It is recommended that all pumps at your care unit are equipped with the same software version.
The availability of the listed features is depending on the configuration of the pump. **Technical Safety Check.**

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**PERFUSOR® SPACE OVERVIEW**

**Arrow up and down**
Scroll through menus, change setting of numbers from 0-9, answer Yes/No questions.

**Arrow left and right**
Select data from a scale and switch between digits when numbers are entered. Open a function while pump is running or stopped with the left arrow key.

- **Yellow LED:** Pre-alarm, reminder alarm
- **Green LED:** Infusing
- **Red LED:** Operating or device alarm
- **Blue LED:** Initiating connection to wireless battery or Space Station

- **Press to reset single values to zero and switch back to the previous screen/menu level.**
- **Drive head with claws to hold the syringe plunger plate and emergency release button.**

- **Press to initiate bolus.**
- **Press to turn pump on/off.**

- **Press to Start/Stop infusion.**
- **Syringe holder locks syringe in position. The drive will automatically move back.**

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

- **Port P2 for power supply, SpaceStation, connection lead (12V), combi lead and further accessory leads (staff call, service)**

- **Port P3 for future options**
Syringe Fixation
Pull and turn the syringe holder to the right to open the green axial fixation (see red arrow). Syringe must be fixed with wings upright in the slot (found to the left hand side of the axial fixation) before closing syringe holder. Make sure that syringe is properly inserted.

Caution: Don’t touch piston brake when moving forward.

Fixation of PoleClamp (Universal Clamp)
Line up bar of pump with bar of PoleClamp and slide PoleClamp forward until locking mechanism clicks. To disconnect, push green locking buttons of top pump device and slide bottom pump forward.

Caution: Avoid external mechanical action.

Transport
A maximum of three pumps (Perfusor® Space or Infusomat® Space) plus one SpaceControl may be stacked together (in ambulance cars or helicopters only one pump). Avoid external mechanical influence.

Locking Devices Together
Line up the bar of the lower pump with the bar of the pump above and slide the lower pump backwards until the lock clicks and the green buttons are above each other. To disconnect, push green locking buttons of top pump device and slide bottom pump forward.

Caution: Avoid external mechanical action.

Pole Fixation
Push the opening of PoleClamp against the vertical pole and lock the screw tightly. Unscrew to release. For vertical position push lever down and rotate either way until lever clicks into notch. Push lever for rotation.

Caution: A maximum of three B. Braun Space pumps can be stacked together only in horizontal pump position when used with the PoleClamp SP.
### SYMBOLS ON PRODUCT

<table>
<thead>
<tr>
<th>Symbol</th>
<th>Explanation</th>
</tr>
</thead>
<tbody>
<tr>
<td><img src="image" alt="Mandatory action" /></td>
<td>Mandatory action: see instruction for use.</td>
</tr>
<tr>
<td><img src="image" alt="See accompanying documents" /></td>
<td>See accompanying documents.</td>
</tr>
<tr>
<td><img src="image" alt="Type CF unit with defibrillation protection" /></td>
<td>Type CF unit with defibrillation protection</td>
</tr>
<tr>
<td><img src="image" alt="Protection class II device" /></td>
<td>Protection class II device</td>
</tr>
<tr>
<td><img src="image" alt="Symbol indicating separate collection for electrical and electronic equipment" /></td>
<td>Symbol indicating separate collection for electrical and electronic equipment (2002/96/EC) only for valid for Europe, not applicable for US</td>
</tr>
<tr>
<td><img src="image" alt="CE 0123" /></td>
<td>CE mark compliant to Directive 93/42/EEC</td>
</tr>
<tr>
<td><img src="image" alt="Temperature Limit" /></td>
<td>Temperature Limit</td>
</tr>
<tr>
<td><img src="image" alt="Moisture Limit" /></td>
<td>Moisture Limit</td>
</tr>
<tr>
<td><img src="image" alt="Limitation of the atmospheric pressure" /></td>
<td>Limitation of the atmospheric pressure</td>
</tr>
<tr>
<td><img src="image" alt="Non-ionizing electromagnetic radiation" /></td>
<td>Non-ionizing electromagnetic radiation</td>
</tr>
<tr>
<td><img src="image" alt="General warning sign" /></td>
<td>General warning sign (e.g. Caution)</td>
</tr>
<tr>
<td><img src="image" alt="Unsafe symbol" /></td>
<td>Unsafe symbol (Do not use in MRI environment)</td>
</tr>
<tr>
<td><img src="image" alt="Batch number" /></td>
<td>Batch number</td>
</tr>
<tr>
<td><img src="image" alt="Serial number" /></td>
<td>Serial number</td>
</tr>
<tr>
<td><img src="image" alt="Catalogue number" /></td>
<td>Catalogue number</td>
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<tr>
<td><img src="image" alt="Manufacturer" /></td>
<td>Manufacturer</td>
</tr>
<tr>
<td><img src="image" alt="Date of manufacture" /></td>
<td>Date of manufacture</td>
</tr>
</tbody>
</table>
PATIENT SAFETY

Intended use

The Perfusor® Space Infusion Syringe Pump System includes an external transportable electronic infusion syringe pump and pump accessories. The system is intended for use on adults, pediatrics, and neonates for the intermittent or continuous delivery of parenteral and enteral fluids through clinically accepted routes of administration. These routes include, but are not limited to intravenous, intra-arterial, subcutaneous, epidural, and enteral. The system is used for the delivery of medications indicated for infusion therapy including but not limited to drugs like anesthetics, sedatives, analgesics, catecholamines, anticoagulants etc.; blood and blood components; Total Parenteral Nutrition (TPN); lipids, and enteral fluids. The Perfusor® Space Infusion Syringe Pump System is intended to be used by trained healthcare professionals in healthcare facilities, home care, outpatient, and medical transport environments.

Using TCI the scope of patients is:

<table>
<thead>
<tr>
<th>Parameter</th>
<th>Minimum</th>
<th>Maximum</th>
</tr>
</thead>
<tbody>
<tr>
<td>Weight [kg]</td>
<td>30</td>
<td>200</td>
</tr>
<tr>
<td>Height [cm]</td>
<td>130</td>
<td>220</td>
</tr>
<tr>
<td>Age [Yrs]</td>
<td>16 (Propofol and Remifentanil)</td>
<td>100</td>
</tr>
<tr>
<td></td>
<td>14 (Sufentanil)</td>
<td></td>
</tr>
</tbody>
</table>

Some parameter sets are using the Lean Body Mass (LBM) to individualize the parameterization. The LBM calculation may furthermore restrict the scope of patients as it will not allow TCI for obese patients.

Using TCI the scope of procedures is:

- Propofol: Anaesthesia and Conscious Sedation
- Remifentanil: Anaesthesia
- Sufentanil: Anaesthesia

The medical specialist must decide on suitability for application on the basis of the warranted properties and the technical data.

Operation

- The initial training of the Perfusor® Space is to be performed by B. Braun sales personnel or other authorized persons.
- After each software update, the user is required to inform himself of the changes to the device and accessories by referring to the Instructions for Use.

⚠️ Caution: Ensure the unit 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.

- Prior to administration, visibly inspect the pump and the accessories (especially the axial fixation) for damage, missing parts or contamination and check audible and visible alarms during selftest.
Not be used adjacent and stacked with other equipment except B. Braun Space devices.

Only connect to patient once the syringe has been inserted correctly and there is proper fixation of the syringe pressure plate by the claws of the drive head. Interrupt connection during syringe change to prevent incorrect dose delivery.

Electronically prime the syringe pump system before starting an infusion or after replacing a near-empty syringe with a replacement syringe.

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 priming and bolusing the pressure limits are set to the maximum level.

Select syringe/catheter suitable for use with the intended medical application.

⚠️ Caution: Position the infusion line free of kinks.

Recommended change of disposable each 24 h (or as per national hygiene regulations).

Installation in medically used rooms must comply with the appropriate regulations (e.g. VDE 0100, VDE 0107 or IEC-publications). Observe national specifications and deviations.

⚠️ Caution: Operate the pump at least 25 cm from flammable anaesthetics to prevent explosion

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

If staff call is used we recommend checking the equipment once after connecting the pump.

Protect the device and the power supply against moisture.

Do not carry the pump device by it’s drive mechanism during transportation.

If the pump device falls or is exposed to force it needs to be examined by the service department.

The displayed data must always be checked by the user prior to making further medical decisions.

During mobile use (homecare, patient transport inside and outside the hospital): Make sure the device is securely fixed and positioned. Positioning changes and severe shock can lead to minor changes in the delivery accuracy and/or unintentional bolus administration.
- A supplemental patient monitoring must be carried out if life-saving medication is performed.
- Avoid applying external force on the drive mechanism during administration.
- In case high potent drugs are given be sure to have a second infusion pump for that drug at hand. The therapy documentation should be suitable to continue the therapy at the second infusion pump.
- Changing the height during a running infusion may lead to flow rate variations.
- Independant of the soft limits the selected values have to be the medically correct ones for the given patient.
- In case values relevant for the dose rate calculation (e.g. body weight) are changing always the flow rate will be updated and the dose rate will be fix.
- Consider startup characteristics before using low infusion rates (0.1ml/h) with critical drugs.

Enteral Nutrition

The Perfusor® Space may be used for enteral nutrition. Do not use enteral fluids for intravenous infusion as this may harm your patient. For this reason only use disposables dedicated and labeled for enteral nutrition.

Other components

- Use only pressure proof and compatible disposable items (min. 2 bar/1500 mm Hg) to avoid influencing performance data - which would result in impairing patient safety.
- 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.
- Refer to the according manufacturer's information for possible incompatibilities of equipment with respect to drugs.
- Use only compatible combinations of equipment, accessories, working parts and disposables with luer lock connectors.
- The use of incompatible disposables may influence the technical specifications of the device.
- Connected electrical equipment must comply with the relevant IEC/EN-specifications (e.g. IEC/EN 60950 for data-processing equipment). The user/operator is responsible for the system configuration if additional equipment is connected. The international standard IEC/EN 60601-1-1 has to be taken into account.
Safety Standards


- The EMC-limits (electro-magnetic compatibility) according to IEC 60601-1-2:2007 and IEC 60601-2-24:2012 are maintained. If the equipment is operated in the vicinity of other equipment which may cause high levels of interference (e.g. HF surgical equipment, nuclear spin tomography units, mobile telephones etc.) may be disturbed. Maintain the protective distances recommended by the manufacturers of these devices.

- The Perfusor® Space fulfils the applicable requirements of EN 13718 to be used in the air, on the water and in difficult terrain. During transport the Perfusor® Space needs to be fixed on a suitable restraint system by means of SpaceStation or Pole Clamp SP. When stored under temperature conditions beyond the defined operating conditions the Perfusor® Space needs to remain under room temperature at least one hour before usage.

Safety Instructions for using PCA

- In case the demand button is used with SpaceStation the PCA pump has to be placed in the lowest slot of the lowest SpaceStation.

- Access to the pump settings can be prohibited by DataLock 3. The code for DataLock level 3 should differ from the one for levels 1 and 2 in case the pump is only allowed to be used by pain management professionals.

- For additional safety the removal of the syringe can be prevented by the use of the Syringe Anti Removal Cap (see accessories) and the locking of the syringe holder. The Syringe Anti Removal Cap is usable for the following syringes: B. Braun Original Perfusor Syringe 50 ml, B. Braun Omnifix 50 ml, BD Plastipak 50/60 ml and Tyco Monoject 50 ml. The locking of the syringe holder is under the pump and is locked by a clockwise turn of 90°. Make sure the syringe holder is safely locked. Opening of the syringe holder may not be possible after locking.

- In case opioids are administered and the Syringe Anti Removal Cap is not in use and the syringe holder is not locked the therapy only should be performed under surveillance of medical staff. This especially is necessary in case non-authorised access to the drug can be anticipated.

- When ending PCA and starting it again the therapy data are set to default values.

- Using the demand button also the patient is a permitted user. With the demand button only a PCA-bolus can be requested. This is limited to pre-defined doses by drug list and pump settings.

- Consider startup characteristics before using low infusion rates (0.1ml/h) with critical drugs.
Safety instructions for using TCI

- TCI should only be performed by experienced anaesthetists being familiar with the principles of TCI and properly trained in using the present device.

- The use of TCI with B. Braun Space does not limit the responsibility of the anaesthetist for administration of drugs. They need to be fully aware of the available literature for any parameter set used in association with a drug and need to refer to the prescribed information for rate and dosing limits.

- Pharmacokinetic and pharmacodynamic interactions among anaesthetic drugs are known, but are not taken into account into the calculation of the plasma and effect site concentrations. They have to be taken into account by the user.

- In particular, the user must be aware that starting the TCI will result in the automatic infusion of a pre-calculated bolus dose followed by an infusion to achieve the selected target concentration.

- It is essential that the user verifies that the patient characteristics and the selected target concentration as well as the resulting dosages conform to the prescribing information of the relevant country.

- B. Braun has verified the accuracy of the mathematical model implementation, the usability as well as pump delivery accuracy.

- While using TCI an appropriate patient monitoring is mandatory.

- Take care of using the right dilution/concentration of the drug and make sure the right dilution is selected at the pump.

- Never administer TCI drugs by a second infusion as long as you use TCI.

- It is possible to completely switch off the TCI mode to avoid the use of TCI accidentally.
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.
At the top of the screen the last therapy is indicated. Yes/No question can be answered by pressing \( \uparrow \) for yes or \( \downarrow \) for no.

Parameters which can be changed (e.g. rate in ml/h) are opened with \( \downarrow \) or \( \uparrow \). When editing parameters, switch digits/levels using \( \leftarrow \), \( \rightarrow \). White background indicates current digit/level. Use \( \uparrow \) or \( \downarrow \) to change current setting. Help text on the bottom/top of the screen indicates options how to proceed (e.g. confirm rate with \( \uparrow \), start infusion with \( \uparrow \) or clear rate by pressing \( \downarrow \)).

All display screen shots are examples and may be different when related to an individual patient and individualized therapy.

Typical display during infusion:

- **Battery status**
- **Connection to SpaceCom in SpaceStation**
- **Set pressure limit and current pressure* (See also OccluGuard)**
- **Therapy profile**
- **Active VTBI- or time preselection**
- **Scrolling arrows indicate pump is infusing**
- **Set rate can be opened with \( \downarrow \)**
- **Unit of drug application**
- **Total volume infused. Alternatively the intermediate volume can be displayed.**
- **Remaining time**
- **Remaining VTBI**

### Display

**Meaning**

At the top of the screen the last therapy is indicated. Yes/No question can be answered by pressing \( \uparrow \) for yes or \( \downarrow \) for no.

Parameters which can be changed (e.g. rate in ml/h) are opened with \( \downarrow \) or \( \uparrow \). When editing parameters, switch digits/levels using \( \leftarrow \), \( \rightarrow \). White background indicates current digit/level. Use \( \uparrow \) or \( \downarrow \) to change current setting. Help text on the bottom/top of the screen indicates options how to proceed (e.g. confirm rate with \( \uparrow \), start infusion with \( \uparrow \) or clear rate by pressing \( \downarrow \)).
Set pressure level with \( \text{l} \) or \( \text{r} \) and confirm by pressing \( \text{k} \). Cancel to edit pressure by using \( \text{c} \).

Pre-alarm s are indicated by the message on the display (e.g. “Syringe nearly empty”), an audible tone and the yellow LED is constantly on. To confirm a pre-alarm press \( \text{ok} \).

In case of an operating alarm (e.g. “Syringe empty”) the infusion stops, an audible tone sounds and the red LED is flashing. Confirm alarm by using \( \text{ok} \). Confirming does not activate an acoustic feedback.

Press and hold \( \text{c} \) for 3 sec to turn pump off. A white bar stretches from left to right and counts down the 3 sec. As long there is a syringe inserted the pump will not turn off but will use standby.
Syringe selection
Prime?
Use last therapy?
Use dose rate calculation?
Use drug library?

Dose
Concentration
Weight
Rate
VTBI
Time

Syringe selection
Prime?
Use last therapy?
Use dose rate calculation?
Use drug library?

Dose Rate Calculation
Drug Library
Change-over from continuous mode to PCA

Options Menu
Occlusion Pressure
OccluGuard
Pressure Leap/Drop
Data Lock
Bolus Rate
KVO-Mode
Contrast
Display Light
Keypad Light
Alarm Volume
Date
Time
Macro Mode
In Dose Mode: Display of flow rate in large scale
Language

Status Menu
Intermediate volume
Intermediate amount
Intermediate time
Total volume
Total amount
Total time
Pressure
OccluGuard Status
Syringe
Battery capacity
Wlan
Version
Drug info
1.1 Start of Infusion

- Ensure correct installation of the pump device. If the pump is connected to mains, the display states information such as the battery status, the mains connection symbol and the last therapy.

- Press \( \text{O} \) to switch on unit. Note the automatic selfcheck: “Selftest active” and the software version are displayed, two audible tones sound and all three LEDs (yellow, green/red and blue) flash once. Information on power supply (battery or mains connection), the set pressure level and the syringe (if syringe already inserted) are displayed. Hence the drive moves backwards.

