Perfusor® Space
and Accessories

Instructions for Use

It is recommended that all pumps at your institution are equipped with the same software version.
CONTENTS

Perfusor® Space Overview.................................................................................3
Symbols on Product ...........................................................................................5
Patient Safety .....................................................................................................6
Menu Structure / Overview .............................................................................13
Menu Structure / Navigation ..........................................................................14
Chapter 1 Operation.......................................................................................16
  1.1 Infusion Start ............................................................................................16
  1.2 Entry with different combinations of Rate, VTBI, and Time ..............17
  1.3 Bolus Application ....................................................................................18
  1.4 Syringe Change and New Therapy Start ..............................................18
  1.5 End of Infusion .......................................................................................19
  1.6 Standby Mode .........................................................................................20

Chapter 2 Advanced Operations ...................................................................21
  2.1 Status Request of Pump when Infusion is Running.........................21
  2.2 Rate, VTBI and Time Change Without Infusion Interruption and Reset of Status Menu Data ...............................................................21

Chapter 3 Special Functions* ........................................................................22
  3.1 Dose Rate Calculation (Overview) .......................................................22
  3.2 Dose Rate Calculation (Operation) .......................................................22
  3.3 Drug Library ...........................................................................................23
  3.4 Instructions for Use of Bar-coding using DoseScan™ and the SpaceStation with SpaceCom .................................................................24

Chapter 4 Options...........................................................................................29
  4.1 Occlusion Pressure ................................................................................29
  4.2 Data lock .................................................................................................29
  4.3 Bolus Rate ..............................................................................................30
  4.4 KVO Mode ..............................................................................................30
  4.5 Contrast / Display Illumination / Keypad Illumination .....................31
  4.6 Alarm Volume .........................................................................................31
  4.7 Date / Time .............................................................................................31
  4.8 Macro Mode ...........................................................................................28

Chapter 5 Alarms............................................................................................32
  5.1 Device Alarm ..........................................................................................32
  5.2 Pre-Alarms and Operating Alarms .......................................................32
  5.3 Reminder Alarms ...................................................................................35
  5.4 Alarm Instruction ...................................................................................35

Chapter 6 Battery Operation and Maintenance ............................................36

Chapter 7 Compatible Syringes ....................................................................38

Chapter 8 Start Up Graphs and Trumpet Curves .........................................40

Chapter 9 Technical Data ..............................................................................41

Chapter 10 TSC** / Service / Training / Disinfecting / Disposal .................44

Chapter 11 Instructions for Use Accessory ..................................................47

Ordering ...........................................................................................................50

Technical Support ..........................................................................................51

* The availability of individual performance features depends on the pump configuration.
** Technical Safety check.
**PERFUSOR® SPACE OVERVIEW**

**Up and down arrow buttons**
Used for scrolling menus, changing settings from 0-9, answering yes/no questions.

**Right and left arrow buttons**
Used for selecting scale values and switching between fields when entering numbers. Press the left arrow button to start a function while the infusion is in operation or to restart when it is interrupted.

**Yellow LED:** Pre-alarm, reminder alarm
**Green LED:** Infusing
**Red LED:** Operating or device alarm
**Blue LED:** Flashes during barcoding function

**Press to initiate bolus.**

**Press to turn pump ON/OFF.**

**Press to Start/Stop the infusion.**

**Drive head with claw mechanism to hold the syringe plunger plate.**

**Protective Cap Note:** Emergency release button on outside of drive head is located below protective cap. Replace Protective cap after pressing.

**The syringe holder locks the syringe in position. To remove the syringe, pull and turn to the right. The drivehead automatically retracts.**

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**Cover of Battery Compartment**
Before changing the battery, always disconnect the pump from the patient and switch the device off.
To remove the battery cover push the button below the battery compartment with the point of a pen and pull the cover away from the device. Slide the green locking mechanism on the back of the battery up and take out the battery pack for replacement.

**P3 port for future options**

**P2 port for AC adapter, SpaceStation, SP connection cable (12V), combination cable and additional accessory cables (staff call, service)**
**Syringe setting**
Pull syringe holder and turn to the right to open the green syringe guide (see red arrow). Syringe must be mounted with flange upright into the slot next to the green syringe guide before the syringe holder is closed. Make sure that the syringe is properly held in place.

**Caution:** Do not touch the parking brake when removing from the syringe holder.

**Attaching the pole clamp (universal clamp)**
Align pole clamp guide slots on the pump with the arms of the pump slots and pole clamp and slide forward until the locking mechanism snaps in place.

To remove, press the release button on the frame, press the handle down and remove the pole clamp.

**Transport**
A maximum of three pumps may be locked together on one pole clamp.

**Lock devices to one another**
Slide the rails of top pump onto slots of bottom pump until they audibly snap in and the green buttons are aligned on top of one another. To separate the units, press the green locking button on the top pump and slide bottom pump forward.

**Caution:** A maximum of three B. Braun Space pumps can be stacked together when used with the PoleClamp SP.

**Caution:** Do not use excessive force when locking pumps together to avoid the possibility of damaging the devices.

**Attachment to IV pole**
Place the pole clamp opening around the IV pole and tighten the screw. Loosen the screw to remove it. To rotate the pump and pole clamp, press the lever (see red arrow) and turn in one of the two directions until the lever clicks into the notch. Press lever to rotate.

**Caution:** Do not lean against the pump as a support while it is attached to the stand! Do not position the pump unit over the patient.
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**List of abbreviations**

- **KVO** = Keep Vein Open
- **LED** = Light-Emitting Diode
  - (indicator lamps)
- **VTBI** = Volume To Be Infused
PATIENT SAFETY

Intended Use
The Perfusor® Space Infusion Syringe Pump System is an electrical, external, syringe infusion pump system indicated for use with adults, pediatrics, and neonates and is intended to provide infusions of parenteral fluids/medications, blood and blood products indicated for infusions through FDA approved routes of administration.

Operation
- Introductory training on the use of the Perfusor Space must be conducted by B. Braun sales representatives or other authorized persons. After each software update, users need to obtain instruction or refer to the user manual on the updates to the device and accessories.
- Prior to start up, check the device and especially the green syringe guide for possible damage. Make note of audio and visual alarms during self-tests.

Warnings

Syringe Size and Selection
- Ensure syringe sizes and models are compatible with the syringe pump (refer to Chapter 7 for more information). Use of incompatible syringes can cause improper pump operation resulting in inaccurate fluid delivery, insufficient occlusion (blockage) sensing, and other potential problems.
- Use the smallest compatible syringe size necessary to deliver the fluid or medication; this is especially important when infusing high risk or life-sustaining medications at low infusion rates (e.g., less than 5 ml per hour, and especially flow rates less than 0.5 ml per hour). Using a larger syringe when infusing at low rates can lead to inadequate syringe pump performance including delivery inaccuracies, delay of therapy and delayed generation of occlusion alarms. This is due to the increased friction and compliance of the syringe plunger tip with larger syringes.
- Use only appropriate syringes/catheters for intended medical use.
- Use only pressure resistant syringes and/or extension sets (min. 2 bar/1500mmHg).

Starting an Infusion or Changing a Syringe
- Electronically prime the syringe pump system before starting an infusion or after replacing a near-empty syringe with a replacement syringe.
  - Verify the fluid flow to the patient is OFF, and if available, use the prime function on the syringe pump to remove any mechanical slack in the system.
• Using the syringe pump's prime feature engages the mechanical components of the pump and decreases the syringe's friction and compliance (i.e., stiffness) to minimize startup delays and delivery inaccuracies, especially at low infusion rates.

• Failure to use the prime feature on the syringe pump after every syringe change and/or tubing change can significantly delay the infusion delivery startup time and lead to delivery inaccuracies.

 ■ During programming and prior to starting an infusion, verify that the syringe size and model on the syringe pump's display screen matches the syringe size and model loaded onto the syringe pump.

 ■ Do not connect to the patient until the syringe has been properly inserted and the syringe plunger plate is properly gripped by the claw on the drive head. Disconnect from the patient when changing the syringe in order to avoid unwanted drug administration.

 ■ Compare the displayed value with the entered value prior to starting infusion.

 ■ Compare physician order to programming parameters including drug name and dosing prior to starting infusion.

Height and Location of Syringe Pump System

■ Ideally, the syringe pump should be level with distal tip of the catheter (e.g., the site of fluid delivery; if accessing a central line the syringe pump should be at the level of the patient’s heart). If the pump height is raised relative to the distal tip of the catheter (e.g., during patient transport), the increase in height of the syringe pump can result in a temporary increase in fluid delivery or bolus until the flow rate stabilizes. Alternatively, if the pump is lowered relative to the distal tip of the catheter, the decrease in height of the syringe pump may result in a decrease in delivery or under-infusion until the flow rate stabilizes.

■ Not to be used adjacent to and/or stacked with other equipment except B. Braun Space devices.

Occlusion Considerations

■ To minimize the amount of time it takes the pump to recognize an occlusion (blockage) and generate an alarm while infusing at low rates (e.g., less than 5 ml per hour, and especially flow rates less than 0.5 ml per hour).

■ Consider the plunger force or occlusion pressure threshold setting and adjust it, as necessary. The lower the plunger force setting or occlusion pressure threshold setting, the shorter the occlusion detection time. However, when infusing viscous or thick fluids (e.g., lipids) the plunger force or occlusion pressure threshold setting may need to be adjusted to reduce false alarms. Please refer to Chapter 4.1 for further detail.
• Use the smallest compatible syringe size necessary to deliver the fluid or medication. This minimizes the amount of friction and compliance (i.e., stiffness) of the syringe plunger tip. Because syringe pumps infuse fluids by precisely controlling the plunger, smaller syringes provide more precise fluid delivery and faster occlusion detection than larger syringes.

• Use the prime feature on the pump when inserting a new syringe and changing a syringe and/or tubing.

• Use accessory devices, which have the smallest internal volume or deadspace (e.g., use microbore tubing when infusing at low rates, shorter length of tubing, etc.). See Chapter 11 for further detail.

