Level 1® H-1200
Fast Flow Fluid Warmer

With:
H-31, Version B, Air Detector/Clamp

REF H-1200 115 V
REF H-1200 230 V

OPERATOR’S MANUAL

P/N 4533708GB Rev. 002
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Level 1® H-1200 Fast Flow Fluid Warmer
Part Number: 4533708GB Rev. 002 (2006-07)

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The products described are covered by one or more of the following U.S. Patent Nos. 4,759,749; 4,878,537; 4,900,308; 5,063,994; 5,097,898; 5,417,274 and 5,512,043.

Manufactured in the U.S.A.
Contents

1 About this Manual 1
2 Description 3
3 Indications for Use 4
4 Important Safety Information 5
   Contraindications 5
   Warnings 5
   Cautions 8
5 Out of the Box—Assembly 9
   Step 1 Verify Components 10
   Step 2 Assemble I.V. Pole to Warming Unit 11
   Step 3 Install the Pressure Chambers 12
   Step 4 Attach the I.V. Bag Hanger 13
   Step 5 Disinfect the Recirculating Solution Reservoir 13
   Step 6 Preliminary Preparation 14
   Step 7 Connect the Pneumatic Tubing 14
   Step 8 Install the Level 1® H-31, Version B, Air Detector/Clamp 16
   Step 9 Perform Electrical Safety Tests 17
6 Principle of Operation 19
   Fluid Warming 19
   Pressurized Fluid Delivery 19
   Air Detector/Clamping 19
7 Controls and Displays 21
   Fluid Warmer Power and Alarm Test Panel 22
   Fluid Warmer Display Panel 23
   Air Detector/Clamp Control Panel and Alarms 24
   Pressure Chamber Control Panel 25
   Interlocks 26
8 Operation 29
   Modes Of Operation 30
      OFF Mode 30
      ON/ Automatic Operation Mode for Fluid Warmer 30
      Alarm Test Mode 31
      Over Temperature Test Mode 31
      Temperature Display 31
      Check Disposables Mode 32
      Add Recirculating Solution Mode 32
      Over Temperature Alarm Mode 33
      Power ON Test for the Air Detector/Clamp 33
      Automatic Operation Air Detector/Clamp 34
      Check Tubing Mode 34
      Air Detected/Clamped Mode 35
      Pressurized Display 35
      Pressurized Mode 35
      Unpressurized Mode 36
9 Operating Instructions

Warnings 37

9.1 Set Up for Use 38
A—Install Disposable Administration Set 38
B—Prime the Disposable Administration Set 39
C—Prime the Patient Line 41
D—Test the Audible and Visual Alarms 41
E—Test the Air Detector/Clamp 42

9.2 Use of the Fluid Warmer 44
Step 1—Load the Pressure Chambers 44
Step 2—Pressurize the Pressure Chambers 44
Step 3—Make Patient Connection 45
Step 4—Replace the Gas Vent/Filter Assembly 45
Step 5—Change the Fluid Bag 45

9.3 Replace the Gas Vent/Filter Assembly 46

9.4 Activated Alarms 47

9.5 After Use 49

10 Troubleshooting 50

General Troubleshooting Guide 50

11 Testing 51

Add Recirculating Solution Alarm 52
Check Disposables Alarm 52
Over Temperature Test 53
Fluid Warmer Alarm Signal Test 53
Performance Testing 54
Cold Start Test 54
Calibration Test 54
Alternative Calibration Test 54
Calibration Test with DSTA-40 54
Proper Calibration of Recirculating Solution Temperature 55

Periodic Electrical Testing 56
Earth Leakage 56
Ground Continuity 56

12 Maintenance 57

Maintenance Performed Prior to Every Use 57
Clean the Exterior 57
General Inspection 57

Maintenance Performed Every 30 Days 57
Lubricate O-Ring Seals 57
Change Recirculating Solution with Distilled Water 58

Maintenance Performed Every 12 Months 58
Disinfect the Recirculating Solution Reservoir 58
Change Recirculating Solution with 0.3% Hydrogen Peroxide/Distilled Water Solution 59
Change O-Rings 59
Clean Fan Filter 59
Inspect Air Detector/Clamp 59
Testing Fluid Warmer Operation 59

Maintenance and Calibration Log 60
Scheduled Maintenance and Calibration Checklist 60
SECTION 1 • About this Manual

About this Manual

These instructions contain important information for safe use of the product. Read the entire operator's manual, including Warnings and Cautions, before using the Level 1® H-1200 Fast Flow Fluid Warmer. Failure to properly follow warnings, cautions, and instructions could result in death or serious injury to the patient.

This Operator’s Manual describes the set-up, use, and maintenance of:

• Level 1® H-1200 Fast Flow Fluid Warmer
• Level 1® H-1000 Fast Flow Fluid Warmer with the Level 1® H-31, Version B, Air Detector/Clamp

Descriptions about the use and functionality of the Level 1® H-1200 Fast Flow Fluid Warmer in this manual are also applicable to the Level 1® H-1000 Fast Flow Fluid Warmer when equipped with the H-2 Pressure Chambers and Level 1® H-31, Version B, Air Detector/Clamp.

The manual is intended for use by individuals trained in the healthcare and biomedical professions.

WARNING!

Read the entire operator’s manual before using the Level 1® H-1200 Fast Flow Fluid Warmer. Failure to properly follow warnings, cautions, and instructions could result in death or serious injury to the patient.

This manual is organized into the following sections:

2 and 3 Description and Indications for Use
These sections provide the purpose and indications for use of the Level 1® H-1200 Fast Flow Fluid Warmer.

4 Important Safety Information
Lists the Contraindications, Warnings, and Cautions associated with the use of the Level 1® H-1200 Fast Flow Fluid Warmer.

5 Out of the Box—Assembly
Guides the user through the installation of the Level 1® H-1200 Fast Flow Fluid Warmer and the Level 1® H-31, Version B, Air Detector/Clamp.
6 Principle of Operation
Provides a functional description of the Level 1® H-1200 Fast Flow Fluid Warmer.

7 Controls and Displays
Provides a description of the function and purpose of the controls, displays, and indicators for the Level 1® H-1200 Fast Flow Fluid Warmer.

8 Operation
Describes Operation, Indicator, and Alarm modes of the Level 1® H-1200 Fast Flow Fluid Warmer.

9 Operating Instructions
Describes the Set Up, Use, and Alarm modes of the Level 1® H-1200 Fast Flow Fluid Warmer.

10 Troubleshooting
Contains information on troubleshooting the Level 1® H-1200 Fast Flow Fluid Warmer. This section also details troubleshooting slow fluid flow rates.

11 Testing
Describes Operational, Performance, and Electrical Tests that are used to verify the proper operation of the Level 1® H-1200 Fast Flow Fluid Warmer.

12 Maintenance
Regular maintenance procedures for every use, 30-day, and 12-month intervals are covered in this section.

13 Limited Warranty
Describes the Limited Warranty and its provisions.

14 Service
Explains Warranty Service and Non-Warranty Work as well as listing Service Contacts.

15 Specifications
Provides physical, environmental, and electrical specifications of the device.

16 Symbols
Lists the symbols and their definitions used with the Fluid Warmer.
SECTION 2 • Description

Description

The Level 1® H-1200 Fast Flow Fluid Warmer (Fluid Warmer) is an I.V. fluid warmer with pressure chambers, air detection, and automatic clamping capability. I.V. fluid and/or blood products are warmed through the use of a sealed heat exchanger through which a recirculating solution flows. Pressure Chambers apply pressurization and deliver the fluids at a fast flow rate. The Air Detector/Clamp monitors for the presence of air in the disposable Gas Vent/Filter Assembly.

This non-invasive method employs single-use, disposable administration sets that include a Gas Vent/Filter Assembly and Heat Exchanger. When air is detected in the Gas Vent/Filter Assembly, the Air Detector/Clamp closes off the patient line and alerts operators to the presence of air with audible and visual alarms. An ultrasonic signal continually passes through the fluid filled Gas Vent/Filter Assembly (GV/FA). As a bolus of air displaces the fluid in the GV/FA, the ultrasonic signal is broken and the clamp closes, stopping the air before it enters the patient line. Audible and visual alarms are activated, notifying the user that the fluid flow has stopped. Clearing the bolus of air and restoring the fluid flow are quickly accomplished without disconnecting from the patient.

Disposable Administration Sets

The installation, set up, and replacement of Level 1® Fast Flow I.V. Disposable Administration Sets (Disposable Sets) follows a four-step sequence that corresponds to numbered blocks on the device. Disposable Sets available for use on the Level 1® H-1200 Fast Flow Fluid Warmer are listed below.

- D-50 / DI-50
- D-60 / DI-60HL
- D-70 / DI-70
- D-100 / DI-100
- D-300 / DI-300

D-series Disposable Sets are for use in the U.S.A. DI-series Disposable Sets are for use in markets outside of the U.S.A.
Section 3: Indications for Use

The Level 1® H-1200 Fast Flow Fluid Warmer (Fluid Warmer) provides a rapid flow of warmed fluids, such as crystalloid or blood product, including red blood cells, as volume replacement for patients suffering from blood loss due to trauma or surgery.

The Fluid Warmer provides fast flow of warmed fluid to re-warm patients during surgery by trained medical personnel.
SECTION 4

Important Safety Information

This section covers information for prescribers and guidelines for safe use of the Level 1® H-1200 Fast Flow Fluid Warmer (Fluid Warmer).

CONTRAINDICATIONS

• Not for use in warming platelets, cryo-precipitates, or granulocyte suspensions.

WARNINGS

Death or serious injury may occur to the patient or user if these warnings are not followed:

• Read and follow all instructions, labeling, and accompanying documents supplied with this medical device. Failure to follow instructions, including all warnings and cautions, could lead to misuse of the device or device malfunction.

• Remove all air from fluid lines before connecting to patient. Failure to do so can result in infusion of air into the patient.

• Replace Gas Vent/Filter Assembly every three hours, or when the filter becomes clogged, or when air is slowly vented. Failure to do so will result in a reduction of flow rate. This may result in inadequate patient treatment.

• The replacement Gas Vent/Filter Assembly must be fully primed before continuing infusion. Failure to do so may allow air to be infused into the patient.

• Do not use the Fluid Warmer in high-energy fields such as: MRI, X-RAY, portable and mobile RF communications equipment, and other such devices. The Fluid Warmer may act as a projectile in a strong magnetic field, cause image artifacts, or not function as intended.
WARNINGS [continued]

• Do not bend the heat exchanger. Bending may damage the heat exchanger allowing communication between the recirculating solution and I.V. fluid path, resulting in the I.V. delivery of inappropriate fluids.

• Blood and blood products could contain pathogenic organisms. Failure to follow institutional policy and procedures for biomedical-hazardous materials could lead to exposure to harmful pathogens.

• When injecting medications into the fluid path, do not inject through the triple lumen tubing of the D/DI-60HL Disposable Set. This may allow communication between the recirculating solution and I.V. fluid path.

• Exposed conductor on MAINS power cord can cause an electrocution hazard. Remove device from service if MAINS power cord has exposed wires.

• Do not re-use partially full fluid bags. Fluid bags that have been partially drained, un-spiked, and then reinstalled may contain air, which if used can result in infusion of air into the patient. Use only new fluid bags from which the air has been removed.

• The tubing must be properly placed in the Clamp Slot of the Air Detector/Clamp. Failure to ensure that the tubing is correctly positioned in the Clamp Slot may result in failure to stop air infusion.

• Activation of the Air Detector/Clamp alarm during infusion indicates that fluid flow has stopped and that immediate operator intervention is required to restore fluid flow.

• Activation of the Over Temperature warning signal indicates that warming has stopped and immediate operator intervention is required to clear the over temperature condition or to take the device out of service.

• The Fluid Warmer is not for use with irrigating tubing, which may not fit into the clamp slot of the Air Detector/Clamp causing diminished flow or a failure to stop flow.
• The Fluid Warmer is for use only with Smiths Medical supplied or approved parts, accessories, and D or DI series Disposable Sets. The device may not function as intended with the use of unapproved parts, accessories, or Disposable Sets.

• Grounding reliability can only be achieved when MAINS power cords are connected to a properly grounded receptacle. Risk of electrical shock exists if the equipment is not connected to a properly grounded receptacle.

• The functional test for the Air Detector/Clamp accessory must be performed before each use. If the device does not function correctly, air in the I.V. line may not be detected, and the patient line may not be clamped. Remove the device from service immediately.

• Use of a bedside leukocyte reduction filter may cause a sudden precipitous drop in blood pressure resulting in respiratory distress, facial flushing, abdominal pain and nausea, and loss of consciousness. Immediately stop transfusion, and follow institution’s protocol for treatment of transfusion reactions.

• If any visual indicator does not illuminate or the audible signal does not sound, do not use the Fluid Warmer. Remove the device from service immediately.