- Press \( \text{C} \) to start with direct entry of therapy parameters or open pump cover and syringe holder to start with syringe insertion.

- Insert syringe with wings of the syringe upright in the slot to the right of the housing. Close syringe holder and pump door. Piston brake moves forward.

  **Caution:** Never leave the pump unattended during syringe loading.

- Confirm syringe type with \( \text{K} \). Type of syringe indicated must coincide with syringe inserted.

- Drive will advance and grip pressure plate of syringe.

  **Caution:** Keep your hands away from advancing device.

  **Note:** Make sure that the piston brake moves back into the syringe holder.

- If the prime function is activated, press \( \text{U} \) to prime infusion set at 1200 ml/h (pressing key once = 1 ml). Interrupt prime function with \( \text{K} \). Repeat procedure until infusion line is fully primed. Then press \( \text{D} \) to proceed.

- Connect with patient.

- Respectively answer questions in Start Up Menu with \( \text{U} \) and \( \text{D} \), until the rate is displayed in the Main Menu.

**Enter infusion rate:**

- Press \( \text{L} \) and set rate using \( \text{Q} \).
Chapter 1

- Press \( \text{SF} \) to commence infusion. Running arrows on display and green LED above display indicate pump is infusing.

**Note:** Stop the infusion at any time by pressing \( \text{SF} \). The pump can be turned off at any time by pressing \( \text{O} \) for 3 sec (Exception: Data lock level 2) and as long a disposable is inserted.

1.2 Entry With Different Combinations of Rate, VTBI (= Volume To Be Infused) and Time

The Perfusor® Space offers the possibility to enter a volume- and time limit in addition to an infusion rate. When two of these parameters are entered, the third is calculated by the pump. If a volume and/or time is preselected, an arrow symbol is placed in front of one of these parameters in the Main Menu. It is called the "target". During the infusion of the pump, this target symbol is displayed next to the moving arrows in the run display (this symbol is not visible in case TCI is used). This indicates that the pump has been programmed, either with a volume- or time limit. The assignment of the target symbol, apparent in the Main Menu, shows the established parameter for the application (VTBI or time). When the rate is changed, the so-called target parameter is principally not adjusted to the new rate but to the parameter which does not have the target symbol in front. After the infusion has started, the remaining VTBI and time are displayed in the status menu and the run display (values are counting down).

1.) Enter VTBI and time: The infusion rate will be calculated and displayed on the bottom of the display.
   **Target: Volume**
   - Select VTBI with \( \text{B} \) and open with \( \text{L} \).
   - Enter VTBI with \( \text{Q} \) and confirm with \( \text{OK} \).
   - Select time with \( \text{B} \) and open with \( \text{L} \).
   - Enter time with \( \text{Q} \) and confirm with \( \text{OK} \).
   Check calculated rate on plausibility.
   Proceed in the same way to calculate 2.) and 3.).

2.) Infusion with volume limit
   Enter rate and VTBI: The infusion time will be calculated and displayed at the bottom of the display.
   **Target: VTBI**

3.) Infusion with time limit
   Enter rate and time: The infusion volume will be calculated and displayed at the bottom of the display.
   **Target: Time**

**Changing already entered values of VTBI and time** (rate, VTBI and time already exist at the point of change):
1.3 Bolus Application

After pressing the button \( \text{nb} \) the bolus unit can be selected by using \( \downarrow \).

Note: The selected unit will not be stored. It is possible to administer a bolus in ml.

There are three ways to administer a bolus:

1.) Manual Bolus: Press \( \text{nb} \). Then press \( \text{ok} \) and hold button. Fluid is administered as long as button is held down. The infused bolus volume is displayed.
   The max. bolus volume is limited to 10 sec. Reaching this limit is indicated by an acoustic signal.

2.) Bolus with volume preselection: Press \( \text{nb} \). Then press \( \leftarrow \) and set bolus dose limit by using \( \text{q} \). Press \( \text{ok} \) to confirm and start bolus. Depending on the service tool settings an acoustic signal will sound after finishing the bolus volume.

3.) Bolus with rate calculation: Press \( \text{nb} \). Then press \( \leftarrow \) and set bolus dose by using \( \text{q} \). Press \( \text{ok} \) to confirm bolus dose. Set time with \( \text{q} \) in which a bolus is to be delivered. Calculated bolus rate is shown on top of the display. Press \( \text{ok} \) to confirm and start bolus.

You can use the service program to enter a default and a maximum bolus rate. Once a new therapy is started the device always returns to the default rate - even if the bolus rate was manually changed beforehand.

Note: If the bolus limit is not entered after pressing \( \text{nb} \), the pump switches back into the run display automatically.

Note: The infused volume during bolus with volume preselection counts up.

In order to purge the line at any time while the pump is stopped press \( \text{u} \). Answer the following question by pressing \( \text{u} \) in order to start the purge process. Cancel by pressing \( \text{u} \) or any other key.

Caution: Take care not to overdose! Given a bolus rate of 1200 ml/h, 1 ml will be administered in just 3 sec. To cancel bolus infusion at any time press \( \text{ok} \).

At low bolus volumes, under dosages due to the start up characteristic of the pump and the tolerances in the infusion system cannot be excluded. Disconnect patient while purging.
1.4 Syringe Change and New Therapy Start

**Note:** To avoid incorrect dosing, always disconnect the pump from the patient when changing the syringe. Never leave the pump device unattended during syringe change. Before inserting a new syringe check if the axial fixation is properly working.

- Press to stop the infusion. The green LED will disappear. Disconnect the pump from the patient.
- Either the drive mechanism moves backwards into starting position when opening syringe holder or the question "Perform change?" must be first answered before the drive moves backwards.
- Open pump door, remove syringe and insert new syringe.
  
  **Note:** In case the plunger head of the syringe is not released anymore by the claws when performing a syringe change, the emergency release button needs to be pressed to release the claws of the drive head. The emergency release button is placed on the outside of the drive head. It can be released with a pointed pen. Then manually open the claws and take out the syringe.

- Close the syringe holder (**Note:** Piston brake must move forward!) and the pump door and confirm the inserted syringe type with . Drive advances and grips pressure plate of syringe.
  
  **Note:** Do not block advancing drive unit with any objects. Piston brake must move backwards into the syringe holder.

- Prime pump if necessary with then press to continue.
- Connect the patient to the pump and check set parameters using .
- Press to start infusion.

**To start a new therapy after a syringe change** (see also Chapter Start Up Graphs and Trumpet Curves):

- Press when pump is in the Main Menu.
- Press and continue to set new therapy parameters with .
- Press to start infusion.

**Note:** A new therapy can be started at any time during a stopped infusion. Press (repeatedly) when the pump is in the Main-, Status- or Options Menu and proceed to follow instructions as described.

1.5 End of Infusion

- Press to stop the infusion. The green LED disappears. Disconnect the pump from the patient.
- Open the syringe holder. Answer the question whether a syringe change should be performed with . The drive moves backwards into the starting position.
Chapter 1

- Open pump cover. Remove the syringe, move the syringe holder into an upright position and close the front door.

- Press \( \odot \) for 3 sec. to switch the pump off. The drive moves into parking position.

  **Note:** The settings will be permanently saved by the switched off device.

  **Note:** Pump cannot be powered off with syringe inserted.

1.6 Standby Mode

In the case of extended interruption, the user has the option to maintain the set values.

- Press \( \odot \) to stop the infusion. Then press \( \odot \) for less than 3 sec.

- Confirm that the pump is supposed to switch into standby by pressing \( \uparrow \).

- The pump is now in Standby.

While the pump is in the standby mode, its display shows the drug and the remaining time for this mode. Change of remaining time by pressing \( \leftarrow \). Exit standby by pressing \( \odot \).

As long as a disposable is inserted in the pump will use standby also in case \( \odot \) is pressed for at least or more than 3 sec.
ADVANCED OPERATIONS

2.1 Status Request of Pump when Infusion is Running

Press \( \circ \) to switch between run display and Main Menu while the device is infusing. Navigate through the menu using \( \triangledown \) to check parameters. In order to check the menu parameters in the Status-/Options Menu, select "Status" respectively "Options" in the Main Menu, open menu with \( \downarrow \) and scroll through menu with \( \uparrow \).

2.2 Rate, VTBI and Time Change Without Infusion Interruption and Reset of Status Menu Data

- Press \( \circ \) when the pump is in the run display in order to switch to the Main Menu. Select rate/VTBI/time with \( \triangledown \) and press \( \downarrow \) in order to open the parameter.
- Enter new value with \( \circ \) and confirm with \( \downarrow \).

Reset Status Menu Data:
The parameters intermediate volume and -time can be reset when the pump is infusing or when the pump is stopped.

- Select "Status" in Main Menu with \( \triangledown \) and press \( \downarrow \).
- Highlight intermediate volume (in ml) or intermediate time (in h:min) with \( \triangledown \) and open parameter with \( \downarrow \).
- Reset values by pressing \( \uparrow \).

Both parameter total volume and -time, are displayed in the pump as "Total" with the according unit and can be reset by starting a new therapy. A second way to reset the parameters when the pump is in the Main Menu: Press \( \circ \), answer question if the last therapy is to be used with \( \uparrow \) and reset the values with \( \downarrow \).

The type of the inserted syringe is displayed in menu item "Syringe" and cannot be changed once it has been confirmed at the beginning of the infusion. The drug info states the drug name, the name of the drug list and its date of origin. The current battery capacity in hours and minutes is displayed in the menu item "Battery Cap." and the current software version in menu item "Version." In-line pressure can also be read in the Status menu in mmHg or Bar depending on the service settings.
3.1 Dosing Units and Dose Rate Calculation (Overview)

The following list shows the units used in the pump:

- **Gram family:** ng, mcg, mg, g
- **Unit family:** mIU, IU, kIU, MIU
- **Equivalents family:** mEq
- **Mole family:** mmol
- **Kilocalorie family:** kcal
- **Milliliter family:** ml, ml/kg

In addition to these dosing units the user can choose:
- Feeding: kcal, mEq, mmol
- Surface related amount units: m²

The pump is calculating the body surface area with the "Dubois" formula (DuBois D, DuBois EF. A formula. Arch Intern Med 1916; 17: 863):

$$\text{BSA (m²)} = 0.007184 \times \text{weight (kg)}^{0.425} \times \text{height (cm)}^{0.725}.$$ 

Check plausibility of calculated body surface area value and resulting delivery rate before starting the infusion, also, if body surface area related dose rate is set by Barcode. The dose rate calculation enables a calculation of the rate in ml/h based on the entered dose parameters.

**Setting parameters:**

1. Concentration as the amount of the active ingredient per volume.
   - Amount of the active ingredient
   - Volume in ml
2. Where necessary: Patient weight or Patient height
   **Note:**
   - Patient weight can be entered in kg, lbs or grams.
   - Patient height is entered in m (is used to calculate BSA)
3. Dose prescription:
   - time related as the amount of the active ingredient per min, h or 24h.
   - time and patient weight related as the amount of the active ingredient per kg per min, h or 24h or BSA.
3.2 Dose Rate Calculation (Operation)

- Select dose rate calculation with ◀.
- Select the unit of the active ingredient with ▼ and confirm it with ◄.
- Enter the concentration by entering the amount of the active ingredient and the volume. In order to do so set the values with ◄ and confirm with OK.
- If the patient weight shall not be entered press ◄.
  Press ▼ to choose “weight” or “surface” and confirm with OK.
- Set the patient weight with ▼ and confirm with OK.
- Select the dose prescription with ◄ and confirm it with ◄.
- Set the dose with ◄ and confirm with OK. The rate will automatically be calculated and displayed at the bottom of the display.
- Check the calculated rate and if necessary the adapted parameters with ◄ on plausibility before starting the infusion with ◄.

Dose can later be changed in the Main Menu in the same way as the rate, VTBI and time (compare 2.2). During TCI mode after a syringe change, the concentration can only be changed in Main Menu. The effect of dose modifications on other parameters is shown at the bottom of the display. Additionally the total and intermediate amount of the infused drug can be taken from the Status Menu. These can be checked and resetted in the same way as the other total and intermediate values.

A deactivation of the dose rate calculation is only possible when the pump is stopped. Press ◄ from Main Menu and then press ◄.

Caution: A change of the patient weight or height will alter the flow rate.

3.3 Drug Library

Up to 1200 drug names including therapy data, information and up to 10 concentrations per drug can be stored in 30 categories. These drugs can be subdivided in 50 care units and 16 Patient Profiles. The loading process into the pump can be performed via a separate PC program (Space Upload Manager & HiBaSeD).

Note: The drug library can be started over the Start Up and Special Functions Menu. The user has to make sure prior to the therapy start that the drug library in the pump complies with the patient target group. The name of the care unit and creation date (see headline) should be checked in the pump.

There are different ways of assigning a drug to an infusion. This can be done while the infusion is running or when the pump is stopped.
On the one hand, a drug name including the according therapy data can be taken from the drug library. On the other hand, if a rate, VTBI and/or time were already defined in the Main Menu, the drug name and the adjusted values of the data set will be loaded. If a dose rate calculation has already been started a belated assignment of the drug name nevertheless is possible.

**Loading a drug (including the according parameters) from the Main Menu:**

- Go to Special Functions Menu and press \(\text{①}\).
- Open the drug library by pressing \(\text{①}\).
- Navigate through the list with \(\text{⑧}\) and select the care unit with \(\text{④}\). If you have already set the care unit once on your pump this step will be skipped for the next time.
- Change the care unit by navigating through the list until "Change care unit" will be displayed. Press \(\text{OK}\) to change the care unit.
- Navigate through the list with \(\text{⑧}\) and select the patient profile with \(\text{④}\). If no profile is set, this step will be skipped.
- Navigate through the list with \(\text{⑧}\) and select in alphabetical order (all drugs) or within a category with \(\text{④}\).
- If different therapies are related to a drug, choose therapy type with \(\text{⑧}\) and confirm with \(\text{⑥}\).
- Confirm the displayed drug information with \(\text{⑦}\).

![Drug Selection Screenshot]

- Decide if the safety limits for the drug are to be applied \(\text{⑦}\) or if only the drug name should be used \(\text{⑦}\).
- Check if the drug short name in the Run Menu is the same as the selected drug. Check the parameter in the Main Menu with \(\text{⑧}\) and start infusion with \(\text{SF}\).

**Note:** If a drug name has been assigned without safety limits, the following hint is provided in the RUN screen:

![Drug Limit Screenshot]

**Note:** Care unit and Patient Profile can not be changed within a therapy.

**Initial Bolus:**

Initial Bolus has to be configured in the Drug List Manager.

- Use the drug library according to the instructions for use.
Select the desired drug with \( \text{\textbullet} \) and press \( \text{\textbullet} \).
Before the initial bolus begins, the bolus menu is displayed to allow editing the bolus with \( \text{\textbullet} \).

Check the parameter and start infusion with \( \text{\textbullet} \).

**Hard Limits:**
If the set rate/dose/bolus volume and bolus rate exceed the values stored in the drug library (hard limits), the drug will be rejected, a hint will be displayed and the pump will fall back into the drug selection. If this occurs while the pump is infusing the pump will continue to administrate.

**Soft Limits:**
For the same parameters so called soft limits can be preset via the Drug List Editor. These can be exceeded without any constraint. The following symbols that describe the status with regard to the soft limits are being displayed:

The infusion is within the range of the minimum and maximum soft limits

The infusion is within the range of the maximum soft limit

The infusion is within the range of the minimum soft limit

Violation of the upper soft limit

Violation of the lower soft limit

No soft limit is defined

Only a drug name is available

(It is possible to select a drug name only from the drug library)

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

**Note:** An adequate monitoring when infusing highly potent drugs is recommended.

**Note:** In case a drug from the drug library is selected and the pump is running under dose rate calculation the initial values will be overwritten by the drug library values if selected.

**Remote Drug Library update from Upload Manager (Space Online Suite)**
The file icon blinks every 2 s. An update is available.
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The Drug Library Upload starts as soon as the pump is in Passive mode.

Note: You can cancel the upload by pressing \( \text{cancel} \).

Please contact your local sales representative in case you like to use Remote Drug Library update.

3.4 Patient Controlled Analgesia (PCA)

For PCA a drug list with at least one drug activating the profile PCA is necessary. By this the conditions for an effective and safe therapy are defined.

Switch on pump with \( \text{on} \) and wait until self-check is finished. Depending on the settings the choice of a drug is offered directly or the pump is in "Main Menu".

Select "Special Functions" with \( \text{select} \) from "Main Menu" and confirm with \( \text{confirm} \).

Select drug list, category and desired drug by using \( \text{select} \). After the selection the pump offers additional drug related information which are confirmed by \( \text{confirm} \).

Select profile PCA by using \( \text{select} \) and confirm with \( \text{confirm} \). The therapy settings stored in the drug list are displayed *.

The therapy can be started now with \( \text{start} \) in case all values are defined.

Depending on the pre-defined settings the therapy is started with an initial bolus and a basal rate or not.