- When addressing or clearing an occlusion:

  • Ensure the fluid flow to the patient is OFF to prevent administering an unintended bolus. An occlusion may pressurize the infusion tubing and syringe, which can result in an unintended bolus of drug when the occlusion is cleared. In order to prevent this additional bolus, disconnect the tubing, or relieve the excess pressure through a stopcock, if present. The healthcare professional should weigh the relative risks of disconnection with the risks of unintended bolus of drug.

  • Be aware that using larger size syringes on a high plunger force setting may produce a larger post occlusion bolus due to excessive syringe plunger tip compliance.

Infusion Pump Use

- The user must always review the data displayed before making further medical decisions.

- Consider additional monitoring which may be required when infusing high risk medications.

- When administering critical drugs, a second pump for the drug should be kept ready.

- Regardless of the soft limits, the values entered for the patient must be medically appropriate.

- The default values and limits of the Drug Library provide a safety net and are not intended to be used to define treatment.

- Check the devices and drugs for possible incompatibilities in the respective manufacturer’s literature.

- In the event that values in the dose rate calculator are changed, the flow rate is automatically updated and the dose rate is fixed.

- To prevent the alarm from sounding in error, do not apply force on the drive head when drugs are being administered.
- Make sure the device is securely mounted and positioned. Changes in location and rough jolts can result in minor changes in the delivery characteristics.

- Where several infusion lines are connected on one single vascular access, the possibility of the lines exerting a mutual influence over each other cannot be excluded.

- The use of incompatible disposables may influence the technical specifications of the device.

- Do not hold the pump by the drive head when transporting.

- If the pump is dropped or is exposed to force, it must be checked by the biomedical engineering department.

- Protect the pump and the AC adapter from moisture.

- Do not allow liquids to enter into or come into contact with any openings or electrical connections on the pump or power supply. Fluid exposure in these areas may result in the risk of short circuit, corrosion or breakdown of sensitive electrical components, and/or electrical shock. If fluid exposure occurs, the device should be swapped out with another device in a manner that presents minimal interruption to patient care. The device should remain unplugged until it can be inspected by a trained technician for any evidence of damage and/or residual moisture which may impair the function of the device.

- Do not operate in the presence of flammable anesthetics or in a hyperbaric oxygen chamber.

- Connected electrical components must satisfy IEC/EN specifications (e.g., IEC/EN 60950 for data processing devices). The system administrator who connects additional devices to the system must adhere to the IEC/EN 60601-1-1 standard.

⚠️ Cautions

Use of Accessory Devices

- Use compatible components which have the smallest internal volume or “deadspace” to minimize residual volumes between the syringe and the patient when administering medications or fluids at low infusion rates (e.g., less than 5 ml per hour, and especially flow rates less than 0.5 ml per hour). This reduces the amount of time it takes for fluid to reach the patient, maintains delivery accuracy, and reduces occlusion detection times. For example:
  - Tubing internal diameter: Small bore or microbore tubing is recommended when infusing at low rates
• Tubing length: Tubing length should be minimized, when possible
• Filters: Internal volume (dosepac) of in-line filters should be minimized
• Connection sites: The number of connection sites such as stopcocks and Y-sites should be limited, and high risk or life-sustaining solutions should be connected as close to the intravenous access site as possible.

■ Avoid use of manifolds with ports containing high pressure valves. High pressure valves require additional pressure (e.g., 50–200 mmHg) to open and allow fluid flow. These high pressure valves may cause a significant delay in therapy followed by a sudden bolus once the valve is opened, particularly at low infusion rates (e.g., less than 5 ml per hour, and especially flow rates less than 0.5 ml per hour)

■ Replace disposables per disposable manufacturer’s labeling.

■ Only use luer lock fittings and syringes as well as compatible combinations of devices, accessories, replacement parts and disposables.

■ If multiple devices/infusion lines are connected, a risk of adverse interactions should be considered.

Starting an Infusion or Changing a Syringe

■ Manually prime the syringe and tubing to remove all air, before connecting to the pump unless using the priming feature on the pump to remove air from the tubing.

■ Position the infusion line so that it is free of kinks.

■ Ensure protective cap is present on the end of the drive head prior to operation of the device. Operation without cap present may result in a device alarm which may cause or contribute to a delay in therapy administration when operating in direct sunlight.

■ If staff call is used, check the equipment once after connecting the pump to ensure staff call is working.

Height and Location of Syringe Pump System

■ If using multiple syringe pumps and it is not clinically feasible to have all pumps level with the distal tip of the catheter (or the site of fluid delivery), place the high risk or life-sustaining medications as close to level with the distal tip of the catheter as possible. When infusing multiple high risk or life sustaining medications, consider placing the ones infusing at the lowest rates as close to the level with the distal tip of the catheter as possible.

■ Minimize the height difference between the pump and the patient and avoid changes in the height of the pump (e.g., during transport of critically ill patients) to prevent unintended fluctuations in the flow rate.
- Make sure that the device is properly positioned and secured. Do not position pump unit above patient or in a position where a patient could come to harm, should the pump fall.

Safety Standards


- The EMC limit values (electro-magnetic compatibility) are in compliance with IEC/EN 60601-1-2 and IEC/EN 60601-2-24. When operating in the vicinity of devices that may cause higher interference emissions (e.g. HF surgical devices, cell phones, etc.), the safety distances recommended for these devices must be observed.
Safety Instructions for using Pole Clamp

1. Line pump up with the Pole Clamp guide rails.
2. Slide pump fully into place onto the guide rails.
3. An audible “Click” should be heard.
4. Test the pump is secure.

⚠️ The pump is now securely attached to Pole Clamp.
- Do not lean on the pump when attached to the Pole Clamp.
- Do not position the pump unit above the patient.

⚠️ DO NOT use any Pole Clamp that shows signs of damage.
- DO NOT use Pole Clamp with missing clamp grids.
The pump can be customized to the user’s needs by deactivating the functions of the start-up and option menus as well as the bolus function via the service program.
Display

Explanations

The current therapy is displayed on the top display line. Yes/no question can be answered by pressing (▲) for yes or (▼) for no.

Parameters that can be changed (e.g., rate in mL/h), are opened with (◄) or (►). Edit parameters, change location/levels with (◄) (►). White background displays current location/level. Use (▲) and (▼) to change current setting. Help text on the top/bottom display line shows options (e.g. confirm rate with (OK), start infusion with (OK), or delete rate with (◄)).

Typical display during operation:

All status information is available in the bottom display line. The desired information can be selected by using (▼) or (▲) and afterward is displayed continuously (e.g. complete drug name, time until syringe is empty, etc.).

was pressed during drug administration. Manual bolus can start with 1200 mL/h by pressing the (OK) (see top display line) or continue by entering a bolus limit with (◄) (see bottom display line).
Display

This message is displayed if an attempt is made to change a value with 4 without authorization.

Set pressure

Select pressure level with 4 or 6 and confirm with OK. Cancel entry with 5.

Alarm

Pre-alarm is displayed by a message on the display (e.g. “syringe almost empty”), a signal tone and a blinking yellow LED. Confirm the pre-alarm with OK.

Pump turns off in 2 sec

By pressing and holding 9, a white bar expands from left to right and counts down 3 seconds. Then the pump turns off. As long as a syringe is inserted, the pump does not turn off; instead it goes to “standby mode.” A standby period of up to 24 hours can be entered.
OPERATION

1.1 Infusion start

- Make sure the pump is properly installed. If the device is connected to the power supply, the display shows the battery status, the power supply icon and the most recent therapy.

- Press \( \) to turn the device on. Note the automated self-test – "Self-test active" – and the software version are displayed; two signal tones sound and all three LEDs (yellow, green/red and blue) light up once. Information concerning power supply (A/C or battery operation), the adjusted pressure level and the syringe type (if the syringe is already inserted) are displayed. Afterward, the drive head retracts.

- Press \( \) to begin entering the therapy parameters or to open the pump cover and syringe holder to insert the syringe.

- Insert the syringe with the syringe flange in a vertical position in the slot to the right of the housing. Then close syringe holder and pump cover. Piston Brake moves forward.

**Caution:** Never leave the pump unattended when inserting the syringe.

- Confirm syringe type with \( \). Syringe type displayed must match the syringe inserted.

- Drive head moves forward and grabs the plunger plate of the syringe.

**Caution:** Keep hands away from the moving drive head.

**Note:** Make sure that the piston brake returns to the syringe holder.

- If the prime function is activated, press \( \) in order to prime the infusion line at the rate shown in the display. Cancel prime function by pressing \( \). Repeat process until the line is completely bled. Then press \( \) to continue.

- Connect to patient.

- If necessary, answer questions in the start-up menu with \( \) and \( \) until the rate in the main menu appears.

**Adjust delivery rate:**

- Press \( \) and enter rate using \( \).

- Press \( \) to start infusion. Continuous arrows on the display and green LED show drug being administered.

**Note:** Continuous infusion can be canceled at any time by pressing \( \). The pump can be turned off at any time by pressing \( \) for 3 seconds (exceptions: data lock levels 2 and 3 or if a disposable is loaded).
1.2 Entries with different rate, volume to be infused and time combinations

The Perfusor® Space allows a volume and time limit to be entered in addition to the delivery rate. If two of these parameters have been entered, then the pump calculates the third parameter. If a volume and/or a time is pre-selected, an arrow symbol is placed in front of one of these parameters in the main menu. This is called the “target.” While the pump delivers the drug, this target symbol appears next to the continuous display arrows on the display. From it, you can see that the pump has been programmed either with a volume or a time limit. The target symbol visible in the main menu shows the parameters (volume or time) for that therapy. In case of changes to the delivery rate, the target parameter is generally not adapted to the new flow rate; rather the parameter is recalculated in front of which no target symbol appears. After the infusion starts, the remaining volume and time values are displayed (values count backwards) in the main menu as well as the delivery display.

1.) Enter volume to be infused and time. Delivery rate is calculated and displayed on the bottom left of the display.
   Target: Volumes
   - Select Volume with and open with .
   - Enter Volume with and confirm with .
   - Select time with and open with .
   - Enter time with and confirm with .

Continue in the same way to calculate 2.) and 3.).

