WARNING
To properly use this medical device, read and comply with these instructions for use.

Babylog VN500
Ventilation unit
SW 2.n
Typographical conventions

1 Consecutive numbers indicate steps of action, with the numbering restarting with “1” for each new sequence of actions.
- Bullet points indicate individual actions or different options for action.
  - Dashes indicate the listing of data, options, or objects.
(A) Letters in parentheses refer to elements in the related illustration.
A Letters in illustrations denote elements referred to in the text.

Any text shown on the screen and any labeling on the device are printed in bold and italics, for example, **PEEP**, **Air** or **Alarms**.

The “greater than” symbol > indicates the navigation path in a dialog window, for example, **System setup > Ventilation > Modes**.

In this example, **System setup** represents the dialog window title, **Ventilation** represents a horizontally aligned tab, and **Modes** a vertically aligned tab.

Screen images

Schematic renderings of screen images are used, which may differ in appearance or in configuration from the actual screen images.

Use of terms

Dräger uses the term “Accessory” not only for accessories in the context of IEC 60601-1, but also for consumable parts, removable parts, and attached parts.

Trademarks

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Safety information definitions

**WARNING**
A WARNING statement provides important information about a potentially hazardous situation which, if not avoided, could result in death or serious injury.

**CAUTION**
A CAUTION statement provides important information about a potentially hazardous situation which, if not avoided, may result in minor or moderate injury to the user or patient or in damage to the medical device or other property.

**NOTE**
A NOTE provides additional information intended to avoid inconvenience during operation.

Definition of the target groups

For this medical device users, service personnel, and experts are defined as target groups.

These target groups have been instructed in the use of the medical device and have the necessary knowledge to use, install, reprocess, maintain or repair the medical device.

Dräger emphasizes that the medical device must be used, installed, reprocessed, maintained or repaired exclusively by defined target groups.

**User**

Users are persons who may use the medical device in accordance with its intended use.

**Service personnel**

Service personnel are persons who are responsible for the maintenance of the medical device towards the operating organization.

Service personnel are persons who may install, reprocess, or maintain the medical device.

**Experts**

Experts are persons who may carry out repair or complex maintenance work on the medical device.
Abbreviations and symbols

For explanations refer to sections "Abbreviations" and "Symbols" in chapter "System overview".
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For your safety and that of your patients

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General safety information

The following WARNING and CAUTION statements apply to general operation of the medical device. WARNING and CAUTION statements specific to subsystems or particular features of the medical device appear in the respective sections of these instructions for use or in the instructions for use of another product being used with this device.

Strictly follow these instructions for use

WARNING
Any use of the medical device requires full understanding and strict observation of all sections of these instructions for use. The medical device must only be used for the purpose specified under "Intended use" on page 18 and in conjunction with appropriate patient monitoring (see page 10). Strictly observe all WARNING and CAUTION statements throughout these instructions for use and all statements on medical device labels. Failure to observe these safety information statements constitutes a use of the medical device that is inconsistent with its intended use.

Maintenance

WARNING
The medical device must be inspected and serviced regularly by service personnel. Repairs or complex maintenance work carried out on the medical device must be performed by experts. Dräger recommends that a service contract is taken out with DrägerService and that all repairs are performed by DrägerService. Dräger recommends that only genuine Dräger parts are used for maintenance. If the above are not complied with, the correct functioning of the medical device may be compromised. Observe chapter "Maintenance".

Safety checks

The medical device must be subject to regular safety checks. See chapter "Maintenance".

Accessories

WARNING
Only the accessories shown in the list of accessories 9039002 (2nd edition or later) have been tested and approved for use with the medical device. Therefore, it is strongly recommended that only these accessories are used in conjunction with the medical device. Otherwise, the correct functioning of the medical device may be compromised.
Not for use in areas of explosion hazard

**WARNING**
This medical device is neither approved nor certified for use in areas where combustible or explosive gas mixtures are likely to occur.

Safe connection to other electrical equipment

**WARNING**
Risk of patient injury
Electrical connections to equipment not listed in these instructions for use or these assembly instructions must only be made when approved by each respective manufacturer.

Device combinations

This device can be operated in combination with other Dräger devices or with devices from other manufacturers. Observe the accompanying documents of the individual devices.

If a device combination is not approved by Dräger, the safety and the correct functioning of the individual devices can be compromised. The operating organization must ensure that the device combination complies with the applicable editions of the relevant standards for medical devices.

Device combinations approved by Dräger meet the requirements of the following standards:

- IEC 60601-1, 3rd edition (general requirements for safety, device combinations, software-controlled functions)
- IEC 60601-1-2 (electromagnetic compatibility)
- IEC 60601-1-8 (alarm systems)

Or:
- IEC 60601-1, 2nd edition (general requirements for safety)
- IEC 60601-1-1 (device combinations)
- IEC 60601-1-2 (electromagnetic compatibility)
- IEC 60601-1-4 (software-controlled functions)
- IEC 60601-1-8 (alarm systems)

Connected devices

**WARNING**
Risk of electric shock and of device malfunction
Any connected devices or device combinations not complying with the requirements mentioned in these instructions for use may compromise the correct functioning of the medical device. Before operating the medical device, strictly comply with the instructions for use of all connected devices or device combinations.
For your safety and that of your patients

Connection to IT network

The connection of the medical device to a network or later changes in the network can result in previously unidentified risks for patients, users and third parties. These risks must be identified and controlled before putting the medical device into operation.

Relevant changes to the network include:
- Configuration changes
- Adding or removing additional equipment
- Update or upgrade of connected devices

Risks

Overloading of the medical device as a result of very high network traffic (e.g., due to "denial of service" attacks) could lead to deactivation of the interfaces. The service functionality would not then be available until the medical device has been restarted. In rare cases, a warm boot may take place and may occur repeatedly.

Patient safety

The design of the medical device, the accompanying documentation, and the labeling on the medical device are based on the assumption that the purchase and the use of the medical device are restricted to persons familiar with the most important inherent characteristics of the medical device. Instructions and WARNING and CAUTION statements are therefore largely limited to the specifics of the Draeger medical device.

These instructions for use do not contain references to various hazards which are obvious to users or references to the consequences of medical device misuse, or to potentially adverse effects in patients with different underlying diseases. Medical device modification or misuse can be dangerous.

CAUTION

Risk of patient injury

Do not make therapeutic decisions based solely on individual measured values and monitoring parameters.

Patient monitoring

The user of the medical device is responsible for choosing suitable monitoring that provides appropriate information about medical device performance and the patient's condition.

Patient safety may be achieved by a wide variety of means ranging from electronic surveillance of medical device performance and patient condition to simple, direct observation of clinical signs.

The responsibility for selecting the best level of patient monitoring lies solely with the user of the medical device.
For your safety and that of your patients

Information on electromagnetic compatibility

General information on electromagnetic compatibility (EMC) according to international EMC standard IEC 60601-1-2:

Electromedical devices are subject to special precautionary measures concerning electromagnetic compatibility (EMC) and must be installed and put into operation in accordance with the EMC information provided in the separate instructions for use "Workstation Critical Care and Workstation Neonatal Care".

Portable and mobile RF communications equipment can affect medical electrical equipment.

WARNING

Do not connect connectors with an ESD warning symbol and do not touch the pins of such connectors without implementing ESD protective measures. Such protective measures may include antistatic clothing and shoes, touching a ground stud before and during connection of the pins, or using electrically insulating and antistatic gloves. All relevant personnel must be instructed in these ESD protective measures.

WARNING

Do not use portable and mobile HF communications equipment, e.g., mobile phones, in the vicinity of the medical device. Maintain separation distances; see EMC information in the separate instructions for use, "Workstation Critical Care and Workstation Neonatal Care".

Disposable articles

WARNING

Risk of patient injury as a result of failure of the accessories

Disposable articles were developed, tested and manufactured for single use only. Reuse, reprocessing or sterilization can lead to a failure of the accessories and cause injuries to the patient.

Do not reuse, reprocess, or sterilize disposable articles.

Sterile accessories

CAUTION

Do not use sterile-packaged accessories if the packaging has been opened, is damaged, or if there are other signs of non-sterility.

Installing accessories

CAUTION

Install accessories to the basic device in accordance with the instructions for use of the basic device. Make sure that there is a safe connection to the basic device system.

Strictly observe instructions for use and assembly instructions.

Storing the instructions for use

CAUTION

The instructions for use must be kept in an accessible location for users.

Training

Training for users is available via the Dräger organization responsible (see www.draeger.com).
## Product-specific safety information

**WARNING**  
This medical device is intended to be used only by trained users.

**WARNING**  
Risk of fire  
The flow sensor can ignite medications or other substances based on highly flammable substances.  
– Do not nebulize medications or other substances that are easily flammable or spray them into the device.  
– Do not use substances containing alcohol.  
– Do not allow flammable or explosive substances to enter the breathing system or the breathing circuit.

**WARNING**  
Risk of fire  
Do not use the medical device in conjunction with flammable gases or flammable solutions that can mix with air, oxygen or nitrous oxide, or other sources of ignition since the medical device could ignite. Do not allow the medical device to come into contact with sources of ignition.

**WARNING**  
Do not use the medical device during magnetic resonance imaging (MRI, NMR, NMI)! This may impair correct functioning of the medical device and endanger the patient.

**WARNING**  
Do not use the medical device in hyperbaric chambers! This may impair correct functioning of the medical device and endanger the patient.

**WARNING**  
Correct functioning of the medical device may be impaired by operation of high-frequency electrosurgery units, defibrillators or short-wave therapy equipment and endanger the patient.

**WARNING**  
Risk of malfunction  
Unauthorized modifications to the medical device lead to malfunctions.  
This medical device must not be modified unless authorized by the manufacturer.

**WARNING**  
Risk of electric shock  
Live components are located under the cover. Do not open the housing of the medical device.

**WARNING**  
Risk of fire  
Do not use the medical device in oxygen-enriched rooms since the medical device could ignite. Medical device malfunctions can increase the O2 concentration in the ambient air. The medical device is only suitable for use in rooms with sufficient ventilation.

**WARNING**  
Do not obstruct the gas inlet for the safety valve. Otherwise, spontaneous breathing via the emergency breathing valve is not possible in the event of a device failure.
WARNING
With neonates, the administration of increased O2 concentrations can lead to retinopathy of prematurity. Use additional monitoring, e.g., external SpO2.

WARNING
Risk of fire
The use of unapproved O2 pressure reducers can lead to excess pressure which can cause a fire.
When supplying the ventilator with oxygen from a compressed gas cylinder, only use pressure reducers compliant with ISO 10524. Slowly open the pressure reducer manually. Do not use tools.

WARNING
Risk of unnoticed change in inspiratory O2 concentration
If an additional flow is delivered by an external flow source, the actual O2 concentration delivered may deviate from the displayed values.
Use additional monitoring, e.g., external SpO2 monitoring, if necessary.

WARNING
Risk of patient injury
If leakages are present, e.g., with non-invasive ventilation, the actual tidal volume may deviate from the measured values for VTe and VTi.
Activate leakage compensation and monitor the measured value for VT. Minimize or remedy all leakages.

WARNING
During HFO, the disconnection detection and MV monitoring are only possible to a limited extent. For this reason, use external monitoring for MV and disconnection during HFO.

WARNING
If a device for nitric oxide (NO) delivery without internal NO monitoring is used, the NO concentration must be monitored separately.

WARNING
To ensure the accuracy of the pressure measurements during HFO, it is necessary to perform a breathing circuit check to determine the resistance and compliance values. Otherwise the mean airway pressure could deviate from the set values.

WARNING
Risk of failure of flow measurement
Deposits that were not removed during reprocessing can damage the measuring wires in the flow sensor or cause a fire.
– Before inserting the flow sensor check for visible damage, soiling, and particles.
– Replace flow sensors when damaged, soiled, or not particle-free.

CAUTION
Keep away from sources of heat such as direct sunlight, heat radiators or spotlights! Otherwise the medical device may become too hot.

CAUTION
Do not obstruct or close off the vents on the medical device. Air must be able to enter freely. Otherwise the medical device may become too hot. An alarm is triggered if the medical device overheats during operation.

CAUTION
Positive-pressure ventilation can lead to negative effects, such as barotrauma or strain on the circulatory system.
For your safety and that of your patients

CAUTION
Risk of patient injury
An additional flow delivered by an external flow source can affect the measured values for airway pressure and flow.

CAUTION
Risk of malfunction
The touch screen has a sensitive surface. Damage to the surface may cause the touch-sensitive controls not to work properly.
Do not operate the screen with sharp objects.

Monitoring ventilation
The following parameters are monitored by the built-in monitoring facilities of Babylog VN500:
– Airway pressure
– Expiratory minute volume
– Respiratory rate
– Apnea alarm time
– Inspiratory O₂ concentration
– End-expiratory CO₂ concentration

Changes in these parameters may be caused by:
– Acute changes in the patient's condition
– Incorrect settings and faulty handling
– Device malfunctions
– Failure of power and gas supplies

If a fault occurs in this equipment, separate measuring instruments must be used.

During O₂ therapy, the monitoring functions of the medical device are restricted. See chapter "O₂ therapy" on page 111.

Back-up ventilation with an independent manual ventilation device

WARNING
If a fault is detected in the medical device, its life-support functions may no longer be assured. Ventilation of the patient using an independent ventilation device must be started without delay, if necessary with PEEP and/or an increased inspiratory O₂ concentration (e.g., with a manual resuscitator).
Handling Infinity ID components

Through ownership or purchase of this medical device equipped with RFID technology, you have only acquired the right to use the medical device and RFID technology in conjunction with products approved by Dräger and in strict compliance with these instructions for use. No intellectual property rights or any rights to the use of the medical device or RFID technology are hereby granted, either explicitly or implicitly, which are contrary to the above-mentioned conditions.

WARNING
Risk of patient injury
Although Babylog VN500 does not exceed the applicable limiting values for electromagnetic fields, radiation can interfere with the functioning of pacemakers. Wearers of pacemakers must keep a distance of at least 25 cm (10 in) between the pacemaker and Babylog VN500.

Emission of high-frequency energy

This medical device is equipped with an RFID (Radio Frequency Identification) system to enable wireless communication with Infinity ID accessories. Any changes or modifications to the RFID system may only be carried out by experts. Otherwise this may compromise patient safety.

This medical device has been designed and manufactured to comply with emission limit values for high-frequency energy. These limiting values are incorporated in international safety standards such as IEC 60601-1-2 (EN 60601-1-2) which have been defined by regulation authorities, such as the Federal Communications Commission (FCC Rules), Industry Canada (Radio Standards Specifications) and the European Telecommunications Standards Institute (ETSI standards).

The RFID system of this medical device complies with Part 15 of the FCC regulations, and its operation is subject to the following conditions:

1. This medical device does not cause any dangerous interference.
2. The medical device is not liable to damage caused by the reception of interference, including interference causing undesired operating conditions.

Dräger hereby declares that the RFID system in the ventilation unit is in compliance with the basic requirements and the other pertinent regulations of Directive 1999/5/EC.
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Application

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Application

Intended use

The Babylog VN500 ventilation unit of the Infinity Acute Care System is intended for the ventilation of pediatric patients and neonates. Babylog VN500 provides mandatory ventilation modes and ventilation modes for supporting spontaneous breathing and also airway monitoring. The Babylog VN500 ventilation unit is used with Dräger Infinity C Series Medical Cockpits. The Babylog VN500 ventilation unit is intended for use in different medical care areas.

Indications for use and contraindications

Indications

The Babylog VN500 ventilation unit is used in combination with Dräger Infinity C Series Medical Cockpits. Babylog VN500 is used for treating patients who require temporary or longer term respiratory support for different medical reasons.

Contraindications

It is the responsibility of the user to select the appropriate respiratory mode for the underlying disease of the patient. For all ventilator settings, the user needs to consider the respiratory status and the general state of health of the patient in order to optimally adapt the ventilation settings to the patient’s condition. Any changes to the patient’s state need to be monitored continuously.

Environment of use

Babylog VN500 is intended for stationary use in hospitals and medical rooms or for patient transportation within the hospital.

Do not use the device in the following environments:

- In hyperbaric chambers
- For magnetic resonance imaging (MRI, NMR, NMI)
- In conjunction with flammable gases or flammable solutions that can mix with air, oxygen or nitrous oxide
- In areas of explosion hazard
- In areas with combustible or explosive substances
- In rooms without sufficient ventilation

Do not operate the device with helium or helium mixtures.
System overview

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How to use the Workstation Neonatal Care

The Workstation Neonatal Care can consist of the following units:
- Infinity C500 (Medical Cockpit)
- Babylog VN500 (ventilation unit)
- Trolley 2 - 90 cm (trolley)
- GS500 (gas supply unit)
- PS500 (power supply unit)
- Transport Supply Unit (transport supply unit)

Before using the Workstation Neonatal Care, carefully read the following instructions for use:
- Instructions for use for "Workstation Critical Care and Workstation Neonatal Care"
- Instructions for use for "Infinity Medical Cockpits"
- Instructions for use for "Babylog VN500"
- Instructions for use for "Transport Supply Unit"

The Workstation Neonatal Care may include additional accessories, see separate list of accessories.
System overview

Babylog VN500 ventilation unit

Front panel

A Operation display of ventilation
   During ventilation, the inspiratory and expiratory phases are indicated by a bar display. The measured values for minute volume MVe and the inspiratory O2 concentration FIO2 are also displayed.

B Infinity ID neonatal expiratory valve with expiratory port Exp. (GAS RETURN)

C Inspiratory unit (safety valve with inspiratory port) Insp. (GAS OUTPUT)

D Gas inlet for the Emergency air intake safety valve, non-tapered connection, do not obstruct (EMERGENCY AIR INTAKE)

E Water trap

F Flap

G Muffler

H Gas outlet Exhaust, non-tapered connection (EXHAUST – NOT FOR SPIROMETER)

I Connections for future extensions

J Nebulizer port (nebulizer gas outlet for pneumatic medication nebulizer)
System overview

Rear view

A  Fuse for the batteries
B  Connection for the neonatal flow sensor V5
C  Connections for future extensions V6, V8
D  Connection for CO2 sensor V7
E  Potential equalization pin
F  Fuse for mains power supply F1, F2
G  Connection for mains power supply

Left side view

A  Connection for system cable to Infinity C500 V1
B  Connections for future extensions V2, V3
C  Connection for nurse call V4
D  Toggle switch
E  Ambient air filter with cover
F  Strain relief for cable
G  Left device flap
System overview

Right side view

A Connection for data cable to the GS500 gas supply unit **V9**
B Connection for gas connection to the GS500 gas supply unit
C Connection for Air compressed gas hose **Air** (FRESH GAS)
D Connection for O2 compressed gas hose **O2** (FRESH GAS)
E Right device flap

Trolley 2 - 90 cm

A Mount for Infinity C500
B Handle
C Trolley column
D Hose hooks
E Alignment aid
F Humidifier holder, can be swiveled
G Universal holder with standard rail
H Double castors with locking brake, set of 4
System overview

GS500 gas supply unit

Back panel

A  Rating plate
B  Gas connection
C  Screws (to hold the side panels in place)
System overview

Range of functions

The functions described correspond to the overall functionality of Babylog VN500. Some functions are only optional and may not be included in the individual device configuration. Optional functions are shown in the separate list of accessories.

Ventilation functions of Babylog VN500

Ventilation modes:
- Pressure-controlled ventilation:
  - PC-SIMV
  - PC-AC
  - PC-CMV
  - PC-APRV
  - PC-PSV
  - PC-HFO
  - PC-MMV
- Support of spontaneous breathing:
  - SPN-CPAP/PS
  - SPN-CPAP/VS
  - SPN-PPS

Additional settings for ventilation:
- Apnea Ventilation
- Flow trigger
- Sigh
- Volume Guarantee
- ATC
- AutoRelease
- HFO-Sigh
- Volume Guarantee (HFO)

Special functions:
- Maneuvers
  - Manual inspiration/hold
  - Suction maneuver
- Medication nebulization

Therapy types:
- Invasive ventilation (Tube)
- O2 Therapy
- Non-invasive ventilation (NIV)

Additional information

For a detailed description of the ventilation modes and the additional settings see page 290. Abbreviations see page 27.

Monitoring

Patient monitoring is supported by the following alarm limit settings:
- Maximum airway pressure Paw
- Expiratory minute volume MVe
- Apnea alarm time Tapn
- Respiratory rate RR
- End-expiratory CO2 concentration etCO2

The inspiratory O2 concentration is monitored by automatically set limits.

Babylog VN500 offers the following displays:
- Curves
- Graphic trends
- Numeric trends
- Loops
- Alarm history
- Logbook
- Numeric parameters
- Preconfigured lists for measured values and set values
- Customized lists for measured values and set values
- Smart Pulmonary View

During non-invasive ventilation and O2 therapy, certain monitoring functions are switched off or can be switched off.
System overview

Power supply

Babylog VN500 is designed for connection to the hospital's mains power supply of 100 to 240 V at 50/60 Hz.

If mains power fails, operation is maintained either via the internal battery of Babylog VN500 or via the PS500 power supply unit.

Gas supply

Babylog VN500 features country-specific connections for the gas supply with oxygen and medical compressed air.

The Workstation Neonatal Care may also be equipped with the GS500 external gas supply unit. GS500 supplies Babylog VN500 with compressed air.

Data transfer

A variety of interfaces can be used for transferring data:
- USB port for data export and configuration exchange using a USB storage medium
- USB port for installation of optional applications via a SIM card reader and a SIM card
- RS232 port on Infinity C500 for data transfer using the MEDIBUS or MEDIBUS.X protocol

Medication nebulizer

For medication nebulization a pneumatic medication nebulizer can be connected.

Attaching accessories

Accessories can be attached to the following holders:
- Universal holder with standard rail (G93140)
- Humidifier holder, can be swiveled (G93111)
- Humidifier holder for the lateral standard rail (8416325)

Observe the permitted maximum distance to the trolley and the permitted maximum load, see "Maximum loads of holders" on page 44.
## System overview

### Abbreviations

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<td>% leak</td>
<td>Leakage in percent</td>
<td>%MVspont</td>
<td>Spontaneous breathing portion of minute volume in percent</td>
</tr>
<tr>
<td>%PEF</td>
<td>Percentage of the peak expiratory flow</td>
<td>%PIF</td>
<td>Percentage of the peak inspiratory flow</td>
</tr>
<tr>
<td>Ah</td>
<td>Ampere hours (output specification for batteries)</td>
<td>Air</td>
<td>Connection for Air compressed gas hose (FRESH GAS)</td>
</tr>
<tr>
<td>ALARM</td>
<td>Acknowledging an alarm message that is no longer active (&quot;Reset&quot;)</td>
<td>Amplitude</td>
<td>Pressure amplitude for HFO (set value)</td>
</tr>
<tr>
<td>RESET</td>
<td>Acknowledging an alarm message that is no longer active (&quot;Reset&quot;)</td>
<td>Amplitude</td>
<td>Pressure amplitude for HFO (set value)</td>
</tr>
<tr>
<td>Ampl hf</td>
<td>Pressure amplitude for HFO (set value)</td>
<td>Ampl hf max</td>
<td>Maximum pressure amplitude for HFO (VG)</td>
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<tr>
<td>Apnea Vent.</td>
<td>Apnea ventilation</td>
<td>APRV</td>
<td>Airway Pressure Release Ventilation</td>
</tr>
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</table>
| ATC          | Automatic Tube Compensation, compensation of the tube resistance | Audio paused | Suppress acoustic alarm for 2 minutes:  
- Infinity C500 (MK31500): with the key  
- Infinity C500 (MS18746): with the key |
| BF           | Insulation class Body Floating | BTPS         | Body Temperature Pressure Saturated, measured values based on the condition of the patient’s lungs, body temperature 37 °C (98.6 °F), water vapor-saturated gas, ambient pressure and mean airway pressure |
| C            | Compliance | C20/Cdyn     | Index of the last 20 % of compliance in relation to the dynamic total compliance |
| Cdyn         | Dynamic compliance | cmH2O        | Unit of measurement for pressure  
1 cmH2O = approx. 1 mbar |
| Compens.     | Degree of tube compensation | COPD         | Chronic Obstructive Pulmonary Disease |
| Cycles sigh  | Number of cycles during a sigh phase (set value) | DCO2         | Dissociation coefficient for CO2 with HFO |
| Device flow  | Delivered inspiratory flow with PC-HFO | DHCP         | Dynamic Host Configuration Protocol |
| E            | Elastance | EIP          | End Inspiratory Pressure |
| EMC          | Electromagnetic compatibility | Emergency air intake | Safety air inlet, inspiratory relief valve (EMERGENCY AIR INTAKE) |
| ESD          | Electrostatic Discharge | ET           | Endotracheal tube |
| etCO2        | End-expiratory CO2 concentration | ETSI         | European Telecommunications Standards Institute |
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<td>Gas outlet (EXHAUST – NOT FOR SPIROMETER)</td>
<td>Insp. flow</td>
<td>Inspiratory flow</td>
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<td>Exp.</td>
<td>Label on the device, Expiratory port (GAS RETURN)</td>
<td>Interval sigh</td>
<td>Time between two sigh phases (set value)</td>
</tr>
<tr>
<td>Exp.</td>
<td>Expiration</td>
<td>IP21</td>
<td>Degree of protection against ingress of liquids and particles</td>
</tr>
<tr>
<td>Exp. term.</td>
<td>Termination criterion in % from the peak expiratory flow</td>
<td>LAN</td>
<td>Local Area Network</td>
</tr>
<tr>
<td>FCC</td>
<td>Federal Communications Commission, regulatory authority for communications devices in the U.S.</td>
<td>MAPhf</td>
<td>Mean airway pressure for HFO (set value)</td>
</tr>
<tr>
<td>fhf</td>
<td>Frequency of oscillation for HFO (set value)</td>
<td>mbar</td>
<td>Millibar, unit of measurement for pressure 1 mbar = approx. 1 cmH2O</td>
</tr>
<tr>
<td>FiO2</td>
<td>Inspiratory O2 concentration (set value)</td>
<td>MEDIBUS</td>
<td>Dräger communication protocol for medical devices</td>
</tr>
<tr>
<td>Flow</td>
<td>Flow (set value)</td>
<td>MEDIBUS.X</td>
<td>Dräger communication protocol for medical devices with a data definition which is standardized across all devices</td>
</tr>
<tr>
<td>Flow Assist</td>
<td>Flow support in SPN-PPS (set value)</td>
<td>mmHg</td>
<td>Unit of measurement for end-expiratory CO2 concentration</td>
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<td>Flow max</td>
<td>Maximum inspiratory flow during NIV (Neo, patient category)</td>
<td>More...</td>
<td>Show more alarms</td>
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<td>Flow trigger</td>
<td>Trigger threshold, sensitivity (set value)</td>
<td>MRI</td>
<td>Magnetic resonance imaging</td>
</tr>
<tr>
<td>FRC</td>
<td>Functional Residual Capacity</td>
<td>MV</td>
<td>Minute volume, leakage-corrected</td>
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<tr>
<td>GS500</td>
<td>Gas supply unit</td>
<td>MV delay</td>
<td>Duration of alarm suppression for MV high and MV low</td>
</tr>
<tr>
<td>HFO</td>
<td>High Frequency Oscillation</td>
<td>MV high</td>
<td>Upper alarm limit for minute volume</td>
</tr>
<tr>
<td>HME</td>
<td>Heat Moisture Exchanger</td>
<td>MV low</td>
<td>Lower alarm limit for minute volume</td>
</tr>
<tr>
<td>hPa</td>
<td>Hectopascal, unit of measurement for pressure 1 hPa = 1 mbar = approx. 1 cmH2O</td>
<td>MVapn</td>
<td>Minute volume during apnea ventilation</td>
</tr>
<tr>
<td>I:E</td>
<td>Ratio of inspiratory time to expiratory time (set value)</td>
<td>MVe</td>
<td>Expiratory minute volume, overall, not leakage-corrected</td>
</tr>
<tr>
<td>I:Ehf</td>
<td>I:E for HFO (set value)</td>
<td>MVemand</td>
<td>Mandatory expiratory minute volume</td>
</tr>
<tr>
<td>I:Espon</td>
<td>I:E during spontaneous breathing</td>
<td>MVespon</td>
<td>Spontaneous expiratory minute volume</td>
</tr>
<tr>
<td>IEC/CEI</td>
<td>Alarm tone in accordance with IEC 60601-1-8</td>
<td>MVi</td>
<td>Inspiratory minute volume, overall, not leakage-corrected</td>
</tr>
<tr>
<td>Insp.</td>
<td>Label on the device, Inspiratory port (GAS OUTPUT)</td>
<td>MVleak</td>
<td>Leakage minute volume</td>
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<td>Neo.</td>
<td>Neonates patient category</td>
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<tr>
<td>NiMH</td>
<td>Nickel-metal hydride (battery technology)</td>
</tr>
<tr>
<td>NIV</td>
<td>Non-Invasive Ventilation</td>
</tr>
<tr>
<td>NMI</td>
<td>Nuclear magnetic imaging</td>
</tr>
<tr>
<td>NMR</td>
<td>Nuclear magnetic resonance</td>
</tr>
<tr>
<td>NO</td>
<td>Nitric oxide</td>
</tr>
<tr>
<td>NTPD</td>
<td>Normal Temperature Pressure Dry, 20 °C (68 °F), 1013 hPa, dry</td>
</tr>
<tr>
<td>O2</td>
<td>Connection for O2 compressed gas hose (FRESH GAS)</td>
</tr>
<tr>
<td>O2 suction</td>
<td>Suction maneuver</td>
</tr>
<tr>
<td>Palv</td>
<td>Alveolar pressure</td>
</tr>
<tr>
<td>Paw</td>
<td>Airway pressure</td>
</tr>
<tr>
<td>Paw high</td>
<td>Upper alarm limit for airway pressure</td>
</tr>
<tr>
<td>PC-AC</td>
<td>Pressure Control-Assist Control, assisted-controlled, pressure-controlled ventilation with back-up respiratory rate</td>
</tr>
<tr>
<td>PC-APRV</td>
<td>Pressure Control-Airway Pressure Release Ventilation, spontaneous breathing under continuous positive airway pressure with brief pressure releases</td>
</tr>
<tr>
<td>PC-CMV</td>
<td>Pressure Control-Continuous Mandatory Ventilation, continuous pressure-controlled ventilation</td>
</tr>
<tr>
<td>PC-HFO</td>
<td>Pressure Control-High Frequency Oscillation, pressure-controlled ventilation with high-frequency oscillation</td>
</tr>
<tr>
<td>PC-MMV</td>
<td>Pressure Control-Mandatory Minute Volume Ventilation, pressure-controlled ventilation to ensure minimum minute ventilation</td>
</tr>
<tr>
<td>PC-PSV</td>
<td>Pressure Control-Pressure Support Ventilation, spontaneous breathing at continuous positive pressure level with pressure support and back-up respiratory rate</td>
</tr>
<tr>
<td>PC-SIMV</td>
<td>Pressure Control-Synchronized Intermittent Mandatory Ventilation, intermittent, triggered, pressure-controlled ventilation</td>
</tr>
<tr>
<td>Ped. pat.</td>
<td>Pediatric patient category</td>
</tr>
<tr>
<td>PEEP</td>
<td>Positive end-expiratory pressure</td>
</tr>
<tr>
<td>Phigh</td>
<td>Upper pressure level in APRV (set value)</td>
</tr>
<tr>
<td>Pinsp</td>
<td>Inspiratory pressure (set value)</td>
</tr>
<tr>
<td>PIP</td>
<td>Peak Inspiratory Pressure</td>
</tr>
<tr>
<td>Plow</td>
<td>Lower pressure level in APRV (set value)</td>
</tr>
<tr>
<td>PmanInsp</td>
<td>Pressure of the breath for manual inspiration during NIV (Neo. patient category, SPN-CPAP ventilation mode)</td>
</tr>
<tr>
<td>Pmax</td>
<td>Maximum allowed airway pressure (set value)</td>
</tr>
<tr>
<td>Pmax/Paw high autoset</td>
<td>Linking the maximum airway pressure to the alarm limit Paw high</td>
</tr>
<tr>
<td>Pmean</td>
<td>Mean airway pressure</td>
</tr>
<tr>
<td>Pmin</td>
<td>Minimum airway pressure</td>
</tr>
<tr>
<td>Pplat</td>
<td>Airway pressure on the plateau</td>
</tr>
<tr>
<td>PS</td>
<td>Pressure Support</td>
</tr>
<tr>
<td>PS500</td>
<td>Power supply unit</td>
</tr>
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<td>Psigh</td>
<td>Inspiratory pressure of sigh for HFO (set value)</td>
</tr>
<tr>
<td>Psupp</td>
<td>Pressure support absolute</td>
</tr>
<tr>
<td>Ptrach</td>
<td>Pressure in the trachea</td>
</tr>
<tr>
<td>R</td>
<td>Total resistance</td>
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<tr>
<td>$r^2$</td>
<td>Correlation coefficient for the calculation method &quot;Least Mean Square&quot; for R, C and TC</td>
<td>SPN-CPAP/PS</td>
<td>Spontaneous-Continuous Positive Airway Pressure/Pressure Support, spontaneous breathing with continuous positive pressure level with or without pressure support</td>
</tr>
<tr>
<td>REF</td>
<td>Material and revision number of the medical device</td>
<td>SPN-CPAP/VS</td>
<td>Spontaneous-Continuous Positive Airway Pressure/Volume Support, spontaneous breathing with continuous positive pressure level with or without volume support</td>
</tr>
<tr>
<td>RF</td>
<td>Radio Frequency</td>
<td>SPN-PPS</td>
<td>Spontaneous-Proportional Pressure Support, spontaneous breathing with flow-proportional and volume-proportional pressure support</td>
</tr>
<tr>
<td>RFID</td>
<td>Radio Frequency Identification</td>
<td></td>
<td></td>
</tr>
<tr>
<td>Rpat</td>
<td>Patient resistance, patient airway resistance</td>
<td>SpO2</td>
<td>Partial $O_2$ saturation</td>
</tr>
<tr>
<td>RR</td>
<td>Respiratory rate (set value)</td>
<td>STPD</td>
<td>Standard Temperature Pressure Dry, 0 °C (32 °F), 1013 hPa, dry</td>
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<tr>
<td>RR high</td>
<td>Upper alarm limit for respiratory rate</td>
<td>Tapn</td>
<td>Apnea alarm time (set value)</td>
</tr>
<tr>
<td>RRapn</td>
<td>Respiratory rate of apnea ventilation (set value)</td>
<td>TC</td>
<td>Time constant tau</td>
</tr>
<tr>
<td>RRmand</td>
<td>Mandatory portion of respiratory rate</td>
<td>TCe</td>
<td>Time constant calculated from VTe and peak expiratory flow</td>
</tr>
<tr>
<td>RRsigh</td>
<td>Respiratory rate of the sighs during HFO (set value)</td>
<td>Tdisconnect</td>
<td>Time for disconnection alarm (set value)</td>
</tr>
<tr>
<td>RRspon</td>
<td>Spontaneous breathing portion of respiratory rate</td>
<td>Te</td>
<td>Expiratory time (set value)</td>
</tr>
<tr>
<td>RRtrig</td>
<td>Portion of mandatory triggered breaths</td>
<td>TGI</td>
<td>Tracheal Gas Insufflation, tracheal gas insufflation</td>
</tr>
<tr>
<td>RSB</td>
<td>Rapid Shallow Breathing, quotient of spontaneous respiratory rate and tidal volume</td>
<td>Thigh</td>
<td>Time of upper pressure level in APRV (set value)</td>
</tr>
<tr>
<td>SIM</td>
<td>Subscriber Identity Module, participant identification</td>
<td>Ti</td>
<td>Inspiratory time (set value)</td>
</tr>
<tr>
<td>Slope</td>
<td>Pressure rise time (set value)</td>
<td>Timax</td>
<td>Maximum inspiratory time for flow during pressure or volume support (set value)</td>
</tr>
<tr>
<td>Slopesigh</td>
<td>Pressure rise time of the sighs during HFO</td>
<td>Tisigh</td>
<td>Inspiratory time of sigh for HFO (set value)</td>
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<td>Smart Pulmonary View</td>
<td>Graphic display of lung characteristics (Lung display)</td>
<td>Tispon</td>
<td>Inspiratory time during spontaneous breathing</td>
</tr>
<tr>
<td>SN</td>
<td>Device serial number</td>
<td>Tisupp</td>
<td>Inspiratory time during pressure support</td>
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<tr>
<td>SPN-CPAP</td>
<td>Spontaneous-Continuous Positive Airway Pressure, spontaneous breathing with continuous positive pressure level</td>
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<td>Tlow</td>
<td>Time of lower pressure level in APRV</td>
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<td>Tlow max</td>
<td>Maximum expiratory time during APRV (set value)</td>
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<tr>
<td>TmanInsp</td>
<td>Duration of the breath for manual inspiration during NIV (patient category Neo., ventilation mode SPN-CPAP)</td>
</tr>
<tr>
<td>Tplat</td>
<td>Time of inspiratory plateau</td>
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<td>Trach.</td>
<td>Tracheostomy tube</td>
</tr>
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<td>Tube Ø</td>
<td>Inner diameter of tube (set value)</td>
</tr>
<tr>
<td>UMDNS</td>
<td>Universal Medical Device Nomenclature System, nomenclature for medical devices</td>
</tr>
<tr>
<td>Un</td>
<td>Rated voltage</td>
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<tr>
<td>USB</td>
<td>Universal Serial Bus, serial bus system</td>
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<td>VG</td>
<td>Volume Guarantee, Volume Guarantee</td>
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<td>VG (HF)</td>
<td>Volume Guarantee for HFO</td>
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<td>Vol. Assist</td>
<td>Volume support in SPN-PPS (set value)</td>
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<td>VRLA</td>
<td>Valve-regulated lead-acid (battery technology)</td>
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<td>VS</td>
<td>Volume Support</td>
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<td>VT</td>
<td>Tidal volume, leakage-corrected</td>
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<td>VTapn</td>
<td>Tidal volume of apnea ventilation (set value)</td>
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<td>VTe</td>
<td>Expiratory tidal volume, not leakage-corrected</td>
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<td>VTmand</td>
<td>Expiratory tidal volume during a mandatory breath</td>
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<td>VTespon</td>
<td>Expiratory tidal volume during a spontaneous breath</td>
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<td>VThf</td>
<td>Tidal volume for HFO (set value for VG (HF))</td>
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<td>VTi</td>
<td>Inspiratory tidal volume, not leakage-corrected</td>
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## Symbols