Before leaving the patient the pump should be put into DataLock level 3 with \( \text{lock} \) in Menu "Options". This is necessary especially in case non-authorised access to the settings can be anticipated.

The code is entered with \( \text{enter} \) and confirmed with \( \text{ok} \). The pump display now may look like this.

*Bolus volume is the volume of a single bolus the patient may demand. Max. Limit is the amount of drug or volume a patient may demand within a certain time in total. Lockout is the time in between two bolus.
In this state the patient is allowed to demand boli. Depending on the status of the therapy these are either administered or denied. Changing the syringe is also possible by using the code for level 1 or level 2. Altering the settings for PCA or other therapies however is only possible with the code for level 3.

The status of the therapy can be checked in the menu „Status“. Enter the „Main Menu“ with and select the “Status“ with .

An acoustic confirmation of demanded boli can be activated and modulated by in Data Lock 3.

Is a demand button connected, the therapy symbol looks like this: .

In case there is no demand button connected the therapy symbol looks like this: .

The demand button is connected to the interface P2 at the rear side of the pump.

Hint: It is possible to start a therapy in continuous mode and switch over to PCA later on (in case the drug is dedicated for use with continuous and PCA application).

SpacePCA–Chart

If is pressed on the RUN screen, the SpacePCA–Chart is displayed:

The bar represents a time axis, with the points above the axis representing the number of boli administered and the points below the axis representing the number of boli refused.

The chart has a 15 minute resolution and shows max of 5 points per 15 minutes. Should more then 5 boli be given or refused in this time, the last point will be turn bold.

Changes to the PCA parameters are displayed as arrowheads at the bottom of the chart.

The A/D–ratio indicates the percentage of administered and demanded boli thus giving an idea about the effectivity of the therapy.
3.5 Target Controlled Infusion (TCI)

Introduction

In TCI the user is defining a desired concentration of drug in the human body (target) rather than an infusion rate. The rates necessary to reach and maintain that said concentration are calculated by the pump using an algorithm based on a three-compartment pharmacokinetic model.

A pharmacokinetic model (PK model) is a mathematic model to predict the concentration of a drug in the human body (e.g. plasma level) after a bolus or a continuous infusion of different duration. A PK model is developed by measurement of plasma level values of a population of patients or volunteers and the respective statistical analysis. A PK model mostly is a 2- or 3-compartment model indicating the volumes of the compartments, indicating rates for the exchange amongst the compartments and indicating rates for elimination / metabolism of the drug.

A PK model can be parameterized to use it for different drugs as long as it is suitable for that said drug. The pharmacokinetic model and its parameters are schematically depicted by the following illustration:

B. Braun Space is offering two modes for TCI:

- TCI by targeting the plasma concentration
  In this mode the user selects the desired concentration of a drug in the blood plasma and the PK model is used to calculate the infusion rates required to achieve that concentration as quick as possible (unless there is no restriction defined by the user).
TCI by targeting the effect-site concentration.
In this mode the user selects the desired concentration of a drug at the site of action and the PK model is used to calculate the infusion rates required to achieve that concentration as quick as possible (unless there is no restriction defined by the user). A certain overshoot of the concentration in the plasma is resulting from this mode.

For effect-site targeting there is a link between pharmacokinetics and pharmacodynamics necessary. As the effect-site compartment is considered to have no volume and the rate constant $k_{1e}$ can be ignored the rate constant $k_{e0}$ is the parameter necessary to perform effect-site TCI. A pharmacokinetic model modified in such way is schematically depicted by the following illustration:

TCI with B. Braun Space is possible with the following drugs: Propofol, Remifentanil and Sufentanil.

For Propofol the user can choose between two parameter sets. The parameter sets used for these drugs are (Not all parameter sets allow effect-site targeting):
<table>
<thead>
<tr>
<th>Drug / Parameter</th>
<th>Propofol Marsh</th>
<th>Propofol Schnider</th>
<th>Remifentanil</th>
<th>Sufentanil</th>
</tr>
</thead>
<tbody>
<tr>
<td>$V_c = V_1$ [ml]</td>
<td>-</td>
<td>-</td>
<td></td>
<td>14,3</td>
</tr>
<tr>
<td>$V_1$ [litre]</td>
<td>0.228 * Weight</td>
<td>4.27</td>
<td>5.1 – 0.0201 * (Age - 40) + 0.072 * (LBM - 55)</td>
<td></td>
</tr>
<tr>
<td>$k_{10}$ [min⁻¹]</td>
<td>0.119</td>
<td>0.443 + 0.0107 *</td>
<td>0.0419</td>
<td>0.0645</td>
</tr>
<tr>
<td></td>
<td>(Weight - 77)</td>
<td>(LBM - 59) + 0.0062 * (Height - 177)</td>
<td></td>
<td></td>
</tr>
<tr>
<td>$k_{12}$ [min⁻¹]</td>
<td>0.112</td>
<td>0.302 - 0.0056 *</td>
<td>0.302 – 0.006 * (Age - 53) / (5.1 - 0.0201 * (Age - 40) + 0.072 * (LBM - 55))</td>
<td>0.1086</td>
</tr>
<tr>
<td>$k_{13}$ [min⁻¹]</td>
<td>0.0419</td>
<td>0.196</td>
<td>[0.076 - 0.00113 * (Age - 40)] / 0.0229</td>
<td></td>
</tr>
<tr>
<td>$k_{21}$ [min⁻¹]</td>
<td>0.055</td>
<td>0.24 * (Age - 53)</td>
<td>0.24 * (Age - 53) / (18.9 - 0.391 * (Age - 53)) / [9.82 - 0.0811 * (Age - 40) + 0.108 * (LBM - 55)]</td>
<td>0.0245</td>
</tr>
<tr>
<td>$k_{31}$ [min⁻¹]</td>
<td>0.0033</td>
<td>0.0035</td>
<td>0.01402 - 0.0002085 * (Age - 40) / 0.0013</td>
<td></td>
</tr>
<tr>
<td>$k_{e0}$ [min⁻¹]</td>
<td>0.26</td>
<td>0.456</td>
<td>0.595 – 0.007 * (Age - 40) / 0.112</td>
<td></td>
</tr>
</tbody>
</table>

**Reference**

- Schnider et al., Anesthesiology, Vol. 88, 1998, 1170-1182
- Schnider et al., Anesthesiology, Vol. 90, 1999, 1502-1516
- Minto et al., Anesthesiology, Vol. 86, 1997, 10-33

**Effect-site targeting**

- No
- Yes
- Yes
- Yes
Drug List

The pre-installed drug list offers the following values:

<table>
<thead>
<tr>
<th>Available Concentrations</th>
<th>Propofol</th>
<th>Remifentanil</th>
<th>Sufentanil</th>
</tr>
</thead>
<tbody>
<tr>
<td>5 mg/ml</td>
<td>20 µg/ml</td>
<td>5 µg/ml</td>
<td></td>
</tr>
<tr>
<td>10 mg/ml</td>
<td>50 µg/ml</td>
<td>50 µg/ml</td>
<td></td>
</tr>
<tr>
<td>20 mg/ml</td>
<td></td>
<td></td>
<td></td>
</tr>
</tbody>
</table>

| Default Max. Rate        | 1.200 ml/h | 1.200 ml/h | 1.200 ml/h |
| Hard Limit Rate          | Max of pump| Max of pump| Max of pump|
| Plasma Limit Default     | 400 %      | 400 %       | 400 %      |
| Plasma Limit Hard Low    | 100 %      | 100 %       | 100 %      |
| Plasma Limit Soft Max    | 450 %      | 450 %       | 450 %      |
| Default Target           | 0.0 µg/ml  | 0.0 ng/ml   | 0.0 ng/ml  |
| Target Soft Max          | 8.0 µg/ml  | 8.0 ng/ml   | 1.0 ng/ml  |
| Target Hard Max          | 15.0 µg/ml | 20.0 ng/ml  | 2.0 ng/ml  |
| Decrement Concentration  | 1.0 µg/ml  | 1.0 ng/ml   | 0.2 ng/ml  |
| Default Parameter Set    | Marsh      | Minto       | Gepts      |

**Important note:** Before installing an additional drug list please contact your local B. Braun representative!

Setting up the pump

For TCI a drug list with at least one drug activating the profile TCI is necessary. The drug list in this version is pre-defined. By this the conditions for an effective and safe therapy are defined.

Switch on pump with \( \text{ } \) and wait until self-check is finished. Insert disposable and use the drug library according to Instructions for Use.

Selecting a drug

Select drug list, category (the TCI drugs need to be selected from the category “TCI”) and desired drug by using \( \text{ } \). In this example: Propofol.

As a next step select the correct dilution (concentration) of the drug to be administered as well as the parameter set (model) and the Mode (Effect-Site Targeting or Plasma Targeting).
These steps are only necessary in case there are different options for that drug.

Input of patient data

Depending on the parameter set one or more of the following data are necessary:
- Weight
- Height
- Gender
- Age

The editor window appears with the initial setting “0” to make sure editing a value takes place (exemption: initial setting for gender is “male”).

Important notes:
- Be sure to enter the data corresponding to the respective patient.
- Once the TCI is started patient data can not be altered!

Editing a target and starting TCI

The editor window for setting the target comes up with the default value from the drug list.

Confirm target with \( \text{ok} \). TCI can be started now with \( \text{sf} \).

After TCI is started the screen looks the following:

The parameter in the lower left corner can be scrolled. The remaining syringe volume depends on the calculated rate. Because of the changing TCI algorithm, rate varies and therefore technically the time for the remaining syringe volume cannot be precisely predicted.
In the top line there is an icon indicating the parameter set and the mode (Mode Indicator) with following meaning:

- “TCI Ma P”: TCI Marsh plasma targeting
- “TCI Sc P”: TCI Schnider plasma targeting
- “TCI Sc E”: TCI Schnider effect-site targeting
- “TCI Mi P”: TCI Minto plasma targeting
- “TCI Mi E”: TCI Minto effect-site targeting
- “TCI Ge P”: TCI Gepts plasma targeting
- “TCI Ge E”: TCI Gepts effect targeting

In the bottom line the status parameters like flow rate, Cp/Ce, infused volume etc. can be displayed. The desired parameter can be selected by using . It is recommended to select Cp/Ce.

In case it is necessary to change the target press to edit the value.

Useful information while pump is running

By pressing additional information can be requested.

Pressing a second time is offering a graphical overview.

The line describes the course of Cp over the time and the area describes the course of Ce over the time. The time window is 20 min (15 min past, 5 min future).

Additional information is left with .

Finishing TCI

There are two possibilities to finish the TCI Therapy (reversion of anaesthesia or sedation):

- Set Target= 0
- Stop pump

It is recommended to simply stop the pump by pressing .

Pressing the pump offers additional information – in this case the information is modified the following way:
After the therapy is ended there are two possibilities:

a) The pump may be used for TCI with the same drug again but with a new patient. In this case, cancel old therapy and use new disposables.

b) The pump may go with the patient but in continuous mode (without TCI).

In both cases the “old” TCI needs to be ended by and selecting “Yes” in this screen by pressing .

In case a) press in the menu – in case b) press .

3.6 Barcoding

The barcoding functionality is included but initially not activated.

Please contact your local sales representative in case you like to use barcoding.

3.7 Ramp and Taper Mode

The Ramp and Taper Mode is designed to deliver infusions with gradual ramp up and taper down rates. The pump automatically calculates the rate increase and decrease required to match the total volume, time and ramp up/ramp down time parameters. It consists of 3 phases.

- Ramp phase: the pump rate is linearly increased until it reaches a predefined rate (plateau rate) in a predefined time (Up-Time)
- Continuous phase: the plateau rate is used as a continuous infusion
- Taper phase: the pump rate is decreased linearly after the continuous phase until the KVO rate is reached or pump is stopped in a predefined time (Down-Time)
Example:

Ramp and Taper should only be performed by an experienced user that is familiar with the principles of the Ramp and Taper function and properly trained in using the present device.

**Note:** The active Ramp and Taper function is always symbolised with an characteristical symbol in the Display ( \( \text{[ ]} / \text{ [ ]} / \text{ [ ]} \)).

**Note:** Bolus function is disabled for Ramp and Taper Mode.

Set Profile Parameters: The therapy can be started directly via the drug library or via the Main Menu/Special functions.

**Starting Ramp and Taper via Drug Library:**

**Note:** Ramp and Taper settings have been configured in the Drug List Manager before and have been uploaded into the pump.

- Switch on pump with \( \text{[ ]} \) and wait until self-check is finished.
- Insert disposable and use the drug library according to the Instructions for Use.
- Select the desired drug with \( \text{[ ]} \) and press \( \text{[ ]} \).

The pump now lists the possible therapy profiles.

- Select "Ramp and Taper Mode" with \( \text{[ ]} \) and press \( \text{[ ]} \).
  The therapy settings for "Ramp and Taper Mode" are shown on the display.
- To change the values, press \( \text{[ ]} \) to change and \( \text{[ ]} \) to confirm.

The pump can be started now by pressing \( \text{[ ]} \).

**Starting Ramp and Taper via Special Function Menu:**

- Switch on pump with \( \text{[ ]} \) and wait until self-check is finished.
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- Insert disposable.
- Go to Special Functions Menu and select Ramp and Taper.
- Press \( \text{↓} \) to enter parameters and \( \text{OK} \) to confirm.
- After entering all desired parameters the pump can be started by pressing \( \text{SF} \).

The status of the therapy is shown in the upper part of the display of the pump by the icon for "Ramp and Taper Mode".

The screen shows the following:

**Ramp phase**

The pump now linearly increases the rate in the predefined time until it reaches the plateau rate and then automatically switches to continuous phase.

**Continuous phase**

The pump continuously infuses the same rate for a predefined time and then automatically switches to taper phase.

**Taper phase**

The pump linearly decreases the rate in the predefined time until it reaches the KVO rate.

**Note:** After starting infusion it is only possible to change rates, time and VTBI in the continuous phase.

By editing (increasing/decreasing) the plateau rate, the therapy is recalculated. With the increase/decrease of the plateau rate the volumes in the ramp phase, the continuous phase and the taper phase are increased/decreased. The continuous phase is shortened/prolonged to infuse the VTBI still completely with the end of the taper phase.

By editing the Ramp/Taper-Time, the therapy is recalculated. The Continuous Phase is extended/shortened to infuse the VTBI still completely until the end of the Taper phase.

By increasing/decreasing the VTBI, the continuous phase is prolonged/shortened to infuse the new entered VTBI completely with the end of the taper phase.

**Note:** The delivery of drugs can be stopped and started again in Ramp and Taper Mode at any time by pressing \( \text{SF} \). Ramp and Taper is stopped immediately without Taper phase and started without a new Ramp phase. This will not have any effect on the settings of the therapy.
Immediate Taper Down

By choosing the Immediate Taper Down Function the therapy can be ended with a taper phase before the originally defined VTBI is completely infused.

- Press \( \text{ during continuous phase.} \)
- Use \( \text{ to select Special Functions and press } \)
- Select Immediate Taper Down Function and confirm with \( \).
- Edit taper time by using \( \) and press \( \text{ to confirm.} \)
  - The pump automatically changes to Taper phase and linearly decreases the rate.

3.8 Program Mode

Program Mode is for infusion requiring a non-standard delivery pattern. The user defines a series of intervals (max. 12 intervals) by certain parameters (rate, time, volume) for each cycle.

The pump automatically gives each programmed period, one after the other.

Example:

![Diagram of Program Mode](image)

Program Mode should only be performed by an experienced user being familiar with the principles of the Program Mode function and properly trained in using the present device.

**Note:** The active Program Mode function always displays this icon in the Display \( \). \)

**Note:** Bolus function is disabled for Program Mode.

Set Profile Parameters: The therapy can be started directly via the drug library or via the Main Menu/Special functions.
Starting Program Mode via Drug Library:

**Note:** Program Mode settings have been configured in the Drug List Manager before and have been uploaded into the pump.

- Switch on pump with \( \text{on} \) and wait until self-check is finished.
- Insert disposable and use the drug library according to Instructions for Use.
- Select the desired drug with \( \text{t} \) and press \( \text{l} \).
- Select Program Mode with \( \text{k} \).

In the following screen the user has to confirm the number of steps for the therapy with \( \text{OK} \).

![Steps](image)

The settings for the steps of the infusion are shown on the display. These settings, configured in the Drug List Editor, need to be confirmed with \( \text{OK} \).

- To change the values, press \( \text{to change} \) and \( \text{OK} \) to confirm.
- Adjust VTBI with \( \text{q} \).

The pump can be started now by pressing \( \text{start} \).

Starting Program Mode via Special Function Menu:

- Switch on pump with \( \text{on} \) and wait until self-check is finished.
- Insert disposable.
- Go to Special Functions Menu and select Program Mode.
- Press \( \text{to enter} \) parameters and \( \text{OK} \) to confirm.
- Adjust VTBI with \( \text{q} \).

After entering all desired parameters the pump can be started by pressing \( \text{start} \).

In the upper part of the display the icon for "Program Mode" appears.

The screen shows the following:

The pump infuses the predefined rate in the predefined time for the current step.