2.) Infusion with volume limit
   Enter rate and volume to be infused: The infusion time is calculated and displayed at the bottom left of the display.
   Target: Volume

3.) Infusion with time limit
   Enter rate and time: The volume is calculated and displayed at the bottom left in the display.
   Target: Time

Changing of volume to be infused and time already entered (rate, volume and time were already entered at the time of the change):

a) Target symbol located in front of volume to be infused:
   - Change in volume to be infused => Change the time. Old and new target: Volume to be infused.
   - Change in time => Change the rate. Old and new target: Volume to be infused.

b) Target symbol is located in front of time:
   - Change in time => Adapting the volume to be infused. Old and new target: Time.
   - Change in volume => Adapting the time. New target: Volume to be infused.

Note: Change in volume and time only possible if the pump has been stopped.
1.3 Bolus Application

There are three different options to administer a bolus:

1.) Press manual bolus: \[\text{Manual bolus} \] . Then press \[\text{OK} \] and hold button down. Bolus is delivered as long as the button is pressed down. Bolus volume delivered is displayed.

The maximum bolus volume is 10% of the syringe size or 10 seconds. If this limit is reached, an audible signal follows.

2.) Bolus with volume pre-selection: Press \[\text{Bolus} \] , press \[\text{Bolus} \] and select bolus limit \[\text{Bolus Limit} \] . Press \[\text{OK} \] to confirm and start the bolus. Regardless of the service program settings, an audible signal sounds after at completion of the bolus.

3.) Bolus with rate calculation: Press \[\text{Bolus} \] , press \[\text{Bolus} \] and select bolus dose by pressing \[\text{Bolus Dose} \]. Press \[\text{OK} \] to confirm the bolus dose. Time in which the bolus should be administered, select by pressing \[\text{Bolus Time} \]. Calculated bolus rate is displayed on the top display line. Press \[\text{OK} \] to confirm and start the bolus.

After the \[\text{Bolus} \] is pressed, the bolus unit can be selected by pressing \[\text{Bolus} \] . This selected unit is stored and used in the future as the default. This means that it is also possible to administer a bolus in mL in the dosage calculation mode.

A default and a maximum bolus rate can be specified via the service program. But the device always returns to the default rate at each startup, even if the bolus rate was previously changed manually.

**Note:** If a bolus cannot be administered after pressing \[\text{Bolus} \] , the pump automatically returns to the delivery display. If a bolus is not completed, the device issues a reminder alarm that has to be confirmed by pressing \[\text{OK} \].

**Note:** For a bolus with a pre-selected volume, the volume counts up.

The pump can be primed at any time by pressing \[\text{Priming} \] while the pump is stopped. Answer the following question by pressing \[\text{Yes} \] in order to start the priming process. Cancel by pressing \[\text{No} \] or any other button.

**Caution:** Do not overdose! At a bolus rate of 1200 mL/h, for example, 1 mL is reached after 3 seconds. Press \[\text{OK} \] to cancel bolus at any time. For low bolus volumes, under dosages cannot be ruled out due to the start-up characteristics of the pump and tolerances in the infusion system. Disconnect patient while priming the system.

1.4 Syringe change and start of new therapy

**Note:** Always disconnect the patient from the device before changing the syringe to prevent an unintended dose. Never leave the pump unattended while replacing the syringe. Before a new syringe is inserted, check the functionality of the green syringe guide.
Press  to stop drug delivery. The green LED disappears. Disconnect patient from device.

Open syringe holder. The drive retracts. If the syringe holder is pulled more than 30 seconds after the pump was stopped, you have to first answer the question of whether to replace a syringe by pressing  before the drive returns (regardless of the time, this always applies in the case of a syringe replacement alarm). If the syringe holder is only briefly pulled (less than a second), answer the question displayed first by pressing  to release the plunger shaft.

Open front cover, remove syringe and insert new one.

**Note:** When removing the syringe, in the event that the plunger plate is not released by the claw, press the emergency release button to unlock the claw on the drive head. The emergency release button is located on the exterior of the drive head. It can be released by pressing with a pointed object (e.g. a pen). The claw can then be opened manually and the syringe removed. The device should then be sent to service for repair.

Close the syringe holder and pump door (**Note:** Plunger brake must move forward) and confirm the syringe type inserted by pressing . Drive moves forward and grabs the plunger plate of the syringe.

**Note:** Do not allow objects to block moving drive. Plunger brake must return automatically to the syringe holder.

If necessary, prime pump by pressing , then press  to continue.

Connect patient and check parameters by pressing .

Begin infusion by pressing .

**To start a new therapy after replacing the syringe:**

If the pump is in the main menu, press .

Press  and enter new therapy parameters by pressing .

Begin infusion by pressing .

**Note:** A new therapy can be started at any time during a stopped infusion. If a pump is in the main status or option menu, press  (repeatedly) and follow the instructions as described.

### 1.5 End of infusion

Press  to stop infusion. The green LED goes out. Disconnect patient from device.

Open syringe holder. Answer question of whether the syringe holder should be opened by pressing . The drive retracts.

Open pump door. Remove syringe, put syringe holder in an upright position and close cover.
Chapter 1

- Press \( \text{①} \) for 3 seconds to turn off pump. Drive automatically parks near the pump housing.

**Note:** The device permanently retains the settings after it is turned off. As long as a syringe is inserted, the "standby mode" is used.

1.6 Standby

In case of a longer delay, the user has the option of retaining the set values.

- Interrupt the infusion by pressing \( \text{②} \). Afterward, press \( \text{①} \) for less than 3 seconds.

- Confirm whether pump should be switched to standby by pressing \( \text{③} \).

- The current setting for standby is displayed. Accept the default setting by pressing \( \text{④} \) or change by pressing \( \text{⑤} \) (0-24 hours) and confirm with \( \text{⑥} \).

- As long as the pump is in "standby," the name of the drug and the remaining time for "standby" are displayed. The remaining time can be changed with \( \text{⑥} \). Exit "standby" with \( \text{②} \).

- As long as a syringe is inserted, the pump remains in "standby" mode. By pressing \( \text{③} \), the system exits "standby." By pressing \( \text{⑥} \), the "standby" duration is entered.
2.1 Querying the pump status during the infusion

While the device is pumping, you can switch between the delivery display and the main menu by pressing \[2\] and navigate through the menu by pressing \[8\] to check the parameters. To check the menu items in the status and options menu, select "status" or "options" in the main menu, select with \[4\] and scroll through the menu \[8\]. All status information is also available, if needed, in the bottom line of the main screen.

2.2 Rate, volume and time change without infusion interruption and resetting of status menu parameters

- If the pump is in the delivery display mode, press \[8\] to switch to the main menu. Select Rate/Volume/Time with \[8\] and press \[4\] to open parameters.
- Enter new value with \[8\] and confirm with \[ok\].

Reset status menu parameters:

The intermediate volume and time parameters can be cleared while the pump is pumping or the infusion is interrupted.

- Select "status" in the main menu by pressing \[8\] and press \[4\].
- Select intermediate volume (in mL) or interim time (in h:min) by highlighting with \[8\] and select parameters by pressing \[4\].
- Reset values by pressing \[ok\].

Both total volume and time parameters are displayed as "total" using the appropriate unit and can be reset by starting a new therapy. Second reset method: If the pump is in the main menu, press \[8\], answer the question concerning the most recent therapy used by pressing \[ok\] and then reset the values with \[ok\].

The type of syringe inserted can be viewed under the menu item "Syringe" and cannot be changed after confirming the syringe item at the beginning of the infusion. The "Drug info" menu lists the drug name, the name of the drug library and its creation date. The current battery capacity is displayed in hours and minutes in the "Battery capacity" menu and the current software version under "Version."
### 3.1 Dosage calculation (Overview)

<table>
<thead>
<tr>
<th>Gram family</th>
<th>$10^6$ ng</th>
<th>$10^3$ mcg</th>
<th>1 mg</th>
<th>$10^{-3}$ g</th>
</tr>
</thead>
<tbody>
<tr>
<td>Unit family</td>
<td>$10^3$ mU</td>
<td>1 U</td>
<td>$10^{-3}$ kU</td>
<td>$10^{-6}$ MU</td>
</tr>
</tbody>
</table>

The dose rate calculation allows the delivery rate to be calculated in mL/h from the dosage parameters entered.

$$\text{Infusion rate} \ [\text{mL/h}] = \frac{\text{Dose}}{\text{Concentration}} \times \frac{[\text{Patient weight} \ (\text{optional})]}{}$$

**Setting parameters:**

1. Concentration as dosage of active ingredient per volume
   - Dosage in μg, mg, mmol, IE or mEq.
   - Volume in mL.
2. If necessary, patient weight in kg or lbs
3. Dose prescription:
   - According to time in dosage per minutes, hour or 24 hours.
   - According to time and patient weight in dosage per kg per minute, hour or 24 hours.

### 3.2 Dose Rate calculation (Application)

- Select dosage calculation with \(\text{Back}^{\text{a}}\).
- Select the unit of the dosage by pressing \(\text{Back}^{\text{b}}\) and confirm with \(\text{OK}\).
- Enter the concentration by entering dosage and volume. Enter the values by pressing \(\text{OK}\) and confirm with \(\text{OK}\).
- If no weight is entered, press \(\text{Down}^{\text{c}}\). For a time- and patient-related calculation, press \(\text{Up}^{\text{d}}\), enter the patient weight with \(\text{OK}\) and confirm with \(\text{OK}\).
- Select the dose prescription by pressing \(\text{Back}^{\text{a}}\) and confirm with \(\text{OK}\).
- Enter the dosage with \(\text{OK}\) and confirm with \(\text{OK}\). The rate is automatically calculated and is displayed at the bottom of the screen.
- Verify the calculated rate displayed and the parameters, if necessary, by pressing \(\text{Back}^{\text{a}}\) prior to start of the infusion with the \(\text{Start}\).

Concentration and dosage can be changed in the main menu at the same time as rate, volume and times are changed (see 2.2). The effects of dosage modifications on other parameters are displayed in the bottom line of the display. The
total and interim quantity of the drug administered can be additionally found in the status menu. These can be checked and reset at the same time as the total and interim values. It is possible to deactivate the dosage calculation in the Stop mode by pressing \( \mathcal{O} \) from the main menu and then pressing \( \mathcal{D} \).