• Do not operate the Fluid Warmer in the presence of a flammable anesthetic mixture with air, oxygen, or nitrous oxide. The risk of explosion exists if the Fluid Warmer is operated in a potentially explosive environment.

• No user-serviceable parts. All service must be performed by Smiths Medical or an authorized representative.

• Do not use Disposable Set if luer caps are not securely in place, else I.V. flow path may not be sterile and may cause death or serious injury.
CAUTIONS

Physical injury to the patient, user, and/or an adverse effect on the device or its performance may occur if these cautions are not followed:

• Do not use the Fluid Warmer if equipment or Disposable Set malfunction is evident.

• To reduce the risk of cross contamination, do not reuse Disposable Sets. Disposable Sets are for single use only.

• When loading fluid bags into Pressure Chambers, choose a hanging hook that allows the bag port to hang freely in the indented slot at the bottom of the chamber door. If bag ports are positioned above this slot, diminished flow could occur.

• Federal law (U.S.A.) restricts this device to sale by or on the order of a physician.
Out of the Box—Assembly

This device must be assembled and tested by authorized Smiths Medical personnel, an authorized distributor of Smiths Medical, or a qualified person prior to placing the device into service.

The following steps describe how to assemble and do preliminary set up of the Level 1® H-1200 Fast Flow Fluid Warmer (Fluid Warmer).

Refer to Step 8 if you need to install the Level 1® H-31, Version B, Air Detector/Clamp to the Level 1® H-1000 Fast Flow Fluid Warmer.

Step 1  Verify components of the Fluid Warmer
Step 2  Assemble I.V. Pole to Warming Unit
Step 3  Install Pressure Chambers
Step 4  Attach the I.V. Bag Hanger
Step 5  Disinfect the Recirculating Solution Reservoir
Step 6  Preliminary Preparation
Step 7  Connect the Pneumatic Tubing
Step 8  Install the Level 1® H-31, Version B, Air Detector/Clamp
Step 9  Perform Electrical Safety Tests

Read through the instructions completely prior to setting up the device.

Note: After unpacking the system, recycle packaging material according to hospital policy for recyclable materials.
Step 1

Verify Components of the Level 1® H-1200 Fast Flow Fluid Warmer

Note: The Level 1® H-31, Version B, Air Detector/Clamp is shipped as a separate accessory only for installation on an existing Level 1® H-1000 Fast Flow Fluid Warmer. Check the contents of all packaging to verify that the following components are present. If any parts are missing or damaged, do not use the Fluid Warmer. Do not substitute parts not supplied by Smiths Medical. Contact Smiths Medical for replacement parts. Below is a listing of the component parts for the Level 1® H-1200 Fast Flow Fluid Warmer.

### Components Checklist

<table>
<thead>
<tr>
<th>Qty</th>
<th>Component</th>
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<tbody>
<tr>
<td></td>
<td>in order of appearance in illustration</td>
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<tr>
<td>2</td>
<td>Pressure Chambers / Contents:</td>
</tr>
<tr>
<td></td>
<td>(2) “U” brackets</td>
</tr>
<tr>
<td></td>
<td>(4) Thumbscrews</td>
</tr>
<tr>
<td>1</td>
<td>I.V. Bag Hanger</td>
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<tr>
<td>1</td>
<td>Fluid Warming Unit</td>
</tr>
<tr>
<td>1</td>
<td>H-31, Version B, Air Detector/Clamp</td>
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<tr>
<td></td>
<td>Accessory Pack</td>
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<tr>
<td></td>
<td>(2) Pan-head screws</td>
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<td></td>
<td>(2) Power Cord Clips</td>
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<td></td>
<td>(2) Flat-head screws</td>
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<tr>
<td>1</td>
<td>Operator’s Manual</td>
</tr>
<tr>
<td>1</td>
<td>I.V. Pole with Flanking Brackets</td>
</tr>
<tr>
<td>1</td>
<td>Accessory Pack / Contents:</td>
</tr>
<tr>
<td></td>
<td>(2) Plastic “J” Clamps</td>
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<tr>
<td></td>
<td>(1) Y Connector</td>
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<td>(1) Black Tubing</td>
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<td></td>
<td>(1) Hex Wrench</td>
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<td>O-Ring Kit / Contents:</td>
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<td>(2) O-Rings</td>
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<td>(1) Hex Wrench</td>
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<td>(1) Instructions for Use</td>
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<tr>
<td>3</td>
<td>Bolts</td>
</tr>
<tr>
<td>3</td>
<td>Washers</td>
</tr>
<tr>
<td>1</td>
<td>I.V. Pole Base</td>
</tr>
</tbody>
</table>
Step 2

Assemble I.V. Pole to the Warming Unit

There are three steps involved in assembling the I.V. Pole to the Warming Unit. The steps are: 1. Assemble the I.V. Pole to the Base, 2. Close the Drain Valve, and 3. Attach the Warming Unit to the Flanking Brackets. Each step is detailed in a short procedure.

2.1 Assemble the I.V. Pole to the Base

1. Locate the I.V. Pole Base (a).

2. Locate the dark-gray extruded I.V. Pole (b) with Flanking Brackets.

3. Place the I.V. Pole Base upright on its wheels, (c) and lock the wheels to prevent movement during set up.

4. Locate three bolts (d) and washers for the pole base.

5. Align the three holes (e) in the I.V. Pole with the three screw holes on the pole base.

6. Slide the I.V. Pole down over the pole base, (f) keeping holes aligned.

7. Guide three bolts and washers through the holes (g) at the base of the pole and tighten.

2.2 Close the Drain Valve

Turn valve, located on the bottom of the device, perpendicular to stem (h) of the Warming Unit as shown.
2.3 Attach the Warming Unit to the Flanking Brackets

1. Align the eight hex screws on the back of the Warming Unit with the eight keyhole notches on the flanking bracket.

2. Slide screw heads down into keyhole notches.

3. Tighten all eight hex screws with the supplied hex wrench and secure in place.

Step 3

Install the Pressure Chambers

1. Locate the two Pressure Chambers.

2. Locate the U-brackets and thumbscrews supplied with the Pressure Chambers.

3. Attach the U-brackets with thumbscrews to the back of the Pressure Chambers, as shown. Keep thumbscrews and brackets loose.

4. Slide one Pressure Chamber with attached U-bracket over the top of each flanking pole.

5. Align the U-bracket slightly below the top of the flanking pole. Tighten the thumbscrews securely.
Step 4

Attach the I.V. Bag Hanger

1. Slide the I.V. Bag Hanger on top of the I.V. Pole.
2. Align with tabs.
3. Press down and snap into place.

Step 5

Disinfect the Recirculating Solution Reservoir

1. Remove the fill-port plug (a) on the reservoir.
2. Prepare a 0.3% hydrogen peroxide/distilled water solution for the reservoir. Mix 140 ml of 3% hydrogen peroxide solution and 1,260 ml of distilled water.
3. Fill the reservoir with 1.4 liters of 0.3% hydrogen peroxide/distilled water solution.
4. Replace the fill-port plug.
5. Insert a Disposable Set into the Fluid Warmer.
6. Insert the power cord into a properly grounded receptacle.
7. Turn the Fluid Warmer ON. Let the solution circulate for a 30-minute disinfection period.
8. Turn the Fluid Warmer OFF.
9. Empty the reservoir.
10. Remove the Disposable Set and discard according to established hospital procedures.
Step 6

Preliminary Preparation

1  Remove the fill-port plug (a) on the front of the warming unit and fill the reservoir to the maximum level with 1.4 liters of one of the following solutions:

- 0.3% Hydrogen Peroxide/Distilled Water Solution
  Mix 140 ml of 3% hydrogen peroxide and 1,260 ml of distilled water.
  **Note:** If this option is selected, the maintenance requirement to change the recirculating solution is once every 12 months. Always use a 0.3% hydrogen peroxide/distilled water solution when refilling the reservoir.

- Distilled Water
  **Note:** If this option is selected, the maintenance requirement to change the recirculating solution is once every 30 days.

2  Replace the fill port plug.

3  Lubricate O-Rings in #1 Block (b) and #2 Block (c). Place a small amount of silicone lubricant, provided in the supplied O-Ring Kit, on a cotton swab and apply all around the inside of each O-Ring.

Step 7

Connect the Pneumatic Tubing

1  Locate the Accessory Pack with the black pneumatic tubing (a), two “J” clamps (b), and one “Y” Connector (c).
2 Locate the orange protective plug in the red ring connector (d) located on the back of the device. Remove the plug by depressing the red plastic ring as you pull the plug out of the connector.

3 Take the pneumatic tubing and press one end of the tubing firmly into the ring connector (d) until it can go no further.

4 Take the “Y” connector and press the other end of the pneumatic tubing into the bottom of the “Y” connector, as shown (e), until it can go no further.

5 Press the pneumatic tubing from the Pressure Chamber into place (f) on the top of the “Y” connector until it can go no further. Repeat this procedure with the pneumatic tubing from the other Pressure Chamber.

6 Remove the protective backing sheet on one “J” clamp, exposing the adhesive side.

7 Carefully position the “J” clamp and press the adhesive side against the gray I.V. Pole in the approximate locations (g)(h) shown.

Press down firmly to secure in place.

Repeat this procedure for the other “J” clamp.

8 Press the pneumatic tubing into place on the “J” clamps.
Step 8

Install the Level 1® H-31, Version B, Air Detector/Clamp

1. Remove 4 screws (a) from the rear of the Fluid Warmer, lower left side, from locations shown in figure.

2. Plug in the power cord (b) of the Level 1® H-31, Version B, Air Detector/Clamp into the auxiliary MAINS outlet located on the bottom of the Fluid Warmer.

   Note: If you turn the Fluid Warmer off and the Air Detector/Clamp does not turn off, contact Smiths Medical or an authorized representative.

3. Loosen the two screws holding the mounting bracket to the rear of the Air Detector/Clamp (do not remove).

4. Align the slot on the Air Detector/Clamp with #3 Block on the Fluid Warmer (c). Fit in place over the block.

5. Align the mounting bracket on the rear of the Air Detector/Clamp with two screw holes, shown (d). Insert two flat-head screws and tighten, securing the Air Detector/Clamp to the Fluid Warmer. Tighten the two screws on the mounting bracket on the Air Detector/Clamp.

6. Locate two power cord clips (e) included with the Air Detector/Clamp and snap onto power cord. Align clips with the two screw holes as shown, insert screws, and tighten.
Step 9

Perform Electrical Safety Tests

Perform all applicable electrical safety tests as required per institutional procedure. These include but are not limited to:

- Leakage current
- Hypot
- Ground bond test

WARNING!

Grounding reliability can only be achieved when MAINS power cords are connected to a properly grounded receptacle. Risk of electrical shock exists if the equipment is not connected to a properly grounded receptacle resulting in death or serious injury to the patient or user.

The Electrical Safety Check must be performed by qualified personnel authorized by the institution to perform such testing. The Safety Check must be performed and documented at least once per year, or according to institutional policy.
Principle of Operation

The schematic illustration on the facing page depicts the Level 1® H-1200 Fast Flow Fluid Warmer’s (Fluid Warmer) operations. The primary operations are described below.

Fluid Warming

The Fluid Warmer utilizes a solution reservoir housed in a controller unit. Recirculating solution is warmed and pumped through a heat exchanger (a part of the Disposable Set). The solution is returned to the reservoir for continuous recirculation and remains isolated from the patient and from the I.V. fluid path. The on-board recirculating solution is heated to a pre-set manufacturer’s temperature set-point. The system continuously monitors and controls the recirculating solution temperature. The Fluid Warmer is designed to shut down and provide audible and visual alarms in the event of an over-temperature condition.

Pressurized Fluid Delivery

The Fluid Warmer provides pressurized fluid delivery through the use of an on-board compressor and two Pressure Chambers. The Pressure Chambers pressurize the fluid bags for fast fluid delivery.

Air Detection/Clamping

The Air Detector/Clamp detects the presence of air in the Gas Vent/Filter Assembly (GV/FA)—a part of the Level 1® D/DI series Disposable Administration Sets (Disposable Sets)—and clamps the patient line. An ultrasonic signal continually passes through the fluid-filled GV/FA. As a bolus of air displaces the fluid in the GV/FA, the ultrasonic signal is broken and the clamp closes, stopping the air before it enters the patient line. Audible and visual alarms are activated, notifying the user that air has been detected and fluid flow has been clamped off. Clearing the bolus of air and restoring the fluid flow are quickly accomplished without disconnecting from the patient.
Section 7

Controls and Displays

Five locations on the Level 1® H-1200 Fast Flow Fluid Warmer (Fluid Warmer) govern how the device is controlled and where function indicators are displayed. They are called-out in the figure on the facing page and are defined in the list below.

