

Instructions for Use and Technical Description



Multicare

Positionable bed for intensive care with scales and without scales



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1 Symbols and Definitions

1.1 Warning Notices

1.1.1 Types of Warning Notices

Warning notices are differentiated by the type of danger using the following key words:

- ► CAUTION warns about the risk of material damage.
- ► WARNING warns about the risk of physical injury.
- ► DANGER warns about the risk of fatal injury.

1.1.2 Structure of Warning Notices



SIGNAL WORDS! Type and source of danger! ► Measures to avoid the danger.

1.2 Instructions

Structure of instructions:

► Perform this step. Results, if necessary.

1.3 Lists

Structure of bulleted lists:

- List level 1
 - □ List level 2
 - List level 3



1.4 Symbols on the Package

| | FRAGILE, HANDLE WITH CARE | |
|-----|----------------------------------|--|
| | THIS WAY UP | |
| | KEEP DRY (PROTECT FROM HUMIDITY) | |
| PAP | PAPER RECYCLING SYMBOL | |
| × | DO NOT USE HAND TRUCK HERE | |
| | DO NOT STACK DURING STORAGE | |



1.5 Symbols and Labels on the Bed

| | READ INSTRUCTIONS FOR USE | |
|--|---|--|
| | THERMAL PROTECTION FOR TRANSFORMER | |
| | WARNING | |
| | ONLY SUITABLE FOR INDOOR USE | |
| Ŕ | APPLIED PARTS TYPE B | |
| $\textcircled{\begin{tabular}{c} \hline \hline$ | SAFETY ISOLATING TRANSFORMER, GENERAL | |
| E212434 | MET LABORATORIES SAFETY MARK | |
| CE | CE MARK (OPTICARE, SYMBIOSO) | |
| | JACK FOR ATTACHMENT OF CONDUCTOR FOR POTENTIAL EQUALISATION | |
| <u></u> = kg | SAFE WORKING LOAD | |
| | WARNING AGAINST CRUSHING OR TRAPPING | |
| | USE MATTRESS RECOMMENDED BY MANUFACTURER | |



| | DO NOT PUT ANY OBJECTS ON UNDERCARRIAGE | |
|---|---|--|
| <u>o⊡</u> ⊒ = kg | MAXIMUM WEIGHT OF PATIENT | |
| 📥 = kg | WEIGHT OF BED | |
| + + + + + + + + + + + + + + + + + + + | DESIGNATION OF HOSPITAL BED FOR ADULTS | |
| Santiged | ANTIBACTERIAL SURFACE FINISH | |
| | WEEE SYMBOL (RECYCLE AS ELECTRONIC WASTE, DO NOT PUT INTO THE HOUSEHOLD WASTE) | |
| EAC | EAC MARKING (EURASIAN CONFORMITY) | |
| | DO NOT POLLUTE THE ENVIRONMENT | |
| S TUCK SUD Protection protocols Select wheter | TÜV MARK | |
| | GO BUTTON (PRESS TO ACTIVATE CONTROL ELEMENT) | |
| бтор | STOP BUTTON (PRESS TO INTERRUPT BED POSITIONING) | |





1.6 Serial Label with UDI

Pictures of serial labels below serve just for explanation of the signs and fields on the serial labels.



Fig. Serial Label with UDI (Multicare with scales)

| 1 | Address of Manufacturer |
|----|--|
| 2 | Manufacturing Date (Year-Month-Day) |
| 3 | DI (Device Identifier) / GTIN (Global Trade Item Number) |
| 4 | 1D Bar code GS1-128 (Serial Number) |
| 5 | Symbols |
| 6 | Configuration number |
| 7 | Electrical Specification |
| 8 | Serial Number |
| 9 | PI (Product Identifier) |
| 10 | 2D Bar Code (GS1 DataMatrix) DI+PI=UDI |

| ⊴⊾⊴ | | |
|------------|---------------------|----------------------------|
| Туре: | WS 17 | 10°C / +40°C |
| Max Min | 250,0 kg 10,0 kg | e = 0,5 kg T = -249,5kg |

 Scales Abbreviations

 Max
 maximum capacity of the weighing instrument

 Min
 minimum capacity of the weighing instrument

 e
 verification scale interval

 T
 tare value

Fig. Serial Label (WS17)



| THIS PRODUCT IS PROTECTED BY ONE OR MORE OF THE FOLLOWING PATENTS: | | |
|---|------------|------------|
| US7492111 | US7647659 | US8024101 |
| US10010465 | US8959680 | US10130538 |
| US9585490 | US8959680 | US10206835 |
| US9271887 | US8452508 | US10335333 |
| US9205011 | US8112836 | US8746710 |
| US10376216 | US10383780 | |

Fig. Label with applied patents

1.7 Acoustic signalisation

| SOUND | MEANING |
|--|--|
| CONTINUOUS SOUND | overheating |
| | accumulator overcurrent |
| | scales overload (only version with scales) |
| | actuator overload |
| BEEP + CONTINUOUS SOUND | Siderail Signal (lateral tilt + head siderail or foot siderail down) |
| REPEATED BEEP: 0,6s sound / 2,6s silence | STOP error (all STOP buttons are disabled) |
| MELODY: 3 beeps, pause, 2 beeps, longer pause, 3 beeps, pause, 2 beeps | Bed Exit Alarm (only version with scales) |
| BEEP lasting 0,3s | confirmation |
| | stopping or locked function |
| | lateral tilt 15° achieved |
| | transition from tilt (lateral tilt, Trendelenburg, Antitrendelenburg) to horizontal position |
| 4 TIMES REPEATED BEEP lasting 0,3s | disconnected from the mains |
| | positioning powered by the accumulator |
| BEEP lasting 0,5s | start of service mode or end of service mode |
| | keyboard error (positioning blocked) |
| BEEP lasting 3s | system error |
| BEEP lasting 5s | SCU disconnected (only if integrated mattress is used) |
| | scale module disconnected (only version with scales) |
| REPEATED BEEP during 3 minutes: 1,1s sound / 1,1s silen- ce | Brake Signal (only version with Brake Signal) |



1.8 Visual signalisation

1.8.1 Mains Power LED (Multiboard, Attendant Control Panel)

| MAINS POWER LED | MEANING |
|-----------------------------------|--|
| lit | connected to the mains |
| flashing: 0,6s lit / 0,6s not lit | keyboard error (flashing inverted to Lock LED) |
| | error (first fault) |
| flashing: 0,1s lit / 0,1s not lit | service mode |
| not lit | disconnected from the mains power |
| | transformer switching error |

1.8.2 Accumulator Indicator (Multiboard, Attendant Control Panel)

| | MEANING |
|-----------------------------------|------------------------------------|
| lit | accumulator disconnected or faulty |
| flashing: 1,6s lit / 0,2s not lit | accumulator deeply discharged |
| flashing: 0,1s lit / 0,1s not lit | accumulator discharged |
| flashing: 0,2s lit / 1,6s not lit | accumulator is charging |
| not lit | accumulator charged |

1.8.3 Lock LED (Attendant Control Panel)

| VISUAL SIGNALISATION | lit | 0,6 | flashing: s lit / 0,6s n | oot lit | not lit |
|---|--------|---------------|-----------------------------|-------------------|----------|
| Thighrest, Calfrest and Bed Extension Lock LED | locked | lock error | keyboard error | motion blocked | unlocked |
| Backrest Lock LED | locked | lock error | keyboard error | motion blocked | unlocked |
| Bed Height, Lateral Tilt, Trendelenburg and Antitrendelen- burg Tilt Lock LED | locked | lock error | keyboard error | motion blocked | unlocked |
| Foot Switch Lock LED | locked | lock error | keyboard error | motion blocked | unlocked |

1.9 Definitions

| Basic Bed Configuration | the pricelist model configuration, not including a mattress |
|----------------------------|--|
| Bed Weight | The value depends on the product configuration, accessories or customer adjustments. |
| Clearance of Undercarriage | the height from the floor to the lowest point of the undercarriage between the castors, for the manipulation of accessories under a braked bed in the standard position |
| Duty Cycle | cycle of operation of the motor: time of activity/time of rest |
| Ergoframe | Ergoframe is the kinematic system of Mattress support platform Adjustment whose effect is the elimination of pressure on the patient's abdomen and pelvic area and frictional forces on the patient's back and legs. |
| Maximum Patient Weight | Maximum Patient Weight depends on the application environment according to IEC 60601-2-52. For application environment 1 (intensive/critical care) and 2 (acute care) reduce Safe working Load by 65 kg. For application environment 3 (long-term care) and 5 (ambulatory care) reduce Safe working Load by 35 kg. |
| Safe Working Load | the highest allowable load on the bed (patient, mattress, accessories and the load supported by those accessories) |
| Siderail Height | the height of the upper crossbar or the edges of the siderails (not the highest point of the siderail controls) from the patient surface |
| Standard Bed Position | The height of the patient surface with regard to the floor is 400 mm The mattress support platform, including the individual parts, has to be in a horizontal (level - 0°) position. The siderails are always locked in the upper position. The basic position of the integrated extension. |
| Adult | Patient having a physical size equal to or more than 146 cm, a mass equal to or more than 40 kg and a body mass index (BMI) equal to or more than 17 (according to IEC 60601-2-52). |



1.10 Abbreviations

| AC (~) | Alternating Current |
|--------|---|
| ALT | Automatic Lateral Therapy |
| CE | European Conformity |
| CPR | Cardiopulmonary Resuscitation |
| dB | Sound Intensity Unit |
| CUC | Control Unit Configuration Number |
| DC () | Direct Current |
| EAC | Euroasian Conformity |
| EMC | Electromagnetic Compatibility |
| FET | Field-effect transistor |
| HF | High Frequency |
| HPL | High Pressure Laminate |
| ICU | Intensive Care Unit |
| INT. | Duty Cycle |
| IP | Ingress Protection |
| IV | Intravenous |
| LED | Light Emitting Diodes |
| ME | Medical Electrical (Equipment) |
| MET | MET Laboratories testing and certifying for the U.S. market |
| OFF | Deactivated |
| ON | Activated |
| ppm | parts per million, millionth (1000 ppm = 0,1%) |
| REF | Reference Number (product type depending on configuration) |
| SCU | System Control Unit |
| SN | Serial Number |
| SWL | Safe Working Load |
| UDI | Unique Device Identification (for medical devices) |
| USB | Universal Serial Bus |
| WEEE | Waste Electrical and Electronic Equipment |



2 Safety Instructions



WARNING! Multicare bed should be left in its lowest position when the patient is unattended in order to reduce risk of injury due to falls!



WARNING!

Siderails of Multicare should be located in the "up" position to reduce the risk of the patient accidentally slipping or rolling off the mattress!



WARNING! Incompatible siderails and mattresses can cause an entrapment hazard!



WARNING! Inappropriate handling of the power supply cord, e. g. by kinking, shearing or other mechanical damages is hazardous!



WARNING! When routing cables from other equipment in the Multicare bed avoid squeezing those between parts of the Multicare bed!



WARNING! Multicare bed should not be used with bed hoists and bed lifts!



WARNING! To avoid risk of electric shock, this equipment must only be connected to a supply mains with protective earth.



WARNING! During specific investigations or treatments the significant risks of reciprocal interference posed by medical electrical equipment may occur.



WARNING! No modification of this equipment is allowed.



WARNING! The bed is intended for adults. Follow chapter Specifications of Use.



WARNING! Incompatible mattresses can create hazards.



WARNING! To avoid the risk of electric shock, this equipment must only be connected to a supply mains with protective earth.





WARNING!

No modification of this equipment is allowed.



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



WARNING!

If this equipment is modified, appropriate inspection and testing must be conducted to ensure continued safe use of the equipment.



WARNING! An additional multiple socket-outlet or extension cord shall not be connected to the medical electrical system.



WARNING! During specific investigations or treatments the significant risks of reciprocal interference posed by ME equipment may occur.



WARNING! Staff expert assessment is needed to consider all individual cases of contraindications!



WARNING!

Certain bed positions are not suitable for specific diagnosis/medical conditions. Fowler position is not suitable for spinal cord injuries! Trendelenburg position is not suitable for patients with higher intracranial pressure!



WARNING!

Length adjustment of the bed must be proportional to the height of patient! Risk of trapping or squeezing because of patient's body constitution disproportionate to the size of mattress support platform!



WARNING!

Any serious incident that has occurred in relation to the device should be reported to the manufacturer and the competent authority of the Member State in which the user and/or patient is established!



WARNING! Only authorised and trained person using the tool is allowed to change fuses and power supplies!



WARNING! This medical device is not intended for oxygen enriched environment!



WARNING! This medical device is not intended for use with flammable substances!



WARNING! This medical device is not portable medical electrical equipment!





WARNING!

Make sure the duty cycle (2 min ON/18 min OFF) is not exceeded during bed positioning!

WARNING!

Patient is allowed to use selected control elements only if hospital personnel had assessed that the patient's physical and psychological state is in accordance with use of them and only if the hospital personnel had trained the patient in accordance with the instructions for use!



WARNING!

Hospital personnel is allowed to use the weighing system (scales) for weighing patients only if they had been trained according to the instructions for use!