**Caution:** A change in the patient’s weight changes the flow rate!

### 3.3 Drug Library

The drug library contains approximately 720 drug names, including related therapy data and information, in 15 categories. Drug information is entered into the pump via the software program "Drug List Editor Space".

**Note:** The drug library can be started in the startup and the special function menu. Prior to starting the therapy, users must ensure that the drug library stored in the pump matches the target patient group to be treated. The name of the drug library (see headline) is displayed on the pump. There are different options for integrating the drug library into the therapy. Integration can be done while a drug is being administered as well as in the pump’s Stop mode.

A drug name with stored therapy data can be selected from the drug library. Or, if a delivery rate, a volume and/or a time were already defined in the main menu, the drug name and the adapted parameters of the data set will be loaded. Even if drug delivery was already started under a dosage rate calculation, it is possible to assign a drug name retroactively.

The following describes how to select a drug, including the parameters stored with it:

- Open the drug library by pressing \( \mathcal{O} \).
- Navigate through the category list with \( \mathcal{A} \) and select the desired drug from the alphabetical list (all drugs) or within a drug category by pressing \( \mathcal{A} \).
- Confirm displayed drug information, if applicable, with \( \mathcal{A} \).
- Check whether the abbreviated drug name in the main menu refers to the selected drug. Check the parameters in the main menu by pressing \( \mathcal{A} \) and start the infusion with \( \mathcal{A} \).

**Hard Limits:**

If the set rate/dosage/bolus volume and bolus rate exceed the hard limits stored in the database, then the drug is rejected, a message appears and the pump returns to the drug selection screen. If this happens during an infusion, the pump continues to deliver the drug.

**Soft Limits:**

For the same parameters, soft limits can be defined via the Drug List Editor. The
pump display shows the following symbols that describe the pump’s status with regard to the soft limits:

- Infusion is within the minimum and maximum soft limit thresholds
- Infusion is within the maximum soft limit threshold
- Infusion is within the minimum soft limit threshold
- Upper soft limit is exceeded
- Lower soft limit is exceeded
- No soft limit is defined

(only one drug name is present (it is possible to load only one drug name from the drug library)

⚠️ Infusions which display the 🗝 symbol next to the drug name DO NOT have the hard or soft limits from the drug library applied to the infusion. If this symbol is displayed, the pump does not provide additional safeguards against an incorrectly programmed infusion.

The limits of the drug library have to comply with the limits of the pump and the disposable.

Note: For life-supporting drugs, additional patient monitoring is strongly advised.

Note: When a drug is selected from the drug database while the pump is in dosage rate mode, the initial values will be overwritten by those from the drug database.

### 3.4 Instructions for Use of Bar-coding Using DoseScan™ and the SpaceStation with SpaceCom

If a hospital is utilizing bar-coding for patient identification, staff identification, and for labeling of patient medications, it is possible to utilize bar-code scanning in order to perform the following workflow options:

- Scan the barcode containing the clinician’s ID into the Perfusor® Space pump and store it in its log file.
- Scan the barcode containing the patient’s ID into the Perfusor® Space pump and store it in its log file.
- Scan the barcodes containing both the patient’s and the clinician’s ID into the Perfusor® Space pump and store those data elements in its log file.
- Scan the patient’s ID bracelet, then scan the patient’s ID on the medication bag, in order to confirm that the proper medication is being administered to the correct patient (i.e. patient matching) and store both in the log file.
Program the Perfusor® Space pump using the DoseScan™ feature, which uses the patient's ID barcode and the barcodes for all of the therapy parameters (i.e., drug name, concentration, dose) on the medication label, in order to program the infusion therapy. Upon completion of scanning, the clinician needs only to confirm the information and start the infusion.

Bar-code scanning requires the use of the B. Braun SpaceStation with SpaceCom and the bar-coding option. Bar-coding can be utilized for any and all pumps contained in the SpaceStation. If a handheld barcode scanner is attached to the SpaceStation with SpaceCom, when a pump is powered on, the first prompt will be requesting a scan as determined in the pump configuration. If a pump is powered on prior to the barcode scanner being attached to SpaceCom the user will need to select DoseScan™ from the menu.

Physician has to provide a confirmed separate prescription documentation (e.g., Medication Application Record) with text in clear for the patient. During the barcode process the user has to check the displayed therapy data in the confirm dialogue of the infusion pump regarding completeness and correctness.

Before starting the infusion, the user has to ensure that the pump therapy configuration conforms the prescription.

The barcode workflow with the infusion pump supports the medication safety instructions, but the normal workflow (without barcode) must not be changed because of usage of the pump feature barcode.

The pump feature patient matching is a comfort feature, which can be deactivated. This does not excuse from existing medication safety instructions.

It has to be ensured that all infusion pump related medication barcode labels on the container are scanned.

It has to be ensured that only to the medication and container related labels are scanned.

Scanning with Clinician ID, Patient ID and with Drug entry matched to the parameters in Drug Library.

Prior to beginning bar-code scanning, insure that at least one Perfusor® Space pump is docked into the SpaceStation, and that the pump is charged. A handheld barcode scanner must also be plugged into the SpaceStation. If a handheld barcode scanner is attached to the SpaceStation with SpaceCom, when a pump is powered on, the first prompt will be requesting a scan as determined in the pump configuration. If a pump is powered on prior to the barcode scanner being attached to SpaceCom the user will need to select DoseScan™ from the menu.

- Load Perfusor® Space pump into SpaceStation.
  - Power on the pump by pressing 

25
■ Load the proper syringe into the pump and confirm proper loading.

■ The display screen of the Perfusor® Space pump will display a prompt “Scan Nurse ID” in the center of the display.

■ With the handheld scanner attached to the SpaceStation, scan the nurse’s ID. The barcode scanner should beep when the barcode is read successfully. The pump will prompt the user for the next required scan.

  Note: Once the first scan occurs, the bar-code scanning process has time limits built in that are designed to require that the process occurs continuously within a short period of time. Should a clinician be interrupted during the scanning process, the process will automatically be aborted and the screen returned to the first step.

■ Next, on the center of the display the prompt “Scan Patient ID” will appear. Scan the ID barcode on the patient’s wristband. The scanner will beep when the barcode is read successfully. The pump will prompt the user for the next required scan.

■ Next, the screen will display the prompt “Scan IV Label – Patient”. On the medication bag, scan the Patient ID first, then scan the Drug ID label to program all of the patient’s drug therapy. The scanner will beep and all medication information from the barcode label will be loaded into the pump, including the concentration and any dose limits that were included in the drug library information. If, for some reason, the medication label scanned does NOT match the Patient ID previously scanned, the screen will display a message “Wrong Barcode” and an alarm tone will sound. The scanning process will not advance until the proper match is obtained between the Patient ID on the wristband and the Patient ID on the medication.

■ If a desired parameter is missing from the IV label or was not scanned correctly the pump will prompt the user to scan the missing information.

■ If no additional information is available the user can override the request by pressing the directed key.

■ The pump will then display the scanned data for confirmation. If it is configured and if a match was found between the Drug data on the label and drug information within the Drug Library, the Drug name and other scanned parameters will appear.

■ Confirm each scanned element by pressing to verify and complete the scanning process. Finally, press to leave the confirmation menu.

■ Begin the infusion by pressing . The green LED will illuminate, and the arrows graphic will be displayed in the upper right, indicating that the pump is infusing.

Scanning with only Patient ID Scan Activated:

If a handheld barcode scanner is attached to the SpaceStation with SpaceCom,
when a pump is powered on, the first prompt will be requesting a scan as determined in the pump configuration. If a pump is powered on prior to the barcode scanner being attached to SpaceCom the user will need to select DoseScan™ from the menu.

- Load Perfusor® Space pump into SpaceStation. Turn on the pump by pressing \( \text{I}\).
- Load the proper syringe into the pump and confirm proper loading.
- The display screen of the Perfusor® Space pump will display a prompt “Scan Patient ID”.
- With the handheld scanner attached to the SpaceStation, scan the patient’s ID. The barcode scanner should beep when the barcode is read successfully.

**Note:** Once the first scan occurs, the bar-code scanning process has time limits built in that are designed to require that the process occurs continuously within a short period of time. Should a clinician be interrupted during the scanning process, the process will automatically be aborted and the screen returned to the first step.

- Upon completion, press \( \text{OK} \) to verify and complete the scanning process.
- Enter any required information on the therapy, including the Volume to be infused (VTBI) and the rate of infusion or time of infusion. When therapy information is complete, the Start icon will be highlighted in the upper right corner of the display.
- Begin the infusion by pressing \( \text{I} \). The green LED will illuminate, and the arrows graphic will be displayed in the upper right, indicating that the pump is infusing.
- Alternately, if the pump has been configured to use the Drug Library, you will be prompted to select a drug from the drug library, confirm the selection, enter a dose, and then start the infusion.

**Scanning with Patient ID and Patient Matching Activated:**

If a handheld barcode scanner is attached to the SpaceStation with SpaceCom, when a pump is powered on, the first prompt will be requesting a scan as determined in the pump configuration. If a pump is powered on prior to the barcode scanner being attached to SpaceCom the user will need to select DoseScan™ from the menu.

- Load Perfusor® Space pump into SpaceStation. Turn on the pump by pressing \( \text{I} \).
- Load the proper IV set or syringe into the pump and confirm proper loading.
The display screen of the Perfusor® Space pump will display a prompt “Scan Patient ID” in the center of the display.

With the handheld scanner attached to the SpaceStation, scan the Patient’s wristband ID. The barcode scanner should beep when the barcode is read successfully. The pump will prompt the user for the next required scan.

**Note:** Once the first scan occurs, the bar-code scanning process has time limits built in that are designed to require that the process occurs continuously within a short period of time. Should a clinician be interrupted during the scanning process, the process will automatically be aborted and the screen returned to the first step.

Next, on the center of the display the prompt “Scan IV Label-Patient” will appear. Scan the ID barcode on the IV label. The barcode scanner should beep when the barcode is read successfully.