1. Fluid Warmer Display Panel
2. Power and Alarm Test Panel
3. Reservoir Level Display
4. Air Detector/Clamp Control Panel
5. Pressure Chamber Control Panel

The Fluid Warmer has five Interlocks, which detect for correct installation of a Disposable Set, that are also defined this section.
Fluid Warmer Power and Alarm Test Panel

The Power and Alarm Test Panel is located on the front of the Fluid Warmer directly above the reservoir fill-cap. This panel contains four pressure-sensitive buttons that are activated when pressed. Refer to the Power Alarm Test Panel (figure on right) whose numbered call outs correspond to a description of the button and the function it performs.

**Button/Function**

1. **Power ON Button**
   The green button on the top-left of the Power and Alarm Test Panel powers on the device. Power is applied to the on-board compressor for the pressure chambers and the Air Detector/Clamp. The green Automatic Operation LED on the Fluid Warmer Display Panel illuminates when the Power ON button is activated, with a Disposable Set in place.

2. **Power OFF Button**
   The Power OFF is the orange button to the right of the Power ON button on the Power and Alarm Test Panel. This button turns power off to the unit. The green Automatic Operation LED on the Display Panel will turn off when this button is pressed.

3. **Over Temperature Test Button**
   The Over Temperature Test is used to confirm the proper operation of the Over Temperature circuitry. Testing the circuitry requires that the Fluid Warmer is at operating temperature (41°C). Once this is established, press and hold the button. Then, release the button. The Over Temperature alarm continues to function. Clear the alarm mode by turning the device off, then back on. See Section 11, *Testing*, for instruction on performing an Over Temperature Test.

4. **Fluid Warmer Alarm Signal Test Button**
   The Fluid Warmer Alarm Signal Test is used to confirm proper operation of the visual and audible alarm indicators. Press and hold this button to test circuitry. Then, release the button. The Over Temperature alarm continues to function. Clear the alarm mode by turning the device off, then back on.

5. **Reservoir Capacity**
   Capacity for recirculating solution reservoir is 1.4 liters. Use recirculating solution. Do not exceed maximum capacity.
Fluid Warmer Display Panel

The Fluid Warmer Display Panel provides continuous information about the operation of the Fluid Warmer. A liquid crystal display (LCD) indicates recirculating solution temperature. Just below the LCD, four light-emitting diodes (LEDs) indicate operation modes for the device. For identification purposes, the diodes are shown illuminated.

1 Recirculating Solution Temperature
The temperature of the recirculating solution is displayed in the LCD panel. The temperature is displayed in degrees celsius.

Note: This is NOT the temperature of fluid delivered to the patient—the display reflects the temperature of the recirculating solution.

2 Automatic Operation LED
The green LED indicator illuminates when the power is ON and the Disposable Set has been properly installed. When lit, this indicates the Fluid Warmer is operating.

3 Check Disposables LED
The yellow LED indicator illuminates and an audible attention signal beeps when the Disposable Set is not properly installed. See the Interlocks description in this section for directions on clearing the Check Disposables alarm.

4 Add Recirculating Solution LED
The yellow LED indicator illuminates and an audible attention signal beeps when reservoir is low. Additional recirculating solution must be added to the reservoir. Maximum capacity for the reservoir is 1.4 liters of recirculating solution.

5 Over Temperature LED
The red LED indicator illuminates and an audible warning signal beeps when the recirculating solution is over the acceptable temperature for safe use.

Reservoir Level Display

The Reservoir Level Display has a clear window for viewing the amount of recirculating solution present in the reservoir. Check the reservoir to ensure the solution level is near the maximum level indicator (α). If the recirculating solution level is too low, the Add Recirculating Solution LED on the Display Panel illuminates and an audible attention signal beeps.
Air Detector/Clamp Control Panel and Alarms

The Air Detector/Clamp Control Panel has three LED indicators that display the operational state of the Air Detector/Clamp. Refer to figure on right.

1 Automatic Operation LED
   The green Automatic Operation LED illuminates when the following conditions are present: The Fluid Warmer power is ON, a Disposable Set is properly installed in the Fluid Warmer and primed, the patient line from the Gas Vent/Filter Assembly is correctly placed in the #3 Clamp Slot, and the Clamp Slot door is closed.

2 Check Tubing LED
   The yellow Check Tubing LED illuminates and an audible attention signal beeps when the patient line tubing from the Gas Vent/Filter Assembly is not correctly placed in the #3 Clamp slot, and when the Clamp Slot door is not closed correctly.

3 Clamped LED
   The red Clamped LED illuminates and an audible warning signal beeps when air is detected in the Gas Vent/Filter Assembly. The patient line is automatically clamped.
Pressure Chamber Control Panel

The Pressure Chamber Control Panel uses a control lever to switch from pressurized to unpressurized mode. A gauge displays pressure levels in the Pressure Chamber.

1 Pressurize / Unpressurize Lever
This lever is used to control the pressure mode in the Pressure Chamber.

a To Pressurize the Pressure Chamber
With Pressure Chamber door closed and latched, slide lever to the left, all the way to the plus (+) pressurized position. This applies 300 mmHg pressure in the Pressure Chamber when the Fluid Warmer is turned ON.

b To Unpressurize the Pressure Chamber
To remove pressure from the Pressure Chamber, slide the lever to the right, all the way over to the minus (–) unpressurized position. Pressure is released on the fluid bag in the Pressure Chamber.

2 Pressure Gauge
This gauge indicates the pressure present in the Pressure Chamber. When the Pressurize lever is in the plus (+) pressurized position and the Fluid Warmer is ON, this gauge displays the operating pressure in the Pressure Chamber. The operating pressure should be 300 mmHg.
Interlocks

The Fluid Warmer has five Interlocks that detect for proper installation of a Disposable Set’s components. Refer to the figure on the facing page to identify the positions of interlocks.

Note: Block 1 is not an Interlock. It cannot detect if a Disposable Set is not correctly installed. It is identified here because it is an essential step for proper installation of the Disposable Set components.

Three Interlocks are located on the Fluid Warmer and check for proper installation of:

1. Heat Exchanger, top end
2. Gas Vent/Filter Assembly
3. Heat Exchanger (guide)

Two Interlocks are located in the Air Detector/Clamp 3 and check for the proper installation of:

4. Patient I.V. Line in the Clamp Slot
5. Door for the Clamp Slot

Interlocks 2, 4, and 5 prevent the Fluid Warmer’s pump from circulating reservoir solution if the Disposable Set’s Heat Exchanger and Gas Vent/Filter Assembly are not installed properly.

If Check Disposable Alarm is Activated on the Fluid Warmer

a. Check Heat Exchanger for proper installation in Block 1, and Interlocks 2 and 5.

b. Press Heat Exchanger down firmly in Block 1 to secure in O-Ring.

c. Press Interlock 2 tab down firmly to engage the Interlock switch.

d. Press Heat Exchanger firmly into Interlock 5.

e. Check Gas Vent/Filter Assembly installation in Interlock 4.

If Check Tubing Alarm is Activated on the Air Detector/Clamp

a. Open the door and check the Patient Line for proper installation in the Clamp Slot at interlock 3.

b. Close the door and check that the tab on the top edge of the door is fully inserted into the Air Detector/Clamp at interlock 3 before pushing the door down to close it.
## Operation

**Level 1** H-1200 Fast Flow Fluid Warmer

<table>
<thead>
<tr>
<th>Function</th>
<th>Interface</th>
<th>Description</th>
<th>Indicator/ Alarm Signal</th>
</tr>
</thead>
<tbody>
<tr>
<td>Fluid Warmer Power and Alarm Test Panel</td>
<td>OFF</td>
<td>None</td>
<td>None</td>
</tr>
<tr>
<td></td>
<td>ON</td>
<td>LEDs display</td>
<td>None</td>
</tr>
<tr>
<td></td>
<td>Alarm Test</td>
<td>LEDs display</td>
<td>1/second</td>
</tr>
<tr>
<td></td>
<td>Over Temperature Alarm Test</td>
<td>LEDs display</td>
<td>1/second</td>
</tr>
<tr>
<td>Fluid Warming</td>
<td>Temperature Display</td>
<td>LCD Readout</td>
<td>None</td>
</tr>
<tr>
<td></td>
<td>Automatic Operation</td>
<td>Green LED</td>
<td>None</td>
</tr>
<tr>
<td></td>
<td>Check disposables</td>
<td>Yellow LED</td>
<td>1/5 seconds</td>
</tr>
<tr>
<td></td>
<td>Add recirculating solution</td>
<td>Yellow LED</td>
<td>1/5 seconds</td>
</tr>
<tr>
<td></td>
<td>Over temperature</td>
<td>Red LED</td>
<td>1/second</td>
</tr>
<tr>
<td>Air Detection/Clamping</td>
<td>Automatic operation</td>
<td>Green LED</td>
<td>None</td>
</tr>
<tr>
<td></td>
<td>Check tubing/patient line</td>
<td>Yellow LED</td>
<td>1/5 seconds</td>
</tr>
<tr>
<td></td>
<td>Air detected/clamped</td>
<td>Red LED</td>
<td>1/second</td>
</tr>
<tr>
<td>Pressurized Fluid Delivery</td>
<td>Pressure gauge</td>
<td>Numbered dial</td>
<td>None</td>
</tr>
<tr>
<td></td>
<td>Pressurized</td>
<td>+ Symbol</td>
<td>None</td>
</tr>
<tr>
<td></td>
<td>Unpressurized</td>
<td>– Symbol</td>
<td>None</td>
</tr>
</tbody>
</table>
The Level 1® H-1200 Fast Flow Fluid Warmer (Fluid Warmer) performs three primary functions; Fluid Warming, Air Detection/Clamping, and Pressurized Fluid Delivery. Functions are monitored and controlled by four interfaces/control panels located on the Fluid Warmer.

The four interfaces are:

- Fluid Warmer Power and Alarm Test Panel
- Fluid Warmer Display Panel
- Air Detector/Clamp Alarm and Control Panel
- Pressure Chamber Control Panel

In the table on the facing page, operation of the device is represented in terms of the four interfaces that control specific device functions. The numbers call-out individual Modes of Operation activated or indicated on the interface.

**Functions**
- Fluid Warming
- Air Detection/Clamping
- Pressurized Fluid Delivery

**Operational modes**
- OFF Mode
- ON/Automatic Operation for Fluid Warmer
- Alarm Test Mode
- Over Temperature Test Mode
- Check Disposables Mode
- Add Recirculating Solution Mode
- Over Temperature Alarm Mode
- Power ON Test for Air Detector/Clamp
- Automatic Operation Air Detector/Clamp
- Check Tubing Mode
- Air Detected/Clamped Mode
- Pressurized Mode
- Unpressurized Mode

The modes of operation are individually defined in the following section. This includes a description of each mode, activation and/or monitoring of the mode, mode characteristics, and clearing of the mode state.
Modes of Operation

WARNING!
If any visual indicator does not illuminate or the audible signal does not sound, do not use the Fluid Warmer. Remove the device from service immediately. Death or serious injury may occur to the patient or user if this warning is not followed.

OFF Mode
Power is off only for a part of the equipment. The MAINS are still connected. Press the OFF button (a) on the Power and Control Panel to turn the device off.

ON/Automatic Operation Mode for Fluid Warmer
The Fluid Warmer enters Automatic Operating mode when a Disposable Set is properly installed and the device is turned ON. This is done by pressing the ON button (b).

Mode characteristics
• The green Automatic Operating LED on the Display Panel illuminates (c).
• Fluid warming begins.
• Pressure infusion is provided by activating the Pressure Chambers.
• Air Detector/Clamp enters the Power ON Test, then enters default Automatic Operation mode.
**Alarm Test Mode**

The Alarm Test mode is used to test the visual and audible indicators of the Level I® H-1200 Fast Flow Fluid Warmer. This mode is entered by pressing and holding the Alarm Test button (d) on the Fluid Warmer’s Control Panel.

**Mode characteristics**
- All visual indicators on the Fluid Warmer’s Display Panel (e) illuminate.
- The Fluid Warmer’s audible alarm beeps.
- When the Alarm Test button is released, the Over Temperature LED and an audible alarm remains active.
- To clear the Over Temperature alarm, turn the Fluid Warmer OFF and then back ON.

---

**Over Temperature Test Mode**

The Over Temperature Test mode is used to test the operation of the Fluid Warmer’s Over Temperature Circuitry. This mode is entered by pressing and holding the Over Temperature Test button (f) on the Fluid Warmer Control Panel with the Fluid Warmer at operating temperature (41°C).

**Mode characteristics**
- The red Over Temperature LED (g) on the Display Panel illuminates.
- An audible warning signal beeps.

**To Clear this mode**
- Turn Fluid Warmer OFF.
- Turn Fluid Warmer back ON.

---

**Temperature Display**

The Temperature Display functions when the Fluid Warmer is powered ON. Temperature is displayed in degrees Celsius.
Check Disposables Mode

The Check Disposables mode of the Fluid Warmer indicates a missing or improperly installed Disposable Set.

