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Introduction

The Alaris® GP Volumetric Pump (hereinafter referred to as ‘Pump’) is a small lightweight volumetric infusion pump that provides accurate and reliable infusions over a range of rates.

The Alaris® GP Editor Software is a medical device accessory, which allows the hospital to develop a best-practice data set of IV medication dosing guidelines for patient-specific care areas. Each data set contains a specific library of drugs, as well as a pump configuration appropriate for the care area.

The hospital defined data set is developed and approved through pharmacy and clinical input, and then transferred into the Alaris® GP Volumetric Pump by qualified technical personnel.

**INTENDED USE:**

The pump is designed to meet the infusion requirements within the operating environment specified in this Directions For Use (DFU) including general wards, critical and intensive care, operating rooms and accident and emergency rooms.

This pump is suitable for use by appropriately trained clinicians or nurses. This pump can be used for Intravenous modes, supporting fluid therapy, drug therapy, blood transfusions and parenteral nutrition.

The Asena® brand name has been recently changed to the Alaris® brand name. This change in brand name has no effect on the intended use or functionality of the product. Recommended disposable products for use with this product may refer to either the Asena® brand name or Alaris® brand name and both types are suitable for use with this infusion pump.

About this Manual

The user must be thoroughly familiar with the pump described in this manual prior to use.

All illustrations used in this manual show typical settings and values which may be used in setting up the functions of the pump. These settings and values are for illustrative use only. The complete range of settings and values are detailed in the specifications section.
Creating a Data Set

To create a data set for the Alaris® GP Volumetric Pump, first the hospital will need to develop, review, approve, upload according to the following process. Refer to the Alaris® GP Editor help file for further details and operating precautions.

1. Create Care Area data set (Using Alaris® GP Editor)
   - **Drug List**: Drug names and concentrations for a data set with default value and maximum limits.
   - Up to 100 unique drug names/drug protocol set-ups.
   - **Pump Configuration***: Pump configuration settings and units for dosing only.

2. Review, approve and export data set (Using Alaris® GP Editor)
   - **Review and Approve**: Entire data set report to be printed, reviewed and signed as proof of approval by an authorised person, according to hospital protocol. Signed printout to be kept safe by hospital. Data set status to be set to Approved (Password is required).
   - **Export**: Export data set for use by the Alaris® GP Transfer Tool, or to back up a data set, or to move the data set to another PC.

3. Upload data set to Alaris® GP Volumetric Pump (Using Alaris® GP Transfer Tool)
   - * See note in Pump Configuration section.

![Data set transfers should only be performed by qualified technical personnel.](image)

*Drug parameters have to be in accordance with local regulation and prescribed information.*
Features of the Alaris® GP Volumetric Pump

Alarm indicator
Flow sensor connector
RS232/Nursecall connector (cover removed for clarity)
Release lever for rotating cam
Folded pole clamp
Rotating cam to lock onto horizontal rectangular bars.
Mains fuses cover
Mains inlet
IR communications port
Potential Equalisation (PE) Connector
Alarm indicator
Display
Softkeys
Chevrons
Mute
Pressure
Battery indicator
On/Off
Door Lever
Handle
Flow sensor connector
RS232/Nursecall connector (cover removed for clarity)
Folded pole clamp
Medical device interface (MDI)
## Controls:

<table>
<thead>
<tr>
<th>Symbol</th>
<th>Description</th>
</tr>
</thead>
<tbody>
<tr>
<td><img src="image" alt="ON/OFF button" /></td>
<td><strong>ON/OFF</strong> button - Press once to switch the pump ON. Press and hold down for approximately 3 seconds to switch the pump OFF.</td>
</tr>
<tr>
<td><img src="image" alt="RUN button" /></td>
<td><strong>RUN</strong> button - Press to start the infusion. The green LED will flash during infusion.</td>
</tr>
<tr>
<td><img src="image" alt="HOLD button" /></td>
<td><strong>HOLD</strong> button - Press to put the infusion on hold. The amber LED will be lit while on hold.</td>
</tr>
<tr>
<td><img src="image" alt="MUTE button" /></td>
<td><strong>MUTE</strong> button - Press to silence alarm for (approximately) 2 minutes. The alarm will resound after this time.</td>
</tr>
</tbody>
</table>
| ![BOLUS button](image) | **BOLUS** button - Press to access **BOLUS** softkey. Press and hold down softkey to operate.  
**BOLUS** - fluid or drug delivered at an accelerated rate.  
- Pump is infusing  
- Infusion set is connected to patient.  
- Volume infused (VI) is added to the total volume infused displayed. |
| ![OPTION button](image) | **OPTION** button - Press to access optional features. |
| ![PRESSURE button](image) | **PRESSURE** button - Use this button to display the pumping pressure and adjust the alarm limit. |
| ![CHEVRON keys](image) | **CHEVRON** keys - Double or single for faster / slower increase / decrease of values shown on display. |
| ![BLANK SOFTKEYS](image) | **BLANK SOFTKEYS** - Use in conjunction with the prompts shown on the display. |

## Indicators:

<table>
<thead>
<tr>
<th>Symbol</th>
<th>Description</th>
</tr>
</thead>
<tbody>
<tr>
<td><img src="image" alt="AC POWER indicator" /></td>
<td><strong>AC POWER</strong> indicator - When illuminated the pump is connected to an AC power supply and the battery is being charged.</td>
</tr>
<tr>
<td><img src="image" alt="BATTERY indicator" /></td>
<td><strong>BATTERY</strong> indicator - When illuminated the pump is running on the internal battery. When flashing the battery power is low with less than 30 minutes of use remaining.</td>
</tr>
</tbody>
</table>
### Labelling Symbols:

<table>
<thead>
<tr>
<th>Symbol</th>
<th>Description</th>
</tr>
</thead>
<tbody>
<tr>
<td><img src="image" alt="Attention Symbol" /></td>
<td>Attention (Consult accompanying document)</td>
</tr>
<tr>
<td><img src="image" alt="Potential Equalisation Symbol" /></td>
<td>Potential Equalisation (PE) Connector</td>
</tr>
<tr>
<td><img src="image" alt="RS232/Nursecall Connector" /></td>
<td>RS232/Nursecall Connector.</td>
</tr>
<tr>
<td><img src="image" alt="Defibrillation-proof Symbol" /></td>
<td>Defibrillation-proof type CF applied part. (Degree of protection against electrical shock)</td>
</tr>
<tr>
<td><img src="image" alt="IPX1 Symbol" /></td>
<td>Protected against vertically falling drops of water</td>
</tr>
<tr>
<td><img src="image" alt="Alternating Current Symbol" /></td>
<td>Alternating Current</td>
</tr>
<tr>
<td><img src="image" alt="CE 0086 Symbol" /></td>
<td>Device complies with the requirements of the EC Directive 93/42/EEC. Registered with the CE Mark.</td>
</tr>
<tr>
<td><img src="image" alt="Date of Manufacture Symbol" /></td>
<td>Date of Manufacture</td>
</tr>
<tr>
<td><img src="image" alt="Manufacturer Symbol" /></td>
<td>Manufacturer</td>
</tr>
<tr>
<td><img src="image" alt="Connector for Flow Sensor Symbol" /></td>
<td>Connector for Flow Sensor</td>
</tr>
<tr>
<td><img src="image" alt="Important Information Symbol" /></td>
<td>Important Information</td>
</tr>
<tr>
<td><img src="image" alt="Not for Municipal Waste Symbol" /></td>
<td>Not for Municipal Waste</td>
</tr>
<tr>
<td><img src="image" alt="Fuse rating Symbol" /></td>
<td>Fuse rating</td>
</tr>
</tbody>
</table>
Main Display - If VTBI has not been set (flow sensor must be used):

![Diagram of display with on hold status]

- **Infusion Status/Drug Name/**Primary or Secondary** (Only if secondary is enabled in the data set)
- **Infusion Rate**
- **Volume Infused**
- **Softkeys**

- **ON HOLD**
  - **Rate**
  - **Volume**

Main Display - If VTBI is set:

![Diagram of display with ADRENALINE*]

- **Infusion Status/Drug Name/**Primary or Secondary** (Only if secondary is enabled in the data set)
- **Infusion Rate**
- **Dose Rate**
- **Volume to be Infused**
- **Volume Infused**
- **Time Remaining**
- **Softkeys**

- **ADRENALINE** *
  - **Rate**
  - **Dose Rate**
  - **Volume to be Infused**
  - **Volume Infused**
  - **Time Remaining**

Screen icon:

<table>
<thead>
<tr>
<th>Icon</th>
<th>Description</th>
</tr>
</thead>
<tbody>
<tr>
<td><img src="clock.png" alt="Clock" /></td>
<td>TIME REMAINING DISPLAY icon - Indicates time remaining before VTBI will be completed. If the time is greater than 24 hours then 24+ will be displayed.</td>
</tr>
<tr>
<td><img src="star.png" alt="Star" /></td>
<td>DRUG PROTOCOL symbol - Indicates drug protocol is in use.</td>
</tr>
<tr>
<td><img src="pressure.png" alt="Pressure" /></td>
<td>PRESSURE INFORMATION icon - Shows the pressure from level 0 being the first bar to level 8. Alarm limits: level 2, 5 or 8.</td>
</tr>
</tbody>
</table>
Operating Precautions

Infusion Sets

- To ensure correct and accurate operation, only use Cardinal Health single use infusion sets described in this Directions For Use.
- It is recommended that infusion sets are changed according to the instructions in the 'Changing the Infusion Set' section. Carefully read the Directions For Use supplied with the infusion set prior to use.
- Use of non-specified infusion sets may impair the operation of the pump and the accuracy of the infusion.
- When combining several apparatus and/or instruments with infusion sets and other tubing, for example via a 3-way tap or multiple infusion, the performance of the pump may be affected and should be monitored closely.
- Uncontrolled flow may result if the infusion set is not properly isolated from the patient i.e. closing a tap in the set or activating an in-line clamp / roller clamp.
- The infusion set may be fitted with an in-line clamp, which can be used to occlude tubing in case it is required to stop fluid flow.
- The Alaris® GP Volumetric Pump is a positive pressure pump, which should use infusion sets fitted with luer lock fittings or equivalent locking connectors.
- To infuse from a burette, close the roller clamp above the burette and open the clamp on the vent on top of the burette.
- Discard infusion set if the packaging is not intact or the protector cap is detached. Ensure sets are not kinked as this may occlude the tubing.

Using Collapsible bags, Glass Bottles & Semi Rigid containers

- It is recommended that the air vent be opened on the Alaris® GP Volumetric Pump set if using glass bottles or semi-rigid containers, to reduce the partial vacuum formed as the fluid is infused from the container. This action will ensure the pump can maintain volumetric accuracy whilst the container empties. The action of opening the air vent for semi-rigid containers should take place after the spiking of the container and priming of the drip chamber.