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<td></td>
<td>Acoustic alarm suppressed for 2 minutes</td>
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<td>Batteries defective or no information available on their charge state</td>
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<td>Group Views, screen displays</td>
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<td>Lower alarm limit</td>
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<td>Group Trends/Data, information on the course of ventilation</td>
<td></td>
<td>Upper alarm limit</td>
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<td>Group Special maneuvers</td>
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<td>Setting or access locked</td>
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<td></td>
<td>Group Alarms</td>
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<td>Expiratory valve locked</td>
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<td>Group Therapy, ventilation parameter settings</td>
<td></td>
<td>Setting or access unlocked</td>
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<tr>
<td></td>
<td>Group configuration, system settings, and settings for sensors</td>
<td></td>
<td>Expiratory valve unlocked</td>
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<td></td>
<td>Group Start/Standby</td>
<td>Exhaust</td>
<td>Gas outlet (EXHAUST – NOT FOR SPIROMETER)</td>
</tr>
<tr>
<td></td>
<td>Switch system on or off (with the key on Infinity C500)</td>
<td></td>
<td>Pediatric patient category (Ped. pat.)</td>
</tr>
<tr>
<td></td>
<td>Alarm limit off</td>
<td></td>
<td>Neonates patient category (Neo.)</td>
</tr>
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<td></td>
<td>Configure trends</td>
<td></td>
<td>Display additional information or open Help</td>
</tr>
<tr>
<td></td>
<td>Save screen display</td>
<td></td>
<td>Hide additional information or close Help</td>
</tr>
<tr>
<td>View 1</td>
<td></td>
<td></td>
<td>Scroll back in tables or lists</td>
</tr>
<tr>
<td>View 2</td>
<td></td>
<td></td>
<td>Scroll forward in tables or lists</td>
</tr>
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<td>View 3</td>
<td></td>
<td></td>
<td>Scroll forward in Help</td>
</tr>
<tr>
<td>Medication nebulizer</td>
<td></td>
<td></td>
<td>Scroll backward in Help</td>
</tr>
<tr>
<td></td>
<td>Charge state of batteries 90 to 100 %</td>
<td></td>
<td>Close dialog window</td>
</tr>
<tr>
<td></td>
<td>Charge state of batteries 60 to &lt;90 %</td>
<td></td>
<td>Active test in the device check</td>
</tr>
<tr>
<td></td>
<td>Charge state of batteries 40 to &lt;60 %</td>
<td></td>
<td>Spontaneous breathing activity by the patient</td>
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<tr>
<td></td>
<td>Charge state of batteries 20 to &lt;40 %</td>
<td></td>
<td>NIV, Non-invasive ventilation</td>
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<table>
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<th>Explanation</th>
<th>Symbol</th>
<th>Explanation</th>
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<tr>
<td>![Audio paused]</td>
<td>Suppress acoustic alarm for 2 minutes (key on Infinity C500 MS18746)</td>
<td></td>
<td>Device switched off (only the following symbol is used in the instructions for use):</td>
</tr>
<tr>
<td>![Suppression]</td>
<td>Suppress acoustic alarm for 2 minutes (key on Infinity C500 MK31500)</td>
<td>![GFCI]</td>
<td>Labeling for FCC approval</td>
</tr>
<tr>
<td>![Mains power supply]</td>
<td>Mains power supply (AC voltage)</td>
<td>![GFCI]</td>
<td>Labeling in accordance with Directive 93/42/EEC concerning medical products</td>
</tr>
<tr>
<td>![GS500]</td>
<td>GS500: Mains power supply (AC voltage, if separate mains plug is available)</td>
<td>![GFCI]</td>
<td>Labeling in accordance with Directive 1999/5/EC on radio equipment and telecommunications terminal equipment</td>
</tr>
<tr>
<td>![Power supply from batteries]</td>
<td>Power supply from batteries</td>
<td>![Serial interface]</td>
<td>Serial interface (on Infinity C500)</td>
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<tr>
<td>![Caution]</td>
<td>Caution: Observe important safety information and precautions in the instructions for use.</td>
<td>![LAN interface]</td>
<td>LAN interface (on Infinity C500)</td>
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<tr>
<td>![Observe the instructions]</td>
<td>Observe the instructions for use</td>
<td>![USB interface]</td>
<td>USB interface (on Infinity C500)</td>
</tr>
<tr>
<td>![Connection for equipotential bonding]</td>
<td>Connection for equipotential bonding</td>
<td>![DVI interface]</td>
<td>DVI interface (on Infinity C500)</td>
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<tr>
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<td>Protective earth</td>
<td>![Attention!]</td>
<td>Attention!</td>
</tr>
<tr>
<td>![Nurse call]</td>
<td>Nurse call</td>
<td>![Nominal weight and maximum weight]</td>
<td>Label regarding intrahospital transport</td>
</tr>
<tr>
<td>![Marking point on the trolley]</td>
<td>Marking point on the trolley – do not lean, press, push or pull against the trolley above the marking points</td>
<td>![Temperature limitation during storage]</td>
<td>Nominal weight and maximum weight (for information, see chapter &quot;Technical data&quot;)</td>
</tr>
<tr>
<td>![ESD warning symbol]</td>
<td>ESD warning symbol</td>
<td>![Ambient pressure]</td>
<td>Temperature limitation during storage</td>
</tr>
<tr>
<td>![ESD warning symbol]</td>
<td>ESD warning symbol</td>
<td>![Relative humidity]</td>
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<tr>
<td>![Information on disposal]</td>
<td>Information on disposal</td>
<td>![Use by]</td>
<td>Relative humidity</td>
</tr>
<tr>
<td>![Manufacturer]</td>
<td>Manufacturer</td>
<td>![Keep dry]</td>
<td>Use by</td>
</tr>
<tr>
<td>![Date of manufacture]</td>
<td>Date of manufacture</td>
<td>![Keep dry]</td>
<td>Keep dry</td>
</tr>
<tr>
<td>![Connection for the neonatal flow sensor]</td>
<td>Connection for the neonatal flow sensor</td>
<td>![Device ready for switch-on]</td>
<td>Device ready for switch-on (only the following symbol is used in the instructions for use):</td>
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Instructions for use Infinity Acute Care System – Babylog VN500 SW 2.n
Operating concept

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Operating concept

Operating concept for Infinity C500

Infinity C500 is the central operating and display unit. The general operating concept is described in the instructions for use for "Infinity Medical Cockpits".

Operating concept for Babylog VN500

The following operating concept only contains the specific information and operating steps for Babylog VN500.

This chapter describes:
- Main screen
- Main menu bar
- Dialogs windows
- Therapy line
- Therapy controls
- Setting ventilation parameters
- Exceeding the set limit of a ventilation parameter
- Direct setting of ventilation parameters (QuickSet)
- Linked setting of ventilation parameters

Main screen

The main screen displays the most important ventilation information at a glance.

A Header bar with the following fields:
- Patient category, see page 66
- System data, e.g., state of charge of the batteries, see page 119
- Therapy status: Therapy type (ventilation or O2 Therapy), ventilation mode and additional settings
- Alarms, messages and instructions for the user, see page 124
- Alarm status

B Monitoring area with curves, loops, trends and measured values, see page 95. The display can be configured, see page 158.

C Main menu bar with buttons for opening dialog windows and activating functions, see page 37.

D Therapy bar with the therapy controls for the ventilation parameters of the active ventilation mode, see page 38.

The main screen can be configured as a Main screen button in the main menu bar to enable direct access. See "Assigning functions to additional buttons" on page 161.
Main menu bar

The main menu bar contains fixed assigned and configurable buttons. The buttons are assigned to various groups. Touching a button opens the corresponding dialog window or activates the corresponding function.

Fixed assigned buttons

<table>
<thead>
<tr>
<th></th>
<th>F</th>
<th>A</th>
</tr>
</thead>
<tbody>
<tr>
<td></td>
<td>G</td>
<td>B</td>
</tr>
<tr>
<td></td>
<td>H</td>
<td>C</td>
</tr>
<tr>
<td></td>
<td></td>
<td>D</td>
</tr>
</tbody>
</table>

A **Alarms**... for setting the alarm limits, displaying the alarm logbook and listing all active alarms, see page 124.

B **Ventilation settings**... for setting the ventilation mode and the ventilation parameters, see page 84.

C **Sensors/Parameters**... for calibrating the sensors and activating or deactivating monitoring, see page 141.

D **System setup**... for configuring the device functions, see page 155.

E **Start/Standby**... for selecting standby mode or starting therapy, see page 114.

F **Views**... for switching to other configured monitoring area views, see page 95.

G **Trends/Data**... for displaying all the measured and set values, logbook, trends and for exporting data, see page 133.

H **Special maneuvers**... for selecting additional functions, e.g. suction maneuver, see page 101, or medication nebulization, see page 103.

Configurable buttons

Additional buttons for directly accessing functions or dialogs can be configured. These buttons are spatially assigned to the corresponding group. See "Assigning functions to additional buttons" on page 161.
Dialog windows

Dialog windows consist of one or several pages which are displayed by touching the corresponding horizontal or vertical tab. Dialog windows contain elements for operating the device and informing the user on current settings. Dialog windows can be opened by touching a button in the main menu bar.

Therapy controls

The therapy controls (A) are used to set the ventilation parameters.

Therapy controls are contained in the therapy bar of the active ventilation mode and in the dialog window for the ventilation settings.

Start-up settings

Arrows beside the scales on the therapy controls indicate the start-up values valid when Babylog VN500 is switched on. These start-up values can be adjusted specifically as required by the hospital. See "Configuring start-up settings for the ventilation parameters" on page 171.

Locking mechanism

The therapy controls in the therapy bar can be locked against the ventilation parameters being changed by accident. See "Locking therapy controls in the therapy bar" on page 164.
Setting ventilation parameters

1. Touch the therapy control. The color turns yellow. The unit of the parameter to be adjusted is displayed in parentheses.
2. Turn the rotary knob to set the value.
3. Press the rotary knob to confirm the value. The color of the therapy control turns dark green.

The following chapters of the instructions for use provide a simplified explanation of these steps: "Use the rotary knob to set and confirm the value."

Exceeding the set limit of a ventilation parameter

When a set limit of a parameter has been reached, Babylog VN500 displays a message.

- Press the rotary knob to exceed the set limit.
- The set limit can be exceeded.

If the maximum set limit for a parameter has been reached, e.g., when it is dependent on other parameters, it is not possible to exceed the set limit.

- Press the rotary knob. Babylog VN500 adopts the maximum possible set value.

Direct setting of ventilation parameters (QuickSet)

When a ventilation parameter is set directly, the changes to a setting become immediately effective for the patient. The user can immediately see the effect the changed setting has on the patient. The finally chosen setting does not have to be confirmed again.

Ventilation parameters can be set directly in all ventilation modes and can be carried out in the dialog window for the ventilation settings. Direct settings are only possible in the therapy bar when the therapy controls are not locked.

O2 and Flow cannot be set directly.

Setting ventilation parameters directly

1. Touch the corresponding therapy control.
2. Press the rotary knob and hold for approximately 3 seconds.

The therapy control changes to dark green with a yellow edge. The direct setting function is now active.

3. Press and hold the rotary knob and turn to set the value.

The set value is immediately effective.

Exceeding the set limit of a parameter with direct setting

When a set limit of a parameter has been reached, Babylog VN500 displays a message.

- Press the rotary knob. Babylog VN500 adopts the maximum possible set value.

5. Press the rotary knob again and turn it.

The set limit can be exceeded.
**Operating concept**

**Linked setting of ventilation parameters**

The linked setting is possible for **PEEP/Pinsp** and for **RR/Ti**.

**Linking PEEP/Pinsp**

1. Touch the therapy control **PEEP** (A) or **Pinsp** (B); the color turns to yellow. The **Link** button (C) is displayed.

2. Touch the **Link** button (C).

3. Turn the rotary knob to set the value for **PEEP** and **Pinsp**. The other value is also automatically changed so that the difference in pressure remains constant.

4. Press the rotary knob to confirm the value.

Both therapy controls turn dark green.

**Linking RR/Ti**

Linking **RR** and **Ti** is done in the same way as linking **PEEP** and **Pinsp**. The I:E ratio remains constant. If the respiratory rate is increased, the inspiratory time is reduced. If the inspiratory time is increased, the respiratory rate is reduced.

**Additional information**

If a condition is reached in which a parameter cannot be changed anymore when setting linked parameters, Babylog VN500 displays a corresponding message in the message field (D).
Assembly and preparation

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**Safety information for assembly and preparation**

**WARNING**
Before each use, reprocess the device and all accessories in accordance with the instructions for use, see "Reprocessing list" on page 236. Observe the hospital hygiene regulations!

**WARNING**
Securely mount Babylog VN500. Check for secure fit. Danger of damage to device or personal injury!

**WARNING**
Risk of tipping over
Do not tilt the device by more than 5°.

**WARNING**
Do not place any containers with liquid on or above the device! Penetrating liquid may cause malfunction of or damage to the device, which may endanger the patient.

**WARNING**
Failure to observe the permitted maximum load and weight distribution may result in the device toppling over. Danger of damage to device or personal injury! Observe the permitted maximum load and weight distribution, see "Maximum load" on page 279.

**CAUTION**
When parking the device, lock all the double castors of the trolley and check that the brakes are working properly.

**Preparation of the trolley**

**CAUTION**
Connect all devices securely to the trolley. Check for secure fit. Danger of damage to device or personal injury!

**WARNING**
Do not use the trolley in the event of visible damage, e.g., damaged double castors! Contact DrägerService.

**WARNING**
Do not lean, press, push or pull against the trolley above the marking points on the trolley. The trolley could topple over.
Assembly and preparation

Connecting the universal holder with standard rail to the trolley

Attach the universal holder with standard rail to the front of the trolley.

1. Unscrew the adjusting screw (A) completely.
2. Attach the right-hand side of the universal holder to the right-hand side of the rail (B). Make sure that the catch of the universal holder is completely behind the alignment aid.
3. Align the universal holder (C) horizontally and press the left-hand side of the universal holder onto the left-hand side of the column.
4. Tighten the adjusting screw (A). Make sure that the catch of the universal holder is completely behind the alignment aid.
5. Check that the universal holder is fixed securely.

Adjusting the height of the universal holder

1. Unscrew the adjusting screw (A).
2. Adjust the height of the universal holder (C).
3. Align the universal holder horizontally.
4. Retighten the adjusting screw (A).

Connecting the humidifier holder to the trolley

The humidifier holder is attached to the front of the trolley. The humidifier holder can be fastened on the left or right-hand side of the trolley column. The attachment of the humidifier holder on the right-hand side is shown.

1. Hold the humidifier holder at the desired height on the guide (A) of the trolley column.
2. Turn the clamping screw (B) to the left until the base (C) fits into the guide of the trolley column.
3. Turn the clamping screw (B) to the right until the humidifier holder is secured firmly in the guide.
4. Move the standard rail (D) to the desired position.
Assembly and preparation

Securing accessories to the standard rail

Maximum loads of holders

The following information applies to the holders:

<table>
<thead>
<tr>
<th>Holder</th>
<th>Position of the holder</th>
<th>Maximum load</th>
<th>Possible accessories</th>
<th>Maximum distance to the lateral standard rail</th>
</tr>
</thead>
<tbody>
<tr>
<td>Universal holder with standard rail (G93140)</td>
<td>On the front of the trolley</td>
<td>10 kg (22 lb)</td>
<td>Breathing gas humidifier, medication nebulizer</td>
<td>–</td>
</tr>
<tr>
<td>Humidifier holder, can be swiveled (G93111)</td>
<td>On the side of the trolley</td>
<td>5 kg (11 lb)</td>
<td>Breathing gas humidifier</td>
<td>–</td>
</tr>
<tr>
<td>Humidifier holder for the lateral standard rail (8416325)</td>
<td>On the lateral standard rails of the ventilation unit(1)</td>
<td>5 kg(2) (11 lb)</td>
<td>Breathing gas humidifier</td>
<td>10 cm (3.9 in)</td>
</tr>
<tr>
<td>IACS hinged arm (MP00690)</td>
<td>On the lateral standard rails of the ventilation unit(1)</td>
<td>1 kg (2.2 lb)</td>
<td>Breathing hoses</td>
<td>100 cm (39.4 in)</td>
</tr>
</tbody>
</table>

1) Maximum load on the lateral standard rails of the ventilation unit: 5 kg (11 lb) on each lateral standard rail
2) If a hinged arm is attached to the lateral standard rails of the ventilation unit in addition to the humidifier holder (8416325), the maximum load of 5 kg (11 lb) per lateral standard rail must be observed. The humidifier holder can then only support 4 kg (8.8 lb).

Securing the compressed gas cylinders to the trolley

Only available with the cylinder holder option

WARNING
Securely attach the compressed gas cylinders to the trolley, using both hook-and-loop straps. Otherwise there is a risk of the trolley toppling over. Danger of damage to device or personal injury!
### Assembly and preparation

**WARNING**
The length of the hook-and-loop straps must match the diameter of the compressed gas cylinders to ensure that the hook-and-loop straps can hold the cylinders securely. If necessary have an appropriate hook-and-loop strap fitted by service personnel. This is essential to ensure that the compressed gas cylinders are properly secured.

Compressed gas cylinders with the following dimensions can be secured:

- **Diameter:** 80 to 176 mm (3.15 to 6.93 in)
- **Length:** 420 to 760 mm (16.54 to 29.92 in)

**WARNING**
Not every combination of compressed gas cylinder diameter and length can be secured. When used in combination with a pressure reducer, the compressed gas cylinder must not come into contact with the console of the trolley. The maximum diameter is 176 mm (6.93 in) when the base of the compressed gas cylinder is resting completely on the base plate of the lower holder or is semi-spherical in shape.

1. Place the cylinders into the mountings on the trolley.
2. Secure each cylinder with 2 hook-and-loop straps (A).
3. Secure the compressed gas hoses by hanging them over the hose hooks (B).

**WARNING**
Position the compressed gas cylinders fitted with pressure reducers in such a way to prevent the pressure reducers from being damaged during transport. The lower part of the trolley is designed to protect against collisions. Take particular care when the compressed gas cylinders being used extend beyond this collision protection.
Preparation and Installation

Preparing the Medical Cockpit

Positioning Infinity C500

Infinity C500 is suitable for positioning on the trolley or on a standard rail.

Positioning the Infinity C500 on the trolley

1. Hook the Infinity C500 holder (A) into the mounting (B) on the trolley.
2. Tighten the locking screw (C).
3. Make sure that Infinity C500 is securely attached to the trolley.

Positioning Infinity C500 on a standard rail

If Infinity C500 is connected to the trolley:

1. Unscrew the locking screw (C).
2. Lift Infinity C500 out of the mounting (B) on the trolley.
3. Hook Infinity C500 into the standard rail.
4. Tighten the locking screw.
5. Make sure that Infinity C500 is securely attached to the standard rail.

Positioning Infinity C500

Tilting the position of Infinity C500

Infinity C500 can be tilted down and up.

1. Press and hold the tilt release button (A).
2. At the same time, tilt Infinity C500 to the desired working position.
3. Release the button and make sure that it engages securely.

Turning Infinity C500

Infinity C500 can be turned by a maximum of 180° counterclockwise or 90° clockwise.

- Turn to the desired working position.
Assembly and preparation

Connecting the system cable

The system cable is connected to Infinity C500 and to Babylog VN500. The system cable is fixed in a clamp.

Connecting the system cable to Infinity C500

On Infinity C500 (MS18746):

1. Unscrew the cover from the socket (A).
2. Insert the system cable connector (B) into the socket (A). Ensure that the connector is inserted with the correct orientation.
3. Screw the cover back on.

On Infinity C500 (MK31500):

1. Open the flap on the left-hand side of Babylog VN500.
2. Run the system cable between Babylog VN500 and the handle.
3. Clip the protective sleeve (C) immediately after the connector (E) onto the system cable (D). Align the protective sleeve so that the slots of the protective sleeve are facing downwards and upwards.
4. Insert the system cable connector (E) into the socket until the connector audibly clicks into place.
5. Insert the protective sleeve (C) into the protective plate (F) at the same time.
6. Turn the protective sleeve (C) by approximately 90° until it clicks into place. The cable is secured.
7. Close the left-hand flap.

Disconnecting the system cable from Babylog VN500

1. Push the locking mechanism on the connector (E) backwards and pull out the connector.
2. Turn the protective sleeve (C) by approximately 90° and withdraw it from the protective plate (F).
Assembly and preparation

Fixing the system cable in the clamp (G)

1. Open the clamp cover (H).
2. Place the system cable into the clamp. Keep the cable length short between the clamp and Babylog VN500.
3. Close the clamp cover (H) and engage. Ensure that the cover engages securely.

Removing the system cable from the clamp

1. Open the clamp cover.
2. Remove the cable from the clamp.
3. Close the clamp cover and engage.

Using the MEDIBUS or the MEDIBUS.X protocol

WARNING
Risk of patient injury
All data transferred via the MEDIBUS interface are for information only and must not be used as the sole basis for diagnostic or therapeutic decisions. The MEDIBUS interface is not intended for use with a distributed alarm system conforming to IEC 60601-1-8:2012.

MEDIBUS and MEDIBUS.X are software protocols for transferring data between Babylog VN500 and an external medical or non-medical device (e.g., patient monitors or computers for data management systems).

Additional information
For MEDIBUS:
"MEDIBUS for V and VN ventilators" (9039527)
"Dräger RS 232 MEDIBUS, Protocol Definition" (9028258)
For MEDIBUS.X:
"MEDIBUS.X, Rules and Standards for Implementation" (9052607)
"MEDIBUS.X, Profile Definition for Data Communication V1.n" (9052608)
Assembly and preparation

Connecting an external device for using MEDIBUS or MEDIBUS.X

On Infinity C500 (MS18746):

- Connect an external device to the COM 1, COM 2 or COM 3 (A) interface of Infinity C500.
  Use MEDIBUS cable 8416326.

Configuring the interface

A description is given in chapter "Configuring interfaces" on page 185.

LAN and USB interfaces of Infinity C500

On Infinity C500 (MS18746):

On Infinity C500 (MK31500):

Use of LAN interfaces (A) of Infinity C500 is permitted exclusively for service purposes.

Only connect the following to the USB port (B):
- USB storage medium
- USB SIM card reader
- Aerogen nebulizer

WARNING

Do not simultaneously touch the connectors of the interfaces and the patient. Risk of electric shock.

WARNING

Risk of voltage surges and device malfunction

Do not connect a device that has its own power supply to the USB port, e.g., a printer or external hard drive.
Preparation and preparation

Preparing the ventilation unit

Preparing the Infinity ID neonatal expiratory valve

WARNING
Only use properly reprocessed expiratory valves which have been sufficiently dried. Otherwise the proper functioning of the device may be impaired and the patient endangered.

The expiratory valve is mounted and then inserted into the ventilation unit.

Fitting the Infinity ID neonatal expiratory valve

1. Fit the diaphragm (A) onto the edge of the expiratory valve housing. Make sure that the diaphragm is fitted properly.
2. If the muffler (B) has been removed, fit the muffler.
3. Fit the water trap container (C).

Open the flap
Open the flap (D) before inserting the expiratory valve.

- Open the flap (D) by lifting the lower edge upwards.
Assembly and preparation

Inserting the expiratory valve into the ventilation unit

1. Turn the locking ring (E) as far as possible to the left.
2. Push the expiratory valve into the fitting.
3. Turn the locking ring (E) as far as it will go to the right until it clicks audibly into place.
4. Check that it is properly secured by gently pulling on the expiratory valve.

Closing the flap

When the Infinity ID neonatal expiratory valve and the muffler are fitted, tilt the flap (D) downwards.

Leave the flap closed during ventilation.
Safety information for the use of HMEs, bacterial filters, and breathing circuits

**WARNING**

**Increased resistance**

Medication nebulization and active humidification may increase the resistance of additional components.

Check the breathing circuit regularly for signs of increased resistance and replace additional components if necessary.

**CAUTION**

Additional components in the breathing circuit such as bacterial filters, HME or CO₂ cuvettes may increase the dead space, compressible volume, and resistance.

Particular care and monitoring are required when using additional components.

Before checking the breathing circuit (see chapter “Getting started”), attach all necessary additional components up to the patient connector.

Additional components in the breathing circuit can increase the inspiratory and expiratory breathing resistance and exceed standard requirements. Examples: Inspiratory and expiratory bacterial filters, HMEs.

Babylog VN500 is designed to minimize the patient’s work of breathing. Operation does therefore not require inspiratory or expiratory bacterial filters. The use of bacterial filters or HMEs requires particular care and monitoring by the user. Especially during medication nebulization and humidification, the resistance of the expiratory bacterial filter may increase gradually.

A higher breathing resistance leads to a greater work of breathing and trigger effort. Under unfavorable conditions, this can lead to an intrinsic PEEP, which can be recognized by the fact that the expiratory flow does not return to "baseline" at the end of expiration. If the PEEP is unacceptably high, this is indicated by an alarm. For additional information, see “Automatic alarm limits” on page 282.

The breathing resistance in the patient connector cannot be monitored directly by Babylog VN500. For this reason:

- Before starting ventilation, determine in standby mode inspiratory and expiratory breathing resistance in the breathing circuit by means of the breathing circuit check.
- Check the condition of the patient and the device's measured values for volume and resistance more frequently.
- Observe the instructions for use for the HMEs, bacterial filters and breathing circuits in use.
Preparation of the breathing gas humidifier
- Prepare the Fisher & Paykel MR850 breathing gas humidifier in accordance with the corresponding instructions for use.

**CAUTION**
Do not use an HME together with a breathing gas humidifier! This can lead to an increased breathing resistance.

Connecting the Fisher & Paykel MR 850 breathing gas humidifier
The breathing gas humidifier can be connected in the following ways:
- on the standard rail of the universal holder
- on the humidifier holder of the trolley
- on the humidifier holder for the lateral standard rail

Connecting the breathing gas humidifier on the universal holder with standard rail

- Clamp the breathing gas humidifier to the standard rail (A) under the ventilation unit and screw firmly into place.

Connecting the breathing gas humidifier to the humidifier holder of the trolley

- Connect the breathing gas humidifier to the humidifier holder of the trolley.
- Tilt the breathing gas humidifier into the correct position.

Attaching the breathing gas humidifier to the humidifier holder for the lateral standard rail
If a compressor is used on the trolley, use the humidifier holder for the lateral standard rail. The holder can be connected to the left-hand or right-hand side of the device.

1. Hook the holder on the lateral standard rail (B) of Babylog VN500. Position the holder on the standard rail so that the flap at the side of the unit can still be opened.
2. Turn the clamping screw (C) until the holder is fixed securely on the rail.
3. Attach the breathing gas humidifier to the mount (D).
Assembly and preparation

Additional information
For the order numbers of the holder for the breathing gas humidifier, see the list of accessories.

Connecting the breathing circuit

WARNING
Do not use antistatic or conductive breathing hoses. The use of these materials increases the risk of electric shock to the patient and of fire in an oxygen-enriched environment.

CAUTION
The sterile packaging of disposable articles must only be opened immediately before use. Otherwise there is a risk of infection.

1 Hang the hinged arm (A) on the lateral standard rail of Babylog VN500 and tighten the screws. Depending on the desired position of the device in relation to the bed, the hinged arm can be fitted to either side of Babylog VN500.

2 Connect breathing hoses to the inspiratory port (B) and to the expiratory port (C).

3 Turn the inspiratory port and expiratory port in the direction of hoses.

A water trap is required for the Fisher & Paykel MR 850 breathing gas humidifier depending on the breathing circuit used.

4 If a water trap is required, install the water trap (D) in a vertical position.

5 Connect the Y-piece (E) to the breathing hoses.

6 Insert the Y-piece or the breathing hoses in the opening of the hinged arm.

Using the Infinity ID breathing circuit

Babylog VN500 recognizes the use of an Infinity ID breathing circuit. The message Infinity ID breathing circuit detected is displayed in the header bar.

The following Infinity ID functions are supported:
– Detection of reversed hoses
– Detection of non-compliance with the settings for the breathing circuit, patient category or humidification type
– Automatic configuration of breathing circuit and humidifier

Automatic configuration of the breathing circuit and the humidifier is only supported in standby mode.

• Fit the Infinity ID breathing hoses in standby mode.

If accessories without RFID functionality are combined with Infinity ID accessories, Infinity ID functions may be restricted or unavailable.

CAUTION
Do not reverse the connections for inspiration (B) and expiration (C). Humidification is ineffective if the connections are reversed.
Setting the breathing circuit

Babylog VN500 supports the user in selecting the breathing circuit on the Start/Standby > Br. circuit/Humidifier page.

- Set the breathing circuit according to the patient category.

Whenever the breathing hoses or the breathing gas humidifier have been changed

- Check the breathing circuit, see "Performing the breathing circuit check" on page 74.

Additional information

For the order numbers of the breathing circuits and the hinged arm, see the list of accessories.

Installing a neonatal flow sensor

**WARNING**

Risk of fire

Residual vapors of easily flammable disinfectants (e.g., alcohols) and deposits that were not removed during reprocessing can ignite when the flow sensor is in use.

- Ensure particle-free cleaning and disinfection.
- After disinfection, allow the flow sensor to air for at least 30 minutes.
- Before inserting the flow sensor check for visible damage and soiling, such as residual mucus, medication aerosols, and particles.
- Replace flow sensors when damaged, soiled, or not particle-free.

The following neonatal flow sensors are available:

- Neonatal flow sensor ISO 15 (8411130)
- Neonatal flow sensor Y-piece (8410185)

If a neonatal flow sensor and HME are used in the Neo. or Ped. pat. patient categories, the HME must be installed between the neonatal flow sensor and the patient connector.

Installing a neonatal flow sensor ISO 15 (8411130)

1. Insert the neonatal flow sensor (A) into the patient connector of the Y-piece (B).
2. Connect plug (C) of the flow sensor cable to the flow sensor.

Installing a neonatal flow sensor Y-piece (8410185)

1. Connect Y-piece with integrated neonatal flow sensor (D) to the breathing hoses.
2. Connect plug (E) of the flow sensor cable to the flow sensor.
Assembly and preparation

Further procedure for both neonatal flow sensors

3 Position patient connector of the Y-piece to point approx. 45° downwards to prevent condensation from forming on the neonatal flow sensor.

4 Run the cables along the breathing hoses to the device.

5 Insert the connector (F) of the flow sensor cable into the socket (G) at the rear of Babylog VN500.

Additional information

For the order numbers of the neonatal flow sensor, see the list of accessories.

Replacing the neonatal flow sensor insert

If Babylog VN500 displays the alarm message Neonatal flow sensor?, the insert of the neonatal flow sensor must be replaced.

1 Disconnect plug (A) of the flow sensor cable from the neonatal flow sensor.

2 Gently press the knobs (B) on both sides while pulling the insert (C) out of the flow sensor housing.

3 Push in new insert (C) until it engages.

4 Connect plug (A) of the flow sensor cable to the neonatal flow sensor.

5 Calibrate the neonatal flow sensor, see page 143.
Installing a CO₂ cuvette and CO₂ sensor

Do not carry out CO₂ measurements on premature infants because the CO₂ cuvette significantly increases the dead space.

1. Insert the cuvette (A) into the patient connector of the Y-piece. The cuvette windows are facing to the side.

2. Fit the CO₂ sensor (B) on the cuvette. The cable is facing towards the device.

3. Insert the connector (C) of the CO₂ sensor into the socket (D) at the rear of Babylog VN500.

4. Select the cuvette type, see page 147.

Additional information

“Information on checking the CO₂ sensor” on page 148.

For the order numbers of the accessories for the "CO₂ monitoring" application, see the list of accessories.
Assembly and preparation

Connecting the mains power supply to Babylog VN500

**WARNING**
Risk of electric shock and of device failure
If the device is connected to a power socket with incorrect mains voltage or without a protective ground, the user can be injured and the device damaged.

Only connect the power cable to power sockets with a protective ground and the correct mains voltage. Observe the technical data.

**NOTE**
The mains power socket in use must be freely accessible during operation.

The mains voltage must conform to the voltage range specified on the rating plate (100 V to 240 V, 50/60 Hz).

1. Plug the appliance socket (A) onto the appliance connector (B).

2. Position the power cable (C) in the clamp (D). Fit the clamp into the housing (E). Tighten the screw (F) (stress relief).

- Insert the mains plug into the mains power socket.

The LED on Infinity C500 lights up green.

Checking the toggle switch on Babylog VN500

**CAUTION**
Do not press the toggle switch during ventilation.

Prerequisite: The flap on the left-hand side of the device is opened.

- Check whether the toggle switch (G) is set to (on).
- If the toggle switch is set to (off), set it to (on).

**WARNING**
Do not simultaneously touch the connectors of the interfaces (H) and the patient. Risk of electric shock.
Assembly and preparation

Connecting the mains power supply to the GS500 gas supply unit

If Babylog VN500 is not equipped with the PS500 power supply unit, and the GS500 gas supply unit features a separate mains plug:

- Plug the mains plug of the gas supply unit into the mains socket.

The LED on GS500 lights up green.

Power supply from batteries

Charge the batteries completely before initial use.

If the mains power fails, operation is maintained either via the internal battery of Babylog VN500 or via the PS500 power supply unit.

Additional information

For additional information, see "Mains power supply / DC power supply" on page 118.

Failure of the power supply

If the mains power fails, operation is maintained via batteries.

If the mains power fails and the batteries are discharged, Babylog VN500 issues a power failure alarm.

The following data are retained even in the event of a power supply failure:
- Set values for ventilation
- Alarm limits
- Set values for monitoring

When the power supply is restored, the device starts automatically with the previous values.

Potential equalization

Differences in electrical potential between devices can be reduced by potential equalization. Potential equalization does not replace the protective ground connection. During operation, the potential equalization connections must be readily accessible and must be removable without tools.

Connecting the potential equalization cable

1. Plug one end of the potential equalization cable fully on to the potential equalization pin on Babylog VN500.
2. Connect the other end of the potential equalization cable to the hospital potential equalization socket.
Assembly and preparation

Connecting the gas supply

<table>
<thead>
<tr>
<th>WARNING</th>
<th>Do not bring any oxygen supply components into contact with oil and grease. Danger of explosion through spontaneous ignition!</th>
</tr>
</thead>
<tbody>
<tr>
<td>WARNING</td>
<td>Only use compressed gases approved for medical use. The compressed gases must be free of dust and oil particles and dry. Otherwise the proper functioning of the device cannot be ensured.</td>
</tr>
</tbody>
</table>

Central gas supply

Prerequisite: The flap on the right-hand side of the device is opened.

1. Screw the Air compressed gas hose to the **Air** (A) connection and the **O2** compressed gas hose to the **O2** connection (B) of Babylog VN500.

2. Plug the probes into the wall terminal units of the central gas supply system.

3. Position the compressed gas hoses over the hose hooks (C). The gas delivered through compressed gas hoses is used as fresh gas (FRESH GAS).

Additional information

For the order numbers of the compressed gas hoses, see the list of accessories.

Gas supply from cylinders

If the central gas supply system fails or is not available, the gas can be supplied from cylinders.

Additional information

Air supply from a gas supply unit (GS500), see "GS500 gas supply unit" on page 110.
Connecting the nurse call

The nurse call is used for transmitting high-priority alarm messages (warning) to a central hospital alarm system.

Safety information for using the nurse call

WARNING
Risk of patient injury
All data transferred via the nurse call are for information only and must not be used as the sole basis for diagnostic or therapeutic decisions. The nurse call is not intended for use with a distributed alarm system conforming to IEC 60601-1-8:2012.

CAUTION
A fault in any of the components in the link between the nurse call and the central hospital alarm system (e.g., in the unit’s electronics for nurse call, in the unit's power supply or in the alarm generator of the central hospital alarm system) can result in failure of the nurse call.

CAUTION
Connection of a nurse call does not relieve staff of their duty to check the monitoring on the device screen at regular intervals. Screen displays must be checked regularly.

CAUTION
All alarms on Babylog VN500 must be checked regularly even when the nurse call is connected. Do not use nurse call as the sole source of alarm information!

Connecting the nurse call to the central hospital alarm system

- The nurse call cable must be connected to the lead to the central hospital alarm system by service personnel.

As soon as Babylog VN500 signals an alarm, the connection between the white cable and the brown cable (NO and COM) is closed and the nurse call is activated.

Connecting the nurse call to the ventilation unit

Prerequisite: The flap on the left-hand side of the device is opened.

1. Plug the nurse call connector (A) into the socket (B) until it engages audibly.

NOTE
The connector must engage audibly into the socket to ensure all alarm messages are transmitted properly.
Assembly and preparation

2 Check the correct operation of connected nurse call system.

Information on the nurse call
High-priority alarm messages (warning) are transmitted to a central hospital alarm system. Medium-priority (caution) and low-priority (note) alarm messages are not transmitted.