- Follow the instructions in the instructions for use carefully.
- Use the bed exclusively if it is in perfect working order.
- If necessary, check the bed functions daily or at each shift change.
- Ensure any user has read and understood this manual completely before operating the product.
- Use the bed exclusively with the correct mains supply.
- Ensure that the bed is operated exclusively by qualified personnel who have been trained according to the instructions for use.
- Ensure that the patient (health permitting) has been informed about the operation of the bed and all applicable safety instructions.
- Move the bed exclusively on even, hard-surfaced floors.
- Replace any damaged parts immediately with original spare parts.
- Ensure that maintenance and installation are performed exclusively by qualified personnel who have been trained by the manufacturer.
- During peak loads or unavoidable excess loads (CPR), place mattress platform in the lowest position.
- Ensure that only one adult patient uses the bed at any time.
- Take care to avoid injuries or squeezing when operating moving parts.
- When using lifting poles or infusion stands, ensure that nothing will be damaged when you move or adjust the bed.
- Brake the castors when the bed is occupied.
- Keep the mattress platform in the lowest position at any time when the healthcare personnel are not trea- ting the patient in order to prevent the patient from falling or injuries.
- Ensure that siderails are operated exclusively by healthcare personnel.
- Never use the bed in areas where there is a hazard of explosion.
- Enable or disable functions on patient controls using the Multiboard panel as appropriate for the patient's physical and mental state. Verify that the function is actually disabled.
- Never handle the mains plug with wet hands.
- Disconnect the product from the mains exclusively by pulling the mains plug.
- When pulling the mains plug, always hold the plug, not the cable.
- Position the mains cable so that there are no loops or kinks in the cable; protect the cable from mechani- cal wear and tear.
- Improper handling of mains cable can cause an electric shock hazard, other serious injuries or damage to the mattress replacement system.
- Ensure that the stipulated duty cycle (on-time) is not exceeded.
- Ensure that the moving parts of the bed are not blocked.
- ▶ To prevent failures, use exclusively the manufacturer's original accessories and mattresses.
- Ensure that the stipulated safe working load is not exceeded.
- If the patient's condition could lead to an entrapment, leave the mattress support platform in the flat position whilst unattended.
- When adjusting the ALT (automatic lateral therapy), ensure that the process does not pose any risk of the patient falling or getting injured, or of any lines or tracheal tubes getting pulled out.
- Adjust bed height to approx. 20 cm below maximum height when transporting the bed in order to facilitate overcoming possible obstacles.
- Do not exceed maximum load of 80 kg for mattress platform extension.
- Ensure that the bed and its components are exclusively modified with the manufacturer's approval.
- ▶ Use the mattress system exclusively as specified in this manual and in perfect working order.
- Use the mattress system exclusively with the correct mains supply.
- Use the mattress system exclusively in its original state and do not modify it in any way.
- ► Have the mattress system used exclusively by or under supervision of trained and qualified nursing per- sonnel.
- ► Have the mattress system serviced and installed exclusively by qualified personnel trained and authorised by the manufacturer.
- Do not use the SCU near flammable gases.



- Use the mattress system exclusively in its original state and do not modify it in any way.
- Have the mattress system used exclusively by or under supervision of trained and gualified nursing per-sonnel. Have the mattress system serviced and installed exclusively by qualified personnel trained and authorised by the manufacturer.
- Do not exceed the maximum patient weight limit (see Mechanical Specifications (OptiCare or Symbioso)).
- Do not use the SCU (System Control Unit) near flammable gases (This does not apply to oxygen cylinders.).
- Never use the mattress replacement system near radiators or other heat sources.
- Never cover the SCU while in use.
- This does not apply to oxygen cylinders.
- Never use the mattress replacement system near radiators or other heat sources.
- Select a suitable location for the placement of bed accessories and other objects to prevent involuntary activation of buttons or controls which may result in the adjustment of bed positioning.
- Do not use the bed in the event parts have been removed (e.g. parts of mattress platform) unless these parts are de signed to be removed (e.g. head and/ or foot board).
- Never place any accessories or handset on the siderails in the area where the integrated siderail controller is located. After each emergency situation always check if any of the controllers (in siderails, handset or ACP) is not involuntarily pressed by the bed accessories or by the mattress.
- The weighing system must be tested at regular intervals and in accordance with the metrological regulations of the relevant country. All testing and certification must be carried out by qualified personnel. The healthcare provider is responsible for ensuring the required testing frequency and testing procedure of the weighing system is carried out.
- To avoid injury or crushing, take extra caution when operating any moving parts of the bed. To avoid unintended activation of moving parts during any use of the bed always check that none of the control elements
- of the bed is not involuntary pressed by persons, mattress or other objects.

3 Intended use

The intended use is the hospitalization of the patient in the intensive and acute care units, which includes above all the following aspects:

- Adjustment of the specific positions needed for the preventive reasons, routine nursing, treatments, mobilization, physiotherapy, examinations, sleeping, and relaxation. These positions are further specified and described in the clinical evaluation of this device, together with their potential clinical outcomes and benefits.
- Providing the safe environment for the patient during all relevant procedures. The particular requirements on patient safety are the subject of the clinical evaluation, including evaluation of the risk/benefit ratio. The relevant safety issues are the part of the risk management file.
- Patient in-bed indoor transport out of the patient room.
- Providing the suitable working conditions for the caregivers to perform the routine and specific tasks during the patient hospitalization.
- Indicative measurement of the patient weight, used as supportive feature without direct diagnostic effect. It helps staff to assess the general patient status and apply the nutrition and medicaments (valid for the version of the beds with in-bed scales).

3.1 User population

- Adult patients (weight >= 40 kg, height >= 146 cm, BMI >= 17) in the intensive and acute care units (Application Environment 1 and 2 as in IEC 60601-2-52)
- Caregivers (nurses, doctors, technical personnel, transport personnel, cleaning personnel)

3.2 Contraindications

- The medical device is not intended for the pediatric patients use.
- Certain positions are not suitable for specific diagnoses/medical conditions (e.g. spinal cord injuries vs. Fowler position, higher ICP patients vs. Trendelenburg). Staff expert assessment / nursing consideration is needed in all individual case of contraindication.

3.3 Operator

- Caregiver
- Patient (based on individual patient status assessment by caregiver the patient can utilize dedicated device functions)

4 Product Description



Fig. Overview of Multicare

- 1. Removable Foot Board with Safety Lock
- 2. Foot Siderail with Integrated Control Panels for Patient
- 3. Split Siderail Head Siderail
- 4. Four-part Mattress Platform with Ergoframe® System (under the mattress)
- 5. Multiboard with LCD Touchscreen
- 6. Quick-Action Panel
- 7. Removable Head Board
- 8. CPR Control Lever Backrest Release
- 9. X-Ray Cassette Holder
- 10. Accessory Holder
- 11. Siderail Release Lever
- 12. Bi-lateral Accessory Rail
- 13. Castor Control Lever
- 14. Castor Diameter 150 mm (5.9 in.) with Main Control Lever
- 15. Mobilift® Handles
- 16. Bumpers

NOTE For safe, easy handling, LINET® recommends having two technicians assemble the bed.

5 Technical Specification

All technical data are rated data and are subject to construction and manufacturing tolerances.



WARNING! If Multicare bed is used with OptiCare integrated mattress replacement system or with Symbioso integrated mattress replacement system, respect values of mechanical and electrical specifications which can harm none of them!

5.1 Identification of Applied Parts (Type B)

All part of the bed (and accessories) the patient can reach are type B Applied Parts.

- Mattress support platform frame, Covers and all Movable Parts
- Head Board and Foot Board
- Siderails
- Mobi-Lift Handles
- Handset

5.2 Scales (only version with scales)

Accuracy of displayed weight values:

- 0,5 kg (1,1 lbs)
- Scales Class III

5.3 Mechanical Specifications (Multicare)

| Parameter | Value |
|--|------------------|
| Dimensions (With Folded-up Siderail) | 215 cm x 105 cm |
| Bed Extension | 0 cm - 22 cm |
| Recommended Mattress Dimensions | 208 cm x 86 cm |
| Maximum Mattress Height | 23 cm |
| Bed Height | 44 cm - 82 cm |
| Siderail length Head section Central section | 54 cm 100 cm |
| Castor (Diameter) | 15 cm |
| Maximum Backrest Angle | 70° |
| Maximum Thighrest Angle | 30° |
| Maximum Calfrest Angle | 38° |
| Maximum Lateral Tilt Angle | 30° |
| Trendelenburg | 13° |
| Anti-Trendelenburg Position | 16° |
| Siderail Height (above Mattress Platform) | 45 cm |
| Bed Weight (Basic Equipment) | 224 kg |
| Safe Working Load | 250 kg |
| Maximum Lifting Pole Load | 75 kg |
| Maximum Patient Weight Application environment 1, 2 Application environment 3, 5 | 185 kg 215 kg |



5.4 Environment Conditions

| Use Conditions | |
|----------------------------------|--------------------|
| Ambient Temperature | 10°C - 40°C |
| Relative Humidity | 30% - 75 % |
| Atmospheric Pressure | 795 hPa - 1060 hPa |
| Storage and Transport Conditions | |
| Ambient Temperature | -20°C - 50°C |
| Relative Humidity | 20% - 90 % |
| Atmospheric Pressure | 795 hPa - 1060 hPa |

5.5 Electrical Specifications (Multicare)

| Parameter | Value |
|-----------------------------------|--------------------------------------|
| Input Voltage | 120 V AC, 50/60 Hz |
| Maximum Power Input | max. 370 VA |
| Ingress Protection | IPX4 |
| Protection Class Applied parts | Class I Type B |
| Electrical Motor Duty Cycle | max. 2 min ON / 18 min OFF |
| Accumulator | Pb ACCU 2 x 12 V / 1,2 Ah / Fuse 15A |
| Fuse | 2x T3.15A L 250 V for 120 V version |

ERGOFRAME

Ergoframe[®] is the kinematic system of Backrest and Thighrest Adjustment resulting in extension of the Mattress support platform in the seat section.

Ergoframe[®] enlarges the space for pelvic area during Auto-contour. Because of increasement of the space the force applied results in decrease of the pressure that can cause pressure injuries in the pelvic area.

Ergoframe maintains a stable ergonomic position of the body and spine of the patient, thus limiting unwanted movement of the patient by moving down or up in beds. Unified movement eliminates the patient's shift over the mattress and thus maintains a uniform position of the patient's body that is not bound to the position of the bed parts.

5.6 Electromagnetic compatibility

Bed is intended for hospitals except for near active HF surgical equipment and the RF shielded room of a medical system for magnetic resonance imaging, where the intensity of EM disturbances is high.

Bed has defined no essential performance.



WARNING!

It is recommended to avoid the use of this device next to or in block with other device, because it could lead to improper operation. If such use is needed, this device and the other equipment should be under surveillance to verify proper operation.

List of used cables:

- 1. Mains cable, maximum length 6 m
- 2. ACP Supervisor control panel, maximum length 3m
- 3. Handset, maximum length 3m



WARNING!

Use of the accessories, converters and other cables, than specified and provided by manufacturer of this bed could lead to increase of electromagnetic emission or lower the electromagnetic immunity of this bed and lead to improper operation.



WARNING!

Mobile RF communication device (including end use devices like antenna cables and external antenna) should not be used closer than 30 cm (12 inches) from any part of this bed Multicare, including cables specified by manufacturer. Otherwise this could lead to deterioration of functionality of this bed.



WARNING!

Do not overload the bed (SWL), respect the duty cycle (INT.) and consider chapter 21 Maintenance in order to maintain the basic safety with regard to electromagnetic disturbances for the expected service life.

5.6.1 Manufacturer instructions – electromagnetic emissions

| Emission test | Compliance |
|---|------------|
| RF emissions CISPR 11 | Group 1 |
| RF emissions CISPR 11 | Class B |
| Harmonic emissions IEC 61000-3-2 | Class A |
| Voltage fluctuations / flicker emissions IEC 61000-3-3 | Complies |



5.6.2 Manufacturer instructions – electromagnetic susceptibility

| Immunity Tests | Compliance level |
|---|---|
| Electrostatic discharge (ESD) | ± 8 kV for contact discharge |
| IEC 61000-4-2 | ± 15 kV for contact discharge |
| Radiated RF | 3 V/m |
| IEC 61000-4-3 | 80 MHz – 2,7 GHz |
| Proximity fields from RF wireless communications equipment | 80 % AM at 1 kHz |
| IEC 61000-4-3 | See Table 1 |
| Fast electrical transients / burst | ±2 kV for power line |
| IEC 61000-4-4 | repetition frequency 100 kHz |
| Surge | ± 1 kV Line-to-line |
| IEC 61000-4-5 | ± 2 kV Line-to-ground |
| Conducted RF IEC 61000-4-6 | 3 V (0,15 MHz – 80 MHz) 6 V in ISM bands between 0,15 MHz and 80 MHz 80 % AM at 1 kHz |
| Power frequency (50/60 Hz) magnetic field IEC 61000-4-8 | 30 A/m |
| Voltage dips, short interruptions on power supply input lines IEC 61000-4-11 | 0 % UT; 0,5 cycle At 0°, 45°, 90°, 135°, 180°, 225°, 270° a 315° 0 % UT; 1 cycle and 70 % UT; 25/30 cycle Single phase: at 0° 0 % UT; 250/300 cycle |

Table 1 - IMMUNITY to RF wireless communications equipment

| Test frequency (MHz) | Band (MHz) | Service | Modulation | Immunity Test Level V/m |
|-------------------------|---------------|---|------------------------------------|----------------------------|
| 385 | 380 - 390 | TETRA 400 | Pulse modulation 18 Hz | 27 |
| 450 | 430 - 470 | GMRS 460, FRS 460 | FM ± 5 kHz deviation 1 kHz sine | 28 |
| 710 745 780 | 704 - 787 | LTE band 13, 17 | Pulse modulation 217 Hz | 9 |
| 810 870 930 | 800 - 960 | GSM 800/900, TETRA 800, iDEN 820, CDMA 850, LTE band 5 | Pulse modulation 18 Hz | 28 |
| 1 720 1 845 1 970 | 1 700 - 1 990 | GSM 1800; CDMA 1900; GSM 1900; DECT; LTE band 1, 3, 4, 25; UMTS | Pulse modulation 217 Hz | 28 |
| 2 450 | 2 400 - 2 570 | Bluetooth, WLAN, 802.11 b/g/n, RFID 2450, LTE band 7 | Pulse modulation 217 Hz | 28 |
| 5 240 5 500 5 785 | 5 100 - 5 800 | WLAN 802.11 a/n | Pulse modulation 217 Hz | 9 |

NOTE There are applied no deviations to requirements of IEC 60601-1-2 ed. 4

NOTE There are no known other measures for keeping the basic safety based on EMC phenomena.

NOTE Beds equipped with integration module meet standard for IEEE 802.11 b/g/n (2400,0 MHz – 2483,5 MHz, modulation DSSS (IEEE 802.11 b), OFDM (IEEE 802.11 g/n) 20MHz bandwidth, EIRP = 0,34 W).

6 Use and Storage Conditions

DANGER!

Danger to life due to electric shock!

To ensure the bed's class I protection against electric shocks:

- Ground the mains.
 - Use exclusively Hospital Grade or Hospital Only receptacles for grounding.



CAUTION! Minimal clearance underneath the bed (standard version with 15 cm castors) is 4.4 cm!

• Observe the path for any obstacles and avoid collisions and possible damages of any bed's part on the undercarriage.

Do not use bed lifts and hoists for lifting the bed (bed hoists for patients are permitted; clearance for the patient hoists is 15 cm).

Multicare with OptiCare or with Symbioso are designed for use in rooms for medical purposes. Electrical installations must therefore meet local norms laying down the necessary conditions for electrical installations.

Disconnect the bed from the mains in exceptional cases (i.e. lightning or earthquake).

Multicare, OptiCare and Symbioso are not suitable for indoor environments:

containing flammable gases (except oxygen cylinders).



