYYYYYYYYYY Hienfificiation 30 + 1 Line 6: YYYYYYYYYYYYYYYYYYYYYYYYYYYYYYYYYYYY					
Line 2: R (\$52) Character 'R" 1 Line 3: ZZ Analyser No 2 + 1 Line 4: YYYYYYYYYYYYYYYYYYYYYYYYYYYYYYYYYYYY	Lino 1 ·	STV (\$02)		Stort of toxt	1
Line 3: zzj Analyser No 2 + 1 Line 4: YYYYYYYYYYYYY Identification No 16 + 1 Line 6: YYYYYYYYYYYYYYYY Identification No 16 + 1 Line 6: zzzzz-RNJ GB rejection & limits 8 + 1 Line 7: zzzzz-RNJ LYC% rejection & limits 8 + 1 Line 9: zzzzz-RNJ LYC% rejection & limits 8 + 1 Line 10: zzzzz-RNJ MONM rejection & limits 8 + 1 Line 11: zzzz-RNJ MONM rejection & limits 8 + 1 Line 12: zzzz-RNJ GRA# rejection & limits 8 + 1 Line 13: zzzz-RNJ NEU½ rejection & limits 8 + 1 Line 14: zzzz-RNJ EOS% rejection & limits 8 + 1 Line 15: zzzz-RNJ EOS% rejection & limits 8 + 1 Line 16: zzzz-RNJ EOS% rejection & limits 8 + 1 Line 17: zzzz-RNJ EOS% rejection & limits 8 + 1 Line 12: zzzzz-RNJ LIC% rejection & limits <td></td> <td></td> <td></td> <td></td> <td></td>					
Line 4: YVYYYYYYYYYYYYYYYYYYYYYYYYYYYYYYYYYYY					
Line 5: YYYYYYYYYYYYYYYYYYYYYYYYYYYYYYYYYYYY		-			
Line 6: zztzzt zzzzz-RNj Time & Date 20+1 Line 7: zzzz - RNj LYC# rejection & limits 8+1 Line 9: zzzz - RNj LYC# rejection & limits 8+1 Line 9: zzzz - RNj LYC# rejection & limits 8+1 Line 10: zzzz - RNj MONN# rejection & limits 8+1 Line 11: zzzz - RNj GRA# rejection & limits 8+1 Line 13: zzzz - RNj GRA# rejection & limits 8+1 Line 14: zzzzz - RNj EOS# rejection & limits 8+1 Line 15: zzzzz - RNj EOS# rejection & limits 8+1 Line 16: zzzzz - RNj EOS# rejection & limits 8+1 Line 12: zzzzz - RNj ALY# rejection & limits 8+1 Line 21: zzzzz - RNj LIC% rejection & limits 8+1 Line 22: zzzzz - RNj LIC% rejection & limits 8+1 Line 23: zzzzz - RNj RET		-	~~~~~		
Line 7: 2222-RN] GB rejection & limits 8 + 1 Line 8: 22222-RN] LYC?# rejection & limits 8 + 1 Line 10: 22222-RN] LYC?# rejection & limits 8 + 1 Line 11: 22222-RN] MONP# rejection & limits 8 + 1 Line 12: 22222-RN] GRA# rejection & limits 8 + 1 Line 13: 22222-RN] GRA# rejection & limits 8 + 1 Line 14: 22222-RN] GRA# rejection & limits 8 + 1 Line 15: 22222-RN] EOS# rejection & limits 8 + 1 Line 16: 22222-RN] EOS# rejection & limits 8 + 1 Line 17: 22222-RN] EOS% rejection & limits 8 + 1 Line 21: 22222-RN] BAS# rejection & limits 8 + 1 Line 22: 22222-RN] LiC% rejection & limits 8 + 1 Line 22: 22222-RN] LiC% rejection & limits 8 + 1 Line 22: 22222-RN] RET* rejection & limits 8 + 1 Line 22:			YYYYYJ		
Line 8: zzzzz-RN LYG# rejection & limits 8 + 1 Line 9: zzzzz-RN LYG% rejection & limits 8 + 1 Line 10: zzzzz-RN MON# rejection & limits 8 + 1 Line 11: zzzzz-RN MON% rejection & limits 8 + 1 Line 12: zzzzz-RN GRA# rejection & limits 8 + 1 Line 13: zzzzz-RN GRA# rejection & limits 8 + 1 Line 14: zzzzz-RN REM rejection & limits 8 + 1 Line 15: zzzzz-RN EOS% rejection & limits 8 + 1 Line 16: zzzzz-RN EOS% rejection & limits 8 + 1 Line 17: zzzzz-RN EOS% rejection & limits 8 + 1 Line 18: zzzzz-RN EOS% rejection & limits 8 + 1 Line 21: zzzzz-RN ALY# rejection & limits 8 + 1 Line 22: zzzzz-RN ALY# rejection & limits 8 + 1 Line 23: zzzzz-RN RET" rejection & limits 8 + 1 Line 24: zzzzz-	Line 6 :	zz/zz/zz-zzhzzmnzzs#]		Time & Date	20 + 1
Line 9: 22222-RN] LYC% rejection & limits 8 + 1 Line 10: 22222-RN] MON% rejection & limits 8 + 1 Line 11: 22222-RN] GRA# rejection & limits 8 + 1 Line 12: 22222-RN] GRA# rejection & limits 8 + 1 Line 13: 22222-RN] GRA# rejection & limits 8 + 1 Line 14: 22222-RN] GRA# rejection & limits 8 + 1 Line 15: 22222-RN] EOS% rejection & limits 8 + 1 Line 16: 22222-RN] EOS% rejection & limits 8 + 1 Line 17: 22222-RN] BAS# rejection & limits 8 + 1 Line 21: 22222-RN] BAS# rejection & limits 8 + 1 Line 22: 22222-RN] LIC% rejection & limits 8 + 1 Line 23: 22222-RN] LIC% rejection & limits 8 + 1 Line 24: 22222-RN] RET rejection & limits 8 + 1 Line 25: 22222-RN] RET rejection & limits 8 + 1 Line 26:	Line 7 :	zzzz-RN]	GB	rejection & limits	8 + 1