The nurse call is also activated when the internal acoustic alarm generator in the device is defective.

If, in the event of an alarm, the key (Audio paused) is pressed, the acoustic alarm on the device and the nurse call are suppressed for 2 minutes.

Additional information
For the order number of the nurse call cable, see the list of accessories.

Closing the flaps at the side of the device
- Close the lateral flaps of the device after preparation.

CAUTION
Keep both lateral flaps on the device closed during operation to prevent accidental actuation of the toggle switch or connections becoming loose.

Intrahospital transport

Transport refers to any movement of the medical device without the patient that does not serve to position the medical device.

Increasing the tipping stability
- Swivel the control and display unit (Medical Cockpit) until it is centrally aligned with the ventilation unit.
- Set the hinged arms to minimum extension.
- Drain the water container of the breathing gas humidifier.
- Secure the breathing gas humidifier to the trolley, not to the lateral standard rails of the ventilation unit.
- Do not attach any additional parts to the lateral standard rails of the ventilation unit.
- If fitted, slide the bed coupling into its retracted position.
- Grasp the trolley handle firmly and push the device in longitudinal direction.

The safety information regarding intrahospital patient transport also applies, see chapter "Intrahospital patient transport" on page 121.
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Safety information on getting started

**WARNING**
Ventilation does not take place in standby mode! The device must only be set to standby mode when no patient is connected to the device. The patient may otherwise be jeopardized.

**CAUTION**
Condensation may form when the device is moved from a cold storage location to a warm environment. Do not switch on the device as otherwise its proper functioning may be adversely affected. Wait until the condensation has dried.

Switching on Babylog VN500 and Infinity C500

Prerequisites:
– Babylog VN500, Infinity C500, PS500 and GS500 are reprocessed and assembled ready for operation.
– The mains power supply and the gas supply are connected.
– The Babylog VN500 toggle switch is set to (on).
– Press the key (A) on Infinity C500.

Babylog VN500 provides you with two options:
– Using the settings of the previous patient (B)
– Admitting a new patient (C)

If the Start dialog is closed using the X button (D), Babylog VN500 adopts the settings of the previous patient.

If a data loss occurs, the previous settings cannot be recovered. The Current patient button (B) is not displayed.

The system is started. The Start dialog is displayed.
Selecting a patient

Using the settings of the previous patient

Prerequisite: The Start dialog is opened.

- Touch the Current patient button (A).

The last used patient-related settings including the alarm limits, application mode and device status are restored. O2 monitoring and flow monitoring are switched on, see "Information on monitoring" on page 142.

The Start/Standby page (B) is displayed. Babylog VN500 is in standby mode.

Admitting a new patient

For a new patient, Babylog VN500 determines the ventilation parameters' start-up settings based on the patient category (factory setting) or the body weight. The factory settings for the settings dependent on patient category and weight can be changed in the System setup dialog window.

The patient category or the body weight can only be changed when a new patient is admitted. In the Ped. pat. patient category, the body height is entered and the ideal body weight is determined from this. In the Neo. patient category, the body weight is entered directly. The weight-dependent setting for a new patient is only possible after selecting Weight in the System setup dialog window.

The alarm limit start-up settings are recalculated according to the customized system configuration.

When a new patient is admitted, the settings and trend data of the previous patient are deleted.

Prerequisite: The Start dialog is opened.

1. Touch the following button for a new patient:
   - New Ped. pat. (A) for new pediatric patients
   - New Neo. (B) for new neonatal patients

   The respective button turns yellow.

2. Confirm with the rotary knob.

The Start/Standby page is displayed. Babylog VN500 is in standby mode.
Ventilation parameter start-up settings by patient category

The Start/Standby page (C) contains the buttons for the patient category:

- New Ped. pat. (D)
- New Neo. (E)

1. Touch the button for the desired patient category (D) or (E).
2. Confirm with the rotary knob.

The ventilation parameters displayed in the lower part of the page (F) are the start-up settings for the selected patient category.

Determining the start-up settings can take up to 5 seconds. No entries can be made during this time.

Ventilation parameter start-up settings by body height/body weight

Prerequisite: In the System setup dialog window, the Weight function was configured and a new patient was admitted.

In the Ped. pat. patient category, the Start/Standby page (C) contains the button for body height (G) and the field for the ideal body weight (H).

1. Touch the button for the body height (G).
2. Set the body height by turning the rotary knob and push to confirm.

Babylog VN500 determines the start-up values for VT, RR, Slope and Flow trigger based on the ideal body weight calculated from the body height. The values for VT and RR are displayed in the lower part of the page (F). The other ventilation parameters displayed in the lower part of the page are start-up settings for the selected patient category.
Getting started

In the Neo. patient category, the patient's body weight is set directly. The Start/Standby page (C) contains the button for this start-up body weight (I).

1. Touch the button for the start-up body weight (I).
2. Using the rotary knob, set the start-up body weight and confirm the value.

The button for the current body weight (J) is displayed. After the patient has been admitted, the current body weight corresponds to the start-up body weight.

Babylog VN500 determines the start-up values for VT, RR, Slope and Flow trigger based on the start-up body weight. The values for VT and RR are displayed in the lower part of the page (F). The other ventilation parameters displayed in the lower part of the page are start-up settings for the selected patient category.

Determining the start-up settings can take up to 5 seconds. No entries can be made during this time.

Setting the body weight during ventilation

As a result of setting the ideal body weight in the Ped. pat. patient category or the current body weight in the Neo. patient category, measurements are displayed relative to the body weight, e.g. VT/kg BW.

Setting the body weight is only possible on the Start/Standby page during ventilation.

In the Ped. pat. patient category:
1. Touch the button for the ideal body weight.
2. Using the rotary knob, set the ideal body weight and confirm the value.

In the Neo. patient category:
1. Touch the button for the current body weight.
2. Using the rotary knob, set the current body weight and confirm the value.

Whenever the patient category has been changed

Check the breathing circuit, see chapter "Performing the breathing circuit check" on page 74.

Additional information

The configuration for the ventilation parameter start-up values by body height/body weight or by patient category is entered on the System setup > Ventilation > Start settings page. See chapter "Configuring start-up settings for the ventilation parameters" on page 171.

For information on configuring customized alarm limits, see chapter "Setting start-up values for alarm limits" on page 165.

For information on starting the therapy, see chapter "Starting therapy" on page 80.
Getting started

Selecting the breathing circuit and the breathing gas humidifier

The breathing circuit and the breathing gas humidifier can only be selected in standby mode.

1 Touch the Start/Standby... button in the main menu bar.
2 Touch the Br. circuit/Humidifier tab (A).

The page for selecting the breathing circuit and the breathing gas humidifier is displayed.

Selecting the breathing circuit from the selection list

3 Touch the button (B).
4 Select the breathing circuit used from the selection list.
5 Confirm with the rotary knob.

To help with the selection, the selected breathing circuit is displayed as a detailed representation (C) and also described as text (D).

Babylog VN500 automatically selects the appropriate humidification type based on the breathing circuit (E) selected. Some breathing circuits provide the selection of HME/Filter and None.

If the breathing circuit used is not included in the selection list

1 Touch the button (B).
2 Select Other from the selection list.
3 Confirm with the rotary knob.
4 Select the humidification type (E):
   – Active humid., exp. unheated
   – Active humid., exp. heated
   – HME/Filter
   – None

   Touch the corresponding button.

Using the user-defined breathing circuit

Prerequisite: The User-defined hose settings function is enabled, see page 169.

1 Touch the button (B).
2 Select User-defined breathing circuit from the selection list.
3 Confirm with the rotary knob.
4 Select the humidification type (E).
5 Perform the breathing circuit check, see page 74.
6 Save the measured values for hose compliance and hose resistance, see page 76.
Infinity ID breathing circuits

When using Infinity ID breathing circuits, the connected hose type as well as the corresponding humidification type are set automatically.

If the message *Infinity ID breathing circuit detected.* is not displayed when an Infinity ID breathing circuit is connected, use a different Infinity ID breathing circuit. If the message is still not displayed, replace the Infinity ID neonatal expiratory valve or inspiratory valve.

Whenever the breathing circuit or the breathing gas humidifier have been changed

- Check the breathing circuit, see "Performing the breathing circuit check" on page 74.
Checking readiness for operation

The system check consists of the following elements:
- Device check
- Breathing circuit check
- Battery check

The battery check must be performed during initial commissioning of the device. For further information on the battery check, see page 250 and page 254.

Safety information on the system check

**WARNING**
Before using on the patient:
- Perform the device check. If a malfunction is detected, do not operate the device! Patient hazard!
- Perform the breathing circuit check to ensure the pressure measurement accuracy. Otherwise the airway pressure may deviate from the set values.

Starting the system check

The system check is only possible in standby mode.

1. Touch the **Start/Standby...** button in the main menu bar.
2. Touch the **System check** tab (A).

Babylong VN500 displays the following on the **Overview** page (B):
- Last device check with date, time, and result (C)
- Last breathing circuit check with date, time, and the amount of leakage determined (D)
- Battery check, result of the last battery check, and date for the next battery check (E)
Performing the device check

The device check is only possible in standby mode.

Keeping the test lung ready

- Pediatric test lung (8409742) for the pediatric and neonatal breathing circuit

The test lung must only be inserted into the patient connector of the Y-piece after instruction by Babylog VN500.

Starting the device check

Prerequisites: The medication nebulizer is not connected. The System check page (A) is opened.

1. Touch the Device check tab (B).
2. Touch the Start button (D).
3. Confirm with the rotary knob.

Test steps in the device check

In the device check the following test steps are performed:
- Auxiliary acoustical alarm (Check of the auxiliary alarm/power failure alarm)
  - There is no need for the user to test other parts of the alarm system, as they are tested in the self-test.
- Breathing circuit connection (visual inspection of breathing circuit)
- Inspect humidifier (visual inspection of breathing gas humidifier)
- CO\textsubscript{2} sensor: Zero calibration
- Neonatal flow sensor: Calibration
- Neonatal flow sensor: Measurement
- Test lung connection
- Gas supply sensors: Calibration
- O\textsubscript{2} supply
- Air supply
- Gas supply unit (if the gas supply unit function is activated)
- Pressure sensor calibration valve
- Expiratory valve (expiratory valve check)
- Safety valve (safety function check)
- O\textsubscript{2} sensor: Calibration
- Nebulizer (medication nebulizer control check)
- Ejector (functional check)
Device check procedure

Babylog VN500 guides the user in the form of a question/answer dialog through the respective test step. The instruction field (E) displays the questions or instructions how to carry out the test steps.

The questions must be answered by touching the Yes (F) or No (G) buttons.

The Next test button (H) can be used to skip the test steps.

A test step is also skipped if the necessary prerequisites have not been met.

The test steps in the device check are displayed with the following symbols:

<table>
<thead>
<tr>
<th>Rotating symbol</th>
<th>Active test step</th>
</tr>
</thead>
<tbody>
<tr>
<td></td>
<td></td>
</tr>
<tr>
<td>Green dot</td>
<td>Correct result</td>
</tr>
<tr>
<td>Red dot</td>
<td>Incorrect result</td>
</tr>
<tr>
<td>Colorless dot</td>
<td>Test step not performed</td>
</tr>
</tbody>
</table>

Repeating test steps in the device check

1. Touch the Repeat button (I).
2. Confirm with the rotary knob.

All test steps that have not yet been performed or that were unsuccessful are repeated.

Aborting the device check

1. Touch the Cancel button (J).
2. Confirm with the rotary knob.

The device check is also canceled when the Device check page is closed. The device check can be continued when the Device check page is opened again.

1. Touch the Repeat button (I).
2. Confirm with the rotary knob.

Test results

The test results obtained from the device check and the calibration and zero-checking values of the sensors remain stored until the next calibration, even if the device is switched off.

Incorrect test steps and remedies

Errors in the following safety-relevant test steps generate the medium-priority alarm message Device check failed:

- Pressure sensor calibration valve
- Expiratory valve
- Safety valve

The alarm cannot be acknowledged. Do not start ventilation!

Errors in non-safety-relevant test steps or test steps that are not performed due to a prerequisite generate the low-priority alarm message Device check incomplete.

The alarm causes and their remedies are displayed on the Current alarms page.
Getting started

The following table shows the remedies for eliminating the errors during the device check:

<table>
<thead>
<tr>
<th>Test step</th>
<th>Remedy</th>
</tr>
</thead>
<tbody>
<tr>
<td>Auxiliary acoustical alarm</td>
<td>Contact DrägerService.</td>
</tr>
<tr>
<td>CO\textsubscript{2} sensor: Zero calibration</td>
<td>Check whether the CO\textsubscript{2} sensor is connected. Wait for the CO\textsubscript{2} sensor to complete its three-minute warm-up phase. Check whether the CO\textsubscript{2} sensor or the cuvette is soiled.</td>
</tr>
<tr>
<td>Neonatal flow sensor: Calibration</td>
<td>Clean the flow sensor. Seal the flow sensor during calibration. Check whether the flow sensor cable is connected.</td>
</tr>
<tr>
<td>Gas supply unit</td>
<td>Check whether the gas connection to the device is kinked. Check whether the data cable is connected. If GS500 is running continuously, shut down and switch off Babylog VN500 (toggle switch to ( \odot )).</td>
</tr>
<tr>
<td>Gas supply sensors: Calibration</td>
<td>Check whether the compressed gas hoses are connected. Shut down Babylog VN500 and switch it off (toggle switch to ( \odot )).</td>
</tr>
<tr>
<td>O\textsubscript{2} supply</td>
<td>Check whether the O\textsubscript{2} compressed gas hose is connected.</td>
</tr>
<tr>
<td>Air supply</td>
<td>Check whether the Air compressed gas hose is connected.</td>
</tr>
<tr>
<td>Pressure sensor calibration valve</td>
<td>Connect the test lung. Check the breathing circuit for leaks. Check whether the compressed gas hoses are connected. Check whether the expiratory valve is properly engaged.</td>
</tr>
<tr>
<td>Expiratory valve</td>
<td>Check whether the water trap is connected. Check whether the expiratory valve is properly engaged.</td>
</tr>
<tr>
<td>Safety valve</td>
<td>Connect the test lung. Check the breathing circuit for leaks. Check whether the compressed gas hoses are connected. Check whether the expiratory valve is properly engaged.</td>
</tr>
<tr>
<td>O\textsubscript{2} sensor: Calibration</td>
<td>Check whether the compressed gas hoses are connected.</td>
</tr>
<tr>
<td>Nebulizer</td>
<td>Prerequisite: The medication nebulizer is not connected. Check whether the compressed gas hoses are connected.</td>
</tr>
<tr>
<td>Ejector</td>
<td>Check whether the compressed gas hoses are connected.</td>
</tr>
</tbody>
</table>

- Eliminate the causes of the error and repeat the test step.
- If the test step still fails, contact DrägerService.
Calibrating the gas supply sensors
The calibration of the gas supply sensors takes approximately 2 minutes. This test step must be performed every 3 months. If a test step does not need to be performed, it can be skipped by pressing No. The test step is still displayed as "successfully completed" (green dot).

If the test step is skipped with Next test, the test step is displayed as "not performed" (colorless dot).

If a complete calibration is necessary after 3 months and the test step is skipped with Next test, the test step is displayed as "failed" (red dot).

Calibrating the O₂ sensor
The O₂ sensor is calibrated during each device check. Regular calibration of the O₂ sensor ensures the specified accuracy.

If the test step is skipped with Next test and the O₂ sensor is not calibrated for 3 months, the accuracy of the O₂ sensor will be reduced. In the parameter field for FiO₂, a question mark will be displayed next to the measurement.

After calibration during the device check the sensor will work again with full accuracy. The measured value is displayed in the parameter field.

If the test step is skipped with Next test, the test step is displayed as "not performed" (colorless dot).

If Babylog VN500 requires the O₂ sensor to be calibrated and the test step is still skipped with Next test, the test step is displayed as failed (red dot).

CAUTION
If the quality of the oxygen from the central gas supply system is inadequate, calibrate the O₂ sensor with an appropriate calibration gas (100 % O₂). Otherwise this may result in an incorrect calibration.

After the device check
The user is requested in field (E) to perform the check of the breathing circuit.

Confirm with Yes (F).

The page for the breathing circuit check is opened.

Performing the breathing circuit check
The check is only possible in standby mode.

The breathing circuit check must be performed after:
- Device check
- Changing the breathing circuit
- Changing the breathing gas humidifier
- Changing the patient category

Test steps during the breathing circuit check
The following test steps are performed:
- Leakage of the breathing circuit
- Compliance of the breathing circuit
- Insp. Resistance
- Exp. Resistance
Starting the breathing circuit check

Prerequisite: The System check page (A) is opened.

1. Touch the Breathing circ. check tab (B).

The values of the last test are displayed (C). If a valid measurement has not yet taken place, the standard values are displayed.

2. Touch the Start button (D).

3. Confirm with the rotary knob.

4. When requested by Babylog VN500 in the instruction field (E): Seal the patient connection port, e.g., with a sterile glove. Confirm with OK (F).

5. When requested, open the patient connection port. Confirm with OK (F).

The current leakage flow is displayed continuously throughout the test. A leakage flow of up to 300 mL/min at a pressure of 60 mbar (60 cmH₂O) is acceptable.

After the leakage test, Babylog VN500 determines the compliance and the inspiratory and expiratory resistance of the breathing circuit.

When changing the breathing circuit and type of humidifier, Babylog VN500 automatically resets the values for hose compliance and hose resistance to default values.

When using Infinity ID breathing circuits, the default values of the breathing circuit detected are used. The leakage measurement becomes invalid.

When the patient category is changed, the breathing circuit that was last used in this category is selected and the corresponding values for hose compliance and hose resistance are used.

The leakage measurement becomes invalid when a new patient is admitted to the same patient category. The values for hose resistance and hose compliance are retained.

It is recommended to perform the breathing circuit check before commencing patient ventilation with a newly started device.

Canceling the breathing circuit check

1. Touch the Cancel button (G).

2. Confirm with the rotary knob.

The leakage measurement becomes invalid. The values for hose resistance and hose compliance are reset to the default values.
Repeating the breathing circuit check

If the breathing circuit is changed after the breathing circuit check, the humidification type or the patient category is changed, the breathing circuit check will have to be repeated.

The breathing circuit check is also necessary when using Infinity ID breathing circuits.

User-defined breathing circuit

Prerequisite: The user-defined breathing circuit has been selected, see “Using the user-defined breathing circuit” on page 68.

The values for hose resistance and hose compliance can be saved and are then available when that breathing circuit is selected again.

Display of results of the breathing circuit check

Prerequisite: The System check page (A) is opened.

- Touch the Check results tab (I).

The detailed results of the check are displayed:
- Compliance [mL/mbar] (J)
- Flow [L/min] (K)
- Inspiratory resistance [mbar/L/s] (L)
- Expiratory resistance [mbar/L/s] (M)

Display of the results of the system check on the Start/Standby page

After the system check, the results are displayed on the Start/Standby > Start/Standby page.

- Result of device check (N)
- Result of breathing circuit check (O)
Checking the switch-over to battery operation

1. Unplug the power plug.
   If there is a PS500 power supply unit present, the device switches over to the PS500 without interruption. If there is no PS500 present or the PS500 is discharged, the device switches over to the internal battery without interruption. The **Battery activated** alarm is displayed.

2. Plug the power plug back in.
   The device switches back to mains operation. The **Battery activated** alarm message goes out.

Checking the alarm signaling

When the system check has been successfully completed, the device is ready for operation. The alarm signaling can be checked additionally.

The description of alarm signaling can be found in chapter “Alarms.” Additional information on alarm criteria can be found in chapter “Alarm – Cause – Remedy.”

High-priority alarm message

1. Start ventilation.

2. After 2 minutes set the upper alarm limit for $MV_e$ to a value below the measured value of $MV_e$.
   The **MV high** alarm is triggered.

Medium-priority alarm message

1. Start ventilation.

2. Switch on the additional setting **Volume Guarantee**.

3. Reduce the upper alarm limit for Paw until the **VT not reached, Pmax active** alarm is triggered.

Low-priority alarm message

1. Start ventilation.

2. In the **Special maneuvers > Maneuvers** dialog window, touch and hold the **Man. insp./hold** button until the **Inspiratory hold interrupted** alarm is triggered.

Checking alarm limits

The alarm limits for a settable alarm can be checked by setting the alarm limits appropriately. When the alarm limit is exceeded, the corresponding alarm is triggered.

Additional information on setting alarm limits can be found in chapter “Setting alarm limits” on page 128.

Test of the acoustic alarm system

The acoustic alarm system need not be tested by the user. The device tests the functions of the acoustic alarm system automatically during the device check.
Getting started

Selecting the Tube or NIV application mode

Babylog VN500 can switch between non-invasive ventilation and tube ventilation.

The application mode can only be selected in standby mode.

1 Touch the **Start/ Standby**... button in the main menu bar.

2 Touch the **Tube/NIV** tab (A).

3 Touch the **Tube** (B) or **NIV** button (C).

4 Confirm with the rotary knob.

Observe the information on changing the application mode!

**CAUTION**
Application mode **NIV** must not be activated with intubated patients.

**WARNING**
Alarm limits and ventilation settings must be checked or set again in order to ensure complete monitoring of ventilation after changing from **NIV** application mode to **Tube** application mode.

Setting parameters for the tube

The inner diameter of the tube and the tube type can be entered for the following functions:

- Display of $P_{trach}$, independent of ATC
- Measurement of patient resistance $R_{pat}$ and the index $C_{20/C_{dyn}}$

If the inner diameter of the tube and the tube type are entered, the measured value $R_{pat}$ corresponds to the patient resistance. $R_{pat}$ and $C_{20/C_{dyn}}$ are only displayed correctly if the inner diameter of the tube and the tube type are entered correctly. The measured value $R$ always corresponds to the total resistance.

Prerequisite: The **Tube/NIV** page (A) is opened. The **Tube** application mode has been selected.

Activating or deactivating the calculation of tracheal pressure

1 Touch the appropriate button (B).

2 Confirm with the rotary knob.

If ATC is switched off, the calculation of tracheal pressure is always deactivated when a new patient is admitted.

Additional information

For information on using the **NIV** application mode for non-invasive ventilation, see "**NIV – Non-invasive ventilation**" on page 92.
Selecting the tube type

In the Neo. patient category, this selection is not available.

1 Touch the appropriate button (C).
2 Confirm with the rotary knob.

Entering the inner diameter of the tube

1 Touch the button (D).
2 Set the value by turning the rotary knob and push to confirm.

Selecting the therapy type

Babylog VN500 can choose between therapy types **Ventilation** and **O2 Therapy**.

The therapy type can only be changed in standby mode.

1 Touch the Start/Standby... button in the main menu bar.

The Start/Standby page (A) is displayed.

<table>
<thead>
<tr>
<th>Start/Standby</th>
</tr>
</thead>
<tbody>
<tr>
<td>A</td>
</tr>
<tr>
<td></td>
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<tr>
<td></td>
</tr>
<tr>
<td></td>
</tr>
<tr>
<td>B</td>
</tr>
<tr>
<td>C</td>
</tr>
</tbody>
</table>

2 Touch the **Ventilation** (B) or **O2 Therapy** button (C).
3 Confirm with the rotary knob.

**Additional information**

"O2 therapy" on page 111.
"Setting ventilation" on page 84.
Getting started

Starting therapy

Before using on the patient
- Carry out a system check to ensure that Babylog VN500 is operating correctly, see page 70.
- Check the therapy settings: Set the alarm limits, see page 128. Set the ventilation modes and ventilation parameters, see “Setting ventilation” on page 84.

Starting ventilation or O2 therapy
1 Touch the Start/Standby... button in the main menu bar.
   The Start/Standby page (A) is displayed.

2 Touch the Start ventilation button (B) and confirm with the rotary knob.
   Babylog VN500 starts the therapy with the set ventilation parameters. The main page for ventilation or O2 therapy is displayed.

Additional information
   The page for the ventilation settings can be opened with the Ventilation settings... button (C).
Displaying the status of accessories

1. Touch the **Start/Standby** button in the main menu bar.
2. Touch the **Accessory status** tab (A).

In field (B) Babylog VN500 displays the time until it is recommended to exchange the accessories.

Sterilization of the expiratory valve or inspiratory valve may gradually impair the operation of RFID transmission. This may mean that Infinity ID breathing circuit functions may not work or may no longer work reliably. The status of the Infinity ID accessories is not displayed.

**Additional information**

The time for the exchange interval can be configured on the **System setup > System status > Exchange intervals** page. See "System status" on page 181.
This page has been left blank intentionally.
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Instructions for use Infinity Acute Care System – Babylog VN500 SW 2.n 83
Setting ventilation

Overview

This chapter describes how to set ventilation modes and general settings as well as additional settings for ventilation parameters.

For a detailed description of the ventilation modes and ventilation parameters, see chapters "Description of the ventilation modes" on page 290 and "Additional settings for ventilation" on page 304.

Opening the Ventilation settings dialog window

The Ventilation settings dialog window can be opened as follows:

- Touch the Ventilation settings... button (A) in the main menu bar.
- Touch the button (B) in the therapy bar.
- Touch the displayed ventilation mode (C) in the header bar.

Babylog VN500 opens the Ventilation settings dialog window.

The page for the active ventilation mode (D) with the General settings tab (F) is displayed by default. The corresponding therapy controls (E) are displayed.

The tab for Additional settings (G) can be used to supplement the active ventilation mode with additional settings.

Selecting ventilation modes

Prerequisite: The General settings page (A) is opened.

The Ventilation settings dialog window contains 5 tabs for selecting the ventilation modes. 4 tabs (B) have ventilation modes permanently assigned to them. The fifth tab (C) can be used to select another ventilation mode, which can be selected from the available ventilation modes.
The following 4 ventilation modes are preset at the factory:
- PC-CMV
- PC-AC
- PC-SIMV
- PC-PSV

For information on changing the assignment of ventilation modes, see "Configuring start-up settings for the ventilation modes" on page 170.

**Selecting an additional ventilation mode in the dialog window**

1. Touch the **Other modes** tab (D).

   ![Ventilation settings](image)

   All the available ventilation modes (E) are displayed.

2. Touch the button for the corresponding ventilation mode. The color of the tab (D) turns yellow.

3. Confirm with the rotary knob.

   The additional ventilation mode is displayed in the fifth tab (C). The ventilation mode is active.

**Changing the ventilation mode**

1. Touch the corresponding tab, e.g., (F). The color of the tab turns yellow.

2. Preset the ventilation parameters if necessary.

3. Confirm with the rotary knob. The color of the tab turns dark green.

   The ventilation mode is active. The settings are applied to the patient.

**Setting ventilation parameters**

Prerequisite: The **General settings** page (A) is opened.

1. Touch the corresponding therapy control, e.g., (B).

2. Set the value by turning the rotary knob and push to confirm.

   The additional ventilation parameters derived from the ventilation parameter are calculated by Babylog VN500 and displayed in the setting assistance field (C).

   Information is displayed in the message field (D), e.g., when the setting limit of a parameter has been reached.

**Setting ventilation parameters in the therapy bar**

The ventilation parameters of the active ventilation mode can also be set with the therapy controls in the therapy bar (E).
Operation

Additional information

"Exceeding the set limit of a ventilation parameter" on page 39.

"Direct setting of ventilation parameters (QuickSet)" on page 39.

"Linked setting of ventilation parameters" on page 40.

General settings for ventilation

The general settings for the ventilation parameters are listed in the following tables:

– Pressure-controlled ventilation modes
– Spontaneous breathing support

**WARNING**
If flow measurement is deactivated for SPN-CPAP, use a separate monitoring device.

**CAUTION**
Only remove the water trap of the expiratory valve briefly during ventilation. Otherwise, ventilation will be impaired.
Pressure-controlled ventilation modes

<table>
<thead>
<tr>
<th>Ventilation parameters</th>
<th>Ventilation mode</th>
</tr>
</thead>
<tbody>
<tr>
<td></td>
<td>PC-SIMV</td>
</tr>
<tr>
<td>FiO2</td>
<td>X</td>
</tr>
<tr>
<td>VT</td>
<td>X(^{1})</td>
</tr>
<tr>
<td>Ti</td>
<td>X</td>
</tr>
<tr>
<td>RR</td>
<td>X</td>
</tr>
<tr>
<td>Slope or Insp. flow(^{2})</td>
<td>X</td>
</tr>
<tr>
<td>Pmax</td>
<td>X(^{3})</td>
</tr>
<tr>
<td>Psp</td>
<td>X(^{4})</td>
</tr>
<tr>
<td>PEEP</td>
<td>X</td>
</tr>
<tr>
<td>ΔPsupp</td>
<td>X</td>
</tr>
<tr>
<td>Tmax</td>
<td>X</td>
</tr>
<tr>
<td>Thigh</td>
<td>X</td>
</tr>
<tr>
<td>Tlow</td>
<td>X(^{6})</td>
</tr>
<tr>
<td>Phigh</td>
<td>X</td>
</tr>
<tr>
<td>Plow</td>
<td>X</td>
</tr>
<tr>
<td>Tlow max</td>
<td>X(^{7})</td>
</tr>
<tr>
<td>Exp. term.</td>
<td>X(^{7})</td>
</tr>
<tr>
<td>ΔEhf</td>
<td>X</td>
</tr>
<tr>
<td>fhf</td>
<td>X</td>
</tr>
<tr>
<td>Amplitude hf</td>
<td>X</td>
</tr>
<tr>
<td>MAPHF</td>
<td>X</td>
</tr>
</tbody>
</table>

1) If VG is switched on
2) Depending on the configuration of Slope adjustment
3) If Pmax/Paw high autoset is activated and ATC or Apnea Ventilation or VG is switched on
4) If VG is switched off
5) In the Neo patient category in the Tube application mode, or in the Ped. pat. patient category in the NIV application mode
6) If AutoRelease is switched off
7) If AutoRelease is switched on
Spontaneous breathing support

<table>
<thead>
<tr>
<th>Ventilation parameters</th>
<th>SPN-CPAP/PS</th>
<th>SPN-CPAP/VS</th>
<th>SPN-CPAP&lt;sup&gt;1)&lt;/sup&gt;</th>
<th>SPN-PPS</th>
</tr>
</thead>
<tbody>
<tr>
<td>FiO2</td>
<td>X</td>
<td>X</td>
<td>X</td>
<td>X</td>
</tr>
<tr>
<td>VT</td>
<td>X</td>
<td></td>
<td></td>
<td></td>
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<tr>
<td>VTmax</td>
<td></td>
<td></td>
<td></td>
<td>X</td>
</tr>
<tr>
<td>Timax</td>
<td>X&lt;sup&gt;2)&lt;/sup&gt;</td>
<td>X&lt;sup&gt;2)&lt;/sup&gt;</td>
<td>X&lt;sup&gt;4)&lt;/sup&gt;</td>
<td></td>
</tr>
<tr>
<td>Slope or Insp. flow&lt;sup&gt;3)&lt;/sup&gt;</td>
<td>X</td>
<td>X</td>
<td>X</td>
<td></td>
</tr>
<tr>
<td>Pmax</td>
<td>X&lt;sup&gt;4)&lt;/sup&gt;</td>
<td>X&lt;sup&gt;4)&lt;/sup&gt;</td>
<td>X&lt;sup&gt;4)&lt;/sup&gt;</td>
<td></td>
</tr>
<tr>
<td>PEEP</td>
<td>X</td>
<td>X</td>
<td>X</td>
<td>X</td>
</tr>
<tr>
<td>ΔPsupp</td>
<td>X</td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>Vol. Assist</td>
<td></td>
<td></td>
<td></td>
<td>X</td>
</tr>
<tr>
<td>Flow Assist</td>
<td></td>
<td></td>
<td></td>
<td>X</td>
</tr>
<tr>
<td>TmanInsp</td>
<td></td>
<td></td>
<td></td>
<td>X</td>
</tr>
<tr>
<td>PmanInsp</td>
<td></td>
<td></td>
<td></td>
<td>X</td>
</tr>
</tbody>
</table>

<sup>1)</sup> only available in the Neo. patient category in the NIV application mode
<sup>2)</sup> in the NIV application mode or in the patient category Neo.
<sup>3)</sup> depending on the configuration of Slope adjustment
<sup>4)</sup> if $P_{max}/P_{aw}$ high autoset is activated
Additional settings for ventilation

Overview of possible supplementary settings

The ventilation modes can be combined with additional settings to optimize ventilation. The table shows the possible additional settings for the respective ventilation mode.

<table>
<thead>
<tr>
<th>Ventilation mode</th>
<th>Apnea Ventilation</th>
<th>Trigger</th>
<th>Sigh</th>
<th>ATC</th>
<th>Volume Guarantee</th>
<th>Auto Release</th>
<th>HFO-Sigh</th>
<th>Volume Guarantee (HFO)</th>
</tr>
</thead>
<tbody>
<tr>
<td>PC-CMV</td>
<td>X</td>
<td>X</td>
<td>X</td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>PC-AC</td>
<td>X</td>
<td>X</td>
<td>X</td>
<td>X</td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>PC-SIMV</td>
<td>X</td>
<td>X</td>
<td>X</td>
<td>X</td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>PC-PSV</td>
<td>X</td>
<td>X</td>
<td>X</td>
<td>X</td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>PC-MMV</td>
<td>X</td>
<td>X</td>
<td>X</td>
<td>X</td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>PC-APRV</td>
<td>X</td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>SPN-CPAP/PS</td>
<td>X</td>
<td>X</td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>SPN-CPAP/VS</td>
<td>X</td>
<td>X</td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>SPN-PPS</td>
<td>X</td>
<td>X</td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>PC-HFO</td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td>X</td>
<td>X</td>
</tr>
</tbody>
</table>

Setting the supplementary settings

Prerequisite: The page with the active ventilation mode is open.

1. Touch the **Additional settings** tab (A).

   The additional settings of the active ventilation mode are displayed.

2. Touch the tab of the respective additional setting (B).

   The page for setting the corresponding parameters is opened.

3. Use the buttons (C) to activate or deactivate the additional setting.

4. Touch the corresponding therapy control (D).

5. Set the value by turning the rotary knob and push to confirm.

Instructions for use Infinity Acute Care System – Babylog VN500 SW 2.n
The **Trigger** and **Apnea Ventilation** additional settings can be configured as buttons in the main menu bar to enable direct access. See "Assigning functions to additional buttons" on page 161.

### Ventilation parameters for the additional settings

**CAUTION**
Undetected worsening of the patient's condition
During high-frequency ventilation with Volume Guarantee, the ventilator automatically compensates for changes in lung mechanics. As a result, a gradual worsening of patient condition may only be detected after a delay.

Observe patient condition. Check ventilation pressure regularly and set the alarm limits accordingly.

**CAUTION**
High trigger sensitivity may lead to auto-triggering of the ventilator.

<table>
<thead>
<tr>
<th>Additional settings</th>
<th>Ventilation parameters</th>
<th>Dependencies, information</th>
</tr>
</thead>
<tbody>
<tr>
<td><strong>Apnea Ventilation</strong></td>
<td></td>
<td></td>
</tr>
<tr>
<td>On/Off</td>
<td></td>
<td></td>
</tr>
<tr>
<td>VTapn</td>
<td></td>
<td></td>
</tr>
<tr>
<td>RRapn</td>
<td></td>
<td></td>
</tr>
<tr>
<td>Pmax</td>
<td>If <strong>Pmax/Paw high autoset</strong> is configured</td>
<td></td>
</tr>
<tr>
<td>PEEP</td>
<td>In PC-APRV</td>
<td></td>
</tr>
<tr>
<td>Flow trigger</td>
<td>In PC-APRV</td>
<td></td>
</tr>
<tr>
<td>Slope</td>
<td>In SPN-PPS</td>
<td></td>
</tr>
<tr>
<td></td>
<td></td>
<td>For configuration of the <strong>Automatic return from Apnea Ventilation</strong> function, see &quot;Configuring general settings&quot; on page 177. For a description, see &quot;Automatic return from apnea ventilation&quot; on page 305.</td>
</tr>
<tr>
<td><strong>Trigger</strong></td>
<td>Flow trigger</td>
<td></td>
</tr>
<tr>
<td><strong>Sigh</strong></td>
<td></td>
<td></td>
</tr>
<tr>
<td>On/Off</td>
<td></td>
<td></td>
</tr>
<tr>
<td>ΔintPEEP</td>
<td></td>
<td></td>
</tr>
<tr>
<td>Interval sigh</td>
<td></td>
<td></td>
</tr>
<tr>
<td>Cycles sigh</td>
<td></td>
<td></td>
</tr>
</tbody>
</table>
### Additional settings

<table>
<thead>
<tr>
<th>ATC</th>
<th><strong>Ventilation parameters</strong></th>
<th><strong>Dependencies, information</strong></th>
</tr>
</thead>
<tbody>
<tr>
<td>On/Off</td>
<td>See chapter &quot;Configuration&quot; on page 155.</td>
<td></td>
</tr>
<tr>
<td>Tube type (ET/Trach.)</td>
<td>Only available in the <strong>Ped. pat.</strong> patient category</td>
<td></td>
</tr>
<tr>
<td>Tube Ø</td>
<td>Inner diameter of the tube</td>
<td></td>
</tr>
<tr>
<td>Compens.</td>
<td>Degree of compensation: &lt;em&gt;Compens. = 100 %&lt;/em&gt; – airway pressure regulation to trachea level</td>
<td></td>
</tr>
<tr>
<td>Pmax</td>
<td>If <strong>Pmax/Paw high autoset</strong> is configured</td>
<td></td>
</tr>
</tbody>
</table>

<table>
<thead>
<tr>
<th>Volume Guarantee</th>
<th><strong>Ventilation parameters</strong></th>
<th><strong>Dependencies, information</strong></th>
</tr>
</thead>
<tbody>
<tr>
<td>On/Off</td>
<td>See chapter &quot;Configuration&quot; on page 155.</td>
<td></td>
</tr>
<tr>
<td>VT</td>
<td></td>
<td></td>
</tr>
<tr>
<td>Pmax</td>
<td>If <strong>Pmax/Paw high autoset</strong> is configured</td>
<td></td>
</tr>
<tr>
<td>Pinsp</td>
<td>If VG is switched off</td>
<td></td>
</tr>
</tbody>
</table>

<table>
<thead>
<tr>
<th>AutoRelease</th>
<th><strong>Ventilation parameters</strong></th>
<th><strong>Dependencies, information</strong></th>
</tr>
</thead>
<tbody>
<tr>
<td>On/Off</td>
<td></td>
<td></td>
</tr>
<tr>
<td>Exp. term.</td>
<td></td>
<td></td>
</tr>
<tr>
<td>Tlow</td>
<td>If AutoRelease is switched off</td>
<td></td>
</tr>
<tr>
<td>Tlow max</td>
<td>If AutoRelease is switched on</td>
<td></td>
</tr>
</tbody>
</table>

<table>
<thead>
<tr>
<th>HFO-Sigh</th>
<th><strong>Ventilation parameters</strong></th>
<th><strong>Dependencies, information</strong></th>
</tr>
</thead>
<tbody>
<tr>
<td>Psigh</td>
<td></td>
<td></td>
</tr>
<tr>
<td>RRaigh</td>
<td></td>
<td></td>
</tr>
<tr>
<td>Tisigh</td>
<td></td>
<td></td>
</tr>
<tr>
<td>Slopesigh</td>
<td>If <strong>Slope</strong> is configured, see page 177</td>
<td></td>
</tr>
</tbody>
</table>

<table>
<thead>
<tr>
<th>Volume Guarantee (HFO)</th>
<th><strong>Ventilation parameters</strong></th>
<th><strong>Dependencies, information</strong></th>
</tr>
</thead>
<tbody>
<tr>
<td>On/Off</td>
<td></td>
<td></td>
</tr>
<tr>
<td>VT</td>
<td></td>
<td></td>
</tr>
<tr>
<td>Ampl hf</td>
<td>Set the amplitude before switching off the Volume Guarantee (HFO)</td>
<td></td>
</tr>
<tr>
<td>Ampl hf max</td>
<td></td>
<td></td>
</tr>
</tbody>
</table>

### Additional information

For a detailed description of the additional settings, see chapter "Additional settings for ventilation" on page 304.
NIV – Non-invasive ventilation

Overview

Babylog VN500 can be used for the ventilation of intubated patients (application mode Tube) and for non-invasive ventilation (application mode NIV).

