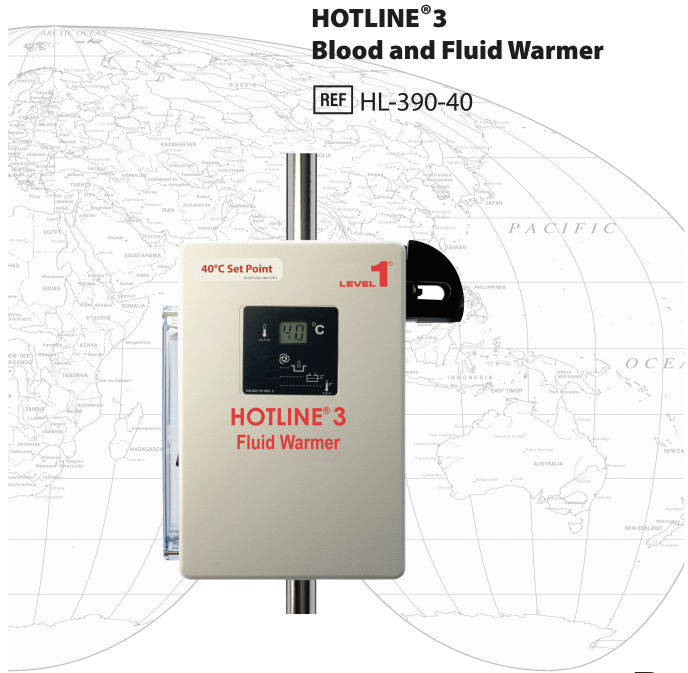
OPERATOR'S MANUAL

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Rx ONLY

smiths medical



HOTLINE[®]3 Blood and Fluid Warmer

REF HL-390-40

OPERATOR'S MANUAL

PN 4534014EN Rev. 002

smiths medical

HOTLINE®3 Blood and Fluid Warmer

Part Number: 4534014EN Rev. 002 (2012-05)

This revision supercedes all previous revisions.

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Contents

1	About this Manual	1
	Indications for Use	1
	Conventions Used in this Manual	1
2	Description	2
	Components	2
	HOTLINE®3 Fluid Warming Set	4
3	Important Safety Information	5
	Additional WARNINGS and CAUTIONS for Accessories	8
4	Assembly Instructions	9
•	Step 1 - Unpack the HOTLINE [®] 3 Warmer	9
	Step 2 - Clamp the HOTLINE®3 Warmer to the I.V. Pole	9
	Step 3 - Disinfect the Reservoir	10
	Step 4 - Fill the Reservoir With Recirculating Solution	11
_	Step 5 - Perform the Electrical Safety Tests	11
5	Principle of Operation	12
_	Infusate Delivery Temperatures	12
6	Operation	13
	Controls and Displays	13
	Display Panel	13
	Power and Alarm Test Panel	14
	Reservoir Level Display Modes of Operation	14 15
	OFF Mode	15
	ON/Operating Mode	15
	Check Disposables Mode	15
	Add Recirculating Solution Mode	16
	Over Temperature Alarm Mode	16
7	Operating Instructions	17
	Step 1 - Set Up the HOTLINE $^{\circledR}$ 3 Warmer	17
	Step 2 - Set Up the HOTLINE $^{ m ext{ iny B}}$ 3 Fluid Warming Set	18
	Step 3 - Connect the Intravenous Administration Set	19
	Step 4 - Using the HOTLINE®3 Warmer	20
	Step 5 - After Use Storage	20 21
8	Troubleshooting	22
9		
9	Testing	24
	Alarm Signal Test	24
	Over Temperature Alarm Test Add Recirculating Solution Test	25
	Check Disposables Test	25 26
	Temperature Verification of the Recirculating Solution	26
	Periodic Electrical Testing	28
	Leakage Current	28

	Ground Bond Test	28
10	Maintenance	29
	Maintenance Performed with Every Use	29
	Clean the Exterior	29
	General Inspection	31
	Maintenance Performed Every 30 Days	31
	Disinfect the Reservoir and Change Recirculating Solution for	
	Distilled Water and 35% Isopropyl Alcohol Solution	31
	Maintenance Performed Every 12 Months	31
	Disinfect the Reservoir and Change Recirculating Solution for	
	0.3% Hydrogen Peroxide Solution	31
	Disinfect the Reservoir and Change the Recirculating Solution	31
	Disinfect the Reservoir	32 32
	Add Recirculating Solution Testing HOTLINE®3 Warmer Operation	33
	Maintenance Log	34
11	Limited Warranty	35
	•	03
12	Service	37
	Non-Warranty Work	37
	Additional Documentation	37
	Disposal Information	38
	Service Contacts	38
	USA/Canada	38
13	Specifications and Accessories	39
	System Specifications	39
	Electromagnetic Compliance	40
	Electromagnetic Environmental Recommendations	41
	Accessories	41
14	Symbols	42

About this Manual

This operator's manual describes the assembly, use, and maintenance of the HOTLINE® 3 Blood and Fluid Warmer. This manual is intended for use by individuals trained in the healthcare and biomedical professions.

WARNING: These instructions contain important information for safe use of the product. Read the entire contents of this operator's manual, including Warnings and Cautions, before using this product. Failure to properly follow warnings, cautions, and instructions could result in death or serious injury to the patient.

Indications for Use

The HOTLINE \$\mathbb{B}\$ 3 Blood and Fluid Warmer is designed for use with the HOTLINE \$\mathbb{B}\$ 3 Fluid Warming Set to warm blood and intravenous (I.V.) fluids and deliver them to the patient's intravenous access site at normothermic temperatures under gravity flow conditions. The HOTLINE \$\mathbb{B}\$ 3 Warmer is intended for use by trained medical personnel to provide routine flow of warmed I.V. fluid.

Conventions Used in this Manual

- The HOTLINE®3 Blood and Fluid Warmer will be referred to as the HOTLINE®3 Warmer.
- The HOTLINE[®]3 L-370 Fluid Warming Set will be referred to as the HOTLINE[®]3 Fluid Warming Set or Disposable Set.

Convention	Description
CONTRAINDICATION	A Contraindication statement alerts the user to conditions when the device should not be used.
WARNING	A Warning statement alerts the user to conditions that may cause death or serious injury to the patient or user.
CAUTION	A Caution statement alerts the user to conditions that may cause malfunction, failure, or damage to the device.

Note: Some HOTLINE[®]3 Warmers may have a 'Converted from HL-90 to HL-390' label on the top of the device. This label indicates that a $HOTLINE^{®}$ HL-90 Warmer was upgraded to a $HOTLINE^{®}$ 3 HL-390 Warmer.

Description

The HOTLINE[®]3 Warmer delivers blood and intravenous fluid at normothermic temperatures by surrounding the sterile intravenous line with a layer of warmed recirculating solution. An onboard recirculating solution supply is heated to 39.5°C \pm 0.5 and circulated through the outer lumen of the HOTLINE[®]3 Fluid Warming Set, which surrounds the intravenous line.

The HOTLINE[®]3 Warmer employs a safe, recirculating solution heating system, inherently free of "hot spots," to actively warm the patient line. Electronic circuitry continuously monitors the recirculating solution temperature. The primary temperature control circuit limits the recirculating solution to 40°C maximum. In the unlikely event of a malfunction of this circuit, a second "watchdog" circuit will visually and audibly alarm and stop the recirculating solution pump if the temperature reaches 41.2°C. Fluid in the HOTLINE[®]3 Fluid Warming Set is never exposed to any damaging or dangerous temperatures while the HOTLINE[®]3 Warmer is operating.

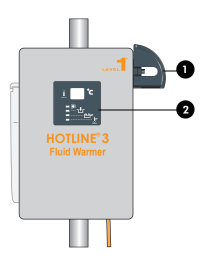
The recirculating solution temperature and visual alarms are indicated on the Display Panel on the front of the HOTLINE®3 Warmer. A green Operating light illuminates on this panel when the HOTLINE®3 Warmer is set up and operating correctly.

