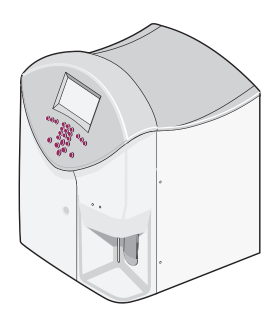
# BECKMAN COULTER™ A<sup>C</sup>•T™ 5diff Hematology Analyzer

# **Service Manual**





PN 4237616B (September 2000) COULTER CORPORATION A Beckman Coulter Company Miami, Florida 33196-2500 USA

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Initial Issue, 3/2000 Released by CN 040130-0003 Software Version 0.11

**Issue B, 07/00** Released by CN 040150-0025 Software version 1.03

The material in the revision B change pages was updated for software version 1.03 and for any hardware changes since revision A. The changes include updating the adjustment procedures for the bath assembly, HGB blank, RBC/PLT gain, WBC/BASO, motor current, thresholds, and the optical bench; updating the replacement procedures for the heater assembly, power supply, start switch, reagent syringes, count syringe, sample prove, waste syringe, 5diff syringe, flow-cell coax, optical bench lamp, diluent reservoir, and sample syringe; updating the procedure for testing and configuring the bar-code reader; updating the parts lists; updating the tubing lists and associated circuit connections; adding procedures for balancing the WBC count, setting the diff+/diff- thresholds, and replacing the new Main card; and adding information on the LX300 + printer.

Changes were made on the following pages:

1.1-1, 1.1-2, 1.1-4 2.1-4, 2.6-1, 2.6-2, 2.6-3, 2.6-4, 2.6-5, 2.8-3, 2.8-11, 2.8-12, 2.8-13, 2.8-15, 2.10-2, 2.10-3 3.1-1, 3.2-1, 3.2-2, 3.2-3, 3.2-6, 3.2-7, 3.2-9, 3.2-10, 3.3-1, 3.3-7, 3.3-8 added 3.3-9 and 3.3-10 4.1-1, 4.1-2, 4.1-3, 4.1-4, 4.2-1, 4.2-2, 4.2-4, 4.2-5, 4.2-6 through 4.2-8, 4.4-1, 4.5-1 through 4.5-4 4.6-1 through 4.6-8, 4.7-2 4.8-1, 4.9-1, 4.9-2, 4.10-1, 4.10-2, 4.11-1, 4.12-1, 4.12-2, 4.13-1; 4.13-2 4.14-1, 4.14-2, 4.17-2, 4.17-3, 4.18-1 4.19-1 through 4.19-8, 4.20-1, 4.20-2, 4.20-3, 4.20-4 deleted 4.20-5 and 4.20-6 4.21-1 through 4.21-3, 4.23-1 through 4.23-6, 4.24-1 through 4.24-6, 4.25-1 through 4.25-8 deleted 4.25-9 through 4.25-12 4.26-1, 4.26-3, 4.26-4, 4.27-1 through 4.27-8 deleted 4.27-9 through 4.27-12 4.29-1, 4.29-3 through 4.29-6, 4.30-1, 4.30-3 through 4.30-5, 4.31-1, 4.31-3, 4.31-4 4.32-1 through 4.32-3, 4.33-1, 4.33-2, 4.35-1, 4.35-2, 4.35-3, 4.35-10, 4.36-1, 4.36-2, 4.36-5, 4.36-6 added 4.37-1 and 4.37-2, 4.38-1 and 4.38-2, 4.39-1 through 4.39-10 5.2-3, 6.3-2 through 6.3-6, 7.3-2 8.1-1 through 8.1-12, 8.2-7, 8.2-8, 8.2-9, 8.2-16, 8.2-17, 8.2-42, 8.2-46, 8.2-48 A.2-9, A.3-3, A.4-1, C.1-1, C.1-2 and C.1-7.

The change page packet also includes the latest revision of the Pneumatic/Hyraulic Schematic, 7616069B.

Changes that are part of the most recent revision are indicated in the printed copy by a bar in the margin of the amended page.

This document applies to the latest software listed and higher versions. When a subsequent software version affects the information in this document, the changes will be included on minor revision change pages or summarized on a Notice of Information Update form and will be released by service memo.

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# 1.1 MANUAL DESCRIPTION

# Scope

This manual provides the reference information and procedures needed for servicing and maintaining the BECKMAN COULTER<sup>TM</sup>  $A^{C} \bullet T^{TM}$  5diff hematology analyzer (hereafter referred to as the  $A^{C} \bullet T$  5diff hematology analyzer or the instrument). It is available both online and in hard copy. The online manual is released on the Service Resource Kit CD-ROM, PN 6417471.

This manual is to be used in conjunction with the following customer documents and does not contain information and procedures already covered in these documents:

| Document                        | Language | Part Number |
|---------------------------------|----------|-------------|
| Operator's Guide                | English  | 4237615     |
|                                 | French   | 4237630     |
|                                 | Italian  | 4237631     |
|                                 | German   | 4237632     |
|                                 | Spanish  | 4237633     |
|                                 | Chinese  | 4237634     |
| Host Transmission Specification | English  | 4277065     |

# **Notification of Updates**

Any service memo that affects the information in this manual will include either minor revision change pages or a Notice of Information Update form for this manual. A Notice of Information Update form will summarize the changes and will list the specific headings, figures, and tables affected.

# **Intended Audience**

To use this manual effectively, you need the following:

- An operator's knowledge of the A<sup>C</sup>•T 5diff hematology analyzer
- A thorough understanding of -
  - Basic electronic and pneumatic principles and devices
  - Reagent systems
  - Quality control
  - Troubleshooting concepts
- The ability to -
  - Use basic mechanical tools and understand related terminology
  - Use a digital voltmeter (DVM)
  - Read pneumatic/hydraulic schematics and understand related terminology
  - Read electronic schematics and understand related terminology

# Organization

The material in this manual is organized into eight chapters and two appendices. To make it easier to access the information:

I

- In the online manual, each page has a Contents button linked to a master table of contents and an Index button linked to an alphabetic index.
- In the printed manual, there is a master table of contents at the beginning of the manual, a chapter-specific table of contents at the beginning of each chapter, and an alphabetic index at the end of the manual.

The chapters / appendices contain:

**Chapter 1, INTRODUCTION -** A brief description of this manual and essential safety information.

**Chapter 2**, **INSTRUMENT DESCRIPTION** - An introduction to the A<sup>C</sup>•T5 diff hematology analyzer and a description of how it functions.

Chapter 3, INSTALLATION PROCEDURES - Installation and verification procedures.

**Chapter 4, SERVICE AND REPAIR PROCEDURES** - The procedures for servicing/repairing the A<sup>C</sup>•T 5diff hematology analyzer.

**Chapter 5**, **MAINTENANCE PROCEDURES** - The procedures for maintaining the A<sup>C</sup>•T 5diff hematology analyzer.

Chapter 6, SCHEMATICS - The schematic diagrams and tubing lists.

Chapter 7, TROUBLESHOOTING - An error message table.

Chapter 8, PARTS LISTS - The master parts list followed by the illustrated parts list.

**Appendix A, QUICK REFERENCE INFORMATION -** Quick reference information: tolerances and limits; connectors, test points and jumpers for the circuit cards; the software menu trees; location diagrams and summarized functions for main analyzer components.

Appendix B, SOFTWARE INTERFACE - Tables of fatal and non-fatal error messages.

Appendix C, FLAG SENSITIVITY AND THRESHOLDS - An overview of the theory including default values; also includes the setup procedures.

**ABBREVIATIONS** - A list of abbreviations, acronyms, and reference designators used in this manual.

## **Numbering Format**

Each chapter of this manual is further divided into topics that are numbered sequentially, beginning at one. The numbering format for the topic heading, which is called the primary heading, is chapter number, decimal point, topic number. For example, the primary heading number for the third topic covered in Chapter 2 is 2.3.

The page, figure, and table numbers are tied directly to the primary heading number. For example, Heading 2.3 begins on page 2.3-1, the first figure under Heading 2.3 is Figure 2.3-1 and the first table under Heading 2.3 is Table 2.3-1.

Note: Primary headings always begin at the top of a right-hand page.

## **Special Headings**

Throughout this manual, WARNING, CAUTION, IMPORTANT, ATTENTION, and Note headings are used to indicate potentially hazardous situations and important or helpful information.

## WARNING

A WARNING indicates a situation or procedure that, if ignored, can cause serious personal injury. The word WARNING is in bold-faced text in the printed manual and is red in the online manual.

## CAUTION

A CAUTION indicates a situation or procedure that, if ignored, can cause damage to the instrument. The word CAUTION is in bold-faced text in the printed manual and is red in the online manual.

## IMPORTANT

An IMPORTANT indicates a situation or procedure that, if ignored, can result in erroneous test results. The word IMPORTANT is in bold-faced text in the printed manual and is red in the online manual.

## ATTENTION

An ATTENTION contains information that is critical for the successful completion of a procedure and/or operation of the instrument. The word ATTENTION is in bold-faced text in the printed manual and is red in the online manual.

## Note

A Note contains information that is important to remember or helpful in performing a procedure.

## Conventions

This manual uses the following conventions.

- Instrument or analyzer refers to the A<sup>C</sup>•T 5diff hematology analyzer.
- Main card refers to the motherboard in the instrument.
- Main Menu refers to the initial menu displayed on the instrument after Startup.
- Each menu option consists of an item number followed by bold, uppercase text. For example, **3. REAGENTS** is the third option on the Main Menu.

**Note:** Both the menu item number and text are displayed on the LCD screen. The item number next to the menu item indicates the numeric pushbutton on the front of the analyzer that can be pressed to select the menu option.

• Keys on the analyzer keypad are in bold, uppercase letters. For example, press **ENTER** indicates the operator should press the **ENTER** pushbutton on the instrument keypad.

- To select a menu item,
  - Use the arrow keys to highlight the desired menu item then press the ENTER pushbutton on the front of the analyzer to select the highlighted option.
     or
  - Simply press the numeric pushbutton (on the front of the analyzer) that correlates with the desired option. This is the faster way to select a menu item.

For example, to select the **3**. **REAGENTS** menu item from the Main Menu, you may:

• Use the down arrow to highlight the **3**. **REAGENTS** option and then press the **ENTER** pushbutton on the front of the analyzer.

or

- Press the pushbutton labeled **3** on the front of the analyzer.
- Select menu item **>>** sub-menu item indicates the software options you have to select, as well as the order in which you should select them. For example, to prime the diluent reagent:

From the Main Menu, select 3. REAGENTS >> 3. PRIME >> 1. DILUENT.

- Italics us used to indicate screen messages. For example: The message *CYCLE IN PROGRESS. PLEASE WAIT* . . . appears on the screen.
- A<sup>C</sup>•T 5diff Rinse reagent is sometimes referred to as Rinse.
- A<sup>C</sup>•T 5diff Fix reagent is sometimes referred to as Fix.
- A<sup>C</sup>•T 5diff Hgb Lyse reagent is sometimes referred to as Hgb Lyse.
- A<sup>C</sup>•T 5diff WBC Lyse reagent is sometimes referred to as WBC Lyse.
- A<sup>C</sup>•T 5diff Diluent reagent is sometimes referred to as Diluent or diluent.
- In the electronic version of the manual:
  - Links to additional information are in blue and are underlined. To access the linked information, select the blue underlined text.
  - The material is divided into many small sections (electronic files) to enhance the loading and accessibility features.
  - Every primary heading is a separate file and whenever possible the amount of material contained within one primary heading is limited to four to ten pages.
  - If a primary heading must be large, such as an illustrated parts list (IPL), invisible breaks are added to the electronic file to further divide it.

Note: Unless you are scrolling, these divisions are invisible. If you choose to scroll through the IPL, you will encounter stop points. When you scroll to the end of a section and encounter a stop point, use the navigation bar to access the next section.

 To move from one section (electronic file) to the next in the HTML version of the manual, use the right and left arrows on the navigation bars displayed at the top and bottom of each section.

# Graphics

All graphics, including screens and printouts, are for illustration purposes only and must not be used for any other purpose.

## 1.2 SAFETY PRECAUTIONS

#### Electronic

**WARNING** Risk of personal injury. Contacting exposed electronic components while the instrument is attached to power can cause personal injury from electric shock. Power down completely before removing covers to access electronic components.

**WARNING** Risk of personal injury or damage to electronic components. While performing maintenance or service on the instrument, rings and other metal jewelry can become caught in the instrument. To avoid personal injury or damage to the instrument, remove rings and other metal jewelry before performing maintenance or service on the electronic components of the instrument.

**CAUTION** Risk of damage to electronic components. If the power is ON while removing or replacing printed circuit cards and components, the instrument could be damaged. To prevent damage to electronic components, always be sure power is OFF before removing or replacing printed circuit cards and components.

**CAUTION** Risk of damage to electronic components. Electrostatic discharge (ESD) can damage add-in circuit cards and other electronic components. If there is a possibility of ESD damage with a procedure, then perform that procedure at an ESD workstation, or wear an antistatic wrist strap attached to a metal part of the chassis connected to an earth ground.

## Biological

**WARNING** Risk of personal injury or contamination. If you do not properly shield yourself while servicing the instrument with the doors open, you may become injured or contaminated. To prevent possible injury or biological contamination, you must wear appropriate safety glasses, a lab coat, and gloves when servicing the instrument with the doors open.

Use care when working with pathogenic materials. Means must be available to decontaminate the instrument, provide ventilation, and to dispose of waste liquid. Refer to the following publications for further guidance on decontamination:

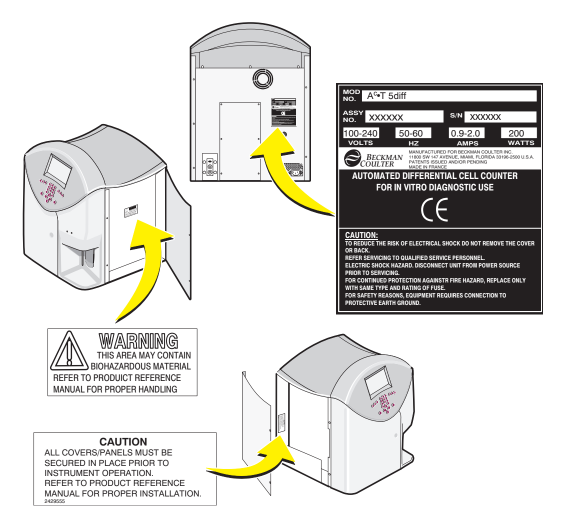
- Biohazards Safety Guide, 1974, National Institute of Health.
- Classifications of Etiological Agents on the Basis of Hazards, 3d ed., June 1974, Center for Disease Control, U.S. Public Health Service.

#### Troubleshooting

Bring the following Warning to the customer's attention before advising that customer to perform any service, maintenance or troubleshooting procedures on the A<sup>C</sup>•T 5diff hematology analyzer. Also, make sure customers are aware of the Warning and information labels shown in Figure 1.2-1.

**WARNING** Risk of personal injury or contamination. If you do not properly shield yourself while performing service, maintenance, and troubleshooting procedures, residual fluids in the instrument could injure or contaminate you. Beckman Coulter recommends that you wear barrier protection, such as appropriate safety glasses, a lab coat, and gloves throughout the performance of service, maintenance, and troubleshooting procedures and residual fluids in the instrument.

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## **INSTRUMENT DESCRIPTION**

# 2.1 INTRODUCTION TO THE A<sup>C</sup>•T 5diff HEMATOLOGY ANALYZER

## Purpose

The purpose of the A<sup>C</sup>•T 5diff hematology analyzer is to identify normal patient results with all normal system-generated parameters and to flag or identify patient results that require additional studies.

## Function

The  $A^{C} \bullet T$  5diff analyzer is a quantitative, fully automated (microprocessor controlled) hematology analyzer and leukocyte differential counter For In Vitro Diagnostic Use in clinical laboratories. The  $A^{C} \bullet T$  5diff hematology analyzer reports a complete blood count (CBC) and white blood cell differential (DIFF) on open-vial, whole-blood specimens.

The CBC consists of white blood cell count (WBC), red blood cell count (RBC), hemoglobin (Hgb), hematocrit (Hct), mean cell volume (MCV), mean cell hemoglobin (MCH), mean cell hemoglobin concentration (MCHC), red cell distribution width (RDW), platelet count (Plt), and mean platelet volume (MPV).

The DIFF (a 5-part leukocyte differential) consists of the percentage (%) and absolute number (#) of the following WBC populations: neutrophils (NE% and NE#), lymphocytes (LY% and LY#), monocytes (MO% and MO#), eosinophils (EO% and EO#), and basophils (BA% and BA#).

Six parameters are qualitative and are For Research Use Only. Not For In Vitro Diagnostic Procedures. These parameters include the plateletcrit (Pct), platelet distribution width (PDW), percentage and absolute number of immature cells (IMM% and IMM#), and percentage and absolute number of atypical lymphocytes (ATL% and ATL#).

## Description

## Components

The A<sup>C</sup>•T 5diff hematology analyzer is contained in one unit, with additional space needed only for the external printer, the diluent reagent container, and a waste container (if used).

## Interaction with the AC•T 5diff Hematology Analyzer

The A<sup>C</sup>•T 5diff analyzer uses an Open-Vial mode of operation. Pressing the aspirate switch (Figure 2.1-1) initiates a cycle. When the sample probe is submerged in a whole-blood specimen and the aspirate switch is pressed, sample is pulled from the specimen tube into the sample probe. As the cycle continues, the instrument then dilutes and analyzes this sample. When the analysis is complete, results appear on a LCD (Figure 2.1-1) and are available to the printer.

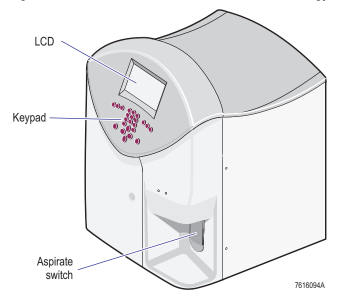


Figure 2.1-1 User Interfaces on the A<sup>C</sup>•T 5diff Hematology Analyzer

You also interact with the instrument through the use of a menu system displayed on a 128 by 240 pixels LCD and a control panel keypad with buttons that are used to setup and operate the instrument (Figure 2.1-1).

Since most input/output functions of the operating system software are controlled by the user, the pushbutton keypad and LCD screen are particularly important because they provide the physical user interface with the software.

See Heading 2.10, SOFTWARE STRUCTURE for more specific information as well as a graphic representation of the available menus and menu items (or options).

# 2

## **Modes of Operation**

The A<sup>C</sup>•T 5diff hematology analyzer has two operating modes: CBC and CBC/DIFF.

#### **CBC Mode**

Twelve parameters are generated in the CBC mode of operation - 10 parameters For In Vitro Diagnostic Use and two qualitative parameters that are For Research Use Only. Not For In Vitro Diagnostic Procedures:

| Parameters<br>(For Research Use Only) |
|---------------------------------------|
| Pct                                   |
| PDW                                   |
|                                       |
|                                       |
|                                       |
|                                       |
|                                       |
|                                       |
|                                       |
|                                       |
|                                       |

#### **CBC/DIFF Mode**

26 parameters are generated in the CBC/DIFF mode of operation - 20 parameters For In Vitro Diagnostic Use and six qualitative parameters that are For Research Use Only. Not For In Vitro Diagnostic Procedures:

| Parameters<br>(For In Vitro Diagnostic Use) | Parameters<br>(For Research Use Only) |
|---|---------------------------------------|
| WBC   | Pct                                   |
| RBC   | PDW                                   |
| Hgb   | IMM% and IMM#                         |
| Hct   | ATL% and ATL#                         |
| MCV   |                                       |
| MCH   |                                       |
| MCHC  |                                       |
| RDW   |                                       |
| Plt   |                                       |
| MPV   |                                       |
| NE% and NE#                                 |                                       |
| LY% and LY#                                 |                                       |
| MO% and MO#                                 |                                       |
| EO% and EO#                                 |                                       |
| BA% and BA#                                 |                                       |
|   |                                       |

## **Reagent Consumption**

I

Table 2.1-1 shows the instrument reagent consumption by cycle.

| Cycle                | A <sup>C</sup> •T 5diff reagents with usage per cycle |          |          | Approximate Duration |          |                      |
|----------------------|---|----------|----------|----------------------|----------|----------------------|
|                      | Diluent   | WBC Lyse | Rinse    | Fix                  | Hgb Lyse |                      |
| CBC                  | 20.5 mL   | 2.1 mL   | 0.9 mL   | Not used             | 0.4 mL   | 1 minute             |
| CBC/DIFF             | 25.6 mL   | 2.1 mL   | 0.9 mL   | 1.0 mL               | 0.4 mL   | 1 minute             |
| Startup <sup>†</sup> | 62.0 mL   | 2.1 mL   | 3.7 mL   | 1.0 mL               | 1.4 mL   | 3 minutes 40 seconds |
| Shutdown             | 25.5 mL   | Not used | 14.0 mL  | Not used             | 1.0 mL   | 2 minutes 45 seconds |
| Prime diluent        | 35.5 mL   | Not used | Not used | Not used             | Not used | 2 minutes 30 seconds |
| Prime rinse          | Not used  | Not used | 25.8 mL  | Not used             | Not used | 1 minute 20 seconds  |
| Prime fix            | Not used  | Not used | Not used | 25.8 mL              | Not used | 1 minute 30 seconds  |
| Prime WBC Lyse       | Not used  | 25.8 mL  | Not used | Not used             | Not used | 1 minute 20 seconds  |
| Prime Hgb Lyse       | 2.5 mL  | Not used | Not used | Not used             | 4.2 mL   | 1 minute             |
| Prime All Reagents   | 23.7 mL   | 16.0 mL  | 16.0 mL  | 16.0 mL              | 4.2 mL   | 3 minutes 20 seconds |
| Extended Cleaning    | 12.5 mL   | Not used | 6.0 mL   | Not used             | Not used | 1 minute 35 seconds  |
| System Reset Cycle   | 24.0 mL   | Not used | 1.4 mL   | Not used             | 1.0 mL   | 1 minute 25 seconds  |

<sup>†</sup> For one background count only. The maximum is three.

## 2.2 OPERATION PRINCIPLES

#### Overview

The A<sup>C</sup>•T 5diff analyzer is a fully automated hematology analyzer providing a complete WBC five-part differential, which is determined simultaneously by the A<sup>C</sup>V Technology (Absorbance Cytochemistry and Volume Technology) and the white blood cell/basophil (WBC/BASO) methodologies.

The A<sup>C</sup>V Technology uses absorbance, cytochemistry, and focused flow impedance. The WBC/BASO methodology uses differential lysis, impedance technology, and differential thresholds. See Table 2.2-1.

| Fluid Dynamics    | Technology  | Measurements   | Output  |
|-------------------|---|--|---|
| Dual Focused Flow | A <sup>C</sup> V Technology                       | Light absorbance of<br>cytochemically-stained<br>cells | Lymphocytes, monocytes,<br>neutrophils, eosinophils,<br>immature cells, and<br>atypical lymphocytes |
| Volume aperture   | Differential lysis using the<br>Coulter Principle | Volume and count                                       | WBC count, basophil<br>percentage, and basophil<br>count  |
| Volume aperture   | Coulter Principle                                 | Volume and count                                       | RBC count, platelet count, and hematocrit   |

Table 2.2-1 AC•T 5diff Analyzer Measurement Technologies

## **Measurement Principles**

## **Coulter Principle**

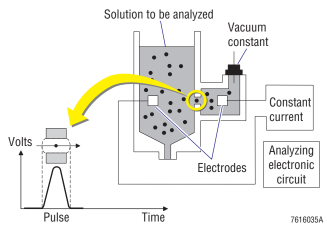
In the A<sup>C</sup>•T 5diff analyzer, the Coulter Principle is used to analyze the final red blood cell and platelet (RBC/Plt) dilution and the WBC/BASO dilution. This electronic method of counting and sizing particles is based on the fact that cells, which are poor conductors of electricity, will interrupt a current flow. The impedance variation generated by the passage of nonconductive cells through a small, calibrated aperture is used to determine the count (number of particles) and size (volume) of the particles passing through the aperture within a given time period.

## **Aperture Sensor System**

The RBC/Plt aperture sensor system determines the cell count and size of red blood cells and platelets. The WBC/BASO aperture sensor system determines the cell count and size of white blood cells. Additionally, the differentiation between basophils and other white blood cells is related to the A<sup>C</sup>•T 5diff WBC Lyse-specific lytic action on the white blood cells in the WBC/BASO bath.

To sense particles using the Coulter Principle (Figure 2.2-1), a current flow is established so changes in that flow can be monitored. In this sensing system, an electrode is placed on each side of the aperture (Figure 2.2-1). The most visible electrode is referred to as the counting head. These electrodes are the conductive metallic housings attached to the front of the RBC and WBC/BASO baths. The second electrode, referred to as the bath electrode, is not as conspicuous. This electrode is located inside the bath. The aperture is located between the counting head and the bath electrode.

#### Figure 2.2-1 Coulter Principle



When the count circuit is activated and an electronically conductive reagent is in the RBC or WBC/BASO bath, an electric current continuously passes through the aperture. Current moving between the two electrodes establishes the electronic flow through the aperture.

Once a sample is aspirated, an aliquot of that aspirated sample is diluted with reagent (an electrolyte) and is delivered to the RBC or WBC/BASO bath using tangential flow, which ensures proper mixing of the dilution. When the cells suspended in the conductive reagent are pulled through a calibrated aperture, the electrical resistance between the two electrodes increases proportionately with the cell volume (Figure 2.2-1).

The resistance creates a pulse that is sensed and counted as a particle by the instrument. The amount of resistance (amplitude of each pulse) is directly related to the size of the particle that produced it.

The generated pulses have a very low voltage, which the amplification circuit increases so that the electronic system can better analyze the pulses and eliminate the background noise.

## **Applying the Coulter Principle**

The  $A^{C} \cdot T$  5diff analyzer makes several dilutions of an aspirated whole-blood sample. The RBC/Plt dilution begins in the DIL1/HGB (first dilution/hemoglobin) bath but is actually analyzed in the RBC bath. The final dilution in the RBC bath is used to determine the cell count and size of red blood cells and platelets.

The WBC/BASO aperture sensor system is directly responsible for determining the cell count and size of white blood cells. The differentiation between basophils and other white blood cells is also related to the A<sup>C</sup>•T 5diff WBC Lyse-specific lytic action on these white blood cells.

Thresholds, which are electronically set size limits, exclude unwanted particles, such as debris, from the analysis. Particles above the threshold are analyzed, and particles below the threshold are excluded.

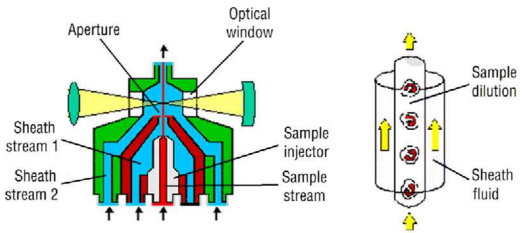
## A<sup>C</sup>V Technology

In the DIFF (differential) bath, 25  $\mu$ L of whole blood is mixed with 1,000  $\mu$ L of A<sup>C</sup>•T 5diff Fix reagent for 12 seconds, then stabilized with 1,000  $\mu$ L of A<sup>C</sup>•T 5diff Diluent for an additional three seconds. This reaction lyses the red blood cells, preserves the leukocytes at their original size, and differentially stains the lymphocytes, monocytes, neutrophils, and eosinophils, with eosinophils staining most intensely. The instrument maintains the reagents and reaction at a regulated temperature of 35°C (95°F).

Lymphocytes, monocytes, neutrophils, and eosinophils each have a unique nuclear and morphology structure and staining intensity; therefore, each cell type absorbs light differently. Each stained cell is individually focused by the Dual Focused Flow (DFF) system and transported through the flow cell using sample pressure and diluent sheath flow.

## **Dual Focused Flow (DFF)**

DFF fluid dynamics uses a hydrodynamic focusing process to focus individual cells or particles in a stream of diluent (Figure 2.2-2). The focused sample stream of the  $A^{C} \cdot T$  5diff analyzer is about 40 µm in diameter.



#### Figure 2.2-2 Dual Focused Flow Process

DFF uses sheath fluid to surround and force cells suspended in diluent to pass one at a time through the center of the flow cell. The first sheath flow focuses the sample through the impedance aperture. The second sheath flow maintains the focused flow of cells as they exit the aperture into the optical flow cell. Hydrodynamic focusing in the flow cell enables accurate and rapid cell-by-cell measurements on a large number of individual cells.

## **Flow Cell**

Sequential analyses for cell volume (impedance) and light absorbance are performed in the flow cell. A total of 72  $\mu$ L of sample is injected through the flow cell for 15 seconds. The flow cell incorporates a 60  $\mu$ m aperture for cellular volume analysis and about a 40  $\mu$ m measurement area for light absorbance.

## **Focused Flow Impedance**

Focused flow impedance technology measures the electrical resistance of a cell as it passes through the aperture in the flow cell. The change in resistance is directly proportional to the volume of the cell.

## **Absorbance Cytochemistry**

As a cell passes through the optical portion of the flow cell, light is scattered in all directions. A sensor detects only forward scattered light. The optical measurement is derived as a function of the amount of light lost due to diffraction and absorbance, as compared to full transmission when no cell is present.

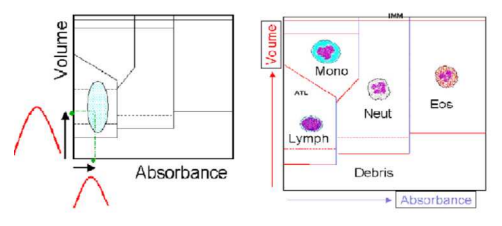
The collected signals are converted into voltage pulses and are processed. The magnitude of the voltage pulses are proportional to the physical and chemical characteristics of the cells being analyzed. Light absorbance is related to cellular contents (granularity, nuclear content, and so forth) after cytochemical staining. These measurements provide the information for lymphocytes, monocytes, neutrophils, and eosinophils, and their precursors.

#### **Signal Processing**

The signals from the flow cell aperture and from the optical measurement are correlated by a window of time. The optical pulse must be detected within 100 to 300 microseconds of the impedance pulse; otherwise, the signal is rejected.

The output signals from the focused flow impedance and the light absorbance measurements are combined to define the WBC differential population clusters. See Figure 2.2-3.





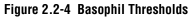
## Thresholds

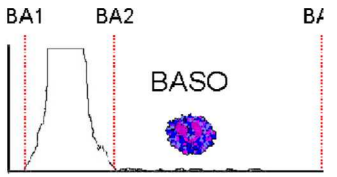
Most of the population partition thresholds are fixed and give the limits of the morphological normality of leukocytes. Changes in the morphology of a population are expressed on the DiffPlot by a shifting of the corresponding population. Volume and absorbance thresholds are used to detect shifting populations.

## WBC/BASO Methodology

In the WBC/BASO bath, 10  $\mu$ L of whole blood is mixed with 2,000  $\mu$ L of A<sup>C</sup>•T 5diff WBC Lyse reagent. This reaction lyses the red blood cells and specifically differentiates between basophils and other leukocytes by volume. The instrument maintains the reagents and reaction at a regulated temperature of 35°C (95°F).

Using a constant vacuum, the instrument then pulls the sample through an 80 µm aperture. As each cell passes through the aperture, a pulse is generated proportional to the cellular volume. The total leukocyte count and basophil percentage are determined by specific thresholds on the WBC/BASO histogram (Figure 2.2-4).





## **Sample Analysis Overview**

## Aspiration

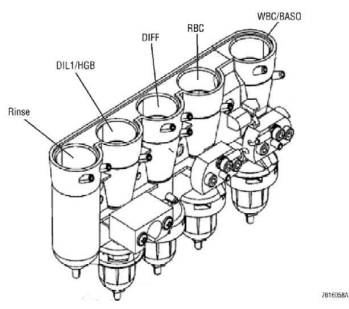
When the sample probe is immersed in a whole-blood specimen and the aspirate switch is pressed, sample is pulled from the tube into the sample probe. Depending on the selected mode of operation, the  $A^{C} \cdot T$  5diff analyzer aspirates either 30 µL (CBC mode) or 53 µL (CBC/DIFF mode) of sample.

The volume of sample aspirated into the sample probe is sufficient to make all the dilutions needed to develop parameter results in the selected mode of operation.

## Dilution

Using the Sequential Dilution System (SDS) technique, the aspirated sample is partitioned as it is distributed to make a series of dilutions in a series of baths (Figure 2.2-5).

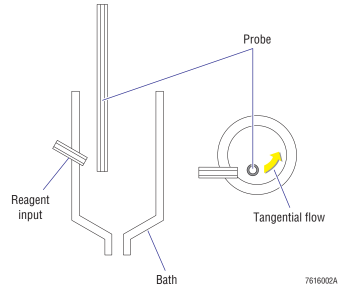
## Figure 2.2-5 Bath Assembly



## Delivery

In the CBC and the CBC/DIFF modes, each aliquotted sample is delivered to its appropriate bath using a tangential flow (Figure 2.2-6) of reagent, which mixes the diluted sample and minimizes viscosity problems.

## Figure 2.2-6 Sample Delivery Using Tangential Flow



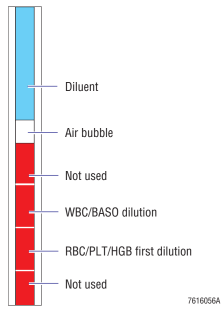
## **Sample Partitioning**

Figure 2.2-7 shows the sample partitioning that occurs in the CBC/DIFF mode. Notice there are three aliquots of the aspirated whole-blood sample that will be used to make dilutions.

#### Figure 2.2-7 CBC/DIFF Mode -**CBC/DIFF Mode** Sample Partitions inside the Probe After aspiration in the CBC/DIFF mode, aliquots of the whole-blood sample are distributed to the various baths as follows (Figure 2.2-5): Diluent The 3 $\mu$ L sample aliquot at the tip of the probe is discarded into the rinse chamber as the Air bubble exterior of the sample probe is rinsed, ensuring sample integrity. Not used 10 µL of sample is delivered to the DIL1/HGB bath for use in preparing the primary RBC/Plt **DIFF** dilution dilution and for measuring the Hgb value. 10 µL of sample is delivered to the WBC/BASO WBC/BASO dilution bath for the WBC/BASO count. 25 µL of sample is delivered to the DIFF bath **RBC/PLT/HGB** first dilution for development of the DiffPlot. 5 µL of remaining sample is discarded into the Not used rinse chamber. 7616001A

Figure 2.2-8 shows the sample partitioning that occurs in the CBC mode. Notice there are only two aliquots of the aspirated whole-blood sample that will be used to make dilutions in this mode of operation. (The DIFF aliquot is not needed in the CBC mode.)

## Figure 2.2-8 CBC Mode -Sample Partitions inside the Probe



## **CBC** Mode

After aspiration in the CBC mode, aliquots of the whole-blood sample are distributed to the various baths as follows (Figure 2.2-5):

- The 3 μL sample aliquot at the tip of the probe is discarded into the rinse chamber as the exterior of the sample probe is rinsed, ensuring sample integrity.
- 10 µL of sample is delivered to the DIL1/HGB bath for use in preparing the primary RBC/Plt dilution and for measuring the Hgb value.
- 10  $\mu$ L of sample is delivered to the WBC/BASO bath for the WBC/BASO count.
- $7 \ \mu L$  of remaining sample is discarded into the rinse chamber.

## 2.3 CYCLE DESCRIPTION

This cycle description focuses on the sequence of the sample probe movement among the baths. It also focuses on the volume of sample and reagents being delivered to make the dilutions needed for sample analysis.

## **Cycle Start Conditions**

Figure 2.3-1 Sample Probe and LED at Start of a Cycle

- The sample probe is in its home position.
- The green LED is glowing indicating the instrument is ready.



Figure 2.3-2 Baths Assembly at Start of a Cycle



• All the baths (except the rinse chamber) are filled with clean diluent.

## **Sample Flow**

Figure 2.3-3 Rinsing Probe Exterior After Aspiration

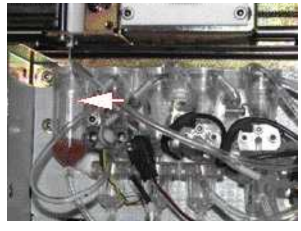
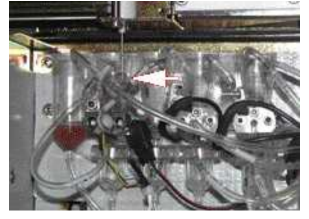
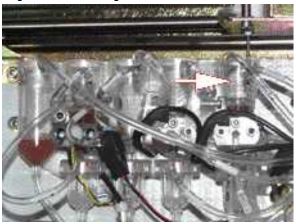


Figure 2.3-4 Making the RBC/PLT First Dilution



- To initiate a cycle, submerge the sample probe in a well-mixed whole-blood specimen and press the aspirate switch to start the cycle.
- All the baths drain.
- A sample of the whole-blood specimen is aspirated.
  - 53  $\mu$ L in the CBC/DIFF mode.
  - $30 \ \mu L$  in the CBC mode.
- The horizontal traverse assembly positions the sample probe over the rinse chamber.
- 3 µL sample aliquot at the tip of the sample probe is discarded into the rinse chamber as the exterior of the sample probe is rinsed.
   Discarding this aliquot helps ensure sample integrity.

- The horizontal traverse assembly positions the sample probe over the DIL1/HGB (first dilution/Hgb) bath.
- The vertical traverse assembly moves the probe downward into the bath. The probe tip is positioned to produce a tangential flow when the sample and diluent are simultaneously dispensed into the bath. For a more detailed description of tangential flow, see Delivery under Heading 2.2, OPERATION PRINCIPLES.
- 10 µL of the whole-blood partitioned for making the first dilution is delivered to the DIL1/HGB bath using a tangential flow of 1.7 mL of diluent.
- The tangential flow of reagent mixes the sample and the diluent. Mixing bubbles enter the bath to make a uniform suspension of cells. This 1:170 dilution is commonly referred to as the first dilution.



- Figure 2.3-5 Making the WBC/BASO Dilution
- The horizontal traverse assembly positions the sample probe over the WBC/BASO bath
- The vertical traverse moves the probe downward into the bath. The tip of the probe is positioned so that a tangential flow occurs as the 10  $\mu$ L of the whole-blood sample and 2.0 mL of WBC Lyse are simultaneously dispensed into the bath.
- The tangential flow of reagent mixes the sample and reagent. Mixing bubbles enter the bath to make a uniform suspension of cells. The WBC Lyse destroys the red blood cells and the specific lytic action on the white blood cells differentiates the basophils from other WBCs.

#### WBC/BASO Bath Dilution

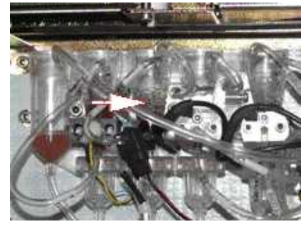
| Whole-blood volume                                 | 10 µL   |
|--|---------|
| Volume of A <sup>C</sup> •T 5diff WBC Lyse reagent | 2000 µL |
| Dilution ratio                                     | 1:200   |

- The horizontal traverse assembly moves the sample probe over the DIFF bath.
- The vertical traverse assembly moves the probe downward into the bath.
- The tip of the probe is positioned so that a tangential flow occurs as 25 µL of the whole-blood sample and 1.0 mL of Fix reagent are simultaneously dispensed into the bath.
- The tangential flow of reagent mixes the sample and the Fix reagent. Mixing bubbles enter the bath to make a uniform suspension of cells. The Fix reagent lyses the red blood cells, stabilizes the WBCs in their native form, and differentially stains the lymphocytes, monocytes, neutrophils, and eosinophils, with the eosinophils staining most intensely.
- After 12 seconds of incubation, the staining process inside the DIFF bath is completed by adding another 1.0 mL of diluent which stops the cytochemical reaction.

## **DIFF Bath Dilution**

| Whole-blood volume                            | 25 µL   |
|---|---------|
| Volume of A <sup>C</sup> •T 5diff Fix reagent | 1000 µL |
| Volume of A <sup>C</sup> •T 5diff Diluent     | 1000 µL |
| Final dilution ratio                          | 1:80    |

## Figure 2.3-6 Making the DIFF Bath Dilution



#### **INSTRUMENT DESCRIPTION** CYCLE DESCRIPTION

Figure 2.3-7 Double Rinse of the Sample Probe

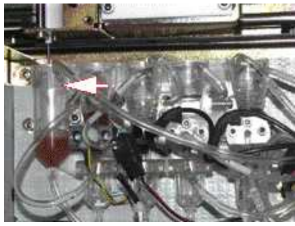
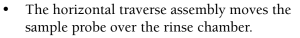
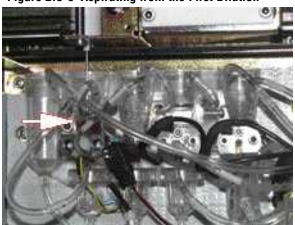


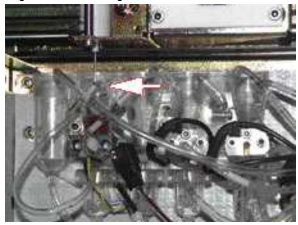
Figure 2.3-8 Aspirating from the First Dilution



•

- A double rinsing (interior and exterior) of the probe removes residual whole-blood sample from inside the probe.
  - In the CBC/DIFF mode, 5 µL is discarded in the rinse chamber.
  - In the CBC mode, 7  $\mu$ L is discarded in the rinse chamber.
- The horizontal traverse assembly moves the sample probe over the DIL1/HGB bath.
- The vertical traverse assembly moves the probe downward into the bath.
- 42.5 µL of the 1:170 first dilution is aspirated into the sample probe.





#### Figure 2.3-9 Rinsing the Outside of the Probe

- While still inside the DIL1/HGB bath, the exterior of the sample probe is rinsed with 0.4 mL of diluent.
- The vertical traverse assembly moves the probe up out of the bath.
- 0.4 mL of Hgb Lyse is added to the bath. The Hgb Lyse reagent rapidly destroys the red blood cells and converts a substantial proportion of the hemoglobin to a stable pigment so a hemoglobin value can be determined.
- Mixing bubbles enter the bath to ensure a uniform dilution.

| DIL1/HGB Bath Dilution                             |         |
|--|---------|
| First dilution                                     | 1:170   |
| Volume of first dilution removed                   | 42.5 μL |
| Volume of A <sup>C</sup> •T 5diff Hgb Lyse reagent | 400 µL  |
| Volume of A <sup>C</sup> •T 5diff Diluent reagent  | 400 µL  |
| Final dilution ratio                               | 1:250   |

- The horizontal traverse assembly moves the sample probe over the RBC bath.
- The vertical traverse assembly moves the probe downward into the bath.
- The tip of the probe is positioned so that a tangential flow occurs as the 42.5 µL of 1:170 dilution obtained from the first dilution in the DIL1/HGB bath and 2.0 mL of diluent are simultaneously dispensed into the bath
- An additional 0.5 mL of diluent is dispensed through the probe at the end of the second dilution.

## **RBC Bath Dilution**

| Volume 1:170 dilution from DIL1/HGB bath          | 42.5 µL  |
|---|----------|
| Volume of A <sup>C</sup> •T 5diff Diluent reagent | 2500 μL  |
| Final dilution ratio                              | 1:10,000 |

## Figure 2.3-10 Making the RBC/Plt Dilution

## 2.4 SAMPLE ANALYSIS

## **RBC and Platelet Analysis**

The RBC/Plt dilution analyzes red blood cells and platelets. This dilution is prepared in two stages – the primary (first) dilution and the secondary (last) dilution.

The primary dilution is made in the DIL1/HGB bath, and the secondary dilution is made in the RBC bath (Figure 2.4-1).

## Figure 2.4-1 Bath Assembly

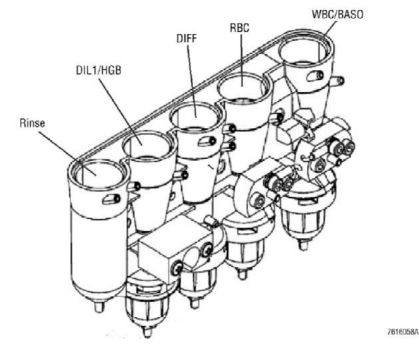


Table 2.4-1 summarizes the technical characteristics required to obtain RBC and Platelet results.

| Dilution Characteristics               |                           |
|--|---------------------------|
| Primary Dilution for RBC and Plt:      |                           |
| Initial volume of whole-blood          | 10 µL                     |
| Volume A <sup>C</sup> •T 5diff diluent | 1700 μL                   |
| Primary dilution ratio                 | 1:170                     |
| Secondary Dilution for RBC and Plt:    |                           |
| Volume of primary dilution             | 42.5 μL                   |
| Volume A <sup>C</sup> •T 5diff diluent | 2500 μL                   |
| Secondary dilution ratio               | 1:58.8                    |
| Final dilution for RBC and Plt results | 1:170 x 1:58.8 = 1:10,000 |
| Reaction temperature                   | 35°C (95°F)               |
| Measurement Characteristics            |                           |
| Method of analysis                     | Coulter Principle         |
| Aperture diameter                      | 50 µm                     |
| Count vacuum                           | 200 mb (5.9 in. Hg)       |
| Count period                           | 2 x 5 seconds             |

#### Table 2.4-1 Technical Characteristics for Obtaining RBC and Platelet Counts

#### Parameter Results Obtained from the RBC/Plt Dilution

This final 1:10,000 RBC/Plt dilution is used to:

- Determine the RBC count.
- Develop the RBC histogram, which is needed to obtain the Hct, MCV, and RDW results.
- Determine the Plt count.
- Develop the Plt histogram, which is needed to obtain the MPV, Pct, and PDW results.

## **Hgb Measurement**

Hemoglobin is determined from the dilution in the DIL1/HGB bath (Figure 2.4-1). This dilution is prepared in two stages – the primary (first) dilution and the secondary (last) dilution.

The primary dilution is made and 42.5  $\mu$ L of that dilution is removed for making the RBC/Plt dilution. A<sup>C</sup>•T 5diff Hgb Lyse and additional diluent are added to make the final 1:250 dilution.

The Hgb concentration is based on the transmittance of light through the optical part of the DIL1/HGB bath using a spectrophotometric technique at a wavelength of 550 nm. The transmittance of the sample dilution is compared to the transmittance of a reagent blank. The system calculates the Hgb using the blank and sample readings.

Table 2.4-2 summarizes the technical characteristics required for measuring hemoglobin.

| Dilution Characteristics  |                   |
|---|-------------------|
| Volume of whole-blood   | 10 µL             |
| Volume A <sup>C</sup> •T 5diff diluent                                    | 1700 μL           |
| Preliminary dilution ratio  | 1:170             |
| Volume of the 1:170 dilution removed<br>(for making the RBC/Plt dilution) | 42.5 µL           |
| Additional volume of A <sup>C</sup> •T 5diff diluent                      | 400 µL            |
| Volume of A <sup>C</sup> •T 5diff Hgb Lyse                                | 400 µL            |
| Final dilution for Hgb determination                                      | 1:250             |
| Reaction temperature  | 35°C (95°F)       |
| Measurement Characteristics   |                   |
| Method of analysis  | Spectrophotometry |
| Wavelength  | 550 nm            |

|

## **WBC Count and Differential**

The WBC count is determined twice using two different methodologies:

- The reference WBC count is the count obtained in the WBC/BASO bath (Figure 2.4-2). The WBC count and the BASO count are determined simultaneously.
- A second WBC count is determined in the flow cell during acquisition of the DiffPlot. The dilution analyzed in the flow cell is prepared in the DIFF bath (Figure 2.4-2).

The WBC counts from the two methodologies are compared and if the results exceed the predefined limits, they will be flagged.

#### Figure 2.4-2 Bath Assembly

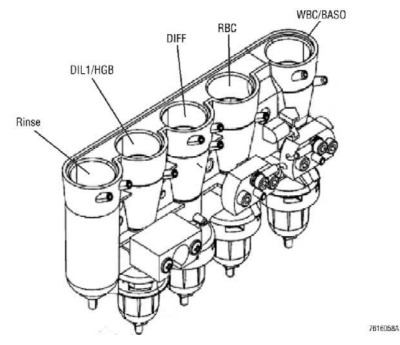


Table 2.4-3 summarizes the technical characteristics required to obtain WBC and BASO results.

| Dilution Characteristics                |                     |
|---|---------------------|
| Volume of whole-blood                   | 10 µL               |
| Volume A <sup>C</sup> •T 5diff WBC Lyse | 2,000 µL            |
| Dilution ratio                          | 1:200               |
| Reaction temperature                    | 35°C (95°F)         |
| Measurement Characteristics             |                     |
| Method of analysis                      | Coulter Principle   |
| Aperture diameter                       | 80 µm               |
| Count vacuum                            | 200 mb (5.9 in. Hg) |
| Count period                            | 2 x 6 seconds       |

## Parameter Results Obtained from the WBC/BASO Dilution

The final 1:200 dilution is used to:

- Determine the WBC count, and
- Develop the WBC/BASO histogram, which is needed to obtain the BASO count.

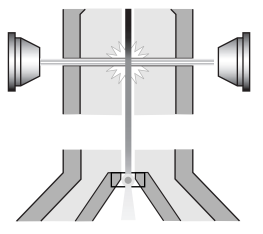
## Differential

Twenty-five microliters (25  $\mu$ L) of whole blood is delivered to the DIFF bath in a flow of A<sup>C</sup>•T 5diff Fix reagent, which lyses the red blood cells, stabilizes the WBC in their native forms, and differentially stains the lymphocytes, monocytes, neutrophils, and eosinophils, with the eosinophils staining most intensely.

The solution is then stabilized with diluent for three seconds and transferred to the measuring bath. See Figure 2.4-3. Each cell is measured in absorbance (cytochemistry) and resistivity (volume).

#### Figure 2.4-3 Flow Cell Operation

2) Second focused flow for optical detection



1) Primary focused flow for impedance

Table 2.4-4 summarizes the technical characteristics required for acquisition of the DiffPlot.

| Dilution Characteristics               |                           |
|--|---------------------------|
| Volume of whole-blood                  | 25 μL                     |
| Volume A <sup>C</sup> •T 5diff Fix     | 1000 μL                   |
| Volume A <sup>C</sup> •T 5diff Diluent | 1000 μL                   |
| Final dilution ratio                   | 1:80                      |
| Reaction temperature                   | 35°C (95°F)               |
| Incubation duration                    | 12 seconds                |
| Measurement Characteristics            |                           |
| Method of analysis                     | Impedance with hydrofocus |
| Aperture diameter                      | 60 µm                     |
| Diameter of the flow                   | 42 µm                     |
| Volume injected                        | 72 µL                     |
| Injection duration                     | 15 seconds                |
| Data accumulation                      | 12 seconds                |

Table 2.4-4 Technical Characteristics for Acquisition of the DiffPlot

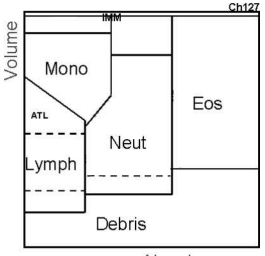
## Parameter Results Obtained from the DIFF Dilution

From these measurements, a DiffPlot is developed with optical transmission (absorbance) on the X-axis and volume on the Y-axis. Figure 2.4-4 shows the DiffPlot regions.

From the DiffPlot, four out of five leukocyte (white blood cell) populations are determined: lymphocytes, monocytes, neutrophils, and eosinophils.

In a typical whole-blood sample, the basophil population (determined in the WBC/BASO bath) is very small compared to the other four white blood cell populations.

Figure 2.4-4 DiffPlot Regions



Absorbance

## **Dilution Summary**

Table 2.4-5 summarizes the dilution characteristics required to obtain CBC and CBC/DIFF parameter results.

| Technical Characteristics<br>for  | Whole-Blood<br>Volume   | Reagent(s)   | Reagent<br>Volume           | Dilution Ratio   | Reaction<br>Temperature |
|---|---|--|-----------------------------|--|-------------------------|
| WBC Count and BASO count<br>(in the WBC/BASO bath)  | 10 µL   | A <sup>C</sup> •T 5diff WBC Lyse   | 2000 µL                     | Final<br>1:200   | 35°C (95°F)             |
| Differential Acquisition with<br>Differential WBC Count<br>(in the DIFF bath)   | 25 μL   | A <sup>C</sup> •T 5diff Fix<br>A <sup>C</sup> •T 5diff Diluent   | 1000 μL<br>1000 μL          | Final<br>1:80  | 35°C (95°F)             |
| Hemoglobin Measurement<br>(in the DIL1/HGB bath)  | 10 μL   | A <sup>C</sup> •T 5diff Diluent<br>After removing 42.5 µL<br>of the 1:170 dilution:<br>A <sup>C</sup> •T 5diff Diluent<br>A <sup>C</sup> •T 5diff Hgb Lyse | 1700 μL<br>400 μL<br>400 μL | Preliminary<br>1:170<br>Final<br>1:250                       | 35°C (95°F)             |
| RBC and PLT Count<br>(in the RBC bath)<br><b>Note:</b> The primary dilution<br>(1:170) is made in the<br>DIL1/HGB bath. | 42.5 μL of the<br>1:170 dilution<br>(from the<br>DIL1/HGB bath) | A <sup>C</sup> •T 5diff Diluent  | 2500 μL                     | Secondary<br>1:58.8<br>1:170 x 1:58.8 =<br>Final<br>1:10,000 | 35°C (95°F)             |

## 2.5 RBC PARAMETER DEVELOPMENT

## **RBC/PIt Dilution**

The final 1:10,000 dilution in the RBC bath contains red blood cells, white blood cells, and platelets. Thresholds are used to separate the platelet pulses, which are much smaller, from the red and white blood cell pulses. Since white blood cells fall in the red blood cell size range, they are counted and sized as RBCs. The WBCs are not sorted out because any interference is usually insignificant; there are normally very few WBCs (thousands) in comparison to the number of RBCs (millions). Only when the white count is markedly elevated is the red cell count or histogram influenced.

## **RBC Count**

The  $A^{C} \cdot T$  5diff hematology analyzer uses duplicate counting criteria, voting criteria, and proprietary flagging information to confirm the parameter result prior to reporting it. To obtain an RBC count result, the instrument compares the data from the two 5-second count periods then votes and rejects any questionable data.

RBC count = Number of cells counted per volume unit x Calibration factor

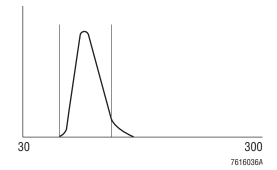
The RBC count is displayed and printed as: RBC = N x  $10^6$  cells /µL.

Note: Cells per microliter (cells/ $\mu$ L) is the US reporting unit format. See Heading A.7 in the Operator's Guide for the other available formats.

## **RBC** Histogram

In addition to being counted, red blood cells are categorized according to size (from 30 fL to 300 fL) by a 256-channel pulse-height analyzer. The pulse-height analyzer uses a number of thresholds to sort the particles into several size (volume) categories and to develop a size distribution curve of the particles. The RBC distribution curve shows cells in their native size. Figure 2.5-1 is an example of an RBC histogram with a normal RBC size distribution.

## Figure 2.5-1 Typical RBC Histogram



## Parameter Results Obtained Using the RBC Histogram

#### **Hct Measurement**

The height of the pulse generated by the passage of a cell through the aperture is directly proportional to the volume of the analyzed red blood cell. The hematocrit (Hct) is the sum of all the digitized pulses. The Hct is displayed and printed as a percentage (%).

**Note:** Percentage (%) is the US reporting unit format. See Heading A.7 in the Operator's Guide for the other available formats.

## **MCV** Calculation

The MCV (Mean Cell Volume) is calculated using the Hct and the RBC count. The MCV is displayed and printed in femtoliters (fL).

**Note:** Femtoliters (fL) is the US reporting unit format. See Heading A.7 in the Operator's Guide for the other available formats.

## **RDW Calculation**

The RDW (Red cell Distribution Width) is an index of the variation or spread in the size of the red blood cells. The study of the RBC distribution detects erythrocyte anomalies linked to anisocytosis and enables the clinician to follow the evolution of the width of the curve relative to the cell number and average volume. Displayed and printed as a percentage, RDW is calculated using the standard deviation (SD) of the RBC population and the MCV.

$$\frac{\text{K SD}}{\text{MCV}} = \text{RDW} (\%)$$

where:

K = System constant

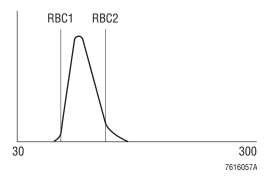
SD = Calculated standard deviation based on the red cell distribution

MCV = Mean Cell Volume of the red cells

## **RBC** Distribution Flags

Once the RBC distribution curve is developed, two positions on the distribution curve are located (Figure 2.5-2):

## Figure 2.5-2 RBC1 and RBC2 Positions - RBC Histogram



## **RBC1 and RBC2 Thresholds**

Thresholds RBC1 and RBC2 define the MICRO and MACRO regions and are calculated based on standard deviation (SD) of the RBC population.

The RBC1 threshold (monitoring area for microcytes) and the RBC2 threshold (monitoring area for macrocytes) identify the points on the curve that are  $\pm 2$  SD from the mean (Figure 2.5-2).

## Flags

Note: MICRO and MACRO flags will be activated in software version 1.0 and higher.

The MICRO flag is generated when the percentage of cells in the microcytic region compared to the total number of RBCs exceeds the preset default limit of 5%. The MACRO flag is generated when the percentage of cells in the macrocytic region compared to the total number of RBCs exceeds the preset default limit of 7.5%. A laboratory may establish its own limits to replace the preset default values.

Note: The MICRO and MACRO flags are independent of the Microcytosis and Macrocytosis flags that are generated from the Low and High patient limits.

## **Hgb Determination**

The hemoglobin (Hgb) released by the lysis of the red blood cells combines with the potassium cyanide to form a stable cyanmethemoglobin compound.

This compound is measured through the optical part of the DIL1/HGB bath using a spectrophotometric technique at a wavelength of 550 nm. Transmittance of the sample dilution is compared with the transmittance of a reagent blank. The system calculates the Hgb using both the blank and sample readings.

The final Hgb result in g/dL represents: absorbance value obtained x Calibration factor.

Hgb is displayed and printed as: Hgb = N g/dL.

**Note:** Grams per deciliter (g/dL) is the US reporting unit format. See Heading A.7 in the Operator's Guide for the other available formats.

## **Hgb Blank Reading**

The Hgb blank value measured during the first patient cycle after a Startup cycle is stored as a reference blank. This blank must be greater than 2.5 Vdc. During each analysis cycle, the instrument checks the measured Hgb blank against the stored Hgb blank reference value using the following formula:

 $(Blank^{Ref} \ge 1/3) + (Blank^{S} \ge 2/3) = Blank^{NR}$ 

where:

Blank<sup>Ref</sup> = Hgb blank reference value

Blank<sup>S</sup> = Hgb blank value from the current cycle

Blank<sup>NR</sup> = New Hgb blank reference value for comparison

**Note:** If the new Hgb blank reference value is within 3% of the old reference value, the Hgb blank reference value is changed to this new value.

## **Sample Reading**

This value is based on the sample, diluent, and Hgb Lyse reagent mixture in the DIL1/HGB bath during sample measurement.

## **Hgb Specific Flags**

If the Hgb blank value is less than 2.5 Vdc, a reject (R) flag occurs on the Hgb value.

If the difference between the new Hgb blank reference value and the original Hgb blank reference value is greater than 3%, a review (R) flag is generated. If three consecutive review

(R) flags occur on the Hgb blank reference value, the  $(\cdots )$  code replaces the Hgb result.

For each Hgb sample read value, the instrument takes three readings. If the difference between these readings exceeds the predefined limits (default setting is 60 A to D units), a voteout (V) flag is generated.

## **MCH and MCHC Calculations**

## **MCH Calculation**

The MCH (Mean Cell Hemoglobin) is calculated from the Hgb value and the RBC count and describes the average weight of hemoglobin in a red cell. The calculation for MCH is:

$$\frac{\text{Hgb}}{\text{RBC}} \times 10 = \text{MCH} (\text{pg})$$

**Note:** Picograms (pg) is the US reporting unit format. See Heading A.7 in the Operator's Guide for the other available formats.

## **MCHC Calculation**

The MCHC (Mean Cell Hemoglobin Concentration) is calculated using the Hgb and Hct values and describes the average concentration of hemoglobin in the red blood cells. The calculation for MCHC is:

 $\frac{\text{Hgb}}{\text{Hct}} \times 100 = \text{MCHC} (\text{g/dL})$ 

Note: Grams per deciliter (g/dL) is the US reporting unit format. See Heading A.7 in the Operator's Guide for the other available formats.

# 2.6 PLATELET PARAMETER DEVELOPMENT

# **RBC/PIt Dilution**

Platelet counting and sizing is also done in the RBC bath. Thresholds separate the platelet pulses, which are much smaller, from the red and white blood cell pulses.

# **PIt Count**

The  $A^{C} \cdot T$  5diff hematology analyzer uses duplicate counting criteria, voting criteria, and proprietary flagging information to confirm the parameter result prior to reporting it. To obtain a Plt count result, the instrument compares the data from the two 5-second count periods then votes and rejects any questionable data.

Plt count = Number of cells counted per volume unit x Calibration factor.

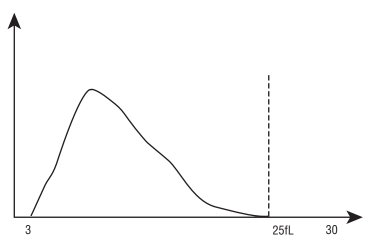
The Plt count is displayed and printed as: Plt = N x  $10^3$  cells /µL.

Note: Cells per microliter (cells/ $\mu$ L) is the US reporting unit format. See Heading A.7 in the Operator's Guide for the other available formats.

# **Platelet Distribution Curve**

Platelets are categorized according to size by a 256-channel pulse-height analyzer. A pulse-height analyzer uses a number of thresholds to sort the particles into several size (volume) categories and to develop a size distribution curve of the particles between 2 fL and 30 fL. The Plt distribution curve shows cells in their native size. Figure 2.6-1 is an example of a Plt histogram with a normal Plt size distribution.





# Parameter Results Obtained Using the Plt Histogram

### **MPV Measurement**

The MPV (Mean Platelet Volume) is measured directly from analysis of the platelet distribution curve. The MPV is displayed and printed in femtoliters (fL).

**Note:** Femtoliters (fL) is the US reporting unit format. See Heading A.7 in the Operator's Guide for the other available formats

## **Pct Calculation**

The Pct (plateletcrit or thrombocrit) is calculated according to the formula:

$$\frac{\text{Plt } (10^3/\mu\text{L}) \times \text{MPV } (\text{fL})}{10,000} = \text{Pct\%}$$

The Pct parameter result is displayed and printed as a percentage (%).

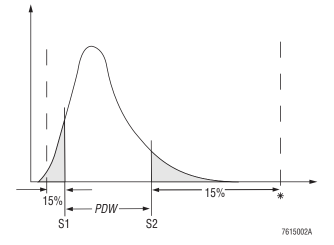
# **PDW Calculation**

PDW (Platelet Distribution Width) is calculated from the Plt histogram as the width of the curve between S1 and S2.

As shown in Figure 2.6-2, S1 and S2 are placed so that:

- 15% of the platelets occur between 2 fL and S1.
- 15% of the platelets occur between S2 and the variable upper threshold.
- Note: This threshold is explained under the Detecting Abnormal Platelet Distributions heading that follows.
- The PDW result is determined on the platelets between S1 and S2.

## Figure 2.6-2 Area of the Plt Histogram Used to Determine the PDW Parameter Result



The PDW parameter result is displayed and printed as a percentage (%).

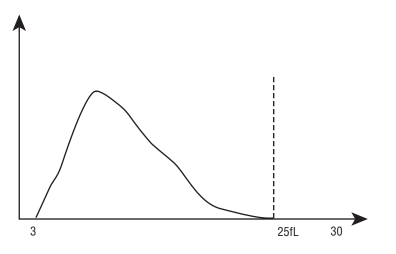
# **Detecting Abnormal Platelet Distributions**

Particles of approximately platelet size can interfere with the platelet histogram and count. Small particles, such as microbubbles or dust, can overlap the low end. Microcytic red cells can intrude at the upper end.

# **Identifying a Normal Distribution**

When a platelet histogram is being evaluated, a mobile threshold can move from its starting position at 25 fL to 18 fL (Figure 2.6-3). The computer searches for a valley between the platelet and red cell populations. If no valley is detected between 18 fL and 25 fL, the threshold remains at 25 fL and no flag is generated.

## Figure 2.6-3 Typical Platelet Distribution Curve



## Interference on the Lower End of the Platelet Distribution Curve

Particles that are approximately platelet size can interfere with the platelet histogram and count. Small particles, such as microbubbles or dust, can interfere at the low end. If the number of pulses in the 2 to 3 fL region is higher than the predefined limits, an SCL flag appears to alert the operator that a significant number of small cells or interference, such as microbubbles, are present.

# Microcytic Interferences on the Upper End of the Platelet Distribution Curve

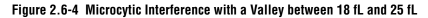
Microcytic red cells can intrude at the upper end of the platelet distribution curve. If the specimen contains microcytes, the A<sup>C</sup>•T 5diff analyzer may be able to successfully eliminate the influence of this interference by repositioning the variable threshold (25 fL threshold) and excluding the microcytes.

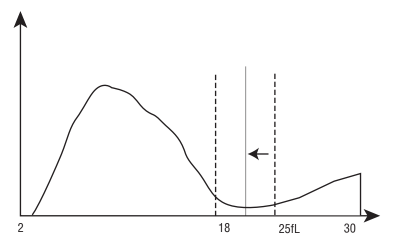
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# Microcytic Interference with a Distinct Valley between 18 fL and 25 fL

If the intrusion of microcytes creates a valley between the 25 fL and the 18 fL thresholds (Figure 2.6-4). The 25 fL threshold is repositioned at the valley to minimize interference to the platelet parameter results. Therefore, the reported platelet results are acceptable. The MIC (microcytes) flag appears to alert the operator that microcytes are present.

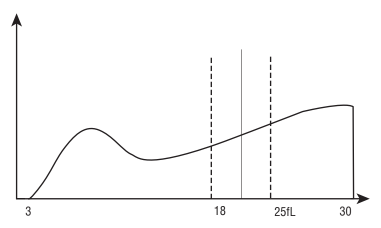




## Microcytic Interference with a Valley below 18 fL

If the microcytes are extremely small so that the valley between the platelet population and the microcyte population falls below the 18 fL limit, the threshold is placed at the 18 fL limit (Figure 2.6-5). The MIC flag appears and the platelet count is flagged to alert the operator that the extremely small microcytes present in this sample could not be eliminated. The platelet count and associated parameters are not reliable and should be verified by an alternative method. To effectively eliminate the microcytes, the Operator's Guide suggests the customer use platelet rich plasma (PRP) or a manual count to verify the results.



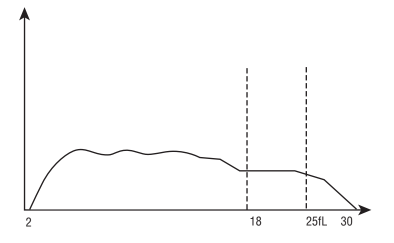


# Interference with No Distinct Valley

Interference present in the upper area of the platelet distribution curve that blends with the platelet population so that there is no clear distinction between the platelets and the interference suggest the presence of schistocytes (fragmented red cells) or platelet aggregates (platelet clumps).

If the threshold cannot be positioned in the 25 fL to 18 fL region, the threshold defaults to the 18 fL position (Figure 2.6-6). The SCH (schistocytes) flag appears and the platelet count is flagged to alert the operator that the interference (which is most likely either schistocytes or platelet clumps) could not be eliminated. The platelet count and associated parameters are not reliable and must be verified using an alternative method.

### Figure 2.6-6 Interference with no Distinct Valley



# 2.7 WBC PARAMETER DEVELOPMENT

## Overview

WBC parameter results are generated from two different dilutions: the 1:200 WBC/BASO dilution which is made and analyzed in the WBC/BASO bath and the 1:80 DIFF dilution which is made in the DIFF bath but analyzed in the flow cell.

# **WBC/BASO** Dilution

The WBC and basophil counts are determined from the 1:200 dilution made in the WBC/BASO bath. To make this dilution, 10  $\mu$ L of whole blood is mixed with 2,000  $\mu$ L of A<sup>C</sup>•T 5diff WBC Lyse reagent. The reaction that occurs lyses the red blood cells and specifically differentiates between basophils and other leukocytes by volume.

# **WBC** Count

The A<sup>C</sup>•T 5diff hematology analyzer uses duplicate counting criteria, voting criteria, and proprietary flagging information to confirm the parameter result prior to reporting it. To obtain an WBC count result, the instrument compares the data from the two 5-second count periods then votes and rejects any questionable data. This is the reference WBC count, which is also the count reported.

A second WBC count is determined in the flow cell during acquisition of the DiffPlot. The two counts are compared and if they differ more than the predefined limit, a flag occurs.

WBC count: Number of cells per volume x calibration factor.

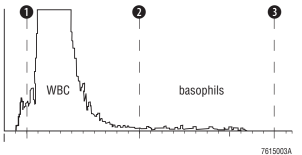
The WBC count is displayed and printed as: WBC = N x  $10^3$  cells /µL.

Note: Cells per microliter (cells/ $\mu$ L) is the US reporting unit format. See Heading A.7 in the Operator's Guide for the other available formats.

## **BASO Count**

Differentiation between basophils and other leukocytes is obtained by means of the A<sup>C</sup>•T 5diff WBC Lyse-specific lytic action. See Figure 2.7-1.

## Figure 2.7-1 Areas Used to Determine WBC and BASO Parameter Results



In Figure 2.7-1, basophils are located in the area between the thresholds labeled **2** and **3**. One hundred percent (100%) of the leukocytes is represented by the total number of nucleated particles plus the basophils within the area between the thresholds labeled **1** and **3**.

The basophil percentage is calculated from the number of particles existing in the area between the thresholds labeled **2** and **3** (Figure 2.7-1)

BASO count = Number of cells per volume x calibration factor in a percentage relative to the number of counted cells (basophils plus other WBC nuclei).

BASO count = 
$$\frac{BASO\%}{WBC\%} \times WBC$$
 count

# **DIFF** Dilution

The data for the DiffPlot is accumulated as the dilution made in the DIFF bath is injected into the flow cell. To make the 1:80 DIFF dilution, 25  $\mu$ L of the whole-blood sample is mixed with 1,000  $\mu$ L of A<sup>C</sup>•T 5diff Fix reagent. The Fix reagent lyses the red blood cells, stabilizes the the white blood cells, and differentially stains the lymphocytes, monocytes, neutrophils, and eosinophils, with the eosinophils staining most intensely. After 12 seconds of incubation, 1,000  $\mu$ L of A<sup>C</sup>•T 5diff Diluent reagent is added to stop the cytochemical reaction. This dilution is injected through the flow cell 15 seconds. For 12 of these 15 seconds, data for developing the DiffPlot is accumulated.

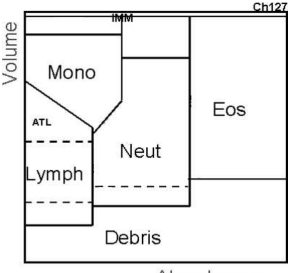
# **DiffPlot Development**

The DiffPlot analysis on the A<sup>C</sup>•T 5diff hematology analyzer is based on three essential principles:

- Dual Focused Flow (DFF) fluid dynamics, which is a process by which individual cells or particles are focused in a stream of diluent (hydrodynamic focusing).
- The volume measurement (Coulter Principle).
- The measurement of transmitted light with zero degree (0°) angle, which permits a response proportional to the internal structure of each cell and its absorbance.

From these measurements, a DiffPlot is developed with optical transmission (absorbance) on the X-axis and volume on the Y-axis. Figure 2.7-2 shows the DiffPlot regions.





Absorbance

# **DiffPlot Regions Defined**

The study of the DiffPlot permits the clear differentiation of four out of five leukocyte populations. In a typical whole-blood sample, the basophil population is very small when compared with the other four white cell populations.

## Neutrophil (Neut)

Neutrophils, with their cytoplasmic granules and segmented nuclei, scatter light according to their morphological complexity. A hypersegmented neutrophil gives an increased optical response when compared to a young neutrophil population. The higher the complexity of the cell, the further to the right they appear in the DiffPlot (Figure 2.7-2).

### Lymphocyte (Lymph)

Lymphocytes, typically being small with regular shape are smaller in volume and lower in absorbance than the other cells, and are positioned in the lower region of the DiffPlot (Figure 2.7-2). Normal lymphocyte populations typically have a homogeneous volume with a Gaussian (bell-shaped) distribution.

Large lymphocytes, reactive lymphoid forms, stimulated lymphocytes and plasma cells are found in the upper portion of the lymphocyte region (Figure 2.7-2).

The lower area of the lymphocyte zone is normally empty; however, when small lymphocytes are present, a population may exist in this area (Figure 2.7-2).

The presence of platelet aggregates is indicated by a distribution pattern that moves from the DiffPlot origin into the lymphocyte region (Figure 2.7-2).

NRBC cytoplasmic membranes lyse like those of mature erythrocytes. The small nuclei that remain appear in the debris and small lymphocyte regions (Figure 2.7-2).

## Monocyte (Mono)

Monocytes are typically large cells with a kidney-shaped nucleus and agranular (granule-free) cytoplasm. These cells neither scatter nor absorb large amounts of light and, therefore, are positioned in the lower end of the absorbance axis. Due to their size, the monocytes are clearly positioned high on the volume axis (Figure 2.7-2).

Very large monocytes may be found in the IMM (immature cell) region.

#### **Eosinophil (Eos)**

With the reagent action, eosinophils are the most intensely stained cells for optical separation. Due to the staining and their size, the eosinophils will show higher absorbance than the neutrophils, but will be of similar volume (Figure 2.7-2).

#### Debris

Platelets and debris from erythrocyte lysis represent the background debris population located in the lower region of the DiffPlot.

# **Immature White Blood Cells**

### **Immature Granulocytes**

Immature granulocytes are detected by their larger volume and by the presence of granules that increase the intensity of the scattered light.

Due to their increased volume and similar absorbance, promyelocytes, myelocytes, and metamyelocytes are located above the neutrophil population and are typically counted as IMM cells. IMM cells are included in the reported neutrophil value.

## **Band Cells**

Band cells are typically larger or of similar size to the neutrophils; however, due to their low level of cellular complexity, they absorb less light. As a result, band cells tend to appear in the region between the neutrophils and the monocytes.

## **Blast Cells**

Blast cells are generally larger than monocytes and have similar absorbance. When blast cells are present, they are generally located above the monocytes, which means they will be included in the IMM cell count.

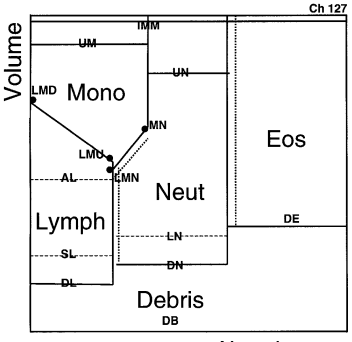
Small blasts will be located between the normal lymphocyte and monocyte populations.

# **DiffPlot Thresholds**

Most of the population partition thresholds are fixed and give the limits of the morphological normality of leukocytes. Changes in the morphology of a population are expressed on the DiffPlot by a shifting of the corresponding population. Volume and absorbance thresholds are used to detect shifting populations. Volume thresholds and definitions are shown in Figure 2.7-3. Absorbance thresholds and definitions are shown in Figure 2.7-4. The NL, NE and MN alarms are also included in Figure 2.7-4.