This chapter describes the use of non-invasive ventilation in the NIV application mode.

The following ventilation modes can be selected in the NIV application mode:

<table>
<thead>
<tr>
<th>Ventilation mode</th>
<th>Patient category</th>
</tr>
</thead>
<tbody>
<tr>
<td>PC-CMV</td>
<td>Ped. pat.</td>
</tr>
<tr>
<td>PC-AC</td>
<td>X</td>
</tr>
<tr>
<td>PC-SIMV</td>
<td>X</td>
</tr>
<tr>
<td>PC-PSV</td>
<td>X</td>
</tr>
<tr>
<td>PC-MMV</td>
<td>X</td>
</tr>
<tr>
<td>PC-APRV</td>
<td>X</td>
</tr>
<tr>
<td>SPN-CPAP</td>
<td>X</td>
</tr>
<tr>
<td>SPN-CPAP/PS</td>
<td>X</td>
</tr>
<tr>
<td>SPN-CPAP/VS</td>
<td>X</td>
</tr>
<tr>
<td>SPN-PPS</td>
<td>X</td>
</tr>
</tbody>
</table>

When the PC-HFO ventilation mode is set and the NIV application mode is activated, Babylog VN500 automatically switches the ventilation mode:
- in the Ped. pat. patient category to the SPN-CPAP/PS ventilation mode
- in the Neo. patient category to the SPN-CPAP ventilation mode

Safety information when using NIV

**CAUTION**
Application mode NIV must not be activated with intubated patients.

**CAUTION**
Use of masks increases the dead space. Observe the mask manufacturer’s instructions!

**NOTE**
Use suitable masks and prongs. Otherwise excessive leakages may occur.

**WARNING**
Avoid high airway pressures. Danger of aspiration!

**WARNING**
Alarm limits and ventilation settings must be checked or set again in order to ensure complete monitoring of ventilation after changing from NIV application mode to Tube application mode.

If automatic tube compensation (ATC) is activated in Tube application mode, it becomes ineffective when switching to NIV application mode.
Operation

Selecting NIV application mode

The application mode can only be selected in standby mode.

1. Touch the Start/Standby... button in the main menu bar.

   Babylog VN500 opens the Start/Standby dialog window. The Start/Standby page is displayed by default.

2. Touch the Standby button and confirm with the rotary knob.

   Babylog VN500 is in standby mode.

3. Touch the Tube/NIV tab (A).

4. Touch the NIV button (B) and confirm with the rotary knob.

   Babylog VN500 is in the NIV application mode.

   Babylog VN500 displays the \( \mathcal{H} \) symbol in the header bar.

   In the Neo. patient category, flow monitoring is deactivated.

Limiting the inspiratory flow in the Neo. patient category

In the Neo. patient category the inspiratory flow can be limited with the Flow max setting. The base flow and the nebulizer flow (if active) are not affected by this setting. If the NIV application mode is selected, the setting is reset to the maximum value.

To limit the maximum flow:

- Touch the Flow max button (C). Set the value by turning the rotary knob and push to confirm.
Starting NIV ventilation

Prerequisite: The Start/Standby dialog window is opened.

1. Touch the Start/Standby tab (A).

2. Touch the Start ventilation button (B) and confirm with the rotary knob.

Babylog VN500 starts the therapy with the set ventilation parameters. The main screen for ventilation is displayed.

Setting ventilation parameters for NIV

- Set the ventilation parameters as described under “Setting ventilation parameters” on page 85.

Therapy control Timax

The therapy control Timax (A) limits the maximum duration of supported breaths (Pressure Support, Volume Support, PPS) because the inspiratory termination criterion may be ineffective with very high leakages.

- Set the value for Timax by turning the rotary knob and push to confirm.

Therapy control TmanInsp and PmanInsp

Prerequisite: The Neo. patient category and the SPN-CPAP ventilation mode are set.

During manual inspiration, the duration of the mandatory breath is determined by the TmanInsp therapy control (B).

- Set and confirm the relevant values using the rotary knob.

During manual inspiration, the pressure of the mandatory breath is determined by the PmanInsp therapy control (C).

Monitoring during NIV

- Use additional monitoring, e.g., external SpO2, if necessary.

A delay time Tdisconnect between 0 and 60 seconds can be set for the lower alarm limit for the airway pressure.

Additional information

“Setting alarm limits” on page 128.
Displaying curves and measured values

Overview
This chapter describes how curves and measured values are displayed on the main screen as well as how to change the screen views during operation.

Changing the screen view
Babylog VN500 displays a preconfigured view on the main screen.
Three hospital-defined views can be created in the System setup dialog window.

Displaying other views
- Touch the Views... button in the main menu bar.
The screen displays the second view.
- Touch the Views... button.
The screen displays the third view.

Changing the display of monitoring fields
The fields can be standard or double in size. The information that can be displayed depends on the size of the fields:

<table>
<thead>
<tr>
<th>Parameter fields</th>
</tr>
</thead>
<tbody>
<tr>
<td><strong>Standard size</strong></td>
</tr>
<tr>
<td>Single parameter</td>
</tr>
<tr>
<td>Two parameters</td>
</tr>
<tr>
<td>Short trend for measured values</td>
</tr>
<tr>
<td>Short trend for set values</td>
</tr>
<tr>
<td><strong>Double size</strong></td>
</tr>
<tr>
<td>Single parameter</td>
</tr>
<tr>
<td>Parameter group</td>
</tr>
<tr>
<td>Loop</td>
</tr>
<tr>
<td>Short trend for measured values</td>
</tr>
<tr>
<td>Short trend for set values</td>
</tr>
</tbody>
</table>

<table>
<thead>
<tr>
<th>Curve fields</th>
</tr>
</thead>
<tbody>
<tr>
<td><strong>Standard size</strong></td>
</tr>
<tr>
<td>Single curve</td>
</tr>
<tr>
<td>Trend for measured values</td>
</tr>
<tr>
<td>Trend for set values</td>
</tr>
<tr>
<td>Tabular trend</td>
</tr>
<tr>
<td>Multi-trend</td>
</tr>
<tr>
<td>Alarm history</td>
</tr>
<tr>
<td><strong>Double size</strong></td>
</tr>
<tr>
<td>Single curve</td>
</tr>
<tr>
<td>Single loop</td>
</tr>
<tr>
<td>Double loops</td>
</tr>
<tr>
<td>Trend for measured values</td>
</tr>
<tr>
<td>Trend for set values</td>
</tr>
<tr>
<td>Tabular trend</td>
</tr>
<tr>
<td>Multi-trend</td>
</tr>
<tr>
<td>Alarm history</td>
</tr>
<tr>
<td>Lung display (Smart Pulmonary View)</td>
</tr>
</tbody>
</table>

The parameters can be displayed in parameter fields (A) and in the curve field (B).
Operation

Selecting the display of parameter fields
1 Touch the parameter field.

The selected parameter field is highlighted.
Babylog VN500 opens the dialog for the contents of the parameter field.

Selecting the display of curve fields
1 Touch the curve field.

The selected curve field is highlighted.
Babylog VN500 opens the dialog for the contents of the curve field.

2 Proceed as described under “Selecting the display of parameter fields”.

Additional information
“Configuring the screen view” on page 158.
“Factory-set screen views” on page 335.

Selecting the field size
2 Touch the 1x button (C) for standard size or 2x (D) for double size.

Selecting the display format
3 Touch the button (E).

The selection list for the display of parameters is displayed according to the selected size of the parameter field.

4 Select the display format and confirm with the rotary knob.

Selecting the parameter
5 Touch the button (F).

The selection list for the displayable parameters is displayed.

6 Select the parameter and confirm it with the rotary knob.

Closing the dialog
7 Touch the button X. The dialog is closed.
Evaluation loops

Displaying a reference loop
- Touch the Ref. button (A).

A loop is recorded and displayed as a reference loop. The date and time of the loop appear beside the button (A). The reference loop is drawn in black. The reference loop remains displayed until the Ref. button (A) is touched again.

Recording the current loop in order to freeze, display and save it afterwards
- Touch the Capture loop button (B).

The current loop is frozen. The loops are drawn in blue. After "freezing", a cursor (C) is displayed which can be moved with the rotary knob. The respective values are displayed (D).

Recording up to 10 loops of mandatory or spontaneous breaths
1 Touch the Draw button (E).
2 Set how many loops should be recorded with the rotary knob and push to confirm.

The set number is displayed in the button.

Additional information
A grid only appears if loops are displayed in the complete curve field.

Freezing waveforms

The Freeze waveforms function can be configured as a button in the main menu bar to enable direct access. See "Assigning functions to additional buttons" on page 161.
- Touch the Freeze waveforms button in the main menu bar.

The current curves are immediately frozen. The cursor (A) displays the time of "freezing" and the value at the cursor position.

To display a measured value at a certain moment in time:
- Position the cursor on the time with the rotary knob.

The measured value or the measured value pair are displayed above the curve.

To cancel the freezing waveforms function:
- Touch the Freeze waveforms button again in the main menu bar.
Smart Pulmonary View

Smart Pulmonary View is a graphic display of the compliance and resistance as well as of the spontaneous and mandatory minute volume.

A double-size curve field must be configured in order to display Smart Pulmonary View. See "Configuring the screen view" on page 158.

A The movement of the diaphragm indicates synchronized mandatory breaths, supported (triggered) breaths, or spontaneous breaths.

B The blue line around the trachea indicates the resistance \( R_{pat} \). The higher the resistance, the thicker the line. The value is also displayed.

C The blue line around the lungs indicates the compliance \( C_{dyn} \). The higher the compliance, the thinner the line. The value is also displayed.

D Diagram displaying the relationship between spontaneous breathing and mandatory ventilation. The following parameters are displayed in different colors:
   - \( V_{Tsp} \) and \( R_{Rsp} \)
   - \( V_{Tma} \) and \( R_{Rmand} \)

Smart Pulmonary View must be calibrated for each new patient. If the measured values for \( R_{pat} \) and \( C_{dyn} \) are outside the current display range, a red line appears and calibration is required.

Babylog VN500 displays the following information:

Touch "Take reference".

Calibrating Smart Pulmonary View:
- Touch the Take reference button (E).

The display range is adapted to the current measured values. The measured values from the last calibration are displayed as a broken line.

Additional information

For a detailed description, see "Smart Pulmonary View" on page 318.
Help

WARNING
Risk of operating error
The Help function is not a substitute for the instructions for use. The instructions for use must be observed to ensure safe operation.

The Help function can be configured as a button in the main menu bar to enable direct access. See "Assigning functions to additional buttons" on page 161.

Opening Help
● Touch the Help... button in the main menu bar.

The following buttons are available in the Help dialog window:
- Home (A) to open the start page
- ← (B) to scroll back
- → (C) to scroll forward
- Content (D) to open the table of contents
- Index (E) to open the index
● Touch the appropriate button.

Closing Help
● Touch the button (F).

Opening Help in the dialog window
The Help function can also be opened in the following dialog windows:
- Ventilation settings
- Special maneuvers > Nebulization
- Special maneuvers > Maneuvers

● Touch the button (G) in the dialog window.
The appropriate section of the Help is displayed.

Closing Help
● Touch the buttons (G) or (H) in the dialog window.
Operation

Maneuvers

Overview

Babylog VN500 permits the following maneuvers on the Special maneuvers > Maneuvers page:
- Manual inspiration – Manual inspiration/hold
- Oxygen enrichment for suction maneuver

Manual inspiration – Manual inspiration/hold

The Manual inspiration/hold maneuver can be activated in all ventilation modes and offers the following options:
- Between two mandatory breaths, a breath can be manually started and held. The pattern of the manually started breath corresponds to the ventilation pattern of the currently active automatic ventilation mode.
- Regardless of the start time, a mandatory breath can be prolonged.

1 Touch the Special maneuvers... button in the main menu bar.

Babylog VN500 opens the Special maneuvers dialog window.

2 Touch the Maneuvers tab (A) if the page is not already preset.

Triggering manual inspiration

- Briefly touch the Man. insp./hold button (B).

Manually extending inspiration

- Touch and hold the Man. insp./hold button (B) for the desired inspiratory time.

Babylog VN500 triggers an extended breath or extends an already triggered mandatory breath.

Babylog VN500 automatically ends inspiration:
- After a maximum of 40 seconds in the Ped. pat. patient category
- After a maximum of 5 seconds in the Neo. patient category

WARNING

The Manual inspiration/hold maneuver must not be used during endotracheal suction. Otherwise negative pressure may jeopardize the patient.

Additional information

The Manual inspiration/hold maneuver can be configured as a Man. insp./hold button in the main menu bar to enable direct access. See "Assigning functions to additional buttons" on page 161.
Oxygen enrichment for suction maneuver

Overview
To avoid hypoxia during endotracheal suction, Babylog VN500 offers a function for oxygen enrichment.

The O2 concentration is increased to the current inspiratory O2 concentration, multiplied by a factor. The factor can be configured, see page 177.

After oxygen enrichment is started, Babylog VN500 ventilates the patient with an increased O2 concentration for an initial oxygen enrichment phase of 180 seconds max. During this time, Babylog VN500 waits for a disconnection.

When the device is disconnected for suction, Babylog VN500 interrupts ventilation. During the suction phase, the acoustic alarms are suppressed so that the suction maneuver is not disturbed.

After suction and automatically recognized reconnection, Babylog VN500 delivers an increased O2 concentration for the final oxygen enrichment phase of 120 seconds.

During suction and for 120 seconds afterwards, the lower alarm limit for the minute volume is switched off.

Initial and final oxygen enrichment are only possible with a fully functioning flow sensor and if flow monitoring is switched on!

WARNING
Select an appropriate suction catheter for suction. Otherwise this may result in a too high negative pressure.

WARNING
Risk of patient injury during suction in a closed breathing circuit

Using closed suction systems produces negative pressure in the patient's airways. This leads to impaired ventilation and therefore to impaired gas exchange.

Observe patient condition.

Before suction
1 Touch the Special maneuvers... button in the main menu bar.
Babylog VN500 opens the Special maneuvers dialog window.
2 Touch the Maneuvers tab (A) if the page is not already preset.

3 Touch the O2 suction button (B) and confirm with the rotary knob.

The oxygen enrichment program is started.

Babylog VN500 ventilates in the set ventilation mode with an increased O2 concentration: 1 to 2 times the current FiO2 concentration.

If PEEP is not set to more than 4 mbar (4 cmH2O), a PEEP of 4 mbar (4 cmH2O) will be applied automatically. This PEEP allows Babylog VN500 to detect disconnection. The other ventilation parameters remain unaffected.
Screen display:

The field (C) in the header bar continuously displays the initial oxygen enrichment phase with the remaining time in seconds.

Initial oxygen enrichment lasts for a maximum of 180 seconds. During this time Babylog VN500 waits for a disconnection for suction. Babylog VN500 terminates the oxygen enrichment if there is no disconnection after the 180 seconds have elapsed.

After disconnection for suction, Babylog VN500 delivers a minimal flow for the duration of disconnection in order to detect automatically the end of the disconnection phase. 120 seconds are available for suctioning. In the header bar, the disconnection phase with the remaining time available for suction is displayed continuously in seconds (C).

**Automatic termination of oxygen enrichment**

If there is no reconnection when the time available (120 seconds) has elapsed, the oxygen enrichment is terminated. All alarms are immediately active again. Babylog VN500 continues to ventilate immediately in the set ventilation mode.

**After reconnection**

After reconnection, Babylog VN500 continues ventilating in the set ventilation mode, except that for 120 seconds an increased O₂ concentration will continue to be delivered for final oxygen enrichment.

In the header bar, the remaining time available for the final oxygen enrichment phase is displayed continuously in seconds.

**Terminating oxygen enrichment prematurely**

- Touch the **O₂ suction** button again and confirm with the rotary knob.

**Additional information**

The suction maneuver can be configured as a **O₂ suction** button in the main menu bar to enable direct access. See "Assigning functions to additional buttons” on page 161.
Medication nebulization

Safety information on medication nebulization

**WARNING**
Risk of fire

The flow sensor can ignite medications or other substances based on highly flammable substances.
- Do not nebulize medications or other substances that are easily flammable or spray them into the device.
- Do not use substances containing alcohol.
- Do not allow flammable or explosive substances to enter the breathing system or the breathing circuit.

**CAUTION**
During medication nebulization, do not use a heat and moisture exchanger (HME) at the Y-piece. The medication will not be appropriately administered to the patient.

**CAUTION**
Do not place a bacterial filter on the nebulizer outlet during nebulization! Bacterial filters may increase the flow resistance and impair ventilation.

**CAUTION**
Remove the medication nebulizer after use. Accidental medication nebulization may impair ventilation.

**CAUTION**
Surplus nebulized medication can affect the ambient air.

**CAUTION**
If no pneumatic medication nebulizer is connected, switch off the nebulization function. Otherwise, Babylog VN500 will deliver a tidal volume that is too low.

**CAUTION**
Ventilation impaired

If unapproved pneumatic medication nebulizers are used, the tidal volume and O2 concentration actually delivered may deviate from the displayed values. Only use the medication nebulizers listed in the current list of accessories.

**CAUTION**
Ventilation impaired

Aerosols may impair the functional integrity of the expiratory valve. When using medication nebulization, shorten the reprocessing cycles for the expiratory valve.

**Failure of the Air supply**

If the Air supply fails during medication nebulization, the medication nebulizer will continue to operate with 100 Vol% O2. In this case, deviations in the inspiratory O2 concentration are possible.
Air supply from the GS500 gas supply unit
If Babylog VN500 is supplied with Air from the GS500 gas supply unit and O2 is supplied from the central gas supply system, the medication nebulizer operates with O2 only.
The measured value $\text{FiO}_2$ indicates the O2 concentration of the gas supplied at the inspiratory port and not the O2 concentration reaching the patient. For deviations, see page 317.

Using a pneumatic medication nebulizer
When the $\text{Insp. flow}$ parameter has been configured instead of the $\text{Slope}$ parameter, 30 % of the set inspiratory time $T_i$ is automatically used during nebulization as the pressure rise time. Configure the $\text{Slope}$ parameter for a direct setting, see page 177. When a ventilation mode is active for which the inspiratory time $T_i$ cannot be set, 0.1 s is used as the slope.
The medication nebulizer nebulizes continuously. The aerosol generated during expiration does not reach the lungs, however.
If Babylog VN500 is supplied with Air and O2 from the central gas supply system, the medication nebulizer is operated with mixed gas at the set O2 concentration. Small deviations in the inspiratory O2 concentration of up to ±4 Vol% are possible. For respiratory rates above 12/min, refer to the graph on page 316.

In order to avoid false alarms and ensure monitoring:
- Use additional monitoring, e.g., external SpO2, if necessary.

Preparing the pneumatic medication nebulizer

**CAUTION**
In order to ensure limitation of the nebulizer flow, only use pneumatic medication nebulizer 8411030.
- Prepare the medication nebulizer in accordance with the corresponding instructions for use.

Installing the medication nebulizer into the breathing circuit

1. Remove the corrugated hose for the breathing circuit (A) from the inspiratory port of the Y-piece and connect it to the inlet port of the medication nebulizer.
2. Fit the corrugated hose (B), length 0.13 m (5.1 in), to the outlet port of the medication nebulizer.
3. Connect the free end of the corrugated hose (B) to the inspiratory port of the Y-piece.

Additional information
Depending on the Dräger breathing circuit used, an adapter is required for the medication nebulizer connection.
Operation

When using on the incubator

- Push the inlet port or the outlet port of the medication nebulizer into the upper hose guide of the incubator.

When using without incubator

1. Press the inlet port or the outlet port of the medication nebulizer into one side of the clip and the expiratory hose into the other.
2. Place the medication nebulizer in the vertical position.

Connecting the nebulizer hose

**WARNING**
The nebulizer port (F) must be used for nebulization only! Otherwise the proper functioning of the device may be disrupted and the patient endangered.

- Connect the nebulizer hose (G) to the nebulizer port (F).

**CAUTION**
Check the correct functioning of the medication nebulizer. Check whether aerosol is generated. A medication nebulizer fault is not detected by Babylog VN500.
**Operation**

**Switching on medication nebulization**

1. Touch the *Special maneuvers*... button in the main menu bar.
2. Touch the *Nebulization* tab (A).
3. Touch the button for the desired nebulization time (B).

Nebulization can be set to 5, 10, 15, or 30 minutes or to continuous nebulization.

**Deactivating flow monitoring**

In the instruction field (C) Babylog VN500 requests that the user switches off flow monitoring.

- Touch the *Off* button (D) and confirm with the rotary knob.

**Removing the neonatal flow sensor from the breathing circuit**

In the instruction field (E) Babylog VN500 requests the user to remove the neonatal flow sensor from the breathing circuit.

**WARNING**

Risk of fire

The measuring wires of the neonatal flow sensor are very hot and may ignite deposits of medication aerosols during nebulization.

- Before medication nebulization, remove the complete ISO 15 neonatal flow sensor, or remove the sensor insert from the neonatal flow sensor Y-piece and insert a sealing plug.
- Use additional monitoring since otherwise the minute volume is not monitored and apnea monitoring is limited.
When using the neonatal flow sensor Y-piece (8410185):

1. Disconnect plug (F) of the flow sensor cable from the neonatal flow sensor (G).

2. Remove the insert (H).

3. Insert the sealing plug (I) (8411024). The sealing plug is a component of the medication nebulizer.

When using the neonatal flow sensor ISO 15 (8411130):

1. Remove the flow sensor (J) from the tube and the Y-piece.

2. Connect the tube (K) to the Y-piece.

- Replace or clean the neonatal flow sensor if there is visible soiling. See "Dismantling the expiratory valve" on page 230.

- Touch the Done button (L).

**During medication nebulization**

Babylog VN500 starts nebulization. The symbol and the remaining nebulization time is displayed in the screen header bar.

Babylog VN500 automatically switches off the medication nebulizer after the set nebulization time has elapsed.

A message indicating that nebulization has been ended appears in the screen header bar.

**During continuous medication nebulization**

The Continuous nebulization in progress message is displayed in the screen header bar.

If the parameter field for continuous nebulization Cont. neb. has been configured for display, the duration of medication nebulization is displayed.
Abort medication nebulization

Touch the Cancel button (A).

Required steps after medication nebulization

1. Remove any residual medication. Observe the instructions for use of the medication nebulizer.

2. If a bacterial filter is used to protect the expiratory valve, exchange or remove the bacterial filter.

3. Reconnect the neonatal flow sensor.
   - When using the neonatal flow sensor Y-piece (8410185):
     - Remove the sealing plug and push the insert back in.
     - Reconnect plug of the flow sensor cable.
   - When using the neonatal flow sensor ISO 15 (8411130):
     - Re-insert the neonatal flow sensor in the Y-piece.

4. Activate flow monitoring with neonatal flow sensor, see page 145.

5. If the insert of the neonatal flow sensor has been replaced, calibrate the neonatal flow sensor, see page 143.

Additional information

The Nebulization maneuver can be configured as a Nebulization button in the main menu bar to enable direct access. See "Assigning functions to additional buttons" on page 161.

Nebulization may lead to increased deposits. Consequently, it may be necessary to change the following components more often:
- Flow sensor
- Expiratory valve

Special maneuvers
Fitting the Aeroneb nebulizer

- Observe the instructions for use of the Aeroneb nebulizer.
- Observe the "Safety information for the use of HMEs, bacterial filters, and breathing circuits" on page 52.
- Observe the "Safety information on medication nebulization" on page 103.
- Do not switch on the Nebulization maneuver on Babylog VN500 as the Aeroneb nebulizer does not require a nebulizer flow from Babylog VN500.

Before nebulization with Aeroneb
1. Deactivate flow monitoring with neonatal flow sensor, see page 145.
2. Remove the neonatal flow sensor from the breathing circuit, see page 106.

WARNING
Risk of fire
The measuring wires of the neonatal flow sensor are very hot and may ignite deposits of medication aerosols during nebulization.
- Before medication nebulization, remove the complete ISO 15 neonatal flow sensor, or remove the sensor insert from the neonatal flow sensor Y-piece and insert a sealing plug.
- Use additional monitoring since otherwise the minute volume is not monitored and apnea monitoring is limited.

After nebulization with Aeroneb
1. If a bacterial filter is used to protect the expiratory valve, exchange or remove the bacterial filter.
2. Reconnect the neonatal flow sensor.

When using the neonatal flow sensor Y-piece (8410185):
- Remove the sealing plug and push the insert back in.
- Reconnect plug of the flow sensor cable.

When using the neonatal flow sensor ISO 15 (8411130):
- Re-insert the neonatal flow sensor in the Y-piece.
3. Activate flow monitoring with neonatal flow sensor, see page 145.
4. Calibrate the neonatal flow sensor, see page 143.

Additional information
- For the order number of the Aeroneb nebulizer, see the list of accessories.
GS500 gas supply unit

In order to ensure continuous Air supply, Babylog VN500 can be equipped with the GS500 gas supply unit. If Babylog VN500 is connected to the central gas supply system, GS500 ensures the supply of Air to the device in the case of failure of the central gas supply system and during intrahospital patient transport.

When Babylog VN500 is in the **PC-HFO** ventilation mode and is supplied from the gas supply unit GS500, it must be connected to an O2 compressed gas cylinder. The ejector is then operated using O2 instead of Air. This leads to a significantly increased O2 consumption which must be observed particularly during the transportation of patients within the hospital.

**Installing the bacterial filter**

- Fit the bacterial filter (A) onto the inspiratory port.

**Using the gas supply unit**

Prerequisite: Functionality of the gas supply unit is activated, see "Configuring supply units" on page 186.

If Babylog VN500 is not connected to the central gas supply system, GS500 starts the supply of Air automatically.

In the event of failure of the central Air supply, or if the probe of the Air compressed gas hose becomes detached from the wall terminal unit of the central gas supply system, Babylog VN500 displays an alarm message. The gas supply unit starts the supply of Air using GS500 after 4 seconds at the latest.

**Switching on the gas supply unit for intrahospital patient transport**

1. Touch the **Special maneuvers...** button in the main menu bar.
2. Touch the **Transport** tab (A).
3. Touch the **On** button (B).
4. Pull out the probe of the Air compressed gas hose from the wall terminal unit of the central gas supply system.
Operation

If the probe of the Air compressed gas hose has not been pulled out within 5 minutes of the gas supply unit being switched on, Babylog VN500 switches off the gas supply unit.

5 Pull the probe of the O₂ compressed gas hose out from the wall terminal unit of the central gas supply system and provide a replacement O₂ supply if necessary.

Switching off the gas supply unit

- Touch the Off button (C).

Additional information

- Deactivating functionality of the gas supply unit, see “Configuring supply units” on page 186.

O₂ therapy

Safety information for O₂ therapy

During O₂ therapy, only the O₂ concentration and the inspiratory pressure are monitored.

Oxygen masks, hoods, or nasal cannulas can be used for O₂ therapy.

**CAUTION**

Only use oxygen masks for the O₂ therapy. Do not use masks for non-invasive ventilation (NIV). Use of unsuitable masks may jeopardize the patient.

**CAUTION**

Internal monitoring is deactivated. Airway pressure and ventilation parameters, e.g., flow, minute volume or apnea are not monitored. Use external SpO₂ monitoring for patients who are dependent on an increased defined O₂ concentration. Otherwise a worsening of the patient's condition cannot be detected.

**NOTE**

If the pressure needed for the set flow exceeds 30 mbar (30 cmH₂O), the device issues an alarm and the safety valve is opened. The cause may be a kinked breathing hose or a blocked mask or nasal cannula.

Observe the specified flows and sizes of the particular accessories.

Preparing O₂ therapy

**WARNING**

Do not use antistatic or conductive breathing hoses. The use of these materials increases the risk of electric shock to the patient and of fire in an oxygen-enriched environment.
Preparing a system with a Fisher & Paykel MR 850 breathing gas humidifier

1. Hang the hinged arm (A) on the rail and tighten the screws. Depending on the desired position of the device in relation to the bed, the hinged arm can be fitted to either side of the device.

2. Fit the breathing hoses (B) for inspiration. The expiratory ports on the device and on the Y-piece remain open!

3. Switch on Babylog VN500. See page 64.

4. Switch Babylog VN500 to standby. See page 114.

5. Activate O2 monitoring. See page 146.

The alarm limits for MVe, RR, Paw, Tapn are not active. The alarm limits for O2 monitoring are automatically set by the device.

Switching on O2 therapy

O2 therapy can only be switched on in standby mode.

1. Touch the Start/Standby... button in the main menu bar.

Babylog VN500 opens the Start/Standby dialog window. The Start/Standby page (A) appears by default.

2. Touch the Standby button (B) and confirm with the rotary knob.

3. Touch the O2 Therapy button (C).

The message field (D) displays the information to use specific masks for O2 therapy.

4. Connect a mask, hood, or nasal cannula for O2 therapy.

5. Touch the Start ventilation button (B) and confirm with the rotary knob.

O2 therapy is switched on. Babylog VN500 displays the main screen with the therapy bar (E) for O2 therapy. The message O2 Therapy is displayed in the header bar (F).

During O2 therapy, the screen display on the main screen cannot be customized.
Operation

Setting FiO2 and flow for O2 therapy

1  Touch the corresponding therapy control in the therapy bar:
   – FiO2 (A)
   – Flow (B)

2  Set the value by turning the rotary knob and push to confirm.

The FiO2 concentration is represented graphically (C).

Switching off O2 therapy

1  Touch the Start/Standby... button.

Babylog VN500 opens the Start/Standby dialog window. The Start/Standby page is displayed by default.

2  Touch the Standby button and confirm with the rotary knob.

Babylog VN500 is in standby mode. O2 therapy is switched off. The therapy type can be switched to ventilation.

Setting O2 and flow in the dialog window

The O2 and flow can also be set in the Ventilation settings dialog window.

– Touch the Ventilation settings... button.

Or

– Touch the ↑ button (D).
Standby mode

Switch to standby mode for the following actions:
- Keep Babylog VN500 ready for operation while the patient is absent
- Change the therapy type between ventilation and O2 therapy
- Change the patient category
- Change the application mode
- Perform the device and breathing circuit check
- Query the status of accessories
- Switch off Babylog VN500

**WARNING**
Ventilation does not take place in standby mode! The device must only be set to standby mode when no patient is connected to the device. The patient may otherwise be jeopardized.

Activating standby mode

1. Touch the **Start/Standby...** button in the main menu bar.

   Babylog VN500 opens the **Start/Standby** dialog window. The **Start/Standby** page (A) appears by default.

2. Touch the **Standby** button (B) and confirm with the rotary knob.

   The message **Standby mode activated** is displayed in the header bar.

3. Touch the **ALARM RESET** button in the header bar and confirm with the rotary knob.

   Babylog VN500 is in standby mode. **Standby** is displayed in the screen header bar.
Continuing the therapy

1  Check the ventilation settings (A) of the current patient.

Change the ventilation settings if necessary. Touch the Ventilation settings button (B).
Babylog VN500 opens the corresponding page.

2  Touch the Start ventilation button (C) and confirm with the rotary knob.

The main screen is displayed; Babylog VN500 continues ventilating.

Additional information

If the patient category or the body weight is changed, Babylog VN500 determines new start-up values for ventilation. See "Admitting a new patient" on page 65.

For information on changing ventilation settings, see "Setting ventilation" on page 84.
Operation

Ending operation

1 Switch Babylog VN500 to standby mode: Touch the Start/Standby... button in the main menu bar. Touch the Standby button and confirm with the rotary knob.

2 Press the key (A) on Infinity C500. Babylog VN500 opens the Shut down device dialog.

3 Touch the OK button (B) and confirm with the rotary knob. Babylog VN500 ends operation.

As soon as the screen is completely dark
- Disconnect the mains plug from the mains power socket.
- Pull the probe of the Air compressed gas hose and the probe of the O2 compressed gas hose out from the wall terminal units of the central gas supply system.

CAUTION
Disconnect the compressed gas hoses from the central gas supply system. Otherwise minute internal leaks could contaminate the central gas supply system through the reverse flow of supply gases.

If Babylog VN500 cannot be switched off on account of a device malfunction

1 Open the device flap on the left side of Babylog VN500.
2 Set the toggle switch to (off).

Once the toggle switch has been pressed and the mains plug is disconnected, Babylog VN500 cannot be switched on.

Placing back into operation

1 Insert the mains plug into the mains power socket.
2 Open the device flap on the left side of Babylog VN500.
3 Set the toggle switch to (on).
4 Switch on Babylog VN500: Press the key on Infinity C500.

When Babylog VN500 is not in standby mode and the button (A) is pressed, the Start/Standby page is opened.
Disconnecting the device from the mains voltage

In the event of device malfunctions or other hazards, the device must be completely disconnected from the mains voltage.

When the toggle switch is at (off), only parts of the device are disconnected from the mains voltage. The batteries continue to be charged. To completely disconnect the device from the mains voltage, unplug the mains plug.

Storing Babylog VN500

Switch Babylog VN500 to energy-saving mode if storing for longer periods.

1 End operation. See "Ending operation" on page 116.

2 Set the toggle switch on the left side of Babylog VN500 to (off) immediately after switching off the device.

3 Disconnect the mains plug from the mains power socket.
Mains power supply / DC power supply

Components and terms

Mains power supply
The device is supplied with mains power via the power cable. Information on voltage ranges and mains power characteristic values can be found in chapter Technical Data, Operating data.

Internal battery
The internal battery is supplied with the device.

PS500 power supply unit
In addition to the internal battery, the device can optionally be equipped with the PS500 power supply unit.

Use of power supplies
The device is supplied with electric power from the following sources in the order stated:
– Mains power
– Batteries in the PS500 (if present)
– Internal battery

The switch-over between these sources takes place without interruption to operation according to the following rules:
– If the mains voltage is sufficient, the power is supplied from the mains.
– If the mains voltage is not sufficient or during a battery check, the power is supplied from the batteries.

Display of power supplies
The power supply is displayed on the Infinity C500 operating and display unit.
On Infinity C500 (MS18746):

On Infinity C500 (MK31500):

A LED for mains power:
– Lights green when mains power is applied and the toggle switch is in the position.
– If the LED does not light up, the device is disconnected from the mains power.

B LED for the internal battery:
– Lights green when the battery charge is greater than approx. 90 %.
– Lights yellow when the battery charge is between approx. 10 % and 90 %.
– Does not light if the internal battery is faulty, discharged or device is switched off with the toggle switch (energy-saving mode).
Battery operation

Alarm messages during battery operation
Switch-over to the batteries is indicated with the alarm message **Battery activated**. The alarm priority can be configured, see "Setting the priority of the battery alarms" on page 167.

Alarm messages are displayed corresponding to the remaining battery capacity in order to warn against the complete discharge of the battery. Alarm messages, see chapter "Alarm – Cause – Remedy."

- Reestablish the mains power supply immediately to avoid interruption of the ventilation functions.