7 Scope of Delivery and Bed Variants

7.1 Delivery

- Upon receipt, check that the shipment is complete as specified on the delivery note.
- Notify the carrier and supplier of any deficiencies or damages immediately as well as in writing or make a note on the delivery note.

7.2 Scope of Delivery

- Multicare medical bed
- Mattress with cover Applied part type B
- SCU (System Control Unit) Applied part type B
- Instructions for use

7.3 Multicare Variants

s = standart

o = optional

Optional bed features:

- Undercarriage
- Standard undercarriage under bed clearance under foot columns 44mm
- □ Higher undercarriage under bed clearance under foot columns 69mm
- OptiCare ready bed with SCU
- with OptiCare
- without OptiCare
- Symbioso
- with Symbioso
- without Symbioso
- Scales
- with scales (with Bed Exit Monitoring)
 without scales (without Bed Exit Monitoring)
- Automatic Lateral Therapy (ALT)
- □ with ALT
- without ALT
- Castors
- □ Tente Integral 150 mm (5.9 in.) double castors (s)
- □ Tente Integral 150 mm (5.9 in.) single castors (o)
- □ retractable fifth castor (o)
- Control Elements
- Multiboard with LCD touchscreen in both head siderails (s)
- Quick-Action panel in both head siderails (s)
- Attendant Control Panel (o)
- □ handset with adapter for simple connection (Plug and Play) (o)
- □ handset with illuminated buttons and adapter for simple connection Plug and Play) (o)
- □ foot control for lateral tilt (o)
- □ foot control for height adjustment (o)
- patient control elements integrated in both foot siderails (s)
- Version without integrated control elements for patients (o)
- Integrated control elements without illumination (o)
- Illuminated integrated control elements for patients (o)
- 1 pair of Mobi-Lift® handles (o)
- i-Brake® (o)
- x-ray cassette holder
- Additional adapter for lifting pole (o)
- LAN/Wi-fi module (o)
- Hercules ready bed (o)
- EMR ready bed (o)
- Nurse call (o)
- i-Drive Power® (o)
- LINIS SafetyPort
- without LINIS SafetyPort (s)
- basic hardware preparation for LINIS SafetyPort (CE06: without Integration Module) (o)
- complete hardware preparation for LINIS SafetyPort (CE31: with Integration Module) (o)



8 Putting into Service



WARNING! Risk of injury when working on the bed!

- Ensure that the bed is disconnected from the mains connection prior to putting into service, putting out of service and maintenance.
- Ensure that the castors are locked prior to putting into service, putting out of service and maintenance.



CAUTION!

Material damage due to incorrect putting into service!

Ensure that putting into service is performed exclusively by manufacturer's customer service or trained hospital personnel.

Set up the bed as follows:

- Unpack the bed.
- Check the delivery (see Scope of Delivery and Bed Variants).
- Remove isolating foil from mains control box (see Accumulator Activation).
- Install equipment and accessories.

► In case of delivery with dismounted head board and foot board, mount the head board and foot board (see Head Board and Foot Board).

- Set-up the bed only on a suitable floor surface (see Transport).
- Ensure the mains cable does not collide or get stretched when adjusting the bed. Check the plug is inserted correctly.
- Do not leave any extension cords or power strips loose on the floor.
- Ensure all the required mechanical and electrical prevention mechanisms are available on site.
- There is no mains switch on the bed, i.e. the mains cable is the only means to isolate the bed from the mains.
- Ensure the mains cable is always accessible.
- ► The plug on the mains cable should only be changed and maintained by qualified and trained service technicians authorised by the manufacturer.



8.1 Accumulator Activation

8.1.1 Placement of Control Section



Fig. Control section placement

8.1.2 Removing the Isolating Foil



Fig. Isolating foil



To remove isolating foil:

- Remove isolating foil from mains control box 1 by pulling strap 2. Check if isolating foil is complete and undamaged.
- If isolating foil is damaged, contact the manufacturer's service department immediately.

NOTE: It is recommended to wear gloves when removing the isolating foil.



8.2 Head Board and Foot Board



Fig. Foot Board Locks



Fig. Installed Head Board

Dismount the foot board as follows:

- Unlock sleeve fittings.
- Pull foot board from sleeve fittings.
- Lock sleeve fittings.

Install the foot board as follows:

- Unlock sleeve fittings.
- Slide foot board into sleeve fittings.
- Lock sleeve fittings.

Dismount the head board as follows:

Pull head board from sleeve fittings.

Install the head board as follows:

Slide head board into sleeve fittings.



8.3 Potential Equalisation

The bed is equipped with a standard protective connector. This connector is used for potential equalisation between the bed and any intravascular or intracardiac device connected to the patient to protect the patient from static electric shocks.





Fig. Potential equalisation connector - female

Fig. Potential equalisation connector - male

Use equalisation connector if:

the patient is connected to any intravascular or intracardiac device.

Before connecting the patient to an intravascular/intracardiac device:

• Connect the ground wire of the device to the potential equalisation connector (male) on the bed on which the patient in question is lying.

- Use a standard hospital connector (female).
- Make sure that the connectors match.
- Make sure that there is no possibility for inadvertent disconnection.

Before moving the bed:

- Disconnect the patient from the intravascular or intracardiac device.
- Disconnect the potential equalisation connector.

8.4 Before Use

Prepare the bed for service as follows:

- Connect the bed to the mains.
- Charge the accumulator.
- Raise and tilt the mattress platform to the highest position.
- Lower and tilt the mattress platform to the lowest position.
- Check that the castors as well as main brake work correctly.
- Check that the bed extension works correctly.
- Check that it is possible to remove the head and foot boards.
- Check all of the functions on the control elements (Multiboard etc.).
- Check that the siderails function properly.
- Dispose of all packaging (see Disposal).

8.5 Transport

For a safe transport, observe the following:

- Ensure that no cables are run over when moving a bed.
- Ensure that the mains cable is attached with a hook (at the head board).
- Ensure that the castors are unlocked before moving the bed during the loading/unloading process (see Castor Control and Bed Transport).
- Move the bed exclusively on suitable floor surfaces.

Suitable surfaces:

- Tile
- Hard linoleum
- Poured flooring

Unsuitable surfaces:

- Too soft, unsealed or defective flooring
- Soft wooden flooring
- Soft and porous stone floors
- Carpeted floors with underlay
- Soft linoleum
 - For longer distances, ensure that the castor steering function (main control) is activated.
 - Ensure that the brakes are released while moving the bed.

8.6 Firmware

The bed includes firmware that can be updated only by an authorised service technician.

This firmware is protected against unauthorised access by mechanical housing (tool is needed to access), by seal (components with processor are sealed), by exclusive compatibility with an authorised software tool and by check of compatibility of the new firmware with the bed.

9 Power Supply Cord (Mains Power Cable)



CAUTION!

Disconnecting bed from the mains does not stop motions of the bed!
 Stop the bed before disconnection bed from the mains.

Attachment plug is means of connecting and disconnecting bed from the mains. Power supply cable (mains power cable) must be attached with a hook at the head board during transport.

Where the integrity of the external protective conductor in the installation or its arrangement is in doubt

operate the bed from internal accumulator only.

10 Accumulator

CAUTION!



Material damage due to temperature difference!

► If there is a considerable temperature difference between the bed and the place of operation (after transport/storage), leave bed unconnected for 24 for the difference to balance itself.

The accumulator supplied with the bed is delivered uncharged. The accumulator serves as a backup during power failures or while transporting the patient.

For declared lifetime period of leaded accumulators is recommended during storage:

- 1) To prevent accumulators from deep discharging and to keep accumulators at least partly charged by regular recharging
- 2) To store accumulators on the places with temperature from 10°C to 40°C
- 3) To prevent accumulators from being in the sunshine

Accumulator lifetime could be up to 5 years if operated under optimum conditions. Accumulator capacity can be significantly reduced if:

- too high ambient temperature
- many accumulator charge/discharge cycles
- recurrence of deep discharge
- bed is often powered only by the accumulator

10.1 Accumulator Operation

- Use only accumulator approved by the manufacturer.
- Check the functionality of accumulators at least once a month in accordance with the user and service manuals and have the accumulator changed if necessary.
- Use exclusively accumulator approved by the manufacturer.

NOTE The service life of the accumulator depends on the frequency and method of use.

The manufacturer will assume no responsibility for any damage to the bed or the accumulator caused by:

- non-observance of the manufacturer's instructions in the instructions for use
- using accumulator not approved by the manufacturer

NOTE The manufacturer provides a 6-month warranty for the full function of the accumulator.

To charge the accumulator:

Connect the bed to the mains.

NOTE Some bed adjustment options are not available without a accumulator, for example, height adjustment under a load of above 200 kg.

The LED indicates the Accumulator charge status:

| Yellow LED | Accumulator charge status |
|--|--|
| Not lit | Accumulator capacity is sufficient (charging completed) |
| Short flashing (shortly lit, longer not lit) (circa 1.8 sec.) | Accumulator is charging - continue charging until the LED is extinguished. In emer- gency cases, the accumulator can be used as a backup power source for a short period. If LED is still flashing after 12 hours of charging or stops flashing, but you can not position with bed, accumulator is defective or broken. Contact manufacturer. |
| Short flashing (0,2s lit, 0,2s not lit) | Only CPR function can be used. |
| Long flashing (longer lit, shortly not lit) (circa 0.2 sec.) | Low accumulator voltage - accumulator can not be used as a backup power supply even for a short period; accumulator is completely discharged or defective (if this type of signalisation persists, it is necessary to replace the accumulator - service action) |
| Lit continuously for several hours (circa 10 hours), when bed is connected to the mains. | Accumulator absence or failure condition (accumulator is connected incorrectly, line between the power supply and accumulator is broken or accumulator fuses are faulty); contact service department of the manufacturer in case of such signa- lisation. |

10.2 Replacing the accumulator



CAUTION!

Damage to the bed due to incorrect accumulator replacement!

- Have the accumulator replaced exclusively by qualified personnel.
- Exclusively use accumulator approved by the manufacturer.



CAUTION! Material damage due to overheating!

If the accumulator is faulty, degassing may occur. In rare cases this might cause deformations of the accumulator case, control panel housing or cable.

- Stop using the bed immediately (see Removing the Bed from Use).
- Inform the manufacturer's service department.



CAUTION!

Risk of reducing accumulator durability due to incorrect use!

- Use bed on accumulator only in crisis situations (e.g.: power blackout, patient complications during transport,
- etc.)
 - After reconnecting bed to the mains charge accumulator to full capacity (see chart Accumulator charge status).
- Have accumulator replaced exclusively by a qualified service organisation.
- For more detailed information on how to replace the accumulator, request service manual from manufacturer.

Status "Faulty accumulator"

The accumulator is regarded as faulty if at least one of the following conditions applies:

- Accumulator charging constantly
- Low voltage on accumulator
- Low charging current of accumulator

This status is indicated by the accumulator status indicator being constantly lit. These statuses are summarised to Linis and written to Blackbox.



To cancel this status:

Press STOP button.

Status "Discharged accumulator"

The accumulator is regarded as discharged if the following condition is met:

- Defined decrease of voltage depending on discharging current
- This status is indicated by the accumulator status indicator flashing quickly.
- The electric CPR position is the only possible position.
- This status will be cancelled automatically when the bed switches to sleep mode.

To cancel this status:

Press STOP button.

10.3 Removing the Bed from Use

Remove the bed from service as follows:

- Disconnect the bed from the mains.
- Disconnect the ground wire.
- Deactivate the accumulator.
- Remove accessories.

To prevent damage during storage:

- Pack or cover the bed and accessories.
- Ensure that storage conditions are the same as the operating conditions.

10.4 Deactivating the Accumulator

To avoid damaging the bed and the environment during storage:

Deactivate the accumulator on the Attendant Control Panel.

To deactivate the accumulator on the supervisor:

- Disconnect the bed from the mains.
- Disconnect the ground wire.
- Activate the keypad by pressing the GO button on the supervisor.

Press the Thighrest Up + Thighrest Down + Trendelenburg Position buttons at the same time and hold them for three seconds.

The accumulator is deactivated.

To activate the accumulator again:

Connect Power Cable to the mains.

11 Manipulation



WARNING! Risk of injury when adjusting the bed!

- Ensure that there are no body parts between the mattress support platform elements and the mattress support platform frame when adjusting the bed.
- Ensure that there are no body parts below the mattress support platform frame before adjusting the bed.

11.1 Siderails



Risk of injury, damaging or involuntary movement of the bed due to incorrect placement of accessories or handset!

Never place any accessories or handset on the siderails in the area where the integrated siderail controller is located.

The split siderails are components of the bed. A pneumatic spring supports the operation of the split siderails. The nursing personnel are responsible for the siderails being folded up while the patient is in bed. The correct placement of handset is shown at following picture.



MANIPULATION

To raise siderails up:

- Grab siderail by Siderail Handle (1).
- Pull siderail up until it latches. You will hear audible "click".

To release siderails down:

- Grab siderail by Siderail Handle (1).
- Press upper edge of siderail inwards.
- ▶ Unlock siderail by pulling Siderail Release Handle (3) to yourself.
- Fold down siderail slowly.



11.2 Castor Control and Bed Transport



CAUTION!

Material damage due to incorrect transport and involuntary movement!

- Prior to transport, ensure that the bed is disconnected from the mains.
- Prior to transport, ensure that the auxiliary outlet plug (if available) is disconnected from the mains.
- Ensure that the castors are locked prior to assembly, disassembly and maintenance.
- Ensure that the castors are locked when the bed is occupied.
- Hang the mains cable on the appropriate hook on the bed during transport.
- ► Have the bed transported exclusively by nursing personnel and by at least 2 persons.

11.2.1 Central Castor Control

The bed is equipped with central castor's control and brake system. The control levers are located in the four corners of the undercarriage.



Castor control lever positions:

1. Forward Movement

The front left castor is locked. The bed moves straight ahead. If the bed is equipped with a fifth castor, this castor determines the direction of movement.

2. Unrestricted Movement

- All of the castors are unlocked.
- 3. Braked
- All of the castors are braked.

Fig. Positions of Castor Control Lever

11.2.2 Bed transport



Transporting the bed:

- Adjust bed height to at least 20 cm (9 in.) below maximum height.
- Push bed by handles on head board or foot board.



11.3 CPR Backrest Release



WARNING! Risk of injury due to lowering the backrest too quickly!

- Ensure that the siderails are in the lowest position.
- Ensure that there are no body parts between the siderails and the backrest.
- Press the backrest down using the mattress guard handle only.