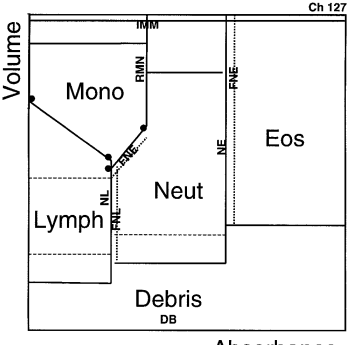
2

Figure 2.7-3 Volume Thresholds



Absorbance

- **DL Threshold** Separates debris and small lymphocytes.
- **DN Threshold** Separates debris and lower neutrophils.
- **SL Threshold** Separates small lymphocytes and lymphocytes.
- **LN Threshold** Separates neutrophils and lower neutrophils.
- **DE Threshold -** Separates debris and eosinophils.
- **LMN Threshold** Intersection point between the lymphocyte, monocyte, and neutrophil thresholds.
- AL Threshold Separates lymphocytes and atypical lymphocytes.
- **LMU Threshold** Lower point on the separation slope between atypical lymphocytes and monocytes.
- **LMD Threshold** Upper point on the separation slope between atypical lymphocytes and monocytes.
- **MN Threshold** Upper point on the separation slope between monocytes and neutrophils.
- **UM Threshold** Separates monocytes and upper monocytes.
- **UN Threshold** Separates neutrophils and upper neutrophils.



# Figure 2.7-4 Absorbance Thresholds / NL, NE and MN Alarms

Absorbance

NL Threshold - Separates lymphocytes and neutrophils.

**RMN Threshold** - Separates upper monocytes and upper neutrophils.

**NE Threshold** - Separates neutrophils and eosinophils.

#### NL, NE and MN Alarms

FNL - # of channels for NL alarm area.

**FNE** - # of channels for NE alarm area.

**FMN** - # of channels for MN alarm area.

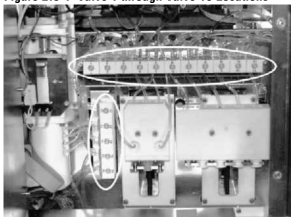
# 2.8 PNEUMATIC/HYDRAULIC SYSTEM

# **Functions of Valves**

Valve blocks are located close to the elements concerned. Five different blocks:

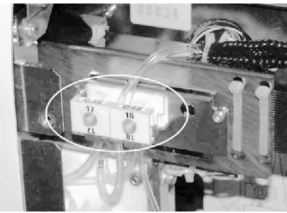
- Valves 1 to 11 (Figure 2.8-1):
  - In the left side compartment behind the Main card,
  - Horizontal block,
  - Above the 5diff syringe and reagent syringes assembly.
- Valves 12 to 16 (Figure 2.8-1):
  - In the left side compartment behind the Main card,
  - Vertical block,
  - Beside the count syringe.

Figure 2.8-1 Valve 1 through Valve 16 Locations



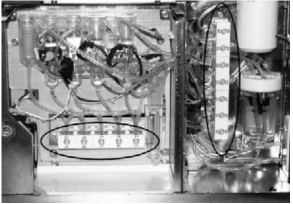
- Valves 17 to 18 (Figure 2.8-2):
  - In the right side compartment,
  - Horizontal block,
  - At the top of the vertical traverse (above the sample syringe assembly).

Figure 2.8-2 Valve 17 and 18 Location



- Valve 19 is not used on the A<sup>C</sup>•T 5diff hematology analyzer.
- Valves 20 to 26 (Figure 2.8-3):
  - In the right side compartment,
  - Vertical block,
  - Beside the waste syringe.
- Valves 27 to 31 (Figure 2.8-3):
  - In the right side compartment (bath enclosure area),
  - Horizontal block,
  - Below the baths assembly.

### Figure 2.8-3 Valve 20 to Valve 31 Locations



See Table 2.8-1 for a description of the functions for each valve in the A<sup>C</sup>•T 5diff hematology analyzer.

| Valve | Function                  | Action  |
|-------|---------------------------|---|
| 1     | Differential diluent      | Select flow cell sheath 2 / DIFF bath                   |
| 2     | Differential diluent      | Select flow cell sheath 1 / sheath 2                    |
| 3     | Differential diluent      | Select input/output for flow cell Diluent syringe       |
| 4     | Flow cell sample supply   | Opens pathway from the DIFF bath to the flow cell       |
| 5     | Flow cell sample injector | Opens waste path for sample injector syringe            |
| 6     | Hgb Lyse syringe flow     | Selects input/output of Hgb Lyse syringe                |
| 7     | Rinse syringe flow        | Selects input/output of Rinse syringe                   |
| 8     | Fix syringe flow          | Selects input/output of Fix syringe                     |
| 9     | Diluent syringe flow      | Selects input/output of Diluent syringe                 |
| 10    | Diluent output control    | Routes diluent to probe rinse block or heating coil     |
| 11    | WBC Lyse syringe flow     | Selects input/output of WBC Lyse syringe                |
| 12    | Rinse output control      | Selects rinse to probe rinse block or WBC/BASO bath     |
| 13    | Count syringe vent        | Opens vent line of count syringe                        |
| 14    | RBC/PLT count valve       | Opens vacuum count line for RBC bath                    |
| 15    | Diluent reservoir vent    | Selects between vacuum and vent for diluent reservoir   |
| 16    | Count syringe drain       | Opens count syringe drain path                          |
| 17    | Probe rinse drain         | Opens drain line for probe rinse block                  |
| 18    | Probe diluent             | Routes diluent/rinse to sample syringe or rinse block   |
| 19    | Spare                     | Not used  |
| 20    | Waste syringe vent        | Opens waste vent (through the rinse chamber)            |
| 21    | Sweep flow diluent        | Routes diluent to heating coil or sweep flow            |
| 22    | Diluent bath select       | Route diluent (via heating coil) to Hgb or RBC bath     |
| 23    | WBC/BASO count vacuum     | Routes vacuum direct or through RBC/PLT count head      |
| 24    | Flow cell drain           | Opens path from flow cell output to DIFF bath for drain |
| 25    | Diluent reservoir input   | Opens diluent source to diluent reservoir               |
| 26    | Waste syringe control     | Selects waste to syringe / syringe waste out            |
| 27    | Rinse chamber drain       | Opens drain path from rinse chamber                     |
| 28    | HGB bath drain            | Opens drain path from Hgb bath                          |
| 29    | DIFF bath drain           | Opens drain path from the DIFF bath                     |
| 30    | RBC bath drain            | Opens drain path from the RBC bath                      |
| 31    | WBC/BASO bath drain       | Opens drain path from WBC/BASO bath drain               |

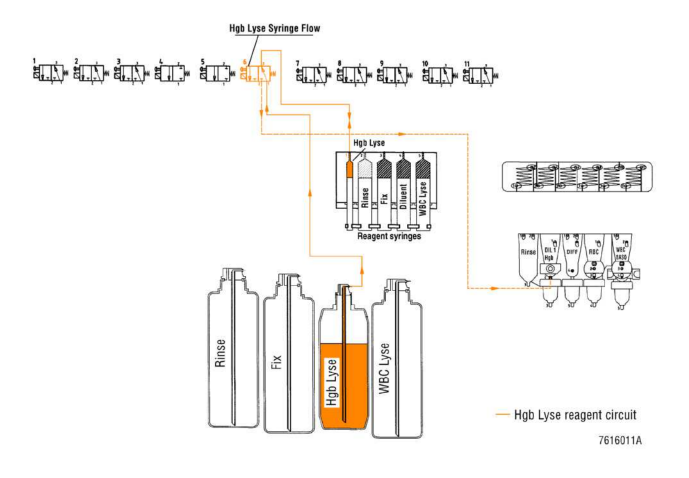
#### Table 2.8-1 Valves and their Functions

# **Pneumatic Diagrams**

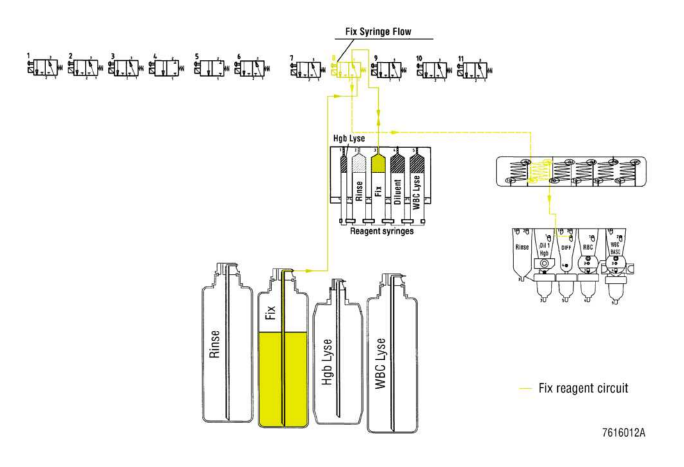
To locate the pneumatic diagram for a desired reagent or the waste circuit, see the designated figure:

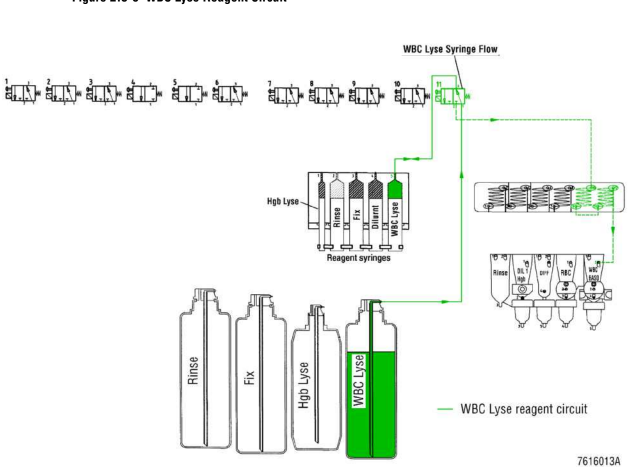
- For the Hgb Lyse reagent circuit, see Figure 2.8-4.
- For the Fix reagent circuit, see Figure 2.8-5.
- For the WBC Lyse reagent circuit, see Figure 2.8-6.
- For the Rinse reagent supply circuit, see Figure 2.8-7.
- For the Probe Rinse reagent circuit, see Figure 2.8-8.
- For the WBC/BASO Rinse reagent circuit, see Figure 2.8-9.
- For the Diluent reagent circuit, see Figure 2.8-10.
- For the Probe Diluent reagent circuit, see Figure 2.8-11.
- For the Bath Diluent reagent circuit, see Figure 2.8-12.
- For the waste circuit, see Figure 2.8-13.

# Figure 2.8-4 Hgb Lyse Reagent Circuit



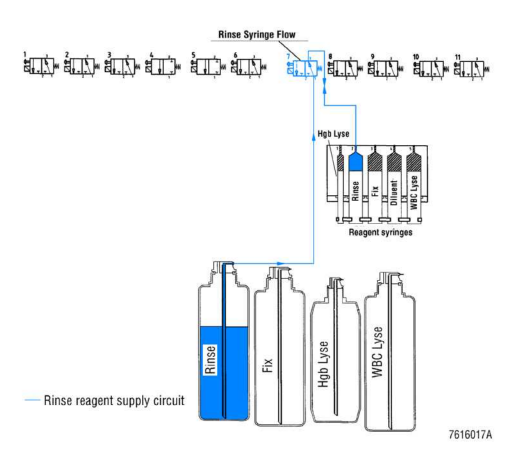




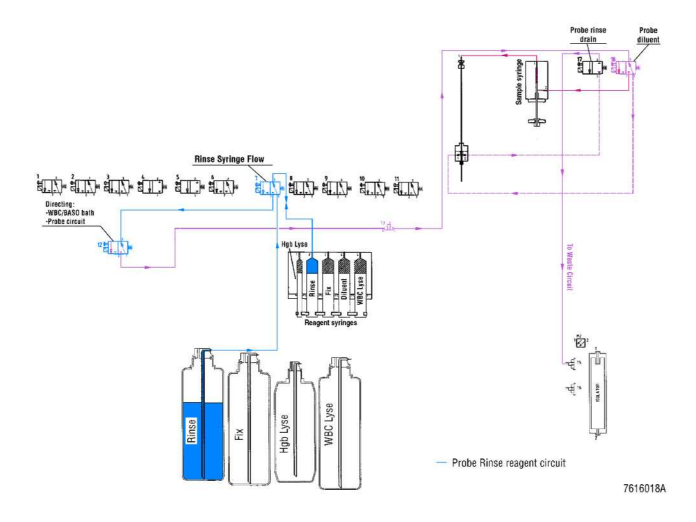


# Figure 2.8-6 WBC Lyse Reagent Circuit

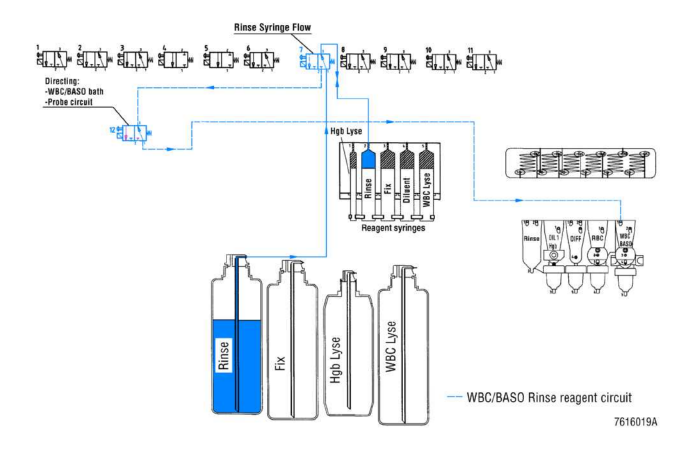










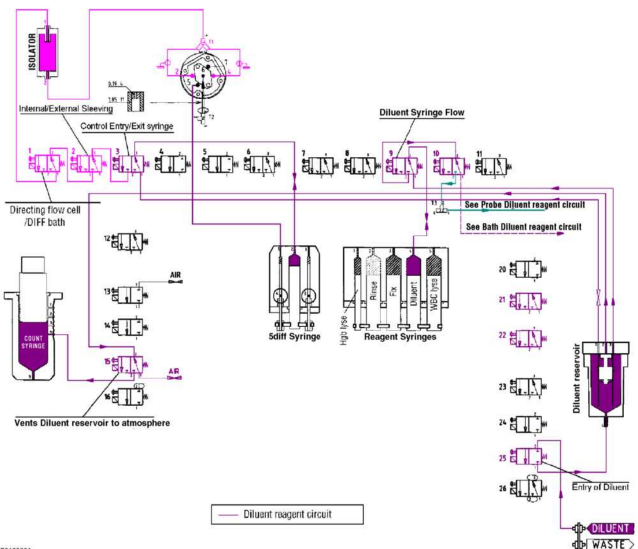


# **Diluter System**

### **Diluent Input** (Figure 2.8-10)

Diluent for the  $A^{C} \cdot T$  5diff hemtology analyzer enters a fitting at the rear of the instrument from a diluent container and is stored in the diluent reservoir. The input tubing from the diluent container should be no longer than 2 meters (78.7 in.) and the top of the container cannot be lower than 80 cm (31.5 in.) from the input fitting on the instrument. Vacuum, produced by the count syringe, is used to draw diluent into the diluent reservoir from the diluent container. This vacuum is applied to port 2 of the diluent reservoir. Solenoid valve 15, Diluent Reservoir Vent, is activated to connect the diluent reservoir to the count syringe. When in the normal inactive state, this valve vents the diluent reservoir to atmosphere. The vent tubing from port 1 of valve 15 is notched to ensure it does not seal against any surface and is routed to the left side drip tray. A float sensor located in the reservoir is used to determine when the reservoir is full, or needs more diluent.

#### Figure 2.8-10 Diluent Reagent Circuit



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Two ports distribute diluent from the diluent reservoir. Port 1 supplies diluent to be used by the flow cell while port 3 supplies diluent for the baths and aspirate probe. Solenoid valve 3 normally connects port 1 of the diluent reservoir to the center syringe piston in the 5diff syringe assembly, allowing the syringe to fill with diluent. For diluent output from the 5diff syringe see 5diff Syringe and Flow Cell.

Solenoid valve 9 connects the diluent reagent syringe to port 3 of the diluent reservoir, and when it is in a normal inactive state, this path is open, allowing the syringe to be filled. For output from the main diluent syringe, see Diluent to Baths and Probe and Probe Rinse.

### **5diff Syringe and Flow Cell**

Diluent for several flow cell requirements originates at the center syringe piston of the 5diff syringe assembly. When solenoid valve 3 is active, valves 1 and 2 have control of the diluent path. If valve 2 is energized, diluent is sent from the large center syringe piston to (and through) the left injector piston. A short upward stroke is used to fill the small left injector piston with clean diluent. Since the center syringe piston is much larger, even a short stroke will produce excess diluent, which exits out the top of the left injector piston and through the flow cell.

When valve 2 is in the normal state, valve 1 determines where diluent is routed. An energized valve 1 allows diluent to be sent through the heater assembly to port 2 of the DIFF bath. This is used for the second part of the DIFF dilution when 1 mL of diluent is added to the sample 12 seconds after the original dilution with Fix. This stops the staining action of the Fix.

Solenoid valve 1 in its normal inactive state creates a diluent path for the flow cell outer sheath. During normal flow cell operation, the center syringe piston (solenoid valve 3 energized and solenoid valves 1 and 2 in a normal inactive state) sends diluent through an electrical isolator and T-fitting to ports 2 and 4 of the flow cell. This creates an outer sheath or fluid pipe around the stream of fluid exiting the flow cell aperture. This sheath is approximately 140 µm in diameter.

During the upward stroke of normal flow cell operation, the 5diff syringe assembly creates the inner sheath flow. Diluent exits from the top of the left injector piston and enters the flow cell at port 5. It then forms the inner sheath, creating a fluid pipe around the injected sample. The sample injector creates sample flow of 40  $\mu$ m in diameter, while the sample and inner sheath is forced through the aperture, which is 60  $\mu$ m in diameter. On exiting the aperture, an outer sheath is created and this double sheath around the sample is called Dual Focused Flow, or DFF.

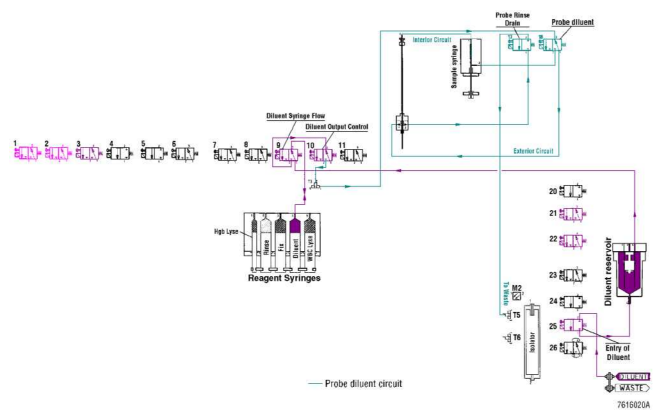
Sample is sent through the flow cell with the right injector piston (see Figure 2.8-13), but it first must be positioned. The sample dilution, created in the DIFF bath, is drained with the waste syringe. The vacuum path to the waste syringe, starting at the bath, is through fluid sensor M1, an energized solenoid valve 4, through the right injector piston (note that the piston is not being used at this time), through an energized solenoid valve 5, T-fitting T6, waste fluid sensor M2, a fluidic/electrical isolator, and solenoid valve 26 in its normal inactive state. Fluid is drained from the bath until sensor M1 detects air. The length and size of the tubing ensures that the sample does not actually reach or enter the right injector piston when sensor M1 detects air and stops flow. This is critical. When the injector piston pushes up, sample in the tubing between T-fitting T2 and port 5 of the right injector piston is sent through the flow cell.

I

# **Probe and Probe Rinse**

When valves 9 and 10 are both energized (Figure 2.8-11), diluent output from the diluent syringe is sent to the probe and probe rinse block by way of solenoid valve 18. An energized solenoid 18 routes diluent through the sample syringe and out the probe for backwash and sometimes dilution. Note that the sample syringe is not used at this time. Since the piston seal is an O-ring at the base of the piston, rather than a seal at the top of the piston, the bottom fitting and top fitting on the syringe assembly have an open fluid path. Diluent is sent through an inactive sample syringe by the diluent syringe. The sample syringe is only used when aspirating or dispensing sample.

### Figure 2.8-11 Probe Diluent Reagent Circuit



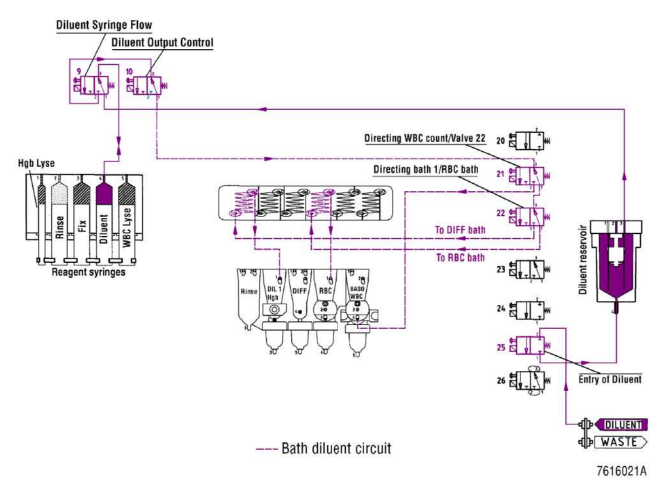
With solenoid valve 18 in its normal inactive state, the diluent is sent to fitting 1 on the probe rinse block. This fluid exits the rinse block from fitting 2 and passes through solenoid valve 17 to the waste system. Usually, this occurs while the probe is moving up through the rinse block to clean blood off the exterior of the probe. However, this is not always the case. On occasion, solenoid 17 is not opened, and the probe does not move. Diluent is forced to flow down the outside of the probe. This is done once to clean the exterior of the probe without moving it and another time to add a small amount of diluent to a dilution.

## **Diluent to Baths**

Energizing solenoid valve 9 routes the diluent syringe output through solenoid valve 10 (Figure 2.8-12). In the normal state of valve 10, diluent is routed for use at the sample baths. There are 3 uses of diluent at the baths, diluent for dilution in the Hgb bath, diluent for dilution in the RBC bath, and rinse for the WBC/BASO counting head.

Solenoid valve 21 routes diluent for dilutions through valve 22 when in a normal inactive state. Valve 22 selects the RBC bath (normal, inactive state) or the Hgb bath (energized state). The fluid paths to the baths both go through a heater block. Each path has a 1 mL coil of fluid in the heater block which allows 1 mL of reagent to be heated to 35°C prior to being delivered to the bath.





When solenoid valve 21 is energized, fluid is sent to the WBC/BASO counting head to rinse the pathway and counting head. This is necessary since fluid in this area is used to flush cells from the rear of the RBC/Plt aperture, a technique called the Rinse Flow System (RFS). During the RBC/Plt count, the WBC/BASO count head is connected to the RBC/Plt count head through an inactive (normal state) valve 23. The vacuum applied to the RBC/Plt aperture actually draws fluid (rinse solution, not diluent) from the WBC/BASO bath, through the WBC/BASO aperture, and past the rear of the RBC/Plt aperture, sweeping away any RBC cells from the rear of the aperture. When solenoid valve 23 is energized, the WBC/BASO counting head is connected directly to the counting syringe, providing vacuum for the WBC/BASO count.

### Waste System

The waste and drain system (Figure 2.8-13) comprises many components. The waste syringe itself is used to drain the baths, and to expel waste from the baths. The count syringe expels any waste that it accumulates during count directly into the waste system through normally closed solenoid valve 16. Waste from the probe rinse block is pushed out by the diluent syringe as well as being evacuated with the waste syringe.

The bath drain system connects each bath, including the rinse chamber, through a normally closed solenoid valve to the waste system with a series of T-fittings. There is a fluid isolator chamber between the baths and the waste syringe and container. This electrically isolates the baths (and flow cell and aspirate probe) from any interference that can be picked up by the external waste system. Draining waste from any bath involves opening the associated solenoid valve while the drain syringe is filling. Solenoid valve 26 is then energized, which connects the waste syringe to the external waste system, and waste is expelled.

There is also a fluid sensor, M2, in the waste system, just before the isolator. It is used to detect whether the waste system has fluid or air at the appropriate times. It is not used to stop a drain action, like M1 does when detecting that sample has been drained from the DIFF bath.

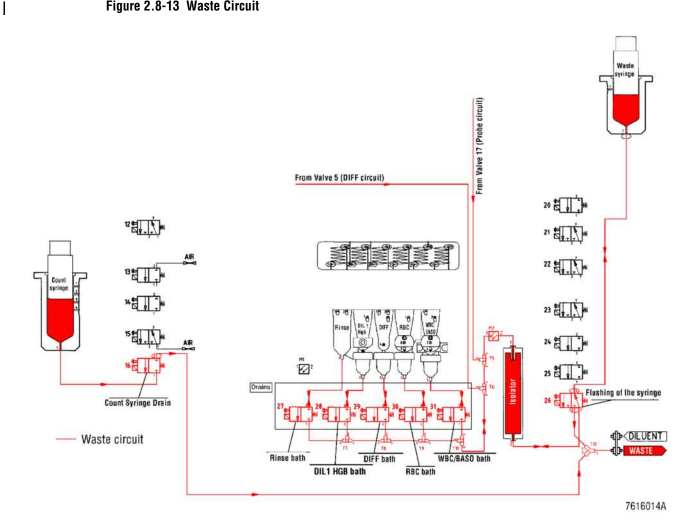


Figure 2.8-13 Waste Circuit

# 2.9 ELECTRONIC SYSTEM

## **Plug/Jack Labels**

In some circumstances, connectors may not have matching plug/jack (P/J) labels. For example, cable connector J5 plugs into board connector J5 on the traverse interconnect card.

# **Optical Preamplifier Card**

This card:

- Performs an absorbance measurement.
- Is a voltage amplifier.
- Contains a photodiode that measures absorbance of light through the cell. The Optical Preamplifier card returns an intensity proportional to the light signal. (The light signal is proportional to the cell size, complexity and staining.)

# LCD and Keypad Card

This card includes the keypad and LCD screen is connected to the Main card using one flat cable through connector J1. The LED card is connected to this card through connector J4.

# **LED Card**

This card supports the LEDs that are used to quickly identify instrument status. These LEDs are located on the front panel.

- A red glowing LED indicates the instrument is busy.
- A green glowing LED indicates the instrument is ready for operation.

# **Motor Interconnect Card**

All the different motors and end-of-run sensors are connected to this card. The Motor Interconnect card is connected to the Main card using one flat cable through connector J2.

## **Traverse Interconnect Card**

All the Traverse module electrical functions are connected to this card. The Traverse Interconnect card is connected to the Main card using one flat cable through connector J5.

# 2.10 SOFTWARE STRUCTURE

### Overview

The operating system software provides instrument I/O, system error checks, data analysis, and individual diluter subroutines or functions. The operating system software also provides diluter functions that energize solenoids, drive motors, and check sensors in the diluter.

### **Menu Trees**

Most I/O functions of the operating system software are controlled by the user. This interaction between the user and the instrument is called the user interface. A pushbutton keypad and an LCD screen provide the physical user interface, while menu items provide the software user interface. These menu items have associated arabic numerals.

The Main Menu consists of five options:

- **1. RUN SAMPLES**
- 2. CALIBRATION
- 3. REAGENTS
- 4. DIAGNOSTICS
- 5. SETUP

## How to Select a Menu Item

An operator may select a menu item two different ways,

- 1. Use the **0** or **0** arrow keys to highlight the desired menu item then press **●** (ENTER pushbutton on the instrument keypad) to select the highlighted option. or
- 2. Simply press the numeric pushbutton (on the instrument keypad) that correlates with the desired option. This is the faster way to select a menu item.

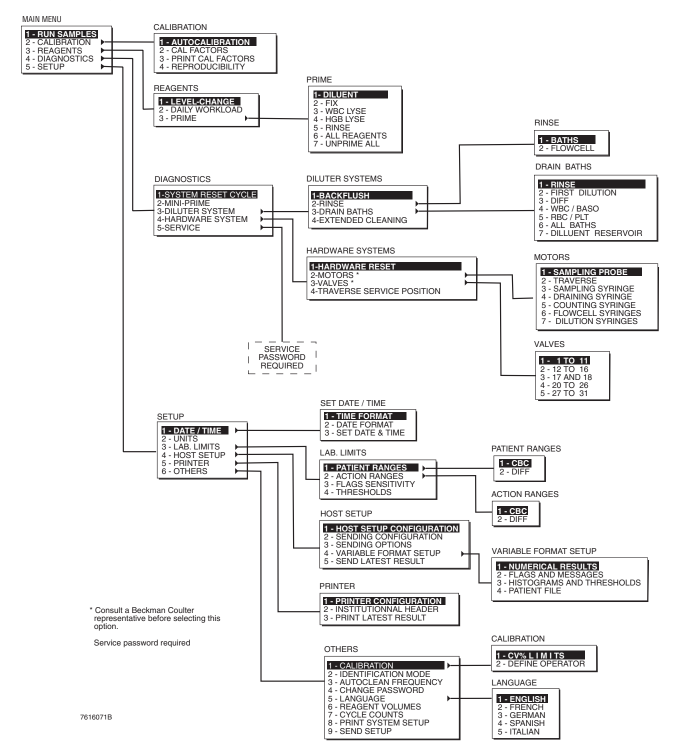
For example, to select the **3**. **REAGENTS** menu item from the Main Menu, an operator may:

- Use the down arrow ♥ to highlight **3. REAGENTS** then press **ENTER** ●. or
- Press the pushbutton labeled **3** on the instrument keypad.

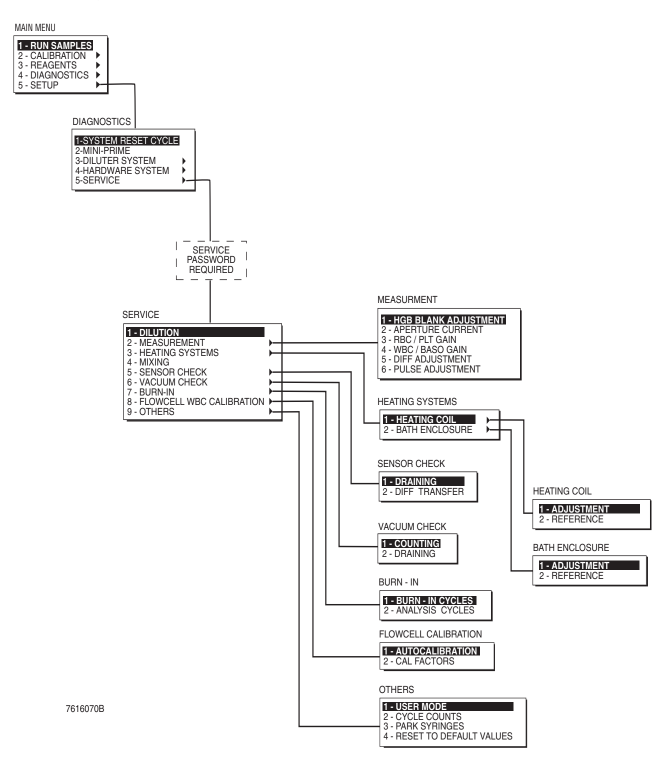
With the exception of Main Menu item **1**. **RUN SAMPLES**, selecting a menu option produces a submenu with a listing of additional options that may be selected.

See Figure 2.10-1 and Figure 2.10-2 for a graphic representation of the menus and menu items (or options) available for the user and service personnel.





#### Figure 2.10-2 Service Menu Tree



#### 3 INSTALLATION PROCEDURES, 3.1-1

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- 3.3-4 Threading the Ribbon, 3.3-2
- 3.3-5 Proper Ribbon Placement, 3.3-3
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- 3.2-1 Whole-Blood Reproducibility CV Limits for 20 Cycles, 3.2-10
- 3.2-2 Calibration Factors Acceptable Range, 3.2-11
- 3.3-1 LX300+ Printer Controls and Indicators, 3.3-7
- 3.3-2 LX300+ Printer Default Settings, 3.3-8

# 3.1 PREINSTALLATION CHECKS

Prior to installing the A<sup>C</sup>•T 5diff hematology analyzer, a pre-site inspection is required to verify the following conditions:

# Environment

The A<sup>C</sup>•T 5diff analyzer should be operated in an indoor location only.

## **Altitude Range**

The A<sup>C</sup>•T 5diff hematology analyzer may be operated at any altitude up to 3,000 meters (9,843 feet). Operation at an altitude over 3,000 meters (9,843 feet) is not recommended.

## **Ambient Temperature**

The ambient operating temperature is 16 to  $34^{\circ}$ C (61 to  $93^{\circ}$ F). If the A<sup>C</sup>•T 5diff analyzer is kept at a temperature less than  $10^{\circ}$ C ( $50^{\circ}$ F), the instrument should be allowed to set at a proper room temperature for one hour before use.

# **Space and Accessibility Requirements**

The  $A^{C} \cdot T$  5diff hematology analyzer should be placed on a clean and level table or work station. Please note that the instrument, printer, and reagents weigh approximately 37 kg (81 lbs).

**WARNING** Risk of operator injury if only one person lifts the instrument. The instrument weighs more than 18 Kg (40 lbs.) and has no lifting handles. To prevent injury, at least two people following necessary safety precautions should lift the instrument together.

The diluent container may be placed on the same level as the instrument or below. If placed on the floor, the top of the diluent container cannot be lower than 80 cm (31.5 inches) from the table level of the instrument. The diluent and waste tubings are limited in length to a maximum of 2 meters (78 inches).

Check the site for proper space allocation (Table 3.1-1).

| Linear Dimensions | Required by Instrument   |
|-------------------|--|
| Height            | 58.0 cm (23.0 inches)  |
| Width             | 44.4 cm (17.5 inches)  |
| Depth             | 50.1 cm (19.8 inches) plus an additional 20.0 cm (8.0 inches) for proper ventilation |

### Table 3.1-1 Space Requirements

## **Electrical Input**

**IMPORTANT** Risk of erroneous results. If an extension cord is used, electrical interference could affect the instrument's operation and results. Locate the instrument close enough to a power outlet that an extension cord is not necessary.

Check for the availability of a power connector. Make sure the instrument is close enough to a power outlet that the ac power cable safely reaches it. The ac power cable is 1.8 meters (6 feet) long and is attached to the back of the instrument, in the lower right corner.

#### **Power Requirements**

Verify the wall socket is an outlet capable of supplying 100 Vac to 240 Vac, from 50 Hz to 60 Hz.

#### Grounding

Proper grounding is required. Verify the ground (earth) for the wall plug is correctly connected to the laboratory grounding electricity installation. If there is no ground, use a ground stake. Current electricity standards must be applied.

#### **Installation Category**

This instrument is designed to be safe for transient voltages according to Installation Category II and Pollution Degree 2.

### **Electromagnetic Environment Check**

The A<sup>C</sup>•T 5diff analyzer produces less than the acceptable level of electromagnetic interference when properly placed. Electromagnetic interferences are limited to levels that allow the correct operation of other instruments conforming to their placement.

To avoid problems, make sure the instrument is not placed near electromagnetic fields or shortwave emissions (such as radar, X-ray machines, scanners, and so forth).

#### **Inspection Report**

Review the findings with your contact person. If deficiencies are present, make sure the customer understands what actions are necessary to meet the specifications for the system. Establish a time frame for completion. Notify your manager if the installation must be rescheduled.

# 3.2 INITIAL SETUP

## **Preinstallation Checks**

If any deficiencies were noted during the preinstallation check, verify they are resolved before installing the instrument.

## **Supplies**

Make sure an adequate supply of reagents, controls, and calibrator are available at the site. For details, see Chapter 1 of the Operator's Guide.

## Unpacking

### Inspection

Inspect all boxes for damage. Notify shipping of external damage.

### **Unpack the Analyzer**

- 1. Unpack the analyzer and place it on the table or bench as determined in the preinstallation site inspection.
- 2. Check the instrument for damage.

### Unpack the Installation Kit, PN XEA484A

- 1. Unpack the Installation kit.
- 2. Using the parts list in Table 8.1-8, ensure that no parts are missing.

### Unpack the Waste Alarm Kit, PN 6912680

- 1. Unpack the Waste Alarm kit.
- 2. Using the packing slip, ensure that no parts are missing.
- 3. Make sure the additional parts needed to complete the installation are available:
  - Two sets of Velcro strips, precut (2" x 1")
    - Hook strip, PN 1017414 (need 2)
    - Loop strip, PN 1017413 (need 2)
  - Tubing, 035 clear polyurethane, 2-inch piece, PN 3202035
  - Feed-through fitting, PN 6216308

### Verify All Caution and Compliance Labels are in Place

1. Verify the caution label on the back of the instrument is in place (Figure 3.2-1).

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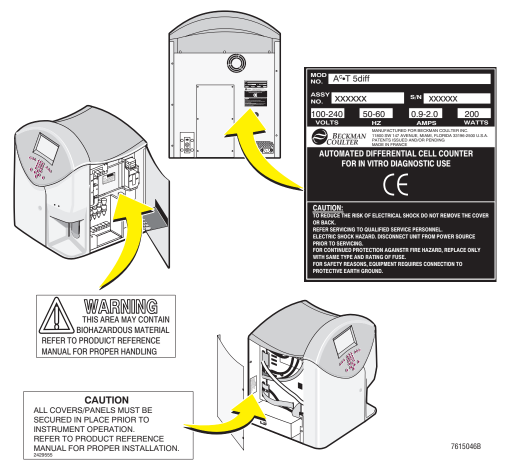
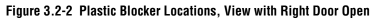
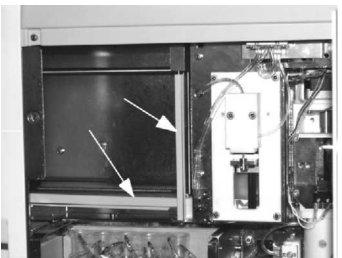


Figure 3.2-1 Warning and Caution Label Locations on the Instrument

- 2. Open the right side door. This door is fastened with two partial-turn, captive-slotted screws.
- 3. Remove the two plastic blockers from the traverse assembly, one from the horizontal and a second from the vertical traverse guide rod. See Figure 3.2-2.





- 4. Manually push the sample probe assembly towards the front of the instrument and verify the warning label is in place (Figure 3.2-1).
- 5. Remove the four hex screws securing the left side panel to the instrument frame and verify the caution label is in place (Figure 3.2-1).
- 6. Replace the left side panel.

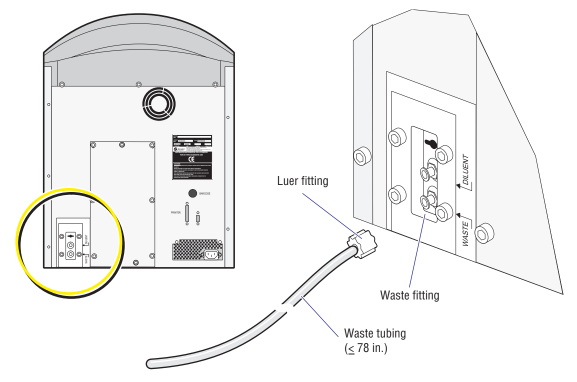
## **Connect the Waste System**

## **Connect the Waste Tubing**

**ATTENTION:** The waste tubing is limited in length to a maximum of 2 meters (78 inches).

- 1. Attach one end of waste tubing (4x6 mm / PN EAE028A) to Luer fitting (PN EAC019A).
- 2. Connect the Luer fitting to the lower waste fitting on the back of the instrument (Figure 3.2-3).

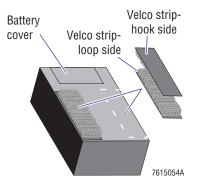
### Figure 3.2-3 Rear Panel Connections



### Install the Waste Alarm

If a 20 L container is used for waste, connect the waste alarm as follows:

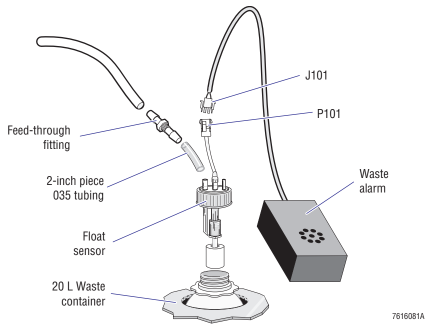
- 1. Remove the battery cover and install the battery in the alarm casing. Replace the cover.
- 2. One at a time, remove the adhesive backing from a loop velcro strip and attach it to the back of alarm casing. The placement must be opposite the battery cover, in the left and right corners (Figure 3.2-4).



#### Figure 3.2-4 Loop-Side Velcro Strip Attachment

- 3. Without removing the adhesive backing, attach the hook fastener of the velcro strips to the loop strips.
- 4. Plug the float sensor connector P101 into the alarm connector J101.
- 5. Invert the float to activate a loud, repetitive beep.
  - If the alarm is working properly, separate the connectors.
  - If a low chirp sound occurs instead, the battery is low. Replace the battery.
- 6. Install the 2-inch piece of 035 tubing (Figure 3.2-5) on the large feed-through fitting (threaded side). Firmly seat.

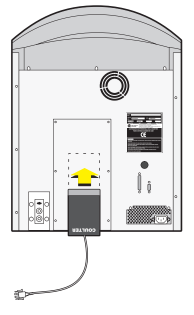
#### Figure 3.2-5 Waste Alarm and Float Sensor Setup



- 7. Install the other end of tubing on the top of float sensor port. Firmly seat.
- Insert the other end of fitting (unthreaded side) into waste tubing.
   Note: Make sure that tubing is installed over both barbs.
- 9. Place the float sensor in a 20-liter waste container. Screw the cap on tightly.

- 10. Install the waste alarm on the back of the instrument as follows:
  - a. While holding the alarm casing with its connector hanging freely downward, remove the adhesive backing from the hook side of the velcro strips.
  - b. Attach the alarm to the rear access panel (Figure 3.2-6).

#### Figure 3.2-6 Position the Waste Alarm on the Rear Access Panel



11. Plug the float sensor connector P101 into the alarm connector J101.

### **Connect the Reagents**

**IMPORTANT** Risk of misleading results. If a reagent pickup tube is contaminated, bacterial and/or fungal growth may occur inside the reagent container. This growth may cause unacceptable background results especially for Plts. When connecting the reagent pickup tubes, the straw portion of the pickup tube should not be touched or laid on an uncovered tabletop. Ensure the reagent pickup tubes remain clean and free of contamination.

#### **Connect the Diluent Tubing**

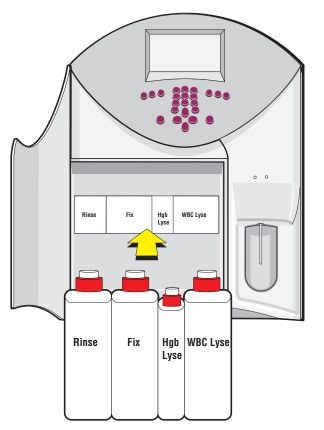
**ATTENTION:** The diluent tubing is limited in length to a maximum of 2 meters (78 inches). The diluent container may be placed on the same level as the instrument or below. If placed on the floor, the top of the diluent container cannot be lower than 80 cm (31.5 inches) from the table level of the instrument.

- 1. Attach one end of diluent tubing (3x6 mm / PN EAE011A) to Luer fitting (PN EAC019A) and the other to pickup tube (PN XEA018A).
- 2. Connect the Luer fitting end to the upper fitting on the rear panel of the instrument (Figure 3.2-3, item 2).
- 3. Insert stopper FBL001A into the container and insert the diluent pickup tube through the stopper.

### **Install the Reagent Bottles**

- 1. Open the reagent compartment door (on the front of the instrument).
  - a. Note the color-coded label on the back panel. The color coding on the instrument label correlates with the color coding on reagent labels (Figure 3.2-7).
  - b. Note the four tubings hanging inside the compartment. Once a bottle stopper is assembled, the assembly is attached to the tubing hanging in its color-coded area and inserted into the reagent bottle. The bottle is then positioned inside the reagent compartment as shown in Figure 3.2-7.

Figure 3.2-7 Reagent Bottle Locations



- 2. Install the WBC Lyse reagent bottle (Figure 3.2-7):
  - a. Loosen the cap of a new WBC Lyse reagent container (yellow label).
  - b. Assemble pickup PN GBG144A with cap PN GAK302A.
  - c. Connect the stopper assembly to the tubing labeled WBC Lyse hanging inside the reagent compartment.
  - d. Without lying the assembly down, remove the cap and ring from the new reagent container and insert the stopper assembly into the WBC Lyse reagent bottle.
  - e. Place the reagent bottle inside the compartment. Position it in front of the yellow WBC Lyse portion of the compartment label.

- 3. Install the Hgb Lyse reagent bottle (Figure 3.2-7):
  - a. Loosen the cap of a new Hgb Lyse reagent container (orange label).
  - b. Assemble pickup PN GBG145A with cap PN GBG155A.
  - c. Connect the stopper assembly to the tubing labeled Hgb Lyse hanging inside the reagent compartment.
  - d. Without lying the assembly down, remove the cap and ring from the new reagent container and insert the stopper assembly into the Hgb Lyse reagent bottle.
  - e. Place the reagent bottle inside the compartment. Position it in front of the orange Hgb Lyse portion of the compartment label.
- 4. Install the Fix reagent bottle (Figure 3.2-7):
  - a. Loosen the cap of a new Fix reagent container (green label).
  - b. Assemble pickup PN GBG144A with cap PN GAK302A.
  - c. Connect the stopper assembly to the tubing labeled Fix hanging inside the reagent compartment.
  - d. Without lying the assembly down, remove the cap and ring from the new reagent container and insert the stopper assembly into the Fix reagent bottle.
  - e. Place the reagent bottle inside the compartment. Position it in front of the green Fix portion of the compartment label.
- 5. Install the Rinse reagent bottle (Figure 3.2-7):
  - a. Loosen the cap of a new Rinse reagent container (blue label)
  - a. Assemble pickup PN GBG144A with cap PN GAK302A.
  - b. Connect the stopper assembly to the tubing labeled Rinse hanging inside the reagent compartment.
  - c. Without lying the assembly down, remove the cap and ring from the new reagent container and insert the stopper assembly into the Rinse reagent bottle.
  - d. Place the reagent bottle inside the compartment. Position it in front of the blue Rinse portion of the compartment label.

### **Install the Printer**

If you have not already done so, install the printer using the procedure under Heading 3.3, PRINTER INSTALLATION. Once the printer is installed, return to this procedure to complete the instrument installation and verification.

### **Power On the Instrument**

- 1. Connect the power cord to the back of the instrument (Figure 3.2-3, item 7).
- 2. Plug the instrument into the ac power source.
- 3. Turn the instrument on.

**Note:** At the factory, a new instrument is left in the Manual Startup mode so the automatic startup is bypassed when the instrument is powered on.

## **Enter Reagent Lot Numbers**

**ATTENTION:** A reagent lot number contains 11 alpha/numeric characters consisting of five numeric digits, an alphabet letter, and five more numeric digits. For example, the lot number for Diluent might be 00102D00002.

- The digits in a lot number are entered using the numeric keypad on the front of the instrument.
- The letter is added using the up arrow key. The first time the up arrow key is pressed, the letter A appears on the screen, the next time the up arrow key is pressed, the letter B appears, and so forth. In the example above, the up arrow key needs to be pressed four times to display the letter D.
- Pressing the right arrow key saves the letter and moves the cursor to the next entry position.
- 1. From the Main Menu, select **3. REAGENTS → 1. LEVEL CHANGE**.
- 2. Press the down arrow key as many times as necessary to highlight the *CHANGE ALL* bar then press **ENTER**. The lot number prompt appears for the reagent.
- 3. Enter the reagent lot number using the instrument keypad as follows:
  - a. Press the numeric keys that correspond with the first five digits.
  - b. Press the up arrow key as many times as necessary to display the required letter.
  - c. Press the right arrow key to save the letter and move to the next entry position.
  - d. Press the numeric keys that correspond with the last five digits.
  - e. Press ENTER to continue to the next reagent.
  - f. Repeat steps a through e for each reagent. When the last lot number is entered, the instrument automatically initiates a prime reagent routine.
- 4. When the prime reagent routine is done,
  - a. Verify all the reagents levels are near 100%.
  - b. Press **ESC** to return to the Reagents menu.

## **Prime the Instrument**

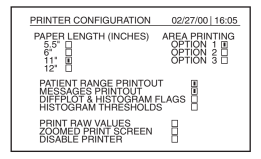
- From the Reagents menu, select 3. PRIME → 6. ALL REAGENTS to prime the reagents again. Note: This second prime is necessary because this an installation. After installation, a reagent line is thoroughly primed and the prime routine done at the end of the lot number change is sufficient.
- 2. Visually inspect the reagent lines and pumps for air bubbles and repeat the priming if air bubbles are still present.

## **Configure the Instrument Printer Settings**

Configure the printer settings, as needed, including.

- Paper length (inches): 5.5 inches, 6 inches, 11 inches, or 12 inches. The paper in the installation kit is 12 inches long.
- Area printing: Options 1 through 3.
- Patient range printout: Prints normal ranges.
- Messages printout: Prints interpretive messages.
- Print Raw Values: Prints raw data. Select this option only for troubleshooting purposes, not for routine operation.
- Zoomed Print Screen: Allows large printout of screen display.
- Disable printer: Does not print the results and does not sound a printer alarm.
- 1. From the Main Menu, select SETUP → PRINTER → PRINTER CONFIGURATION (Figure 3.2-8).

#### Figure 3.2-8 Printer Configuration Menu



2. At the instrument keypad, use the arrow keys to move the blinking cursor into the box next to the desired option.

**Note:** If you press the decimal point key too long, the dot may appear then disappear. The dot inside the box indicates the option is selected.

- a. If the box is empty, pressing the decimal point key places a dot inside the box. The dot indicates the option is selected.
- b. If the box has a dot, the option is already selected. Pressing the decimal point key removes the dot which de-selects the option.
- 3. When all desired selections are made, press **ESC** to save and exit.
- 4. Press **ESC** as many times as necessary to return to the desired menu.
- 5. Printer configuration is detailed in Appendix A of the Operator's Guide.

### Set the User Mode

At the factory, a new instrument is left in the Manual Startup mode so the automatic startup is bypassed when the instrument is powered on. To make sure the system is primed and operational after the Customer has turned the power off then back on, the Manual Startup mode should be disabled.

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Disable the Manual Startup mode as follows:

- 1. From the Service menu, select **9. OTHERS → 1. USER MODE**. The blinking cursor is inside the *MANUAL STARTUP* box. If you look closely, you should also be able to distinguish a dot inside the box.
- 2. At the instrument keypad, press the decimal point key. Make sure the dot no longer appears inside the *MANUAL STARTUP* box. The automatic startup is reactivated.

Note: If you press the decimal point too long, the dot may reappear. The dot inside the *MANUAL STARTUP* box indicates the Manual Startup mode is selected.

3. Press **ESC** to save the selection.

**IMPORTANT** Risk of erroneous Hgb results. The right side door must be closed during Startup. Bypassing the right side door interlock and running the Startup with the right side door open may generate a  $(\cdot \cdot \cdot)$  code for the Hgb blank. Make sure the right side door is closed at least 5 minutes before running a Startup.

4. Turn the instrument OFF for about five seconds, then turn the instrument back ON again. The power ON sequence should now perform a Startup and background cycle. This sequence also establishes a Hgb blank reference which is used as a Hgb blank check during normal sample analysis.

## Verification

- 1. Verify the Startup results passed. If the results do not pass, press **STARTUP** on the instrument keypad to repeat the Startup and background check.
- 2. Use a fresh normal whole-blood specimen to check instrument reproducibility.
  - a. From the Main Menu, select **2**. CALIBRATION **•• 4**. REPRODUCIBILITY.
  - b. Run eleven samples and then delete the first one for a ten-shot reproducibility.
  - c. Verify the reproducibility results are within acceptable limits. See Table 3.2-1.

| Parameter | %CV    | Test Level                    |
|-----------|--------|-------------------------------|
| WBC       | <2.0%  | at 10.0 x 10 <sup>3</sup> /µL |
| RBC       | <2.0%  | at 5.00 x 10 <sup>6</sup> /µL |
| Hgb       | <1.0%  | at 15.0 g/dL                  |
| Hct       | <2.0%  | at 45.0%                      |
| MCV       | <1.0%  | at 90.0 fL                    |
| Plt       | < 5.0% | at 300 x 10 <sup>3</sup> /µL  |

Table 3.2-1 Whole-Blood Reproducibility CV Limits for 20 Cycles

3. Have the Customer perform the calibration procedure using the instructions in Chapter 7 of the Operator's Guide.

4. Verify the calibration factors are acceptable according to Table 3.2-2.

| Parameter | Minimum<br>Acceptable Value | Maximum<br>Acceptable Value |
|-----------|-----------------------------|-----------------------------|
| WBC       | 90                          | 200                         |
| RBC       | 160                         | 290                         |
| Hgb       | 25.0                        | 55.0                        |
| Hct       | 160                         | 290                         |
| Plt       | 180                         | 400                         |
| RDW       | 0.1                         | 0.9                         |

## Table 3.2-2 Calibration Factors - Acceptable Range

# **INSTALLATION PROCEDURES** *INITIAL SETUP*

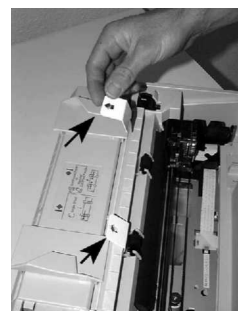
# 3.3 PRINTER INSTALLATION

# EPSON<sup>®</sup> LX<sup>™</sup>- 300 and LX<sup>™</sup>- 300+ Printer Connection

# **Unpack the Printer**

- 1. Unpack the printer and set it on a flat, stable surface.
- 2. Carefully remove all packing materials including the tabs shown in Figure 3.3-1.

# Figure 3.3-1 Carefully Remove All Packing Materials



# Install the Knob

Install the paper-feed knob (Figure 3.3-2).

Figure 3.3-2 Paper-Feed Knob Installation



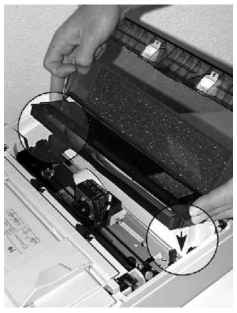
# Install the Ribbon Cartridge

1. Open the printer cover to the upright position, then pull it up to remove it.

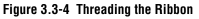
**CAUTION** Risk of damage to the printer. Never move the print head while the printer is turned on. Do not move the print head unless the power is turned OFF.

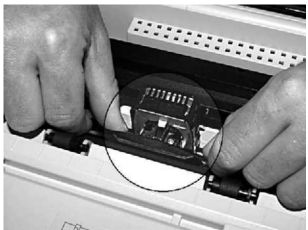
- 2. Slide the printer head to the middle of the printer.
- 3. Locate the ribbon cartridge and turn the ribbon-tightening know in the direction of the arrow to remove any slack from the ribbon.
- 4. Insert the ribbon cartridge into the printer as shown in Figure 3.3-3. Press both sides of the cartridge firmly to fit the plastic hooks into the printer slots.

Figure 3.3-3 Insert the Ribbon Cartridge



5. Guide the ribbon between the print head and ribbon guide (Figure 3.3-4).

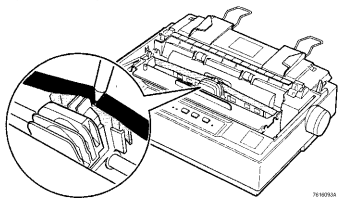




6. Make sure the ribbon is positioned between the print head and the ribbon guide. You may use a pointed object, such as a ball point pen to help guide it into place (Figure 3.3-5). Once in place, turn the ribbon tightening knob to help position the ribbon. It should not be twisted or creased.

**Note:** This ribbon placement can also verified later when the paper is installed. If the ribbon is properly installed, the paper and ribbon do not make direct contact.

#### Figure 3.3-5 Proper Ribbon Placement



7. Manually slide the printer head from side to side to make sure it moves smoothly.

#### **Connect the Printer**

**CAUTION** Risk of damage to the instrument. Connecting the printer to the instrument with the power on could damage the instrument. Ensure the instrument's power is OFF before connecting the printer.

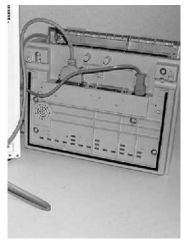
- 1. Make sure the instrument's power is off.
- 2. Locate the printer cable delivered with the instrument.
- 3. Notice that one connector has two screws and the other connector has two clips.
- 4. At the back of the instrument, attach the connector with the two screws to the connector located in the lower right corner of the instrument (Figure 3.3-6).

#### Figure 3.3-6 Cable Connection at the Instrument



- 5. Tighten the screws to secure the cable connection.
- 6. At the printer, lift the printer and set it on its side as shown in Figure 3.3-7.

#### Figure 3.3-7 Cable Connections



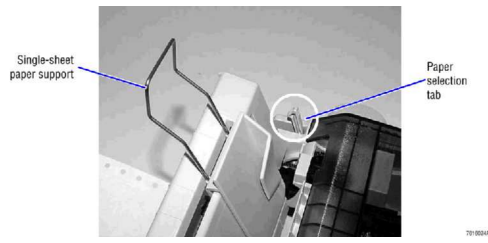
- 7. Attach the other end of the cable to the printer as shown in Figure 3.3-7. Lock the connector in place with its two side clips.
- If a 220 Vdc printer is being installed, insert the power cord in the printer (Figure 3.3-7).
   Note: The power cord for the 120 Vdc printer is already attached.
- 9. If the laboratory will be using continuous feed printing, guide the interface cable and the power cord through the cable slots on the left and right sides of the printer to keep the cables from blocking the paper supply. Both cables can be routed to one side.
- 10. Set the printer flat on the table.
- 11. Plug the printer power cord into the ac power source.

## **Paper Feed Options**

An operator may choose continuous feed printing or printing on single sheets of paper. Set up the option your customer desires to use on a routine basis.

### **Single Sheet Paper Feed Setup**

- 1. Install the paper supports as shown in Figure 3.3-8. Use this support only when printing on single sheets of paper is desired.
- 2. Set the paper selection tab to single sheet Figure 3.3-8.



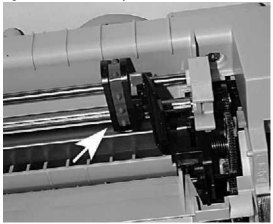
#### Figure 3.3-8 Paper Support for Printing Single Sheets of Paper

3. See the printer user manual for additional information.

#### **Loading Continuous Feed Paper Feed**

- 1. Place the paper selection tab to continuous feed Figure 3.3-8.
- 2. Make sure the printer is turned off.
- 3. Remove the paper guide that separates the incoming paper from the printed paper.
- 4. Release the sprocket units by pulling the sprocket locks forward (Figure 3.3-9).

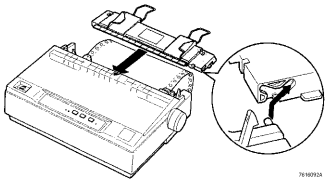
#### Figure 3.3-9 Printer Sprockets



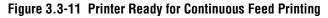
- 5. The printer prints to the right of the position marked 0. Slide the left sprocket unit to position the paper then push the sprocket lock lever back to lock it in place. Slide the right sprocket unit to match the width of the paper, but do not lock it. Move the paper support so it is midway between the sprocket units.
- 6. Make sure the paper has a clean, straight leading edge. Open the sprocket covers. Fit the first holes of the paper over the sprocket pins and then close the sprocket covers.
- 7. Slide the right sprocket unit to remove any slack in the paper and lock it in place. Now the paper is in the paper-park position.

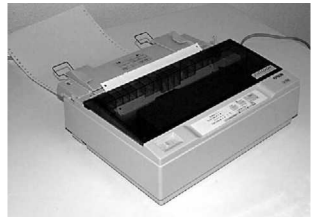
8. To separate the incoming paper from the printed paper, attach the paper guide by holding it horizontally and fitting its notches over the printer mounting posts (Figure 3.3-10). Slide the paper guide towards the front of the printer until you feel it click.

Figure 3.3-10 Replace Paper Guide



- 9. Slide the left and right paper edge guides (used for single sheets) to the center.
- 10. Close the printer cover (Figure 3.3-11).





11. Turn the printer ON.

**CAUTION** Risk of damage to the printer. Use the knob on the right side of the printer only to clear paper jams and only when the printer is off. Otherwise, you may damage the printer or cause it to lose the top-of-form position. Do not turn the printer knob unless the power is turned OFF.

- 12. Press the LF/FF button to feed the paper to the loading position.
- 13. Verify that the paper and the ribbon are not touching each other. If they are in direct contact, the ribbon is not installed correctly. Reposition the ribbon between the print head and ribbon guide (Figures 3.3-4 and 3.3-5).

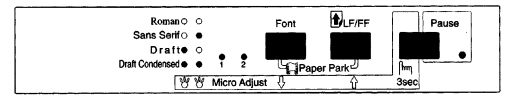
## **Configure the Printer**

## LX300 Printer

**ATTENTION:** The LX300 printer must be set to Draft mode to ensure printouts, such as patient result reports, are formatted correctly.

At the printer control panel (Figure 3.3-12), verify the printer is set to the Draft mode (the LED above the number 2 is glowing). If the LED is not glowing, press the **FONT** key until it does glow.

#### Figure 3.3-12 LX300 Printer Control Panel



When the printer receives data, it will begin printing automatically. For addional information, see the printer user manual.

## LX300+ Printer

At the printer control panel (Figure 3.3-13), press and hold the **TEAR OFF** button while switching the printer On to access the parameter setup meni. Follow the printed instructions to setup the printer. The printer control functions and default settings are shown in Table 3.3-1 and Table 3.3-2.

#### Figure 3.3-13 LX300+ Printer Control Panel

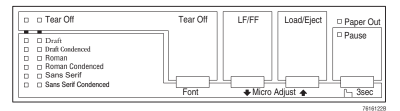


Table 3.3-1 LX300+ Printer Controls and Indicators

| Control/Indicator | Function  |
|-------------------|---|
| LF/FF button      | • Feeds paper line by line when pressed and released.   |
|                   | • Ejects a single sheet or advances continuous paper to the next top-of-form position when held down. |
| Load/Eject button | Loads a single sheet of paper.  |
|                   | • Ejects a single sheet of paper if a sheet is loaded.  |
|                   | • Loads continuous paper backwards to the standby position.   |
|                   | • Feeds continuous paper backwards to the standby position.   |

| Control/Indicator | Function  |
|-------------------|---|
| Paper Out light   | • On when no paper is loaded in the selected paper source or paper is not loaded correctly.                 |
|                   | • Flashes when paper has not been fully ejected or a paper jam has occurred.                                |
| Pause light       | On when printer is paused.  |
|                   | • Flashes when the printer is in the Micro Adjust mode.   |
|                   | • Flashes when the print head has overheated.   |
| Pause button      | • Stops printing temporarily and resumes printing when pressed again.                                       |
|                   | When pressed for 3 seconds, turns on the Micro Adjust mode*. To turn off the mode, press again.             |
| Tear Off button†  | Advances continuous paper to the tear-off position.   |
|                   | • Feeds continuous paper backward from the tear-off position to the top-of-form position.                   |
| Tear Off lights†  | • Lights when continuous paper is in the tear-off position, otherwise the lamps indicate the selected font. |

Table 3.3-1 LX300+ Printer Controls and Indicators (Continued)

position.
† In Micro Adjust mode, press the Tear Off button to select the font to use for printing. The Tear Off lights turn on, off or flash to indicate the selected font.

#### Table 3.3-2 LX300+ Printer Default Settings

| Page length for tractor         | 6 inch     |
|---------------------------------|------------|
| Skip over perforation           | Off        |
| Auto tear Off                   | Off        |
| Auto line feed                  | Off        |
| Print direction                 | Bi-D       |
| Software                        | ESC/P      |
| 0 slash                         | 0          |
| High speed draft                | On         |
| I/F mode                        | Auto       |
| Auto I/F wait time              | 10 seconds |
| Baud rate                       | 19200BPS   |
| Parity                          | None       |
| Parallel I/F bidirectional mode | On         |
| Packet mode                     | Auto       |
|                                 |            |

|  | ,             |
|--|---------------|
| Character table                              | PC 437        |
| International character set for Italic table | Italic U.S.A. |
| Manual feed wait time                        | 1.5 seconds   |
| Buzzer                                       | On            |
| Auto CR (IBM 2380 Plus)                      | Off           |
| IBM character table                          | Table2        |
|  |               |

#### Table 3.3-2 LX300+ Printer Default Settings (Continued)

# **Complete the Instrument Installation**

Return to Heading 3.2, INITIAL SETUP and continue the installation starting at the Power On the Instrument heading.

|

# **INSTALLATION PROCEDURES** *PRINTER INSTALLATION*

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# 4.1 GUIDELINES FOR SERVICING THE A<sup>C</sup>•T 5diff HEMATOLOGY ANALYZER

# **General Guidelines**

# **Safety Precautions**

Review and heed the general safety warnings and cautions listed under Heading 1.2, SAFETY PRECAUTIONS.

### Accessibility

Ensure there is adequate space to work and to access the instrument components safely.

# **Electronic Precautions**

**WARNING** Risk of personal injury. Contacting exposed electronic components while the instrument is attached to power can cause personal injury from electric shock. Power down completely before removing covers to access electronic components.

**CAUTION** Risk of damage to electronic components. If the power is ON while removing or replacing electronic components, the instrument could be damaged. To prevent damage to electronic components, always be sure power is OFF before removing or replacing printed circuit cards and components.

Before disconnecting or reconnecting any electronic component, turn the instrument off and disconnect the power cord from the instrument or the wall outlet. See Power Down / Power Up the Instrument in this section.

#### **Environment Protection**

If the A<sup>C</sup>•T 5diff analyzer is old and ready for disposal, the instrument and its accessories must be collected by a company that specializes in the elimination or the recycling of laboratory equipment according to the legislation.

# Procedures

**WARNING** Risk of personal injury or contamination. If you do not properly shield yourself while servicing the instrument with the doors or panels open, you may become injured or contaminated. To prevent possible injury or biological contamination, you must wear appropriate safety glasses, a lab coat, and gloves when servicing this instrument with the doors or panels open.

Adjustment and maintenance procedures that need to be done on the A<sup>C</sup>•T 5diff hematology analyzer are in this section, including:

- Hydraulic maintenance and adjustments.
- Pneumatic maintenance and adjustments.
- Power supply maintenance and adjustments.
- Electronic maintenance and adjustments.

Note: Read each procedure entirely before beginning the service or repair.

#### **Tools and Supplies**

You can do most procedures using the standard Service Tool Kit and a digital voltmeter (DVM). Any special tools, supplies, or equipment required are identified under the Tools/Supplies Needed heading at the beginning of the procedure.

#### **Instrument Performance Verification**

When a service/repair procedure requires some type of instrument performance verification upon completion, a Verification heading is provided with the necessary steps that must be completed.

#### Service Password

**CAUTION** Risk of instrument damage. Do not disclose the Service password to a Customer. An untrained person with access to the Service menu may activate routines in an unauthorized order which may result in damage to critical systems such as the sample probe.

When performing some of the service and maintenance procedures, it will be necessary to use the Service menu. Access to this Service menu requires a password (which is actually a number) 239. The sequence for accessing the Service menu is:

- 1. From the Main Menu, select **4. DIAGNOSTICS** → **5. SERVICE**. The *SERVICE PASSWORD* prompt appears.
- 2. At the numeric keypad, press [2] [3] [9] then ENTER. The Service menu appears.
- 3. From the service menu, select the desired option.

#### **User Mode**

#### How to Disable the Right Side Door Interlock

**WARNING** Risk of personal injury. When SERVICE is selected as the User mode, the right side door interlock is bypassed allowing instrument operation with the right side door open. Avoid any unnecessary contact with the sample probe. The probe is sharp and may contain biohazardous materials. When the service call is complete, make sure the interlock is reactivated to ensure the Customer is not accidently injured by the sample probe or its movement.

*SERVICE* must be the selected User mode to operate the instrument with the right side door open. When the instrument is set to the Service mode, the interlock for the right side door is bypassed. To set the User mode to service:

- 1. From the Service menu, select **9. OTHERS → 1. USER MODE.**
- 2. At the instrument keypad,
  - a. Press the down arrow key to move the blinking cursor to the SERVICE box.
  - b. Press the decimal point key. Make sure a dot appears inside the box.

**Note:** If you press the decimal point too long, the dot may disappear. The dot inside the *SERVICE* box indicates the right side door interlock is bypassed.

- c. Press **ESC** to exit the User Mode screen.
- d. Press **ESC** as many times as necessary to return to the desired menu.

# How to Reactivate the Right Side Door Interlock

When the service call is complete, make sure the right side door interlock is reactivated to ensure the Customer is not accidently injured by the sample probe or its movement. To reactivate the right side door interlock:

- 1. From the Service menu, select 9. OTHERS → 1. USER MODE.
- 2. At the instrument keypad,
  - a. Press the down arrow key to move the blinking cursor to the SERVICE box.
  - b. Press the decimal point key or the **DEL** key. Make sure the dot no longer appears inside the *SERVICE* box. The right side door interlock is reactivated.

**Note:** If you press the decimal point too long, the dot may reappear. The dot inside the *SERVICE* box indicates the right side door interlock is bypassed.

c. Press **ESC** as many times as necessary to return to the Main Menu.

#### **Power Down / Power Up the Instrument**

#### **Purpose**

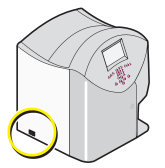
**WARNING** Risk of personal injury. Contacting exposed electronic components while the instrument is attached to power can cause personal injury from electric shock. Power down completely before removing covers to access electronic components.

Following the Power Down procedure ensures all power is removed from the instrument, preventing personal injury from electronic shock.

#### **Power Down**

1. Switch the Power On/Off rocker switch from ON (-) to OFF (**O**). This rocker switch is located at the base of the left side panel.

#### Figure 4.1-1 Location of the Power On/Off Rocker Switch



2. Unplug the ac power cord. Either remove the cord from the instrument (at the back panel, in the lower right corner) or from the ac wall outlet.

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# **Power Up**

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- 1. Plug the ac power cord into the instrument (at the back panel, in the lower right corner) or the ac wall outlet, as applicable.
- 2. Switch the Power On/Off rocker switch from OFF (**O**) to ON (-). This rocker switch is located at the base of the left side panel.

**Note:** An automatic Startup routine and background check is performed. If *MANUAL STARTUP* is the selected User Mode, an automatic Startup and background check does not occur. To initiate a Startup routine and background check, you must press the **STARTUP** button on the instrument keypad.

# **Reset the Instrument**

Two instrument resets are available: System Reset Cycle and Hardware Reset.

# System Reset Cycle

From the Main Menu, selecting **4**. **DIAGNOSTICS** → **1**. **SYSTEM RESET CYCLE** initiates the following actions:

- Checks all stepper motor positions
- Checks all sensors
- Drains and rinses the baths
- Cleans the sample probe
- Checks reagent volumes
- Fills the diluent reservoir
- Checks all mechanical functions

#### Hardware Reset

From the Main Menu, selecting **4**. **DIAGNOSTICS** → **4**. **HARDWARE SYSTEMS** → **1**. **HARDWARE RESET** moves all stepper motors to their home position.

# 4.2 OPENING OR REMOVING INSTRUMENT DOORS, PANELS, AND COVERS

### Purpose

Use the procedures in this section for accessing instrument components. This access includes:

- Opening the Right Side Door
- Removing the Left Side Panel
- Removing the Rear Access Panel
- Removing the Top Cover
- Removing the Front Cover

#### **Tools/Supplies Needed**

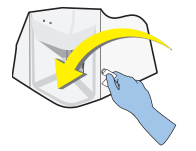
- □ Allen wrenches, 2.5 mm and 3 mm
- □ Screwdriver, T10 torx
- □ Locker key (provided)

# **Opening the Right Side Door**

The pneumatic access door on the right side of the instrument allows the operator to access hydraulic assemblies for maintenance operations. This area is also called the bath enclosure. It is mandatory to keep the door locked during a cycle to ensure proper heating of the dilutions inside the baths. The door is secured using two captive screws that require a special key.

To open the door, place the edge of the key inside the screw slot and turn the captive screw counterclockwise (Figure 4.2-1).

#### Figure 4.2-1 Opening the Right Side Door



#### **Bypassing the Right Side Door interlock**

**WARNING** Risk of personal injury. When SERVICE is selected as the User mode, the right side door interlock is bypassed allowing instrument operation with the right side door open. Avoid any unnecessary contact with the sample probe. The probe is sharp and may contain biohazardous materials. When the service call is complete, make sure the interlock is reactivated to ensure the Customer is not accidently injured by the sample probe or its movement.

*SERVICE* must be the selected User mode to operate the instrument with the right side door open. When the instrument is set to the Service mode, the interlock for the right side door is bypassed. To set the User mode to service:

- 1. From the Service menu, select **9. OTHERS → 1. USER MODE.**
- 2. At the instrument keypad,
  - a. Press the down arrow key to move the blinking cursor to the SERVICE box.
  - b. Press the decimal point key. Make sure a dot appears inside the box.

**Note:** If you press the decimal point too long, the dot may disappear. The dot inside the *SERVICE* box indicates the right side door interlock is bypassed.

- c. Press **ESC** to exit the User Mode screen.
- d. Press **ESC** as many times as necessary to return to the desired menu.
- 3. When the service call is complete, make sure the right side door interlock is reactivated to ensure the Customer is not accidently injured by the sample probe or its movement. To reactivate the right side door interlock:
  - a. From the Service menu, select 9. OTHERS → 1. USER MODE.
  - b. At the instrument keypad,
    - 1) Press the down arrow key to move the blinking cursor to the SERVICE box.
    - 2) Press the decimal point key or the **DEL** key. Make sure the dot no longer appears inside the *SERVICE* box. The right side door interlock is reactivated.

**Note:** If you press the decimal point too long, the dot may reappear. The dot inside the *SERVICE* box indicates the right side door interlock is bypassed.

3) Press **ESC** as many times as necessary to return to the Main Menu.

#### **Removing the Left Side Panel**

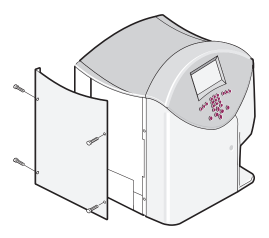
#### Removal

Remove the left side panel to gain access to the Main card:

- 1. Turn the instrument off and unplug the power cord from the instrument or the wall.
- 2. Remove the left side panel (Figure 4.2-2). Use a 3 mm hex key to remove the four hex screws securing the panel to the instrument frame.
- 3. Open the Main card door and carefully anchor the door behind the white plastic catch to keep it open.

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#### **Opening the Main Card Door**

Opening the Main card door allows the operator to access hydraulic assemblies for maintenance operations.

1. To the right of the Main card, turn the two captive knobs counterclockwise to release the Main card door.

**CAUTION** A flat cable is connected to the rear of the Main card. Be careful when you open the door (where the Main card is attached) that you do not disconnect or damage this cable.

2. When opening the Main card door, carefully anchor the door behind the white plastic catch to keep it open.

#### Installation

- 1. Lift the white plastic catch to release the Main card.
- 2. Close the Main card door and turn the two captive knobs clockwise to secure it.
- 3. Replace the left side panel and install the four hex screws removed earlier.
- 4. Reconnect the power cord.
- 5. Turn the instrument on.

#### **Removing the Rear Access Panel**

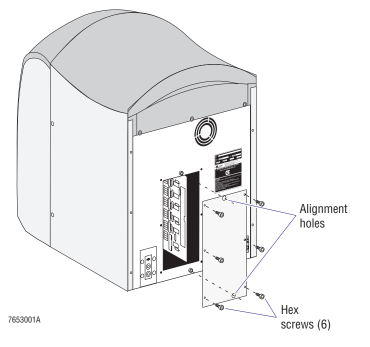
The back panel provides access to the motors and sensor connectors on the Motor Interconnect card.

### Removal

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Use a 3.0 mm Allen key to remove the six hex screws securing the panel to the back of the instrument (Figure 4.2-3). Set the panel aside.



# Figure 4.2-3 Rear Access Panel Screw Locations

#### Installation

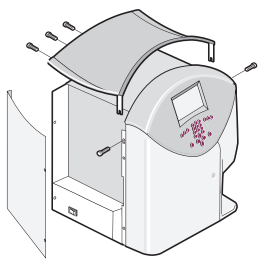
This panel is not reversible, When replacing this panel, make sure the center opening at the top of the panel and the opening at the bottom of the panel are aligned as shown in Figure 4.2-3.