When battery supply is no longer needed, recharge the batteries, see chapter "Battery charging".

Operating time during battery operation
The operating time depends on the following battery factors:
- Age
- Utilization (frequency, duration, and power consumption)
- Battery charge
- Ambient temperature

For operating times when batteries are fully charged and new, and ventilation is typical, see chapters "Battery ageing" on page 324 and "Batteries" on page 276.

Observe the maintenance intervals.

**NOTE**
The LED □□□ indicates only the battery charge of the internal battery, even when the PS500 is present.

Battery charging

The batteries are charged when the device is supplied with mains voltage. The ■ symbol (A) is displayed in the screen header bar.

The batteries are charged in the following order:
- Internal battery
- Batteries in the PS500 (if present)

Charging times
For information on the charging times, see page 277.

Battery charge indication on the screen
The battery charge is indicated by the symbol (B) in the header bar on the screen. The battery charge indication applies to both charging and discharging. When the batteries are being charged, the last segment in the battery symbol flashes white.

<table>
<thead>
<tr>
<th>Symbol</th>
<th>Battery charge</th>
</tr>
</thead>
<tbody>
<tr>
<td>■□□□□</td>
<td>90 to 100 %</td>
</tr>
<tr>
<td>□□□□□</td>
<td>60 to &lt;90 %</td>
</tr>
<tr>
<td>□□□□</td>
<td>40 to &lt;60 %</td>
</tr>
<tr>
<td>□□□□</td>
<td>20 to &lt;40 %</td>
</tr>
<tr>
<td>□□□□</td>
<td>&lt;20 %, flashes light and dark red in 1-second pulses</td>
</tr>
<tr>
<td>□□□□</td>
<td>Batteries faulty or no information available on the battery charge</td>
</tr>
</tbody>
</table>

The battery charge indication applies to both charging and discharging.

The battery charge indication always shows the total battery charge that is available. If there is a PS500 present, the battery charge available from the internal battery and the PS500 will be displayed.
A flashing \(\text{⚠️}\) symbol (C) indicates the following:
- The battery check is running.
- The interval for the battery check has expired.
- The last battery check failed.
- Battery replacement is recommended.

**Parameter field Battery**

In addition to the battery charge indication, the Battery parameter field can be configured.

The Battery parameter field contains the following information:
- **PS500** (if present)
  - Operating time in minutes (value corresponds to the operating time when the battery is used at the present power consumption) (A)
  - Battery charge in percent (B)
- **Internal** (internal battery)
  - Operating time in minutes (value corresponds to the operating time when the battery is used at the present power consumption) (C)
  - Battery charge in percent (D)

**NOTE**

Ageing and use of the batteries can result in a shorter operating time compared with new batteries.

Depending on the battery used, the battery charge is indicated to the nearest 5 or 10 minutes.

It is always the minimum calculated operating time that is displayed.

**Care and maintenance of the batteries**

Take note of the following points to limit premature ageing of the battery:
- Operate the device under the stated ambient conditions
- Avoid storing the device with discharged or partially discharged batteries
- Connect the device to the mains power supply after battery operation
- Avoid shocks and vibrations
- Perform the recommended battery checks
Intrahospital patient transport

**WARNING**
Do not tilt the device by more than 10°. Failure to observe this may result in the device toppling over. Danger of damage to device or personal injury!

**WARNING**
The device must not be placed on the bed while transferring a patient within the hospital. The device could topple over or fall down. Danger of damage to device or personal injury!

**WARNING**
Do not lean, press, push or pull against the trolley above the marking points on the trolley. The trolley could topple over.

**WARNING**
Do not move trolley faster than at a walking pace. There is an increased danger of the trolley toppling over at thresholds, uneven surfaces and ramps. Reduce the speed of transport further. Danger of damage to equipment!

**WARNING**
Two people are always required to move the device. Otherwise there is an increased risk of the device toppling over.

**WARNING**
Make sure to securely hold onto the handle of the trolley whenever moving or positioning the device. Otherwise there is an increased risk of the device toppling over.

**WARNING**
Patient hazard due to discharged batteries. Only start transporting patients when the batteries are sufficiently charged.

When transporting a patient within the hospital, the user must ensure that the patient is monitored at all times.

When transporting the patient within the hospital, grasp the trolley handle firmly and push the device in longitudinal direction.

### Using Babylog VN500 with a safety bar

**CAUTION**
During intrahospital patient transport, Babylog VN500 must be used with a safety bar (A) in order to prevent accidental disconnection of the breathing hoses or damage to the inspiratory port and the expiratory port.

*Image of safety bar with label A.*

### Increasing the toppling stability during intrahospital patient transport

To ensure that the equipment cannot topple over, the accessories must be moved to the most advantageous position:

1. Hinged arm set to minimum deflection.
2. Hoses and cables hooked as close as possible to the trolley.
3. Humidifier secured to the trolley, not to the lateral rails of Babylog VN500.
Operation

Additional information

Air supply from the GS500 gas supply unit, see "GS500 gas supply unit" on page 110.

Power supply, see "Mains power supply / DC power supply" on page 118.

For the order number of the safety bar, see the list of accessories.
# Alarms

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<th>Page</th>
</tr>
</thead>
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<td>131</td>
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<td>132</td>
</tr>
</tbody>
</table>
**Alarms**

**Overview**

Alarms are issued acoustically and visually. The alarm tone can be suppressed for 2 minutes. The *Alarms* dialog window provides the following functions for selection:

- Setting alarm limits
- Displaying current alarms
- Alarm history
- Alarm settings

**Display of alarms**

In the event of an alarm, the system displays the relevant alarm message in the alarm message field (A). If the parameter field (B) is configured to display an individual parameter, the parameter field (B) of the parameter triggering the alarm flashes. The alarm bar (D) flashes in the color of the corresponding alarm priority.

If the alarm message field (A) contains more alarms than can be displayed, the *More...* button (C) appears in the header bar. Touching this button opens the page containing all the active alarms.
Alarms

Alarm priorities

A certain priority, indicating the urgency, is assigned to each alarm.

The following table shows the differences between the alarm priorities with respect to identification and the action required.

<table>
<thead>
<tr>
<th>Alarm priority</th>
<th>Identification</th>
<th>Action required</th>
</tr>
</thead>
<tbody>
<tr>
<td>High</td>
<td>Red</td>
<td>!!! Immediate action is necessary in order to avert an acute danger</td>
</tr>
<tr>
<td>Medium</td>
<td>Yellow</td>
<td>!! Prompt action is necessary in order to avert a danger</td>
</tr>
<tr>
<td>Low</td>
<td>Turquoise</td>
<td>! Attention is necessary, but a delayed response is sufficient</td>
</tr>
</tbody>
</table>

Optical alarm signals

The following optical alarm signals are displayed in the event of an alarm.

<table>
<thead>
<tr>
<th>Alarm priority</th>
<th>Alarm message field</th>
<th>Parameter field</th>
<th>Alarm bar</th>
</tr>
</thead>
<tbody>
<tr>
<td>High</td>
<td>Alarm message on a red background</td>
<td>Flashes red</td>
<td>Flashes red</td>
</tr>
<tr>
<td>Medium</td>
<td>Alarm message on a yellow background</td>
<td>Flashes yellow</td>
<td>Flashes yellow</td>
</tr>
<tr>
<td>Low</td>
<td>Alarm message on a turquoise background</td>
<td>--</td>
<td>--</td>
</tr>
</tbody>
</table>

If several alarms occur at the same time, the two most urgent alarms are displayed first.

High-priority alarm messages that are no longer active are displayed in the background color of the alarm message field.

In the tables for Current alarms and Alarm history the priority of the alarm messages is also indicated by exclamation marks.

Acoustic alarm signals

Babylog VN500 generates different alarm tone sequences to indicate alarms acoustically. The alarm tone sequences can be configured, see “Selecting alarm tone sequences” on page 167.
## Displaying information on alarms

### Displaying current alarms

To display the current alarms, proceed as follows:

1. Touch the alarm message in the header bar.
   Or
2. Touch the **Alarms...** button in the main menu bar.

3. Touch the **Current alarms** tab (A).

All the current alarm messages are displayed chronologically with the corresponding duration, priority and alarm message text in the list (B).

### Displaying the cause and remedy for an alarm

1. Touch the alarm message or select it in the list (B) with the rotary knob.
2. Touch the button (C).
   This displays the cause and remedy for the alarm message selected.
3. Eliminate the fault.

### Acknowledging an alarm message that is no longer active

After the fault has been eliminated, the alarm tone is silenced. Medium- and low-priority alarm messages expire automatically. High-priority alarm messages continue to be displayed for information after the cause of the alarm has been eliminated and need to be acknowledged.

1. Touch the **ALARM RESET** button (A) in the header bar.
2. Confirm with the rotary knob.

### Acknowledging all alarm messages that are no longer active

Prerequisite: The **Current alarms** page (A) is opened.

1. Touch the **Reset all** button (B).
2. Confirm with the rotary knob.

The acknowledgeable messages are deleted in the header bar and in the list containing the current alarms. However, Babylog VN500 records all alarm messages in the alarm history.

### Additional information

For a list of causes and remedies, see chapter "Alarm – Cause – Remedy" on page 189.
Alarm history

The alarm history records all alarm messages in chronological order.

The entries in the alarm history are also retained after the device has been switched off and on again or following a power supply failure.

The alarm history is part of the logbook. The length of the alarm history depends on the number of logbook entries.

When the logbook reaches its maximum size, the oldest entry in the logbook is deleted as each new entry is logged.

Switching the device off and on are not recorded in the logbook.

1 Touch the Alarms... button in the main menu bar.

2 Touch the Alarm history tab (A).

3 Use the buttons (B) to scroll in the alarm history.
Setting alarm limits

- Touch the Alarms... button in the main menu bar.

The Limits page (A) appears by default.

The alarm limit settings and the current measured value are displayed.

(B) $\mathcal{R}$ : Upper alarm limit

(C) Current value : Current measured value

(D) $\mathcal{S}$ : Lower alarm limit

Additional information

The start-up values for the alarm limits can be configured specifically as required by the hospital concerned, see page 165.

The alarm limits are displayed depending on the ventilation parameter in the parameter field.

Deactivating alarm limits

**WARNING**

Alarms must only be deactivated if the safety of the patient is not jeopardized by the absence of an alarm!

The following alarm limits can be deactivated:

<table>
<thead>
<tr>
<th>Patient category</th>
<th>Invasive ventilation</th>
<th>Non-invasive ventilation</th>
</tr>
</thead>
<tbody>
<tr>
<td>Ped. pat.</td>
<td>RR high</td>
<td>MV low</td>
</tr>
<tr>
<td></td>
<td>–</td>
<td>RR high</td>
</tr>
<tr>
<td></td>
<td>–</td>
<td>Tapn</td>
</tr>
<tr>
<td>Neo.</td>
<td>MV low</td>
<td>–</td>
</tr>
<tr>
<td></td>
<td>RR high</td>
<td>RR high</td>
</tr>
<tr>
<td></td>
<td>Tapn</td>
<td>Tapn</td>
</tr>
</tbody>
</table>

**WARNING**

The alarm limits must be set to meet the needs of the therapy required by the current patient. The patient may otherwise be jeopardized. Setting extreme alarm limits can render the alarm system useless.

**How to set an alarm limit**

Prerequisite: The Limits page (A) is opened.

1. Touch the corresponding button for the alarm limit.

2. Set the value by turning the rotary knob and push to confirm.

**How to deactivate an alarm limit**

1. Touch the corresponding button for the alarm limit.

2. Continue turning the rotary knob until Off is displayed instead of the value.

3. Confirm with the rotary knob.

The alarm limit is deactivated. Babylog VN500 displays the $\mathcal{X}$ symbol in the header bar and the deactivated alarm limit. The header bar can display up to 5 deactivated alarm limits.
Alarms

Response to power failure

Alarm limits are also retained in the event of a power failure, e.g., caused by a defective internal battery.

Display of alarm limits in the parameter field

If the alarm limits are assigned to a ventilation parameter, the alarm limits are displayed in the parameter fields for single parameters (standard and double size).

The following assignments have been defined:

<table>
<thead>
<tr>
<th>Alarm limits</th>
<th>Measured values</th>
</tr>
</thead>
<tbody>
<tr>
<td>MV high, MV low</td>
<td>MVe</td>
</tr>
<tr>
<td>Paw high</td>
<td>PIP</td>
</tr>
<tr>
<td>RR high</td>
<td>RR</td>
</tr>
</tbody>
</table>
Setting the volume of the alarm tone

**WARNING**
Unnoticed alarms in loud environments
Alarm situations are not recognized.
Set the volume of the alarm tone so that alarms can be heard.

1. Touch the **Alarms**... button in the main menu bar.
2. Touch the **Settings** tab (A).
3. Touch the button (B).
4. Set the volume of the alarm tone by turning the rotary knob and push to confirm.

During the automatic switch-over between day and night modes, the alarm tone volume setting is overwritten by the volumes defined for these times. An automatic increase in volume can be activated. See "Setting the alarm tone" on page 167.

The lower value for the volume of the alarm tone is limited by the configured minimum volume of the alarm tone. The minimum volume can be configured on the **System setup > Alarms > Alarm vol./tone** page, see "Setting the alarm tone" on page 167.

**Additional information**

The **Settings** page can be configured for direct access into the main menu bar as the **Alarm volume** button. See "Assigning functions to additional buttons" on page 161.
Suppressing the alarm tone

The alarm tone can be suppressed for a maximum of 2 minutes.

- Press the (Audio paused) key (A).
  
  This suppresses the acoustic alarm for 2 minutes. Babylog VN500 displays the symbol in the header bar and the remaining time for the suppressed alarm tone.
  
  If an alarm with a higher priority appears during this time, the alarm tone sounds once.
  
  If the fault triggering the alarm is not eliminated after 2 minutes, the alarm tone sounds again.
  
  Reactivate the alarm tone before the suppression time has elapsed:
  - Press the (Audio paused) key (A) again.

Position of the user to the alarm system

The optical alarm signals are designed as follows:

- At a distance of 4 m (157 in) it is possible to recognize which device is generating an alarm.
- At a distance of 1 m (39 in) the alarm message can be read clearly.

The alarm volume can be set so that the acoustic alarm signals can be heard in the vicinity of the device, see “Setting the alarm tone” on page 167.
Alarms

Failure of the acoustic alarm

If the loudspeaker for acoustic alarm signaling (main alarm) fails on account of a defect, an intermittent tone will be generated by the loudspeaker for the auxiliary alarm.

This intermittent tone is also used for the power failure alarm.

Additional information on the power failure alarm

See “Failure of the power supply” on page 59.
Trends and data

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Trends and data

Overview

Babylog VN500 saves measured value and trend data. Trends are displayed in the form of a graphic or a table. The following can be displayed: current measured values, settings and hospital-specific combinations of measured and set values. The logbook can save up to a maximum of 5000 entries. Data can be exported with a USB storage medium.

The Trends/Data dialog window provides the following functions for selection:

- Display trends
- Display data
- Logbook
- Data export

Displaying trends

Trends are displayed as a graphic or a table. Trends are recorded for up to 7 days.

In graphic trends, measured values are displayed in blue and set values in green. In the apnea trend, the number of the apneic events that occurred per minute is represented as a histogram.

In tabular trends, measured values are displayed in blue lettering and set values in green lettering.

Displaying an additional graphic trend

Prerequisite: The Trends page (A) is opened.

- Touch the Graphics 2 tab (C).

Each page contains 2 graphic trend displays (D).

Graphic trends

1. Touch the Trends/Data... button in the main menu bar.

Babylog VN500 opens the Trends page (A) with the Graphics 1 page (B).
Selecting parameters for the graphic trend display

Prerequisite: The Graphics 1 or Graphics 2 page is open.

1 Touch the button (E).

The Setup dialog is displayed with the buttons for Meas. (F) and Settings (G).

The measurements (F) are divided into the following parameter types:
- Pressures
- Minute vol.
- Volume/Flow
- Gases
- Timing/Cycl.
- Others
- Events

The settings (G) are divided into the following parameter types:
- Pressures
- Volume/Flow
- Gases
- Timing/Cycl.
- Others

2 Touch the appropriate button for measurements or settings.

Another dialog containing all the parameters of the selected parameter type is displayed.

3 Touch the desired parameter.

4 Confirm with the OK button (H).

The dialog for the group selected is closed.

A maximum of 3 parameters can be selected for each graphic trend display. If 3 parameters are already selected, one parameter must be deselected before selecting a new parameter.

5 Select further parameters according to step 2 to 4.

6 Confirm the parameter selection with the OK button (H).

The selected parameters are displayed in the trend display. The Setup dialog is closed.

The selection can be canceled with Cancel (I). The previous selection is displayed in the graphic trend.

Clear (J) can be used to delete all parameter selections made.

Deselecting a parameter in the trend display

Touch the parameter to be deselected in the parameter type dialog. The button turns pale green.
Selecting a time interval for the graphic trend display

Prerequisite: The Graphics 1 or Graphics 2 page is open.

1. Touch the button for the time interval (K).
2. Select the time interval from the selection list (2, 4, 8, 12 hours; 1 day, 7 days).

Displaying the value of a parameter at a certain moment in time

- Position the cursor (L) on the time by turning the rotary knob or touching the time.

The parameter value and the marked time are displayed (M).

The marked time in the trend display also corresponds with the marked row of this time in the logbook.

Changing the displayed time period

- Touch the buttons in the scrollbar (N) or turn the rotary knob.

Apnea trend, apnea ventilation trend

In the apnea trend, the number of the apneic events that occurred per minute is represented as a histogram. The number per minute is represented as a bar height. If an apnea lasts longer than one minute, the apnea is only counted once in the period of occurrence.

In the apnea ventilation trend, the system displays whether or not apnea ventilation is activated.

Prerequisite: The Graphics 1 or Graphics 2 page is open.

1. Touch the button.
2. In the Setup dialog window under Meas., touch the Events parameter type.
3. Select the Apnea or Apnea Vent. event.

Additional information

The apnea trend is only recorded when apnea ventilation is switched off.

The duration of an apnea is displayed only in the alarm history.
**Tabular trend**

Babylog VN500 displays the trends of all parameters in a table. The parameters that are first displayed correspond with the parameters configured specifically for the hospital. These are followed by all measured values, and then all set values.

1. Touch the **Trends/Data...** button in the main menu bar.
2. Touch the **Table** tab (A).

The trend values for the parameters (B) with the units are displayed in 7 to 8 time columns (C). Use the buttons (F) to scroll in the trend table.

### Selecting a time interval for the tabular trend display

1. Touch the button for the time interval (D).
2. Select the time interval from the selection list (5, 10, 30 minutes; 1, 2, 6, 12 hours; 1 day).

The tabular trends are available for the following times according to the selected time interval:

<table>
<thead>
<tr>
<th>Time interval</th>
<th>Availability</th>
</tr>
</thead>
<tbody>
<tr>
<td>5, 10, 30 minutes</td>
<td>1 day</td>
</tr>
<tr>
<td>1 hour</td>
<td>2 days</td>
</tr>
<tr>
<td>2, 6, 12 hours, 1 day</td>
<td>7 days</td>
</tr>
</tbody>
</table>

### Displaying the value of a parameter at a certain moment in time

- Position the cursor (E) on the time by turning the rotary knob or touching the time.

### Additional information

"Configuring the display of hospital-specific measured values and settings" on page 160.

The **Table** page can be configured as the **Trends table** button in the main menu bar to enable direct access. See "Assigning functions to additional buttons" on page 161.
Displaying data

The following data can be displayed:
- Hospital-specific data
- Measured values 1
- Measured values 2
- Set values

Measured values are displayed on a blue background and set values on a green background.

Displaying hospital-specific data

1. Touch the **Trends/Data...** button in the main menu bar.
2. Touch the **Values** tab (A).

Babylog VN500 opens the page containing the current, hospital-specific measured and set values (B).

Babylog VN500 displays the hospital-specific measured and set values (C) selected in the system setup.

Additional information

"Configuring the display of hospital-specific measured values and settings" on page 160.

Displaying all measured values

Prerequisite: The **Values** page (A) is opened.

- Touch the **Values 1 (B)** or **Values 2 (C)** tab.

Displaying set values

Prerequisite: The **Values** page (A) is opened.

- Touch the **Settings** tab (B).

Additional information

The **Values** page can be configured as the **Values** button in the main menu bar to enable direct access. See "Assigning functions to additional buttons" on page 161.
Displaying the logbook

The logbook records changes, events and alarms in chronological order. A maximum of 5000 logbook entries is possible. Events include, for example, use of the medication nebulizer or flow calibration. For alarms only the occurrence of the alarm condition is recorded, not its termination.

The entries in the logbook are also retained after the device has been switched off and on again or following a power supply failure.

1. Touch the Trends/Data... button in the main menu bar.
2. Touch the Logbook tab (A).

Displaying the setting parameters at another moment in time

- Select the row by turning the rotary knob or touching the row.

With the button (D) the cursor will be moved backwards or forwards by at least 24 hours.

Additional information

The Logbook page can be configured as the Logbook button in the main menu bar to enable direct access. See "Assigning functions to additional buttons" on page 161.

Babylog VN500 opens the logbook. The cursor (B) marks a row in the logbook. The marked row corresponds with the cursor position in the trend display.

For the marked row Babylog VN500 displays all the set values of the ventilation mode effective at this time in the field (C).
Data export

The data export takes place via a USB storage medium. A maximum of 5000 logbook entries from the last 7 days can be exported.

1. Insert the USB storage medium into a USB port on Infinity C500.
2. Touch the Trends/Data... button in the main menu bar.
3. Touch the Export data tab (A).

The following data can be exported:
- Current settings and measured values (B)
- Results obtained from the device check (C)
- Results obtained from the breathing circuit check (D)
- Logbook 1 day or 7 days (E)
- Alarm history 1 day or 7 days (F)
- Trends 1 day or 7 days (G)

4. Touch the appropriate button for the export of the related data.
5. Touch the All data button (I) for the export of all the data.

The data is exported to the USB storage medium. After the successful completion of the data export, Babylog VN500 displays a message in the message field (H).

After the data export
- Remove the USB storage medium from the USB port after waiting at least 2 seconds.

If data export was not successful
If data export fails owing to the USB storage medium being full, the buttons are deactivated.
- Remove the USB storage medium from the USB port and use a different USB storage medium.

Additional information
The buttons are deactivated when a USB storage medium is not connected.
The exported files can only be viewed with a Unicode-enabled editor and a Unicode font.
An import into word processors or spreadsheets is possible.
Monitoring

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Information on monitoring

Monitoring is activated at the factory. Each monitoring function can be deactivated separately. O2 monitoring and flow monitoring are switched on after the device is switched on.

Possible displays for measured values

Instead of a measured value, the following displays are possible in the parameter fields or tables:

<table>
<thead>
<tr>
<th>Display</th>
<th>Cause</th>
</tr>
</thead>
<tbody>
<tr>
<td>Off</td>
<td>Monitoring deactivated by user</td>
</tr>
<tr>
<td>ERR</td>
<td>Sensor error</td>
</tr>
<tr>
<td>CAL</td>
<td>Calibration active, no measured value display possible</td>
</tr>
<tr>
<td>Measured value?</td>
<td>Reduced sensor accuracy</td>
</tr>
<tr>
<td>No measured value</td>
<td>Prerequisites for measurement or calculation currently not met</td>
</tr>
<tr>
<td>+++</td>
<td>Measured value above specified measurement range</td>
</tr>
<tr>
<td>---</td>
<td>Measured value below specified measurement range</td>
</tr>
</tbody>
</table>

Display of etCO2 measurements

The measured value for etCO2 can be displayed in Vol%, kPa or mmHg. The display is configurable, see "Configuring units" on page 184.

Information on the sensors used

Babylog VN500 uses the following sensors for measurement and monitoring purposes:
- Neonatal flow sensor
- O2 sensor
- Pressure sensor
- CO2 sensor

CAUTION

Regular calibration is essential to ensure that the sensors deliver reliable and accurate results. Otherwise the proper functioning of the device may be impaired.

Automatic calibration of the pressure sensors takes place immediately and an hour after the device has been switched on, afterwards every 12 hours.

For calibrating or checking the other sensors, see:
- "Calibrating the neonatal flow sensor" on page 143
- "Calibrating the O2 sensor" on page 146
- "Information on checking the CO2 sensor" on page 148

The calibration or zero-checking values of the sensors that were last determined remain stored until the next calibration or zero check, even if the device is switched off.
Flow monitoring

Information on flow monitoring

The measured values for $MVe$ and $VTe$ are not leakage-corrected and are therefore lower than the actual minute and tidal volumes applied to the patient if a leakage occurs. When leakage compensation is activated, the measured volume and flow values as well as the curves for flow and volume are displayed with leakage correction.

Babylog VN500 compensates for leakages up to 100% of the set tidal volume $VT$. Pressure-controlled ventilation is recommended in the case of larger leakages.

In order to avoid false alarms and assure proper monitoring, the following settings are required:

- Adjust both alarm limits for $MVe$ in line with the current value.
- Use additional monitoring, e.g., external SpO2, if necessary.

**CAUTION**

Patient hazard!

Use additional external monitoring during ventilation with very low tidal volumes.

Before each manual calibration, whether started from the device check or from the Sensors/Parameters dialog window, Babylog VN500 automatically cleans the neonatal flow sensor by heating.

**WARNING**

Risk of fire

Residual vapors of easily flammable disinfectants (e.g., alcohols) and deposits that were not removed during reprocessing can ignite when the flow sensor is in use.

- Ensure particle-free cleaning and disinfection.
- After disinfection, allow the flow sensor to air for at least 30 minutes.
- Before inserting the flow sensor check for visible damage and soiling, such as residual mucus, medication aerosols, and particles.
- Replace flow sensors when damaged, soiled, or not particle-free.

Calibrating the neonatal flow sensor

Calibration of the neonatal flow sensor corresponds to a zero calibration.

Manual calibration of the neonatal flow sensor is necessary:
- During the device check and before use
- At least once a day
- After replacing the neonatal flow sensor
- After medication nebulization

Recalibration is not necessary if the neonatal flow sensor has been unplugged only briefly.
### Starting calibration of the neonatal flow sensor

1. Touch the **Sensors/Parameters**... button in the main menu bar.

   Babylog VN500 opens the **Sensors/Parameters** dialog window. The **Neonatal flow sensor** page (A) appears by default.

2. Touch the **Y flow sensor** (E) or **ISO 15 flow sensor** button (F).

3. Touch the **Start** button (B).

   The instruction field (G) displays the instructions for performing calibration. Button (H) is preselected.

### Removing the neonatal flow sensor

4. Remove the tube connector.

5. Put on a sterile glove.

6. Seal the neonatal flow sensor ISO 15 (I) or neonatal flow sensor Y-piece (J).

   This ensures that the requirement for calibration (flow = 0) is met.

### Performing calibration

7. Press the rotary knob.

   Babylog VN500 calibrates the neonatal flow sensor.

### After calibration of the neonatal flow sensor

8. Connect the tube connector.

### Setting the flow trigger

- Touch the **Trigger** button (K).

   Babylog VN500 opens the page for setting the flow trigger. For additional information, see "Additional settings for ventilation" on page 89.

### Additional information

The **Neonatal flow sensor** page can be configured as the **Neonatal flow sensor** button in the main menu bar to enable direct access. See "Assigning functions to additional buttons" on page 161.
Deactivating or activating flow monitoring

The ventilation functions and ventilation monitoring are limited when flow monitoring is deactivated.

**WARNING**
If flow and volume monitoring is deactivated, ensure that appropriate substitute monitoring is available immediately. The patient may otherwise be jeopardized.

**WARNING**
No apnea monitoring takes place when flow monitoring is deactivated. Use an independent apnea monitoring.

**CAUTION**
Patient-triggered ventilation is not possible when flow monitoring with the neonatal flow sensor is deactivated.

Flow monitoring can be deactivated, e.g.:
- If medication nebulization is being performed
- To permit ventilation in the event of major tube leakage
- If the flow sensor has failed but cannot be replaced immediately.
A defective or disconnected flow sensor can lead to deviations in the minute and tidal volumes or to auto-triggering.

In the **Neo**. patient category, Babylog VN500 deactivates flow monitoring when changing to the **NIV** application mode.

### Deactivating flow monitoring with neonatal flow sensor

1. Touch the **Sensors/Parameters...** button in the main menu bar.
2. Touch the **Neonatal flow sensor** tab (A).
3. Touch the **Off** button (B) and confirm with the rotary knob.

Babylog VN500 displays the following information in the message field (C): **External monitoring must be used!**

Flow monitoring with the neonatal flow sensor is deactivated. Babylog VN500 displays the symbol \( \text{Flow} \) in the header bar. The measured values are no longer displayed. The alarm function is deactivated.

### Activating flow monitoring with neonatal flow sensor

Reactivate flow monitoring after exchanging the neonatal flow sensor or as soon as possible.

Prerequisite: The **Neonatal flow sensor** page (A) is opened.
- Touch the **On** button (D) and confirm with the rotary knob.

Flow monitoring is activated.
O2 monitoring

Calibrating the O2 sensor

The O2 sensor is calibrated during the device check. Regular calibration during the device check ensures the specified accuracy. If the O2 sensor is not calibrated for 3 months, the accuracy of the O2 sensor will be reduced. The parameter field for the O2 concentration displays a question mark in addition to the measured value.

After calibration during the device check the sensor will work again with full accuracy.

CAUTION
If the quality of the oxygen from the central gas supply system is inadequate, calibrate the O2 sensor with calibration gas (100 % O2). Otherwise this may result in an incorrect calibration.

Additional information

"Performing the device check“ on page 71.

The O2 sensor page can be configured as the O2 sensor button in the main menu bar to enable direct access. See "Assigning functions to additional buttons" on page 161.

The O2 sensor is deactivated in standby mode. When the therapy is started, the O2 concentration is not displayed until after about 5 seconds.

Deactivating or activating O2 monitoring

WARNING
If O2 monitoring is deactivated, ensure that appropriate substitute monitoring is available immediately. The patient may otherwise be jeopardized.

O2 monitoring can be replaced by appropriate substitute monitoring. Set the O2 alarm limits for the substitute monitoring according to the set value for FiO2:

| FiO2 <60 Vol% | O2 ±4 Vol% |
| FiO2 ≥60 Vol% | O2 ±6 Vol% |

Deactivating O2 monitoring

1 Touch the Sensors/Parameters... button in the main menu bar.
2 Touch the O2 sensor tab (A).
3 Touch the Off button (B) and confirm with the rotary knob.

Babyllog VN500 displays the following information in the message field (C): **External monitoring must be used!**

O2 monitoring is deactivated. Babyllog VN500 displays the symbol FiO2 in the header bar. The measured values are no longer displayed. The corresponding alarm function is deactivated.
Activating O2 monitoring

Reactivate O2 monitoring as soon as possible.
Prerequisite: The O2 sensor page (A) is opened.

- Touch the On button (D) and confirm with the rotary knob.

O2 monitoring is activated.

CO2 monitoring

Information on CO2 monitoring

The CO2 sensor page can be configured for direct access into the main menu bar as the CO2 sensor button. See “Assigning functions to additional buttons” on page 161.

Selecting the cuvette type

The following cuvettes can be used:
- Reusable cuvettes
- Disposable cuvettes

The cuvette type used must be selected on the Zero calib. on/off page.

CAUTION

The cuvette windows of the reusable cuvette have different optical properties to the cuvette windows of the disposable cuvette. The cuvette type used must therefore be selected correctly on the Zero calib. on/off page. Otherwise the zero point will be shifted by up to 8 mmHg CO2.

1. Touch the Sensors/Parameters... button in the main menu bar.
2. Touch the CO2 sensor tab (A).

The Zero calib. on/off page (B) appears by default.

3. Touch the Reusable (C) or Disposable button(D).

If the selected cuvette does not correspond to the cuvette used, the alarm message Check CO2 cuvette is displayed.
Monitoring

Information on checking the CO₂ sensor

The CO₂ sensor is factory-calibrated and can be used on any Babylog VN500.

Information on checking the zero indication and zero calibration

When checking the zero indication or performing zero calibration, the CO₂ concentration in the cuvette or in the cuvette slot of the sensor must not be higher than the background concentration in rooms of approximately 0.4 mmHg or 0.05 Vol%.

For this reason, do not breathe on or into the cuvettes or into the cuvette slot.

The following checks are required for the CO₂ sensor:

<table>
<thead>
<tr>
<th>Check</th>
<th>When required?</th>
</tr>
</thead>
<tbody>
<tr>
<td>Check CO₂ zero indication in ambient air</td>
<td>Required before measurement and when changing the CO₂ sensor to another ventilation unit.</td>
</tr>
<tr>
<td>Perform a CO₂ zero calibration</td>
<td>If the CO₂ zero indication in ambient air is not between 0 and 1 mmHg (or 0 and 0.1 Vol%, or 0 and 0.1 kPa).</td>
</tr>
<tr>
<td>Check calibration of the CO₂ sensor with a test filter</td>
<td>Required in intervals of one month.</td>
</tr>
<tr>
<td>Check calibration of the CO₂ sensor with test gas</td>
<td>When the test values are not adhered to during the calibration check with test gas.</td>
</tr>
<tr>
<td>Perform calibration of the CO₂ sensor</td>
<td>When the test values are not adhered to during the calibration check with test gas.</td>
</tr>
</tbody>
</table>

Zero calibration in ambient air, calibration check with test filter or test gas and calibration of the CO₂ sensor can be performed during ventilation.

Information on the alarm messages issued during CO₂ monitoring

This information refers to the alarm messages which are generated due to a soiled cuvette or sensor.

Alarm message Clean CO₂ cuvette

If the Clean CO₂ cuvette message is displayed, the following panes may be soiled:

- Cuvette (disposable or reusable cuvette)
- CO₂ sensor

1. Clean the cuvette or use another cuvette.
   - When using reusable cuvettes, insert a clean reusable cuvette.
   - When using disposable cuvettes, insert a new disposable cuvette.

2. Clean the CO₂ sensor.

Alarm message CO₂ zero calibration?

If the CO₂ zero calibration? message is displayed or if incorrect measured values are suspected, e.g., etCO₂ values too low or inspiratory values too high, then proceed as follows:

1. Check whether the cuvette windows are soiled.
2. Clean the soiled windows. Or, if a reusable cuvette was used previously, use a clean reusable cuvette. If a disposable cuvette was used previously, use a new disposable cuvette.

If the cuvette windows are extremely soiled, e.g., from deposits due to medication nebulization, this may result in a zero shift. The CO₂ measured values may be incorrect even before insufficient measuring light causes the Clean CO₂ cuvette message to appear.

If the CO₂ zero calibration? message does not disappear or if the measured CO₂ values remain suspect, a zero calibration must be performed.
Checking the CO₂ zero indication

The check of the CO₂ zero indication in ambient air is performed with a clean CO₂ sensor that is placed on the cuvette used for measurement.

Prerequisite: Babylog VN500 is switched on and at least the three-minute warm-up phase for the CO₂ sensor has elapsed. After 3 minutes, the measured CO₂ values will be inside the specified tolerance range.

1. Fit the CO₂ sensor on the cuvette.
2. Select the cuvette type, see page 147.
3. To display CO₂ measured values as a curve, see "Changing the display of monitoring fields" on page 95.
4. Remove the CO₂ sensor with the cuvette from the breathing circuit and hold it in ambient air. Do not breathe on or into the cuvette.
   Instead of the cuvette from the breathing circuit, another clean cuvette of the selected type (disposable or reusable) can be used.
5. Observe the measured CO₂ value. If 0 to 1 mmHg (or 0 to 0.1 Vol% or 0 to 0.1 kPa) is not displayed in the ambient air, perform a zero calibration.

Performing a CO₂ zero calibration

Zero calibration is performed in ambient air and with a clean CO₂ sensor which is removed from the cuvette.

Prerequisite: Babylog VN500 is switched on and at least the three-minute warm-up phase for the CO₂ sensor has elapsed. After 3 minutes, the measured CO₂ values will be inside the specified tolerance range.

Starting zero calibration

1. Touch the Sensors/Parameters... button in the main menu bar.
2. Touch the CO₂ sensor tab (A).
   The Zero calib. on/off page (B) appears by default.
3. Touch the Start button (C).
   When requested by Babylog VN500:
   4. Remove the CO₂ sensor (D) from the cuvette (E).
   5. Confirm with the rotary knob.
   Babylog VN500 performs the zero calibration and displays the message Calibration in progress.
Monitoring

If zero calibration was successful
After approximately 5 seconds, Babylog VN500 reports *Zero calibration successful.*
- Fit the CO₂ sensor (D) back on the cuvette (E).

If zero calibration was not successful
Babylog VN500 reports *Zero calibration failed.*
- Repeat zero calibration.

If zero calibration is still impossible
1. Check whether the sensor is soiled and clean it if necessary. If the sensor is faulty, replace it.
2. Repeat zero calibration.

Checking the calibration of the CO₂ sensor with a test filter
Perform the calibration check of the CO₂ sensor with a test filter at intervals of one month.

Before the check
Prerequisite: Babylog VN500 is switched on and at least the three-minute warm-up phase for the CO₂ sensor has elapsed.
- Perform CO₂ zero calibration in ambient air.

Starting the calibration check of the CO₂ sensor with a test filter
Prerequisite: The CO₂ sensor page (A) is opened.
1. Touch the *Check sensor* tab (B).
2. Remove the sensor from the cuvette and connect it to the test filter (E) on the sensor cable.
3. Touch the *Filter check* button (C) and confirm with the rotary knob.
Babylog VN500 starts the check and displays the progress and result of the check in the message field (D).