Line 10: zzzzz-RN MON# rejection & limits 8 + 1 Line 11: zzzzz-RN GRA% rejection & limits 8 + 1 Line 12: zzzzz-RN GRA% rejection & limits 8 + 1 Line 13: zzzzz-RN GRA% rejection & limits 8 + 1 Line 14: zzzzz-RN NEU# rejection & limits 8 + 1 Line 15: zzzz-RN NEU# rejection & limits 8 + 1 Line 16: zzzzz-RN EOS# rejection & limits 8 + 1 Line 17: zzzzz-RN EOS# rejection & limits 8 + 1 Line 18: zzzzz-RN BAS% rejection & limits 8 + 1 Line 20: zzzzz-RN BAS% rejection & limits 8 + 1 Line 21: zzzzz-RN LiC# rejection & limits 8 + 1 Line 22: zzzzz-RN LiC# rejection & limits 8 + 1 Line 23: zzzzz-RN RET* rejection & limits 8 + 1 Line 26: zzzzz-RN RET* rejection & limits 8 + 1 Line 27: zzzz	Line 8 :	zzzz-RN]	LYC#	rejection & limits	8 + 1
Line 11: zzzzz-RN MON% rejection & limits 8 + 1 Line 12: zzzzz-RN GRA# rejection & limits 8 + 1 Line 13: zzzzz-RN GRA% rejection & limits 8 + 1 Line 14: zzzzz-RN NEU# rejection & limits 8 + 1 Line 15: zzzz-RN NEU% rejection & limits 8 + 1 Line 16: zzzzz-RN EOS# rejection & limits 8 + 1 Line 17: zzzz-RN EOS# rejection & limits 8 + 1 Line 18: zzzz-RN EOS# rejection & limits 8 + 1 Line 21: zzzzz-RN BAS# rejection & limits 8 + 1 Line 22: zzzzz-RN LIC# rejection & limits 8 + 1 Line 23: zzzzz-RN LIC# rejection & limits 8 + 1 Line 24: zzzzz-RN RET* rejection & limits 8 + 1 Line 25: zzzzz-RN RET* rejection & limits 8 + 1 Line 26: zzzzz-RN RET* rejection & limits 8 + 1 Line 27: zzzzz-	Line 9 :	zzzz-RN]	LYC%	rejection & limits	8 + 1
Line 12: zzzzz-RN] GRA# rejection & limits 8 + 1 Line 13: zzzzz-RN] GRA% rejection & limits 8 + 1 Line 15: zzzzz-RN] NEU% rejection & limits 8 + 1 Line 16: zzzzz-RN] EOS# rejection & limits 8 + 1 Line 16: zzzzz-RN] EOS# rejection & limits 8 + 1 Line 16: zzzzz-RN] BAS# rejection & limits 8 + 1 Line 17: zzzzz-RN] BAS# rejection & limits 8 + 1 Line 21: zzzzz-RN] BAS# rejection & limits 8 + 1 Line 21: zzzzz-RN] ALY# rejection & limits 8 + 1 Line 22: zzzzz-RN] ALY# rejection & limits 8 + 1 Line 23: zzzzz-RN] RET rejection & limits 8 + 1 Line 24: zzzzz-RN] RET rejection & limits 8 + 1 Line 25: zzzzz-RN] RET rejection & limits 8 + 1 Line 26: zzzzz-RN] RET rejection & limits 8 + 1 Line 27:	Line 10 :	zzzz-RN]	MON#	rejection & limits	8 + 1
Line 13: zzzzz-RN] GRA% rejection & limits 8 + 1 Line 14: zzzzz-RN] NEU# rejection & limits 8 + 1 Line 15: zzzzz-RN] EOS# rejection & limits 8 + 1 Line 16: zzzzz-RN] EOS# rejection & limits 8 + 1 Line 17: zzzzz-RN] EOS# rejection & limits 8 + 1 Line 18: zzzzz-RN] BAS# rejection & limits 8 + 1 Line 20: zzzzz-RN] ALY# rejection & limits 8 + 1 Line 21: zzzzz-RN] ALY# rejection & limits 8 + 1 Line 22: zzzzz-RN] LIC% rejection & limits 8 + 1 Line 23: zzzzz-RN] LIC% rejection & limits 8 + 1 Line 24: zzzzz-RN] RET* rejection & limits 8 + 1 Line 25: zzzzz-RN] RET* rejection & limits 8 + 1 Line 26: zzzzz-RN] RET rejection & limits 8 + 1 Line 27: zzzzz-RN] RET rejection & limits 8 + 1 Line 27:	Line 11 :	zzzz-RN]	MON%	rejection & limits	8 + 1
Line 14: zzzzz-RN] NEU# rejection & limits 8 + 1 Line 15: zzzzz-RN] EOS# rejection & limits 8 + 1 Line 16: zzzzz-RN] EOS% rejection & limits 8 + 1 Line 17: zzzzz-RN] EOS% rejection & limits 8 + 1 Line 18: zzzzz-RN] BAS% rejection & limits 8 + 1 Line 20: zzzzz-RN] BAS% rejection & limits 8 + 1 Line 21: zzzzz-RN] ALY# rejection & limits 8 + 1 Line 22: zzzzz-RN] LIC% rejection & limits 8 + 1 Line 23: zzzzz-RN] RET* rejection & limits 8 + 1 Line 24: zzzzz-RN] RET* rejection & limits 8 + 1 Line 25: zzzzz-RN] RET* rejection & limits 8 + 1 Line 26: zzzzz-RN] REC rejection & limits 8 + 1 Line 27: zzzzz-RN] REC rejection & limits 8 + 1 Line 28: zzzzz-RN] REC rejection & limits 8 + 1 Line 31:	Line 12 :	zzzz-RN]	GRA#	rejection & limits	8 + 1
Line 15: zzzzz-RN] NEU% rejection & limits 8 + 1 Line 16: zzzzz-RN] EOS# rejection & limits 8 + 1 Line 16: zzzzz-RN] EOS% rejection & limits 8 + 1 Line 17: zzzzz-RN] EOS% rejection & limits 8 + 1 Line 18: zzzzz-RN] BAS% rejection & limits 8 + 1 Line 21: zzzzz-RN] ALY# rejection & limits 8 + 1 Line 22: zzzzz-RN] ALY# rejection & limits 8 + 1 Line 23: zzzzz-RN] LIC# rejection & limits 8 + 1 Line 24: zzzzz-RN] RET* rejection & limits 8 + 1 Line 25: zzzzz-RN] RET* rejection & limits 8 + 1 Line 26: zzzzz-RN] RET* rejection & limits 8 + 1 Line 27: zzzzz-RN] RET rejection & limits 8 + 1 Line 28: zzzzz-RN] MCV rejection & limits 8 + 1 Line 31: zzzzz-RN] MCV rejection & limits 8 + 1 Line 32:	Line 13 :	zzzz-RN]	GRA%	rejection & limits	8 + 1