# **Removing the Top Cover**

The top cover is secured to the instrument frame with five hex screws; three screws at the back of the instrument and one hex screw on each side of the instrument. The right door must be opened and the left panel must be removed before removing the top cover.

#### Removal

- 1. Turn the instrument off and unplug the power cord from the instrument or the wall.
- 2. Remove the left side panel. See Removing the Left Side Panel in this section.
- 3. In the left side compartment, remove the hex screw in the front upper corner (Figure 4.2-4).



#### Figure 4.2-4 Top Cover - Side Screw Locations

- 4. Open the right side door. See Opening the Right Side Door in this section.
- 5. In the right side compartment, remove the hex screw in the front upper corner (Figure 4.2-4).
- 6. At the rear of the instrument, remove the three hex screws securing the top cover to the instrument frame (Figure 4.2-4).
- 7. Carefully slide the cover back and off the instrument.
- 8. Set the cover aside.

#### Installation

- 1. Carefully position the top cover back on the instrument.
- 2. In the left side compartment, replace the hex screw in the front upper corner.
- 3. Open the right side door and replace the hex screw in the front upper corner.
- 4. At the rear of the instrument, replace the three hex screws that secure the top cover to the instrument frame.
- 5. Replace the left side panel and install the four hex screws removed earlier.
- 6. Close the right side door.
- 7. Reconnect the power cord.
- 8. Turn the instrument on.

#### **Removing the Front Cover**

The front cover requires that the right side door be opened, and the left side panel and top cover be removed.

#### Removal

- 1. Turn the instrument off and disconnect the power cord from the instrument or the wall.
- 2. Remove the top cover from the instrument. See Removing the Front Cover in this section.

- 3. Open the reagent door and remove the four bottles.
- 4. Disconnect the tubing from the stopper.

**CAUTION** Risk of damage to the reagent compartment door. The two left screws inside the reagent compartment not only secure the compartment to the instrument frame but also secure the reagent compartment door. When these screws are removed, the door may fall. If the door becomes bent, it may not close properly. When removing the two left screws, hold the reagent compartment door securely to prevent it from falling when it detaches.

5. Unscrew the four hex screws shown in Figure 4.2-5. When removing the two left screws, hold the reagent compartment door securely to prevent it from falling when it detaches from the instrument.



#### Figure 4.2-5 Reagent Compartment Screw Locations

- 6. Make sure the sample probe is inside its housing then push the sample probe housing towards the back of the instrument.
- 7. Unscrew several turns the two torx screws shown in Figures 4.2-6 and 4.2-7.

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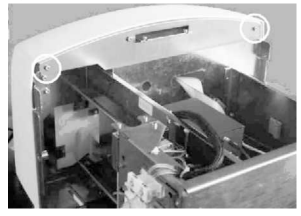






8. Unscrew the two hex screws shown in Figure 4.2-8.

#### Figure 4.2-8 Front Panel Screw Locations



9. Remove the flat connector attached to Keypad and LCD card (Figure 4.2-9).

Figure 4.2-9 Connector for the Keypad and LCD Card



- 10. Lift the front cover up and away slightly until the torx screws clear their holes then lift the cover off the instrument.
- 11. Set the cover aside in a safe place where it will not get damaged.

#### Installation

- 1. Position the front cover back on the instrument. Make sure the holes for the two torx screws are aligned with the screws that remained in the instrument.
- 2. Reattach the flat connector attached to Keypad and LCD card (Figure 4.2-9).
- 3. Replace the two hex screws located at the top inside the instrument (Figure 4.2-8).
- 4. Tighten the two torx screws shown in Figures 4.2-6 and 4.2-7.
- 5. Open the reagent door replace the four hex screws (Figure 4.2-5).
- 6. Reconnect the reagent tubing to its stopper and place the bottle back inside the reagent compartment (match the color coding on the bottle label with the label on the panel).

- 7. Replace the top cover. Under Removing the Front Cover, see the installation instructions.
- 8. Reconnect the power cord.
- 9. Turn the instrument on.

# 4.3 PREPARATION TO SHIP THE INSTRUMENT

### Purpose

Use this procedure to clean and properly prepare the instrument for shipping.

# **Tools/Supplies Needed**

- □ Fungicidal, bactericidal, virus killing detergent spray, non-corrosive for metals, non-plastic altering
- □ High quality, fragrance-free bleach (10-12% sodium hypochlorite available chloride)
- Distilled water
- □ Absorbent paper
- □ Two 500 mL glass beakers or flasks

# Bleach the Baths (20 minutes)

- 1. If OFF, turn the instrument ON.
- 2. From the Main Menu, select **4. DIAGNOSTICS → 3. DILUTER SYSTEMS → 4. EXTENDED CLEANING**. The Extended Cleaning prompt appears.
- 3. Press **ENTER** and the message CYCLE IN PROGRESS. PLEASE WAIT. . . appears.

**WARNING** Risk of contamination. If you do not properly shield yourself while decontaminating the instrument, you may become contaminated. To prevent possible biological contamination, you must wear appropriate safety glasses, a lab coat, and gloves when performing this procedure.

- 4. When the message POUR 3 mL OF EXTENDED CLEANING REAGENTS INTO BATHS appears,
  - a. Open the right side door.
  - b. Pour 3 mL of high quality, fragrance-free bleach (10-12% sodium hypochlorite available chloride) into each bath.
  - c. Close the right side door.
  - d. Press **ENTER** to continue. Allow the instrument to complete the cleaning procedure. It takes about five minutes for the cycle to complete. The system will automatically flush to remove the bleach poured in the baths earlier.
- 5. When the cycle is finished, turn the instrument OFF and unplug the power cord from the instrument or the wall.
- 6. Open the right side door.
- 7. Spray the bactericidal cleaner on all biohazard areas and wait for 10 minutes (assemblies in contact with the biohazard materials such as instrument cover, tube holder, keypad, start key, assemblies close to the sample probe, and so forth).

# **Clean the External Surfaces (20 minutes)**

- 1. Make a 1:5 bleach solution: 4 parts distilled water to 1 part high quality, fragrance-free bleach (10-12% sodium hypochlorite available chloride).
- 2. Clean the work area around the instrument.

**ATTENTION:** Do not use a sponge or cloth to clean instrument assemblies. Use absorbent paper towels that can be discarded in a biohazard container. For small or sensitive assemblies, use lint-free tissues.

- 3. Use a spray detergent to remove blood stains and salt marks from the following assemblies:
  - Outer surfaces of the instrument (covers, LCD, reagent locations)
  - Keypad
  - Waste connector plug
  - Liquid valve push button
  - Assemblies close to the sample probe
  - Tube holder assembly
  - Overflow trays

**ATTENTION:** Do not bleach stainless steel components if the ambient room temperature is more than  $30^{\circ}$  C ( $86^{\circ}$  F).

- 4. Disinfect, with the diluted bleach solution, all assemblies that have been in contact with biohazard materials.
- 5. Reinstall all the assemblies and set the instrument back to its initial configuration.

#### Clean the Tubing and Chambers (60 minutes)

#### Preparation

- 1. Locate two containers such as a glass beaker or flask that will hold a little more than 500 mL of liquid. Select containers that can be placed in front of the reagent compartment when the front door is open.
  - a. In one container, prepare approximately 500 mL of a 1:10 bleach solution: 9 parts distilled water to 1 part high quality, fragrance-free bleach (10-12% sodium hypochlorite available chloride).
  - b. Pour 500 mL of distilled water into the second container.
- 2. Turn the instrument ON.
- 3. Remove each pickup tube from its reagent container and place the tube in the diluted bleach. Don't forget the diluent pickup tube.
- 4. At the Main Menu, select **3**. **REAGENTS** → **3**. **PRIME** → **6**. **ALL REAGENTS** to pull the diluted bleach into the instrument.
- 5. Fill a glass or plastic tube with a 1:5 bleach solution: 4 parts distilled water to 1 part high quality, fragrance-free bleach (10-12% sodium hypochlorite available chloride).

#### **Cycle Routine**

- 1. From the Service menu, select **7. BURN-IN**.
- 2. Set the number of burn-in cycles to **15**.
- 3. Cycle the 1:5 bleach solution to initiate the burn-in function. The instrument will cycle 15 times. Do not attempt to stop the cycles. Let the instrument operate until it stops.

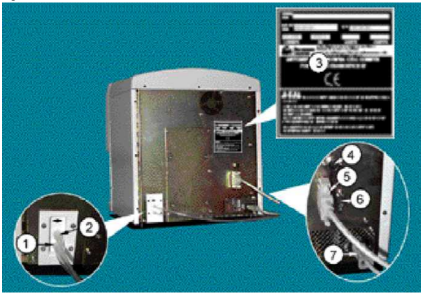
#### **Drain and Rinse**

- 1. Remove the reagent pickup tube assemblies from the diluted bleach and wrap the tubes in absorbent paper.
- 2. From the Main Menu, select **3**. **REAGENTS** → **3**. **PRIME** → **6**. **ALL REAGENTS** to drain the diluted bleach from the system.
- 3. When the prime cycles are complete, select **6**. **ALL REAGENTS** again to ensure the bleach is removed from the system.
- 4. Exchange the diluted bleach container with the vessel containing distilled water and place the pickup tubes in the distilled water.
- 5. From the Prime menu, select 6. ALL REAGENTS to rinse the system with distilled water.
- 6. Remove the reagent pickup tubes from the distilled water and wrap the tubes in absorbent paper.
- 7. From the Prime menu, select **6. ALL REAGENTS** to drain the distilled water from the system.
- 8. When the prime cycles are complete, select **6**. **ALL REAGENTS** again to ensure the distilled water is removed from the system.

**WARNING** When this process is complete, open the right side door and verify the DIFF bath is empty.

- 9. Turn the instrument OFF.
- 10. Remove the diluent input tubing (Figure 4.3-1, item 2).

Figure 4.3-1 Rear Panel Connections



**WARNING** Risk of contamination. If you do not properly shield yourself while removing the waste tubing, you may become contaminated. To prevent possible biological contamination, you must wear appropriate safety glasses, a lab coat, and gloves when removing this tubing.

- 11. Remove the waste output tubing (Figure 4.3-1, item 1).
- 12. Disconnect any cables attached to the back of the instrument including the:
  - a. Bar-code reader connector, if attached (Figure 4.3-1, item 4).
  - b. Printer connector (Figure 4.3-1, item 5).
  - c. RS232C output, if attached (Figure 4.3-1, item 6).
  - d. Power supply cable (Figure 4.3-1, item 7).
- 13. Verify that all four reagent bottles are removed from the reagent compartment.
- 14. Close all the instrument doors.
- 15. Pack the instrument in its original box, if available.

# 4.4 FLOW CELL CHECKS AND ADJUSTMENTS

#### Purpose

Use this adjustment procedure when the flow cell is moved or replaced, the DIFF tubing is changed, or the DiffPlot does not look correct. No adjustment should be required when the DIFF lamp is replaced.

To adjust the optical bench assembly, 5  $\mu$ L of RBC/PLT latex is aspirated and diluted in the DIFF bath with 2 mL of diluent. This dilution is then injected into the flow cell. Once the Diff Adjustment screen appears, measurements are continuously displayed for 27 seconds with updates occurring every 700 microseconds. Adjustments may be made during this 27 seconds as needed. Three audible beeps indicate the end of the adjustment period. Do not make adjustments once these beeps are heard. The screen is no longer being updated.

# **Tools/Supplies Needed**

- □ Allen wrench, 3 mm
- □ Plastic potentiometer adjustment tool, PN 5415364
- □ RBC/PLT latex, PN LAD002AS
- □ Several fresh whole-blood specimens

### **Preparation**

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- 1. If off, turn the instrument on.
- From the Main Menu, select 4. DIAGNOSTICS → 3. DILUTER SYSTEM → 2. RINSE →
   2. FLOWCELL to remove air bubbles clinging to the inner optical surfaces.
- 3. Press **ESC** twice to return to the Diagnostics menu then select **5**. **SERVICE**.

#### **Flow Cell Checks**

- 1. From the Service menu, select **2. MEASUREMENT → 5. DIFF ADJUSTMENT**.
- 2. When the DIFF ADJUSTMENT prompt appears, press ENTER to continue.

**IMPORTANT** Risk of misleading results. The RBC/PLT latex particles tend to clump as they settle out of solution. Clumped latex particles will affect adjustment results. Mix the RBC/PLT latex vigorously before use. A vortex may be used. Remix the latex thoroughly before each sampling.

- 3. Mix the RBC/PLT latex vigorously. Use a vortex, if available.
- 4. When the *PLEASE SAMPLE LATEX* prompt appears, present the vial of well-mixed latex particles to the probe and press the aspirate switch to initiate aspiration and the routine.
- 5. When the Diff Adjustment screen appears (Figure 4.4-1), verify each result is acceptable according to the following criteria:

Figure 4.4-1 Diff Adjustment Screen

| DIFF ADJUSTMENT          | 1/27/00 16:05 PM |
|--------------------------|------------------|
| DIFF LAMP                | 6.20 VOLTS       |
| 4.00<br>TRANSFER TIME    | 8.00             |
|                          | 212              |
| 100<br>RESISTIVE CHANNEL | 300              |
|                          | 50               |
| 5<br>ABSORBANCE CHANNEL  | 70               |
|                          | 180              |
| 10                       | 240              |
| CYCLE IN PROGRESS        |                  |
|                          | 56%              |
|                          |                  |

- 6. Verify the *DIFF LAMP* value displayed on the screen is between **5.50** and **6.50**.
  - If the value on the screen is within the acceptable range, go to step 7.
  - If the value on the screen is outside this acceptable range or or if you desire a value closer to the target value (6.0), go to the DIFF Lamp Voltage Adjustment heading.
- 7. Verify the *TRANSFER TIME* value is between **150** and **250**.
  - If the value on the screen is within the acceptable range, go to step 8.
  - If the *TRANSFER TIME* value is outside the acceptable range, there is a problem. Complete the instructions under Heading 4.36, OPTICAL BENCH PRELIMINARY ADJUSTMENTS before proceeding.

**Note:** When cells are not optically detected, the *TRANSFER TIME* value displayed on the screen is 100 and backlighted. The backlighted number indicates the value is less than 100, and may actually be zero. If the flow cell is so far out of alignment that no cells are detected, the procedure under Heading 4.36 provides a series of preliminary adjustments for the flow cell and optics lamp.

When the *TRANSFER TIME* value is greater than 100 but less than 149, the timing is outside the acceptable range but the number greater than 100 indicates that cells are definitely being detected. The *TRANSFER TIME* is controlled by the height of the flow cell relative to the level of the light beam. Since the light beam is at a fixed height with no vertical adjustment knob for the flow cell, this is a difficult adjustment. The height of the flow cell is controlled by the spacers or shims placed under the flow cell mounting block. Each shim moves the flow cell up a distance that equates to a 30 microsecond drop in the *TRANSFER TIME* value. A new flow cell is shipped with matching shims that should set it to the proper height. As a result, it is critical that these shims are always replaced anytime the flow cell is removed and replaced.

- 8. Verify the RESISTIVE CHANNEL value is between 45 and 55.
  - If the value on the screen is within the acceptable range, go to step 9.
  - If the value on the screen is outside this acceptable range or if you desire a value closer to the target value (50), go to the Resistive Channel Adjustment heading.
- 9. Verify the ABSORBANCE CHANNEL value is greater than **170**.
  - If the value on the screen is acceptable, go to the Final Verification heading at the end of this section.
  - If the value on the screen is less than 170 or if you desire a higher value, go to the Absorbance Channel Adjustment heading.

# **DIFF Lamp Voltage Adjustment**

**Note:** If you are only interested in the *DIFF LAMP* voltage, it it not necessary to aspirate latex. However, if you change the lamp voltage, the Transfer Time, Resistive Channel, and Absorbance Channel values must be checked using latex particles.

#### Preparation

- 1. Remove the left side panel and the top cover from the instrument. For details, see Heading 4.2. Set the left side panel and top cover aside.
- 2. Locate potentiometer R11 on the optical bench assembly (Figures 4.4-2 and 4.4-3). This is the potentiometer you will use to adjust the DIFF lamp voltage.

#### Figure 4.4-2 Potentiometer R11 - Location on the Optical Bench Assembly

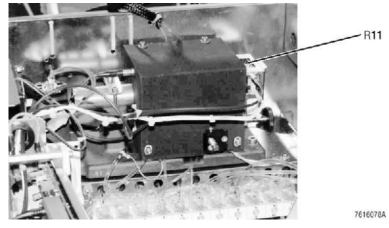
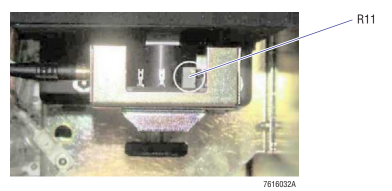


Figure 4.4-3 Potentiometer R11 Location - Top View from the Front of the Instrument



#### Adjustment

1. From the Service menu, select **2. MEASUREMENT → 5. DIFF ADJUSTMENT**.

**Note:** If the Diff Adjustment screen is currently displayed, press **ESC** to return to the Measurement Menu then select **5**. **DIFF ADJUSTMENT**.

2. When the *DIFF ADJUSTMENT* prompt appears, press **ENTER** to continue.

**IMPORTANT** Risk of misleading results. The RBC/PLT latex particles tend to clump as they settle out of solution. Clumped latex particles will affect adjustment results. Mix the RBC/PLT latex vigorously before use. A vortex may be used. Remix the latex thoroughly before each sampling.

- 3. Mix the RBC/PLT latex vigorously. Use a vortex, if available.
- 4. When the *PLEASE SAMPLE LATEX* prompt appears, present the vial of well-mixed latex particles to the probe and press the aspirate switch to initiate aspiration and the routine
- 5. When the Diff Adjustment screen appears, monitor the *DIFF LAMP* value on the screen. 6.00 Vdc is the target value. The acceptable range is 5.50 to 6.50 Vdc.
- 6. If a change is desired, adjust potentiometer R11 (Figure 4.4-3) to produce a readout of 6.00 Vdc.

**Note:** Measurements are continuously displayed for 27 seconds. The readout updates every 700 microseconds. Do not continue to make adjustments after the three audible beeps.

- 7. If more time is needed to make the adjustment,
  - a. When the *CYCLE IN PROGRESS* timing bar disappears, press **ESC** to return to the Measurement menu.
  - b. At the Measurement menu, select 5. DIFF ADJUSTMENT.
  - c. When the DIFF ADJUSTMENT prompt appears, press ENTER to continue.
  - d. Mix the RBC/PLT latex vigorously. Use a vortex, if available.
  - e. When the *PLEASE SAMPLE LATEX* prompt appears, present the vial of well-mixed latex particles to the probe and press the aspirate switch to initiate aspiration and the routine.
  - f. When the Diff Adjustment screen appears, the monitoring and adjustment process can be continued for another 27 seconds until the three beeps sound.

#### **Interim Verification**

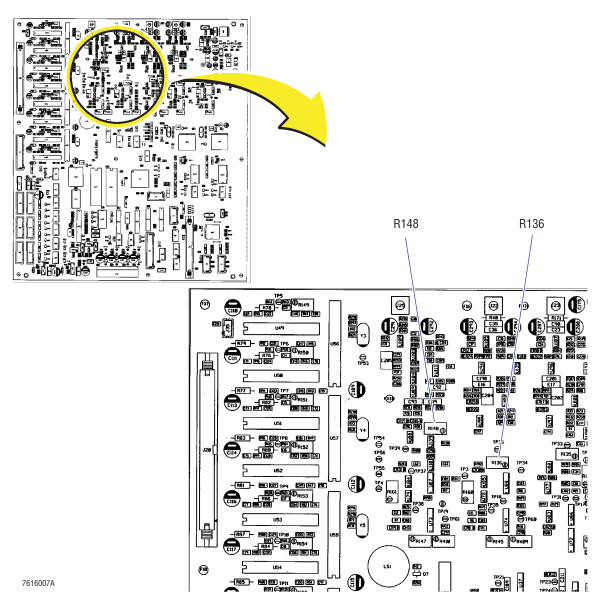
When the *DIFF LAMP* value is within the acceptable range of 5.50 to 6.50 Vdc, return to the Flow Cell Checks heading to verify this new setting and make sure all the other values are acceptable.

# **Resistive Channel Adjustment**

# Preparation

- 1. If you have not already done so, remove the left side panel from the instrument. For details, see Heading 4.2. Set the panel aside.
- 2. Locate potentiometer R136 on the Main card. See Figure 4.4-4.

#### Figure 4.4-4 Main Card Flow Cell Adjustments



#### Adjustment

1. From the Service menu, select **2. MEASUREMENT → 5. DIFF ADJUSTMENT**.

Note: If the Diff Adjustment screen is currently displayed, press **ESC** to return to the Measurement Menu then select **5**. **DIFF ADJUSTMENT**.

2. When the *DIFF ADJUSTMENT* prompt appears, press **ENTER** to continue.

**IMPORTANT** Risk of misleading results. The RBC/PLT latex particles tend to clump as they settle out of solution. Clumped latex particles will affect adjustment results. Mix the RBC/PLT latex vigorously before use. A vortex may be used. Remix the latex thoroughly before each sampling.

- 3. Remix the RBC/PLT latex vigorously. Use a vortex, if available.
- 4. Present the vial of well-mixed latex particles to the probe and press the aspirate switch to initiate aspiration and the adjustment routine.
- 5. When the Diff Adjustment screen appears, monitor the *RESISTIVE CHANNEL* value on the screen. 50 is the target value. The acceptable range is 45 to 55.

**Note:** If the values are fluctuating, bubbles may be entering the flow cell. Purge the flow cell as follows:

From the Main Menu, select **4**. **DIAGNOSTICS** → **3**. **DILUTER SYSTEM** → **2**. **RINSE** → **2**. **FLOWCELL** to remove air bubbles clinging to the inner optical surfaces.

6. If a change is desired, adjust potentiometer R136 until the channel is set at 50.

**Note:** Measurements are continuously displayed for 27 seconds. The readout updates every 700 microseconds. Do not continue to make adjustments after the three audible beeps.

- 7. If more time is needed to make the adjustment,
  - a. When the *CYCLE IN PROGRESS* timing bar disappears, press **ESC** to return to the Measurement menu.
  - b. At the Measurement menu, select **5. DIFF ADJUSTMENT**.
  - c. When the DIFF ADJUSTMENT prompt appears, press ENTER to continue.
  - d. Mix the RBC/PLT latex vigorously. Use a vortex, if available.
  - e. When the *PLEASE SAMPLE LATEX* prompt appears, present the vial of well-mixed latex particles to the probe and press the aspirate switch to initiate aspiration and the routine.
  - f. When the Diff Adjustment screen appears, the monitoring and adjustment process can be continued for another 27 seconds until the three beeps sound.

#### **Interim Verification**

When the *RESISTIVE CHANNEL* value is within the acceptable range of 45 to 55, return to the Flow Cell Checks heading to verify this new setting and make sure all the other values are acceptable.

# Absorbance Channel Adjustment

During this adjustment, the flow cell is repositioned so that the focal point of the light beam (which is fixed) is properly positioned inside the flow cell. The goal is to maximize the absorbance using a two-positional adjustment.

### Preparation

- 1. If you have not already done so, remove the left side panel and the top cover from the instrument. For details, see Heading 4.2. Set the left side panel and top cover aside.
- 2. At the optical bench assembly, locate the front adjustment knob (Figure 4.4-5) and the side adjustment screw (Figure 4.4-6).
  - The Absorbance Channel adjustment is made using the front knob (Figure 4.4-5) or side screw (Figure 4.4-6) to move the flow cell.
  - An adjustment to move the flow cell along the Y-axis (front or back) is made using the front knob.
  - An adjustment to move the flow cell along the X-axis (right or left) is made using the side screw.

Figure 4.4-5 Front Adjustment Knob - Optical Bench

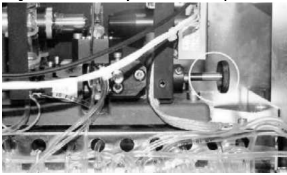
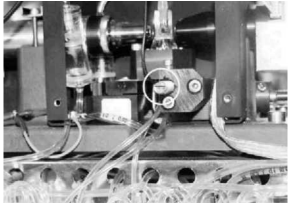


Figure 4.4-6 Side Adjustment Screw - Optical Bench



#### Adjustment

1. From the Service menu, select **2. MEASUREMENT → 5. DIFF ADJUSTMENT**.

Note: If the Diff Adjustment screen is currently displayed, press **ESC** to return to the Measurement Menu then select **5**. **DIFF ADJUSTMENT**.

2. When the DIFF ADJUSTMENT prompt appears, press ENTER to continue.

**IMPORTANT** Risk of misleading results. The RBC/PLT latex particles tend to clump as they settle out of solution. Clumped latex particles will affect adjustment results. Mix the RBC/PLT latex vigorously before use. A vortex may be used. Remix the latex thoroughly before each sampling.

- 3. Remix the RBC/PLT latex vigorously. Use a vortex, if available.
- 4. Present the vial of well-mixed latex particles to the probe and press the aspirate switch to initiate aspiration and the adjustment routine.

- 5. When the Diff Adjustment screen appears, verify the *ABSORBANCE CHANNEL* value is greater than 170.
- 6. If a change is desired, use the front knob (Figure 4.4-5) and the side screw (Figure 4.4-6) to adjust the channel higher than 170.

Note: Measurements are continuously displayed for 27 seconds. The readout updates every 700 microseconds. Do not continue to make adjustments after the three audible beeps. If you have difficulties adjusting the absorbance high enough and the *DIFF LAMP* voltage value is acceptable, use potentiometer R148 on the Main card (Figure 4.4-4) to increase the gain. Final adjustment of potentiometer R148 is done during verification with fresh, normal whole-blood specimens.

- 7. If more time is needed to make the adjustment,
  - a. When the *CYCLE IN PROGRESS* timing bar disappears, press **ESC** to return to the Measurement menu.
  - b. At the Measurement menu, select **5**. **DIFF ADJUSTMENT**.
  - c. When the *DIFF ADJUSTMENT* prompt appears, press **ENTER** to continue.
  - d. Mix the RBC/PLT latex vigorously. Use a vortex, if available.
  - e. When the *PLEASE SAMPLE LATEX* prompt appears, present the vial of well-mixed latex particles to the probe and press the aspirate switch to initiate aspiration and the routine.
  - f. When the Diff Adjustment screen appears, the monitoring and adjustment process can be continued for another 27 seconds until the three beeps sound.

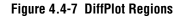
#### **Interim Verification**

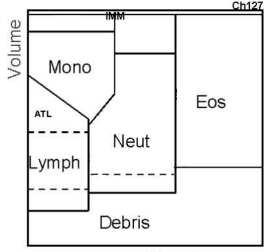
When the *ABSORBANCE CHANNEL* value is greater than 170, return to the Flow Cell Checks heading to verify this new setting and make sure all the other values are acceptable.

# **Final Verification**

When all adjustments are complete,

- 1. If the optical bench cover was removed, replace the cover and secure it with the four screws removed earlier.
- 2. Make sure the right side door is closed.
- 3. Cycle several fresh normal whole-blood specimens (five different specimens, if possible) and verify the DiffPlot looks acceptable. It is particularly important that the lymphocyte and neutrophil population positions (Figure 4.4-7) meet the following criteria:





Absorbance

• The lymphocyte population must fall between the dotted lines representing normal lymphocytes (area labeled Lymph). Very few, if any, cells should be located in the small lymphocyte area (unlabeled area under the lower dotted line) or in the atypical lymphocyte area (area labeled ATL above the upper dotted line). See Figure 4.4-7.

Move the lymphocyte population up or down by adjusting potentiometer R136 as needed (Figure 4.4-4). Base this resistive channel adjustment on several bloods, just in case the population of a specific blood is not as normal as you think.

• The vertical line separating the lymphocyte (area labeled Lymph) and neutrophil (Neut) populations must bisect those two populations. See Figure 4.4-7.

If needed, use potentiometer R148 (Figure 4.4-4) to adjust the absorbance channel until the two populations are distinct and separated by the line.

- 4. If it was necessary to adjust either the resistive channel or the absorbance channel, recheck the diff adjustments as follows:
  - a. From the Service menu, select **2. MEASUREMENT → 5. DIFF ADJUSTMENT**.

**Note:** If the Diff Adjustment screen is currently displayed, press **ESC** to return to the Measurement Menu then select **5**. **DIFF ADJUSTMENT**.

b. When the *DIFF ADJUSTMENT* prompt appears, press **ENTER** to continue.

**IMPORTANT** Risk of misleading results. The RBC/PLT latex particles tend to clump as they settle out of solution. Clumped latex particles will affect adjustment results. Mix the RBC/PLT latex vigorously before use. A vortex may be used. Remix the latex thoroughly before each sampling.

- c. Mix the RBC/PLT latex vigorously. Use a vortex, if available.
- d. When the *PLEASE SAMPLE LATEX* prompt appears, present the vial of well-mixed latex particles to the probe and press the aspirate switch to initiate aspiration and the routine.
- e. Verify the channel results appearing on the Diff Adjustment screen are acceptable:
  - *RESISTIVE CHANNEL* value is between **45** and **55**.
  - *ABSORBANCE CHANNEL* value is greater than **170**.

Note: If any value falls outside the acceptable range, the whole-blood specimens may not be normal or there may be an instrument problem affecting the DiffPlot. Timing, dilutions, noise, bubbles, or reagent can affect performance. Troubleshoot as needed.

- 5. When system performance is acceptable,
  - a. Carefully close the Main card door. Turn the two captive knobs clockwise to secure the door.
  - b. Replace the top cover if it was removed. See Heading 4.2 as needed.

**ATTENTION:** When replacing the left side panel with the instrument powered on, avoid accidently turning the instrument off again by carefully positioning the opening for the power on/off rocker switch over the switch as you position the panel on the instrument frame.

- c. Replace the left side panel and install the four hex screws that secure it to the instrument frame.
- 6. Perform a system verification. See Heading 5.3, SYSTEM VERIFICATION PROCEDURES.

# 4.5 BATHS ASSEMBLY ALIGNMENT CHECK AND ADJUSTMENT

#### Purpose

Use this procedure to properly position the baths assembly anytime the assembly is either moved or replaced. Avoid any unnecessary contact with the sample probe. The probe is sharp and may contain biohazardous materials.

# **Tools/Supplies Needed**

- □ Allen wrench, 3 mm
- □ Feeler gauges, optional
- □ Flashlight, optional

# Preparation

- 1. If off, turn the instrument on.
- 2. From the Main Menu, select 4. DIAGNOSTICS → 4. HARDWARE SYSTEMS → 4. TRAVERSE SERVICE POSITION.
- 3. When the *TRAVERSE SERVICE POSITION* prompt appears, press **ENTER** to retract the probe and move the housing over the bath assembly.
- 4. Turn the instrument off and unplug the power cord from the instrument or the wall.
- 5. Open the pneumatic access door (right side of the instrument).

# **Alignment Check**

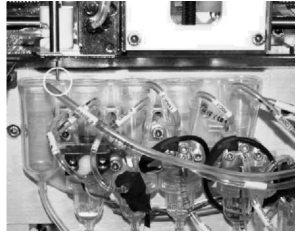
**WARNING** Risk of personal contamination. If you do not properly shield yourself while servicing the baths assembly, you may become contaminated. To prevent possible biological contamination, you must wear appropriate safety glasses, a lab coat, and gloves when servicing the baths assembly.

1. Using the horizontal traverse belt, manually position the sample probe housing over the inside rim of the rinse bath.

2. Gently push down on the top of the sample probe until the tip of the probe rests on the inside rim of the rinse bath (Figure 4.5-1 and Figure 4.5-2).

Figure 4.5-1 Sample Probe Position at the Rinse Bath

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#### Figure 4.5-2 Close-up of Probe Position



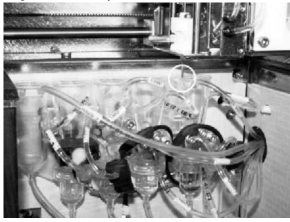
**CAUTION** Risk of damage to the sample probe. If power is restored to the instrument and the baths assembly is mounted too high, the sample probe may become bent when it hits the bath edges as it moves from bath to bath. Do not restore instrument power until the baths assembly is aligned.

3. Verify the tip of the sample probe evenly clears the top of each bath:

**ATTENTION:** If you have removed and replaced the baths assembly, be careful that you do not bend the probe as you move it towards the WBC/BASO bath.

- a. Without lifting the probe, gently push the probe housing towards the inside edge of the last bath, the WBC/BASO bath.
- b. Note the distance between the tip of the probe and the top of each bath as you move towards the WBC/BASO bath (Figures 4.5-3 and 4.5-4).

Figure 4.5-3 Sample Probe at the WBC/BASO Bath



4. If the distance differs, do the Alignment Adjustment that follows.

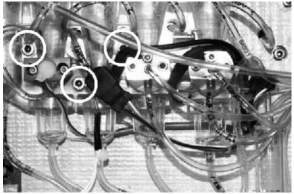


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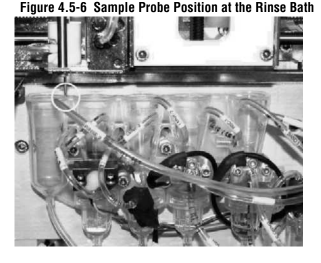
# **Alignment Adjustment**

1. Loosen, but do not remove, the three hex screws securing the baths assembly to the support panel (Figure 4.5-5). Use a 3 mm Allen wrench.

#### Figure 4.5-5 Location of Screws Securing the Baths Support Panel



- 2. Manually push the sample probe housing to the outside edge of the rinse bath. Move the baths assembly up or down as necessary until the tip rests gently on the outside rim of the rinse bath.
- 3. While continuing to push the sample probe housing over the rinse bath to the inside edge of the bath, adjust the baths assembly up or down as needed so that the sample probe tip comes to rest gently on the inside rim of the rinse bath as shown in (Figures 4.5-6 and 4.5-7).





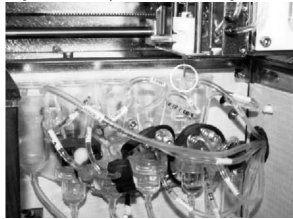


- 4. Gently tighten the center and left screws (See Figure 4.5-5).
- 5. Without lifting the probe, gently push the probe housing backwards towards the inside edge of the last bath, the WBC/BASO bath. Reposition the baths assembly as needed to clear the edge of each bath.

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6. Verify the sample probe is positioned at the WBC/BASO bath as shown in Figure 4.5-8 and Figure 4.5-9.

Figure 4.5-8 Sample Probe Position, Right Side



# Figure 4.5-9 Close-up of Acceptable Probe Position



- 7. Gently tighten the right screw on the baths assembly (See Figure 4.5-5).
- 8. Make sure all three screws are tight so that the baths assembly is secure on the instrument's frame.
- 9. Carefully move the sample probe housing over the outer edge of the rinse bath. The probe should equally clear all baths. If not, loosen the nearest screw and reposition the baths assembly until the distance between the tip of the probe and the top of each bath is the same.

#### Verification

1. Go back to the Alignment Check heading and perform this check to verify proper alignment.

**CAUTION** It is mandatory to perform the probe adjustment procedure after this adjustment.

- 2. When the adjustment is correct
  - a. Reconnect the power cord.
  - b. Turn the instrument on. An automatic startup and background check are performed.
     Note: When the startup routine and background check are done, the sample probe will be back in its home position.
  - c. Go to Heading 4.6, SAMPLE PROBE CHECKS AND ADJUSTMENTS and perform the probe adjustment procedure.

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# 4.6 SAMPLE PROBE CHECKS AND ADJUSTMENTS

### Purpose

**IMPORTANT** Check the baths assembly alignment before checking or adjusting the sample probe position. See Heading 4.5, BATHS ASSEMBLY ALIGNMENT CHECK AND ADJUSTMENT.

Use this procedure to check the sample probe home position and the sample probe position inside each bath. An adjustment procedure is provided after the check procedures. The check procedures may be done at any time, but the adjustment procedures should be performed only if the check fails.

From the Service menu, selecting **1**. **DILUTION** provides the options needed to adjust the traverse and sample probe (Figure 4.6-1.)

Figure 4.6-1 Dilution Screen

| DILUTION  | 1           | /27/00   16:05 PM               |
|---|-------------|---------------------------------|
| PROBE HOME<br>TRAVERSE HOME                           | 35<br>20    | RUN                             |
| PROBE POS.<br>TRAVERSE POS.                           | 476<br>1132 | RUN                             |
| WBC / BASO TRAVE<br>FLOWCELL TRAVE<br>RBC TRAVERSE PC | RSE POS.    | 1060<br>737<br>734<br>RUN CYCLE |
|   |             | dilution.eps                    |

- **PROBE HOME**: Allow adjustment to get the best probe extremity cleaning.
- **TRAVERSE HOME**: Default value is 20 and does not need to be changed.
- **PROBE POS.**: Probe up/down adjustment.
- **TRAVERSE POS.**: Probe left/right position based on Traverse Home.
- **RUN**: Upper prompt positions the probe to determine if the tip of the sample probe is properly positioned 9.4 mm from its guide.

- **RUN**: Lower prompt positions the probe to determine if the tip of the sample probe is properly aligned with the DIFF bath at port 3.
- **RUN CYCLE**: Selection initiates a routine that provides a way to determine if the tip of the sample probe is properly aligned inside each of the baths. This routine is similar to an actual sample cycle.
- WBC/BASO TRAVERSE POS.: Probe left/right position in the WBC/BASO bath relative to the DIL1/HGB bath position. Adjusts distance from DIL1/HGB bath to WBC/BASO bath.
- **FLOWCELL TRAVERSE POS**.: Probe left/right position in the DIFF bath relative to the WBC/BASO bath. Adjusts distance from WBC/BASO bath to DIFF bath.
- **RBC TRAVERSE POS.**: Probe left/right position in the RBC bath relative to the DIFF bath. Adjusts distance from DIFF bath to RBC bath.

# **Tools/Supplies Needed**

- □ Feeler adjustment gauge, 9.4 mm, or a sturdy piece of paper cut exactly 9.4 mm (0.37 in.) wide and approximately 76.2 mm (3.0 in.) long
- □ Plastic transfer pipet
- □ Flashlight, optional
- □ Jeweler's loop (x5 power magnifier with 2-inch focal length), optional

# **Sample Probe Checks**

Use this procedure to check the home position of the sample probe and the probe's position inside each bath.

#### **Home Position Check**

- 1. From the Service menu, select **1. DILUTION**.
- 2. At the Dilution screen, select the upper *RUN* option and press **ENTER**.
- 3. Open the right side door
- 4. Verify the distance between the tip of the sample probe and the traverse is about 9.4 mm. Use either the feeler gauge or the 9.4 mm side of the paper you may have prepared. See Figures 4.6-2 and 4.6-3.

# Figure 4.6-2 Acceptable Distance between the Sample Probe Tip and the Traverse

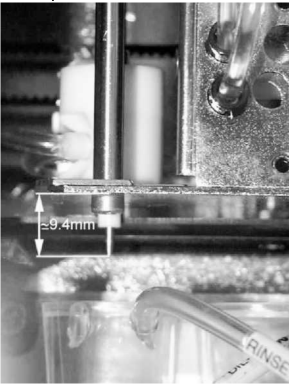


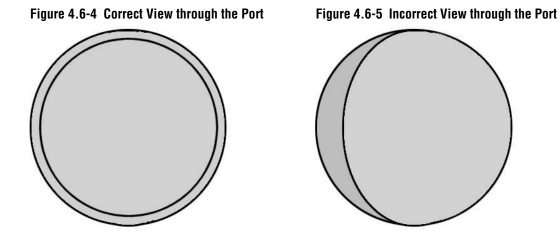
Figure 4.6-3 Measuring the Distance between the Sample Probe Tip and the Traverse



- If the distance is approximately 9.4 mm, press **ESC** then proceed to the heading Inside Bath Position Check.
- If the distance is unacceptable, continue to the Home Position Adjustment heading to perform the adjustment procedure.

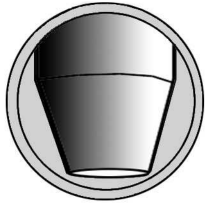
#### **Inside Bath Position Check**

- 1. At the Dilution screen, select the *RUN CYCLE* option at the bottom of the screen.
- 2. Remove the tubing connected to port 1 of the DIL1/HGB bath, port 3 of the DIFF bath, port 1 of the RBC bath and port 2 of the WBC/BASO bath. These ports are located on the right side of each bath about 1cm (1/2 inch) from the top.
- 3. Clean the ports of any fluid that may affect your line of sight through the port. A plastic transfer pipet may be used to clear the ports.
- 4. With the following considerations in mind, evaluate the position of the probe tip in relation to the bath port.
  - Consider using a jeweler's loop (x5 power magnifier with 2-inch focal length) and a flashlight. These tools greatly enhance the image.
  - To make a proper judgement, your eyes must be correctly positioned.
  - The correct position for your eyes is when the port looks like Figure 4.6-4. If the circles are not centered (as in Figure 4.6-5), reposition your gaze until the circles are centered (as in Figure 4.6-4).



• As you gaze through the port, the tip of the sample probe should mimic the position of the tip shown in Figure 4.6-6, in each bath. The first bath is the DIL1/HGB bath. Press **ENTER** to advance to the WBC/BASO bath and then again for the DIFF bath and the RBC bath. Check that the probe position is correct in each bath.

Figure 4.6-6 Properly Adjusted Sample Probe Tip



5. If the probe is properly adjusted for all baths, reconnect the tubing to the bath ports. If the probe is out of adjustment, continue on to the heading Inside Bath Position Adjustment and follow the instructions to properly adjust the position of the probe.

# Sample Probe Adjustments

Use this procedure to adjust the home position of the sample probe and the probe's position inside each bath.

#### **Home Position Adjustment**

**CAUTION** Check the probe position using the RUN functions before making an adjustment.

- 1. Thoroughly clean the exterior of the probe before making any adjustments.
- 2. At the Dilution screen, select the upper *RUN* option and press **ENTER**.
- 3. Use the 9.4 mm side of the paper to recheck the distance between the tip of the sample probe and the traverse to confirm the gap is incorrect. See Figures 4.6-2 and 4.6-3.
- 4. Press **ESC** to cancel the previous function.
- 5. At the Dilution screen,
  - a. Adjust the PROBE HOME value as needed.
    - If probe tip is too low (gap >9.4 mm), increase the PROBE HOME value.
    - If probe tip is too high (gap <9.4 mm), decrease the PROBE HOME value.
  - b. Select the upper *RUN* option and press **ENTER** to rerun the routine.
- 6. Use the feeler gauge or the 9.4 mm side of the paper to recheck the distance between the tip of the sample probe and the traverse.
  - If the gap is now correct, press **ESC** then proceed to the heading Inside Bath Position Adjustment.
  - If the gap is still incorrect, repeat steps 4 through 6 until the gap is approximately 9.4 mm.

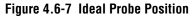
#### **Inside Bath Position Adjustment**

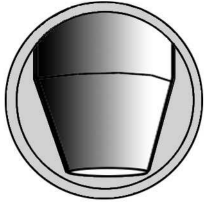
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**IMPORTANT** Do not perform this probe adjustment procedure unless it is absolutely necessary.

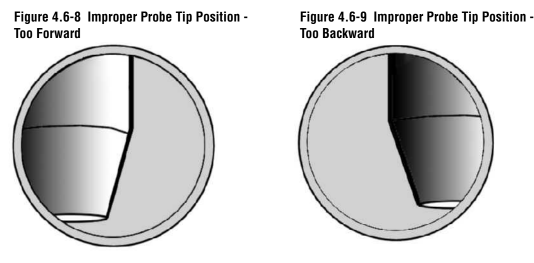
1. At the Dilution screen, highlight the RUN CYCLE option and press ENTER.

**ATTENTION:** As you complete this procedure, it is imperative that as you gaze through the port the circles are centered. When the sample probe is properly adjusted, the tip of the probe should mimic the view shown in Figure 4.6-7.



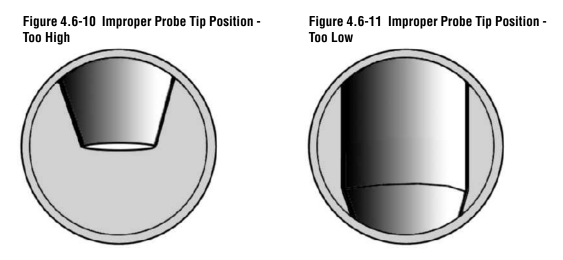


- 2. The first bath that the probe stops at is the DIL1/HGB bath. Evaluate the position of the probe tip in relation to the bath port.
  - Is it a horizontal positioning problem? Is the probe is too forward (Figure 4.6-8) or too backward (Figure 4.6-9)?



Note: The TRAVERSE POS. value moves the sample probe to the left or right.

• Is it a vertical positioning problem? Is the probe too high (Figure 4.6-10) or too low (Figure 4.6-11)?



Note: The PROBE POS. value moves the sample probe up or down.

- 3. Determine if changes to the position are required. If they are, press **ENTER** several times to advance through the cycle. At the Dilution screen, adjust the *PROBE POS*. or the *TRAVERSE POS*. values as needed.
  - The *TRAVERSE POS*. value moves the sample probe to the left or right (horizontal positioning).
    - If the probe is too forward (Figure 4.6-8), increase *TRAVERSE POS*. value.
    - ▶ If the probe is too backward (Figure 4.6-9), decrease *TRAVERSE POS*. value.
  - The PROBE POS. value moves the sample probe up or down (vertical positioning).
    - If the probe is too high (Figure 4.6-10), increase the PROBE POS. value.
    - If the probe is too low (Figure 4.6-11), decrease the PROBE POS. value.
- 4. Highlight and ENTER the RUN CYCLE function again.
- 5. If the probe position in the DIL1/HGB bath is acceptable, press the **ENTER** key once to advance to the WBC/BASO bath.
- 6. Check the probe position in the WBC/BASO bath. At this point, there should be no problem with height. If there is, the bath assembly is not level and proceed to the heading BATHS ASSEMBLY ALIGNMENT CHECK AND ADJUSTMENT. When level, start this procedure again at step 1.
  - If the lateral probe position is not correct, adjust the WBC/BASO Traverse Position value. You must press **ENTER** several times and advance through the cycle to make any changes. Increase the number to move the probe farther back, or to the right, decrease to move frontward of to the left. Note that it is rare to change from the default value of 1060.
  - Any changes to this value will almost certainly require changes to the flow cell traverse position and the RBC traverse position, since these are relative moves from the previous bath position.
- 7. Highlight and ENTER the RUN CYCLE function again.
- 8. Advance to the WBC/BASO bath, if not already there, and check the probe position in each bath. If no changes are required, press **ENTER** to advance to the DIFF bath.

- 9. Check the position in the DIFF bath. The default value is 737 and is usually acceptable. If changes are required, press **ENTER** several times to advance through the cycle. Since the probe was moving back to front, increasing this number moves the probe towards the front or to the left, decreasing the number moves the probe towards the back or to the right. If there were no changes, go to step 11.
- 10. Highlight and ENTER the RUN CYCLE function again.
- 11. Advance to the DIFF bath, if not already there, checking the probe position in each bath. If the positions are acceptable, and no further changes are made, press **ENTER** and advance to the RBC bath.
- 12. Check the position in the RBC bath. The default value of 734 is usually good. If a change is required, press **ENTER** several times to exit the cycle and adjust the RBC Traverse Position value as required. Highlight and **ENTER** the *RUN CYCLE* function again. Check the probe position in each bath.

**WARNING** Risk of contamination. If one or more tubings are left unattached to their respective bath ports and you run a cycle, fluid will extrude out the ports onto the exterior of the baths, the counting heads, and valves. It may result in the need for extensive cleanup inside the bath enclosure. To prevent possible biological contamination, you must wear appropriate safety glasses, a lab coat, and gloves.

**CAUTION** Risk of component damage to electronic components. If one or more tubings are left unattached to their respective bath ports and you run a cycle, fluid will extrude out the ports into the bath enclosure. Excessive moisture may damage the Hgb circuitry and solenoid valves. Thoroughly wipe dry electronic components as quickly as possible.

**IMPORTANT** Risk of misleading results. If one or more tubings are left unattached to their respective bath ports and you run a cycle, fluid will extrude out the ports onto the exterior of the baths, the counting heads, and valves. Moisture on these components affects parameter results. Thoroughly wipe components dry before reporting patient results.

- 13. When no changes are required, replace each tubing back on its designated bath port.
- 14. Perform a system verification. See Heading 5.3, SYSTEM VERIFICATION PROCEDURES.

**SERVICE AND REPAIR PROCEDURES** SAMPLE PROBE CHECKS AND ADJUSTMENTS

### 4.7 HGB BLANK ADJUSTMENT

### Purpose

Use this procedure to adjust the Hgb blank voltage. This adjustment must be made under stabilized thermal conditions.

### **Tools/Supplies Needed**

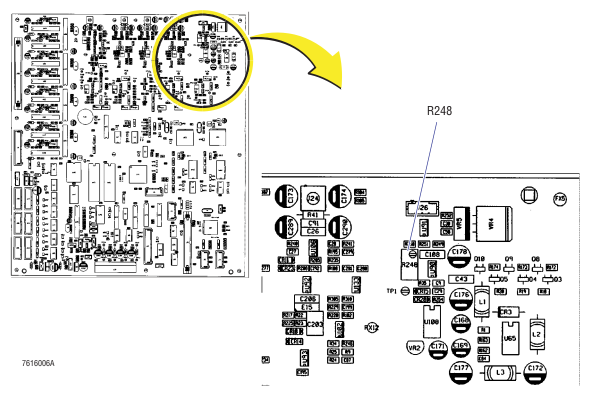
- □ Allen wrench, 3 mm
- □ Plastic potentiometer adjustment tool, PN 5415364

### Preparation

**IMPORTANT** The Hgb blank voltage adjustment must be made under stabilized thermal conditions. Make sure the right side door is closed before starting this adjustment. After completing a Startup, wait at least 5 minutes before performing this adjustment procedure.

- 1. Close the right side door to stabilize the temperature inside the bath enclosure.
- 2. Remove the left side panel to access the Main card. For details, see Heading 4.2.
- 3. Locate potentiometer R248 in the upper right quadrant of the Main card (Figure 4.7-1). This is the adjustment potentiometer.





### Adjustment

- 1. From the Service menu, select **2**. **MEASUREMENT → 1**. **HGB BLANK ADJUSTMENT.**
- 2. When the HGB BLANK ADJUSTMENT prompt appears, press ENTER to continue.

**Note:** Selecting this option initiates a routine that drains and rinses the DIL1/HGB bath and continuously displays (for 20 seconds) the Hgb blank voltage used by the converter. Three audible beeps signal the end of the 20 seconds.

3. While monitoring the Hgb Adjustment screen, adjust potentiometer R248 until the voltage reads 4.7 Vdc ±0 Vdc.

### Verification

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- 1. Run a Startup and verify the Hgb blank result is acceptable.
- 2. Run a whole-blood sample to establish a new Hgb reference blank.
- 3. Replace the left side panel.

### 4.8 APERTURE CURRENT CHECK

### Purpose

Use this procedure to verify the voltages needed to generate aperture current are present and at sufficient levels. These voltages are not adjustable.

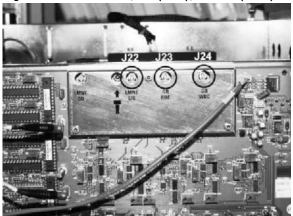
### **Tools/Supplies Needed**

- □ Allen wrench, 3 mm
- Digital voltmeter
- Old coax

### Procedure

- 1. Open the left side panel to access the Main card. For details, see Heading 4.2.
- 2. On the Main card, disconnect the LMNE CIS (for DIFF analysis), GR RBC, and GB WBC coaxes (Figure 4.8-1).

### Figure 4.8-1 LMNE CIS, GR (RBC), and GB (WBC) Coax Locations



- 3. From the Service menu, select **2**. MEASUREMENT → **2**. APERTURE CURRENT.
- 4. When the Aperture Current prompt appears, press ENTER.

**CAUTION** Do not damage connectors while measuring.

**IMPORTANT** Use an old coax that you have previously cut to check these voltages with the voltmeter. A flow cell coaxial cable with the T- connector removed is ideal.

- 5. When the *CHECK APERTURE CURRENT* prompt appears, use a DVM to verify the voltages at J22, J23 and J24 (Figure 4.8-1) are approximately 60 Vdc. These voltages are not adjustable.
- 6. Press **ENTER** to exit the routine.
- 7. Reconnect the LMNE CIS, GR RBC and GB WBC coaxes (Figure 4.8-1).

**ATTENTION:** When replacing the left side panel with the instrument powered on, avoid accidently turning the instrument off by carefully positioning the opening for the power on/off rocker switch over the switch as you position the panel on the instrument frame.

8. Replace the left side panel and install the four hex screws that secure it to the instrument frame.

#### 4.9 **RBC/PLT GAIN ADJUSTMENT**

### Purpose

Use this procedure to adjust the RBC and Plt gains. A special cycle automatically makes a 1:220 dilution of the RBC/PLT latex (10 µL of latex to 2.2 mL of diluent). A special count program carries out the calculations of the mean volume in the predefined zones and displays them every 700 ms. The duration of the measurement cycle is 21 seconds.

### **Tools/Supplies Needed**

- □ Allen wrench, 3.0 mm
- □ RBC/PLT latex, PN LAD002AS
- □ Plastic potentiometer adjustment tool, PN 5415364

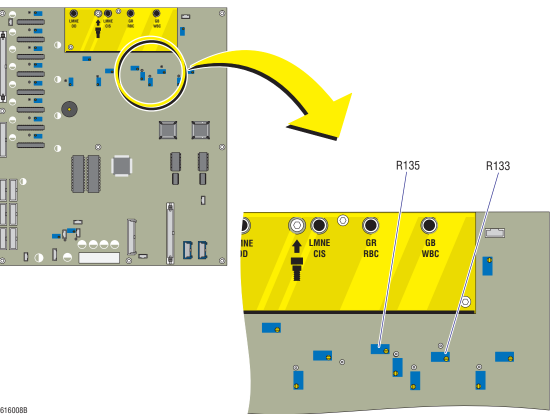
### Procedure

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### Preparation

- 1. Open the left side panel to access the Main card. For details, see Heading 4.2.
- 2. Locate potentiometers R133 for the RBC adjustment and R135 for the PLT adjustment in the upper right quadrant of the Main card (Figure 4.9-1).

### Figure 4.9-1 Main Card RBC/PLT Gain Adjustments



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- 3. From the Service menu, select 2. MEASUREMENT >> 3. RBC / PLT GAIN.
- 4. When the *RBC/PLT GAIN* prompt appears, press **ENTER** to continue.

**IMPORTANT** Risk of misleading results. The RBC/PLT latex particles tend to clump as they settle out of solution. Clumped latex particles will affect adjustment results. Mix RBC/PLT latex vigorously before use. A vortex may be used. Remix the latex thoroughly before each sampling.

- 5. Mix the RBC/PLT latex vigorously. Use a vortex, if available.
- 6. When the *PLEASE SAMPLE LATEX* prompt appears, present the vial of well-mixed latex particles to the probe and press the aspirate switch to initiate aspiration and the routine.

### **Adjustments**

- 1. While monitoring the RBC bar graph on the RBC/PLT Gain screen, adjust potentiometer R133 to read 78 (target value for RBC).
- 2. While monitoring the PLT bar graph on the RBC/PLT Gain screen, adjust potentiometer R135 to read 112 (target value for PLT).

#### **WBC/BASO GAIN ADJUSTMENT** 4.10

### Purpose

Use this procedure to adjust the WBC and BASO gains. A special cycle automatically makes a 1:733 dilution of the RBC/PLT latex (3 µL of latex to 2.2 mL of diluent). A special count program carries out the calculations of the mean volume in the predefined zones and displays them every 700 ms. The duration of the measurement cycle is 21 seconds.

### **Tools/Supplies Needed**

- □ Allen wrench, 3 mm
- □ RBC/PLT latex, PN LAD002AS
- □ Plastic potentiometer adjustment tool, PN 5415364

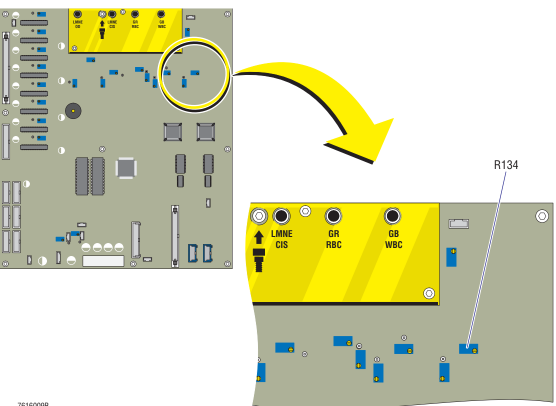
### Procedure

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### Preparation

- 1. Open the left side panel to access the Main card. For details, see Heading 4.2.
- 2. Locate potentiometer R134 in the upper right quadrant of the Main card (Figure 4.10-1). This is the adjustment potentiometer.

### Figure 4.10-1 Main Card WBC/BASO Gain Adjustment



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- 3. From the Service menu, select **2. MEASUREMENT >> 4. WBC / BASO GAIN.**
- 4. When the WBC/BASO GAIN prompt appears, Press ENTER to continue.

**IMPORTANT** Risk of misleading results. The RBC/PLT latex particles tend to clump as they settle out of solution. Clumped latex particles will affect adjustment results. Mix RBC/PLT latex vigorously before use. A vortex may be used. Remix the latex thoroughly before each sampling.

- 5. Mix the RBC/PLT latex vigorously. Use a vortex, if available.
- 6. When the *PLEASE SAMPLE LATEX* prompt appears, present the vial of well-mixed latex particles to the probe and press the aspirate switch to initiate aspiration and the routine.

### Adjustment

While monitoring the WBC/BASO bar graph on the WBC/BASO Gain screen, adjust potentiometer R134 to read 102 (target value).

### 4.11 DRAIN SENSOR ADJUSTMENT

### Purpose

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Use this procedure to adjust the drain sensor. As preparation for this adjustment, this routine automatically empties and fills the drain cell.

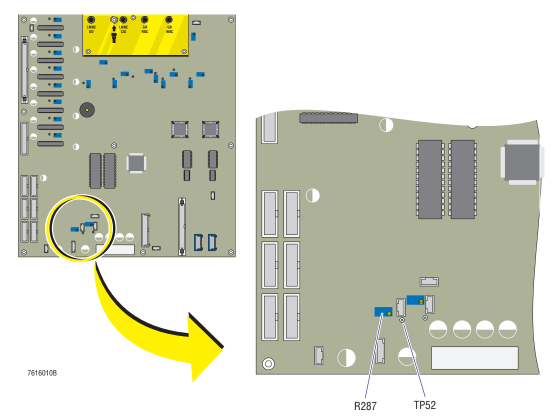
### **Tools/Supplies Needed**

- □ Allen wrench, 3 mm
- □ Plastic potentiometer adjustment tool, PN 5415364
- Digital voltmeter (DVM)

### Preparation

- 1. Remove the four hex screws securing the left side panel to the instrument frame. Set the door aside.
- 2. Locate potentiometer R287 and test point TP52 in the lower left quadrant of the Main card (Figure 4.11-1).

### Figure 4.11-1 Main Card Drain Sensor Adjustment



### Adjustment

- 1. From the Service menu, select **5**. **SENSOR CHECK → 1**. **DRAINING**. The Draining prompt appears on the screen.
- 2. Press **ENTER** to continue. The *CYCLE IN PROGRESS* bar graph appears and advances to approximately 48% while draining the sensor.
- 3. When the ADJUST SENSOR VOLTAGE prompt appears,
  - a. Use a DVM to verify the voltage at TP52 is  $4.5 \pm 0.3$ .
  - b. Adjust the voltage at potentiometer R287 as needed.
  - c. Press **ENTER** to continue. The *CYCLE IN PROGRESS* bar graph advances to approximately 53% as the sensor is filled with diluent.
- 4. When the CHECK SENSOR VOLTAGE prompt appears,
  - a. Make sure the voltage at TP52 is <1.0 Vdc.
  - b. Press **ENTER** to continue. The *CYCLE IN PROGRESS* bar graph advances to 100% as the routine is completed.

### Wrap Up

- 1. Press **ESC** as many times as necessary to return to the Main Menu.
- 2. Replace the left side panel and install the four hex screws that secure it to the instrument frame.

### 4.12 TRANSFER SENSOR ADJUSTMENT

### Purpose

Use this procedure to adjust the transfer sensor. As preparation for this adjustment, this routine automatically empties then fills the drain sensor. This sensor controls the transfer of the diluted sample from the Diff bath to the flow cell.

### **Tools/Supplies Needed**

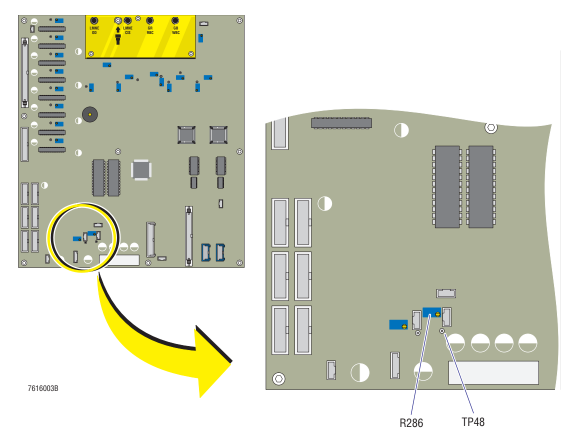
- □ Allen wrench, 3 mm
- □ Plastic potentiometer adjustment tool, PN 5415364
- Digital voltmeter (DVM)

### Preparation

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- 1. Remove the four hex screws securing the left side panel to the instrument frame. Set the door aside.
- 2. Locate potentiometer R286 and test point TP48 in the lower left quadrant of the Main card (Figure 4.12-1).

### Figure 4.12-1 Main Card Transfer Sensor Adjustment



### Adjustment

- 1. From the Service menu, select **5**. **SENSOR CHECK → 2**. **DIFF TRANSFER.** The Diff Transfer prompt appears on the screen.
- 2. Press **ENTER** to continue. The *CYCLE IN PROGRESS* bar graph appears and advances to approximately 48% while draining the sensor.
- 3. When the ADJUST SENSOR VOLTAGE prompt appears,
  - a. Use a DVM to verify the voltage at TP48 is  $4.5 \pm 0.3$ .
  - b. Adjust the voltage at potentiometer R286 as needed.
  - c. Press **ENTER** to continue. The *CYCLE IN PROGRESS* bar graph advances to approximately 53% as the sensor is filled with diluent.
- 4. When the CHECK SENSOR VOLTAGE prompt appears,
  - a. Make sure the voltage at TP48 is <1.0 Vdc.
  - b. Press **ENTER** to continue. The *CYCLE IN PROGRESS* bar graph advances to 100% as the routine is completed.

### Wrap Up

- 1. Press **ESC** as many times as necessary to return to the Main Menu.
- 2. Replace the left side panel and install the four hex screws that secure it to the instrument frame.

### 4.13 MOTOR CURRENT ADJUSTMENT

### Purpose

Use this procedure to check or adjust the voltages for the motors that supply power for the various syringe assemblies and the sampling carriage on the Traverse module.

### **Tools/Supplies Needed**

- □ Allen wrench, 3 mm
- □ Plastic potentiometer adjustment tool, PN 5415364
- Digital voltmeter (DVM)

### Procedure

- 1. Open the left side door to access the Main card.
- 2. Check and adjust, when necessary, the voltages in Table 4.13-1 using the designated potentiometers and test points located in the upper left quadrant of the Main card (Figure 4.13-1).

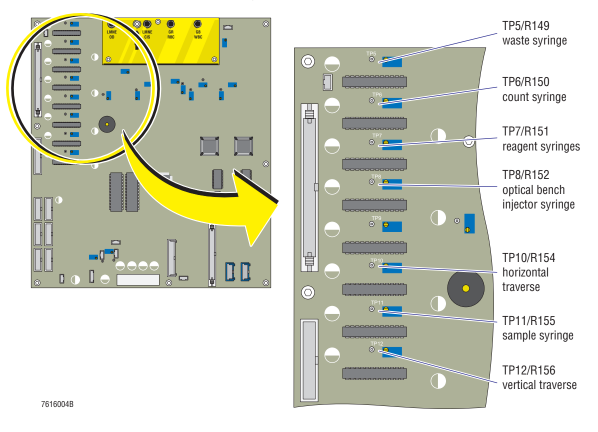
### Table 4.13-1 Motor Voltage Limits

| Motors                         | Test Point | Voltage       | Potentiometer |
|--------------------------------|------------|---------------|---------------|
| Drain syringe                  | TP5        | 4 V ±0.05 V   | R149          |
| Count syringe                  | TP6        | 4 V ±0.05 V   | R150          |
| Reagent syringes               | TP7        | 4 V ±0.05 V   | R151          |
| Optical bench injector syringe | TP8        | 3 V ±0.05 V   | R152          |
| Horizontal traverse            | TP10       | 3 V ±0.05 V   | R154          |
| Sample syringe                 | TP11       | 2 V ± 0.05 V  | R155          |
| Vertical traverse              | TP12       | 4.5 V ±0.05 V | R156          |

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### Figure 4.13-1 Main Card Motor Current Adjustments

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### 4.14 THRESHOLD ADJUSTMENTS

### Purpose

Use this procedure to check or adjust the threshold voltages as needed.

### **Tools/Supplies Needed**

- □ Allen wrench, 3 mm
- □ Plastic potentiometer adjustment tool, PN 5415364
- Digital voltmeter

### Procedure

- 1. Open the left side door to access the Main card.
- 2. Check and adjust, when necessary, the voltages in Table 4.14-1 using the designated potentiometers and test points in the upper area of the Main card (Figure 4.14-1).

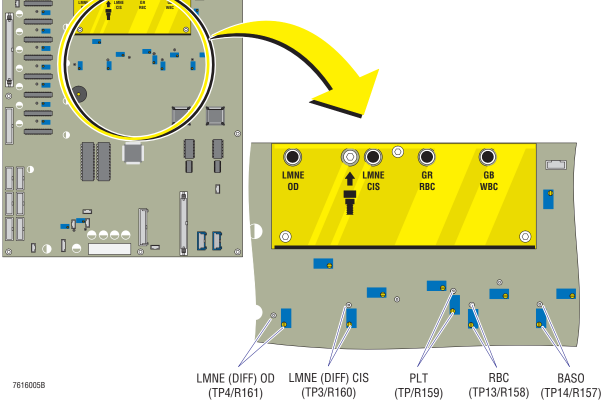
### Table 4.14-1 Threshold Voltage Limits

| Threshold           | Test Point | Voltage   | Potentiometer |
|---------------------|------------|-----------|---------------|
| BASO                | TP14       | 300 mV ±5 | R157          |
| RBC                 | TP13       | 300 mV ±5 | R158          |
| PLT                 | TP2        | 300 mV ±5 | R159          |
| LMNE CIS (for DIFF) | TP3        | 650 mV ±5 | R160          |
| LMNE OD (for DIFF)  | TP4        | 350 mV ±5 | R161          |

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### 4.15 REAGENT TEMPERATURE CHECK AND ADJUSTMENT

### Purpose

Use this procedure to check proper operation of the heating coil and associated system components. This check and adjustment must be made under stabilized thermal conditions.

### **Tools/Supplies Needed**

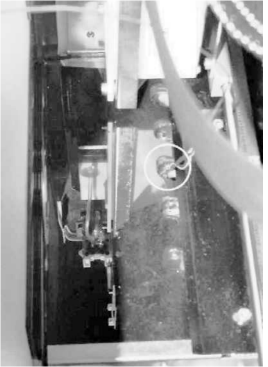
- □ Thermometer probe
- Digital voltmeter

### **Reagent Temperature Check**

- 1. Open the right side door.
- 2. Place the thermometer probe into the upper portion of the fluid in the DIFF bath. Avoid touching the sides and bath electrodes (Figure 4.15-1).

If using a wire bead, bend and insert the bead through the top of the unused fitting of the bath.

### Figure 4.15-1 Thermometer Probe inside the DIFF Bath



3. Insert the thermometer probe leads in the DVM.

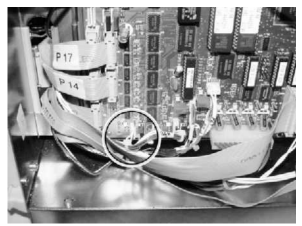
**IMPORTANT** The reagent temperature check must be done under stabilized thermal conditions. Make sure the right side door is closed before starting this check. After completing a startup, wait at least 5 minutes before performing this temperature check.

- 4. Carefully close the right side door to ensure the thermometer probe remains submerged in the reagent.
- 5. Open the left side door.
- 6. In the lower left quadrant of the Main card, locate the LED near connector J31 (Figure 4.15-2) and verify the LED is flashing (rapidly).

Note: This LED indicates heating status:

- When the LED is off, the heating coil is off. When the LED is on, the heating coil is on.
- A slow blinking LED indicates the heating coil is in the process of increasing or decreasing the temperature.
- When the LED is flashing rapidly, the heating coil is maintaining the target temperature.

Figure 4.15-2 Main Card Heating Status LED Location



- 7. From the Service menu, select **3. HEATING SYSTEMS >> 1. HEATING COIL >> 2. REFERENCE**.
- 8. At the Reference screen,
  - a. Set the RUN CYCLE NUMBER to **5**.

**IMPORTANT** Make sure the thermometer probe is submerged in the liquid before starting this routine.

b. Select *RUN* then press **ENTER**.

- 9. Each time the instrument beeps, compare the *REFERENCE TEMPERATURE* with the *RUNNING TEMPERATURE LIMITS* displayed on the screen. Monitor the temperature reading through at least 10 beeps.
  - If the temperature remains within the limits all 10 times, the heating system is working properly. Remove the temperature probe and close the right and left side doors.
  - If the temperature falls outside the displayed limits, perform the Reagent Temperature Adjustment procedure.

### **Reagent Temperature Adjustment**

ATTENTION: Always perform the Reagent Temperature Check before making this adjustment.

### Preparation

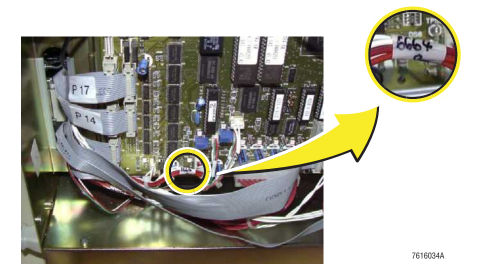
1. Perform a Reagent Temperature Check if you have not already so.

**IMPORTANT** The reagent temperature adjustment must be done under stabilized thermal conditions. Make sure the right side door is closed before starting this adjustment. After completing a startup, wait at least 5 minutes before performing this temperature adjustment.

- 2. From the Service Menu, select **3**. **HEATING SYSTEMS** → **1**. **HEATING COIL** → **1**.**ADJUSTMENT**. The prompt *TEMP*. *ADJ VALUE XXXX* appears.
- 3. In the lower left quadrant of the Main card, locate connector J31 and find the temperture value written on the cable (Figure 4.15-3).

Note: Make sure you read the correct value, the sticker may be oriented up or down.

### Figure 4.15-3 Location of the Label Containing the Temperature Value for the Heater Assembly



- 4. Verify the temperature value on the screen agrees with the temperature value on the cable.
  - If the temperature values agree, press **ESC** to return to Heating Coil menu then proceed to the Adjustment procedure that follows.
  - If the temperature values do not agree, change the screen value to match the value written on the cable, press **ESC** to return to the Heating Coil menu, then continue on to the Adjustment procedure that follows.

### Adjustment

- 1. From the Heating Coil menu, select **2. REFERENCE**.
- 2. At the Reference screen,
  - a. Set the *RUN CYCLE NUMBER* to **5**.

**IMPORTANT** Make sure the thermometer probe is submerged in the liquid before starting this routine.

- b. Select *RUN* then press **ENTER**.
- 3. Each time the instrument beeps, compare the *REFERENCE TEMPERATURE* with the *RUNNING TEMPERATURE LIMITS* displayed on the screen. Monitor the temperature reading through at least 10 beeps.
  - If the temperature remains within the limits all 10 times, the heating system is working properly. Remove the temperature probe and close the left side door.
  - If the temperature falls outside the displayed limits, go to step 4.
- 4. Adjust the *TEMPERATURE REFERENCE* value as needed.

Note: When the temperature reference is increased, the reagent preheating temperature increases; when decreased, the reagent preheating temperature decreases. As the requested temperature change occurs, the heating status LED is blinking slowly.

- 5. In the lower left quadrant of the Main card, monitor the heating status LED (Figure 4.15-2).
- 6. When the heating status LED is flashing rapidly (indicating the target temperature is achieved), repeat steps 2 through 5 until you obtain an acceptable temperature. If it is not possible to properly adjust the temperature, change the heating coil using the instructions under Heading 4.19, HEATER ASSEMBLY REPLACEMENT.

## 4.16 BATH ENCLOSURE TEMPERATURE CHECK AND ADJUSTMENT

### Purpose

Use this procedure to check proper operation of the bath enclosure fan and associated system components. These checks and adjustments must be made under stabilized thermal conditions.

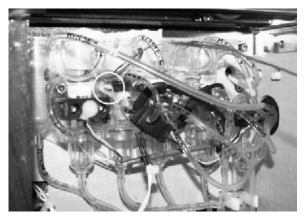
### **Tools/Supplies Needed**

- □ Thermometer probe
- Digital voltmeter

### **Bath Enclosure Temperature Check**

- 1. Open the right side door.
- 2. Place the thermometer probe close to the temperature sensor (Figure 4.16-1).

### Figure 4.16-1 Temperature Sensor Location - View with the Right Side Door Open



3. Insert the thermometer probe leads in the DVM.

**IMPORTANT** The bath enclosure temperature check must be done under stabilized thermal conditions. Make sure the right side door is closed before starting this check. After completing a startup, wait at least 5 minutes before performing this temperature check.

- 4. Carefully close the right side door to ensure the thermometer probe remains close to the temperature sensor.
- 5. From the Service menu, select **3. HEATING SYSTEMS >> 2. BATH ENCLOSURE >> 2. REFERENCE**.
- 6. At the Reference screen,
  - a. Set the RUN CYCLE NUMBER to **5**.

**IMPORTANT** Make sure the thermometer probe is close to the temperature sensor before starting this routine.

b. Select *RUN* then press **ENTER**.

- 7. Each time the instrument beeps, compare the *REFERENCE TEMPERATURE* with the *RUNNING TEMPERATURE LIMITS* displayed on the screen. Monitor the temperature reading through at least 10 beeps.
  - If the temperature remains within the limits all 10 times, the heating system is working properly. Remove the temperature probe.
  - If the temperature falls outside the displayed limits, perform the Bath Enclosure Temperature Adjustment procedure.

### **Bath Enclosure Temperature Adjustment**

**ATTENTION:** Always perform the Bath Enclosure Temperature Check before making this adjustment.

### Preparation

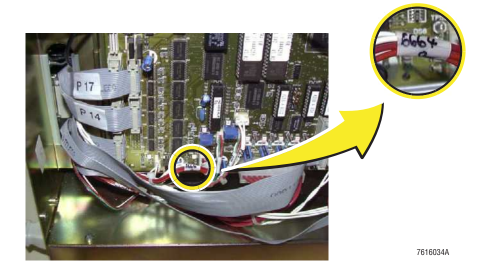
1. Perform a Bath Enclosure Temperature Check if you have not already so.

**IMPORTANT** The bath enclosure temperature adjustment must be done under stabilized thermal conditions. Make sure the right side door is closed before starting this adjustment. After completing a startup, wait at least 5 minutes before performing this temperature adjustment.

- 2. From the Service menu, select **3**. **HEATING SYSTEMS** → **2**. **BATH ENCLOSURE** → **1**.**ADJUSTMENT**. The prompt *TEMP*. *ADJ VALUE XXXX* appears.
- 3. In the lower left quadrant of the Main card, locate connector J33 and find the temperature value written on the cable (Figure 4.16-2).

Note: Make sure you read the correct value, the sticker may be oriented up or down.