Mode characteristics
- The Check Disposables yellow LED (a) on the Fluid Warmer’s Display Panel is illuminated.
- An audible attention signal beeps.
- Reservoir solution circulation is stopped; fluid warming stops.
- Pressure Chambers continue to operate.

To Clear this mode
Install a disposable or check the disposable installation as follows:
- Check the position of the disposable in the #1 Block.
- Make sure the heat exchanger is seated in the heat exchanger guide.
- Press down firmly on the #2 Block.
- Check the position of the Gas Vent/Filter Assembly in the #4 Interlock.

Add Recirculating Solution Mode

The Add Recirculating Solution mode of the Fluid Warmer indicates that the solution level in the recirculating solution reservoir is below its minimum level.

Mode characteristics
- The yellow Add Recirculating Solution LED (b), on the Fluid Warmer’s Display Panel, is illuminated
- An audible warning signal beeps.
- Solution circulation stops, fluid warming stops.
- Pressure Chambers continue to operate.
- Fluid flow to the patient continues.

To Clear this mode
Add recirculating solution to the Fluid Warmer’s reservoir.
Over Temperature Alarm Mode

The Over Temperature Alarm mode is entered when the temperature of the recirculating solution reservoir is at or above 43.9°C.

**WARNING!**

Activation of the Over Temperature warning signal indicates that warming has stopped and immediate operator intervention is required. Failure to clear the over temperature condition or to take the device out of service may result in patient death or serious injury.

**Mode characteristics**

- The Over-Temperature LED warning light illuminates (c).
- An audible warning signal beeps.
- Solution circulation is stopped; fluid warming stops.
- Pressure Chambers continue to operate.
- Fluid flow to the patient continues.

**To Clear Over Temperature Alarm mode**

Do the following:

- Turn OFF the Fluid Warmer to clear the alarm.
- Turn the power back ON.

Power ON Test for Air Detector/Clamp

The **Power ON Test** for the Air Detector/Clamp is activated when the Fluid Warmer is turned ON. This test activates the Air Detector/Clamp’s visual and audible indicators.

**Mode characteristics**

- Audible alarm indicator beeps.
- All LED indicators on the Air Detector/Clamp Control Panel illuminate:
  - Automatic Operation LED indicator - Green
  - Check Tubing LED indicator - Yellow
  - Clamped LED indicator - Red

At the end of the Power ON Test the Air Detector/Clamp enters Automatic Operation mode. This is the default mode for the Air Detector/Clamp.
Automatic Operation Air Detector/Clamp

In Automatic Operating mode, the Air Detector/Clamp monitors for the presence of air in the Disposable Set’s Gas Vent/Filter Assembly (GV/FA). If air is detected the patient line is clamped off and an audible alarm beeps. The Air Detector/Clamp goes into Automatic Operation mode when the Fluid Warmer is turned ON and Disposable Set is properly installed and primed—no air is present in the GV/FA.

Mode characteristics

- The green Automatic Operation LED (1) is illuminated.
- Monitoring for air in the Gas Vent/Filter Assembly is active.
- Fluid is ready to be delivered to the patient when in Automatic Operating mode.

Check Tubing Mode

The Air Detector/Clamp’s Check Tubing mode is entered when the patient line from the Gas Vent/Filter Assembly is not properly installed in the Air Detector/Clamp Slot and when the Clamp Slot door is not closed correctly.

Mode characteristics

- The yellow Check Tubing LED (b) on the Air Detector/Clamp Control Panel is illuminated.
- An audible low-priority warning signal beeps.

To Clear this mode

- Place the patient line from the Gas Vent/Filter Assembly in the #3 Clamp Slot of the Air Detector/Clamp and close the Clamp Slot door.
Air Detected/Clamped Mode

In this mode the Air Detector/Clamp is activated when the ultrasonic sensor detects the presence of air in the Gas Vent/Filter Assembly. Auditory and visual warnings are activated and the clamp is closed, preventing passage of fluid through the patient line.

Mode characteristics

• The patient line is clamped until alarm condition is removed.
• The red Clamped warning indicator LED (c) illuminates.
• The Air Detector/Clamp warning signal beeps.

To Clear this mode

Perform the steps in Section 9, Operating Instructions, under the heading: Clear the “Air Detected” Alarm mode.

Pressure Display

The Pressure Display gauge (a) functions when the device is ON. The graduated indicator represents the level of pressure applied to the fluid bags. Pressure should be in the range of 280-300 mmHg pressure.

Pressurized Mode

The H-2 Pressure Chambers deliver fluids at an increased flow rate with the application of 300 mmHg pressure upon the fluid bags. Flow rate varies according to fluid type and viscosity, temperature, Disposable Set used, and the amount of clamping applied to the roller clamps.

Pressurized infusion is enabled when:

• A fluid bag is installed in the Pressure Chamber.
• The Fluid Warmer is turned ON.
• The Pressure Chamber switch is placed in the plus (++) pressurized position (b).

Mode characteristics

• The Fluid Warmer must be ON for the Pressure Chamber to work.
• Operating pressure should be between 280-300 mmHg.
• Pressure is not adjustable on the Pressure Chamber.
• Pressure is applied to the fluid bag in the Pressure Chamber.
• Pressure is indicated on the pressure gauge.
To Exit this mode
• Place the Pressure Chamber switch in the minus (−) unpressurized position (c).

Unpressurized Mode

Disable pressurization by moving the switch on the Pressure Chamber to the minus (−) unpressurized position (c).

Mode characteristics
• Pressure is released from the Pressure Chamber.
• A fluid bag can be removed or loaded into the Pressure Chamber.
SECTION 9

Operating Instructions

The Operating Instructions are grouped into five segments. Read through each section BEFORE performing a procedure.

WARNINGs

• For use only with Smiths Medical supplied or approved parts, accessories and Level 1® D/DI series Disposable Administration Sets (Disposable Sets). The device will not function as intended with the use of unapproved parts, accessories, or Disposable Sets resulting in death or serious injury to the patient.

• When injecting medications into the fluid path, do not inject through the triple-lumen tubing of the Level 1® D/DI-60HL Disposable Set. This may allow communication between the recirculating solution path and I.V. fluid path, which could result in death or serious injury to the patient.

• Replace Gas Vent/Filter Assembly (GV/FA) every three hours, or when the filter becomes clogged, or when air is slowly vented. Failure to do so will result in a reduction of flow rate. This may result in inadequate patient treatment resulting in death or serious injury to the patient.

• Replacement Gas Vent/Filter Assembly must be fully primed before continuing infusion. Failure to do so may allow air to be infused into the patient resulting in death or serious injury to the patient.

• Grounding reliability can only be achieved when MAINS power cords are connected to a properly grounded receptacle. Risk of electrical shock exists if the equipment is not connected to a properly grounded receptacle resulting in death or serious injury to the patient or user.

• Do not bend the heat exchanger. Bending may damage the heat exchanger allowing communication between the recirculating solution path and I.V. fluid path, resulting in the I.V. delivery of inappropriate fluids which could result in death or serious injury to the patient.

• The Gas Vent/Filter Assembly (GV/FA) must be oriented properly in the Detector Head. Failure to do so could result in air not venting from the GV/FA which may cause the Air Detector/Clamp to malfunction resulting in death or serious injury to the patient.

• The tubing must be properly placed in the Clamp Slot of the Air Detector/Clamp. Failure to ensure that the tubing is correctly positioned in the Clamp Slot may result in failure to stop air infusion which may result in patient death or serious injury.
9.1 Set Up for Use

**WARNINGS**

- Read and follow all instructions, labeling, and accompanying documents supplied with this medical device. Failure to follow instructions, including all warnings and cautions, could lead to misuse of the device or device malfunction.

- Disposable Sets are supplied with a sterile fluid path which may be compromised if the caps are not in place. Do not use administration sets if luer and spike caps are not securely in place, else flow path may not be sterile and may cause death or serious injury to the patient.

- Disposable Sets are for single use only. To reduce the risk of cross contamination, do not reuse Disposable Sets.

**A—Install the Disposable Administration Set**

The installation sequence for the Disposable Administration Set corresponds to the numbered Blocks marked 1-2-3-4 on the Fluid Warmer.

Remove the Disposable Set from its packaging and review the Instructions for Use provided. Do not remove spike caps or luer caps at this time.

1. Push the bottom end of the Heat Exchanger (a) [the end near the Gas Vent/Filter Assembly] into #1 Block (b). Press the Heat Exchanger down firmly to properly seat in the block.
   
   **Note:** D/DI-60HL Disposable Sets require the Heat Exchanger to be placed with the Patient Line extending to the left.

2. Slide #2 Block up (c). Snap Heat Exchanger into guide (d). Press firmly into place to ensure it is properly seated. Slide #2 Block down (e), push down firmly to secure.
3 Move the pinch clamp on the Patient Line to a point midway between the Gas Vent/Filter and the luer connector. Close the pinch clamp.

**Note:** Level 1® D/DI-60HL Disposable Sets do not have a pinch clamp on the Patient Line. In this case, move the pinch clamp below the Gas Vent/Filter Assembly (GV/FA) to a midway point between the GV/FA and the luer connector. Then, close the pinch clamp.

4 Install the Gas Vent/Filter Assembly. Refer to the series of figures on the left.

   a Open the #3 Clamp Slot door by pushing down on tab, (a) and lifting up the front of the door (b).
   b Pull the door down (c) and away from the Clamp Slot.
   c Insert the Patient Line in the Clamp Slot (d) and push it back into the slot.
   d Hold the Patient Line in the Clamp Slot and push the door up (e) to engage the top hinge. Then push the front of the door down to close it.
   e Pull the Patient Line to the right (f) to align it in the Clamp Slot without kinking.
   f Align the Gas Vent/Filter Assembly to the #4 Block, (g) and press it into place.

---

**B—Prime the Disposable Administration Set**

1 Close the Disposable Set clamps above the Heat Exchanger.
   - For D/DI-50, D/DI-60HL, D/DI-70 Disposable Sets, 
     - close pinch clamps below the bag spikes, and
     - close roller clamp below the drip chamber (a).
   - For D/DI-100, D/DI-300 Disposable Sets
     - close ratchet clamps below the drip chambers.

2 Remove all air from the fluid bag:

   a Invert solution bag.
   b Use aseptic technique. Pierce membrane of bag port with spike of Disposable Set. Then withdraw spike.
   c Squeeze bag to exhaust ALL air.
   d Place spike in bag port. Do not allow air to re-enter bag.
3 Remove spike cap and insert spike into the port of an air-free fluid bag. Repeat this step for each fluid line to be used.

4 Slide lever on H-2 Pressure Chamber to the minus (–) unpressurized position.

5 Hang spiked fluid bag/s in Pressure Chamber:
   a Release hinged latch, open door and hang fluid bag inside on tab appropriate for bag size.
   b Close the door and secure latch.
   c Injection and Spike ports on the fluid bag should extend from opening at the bottom of the Pressure Chambers without being obstructed.

   Note: When installing Level 1® D/DI-300 series Disposable Sets, hang the third fluid bag from the I.V. pole.

---

**CAUTIONS**

- When loading fluid bags into H-2 Pressure Chambers, choose a hanging hook that allows the bag port to hang freely in the indented slot at the bottom of the chamber door. If bag ports are positioned above this slot, diminished flow could occur.
- Do not overextend I.V. pole; If I.V. pole is overextended the Fluid Warmer may become unstable.

6 Open clamp above Drip Chamber on Level 1® D/DI-50, D/DI-60HL, and D/DI-70 Disposable Administration Sets for each I.V. fluid bag being used to prime the drip chamber.

7 Prime Drip Chamber by squeezing drip chamber until one-third to one-half full of fluid.

   Note:  
   • D/DI-50, D/DI-60HL, D/DI-70 series use a single drip chamber.
   • D/DI-100 and D/DI-300 series use a separate drip chamber for each spiked bag. The clamp is below the drip chamber.

8 Open remaining clamps above the Heat Exchanger. Fluid flows into the Gas Vent/Filter Assembly (GV/FA).

9 Vigorously tap the Gas Vent/Filter Assembly to dislodge air bubbles from filter screen.

10 Press the green Power ON button, located on the Power and Alarm Test Panel, to turn ON the Fluid Warmer.
• The Air Detector/Clamp runs a Power ON Test.
• All Air Detector/Clamp indicator LEDs illuminate.
• The audible warning beeps.
• Upon completion of the Power On Test the Air Detector/Clamp enters Operation mode with the Automatic Operation LED illuminated.
• If the Disposable Set is incorrectly installed, the Fluid Warmer’s Check Disposables attention indicator illuminates and the audible attention signal beeps. Check the installation of the Disposable Set following the directions provided in Section 6 Controls and Displays, on Interlocks.

C—Prime the Patient Line

1. Remove the male luer cap from the distal end of the Patient Line.

   Note: On Level 1® D/DI-60HL Disposable Administration Sets (D/DI-60 HL Disposable Sets), verify that no recirculating solution comes out of the distal end of the Patient Line.