Steps for Collapsible bags

Follow steps 1 to 3 as shown for the semi-rigid containers, however do not open vent as in step 4, but prime the set as per step 5. Ensure the bag outlet is fully pierced before filling the drip chamber.

Steps for Semi-rigid containers

1. Close the roller clamp
2. Spike the container
3. Fill drip chamber to fill line
4. Open the air vent to allow pressure equalisation - ready for infusion
5. Prime the set by opening / closing the roller clamp

Operating Environment

- When using any infusion pump in conjunction with other pumps or devices requiring vascular access, extra care is advised. Adverse delivery of medication or fluids can be caused by the substantial variation in pressures created within the fluid channels of such pumps. Typical examples of those pumps are used during dialysis, bypass or cardiac assist applications.
- The pump is suitable for use in all establishments, including domestic establishments and those directly connected to the public low voltage power supply network that supplies buildings used for domestic purposes.
- This pump is not intended to be used in the presence of a flammable anaesthetic mixture with air or oxygen or nitrous oxide.

Operating Pressure

- The pumping pressure alarm system is not designed to provide protection against, or detection of extravasation or tissue, complications which can occur.
**Alarm Conditions**

- Several alarm conditions detected by this pump will stop the infusion and generate visual and audible alarms. Users must perform regular checks to ensure that the infusion is progressing correctly and no alarms are operating.

**Electromagnetic Compatibility and Interference**

- This pump is protected against the effects of external interference, including high energy radio frequency emissions, magnetic fields and electrostatic discharge (for example, as generated by electrosurgical and cauterising equipment, large motors, portable radios, cellular telephones etc.) and is designed to remain safe when unreasonable levels of interference are encountered.

- In some circumstances the pump may be affected by an electrostatic discharge through air at levels close to or above 15kv; or by radio frequency radiation close to or above 10v/m. If the pump is affected by this external interference the pump will remain in a safe mode; the pump will duly stop the infusion and alert the user by generating a combination of visual and audible alarms. Should any encountered alarm condition persist even after user intervention, it is recommended to replace that particular pump and quarantine the pump for the attention of appropriately trained technical personnel.

- This pump is a CISPR 11 Group 1 Class B device and uses RF energy only for its internal function in the normal product offering. Therefore, its RF emissions are very low and are not likely to cause any interference with the nearby electronic equipment. However, this pump emits a certain level of electromagnetic radiation which is within the levels specified by IEC/EN60601-2-24 and IEC/EN60601-1-2. If the pump interacts with other equipment, measures should be taken to minimise the effects, for instance by repositioning or relocation.

**Earth Conductor**

- The Alaris® GP Volumetric Pump is a Class I device, therefore must be earthed when connected to an AC power supply.
- This pump also has an internal power source.
- When connected to an external power source, a three-wire (Live, Neutral, Earth) supply must be used. If the integrity of the external protective conductor on the AC power cable has been compromised, the pump should be disconnected from the AC power source and operated utilising the internal battery.

**Hazards**

- An explosion hazard exists if the pump is used in the presence of flammable anaesthetics. Exercise care to locate the pump away from any such hazardous sources.

- Dangerous Voltage: An electrical shock hazard exists if the pump's casing is opened or removed. Refer all servicing to qualified service personnel.

- Do not open the RS232/Nurse Call protective covering when not in use. Electrostatic discharge (ESD) precautions are required when connecting RS232/Nurse Call. Touching the pins of the connectors may result in ESD protection failure. It is recommended that all actions must be taken by appropriately trained personnel.

- If this pump is dropped, subjected to excessive moisture, fluid spillage, humidity or high temperature, or otherwise suspected to have been damaged, remove it from service for inspection by a qualified service engineer. When transporting or storing the pump, use original packaging where possible, and adhere to temperature, humidity and pressure ranges stated in the Specifications section and on the outer packaging.

- If this pump behaves abnormally, remove from service and contact a qualified service engineer.

**Latex Content**

- The Alaris® GP Volumetric Pump does not contain any latex.
Getting Started

Before operating the pump read this Directions For Use (DFU) manual carefully.

Initial Set Up

1. Check that the pump is complete, undamaged and that the voltage rating specified on the label is compatible with your AC power supply.

2. Items supplied are:
   - Alaris® GP Volumetric Pump
   - Directions For Use (CD)
   - AC Power Cable (as requested)
   - Protective Packaging
   - Alaris® GP Editor Software (including the Alaris® GP Transfer Tool) - per hospital

3. Connect the pump to the AC power supply for at least 2½ hours to ensure that the internal battery is charged (verify that the \( \bigcirc \) is lit).

4. On initial start-up the pump will display the Select Language screen. Select the required language from the list displayed using the \( \bigtriangleup \bigtriangleup \bigtriangledown \bigtriangledown \) keys.

5. Press the OK softkey to confirm your selection.

The Alaris® GP Editor Software can be used to create an approved data set that can be uploaded into the pump. However, a default data set is already installed in the pump (See details below).

The pump will automatically operate from its internal battery if the pump is switched on without being connected to the power supply.

Should the pump fail to perform correctly, replace in its original protective packaging, where possible and contact a qualified service engineer for investigation.

Factory Default Data Set

The Alaris® GP Volumetric Pump is supplied with the following factory default data set

<table>
<thead>
<tr>
<th>Parameter</th>
<th>Factory Default Setting</th>
<th>Default Units Enabled for Dosing Only:</th>
</tr>
</thead>
<tbody>
<tr>
<td>AC Fail Warning</td>
<td>Enabled</td>
<td>( \mu g/\text{min} )</td>
</tr>
<tr>
<td>Alarm Volume</td>
<td>Medium</td>
<td>( \mu g/\text{h} )</td>
</tr>
<tr>
<td>Pressure Default</td>
<td>L5</td>
<td>( mg/\text{h} )</td>
</tr>
<tr>
<td>Pressure Max</td>
<td>L8</td>
<td>( g/\text{h} )</td>
</tr>
<tr>
<td>Rate Titration</td>
<td>Disabled</td>
<td>( U/\text{h} )</td>
</tr>
<tr>
<td>Infusion Rate Max</td>
<td>1200ml/h</td>
<td>( \text{mmol}/\text{h} )</td>
</tr>
<tr>
<td>Bolus Mode</td>
<td>Enabled</td>
<td>( \text{ng}/\text{kg}/\text{min} )</td>
</tr>
<tr>
<td>Bolus Rate Default</td>
<td>500ml/h</td>
<td>( \mu g/\text{kg}/\text{min} )</td>
</tr>
<tr>
<td>Bolus Rate Max</td>
<td>12000ml/h</td>
<td>( \mu g/\text{kg} )</td>
</tr>
<tr>
<td>Bolus Volume Max</td>
<td>5ml</td>
<td>( mg/\text{kg}/\text{min} )</td>
</tr>
<tr>
<td>Weight Default</td>
<td>1kg</td>
<td>( mg/\text{kg} )</td>
</tr>
<tr>
<td>AIL Limit Max</td>
<td>100( \mu l )</td>
<td>( U/\text{kg} )</td>
</tr>
<tr>
<td>VTBI Max</td>
<td>9999ml</td>
<td>( \text{mmol}/\text{kg}/\text{min} )</td>
</tr>
<tr>
<td>Secondary Infusion</td>
<td>Disabled</td>
<td>( \text{mmol}/\text{kg}/\text{h} )</td>
</tr>
</tbody>
</table>
Pole Clamp Installation

A pole clamp is fitted to the rear of the pump and will provide secure fixing to vertical I.V. poles of a diameter between 15 and 40 mm.

1. Pull the folded pole clamp towards you and unscrew the clamp to leave enough room for the size of the pole.
2. Place pump around pole and tighten screw until the clamp is secured to the pole.

*Alaris® DS Docking Station, Asena® IDS Docking Station & Alaris® Gateway Workstation.
Pump can only be mounted on the horizontal section of the docking stations listed above.

Docking Station/Workstation* or Equipment Rail Installation

The rotating cam can be fitted to the rectangular bar on the Docking Station/Workstation* or equipment rails measuring 10mm by 25mm.

1. Align the rotating cam on the rear of the pump with the rectangular bar on the Docking Station/Workstation* or the equipment rail.
2. Push the pump firmly onto the rectangular bar or equipment rail.
3. To release, push the release lever and pull the pump forwards.

*It is recommended that infusion bags be located on a hanger directly above the pump with which they are being used. This minimises the potential for confusion of infusion sets when multiple volumetric pumps are used.

Never mount the pump such that the infusion stand becomes top heavy or unstable.
Ensure pole clamp is folded away and stored within recessed area at the rear of the pump before connecting to a Docking Station/Workstation* or when not in use.
ALARIS® SAFETY CLAMP**:

Safety Clamp Frame

Safety Clamp Tab

Safety Clamp Slider

SAFETY CLAMP IN NON OCCLUDED POSITION:

When a new infusion set is removed from packaging the Safety Clamp will be in this position*:

Clamp in NON OCCLUDED POSITION

FLOW ENABLED

* This is necessary to avoid tube damage during storage and to ensure correct sterilisation and allows immediate priming.

SAFETY CLAMP IN OCCLUDED POSITION:

After infusion set is loaded into the pump, opening the door activates door hooks which will pull the Safety Clamp slider out, as shown:

Clamp is in OCCLUDED POSITION

NO FLOW

MANUALLY OPERATING THE SAFETY CLAMP

To move the slider into the non occluded position manually, push up Safety Clamp Tab and push Safety Clamp Slider completely into Frame:

1. Push up

2. Push

Pushing on the Safety Clamp Slider enables full set flow to the patient. Therefore it is recommended to always close the roller clamp as well. However, if gravity infusion is required, push up Safety Clamp Tab and push orange Safety Clamp Slider completely into Frame to enable flow. The gravity infusion can be regulated using the roller clamp on the set.

** - Hereinafter referred to as to as 'Safety Clamp'.
Getting Started (Continued) - Loading an Infusion Set

Ensure the appropriate infusion set for the fluid/drug to be infused has been selected.
Follow the instructions supplied with the individual infusion set.
Only use Alaris® GP Volumetric Pump infusion sets. (Refer to ‘Infusion Sets’ section of the DFU)
Position the fluid container to avoid spillage onto the pump.
Ensure that the tubing is inserted completely into the top set retainer through to the tubing guide avoiding any slack.

Loading an Infusion Set: Alaris® Safety Clamp in the NON OCCLUDED position - FLOW ENABLED

1. Remove infusion set from package and close roller clamp.

2. Insert the bag spike into the fluid container and hang appropriately.
   At a minimum height of 300 mm above the pump.