### Figure 4.16-2 Location of the Label Containing the Temperature Value for the Temperature Sensor



- 4. Verify the temperature value on the screen agrees with the temperature value on the cable.
  - If the temperature values agree, press **ESC** to return to Bath Enclosure menu then proceed to the Adjustment procedure that follows.
  - If the temperature values do not agree, change the screen value to match the value written on the cable, press **ESC** to return to the Bath Enclosure menu, then continue on to the Adjustment procedure that follows.

### Adjustment

- 1. From the Bath Enclosure menu, select **2. REFERENCE**.
- 2. At the Reference screen,
  - a. Set the RUN CYCLE NUMBER to **5**.

**IMPORTANT** Make sure the thermometer probe is close to the temperature sensor before starting this routine.

- b. Select *RUN* then press **ENTER**.
- 3. Each time the instrument beeps, compare the *REFERENCE TEMPERATURE* with the *RUNNING TEMPERATURE LIMITS* displayed on the screen. Monitor the temperature reading through at least 10 beeps.
  - If the temperature remains within the limits all 10 times, the bath enclosure temperature monitoring system is working properly. Remove the temperature probe and close the right side door.
  - If the temperature falls outside the displayed limits, go to step 4.
- 4. Adjust the *REFERENCE TEMPERATURE* value as needed.
  - This adjustment controls the operation of a bath enclosure fan and heater.
  - If the reference temperature is increased, the fan and a heater mechanism inside the fan is turned on until the ambient bath enclosure temperature reaches the desired temperature.
  - If the reference temperature is decreased, the fan is turned on to pull atmospheric air into the bath enclosure until the ambient temperature inside the enclosure reaches the desired temperature.
  - If it is not possible to properly adjust the temperature, check fan operation.

**SERVICE AND REPAIR PROCEDURES** BATH ENCLOSURE TEMPERATURE CHECK AND ADJUSTMENT

### 4.17 VACUUM CHECKS AND ADJUSTMENTS

### Purpose

Use this procedure to check waste or count syringe vacuum and to adjust the count syringe vacuum, as needed. Since altitude affects vacuum, these vacuum checks should be performed anytime an instrument is being installed in a location that is either above or below sea level.

### **Tools/Supplies Needed**

□ External digital pressure/vacuum gauge; hereafter, referred to as a vacuum meter.

### Waste Syringe Vacuum Check

- 1. Open the right side door.
- 2. Remove the tubing attached to the side of the waste syringe and replace it with a tubing attached to the vacuum meter, as shown in Figure 4.17-1.

### Figure 4.17-1 Attach the Vacuum Meter to the Waste Syringe



**IMPORTANT** Risk of misleading results. The syringe creates a finite vacuum. It does not have continuous flow and evacuation like a pump. If the tubing attached to the vacuum meter is too large, the volume of the tubing adds to the volume in the syringe and may produce falsely low readings.

- 3. From the Service menu, select 6. VACUUM CHECK → 2. DRAINING.
- 4. When the *CHECK VACUUM* prompt appears on the screen, check the vacuum reading on the vacuum meter. The vacuum must be approximately 260 mb (7.7-inches Hg) and must be stable.

Note: No vacuum adjustment is available for the waste syringe.

- 5. Disconnect the vacuum meter and reattach the tubing on the waste syringe.
- 6. Close the right side door.

### **Count Syringe Vacuum Check and Adjustment**

### **Count Syringe Vacuum Check**

1. Remove the left side panel. For details, see Heading 4.2.

**CAUTION** A flat cable is connected to the rear of the Main card. Be careful when you open the door (where the Main card is attached) that you do not disconnect or damage this cable.

- 2. In the left side compartment,
  - a. Turn the two captive knobs counterclockwise to release the Main card door. These knobs are located to the right of the Main card.
  - b. Open the Main card door and carefully anchor the door behind the white plastic catch to keep it open. The optical bench is exposed.

**IMPORTANT** Risk of misleading results. The syringe creates a finite vacuum. It does not have continuous flow and evacuation like a pump. If the tubing attached to the vacuum meter is too large, the volume of the tubing adds to the volume in the syringe and may produce falsely low readings.

3. At the count syringe, disconnect the side tubing nearest the bottom of the syringe and replace it with a tubing attached to the vacuum meter, as shown in Figure 4.17-2.

#### Figure 4.17-2 Attach the Vacuum Meter to the Count Syringe



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- 4. From the Service menu, select 6. VACUUM CHECK >> 1. COUNTING.
- 5. When *VACUUM XXX* prompt appears on the screen, the down arrow key select the *RUN* prompt.
- 6. When the *CHECK VACUUM* (*XXX MB*) prompt appears, check the vacuum reading on the vacuum meter. The vacuum must be approximately 220 mb (6.5-inches Hg) and stable.
  - If the vacuum level is correct and stable, go to step 7.
  - If the vacuum level is too low or unstable, press **ESC** to erase the *CHECK VACUUM* (*XXX MB*) prompt then go to the Count Syringe Vacuum Adjustment heading to make the appropriate adjustment.
- 7. Disconnect the vacuum meter and reattach the tubing on the count syringe.
- 8. Close the Main card door. Turn the two captive knobs clockwise to secure the door.

**ATTENTION:** When replacing the left side panel with the instrument powered on, avoid accidently turning the instrument off by carefully positioning the opening for the power on/off rocker switch over the switch as you position the panel on the instrument frame.

- 9. Replace the left side panel and install the four hex screws that secure it to the instrument frame.
- 10. Press **ESC** as many times as necessary to return to the Main Menu.
- 11. Continue operation.

### **Count Syringe Vacuum Adjustment**

- 1. At the Counting screen, use the up arrow to move the cursor to the *VACUUM XXX* prompt then change the step value for vacuum.
  - If the vacuum is too low, increase the step value.
  - If the vacuum is too high, decrease the step value.
- 2. Select the RUN prompt.
- 3. When the prompt *CHECK VACUUM* (*XXX MB*) appears, check the vacuum reading on the vacuum meter. The vacuum must be approximately 220 mb (6.5-inches Hg) and stable.
  - If the vacuum level is correct and stable, go to step 4.
  - If the vacuum level is still unacceptable, press **ESC** and repeat steps 1 through 3 until the vacuum is stable at approximately 220 mb (6.5-inches Hg).
- 4. Disconnect the vacuum meter and reattach the tubing on the count syringe.
- 5. Close the Main card door. Turn the two captive knobs clockwise to secure the door.

**ATTENTION:** When replacing the left side panel with the instrument powered on, avoid accidently turning the instrument off by carefully positioning the opening for the power on/off rocker switch over the switch as you position the panel on the instrument frame.

- 6. Replace the left side panel and install the four hex screws that secure it to the instrument frame.
- 7. Press **ESC** as many times as necessary to return to the Main Menu.
- 8. Continue operation.

# **SERVICE AND REPAIR PROCEDURES** VACUUM CHECKS AND ADJUSTMENTS

### 4.18 MIX BUBBLE ADJUSTMENT

### Purpose

Mixing bubbles are factory adjusted and normally do not require further adjustments; however, if an adjustment is necessary follow this procedure.

### **Tools/Supplies Required**

None

### Procedure

- 1. From the Service menu, select 4. MIXING.
- 2. At the Mixing screen, enter a new step value to increase or decrease the mixing, as needed. Use the values in Table 4.18-1 as a guide.
  - To increase bubbling, increase the number of steps.
  - To decrease bubbling, decrease the number of steps.

### Table 4.18-1 Mixing Bubble Limits

| Mixing Bath    | Low Limit | Normal | High Limit |
|----------------|-----------|--------|------------|
| FIRST DILUTION | 100       | 300    | 400        |
| DIFF           | 100       | 300    | 400        |
| WBC/BASO       | 100       | 300    | 400        |
| HGB LYSE       | 300       | 400    | 500        |

3. Verify the final step values are within limits (Table 4.18-1).

### **SERVICE AND REPAIR PROCEDURES** *MIX BUBBLE ADJUSTMENT*

## HEATER ASSEMBLY REPLACEMENT

### **Purpose**

4.19

Use this procedure to replace the heater assembly.

### **Tools/Supplies Needed**

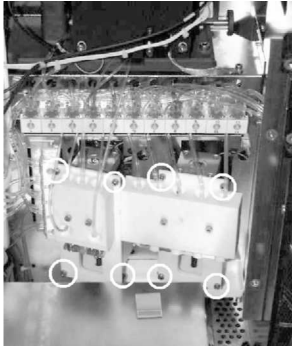
- □ Allen wrench, 3 mm
- □ Flat-blade screwdriver
- □ Torx keys
- □ Heater reagent coil assembly, PN XDA625AS

### Removal

- 1. To access the heater assembly:
  - a. Remove the front panel. Follow cover removal under Heading 4.2.
  - b. Pull the reagent syringes and the 5 diff syringe to free space behind (no need to disconnect tubes).

**ATTENTION:** The syringe assemblies uses captive hex screws mounted inside rubber shock mounts. It is recommended that you use only four turns to either loosen or tighten these screws. Turning the hex screw too many counterclockwise rotations may separate the screw from the rubber shock mount. If all Service Representatives consistently use four turns to remove or install these hex screws, it is unlikely that a rubber shock mount will separate from its hex screw and fall inside the instrument.

c. Locate the six hex screws (CHC M4x16) shown in Figure 4.19-1. These are captive screws anchored inside rubber shock mounts.



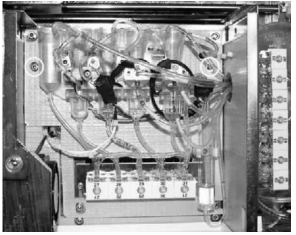
### Figure 4.19-1 Captive Hex Screw Locations - Reagent Syringes and 5diff Syringe

d. Loosen each hex screw with four counterclockwise rotations.

**Note:** If a screw is not released in four rotations, the screw was overtightened the last time the syringe assembly was serviced. From this point, make a single rotation then check to see if the mount is free. Repeat this sequence of making a single rotation and checking until the mount is free.

- 2. Open the right side door.
- 3. Disconnect the following tubings:
  - At the WBC/BASO bath, disconnect the tubing attached to port 2.
  - At the RBC bath, disconnect the tubing attached to port 1.
  - At the DIFF bath, disconnect the tubing attached to ports 2 and 3.
  - At the DIL1/HGB bath, disconnect the tubing attached to port 1.
- 4. Unscrew the two CHC M3x6 screws shown in Figure 4.19-2.

### Figure 4.19-2 CHC M3 x 6 Screw Locations



5. The heater assembly is free. Lay the assembly on the instrument frame (Figure 4.19-3).

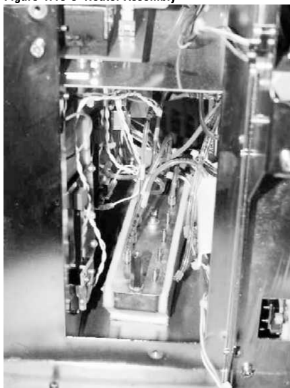


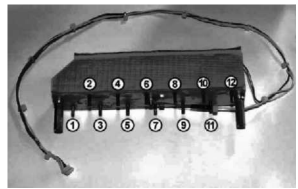
Figure 4.19-3 Heater Assembly

- 6. Move the heater assembly as close as possible to the opening in the instrument's frame.
- 7. Using Figure 4.19-4 as a guide, disconnect the tubings attached to ports 1, 3, 5, 7, and 10 on the heater assembly.

Note: The ports are associated with the following components:

- The tubing disconnected from port 1 is from valve 22, port 2.
- The tubing disconnected from port 3 is from valve 8, port 2.
- The tubing disconnected from port 5 is from T4.
- The tubing disconnected from port 7 is from valve 22, port 1.
- The tubing disconnected from port 10 is from valve 11, port 2.

### Figure 4.19-4 Heater Assembly - Tubing Port Locations



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8. Locate connector P33 in the lower left quadrant of the Main card (Figure 4.19-5).

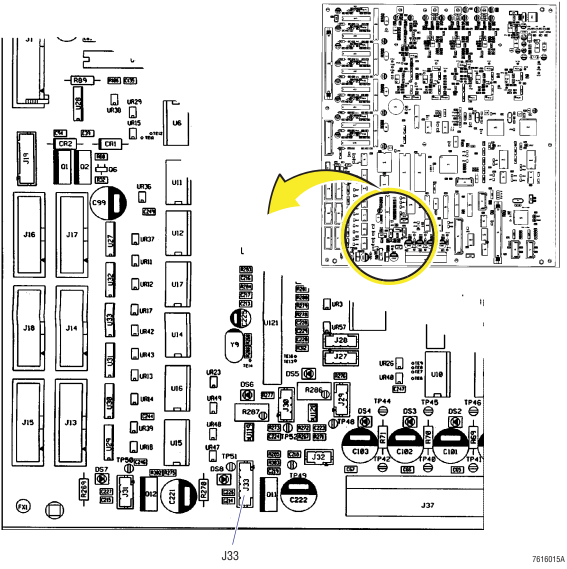


Figure 4.19-5 Main Card Heater Assembly Replacement

9. Remove the heater assembly cable attached to J33.

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10. Pull the disconnected cable (and tubings) through the instrument and remove the heater assembly (Figure 4.19-6).

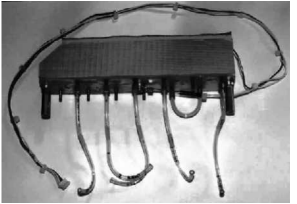


Figure 4.19-6 Heater Assembly After Removal

11. Place the heater assembly on a stack of absorbent paper towels to drain it.

#### Installation

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- 1. Place the new heater assembly next to the old heater assembly. Make sure they are oriented in the same direction (cables on the same side, ports facing the same direction).
- 2. One by one, disconnect a tubing from the old heater and attach it in the same location on the new heater. Use Figure 4.19-7 as a guide if needed.

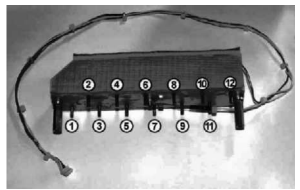


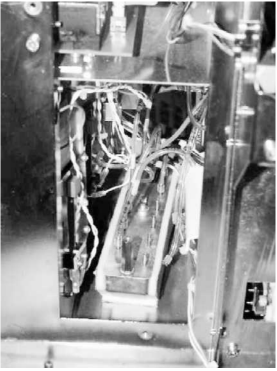
Figure 4.19-7 Heater Assembly - Tubing Port Locations

**Note:** The tubings being removed from the ports on the old heater and attached to the ports on new heater are associated with the following components:

- The tubing attached to port 2 will later be attached to the diluent bath, DIL1/HGB.
- The tubing attached to port 4 will later be attached to the DIFF bath, DIFF 3.
- The tubing attached to port 6 will later be attached to the DIFF bath, DIFF 2.
- The tubing attached to port 8 will later be attached to the RBC bath, RBC 1.
- The tubing attached to port 9 loops to port 11 on the heater assembly.
- The tubing attached to port 12 will later be attached to the WBC/BASO bath, WBC/BASO 2.

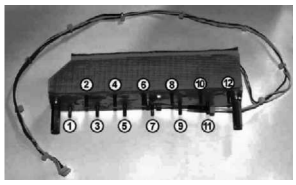
- 3. Discard the old heater assembly.
- 4. Position the new heater assembly (with the old tubing attached) inside the instrument frame (Figure 4.19-8).

#### Figure 4.19-8 Heater Assembly - Orientation Inside the Instrument



- 5. Using Figure 4.19-9 as a guide, one by one, locate each tubing removed from the old heater assembly and attach the tubing to its designated port on the new heater assembly:
  - a. Locate the tubing attached to valve 11, port 2. Attach this tubing to port 10 of the heater assembly.
  - b. Locate the tubing attached to valve 22, port 1. Attach this tubing to port 7 of the heater assembly.
  - c. Locate the tubing attached to T4. Attach this tubing to port 5 of the heater assembly.
  - d. Locate the tubing attached to valve 8, port 2. Attach this tubing to port 3 of the heater assembly.

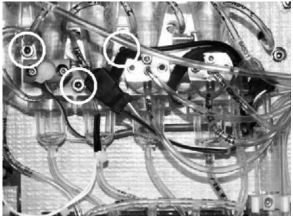
#### Figure 4.19-9 Heater Assembly - Port Locations



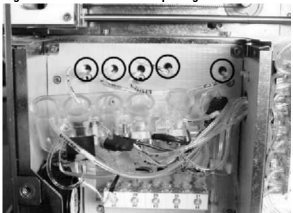
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- e. Locate the tubing attached to valve 22, port 2. Attach this tubing to port 1 of the heater assembly.
- 6. Remove the three nuts shown in Figure 4.19-10. Removing these nuts dismantles the bath assembly from the instrument frame so that tubings attached to the new heater assembly can be routed through the panel openings for attachment to the designated bath.

Figure 4.19-10 Baths Assembly Support Panel Nut Locations



7. Lift and position the heater assembly so that each tubing disconnected during the removal procedure can be routed through the support panel opening closest to the bath where the tubing must be attached (Figure 4.19-11).



#### Figure 4.19-11 Location of Openings in the Baths Assembly Support Panel

- 8. From left to right, locate the tubing on the heater assembly and route the tubing through the panel opening closest to the bath:
  - Route the tubing attached to heater assembly port 2 through the first opening for attachment to the DIL1/HGB bath.
  - Route the tubing attached to heater assembly port 4 through the second opening for attachment to the DIFF bath, DIFF 3.
  - Route the tubing attached to heater assembly port 6 through the third opening for attachment to the DIFF bath, DIFF 2.

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- Route the tubing attached to heater assembly port 8 through the fourth opening for attachment to the RBC bath, RBC 1.
- Route the tubing attached to heater assembly port 12 through the fifth opening for attachment to the BASO bath, BASO 2.
- 9. Secure the two heater assembly screws (Figure 4.19-12).

Figure 4.19-12 Heater Assembly Screw Locations



- 10. At the Main card, reconnect the cable attached to the heater assembly to connector J33 (Figure 4.19-5).
- 11. Secure the reagent syringes and the 5diff syringe back inside the instrument:
  - a. Locate the six captive hex screws (Figure 4.19-1). The syringe assemblies should be flush against the instrument frame.

**ATTENTION:** It is recommended that you use only four clockwise turns to tighten these captive hex screws. As each hex screw is tightened, its rubber shock mount expands to secure the syringe assembly to the instrument frame. If all Service Representatives consistently use four turns to install these hex screws, it is unlikely that a rubber shock mount will separate from its hex screw and fall inside the instrument the next time the assembly is removed.

- b. Tighten the six hex screws using four clockwise rotations.
- 12. Replace the front panel. Follow Heading 4.2, OPENING OR REMOVING INSTRUMENT DOORS, PANELS, AND COVERS.
- 13. Do a probe to bath alignment check. Go to Heading 4.5, BATHS ASSEMBLY ALIGNMENT CHECK AND ADJUSTMENT.
- 14. When startup and prime cycles are done, verify there are no leaks.

#### 4.20 POWER SUPPLY REPLACEMENT

#### Purpose

Use this procedure to remove and replace the power supply as needed.

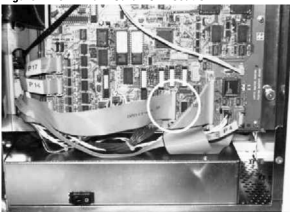
#### **Tools/Supplies Needed**

- □ Allen wrench, 3 mm
- □ Flat-blade screwdriver
- Dever Supply, PN 1xDBN 004 A

#### Removal

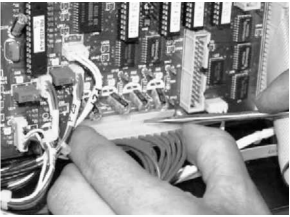
- 1. Turn the instrument off and disconnect the power cord.
- 2. Open the cover.
- 3. Open the left side door.
- 4. At the bottom of the Main card,
  - a. Disconnect the printer RS flat cable attached to connector J2 (Figure 4.20-1).

#### Figure 4.20-1 Main Card - J2 Location



b. With a flat-blade screwdriver, disconnect the power supply cable attached to connector J37 (Figure 4.20-2).

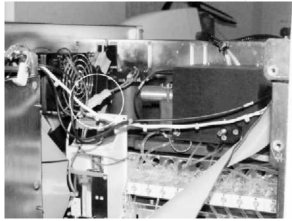
#### Figure 4.20-2 Main Card - J37 Location



**CAUTION** A flat cable is connected to the rear of the Main card. Be careful when you open the door (where the Main card is attached) that you do not disconnect or damage this cable.

- 5. Open the Main card door (block it open).
- 6. Disconnect the optical bench lamp supply cable (Figure 4.20-3).

#### Figure 4.20-3 Optical Module - Lamp Supply Cable Location



7. At the rear panel of the instrument, remove the two CHC M3x6 screws securing the power supply to the rear panel (Figure 4.20-4).

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Figure 4.20-4 Location of Screws Securing the Power Supply to the Rear Panel

- 8. Remove the rear panel and set it aside.
- 9. Remove the two CHC M3x6 front screws securing the power supply to the instrument frame (Figure 4.20-5).

#### Figure 4.20-5 Screws Securing the Power Supply to the Instrument Frame



**CAUTION** Make sure the lamp power supply cable is disconnected before removing the power supply.

10. Remove the power supply.

#### Installation

- 1. Position the new power supply inside the instrument.
- 2. Secure the power supply to the instrument frame using the two CHC M3x6 front screws removed earlier (Figure 4.20-5).

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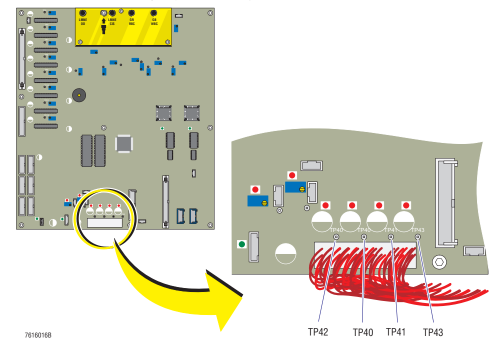
- 3. Make sure the lamp supply cable is routed towards the optical bench then replace the rear panel.
- 4. Secure the power supply to the rear panel using the two CHC M3x6 screws removed earlier (Figure 4.20-4).
- 5. Connect the optical bench lamp supply cable (Figure 4.20-3).
- 6. Close the Main card door.
- 7. At the bottom of the Main card,
  - a. Connect the power supply cable to connector J37 (Figure 4.20-2).
  - b. Connect the printer RS flat cable to connector J2 (Figure 4.20-1).
- 8. Insert the power supply connector into the socket on the rear instrument panel. Make sure the plug end is in the wall socket.
- 9. Turn the instrument on.
- 10. Replace the left side panel.

#### Verification

At the Main card, check following voltages (Figure 4.20-6):

| Test Point | Designation       | Target Voltage |
|------------|-------------------|----------------|
| TP40       | 5V Power supply   | +5 V           |
| TP41       | 12V Power supply  | +12 V          |
| TP42       | 24V Power supply  | +24 V          |
| TP43       | -12V Power supply | -12 V          |

#### Figure 4.20-6 Main Card Heater Assembly Replacement Adjustment



#### 4.21 START SWITCH REPLACEMENT

#### Purpose

Use this procedure to remove then replace the Start switch anytime it needs to be changed or dismantled.

#### **Tools/Supplies Needed**

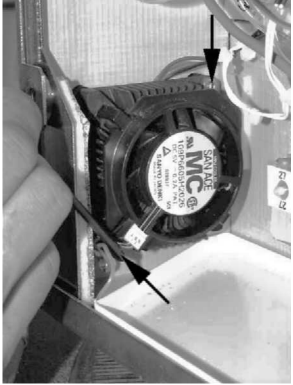
- □ Allen wrenches
- □ Torx keys
- □ Flat-blade screwdriver

#### Removal

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- 1. To access the start switch, remove the front panel. See Heading 4.2, OPENING OR REMOVING INSTRUMENT DOORS, PANELS, AND COVERS, if needed.
- 2. Open the pneumatic access door (right side of the instrument).
- 3. Remove the enclosure fan that contains a thermostat (Figure 4.21-1).
  - a. Using a flat-blade screwdriver, remove the ground from the fan assembly.
  - b. Remove the four screws securing the fan to the front panel.

#### Figure 4.21-1 Fan Removal - Right Side Compartment



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4. Remove the two screws securing the start switch to the instrument's frame (Figure 4.21-2).

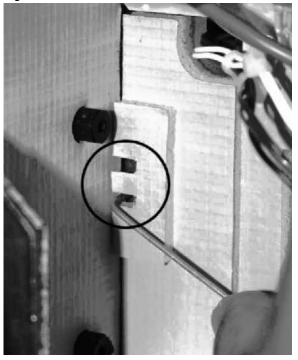
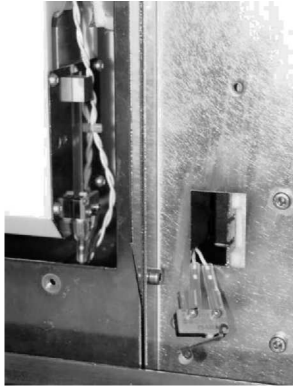


Figure 4.21-2 Start Switch Screw Locations - With Fan Removed

5. Disconnect the start switch (Figure 4.21-3).

Figure 4.21-3 Disconnected Start Switch - Front View with Front Panel Removed



6. Remove the start switch.

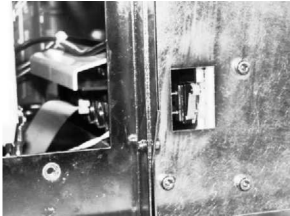
#### Installation

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**IMPORTANT** The Start switch is connected in the normally open position.

- 1. Connect the new start switch to J34 on the Main card.
- 2. Position the start switch (with the wheel at the top) in the two slots (Figure 4.21-4).

#### Figure 4.21-4 Start Switch Orientation



- 3. Install the two screws that secure the switch to the instrument frame.
- 4. With the front panel in position, verify that the start cycle key is operating normally.
  - When the Start switch is installed correctly, you will hear a click when the sample bar is pressed and another click when the sample bar is released.
  - If you do not hear the two clicks, move the switch forward or backward until the location is correct.

**WARNING** Use the washer to connect the ground wiring to the fan.

- 5. Reinstall the fan.
- 6. Reassemble the front panel. See Heading 4.2, OPENING OR REMOVING INSTRUMENT DOORS, PANELS, AND COVERS, if needed.

## **SERVICE AND REPAIR PROCEDURES** *START SWITCH REPLACEMENT*

#### 4.22 OPTICAL BENCH DISASSEMBLY AND REPLACEMENT

#### Purpose

Use this procedure to remove, replace, and control the optical bench assembly.

#### **Tools/Supplies Needed**

□ Allen wrenches, 3.0 mm and 5.0 mm

#### Removal

- 1. Turn the instrument off and unplug the power cord from the instrument or the wall.
- 2. Remove the left side panel and the top cover from the instrument. For details, see Heading 4.2. Set the left side panel and top cover aside.

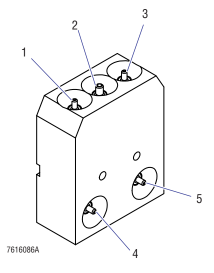
**CAUTION** A flat cable is connected to the rear of the Main card. Be careful when you open the door (where the Main card is attached) that you do not disconnect or damage this cable.

- 3. In the left side compartment,
  - a. Turn the two captive knobs counterclockwise to release the Main card door. These knobs are located to the right of the Main card.
  - b. Open the Main card door and carefully anchor the door behind the white plastic catch to keep it open.

**ATTENTION:** The two tubings disconnected from the 5diff syringe are attached to the flow cell. To ensure proper reconnection, you may want to label each tubing with its port number before disconnecting the tubing from the 5diff syringe assembly.

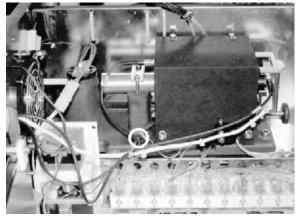
4. At the 5diff syringe, disconnect the tubings attached to ports 1 and 5.

#### Figure 4.22-1 5diff Syringe Port Locations



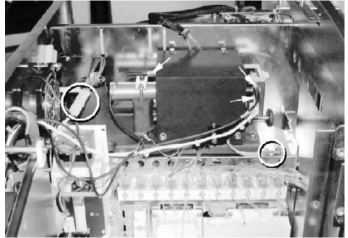
- 5. At the 11-valve assembly,
  - a. Disconnect the tubing attached to solenoid valve 1, port 1.
  - b. Disconnect the tubing attached to solenoid valve 4, port 2.
- 6. Disconnect the tubing attached to the ground fitting (Figure 4.22-2).

#### Figure 4.22-2 Optical Bench - Ground Fitting Location



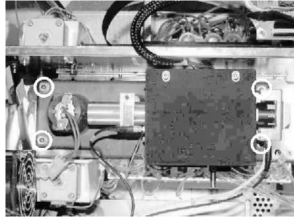
- 7. Disconnect the following components. See Figure 4.22-3 for disconnection locations (from left to right):
  - Lamp supply connector
  - Optical Preamplifier card supply
  - LMNE CIS and LMNE OD coaxes
  - Optical bench grounding wire

#### Figure 4.22-3 Disconnection Sites for Named Components



**ATTENTION:** The optical bench assembly uses captive hex screws mounted inside rubber shock mounts. It is recommended that you use only four turns to either loosen or tighten these screws. Turning the hex screw more than four counterclockwise rotations may separate the screw from the rubber shock mount. If all Service Representatives consistently use four turns to remove or install these hex screws, it is unlikely that a rubber shock mount will separate from its hex screw and fall inside the instrument.

8. Locate the four hex screws circled in Figure 4.22-4. These are captive screws anchored inside rubber shock mounts.



#### Figure 4.22-4 Captive Screw Locations - Optical Bench Assembly

9. Loosen each hex screw with four counterclockwise rotations (Figure 4.22-4). Use a 5.0 mm Allen wrench.

**Note:** If a screw is not released in four rotations, the screw was overtightened the last time the optical bench assembly was serviced. From this point, make a single rotation then check to see if the mount is free. Repeat this sequence of making a single rotation and checking until the mount is free.

10. Gently remove the optical bench from the instrument.

#### Installation

1. Locate the new optical bench assembly and position it inside the instrument.

**Note:** Make sure the rubber shock mounts are positioned behind the panel. The optical bench assembly should be flush against the instrument.

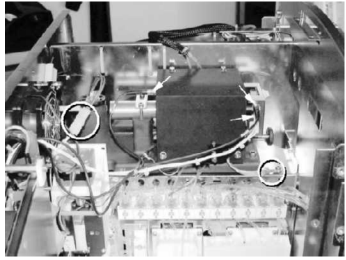
**ATTENTION:** It is recommended that you use only four clockwise turns to tighten the captive hex screws. As each hex screw is tighten, its rubber shock mount expands to secure the optical bench assembly to the instrument. If all Service Representatives consistently use four turns to install these hex screws, it is unlikely that a rubber shock mount will separate from its hex screw and fall inside the instrument the next time the assembly is removed.

Although you must be careful to not overtighten the rubber shock mounts, make sure the optical bench is mounted securely. The assembly should not move if you try and lift it.

2. Tighten the four captive hex screws using four clockwise rotations (Figure 4.22-4). Use a 5.0 mm Allen wrench.

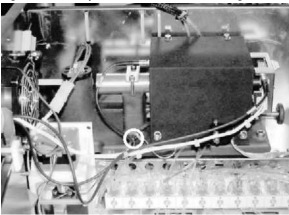
- 3. Reconnect the following components. See Figure 4.22-5 for connection locations (from left to right):
  - Lamp supply connector
  - Optical Preamplifier card supply
  - LMNE CIS and LMNE OD coaxes
  - Optical bench grounding wire

#### Figure 4.22-5 Connection Sites for Named Components



4. Reconnect the tubing attached to the ground fitting (Figure 4.22-6).

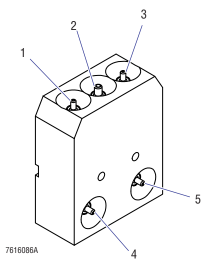
#### Figure 4.22-6 Optical Bench - Ground Fitting Location



- 5. Reconnect the two hydraulic tubings as follows:
  - a. Attach the tubing connected to the isolator chamber to solenoid valve 1, port 1.
  - b. Attach the tubing connected to the flow cell (T-connector) to solenoid valve 4, port 2.

6. At the 5diff syringe, reattach the two flow cell tubings to their proper ports, port 1 and port 5.





7. When the installation is completed, go to Heading 4.4, FLOW CELL CHECKS AND ADJUSTMENTS to ensure proper flow cell operation.

**SERVICE AND REPAIR PROCEDURES** OPTICAL BENCH DISASSEMBLY AND REPLACEMENT

### 4.23 BAR-CODE READER TESTING AND CONFIGURATION

#### Purpose

Use this procedure to test, troubleshoot, and reconfigure the bar-code reader.

#### **Read Test**

Verify the bar-code reader is working properly by successfully reading the test labels. That is, when the bar-code reader is programmed to the default configuration, the reader successfully reads all the labels in Table 4.23-1 and the Codabar label in Table 4.23-2.

If the bar-code reader fails to read the bar-code labels as specified, restore the reader to the default settings, perform the Read Test again, then program each bar-code option per lab requirements.

**IMPORTANT** There is a risk of sample misidentification if the entire bar code is not captured with the bar-code reader, especially with the Interleaved 2-of-5 bar-code format. Position the bar-code reader over the label to capture the entire bar-code identification. Otherwise, part of the identification may not be scanned, resulting in misidentification. Just as an operator must pass the bar-code reader over the bar-code label on the specimen tube to capture the entire bar-coded sample ID, make sure the entire label is read to avoid possible misidentification.





Code 128



Code 39 If this label is read with Check Digit disabled, the last character "\$" is also displayed.



Interleaved 2-of-5 Reads 11 characters with Check Digit or reads 12 characters without Check Digit.



EAN 8 Reads 123456770



EAN13 Reads 12345678901228

#### Table 4.23-2 Test Labels Without Check Digit



Code 39 Label will not read if scanner is programmed to default condition

# A123123A

Codabar

#### **Default Settings**

Do this procedure when the Read Test fails or you want to restore the default settings to the bar-code reader. The default values are shown below.

| Setting                    | Code 128●         | Code 39          | Codabar          | l 2-of-5         | EAN 8             | EAN 13            |
|----------------------------|-------------------|------------------|------------------|------------------|-------------------|-------------------|
| Character Length           | 1 to 16           | 1 to 16          | 3 to 16          | 118              | 7                 | 12                |
| Check Digit (Checksum)     | Always<br>Enabled | Enabled          | Not<br>Available | Enabled          | Always<br>Enabled | Always<br>Enabled |
| Start/Stop Equality Check  | Not<br>Available  | Not<br>Available | Enabled          | Not<br>Available | Not<br>Available  | Not<br>Available  |
| Start/Stop Equality Output | Not<br>Available  | Not<br>Available | Disabled         | Not<br>Available | Not<br>Available  | Not<br>Available  |

• Code 128 provides excellent density, alphanumeric characters, and good security. Recommend using this symbology if using barcodes for the first time, and if compatible with other bar code systems used in your lab.

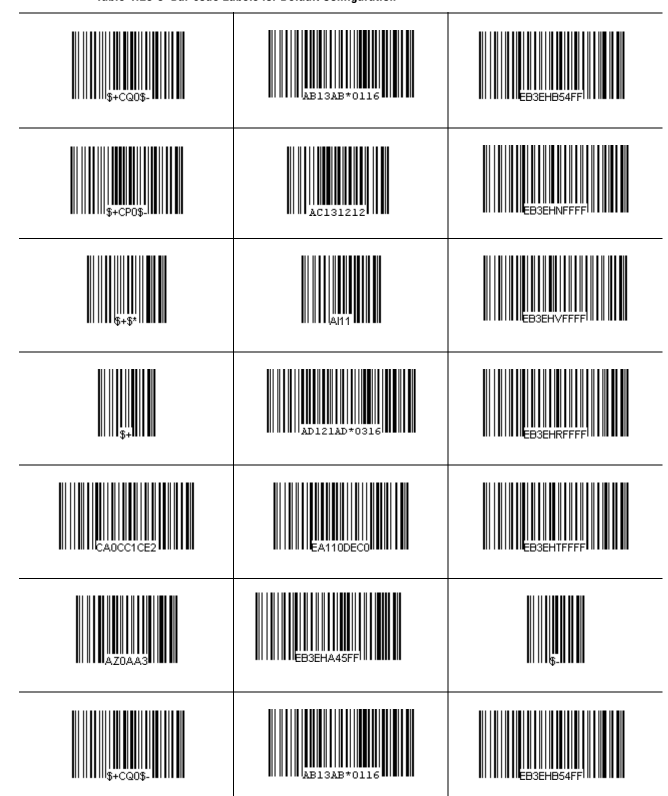
• For increased sample identification integrity, always use Check Digit (Checksum).

• Number of characters for I 2-of-5 can be programmed for other lengths, including variable length. However, the variable length is NOT recommended for I 2-of-5 due to the possibility of capturing a partial read of the bar code label.

- 1. If the instrument is on, turn the instrument off then back on again before starting programming.
- 2. Read in the bar-code labels in Table 4.23-3 from top to bottom and left to right. Bar codes with \$+ and \$- will sound multiple beeps when read. Other codes will only sound a single beep.
- 3. When the last label is read, do the Read Test again.

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#### Table 4.23-3 Bar-code Labels for Default Configuration



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Read in one of the Code 39 labels below to change the Check Digit option.





#### Codabar - Start/Stop Equality Check/Output

If you want to change the Start/Stop Equality Check/Output default setting (*start/stop equality check but no transmissions*), read in one of the labels below.



No Start/Stop Equality Check Nor Transmission



Start/Stop Equality Check But No Transmission



No Start/Stop Equality Check But Transmission



Start/Stop Equality Check And Transmission

#### **Interleaved 2-of-5 Options**

If you want to change the Interleaved 2-of-5 default setting (check digit with 11 digits), read in one of the labels for check digit control and select one of the labels to set the number of digits.

Note: To increase sample identification integrity, use fixed length digits with Check Digit.

When check digit is enabled, the available digits are: 3, 5, 7, 9, 11, 13, 15, and variable (3 to 15).

When check digit is disabled, the available digits are: 4, 6, 8, 10, 12, 14, 16 and variable (4 to 16).

Variable length digits are NOT recommended for Interleaved 2-of-5 bar codes. If the test label fails to read, reset the scanner by turning the instrument off then on and repeating the programming sequence.

| Number of<br>Digits | With Check Digit Control                                  | No Check Digit Control                                    | Fixed Digit Test Labels |
|---------------------|---|---|-------------------------|
|                     | Read this label first, then ONE of the other labels below | Read this label first, then ONE of the other labels below |                         |
|                     | \$+ACO\$-   | \$+&C0\$-   |                         |
| 3* or 4†            | \$+AC130404\$-  | \$+AC110404\$-  | 1236                    |
| 5* or 6†            | \$+AC130606\$-  | \$+AC110606\$-  | 123457                  |
| 7* or 8†            | \$+AC130808\$-  | \$+AC110808\$-  | 12345670                |



\*= With Check Digit Control †= No Check Digit Control

#### 4.24 REAGENT SYRINGES ASSEMBLY REPLACEMENTS

#### Purpose

Use this procedure to remove and replace the reagent syringe pistons, O-rings, or washers.

The reagent syringes assembly consists of five reagent syringes. With the exception of the Hgb Lyse syringe (the syringe with the smallest diameter), each syringe piston has both a washer and an O-ring. The Hgb Lyse syringe has an O-ring but no washer.

When performing this procedure to replace an O-ring or a washer, replace both. When replacing a syringe piston, replace its O-ring and washer as well.

#### **Tools/Supplies Needed**

- □ Allen wrench, 3 mm
- □ Torque screwdriver, 2.5 mm and T10 torx

Note: 400 mN.m (56.8 ozf.in) torque is required in this procedure.

- □ Absorbent paper towels
- □ Replacement parts, available as needed
  - Washer and silicone O-ring, PN XDA622A
     Note: Four sets are needed when performing 6-months, 1-year, or every 2-years maintenance.
  - Reagent syringe piston (except Hgb Lyse), PN GBC030A
     Note: Four syringe pistons are needed when performing two-year maintenance.
  - Hgb Lyse reagent syringe piston, PN GBC031A
  - Silicone O-ring for the Hgb Lyse reagent syringe piston, PN FAA065A
  - Silicone grease, PN XEA019A

#### Preparation

- 1. If off, turn the instrument on.
- 2. From the Service menu, select 9. OTHERS → 3. PARK SYRINGES.
- 3. Turn the instrument off and disconnect the power cord.
- 4. Remove the left side door.

**CAUTION** A flat cable is connected to the rear of the Main card. Be careful when you open the door (where the Main card is attached) that you do not disconnect or damage this cable.

5. Open the Main card door and anchor the door so that it remains open.

#### Removal

1. Disconnect the tubing attached to port 3 at valves 6, 7, 8, 9, and 11 (Figure 4.24-1):

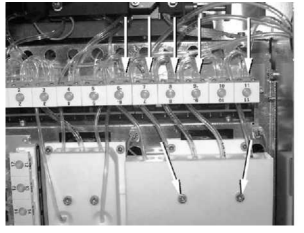
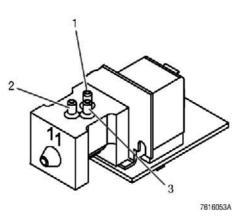


Figure 4.24-1 Valve and Screw Locations - Left Side View



- 2. Remove the two hex screws securing the reagent syringes assembly (Figure 4.24-1). Use a 3 mm Allen wrench.
- 3. Gently remove the reagent syringes assembly from the instrument.
- 4. Hold the reagent syringes assembly over a waste container and drain the syringes by manually pushing the pistons up and down inside the syringe barrels.

Note: Selecting the **PARK SYRINGES** option at the beginning of this procedure, activated an upward movement of the syringe pistons dispensing most of the reagent contained inside the syringe. This manual up and down movement of the pistons is done to drain residual reagent so that all syringes are relatively dry before proceeding.

- 5. Place the drained assembly on absorbent paper.
- 6. Wipe the solenoid valves with a lint-free tissue to remove any reagent that dripped on the valves.

#### **O-Ring, Washer, and Piston Replacement**

If you are replacing an O-ring or washer, replace both. When replacing a syringe piston, replace the O-ring and washer too. Complete needed replacements before proceeding to the Installation instructions.

**CAUTION** With the exception of the Hgb Lyse reagent piston, each syringe piston has both a washer and an O-ring. Compression of the O-ring is critical for a good piston seal. For this reason, the thickness of the washer is matched to the O-ring thickness. As a result, it is important to keep the O-ring and its matching washer together. The Hgb Lyse reagent syringe uses an O-ring but does not use a washer on its piston. See Figure 4.24-2.



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#### Figure 4.24-2 O-rings and Washers - Reagent Syringes Assembly

1. Remove the nine hex screws (CHC M3x12) and the two torx screws (FX M3x12) that secure the bottom plate.

**ATTENTION:** When replacing parts, remove syringes one at a time and in order. Run fingers along piston to check for scratches or scores. The Fix syringe may be slightly discolored.

- 2. If you only need to replace an O-ring or washer, go to step 3. If you need to replace a syringe piston, go to step 4.
- 3. One at a time, replace an old O-ring and washer as follows:
  - a. Remove and discard the old washer and O-ring.

Note: The Hgb Lyse reagent piston has an O-ring but does not have a washer.

- b. Locate the replacement O-ring and washer.
- c. Use a small amount of silicone grease between two fingers to lubricate the O-ring.
- d. Place the O-ring on the syringe piston, followed by the washer (Figure 4.24-2).Note: The Hgb Lyse reagent piston does not have a washer.
- e. Repeat step 3 as many times as necessary then proceed to the IMPORTANT message just before step 5.
- 4. One at a time, replace the old piston, O-ring, and washer as follows:
  - a. Remove and discard the old washer and O-ring.

Note: The Hgb Lyse reagent piston has an O-ring but does not have a washer.

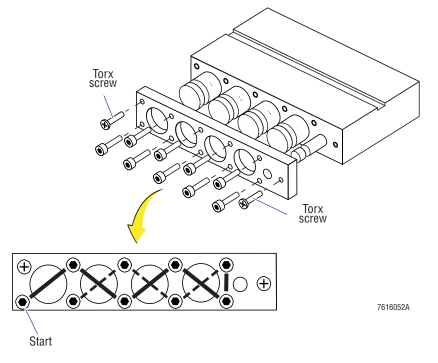
- b. Remove and discard the old syringe piston.
- c. Clean the barrel inside the assembly with lint-free tissues.
- d. Locate the replacement piston, O-ring, and washer.
- e. Use a small amount of silicone grease between two fingers to lubricate the long section of the piston.
- f. Insert the lubricated piston into its barrel.
- g. Use a small amount of silicone grease between two fingers to lubricate the new O-ring.

- h. Place the O-ring on the syringe piston, followed by the washer (Figure 4.24-2).Note: The Hgb Lyse reagent piston does not have a washer.
- i. Repeat step 4 as many times as necessary.

**IMPORTANT** Risk of misleading results. Restricted piston movement affects dilution ratios which may affect final parameter results. The bottom plate on the reagent syringes assembly must be properly aligned to ensure unrestricted movement of the syringe pistons. To ensure alignment, the bottom plate must be installed in two stages with the two torx screws being installed first followed by installation of the nine hex screws. These nine hex screws must be tightened in a zigzag, crisscross pattern to prevent skewing (Figure 4.24-3).

- 5. When all reagent syringes assembly replacements are complete, replace the bottom plate as follows:
  - a. Position the reagent syringes assembly so that the smallest piston is to your right, as shown in Figure 4.24-2.
  - b. Carefully replace the bottom plate. Make sure all O-rings and washers are properly seated inside their respective slots.
  - c. Loosely install the two torx screws one screw in the center opening to the right of the small piston; the second, in the upper left corner (Figure 4.24-3).
  - d. Loosely install the nine hex screws.

#### Figure 4.24-3 Bottom Plate Screw Locations and Tightening Patterns



ATTENTION: Torque needed for the two torx screws is 400 mN.m (56.8 ozf.in).

e. Tighten the two torx screws (Figure 4.24-3). Use a torque driver with a T10 torx bit to tighten the screws to 400 mN.m (56.8 ozf.in).

**ATTENTION:** Torque needed for the nine hex screws is 400 mN.m (56.8 ozf.in). These hex screws must be tighten in a zigzag, crisscross pattern to prevent skewing and ensure proper alignment of the bottom plate (Figure 4.24-3).

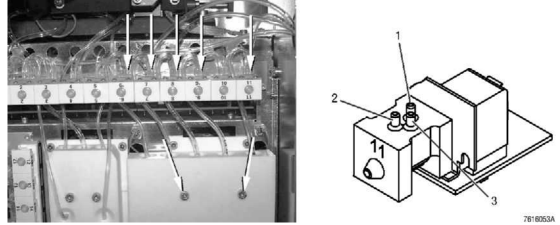
- f. Tighten the nine hex screws in a zigzag, crisscross pattern (Figure 4.24-3).
  - Use a 2.5 mm torque screwdriver to tighten each screw to 400 mN.m (56.8 ozf.in).
  - Start in the lower left corner and use a zigzag pattern (from left to right) to tighten every other screw. Then reverse the zigzag pattern. Start in the upper right corner and use a right-to-left zigzag pattern to tighten the remaining loose screws (as you crisscross the previous zigzag pattern).

#### Installation

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- 1. Align the reagent syringes assembly back inside the instrument. Make sure the channel on the back of the assembly housing fits securely on the motor housing guide.
- 2. Replace the two hex screws that secure the reagent syringes assembly to the instrument (Figure 4.24-4). Use a 3 mm Allen wrench.

#### Figure 4.24-4 Valve and Screw Locations - Left Side View



3. Thread each tubing up through the hole and reconnect at port 3 of valves 6, 7, 8, 9 and 11 (Figure 4.24-4).

At the reagent syringes assembly, from left to right:

- Attach the first tubing (from the Hgb Lyse reagent syringe) to LV6, port 3.
- Attach the second tubing (from the Rinse reagent syringe) to LV7, port 3.
- Attach the third tubing (from the Fix reagent syringe) to LV8, port 3.
- Attach the fourth tubing (from the Diluent reagent syringe) to LV9, port 3.
- Attach the fifth tubing (from the WBC Lyse reagent syringe) to LV11, port 3.

#### Verification

- 1. Reconnect the power cord.
- 2. Turn the instrument on.

- 3. When the startup routine and background check are done, verify the reagent syringes assembly is not leaking.
- 4. If no leakage is seen, prime the reagents.
   From the Main Menu, select 3. REAGENTS → 3. PRIME → 6. ALL REAGENTS.
- 5. When the prime cycles are done, check the reagent syringes assembly for leaks.
- 6. If no leaks are detected, close the Main card door and reattach the left side door.
- 7. Perform a system verification. See Heading 5.3, SYSTEM VERIFICATION PROCEDURES.

#### 4.25 COUNT SYRINGE COMPONENT REPLACEMENTS

#### Purpose

Use this procedure to remove and replace the count syringe piston, O-ring, or washer. When performing this procedure to replace an O-ring or a washer, replace both. When replacing the count syringe piston, replace its O-ring and washer too.

#### **Tools/Supplies Needed**

- □ Allen wrenches, 2.5 mm, 3 mm
- □ T10 torx driver
- □ Absorbent paper towels
- **C** Replacement components, available as needed
  - Washer and O-ring, PN XDA621A
  - Count syringe piston, PN GBG052A
  - Silicone grease, PN XEA019A

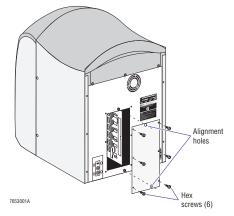
#### Preparation

- 1. If off, turn the instrument on.
- 2. From the Service menu, select 9. OTHERS >> 3. PARK SYRINGES.
- 3. Turn the instrument off.
- 4. Remove the left side door.

**CAUTION** A flat cable is connected to the rear of the Main card. Be careful when you open the door (where the Main card is attached) that you do not disconnect or damage this cable.

- 5. Open the Main card door and anchor the door so that it remains open.
- 6. At the rear of the instrument,
  - a. Disconnect the power cord.
  - b. Remove the six hex screws securing the rear access panel (Figure 4.25-1). Use a 3 mm Allen wrench.

#### Figure 4.25-1 Rear Access Panel Screw Locations



7. Remove the rear access panel and set it aside.

#### Removal

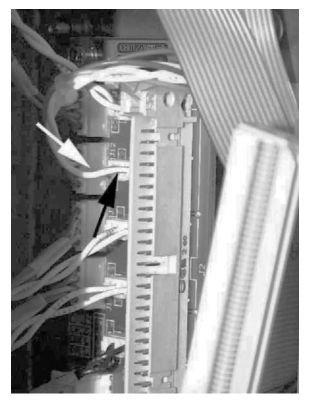
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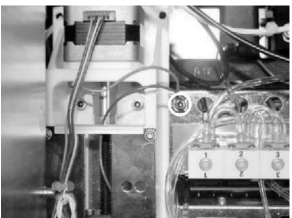
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- 1. At the Motor Interconnect card (Figure 4.25-2):
  - a. Disconnect the large (unlabeled) cable connector from the J2 card connector on the card. This allows better access to the individual motor and sensor connectors.
  - b. Disconnect sensor connector P7. In Figure 4.25-2, the dark arrow is pointing to this connector. (The J7 connector on the card is oriented in a horizontal position behind the large connector on the Motor Interconnect card. It is the second horizontal connector from the top of the Motor Interconnect card.)
  - c. Disconnect motor connector P12. In Figure 4.25-2, the light arrow is pointing to this connector. (The J12 connector on the card is oriented in a vertical position, the second vertical connector from the top of the Motor Interconnect card.)

#### Figure 4.25-2 Motor Interconnect Card - Count Syringe Connector Locations

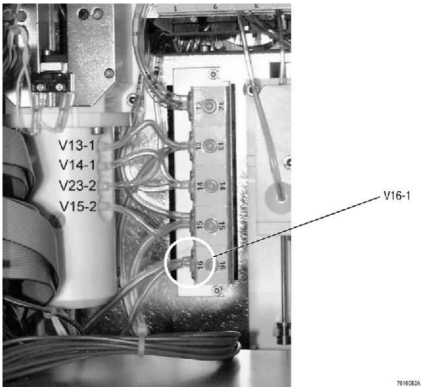


- 2. In the left side compartment,
  - a. Locate the count syringe assembly.
  - b. Remove the ground wire from the instrument's frame (Figure 4.25-3).



#### Figure 4.25-3 Count Syringe - Ground Wire Location

- 3. Place absorbent paper under the count syringe assembly and the 5-valve assembly to the right of the syringe assembly.
- 4. Remove the four tubings attached to the fittings on the right side of the syringe housing (Figure 4.25-4).



#### Figure 4.25-4 Count Syringe Housing - Tubing Locations

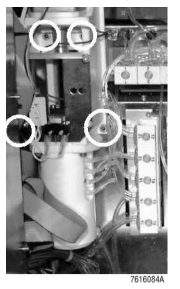
5. Locate the tubing attached to the bottom fitting. Trace this tubing to its connection point on solenoid valve 16. Remove the tubing from LV16, port 1.

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**ATTENTION:** The count syringe assembly uses captive hex screws mounted inside rubber shock mounts. It is recommended that you use only four turns to either loosen or tighten these screws. Turning the hex screw too many counterclockwise rotations may separate the screw from the rubber shock mount. If all Service Representatives consistently use four turns to remove or install these hex screws, it is unlikely that a rubber shock mount will separate from its hex screw and fall inside the instrument.

6. Locate the four hex screws (CHC M4x16) shown in Figure 4.25-5. These are captive screws anchored inside rubber shock mounts. Use a 3 mm Allen wrench.



#### Figure 4.25-5 Count Syringe - Captive Screw Locations

7. Loosen each hex screw with four counterclockwise rotations (Figure 4.25-5).

Note: If a screw is not released in four rotations, the screw was overtightened the last time the count syringe assembly was serviced. From this point, make a single rotation then check to see if the mount is free. Repeat this sequence of making a single rotation and checking until the mount is free.

**ATTENTION:** Ensure the electronic cables are free and unrestricted as you remove the count syringe assembly from the instrument. Do not continue to pull on the count syringe assembly if these cables become restricted. Locate and free the restriction before proceeding.

- 8. Gently remove the count syringe from the instrument. Make sure the electronic cables remain unrestricted.
- 9. Hold the count syringe assembly over a waste container and push the piston up and down several times to drain the syringe barrel.

Note: Selecting the **PARK SYRINGES** option at the beginning of this procedure, activated an upward movement of the syringe piston which expelled most of the reagent inside this syringe. This manual up and down movement of the piston is done to drain residual liquid so that the syringe is relatively dry before proceeding.

10. Place the drained assembly on absorbent paper.

#### **O-ring, Washer, and Piston Replacement**

- If you only need to replace an O-ring on a piston or a washer on a housing, replace both using the instructions under the O-ring and Washer Replacement Only heading.
- If you need to replace the syringe piston, replace the O-ring and washer too using the instructions under the Piston Replacement heading.

#### **Piston Replacement**

1. Remove the four torx screws securing the syringe guide plate (Figure 4.25-6). Use a T10 torx driver.

#### Torx Piston screws (3) screws (4) Ground 1 wire Shaft housing É 9 Syringe guide plate 0-ring Syringe Anti-extrusion piston ring (new style assemblies only) Washer in the second Syringe cover Syringe cover screws (4) 7616113B

Figure 4.25-6 Count Syringe - Piston, O-ring, and Washer Replacement

- 2. Remove the syringe guide plate.
- 3. Remove the four hex screws (CHC M4x16) located around the perimeter of the syringe cover. Use a 3 mm Allen wrench.
- 4. Remove the cover to expose the syringe piston, O-ring, washer and anti-extrusion ring.
- 5. Remove and discard the O-ring and washer but retain the anti-extrusion ring.

ATTENTION: Be careful to keep the syringe drive nut in place on the motor lead screw.

- 6. Manually move piston up to expose the three piston screws.
- 7. Remove the three piston screws (CHC M3x16). Use a 2.5 mm Allen wrench (one screw has the ground wire). Make sure the shaft housing remains stable.
- 8. Pull the piston off the shaft housing and remove it by passing it through the end of the frame.

- 9. Replace the old piston, O-ring, and washer as follows (Figure 4.25-6):
  - a. Clean the barrel inside the syringe assembly with lint-free tissues.
  - b. Locate the replacement piston, O-ring, and washer.
  - c. Position the new piston in the housing and replace the three piston screws (with the ground wire attached). Use a 2.5 mm Allen wrench.
  - d. Use a small amount of silicone grease between two fingers to lubricate the long section of the piston and the new O-ring.
  - e. Place the O-ring on the syringe piston.

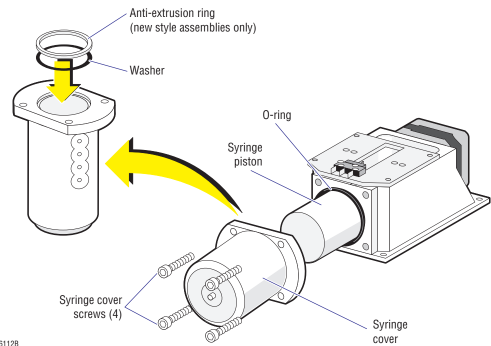
**IMPORTANT** Risk of erroneous results. The washer must be properly seated in the rim of the syringe housing before reassembling the piston and housing.

- f. Seat washer on rim of syringe housing followed by the anti-extrusion ring.
- 10. Replace the syringe cover. Make sure the notches on the cover and syringe are aligned. Reinstall the four hex screws around the perimeter of the syringe cover. Use a 3 mm Allen wrench. Make sure washer is not pinched.
- 11. Position the guide plate on the syringe assembly (notched corners toward the bottom) then secure the plate to the assembly with the four torx screws. Use a T10 torx driver.
- 12. Go to the Installation heading.

#### **O-ring and Washer Replacement Only**

1. Remove the four hex screws (CHC M4x16) located around the perimeter of the syringe cover (Figure 4.25-7).

#### Figure 4.25-7 Count Syringe - O-ring and Washer Replacement



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- 2. Remove the cover to expose the syringe piston, O-ring, washer and anti-extrusion ring.
- 3. Remove and discard the O-ring and washer but retain the anti-extrusion ring.
- 4. Replace the old O-ring and washer as follows (Figure 4.25-7):
  - a. Locate the replacement O-ring and washer.
  - b. Use a small amount of silicone grease between two fingers to lubricate the O-ring.
  - c. Place the O-ring on the syringe piston.

**IMPORTANT** Risk of erroneous results. The washer must be properly seated in the rim of the syringe housing before reassembling the piston and housing.

- d. Seat washer on rim of syringe housing followed by the anti-extrusion ring.
- 5. Replace the syringe cover. Make sure the notches on the cover and syringe are aligned. Reinstall the four hex screws around the perimeter of the syringe cover. Use a 3 mm Allen wrench. Make sure washer is not pinched.

#### Installation

1. Position the count syringe assembly back inside the instrument.

**Note:** Make sure the rubber shock mounts are positioned behind the panel. The syringe assembly should be flush against the instrument frame.

**ATTENTION:** It is recommended that you use only four clockwise turns to tighten these captive hex screws. As each hex screw is tighten, its rubber shock mount expands to secure the syringe assembly to the instrument frame. If all Service Representatives consistently use four turns to install these hex screws, it is unlikely that a rubber shock mount will separate from its hex screw and fall inside the instrument the next time the assembly is removed.

- 2. Tighten the four captive hex screws using four clockwise rotations (Figure 4.25-5). Use a 3 mm Allen wrench.
- 3. Reattach the four tubings to the fittings on the right side of the syringe housing (Figure 4.25-8).

From the top fitting to the bottom fitting, attach the tubings as follows:

- Attach the tubing connected to solenoid valve 13, port 1 to the top fitting on the count syringe housing.
- Attach the tubing connected to solenoid valve 14, port 1 to the second fitting on the count syringe housing.
- Attach the tubing connected to solenoid valve 23, port 2 to the third fitting on the count syringe housing.
- Attach the tubing connected to solenoid valve 15, port 2 to the bottom fitting on the count syringe housing.
- 4. Route the tubing attached to the bottom fitting of the syringe housing to solenoid valve 16. Attach the tubing to port 1.
- 5. Reconnect the ground wire to the instrument frame (Figure 4.25-3).
- 6. Route the motor and sensor cables through the opening to the left of the count syringe assembly. Route the connectors over to the Motor Interconnect card.

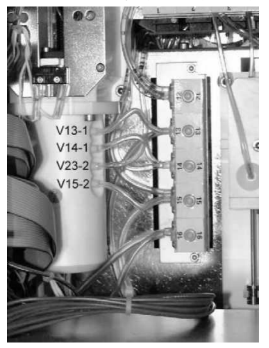


Figure 4.25-8 Count Syringe Housing - Tubing Locations

- 7. At the Motor Interconnect card (Figure 4.25-2):
  - a. Reattach motor connector P12. In Figure 4.25-2, the light arrow is pointing to this connector. (The J12 connector on the card is oriented in a vertical position near the top of the Motor Interconnect card).
  - b. Reattach sensor connector P7. In Figure 4.25-2, the dark arrow is pointing to this connector. (The J7 connector on the card is oriented in a horizontal position behind the large connector on the Motor Interconnect card).
  - c. Reattach the large, unlabeled cable connector to the J2 card connector.
- 8. Replace the rear access panel using the six hex screws removed earlier (Figure 4.25-1).
- 9. Reconnect the power cord.

#### Verification

- 1. Turn the instrument on. An automatic startup and background check are performed.
- 2. When the startup routine and background check are done, verify the count syringe is not leaking.
- 3. If no leaks are detected, close the Main card door and reattach the left side door.
- 4. Perform a system verification. See Heading 5.3, SYSTEM VERIFICATION PROCEDURES.

### 4.26 SAMPLE PROBE AND RINSE BLOCK ASSEMBLY COMPONENT REPLACEMENTS

#### Purpose

Use this procedure to remove and replace the sample probe or any part of the rinse block assembly including the probe guide, O-ring, or rinse block.

#### **Tools/Supplies Needed**

□ Torque screwdriver, 2.5 mm, hex-ball

Note: 100 mN.m (14.2 ozf.in) torque is required in this procedure.

- □ Replacement parts, available as needed
  - Sample probe, PN XDA619AS
  - Sample probe guide, PN GBG091A
  - O-ring, flurocarbon, PN FAA053A
  - Rinse block, PN GBG090A
  - Silicone grease, PN XEA019A

#### Preparation

- 1. If off, turn the instrument on.
- 2. From the Main Menu, select **4. DIAGNOSTICS → 4. HARDWARE SYSTEMS → 4. TRAVERSE SERVICE POSITION**.
- 3. Turn the instrument off.
- 4. Open the pneumatic access door (right side of the instrument).

#### Removal

**ATTENTION:** Note how the existing sample probe and rinse block assembly are seated.

1. Loosen the two hex screws (CHC M3x25) securing the sample probe rinse block assembly on the traverse (Figure 4.26-1). Use a 2.5 mm hex-ball Allen wrench.

#### Figure 4.26-1 Probe Rinse Block Screw Locations



2. Lift the sample probe lock-lever to free the probe then remove both the probe and rinse block assembly at the same time (Figures 4.26-2 and 4.26-3).

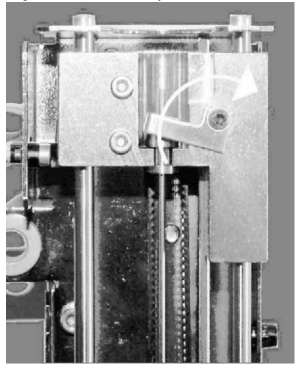
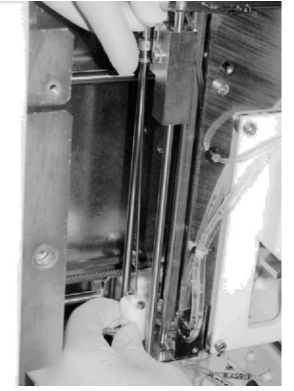


Figure 4.26-2 Lift the Sample Probe Lock-Lever



- 3. Separate the sample probe and the rinse block assembly.
  - If the sample probe needs replaced, go to the Sample Probe Replacement heading.
  - If one or more rinse block assembly components need replaced,
    - Place the probe back on its support and close the lock-lever.
    - Go to the Rinse Block Assembly Component Replacement heading.

#### Sample Probe Replacement

- 1. Remove the tubing attached to the top of the old probe.
- 2. Attach the new probe.
  - If replacements in the rinse block assembly are required,
    - Place the probe back on its support and close the lock.
    - Go to the Rinse Block Assembly Component Replacement heading.
  - If no other replacements are needed, go to the Installation heading.

### **Rinse Block Assembly Component Replacement**

The rinse block assembly consists of three components: (from top to bottom) the probe guide, a fluorocarbon O-ring, and the rinse block. Complete needed replacements before proceeding to the Installation instructions.



- 1. Remove the two hex screws.
- 2. Separate the rinse block assembly and replace components as needed. See step 3 to replace the rinse block and step 4 to replace the O-ring or probe guide.

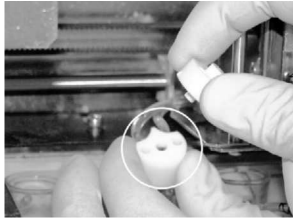
**IMPORTANT** Risk of erroneous results due to pinching of the tubing. When reattaching tubing from Valves 17 and 18 to new rinse block, route tubing from Valve 17 on inside of guide post, and tubing from Valve 18 around outside of guide post.

- 3. To replace the rinse block:
  - a. Place the new rinse block next to the old rinse block. Make sure the new rinse block is oriented the same as the old with the top up and the ports towards the instrument.
  - b. Remove tubing from the old rinse block. Route tubing from Valve 18 around outside of guide post and attach to port 1 on the new rinse block. Route tubing from Valve 17 on inside of guide post and attach to port 2 on the new rinse block.
- 4. To replace the O-ring:
  - a. Remove and discard the old O-ring.
  - b. Clean then dry the two white blocks. Use a twisted lint free tissue to clean the center opening.
  - c. Locate the replacement O-ring.
  - d. Use a small amount of silicone grease between two fingers to lubricate the O-ring.
  - e. Place the new O-ring inside the probe guide (top block).

Note: Use a new probe guide if needed.

- 5. When all rinse block component replacements are complete,
  - a. Reassemble the rinse block (Figure 4.26-4).

#### Figure 4.26-4 Reassemble the Rinse Block Assembly



- b. Route the two hex screws through the openings at the top of the assembly.
- c. Proceed to the Installation heading to complete this procedure.

#### Installation

- 1. Lift the probe lock-lever and remove the probe from its support (Figure 4.26-2).
- 2. Position the probe inside the probe guide, using care not to bend the probe.
- 3. Lift the probe lock and return the probe and the rinse block assembly back in its original position on the traverse. Position the notch and tubing towards the front of the instrument.
- 4. Seat the collar of the sample probe by gently pushing it into place then close the lock-lever.

ATTENTION: Torque needed for the rinse block assembly screws is 100 mN.m (14.2 ozf.in).

5. Tighten the two hex screws to secure the rinse block assembly to the traverse carriage (Figure 4.26-1). Use a 2.5 mm, hex-ball torque screwdriver to tighten the screws to 100 mN.m (14.2 ozf.in).

#### Verification

- 1. Turn the instrument on.
- 2. When the startup routine and background check are done, verify the sample probe and the rinse block assembly are not leaking.
- 3. Close the right side door.
- 4. Perform a system verification. See Heading 5.3, SYSTEM VERIFICATION PROCEDURES.

# 4.27 WASTE SYRINGE COMPONENT REPLACEMENTS

#### Purpose

Use this procedure to remove and replace the waste syringe piston, O-ring, or washer. When performing this procedure to replace an O-ring or a washer, replace both. When replacing the waste syringe piston, replace its O-ring and washer as well.

#### **Tools/Supplies Needed**

- □ Allen wrenches, 2.5 mm, 3 mm
- □ Cutting pliers
- □ T10 torx driver
- □ Absorbent paper towels
- □ Replacement components, available as needed
  - Washer and O-ring, PN XDA621A
  - Waste syringe piston, PN GBG052A
  - Silicone grease, PN XEA019A

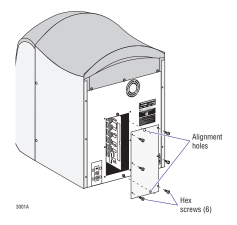
#### Preparation

- 1. Turn the instrument off.
- 2. Remove the left side door.

**CAUTION** A flat cable is connected to the rear of the Main card. Be careful when you open the door (where the Main card is attached) that you do not disconnect or damage this cable.

- 3. Open the Main card door and anchor the door so that it remains open.
- 4. At the rear of the instrument,
  - a. Disconnect the power cord.
  - b. Remove the six hex screws securing the rear access panel (Figure 4.27-1). Use a 3 mm Allen wrench.

#### Figure 4.27-1 Rear Access Panel Screw Locations



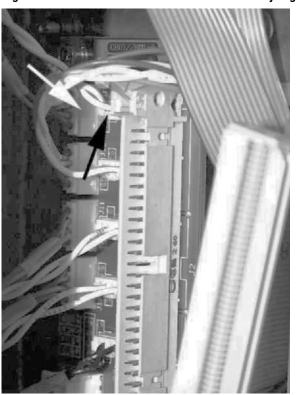
5. Remove the rear access panel and set it aside.

#### Removal

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- 1. At the Motor Interconnect card (Figure 4.27-2),
  - a. Disconnect the large, unlabeled cable connector from the J2 card connector. This allows better access to the individual motor and sensor connectors.
  - b. Disconnect sensor connector P8. In Figure 4.27-2, the dark arrow is pointing to this connector. (The J8 connector on the card is oriented in a horizontal position behind the large connector on the Motor Interconnect card.)
  - c. Disconnect motor connector P13. In Figure 4.27-2, the light arrow is pointing to this connector. (The J13 connector on the card is oriented in a vertical position near the top of the Motor Interconnect card.)



#### Figure 4.27-2 Motor Interconnect Card - Waste Syringe Connector Locations

- 2. In the left side compartment,
  - a. Locate the waste syringe assembly.
  - b. Cut the tie wrap (Figure 4.27-3).
  - a. Remove the ground wire from the instrument's frame.



Figure 4.27-3 Waste Syringe - Tie Wrap and Ground Wire Location

**WARNING** Risk of personal contamination. If you do not properly shield yourself while servicing the waste syringe assembly, you may become contaminated. To prevent possible biological contamination, you must wear appropriate safety glasses, a lab coat, and gloves when servicing the waste syringe assembly.

- 3. Place absorbent paper under the waste syringe and the 7-valve assembly to its left.
- 4. At the waste syringe, disconnect the two tubings as follows:
  - a. Remove the tubing attached to the left side fitting (port 1).
  - b. Locate the tubing attached to the bottom fitting (port 2). Trace this tubing to its connection point on LV26. Remove the tubing from LV26, port 3.

**Note:** To minimize waste exposure, use a hemostat to clamp this tubing between the 7-valve assembly and the waste syringe <u>before</u> removing the tubing from LV26.

**ATTENTION:** The waste syringe assembly uses captive hex screws mounted inside rubber shock mounts. It is recommended that you use only four turns to either loosen or tighten these screws. Turning the hex screw more than four counterclockwise rotations may separate the screw from the rubber shock mount. If all Service Representatives consistently use four turns to remove or install these hex screws, it is unlikely that a rubber shock mount will separate from its hex screw and fall inside the instrument.

5. Locate the four hex screws (CHC M4x16) circled in Figure 4.27-4. These are captive screws anchored inside rubber shock mounts. Use a 3 mm Allen wrench.

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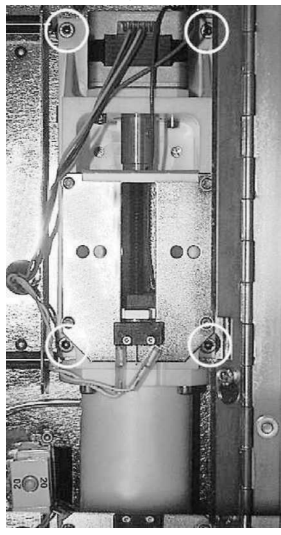


Figure 4.27-4 Waste Syringe - Captive Screw Locations

6. Loosen each hex screw with four counterclockwise rotations (Figure 4.27-4).

**Note:** If a screw is not released in four rotations, the screw was overtightened the last time the waste syringe assembly was serviced. From this point, make a single rotation then check to see if the mount is free. Repeat this sequence of making a single rotation and checking until the mount is free.

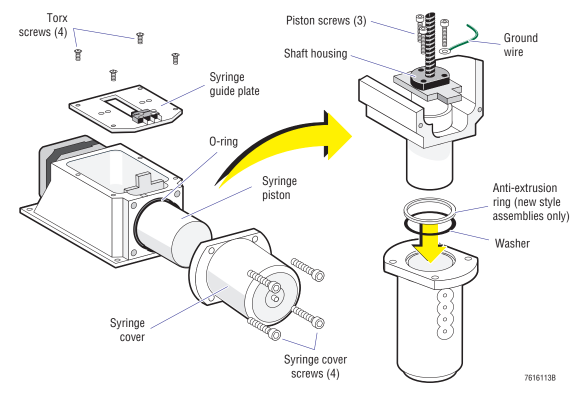
**ATTENTION:** Ensure the electronic cables are free and unrestricted as you remove the waste syringe assembly from the instrument. Do not continue to pull on the waste syringe assembly if these cables become restricted. Locate and free the restriction before proceeding.

7. Gently remove the waste syringe from the instrument. Make sure the electronic cables remain unrestricted.

- 8. Hold the waste syringe assembly over a biohazardous waste container and push the piston up and down several times to drain the syringe barrel.
- 9. Place the drained assembly on absorbent paper.

#### **Piston Replacement**

1. Remove the four torx screws securing the syringe guide plate (Figure 4.27-5). Use a T10 torx driver.



#### Figure 4.27-5 Waste Syringe - Piston, O-ring, and Washer Replacement

- 2. Remove the syringe guide plate.
- 3. Remove the four hex screws (CHC M4x16) located around the perimeter of the syringe cover. Use a 3 mm Allen wrench.
- 4. Remove the cover to expose the syringe piston, O-ring, washer and anti-extrusion ring.
- 5. Remove and discard the O-ring and washer but retain the anti-extrusion ring.

**ATTENTION:** Be careful to keep the syringe drive nut in place on the motor lead screw.

- 6. Manually move piston up to expose the three piston screws.
- 7. Remove the three piston screws (CHC M3x16). Use a 2.5 mm Allen wrench (one screw has the ground wire). Make sure the shaft housing remains stable.
- 8. Pull the piston off the shaft housing and remove it by passing it through the end of the frame.

- 9. Replace the old piston, O-ring, and washer as follows (Figure 4.27-5):
  - a. Clean the barrel inside the syringe assembly with lint-free tissues.
  - b. Locate the replacement piston, O-ring, and washer.
  - c. Position the new piston in the housing and replace the three piston screws (with the ground wire attached). Use a 2.5 mm Allen wrench.
  - d. Use a small amount of silicone grease between two fingers to lubricate the long section of the piston and the new O-ring.
  - e. Place the O-ring on the syringe piston.

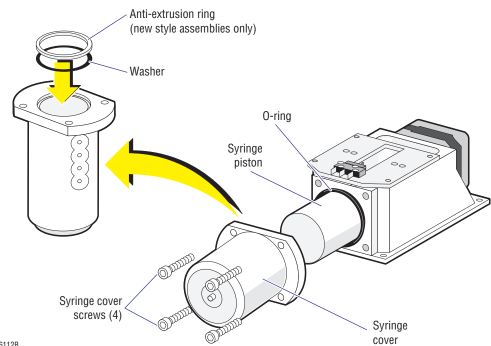
**IMPORTANT** Risk of erroneous results. The washer must be properly seated in the rim of the syringe housing before reassembling the piston and housing.