   ! WARNING

   If fluid exits in the Patient Line or the D/DI-60HL Disposable Set, replace the Disposable Set.

2. Open the pinch clamp below the Gas Vent/Filter Assembly.

3. Allow fluid to flow until no air is observed in the Patient Line and the line is primed with fluid. Then, close the pinch clamp, roller clamp, or ratchet clamp on the Patient Line.

   Note: On D/DI-60HL Disposable Sets close roller clamp below Drip Chamber.

D—Test the Audible and Visual Alarms

Test the visual and audible alarm signals by performing the following steps.

1. Press and hold the Alarm Test button on the Fluid Warmer’s Power and Alarm Test Panel.
   • All Fluid Warmer visual alarm LEDs illuminate and the audible alarm signal beeps.

2. Release the Alarm Test button; the Over Temperature alarm continues.
3 Clear the Over Temperature alarm condition.
   • Turn the Fluid Warmer OFF, then ON.
   • The Air Detector/Clamp runs a Power On Test.
   • The Air Detector/Clamp goes into Automatic operation.

E—Test the Air Detector/Clamp

1 Slide the lever on the Pressure Chambers to the plus (+) pressurized position to pressurize fluid delivery.

2 Move the top of the Gas Vent/Filter Assembly away from the Air Detector sensor as shown.

3 The following occurs:
   • The patient line clamp closes.
   • The red Clamped indicator LED illuminates.
   • The audible warning signal beeps.
   • Fluid Warmer disposable alarm activates.

4 Open the pinch clamp, roller clamp, or ratchet clamp on the Patient Line to verify that fluid does not flow.

5 Return to normal operation by pressing the top of the Gas Vent/Filter Assembly back into the #4 Block.

6 The Air Detector/Clamp resumes Automatic Operation mode.
   • The green Automatic Operation LED on the Air Detector/Clamp Control Panel illuminates.

7 The Fluid Warmer is now ready for patient connection. Unclamp Patient Line to begin infusion.

   Note: On D/DI-60HL Disposable Sets open roller clamp below drip chamber.

Conclusion

This concludes Section 9.1, Set Up for Use. Operators can proceed to the next Section 9.2, Use of the Fluid Warmer.
WARNINGS

• Remove all air from fluid bags before connecting to patient. Failure to do so can result in infusion of air into the patient resulting in death or serious injury to the patient.

• Do not reuse partially full fluid bags. Fluid bags that have been partially drained, un-spiked, and then reinstalled may contain air, which if used can result in infusion of air into the patient resulting in death or serious injury to the patient. Use only new fluid bags from which the air has been removed.

• Do not leave the tubing in a closed Pinch Clamp for longer than three hours, as this may cause tubing deformation that could result in diminished flow resulting in patient death or serious injury.

• Replace Gas Vent/Filter Assembly every three hours, or when the filter becomes clogged, or when air is slowly vented. Failure to do so will result in a reduction of flow rate. This may result in inadequate patient treatment resulting in death or serious injury to the patient.

• The Replacement Gas Vent/Filter Assembly must be fully primed before continuing infusion. Failure to do so may allow air to be infused into the patient which could result in patient death or serious injury.

CAUTION

• When loading fluid bags into Pressure Chambers, choose a hanging hook that allows the bag port to hang freely in the indented slot at the bottom of the chamber door. If bag ports are positioned above this slot, diminished flow could occur.
9.2 Use of the Fluid Warmer

Use of the Fluid Warmer requires that the steps in Section 9.1, Set Up for Use have been completed.

Overview

Use of the Fluid Warmer involves the following steps:
1—Load the Pressure Chambers
2—Pressurize the Pressure Chambers
3—Make patient connection, begin infusion
4—Replace Gas Vent/Filter Assembly
5—Change fluid bag

Step 1—Load the Pressure Chambers

a Turn the hinged latch on the right side of the Pressure Chamber outward. Open the door.

b Hang a solution bag on the appropriate hanging hook inside the door. The Pressure Chamber can hold bags of varying sizes.

- On the inside of the Pressure Chamber door are hooks for bags smaller than 1000ml.
- On the top of the Pressure Chamber door are hooks appropriate for 1000ml bags.
- Bags from different fluid manufacturers vary somewhat in their dimensions.
- Choose a hanging hook that allows the bag drain port to hang freely in the indented slot at the bottom of the Pressure Chamber door.

c Close the door and secure side latch.

Step 2—Pressurize the Pressure Chambers

a Turn ON the Pressure Chamber by moving the lever located at the top of the Pressure Chambers over to the plus (+) pressurized position.

b Check gauge to ensure pressure of 280-300 mmHg is achieved.
  - Pressure in the chambers is not adjustable.

Note: Power must be ON for the Pressure Chambers to operate.
**WARNING!**

Blood and blood products could contain pathogenic organisms. Failure to follow Institutional policy and procedures for biomedical-hazardous materials could lead to exposure to harmful pathogens which could result in user death or serious injury.

---

**Step 3—Make Patient Connection**

Make patient connection and begin infusion.

**Step 4—Replace the Gas Vent/Filter Assembly**

Replace the Gas Vent/Filter Assembly every 3 hours, when the filter becomes clogged, or if air is venting slowly. Refer to Section 9.3, *Replace the Gas Vent/Filter Assembly.*

**Step 5—Change the Fluid Bag**

a. Move the lever on the Pressure Chamber over to the minus (−) unpressurized position. This will release the pressure in the Pressure Chamber and deflate the bladder.

   Close ratchet clamp under empty bag.

b. Open door and remove the fluid bag from the Pressure Chamber.

c. Remove the spike from the used fluid bag.

d. Remove any air from the new fluid bag and spike the fluid bag.

e. Hang the new fluid bag in the Pressure Chamber. Close and latch the door.

f. Move the lever on the Pressure Chamber over to the plus (+) pressurized position to pressurize the chamber.

g. Open ratchet clamp.
9.3 Replace the Gas Vent/Filter Assembly

Use one of the following Gas Vent/Filter Assemblies:

- F-30 Filter with Gas Vent for D/DI-300 Disposable Sets
- F-10 Filter with Gas Vent for all other Fast Flow Disposable Sets

1 Close all clamps above and below the Gas Vent/Filter Assembly on D-series Disposable Set and on new Gas Vent/Filter Assembly.

2 Turn the Fluid Warmer OFF.

3 Remove used Gas Vent/Filter Assembly while still connected to the Disposable Set:
   a Remove Gas Vent/Filter Assembly from the #4 Block.
   b Open the #3 Clamp Slot door and remove the Patient Line from the Clamp Slot.

4 Install the new Gas Vent/Filter Assembly:
   a Open the Clamp Slot door.
   b Insert outlet tube in the Clamp Slot.
   c Close the Clamp Slot door.
   d Align the outlet tube in the Clamp Slot without kinking.
   e Press the Gas Vent/Filter Assembly into the #4 Block.

5 Using aseptic technique:
   a Disconnect upper luer fitting from the used Gas Vent/Filter Assembly.
   b Remove upper luer end cap from new Gas Vent/Filter Assembly.
   c Connect the Disposable Set to new Gas Vent/Filter Assembly inlet.

6 With Gas Vent/Filter Assembly clamped, open the clamp above Gas Vent/Filter Assembly. The Gas Vent/Filter Assembly will self prime.

7 Turn power ON.

8 Remove the end cap on the outlet tube. Slowly release the outlet tube clamp and allow outlet tube to fill completely.

9 Holding the used Gas Vent/Filter Assembly horizontally, disconnect lower patient line luer lock. Connect used filter luer-lock fittings together and discard.

10 Connect outlet tube luer lock to patient line.

11 Open clamps below the Gas Vent/Filter Assembly and resume infusion.
9.4 Activated Alarms

Refer to Section 8, Operation for information on identifying Alarm states, conditions that activate them, and methods for clearing Alarm states.

A. “Air Detected” Alarm mode

Detection of air in the Gas Vent/Filter Assembly results in the following:

• The patient line is clamped off 
• The red Clamped LED warning signal illuminates 
• The audible warning signal beeps

WARNING!

Activation of the Air Detector/Clamp Alarm during infusion indicates that fluid flow has stopped and that immediate operator intervention is required to restore fluid flow. Failure to reinstate flow (after purging any air or foam) may result in patient death or serious injury.

B. Clear the “Air Detected” Alarm mode:

1. Close the following clamp/s:
   • The clamp under the I.V. bag/s
   • The clamp below or after the Gas Vent/Filter Assembly
   • The Roller Clamp on the D/II series Disposable Set

2. Remove pressure from the Pressure Chambers by moving the lever to the minus (–) unpressurized position.

3. Inspect the entire circuit for the presence of air, locate the source of the air, and correct this condition.

Note: Air or foam may have been vented through the Gas Vent/Filter Assembly.
4 Remove any remaining air from the circuit:
   a Insert spike into an air-free bag/s of I.V. solution.
   b Place the I.V. bag/s in the Pressure Chamber/s.
   c Move the lever to the plus (+) pressurized position to
      pressurize the Chamber/s.
   d Prime the drip chamber/s and open the Disposable Set clamps
      above the Gas Vent/Filter Assembly.
   e Fluid flows freely through the tubing. Air in the I.V. line is
      vented out through the Gas Vent/Filter Assembly. When air is
      no longer present in the Gas Vent/Filter Assembly, the
      Clamp opens and resumes automatic operation mode.

**Note:** *If air is not freely vented, replace the Gas Vent/Filter
Assembly.*

Refer to Section 9.3, *Replace the Gas Vent/Filter Assembly.*

5 If no warning signals are active, the Fluid Warmer with Air
Detection and Pressure Chambers is ready for use.

6 Open the remaining Disposable Set clamps, slowly open the
roller clamp, and reestablish fluid flow to the patient.

For more information on the following alarm conditions, refer to Section 8,
*Operation:*

- Over Temperature Alarm
- Check Disposables Alarm
- Check Tubing Alarm
- Add Recirculating Solution Alarm
9.5 After Use

1 Discontinue infusion.

2 Turn the Fluid Warmer OFF.

3 Release chamber pressure before opening the Pressure Chamber door:
   a Move the lever to the minus (–) unpressurized position. This will release the pressure in the chamber and deflate the bladder.
   b Open door and remove the fluid bag.

4 Remove the Disposable Set from the Fluid Warmer and the Air Detector/Clamp Assembly.

5 Properly discard the Disposable Set in containers marked for biohazardous materials. Dispose by incineration, or follow hospital policies and procedures applicable for the disposal of biohazardous material.

6 Visually check the condition of the device. Remove from service any unit that shows physical damage.

7 Clean the device with warm soapy water.
Troubleshooting

Only authorized personnel should perform any routine maintenance and repairs to the Level 1® H-1200 Fast Flow Fluid Warmer.

The following two tables feature general troubleshooting information along with slow flow rate troubleshooting.

**General Troubleshooting Guide**

<table>
<thead>
<tr>
<th>Problem</th>
<th>Check the following:</th>
</tr>
</thead>
<tbody>
<tr>
<td>No Power</td>
<td>Check to see if unit is plugged in and power is turned ON. Be sure the unit is plugged into a working MAINS receptacle.</td>
</tr>
<tr>
<td>Disposables Alarm beeps</td>
<td>Check to see that the Heat Exchanger and Gas Vent/Filter Assembly are properly installed.</td>
</tr>
<tr>
<td>Add Recirculating Solution Alarm</td>
<td>Fill reservoir to the maximum reservoir level with recirculating solution.</td>
</tr>
<tr>
<td>Over - Temperature Alarm beeps</td>
<td>Press OFF button to clear alarm. Then, press ON button to power on. If the Fluid Warmer continues to alarm, discontinue use of the medical device and remove from service. Contact Smiths Medical or an authorized representative for service.</td>
</tr>
<tr>
<td>Hot Cabinet</td>
<td>Check the air inlet on the bottom of the unit. Remove any blockage or dust to insure adequate air flow.</td>
</tr>
<tr>
<td>Heat Exchanger hard to install</td>
<td>Lubricate O-Rings in #1 and #2 block heat exchanger sockets with silicone lubricant. Silicone lubricant part # 80-04-002.</td>
</tr>
<tr>
<td>Loud Compressor</td>
<td>Verify pneumatic tubing is fully seated into fittings.</td>
</tr>
</tbody>
</table>
| LED doesn’t light up during set up (on Fluid Warmer or Air Detector/Clamp) | 1. Verify unit is plugged into MAINS.  
  2. If still no LED illumination, discontinue use of the medical device and remove from service.                                                       |
| Tubing doesn’t fit in Air Detector/Clamp     | Verify tubing is Level 1® D/DI series tubing.                                                                                                            |
### Slow Flow Rate Troubleshooting