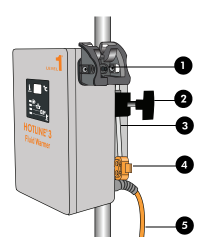
Components

The HOTLINE®3 components are called-out in the following series of figures.

Front View

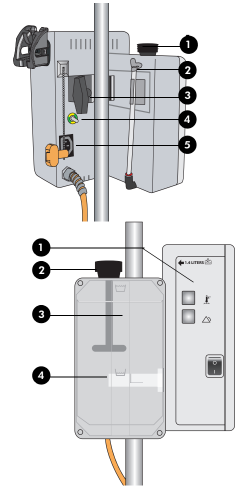
- 1 Socket for HOTLINE®3 Fluid Warming Set with the reflux plug in place
- 2 Display Panel





Right Side View

- **1** Socket with the reflux plug removed
- 2 Clamp for I.V. pole
- **3** Drain tube in tube holder
- 4 Reflux plug
- **5** Power cord



Rear View

- **1** Fill-port plug
- **2** Drain tube in tube holder
- 3 Clamp for I.V. pole
- **4** Protective earth terminal
- **5** Auxiliary electrical outlet (uncovered)

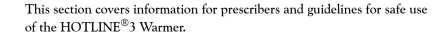
Left Side View

- 1 Power and Alarm Test Panel
- **2** Fill-port plug
- **3** Reservoir, contains recirculating solution
- **4** Float switch (inside reservoir)

HOTLINE®3 Fluid Warming Set

HOTLINE[®]3 L-370 Fluid Warming Set is an individually packed, single-use disposable with a Sterile Fluid Path. The priming volume is approximately 20 ml. The HOTLINE[®]3 Fluid Warming Set has a connector that plugs into the socket on the right side of the HOTLINE[®]3 Warmer. The connector is designed with a support guide to facilitate installation and two locking tabs to ensure that HOTLINE[®]3 Fluid Warming Set is securely fastened in the socket. This is the only connection necessary to provide the warming function. The HOTLINE[®]3 Fluid Warming Set is easily unplugged from the HOTLINE[®]3 Warmer and discarded.

Important Safety Information



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.

- These instructions contain important information for safe use of the product. Read the entire contents of this operator's manual, including Warnings and Cautions, before using this product.
 Failure to properly follow warnings, cautions, and instructions could result in death or serious injury to the patient.
- The HOTLINE®3 Fluid Warming Set, L-10, PC-8, and YC-8 are single-use devices and are not intended for re-sterilization.
- Do not use HOTLINE®3 Fluid Warming Set, L-10, PC-8, and YC-8 if the caps are not securely in place, else the I.V. flow path may not be sterile.
- The HOTLINE[®]3 Warmer is for use only with Smiths Medical supplied or approved parts, accessories, and Disposable Sets.
 The device may not function as intended with the use of unapproved parts, accessories, or Disposable Sets.
- 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.
- Set-up, priming, and use require aseptic technique as per applicable institutional policies and procedures.

Continued

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WARNINGS

- Prime the recirculating solution path before connecting to the intravenous extension set. This is to confirm that there is not a breach between the recirculating solution path and intravenous path. If fluid exits the patient end of the HOTLINE[®]3 Fluid Warming Set before connecting to the intravenous administration set, remove and replace HOTLINE[®]3 Fluid Warming Set.
- Remove all air from the HOTLINE®3 Fluid Warming Set, L-10, PC-8, and YC-8 before connecting to the patient. Failure to do so may result in introduction of air to the patient.
- To reduce the risk of outgassed microbubbles entering patient vasculature, an L-10 Gas Vent may be used with the HOTLINE®3 Fluid Warming Set.
- Not for use with pressure devices generating over 300 mmHg.
 Pressure greater than 300 mmHg may compromise the integrity of the HOTLINE®3 Fluid Warming Set.
- Do not stick the HOTLINE®3 Fluid Warming Set with needles, as
 this will breach the I.V. path and compromise the integrity of the
 patient intravenous line. If a Disposable Set with a breached
 recirculating solution path/intravenous path is used, then
 patient illness may occur because of the HOTLINE®3 Warmer's
 recirculating solution entering the patient's bloodstream.
- 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 remove the device from service.
- If any visual indicator does not illuminate or the audible signal does not sound, do not use the HOTLINE®3 Warmer. Remove the device from service immediately.
- Do not operate the HOTLINE®3 Warmer in the presence of a flammable anesthetic mixture with air, oxygen, or nitrous oxide. The risk of explosion exists if the HOTLINE®3 Warmer is operated in a potentially explosive environment.
- Do not use the HOTLINE®3 Warmer in high-energy fields such as: MRI, X-RAY, portable and mobile RF communications equipment, and other such devices. The HOTLINE®3 Warmer may act as a projectile in a strong magnetic field, cause image artifacts, or not function as intended.

Continued

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WARNINGS

- Exposed conductor on MAINS power cord can cause an electrocution hazard. Remove device from service if the MAINS power cord has exposed wires.
- Grounding reliability can only be achieved when the MAINS power cord is connected to a properly grounded receptacle.
 Risk of electrical shock exists if the equipment is not connected to a properly grounded receptacle.
- Do not mount the HOTLINE®3 Warmer more than 107cm (42") above the floor. For convenience, 107cm (42") is indicated on the HOTLINE®3 Warmer power cord by a black mark. Mounting the HOTLINE®3 Warmer above 107cm (42") may result in instability of the pole and tipping.
- Ensure that the HOTLINE®3 Warmer clamp is screwed tightly onto the I.V. pole. Failure to securely mount the HOTLINE®3 Warmer onto the I.V. pole may cause the HOTLINE®3 Warmer to slide down the I.V. pole.
- Do not use the HOTLINE®3 Warmer if equipment or Disposable Set malfunction is evident.
- No user-serviceable parts. All service must be performed by Smiths Medical or competent personnel.
- No modification of this equipment is allowed.

CAUTIONS

Malfunction, failure, or damage to the device may occur if these cautions are not followed.

- Federal (U.S.A.) law restricts this device to sale by or on the order of a physician.
- Do not autoclave or immerse any part of the HOTLINE®3
 Warmer in liquids, which may cause damage and improper functioning.
- Never use organic solvents (e.g., acetone), strong acids, or bases to clean any portion of the HOTLINE®3 Warmer.

Continued

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CAUTIONS

- Do not place the HOTLINE®3 Warmer directly under a faucet or use a faucet sprayer to rinse. Never spray cleaning or other fluids into openings on the HOTLINE®3 Warmer or into the external connectors.
- This device is cooled by convection. Be sure the air vents on the bottom and the back of the device are kept clear.
- Do not fill the HOTLINE®3 Warmer reservoir with a HOTLINE®3
 Fluid Warming Set in place. Failure to remove the HOTLINE®3
 Fluid Warming Set before the fill procedure may result in an air lock in the HOTLINE®3 Warmer.
- Medical devices require specific material characteristics to perform as intended. These characteristics have been verified for single use only. Any attempt to re-process the device for subsequent re-use may adversely affect the integrity of the device or lead to deterioration in performance.

Additional WARNINGS and CAUTIONS for Accessories

WARNINGS for the L-10 Gas Vent

- Do not tape over vents, else air will not be vented.
- Not for use with volumetric infusion pumps, hand pumps, or syringes. These may compromise the integrity of the L-10 Gas Vent or HOTLINE®3 Fluid Warming Set.
- When the L-10 Gas Vent is in use, it should be placed at or below the heart level. Do not raise the gas vent above the patient's heart level. If the gas vent is raised above heart level, air may be entrained into the infusion line, possibly causing air embolism, resulting in serious injury or death.

CAUTIONS for the L-10 Gas Vent

 This product contains natural rubber latex, which may cause allergic reactions.

Assembly Instructions

Read through the instructions completely prior to setting up the $HOTLINE^{\circledR}3$ Warmer.