- f. Seat washer on rim of syringe housing followed by the anti-extrusion ring.
- 10. Replace the syringe cover. Make sure the notches on the cover and syringe are aligned. Reinstall the four hex screws around the perimeter of the syringe cover. Use a 3 mm Allen wrench. Make sure washer is not pinched.
- 11. Position the guide plate on the syringe assembly (notched corners toward the bottom) then secure the plate to the assembly with the four torx screws. Use a T10 torx driver.
- 12. Go to the Installation heading.

#### **O-ring and Washer Replacement Only**

1. Remove the four hex screws (CHC M4x16) located around the perimeter of the syringe cover (Figure 4.27-6).

#### Figure 4.27-6 Waste Syringe - O-ring and Washer Replacement



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- 2. Remove the cover to expose the syringe piston, O-ring, washer and anti-extrusion ring.
- 3. Remove and discard the O-ring and washer but retain the anti-extrusion ring.
- 4. Replace the old O-ring and washer as follows (Figure 4.27-6):
  - a. Locate the replacement O-ring and washer.
  - b. Use a small amount of silicone grease between two fingers to lubricate the O-ring.
  - c. Place the O-ring on the syringe piston.

**IMPORTANT** Risk of erroneous results. The washer must be properly seated in the rim of the syringe housing before reassembling the piston and housing.

- d. Seat washer on rim of syringe housing followed by the anti-extrusion ring.
- 5. Replace the syringe cover. Make sure the notches on the cover and syringe are aligned. Reinstall the four hex screws around the perimeter of the syringe cover. Use a 3 mm Allen wrench. Make sure washer is not pinched.

#### Installation

1. Position the waste syringe assembly back inside the instrument.

**Note:** Make sure the rubber shock mounts are positioned behind the panel. The syringe assembly should be flush against the instrument frame.

**ATTENTION:** It is recommended that you use only four clockwise turns to tighten these captive hex screws. As each hex screw is tighten, its rubber shock mount expands to secure the syringe assembly to the instrument frame. If all Service Representatives consistently use four turns to install these hex screws, it is unlikely that a rubber shock mount will separate from its hex screw and fall inside the instrument the next time the assembly is removed.

- 2. Tighten the four captive hex screws using four clockwise rotations (Figure 4.27-4). Use a 3 mm Allen wrench.
- 3. Reconnect the two hydraulic tubings as follows:
  - a. Attach the tubing connected to S20 (port 1) to the left side fitting (port 1) on the waste syringe assembly.
  - b. Route the tubing attached to the bottom fitting (port 2) to S26. Attach the tubing (with the collar) to port 3.
- 4. Reconnect the grounding wire to the instrument frame (Figure 4.27-3).
- 5. Route the motor and sensor cables through the opening to the left of the waste syringe assembly (Figure 4.27-3). Route the connectors over to the Motor Interconnect card.

- 6. At the Motor Interconnect card (Figure 4.27-2),
  - a. Reattach motor connector P13. In Figure 4.27-2, the light arrow is pointing to this connector. (The J13 connector on the card is oriented in a vertical position near the top of the Motor Interconnect card).
  - b. Reattach sensor connector P8. In Figure 4.27-2, the dark arrow is pointing to this connector. (The J8 connector on the card is oriented in a horizontal position behind the large J2 connector on the Motor Interconnect card).
  - c. Reattach the large unlabeled cable connector to the J2 card connector.
- 7. Replace the rear access panel using the six hex screws removed earlier (Figure 4.27-1).
- 8. Reconnect the power cord.

#### Verification

- 1. Turn the instrument on. An automatic startup and background check are performed.
- 2. When the startup routine and background check are done, open the right side door and verify the waste syringe is not leaking.
- 3. If no leaks are detected, perform a system verification. See Heading 5.3, SYSTEM VERIFICATION PROCEDURES.

### 4.28 CLEANING THE BATH ENCLOSURE

#### Purpose

Use this procedure to clean bath enclosure parts with diluted bleach. This procedure provides instructions for cleaning the:

- Black plastic that covers the baths (attached to the right side door).
- White plastic under the baths.
- Top of the hemoglobin bath.

#### **Tools/Supplies Needed**

- □ High quality, fragrance-free bleach (10-12% sodium hypochlorite available chloride)
- Deionized water
- □ Absorbent paper towels
- □ Lint-free tissues

#### Procedure

- 1. Prepare a 1:5 bleach solution: 4 parts deionized water to 1 part high quality, fragrance-free bleach (10-12% sodium hypochlorite available chloride).
- 2. Turn the instrument off.
- 3. Open the pneumatic access door (right side of the instrument).

**WARNING** Risk of personal contamination. If you do not properly shield yourself while servicing the waste syringe assembly, you may become contaminated. To prevent possible biological contamination, you must wear appropriate safety glasses, a lab coat, and gloves when servicing the waste syringe assembly.

**ATTENTION:** Do not use a sponge or cloth to clean instrument assemblies. Use absorbent paper towels that can be discarded in a biohazardous container. For small or sensitive assemblies such as the top of the hemoglobin bath, use lint-free tissues.

- 4. Pour the diluted bleach on a soft paper towel and thoroughly clean the black plastic that covers the baths (attached to the right side door) and the white plastic under the baths.
- 5. Wipe the plastic dry with a clean paper towel.
- 6. Discard the towels in a biohazardous container.
- 7. Wet a lint-free tissue with the diluted bleach and clean the top of the hemoglobin bath.
- 8. Wipe the top of the bath dry with a clean lint-free tissue.
- 9. Discard the tissues in a biohazardous container.
- 10. Close the pneumatic access door (right side of the instrument).
- 11. Turn the instrument on. An automatic startup and background check is performed.
- 12. When the startup routine and background check are done, resume normal operation.

# **SERVICE AND REPAIR PROCEDURES** *CLEANING THE BATH ENCLOSURE*

# 4.29 5diff SYRINGE ASSEMBLY REPLACEMENTS

#### Purpose

The 5diff syringe consists of one large center piston with a smaller injector piston on each side. Use this procedure to remove and replace the 5diff syringe O-rings associated with this center piston and the two injector pistons. Two types of O-rings are used in the 5diff syringe. The large silicone O-ring is used for the center piston and two smaller fluorocarbon O-rings are used on each injector piston. Although this procedure focuses on O-ring replacement, it could be used as a guide in replacing any component inside the 5diff syringe housing or for replacing the entire 5diff syringe assembly.

#### **Tools/Supplies Needed**

- □ Allen wrenches, 2.5 mm, 3 mm
- □ Torque screwdriver, 2.5 mm and T10 torx

Note: 400 mN.m (56.8 ozf.in) torque is required in this procedure.

- □ Replacement components
  - Silicone O-ring, PN FAA040A
  - Four fluorocarbon O-rings, PN FAA067A
  - Silicone grease, PN XEA019A

#### Preparation

- 1. If off, turn the instrument on.
- 2. From the Service menu, select 9. OTHERS → 3. PARK SYRINGES.
- 3. Turn the instrument off and unplug the power cord from the instrument or the wall.
- 4. Remove the left side panel from the instrument. For details, see Heading 4.2. Set the panel aside.

**CAUTION** A flat cable is connected to the rear of the Main card. Be careful when you open the door (where the Main card is attached) that you do not disconnect or damage this cable.

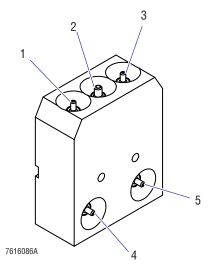
- 5. In the left side compartment,
  - a. Turn the two captive knobs counterclockwise to release the Main card door. These knobs are located to the right of the Main card.
  - b. Open the Main card door and carefully anchor the door behind the white plastic catch to keep it open.

#### Removal

**WARNING** Risk of personal contamination. If you do not properly shield yourself while servicing the 5diff syringe assembly, you may become contaminated. To prevent possible biological contamination, you must wear appropriate safety glasses, a lab coat, and gloves when servicing the 5diff syringe assembly.

- 1. At the 5diff syringe assembly, disconnect the two tubings associated with the flow cell:
  - a. Disconnect the tubing attached to port 1 (Figure 4.29-1).
  - b. Disconnect the tubing attached to port 5 (Figure 4.29-1).

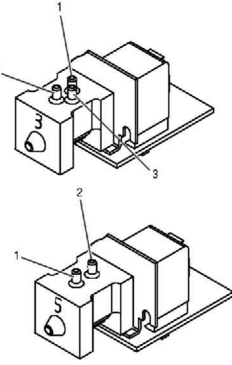




- 2. Disconnect the other three tubings at the 11-valve assembly (Figure 4.29-2):
  - a. Disconnect the tubing attached to valve 2, port 2.(The other end of this tubing is attached to 5diff syringe port 4, the lower left port.)
  - b. Disconnect the tubing attached to valve 3, port 3.(The other end of this tubing is attached to 5diff syringe port 2, upper middle port.)
  - c. Disconnect the tubing attached to valve 5, port 1.(The other end of this tubing is attached to 5diff syringe port 3, upper right port.)

Figure 4.29-2 Valve and Tubing Locations - Left Side View





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3. Remove the two screws on the front of the 5diff syringe housing (Figure 4.29-3). Use a 3.0 mm Allen wrench.



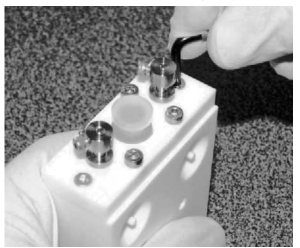
Figure 4.29-3 Screw Locations - 5diff Syringe

- 4. Remove the 5diff syringe assembly.
- 5. Place the syringe assembly on absorbent paper.

#### **O-ring Replacements**

- 1. At the bottom plate (Figure 4.29-4),
  - a. Remove the two torx screws (FX M3x12) that are flush with the bottom plate. Use a T10 torx driver.
  - b. Remove the four hex screws (CHC M3x12) using a 2.5 mm Allen wrench.

#### Figure 4.29-4 Bottom Plate - 5diff Syringe



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2. Separate the bottom plate (and attachments) from the 5diff syringe housing (Figure 4.29-5).

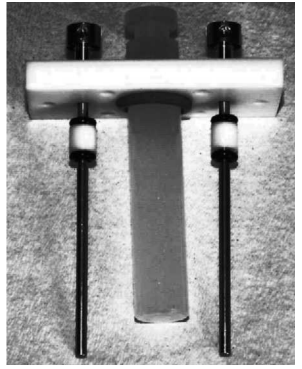


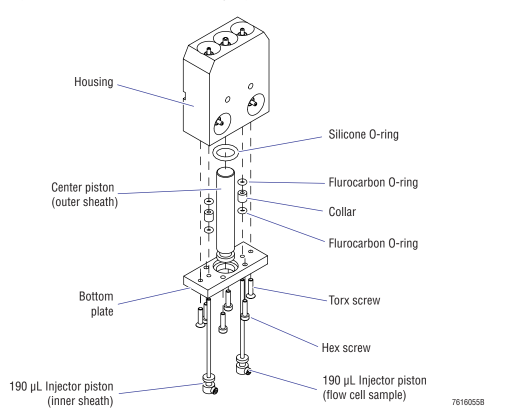
Figure 4.29-5 Bottom Plate and Attachments - Housing Removed

- 3. Clean then dry the outside of the 5diff syringe housing.
- 4. Replace the O-rings on the 190  $\mu$ L injector piston located on each side of the center syringe piston (Figure 4.29-6):
  - a. Remove the two O-rings and the collar from each injector piston.
  - b. Discard the four old O-rings. Save the two collars.
  - c. Clean the outside of the injector pistons, the center piston, and the bottom plate.
  - d. Locate the four replacement fluorocarbon O-rings.
  - e. Use a small amount of silicone grease between two fingers to lubricate an O-ring then place it on the injector piston.
  - f. Place the collar on the injector piston.
  - g. Use a small amount of silicone grease between two fingers to lubricate another O-ring then place it on the injector piston.
  - h. Repeat steps e through g to replace the O-rings on the other injector piston.

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#### Figure 4.29-6 Illustrated Parts - 5diff Syringe

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- 5. Use a small amount of silicone grease between two fingers to lubricate the silicone O-ring before installing this O-ring on the syringe piston.
- 6. Once all replacements are complete, replace the bottom plate as follows (Figure 4.29-6):
  - a. Carefully replace the bottom plate. Make sure the piston O-ring is properly seated inside the slotted opening on the bottom plate.
  - b. Loosely install the two torx screws in the two outside openings.
  - c. Loosely install the four hex screws.

ATTENTION: Torque needed for the two torx screws is 400 mN.m (56.8 ozf.in).

d. Tighten the two torx screws. Use a torx driver with a T10 torx bit to tighten the screws to 400 mN.m (56.8 ozf.in).

**ATTENTION:** Torque needed for the four hex screws is 400 mN.m (56.8 Ozf.in). These hex screws must be tightened in a crisscross pattern to prevent skewing and ensure alignment of the bottom plate.

- e. Tighten the four hex screws in a crisscross pattern.
  - Use a 2.5 mm torque screwdriver to tighten each screw to 400 mN.m (56.8 ozf.in).
  - Start by tightening the lower left hex screw then move to the upper right. Continue the process by tightening the upper left hex screw then move to the lower right.

#### Installation

- 1. Position the 5diff syringe assembly back inside the instrument. Make sure the channel on the back of the assembly housing fits securely on the motor housing guide. Use a 3 mm Allen wrench.
- 2. Replace the two hex screws that secure the 5diff syringe assembly to the instrument.
- 3. At the 5diff syringe assembly, attach the two tubings associated with the flow cell (Figure 4.29-1):
  - a. Connect the loose tubing near the top of assembly to port 1 (upper left port).
  - b. Connect the loose tubing near the bottom of the assembly to port 5 (lower right port).
- 4. Attach the other three tubings at the 11-valve assembly (Figure 4.29-2):
  - a. Connect the tubing (with a collar) that is attached to the upper middle port (port 2) on the 5diff syringe assembly to valve 3, port 3.
  - b. Connect the tubing that is attached to the upper right port (port 3) on the 5diff syringe assembly to valve 5, port 1.
  - c. Connect the tubing that is attached to the lower left port (port 4) on the 5diff syringe assembly to valve 2, port 2.

#### Verification

- 1. Reconnect the power cord.
- 2. Turn the instrument on. An automatic startup and background check is performed.
- 3. When the startup routine and background check are done, verify the 5diff syringe is not leaking.
- 4. If no leaks are detected, close the Main card door and reattach the left side door.
- 5. Perform a system verification. See Heading 5.3, SYSTEM VERIFICATION PROCEDURES.

# 4.30 FLOW CELL COAXIAL CABLE REPLACEMENT

#### Purpose

Use this procedure to replace the flow cell coaxial cable located inside the optical bench.

#### **Tools/Supplies Needed**

- □ Allen wrench, 3 mm
- □ Small Phillips-head screwdriver
- □ Lint-free tissues
- □ DIFF flow cell coaxial cable, PN XBA399A

#### Preparation

- 1. Turn the instrument off and unplug the power cord from the instrument or the wall.
- 2. Remove the left side panel and top cover from the instrument. For details, see Heading 4.2. Set the top cover and left side panel aside.

**CAUTION** A flat cable is connected to the rear of the Main card. Be careful when you open the door (where the Main card is attached) that you do not disconnect or damage this cable.

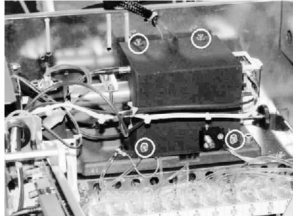
- 3. In the left side compartment,
  - a. Turn the two captive knobs counterclockwise to release the Main card door. These knobs are located to the right of the Main card.
  - b. Open the Main card door and carefully anchor the door behind the white plastic catch to keep it open. The optical bench is exposed.

#### Removal

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1. Remove the four screws securing the optical bench cover (Figure 4.30-1).

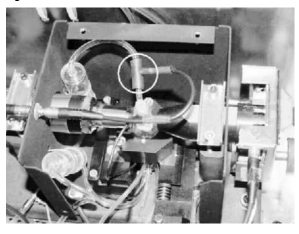
#### Figure 4.30-1 Optical Bench Cover Screw Locations



2. Locate the T-connector inside the optical bench (Figure 4.30-2).

**Note:** Take time to examine the T-connector and its attachments. Notice a cable is inserted in the base of the T-connector. The top of the T-connector links two liquid tubings - the tubing from the isolator chamber and the tubing to the flow cell.

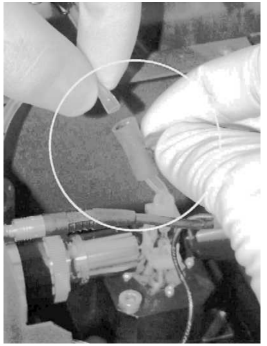
#### Figure 4.30-2 T-Connector Location



**IMPORTANT** Risk of compromising results if liquid splashes on the surface of the flow cell. Wet or dried liquid on the flow cell can affect output. Do not allow liquid to splash on the surface of the flow cell.

- 3. While holding a few lint-free tissues under the T-connector, disconnect the tubing coming from the isolator chamber (Figure 4.30-3). A few diluent drops will leak out.
- 4. Verify the sleeving is still attached to the tubing. If not, remove the sleeving from the T-connector and place it back on the tubing. This sleeve is needed to prevent leakage.

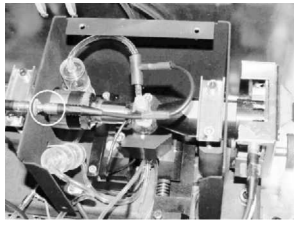
#### Figure 4.30-3 Disconnecting the Isolator Chamber from the T-connector



ATTENTION: Do not remove the tubing directly attached to the flow cell.

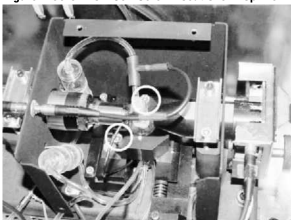
- 5. On the other side of the T-connector (at the T-connector, not the flow cell), disconnect the tubing going to the flow cell.
- 6. Verify the sleeving is still attached to the tubing. If not, remove the sleeving from the T-connector and place it back on the tubing. This sleeve is needed to prevent leakage.
- 7. Disconnect the coaxial cable from the coaxial connector on the optical bench housing.

#### Figure 4.30-4 Coaxial Cable Connector - Top View



**IMPORTANT** Risk of compromising results if the flow cell surface is smudged. Fingerprints or other smudges on the flow cell can affect output. Do not touch the surface of the flow cell. Handle the black mount only when moving the flow cell.

- 8. Disconnect the ground fitting under the flow cell as follows:
  - a. Locate and remove the two screws shown in Figure 4.30-5. Use a 3 mm Allen wrench.



#### Figure 4.30-5 Flow Cell Screw Locations - Top View

- b. Using care not to touch the surface of the flow cell, slide the flow cell forward to separate it from its mount.
- c. Turn the flow cell slowly. Make sure the plastic shims remain in the instrument.

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d. Remove the ground screw using a small Phillips-head screwdriver (Figure 4.30-6).





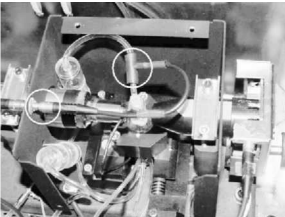
#### Replacement

- 1. Connect the ground ring on the new DIFF coaxial cable to the flow cell as follows:
  - a. Position the ground ring on the bottom of the flow cell assembly and secure with the screw removed earlier. Use a small Phillips-head screwdriver (Figure 4.30-6).
  - b. Turn the flow cell slowly and slide it inside its housing.
  - c. Install the two screws that secure the flow cell to the optical bench (Figure 4.30-5). Use a 3 mm Allen wrench.
- 2. Attach the coaxial cable to the coaxial connector on the optical bench housing (Figure 4.30-7).
- 3. Attach the new DIFF coaxial cable, at the T-connector (Figure 4.30-7):

Note: These tubings are sleeved to prevent leakage.

- a. Insert the tubing and sleeve from the isolator chamber into one leg of the T-connector.
- b. Insert the tubing and sleeve from the flow cell into the other leg of the T-connector.

#### Figure 4.30-7 Connector Location - Top View



- 4. Reconnect the power cord.
- 5. Turn the instrument on. An automatic Startup and background check is performed.
- 6. When the Startup routine and background check are done, press **ESC** to display the Main Menu.
- From the Main Menu, select 4. DIAGNOSTICS → 3. DILUTER SYSTEMS → 2. RINSE →
   2. FLOWCELL to get rid of the air bubbles stuck to the inner optical surfaces.
- 8. When the flow cell routine is done, verify there are no leaks and that the flow cell contains no (or just a very few) air bubbles.
- 9. Replace the optical bench cover (Figure 4.30-1).

#### Verification

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Go to Heading 4.4 and do a flow cell adjustment check. If an adjustment value is outside the acceptable range, do the corresponding adjustment.

**SERVICE AND REPAIR PROCEDURES** *FLOW CELL COAXIAL CABLE REPLACEMENT* 

## 4.31 OPTICAL BENCH LAMP REPLACEMENT

#### Purpose

Use this procedure to replace the optical bench lamp when the flow cell lamp fails.

#### **Tools/Supplies Needed**

- □ Allen wrenches, 3 mm, 2.5 mm, 2 mm
- □ Optical bench lamp. PN DAJ007A

#### Preparation

- 1. Turn the instrument off and unplug the power cord from the instrument or the wall.
- 2. Remove the left side panel and top cover from the instrument. For details, see Heading 4.2. Set the left side panel and top cover aside.

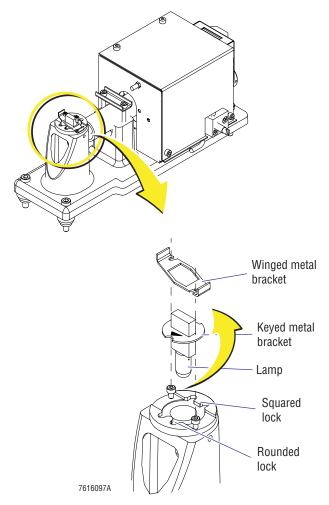
**CAUTION** A flat cable is connected to the rear of the Main card. Be careful when you open the door (where the Main card is attached) that you do not disconnect or damage this cable.

- 3. In the left side compartment,
  - a. Turn the two captive knobs counterclockwise to release the Main card door. These knobs are located to the right of the Main card.
  - b. Open the Main card door and carefully anchor the door behind the white plastic catch to keep it open. The optical bench is exposed.

**WARNING** Risk of personal injury due to hot surfaces within the instrument. Use care when working in this area. The lamp and some of the surfaces may be very hot and can burn you.

4. At the rear of the optical bench assembly (as view from the front of the instrument), locate the existing lamp and examine how it is seated. Figure 4.31-1 shows an exploded view of the assembly.





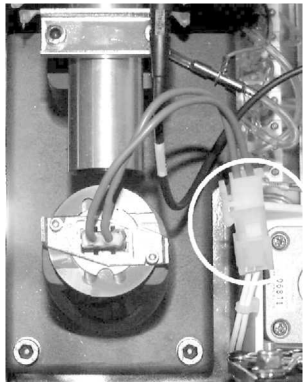
- 5. Notice the positioning of the two metal brackets (Figure 4.31-1):
  - The top metal "winged" bracket is secured underneath a screw near each end.
  - The bottom metal bracket, which is part of the lamp, is keyed to ensure proper positioning of the lamp. Note the two different notches:
    - One notch is a semicircle that matches the rounded raised area (lock) on the housing.
    - On the opposite side is a squared notch that matches a squared raised area (lock) on the housing.

#### Removal

**WARNING** Risk of personal injury due to hot surfaces within the instrument. Use care when working in this area. The lamp and some of the surfaces may be very hot and can burn you.

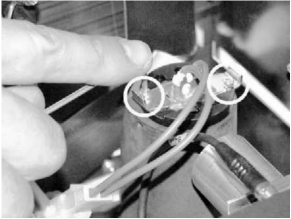
1. At the power connector, disconnect the lamp from the power supply (Figure 4.31-2).

Figure 4.31-2 Power Connector Location - Top View



2. At the top of the lamp housing, loosen the two screws a few turns (Figure 4.31-3). It is not necessary to remove the screws. Use a 2 mm hex key.

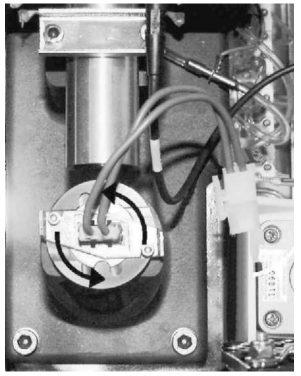




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3. Turn the winged metal bracket counterclockwise to unlock the lamp (Figure 4.31-4).





- 4. Lift the lamp and the bracket out of the housing.
- 5. Separate the winged metal bracket from the lamp and cable assembly.
- 6. Save the winged metal bracket. Discard the old lamp assembly.

#### Lamp Replacement

**IMPORTANT** Risk of compromising output of the new lamp if the surface is smudged. Fingerprints or other smudges on the lamp can affect output. Do not touch the surface of the lamp.

- 1. Using care not to touch the surface of the new lamp, place the old winged metal bracket over the new lamp.
- 2. Using the notches on the new lamp as a guide, position the new lamp assembly inside the housing. When the positioning is correct, each notch fits snugly against its raised area like a lock and key.
- 3. Turn the winged metal bracket clockwise until it is seated under the two screws.
- 4. Tighten the two screws (Figure 4.31-3).
- 5. Reconnect the lamp to the Power Supply (Figure 4.31-2).

#### Verification

- 1. Reconnect the power cord.
- 2. Close the right side door.
- 3. Turn the instrument on.
- 4. While the instrument is performing a Startup and background check, verify the new lamp is lighted.
- 5. When the Startup routine and background check are done, replace all covers and close all doors.
  - a. Gently close the Main card door. Turn the two captive knobs clockwise to secure the door.
  - b. Replace the top cover and install the five hex screws that secure it to the instrument frame. See Heading 4.2 as needed.

**ATTENTION:** When replacing the left side panel with the instrument powered on, avoid accidently turning the instrument off again by carefully positioning the opening for the power on/off rocker switch over the switch as you position the panel on the instrument frame.

- c. Replace the left side panel and install the four hex screws that secure it to the instrument frame.
- d. Close the right side door.
- 6. Perform a system verification. See Heading 5.3, SYSTEM VERIFICATION PROCEDURES

# **SERVICE AND REPAIR PROCEDURES** *OPTICAL BENCH LAMP REPLACEMENT*

# 4.32 DILUENT RESERVOIR REPLACEMENTS

#### Purpose

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Use this procedure to replace the O-ring and washer in the diluent reservoir. When performing this procedure to replace an O-ring or washer, replace both.

#### **Tools/Supplies Needed**

- □ Allen wrench, 3 mm
- □ Torque screwdriver, 2.5 mm hex-ball

Note: 120 mN.m (17.0 ozf.in) torque is required in this procedure.

□ O-ring and washer, PN - XEA286AS

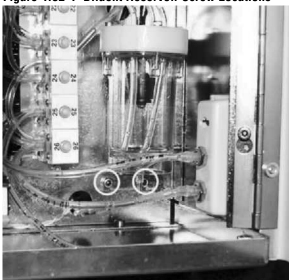
#### Preparation

- 1. If off, turn the instrument on.
- 2. From the Main Menu, select 4. DIAGNOSTICS → 3. DILUTER SYSTEMS → 3. DRAIN BATHS → 7. DILUENT RESERVOIR.
- 3. When the cycle is finished, open the right side door and verify the diluent reservoir is empty.
- 4. Turn the instrument off.

#### Removal

- 1. Locate the tubing attached to the bottom fitting under the reservoir tank. Trace this tubing to its connection point on solenoid valve 25. Remove the tubing from LV25, port 1.
- 2. Loosen the two screws under the diluent reservoir (Figure 4.32-1). It only takes a few turns to release the diluent reservoir from its support. Use a 3.0 mm Allen key.

#### Figure 4.32-1 Diluent Reservoir Screw Locations



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3. Carefully remove the reservoir and bottom bracket. Be careful that no other tubings disconnect from the reservoir.

#### **O-ring and Washer Replacement**

- 1. Remove the four screws on top of the cap using a 2.5 mm Allen wrench then remove the cap.
- 2. Remove and discard the old O-ring and washer.
- 3. Clean then dry the cap and the reservior.
- 4. Locate the replacement O-ring and washer.
- 5. Use a small amount of silicone grease between two fingers to lubricate the O-ring.
- 6. Place the new washer and O-ring securely inside the reservior cap (Figure 4.32-2).

#### Figure 4.32-2 Diluent Reservoir O-ring and Washer Positioning



- 7. Position the cap back on the reservoir body with the interior support connection turned towards the inside of the instrument and the extended tubings in the front (Figure 4.32-2).
- 8. When the cap is in position, place the four reservoir screws back in the cap then loosely tighten the four screws.

ATTENTION: Torque needed for the four hex screws (CHC M3x20) is 120 mN.m (17.0 ozf.in).

9. Tighten the four hex screws to secure the cap to the reservoir. Use a 2.5 mm hex-ball torque screwdriver to tighten the screws to 120 mN.m (17.0 ozf.in).

#### Installation

- 1. Position the diluent reservoir and bracket back in its original location.
- 2. Tighten the two hex screws that attach the reservior to its support (Figure 4.32-1).
- 3. Reattach the drain tubing under the diluent reservoir to solenoid valve 25, port 1.

#### Verification

- 1. Reconnect the power cord.
- 2. Close the right side door.
- 3. Turn the instrument on. An automatic startup and background check is performed.
- 4. When the startup routine and background check are done, verify the diluent reservoir is not leaking.
- 5. If no leaks are detected, perform a system verification. See Heading 5.3, SYSTEM VERIFICATION PROCEDURES.

# **SERVICE AND REPAIR PROCEDURES** DILUENT RESERVOIR REPLACEMENTS

## 4.33 SAMPLE SYRINGE ASSEMBLY REPLACEMENTS

#### Purpose

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Use this procedure to remove and replace the two O-rings on the injector piston of the sample syringe.

#### **Tools/Supplies Needed**

- □ Allen wrench, 3 mm
- □ Torque screwdriver, T10 torx

Note: 400 mN.m (56.8 ozf.in) torque is required in this procedure.

- □ Absorbent paper towels
- □ Replacement parts: O-ring, FAA064A (need 2)

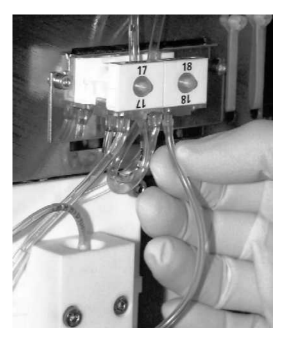
#### Preparation

- 1. If off, turn the instrument on.
- 2. From the Service menu, select 9. OTHERS → 3. PARK SYRINGES.
- 3. Turn the instrument off.
- 4. Open the right side door.

#### Removal

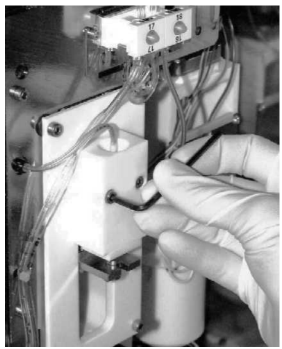
- 1. Manually push the sample probe housing towards the rear of the instrument until the sample probe is positioned over the rinse chamber.
- 2. Disconnect the tubing at solenoid valve 18, port 2 (Figure 4.33-1). The diluent drains through the sample syringe into the rinse chamber.

#### Figure 4.33-1 Disconnect Tubing at Valve 18, Port 2



3. Remove the two hex screws securing the sample syringe housing to the instrument frame (Figure 4.33-2). Use a 3 mm Allen wrench.

Figure 4.33-2 Screw Locations - Sample Syringe Housing



4. Disconnect the tubing attached to the syringe output port (Figure 4.33-3). The other end of this tubing is connected to the top port of the sample probe.

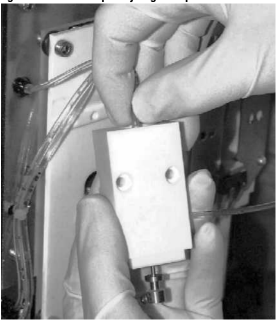


Figure 4.33-3 Sample Syringe Output Port

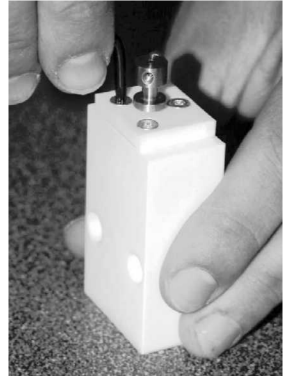
5. Remove the housing from the instrument and place it on absorbent paper towels.



## **O-ring Replacements**

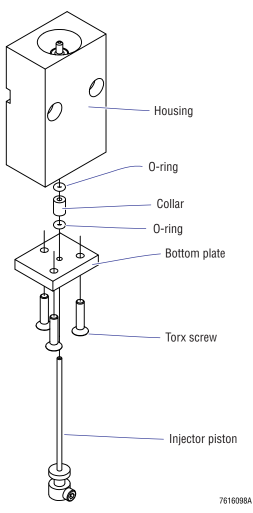
1. At the bottom plate, remove the three torx screws (FX M3x12) using a T10 Torx screwdriver (Figure 4.33-4).

Figure 4.33-4 Screw Locations on the Bottom of the Sample Syringe Housing



- 2. Separate the bottom plate (and attachments) from the sample syringe housing.
- 3. Clean then dry the outside of the sample syringe.
- 4. Replace the O-rings on the injector piston (Figure 4.33-5):
  - a. Remove the two O-rings and the collar from the injector piston.
  - b. Discard the old O-rings. Save the collar.
  - c. Clean the outside of the injector piston and the bottom plate.
  - d. Locate the two replacement O-rings.
  - e. Use a small amount of silicone grease between two fingers to lubricate an O-ring then place it on the injector piston.
  - f. Place the collar on the injector piston.
  - g. Use a small amount of silicone grease between two fingers to lubricate another O-ring then place it on the injector piston.





- 5. Once the replacements are complete, replace the bottom plate as follows:
  - a. Carefully replace the bottom plate.
  - b. Loosely install the three torx screws (Figure 4.33-4).

ATTENTION: Torque needed for the three torx screws is 400 mN.m (56.8 ozf.in).

c. Tighten the three torx screws. Use a torx driver with a T10 torx bit to tighten the screws to 400 mN.m (56.8 ozf.in).

#### Installation

- 1. Position the sample syringe back inside the instrument.
- 2. Replace the two hex screws that secure the sample syringe to the instrument (Figure 4.33-2).
- 3. Reattach the tubing connected at the top of the sample probe to the output port on the top of the syringe housing.
- 4. Reattach the tubing connected at the side of the sample syringe to solenoid valve 18, port 2.

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#### Verification

- 1. Reconnect the power cord.
- 2. Close the right side door.
- 3. Turn the instrument on. An automatic startup and background check is performed.
- 4. When the startup routine and background check are done, verify the sample syringe is not leaking.
- 5. If no leaks are detected, perform a system verification. See Heading 5.3, SYSTEM VERIFICATION PROCEDURES.

**SERVICE AND REPAIR PROCEDURES** SAMPLE SYRINGE ASSEMBLY REPLACEMENTS

## 4.34 DRAINING BATH REPLACEMENTS

#### Purpose

Use this procedure to replace the O-rings on the draining baths.

#### **Tools/Supplies Needed**

□ O-ring, FAA066A (one for each bath)

#### Preparation

- 1. If off, turn the instrument on.
- 2. From the Main Menu, select 4. DIAGNOSTICS → DILUTER SYSTEMS → 3. DRAIN BATHS → 6. ALL BATHS.
- 3. Open the right side door.
- 4. Verify all the draining baths are empty.
- 5. Turn the instrument off.

#### **O-ring Replacements**

1. Locate the replacement O-rings.

**WARNING** Risk of personal contamination. If you do not properly shield yourself while servicing the drains baths, you may become contaminated. To prevent possible biological contamination, you must wear appropriate safety glasses, a lab coat, and gloves when servicing the drain baths.

**ATTENTION:** When removing a draining bath, the diffuser (which resembles a three-leg stool) may pop out of the bath. If this happens, the diffuser must be retrieved, rinsed with distilled water, and placed back inside the draining bath.

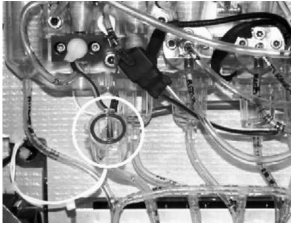
2. Grasp the bottom of the drain bath between two fingers and gently turn the bath to loosen then remove it from its support (Figure 4.34-1).

#### Figure 4.34-1 Removing a Draining Bath



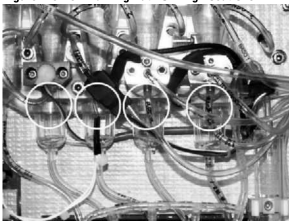
- 3. Remove and discard the old O-ring (Figure 4.34-2).
- 4. Clean then dry the top of the bath.
- 5. Make sure the diffuser is seated properly inside the draining bath. The circular part of the diffuser must rest on the bottom of the draining bath with the tabs extended upward. (Proper placement of diffuser resembles a three-legged stool turned upside down.)
- 6. Replace the O-ring (Figure 4.34-2)

#### Figure 4.34-2 Draining Bath O-ring Placement



- 7. Position the draining bath back on its support then gently rotate the bath to tighten it.
- 8. Repeat steps 2 through 7 at each draining bath (Figure 4.34-3).

Figure 4.34-3 Draining Bath O-Ring Locations



#### Verification

- 1. Close the right side door.
- 2. Turn the instrument on. An automatic startup and background check is performed.
- 3. When the startup routine and background check are done, verify the draining baths are not leaking.
- 4. If no leaks are detected, perform a system verification. See Heading 5.3, SYSTEM VERIFICATION PROCEDURES.

## 4.35 O-RING REPLACEMENTS IN THE COUNTING BATHS (RBC and WBC/BASO Baths)

#### Purpose

Use this procedure to replace the coaxial cable O-ring and the two counting head aperture O-rings. This procedure instructs you to bypass the right side door interlock which allows you to observe instrument operation with the right side door open. Avoid any unnecessary contact with the sample probe. The probe is sharp and may contain biohazardous materials. When the procedure is complete, make sure the interlock is reactivated to ensure the customer is not injured by the sample probe or its movement.

#### **Tools/Supplies Needed**

- □ Allen wrenches, 3 mm, 2.5 mm, 2 mm
- Torque screwdriver

Note: 100 mN.m (14.2 Ozf.in) torque is required in this procedure.

- □ Hemostat or tweezers with a pointed end
- □ Small side cutters
- □ Protective cover for the bath electrode

**Note:** If you do not already have this special tool, you will need a micropipette tip and a scalpel or sharp knife.

- □ White paper towel or cloth
- □ Cotton-tip applicator stick
- □ High quality, fragrance-free bleach (10-12% sodium hypochlorite available chloride)
- Distilled water
- □ Replacement parts
  - Coaxial cable O-ring, PN FAA046A (need 1 for each bath)
  - Aperture O-ring, PN GBG156A (need 2 for each bath)

#### Preparation

- 1. Locate the replacement coaxial cable O-rings (one needed for each bath) and the aperture O-rings (two needed for each bath).
- 2. If off, turn the instrument on.
- 3. From the Service menu, bypass the right side door interlock:
  - a. Select 9. OTHERS → 1. USER MODE.
  - b. Press the down arrow key to highlight the *SERVICE* option.

Note: This option bypasses the right side door interlock allowing you to observe instrument operation with the right side door open.

- c. Press the decimal point key then **ESC** to select the Service option.
- d. Press **ESC** two more times to return to the Diagnostics menu.
- 4. From the Diagnostics menu, select **3**. **DILUTER SYSTEMS >> 3**. **DRAIN BATHS >> 6**. **ALL BATHS**.
- 5. Open the right side door and verify all the baths are empty.
- 6. Turn the instrument off and disconnect the power cord.

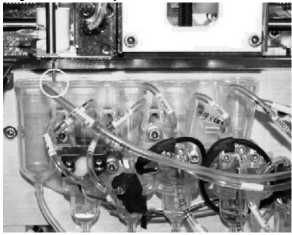
7. Remove the left side panel and set it aside.

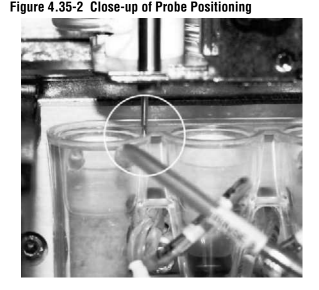
#### Removal

**WARNING** Risk of personal contamination. If you do not properly shield yourself while servicing the bath assembly, you may become contaminated. To prevent possible biological contamination, you must wear appropriate safety glasses, a lab coat, and gloves when servicing the bath assembly.

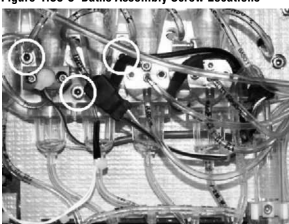
- 1. Before removing the baths assembly, position the sample probe to provide a rough alignment guide when the baths assembly is replaced at the end of this procedure.
  - a. Manually push the sample probe housing over the rinse bath.
  - b. Gently push down on the top of the sample probe until the tip of the probe rests on the inside rim of the rinse bath (Figures 4.35-1 and 4.35-2).

Figure 4.35-1 Sample Probe Position at the Rinse Bath





- c. Without moving the probe position, manually push the sample probe as far forward as possible to the front of the instrument. At the end of this procedure, the positioning of this probe will provide a guide for aligning the baths assembly.
- 2. Place a piece of white paper or cloth under the baths assembly.
- 3. Record the tubing positions before dismantling the baths assembly.
- 4. Disconnect all bath tubings except the waste tubings.
- 5. Remove the three screws securing the baths assembly to the instrument panel (Figure 4.35-3). Use a 3 mm Allen wrench.



#### Figure 4.35-3 Baths Assembly Screw Locations

#### **O-ring Replacements**

#### **Replacing the Coaxial Cable O-ring on the Bath Electrode**

1. Loosen the two hex screws on the back of the RBC bath (Figure 4.35-4). Use a 2 mm Allen wrench.

#### Figure 4.35-4 Electrode Screw Locations



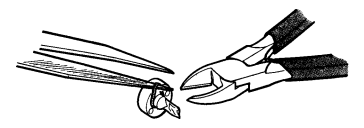
2. Carefully remove the electrode from the bath.

**CAUTION** Risk of damage to the bath electrode. When removing the coaxial cable O-ring, you can accidently cut the bath electrode. Make sure the O-ring is sufficiently clear of the electrode before cutting the O-ring.

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- 3. Remove the coaxial cable O-ring from the bath electrode as follows (Figure 4.35-5):
  - a. Use a small, pointed-tip tweezers to grasp the O-ring (Figure 4.35-5).
  - b. While gently lifting the O-ring to separate it from the electrode, carefully cut the O-ring with a pair of small side cutters (Figure 4.35-5). Do not clip the electrode.

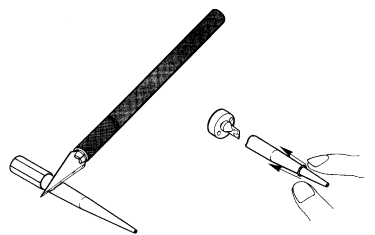
Figure 4.35-5 Removing the Coaxial Cable O-ring from the Bath Electrode



**CAUTION** Risk of damage to the bath electrode. When replacing the coaxial cable O-ring, you can accidently break off the bath electrode. Use a plastic micropipette tip (cut to fit over the electrode) to protect the bath electrode as you seat the O-ring.

- 4. If you have not already done so, use a plastic micropipette tip to make a protective cover for the bath electrode (Figure 4.35-6).
  - Use a scalpel or other sharp knife to cut the plastic micropipette tip.
  - Shorten the tip until it fits securely over the bath electrode.

#### Figure 4.35-6 Making a Protective Cover from a Micropipette Tip



- 5. Position the new O-ring over the bath electrode as follows:
  - a. Place the coaxial cable O-ring on the protective cover (Figure 4.35-6).
  - b. Carefully position the protective cover over the bath electrode.
  - c. Push the O-ring from the plastic cover onto the bath electrode.
  - d. .Carefully remove the plastic cover from the electrode. Make sure the O-ring remains on the bath electrode.
  - e. Verify the O-ring is properly seated.

6. Set the bath electrode aside. While the bath is disassembled, replace the aperture O-rings in the counting head.

#### **Replacing the Aperture O-rings in the Counting Head**

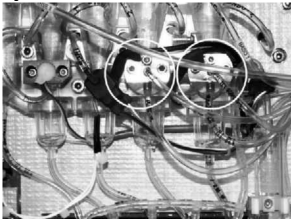
**IMPORTANT** Risk of misleading results. The 50  $\mu$ m RBC/PLT aperture and the 80  $\mu$ m WBC/BASO aperture are not interchangeable. Disassemble the RBC/PLT counting head, replace the aperture O-ring, and reassemble the counting head <u>before</u> disassembling the WBC/BASO counting head.

**ATTENTION:** Be careful when handling an aperture, it is very small so it is easy to drop and easy to lose. It is recommended that you work over a white surface so the aperture can be easily seen if it is dropped.

- 1. If you have not already done so, place a piece of white paper or cloth under the baths assembly.
- 2. Carefully dismantle the RBC/PLT counting head (Figure 4.35-7). Use a 2.5 mm Allen key.

Note: In Figure 4.35-7, the RBC/PLT counting head is on the left and the WBC/BASO counting head is on the right.

#### Figure 4.35-7 Bath Electrode Locations



3. When the counting head is separated from the bath, one of the O-rings may remain inside the counting head. If this happens, remove and discard the O-ring and clean the counting head.

O-RING REPLACEMENTS IN THE COUNTING BATHS (RBC and WBC/BASO Baths)

4. Figure 4.35-8 shows the aperture and O-ring location on the bath when the counting head is removed.



Figure 4.35-8 Location of the Aperture and Its Two O-Rings

**CAUTION** Risk of damage to the bath or aperture. Using a sharp instrument inside the bath may damage the inside of the bath and the aperture. Never use a sharp instrument inside the bath.

- 5. Remove the aperture and the O-ring from the bath as follows:
  - a. Locate a cotton-tip applicator stick.
  - b. Hold the bath close to the surface of the white paper towel.
  - c. Insert the cotton tip of the applicator stick through the opening created when the bath electrode was removed (at the rear of the baths assembly).
  - d. Gently push the aperture and O-ring onto the white paper surface.
- 6. Place the aperture in a small amount of distilled water to keep it moist and discard the old O-ring(s).
- 7. Clean the bath and counting head as follows:
  - a. Pour diluted bleach on a soft, clean paper towel.
  - b. Clean the bath and the counting head using the wet paper towel.
  - c. Rinse the surfaces thoroughly with distilled water.
  - d. Dry the counting head and the exterior of the bath with a soft paper towel.

8. Install the new O-rings and replace old aperture as follows:

**Note:** When completed, the aperture should be sandwiched between the two aperture O-rings.

- a. Moisten the tip of your finger with distilled water then pick up one of the aperture O-rings using your moistened finger.
- b. Pass your finger (with the clinging O-ring) across the bath opening to seat the O-ring in the groove.

**CAUTION** Risk of damage to the aperture. Do not handle the aperture using a hard instrument or tool. Handle the aperture with your fingers. Clean the aperture gently between your two fingers.

- c. Pick up the aperture with your finger tip then gently rub it between two fingers to clean it. Rinse the aperture with distilled water.
- d. Position the aperture on the tip of your moistened finger.
- e. Pass your finger (with the clinging aperture) across the bath opening to seat the aperture on the aperture O-ring already seated in the bath opening.
- f. Moisten the second aperture O-ring and seat it inside the counting head.

**CAUTION** Risk of damage to the aperture. If too much pressure is applied as the bath components are being reassembled, the aperture may break. To avoid applying too much pressure on the bath (and the aperture), reconnect the counting head tubing <u>before</u> attaching the counting head back on the bath.

- 9. Reconnect the counting head tubings.
- 10. Carefully position the counting head back on the bath. Make sure the O-rings and aperture are properly seated and do not become crimped as you position the counting head on the bath.
- 11. Loosely install the two hex screws in the counting head.

**CAUTION** Risk of damage to the aperture. Overtightening the counting head hex screws can break the aperture. Tighten these screws to a torque of 100mN.m (14.2 ozf.in) using a torque screwdriver with a 2.5 mm bit.

**ATTENTION:** Torque needed for the RBC (and WBC/BASO) bath hex screws is 100 mN.m (14.2 ozf.in).

- 12. Tighten the two counting head hex screws. Use a 2.5 mm torque screwdriver to tighten each screw to 100 nN.m (14.2 ozf.in).
- 13. Install the coaxial cable electrode that was set aside earlier:
  - a. Carefully insert the bath electrode inside the opening at the rear of the bath.
  - b. Loosely install the two hex screws.
  - c. Tighten the two hex screws. Use a 2 mm torque screwdriver to tighten each screw to 100 nN.m (14.2 ozf.in).
- 14. Repeat steps 2 through 13 on the WBC/BASO counting head. See Figure 4.35-7. (The WBC/BASO counting head is the one on the right.)

## Installation

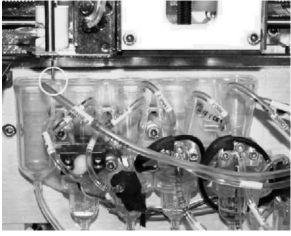
- 1. Using your recorded tubing positions, reattach the bath tubings.
- 2. Being careful to not crimp any tubings, position the baths assembly against its support panel.
- 3. Replace and loosely tighten the three screws removed earlier (Figure 4.35-3).
- 4. Make sure any liquid inside the bath compartment is wiped dry, especially any liquid on the solenoid valves.

## Align the Bath Assembly

**CAUTION** Risk of damage to the sample probe. If power is restored to the instrument and the baths assembly is mounted too high, the sample probe may become bent when it hits the bath edges as it moves from bath to bath. Do not restore instrument power until the baths assembly is aligned.

- 1. Manually push the sample probe housing to the outside edge of the rinse bath. Move the baths assembly up or down as necessary until the tip rests gently on the outside rim of the rinse bath.
- 2. While continuing to push the sample probe housing over the rinse bath to the inside edge of the bath, adjust the bath assembly up or down as needed so that the sample probe tip comes to rest gently on the inside rim of the rinse bath as shown in (Figures 4.35-9 and 4.35-10).

Figure 4.35-9 Sample Probe Position at the Rinse Bath

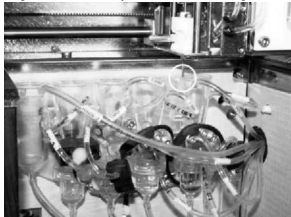


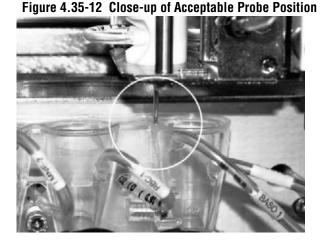


- 3. Gently tighten the center and left screws.
- 4. Without lifting the probe, gently push the probe housing towards the inside edge of the last bath, the WBC/BASO bath. Reposition the baths assembly as needed to clear the edge of each bath.

5. Verify the sample probe is positioned at the WBC/BASO bath as shown in Figures 4.35-11 and 4.35-12.

Figure 4.35-11 Sample Probe Position, Right Side





- 6. Gently tighten the right screw on the baths assembly.
- 7. Make sure all three screws are tight so that the baths assembly is secure on the instrument's frame.
- 8. Carefully move the sample probe housing over the outer edge of the rinse bath. The probe should equally clear all baths. If not, loosen the nearest screw and reposition the baths assembly until the distance between the tip of the probe and the top of each bath is the same.

#### Verification

- 1. Reconnect the power cord.
- 2. Turn the instrument on.

**WARNING** Risk of personal injury. The right side door interlock is bypassed which allows you to observe instrument operation with the right side door open. Avoid any unnecessary contact with the sample probe. The probe is sharp and may contain biohazardous materials. When the procedure is complete, make sure the interlock is reactivated to ensure the Customer is not accidently injured by the sample probe or its movement.

- 3. On the instrument keypad, press the **STARTUP** button to initiate the startup cycles and background check.
- 4. As the startup routine and background check are being done,
  - a. Observe operation inside the right side compartment to verify there are no leaks. Look at the entire compartment not just the baths assembly. If a tubing is connected to a wrong port, the liquid in that tubing will be sent to an area that is not meant to receive it.
  - b. Check for leaks inside the left compartment.

- 5. From the Service menu, select **9. OTHERS → 1. USER MODE** and reconnect the right side door interlock.
  - a. Press the up arrow key to highlight the *MANUAL* option.
  - b. Press the decimal point key then **ESC** to select the Manual option.
  - c. Press **ESC** again to return to the Service menu.

**CAUTION** It is mandatory to perform the probe adjustment procedure after this adjustment.

- 6. Check the sample probe position inside each bath.
  - Under Heading 4.6, SAMPLE PROBE CHECKS AND ADJUSTMENTS, do the Inside Bath Position Check.
  - Perform the adjustment if needed.
- 7. When the probe position inside each bath is correct, perform a system verification. See Heading 5.3, SYSTEM VERIFICATION PROCEDURES.

## 4.36 OPTICAL BENCH PRELIMINARY ADJUSTMENTS

#### Purpose

Use this procedure to make preliminary adjustments to the flow cell and optics in the optical bench assembly. These adjustments may be necessary when the:

- Optical bench assembly is replaced.
- The flow cell is replaced.
- Transfer Time value is outside the acceptable range.
- Absorbance Channel value is outside the acceptable range.
- Optical lamp is replaced.

#### **Tools/Supplies Needed**

□ Allen wrenches, 3 mm, 2 mm

#### Preparation

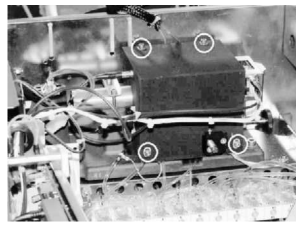
I

- 1. Turn the instrument off and unplug the power cord from the instrument or the wall.
- 2. Remove the left side panel and the top cover from the instrument. For details, see Heading 4.2. Set the left side panel and top cover aside.

**CAUTION** A flat cable is connected to the rear of the Main card. Be careful when you open the door (where the Main card is attached) that you do not disconnect or damage this cable.

- 3. In the left side compartment,
  - a. Turn the two captive knobs counterclockwise to release the Main card door. These knobs are located to the right of the Main card.
  - b. Open the Main card door and carefully anchor the door behind the white plastic catch to keep it open. The optical bench is exposed.
- 4. At the optical bench,
  - a. Remove the four hex screws shown in Figure 4.36-1.
  - b. Carefully remove the optical bench cover and set it aside.

#### Figure 4.36-1 Optical Bench Cover Screw Locations



#### Verify the Flow Cell is Free of Bubbles

- 1. Turn the instrument on. An automatic startup and background check is performed.
- From the Main Menu, select 4. DIAGNOSTICS → 3. DILUTER SYSTEM → 2. RINSE →
   2. FLOWCELL to remove air bubbles clinging to the inner optical surfaces.
- 3. When the routine is complete, verify the flow cell is bubble-free. (An occasional air bubble is acceptable.)
- 4. Press **ESC** as many times as necessary to return to the Main Menu.

#### **Course Adjustments**

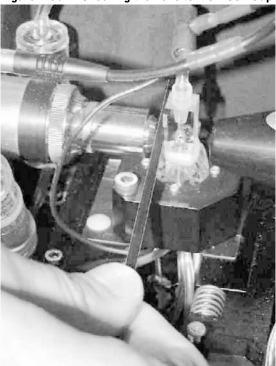
**ATTENTION:** Course adjustment of the flow cell positions the lens and flow cell system for final alignment with the Absorbance Channel adjustment. **Course adjustments may not be necessary.** If the Absorbance Channel value and the Transfer Time value are acceptable, course adjustments are not needed.

#### Y-Axis Adjustment

**CAUTION** Never dismantle the lens assembly or even unlock it. The adjustment is factory made and cannot be performed in the field.

**CAUTION** If you use a 3 mm Allen wrench to check the gap, be careful that you do not damage or scratch the flow cell or lens assembly.

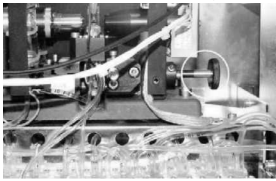
Use a 3 mm shim or Allen wrench to verify the gap between the lens assembly and the flow cell is approximately 3 mm (Figure 4.36-2). The final adjustment will be close to this position.



#### Figure 4.36-2 Checking the Lens to Flow Cell Gap

- If the gap is approximately 3 mm, go to the X-Axis Adjustment heading.
- If the gap is not close to 3 mm, use the front knob (Figure 4.36-3) to move the flow cell forward or backwards (along the Y-axis) as needed.

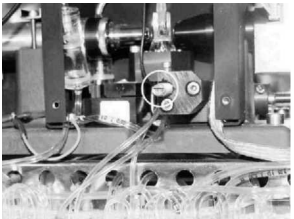
#### Figure 4.36-3 Front Knob for Y-Axis Adjustment



#### **X-Axis Adjustment**

The lens assembly is fixed. During this course adjustment, the flow cell is repositioned as needed along its X-axis so that the focal point of the light beam coming from the lens assembly is properly positioned inside the flow cell.

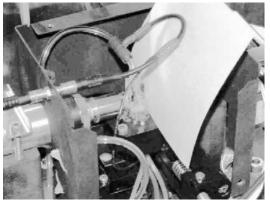
- 1. From the rear of the instrument, look down the lens assembly and evaluate the position of the flow cell in relation to the lens assembly. The flow cell must be centered with the lens.
- 2. If the flow cell is located to the left or to the right of the lens, use the side screw (Figure 4.36-4) to move the flow cell left or right (along the X-axis) so that the flow cell is roughly centered with the lens.



#### Figure 4.36-4 Side Screw - X-Axis Adjustment

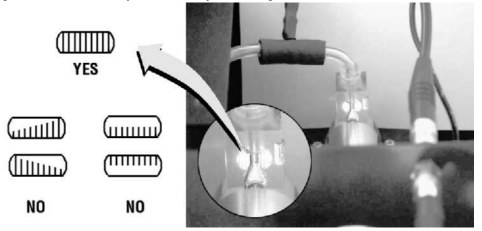
3. Place a piece of white paper between the flow cell and the reception diode as shown in Figure 4.36-5.

Figure 4.36-5 How to Position the White Paper



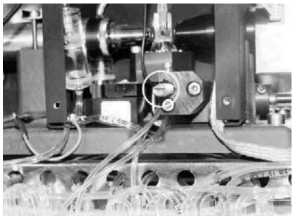
- 4. From the rear of the instrument, look for a light spot on the white paper.
  - If the lamp filament image on the white paper matches the ideal image seen in Figure 4.36-6, go to step 5.
  - If the image does not match the ideal projection seen in Figure 4.36-6, go to the Lamp Alignment heading that follows before proceeding.

Figure 4.36-6 Ideal Lamp Filament Projection Image



5. Use the side screw (Figure 4.36-7) to position the flow cell in the center of the light spot.

Figure 4.36-7 Side Screw - X-Axis Adjustment



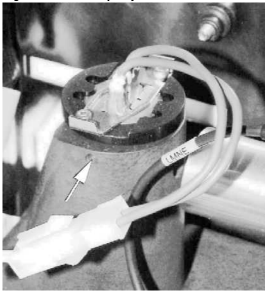
6. Go to Heading 4.4, FLOW CELL CHECKS AND ADJUSTMENTS to ensure the diff adjustment values are within acceptable limits.

#### Lamp Alignment

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**CAUTION** A small 2 mm setscrew in the lamp housing allows adjustment of the optics lamp. This adjustment is factory made and should not be changed unless absolutely neceessary.

The optics lamp can be aligned using a 2 mm setscrew in the lamp housing (Figure 4.36-8). When this setscrew is loosened, the entire lamp, with its base, can be turned or moved up and down. Make this adjustment only if the previous flow cell adjustments did not produce a proper filament projection.

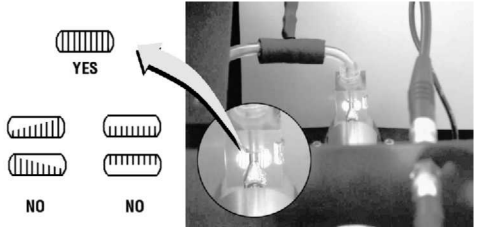


#### Figure 4.36-8 Lamp Adjustment Screw

1. Use a 2 mm Allen wrench to loosen the setscrew holding the lamp base in place (Figure 4.36-8). Do not remove the setscrew.

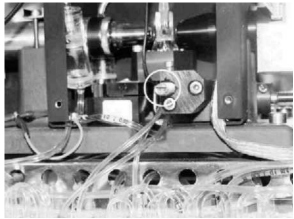
2. While rotating the entire lamp with its base, move the lamp up or down as needed until the lamp filament projection image matches the ideal projection image seen in Figure 4.36-9.

#### Figure 4.36-9 Ideal Lamp Filament Projection Image



- 3. When the light spot matches the image seen in Figure 4.36-9, tighten the setscrew.
- 4. Use the side screw (Figure 4.36-10) to position the flow cell in the center of the light spot.

#### Figure 4.36-10 Side Screw - X-Axis Adjustment



5. Go to Heading 4.4, FLOW CELL CHECKS AND ADJUSTMENTS to ensure the diff adjustment values are within acceptable limits.

## 4.37 FLOW CELL WBC BALANCE

#### Purpose

Use this procedure to balance the WBC count generated in the flow cell with the WBC count generated and reported from the WBC/Baso bath. A comparison of these two counts is made to ensure that there are no problems with either system. When they are too far apart, the diff+/diff- flags are generated. These adjustments may be necessary when:

- the optical bench assembly is replaced.
- the flow cell is replaced.
- the flow cell aperture gain is adjusted.
- too many diff+ or diff- flags.
- WBC aperture and calibration have changed.

#### **Tools/Supplies Needed**

□ Fresh blood samples

#### Procedure

1. From the Main Menu, select

#### 4. Diagnostics → 5. Service → 8. Flowcell WBC Calibration → 1. Autocalibration

The WBC/Flow Cell Balance screen is displayed (Figure 4.37-1).



| WBC     FLOWCELL     WBC       RUN AT LEAST 5 SAMPLES<br>AND PRESS ENTER TO CALIBRATE       STAT     WBC     FLOWCELL     WBC       CURR     1.00     135       NEW         MEAN | A | UTO      | CALIE   | BRATION |       | 02/2 | 7/00   16: | 05 |
|--|---|----------|---------|---------|-------|------|------------|----|
| AND PRESS ENTER TO CALIBRATE STAT WBC FLOWCELL WBC CURR 1.00 135 NEW   |   |          |         | WBC     | FLOWC | ELL  | WBC        | ļ  |
| CURR 1.00 135<br>NEW   |   |          |         |         |       |      | ATE        |    |
|  |   | CU<br>NE | RR<br>W |         | 135   | ELL  | WBC        |    |

- 2. Run a fresh normal blood sample at least 5 times. A mean will be established after 3 runs and a new flow cell WBC count calibration factor will be calculated. The raw flow cell WBC count is multiplied by this factor before being matched with the printed WBC count from the WBC bath.
- 3. Press **ENTER** when the runs are complete. You will then be given the option of accepting (**ENTER**) or rejecting (**ESC**) the new factor. There should be no flagging for the WBC results. If there is, do not accept the new factor. Repeat the procedure, possibly with a different fresh normal blood.

### **SERVICE AND REPAIR PROCEDURES** *FLOW CELL WBC BALANCE*

## 4.38 SETTING diff+/diff- THRESHOLDS

#### Purpose

Use this procedure to check or set the sensitivity for the diff+ and diff- flags. These flags indicate a difference between the WBC count arrived at in the flow cell and the WBC count derived in the bath.

#### **Tools/Supplies Needed**

□ None

#### Procedure

1. From the Main Menu, select

#### 4. Diagnostics → 5. Service → 8. FlowCell WBC Calibration → 2. Cal Factors

The diff Flag Sensitivity screen is displayed (Figure 4.38-1):

#### Figure 4.38-1 diff Flag Sensitivity Screen

| CAL FACTORS  | 02 / 27 / 00   16:05 |
|--|----------------------|
| %WBC BAL. 1 50.0<br>%WBC BAL. 2 20.0<br>%WBC BAL. 3 15.0 |                      |
| FLOW CELL WBC 0.95                                       |                      |
| WBC FLOW CELL BALANCE                                    |                      |
|  |                      |
|  | 7616104E             |

2. WBC FLOWCELL BALANCE LIMIT (%) XX.X lines set the sensitivity threshold for the diff+/diff- flag, as a percentage. The default values are 50%, 20%, and 15% for BAL 1, BAL 2, and BAL 3 respectively.

For example, for WBC BAL 2, the diff+/diff- flag will be triggered when enabled, if the (calibration factor corrected)WBC count from the flow cell is more than 20% different from the (calibration factor corrected) WBC BAL 2 count at the WBC/Baso bath.

The *FLOWCELL WBC XXX* line indicates the flow cell WBC calibration factor that was determined by Heading 4.37, FLOW CELL WBC BALANCE. The flow cell WBC calibration factor can be altered or manually entered on this screen.

The last line, *WBC FLOWCELL BALANCE* allows the diff+/diff- flagging to be disabled by clearing the enable/disable button.

# **SERVICE AND REPAIR PROCEDURES** SETTING diff+/diff- THRESHOLDS

## 4.39 MAIN CARD REPLACEMENT AND SOFTWARE TRANSFER

#### Purpose

Use this procedure to replace the Main Card and transfer the software EPROMs to the new card. The new card is then checked to ensure the settings are correct.

If a software upgrade is also being performed, refer to the appropriate service memo.

#### **Tools/Supplies Needed**

- □ Allen wrench
- □ IC puller, PN 5450537
- □ External digital pressure/vacuum gauge (referred to as vacuum meter)
- □ Voltmeter

#### **Preliminary Setup**

- Set the instrument to the Manual Startup mode.
   From the Main Menu, select
   4. Diagnostics >> 5. Service >> 9. Others >> 1. User Mode then Manual Startup.
- 2. Print out the system settings before removing the Main card.
  - From the Main Menu, select

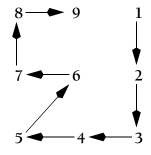
5. Setup № 6. Others № 8. Print System Setup.

Note: After the new card is installed, the new settings are compared to these settings and adjustments made as necessary.

#### **Removing Main Card**

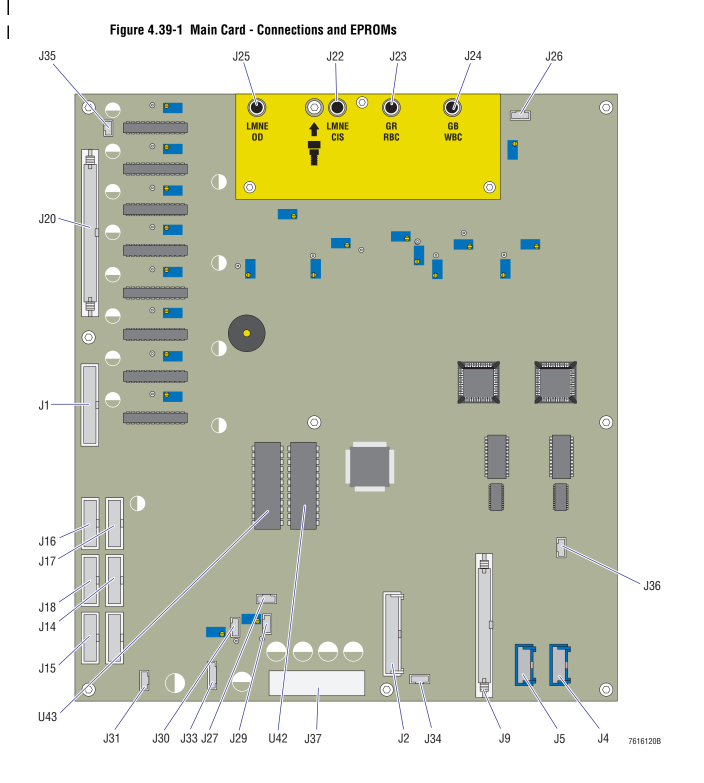
- 1. Turn off the instrument and unplug the power cord from the wall outlet.
- 2. Remove the left side panel (Heading 4.2).
- 3. Starting at the top left corner of the Main card (Figure 4.39-1), remove the cable plugs from the connections shown in Table 4.39-1.
- 4. Using a 3-mm Allen wrench, remove the nine screws from the Main card in accordance with the sequence shown below.

**ATTENTION:** At last position (9), do not allow the screw to drop into shield. Rather, hold the wrench against the screw while removing the card. Turn the card down to allow the screw to drop through the hole in shield.



**SERVICE AND REPAIR PROCEDURES** MAIN CARD REPLACEMENT AND SOFTWARE TRANSFER

Remove the Main card. 5.



| Cable Plug  | Color                 | Number of Wires<br>(type) | Label (on board)                                 |
|---|-----------------------|---------------------------|--|
| P35   | red, blk              | 2                         | J35 (3-pin connector)                            |
| LMNE OD (P25)<br>LMNE CIS (P22)<br>GR RBC (P23)<br>GB WBC (P24) | blk                   | (coax)                    | LMNE OD<br>LMNE CIS<br>GR RBC<br>GB WBC          |
| P26   | wht, blu, yel, brn    | 4                         | J26  |
| P36   | wht                   | 4                         | J36  |
| P4  | gry                   | (10-wire ribbon)          | J4   |
| P5  | wht                   | 4                         | J5 (10-pin connector)                            |
| P9  | gry                   | (ribbon)                  | J9 (40-pin connector)                            |
| P34   | wht                   | 2                         | J34 (4-pin connector)                            |
| P2  | gry                   | (ribbon)                  | J2 (26-pin connector)                            |
| P37* (pwr supply)   | red                   | 12                        | J37  |
| P27   | wht                   | 2                         | J27 (4-pin connector)                            |
| P29   | red, grn, blk, wht    | 4                         | J29  |
| P30   | red, grn, blk, wht    | 4                         | J30  |
| P33†  | red (3), wht (2), blk | 6                         | J33  |
| P31†  | red (2), wht (2)      | 4                         | J31  |
| P15, P14<br>P18, P17, P16                                       | gry                   | (ribbon)                  | J15, J14<br>J18, J17, J16<br>(14-pin connectors) |
| P1  | blk                   | ribbon                    | J1 (26-pin connector)                            |
| P20   | gry                   | ribbon                    | J20 (52-pin connector)                           |

\* P37 plug holes are offset to the top. DO NOT use excessive force when installing plug. † Plug label shows thermistor values in ohms.

#### **Transferring EPROMs to Replacement Card**

Note: Software EPROMs are not provided with the replacement card.

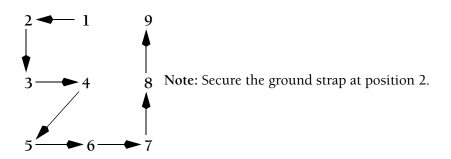
**CAUTION** DO NOT pry up EPROMs with screwdriver. Prying up the EPROMs can damage the lands on the card. Remove the EPROMs with an IC puller.

- 1. Using the IC puller, remove the EPROMs U42 (right) and U43 (left) from the old Main card. See Figure 4.39-1 for location. Set the puller over the EPROM and pull straight out.
- 2. Check the EPROM labels for proper identification, software version and addresses (U42 is labeled "odd" and U43 labeled "even").
- 3. Insert U42 (right) and U43 (left) into their plug-ins on the card, NOTCH UP.

### **Installing Main Card**

1. Using a 3-mm Allen wrench, install the nine screws to secure the Main card to the instrument in accordance with the sequence shown below.

**ATTENTION:** At position 1, start threading the screw through the shield into the card first, then hold the wrench against the screw and mount the card on the instrument. Complete threading the screw but leave it loose for aligning other screws.



2. Install all connections listed in Table 4.39-1.

**CAUTION** If, following the Beckman Coulter logo the instrument begins a **rinse cycle**, immediately TURN OFF the instrument. Damage can occur to the probe if the instrument was not set for Manual Startup mode.

- 3. Connect power cord to ac wall outlet and turn on the instrument. The *Beckman Coulter* logo, along with the revision level of the software (x.xx) is displayed on the screen in approximately 7 seconds. Continue to check the instrument settings as outlined in "Verifying System Configuration" and "Verifying Main Card Settings" below.
  - a. If only a single line appears on the screen, turn off the instrument and check the positions of the EPROMs (U43 and U42); they may be reversed.
  - b. If the instrument begins a **rinse cycle**:
    - 1) Turn off the instrument.
    - Open the right side door and turn on the instrument. One or both of the folowing messages is displayed:
       SYSTEM ERROR RUN SYSTEM RESET CYCLE
       BATH ENCLOSURE DOOR OPEN
    - 3) Press <sup>ESC</sup> to access the Main Menu and select
      4. Diagnostics → 5. Service → 9. Others → 1. User Mode then Manual Startup.
- 4. Power Off, close the right side door and power On again. The instrument should not attempt to do a rinse cycle.

#### Verifying System Configuration

:

Use this procedure when replacing a Main card and the configuration of the replacement card is unknown.