The bed permits quick, mechanical lowering of the backrest for emergency resuscitation (CPR) procedures.



Fig. CPR lever (release handle)

Set the position as follows:

- Pull and hold release handle.
- Press backrest down.
11.4 Control Elements

The bed is operated by different control elements.

Control elements depending on the model:

- Multiboard with LCD touchscreen in both head siderails
- Quick-Action panel in both head siderails
- Additional supervisor
- Handset
- Handset with adapter for easy connection (Plug and Play)
- Handset with illuminated buttons
- Foot control for lateral tilt
- Foot control for height adjustment
- Patient control elements integrated in both foot siderails

Disabling individual functions on the Multiboard will affect all control elements.

If the bed does not react to individual position settings:

Check whether the function is disabled on the Attendant Control Panel.

11.4.1 Multiboard with LCD Touchscreen in Both Head Siderails

The Multiboard is the main control element. It is integrated in the outside of both head siderails.

Ensure that exclusively nursing staff trained for critical care operate the Multiboard.



7.

8.

9.

Fig. Multiboard

- 1. Calfrest Position Buttons
- 2. Thighrest Adjustment Button
- 3. Backrest Adjustment Button
- 4. LCD Touchscreen
- 5. Buttons Mattress Platform Extension
- 6. Buttons Longitudinal Tilt Adjustment
- Accumulator Charge Status LED
- Mains Power LED
- **Button Cardiac Chair Position**
- 10. Button CPR (Resuscitation) Position
- 11. Central STOP Button
- 12. GO Button



Central STOP Button

The central STOP button immediately interrupts all bed movements in case of unauthorized bed positioning or an electronic failure. Pressing the central STOP button for at least 0.3 seconds immediately stops all electronic bed movements.

Activating GO Button

The GO button activates the keypad or the touchscreens of all control elements.

A GO button is included on a number of different control elements. The function of the GO button is identical on all control elements. Pressing a function button will keep the keypad active for another 3 minutes.

During this time the following is possible:

- Adjusting individual mattress platform elements by pressing the corresponding function buttons.
- Disabling individual functions with the lock buttons.

Each time a function button is pressed, the keypad will remain active for another 3 minutes.

Function Buttons

The positioning function buttons 1, 2, 3, 5 and 6 adjust the position of the backrest, thighrest and calfrest as well as the tilting and extending of the mattress platform. The buttons 9 and 10 allow adjusting the Cardiac Chair and CPR positions.

Button CPR (Resuscitation) Position

If the bed is equipped with OptiCare mattress or with Symbioso mattress, pressing button 10 will also deflate the mattress.

NOTE Pressing two function buttons at the same time will be recognized as an error by the controller. The controller will interrupt immediately all bed movements immediately.

Set the position as follows:

- Activate the keypad by pressing the GO button.
- Press and hold function button until desired position is reached.

Mains power LED

| Status | Meaning |
|--------------|-----------------------------|
| lit LED | connected to the mains |
| unlit LED | disconnected from the mains |
| flashing LED | system error |



11.4.2 Attendant Control Panel

The Attendant Control Panel is an optional control element. The Attendant Control Panel can be hung from the foot board if required. It is possible to hold the additional supervisor in the hand while operating.



Fig. Attendant Control Panel

- 1. Button and LED Thighrest, Calfrest and Extension Lock 10.
- 2. Button Thighrest Adjustment
- 3. Button and LED Backrest Lock
- 4. Button Backrest Adjustment
- 5. Button and LED Height/Tilt Lock
- 6. Buttons Height Adjustment
- 7. Buttons Calfrest Position
- 8. Buttons Mattress Platform Extension
- 9. Button and LED Foot Control Lock

- Buttons Longitudinal Tilt
- 11. Button Cardiac Chair Position
- 12. Buttons ALT
- 13. Button Trendelenburg Position
- 14. LED Mains Power
- 15. LED Accumulator Charge Status
- 16. Button CPR (Resuscitation) Position
- 17. GO Button

To set position:

- Activate the keypad by pressing the GO button.
- Press and hold corresponding button until desired position is reached.

Mains power LED

| Status | Meaning |
|--------------|-----------------------------|
| lit LED | connected to the mains |
| unlit LED | disconnected from the mains |
| flashing LED | system error |



11.4.3 Handset

A handset is included with the bed as an optional feature. The position of the handset depends on the patient's condition. The handset is available with and without button illumination. The illumination is activated for 7s if any button was pressed and the illumination is activated for 3 minutes if GO Button was pressed. The functions of both handsets are identical.



- 1. Buttons Thighrest Position
- 2. LED Thighrest/Backrest Lock
- 3. Button Backrest Position
- 4. GO Button
- 5. Button Autocontour
- 6. Button Flashlight
- 7. LED Height Lock
- 8. Buttons Height Adjustment

To switch on the flashlight:

Press flashlight button 6.

Set the position as follows:

- Activate the keypad by pressing the GO button.
- Press and hold function button until desired position is reached.

NOTE Depending on the patient's condition, the nursing staff decides whether the patient is allowed to adjust the bed's position.

If required, prevent the patient from adjusting the bed as follows: Disable functions.

Disable function

NOTE An adapter for the handset is available. The adapter enables quick installation and removal (e.g. replacing a defective handset, using the handset for another bed).

Fig. Handset

11.4.4 Patient Control Panels

The patient control panels integrated in the foot siderails allow the patient to adjust the positions of the Backrest, Thighrest and Autocontour.



Fig. Patient Control Panels in the foot siderails

1. Autocontour Adjustment Button - (simultaneous movement of the Backrest and Thighrest) - DOWN

- 2. GO Button (activation of the control panel)
- 3. Autocontour Adjustment Button (simultaneous movement of the Backrest and Thighrest) UP
- 4. Backrest Adjustment Button UP
- 5. Backrest Adjustment Button DOWN
- 6. Thighrest Adjustment Button UP
- 7. Thighrest Adjustment Button DOWN

NOTE Keyboards are optionally illuminated. The illumination is activated for 7s if any button was pressed and the illumination is activated for 3 minutes if GO Button was pressed.

NOTE Functions on the Patient Control Panel in the foot siderails are disabled when the foot siderail is in lower position.



11.4.5 Bed Height Foot Control

The foot control is optional and allows setting the height of the bed with one's feet.



- Protection Frame against Unwanted Activation 1.
 - Foot Switch Raise Mattress Platform
- Foot Switch Examination Position 3.
- 4. Foot Switch Lower Mattress Platform

Set the position as follows:

2.

- Press foot switch 2, 3 or 4 to activate foot control.
- Press and hold foot switch until desired position is reached.

NOTE: It is possible to activate foot control by pressing GO button on the control elements of the bed then it is not needed to activate the foot control by buttons 2, 3 or 4.

Fig. Foot Control Bed Height

11.4.6 Lateral Tilt Foot Control

The foot control is optional and allows setting the lateral tilt of the bed with the feet.



- Protection Frame against Unwanted Activation
- Foot Switch Tilt Right 2.
- 3. Foot Switch GO 4.

1.

Foot Switch Tilt Left

Set the position as follows:

- Activate the keypad by pressing the GO button.
- Press and hold foot switch until desired position is reached.

Fig. Foot Switch Lateral Tilt

11.4.7 Quick-Action Panels

The Quick-Action panels integrated in the head sections of the siderails allow the nursing personnel and the patient to adjust the bed height.



Fig. Quick-Action Panel

- Buttons Bed Height Adjustment
- 2. GO Button

1.

Set position as follows:

- Activate the keypad by pressing the GO button.
 - Press and hold function button until desired position is reached.

11.5 LCD Touchscreen

The LCD touchscreen is a part of the Multiboard integrated in the head siderail. Depending on the current function, the LCD touchscreen shows different screens. Every screen will display a status bar in the top and a menu bar in the bottom. The status bar shows date and time. The menu bar allows selecting other screens. Green strips above individual icons in the menu bar indicate the active functions in the respective screens.

11.5.1 Menu Bar Icons

| •••• | Positioning Screen Icon |
|------|---|
| 0 | Lock Screen Icon |
| | Scales Screen Icon |
| للج | Bed Exit Monitoring Screen Icon |
| | Transport Mode Screen Icon |
| | Automatic Lateral Therapy Screen Icon |
| | Mattress Screen Icon (Symbioso or OptiCare integrated mattresses) |
| | Help Screen Icon (Screen with instructions for user) Help Screen shows instructions concerning a screen set before Help Screen Icon was pressed. If Settings Screen was set before Help Screen an introduction of the Help Screen is displayed. Use arrows to move between pages of the Help Screen. |
| 000 | Settings Screen Icon |



11.5.2 Positioning Screen



WARNING!

Risk of injury or patient falling out of bed due to lateral tilt!

- Ensure that siderails on the respective side are folded up.
 - Ensure that the patient will not fall out of the bed.

Positioning Screen allows setting certain special positions of the bed and indicates the tilt angle. Bed positioning which depends on columns is continuous.



- History of Backrest adjustment Icon
- Mobilisation Position Icon
 - Notification of expired servicing period
 - Lateral Tilt Icons
- Trendelenburg Position Icon
- Date
- Backrest Angle Indicator
- Autocontour Icons
- Longitudinal Tilt Angle Indicator
- Settings Screen Icon
- Lateral Tilt Angle Indicator

Fig. Positioning Screen

Setting positions:

- Activate touchscreen by pressing GO button.
- Press and hold respective icon until desired position is reached.

The respective indicator indicates the tilt angle or the backrest angle

NOTE Selecting the lateral tilt is exclusively possible if the siderail on the respective side is folded up.

Possible positions:

- Lateral Tilt
- $\hfill \Box$ Allows optimizing the pulmonary function.
- Prevents decubitus.
- Tilting the mattress platform to the left or the right.
- Autocontour
- Raising or lowering the backrest and thighrest.
- Mobilization position
- □ Makes it easier for the patient to get out of the bed.
- Backrest upright
- Mattress platform in lowest position
- Trendelenburg position
- Provides anti-shock conditions for the patient.
- Backrest 30°
- Provides optimum conditions for easier patient ventilation.

NOTE: During continuous positioning Backrest stops automatica¬IIy in 30 and 45 degrees. To continue in positioning press corresponding button once more.

Notification of expired servicing period:

Flashing "Clock and spanner" symbols in the upper left corner mean that the recommended period for safety check has expired. Contact your servicing specialist and plan the next safety check.

INET

The history chart for back part positioning shows:

- the date
- time spent in position at an angle of at least 30° over the last 24 hours
- time spent in position at an angle of at least 45° over the last 24 hours

11.5.3 Lock Screen

Lock allows locking all or individual positioning functions.



- Thighrest Lock Icon
- Height and Tilt Lock Icon
- Backrest Lock Icon

1.

2. 3.

4.

5.

- Foot Control Lock Icon
- Central Lock Icon

Fig. Lock Screen

Locking individual positioning functions:

- Activate touchscreen by pressing GO button.
- Press icon or icons of functions to lock.
- Selected functions are locked. Icons of locked functions are highlighted in yellow.

A light green line appears above the Function Lock icon in the Menu Bar.

Locking all functions:

- Activate touchscreen by pressing GO button.
- Press Central Lock Icon 5.

All positioning functions are locked.

11.5.4 Settings Screen

Settings Screen allows setting the following parameters: Date, Year, Time, Language, New Patient



Weight Value Hidding Icon (ON/OFF) 1. 2. **Day Setting Icons**

- 3. Month Setting Icons
- 4. Year Setting Icons
- Hour Setting Icons 5. 6.
 - Minute Setting Icons
- Date & Time Format Icons (24 hours : minutes 7. + Day. Month. Year OR 24 hours : minutes + Month. Day. Year OR 12 hours : minutes (with AM or PM) + Month. Day. Year)
- 8. Language Icon (language of Help Screen) Mattress OFF Icon (to log off an integrated 9.
- mattress)
- New Patient Icon 10.
- Touchscreen Sound Response Icon (ON/OFF) 11.

Fig. Settings Screen

Setting date, year or time:

Press corresponding icon (2 or 3 or 4 or 5 or 6).

Setting language of the Help Screen:

Press icon 8 repeatedly until desired language is displayed.

New patient:

It is recommended to use New Patient function when replacing patients. The New Patient function is enabled when the bed is loaded with at least 35 kg.

Button performs following functions:

- Scales Taring (only if the scales are stabilized)
- Deletes Scales history
- Deletes ALT history
- Deletes History of Backrest adjustment
- Sets CLP with level 2 (only Symbioso)
- Deletes Pressure Comfort Setting (only OptiCare)
- Turns ON MCM
- Turns ON Fowler Boost function (only Symbioso)
- Turns OFF Sleep Mode (only Symbioso)
- Sets MCM with HIGH level (only OptiCare)

Use the function as follows:

- Wait until the scales are stabilized.
- Press GO button.
- Press and hold icon 10 until it turns yellow.
- The Scales display is shown on LCD.

Scales have been tared, history deleted and it is possible to place new patient on the bed.

Logging out the mattress:

When replacing the integrated mattress (OptiCare or Symbioso) by a standard mattress, it is necessary to log out the integrated mattress.

Press and hold Mattress OFF Icon 9 until vertical bar timer runs to zero and mattress icon disappears.

NOTE Logging out OptiCare mattress or Symbioso mattress to use a standard mattress instead will disable the Manual CPR pop-up menu.

11.5.5 Help Screen

Help Screen shows instructions for users.

Help Screen is available in Czech, English, Italian, French, Spanish, Swedish, Dutch, Brazilian Portuguese, Finnish, Danish and German. Use Settings Screen to set a required language.