Step 1 - Unpack the HOTLINE®3 Warmer

- 1 Open the shipping carton and remove the HOTLINE®3 Warmer.
- **2** Check the contents of the package to verify the following components are present:
 - HOTLINE®3 Warmer
 - Operator's Manual
 - HOTLINE[®] 3 Inspection/Test Form
- **3** Examine the HOTLINE[®]3 Warmer for damage. If any components appear damaged, do not use the HOTLINE[®]3 Warmer. Contact Smiths Medical for a replacement.

Note: After unpacking the HOTLINE[®]3 Warmer, recycle packaging material according to hospital policy for recyclable materials.

Step 2 - Clamp the HOTLINE®3 Warmer to the I.V. Pole

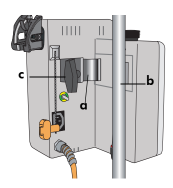
WARNINGS

- Ensure that the HOTLINE®3 Warmer pole clamp is screwed tightly onto the I.V. pole. Failure to securely mount the HOTLINE®3 Warmer onto the I.V. pole may cause the HOTLINE®3 Warmer to slide down the pole and may injure the patient or user.
- Do not mount the HOTLINE®3 Warmer more than 107cm (42") above the floor. For convenience, 107cm (42") is indicated on the HOTLINE®3 Warmer line cord by a black mark. Mounting the HOTLINE®3 Warmer above 107cm (42") may result in instability of the pole and tipping that may injure the patient or user.

CAUTIONS

 This device is cooled by convection. Be sure the air vents on the bottom and the back of the device are kept clear.

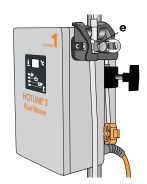
- 1 Slide the clamp (a) on the HOTLINE®3 Warmer over the I.V. pole (b) and tighten the clamp screw (c) firmly.
- **2** Check the tightness of the HOTLINE[®]3 Warmer to ensure it is securely clamped to the pole.

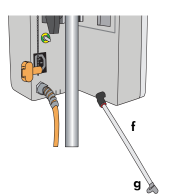


Step 3 - Disinfect the Reservoir

- 1 Prepare a 0.3% hydrogen peroxide solution by mixing 140 ml of 3% hydrogen peroxide solution and 1,260 ml of distilled water.
- **2** Remove the reflux plug from the socket if required, and then remove the fill-port plug (**d**) and fill the reservoir with 1.4 liters of 0.3% hydrogen peroxide solution.
- **3** Replace the fill-port plug (**d**).
- **4** Insert a HOTLINE[®]3 Fluid Warming Set (**e**) into the socket.
- **5** Plug the HOTLINE®3 Warmer into properly grounded power outlet.
- **6** Turn the HOTLINE® 3 Warmer ON and let the solution circulate for a 30-minute disinfection period.
- **7** Turn the HOTLINE®3 Warmer OFF.
- **8** Invert the drain tube (**f**) and place a container under the end of the tube. Remove the end cap (**g**) and drain the recirculating solution into the container.
- **9** When all the recirculating solution has drained from the reservoir, replace the end cap and insert the drain tube back in the holder.
- **10** Remove the HOTLINE[®]3 Fluid Warming Set. Dispose of the HOTLINE[®]3 Fluid Warming Set in a safe manner according to local guidelines for disposal of contaminated medical waste.







Step 4 - Fill the Reservoir With Recirculating Solution

WARNINGS

Do not fill the HOTLINE®3 Warmer reservoir with a HOTLINE®3
 Fluid Warming Set in place. Failure to remove the HOTLINE®3
 Fluid Warming Set before the fill procedure may result in an air lock in the HOTLINE®3 Warmer.

Recirculating Solution Protocols

Use one of the following solutions for the reservoir.

Recirculating Solution	Preparation	Maintenance
0.3% Hydrogen Peroxide Solution	Mix 140 ml of 3% hydrogen peroxide with 1,260 ml of distilled water.	Replace solution and disinfect reservoir every 12 months.
Distilled Water	Use distilled water.	Replace solution and disinfect reservoir every 30 days.
35% Isopropyl Alcohol Solution	Mix 700 ml of 70% isopropyl alcohol with 700 ml of distilled water.	Replace solution and disinfect reservoir every 30 days.

Note: Use distilled water only, not tap water. Failure to do so may cause build-up of mineral deposits in the recirculating solution path, which may impair heater performance.

- **1** Prepare the recirculating solution.
- **2** Remove the fill-port plug (a).
- **3** Fill the reservoir with 1.4 liters of recirculating solution.
- **4** Replace the fill-port plug.

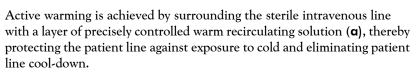
Step 5 - Perform the Electrical Safety Tests

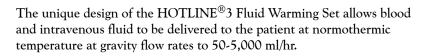
Perform all applicable electrical safety tests as required per institutional procedure. Refer to Section 9, *Testing*, for more information about electrical safety testing.



Principle of Operation

HOTLINE[®]3 Warmer delivers blood and intravenous fluid at normothermic temperatures under routine, gravity flow rates. Conventional fluid warming systems suffer from cool-down between the warmer and the patient connection. HOTLINE[®]3 Warmer overcomes this problem by providing active warming of the patient line all the way to the patient connection.

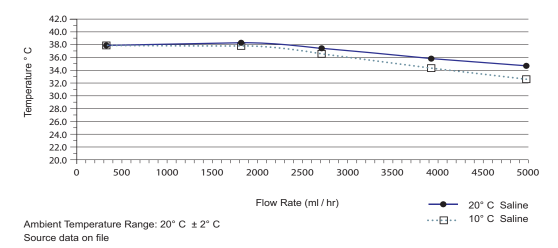


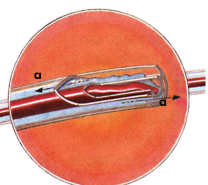




The following table shows the typical infusate delivery temperatures at the patient end of an L-370 HOTLINE®3 Fluid Warming Set.

Note: The setpoint temperature of the recirculating solution is 40.0°C.





Operation

This section describes the controls and displays that monitor and control the HOTLINE[®]3 Warmer, and the modes of operation.

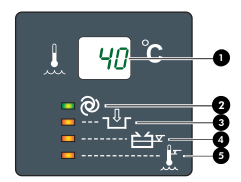
Controls and Displays

- Display Panel
- Power and Alarm Test Buttons
- Reservoir Level Display

Display Panel

The Display Panel is located on the front of the HOTLINE[®]3 Warmer and provides continuous information about the operation of the HOTLINE[®]3 Warmer. A liquid crystal display (LCD) indicates recirculating solution temperature. Just below the LCD, four lightemitting diodes (LEDs) indicate operation modes for the HOTLINE[®]3 Warmer.

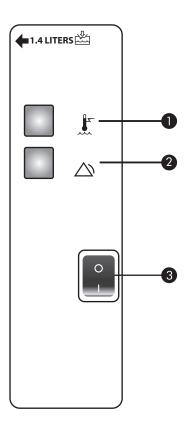
- Recirculating Solution Temperature The temperature is displayed in degrees Celsius.
- **2 ON/Operation** The green LED illuminates when the power is turned on and the HOTLINE®3 Fluid Warming set is properly installed.
- **3** Check Disposables The red LED illuminates and an audible attention signal beeps when the HOTLINE®3 Fluid Warming Set is not properly installed.
- **4** Add Recirculating Solution The red LED illuminates and an audible attention signal beeps when the level in the reservoir is low and additional recirculating solution must be added.
- **5 Over Temperature** The red LED illuminates and an audible warning signal beeps when the recirculating solution is over the acceptable temperature for safe use.



Power and Alarm Test Panel

The Power and Alarm Test Panel is located on the left side of the $HOTLINE^{\circledR}3$ Warmer next to the reservoir. This panel contains two pressure-sensitive buttons that are activated when pressed, and the ON/OFF switch.

