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Section 1

INTRODUCTION

The LifeCare® PCA Plus and Plus II Series Infusers, herein referred to as the infuser, provide analgesic delivery to patients through the intravenous or epidural routes in a wide range of clinical settings.

1.1 SCOPE

This manual is organized into 11 sections as follows:

- Section 1 Introduction
- Section 2 Warranty
- Section 3 System Operating Manual
- Section 4 Theory of Operation
- Section 5 Maintenance and Service Tests
- Section 6 Troubleshooting
- Section 7 Replaceable Parts and Repairs
- Section 8 Specifications
- Section 9 Drawings
- Section 10 Index
- Technical Service Bulletins

If a problem in the infuser operation cannot be resolved using the information in this manual, contact Abbott Laboratories Technical Support Operations (see Section 6.1, Technical Assistance).

The system operating manual contains specific instructions for operating the infuser. Provision is made for inclusion of the system operating manual in Section 3 of this manual.

Note: Figures are rendered as graphic representations to approximate the actual product; therefore, figures may not exactly reflect the product. Display screens and touchswitch labels may vary slightly, depending on the configuration of the infuser in use.
1.2 CONVENTIONS

The conventions listed in *Table 1-1, Conventions*, are used throughout this manual.

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<td>Remove power supply PWA and CPU/display PWA as described in <em>Section 7.3.15, Power Supply PWA Replacement</em></td>
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<td>Touchswitch labels on the infuser are described in all caps and enclosed in brackets</td>
<td>[LOADING DOSE]</td>
</tr>
<tr>
<td><strong>ALL CAPS</strong></td>
<td>Screen displays (as appropriate)</td>
<td>ADMINISTER LOADING DOSE NOW?</td>
</tr>
<tr>
<td><strong>Bold</strong></td>
<td>Emphasis</td>
<td>CAUTION: Infuser damage may occur unless proper care is exercised during unpacking, inspection, and self test</td>
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Throughout this manual, warnings, cautions, and notes are used to emphasize important information as follows:

**WARNING**

A WARNING CONTAINS SPECIAL SAFETY EMPHASIS AND MUST BE OBSERVED AT ALL TIMES. FAILURE TO OBSERVE A WARNING IS POTENTIALLY LIFE THREATENING.

**CAUTION:** A CAUTION usually appears in front of a procedure or statement and contains information that could prevent irreversible equipment damage or failure.

**Note:** A note highlights information that helps explain a concept or procedure.

1.3 ACRONYMS AND ABBREVIATIONS

Acronyms and abbreviations used in this manual are as follows:

- **AC** Alternating current
- **ADC** Analog-to-digital converter
- **BRD** Bus read
- **BWR** Bus write
<table>
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<td>Centimeter</td>
</tr>
<tr>
<td>CPU</td>
<td>Central processor unit</td>
</tr>
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<td>Chip select</td>
</tr>
<tr>
<td>dB</td>
<td>Decibel</td>
</tr>
<tr>
<td>DC</td>
<td>Direct current</td>
</tr>
<tr>
<td>DISBATT</td>
<td>Disable battery</td>
</tr>
<tr>
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<td>Digital pressure meter</td>
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<tr>
<td>ECG</td>
<td>Electroencephalogram</td>
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<td>Electrocardiograph</td>
</tr>
<tr>
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<td>Hour</td>
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</tbody>
</table>
1.4

USER QUALIFICATION

The infusers are for use at the direction or under the supervision of licensed physicians or by licensed or certified healthcare professionals who are trained in the use of the infusers and the administration of parenteral or enteral fluids or drugs.

1.5

ARTIFACTS

Nonhazardous, low-level electrical potentials are commonly observed when fluids are administered using infusion devices. These potentials are well within accepted safety standards, but may create artifacts on voltage-sensing equipment such as ECG, EMG, and EEG machines. These artifacts vary at a rate that is associated with the infusion rate. If the monitoring machine is not operating correctly or has loose or defective connections to its sensing electrodes, these artifacts may be accentuated so as to simulate actual physiological signals. To determine if the abnormality in the monitoring equipment is caused by the infusion device instead of some other source in the environment, set the infusion device so that it is temporarily not delivering fluid. Disappearance of the abnormality indicates that it was probably caused by electronic noise generated by the infusion device. Proper setup and maintenance of the monitoring equipment should
eliminate the artifact. Refer to the appropriate monitoring system documentation for setup and maintenance instructions.

1.6

INSTRUMENT INSTALLATION PROCEDURE

CAUTION: Infuser damage may occur unless proper care is exercised during product unpacking, inspection, and self test. The battery may not be fully charged upon receipt of the infuser. Do not place the infuser in service if it fails the self test.

The infuser installation procedure consists of unpacking, inspection, and self test.

Note: Do not place the infuser in service if the battery is not fully charged. To make certain the battery is fully charged, connect the infuser to AC power for 24 hours.

1.6.1

UNPACKING

Inspect the infuser shipping container as detailed in Section 1.6.2, Inspection. Use care when unpacking the infuser. Retain the packing slip and save all packing material in the event it is necessary to return the infuser to the factory. Verify that the shipping container contains a copy of the system operating manual.

1.6.2

INSPECTION

Inspect the infuser shipping container for damage prior to opening. Should any damage be found, contact the delivering carrier immediately.

Inspect the infuser for evidence of damage. Do not use the infuser if it appears to be damaged. Should damage be found, contact Abbott Laboratories (see Section 6.1, Technical Assistance).

Inspect the infuser periodically for signs of defects such as worn accessories, broken connections, or damaged cable assemblies. Also inspect the infuser after repair or during cleaning. Replace any damaged or defective external parts.
1.6.3

**SELF TEST**

**CAUTION:** Do not place the infuser in service if the self test fails.

To conduct the infuser self test, proceed as follows:

1. Unlock and open the security door.
2. Connect AC power cord to a properly grounded AC outlet. Confirm that the AC power symbol on the front panel is illuminated.
3. Press and release the [ON] touchswitch. The infuser performs a self test verifying its functional integrity. If the infuser fails the self test, do not place it in service; contact Abbott Laboratories.
4. *PCA Plus II only:* Set the real time clock as described in Section 7.2.9, Setting the Real Time Clock (PCA Plus II).

**Note:** To assure the battery is fully charged, connect the infuser to AC power for 24 hours before placing it in service.
Section 2

WARRANTY

Subject to the terms and conditions herein, Abbott Laboratories, herein referred to as Abbott, warrants that (a) the product shall conform to Abbott's standard specifications and be free from defects in material and workmanship under normal use and service for a period of one year after purchase, and (b) the replaceable battery shall be free from defects in material and workmanship under normal use and service for a period of 90 days after purchase. Abbott makes no other warranties, express or implied, as to merchantability, fitness for a particular purpose, or any other matter.

Purchaser’s exclusive remedy shall be, at Abbott’s option, the repair or replacement of the product. In no event shall Abbott’s liability arising out of any cause whatsoever (whether such cause be based in contract, negligence, strict liability, other tort or otherwise) exceed the price of such product, and in no event shall Abbott be liable for incidental, consequential, or special damages or losses or for lost business, revenues, or profits. Warranty product returned to Abbott must be properly packaged and sent freight prepaid.

The foregoing warranty shall be void in the event the product has been misused, damaged, altered, or used other than in accordance with product manuals so as, in Abbott’s judgment, to affect its stability or reliability, or in the event the serial or lot number has been altered, effaced, or removed.

The foregoing warranty shall also be void in the event any person, including the Purchaser, performs or attempts to perform any major repair or other service on the product without having been trained by an authorized representative of Abbott and using Abbott documentation and approved spare parts. For purposes of the preceding sentence, "major repair or other service" means any repair or service other than the replacement of accessory items such as batteries, flow detectors, detachable AC power cords, and patient pendants.

In providing any parts for repair or service of the product, Abbott shall have no responsibility or liability for the actions or inactions of the person performing such repair or service, regardless of whether such person has been trained to perform such repair or service. It is understood and acknowledged that any person other than an Abbott representative performing repair or service is not an authorized agent of Abbott.
This page intentionally left blank.
A copy of the system operating manual is included with every infuser. Insert a copy here for convenient reference. If a copy of the system operating manual is not available, contact Abbott Laboratories Technical Support Operations (see Section 6.1, Technical Assistance).
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THEORY OF OPERATION

This section describes the principles of operation for the LifeCare PCA Plus and Plus II Series Infusers. Related drawings are provided in Section 9, Drawings.

4.1 GENERAL FUNCTIONS

This section describes the general functions of the infuser (see Figure 9-2, Block Diagram, PCA II).

4.1.1 STARTING OPERATION

The infuser is powered on by inserting or removing a vial, or by pressing the [ON] touchswitch. Once powered on, the infuser begins a warm-start or a cold-start sequence, depending on the time elapsed since the [OFF/RECHG] touchswitch was last pressed.

4.1.1.1 WARM-START SEQUENCE

The infuser initiates a warm-start sequence if it is powered on within 60 minutes of previous operation. In the warm-start sequence, all previous therapy parameters and dose history information are retained in memory. Before the infuser can operate, previous settings must be confirmed or cleared. If previous settings are cleared, the user must program the infuser with new therapy parameters.

4.1.1.2 COLD-START SEQUENCE

The infuser initiates a cold-start sequence if it is powered on after 60 minutes since previous operation. The infuser also initiates a cold-start sequence if the [OFF/RECHG] touchswitch is pressed for approximately four seconds to clear all memory.

After 60 minutes, most major electronic circuits turn off and all therapy settings and dose history data are lost. During a cold-start sequence, a self test is performed to verify the functional integrity of the infuser. After successful completion of the self test, the infuser must be reprogrammed with new therapy parameters.
4.1.2 SETUP

During setup, the liquid crystal display (LCD) screen displays prompts for each step of programming the infuser. The infuser must be programmed before it will operate.

4.1.3 DELIVERY MECHANISM

The delivery mechanism is actuated by a four-phase stepper motor that controls delivery. The delivery mechanism is constantly monitored by the infuser electronics.

4.1.4 PATIENT CONTROLLED ANALGESIA (PCA)

When the infuser is operating in either the PCA or the PCA + CONTINUOUS modes, the patient can initiate a request for analgesia by pressing the patient pendant pushbutton. The patient request is answered based on the status of the lockout interval and the four-hour dose limit. A dose is delivered only if the lockout interval has expired and the four-hour dose limit has not been reached. Timing of the lockout interval and the four-hour dose limit is programmed by the medical professional, and is maintained by the central processor unit (CPU). The four-hour dose limit is the maximum PCA dose deliverable by the infuser in any rolling four-hour period.

Note: The PCA Plus II will not discontinue an individual dose upon reaching the four-hour dose limit; the dose continues until completion.

4.1.5 STATUS MESSAGES

Infuser status is displayed on the LCD screen during normal operation. Status messages are listed in Table 4-1, Status Messages and Definitions.

<table>
<thead>
<tr>
<th>Table 4-1. Status Messages and Definitions</th>
</tr>
</thead>
<tbody>
<tr>
<td><strong>Message</strong></td>
</tr>
<tr>
<td>--------------</td>
</tr>
<tr>
<td>DOOR LOCKED</td>
</tr>
<tr>
<td>PATIENT LOCKOUT</td>
</tr>
<tr>
<td>4 HR LIM REACHED</td>
</tr>
</tbody>
</table>
### Table 4-1. Status Messages and Definitions

<table>
<thead>
<tr>
<th>Message</th>
<th>Definition</th>
</tr>
</thead>
<tbody>
<tr>
<td>* * *</td>
<td>Indicates that patient can initiate a dose in PCA or PCA + CONTINUOUS mode</td>
</tr>
<tr>
<td>PCA</td>
<td>Displays when infuser is in PCA mode</td>
</tr>
<tr>
<td>CONTINUOUS</td>
<td>Displays when infuser is in CONTINUOUS mode</td>
</tr>
<tr>
<td>PCA + CONTINUOUS</td>
<td>Displays when infuser is delivering a PCA dose in addition to a continuous</td>
</tr>
<tr>
<td></td>
<td>delivery in the PCA + CONTINUOUS mode</td>
</tr>
</tbody>
</table>

---

### 4.2 ELECTRONICS OVERVIEW

For interconnect diagrams of the LifeCare PCA Plus and Plus II Series Infuser, see Figure 9-3, Interconnect Schematic, PCA II.

**Note:** A vinculum (overscore) above a signal designation (e.g. BWR) denotes an active low signal.

The infuser electronics system contains the CPU/display printed wiring assembly (PWA) and the power supply PWA. The microprocessor on the CPU/display PWA communicates with components on both PWAs through an 8-bit data bus (D7-D0) and a 16-bit address bus (A15-A0).

Low-level signals from the front panel touchswitches and sensor switches are input directly by the CPU/display PWA where they are placed in buffers on the microprocessor data bus. Data buffers on the CPU/display PWA also receive input from the power supply PWA.

Primary AC power is routed through line fuses F1 and F2 to the primary of the power transformer T1. Low level alternating current (AC) from the secondary of T1 is applied to a rectifier, CR14, on the power supply PWA. The direct current (DC) output of CR14 connects to the battery input and battery charger circuitry on the power supply PWA. The power supply PWA provides +5 VDC logic power, drivers for the drive motor, power to the audible alarm, access to test points throughout the system through a test port, and a printer/Dataway port.

---

### 4.2.1 CPU/DISPLAY PWA

For a schematic of the CPU/display PWA, see Figure 9-4, CPU, PWA Schematic, LC 4100/4200.

The CPU/Display PWA contains the following:

- 8032 single-chip microprocessor containing read-only memory (ROM), random-access memory (RAM), an 8-bit input/output (I/O) port, two timer circuits, 64K addressing capability, and an 8-bit data bus
32K erasable/programmable read-only memory (EPROM) for firmware program storage, including all CPU firmware

8K static RAM mounted in a socket that contains a real-time clock chip that can be read from and written to by the 8032 microprocessor. A battery in the socket supplies power to the real-time clock when logic power is turned off

Watchdog circuit that sounds an alarm if the 8032 microprocessor does not execute a code as designed or does not start up properly

DC-to-AC converter that provides power for the electroluminescent panel on the LCD screen

LCD containing its own controlling circuitry on a PWA connected to the CPU/display PWA

Light emitting diode (LED) displays: five seven-segment LEDs, one single LED, and two multi-LEDs, along with driving, multiplexing, and monitoring components

Eight-channel analog-to-digital converter that monitors the +5 VDC power supply, the +2.5 VDC precision voltage reference (used to regulate the +5 VDC supply and the battery monitoring circuits), and the current consumption of the LEDs

4.2.1.1
CPU/DISPLAY PWA SYSTEM INTERFACES

The CPU/display PWA interfaces with the front panel touchswitches, the LCD screen, and the power supply (see Figure 9-3, Interconnect Schematic, PCA II).

The 12 front panel touchswitches interface with the CPU/display PWA via P/J13. The YES/ENTER, REV/CHG, ONSW, and OFFSW touchswitch signals are routed off the CPU/display PWA via P/J12 to the power supply connector P/J7. The YES/ENTER and REV/CHG signals input the data buffer U8 on the power supply PWA. A similar data buffer, U4, on the CPU/display PWA inputs the remaining eight touchswitch signals. U8 and U4 place data on the system data bus (D0-D7). When one of the touchswitches is pressed, a ground is applied to the corresponding touchswitch signal line, which provides an active low signal at the appropriate data buffer input pin. The ONSW signal is used on the power supply PWA as an input to the ON/OFF latch consisting of U14B and U14C. The OFFSW line is routed to the test port via connector P/J9, pin 8.

The LCD screen interfaces with the CPU/display PWA via P/J15 and P/J14. AC power from the DC-to-AC converter U17 is supplied to the LCD screen via P/J15. The LCD screen also receives addresses A0 and A1 and data bits D0 through D7 via P/J14 for display control.

The primary interface between the CPU/display PWA and the power supply PWA interface is through the power supply PWA connector P/J12 on the CPU/display PWA which connects via a ribbon cable with connector P/J7 on the power supply PWA. The CPU/display PWA interface consists of address lines A1 and A2, data lines D0 through D7, +5 VDC power, +2.5 VDC reference voltage, and various signals.
4.2.1.2 MICROPROCESSOR CIRCUITRY

The microprocessor circuitry consists of the following:

- 8032 single-chip CPU, U11
- 10-MHz crystal oscillator, Y1
- Bus transceiver, U12
- Transparent latch, U13
- 8K x 8 RAM chip, U21
- 32K x 8 EPROM chip, U22

U11, the 8032 single-chip CPU, controls all functions of the infuser. U11 has internal RAM for the program stack and for storage of temporary variables. U11 uses a multiplexed 16-bit address bus, an 8-bit data bus, and a built-in 8-bit I/O port.

The microprocessor is clocked by Y1, a parallel-resonant 10-MHz crystal oscillator. Test points E1-E2 and E3-E4 are used to disconnect Y1 during depot level repair in order to single-step U11 for control.

All operating parameters for the infuser are stored in U21, the nonvolatile 8K x 8 RAM chip, which decodes the 8032 address bits (A0 through A14) to read from or write to the system data bus (data bits D0 through D7). Read and write operations are controlled by the BRD and BWR signals derived from the CPU RD and WR signals.

U21 is mounted on a socket that contains a lithium battery, a voltage monitor chip, and a real-time clock chip. The lithium battery is used to retain the data stored in RAM during short periods of infuser power loss. The voltage monitor chip monitors the power supplied to the socket and connects the lithium battery to the real-time clock chip if the logic power supply shuts down.

U22 is a 32K x 8 EPROM that stores the infuser software program. U22 places firmware instructions on the system data bus according to the status of address lines A0 through A15 when enabled by the PSEN signal from the CPU.

The octal tri-state bus receiver, U12, and the octal D-type transparent latch, U13, provide an interface between the CPU and the system data and address busses. U12 and U13 connect to the CPU lines AD0 through AD7. U12 uses AD0 through AD7 as a data interface between the CPU and the system data bus (D0 through D7), transmitting data to and receiving data from the various components throughout the system that connect to the data bus. U13 latches in these lines and uses them with CPU lines AD8 through AD15 for the system address bus, consisting of addresses A0 through A15.

4.2.1.3 CHIP SELECT CIRCUITRY

U6 is a 3-to-8 multiplexer that decodes address bits A6 through A4 when gated by A7 and A15 to provide eight I/O chip select signals, CS0 through CS7. These chip select signals, the bus read command (BRD), and the bus write command (BWR) are used as system control signals (see Table 4-2, CPU/Display PWA Control Signals).
### Table 4-2. CPU/Display PWA Control Signals

<table>
<thead>
<tr>
<th>Signal</th>
<th>Function</th>
</tr>
</thead>
<tbody>
<tr>
<td>CS0</td>
<td>NANDed with BWR to gates U2 and U14</td>
</tr>
<tr>
<td>CS1</td>
<td>NANDed with BRD to enable output of U19</td>
</tr>
<tr>
<td>CS2</td>
<td>NANDed with BRD to enable output of U19</td>
</tr>
<tr>
<td>CS3</td>
<td>Routed to power supply PWA; used as chip select input (CS) to PPI U7</td>
</tr>
<tr>
<td>CS4</td>
<td>ANDed with BRD to gate front panel switch data buffer U4</td>
</tr>
<tr>
<td>CS5</td>
<td>CPU bus read command (a function of the CPU RD command)</td>
</tr>
<tr>
<td>BRD</td>
<td>CPU bus read command (a function of the CPU RD command)</td>
</tr>
<tr>
<td>BWR</td>
<td>CPU bus read command (a function of the CPU RD command)</td>
</tr>
</tbody>
</table>

#### 4.2.1.4 ANALOG-TO-DIGITAL CONVERTER (ADC)

The ADC, U19, inputs the enable signals to the LEDs. The ADC provides data outputs that represent LED display status to the microprocessor data bus by decoding addresses A0, A1, and A2. The ADC provides a digital representation of eight analog inputs (IN0 through IN7) to U11. IN0 monitors the current consumption of U3 and the LED display buffer through differential amplifier U18A. Analog inputs IN1 through IN6 monitor the outputs of U3 (cathodes of the LED displays) to check for shorted or open LEDs. IN7 monitors the +2.5 VDC precision voltage reference to compare it against the supply voltage applied to the ADC.

#### 4.2.1.5 LCD DISPLAY INTERFACE

The LCD subassembly connects to the system data bus, address bits A1 through A0, and an I/O chip select. The LCD background lighting is provided by an electroluminescent panel powered by a DC-to-AC converter at U17. The converter converts the +5 VDC to approximately 90 VAC. The background lighting is turned off by disabling transistor Q2 two minutes after a touchswitch is pressed or 30 seconds after the door is locked. The viewing angle is set to provide maximum contrast by adjusting potentiometer R19. Potentiometer R19 is factory adjusted to give maximum contrast at a viewing angle perpendicular to the LCD screen. U10, a DC-to-DC converter, provides -5 VDC to allow a full 10 volt adjustment range of R19.
4.2.1.6 LED DISPLAY CIRCUIT

The LEDs are driven on a time-multiplexed basis. The pattern of segments to be illuminated is latched into U14 by U11. During the same CPU cycle, the seven-segment LED that displays the pattern is enabled by address bits written to the latch at U2, which is buffered by U3.

The anodes of the single LED that provides the decimal point and the LED array that illuminates the battery symbol are illuminated by the DP signal from the U14 latch. Information represented by the DP signal is displayed by the decimal point when it is enabled with the tenths display and by the battery LED when it is enabled with the units display.

4.2.1.7 WATCHDOG CIRCUIT

The watchdog circuit consists of the dual one shot at U15 and the two NAND gates from U16 that form a set/reset flip flop. The one shot is periodically triggered by U11 as part of its firmware program. Before the one shot times out and causes the flip flop to set the MALF line, U11 must activate RWAT, which resets the flip flop and prevents it from activating MALF. The watchdog circuit assures that the firmware contains the correct code and that U11 is executing code. The watchdog circuit also assures that U11 is the correct length and that the real-time clock, which sets U11 to activate RWAT, is keeping correct time and can be accessed by U11.

4.2.2 POWER SUPPLY PWA

For schematics of the power supply PWA, see Figure 9-5, Power Supply PWA Schematic, PCA II.

The power supply PWA contains the following:

- Power supply circuitry that converts the low-voltage (11 to 17 VAC) from the line transformer to regulated DC
- Battery charging and monitoring circuitry that keeps the battery at full charge and provides the LOW BATT alarm
- 8255 programmable peripheral interface (PPI), U7, that contains three 8-bit I/O ports to allow U11 to control the motor, read data from and send data to the printer port, and read the state of the switches
- Set/reset flip flop (ON/OFF latch), U14B/U14C, that monitors the state of the vial and injector switches and the [ON] touchswitch to control power to the logic circuitry, and is powered by the battery when AC power is off
- Motor drivers U17A, U17B, U17C, and U17D that step the motor through its phases, and motor monitoring circuitry U15A, U15B, U15C, U15D that detects driver and motor malfunctions
4.2.2.1
POWER SUPPLY PWA SYSTEM INTERFACES

The power supply PWA receives input from either the isolation transformer T1 or the battery pack via P/J3. The power supply PWA interfaces with the CPU/display PWA through a ribbon cable connected at P/J7, which consists of the microprocessor data bus lines, address lines A0 and A1, +5 VDC power, and various signal lines.

The power supply PWA receives the VIAWSW signal from P/J6 and the other microswitch signals (PRESSURES, SYRINGELO, SYRINGE EMPTYSW, and INJECTORSW) from P/J4. The motor drive signals and the shaft sensor feedback signals are routed to the power supply PWA via P/J5.