Only the VTBI may be changed during an infusion that is running.

- Press \( \text{to check} \) upcoming Program Mode intervals in Main Menu.

It is possible to cancel one step of the running therapy. All following steps in the programmed sequence persist.

- Go to Main Menu by pressing \( \text{to go} \).
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- Use  to navigate through the Main Menu and select Current with  
- For checking upcoming intervals press  
- Select "Program Parameters" with  
- Go through all interval steps with  

Note: The delivery of drugs can be stopped and started again in the Program Mode at any time by pressing  . This will not have any effect on the settings of the therapy.

Number of cycles is defined by VTBI. Take care to set the VTBI in the correct relation to the volume of one Cycle. VTBI may need to be adjusted after changing the intervals.

The Main menu informs about the current interval. The configured parameters can be checked by Program Parameter Menu in Main.

3.9 Intermittent Mode

The Intermittent Mode consists of 2 phases. This phases will be repeated.
- Bolus phase: the configured bolus is active
- Rate phase: time during the therapy in which the entered rate is active

Example:

![Diagram of Intermittent Mode]

Intermittent Mode should only be performed by an experienced user being familiar with the principles of the Intermittent Mode and properly trained in using the present device.

Note: The active Multi Dose Mode function always displays this icon in the Display.

Note: Regular Bolus function is disabled for Intermittent Mode.
In Intermittent Bolus the bolus service settings are active. The pressure level is automatically set to max value.

Set Profile Parameters: The therapy can be started directly via the drug library or via the Main Menu/Special functions.

Starting Intermittent Mode via Drug Library:

**Note:** Intermittent Mode settings have been configured in the Drug List Manager before and have been uploaded into the pump.

- Switch on pump with and wait until self-check is finished.
- Insert disposable and use the drug library according to Instructions for Use.
- Select the desired drug with and press .

The pump now offers the possible therapy profiles.

- Select “Intermittent Mode” with and press . The therapy settings for “Intermittent Mode” are shown on the display.
- For changing the parameters, press to change and to confirm.

**Note:** Bolus rate is calculated by editable parameters. These parameters have to be checked by the user before starting the infusion.

The pump can be started now by pressing .

Starting Intermittent Mode via Special Function Menu:

- Switch on pump with and wait until self-check is finished.
- Insert disposable.
- Go to Special Functions Menu and select Intermittent Mode.
- Press to enter parameters and to confirm.

After entering all desired parameters the pump can be started by pressing .

In the upper part of the display the icon for “Intermittent Mode” appears.

In bolus phase the screen shows the following:

The pump now delivers the predefined bolus.

After the bolus phase the pump switches to rate phase and the screen shows the following:

The pump now delivers the predefined rate.
Note: To cancel bolus infusion in the Intermittet Bolus therapy at any time it is only possible with sf.

Note: The delivery of drugs can be stopped and started again in the Intermittent Mode at any time by pressing sf. During infusion it is possible to change the bolus volume, amount, VTBI as well as the time interval.

- Press $\cdot$.
- Use $\cdot$ to navigate through the parameter list and select the parameter to be changed with $\cdot$.
- Enter the new value and press $\cdot$.

The pump continues infusion.

Changing the bolus after start:

If the user edits the bolus the therapy progression changes.

- Press $\cdot$.
- Use $\cdot$ to select Bolus and press $\cdot$.
- Change Bolus by using $\cdot$ and press $\cdot$ to confirm.

The pump automatically recalculates all other settings of the therapy.

Changing the time interval after start:

If the user edits the time interval the therapy progression changes.

- Press $\cdot$.
- Use $\cdot$ to select Interval and press $\cdot$.
- Change Interval by using $\cdot$ and press $\cdot$ to confirm.

The pump automatically recalculates all other settings of the therapy.

### 3.10 Dose Over Time

Dose Over Time is used to administer a specific dose of antibiotics in a specific time. Dose Over Time is an own therapy and cannot be used in combination with another therapy. It can only be activated via the Drug List Manager. It can be used for standard infusion.

The active Dose Over time function is always symbolised with a characteristic symbol in the Display (•).

Note: Dose Over Time should only be performed by experienced users being familiar with the principles of the Dose Over Time function and properly trained in using the present device.

The infusion rate in Dose Over Time can not be changed. This parameter is a result of the total dose and the infusion time setting. Directly, after the Drug
selection, the infusion time and the total dose intended to be infused have to be set. If the drug library contains default values for these parameters, the default values are used as preset values.

If changes are necessary during infusion, the delivery can be controlled by changing the time. The pump calculates the new rate by using the remaining total dose and the remaining time. In the Main Menu total dose, time and VTBI can be changed, also during RUN-Mode. Other parameters (dose rate, basal rate, concentration, patient weight and patient height) cannot be changed.

**Note:** The KVO function and Bolus function are disabled during Dose Over Time.

**Note:** The feature Dose Over Time always requires the usage of dosing units (i.e., mg or mg/kg patient weight).

Before using Dose Over Time contact your local B. Braun representative!

**Starting Dose Over Time via Drug Library:**

**Note:** Dose Over Time settings have been configured in the Drug List Manager before and have been uploaded into the pump.

- Switch on pump with \(\circ\) and wait until self-check is finished.
- Insert disposab le and use the drug library according to the Instructions for Use.
- Select a drug by using \(\text{D}\) and press \(\text{↓}\).

The pump now offers the possible therapy profiles. Select “Dose over Time” with \(\text{D}\) and press \(\text{↓}\).

The editor for Total Dose is shown if a drug with therapy Dose over Time is selected from drug library and no default value for Total Dose was entered in library. The editor is also shown if the Total Dose is edited in the Main menu.

The editor for Time is shown if a drug with therapy Dose over Time is selected from drug library and no default value for Time was entered in library. The editor is also shown if the Time is edited in Main Menu.

Enter the total dose, if necessary, and confirm with \(\text{OK}\).

Enter the time, if necessary, and confirm with \(\text{OK}\).
The VTBI is calculated automatically and the following screen is displayed:

- Check calculated rate by using \( \text{key} \) for plausibility
- Start Dose Over Time by pressing \( \text{key} \).

Run Menu: The time is used to control the therapy. For this reason the remaining time is shown in big digits in menu Run. The parameter in the lower left corner can be scrolled. Set to Rate when leaving the pump.

Note: It is always possible to press the key \( \text{key} \) in the Run Menu and edit or check values in the Main Menu while the pump is delivering.

### 3.11 Take Over Mode (TOM)

Take Over Mode is a feature to support the user during syringe changes by automatically starting a second Perfusor® Space pump when the first has run empty. The second pump automatically takes over the infusion rate from the first pump.
Activation:

- Start an infusion of the desired medication from the drug library on a Perfusor® Space pump (see Section 3.3).
- Place a second a Perfusor® Space pump in an adjacent slot of the SpaceStation (either above or below).
  Note: Make sure that the pumps are correctly inserted in the SpaceStation.
- Navigate to the Drug Library on the second pump (Note: The drug library can be started over the Start Up or Special Functions Menu).
- Navigate through the list with ☐ and select the Care Unit with ◀. The Care Unit of the second pump must be the same as the first.
  Note: If you have already set the Care Unit once on your pump this step will be skipped for the next time.
- Navigate through the list with ☐ and select the patient profile with ◀. The patient profile of the second pump must be the same as the first.
  Note: If no profile is set, this step will be skipped.
- Navigate through the list with ☐ and select in alphabetical order (all drugs) or within a category with ◀. The drug selected in the second pump must be the same as the first.
- Navigate through the list with ☐ and select a concentration with ◀. The concentration in the second pump must be the same as the first.
- Confirm 'Use Take Over Mode' with ⬆.
- Check IV line of TOM2 is connected to the patient and that stopcocks are open.

Symbols:

TOM1 first Perfusor® Space pump
TOM2 second Perfusor® Space pump
Deactivation:

- Press the button on the pump.

Take Over Phase:

When the syringe is nearly empty, a pre-alarm will sound on the pump.

When the syringe is empty, the pump will automatically start infusing at the correct rate.

Note: Start-up behaviour is not influenced by TOM. See Chapter Start Up Graphs and Trumpet Curves.

Note: Please use a separate patient connection for Take Over Mode infusion (e.g. smallbore extension set) or use a back check valve for lines at the same access which are not used for Take Over Mode.

Note: When both pumps are turned off (e.g. for changing the syringe), pumps will be turned on again, both will start as a normal pumps and not in TOM modus with the latest TOM setting.

When both pumps are put in Standby during infusion and both are turned on again, both pumps will start with the latest drug library setting. TOM has to be activated separately.

TOM Requirements:

TOM will only be offered if the following requirements are met:

- Same drug selected on both pumps.
- Same drug concentration selected on both pumps.
First Perfusor® Space pump must have a running infusion with a drug from drug library.

TOM Hints:
The following TOM hints are to be observed:

<table>
<thead>
<tr>
<th>TOM Hint</th>
<th>Recommendation</th>
</tr>
</thead>
<tbody>
<tr>
<td>Take Over Mode not possible, Other pump not in Run menu</td>
<td>Ensure first Perfusor® Space pump is infusing</td>
</tr>
<tr>
<td>Take Over Mode not possible, Other pump in wrong therapy</td>
<td>Ensure first Perfusor® Space pump must be running in ‘continuous mode’ (i.e. ml/h or a dose rate; not KVO, PCA etc.)</td>
</tr>
<tr>
<td>Take Over Mode not possible, DataLock active in other pump</td>
<td>Deactivate Data Lock</td>
</tr>
<tr>
<td>Take Over Mode not possible, Data connection lost</td>
<td>Data connection must be active between pumps – check the positioning of pumps in the SpaceStation</td>
</tr>
<tr>
<td>Take Over Mode not possible, Different syringe sizes</td>
<td>Ensure both pumps must have a syringe of the same size</td>
</tr>
<tr>
<td>Take Over Mode not possible, Different care units selected</td>
<td>Ensure both pumps have the same Care Unit selected</td>
</tr>
<tr>
<td>Take Over Mode not possible, Different patient profiles</td>
<td>Ensure same patient profile selected on both pumps</td>
</tr>
<tr>
<td>Take Over Mode not possible, Software update required</td>
<td>Both pumps must have the same software version – contact your service department</td>
</tr>
<tr>
<td>Take Over Mode not possible, Mod.data update required</td>
<td>Both pumps must have the same modification data – contact your service department</td>
</tr>
<tr>
<td>Take Over Mode not possible, Invalid config. ‘Stop at syr. end’</td>
<td>Both pumps must have the same ‘Stop at syr. end’ settings – contact your service department</td>
</tr>
<tr>
<td>Take Over Mode not possible, TOM not enabled in other pump</td>
<td>Both pumps must have TOM activated – contact your service department</td>
</tr>
<tr>
<td>Take over of infusion has failed</td>
<td>Manually start second pump if appropriate</td>
</tr>
</tbody>
</table>

More information regarding alarm hints may be found in chapter 5.
Changes in TOM system:

<table>
<thead>
<tr>
<th>Change</th>
<th>Reaction</th>
</tr>
</thead>
<tbody>
<tr>
<td>Rate changed in <strong>TOM1</strong> pump</td>
<td>No user interaction necessary, <strong>TOM2</strong> will start infusion at new rate when syringe is empty.</td>
</tr>
<tr>
<td><strong>TOM1</strong> pump is stopped</td>
<td><strong>TOM2</strong> pump shows “connection lost – TOM aborted” alarm. TOM may be reactivated by pressing <strong>OK</strong> and then <strong>△</strong> when prompted “Return to Take Over M.”</td>
</tr>
<tr>
<td><strong>TOM4</strong> pump is put in standby</td>
<td></td>
</tr>
<tr>
<td>VTBI ended in <strong>TOM1</strong></td>
<td></td>
</tr>
</tbody>
</table>
AUTOPROGRAMMING

Note: All normal pump functions remain in place when orders are received via autoprogramming.

The pump can accept drug orders wirelessly from the EHR system or from SpaceStation with SpaceCom. The workflow to accept an order wirelessly will vary depending on your EHR vendor.

- Using the hand held device or lap top, review the order and follow your hospital protocol for scanning the bag/syringe, pump, patient and nurse (optional).
- Once order is confirmed on the hand held or laptop, prompt EHR to send order directly to pump. The order will arrive and appear on the pump within 10 seconds.
- Ensure pump is in the Main Menu, passive mode or Standby.
- New Order message will appear with drug name and mode.

![Order received for PRIM: Normal Saline 0.9%](image1)

- Press OK to accept or C key to cancel order and respond to prompt.
- Select Care Unit and Patient Profile as in Drug Library programming.
- Pump will search for Drug Library match.

Note: If no drug library match, which may be due to no matching name, concentration or dosing units, pump displays reason for no match and depending on your hospitals configuration either allows manual programming outside the drug library or rejects order completely. An order that is confirmed outside the drug library will have a triangle with an exclamation point on display to indicate there are no drug library settings.

![No concentration match found in drug from library Confirm](image2)

- Scroll to each value to confirm using arrow keys.

![Check values](image3)
Note: Order may be cancelled prior to confirming order.

Order: Normal S:
Cancel incoming order? Yes ▲No ▼

Once all values are confirmed, the Main Menu is displayed.

Note: Soft Limit alert will be issued if value exceeds any soft limits set in drug library, soft limit may be overridden or value re-programmed per institutional policy. Order will be rejected if hard limit is exceeded. (except in circumstance where pump service program is not set to perform drug library match for auto-programming).

For PRIMary Orders (Either 'Continuous' oder 'Dose over Time'):

Note: The first order sent send as 'Continuous' is always considered as the PRIMary infusion, subsequent orders will be considered PIGGYback.

Note: Order sent as 'Dose over Time' is always considered the PRIMary infusion, no subsequent order can be received. Additional, no updates can be received for 'Dose over Time'.

- Press Start/Stop key to start infusion.

Updates to Current Primary Infusion

Updates may be received for PRIMary infusions while pump is running or stopped and while in PRIMary or PIGGYback.

While in PRIMary:

- Update icon will appear on display, follow on screen prompts to accept or cancel the order. Confirmation screen will indicate both OLD and NEW value for parameter(s) that changed.

While in PIGGYback:

- Message will appear on top of display indicating update is available for PRIMary.
- Press key to view order.
Follow prompt, pressing 🍃 to accept order or 🍇 key to cancel and hold order for later.

New Primary Infusion:
- To accept a new PRIMary order, stop infusion and clear current PRIMary infusion by pressing 🍇 key and responding “yes” to clear current infusion.

PIGGYback Orders:
Orders received after PRIMary has been set will be for PIGGYback infusions, follow prompts on screen to stop the PRIMary to accept the PIGGYback order.

- Confirm order values as above for PRIMary orders.
- Respond to prompts to check bag height and clamps prior to starting PIGGYback.

New PIGGYback order while PIGGYback is Infusing:
- Follow display prompts to stop current infusion.

**Note:** A PIGGYback order may be held for later by pressing 🍇 key to cancel order and answering yes to “hold for later”.

Order for SEC: cefaZOL
Hold order for later? Yes 🌟 No 🍇
Note: Changing values on any incoming order may only be done after confirming all values. Once all values are confirmed you may scroll to any value and open editor with ↓ to change value. Alternately, order may be cancelled and request made for revised order to be sent.

Note: If pump is placed in standby while order is pending new order will flash on top of stand by display, press ✂ key to accept order (pump will come out of standby).
The options functions may be selected and changed while the pump is infusing or stopped. To edit a menu item, select “Options” in the Main Menu and press . Then select desired function with and follow the Instructions for Use as described.

5.1 Occlusion Pressure

The higher the pressure level is set at, the higher the pressure level must rise before triggering an occlusion pressure alarm.

Using the occlusion pressure, the alarm sounding period can be kept short in the event of system occlusions. It generally applies that the set pressure should always be set higher than the system pressure. If pressure alarms occur in a pressure level without a system occlusion, the pressure level must be adjusted upward. In order to be able to ensure short alarm times, a low pressure level should be started with and the pressure level increased until the syringe starts up.

Depending on the different influences, such as friction fluctuations of the syringes, tube length, tube diameter, viscosity of the liquid and the filter used in the system set-up, adjustments may need to be made to the pressure level.

- Enter pressure in Options Menu by pressing .
- Choose between nine pressure levels (1=lowest level; 9=highest level) by pressing or and confirm entry with . Pressure levels and equivalent mmHg are displayed when left arrow is pressed while in pressure menu.

Note: The pressure will remain at set level until changed by user unless the drug selected had a pressure level set in the drug library. When pump is powered off pressure level returns to default value set in service program when powered back on unless drug selected has a different pressure level set in the drug library.

The top line is the current infusion pressure. The bottom dashed line shows the pressure alarm setting, currently 5 out of 9 which is represented by 5 dashed. The picture shows a current pressure of ~30% of the pressure level 5.

If occlusion pressure levels lower than the level 1 are needed, it can be activated via the service tool.
Please contact your local sales representative for further data if you use pressure levels below lever 1.

The editor is extended by maximal 3 dashes.

Confirm new pressure level with \( \text{OK} \) and go back to Options Menu.