- 1 Over Temperature Alarm Test Button The Over Temperature Alarm Test is used to confirm the proper operation of the Over Temperature circuitry.
- **2 Alarm Signal Test Button** The Alarm Signal Test is used to confirm proper operation of the visual and audible alarms.
- **3 Power ON/OFF Switch** The black switch toggles to turn power ON and OFF.



Reservoir Level Display

The reservoir for the recirculating solution is located on the left side of the HOTLINE $^{\circledR}$ 3 Warmer, next to the Power and Alarm Test Panel. The level of the recirculating solution is visible in the reservoir. Two symbols indicate the maximum (α) and minimum (b) solution level requirements.



Modes of Operation

The HOTLINE®3 Warmer operation is defined in the following modes:

- OFF Mode
- ON/Operating Mode
- Check Disposables Mode
- Add Recirculating Solution Mode
- Over Temperature Alarm Mode

The description of each mode includes a definition of the mode, activation and/or monitoring of the mode, mode characteristics, and method to clear the mode state.



OFF Mode

The power switch is in the OFF position (α) and the HOTLINE[®]3 Warmer is turned off.



ON/Operating Mode

The power switch is in the ON position (**b**) and the HOTLINE[®]3 Fluid Warming Set has been properly installed.



Mode Characteristics

- The green Operating LED (c) illuminates.
- The recirculating solution temperature display will begin to increase.
- The recirculating solution path in the HOTLINE®3 Fluid Warming Set will automatically prime.



Check Disposables Mode

The Check Disposables mode indicates a missing or improperly installed HOTLINE®3 Fluid Warming Set.

Mode characteristics

- The green Operating LED on the Display Panel turns off.
- The red Check Disposables LED (d) on the Display Panel illuminates.
- The audible alarm sounds and repeats approximately every two seconds.

• The recirculating solution stops circulating.

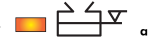
To clear this mode, check that the connector on the HOTLINE®3 Fluid Warming Set is firmly inserted in the socket.

Add Recirculating Solution Mode

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

Mode characteristics

- The green Operating LED on the Display Panel turns off.
- The red Add Solution LED (α) on the Display Panel illuminates.



- The audible alarm sounds and repeats approximately every two seconds.
- The recirculating solution stops circulating.

To clear this mode, add recirculating solution to the reservoir.

Over Temperature Alarm Mode

The Over Temperature Alarm mode indicates that the temperature of the recirculating solution is at or above 42.2°C.

Mode characteristics

- The green Operating LED on the Display Panel turns off.
- The red Over Temperature LED (**b**) on the Display Panel illuminates.



- The audible alarm sounds and repeats approximately every two seconds.
- The recirculating solution stops circulating.

For instructions to clear this mode, see Section 8, Troubleshooting.

Operating Instructions

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

WARNINGS

- Set-up, priming, and use require aseptic technique as per applicable institutional policies and procedures. Death or serious injury may occur to the patient or user if this warning is not followed.
- 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.
- Do not fill the HOTLINE®3 reservoir with a HOTLINE®3 Fluid Warming Set in place. Failure to remove the HOTLINE®3 Fluid Warming Set before the fill procedure may result in an air lock in the HOTLINE®3 Warmer.

Step 1 - Set Up the HOTLINE®3 Warmer



- 1 Check that the level is above the minimum level mark (a) on the reservoir. Add recirculating solution to the reservoir through the fill-port if required.
- **2** Check the condition of the HOTLINE[®]3 Warmer with a visual inspection before using. Remove from service any HOTLINE[®]3 Warmer that shows physical damage.
- **3** Plug the HOTLINE®3 Warmer into properly grounded power outlet.

Step 2 - Set Up the HOTLINE®3 Fluid Warming Set

WARNINGS

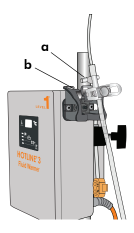
- The HOTLINE®3 Fluid Warming Set is a single-use device and is not intended for re-sterilization. Death or serious injury may occur to the patient or user if this warning is not followed.
- Do not use HOTLINE®3 Fluid Warming Set, L-10, PC-8, and YC-8 if the caps are not securely in place, else the I.V. flow path may not be sterile and may cause death or serious injury.
- Prime the recirculating solution path before connecting to the intravenous extension set. This is to confirm that there is not a breach between the recirculating solution path and intravenous path. If fluid exits the patient end of the HOTLINE®3 Fluid Warming Set before connecting to the intravenous extension set, remove and replace HOTLINE®3 Fluid Warming Set. Death or serious injury may occur to the patient or user if this warning is not followed.

To set up the HOTLINE[®]3 Fluid Warming Set, you will need the following:

- HOTLINE®3 Warmer
- Intravenous administration set
- Intravenous fluid or blood
- Extension Set, 20cm (8") or less in length (optional)
- **1** Remove the reflux plug (if present) from the socket on the right side of the HOTLINE[®] 3 Warmer.
- **2** Plug the connector on the HOTLINE®3 Fluid Warming Set (a) into the socket.

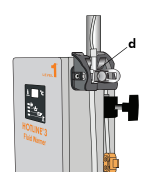
Note: Do not squeeze the tabs when inserting the connector.

a Insert the support guide (**a**) on the connector into the groove (**b**) on the top of the socket.



b Push on the center (**c**) of connector until it clicks in the socket (**d**).





- **3** Turn ON the power switch.
 - The green Operating LED on the Display Panel illuminates.
 - The recirculating solution temperature display will begin to increase.
 - The recirculating solution path in the HOTLINE[®]3 Fluid Warming Set will automatically prime.
- **4** Remove the end cap and inspect the patient end of the HOTLINE®3 Fluid Warming Set for leaks to confirm the integrity of the intravenous pathway.

Step 3 - Connect the Intravenous Administration Set

WARNINGS

- Remove all air from the HOTLINE®3 Fluid Warming Set, L-10, PC-8, and YC-8 before connecting to the patient. Failure to do so may result in introduction of air to the patient, which may contribute to serious patient injury or death.
- Do not stick the HOTLINE®3 Fluid Warming Set with needles, as
 this will breach the I.V. path and compromise the integrity of the
 patient intravenous line. If a Disposable Set with a breached
 recirculating solution path/intravenous path is used, then
 patient illness may occur because of the HOTLINE®3 Warmer's
 recirculating solution entering the patient's blood stream.
 - 1 Connect the I.V. fluid and the intravenous administration set to the HOTLINE®3 Fluid Warming Set.
 - **2** Fully prime the intravenous administration set, the HOTLINE[®]3 Fluid Warming Set, and patient extension set (if used).

3 Connect the distal end of the HOTLINE®3 Fluid Warming Set to the patient's intravenous access site without entrapping air.

Step 4 - Using the HOTLINE®3 Warmer

WARNINGS

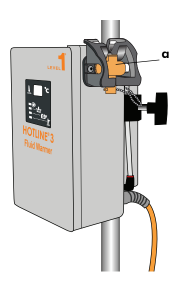
- Activation of the Over Temperature warning signal indicates that warming has stopped and immediate operator intervention is required to clear the over temperature condition or to remove the device from service. Death or serious injury may occur to the patient or user if this warning is not followed.
- If any visual indicator does not illuminate or the audible signal does not sound, do not use the HOTLINE®3 Warmer. Remove the device from service immediately. Death or serious injury may occur to the patient or user if this warning is not followed.
- Not for use with pressure devices generating over 300 mmHg.
 Pressure greater than 300 mmHg may compromise the integrity of the HOTLINE®3 Fluid Warming Set.
 - 1 Wait until the recirculating solution temperature display reaches 39°C, which indicates the HOTLINE®3 Warmer is ready for use.
 - **2** Adjust the rate of I.V. flow using the clamp on the intravenous administration set.

Note: Do not kink the Disposable Set. Do not restrict the circulation of the solution through the tubing.

Step 5 - After Use

WARNINGS

- 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.
 - **1** Turn OFF the power switch.
 - **2** Remove the HOTLINE®3 Fluid Warming Set.
 - **a** Squeeze the tabs on the connector.
 - **b** Lift up and pull the connector away from the socket.