Line 15: zzzzz-RN] NEU% rejection & limits 8 + 1 Line 16: zzzzz-RN] EOS# rejection & limits 8 + 1 Line 16: zzzzz-RN] EOS% rejection & limits 8 + 1 Line 17: zzzzz-RN] EOS% rejection & limits 8 + 1 Line 18: zzzzz-RN] BAS% rejection & limits 8 + 1 Line 21: zzzzz-RN] ALY# rejection & limits 8 + 1 Line 22: zzzzz-RN] ALY# rejection & limits 8 + 1 Line 23: zzzzz-RN] LIC# rejection & limits 8 + 1 Line 24: zzzzz-RN] RET* rejection & limits 8 + 1 Line 25: zzzzz-RN] RET* rejection & limits 8 + 1 Line 26: zzzzz-RN] RET* rejection & limits 8 + 1 Line 27: zzzzz-RN] RET rejection & limits 8 + 1 Line 28: zzzzz-RN] MCV rejection & limits 8 + 1 Line 31: zzzzz-RN] MCV rejection & limits 8 + 1 Line 32:	Line 14 :	zzzz-RN]	NEU#	rejection & limits	8 + 1
Line 16: zzzzz-RN] EOS# rejection & limits 8 + 1 Line 17: zzzzz-RN] EOS% rejection & limits 8 + 1 Line 18: zzzzz-RN] BAS# rejection & limits 8 + 1 Line 18: zzzzz-RN] BAS# rejection & limits 8 + 1 Line 20: zzzzz-RN] ALY# rejection & limits 8 + 1 Line 21: zzzzz-RN] ALY% rejection & limits 8 + 1 Line 22: zzzzz-RN] LIC# rejection & limits 8 + 1 Line 23: zzzzz-RN] LIC# rejection & limits 8 + 1 Line 24: zzzzz-RN] RET* rejection & limits 8 + 1 Line 25: zzzzz-RN] RET* rejection & limits 8 + 1 Line 26: zzzzz-RN] RET rejection & limits 8 + 1 Line 27: zzzzz-RN] HCT rejection & limits 8 + 1 Line 31: zzzzz-RN] MCV rejection & limits 8 + 1 Line 32: zzzzz-RN] MCH rejection & limits 8 + 1 Line 32:	Line 15 :	-	NEU%	-	8 + 1
Line 17: zzzzz-RN EOS% rejection & limits 8 + 1 Line 18: zzzzz-RN BAS# rejection & limits 8 + 1 Line 19: zzzzz-RN BAS% rejection & limits 8 + 1 Line 20: zzzzz-RN ALY4 rejection & limits 8 + 1 Line 21: zzzzz-RN ALY4 rejection & limits 8 + 1 Line 22: zzzzz-RN LIC# rejection & limits 8 + 1 Line 23: zzzzz-RN LIC# rejection & limits 8 + 1 Line 24: zzzzz-RN RET* rejection & limits 8 + 1 Line 25: zzzzz-RN RET* rejection & limits 8 + 1 Line 26: zzzzz-RN RET* rejection & limits 8 + 1 Line 27: zzzzz-RN REC rejection & limits 8 + 1 Line 28: zzzzz-RN HGB rejection & limits 8 + 1 Line 31: zzzzz-RN MCV rejection & limits 8 + 1 Line 32: zzzzz-RN MCV rejection & limits 8 + 1 Line 33: zzzzz-R	Line 16 :		EOS#	-	8 + 1
Line 18: zzzzz-RN BAS# rejection & limits 8+1 Line 19: zzzzz-RN BAS% rejection & limits 8+1 Line 20: zzzzz-RN ALY# rejection & limits 8+1 Line 21: zzzzz-RN ALY% rejection & limits 8+1 Line 21: zzzzz-RN ALY% rejection & limits 8+1 Line 22: zzzzz-RN LIC# rejection & limits 8+1 Line 23: zzzzz-RN LIC# rejection & limits 8+1 Line 24: zzzzz-RN RET* rejection & limits 8+1 Line 25: zzzzz-RN RET* rejection & limits 8+1 Line 26: zzzzz-RN RET* rejection & limits 8+1 Line 27: zzzzz-RN REC rejection & limits 8+1 Line 28: zzzzz-RN MCV rejection & limits 8+1 Line 30: zzzzz-RN MCV rejection & limits 8+1 Line 31: zzzzz-RN MCHC rejection & limits 8+1 Line 32: zzzzz-RN MCHC		•		•	
Line 19: zzzzz-RN] BAS% rejection & limits 8 + 1 Line 20: zzzzz-RN] ALY# rejection & limits 8 + 1 Line 21: zzzzz-RN] ALY# rejection & limits 8 + 1 Line 21: zzzzz-RN] LIC# rejection & limits 8 + 1 Line 22: zzzzz-RN] LIC# rejection & limits 8 + 1 Line 23: zzzzz-RN] LIC# rejection & limits 8 + 1 Line 24: zzzzz-RN] RET* rejection & limits 8 + 1 Line 25: zzzzz-RN] RET* rejection & limits 8 + 1 Line 26: zzzzz-RN] RET* rejection & limits 8 + 1 Line 27: zzzzz-RN] RET* rejection & limits 8 + 1 Line 28: zzzzz-RN] HCT rejection & limits 8 + 1 Line 31: zzzzz-RN] MCV rejection & limits 8 + 1 Line 32: zzzzz-RN] MCV rejection & limits 8 + 1 Line 32: zzzzz-RN] MCV rejection & limits 8 + 1 Line 32:		•		•	