The audible alarm signal, audible level switch inputs, door switch signal, and patient pendant input interface with the power supply PWA through a connector, P/J8. P/J8 also routes lines to the printer/Dataway connector J10.

Test port J11 interfaces with the power supply PWA via P/J9 to provide external access to test points throughout the system.

4.2.2.2
UNREGULATED DC VOLTAGE

The output of AC power transformer T1 is applied to CR14 via connector P/J3. CR14 is a full-wave rectifier; capacitor C16 filters the resulting DC voltage. When AC power is on, operational amplifier U12A, transistors Q4 and Q2, and voltage reference U18 charge the battery. Transistor Q1 gates the unregulated voltage to U9 and U10, which are +5 VDC regulators for the logic power and motor power.

Operational amplifier U3 monitors the unregulated voltage and provides DISBATT and LOBAT signals to U11. The DISBATT and LOBAT signals are derived from two voltage thresholds produced from a precision resistor network.

4.2.2.3
ON/OFF LATCH

Unregulated DC voltage is present in the power supply when either AC or battery power is available. A supply of +5 VDC is available from the unregulated DC voltage by precision voltage references U5 and U6. The voltage powers U13 and U14, which make up the ON/OFF latch. The ON/OFF latch is a set/reset flip flop set by the injector or vial switches being closed or by the [ON] touchswitch being pressed. The output of the ON/OFF latch drives the gate of Q5. Q5 drives the base of Q1, which connects the +5 VDC logic and motor regulators with the unregulated DC voltage. Gates from U13 and U14 prevent the [OFF/RECHG] touchswitch from resetting the ON/OFF latch when a malfunction is being reported via the MALF line.

4.2.2.4
PRINTER DATA LATCH

The printer data latch latches data from U11 to send out of the printer port as printer data.
4.2.2.5

**MOTOR DRIVER**

The gates of U17 drive transistors Q9 through Q6 from the 8255 port A outputs and are disabled by MALF, preventing motor movement during a malfunction. The gates of U15 generate the MOTORBAD signal if the inputs to U17 are not of opposite polarity to the drive signals to the motor.

4.2.2.6

**8255 PROGRAMMABLE PERIPHERAL INTERFACE**

The 8255 programmable peripheral interface chip, U7, is an I/O peripheral that is written to and read from by U11 via the microprocessor data bus when addresses A0 and A1 are properly decoded. U7 contains three 8-bit I/O ports: port A, port B, and port C.

Port A (PA0 through PA7) is used for motor control, the strobe for the printer port, and the signal that enables the LCD backlighting. Port B is used for the printer data input and parity bit I/O. Port C is a general purpose input port that monitors most critical switches and indicators.

4.2.3

**BATTERY BOOST CHARGER PWA**

For a schematic of the battery boost charger PWA, see *Figure 9-6, Current Boost Charger Schematic, MDL 4/PCA*.

The battery boost charger PWA is installed in infusers with the following final assembly numbers and service revision letters:

- 850-04250-005 (L and above)
- 850-04250-008 (C and above)

**Note:** The final assembly number and service revision letter are located on the lower right side on the back of the infuser.

The battery boost charger PWA contains the following circuitry:

- Differential amplifier, U3
- 20 mA shut-off circuitry, Q2
- Window comparator with hysteresis, U1
- 60 minute battery charger timer, U2
- 200 mA constant current source (transistors Q7, Q8, Q9) and associated logic (transistors Q3, Q4, Q5, Q6)
- AC detector, transistor Q1

The battery boost charger PWA functions during infuser AC power and battery power operation, as described in the following sections.
4.2.3.1 AC OPERATION

Voltage detector U1, with associated capacitors and resistors, functions as a window comparator with hysteresis. When the battery is initially connected, the output of U1 is at logic high. After the battery voltage reaches 10 VDC, the output of U1 goes low, disabling transistors Q3 and Q4 and triggering the timer, U2.

The clock frequency of U2 is determined by C5 and R9; time-out is pending the biasing of the U2 program inputs A, B, C, and D. Upon time-out, the DECODE signal on output pin 13 of U2 goes high; this signal disables Q5, removes the supply to Q9, and causes battery charging to stop. The DECODE signal also locks in U2, preventing operation by placing a logic high on the SET input, pin 1, of U2.

The AC/DC detector consists of transistor Q1 and associated resistors. When the infuser operates on AC power, the collector of Q1 is at logic low, enabling timer U2 and transistor Q6. During DC operation, the +VRDC signal is at ground level and keeps Q1 off.

Current is limited to 200 mA by the quotient of the VBE of transistor Q8 divided by R17. Transistor Q7 acts as a power switch that is enabled and disabled by transistor Q9.

The IC, U3-B, acts as a differential amplifier with a gain of 20 decibels (dB). Current is sensed across resistor R62 in the battery charging circuitry of the power supply PWA. When current drops below approximately 20 mA, the output of U3-B, pin 1 (which is also the input to comparator U3-A, pin 6), is at 100 mV or below. Pin 3 of U3-A is referenced at 500 mV from the battery charger circuitry on the power supply PWA. Also, when current drops below 20 mA, U3-A switches to a logic high and turns on transistor Q2, disabling the charging circuit (Q1 and Q19) on the power supply PWA. Resistor R26 and capacitor C8 act as a noise filter to Q8.

4.2.3.2 DC OPERATION

When the infuser operates on battery power and the battery pack drains to approximately 8 VDC, pin 4 of the voltage detector U1 switches to logic high; this resets the timer U2, enabling transistors Q3, Q4, and Q5. Transistor Q6 remains disabled because Q1 is disabled.

4.3 MECHANICS OVERVIEW

Mechanical elements of the infuser consist of the slide assembly, the syringe low (-008 and lower) and syringe empty alarm system, the occlusion pressure alarm system, and the vial and injector sensor alarm system.
4.3.1

SLIDE ASSEMBLY

The main components of the slide assembly are shown in Figure 4-1, Infuser Slide Assembly (-008 and Lower), and Figure 4-2, Infuser Slide Assembly (-010 and Higher). The following sections describe the operation of the slide assembly.
4.3.1.1 MOTOR ASSEMBLY

The motor assembly consists of a four-phase stepper motor housed in the motor case (see Figure 4-1, Infuser Slide Assembly, (-008 and Lower)). The motor case attaches to the gearbox. The gearbox consists of reduction gears that provide a 50-to-1 reduction in the rotation of the motor shaft. The reduction gears drive the lead screw. Fifty revolutions of the motor produce one revolution of the lead screw.

The stepper motor receives its commands (MOTO1, MOTO2, MOTO3, and MOTO4) from motor drivers on the power supply PWA. The MOTO1, MOTO2, MOTO3, and MOTO4 commands are issued by the PPI U7 under control of the microprocessor on the CPU/display PWA.
Data buffer U4 on the CPU/display PWA inputs signals from the touchswitches on the infuser front panel. The touchswitch signals represent delivery data placed on the microprocessor data bus by U4 (D0 through D7) when U4 is gated by the ANDed product of the bus read command (BRD) and chip select signal CS6. Digitized delivery data is interpreted by the microprocessor according to the program in EPROM. The data provides the stepper motor commands for the delivery data set up by the touchswitches.

A reduced-diameter extension of the gear box shaft protrudes through the bottom of the gearbox; this extension is flattened to allow an opto switch mounted on a bracket on the motor case to sense rotation of the lead screw. The opto switch senses the flattened area of the extension. For each rotation of the lead screw, the opto switch generates a signal, SHAFT, which is routed to data buffer U8 on the power supply PWA.

The SHAFT signal is read by the microprocessor on the CPU/display PWA as data bit D6 when U8 is gated by the SWSTSEL signal from the CPU/display PWA. The SHAFT signal provides a feedback loop to allow the microprocessor to control the rotation of the lead screw for the delivery data set up by the touchswitches.

### 4.3.1.2 SLIDE MECHANISM

The slide block connects to the lead screw by a half nut. The surface of the half nut that contacts the lead screw is machined so that it engages the threads on the lead screw. When the lead screw is rotated and the half nut is engaged, the slide block moves downward towards the motor assembly. Squeezing the arm above the cradle with no syringe mounted in the cradle causes the half nut to disengage from the lead screw. The slide mechanism can then be moved up and down on the two guide shafts to the full extent of cradle travel.

### 4.3.1.3 CRADLE ASSEMBLY

When an inverted vial is inserted in the cradle assembly, the injector flange is held stationary in the injector flange clamp. When the slide block is driven downward by the rotation of the lead screw, the vial is forced downward and the stationary injector plunger moves further into the vial. Fluid in the vial is forced into the administration set in an amount proportional to the downward movement of the cradle and slide mechanism.

### 4.3.2 SYRINGE LOW AND SYRINGE EMPTY ALARM SYSTEM

**Note:** Software version 3.1 and higher deactivates the syringe low switch.

A syringe status microswitch mounted to the slide assembly provides a signal to the power supply supply PWA when activated by the downward movement of the cradle assembly (see Figure 4-3, Syringe Low and Syringe Empty Alarm System (-008 and Lower) and Figure 4-4, Syringe Empty Alarm System (-010 and Higher)). When activated, the syringe empty switch sends the SYRINGE EMPTYSW signal to data buffer U8 on the power supply PWA. Data buffer U8 places switch status data on the microprocessor data bus as data bits D1 and D2 for processing by the CPU/Display PWA microprocessor when the buffer is gated by SWSTSEL.
On infusers with final assembly number -008 and lower, two bracket arms attach to the slide block; two syringe status microswitches mount on a bracket attached to the base of the slide assembly. The two bracket arms are oriented to the switches so that downward movement of the slide block causes the bracket arms to press against actuator arms on the switches. The inner microswitch (the syringe low switch) is closed first, by the longer, innermost arm, sending the SYRINGE LOSW signal to the power supply PWA. Further downward movement causes the shorter of the two bracket arms (the outer arm) to press the actuator arm on the outer switch (the syringe empty switch), which sends the SYRINGE EMPTYSW signal to the power supply PWA.

**Note:** Later configurations of -008 and lower have one bracket arm containing both actuators, allowing both the syringe low and syringe empty alarms to be adjusted with a single adjustment on the one bracket arm.

The SYRINGE LOSW signal occurs when the distance between the bottom of the syringe vial stop and the bottom of the syringe injector flange groove is 4.628 ± 0.062 inches (11.57 ± 0.155 centimeters (cm)), which indicates approximately 5 mL of fluid is left in the vial. The SYRINGE LOSW signal generates the syringe low alarm and the LOW SYRINGE message on the LCD.

The SYRINGE EMPTYSW signal occurs when the distance between the bottom of the syringe vial stop and the bottom of the syringe injector groove is 4.328 ± 0.062 inches (10.82 ± 0.155 cm), which indicates approximately 1 mL remaining in the vial. The SYRINGE EMPTYSW signal generates the syringe empty alarm and the EMPTY SYRINGE message on the LCD.

On infusers with final assembly number -010 and higher, the slide block moves down on the switch, causing the switch arm to activate the syringe empty switch.

**Note:** Infusers with final assembly number -010 and higher do not have a syringe low switch.

---

**Figure 4-3. Syringe Low and Syringe Empty Alarm System (-008 and Lower)**
4.3.3 OCCLUSION PRESSURE ALARM SYSTEM

The OCCLUSION alarm occurs when a backpressure of $15 \pm 5$ psig ($103.5 \pm 34.5$ kPa) is exceeded. Backpressure is sensed by the occlusion pressure microswitch on the top plate of the slide assembly (see Figure 4-5, Occlusion Pressure Alarm System (-008 and Lower) and Figure 4-6, Occlusion Pressure Alarm System (-010 and Higher)).

During a normal delivery cycle, rotation of the lead screw moves the slide block downward, which moves the cradle assembly downward and forces fluid out of the syringe. If there is an occlusion in the administration set, backpressure builds up in the delivery system and downward motion of the slide block is restricted.

A longitudinal slack in the lead screw exists so that when the downward movement of the slide block is restricted, rotation of the lead screw forces the slide block upwards against a compression spring between the top of the lead screw and the top plate. When a given amount of pressure is applied against the compression spring, the upper end of the lead screw presses against the switch actuator and causes it to pivot upward, thereby closing the occlusion alarm switch. On infusers with final assembly number -010 and higher, the switch opens.

When the occlusion alarm switch closes, the PRESSURESW signal goes active low. The PRESSURESW signal is applied to the input of data buffer U8 on the power supply supply PWA. When U8 is gated by the SWSTSEL signal, the microprocessor on the CPU/Display PWA reads data bit D0 for occlusion pressure status and generates an OCCLUSION alarm. In setup mode, the OCCLUSION alarm is cleared by releasing the cradle assembly. In patient mode, the OCCLUSION alarm is cleared by opening the front door and releasing the cradle assembly. The [SILENCE] touchswitch may be pressed to mute the occlusion audible alarm for one minute.
Occlusion pressure is adjusted as described in Section 7.2.7, Occlusion Alarm Switch Test and Adjustment.

Figure 4-5. Occlusion Pressure Alarm System (-008 and Lower)
4.3.4
VIAL AND INJECTOR SENSOR ALARM SYSTEM

The vial sensor microswitch between the cradle assembly and slide block senses the presence or absence of a vial in the cradle assembly. The injector sensor switch behind the injector flange clamp senses whether or not the vial injector flange is properly seated in the injector flange clamp when a vial is clamped in the cradle assembly. When open, the vial sensor switch generates the CHECK VIAL alarm.

4.3.4.1
CHECK VIAL ALARM

When a vial is properly seated in the cradle assembly, the vial sensor rod is forced towards the rear of the infuser and presses against the vial sensor microswitch actuator arm, causing the switch to close. If the vial is improperly seated (i.e., the vial sensor rod is not forced rearward and the switch remains open) and the injector flange is properly seated, an attempt to start the infuser causes logic circuitry on the power supply supply PWA to sense the VIALSW signal and set the ON/OFF latch (U14B, U14C). When the ON/OFF latch is set, the infuser is prevented from starting, the audible alarm sounds, and the CHECK VIAL message is displayed on the LCD. The [SILENCE] touchswitch has no effect on this alarm; the alarm is cleared by properly seating the vial in the cradle assembly.
4.3.4.2
CHECK INJECTOR ALARM

When a vial is mounted in the cradle assembly and the injector flange is properly seated in the injector flange clamp, the vial sensor rod is forced towards the rear of the infuser and presses against the injector sensor microswitch, which causes the switch to close. If a vial is installed in the cradle, but the injector flange is improperly seated (i.e., the vial sensor rod is not forced rearward and the switch remains open), and an attempt is made to start the infuser, logic circuitry on the power supply supply PWA senses the status of the INJECTORSW signal and sets the ON/OFF latch (U14B, U14C). When the ON/OFF latch is set, the infuser is prevented from starting, the audible alarm sounds, and the CHECK INJECTOR message is displayed on the LCD. The [SILENCE] touchswitch has no effect on this alarm; the alarm is cleared by properly seating the injector flange in the injector flange clamp.

4.3.4.3
CHECK SYRINGE ALARM

When no syringe is installed in the infuser, or when a syringe is installed, but both the vial and the injector flange are improperly seated, the vial sensor switch and the injector sensor microswitch remain open. If an attempt is made to start the infuser under these conditions, logic circuitry on the power supply supply PWA senses the status of the INJECTORSW signal and sets the ON/OFF latch (U14B, U14C). When the ON/OFF latch is set, the infuser is prevented from starting, the audible alarm sounds, and the CHECK SYRINGE message is displayed on the LCD. The [SILENCE] touchswitch has no effect on this alarm; the alarm is cleared by installing a syringe so that the vial is properly seated in the cradle assembly and the injector flange is properly seated in the injector flange clamp.

4.4
BATTERY OPERATION OVERVIEW

The infuser is intended to operate on battery power on an exception basis only, such as emergency backup or temporary portable operation. Examples of emergency backup include AC power failure or inadvertent disconnection of the AC power cord. Temporary portable operation includes patient transfer from one location to another.

The infuser should be connected to AC power whenever possible to allow the battery to remain fully charged. The battery indicator illuminates when the infuser is operating on battery power.

Factors that most commonly affect battery life are the depth and frequency of discharge and the length of the recharge period. As a general rule, the more often the battery is discharged and recharged, the sooner it will need replacement. The primary cause of damage is leaving the battery in a less than fully charged state for any period of time. Battery damage can occur in a matter of hours and cause a permanent loss of battery capacity. The amount of lost capacity depends on the degree of discharge, the storage temperature, and the length of time the battery was stored in a discharged state.

Note: A permanently damaged battery cannot be recharged to full capacity.
When the battery discharges below an acceptable level while the infuser is operating, an alarm sounds and the LOW BATTERY message displays. Although it is not recommended to continue operating the infuser on battery power at this point, the battery will continue providing power until discharged; at this point, the infuser enters the battery discharged mode and operation ceases.

**CAUTION: As soon as the LOW BATTERY alarm occurs, connect the infuser to AC power.**

Recharging occurs any time the infuser is connected to AC power. It is recommended that the infuser be connected to AC power whenever practicable to maximize available battery charge during patient transport or ambulation. The power switch does not have to be on for the battery to recharge. Recharging while the infuser is operating is rate dependent.
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Section 5

MAINTENANCE AND SERVICE TESTS

This section contains preventive maintenance information, a performance verification test (PVT), and battery maintenance information for the PCA Plus and Plus II Series Infuser.

5.1

PREVENTIVE MAINTENANCE

A preventive maintenance program promotes longevity and trouble-free operation of the infuser. Such a program should include periodic inspection of the infuser, exterior cleaning and sanitizing, and checking for proper operation of the infuser by performing the PVT in Section 5.2.

As a minimum requirement, clean the infuser after each use. Establish a regular cleaning schedule during use. In addition, clean the infuser and perform the PVT as part of any scheduled service or after any repair procedure.

5.1.1

INSPECTING THE INFUSER

Periodically inspect the infuser for signs of defects such as worn accessories, broken instrument connections, or damaged cables. Such an inspection is also applicable after repairing or during cleaning. Replace any damaged or defective external parts.

Inspect the following areas for missing or damaged parts and for cosmetic defects:

- Labels
- Cords
- Switches and touchswitches
- Velcro® straps
- External screws
- Case
- Pole clamp and pads
- Front panel
- Security door
- Accessories
5.1.2
CLEANING THE INFUSER

The following procedures are designed to maintain the infusion system, sustain infuser longevity, and promote trouble-free instrument operation.

Follow hospital protocol for establishing the infuser cleaning schedule.

**WARNING**

DISCONNECT THE INFUSER FROM AC POWER PRIOR TO CLEANING THE INSTRUMENT. FAILURE TO COMPLY WITH THIS WARNING COULD RESULT IN ELECTRICAL SHOCK.

**CAUTION:** Do not immerse the infuser in liquids. Immersion could damage the instrument. Do not allow liquids to enter the infuser electronics compartment.

**CAUTION:** Do not spray cleaning solutions toward any openings in the infuser.

**CAUTION:** Certain cleaning and sanitizing compounds may slowly degrade components made from some plastic materials. Using abrasive cleaners or cleaning solutions not recommended by Abbott Laboratories may result in product damage and, potentially, void the product warranty. Do not use compounds containing combinations of isopropyl alcohol and dimethyl benzyl ammonium chloride.

Clean the exposed surfaces of the infuser with a soft, lint-free cloth dampened with one of the cleaning solutions listed in Table 5-1, *Cleaning Solutions*, or a mild solution of soapy water. Remove soap residue with clear water. Do not use solvents that are harmful to plastic, such as isopropyl alcohol or acetone. Do not use abrasive cleaners.

**CAUTION:** To avoid infuser damage, cleaning solutions should be used only as directed in Table 5-1. The disinfecting properties of cleaning solutions vary; consult the manufacturer for specific information.

**Table 5-1. Cleaning Solutions**

<table>
<thead>
<tr>
<th>Cleaning Solution</th>
<th>Manufacturer</th>
<th>Preparation</th>
</tr>
</thead>
<tbody>
<tr>
<td>Vesphene® II se</td>
<td>Calgon Vestal Laboratories</td>
<td>Per manufacturer’s recommendation</td>
</tr>
<tr>
<td>Manu-Klenz®</td>
<td>Calgon Vestal Laboratories</td>
<td>Per manufacturer’s recommendation</td>
</tr>
<tr>
<td>Formula C™</td>
<td>Diversey Corporation</td>
<td>Per manufacturer’s recommendation</td>
</tr>
<tr>
<td>Super Edisonite®</td>
<td>S. M. Edison Chemical Co.</td>
<td>Per manufacturer’s recommendation</td>
</tr>
<tr>
<td>Household bleach</td>
<td>Various</td>
<td>Per hospital procedures; do not exceed one part bleach in ten parts water</td>
</tr>
<tr>
<td>LifeCare® Germicidal Towelette</td>
<td>Manufactured for Abbott Laboratories</td>
<td>Per manufacturer’s recommendations; use undiluted</td>
</tr>
</tbody>
</table>
5.1.3
SANITIZING THE INFUSER

Sanitize the external surfaces of the infuser using a cleaning solution listed in Table 5-1, Cleaning Solutions.

CAUTION: Do not sterilize the infuser using heat, steam, ethylene oxide (ETO), or radiation; these methods may cause the instrument to malfunction.

Note: Not all cleaning solutions are sanitizers. Check product labeling.

5.2
PERFORMANCE VERIFICATION TEST

As a part of a preventive maintenance schedule, it is recommended that the performance verification test (PVT) be conducted periodically per hospital procedures for compliance to accreditation requirements.

The PVT is used for overall verification of infuser performance and as a diagnostic tool during infuser troubleshooting. The PVT can be used for diagnostic purposes during the troubleshooting of a malfunctioning infuser, and for verification of the overall performance of an infuser as part of a preventive maintenance schedule. In addition, the PVT should be used for performance verification before an infuser is returned to service after repair. If any malfunction is detected as a result of the PVT, refer to Table 6-3, Troubleshooting with the PVT.

Note: The PVT must be performed exactly as described in this manual to assure effective and reliable product evaluation information.

5.2.1
EQUIPMENT REQUIRED

The following equipment, or equivalent, is required to perform the PVT:

- Door key
- PCA vial, standard, List No. 6021-03
- PCA vial, empty, List No. 6021, modified to one ring
- PCA set, List No. 3559-01
- Graduate or marked test tube, readable to 0.2 mL increments or smaller
- Three-way stopcock, List No. 3233
- Digital pressure meter (DPM), (0 to 50 psig), Bio-Tek® DPM II
- Safety analyzer, Dynatech Nevada® 231D
- Butterfly, 21-gauge, List No. 4492
- Test cable, fitted with phone jack with individual banana plugs (compatible with patient control connector on rear of infuser) P/N 561-88416-001
- Parallel network, P/N 561-88419-001
- Digital multimeter (DMM) Fluke® Model 77
Silicone oil, Dow Corning® No. 360, P/N 743-35070-001
X-acto® knife

5.2.2 INSPECTION

Inspect the areas listed in Section 5.1.1, Inspecting the Infuser, for missing or damaged parts and for any cosmetic defects.

5.2.3 INFUSER TEST SETUP

WARNING DURING TESTING, DO NOT CONNECT THE INFUSER TO A PATIENT.

To set up the infuser, proceed as follows:

1. Using the dual-lock mechanism, secure the infuser to an IV pole.

   **Note:** When the security door is locked, the infuser locks to the pole clamp and prevents its removal without a key.

2. Connect the system to AC power unless otherwise specified.
3. Connect the appropriate Abbott PCA set to an Abbott 30 mL PCA vial/injector.
4. Prime the vial and administration set. Hold the vial vertically with the administration set extending from the top. Slowly push down on the injector until all air is cleared from the vial and administration set.