3. Fill the drip chamber to the fill line if shown.
   (Approximately half full) Refer to operating precaution section 'Using Collapsible bags, Glass Bottles & Semi-Rigid containers'.

4. Open roller clamp and prime set slowly (to prevent air bubbles) ensuring all air is removed.

5. Close roller clamp.

6. Switch the pump on. Open door and load infusion set as follows:
   • Fit blue adaptor of infusion set into blue top set retainer.
   • Insert orange safety clamp into orange retainer.
   • Ensure infusion set is fully inserted into tubing guide.

7. Close door and open roller clamp. Ensure no drops are falling in the drip chamber.

8. Ensure all air is removed from the set. Connect the infusion set to the patient access device.

Loading an Infusion Set: Alaris® Safety Clamp in the OCCLUDED position - NO FLOW

1. Follow steps 1 to 4 as above where necessary.

2. Ensure roller clamp is closed.

3. Open door and load infusion set as follows:
   • Fit blue adaptor on infusion set into blue top set retainer.
   • Insert orange safety clamp (leaving slider extended) in the occluded position into orange retainer.

   Pushing on the Safety Clamp Slider may lead to uncontrolled flow to the patient. Therefore, always close the roller clamp before pushing on the safety clamp slider.

4. Ensure infusion set is fully inserted into tubing guide.

5. Close door and open roller clamp. Ensure no drops are falling in the drip chamber.

6. Ensure all air is removed from the set. Connect the infusion set to the patient access device.

Ensure the appropriate infusion set for the fluid/drug to be infused has been selected.
Follow the instructions supplied with the individual infusion set.
Only use Alaris® GP Volumetric Pump infusion sets. (Refer to ‘Infusion Sets’ section of the DFU)
Position the fluid container to avoid spillage onto the pump.
Ensure that the tubing is inserted completely into the top set retainer through to the tubing guide avoiding any slack.
Getting Started (Continued) - Starting the Infusion

**PRIME AND LOAD THE SET (Refer to 'Loading an Infusion Set')**

1. Ensure the pump is connected to an AC power supply (also operates from battery).
2. Connect flow sensor, if required. (See 'Flow Sensor Operation')
3. Press the \( \text{\text{OK}} \) key.

   *The pump will run a short self-test. Check two beeps are activated during this test.*
   *Check the displayed date and time are correct.*
   *Check display shows the data set name and version number.*

   **NOTE:** The pump starts up and displays previous settings.

- **No Drug Name**
- **ml/h**
- **Drug Name***

### CLEAR SETUP?

<table>
<thead>
<tr>
<th>RATE</th>
<th>300 ml/h</th>
</tr>
</thead>
<tbody>
<tr>
<td>VTBI</td>
<td>46.5 ml</td>
</tr>
<tr>
<td>VOLUME</td>
<td>3.5 ml</td>
</tr>
<tr>
<td>CLEAR</td>
<td>KEEP</td>
</tr>
</tbody>
</table>

- **Drug Protocol**
- **Dosing Only**

### CLEAR SETUP?

<table>
<thead>
<tr>
<th>RATE</th>
<th>150 ml/h</th>
</tr>
</thead>
<tbody>
<tr>
<td>CONC</td>
<td>0.01 mg/ml</td>
</tr>
<tr>
<td>VOLUME</td>
<td>1.4 ml</td>
</tr>
<tr>
<td>CLEAR</td>
<td>KEEP</td>
</tr>
</tbody>
</table>

- **Primary/Secondary**
- **Drug Names***

### CLEAR SETUP?

<table>
<thead>
<tr>
<th>RATE</th>
<th>300 ml/h</th>
</tr>
</thead>
<tbody>
<tr>
<td>VTBI</td>
<td>50.0 ml</td>
</tr>
<tr>
<td>VOLUME</td>
<td>0.0 ml</td>
</tr>
<tr>
<td>CLEAR</td>
<td>KEEP</td>
</tr>
</tbody>
</table>

If secondary infusions have been enabled in the data set, then 'PRIMARY' may also alternate.

### CLEAR SETUP?

- Selecting **KEEP** will retain all previous rate and volume settings.
- Selecting **CLEAR** will automatically reset the rate and volume settings to zero and the **INFUSION SETUP - SELECT** screen will be displayed:

5. If **CLEAR** was selected, choose from either **ml/h, DOSING ONLY** or **DRUGS (A-Z)** and press **OK** to confirm. Then follow the prompts as required. (Refer to 'Basic Features -Drugs and Dosing' section)

6. Clear **VOLUME** infused, if required. (Refer to 'Clear Volume Infused' section, this is recommended for a new patient or when a new infusion is set-up.)

7. Enter **VTBI** (if required) by selecting **VTBI** softkey on main display. (Refer to 'Setting a VTBI' section)

   Set **VTBI** by using the **BAGS** option and/or \( \text{\text{OK}} \) keys and press **OK** to confirm.

8. Enter or adjust the **RATE** (if necessary) using the \( \text{\text{OK}} \) keys.

9. Press \( \text{\text{OK}} \) key to start the infusion. **INFUSING** will be displayed.

   **NOTE:** The green run button LED will flash to show that the pump is infusing.

---

* If a drug name is selected, then 'CLEAR SETUP?' will alternate with the drug name. If secondary infusions have been enabled in the data set, then 'PRIMARY' may also alternate.
Basic Features - Drugs and Dosing

The following options enable the pump to be set-up for use with a specific drug name and/or drug protocol. Drugs are pre-configured in the Alaris® GP Editor to enable rapid selection of the drug name, dosing units and default rate. For increased security using a configured drug, maximum and minimum safety limits are programmable for concentration and dose rates. (Using the Alaris® GP Editor software)

When adjusting an infusion using the dose rate, the display may not show any corresponding changes to the infusion rate in ml/h. This does not affect the accuracy of the infusion.

Selecting the INFUSION SETUP

1. Press the \( \uparrow \) button to first access the options menu.
2. Drugs and dosing set-up options are available by selecting INFUSION SETUP from the list using the \( \uparrow \downarrow \) keys.
3. Select from the list of the options (ml/h, DOSING ONLY or DRUGS) as detailed below and press the OK softkey to confirm the selection.

<table>
<thead>
<tr>
<th>ml/h</th>
</tr>
</thead>
<tbody>
<tr>
<td>ml/h</td>
</tr>
<tr>
<td>DOSING ONLY</td>
</tr>
<tr>
<td>DRUGS: A B C D E F</td>
</tr>
<tr>
<td>G H I J K L M</td>
</tr>
<tr>
<td>N O P Q R S</td>
</tr>
<tr>
<td>T U V W X Y Z</td>
</tr>
</tbody>
</table>

SELECT WITH \( \uparrow \downarrow \)

| OK |

1. Select ml/h from the list using the \( \uparrow \downarrow \) keys (if necessary).
2. Press OK to confirm.
3. Enter the ml/h rate as prompted on the display in the next screen.

Dosing Only

1. Select DOSING ONLY from the list using the \( \uparrow \downarrow \) keys.
2. Press OK to confirm.
3. Select the dosing units from the list using the \( \uparrow \downarrow \) keys, press OK to confirm.
4. Enter WEIGHT\(^1\) using the \( \uparrow \downarrow \) keys, press OK to confirm.
5. Use the \( \uparrow \downarrow \) keys to select the TOTAL VOLUME\(^2\), press OK to confirm.
6. Enter DRUG AMOUNT using the \( \uparrow \downarrow \) keys and if units need to be changed, select UNITS which will scroll through the units available. Press OK to confirm selection.
7. A summary of the DOSING ONLY information is displayed, to CONFIRM? all details shown press OK. The BACK softkey may be used at any time to return to the previous screen.

Drugs

1. Select the required DRUGS alphabetical row from the list using the \( \uparrow \downarrow \) keys.
2. Press OK to confirm.
3. Select the drug from the displayed list using the \( \uparrow \downarrow \) keys, press OK to confirm.
4. Enter WEIGHT\(^1\) using the \( \uparrow \downarrow \) keys, press OK to confirm.
5. Use the \( \uparrow \downarrow \) keys to enter the TOTAL VOLUME\(^2\), press OK to confirm.
6. Enter DRUG AMOUNT using the \( \uparrow \downarrow \) keys, press OK to confirm selection.
7. A summary of the DRUG information is displayed, to CONFIRM? all details shown press OK. The BACK softkey may be used at any time to return to the previous screen.

1 - Only displayed if weight based units are used.
2 - Total Volume = Drug Volume + Diluent Volume i.e. Total Volume of fluid in the fluid container after a drug is added.

A - When adjusting an infusion using the dose rate, the display may not show any corresponding changes to the infusion rate in ml/h. This does not affect the accuracy of the infusion.
Basic Features

Clear Volume Infused

This option enables the volume infused to be cleared.
1. Press the VOLUME softkey on main display to show the clear VOLUME INFUSED option.
2. Press the CLEAR softkey to clear the volume infused. Press the QUIT softkey to retain the volume.

NOTE: On completion of VTBI pump will continue to infuse at KVO rate.

Setting a VTBI

A VTBI can only be set if the pump is on hold, then as follows by either:
1. Using the keys:
   a) Press the VTBI softkey on main display to enter the volume to be infused screen.
   b) Enter the volume to be infused using the keys and press OK to confirm.

OR
2. Using the BAGS softkey:
   a) Press the VTBI softkey on main display to enter the volume to be infused screen.
   b) Select the BAGS softkey, select the required bag volume using the keys and press OK to confirm the selection.
   c) Press OK to confirm again, or adjust the VTBI using the keys.

KVO (Keep Vein Open) Rate

At the end of VTBI, the pump will first display VTBI DONE/INFUSING KVO. Press CANCEL to display KVO screen.
The pump continues to infuse at a very low rate (refer to “Specifications” section of this DFU). KVO is used to keep the patients vein open, in order to prevent blood clots and catheter occlusions.
NOTE: If the KVO rate (5ml/h) is greater than the set infusion parameters then the pump will continue to infuse at the set infusion rate. The KVO rate will flash on screen to indicate this is not the usual infusion rate.
The pump will beep every 5 seconds while in KVO mode.

Pressure

To check and adjust the pressure level, press the button. The display will change to show the current pumping pressure level and the pressure alarm limit. The pressure alarm limit can be set via the Alaris® GP Editor.
1. Press the keys to increase or decrease the alarm limit (L2,L5 or L8). The new limit will be indicated on the display.
2. Press OK to exit the screen.

The pressure alarm limit is auto adjusted and is fixed at level 8 (L8) for rates above 200ml/h.
The interpretation of pressure readings and occlusion alarms are the responsibility of the clinician depending on the specific application.
Basic Features

Bolus Infusions

**Bolus** - Administering a controlled volume of fluid or drug at an increased rate for diagnostic or therapeutic purposes. The pump should always be infusing and always attached to the patient. (Drugs given by an IV bolus could achieve immediate and high drug concentration levels.)