<table>
<thead>
<tr>
<th>Problem</th>
<th>Check the following:</th>
</tr>
</thead>
<tbody>
<tr>
<td>Pressure Chambers not activated</td>
<td>Verify the Fluid Warmer is ON. Verify the Pressure Chamber levers are fully in the plus (+), pressurized position. Verify pneumatic tubing is fully seated into fittings on the Pressure Chambers and Fluid Warmer.</td>
</tr>
<tr>
<td>Blood develops particulate</td>
<td>Stored blood develops particulate. If blood is older than 5-7 days, consider using Level 1® PF-1 pre-filters.</td>
</tr>
<tr>
<td>Clogged filter</td>
<td>Change Gas Vent/Filter Assembly if filter becomes clogged.</td>
</tr>
<tr>
<td>Fluid bag not fully spiked or bag port twisted</td>
<td>Make sure the membrane of the fluid bag port is fully pierced by the bag spike and that the neck of the bag port is not twisted.</td>
</tr>
<tr>
<td>Non-Level 1® bag port filters</td>
<td>40 to 80 micron filters used between the bag port and the spikes of the Disposable Set may restrict flow. Consider using Level 1® PF-1 pre-filters (340 micron).</td>
</tr>
<tr>
<td>Clamps partly engaged</td>
<td>Verify all clamps are fully open.</td>
</tr>
<tr>
<td>Clamps left in the clamped position for long periods</td>
<td>Leaving clamps fully clamped for long periods will cause the tubing to become deformed. Do not leave clamps closed for extended periods of time.</td>
</tr>
<tr>
<td>Kinked tubing</td>
<td>Verify that no tubing kinks are present.</td>
</tr>
<tr>
<td>Air trapped on filter screen of Gas Vent/Filter Assembly</td>
<td>Remove Gas Vent/Filter Assembly and tap against the cabinet of the Fluid Warmer to dislodge air bubbles, allowing them to vent. If this fails to correct problem, replace with new Gas Vent/Filter Assembly.</td>
</tr>
<tr>
<td>Non-high flow extension lines</td>
<td>Use only extension lines with an inner diameter of 0.103&quot; (3.3mm) or larger, such as Level 1® X-36 or Y-30 extension sets.</td>
</tr>
<tr>
<td>Stopcock</td>
<td>Use only Level 1® SC-3 (9 french inner diameter).</td>
</tr>
<tr>
<td>Small gauge needle or catheter</td>
<td>Use large bore needles or catheters to maximize flow rates.</td>
</tr>
</tbody>
</table>
Testing

This unit should be tested by hospital biomedical personnel prior to placing it in service. All testing and maintenance should be performed by qualified personnel. If qualified personnel are not available, contact Smiths Medical. If the Level 1® H-1200 Fast Flow Fluid Warmer (Fluid Warmer) and installed accessories do not pass the tests, discontinue use of the medical device and remove from service. Contact Smiths Medical or an authorized representative for service. Testing requires a Level 1® Disposable Administration Set (Disposable Set) to be installed in the Fluid Warmer.

Add Recirculating Solution Alarm

The Fluid Warmer is equipped with a float switch that senses the level of the recirculating solution in the reservoir. When the solution level is too low, an LED on the Display Panel illuminates and an audible alarm beeps. In the Add Recirculating Solution Alarm mode, the circulating pump is not running. With a Disposable Set in place and the unit turned on, test the Add Recirculating Solution Alarm by draining the solution until the level has dropped below the minimum reservoir level. The Add Recirculating Solution Alarm should activate. To drain recirculating solution from the Fluid Warmer, turn the drain valve, on the bottom of the unit 90 degrees clockwise and allow some of the recirculating solution to drain into a container.

Check Disposables Alarm

Five Interlocks detect the proper installation of an Disposable Set in the Fluid Warmer. If a Disposable Set is not properly installed and the power is ON, an indicator will illuminate and an audible alarm beeps. With the Fluid Warmer ON, the Interlocks should be tested one at a time by performing the following steps.

1. Top Heat Exchanger Socket – Slide the #2 Block up slowly. The Check Disposables indicator will illuminate and the audible alarm beeps.

2. Heat Exchanger Interlock – Gently pull on the middle of the heat exchanger. The Check Disposables indicator will illuminate and the audible alarm beeps.
3 Gas Vent/Filter Assembly Interlock – Pull the top of the Gas Vent/Filter Assembly from the #4 Block. The Check Disposables indicator will illuminate and the audible alarm beeps.

4 #3 Check Tubing Interlock on the Air Detector/Clamp – Remove the tubing from the Clamp Slot and close the door. The Check Tubing alarm signal on the Air Detector/Clamp is activated and the audible alarm beeps.

5 #3 Check Door Closed Interlock on the Air Detector/Clamp – With tubing installed in the Clamp Slot, open the door. The Check Tubing alarm signal on the Air Detector/Clamp is activated and the audible alarm beeps.

**Over Temperature Test**

Do the following
1 Insure that the Fluid Warmer is at operating temperature (41°C).
2 Press and hold the Over Temperature Test button (a).
3 The Over Temperature LEDs illuminate, and an audible alarm beeps.
4 Release the Over Temperature Test button.
5 Over Temperature LED and audible alarm signal remains active.

Clear the Alarm mode
1 Turn the Fluid Warmer OFF.
2 Turn the Fluid Warmer back ON.

**Fluid Warmer Alarm Signal Test**

The Alarm Test button is used to confirm proper operation of the visual and audible alarm indicators.

Do the following
1 Press and hold the Alarm Test button (b).
2 The LED illuminates, and an audible alarm beeps.
3 Release the Alarm Test button.
4 Over Temperature LED remains lit, and the audible alarm continues to beep.

Clear the Alarm mode
1 Turn the Fluid Warmer OFF.
2 Turn the Fluid Warmer back ON.
Performance Testing

Cold Start Test
Store the Fluid Warmer unit in a room where the room temperature is approximately 21°C (70°F).

1. Put a Disposable Set in place.
2. Record the start time.
3. Turn the Power button ON.
   • The green system operational indicator illuminates.
   • The Air Detector/Clamp goes through the Power ON Test.
   • Rapidly rising numbers will appear on the recirculating solution temperature display.
   • Within 60 seconds (±) the display should read at least 30°C.
   • In 3 to 10 minutes the display should read 41°C.

The Temperature Set Point is 41.7°C (+/- 0.3°C).

**Note:** If the Fluid Warmer does not pass the Cold Start Test, it should be removed from use and returned to Smiths Medical or an authorized representative for service.

Calibration Test
One approved way to confirm proper calibration of the recirculating solution temperature is to use the Level 1® Thermal Calibration Well (TCW) (Part Number 80-03-002). A replacement filter (F-10 or F-30) must be in place to utilize the TCW.

**Note:** Use of the TCW requires a digital thermometer NIST traceable and accurate within 0.1°C. Required probe size: 0.099” OD (Outside Diameter) maximum (0.25 cm) 0.50” — 1.50” long (1.27—3.81cm).

Alternative Calibration Test
Another approved way to confirm proper calibration of the recirculating solution temperature is to use the Level 1® DSTA-40 TEMPCHECK (Part Number 80-01-004).

Calibration Test with DSTA-40
The DSTA-40 TEMPCHECK (DSTA-40) is an electronic thermometer used to verify the recirculating solution operating temperature of the Fluid Warmer. The DSTA-40 uses thermistor technology to sense the temperature of the recirculating solution temperature for the Fluid Warmer. The recirculating solution temperature is displayed by a liquid crystal display (LCD). The DSTA-40 is powered from the auxiliary outlet of the Fluid Warmer. No batteries are required.

**Note:** The Level 1® DSTA-40 TEMPCHECK is available for purchase from Smiths Medical.
Proper Calibration of Recirculating Solution Temperature

Confirm proper calibration of recirculating solution temperature by performing the following steps:

1. Plug the DSTA-40 into the auxiliary outlet located on the bottom of the Fluid Warmer.

2. Install the DSTA-40 in the heat exchanger position (#1 Block and #2 Block).

3. Install a test filter in the #4 Block.

4. Turn the Fluid Warmer ON. Allow to warm up until the DSTA-40 temperature display stabilizes.

5. Compare the DSTA-40 temperature display with the temperature display on the Fluid Warmer’s Display Panel.

**Note:** The Fluid Warmer’s temperature display must read within 0.3°C of the Level 1® DSTA-40 TEMPCHECK display on a properly calibrated unit.

If the DSTA-40 reads 41.7°C and the Fluid Warmer’s temperature display is within the range of 41.4°C-42.0°C, the calibration is OK.
Periodic Electrical Testing

Earth Leakage

The Fluid Warmer must be tested in accordance with EN 60601-1. The earth leakage current test should be performed with the immersion heater circuit in the full ON condition; for this reason the leakage current test should be performed on units which have room temperature recirculating solution in the reservoir. Units not meeting this standard should be returned to Smiths Medical or an authorized representative for service.

Ground Continuity

The Fluid Warmer must be tested in accordance with EN 60601-1.

---------------------------------------------------------------------

⚠️ WARNING!

Grounding reliability can be achieved only when MAINS power cords are connected to a properly grounded receptacle. Risk of electrical shock exists if the equipment is not connected to a properly grounded receptacle resulting in death or serious injury to the patient or user.

---------------------------------------------------------------------
Maintenance

Only authorized personnel should perform any routine maintenance and repairs to the Level 1® H-1200 Fast Flow Fluid Warmer (Fluid Warmer). Maintenance is scheduled prior to each use, every 30 days, and annually. The tasks are detailed below.

**Note:** If distilled water is used as the recirculating solution, change the solution every 30 days. If a 0.3% hydrogen peroxide/distilled water solution is used as the recirculating solution, change the solution every 12 months, and during the 12 month period, always refill the reservoir with a 0.3% hydrogen peroxide/distilled water solution.

**Maintenance Performed Prior to Every Use**

Clean and inspect the Fluid Warmer with Air Detector/Clamp and Pressure Chambers after each use.

**Clean the Exterior**

- Clean the entire Fluid Warmer after every use. Use only mild detergents, water, and a soft cloth.
- To disinfect external surfaces, use 10% bleach/distilled water solution.

**General Inspection**

- Check the condition of the Fluid Warmer with a visual inspection. Remove from service any unit that shows physical damage.
- If the Disposable Set does not install easily, lubricate the O-Rings as directed in the following section.

**Maintenance Performed Every 30 Days**

**Lubricate O-Ring Seals**

It is not necessary to disassemble the blocks to lubricate the O-Rings.

1. Place a small amount of silicone grease on a cotton swab.

2. Apply the silicone grease along the O-Rings in the bottom #1 Block, and top #2 Block Heat Exchanger sockets.
Change Recirculating Solution with Distilled Water

1. Place a container under the drain valve of the Fluid Warmer.
2. Drain the recirculating solution by turning the drain valve clockwise 90 degrees.
3. When all solution has drained from the reservoir, close the drain valve.
4. Refill the reservoir with distilled water. The reservoir holds 1.4 liters.

Maintenance Performed Every 12 Months

Disinfect the Reservoir

1. Place a container under the drain valve of the Fluid Warmer.
2. Drain the recirculating solution by turning the drain valve clockwise 90 degrees.
3. When all solution has drained from the reservoir, close the drain valve.
4. Remove the fill-port plug on the reservoir.
5. Prepare a 0.3% hydrogen peroxide/distilled water solution. Mix 140 ml of 3% hydrogen peroxide solution and 1,260 ml of distilled water.
6. Fill the reservoir with 1.4 liters of 0.3% hydrogen peroxide/distilled water solution.
7. Replace the fill-port plug.
8. Insert a Disposable Set into the Fluid Warmer.
9. Turn the Fluid Warmer ON. Let the solution circulate for a 30-minute disinfection period.
10. Turn the Fluid Warmer OFF.
11. Empty the reservoir.
12. Remove the Disposable Set and discard according to established hospital procedures.
Change Recirculating Solution with a 0.3% Hydrogen Peroxide/Distilled Water Solution

1. Place a container under the drain valve of the Fluid Warmer.
2. Drain the recirculating solution by turning the drain valve clockwise 90 degrees.
3. When all the solution has drained from the reservoir, close the drain valve.
4. Prepare a 0.3% hydrogen peroxide/distilled water solution. Mix 140 ml of 3% hydrogen peroxide solution and 1,260 ml of distilled water.
5. Fill the reservoir with 0.3% hydrogen peroxide/distilled water solution. The reservoir holds 1.4 liters.

Change O-Rings

Change the O-Rings in the #1 Block and #2 Block.

1. Remove each O-Ring from its socket by pulling it out with a pair of needle-nose pliers or by prying it out with a small screwdriver.
2. Lubricate the new O-Rings from the O-Ring Kit.
3. Press each O-Ring into its socket.

Clean Fan Filter

The Fan Filter (a) is located on the bottom of the Fluid Warmer. The fan guard snaps in place.

1. Remove four screws and unsnap the fan guard from the bottom of the unit.
2. Clean the filter with warm soapy water.
3. Replace the fan guard and filter.