Line 20: zzzzz-RN] ALY# rejection & limits 8+1 Line 21: zzzzz-RN] ALY% rejection & limits 8+1 Line 22: zzzzz-RN] LIC# rejection & limits 8+1 Line 22: zzzzz-RN] LIC# rejection & limits 8+1 Line 24: zzzzz-RN] LIC# rejection & limits 8+1 Line 25: zzzzz-RN] RET* rejection & limits 8+1 Line 26: zzzzz-RN] RET* rejection & limits 8+1 Line 27: zzzzz-RN] RET* rejection & limits 8+1 Line 28: zzzzz-RN] RET rejection & limits 8+1 Line 29: zzzzz-RN] REC rejection & limits 8+1 Line 31: zzzzz-RN] MCV rejection & limits 8+1 Line 32: zzzzz-RN] MCH rejection & limits 8+1 Line 32: zzzzz-RN] MCH rejection & limits 8+1 Line 34: zzzzz-RN] MCH rejection & limits 8+1 Line 36: zzzzzz-RN] <		-		-	
Line 21: zzzzz-RN] ALY% rejection & limits 8 + 1 Line 22: zzzzz-RN] LIC# rejection & limits 8 + 1 Line 23: zzzzz-RN] LIC% rejection & limits 8 + 1 Line 24: zzzzz-RN] RET* rejection & limits 8 + 1 Line 25: zzzzz-RN] RET* rejection & limits 8 + 1 Line 26: zzzzz-RN] RET* rejection & limits 8 + 1 Line 27: zzzzz-RN] RET* rejection & limits 8 + 1 Line 28: zzzzz-RN] RET* rejection & limits 8 + 1 Line 29: zzzzz-RN] HGB rejection & limits 8 + 1 Line 30: zzzzz-RN] MCV rejection & limits 8 + 1 Line 31: zzzzz-RN] MCV rejection & limits 8 + 1 Line 32: zzzzz-RN] MCV rejection & limits 8 + 1 Line 32: zzzzz-RN] MCV rejection & limits 8 + 1 Line 33: zzzzz-RN] PLT rejection & limits 8 + 1 Line 35:				•	
Line 22: zzzzz-RN] LIC# rejection & limits 8 + 1 Line 23: zzzzz-RN] LIC% rejection & limits 8 + 1 Line 24: zzzzz-RN] RET* rejection & limits 8 + 1 Line 25: zzzzz-RN] RET* rejection & limits 8 + 1 Line 26: zzzzz-RN] RET* rejection & limits 8 + 1 Line 27: zzzzz-RN] RET* rejection & limits 8 + 1 Line 28: zzzzz-RN] RET* rejection & limits 8 + 1 Line 27: zzzzz-RN] HGB rejection & limits 8 + 1 Line 27: zzzzz-RN] HGB rejection & limits 8 + 1 Line 30: zzzzz-RN] HCT rejection & limits 8 + 1 Line 31: zzzzz-RN] MCV rejection & limits 8 + 1 Line 32: zzzzz-RN] MCHC rejection & limits 8 + 1 Line 33: zzzzz-RN] RDW rejection & limits 8 + 1 Line 36: zzzzz-RN] PLT rejection & limits 8 + 1 Line 37:		•		-	
Line 23 : ZZZZ-RN] LIC% rejection & limits 8 + 1 Line 24 : ZZZZZ-RN] RET* rejection & limits 8 + 1 Line 25 : ZZZZZ-RN] RET* rejection & limits 8 + 1 Line 26 : ZZZZZ-RN] RET* rejection & limits 8 + 1 Line 26 : ZZZZZ-RN] RET* rejection & limits 8 + 1 Line 27 : ZZZZZ-RN] RET* rejection & limits 8 + 1 Line 28 : ZZZZZ-RN] REC rejection & limits 8 + 1 Line 30 : ZZZZZ-RN] HGB rejection & limits 8 + 1 Line 31 : ZZZZZ-RN] MCV rejection & limits 8 + 1 Line 32 : ZZZZZ-RN] MCV rejection & limits 8 + 1 Line 33 : ZZZZZ-RN] MCHC rejection & limits 8 + 1 Line 36 : ZZZZZ-RN] RET* rejection & limits 8 + 1 Line 36 : ZZZZ-RN] PLT rejection & limits 8 + 1 Line 37 : ZZZZZ-RN] PCT rejection & limits 8 + 1 Line 38		-		•	
Line 24 :ZZZZZ-RN]RET*rejection & limits8 + 1Line 25 :ZZZZZ-RN]RET*rejection & limits8 + 1Line 26 :ZZZZZ-RN]RET*rejection & limits8 + 1Line 27 :ZZZZZ-RN]RET*rejection & limits8 + 1Line 28 :ZZZZZ-RN]RECrejection & limits8 + 1Line 29 :ZZZZZ-RN]HGBrejection & limits8 + 1Line 30 :ZZZZZ-RN]HCTrejection & limits8 + 1Line 31 :ZZZZZ-RN]MCVrejection & limits8 + 1Line 32 :ZZZZZ-RN]MCHrejection & limits8 + 1Line 33 :ZZZZZ-RN]MCHrejection & limits8 + 1Line 36 :ZZZZZ-RN]RDWrejection & limits8 + 1Line 36 :ZZZZZ-RN]PLTrejection & limits8 + 1Line 36 :ZZZZZ-RN]PCTrejection & limits8 + 1Line 37 :ZZZZZ-RN]PCTrejection & limits8 + 1Line 38 :ZZZZZ-RN]PCTrejection & limits8 + 1Line 39 :ZZZZZ-RN]PDWrejection & limits8 + 1Line 40 :ABCDEFGHIJKLMNOPQRSTU]WBC 5 DIFF Flags21 + 1Line 41 :LMMGGG]WBC LMG Flags6 + 1Line 42 :PSM]PLT Flags3 + 1Line 43 :CRC1				-	
Line 25:ZZZZZ-RN]RET*rejection & limits8 + 1Line 26:ZZZZZ-RN]RET*rejection & limits8 + 1Line 27:ZZZZZ-RN]RET*rejection & limits8 + 1Line 28:ZZZZZ-RN]RBCrejection & limits8 + 1Line 29:ZZZZZ-RN]HGBrejection & limits8 + 1Line 30:ZZZZZ-RN]HCTrejection & limits8 + 1Line 31:ZZZZZ-RN]HCTrejection & limits8 + 1Line 32:ZZZZZ-RN]MCHrejection & limits8 + 1Line 33:ZZZZZ-RN]MCHrejection & limits8 + 1Line 34:ZZZZZ-RN]RDWrejection & limits8 + 1Line 36:ZZZZZ-RN]RDWrejection & limits8 + 1Line 36:ZZZZZ-RN]PLTrejection & limits8 + 1Line 36:ZZZZZ-RN]PLTrejection & limits8 + 1Line 37:ZZZZZ-RN]PCTrejection & limits8 + 1Line 38:ZZZZZ-RN]PCTrejection & limits8 + 1Line 39:ZZZZZ-RN]PDWrejection & limits8 + 1Line 40:ABCDEFGHIJKLMNOPQRSTU]WBC 5 DIFF Flags21 + 1Line 41:LMMGGG]WBC 1MG Flags6 + 1Line 42:PSM]PLT Flags3 + 1Line 43:CRC1		•		-	