In the Run Menu the top line shows the current infusion pressure. The bottom line and the 3 dashes before the symbol shows the pressure alarm settings.

5.2 OccluGuard & Pressure Leap/Drop detection

OccluGuard

OccluGuard speeds up time to alarm when an occlusion is present. Occlusions can be caused by problems in IV access (e.g. a blocked catheter), problems in the infusion setup (e.g. closed stopcocks) or ‘syringe occlusions’ i.e. Due to varying syringe tolerances of syringes from other manufacturers, an OccluGuard alarm may occur because of high syringe friction forces. OccluGuard can be used with all syringe sizes and drugs, but is ideally suited to infusions at low rates and/or with drugs of short half life (e.g. Catecholamines).

OccluGuard activation / deactivation from the Main Menu

- Go to Options Menu and press \( \leftarrow \).
- Navigate through the list with \( \uparrow \) and select OccluGuard.
- OccluGuard can be activated with \( \uparrow \) and deactivated with \( \downarrow \).

Pressure Leap/Drop detection

The pressure leap/drop software detects sudden increases and decreases in infusion pressure respectively which can be caused by problems in IV access, or changes in pump position in the SpaceStation.

Pressure Leap/Drop detection activation / deactivation from the Main Menu

- Go to Options Menu and press \( \leftarrow \).
- Navigate through the list with \( \uparrow \) and select “Pr. leap/drop”.

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- Navigate to Off with ← → and press OK to deactivate pressure leap/drop.
- Navigate to high (2mmHg), medium (8mmHg) or low (20mmHg) with ← → and press OK to activate pressure leap/drop.

Note: after a restart of the pump, these settings remain at the levels set before the restart.

Area of application

OccluGuard and pressure leap/drop are active below the following infusion rates. Should the rates increase, the OccluGuard Inactive symbol (X) is shown in run screen.

<table>
<thead>
<tr>
<th>Syringe Size</th>
<th>Maximum Rate (typical)</th>
</tr>
</thead>
<tbody>
<tr>
<td>50 ml</td>
<td>30 ml/h</td>
</tr>
<tr>
<td>20 ml</td>
<td>14 ml/h</td>
</tr>
<tr>
<td>10 ml</td>
<td>9.8 ml/h</td>
</tr>
</tbody>
</table>

Symbols

OccluGuard symbol
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### OPTIONS

**OccluGuard Status**

<table>
<thead>
<tr>
<th>OccluGuard Symbol</th>
<th>Meaning</th>
<th>Recommendation</th>
</tr>
</thead>
<tbody>
<tr>
<td><img src="image" alt="ok" /></td>
<td>OccluGuard is active. Infusion is running stably</td>
<td>n/a</td>
</tr>
<tr>
<td><img src="image" alt="clock" /></td>
<td>Pending – OccluGuard has not enough data</td>
<td>n/a</td>
</tr>
<tr>
<td><img src="image" alt="times" /></td>
<td>OccluGuard Inactive</td>
<td>OccluGuard will automatically reactivate as soon as infusion rate drops below threshold levels – see above.</td>
</tr>
<tr>
<td><img src="image" alt="bell" /></td>
<td>Pressure rise detected</td>
<td>Occlusion has been detected</td>
</tr>
<tr>
<td><img src="image" alt="no symbol" /></td>
<td>OccluGuard is deactivated</td>
<td>Activate OccluGuard – see below</td>
</tr>
</tbody>
</table>

**Pressure Leap/Drop Symbol**

<table>
<thead>
<tr>
<th>Pressure leap/drop Symbol</th>
<th>Meaning</th>
<th>Recommendation</th>
</tr>
</thead>
<tbody>
<tr>
<td><img src="image" alt="bell" /></td>
<td>Sudden pressure leap detected</td>
<td>Check IV access and IV setup</td>
</tr>
<tr>
<td><img src="image" alt="bell" /></td>
<td>Sudden pressure drop detected</td>
<td>Check IV access and IV setup</td>
</tr>
</tbody>
</table>

**Note:**

- The OccluGuard status can be checked in the status menu.
- **Perfusor® Space** continues to infuse during OccluGuard and Pressure leap/drop alarms.
- The existing occlusion alarm pressure levels are unaffected by OccluGuard.
When a change is made to the infusion system (e.g. addition or removal of a pump to a SpaceStation, a change of infusion rate, a bolus application) the OccluGuard and pressure leap/drop are temporarily set to 'pending' (_succes) to allow the system to reach a hydrostatic balance, and so prevent false alarms.

### 5.3 Data Lock

The data lock function protects the device against unauthorized access. It is recommended to adapt the four digit code for level 1 and 2 from the default setting (9119), using the service program. There are three security levels.

**Level 1:**
A modification of values as well as a bolus application are not possible but a change of the disposable can be conducted. It is possible to navigate through all menus and status data can be checked. Starting, interrupting and switching the pump off is possible.

**Level 2:**
This level has the same performance characteristic as described under level 1 and additional will not allow the change of disposable. In order to prevent a data lock alarm the correct code must be entered within 20 sec after the pump was stopped. Changing the disposable and switching the pump off is only possible after the code was entered.

**Level 3:**
This level will allow starting and stopping the pump as well as switching off. The code for this level may be different for each drug and is defined in the drug list. A change of the syringe, however, is possible by using the code defined for the other levels. An overview about the differences between the levels 1, 2 and 3 is given by the following table.

<table>
<thead>
<tr>
<th>Event</th>
<th>Level 1</th>
<th>Level 2</th>
<th>Level 3</th>
</tr>
</thead>
<tbody>
<tr>
<td>Change of disposable</td>
<td>✔️</td>
<td>✗</td>
<td>✔️ with code for level 1/2</td>
</tr>
<tr>
<td>Start of infusion</td>
<td>✔️</td>
<td>✗</td>
<td>✗</td>
</tr>
<tr>
<td>Change of parameters</td>
<td>✗</td>
<td>✗</td>
<td>✗</td>
</tr>
<tr>
<td>Stop of infusion</td>
<td>✔️</td>
<td>✔️</td>
<td>✔️</td>
</tr>
<tr>
<td>Switching off pump / Standby</td>
<td>✔️</td>
<td>✗</td>
<td>✗</td>
</tr>
<tr>
<td>PCA bolus with pump-based bolus button</td>
<td>✗</td>
<td>✗</td>
<td>✔️</td>
</tr>
<tr>
<td>Customisable screen</td>
<td>✗</td>
<td>✗</td>
<td>✔️</td>
</tr>
<tr>
<td>Acoustic feedback of demanded boli</td>
<td>✗</td>
<td>✗</td>
<td>✔️</td>
</tr>
<tr>
<td>Indicates denied PCA boli</td>
<td>✔️</td>
<td>✔️</td>
<td>✗</td>
</tr>
</tbody>
</table>


✔️ = possible | ✗ = not possible | ✗= followed by standby-alarm
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Activation of the function:

- Open data lock in Options Menu with \( \textgreater \).
- Select between level 1, 2 or 3 (if activated) with \(<\) and \(>\) and confirm with \(\text{OK}\).
- Enter code with \(\text{OK}\) and press \(\text{OK}\) in order to activate data lock.

Changes to the protected values and the bolus function which are marked with \(\text{y}\) are only possible after entering the code. After 20 sec in the Main Menu, Status Menu, Special Functions Menu and Options Menu the lock will be activated again. If the wrong code is entered twice the pump will switch into the last menu. If the wrong code is entered twice again the pump will go into an audible alarm, a nurse call will go off and the yellow LED blinks. If a target value was reached while data lock is active a new start of the pump is only possible after entering the code.

In order to deactivate the function, select “Off” in the data lock, press \(\text{OK}\), enter the code and press \(\text{OK}\) again.

5.4 Bolus Rate

- Open bolus rate in Options Menu with \(\textless\).
- Change bolus rate with \(\text{Q}\) and confirm setting with \(\text{OK}\).

*Note:* Set bolus rate according to therapy requirements. Take care not to overdose!

Given a bolus rate of 1800 ml/h, e.g. 0,5 ml are reached within just one second.

5.5 KVO-Mode

After reaching a preselected VTBI/time, the pump can continue the infusion with a predefined KVO-rate (see “Technical Data”). The duration of the KVO-infusion is set via the service program.

- Open KVO-Mode in Options Menu with \(\textless\).
- Answer Yes/No question with \(\text{a}\), to activate KVO.

5.6 Contrast / Display Light / Keypad Light

Contrast as well as display- and keypad light can be adjusted individually according to the lighting conditions.

- Open contrast/display light/keypad light in Options Menu by pressing \(\text{q}\).
- Choose between 9 contrast- and display light levels with \(<\) or \(>\) and confirm with \(\text{OK}\). For use with light sensitive drugs the keypad- respectively syringe light can be completely turned off.
5.7 Alarm Volume

Chose between 9 different alarm volume levels.

- Open alarm volume in Options Menu with ▼.
- Set volume with ▼ or ▶ and confirm entry with OK.

5.8 Date / Time

- Open date/time in Options Menu with ▼.
- Change date/time with ▼▼ and confirm with OK.

5.9 Macro Mode

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

- Open macro mode in Options Menu with ▼.
- Answer Yes/No question by pressing ▲ to activate the macro mode.

For quick activation of macro mode: Press and hold ▶ while the pump is infusing until the font size changes.

5.10 Language

This function enables a change of the pump language.

- Open language in the Options Menu with ▼.
- Select language with ▶, then press ▼.
- Confirm Yes/No with ▲.
ALARMS

The Perfusor® Space is equipped with a 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>off</td>
<td>device alarm and alarm code (see service program)</td>
</tr>
<tr>
<td>Operating Alarm</td>
<td>yes</td>
<td>flashes</td>
<td>off</td>
<td>see alarm description</td>
</tr>
<tr>
<td>Pre-Alarm</td>
<td>yes</td>
<td>off</td>
<td>constant on</td>
<td>see alarm description</td>
</tr>
<tr>
<td>Reminder Alarm</td>
<td>yes</td>
<td>off</td>
<td>constant on</td>
<td>see alarm description</td>
</tr>
<tr>
<td>Alarm Hint</td>
<td>no</td>
<td>off</td>
<td>off</td>
<td>see alarm description</td>
</tr>
</tbody>
</table>

<table>
<thead>
<tr>
<th>Audible signal</th>
<th>Optical signal</th>
<th>Staff call</th>
<th>User confirmation</th>
</tr>
</thead>
<tbody>
<tr>
<td>Red LED</td>
<td>Yellow LED</td>
<td>Text</td>
<td></td>
</tr>
</tbody>
</table>

6.1 Device Alarms

When a device alarm occurs the infusion is immediately stopped. Press \( \text{O} \) to switch off the device. Then switch the device on again. In case of a repeated device alarm you must disconnect from the patient, open the front door of the pump and take out the disposable. The device needs to be handed to the service department.

6.2 Pre-Alarms and Operating Alarms

Pre-alarm:
Pre-alarm occurs a few minutes (dependable on service settings, excluding OccluGuard and pressure leap/drop pre-alarm) prior to operating alarms. During pre-alarm an audible tone sounds, the yellow LED is constantly on and a staff call is activated (optional). The display message varies depending on the alarm reason. The signal tone and the staff call are turned off with \( \text{O} \). Display and LED stay in pre-alarm until the operating alarm goes off. Pre-alarm don’t lead to an interruption of the infusion.
<table>
<thead>
<tr>
<th>Display message</th>
<th>Pre-alarm reason</th>
</tr>
</thead>
<tbody>
<tr>
<td>&quot;Syringe nearly empty&quot;</td>
<td>Very little fluid is left in syringe.</td>
</tr>
<tr>
<td>&quot;VTBI near end&quot;</td>
<td>The preselected volume is nearly infused.</td>
</tr>
<tr>
<td>&quot;Time near end&quot;</td>
<td>The preselected time is almost over.</td>
</tr>
<tr>
<td>&quot;Battery nearly empty&quot;</td>
<td>The battery is almost discharged.</td>
</tr>
<tr>
<td>&quot;KVO mode&quot;</td>
<td>Volume/time are reached and the pump continues the infusion at the KVO-rate.</td>
</tr>
<tr>
<td>&quot;Communication error&quot;</td>
<td>The pump is located in a system in which at least one device is incompatible or defective. The use of this device in a system is not permitted. The system is to be checked by a service technician.</td>
</tr>
<tr>
<td>&quot;Pressure rise detect.&quot;</td>
<td>OccluGuard has detected an occlusion. Check IV access, IV setup and syringe for cause of occlusion. Should the cause of the alarm be removed, the alarm will stop automatically. Due to varying syringe tolerances of syringes from other manufacturers, a pressure alarm may occur because of high syringe friction forces.</td>
</tr>
<tr>
<td>&quot;Pressure drop detect.&quot;</td>
<td>A sudden pressure drop has been detected – check IV access.</td>
</tr>
<tr>
<td>&quot;TOM pending&quot;</td>
<td>Very little fluid is left in syringe, infusion will be handed over to second Perfusor® Space pump when syringe is empty (Take Over Mode only).</td>
</tr>
<tr>
<td>&quot;TOM aborted&quot;</td>
<td>Take Over Mode has been aborted (Take Over Mode only)</td>
</tr>
</tbody>
</table>

Except OccluGuard and pressure leap/drop pre-alarms, a stopwatch on the display counts down the remaining time (depending on the service program, between 0-30 min). After that, the pump changes to the operating alarm.

The pre-alarms “VTBI near end” (volume preselection) and “Time near end” (time preselection) can be deactivated via the service program.

**Operating alarms:**
Operating alarms lead to an interruption of the infusion. An audible tone sounds, the red LED flashes and a staff call is activated.

The display states "Alarm", the reason for the operating alarm and gives the option to either confirm the alarm by pressing **OK** or mute the alarm by pressing **Mute**. If the alarm is muted, the alarm message will remain on the screen until it is confirmed by pressing **OK**. After 2 minutes, if the alarm has not been confirmed by pressing **OK**, the audible tone will sound again. The alarm tone,
the alarm message, and the staff call (optional) are all cleared by pressing \textit{OK}. Corrections should be made in accordance with the alarm reason.

<table>
<thead>
<tr>
<th>Display message</th>
<th>Alarm reason</th>
</tr>
</thead>
<tbody>
<tr>
<td>“Syringe empty”</td>
<td>There is no fluid left in the syringe. Due to varying syringe tolerances of syringes from other manufacturers, some fluid may be left inside the syringe. Restarting the infusion leads to a complete depletion of the syringe and shut-off via the pressure sensor. Perform syringe change as described in 1.4.</td>
</tr>
<tr>
<td>“VTBI infused”</td>
<td>The preselected volume was infused. Continue therapy or select new therapy.</td>
</tr>
<tr>
<td>“Time expired”</td>
<td>The preselected time has ended. Continue therapy or select new therapy.</td>
</tr>
<tr>
<td>“Battery empty”</td>
<td>The battery pack is discharged. Connect device with mains and/or exchange battery pack. The battery alarm will be on for 3 min. Then the pump will automatically turn off.</td>
</tr>
<tr>
<td>“KVO finished”</td>
<td>KVO is reached. Continue with old or set new therapy.</td>
</tr>
<tr>
<td>“Pressure high”</td>
<td>An occlusion occurred in the system. The set pressure level was exceeded. A bolus reduction is automatically initiated by the pump. Check if syringe is empty, kinks are in tubing and tubing isn’t damaged, IV patency and filter patency. Increase occlusion pressure if necessary. Due to varying syringe tolerances of syringes from other manufacturers, a pressure alarm may occur because of high syringe friction forces.</td>
</tr>
<tr>
<td>“Syringe not correctly inserted”</td>
<td>The wings of the syringe are not properly inserted. Insert syringe according to description in “Overview Perfusor® Space” as well as 1.1.</td>
</tr>
<tr>
<td>“Syringe holder”</td>
<td>The syringe holder was opened during a running infusion. Close syringe holder.</td>
</tr>
<tr>
<td>“Battery cover removed”</td>
<td>The battery cover is not properly engaged on the battery compartment. When pushing on the battery cover listen for “click”.</td>
</tr>
<tr>
<td>“Drive blocked”</td>
<td>An external interference kept the drive unit from advancing. Basically prevent all external interferences. Consider “Patient Safety”.</td>
</tr>
</tbody>
</table>
### “Calibrate device”
- Pump calibration data have changed (e.g. after an update).
- Recalibrate device via the service program.

### “Claw malfunction”
- The emergency release button was pressed and the claws manually opened.
- Take out syringe and contact technical service department.

### “Plunger plate not prop. fixed”
- The syringe plunger plate does not attach to the plunger plate sensor on the pump.
- Check system for negative pressure and eliminate cause. Consider “Patient Safety”.

### “Standby Time expired”
- The set standby time has ended.
- Set new time or continue with previously set therapy.

### “No battery inserted”
- It is not possible to use the pump without a battery pack.
- Turn off pump and insert battery pack according to description “Overview Perfusor® Space”.