If the check was successful
Babylog VN500 displays the message *Filter check successful.* The test value is within the permissible tolerance.
- Fit the CO₂ sensor back on the cuvette.

If the check was not successful
Babylog VN500 displays the message *Filter check failed.* The test value is outside the permissible tolerance.
- Check the CO₂ calibration with test gas.
Checking the calibration of the CO2 sensor with test gas

Perform the check when the test values are not within the permitted tolerance during the calibration check of the CO2 sensor with test filter.

**CAUTION**
For the check and calibration only use a test gas which consists of CO2 and N2! Otherwise display deviations of ±0.5 Vol% CO2 may occur.

**Before the check**

Prerequisite: Babylog VN500 is switched on and at least the three-minute warm-up phase for the CO2 sensor has elapsed.

- Perform CO2 zero calibration in ambient air.

**Connecting the test gas supply**

The test gas must consist only of CO2 and N2!

1. Use the reusable cuvette from the calibration set!

At the start of the check with test gas, Babylog VN500 automatically sets the cuvette type to *Reusable*.

2. Connect the test gas cylinder (A) and the cuvette (B) of the calibration set to the hose (C).

3. Fit the CO2 sensor (D) on the cuvette (B) from the calibration set.

4. Read the CO2 concentration of the test gas from the test gas cylinder (A).

5. Open the test gas cylinder (E) and set the test gas flow to 0.1 L/min.

**Starting the calibration check of the CO2 sensor with test gas**

Prerequisite: The **CO2 sensor** page (F) is opened.

1. Touch the **Check sensor** tab (G).

2. Touch the **Gas check** button (H).

Babylog VN500 displays the measured CO2 concentration (I).

About 1 minute after the test gas flow has been set, the measured CO2 value must match the CO2 content of the test gas read from the test gas cylinder with a tolerance of ±0.2 Vol%.

Babylog VN500 terminates the check with test gas approx. 1 minute after the start.

3. Close the test gas cylinder again.

If the test value is outside the permitted tolerance, the CO2 sensor must be recalibrated with test gas.
After the calibration check of the CO2 sensor with test gas
The cuvette type is automatically reset to the previously set cuvette type.
- Fit the CO2 sensor back on the cuvette in the breathing circuit.

Performing the calibration of the CO2 sensor
The CO2 sensor must be calibrated if the test values are not within the permitted tolerance during the calibration check with test gas.

CAUTION
For the check and calibration only use a test gas which consists of CO2 and N2! Otherwise display deviations of ±0.5 Vol% CO2 may occur.

Before calibration
Prerequisite: Babylog VN500 is switched on and at least the three-minute warm-up phase for the CO2 sensor has elapsed.
- Perform CO2 zero calibration in ambient air.

Connecting the test gas supply
The test gas must consist only of CO2 and N2!
1 Use the reusable cuvette from the calibration set!
At the start of calibration, Babylog VN500 automatically sets the cuvette type to Reusable.

Starting calibration of the CO2 sensor with test gas
Prerequisite: The CO2 sensor page (F) is opened.
1 Touch the Calibration tab (G).

Sensors/Parameters

J

2 Connect the test gas cylinder (A) and the cuvette (B) of the calibration set to the hose (C).
3 Fit the CO2 sensor (D) on the cuvette (B) from the calibration set.
4 Read the CO2 concentration of the test gas from the test gas cylinder (A).
5 Open the test gas cylinder (E) and set the test gas flow to 0.1 L/min.

2 Touch the CO2 sensor therapy control (H). Enter the value for the CO2 concentration in the test gas with the rotary knob and confirm.
3 About 1 minute after setting the test gas flow, touch the Start button (I) and confirm with the rotary knob.
Babylog VN500 starts the calibration of the CO2 sensor and displays the progress and result of the calibration in the message field (J).

4 Close the test gas cylinder again.

If calibration was successful
Babylog VN500 displays the message **CO2 sensor calib. with test gas successful.**
The cuvette type is automatically reset to the previously set cuvette type.
- Fit the CO2 sensor back on the cuvette in the breathing circuit.

If calibration was not successful
Babylog VN500 displays the message **Calibration of CO2 sensor with test gas failed.**
If calibration failed, the following causes are possible:

<table>
<thead>
<tr>
<th>Cause</th>
<th>Remedy</th>
</tr>
</thead>
<tbody>
<tr>
<td>The CO2 concentration entered does not match the value on the test gas cylinder.</td>
<td>Check the CO2 concentration entered and repeat calibration of the CO2 sensor.</td>
</tr>
<tr>
<td>The test gas cylinder is empty.</td>
<td>Use a new test gas cylinder and repeat calibration of the CO2 sensor.</td>
</tr>
<tr>
<td>The CO2 sensor is soiled.</td>
<td>Clean the CO2 sensor and repeat calibration of the CO2 sensor.</td>
</tr>
<tr>
<td>The CO2 sensor is faulty.</td>
<td>Replace the CO2 sensor and check the CO2 zero indication.</td>
</tr>
</tbody>
</table>

**Resetting the calibration of the CO2 sensor**
If problems occurred during calibration, the sensor can be reset to the delivery default values.
Prerequisite: The **Calibration** page is opened.
1 Touch the **Reset calibration** button (K) and confirm with the rotary knob.
After approximately 5 seconds, the factory-set calibration value is effective again and must be checked with the test filter.
2 Check the calibration of the CO2 sensor with test filter, see page 150.

**Deactivating or activating CO2 monitoring**
Deactivate CO2 monitoring when a defective CO2 sensor cannot immediately be exchanged or the CO2 measured values are currently not needed.

Deactivating CO2 monitoring
1 Touch the **Sensors/Parameters...** button in the main menu bar.
2 Touch the **CO2 sensor** tab (A).
3 Touch the **Zero calib. on/off** tab (B) if the page is not already preset.
4 Touch the **Off** button (C) and confirm with the rotary knob.

CO2 monitoring is deactivated. Babylog VN500 displays **Off** in the CO2 parameter field. The measured values are no longer displayed. The alarm function is deactivated.
Activating CO₂ monitoring

Prerequisite: The Zero calib. on/off page (B) is opened.

- Touch the On button (D) and confirm with the rotary knob.

CO₂ monitoring is activated.
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Information on configuration

The **System setup** dialog window provides the user with the following configuration options:

- Screen layout
- Alarms
- Ventilation
- Config. exchange (Importing and exporting configurations)
- Applications
- System status
- Exchange intervals
- System

To prevent unauthorized adjustments, the following pages are password-protected:

- Screen layout > Views
- Alarms
- Ventilation
- Config. exchange
- Applications
- Exchange intervals

The password only needs to be entered once as long as the **System setup** dialog window remains open.

For additional information on the password, see page 341.

Configuring the screen display

The following settings can be configured on the **System setup > Screen layout** page:

- General settings (General settings)
- Views
- Customized data (Customized data)
- Config. buttons (Configurable buttons)
- Trends graphic 1
- Trends graphic 2
- Therapy bar

The customized settings for Trends graphic 1 and Trends graphic 2 become effective with the admission of a new patient. The other customized screen display settings are immediately effective.

- Touch the **System setup...** button.

Babylog VN500 opens the **System setup** dialog window. The **Screen layout** page is displayed by default.
### Adjusting illumination and brightness

Prerequisite: The **Screen layout** page (A) is opened.

1. Touch the **General settings** tab (B).

#### Automatic changeover of day and night mode

If the automatic changeover of day and night mode is switched on, the system will automatically change over the following settings:
- Illumination of the screen
- Volume of the alarm tone
- Automatic increase of the alarm tone volume

- Touch button **On** (C) or **Off** (D).

#### Selecting the time period for screen illumination at night

The illumination of the screen is reduced with a dark background color for the time period entered.

Hours (E) : minutes (F) to hours (G) : minutes (H).

1. Touch the appropriate button.
2. Set the time by turning the rotary knob and push to confirm.

If the automatic changeover for the illumination of the screen is switched on, the system will change over at the times entered.

The **Day/Night** button in the main menu bar can be configured to enable direct access to the reduced screen illumination mode that uses a dark background color, see page 161.

### Adjusting screen brightness

The screen brightness can be adjusted automatically or manually.

#### Activating automatic brightness adjustment:

- Touch the **Automatic** button (I).

#### Adjusting brightness manually:

1. Touch the **Manual** button (J).
2. Touch the button (K).
3. Set the value by turning the rotary knob and push to confirm.

### Adjusting automatic screen dimming

Automatic dimming of the screen can be set for standby mode and battery operation.

1. Touch the **On** button (L).
2. Touch the button (N).
3. Set the value by turning the rotary knob and push to confirm.

#### Switching off automatic screen dimming:

- Touch the **Off** button (M).
Configuration

Configuring the screen view

Prerequisite: The Screen layout page (A) is opened.

1  Touch the Views tab (B).
2  Enter password and confirm with Enter.

The Views page is displayed.

Adjusting the selected view

7  Touch a field in the view (E).
The dialog for the field contents is displayed.

8  Select the parameter, display format and display size for curves and parameter fields.
See "Changing the display of monitoring fields" on page 95.

Locking the view against overwriting

●  Touch the button (F).
The selected view is locked and cannot be changed. The display of the monitoring fields cannot be changed on the main page.

Deactivating the lock

●  Touch the button (G).

Saving the view

1  Touch the button (H).
2  Confirm with the rotary knob.
The current view for the selected view (View 1 to View 3) is saved.

Reset current view

Each view can be reset individually, either to factory or saved settings. The view must not be locked.

Loading factory settings

1  Touch the Dräger default button (I).
2  Confirm with the rotary knob.

For information on the factory settings for the views, see chapter "Factory-set screen views" on page 335.

Loading saved settings

1  Touch the Load button (J).
2  Confirm with the rotary knob.
### Overview of format templates

The following format templates are available for selection:

![Format templates diagram](image)

The (K), (L) and (M) fields can be configured with customized settings. All fields can also be configured without contents. The following settings are possible:

<table>
<thead>
<tr>
<th>K</th>
<th>L</th>
<th>M</th>
</tr>
</thead>
<tbody>
<tr>
<td><strong>Field size 1x</strong></td>
<td><strong>Field size 2x</strong></td>
<td><strong>Field size 1x</strong></td>
</tr>
<tr>
<td>Waveform</td>
<td>Waveform</td>
<td>Single parameter</td>
</tr>
<tr>
<td>Trends (meas.)</td>
<td>Loop</td>
<td>Double parameter</td>
</tr>
<tr>
<td>Trends (settings)</td>
<td>Double loop</td>
<td>Trends (meas.)</td>
</tr>
<tr>
<td>Trends table</td>
<td>Trends (meas.)</td>
<td>Trends (settings)</td>
</tr>
<tr>
<td>Multi Trend</td>
<td>Trends (settings)</td>
<td></td>
</tr>
<tr>
<td>Alarm history</td>
<td>Trends table</td>
<td></td>
</tr>
<tr>
<td>Lung display (Smart Pulmonary View)</td>
<td>Multi Trend</td>
<td></td>
</tr>
<tr>
<td></td>
<td>Alarm history</td>
<td></td>
</tr>
<tr>
<td></td>
<td>Lung display (Smart Pulmonary View)</td>
<td></td>
</tr>
</tbody>
</table>
Configuring the display of hospital-specific measured values and settings

A maximum of 20 measured and set values can be grouped together. The hospital-specific measured and set values are displayed on the **Trends/Data > Values > Customized data** page.

Prerequisite: The **Screen layout** page (A) is opened.

1. Touch the **Customized data** tab (B).

**System setup**

Selecting a row in the list

- Turn the rotary knob until the desired row is marked in column 1 or 2 (C) or touch the row.

Configuring the display of hospital-specific measured values

Prerequisite: The desired row is marked.

1. Touch the **Values** button (D).
2. Select the parameter from the selection list with the rotary knob and push to confirm.

Configuring the display of hospital-specific set values

Prerequisite: The desired row is marked.

1. Touch the **Settings** button (E).
2. Select the parameter from the selection list with the rotary knob and push to confirm.

**Additional information**

Measured values are displayed in the list with a blue background color and set values with a green background color.
Assigning functions to additional buttons

Additional buttons can be assigned in the main menu bar to enable direct access to a function or to directly open a page. The buttons are spatially assigned to the corresponding group.

Prerequisite: The Screen layout page (A) is opened.

1. Touch the Config. buttons tab (B).

![Diagram of System setup with tabs and columns]

In the first column (C), the buttons can be selected for the left column of the main menu bar. In the second column (D), the buttons can be selected for the right column of the main menu bar.

2. Touch the button.

3. Select the desired button from the selection list with the rotary knob and push to confirm.

Babylog VN500 displays the selected button in the main menu bar.

Additional information

For information on the display of additional buttons and their location in the main menu bar, see "Main menu bar structure" on page 331.
Selecting parameters for the graphic trend display

The graphic trend display for the Trends/Data > Trends > Graphics 1 and Graphics 2 pages can be configured. The settings become effective with the admission of a new patient.

Prerequisite: The Screen layout page (A) is opened.

1. Touch the Trends graphic 1 (B) or Trends graphic 2 (C) tab.

Each page contains 2 graphic trend displays (D). A maximum of 3 parameters can be configured for each trend display.

Configuring the trend display

1. Touch the button (E).

Babylog VN500 opens the Setup dialog with the buttons for Meas. (F) and Settings (G).

The measured values (F) are divided into the following parameter types:
- Pressures
- Minute vol.
- Volume/Flow
- Gases
- Timing/Cycl.
- Others
- Events

The settings (G) are divided into the following parameter types:
- Pressures
- Volume/Flow
- Gases
- Timing/Cycl.
- Others

2. Touch the appropriate button for measured values or set values.
Babylog VN500 opens another dialog containing all the parameters of the selected parameter type (example Volume/Flow).

3 Touch the parameter. The button turns dark green.
4 Confirm with the OK button (K).

The dialog for the parameter type is closed.
A maximum of 3 parameters can be selected for each graphic trend display.
5 Select further parameters according to step 2 to 4.
6 Confirm the selected measured values and set values with the OK button (H).

The selected parameters are displayed in the trend display. The Setup dialog is closed.

The selection can be canceled with Cancel (I). The previous selection is displayed in the graphic trend. Clear (J) can be used to delete all parameter selections made.

Deselecting a parameter in the trend display

● Touch the parameter to be deselected in the parameter type dialog. The button turns pale green.

Selecting a time interval

● Touch the button (L). Select the time interval from the selection list: 2, 4, 8, 12 hours, 1 day, 7 days.
Locking therapy controls in the therapy bar

The therapy controls in the therapy bar can be locked to prevent accidental changes from being made to the ventilation parameters.

Prerequisite: The Screen layout page (A) is opened.

1 Touch the Therapy bar tab (B).

2 Touch the On button (C).

The therapy controls in the therapy bar are locked. The ventilation parameters can only be changed in the Ventilation settings dialog window.

Canceling the lock

• On the Therapy bar page, touch the Off button (D).
Configuring alarm settings

The customized settings for the start-up values of the alarm limits become effective with the admission of a new patient. The customized alarm tone settings are effective immediately depending on the time of day. The selection of the alarm tone sequence is effective immediately.

1. Touch the System setup... button in the main menu bar.
2. Touch the Alarms tab.
3. Enter password and confirm with Enter.

Babylog VN500 displays the following configurable alarm settings in the overview:
- Start-up values for alarm limits
- Alarm volume and alarm tone

CAUTION

If several devices of the same type are used on a ward, the alarm defaults must be configured identically on all devices. The patient may otherwise be jeopardized.

Prerequisite: The Alarms page (A) is opened.
1. Touch the Preset limits tab (B).

Selecting the factory settings

- Touch the Dräger default button (E) and confirm with the rotary knob.

The Dräger default button also resets other start-up settings on the Ventilation page and the Alarms page to the factory settings.
Table of alarm limits

The following table lists the alarm limits with the setting range and the factory-set start-up values (Dräger default).

<table>
<thead>
<tr>
<th>Parameter</th>
<th>Setting range</th>
<th>Factory-set start-up value (Dräger default)</th>
</tr>
</thead>
<tbody>
<tr>
<td>MVe</td>
<td>1 to 100 %</td>
<td>(VT x RR) +50 %</td>
</tr>
<tr>
<td>MVe</td>
<td>Off, 1 to 100 %</td>
<td>(VT x RR) –20 %</td>
</tr>
<tr>
<td>MV delay</td>
<td>0 to 20 seconds</td>
<td>0 seconds</td>
</tr>
<tr>
<td>MV delay</td>
<td>0 to 20 seconds</td>
<td>0 seconds</td>
</tr>
<tr>
<td>Paw</td>
<td>7 to 105 mbar (7 to 105 cmH2O)</td>
<td>30 mbar (30 cmH2O)</td>
</tr>
<tr>
<td>RR</td>
<td>1 to 100 %, Off</td>
<td>RR +20 %</td>
</tr>
<tr>
<td>Tapn</td>
<td>5 to 60 seconds</td>
<td>15 seconds</td>
</tr>
<tr>
<td>etCO2</td>
<td>0.1 to 13.1 Vol%</td>
<td>8.0 Vol%</td>
</tr>
<tr>
<td></td>
<td>1 to 98 mmHg</td>
<td>60 mmHg</td>
</tr>
<tr>
<td></td>
<td>0.1 to 13.3 kPa</td>
<td>–</td>
</tr>
<tr>
<td>etCO2</td>
<td>0 to 13.0 Vol%</td>
<td>4.0 Vol%</td>
</tr>
<tr>
<td></td>
<td>0 to 97 mmHg</td>
<td>30 mmHg</td>
</tr>
<tr>
<td></td>
<td>0 to 13.2 kPa</td>
<td>–</td>
</tr>
</tbody>
</table>

Additional information

The alarm limits for the minute volume are set as a percentage of the start-up value (VT x RR). To configure RR, see “Configuring start-up settings for the ventilation parameters” on page 171.

For an overview of the device’s internal alarm limits, see chapter “Automatic alarm limits” on page 282.
Setting the alarm tone

**WARNING**
Unnoticed alarms in loud environments
Alarm situations are not recognized.
Set the volume of the alarm tone so that alarms can be heard.

Prerequisite: The **Alarms** page (A) is opened.
1 Touch the **Alarm vol./tone** tab (B).

**Setting the minimum alarm volume**
Configuring the minimum alarm volume sets the lower limit of the factory setting for the volume of the alarm tone (10 to 100 %). This allows the setting range to be adjusted to the acoustical situation at operation site.
2 Touch the button (C).
3 Set the value for the minimum volume by turning the rotary knob and push to confirm.

**Setting the volume for day or night**
4 For the day setting, touch the button (D).
5 Set the value for the sound level by day by turning the rotary knob and push to confirm.
6 For the night setting, touch the button (E).
7 Set the value for the sound level by night by turning the rotary knob and push to confirm.

**Activating the automatic sound level increase**
The **Auto increase** function can be set separately for day and night.
- Touch the appropriate button **Auto increase** (F).

**Selecting alarm tone sequences**
Babylóg VN500 offers the following alarm tone sequences:
- Touch the appropriate button.

**Setting the priority of the battery alarms**
The device offers the following priorities for battery alarms:
- Touch the appropriate button.

The **Battery activated** alarm message indicating the changeover to battery operation can be configured as a high or medium priority alarm when **Dräger ventilation** is selected.
- Touch the **Medium** (K) or **High** (L) button and confirm.
Depending on the setting (IEC/CEI or Dräger ventilation), alarm messages have the following priority:

<table>
<thead>
<tr>
<th>Alarm message</th>
<th>Priority IEC/CEI</th>
<th>Priority Dräger ventilation</th>
</tr>
</thead>
<tbody>
<tr>
<td>Battery activated</td>
<td>Low-priority alarm message</td>
<td>High-priority or medium-priority alarm message</td>
</tr>
<tr>
<td>Battery low</td>
<td>Medium-priority alarm message</td>
<td>High-priority alarm message</td>
</tr>
<tr>
<td>Battery discharged</td>
<td>High-priority alarm message</td>
<td>High-priority alarm message</td>
</tr>
</tbody>
</table>

**Configuring the confirmation prompt**

The display of messages and alarms requesting confirmation of ventilation settings can be switched on (M) or off (N).

- Touch the appropriate button and confirm.

**Configuring ventilation settings**

The following ventilation configurations are possible:
- Configuration of patient category for start-up
- Configuration of main ventilation modes
- Configuration of start-up ventilation settings
- Configuration of general settings for ventilation
- Configuration of settings for maneuvers

The customized ventilation settings become effective with the admission of a new patient.

1. Touch the **System setup...** button in the main menu bar.
2. Touch the **Ventilation** tab.
3. Enter password and confirm with **Enter**.
**Configuration**

**Configuring start-up settings for the patient category**

Prerequisite: The Ventilation page (A) is opened.

1. Touch the Patient category tab (B).

2. Touch the button (C).

Babylog VN500 opens the selection list. The following patient categories are available for selection:

- Pediatric patients only
- Pediatric patients, neonates
- Neonates only

3. Select the patient category with the rotary knob and push to confirm.

Babylog VN500 displays the buttons for the selected patient category on the Start and Start/Standby pages.

**Configuring a user-defined breathing circuit**

When the User-defined hose settings function is activated, a user-defined breathing circuit can be selected on the Start/Standby... > Br. circuit/Humidifier page.

Activating a user-defined breathing circuit:

- Touch the On button (D).

Deactivating a user-defined breathing circuit:

- Touch the Off button (E).

**Additional information**

Using the user-defined breathing circuit, see page 68.
**Configuration**

**Configuring start-up settings for the ventilation modes**

**CAUTION**
If the ventilation start-up values are configured differently to the Dräger standard values, this configuration must be identical on all Babylog VN500 belonging to a ward. The patient may otherwise be jeopardized.

Prerequisite: The Ventilation page (A) is opened.

1. Touch the Modes tab (B).

Babylog VN500 displays the start mode (C) and 3 ventilation modes (D). These ventilation modes are displayed in the Ventilation settings dialog window after Babylog VN500 has been started.

The ventilation mode (E) configured under Other modes is displayed as an additional mode for information purposes and can be changed in the Ventilation settings dialog window.

2. Touch the appropriate button.

Babylog VN500 opens the ventilation mode selection list.

3. Select the mode with the rotary knob and push to confirm.

If --- is configured for a ventilation mode, the corresponding page is not available in the Ventilation settings dialog window.

The same ventilation mode cannot be configured on 2 buttons.

The button with the active ventilation mode is highlighted in gray and cannot be changed. The button assignment can only be changed when another ventilation mode is activated in the Ventilation settings dialog window.

Depending on configuration, the number of displayed ventilation modes can vary between 1 and 4.
Configuring start-up settings for the ventilation parameters

Prerequisite: The Ventilation page (A) is opened.

1. Touch the Start settings tab (B).

Babylog VN500 displays the following pages for the ventilation start-up settings:
- VT, RR, Trigger
- Pressures, O₂, I:E
- Other settings
- ATC
- HFO

The VT, RR, Trigger page (C) appears by default.

Setting start-up values for VT, RR, Slope, and Flow trigger

Depending on the patient category or the patient’s weight, these start-up values can be set:
- VT
- RR
- Slope
- Flow trigger

Setting start-up values depending on the patient category

1. If not yet preset, touch the Patient button (D) and confirm with the rotary knob.

Babylog VN500 displays the start-up values for the different patient categories (E).

2. Touch the appropriate button (E).

3. Set the value by turning the rotary knob and push to confirm.

After the start of ventilation, Babylog VN500 begins ventilation with the start-up values depending on the patient category set on the Start/Standby page.

Setting start-up values depending on the body height/weight

Prerequisite: The VT, RR, Trigger page (C) is opened.

1. Touch the Weight button (F) and confirm with the rotary knob.

2. Touch the appropriate button (G).

3. Set the value by turning the rotary knob and push to confirm.

After the start of ventilation, Babylog VN500 begins ventilation with the start-up values depending on the body height set on the Start/Standby page and the ideal body weight derived from that, or with the set start-up body weight in the Neo patient category.
Configuration

Selecting the factory settings

- Touch the Dräger default button (H) and confirm with the rotary knob.

The Dräger default button also resets other start-up settings on the Ventilation page and the Alarms page to the factory settings.

Tables for start-up values

The following tables show the factory-set start-up values (Dräger default) for VT, RR, Slope and Flow trigger.

The following table applies to the selection of start-up values depending on the patient category:

<table>
<thead>
<tr>
<th>Patient category</th>
<th>VT (mL)</th>
<th>RR (1/min)</th>
<th>Slope (s)</th>
<th>Flow trigger (L/min)</th>
</tr>
</thead>
<tbody>
<tr>
<td>Neo.</td>
<td>5.0</td>
<td>60</td>
<td>0.1</td>
<td>0.3</td>
</tr>
<tr>
<td>Ped. pat.</td>
<td>50</td>
<td>29</td>
<td>0.2</td>
<td>1.0</td>
</tr>
</tbody>
</table>

The following table applies to the selection of start-up values depending on the body weight according to the Radford nomogram:

<table>
<thead>
<tr>
<th>Weight (kg)</th>
<th>VT (mL)</th>
<th>RR (1/min)</th>
<th>Slope (s)</th>
<th>Flow trigger (L/min)</th>
</tr>
</thead>
<tbody>
<tr>
<td>0.5</td>
<td>3.0</td>
<td>100</td>
<td>0.05</td>
<td>0.2</td>
</tr>
<tr>
<td>5</td>
<td>36</td>
<td>32</td>
<td>0.2</td>
<td>1.0</td>
</tr>
<tr>
<td>15</td>
<td>110</td>
<td>26</td>
<td>0.2</td>
<td>1.0</td>
</tr>
</tbody>
</table>
Setting start-up values for pressures, FiO2 and I:E
Prerequisite: The *Start settings* page (A) is opened.

1. Touch the *Pressures, O2, I:E* tab (B).

2. Touch the corresponding button for the parameters:
   - *Pressures* (C)
   - *APRV pressures* (D)
   - *Insp. flow* (E)
   - *FiO2* (F)
   - *I:E* (G)

3. Set the value by turning the rotary knob and push to confirm.

Selecting the factory settings
- Touch the *Dräger default* button (H) and confirm with the rotary knob.

The *Dräger default* button also resets other start-up settings on the *Ventilation* page and the *Alarms* page to the factory settings.

<table>
<thead>
<tr>
<th>Parameter</th>
<th>Factory-set start-up value (Dräger default)</th>
</tr>
</thead>
<tbody>
<tr>
<td>PEEP</td>
<td>5 mbar (5 cmH2O)</td>
</tr>
<tr>
<td>ΔPsupp</td>
<td>0 mbar (0 cmH2O)</td>
</tr>
<tr>
<td>Pinsp</td>
<td>15 mbar (15 cmH2O)</td>
</tr>
<tr>
<td>Pmax</td>
<td>40 mbar (40 cmH2O)</td>
</tr>
<tr>
<td>Flow</td>
<td>5 mbar (5 cmH2O)</td>
</tr>
<tr>
<td>Phigh</td>
<td>15 mbar (15 cmH2O)</td>
</tr>
<tr>
<td>Insp. flow</td>
<td>neo.: 6 L/min</td>
</tr>
<tr>
<td>Ped. pat.</td>
<td>10 L/min</td>
</tr>
<tr>
<td>FiO2</td>
<td>21 Vol%</td>
</tr>
<tr>
<td>I:E</td>
<td>1:2</td>
</tr>
</tbody>
</table>
Defining the start-up setting of the additional settings

Prerequisite: The Start settings page (A) is opened.

1 Touch the Other settings tab (B).

The following settings can be switched on or off:

– **Volume Guarantee** (C)
– **Apnea Ventilation** (D)

2 Touch button **On** or **Off**.

3 Confirm with the rotary knob.

A start-up value can be set for the expiratory termination criterion **Exp. term.** (E):

4 Touch the button (E).

5 Set the value by turning the rotary knob and push to confirm.

If the Dräger default button is touched on another page, e.g. the Ventilation > Start settings page or the Alarms page, the settings are also set to the factory settings.
Defining start-up settings for tube compensation

Prerequisite: The Start settings page (A) is opened.

1. Touch the ATC tab (B).

The following settings can be switched on or off:
- Tube comp. (ATC) (C)
- Expiratory compensation (D)

2. Touch the On or Off button and confirm with the rotary knob.

Inspiratory compensation can be selected for spontaneous and mandatory or only spontaneous breaths:
- Spon + mand (E)
- Only spon (F)

3. Touch the appropriate button and confirm with the rotary knob.

In the Ped. pat. patient category, select the tube type:

4. Touch the ET (G) or Trach. (H) button and confirm.

Enter the tube diameter (I) according to the selected tube type:
- ET: 2 to 8 mm
- Trach.: 2.5 to 8 mm

In the Neo. patient category, only the ET tube type (G) is available.

Enter the tube diameter (I):
- ET: 2 to 5 mm

5. Touch the relevant button for the patient category.

6. Set the value for the tube diameter by turning the rotary knob and push to confirm.

Enter degree of compensation (J) for the respective patient category: 0 to 100 %

7. Touch the relevant button for the patient category.

8. Set the value for the degree of compensation by turning the rotary knob and push to confirm.

Babylog VN500 starts with the start-up settings selected for the ventilation parameters.

The customized settings for inspiratory and expiratory compensation are immediately effective when ATC is set.

If the Dräger default button is touched on another page, the settings for inspiratory and expiratory compensation are also set to the factory settings.
Defining start-up settings for high-frequency oscillation (HFO)

Prerequisite: The **Start settings** page (A) is opened.

1. Touch the **HFO** tab (B).

2. Touch the corresponding button for the parameters:
   - \( V_{Thf} \) (C)
   - \( f_{hf} \) (D)
   - \( I:Ehf \) (E)
   - \( Ampl \ hf \) (F)

3. Set the value by turning the rotary knob and push to confirm.
Configuring general settings

Prerequisite: The Ventilation page (A) is opened.

1 Touch the General settings tab (B).

The following settings can be switched on or off:
- Leakage Compensation (C)
- Automatic return from Apnea Ventilation (D)
- Apnea Ventilation alarm (E)
- Pmax/Paw high autoset (F)
- Anti Air Shower (G)

2 Touch the On or Off button as appropriate and confirm with the rotary knob.

Select Slope adjustment (H):
- Slope (I)
- Insp. flow (J)

3 Touch the appropriate button and confirm with the rotary knob.

Babylog VN500 starts with the selected settings.

For further information on the Anti Air Shower function, see page 321.

Setting a maneuver

Prerequisite: The Ventilation page (A) is opened.

1 Touch the Maneuver tab (B).

Setting the FiO2 concentration for the suction maneuver

For the suction maneuver, FiO2 is set based on the current FiO2 concentration using a factor between 1.0 and 2.0.

2 Touch button (C) or (D).

3 Set the factor by turning the rotary knob and push to confirm.

Babylog VN500 starts with the selected start-up settings.
### Importing and exporting configurations

Babylog VN500 can export the device configuration on a USB storage medium. The configuration saved on the USB storage medium can be imported to other Babylog VN500 devices.

The following settings from the system configuration are exported and imported:

<table>
<thead>
<tr>
<th>Screen layout</th>
<th>General settings</th>
</tr>
</thead>
<tbody>
<tr>
<td>Views (^1)</td>
<td>Config. buttons</td>
</tr>
<tr>
<td>Customized data</td>
<td>Trends graphic 1</td>
</tr>
<tr>
<td></td>
<td>Trends graphic 2</td>
</tr>
<tr>
<td></td>
<td>Therapy bar</td>
</tr>
</tbody>
</table>

<table>
<thead>
<tr>
<th>Alarms</th>
<th>Preset limits</th>
</tr>
</thead>
<tbody>
<tr>
<td></td>
<td>Alarm vol./tone</td>
</tr>
</tbody>
</table>

<table>
<thead>
<tr>
<th>Ventilation</th>
<th>Patient category</th>
</tr>
</thead>
<tbody>
<tr>
<td>Modes</td>
<td>Start settings</td>
</tr>
<tr>
<td></td>
<td>VT, RR, Trigger</td>
</tr>
<tr>
<td></td>
<td>Pressures, O2, I:E</td>
</tr>
<tr>
<td></td>
<td>Other settings</td>
</tr>
<tr>
<td></td>
<td>ATC</td>
</tr>
</tbody>
</table>

<table>
<thead>
<tr>
<th>System status</th>
<th>Exchange intervals</th>
</tr>
</thead>
</table>

<table>
<thead>
<tr>
<th>System</th>
<th>Country</th>
</tr>
</thead>
<tbody>
<tr>
<td></td>
<td>Units</td>
</tr>
</tbody>
</table>

<table>
<thead>
<tr>
<th>Interface</th>
<th>LAN</th>
</tr>
</thead>
<tbody>
<tr>
<td></td>
<td>COM</td>
</tr>
<tr>
<td></td>
<td>External display</td>
</tr>
</tbody>
</table>

\(^1\) Views are only exported if the view configured was first saved on the Screen layout page. When a configuration is imported, all the current views are overwritten, including the locked views.
Preparing the configuration exchange

- Insert the USB storage medium into a USB port on Infinity C500.

1. Touch the **System setup...** button in the main menu bar.

2. Touch the **Config. exchange** tab (A).

Importing a configuration from a USB storage medium to the device

A configuration can only be imported in standby mode.

1. Switch Babylog VN500 to standby mode.

2. Select a configuration from the USB storage medium (C).

3. Touch the **Import** button (E).

4. Confirm with the rotary knob.

If there is no valid configuration saved on the USB storage medium, the system issues a message.

After the import, Babylog VN500 is switched off automatically.

5. Switch Babylog VN500 on again.

Exporting a configuration from the device to a USB storage medium

1. Touch the **Export** button (F).

2. Confirm with the rotary knob.

If the USB storage medium already contains a configuration, a message appears stating that this configuration will be overwritten.

No export is possible onto the USB storage medium if it is full. The **Export** button (F) is grayed out and cannot be activated.

Babylog VN500 reports the completion of the configuration with a low-priority alarm.

6. Check the settings of the imported configuration.

**System setup**

A **B** Configuration on the device with the date of export

C Existing configurations on the USB storage medium

D Selected configuration on the USB storage medium with the date of export

E **Import** button

F **Export** button

D E A

C B

Instructions for use Infinity Acute Care System – Babylog VN500 SW 2.n
Installing applications

Babylog VN500 can be supplemented with additional Dräger applications. The applications are installed with a SIM card.

1. Insert the SIM card into the USB SIM card reader.

2. Insert the USB SIM card reader into a USB port on Infinity C500.

3. Touch the **System setup...** button in the main menu bar.

4. Touch the **Applications** tab (A).

5. Enter password and confirm with **Enter**.

Babylog VN500 displays the already installed applications (B) and the applications available on the SIM card (C).

**Installing applications**

1. Touch the **Install** button (D).

2. Select the application from the list (C) with the rotary knob and push to confirm.

3. Install the next application (repeat steps 1 to 3).

4. After all applications are installed, restart Babylog VN500.

The installed applications are displayed in the list (B).

**Additional information**

The **Applications** page can be configured as a button in the main menu bar to enable direct access. See "Assigning functions to additional buttons" on page 161.
System status

The **System status** page contains the following information:
- General status information on maintenance and operating hours
- Exchange intervals

Displaying general status information

1. Touch the **System setup** button in the main menu bar.
2. Touch the **System status** tab (A).
3. Enter password and confirm with **Enter**.
4. Touch the **General status** tab (B).

The following information is displayed:
- Next service due (C)
- Cockpit (D)
  - Operating hours: Standby
  - Operating hours: Running
- Ventilation unit (E)
  - Operating hours: Standby
  - Operating hours: Running
  - Internal battery installation date
- Gas supply unit (GS500) (F)
  - Operating hours: Blower
  - Installation date
- Power supply unit (PS500) (G), if present
  - Installation date of the batteries in the PS500
Configuration

Configuring exchange intervals

The user can configure how much of the period of use elapses before Babylog VN500 displays a message indicating that the next exchange of an accessory is due. This depends on the device type and software version.

The exchange interval must be defined in accordance with the applicable hygiene guidelines or in accordance with the specifications of the corresponding accessory’s instructions for use.

**WARNING**  
Risk of inappropriate operating life  
Exchange monitoring only considers the actual period of use and not the current status of the Infinity ID accessory and therefore does not release the user from the responsibility of periodically checking the accessory.

The exchange interval set for exchange monitoring does not guarantee that the accessory will last until the exchange interval has expired.

Opening the exchange interval page

1. Touch the **System setup...** button in the main menu bar.
2. Touch the **System status** tab (A).
3. Enter password and confirm with **Enter**.
4. Touch the **Exchange intervals** tab (B).

The exchange interval and the service life already elapsed for the accessory (C) are displayed.

Setting the exchange intervals

5. Touch the button (D).
6. Set the value by turning the rotary knob and push to confirm.

The settings are effective immediately.

No display of exchange intervals

- Touch the button (D). Set **Off** by turning the rotary knob and push to confirm.

Additional information

Babylog VN500 displays the remaining period of use for the accessories on the **Start/Standby > Accessory status** page.

Sterilization of the expiratory valve or inspiratory valve may gradually impair the operation of RFID transmission. This may mean that Infinity ID breathing circuit functions may not work or may no longer work reliably. The period of use for the Infinity ID accessories is displayed with ---.
System settings

The following system settings can be configured:

- Country
- Units
- Interface (interfaces)
- Supply units (supply units)
- Service

The customized settings are immediately effective.

1. Touch the **System setup...** button.
2. Touch the **System** tab.

Babylog VN500 displays the following configurable settings in an overview:

- Language, date and time
- Units for measured values and settings
- Network and serial interfaces
- GS500
- Service information

Selecting country-specific settings

Prerequisite: The **System** page (A) is opened.