- **3** Insert the reflux plug (a) into the socket.
- **4** After use, handle and dispose of the HOTLINE®3 Fluid Warming Set in a safe manner according to local guidelines for disposal of contaminated medical waste.
- **5** Wipe down the external surfaces of the HOTLINE[®] 3 Warmer with mild liquid detergent soap and warm tap water mixture and a soft cloth or sponge. See Section 10, *Maintenance*, for more details about cleaning and external disinfection.

Storage

Store the HOTLINE[®]3 Warmer in a cool, dry place. Do not expose to extreme temperatures. See Section 13, *Specifications*, for more details.

Troubleshooting

Only competent personnel should perform any routine maintenance and repairs to the $HOTLINE^{\circledR}3$ Warmer.

Problem	Check the following		
No power	 Confirm that the HOTLINE[®]3 Warmer is plugged in properly. Confirm that the power switch is in the ON position. Note: If the HOTLINE[®]3 Warmer is plugged in and the power switch is turned ON, the green or red LED will illuminate. 		
Check Disposables alarm	Confirm that the HOTLINE® Fluid Warming Set is properly installed. 1 Push the connector firmly into the socket on the rig side of the HOTLINE®3 Warmer. Note: Turn OFF the power switch before replacing the HOTLINE®3 Fluid Warming Set. 2 If the alarm is not cleared, replace the HOTLINE®3 Fluid Warming Set. Turn ON the power switch and verify that the alarm has cleared. 3 If the alarm is not cleared, remove the HOTLINE®3 Warmer from service.		
Add Recirculating Solution alarm	Check the level in the reservoir Turn OFF the power switch, remove the HOTLINE®3 Fluid Warming Set if installed, add recirculating solution to the maximum level.		
Over Temperature alarm	 Check the HOTLINE®3 Fluid Warming Set for kinks or other restrictions. Check for air lock: Turn the power switch OFF, remove the HOTLINE®3 Fluid Warming Set, and gently shake HOTLINE®3 Warmer to dislodge air. Plug in the HOTLINE®3 Fluid Warming Set and turn power switch ON. If the alarm is not cleared, remove the HOTLINE®3 Warmer from service and return it for repair or replacement. 		

Problem	Check the following		
Hot cabinet	heck for blocked air vents on the bottom or the back of e HOTLINE [®] 3 Warmer. Note: Room temperature above 40°C may cause the HOTLINE [®] 3 Warmer to shut down and the Over Temperature alarm to activate. In this situation, turn the power switch OFF and allow the HOTLINE [®] 3 Warmer to cool down before returning it to service.		
Recirculating solution leaks at the socket where the HOTLINE®3 Fluid Warming Set plugs into the HOTLINE®3 Warmer	Replace the HOTLINE [®] 3 Fluid Warming Set.		
Electrical interference - receiving or transmitting	 Move the HOTLINE®3 Warmer away from the device in question. Plug the HOTLINE®3 Warmer into a separate electrical circuit. If the problem continues, notify Smiths Medical or your local Smiths Medical distributor. 		

Testing

The HOTLINE[®]3 Warmer should be tested by hospital biomedical personnel prior to placing it in service. All testing and maintenance should be performed by competent personnel. If competent personnel are not available, contact Smiths Medical or your local Smiths Medical distributor.

If the HOTLINE[®]3 Warmer and any installed accessories do not pass any of the listed tests, discontinue use of the HOTLINE[®]3 Warmer and remove from service. Contact Smiths Medical or your local Smiths Medical distributor.

WARNINGS

 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.

Note: Alarm testing requires a HOTLINE®3 Fluid Warming Set to be installed and that the HOTLINE®3 Warmer be turned ON and in the Operating mode.

Alarm Signal Test

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



- 1 Press and hold the Alarm Test button (a).
- **2** Observe the following:
 - The green Operating LED turns off.
 - Three red LEDs (Check Disposables, Add Solution, and Over Temperature) illuminate.
 - The audible alarm sounds and repeats approximately every two seconds.



Over Temperature Alarm Test

The HOTLINE[®]3 Warmer should be running at an operating temperature of approximately 39°C to 40°C.

- **1** Press and hold the Over Temperature Alarm Test button (a).
- **2** Observe the following:
 - The recirculating solution Over Temperature Alarm activates at 41°C.
 - The green Operating LED turns off.
 - The red Over Temperature LED (b) illuminates.
 - The audible alarm sounds and repeats approximately every two seconds.
- **3** Stop pressing the Over Temperature Alarm Test button to stop the test.



Add Recirculating Solution Test

The HOTLINE®3 Warmer is equipped with a float switch, which senses the recirculating solution level in the reservoir. When the recirculating solution is too low, the Add Recirculating Solution Alarm will activate.

- **1** Remove the fill-port plug (**c**) on the reservoir.
- **2** Gently depress the float switch (**d**). (This action will simulate the low solution condition.)

Note: Use a non-metal tool to depress the float switch because the float switch contains a magnet.



- **3** Observe the following:
 - The green Operating LED turns off.
 - The red Add Recirculating Solution LED (e) illuminates.
 - The audible alarm sounds and repeats approximately every two seconds.

Check Disposables Test

An interlock switch/sensor, located in the socket on the right side of the HOTLINE[®]3 Warmer, senses a properly installed HOTLINE[®]3 Fluid Warming Set. When the switch does not sense a HOTLINE[®]3 Fluid Warming Set, the Check Disposables alarm activates.

- 1 Slowly remove the HOTLINE®3 Fluid Warming Set (a) from the HOTLINE®3 Warmer socket.
- **2** Observe the following actions:
 - The green Operating LED turns off.
 - The red Check Disposables LED (b) illuminates.
 - The audible alarm sounds and repeats approximately every two seconds.

Note: In any alarm condition, the pump should not be running. A small amount of solution dripping from the disconnection is normal and should stop in a few seconds.

HOTUNE'S Plus Welling

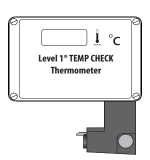


Temperature Verification of the Recirculating Solution

Use the Level 1^{\otimes} TEMP CHECK Thermometer (HLTA-390) to verify the displayed recirculating solution temperature. Other methods of temperature verification may be inaccurate.

TEMP CHECK provides an accurate reading of the highest temperature of the recirculating solution. Because the temperature of the reservoir is typically 0.5°C to 2.0°C lower than the temperature from the heater, and the temperature of the recirculating solution begins to drop due to the effect of ambient temperature on the HOTLINE®3 Fluid Warming Set, the highest temperature of the solution is just after it leaves the heater. During the temperature verification test, the TEMP CHECK is positioned on the right side of the HOTLINE®3 Warmer attached to the socket and senses the solution just after it leaves the heater and before it enters the HOTLINE®3 Fluid Warming Set.

Refer to the TEMP CHECK HLTA-390 Thermometer Operator's Manual for complete Temperature Verification and Calibration Instructions.



To verify the recirculating solution temperature, you will need the following:

- TEMP CHECK (HLTA-390)
- HOTLINE®3 Warmer

To Verify the Recirculating Solution Temperature:

- 1 Plug the HOTLINE[®] 3 Warmer into a power outlet.
- **2** Place the TEMP CHECK on the top right corner of the HOTLINE[®]3 Warmer and plug it into the socket on the right side of the HOTLINE[®]3 Warmer.
- **3** Remove the black label from the auxiliary outlet on the back of the HOTLINE[®]3 Warmer and plug in the TEMP CHECK power cord.

Note: The auxiliary outlet is for use only with Smiths Medical accessories.

- **4** Turn ON the HOTLINE[®] 3 Warmer. Allow 15 minutes for the temperature to stabilize.
- **5** If the TEMP CHECK display indicates a temperature between 39°C and 40°C, and the HOTLINE®3 Warmer display equals the TEMP CHECK display, recirculating solution verification is complete. Refer to the TEMP CHECK Manual for OVERTEMP ALARM verification.
- **6** If the TEMP CHECK display does not indicate a temperature between 39°C and 40°C, refer to the TEMP CHECK Manual for calibration instructions.