Line 26 :zzzzz-RN]RET* RET*rejection & limits8 + 1Line 27 :zzzzz-RN]RBCrejection & limits8 + 1Line 28 :zzzzz-RN]HGBrejection & limits8 + 1Line 29 :zzzzz-RN]HGBrejection & limits8 + 1Line 30 :zzzzz-RN]HCTrejection & limits8 + 1Line 31 :zzzzz-RN]MCVrejection & limits8 + 1Line 32 :zzzzz-RN]MCHrejection & limits8 + 1Line 33 :zzzzz-RN]MCHCrejection & limits8 + 1Line 34 :zzzzz-RN]RDWrejection & limits8 + 1Line 35 :zzzzz-RN]RET*rejection & limits8 + 1Line 36 :zzzzz-RN]PLTrejection & limits8 + 1Line 37 :zzzzz-RN]PCTrejection & limits8 + 1Line 38 :zzzzz-RN]PCTrejection & limits8 + 1Line 39 :zzzzz-RN]PDWrejection & limits8 + 1Line 40 :ABCDEFGHIJKLMNOPQRSTU]WBC 5 DIFF Flags21 + 1Line 41 :LMMGGG]WBC LMG Flags6 + 1Line 42 :PSM]PCTYBC LMG Flags6 + 1Line 43 :CRC11		-		-	
Line 27 :zzzzz-RN]RET*rejection & limits8 + 1Line 28 :zzzzz-RN]RBCrejection & limits8 + 1Line 29 :zzzzz-RN]HGBrejection & limits8 + 1Line 30 :zzzzz-RN]HCTrejection & limits8 + 1Line 31 :zzzzz-RN]MCVrejection & limits8 + 1Line 32 :zzzzz-RN]MCHrejection & limits8 + 1Line 33 :zzzzz-RN]MCHCrejection & limits8 + 1Line 34 :zzzzz-RN]RDWrejection & limits8 + 1Line 35 :zzzzz-RN]RDWrejection & limits8 + 1Line 36 :zzzzz-RN]PLTrejection & limits8 + 1Line 37 :zzzzz-RN]PCTrejection & limits8 + 1Line 38 :zzzzz-RN]PCTrejection & limits8 + 1Line 39 :zzzzz-RN]PDWrejection & limits8 + 1Line 41 :LMMGGG]WBC 5 DIFF Flags21 + 1Line 42 :PSM]WBC 5 DIFF Flags6 + 1Line 42 :PSM]Line 45 :3 + 1		-		•	
Line 28 :zzzzz-RN]RBCrejection & limits8 + 1Line 29 :zzzzz-RN]HGBrejection & limits8 + 1Line 30 :zzzzz-RN]HCTrejection & limits8 + 1Line 31 :zzzzz-RN]MCVrejection & limits8 + 1Line 32 :zzzzz-RN]MCHrejection & limits8 + 1Line 33 :zzzzz-RN]MCHCrejection & limits8 + 1Line 34 :zzzzz-RN]RDWrejection & limits8 + 1Line 35 :zzzzz-RN]RDWrejection & limits8 + 1Line 36 :zzzzz-RN]PLTrejection & limits8 + 1Line 37 :zzzzz-RN]PLTrejection & limits8 + 1Line 38 :zzzzz-RN]PCTrejection & limits8 + 1Line 39 :zzzzz-RN]PCTrejection & limits8 + 1Line 40 :ABCDEFGHIJKLIMNOPQRSTU]WBC 5 DIFF Flags21 + 1Line 41 :LMMGGG]WBC LMG Flags6 + 1Line 42 :PSM]PLT Flags3 + 1		•		-	
Line 29 :zzzzz-RN]HGBrejection & limits8 + 1Line 30 :zzzzz-RN]HCTrejection & limits8 + 1Line 31 :zzzzz-RN]MCVrejection & limits8 + 1Line 32 :zzzzz-RN]MCHrejection & limits8 + 1Line 33 :zzzzz-RN]MCHCrejection & limits8 + 1Line 34 :zzzzz-RN]MCHCrejection & limits8 + 1Line 35 :zzzzz-RN]RET*rejection & limits8 + 1Line 36 :zzzzz-RN]PLTrejection & limits8 + 1Line 37 :zzzzz-RN]MPVrejection & limits8 + 1Line 38 :zzzzz-RN]PCTrejection & limits8 + 1Line 39 :zzzzz-RN]PDWrejection & limits8 + 1Line 40 :ABCDEFGHIJKLMNOPQRSTUJWBC 5 DIFF Flags21 + 1Line 41 :LMMGGG]WBC 5 DIFF Flags6 + 1Line 42 :PSM]PLT Flags3 + 1					
Line 30 :zzzzz-RN]HCTrejection & limits8 + 1Line 31 :zzzzz-RN]MCVrejection & limits8 + 1Line 32 :zzzzz-RN]MCHrejection & limits8 + 1Line 33 :zzzzz-RN]MCHCrejection & limits8 + 1Line 34 :zzzzz-RN]RDWrejection & limits8 + 1Line 35 :zzzzz-RN]RET*rejection & limits8 + 1Line 36 :zzzzz-RN]PLTrejection & limits8 + 1Line 37 :zzzzz-RN]PCTrejection & limits8 + 1Line 38 :zzzzz-RN]PCTrejection & limits8 + 1Line 39 :zzzzz-RN]PCTrejection & limits8 + 1Line 40 :ABCDEFGHIJKLMNOPQRSTUJWBC 5 DIFF Flags21 + 1Line 41 :LMMGGG]WBC LMG Flags6 + 1Line 42 :PSM]PLT Flags3 + 1Line 43 :CRC1	Line 28 :	zzzz-RN]	RBC	rejection & limits	8 + 1
Line 31 : zzzzz-RN] MCV rejection & limits 8 + 1 Line 32 : zzzzz-RN] MCH rejection & limits 8 + 1 Line 33 : zzzzz-RN] MCHC rejection & limits 8 + 1 Line 34 : zzzzz-RN] RDW rejection & limits 8 + 1 Line 35 : zzzzz-RN] RET* rejection & limits 8 + 1 Line 36 : zzzzz-RN] PLT rejection & limits 8 + 1 Line 37 : zzzzz-RN] PLT rejection & limits 8 + 1 Line 38 : zzzzz-RN] PCT rejection & limits 8 + 1 Line 39 : zzzzz-RN] PCT rejection & limits 8 + 1 Line 39 : zzzzz-RN] PDW rejection & limits 8 + 1 Line 40 : ABCDEFGHIJKLMNOPQRSTUJ PDW rejection & limits 8 + 1 Line 41 : LMMGGG] Line 42 : PSM] NBC LMG Flags 6 + 1 PLT Flags 3 + 1 Line 43 : CRC 1	Line 29 :	zzzz-RN]	HGB	rejection & limits	8 + 1