11.5.6 Pop-Ups

| Pop-Up | Meaning | Required Action |
|---------|--|---|
| θ | Function locked. | Unlock function if required! |
| L X-RAY | Incorrectly inserted X-Ray Cassette Holder. | Insert X-Ray Cassette Holder correctly! |
| ٢ | GO Button not activated. | Press GO Button! |
| | Lateral Tilt disabled when siderail folded down. | Raise siderail up to enable additional Lateral Tilting. |



| \$2-4-2 1 | Positioning blocked to avoid collision of the bed with floor or collision of the bed with bed equipments. | To continue in positioning, adjust the bed so that there is no collision. |
|-----------------------|---|---|
| └── ○ 0° - ••• | Horizontal position was reached during tilting. | Press corresponding button to continue in positioning. |
| 15° 0' | Lateral Tilt stopped in 15° when Patient Transfer Mode is activated. | For information only. |
| ••• 15° •• 0° | Maximum Lateral tilt was adjusted by foot control (15°). | For information only. |
| 30° 0° | Lateral Tilt limited to 15 degrees when Backrest Angle is in 30 degrees or more. | Lower the Backrest to continue in Lateral Tilting. |
| Kg/lb | Safe Working Load exceeded (more than 10 kg over Safe Working Load). | Remove load! |
| 2 15° | Maximum Lateral Tilt 15 degrees (Load more than 150 kg). | Remove load to enable Lateral Tilting again! |
| kg/lb O° | Lateral Tilt disabled (Load more than 200 kg). | Remove load to enable Lateral Tilting again! |
| * * | System Control Unit (OptiCare or Sym- bioso) disconnected. | Check that bed is connected to the mains power and contact service department approved by manufacturer if the bed with Symbioso integrated mattress is conne- cted to the mains power and this pop-up remains. |
| | The mattress deflating failed. | Use manual CPR! (System Control Unit of the integra- ted mattress is disconnected.) |
| | System Fatal Error. | Contact service department approved by manufactu- rer. |
| | Overwrite confirmation / Confirmation of changes to settings. | Select tick (✓) for "yes" or cross (×) for "no". |
| BED EXIT IS OFF | Scale module disconnected and Bed Exit monitoring disabled. | Contact service department approved by manufactu- rer. |



| * ** | Antitrendelenburg Tilt and Trendelenburg Tilt disabled during Lateral Tilt. | Undo the Lateral Tilt to continue with Antitrendelen- burg Tilt or Trendelenburg Tilt. |
|----------------------|--|---|
| | ALT can´t be running from the accumu- lator. | Connect bed to the mains. |
| <u> </u> | ALT Error. | Contact service department approved by manufactu- rer. |
| 4. 4. 4. 4. | Column Unit Error. | Contact service department approved by manufactu- rer. |
| OPEN | The mattress deflating failed. | Open manual CPR to deflate the mattress! (Automatic deflation is not available.) |

12ALT (Automatic Lateral Therapy)



WARNING!

Risk of injury due to lateral tilt!

- Ensure that the tilting bed does not interfere with the functioning of cannulas, intubation tubes etc.
- Ensure that the tilting bed does not collide with any objects.
- ► Interrupt ALT immediately if the patient's condition worsens, a device or the bed is damaged or any risks to the patient are detected.

ALT allows tilting the mattress platform in order to optimise the patient's pulmonary function and prevent decubitus. Speed of the ALT cycle minimalizes the shock effect and is in accordance with patient comfort.

Before starting ALT:

- Ensure that siderails are folded up.
- Always use LINET ® stabilising ALT pads for positioning patient in centre of bed. (see Stabilising ALT Pads)
 - Always use LINET ® tube holder to prevent extubation (see Ventilation Circuit Holder).
- Ensure that IV lines, breathing tubes etc. are not obstructed and work correctly.
- Reset bed to initial position.

12.1 Resetting Bed to Initial Position

Resetting the bed to its initial position helps to avoid collisions of movable bed parts.



To adjust the bed initial position:

Activate touchscreen by pressing GO button.

Press and hold the single icon on the screen until the Initial Position is reached.

Once the initial position is reached, the bed will stop moving automatically.



- ALT history Icon
- Cycle Setting Time Icons
- Cycle Setting Angle Icons
- Cycle Counter

1.

2.

3.

4.

5.

6.

- TEST Icon (Activation)
 - Backrest Angle Indicator (over 30°)

Fig. ALT Screen - Cycle Values Definition and Test

Defining values of ALT cycle:

- Activate touchscreen by pressing GO button.
- Set time value by pressing and holding one of icons 2 until desired time value is reached.
- Set angle value by pressing and holding one of icons 3 until desired angle value is reached.

NOTE It is possible to change the time values in 5-minute steps to up to 30 min. It is possible to change the angle values continuously up to 30 degrees.

12.2 Test ALT cycle

The ALT test cycle is obligatory and serves to prevent risks such as collisions of moving bed parts, extubation of the patient or disconnection of the ventilation circuit or the cannulas.

During the test cycle, the bed goes through every defined ALT angle and stops at every defined angle level.

Perform ALT test cycle:

- Press icon 5 to start test cycle.
- ▶ Hold icon 5 until test cycle is finished.

Acoustic signal sounds.

Icon Start appears instead of icon Test 5.

12.3 ALT Cycle

After performed test the TEST Icon becomes the Activation Icon to start ALT Cycle.

During an ongoing ALT Cycle the Activation Icon becomes the STOP Icon to stop the ongoing ALT Cycle. During the ongoing ALT Cycle a countdown to the next ALT movement is displayed on the left from the STOP Icon.



Fig. ALT Screen – Stop Icon

To stop cycle:

Press STOP Icon.
Cycle is stopped.

ALT countdown:

The ALT function comes with an automatic countdown of the time remaining until the next lateral bed movement. This is convenient for planning any necessary activities with the patient.



| | | 11:31 | 04.09 | .2010 |
|----|-----------|--------|----------|-------|
| | Date | Time 🔥 | Cycles 🛝 | |
| 03 | .03. 2000 | 00:00 | 3 | |
| 04 | .03. 2000 | 03:00 | 43 | |
| | | | | |
| | | | | |
| | | | | |
| | | | | |
| | | | | _ |
| | | | | _ |
| | | | | _ |
| | | | | |

Fig. ALT history screen

13 Scales Control

13.1 Scales Screen (WS17)

Multicare is equipped with a weighing system that allows weighing the patient in bed. The control and display elements for the weighing system are on the Scales screen on the LCD touchscreen.



- 1. History Subscreen Icon
- 2. Primary display Current Weight Value
- 3. Stabilized Scales Status Icon
- 4. Secondary display Weight Value

Change 5. HC

Displaying ALT history:

Press ALT history Icon.

- HOLD Icon
- 6. ZERO/T Icon (zero Secondary display)
- 7. CANCEL lcon

8. ZERO Icon (zero or tare scales - Primary display)

9. Save Weight Value Icon

13.1.1 Preparation

Install mattress and accessories to prepare bed before patient admission and using the scales.



CAUTION!

Incorrect use of scales due to incomplete preparation!

Before each patient admission tare the scales.

13.1.2 Taring

Taring can be done in range of 5kg to 249,5 kg. Taring is used to set "0" on the display before placing the patient on the bed. Taring must be done with an unloaded bed with mattress, bed sheets, pillows and necessary accessories, without patient. It is recommended to position mattress platform about 8 in. above the lowest horizontal position.



To tare weight:

- Ensure that nothing and nobody touches the bed except you.
- Press and hold icon 8 (ZERO) for 0.5s. The secondary display starts blinking. Hold the icon for another second until the primary display starts to blink.
- Press icon 8 to confirm taring. "0" is shown on both displays. Place the patient on the bed.

To cancel taring:

Press icon 7 while taring.

13.1.3 Displaying

Primary display with Current Weight Value **2** displays the calibrated and metrological certified weight value. Secondary display with Weight Value Change **4** shows the weight difference as compared to the last ZERO or HOLD setting.

Verification Scale Interval is 0.5 kg.

13.1.4 Hold Mode

Hold mode must be used only when the scales are stabilized (icon **3** is shown on the display). It allows adding or removing bed accessories and other items without changing the weight value.

To activate Hold mode:

- Wait until the scales are stabilized. The icon 3 will be illuminated when the scales are stabilized.
- Press icon 5.
- The display shows "HOLD".
- Add or remove required accessories.

To deactivate Hold mode:

- After adding or removing accessories wait until the scales are stabilized. When the scales are stabilized the icon 3 is illuminated.
- Press button 5.
- Display shows the original weight.
- To deactivate Hold mode without fixing the weight value:
- Press button 7.

13.1.5 Saving Weight of Patient

Icon 9 allows saving one particular weight value of patient each day. Saved value will be displayed in History Subscreen.

To save weight of patient:

- First save of the day:
 - Press button 9.

Second (and all next) save of the day:

- Press button 9.
- Confirm saving of the weight on the pop-up by pressing green tick.

-or-

• Deny saving of the weight on the pop-up by pressing red cross.

13.1.6 History Subscreen

To open History Subscreen:

Press icon 1



Fig. Pop-up (Confirm or Deny Saving)





- Unit of Weight Icon (kg / lbs) 1.
 - Scales Screen Icon Graph of Measuring History
- 3.

To return to Scales Screen:

Press icon 2.

2.

Fig. History Subscreen

13.1.7 Bed Overload

If the bed load is over 550 lbs.:

"Hi" is shown on the display.

NOTE: If the bed is overloaded it is impossible to position or manipulate the bed until overloading is removed.

NOTE: Bed overloading always has higher priority than HOLD Mode and Taring functions.

13.1.8 Bed Underload

If the bed is underloaded (factory zero - 5kg.):

Display shows "Lo".

13.1.9 Weighing in tilt

The bed can be weighed in tilt. Accuracy is guaranteed by the spirit level, which is located under the head board of the bed. If the bubble is in the highlighted circle then weighing is accurate.

13.1.10 Zeroing Scales

Zeroing is only possible in a range of ± 5 kg. Zeroing is used to reset weight on the display and set up user zero, which sets the maximum weight range of the scales system.

Zeroing must be done with an empty, unloaded bed, without the mattress and accessories. Zeroing is done after installation, weight verification or servicing.

To zero scales:

- Remove all accessories and the mattress from the bed. Position the bed about 8 in. above the lowest
- position and the mattress platform in the horizontal position. Ensure that nothing and nobody touches the bed except you. Press icon/button 8 (ZERO) for 0.5s. Both displays will start to blink.
- Press icon 8 to confirm zeroing.
 - "0" is shown on the display and an acoustic signal confirms zeroing.

To cancel zeroing:

Press icon 7 while zeroing.

NOTE This function is designed only for metrological certified calibration in recommended periods.

14 Bed Exit Monitoring

Multicare bed is equipped with a Bed Exit Monitoring system that monitores patient's presence in bed and triggers alarms when patient is not present in bed in ordered position. Use Multiboard display to control the Bed Exit Monitoring.

14.1 Preparation

- Place a patient on the bed with suitable mattress.
- Place the patient towards the middle of the bed for the correct function of the Bed Exit Monitoring in Inner Zone.

14.2 Activation



- 1. Notification of Not Loaded Bed
- 2. ON Icon (activation)
- 3. Bed Exit Monitoring Screen Icon

Fig. Bed Exit Monitoring Screen (OFF)

To enter Bed Exit Monitoring Screen:

Press Bed Exit Monitoring Screen Icon (3).

To activate Bed Exit Monitoring:

Press ON Icon when patient is on the bed.

Bed Exit Monitoring Screen is displayed with Bed Picture and Patient Picture.

If you press ON Icon without patient on the bed, Notification of Not Loaded Bed (1) is displayed and the Bed Exit Monitoring is not activated. Minimum patient weight for Bed Exit Monitoring is 35 kg.



14.3 Control of Bed Exit Monitoring



| 1. | Bed Picture |
|----|--------------------------|
| 2. | Inner Zone Icon |
| 3. | Outer Zone Icon |
| 4. | ON Icon (activation) |
| 5. | OFF Icon (deactivation) |
| 6. | PAUSE Icon |
| 7. | VOLUME Icon |
| 7. | VOLUME Icon |
| 8. | Volume Levels (3 levels) |
| 9. | Patient Picture |

Fig. Bed Exit Monitoring Screen (ON)

14.4 Monitored Zone

Bed Exit Monitoring provides monitoring of Inner Zone or monitoring of Outer Zone. Inner Zone covers the mattress support platform without margins around siderails, head board and foot board. Outer Zone covers the mattress support platform up to the margins. Monitoring of Inner Zone is set by default.

To set monitoring of Outer Zone:

Press Outer Zone Icon (3).

To set monitoring of Inner Zone:

Press Inner Zone Icon (2).

14.5 PAUSE

During PAUSE mode the Bed Exit Monitoring is temporarily interrupted and alarms are not activated. PAUSE period is terminated automatically and the Bed Exit Monitoring is reactivated again when patient returns just to the selected zone.

To PAUSE Bed Exit Monitoring:

Press PAUSE Icon (6).

PAUSE Icon turns yellow and countdown timer (15 min) appears on the display above the PAUSE Icon. After terminated PAUSE period when patient is in ordered position the Bed Exit Monitoring is reactivated again.

To extend the PAUSE period:

Press PAUSE Icon (6) again to extend the countdown to 15 minutes period again.

- To terminate the PAUSE period:
- Press ON Icon (4).



14.6 ALARM

Audible alarm is triggered when patient has left selected monitored zone or when PAUSE period is terminated and patient is not just in the selected zone. During this alarm a text "BED EXIT ALARM" in a red rectangle is displayed on the Bed Exit Monitoring Screen. Red strip is displayed above the Bed Exit Monitoring Screen Icon during this alarm.

To stop Alarm:

Press OFF Icon (5).

Bed Exit Monitoring is deactivated and text "BED EXIT IS OFF" in a yellow field is displayed on the screen. The audible alarm is muted.

To pause Alarm:

Press PAUSE Icon (6).



Fig. Bed Exit Alarm (alert)

Countdown timer (15 min) appears on the display above the yellow PAUSE Icon. The audible alarm is muted. If the audible Bed Exit Alarm is paused it is not possible to restore the Bed Exit Monitoring with ON Icon (4). In the case ON Icon is pressed a long beep sounds and PAUSE period continues.

Alarm Volume

It is possible to set Alarm Volume before and during triggered audible alarm. Maximum Alarm Volume Level is set by default. It is not possible to mute completely the audible alarm with this volume setting.

To lower Alarm Volume Level:

Press Volume Icon (7).

Picture with lower Alarm Volume Level (8) appears on the display. Volume is lowered.

To return to Maximum Alarm Volume Level:

Press Volume Icon (7) after Minimum Alarm Volume Level has been reached.

Picture with the 3 volume levels appears on the display.



Fig. Volume Icon with 3 Volume Levels

| 1. | Minimum Volume |
|----|-----------------|
| 2. | Moderate Volume |

3. Maximum Volume

14.7 Deactivation

To deactivate Bed Exit Monitoring:

Press OFF Icon (5).
Text "BED EXIT IS OFF" in a yellow field is displayed on the screen.

14.8 Fault Status (disconnected from the mains)

When the bed is disconnected from the mains the Bed Exit Monitoring system is out of order. If Bed Exit Monitoring is turned on and the bed is disconnected from the mains a fault status appears with text "BED EXIT IS OFF" in a red rectangle. During this Fault Status a beeping sounds for 30 seconds and a louder audible alarm follows.

To mute the audible alarm:

Press the icon on the right side of the red rectangle displayed during this fault status.