1. Reset values to default. From the Main Menu, select

#### 4. Diagnostics >> 5. Service >> 9. Others >> 4. Setting to Default Values.

The instrument prints the current values and then resets to the defaults. Enter the settings as listed in Table 4.39-2 and compare them to the original system setup (printed in step 2 of "Preliminary Setup").

| Parameter                    | From Main Menu  |
|------------------------------|---|
| Date Format                  | 5. Setup ↦ 1. Date/Time ↦ 2. Date Format                                    |
| Time Format                  | 5. Setup ↦ 1. Date/Time ↦ 1.Time Format                                     |
| Units                        | 5. Setup ↦ 2. Units   |
| Language                     | 5. Setup ↦ 6. Others ↦ 5.Language.  |
|                              | If American English, set "USA" to selected or (enable).                     |
| Manual Startup               | 4. Diagnostics ↦ 5. Service ↦ 9. Others ↦ 1. User Mode then Manual Startup  |
| Service                      | 4. Diagnostics ↦ 5. Service ↦ 9. Others ↦ 1. User Mode                      |
| Identification Mode          | 5. Setup № 6. Others № 2. Identification Mode                               |
| Bar Code with Checksum       | 5. Setup ↦ 6. Others ↦ 2. Identification Mode                               |
| Display DIFF #               | 5. Setup ↦ 6. Others ↦ 2. Identification Mode                               |
| Enable ATL, IMM, PCT,<br>PDW | 5. Setup № 6. Others № 2. Identification Mode                               |
| Sequence Number              | 5. Setup ↦ 6. Others ↦ 2. Identification Mode.                              |
|                              | Resets to "1".  |
| Identification Number        | 5. Setup ↦ 6. Others ↦ 2. Identification Mode.                              |
|                              | Resets to "1".  |
| Reset Time                   | 5. Setup ↦ 6. Others ↦ 2. Identification Mode                               |
| Operators                    | 5. Setup ↦ 6. Others ↦ 1. Calibration ↦ 2. Define Operators                 |
| Calibration Factors          | 2. Calibration → 2. Cal Factors   |
| CV% Limits                   | 5. Setup ↦ 6. Others ↦ 1. Calibration ↦ 1. CV% Limits                       |
| Flow Cell WBC Calibration    | 4. Diagnostics ↦ 5. Service ↦ 8. Flowcell WBC Calibrationn ↦ 2. Cal Factors |
|                              | %WBC BA1 50.0   |
|                              | %WBC BA2 20.0   |
|                              | %WBC BA3 15.0   |
|                              | FLOW CELL WBC X.XX  |
|                              | WBC FLOW CELL BALANCE (enable)  |
| Daily Workload               | 3. Reagents ↦ 2. Daily Workload   |

#### Table 4.39-2 A<sup>C</sup>•T 5diff Menu Paths - System Settings

| Parameter                | From Main Menu  |  |  |
|--------------------------|---|--|--|
| Cycle Counts             | 5. Setup ↦ 6. Others ↦ 7. Cycle Counts  |  |  |
|                          | Resets to "0".  |  |  |
| Autoclean Frequency      | 5. Setup ↦ 6. Others ↦ 3. Autoclean Frequency   |  |  |
| Burn In                  | 4. Diagnostics ↦ 5. Service ↦ 7. Burn-In ↦ 1. Burn-In Cycles  |  |  |
| Heating Coil Thermistor  | 4. Diagnostics ↦ 5. Service ↦ 3. Heating Systems ↦ 1. Heating Coil ↦ 1. Adjustment.   |  |  |
| Value                    | Press ENTER for calculation after updating the value.   |  |  |
| Heating Coil Reference   | 4. Diagnostics ↦ 5. Service ↦ 3. Heating Systems ↦ 1. Heating Coil ↦ 2. Reference.  |  |  |
| Temperature              | Adjust reference temperature setting.   |  |  |
| Bath Enclosure           | 4. Diagnostics ↦ 5. Service ↦ 3. Heating Systems ↦ 2. Bath Enclosure ↦ 1. Adjustment  |  |  |
| Thermistor Value         | Press ENTER for calculation after updating the value.   |  |  |
| Bath Enclosure Reference | 4. Diagnostics ↦ 5. Service ↦ 3. Heating Systems ↦ 2. Bath Enclosure ↦ 2. Reference   |  |  |
| Temperature              | Adjust reference temperature setting.   |  |  |
| Dilution                 | 4. Diagnostics ↦ 5. Service ↦ 1. Dilution   |  |  |
| Mixing                   | 4. Diagnostics ↦ 5. Service ↦ 4. Mixing   |  |  |
| Vacuum                   | 4. Diagnostics ↦ 5. Service ↦ 6. Vacuum Check ↦ 1. Counting   |  |  |
| Pulse Adjustment         | 4. Diagnostics ↦ 5. Service ↦ 2. Measurement ↦ 6. Pulse Adjustment  |  |  |
| Reagent Volumes          | 5. Setup ↦ 6. Others ↦ 6. Reagent Volumes   |  |  |
| Reagent Lot Numbers      | 3. Reagents → 1. Level Change then select Change All.   |  |  |
|                          | Enter the lot numbers from the old system settings. After the last number is entered, the instrument primes all the reagents.                               |  |  |
|                          | <b>Note:</b> If new or full reagent containers are not placed in the system, the cycle count will no represent the actual volume of reagent in the bottles. |  |  |
| Printer Configuration    | 5. Setup ↦ 5. Printer ↦ 1. Printer Configuration.   |  |  |
|                          | In addition, Rev 1.03 or later has Histograms Thresholds as an option.  |  |  |
| Institutional Header     | 5. Setup ↦ 5. Printer ↦ 2. Institutional Header   |  |  |
| Sending Options          | 5. Setup ↦ 4. Host Setup ↦ 3. Sending Options   |  |  |
| Host Setup Configuration | 5. Setup ↦ 4. Host Setup ↦ 1. Host Setup Configuration  |  |  |
| Format                   | 5. Setup ↦ 4. Host Setup ↦ 2. Sending Configuration   |  |  |
| Variable Format Setup    | 5. Setup ↦ 4. Host Setup ↦ 4. Variable Format Setup   |  |  |
|                          |   |  |  |

#### Table 4.39-2 AC•T 5diff Menu Paths - System Settings (Continued)

#### Patient Ranges, Action Ranges and Thresholds

If the instrument was using the default values for *Patient Ranges, Action Ranges* and *Thresholds* (on all 9 ranges), then the software retains these values. If the values were not set to default, then re-enter the values from the old system setup.

Note: Rev 1.03 Software defaults to "Simple Flagging" and no "Threshold Flags." To set the DIFF Plot and Histogram Flags to On, from the Main Menu, select

#### 5. Setup >> 5. Printer >> 1. Printer Configuration then enable DIFF Plot and Histogram Flags.

# **Verifying Main Card Settings**

Use this procedure to verify the Main card settings after the replacement of the card.

1. Compare the settings listed in Table 4.39-3 to the readings taken from the Main card and adjust as required. Use Figure 4.39-2 to locate the potentiometer and corresponding test point.

| • |  |
|---|--|
| ٠ |  |

| Table 4.39-3         AcT 5diff - Main Card Settings |
|---|
|---|

| Function (Heading)                      | Adjustn              | nent           | Target        | Range   | Test Point   |
|---|----------------------|----------------|---------------|---------|--|
| Motor Current Adjustment                | Draining Syringe     | R149           | 4.0 V         | ±0.05 V | TP5  |
| (4.13)                                  | Counting Syringe     | R150           | 4.0 V         | ±0.05 V | TP6  |
|   | Diluter Syringe      | R151           | 4.0 V         | ±0.05 V | TP7  |
|   | Injector Syringe     | R152           | 3.0 V         | ±0.05 V | TP8  |
|   | Horizontial Carriage | R154           | 3.0 V         | ±0.05 V | TP10   |
|   | Sample Syringe       | R155           | 2.0 V         | ±0.05 V | TP11   |
|   | Probe Carriage       | R156           | 4.5 V         | ±0.05 V | TP12   |
| Threshold Adjustments                   | BASO                 | R157           | 300 mV        | ±5 mV   | TP14   |
| (4.14)                                  | RBC                  | R158           | 300 mV        | ±5 mV   | TP13   |
|   | PLT                  | R159           | 300 mV        | ±5 mV   | TP2  |
|   | FLOW CIS             | R160           | 650 mV        | ±5 mV   | TP3  |
|   | FLOW OD              | R161           | 350 mV        | ±5 mV   | TP4  |
| Drain Sensor Adjustment                 | Drain Sensor         | R287           | 4.5 V (empty) | ±0.3 V  | TP52   |
| (4.11)                                  |                      |                | <1.0 V (full) |         | (under P30)  |
| DIFF Bath Drain Sensor                  | Diff Bath Drain      | R286           | 4.5 V (empty) | ±0.3 V  | TP48   |
| Adjustment (4.12)                       | Sensor               |                | <1.0 V (full) |         | (under P29)  |
| Vacuum Checks and<br>Adjustments (4.17) | Counting Syringe     | Step value     | 6.5 inches Hg | stable  | test box attached<br>to bottom port<br>on side of<br>syringe |
| Vacuum Checks and<br>Adjustments (4.17) | Draining Syringe     | Non-adjustable | 7.7 inches Hg | stable  | test box attached<br>to top port on<br>side of syringe       |
| HGB Blank Adjustment (4.7)              |                      | R248           | 4.7 V         | ±0.3 V  | screen   |
| Aperture Current Check<br>(4.8)         |                      | Non-adjustable | 60 V          |         | J22, J23, J24  |
| RBC/PLT Gain Adjustment                 | RBC Gain             | R133           | Channel 78    |         | screen   |
| (4.9)                                   | PLT Gain             | R135           | Channel 112   |         | screen   |
| WBC/BASO Gain<br>Adjustment (4.10)      |                      | R134           | Channel 102   |         | screen   |

| Function (Heading)                     | Adjustn                               | nent | Target                              | Range                      | Test Point  |  |
|--|---------------------------------------|------|-------------------------------------|----------------------------|-------------|--|
| DIFF Lamp Voltage<br>Adjustment (4.4)  |                                       | R11  | 6.0v                                | 5.50 - 6.50 V              | screen      |  |
| Transfer Time (4.4)                    |                                       | shim | 200                                 | 150 - 250                  | screen      |  |
| Resistive Channel                      |                                       | R136 | 50                                  | 45 - 55                    | screen      |  |
| Adjustment (4.4)                       | Final Gain Using<br>Fresh Whole Blood | R136 | good<br>scatterplot<br>distribution |                            | scatterplot |  |
| Absorbance Channel<br>Adjustment (4.4) |                                       | R148 | >170                                | adjust to max<br>170 - 190 | screen      |  |
|  | Final Gain Using<br>Fresh Whole Blood | R148 | good<br>scatterplot<br>distribution |                            | scatterplot |  |

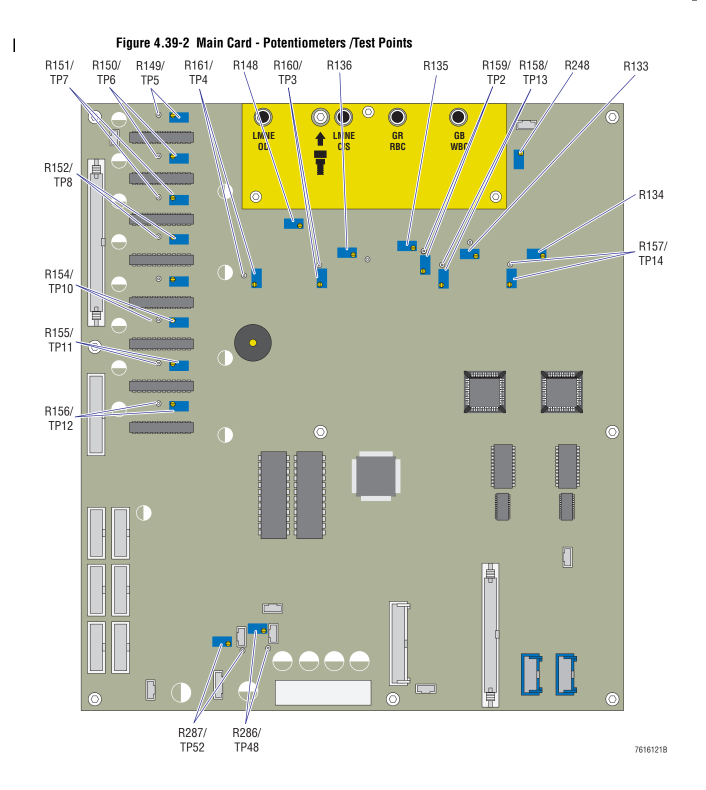
- 2. Use a fresh normal whole-blood specimen to check instrument reproducibility.
  - a. From the Main Menu, select **2. CALIBRATION → 4. REPRODUCIBILITY**.
  - b. Run eleven samples and then delete the first one for a ten-shot reproducibility.
  - c. Verify the reproducibility results are within acceptable limits. See Table 4.39-4.

Table 4.39-4 Whole-Blood Reproducibility CV Limits for 20 Cycles

| Parameter | %CV   | Test Level                    |
|-----------|-------|-------------------------------|
| WBC       | <2.0% | at 10.0 x 10 <sup>3</sup> /µL |
| RBC       | <2.0% | at 5.00 x 10 <sup>6</sup> /µL |
| Hgb       | <1.0% | at 15.0 g/dL                  |
| Hct       | <2.0% | at 45.0%                      |
| MCV       | <1.0% | at 90.0 fL                    |
| Plt       | <5.0% | at 300 x 10 <sup>3</sup> /µL  |

3. Run controls and confirm results fall within expected ranges.

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**SERVICE AND REPAIR PROCEDURES** MAIN CARD REPLACEMENT AND SOFTWARE TRANSFER

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# TABLES

5.1-1 Maintenance Schedule, 5.1-1

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# 5.1 RECOMMENDED MAINTENANCE SCHEDULE

Table 5.1-1 lists the maintenance procedures for the  $A^{C\bullet}T$  5diff hematology analyzer and when they should be performed. Maintenance is based on the number of cycles; therefore, the frequency a maintenance procedure needs to be done is driven by the customer workload. The maintenance schedule in Table 5.1-1 is the customer workload translated into a time schedule.

| Workload (Cycles Per Day)                       | <30 |    | 30 to 120 |    |     |    | >120 |    | PMI Kit                           |
|---|-----|----|-----------|----|-----|----|------|----|-----------------------------------|
| Maintenance Frequency                           | 1Y  | 2Y | 6M        | 1Y | 18M | 2Y | 6M   | 1Y |                                   |
| Replace rinse block O-ring and washer           | Х   | Х  | Х         | Х  | Х   | Х  | Х    | Х  | 6 months                          |
| Replace reagent syringe O-rings and washers     | Х   | Х  | Х         | Х  | Х   | Х  | Х    | Х  | Maintenance Kit,<br>PN - XEA485AS |
| Replace waste syringe O-ring and washer         | Х   | Х  | Х         | Х  | Х   | Х  | Х    | Х  | 1 N - XLA+03A3                    |
| Replace count syringe O-ring and washer         | Х   | Х  | Х         | Х  | Х   | Х  | Х    | Х  |                                   |
| Clean the bath enclosure                        | Х   | Х  | Х         | Х  | Х   | Х  | Х    | Х  |                                   |
| Replace sample syringe O-rings                  | Х   | Х  |           | Х  |     | Х  | Х    | Х  | 1 year                            |
| Replace 5diff syringe O-rings                   | Х   | Х  |           | Х  |     | Х  | Х    | Х  | Maintenance Kit,<br>PN - XEA486AS |
| Replace draining chamber O-rings                | Х   | Х  |           | Х  |     | Х  | Х    | Х  |                                   |
| Replace optical bench lamp                      | Х   | Х  |           | Х  |     | Х  | Х    | Х  |                                   |
| Replace flow cell coaxial cable                 | Х   | Х  |           | Х  |     | Х  | Х    | Х  |                                   |
| Replace diluent reservoir O-ring and washer     | Х   | Х  |           | Х  |     | Х  | Х    | Х  |                                   |
| Replace counting head coax and aperture O-rings | Х   | Х  |           | Х  |     | Х  | Х    | Х  |                                   |
| Replace sample probe and rinse block            |     | Х  |           |    |     | Х  |      | Х  | Every 2 years                     |
| Replace reagent syringe pistons                 |     | Х  |           |    |     | Х  |      | Х  | Maintenance Kit,<br>PN - XEA581AS |
| Replace waste syringe piston                    |     | Х  |           |    |     | Х  |      | Х  |                                   |
| Replace count syringe piston                    |     | Х  |           |    |     | Х  |      | Х  |                                   |

#### Table 5.1-1 Maintenance Schedule

**MAINTENANCE PROCEDURES** RECOMMENDED MAINTENANCE SCHEDULE

# 5.2 MAINTENANCE WORKLIST

#### Purpose

Use this procedure as a guide when the 6-months, 1-year, or every 2-years maintenance is needed. 6-months maintenance mainly consists of replacing O-rings and washers. The 1-year maintenance is more extensive and includes all the procedures that would typically be done during the 6-months maintenance. The every 2-years maintenance is the most extensive because it also includes all the 6-months and 1-year maintenance procedures.

A worklist is provided to guide you through the various procedures and checks that need to be done to complete a scheduled maintenance. Make a copy of this worklist before starting the maintenance procedures and use it as a checklist to help you keep track of your progress. You may also use this worklist as a record if you sign and date it.

#### **Tools/Supplies Needed**

- □ 6 months maintenance kit, PN XEA485AS
- □ Allen wrench, 2.5 mm and 3.0 mm
- Torque screwdriver, 2.5 mm hex-ball and T10 torx
   Note: 100 mN.m (14.2 ozf.in) and 400 mN.m (56.8 ozf.in) torques are required.
- □ Cutting pliers
- □ High quality, fragrance-free bleach (10-12% sodium hypochlorite available chloride)
- Distilled water
- □ Absorbent paper towels
- □ Lint-free tissues

#### Additional Tools or Supplies Needed for 1-Year Maintenance

- □ 1 year maintenance kit, PN XEA486AS
- □ Small Phillips-head screwdriver
- □ Allen wrench, 2 mm
- □ Hemostat or tweezers with a pointed end
- □ Small side cutters
- □ White paper towel or cloth
- □ Cotton-tip applicator swab

#### Additional Tools or Supplies Needed for the Every 2-Years Maintenance

□ Every 2 years maintenance kit, PN - XEA581AS

#### **User Mode**

While performing these maintenance procedures, you are instructed to set the User mode to Service. In this mode, the right side door interlock is bypassed. When the instrument is operating with the right side door open, it is important to avoid any unnecessary contact with the sample probe. The probe is sharp and may contain biohazard materials. When the service call is complete, make sure the interlock is reactivated to ensure the Customer is not accidently injured by the sample probe or its movement.

### **Worklist Instructions**

#### **Overview**

The maintenance worklist is a composite guide for the 6-months, 1-year, and every 2-years maintenance procedures. After providing you with the steps required to prepare the instrument for maintenance, the worklist directs you to the individual service and repair procedures needed to complete maintenance in the left compartment. When these replacements are complete, an interim verification check is done to ensure there is no leakage before moving to the right compartment. After performing replacements inside the right compartment, another verification is done to again ensure there is no leakage. When all the required maintenance procedures are completed, an overall system verification is done to ensure proper operation.

#### **Maintenance Category Identifier**

Since this worklist is a composite of the three possible maintenance schedules, a block containing the maintenance category appears before each procedure:

**[6 months / 1 year / 2 years]** indicates a procedure that must be completed each time maintenance is done on an  $A^{C} \bullet T$  5diff hematology analyzer.

**[1 year / 2 years]** indicates a procedure that is completed only when the 1 year or every 2 years maintenance is done. When doing a 6 months maintenance, this procedure is not required.

**[6 months]** appears only once to direct you to perform the System Verification Procedures when you are doing a 6 months maintenance.

If you are doing a 6 months maintenance, follow the instructions in the rows marked **[6 months / 1 year / 2 years]** or **[6 months]**. If you are doing a 1 year or every 2 years maintenance, follow the instructions in the rows marked **[6 months / 1 year / 2 years]** or **[1 year / 2 years]**.

#### **Basic Instructions**

Preparation instructions for performing the required maintenance are provided at the beginning of the worklist. These instructions supersede the preparation instructions in the individual service and repair procedures. As a result, when you are directed to a particular service and repair procedure, you are usually directed to begin at the **Removal** heading and further directed as to what subsequent headings need to be completed. In most cases, you will not complete the Verification in the individual procedures. When all the required maintenance procedures are completed, you will be directed to do an overall system verification to ensure the instrument is operating properly before you close out the service call. It is important for you understand that by carefully reading and following the directions associated with each procedure you will avoid doing unnecessary work.

Before starting a maintenance call, make a copy of the complete worklist that begins on the next page. Since this worklist consolidates all the individual procedures, it is important to progress through the procedures in their designated order. Use this worklist as a guide to ensure all needed maintenance procedures are completed. As you finish each task, place a checkmark ( $\checkmark$ ) next to the completion statement to keep track of your progress as you work through the various procedures.

# **MAINTENANCE WORKLIST**

Copy this worklist and use it as a guide to ensure all needed maintenance procedures are completed. In the electronic version of this manual, a file (worklist.pdf) is provided for printing. As you finish each task, place a checkmark ( $\checkmark$ ) next to the completion statement to keep track of your progress as you work through the various procedures.

# **Replacements in the Left Side Compartment**

#### [6 months / 1 year / 2 years]

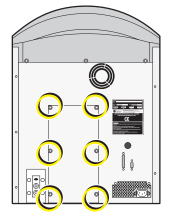
#### Preparation

- 1. Use the maintenance kit part listings to ensure no parts are missing from the kit.
  - If performing a 6-months maintenance, use Table 8.1-10.
  - If performing the 1-year maintenance, use Table 8.1-10 and 8.1-11.
  - If performing every 2-years maintenance, use Table 8.1-10, 8.1-11, and 8.1-12.
- 2. If off, turn the instrument on.
- 3. From the Service menu, select 9. OTHERS → 1. USER MODE.
- 4. At the instrument keypad,
  - a. Press the down arrow key to move the blinking cursor to the SERVICE box.
  - b. Press the decimal point key. Make sure a dot appears inside the box.

Note: If you press the decimal point too long, the dot may disappear. The dot indicates the right side door interlock is bypassed.

- c. Press **ESC** to return to the Others menu.
- 5. From the Others menu, select **3. PARK SYRINGES**. When the park syringes prompt appears, press **ENTER**.
- 6. When the park syringes routine is complete, turn the instrument off.
- 7. At the rear of the instrument,
  - a. Disconnect the power cord.
  - b. Remove the six hex screws securing the rear access panel.

**Note:** Use a 3 mm hex key to remove any hex screws securing an instrument cover. For details, see Heading 4.2.



c. Remove the rear access panel and set it aside.

8. Remove the four hex screws securing the left side panel to the instrument frame and set the door aside.

**CAUTION** A flat cable is connected to the rear of the Main card. Be careful when you open the door (where the Main card is attached) that you do not disconnect or damage this cable.

- 9. In the left side compartment,
  - a. Turn the two captive knobs counterclockwise to release the Main card door. These knobs are located to the right of the Main card.
  - b. Open the Main card door and carefully anchor the door behind the white plastic catch to keep it open.

#### [1 year / 2 years]

- 10. Remove the top cover:
  - a. In the left side compartment, remove the hex screw in the front upper corner.
  - b. Open the right side door and remove the hex screw in the front upper corner.
  - c. At the rear of the instrument, remove the three hex screws securing the top cover to the instrument frame.
  - d. Carefully remove the top cover and set it aside.

#### Preparation complete.

#### [6 months / 1 year / 2 years]

#### Reagent Syringes Assembly Replacement Parts

From the 6 months maintenance kit:

- □ Washer and silicone O-ring, PN - XDA622A (4 sets)
- □ Silicone O-ring for the Hgb Lyse reagent syringe piston, PN FAA065A

□ Silicone grease, PN - XEA019A

#### Procedure

From the every 2 years maintenance kit:

- Reagent syringe piston, PN - GBC030A (need 4)
- □ Hgb Lyse reagent syringe piston, PN - GBC031A

Under Heading 4.24, REAGENT SYRINGES ASSEMBLY REPLACEMENTS, complete the following sections:

\_\_\_\_\_ Removal

\_\_\_\_\_ O-Ring, Washer, and Piston Replacement

- If performing the 6 months or 1 year maintenance, replace the O-rings and washers only. Do not replace the pistons.
- If performing the every 2 years maintenance, replace all pistons, O-rings, and washers.
- \_\_\_\_ Installation

All reagent syringes assembly replacements are complete.

#### [1 year / 2 years]

#### **5diff Syringe Assembly**

#### **Replacement Parts**

From the 6 month maintenance kit:

□ Silicone grease, PN - XEA019A

From the 1 year maintenance kit:

- □ Silicone O-ring, PN FAA040A
- □ Fluorocarbon O-rings, PN FAA067A (need 4)

From the every 2 years maintenance kit:

□ Count syringe piston, PN - GBG052A

#### **Procedure**

Under Heading 4.29, 5diff SYRINGE ASSEMBLY REPLACEMENTS, complete the following sections:

\_\_\_\_\_ Removal

**O-ring Replacements** Note: Replace five O-rings, one silicone and four fluorocarbon. Installation

All five O-rings are replaced.

#### [6 months / 1 year / 2 years]

#### **Count Syringe Assembly**

#### **Replacement Parts**

From the 6 months maintenance kit:

- □ Washer and O-ring, PN XDA621A
- □ Silicone grease, PN XEA019A
- **Procedure**

Under Heading 4.25, COUNT SYRINGE COMPONENT REPLACEMENTS, complete the following sections:

Removal

- \_\_\_\_ O-ring, Washer, and Piston Replacement
  - If performing the 6 months or 1 year maintenance, replace the O-rings and • washers only. Do not replace the pistons.
  - If performing the every 2 years maintenance, replace all pistons, O-rings, and washers.

Installation

All count syringe replacements are complete.

#### **Interim Verification Check**

- 1. Reconnect the power cord.
- 2. Turn the instrument on.
- 3. Press the **STARTUP** button on the instrument keypad.
- 4. When the Startup routine and background check are done,
  - If performing the 6 months maintenance, verify the reagents syringe assembly and the count syringe assembly are not leaking.
  - If performing the 1 year or every two years maintenance, verify the reagents syringe assembly, the 5diff syringe assembly, and the count syringe assembly are not leaking.
- 5. Open the right side door and check the right compartment for leaks. A tubing connected on the wrong port inside the left compartment may cause leaking inside the right compartment.
- 6. If no leakage is seen, prime the reagents.
  From the Main Menu, select 3. REAGENTS → 3. PRIME → 6. ALL REAGENTS.
- 7. When the prime cycles are done,
  - a. Check the reagents syringe assembly, the 5diff syringe assembly (if applicable), and the count syringe assembly for leaks.
  - b. Check the right side compartment for leaks.
- 8. If no leaks are detected,
  - a. Turn the instrument off.
  - b. Disconnect the power cord.
  - c. Do the electronic replacements that follow.

Syringe assemblies inside the left compartment are not leaking.

#### [1 year / 2 years]

#### Replace the Optical Bench Lamp

#### **Replacement Part**

From the 1 year maintenance kit:

□ Optical bench lamp, PN - DAJ007A

#### Procedure

Under Heading 4.31, OPTICAL BENCH LAMP REPLACEMENT, complete the following sections:

\_\_\_\_\_ Removal

\_\_\_\_ Lamp Replacement

Optical bench lamp is replaced.

#### [1 year / 2 years]

#### **Replace the Flow Cell Coaxial Cable**

#### **Replacement Part**

From the 1 year maintenance kit:

□ DIFF flow cell coaxial cable, PN - XBA399A

#### Procedure

Under Heading 4.30, FLOW CELL COAXIAL CABLE REPLACEMENT, complete the following sections:

\_\_\_\_\_ Removal

\_\_\_\_\_ Replacement

\_\_\_\_\_ Under Heading 4.4, do the Flow Cell Checks.

Note: If a value is outside acceptable limits, do the appropriate adjustment.

Flow cell coaxial cable is replaced and the flow cell checks are within acceptable limits.

# **Replacements in the Right Side Compartment**

#### [6 months / 1 year / 2 years]

#### Preparation

- 1. From the Service menu, select **9. OTHERS → 3. PARK SYRINGES**. When the park syringes prompt appears, press **ENTER**.
- 2. When the park syringes routine is complete, turn the instrument off.
- 3. Disconnect the power cord.
- 4. Open the pneumatic access door (right side of the instrument).
- 5. Manually push down on the waste syringe piston until it reaches its lowest position.

#### Preparation complete.

#### Sample Probe and Rinse Block Assembly

#### **Replacement Parts**

From the 6 month maintenance kit:

From the 2 year maintenance kit:

□ O-ring, fluorocarbon, PN - FAA053A

□ Silicone grease, PN - XEA019A

- □ Sample probe, PN XDA619AS
- □ Sample probe guide, PN GBG091A

#### Procedure

In the right side compartment, manually push the sample probe housing towards the rear of the instrument then under Heading 4.26, SAMPLE PROBE AND RINSE BLOCK ASSEMBLY COMPONENT REPLACEMENTS, complete the following sections:

\_\_\_\_\_ Removal

\_\_\_\_\_ Rinse Block Assembly Component Replacement

- If performing the 6 months or 1 year maintenance, replace the O-ring only.
- If performing the every 2 years maintenance, replace the sample probe, the probe guide, and its O-ring.

\_\_\_\_ Installation

All needed components are replaced.

#### [1 year / 2 years]

#### Sample Syringe Assembly

#### **Replacement Part**

From the 1 year maintenance kit:

□ Fluorocarbon O-ring, PN - FAA064A (need 2)

#### Procedure

Under Heading 4.33, SAMPLE SYRINGE ASSEMBLY REPLACEMENTS, complete the following sections:

\_\_\_\_\_ Removal

\_\_\_\_\_ O-ring Replacements

\_\_\_\_\_ Installation

#### O-rings replaced.

#### Waste Syringe Assembly

#### Replacement Parts

From the 6 months maintenance kit:

From the every 2 years maintenance kit:

□ Waste syringe piston, PN - GBG052A

- □ Washer and O-ring, PN XDA621A
- □ Silicone grease, PN XEA019A

#### Procedure

Under Heading 4.27, WASTE SYRINGE COMPONENT REPLACEMENTS, complete the following sections:

\_\_\_\_\_ Removal

\_\_\_\_\_ O-ring, Washer, and Piston Replacement

- If performing the 6 months or 1 year maintenance, replace the O-ring and washer only. Do not replace the piston.
- If performing the every 2 years maintenance, replace the syringe piston, O-ring, and washer.

\_\_\_\_ Installation

All waste syringe replacements are complete.

#### [1 year / 2 years]

#### **Diluent Reservoir**

#### **Replacement Parts**

From the 1 year maintenance kit:

□ O-ring and washer, PN - XEA286AS

#### Procedure

Under Heading 4.32, DILUENT RESERVOIR REPLACEMENTS, complete the following sections:

\_\_\_\_\_ Removal

\_\_\_\_\_ O-ring and Washer Replacement

\_\_\_\_\_ Installation

#### O-ring and washer replaced.

#### **Interim Verification Check**

- 1. Reconnect the power cord.
- 2. Turn the instrument on.

**WARNING** Risk of personal injury. The right side door interlock is bypassed which allows you to observe instrument operation with the right side door open. Avoid any unnecessary contact with the sample probe. The probe is sharp and may contain biohazardous materials.

- 3. Press the **STARTUP** button on the instrument keypad.
- 4. When the Startup routine and background check are done,
  - If performing the 6 months maintenance, verify the waste syringe is not leaking.
  - If performing the 1 year or every two years maintenance, verify the waste syringe, the diluent reservoir, and the sample syringe are not leaking.
- 5. Check the left side compartment for leaks. A tubing connected on the wrong port inside the right compartment may cause leaking inside the left side compartment.
- 6. From the Main Menu, select **4. DIAGNOSTICS → 4. HARDWARE SYSTEMS → 4. TRAVERSE SERVICE POSITION**.
  - a. When the traverse service position prompt appears, press ENTER.
  - b. Verify the sample probe rinse block is not leaking.
  - c. Press **ESC** twice to return to the Diagnostics menu.

#### Assemblies inside the right and left compartments are not leaking.

#### [1 year / 2 years]

#### Preparation

- From the Diagnostics menu, select 3. DILUTER SYSTEMS → 3. DRAIN BATHS → 6. ALL BATHS.
- 2. Verify all baths (including the draining baths and rinse chamber) are empty.
- 3. Turn the instrument off.
- 4. Disconnect the power cord.
- 5. Do the draining bath and counting head replacements that follow.

#### Preparation complete.

#### [1 year / 2 years]

#### Draining Baths

#### **Replacement Parts**

From the 1 year maintenance kit:

□ O-ring, PN - FAA066A (one for each bath)

#### Procedure

Under Heading 4.34, DRAINING BATH REPLACEMENTS, complete:

\_\_\_\_\_ O-ring Replacements

O-rings are replaced.

#### [1 year / 2 years]

#### **Counting Heads**

#### **Replacement Parts**

From the 1 year maintenance kit:

□ Coaxial cable O-ring, PN - FAA046A (need 1 for each bath)

Aperture O-ring, PN - GBG156A (need 2 for each bath)

#### Procedure

Under Heading 4.35, O-RING REPLACEMENTS IN THE COUNTING BATHS (RBC and WBC/BASO Baths), complete the following sections:

\_\_\_\_\_ Removal

\_\_\_\_\_ O-ring Replacements

\_\_\_\_\_ Installation

\_\_\_\_\_ Align the Bath Assembly

\_\_\_\_\_ Verification which also includes the SYSTEM VERIFICATION PROCEDURES.

Coaxial cable and aperture O-rings are replaced. None of the baths are leaking.

\_\_\_\_\_ All system verification checks are acceptable.

#### [6 months / 1 year / 2 years]

#### Clean the Bath Enclosure

Under Heading 4.28, CLEANING THE BATH ENCLOSURE, complete the procedure as written.

\_\_\_\_\_ Bath enclosure cleaned as directed.

# Wrap Up

#### [6 months]

**WARNING** Risk of personal injury. The right side door interlock is bypassed which allows you to observe instrument operation with the right side door open. Avoid any unnecessary contact with the sample probe. The probe is sharp and may contain biohazardous materials.

Perform a system verification as instructed under Heading 5.3, SYSTEM VERIFICATION PROCEDURES.

#### All system verification checks are acceptable.

#### [1 year / 2 years]

- 1. Replace the top cover:
  - a. Carefully position the top cover on the instrument.
  - b. In the left side compartment, replace the hex screw in the front upper corner.
  - c. Open the right side door and replace the hex screw in the front upper corner.
  - d. At the rear of the instrument, replace the three hex screws that secure the top cover to the instrument frame.

#### [6 months / 1 year / 2 years]

- 2. Replace the rear access panel using the six hex screws removed earlier.
- 3. In the left side compartment,
  - a. Lift the white plastic catch to release the Main card.
  - b. Close the Main card door and turn the two captive knobs clockwise to secure it.
- 4. Replace the left side panel and install the four hex screws removed earlier.
- 5. Close the right side door.

**WARNING** Risk of personal injury. The right side door interlock is bypassed allowing operation with the right side door open. The probe is sharp and may contain biohazardous materials. When all the maintenance procedures are complete, make sure the interlock is reactivated to ensure the Customer is not accidently injured by the sample probe or its movement.

- 6. From the Service menu, select **9. OTHERS → 1. USER MODE.**
- 7. At the instrument keypad,
  - a. Press the down arrow key to move the blinking cursor to the *SERVICE* box.
  - b. Press the decimal point or **DEL** key. Make sure the dot no longer appears inside the *SERVICE* box. The right side door interlock is reactivated.

**Note:** If you press the decimal point too long, the dot may reappear. The dot indicates the right side door interlock is bypassed.

c. Press **ESC** as many times as necessary to return to the Main Menu.

All panels and covers are replaced. The right side door interlock is reactivated.

| <br>6 months maintenance procedures are completed.        |
|---|
| <br>1 year maintenance procedures are completed.          |
| <br>Every two years maintenance procedures are completed. |
| Date of Completion<br>Institution<br>Completed by         |
| <br>Date of Completion                                    |

# 5.3 SYSTEM VERIFICATION PROCEDURES

#### Purpose

Use this procedure at the end of each service call to verify the A<sup>C</sup>•T 5diff hematology analyzer is operating correctly.

#### **Tools/Supplies Needed**

- □ Fresh, normal whole- blood specimens
- □ Calibration blood specimens

#### **Preparation**

**WARNING** Risk of personal injury or contamination. If you do not properly shield yourself while servicing the instrument, you may become injured or contaminated. To prevent possible injury or biological contamination, you must wear appropriate safety glasses, a lab coat, and gloves when servicing the instrument.

**IMPORTANT** Risk of misleading results. This procedure must be performed on a clean instrument. If it is suspected that the instrument is not perfectly clean, perform the PREPARATION TO SHIP THE INSTRUMENT procedure under Heading 4.3.

#### Startup

Run a Startup cycle and background check.

#### Reproducibility

Use one fresh, normal whole-blood specimen to check instrument reproducibility.

- 1. From the Main Menu, select 2. CALIBRATION → 4. REPRODUCIBILITY.
- 2. Do a 20 sample reproducibility study in the CBC mode of operation. Verify the results from these runs are not flagged.
- 3. Verify the reproducibility results are within acceptable limits. See Table A.1-6.
- 4. Run a fresh, normal whole-blood specimen in the CBC/DIFF to verify normal specimens are not flagging in the CBC/DIFF mode.

### Calibration

#### Autocalibration

- 1. From the Main Menu, select **2. CALIBRATION → 1. AUTOCALIBRATION**.
- 2. At the Autocalibration screen, change the lot number, expiration date, or target values as necessary.

#### **Run Calibration**

- 1. Prepare the calibrator according to the specific instructions (temperature, mixing, and so forth).
- 2. Open the vial and submerge the sample probe inside the vial.
- 3. Press the aspirate switch located behind the probe.
- 4. When the cycle LED stops flashing, remove the vial and replace the cap on the calibrator.
- 5. When the analysis cycle ends, the result is displayed.

**IMPORTANT** The calibration of the A<sup>C</sup>•T 5diff hematology analyzer can be performed on 3 to 11 analyses. To obtain the best calibration possible, it is recommended to run the calibration blood a minimum of five times. The autocalculation module performs statistics on these results to obtain the best calibration factors.

6. Remix the calibrator and repeat steps 2 through 5 until at least three, but no more than 11 calibrator samples are analyzed. It is recommended that you run the calibrator at least five times to achieve the best calibration.

Note: After three runs, the instrument calculates calibration statistics.

#### **Calibration Passed**

Calibration passes if the statistics are within the acceptable limits:

- Coefficient of variation (CV) is within the limits setup as described.
- The percentage difference between the target and the mean value is less than 20.
- 1. At the instrument keypad,
  - a. Press Enter to save the new calibration factors.
  - b. Press **Enter** again to print the new calibration factors.
- 2. Check that the calibration factors are within the acceptable ranges listed in Table A.1-7.

#### Step-By-Step Cycle Check

Use this step-by-step description of the cycle as a guide to help you verify proper instrument operation at the end of a service call. This check covers the four principal phases of a cycle:

- 1. Sample preparation (making the dilutions)
- 2. Count and measurement of the WBC group
- 3. RBC/PLT group count
- 4. Filling the diluent reservoir

A special cycle is required when no other cycle is started during RBC/PLT count or during the filling of the diluent reservoir: rinsing of the RBC bath.

All phases are described in this procedure. Use Figures A.3-1 and A.3-2 If you need help locating an assembly.

#### Preparation

- 1. Remove the four hex screws securing the left side panel to the instrument frame and set the door aside.
- 2. If you have not already done so, change the User mode to Service. This bypasses the right side door interlock so you can cycle with the right side door open.

From the Service menu, select **9**. **OTHERS → 1**. **USER MODE**. For details, see User Mode under Heading 4.1.

3. Open the right side door.

#### **Instrument At Rest**

When the A<sup>C</sup>•T 5diff analyzer is at rest (power is ON but the instrument is not cycling), verify:

- 1. The sample probe is in its home position and the green LED is glowing.
- 2. In the right compartment, verify all the baths (except the rinse chamber) are filled with diluent and the piston inside the waste syringe is up.
- 3. Inside the left compartment, verify the pistons inside the count syringe, 5diff syringe, and reagent syringes assemblies are down.

**Note:** It will be necessary to cycle a whole-blood specimen several times to see the following actions.

#### Sample Preparation (Making the Dilutions)

**WARNING** Risk of personal injury. When SERVICE is selected as the User mode, the right side door interlock is bypassed allowing instrument operation with the right side door open. Avoid any unnecessary contact with the sample probe. The probe is sharp and may contain biohazardous materials. When the service call is complete, make sure the interlock is reactivated to ensure the Customer is not accidently injured by the sample probe or its movement.

Verify sample probe movement and dilution preparation:

- □ Press the aspirate switch. All the baths are drained (waste syringe).
- $\Box$  53 µL whole-blood sample is aspirated (sample syringe).
- □ Sample probe moves over the rinse chamber (sample probe, probe motor, traverse motor).
- $\Box$  3 µL sample aliquot at the tip of the sample probe is discarded into the rinse chamber as the exterior of the sample probe is rinsed (sample syringe, diluent reagent syringe).
- □ Sample probe moves from the rinse chamber to the first dilution (DIL1/HGB) bath (sample probe motor, traverse motor).
- □ 10 µL of whole-blood partitioned for making the first dilution is delivered to the DIL1/HGB bath using a tangential flow of 1.7 mL of diluent (sample syringe, diluent reagent syringe). Mixing bubbles enter the bath. This is the primary 1:170 dilution (or first dilution).
- □ Sample probe moves from the first dilution (DIL1/HGB) bath to the WBC/BASO bath (sample probe motor, traverse motor).
- 10 µL of whole-blood partitioned for making the WBC/BASO dilution is delivered to the WBC/BASO bath using a tangential flow of 2.0 mL of WBC Lyse (sample syringe, WBC Lyse reagent syringe). Mixing bubbles enter the bath. This is a 1:200 dilution.
- □ Sample probe moves from the WBC/BASO bath to the DIFF bath (sample probe motor, traverse motor).
- 25 µL of whole-blood partitioned for making the DIFF dilution is delivered to the DIFF bath using a tangential flow of 1.0 mL of Fix reagent (sample syringe, Fix reagent syringe). Mixing bubbles enter the bath.
- □ Sample probe moves from the DIFF bath to the rinse chamber (sample probe motor, traverse motor).
- **□** 5 μL of residual blood is expelled during the double rinsing (interior and exterior) of the sample probe (sample syringe, diluent reagent syringe).
- After 12 seconds of incubation, the dilution inside the DIFF bath is completed by adding another 1.0 mL of diluent which stops the cytochemical reaction (5diff syringe).

Making the DIFF dilution detailed:

- Sample probe is positioned opposite the entry port for the Fix reagent.
- Solenoid valve 8 is energized which also sets up the reagent syringes assembly movement to dispense 1.0 mL of Fix reagent.
- 25 µL of whole-blood sample is simultaneous delivered to the DIFF bath.
- Solenoid valve 8 de-energizes.
- DIFF dilution incubates for 12 seconds.
- Solenoid valves 1 and 3 are energized which sets up the 5diff syringe to dispense 1.0 mL of diluent is added to the DIFF bath to stop the cytochemical reaction.
- Solenoid valves 1 and 3 are de-energized.

Making the second dilution for RBC and PLT analysis:

- □ Sample probe moves from the rinse chamber the first dilution (DIL1/HGB) bath (sample probe motor, traverse motor).
- $\Box$  42.5 µL of the primary 1:170 dilution is aspirated (sample syringe).
- □ Sample probe exterior is rinsed (in the first dilution bath) with 0.4 mL of diluent (diluent reagent syringe).
- □ Sample probe moves from the first dilution (DIL1/HGB) bath to the RBC bath (sample probe motor, traverse motor).
- 42.5 µL of the first 1:170 dilution is delivered to the RBC bath using a tangential flow of 2.0 mL of diluent (sample syringe, diluent reagent syringe). The tangential flow of reagent mixes the 1:170 dilution sample and the diluent to make a uniform suspension of cells. Mixing bubbles also enter the bath. This is the final RBC/Plt dilution.
- □ An additional 0.5 mL of diluent is dispensed through the probe at the end of the second dilution.

Hemoglobin measurement:

Once the 42.5 µL of dilution is removed, the exterior of the sample probe is rinsed with 0.4 mL of diluent. 0.4 mL of Hgb Lyse reagent is then dispensed into the first dilution (DIL1/HGB) bath. Mixing bubbles enter the bath. The Hgb Lyse reagent destroys the red blood cells and converts a substantial proportion of the hemoglobin to a stable pigment so a hemoglobin value can be determined on the final 1:250 dilution.

#### **Count and Measurement of the WBC Group**

DiffPlot acquisition and WBC count:

- □ Loading of the vacuum injection circuit of the optical bench (waste syringe, energized solenoid valves 4 and 5).
- □ Start flow in the flow cell (approximately 2 seconds).
- □ Injection into the flow cell is 15 seconds but acquisition of data for the DiffPlot and WBC count occurs only 12 seconds.

Simultaneously, the WBC and BASO counts are gathered.

- Rinse of the counting head (the dilution pushed into position, solenoid valves 9, 10, 21, 23 and 13 are energized).
- □ Start the vacuum and first part of count (duration of 6 seconds, solenoid valve 23 energized).
- □ Rinsing of counting head and adjustment of the vacuum count.
- □ Second part of count (duration of 6 seconds).

After this first series of measurements:

- □ Measurement of the hemoglobin.
- □ Rinsing of the WBC/BASO bath with a mixture of 1.06 mL of Rinse reagent and 1.44 mL of diluent.
- □ Rinsing of the WBC/BASO counting head.

#### **RBC/PLT Group Count**

- □ Rinsing the RBC/PLT counting head.
- □ Start the vacuum for the RBC/PLT counting head.
- □ First series of counts (duration of 5 seconds, solenoid valve 14 energized).
- □ Rinsing of the RBC/PLT counting head and adjustment of the vacuum.
- □ Second series of counts (duration of 5 seconds).

#### Permanent Rinse Flow (PRF)

- □ The WBC/BASO and the RBC/PLT counting heads are *in series* during the RBC/PLT count. The count flow from the WBC bath circulates in the RBC/PLT counting head and limits the possibility of the circulation of the RBC behind the count orifice.
- □ The PRS functions similar to the autocleaning function in that it acts like a cleaning solution being drawn through the WBC/BASO aperture during the entire duration of the RBC and PLT counts.

#### **Filling of Diluent Reservoir**

- □ At the end of the RBC/PLT count, the RBC count vacuum is used for filling the diluent reservoir. Solenoid valves 15 and 25 are energized.
- □ To improve the filling speed, the syringe is initialized for a few seconds then drawn again to complete the filling.
- □ A float located in the tank controls the level of liquid.

#### **Completing the Cycle**

- □ To maximize mechanical resources, rinsing the RBC bath is normally programmed for the beginning of the succeeding cycle (which can be started during the RBC and PLT count).
- □ If no other cycle is started during the RBC/PLT count or during the filling of the diluent reservoir, a special rinsing sequence is initiated:
  - The sample signal light changes to red and the instrument is unavailable for a few seconds.
  - This sequence also involves draining of the RBC bath followed with a refilling with the diluent.

#### Wrap Up

- 1. Replace the left side panel. Secure the panel using the four hex screws removed earlier.
- 2. Close the right side door.
- 3. Make sure the right side door interlock is reactivated. See User Mode under Heading 4.1, as needed.
- 4. Turn the instrument off for 5 seconds then turn the instrument back on. An automatic Startup routine and background check is performed.
- 5. When the Startup and background checks are complete, verify the results are acceptable.

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- 6.2 PNEUMATIC / HYDRAULIC SCHEMATIC, 6.2-1 Layout, 6.2-1 Color Coding, 6.2-1 Tubing Designations, 6.2-1 Solenoid Valves, 6.2-1
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## **ILLUSTRATIONS**

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# CONTENTS

6

# 6.1 ENGINEERING SCHEMATIC DRAWINGS

This chapter contains a variety of troubleshooting aids including fold-out engineering schematics and interconnect diagrams. Details concerning the instrument's pneumatic/hydraulic tubing and connections are also provided.

Because the engineering schematics are insserted into the document and are not assigned page numbers or figure numbers, they are not included in the table of contents or the index. In the text, they are referenced by their name.

### **Engineering Schematics**

This chapter contains two schematics:

- Pneumatic/hydraulic schematic
- Interconnect schematic

The schematic revision levels were current on the date this revision of the manual was released. They will be updated again to the latest revision level whenever this manual is revised.

#### Additional Pneumatic/Hydraulic Information

#### **Tubings and Connectors**

In addition to the pneumatic/hydraulic schematic, a listing of the tubings and connectors used in the instrument are detailed in Table 6.3-1. This table also includes part numbers.

#### Pneumatic/Hydraulic Circuit Connections

Table 6.3-2 details the connections within each circuit including the diameter and length of tubing needed to make the connection. Tubing part numbers can be located on Table 6.3-1.

#### Additional Interconnect Information

Some components on the interconnect engineering schematic are detailed in diagrams. These interconnect diagrams are located under Heading 6.5.

SCHEMATICS ENGINEERING SCHEMATIC DRAWINGS

6.1-2

# 6.2 PNEUMATIC / HYDRAULIC SCHEMATIC

#### Layout

The pneumatic/hydraulic schematic for the  $A^{C} \cdot T$  5diff analyzer is laid out like the instrument. On the left side of the schematic are the assemblies you find in the left side compartment, including the count syringe, the 5diff syringe, the reagent syringes assembly, the flow cell, and the associated solenoid valves. The right side of the schematic shows the assemblies located behind the right side door, including the baths assembly, the diluent reservoir, the waste syringe, the sample syringe, and the sample probe and rinse block assembly.

#### **Color Coding**

The pneumatic/hydraulic schematic is color coded to correlate with the color used on the reagent labels. The arrangement of reagents bottles correlates with their arrangement inside the reagent compartment.

- A<sup>C</sup>•T 5diff Rinse lines are blue.
- A<sup>C</sup>•T 5diff Fix reagent lines are yellow.
- A<sup>C</sup>•T 5diff Hgb Lyse reagent lines are orange.
- A<sup>C</sup>•T 5diff WBC Lyse reagent lines are green.
- The A<sup>C</sup>•T 5diff Diluent reagent lines are magenta.
- All waste lines are color-coded red.

#### **Tubing Designations**

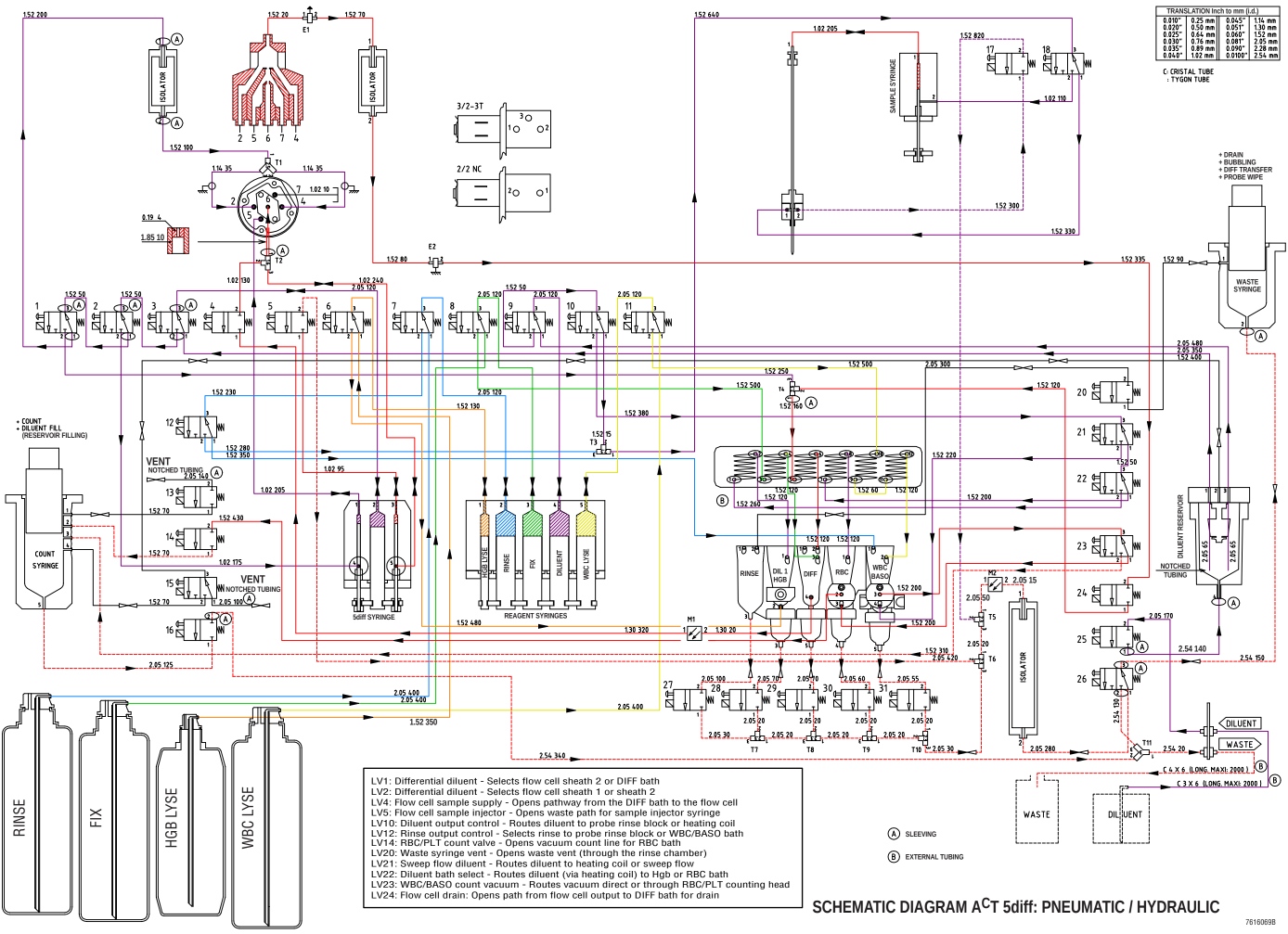
All tubing on this instrument is considered critical length. Notice there are two numbers associated with each tubing. The first number is the internal diameter of the tubing in millimeters; the second, is the length of the tubing in millimeters. For example, locate the count syringe. Notice the tubing attached to the bottom fitting also attaches to port 1 of solenoid valve 16. The number 2.05 125 is displayed above that line. This number indicates the tubing has a 2.05 mm diameter and is 125 mm long. A box containing a translation of inches to millimeter is located In the upper right corner. Although you may find this helpful, it does not contain all tubing measurements. The numbers will be close but, as in our example, not exact. Since the tubing is critical length, you may want to have a metric ruler available.

Most of the tubing used on this instrument is Tygon tubing. The legend in the upper right corner lists a second tubing, Cristal tubing which is used in limited quantities. In the lower right corner of the schematic, notice the waste container and the diluent container both use Cristal tubing. The Cristal tubing is designated by the letter C. The tubing used for the waste line has a larger diameter (4 mm) than the tubing used for the diluent (3 mm). These tubings are not interchangeable.

#### **Solenoid Valves**

Two types of solenoid valves are used on this instrument, a normally-closed two-way valve and a three-way valve. Inside the box to the right of the reagent bottle, LV designates the electronic valve or solenoid valve. Valve numbers may also be preceded by a V, an S, and an SL designation.

SCHEMATICS PNEUMATIC / HYDRAULIC SCHEMATIC





6

## 6.3 PNEUMATIC/HYDRAULIC TUBINGS AND CONNECTIONS

## **Tubings and Connectors List**

The tubings and connectors used in the A<sup>C</sup>•T 5diff hematology analyzer are detailed in Table 6.3-1. This table also includes part numbers.

#### Table 6.3-1 Instrument Tubing and Connectors

| Designation                      | Part Number | Diameter | Length | Quantity |
|----------------------------------|-------------|----------|--------|----------|
| Y-connector                      | EAB021A     | 3        |        | 1        |
| Y-connector                      | EAB026A     | 2.5      |        | 1        |
| T410 connector                   | EAB033A     | 1.6      |        | 1        |
| T220 connector                   | EAB035A     | 2.3      |        | 8        |
| 0.040" Tygon <sup>®</sup> tubing | EAE005A     | 1.02     | 10     | 1        |
| 0.040" Tygon tubing              | EAE005A     | 1.02     | 95     | 1        |
| 0.040" Tygon tubing              | EAE005A     | 1.02     | 110    | 1        |
| 0.040" Tygon tubing              | EAE005A     | 1.02     | 130    | 1        |
| 0.040" Tygon tubing              | EAE005A     | 1.02     | 175    | 1        |
| 0.040" Tygon tubing              | EAE005A     | 1.02     | 205    | 2        |
| 0.040" Tygon tubing              | EAE005A     | 1.02     | 240    | 1        |
| 0.051" Tygon tubing              | EAE006A     | 1.30     | 20     | 1        |
| 0.051" Tygon tubing              | EAE006A     | 1.30     | 140    | 1        |
| 0.051" Tygon tubing              | EAE006A     | 1.30     | 320    | 1        |
| 0.060" Tygon tubing              | EAE007A     | 1.52     | 15     | 1        |
| 0.060" Tygon tubing              | EAE007A     | 1.52     | 20     | 1        |
| 0.060" Tygon tubing              | EAE007A     | 1.52     | 50     | 4        |
| 0.060" Tygon tubing              | EAE007A     | 1.52     | 60     | 1        |
| 0.060" Tygon tubing              | EAE007A     | 1.52     | 70     | 4        |
| 0.060" Tygon tubing              | EAE007A     | 1.52     | 80     | 1        |
| 0.060" Tygon tubing              | EAE007A     | 1.52     | 90     | 1        |
| 0.060" Tygon tubing              | EAE007A     | 1.52     | 100    | 1        |
| 0.060" Tygon tubing              | EAE007A     | 1.52     | 120    | 6        |
| 0.060" Tygon tubing              | EAE007A     | 1.52     | 130    | 1        |
| 0.060" Tygon tubing              | EAE007A     | 1.52     | 160    | 1        |
| 0.060" Tygon tubing              | EAE007A     | 1.52     | 200    | 4        |
| 0.060" Tygon tubing              | EAE007A     | 1.52     | 220    | 1        |
| 0.060" Tygon tubing              | EAE007A     | 1.52     | 230    | 1        |
| 0.060" Tygon tubing              | EAE007A     | 1.52     | 250    | 2        |
| 0.060" Tygon tubing              | EAE007A     | 1.52     | 280    | 1        |

| Designation         | Part Number | Diameter | Length | Quantity |
|---------------------|-------------|----------|--------|----------|
| 0.060" Tygon tubing | EAE007A     | 1.52     | 300    | 1        |
| 0.060" Tygon tubing | EAE007A     | 1.52     | 310    | 1        |
| 0.060" Tygon tubing | EAE007A     | 1.52     | 330    | 1        |
| 0.060" Tygon tubing | EAE007A     | 1.52     | 335    | 1        |
| 0.060" Tygon tubing | EAE007A     | 1.52     | 350    | 2        |
| 0.060" Tygon tubing | EAE007A     | 1.52     | 380    | 1        |
| 0.060" Tygon tubing | EAE007A     | 1.52     | 400    | 1        |
| 0.060" Tygon tubing | EAE007A     | 1.52     | 430    | 1        |
| 0.060" Tygon tubing | EAE007A     | 1.52     | 480    | 1        |
| 0.060" Tygon tubing | EAE007A     | 1.52     | 500    | 2        |
| 0.060" Tygon tubing | EAE007A     | 1.52     | 640    | 1        |
| 0.060" Tygon tubing | EAE007A     | 1.52     | 820    | 1        |
| 0.081" Tygon tubing | EAE008A     | 2.05     | 15     | 1        |
| 0.081" Tygon tubing | EAE008A     | 2.05     | 20     | 8        |
| 0.081" Tygon tubing | EAE008A     | 2.05     | 30     | 2        |
| 0.081" Tygon tubing | EAE008A     | 2.05     | 50     | 1        |
| 0.081" Tygon tubing | EAE008A     | 2.05     | 55     | 1        |
| 0.081" Tygon tubing | EAE008A     | 2.05     | 60     | 1        |
| 0.081" Tygon tubing | EAE008A     | 2.05     | 65     | 2        |
| 0.081" Tygon tubing | EAE008A     | 2.05     | 70     | 2        |
| 0.081" Tygon tubing | EAE008A     | 2.05     | 100    | 1        |
| 0.081" Tygon tubing | EAE008A     | 2.05     | 100    | 1        |
| 0.081" Tygon tubing | EAE008A     | 2.05     | 120    | 5        |
| 0.081" Tygon tubing | EAE008A     | 2.05     | 125    | 1        |
| 0.081" Tygon tubing | EAE008A     | 2.05     | 140    | 1        |
| 0.081" Tygon tubing | EAE008A     | 2.05     | 170    | 1        |
| 0.081" Tygon tubing | EAE008A     | 2.05     | 280    | 1        |
| 0.081" Tygon tubing | EAE008A     | 2.05     | 300    | 1        |
| 0.081" Tygon tubing | EAE008A     | 2.05     | 350    | 1        |
| 0.081" Tygon tubing | EAE008A     | 2.05     | 400    | 3        |
| 0.081" Tygon tubing | EAE008A     | 2.05     | 420    | 1        |
| 0.081" Tygon tubing | EAE008A     | 2.05     | 480    | 1        |
| 0.045" Tygon tubing | EAE033A     | 1.14     | 35     | 2        |
| 0.100" Tygon tubing | EAE034A     | 2.54     | 20     | 1        |
| 0.100" Tygon tubing | EAE034A     | 2.54     | 130    | 1        |

Table 6.3-1 Instrument Tubing and Connectors (Continued)

I

|

I

| Designation          | Part Number | Diameter | Length | Quantity |
|----------------------|-------------|----------|--------|----------|
| 0.100" Tygon tubing  | EAE034A     | 2.54     | 140    | 1        |
| 0.100" Tygon tubing  | EAE034A     | 2.54     | 150    | 1        |
| 0.100" Tygon tubing  | EAE034A     | 2.54     | 340    | 1        |
| 0.0075" Tygon tubing | EAE047A     | 0.19     | 4      | 1        |
| 0.073" Tygon tubing  | EAE049A     | 1.85     | 10     | 1        |
| Sleeving             | GAL098A     |          |        | 27       |
| Tube shielding       | XBA403A     |          |        | 1        |

 Table 6.3-1 Instrument Tubing and Connectors (Continued)

## **Pneumatic/Hydraulic Circuit Connections**

In this table, each circuit connection has several descriptions. As you review the table, you will notice that many of the descriptions have a name or number, followed by a line, followed by a number. LV13\_1 is a typical example of this format. LV13\_1 is read as port 1 of valve 13 or valve 13, port 1.

| Circuit | From                            | F.S. | Diameter | Length | T.S. | То                  |
|---------|---------------------------------|------|----------|--------|------|---------------------|
| AIR     | Atmosphere                      |      | 2.05     | 140    |      | LV13_2              |
|         | LV13_1                          |      | 1.52     | 70     |      | Count syringe_1     |
|         | Atmosphere                      |      | 2.05     | 100    |      | LV15_1              |
|         | LV15_3                          |      | 1.52     | 400    |      | Diluent reservoir_2 |
|         | LV15_2                          |      | 1.52     | 70     |      | Count syringe_4     |
|         | Rinse_2 chamber<br>(Atmosphere) |      | 2.05     | 300    |      | LV20_2              |
|         | LV20_1                          |      | 1.52     | 90     |      | Waste syringe_1     |

Table 6.3-2 Circuit Connections

I

|

| Circuit         | From                | F.S. | Diameter | Length | T.S. | То                  |
|-----------------|---------------------|------|----------|--------|------|---------------------|
| DILUENT REAGENT | Diluent Input       |      | 2.05     | 170    |      | LV25_2              |
|                 | LV25_1              | S    | 2.54     | 140    | S    | Diluent reservoir_4 |
|                 |                     |      | 2.05     | 65     |      | Diluent reservoir_1 |
|                 | Diluent reservoir_1 |      | 2.05     | 350    |      | LV3_1               |
|                 | LV3_3               | S    | 2.05     | 120    |      | Injector syringe_2  |
|                 | LV3_2               | S    | 1.52     | 50     | S    | LV2_3               |
|                 | LV2_1               | S    | 1.52     | 50     | S    | LV1_3               |
|                 | LV1_1               | S    | 1.52     | 200    | S    | Isolator_1          |
|                 | Isolator_2          | S    | 1.52     | 100    |      | T1_1                |
|                 | T1_2                |      | 1.14     | 35     |      | DIFF flow cell_4    |
|                 | T1_2                |      | xba403a  |        |      | DIFF flow cell_4    |
|                 | T1_3                |      | 1.14     | 35     |      | DIFF flow cell_2    |
|                 | T1_3                |      | xba403a  |        |      | DIFF flow cell_2    |
|                 | LV1_2               |      | 1.52     | 250    |      | T4_3                |
|                 | LV2_2               |      | 1.02     | 175    |      | Injector syringe_4  |
|                 | Injector syringe_1  |      | 1.02     | 205    |      | DIFF flow cell_5    |
|                 | DIFF flow cell_7    |      | 1.02     | 10     |      | (Cap)               |
|                 |                     |      | 2.05     | 65     |      | Diluent reservoir_3 |
|                 | Diluent reservoir_3 |      | 2.05     | 480    |      | LV9_1               |
|                 | LV9_3               |      | 2.05     | 120    |      | Reagent syringe_4   |
|                 | LV9_2               |      | 1.52     | 50     |      | LV10_3              |
|                 | LV10_1              |      | 1.52     | 15     |      | T3_2                |
|                 | T3_1                |      | 1.52     | 640    |      | LV18_3              |
|                 | LV18_1              |      | 1.52     | 330    |      | Probe rinse block_1 |
|                 | Probe rinse block_2 |      | 1.52     | 300    |      | LV17_1              |
|                 | LV17_2              |      | 1.52     | 820    |      | T5_2                |
|                 | LV18_2              |      | 1.02     | 110    |      | Sample syringe_2    |
|                 | LV10_2              |      | 1.52     | 380    |      | LV21_3              |
|                 | LV21_1              |      | 1.52     | 50     |      |                     |
|                 | LV22_1              |      | 1.52     | 200    | S    | Reagent heater_7    |
|                 | Reagent heater_8    | S    | 1.52     | 120    |      | RBC bath_1          |
|                 | LV22_2              |      | 1.52     | 250    | S    | Reagent heater_1    |
|                 | Reagent heater_2    | S    | 1.52     | 120    |      | DIL1/HGB bath_1     |
|                 | LV21_2              |      | 1.52     | 220    |      | WBC/BASO bath_4     |

Table 6.3-2 Circuit Connections (Continued)

I

| Circuit          | From                    | F.S. | Diameter | Length | T.S. | То                  |
|------------------|-------------------------|------|----------|--------|------|---------------------|
| RINSE REAGENT    | Rinse reagent bottle    |      | 2.05     | 400    |      | LV7_1               |
|                  | LV7_3                   |      | 2.05     | 120    |      | Reagent syringe_2   |
|                  | LV7_2                   |      | 1.52     | 230    |      | LV12_3              |
|                  | LV12_1                  |      | 1.52     | 350    |      | WBC/BASO bath_1     |
|                  | LV12_2                  |      | 1.52     | 280    |      | T3_3                |
| FIX REAGENT      | Fix reagent bottle      |      | 2.05     | 400    |      | LV8_1               |
|                  | LV8_3                   |      | 2.05     | 120    |      | Reagent syringe_3   |
|                  | LV8_2                   |      | 1.52     | 500    | S    | Reagent heater_3    |
|                  | Reagent heater_4        | S    | 1.52     | 120    |      | DIFF bath_3         |
| WBC LYSE REAGENT | WBC Lyse reagent bottle |      | 2.05     | 400    |      | LV11_1              |
|                  | LV11_3                  |      | 2.05     | 120    |      | Reagent syringe_5   |
|                  | LV11_2                  |      | 1.52     | 500    | S    | Reagent heater_10   |
|                  | Reagent heater_9        | S    | 1.52     | 60     | S    | Reagent heater_11   |
|                  | Reagent heater_12       | S    | 1.52     | 120    |      | WBC/BASO bath_2     |
| HGB LYSE REAGENT | Hgb Lyse reagent bottle |      | 1.52     | 350    |      | LV6_1               |
|                  | LV6_3                   |      | 1.52     | 130    |      | Reagent syringe_1   |
|                  | LV6_2                   |      | 1.52     | 480    |      | DIL1/HGB bath_2     |
| SAMPLING         | Probe_1                 |      | 1.02     | 205    |      | Sample syringe_1    |
| DIFF COUNTING    | DIFF bath_4             |      | 1.30     | 20     |      | M1_2 photocell      |
|                  | M1_1 photocell          |      | 1.30     | 320    |      | LV4_1               |
|                  | LV4_2                   |      | 1.02     | 130    |      | T2_2                |
|                  | T2_3                    |      | 1.02     | 240    |      | Injector syringe_5  |
|                  | Injector syringe_3      |      | 1.02     | 95     |      | LV5_1               |
|                  | LV5_2                   |      | 2.05     | 420    |      | T6_2                |
|                  | T2_1                    | S    | 1.85     | 10     |      | DIFF flow cell_6    |
|                  |                         |      | 0.19     | 4      |      | DIFF flow cell_6    |
|                  | DIFF flow cell_output   |      | 1.52     | 20     |      | E1_1 anode fitting  |
|                  | E1_2 anode fitting      |      | 1.52     | 70     |      | Isolator_1          |
|                  | Isolator_2              |      | 1.52     | 80     |      | E2_1 ground fitting |
|                  | E2_2 ground fitting     |      | 1.52     | 335    |      | LV24_2              |
|                  | LV24_1                  |      | 1.52     | 120    |      | T4_2                |
|                  | T4_1                    |      | 1.52     | 160    | S    | Reagent heater_5    |
|                  | Reagent heater_6        | S    | 1.52     | 120    |      | DIFF bath_2         |

Table 6.3-2 Circuit Connections (Continued)

I

I

| Circuit          | From            | F.S. | Diameter | Length | T.S. | То              |
|------------------|-----------------|------|----------|--------|------|-----------------|
| WBC/RBC COUNTING | WBC/BASO bath_3 |      | 1.52     | 200    |      | LV23_3          |
|                  | LV23_1          |      | 1.52     | 200    |      | RBC bath_3      |
|                  | RBC bath_2      |      | 1.52     | 430    |      | LV14_2          |
|                  | LV14_1          |      | 1.52     | 70     |      | Count syringe_2 |
|                  | LV23_2          |      | 1.52     | 310    |      | Count syringe_3 |
| WASTE            | Rinse chamber_3 |      | 2.05     | 100    |      | LV27_2          |
|                  | LV27_1          |      | 2.05     | 30     |      | T7_3            |
|                  | DIL1/HGB bath_3 |      | 2.05     | 70     |      | LV28_2          |
|                  | LV28_1          |      | 2.05     | 20     |      | T7_2            |
|                  | T7_1            |      | 2.05     | 20     |      | T8_3            |
|                  | DIFF bath_5     |      | 2.05     | 70     |      | LV29_2          |
|                  | LV29_1          |      | 2.05     | 20     |      | T8_2            |
|                  | T8_1            |      | 2.05     | 20     |      | T9_3            |
|                  | RBC bath_4      |      | 2.05     | 60     |      | LV30_2          |
|                  | LV30_1          |      | 2.05     | 20     |      | T9_2            |
|                  | T9_1            |      | 2.05     | 20     |      | T10_2           |
|                  | WBC/BASO bath_5 |      | 2.05     | 55     |      | LV31_2          |
|                  | LV31_1          |      | 2.05     | 20     |      | T10_1           |
|                  | T10_3           |      | 2.05     | 30     |      | T6_3            |
|                  | T6_1            |      | 2.05     | 20     |      | T5_3            |
|                  | T5_1            |      | 2.05     | 50     |      | M2_1 photocell  |
|                  | M2_2 photocell  |      | 2.05     | 15     |      | Isolator_1      |
|                  | Isolator_2      |      | 2.05     | 280    |      | LV26_1          |
|                  | LV26_3          | S    | 2.54     | 150    | S    | Waste syringe_2 |
|                  | LV26_2          | S    | 2.54     | 130    |      | T11_3           |
|                  | Count syringe_5 |      | 2.05     | 125    | 1    | LV16_1          |
|                  | LV16_2          | S    | 2.54     | 340    |      | T11_2           |
|                  | T11_1           |      | 2.54     | 20     |      | Waste output    |

Table 6.3-2 Circuit Connections (Continued)

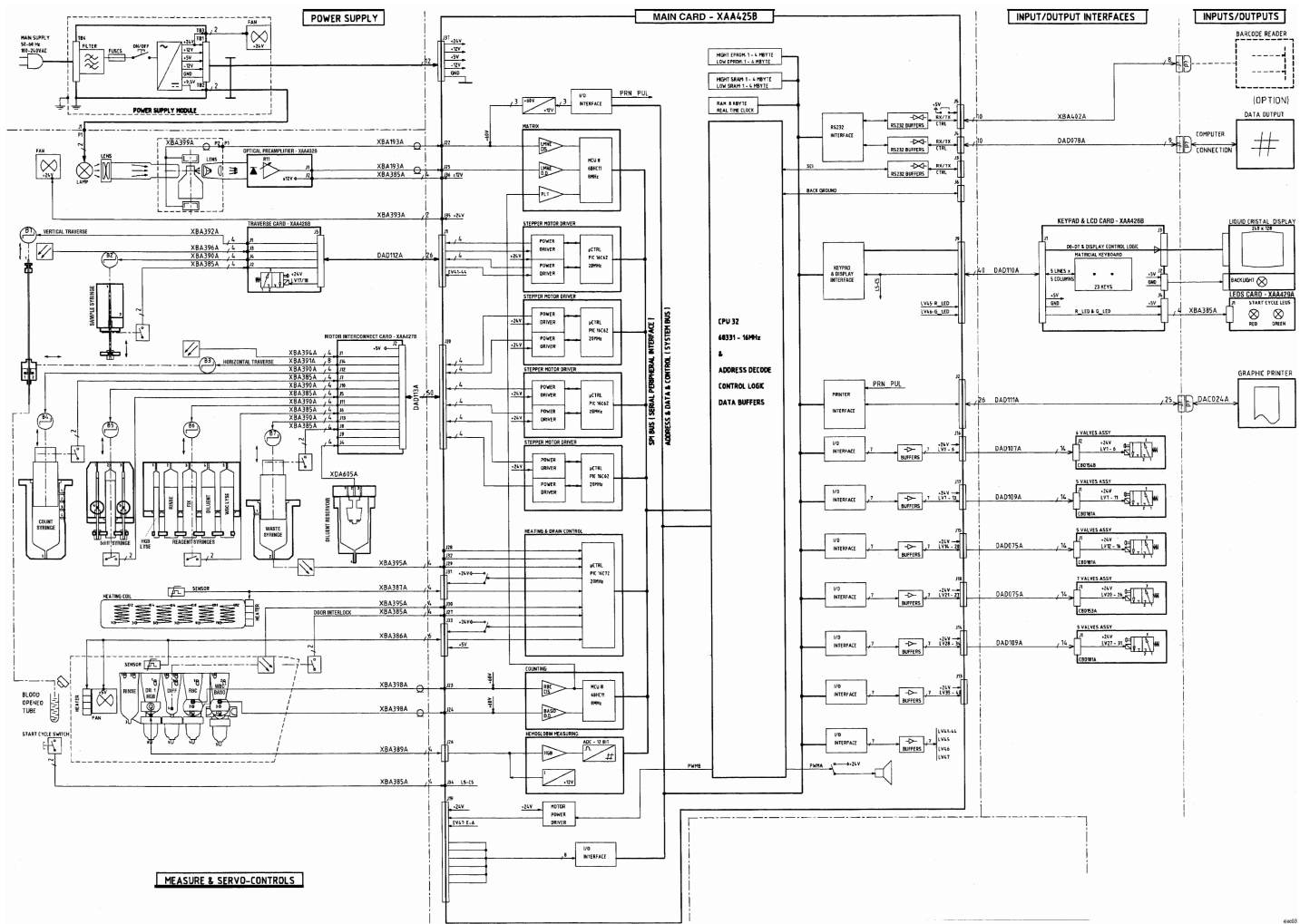
# 6.4 INTERCONNECT DIAGRAM

The fold-out interconnect diagram for the A<sup>C</sup>•T 5diff analyzer is divided into five sections using a dash/dot line. These five sections include:

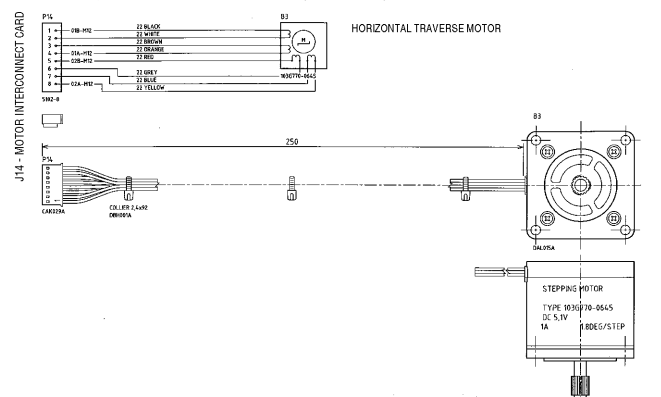
- Power Supply module in the upper left corner.
- Measurement components on the left, under the Power Supply module.
- Main card in the center.
- Input/output interfaces to the right of the Main card.
- System inputs and outputs.

This interconnect diagram is a helpful troubleshooting tool, especially when troubleshooting an electronic problem.