Inspect Air Detector/Clamp

1. Check Clamp Cover Door for proper closure.
2. Check the Clamp Cover Door, Clamp Slot and Detector Head for structural integrity.

Testing Fluid Warmer Operation

Perform all the tests described in the testing section of this manual. See Section 11, Testing.

- Add Recirculating Solution Alarm
- Check Disposables Alarm
- Over Temperature Alarm
- Performance Testing
### Maintenance and Calibration Log

All maintenance and testing should be done by qualified personnel. Regularly scheduled maintenance ensures proper functioning of the equipment. Refer to the table below for required tasks and frequency of routine maintenance.

### Scheduled Maintenance and Calibration Checklist

<table>
<thead>
<tr>
<th>Task</th>
<th>Every Use</th>
<th>Every 30 Days</th>
<th>Every 12 Months</th>
</tr>
</thead>
<tbody>
<tr>
<td>Clean Exterior</td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>General Inspection</td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>Change Distilled Water</td>
<td></td>
<td></td>
<td>☐</td>
</tr>
<tr>
<td>Lubricate O-Rings</td>
<td></td>
<td></td>
<td>☐</td>
</tr>
<tr>
<td>Disinfect Recirculating Solution Reservoir</td>
<td></td>
<td></td>
<td>☐</td>
</tr>
<tr>
<td>Change 0.3% Hydrogen Peroxide Solution</td>
<td></td>
<td></td>
<td>☐</td>
</tr>
<tr>
<td>Replace O-Rings</td>
<td></td>
<td></td>
<td>☐</td>
</tr>
<tr>
<td>Clean Fan Filter</td>
<td></td>
<td></td>
<td>☐</td>
</tr>
<tr>
<td>Test Over Temperature Alarm</td>
<td></td>
<td></td>
<td>☐</td>
</tr>
<tr>
<td>Test Add Recirculating Solution Alarm</td>
<td></td>
<td></td>
<td>☐</td>
</tr>
<tr>
<td>Test Disposable Alarm</td>
<td></td>
<td></td>
<td>☐</td>
</tr>
<tr>
<td>Test Fluid Warmer Alarm Signal</td>
<td></td>
<td></td>
<td>☐</td>
</tr>
<tr>
<td>Verify Temperature Calibration</td>
<td></td>
<td></td>
<td>☐</td>
</tr>
<tr>
<td>Electrical Safety Tests</td>
<td></td>
<td></td>
<td>☐</td>
</tr>
</tbody>
</table>
The Level 1® H-1200 Fast Flow Fluid Warmer (Fluid Warmer) and the Level 1® H-31, Version B, Air Detector/Clamp (Air Detector/Clamp) are warranted by Smiths Medical ASD, Inc., (Smiths Medical) to be free from defects in material or workmanship for a period of 1 year (12 months) from the date of shipment to the customer.

If the customer finds any Fluid Warmer or Air Detector/Clamp to have such defects during this period, it should be returned to the address provided in the Service section of this manual. At the option of Smiths Medical, the product will either be repaired or replaced by a new Fluid Warmer or Air Detector/Clamp, and returned to the customer. Provided Smiths Medical confirms that there were defects in the Fluid Warmer or Air Detector/Clamp, Smiths Medical will also refund the customer’s reasonable cost of returning the Fluid Warmer or Air Detector/Clamp for repair.

This warranty will not apply in respect of any Fluid Warmer or Air Detector/Clamp product which does not have its original serial number plates intact. Nor will this warranty apply to any damage or defect caused by misuse of the product, by careless or deliberate mistreatment of the product, or by any impact to the product.

In no event will Smiths Medical or its distributors be liable for consequential or economic loss incurred by the customer. The liability of Smiths Medical and its distributors for any defect in the Fluid Warmer or Air Detector/Clamp product will be limited to the invoice value of the product.

This warranty does not affect any warranty or guarantee to which the customer is irrevocably entitled by virtue of any applicable law. With that proviso, this warranty replaces all other express or implied warranties, representations, or indemnities to which the customer may otherwise be entitled by virtue of any law, trade practice, or otherwise.
All service must be performed by Smiths Medical or an authorized service representative. Service by any other person or organization voids the warranty and transfers liability for malfunctions of the device to the servicing organization.

**Warranty Service**

Units received for repair which have not been obviously abused or impact damaged and are still under warranty will be promptly repaired and returned at no charge. See the limited warranty section of this manual. A no-charge purchase order is requested for tracking.

**Non-Warranty Work**

Units received which have suffered obvious abuse or impact damage and units no longer under warranty will be promptly inspected and a verbal estimate of repair cost will be supplied. A purchase order will be required from the hospital consistent with the verbal estimate. A written estimate will be provided upon request.

Before returning your Level 1® H-1200 Fluid Warmer (Fluid Warmer) or Level 1® H-31, Version B, Air Detector/Clamp for service, contact Smiths Medical for Returned Goods Authorization. Be sure that ALL recirculating solution is drained from the unit before packing the Fluid Warmer for shipment.

**Note:** The Fluid Warmer must be cleaned and disinfected for repair shipment or it will be immediately returned as received.

**Additional Documentation**

Upon request Smiths Medical will provide the following documentation:

- Circuit diagrams
- Components parts list(s)
- Description of function
- Service and calibration instructions
Disposal Information

The Level 1® H-1200 Fast Flow Fluid Warmer contains lead that is used in solder of electric assembly. When you are ready to dispose of the device, observe federal, state, and local codes or requirements for disposal of hazardous materials and for recycling of solid waste materials that may impact the environment.

Service Contacts

Contact your Smiths Medical Technical Service Department or Smiths Medical distributor at:

USA/Canada

Smiths Medical ASD, Inc.
160 Weymouth Street
Rockland, MA 02370 USA

USA/Canada 1-800-258-5361
International +1-781-878-8011

European Representative

Smiths Medical International Ltd
Colonial Way, Watford,
Herts, WD24 4LG, UK

Tel +44 (0) 1923 246434
Fax +44 (0) 1923 240273

Australian Representative

Smiths Medical Australasia Pty. Ltd.
61 Brandl Street
Eight Mile Plains, QLD 4113, Australia

Tel +61 (0) 7 3340 1300
Fax +61 (0) 7 3340 1399
### System Specifications

<table>
<thead>
<tr>
<th>Physical</th>
<th>Measurement</th>
</tr>
</thead>
<tbody>
<tr>
<td>Height, Overall</td>
<td>67 inches (1.7 m)</td>
</tr>
<tr>
<td>Width, Overall</td>
<td>20 inches (51 cm)</td>
</tr>
<tr>
<td>Depth, Overall</td>
<td>20 inches (51 cm)</td>
</tr>
<tr>
<td>Weight Assembled; Dry</td>
<td>63 pounds (28.5 kg)</td>
</tr>
<tr>
<td>Recirculating Solution Capacity</td>
<td>0.37 gallons (1.4 L)</td>
</tr>
<tr>
<td>Air Source Pressure</td>
<td>300 (294 ± 6) mm/Hg</td>
</tr>
</tbody>
</table>

<table>
<thead>
<tr>
<th>Environmental</th>
<th>Temperature</th>
<th>Humidity [%]</th>
</tr>
</thead>
<tbody>
<tr>
<td>Operation</td>
<td>10°C to 40°C</td>
<td>10 to 95</td>
</tr>
<tr>
<td>Transportation</td>
<td>-18°C to 60°C</td>
<td>5 to 95</td>
</tr>
<tr>
<td>Storage</td>
<td>5°C to 40°C</td>
<td>5 to 95</td>
</tr>
</tbody>
</table>

<table>
<thead>
<tr>
<th>Thermal</th>
<th></th>
<th></th>
</tr>
</thead>
<tbody>
<tr>
<td>Temperature Set Point</td>
<td>41.7°C ± 0.3°C</td>
<td></td>
</tr>
<tr>
<td>Over Temperature Set Point</td>
<td>43.9°C ± 0.1°C</td>
<td></td>
</tr>
</tbody>
</table>

<table>
<thead>
<tr>
<th>Electrical</th>
<th>Type</th>
</tr>
</thead>
<tbody>
<tr>
<td>Protection Against Electrical Shock</td>
<td>Class I Equipment</td>
</tr>
<tr>
<td>Type BF Equipment</td>
<td></td>
</tr>
<tr>
<td>Mode of Operation</td>
<td>Continuous</td>
</tr>
<tr>
<td>Type of Current</td>
<td>Alternating</td>
</tr>
<tr>
<td>Ingress Protection Rating</td>
<td>IPX1</td>
</tr>
<tr>
<td>MAINS Power Input (REF H-1200 115 V)</td>
<td>115 VAC, 60 Hz, 12 Amps</td>
</tr>
<tr>
<td>MAINS Power Input (REF H-1200 230 V)</td>
<td>230 VAC, 50 Hz, 6.3 Amps</td>
</tr>
<tr>
<td>Auxiliary MAINS Outlet (REF H-1200 115 V)</td>
<td>1.5 Amps</td>
</tr>
<tr>
<td>Auxiliary MAINS Outlet (REF H-1200 230 V)</td>
<td>0.75 Amps</td>
</tr>
</tbody>
</table>
Electromagnetic Environment Recommendations

The Fluid Warmer is intended for use in an electromagnetic environment in which radiated RF disturbances are controlled. The customer or the user of the Fluid Warmer can help prevent electromagnetic interference by maintaining a minimum distance between portable and mobile RF communications equipment (transmitters) and the Fluid Warmer as recommended below, according to the maximum output power of the communications equipment.

<table>
<thead>
<tr>
<th>Rated maximum output power of transmitter W</th>
<th>Separation distance according to frequency of transmitter m</th>
</tr>
</thead>
<tbody>
<tr>
<td></td>
<td>150 kHz to 80 MHz</td>
</tr>
<tr>
<td></td>
<td>(d = \frac{3.5}{V1}\sqrt{\frac{P}{E}})</td>
</tr>
<tr>
<td>.01</td>
<td>.116</td>
</tr>
<tr>
<td>.1</td>
<td>.368</td>
</tr>
<tr>
<td>1</td>
<td>1.16</td>
</tr>
<tr>
<td>10</td>
<td>3.69</td>
</tr>
<tr>
<td>100</td>
<td>11.66</td>
</tr>
</tbody>
</table>

For transmitters rated at a maximum output power not listed above, the recommended separation distance \(d\) in metres (m) can be estimated using the equation applicable to the frequency of the transmitter, where \(P\) is the maximum output power rating of the transmitter in watts (W) according to the transmitter manufacturer.

**NOTE 1** At 80 MHz and 800 MHz, the separation distance for the higher frequency range applies.

**NOTE 2** These guidelines may not apply in all situations. Electromagnetic propagation is affected by absorption and reflection from structures, objects and people.

Disposable Administration Set Specifications

<table>
<thead>
<tr>
<th>D-50 / DI-50</th>
<th>170 Micron</th>
<th>56 ml</th>
</tr>
</thead>
<tbody>
<tr>
<td>Filter Size</td>
<td>170 Micron</td>
<td></td>
</tr>
<tr>
<td>Priming Volume</td>
<td>56 ml</td>
<td></td>
</tr>
<tr>
<td>System 1200 Normothermic (35 – 41°C)</td>
<td></td>
<td></td>
</tr>
<tr>
<td>Fluid Delivery Range</td>
<td></td>
<td></td>
</tr>
<tr>
<td>10°C Input:</td>
<td>40 ml/min. to 300 ml/min.</td>
<td></td>
</tr>
<tr>
<td>20°C Input:</td>
<td>40 ml/min. to 400 ml/min.</td>
<td></td>
</tr>
<tr>
<td>Maximum Flow Rate</td>
<td>500 ml/min.</td>
<td></td>
</tr>
<tr>
<td>(crystalloid, 300 mmHg, 14g catheter)</td>
<td></td>
<td></td>
</tr>
</tbody>
</table>

<table>
<thead>
<tr>
<th>D-60 / DI-60HL</th>
<th>170 Micron</th>
<th>74 ml</th>
</tr>
</thead>
<tbody>
<tr>
<td>Filter Size</td>
<td>170 Micron</td>
<td></td>
</tr>
<tr>
<td>Priming Volume</td>
<td>74 ml</td>
<td></td>
</tr>
<tr>
<td>System 1200 Normothermic (35 – 41°C)</td>
<td></td>
<td></td>
</tr>
<tr>
<td>Fluid Delivery Range</td>
<td></td>
<td></td>
</tr>
<tr>
<td>10°C Input:</td>
<td>75 ml/hour to 530 ml/min.</td>
<td></td>
</tr>
<tr>
<td>20°C Input:</td>
<td>75 ml/hour to 530 ml/min.</td>
<td></td>
</tr>
<tr>
<td>Maximum Flow Rate</td>
<td>530 ml/min.</td>
<td></td>
</tr>
<tr>
<td>(crystalloid, 300 mmHg, 8.5 F catheter)</td>
<td></td>
<td></td>
</tr>
</tbody>
</table>
### Disposable Administration Set Specifications (continued)