Line 32 :zzzzz-RN]MCHrejection & limits8 + 1Line 33 :zzzzz-RN]MCHCrejection & limits8 + 1Line 34 :zzzzz-RN]RDWrejection & limits8 + 1Line 35 :zzzzz-RN]RET*rejection & limits8 + 1Line 36 :zzzzz-RN]PLTrejection & limits8 + 1Line 36 :zzzzz-RN]PLTrejection & limits8 + 1Line 37 :zzzzz-RN]PCTrejection & limits8 + 1Line 38 :zzzzz-RN]PCTrejection & limits8 + 1Line 39 :zzzzz-RN]PDWrejection & limits8 + 1Line 40 :ABCDEFGHIJKLMNOPQRSTU]WBC 5 DIFF Flags21 + 1Line 41 :LMMGGG]WBC LMG Flags6 + 1Line 42 :PSM]PLT Flags3 + 1	Line 30 :	zzzz-RN]	HCT	rejection & limits	8 + 1
Line 33 :zzzzz-RN]MCHC RDWrejection & limits8 + 1Line 34 :zzzzz-RN]RDWrejection & limits8 + 1Line 35 :zzzzz-RN]RET*rejection & limits8 + 1Line 36 :zzzzz-RN]PLTrejection & limits8 + 1Line 37 :zzzzz-RN]MPVrejection & limits8 + 1Line 38 :zzzzz-RN]PCTrejection & limits8 + 1Line 39 :zzzzz-RN]PCTrejection & limits8 + 1Line 40 :ABCDEFGHIJKLMNOPQRSTUJWBC 5 DIFF Flags21 + 1Line 41 :LMMGGG]WBC LMG Flags6 + 1Line 42 :PSMJPLT Flags3 + 1	Line 31 :	zzzz-RN]	MCV	rejection & limits	8 + 1
Line 34 :zzzzz-RN]RDW RET*rejection & limits rejection & limits8 + 1Line 35 :zzzzz-RN]PLT MPVrejection & limits rejection & limits8 + 1Line 36 :zzzzz-RN]PLT MPVrejection & limits rejection & limits8 + 1Line 37 :zzzzz-RN]PCT PCTrejection & limits rejection & limits8 + 1Line 38 :zzzzz-RN]PCT PDWrejection & limits rejection & limits8 + 1Line 39 :zzzzz-RN]PDWrejection & limits rejection & limits8 + 1Line 40 :ABCDEFGHIJKLMNOPQRSTUJ Line 41 :WBC 5 DIFF Flags VBC LMG Flags A + 121 + 1Line 42 :PSM]PLT Flags3 + 1Line 43 :CRC1	Line 32 :	zzzz-RN]	MCH	rejection & limits	8 + 1
Line 35 :zzzzz-RN]RET*rejection & limits8 + 1Line 36 :zzzzz-RN]PLTrejection & limits8 + 1Line 37 :zzzzz-RN]MPVrejection & limits8 + 1Line 38 :zzzzz-RN]PCTrejection & limits8 + 1Line 39 :zzzzz-RN]PDWrejection & limits8 + 1Line 40 :ABCDEFGHIJKLMNOPQRSTUJWBC 5 DIFF Flags21 + 1Line 41 :LMMGGGJWBC LMG Flags6 + 1Line 42 :PSMJPLT Flags3 + 1Line 43 :CRC1	Line 33 :	zzzz-RN]	MCHC	rejection & limits	8 + 1
Line 36 :zzzzz-RN]PLTrejection & limits8 + 1Line 37 :zzzzz-RN]MPVrejection & limits8 + 1Line 38 :zzzzz-RN]PCTrejection & limits8 + 1Line 39 :zzzzz-RN]PDWrejection & limits8 + 1Line 40 :ABCDEFGHIJKLMNOPQRSTUJWBC 5 DIFF Flags21 + 1Line 41 :LMMGGGJWBC LMG Flags6 + 1Line 42 :PSMJPLT Flags3 + 1	Line 34 :	zzzz-RN]	RDW	rejection & limits	8 + 1
Line 37 :zzzzz-RN]MPVrejection & limits8 + 1Line 38 :zzzzz-RN]PCTrejection & limits8 + 1Line 39 :zzzzz-RN]PDWrejection & limits8 + 1Line 40 :ABCDEFGHIJKLMNOPQRSTU]WBC 5 DIFF Flags21 + 1Line 41 :LMMGGG]WBC LMG Flags6 + 1Line 42 :PSM]PLT Flags3 + 1Line 43 :CRC1	Line 35 :	zzzz-RN]	RET*	rejection & limits	8 + 1
Line 37 :zzzzz-RN]MPVrejection & limits8 + 1Line 38 :zzzzz-RN]PCTrejection & limits8 + 1Line 39 :zzzzz-RN]PDWrejection & limits8 + 1Line 40 :ABCDEFGHIJKLMNOPQRSTU]WBC 5 DIFF Flags21 + 1Line 41 :LMMGGG]WBC LMG Flags6 + 1Line 42 :PSM]PLT Flags3 + 1Line 43 :CRC1	Line 20 i			vois stien 9 linsite	0 . 4
Line 38 :zzzzz-RN]PCT PDWrejection & limits rejection & limits8 + 1Line 39 :zzzzz-RN]PDWrejection & limits rejection & limits8 + 1Line 40 :ABCDEFGHIJKLMNOPQRSTU]WBC 5 DIFF Flags VBC LMG Flags21 + 1Line 41 :LMMGGG]WBC LMG Flags PLT Flags6 + 1Line 42 :PSM]PLT Flags3 + 1		-			
Line 39 :zzzzz-RN]PDWrejection & limits8 + 1Line 40 :ABCDEFGHIJKLMNOPQRSTU]WBC 5 DIFF Flags21 + 1Line 41 :LMMGGG]WBC LMG Flags6 + 1Line 42 :PSM]PLT Flags3 + 1Line 43 :CRC1		-		•	
Line 40 :ABCDEFGHIJKLMNOPQRSTU]WBC 5 DIFF Flags21 + 1Line 41 :LMMGGG]WBC LMG Flags6 + 1Line 42 :PSM]PLT Flags3 + 1Line 43 :CRC1		=		-	
Line 41 : LMMGGG] WBC LMG Flags 6 + 1 Line 42 : PSM] PLT Flags 3 + 1 Line 43 : CRC 1	Line 39 :	zzzz-RNJ	PDW	rejection & limits	8 + 1
Line 41 : LMMGGG] WBC LMG Flags 6 + 1 Line 42 : PSM] PLT Flags 3 + 1 Line 43 : CRC 1	Line 40 :	ABCDEFGHIJKLMNOPQRSTU]		WBC 5 DIFF Flags	21 + 1
Line 43 : CRC 1	Line 41 :	-		_	6 + 1
Line 43 : CRC 1		•		PLT Flags	
		-		-	
Line 44 : ETX (\$03) end of text 1	Line 43 :	CRC			1
	Line 44 :	ETX (\$03)		end of text	1