If, for some reason, the Patient ID on the medication label scanned does NOT match the Patient ID previously scanned, the screen will display a message “Wrong Barcode” and an alarm tone will be sound. The scanning process will not advance until the proper match is obtained between the Patient ID on the wristband and the Patient ID on the medication.

Upon completion, press **OK** to verify and complete the scanning process.

Enter any required information on the therapy, including the Volume to be infused (VTBI) and the rate of infusion or time of infusion. When therapy information is complete, the Start icon will be highlighted in the upper right corner of the display.

Begin the infusion by pressing ****. The green LED will illuminate, and the arrows graphic will be displayed in the upper right, indicating that the pump is infusing.
OPTIONS

You can select and change the option functions during therapy or if the infusion is interrupted. To edit a menu item, select “options” in the main menu and press (4). Then select the desired function with (3) and follow the instructions as described.

4.1 Occlusion pressure

The higher the pressure level is set, the higher the system pressure has to rise to trigger a pressure alarm.

- Select pressure in the options menu by pressing (4).
- Select between nine pressure levels (1=lowest level; 9=highest level) by pressing (4) or (5) and confirm with (OK).

4.2 Data lock

The data lock function protects the device from unauthorized access. A four-digit code, which can be changed in the service program, activates this function. There are three lock levels.

Level 1:
Values or bolus cannot be changed, but syringes can be. Navigation is allowed through all menus and status data can be queried. The pump can also be started, stopped and shut down.

Level 2:
This level has the same permissible options as Level 1, but does not permit the syringe to be changed. To prevent a data lock alarm, the user has to enter the correct code within 20 seconds after the pump was stopped. Only after entering the code can the syringe be changed and the pump turned off.

Level 3:
This level allows the pump to be started and stopped as well as shut down. The code for this level can be different for each drug and is defined in the drug library. However, the syringe can be changed with the code that is used for the other levels. The following table lists the different options for Levels 1, 2 and 3:
<table>
<thead>
<tr>
<th>Action/Event</th>
<th>Level 1</th>
<th>Level 2</th>
<th>Level 3</th>
</tr>
</thead>
<tbody>
<tr>
<td>Change syringe</td>
<td>✓</td>
<td></td>
<td>✓ with code for level 1/2</td>
</tr>
<tr>
<td>Infusion start</td>
<td>✓</td>
<td></td>
<td></td>
</tr>
<tr>
<td>Change in the parameters</td>
<td></td>
<td></td>
<td>✓</td>
</tr>
<tr>
<td>Infusion stop</td>
<td>✓</td>
<td>✓</td>
<td></td>
</tr>
<tr>
<td>Pump shut-down/standby</td>
<td>✓</td>
<td></td>
<td>✓</td>
</tr>
<tr>
<td>Customizable Screen</td>
<td></td>
<td></td>
<td>✓</td>
</tr>
<tr>
<td>Audible feedback with bolus request</td>
<td></td>
<td></td>
<td>✓</td>
</tr>
</tbody>
</table>

✓ = Possible | ✗ = Not possible | ⚠= Followed by “standby” alarm

**Activation of the function:**

- Select data lock in the options menu by pressing ⬅.
- With ⬅ and ➤ switch between Level 1, 2 or 3 (if activated) and confirm with OK.
- Enter code with ☢ and press OK to activate the data lock function.

The values as well as the bolus function, which is identified by ☢, can only be changed by entering the code. From the overviews, main status, special functions and options menus, the lock is reactivated after 20 seconds. If the code is incorrectly entered twice, the pump switches back to the last menu. If the code is again entered incorrectly twice, the pump sounds an audible alarm, triggers a nurse call and the yellow LED begins to blink. If the target value is reached while the data lock is active, a restart is only possible after entering the code.

To deactivate the function, select “Off” from the data lock function list, press OK, enter the code and confirm again with OK.

### 4.3 Bolus rate

- Select bolus rate in the options menu by pressing ⬅.
- Change the bolus rate with ☢ and confirm setting with OK.

**Note:** Adjust the bolus rate to the therapy requirements. Take care not to overdose! For very high bolus rates (e.g. 1800 mL/h), 5 mL is administered within just one second.

### 4.4 KVO mode (Keep Vein Open)

After the pre-selected volume to be infused or the pre-selected time has run out, the pump can continue delivering at a pre-defined KVO rate (see Technical data). The duration of the KVO delivery is defined in the service program.
4.5 Contrast / Display Illumination / Keypad Illumination

Contrast, display and keypad illumination can be adjusted individually according to lighting conditions.

- Select contrast/display illumination/keypad illumination in the options menu by pressing \( < \).
- Select between the 9 contrast and display illumination levels with \( < \) or \( > \) and confirm with \( OK \). If light-sensitive drugs are used, you can turn off the keypad or the syringe illumination.

4.6 Alarm Volume

There are 9 different volume levels to choose from.

- Select volume in the options menu by pressing \( < \).
- Confirm volumes with \( < \) or \( > \) and confirm with \( OK \).

4.7 Date / Time

- Open date/time in options menu by pressing \( < \).
- Press \( < \) or \( > \) to change date/time and confirm the setting with \( OK \).

4.8 Macro mode

The delivery rate appears larger on the display when macro mode is activated and the pump is infusing.

- Select macro mode in the options menu with \( < \).
- Answer yes/no questions by pressing \( \uparrow \) to activate macro mode.

Quick selection of the macro mode: While the pump is administering the drug, press \( \uparrow \) and hold down until the font size changes.
The Perfusor® Space is equipped with an audible and optical alarm signal.

<table>
<thead>
<tr>
<th>Alarm type</th>
<th>Audible signal</th>
<th>Optical signal</th>
<th>Staff Call</th>
<th>User confirmation</th>
</tr>
</thead>
<tbody>
<tr>
<td>Device alarm</td>
<td>Yes</td>
<td>flashes</td>
<td>Device alarm and alarm code (see service manual)</td>
<td>Press (\text{OK}) and follow the instruction on the display.</td>
</tr>
<tr>
<td>Operating alarm</td>
<td>Yes</td>
<td>flashes</td>
<td>See alarm description</td>
<td>Press (\text{OK}) to acknowledge the audible alarm, alarm text and nurse call. The red LED remains on until the infusion is restarted.</td>
</tr>
<tr>
<td>Pre-alarm</td>
<td>Yes</td>
<td>off</td>
<td>See alarm description</td>
<td>Press (\text{OK}) to turn off the alarm tone and nurse call.</td>
</tr>
<tr>
<td>Reminder alarm</td>
<td>Yes</td>
<td>off</td>
<td>See alarm description</td>
<td>Press (\text{OK}) to turn off the alarm tone, the nurse call and delete the alarm text.</td>
</tr>
<tr>
<td>Alarm reference</td>
<td>Yes</td>
<td>off</td>
<td>See alarm description</td>
<td>No</td>
</tr>
</tbody>
</table>

### 5.1 Device alarms

In case of a device alarm, the infusion is immediately stopped. Press \(\text{OK}\) to turn off the device. Then turn the device back on. If the device alarm comes on again, disconnect the patient’s IV line, open front cover and remove the syringe. The device must then be serviced.

### 5.2 Pre-Alarms and operating alarm

Pre-alarm:
Pre-alarms occur several minutes (regardless of service settings) prior to the operating alarms. With pre-alarms, an audible signal sounds, the yellow LED blinks and the staff call (optional) is issued. The display varies depending on the cause of the pre-alarm. The signal tone and staff call are turned off by pressing \(\text{OK}\). The screen display and LED remain in pre-alarm mode until the operating alarm initiates. If multiple pre-alarms are triggered simultaneously, the staff call and the audible signal do not go off until the last pre-alarm is exited. Pre-alarms do not cause drug delivery to stop.
### Chapter 5

<table>
<thead>
<tr>
<th>Display message</th>
<th>Cause of pre-alarm</th>
</tr>
</thead>
<tbody>
<tr>
<td>“Syringe almost empty”</td>
<td>Little solution left in the syringe.</td>
</tr>
<tr>
<td>“VTBI almost reached”</td>
<td>Pre-selected volume almost infused.</td>
</tr>
<tr>
<td>“Time near end”</td>
<td>Pre-selected time almost expired.</td>
</tr>
<tr>
<td>“Battery nearly empty”</td>
<td>Battery almost completely discharged.</td>
</tr>
<tr>
<td>“KVO active”</td>
<td>VTBI/time are expired and pump continues to run at KVO rate.</td>
</tr>
<tr>
<td>“Communications error“</td>
<td>The pump is integrated into a system in which one device is incompatible or defective. Operation of this pump is not permitted in the system. The system has to be checked by a service technician.</td>
</tr>
</tbody>
</table>

A countdown clock in the display counts down the remaining pre-alarm time (3 to 30 minutes, depending on service program). Afterward, the pump goes into operating alarm mode.

Pre-alarms “VTBI” almost reached (volume pre-selection) and “Time near end” (time pre-selection) can be (de) activated via the service program.

### Operating alarms:
When operating alarms occur, the infusion stops. An audible signal sounds, the red LED blinks and a nurse call is issued. The display shows “alarm” and the cause of the alarm. The signal tone and nurse call can be turned off by pressing OK. Corrective actions must be implemented based on the cause of the alarm.

<table>
<thead>
<tr>
<th>Display message</th>
<th>Cause of pre-alarm and corrective measures</th>
</tr>
</thead>
<tbody>
<tr>
<td>“Syringe empty”</td>
<td>No remaining infusion solution in the syringe. Due to different manufacturing syringe tolerances, some liquid may remain in the syringe. Restarting results in the syringe being completely emptied and shuts down via the pressure sensor. Change the syringe as described under 1.4.</td>
</tr>
<tr>
<td>“VTBI infused “</td>
<td>Pre-selected “VTBI” is infused. Continue with therapy or select new therapy.</td>
</tr>
<tr>
<td>“Time expired”</td>
<td>Pre-selected time has expired. Continue with therapy or select a new therapy.</td>
</tr>
<tr>
<td>“Battery empty”</td>
<td>The battery is discharged. Connect device to power supply and/or change battery. Battery alarm sounds for 3 minutes. Afterward, the pump shuts itself off.</td>
</tr>
<tr>
<td>“Pressure high“</td>
<td>Occlusion in the system has occurred. The set pressure level has been exceeded. The pump automatically reduces the bolus. Check whether the syringe is empty, make sure the line is free of kinks and undamaged, and that IV and filter are clear.</td>
</tr>
</tbody>
</table>
If applicable, increase the shut-down pressure. Due to different syringe threshold settings, a pressure alarm can go off depending on the high syringe frictional forces.