<table>
<thead>
<tr>
<th>Model</th>
<th>Filter Size</th>
<th>Priming Volume</th>
<th>System</th>
<th>Fluid Delivery Range</th>
<th>Maximum Flow Rate</th>
</tr>
</thead>
<tbody>
<tr>
<td>D-70 / DI-70</td>
<td>170 Micron</td>
<td>70 ml</td>
<td>1200 Normothermic (35 – 41°C)</td>
<td>10°C Input: 50 ml/min. to 525 ml/min.</td>
<td>590 ml/min.</td>
</tr>
<tr>
<td></td>
<td></td>
<td></td>
<td>Fluid Delivery Range</td>
<td>20°C Input: 30 ml/min. to 525 ml/min.</td>
<td></td>
</tr>
<tr>
<td></td>
<td></td>
<td></td>
<td>Maximum Flow Rate</td>
<td></td>
<td>(crystalloid, 300 mmHg, 14g catheter)</td>
</tr>
<tr>
<td></td>
<td></td>
<td></td>
<td><strong>Fluid Delivery Range</strong></td>
<td></td>
<td></td>
</tr>
<tr>
<td></td>
<td></td>
<td></td>
<td>10°C Input:</td>
<td>30 ml/min. to 650 ml/min.</td>
<td></td>
</tr>
<tr>
<td></td>
<td></td>
<td></td>
<td>20°C Input:</td>
<td>30 ml/min. to 950 ml/min.</td>
<td></td>
</tr>
<tr>
<td></td>
<td></td>
<td></td>
<td>Maximum Flow Rate</td>
<td>950 ml/min.</td>
<td></td>
</tr>
<tr>
<td></td>
<td></td>
<td></td>
<td>(crystalloid, 300 mmHg, 8.5 F catheter)</td>
<td></td>
<td></td>
</tr>
<tr>
<td>D-100 / DI-100</td>
<td>170 Micron</td>
<td>65 ml</td>
<td>1200 Normothermic (35 – 41°C)</td>
<td>10°C Input: 30 ml/min. to 650 ml/min.</td>
<td>950 ml/min.</td>
</tr>
<tr>
<td></td>
<td></td>
<td></td>
<td>Fluid Delivery Range</td>
<td>20°C Input: 30 ml/min. to 950 ml/min.</td>
<td>(crystalloid, 300 mmHg, 8.5 F catheter)</td>
</tr>
<tr>
<td></td>
<td></td>
<td></td>
<td>Maximum Flow Rate</td>
<td>1400 ml/min.</td>
<td></td>
</tr>
<tr>
<td></td>
<td></td>
<td></td>
<td></td>
<td>(crystalloid, 300 mmHg, 8.5 F catheter)</td>
<td></td>
</tr>
</tbody>
</table>
## Symbols

<table>
<thead>
<tr>
<th>Symbol</th>
<th>Meaning</th>
</tr>
</thead>
<tbody>
<tr>
<td>Type BF Equipment</td>
<td></td>
</tr>
<tr>
<td>Protected Against Dripping Water</td>
<td></td>
</tr>
<tr>
<td>Catalog Number</td>
<td></td>
</tr>
<tr>
<td>Serial Number</td>
<td></td>
</tr>
<tr>
<td>Part Number</td>
<td></td>
</tr>
<tr>
<td>Batch Code</td>
<td></td>
</tr>
<tr>
<td>Authorized Representative in the European Community</td>
<td></td>
</tr>
<tr>
<td>Manufacturer</td>
<td></td>
</tr>
<tr>
<td>Date of Manufacture</td>
<td></td>
</tr>
<tr>
<td>Quantity</td>
<td></td>
</tr>
<tr>
<td>Protective Earth [Ground]</td>
<td></td>
</tr>
<tr>
<td>Alternating Current</td>
<td></td>
</tr>
<tr>
<td>Do Not Reuse</td>
<td></td>
</tr>
<tr>
<td>Attention, see instructions for use</td>
<td></td>
</tr>
</tbody>
</table>
### Symbols

<table>
<thead>
<tr>
<th>Symbol</th>
<th>Meaning</th>
</tr>
</thead>
<tbody>
<tr>
<td>![Triangle]</td>
<td>Electrical Shock Hazard</td>
</tr>
<tr>
<td>![Latex]</td>
<td>Latex Free</td>
</tr>
<tr>
<td>![Sterilized]</td>
<td>Sterilised using ethylene oxide</td>
</tr>
<tr>
<td>![Rx]</td>
<td>Caution: Federal law (U.S.A.) restricts this device to sale by or on the order of a physician.</td>
</tr>
<tr>
<td>![Class 1]</td>
<td>Device is a class type 1 equipment.</td>
</tr>
<tr>
<td>![Protective Earth Terminal]</td>
<td>Protective earth terminal</td>
</tr>
<tr>
<td>![Instruction]</td>
<td>Consult instructions for use</td>
</tr>
<tr>
<td>![Do Not Use]</td>
<td>Do not use if package is damaged.</td>
</tr>
<tr>
<td>![Temperature Limitation]</td>
<td>Temperature limitation</td>
</tr>
<tr>
<td>![Use By]</td>
<td>Use by</td>
</tr>
<tr>
<td>![Recyclable Product]</td>
<td>Recyclable Product</td>
</tr>
<tr>
<td>![Pressure Gauge]</td>
<td>Pressure Gauge</td>
</tr>
<tr>
<td>![Pressurize]</td>
<td>Pressurize</td>
</tr>
<tr>
<td>![Unpressurize]</td>
<td>Unpressurize</td>
</tr>
<tr>
<td>![Do Not Bend]</td>
<td>Do not bend heat exchanger</td>
</tr>
</tbody>
</table>
### Symbols

<table>
<thead>
<tr>
<th>Symbol</th>
<th>Meaning</th>
</tr>
</thead>
<tbody>
<tr>
<td><img src="image" alt="Alarm Test" /></td>
<td>Alarm Test</td>
</tr>
<tr>
<td><img src="image" alt="ON MAINS are connected" /></td>
<td>ON — Only for a part of the equipment. MAINS are connected</td>
</tr>
<tr>
<td><img src="image" alt="OFF MAINS are still connected" /></td>
<td>OFF — Only for a part of the equipment. The MAINS are still connected</td>
</tr>
<tr>
<td><img src="image" alt="Automatic Operation" /></td>
<td>Automatic Operation</td>
</tr>
<tr>
<td><img src="image" alt="Recirculating Solution Temperature" /></td>
<td>Recirculating Solution Temperature</td>
</tr>
<tr>
<td><img src="image" alt="Over Temperature Test" /></td>
<td>Over Temperature Test (Recirculating Solution Over Temperature)</td>
</tr>
<tr>
<td><img src="image" alt="Add Recirculating Solution" /></td>
<td>Add Recirculating Solution</td>
</tr>
<tr>
<td><img src="image" alt="Check Disposables, Check Tubing" /></td>
<td>Check Disposables, Check Tubing</td>
</tr>
<tr>
<td><img src="image" alt="Clamped" /></td>
<td>Clamped</td>
</tr>
<tr>
<td><img src="image" alt="Maximum Reservoir Level" /></td>
<td>Maximum Reservoir Level</td>
</tr>
<tr>
<td><img src="image" alt="Minimum Reservoir Level" /></td>
<td>Minimum Reservoir Level</td>
</tr>
<tr>
<td><img src="image" alt="Device has been tested by National Technical Systems, a nationally recognized technical lab, to meet U.S. requirements for safety." /></td>
<td>Device has been tested by National Technical Systems, a nationally recognized technical lab, to meet U.S. requirements for safety.</td>
</tr>
<tr>
<td><img src="image" alt="CE Mark and Notified Body number (0473 indicates AMTAC)" /></td>
<td>CE Mark and Notified Body number (0473 indicates AMTAC)</td>
</tr>
<tr>
<td><img src="image" alt="Humidity Limitation" /></td>
<td>Humidity Limitation</td>
</tr>
<tr>
<td><img src="image" alt="Collect separately for electrical and electronic equipment." /></td>
<td>Collect separately for electrical and electronic equipment.</td>
</tr>
</tbody>
</table>
A
Add Recirculating Solution indicator 23
Add Recirculating Solution Mode 32
After Use 49
Air Detected Alarm mode
Clear 47
Detected 47
Air Detected/Clamped Mode 35
Air Detection/Clamp
Description 3
Principles of operation 19
Air Detector/Clamp Control Panel 24
Alarm Test Mode 31
Alarms
Add Recirculating Solution 32
Air Detected 47
Air detected/Clamped 35
Check Disposables 32
Check Tubing 34
Over Temperature 33
Summary table 28
Assembly instructions 9
Disinfect recirculating solution reservoir 13
Fluid Warmer 9
H-31, B Air Detector/Clamp 16
Automatic Operation Air Detector/Clamp 34
Automatic Operation indicator
Air Detector/Clamp 23
Fluid Warmer 23
B
Button functions 22
C
Calibration of recirculating solution temperature 55
Change O-Rings 59
Change recirculating solution
Distilled water 58
0.3% hydrogen peroxide/distilled water solution 59
Check Disposables indicator 23
Check Disposables Mode 32
Check Tubing indicator 24
Check Tubing Mode 34
Clamped indicator 24
Clean fan filter 59
Clean the exterior 57
Components list 10
Contraindications 5
Controls and displays 21
D
Description 3
Air Detector/Clamp 3
Disposable Administration Set 3
Fluid Warmer 3
Pressure Chambers 3
Disinfect recirculating solution reservoir 58
During assembly 13
Disposable Administration Set Description 3
Install 38
Prime 39
Disposal information 64
Drain valve 11
DSTA-040 calibration test 54
E
Electrical safety tests 17
Electrical testing 56
Electromagnetic Environment Recommendations 65
F
Flanking brackets 12
Fluid Bag
Change 43
Fluid Warmer
Description 3
Principles of operation 19
Fluid Warmer
Description 3
Use of 44
Fluid Warmer Alarm Signal Test button 22
Fluid Warmer Display Panel 23
G
Ground continuity 56
H
H-31, B Air Detector/Clamp
Assembly instructions 16
I
Indications for use 4
Interlocks for Disposable Set installation 26
L
Log
Maintenance and calibration 60
Lubricate O-Ring seals 57
M
Maintenance 57
Every 12 months 58
Every 30 days 57
Log/Schedule 60
Prior to every use 57
Modes of Operation 29
Add Recirculating Solution Mode 32
Air Detected/Clamped Mode 35
Alarm Test Mode 31
Automatic Operation Air Detector/Clamp 34
Check Disposables Mode 32
Check Tubing Mode 34
OFF Mode 30
ON/Automatic Operation for Fluid Warmer 30
Over Temperature Alarm Mode 33
Over Temperature Test Mode 31
Power ON Test for Air Detector/Clamp 33
Pressurized Mode 35
Unpressurized Mode 36
O
OFF Mode 30
ON/Automatic Operation for Fluid Warmer 30
Operating Instructions 37
Operation Display/Panel summary 28
O-Rings
Lubricate 57
Over Temperature Alarm Mode 33
Over Temperature indicator 23
Over Temperature Test button 22
Over Temperature Test Mode 31
P
Panels
Air Detection/Clamp Control 24
Fluid Warmer Display 23
Power and Alarm Test 22
Pressure Chamber control 25
Performance testing 54
Pneumatic tubing 14
Power and Alarm Test Panel 22
Power OFF button 22
Power ON button 22
Power ON Test for Air Detector/Clamp 33
Pressure Chamber Control Panel 25
Pressure Chambers
Description 3
Install 12
Load for use 44
Pressurize for use 44
Principles of operation 19
Index

Pressure gauge 25
Pressurize Pressure Chambers 44
Pressurize/Unpressurize lever 25
Pressurized Mode 35
Prime
    Disposable Administration Set 39
    Patient Line 41
Principles of operation 19

R
Recirculating solution temperature calibration 54
Recirculating Solution Temperature display 23
Replace Gas Vent/Filter Assembly 46
Reservoir Capacity 22
Reservoir level display 23

S
Safety information 5
Service information 62
    contacts 63
Set Up for use 38
Specifications 64
    Disposable Set 65
    system 64
Symbols 67

T
TEMPCHECK calibration test 54
Temperature Display 23, 33
Test Air Detector/Clamp 42
Test Fluid Warmer Alarms 41
Testing 52
    Add Recirculating Solution Alarm 52
    Air Detector/Clamp 42
Check Disposables Alarm 52
    Electrical 56
    Fluid Warmer Alarm 53
    Ground continuity 56
    Maintenance requirement 59
    Over Temperature test 53
    Performance 54
Troubleshooting
    General 50
    Slow flow rate 51

U
Unpressurized Mode 36

W
Warranty information 61

H-1200 FAST FLOW FLUID WARMER | Operator’s Manual 71