### “Data were reset”
- Therapy and pump settings could not be restored.
- Enter therapy and pump settings anew.

### “Therapy data were reset”
- Therapy data could not be restored.
- Enter therapy anew.

### “Data Lock”
- An attempt was made to stop or switch the pump off without entering the code.
- Enter the correct code in order to continue the therapy or in order to turn the pump off.

### “Connection lost – TOM aborted”
- Data connection between TOM pumps in the SpaceStation has been lost and TOM has been aborted (Take Over Mode only).
- TOM may be reactivated by pressing [OK] and then ▲ when prompted “Return to Take Over M.”

### “Infusion taken over by other pump”
- Infusion been handed over to second Perfusor® Space pump (Take Over Mode only)

The red LED extinguishes with the acknowledgement of the alarm.

**Caution:** If a wrench ⚒ is displayed and/or a yellow, red and blue LED blink then the pump is in the service mode and is not permitted to be used on a patient. The pump is then to be checked by a service technician.

**Caution:** If the 🔴 is constantly displayed in the headline, the audible alarm for the pre- and operating alarm is silenced for a predefined time via service tool. Only the visual alarm is still displayed at the pump. After the predefined time is elapsed the pump provides the audible alarm.
6.3 Reminder Alarms

Reminder alarms only occur in two cases:
1. A syringe is inserted, the pump doesn’t administrate, no value is being edited and the device is not operated for two minutes.
   An acoustic tone sounds, the yellow LED is constantly on and a staff call is activated.
   a) The display states “Reminder alarm!”
   b) The display states “Config. not finished!”
      Confirm alarm with \( \text{OK} \) and continue to set therapy/Start Up configuration.
2. A value edition was started but not finished and confirmed. This is also possible with a missing disposable.
   An acoustic tone sounds, the display states “Value not accepted”, the yellow LED is constantly on and a staff call is activated.
   Confirm alarm with \( \text{OK} \) and continue to set therapy.

6.4 Alarm Hints

If improper entries are made the display states corresponding hints (e.g. “Bol.rate out of range”; “Download failed”; “The parameter can not be modified”). These hints disappear after a few seconds and don’t need to be confirmed.
BATTERY OPERATION AND MAINTENANCE

The battery has an operating lifetime of 8 hours at 25 ml/h when new. For optimal treatment of the battery, the device is equipped with protection against overcharge and deep depletion. The battery pack is charged by the pump during connection to mains. When disconnected from mains or in case of power failure, the pump automatically switches to battery power.

Note: Prior to a longer storage of the pump (5 months) the battery pack must be completely charged and then removed from the pump. Before changing the battery, always disconnect the pump from the patient and switch off the device.

The battery status indicator is a trend display (low, medium, high). For more detailed information on the current battery capacity (operating time in hours and minutes) please refer to menu item “Batt. Cap.” in the Status Menu of the Perfusor® Space.

Caution: The display of the battery operating time on the pump is an approximate value based on the current delivery rate. Changes in the delivery rate may affect the battery operating time.

Syringe change procedures require a high power consumption. A sudden break down of the battery operating time can be possible with an aged battery. In this case the battery has to be replaced by a new one.

If highly potent drugs are to be given over an extended time without mains power, it is recommended to have a fully charged reserve pump at hand.

Note: In case of ESD, Pump may need plugged into wall outlet to re-start the battery.

Attention: If the battery module is stored for long periods of time outside the pump, it is recommended to fully charge the battery and store it at room temperature.

Caution: Batteries may explode or leak if they are opened or incinerated. Consider disposal directions!

Important information for battery self-check:

If the battery symbol is blinking during mains operation, the battery has less then 30 minutes remaining capacity. In this case, the pump should not be disconnected from mains. If it is necessary to disconnect the pump from mains power for urgent reasons, the user should check to ensure if the battery capacity is sufficient for the proposed use. When the battery symbol blinks permanently (>1h), the battery must be checked by a technician and replaced if necessary.

Directions for optimal battery use:

The actual battery life may vary due to
- ambient temperature
- varying load (e.g. frequent boluses).

The optimal life time of a battery pack will only be reached if it’s completely discharged from time to time. A maintenance mode which conducts this battery maintenance is built in. This function should be activated once a month. Furthermore:
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- If a battery, which is not completely discharged, is charged several times, its capacity can be reduced.
- Under normal temperature conditions a battery can be charged and discharged approx. 500 times before its lifetime decreases.
- When the pump is not connected to mains power the battery discharges itself slowly. This can occur even when the pump is not operating.
- The battery operating time can only be realized if the pump operates continuously with a fully charged battery at room temperature.

Battery maintenance:
To accurately balance the battery capacity a cyclical battery maintenance is necessary. The frequency of battery maintenance may be set in the service tool. The battery maintenance mode detects a possible capacity loss (e.g. through ageing of the battery pack) and then the capacity/running time will be calculated anew. After a longer storage time or a longer operation without battery maintenance it can happen that the battery pre-alarm time can no longer be maintained. In this case it is necessary to perform a battery maintenance.

To initiate the discharge process the message „Battery maintenance“ and the OK–key will be displayed after switching the pump off. By pressing OK and the discharge process will start. The process is interrupted by switching the pump on again. If the battery maintenance is to be continued a new activation is necessary. After completely discharging the battery it will be completely charged again. The total duration of the battery maintenance process takes approx. twelve hours.

Caution: Please take into account that, if the battery maintenance has not been completed there is a possibility of a reduced battery operating time.

Replacing batteries:
The Battery Pack SP can be exchanged by any user. No special qualification is required.

All rechargeable batteries exhibit a reduction in capacity as they age. This aging is dependent on several factors including charging cycles, temperature and battery usage.

It is recommended to periodically check the function of the battery. A battery should no longer be used if a change of syringe leads to a “Battery nearly empty“ or a “Battery empty” alarm when it is fully charged.

Caution: Batteries may explode or leak if they are opened or incinerated. Consider disposal directions!
COMPATIBLE SYRINGES

The syringe types listed in the following tables can be used with the Perfusor® Space. Please refer to the listed material number (Mat. No.) to ensure specific syringe brand compatibility.

The Time to Occlusion alarm has been measured at 5 ml/h. The measured data are typical values which may vary because of possible syringe tolerances.

**Note:** At a rate of 0,01 ml/h, the time of occlusion alarm is > 3 hours.

<table>
<thead>
<tr>
<th>Manufacturer / item #</th>
<th>Syringe Size</th>
<th>Pressure Level</th>
<th>Max. Time to occlusion at 0.01 ml/h</th>
<th>Max. Time to occlusion at 1 ml/h</th>
<th>Max. Time to occlusion at 5 ml/h</th>
<th>Max. Post occlusion bolus volume</th>
</tr>
</thead>
<tbody>
<tr>
<td>B. Braun Omnifix®</td>
<td></td>
<td></td>
<td>hr:min:sec</td>
<td>hr:min:sec</td>
<td>hr:min:sec</td>
<td>ml</td>
</tr>
<tr>
<td>4617022V</td>
<td>3 ml</td>
<td>P1</td>
<td>&gt; 3 h</td>
<td>---</td>
<td>00:00:45</td>
<td>&lt; 0.1</td>
</tr>
<tr>
<td></td>
<td></td>
<td>P9</td>
<td>&gt; 3 h</td>
<td>---</td>
<td>00:02:30</td>
<td>&lt; 0.1</td>
</tr>
<tr>
<td>4617053V</td>
<td>5 ml</td>
<td>P1</td>
<td>&gt; 3 h</td>
<td>---</td>
<td>00:00:45</td>
<td>&lt; 0.1</td>
</tr>
<tr>
<td></td>
<td></td>
<td>P9</td>
<td>&gt; 3 h</td>
<td>---</td>
<td>00:03:00</td>
<td>0.1</td>
</tr>
<tr>
<td>4617100V</td>
<td>10 ml</td>
<td>P1</td>
<td>&gt; 3 h</td>
<td>---</td>
<td>00:01:30</td>
<td>&lt; 0.1</td>
</tr>
<tr>
<td></td>
<td></td>
<td>P9</td>
<td>&gt; 3 h</td>
<td>---</td>
<td>00:03:40</td>
<td>0.1</td>
</tr>
<tr>
<td>4617207V</td>
<td>20 ml</td>
<td>P1</td>
<td>&gt; 3 h</td>
<td>---</td>
<td>00:01:00</td>
<td>&lt; 0.1</td>
</tr>
<tr>
<td></td>
<td></td>
<td>P9</td>
<td>&gt; 3 h</td>
<td>---</td>
<td>00:06:00</td>
<td>0.1</td>
</tr>
<tr>
<td>4617304F</td>
<td>30 ml</td>
<td>P1</td>
<td>&gt; 3 h</td>
<td>---</td>
<td>00:01:00</td>
<td>&lt; 0.1</td>
</tr>
<tr>
<td></td>
<td></td>
<td>P9</td>
<td>&gt; 3 h</td>
<td>---</td>
<td>00:09:00</td>
<td>0.2</td>
</tr>
<tr>
<td>4617509F</td>
<td>50 ml</td>
<td>P1</td>
<td>&gt; 3 h</td>
<td>00:12:50</td>
<td>00:02:30</td>
<td>0.2</td>
</tr>
<tr>
<td></td>
<td></td>
<td>P9</td>
<td>&gt; 3 h</td>
<td>01:30:30</td>
<td>00:16:00</td>
<td>0.2</td>
</tr>
</tbody>
</table>

Please contact your local sales representative for further data of other syringe manufacturer.

**Note:** Values for “Max. time to occlusion at 1ml/h” are given for representative syringe types only.

**Note:** Values for “Max. Post occlusion bolus volume” are given for small bore and microbore tubing only. For standard bore tubing the values can be up to 0.4 ml in addition, depending on the pressure settings.
### COMPATIBLE SYRINGES

#### Chapter 8

**Manufacturer:** B. Braun

<table>
<thead>
<tr>
<th>Syringe Type</th>
<th>Omnifix 2 ml</th>
<th>Omnifix 5 ml</th>
<th>Omnifix 20 ml</th>
<th>Omnifix 30 ml</th>
<th>Omnifix 50 ml</th>
</tr>
</thead>
<tbody>
<tr>
<td>B. Braun</td>
<td>4617029V</td>
<td>4617053V</td>
<td>4617207V</td>
<td>4617304F</td>
<td>4617509F</td>
</tr>
<tr>
<td>Mat. No. 1)</td>
<td>typ.</td>
<td>typ.</td>
<td>typ.</td>
<td>typ.</td>
<td>typ.</td>
</tr>
<tr>
<td>Time to Occl. 2)</td>
<td>0:39</td>
<td>0:58</td>
<td>1:04</td>
<td>1:13</td>
<td>1:16</td>
</tr>
</tbody>
</table>

**Manufacturer:** B. Braun

<table>
<thead>
<tr>
<th>Syringe Type</th>
<th>Omnifix 3ml 3)</th>
<th>Omnifix 10ml LL</th>
</tr>
</thead>
<tbody>
<tr>
<td>B. Braun</td>
<td>4617022V</td>
<td>4617100V</td>
</tr>
<tr>
<td>Mat. No. 1)</td>
<td>A/P 4617022V-03</td>
<td>A/P 4617100V-03</td>
</tr>
<tr>
<td>US 4610303V-02</td>
<td>US 4617100V-02</td>
<td></td>
</tr>
<tr>
<td>Time to Occl. 2)</td>
<td>typ.</td>
<td>typ.</td>
</tr>
<tr>
<td>P 1</td>
<td>0:25</td>
<td>0:53</td>
</tr>
<tr>
<td>P 9</td>
<td>1:43</td>
<td>3:50</td>
</tr>
</tbody>
</table>

**Manufacturer:** B. Braun

<table>
<thead>
<tr>
<th>Syringe Type</th>
<th>OPS 20 ml</th>
<th>OPS 50 ml</th>
</tr>
</thead>
<tbody>
<tr>
<td>B. Braun</td>
<td>8728615</td>
<td>8728810F-06</td>
</tr>
<tr>
<td>Mat. No. 1)</td>
<td>typ.</td>
<td>typ.</td>
</tr>
<tr>
<td>Time to Occl. 2)</td>
<td>00:50</td>
<td>05:50</td>
</tr>
<tr>
<td>P 1</td>
<td>1:34</td>
<td>15:27</td>
</tr>
<tr>
<td>P 9</td>
<td></td>
<td></td>
</tr>
</tbody>
</table>
## Chapter 8

### COMPATIBLE SYRINGES

**Manufacturer:**
**Medtronic / Covidien**

<table>
<thead>
<tr>
<th>Syringe Type</th>
<th>Medtronic / Covidien</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></td>
<td>8881-513934</td>
<td>8881-516937</td>
<td>8881-512878</td>
<td>8881-520657</td>
<td>8881-535762</td>
<td>8881-560125</td>
</tr>
<tr>
<td></td>
<td></td>
<td>8881-713005</td>
<td>8881-716008</td>
<td>8881-712023</td>
<td></td>
<td></td>
<td>8881-760089</td>
</tr>
</tbody>
</table>

<table>
<thead>
<tr>
<th>Time to Occl. 2)</th>
<th></th>
<th></th>
<th></th>
<th></th>
<th></th>
<th></th>
<th></th>
</tr>
</thead>
<tbody>
<tr>
<td>P 1 [mm:ss]</td>
<td>typ.</td>
<td>typ.</td>
<td>typ.</td>
<td>typ.</td>
<td>typ.</td>
<td>typ.</td>
<td>typ.</td>
</tr>
<tr>
<td>P 9 [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>
<td>1:35</td>
</tr>
<tr>
<td></td>
<td>1:17</td>
<td>2:07</td>
<td>3:45</td>
<td>4:49</td>
<td>11:50</td>
<td>15:46</td>
<td></td>
</tr>
</tbody>
</table>

**Manufacturer:**
**Becton Dickinson**

<table>
<thead>
<tr>
<th>Syringe Type</th>
<th>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></td>
<td>309585</td>
<td>309603</td>
<td>305959</td>
<td>309661</td>
<td>309662</td>
<td>309653</td>
</tr>
<tr>
<td></td>
<td></td>
<td>300910</td>
<td>300911</td>
<td></td>
<td>300629</td>
<td>300863</td>
<td>300865</td>
</tr>
</tbody>
</table>

<table>
<thead>
<tr>
<th>Time to Occl. 2)</th>
<th></th>
<th></th>
<th></th>
<th></th>
<th></th>
<th></th>
<th></th>
</tr>
</thead>
<tbody>
<tr>
<td>P 1 [mm:ss]</td>
<td>typ.</td>
<td>typ.</td>
<td>typ.</td>
<td>typ.</td>
<td>typ.</td>
<td>typ.</td>
<td>typ.</td>
</tr>
<tr>
<td>P 9 [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>
<td></td>
</tr>
<tr>
<td></td>
<td>1:15</td>
<td>1:34</td>
<td>3:27</td>
<td>6:30</td>
<td>6:36</td>
<td>15:34</td>
<td></td>
</tr>
</tbody>
</table>

**vManufacturer:**
**Becton Dickinson**

<table>
<thead>
<tr>
<th>Syringe Type</th>
<th>B-D EU/USA</th>
<th>Plastipak BD 30 ml</th>
<th>BD Luer Lock 3 ml A/P</th>
<th>BD Luer Lock 5 ml A/P</th>
<th>BD Luer Lock 10 ml A/P</th>
<th>BD Luer Lock 20 ml A/P</th>
<th>BD Luer Lock 50 ml A/P</th>
</tr>
</thead>
<tbody>
<tr>
<td>Mat. No. 1)</td>
<td></td>
<td>301229</td>
<td>302113</td>
<td>302135</td>
<td>300149</td>
<td>300141</td>
<td>300144</td>
</tr>
</tbody>
</table>

<table>
<thead>
<tr>
<th>Time to Occl. 2)</th>
<th></th>
<th></th>
<th></th>
<th></th>
<th></th>
<th></th>
<th></th>
</tr>
</thead>
<tbody>
<tr>
<td>P 1 [mm:ss]</td>
<td>1:25</td>
<td>0:24</td>
<td>0:28</td>
<td>0:50</td>
<td>1:11</td>
<td>3:17</td>
<td></td>
</tr>
<tr>
<td>P 9 [mm:ss]</td>
<td>8:50</td>
<td>1:04</td>
<td>1:22</td>
<td>2:36</td>
<td>5:03</td>
<td>16:36</td>
<td></td>
</tr>
</tbody>
</table>
Chapter 8

**COMPATIBLE SYRINGES**

### Manufacturer: BD Precise

<table>
<thead>
<tr>
<th>Syringe Type</th>
<th>3 ml</th>
<th>5 ml</th>
<th>10 ml</th>
<th>20 ml</th>
<th>30 ml</th>
<th>50 ml</th>
</tr>
</thead>
<tbody>
<tr>
<td>BD Precise</td>
<td>SS+03L1</td>
<td>SS+05L1</td>
<td>SS*10LE1</td>
<td>SS+20L1</td>
<td>SS+30L1</td>
<td>SS+50L1</td>
</tr>
<tr>
<td>Time to Occl.</td>
<td>typ.</td>
<td>typ.</td>
<td>typ.</td>
<td>typ.</td>
<td>typ.</td>
<td>typ.</td>
</tr>
<tr>
<td>P 1</td>
<td>0:43</td>
<td>0:35</td>
<td>0:55</td>
<td>2:12</td>
<td>2:25</td>
<td>3:01</td>
</tr>
<tr>
<td>P 9</td>
<td>1:17</td>
<td>1:16</td>
<td>4:48</td>
<td>7:53</td>
<td>8:18</td>
<td>16:55</td>
</tr>
</tbody>
</table>