<table>
<thead>
<tr>
<th>Alarm Description</th>
<th>Description</th>
</tr>
</thead>
<tbody>
<tr>
<td>“KVO finished”</td>
<td>KVO time has expired. Continue with therapy or select new therapy.</td>
</tr>
<tr>
<td>“Syringe not correctly inserted”</td>
<td>The syringe tabs are not properly positioned and syringe not properly secured. Insert as described under “Overview of Perfusor® Space” as well as 1.1.</td>
</tr>
<tr>
<td>“Syringe holder”</td>
<td>The syringe holder was opened during an infusion. Close the syringe holder.</td>
</tr>
<tr>
<td>“Battery cover removed”</td>
<td>The battery cover is not properly attached to the battery compartment. When closing the battery cover, make sure you hear it snap in.</td>
</tr>
<tr>
<td>“Drive blocked”</td>
<td>The drive head was blocked from moving forward due to an external interference. Take care to avoid obstructions.</td>
</tr>
<tr>
<td>“Calibrate device”</td>
<td>Pump calibration data has changed (e.g. after an update). Recalibrate device with service program.</td>
</tr>
<tr>
<td>“Claw malfunction”</td>
<td>The emergency release button was pressed and the claw manually opened. Remove syringe and inform technical service.</td>
</tr>
<tr>
<td>“Plunger plate has no contact”</td>
<td>The syringe’s plunger plate has no contact with the plunger plate sensor. Check system for low pressure and remove the cause of low pressure.</td>
</tr>
<tr>
<td>“Standby time expired”</td>
<td>The standby time has expired. Enter new time or continue with the previous therapy.</td>
</tr>
<tr>
<td>“No battery in the device”</td>
<td>Pump operation without battery is not possible. Turn off pump and insert battery as described in “Overview of Perfusor® Space.”</td>
</tr>
<tr>
<td>“Data was reset”</td>
<td>Therapy and pump settings could not be restored. Re-enter therapy and pump data.</td>
</tr>
<tr>
<td>“Therapy data was reset”</td>
<td>Therapy data could not be restored. Re-enter therapy.</td>
</tr>
<tr>
<td>“Data lock”</td>
<td>Someone attempted to stop or turn off the pump without entering the proper code. Enter the correct code to continue with the therapy or to turn off the pump. The red LED does not go off until drug delivery begins or after the pump is turned off.</td>
</tr>
</tbody>
</table>
Caution: If a wrench appears and/or the yellow, red and blue LEDs blink, this means that the pump is in service mode and is cannot be used on the patient. The pump must be checked by a service technician.

5.3 Reminder alarms

Reminder alarms occur in two cases:

1. A syringe is inserted, the pump does not deliver, no value is edited and the device is not operated for two minutes.
   A signal tone sounds, the yellow LED blinks and a staff call is issued.
   a) The display shows "Reminder alarm!"
   b) The display shows "Configuration not completed!"
   Confirm alarm with (OK) and continue by pressing therapy entry/start-up.

2. Value entry was begun, but not completed and confirmed. This is also possible in the case of a missing syringe. A signal tone sounds, the display shows "Value not accepted," the yellow LED blinks and a staff call is issued. Confirm alarm by pressing (OK) and continue with therapy entry.

5.4 Alarm instructions

In case of incorrect entries, the display will show relevant instructions (e.g. "Caution: rate is out of range" or "The parameter cannot be changed") and a signal tone sounds. These warnings disappear after a few seconds without having to be confirmed.
BATTERY OPERATION AND MAINTENANCE

The Perfusor® Space is equipped with a NiMH-battery that has an operating life of 8 hours at a 25 ml/h delivery rate. For optimal life of the battery, the device is equipped with protection against overcharging and over-discharging. The device charges the battery when plugged into the wall outlet. In case of a power outage or removing from the power supply, the pump automatically switches over to battery operation.

Note: If the pump is to be stored unused for a longer period (> 2 weeks), the battery should be completely charged and then removed from the pump. Before removing the battery, the patient has to be disconnected and the pump shut off.

The battery status display indicates the battery power (low, medium, high). More detailed information on the battery capacity (operating time in hours and minutes) can be found in the menu item "Battery capacity" in the status menu of the Perfusor® Space.

Important information for the battery self-test:

If the battery status symbol blinks while operating from the wall outlet, the battery is either discharged or used up. In this case, the pump should not be disconnected from the outlet. If the pump has to be disconnected from power because of an emergency, check to make sure that the battery capacity is sufficient for intended use. If the battery status icon blinks continuously (> 1 h), the battery must be checked by a technician and replaced if necessary.

Tips for optimal battery operation:

The service life of a battery can vary due to

- Ambient temperature
- Varying load (e.g. frequent boluses).

You can extend the battery’s service life by discharging it and then fully recharging it. For this purpose, the pump has to be operated in battery maintenance mode. To charge the battery, the pump has to be connected to the power supply for at least 6 hours. This procedure should be performed once a month. Moreover, please observe the following:

- If possible, only charge the battery if it was completely discharged.
- If a battery that was not completely discharged is charged multiple times, this then reduces its capacity. Its original capacity can be achieved again by completely discharging the battery and then recharging it.
- Under normal temperature conditions, a battery can be charged 500 times on average and then discharged again until its service life decreases.
- If the pump is not connected to the wall outlet, the battery discharges and can be fully depleted after a month, even when the device is not in operation. If this happens, the battery cannot reach its original capacity after just one charge, rather it will take multiple cycles of charging and discharging.
- Optimal battery life can only be achieved if the battery is kept charged and the pump is in continual operation at room temperature. The pump’s battery display is an approximate value based on the current delivery rate.
If the battery begins to show signs of fatigue, its "battery display" can differ from the current achievable operating time. **Caution:** Batteries can explode or leak if they are opened or burned. Therefore, please follow appropriate disposal regulations in your institution.

**Battery maintenance:**

To accurately maintain battery capacity, routine battery maintenance is necessary. It is advised for the pump that battery maintenance be performed every 30 days as a standard procedure. Routine battery maintenance can determine possible capacity loss (e.g. through battery's aging) and rebalances capacity/duration. To initiate the discharge procedure, after turning off the pump, the "Battery Maintenance" display and 🔄-button will appear. By pressing OK and 🔄, discharge is initiated. If the pump is turned on again, this procedure is interrupted. To continue battery maintenance, it has to be reactivated. After complete discharge, the battery must be completely recharged again. The entire battery maintenance procedure takes about 12 hours. **Caution:** When pump is started, please make note of the shortened battery run-time in the event that the battery maintenance procedure was not completed.
COMPATIBLE SYRINGES

The syringe types listed in the following tables can be used with the Perfusor® Space. Because B. Braun has no influence on the quality of syringes offered from other manufacturers. Quality inconsistencies can lead to changes in the technical properties of the pump. B. Braun is not liable for deviations caused by the use of a syringe from manufacturers other than B. Braun’s approved table. In case of such quality fluctuations, please contact the syringe manufacturer.

Please refer to the listed item number (Item No.1) to be sure of specific brand compatibility.

The alarm actuation times2 after a system breech were measured at 5mL/h. All measured data in the tables are typical average values that can deviate above and below due to possible syringe variations.

Manufacturer:
B. Braun

<table>
<thead>
<tr>
<th>Syringe Type</th>
<th>Omnifix™ 2/3 mL</th>
<th>Omnifix™ 5 mL</th>
<th>Omnifix™ 10 mL</th>
<th>Omnifix™ 20 mL</th>
<th>Omnifix™ 30 mL</th>
<th>Omnifix™ 50 mL</th>
</tr>
</thead>
<tbody>
<tr>
<td>Mat. no.1)</td>
<td>4610303-02</td>
<td>4617053V-02</td>
<td>4617100V-02</td>
<td>4617207V-02</td>
<td>4617034F-02</td>
<td>4617509F-02</td>
</tr>
<tr>
<td>Time to Occl.2)</td>
<td>typ.</td>
<td>typ.</td>
<td>typ.</td>
<td>typ.</td>
<td>typ.</td>
<td>typ.</td>
</tr>
<tr>
<td>P 1 [mm:ss]</td>
<td>0:39</td>
<td>0:58</td>
<td>0:47</td>
<td>1:04</td>
<td>1:13</td>
<td>1:16</td>
</tr>
</tbody>
</table>

Manufacturer:
Covidien AG

<table>
<thead>
<tr>
<th>Syringe Type</th>
<th>Monoject® 3 mL</th>
<th>Monoject® 6 mL</th>
<th>Monoject® 12 mL</th>
<th>Monoject® 20 mL</th>
<th>Monoject® 35 mL</th>
<th>Monoject® 50/60 mL</th>
</tr>
</thead>
<tbody>
<tr>
<td>Mat. no.1)</td>
<td>1180300777</td>
<td>8881-516937</td>
<td>8881-512878</td>
<td>8881-520657</td>
<td>8881-535762</td>
<td>8881-560125</td>
</tr>
<tr>
<td>Time to Occl.2)</td>
<td>typ.</td>
<td>typ.</td>
<td>typ.</td>
<td>typ.</td>
<td>typ.</td>
<td>typ.</td>
</tr>
<tr>
<td>P 1 [mm:ss]</td>
<td>0:41</td>
<td>0:50</td>
<td>1:07</td>
<td>1:13</td>
<td>1:27</td>
<td>1:35</td>
</tr>
</tbody>
</table>