To remove this fault status:

Connect the bed to the mains again.



Fig. Bed Exit is OFF (disconnected from the mains)



15 Patient Transfer

This setting allows transferring the patient from the bed to a stretcher or another bed by tilting the bed laterally while the siderails are lowered.



Fig. Patient Transfer Screen

Activating patient transfer setting:

► Press Tick Icon (✓).

Deactivating patient transfer setting:

Press Cross Icon (*).

Transferring patient:

- Fold down siderail.
 - Position stretcher or other bed beside bed.
- Slide transfer board or other transfer aid under patient.
- Activate patient transfer setting.

Maximum load of 5 kg (11.02 lbs) without leverage Maximum load of hook pair 10 kg (22.05 lbs)

Accessories for hanging on the accessory rail:

Urine bag holder Redon bottle basket Stainless steel rails

Move patient from bed to stretcher or other bed.

NOTE The patient transfer setting is deactivated automatically after 3 min. It is possible to reactivate the setting.

Load capacity:

16 Equipment

16.1 Accessory rails



Fig. Accessory Rail

16.2 i-Brake® (optional)

It is possible to equip the bed with an automatic castor brake. The automatic castor brake prevents injuries of pati- ents and staff due to an unbraked bed.

The brakes are activated automatically 60 seconds after the bed is plugged in, and 60 seconds after they have been released if the bed is not being moved.

It is possible to activate the brakes manually as well.

16.3 Retractable 5th wheel i-Drive® (optional)

It is possible to equip the bed with a 5th wheel in the chassis centre. The 5th wheel helps to steer and manoeuvre the bed in long corridors and small rooms.

If the bed is plugged in, the 5th wheel automatically retracts. In this position, the 5th wheel does not obstruct access to any devices under the chassis.

To activate the 5th wheel i-Drive®:

- Disconnect the bed from the mains.
- Adjust the castor control so that the green lever points down



16.4 Mobi-Lift®

Mobi-Lift® is optional. It serves as a support handle to enhance the patient's safety when getting up. Mobi-Lift® is a support handle with a built-in height adjustment button. It allows the patient to raise and lower the mattress platform.



Fig. Mobi-Lift® Support Handle

16.4.1 Using the Support Handles



WARNING!

Risk of injury due to slipping or falling when standing up!

- Ensure that the support handles are completely inserted in the sleeve fittings.
- Ensure that no bed linen is caught between the sleeve fitting and the support handle.

To adjust the support handle:

- Lift the handle up towards the bed.
- Push the handle into the sleeve fitting as far as it will go.

To adjust the height of the mattress platform:

- Press GO button on any control element.
- Press the button to adjust the height.

16.5 Safety Night Light

It helps the nursing staff as well as the patient to orientate.

NOTE The night light is turned off during accumulator operation.

16.6 LINIS SafetyPort (optional)

LINIS SafetyPort is a medical device data system for capturing and transferring data from LINET beds into SafetyPort Dashboard and third party systems, including nurse calls, EHR and digital whiteboards. Data collection and evaluation takes place at one central location for all beds connected to the system simultaneously. The records are completely anonymous and the system does not work with any personally identifiable information.



The customer can decide which data will be sent to the 3rd party system and adjust their sending period. LINIS SafetyPort is intended to be used to increase efficiency of healthcare personnel workflows by saving their time spent on documentation and eliminating errors. This is achieved by automated recording of different parameters of medical beds and their subsequent transfer to various hospital systems in HL7 format. Optional feature LINIS SafetyPort Dashboard is intended to save time the healthcare personnel spends on checking different beds at their workspace and to provide them with both near-real-time data and their aggregation to be able to check the history of provided care. LINIS SafetyPort may be used in various healthcare environments, including both intensive and non-intensive care units as well as units providing speciality care to a broad population of patients. The product is intended to be used by variety of healthcare personnel who have the cognitive skills to operate the product and are trained to use the product. LINIS SafetyPort is not an alarm system and the use of this product for this purpose means incorrect use.

16.7 Nurse Call



WARNING!

Functions of the Nurse Call system depends on the local hospital information system!
Ensure the Nurse Call system is compatible with local hospital information system!



CAUTION!

Ability to use the Nurse Call correctly increases patient's safety!
Hospital staff should inform patient how to use the Nurse Call system!

Intended use

Nurse Call system is intended for sending signals from the bed to hospital staff. Nurse Call system can be used by hospital staff and by patient.

Positions of the Nurse Call control elements

The buttons for activating the Nurse Call function are located on the inner and outer sides of the head siderails. Speakers and microphones are located on the inner sides of the head siderails.



Fig. Nurse Call Control Elements (inner side of foot siderail)



Fig. Nurse Call Control Elements (inner side of head siderail)

Activating the Nurse Call function:

Press Nurse Call button (1).

The patient can speak into the microphone (2) located on the inner side of the head siderails.

16.8 i-Drive Power (optional)

16.8.1 i-Drive Power System - Basic Description

It is possible to equip the bed with the i-Drive Power wheel. The i-Drive Power helps hospital staff to drive the bed during patient transport with minimal manpower.

The i-Drive wheel is located in the center of the bed under the undercarriage. i-Drive Power is equipped with its own accumulator and charger and it is not dependent on the bed functions so, if discharged you can still use the bed functions. The bed is equipped with one i-Drive controller. i-Drive is oriented in straight direction of the bed.

16.8.2 Safety instruction for i-Drive Power

- Follow the instructions carefully.
- Ensure that the bed is operated exclusively by qualified staff.
- Make sure the siderails are pulled up during the transport.
- Never use bed positioning buttons during transport.
- Never use Fast forward button when descending. The Fast forward button is recommended for use when ascending as it is more efficient.
- Special precaution need to be considered when reversing. Always keep distance from the bed and never use reverse button when descending or ascending.
- Do not use Free Drive to transport on a slope greater than 1 degree unless adequate personnel are available to manage safe bed transport.
- Never use the i-Drive Power to drive the bed up or down the slope that exceeds 6 degrees.
- Never leave the bed with an activated i-Drive Power system without supervision of the trained staff.
- Always use the regular mechanical brake system to brake and stabilize the bed.
- Pay increased attention when driving the bed using i-Drive Power. Be aware of people and objects in close proximity and avoid collision with them by careful driving, especially by appropriate speed control.
- Make sure the bed is unplugged and bed brakes are released before using i-Drive Power.
- Push the emergency stop drive button if immediate movement interruption is needed (e.g. to avoid collision with other persons or objects).
- Retract the i-Drive Power wheel to the undercarriage when parking. This will prevent misuse when unbraking and braking the bed.
- The i-Drive Power electromagnetic brake is designed just for temporary bed stop and not for the permanent parking.
- Switch off the i-Drive Power accumulator prior to long-term storage or transport (see chapter 6.1).
- Push the emergency retraction button under the chassis cover to retract the i-Drive Power wheel
- in case an of i- Drive Power system failure. This will enable moving the bed to a safe area manually without using i-Drive Power.
- Retract the i-Drive Power wheel to the undercarriage every time you intend to move the bed sideways.
- Pay attention to the LED accumulator status indicator and plan your drive using the i-Drive Power accordingly. Insufficient accumulator capacity can cause unexpected complications and risks during the drive.
- Always plug the bed in when you finish your drive in order to recharge the accumulator and keep your bed ready to go using the i-Drive Power.
- The i-Drive Power accumulator must be replaced every 2 years to maintain proper functions of the i-Drive Power.

16.8.3 Specifications of Use



WARNING!

Risk of injury due to careless driving!

- Always drive safely and carefully.
- Observe the path for any obstacles and avoid collisions.
- Ensure there are no people in your way.
- Manipulate with the bed carefully not to drive over any staff or patients.



CAUTION!

Maximal clearance underneath the bed (with 15 cm castors) is 14 cm!

Observe the path for any obstacles and avoid collisions.

Intended use:

bed transport (with or without patient)

Unintended use:

- riding the bed
- other usage than described in instructions for use



NOTE Each bed can transport only one patient at a time and cannot be used to transport other items (except bed accessories in secured position).

NOTE For information concerning uses other than those outlined in the "Specifications of Use" section above, please contact LINET ®.

16.8.4 Manipulation



CAUTION! Damage to i-Drive Power main control panel cable due to wrong cable placement!

Ensure that the main control panel connecting cable (13) is placed exactly as on the following picture.



CAUTION! Material damage due to incorrect use!

Do not hang anything on the main control panel and its cable!









- 1. Safety Sense (touch sensor)
- 2. Main control panel
- 3. Activation panel
- 4. GO indicator
- 5. Fault indicator
- 6. Stop drive button
- 7. Fast forward button
- 8. Forward button
- 9. Reverse button
- 10. Accumulator status and fault indicator
- 11. i-Drive wheel Activation button
- 12. i-Drive wheel Retraction and Deactivation button
- 13. Main control panel cable correct cable placement



NOTE The i-Drive Power controller cannot control the bed functions. Control the bed using the bed control elements.

NOTE The main control panel is enhanced with a touch sensor (1); your hand must always be in contact with the i-Drive Power control panel to use the functions. If released, the i-Drive Power will stop.

NOTE Raising and lowering of the i-Drive wheel is electrically controlled by the i-Drive activation panel.

16.8.5 Powered Drive



CAUTION!

- Damage to property due to incorrect transport and involuntary movement!
- Prior to transport, ensure that the bed is disconnected from the mains.
- Prior to transport, ensure that the auxiliary outlet plug (if available) is disconnected from the mains.
- Ensure that the castors are locked prior to assembly, disassembly and maintenance (e.g.: i-Drive Power maintenance).
 - Ensure that the castors are locked when the bed is occupied.
 - Hang the mains cable on the appropriate hook on the bed during transport.
- 1. Check, if the mains switch of i-Drive Power is activated.
- 2. Press the ON (11) button on the Activation panel. The i-Drive wheel will lower and the GO indicator (4) will flash.
- 3. Place your hand on the Safety Sense touch sensor (1) and push the buttons 7 or 8 for forward motion, or 9 for reverse motion. Your hand must be placed on the Safety Sense sensor to use the i- Drive Power, if released, the i-Drive Power will stop.
- 4. The i-Drive motor is immediately stopped and the electric brake is activated after pressing the red stop drive button (6) when braking or in emergency.
- 5. i-Drive Power control system is automatically deactivated and the electric brake is activated if no i-Drive function is used for 3 minutes. This is signalized by the green indicator (4) which is extinguished after 3 minutes.
- NOTE Your hand must be placed on the Safety Sense panel to use the i-Drive Power.
- **NOTE** i-Drive Power is not designed for ascending or descending a slope greater than 6° or longer than 65 ft. (20 m), especially when loaded. The support of personnel is needed when ascending or descending with a full SWL.
- **NOTE** The i-Drive wheel has an electromagnetic brake for emergency or normal stopping of the bed. When parking it is always necessary, for safety reasons, to use the bed brakes (see chapter: Castor control and bed transport) which will brake all four bed castors.
- **NOTE** When i-Drive wheel is lowered, it is not possible to move the bed to the sideways. Press the OFF button to retract the wheel, release the castors to the neutral position and then move the bed to any direction required.

16.8.6 Braking

-or-

-or-

- 1. Press and hold the stop drive button (6) to brake immediately.
- 2. Press and hold the reverse button (9) to brake slowly (Press the Forward button to brake when reversing)
- 3. Release your hand f rom the touch sensor area (1) and i-Drive Power will brake automatically.
- **NOTE** Always brake the bed when not transporting by using the castor control lever. The i-Drive brake is not designed to permanently brake the bed.
- **NOTE** In a crisis situation (e.g. acceleration when driving down a steep slope) i-Drive dual braking prevents acceleration and slows down bed movement. However, it is not guaranteed the bed will stop by itself without personnel support (using stop drive button and castor control lever).
- NOTE When descending, it is possible to actively brake using the opposite direction button to slow.

16.8.7 i-Drive Power Activation/Deactivation





Fig. i-Drive mains switch

To activate the i-Drive Power:

- 1. Check, if the mains switch of i-Drive Power is activated (1).
- 2. Press the Activation button ON located on the activation panel. The i-Drive wheel will lower and the green indicator will flash.

To deactivate the i-Drive Power:

- 1. Retract the i-Drive wheel using the Retraction button located on the activation panel.
- 2. Deactivate the i-Drive using the mains switch (1).

Emergency i-Drive Power wheel retraction:

- 1. Press any GO button on the bed.
- 2. Deactivate the i-Drive using the mains switch (1).
- 3. Press the emergency retraction button (2).
- **NOTE** Use emergency retraction in case of accumulator discharge or drive malfunction to move the bed to a safe area manually without using i-Drive Power.

16.8.8 Free Drive

The i-Drive motor is equipped with free drive, which is active after pressing the forwards (7 or 8) or backwards (9) buttons (until user holds the touch sensor area).

Free Drive is deactivated and the brake is activated when the direction of motion is changed. This is feature for lowering the risks when going to a slope.

16.8.9 Accumulator



Accumulator charge status:

1. While this indicator is flashing, the accumulator is critically dis-

charged. 2.

3.

4.

- 50% 75%
 - 100% the accumulator is charged

Fig. Accumulator indicator levels

To charge the accumulator:

- Connect the bed main cable to mains power.
- i-Drive will be charged (with the accumulator discharged, the charging may take up to 9 hours).

NOTE Accumulator charge values are just informational. Accumulator life is reduced when the accumulator is allowed to discharge completely.



16.8.10 Fault Signalization

The system is protected against failure states, by stopping and braking the drive system, and respective signalization. The fault indicator flashing briefly and the accumulator indicator shows the fault state. Some defects are cleared automatically (e.g.: drive overheating).

| Error | LED1 | LED2 | LED3 | LED4 |
|--|------|------|------|------|
| Drive overheated* | Off | Off | Off | On |
| Electronics overheated* | Off | Off | On | Off |
| Brake error | Off | Off | On | On |
| Retraction not completed | Off | On | Off | Off |
| 5V off limits | Off | On | Off | On |
| FETclosingpenetrated | Off | On | On | Off |
| Control circuit overheated | Off | On | On | On |
| Controlcircuiterror | On | Off | Off | Off |
| Activation button stuck | On | Off | Off | On |
| Retraction button stuck | On | Off | On | Off |
| Activebuttonafterstart | On | Off | On | On |
| * An acoustic signal occurs before the drive is blocked (short acoustic signalization) | | | | |

NOTE LED indicators are numbered from the left.