### Manufacturer: Codan

<table>
<thead>
<tr>
<th>Syringe Type</th>
<th>2 ml</th>
<th>5 ml</th>
<th>10 ml</th>
<th>20 ml</th>
<th>30/35 ml</th>
<th>50/60 ml</th>
</tr>
</thead>
<tbody>
<tr>
<td>Time to Occl.</td>
<td>typ.</td>
<td>typ.</td>
<td>typ.</td>
<td>typ.</td>
<td>typ.</td>
<td>typ.</td>
</tr>
<tr>
<td>P 1</td>
<td>0:07</td>
<td>0:09</td>
<td>0:19</td>
<td>0:36</td>
<td>0:45</td>
<td>1:48</td>
</tr>
<tr>
<td>P 9</td>
<td>0:58</td>
<td>1:18</td>
<td>2:23</td>
<td>4:14</td>
<td>4:22</td>
<td>11:41</td>
</tr>
</tbody>
</table>

### Manufacturer: Fresenius

<table>
<thead>
<tr>
<th>Syringe Type</th>
<th>Injectomat 50 ml</th>
</tr>
</thead>
<tbody>
<tr>
<td>Fresenius</td>
<td>9000701</td>
</tr>
<tr>
<td>Mat. No.</td>
<td>9000701</td>
</tr>
<tr>
<td>Time to Occl.</td>
<td>typ.</td>
</tr>
<tr>
<td>P 1</td>
<td>4:37</td>
</tr>
<tr>
<td>P 9</td>
<td>21:09</td>
</tr>
</tbody>
</table>

### Manufacturer: Becton–Dickinson

<table>
<thead>
<tr>
<th>Syringe Type</th>
<th>BD Precise 50 ml A/P</th>
<th>BD Precise 20 ml A/P</th>
</tr>
</thead>
<tbody>
<tr>
<td>B–D Precise</td>
<td>300144</td>
<td>300141</td>
</tr>
<tr>
<td>Mat. No.</td>
<td>300144</td>
<td>300141</td>
</tr>
<tr>
<td>Time to Occl.</td>
<td>03:17</td>
<td>01:11</td>
</tr>
<tr>
<td>P 1</td>
<td>16:36</td>
<td>05:03</td>
</tr>
</tbody>
</table>
Syringes not specified in IEC/EN 60601–2–24

Nutrition pumps, in contrast to infusion pumps, are not classified as Class IIb according to the infusion pump norm IEC/EN 60601–2–24. There are therefore no direct guidelines concerning the technical characteristics (accuracy of infusion rate, alarm parameters etc) of the relevant disposables.

The syringes types listed in the following tables can be used with the Perfusor® Space. However, due to the relatively high tolerances allowed in the disposables, the accuracy of infusion rate and the alarm parameters are not comparable with infusion syringes.

Precautions must be taken as follows:

- Consider the risks involved in the use of a non LuerLock connection prior to the therapy
- Permanently observe the connection between the syringe and the infusion set, in order to detect any break in the connection
- If syringe and tubing line are not fixed, a disconnection can occur possibly leading to air infusion, reverse infusion, under- /over delivery and/or nonsterility.

<table>
<thead>
<tr>
<th>Manufacturer</th>
<th>Sizes (ml)</th>
</tr>
</thead>
<tbody>
<tr>
<td>Polfa</td>
<td>60, 20, 10, 5, 3</td>
</tr>
<tr>
<td>NeoMed</td>
<td>60, 35, 20, 12, 6, 3</td>
</tr>
<tr>
<td>Vygon</td>
<td>Nutrisafe: 60, 35, 20, 10, 5  C-Gon: 60, 20, 5</td>
</tr>
</tbody>
</table>

Complete list available on request.
The graphs show the accuracy/uniformity of flow in relation to time. They allow for the following:

The delivery behaviour or delivery precision is essentially influenced by the type of (disposable syringe) used. Deviations from the technical data of the pump cannot be guaranteed as the manufacturer may change syringe specification significant to system accuracy without prior notification.

System accuracy is ± 2% typical by volume as measured using the trumpet curve test method defined in IEC 60601-2-24 at rates of 1ml/h (23°C) and when the pump is used with recommended syringes.
Trumpet Curves
Measured values for second hour in each case.
Measurement interval $\Delta t = 0.5$ min
Observation interval $p \times \Delta t [\text{min}]$ 

Start Up Graphs
Measurement interval $\Delta t = 0.5$ min
Measurement duration $T = 120$ min
Flow $Q_i$ (ml/h) 

Advice on choosing the right syringe size
The device is to be used with the smallest possible, clinically acceptable, syringe. This is particularly important when infusing critical medication with a short biological half-life at a low rate.

Using large syringes for very slow infusions can affect the technical characteristics of the device and lead to rate fluctuations, lengthened start-up times and delayed detection of occlusions.

Recommendation

<table>
<thead>
<tr>
<th>Syringe size</th>
<th>50/60 ml</th>
<th>30 ml</th>
<th>20 ml</th>
<th>10 ml</th>
<th>5 ml</th>
<th>3 ml</th>
</tr>
</thead>
<tbody>
<tr>
<td>Recommended minimum rate*</td>
<td>1 ml/h</td>
<td>1 ml/h</td>
<td>0.5 ml/h</td>
<td>0.2 ml/h</td>
<td>0.1 ml/h</td>
<td>0.1 ml/h</td>
</tr>
</tbody>
</table>

* The adjustment of smaller infusion rates is possible in general. However it may influence the start-up characteristic and accuracy in respect to the trumpet curve.

Advice on start-up
Every syringe has certain tolerances in its start-up characteristics (dependant on manufacturer, bung material, siliconisation of the cylinder etc). To minimise the start-up delay, we recommend choosing an adequately sized syringe and manipulating the plunger back and forth before inserting it. This ensures the stick-slip effect of the rubber bung is reduced.

To further reduce start-up delays, the device is fitted with a start-up acceleration function.
## TECHNICAL DATA

<table>
<thead>
<tr>
<th>Type of unit</th>
<th>Infusion Syringe Pump</th>
</tr>
</thead>
<tbody>
<tr>
<td>Classification</td>
<td>💪 Defibrillator–proof; CF equipment</td>
</tr>
<tr>
<td></td>
<td>☐ Protective Class II; Protective Class I in combination with SpaceStation</td>
</tr>
<tr>
<td>Class (acc. to Directive 93/42 EEC)</td>
<td>IIb</td>
</tr>
<tr>
<td>Moisture protection</td>
<td>IP 22 (fluid protected for horizontal usage)</td>
</tr>
<tr>
<td>External power supply:</td>
<td></td>
</tr>
<tr>
<td>■ Rated voltage</td>
<td>Via B. Braun SpaceStation or optional mains adaptor (rated voltage 100 ... 240 V AC~ 50/60 Hz) for stand alone operation</td>
</tr>
<tr>
<td>■ External low voltage</td>
<td>11 ... 16 V DC — via Connection Lead SP 12 V or via SpaceStation</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>
</tbody>
</table>

### Operating conditions:

- Relative humidity: 30 % ... 90 % (without condensation)
- Temperature: +5 ... +40 °C
- Atmospheric pressure: 500 ... 1060 mbar

### Storage conditions:

- Relative humidity: 20 % ... 90 % (without condensation)
- Temperature: -20 ... +55 °C
- Atmospheric pressure: 500 ... 1060 mbar

<table>
<thead>
<tr>
<th>Type of battery pack (rechargeable)</th>
<th>Li-Ion</th>
<th>NiMHH</th>
</tr>
</thead>
<tbody>
<tr>
<td>Operating time of rechargeable battery</td>
<td>Li-Ion</td>
<td>Wireless active Perfusor® at 5ml/h typ. 3 hours</td>
</tr>
<tr>
<td></td>
<td></td>
<td>Wireless active Perfusor® at 25ml/h typ. 2.5 hours</td>
</tr>
<tr>
<td></td>
<td></td>
<td>Wireless inactive Perfusor® at 5ml/h typ. 17 hours</td>
</tr>
<tr>
<td></td>
<td></td>
<td>Wireless inactive Perfusor® at 25ml/h typ. 10 hours</td>
</tr>
<tr>
<td></td>
<td></td>
<td>NiMHH at 5ml/h typ. 19 hours</td>
</tr>
<tr>
<td></td>
<td></td>
<td>at 25ml/h typ. 10 hours</td>
</tr>
<tr>
<td>Recharging time</td>
<td>Approx. 6 hours</td>
<td></td>
</tr>
<tr>
<td>Weight</td>
<td>Approx. 1.4 kg</td>
<td></td>
</tr>
<tr>
<td>Dimensions (W x H x D)</td>
<td>249 x 68 x 152 mm</td>
<td></td>
</tr>
</tbody>
</table>
**Volume preselection**

0.1 - 99.99 ml in increments of 0.01 ml
100.0 - 999.0 ml in increments 0.1 ml
1000 - 9999 ml in increments 1 ml

**Time preselection**

00:01 – 99:59 h

**Accuracy of set delivery rate**

± 2 % according to IEC/EN 60601-2-24

**Occlusion alarm pressure**

9 levels up to 1.2 bar

**Max. Volume in case of single fault condition**

For incorrect dosages of 0.1 ml due to malfunctions of the device the pump will automatically shut off.

**Technical inspection (safety check)**

Every 2 years

**Multiple lines connected to one patient port**

Connecting multiple infusion lines with different flow rates may affect the rate for all infusions past the point of connection.

**Selectable delivery rates**

Continuous infusion rate range / bolus rates in dependence on syringe sizes:

<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>30/35</td>
<td>0.01 - 999.9</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>

**Rate increments**

0.01 - 99.99 ml/h in increments of 0.01 ml/h
100.0 - 999.9 ml/h in increments of 0.1 ml/h

**Accuracy of bolus infusion**

typ. ± 2 %

**Max. bolus after bolus reduction**

≤ 0.2 ml

**KVO-rate**

Delivery rate > 10 ml/h: KVO-rate 3 ml/h
Delivery rate < 10 ml/h: KVO-rate 1 ml/h
Delivery rate < 1 ml/h: KVO-rate = set rate (default setting 0.1 ml/h)

**Computer connection**

USB connection in combination with B. Braun interface lead CAN SP (8713230) including electrical insulation. Please pay attention to safety notices.
| History protocol | < 3000 last history entries.  
| | 100 events for system diagnose.  
| | Refer to separate documents of the History Viewer for closer information.  
| Alarm volume | 9 levels from 1 (59dBA) to 9 (74dBA)  

- Use only pressure proof and compatible disposable items (min. 2 bar/1500 mm Hg) to avoid influencing performance data - which would result in impairing patient safety.
- Only use combined with approved devices/accessories by the manufacturer, otherwise this may lead to higher emission or reduced immunity.
- Use only compatible combinations of equipment, accessories, working parts and disposables with luer lock connectors.

**Essential Performance for Infusion pumps:**
- Infusion of liquids without variation of infusion rate
- Pressure limitation as protection from the bursting of the infusion line
- Protection from air-infusion
- Protection against unintended bolus volumes and occlusion (added by IEC 60601-2-24)
- Alarm signal of high priority (added by IEC 60601-2-24)
EMC (ELECTROMAGNETIC COMPATIBILITY)

Guidance and manufacturer’s declaration on electromagnetic compatibility

<table>
<thead>
<tr>
<th>Emissions test</th>
<th>Compliance</th>
<th>Electromagnetic environment guidance</th>
</tr>
</thead>
<tbody>
<tr>
<td>RF emissions</td>
<td>Group 1</td>
<td>The Space System uses RF energy only for its internal function. Therefore, its RF emissions are very low and are not likely to cause any interference in nearby electronic equipment. If WLAN-Module is installed within Battery module (8713182A) or WLAN USB Stick for SpaceCom (8713185) is used RF energy is transmitted by the Space System. Refer to technical data of Battery-Pack SP with Wifi IUf and/or SpaceStation and SpaceCom for details.</td>
</tr>
<tr>
<td>CISPR 11</td>
<td></td>
<td>The Space System or any component 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.</td>
</tr>
<tr>
<td>Harmonic emissions</td>
<td>Class B [Note 2]</td>
<td>Note 1: Maximum emissions are measured with a complete system (SpaceStation and components). Note 2: If Class A equipment is attached to the Space System, the Space System will become Class A too. This equipment/system may cause radio interference or may disrupt the operation of nearby equipment. It may be necessary to take mitigation measures, such as re-orienting or relocating the Space System or shielding the location.</td>
</tr>
<tr>
<td>IEC 61000-3-2</td>
<td></td>
<td></td>
</tr>
<tr>
<td>Voltage fluctuations /</td>
<td>Complies</td>
<td></td>
</tr>
<tr>
<td>flicker emissions</td>
<td></td>
<td></td>
</tr>
<tr>
<td>IEC 61000-3-3</td>
<td></td>
<td></td>
</tr>
</tbody>
</table>
### Guidance and manufacturer’s declaration – electromagnetic immunity

The Space System is intended for use in the electromagnetic environment specified below. The customer or the user of the Space System or any component should assure that it is used in such an environment.

<table>
<thead>
<tr>
<th>Immunity test</th>
<th>test level</th>
<th>Compliance level</th>
<th>Electromagnetic environment – guidance</th>
</tr>
</thead>
</table>
| Electrostatic discharge (ESD) according IEC 60601-4-2 | contact IEC 60601-1-2: ±6KV IEC 60601-2-24: ±8KV air IEC 60601-1-2: ±8KV IEC 60601-2-24: ±15KV | ±6KV no disturbances ±8KV stop with alarm possible ±8KV no disturbances ±15KV stop with alarm possible | Floors should be wood, concrete or ceramic tile. If floors are covered with synthetic material, the relative humidity should be at least 30%.
| Electrostatic transient / burst according IEC 61000-4-4 | ± 2 kV for power supply lines ± 1 kV for input/output lines | ±2KV ±1KV | A/C power quality should be that of a typical commercial or hospital environment.
| Surge according IEC 61000-4-5                     | differential mode ± 1KV common mode ±2KV | ±1KV ±2KV | A/C power quality should be that of a typical commercial or hospital environment.
| Voltage dips, short interruptions and voltage variations on power supply input lines according IEC 61000-4-11 | < 5 % UT (>95 % dip in UT ) for 0,5 cycle 40 % UT (60 % dip in UT ) for 5 cycles 70 % UT (30 % dip in UT ) for 25 cycles < 5 % UT (>95 % dip in UT ) for 5 sec <5% UT for 5 s (>95% dip) | complies by use of internal battery | A/C power quality should be that of a typical commercial or hospital environment. If the user of the Space System requires continued operation during long time A/C power interruptions, it is recommended that the Space System or component be powered from an uninterruptible power supply or a battery.
| Power frequency (50/60 Hz) magnetic field according IEC 61000-4-8 | 3 A/m | 400 A/m | Power frequency magnetic fields should be at levels characteristic of a typical location in a typical commercial or hospital environment.

**Note:** Different test values of IEC 60601-2-24 are marked in the table. At the test values no dangerous disturbances occurred at the lower test values of IEC 60601-1-2.
## Guidance and manufacturer’s declaration – electromagnetic immunity

The Space System is intended for use in the electromagnetic environment specified below. The customer or the user of the Space System or any component should assure that it is used in such an environment.