Periodic Electrical Testing

Electrical Safety Tests must be performed by competent personnel authorized by the institution to perform such testing. The Safety Tests must be performed and documented at least once per year, or according to institutional policy. These tests include but are not limited to:

- Leakage current
- Ground bond test

Leakage Current

Leakage current must be tested according to methods and pass/fail criteria described in UL 2601-1 or EN 60601-1. Leakage current must be performed with the heater circuit in the full ON condition. To achieve this condition, perform the test when the reservoir is at room temperature. When the HOTLINE®3 Warmer is first turned on and the temperature is rapidly rising, but still below 39°C, the heater circuit is in a full ON condition.

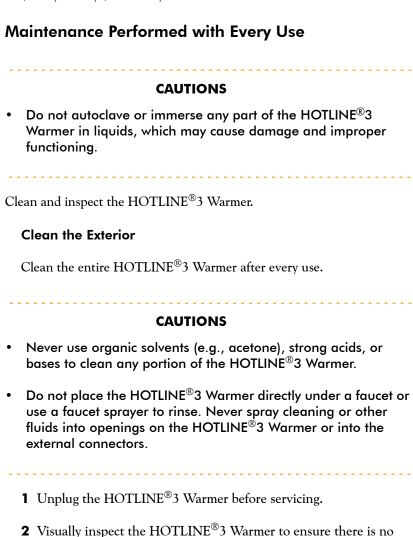
Note: The HOTLINE®3 Warmer is equipped with sensing interlocks. A HOTLINE®3 Fluid Warming Set is required to correctly operate the HOTLINE®3 Warmer and perform leakage current testing. Do not defeat the sensing interlocks or try to operate the HOTLINE®3 without a HOTLINE®3 Fluid Warming Set in place.

Ground Bond Test

Ground bond test must be tested according to methods and pass/fail criteria described in UL 2601-1 or EN 60601-1.

Maintenance

Only competent personnel should perform any routine maintenance and repairs to the HOTLINE®3 Warmer. Maintenance is scheduled with each use, every 30 days, and every 12 months. The tasks are described below.



- visible damage or deterioration of the enclosure such as cracks, or deterioration of the labels and power cord. Do not clean if there is a
 - defect. Contact Smiths Medical or your local Smiths Medical distributor.

- **3** Immerse a soft cloth or sponge as an applicator into the cleaning solution consisting of mild liquid detergent soap and warm tap water mixture. Squeeze out excess solution so that the applicator is not dripping. Wipe or scrub the entire surface of the enclosure and control panels. Use a soft brush to clean the power cord if necessary.
- **4** Rinse a separate soft cloth or sponge in room temperature running potable water. Squeeze out excess water so that the applicator is not dripping. Wipe all of the aforementioned surfaces. Repeat rinsing the cloth or sponge several times with fresh running water during this process to insure all visible residue is removed.
- **5** Dry the item with a hand towel or soft cloth.
- **6** Visually inspect the HOTLINE®3 Warmer and its components to insure that they have been thoroughly cleaned. Repeat cleaning procedure if necessary.
- **7** After thoroughly cleaning the HOTLINE®3 Warmer, perform disinfection if required.
- **8** If it is hospital policy to perform disinfection as part of reprocessing, then follow your institution's guidelines for disinfecting of the surfaces of non-critical medical devices. The list below includes low-level disinfectants that are commonly used in the medical community and high-level disinfectants that are claimed by the manufacturer. The effectiveness of these listed disinfectants should be validated using the hospital procedures.

The following disinfectant agents can be used without causing damage to the enclosure:

Low Level Disinfectants:

• fantastik® All Purpose Cleaner

High Level Disinfectants:

- 1.56% Phenol (e.g., Sporicidin®)
- 3.4% Glutaraldehyde (e.g., CIDEX® Plus)
- 10% Bleach solution
- 1% Ammonia solution
- Surface disinfectants compatible with plastic materials.
- **9** Rinsing of the disinfectant residue should be done using a soft cloth or sponge as the applicator.

General Inspection

Check the condition of the HOTLINE $^{\otimes}$ 3 Warmer with a visual inspection before using. Remove from service any HOTLINE $^{\otimes}$ Warmer that shows physical damage.

Maintenance Performed Every 30 Days

Disinfect the Reservoir and Change Recirculating Solution for Distilled Water and 35% Isopropyl Alcohol Solution

Refer to *Disinfect the Reservoir and Change the Recirculating Solution* procedure in this section.

Maintenance Performed Every 12 Months

Disinfect the Reservoir and Change Recirculating Solution for 0.3% Hydrogen Peroxide Solution

Refer to *Disinfect the Reservoir and Change the Recirculating Solution* procedure in this section.

Disinfect the Reservoir and Change the Recirculating Solution

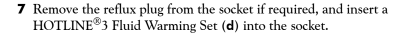
Disinfect the reservoir and change the recirculating solution every 30 days or every 12 months based on the recirculating solution used for the HOTLINE[®]3 Warmer. Refer to the following table for the maintenance schedule.

Recirculating Solution	Preparation	Maintenance
0.3% Hydrogen Peroxide Solution	Mix 140 ml of 3% hydrogen peroxide with 1,260 ml of distilled water.	Replace solution and disinfect reservoir every 12 months.
Distilled Water	Use distilled water.	Replace solution and disinfect reservoir every 30 days.
35% Isopropyl Alcohol Solution	Mix 700 ml of 70% isopropyl alcohol with 700 ml of distilled water.	Replace solution and disinfect reservoir every 30 days.

Note: Use distilled water only, not tap water. Failure to do so may cause build-up of mineral deposits in the recirculating solution path, which may impair heater performance.

Disinfect the Reservoir

- 1 Unplug the HOTLINE®3 Warmer before servicing.
- **2** Remove the drain tube from the holder on the rear of the HOTLINE[®]3 Warmer.
- **3** Invert the drain tube (a) and place a container under the end of the tube. Remove the end cap (b) and drain the recirculating solution into the container.
- **4** When all the recirculating solution has drained from the reservoir, replace the end cap and insert the drain tube back in the holder.
- **5** Prepare a 0.3% hydrogen peroxide solution by mixing 140 ml of 3% hydrogen peroxide solution and 1,260 ml of distilled water.
- **6** Remove the fill-port plug (**c**), fill the reservoir with the hydrogen peroxide solution, and replace the fill-port plug.



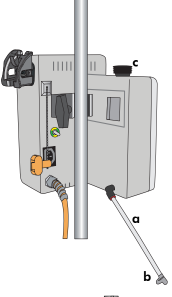
- **8** Turn the HOTLINE[®]3 Warmer ON, and let the recirculating solution circulate for a 30-minute disinfection period.
- **9** Turn the HOTLINE®3 Warmer OFF and unplug the power cord.
- **10** Empty the reservoir.
- **11** Remove the HOTLINE[®]3 Fluid Warming Set and discard according to established hospital procedures.

These suggested instructions are designed to be used in conjunction with established hospital procedures.

Add Recirculating Solution

CAUTIONS

 Do not fill the HOTLINE[®]3 Warmer reservoir with a HOTLINE[®]3
 Fluid Warming Set or a TEMP CHECK in place. Failure to
 remove the HOTLINE[®]3 Fluid Warming Set before the fill
 procedure may result in an air lock in the HOTLINE[®]3 Warmer.





- **1** Prepare the recirculating solution.
- **2** Remove the fill-port plug.
- **3** Fill the reservoir with 1.4 liters of recirculating solution.
- **4** Replace the fill-port plug.

Testing HOTLINE®3 Warmer Operation

Perform all the tests described in the testing section of this manual. See Section 9, *Testing*. The Scheduled Maintenance Checklist below also lists the tests.