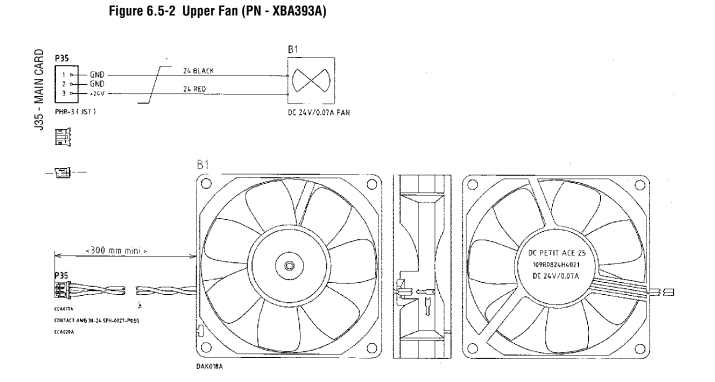
SCHEMATICS INTERCONNECT DIAGRAM

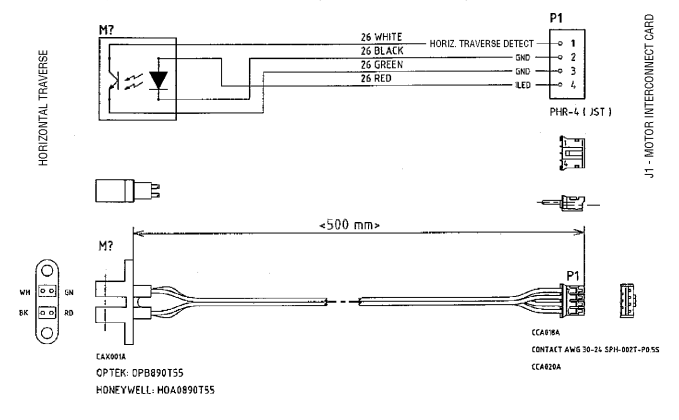


# 6.5 MOTORS AND CABLES



#### Figure 6.5-1 Horizontal Traverse Motor (PN - XBA391A)





#### Figure 6.5-3 Horizontal Traverse Sensor (IR Sensor, PN - XBA394AS)

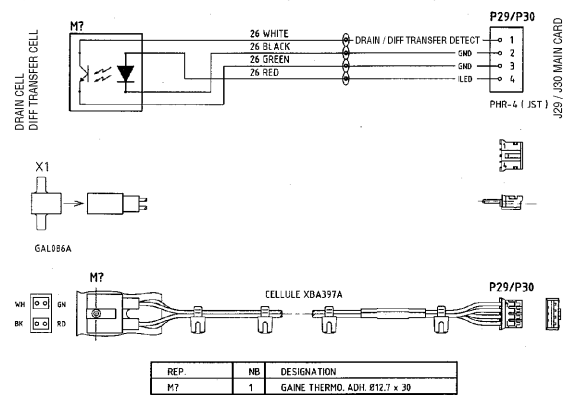
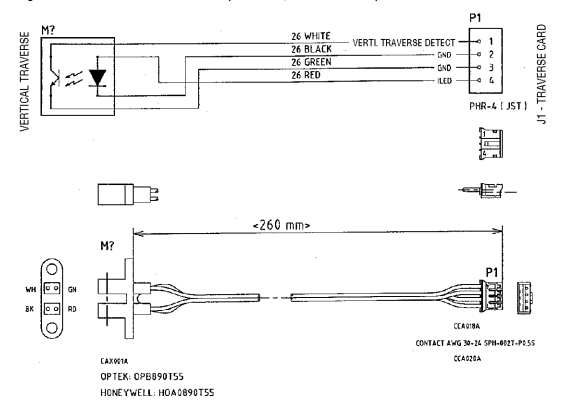
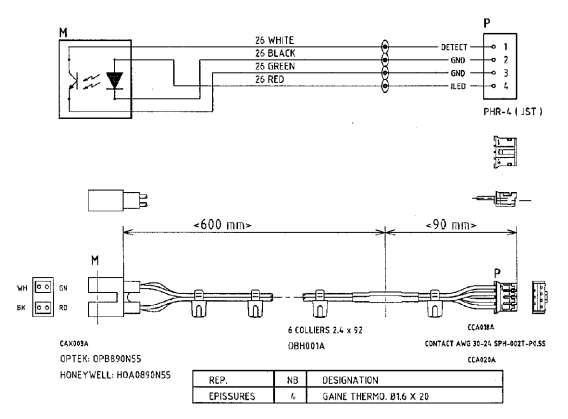


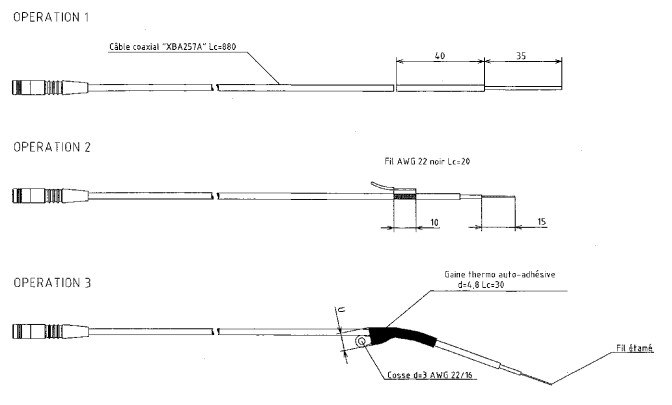
Figure 6.5-4 Bath Drain and DIFF Transfer Sensor (IR Sensor, PN - XBA395AS)

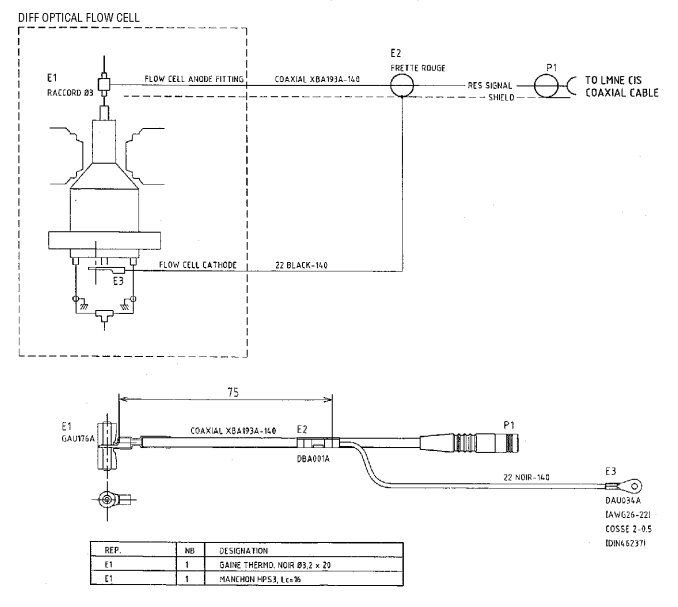


#### Figure 6.5-5 Vertical Traverse Sensor (IR Sensor, PN - XBA396AS)

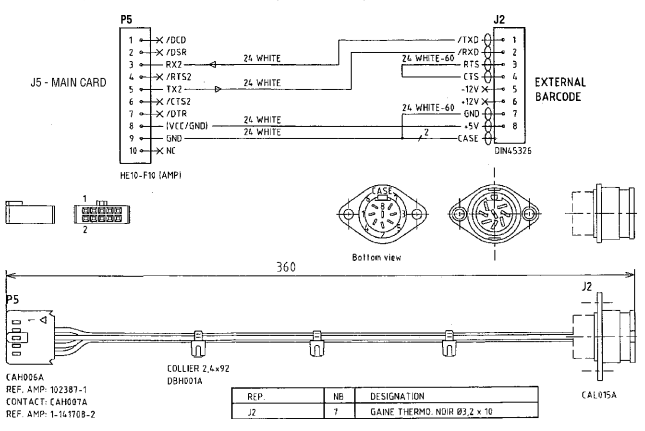






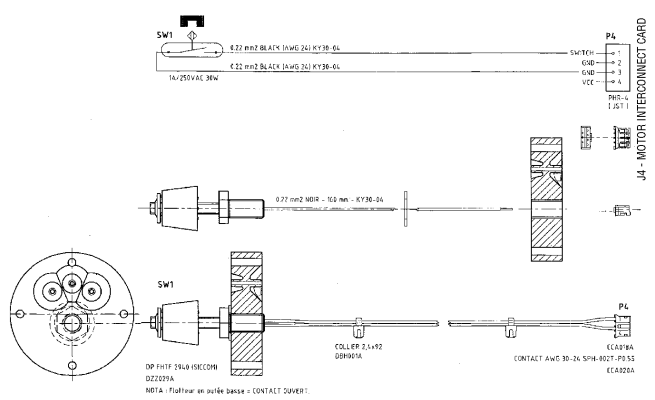


#### Figure 6.5-8 DIFF Flow Cell Coaxial Cable (PN - XBA399AS)



#### Figure 6.5-9 Bar-Code Reader Cable (PN - XBA402AS)

### Figure 6.5-10 Diluent Level Sensor (PN - XDA605AS)



- 7 TROUBLESHOOTING, 7.1-1
  - 7.1 ERROR MESSAGES, 7.1-1
  - 7.2 CHECKING THE MOTORS, 7.2-1 Purpose, 7.2-1 Tools/Supplies Needed, 7.2-1 Preparation, 7.2-2 To Check All Motors, 7.2-2 To Only Check Motors in the Right Side Compartment, 7.2-2 To Only Check Motors in the Left Side Compartment, 7.2-2 Motors Check, 7.2-3
  - 7.3 CHECKING THE VALVES, 7.3-1

Purpose, 7.3-1 Tools/Supplies Needed, 7.3-1 Preparation, 7.3-1 To Only Check Valves in the Left Side Compartment, 7.3-1 To Only Check Valves in the Right Side Compartment, 7.3-1 Valves Check, 7.3-2

Wrap Up, 7.3-3

## **ILLUSTRATIONS**

- 7.3-1 Valve 1 through 16 Locations, 7.3-2
- 7.3-2 Valves 17 and 18 Location, 7.3-2
- 7.3-3 Valves 20 through 31 Locations, 7.3-2

## TABLES

7.1-1 Error Messages, 7.1-1

# CONTENTS

# 7.1 ERROR MESSAGES

## Table 7.1-1 Error Messages

| Displayed Message                             | ayed Message Problem  |   |
|---|---|---|
| AT LEAST 3 TAGGED RESULTS<br>REQUIRED         | At least 3 results are required for calibration calculations and less than 3 have been run.   | Run at least three results for calculation results to be generated.   |
| BATH ENCLOSURE DOOR OPENED                    | If a cycle is attempted while the right side door is open, this message is generated.   | Close door to continue.   |
| DATA NOT SAVED, VALUE OUT OF<br>RANGE         | The value typed in is not an acceptable value. It may be out of an expected range or an incorrect data type.                                  | Re-enter the data   |
| DRAIN TIMEOUT                                 | Draining problems.  | <ol> <li>Run a System Reset cycle.</li> <li>If the problem persists, call your<br/>Beckman Coulter Representative.</li> </ol>             |
| ENTER AN IDENTIFICATION                       | An ID is required to run an analysis in the Manual ID mode.   | Enter the specimen ID.  |
| INCORRECT DATE ENTRY                          | The value entered is not a valid date.  | Enter a valid date.   |
| INCORRECT TIME ENTRY                          | The time entered is not a valid time.   | Enter a valid time.   |
| <i>NO ACK CHARACTER RECEIVED ON<br/>RS232</i> | There is a problem with the communication or handshaking to the host computer.  | Check that the protocol that has been<br>set up in the host transmission screen<br>matches the protocol expected by the<br>host computer. |
| NO DILUENT, CHECK LEVEL                       | The diluent reservoir is not able to fill.  | Check the diluent level and replace diluent if necessary.   |
| <i>NO ENQ CHARACTER RECEIVED ON<br/>RS232</i> | There is a problem with the communication or handshaking to the host computer.  | Check that the protocol that has been<br>setup in the host transmission screen<br>matches the protocol expected by the<br>host computer.  |
| PRINTER ERROR, CHECK PAPER                    | An error indication has been sent from<br>the Printer to the instrument, usually a<br>paper out message.                                      | <ol> <li>Ensure there is paper in the Printer.</li> <li>Check the Printer user's manual for<br/>other Printer errors.</li> </ol>          |
| REAGENT LOW LEVEL [REAGENT<br>NAME]           | The calculated reagent level for the specified reagent indicates no reagent.  | <ol> <li>Check the reagent level and replace<br/>the reagent if necessary.</li> <li>Update the reagent level.</li> </ol>                  |
| REAGENTS LOW LEVEL                            | This message is given at the end of<br>startup if there is not enough reagent<br>left to complete the daily workload that<br>has been set up. | Monitor the reagent levels, or change the reagents and update the levels.   |

| Displayed Message   | Problem   | Corrective Action  |
|---|---|--|
| SYSTEM ERROR, RUN SYSTEM<br>RESET CYCLE                                   | <ul> <li>During a cycle, a system error of the following type has caused the system to stop:</li> <li>A motor has not returned to its home sensor when expected.</li> <li>A drain problem has been detected at one of the two drain sensors.</li> <li>The right side door has been opened during a cycle, losing temperature control at the baths.</li> </ul> | <ol> <li>Check the specified motor to make<br/>sure it is not jammed, and there are<br/>no signs of physical damage to<br/>motor mechanism or sensor.</li> <li>Check the bath compartment for<br/>leaks, plugs, loose tubing. Ensure<br/>that there is sufficient reagent and<br/>that the system is primed.</li> <li>Ensure that there is no physical<br/>damage to the microswitch that<br/>senses closure of the right side<br/>door and that the door is<br/>completely closed.</li> <li>Run an System Reset cycle after<br/>checking/correcting any<br/>instrument problems.</li> </ol> |
| TEMPERATURE OUT OF RANGE  | The temperature in the counting bath compartment is outside of the acceptable range.  | <ol> <li>Ensure the sure right side door is<br/>closed.</li> <li>Wait a few minutes.</li> <li>If problem persists, call your<br/>Beckman Coulter Representative.</li> </ol>  |
| THE PRINTER IS DISCONNECTED,<br>SWITCHED OFF, OR HAS NOT BEEN<br>SELECTED | No or incorrect communication<br>between Printer and instrument   | <ol> <li>Ensure the cable is properly<br/>connected.</li> <li>Ensure the Printer is turned on.</li> <li>Ensure the Printer is online or<br/>selected.</li> </ol>   |
| TIMEOUT OVERFLOW ON RS232   | There is a problem with the communication or handshaking to the host computer.  | Check that the protocol that has been<br>setup in the host transmission screen<br>matches the protocol expected by the<br>host computer.   |
| USER PASSWORD   | A password is required to perform the requested action.   | Enter user password.   |
| WRITE ERROR ON RS232  | There is a problem with the communication or handshaking to the host computer.  | Check that the protocol that has been<br>set up in the host transmission screen<br>matches the protocol expected by the<br>host computer.  |
| XXX NOT REACHING HOME<br>Note: XXX = name of motor.                       | Motor did not reach home sensor.  |  |

Table 7.1-1 Error Messages (Continued)

# 7.2 CHECKING THE MOTORS

#### Purpose

Use this procedure to individually verify motor operation is smooth and regular for one or more of seven different stepper motors including the:

**Note:** If you need help locating the various assemblies, see Figures A.3-1 and A.3-2. These figures show the assembly the motor controls, not the actual motor being checked.

- Sample probe motor. This motor is responsible for moving the sample probe up and down. This is also called the vertical traverse motor. You can verify operation of this motor without opening any compartments.
- **Traverse motor**. This motor is responsible for the left/right movement of the sample probe (inside its housing) over the baths assembly. This motor is also called the horizontal traverse motor. You need to access the right side compartment to verify operation of this motor.
- **Sample syringe motor.** This motor is responsible for aspirating, partitioning, and delivering the proper volume of sample. You need to access the right side compartment to verify operation of this motor by observing the sample syringe piston movement.
- Waste syringe motor. This motor is responsible for the draining the baths and various chambers. It also provides mixing bubbles and the vacuum needed to pull the DIFF specimen from the DIFF bath towards the flow cell injector syringe. You need to access the right side compartment to verify operation of this motor by observing waste syringe piston movement.
- **Count syringe motor.** This motor is responsible for supplying the aperture vacuum needed for counting as well as the vacuum needed to fill the diluent reservoir. You need to access the left side compartment to verify operation of this motor by observing count syringe piston movement.
- **5diff syringe motor**. This motor is responsible for the correct proportioning of the stop diluent in the DIFF bath and for injecting the sample into the flow cell. You need to access the left side compartment to verify operation of this motor by observing 5diff syringe piston movement.
- **Reagent syringes motor.** This motor is responsible for the correct distribution of the different reagents including Hgb Lyse, Rinse, Fix, Diluent, and WBC Lyse. You need to access the left side compartment to verify operation of this motor by observing the piston movements of the reagent syringes.

#### **Tools/Supplies Needed**

Allen wrenches, 2.5 mm and 3.0 mm , may be required.

## Preparation

Determine which motor(s) you want to check so you know which door to open or panel to remove.

#### **To Check All Motors**

- 1. Open the right side door.
- 2. Remove the left side panel and set it aside. For details, see Heading 4.2.

**CAUTION** A flat cable is connected to the rear of the Main card. Be careful when you open the door (where the Main card is attached) that you do not disconnect or damage this cable.

- 3. In the left side compartment,
  - a. Turn the two captive knobs counterclockwise to release the Main card door. These knobs are located to the right of the Main card.
  - b. Open the Main card door and carefully anchor the door behind the white plastic catch to keep it open.
- 4. Go to the Motors Check heading that follows.

#### To Only Check Motors in the Right Side Compartment

To check the sample probe, traverse assembly, sample syringe, or waste syringe motor:

- 1. Open the right side door of the instrument.
- 2. Go to the Motors Check heading that follows.

#### To Only Check Motors in the Left Side Compartment

To check the count syringe, 5diff syringe, or reagent syringes motor:

1. Remove the left side panel and set it aside. For details, see Heading 4.2.

**CAUTION** A flat cable is connected to the rear of the Main card. Be careful when you open the door (where the Main card is attached) that you do not disconnect or damage this cable.

- 2. In the left side compartment,
  - a. Turn the two captive knobs counterclockwise to release the Main card door. These knobs are located to the right of the Main card.
  - b. Open the Main card door and carefully anchor the door behind the white plastic catch to keep it open.
- 3. Go to the Motors Check heading that follows.

#### **Motors Check**

1. From the Main Menu, select **DIAGNOSTICS >> HARDWARE SYSTEMS >> MOTORS**.

Note: Motors must be checked individually. If more than one motor needs checked, select one motor and verify its operation before proceeding to the next.

- 2. From the Motors menu, select the motor you want to check:
  - 1. SAMPLING PROBE (same as Vertical Traverse or Sample Probe)
  - 2. TRAVERSE (same as Horizontal Traverse)
  - 3. SAMPLING SYRINGE (same as Sample Syringe)
  - 4. DRAINING SYRINGE (same as Waste Syringe)
  - 5. COUNTING SYRINGE (same as Count Syringe
  - 6. FLOWCELL SYRINGES (same as 5diff Syringe)
  - 7. DILUTION SYRINGES (same as Reagent Syringes assembly)
- 3. Verify the motor is operating smoothly and regularly by observing the movement of the respective components. See Figures A.3-1 and A.3-2 if you need help locating these assemblies.
  - If checking the **SAMPLING PROBE** motor, make sure the sample probe is moving up and down properly (vertical traverse movement check).
  - If checking the **TRAVERSE** motor, make sure the left/right movement of the sample probe (inside its housing) over the baths assembly is correct (horizontal traverse movement check).
  - If checking the **SAMPLING SYRINGE** motor, make sure the sample syringe piston is moving up and down properly.
  - If checking the **DRAINING SYRINGE** motor, make sure the waste syringe piston is moving up and down properly.
  - If checking the **COUNTING SYRINGE** motor, make sure the count syringe piston is moving up and down properly.
  - If checking the **FLOWCELL SYRINGE** motor, make sure the 5diff syringe piston is moving up and down properly.
  - If checking the **DILUTION SYRINGES** motor, make sure all five syringe pistons in the reagent syringes assembly are moving up and down properly.

#### Wrap Up

When motor performance is acceptable, close all doors and panels:

- 1. If the right side compartment was opened, close the right side door.
- 2. If the left side compartment was opened,
  - a. Close the Main card door. Turn the two captive knobs clockwise to secure the door.

**ATTENTION:** When replacing the left side panel with the instrument powered on, avoid accidently turning the instrument off again by carefully positioning the opening for the power on/off rocker switch over the switch as you position the panel on the instrument frame.

b. Replace the left side panel and install the four hex screws that secure it to the instrument frame.

# 7.3 CHECKING THE VALVES

#### Purpose

Use this procedure to verify a valve's operation is smooth and regular. When activated, all the valves in the assembly should be activated. There are five valve assemblies in the instrument including:

- One 11-valve assembly (valve numbers 1 through 11) located in the left side compartment.
- One 5-valve assembly (valve numbers 12 through 16) located in the left side compartment.
- One 2-valve assembly (valve numbers 17 and 18) located in the right side compartment.
- One 7-valve assembly (valve numbers 20 through 26) located in the right side compartment.
- One 5-valve assembly (valve numbers 27 through 31) located in the right side compartment.

Note: Valve 19 is not used on the A<sup>C</sup>•T 5diff hematology analyzer.

#### **Tools/Supplies Needed**

□ Allen wrenches, 2.5 mm and 3.0 mm, may be required.

#### Preparation

Determine which valve or valves you want to check so you know which door to open or panel to remove.

#### To Only Check Valves in the Left Side Compartment

To check one or more valves with a number 1 through 16:

1. Remove the left side panel and set it aside. For details, see Heading 4.2.

**CAUTION** A flat cable is connected to the rear of the Main card. Be careful when you open the door (where the Main card is attached) that you do not disconnect or damage this cable.

- 2. In the left side compartment,
  - a. Turn the two captive knobs counterclockwise to release the Main card door. These knobs are located to the right of the Main card.
  - b. Open the Main card door and carefully anchor the door behind the white plastic catch to keep it open.
- 3. Go to the Valves Check heading that follows.

#### To Only Check Valves in the Right Side Compartment

To check one or more valves with a number 17 or higher:

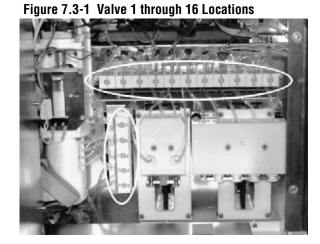
- 1. Open the right side door of the instrument.
- 2. Go to the Valves Check heading that follows.

## **Valves Check**

- 1. From the Main Menu, select **DIAGNOSTICS → HARDWARE SYSTEMS → VALVES**.
- 2. From the Valves menu, select the range of valves you want to check:

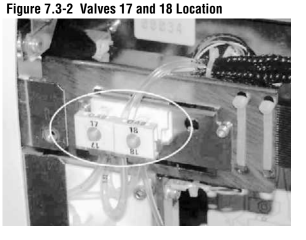
a. 1 to 11 b. 12 to 16

(Valves are located inside the left compartment.)



#### c. 17 and 18

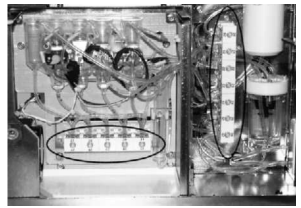
(Valves are located inside the right compartment, just above the sample syringe assembly.)



d. 20 to 26 e. 27 to 31

(Valves are located inside the right compartment.)





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## Wrap Up

When valve performance is acceptable, close all doors and panels:

- 1. If the right side compartment was opened, close the right side door.
- 2. If the left side compartment was opened,
  - a. Close the Main card door. Turn the two captive knobs clockwise to secure the door.

**ATTENTION:** When replacing the left side panel with the instrument powered on, avoid accidently turning the instrument off again by carefully positioning the opening for the power on/off rocker switch over the switch as you position the panel on the instrument frame.

b. Replace the left side panel and install the four hex screws that secure it to the instrument frame.

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# 8.1 MASTER PARTS LISTS

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The parts listed in this section are divided into tables by category. Table 8.1-1 lists the categories and their table numbers.

Within each table the part numbers are listed in numeric order. When applicable, a component is cross referenced to its illustration in Heading 8.2.

Table 8.1-1 Part Categories

| Category                                 | Table | Category                                     | Table  |
|--|-------|--|--------|
| Return Parts                             | 8.1-2 | Waste Alarm Kit, PN - 6912680                | 8.1-9  |
| Nonreturn Parts                          |       | 6 Month Maintenance Kit, PN - XEA485AS       | 8.1-10 |
| Peripherals, Accessories and Consumables |       | 1 Year Maintenance Kit, PN - XEA486AS        | 8.1-11 |
| Tools                                    |       | Every 2 Years Maintenance Kit, PN - XEA581AS | 8.1-12 |
| Fitting Kit Parts PN - XEA311AS          |       | 100 mN-m Torque Driver Kit, - PN 6915456     | 8.1-13 |
| Screws Kit Parts PN - XEA293AS           |       | 400 mN-m Torque Driver Kit, - PN 6915457     | 8.1-14 |
| Installation Kit, PN - XEA484AS          |       | Assorted Tools Kit, - PN 6915458             | 8.1-15 |

#### Table 8.1-2 Return Parts

| Part Number | Description                       | Figure | Item |
|-------------|-----------------------------------|--------|------|
| DBN004A     | Power Supply, module              |        |      |
| XAA423BS    | Preamp, optical signal assembly   | 8.2-40 | 1    |
| XAA477CS    | PCB, Main card (without software) |        |      |
| XAA427BS    | PCB, Motor interface card         |        |      |
| XAA461A     | PCB, Keypad/LCD Display           |        |      |
| XDA600A     | Optical bench, module             |        |      |

**PARTS LISTS** MASTER PARTS LISTS

#### Table 8.1-3 Nonreturn Parts

| Part Number | Description   | Figure                    | lten        |
|-------------|---|---------------------------|-------------|
| 1017413     | Velcro (loop) strip, precut (2 in. x 1 in.), 2 each (used with installation of waste alarm) |                           |             |
| 1017414     | Velcro (hook) strip, precut (2 in. x 1 in.), 2 each (used with installation of waste alarm) |                           |             |
| 3202035     | Tubing, 035 clear polyurethane, 2-inch piece (used with installation of waste alarm)        |                           |             |
| 6011001     | Tie wrap, small   |                           |             |
| 6011002     | Tie wrap, large   |                           |             |
| 6216308     | Feed-through fitting (used with installation of waste alarm)                                |                           |             |
| 6912680     | Waste alarm, kit. Table 8.1-9 lists the contents of this kit.                               |                           |             |
| CAE006A     | Switch, microswitch XC5-81-82   | 8.2-42<br>8.2-44          | 3<br>3      |
| CAE010A     | Switch, microswitch XC5-81  | 8.2-8<br>8.2-16<br>8.2-22 | 7<br>7<br>5 |
| CAY012A     | LCD, display screen   |                           |             |
| DAC011A     | Cable, power, Europe  |                           |             |
| DAC012A     | Cable, power USA  |                           |             |
| DAD113A     | Cable, motor interface to Main Card   |                           |             |
| DAJ007A     | Lamp assembly for Optical module  | 8.2-38                    | 1           |
| DBK004A     | Holder, adhesive, tie wrap, o.d.3 mm, package of 10   |                           |             |
| DZZ015A     | Ferrite shield (for printer cable)  |                           |             |
| DZZ018A     | Ferrite shield (for coax cables)  |                           |             |
| EAC008A     | Fitting, antirotation washer, package of 5  | 8.2-49                    | 11          |
| EAC010A     | Fitting, Luer, female, i.d. 3 mm, package of 5  | 8.2-49                    | 10          |
| EAC019A     | Fitting, Luer, male, i.d. 3 mm, package of 5  |                           |             |
| EAE005AS    | Tubing, Tygon, 1.016 mm i.d. (0.040 in.), 2 meter length                                    |                           |             |
| EAE006AS    | Tubing, Tygon, 1.295 mm i.d. (0.051 in.), 2 meter length                                    |                           |             |
| EAE007AS    | Tubing, Tygon, 1.52 mm i.d. (0.060 in.), 2 meter length                                     |                           |             |
| EAE008AS    | Tubing, Tygon, 2.05 mm i.d. (0.081 in.), 2 meter length                                     | 8.2-7                     | 3           |
| EAE011AS    | Tubing, crystal, 3x6 mm (i.d. x o.d.), 10 meter length                                      |                           |             |
| EAE028AS    | Tubing, crystal, 4x6 mm (i.d. x o.d.), 2 meter length                                       |                           |             |
| EAE034AS    | Tubing, Tygon, 2.54 mm i.d. (0.100 in.), 2 meter length                                     |                           |             |
| FAA040A     | O-ring, 5diff syringe, 12.1 mm diameter, package of 10                                      | 8.2-15                    | 2           |
| FAA046A     | O-ring, coaxial cable, package of 10  |                           |             |
| FAA053A     | O-ring, probe rinse block, package of 10  | 8.2-27                    | 3           |
| FAA064A     | O-ring, sample syringe, package of 10   | 8.2-21                    | 2           |
| FAA065A     | O-ring, reagent syringe, 6.3 mm diameter, package of 10                                     | 8.2-13                    | 8           |

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| Part Number | Description   | Figure | ltem |
|-------------|---|--------|------|
| FAA066A     | O-ring, bath drain/debubble chamber, package of 12      | 8.2-46 | 10   |
|             |   | 8.2-48 | 10   |
| FAA067A     | O-ring, 5diff syringe, 2.4 mm diameter, package of 10   | 8.2-15 | 3    |
| FAK001A     | Aperture, RBC/Plt, 50 µ                                 | 8.2-46 | 4    |
| FAK003A     | Aperture, WBC/BASO, 80 µ                                | 8.2-48 | 4    |
| FAL009A     | Nut, shock mount, for most subassemblies, package of 12 | 8.2-8  | 1    |
|             |   | 8.2-16 | 1    |
|             |   | 8.2-22 | 10   |
| FAL010A     | Nut, shock mount, for optics module, package of 12      |        |      |
| FAM004A     | Foot, plastic, main chassis, package of 8               |        |      |
| FBL001A     | Stopper, rubber with two holes, used for diluent pickup |        |      |
| FBR011A     | Belt, probe vertical, 364 tooth                         | 8.2-30 | 1    |
| FBR012A     | Belt, traverse horizontal, 544 tooth                    | 8.2-35 | 1    |
| GAK302A     | Cap, reagent bottle, 40 mm diameter, package of 12      |        |      |
| GAL094A     | Chamber, diluent reservoir                              | 8.2-7  | 4    |
| GAL098A     | Sleeve, tubing/fitting compression collar               |        |      |
| GBC015A     | Clip, long isolator chamber holder, package of 10       | 8.2-45 | 6    |
| GBC030A     | Piston, reagent syringe, large                          | 8.2-13 | 3    |
| GBC031A     | Piston, reagent syringe, lyse                           | 8.2-13 | 7    |
| GBG003A     | Chamber, drain and debubble                             | 8.2-46 | 12   |
|             |   | 8.2-48 | 12   |
| GBG007A     | Diffuser, drain and debubble, set of 4                  | 8.2-46 | 11   |
|             |   | 8.2-48 | 11   |
| GBG013A     | Drip tray, overflow, bath enclosure                     |        |      |
| GBG022A     | Cover, bath enclosure                                   |        |      |
| GBG205A     | Cover, solenoid drivers, bank 27-31                     | 8.2-42 | 7    |
| GBG033A     | Syringe, reagent, body                                  | 8.2-13 | 1    |
| GBG037A     | Syringe, 5diff, body                                    | 8.2-15 | 1    |
| GBG040A     | Piston, 5diff syringe                                   | 8.2-15 | 5    |
| GBG042A     | Syringe, 5diff, O-ring collar                           | 8.2-15 | 4    |
| GBG044A     | Syringe, sample, body                                   | 8.2-21 | 1    |
| GBG048A     | Syringe, sample, O-ring collar                          | 8.2-21 | 3    |
| GBG052A     | Piston, count/waste syringe                             | 8.2-11 | 4    |
| GBG053A     | Syringe, count, body                                    | 8.2-9  | 3    |
| GBG054A     | Syringe, waste, body                                    | 8.2-17 | 4    |
| GBG090A     | Sample probe, rinse block                               | 8.2-27 | 2    |

# Table 8.1-3 Nonreturn Parts (Continued)

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| Part Number | Description                                       | Figure | Item |
|-------------|---|--------|------|
| GBG091A     | Sample probe, guide                               | 8.2-27 | 4    |
| GBG093A     | Pulley, belt freewheel                            | 8.2-31 | 4    |
|             |   | 8.2-36 | 3    |
| GBG138A     | Key, right side door                              |        |      |
| GBG144A     | Pickup, reagent bottle, 27 mm stopper             |        |      |
| GBG145A     | Pickup, reagent bottle, 20 mm stopper             |        |      |
| GBG147A     | Drip tray, reagents syringe                       |        |      |
| GBG155A     | Cap, reagent bottle, 25 mm diameter, package of 4 |        |      |
| GBG156A     | O-ring, aperture, package of 12                   | 8.2-46 | 5    |
|             |   | 8.2-48 | 5    |
| GBG157A     | Counting head (internal electrode)                | 8.2-46 | 7    |
|             |   | 8.2-48 | 7    |
| GBG210A     | Syringe body (new style)                          |        |      |
| GBG211A     | Chamber, counting syringe (new style)             |        |      |
| GBG212A     | Chamber, drain syringe (new style)                |        |      |
| GBG219A     | Anti-extrusion ring                               | 8.2-9  | 5    |
|             |   | 8.2-17 | 5    |
| GBG225A     | Keypad, silicone                                  |        |      |
| HAE026B     | Label, valves, 1-10, package of 10                |        |      |
| HAE027B     | Label, valves, 11-20, package of 10               |        |      |
| HAE028B     | Label, valves, 21-30, package of 10               |        |      |
| HAE029B     | Label, valves, 31-40, package of 10               |        |      |
| HAX0012     | Label, tubing, reagent lines                      |        |      |
| HAX0013     | Label, tubing, bath enclosure lines               |        |      |
| XAA468A     | PCB, LED card, aspiration indicator               | 8.2-41 | 2    |
| XBA144A     | Cable, reagent ground fitting                     |        |      |
| XBA193A     | Cable, coaxial, flow cell optical                 |        |      |
| XBA386A     | Cable, bath enclosure harness                     |        |      |
| XBA389A     | Photometer, HGB diode and preamp assembly         | 8.2-47 | 4    |
| XBA390A     | Motor, for syringe assembly                       | 8.2-10 | 1    |
|             |   | 8.2-18 | 1    |
|             |   | 8.2-22 | 1    |
| XBA391A     | Motor, traverse, horizontal                       | 8.2-34 | 1    |
| XBA392A     | Motor, sample probe, vertical                     | 8.2-31 | 9    |
|             |   | 8.2-32 | 1    |
| XBA393A     | Fan, main chassis, 24 V                           | 8.2-49 | 6    |

| Part Number | Description  | Figure | ltem |
|-------------|--|--------|------|
| XBA394A     | Sensor, traverse horizontal, home                                    | 8.2-36 | 7    |
| XBA395A     | Sensor, fluid, bath drain  |        |      |
| XBA396A     | Sensor, sample probe vertical, home                                  | 8.2-33 | 1    |
| XBA398B     | Cable, coaxial, with RBC/WBC bath electrode                          | 8.2-46 | 2    |
|             |  | 8.2-48 | 2    |
| XBA399A     | Cable, coaxial, flow cell volume                                     |        |      |
| XBA403A     | Shield, flow cell tubing ground                                      | 8.2-39 | 2    |
| XCA166A     | Chamber, isolator (long)   | 8.2-45 | 5    |
| XCA167A     | Chamber, isolator (small)  |        |      |
| XDA481B     | Valve, liquid. 2-way, normally closed, w/o coil                      |        |      |
| XDA483B     | Valve, liquid, 3-way, w/o coil                                       |        |      |
| XDA591AS    | Syringe, 5diff assembly  | 8.2-14 | 2    |
|             |  | 8.2-15 | 10   |
| XDA592AS    | Syringe, reagent assembly  | 8.2-12 | 3    |
|             |  | 8.2-13 | 9    |
| XDA593AS    | Syringe, sampling assembly   | 8.2-20 | 2    |
| XDA594AS    | Motor, assembly, for 5diff syringe                                   | 8.2-14 | 1    |
| XDA595AS    | Motor, assembly, for reagent syringe                                 | 8.2-12 | 1    |
| XDA596AS    | Motor, assembly, for sample syringe                                  | 8.2-20 | 1    |
| XDA597BS    | Syringe, waste - complete assembly                                   | 8.2-16 | 2    |
|             |  | 8.2-17 | 1    |
| XDA598BS    | Syringe, vacuum - complete assembly                                  | 8.2-8  | 2    |
|             |  | 8.2-9  | 1    |
| XDA601AS    | Flow cell, assembly  | 8.2-39 | 1    |
| XDA602B     | Bath, assembly, 3-baths and rinse chamber (includes long coax cable) | 8.2-47 | 3    |
| XDA605A     | Reservoir, diluent, assembly   | 8.2-6  | 1    |
|             |  | 8.2-7  | 8    |
| XDA610B     | Bath, WBC/BASO (includes long coax cable)                            | 8.2-47 | 1    |
| XDA611CS    | Valve, liquid, 11-valve assembly (1-11)                              | 8.2-1  | 1    |
| XDA612CS    | Valve, liquid, 5-valve assembly (12-16)                              | 8.2-2  | 1    |
| XDA613CS    | Valve, liquid, 2-valve assembly (17-18)                              | 8.2-3  | 1    |
|             |  | 8.2-19 | 2    |
| XDA614CS    | Valve, liquid, 7-valve assembly (20-26)                              | 8.2-4  | 1    |
| XDA615CS    | Valve, liquid, 5-valve assembly (27-31)                              | 8.2-5  | 1    |
|             |  | 8.2-45 | 4    |
| XDA616AS    | Piston, 5diff syringe, 190 µL needle                                 | 8.2-15 | 9    |

# Table 8.1-3 Nonreturn Parts (Continued)

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| Part Number | Description   | Figure | ltem |
|-------------|---|--------|------|
| XDA617AS    | Piston, sample syringe, 100 µL needle   | 8.2-21 | 6    |
| XDA618AS    | Guide, sample probe retainer  | 8.2-26 | 1    |
| XDA619AS    | Probe, sample   | 8.2-28 | 1    |
| XDA621A     | O-ring, with matched washer, count/waste syringe, package of 10   | 8.2-9  | 2    |
|             |   | 8.2-13 | 2    |
|             |   | 8.2-17 | 2    |
| XDA622A     | O-ring, with matched washer, for reagent syringe, package of 10   |        |      |
| XDA623AS    | Panel, reagent/waste fittings assembly  | 8.2-49 | 14   |
| XDA625AS    | Heater, reagent coil assembly   | 8.2-45 | 1    |
| XDA626AS    | Cover, diluent reservoir  | 8.2-7  | 2    |
| XDA657B     | Bath assembly, including WBC/Baso bath (includes long coax cable)   |        |      |
| XEA018A     | Pickup tube, diluent (needs stopper)  |        |      |
| XEA286AS    | Kit, O-ring and washer  | 8.2-7  | 7    |
| XEA293AS    | Kit, screws. Table 8.1-7 lists the contents of this kit.  |        |      |
|             | <b>Note:</b> The Screws kit contains the most common screws used in the A <sup>C</sup> •T 5diff analyzer, plus nuts and washers.  |        |      |
| XEA311AS    | Kit, fittings. Table 8.1-6 lists the contents of this kit.  |        |      |
|             | <b>Note:</b> The Fitting kit contains the most common pneumatic and hydraulic parts used in the A <sup>C</sup> •T 5diff analyzer. |        |      |
| XEA484A     | Kit, installation (comes with instrument). Table 8.1-8 lists the contents of this kit.  |        |      |
| XEA485AS    | Kit, maintenance, 6 month. Table 8.1-10 lists the contents of this kit.   |        |      |
| XEA486AS    | Kit, maintenance, 1 year. Table 8.1-11 lists the contents of this kit.  |        |      |
| XEA487AS    | Fan, bath enclosure temperature control   | 8.2-43 | 2    |
| XEA488AS    | Cable, heating coil harness   |        |      |
| XEA581AS    | Kit, maintenance, every two years. Table 8.1-12 lists the contents of this kit.   |        |      |
| XEA616AS    | Tubing, with sleeve, for flow cell sample port 6  |        |      |
| XEA643AS    | Kit, software PROMS, A <sup>C</sup> •T 5diff v1.03  |        |      |

## Table 8.1-3 Nonreturn Parts (Continued)

# $\label{eq:constraint} \textbf{Table 8.1-4} \hspace{0.1 in} \textbf{Peripherals, Accessories and Consumables}$

| Part Number | Description  |
|-------------|--|
| 2016891     | Ribbon, black print, for EPSON LX300                   |
| 4237615     | Operator's Manual, A <sup>C</sup> •T 5diff (English)   |
| 4237616     | Service Manual, A <sup>C</sup> •T 5diff (English only) |
| 4237630     | Operator's Manual, A <sup>C</sup> •T 5diff (French)    |
| 4237631     | Operator's Manual, A <sup>C</sup> •T 5diff (Italian)   |
| 4237632     | Operator's Manual, A <sup>C</sup> •T 5diff (German)    |

| Part Number | Description  |
|-------------|--|
| 4237633     | Operator's Manual, AC+T 5diff (Spanish)              |
| 4237634     | Operator's Manual, A <sup>C</sup> •T 5diff (Chinese) |
| 7547175     | Calibrator, A <sup>C</sup> •T 5diff Cal              |
| 7547176     | Control, A <sup>C</sup> •T 5diff                     |
| 8547168     | Reagent, A <sup>C</sup> •T 5diff Hgb Lyse            |
| 8547169     | Reagent, A <sup>C</sup> •T 5diff Diluent             |
| 8547170     | Reagent, A <sup>C</sup> •T 5diff WBC Lyse            |
| 8547171     | Reagent, A <sup>C</sup> •T 5diff Fix                 |
| 8547176     | Reagent, A <sup>C</sup> •T 5diff Rinse               |
| CBE070AS    | Printer Head, Epson LX300+                           |
| FBH016A     | Cover, dust  |
| LAD002AS    | Latex particles, RBC/PLT (used for all procedures)   |
| LAM004A     | Grease, silicone                                     |
| XAA473A     | Printer, Epson LX300+ 220 V                          |
| XAA476A     | Printer, Epson LX300+ 110 V                          |
| XEA019A     | Grease, silicone, for syringe assembly               |
| XEA381AS    | Grease, for mechanical assemblies                    |
| XEA587AS    | Bar code reader option                               |

Table 8.1-4 Peripherals, Accessories and Consumables (Continued)

## Table 8.1-5 Tools

|   | Part Number | Description  |
|---|-------------|--|
|   | 5415407     | Set, Allen wrenches, balldriver, metric                                |
|   | 5450517     | Screwdriver, 2.5 mm hex, balldriver                                    |
| l | 5450518     | Screwdriver, 3.0 mm hex, balldriver                                    |
|   | 5450519     | Bit, 2.5 mm hex balldriver, for 1/4 inch drive                         |
|   | 5450520     | Bit, 3.0 mm hex balldriver, for 1/4 inch drive                         |
| l | 5450521     | Screwdriver, torque, preset to 100 mN-m (14.2 oz.f-in), 1/4 inch drive |
|   | 5450522     | Screwdriver, torque, preset to 400 mN-m (56.8 oz.f-in), 1/4 inch drive |
| l | 5450532     | Screwdriver, T10 Torx  |
|   | 5450533     | Bit, T10 Torx, for 1/4 inch drive                                      |
|   | 5450535     | Screwdriver, small, Phillips 00  |
|   | 5450536     | Magnifier, jeweller's loupe, 5X with 2 in. focal length                |
| l | 5450537     | Extractor, chip remover, for U42 and U43                               |
|   | 5450540     | Gauge, feeler adjustment, 9.4 mm and 3.0 mm                            |

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## Table 8.1-5 Tools (Continued)

| Part Number | Description   |
|-------------|---|
| 6915456     | Kit, 100 mN-m torque driver with 3 bits. Table 8.1-13 lists the contents of this kit.         |
| 6915457     | Kit, 400 mN-m torque driver with 3 bits. Table 8.1-14 lists the contents of this kit.         |
| 6915458     | Kit, assorted tools for A <sup>C</sup> •T 5diff. Table 8.1-15 lists the contents of this kit. |
| MAB002A     | Allen wrench, L-shape, 2.5 mm   |
| MAB018A     | Allen wrench, L-shape, 3.0 mm   |
| MAB090A     | Torx key, L-shape, T10  |
| XDA555AS    | Knob, used to adjust flowcell   |
| XEA585AS    | Torque screwdriver, A300, adjustable torque, 100 mN-m, 1/8 inch drive                         |
| XEA587AS    | Torque screwdriver, A301, adjustable torque, 400 mN-m, 1/4 inch drive                         |

#### Table 8.1-6 Fitting Kit Parts PN - XEA311AS

| Part Number | Description                                    | Quantity |
|-------------|--|----------|
| EAA005A     | Fitting, straight connector, 01.6 mm 1/8       | 10       |
| EAA006A     | Fitting, straight connector                    | 10       |
| EAA009A     | Fitting, bent connector, i.d.1.5 10/32         | 10       |
| EAA013A     | Fitting, bent connector, 1/8                   | 10       |
| EAA014A     | Fitting, straight connector, 1/8               | 10       |
| EAB002A     | Fitting, L-shaped connector                    | 10       |
| EAB003A     | Fitting, connector04 mm H19.5 20               |          |
| EAB005A     | Fitting, T-connector, 01.6 mm T10-6 30         |          |
| EAB006A     | Fitting, T-connector, 02.3 mm 30               |          |
| EAB009A     | Fitting, straight connector, 01.6 mm 30        |          |
| EAB010A     | Fitting, straight connector, 02.3 mm           | 30       |
| EAB014A     | Fitting, straight connector, Y, 01.5 mm/1.5 mm | 10       |
| EAB021A     | Fitting, Y-connector, 03 mm 3                  |          |
| EAB026A     | Fitting, Y-connector, 02.5 mm 3                |          |
| EAB035A     | Fitting, T-connector, 02.3 mm T220-610         |          |

#### Table 8.1-7 Screws Kit Parts PN - XEA293AS

| Part Number | Description      | Quantity |
|-------------|------------------|----------|
| KAA002A     | Screw, hex M3x6  | 20       |
| KAA003A     | Screw, hex M3x8  | 20       |
| KAA005A     | Screw, hex M3x12 | 20       |

| Part Number | Description                    | Quantity |  |  |
|-------------|--------------------------------|----------|--|--|
| KAA006A     | Screw, hex M3x14               | 20       |  |  |
| KAA009A     | Screw, hex M3x20               |          |  |  |
| KAA011A     | Screw, hex M3x30               | 20       |  |  |
| KAA015A     | Screw, hex M4x8                | 20       |  |  |
| KAA016A     | Screw, hex M4x12               | 20       |  |  |
| KAA017A     | Screw, hex M4x16               | 20       |  |  |
| KAA021A     | Screw, hex M5x10               | 20       |  |  |
| KAA028A     | Screw, hex M4x20               | 20       |  |  |
| KAB002A     | Screw, FHC M3x6                | 20       |  |  |
| KAB003A     | Screw, FHC M3x8                | 5        |  |  |
| KAB004A     | Screw, FHC M3x10               | 20       |  |  |
| KAB005A     | Screw, FHC M3x12               | 20       |  |  |
| KAB016A     | Screw, torx M3x8               | 5        |  |  |
| KAC002A     | Screw, TC M3x5                 | 20       |  |  |
| KAE003A     | Screw, BHC M4x12 screw         | 20       |  |  |
| KAH001A     | Nut, HU M3                     | 20       |  |  |
| KAH002A     | Nut, HU M4                     | 20       |  |  |
| KAH003A     | Nut, HU M5                     | 20       |  |  |
| KAH024A     | Nut, 1/4-28 (US std)           | 10       |  |  |
| KAJ001A     | Washer, M diameter 0.3         | 20       |  |  |
| KAJ002A     | Washer, M diameter 0.4         | 20       |  |  |
| KAJ003A     | Washer, M diameter 0.5         | 20       |  |  |
| KAJ007A     | Washer, AZ diameter 0.3        | 20       |  |  |
| KAJ008A     | Washer, Z diameter 0.4         | 20       |  |  |
| KAJ009A     | Washer, AZ diameter 0.5        | 20       |  |  |
| KAJ010A     | Washer, W grower diameter 0.5  | 20       |  |  |
| KAM002A     | Collar, plastic, M3x6 Lg.8     | 10       |  |  |
| KAM003A     | Collar, plastic, M4x6 Lg.8     | 10       |  |  |
| KAM004A     | Collar, plastic, M3.5x6 Lg.6   |          |  |  |
| KAM005A     | Collar, plastic, M3.5x6 Lg.8   | 10       |  |  |
| KAM006A     | Collar, plastic, M3.5x6 Lg.12  |          |  |  |
| KAM010A     | Collar, plastic, M3x6 Lg.10    | 10       |  |  |
| KAM011A     | Collar, plastic, M3.5x6 Lg.5.7 | 10       |  |  |
| KAM013A     | Collar, plastic, M3.5x6 Lg.10  | 10       |  |  |
| KAM017A     | Collar, brass, M3.5x6 Lg.3     | 10       |  |  |

Table 8.1-7 Screws Kit Parts PN - XEA293AS (Continued)

| Part Number | Description                               | Quantity |  |  |
|-------------|---|----------|--|--|
| DAJ007A     | Lamp, 20 W, 9.5 Vdc                       | 1        |  |  |
| DBH001A     | Tie wrap LA=2.4 L=92                      |          |  |  |
| DBH002A     | Tie wrap LA=3.6 L=140                     | 3        |  |  |
| DBK003A     | Adhesive holder 3 mm                      | 1        |  |  |
| EAE007A     | Tygon tube 1.52 mm (0.060 inch)           | 2        |  |  |
| EAE006A     | Tygon tube 1.295 mm (0.051 inch)          | 2        |  |  |
| EAE034A     | Tygon tube 2.54 mm (0.100 inch)           | 2        |  |  |
| EAE005A     | Tygon tube 1.02 mm (0.040 inch)           | 2        |  |  |
| EAE008A     | Tygon tube 2.05 mm (0.081 inch)           | 2        |  |  |
| EAE011A     | Cristal tube 3 mm (i.d.) x 6 mm (o.d.)    | 4        |  |  |
| EAE028A     | Cristal tube 4 mm (i.d.) x 6 mm (o.d.)    | 4        |  |  |
| EAC019A     | Luer male connector, i.d. 3 mm            | 3        |  |  |
| EAC010A     | Luer bushing, i.d. 3 mm                   | 2        |  |  |
| EAB021A     | Fitting, Y-connector 3 mm                 | 1        |  |  |
| EAB033A     | Fitting, T-connector (small) 1            |          |  |  |
| EAB035A     | Fitting, T-connector 2.3 mm T220-6 inches |          |  |  |
| FAA053A     | O-ring, 1.40 x 1.25, fluocarbon           | 1        |  |  |
| FBH016A     | Instrument cover                          |          |  |  |
| FBL001A     | Rubber cap 2 holes                        |          |  |  |
| GBG138A     | Tool, right access panel fasteners        | 1        |  |  |
| GBG156A     | EPO counting head Joint                   | 2        |  |  |
| GBG144A     | Reagent straw cap 027                     | 3        |  |  |
| GBG145A     | Reagent straw cap 020                     | 1        |  |  |
| GAK302A     | Factory bottle cap                        | 3        |  |  |
| GBG155A     | Cap 025                                   | 1        |  |  |
| MAB018A     | Bent allen key 3 mm                       | 1        |  |  |
| MAB002A     | Bent allen key 2.5 mm                     | 1        |  |  |
| MAB090A     | Bent Torx key T10                         | 1        |  |  |
| XDA619AS    | Equipped probe                            | 1        |  |  |
| XDA483B     | Valve 3V (no solenoid)                    | 1        |  |  |
| XDA621A     | 0-ring 30.80 x 3.80 + wedge               | 1        |  |  |
| XEA018A     | Diluent pickup tube, 360 mm length        | 1        |  |  |
| XEA019A     | Grease KM 1011                            | 1        |  |  |

#### Table 8.1-8 Installation Kit, PN - XEA484AS

#### Table 8.1-9 Waste Alarm Kit, PN - 6912680

| Part Number | Description                   | Quantity |
|-------------|-------------------------------|----------|
| 6706397     | Alarm, waste, with 9V battery | 1        |
| 6856742     | Sensor, float                 | 1        |

#### Table 8.1-10 6 Month Maintenance Kit, PN - XEA485AS

| Part Number | Description  | Quantity |
|-------------|--|----------|
| FAA053A     | O-ring, fluorocarbon, rinse block assembly   | 1        |
| FAA065A     | ing, silicone, Hgb Lyse reagent syringe 1  |          |
| XDA621A     | ring and washer, matched set, waste or count syringe 2                             |          |
| XDA622A     | O-ring and washer, matched set, for all reagent syringes (except Hgb Lyse syringe) | 4        |
| XEA019A     | Silicone grease  | 1        |

#### Table 8.1-11 1 Year Maintenance Kit, PN - XEA486AS

| Part Number | Description   |   |  |
|-------------|---|---|--|
| DAJ007A     | Lamp, optical bench, 20 W, 9.5 Vdc                  | 1 |  |
| FAA040A     | O-ring, silicone, 5diff syringe                     | 1 |  |
| FAA046A     | O-ring, coaxial cable, counting head                | 2 |  |
| FAA064A     | O-ring, fluorocarbon, sample syringe 2              |   |  |
| FAA066A     | 0-ring, silicon, draining bath 4                    |   |  |
| FAA067A     | O-ring, fluorocarbon, 5diff syringe 4               |   |  |
| GBG156A     | O-ring, aperture, counting head 4                   |   |  |
| XBA399A     | Cable, coaxial, DIFF flow cell 1                    |   |  |
| XEA286AS    | O-ring and washer, matched set, diluent reservoir 1 |   |  |

#### Table 8.1-12 Every 2 Years Maintenance Kit, PN - XEA581AS

| Part Number | Description                        | Quantity |
|-------------|------------------------------------|----------|
| GBC031A     | Piston, Hgb Lyse reagent syringe   | 1        |
| GBC030A     | stons, reagent syringes assembly 4 |          |
| GBG052A     | Piston, waste or count syringe 2   |          |
| GBG091A     | Block, sample probe guide          | 1        |
| XDA619AS    | Probe, sample 1                    |          |

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#### Table 8.1-13 100 mN-m Torque Driver Kit, - PN 6915456

|   | Part Number | Description  | Quantity |
|---|-------------|--|----------|
| I | 5450519     | Bit, 2.5 mm hex balldriver, for 1/4 inch drive                         | 1        |
| I | 5450520     | Bit, 3.0 mm hex balldriver, for 1/4 inch drive                         | 1        |
| I | 5450521     | Screwdriver, torque, preset to 100 mN-m (14.2 oz.f-in), 1/4 inch drive | 1        |
|   | 5450533     | Bit, T10 Torx, for 1/4 inch drive                                      | 1        |

#### Table 8.1-14 400 mN-m Torque Driver Kit, - PN 6915457

|    | Part Number | Description  | Quantity |
|----|-------------|--|----------|
|    | 5450519     | Bit, 2.5 mm hex balldriver, for 1/4 inch drive                         | 1        |
|    | 5450520     | Bit, 3.0 mm hex balldriver, for 1/4 inch drive                         | 1        |
|    | 5450522     | Screwdriver, torque, preset to 400 mN-m (56.8 oz.f-in), 1/4 inch drive | 1        |
| I. | 5450533     | Bit, T10 Torx, for 1/4 inch drive                                      | 1        |

#### Table 8.1-15 Assorted Tools Kit, - PN 6915458

|  | Part Number                                     | Description   | Quantity |
|--|---|---|----------|
|  | 5415407 Set, Allen wrenches, balldriver, metric |   | 1        |
|  | 5450517   | Screwdriver, 2.5 mm hex, balldriver                     | 1        |
|  | 5450518   | Screwdriver, 3.0 mm hex, balldriver                     | 1        |
|  | 5450532   | Screwdriver, T10 Torx                                   | 1        |
|  | 5450535   | Screwdriver, small, Phillips 00                         | 1        |
|  | 5450536   | Magnifier, jeweller's loupe, 5X with 2 in. focal length | 1        |
|  | 5450537   | Extractor, chip remover, for U42 and U43                | 1        |

# 8.2 ILLUSTRATED PARTS

The exploded views in this section are shown for informational purposes only. Many of the subcomponents are not available. Consult the MASTER PARTS LISTS for part availability.

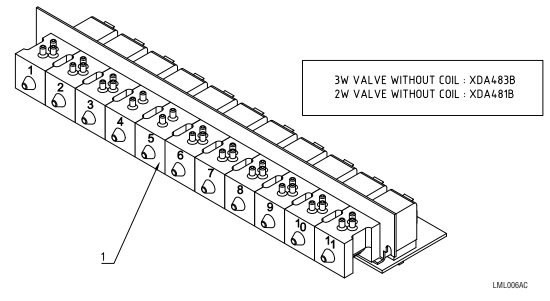
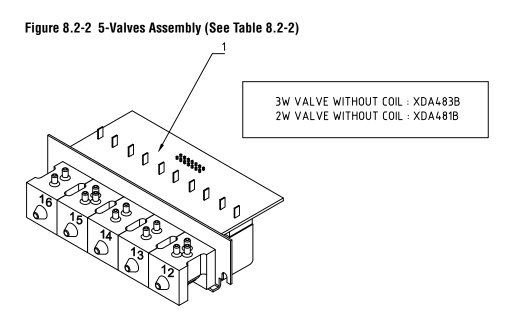


Figure 8.2-1 11-Valves Assembly (See Table 8.2-1)

Table 8.2-1 11-Valves Assembly (See Figure 8.2-1)

| ltem | Part Number | Description                             |
|------|-------------|---|
| 1    | XDA611CS    | Valve, liquid, 11-valve assembly (1-11) |



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| ltem | Part Number | Description                             |
|------|-------------|---|
| 1    | XDA612CS    | Valve, liquid, 5-valve assembly (12-16) |



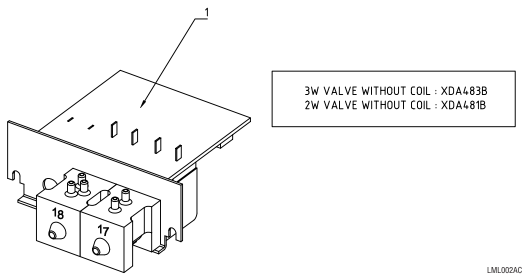
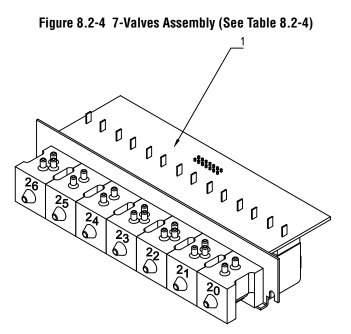


Table 8.2-3 2-Valves Assembly (See Figure 8.2-3)

| ltem | Part Number | Description                             |
|------|-------------|---|
| 1    | XDA613CS    | Valve, liquid, 2-valve assembly (17-18) |



3W VALVE WITHOUT COIL : XDA483B 2W VALVE WITHOUT COIL : XDA481B

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#### Table 8.2-4 7-Valves Assembly (See Figure 8.2-4)

| ltem | Part Number | Description                             |
|------|-------------|---|
| 1    | XDA614CS    | Valve, liquid, 7-valve assembly (20-26) |



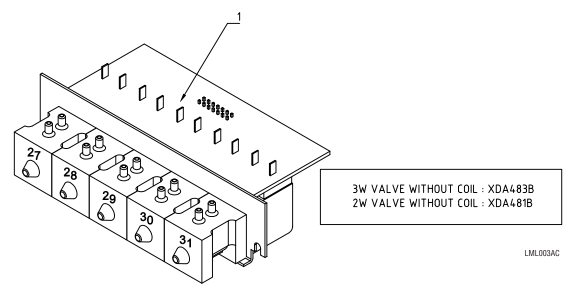


Table 8.2-5 5-Valves Assembly (See Figure 8.2-5)

| ltem | Part Number | Description                             |
|------|-------------|---|
| 1    | XDA615CS    | Valve, liquid, 5-valve assembly (27-31) |

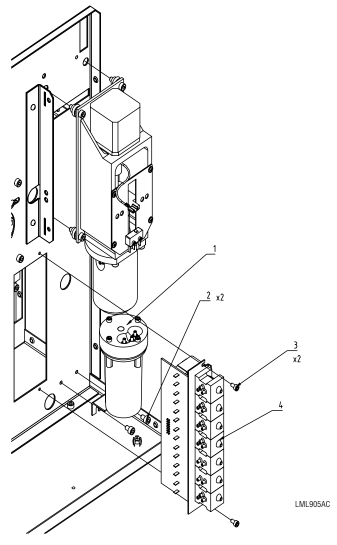
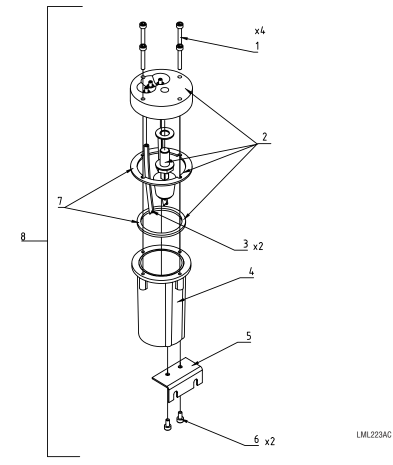


Figure 8.2-6 Right Side Compartment, Lower Rear Area (See Table 8.2-6)

Table 8.2-6 Right Side Compartment, Lower Rear Area (See Figure 8.2-6)

| ltem | Part Number | Description                             |
|------|-------------|---|
| 1    | XDA605A     | Reservoir, diluent, assembly            |
| 2    | KAA013A     | Screw, hex M4x6                         |
| 3    | KAA002A     | Screw, hex M3x6                         |
| 4    | XDA603CS    | Valve, liquid, 7-valve assembly (20-26) |



## Figure 8.2-7 Diluent Reservoir Assembly (See Table 8.2-7)

Table 8.2-7 Diluent Reservoir Assembly (See Figure 8.2-7)

| ltem | Part Number | Description   |
|------|-------------|---|
| 1    | KAA009A     | Screw, hex M3x20  |
| 2    | XDA626AS    | Cover, diluent reservoir                                |
| 3    | EAE008A     | Tubing, Tygon, 2.05 mm i.d. (0.081 in.), 2 meter length |
| 4    | GAL094A     | Chamber, diluent reservoir                              |
| 5    | GBG056A     | Brace, reservoir, diluent                               |
| 6    | KAA002A     | Screw, hex M3x6   |
| 7    | XEA286AS    | Kit, O-ring and washer                                  |
| 8    | XDA605A     | Reservoir, diluent, assembly                            |

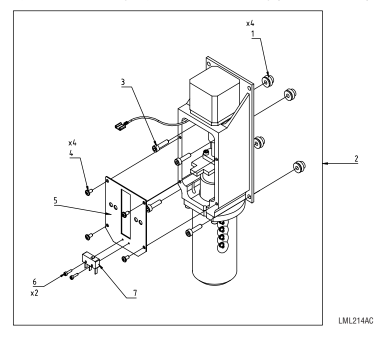
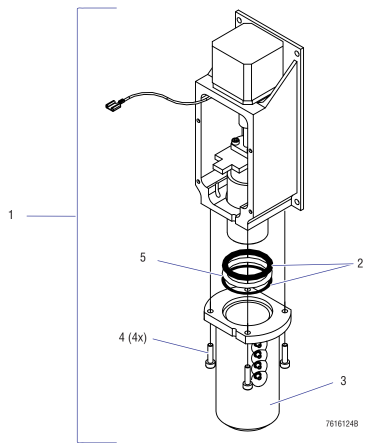


Figure 8.2-8 Count Syringe and Motor Assembly (See Table 8.2-8)

| ltem | Part Number          | Description   |
|------|----------------------|---|
| 1    | FAL009A              | Nut, shock mount, package of 12 - for most subassemblies  |
| 2    | XDA598AS<br>XDA598BS | Syringe, vacuum - complete assembly<br>Syringe, vacuum - complete assembly (includes anti-extrusion ring) |
| 3    | KAA017A              | Screw, hex M4x16  |
| 4    | KZZ022A              | Screw, auto-threaded  |
| 5    | GBG027A              | Plate, microswitch, motor   |
| 6    | KAA040A              | Screw, hex M2x8   |
| 7    | CAE010A              | Switch, microswitch XC5-81  |



# Figure 8.2-9 Count Syringe Assembly (See Table 8.2-9)

 Table 8.2-9
 Count Syringe Assembly (See Figure 8.2-9)

| ltem | Part Number | Description  |
|------|-------------|--|
| 1    | XDA598AS    | Syringe, vacuum - complete assembly                                |
|      | XDA598BS    | Syringe, vacuum - complete assembly (includes anti-extrusion ring) |
| 2    | XDA621A     | O-ring, with matched washer, count/waste syringe, package of 10    |
| 3    | GBG053A     | Syringe, count, body   |
| 4    | KA017A      | Screw, hex M4x16   |
| 5    | GBG219A     | Anti-extrusion ring  |

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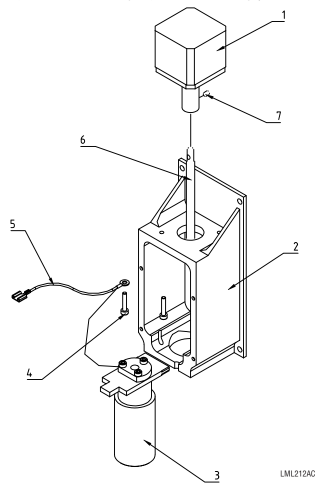


Figure 8.2-10 Count Syringe Motor Assembly (See Table 8.2-10)

Table 8.2-10 Count Syringe Motor Assembly (See Figure 8.2-10)

| ltem | Part Number | Description  |
|------|-------------|--|
| 1    | XBA390A     | Motor, for syringe assembly                        |
| 2    | GBG050A     | Body, count syringe                                |
| 3    |             | Piston assembly, count syringe (See Figure 8.2-11) |
| 4    | KAA007A     | Screw, hex M3x16                                   |
| 5    |             | Grounding wire                                     |
| 6    | GBG029A     | Syringe screw                                      |
| 7    | KAD016A     | Screw, HC M4x6                                     |

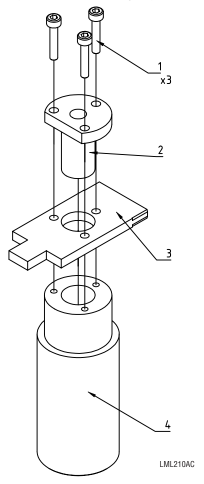


Figure 8.2-11 Count Syringe Piston Assembly (See Table 8.2-11)

Table 8.2-11 Count Syringe Piston Assembly (See Figure 8.2-11)

| ltem | Part Number | Description                      |
|------|-------------|----------------------------------|
| 1    | KAA007A     | Screw, hex M3x16                 |
| 2    | GBG029A     | Syringe nut                      |
| 3    | GBG051A     | Guide plate, count/waste syringe |
| 4    | GBG052A     | Piston, count/waste syringe      |

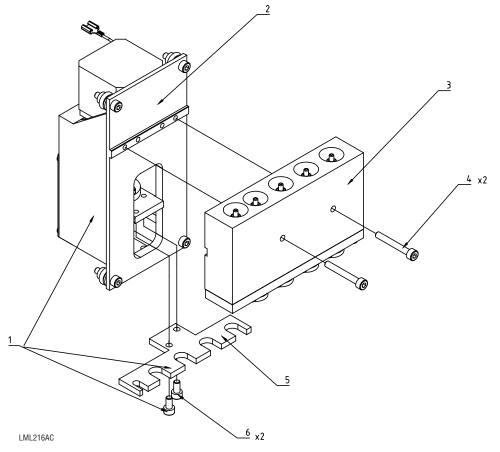


Figure 8.2-12 Reagent Syringes and Motor Assembly (See Table 8.2-12)

| Table 8.2-12 | <b>Reagent Syringes</b> | and Motor Ass | sembly (See | Figure 8.2-12)                        |
|--------------|-------------------------|---------------|-------------|---------------------------------------|
|              |                         |               |             | · · · · · · · · · · · · · · · · · · · |

| ltem | Part Number | Description                           |
|------|-------------|---------------------------------------|
| 1    | XDA595AS    | Motor, assembly - for reagent syringe |
| 2    |             | Motor - for reagent syringe           |
| 3    | XDA592AS    | Syringes, reagent assembly            |
| 4    | KAA030A     | Screw, hex M4x30                      |
| 5    | GBG035A     | Guide plate, reagent syringe          |
| 6    | KAA015A     | Screw, hex M4x8                       |

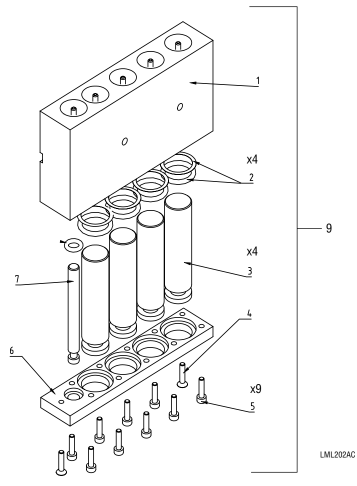


Figure 8.2-13 Reagent Syringes Assembly (See Table 8.2-13)

Table 8.2-13 Reagent Syringes Assembly (See Figure 8.2-13)

| ltem | Part Number | Description   |
|------|-------------|---|
| 1    | GBG033A     | Syringe, reagent, body  |
| 2    | XDA621A     | O-ring, with matched washer, count/waste syringe, package of 10 |
| 3    | GBC030A     | Piston, reagent syringe, large                                  |
| 4    | KAB017A     | Screw, torx M3x12   |
| 5    | KAA005A     | Screw, hex M3x12  |
| 6    | GBG034A     | Syringe, reagent, top   |
| 7    | GBC031A     | Piston, reagent syringe, lyse                                   |
| 8    | FAA065A     | O-ring, reagent syringe, 6.3 mm diameter, package of 10         |
| 9    | XDA592AS    | Syringe, reagent assembly                                       |

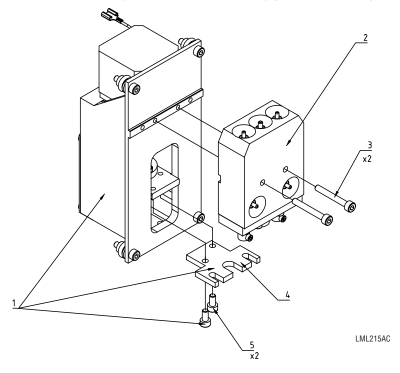




Table 8.2-14 5diff Syringe and Motor Assembly (See Figure 8.2-14)

| ltem | Part Number | Description                         |
|------|-------------|-------------------------------------|
| 1    | XDA594AS    | Motor, assembly - for 5diff syringe |
| 2    | XDA591AS    | Syringe, 5diff assembly             |
| 3    | KAA030A     | Screw, hex M4x30                    |
| 4    | GBG039A     | Guide plate, syringe                |
| 5    | KAA015A     | Screw, hex M4x6                     |

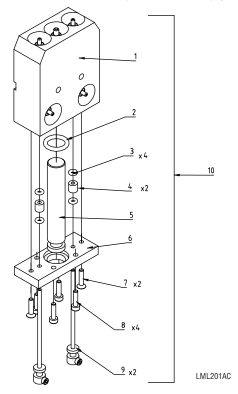


Figure 8.2-15 5diff Syringe Assembly (See Table 8.2-15)

Table 8.2-15 5diff Syringe Assembly (See Figure 8.2-15)

| ltem | Part Number | Description  |
|------|-------------|--|
| 1    | GBG037A     | Syringe, 5diff, body                                   |
| 2    | FAA040A     | O-ring, 5diff syringe, 12.1 mm diameter, package of 10 |
| 3    | FAA067A     | O-ring, 5diff syringe, 2.4 mm diameter, package of 10  |
| 4    | GBG042A     | Syringe, 5diff, O-ring collar                          |
| 5    | GBG040A     | Piston, 5diff syringe                                  |
| 6    | GBG038A     | Syringe top  |
| 7    | KAB017A     | Screw, torx M3x12                                      |
| 8    | KAA005A     | Screw, hex M3x12                                       |
| 9    | XDA616AS    | Piston, 5diff syringe, 190 µL needle                   |
| 10   | XDA591AS    | Syringe, 5diff assembly                                |

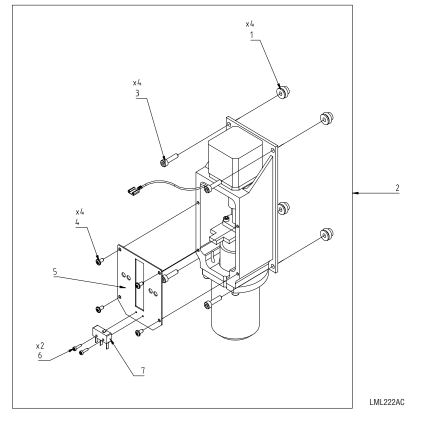
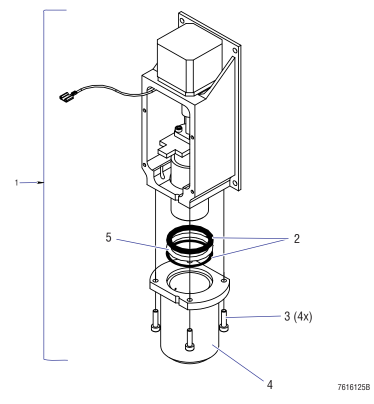


Figure 8.2-16 Waste Syringe and Motor Assembly (See Table 8.2-16)

Table 8.2-16 Waste Syringe and Motor Assembly (See Figure 8.2-16)

| ltem | Part Number | Description   |
|------|-------------|---|
| 1    | FAL009A     | Nut, shock mount, package of 12 - for most subassemblies          |
| 2    | XDA597AS    | Syringe, waste - complete assembly                                |
|      | XDA597BS    | Syringe, waste - complete assembly (includes anti-extrusion ring) |
| 3    | KAA017A     | Screw, hex M4x16  |
| 4    | KZZ022A     | Screw, auto-threaded  |
| 5    | GBG027A     | Plate, microswitch, motor   |
| 6    | KAA040A     | Screw, hex M2x8   |
| 7    | CAE010A     | Switch, microswitch XC5-81  |



# Figure 8.2-17 Waste Syringe Assembly (See Table 8.2-17)

Table 8.2-17 Waste Syringe Assembly (See Figure 8.2-17)

| ltem | Part Number          | Description   |
|------|----------------------|---|
| 1    | XDA597AS<br>XDA597BS | Syringe, waste - complete assembly<br>Syringe, waste - complete assembly (includes anti-extrusion ring) |
| 2    | XDA621A              | O-ring, with matched washer, count/waste syringe, package of 10   |
| 3    | KAA017A              | Screw, hex M4x16  |
| 4    | GBG054A              | Syringe, waste, body  |
| 5    | BGG219A              | Anti-extrusion ring   |

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## Figure 8.2-18 Syringe Motor (See Table 8.2-18)

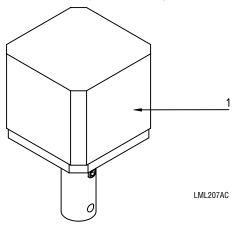


Table 8.2-18 Syringe Motor (See Figure 8.2-18)

| ltem | Part Number | Description                  |
|------|-------------|------------------------------|
| 1    | XBA390A     | Motor - for syringe assembly |



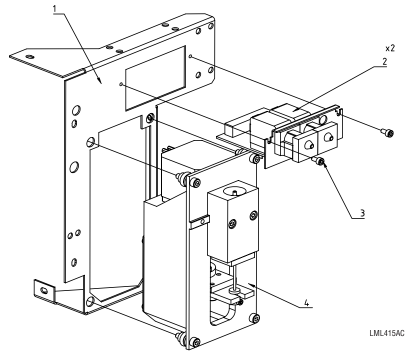


Table 8.2-19 Sample Motor (See Figure 8.2-19)

| ltem | Part Number | Description   |
|------|-------------|---|
| 1    | GBG086A     | Support plate, sample syringe                         |
| 2    | XDA613CS    | Valve, liquid, 2-valve assembly (17-18)               |
| 3    | KAA002A     | Screw, hex M3x6                                       |
| 4    |             | Sample Syringe and Motor Assembly (See Figure 8.2-20) |

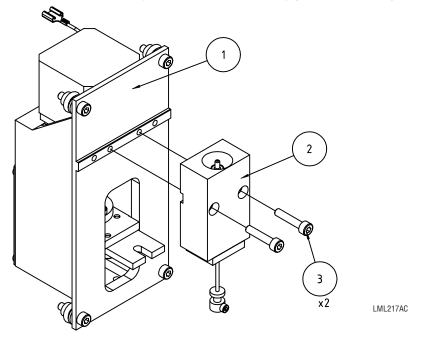


Figure 8.2-20 Sample Syringe and Motor Assembly (See Table 8.2-20)