The bolus feature can be configured via the Alaris® GP Editor to:

a) BOLUS Disabled
b) BOLUS Enabled

**BOLUS Disabled**

If configured to Disabled, pressing the button will have no effect and the pump will continue to infuse at the set rate.

**BOLUS Enabled**

Press and hold the (flashing) BOLUS softkey to deliver the required bolus. The bolus rate can be adjusted. The bolus volume is limited in the configuration via Alaris® GP Editor.

1. During infusion press the button once to display the bolus screen.
2. Use the keys to adjust the bolus rate if required.
3. To deliver the bolus press and hold the BOLUS softkey. During the bolus, the volume being infused is displayed. When the desired bolus volume has been delivered or the bolus volume limit is reached, release the softkey. The bolus volume is added to the total volume infused displayed.

If the volume to be infused (VTBI) is reached during a bolus, the VTBI complete alarm will sound. Press to silence the alarm or CANCEL to acknowledge the alarm. See VTBI section for more details on VTBI operation. When using infusion set 63280NY the maximum infusion rate is 150ml/h.

**Rate Titration**

If Rate Titration is enabled (via the Alaris® GP Editor) the infusion rate or dose rate (if available) can be adjusted while infusing.

1. Select the new rate using the keys.
   
   The message <TITRATE PRESS TO CONFIRM > will flash on screen and the pump continues to infuse at the original rate.

2. Press the button to confirm the new infusion rate and start infusing at the new rate.

If Rate Titration is disabled the rate can only be adjusted whilst ON HOLD:

1. Press the button to put the pump ON HOLD.
2. Select the new rate using the keys.
3. Press the button to start infusing at the new rate.
To set doserate or flowrate in precise increments it may be necessary to switch between the rate adjust options SET BY DOSERATE and SET BY ml/h. An arrow to the left of the rate display shows the rate changed when the keys are used to increase/decrease the infusion rate.

To set a doserate precisely the arrow must be pointing to the doserate (for example: mg/kg/h); the flowrate will be calculated from the doserate.

To precisely set a flowrate the arrow must be pointing to flowrate (ml/h); the doserate will be calculated from the flowrate.

Selecting the SET BY ml/h Option
1. Press the button to access the options menu.
2. Select the SET BY ml/h option using the keys and press the OK softkey indicated on the screen. This will select the set by flowrate option, the arrow on the display will automatically select the flowrate, the flowrate can be adjusted if necessary.

Selecting the SET BY DOSERATE Option
1. Press the button to access the options menu.
2. Select the SET BY DOSERATE option using the keys and press the OK softkey indicated on the screen. This will select the set by doserate option, the arrow on the display will automatically select the doserate, the doserate can be adjusted if necessary.

Clearing the drug name is only available if drug name only has been selected:
1. Press to put the pump ON HOLD.
2. Press the button to access the options menu.
3. Select Drug Name Only using the keys, press OK to confirm.
4. Select Clear Drug Name (displayed if a name only is selected) using the keys. Press the OK softkey to confirm the selection.

If a secondary infusion has already been setup, then access to the primary infusion setup is as follows:
1. Press to put the pump ON HOLD.
2. Press the button to access the options menu.
3. Select Primary Setup and press the OK softkey to confirm. Make changes to the primary setup as necessary.

To setup a secondary infusion refer to the next page.

To review currently selected dosing information:
1. Press the button to first access the options menu.
2. Select DOSING SUMMARY.
3. Review the information and then press the QUIT softkey.

To review pump information:
1. Press the button to access the options menu.
2. Select PUMP DETAILS.
3. Review the information and then press the QUIT softkey.
Basic Features - Secondary (Piggyback) Infusions

Secondary (or piggyback) Infusion mode is only available if it has been configured. The application of secondary infusions should be limited to the intermittent therapy of medications which are not sensitive to the total time required to complete an infusion.

- Typically antibiotics may be infused using a secondary infusion, where the primary infusion is limited to maintenance fluid. If intending to use the secondary infusion facility, the primary infusion should be a maintenance fluid only and is not indicated for drug therapy.
- The application of secondary infusions for delivery of critical drugs, particularly those with a short half life, is NOT indicated for use. These drugs should be administered through a dedicated pump channel.
- Dependent upon factors such as fluid viscosity, the secondary infusion rate, head height between the secondary and primary fluid containers and the use of clamps, flow may occur from the primary fluid container during a secondary infusion. This could result in drug remaining in the container at the end of the secondary infusion, delaying its delivery for a period of time which is dependent upon the primary infusion rate. For example, a secondary infusion of 250ml at 300ml/h could result in approximately 33ml remaining, requiring up to 25 minutes additional time to complete the delivery, assuming a primary infusion rate of 80ml/h (and the use of a 72213N-0006 secondary infusion set and its supplied extension hook). Therefore it is recommended that flow sensors (if used) are disconnected from the pump during secondary infusions.
- Regular monitoring for unexpected primary flow is recommended. If flow from the primary fluid container is not desired during secondary infusion and/or the patient is sensitive to fluid balance, the clamp on the primary infusion set should be closed. Check that no drops fall in the primary drip chamber.
- On completion of the primary infusion the pump will continue at Keep Vein Open rate (KVO) rate.

Setting up a secondary infusion:
1. Ensure Primary infusion has been setup in ml/h (rate > 0ml/h).
2. Press to put the pump ON HOLD.
3. Press to access the OPTIONS screen.
4. Select SECONDARY SETUP, press OK to confirm.
5. Select either NO DRUG NAME or DRUGS A-Z. Press OK to confirm either selection.
6. Enter the secondary RATE using the keys.
7. Press OK to confirm.
8. Set VTBI using the keys. (Refer to 'Setting a VTBI' section)
9. Press OK to confirm.
11. If correct, press OK to continue, or BACK to adjust VTBI or RATE of the SECONDARY mode.
12. Press to start the infusion in secondary mode.
   An ADVISORY screen will be displayed - ENSURE SECONDARY INFUSION SET OPEN.
13. Press OK to start infusing at the displayed rate.

Setting up a subsequent secondary infusion:
On completion of the secondary VTBI, the pump will automatically transition to the primary infusion. (An audible ‘BEEP’ will be heard)
1. Press to place the primary infusion ON HOLD.
2. Follow instructions 3 to 13 of 'Setting up a secondary infusion'.

Typical Secondary infusions:

IV Pole

Extension Hook (approx. 26cm)

 Normally included with the secondary Infusion set. Primary fluid container must hang lower to allow the secondary infusion to run and primary infusion to restart on completion of the secondary infusion.

Primary Fluid Container

Primary Infusion set
e.g. 63420E with an upper Y-Site (SmartSite® Needle-Free Valve).

In-line Clamp

Check Valve

Prevents secondary infusions from flowing back up the primary infusion set instead of to the patient.

Ensure primary set has a backcheck valve upstream from the Y-site.
Changing the Fluid Container

1. Press 🔄 to put the pump ON HOLD.
2. Remove bag spike on infusion set from empty / used container. Discard empty / used container according to hospital protocol.
3. Insert spike into new container.
4. Squeeze the drip chamber approximately half full or up to fill line (if the drip chamber is marked) with fluid.
5. Restart infusion, see “Getting Started”.

Changing the Infusion Set

1. Press 🔄 to put the pump ON HOLD.
2. Close in-line clamp and ensure the access to the patient is isolated.
3. Disconnect the infusion set from the patient.
4. Open pump door and remove infusion set from the pump and discard the set and fluid container according to hospital protocol.
5. Prepare the new infusion set, load infusion set into pump and close the door, see “Loading the Infusion Set”.
6. Restart infusion, see “Getting Started”.

When changing the infusion set or the fluid container use aseptic technique according to hospital protocol.

It is recommended that infusion sets are changed in accordance with the Directions For Use. Carefully read the Directions For Use supplied with the infusion set prior to use.
The set change interval is 24 hours.

SmartSite® Needle-Free System Instructions

SmartSite® Needle-Free Valve is designed to permit safe gravity flow and automated flow, injection and aspiration of fluids without the use of needles by utilising luer lock and luer slip connectors.

Precautions:

- Discard if packaging is not intact or protector caps are unattached.
- If Needle-Free Valve is accessed by a needle in an emergency the valve will be damaged causing leakage. Replace Needle-Free Valve immediately.
- Needle-Free Valve contraindicated for blunt cannula system.
- DO NOT leave slip luer syringes unattended.

DIRECTIONS - Use Aseptic Technique

1. Prior to every access, swab top of Needle-Free Valve port with 70% Isopropyl alcohol (1-2 seconds) and allow to dry (approximately 30 seconds).
   NOTE: Dry time is dependent on temperature, humidity, ventilation of the area.
2. Prime valve port. If applicable, attach syringe to Needle-Free Valve port and aspirate miniscule air bubbles.
3. Replace every 72 hours for stand alone valves. However, if the valve is part of the set, then the set change interval is as per the complete set or 100 activations which ever occurs first. For infusions of blood, blood products or lipid emulsions replace every 24 hours.
   NOTE: During use of Needle-Free Valve port, fluid may be observed between the housing and blue piston. This fluid does not enter the fluid path and requires no action. For product questions or needle-free valve educational materials contact your Cardinal Health, Alaris® Products representative. The Center for Disease Control, Intravenous Nurses Society (USA) and other organizations publish guidelines useful in developing facility guidelines. Consult facility protocols.
Service Configuration Mode

This section comprises of a list of options which can be configured. Some can be entered via the pump SERVICE CONFIGURATION menu (available in Technician Mode) and others through the Alaris® GP Editor Software.

Enter the access code on Alaris® GP Volumetric Pump for SERVICE mode, then select SERVICE CONFIGURATION, see the Technical Service Manual for details.

Use Alaris® GP Editor to configure the pump configuration, drug list and units enabled for each data set.

Access codes should only be entered by qualified technical personnel.

Date & Time

1. Select DATE & TIME from the SERVICE CONFIGURATION menu using the keys and press the OK softkey.
2. Press the OK softkey to confirm.
3. Use the keys to adjust the date displayed, pressing the NEXT softkey to access the next field.
4. When the correct date and time are displayed press the OK softkey to return to the SERVICE CONFIGURATION menu.
5. Press the QUIT softkey to return to the SERVICE menu and press to exit and power down.

Pump Reference Text

This option is used to add reference text to be shown on the pump start up display.

1. Select PUMP REFERENCE from the SERVICE CONFIGURATION menu using the keys and press the OK softkey.
2. Use the keys to enter the text and NEXT to move to the next character.
3. When the desired text has been selected press OK softkey to return to the SERVICE CONFIGURATION menu.
4. Press QUIT to exit back to the main SERVICE menu and press to exit and power down.