1. Touch the **Country** tab (B).

Selecting the screen text language

Babylog VN500 is factory set to the customer's own language. The current language is displayed in the field (C).

Selecting a different language:

2. Touch the ▼ button (D).

Babylog VN500 opens the selection list containing the available languages.

3. Select the language with the rotary knob and push to confirm.

Setting the date and time

Babylog VN500 does not change over automatically between daylight saving time and standard time. The user must change the time manually. Otherwise the times will be incorrect on the screen and for saved values and actions (e.g., in the logbook).

Changing the system time changes the time displayed in trends, logbook, alarm history, maneuver measured values and reference loops. The data saved up to the change is displayed with the system time up till then.

1. Touch the appropriate button:
   - Day (E)
   - Month (F)
   - Year (G)
   - Hours (H)
   - Minutes (I)

   The order of the buttons (E) and (F) varies depending on language.

2. Set the value by turning the rotary knob and push to confirm.

3. After completing all the settings for the date and time, touch the **Apply** button (J).
Configuration

Entering the height above sea level

The ambient pressure is considered in the calculation of measured values. The ambient pressure sensor is checked for plausibility using the entered height above sea level. Incorrect entries can mean that the ambient pressure sensor is recognized as incorrect.

1. Touch the button (K).

2. Set the height by turning the rotary knob and push to confirm.

Configuring units

Prerequisite: The System page (A) is opened.

1. Touch the Units tab (B).

The units for the following parameters can be selected.

- **Airway pressure** (C) in mbar or cmH2O
- **Height** (D) in m, cm or feet, inch
- **CO2** (E) in Vol% or mmHg or kPa

The unit selected for the CO2 measured value is adopted for the alarm limit.

2. Touch the relevant button for the unit.
Configuring interfaces

The communication settings can be configured to enable connection to a network and data exchange with other devices.

LAN

Use of LAN ports is exclusively permitted for service purposes. Parameters must be set for connection to a network.

Prerequisite: The System page (A) is opened.

1 Touch the Interface tab (B).

The LAN page (C) appears by default. The settings are displayed. DHCP (D) must be deactivated in order to change the settings.

Deactivating DHCP (D):

1 Touch the Off button.

2 Touch the relevant button for the network parameters:
   – IP address (E)
   – Subnet mask (F)
   – Gateway (G)

3 Enter the login details using the rotary knob and confirm.

4 Touch the Install button (H).

Activating DHCP (D):

5 Touch the On button.

Serial interfaces

The serial interfaces (COM 1, COM 2 and COM 3) are used for data exchange with MEDIBUS-capable display devices, e.g., patient monitor or patient data management system.

Prerequisite: The System page (A) is opened.

1 Touch the Interface tab (B).

2 Touch the COM tab (C).

The settings for COM 1, COM 2, and COM 3 are displayed. MEDIBUS or MEDIBUS.X can be selected for the Protocol parameter.

3 Touch the relevant button for the interface parameters:
   – Protocol (D)
   – Baud rate (E)
   – Parity (F)
   – Stop bit (G)

4 Select the setting with the rotary knob and push to confirm.
**External screen**

If a second screen is connected to Infinity C500, the user has to define whether the screen is analog or digital.

Prerequisite: The *System* page (A) is opened.

1. Touch the *Interface* tab (B).
2. Touch the *External display* tab (C).
3. Touch the *Digital* (D) or *Analog* (E) button.

**Additional information**

The serial interface connectors are located on the rear of Infinity C500.

---

**Configuring supply units**

**Functionality of the GS500 gas supply unit**

The functionality of the gas supply unit can be deactivated if Babylog VN500 is equipped with a gas supply unit that is currently not supposed to be used.

Prerequisite: The *System* page (A) is opened.

- Touch the *Supply units* tab (B).
- Touch the *Off* button (C).

The gas supply unit is no longer available. In the device check, the system does not display the *Gas supply unit* test step.

Activating the functionality of the gas supply unit:

1. Touch the *On* button (D).

Babylog VN500 displays in the message field (E) that the device check has to be carried out.

2. Perform device check.

**Additional information**

For information on using the gas supply unit, see "GS500 gas supply unit" on page 110.
Service dialog

The service dialog is password-protected and reserved for DrägerService or experts.

For further information on Remote Service, see chapter "Remote Service" on page 249.
This page has been left blank intentionally.
The alarm messages are displayed in the message field of the header bar in hierarchical order. See "Display of alarms" on page 124.

In order to classify the alarms within an alarm category, internal priority numbers are given after the exclamation marks in the table below. The most critical alarm is awarded the number 255. The priority of the alarm decreases the lower the number is.

In the following table, the alarm messages are listed in alphabetical order. If an alarm occurs, the table helps to identify causes and remedies. The different causes and remedies should be worked through in the order listed until the alarm has been resolved.

The acknowledgeable alarm messages can be found in the chapter "Alarm – Cause – Remedy". For alarm messages that can be acknowledged, the "Remedy" column in the table contains information stating that the alarm message can be acknowledged by pressing the **ALARM RESET** button and confirming with the rotary knob.

The following alarm messages that can be acknowledged are not listed:

- **Suction maneuver overused?**
- **PEEP high (!!)**
### Alarm – Cause – Remedy

<table>
<thead>
<tr>
<th>Alarm priority</th>
<th>Alarm message</th>
<th>Cause</th>
<th>Remedy</th>
</tr>
</thead>
<tbody>
<tr>
<td>! 060</td>
<td>Accessory ID detection failed</td>
<td>Accessory ID detection malfunction.</td>
<td>Ventilation can be started without ID functions.</td>
</tr>
<tr>
<td></td>
<td></td>
<td></td>
<td>Call DrägerService.</td>
</tr>
<tr>
<td>!! 110</td>
<td>Air and O2 supply insufficient</td>
<td>Air and O2 supply insufficient to deliver required flow and pressure for HFO.</td>
<td>Check connections to Air and O2 supply.</td>
</tr>
<tr>
<td></td>
<td></td>
<td></td>
<td>Make sure supply pressures are greater than 3 bar (43.5 psi).</td>
</tr>
<tr>
<td></td>
<td></td>
<td></td>
<td>Downgrade alarm priority by touching &quot;ALARM RESET&quot; button and confirm with rotary knob.</td>
</tr>
<tr>
<td>!! 110</td>
<td>Air pressure regulation failed</td>
<td>Internal Air supply pressure too high.</td>
<td>No HFO possible.</td>
</tr>
<tr>
<td></td>
<td></td>
<td></td>
<td>Call DrägerService.</td>
</tr>
<tr>
<td>!! 110</td>
<td>Air supply insufficient</td>
<td>Air supply insufficient to deliver required flow and pressure for HFO.</td>
<td>Check connection to Air supply.</td>
</tr>
<tr>
<td></td>
<td></td>
<td></td>
<td>Make sure that the supply pressure is greater than 3 bar (43.5 psi).</td>
</tr>
<tr>
<td></td>
<td></td>
<td></td>
<td>Acknowledge message by touching &quot;ALARM RESET&quot; and confirm.</td>
</tr>
<tr>
<td></td>
<td></td>
<td></td>
<td>Use external O2 monitoring.</td>
</tr>
</tbody>
</table>
### Alarm – Cause – Remedy

<table>
<thead>
<tr>
<th>Alarm priority</th>
<th>Alarm message</th>
<th>Cause</th>
<th>Remedy</th>
</tr>
</thead>
<tbody>
<tr>
<td>! 100</td>
<td>Air supply low, GS500 active</td>
<td>Air supply insufficient to deliver the required flow and pressure. Air is supplied by the gas supply unit GS500. Air supply is not required when FiO2 = 100 Vol%.</td>
<td>Check connection to Air supply. Make sure supply pressure is greater than 3 bar (43.5 psi). Consider readjusting ventilation settings. Remove connection to Air supply if alarm condition persists (to avoid reverse flow into the Air supply).</td>
</tr>
<tr>
<td></td>
<td>Central Air supply insufficient. Gas delivery system is supplied with Air delivered by GS500.</td>
<td></td>
<td>Check connection to central air supply and to gas supply unit GS500. Make sure that the supply pressure is greater than 3 bar (43.5 psi). Adjust ventilation settings, if necessary.</td>
</tr>
<tr>
<td>!!! 190</td>
<td>Airway obstructed?</td>
<td>The ventilation unit applies only a very small volume with each mechanical breath. The tube or mask could be blocked. Patient breathes against the mechanical breaths during pressure-controlled ventilation.</td>
<td>Check patient condition. Check tube or mask. Check patient condition. Check ventilation settings.</td>
</tr>
<tr>
<td>Alarm priority</td>
<td>Alarm message</td>
<td>Cause</td>
<td>Remedy</td>
</tr>
<tr>
<td>----------------</td>
<td>---------------</td>
<td>-------</td>
<td>--------</td>
</tr>
</tbody>
</table>
| !!! 205        | Airway pressure high | Breathing hose kinked. | Check breathing circuit.  
Check tube or mask.  
The upper alarm limit for the airway pressure has been exceeded. The patient is breathing against the ventilation unit or coughing.  
Check patient condition.  
Check ventilation settings.  
Adjust alarm limit if necessary. |
| !!! 200        | Airway pressure low | Leakage or disconnection. | Check breathing circuit for tight connections.  
Check whether the expiratory valve is properly engaged.  
Make sure that the tube or mask is connected correctly. |
| !!! 140        | Airway pressure negative | Airway pressure has fallen below –10 mbar (–10 cmH2O). | Disconnect tube for suctioning.  
Check patient condition.  
Check ventilation settings.  
The breathing hose is connected to the expiratory valve during O2 therapy.  
Connect breathing hose to the inspiratory valve. |
| !!! 140        | Airway pressure negative (averaged) | Average airway pressure has fallen below –2 mbar (–2 cmH2O). | Disconnect tube for suctioning.  
Check patient condition.  
Check ventilation settings. |
| !! 140         | Airway pressure negative (averaged) | Average airway pressure has fallen below –2 mbar (–2 cmH2O). | Disconnect tube for suctioning.  
Check patient condition.  
Check ventilation settings. |
<table>
<thead>
<tr>
<th>Alarm priority</th>
<th>Alarm message</th>
<th>Cause</th>
<th>Remedy</th>
</tr>
</thead>
<tbody>
<tr>
<td>! 200</td>
<td>Alarm limit not confirmed</td>
<td>One or more alarm limits have been changed but not confirmed.</td>
<td>If necessary, change these alarm limits and confirm the change with the rotary knob.</td>
</tr>
<tr>
<td>! 120</td>
<td>Alarm system failure</td>
<td>Failure of primary alarm speaker. In case of an alarm situation, the auxiliary acoustical alarm will sound.</td>
<td>To continue ventilation with this device, continuously monitor the device functions. Call DrägerService.</td>
</tr>
<tr>
<td>!! 100</td>
<td>Ambient pressure sensor?</td>
<td>Altitude setting deviates too much from measured ambient pressure.</td>
<td>Check altitude setting and adjust if necessary. If the setting has been adjusted, the device check must be repeated. Check ambient pressure sensor failure.</td>
</tr>
<tr>
<td>!!! 181</td>
<td>Apnea</td>
<td>The patient has stopped breathing.</td>
<td>Check patient condition. Apply controlled ventilation if necessary. Check patient condition. Check breathing circuit. Check tube or mask. Flow sensor is not calibrated or faulty.</td>
</tr>
<tr>
<td>Alarm priority</td>
<td>Alarm message</td>
<td>Cause</td>
<td>Remedy</td>
</tr>
<tr>
<td>----------------</td>
<td>---------------</td>
<td>-------</td>
<td>--------</td>
</tr>
<tr>
<td>!! 230</td>
<td>Apnea Ventilation</td>
<td>Due to detected apnea, the ventilation unit has automatically switched to Apnea Ventilation.</td>
<td>Check patient condition. Check tube or mask. Check ventilation settings and patient condition. Return to the original ventilation mode by touching the &quot;Apn. Vent. reset&quot; button and confirm with rotary knob.</td>
</tr>
<tr>
<td>! 020</td>
<td>Application already installed</td>
<td>Application is already installed.</td>
<td>Acknowledge message by touching &quot;ALARM RESET&quot; button and confirm with rotary knob.</td>
</tr>
<tr>
<td>! 020</td>
<td>Application transfer failed</td>
<td>Invalid application.</td>
<td>Acknowledge message by touching &quot;ALARM RESET&quot; button and confirm with rotary knob. Call DrägerService.</td>
</tr>
<tr>
<td>!! 050</td>
<td>&quot;Audio paused&quot; key used too often</td>
<td>The &quot;Audio paused&quot; key is either faulty or was pressed more than 80 times per hour.</td>
<td>The function of the &quot;Audio paused&quot; key is not available while the defect exists. If the defect cannot be remedied, call DrägerService.</td>
</tr>
<tr>
<td>!! 050</td>
<td>&quot;Audio paused&quot; overused or stuck</td>
<td>The &quot;Audio paused&quot; button is either stuck or faulty or was pressed for more than 6 seconds.</td>
<td>Ventilation functions are not affected. Do not press the &quot;Audio paused&quot; button longer than 6 seconds. If the error persists, call DrägerService.</td>
</tr>
</tbody>
</table>
### Alarm – Cause – Remedy

<table>
<thead>
<tr>
<th>Alarm priority</th>
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<th>Remedy</th>
</tr>
</thead>
</table>
| !! 120        | Auxiliary acoustical alarm failure | Failure of auxiliary alarm speaker.  
- In case of mains failure and discharged battery, there is no power failure alarm.  
- In case of faulty primary alarm speaker, there is no acoustical alarm at all. | To continue ventilation with this device, continuously monitor the device functions.  
Downgrade alarm priority by touching "ALARM RESET" button and confirm with rotary knob.  
Call DrägerService. |
| !!! 160       | Battery activated | The ventilation unit is powered by the battery as there is no mains power supply. | Connect device to the mains power supply. |
| !! 200        | Battery activated | The ventilation unit is powered by the battery as there is no mains power supply. | Connect device to the mains power supply. |
| ! 201         | Battery activated | The ventilation unit is powered by the battery as there is no mains power supply. | Connect device to the mains power supply. |
| !! 127        | Battery charging deferred | Battery charging is deferred to prevent battery overheating.  
The device can be used normally. | Battery charging continues automatically and is indicated by a flashing segment in the battery symbol. |
<p>| ! 100         | Battery check in progress | The battery check has been started. | Wait until the battery check is completed. In the event of mains power supply failure, battery operation is limited. |
| ! 100         | Battery check recommended | The interval for the battery check has been exceeded. | Perform the battery check. |
| !!! 254       | Battery discharged | The remaining calculated operating time of the battery is less than 5 minutes. | Connect device immediately to the mains power supply. |</p>
<table>
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<tr>
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<th>Remedy</th>
</tr>
</thead>
<tbody>
<tr>
<td>!!</td>
<td>120 Battery failure</td>
<td>Battery operation is not available in the event of mains power supply failure.</td>
<td>To continue ventilation with this device, continuously monitor the device functions. Call DrägerService.</td>
</tr>
<tr>
<td>!!!</td>
<td>250 Battery low</td>
<td>The remaining calculated operating time of the battery is less than 10 minutes.</td>
<td>Connect device to the mains power supply.</td>
</tr>
<tr>
<td>!!</td>
<td>251 Battery low</td>
<td>The remaining calculated operating time of the battery is less than 10 minutes.</td>
<td>Connect device to the mains power supply.</td>
</tr>
<tr>
<td>!!</td>
<td>105 Breath. circ. does not fit to patient category</td>
<td>Connected breathing circuit does not fit to selected patient category.</td>
<td>Use suitable breathing circuit or select correct patient category.</td>
</tr>
<tr>
<td>!!</td>
<td>100 Breathing circuit does not match config.</td>
<td>Breathing circuit has been exchanged. The new breathing circuit does not match the one that was used before.</td>
<td>Check breathing circuit. Acknowledge message by pressing &quot;ALARM RESET&quot; and confirm.</td>
</tr>
<tr>
<td>!</td>
<td>060 Breathing circuit ID invalid</td>
<td>Accessory ID detection failed. No automatic adjustment of breathing circuit properties.</td>
<td>Replace ID Breathing Circuit or perform breathing circuit check. Acknowledge message by touching &quot;ALARM RESET&quot; button and confirm with rotary knob.</td>
</tr>
<tr>
<td></td>
<td></td>
<td>Accessory ID detection failed. Breathing circuit exchange interval cannot be monitored.</td>
<td>Replace Infinity ID Breathing Circuit or acknowledge message by touching &quot;ALARM RESET&quot; button and confirm with rotary knob.</td>
</tr>
</tbody>
</table>
## Alarm – Cause – Remedy

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<tbody>
<tr>
<td>!! 205</td>
<td>Breathing hose kinked</td>
<td>The pressure at the inspiratory port is greater than 30 mbar (30 cmH2O), e.g., due to a kinked or blocked hose, or a blocked mask.</td>
<td>Check breathing circuit.</td>
</tr>
<tr>
<td></td>
<td></td>
<td></td>
<td>Check mask.</td>
</tr>
<tr>
<td>!! 105</td>
<td>Breathing hoses interchanged</td>
<td>Inspiratory and expiratory limbs of the breathing circuit are connected reversely to the ventilation unit.</td>
<td>Connect inspiratory and expiratory limbs of the breathing circuit correctly.</td>
</tr>
</tbody>
</table>
| !!! 240        | Calibration of gas delivery system required | Technical malfunction detected during operation.  
Calibration of gas delivery system failed.  
Recalibration necessary.  
Ventilation not possible. | Disconnect patient from the device and continue ventilation without delay using another independent ventilator.  
Perform device check. |
<table>
<thead>
<tr>
<th>Alarm priority</th>
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<th>Remedy</th>
</tr>
</thead>
<tbody>
<tr>
<td>! 012</td>
<td>Calibration of gas delivery system required</td>
<td>Technical malfunction detected in standby mode.</td>
<td>Perform device check.</td>
</tr>
<tr>
<td></td>
<td></td>
<td>Calibration of gas delivery system failed.</td>
<td>Do not start with ventilation before device check is performed:</td>
</tr>
<tr>
<td></td>
<td></td>
<td>Recalibration necessary.</td>
<td>Ventilation will not be possible.</td>
</tr>
<tr>
<td></td>
<td>Technical malfunction detected in standby mode.</td>
<td></td>
<td>Perform device check.</td>
</tr>
<tr>
<td></td>
<td></td>
<td>Calibration of gas delivery system is due.</td>
<td></td>
</tr>
<tr>
<td></td>
<td></td>
<td>Accuracy of gas delivery system could be impaired.</td>
<td></td>
</tr>
<tr>
<td></td>
<td></td>
<td>Recalibration necessary.</td>
<td></td>
</tr>
<tr>
<td></td>
<td>Technical malfunction detected in standby mode.</td>
<td></td>
<td>Perform device check.</td>
</tr>
<tr>
<td></td>
<td></td>
<td>Calibration of gas delivery system failed.</td>
<td>Do not start with ventilation before device check is performed:</td>
</tr>
<tr>
<td></td>
<td></td>
<td></td>
<td>Ventilation will not be possible.</td>
</tr>
<tr>
<td></td>
<td></td>
<td></td>
<td>If alarm cannot be resolved by performing device check, call DrägerService.</td>
</tr>
</tbody>
</table>
### Alarm – Cause – Remedy

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</tr>
</thead>
<tbody>
<tr>
<td>!!! 228</td>
<td>Calibration of neo. flow sensor required</td>
<td>Calibration data is corrupted.</td>
<td>Patient category “Neonates”:&lt;br&gt;- Calibrate neonatal flow sensor.&lt;br&gt;- If calibration was not successful, deactivate neonatal flow monitoring and use external flow monitoring.&lt;br&gt;- Call DrägerService.</td>
</tr>
<tr>
<td>!! 115</td>
<td>Calibration of neo. flow sensor required</td>
<td>After switching on the ventilation unit, the neonatal flow sensor needs to be calibrated.</td>
<td>Calibrate neonatal flow sensor.</td>
</tr>
<tr>
<td>!!! 228</td>
<td>Calibration of neonatal flow sensor failed</td>
<td>Calibration of neonatal flow sensor failed.</td>
<td>Calibrate neonatal flow sensor.&lt;br&gt;- Seal neonatal flow sensor properly during calibration.</td>
</tr>
<tr>
<td></td>
<td>Neonatal flow sensor malfunction.</td>
<td></td>
<td>Replace neonatal flow sensor or sensor insert and calibrate the new sensor.</td>
</tr>
</tbody>
</table>
## Alarm – Cause – Remedy

<table>
<thead>
<tr>
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<th>Cause</th>
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</tr>
</thead>
<tbody>
<tr>
<td>!! 100</td>
<td>Check CO2 cuvette</td>
<td>The selected type of CO2 cuvette is not correct.</td>
<td>Select the correct type of CO2 cuvette.</td>
</tr>
<tr>
<td></td>
<td></td>
<td>CO2 cuvette or sensor soiled.</td>
<td>Clean the CO2 cuvette or sensor.</td>
</tr>
<tr>
<td></td>
<td></td>
<td>CO2 sensor drift.</td>
<td>Perform zero calibration.</td>
</tr>
<tr>
<td></td>
<td></td>
<td>Inspiratory CO2 concentration high.</td>
<td>Check ventilation settings. Check patient condition.</td>
</tr>
<tr>
<td>!! 140</td>
<td>Check settings</td>
<td>Loss of stored data was detected.</td>
<td>Check all settings and adjust if necessary. Acknowledge message by</td>
</tr>
<tr>
<td></td>
<td></td>
<td></td>
<td>touching &quot;ALARM RESET&quot; button and confirm with rotary knob.</td>
</tr>
<tr>
<td>!! 252</td>
<td>Check ventilation settings</td>
<td>Due to data loss, the device uses previous</td>
<td>Check all therapy settings and adjust them if necessary. Acknowledge</td>
</tr>
<tr>
<td></td>
<td></td>
<td>settings.</td>
<td>message by touching &quot;ALARM RESET&quot; button and confirm with rotary knob.</td>
</tr>
<tr>
<td>!! 252</td>
<td>Check ventilation settings</td>
<td>While adjusting ventilation settings or alarm</td>
<td>The device may apply default settings. Check ventilation settings and</td>
</tr>
<tr>
<td></td>
<td></td>
<td>limits, a power interruption occurred.</td>
<td>alarm limits. Acknowledge message by touching &quot;ALARM RESET&quot; button and</td>
</tr>
<tr>
<td></td>
<td></td>
<td></td>
<td>confirm with rotary knob.</td>
</tr>
<tr>
<td></td>
<td></td>
<td>Data loss.</td>
<td>The device may apply default settings. Check ventilation settings and</td>
</tr>
<tr>
<td></td>
<td></td>
<td></td>
<td>alarm limits. Acknowledge message by touching &quot;ALARM RESET&quot; button and</td>
</tr>
<tr>
<td></td>
<td></td>
<td></td>
<td>confirm with rotary knob.</td>
</tr>
<tr>
<td>Alarm priority</td>
<td>Alarm message</td>
<td>Cause</td>
<td>Remedy</td>
</tr>
<tr>
<td>---------------</td>
<td>---------------</td>
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</tr>
<tr>
<td>!!! 144</td>
<td>Clean CO2 cuvette</td>
<td>Cuvette or sensor window is soiled, e.g. with deposits due to nebulization.</td>
<td>Use clean cuvette and/or clean CO2 sensor.</td>
</tr>
<tr>
<td>!!! 145</td>
<td>CO2 measurement failed</td>
<td>CO2 sensor faulty.</td>
<td>Replace faulty CO2 sensor.</td>
</tr>
<tr>
<td></td>
<td></td>
<td>CO2 measurement incorrect.</td>
<td>Use external CO2 monitoring and deactivate integrated CO2 monitoring. Call DrägerService.</td>
</tr>
<tr>
<td>!!! 146</td>
<td>CO2 sensor?</td>
<td>Plug of CO2 sensor was removed during operation.</td>
<td>Reinsert plug.</td>
</tr>
<tr>
<td></td>
<td></td>
<td>CO2 sensor not positioned on cuvette.</td>
<td>Place CO2 sensor on cuvette.</td>
</tr>
<tr>
<td></td>
<td></td>
<td>CO2 sensor faulty.</td>
<td>Replace faulty CO2 sensor.</td>
</tr>
<tr>
<td>!!! 142</td>
<td>CO2 zero calibration?</td>
<td>Zero point of the CO2 sensor is outside of the tolerance range.</td>
<td>Perform zero calibration.</td>
</tr>
<tr>
<td></td>
<td></td>
<td>Cuvette or sensor window is soiled, e.g. with deposits due to nebulization.</td>
<td>Use clean cuvette and/or clean CO2 sensor.</td>
</tr>
<tr>
<td>! 100</td>
<td>Cockpit restarted</td>
<td>Internal communication error caused restart of the cockpit.</td>
<td>Check all therapy settings and adjust them if necessary. Acknowledge message by touching &quot;ALARM RESET&quot; button and confirm with rotary knob.</td>
</tr>
<tr>
<td>! 100</td>
<td>Continuous nebulization activated</td>
<td>Continuous nebulization was activated by the user.</td>
<td>To end continuous nebulization, press the &quot;Cancel&quot; button if required.</td>
</tr>
</tbody>
</table>
## Alarm – Cause – Remedy

<table>
<thead>
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</tr>
</thead>
<tbody>
<tr>
<td>!!! 252</td>
<td>Data loss</td>
<td>Loss of stored data was detected.</td>
<td>To continue ventilation with this device, continuously monitor the device functions. Downgrade alarm priority by touching &quot;ALARM RESET&quot; button and confirm with rotary knob. Call DrägerService.</td>
</tr>
<tr>
<td>!! 252</td>
<td>Data loss</td>
<td>Loss of stored data was detected.</td>
<td>To continue ventilation with this device, continuously monitor the device functions. Downgrade alarm priority by touching &quot;ALARM RESET&quot; button and confirm with rotary knob. Call DrägerService.</td>
</tr>
<tr>
<td>!! 240</td>
<td>Device check failed</td>
<td>A safety-related failure was detected during device check.</td>
<td>Do not use this device for ventilation therapy. Call DrägerService. Check assembly and position of expiratory valve. Replace expiratory valve if required. Do not use this device for ventilation therapy unless the device check was repeated successfully.</td>
</tr>
<tr>
<td>! 100</td>
<td>Device check incomplete</td>
<td>Device check not completely performed or partially unsuccessful.</td>
<td>Perform device check. Acknowledge message by touching &quot;ALARM RESET&quot; button and confirm with rotary knob.</td>
</tr>
</tbody>
</table>
### Alarm – Cause – Remedy

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<tbody>
<tr>
<td>!!! 253</td>
<td>Device failure</td>
<td>A system failure was detected.</td>
<td>Disconnect patient from the device and continue ventilation without delay using another independent ventilator. Call DrägerService.</td>
</tr>
<tr>
<td>!!! 253</td>
<td>Device failure</td>
<td>Due to missing measurements, ventilation is not possible anymore.</td>
<td>Immediately disconnect the patient from the device and continue ventilation without delay using another independent ventilator. Call DrägerService.</td>
</tr>
<tr>
<td>!!! 253</td>
<td>Device failure (1)</td>
<td>Internal safety system failure.</td>
<td>Disconnect patient from the device and continue ventilation without delay using another independent ventilator. Call DrägerService.</td>
</tr>
<tr>
<td>!!! 253</td>
<td>Device failure (10)</td>
<td>A failure was detected by the safety software system.</td>
<td>Disconnect patient from the device and continue ventilation without delay using another independent ventilator. Call DrägerService.</td>
</tr>
<tr>
<td>!!! 253</td>
<td>Device failure (11)</td>
<td>A failure was detected during the start-up phase.</td>
<td>Disconnect patient from the device and continue ventilation without delay using another independent ventilator. Call DrägerService.</td>
</tr>
<tr>
<td>!!! 253</td>
<td>Device failure (12)</td>
<td>A system failure was detected.</td>
<td>Disconnect patient from the device and continue ventilation without delay using another independent ventilator. Call DrägerService.</td>
</tr>
</tbody>
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### Alarm – Cause – Remedy

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<tbody>
<tr>
<td>!! 090</td>
<td>Device failure (13)</td>
<td>The broken wire detection for the flow sensor is faulty.</td>
<td>Ventilation functions are not affected. Call DrägerService.</td>
</tr>
<tr>
<td>!!! 253</td>
<td>Device failure (2)</td>
<td>Internal safety system failure.</td>
<td>Do not use this device for ventilation therapy. Call DrägerService.</td>
</tr>
<tr>
<td>!!! 253</td>
<td>Device failure (3)</td>
<td>Internal communication failure.</td>
<td>Disconnect patient from the device and continue ventilation without delay using another independent ventilator. Call DrägerService.</td>
</tr>
<tr>
<td>!!! 253</td>
<td>Device failure (4)</td>
<td>Defective system data storage media detected.</td>
<td>Disconnect patient from the device and continue ventilation without delay using another independent ventilator. Switch off the device. Call DrägerService.</td>
</tr>
<tr>
<td>!!! 253</td>
<td>Device failure (5)</td>
<td>Gas delivery system faulty.</td>
<td>Disconnect patient from the device and continue ventilation without delay using another independent ventilator. Call DrägerService.</td>
</tr>
<tr>
<td>!!! 253</td>
<td>Device failure (6)</td>
<td>Gas delivery system faulty.</td>
<td>Disconnect patient from the device and continue ventilation without delay using another independent ventilator. Call DrägerService.</td>
</tr>
<tr>
<td>!!! 253</td>
<td>Device failure (7)</td>
<td>Gas delivery system faulty.</td>
<td>Disconnect patient from the device and continue ventilation without delay using another independent ventilator. Call DrägerService.</td>
</tr>
<tr>
<td>Alarm priority</td>
<td>Alarm message</td>
<td>Cause</td>
<td>Remedy</td>
</tr>
<tr>
<td>---------------</td>
<td>--------------------------------------</td>
<td>----------------------------------------------------------------------</td>
<td>------------------------------------------------------------------------</td>
</tr>
<tr>
<td>!!! 253</td>
<td>Device failure (8)</td>
<td>Test alarm which should only be triggered during maintenance.</td>
<td>Disconnect patient from the device and continue ventilation without delay using another independent ventilator. Call DrägerService.</td>
</tr>
<tr>
<td>!! 100</td>
<td>Device failure (9)</td>
<td>No mass storage device found.</td>
<td>Acknowledge message by touching &quot;ALARM RESET&quot; button and confirm with rotary knob. Call DrägerService.</td>
</tr>
<tr>
<td>!!! 200</td>
<td>Device temperature high</td>
<td>The internal device temperature is too high.</td>
<td>Disconnect patient from the device and continue ventilation without delay using another independent ventilator. Switch off the device. Call DrägerService.</td>
</tr>
<tr>
<td>!! 141</td>
<td>Device temperature measurement failed</td>
<td>Failure of the internal breathing-gas temperature measurement.</td>
<td>To continue ventilation with this device, use external breathing gas temperature monitoring. Call DrägerService.</td>
</tr>
<tr>
<td></td>
<td></td>
<td>In case of a too high breathing-gas temperature, there is no alarm.</td>
<td></td>
</tr>
<tr>
<td></td>
<td></td>
<td>Failure of the internal temperature measurement.</td>
<td>To continue ventilation with this device, continuously monitor the device functions. Call DrägerService.</td>
</tr>
<tr>
<td></td>
<td></td>
<td>No alarm in case of a too high device temperature.</td>
<td></td>
</tr>
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<tr>
<td>!!! 200</td>
<td>Disconnection?</td>
<td>Leakage or disconnection.</td>
<td>Check breathing circuit for tight connections.</td>
</tr>
<tr>
<td></td>
<td></td>
<td></td>
<td>Check whether the expiratory valve is properly engaged.</td>
</tr>
<tr>
<td></td>
<td></td>
<td></td>
<td>Make sure that the tube or mask is connected correctly.</td>
</tr>
<tr>
<td>!!! 138</td>
<td>etCO₂ high</td>
<td>Upper alarm limit for end-expiratory CO₂ concentration has been exceeded.</td>
<td>Check patient condition.</td>
</tr>
<tr>
<td></td>
<td></td>
<td></td>
<td>Check ventilation settings.</td>
</tr>
<tr>
<td></td>
<td></td>
<td></td>
<td>Adjust alarm limit if necessary.</td>
</tr>
<tr>
<td></td>
<td></td>
<td></td>
<td>Perform CO₂ zero calibration if necessary.</td>
</tr>
<tr>
<td></td>
<td></td>
<td></td>
<td>Check whether the cuvette windows are soiled.</td>
</tr>
<tr>
<td>!!! 138</td>
<td>etCO₂ low</td>
<td>Lower alarm limit for end-expiratory CO₂ concentration has been exceeded.</td>
<td>Check patient condition.</td>
</tr>
<tr>
<td></td>
<td></td>
<td></td>
<td>Check ventilation settings.</td>
</tr>
<tr>
<td></td>
<td></td>
<td></td>
<td>Adjust alarm limit if necessary.</td>
</tr>
<tr>
<td></td>
<td></td>
<td></td>
<td>Perform CO₂ zero calibration if necessary.</td>
</tr>
<tr>
<td></td>
<td></td>
<td></td>
<td>Check whether the cuvette windows are soiled.</td>
</tr>
<tr>
<td>!!! 220</td>
<td>Expiratory valve faulty</td>
<td>Expiratory valve is not properly connected to the socket.</td>
<td>Insert expiratory valve correctly.</td>
</tr>
<tr>
<td></td>
<td></td>
<td></td>
<td>Expiratory valve faulty. Replace expiratory valve.</td>
</tr>
<tr>
<td>!!! 105</td>
<td>Expiratory valve incompatible</td>
<td>Incompatible expiratory valve connected to the socket.</td>
<td>Replace expiratory valve.</td>
</tr>
</tbody>
</table>

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Instructions for use Infinity Acute Care System – Babylog VN500 SW 2.n
<table>
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<th>Alarm priority</th>
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<th>Remedy</th>
</tr>
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<tbody>
<tr>
<td>!! 100</td>
<td>Expiratory valve incompatible</td>
<td>Incompatible expiratory valve connected to the socket.</td>
<td>Replace expiratory valve.</td>
</tr>
<tr>
<td>!!! 130</td>
<td>FiO2 high</td>
<td>O2 sensor is not calibrated.</td>
<td>Calibrate O2 sensor.</td>
</tr>
<tr>
<td></td>
<td></td>
<td>Mixer function faulty.</td>
<td>Call DrägerService.</td>
</tr>
<tr>
<td>!!! 130</td>
<td>FiO2 low</td>
<td>O2 sensor is not calibrated.</td>
<td>Calibrate O2 sensor.</td>
</tr>
<tr>
<td></td>
<td></td>
<td>Mixer function faulty.</td>
<td>Call DrägerService.</td>
</tr>
<tr>
<td>!! 100</td>
<td>Flow measurement inaccurate</td>
<td>Flow sensor is not calibrated or faulty.</td>
<td>Calibrate flow sensor and replace it if necessary.</td>
</tr>
<tr>
<td></td>
<td></td>
<td>Water in flow sensor.</td>
<td>Drain water trap of breathing circuit. Dry flow sensor.</td>
</tr>
<tr>
<td></td>
<td></td>
<td>Flow measurement is not reliable. Expiratory minute volume exceeds</td>
<td>To continue ventilation with this device, use external flow monitoring and deactivate integrated flow monitoring. This could impair the quality of ventilation. Call DrägerService.</td>
</tr>
<tr>
<td></td>
<td></td>
<td>minute volume delivered by the ventilation unit.</td>
<td></td>
</tr>
<tr>
<td>!!! 110</td>
<td>GS500 communication failure</td>
<td>Communication to gas supply unit GS500 lost.</td>
<td>Check communication connection to gas supply unit GS500.</td>
</tr>
<tr>
<td></td>
<td></td>
<td>Acknowledge message by pressing &quot;ALARM RESET&quot; and confirm.</td>
<td></td>
</tr>
<tr>
<td></td>
<td></td>
<td>Call DrägerService.</td>
<td></td>
</tr>
<tr>
<td>Alarm priority</td>
<td>Alarm message</td>
<td>Cause</td>
<td>Remedy</td>
</tr>
<tr>
<td>---------------</td>
<td>--------------------------------</td>
<td>----------------------------------------------------------------------</td>
<td>------------------------------------------------------------------------</td>
</tr>
<tr>
<td>! 110</td>
<td>GS500 communication failure</td>
<td>Communication to gas supply unit GS500 lost.</td>
<td>Check communication connection to gas supply unit GS500. Acknowledge message by pressing &quot;ALARM RESET&quot; and confirm. Call DrägerService.</td>
</tr>
<tr>
<td>!!! 100</td>
<td>GS500 failure</td>
<td>Air supply insufficient to deliver required flow and pressure.</td>
<td>Check connection to gas supply unit GS500. If this condition persists, call DrägerService.</td>
</tr>
<tr>
<td>!!! 110</td>
<td>GS500 internal failure</td>
<td>Gas supply unit GS500 has reported a failure.</td>
<td>Shut down ventilation unit. Switch toggle switch to &quot;Off&quot; to disconnect ventilation unit from power supply. Switch toggle switch to &quot;On&quot; and restart ventilation unit. If this condition persists, call DrägerService.</td>
</tr>
<tr>
<td>!! 110</td>
<td>GS500 internal failure</td>
<td>Gas supply unit GS500 has reported a failure.</td>
<td>Shut down ventilation unit. Switch toggle switch to &quot;Off&quot; to disconnect ventilation unit from power supply. Switch toggle switch to &quot;On&quot; and restart ventilation unit. If this condition persists, call DrägerService.</td>
</tr>
<tr>
<td>!!! 110</td>
<td>GS500 temperature too high</td>
<td>Gas supply unit GS500 temperature is too high.</td>
<td>Shut down ventilation unit. Switch toggle switch to &quot;Off&quot;. Call DrägerService.</td>
</tr>
</tbody>
</table>
### Alarm – Cause – Remedy