<table>
<thead>
<tr>
<th>Immunity test</th>
<th>test level IEC 60601-1-2</th>
<th>Compliance level</th>
<th>Electromagnetic environment – guidance</th>
</tr>
</thead>
<tbody>
<tr>
<td>Conducted electromagnetic RF fields according IEC 61000-4-6</td>
<td>IEC 60601-1-2: 3 V eff normal and 10 V eff in ISM frequency band</td>
<td>10 V eff 150 KHz to 800 MHz</td>
<td>Portable and mobile RF communications equipment should be used no closer to any part of the Space System or its components, including cables, than the recommended separation distance calculated from the equation applicable to the frequency of the transmitter.</td>
</tr>
<tr>
<td>Radiated electromagnetic RF fields according IEC 61000-4-3</td>
<td>IEC 60601-2-24: 10 V eff 150 KHz to 800 MHz</td>
<td>10 V/m 80 MHz to 2.5 GHz</td>
<td><strong>Recommended separation distance</strong></td>
</tr>
<tr>
<td></td>
<td></td>
<td></td>
<td>d = 1.2 \sqrt{P}</td>
</tr>
<tr>
<td></td>
<td></td>
<td></td>
<td>Field strengths should be less than 10 V/m</td>
</tr>
<tr>
<td></td>
<td></td>
<td></td>
<td>d = 1.2 \sqrt{P}</td>
</tr>
<tr>
<td></td>
<td></td>
<td></td>
<td>80 MHz to 800 MHz</td>
</tr>
<tr>
<td></td>
<td></td>
<td></td>
<td>d = 2.3 \sqrt{P}</td>
</tr>
<tr>
<td></td>
<td></td>
<td></td>
<td>800 MHz to 2.5 GHz</td>
</tr>
<tr>
<td></td>
<td></td>
<td></td>
<td>where p is the maximum output power rating of the transmitter in watts (W) according to the transmit-ter manufacturer and d is the recom-mended separation distance in meters (m).</td>
</tr>
<tr>
<td></td>
<td></td>
<td></td>
<td>Field strengths from fixed RF transmitters, as determined by an electromagnetic site survey, should be less than the compliance level in each frequency range.</td>
</tr>
<tr>
<td></td>
<td></td>
<td></td>
<td>Interference may occur in the vicinity of equipment marked with the following symbol:</td>
</tr>
</tbody>
</table>

### Notes:

**NOTE 1:** At 80 MHz and 800 MHz, the higher frequency range applies.

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

**NOTE 3:** See next page.
NOTE 3: Different test values of IEC 60601-2-24 are marked in the table. At these test values no dangerous disturbances are allowed while at the lower test values of IEC 60601-1-2. Field strengths from fixed transmitters, such as base stations for radio (cellular/cordless) telephones and land mobile radios, amateur radio, AM and FM radio broadcast and TV broadcast cannot be predicted theoretically with accuracy. To assess the electromagnetic environment due to fixed RF transmitters, an electromagnetic site survey should be considered. If the measured field strength in the location in which the SpaceSystem is used exceeds the applicable RF compliance level above, the Space System should be observed to verify normal operation. If abnormal performance is observed, additional measures may be necessary, such as re-orienting or relocating the Space System.

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

| rated power of the ratio transmitter \[W\] | Separation distance according to frequency of transmitter \[m\] |
|-----------------|-----------------|-----------------|
|                 | 150 kHz bis 80 MHz | 80 MHz bis 800 MHz | 800 MHz bis 2,5 GHz |
| 0,01            | 0,12             | 0,12             | 0,23             |
| 0,1             | 0,38             | 0,38             | 0,73             |
| 1               | 1,2              | 1,2              | 2,3              |
| 10              | 3,8              | 3,8              | 7,27             |
| 100             | 12               | 12               | 23               |

NOTE 1: For transmitters rated at a maximum power output not listed above, the recommended separation distance \(d\) in meters \(m\) can be determined using the equation applicable to the frequency of the transmitter, where \(P\) is the maximum output power rating of the transmitter in watts \(W\) according to the transmitter manufacturer.

NOTE 2: An additional factor of \(10/3\) is used in calculating the recommended separation distance for transmitters in the frequency range 0.15 MHz to 2.5 GHz to decrease the likelihood that mobile/portable communications equipment could cause interference if it is inadvertently brought into patient areas.

NOTE 3: These guidelines may not apply in all situations. Electromagnetic propagation is affected by absorption and reflection from structures, objects and people.
Responsibility of the Manufacturer

The manufacturer, assembler, installer or importer is responsible for the effects on safety, reliability and performance of the equipment only if:

- assembly operations, extensions, re-adjustments, modifications or repairs are carried out by authorized personnel,
- the electrical installation of the relevant room complies with the appropriate requirements (e.g. VDE 0100, 0107 and/or the IEC-publications resp. national requirements),
- the equipment is used in accordance with the Instructions for Use and
- the Technical Safety Checks are carried out regularly.

Warranty

B. Braun provides 24 months warranty, as from the date of delivery, for every Perfusor® Space (12 months for every Battery-Pack SP). This covers repair or replacement of parts damaged as a result of design/manufacturing errors or material defects. Modifications or repairs to the unit undertaken by the user/operator or by third parties invalidate the warranty.

The warranty does not cover the following:
Elimination of faults attributable to incorrect/unauthorized handling, or to normal wear and tear.
Defective rechargeable battery packs can be returned to B. Braun for further disposal.

WARNING: Do not modify this equipment without authorization of the manufacturer.

Labeling of electric and electronic devices according to directive 2012/19/EU (WEEE).

Training

B. Braun offers a training. Please ask your local representative for further details.

Technical Safety Check* / Service

The Technical Safety Check is recommended to be carried out every 2 years and should be documented. Servicing work must be carried out exclusively by trained personnel.
Check regularly

Check for cleanliness, completeness and damage. Use only according to Instructions for Use. During an exchange interval of the disposable the pump has to perform a self-test. Check the following items each time the pump is switched on: self-check, audible alarm, process- and alarm control indication.

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 all exposed surfaces using a clean, soft, lint-free cloth dampened with a mild cleaning solution of warm, soapy water. Make sure to remove any visible residue from all surfaces prior to disinfecting. Do not spray disinfectants directly on the pump, use a soft, low lint cloth dampened but not saturated with product. After cleaning and disinfecting allow device to dry for at least 20 minutes prior to use. Wipe magnifying- and displayglas on front of pump door only with a soft cloth.

Note: Keep instrument upright and do not allow any part of instrument to become saturated with or submersed 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.

Substances of the listed groups are approved for cleaning and regular disinfection routines of surfaces according to recommendation of disinfectant manufacturer.

<table>
<thead>
<tr>
<th>Group</th>
<th>Active Substance</th>
</tr>
</thead>
<tbody>
<tr>
<td>Quaternary ammonium compounds</td>
<td>DDAC (Didecyldimethylammoniumchlorid)</td>
</tr>
<tr>
<td></td>
<td>BAC (Benzalkoniumchlorid)</td>
</tr>
<tr>
<td>Aldehydes</td>
<td>Glutaral</td>
</tr>
<tr>
<td></td>
<td>Glyoxal</td>
</tr>
<tr>
<td>Peroxides</td>
<td>Hydrogen Peroxide</td>
</tr>
<tr>
<td>Active chlorine</td>
<td>Sodium Hypochlorite</td>
</tr>
<tr>
<td>Acid</td>
<td>Citric Acid</td>
</tr>
</tbody>
</table>

**Note:** Do not use Hexaquart® or other alkylamine containing disinfectants. Recommended: disinfectant for wiping available from B. Braun: Meliseptol® Foam pure, Melsitt 10% and Melsept SF 10%.

**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**

The pumps as well as battery packs can be returned to B. Braun for further disposal. When taking care of disposing of disposables as well as infusion solutions, please consider the applicable hygiene and disposal regulations.

**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. Contact the service department. Testing the proper function of the device should be performed before initial use. This is even ruled by law in several countries. A respective form can be obtained from B. Braun.

**Included in Delivery**

Perfusor® Space, Battery-Pack SP, Instructions for Use-Set.
INSTRUCTIONS FOR USE ACCESSORY

SpaceStation (8713140)

Station for up to four pumps. For further information see Instructions for Use of SpaceStation.

SpaceCover Standard (8713147)
SpaceCover Comfort (8713145)

Cover to be placed on upper SpaceStation incl. built-in handle. The SpaceCover Comfort additionally includes a central alarm management and alarm LEDs.

PoleClamp SP (8713130)

A maximum of three B. Braun Space pumps and one SpaceControl can be stacked together when used with the PoleClamp SP. For detailed instructions on secure fixation of the PoleClamp SP please refer to "Overview Perfusor® Space" and "Patient Safety".

Power Supply SP III (8713110D - 8713123D)

The Power Supply SP is adequate to supply power for a single pump and one SpaceControl.

1.) Connect plug of Power Supply SP with socket P2 on back of pump (ensure that plug "clicks").
2.) Push power plug into wall outlet.

Note: For disconnection from pump, press lever on plug down.
A maximum of three plugs can be stacked upon each other in socket P2.

Prior to use, visibly inspect the power supply and if reject if damaged.

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

Combi Lead SP 12 V (8713133)

The Combi Lead SP can connect up to three pumps. All pumps can then be operated via the Connection Lead SP (12 V).

1.) Connect plug of the Combi Lead SP 12 V with the socket P2 on the back of the pump.
2.) Connect plug of Connection Lead SP with Combi Lead SP.
3.) Push plug of Connection Lead SP into 12 V connector.

Note: A maximum of three plugs can be stacked upon each other in socket P2.

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

For further information on the Battery-Pack SP (NiMH) see "Battery Operation".

Battery-Pack SP (Lilon) incl. Pin and WiFi (8713182A)

For further information see Instructions for Use of "Battery Pack SP with WiFi".

Interface Lead CAN SP (8713230)

Interface Lead CAN SP is needed in order to set up a connection between the SpaceStation/pump and the computer outlet (for service requirements).

1.) Push plug into socket F3 on the SpaceStation or P2 on the pump and connect with the CAN/USB converter.

2.) Connect CAN/USB converter to computer outlet as described in the Instructions for Use manual.

Caution: The Interface Lead CAN SP is only to be used by the service department; never use while patient is connected.

Note: A maximum of three plugs can be stacked upon each other in socket P2.

Interface Lead RS232 SP (8713234)

Interface Lead RS232 SP is needed in order to set up a connection between the Space pump and the computer outlet (for service requirements).

1.) Push plug into socket P2 on the pump and connect with the Interface Lead RS232 SP.

2.) Connect Interface Lead RS232 SP to computer outlet as described in the Instructions for Use manual.

Caution: The Interface Lead RS232 SP is only to be used by the service department; never use while patient is connected.

Note: A maximum of three plugs can be stacked upon each other in socket P2.
Connection Lead SP (12 V) (8713231)

Install the Connection Lead SP (12 V) in the following way:

1.) Connect plug to socket P2 on back of pump or F3 on SpaceStation respectively.
2.) Put the connection lead into the car socket.
3.) If necessary, remove red adaptor of motor vehicle connector by slightly turning and simultaneously pulling.

The green LED of the electronic box shows the operating voltage. The mains connector can easily be replaced by another plug if required.

**Caution:** Do not connect the pump to a patient during external car battery charging!

**Note:** A maximum of three plugs can be stacked upon each other in socket P2.

Connection Lead for Staff Call SP (8713232)

To connect the Perfusor® Space to staff call, use the Connection Lead for Staff Call SP. The staff call needs to comply with the requirements of VDE 0834 (consider country specific regulations).

**Note:** Test staff call signalling before every use.

The Perfusor® Space offers three different staff call operating modes. They are displayed in the signalling scheme. Consider the staff call of the hospital when choosing an operating mode. Choose the operating mode via the service program.

<table>
<thead>
<tr>
<th>Mode</th>
<th>Alarm</th>
<th>Operation</th>
</tr>
</thead>
<tbody>
<tr>
<td>static without Off-Alarm</td>
<td><img src="image1.png" alt="Image" /></td>
<td><img src="image2.png" alt="Image" /></td>
</tr>
<tr>
<td>dynamic without Off-Alarm</td>
<td><img src="image3.png" alt="Image" /></td>
<td><img src="image4.png" alt="Image" /></td>
</tr>
<tr>
<td>dynamic with Off-Alarm</td>
<td><img src="image5.png" alt="Image" /></td>
<td><img src="image6.png" alt="Image" /></td>
</tr>
</tbody>
</table>

*) in the mode static without Off-Alarm, the staff call can be suppressed with OK
Caution: The user should always closely observe the local pump alarms as well.

Note: A maximum of three plugs can be stacked upon each other in socket P2.

Technical Data

<table>
<thead>
<tr>
<th></th>
<th>Connecting Wire</th>
</tr>
</thead>
<tbody>
<tr>
<td>Alarm</td>
<td>disconnected</td>
</tr>
<tr>
<td>Operation</td>
<td>connected</td>
</tr>
</tbody>
</table>

Polarity of connexion is arbitrary:

max. 24 V / 0.5 A / 12 VA
PCA-Accessories

- Space PCA-Kit (REF 8713554) consisting of:
  - Demand button
  - Hook and loop tape for fixation of the demand button at the patient’s arm
  - Line fixation connection between hook and loop tape and demand button
  - Metal clip alternatively for fixation at the bed sheet
  - Cable strap for wrapping the cable of the demand button
  - PCA-Key for locking the syringe holder or the Syringe Anti Removal Cap

- Syringe Anti Removal Cap PSP (REF 8713556)

Fixation of the demand button:
- at the wrist: or at the bed sheet:

Usage of the cable strap:

Usage of the Syringe Anti Removal Cap PSP:
The Syringe Anti Removal Cap PSP is slided over the drive head from the front and is fixed with the PCA-key (270° clockwise rotation). Mind the markings – make sure it is securely locked. Dismantling: counter clockwise rotation of 270°. Push to the left and disengage.

Caution: When Syringe Anti Removal Cap is used always change the syringe as soon the “syringe empty” alarm appears.
B. Braun Perfusor® Space (100 – 240 V) .................................................. 8713030

Recommended accessories for the B. Braun Perfusor® Space:

SpaceStation .................................................................................. 8713140
SpaceCover Standard ................................................................. 8713147
SpaceCover Comfort ................................................................. 8713145
PoleClamp SP .......................................................... 8713130
Power Supply SP EU III .................................................. 8713110D
Power Supply SP EU III 3.0m .................................................. 8713123D
Power Supply SP GB III .............................................. 8713111D
Power Supply SP US III .................................................. 8713112D
Power Supply SP AU III .................................................. 8713113D
Power Supply SP-RSA plug III ........................................ 8713115D
Power supply SP CN III .................................................. 8713117D
Power Supply SP DK III .................................................. 8713118D
Power Supply SP BR III .................................................. 8713119D
Power Supply SP KR III .................................................. 8713120D
Combi Lead SP 12 V .......................................................... 8713133
Battery-Pack SP (NiMH) .................................................. 8713180
Battery-Pack SP (NiMH) incl. Pin ........................................ 8713180A
Battery-Pack SP (LiIon) incl. Pin and WiFi ...................... 8713182A
Interface Lead CAN SP .................................................. 8713230
Connection Lead SP (12 V) ............................................... 8713231
Connection Lead for Staff Call SP .................................... 8713232
Interface Lead RS232 SP .................................................. 8713234
Space PCA Kit .............................................................. 8713554
Syringe Anti Removal Cap PSP ........................................ 8713556
PCA Lockbox ................................................................. 8713557

Original Perfusor® Syringes:

20ml, without needle .................................................. 8728615
20ml, with aspiration needle ................................. 8728623
50ml, without needle .................................................. 8728844F
50 ml, without needle .................................................. 8728844F-06
50 ml, with aspiration needle ........................................ 8728810F-06
50 ml, with aspiration needle and 15 my particle filter .... 8728852F-06
50 ml, with aspiration needle, EPLANESTH (yellow cylinder) .... 8728801F-06
50ml, with aspiration needle ...............................................................8728810F
50ml, with aspiration needle and 15 µm particle filter ......................8728852F
50ml, light protection orange, aspiration needle and 15 µm particle filter .......................................................................................8728861F-06
50ml, yellow inked cylinder and aspiration needle .........................8728801F

**Omnifix® syringes**

Omnifix® 50/60 ml Luer Lock.................................................................4617509F
Omnifix® 30 ml Luer Lock....................................................................4617304F
Omnifix® 20 ml Luer Lock.................................................................4617207V
Omnifix® 10 ml Luer Lock.................................................................4617100V
Omnifix® 5 ml Luer Lock ....................................................................4617053V
Omnifix® 3 ml Luer Lock ....................................................................4617022V
Omnifix® 2 ml Luer Lock ....................................................................4617029V
Omnifix® 50 ml Luer Lock, UV-protect .................................................4617510F-06

**Original Perfusor lines made of PE (polyethylen). Tube diameter 0.9 mm**

50 cm, Luer Lock.................................................................8255059
100 cm, Luer Lock ....................................................................8255067
150 cm, Luer Lock ....................................................................8722935
200 cm, Luer Lock ....................................................................8723060
250 cm, Luer Lock ....................................................................8272565
300 cm, Luer Lock ....................................................................8250146

**Original Perfusor lines made of PVC. Tube diameter 1.5 mm**

50 cm, Luer Lock ....................................................................8255172
150 cm, Luer Lock ....................................................................8722960
200 cm, Luer Lock ....................................................................8722862
250 cm, Luer Lock ....................................................................8255490
300 cm, Luer Lock ....................................................................8255253

**Original Perfusor lines made of PVC. Tube diameter 0.9 mm**

75 cm, Luer Lock ....................................................................8722870
150 cm, Luer Lock ....................................................................8255504
Specific Original Perfusor lines
Transparent UV - light protected, 150 cm,
PE - tube 0.9 mm, Luer Lock ..............................................................8723017
Black PE tube 0.9 mm, 150 cm, Luer Lock.......................................8723010
Black PVC tube 1,5 mm, 150 cm, Luer Lock.................................8722919

Type PCA, PVC tube 1,5 mm, 150 cm, Luer Lock..........................8726019
with 0.2 µm Sterifix filter, PVC tube 1,5 mm,
200cm, Luer Lock .................................................................8723001
with SafeSite valve, PVC tube 1,5 mm,
150 cm, Luer Lock.................................................................8722820