Language

This option is used to set the language of messages shown on the pump display.

1. Select LANGUAGE from the SERVICE CONFIGURATION menu using the keys and press the OK softkey.
2. Use the keys to select the language.
3. When the desired language has been selected press OK softkey to return to the SERVICE CONFIGURATION menu.
4. Press QUIT to exit back to the main SERVICE menu and press to exit and power down.

Backlight & Contrast

This option is used to set the backlight and contrast on the pump display.

1. Select BACKLIGHT & CONTRAST from the SERVICE CONFIGURATION menu using the keys and press the OK softkey.
2. Use the keys to adjust BACKLIGHT, CONTRAST & DIMMING. The contrast of the display will change when scrolling through the numbers. (Use PARAM to scroll between each option)
3. When the desired value has been reached press the OK softkey, then QUIT to get back to the SERVICE menu and press to exit and power down.
Pump Configuration available via the Alaris® GP Editor Software

The following options are only configurable via the Alaris® GP Editor Software (PC based), see Alaris® GP Editor help files for further details.

GENERAL SETTINGS:
- **AC Fail Warning**: Warning to indicate that the AC Power has been disconnected and the pump is operating on battery power.
- **Alarm Volume**: The audio volume of the pump used for alarms and warnings.

PRESSURE SETTINGS:
- **Pressure Default**: The default occlusion alarm limit.
- **Pressure Max**: The maximum occlusion pressure alarm limit.

RATE SETTINGS:
- **Rate Titration**: Allows the adjustment of the infusion rate while the pump is infusing, without putting the pump on hold.
- **Infusion Rate Max**: The maximum permissible infusion rate.

BOLUS SETTINGS:
- **Bolus Mode***: Allows use of the bolus feature.
- **Bolus Rate Max**: The maximum permissible bolus rate.
- **Bolus Rate Default***: The default values for bolus rates.
- **Bolus Volume Max***: The maximum permissible bolus volume in a session.

PATIENT SETTINGS:
- **Weight Default**: The default patient weight.

AIR-IN-LINE SETTINGS:
- **AIL Limit Max**: The single bubble AIL setting.

VTBI SETTINGS:
- **VTBI Max**: The maximum permissible setting for the Volume To Be Infused (VTBI).

SECONDARY INFUSION SETTINGS:
- **Secondary Infusion**: Allows the use of a secondary infusion (Piggyback) in the same pump channel.
- **Sec. VTBI Max**: The maximum permissible setting for the Volume To Be Infused for secondary infusions.
- **Sec. Infusion Rate Max**: The maximum permissible infusion rate for secondary infusions.

* These settings may be overriden by drug list settings.
Drug List available via the Alaris® GP Editor Software

The following drug parameters are only configurable via the Alaris® GP Editor Software (PC based), see Alaris® GP Editor help files for further details.

**CONCENTRATION SETTINGS:**
- **Concentration**
  Specifies the drug concentration.
- **Concentration Min**
  The weakest permissible concentration for this drug (amount of drug per ml).
- **Concentration Max**
  The strongest permissible concentration for this drug (amount of drug per ml).

**DOSE RATE SETTINGS:**
- **Weight Based Units**
  Selects weight based or non-weight based units.
- **Dose Rate Default**
  The default dose rate for infusing this drug.
- **Dose Rate Units**
  The unit for dose rate parameters.
- **Dose Rate Max**
  The maximum permissible dose rate for infusing this drug.

**BOLUS SETTINGS:**
- **Bolus Mode**
  Allows the use of the bolus feature for this drug.
- **Bolus Rate Default**
  The default value for bolus rate for this drug.
- **Bolus Volume Max**
  The maximum permissible bolus volume per bolus session, for this drug.

* These settings override pump configuration settings.
Alarms stop the infusion and are indicated by a combination of an audible sound, flashing red alarm indicator and a message on the display.

1. Check the display for an alarm message and review table below for cause and action. Press \[\text{c}\] to silence the sound for 2 minutes, \[\text{CANCEL}\] to clear the message.

2. When the cause of the alarm has been rectified, press the \[\text{c}\] key to resume the infusion. (Exceptions are DO NOT USE & BATTERY EMPTY)

<table>
<thead>
<tr>
<th>Display</th>
<th>Infusion Status</th>
<th>Cause</th>
<th>Action</th>
</tr>
</thead>
</table>
| AIR IN LINE | Infusion stopped | Single air bubble exceeds alarm limit. Set not fitted correctly into air in line detector. | • Assess the amount of air detected by air in line detector.  
• Opening the door may cause an air bubble to rise in the set. Check set for air.  
• Remove air according to hospital policy.  
• Ensure set is fitted correctly in the air in line detector.  
• Check level of fluid in container.  
• Check enough fluid left in drip chamber.  
• Restart infusion. |
| AIR IN LINE | Infusion stopped | Accumulated air bubbles exceeds alarm limit. (Multiple bubbles smaller than the single bubble alarm limit, which has been detected over a 15 min. window and >1ml.) | • Review infusion set for air bubbles and take appropriate action.  
• Check level of fluid in container.  
• Check enough fluid left in drip chamber.  
• Restart infusion. |
| DOOR OPEN | Infusion stopped | Door was opened during an infusion. | • Close door or clamp infusion set using roller clamp.  
• Restart infusion. |
| DOWNSTREAM OCCLUSION | Infusion stopped | A blockage has occurred downstream. | • Check fluid path between pump and patient for clamps, connectors, kinks or blockages.  
• Examine access site for signs of complications (redness, swelling, pain, heat). |
| UPSTREAM OCCLUSION | Infusion stopped | A blockage has occurred upstream. Possible container empty. | • Check set above the pump.  
• Check all clamps above pump.  
• Check fluid level in container.  
• Ensure drip chamber is half filled.  
• Ensure that the bag spike is inserted correctly.  
• Ensure air vent on drip chamber is open on all glass and semi rigid containers. |
| NO FLOW | Infusion stopped | Flow sensor detects no flow. | • Check flow sensor.  
• Check fluid level in container.  
• Ensure all clamps above pump are open.  
• Ensure drip chamber is half filled.  
• Ensure that the bag spike is inserted correctly.  
• Check flow sensor is clean. |
| FLOW ERROR | Infusion stopped | Gross difference between detected drops and expected amount of drops. | • Clamp infusion set using roller clamp.  
• Check flow sensor.  
• Check fluid level in drip chamber. |
| FLOW ERROR (In secondary infusion mode only) | Infusion stopped | Unexpected drops detected. | • Hang secondary container above primary.  
• Check drops are from secondary container when infusing.  
• Flow sensor disconnection is recommended. |
| FREE FLOW | Infusion stopped | Uncontrolled flow possible. | • Clamp infusion set using roller clamp.  
• Remove pump from use. |
| BATTERY EMPTY | Infusion stopped | The internal battery is exhausted. The pump will automatically switch off in the immediate future. | • Connect to power supply immediately or switch pump off. |
| SAFETY CLAMP | Pump on hold | Safety clamp broken or missing. | • Clamp infusion set using roller clamp.  
• Replace infusion set.  
• Investigate and correct set loading. |
| SET MISLOAD | Pump on hold | Set loaded incorrectly. | • Clamp infusion set using roller clamp.  
• Investigate and correct set loading. |
| FLOW SENSOR DISCONNECT | Infusion stopped | Flow sensor unplugged during infusion. | • Check / replace flow sensor or set VTBI. |
### Warnings

Warnings alert the user but may not stop the infusion and are indicated by an audible sound, a flashing amber warning indicator and a message on the display or both.

1. Check the display for a warning message. Press \( \square \) to silence the sound for 2 minutes, **CANCEL** to clear the message.
2. Rectify the cause of the warning or proceed with caution.

#### Warnings:

<table>
<thead>
<tr>
<th>Display</th>
<th>Infusion Status</th>
<th>Cause</th>
<th>Action</th>
</tr>
</thead>
</table>
| WRONG SET | Pump on hold | Safety clamp not detected. | - Clamp infusion set using roller clamp.  
- Check set and close door.  
- Replace infusion set. (If necessary) |
| DOOR CLOSE INCOMPLETE | Pump on hold | Safety clamp in non-occluded position with door open or obstructed. | - Clamp infusion set using roller clamp.  
- Investigate and correct set loading.  
- Close door. |
| DO NOT USE | Pump on hold / infusion stopped | Internal error has occurred. | - Remove pump from use. |
| LEVER OPEN | Infusion stopped | Door lever is open | - Check door lever.  
- Check lever hooks.  
- Check lever is not obstructed, if so, free obstruction. |

### Alarms (Continued)

<table>
<thead>
<tr>
<th>Display</th>
<th>Infusion Status</th>
<th>Cause</th>
<th>Action</th>
</tr>
</thead>
</table>
| BATTERY LOW | Infusion continues | Less than 30 minutes of battery life remaining. | - Connect to power supply.  
- Check power cable. |
| AC POWER FAIL | Infusion continues* | AC power disconnected or failed. | - Reconnect to power supply. |
| VTBI DONE | Infusing KVO | Intended VTBI completed. | - Set new VTBI or clear VTBI. |
| AIR-IN-LINE | Pump on hold | Air detected in infusion set at the start of infusion.  
Set not fitted correctly into air in line detector. | - Ensure set is fitted correctly in the air in line detector.  
- Assess air in infusion set.  
- Check fluid level in drip chamber.  
- Check level of fluid in container. |
| SET CLOCK | Pump on hold | Date / time not set. | - A qualified service engineer must set date / time.  
- Press cancel softkey to continue. |
| TITRATION | Infusion continues | Rate titration not confirmed. | - Confirm or cancel new rate. |

* If pump was on hold the alarm will still be activated but this message will not be displayed.
Prompts: alert the user but may not stop the infusion and are indicated by an audible sound, a flashing amber warning indicator and a message on the display or both.

1. Check the display for a prompt message. Press 🎎 to silence the sound for 2 minutes, CANCEL to clear the message.
2. Rectify the cause of the prompt or proceed with caution.