   **CAUTION:** Make certain all caps on the vial and the administration set are removed and all clamps are open when priming syringe.

5.2.4 SERVICE TEST MODE TEST

*Table 5-2, Service Tests,* and *Table 5-3, Service Options,* provide a list of tests and options available in the service test mode. Each test displays prompting information on the LCD screen to serve as a guide through the test. Prompts and descriptions are shown in *Table 5-2.*

**Note:** The patient pendant must be connected to the infuser to complete the service test mode test.

To enter the service test mode, proceed as follows:

1. Confirm that the AC power symbol on the front panel is illuminated.
2. Turn the infuser off and enter the storage mode by opening the security door and pressing the [OFF/RECHG] touchswitch for approximately six seconds. The LCD screen should darken.
3. Press and hold the [YES/ENTER] touchswitch and the [ON] touchswitch simultaneously for approximately three seconds.

The LCD screen displays INT RAM TEST until **SERVICE TEST** appears. The LCD screen also displays the current software version and the total elapsed days since the infuser was placed in service.

**Note:** Tests are self-prompting, giving the option of repeating the test, or going to the next test.

<table>
<thead>
<tr>
<th>Table 5-2. Service Tests</th>
</tr>
</thead>
<tbody>
<tr>
<td><strong>Test</strong></td>
</tr>
<tr>
<td>Software version</td>
</tr>
<tr>
<td>Elapsed days</td>
</tr>
<tr>
<td>Internal RAM</td>
</tr>
<tr>
<td>External RAM ADRS</td>
</tr>
<tr>
<td>External RAM cell</td>
</tr>
<tr>
<td>CPU</td>
</tr>
<tr>
<td>ROM</td>
</tr>
<tr>
<td>Real-time clock</td>
</tr>
<tr>
<td>LED</td>
</tr>
<tr>
<td>LCD</td>
</tr>
<tr>
<td>Keypad</td>
</tr>
<tr>
<td>Indicator</td>
</tr>
<tr>
<td>Alarm</td>
</tr>
<tr>
<td>Motor rotation</td>
</tr>
</tbody>
</table>
### Table 5-2. Service Tests

<table>
<thead>
<tr>
<th>Test</th>
<th>Description</th>
</tr>
</thead>
<tbody>
<tr>
<td>Security door</td>
<td>Test verifies the microcomputer recognizes when the security door is locked or unlocked</td>
</tr>
<tr>
<td>Syringe test</td>
<td>Test verifies the microcomputer can detect the presence and proper positioning of the vial and injector in the syringe driver mechanism</td>
</tr>
<tr>
<td>Empty syringe</td>
<td>Test verifies the microcomputer detects an empty syringe</td>
</tr>
<tr>
<td>Low syringe</td>
<td>Test verifies the microcomputer detects a low syringe (PCA Plus only)</td>
</tr>
</tbody>
</table>

### Table 5-3. Service Options

<table>
<thead>
<tr>
<th>Option</th>
<th>Description</th>
</tr>
</thead>
<tbody>
<tr>
<td>Patient pendant tone selection</td>
<td>Selection provides one of the following options:</td>
</tr>
<tr>
<td></td>
<td>1. Tone with successful PCA dose only (default)</td>
</tr>
<tr>
<td></td>
<td>2. Tone on all attempted PCA doses (placebo)</td>
</tr>
<tr>
<td>12 hour clock selection (PCA Plus only); 12/24 hour clock selection (PCA Plus II Series)</td>
<td>Selection provides one of the following options:</td>
</tr>
<tr>
<td></td>
<td>1. Clock format: 12 hour</td>
</tr>
<tr>
<td></td>
<td>2. Clock format: 24 hour</td>
</tr>
<tr>
<td>Select RX concentration (3.1 software and higher)</td>
<td>Selection provides one of the following options:</td>
</tr>
<tr>
<td></td>
<td>1. Milligrams only</td>
</tr>
<tr>
<td></td>
<td>2. Milligrams and micrograms</td>
</tr>
<tr>
<td></td>
<td>3. Drugs, milligrams, and micrograms</td>
</tr>
</tbody>
</table>

### 5.2.5 DELIVERY ACCURACY TEST

To perform the delivery accuracy test, proceed as follows:

1. Using a PCA vial/syringe with a primed PCA administration set and a 21-gauge butterfly, insert the vial into the infuser vial holder and injector cradle.
2. Purge the infuser at start up, then set up the infuser for 1 mg/mL. Place the butterfly in the graduate. Initiate two loading doses of 10 mL each.
3. Verify the infuser delivers $20 \pm 1$ mL in the graduate.
5.2.6

OCCLUSION TEST

To perform the occlusion test, proceed as follows:

1. Remove the plunger from the syringe. Using an X-acto knife, cut the two top ribs off the syringe plunger, as shown in Figure 5-1, Modification of Syringe Plunger. Make certain that no rough material is loose on the plunger.

2. Apply a light coating of silicone oil to the remaining rib. Replace the plunger in the syringe.

3. Insert the water-filled modified vial with the primed PCA administration set into the cradle assembly. Lubricate the vial with silicone oil before each use.

4. Attach the DPM to the distal end of the administration set through the three-way stopcock.

5. Allow the infuser self test to complete. When the LCD screen displays PURGE THE SYSTEM NOW?, press the [SILENCE/NO] touchswitch.


9. Observe fluid discharge at the end of the stopcock, then close the stopcock.

10. Verify the infuser sounds an alarm and flashes the OCCLUSION message when the pressure gauge indicates the following:

- 008 and lower: 14.0 to 19.0 psig (96.6 to 131.1 kPa)
- 010 and higher: 12.5 to 17.5 psig (86.25 to 120.75 kPa)

11. Open the stopcock to clear the OCCLUSION alarm.

12. Press the [HISTORY] touchswitch and check for correct time; if incorrect, refer to the system operating manual for information on resetting the time.

13. Clear all dose history data from memory by holding down the [OFF/RECHG] touchswitch until the LCD screen, including the backlighted grid, goes blank.

Figure 5-1. Modification of Syringe Plunger
5.2.7

**PCA + CONTINUOUS TEST**

To perform the PCA + CONTINUOUS test, proceed as follows:

1. Remove the modified vial from the cradle assembly and set up a standard water-filled vial with a 21-gauge butterfly on the distal end of the set.

   **CAUTION:** Do not remove the protective cover from the butterfly needle.

2. Insert the vial into the cradle assembly.

3. When the LCD screen displays **PURGE THE SYSTEM NOW?**, press the [SILENCE/NO] touchswitch.


5. When the LCD screen displays **ADMINISTER LOADING DOSE NOW?**, press the [YES/ENTER] touchswitch.


7. When the LCD screen displays **SELECT MODE PCA ONLY?**, press the [SILENCE/NO] touchswitch.

8. When the LCD screen displays **SELECT MODE CONTINUOUS?**, press the [SILENCE/NO] touchswitch.


10. Select a lockout interval of five minutes, then press [YES/ENTER].

11. Select a continuous rate of 20 mg/hr. then press [YES/ENTER].

12. When the LCD screen displays **4 HOUR DOSE LIMIT SET?**, press [YES/ENTER].

13. Select a 4 **HOUR DOSE LIMIT** of 1.5 mg, then press [YES/ENTER].

14. Press the [HISTORY] touchswitch to confirm that the infuser settings are as specified. Close and lock the security door; the display should read: DOOR LOCKED - TOTAL DELIVERED 1 mg.

15. Press the [RESET/START] touchswitch. Verify that the walking bar appears in the LED window, indicating CONTINUOUS mode delivery. Verify that three asterisks appear in the upper left corner. The asterisks indicate PCA dosing is available.

16. Press the patient pendant pushbutton to deliver a PCA dose; verify that a beep sounds and PCA + CONTINUOUS displays. Verify the walking bar appears in the LED window.

17. After approximately two minutes, verify the four hour dose limit has been reached and the KVO rate has begun. Verify the TOTAL DELIVERED displays 2.5 mg.

   **Note:** If delivered prior to setup, the loading dose is not included in the four hour dose limit.

18. Press the patient pendant pushbutton three times; the infuser should not respond.

19. Unlock the security door and remove the injector from the holder; verify that the CHECK INJECTOR message flashes. Reseat injector and verify that the CHECK INJECTOR message disappears.
20. With the security door unlocked, pull the top of the vial away from the vial holder; verify that the CHECK VIAL message flashes. Reinsert the vial and verify that the CHECK VIAL message disappears.

21. Remove the vial from the cradle. Clear all dose history data from memory by holding down the [OFF/RECHG] touchswitch until the LCD screen, including the backlighted grid, goes blank.

5.2.8

PATIENT CONTROL JACK TEST

To perform the patient control jack test, proceed as follows:

1. Disconnect the patient pendant cable from the patient control jack on the back of the infuser.
2. Connect the test cable phone jack to the patient control connector; connect the leads on the other end of the cable to the terminals of the DMM as appropriate. Set the DMM to 10 volt DC scale.
3. Turn on the infuser and verify a DMM reading of greater than or equal to 4.5 VDC. Remove the test cable from the infuser.

5.2.9

PATIENT PENDANT ASSEMBLY TEST

To perform the patient pendant assembly test, proceed as follows:

1. Disconnect the patient pendant from the patient control jack on the back of the infuser.
2. Unscrew the shield from the patient pendant phono plug.
3. Set the DMM to measure continuity, then attach a DMM test lead to each patient pendant phone plug connection (polarity is not critical).
4. Press and hold the patient pendant switch. Verify the DMM displays continuity (0 ohm) and the continuity beeper sounds.

**Note:** A value of up to 3 ohms may be indicated due to patient pendant cable assembly internal resistance.

5. Press and hold the patient pendant switch. Grasp the patient pendant cable approximately 4 inches (10 cm) above the patient pendant switch and rotate the cable three to four times while bending it approximately 30 degrees. Verify the DMM continues to display continuity.
6. Press and hold the patient pendant switch. Grasp the patient pendant cable approximately 4 inches (10 cm) above the phone plug and rotate the cable three to four times while bending it approximately 30 degrees. Verify the DMM continues to display continuity.
8. Connect the patient pendant cable to the patient control jack. Secure the locking nut (if applicable).
5.2.10

BATTERY CHARGER TEST

To perform the battery charger test, proceed as follows:

1. Disconnect the infuser from AC power.
2. Remove the battery cover and disconnect the battery.
3. Connect the parallel network to the charging circuit connector.
4. Connect the infuser to AC power. Power on the infuser. Measure voltage across the network; the DMM should read 9.4 ± 0.15 VDC (14.5 ± 2 VDC with a battery boost charger PWA).

**Note:** The infuser must be powered on before obtaining the DMM reading.

**Note:** See Section 4.2.3, Battery Boost Charger PWA, to identify infuser configurations with a battery boost charger PWA.

5. Remove the parallel network. Reconnect and re-install the battery; make certain the battery wires are not pinched. Re-install the battery cover.

5.2.11

ELECTRICAL SAFETY TEST

To perform the electrical safety test, proceed as follows:

1. Connect the AC power cord to an approved safety analyzer. Leakage current must not exceed 50 microamperes, but must be greater than 2 microamperes (open ground).

2. Using the safety analyzer, measure the resistance between the ground lug of the AC connector and exposed metal parts, such as the door key lock, patient pendant connector, or the exposed screw heads on the pole clamp. Note that the door hinge is not grounded. Resistance should not exceed 0.1 ohm.

**Note:** Exposed metal on the infuser is isolated from ground.
5.2.12

PRINTER TEST

To perform the printer test, proceed as follows:

1. Connect the infuser to AC power.
2. Connect the printer cable (Centronics interface) to the printer port located on the back of the infuser.
3. Power on the printer and verify it is on line.
4. Open the infuser security door, then insert the syringe into the cradle assembly. Verify the infuser begins the self test.
5. After the self test completes, the PURGE SYSTEM NOW? prompt displays. Press the [PRINT] touchswitch.
6. Confirm END OF RECORD is printed on the history and event log printout. It may be necessary to take the printer off line, then form feed the last partial page of the printout from the printer.
7. Compare the printout to Figure 5-2, Sample Printer Test. Date, time, and parameters will vary.

Note: Depending on the infuser configuration or the software revision level, the printout may not have an HOUR-BY-HOUR: entry.
5.2.13 END OF PERFORMANCE VERIFICATION TEST

At the end of the PVT, clear all dose history data from memory by holding down the [OFF/RECHG] touchswitch until the display goes blank. If all tests have been successful, return the infuser to service. If any of the tests fail, refer to the troubleshooting information in Section 6, Troubleshooting, or contact Abbott Laboratories.
5.3

PVT DATA FORM

List Number: ________________ Serial Number: ________________

Inspection
1. Verify that all labels are on the infuser. Pass___ Fail___
2. Inspect the electrical cord for damage or foreign material. Pass___ Fail___
3. Verify that the control panel switches have no cracks or other damage. Pass___ Fail___
4. Verify that the two Velcro straps are present and not damaged. Pass___ Fail___
5. Verify that all screws are secured and tight. Pass___ Fail___
6. Inspect the case for cracks or stains. Pass___ Fail___
7. Inspect the pole clamp and pads for damage. Pass___ Fail___
8. Verify that the front panel has no cracks or cosmetic defects. Pass___ Fail___
9. Verify that the two bottom pressure pads are present and do not show excessive wear. Pass___ Fail___
10. Inspect the security door for any cracks or other damage. Pass___ Fail___
11. Inspect the patient pendant for cracked connector housing or cable damage. Pass___ Fail___

Service Test Mode
1. Record software version _________
2. All tests in service mode pass successfully. Pass___ Fail___

Delivery Accuracy Test
1. Set up PCA for a concentration of 1 mg/mL. Deliver two loading doses of 10.0 mL (Total = 20.0 mL) and verify delivery accuracy. Record __________ mL (specification = 19.0 to 21.0 mL) Pass___ Fail___

Occlusion Test
1. Initiate 5 mg loading dose and verify occlusion alarm and flashing OCCLUSION message. Record __________ psig Specification:
   -008 and below: 14.0 to 19.0 psig (96.6 to 131.1 kPa)
   -010 and above: 12.4 to 17.5 psig (86.25 to 120.75 kPa) Pass___ Fail___

PCA + Continuous Test
1. Verify the walking bar is displayed during continuous mode delivery. Pass___ Fail___
2. Verify that three asterisks appear in the upper left corner, indicating PCA dosing available. Pass___ Fail___
3. Verify a beep occurs and PCA + CONTINUOUS is displayed when the pendant is pressed. Pass___ Fail___
4. Approximately two minutes after pressing the patient pendant, verify that 4 HR LIM REACHED is displayed, KVO rate begins, and TOTAL DELIVERED display reads 2.5 mg. Pass___ Fail___
5. Verify that after pressing patient pendant three more times, there is no response from the infuser. Pass___ Fail___
6. Verify that by pulling the syringe vial away from the holder, a check vial alarm occurs. Pass___ Fail___
7. Verify that by pulling the syringe injector away from the injector holder, the check injector alarm occurs. Pass___ Fail___
Patient Control Jack Test
1. Verify that the patient control input measures greater than or equal to 4.50 VDC. Pass___ Fail___

Patient Pendant Assembly Test
1. Connect DMM to patient pendant phone plug connections. Set DMM to measure continuity. Press and hold the patient pendant switch and verify the DMM displays continuity. Pass___ Fail___
2. Press and hold the patient pendant switch while rotating the control cable. Verify the DMM displays continuity. Pass___ Fail___
3. Press and hold the patient pendant switch while rotating the control cable. Verify the DMM displays continuity. Pass___ Fail___

Battery Charger Test
1. Connect the parallel network and measure the charger voltage across the network. Record __________ VDC Specification:
   9.4 ± 0.15 VDC without battery boost circuit
   14.5 ± 2.0 VDC with battery boost circuit Pass___ Fail___

Electrical Safety Test
1. Record leakage current. __________ Acceptable result 50 µA. Pass___ Fail___
2. Record ground lug resistance. __________ Acceptable result 0.1 ohms. Pass___ Fail___

Printer Test
1. Confirm END OF RECORD is printed on the history and event log printout. Pass___ Fail___
2. Compare the printout to Figure 5-2, Sample Printer Test. Pass___ Fail___

Performance Verification Test performed by __________________________Date:_____________

Test Equipment:
Pressure Meter # __________
Safety Analyzer # __________
DMM # _____________________
Section 6
TROUBLESHOOTING

This section contains information on obtaining technical assistance from Abbott Laboratories Technical Support Operations, audible alarm and alarm messages, alarm code history, and troubleshooting the infuser.

6.1
TECHNICAL ASSISTANCE

For technical assistance, product return authorization, and to order parts, accessories, or manuals within the United States, contact Abbott Laboratories Technical Support Operations.

1-800-241-4002

Send all authorized, prepaid returns to the following address:

Abbott Laboratories
Technical Support Operations
755 Jarvis Drive
Morgan Hill, California 95037

From outside the United States, contact the nearest Abbott Laboratories representative.

Contact the Abbott Laboratories professional hospital representative or Abbott Laboratories Technical Support Operations for information about the Abbott Laboratories Extended Warranty Program for the PCA Plus and Plus II Series Infuser.

6.2
DIAGNOSTIC MODE

Note: A printer must be connected to the infuser when using the diagnostic mode.

The PCA Plus II Series Infuser with 3.1 software and above has a diagnostic mode that allows retrieval of the last 200 events and malfunction codes. This information is retained indefinitely in a battery-backed RAM on the CPU/display PWA.

To access diagnostic information, the infuser must be in storage mode (cold-start condition) (see Section 4.1.1.2, Cold-Start Sequence).

Simultaneously press and hold the [PURGE SYSTEM] and [ON] touchswitches for approximately three seconds; the LCD screen first displays INT RAM TEST, then DIAGNOSTIC MODE. The LCD screen also displays the software revision and the total elapsed days since the infuser was placed into service.
Press the [YES/ENTER] touchswitch, then press [PRINT] to print out the event log, or press [SILENCE/NO] to continue to the malfunction log screen.

The diagnostic mode event log lists the last 200 events. The malfunction log lists the last 200 malfunction codes (see Figure 6-1, Sample Event Log and Malfunction Log).

![Figure 6-1: Sample Event Log and Malfunction Log](image)

### 6.3 ALARM AND DISPLAY MESSAGE

#### OVERVIEW

Under most alarm conditions, the infuser stops operating, generates an audible alarm, and displays an alarm message on the LCD screen.

**CAUTION:** For those patients who are likely to be adversely affected by unintended operations and failures, including interrupted medication or fluid delivery from the device, close supervision and provision for immediate corrective action should be provided.

**CAUTION:** Do not return the infuser to service after an alarm or malfunction without determining and correcting the cause of the alarm or malfunction.
6.3.1

AUDIBLE ALARMS

The infuser sounds an alarm in the event of any abnormal condition. The alarm volume can be set to LOW, MEDIUM, or HIGH by a three-way switch on the back of the infuser.

The infuser produces five different tones, as follows:

1. A single, short tone indicates the infuser senses a touchswitch being pressed.
2. Two short tones indicate the start of a power-up sequence.
3. Three short tones indicate that a touchswitch was pressed that is invalid for the infuser’s current state of operation or is not an allowable response to a prompt on the LCD screen.
4. Two short-tone, long-tone sequences, repeating every few seconds, indicate an alarm condition. The LCD screen displays the type of alarm in flashing letters.
5. A continuous tone indicates an infuser malfunction. The LCD screen may or may not display further information, depending on the malfunction.

6.3.2

ALARM CAUSES AND ACTIONS

The infuser will stop delivery when any of the following alarms occur:

- CHECK VIAL
- CHECK INJECTOR
- CHECK SYRINGE
- EMPTY SYRINGE
- OCCLUSION
- MALFUNCTION

These alarms, their possible causes, and corrective actions are detailed in Table 6-2, Infuser Troubleshooting.

6.3.3

MALFUNCTION CODES

Table 6-1, Malfunction Codes, describes the malfunction codes as they appear when an alarm sounds and MALFUNCTION appears on the LCD screen.
<table>
<thead>
<tr>
<th>Code</th>
<th>Malfunction</th>
<th>Corrective Action</th>
</tr>
</thead>
<tbody>
<tr>
<td>1A</td>
<td>MALF line went low and could not return to normal</td>
<td>Replace CPU/display PWA</td>
</tr>
<tr>
<td>1B</td>
<td>MALF line went low and could not return to normal</td>
<td>Replace CPU/display PWA</td>
</tr>
<tr>
<td>1C</td>
<td>MALF line did not go low soon enough after power-up</td>
<td>Replace CPU/display PWA</td>
</tr>
<tr>
<td>1D</td>
<td>Malfunction line went high too soon</td>
<td>Replace CPU/display PWA</td>
</tr>
<tr>
<td>1E</td>
<td>Watchdog circuit failure during real-time operation</td>
<td>Replace CPU/display PWA</td>
</tr>
<tr>
<td>2A</td>
<td>Power-up diagnostic failed</td>
<td>Replace CPU/display PWA</td>
</tr>
<tr>
<td>2B</td>
<td>Nonvolatile RAM checksum test failed</td>
<td>Replace CPU/display PWA</td>
</tr>
<tr>
<td>3A</td>
<td>Motor failed to rotate</td>
<td>Lubricate lead screw</td>
</tr>
<tr>
<td></td>
<td></td>
<td>Replace power supply PWA</td>
</tr>
<tr>
<td></td>
<td></td>
<td>Replace motor assembly</td>
</tr>
<tr>
<td>3B</td>
<td>MOTOR BAD signal from motor driver circuitry</td>
<td>Replace power supply PWA</td>
</tr>
<tr>
<td></td>
<td></td>
<td>Replace motor assembly</td>
</tr>
<tr>
<td>4A</td>
<td>LCD timeout error</td>
<td>Replace CPU/display PWA</td>
</tr>
<tr>
<td>4B</td>
<td>LCD data readback error</td>
<td>Replace CPU/display PWA</td>
</tr>
<tr>
<td>4C</td>
<td>Nonvolatile RAM test failed</td>
<td>Replace CPU/display PWA</td>
</tr>
<tr>
<td>4D</td>
<td>EPROM checksum test failed</td>
<td>Replace CPU/display PWA</td>
</tr>
<tr>
<td>4E</td>
<td>Not used</td>
<td></td>
</tr>
<tr>
<td>4F</td>
<td>LED segment defective</td>
<td>Replace CPU/display PWA</td>
</tr>
<tr>
<td>4G</td>
<td>Real-time clock stopped</td>
<td>Replace CPU/display PWA</td>
</tr>
<tr>
<td>4H</td>
<td>Touchswitch stuck on front panel</td>
<td>Replace front panel</td>
</tr>
<tr>
<td></td>
<td></td>
<td>Replace CPU/display PWA</td>
</tr>
<tr>
<td>4I</td>
<td>Not used</td>
<td></td>
</tr>
<tr>
<td>4J</td>
<td>Not used</td>
<td></td>
</tr>
<tr>
<td>4K</td>
<td>LED error</td>
<td>Replace CPU/display PWA</td>
</tr>
</tbody>
</table>
### Table 6-1. Malfunction Codes

<table>
<thead>
<tr>
<th>Code</th>
<th>Malfunction</th>
<th>Corrective Action</th>
</tr>
</thead>
<tbody>
<tr>
<td>4L</td>
<td>Voltage error</td>
<td>Replace power supply PWA</td>
</tr>
<tr>
<td>4M</td>
<td>Not used</td>
<td></td>
</tr>
<tr>
<td>4N</td>
<td>Error in timer T1</td>
<td>Replace CPU/display PWA</td>
</tr>
<tr>
<td>4O</td>
<td>Error in timer T2</td>
<td>Replace CPU/display PWA</td>
</tr>
<tr>
<td>4P</td>
<td>Program error</td>
<td>Replace CPU/display PWA</td>
</tr>
<tr>
<td>4Q</td>
<td>CPU stack error</td>
<td>Replace CPU/display PWA</td>
</tr>
<tr>
<td>4R</td>
<td>Analog-to-digital converter error</td>
<td>Replace CPU/display PWA</td>
</tr>
<tr>
<td>5A</td>
<td>Undetermined error</td>
<td>Contact Abbott Laboratories</td>
</tr>
</tbody>
</table>

### 6.4 INFUSER TROUBLESHOOTING

**Table 6-2. Infuser Troubleshooting** shows failure symptoms and probable causes. **Table 6-2** is organized by the alarm messages that appear on the infuser LCD screen.