Maintenance Log

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

Maintenance Checklist

Task	Every Use	Every 30 Days	Every 12 Months
Clean the Exterior			
General Inspection			
Disinfect the Reservoir and Change Distilled Water or Isopropyl Alcohol solution			
Disinfect the Reservoir and Change the Hydrogen Peroxide Solution			
Alarm Signal Test			
Add Recirculating Solution Test			
Check Disposables Test			
Over Temperature Alarm Test			
Verify Temperature Calibration			
Electrical Safety Tests			

Limited Warranty

Smiths Medical ASD, Inc. (the "Manufacturer") warrants to the Original Purchaser that the HOTLINE[®]3 Blood and Fluid Warmer (the "HOTLINE[®]3 Warmer"), not including accessories, shall be free from defects in materials and workmanship under normal use, if used in accordance with this Operator's Manual, for a period of one year from the actual date of sale to the Original Purchaser. THERE ARE NO OTHER WARRANTIES.

This warranty does not cover normal wear and tear and maintenance items, and excludes any accessory items or equipment used with the HOTLINE[®]3 Warmer.

Subject to the conditions of and upon compliance with this Limited Warranty, the Manufacturer will repair or replace at its option without charge (except for a minimal charge for postage and handling) any HOTLINE[®]3 Warmer (not including accessories) which is defective if a claim is made during such one-year period.

The following conditions, procedures, and limitations apply to the Manufacturer's obligation under this warranty:

- A. Parties Covered by this Warranty: This warranty extends only to the Original Purchaser of the HOTLINE[®] 3 Warmer. This warranty does not extend to subsequent purchasers. The Original Purchaser may be medical personnel, a hospital, or institution which purchases HOTLINE[®] 3 Warmers for treatment of patients. The Original Purchaser should retain the invoice or sales receipt as proof as to the actual date of purchase.
- B. Warranty Performance Procedure: Notice of the claimed defect must be made in writing or by telephone to the Manufacturer as follows: Customer Service Department, Smiths Medical ASD, Inc., 160 Weymouth Street, Rockland, MA 02370, (800) 258-5361. Notice to the Manufacturer must include date of purchase, model and serial number, and a description of the claimed defect in sufficient detail to allow the Manufacturer to determine and facilitate any repairs which may be necessary. AUTHORIZATION MUST BE OBTAINED PRIOR TO RETURNING THE HOTLINE®3 WARMER. If authorized, the HOTLINE®3 Warmer must be properly and

carefully packaged and returned to the Manufacturer, postage prepaid. Any loss or damage during shipment is at the risk of the sender.

- C. Conditions of Warranty: The warranty is void if the HOTLINE[®]3 Warmer has been 1) repaired by someone other than the Manufacturer or its authorized agent; 2) altered so that its stability or reliability is affected; 3) misused; or 4) damaged by negligence or accident. Misuse includes, but is not limited to, use not in compliance with the Operator's Manual or use with non-approved accessories. Removal or damage to the HOTLINE[®]3 Warmer's serial number will invalidate this warranty.
- D. **Limitations and Exclusions:** Repair or replacement of the HOTLINE[®]3 Warmer or any component part thereof is the EXCLUSIVE remedy offered by the Manufacturer. The following exclusions and limitations shall apply:
 - No agent, representative, or employee of the Manufacturer has authority to bind the Manufacturer to any representation or warranty, expressed or implied.
 - 2. THERE IS NO WARRANTY OF MERCHANTABILITY OR FITNESS OR USE OF THE HOTLINE® 3 WARMER FOR ANY PARTICULAR PURPOSE.
 - 3. The HOTLINE[®] 3 Warmer can only be used under the supervision of medical personnel whose skill and judgment determine the suitability of the HOTLINE[®] 3 Warmer for any particular medical treatment.
 - 4. All recommendations, information, and descriptive literature supplied by the Manufacturer or its agents are believed to be accurate and reliable, but do not constitute warranties.

The Manufacturer disclaims responsibility for the suitability of the HOTLINE®3 Warmer for any particular medical treatment or for any medical complications resulting from the use of the HOTLINE®3 Warmer. The Manufacturer shall not be responsible for any incidental damages or consequential damages to property, loss of profits, or loss of use caused by any defect or malfunction of the HOTLINE®3 Warmer.

This warranty gives the Original Purchaser specific legal rights, and the Original Purchaser may have other legal rights which may vary from state to state.

Service

WARNINGS

 No user-serviceable parts. All service must be performed by Smiths Medical or competent personnel. Death or serious injury may occur if this warning is not followed.

All service must be performed by Smiths Medical or competent personnel. Service by any other person or organization voids the warranty and transfers liability for malfunctions of the device to the servicing organization.

Non-Warranty Work

Devices received that are no longer under warranty can be returned for repair at a cost. The device will be promptly inspected and a verbal estimate of the repair cost will be provided. A purchase order will be required from the original purchaser consistent with the verbal estimate. A written estimate will be provided upon request

Before returning the HOTLINE $^{\circledR}$ 3 Warmer for service, contact Smiths Medical for Returned Goods Authorization. Be sure that ALL recirculating solution is drained from the device before packing the HOTLINE $^{\circledR}$ 3 Warmer for shipment.

Note: The HOTLINE® 3 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

Observe national and local codes or requirements for disposal of contaminated materials and for recycling 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 Tel: 1-800-258-5361 (US/CA)

Tel: +1-781-878-8011

www.smiths-medical.com

Specifications and Accessories

System Specifications

Standard Compliance	Guidelines	
Product Safety	IEC 60601-1:2005	5
EMC	EN 60601-1-2, FC Class B	CC 47 CFR Part 15,
Enclosure Protection	IEC 60529 IP Code: IPX1	
Fluid Warmers	ASTM F2172-02	
Physical	Dimensions	
Height, Overall	24.1 cm	(9.5 inches)
Width, Overall	23.1 cm	(9.11 inches)
Depth, Overall	17.8 cm	(7.0 inches)
Weight, Dry	3.5 Kg	(7.6 lbs)
Weight, Wet (with recirculating solution)	5.0 Kg	(11.0 lbs)
Weight, Shipping	3.6 Kg	(7.95 lbs)
Recirculating Solution Capacity	1.4 L	(0.37 gallons)
Maximum Height on I.V. Pole	107 cm	(42 inches)
Environmental	Temperature	Humidity [%]
Operation	10°C to 45°C	10 to 95
Transportation	-18°C to 60°C	5 to 90
Storage	-18°C to 60°C	5 to 90
Thermal	Temperature	
Temperature Set Point	40.0°C ± 0.1°C	
Over Temperature Set Point	41.2°C	
Electrical	Туре	
MAINS Power Input: 115V	115VAC, 50/60 H	z, 3.0 Amps
MAINS Auxiliary Supply Power Output: 115V	115VAC, 50/60 H	z, 1.0 Amps

Electrical	Туре
Protection Against Electrical Shock	Class 1 Equipment, Type BF
Mode of Operation	Continuous
Type of Current	Alternating
Ingress Protection Rating	IPX1
Performance	
Recirculating Solution	Recirculating solution temperature reaches
Temperature	37°C from ambient in about 4 minutes
Normothermic Flow Rates	At gravity flow rates to 5,000 ml per hour

Electromagnetic Compliance

HOTLINE®3 Warmer is certified to be in compliance with the European Communities Council Directive relating to Electromagnetic Compatibility (EMC): (89/336/EEC). Test methods and acceptance criteria as specified in EN 60601-1-2 demonstrate conformance.

Electromagnetic Environmental Recommendations

Recommended separation distances between portable and mobile RF communications equipment and the HOTLINE®3 Warmer

The HOTLINE®3 Warmer is intended for use in an electromagnetic environment in which radiated RF disturbances are controlled. The customer or the user of the HOTLINE®3 Warmer can help prevent electromagnetic interference by maintaining a minimum distance between portable and mobile RF communications equipment (transmitters) and the HOTLINE®3 Warmer as recommended below, according to the maximum output power of the communications equipment.