**Prompts:**

<table>
<thead>
<tr>
<th>Display</th>
<th>Infusion Status</th>
<th>Cause</th>
<th>Action</th>
</tr>
</thead>
<tbody>
<tr>
<td>ATTENTION</td>
<td>Pump on hold</td>
<td>Pump left on hold for 2 minutes without starting the infusion.</td>
<td>• Review pump setup.</td>
</tr>
<tr>
<td></td>
<td></td>
<td></td>
<td>• Start infusion or turn off pump.</td>
</tr>
<tr>
<td>SET VTBI</td>
<td>Pump on hold</td>
<td>No VTBI / flow sensor.</td>
<td>• Set VTBI or fit flow sensor.</td>
</tr>
<tr>
<td>SET NOT FITTED</td>
<td>Pump on hold</td>
<td>No infusion set fitted.</td>
<td>• Fit infusion set.</td>
</tr>
</tbody>
</table>

**Restarting an Infusion following an Air-in-Line Alarm**

The pump may be restarted by opening the door, assessing and removing any air from the tubing guide area and in the infusion set on the patient side of the pump (if required) according to hospital policy. Close the door and cancel the air-in-line alarm. Restarting the infusion will reactivate the air-in-line system and will alarm if the preset air-in-line limit is exceeded.
Flow Sensor Operation (Optional)

The flow sensor automatically monitors the infusion flow rate through the drip chamber. The flow sensor will cause the pump to alarm if a significant deviation from the infusion rate occurs. The flow sensor will also be able to detect empty containers. For this reason we recommend use of a flow sensor wherever possible excluding secondary infusions.

1. Plug the flow sensor into the flow sensor interface located on the top rear part of the pump.
2. Attach the IVAC® Model 180 Flow Sensor to the drip chamber of the infusion set, by pulling back the handles. Refer to the illustration above.
3. Proceed with load, priming, and set-up instructions as described in section “Getting Started”.

NOTE: Ensure drip chamber is half full and upright.

Always attach the flow sensor before you start an infusion.
Avoid using the flow sensor in direct sunlight.
Always ensure lens is clean.

Always replace the flow sensor interface cover when the flow sensor is disconnected.
Infusion Sets - Standard Sets

The Alaris® GP Volumetric Pump uses standard, single-use, disposable infusion sets. The user is responsible for verifying the suitability of a product used, if it is not recommended by Cardinal Health.

For availability please contact your local Cardinal Health, Alaris® Product representative.

<table>
<thead>
<tr>
<th>Part Number</th>
<th>Description</th>
<th>Details</th>
</tr>
</thead>
<tbody>
<tr>
<td>60073</td>
<td>2 Injection Ports</td>
<td>260cm</td>
</tr>
<tr>
<td>60093E</td>
<td>2 SmartSite® Needle-Free Valve Ports.</td>
<td>15 Micron Filter, 1 Backcheck Valve, 260cm</td>
</tr>
<tr>
<td>60123E</td>
<td>2 SmartSite® Needle-Free Valve Ports.</td>
<td>1.2 &amp; 15 Micron Filter, 265cm</td>
</tr>
<tr>
<td>60293E</td>
<td>2 SmartSite® Needle-Free Valve Ports.</td>
<td>1 Backcheck Valve, No Filter, 260cm</td>
</tr>
<tr>
<td>60693</td>
<td>1 Injection Port</td>
<td>15 Micron Filter, 255cm</td>
</tr>
<tr>
<td>60693E</td>
<td>1 SmartSite® Needle-Free Valve Port.</td>
<td>15 Micron Filter, 255cm</td>
</tr>
<tr>
<td>60793</td>
<td>2 Injection Ports</td>
<td>15 Micron Filter, 255cm</td>
</tr>
<tr>
<td>60793E</td>
<td>2 SmartSite® Needle-Free Valve Ports.</td>
<td>15 Micron Filter, 255cm</td>
</tr>
<tr>
<td>60903</td>
<td>15 Micron Filter</td>
<td>250cm</td>
</tr>
<tr>
<td>60593</td>
<td>15 Micron Filter</td>
<td>260cm</td>
</tr>
<tr>
<td>60173E</td>
<td>1 SmartSite® Needle-Free Valve Port.</td>
<td>No Filter, 260cm</td>
</tr>
<tr>
<td>63120V</td>
<td>1 Split Septum Injection Port</td>
<td>1 Backcheck Valve, No Filter, 305cm</td>
</tr>
</tbody>
</table>

Check infusion set materials and drug compatibility before selecting an infusion set. It is recommended that infusion sets are changed according to the instructions in the 'Changing the Infusion Set' section. Carefully read the Directions For Use supplied with the infusion set prior to use.
Infusion Sets - Standard and Blood Sets

The Alaris® GP Volumetric Pump uses standard, single-use, disposable infusion sets. The user is responsible for verifying the suitability of a product used, if it is not recommended by Cardinal Health.

**For availability please contact your local Cardinal Health, Alaris® Product representative.**

**Alaris® GP standard infusion sets**

<table>
<thead>
<tr>
<th>Code</th>
<th>Description</th>
<th>Filters</th>
<th>Length</th>
</tr>
</thead>
<tbody>
<tr>
<td>63200NY</td>
<td>• No Filter • Length: 260cm</td>
<td></td>
<td></td>
</tr>
<tr>
<td>63110V</td>
<td>• 2 Split Septum Injection Ports • No Filter • Length: 290cm</td>
<td></td>
<td></td>
</tr>
<tr>
<td>63401E</td>
<td>• 1 SmartSite® Needle-Free Valve Port • No Filter • Length: 275cm</td>
<td></td>
<td></td>
</tr>
<tr>
<td>63420E</td>
<td>• 2 SmartSite® Needle-Free Valve Ports • 1 Backcheck Valve • No Filter • Length: 295cm</td>
<td></td>
<td></td>
</tr>
</tbody>
</table>

**Alaris® GP blood infusion sets**

<table>
<thead>
<tr>
<th>Code</th>
<th>Description</th>
<th>Filters</th>
<th>Length</th>
</tr>
</thead>
<tbody>
<tr>
<td>60393</td>
<td>• 2 Injection Ports • 200 Micron Filter. • Length: 270cm</td>
<td></td>
<td></td>
</tr>
<tr>
<td>60393E</td>
<td>• 2 SmartSite® Needle-Free Valve Ports. • 200 Micron Filter. • Length: 270cm</td>
<td></td>
<td></td>
</tr>
<tr>
<td>60893</td>
<td>• 1 Injection Port • 200 Micron Filter. • Length: 255cm</td>
<td></td>
<td></td>
</tr>
<tr>
<td>60894</td>
<td>• 1 Injection Port • 200 Micron Filter. • Length: 255cm</td>
<td></td>
<td></td>
</tr>
<tr>
<td>60980</td>
<td>• Twin Spike • 1 Injection Port • 200 Micron Filter. • Length: 250cm</td>
<td></td>
<td></td>
</tr>
<tr>
<td>63477E</td>
<td>• 2 Non-Vented Spikes. • 180 Micron Filter • Length: 305cm • 1 SmartSite® Needle-Free Valve Port.</td>
<td></td>
<td></td>
</tr>
</tbody>
</table>

---

Check infusion set materials and drug compatibility before selecting an infusion set. It is recommended that infusion sets are changed according to the instructions in the ‘Changing the Infusion Set’ section. Carefully read the Directions For Use supplied with the infusion set prior to use.

Please note these drawings are not to scale.
The Alaris® GP Volumetric Pump uses standard, single-use, disposable infusion sets. The user is responsible for verifying the suitability of a product used, if it is not recommended by Cardinal Health.

For availability please contact your local Cardinal Health, Alaris® Product representative.

### Infusion Sets - Opaque, Low Sorbing, Burette and Secondary Sets

#### Alaris® GP light resistant infusion sets

<table>
<thead>
<tr>
<th>Item Code</th>
<th>Features</th>
</tr>
</thead>
</table>
| 60643     | - 15 Micron Filter.  
            - PVC tubing  
            - Length: 250cm |

#### Alaris® GP burette infusion sets

<table>
<thead>
<tr>
<th>Item Code</th>
<th>Features</th>
</tr>
</thead>
</table>
| 60103E    | - 2 SmartSite® Needle-Free Valve Port.  
            - 1 Burette (150ml).  
            - Length: 270cm |
| 63441E    | - 4 SmartSite® Needle-Free Valve Port.  
            - 1 Burette (150ml).  
            - Length: 330cm |

#### Alaris® GP low sorbing infusion sets

<table>
<thead>
<tr>
<th>Item Code</th>
<th>Features</th>
</tr>
</thead>
<tbody>
<tr>
<td>60953</td>
<td>- Length: 260cm</td>
</tr>
<tr>
<td>63260NY</td>
<td>- Length: 295cm</td>
</tr>
</tbody>
</table>

#### Alaris® GP syringe adapter infusion sets

<table>
<thead>
<tr>
<th>Item Code</th>
<th>Features</th>
</tr>
</thead>
</table>
| 63280NY   | - Length: 270cm  
            Restricted to maximum infusion rate of 150ml/h |

#### Alaris® GP secondary infusion set

<table>
<thead>
<tr>
<th>Item Code</th>
<th>Features</th>
</tr>
</thead>
</table>
| 72213N-0006 | - Male luer and hanger  
            - Length: 79cm |

Check infusion set materials and drug compatibility before selecting an infusion set. It is recommended that infusion sets are changed according to the instructions in the 'Changing the Infusion Set' section. Carefully read the Directions For Use supplied with the infusion set prior to use.

Please note these drawings are not to scale.
Associated Products

- The Alaris® DS Docking Station
- The Asena® IDS Docking Station
- The Alaris® Gateway Workstation
Maintenance

Routine Maintenance Procedures

To ensure that this pump remains in good operating condition, it is important to keep it clean and carry out the routine maintenance procedures described below. All servicing should only be performed by a qualified service engineer with reference to the Technical Service Manual (TSM). Circuit diagrams and components parts lists and all other servicing information which will assist the qualified service engineer in performing repair of the parts designated as repairable are available upon request from Cardinal Health.

If the pump is dropped, damaged, subjected to excessive moisture or high temperature, immediately take it out of service for examination by a qualified service engineer.

All preventative and corrective maintenance and all such activities shall be performed at a compliant work place in accordance with the information supplied. Cardinal Health, Alaris® Products will not be responsible should any of these actions be performed outside the instructions or information supplied by Cardinal Health.

Refer to the Technical Service Manual for the access code for technical service.

<table>
<thead>
<tr>
<th>INTERVAL</th>
<th>ROUTINE MAINTENANCE PROCEDURE</th>
</tr>
</thead>
<tbody>
<tr>
<td>As per Hospital policy.</td>
<td>Thoroughly clean external surfaces of the pump before and after prolonged period of storage.</td>
</tr>
</tbody>
</table>
| At least once per year (Refer to TSM for identification of parts) | 1. Inspect AC power supply plug and cable for damage.  
3. Operate the pump on battery power until the battery low alarm then charge the battery to confirm battery operation and charging. |

Please refer to Technical Service Manual for calibration procedures. The units of measurement used in the calibration procedure are standard SI (The International System of Units) units.

Replacing the Mains Fuses

If the pump continually illuminates the battery symbol and the AC power indicator light does not illuminate when the pump is connected to the AC power supply and switched ON, either the power supply fuse in the AC plug, if fitted, or the internal fuses may have blown.