Before troubleshooting, turn the infuser power off, then on. Allow the self test to complete. If an alarm or malfunction persists, carefully inspect the infuser for damage as described in **Section 5.1.1, Inspecting the Infuser**. If alarm or malfunction persists, perform the corrective action specified in **Table 6-2**, or contact Abbott Laboratories.

### Table 6-2. Infuser Troubleshooting

<table>
<thead>
<tr>
<th>Alarm Message/ Fault Symptom</th>
<th>Possible Cause</th>
<th>Corrective Action</th>
</tr>
</thead>
<tbody>
<tr>
<td>CHECK VIAL</td>
<td>Vial is improperly installed or missing</td>
<td>Properly insert vial in the vial cradle <em>(see system operating manual)</em></td>
</tr>
<tr>
<td></td>
<td></td>
<td>Adjust vial sensor switch <em>(see Section 7.2.4)</em></td>
</tr>
<tr>
<td>CHECK INJECTOR</td>
<td>Injector is improperly installed or missing</td>
<td>Properly insert injector in the driver/retainer assembly <em>(see system operating manual)</em></td>
</tr>
<tr>
<td></td>
<td></td>
<td>Adjust injector sensor switch <em>(see Section 7.2.3)</em></td>
</tr>
<tr>
<td>Alarm Message/ Fault Symptom</td>
<td>Possible Cause</td>
<td>Corrective Action</td>
</tr>
<tr>
<td>-------------------------------</td>
<td>----------------</td>
<td>-------------------</td>
</tr>
<tr>
<td>CHECK SYRINGE</td>
<td>Syringe (vial and injector) improperly installed or missing</td>
<td>Properly insert syringe in the cradle (see system operating manual)</td>
</tr>
<tr>
<td>LOW SYRINGE (PCA Plus only)</td>
<td>Approximately 5 mL of solution remaining in syringe</td>
<td>Note alarm and press [SILENCE/NO] touchswitch to silence alarm</td>
</tr>
<tr>
<td>EMPTY SYRINGE</td>
<td>Syringe empty (approximately 1 mL may remain)</td>
<td>Press [SILENCE/NO] touchswitch to silence alarm. Unlock and open security door, remove empty syringe and turn infuser off. Prepare and insert vial/injector. Complete setup (see system operating manual)</td>
</tr>
<tr>
<td>CHECK SETTINGS</td>
<td>Security door closed and locked before setup was complete</td>
<td>Open security door and assure settings are properly entered</td>
</tr>
<tr>
<td>CHECK 4-HR LIMIT</td>
<td>Four-hour dose limit is set less than PCA dose</td>
<td>Confirm PCA dose is less than or equal to four-hour dose limit</td>
</tr>
<tr>
<td>DOOR OPEN</td>
<td>Security door has been left open for more than two minutes without a touchswitch being pressed</td>
<td>Close and lock security door after all parameters are entered</td>
</tr>
<tr>
<td>Alarm Message/ Fault Symptom</td>
<td>Possible Cause</td>
<td>Corrective Action</td>
</tr>
<tr>
<td>------------------------------</td>
<td>----------------</td>
<td>------------------</td>
</tr>
<tr>
<td>OCCLUSION</td>
<td>Kinked or otherwise occluded tubing, occluded venipuncture device, or closed slide clamp</td>
<td>Open security door if closed. Relieve backpressure by squeezing and releasing the cradle release handles</td>
</tr>
<tr>
<td></td>
<td>Occlusion pressure out of calibration</td>
<td>Adjust occlusion pressure (see Section 7.2.7.3 or Section 7.2.7.4)</td>
</tr>
<tr>
<td></td>
<td></td>
<td>Replace occlusion pressure switch (see Section 7.3.26)</td>
</tr>
<tr>
<td>INFUSER IN RESET</td>
<td>[RESET/START] touched when security door locked (in CONTINUOUS or PCA+ CONTINUOUS mode) and delivery in progress</td>
<td>Press the [RESET/START] touchswitch to resume delivery</td>
</tr>
<tr>
<td>Note: INFUSER IN RESET message also appears after an alarm condition has been cleared. Press the [RESET/START] touchswitch to resume delivery</td>
<td></td>
<td></td>
</tr>
<tr>
<td>LOW BATTERY (flashing battery symbol)</td>
<td>Battery is near discharge level</td>
<td>Connect infuser to AC power</td>
</tr>
<tr>
<td>Note: See Section 4.4, Battery Operation Overview, for detailed information about battery-powered operation</td>
<td></td>
<td></td>
</tr>
<tr>
<td>No battery operation or short operating time to LOW BATTERY alarm</td>
<td>Battery pack not fully charged</td>
<td>Recharge battery pack for 16 hours</td>
</tr>
<tr>
<td></td>
<td>Battery pack is defective or has exceeded useful life</td>
<td>Test battery charging circuit (see Section 5.2.10). If test passes, replace battery pack (see Section 7.3.5). If test fails, replace power supply PWA (see Section 7.3.15)</td>
</tr>
<tr>
<td></td>
<td></td>
<td>Replace battery pack (see Section 7.3.5)</td>
</tr>
</tbody>
</table>
### Table 6-2. Infuser Troubleshooting

<table>
<thead>
<tr>
<th>Alarm Message/ Fault Symptom</th>
<th>Possible Cause</th>
<th>Corrective Action</th>
</tr>
</thead>
<tbody>
<tr>
<td>No AC or battery operation</td>
<td>Fuse blown</td>
<td>Replace fuse <em>(see Section 7.3.6)</em></td>
</tr>
<tr>
<td></td>
<td>Defective power supply PWA</td>
<td>Measure AC voltage between J3 pins 2 and 3. If voltage is between 11 and 17 VAC, replace power supply PWA <em>(see Section 7.3.15)</em>. If voltage is not between 11 and 17 VAC, contact Abbott Laboratories</td>
</tr>
<tr>
<td>MALFUNCTION XX (Error code displayed in place of XX)</td>
<td>Infuser self-test programs have detected an internal failure</td>
<td>Note malfunction code displayed. Press [OFF/RECHG] touchswitch to silence alarm; turn infuser off. Remove infuser from service and refer to <em>Table 6-1</em></td>
</tr>
<tr>
<td>No audible alarm or audible alarm level control</td>
<td>Defective audible level switch</td>
<td>Replace audible level switch <em>(see Section 7.3.36)</em></td>
</tr>
<tr>
<td></td>
<td>Defective power supply PWA</td>
<td>Replace power supply PWA <em>(see Section 7.3.15)</em></td>
</tr>
</tbody>
</table>

### 6.5 PVT TROUBLESHOOTING

*Table 6-3. Troubleshooting with the PVT* describes failures that may be detected during the PVT *(see Section 5.2)*. If an error code displays, see *Section 6.3, Alarm and Display Message Overview*.

<table>
<thead>
<tr>
<th>PVT Test Failure</th>
<th>Possible Cause</th>
<th>Corrective Action</th>
</tr>
</thead>
<tbody>
<tr>
<td>Service test mode test</td>
<td>Infuser not in cold start</td>
<td>Hold [OFF/RECHG] for six seconds</td>
</tr>
<tr>
<td><em>Section 5.2.4</em></td>
<td>Defective touchswitch</td>
<td>Replace front panel <em>(see Section 7.3.40)</em></td>
</tr>
<tr>
<td></td>
<td>Defective CPU/display PWA</td>
<td>Replace CPU/display PWA <em>(see Section 7.3.16)</em></td>
</tr>
<tr>
<td>PVT Test Failure</td>
<td>Possible Cause</td>
<td>Corrective Action</td>
</tr>
<tr>
<td>----------------------</td>
<td>---------------------------------</td>
<td>-----------------------------------------------------------------------------------</td>
</tr>
<tr>
<td>Software version</td>
<td>Defective CPU/display PWA</td>
<td>Replace CPU/display PWA (see Section 7.3.16)</td>
</tr>
<tr>
<td>Elapsed days</td>
<td>Defective CPU/display PWA</td>
<td>Replace CPU/display PWA (see Section 7.3.16)</td>
</tr>
<tr>
<td>Internal RAM</td>
<td>Defective CPU/display PWA</td>
<td>Replace CPU/display PWA (see Section 7.3.16)</td>
</tr>
<tr>
<td>External RAM ADRS</td>
<td>Defective CPU/display PWA</td>
<td>Replace CPU/display PWA (see Section 7.3.16)</td>
</tr>
<tr>
<td>External RAM cell</td>
<td>Defective CPU/display PWA</td>
<td>Replace CPU/display PWA (see Section 7.3.16)</td>
</tr>
<tr>
<td>CPU</td>
<td>Defective CPU/display PWA</td>
<td>Replace CPU/display PWA (see Section 7.3.16)</td>
</tr>
<tr>
<td>ROM</td>
<td>Defective CPU/display PWA</td>
<td>Replace CPU/display PWA (see Section 7.3.16)</td>
</tr>
<tr>
<td>Real time clock</td>
<td>Defective CPU/display PWA</td>
<td>Replace CPU/display PWA (see Section 7.3.16)</td>
</tr>
<tr>
<td>LED</td>
<td>Defective CPU/display PWA</td>
<td>Replace CPU/display PWA (see Section 7.3.16)</td>
</tr>
<tr>
<td>LCD</td>
<td>Defective CPU/display PWA</td>
<td>Replace CPU/display PWA (see Section 7.3.16)</td>
</tr>
<tr>
<td>Keypad</td>
<td>Loose connection/damaged ribbon cable</td>
<td>Check ribbon cable from touchswitch to J13 on CPU/display PWA. Replace front panel if necessary (see Section 7.3.40)</td>
</tr>
<tr>
<td></td>
<td></td>
<td>Check ribbon cable from J12 on CPU/display PWA to J7 on power supply PWA. Replace ribbon cable if necessary (see Section 7.3.19)</td>
</tr>
<tr>
<td></td>
<td>Defective touchswitch</td>
<td>Replace front panel (see Section 7.3.40)</td>
</tr>
<tr>
<td></td>
<td>Defective CPU/display PWA</td>
<td>Replace CPU/display PWA (see Section 7.3.16)</td>
</tr>
<tr>
<td>PVT Test Failure</td>
<td>Possible Cause</td>
<td>Corrective Action</td>
</tr>
<tr>
<td>------------------</td>
<td>----------------</td>
<td>-------------------</td>
</tr>
<tr>
<td>Indicator</td>
<td>Loose connection/defective ribbon cable</td>
<td>Check ribbon cable from J12 on CPU/display PWA to J7 on power supply PWA</td>
</tr>
<tr>
<td></td>
<td>Defective CPU/display PWA</td>
<td>Replace CPU/display PWA (see Section 7.3.16)</td>
</tr>
<tr>
<td></td>
<td>Defective power supply PWA</td>
<td>Replace power supply PWA (see Section 7.3.15)</td>
</tr>
<tr>
<td>Alarm</td>
<td>Loose connection/defective ribbon cable</td>
<td>Check connection at J8 on power supply PWA</td>
</tr>
<tr>
<td></td>
<td>Defective alarm volume switch</td>
<td>Replace alarm volume switch or interconnect PWA (see Section 7.3.36 or Section 7.3.37)</td>
</tr>
<tr>
<td></td>
<td>Defective piezoelectric alarm</td>
<td>Replace piezoelectric alarm or interconnect PWA (see Section 7.3.34 or Section 7.3.37)</td>
</tr>
<tr>
<td></td>
<td>Defective power supply PWA</td>
<td>Replace power supply PWA (see Section 7.3.15)</td>
</tr>
<tr>
<td></td>
<td>Defective CPU/display PWA</td>
<td>Replace CPU/display PWA (see Section 7.3.16)</td>
</tr>
<tr>
<td>Motor rotation</td>
<td>Lead screw dirty or needs lubrication</td>
<td>Lubricate lead screw (see Section 7.2.11)</td>
</tr>
<tr>
<td></td>
<td>Loose connection</td>
<td>Check connector J5 on power supply PWA</td>
</tr>
<tr>
<td></td>
<td>Defective power supply PWA</td>
<td>Replace power supply PWA (see Section 7.3.15)</td>
</tr>
<tr>
<td></td>
<td>Defective motor assembly</td>
<td>Replace motor assembly (see Section 7.3.28 or Section 7.3.29)</td>
</tr>
<tr>
<td>PVT Test Failure</td>
<td>Possible Cause</td>
<td>Corrective Action</td>
</tr>
<tr>
<td>------------------</td>
<td>----------------</td>
<td>------------------</td>
</tr>
<tr>
<td>Security door</td>
<td>Loose connection</td>
<td>Check connector J8 and J7 on power supply PWA</td>
</tr>
<tr>
<td></td>
<td>Door lock switch out of adjustment or defective</td>
<td>Adjust or replace door lock switch (see Section 7.3.41)</td>
</tr>
<tr>
<td></td>
<td>Defective power supply PWA</td>
<td>Replace power supply PWA (see Section 7.3.15)</td>
</tr>
<tr>
<td>Syringe test</td>
<td>Loose connection</td>
<td>Check connector J6 or J4 on power supply PWA</td>
</tr>
<tr>
<td></td>
<td>Injector sensor switch defective</td>
<td>Replace injector sensor switch (see Section 7.3.27)</td>
</tr>
<tr>
<td></td>
<td>Vial sensor switch out of adjustment or defective</td>
<td>Adjust or replace vial sensor switch (see Section 7.3.22)</td>
</tr>
<tr>
<td></td>
<td>Defective power supply PWA</td>
<td>Replace power supply PWA (see Section 7.3.15)</td>
</tr>
<tr>
<td>Low/empty syringe</td>
<td>Loose connection</td>
<td>Check connector J4 on power supply PWA</td>
</tr>
<tr>
<td></td>
<td>Syringe low sensor switch out of adjustment or defective (PCA only)</td>
<td>Adjust or replace syringe low switch (see Section 7.3.25)</td>
</tr>
<tr>
<td></td>
<td>Syringe empty sensor switch out of adjustment or defective</td>
<td>Adjust or replace syringe empty switch (see Section 7.3.25)</td>
</tr>
<tr>
<td></td>
<td>Defective power supply PWA</td>
<td>Replace power supply PWA (see Section 7.3.15)</td>
</tr>
<tr>
<td>Patient pendent tone selection</td>
<td>Defective CPU/display PWA</td>
<td>Replace CPU/display PWA (see Section 7.3.16)</td>
</tr>
<tr>
<td>12/24 hour clock selection (PCA Plus)</td>
<td>Defective CPU/display PWA</td>
<td>Replace CPU/display PWA (see Section 7.3.16)</td>
</tr>
<tr>
<td>Select RX concentration (3.1 software and higher)</td>
<td>Incorrect software revision</td>
<td>Verify software is 3.1 or higher</td>
</tr>
<tr>
<td>PVT Test Failure</td>
<td>Possible Cause</td>
<td>Corrective Action</td>
</tr>
<tr>
<td>------------------------------------------</td>
<td>---------------------------------------</td>
<td>--------------------------------------------------------</td>
</tr>
<tr>
<td>Delivery accuracy test</td>
<td>PCA set not primed properly</td>
<td>Prime set/check clamps</td>
</tr>
<tr>
<td></td>
<td>Defective power supply PWA</td>
<td>Replace power supply PWA (see Section 7.3.15)</td>
</tr>
<tr>
<td></td>
<td>Defective motor assembly</td>
<td>Replace motor assembly (see Section 7.3.28 or Section 7.3.29)</td>
</tr>
<tr>
<td>Occlusion test</td>
<td>Syringe not modified properly</td>
<td>Check modification of syringe plunger</td>
</tr>
<tr>
<td></td>
<td>Set not installed or primed properly</td>
<td>Prime set</td>
</tr>
<tr>
<td></td>
<td>Occlusion pressure switch out of</td>
<td>Adjust or replace occlusion pressure switch (see Section 7.2.7.3 or Section 7.2.7.4 and Section 7.3.26)</td>
</tr>
<tr>
<td></td>
<td>adjustment or defective</td>
<td>Replace power supply PWA (see Section 7.3.15)</td>
</tr>
<tr>
<td>PCA + continuous test</td>
<td>Set not installed or not primed</td>
<td>Check installation and prime set</td>
</tr>
<tr>
<td></td>
<td>properly</td>
<td>Check infuser settings</td>
</tr>
<tr>
<td></td>
<td>Infuser not set up properly</td>
<td></td>
</tr>
<tr>
<td>Patient control jack test</td>
<td>Loose connection</td>
<td>Check connector J8 on the power supply PWA</td>
</tr>
<tr>
<td></td>
<td>Defective interconnect PWA cable</td>
<td>Replace interconnect PWA cable (see Section 7.3.38)</td>
</tr>
<tr>
<td></td>
<td>Defective patient pendant jack</td>
<td>Replace patient pendant jack or interconnect PWA (see Section 7.3.35 or Section 7.3.37)</td>
</tr>
<tr>
<td>Patient pendant assembly test</td>
<td>Defective patient pendant assembly</td>
<td>Replace patient pendant assembly</td>
</tr>
</tbody>
</table>
### Table 6-3. Troubleshooting with the PVT

<table>
<thead>
<tr>
<th>PVT Test Failure</th>
<th>Possible Cause</th>
<th>Corrective Action</th>
</tr>
</thead>
<tbody>
<tr>
<td>Battery charger test</td>
<td>Loose connection</td>
<td>Check connector J3 on the power supply PWA</td>
</tr>
<tr>
<td>Section 5.2.10</td>
<td></td>
<td>Check connector P1 and P2 on boost charger PWA</td>
</tr>
<tr>
<td></td>
<td>Defective boost charger PWA</td>
<td>Replace boost charger PWA (see Section 7.3.39)</td>
</tr>
<tr>
<td></td>
<td>Defective power supply PWA</td>
<td>Replace power supply PWA (see Section 7.3.15)</td>
</tr>
<tr>
<td>Electrical safety test</td>
<td>Insufficient earth connection</td>
<td>Check for adequate earth connection</td>
</tr>
<tr>
<td>Section 5.2.11</td>
<td>Defective AC power cord plug</td>
<td>Replace AC power cord plug (see Section 7.3.13)</td>
</tr>
<tr>
<td></td>
<td>Defective AC power cord</td>
<td>Replace AC power cord (see Section 7.3.12)</td>
</tr>
</tbody>
</table>
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Section 7

REPLACEABLE PARTS AND REPAIRS

This section itemizes all parts and subassemblies of the infuser that are repairable within the scope of this manual. In addition, this section describes replacement procedures for all listed parts.

**WARNING**
POSSIBLE EXPLOSION HAZARD IF THE INFUSER IS SERVICED IN THE PRESENCE OF FLAMMABLE ANESTHETICS.

**WARNING**
UNLESS OTHERWISE INDICATED, DISCONNECT INFUSER FROM AC POWER BEFORE PERFORMING ADJUSTMENT OR REPLACEMENT PROCEDURES.

CAUTION: Use proper ESD grounding techniques when handling components. Wear an antistatic wrist strap and use an ESD-protected workstation. Store the PWA in an antistatic bag before placing it on any surface.

7.1

REPLACEABLE PARTS LIST

Replaceable parts for the infuser are itemized in the spare parts price list and are identified in Figure 9-1, IPB for the Infuser. Table 9-2, IPB for the Infuser, identifies each infuser part by an index number that correlates to Figure 9-1.

To request a copy of the current spare parts price list, contact Abbott Laboratories. For convenient reference, insert a copy of the spare parts price list here.
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7.2
ADJUSTMENT PROCEDURES

Unless otherwise indicated, the following procedures require an empty syringe to be installed in the cradle.

---

**WARNING**

UNLESS OTHERWISE INDICATED, DISCONNECT INFUSER FROM AC POWER BEFORE PERFORMING ADJUSTMENT PROCEDURES.

---

7.2.1
REQUIRED TOOLS AND MATERIALS

The following standard handtools, special tools, and materials, or equivalents, are required for the adjustment procedures in this section. The tools and materials required for a specific adjustment procedure are listed at the beginning of each procedure.

- Set of Allen wrenches
- Set of nutdrivers
- Set of open end wrenches
- Flat blade screwdriver
- No. 0 Phillips screwdriver
- No. 2 Phillips screwdriver
- X-acto knife
- Digital pressure meter (DPM), Bio-Tek DPM II
- Red GLPT insulating varnish
- Water-filled syringe
- Empty syringe
- Modified syringe
- Three-way stopcock
- IV administration set
- Permanent marker
- Syringe limit gauge
- Trim pot tool
- Silicone oil, Dow Corning® 360 Mineral Fluid
- Grease, Braycote® 804
- Needle nose pliers
- Electro-Wash 2000® or isopropyl alcohol
- Lint-free cloth or cotton swabs
- Small brush
7.2.2  
SEPARATING FRONT AND REAR CASE ASSEMBLIES

CAUTION: Use proper ESD grounding techniques when handling components. Wear an antistatic wrist strap and use an ESD-protected workstation. Store the PWA in an antistatic bag before placing it on any surface.

The infuser front and rear case assemblies must be separated before performing any adjustment procedure.

The recommended tools for this procedure are as follows: No. 2 Phillips screwdriver, 1/4 inch nutdriver, 5/16 inch nutdriver, and permanent marker.

To separate the infuser front and rear case assemblies, refer to Figure 7-1, Separating Front and Rear Case Assemblies, then proceed as follows:

1. Disconnect the infuser from AC power.
2. Set the infuser upright on a flat surface. Confirm the security door is unlocked.
3. Using a No. 2 Phillips screwdriver, remove the AC power cord retaining plate by removing the screw that attaches the retaining plate to the back of the infuser.
4. Using a No. 2 Phillips screwdriver, remove the long flat-head Phillips screws from each corner of the infuser rear case assembly.
5. Separate the front case assembly from the rear case assembly.
6. Disconnect the 28 pin, 2 row connector from J8 on the power supply PWA.

   Note: Prior to disconnecting a connector, verify it is numbered. If the connector contains no identifier, number it with a permanent marker.

7. Disconnect the 16 pin connector from J9 on the power supply PWA.
8. Disconnect the 4 pin connector from J3 on the power supply PWA.
9. 3.1 Series only: Disconnect the 4 pin and 6 pin ribbon cables from J1 and J2 on the battery boost charger PWA at the back of the rear case assembly.

   Note: Ribbon cables are hard-wired to the power supply PWA. Do not damage the cables when disconnecting them.

10. Place the front assembly on the work surface.
11. Using a 5/16 inch nutdriver, remove the nut and star lockwasher securing the ground wire (green with yellow stripe) to the motor gear case.
12. Using a 5/16 inch nutdriver, remove the nut securing the ground wire (green with yellow stripe) to the front panel. The nut may be hidden by the ribbon cable connecting the bottom of the two PWAs.
13. Disconnect connectors J4, J5, J6, and J7 on the power supply PWA.
14. Re-assemble the infuser in the exact reverse order of disassembly.
15. Close and lock the security door.