Rated maximum	Separation distance according to frequency of transmitter m		
output power of transmitter	150 kHz to 80 MHz d=[3.5/V1]√P	80 MHz to 800 MHz d=[3.5/E1]√P	800 MHz to 2.5 GHz d=[7/E1]√P
0.01	0.116	0.116	0.233
0.1	0.368	0.368	0.737
1	1.16	1.16	2.33
10	3.69	3.69	7.38
100	11.66	11.66	23.33

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.

Accessories

REF	Product Description
L-370	HOTLINE®3 Fluid Warming Set
L-10	Gas Vent
PC-8	T-Connector, 20.3 cm (8") Patient Lead with Injection Port
YC-8	Y-Connector, 20.3 cm (8") Patient Lead with Injection Port

Symbols

Symbols	Definitions
0 -	Power switch in the ON position
0	Power switch in the OFF position.
<u></u>	ON/Operation
L GO C	Reservoir Temperature Display
	Add Recirculating Solution
_	Check Disposables
	Over Temperature
	Maximum Reservoir Level
<u></u>	Minimum Reservoir Level
\triangle	Alarm Test
*	Type BF Equipment
	Warning: Do not stick HOTLINE®3 tubing with needles. Patient injury or death could result.

Symbols	Definitions
IPX1	Protected Against Dripping Water
REF	Catalog Number
SN	Serial Number
PN	Part Number
LOT	Batch Code
EC REP	Authorized Representative in the European Community
	Manufacturer
	Date of Manufacture
	Quantity
	Protective Earth [Ground]
\sim	Alternating Current
2	Do Not Reuse
	Consult instructions for use (The symbol appears on the device with a blue background.)
	Consult instructions for use
\triangle	Caution
Â	Electric Shock Hazard
(ATEX)	Not made with natural rubber latex

Symbols	Definitions
LATEX	Caution: This product contains natural rubber latex which may cause allergic reactions.
RX	Caution: Federal (U.S.A.) law restricts this device to sale by or on the order of a physician.
CLASS 1	Device is class type 1 equipment
	Protective earth terminal
	Do not use if package is damaged.
STERILE EO	Sterile fluid path, ethylene oxide gas sterilized
	Temperature Limitation
<u>%</u>	Humidity Limitation
\sum	Use by
*	Keep away from sunlight
*	Keep dry
	Recyclable Product
C SUD US	Device has been tested by TÜV SÜD America, a nationally recognized technical laboratory, to meet all requirements for safety.
AZS	Device has been tested by National Technical Systems, a nationally recognized technical laboratory, to meet U.S. requirements for safety.
	Collect separately for electrical and electronic equipment.
PHT DEHP	Contains or Presence of Phthalate: bis(2-ethylhexyl) phthalate (DEHP)

Numerics D 0.3% Hydrogen peroxide solution Description maintenance schedule 31 components 2 **HOTLINE 3 Fluid Warming Set 4** preparation 11, 31 35% Isopropyl alcohol solution **HOTLINE 3 Warmer 2** maintenance schedule 31 Disinfect the reservoir 10 Display Panel 2, 13 preparation 11, 31 description 13 Disposal information 38 contaminated medical waste 10, 21 Accessories list 41 electrical device 38 Add Recirculating Solution mode 16 Distilled water Add Recirculating Solution test 25 maintenance schedule 31 Add Solution LED 16 Drain tube 3 Additional documentation 37 After Use 20 E Alarm signal test 24 Alarm Signal Test button 14 Electrical safety tests 11, 28 Alcohol (isopropyl) solution 11, 31 Electrical specifications 39 Anesthetic use warning 6 Electromagnetic compliance 40 Assembly instructions 9 Electromagnetic environmental recommendations Auxiliary electrical outlet 3 Environmental specifications 39 C Canada service contact 38 Cautions 7 Fill-port plug 3 L-10 Gas Vent 8 Float switch 3 Change the recirculating solution 31 Check Disposables LED 15 G Check Disposables mode 15 Grounding reliability 7, 17 Check Disposables Test 26 Guidelines for safe use 5 Clamp for I.V. pole 3 Clean exterior surfaces 29 Н Components 2 description 2 **HOTLINE 3 Fluid Warming Set** Connect the intravenous administration set 19 priming volume 4 Contaminated medical waste Hydrogen peroxide solution disposal information 21 preparation 10, 11, 31, 32 Contents list 9 Contraindications 5 Controls 13 I.V. pole mounting height restrictions 9 Conventions used in manual 1 Important safety information 5 Indications for use 1 Infusate delivery temperatures 12 Intravenous administration set connect 19

Isopropyl alcohol solution 11, 31	R
L-10 Gas Vent warnings and cautions 8	Recirculating solution change 31 maintenance schedule 11, 31 preparation 11, 31
LCD 13	protocols 11
LEDs 13	Reflux plug 3 Reservoir 3
Light-emitting diodes (LEDs) 13 Liquid crystal display (LCD) 13	fill with recirculating solution 11
Elquid Crystal display (LCD) 15	Reservoir level display 14
M	reservoir lever display 17
M	S
Maintenance 29	
performed every 12 months 31	Service 37
performed every 30 days 31	Service contacts 38
performed with every use 29	Set Up the HOTLINE 3 Fluid Warming Set 18
Maintenance schedule	Set Up the HOTLINE 3 Warmer 17
recirculating solution 11, 31	Socket 2
Modes of operation 15	Specifications 39 electrical 39
Mounting to I.V. pole 9	environmental 39
N	performance 40
14	physical 39
Non-Warranty work 37	thermal 39
	Standard compliance guidelines 39
0	Storage 21
OFF mode 15	Symbols 42
ON/Operating mode 15	System specifications 39
Operating LED 15	
Operation modes 13, 15	T
Over Temperature Alarm mode 16	Tomporeture
Over Temperature LED 16	Temperature display 13
Over Temperature test 25	Over Temperature Alarm 16
Over Temperature Test button 14	Temperature verification of the recirculating
	solution 26
P	Testing 24
Performance specifications 40	Thermal specifications 39
Physical specifications 39	Troubleshooting 22
Power and Alarm Test Panel 14	Add Recirculating Solution alarm 22
Power cord 3	Check Disposables alarm 22
Power ON/OFF switch 14	electrical interference 23
priming volume	hot cabinet 23
HOTLINE 3 Fluid Warming Set 4	no power 22
Principle of operation 12	Over Temperature alarm 22
	recirculating solution leaks 23

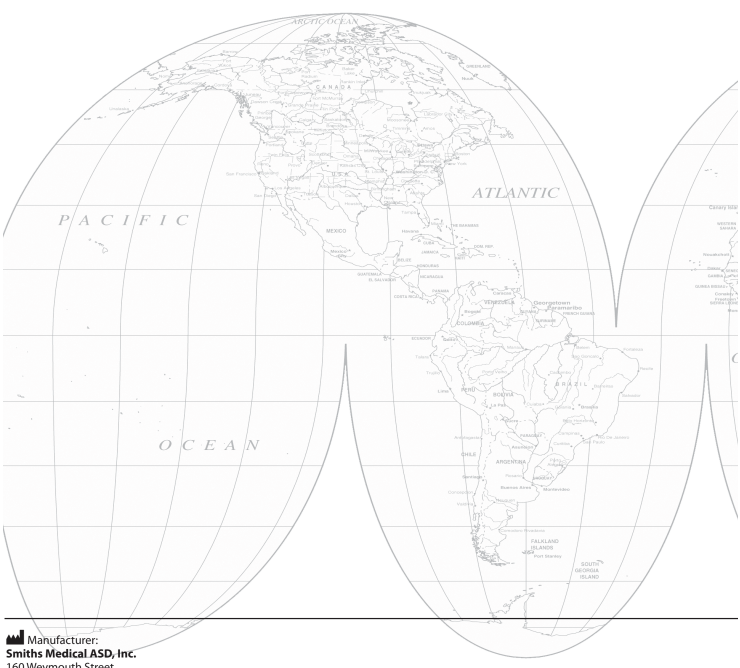
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USA service contact 38 Using the HOTLINE 3 Warmer 20

W

Warnings 5 L-10 Gas Vent 8 Warranty 35

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