First check the power supply fuse in the AC mains plug, if fitted. If the AC power indicator light does not illuminate remove the pump from service. It is recommended that the AC power fuses are only replaced by a qualified service engineer.

The fuses in the pump should only be replaced by a qualified service engineer. For further information regarding the replacement of fuses refer to the Technical Service Manual.

If the fuses continue to blow, an electrical fault may have occurred and the pump and power supply should be checked out by a qualified service engineer.

Battery Operation

The internal rechargeable battery allows continued operation when the AC power is unavailable, for example during patient transfer or AC power failure. A fully charged battery will provide over 4 hours of operation at typical infusion rates. When connected to the AC power supply for 4 hours, (whether the pump is in use or not) a new battery pack will be fully charged.

The battery is maintenance free, sealed Nickel Metal Hydride and requires no routine servicing. However, to achieve optimum operation, ensure that the battery is fully recharged after full discharge, before storage, and at regular 3 month intervals during storage. Charge retention will eventually degrade. Where retention is critical the internal battery should be replaced every 3 years.

It is recommended that only a qualified service engineer replaces the battery. For further information regarding the replacement of batteries refer to the Technical Service Manual.

Test Routines

The test routines are designed to allow confirmation of many of the pump functions, defaults and calibrations without requiring internal inspection. They do not represent a full calibration check.

See the Technical Service Manual for a complete list of the test procedures, access codes and calibration procedures.
Cleaning and Storage

Cleaning the pump: -
Before the transfer of the pump to a new patient and periodically during the use, clean the pump by wiping over with a lint-free cloth lightly dampened with warm water and a standard disinfectant / detergent solution.

Cleaning and storing the infusion set: -
The infusion set is a disposable single use item and should be discarded after use according to hospital protocol.

Cleaning the door: -
Refer to the Technical Service Manual for information for removing the door to facilitate cleaning of the fluid path, the use of a screwdriver (torx) is required and should only be carried out by a qualified service engineer.

Cleaning the Flow Sensor: -
Before the transfer of the flow sensor to a new infusion set and periodically during use, clean the flow sensor by wiping over with a lint-free cloth lightly dampened with warm water and a standard disinfectant / detergent solution. Ensure the connector does not get wet. Dry flow sensor before use.

To aid cleaning of flow sensors which have been heavily soiled, contaminated or if the handle operation is not free moving, then the flow sensor may be immersed and soaked in clean soapy water (see A). The inside of the spring mechanism can be cleaned by activating it whilst submerged in the water.

After cleaning, the sensor should be allowed to dry fully prior to use.

The plug of the flow sensor must not be immersed in water as damage will occur.

Disposal

Information on Disposal for Users of Waste Electrical & Electronic Equipment

This symbol on the product and/or accompanying documents means that used electrical and electronic products should not be mixed with household waste.

If you wish to discard electrical and electronic equipment, please contact your Cardinal Health affiliate office or distributor for further information.

Disposing of this product correctly will help to save valuable resources and prevent any potential negative effects on human health and the environment which could otherwise arise from inappropriate waste handling.

Information on Disposal in Countries outside the European Union

This symbol is only valid in the European Union. The product should be disposed of taking environmental factors into consideration. To ensure no risk or hazard, remove the internal rechargeable battery and the Nickel Metal Hydride battery from the control board and dispose of as outlined by the local country regulations. All other components can be safely disposed of as per local regulations.
Specifications

Electrical Protection
Class I, Type CF (Defibrillation-proof)

Electrical/Mechanical Safety

Electro Magnetic Compatibility (EMC)

Electrical Safety
Typical earth leakage current 78μA.
Typical Enclosure Leakage Current (Normal Condition) = 0μA
Typical Protective Earth Resistance = 32mOhms
The above measurements are for guidance only, IEC/EN60601-1
limits are defined below:
Earth Leakage Current (Normal Condition) <= 500μA
Enclosure Leakage Current (Normal Condition) <= 100μA
Protective Earth Resistance <= 200mOhms
Classification - Continuous mode of operation, Portable

AC Power Supply -
100 - 230 VAC, 50 - 60Hz, 60VA (Maximum).

Fuse Type -
2 X T 1.25 A, slow blowing.

Dimensions -
148mm (w) x 225mm (h) x 148mm (d). Weight: approx. 2.5kg
(excluding power cable).

Protection against fluid ingress -
IPX1 - Protected against vertically falling drops of water.

BATTERY SPECIFICATIONS -
Rechargeable NiMH (Nickel Metal Hydride). Automatically charges
when the pump is connected to AC power.

Battery Life - 4 hours @ 125ml/h

Battery Charging - 2.5 hours to 95%.

Alarm Conditions -

<table>
<thead>
<tr>
<th>Warnings</th>
<th>Alarms</th>
</tr>
</thead>
<tbody>
<tr>
<td>AC POWER FAIL</td>
<td>AIR IN LINE (SINGLE BUBBLE)</td>
</tr>
<tr>
<td>VTBI DONE</td>
<td>AIR IN LINE (ACCUMULATED)</td>
</tr>
<tr>
<td>BATTERY LOW</td>
<td>DOOR OPEN</td>
</tr>
<tr>
<td>AIR-IN-LINE</td>
<td>DOWNSTREAM OCCLUSION</td>
</tr>
<tr>
<td>TITRATION</td>
<td>UPSTREAM OCCLUSION</td>
</tr>
<tr>
<td>SET CLOCK</td>
<td>NO FLOW</td>
</tr>
<tr>
<td></td>
<td>FLOW ERROR</td>
</tr>
<tr>
<td></td>
<td>FREE FLOW</td>
</tr>
<tr>
<td></td>
<td>ATTENTION</td>
</tr>
<tr>
<td></td>
<td>BATTERY EMPTY</td>
</tr>
<tr>
<td></td>
<td>SAFETY CLAMP</td>
</tr>
<tr>
<td></td>
<td>SET MISLOAD</td>
</tr>
<tr>
<td></td>
<td>FLOW SENSOR</td>
</tr>
<tr>
<td></td>
<td>DISCONNECTED</td>
</tr>
<tr>
<td></td>
<td>WRONG SET</td>
</tr>
<tr>
<td></td>
<td>DOOR CLOSE INCOMPLETE</td>
</tr>
<tr>
<td></td>
<td>DO NOT USE</td>
</tr>
<tr>
<td></td>
<td>LEVER OPEN</td>
</tr>
</tbody>
</table>

Memory Retention -
The electronic memory of the pump will be retained for more
than 2 years with normal use.

Air Sensor - Integral Ultrasonic Sensor.

Environmental Specifications

<table>
<thead>
<tr>
<th>Condition</th>
<th>Operating</th>
<th>Transport &amp; Storage</th>
</tr>
</thead>
<tbody>
<tr>
<td>Temperature</td>
<td>+5°C - +40°C</td>
<td>-20°C - +50°C</td>
</tr>
<tr>
<td>Humidity</td>
<td>20% - 90%*</td>
<td>15% - 95%*</td>
</tr>
<tr>
<td>Atmospheric Pressure</td>
<td>700hPa - 1060 hPa</td>
<td>500hPa - 1060hPa</td>
</tr>
</tbody>
</table>

*Non condensing.
IrDA, RS232 and Nursecall Feature

The IrDA (or RS232 / Nursecall optional feature) is a feature on Alaris® GP Volumetric Pump that allows the pump to be connected to an external device for the purpose of data communication.

**IrDA / RS232 / Nursecall Feature**

**RS232 / Nursecall Connection Data**

<table>
<thead>
<tr>
<th>Nursecall Specification</th>
</tr>
</thead>
<tbody>
<tr>
<td><strong>Connector</strong></td>
</tr>
<tr>
<td><strong>TXD/RXD</strong></td>
</tr>
<tr>
<td><strong>Baud Rate</strong></td>
</tr>
<tr>
<td><strong>Start Bits</strong></td>
</tr>
<tr>
<td><strong>Data Bits</strong></td>
</tr>
<tr>
<td><strong>Parity</strong></td>
</tr>
<tr>
<td><strong>Stop Bits</strong></td>
</tr>
</tbody>
</table>

**Nurse Call Relay Contacts**

Pins 1, 8 + 9, 30V dc, 1A rating

---

**IrDA**

<table>
<thead>
<tr>
<th>Parameter</th>
<th>Specification</th>
</tr>
</thead>
<tbody>
<tr>
<td><strong>Baud Rate</strong></td>
<td>115k Baud</td>
</tr>
<tr>
<td><strong>Start Bits</strong></td>
<td>1 Start Bit</td>
</tr>
<tr>
<td><strong>Data Bits</strong></td>
<td>8 Data Bits</td>
</tr>
<tr>
<td><strong>Parity</strong></td>
<td>No Parity</td>
</tr>
<tr>
<td><strong>Stop Bits</strong></td>
<td>1 Stop Bit</td>
</tr>
</tbody>
</table>

---

The nursecall interface provides a remote backup to the internal audible alarm. It should not be relied upon to replace monitoring of the internal alarm. Refer to the Technical Service Manual for further information regarding the RS232 interface.