To verify successful separation and re-assembly of the infuser front and rear case assemblies, perform the PVT as described in Section 5.2.
7.2.3 INJECTOR SENSOR SWITCH ADJUSTMENT

CAUTION: Use proper ESD grounding techniques when handling components. Wear an antistatic wrist strap and use an ESD-protected workstation. Store the PWA in an antistatic bag before placing it on any surface.

The injector sensor switch is adjustable on most PCA Plus Series Infusers with final assemblies 850-04250-002 through 850-04250-008. PCA Plus II Series Infusers with final assemblies 850-04250-010 and higher do not contain an injector sensor switch bracket; therefore, the injector switch is not adjustable. Some configurations of PCA Plus Series Infusers with final assembly numbers 850-04250-006 through 850-04250-008 may not contain an injector sensor switch bracket.

Recommended tools for this procedure are as follows: No. 2 Phillips screwdriver, 1/4 inch nutdriver, 3/16 inch nutdriver, 5/16 inch nutdriver, and red GLPT insulating varnish.

To adjust the injector sensor switch, refer to Figure 7-2, Injector Sensor Switch (-008 and Lower) or Figure 7-3, Injector Sensor Switch (-010 and Higher), then proceed as follows:

1. Disconnect the infuser from AC power.
2. Separate the infuser front and rear case assemblies as described in Section 7.2.2, Separating Front and Rear Case Assemblies.
3. Using a 3/16 inch nutdriver, loosen the two switch mounting screws until the switch holder can be easily moved; do not loosen the screws so much that the switch bracket rocks from side to side.

4. Position the switch bracket parallel to the surface and 1/32 inch from the infuser case; tighten the screws.

5. Apply red GLPT insulating varnish to the screw heads.

6. Re-assemble infuser in exact reverse order of disassembly.

7. Close and lock security door.

To verify successful adjustment of the injector sensor switch, perform the PVT as described in Section 5.2.

Figure 7-2. Injector Sensor Switch (-008 and Lower)
7.2.4
VIAL SENSOR SWITCH ADJUSTMENT

CAUTION: Use proper ESD grounding techniques when handling components. Wear an antistatic wrist strap and use an ESD-protected workstation. Store the PWA in an antistatic bag before placing it on any surface.

The recommended tools for this procedure are as follows: No. 2 Phillips screwdriver, 1/4 inch nutdriver, 3/16 inch nutdriver, 5/16 inch nutdriver, and red GLPT insulating varnish.

To adjust the vial sensor switch, refer to Figure 7-4, Vial Sensor Switch and Syringe Low Switch (-008 and Lower), then proceed as follows:

Note: Infusers with final assembly number -010 and higher do not have a vial sensor switch bracket (see Figure 7-5, Vial Sensor Switch (-010 and Higher). No adjustment is required on infusers with final assembly number -010 and higher.

1. Disconnect the infuser from AC power.
2. Separate the infuser front and rear case assemblies as described in Section 7.2.2, Separating Front and Rear Case Assemblies.
3. Loosen the two vial sensor switch adjustment screws with a 3/16 inch nutdriver until the switch holder can be easily moved; do not loosen the screws so much that the switch bracket rocks from side to side.
4. Place the switch bracket upright. Move the switch up slowly until it clicks off, then move the switch down slowly until it clicks on. Move the switch down further
(approximately 1/16 inch) to verify the switch detects a vial. Tighten the screws while keeping the switch bracket level.

5. Apply red GLPT insulating varnish to the screw heads.
6. Re-assemble the infuser in the exact reverse order of disassembly.
7. Close and lock the security door.

To verify successful adjustment of the vial sensor switch, perform the PVT as described in Section 5.2.

Figure 7-4. Vial Sensor Switch and Syringe Low Switch (-008 and Lower)
SYRINGE LOW SWITCH ADJUSTMENT (PCA PLUS ONLY)

CAUTION: Use proper ESD grounding techniques when handling components. Wear an antistatic wrist strap and use an ESD-protected workstation. Store the PWA in an antistatic bag before placing it on any surface.

The recommended tools for this procedure are as follows: No. 2 Phillips screwdriver, 1/4 inch nutdriver, 3/16 inch nutdriver, 5/16 inch nutdriver, syringe limit gauge, and red GLPT insulating varnish.

To adjust the syringe low switch, refer to Figure 7-4, Vial Sensor Switch and Syringe Low Switch (-008 and Lower), then proceed as follows:

1. Disconnect the infuser from AC power.
2. Separate the infuser front and rear case assemblies as described in Section 7.2.2, Separating Front and Rear Case Assemblies.
3. Install the syringe limit gauge in the cradle. Squeeze the cradle clamps and move the cradle up to the limit allowed by the gauge. Verify that the gauge dial reads approximately 0.05 inch (0.127 cm).
4. Start the infuser using any drug/concentration setting. Purge the system until the gauge dial reads approximately 0.1 inch.
5. Using a 3/16 inch nutdriver, loosen the two screws securing the syringe low switch bracket. Move the switch bracket until the switch clicks on. Tighten the screws.

6. Move the cradle up to the limit allowed by the gauge. Program the infuser for normal operation and verify that the syringe low alarm occurs when the gauge dial reads between 0.05 inch and 0.15 inch.

7. Apply red GLPT insulating varnish to the screw heads.

8. If Section 7.2.6, Syringe Empty Switch Adjustment, is not performed, re-assemble the infuser in the exact reverse order of disassembly.

9. Close and lock the security door.

To verify successful adjustment of the syringe low switch, perform the PVT as described in Section 5.2.

7.2.6 SYRINGE EMPTY SWITCH ADJUSTMENT

**CAUTION:** Use proper ESD grounding techniques when handling components. Wear an antistatic wrist strap and use an ESD-protected workstation. Store the PWA in an antistatic bag before placing it on any surface.

The recommended tools for this procedure are as follows: No. 2 Phillips screwdriver, 1/4 inch nutdriver, 3/16 inch nutdriver, 5/16 inch nutdriver, syringe limit gauge, and red GLPT insulating varnish.

To adjust the syringe empty switch, refer to Figure 7-6, Syringe Empty Switch (-008 and Lower) or Figure 7-7, Syringe Empty Switch (-010 and Higher), then proceed as follows:

1. Disconnect the infuser from AC power.

2. Separate the infuser front and rear case assemblies as described in Section 7.2.2, Separating Front and Rear Case Assemblies.

3. Install the syringe limit gauge in the cradle. Squeeze the cradle clamps and move the cradle until the dial reads approximately 0.3 inch. When the LCD screen displays PURGE THE SYSTEM NOW?, press the [YES/ENTER] touchswitch. Purge the system until the gauge reads approximately 0.39 inch. If the empty syringe alarm is activated before the system is purged, loosen the syringe empty adjustment screws and move the switch bracket up until the switch clicks off and the alarm stops.

4. Loosen the adjustment screws as necessary. Move the switch bracket down slowly until the switch clicks on. Tighten the screws.

5. Adjust the cradle until the gauge dial reads approximately 0.3 inch. Set the infuser for PCA only mode delivery. When the LCD screen displays SELECT MODE PCA ONLY?, press the [YES/ENTER] touchswitch.

6. Set a PCA dose of 5 mg.

7. Close and lock the security door.

8. When three asterisks (***') appear on the LCD screen, press the patient pendant pushbutton.

9. Verify the empty syringe alarm occurs when the dial reads between 0.39 and 0.41 inch. If the alarm does not occur within these parameters, return to Step 3 and vary the setting to less than or greater than 0.39 inch, as appropriate.

10. Tighten the screws. Apply red GLPT insulating varnish to the screw heads.
11. Re-assemble the infuser in the exact reverse order of disassembly.

12. Close and lock the security door.

To verify successful adjustment of the syringe empty switch, perform the PVT as described in Section 5.2.

Figure 7-6. Syringe Empty Switch (-008 and Lower)
7.2.7 OCCLUSION ALARM SWITCH TEST AND ADJUSTMENT

The recommended tools for this procedure are as follows: No. 0 Phillips screwdriver, No. 2 Phillips screwdriver, 1/4 inch nutdriver, 3/16 inch nutdriver, 5/16 inch nutdriver, two 9/32 inch open end wrenches, 5/64 inch Allen wrench, X-acto knife, red GLPT insulating varnish, water-filled syringe, DPM, three-way stopcock, and IV administration set.

Note: The following procedure requires a modified syringe to assure that the friction between the syringe plunger and the inside surface of the vial will not cause false occlusion pressure readings.

7.2.7.1 MODIFYING THE SYRINGE

To modify the syringe, refer to Figure 7-8, Modification of Syringe Plunger, then proceed as follows:

1. Remove the plunger from the syringe. Using an X-acto knife, cut the two top ribs off the syringe plunger, as shown in Figure 7-8. Make certain that no rough material is loose on the plunger.

2. Apply a light coating of silicone oil to the remaining rib. Replace the plunger in the syringe.
OCCLUSION ALARM SWITCH TEST

To perform the occlusion alarm switch test, refer to Figure 7-9, Occlusion Alarm Test Setup, then proceed as follows:

1. Disconnect the infuser from AC power.
2. Using a three-way stopcock, attach the DPM to the end of the patient line.
3. Install a water-filled syringe with modified plunger and an IV administration set into the infuser.
4. Purge the set until water flows out of the stopcock valve. Close the valve. When the LCD screen displays DRUG: Rx MORPHINE 1 MG/ML? YES OR NO, press [YES/ENTER].
5. Program a loading dose of 10 mg. Let the infuser operate until an occlusion alarm sounds.
6. Note the reading on the meter when the occlusion alarm occurs. If the reading is not between 14 and 19 psig (96.6 and 131.1 kPa) for -008 and lower or 12.5 and 17.5 psig (86.25 and 120.75 kPa) for -010 and higher, perform the occlusion alarm adjustment procedure in Section 7.2.7.3, Occlusion Alarm Switch and Pressure Adjustment (-008 and Lower) or 7.2.7.4, Occlusion Alarm Switch and Pressure Adjustment (-010 and Higher).

If the reading is between the specified parameters, perform the PVT as described in Section 5.2.
7.2.7.3
OCCLUSION ALARM SWITCH AND PRESSURE ADJUSTMENT (-008 AND LOWER)

CAUTION: Use proper ESD grounding techniques when handling components. Wear an antistatic wrist strap and use an ESD-protected workstation. Store the PWA in an antistatic bag before placing it on any surface.

To adjust the occlusion alarm switch and pressure, refer to Figure 7-10, Occlusion Alarm Pressure Adjustment (-008 and Lower), then proceed as follows:

1. Separate the infuser front and rear case assemblies as described in Section 7.2.2, Separating Front and Rear Case Assemblies.
2. Examine the position of the occlusion alarm switch. If the switch button is in contact with the switch actuator, proceed to Step 5. If the switch button is not making contact with the switch actuator, loosen the bracket locknut at the top of the bracket, using an 11/32 inch open end wrench.
3. Using a 9/32 inch Allen wrench, adjust the set screw so the switch button makes contact with the switch actuator, but does not open the switch.
4. Retighten the bracket locknut, using the 11/32 inch open end wrench, and apply red GLPT insulating varnish to prevent the set screw from turning.
5. Using a 5/16 inch open end wrench, loosen the lower locknut below the compression spring.

6. Connect the cables from the rear case assembly to the PWAs and slide assembly to allow the infuser to operate with the case open. Support the front case assembly to keep it upright for the adjustment procedure.

7. Install a water-filled syringe with a modified plunger into the infuser; connect an IV administration set, DPM, and stopcock. Open the stopcock valve.

8. Purge the system until water flows out of the stopcock valve. Close the valve. When the LCD screen displays DRUG: Rx MORPHINE 1 MG/ML? YES OR NO, press [YES/ENTER]. Program a loading dose of 10 mg. Let the infuser operate until an occlusion alarm sounds or the pressure reaches 19 psig (131.1 kPa).

9. If the occlusion alarm sounds before the pressure reaches 14 psig (96.6 kPa), relieve the fluid pressure by opening the stopcock valve. Tighten the upper locknut one-half turn against the compression spring. Repeat Step 7.

10. If the occlusion alarm does not sound at 19 psig (131.1 kPa), relieve the fluid pressure, loosen the upper locknut one-half turn. Repeat Step 7.

11. When the infuser occludes at between 14 and 19 psig (96.6 and 131.1 kPa), retighten the lower locknut against the upper locknut; use one 5/16 inch open end wrench to hold the upper locknut and another 5/16 inch open end wrench to tighten the lower locknut against the upper locknut.

12. Close and lock the security door.

To verify successful adjustment of the occlusion alarm switch and pressure, perform the PVT as described in Section 5.2.

Figure 7-10. Occlusion Alarm Pressure Adjustment (-008 and Lower)
7.2.7.4

**OCCLUSION ALARM SWITCH AND PRESSURE ADJUSTMENT (-010 AND HIGHER)**

To adjust the occlusion alarm switch and pressure, refer to *Figure 7-11, Occlusion Alarm Pressure Adjustment (-010 and Higher)*, then proceed as follows:

1. Separate the infuser front and rear case assemblies as described in *Section 7.2.2, Separating Front and Rear Case Assemblies*.
2. Examine the position of the occlusion alarm switch. If the switch actuator is in contact with the switch button, proceed to Step 3. If the switch actuator is not making contact with the switch button, use a No. 0 Phillips screwdriver to loosen the screw on the switch actuator. Adjust the screw until the switch actuator contacts the switch button, but does not trip the switch.
3. Connect the cables from the rear case assembly to the PWAs and slide assembly to allow the infuser to operate with the case open. Support the front case assembly to keep it upright for the adjustment procedure.
4. Install a water-filled syringe with modified plunger into the infuser; connect an IV administration set, DPM, and stopcock. Open the stopcock valve.
5. Purge the system until water flows out of the stopcock valve. Close the valve. When the LCD screen displays DRUG: Rx MORPHINE 1 MG/ML ? YES OR NO, press [YES/ENTER]. Set and initiate a loading dose of 10 mg. Allow the infuser to operate until an occlusion alarm sounds or the pressure reaches 17.5 psig (120.75 kPa).
6. If the occlusion alarm sounds before the pressure reaches 12.5 psig (86.25 kPa), relieve the fluid pressure by opening the stopcock valve. Loosen the set screw on the thumb nut and turn the thumb nut clockwise to tighten. Repeat Step 4.
7. If the occlusion alarm does not sound at or before 17.5 psig (120.75 kPa), relieve the fluid pressure by opening the stopcock valve. Loosen the set screw on the thumb nut and turn the thumb nut counterclockwise to loosen. Repeat Step 4.
8. When the infuser occludes at between 12.5 and 17.5 psig (86.25 and 120.75 kPa), retighten the set screw on the thumb nut.
9. Close and lock the security door.

To verify successful adjustment of the occlusion switch and pressure, perform the PVT as described in *Section 5.2*. 

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Section 7 REPLACEABLE PARTS AND REPAIRS
7.2.8

LCD INTENSITY ADJUSTMENT

CAUTION: Use proper ESD grounding techniques when handling components. Wear an antistatic wrist strap and use an ESD-protected workstation. Store the PWA in an antistatic bag before placing it on any surface.

The recommended tool for this procedure is a trim pot tool.

To adjust the LCD intensity, refer to Figure 7-12, CPU/Display PWA, LED, and LCD Assembly.

1. Remove the power supply PWA as described in Section 7.3.15, Power Supply PWA Replacement.
2. Remove the four spacers from the CPU/display PWA.
3. Lift the CPU/display PWA from the front enclosure. Remove the flexible circuit cable from J13 on the CPU/display PWA.
4. Position the power supply PWA and the CPU/display PWA so the resistor R19 adjustment screw is accessible. Connect P3 of the power cable to J3 on the power supply PWA. Connect the 40 pin ribbon cable to J7 on the power supply PWA.

   CAUTION: Assure the power supply PWA and the CPU/display PWA remain separated during the adjustment procedure.

5. Connect the infuser to AC power. Using the trim pot tool, adjust resistor R19 so the LCD screen can be read from a distance of approximately three feet.
6. Disconnect the infuser from AC power. Connect the flexible circuit cable to J13. Insert the CPU/display PWA in the front enclosure; replace the four spacers.
7. Replace the power supply PWA in the exact reverse order of its removal.

To verify successful adjustment of the LCD intensity, perform the PVT as described in Section 5.2.

---

7.2.9 SETTING THE REAL TIME CLOCK (PCA PLUS II)

No tools are required for this procedure.

The PCA Plus II Series Infuser real-time clock tracks time. Infuser operations require that the real-time clock be operating properly. The real-time clock must be set under the following circumstances:

- Initial installation
- Time change
- Following complete battery discharge

At start up, after successful completion of the self test, the time and date are displayed on the LCD screen. To reset the real time clock, proceed as follows:
1. Press the [NO] touchswitch to advance beyond the CLEAR? screen.
2. Press the [NO] touchswitch to advance beyond the PURGE SYSTEM NOW screen.
3. Press the [YES/ENTER] touchswitch to select the first available drug screen.
4. Press the [NO] touchswitch to advance beyond the ADMINISTER LOADING DOSE screen.
5. Press the [YES/ENTER] touchswitch to select PCA ONLY mode.
6. Press the [YES/ENTER] touchswitch to select the displayed PCA dose.
7. Press the [YES/ENTER] touchswitch to select the displayed LOCKOUT INTERVAL.
8. Press the [NO] touchswitch to advance beyond the 4 HOUR DOSE LIMIT.
9. Press the [REVIEW CHANGE] touchswitch to advance beyond the PCA SETTINGS screen.
10. Press the [NO] touchswitch to advance beyond the CHANGE? ANY SETTINGS;MODE screen.
11. Press the [YES/ENTER] touchswitch to select the CHANGE? screen to reset time and date.

   **Note:** If the fields stop flashing before the correct time and date are set, press the [NO] touchswitch to restart the flashing.

12. Press the [↑] or [↓] touchswitch to enter the correct value.
13. Press the [REVIEW CHANGE] touchswitch to advance the flashing field to the next field to be changed.
14. When all fields have been set, the LCD screen displays ACCEPT?. Press the [YES/ENTER] touchswitch to accept the values.

Setting the real time clock is a routine maintenance procedure and no verification procedure is normally required. However, if the infuser may have been damaged during this procedure, perform the PVT as described in Section 5.2.

### 7.2.10 DOOR LOCK SWITCH ADJUSTMENT

The recommended tool for this procedure is needle nose pliers.

To adjust the door lock switch, proceed as follows:

1. Disconnect the infuser from AC power.
2. Separate the infuser case assemblies as described in Section 7.2.2, Separating Front and Rear Case Assemblies.
3. Press the door lock actuator arm and listen for switch actuation.
4. Close and lock the security door and confirm the door hook and the door lock actuator hold the door closed.
5. With the security door closed, turn the key to the unlocked and locked positions while listening for switch actuation. If the door lock switch is adjusted properly, switch actuation should be heard each time the door is locked or unlocked. If the door lock switch does not require adjustment, proceed to Step 7.
6. Using needle nose pliers, carefully bend the door lock actuator or the switch actuator arm (as applicable) until proper adjustment is achieved. Be careful not to damage the switch.
7. Re-assemble the infuser in the exact reverse order of disassembly.
8. Close and lock the security door.

To verify successful adjustment of the door lock switch, perform the PVT as described in Section 5.2.

### 7.2.11 LEAD SCREW LUBRICATION

**Recommended tools for this procedure are as follows:** Electro-Wash 2000 or isopropyl alcohol, lint-free cloth or cotton swabs, grease, and a small brush.

1. Disconnect the infuser from AC power.
2. Separate the infuser case assemblies as described in Section 7.2.2, Separating Front and Rear Case Assemblies.
3. Using Electro-Wash 2000 or isopropyl alcohol and a small brush or cotton swabs, remove all old grease and residue from the lead screw.
   
   **Note:** Isopropyl alcohol may be substituted for Electro-Wash 2000; however, if using isopropyl alcohol, use extra care to assure that all residual lubricant is removed.

4. Apply a thin coating of grease and work it into the threads along the length of the shaft. Do not fill the threads. Move the slide clamp as necessary to clean and lubricate the length of the lead screw.
5. Re-assemble the infuser case assemblies in the exact reverse order of disassembly.

To verify successful lead screw lubrication, perform the PVT as described in Section 5.2.

### 7.3 REPLACEMENT PROCEDURES

This section contains safety and equipment precautions, required tools and materials, and step-by-step replacement procedures for the infuser. Unless otherwise stated, always perform the PVT after replacement procedures.

#### 7.3.1 SAFETY AND EQUIPMENT PRECAUTIONS

Before opening the front enclosure of the infuser, take all necessary precautions for working on high-voltage equipment.

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**WARNING**

UNLESS OTHERWISE INDICATED, DISCONNECT THE INFUSER FROM AC POWER BEFORE PERFORMING ANY REPLACEMENT PROCEDURE.
CAUTION: Use proper ESD grounding techniques when handling components. Wear an antistatic wrist strap and use an ESD-protected workstation. Store the PWA in an antistatic bag before placing it on any surface.

7.3.2
REQUIRED TOOLS AND MATERIALS

The following tools and materials, or equivalents, are required for the replacement procedures in this section. In addition, at the beginning of each procedure is a list of tools and materials required for that specific procedure.

- Set of Allen wrenches
- Set of nutdrivers
- 7/8 inch open end wrench
- Set of box wrenches (3/16 inch to 1/4 inch)
- Small flat blade screwdriver
- Standard flat blade screwdriver
- No. 0 Phillips screwdriver
- No. 1 Phillips screwdriver
- No. 2 Phillips screwdriver
- Electricians knife
- X-acto knife
- Wire cutters
- Wire strippers
- Soldering iron
- Solder, tin/lead, Sn63/Pb37, RMA type only
- Shrink tubing, 1/8 inch
- Small tie wraps
- Isopropyl alcohol
- Grease, Braycote 804
- Heat gun (optional)

7.3.3
SEPARATING FRONT AND REAR CASE ASSEMBLIES FOR REPLACEMENT PROCEDURES

Several of the replacement procedures in this section require separating the infuser front and rear case assemblies. Refer to Section 7.2.2, Separating Front and Rear Case Assemblies, as indicated in the following procedures.

7.3.4
BATTERY PACK COVER REPLACEMENT

The recommended tool for this procedure is a 1/4 inch nutdriver.
To replace the battery pack cover, refer to Figure 7-13, Battery Pack Replacement, then proceed as follows:

1. Disconnect the infuser from AC power.
2. Place the infuser face down on a soft surface. Using a 1/4 inch nutdriver, remove the two 6-32 hex head screws from the bottom cover.
3. Remove the battery pack cover and install the replacement battery pack cover.
4. Replace the two screws removed in Step 2.

Replacement of the battery pack cover is a routine maintenance procedure and no verification procedure is normally required. However, if the infuser may have been damaged during this procedure, perform the PVT as described in Section 5.2.

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7.3.5

BATTERY PACK REPLACEMENT

The recommended tool for this procedure is a 1/4 inch nutdriver.

To replace the battery pack, refer to Figure 7-13, Battery Pack Replacement, then proceed as follows:

1. Disconnect the infuser from AC power.
2. Place the infuser face down on a soft surface. Using a 1/4 inch nutdriver, remove the two 6-32 hex head screws from the bottom cover.
3. Remove the battery pack cover.
4. Separate the battery pack cable from the charger circuit cable. Remove the battery and dispose of it in accordance with local battery disposal practices.
5. Connect the leads of the new battery pack to the battery connector; make certain that the connection is tight.
6. Install the replacement battery pack into the housing so that the bottoms of batteries are visible; make certain that the wires are not kinked or crushed under the batteries.
7. Replace the bottom cover; make certain the battery leads are tucked inside.
8. Replace the two screws removed in Step 2.
9. *PCA Plus II only:* Reset the real time clock as described in Section 7.2.9, Setting the Real Time Clock (PCA Plus II).