1.2.3. End of communication key

- (]) :	Carriage return \$0D

- CRC : the exclusive «OR» of all the transmitted bytes, except ETX and STX, then the inclusive «OR» with a \$40 value.
- zz : Number of the analyser.

2. STANDARD FORMAT

The **standard format** can have a different a different number of fields according to the transmitted items setup by the user (results, curves, flags, etc...).

2.1 Message Structure

STX

Size + carriage return.

Identifier followed by heading title + carriage return. Identifier followed by the Information associated to the heading title + carriage return. Remainder of the other Identifiers and Informations associated to the heading title + carriage returns.

Other heading titles + associated Informations

.....

Identifier followed by the Checksum + carriage return ETX

2.2. Details about the structure

Size :	5 bytes representing the total amount of the data except STX and ETX.
Heading title :	An 8 characters chain preceded by a space, indicating the associated data
	type.
Identifier :	1 byte (moving about \$21 to \$FF, it describes the information type which
	follows this indicator).
CheckSum :	Sum modulo 65535 of all the characters except ETX, STX and all the infor mations linked to the checksum (identifier - space - checksum) in the
	hexadecimal format on 4 bytes, preceded by a space.

2.3. Identifier list and their formats

2.3.1. Hematologyc numeric parameters

2.3.1.1. Format description

• Numerical field

For all indicated parameters from \$21 to \$43, the format is a numerical field of 5 digits completed with zeros on the left side (ex. : 04.55) and preceded by a space. The unit is the one chosen by the operator.

When the parameter cannot be calculated by the analyzer, the field is replaced with (--.--).

• Parameter status

Following the numerical field, a first digit gives the counting rejection status or the suspicion, a second one gives the parameter value status according to high and low normalities, and to the overloading capacities.

First digit (letter)	correspondance
R	Parameter rejected for a counting default
S	Suspicious parameter value
'space'	No anomaly observed
Second digit (letter)	correspondance
I	Parameter < to the low normal value
'space'	Parameter normal value
h	Parameter > to the high value
0	Parameter exceeding the capacity

• Example

5.5 millions RBC with a counting error in the standard units :

\$32 \$20 \$30 \$35 \$2E \$35 \$30 \$52 \$68 \$0D or **«2 05.5Rh»** + carriage return.

The length is fixed and is worth 2+7+1, that is to say 10 bytes for one parameter.