The assessment for the suitability of any software used in the clinical environment to control receive data from the pump lies with the user of the equipment. This software should include detection of the disconnection or other failure of the RS232 cable. Any connected analogue and digital components are required to meet IEC/EN60950 for data processing and IEC/EN60601 for medical devices. Anyone connecting additional devices to the signal input or output is a system configurator and responsible for meeting the requirements of the system standard IEC/EN60601-1-1. To connect to the RS232 port use spare part 1000SP01183 - RS232 cable.
**Infusion Specifications**

**System Accuracy:**
Rate Accuracy is ±5.0% [95% Confidence Interval / 80% Population], achieved under nominal conditions²,⁴

**Occlusion Alarm Limits**
Achieved under nominal conditions²,⁵

<table>
<thead>
<tr>
<th>Level</th>
<th>L2 - Low</th>
<th>L5 - Medium</th>
<th>L8 - High &lt;= 200 ml/h</th>
<th>L8 - High &gt; 200 ml/h</th>
</tr>
</thead>
<tbody>
<tr>
<td>Pressure (mmHg) approx.</td>
<td>230 ± 135</td>
<td>460 ± 185</td>
<td>725 ± 200</td>
<td>950 ± 300</td>
</tr>
</tbody>
</table>

**Maximum Occlusion Alarm Pressure:** 1250 mmHg

**Post Occlusion Bolus:**
Bolus volume generated at 25 ml/h when the minimum occlusion alarm threshold is reached <0.45 ml
Bolus volume generated at 25 ml/h when the maximum occlusion alarm threshold is reached <0.95 ml

**Bolus Volume Accuracy:**
Typical: -4.1%, Max: -3.2%, Min: -5.5% 1ml @ 10ml/h
Typical: -1.3%, Max: -0.9%, Min: -1.6% 100ml @ 1200ml/h

**Administering a Bolus**

<table>
<thead>
<tr>
<th>Parameter</th>
<th>Range</th>
</tr>
</thead>
<tbody>
<tr>
<td>Bolus Rate</td>
<td>10 - 1200ml/h in steps of 10ml/h</td>
</tr>
<tr>
<td>Bolus Volume Displayed</td>
<td>0.0ml - 100.0ml/h in steps of 0.1ml</td>
</tr>
</tbody>
</table>

**Starting the Infusion / Set-up**

<table>
<thead>
<tr>
<th>Infusion Parameter</th>
<th>Range</th>
</tr>
</thead>
<tbody>
<tr>
<td>Infusion Rate</td>
<td>1.0 - 99.9ml/h in steps of 0.1ml/h &amp; 100 - 999ml/h in steps of 1ml/h</td>
</tr>
<tr>
<td>VTBI Primary</td>
<td>(0 - OFF), 1 - 9999ml</td>
</tr>
<tr>
<td>VI (Total)</td>
<td>0.1 - 9999ml</td>
</tr>
</tbody>
</table>

**Maximum time for activation of occlusion alarm:**
At Maximum Pressure, time to alarm at 1.0ml/h is nominally 65 [±4] minutes (Maximum <95 min)
At Minimum Pressure, time to alarm at 1.0ml/h is nominally 16 [±2] minutes (Maximum <28 min)

At Maximum Pressure, time to alarm at 25ml/h is nominally 119 [±7] seconds (Maximum <3 min)
At Minimum Pressure, time to alarm at 25ml/h is nominally 29 [±3] seconds (Maximum <50 sec)

**Air in Line Accuracy:**
Single bubble (configurable)
50 μl Setting Measured nominally 54 [±1] μl
100 μl Setting Measured nominally 108 [±1] μl
250 μl Setting Measured nominally 262 [±3] μl
500 μl Setting Measured nominally 514 [±2] μl
Achieved under nominal conditions²
Bubble accumulation: 1ml over a 15 minute window.

**Critical Volume**
The maximum volume infused following a single fault condition is for rates < 10ml/h: +/- 0.25 ml, rates < 100ml/h: +/- 0.5ml, rates ≥100ml/h: +/- 2 ml

**Set based, pump activated Safety Clamp Device to prevent free flow**

Notes:
1. All accuracy specifications are with a 95% confidence interval / 95% population, unless stated otherwise.
2. Nominal conditions are defined as:
   - Set Rate: 1 to 1200 ml/h;
   - Recommended disposable: 60593;
   - Needle: 18 gauge x 40 mm;
   - Solution Type: De-ionized & Degassed Water;
   - Temperature: 23°C ± 2°C
   - Fluid Head Height: +300 ± 30 mm;
   - Back Pressure: 0 ± 10 mmHg.

3. Tested using Distilled water, 20% lipid, 50% glucose, 0.9% Normal Saline and 5% Alcohol solutions.
4. Solution Type: All fluids
   Duration: Over 24 hours of continuous use
   Back Pressure: -100 to +800 mmHg

4. For all conditions⁶ the system accuracy will change by the following percentages:
   - Temperature: nominally -5.7 (±1.5)% at 5 °C and nominally +0.3 (±1.7)% at 40 °C
   - Fluid Head Height: nominally -3.4 (±1.3)% at -0.5 m and 0.0 (±1.1)% at +0.5 m
   - Duration: nominally -1.1 [±0.2] % over 24 hours of continuous use
   - Back Pressure: nominally +2.0 (±1.3)% at -100 mmHg, -13.4 (±1.8)% at +800 mmHg respectively
   Environmental specifications @ 700hPa (± 5%) @ 25 ml/h

5. For all conditions⁶ the occlusion pressure accuracy will change by the following:
   - Temperature: Low setting nominally 7 (±12) mmHg at 5 °C and nominally -24 (±17) mmHg at 40 °C respectively
   - Normal setting nominally 4 (±16) mmHg at 5 °C and -41 (±18) mmHg at 40 °C respectively
   - High Pressure nominally 4 (±14) mmHg at 5 °C and -38 (±21) mmHg at 40 °C respectively

6. All conditions are defined as:
   - Temperature: 5°C to 40°C;
   - Fluid Head Height: -0.5 to +0.5 m;

The specified accuracy may not be maintained if the above conditions are not met, see notes 1 to 6.
Trumpet & Flow Rate Curves

In this pump, as with all infusion systems, the action of the pumping mechanism and variations cause short-term fluctuations in rate accuracy.

The following curves show typical performance of the system in two ways: 1) the accuracy of fluid delivery over various time periods is measured (trumpet curves), and 2) the delay in onset of fluid flow when infusion commences (start-up curves).

Trumpet curves are named for their characteristic shape. They display discrete data averaged over particular time periods or 'observation windows', not continuous data versus operating time. Over long observation windows, short term fluctuations have little effect on accuracy as represented by the flat part of the curve. As the observation window is reduced, short term fluctuations have greater effects as represented by the "mouth" of the trumpet.

Knowledge of system accuracy over various observation windows may be of interest when certain drugs are being administered. Short term fluctuations in rate accuracy may have clinical impact depending on the half-life of the drug being infused and the degree of intervascular integration, the clinical effect cannot be determined from the trumpet curves alone.

The start-up curves represent continuous flow versus operating time for two hours from the start of the infusion. They exhibit the delay in onset of delivery due to mechanical compliance and provide a visual representation of uniformity. Trumpet curves are derived from the second hour of this data. Tests performed per IEC/EN60601-2-24 standard.

Note: The typical flow rate and trumpet curves are achieved using a recommended infusion set.

[Graphs and data tables showing flow rate variations over time and observation windows]
Alaris® Infusion System
Range of products in the Alaris® Infusion System product family are:

<table>
<thead>
<tr>
<th>Part Number</th>
<th>Description</th>
</tr>
</thead>
<tbody>
<tr>
<td>80013UN01</td>
<td>Alaris® GS Syringe Pump</td>
</tr>
<tr>
<td>80023UN01</td>
<td>Alaris® GH Syringe Pump</td>
</tr>
<tr>
<td>80033UND1</td>
<td>Alaris® CC Syringe Pump</td>
</tr>
<tr>
<td>80043UN01</td>
<td>Alaris® TIVA Syringe Pump</td>
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<tr>
<td>80053UN01</td>
<td>Alaris® PK Syringe Pump</td>
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<tr>
<td>80063UN01</td>
<td>Alaris® GP Volumetric Pump</td>
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<tr>
<td>80033UND1-G</td>
<td>Alaris® CC Syringe Pump with Guardrails® Safety Software</td>
</tr>
<tr>
<td>80023UN01-G</td>
<td>Alaris® GH Syringe Pump with Guardrails® Safety Software</td>
</tr>
<tr>
<td>80083UN00-xx*</td>
<td>Alaris® DS Docking Station</td>
</tr>
<tr>
<td>80093UN0x-xx*</td>
<td>Asena® IDS Docking Station</td>
</tr>
<tr>
<td>80203UNS0x-xx*</td>
<td>Alaris® Gateway Workstation</td>
</tr>
</tbody>
</table>

Note: All Alaris® Syringe Pumps are also available without an RS232 option fitted, contact local customer services representative to obtain part number details.

* For Docking Stations and Workstation contact local customer services representative to obtain configurations availability and part numbers.

Spare Parts
A comprehensive list of spare parts for this pump is included within the Technical Service Manual.
The Technical Service Manual (1000SM00013) will be available in electronic format on the World Wide Web at:
www.cardinalhealth.co.uk/alaris.
A username and password are required to access our manuals. Please contact a local customer services representative to obtain login details.

<table>
<thead>
<tr>
<th>Part Number</th>
<th>Description</th>
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</thead>
<tbody>
<tr>
<td>1000SP00487</td>
<td>Internal Battery Pack</td>
</tr>
<tr>
<td>1000SP01183</td>
<td>RS232 Cable</td>
</tr>
<tr>
<td>1001FAOPT91</td>
<td>AC Power Lead - UK</td>
</tr>
<tr>
<td>1001FAOPT92</td>
<td>AC Power Lead - European</td>
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Alaris® GP Editor Software
The following item may be useful when using the Alaris® GP Volumetric Pump.

<table>
<thead>
<tr>
<th>Part Number</th>
<th>Description</th>
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<tbody>
<tr>
<td>1000SP01310</td>
<td>Alaris® GP Editor PC Software Kit</td>
</tr>
</tbody>
</table>
Service Contacts

For service contact your local Affiliate Office or Distributor:

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Freephone: 0508 422734  
Fax: 09 270 6285

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Sverige.  
Tel: (46) 8 544 43 200  
Fax: (46) 8 544 43 225
**Warranty**

Cardinal Health, Alaris® Products ("Cardinal Health") warrants that:

(A) Each new infusion instrument (pump, controller or peripheral instrument) is free from defects in material and workmanship under normal use and service for a period of two (2) years from the date of delivery by Cardinal Health to the original purchaser.

(B) Each new accessory is free from defects in material and workmanship under normal use and service for a period of ninety (90) days from the date of delivery by Cardinal Health to the original purchaser.

(C) Each Mains Cable, Battery, Flow Sensor (ECD) and non-disposable probe is free from defects in material and workmanship under normal use and service for a period of ninety (90) days from the date of delivery by Cardinal Health to the original purchaser.

(D) Each new Thermometer is free from defects in material and workmanship under normal use and service for a period of one (1) year from the date of delivery by Cardinal Health to the original purchaser.

If any product requires repair during the applicable warranty period, the purchaser should communicate directly with its local Cardinal Health service centre to determine the appropriate service facility. Except as provided otherwise in this warranty, repair or replacement will be carried out at Cardinal Health's expense. The product requiring service should be returned promptly, properly packaged, and postage prepaid by purchaser. Loss or damage in return shipment to Cardinal Health shall be at purchaser's risk.

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(B) altered in any way so as to affect, in Cardinal Health's judgement the stability or reliability of the product or has had the product's serial or lot number altered, effaced or removed;

(C) subjected to misuse or negligence or accident; or

(D) improperly maintained or used in any manner other than in accordance with the written instructions furnished by Cardinal Health.

This warranty is in lieu of all other warranties, express or implied, and of all other obligations or liabilities of Cardinal Health, and Cardinal Health neither assumes nor authorises any representative or other person to assume for it any other liability in connection with the sale of Cardinal Health products.

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**Document History**

<table>
<thead>
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<th>CO Number</th>
<th>Date</th>
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<td>Dec 2006</td>
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