To verify successful replacement of the battery pack, press the [ON] touchswitch with the infuser disconnected from AC power. Verify the front panel battery symbol illuminates.

Replacement of the battery pack is a routine maintenance procedure and no additional verification procedure is normally required. However, if the infuser may have been damaged during this procedure, perform the PVT as described in Section 5.2.

### 7.3.6 FUSE REPLACEMENT

The recommended tools for this procedure are as follows: No. 2 Phillips screwdriver and small flat blade screwdriver.

To replace the fuses, refer to *Figure 7-14, Replaceable Parts on Back of Infuser*, then proceed as follows:

1. Disconnect the infuser from AC power.
2. Place the infuser face down on a soft surface and remove the 6-32 x 1/4 inch flat head Phillips screw that attaches the AC power cord retaining plate to the infuser rear case.
3. To access the fuses, remove the AC power cord from its receptacle by grasping the plug. Do not pull on the cord.
4. Locate the plastic fuseholder directly above the AC power receptacle. Insert a small flat blade screwdriver between the locking tab at one end the fuseholder and the infuser housing. Press the tab toward the center of the fuseholder to release it. The fuseholder will move slightly outward when released from the one side.
5. Repeat Step 4 to release the other locking tab. The spring-loaded fuseholder will release and move outwards. Grasp both locking tabs and remove the fuseholder from the receptacle.
6. Remove fuses and replace with approved fuses only *(see Section 8, Specifications).* Do not use any other fuse types.
7. Insert the fuseholder into the receptacle, then press the fuseholder against the locking tabs until it clicks into position.
8. Replace the AC power cord by connecting it to the infuser AC power receptacle.
9. Re-install the AC power cord retaining plate with the screw removed in Step 2.

To verify successful replacement of the fuses, perform the PVT as described in *Section 5.2.*
7.3.7 POLE CLAMP ASSEMBLY REPLACEMENT

The recommended tool for this procedure is a No. 2 Phillips screwdriver.

To replace the pole clamp assembly, refer to Figure 7-14, Replaceable Parts on Back of Infuser, then proceed as follows:

1. Place the infuser face down on a soft surface.
2. Remove the four 5/8 inch Phillips mounting screws and associated lock washers.
3. Remove the pole clamp assembly and install the replacement pole clamp assembly using the screws and lockwashers removed in Step 2.

Replacement of the pole clamp assembly is a routine maintenance procedure and no verification procedure is normally required. However, if the infuser may have been damaged during this procedure, perform the PVT as described in Section 5.2.
7.3.8
VELCRO STRAP AND RETAINER PLATE REPLACEMENT

Velcro straps and retainer plates on the rear of the infuser secure the patient pendant and the AC power cord. This procedure details the replacement of the Velcro strap and retainer plate in both locations.

The recommended tools for this procedure are as follows: No. 2 Phillips screwdriver and X-acto knife.

To replace the Velcro strap and retainer plate, refer to Figure 7-14, Replaceable Parts on Back of Infuser, then proceed as follows:

1. Remove the two screws that attach the Velcro strap and retainer plate to the back of the infuser. Remove the retainer plate and strap. Do not discard the strap.

   **Note:** The replacement Velcro strap does not have holes for mounting screws. The holes must be cut at the time of installation.

2. Place the replacement Velcro strap, with fuzzy side down, on the work surface. Place the retainer plate on the strap in the exact location as on the old strap. Use the retainer plate as a template to mark hole locations on the strap.

3. Using an X-acto knife, cut holes in the replacement strap at the marked locations.

4. Replace the retainer plate if damaged.

5. Install the replacement strap and retainer plate using the two screws removed in Step 1.

Replacement of the Velcro strap and retainer plate is a routine maintenance procedure and no verification procedure is normally required. However, if the infuser may have been damaged during this procedure, perform the PVT as described in Section 5.2.

7.3.9
SECURITY DOOR REPLACEMENT

The recommended tool for this procedure is a No. 2 Phillips screwdriver.

To replace the security door, proceed as follows:

1. Unlock and open the security door.

2. Open the door completely (approximately 270 degrees) to reveal four round head, Phillips mounting screws.

3. Using a Phillips screwdriver, remove the four screws and remove the door.

4. Attach the replacement door using the screws removed in Step 3.

5. Close and lock the security door. Verify that the door aligns with the door latch assembly and that the door is locked.

Replacement of the security door is a routine maintenance procedure and no verification procedure is normally required. However, if the infuser may have been damaged during this procedure, perform the PVT as described in Section 5.2.
### 7.3.10 SECURITY DOOR LATCH REPLACEMENT

The recommended tool for this procedure is a No. 2 Phillips screwdriver.

To replace the security door latch, proceed as follows:

1. Unlock and open the security door.
2. From inside of the door, remove the flat head Phillips screw that attaches the latch to the door.
3. From the front of the door, peel back the nameplate label and remove the latch.
4. Install the replacement latch and re-attach it from inside the door using the attachment screw that was removed in Step 2.
5. Press the nameplate label back in place.
6. Close and lock the security door.

Replacement of the security door latch is a routine maintenance procedure and no verification procedure is normally required. However, if the infuser may have been damaged during this procedure, perform the PVT as described in Section 5.2.

### 7.3.11 PATIENT PENDANT CONTROL CABLE REPLACEMENT

To replace the patient pendant control cable, disconnect the cable from the PATIENT CONTROL jack on the rear of the infuser and connect the new cable.

Replacement of the patient pendant control cable is a routine maintenance procedure and no verification procedure is normally required. However, if the infuser may have been damaged during this procedure, perform the PVT as described in Section 5.2.

### 7.3.12 AC POWER CORD REPLACEMENT

The recommended tool for this procedure is a No. 2 Phillips screwdriver.

To replace the AC power cord, Refer to Figure 7-14, Replaceable Parts on Back of Infuser, then proceed as follows:

1. Using a No. 2 Phillips screwdriver, remove the screw that attaches the AC power cord retaining plate to the rear of the infuser.
2. Disconnect the AC power cord by grasping the plug. Do not pull on the power cord.
3. Replace the AC power cord by connecting it to the infuser AC power receptacle.
4. Re-install the power cord retaining plate with the screw removed in Step 1.
Replacement of the AC power cord is a routine maintenance procedure and no additional verification procedure is normally required. However, if the infuser may have been damaged during this procedure, perform the PVT as described in Section 5.2.

### 7.3.13 AC POWER CORD PLUG REPLACEMENT

In the event that the AC power cord plug becomes damaged, obtain a hospital-grade replacement plug. The exact procedure for replacing the plug depends upon the construction of the replacement plug. The following procedure is intended only as a guide, and not as a specific method for plug replacement.

The recommended tools for this procedure are as follows: wire cutters, screwdriver, and electricians knife.

To replace the AC power cord plug, proceed as follows:

1. Remove the damaged plug by cutting the cord near the plug.
2. Disassemble the replacement plug to access the plug terminals and to estimate the amount of insulation that needs to be removed from the AC power cord.
3. Using an electricians knife, remove the appropriate amount of outer insulation from the end of the power cord to expose the three wires in the cable. Note that the two AC wires are black and white and the ground wire is green. Using wire strippers, remove sufficient insulation (approximately 1/4 inch) from these wires to permit connection of bare connectors to the replacement plug.
4. Connect the wires to the replacement plug, as appropriate; assure that the ground wire is connected to the ground lug on the plug.
5. Re-assemble the replacement plug.
6. Conduct the electrical safety test in Section 5.2.11, Electrical Safety Test.

To verify successful replacement of the AC power cord plug, perform the PVT as described in Section 5.2.

### 7.3.14 GASKET SEAL REPLACEMENT

CAUTION: Use proper ESD grounding techniques when handling components. Wear an antistatic wrist strap and use an ESD-protected workstation. Store the PWA in an antistatic bag before placing it on any surface.

The recommended tools for this procedure are as follows: No. 2 Phillips screwdriver, 1/4 inch nutdriver, 5/16 inch nutdriver, and isopropyl alcohol.

To replace the gasket seal, proceed as follows:

1. Disconnect the infuser from AC power.
2. Separate the infuser front and rear case assemblies as described in Section 7.2.2, Separating Front and Rear Case Assemblies. Do not disconnect ribbon cables as part of this procedure.
3. Set the infuser rear case assembly upright with the security door open.
4. Remove the gasket seal and install the replacement gasket seal.
5. Wipe the replacement gasket seal with isopropyl alcohol.
6. Re-assemble the infuser case assemblies in the exact reverse order of disassembly.
7. Close and lock the security door.

To verify successful replacement of the gasket seal, perform the PVT as described in Section 5.2.

7.3.15
POWER SUPPLY PWA REPLACEMENT

CAUTION: Use proper ESD grounding techniques when handling components. Wear an antistatic wrist strap and use an ESD-protected workstation. Store the PWA in an antistatic bag before placing it on any surface.

The recommended tools for this procedure are as follows: No. 2 Phillips screwdriver, 1/4 inch nutdriver, 5/16 inch nutdriver, 1/4 inch open end wrench, and Allen wrenches.

To replace the power supply PWA, refer to Figure 7-15, Power Supply PWA and CPU/Display PWA, then proceed as follows:

1. Disconnect the infuser from AC power.
2. Separate the infuser front and rear case assemblies as described in Section 7.2.2, Separating Front and Rear Case Assemblies.
3. Using a 1/4 inch nutdriver, remove the four hex head screws securing the power supply PWA to the spacers. If necessary, hold the power supply PWA spacers with a 1/4 inch open end wrench while removing the screws.
4. Lift the power supply PWA from the spacers.
5. Install the replacement power supply PWA in the exact reverse order of removal.
6. Re-assemble the infuser case assemblies in the exact reverse order of disassembly.
7. Close and lock the security door.

To verify successful replacement of the power supply PWA, perform the PVT as described in Section 5.2.
7.3.16 CPU/DISPLAY PWA REPLACEMENT

CAUTION: Use proper ESD grounding techniques when handling components. Wear an antistatic wrist strap and use an ESD-protected workstation. Store the PWA in an antistatic bag before placing it on any surface.

The recommended tools for this procedure are as follows: No. 2 Phillips screwdriver, 1/4 inch nutdriver, 5/16 inch nutdriver, 1/4 inch open end wrench, and Allen wrenches.

To replace the CPU/display PWA, refer to Figure 7-15, Power Supply PWA and CPU Display PWA, then proceed as follows:

1. Disconnect the infuser from AC power.
2. Separate the infuser front and rear case assemblies as described in Section 7.2.2, Separating Front and Rear Case Assemblies.
3. Remove the power supply PWA as described in Section 7.3.15, Power Supply PWA Replacement.
4. Using a 1/4 inch nutdriver, remove the four CPU/display PWA spacers. Set the nylon washers aside for re-assembly.
5. Pivot the CPU/display PWA on its right edge to access the flexible circuit cable.
6. Loosen the flexible circuit cable from J13 on the CPU/display PWA.
7. Remove the 40 pin ribbon cable from J12 on the CPU/display PWA.
8. Remove the CPU/display PWA from the front panel assembly.
9. Reconnect P3 of the power cable to J3 of the power supply PWA.
10. Using a 5/16 inch nutdriver, reconnect the ground wire to the motor gear case.
11. Reconnect the infuser to AC power and press the [ON] touchswitch. Verify the LCD screen can be read from a distance of approximately three feet. If necessary, adjust the LCD intensity (see Section 7.2.8, LCD Intensity Adjustment).
12. Disconnect the infuser from AC power.
13. Re-assemble the infuser in the exact reverse order of disassembly.
14. Close and lock the security door.

To verify successful replacement of the CPU/display PWA, perform the PVT as described in Section 5.2.

7.3.17 LED REPLACEMENT

**CAUTION:** Use proper ESD grounding techniques when handling components. Wear an antistatic wrist strap and use an ESD-protected workstation. Store the PWA in an antistatic bag before placing it on any surface.

The recommended tools for this procedure are as follows: No. 2 Phillips screwdriver, 1/4 inch nutdriver, 5/16 inch nutdriver, 1/4 inch open end wrench, and Allen wrenches.

To replace any of the LEDs on the CPU/display PWA, refer to Figure 7-12, CPU/Display PWA, LED, and LCD Assembly, then proceed as follows:

1. Disconnect the infuser from AC power.
2. Separate the infuser front and rear case assemblies as described in Section 7.2.2, Separating Front and Rear Case Assemblies.
3. Remove the power supply PWA as described in Section 7.3.15, Power Supply PWA Replacement, and the CPU/display PWA as described in Section 7.3.16, CPU/Display PWA Replacement.
4. Remove the LED and install the replacement LED. The LEDs are socketed and may initially be difficult to remove; use care not to bend the connector pins.

**Note:** To assure proper alignment when replacing a seven-segment LED (DS2, DS3, DS4, DS5, or DS7), make certain the decimal point is in the lower right corner.

5. Re-install the power supply PWA and the CPU/display PWA.
6. Re-assemble the infuser case assemblies in the exact reverse order of disassembly.
7. Close and lock the security door.

To verify successful replacement of the LED, perform the PVT as described in Section 5.2.
7.3.18
LCD ASSEMBLY REPLACEMENT

CAUTION: Use proper ESD grounding techniques when handling components. Wear an antistatic wrist strap and use an ESD-protected workstation. Store the PWA in an antistatic bag before placing it on any surface.

The recommended tools for this procedure are as follows: No. 0 Phillips screwdriver, No. 2 Phillips screwdriver, 1/4 inch nutdriver, and 5/16 inch nutdriver.

To replace the LCD assembly, refer to Figure 7-12, CPU/Display PWA, LED, and LCD Assembly, then proceed as follows:

1. Disconnect the infuser from AC power.
2. Separate the infuser front and rear case assemblies as described in Section 7.2.2, Separating Front and Rear Case Assemblies.
3. Remove the power supply PWA as described in Section 7.3.15, Power Supply PWA Replacement, and the CPU/display PWA as described in Section 7.3.16, CPU/Display PWA Replacement.
4. Disconnect the LCD ribbon cable from J15 on the CPU/display PWA.
5. Pull the adhesive-backed plastic insulator from the LCD window.
6. Using a No. 0 Phillips screwdriver, remove the screws from each corner of the LCD assembly.
7. Grasp the CPU/display PWA and carefully remove the LCD assembly out of the connector J14 socket. Do not bend the connector pins.
8. Mount the adhesive-backed plastic insulator on the replacement LCD; center the insulator window on the LCD so the insulator covers the edges of the metal bracket.
9. Re-install the LCD assembly, power supply PWA, and CPU/display PWA.
10. Re-assemble the infuser case assemblies in the exact reverse order of disassembly.
11. Close and lock the security door.

To verify successful replacement of the LCD assembly, perform the PVT as described in Section 5.2.

7.3.19
40 PIN RIBBON CABLE REPLACEMENT

CAUTION: Use proper ESD grounding techniques when handling components. Wear an antistatic wrist strap and use an ESD-protected workstation. Store the PWA in an antistatic bag before placing it on any surface.

The recommended tools for this procedure are as follows: No. 2 Phillips screwdriver, 1/4 inch nutdriver, 5/16 inch nutdriver, 1/4 inch open end wrench, and Allen wrenches.

To replace the 40 pin ribbon cable, refer to Figure 7-15, Power Supply PWA and CPU/Display PWA, then proceed as follows:

1. Disconnect the infuser from AC power.
2. Separate the infuser front and rear case assemblies as described in Section 7.2.2, Separating Front and Rear Case Assemblies.
3. Remove the power supply PWA as described in *Section 7.3.15, Power Supply PWA Replacement*, and the CPU/display PWA as described in *Section 7.3.16, CPU/Display PWA Replacement*.

4. Remove the 40 pin ribbon cable from J7 on the power supply PWA and J12 on the CPU/display PWA; install the replacement 40 pin ribbon cable.

   **Note:** Connectors are keyed for pin 1.

5. Re-install the power supply PWA and the CPU/display PWA.

6. Re-assemble the infuser case assemblies in the exact reverse order of disassembly.

7. Close and lock the security door.

To verify successful replacement of the 40 pin ribbon cable, perform the PVT as described in *Section 5.2.*

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### 7.3.20

**15 CONDUCTOR DIAGNOSTIC CABLE REPLACEMENT**

**CAUTION:** Use proper ESD grounding techniques when handling components. Wear an antistatic wrist strap and use an ESD-protected workstation. Store the PWA in an antistatic bag before placing it on any surface.

The recommended tools for this procedure are as follows: No. 2 Phillips screwdriver, 1/4 inch nutdriver, 3/16 inch nutdriver, 5/16 inch nutdriver, 1/4 inch open end wrench, and Allen wrenches.

To replace the 15 conductor diagnostic cable, refer to *Figure 7-16, 15 Conductor Diagnostic Cable*, then proceed as follows:

1. Disconnect the infuser from AC power.
2. Separate the infuser front and rear case assemblies as described in *Section 7.2.2, Separating Front and Rear Case Assemblies*.
3. Disconnect the diagnostic cable from J9 on the power supply PWA.
4. Using a 3/16 inch nutdriver and a flat blade screwdriver, remove the two hex head screws and nuts that retain the diagnostic connector on the transformer/receptacle assembly.
5. Disconnect the cable from the diagnostic connector and remove it from the adhesive pad above the AC power connector next to the battery case. If the adhesive pad is worn, remove and replace it.
6. Press the connector on the replacement cable into place on the diagnostic connector; use caution not to bend the connector pins. Snap the retainer clip into position on the diagnostic connector.
7. Fold the cable 90 degrees and press it on the adhesive pads to assure clearance of the slide assembly during re-assembly *(see Figure 7-16)*.
8. Re-assemble the infuser case assemblies in the exact reverse order of disassembly.
9. Close and lock the security door.

To verify successful replacement of the 15 conductor diagnostic cable, perform the PVT as described in *Section 5.2.*
7.3.21  
CRADLE ASSEMBLY REPLACEMENT

There are two versions of the cradle assembly. Refer to Figure 7-17, Cradle Assembly Replacement, which details the differences between the cradle assemblies.

The recommended tool for this procedure is a 5/64 inch Allen wrench or a No. 2 Phillips screwdriver.

To replace the cradle assembly, refer to Figure 7-17, then proceed as follows:

1. Disconnect the infuser from AC power.
2. Open the security door.
3. Place the infuser face up on a soft surface.
4. Using a 5/64 inch Allen wrench, remove the two Allen screws that secure the cradle assembly. Remove the cradle assembly.

**Note:** Infusers -010 and higher are configured with Phillips screws securing the cradle assembly. If replacing the cradle assembly for infusers -010 or higher, use a No. 2 Phillips screwdriver to remove the screws.

5. Install the replacement cradle assembly using the screws removed in Step 4.
6. Close and lock the security door.
To verify successful replacement of the cradle assembly, perform the PVT as described in Section 5.2.

**Figure 7-17. Cradle Assembly Replacement**

### 7.3.22 VIAL SENSOR SWITCH REPLACEMENT

**CAUTION:** Use proper ESD grounding techniques when handling components. Wear an antistatic wrist strap and use an ESD-protected workstation. Store the PWA in an antistatic bag before placing it on any surface.

The recommended tools for this procedure are as follows: No. 2 Phillips screwdriver, 1/4 inch nutdriver, 3/16 inch nutdriver, 5/16 inch nutdriver, soldering iron, and solder.

To replace the vial sensor switch, refer to Figure 7-17, Cradle Assembly Replacement, and Figure 7-4, Vial Sensor Switch and Syringe Low Switch (-008 and Lower) or Figure 7-5, Vial Sensor Switch (-010 and Higher), then proceed as follows:

1. Disconnect the infuser from AC power.
2. Separate the infuser front and rear case assemblies as described in Section 7.2.2, Separating Front and Rear Case Assemblies.

3. Place the front case assembly face down on a soft surface.

4. -008 and Lower Only: Using a 3/16 inch nutdriver, remove the two vial sensor adjustment screws that attach the vial sensor switch bracket and switch to the slide assembly.

5. Using a Phillips screwdriver, remove the screw that attaches the vial sensor switch cradle clip to the slide assembly.

6. Unsolder the two wires from the vial sensor switch.

7. Solder wires to the replacement switch as follows: red wire to lower switch terminal (common) and white wire to center (N/O) terminal.

8. Replace the cradle clip removed in Step 5.

9. Install the replacement vial sensor switch. Mount the switch using the two screws removed in Step 4. -008 and Lower Only: Mount the switch bracket to the slide assembly.

10. Adjust the vial sensor switch as described in Section 7.2.4, Vial Sensor Switch Adjustment, as applicable.

11. Re-assemble the infuser case assemblies in the exact reverse order of disassembly.

12. Close and lock the security door.

To verify replacement of the vial sensor switch, perform the PVT as described in Section 5.2.

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7.3.23

SLIDE CLAMP KNOB AND LEVER REPLACEMENT (-008 AND LOWER)

CAUTION: Use proper ESD grounding techniques when handling components. Wear an antistatic wrist strap and use an ESD-protected workstation. Store the PWA in an antistatic bag before placing it on any surface.

The recommended tools for this procedure are as follows: 3/32 inch Allen wrench, No. 2 Phillips screwdriver, 1/4 inch nutdriver, and 5/16 inch nutdriver.

To replace the slide clamp knobs and levers, refer to Figure 7-18, Location of Slide Clamp Handle Allen Screws (-008 and Lower), then proceed as follows:

Note: Infusers with final assembly numbers -010 and higher do not contain slide clamp handle Allen screws.

1. Disconnect the infuser from AC power.

2. Separate the infuser front and rear case assemblies as described in Section 7.2.2, Separating Front and Rear Case Assemblies.

3. Move the cradle to the top of its travel path. Using a 3/32 inch Allen wrench, remove the two Allen screws and lock washers that secure the lever of the left slide clamp knob.

4. Remove the two Allen screws and lock washers that secure the lever of the right slide clamp knob. Press the two knobs together and remove them. Slowly release the tension of the springs between the slide clamp knobs. Note the configuration of the spring assembly (two per handle) and spring retainer sleeves.
5. Install the spring and spring retainer in the replacement knob.
6. Press the knobs together and re-install.
7. Re-assemble the infuser case assemblies in the exact reverse order of disassembly.
8. Close and lock the security door.

To verify successful replacement of the slide clamp knob and lever, perform the PVT as described in Section 5.2.

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**Figure 7-18. Location of Slide Clamp Handle Allen Screws (-008 and Lower)**

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**7.3.24 SLIDE CLAMP LEVER REPLACEMENT (-010 AND HIGHER)**

**CAUTION:** Use proper ESD grounding techniques when handling components. Wear an antistatic wrist strap and use an ESD-protected workstation. Store the PWA in an antistatic bag before placing it on any surface.

Recommended tools for this procedure are as follows: No. 2 Phillips screwdriver, 1/4 inch nutdriver, and 5/16 inch nutdriver.

To replace the slide clamp levers, refer to [Figure 7-11, Occlusion Alarm Pressure Adjustment (-010 and Higher)], then proceed as follows:

1. Disconnect the infuser from AC power.
2. Separate the infuser front and rear case assemblies as described in Section 7.2.2, Separating Front and Rear Case Assemblies.
3. Remove the cradle assembly as described Section 7.3.21, Cradle Assembly Replacement.
4. Lift the slide clamp lever off the pivot post and pull through the front of the infuser.
5. Install the replacement slide clamp lever.
6. Re-install the cradle assembly.
7. Re-assemble the infuser in the exact reverse order of disassembly.
8. Close and lock the security door.