16.8.11 Light Indicators

| Indicator | Meaning |
|--|--|
| Go Indicator ► Constantly lit ► Flashing | Hand is on touch sensor; drive wheel is ready for use. Hand is not on touch sensor; i-Drive is not ready for use. |
| Fault Indicator | |
| Constantly lit | i-Drive cannot be activated (i-Drive wheel is not lowered, castor control lever is braked, bed is connected to the mains). |
| ► Flashing | |
| | System is faulty (indicated on accumulator status indicator, see service manual) |
| | -or- i-Drive control box heat protection is activated |

16.8.12 Technical Specifications

| Specification | Value |
|---|------------------|
| i-Drive wheel diameter | 8,27 in. |
| Max. fast forward speed (flat ground, loaded) | 4,43 Km/h (±15%) |
| Max. forward speed (flat ground, loaded) | 2,16 Km/h (±15%) |
| Max. reverse speed (flat ground, loaded) | 2,16 Km/h (±15%) |
| Max. angle of ascent | 6° |
| Noise level (when retracting the drive wheel) | 65 dB |

16.8.13 Electrical specification

| Specification | Value |
|-----------------------|--------------------------------------|
| Accumulator Voltage | 36 V DC, Capacity: 12 Ah |
| Maximum Power Input | 300 W |
| Fuse Accumulator fuse | pipe fuse T 3.15 A MDP 030 (30 A) |



16.8.14 I-Drive Power Maintenance

Periodical maintenance of the i-Drive Power must be done by qualified service technician or authorized service organization at least once a year.

Service technician must check the following:

- accumulator status and eventual replacement of accumulator (after maximum of three years of duty)
- gas spring replace if necessary (after maximum of three years of duty)
- ▶ i-Drive Power wheel replace if necessary
- ▶ lifting mechanism grease if necessary
- cables, control elements replace if necessary
- i-Drive Power function

NOTE To continue maintenance please see chapter Maintenance.

16.9X-Ray Lung Examination

The backrest of the bed consists of HPL and is x-ray translucent. The bed is equipped with an x-ray cassette hol- der with 2 U-profiles under the backrest. This design allows taking x-ray images of the patient's lungs without moving the patient manually.



Fig. X-Ray Lung Examination

16.9.1 Necessary Steps before the Examination

NOTE This procedure is suitable for patients who cannot be moved due to critical conditions (e. g. internal bleeding).

- Make sure that patient is in centre of bed.
- Make sure that backrest is in lowest position and siderails are folded up.
- Pull out x-ray cassette holder.
- Insert x-ray cassette (format 43×35 cm (16.93 in. x 13.78 in.)).
- Push back x-ray cassette holder with x-ray cassette so that the cassette centre indicator is exactly under the edge of the mattress platform.
- Correct position of x-ray cassette holder using the tooth mechanism so that the upper edge of the x-ray cassette is exactly under the patient's shoulder line.
- Adjust parameters of the x-ray device.

16.9.2 Examination with C-arm

Backrest and seat of the bed are x-ray translucent. The bed is equipped with a column construction. This design allows C-arm--assisted operations (mainly cardiological operations such as temporary external cardiostimulation) without moving the patient. The x-ray tube of the C-arm is located between the undercarriage and the mattress platform.

Necessary Steps before the Operation

- Make sure that backrest is in highest position and siderails are folded up.
- Position upper part of C-arm (sensor and indicator) above the patient's chest.

16.10 Auxiliary Outlet (120V)



DANGER! Danger to life due to incorrect use!

- Do not use auxiliary outlet for life-sustaining equipment!
- Ensure that total leakage current in the chassis does not exceed 100 µA!
- Use only hospital grade or hospital only medical devices!
- Never repair and never open the mains socket outlet!
 - Call service department instead!



DANGER!

Danger to life due to damaged cables or faulty grounding!

- Do not use damaged cables!
 - Use plastic hooks on head end to secure cables when moving the bed!
- Check grounding regularly!



DANGER!

Danger to life due to incorrect use!

- The mains socket outlet shall be used within its marked load rating 10 A!
- Use the mains socket outlet for supplying equipment which is intended to form part of the medical system only!



DANGER!

Risk of material damage due to objects on the undercarriage cover!

- Operator shall not touch parts of the mains socket outlet and the patient simultaneously!
- Keep the mains socket outlet in factory position and permanently attached to the bed, never remove it. Call service department instead!
- Do not use the bed if the mains socket outlet is located or positioned on the floor! Call service department immediately!

Intended use

An auxiliary power supply outlet for medical devices (hospital grade only) is located under the calfrest on the right side.



Fig. Position of the Auxiliary Outlet (120V)



17 Mattress

Multicare bed is designed for passive and active mattresses from LINET portfolio.



CAUTION!

Incompatibility with bed due to incorrect mattress dimensions!

Check maximum approved mattress dimensions (chapter Technical Specification).

The manufacturer recommends the use of the following mattress systems on the Multicare bed:

PASSIVE MATTRESSES

- CliniCare 10
- CliniCare 20
- CliniCare 30

ACTIVE MATTRESSES

- Virtuoso (not integrated)
- Symbioso (integrated)
- OptiCare (integrated)

17.1 Passive Mattress

Recommended Passive Mattresses are equipped with straps (1) intended for fixing mattress on the Mattress support platform.



Fig. Bottom of Passive Mattress

17.1.1 Straps with side release buckles

To fix mattress on the Mattress Support Platform:

- Run three straps through the three corresponding holes in the covers of Mattress Support Platform.
- Run these three straps under the bars of the Mattress Support Platform.
- Lock the three side release buckles by connecting their male and female parts together.

To remove mattress from the Mattress Support Platform:

- Release the three buckles by pressing them from both sides and by disconnecting their male and female parts.
- Pull these three straps out of the Mattress Support Platform.
- Remove mattress from the Mattress Support Platform.



17.2 Active Mattress (not integrated)



WARNING!

Follow instructions for use of a compatible active mattress carefully!



CAUTION!

Risk of material damage due to an incorrect fixation of compatible active mattress on the mattress support platform!

► Adjust the bed to maximum Cardiac Chair Position before fixing all the straps of the inflated mattress to the mattress support platform!

Installation instructions:

- Remove any existing mattress.
- ▶ Observe mattress dimensions and its orientation before putting it on the Mattress support platform.
- ▶ Place SCU on the foot board of the bed or on the floor.



Fig. Bottom of Active Mattress (not integrated)

17.3 OptiCare and Symbioso (integrated mattress)



WARNING!

Follow instructions for use of the OptiCare integrated mattress replacement system or instructions for use of the Symbioso integrated mattress replacement system carefully!



CAUTION!

Risk of material damage due to an incorrect fixation of compatible integrated mattress on the mattress support platform!

► Adjust the bed to maximum Cardiac Chair Position before fixing all the straps of the inflated mattress to the mattress support platform!



WARNING!

OptiCare mattress and Symbioso mattress are compatible with System Control Units delivered by manufacturer only!

Do not use any other System Control Units with OptiCare mattress or with Symbioso mattress!



CAUTION!

Material damage due to incorrect installation of SCU!

If the SCU does not come factory-fitted, have it installed by a service engineer authorised by LINET ®.



18 Accessories



WARNING!

Risk of injury due to incompatible accessories!

Use exclusively original accessories from the manufacturer.

The manufacturer is not responsible for the use of unapproved accessories.



WARNING!

Risk of injury due to damaged accessories!

Use exclusively accessories in perfect condition.

18.1 Lifting Pole

To ensure safe use of the lifting pole:

- Never exceed the maximum load of 75 kg (165.35 lbs).
- Never use the lifting pole for rehabilitation exercises.
- To prevent the bed from tipping over, ensure that the lifting pole does not project out from the bed.
- Replace plastic handle every 4 years.

To install the lifting pole:

- Insert lifting pole in corresponding sleeve fitting on lifting pole adapter at head end.
- Ensure that safety pin locks into place.

A plastic grab handle with an adjustable strap shall be attached to the lifting pole.

- **NOTE** The lifting pole adapter is optional. It is necessary to specify this feature in the order.
- **NOTE** The date of manufacture is marked on the grab handle. LINET® recommends replacing the plastic grab handle every four years.



18.2 Infusion Stands



WARNING!

Risk of injury due to use of incorrect accessories or because of incorrect use!

- Infusion Stands must only be used for their intended use. Always read the instructions for use!
- Only mount an infusion pump to the lower (wider) telescopic section of an infusion stand above the head board/ foot board.
 - Never mount an infusion pump to the upper (thiner) telescopic section of an infusion stand.
- Ensure the infusion pump will not collide with any movable parts of the bed (especially backrest part) or with the patient. This must be verified during installation.
- Do not over tighten the infusion pump clamps during fitment. Over tightening may damage the infusion stand.
- Infusion pump can be only used if the infusion stand is fitted in the accessory holder socket in the head end on the undercarriage of the bed.
- Do not use the infusion stand as driving/pushing device during the bed transport.

Infusion stands can be fitted to the head and foot board by either fitting into the IV/Infusion sockets mounted on the bed or using alternative accessory holder socket in the head end on the undercarriage of the bed.

 Use exclusively infusion stands with 4 hooks for hanging IV bags or baskets for intravenous solutions.
Ensure the infusion stand individual hook 2kg maxi-

mum Safe Working Load is not exceeded.

• Ensure the infusion stand 20kg maximum Safe Working Load is not exceeded.

► The total maximum loading of the IV/Infusion poles must not exceed 20 Kg (44.1lbs).





Fig. Infusion Stand

Fig. Infusion Pump – Correct Fitment

18.3 Stabilising ALT Pads



Fig. Stabilising Pads

The stabilising pads ensure a stable position of the patient during ALT in order to prevent extubation or disconnection of IV lines or other equipment.

Stabilising pad set:

- 2 lateral arm pads
 - 2 lateral leg pads
 - 2 head pads
 - 1 internal leg pad
 - Always use LINET [®] stabilising ALT pads for positioning patient in centre of bed during ALT.

Applying pads:

- Position the patient in the middle of the bed.
- Place lateral pads between patient and siderails.
- Attach head pads to arm pads with Velcro.
- Place internal pad between the patient's legs.
- Tilt mattress platform left and right by 30° to check if the patient's position is stable.
- The position is stable if the patient does neither shift nor turn over.

18.4 Ventilation Circuit Holder

The ventilation circuit holder prevents an extubation.

Always use LINET ® ventilation circuit holder to prevent extubation during ALT.



Applying ventilation circuit holder:

Put ventilation circuit holder in hole on right or left of head

- end.Fasten ventilation circuit holder with wing screw provided.
 - Put intubation tube through plastic head of ventilation circuit holder.
 - Tilt mattress platform left and right by 30° to check if intubation tube is fastened securely. The fastening is secure if no parts the ventilation circuit are disconnected.

Fig. Ventilation Circuit Holder

18.5 Monitor Tray

The monitor tray is suitable for transporting monitors with a weight of up to 15 kg (33.07 lbs).



Fig. Monitor Tray

Installing the monitor tray:

Insert two vertical monitor tray tubes into corner sleeves on foot end.

Fixate monitor with safety belts in order to avoid any damage during transport.

18.6 Oxygen Bottle Holders



WARNING!

Risk of injury with oxygen bottle holder due to incorrect use or due to careless driving!

- Ensure the oxygen bottle holder is correctly fitted in correct position.
- It is necessary to place oxygen bottle holder (with or without O2 bottle) before transport to secure transport position.
- Be aware of people or objects in close proximity when driving or manipulating the bed equipped with oxygen bottle holder.
- Secure the oxygen bottles against falling or involuntary movement with rubber strap.
- Place the oxygen bottle holder on the bed by instructions in the following text.
- Ensure the oxygen bottle valve will not get damaged by careless or incorrect manipulation or placement.

The oxygen bottle holders are suitable for transporting oxygen bottles with a weight of up to 15 kg (33.07 lbs) and a volume of 5 litres.



Version A

Put oxygen bottle holder on transversal profile behind head end.

NOTE Using oxygen bottle holder 4MAR2010PC004 is not possible if the bed is equipped with an additional adapter for a lifting pole.

Fig. Oxygen Bottle Holder A

Version B

Version C

- Put holder on sleeve fittings in multifunctional accessory adapter on chassis.
- Ensure the locking pin of oxygen bottle holder B is locked in sleeve fitting.



Fig. Oxygen Bottle Holder B – correct fitment

Put oxygen bottle

holder on all 4 accessory adapters on chassis.



Fig. Oxygen Bottle Holder B – incorrect fitment



Fig. Oxygen Bottle Holder C



18.7 Protector



WARNING! Bisk of injury due to the r

Risk of injury due to the patient falling off the bed!

- Ensure that the Protector is installed securely.
- Always check that the siderails are properly locked.

The Protector is an optional accessory for the Multicare bed. The main purpose of the Protector is to reduce the risk of patients falling off the bed.



Fig. Multicare bed with Protector

- 1. Inserting the Protector into the casing in the protective ring on the corner
- 2. The Protector inserted in the casing
- 3. The fixing element attached to the telescopic profile of the bed extension
- 4. The Protector attached to the Multicare bed (The Protector can also be used on expanded beds.)

Attach the Protector to the bed as follows:

- Insert the Protector pin into the casing in the protective ring at the corner of the foot end of the bed (1).
- Ensure that the fixing element is secured to the telescopic profile of the bed extension (3).

Remove the Protector from the bed as follows:

- Grasp the upper end of the Protector.
- Remove the Protector from the casing.



18.8 Hercules



WARNING! Ensure that Hercules is operated exclusively by qualified personnel.



WARNING!

Hospital staff is responsible for the patient during his or her repositioning. The patient should not be left unattended on the bed during his or her repositioning!



WARNING!

In Backrest angle of 30 degrees or more it is not possible to use Hercules which is indicated by LED on the side of Hercules. Follow the instructions for use of the Hercules!



WARNING! Do not use Hercules without the gas spring securely installed!



WARNING! In order to facilitate CPR Backrest Release push the Backrest down using head siderail!



WARNING!

Risk of squeezing between head siderails and sides of the Hercules! Manipulate carefully with head siderails when Hercules is installed!

Hercules Patient Repositioner is intended for Hercules ready Multicare bed. Hercules is compatible also with a specific configuration of the OptiCare integrated mattress. For detailed information about use of the Hercules with OptiCare integrated mattress follow the instructions for use of the OptiCare. Installation of Hercules must be done by qualified service technician authorized by the manufacturer. For detailed information about Hercules follow the instructions for use of this product.

Purpose:

Hercules is intended to assist caregivers with up-in-bed patient repositioning. Hercules repositioning system shall be used when bed is in horizontal and flat position. Hercules can be used with compatible configuration of the OptiCare integrated mattress only if OptiCare is in MAX Mode.