Dead volume of Monoject 3 mL (1180300777) is 0.46 mL.
## Chapter 7

### Manufacturer:
**Becton Dickinson**

<table>
<thead>
<tr>
<th>Syringe Type B-D EU/USA</th>
<th>Plastipak® 3 mL</th>
<th>Plastipak® 5 mL</th>
<th>Plastipak® 10 mL</th>
<th>Plastipak® 20 mL</th>
<th>Plastipak® 30 mL</th>
<th>Plastipak® 50/60 mL</th>
</tr>
</thead>
<tbody>
<tr>
<td>Mat. no.¹)</td>
<td>309585</td>
<td>309603</td>
<td>309604</td>
<td>309661</td>
<td>309650</td>
<td>309653</td>
</tr>
<tr>
<td>Time to Occl.²)</td>
<td>typ.</td>
<td>typ.</td>
<td>typ.</td>
<td>typ.</td>
<td>typ.</td>
<td>typ.</td>
</tr>
<tr>
<td>P 1 [mm:ss]</td>
<td>0:53</td>
<td>0:55</td>
<td>1:15</td>
<td>2:05</td>
<td>2:14</td>
<td>2:53</td>
</tr>
<tr>
<td>P 9 [mm:ss]</td>
<td>1:15</td>
<td>1:34</td>
<td>3:37</td>
<td>6:30</td>
<td>6:36</td>
<td>15:34</td>
</tr>
</tbody>
</table>

### Manufacturer:
**TERUMO**

<table>
<thead>
<tr>
<th>Syringe Type TERUMO USA</th>
<th>3 mL</th>
<th>5 mL</th>
<th>10 mL</th>
<th>20 mL</th>
<th>30 mL</th>
<th>60 mL</th>
</tr>
</thead>
<tbody>
<tr>
<td>Mat. no.¹)</td>
<td>3SS*03L</td>
<td>3SS*05L</td>
<td>3SS*10L</td>
<td>3SS*20L</td>
<td>1SS*30L</td>
<td>3SS*60L</td>
</tr>
<tr>
<td>Time to Occl.²)</td>
<td>typ.</td>
<td>typ.</td>
<td>typ.</td>
<td>typ.</td>
<td>typ.</td>
<td>typ.</td>
</tr>
<tr>
<td>P 1 [mm:ss]</td>
<td>0:43</td>
<td>0:35</td>
<td>0:55</td>
<td>2:12</td>
<td>2:25</td>
<td>3:34</td>
</tr>
<tr>
<td>P 9 [mm:ss]</td>
<td>1:17</td>
<td>1:16</td>
<td>4:48</td>
<td>7:53</td>
<td>8:18</td>
<td>17:03</td>
</tr>
</tbody>
</table>
START UP GRAPHS AND TRUMPET CURVES

These graphics show the accuracy/uniformity of the flow, depending on the time. The delivery behaviour or delivery precision is influenced by the type of the disposable used. Deviations from the technical data of the pump cannot be excluded if lines (disposables) other than B.Braun approved components are used.

**Trumpet Curves**

- Each value measured in the second hour.
- Measurement interval $\Delta t = 0.5$ min
- Monitoring interval $p \times \Delta t$ [min]

**Start-up Curves**

- Measurement interval $\Delta t = 0.5$ min
- Measurement duration $T = 120$ min
- Flow $Q_i$ (mL/h)
# TECHNICAL DATA

<table>
<thead>
<tr>
<th>Device type</th>
<th>Infusion Syringe Pump</th>
</tr>
</thead>
<tbody>
<tr>
<td>Classification (according to IEC/EN 60601-1)</td>
<td>☑️ defibrillation-protected; Type CF ☐ protection rating II; protection rating I combined with SpaceStation</td>
</tr>
<tr>
<td>Class (in accordance to guideline 93/42 EEC)</td>
<td>IIb</td>
</tr>
<tr>
<td>Moisture protection</td>
<td>IP 22 (protected from drip water if positioned horizontally for general use)</td>
</tr>
<tr>
<td>External power supply:</td>
<td>via B. Braun SpaceStation or optional with separate power cord (rated voltage 100 ... 240 V AC- , 50/60 Hz) for standalone operation</td>
</tr>
<tr>
<td>■ Rated voltage</td>
<td>11 ... 16 V DC === via Connection cable SP 12 V or via SpaceStation</td>
</tr>
<tr>
<td>■ External low voltage</td>
<td></td>
</tr>
<tr>
<td>Staff call</td>
<td>Max. 24 V / 0.5 A / 24 VA (VDE 0834)</td>
</tr>
<tr>
<td>EMC</td>
<td>IEC/EN 60601-1-2 / 60601-2-24</td>
</tr>
<tr>
<td>Time of operation</td>
<td>100 % (continuous operation)</td>
</tr>
<tr>
<td>Operating conditions:</td>
<td></td>
</tr>
<tr>
<td>■ Relative humidity</td>
<td>30 % ... 90 % (without condensation)</td>
</tr>
<tr>
<td>■ Temperature</td>
<td>+41 ... +104° F</td>
</tr>
<tr>
<td>■ Atmospheric pressure</td>
<td>500 ... 1060 mbar</td>
</tr>
<tr>
<td>Storage conditions:</td>
<td></td>
</tr>
<tr>
<td>■ Relative humidity</td>
<td>20 % ... 90 % (without condensation)</td>
</tr>
<tr>
<td>■ Temperature</td>
<td>-4 ... +131° F</td>
</tr>
<tr>
<td>■ Atmospheric pressure</td>
<td>500 ... 1060 mbar</td>
</tr>
<tr>
<td>Battery type (re-chargeable)</td>
<td>NiMH</td>
</tr>
<tr>
<td>Battery run-time</td>
<td>Approx. 8 hours at 25 mL/h</td>
</tr>
<tr>
<td>Recharge time</td>
<td>Approx. 6 hours</td>
</tr>
<tr>
<td>Weight</td>
<td>Approx. 3.08 lbs. = Approx 1.4 kg</td>
</tr>
<tr>
<td>Dimensions (W x H x D)</td>
<td>Approx 9.8X2.6X5.9 inches = Approx 249 x 68 x 152 mm</td>
</tr>
<tr>
<td>Pre-selected volume</td>
<td>0.1 – 99.99 mL in increments of 0.01 mL 100.0 – 999.0 mL in increments 0.1 mL 1,000 – 9,999 mL in increments 1 mL</td>
</tr>
<tr>
<td>Pre-selected time</td>
<td>00:01 – 99:59 h</td>
</tr>
<tr>
<td>Delivery accuracy</td>
<td>± 2 % according to IEC/EN 60601-2-24</td>
</tr>
</tbody>
</table>
Occlusion alarm button 9 stages from 75 mmHg to 900 mmHg

Alarm sounds in case of incorrect delivery In case of incorrect dosage of 0.1 mL due to a pump malfunction, the pump turns off automatically

Technical inspection (Safety check) Every 2 years

Adjustable delivery rates

<table>
<thead>
<tr>
<th>Syringe sizes</th>
<th>Cont. Rates*</th>
<th>Bolus Rates</th>
</tr>
</thead>
<tbody>
<tr>
<td>[mL]</td>
<td>[mL/h]</td>
<td>[mL/h]</td>
</tr>
<tr>
<td>50/60</td>
<td>0.01 – 200</td>
<td>1 – 1800</td>
</tr>
<tr>
<td></td>
<td>Optional</td>
<td></td>
</tr>
<tr>
<td></td>
<td>0.01 – 999.9</td>
<td></td>
</tr>
<tr>
<td>30/35</td>
<td>0.01 – 100</td>
<td>1 – 1200</td>
</tr>
<tr>
<td>20</td>
<td>0.01 – 100</td>
<td>1 – 800</td>
</tr>
<tr>
<td>10/12</td>
<td>0.01 – 50</td>
<td>1 – 500</td>
</tr>
<tr>
<td>5/6</td>
<td>0.01 – 50</td>
<td>1 – 300</td>
</tr>
<tr>
<td>2/3</td>
<td>0.01 – 25</td>
<td>1 – 150</td>
</tr>
</tbody>
</table>

Delivery rates increments 0.01* – 99.99 mL/h in stages from 0.01 mL/h
100.0 – 999.9 mL/h in stages from 0.1 mL/h

Delivery accuracy with bolus typ. ± 2 %

Max. bolus volume according to bolus reduction ≤ 0.2 mL

KVO rate
Rate ≥ 10 mL/h: KVO rate 3 mL/h
Rate < 10 mL/h: KVO rate 1 mL/h
Rate < 1 mL/h: KVO rate = set rate (factory default setting 0.1 mL/h)

PC (Computer) connection USB connection with B. Braun port cable (8713230) with galvanic separation. Please pay attention to safety instructions.

History log 1,000 most recent history entries
100 events for system diagnosis
You can find more information from the separate documents on the History Viewer

*Factory pre-setting allows delivery rates to be entered starting at 0.1 mL/h.

Caution: If a wrench is displayed and/or the yellow, red and blue LEDs blink, then the pump is in the service mode and cannot be used on a patient. The pump must then be checked by a service technician.
Use only compatible combinations of equipment, accessories, working parts and disposables with luer lock connectors.

Only use combined with approved devices/accessories by the manufacturer, otherwise this may lead to higher emission or reduced immunity.

Dosing families

The following table shows the dosing units of the gram and unit family and their conversion used in the pump:

<table>
<thead>
<tr>
<th>Gram family</th>
<th>$10^6$ ng</th>
<th>$10^3$ mcg</th>
<th>1 mg</th>
<th>$10^{-3}$ g</th>
</tr>
</thead>
<tbody>
<tr>
<td>Unit family</td>
<td>$10^3$ mU</td>
<td>1 U</td>
<td>$10^{-3}$ kU</td>
<td>$10^{-6}$ MU</td>
</tr>
</tbody>
</table>

Equivalents family = mEq
Mole family = mmol
Kilocalorie family = kcal
Milliliter family = ml

In addition to these dosing units the user can choose:

\[
\text{Infusion rate [ml/h]} = \frac{\text{Dose}}{\text{Concentration}} \times \left[ \text{Patient weight (optional)} \right]
\]
Training

B. Braun offers training and in-servicing. Please ask your local representative for more details.

Technical Safety Check* / Service

The technical safety check is required every two years, and should be documented. Service work may only be performed by technicians authorized or trained by B. Braun.
Routine Checks

Check regularly for cleanliness, completeness and signs of damage. Operate according to this Instruction for Use. The pump has to run through the power-on test each time a syringe is changed. When turning on, check self-check, alarm sound, operating and alarm check displays.