To verify successful replacement of the slide clamp knob and lever, perform the PVT as described in Section 5.2.

### 7.3.25

**SYRINGE LOW AND SYRINGE EMPTY SWITCH REPLACEMENT**

**CAUTION:** Use proper ESD grounding techniques when handling components. Wear an antistatic wrist strap and use an ESD-protected workstation. Store the PWA in an antistatic bag before placing it on any surface.

The syringe low switch is functional only on PCA Plus Series Infusers and does not require replacement on PCA Plus II Series Infusers.

The recommended tools for this procedure are as follows: small flat blade screwdriver, No. 2 Phillips screwdriver, 1/4 inch nutdriver, 5/16 inch nutdriver, soldering iron, solder, and small tie wrap.

To replace the syringe low or syringe empty switch, refer to Figure 7-4, Vial Sensor Switch and Syringe Low Switch (-008 and Lower), Figure 7-6, Syringe Empty Switch (-008 and Lower), or Figure 7-7, Syringe Empty Switch (-010 and Higher), then proceed as follows:

1. Disconnect the infuser from AC power.
2. Separate the infuser front and rear case assemblies as described in Section 7.2.2, Separating Front and Rear Case Assemblies.
3. Place the front case assembly face down on a soft surface.
4. **-008 and lower only:** Using a 1/4 inch nutdriver or a standard screwdriver, remove the two screws from the switch mounting bracket. Remove the switch mounting bracket.
5. Using a small flat blade screwdriver and a 1/4 inch nutdriver, remove the two screws that attach the switches to the mounting bracket.
6. Cut and remove the tie wrap that ties the switch wires to the wiring harness. Unsolder the wires from the switch.
   
   **Note:** The syringe low switch is mounted next to the mounting bracket, and the syringe empty switch is mounted on top of the syringe low switch.

7. Solder the wires to the replacement switch as follows: black wire (common) to the center terminal of both switches; yellow wire to the top terminal of the syringe low switch; and green wire to the top terminal of the syringe empty switch.
8. Re-install the switches on the switch mounting bracket (-008 and lower only). Note the two flat washers between the switches.
9. Re-attach the switch mounting bracket (-008 and lower) or switch (-010 and higher) to the slide assembly.

10. Tie the switch wires to the wiring harness with a small tie wrap.

11. Adjust the syringe low switch as described in Section 7.2.5, Syringe Low Switch Adjustment (PCA Plus Only), as required. Adjust the syringe empty switch as described in Section 7.2.6, Syringe Empty Switch Adjustment, as required.

12. Re-assemble the infuser case assemblies in the exact reverse order of disassembly.

13. Close and lock the security door.

To verify successful replacement of the syringe low and syringe empty switch, perform the PVT as described in Section 5.2.

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## 7.3.26 OCCLUSION PRESSURE SWITCH REPLACEMENT

**CAUTION:** Use proper ESD grounding techniques when handling components. Wear an antistatic wrist strap and use an ESD-protected workstation. Store the PWA in an antistatic bag before placing it on any surface.

The recommended tools for this procedure are as follows: small flat blade screwdriver, No. 2 Phillips screwdriver, 1/4 inch nutdriver, 5/16 inch nutdriver, soldering iron, solder, and small tie wrap.

To replace the occlusion pressure switch, proceed as follows:

1. Disconnect the infuser from AC power.
2. Separate the infuser front and rear case assemblies as described in Section 7.2.2, Separating Front and Rear Case Assemblies.
3. Place the front case assembly face down on a soft surface.
4. Cut the tie wrap on the three wires connected to the occlusion pressure switch. Unsolder the wires from the switch terminals.
5. Using a small flat blade screwdriver, remove the two screws that attach the occlusion switch to the lower end of the slide assembly. Remove the switch.
6. Install the replacement switch using the screws removed in Step 5.
7. Solder the wires to the replacement switch as follows:
   - **-008 and Lower:** Orange wire to front terminal, black wire (common) to center terminal, and blue wire to front terminal.
   - **-010 and Higher:** Orange wire to front terminal, blue wire (common) to center terminal, and black wire to front terminal.
8. Place a small tie wrap around the switch wires.
9. Perform the occlusion alarm switch test and adjustment procedure in Section 7.2.7, Occlusion Alarm Switch Test and Adjustment, as applicable.
10. Re-assemble the infuser case assemblies in the exact reverse order of disassembly.
11. Close and lock the security door.

To verify successful replacement of the occlusion pressure switch, perform the PVT as described in Section 5.2.
7.3.27

INJECTOR SENSOR SWITCH REPLACEMENT

CAUTION: Use proper ESD grounding techniques when handling components. Wear an antistatic wrist strap and use an ESD-protected workstation. Store the PWA in an antistatic bag before placing it on any surface.

The recommended tools for this procedure are as follows: No. 2 Phillips screwdriver, 1/4 inch nutdriver, 5/16 inch nutdriver, soldering iron, and solder.

To replace the injector sensor switch, refer to Figure 7-2, Injector Sensor Switch (-008 and Lower) or Figure 7-3, Injector Sensor Switch (-010 and Higher), then proceed as follows:

1. Disconnect the infuser from AC power.
2. Separate the infuser front and rear case assemblies as described in Section 7.2.2, Separating Front and Rear Case Assemblies.
3. Place the front case assembly face down on a soft surface.
4. Using a 3/16 inch nutdriver, remove the two switch mounting screws that attach the injector sensor switch bracket (as applicable) and switch to the injector flange clamp. Remove the injector sensor switch.

   Note: Some infusers do not contain an injector sensor switch bracket.

5. Unsolder the wires from the injector sensor switch terminals.
6. Solder the wires to replacement injector sensor switch as follows: two black wires to center terminal and brown wire to outside switch terminal.
7. Mount the injector sensor switch and switch bracket (as applicable) to the injector flange clamp using the screws removed in Step 4.
8. Adjust the injector sensor switch as described in Section 7.2.3, Injector Sensor Switch Adjustment.
9. Re-assemble the infuser in the exact reverse order of disassembly.
10. Close and lock the security door.

To verify successful replacement of the injector sensor switch, perform the PVT as described in Section 5.2.

7.3.28

MOTOR ASSEMBLY REPLACEMENT (-008 AND LOWER)

CAUTION: Use proper ESD grounding techniques when handling components. Wear an antistatic wrist strap and use an ESD-protected workstation. Store the PWA in an antistatic bag before placing it on any surface.

The recommended tools for this procedure are as follows: 7/64 inch Allen wrench, No. 2 Phillips screwdriver, 1/4 inch nutdriver, and 5/16 inch nutdriver.

To replace the motor assembly, refer to Figure 7-19, Motor Assembly Replacement (-008 and Lower), then proceed as follows:
1. Disconnect the infuser from AC power.
2. Separate the infuser front and rear case assemblies as described in Section 7.2.2, Separating Front and Rear Case Assemblies.
3. Remove the slide clamp knobs/levers as described in Section 7.3.23, Slide Clamp Knob and Lever Replacement (-008 and Lower).
4. Remove the cradle assembly as described in Section 7.3.21, Cradle Assembly Replacement.
5. Remove the slide assembly as described in Section 7.3.30, Slide Assembly Replacement (-008 and Lower).
6. Remove the two 7/64 inch Allen screws that attach the motor gear case to the slide assembly. Remove motor assembly.

**Note:** A coupling tube and a coupling clip come off with the motor assembly.

7. Place the coupling clip inside the coupling tube and install both parts on the motor gear case shaft.
8. Line up the flats of the lead screw and the motor gear case shafts. Push the motor assembly onto the lead screw shaft. If necessary, move the motor assembly from side to side until the coupling pin engages the flats of the lead screw and motor gear case shafts. When the motor assembly is flush against the bottom of the slide, align the motor gear case holes with the holes tapped in the bottom of the slide.
9. Re-install the two 7/64 inch Allen screws removed in Step 6.
10. Re-assemble the infuser in the exact reverse order of disassembly.
11. Close and lock the security door.

To verify successful replacement of the motor assembly, perform the PVT as described in Section 5.2.

![Figure 7-19. Motor Assembly Replacement (-008 and Lower)](image-url)
CAUTION: Use proper ESD grounding techniques when handling components. Wear an antistatic wrist strap and use an ESD-protected workstation. Store the PWA in an antistatic bag before placing it on any surface.

The recommended tools for this procedure are as follows: 7/64 inch Allen wrench, No. 2 Phillips screwdriver, 1/4 inch nutdriver, and 5/16 inch nutdriver.

To replace the motor assembly, refer to Figure 7-20, Motor Assembly Replacement (-010 and Higher), then proceed as follows:

1. Disconnect the infuser from AC power.
2. Separate the infuser front and rear case assemblies as described in Section 7.2.2, Separating Front and Rear Case Assemblies.
3. Remove the slide clamp levers as described in Section 7.3.24, Slide Clamp Lever Replacement (-010 and Higher).
4. Remove the cradle assembly as described in Section 7.3.21, Cradle Assembly Replacement.
5. Disconnect the connectors from the power supply PWA and CPU/display PWA.
6. Using a 5/16 inch nutdriver, remove the three hex-head screws that mount the slide assembly to the front case (see Figure 7-21, Location of Slide Assembly Mounting Bolts).
7. Lift the slide assembly from the front case and place it on its right side on a soft surface. Do not damage the vial switch plunger that protrudes from the slide assembly; do not damage the injector switch wires that remain attached.
8. Remove the two 6-32 Phillips screws and nuts that hold the motor assembly to the motor support.
9. Remove the motor assembly and coupler.
10. Install the replacement motor assembly in the slide assembly with the coupler. Replace the coupler at this point, if necessary.
11. Re-assemble the infuser in the exact reverse order of disassembly.
12. Close and lock the security door.

To verify successful replacement of the motor assembly, perform the PVT as described in Section 5.2.
Figure 7-20. Motor Assembly Replacement (-010 and Higher)

Figure 7-21. Location of Slide Assembly Mounting Bolts
7.3.30
SLIDE ASSEMBLY REPLACEMENT (-008 AND LOWER)

CAUTION: Use proper ESD grounding techniques when handling components. Wear an antistatic wrist strap and use an ESD-protected workstation. Store the PWA in an antistatic bag before placing it on any surface.

The recommended tools for this procedure are as follows: 9/64 inch Allen wrench, No. 2 Phillips screwdriver, 1/4 inch nutdriver, 3/16 inch nutdriver, 5/16 inch nutdriver, and grease.

To replace the slide assembly, refer to Figure 7-21, Location of Slide Assembly Mounting Bolts, then proceed as follows:

1. Disconnect the infuser from AC power.
2. Separate the infuser front and rear case assemblies as described in Section 7.2.2, Separating Front and Rear Case Assemblies.
3. Remove the following using the procedures described in the corresponding sections:
   - Slide clamp knobs and levers - Section 7.3.23
   - Cradle assembly - Section 7.3.21
   - Vial sensor switch - Section 7.3.22
   - Syringe low and syringe empty switches - Section 7.3.25
   - Occlusion pressure switch - Section 7.3.26
4. Using a 5/16 inch nutdriver, remove the three hex head bolts that mount the slide assembly to the front case.
5. Lift the slide assembly from the front case and place it on its right side on a soft surface; use care not to damage the vial switch that protrudes from the slide assembly.
6. Remove the injector sensor switch as described in Section 7.3.27, Injector Sensor Switch Replacement.
7. Remove the plastic cable clamp from the slide assembly; remove the switch wiring harness.
8. Remove the motor assembly from the slide assembly as described in Section 7.3.28, Motor Assembly Replacement (-008 and lower).
9. Mount the motor assembly on the replacement slide assembly.
10. Replace the injector sensor switch removed in Step 6.
11. Using the three hex head bolts removed in Step 4, mount the replacement slide assembly on the front case. Lubricate the lead screw with grease.
12. Re-install the occlusion pressure switch, syringe low switch, syringe empty switch, and vial sensor switch in the exact reverse order of disassembly.
13. Re-attach the plastic cable clamp removed in Step 7 on the replacement slide assembly.
14. Re-install the cradle assembly, and the slide clamp knobs and levers in the exact reverse order of disassembly.
15. Adjust the following switches using the procedures described in the corresponding sections:
   - Injector sensor switch - Section 7.2.3
   - Vial sensor switch - Section 7.2.4
16. Re-assemble the infuser case assemblies in the exact reverse order of disassembly.
17. Close and lock the security door.

To verify successful replacement of the slide assembly, perform the PVT as described in Section 5.2.

7.3.31
SLIDE ASSEMBLY REPLACEMENT (-010 AND HIGHER)

CAUTION: Use proper ESD grounding techniques when handling components. Wear an antistatic wrist strap and use an ESD-protected workstation. Store the PWA in an antistatic bag before placing it on any surface.

The recommended tools for this procedure are as follows: 9/64 inch Allen wrench, No. 2 Phillips screwdriver, 1/4 inch nutdriver, 3/16 inch nutdriver, 5/16 inch nutdriver, and grease.

To replace the slide assembly, refer to Figure 7-21, Location of Slide Assembly Mounting Bolts, then proceed as follows:

1. Disconnect the infuser from AC power.
2. Separate the infuser front and rear case assemblies as described in Section 7.2.2, Separating Front and Rear Case Assemblies.
3. Remove the cradle assembly as described in Section 7.3.21, Cradle Assembly Replacement.
4. Remove the slide clamp levers as described in Section 7.3.24, Slide Clamp Lever Replacement (-010 and Higher).
5. Using a No. 1 Phillips screwdriver and a 1/8 inch nutdriver, remove the two wire harnesses that contain the occlusion pressure switch, vial sensor switch, injector sensor switch, and empty vial/syringe switch. Mark P/J numbers on the connectors prior to removing, if necessary. Cut the tie wraps to remove the harnesses, if necessary.
6. Using a 5/16 inch nutdriver, remove the three hex-head screws that mount the slide assembly to the front case.
7. Lift the slide assembly out of the front case and place it on its right side on a soft surface; use care not to damage the vial switch plunger that protrudes from the slide assembly.
8. Install the replacement slide assembly and replace the screws removed in Step 6. Lubricate the lead screw with grease.
9. Re-install the wire harnesses in the exact reverse order of disassembly.
10. Reconnect the ribbon cables to the power supply PWA.
11. Replace the slide clamp knobs/levers removed in Step 4.
12. Replace the cradle assembly removed in Step 3.
13. Adjust the following switches using the procedures described in the corresponding sections:
   - Injector sensor switch - Section 7.2.3
Vial sensor switch - Section 7.2.4
- Syringe low switch (as applicable) - Section 7.2.5
- Syringe empty switch - Section 7.2.6
- Occlusion pressure switch - Section 7.2.7.4

14. Re-assemble the infuser case assemblies in the exact reverse order of disassembly.
15. Close and lock the security door.

To verify successful replacement of the slide assembly, perform the PVT as described in Section 5.2.

### 7.3.32
**INJECTOR FLANGE CLAMP REPLACEMENT**

**CAUTION:** Use proper ESD grounding techniques when handling components. Wear an antistatic wrist strap and use an ESD-protected workstation. Store the PWA in an antistatic bag before placing it on any surface.

Two configurations of injector flange clamps are used, depending on the final assembly number of the infuser. This procedure may be followed for both configurations.

The recommended tools for this procedure are as follows: No. 2 Phillips screwdriver, 1/4 inch nutdriver, 3/16 inch nutdriver, and 5/16 inch nutdriver.

To replace the injector flange clamp, refer to Figure 7-17, Cradle Assembly Replacement, then proceed as follows:

1. Disconnect the infuser from AC power.
2. Separate the infuser front and rear case assemblies as described in Section 7.2.2, Separating Front and Rear Case Assemblies.
3. Place the front case assembly face down on a soft surface.
4. Remove the slide assembly as described in Section 7.3.30, Slide Assembly Replacement (-008 and Lower), or Section 7.3.31, Slide Assembly Replacement (-010 and Higher).
5. Using a 3/16 inch nutdriver, remove the two switch mounting screws that attach the injector sensor switch bracket (as applicable) and switch to the slide assembly. Remove the injector sensor switch.

**Note:** Some infusers do not contain an injector sensor switch bracket.

6. Using a 1/4 inch nutdriver, remove the screws that attach the injector flange clamp to the front case.
7. Mount the replacement injector flange clamp to the front case using the screws removed in Step 6.
8. Mount the injector sensor switch and switch bracket (as applicable) to the injector flange clamp using the screws removed in Step 5.
9. Re-install the slide assembly as described in Section 7.3.30, or Section 7.3.31.
10. Adjust the injector sensor switch as described in Section 7.2.3, Injector Sensor Switch Adjustment.
11. Re-assemble the infuser case assemblies in the exact reverse order of disassembly.
12. Close and lock the security door.

To verify successful replacement of the injector flange clamp, perform the PVT as described in Section 5.2.

7.3.33

PRINTER/DATAWAY CONNECTOR AND CONTROL PANEL HARNESS REPLACEMENT

CAUTION: Use proper ESD grounding techniques when handling components. Wear an antistatic wrist strap and use an ESD-protected workstation. Store the PWA in an antistatic bag before placing it on any surface.

The recommended tools for this procedure are as follows: No. 2 Phillips screwdriver, 3/16 inch nutdriver, 1/4 inch nutdriver, 5/16 inch nutdriver, 5/16 inch box wrench, 1/2 inch box wrench, and small flat blade screwdriver.

To replace the printer/Dataway connector and control panel harness, refer to Figure 7-22, Printer/Dataway Connector, Control Panel Harness, and Piezoelectric Alarm Replacement, then proceed as follows:

1. Disconnect the infuser from AC power.
2. Separate the infuser front and rear case assemblies as described in Section 7.2.2, Separating Front and Rear Case Assemblies.
3. Place the infuser rear case assembly on a soft surface with interior facing up.
4. Disconnect the harness connector from J8 on the power supply PWA.
5. Pull the adhesive-backed tie wrap clamp from the rear case. Cut and remove the tie wrap that attaches the door lock switch wires to the rear case.
6. Using a small flat blade screwdriver and a 1/4 nutdriver, remove the two screws that attach the door lock switch to the door latch.
7. To ease disassembly and re-assembly, remove the eight Phillips screws that attach the control panel to the rear case. Remove the control panel.
8. Using a small flat blade screwdriver and a 3/16 inch nutdriver, remove the two screws that attach the printer/Dataway connector (J10) to the control panel.
9. Using a 5/16 inch nutdriver, remove the nut that attaches the green wire to the control panel.
10. Using a 3/16 inch nutdriver, remove the screws that attach the piezoelectric alarm to the control panel. The longer of the two screws is used in the upper mounting hole to accommodate the plastic cable clip.
11. Using a 5/16 inch box wrench, remove the retainer nut that attaches the audible level switch to the control panel.
12. Using a 1/2 inch box wrench, remove the retainer nut that attaches the patient pendant jack to the control panel.
13. Remove the plastic cable clip that attaches the harness to the piezoelectric alarm mounting screw; place the clip on the replacement harness in the same location.
14. Using a 1/2 inch box wrench, attach the patient pendant jack on the replacement harness to the control panel.
15. Using a 5/16 inch box wrench, attach the audible level switch on the replacement harness to the control panel.

16. Using a 5/16 inch box wrench, attach the piezoelectric alarm on the replacement harness to the control panel. Attach the harness clamp to the alarm with the longer mounting screw in the upper position.

17. Using a small flat blade screwdriver and a 3/16 inch nutdriver, attach the printer/Dataway connector (J10) on the replacement harness to the control panel.

18. Remount the control panel on the rear case with the eight Phillips screws removed in Step 7.

19. Using a 5/16 inch nutdriver, attach the green ground wire in the replacement harness to the control panel.

20. Replace the adhesive-backed tie wrap clamp removed in Step 5; use care to place it in the same location. Using the tie wrap, tie the replacement harness to the clamp.

21. Connect connector P8 on the replacement harness to J8 on the power supply PWA.

22. Re-assemble the infuser case assemblies in the exact reverse order of disassembly.

23. Close and lock the security door.

To verify successful replacement of the printer/Dataway connector and control panel harness, perform the PVT as described in Section 5.2.
7.3.34

PIEZOELECTRIC ALARM REPLACEMENT

CAUTION: Use proper ESD grounding techniques when handling components. Wear an antistatic wrist strap and use an ESD-protected workstation. Store the PWA in an antistatic bag before placing it on any surface.

This procedure applies to infusers with final assembly numbers 850-04250-002 through -007. Infusers with final assembly number 850-04250-008 and higher require replacement of the Interconnect PWA if the piezoelectric alarm is defective (see Section 7.3.37, Interconnect PWA Replacement).

The recommended tools for this procedure are as follows: No. 2 Phillips screwdriver, 1/4 inch nutdriver, 5/16 inch nutdriver, wire cutters, soldering iron, solder, 1/8 inch shrink tubing, and small tie wrap.

To replace the piezoelectric alarm, refer to Figure 7-22, Printer/Dataway Connector, Control Panel Harness, and Piezoelectric Alarm Replacement, then proceed as follows:

1. Disconnect the infuser from AC power.
2. Separate the infuser front and rear case assemblies as described in Section 7.2.2, Separating Front and Rear Case Assemblies.
3. Using a 3/16 inch nutdriver, remove the screws that attach the piezoelectric alarm to the control panel. The longer of the two screws is used in the upper mounting hole to accommodate the plastic cable clip.
4. Cut the cable tie nearest to the piezoelectric alarm on the wire harness. Note that the piezoelectric alarm has a black wire and a red wire. The red wire connects to the gray wire on the wire harness; the black wire connects to the white wire on the wire harness. Remove the shrink wrap from the solder joints and unsolder the wires.
5. Remove the piezoelectric alarm.
6. Slip one inch of 1/8 inch shrink tubing over the gray and white wires on the wire harness.
7. Solder the red wire of the replacement piezoelectric alarm to the gray wire on the harness. Solder the black wire of the replacement piezoelectric alarm to the white wire on the harness.
8. Slip the shrink tubing over the solder joints and shrink into place.
9. Mount the piezoelectric alarm to the control panel with the screws removed in Step 3. Place the clamp around the harness wires routed to the rear of the printer connector on the control panel. Attach the plastic cable clip with the upper screw.
10. Install a new tie wrap on the wire harness; use care to place it in the same location.
11. Re-assemble the infuser case assemblies in the exact reverse order of disassembly.
12. Close and lock the security door.

To verify successful replacement of the piezoelectric alarm, perform the PVT as described in Section 5.2.
7.3.35  
PATIENT PENDANT JACK REPLACEMENT

CAUTION: Use proper ESD grounding techniques when handling components. Wear an antistatic wrist strap and use an ESD-protected workstation. Store the PWA in an antistatic bag before placing it on any surface.

This procedure applies to infusers with final assembly numbers 850-04250-002 through -007. Infusers with final assembly number 850-04250-008 and higher require replacement of the Interconnect PWA if the patient pendant jack is defective (see Section 7.3.37, Interconnect PWA Replacement).

The recommended tools for this procedure are as follows: No.2 Phillips screwdriver, 1/4 inch nutdriver, 5/16 inch nutdriver, 1/2 inch box wrench, soldering iron, and solder.