Table 8.2-20 Sample Syringe and Motor Assembly (See Figure 8.2-20)

| ltem | Part Number | Description  |
|------|-------------|--|
| 1    | XDA596AS    | Motor, assembly - for sample syringe (See Figure 8.2-22) |
| 2    | XDA593AS    | Syringe, sampling assembly (See Figure 8.2-21)           |
| 3    | KAA028A     | Screw, hex M4x20   |

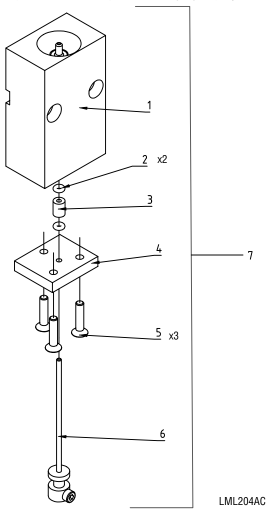


Figure 8.2-21 Sample Assembly Syringe (See Table 8.2-21)

Table 8.2-21 Sample Assembly Syringe (See Figure 8.2-21)

| ltem | Part Number | Description                           |
|------|-------------|---------------------------------------|
| 1    | GBG044A     | Syringe, sample, body                 |
| 2    | FAA064A     | O-ring, sample syringe, package of 10 |
| 3    | GBG048A     | Syringe, sample, O-ring collar        |
| 4    | GBG045A     | Syringe top, sampling                 |
| 5    | KAB017A     | Screw, torx M3x12                     |
| 6    | XDA617AS    | Piston, sample syringe, 100 µL needle |
| 7    | XDA593AS    | Syringe, sampling assembly            |

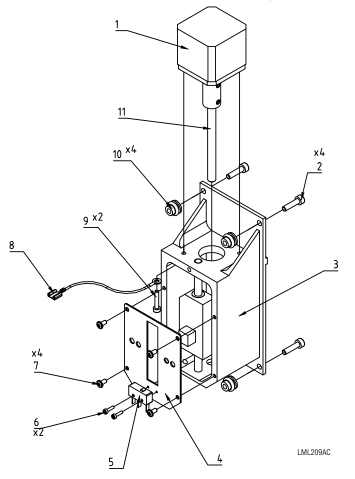
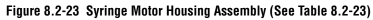


Figure 8.2-22 Sample Syringe Motor Assembly (See Table 8.2-22)

Table 8.2-22 Sample Syringe Motor Assembly (See Figure 8.2-22)

| ltem | Part Number | Description  |
|------|-------------|--|
| 1    | XBA390A     | Motor - for syringe assembly                             |
| 2    | KAA017A     | Screw, hex M4x16   |
| 3    |             | Syringe Motor Housing Assembly (See Figure 8.2-23)       |
| 4    | GBG027A     | Plate, microswitch, motor                                |
| 5    | CAE010A     | Switch, microswitch XC5-81                               |
| 6    | KAA040A     | Screw, hex M2x8  |
| 7    | KZZ022A     | Screw, auto-threaded                                     |
| 8    |             | Grounding wire   |
| 9    | KAA007A     | Screw, hex M3x16   |
| 10   | FAL009A     | Nut, shock mount, package of 12 - for most subassemblies |
| 11   | GBG025A     | Syringe screw  |



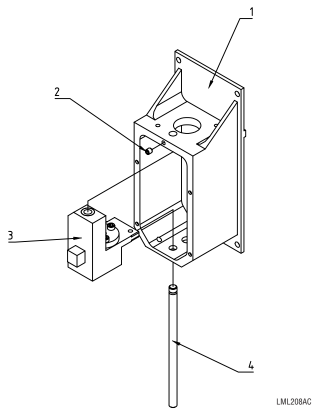


Table 8.2-23 Syringe Motor Housing Assembly (See Figure 8.2-23)

| ltem | Part Number | Description                                   |
|------|-------------|---|
| 1    | GBG028A     | Motor housing, syringe                        |
| 2    | KAD016A     | Screw, HC M4x6                                |
| 3    |             | Syringe Motor Guide Block (See Figure 8.2-24) |
| 4    | GBG030A     | Motor guide, assembly                         |

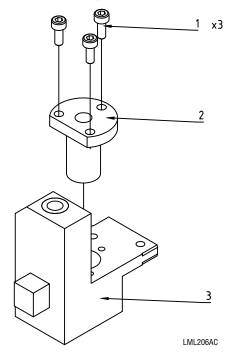


Figure 8.2-24 Syringe Motor Guide Block Assembly (See Table 8.2-24)

Table 8.2-24 Syringe Motor Guide Block Assembly (See Figure 8.2-24)

| ltem | Part Number | Description                |
|------|-------------|----------------------------|
| 1    | KAA003A     | Screw, hex M3x8            |
| 2    | GBG029A     | Syringe nut                |
| 3    | GBG031A     | Motor guide block, syringe |

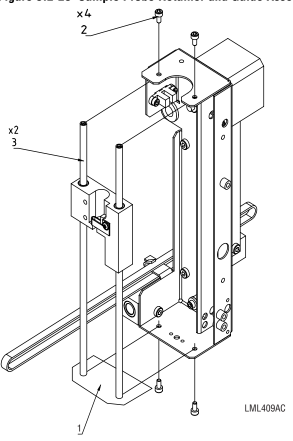


Figure 8.2-25 Sample Probe Retainer and Guide Assembly (See Table 8.2-25)

 Table 8.2-25
 Sample Probe Retainer and Guide Assembly (See Figure 8.2-25)

| ltem | Part Number | Description            |
|------|-------------|------------------------|
| 1    | GBG162A     | Centering sheet, steel |
| 2    | KAA002A     | Screw, hex M3x6        |
| 3    | GBG081A     | Needle axis            |

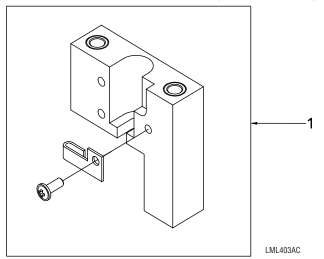


Figure 8.2-26 Sample Probe Retainer (See Table 8.2-26)

| Table 8 2-26 | Sample Probe  | Retainer (S | See Figure  | 8 2-26) |
|--------------|---------------|-------------|-------------|---------|
|              | oampic i iouc | metanier (t | occ i iyuic | 0.2-20) |

| Item | Part Number | Description                  |
|------|-------------|------------------------------|
| 1    | XDA618AS    | Guide, sample probe retainer |

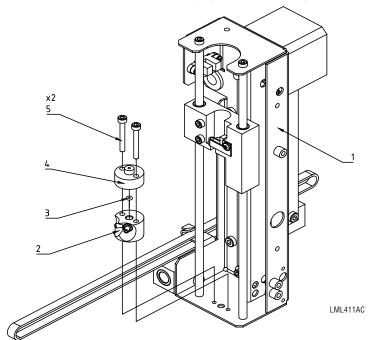


Figure 8.2-27 Rinse Block Assembly (See Table 8.2-27)

Table 8.2-27 Rinse Block Assembly (See Figure 8.2-27)

| ltem | Part Number | Description   |
|------|-------------|---|
| 1    |             | Traverse vertical movement components - Home sensor (See Figure 8.2-33) |
| 2    | GBG090A     | Sample probe, rinse block   |
| 3    | FAA053A     | O-ring, probe rinse block, package of 10                                |
| 4    | GBG091A     | Sample probe, guide   |
| 5    | KAA010A     | Screw, hex M3x25  |

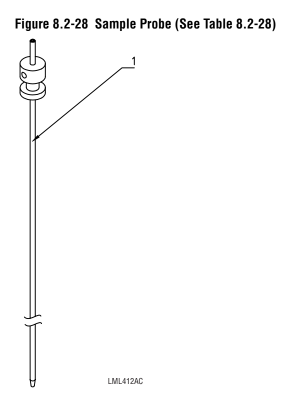


Table 8.2-28 Sample Probe (See Figure 8.2-28)

| ltem | Part Number | Description   |
|------|-------------|---------------|
| 1    | XDA619AS    | Probe, sample |

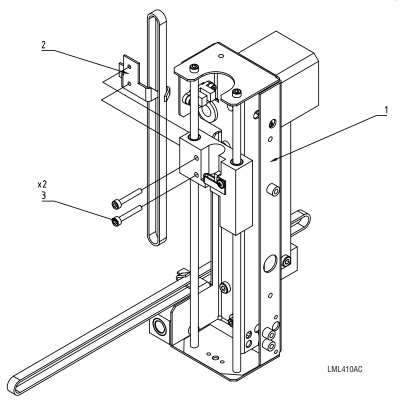


Figure 8.2-29 Traverse Vertical Movement Components - Belt Retainer (See Table 8.2-29)

Table 8.2-29 Traverse Vertical Movement Components - Belt Retainer (See Figure 8.2-29)

| ltem | Part Number | Description   |
|------|-------------|---|
| 1    |             | Traverse vertical movement components - Home sensor (See Figure 8.2-33) |
| 2    |             | Belt, probe vertical, assembly, 364 tooth                               |
| 3    | KAA009A     | Screw, hex M3x20  |

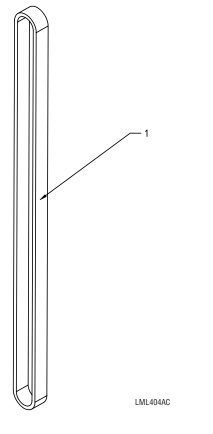


Figure 8.2-30 Vertical Traverse Vertical Movement Components - Belt (See Table 8.2-30)

| ltem | Part Number | Description                     |
|------|-------------|---------------------------------|
| 1    | FBR011A     | Belt, probe vertical, 364 tooth |

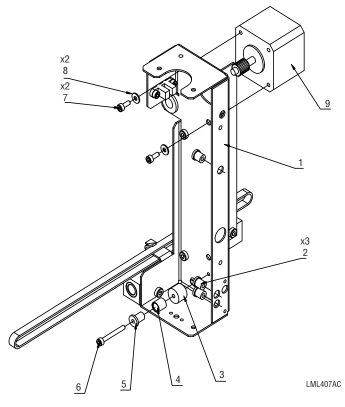
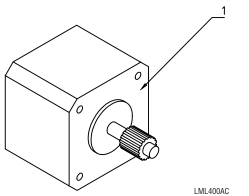
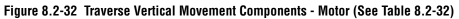


Figure 8.2-31 Traverse Vertical Movement Components - Motor and Pulley (See Table 8.2-31)

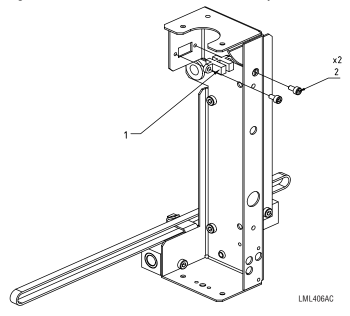
Table 8.2-31 Traverse Vertical Movement Components - Motor and Pulley (See Figure 8.2-31)

| ltem | Part Number | Description   |
|------|-------------|---|
| 1    |             | Traverse vertical movement components - Home sensor (See Figure 8.2-33) |
| 2    | DBE014A     | Wire guide  |
| 3    | GBC147A     | Pulley holder   |
| 4    | GBG093A     | Pulley, belt freewheel  |
| 5    | FAG011A     | Bearing   |
| 6    | KAA010A     | Screw, hex M3x25  |
| 7    | KAA002A     | Screw, hex M3x6   |
| 8    | KAJ001A     | Washer, M diameter 0.3  |
| 9    | XBA392A     | Motor, sample probe, vertical   |





| ltem | Part Number | Description                   |
|------|-------------|-------------------------------|
| 1    | XBA392A     | Motor, sample probe, vertical |



## Figure 8.2-33 Traverse Vertical Movement Components - Home Sensor (See Table 8.2-33)

Table 8.2-33 Traverse Vertical Movement Components - Home Sensor (See Figure 8.2-33)

| ltem | Part Number | Description                         |
|------|-------------|-------------------------------------|
| 1    | XBA396A     | Sensor, home, sample probe vertical |
| 2    | KAA002AA    | Screw, hex M3x6                     |

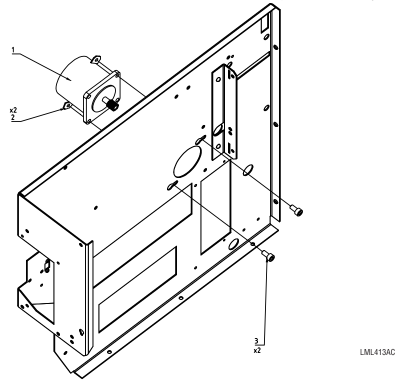


Figure 8.2-34 Traverse Horizontal Movement Components - Motor (See Table 8.2-34)

Table 8.2-34 Traverse Horizontal Movement Components - Motor (See Figure 8.2-34)

| ltem | Part Number | Description                 |
|------|-------------|-----------------------------|
| 1    | XBA391A     | Motor, traverse, horizontal |
| 2    | GBG095A     | Nut, plate M5               |
| 3    | KAA021A     | Screw, hex M5x10            |

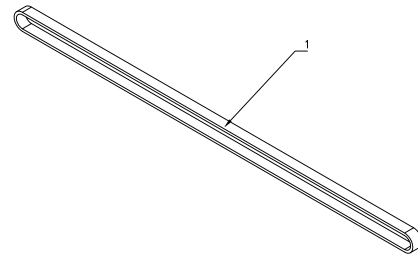


Figure 8.2-35 Traverse Horizontal Movement Components - Belt (See Table 8.2-35)t

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Table 8.2-35 Traverse Horizontal Movement Components - Belt (See Figure 8.2-35)

| ltem | Part Number | Description                          |
|------|-------------|--------------------------------------|
| 1    | FBR012A     | Belt, traverse horizontal, 544 tooth |

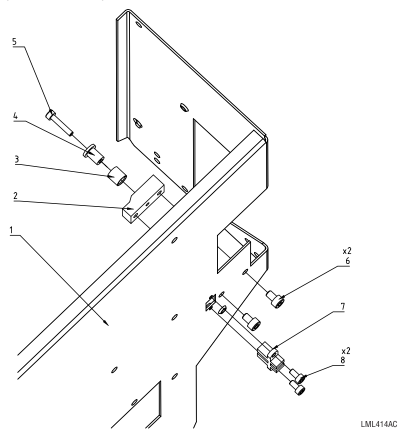


Figure 8.2-36 Traverse Horizontal Movement Components - Free Wheel and Home Sensor (See Table 8.2-36)

| Table 8.2-36 Traverse Horizontal Movement Components - Free Wheel and Home | e Sensor |
|--|----------|
| (See Figure 8.2-36)  |          |

| ltem | Part Number | Description                       |
|------|-------------|-----------------------------------|
| 1    |             | Frame assembly                    |
| 2    | GBG146A     | Freewheel strengthener            |
| 3    | GBG093A     | Pulley, belt freewheel            |
| 4    | FAG011A     | Bearing                           |
| 5    | KAA007A     | Screw, hex M3x16                  |
| 6    | KAA013A     | Screw, hex M4x6                   |
| 7    | XBA394A     | Sensor, home, traverse horizontal |
| 8    | KAA002A     | Screw, hex M3x6                   |

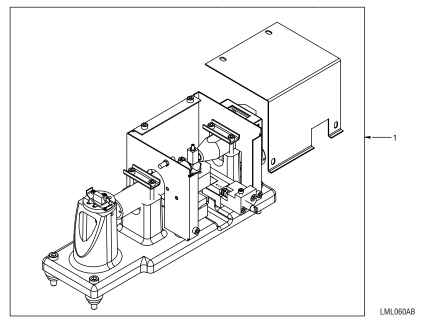


Figure 8.2-37 Optical Bench Assembly (See Table 8.2-37)

Table 8.2-37 Optical Bench Assembly (See Figure 8.2-37)

| ltem | Part Number | Description            |
|------|-------------|------------------------|
| 1    | XDA600AS    | Optical bench assembly |



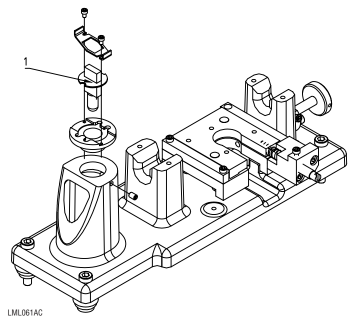


Table 8.2-38 Optical Bench Lamp (See Figure 8.2-38)

| ltem | Part Number | Description        |
|------|-------------|--------------------|
| 1    | DAJ007A     | Optical bench lamp |

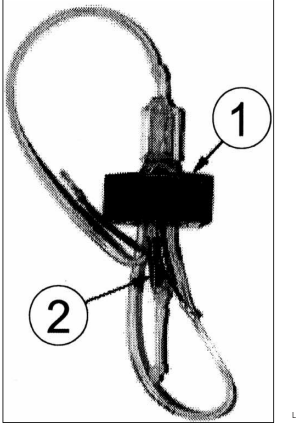
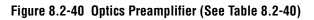


Figure 8.2-39 DIFF Flow Cell Assembly (See Table 8.2-39)

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Table 8.2-39 DIFF Flow Cell Assembly (See Figure 8.2-39)

| ltem | Part Number | Description                     |
|------|-------------|---------------------------------|
| 1    | XDA601AS    | Flow cell, assembly             |
| 2    | XBA403A     | Shield, flow cell tubing ground |



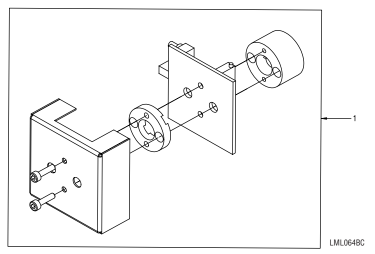
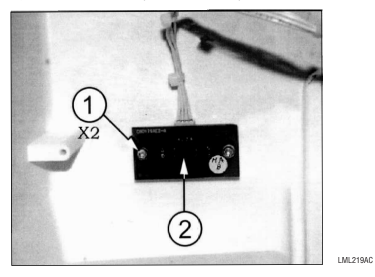


Table 8.2-40 Optics Preamplifier (See Figure 8.2-40)

| ltem | Part Number | Description                     |
|------|-------------|---------------------------------|
| 1    | XAA423BS    | Preamp, optical signal assembly |

# Figure 8.2-41 LED Card (See Table 8.2-41)



# Table 8.2-41 LED Card (See Figure 8.2-41)

| ltem | Part Number | Description                         |
|------|-------------|-------------------------------------|
| 1    | KZZ022A     | Screw, auto-threaded                |
| 2    | XAA468A     | PCB, LED card, aspiration indicator |

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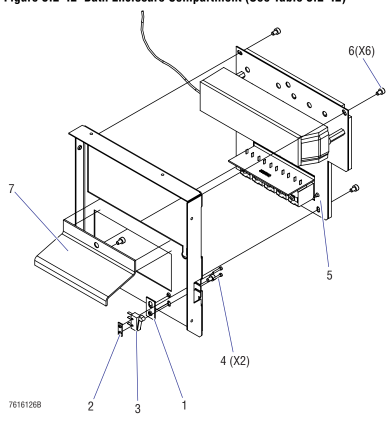


Figure 8.2-42 Bath Enclosure Compartment (See Table 8.2-42)

Table 8.2-42 Bath Enclosure Compartment (See Figure 8.2-42)

| ltem | Part Number | Description                            |
|------|-------------|--|
| 1    | GBG020A     | Nut, plate, bath enclosure compartment |
| 2    | GBD498A     | Nut, plate, microswitch                |
| 3    | CAE006A     | Switch, microswitch XC5-81-82          |
| 4    | KAA041A     | Screw, hex M2x12                       |
| 5    | KZZ022A     | Screw, auto-threaded                   |
| 6    | KAA013A     | Screw, hex M4x6                        |
| 7    | GBG205A     | Cover, solenoid drivers, bank 27-31    |

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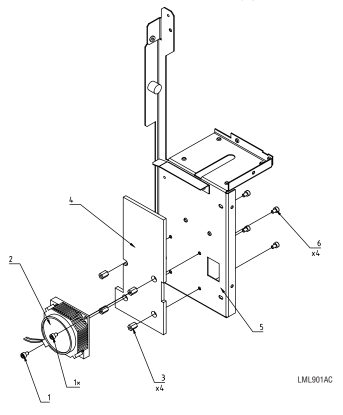


Figure 8.2-43 Bath Enclosure Fan Assembly (See Table 8.2-43)

Table 8.2-43 Bath Enclosure Fan Assembly (See Figure 8.2-43)

| ltem | Part Number | Description                             |
|------|-------------|---|
| 1    | KAA013A     | Screw, hex M4x6                         |
| 1x   | KAA015A     | Screw, hex M4x8                         |
| 2    | XEA487AS    | Fan, bath enclosure temperature control |
| 3    | KAN023A     | Collar                                  |
| 4    | GBG019A     | Insulated plate                         |
| 5    | GBG063A     | Steel                                   |
| 6    | KAA013A     | Screw, hex M4x6                         |

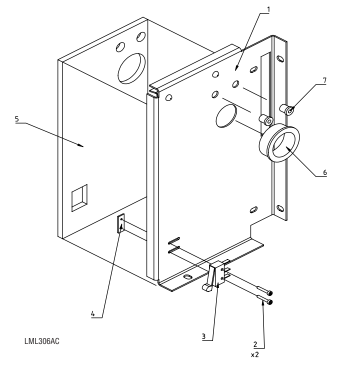
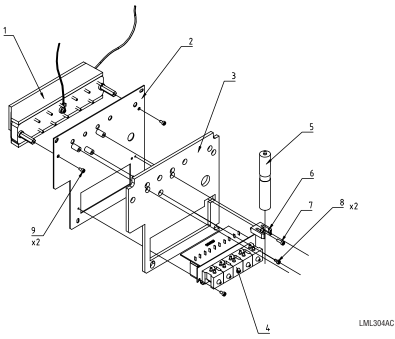


Figure 8.2-44 Bath Enclosure Door Interlock (See Table 8.2-44)

Table 8.2-44 Bath Enclosure Door Interlock (See Figure 8.2-44)

| ltem | Part Number | Description                           |
|------|-------------|---------------------------------------|
| 1    | GBG012A     | Backplate, bath enclosure             |
| 2    | KAA041A     | Screw, hex M2x12                      |
| 3    | CAE006A     | Switch, microswitch XC5-81-82         |
| 4    | GBD498A     | Nut, microswitch                      |
| 5    | GBG018A     | Plate, bath enclosure, rear isolation |
| 6    | DBE026A     | Thrust                                |
| 7    | FAM006A     | Bushing                               |



# Figure 8.2-45 Reagent Heating Coil Assembly (See Table 8.2-45)

Table 8.2-45 Reagent Heating Coil Assembly (See Figure 8.2-45)

| ltem | Part Number | Description                                       |  |
|------|-------------|---|--|
| 1    | XDA625AS    | Heater, reagent coil assembly                     |  |
| 2    | GBG009A     | Holder clip, chamber                              |  |
| 3    | GBG014A     | Chamber plate, insulated                          |  |
| 4    | XDA615CS    | Valve, liquid, 5-valve assembly (27-31)           |  |
| 5    | XCA166A     | Chamber, isolator (long)                          |  |
| 6    | GBC015A     | Clip, long isolator chamber holder, package of 10 |  |
| 7    | KAA003A     | Screw, hex M3x8                                   |  |
| 8    | KAA002A     | Screw, hex M3x6                                   |  |
| 9    | KAA013A     | Screw, hex M4x6                                   |  |

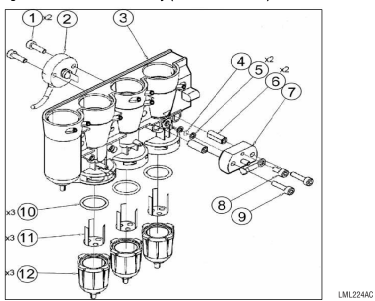


Figure 8.2-46 Baths Assembly (See Table 8.2-46)

Table 8.2-46 Baths Assembly (See Figure 8.2-46)

| ltem | Part Number | Description  |  |
|------|-------------|--|--|
| 1    | KAA004A     | Screw, hex M3x10                                   |  |
| 2    | XBA398B     | Cable, coaxial, with RBC/WBC bath electrode        |  |
| 3    | GBG001A     | Four-bath unit                                     |  |
| 4    | FAK001A     | Aperture, RBC/Plt, 50 μ                            |  |
| 5    | GBG156A     | O-ring, aperture, package of 12                    |  |
| 6    | GBG004A     | Collar   |  |
| 7    | GBG157A     | Counting head (internal electrode)                 |  |
| 8    | KAA002A     | Screw, hex M3x6                                    |  |
| 9    | KAA005A     | Screw, hex M3x12                                   |  |
| 10   | FAA066A     | O-ring, bath drain/debubble chamber, package of 12 |  |
| 11   | GBG007A     | Diffuser, drain and debubble, set of 4             |  |
| 12   | GBG003A     | Chamber, drain and debubble                        |  |

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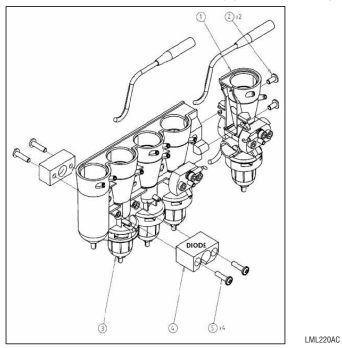


Figure 8.2-47 Hgb Photometer Assembly (See Table 8.2-47)

Table 8.2-47 Hgb Photometer Assembly (See Figure 8.2-47)

| ltem | Part Number | Description                               |  |
|------|-------------|---|--|
| 1    | XDA610A     | Bath, WBC/BASO                            |  |
| 2    | KZZ022A     | Screw, auto-threaded                      |  |
| 3    | XDA602A     | Bath, assembly, 3-baths and rinse chamber |  |
| 4    | XBA389A     | Photometer, Hgb diode and preamp assembly |  |
| 5    | KZZ026A     | Screw, auto-threaded                      |  |

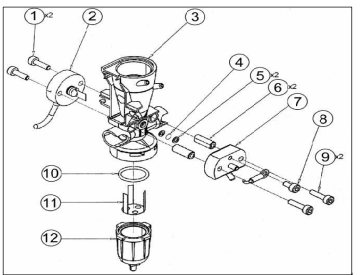


Figure 8.2-48 WBC/BASO Bath Assembly (See Table 8.2-48)

| ltem | Part Number | Description  |  |
|------|-------------|--|--|
| 1    | KAA004A     | Screw, hex M3x10                                   |  |
| 2    | XBA398B     | Cable, coaxial, with RBC/WBC bath electrode        |  |
| 3    | GBG002A     | Bath, WBC/Hgb/BASO                                 |  |
| 4    | FAK003A     | Aperture, WBC/BASO, 80 µm                          |  |
| 5    | GBG156A     | O-ring, aperture, package of 12                    |  |
| 6    | GBG004A     | Collar   |  |
| 7    | GBG157A     | Counting head (internal electrode)                 |  |
| 8    | KAA002A     | Screw, hex M3x6                                    |  |
| 9    | KAA005A     | Screw, hex M3x12                                   |  |
| 10   | FAA066A     | O-ring, bath drain/debubble chamber, package of 12 |  |
| 11   | GBG007A     | Diffuser, drain and debubble, set of 4             |  |
| 12   | GBG003A     | Chamber, drain and debubble                        |  |

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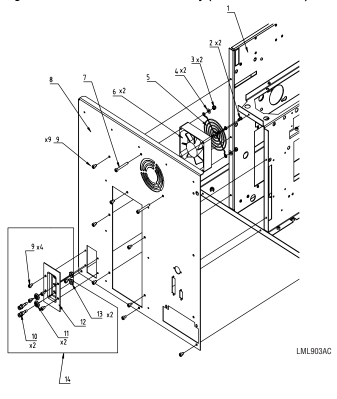


Figure 8.2-49 Rear Frame Assembly (See Table 8.2-49)

Table 8.2-49 Rear Frame Assembly (Figure 8.2-49)

| ltem | Part Number | Description                                    |  |
|------|-------------|--|--|
| 1    |             | Frame assembly                                 |  |
| 2    | DBK021A     | Rivets   |  |
| 3    | KAH018A     | Nut, M4  |  |
| 4    | KAJ002A     | Washer, M diameter 0.4                         |  |
| 5    | FAN001A     | Fan protector                                  |  |
| 6    | XBA393A     | Fan, main chassis, 24V                         |  |
| 7    | KAA031A     | Screw, hex M4x35                               |  |
| 8    | GBG058A     | Plate, rear frame                              |  |
| 9    | KAA013A     | Screw, hex M4x6                                |  |
| 10   | EAC010A     | Fitting, Luer, female, i.d. 3 mm, package of 5 |  |
| 11   | EAC008A     | Fitting, antirotation washer, package of 5     |  |
| 12   | GBG071A     | Receptacle                                     |  |
| 13   | KAH024A     | Nut 1/4-28 (US std)                            |  |
| 14   | XDA623AS    | Panel, reagent/waste fittings assembly         |  |

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# DN A

# A.1 TOLERANCE AND LIMITS

## Table A.1-1 Flow Cell Adjustment Limits

| Parameter          | Target Value | Acceptable Range |
|--------------------|--------------|------------------|
| DIFF LAMP          | 6.00         | 5.50 to 6.50     |
| TRANSFER TIME      | 200          | 150 to 250       |
| RESISTIVE CHANNEL  | 50           | 45 to 55         |
| ABSORBANCE CHANNEL | 180          | 170 to 190       |

### Table A.1-2 Motor Voltage Limits

| Motor                          | Test Point | Voltage      | Potentiometer |
|--------------------------------|------------|--------------|---------------|
| Waste syringe                  | TP5        | 4 V ± 0.05 V | R149          |
| Count syringe                  | TP6        | 4 V ± 0.05 V | R150          |
| Dilutor syringe                | TP7        | 4 V ± 0.05 V | R151          |
| Optical bench injector syringe | TP8        | 3 V ± 0.05 V | R152          |
| Horizontal traverse            | TP10       | 3 V ± 0.05 V | R154          |
| Sample syringe                 | TP11       | 2 V ± 0.05 V | R155          |
| Probe carriage                 | TP12       | 5 V ± 0.05 V | R156          |

#### Table A.1-3 Thresholds Voltage Limits

| Threshold       | Test Point | Voltage    | Potentiometer |
|-----------------|------------|------------|---------------|
| BASO            | TP14       | 300 mV ± 5 | R157          |
| RBC             | TP13       | 300 mV ± 5 | R158          |
| PLT             | TP2        | 300 mV ± 5 | R159          |
| LMNE (DIFF) CIS | TP3        | 650 mV ± 5 | R160          |
| LMNE (DIFF) OD  | TP4        | 350 mV ± 5 | R161          |

## Table A.1-4 Mixing Bubble Limits

| Mixing Bath    | Low Limit | Normal | High Limit |
|----------------|-----------|--------|------------|
| FIRST DILUTION | 100       | 300    | 400        |
| DIFF           | 100       | 300    | 400        |
| WBC/BASO       | 100       | 300    | 400        |
| HGB LYSE       | 300       | 400    | 500        |

| Test Point | Designation       | Target Value |
|------------|-------------------|--------------|
| TP40       | 5V Power supply   | +5V          |
| TP41       | 12V Power supply  | +12V         |
| TP42       | 24V Power supply  | +24V         |
| TP43       | -12V Power supply | -12V         |

Table A.1-5 Power Supply Voltages

## Table A.1-6 Whole-Blood Reproducibility CV Limits for 20 Cycles

| Parameters | %CV | Test Level                           |
|------------|-----|--------------------------------------|
| WBC        | <2% | at 10.0 x 10 <sup>3</sup> cells / µL |
| RBC        | <2% | at 5.00 x 10 <sup>6</sup> cells / µL |
| Hgb        | <1% | at 15.0 g/dL                         |
| Hct        | <2% | at 45.0%                             |
| MCV        | <1% | at 90.0 fL                           |
| Plt        | <5% | at 300 x 10 <sup>3</sup> cells / µL  |

Table A.1-7 Calibration Factor Limits

| Parameter | Target Value | Minimum<br>Acceptable Value | Maximum<br>Acceptable Value |
|-----------|--------------|-----------------------------|-----------------------------|
| WBC       | 137          | 90                          | 200                         |
| RBC       | 225          | 160                         | 290                         |
| Hgb       | 40.0         | 25.0                        | 55.0                        |
| Hct       | 220          | 160                         | 290                         |
| PLT       | 290          | 180                         | 400                         |
| RDW       | 0.3          | 0.1                         | 0.9                         |

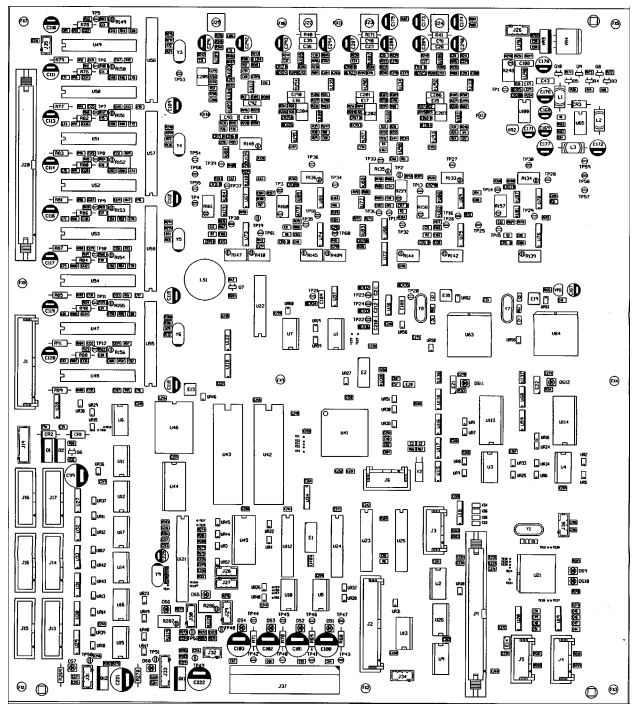


# A.2 CIRCUIT CARD LAYOUTS WITH KEY COMPONENT DESCRIPTIONS

#### **Main Card**

#### **Component Locations**

Figure A.2-1 Main Card Components



# **QUICK REFERENCE INFORMATION** *CIRCUIT CARD LAYOUTS WITH KEY COMPONENT DESCRIPTIONS*

#### **Test Points**

#### Table A.2-1 Main Card Test Points

| Test Point | Designation                          | Potentiometer | Target | Tolerance | Remarks          |
|------------|--------------------------------------|---------------|--------|-----------|------------------|
| TP1        | HB Gain adjustment                   | R248          | 4.7 mV | ±0.1 mV   |                  |
| TP2        | PLT Threshold adjustment             | R159          | 300 mV | ±5 mV     |                  |
| TP3        | LMNE (DIFF) CIS Threshold adjustment | R160          | 650 mV | ±5 mV     |                  |
| TP4        | LMNE (DIFF) OD Threshold adjustment  | R161          | 350 mV | ±5 mV     |                  |
| TP5        | Waste syringe motor                  | R149          | 3 V    | ±50mV     |                  |
| TP6        | Count syringe motor                  | R150          | 3 V    | ±50mV     |                  |
| TP7        | Reagent syringes motor               | R151          | 3 V    | ±50mV     |                  |
| TP8        | Injection syringe motor              | R152          | 3 V    | ±50mV     |                  |
| TP9        | Not used                             |               |        |           |                  |
| TP10       | Horizontal traverse motor            | R154          | 3 V    | ±50mV     |                  |
| TP11       | Sample syringe motor                 | R155          | 2 V    | ±50mV     |                  |
| TP12       | Vertical traverse motor              | R156          | 3 V    | ±50mV     |                  |
| TP13       | RBC Threshold adjustment             | R158          | 300 mV | ±5mV      |                  |
| TP14       | BASO Threshold adjustment            | R157          | 300 mV | ±5mV      |                  |
| TP15       | BASO Comparator                      |               |        |           |                  |
| TP16       | RBC Comparator                       |               |        |           |                  |
| TP17       | PLT Comparator                       |               |        |           |                  |
| TP18       | LMNE (DIFF) CIS Comparator           |               |        |           |                  |
| TP19       | LMNE (DIFF) OD Comparator            |               |        |           |                  |
| TP20       | RBC line height adjustment           | R133          | XX V   | ±50 mV    | Factory adjusted |
| TP21       | BASO gain adjustment                 | R134          | XX V   | ±50 mV    | Factory adjusted |
| TP22       | PLT line height adjustment           | R135          | XX V   | ±50 mV    | Factory adjusted |
| TP23       | DIFF Resistive gain adjustment       | R136          | XX V   | ±50 mV    | Factory adjusted |
| TP24       | DIFF Optical gain adjustment         | R148          | XX V   | ±50 mV    | Factory adjusted |
| TP25       | RBC line width adjustment            |               |        |           |                  |
| TP26       | RBC line reject adjustment           | R142          | 10 µs  | ±0.5 μs   | Factory adjusted |
| TP27       | RBC line pulse adjustment            |               |        |           |                  |
| TP28       | BASO line width adjustment           |               |        |           |                  |
| TP29       | BASO line reject adjustment          | R139          | 10 µs  | ±0.5 µs   | Factory adjusted |
| TP30       | BASO line pulse adjustment           |               |        |           |                  |
| TP31       | PLT line width adjustment            |               |        |           |                  |
| TP32       | PLT line reject adjustment           | R144          | 10 µs  | ±0.5 μs   | Factory adjusted |
| TP33       | PLT line pulse adjustment            |               |        |           |                  |
| TP34       | DIFF Resistive line width adjustment |               |        |           |                  |



| Test Point | Designation  | Potentiometer | Target | Tolerance | Remarks          |
|------------|--|---------------|--------|-----------|------------------|
| TP35       | DIFF Resistive line reject adjustment                | R145          | 15 µs  | ±0.5 μs   | Factory adjusted |
| TP36       | DIFF Resistive line pulse adjustment                 |               |        |           |                  |
| TP37       | DIFF Optical line width adjustment                   |               |        |           |                  |
| TP38       | DIFF Optical line reject adjustment                  | R147          | 5 µs   | ±0.5 μs   | Factory adjusted |
| TP39       | DIFF Optical line pulse adjustment                   |               |        |           |                  |
| TP40       | 5V Power supply                                      |               | 5 V    |           |                  |
| TP41       | 12V Power supply                                     |               | 12 V   |           |                  |
| TP42       | 24V Power supply                                     |               | 24 V   |           |                  |
| TP43       | -12V Power supply                                    |               | -12 V  |           |                  |
| TP44       | GROUND   |               |        |           |                  |
| TP45       | GROUND   |               |        |           |                  |
| TP46       | GROUND   |               |        |           |                  |
| TP47       | GROUND   |               |        |           |                  |
| TP48       | Waste sensor   | R286          | <1 V   |           | With water       |
| TP49       | Not used   |               |        |           |                  |
| TP50       | Reagent heating system temperature voltage           |               |        |           |                  |
| TP51       | Thermostated room heating system temperature voltage |               |        |           |                  |
| TP52       | DIFF drain sensor                                    | R287          | <1 V   |           | With water       |
| TP53       | DIFF Lamp  | R11           | 6 V    |           |                  |
| TP54       | GROUND   |               |        |           |                  |
| TP55       | GROUND   |               |        |           |                  |
| TP56       | GROUND   |               |        |           |                  |
| TP57       | GROUND   |               |        |           |                  |
| TP58       | GROUND   |               |        |           |                  |
| TP59       | GROUND   |               |        |           |                  |
| TP60       | DIFF Resistive line reject2 adjustment               | R409          | 50µs   | ±2 μs     | Factory adjusted |
| TP61       | DIFF Resistive line reject2 adjustment               | R410          | 250µs  | ±5 μs     | Factory adjusted |

## Table A.2-1 Main Card Test Points (Continued)

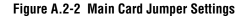
#### Potentiometers

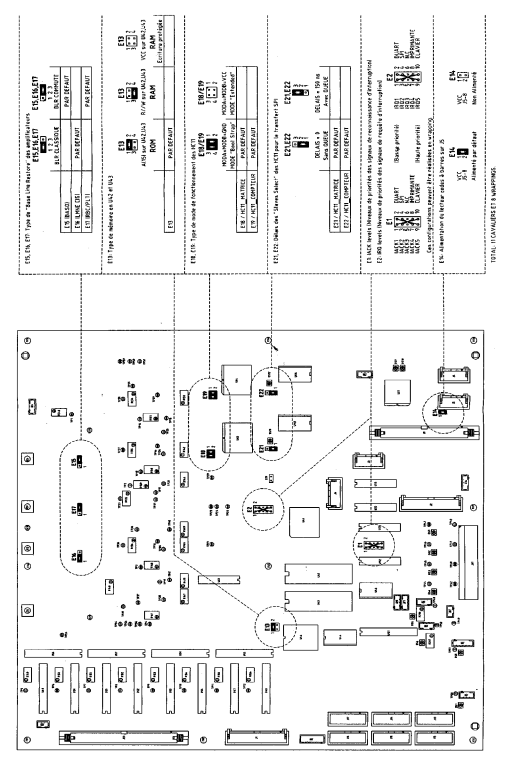
#### Table A.2-2 Main Card Potentiometers

| Potentiometer | Designation                           | Test Point | Target | Tolerance | Remarks          |
|---------------|---------------------------------------|------------|--------|-----------|------------------|
| R133          | RBC line height adjustment            | TP20       |        |           | Factory adjusted |
| R134          | BASO gain adjustment                  | TP21       |        |           | Factory adjusted |
| R135          | PLT line height adjustment            | TP22       |        |           | Factory adjusted |
| R136          | DIFF Resistive gain adjustment        | TP23       |        |           | Factory adjusted |
| R139          | BASO line reject adjustment           | TP29       |        |           | Factory adjusted |
| R142          | RBC line reject adjustment            | TP26       |        |           | Factory adjusted |
| R144          | PLT line reject adjustment            | TP32       |        |           | Factory adjusted |
| R145          | DIFF Resistive line reject adjustment | TP35       |        |           | Factory adjusted |
| R147          | DIFF Optical line reject adjustment   | TP38       |        |           | Factory adjusted |
| R148          | DIFF Optical gain adjustment          | TP24       |        |           | Factory adjusted |
| R149          | Waste syringe motor                   | TP5        | 3 V    | ±50 mV    |                  |
| R150          | Count syringe motor                   | TP6        | 3 V    | ±50 mV    |                  |
| R151          | Reagent syringes motor                | TP7        | 3 V    | ±50 mV    |                  |
| R152          | Injection syringe motor               | TP8        | 3 V    | ±50 mV    |                  |
| R154          | Horizontal traverse motor             | TP10       | 3 V    | ±50 mV    |                  |
| R155          | Sample syringe motor                  | TP11       | 2 V    | ±50 mV    |                  |
| R156          | Vertical traverse motor               | TP12       | 3 V    | ±50 mV    |                  |
| R157          | BASO Threshold adjustment             | TP14       | 300 mV | ±5 mV     |                  |
| R158          | RBC Threshold adjustment              | TP13       | 300 mV | ±5 mV     |                  |
| R159          | PLT Threshold adjustment              | TP2        | 300 mV | ±5 mV     |                  |
| R160          | LMNE (DIFF) CIS Threshold adjustment  | TP3        | 650 mV | ±5 mV     |                  |
| R161          | LMNE (DIFF) OD Threshold adjustment   | TP4        | 350 mV | ±5 mV     |                  |
| R248          | HB Gain adjustment                    | TP1        | 4.7 mV | ±0.1 mV   |                  |

A

#### **Jumper Settings**



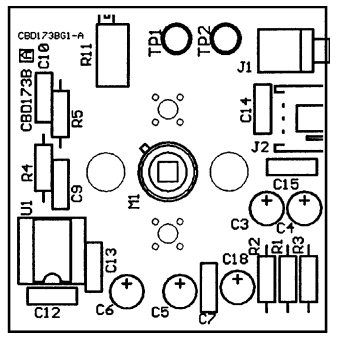


Note: If you have an initialization problem with the 68HC11 Microcontrolers, remove the E20 jumper to reset  $A^{C} \bullet T$  5diff analyzer. (The instrument must be turned ON).

### **Optical Preamplifier Card**

#### **Component Locations**

Figure A.2-3 Optical Preamplifier Card Components



#### Connectors

Table A.2-3 Connectors on the Optical Preamplifier Card

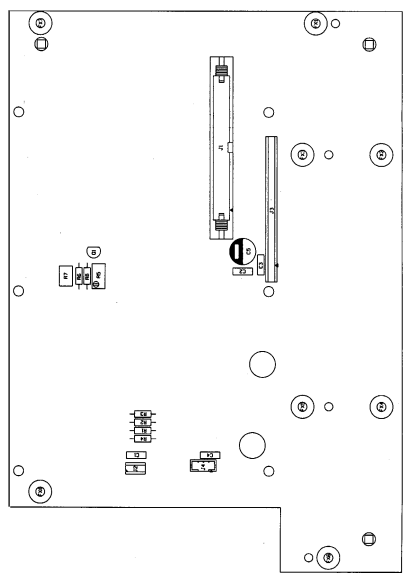
| Card Label Description |                       |
|------------------------|-----------------------|
| J1                     | Optical output signal |
| J2                     | Power to card         |



### LCD and Keypad Card

### **Component Locations**





#### Connectors

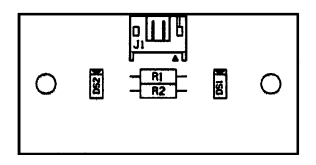
Table A.2-4 Connectors on the Keypad and LCD Card

| Card Label | Card Label Description           |  |  |
|------------|----------------------------------|--|--|
| J1         | Main interface to main card      |  |  |
| J2         | Backlight power to LCD           |  |  |
| J3         | Data out to LCD                  |  |  |
| J4         | Power to aspirate indicator card |  |  |

### LED Card

#### **Component Locations**

Figure A.2-5 LED Card Components



#### Connectors

#### Table A.2-5 Connector on the LED Card

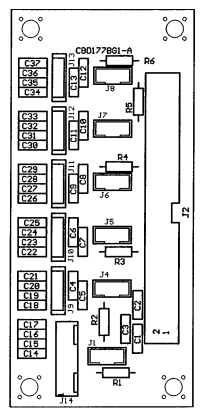
| Card Label | Description          |  |
|------------|----------------------|--|
| J1         | Power/signal to LEDs |  |



### **Motor Interconnect Card**

### **Component Locations**

Figure A.2-6 Motor Interconnect Card Components



#### Connectors

#### Table A.2-6 Connectors on the Motor Interconnect Card

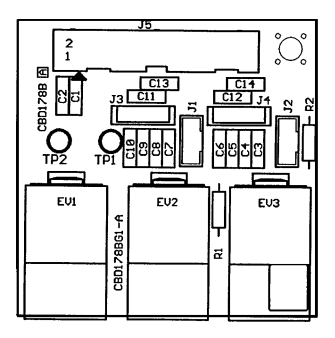
| Card Label | Description                     |
|------------|---------------------------------|
| J1         | Traverse horizontal home sensor |
| J2         | Power/signals from Main card    |
| J4         | Float sensor                    |
| J5         | Injector syringe home switch    |
| J6         | Reagent syringe home switch     |
| J7         | Count syringe home switch       |
| J8         | Waste syringe home switch       |
| J9         | Not used                        |
| J10        | Injector syringe motor          |
| J11        | Reagent syringe motor           |
| J12        | Count syringe motor             |
| J13        | Waste syringe motor             |
| J14        | Traverse horizontal motor       |

I

### **Traverse Card**

#### **Component Locations**

Figure A.2-7 Traverse Card Components



#### Connectors

| Table A.2-7 | <b>Connectors on the</b> | Traverse Card |
|-------------|--------------------------|---------------|
|             |                          | maronoo oana  |

| Card Label | Description                   |
|------------|-------------------------------|
| J1         | Traverse vertical motor       |
| J2         | Sample syringe home switch    |
| J3         | Traverse vertical home sensor |
| J4         | Sample syringe motor          |
| J5         | Power/interface to Main card  |



## A.3 AC•T 5diff MODULE LOCATIONS AND FUNCTIONS

#### Overview

Most functions are accomplished by fluidic components that are interconnected by tubing and controlled by timed solenoid signals. This section briefly describes the functions of these fluidic components and shows their locations.

#### **Analyzer Modules**

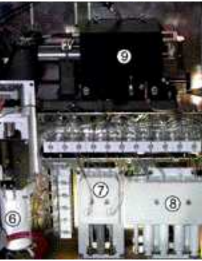
The A<sup>C</sup>•T 5diff hematology analyzer consists of nine mechanical and hydraulic modules. These modules are identified on Table A.3-1. A description of the module's primary functions is also included. Figures A.3-1 and A.3-2 show each module's location inside the instrument.

#### Mechanical and Hydraulic Modules Locations

Figure A.3-1 View of an A<sup>C</sup>•T 5diff Hematology Analyzer with the Right Side Door Open



Figure A.3-2 View of an ACoT 5diff Hematology Analyzer with the Left Side Panel Removed





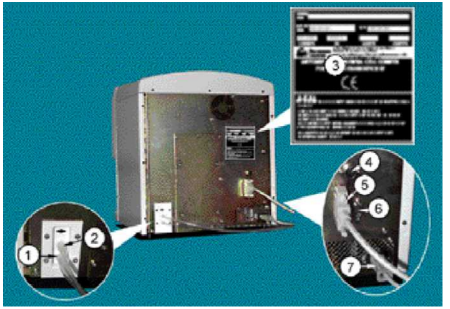
| Figure<br>Reference Module Functions |                           |  |  |
|--------------------------------------|---------------------------|--|--|
| Figure A.3-1, 1                      | Traverse Assembly         | • Ensures sample probe positioning for the different sample stages and distribution.                                 |  |
|                                      |                           | Supports the sample syringe and the distribution of the blood.   |  |
| Figure A.3-1, 2                      | Sample Syringe Assembly   | • Takes the sample and distributes the portions into the different baths.  |  |
|                                      |                           | • Removes a sample from the first dilution (made in the DIL1/HGB bath) and dispenses it into the RBC bath.           |  |
| Figure A.3-1, 3                      | Baths Assembly inside the | Receives the different rinsing and dilutions.  |  |
|                                      | Bath Enclosure            | Ensures the temperature control of the dilutions.  |  |
|                                      |                           | Ensures the counts for WBC, BASO, RBC, Plt, and Hct.   |  |
|                                      |                           | Ensures Hgb determination.   |  |
| Figure A.3-1, 4                      | Waste Syringe             | Drains the different baths.  |  |
|                                      |                           | <ul> <li>Provides mixing bubbles to the mixtures.</li> </ul>   |  |
|                                      |                           | • Provides the vacuum needed to pull the DIFF specimen from the DIFF bath towards the flow cell injector.            |  |
| Figure A.3-2, 5                      | Diluent Reservoir         | Holds the diluent needed for an analysis cycle.  |  |
|                                      |                           | • Eliminates the risk of diluent degassing as it is being aspirated by the syringes.                                 |  |
|                                      |                           | Note: Reservoir is vacuum filled by the count syringe.   |  |
| Figure A.3-1, 6                      | Count Syringe             | • Provides the vacuum needed for the WBC and the BASO counts.  |  |
|                                      |                           | Provides the vacuum needed for the RBC and PLT counts.   |  |
|                                      |                           | Provides the vacuum needed for filling the diluent reservoir.  |  |
| Figure A.3-2, 7                      | 5diff Syringe Assembly    | <ul> <li>Ensures correct proportioning of the stop diluent in the DIFF preparation bath.</li> </ul>                  |  |
|                                      |                           | Injects the sample into the flow cell.   |  |
|                                      |                           | Injects the interior and exterior gains into the flow cell.  |  |
| Figure A.3-2, 8                      | Reagent Syringes Assembly | Ensures the correct distribution of the different reagents including:  |  |
|                                      |                           | <ul> <li>Lysing agent for hemoglobin (A<sup>C</sup>•T 5diff Hgb Lyse).</li> </ul>                                    |  |
|                                      |                           | ► Cleaning reagent (A <sup>C</sup> •T 5diff Rinse).  |  |
|                                      |                           | <ul> <li>Lysing agent for the DIFF (A<sup>C</sup>•T 5diff Fix).</li> </ul>   |  |
|                                      |                           | <ul> <li>Diluent used for the dilutions (A<sup>C</sup>•T 5diff Diluent) except the DIFF<br/>stop diluent.</li> </ul> |  |
|                                      |                           | <ul> <li>Lysing agent for WBC/BASO (A<sup>C</sup>•T 5diff WBC Lyse).</li> </ul>                                      |  |
| Figure A.3-2, 9                      | Optical Bench Assembly    | Provides support and ensures adjustment of the flow cell.  |  |
|                                      |                           | • Provides support and ensures adjustment of the optics lamp.  |  |
|                                      |                           | <ul> <li>Provides support and ensures adjustment of the optical and electronic elements.</li> </ul>                  |  |

#### Table A.3-1 Mechanical and Hydraulic Modules

I

### **Rear Panel**

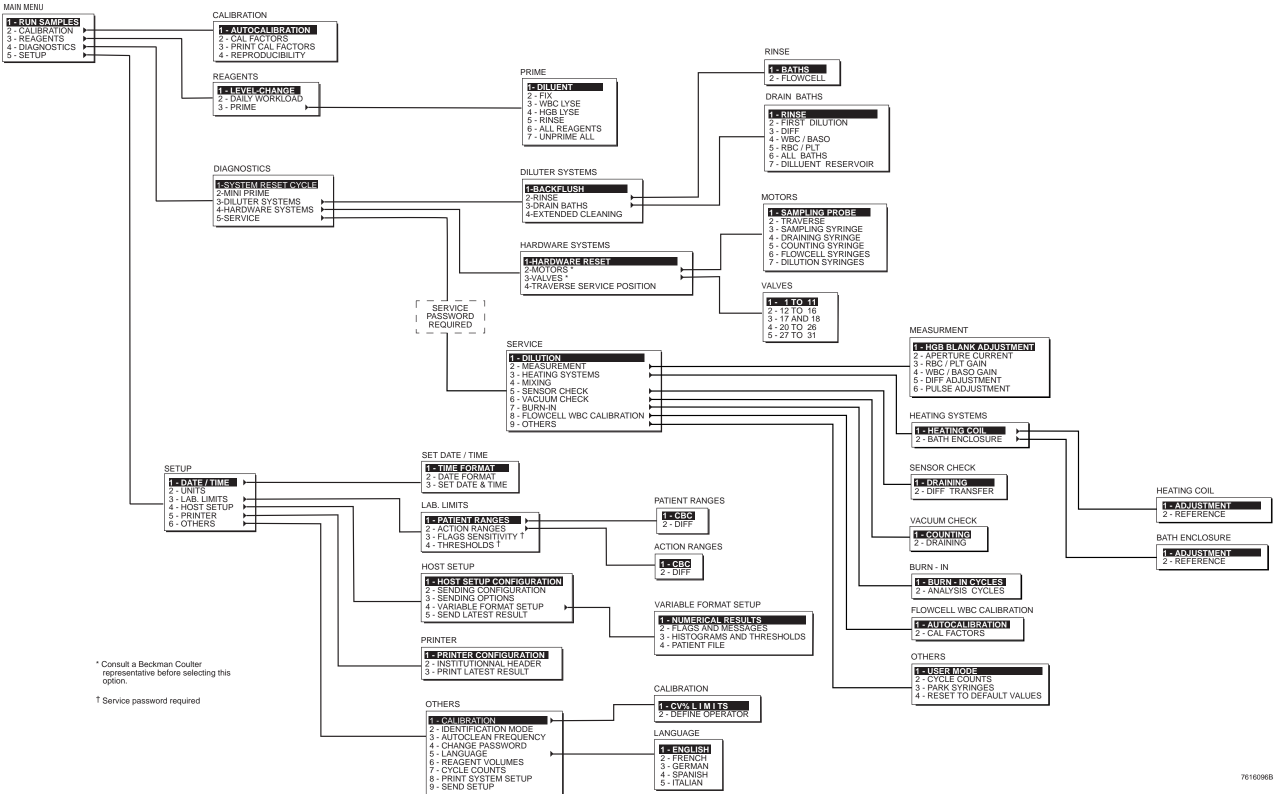




| Figure Reference | Attachments to the Rear Panel |
|------------------|-------------------------------|
| Figure A.3-3, 1  | Waste output                  |
| Figure A.3-3, 2  | Diluent input                 |
| Figure A.3-3, 3  | Serial number label           |
| Figure A.3-3, 4  | Bar-code reader connector     |
| Figure A.3-3, 5  | Printer connector             |
| Figure A.3-3, 6  | RS232C output                 |
| Figure A.3-3, 7  | Power cord                    |

#### **SOFTWARE MENU TREE** A.4

#### Figure A.4-1 Software Menu Tree



### **QUICK REFERENCE INFORMATION** SOFTWARE MENU TREE $|\mathbf{A}|$

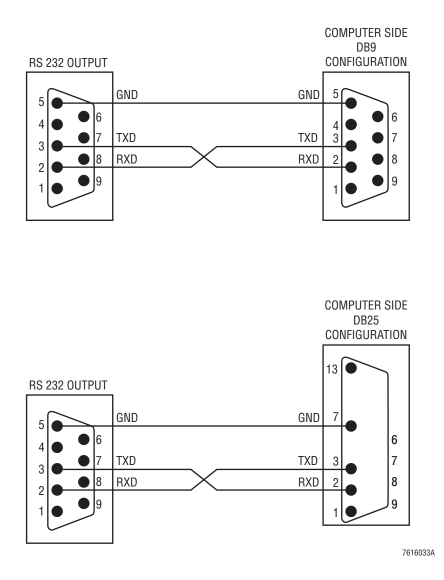
**QUICK REFERENCE INFORMATION** SOFTWARE MENU TREE

### **B.1** FORMAT

See Host Transmission Specification PN 4277065A.

**SOFTWARE INTERFACE** *FORMAT* 

### **B.2 PACKET TYPE PIN ASSIGNMENTS**





### C.1 OVERVIEW

The instrument flags a sample if its results exceed specific criteria defined within the software.

The values used to position the thresholds, which separate different cell populations and determine if a flag should be triggered, are selected to provide optimal population separation and flagging under normal operating conditions.

The software provides you with the ability to modify the values used to position the thresholds between populations and to alter the level at which a flag associated with a population is triggered.

Changing any of the values that define threshold positions or changing the flag sensitivity values from the default values will alter the parameter values reported and whether or not a flag is associated with a result.

#### **Flag Sensitivity**

A result outside the flag sensitivity range triggers a flag to appear with the parameter result. The flag occurs if either the percentage or absolute count value is exceeded.

You can adjust a flag's sensitivity for percentage and/or absolute number. Table C.1-1 shows the flag sensitivity default values.

| Flag Region                  | Percentage of Cells | Absolute Number of Cells |
|------------------------------|---------------------|--------------------------|
| DB (Debris)                  | 100                 | 120                      |
| SL (Small Lymphocytes)       | 100                 | 50                       |
| SL1 (Small Lymphocytes 1)    | 5                   | 45                       |
| NL (Neutrophil / Lymphocyte) | 3                   | 120                      |
| MN (Monocyte / Neutrophil)   | 100                 | 120                      |
| UM (Upper Monocyte)          | 1.1                 | 999                      |
| LN (Lower Neutrophil)        | 2.5                 | 999                      |
| UN (Upper Neutrophil)        | 1.1                 | 999                      |
| NE (Neutrophil / Eosinophil) | 1.1                 | 60                       |
| ATL (Atypical Lymphocytes)   | 2                   | 200                      |
| *WBC                         | 3.5                 | 999                      |
| MICRO (Microcytes)           | 5*                  | Not applicable           |
| MACRO (Macrocytes)           | 7.5*                | Not applicable           |
| Hgb M                        | 3.0%                | Not applicable           |
| Hgb B                        | Not applicable      | 60 (A to D units)        |

Table C.1-1 Flag Sensitivity Default Values

**Note:** Hgb M - defines the allowable difference between the three readings taken on the sample. Hob B - defines the allowable difference between the reference blank and sample blank.

\* MICRO and MACRO flags are activated in software version 1.03 and higher.

- For a definition of each flag, see Chapter 6 of the Operator's Guide.
- If you need to change a flag sensitivity value, use the procedure under Heading C.2.

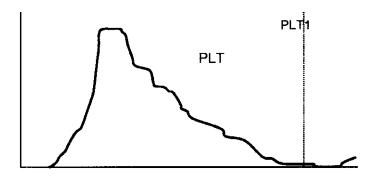
#### Thresholds

The values used to position the thresholds, which separate different cell populations and determine if a flag should be triggered, are selected to provide optimal population separation and flagging under normal operating conditions.

#### **Pit Threshold**

The PLT threshold (PLT 1) is the number of the last channel used to calculate the platelet count (Figure C.1-1). The factory setting for the Plt threshold is 197.

#### Figure C.1-1 PLT Threshold

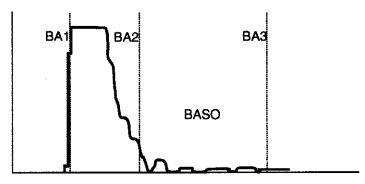


If you must change the platelet threshold, use the procedure under Heading C.3. However, be aware that changing this value from the default value will alter parameter values reported and whether or not those results are flagged.

#### WBC and BASO Thresholds

Differentiation between basophils and other leukocytes is obtained by means of the A<sup>C</sup>•T 5diff WBC Lyse-specific lytic action. Thresholds BA1, BA2, and BA3 allow the determination of the basophil population in relationship to the total number of white blood cells (Figure C.1-2).





In Figure C.1-2, all leukocytes are shown between the BA1 and BA3 thresholds. \*WBC absolute value is calculated between channel 0 and the BA1 threshold.

The percentage of basophils is calculated according to the number of particles from the BA2 threshold to the BA3 threshold. These thresholds are factory-set to the values shown in Table C.1-2.

| Table C.1-2 | WBC/BASO | Factory-Set | Threshold | Values |
|-------------|----------|-------------|-----------|--------|
|-------------|----------|-------------|-----------|--------|

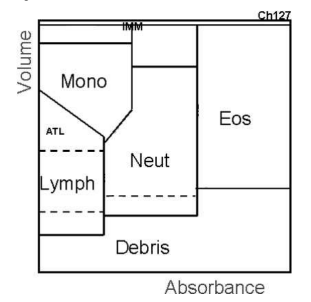
| Threshold | Purpose   | Channel |
|-----------|---|---------|
| 0         | *WBC # counting area  | 0       |
| BA1       | Separation threshold between the *WBC # counting area and the WBC | 35      |
| BA2       | Separation threshold between the WBC and basophils                | 110     |
| BA3       | End of the basophil counting area                                 | 240     |

If you must change the BA1, BA2, and BA3 thresholds, use the procedure under Heading C.3. However, be aware that changing these values from the default values will alter the parameter values reported and whether or not those results are flagged.

#### **DiffPlot Thresholds**

On the DiffPlot, both the X-axis and Y-axis are divided into 128 channels, numbered from 0 to 127 (Figure C.1-3).

#### Figure C.1-3 DiffPlot



Although invisible, there are 13 vertical grids on the DiffPlot's X-axis and 13 horizontal grids on the Y-axis. These grids form channels that can be located and given numbers. The origin or first location is Channel 0, at the bottom left corner. The fourth location of the DiffPlot is channel 30, and so forth. Threshold adjustment is expressed in channels.

Adjustment of the thresholds may be considered to:

- Improve the separation between different cell populations that may vary according to the anti-coagulant used or the instrument's internal adjustment.
- Modify the flag areas to improve detection sensitivity. You must also readjust the value of the respective flags.
- Modify one or several DiffPlot regions to precisely define a specific population for research purposes.

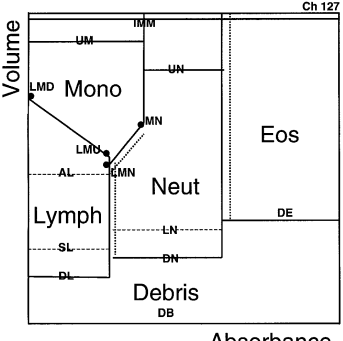
There are three categories of threshold adjustment limits:

- Volume thresholds on the Y-axis.
- Absorbance thresholds on the X-axis.
- FNL, FNE, FMN thresholds which indicate the width of the channel in the NL, NE, and MN alarm areas.

#### **DiffPlot - Volume Thresholds**

The volume thresholds on the Y-axis are identified in Figure C.1-4 and explained in Table C.1-3.





Absorbance

| Threshold | Purpose   | Default<br>Channel | Low Limit | High Limit  |
|-----------|---|--------------------|-----------|-------------|
| DL        | Separates debris and small lymphocytes.                                       | 22                 | 0         | SL          |
| DN        | Separates debris and lower neutrophils.                                       | 25                 | DL        | DE          |
| SL        | Separate lower lymphocytes and lymphocytes.                                   | 30                 | NL        | AL          |
| LN        | Separates neutrophils and small neutrophils.                                  | 35                 | DN        | LMN         |
| DE        | Separates debris and eosinophils.   | 48                 | DN        | Channel 127 |
| LMN       | Intersection point between the LY, MO, and NE thresholds.                     | 70                 | LN        | LMU         |
| AL        | Separates lymphocytes and atypical lymphocytes.                               | 68                 | SL        | LMU         |
| LMU       | Lower dot on the separation slope between atypical lymphocytes and monocytes. | 78                 | AL        | LMD         |
| LMD       | Upper dot on the separation slope between atypical lymphocytes and monocytes. | 90                 | LMU       | UM          |
| MN        | Upper dot on the separation slope between monocytes and neutrophils.          | 90                 | LMN       | UM          |
| UM        | Separates monocytes and upper monocytes.                                      | 118                | LMD       | Channel 127 |
| UN        | Separates neutrophils and upper neutrophils.                                  | 118                | MN        | Channel 127 |

Table C.1-3 DiffPlot - Volume Thresholds (Y-Axis)

#### **DiffPlot - Absorbance Thresholds**

The absorbance thresholds on the X-axis are identified in Figure C.1-5 and explained in Table C.1-4.

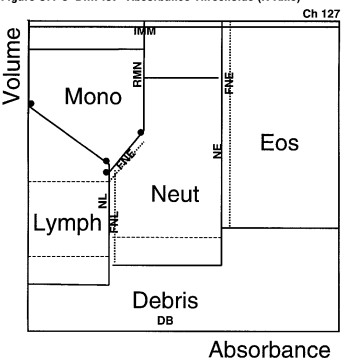


Figure C.1-5 DiffPlot - Absorbance Thresholds (X-Axis)

| Table C.1-4 | <b>DiffPlot - Absorbance</b> | Thresholds (X-Axis) |
|-------------|------------------------------|---------------------|
|-------------|------------------------------|---------------------|

| Thresholds | Purpose  | Default<br>Channel | Low Limit | High Limit  |
|------------|--|--------------------|-----------|-------------|
| NL         | Separates lymphocytes and neutrophils.           | 29                 | 0         | RMN         |
| RMN        | Separates upper monocytes and upper neutrophils. | 51                 | NL        | NE          |
| NE         | Separates neutrophils and eosinophils.           | 78                 | RMN       | Channel 127 |

#### NL, NE, and MN Alarms

FNL, FNE, FMN thresholds indicate the width in channel of the NL, NE, and MN alarm areas. The alarms are identified in Figure C.1-5 and explained in Table C.1-5.

| Threshold | Purpose                                 | Default Channel |
|-----------|---|-----------------|
| FNL       | Channel number for the NL alarm region. | 2               |
| FNE       | Channel number for the NE alarm region. | 2               |
| FMN       | Channel number for the MN alarm region. | 2               |

Table C.1-5 DiffPlot - FNL, FNE, and FMN Thresholds

#### **RBC** Histogram

RDI (Red Cell Distribution Index) threshold indicates the width in channel numbers of the RBC1 and RBC2 thresholds used in MICRO and MACRO calculation. The default value is 12.2.

### FLAG SENSITIVITY AND THRESHOLDS Overview

### C.2 SETTING FLAG SENSITIVITY

#### Purpose

**IMPORTANT** Risk of inaccurate flagging. Use extreme caution when doing this procedure, flag sensitivity settings are typically not changed from the factory default settings.

It is strongly recommended that **no changes** be made to the default values. Changing the flag sensitivity values from the default values will alter whether or not a flag will be associated with a result.

Use this procedure only if changing the flag sensitivity values is absolutely necessary. You may also use this procedure to re-enter the flag sensitivity default values listed in Table C.1-1.

#### **Tools/Supplies Needed**

Service password (If needed, see Service Password under Heading 4.1.)

#### Procedure

- 1. From the Main menu, select **SETUP → LAB. LIMITS → FLAGS SENSITIVITY**. The *SERVICE PASSWORD* prompt appears.
- 2. At the instrument keypad, press the numeric keys correlating to the Service password then press **ENTER**. The Flags Sensitivity screen appears (Figure C.2-1).

Figure C.2-1 Flags Sensitivity Screen

| FLAGS SENSITIVITY        |                  | 12 / 07 / 9          |          |
|--------------------------|------------------|----------------------|----------|
| DIFF REJECT              |                  | ENSIIIVIIY           | I        |
| DB % 100.0<br>SL % 100.0 | # 120 *V<br># 50 | VBC% 3.5             | # 999    |
| SL1 % 5.0<br>NL % 3.0    | # 45 M           | CRO 5.0<br>ACRO 20.0 |          |
| MN % 100.0               | # 120            |                      |          |
| UM % 1.1<br>LN % 2.5     | # 999 HO         | GBM% 3.0<br>GBB# 60  |          |
| UN % 1.1<br>NE % 1.1     | # 999<br># 60    |                      |          |
|                          |                  |                      | 7616127F |

- 3. Move the cursor to the value to be changed.
- 4. Edit the value using the numeric keypad.
- 5. Press **ENTER** to accept the new value and move to the next field.
- 6. Repeat steps 3 through 5 as needed to change additional values.
- 7. Select the next range and repeat steps 3 through 7 as required.
- 8. When all changes are completed, press **ESC** as many times as necessary to return to the Main Menu.

#### Verification

Run several normal and abnormal whole-blood specimens to verify the results are accurate with proper flagging, as applicable.

### **FLAG SENSITIVITY AND THRESHOLDS** SETTING FLAG SENSITIVITY

### C.3 SETTING THRESHOLDS

#### Purpose

**IMPORTANT** Risk of inaccurate results. Use extreme caution when doing this procedure, threshold settings are typically not changed from the factory default settings.

It is strongly recommended that **no changes** be made to the default values. By changing the thresholds, you are altering the populations. Therefore, changing any of the values that define threshold positions will ultimately alter the parameter values reported and whether or not a flag is associated with a result.

Use this procedure only if changing the threshold values is absolutely necessary. You may also use this procedure to re-enter the threshold default values listed in Appendix C:

- Table C.1-2, WBC/BASO Factory-Set Threshold Values
- Table C.1-3, DiffPlot Volume Thresholds (Y-Axis)
- Table C.1-4, DiffPlot Absorbance Thresholds (X-Axis)
- Table C.1-5, DiffPlot FNL, FNE, and FMN Thresholds

#### **Tools/Supplies Needed**

□ Service password (If needed, see Service Password under Heading 4.1.)

#### **Procedure**

- 1. From the Main menu, select **SETUP → LAB. LIMITS → THRESHOLDS**. The *SERVICE PASSWORD* prompt appears.
- 2. At the instrument keypad, press the numeric keys correlating to the Service password then press **ENTER**. The Thresholds screen appears (Figure C.3-1).

#### Figure C.3-1 Thresholds Screen

| THRESHO                   | THRESHOLDS 12 / 07 / 99   16:05 |   |
|---------------------------|---------------------------------|---|
|                           | THRE                            | SHOLDS 1                                    |
| BA1 35                    | BA2 110                         | BA3 240                                     |
| DL 22<br>UN 118<br>LMD 90 | DN 25<br>SL 30<br>LMN 70        | DE 48 LN 35<br>AL 68 LMU 68<br>MN 90 UM 118 |
| NL 29                     | NE 78                           | UMN 51                                      |
| FNL 2                     | FNE 2                           | FMN 2                                       |
| RDI 12.2                  |                                 |   |
|                           |                                 | 7616128B                                    |

- 3. Move the cursor to the value to be changed.
- 4. Edit the value using the numeric keypad.
- 5. Press **ENTER** to accept the new value and move to the next field.
- 6. Repeat steps 3 through 5 as needed to change additional values.
- 7. Select the next range and repeat steps 3 through 7 as required.
- 8. When all changes are completed, press **ESC** as many times as necessary to return to the Main Menu.

#### Verification

Run several normal and abnormal whole-blood specimens to verify the results are accurate with proper flagging, as applicable.



#### ABBREVIATIONS, ABBREVIATIONS-1

### CONTENTS

The following list is a composite of the abbreviations, acronyms and reference designators used in this manual. When the same abbreviation (or reference designator) is used for more than one word (or type of component), all meanings relevant to this manual are included, separated by semicolons.

## SYMBOLS

°C - degrees Celsius °F - degrees Fahrenheit > - greater than ≥ - greater than or equal to < - less than ≤ - less than or equal to µA - micro amperes µL - microliters µs - microseconds - - minus % - percent + - plus ± - plus or minus

### A

ac - alternating current ATL - atypical lymphocytes

## B

BA - basophil

## C

cm - centimeters CTN -CV - coefficient of variant; check valve

## D

dc - direct current DB - debris Dil - diluent DVM - digital voltmeter DFF - dual focused flow

### Ε

ESD - electrostatic discharge

EV -EOS - eosinophil

## F

F - fuse fL - femtoliters ft - feet

### G

g - grams g/dL - grams per deciliter

### H

Hct - hematocrit Hg - mercury Hgb -hemoglobin Hz - hertz

## I

i.d. - internal diameter in. - inches in. Hg - inches of mercury I/O - input/output IMM - immature cell

### J

J - receptacle connector

### K

K - kilos KΩ - kilohms KHz - kilohertz

### L

LCD - liquid crystal display LED - light emitting diode LN - lower neutrophil LYMPH -lymphocyte

### М

 $M\Omega$  - megohms M - motor mA - milli-amperes max - maximum MB - megabytes MACRO - macrocytes MCH - mean cell hemoglobin MCHC - mean cell hemoglobin concentration MCV - mean corpuscle volume MICRO - microcytes MN - monocyte/neutrophil) MONO - monocyte MHz - megahertz mb - millibars mL - milliliters mm - millimeters mN.m - milliNewton meter

### N

NE - neutrophil /eosinophil) NEUT - neutrophil NL - neutrophil /lymphocyte nm - nanometers

### 0

o.d. - outside diameter ozf.in - ounce force inch

### Ρ

P - plug connector
PC - printed circuit
PCB - printed circuit board
Plt - platelet count
PN - part number
ppm - pages per minute

### Q

QA - quality assurance QC - quality control

## R

R - resistor RAM - random access memory RBC - red blood cell count RDW - red cell distribution width RF - reference RH - relative humidity RN - resistor ROM - read only memory RPWV - red pulse-width value RS-232 - Electronic Industries Association standard governing interface between data processing and data communications equipment

## S

S - switch; solenoid SLO-BLO - slow blow SW - switch SL - small lymphocytes SL1 - small lymphocytes 1

## Т

Temp - temperature TP - test point

### U

UM - upper monocyte UN - upper neutrophil

### V

V - volts Vac - vacuum; volts alternating current Vdc - volts direct current

### W

W - watt

WBC - white blood cell count WM - wire marker WMCV - white mean corpuscle value WPWV - white pulse-width voltage

## X

X - jumper; plugged

x - times

### Numerics

5diff syringe, 4.29-1 5-part leukocyte differential, 2.1-1 +5V voltage, 4.20-4 +12V voltage, 4.20-4 -12V voltage, 4.20-4 +24V voltage, 4.20-4 35°C (95°F), regulated temperature of, 2.2-5

## A

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