Description:

Hercules system consists of Hercules Drive, Hercules Dream Sleep Surface / Hercules dream Gel Sleep Surface / OptiCare integrated mattress and Hercules Dream Sheet.

Placement:

Hercules is located at the end of the Backrest.



Fig. Hercules with original Morel mattress


19Cleaning/Disinfection

Antibacterial surface finish:

Selected parts of the Multicare bed are treated against the spread of bacteria with certified technology by Sanitized[®]. This technology supplements regular bed disinfection procedures. Regular bed cleaning cannot be omitted relying only on the antibacterial surface finish. Clean the bed according to the following instructions.



WARNING!

Risk of injury due to accidental bed movement!

Always disable the function buttons when cleaning between the undercarriage and mattress platform.



CAUTION!

Material damage due to incorrect cleaning/disinfection!

- Do not use washing machines.
- Do not use pressure or steam cleaners.
- Exclusively use the recommended cleaning agents.
- Follow the instructions and observe the dosages recommended by the manufacturer.
- Ensure that disinfectants are selected and applied exclusively by qualified hygiene experts.
- Check if used cleaning agents and disinfectants are compatible with materials that the product consists of! For information see the following table.

| BED COMPONENTS THAT ARE INTENDED TO BE CLEANED | MATERIALS (SURFACES OF THE MENTIONED BED COMPONENTS) | |
|--|--|--|
| Do not clean what is not mentioned in this column! | Competent user is responsible for check if used cleaning agents and disinfectants are compatible with mentioned materials! | |
| Head board and foot board | Polypropylene with antibacterial surface (PP) | |
| Head siderails and foot siderails | Polypropylene with antibacterial surface (PP) | |
| Decors (head board, foot board, head siderails, foot siderails) | Acrylonitrile butadiene styrene (ABS) | |
| Mattress support platform cover (Backrest) | High Pressure Laminate (HPL) | |
| Mattress support platform covers (Thighrest, Calfrest) | High Pressure Laminate (HPL) + Lacquered steel | |
| Mattress support platform cover (Seat section) | High Pressure Laminate (HPL) | |
| Covers of frame of the mattress supp- ort platform | Polypropylene (PP) + Acrylonitrile butadiene styrene (ABS) | |
| Frame of the mattress support plat- form | Lacquered steel | |
| Castors | Polypropylene (PP) | |
| Columns | Oxidized aluminium alloy | |
| Siderail release mechanisms | Lacquered aluminium (Al) | |
| Undercarriage cover | Acrylonitrile butadiene styrene (ABS) | |
| Corners and corner covers | Polypropylene (PP) | |
| Handles of head board lock and foot board lock | Polyamide (PA) | |
| Corner bumpers | Polypropylene (PP) | |
| Keyboards (Attendant Control Panel, Handset, control elements integrated in the siderails) | Autotex film with antibacterial surface | |
| CPR levers | Polyamide with antibacterial surface (PA) | |
| Mobi-Lift [®] handles | Polyamide with antibacterial surface (PA) | |
| Accessory rail | Lacquered steel + Polyethylene (PE) | |
| Labels | siliconized paper with lamination or with resin | |

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For safe and gentle cleaning:

- Do not use any strong acids or bases (optimum pH range 6 8).
- Exclusively use detergents that are suitable for cleaning medical equipment.
- Do not use abrasive powders, steel wool, or other materials and cleaning agents that might damage the mattress replacement system.
- Never use any corrosive or caustic detergents.
- Never use detergents that deposit calcium carbonate.
- Never use detergents with solvents that might affect the structure and consistency of the plastics (benzene, toluene, acetone, etc.).
- Clean electrical components carefully and allow them to dry completely.
- Do not immerse SCU in water or steam-clean it.
- Observe local directives regarding infection control.
- Make sure any cleaning agent used is approved by:
- the facility in which the mattress replacement system is to be used.
- by the environmental protection agency of the country in which the mattress replacement system is to be used.

LINET ® recommends the following cleaning agents:

| Parts to be cleaned | Cleaning agents | |
|--|--|--|
| Multicare hospital bed | Mikrozid, Terralin Protect, Thermosept (Schülke & Mayr) Bacillol AF, Bacillol Rasant, Dismozon Pur, Microbac Forte, Neodisher Dekonta (BODE Chemie) Lysoformin 3000, Lysoform Spezial (LYSOFORM) Incidin plus, Incidin rapid (Ecolab) Perform, TPH protect (Schülke) | |
| Mattress cover base, comforter covers, air cells, foam base, SCU | standard hospital detergents alcohol or chlor based desinfections | |
| Mattress cover top | standard hospital detergents alcohol and quaternary ammonium-based disinfectants | |

19.1 Cleaning (Multicare)

Prepare for cleaning as follows:

- Put the mattress platform in the highest position.
- Adjust the back and thighrests so that the reverse sides are accessible.
- Disable the function buttons on the control elements using the Attendant Control Panel.
- Disable the foot controls using the Attendant Control Panel.
- Disconnect the bed from the mains.
- Move the bed to the location where it will be cleaned.
- Lock the brakes on the bed.

19.1.1 Daily Cleaning

Clean the following bed parts:

- All control elements for adjusting the bed
- All handles
- CPR release handle
- Head Board and Foot Board
- Siderails (in highest position)
- Freely accessible mattress surface
- Mobi-Lift®
- Accessory rails

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19.1.2 Cleaning before Changing Patients

Clean the following bed parts:

- All control elements for adjusting the bed
- All handles
- CPR release handle П
- Head Board and Foot Board
- Siderails (in highest position)
- Freely accessible mattress surface ►
- Mobi-Lift® ►
- Accessory rails
- All plastic mattress platform covers
- Plastic undercarriage covers
- Telescopic columns
- Mattress on all sides
- Freely accessible metal parts of mattress platform
- Cable ducts
- Lifting pole sleeve fitting
- Infusion stand sleeve fitting
- ► **Bumpers**
- ► Castors
- Brakes

19.1.3 Complete Cleaning and Disinfection

Clean the following bed parts:

- All control elements for adjusting the bed
- All handles
 - CPR release handle
 - Head Board and Foot Board
- Siderails (in highest position)
- Freely accessible mattress surface
- Mobi-Lift®
- Accessory rails
- All plastic mattress platform covers
- Plastic undercarriage covers
- Telescopic columns
- Mattress on all sides
- Freely accessible metal parts of mattress platform
- Cable ducts
- Lifting pole sleeve fitting
- Infusion stand sleeve fitting
- **Bumpers**
- Castors
- Brakes
- Interior parts
- (accessible after removing mattress platform covers)

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20 Troubleshooting



DANGER! Danger to life due to electric shock!

- If a fault occurs ensure the electric motor, power box and other electrical parts checked by qualified personnel only.
- Do not open protective covers of the electric motor or power box.

| Error/Fault | Cause | Solution |
|--|---|--|
| Adjusting with position buttons not possible | GO button was not pressed | Press the GO button. |
| | Function disabled on Attendant Control Panel | Enable disabled function. |
| | Drive motors have no power, Defective drive motors, Defective accumulator | Check the mains connection. Notify the service department of manufacturer. |
| | Plug inserted incorrectly | Insert the mains plug correctly. |
| | Faulty power source | Notify the service department of manufacturer. |
| | Faulty control element | Notify the service department of manufacturer. |
| Faulty mattress platform height/tilt adjus- tment | There is an object on the undercarriage cover | Remove the object. |
| | Function disabled on Attendant Control Panel | Enable disabled function. |
| | Drive motors have no power, Defective drive motors, Defective accumulator | Check the mains connection. Notify the service department of manufacturer. |
| | Plug inserted incorrectly | Insert the mains plug correctly. |
| | Faulty power source | Notify the service department of manufacturer. |
| | Faulty control element | Notify the service department of manufacturer. |
| Lowering backrest from the upright position not possible | There is an object under the backrest or in the drive mechanism | Remove the object. |
| | Locking handle is defective | Notify the service department of manufacturer. |
| Adjusting siderails not possible | The siderail lock is dirty | Clean the locking mechanism. |
| | Release handle is defective | Notify the service department of manufacturer. |
| Faulty brakes | Dirt blocking brakes mechanically | Clean the brake system. |
| | The brake mechanism is defective | Notify the service department of manufacturer. |



21 Maintenance

WARNING!



Risk of injury when working on the bed!

- Ensure that the bed is disconnected from the mains power prior to installation, putting into service, maintenance and deinstallation.
- Ensure that the castors are locked prior to installation, putting into service, maintenance and deinstallation.



WARNING!

Risk of injury due to defective bed!

- Have a defective bed repaired immediately.
- If the defect cannot be repaired, do not use the bed.



CAUTION!

Material damage due to incorrect maintenance!

- Ensure that maintenance is performed exclusively by manufacturer's customer service or by authorised service personnel certified by the manufacturer.
- If the defect cannot be repaired, do not use the bed.

LINET ® recommends attaching the maintenance plaque to the bed.

21.1 Regular maintenance

- Check regularly movable parts for wear.
- Perform regularly visual check of the product (with delivery note if necessary).
- Ask service department of the manufacturer for addition of the original spare parts if some product parts are missing.
- Ask service department of the manufacturer for replacement of any damaged product parts by the original spare parts.
- Check that the accumulator is working properly. Disconnect the bed from the mains power to check signalisation of accumulator indicator according to the instructions for use.
- Have the accumulator replaced if it is not working properly.
- Check regularly that all accessories are working properly.
- Replace damaged accessories immediately.

21.2 Spare Parts

The serial label is located on the frame of the mattress support platform. The serial label contains information for claims and ordering replacement parts.

Information about spare parts is available from:

- Manufacturer's customer service
- Sales department

21.3 Safety Technical Checks



WARNING! Risk of injury due to incorrect safety technical checks!

Ensure that safety technical checks are performed exclusively by manufacturer's customer service or by authori-

- sed service personnel certified by the manufacturer.
 - Ensure that the safety technical checks are recorded in the service and maintenance log.

Safety technical check of the medical bed must be performed at least once every 12 months.

The procedure for performing the safety technical check is stipulated in EN 62353:2014.

NOTE On request, the manufacturer will provide service documentation (e.g. circuit diagrams, component part lists, descriptions, calibration instructions etc.) for service personnel for the repair of ME equipment designated by the manufacturer as repairable by service personnel.

22 Disposal

22.1 Environment Protection

The company LINET® is aware of the importance of environmental protection for future generations. Within this company the environmental management system is applied in accordance with the internationally agreed standard ISO 14001. The compliance with this standard is annually tested by the external audit executed by an authorised company. Based on the Directive No. 2002/96/ EC (Directive **WEEE** - Waste, Electric and Electronic Equipments) the company LINET, s. r. o. is registered in the List of Electric and Electronic Equipment Producers (**Seznam výrobců elektrozařízení**) on the Ministry of the Environment of the Czech Republic (Ministerstvo životního prostředí).

Materials used in this product are not environmentally hazardous. LINET® products meet valid requirements of national and European legislation in the areas of **RoHS** and **REACH**, so they do not contain any prohibited substances in excess quantities. None of the wooden parts is made of tropical wood (such as mahogany, rosewood, ebony, teak etc.) or made of timber from the Amazon region or from similar rainforests. Product noise (sound pressure level) meets requirements of the regulations for the protection of public health against undesirable effects of noise and vibration in protected interior spaces of buildings (according to standard IEC 60601-2-52). Used packaging materials are in accordance with requirements of the Packaging Act (**Zákon o obalech**). For disposal of packaging materials after installation of products contact your sales representative or manufacturer's customer service about the possibility of a free take-back of packaging through an authorized company (more details on **www.LINET.cz**).

22.2 Disposal

The main objective of the obligations arising from the European Directive No. 2012/19/EU on Waste, Electric and Electronic Equipments (nationally regulated in Act No. 185/2001 Coll. as amended. On Waste and in Decree of the Ministry of the Environment No. 352/2005 Coll. as amended), is to increase the re-use, material recovery and recovery of electric and electronic equipment at the required level, thereby avoiding the production of waste and thereby avoiding the possible harmful effects of hazardous substances contained in electric and electronic equipment on human health and the environment. LINET® electric and electronic equipments that have a built-in accumulator or accumulator are designed so that the used accumulator or accumulators can be safely removed by LINET® qualified service technicians. There is an information about its type on the built-in accumulator or accumulator.

22.2.1 Within Europe

To dispose of the electric and electronic equipment:

- The electric and electronic equipment must not be disposed of as household waste.
- Dispose of this equipment at designated collection points or take-back points.

To dispose of the other equipment:

- The equipment must not be disposed of as household waste.
- Dispose of this equipment at designated collection points or take-back points.

LINET® participates in a collective system with take-back company REMA System (see **www.remasystem.cz/sberna-mista**/). By bringing electric and electronic equipment to a take-back point, you participate in recycling and you save primary raw material resources while protecting your environment from effects of unprofessional disposal.

22.2.2 Outside Europe

- Dispose of the product or its components in accordance with local laws and regulations!
- Hire an approved waste disposal company for disposal!



23 Warranty

LINET ® will only be held responsible for the safety and reliability of products that are regularly serviced, maintained and used in accordance with the safety guidelines.

Should a serious defect arise that cannot be repaired during maintenance:

Do not continue to use the bed.

Warranty duration is subject to individual purchasing agreements with a minimum length of 12 months.

The warranty covers all material and manufacturing-related failures and errors. Failures and errors caused by incorrect use and external effects are not covered. Justified complaints will be fixed free of charge during the warranty period. Proof of purchase, with the date of purchase, is required for all warranty service. Our standard terms and conditions apply.

24 Standards and Regulations

24.1 Multicare

The Multicare bed comlies with the following standards:

- IEC 60601-1
- CAN/CSA C22.2 NO. 60601-1
- ANSI/AAMI ES60601-1
- IEC 60601-1-2
- IEC 60601-1-6
- IEC 60601-2-52
- ISO 14971

24.2 OptiCare and Symbioso

The OptiCare and Symbioso integrated mattress replacement system complies with the following standards:

- IEC 60601-1
- CAN/CSA C22.2 NO. 60601-1
- ANSI/AAMI ES60601-1
- IEC 60601-1-2
- IEC 60601-1-6
- ISO 10993-5
- ISO 10993-10

24.3 Manufacturer

The manufacturer adheres to a certified quality management system in compliance with the following standards:

- ISO 9001
- ISO 14001
- ISO 13485
- MDSAP (Medical Device Single Audit Program)