To replace the patient pendant jack, refer to Figure 7-22, Printer/Dataway Connector, Control Panel Harness, and Piezoelectric Alarm Replacement, then proceed as follows:

1. Disconnect the infuser from AC power.
2. Separate the infuser front and rear case assemblies as described in Section 7.2.2, Separating Front and Rear Case Assemblies.
3. Place the infuser rear case assembly on a soft surface with the interior facing up. Remove the adhesive-backed tie wrap clamp from the rear case.
4. Remove the eight Phillips screws that attach the control panel to the rear case. Remove the control panel.
5. Unsolder the harness wires from the patient pendant jack.
6. Using a 1/2 inch box wrench, remove the nut that attaches the patient pendant jack to the control panel. Remove the patient pendant jack.
7. Install the replacement patient pendant jack on the control panel. Using a 1/2 inch box wrench, secure the jack with the retaining nut.
8. Solder the wires from the harness to the patient pendant jack as follows: white wire with red stripe to the ring terminal, white wire with green stripe to the tip terminal.
9. Using a 1/2 inch box wrench, remove the retainer nut that attaches the patient pendant jack to the control panel.
10. Remount the control panel on the rear case with the eight Phillips screws removed in Step 4.
11. Replace the adhesive-backed tie wrap clamp on the inside of the rear case; use care to place it in the same location.
12. Re-assemble the infuser case assemblies in the exact reverse order of disassembly.
13. Close and lock the security door.

To verify successful replacement of the patient pendant jack, perform the PVT as described in Section 5.2.
7.3.36

AUDIBLE LEVEL SWITCH REPLACEMENT

CAUTION: Use proper ESD grounding techniques when handling components. Wear an antistatic wrist strap and use an ESD-protected workstation. Store the PWA in an antistatic bag before placing it on any surface.

This procedure applies to infusers with final assembly numbers 850-04250-002 through -007. Infusers with final assembly number 850-04250-008 and higher require replacement of the Interconnect PWA if the audible level switch is defective (see Section 7.3.37, Interconnect PWA Replacement).

The recommended tools for this procedure are as follows: No. 2 Phillips screwdriver, 1/4 inch nutdriver, 5/16 inch nutdriver, soldering iron, and solder.

To replace the audible level switch, proceed as follows:

1. Disconnect the infuser from AC power.
2. Separate the infuser front and rear case assemblies as described in Section 7.2.2, Separating Front and Rear Case Assemblies.
3. Place the infuser rear case assembly on a soft surface with the interior facing up. Remove the adhesive-backed tie wrap clamp from the rear case.
4. Remove the eight Phillips screws that attach the control panel to the rear case. Remove the control panel.
5. Unsolder the harness wires from the audible level switch terminals.
6. Using a 5/16 inch nutdriver, remove the nut that attaches the audible level switch to the control panel.
7. Install the replacement audible level switch in the control panel. Using a 1/2 inch box wrench, secure the switch with the retaining nut.
8. Solder the wires from the harness to the audible level switch as follows: white wire with orange stripe to the upper terminal, white wire with black stripe to the center terminal, white wire with brown stripe to the lower terminal.
9. Remount the control panel on the rear case with the eight Phillips screws removed in Step 4.
10. Replace the adhesive-backed tie wrap clamp on the inside of the rear case; use care to place it in the same location.
11. Re-assemble the infuser case assemblies in the exact reverse order of disassembly.
12. Close and lock the security door.

To verify successful replacement of the audible level switch, perform the PVT as described in Section 5.2.
7.3.37
INTERCONNECT PWA REPLACEMENT

CAUTION: Use proper ESD grounding techniques when handling components. Wear an antistatic wrist strap and use an ESD-protected workstation. Store the PWA in an antistatic bag before placing it on any surface.

This procedure applies to infusers with final assembly number 850-04250-008 and higher.

The recommended tools for this procedure are as follows: No. 2 Phillips screwdriver, 1/4 inch nutdriver, 5/16 inch nutdriver, wire cutters, wire strippers, soldering iron, solder, and approximately one inch of 1/8 inch shrink tubing.

To replace the interconnect PWA, refer to Figure 7-23, Interconnect PWA and Battery Boost Charger PWA Replacement, then proceed as follows:

1. Disconnect the infuser from AC power.
2. Separate the infuser front and rear case assemblies as described in Section 7.2.2, Separating Front and Rear Case Assemblies.
3. Disconnect the interconnect PWA cable from J10, the interconnect PWA, and J8 on the power supply PWA.
4. Using a 5/16 inch nutdriver, remove the nut that attaches the green ground wire to the control panel.
5. Using wire cutters, cut the two wires that connect the door lock switch to the interconnect PWA. Cut the wires between the interconnect PWA and the cable clamps mounted to the rear case.
6. Remove the eight Phillips screws that attach the control panel and interconnect PWA to the rear case. Remove the control panel and interconnect PWA.
7. Install the replacement control panel and interconnect PWA using the eight Phillips screws removed in Step 6.
8. Using wire strippers, strip approximately 1/2 inch of insulation from the two wires that connect the door lock switch to the interconnect PWA.
9. Slip one inch of 1/8 inch shrink tubing over the door lock switch wires. Solder the wires to the corresponding wires on the interconnect PWA. Slip the shrink tubing over the solder joint and shrink in place.
10. Attach the green ground wire removed in Step 4. Connect the ribbon cable disconnected in Step 3.
11. Re-assemble the infuser case assemblies in the exact reverse order of disassembly.
12. Close and lock the security door.

To verify successful replacement of the interconnect PWA, perform the PVT as described in Section 5.2.
7.3.38
INTERCONNECT PWA CABLE REPLACEMENT

CAUTION: Use proper ESD grounding techniques when handling components. Wear an antistatic wrist strap and use an ESD-protected workstation. Store the PWA in an antistatic bag before placing it on any surface.

This procedure applies to infusers with final assembly number 850-04250-008 and higher.

The recommended tools for this procedure are as follows: No. 2 Phillips screwdriver, 1/4 inch nutdriver, and 5/16 inch nutdriver.

To replace the interconnect PWA cable, refer to Figure 7-23, Interconnect PWA and Battery Boost Charger PWA Replacement, then proceed as follows:

1. Disconnect the infuser from AC power.
2. Separate the infuser front and rear case assemblies as described in Section 7.2.2, Separating Front and Rear Case Assemblies.
3. Disconnect the interconnect PWA cable from connector J10 on the interconnect PWA and from J8 on the power supply PWA.
4. Connect the replacement interconnect PWA cable to connectors J8 and J10.
5. Re-assemble the infuser case assemblies in the exact reverse order of disassembly.
6. Close and lock the security door.
To verify successful replacement of the interconnect PWA cable, perform the PVT as described in Section 5.2.

7.3.39
BATTERY BOOST CHARGER PWA REPLACEMENT

**CAUTION:** Use proper ESD grounding techniques when handling components. Wear an antistatic wrist strap and use an ESD-protected workstation. Store the PWA in an antistatic bag before placing it on any surface.

The recommended tools for this procedure are as follows: No. 1 Phillips screwdriver, No. 2 Phillips screwdriver, 1/4 inch nutdriver, and 5/16 inch nutdriver.

To replace the battery boost charger PWA, refer to Figure 7-23, Interconnect PWA and Battery Boost Charger PWA Replacement, then proceed as follows:

1. Disconnect the infuser from AC power.
2. Separate the infuser front and rear case assemblies as described in Section 7.2.2, Separating Front and Rear Case Assemblies.
3. Disconnect P1 and P2 from J1 and J2 on the battery boost charger PWA.
4. Using a No. 1 Phillips screwdriver, remove the two 6-32 screws that hold the battery boost charger PWA on the control panel assembly. Remove the battery boost charger PWA.
5. Mount the replacement battery boost charger PWA on the control panel assembly with the screws removed in Step 4.
6. Re-assemble the infuser case assemblies in the exact reverse order of disassembly.
7. Close and lock the security door.

To verify successful replacement of the battery boost charger PWA, perform the PVT as described in Section 5.2.

7.3.40
FRONT PANEL REPLACEMENT

**CAUTION:** Use proper ESD grounding techniques when handling components. Wear an antistatic wrist strap and use an ESD-protected workstation. Store the PWA in an antistatic bag before placing it on any surface.

The recommended tools for this procedure are as follows: No. 2 Phillips screwdriver, 1/4 inch nutdriver, 5/16 inch nutdriver, 1/4 inch open end wrench, Allen wrenches, X-acto knife, and isopropyl alcohol.

To replace the front panel, refer to Figure 7-24, Front Panel Replacement, then proceed as follows:

1. Disconnect the infuser from AC power.
2. Separate the infuser front and rear case assemblies as described in Section 7.2.2, Separating Front and Rear Case Assemblies.
3. Remove the power supply PWA as described in Section 7.3.15, Power Supply PWA Replacement, and the CPU/display PWA as described in Section 7.3.16, CPU/Display PWA Replacement.

4. Using a 1/4 inch nutdriver, remove the four hex head screws that attach the front panel assembly to the front case. Remove the front panel assembly.

5. Using an X-acto knife, pry the front panel from the sub panel. Remove the old adhesive from the front panel with isopropyl alcohol.

6. Remove the protective backing from the replacement front panel, then place the replacement front panel over the sub panel. Make certain that the front panel is properly seated on the sub panel, then press it into place. Make certain that no air bubbles are trapped between the two surfaces and that the surfaces bond securely.

7. Re-install the power supply PWA and the CPU/display PWA.

8. Re-assemble the infuser case assemblies in the exact reverse order of disassembly.

9. Close and lock the security door.

To verify successful replacement of the front panel, perform the PVT as described in Section 5.2.

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**Figure 7-24. Front Panel Replacement**

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**7.3.41**

**DOOR LOCK SWITCH REPLACEMENT**

**CAUTION:** Use proper ESD grounding techniques when handling components. Wear an antistatic wrist strap and use an ESD-protected workstation. Store the PWA in an antistatic bag before placing it on any surface.

The recommended tools for this procedure are as follows: No. 2 Phillips screwdriver, small flat blade screwdriver, 1/4 inch nutdriver, soldering iron, and solder.
To replace the door lock switch, refer to Figure 7-25, Door Latch Assembly (Type A), or Figure 7-26, Door Latch Assembly (Type B), then proceed as follows:

1. Disconnect the infuser from AC power.
2. Separate the infuser front and rear case assemblies as described in Section 7.2.2, Separating Front and Rear Case Assemblies.
3. Using a small flat blade screwdriver and a 1/4 inch nutdriver, remove the door lock switch from the door latch assembly.

   **Note:** A small flat blade screwdriver is required only for infusers with final assembly number 850-04250-008.

4. Unsolder the wires from the door lock switch.
5. Mount the replacement door lock switch on the door latch assembly.
6. Solder the wires to the terminal of the replacement door lock switch as follows: white wire with green and brown stripes to the center terminal, white wire with blue and brown stripes to the rear terminal.
7. Re-assemble the infuser case assemblies in the exact reverse order of disassembly.
8. Close and lock the security door.

To verify successful replacement of the door lock switch, perform the PVT as described in Section 5.2.

**Figure 7-25. Door Latch Assembly (Type A)**
7.3.42 DOOR LOCK ASSEMBLY REPLACEMENT

CAUTION: Use proper ESD grounding techniques when handling components. Wear an antistatic wrist strap and use an ESD-protected workstation. Store the PWA in an antistatic bag before placing it on any surface.

The recommended tools for this procedure are as follows: No. 2 Phillips screwdriver, small flat blade screwdriver, 1/4 inch nutdriver, 5/16 inch nutdriver, 7/16 inch box wrench, and 7/8 inch open end wrench.

To replace the door lock assembly, refer to Figure 7-27, Door Lock Assembly, then proceed as follows:

1. Disconnect the infuser from AC power.
2. Separate the infuser front and rear case assemblies as described in Section 7.2.2, Separating Front and Rear Case Assemblies.
3. Remove the door lock switch from the door lock assembly as described in Section 7.3.41, Door Lock Switch Replacement.
4. Using a 7/16 inch box wrench, remove the nut from the end of the door lock cylinder. Remove the lock washer, the door latch pawl, and the locking washer.
5. Using a 7/8 inch open end wrench, remove the retainer nut that attaches the lock cylinder to the door lock assembly. Remove the lock cylinder, then remove the door lock assembly from the rear case.
6. Install the new door lock cylinder into the double-D hole in the cover and door lock assembly. Confirm the keyway slot is in the upward (12 o’clock) position.

Figure 7-26. Door Latch Assembly (Type B)
7. Assemble the retainer nut on the door lock cylinder and tighten it against the door lock assembly. Replace the locking washer, latch pawl, lock washer, and nut on the door lock cylinder. Confirm the latch pawl hook is in the upward position.

8. Attach the replacement door lock assembly to the rear case in the exact reverse order of disassembly.

9. Re-assemble the infuser case assemblies in the exact reverse order of disassembly.

10. Close and lock the security door.

To verify successful replacement of the door lock assembly, perform the PVT as described in Section 5.2.

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7.3.43

CHANGING THE KEY NUMBER

CAUTION: Use proper ESD grounding techniques when handling components. Wear an antistatic wrist strap and use an ESD-protected workstation. Store the PWA in an antistatic bag before placing it on any surface.

Note: This procedure is applicable only to locks with the numbers 1 through 8 engraved on the lock face.

Recommended tools for this procedure are as follows: master key (black), and keys numbered 2 through 8.

To change a key number, proceed as follows:

1. Remove the door lock cylinder as described in Section 7.3.42, Door Lock Assembly Replacement.

2. Completely insert the master key (black) into the keylock assembly at position 1. Confirm the longer wing of the master key is positioned upward (12 o’clock).

3. Turn the master key clockwise to the new key number position.

4. Remove the master key from the keylock assembly.

5. Completely insert the new position combination key into the keylock assembly. Confirm an audible click sounds.
6. To confirm the keylock is set correctly, insert the key and lock, then unlock the security door.

7. If the key does not unlock the security door, reinsert the master key into the keylock with the longer wing positioned at the new key number position.

8. Turn the master key counterclockwise to position 1. Repeat Steps 3 through 5.

9. Tag the PCA infuser with the new key number.

10. Reassemble the infuser in the exact reverse order of disassembly.

Perform the PVT as described in Section 5.2.

7.3.44

DOOR LATCH ASSEMBLY REPLACEMENT

CAUTION: Use proper ESD grounding techniques when handling components. Wear an antistatic wrist strap and use an ESD-protected workstation. Store the PWA in an antistatic bag before placing it on any surface.

The recommended tools for this procedure are as follows: No. 2 Phillips screwdriver, small flat blade screwdriver, 1/8 inch nutdriver, 1/4 inch nutdriver, 5/16 inch nutdriver, 7/16 inch box wrench, and 7/8 inch open end wrench.

To replace the door latch assembly, refer to Figure 7-25, Door Latch Assembly (Type A), and Figure 7-28, Door Latch Torsion Spring Placement (Type A), or Figure 7-26, Door Latch Assembly (Type B), then proceed as follows:

1. Disconnect the infuser from AC power.

2. Separate the infuser front and rear case assemblies as described in Section 7.2.2, Separating Front and Rear Case Assemblies.

3. Using a small flat blade screwdriver and a 1/4 inch nutdriver, remove the door lock switch from the door latch assembly.

   **Note:** A small flat blade screwdriver is required only for infusers with final assembly number 850-04250-008.

4. Make sure the key is in the upward (12 o'clock) position, then remove the two 1/4 inch hex head screws that attach the door latch and door latch spacer (as applicable) to the support plate. Remove the door latch assembly.

5. For type B locks only, remove the extension spring from the door lock bracket and latch (see Figure 7-26).

6. Remove the retaining ring from the pivot pin. Remove the pivot pin, torsion spring (type A locks only), and the door lock actuator.

7. Reassemble the door latch assembly in the exact reverse order of disassembly; for type A locks make certain the torsion spring is compressed, as shown in Figure 7-28, before replacing the pivot pin.

8. Re-install the door lock switch and reassemble the infuser in the exact reverse order of disassembly.

9. Close and lock the security door.

To verify successful installation of the door latch assembly, perform the PVT as described in Section 5.2.
7.3.45

REAR CASE ASSEMBLY REPLACEMENT

CAUTION: Use proper ESD grounding techniques when handling components. Wear an antistatic wrist strap and use an ESD-protected workstation. Store the PWA in an antistatic bag before placing it on any surface.

Note: The infuser serial number is required when ordering a new rear case. The serial number is located on the back of the infuser. Abbott Laboratories will place a duplicate serial number label on the replacement rear case.

The recommended tools for this procedure are as follows: No. 2 Phillips screwdriver, 3/16 inch nutdriver, 1/4 inch nutdriver, 5/16 inch nutdriver, 5/16 inch box wrench, 1/2 inch box wrench, and small flat blade screwdriver.

To replace the rear case, proceed as follows:

1. Disconnect the infuser from AC power.
2. Verify that the serial number on the replacement rear case matches the serial number on the original rear case.
3. Separate the infuser front and rear case assemblies as described in Section 7.2.2, Separating Front and Rear Case Assemblies.
4. Remove the following using the procedures described in the corresponding sections:
   - Battery pack cover - Section 7.3.4
   - Battery pack - Section 7.3.5
   - Pole clamp assembly - Section 7.3.7
   - Velcro strap and retainer plate - Section 7.3.8
   - Security door - Section 7.3.9
   - AC power cord - Section 7.3.12
   - AC power cord plug - Section 7.3.13
   - Printer/Dataway connector and control panel harness - Section 7.3.33
   - Battery Boost charger PWA (if applicable) - Section 7.3.39
5. Re-assemble the rear case assembly in the exact reverse order of disassembly.
6. Re-assemble the infuser in the exact reverse order of disassembly.
7. Close and lock the security door.

To verify successful replacement of the rear case assembly, perform the PVT as described in Section 5.2.

7.3.46
FRONT CASE ASSEMBLY REPLACEMENT

CAUTION: Use proper ESD grounding techniques when handling components. Wear an antistatic wrist strap and use an ESD-protected workstation. Store the PWA in an antistatic bag before placing it on any surface.

The recommended tools for this procedure are as follows: 1/4 inch nutdriver, X-acto knife, and isopropyl alcohol.

To replace the front case assembly, proceed as follows:

1. Remove the power supply PWA as described in Section 7.3.15, Power Supply PWA Replacement, and the CPU/display PWA as described in Section 7.3.16, CPU/Display PWA Replacement.
2. Remove the slide assembly as described in Section 7.3.30, Slide Assembly Replacement (-008 and Lower), or Section 7.3.31, Slide Assembly Replacement (-010 and Higher).
3. Using a 1/4 inch nutdriver, remove the four hex head screws that attach the front panel assembly to the front case. Remove the front panel assembly and install on the new case assembly.
4. Install the slide assembly into the new front case as described in Section 7.3.30 or Section 7.3.31.
5. Re-install the power supply PWA and the CPU/display PWA in the exact reverse order of disassembly.
6. Re-assemble the infuser in the exact reverse order of disassembly.

To verify successful front case assembly replacement, perform the PVT as described in Section 5.2.
7.3.47

RUBBER FOOT PAD REPLACEMENT

The recommended tools for this procedure are as follows: 3/8 inch wood chisel or X-acto knife, and isopropyl alcohol.

To replace the rubber foot pad, refer to Figure 7-13, Battery Pack Replacement, then proceed as follows:

1. Place the infuser on its side.
2. Using a 3/8 inch wood chisel or an X-acto knife, remove the rubber foot pad and scrape the enclosure recess to remove adhesive residue.
   
   **Note:** Each adhesive-backed rubber foot pad is bonded in its recess; do not damage the recess.

3. Using isopropyl alcohol, clean any adhesive residue from the enclosure recess.
4. Remove the protective backing from the self-adhesive surface of the replacement foot pad and press the pad in place.
5. After approximately five minutes, verify the foot pad is secure.

Replacement of a rubber foot pad is a routine maintenance procedure and no verification procedure is normally required. However, if the infuser may have been damaged during a rubber foot pad replacement, perform the PVT as described in Section 5.2.
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Section 8
SPECIFICATIONS

8.1
PCA PLUS

**Dimensions:**
- Height: 13 inches (33 cm)
- Width: 8 inches (20 cm)
- Depth: 6 inches (15 cm)

**Weight:** Approximately 15 pounds (7 kilograms)

**Power Requirements:** 110-120 VAC, 50/60 Hz, 25 watts

**Fuses:** Slo-blo, 0.25 A, 250 VDC

**Power Cord Plug:** Hospital grade (3 pin)

**Case Material:** High density polyurethane structural foam

**Battery:** 8 VDC, 3.0 ampere hour, 4 cell, sealed lead acid, rechargeable battery pack

**Battery Life:** Approximately four hours of operation with up to 30 mL of delivery when fully charged

**PCA Only Mode:** Approximately 1 mL in 35 seconds

**Continuous Only Mode:** Variable from 0.5 x concentration to 20 x concentration (mg/hr)

**PCA+Continuous** Variable from 0.5 x concentration to 20 x concentration (mg/hr), plus PCA rate

**Lockout Interval Range** 5 to 100 minutes in 1 minute increments

**Backpressure Range** 10 to 20 psig

**Operating Environment** 10 to 40 degrees C (50 to 104 degrees F) ambient temperature
- 5 to 95 percent relative humidity, noncondensing

**Electrical Safety:** Meets UL 544, ANSI SLC, and CSA 22.2 standards

**Administration Sets:** Use only compatible Abbott PCA sets with integral anti-siphon valve
8.2 PCA PLUS II

**Dimensions:**
- Height: 13 inches (33 cm)
- Width: 8 inches (20 cm)
- Depth: 6 inches (15 cm)

**Weight:**
Approximately 15 pounds (7 Kilograms)

**Power Requirements:**
110-120 VAC, 50/60 Hz, 25 watts

**Fuses:**
Slo-blo, 0.25 A, 250 VDC

**Power Cord Plug:**
Hospital grade (3 pin)

**Case Material:**
High density polyurethane structural foam

**Battery:**
8 VDC, 3.0 ampere hour, 4 cell, sealed lead acid, rechargeable battery pack

**Battery Life:**
Approximately four hours of operation with up to 30 mL of delivery when fully charged

**PCA Only Mode:**
Approximately 1 mL in 35 seconds

**Continuous Only Mode:**
Variable from 0.1 x concentration to 20 x concentration (mg/hr or µg/hr)

**PCA+Continuous**
Variable from 0.1 x concentration to 20 x concentration (mg/hr or µg/hr), plus PCA rate

**Lockout Interval Range**
5 to 100 minutes in 1 minute increments

**Backpressure Range**
10 to 20 psig

**Operating Environment**
10 to 40 degrees C (50 to 104 degrees F) ambient temperature
5 to 95 percent relative humidity, noncondensing

**Electrical Safety:**
Meets UL 544, ANSI SLC, and CSA 22.2 standards

**Administration Sets:**
Use only compatible Abbott PCA sets with integral anti-siphon valve
Section 9

DRAWINGS

*Figure 9-1, IPB for the Infuser*, through *Figure 9-7, Interconnect PWA Schematic*, detail the infuser through illustrated parts breakdown (IPB), interconnect, and schematic diagrams. *Table 9-1, Drawings*, lists drawings by figure number, title, and drawing number. *Table 9-2, IPB for the Infuser*, identifies infuser parts by index numbers that correlate to *Figure 9-1*.

**Note:** Figures listed in *Table 9-1* are rendered as graphic representations to approximate actual product; therefore, figures may not exactly reflect the product. Drawings and schematics in *Section 9* are provided as information only; drawings and schematics may not exactly reflect current product configurations.

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Morgan Hill, California 95037

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CAUTION: Federal (USA) law restricts this device to sale by or on the order of a physician or other licensed practitioner.

WARNING
POSSIBLE EXPLOSION HAZARD EXISTS IF INFUSION SYSTEM IS USED IN THE PRESENCE OF FLAMMABLE ANESTHETICS.

Covered by one or more of the following U.S. patents: 4,336,800; 4,453,932

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Attention, consult accompanying documents

IEC 601-1-1 Classification: Class 1, Type CF, Drip-Proof Medical Equipment

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