Cleaning and Disinfecting

⚠️ Caution: Before disinfecting the pump, always disconnect the pump from the patient, switch off the device and disconnect from power and other devices (e.g. staff call).

Clean external surfaces of the Perfusor® Space with using mild soap suds. The housing of Perfusor® Space can be disinfected using 1-Propanol or didecyl dimethyl ammonium chloride. Do not spray disinfectants directly on the pump, use a soft, low lint cloth dampened but not saturated with product. After cleaning allow device to dry for at least 20 minutes prior to use.

Note: Do not use Hexaquart®, other alkylamine containing disinfectants or chloride products (bleach).

Note: Keep instrument upright and do not allow any part of instrument to become saturated with or submerged in fluid during cleaning operation.

Do not allow moisture or detergents to come into contact with the electrical connections of the device (P2 or P3 connectors) or any device openings. To reduce the likelihood of moisture ingress into the electrical connectors, the P2 connector of a power supply or combi cable may be used to cover the connections during cleaning operations. Ensure that any connectors used to cover are not connected to a wall outlet or other electrical source. Once the cleaning has been completed, remove the connector and inspect all connectors for residual moisture and evidence of damage or breakdown to the plating on the connectors. Allow any residual moisture to evaporate before plugging the device into a wall outlet. Replace any connectors which exhibit damage or evidence of plating breakdown prior to returning the device to service. Utilize electrical contact cleaner that does not react with plastics to remove any deposits of material which may be present inside the electrical connectors as required.

⚠️ Caution: Do not allow liquids to enter into or come into contact with any openings or electrical connections on the pump or power supply. Fluid exposure in these areas may result in the risk of short circuit, corrosion or breakdown of sensitive electrical components, and/or electrical shock. If fluid exposure occurs, the device should be swapped out with another device in a manner that presents minimal interruption
to patient care. The device should remain unplugged until it can be inspected by a trained technician for any evidence of damage and/or residual moisture which may impair the function of the device.

Note: The use of unapproved cleaners and failure to follow the disinfection procedures and the manufacturer's recommended dilutions can result in an instrument malfunction or product damage and could void the warranty.

Disposal

B. Braun accepts pumps as well as batteries for proper disposal. To dispose of syringe as well as infusion solutions, follow applicable hygiene and disposal regulations provided by your institution.

Inspection on Delivery

Despite careful packaging, the risk of damage during transport cannot be entirely prevented. Upon delivery, please check that all items are present. Do not use a damaged device.

The device should be tested for proper functioning before initial use.

A respective form can be obtained from B. Braun.

Included in Delivery

Perfusor® Space, SP battery pack, Instructions for Use.
INSTRUCTIONS FOR USE ACCESSORIES

SpaceStation (8713140U)

Station that can hold up to four B. Braun Space pumps. For more information, please refer to the SpaceStation user manual.

SpaceCover Comfort (8713145U)

Cover for placing on the top of SpaceStation, includes center carrying handle and includes a central alarm management and alarm LED.

PoleClamp SP (8713130)

Maximum three B. Braun Space pumps can be stacked on top of one another and secured and transported with the SP pole clamp. To securely attach the SP pole clamp, please observe the “Perfusor® Space Overview” and “Patient Safety.”

Power Supply SP (8713112D)

The Power Supply SP can supply power for a single pump.

1.) Connect P2 plug of Power Supply SP with P2 socket on back of pump (ensure that plug is inserted straight and “clicks”).

2.) Push power plug into wall outlet.

Note: To disconnect plug from pump, firmly grip connector and pull plug straight away from the pump. DO NOT PULL ON THE CORD TO REMOVE THE PLUG CONNECTOR.

A maximum of three pumps may be stacked and connected to one plug with the use of the Combi Lead (8713133).

Technical Data: 100 – 240V AC~, 50/60 Hz, 0.4–0.2A

Combi Lead SP Cable 12 V (8713133)

Use the Combi lead SP combination cable to connect up to three pumps. All pumps can be operated via the SP connection line (12 V).

1.) Connect the combination cable plug to the P2 socket on the back of the pump.

2.) Connect the SP connection line to the SP combination cable.

3.) Plug connector of the SP connection line into the 12 V socket.
Note: A maximum of three plugs can be plugged into the P2 socket on top of one another.

Battery Pack incl. Pin (NiMH) (8713180A)

For more information on the SP battery pack, see "Battery Operation."

Interface Lead CAN SP (8713230)

The Interface Lead CAN SP is needed to establish a connection between the SpaceStation/Pump and the computer output for servicing.

1. Plug connector into the F2 socket on the SpaceStation or P2 on the pump and connect this with the CAN/USB converter.
2. Connect the CAN/USB converter with the computer output as described in the relevant user manual.

Caution: Interface Lead CAN SP is only meant for servicing; do not use during patient application.

Note: A maximum of three connectors can be connected on top of one another in the P2 socket.

Connection Lead for Staff Call SP (8713232)

The SP nurse call connection line has to be used to connect the Perfusor® Space to a call system. The call system must comply with the provisions of the VDE 0834 (note country-specific regulations).

Note: Prior to each application, test the nurse call signal to make sure it works. The Perfusor® Space offers three different nurse call operating modes. They are shown in the signalization diagram. When selecting an operating mode, pay attention to the technology of the nurse call system. The operating mode can be set via the service program.
Caution: Because the staff call can fail and may remain unnoticed, the user is responsible for monitoring the local alarms. (Staff call is not checked during the pump self-test.)

Note: A maximum of three connectors can be plugged into the P2 socket on top of one another.

Technical Data

<table>
<thead>
<tr>
<th>Connecting Wire</th>
<th>Alarm</th>
<th>Operation</th>
</tr>
</thead>
<tbody>
<tr>
<td>White and Green</td>
<td></td>
<td></td>
</tr>
<tr>
<td>White and Brown</td>
<td></td>
<td></td>
</tr>
<tr>
<td>disconnected</td>
<td></td>
<td>connected</td>
</tr>
<tr>
<td>connected</td>
<td></td>
<td></td>
</tr>
</tbody>
</table>

Polarity of connections is arbitrary:
max. 24 V / 0.5 A / 12 VA
B. Braun Perfusor® Space (100 – 240 V)..............................871 3030U

Recommended accessories for the Perfusor® Space:
SpaceStation .................................................................8713140U
SpaceCover Comfort .....................................................8713145U
Pole Clamp SP ...............................................................8713130
Power Supply SP US III ...................................................8713112D
Combi Lead SP Cable 12 V ................................................8713133
Battery Pack incl. Pin (NiMH) ...........................................8713180A
Interface Lead CAN SP .....................................................8713230
Connection Lead for Staff Call SP .................................8713232

Omnifix™ Syringes:
Omnifix™ 50/60 mL Luer Lock .......................................4617509F-02
Omnifix™ 30 mL Luer Lock ..............................................4617309F-02
Omnifix™ 20 mL Luer Lock ..............................................4617207V-02
Omnifix™ 10 mL Luer Lock ..............................................4617100V-02
Omnifix™ 5 mL Luer Lock ................................................4617053V-02
Omnifix™ 2/3 mL Luer Lock .............................................4610303-02

Microbore tubing with male and female luer locks:
36", 0.023" ID. Priming volume: 0.3 mL .....................V6510
60", 0.023" ID. Priming volume: 0.4 mL .....................V6512
60", 0.023" ID. with 22 ga. Protected Needle.
Priming Volume: 0.4 mL ..............................................V6512-22
60", 0.050" ID. Priming volume: 2.5 mL.
Use with higher flow rates and viscous fluids: ............V6516

Microbore Extension sets with PVC free fluid path,
male and female luer locks:
36", 0.02" ID. Priming volume: 0.3 mL .......................V6200
36", 0.03" ID. Priming volume: 0.6 mL
Use with higher flow rates and viscous fluids: ..........V6230
60", 0.02" ID. Priming Volume: 0.5 mL .......................V6212
60", 0.02" ID with 22 ga. Protected Needle.
Priming Volume: 0.5 mL ..............................................V6212-22
60", 0.003" ID. Priming volume: 0.8 mL.
Use with higher flow rates and viscous fluids: ..........V6213
60", 0.03" ID. Priming volume: 1.3 mL with 0.2 micron filter ....V6215
Technical Support

If the pump fails to respond to the operating or troubleshooting procedures listed in this Instructions for Use and the cause cannot be determined, discontinue use and forward it to an authorized B. Braun Service Center.

Should it be necessary to return the pumps for repair, contact Technical Support at B. Braun Customer Support at (800) 627-PUMP (7867). A Returned Materials Authorization number will be provided. Carefully pack up the pump (preferably in the original packing), and ship to the address below. B. Braun can not assume any responsibility for loss or damage to returned instruments while they are in transit.

Service and product performance information, operation training, service training, and service manuals may be obtained from the manufacturer by contacting:

B. Braun Medical Inc.
1601 Wallace Drive, Suite 150
Carrollton, TX 75006
Attn: Service Manager
or call (800) 627-PUMP (7867)

Product complaints may be sent to the Quality Assurance Manager at the above address.

With each complaint, please include:

- the pump’s serial number and software revision,
- a description of the difficulty experienced,
- the pressure limit setting,
- the rate setting,
- the initial volume(s) to be infused,
- the amount of time between the start of the infusion and the time the difficulty was noticed,
- the message displayed at the time the difficulty occurred,
- the catalog and lot number of the set(s) in use,
- the diagnostic code (if applicable), and
- any other information which may aid in the investigation of the complaint.

Authorization to return products must be received from B. Braun prior to shipment. Please contact Customer Service at the above phone number for a Returned Materials Authorization Number.

Clinical Support

The customer may speak with a Registered Nurse for the clarification of operating instructions or clinical applications for the Space/Pump/etc. A (Clinical Support Specialist) Nurse Consultant may be reached at (800) 854-6851.