2.3.1.2. Identifier list

Ident	dentifiers Parameters		Units
\$21	!	WBC	Standard - SI g/dl - SI mmoles
\$22	"	Lymphocytes	(#)
\$23	#		(%)
\$24	\$	Monocytes	(#)
\$25	%		(%)
\$26	&	Granulocytes	(#)
\$27	1		(%)

Ident	ifiers	Parameters	Units
\$32	2	RBC	Standard - SI g/dl - SI mmoles
\$33	3	HCB	
\$34	4	НСТ	
\$35	5	MCV	
\$36	6	МСН	
\$37	7	МСНС	
\$38	8	RDW	

Ident	ifiers	Parameters	Units
\$40	0	PLT	Standard - SI g/dl - SI mmoles
\$41	А	MPV	
\$42	В	ТНТ	
\$43	С	PDW	

2.3.2. Pathology

* Flags associated with parameters

• Format description

Flags are transmitted in a comprehensive mode, preceded by a space (same presentation than on the screen, i.e. dependant from the language) 2 characters which are replaced with spaces when the flag has not been detected.

• Identifier list

Ident	tifiers	Parameters	Formats	Length
\$50	Р	WBC or LMG	L1M1M2G1G2G3	2 + 12 + 1
\$53	S	PLT	PcScMc	2 + 6 + 1

2.3.3. Histograms and matrix

2.3.3.1. Format description

• Histograms

Histograms are transmitted on 128 or 256 channels, preceded by a space. They are automatically rescaled to a 223 maximum amplitude value.

The zero amplitude value is \$20, the maximum amplitude value is \$FF.

2.3.3.2. Identifier list

Identi	ifiers	Parameters	Formats	Length
\$57	W	WBC	Amplitude of each channel	2 + 128 + 1
\$58	х	RBC	n	"
\$59	Y	PLT	n	"
\$5D	1	WBC thresholds	5 thresholds	1 + 20 + 1
\$5E	^	RBC thresholds	2 thresholds	1 + 8 + 1
\$5F	_	PLT thresholds	1 threshold	1 + 4 + 1

2.3.4. Patient result identification

2.3.4.1. Format description

All the described fields have a fixed size character chain type and are completed with spaces for the non significant informations.

Iden	tifiers	Correspondance	Formats	Length
\$70	р	Analyzer number	01	2 + 2 + 1
\$71	q	Analysis date and time	94/06/06 13h15mn31s	2 + 19 + 1
\$73	S	Analyzer sequence number	0128	2 + 4 + 1
\$74	t	Sampling mode	'O' : open tube 'C' : close tube	2 + 1 + 1
\$75	u	Identification number	1450302154275-42	2 + 16 + 1
\$76	v	Identification	SMITH Ronald	2 + 30 + 1
\$80	ç	Analysis type	Defined on 1 character (see description) 'A' CBC analysis 'D' LMG analysis From 'G' to 'Z' can be configurated by the user	2 + 1 + 1

2.3.4.2. Identifier list

2.3.4.3. Analysis type (\$80)

This identifier defines the analysis type CBC, LMG to carried out on the sample. It also provides the analysis of one or several specific parameters.

The CBC analysis includes the 12 parameters of the CBC's count.

The LMG analysis includes the CBC analysis and the % and # of the 6 WBC populations.

2.4. Heading title

The heading title provides the data of the whole message : current hematological results or results coming from statistics. This heading title is able to drive commands that can be interpreted by the analyser or by an external computer.

Analyzers being able to communicate in the bidirectionnal mode and supporting the remote control mode, can interprete the heading title and runs the corresponding actions.

This string is a 8 characters length, preceded by a space, containing data that follows, or the command type to be carried out. The identifier is \$FF.

Iden	tifier	Correspondance	Format	Length
\$FF		type of data packet	String of characters	2 + 8 + 1

Data exported by the analyzer

Data packet string	Use
RESULT	Hematological result transmission on a routine mode

2.5. Other identifiers

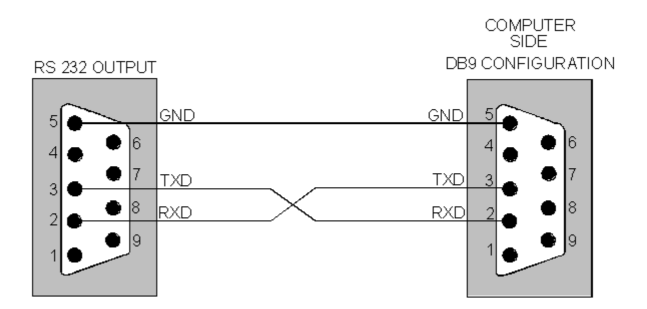
Ident	ifiers	Correspondance	Format	Length
\$FC		Number	On 8 bytes	2 + 8 + 1
\$FD		Checksum value	hexadecimal on 4 Bytes	2 + 4 + 1

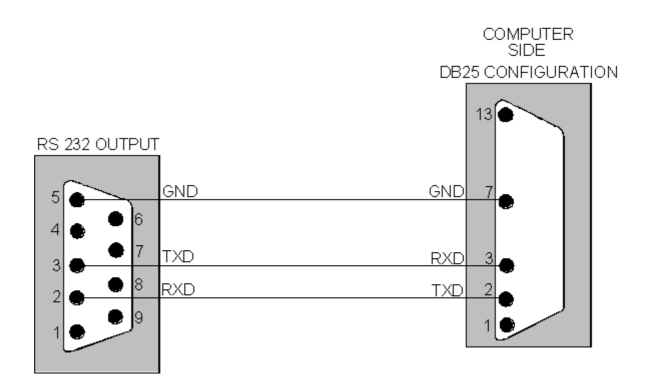
Identifier \$FC : This identifier allows the number transmission that could be an error #, a position #, a burn-in sequence # or an hexadecimal status (see "Error list"). Identifier \$FD : Checksum value : see chapter on the message constitution.

Error list

Error N°	event linked to the analyzer
1	Operating temperature out of limits

3. PIN ASSIGNMENTS





RAA 009 A Ind.B

5. TRAINING SLIDES