<table>
<thead>
<tr>
<th>Alarm priority</th>
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</tr>
</thead>
<tbody>
<tr>
<td>!!! 249</td>
<td>HFO not possible</td>
<td>Gas supply unable to deliver required flow and pressure for active expiration during HFO.</td>
<td>Check connections to Air and O₂ supply. Make sure supply pressures are greater than 3 bar (43.5 psi). If supply pressures are correct: Call DrägerService. When the gas supply unit is active, change to a different ventilation mode.</td>
</tr>
<tr>
<td>! 060</td>
<td>ID tag of expiratory valve faulty</td>
<td>Accessory ID detection failed. Expiratory valve exchange interval cannot be monitored.</td>
<td>Replace Infinity ID Expiratory Valve or acknowledge message by touching &quot;ALARM RESET&quot; button and confirm with rotary knob.</td>
</tr>
<tr>
<td>! 020</td>
<td>Import failed, check settings</td>
<td>Configuration import failed.</td>
<td>Check all settings and adjust if necessary. Acknowledge message by touching &quot;ALARM RESET&quot; button and confirm with rotary knob.</td>
</tr>
<tr>
<td>! 020</td>
<td>Import successful, check settings</td>
<td>Configuration import was successful.</td>
<td>Check all settings and adjust if necessary. Acknowledge message by touching &quot;ALARM RESET&quot; button and confirm with rotary knob.</td>
</tr>
<tr>
<td>! 150</td>
<td>Inspiratory hold interrupted</td>
<td>The &quot;Man. insp./hold&quot; button was pressed too long.</td>
<td>Release &quot;Man. insp./hold&quot; button.</td>
</tr>
<tr>
<td>!! 210</td>
<td>Internal battery activated</td>
<td>The batteries of PS500 are depleted. Power supply is provided by the internal battery.</td>
<td>Connect device to the mains power supply.</td>
</tr>
</tbody>
</table>
### Alarm – Cause – Remedy

<table>
<thead>
<tr>
<th>Alarm priority</th>
<th>Alarm message</th>
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</tr>
</thead>
<tbody>
<tr>
<td>!! 120</td>
<td>Internal power supply failure</td>
<td>Technical failure detected.</td>
<td>To continue ventilation with this device, continuously monitor the device functions. Call DrägerService.</td>
</tr>
<tr>
<td>! 140</td>
<td>Leakage</td>
<td>Only monitored for intubated patients! The measured relative leakage exceeds 55 %.</td>
<td>Check for leakages in breathing circuit. Make sure that the tube is connected correctly.</td>
</tr>
<tr>
<td>!!! 205</td>
<td>Mean airway pressure high</td>
<td>Mean airway pressure exceeds the set value by more than 5 mbar (5 cmH₂O).</td>
<td>Check patient condition. Check ventilation settings.</td>
</tr>
<tr>
<td>!!! 200</td>
<td>Mean airway pressure low</td>
<td>Measured Pmean is 7 mbar (7 cmH₂O) lower than set Pmean.</td>
<td>Check breathing circuit for tight connections. Check whether the expiratory valve is properly engaged. Make sure that the tube is connected correctly.</td>
</tr>
<tr>
<td>! 008</td>
<td>MEDIBUS communication failed</td>
<td>MEDIBUS communication failure.</td>
<td>Ventilation functions are not affected. Check MEDIBUS connection. Check MEDIBUS settings.</td>
</tr>
</tbody>
</table>
### Alarm – Cause – Remedy

<table>
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<tr>
<th>Alarm priority</th>
<th>Alarm message</th>
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</tr>
</thead>
<tbody>
<tr>
<td>!!! 160</td>
<td>MV high</td>
<td>The minute volume exceeds the upper alarm limit.</td>
<td>Check patient condition.</td>
</tr>
<tr>
<td></td>
<td></td>
<td></td>
<td>Check ventilation settings.</td>
</tr>
<tr>
<td></td>
<td></td>
<td></td>
<td>Adjust alarm limit if necessary.</td>
</tr>
<tr>
<td></td>
<td></td>
<td>Water in flow sensor.</td>
<td>Drain water trap of breathing circuit. Dry flow sensor.</td>
</tr>
<tr>
<td></td>
<td></td>
<td>Flow sensor is not calibrated or faulty.</td>
<td>Calibrate flow sensor and replace it if necessary.</td>
</tr>
</tbody>
</table>
### Alarm – Cause – Remedy

<table>
<thead>
<tr>
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<th>Remedy</th>
</tr>
</thead>
<tbody>
<tr>
<td>!!! 160</td>
<td>MV low</td>
<td>The minute volume has fallen below the lower alarm limit.</td>
<td>Check patient condition.</td>
</tr>
<tr>
<td></td>
<td></td>
<td></td>
<td>Check ventilation settings.</td>
</tr>
<tr>
<td></td>
<td></td>
<td></td>
<td>Adjust alarm limit if necessary.</td>
</tr>
<tr>
<td></td>
<td>Obstruction.</td>
<td>Check patient condition.</td>
<td>Check breathing circuit.</td>
</tr>
<tr>
<td></td>
<td></td>
<td></td>
<td>Check tube or mask.</td>
</tr>
<tr>
<td></td>
<td>Flow sensor is not calibrated or faulty.</td>
<td>Calibrate flow sensor and replace it if necessary.</td>
<td></td>
</tr>
<tr>
<td></td>
<td>Leakage or disconnection.</td>
<td>Check breathing circuit for tight connections.</td>
<td>Check whether the expiratory valve is properly engaged.</td>
</tr>
<tr>
<td></td>
<td></td>
<td></td>
<td>Make sure that the tube or mask is connected correctly.</td>
</tr>
<tr>
<td></td>
<td>Device failure.</td>
<td>Disconnect patient from the device and continue ventilation without delay using another independent ventilator.</td>
<td>Call DrägerService.</td>
</tr>
<tr>
<td>Alarm priority</td>
<td>Alarm message</td>
<td>Cause</td>
<td>Remedy</td>
</tr>
<tr>
<td>---------------</td>
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</tr>
<tr>
<td>!! 110</td>
<td>Nebulization canceled</td>
<td>Air and O2 supply insufficient to deliver required flow and pressure for nebulization. Nebulization canceled.</td>
<td>Check connections to Air and O2 supply. Make sure supply pressures are greater than 3 bar (43.5 psi). Acknowledge message by touching &quot;ALARM RESET&quot; button and confirm with rotary knob.</td>
</tr>
<tr>
<td></td>
<td>Inspiratory flow insufficient for nebulization.</td>
<td>Increase inspiratory flow to more than 6 L/min for neonates and pediatric patients. Acknowledge message by touching &quot;ALARM RESET&quot; button and confirm with rotary knob.</td>
<td></td>
</tr>
<tr>
<td></td>
<td>Air supply insufficient to deliver required flow and pressure for nebulization.</td>
<td>Check connection to Air supply. Make sure supply pressure is greater than 3 bar (43.5 psi). Acknowledge message by touching &quot;ALARM RESET&quot; button and confirm with rotary knob.</td>
<td></td>
</tr>
<tr>
<td></td>
<td>O2 supply insufficient to deliver required flow and pressure for nebulization.</td>
<td>Check connection to O2 supply. Make sure supply pressure is greater than 3 bar (43.5 psi). Acknowledge message by touching &quot;ALARM RESET&quot; button and confirm with rotary knob.</td>
<td></td>
</tr>
</tbody>
</table>
## Alarm – Cause – Remedy

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</tr>
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<tbody>
<tr>
<td></td>
<td>Internal supply pressures too high.</td>
<td>Acknowledge message by touching &quot;ALARM RESET&quot; button and confirm with rotary knob.</td>
<td>Call DrägerService.</td>
</tr>
<tr>
<td></td>
<td>Air and O2 supply inappropriate to deliver required flow and pressure for nebulization.</td>
<td>Nebulization canceled.</td>
<td></td>
</tr>
<tr>
<td></td>
<td>Neonatal flow monitoring active.</td>
<td>Deactivate neonatal flow monitoring and remove neonatal flow sensor.</td>
<td></td>
</tr>
<tr>
<td></td>
<td>Nebulization is only possible if neonatal flow monitoring is deactivated and neonatal flow sensor is removed from breathing circuit.</td>
<td>Acknowledge message by touching &quot;ALARM RESET&quot; button and confirm with rotary knob.</td>
<td></td>
</tr>
<tr>
<td></td>
<td>Incompatible ventilation mode.</td>
<td>Select an appropriate ventilation mode.</td>
<td></td>
</tr>
<tr>
<td></td>
<td>Nebulization is only possible in pressure-controlled ventilation modes without Volume Guarantee.</td>
<td>Acknowledge message by touching &quot;ALARM RESET&quot; button and confirm with rotary knob.</td>
<td></td>
</tr>
<tr>
<td>I 100</td>
<td>Nebulization finished or canceled.</td>
<td>Install neonatal flow sensor. Switch on neonatal flow monitoring. Acknowledge message by pressing &quot;ALARM RESET&quot; and confirm.</td>
<td></td>
</tr>
<tr>
<td>Alarm priority</td>
<td>Alarm message</td>
<td>Cause</td>
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</tr>
</tbody>
</table>
| !! 100         | Nebulizer uses Air only | O2 supply insufficient to deliver required flow and pressure for nebulization.  
Nebulizer is supplied with Air only.  
Increased deviation from the set FiO2. | Check connection to O2 supply.  
Make sure supply pressure is greater than 3 bar (43.5 psi).  
Downgrade alarm priority by touching "ALARM RESET" button and confirm with rotary knob. |
| !! 100         | Nebulizer uses O2 only | Air supply insufficient to deliver required flow and pressure for nebulization.  
Nebulizer is supplied with O2 only.  
Increased deviation from the set FiO2. | Check connection to Air supply.  
Make sure supply pressure is greater than 3 bar (43.5 psi).  
Downgrade alarm priority by touching "ALARM RESET" button and confirm with rotary knob. |
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</thead>
<tbody>
<tr>
<td>228</td>
<td>Neonatal flow measurement failed</td>
<td>Neonatal flow measurement malfunction</td>
<td>In case of modes with tidal volume or trigger setting:</td>
</tr>
<tr>
<td></td>
<td></td>
<td></td>
<td>Check ventilation settings.</td>
</tr>
<tr>
<td></td>
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<td></td>
<td>Change ventilation mode if required.</td>
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<tr>
<td></td>
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<td></td>
<td>Use external flow monitoring and deactivate the integrated flow monitoring.</td>
</tr>
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<td></td>
<td></td>
<td></td>
<td>Call DrägerService.</td>
</tr>
<tr>
<td></td>
<td></td>
<td></td>
<td><strong>In case of modes without tidal volume or trigger setting:</strong></td>
</tr>
<tr>
<td></td>
<td></td>
<td></td>
<td>Ventilation functions are not affected.</td>
</tr>
<tr>
<td></td>
<td></td>
<td></td>
<td>To continue ventilation with this device, use external flow monitoring and deactivate the integrated neonatal flow monitoring.</td>
</tr>
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<td></td>
<td></td>
<td></td>
<td>Call DrägerService.</td>
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</table>
Alarm – Cause – Remedy

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</thead>
<tbody>
<tr>
<td>!!! 228</td>
<td>Neonatal flow sensor failure</td>
<td>Neonatal flow sensor cable faulty.</td>
<td>Replace neonatal flow sensor cable.</td>
</tr>
<tr>
<td></td>
<td></td>
<td>Neonatal flow sensor faulty.</td>
<td>Replace neonatal flow sensor or sensor insert and calibrate the new sensor.</td>
</tr>
<tr>
<td></td>
<td></td>
<td>Neonatal flow measurement</td>
<td>In case of modes with tidal volume or trigger setting:</td>
</tr>
<tr>
<td></td>
<td></td>
<td>malfunction.</td>
<td>Check ventilation settings.</td>
</tr>
<tr>
<td></td>
<td></td>
<td></td>
<td>Change ventilation mode if required.</td>
</tr>
<tr>
<td></td>
<td></td>
<td></td>
<td>Use external flow monitoring and deactivate the integrated flow monitoring.</td>
</tr>
<tr>
<td></td>
<td></td>
<td></td>
<td>Call DrägerService.</td>
</tr>
<tr>
<td></td>
<td></td>
<td>In case of modes without tidal</td>
<td>Ventilation functions are not affected.</td>
</tr>
<tr>
<td></td>
<td></td>
<td>volume or trigger setting:</td>
<td>To continue ventilation with this device, use external flow monitoring and deactivate the</td>
</tr>
<tr>
<td></td>
<td></td>
<td></td>
<td>integrated neonatal flow monitoring.</td>
</tr>
<tr>
<td></td>
<td></td>
<td></td>
<td>Call DrägerService.</td>
</tr>
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<tbody>
<tr>
<td>! 100</td>
<td>Neonatal flow sensor replaced?</td>
<td>Reconnection of the neonatal flow sensor detected.</td>
<td>Confirm message if calibrated neonatal flow sensor is still used. Otherwise calibrate neonatal flow sensor.</td>
</tr>
<tr>
<td></td>
<td></td>
<td>Neonatal flow monitoring was temporarily deactivated.</td>
<td>Confirm message if calibrated neonatal flow sensor is still used. Otherwise calibrate neonatal flow sensor.</td>
</tr>
<tr>
<td>!! 115</td>
<td>Neonatal flow sensor soiled</td>
<td>Water or secretion in the neonatal flow sensor.</td>
<td>Replace neonatal flow sensor or sensor insert and calibrate the new sensor.</td>
</tr>
<tr>
<td>!!! 229</td>
<td>Neonatal flow sensor?</td>
<td>Neonatal flow sensor is not connected.</td>
<td>Check connections of the neonatal flow sensor and cable.</td>
</tr>
<tr>
<td></td>
<td></td>
<td>Neonatal flow sensor malfunction.</td>
<td>Replace neonatal flow sensor or sensor insert and calibrate the new sensor.</td>
</tr>
<tr>
<td>!!! 140</td>
<td>Neonatal flow sensor?</td>
<td>Neonatal flow sensor not installed in the breathing circuit.</td>
<td>Check whether the neonatal flow sensor is fitted correctly.</td>
</tr>
<tr>
<td></td>
<td></td>
<td></td>
<td>Replace neonatal flow sensor if necessary.</td>
</tr>
<tr>
<td>!!! 250</td>
<td>No Air supply</td>
<td>Air supply insufficient to deliver required flow and pressure.</td>
<td>Check connection to Air supply. Make sure supply pressure is greater than 3 bar (43.5 psi). Consider readjusting ventilation settings. Remove connection to Air supply if alarm condition persists (to avoid reverse flow into the Air supply).</td>
</tr>
</tbody>
</table>

Instructions for use Infinity Acute Care System – Babylog VN500 SW 2.n
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</tr>
</thead>
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<tr>
<td>! 100</td>
<td>No Air supply</td>
<td>Air supply insufficient. If FiO2 = 100 Vol%, Air supply is not required.</td>
<td>Check connection to Air supply. Make sure supply pressure is greater than 3 bar (43.5 psi). Consider readjusting ventilation settings. Remove connection to Air supply if alarm condition persists (to avoid reverse flow into the Air supply).</td>
</tr>
<tr>
<td>!!! 250</td>
<td>No O2 supply</td>
<td>O2 supply insufficient to deliver required flow and pressure. Gas delivery system supplied with Air only. Ventilation continues with Air only.</td>
<td>Check connection to O2 supply. Make sure supply pressure is greater than 3 bar (43.5 psi). Consider readjusting ventilation settings. Remove connection to O2 supply if alarm condition persists (to avoid reverse flow into the O2 supply).</td>
</tr>
<tr>
<td>! 100</td>
<td>No O2 supply</td>
<td>O2 supply insufficient. If FiO2 = 21 Vol%, O2 supply is not required.</td>
<td>Check connection to O2 supply. Make sure supply pressure is greater than 3 bar (43.5 psi). Consider readjusting ventilation settings. Remove connection to O2 supply if alarm condition persists (to avoid reverse flow into the O2 supply).</td>
</tr>
</tbody>
</table>
## Alarm – Cause – Remedy

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<tbody>
<tr>
<td>!! 119</td>
<td>Nurse call failure</td>
<td>Technical failure detected.</td>
<td>To continue ventilation with this device, continuously monitor the device functions. Call DrägerService.</td>
</tr>
<tr>
<td>!! 110</td>
<td>O₂ and Air supply pressures differ too much</td>
<td>The difference between O₂ supply pressure and Air supply pressure can lead to an incorrect O₂ concentration during nebulization.</td>
<td>Check connections to Air and O₂ supply. Make sure supply pressures are greater than 3 bar (43.5 psi). Acknowledge message by touching &quot;ALARM RESET&quot; button and confirm with rotary knob.</td>
</tr>
<tr>
<td>!!! 132</td>
<td>O₂ measurement failed</td>
<td>O₂ measurement failed.</td>
<td>To calibrate the O₂ sensor, perform the device check. Ventilation can be continued also if the alarm does not disappear. Use external O₂ monitoring and deactivate the integrated O₂ monitoring. Call DrägerService.</td>
</tr>
<tr>
<td>!! 110</td>
<td>O₂ pressure regulation failed</td>
<td>Internal O₂ supply pressure too high. O₂ pressure regulator failure.</td>
<td>No HFO possible. Call DrägerService.</td>
</tr>
<tr>
<td>!! 040</td>
<td>Oxygenation maneuver failed</td>
<td>Internal error during oxygenation maneuver.</td>
<td>Do not perform suction maneuver until the device was checked. Call DrägerService.</td>
</tr>
</tbody>
</table>
### Alarm – Cause – Remedy

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<tr>
<td>!!! 140</td>
<td>PEEP high</td>
<td>Expiratory valve or breathing circuit obstructed.</td>
<td>Check breathing circuit and expiratory valve.</td>
</tr>
<tr>
<td></td>
<td></td>
<td>Expiratory resistance increased.</td>
<td>Check viral/bacterial filter. Replace it if necessary.</td>
</tr>
<tr>
<td></td>
<td></td>
<td>Device failure.</td>
<td>Disconnect patient from the device and continue ventilation without delay using another independent ventilator. Call DrägerService.</td>
</tr>
<tr>
<td>!! 140</td>
<td>PEEP high</td>
<td>Expiratory valve or breathing circuit obstructed.</td>
<td>Check breathing circuit and expiratory valve.</td>
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<tr>
<td></td>
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<td>Expiratory resistance increased.</td>
<td>Check viral/bacterial filter. Replace it if necessary.</td>
</tr>
<tr>
<td></td>
<td></td>
<td>Device failure.</td>
<td>Disconnect patient from the device and continue ventilation without delay using another independent ventilator. Call DrägerService.</td>
</tr>
<tr>
<td>!!! 140</td>
<td>PEEP low</td>
<td>Measured PEEP is 3 mbar (3 cmH2O) less than set PEEP.</td>
<td>Check breathing circuit for tight connections.</td>
</tr>
<tr>
<td></td>
<td></td>
<td></td>
<td>Check whether the expiratory valve is properly engaged.</td>
</tr>
<tr>
<td></td>
<td></td>
<td></td>
<td>Make sure that the tube or mask is connected correctly.</td>
</tr>
<tr>
<td>!! 210</td>
<td>Perform device and breathing circuit check</td>
<td>Device check and breathing circuit check must be performed before operation.</td>
<td>Perform device check. Acknowledge message by touching &quot;ALARM RESET&quot; button and confirm with rotary knob.</td>
</tr>
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## Alarm - Cause - Remedy

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<tr>
<td>!!! 140</td>
<td>Plow high</td>
<td>Expiratory valve or breathing circuit obstructed.</td>
<td>Check breathing circuit and expiratory valve.</td>
</tr>
<tr>
<td></td>
<td></td>
<td></td>
<td>Check for condensate.</td>
</tr>
<tr>
<td></td>
<td></td>
<td>Expiratory resistance increased.</td>
<td>Check viral/bacterial filter. Replace it if necessary.</td>
</tr>
<tr>
<td></td>
<td></td>
<td>Device failure.</td>
<td>Disconnect patient from the device and continue ventilation without delay using another independent ventilator.</td>
</tr>
<tr>
<td></td>
<td></td>
<td></td>
<td>Call DrägerService.</td>
</tr>
<tr>
<td></td>
<td></td>
<td>Not monitored if AutoRelease is enabled.</td>
<td>To enable monitoring switch off AutoRelease or increase Tlow to &gt;1 second.</td>
</tr>
<tr>
<td></td>
<td></td>
<td>Not monitored if Tlow is set to less than 1 second.</td>
<td></td>
</tr>
<tr>
<td>!!! 140</td>
<td>Plow low</td>
<td>Measured Plow is 5 mbar (5 cmH2O) less than set Plow.</td>
<td>Check breathing circuit for tight connections.</td>
</tr>
<tr>
<td></td>
<td></td>
<td></td>
<td>Check whether the expiratory valve is properly engaged.</td>
</tr>
<tr>
<td></td>
<td></td>
<td></td>
<td>Make sure that the tube or mask is connected correctly.</td>
</tr>
<tr>
<td>! 140</td>
<td>Pressure limited</td>
<td>The pressure of a breath is limited by the set &quot;Paw high&quot; limit or Pmax.</td>
<td>Check patient condition.</td>
</tr>
<tr>
<td></td>
<td></td>
<td></td>
<td>Check ventilation settings.</td>
</tr>
<tr>
<td></td>
<td></td>
<td></td>
<td>Adjust &quot;Paw high&quot; alarm limit or Pmax.</td>
</tr>
<tr>
<td>!!! 238</td>
<td>Pressure measurement failed</td>
<td>Pressure measurement malfunction.</td>
<td>Disconnect patient from the device and continue ventilation without delay using another independent ventilator.</td>
</tr>
<tr>
<td></td>
<td></td>
<td></td>
<td>Call DrägerService.</td>
</tr>
</tbody>
</table>
### Alarm – Cause – Remedy

<table>
<thead>
<tr>
<th>Alarm priority</th>
<th>Alarm message</th>
<th>Cause</th>
<th>Remedy</th>
</tr>
</thead>
<tbody>
<tr>
<td>!! 140</td>
<td>Pressure measurement impaired</td>
<td>Pressure measurement malfunction. Accuracy of measured values based on pressure could be impaired. To continue ventilation with this device, continuously monitor the device functions. Call DrägerService.</td>
<td></td>
</tr>
<tr>
<td>!! 100</td>
<td>Pressure measurement inaccurate</td>
<td>Fluid in expiratory valve. Replace expiratory valve. Clean and dry used one.</td>
<td></td>
</tr>
<tr>
<td></td>
<td></td>
<td>Breathing circuit check has not been performed. Perform or repeat breathing circuit check.</td>
<td></td>
</tr>
<tr>
<td></td>
<td></td>
<td>The inspiratory or expiratory hose is obstructed. Check breathing circuit.</td>
<td></td>
</tr>
<tr>
<td></td>
<td></td>
<td>Pressure measurement failure. Disconnect patient from the device and continue ventilation without delay using another independent ventilator. Call DrägerService.</td>
<td></td>
</tr>
<tr>
<td>!! 140</td>
<td>Pressure sensor? Ventilation impaired</td>
<td>Ventilation patterns for which a pressure sensor is necessary cannot be performed. The ventilation unit applies back-up ventilation. To continue ventilation with this device, use external pressure monitoring. Call DrägerService.</td>
<td></td>
</tr>
<tr>
<td>!! 100</td>
<td>Product test: Not for clinical use</td>
<td>License for product test is installed.</td>
<td>Call DrägerService.</td>
</tr>
</tbody>
</table>
### Alarm – Cause – Remedy

<table>
<thead>
<tr>
<th>Alarm priority</th>
<th>Alarm message</th>
<th>Cause</th>
<th>Remedy</th>
</tr>
</thead>
<tbody>
<tr>
<td>!!! 150</td>
<td>Respiratory rate high</td>
<td>The patient is breathing at a high respiratory rate.</td>
<td>Check patient condition.</td>
</tr>
<tr>
<td></td>
<td></td>
<td></td>
<td>Check ventilation settings or spontaneous respiratory rate.</td>
</tr>
<tr>
<td></td>
<td></td>
<td></td>
<td>Adjust alarm limit if necessary.</td>
</tr>
<tr>
<td></td>
<td></td>
<td>The set respiratory rate exceeds upper alarm limit.</td>
<td>Adjust the respiratory rate or the upper alarm limit for the respiratory rate.</td>
</tr>
<tr>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td></td>
<td></td>
<td>Auto triggering caused by water in the breathing circuit.</td>
<td>Drain water trap of breathing circuit. Dry flow sensor.</td>
</tr>
<tr>
<td></td>
<td></td>
<td></td>
<td>Check breathing circuit.</td>
</tr>
<tr>
<td>! 100</td>
<td>Restart of ventilation unit</td>
<td>Technical failure detected. Last restart was delayed.</td>
<td>Downgrade alarm priority by touching &quot;ALARM RESET&quot; button and confirm with rotary knob.</td>
</tr>
<tr>
<td></td>
<td>delayed</td>
<td></td>
<td>To continue ventilation with this device, continuously monitor the device functions.</td>
</tr>
<tr>
<td></td>
<td></td>
<td></td>
<td>Call DrägerService.</td>
</tr>
<tr>
<td>!! 050</td>
<td>Rotary knob stuck or pressed</td>
<td>The rotary knob is either faulty or was pressed for more than 20 seconds without turning.</td>
<td>If you are still pressing the rotary knob, release it. Otherwise press and turn rotary knob repeatedly. If alarm condition persists, settings cannot be adjusted anymore.</td>
</tr>
<tr>
<td></td>
<td>too long</td>
<td></td>
<td>Disconnect patient from the device and continue ventilation without delay using another independent ventilator. Call DrägerService.</td>
</tr>
</tbody>
</table>
## Alarm – Cause – Remedy

<table>
<thead>
<tr>
<th>Alarm priority</th>
<th>Alarm message</th>
<th>Cause</th>
<th>Remedy</th>
</tr>
</thead>
</table>
| !! 050         | Rotary knob used too often         | The rotary knob is either faulty or was pressed more than 5 times per second. | Press and turn rotary knob repeatedly.  
If alarm condition persists, settings cannot be adjusted anymore.  
Disconnect patient from the device and continue ventilation without delay using another independent ventilator.  
Call DrägerService. |
| ! 100          | Service date approaching           | Service date is almost reached.                                      | Acknowledge message by touching "ALARM RESET" button and confirm with rotary knob.  
Call DrägerService. |
| ! 100          | Service date reached               | Service is due.                                                      | Acknowledge message by touching "ALARM RESET" button and confirm with rotary knob.  
Call DrägerService. |
| ! 200          | Setting not confirmed              | One or more settings have been changed but not confirmed.            | If necessary, change these settings and confirm the change with the rotary knob. |
| !!! 255        | Standby mode activated             | Device has been switched to standby mode.                            | Acknowledge standby mode by touching "ALARM RESET" button and confirm with rotary knob. |
| !! 040         | Suction maneuver failed            | Internal error during suction maneuver.                              | Do not perform suction maneuver until the device was checked.  
Call DrägerService. |
| ! 140          | Suction maneuver overused?         | The suction maneuver has been performed more than 5 times within an hour. | Perform suction maneuver less frequently. |
## Alarm – Cause – Remedy

<table>
<thead>
<tr>
<th>Alarm priority</th>
<th>Alarm message</th>
<th>Cause</th>
<th>Remedy</th>
</tr>
</thead>
<tbody>
<tr>
<td>! 200</td>
<td>Ventilation mode not confirmed</td>
<td>The ventilation mode has been changed but not confirmed.</td>
<td>If necessary, change the ventilation mode and confirm the change with the rotary knob.</td>
</tr>
<tr>
<td>!! 255</td>
<td>Ventilation unit restarted</td>
<td>Internal communication error caused restart of the ventilation unit.</td>
<td>Check all therapy settings and adjust them if necessary. Acknowledge message by touching &quot;ALARM RESET&quot; button and confirm with rotary knob.</td>
</tr>
<tr>
<td>! 190</td>
<td>VT limited</td>
<td>The delivered tidal volume exceeds the set maximum tidal volume.</td>
<td>Check patient condition. Check ventilation settings.</td>
</tr>
<tr>
<td>!! 166</td>
<td>VT low</td>
<td>The applied VT has been less than 90% of the set VT for more than five (pediatric patients) or eight (neonatal patients) consecutive breaths.</td>
<td>Check ventilation settings. Check patient condition. Adjust &quot;Paw high&quot; alarm limit or Pmax.</td>
</tr>
<tr>
<td>! 140</td>
<td>VT not reached, leakage</td>
<td>Set volume cannot be reached. Flow delivery terminated.</td>
<td>Check for leakages in breathing circuit. Make sure that the tube or mask is connected correctly.</td>
</tr>
</tbody>
</table>
## Alarm – Cause – Remedy

<table>
<thead>
<tr>
<th>Alarm priority</th>
<th>Alarm message</th>
<th>Cause</th>
<th>Remedy</th>
</tr>
</thead>
<tbody>
<tr>
<td>!! 140</td>
<td>VT/h not reached</td>
<td>Tidal volume could not be reached due to increased resistance.</td>
<td>Check patient condition.</td>
</tr>
<tr>
<td></td>
<td></td>
<td></td>
<td>Check ventilation settings.</td>
</tr>
<tr>
<td></td>
<td></td>
<td>Tidal volume could not be reached due to pressure limitation of the device.</td>
<td>Reduce f/h or increase I:Eh to 1:1.</td>
</tr>
<tr>
<td>!! 100</td>
<td>Wrong or invalid applications found</td>
<td>Wrong or defective application card.</td>
<td>Call DrägerService.</td>
</tr>
</tbody>
</table>
This page has been left blank intentionally.
Reprocessing

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Disassembly

Observe before disassembly
1. Switch off the device and all devices connected to it.
2. Disconnect the mains plugs.
3. Drain the water traps and the breathing hoses.
4. Drain the water reservoir of the breathing gas humidifier.

Disconnecting the breathing circuit
1. Pull the breathing hoses from the inspiratory port and the expiratory port (A).
2. If fitted: Remove the water trap (B) from the breathing hose.
3. Remove the water trap container from the water trap. Empty the water trap container.

Dismantling the expiratory valve

Removing the expiratory valve
1. Turn the locking ring (A) as far as possible to the left.
2. Remove the expiratory valve from the fitting.

Dismantling the expiratory valve
1. Remove the muffler (B) from the expiratory valve.
2. Remove the diaphragm (C). Do not dismantle the diaphragm any further.
3. Remove the water trap container (D). Empty the water trap container.
Reprocessing

Dismantling the CO₂ sensor

1. Remove the CO₂ sensor plug from the socket.
2. Remove the CO₂ sensor (A) from the cuvette.
3. Remove the cuvette (B) from the patient connector of the Y-piece.

Dismantling the neonatal flow sensor

- Disconnect the sensor plug on the rear of the device.

Dismantling the neonatal flow sensor ISO 15 (8411130)

1. Remove the flow sensor housing (A) from the Y-piece.
2. Disconnect plug (D) of the flow sensor cable from the neonatal flow sensor.
3. Gently press the knobs (B) on both sides while pulling the insert (C) out of the flow sensor housing.

Dismantling the neonatal flow sensor Y-piece (8410185)

1. Pull the Y-piece (A) out of the breathing hoses.
2. Disconnect plug (D) of the flow sensor cable from the neonatal flow sensor.
3. Gently press the knobs (B) on both sides while pulling the insert (C) out of the Y-piece.

Dismantling the inspiratory unit

When the inspiratory unit must be reprocessed:
The inspiratory unit must only be reprocessed when patient gas has passed through the safety valve. In the case of spontaneously breathing patients, this can occur in the following situations:
- Excess pressure in the system caused by a kink in the expiratory hose
- Failure of both supply gases
- Complete failure of the power supply (failure of mains power supply and discharged or faulty batteries)
Removing the inspiratory unit

The inspiratory unit must only be removed when the device is switched off.

1. Press and hold the locking lever (A) on the underside of the inspiratory unit.
2. Simultaneously turn the inspiratory unit (B) approx. 20° counterclockwise.
3. Remove the inspiratory unit from the fitting.

Dismantling the inspiratory unit

1. Remove the diaphragm with adapter (C) from the fitting of the inspiratory unit.
2. Do not dismantle the inspiratory unit any further.

Dismantling the pneumatic medication nebulizer

1. Remove the nebulizer hose (A) from the medication nebulizer (B) and from the nebulizer port on the device.
2. Remove the medication nebulizer (B) from the breathing circuit.
3. Remove the corrugated hose for the breathing circuit (C) from the inlet port.
4. Remove the corrugated hose (D) from the outlet port.
5. Dismantle the medication nebulizer in accordance with the corresponding instructions for use.

Dismantling other accessories

- Dismantle the breathing gas humidifier, the Aeroneb nebulizer and the bacterial filter in accordance with the corresponding instructions for use.
Reprocessing

Information on reprocessing

Instructions for reprocessing are based on internationally accepted guidelines, e.g., standard ISO 17664.

The components through which contaminated breathing gas passes during normal operation and in the event of a fault must be reprocessed. In normal operation breathing gas passes through the expiratory valve or the expiratory valve with ejector and muffler and other accessories in the expiratory path. In the event of a fault, the inspiratory unit and other accessories in the inspiratory path can become contaminated.

Safety information

**WARNING**
Risk due to inappropriately reprocessed products

Reusable products must be reprocessed, otherwise there is an increased risk of infection and the products may no longer function correctly.

- Observe the hygiene regulations and reprocessing regulations of the healthcare facility.
- Observe national hygiene regulations and reprocessing regulations.
- Use validated procedures for reprocessing.
- Reprocess reusable products after every use.
- Observe the manufacturer’s instructions for cleaning agents, disinfectants, and reprocessing devices.

**CAUTION**
Risk due to faulty products

Signs of wear, e.g., cracks, deformation, discoloration, or peeling, may occur with reprocessed products.

Check the products for signs of wear and replace them if necessary.

**CAUTION**
Health hazard

Do not sterilize parts in ethylene oxide. Ethylene oxide may diffuse into the parts.

**CAUTION**
For infectious patients, all parts that come into contact with breathing gas also have to be sterilized after disinfection and cleaning.
Reprocessing

Safety information on disposable articles

**WARNING**
Risk of patient injury as a result of failure of the accessories

Disposable articles were developed, tested and manufactured for single use only. Reuse, reprocessing or sterilization can lead to a failure of the accessories and cause injuries to the patient.

Do not reuse, reprocess, or sterilize disposable articles.

**CAUTION**
Risk of failure of flow measurement

Improper reprocessing and soiling, such as deposits or particles, can damage the flow sensor:
- No machine cleaning or disinfection of the sensor insert
- No plasma sterilization or radiation sterilization
- No compressed air
- No water jets, compressed air, brushes or the like when cleaning the sensor insert
- No ultrasonic bath
- Clean and disinfect the flow sensor in accordance with the corresponding instructions for use.
- For disinfecting the flow sensor use only clean disinfectant solutions.

**NOTE**
- Do not use brushes for reprocessing the sensor insert and do not use a syringe on the sensor insert.
- For reprocessing the housing use lint-free brushes only.

Safety information on the neonatal flow sensor

**WARNING**
Risk of fire

Residual vapors of easily flammable disinfectants (e.g., alcohols) and deposits that were not removed during reprocessing can ignite when the flow sensor is in use.
- Ensure particle-free cleaning and disinfection.
- After disinfection, allow the flow sensor to air for at least 30 minutes.
- Before inserting the flow sensor check for visible damage and soiling, such as residual mucus, medication aerosols, and particles.
- Replace flow sensors when damaged, soiled, or not particle-free.
Reprocessing

Classifications for reprocessing

Classification of medical devices

Medical devices and their components are classified according to the way they are used and the resulting risk.

<table>
<thead>
<tr>
<th>Classification</th>
<th>Definition</th>
</tr>
</thead>
<tbody>
<tr>
<td>Non-critical</td>
<td>Components that come only into contact with skin that is intact</td>
</tr>
<tr>
<td>Semi-critical (A, B)</td>
<td>Components that carry breathing gas or come into contact with mucous membranes or pathologically altered skin</td>
</tr>
<tr>
<td>Critical (A, B, C)</td>
<td>Components that penetrate skin or mucous membranes or come into contact with blood</td>
</tr>
</tbody>
</table>

Classification of device-specific components

Observe the instructions for use for the components.

The following classification is a recommendation from Dräger.

**Non-critical**
- Babylog VN500 ventilation unit
- Medical Cockpit display unit
- GS500 gas supply unit
- PS500 power supply unit
- Trolley with accessory mounts
- System cable
- Compressed gas hoses
- CO2 sensor
- Connection cable for the neonatal flow sensor
- Data link cable

**Semi-critical A**
- Reusable cuvette for the CO2 sensor
- Neonatal flow sensor Y-piece or ISO 15, including individual parts

**Semi-critical B**
- Infinity ID neonatal expiratory valve, including individual parts
- Inspiratory unit, including individual parts
## Reprocessing

### Reprocessing list

<table>
<thead>
<tr>
<th>Components</th>
<th>Surface disinfection with cleaning</th>
<th>Manual cleaning followed by disinfection by immersion</th>
<th>Machine cleaning with thermal disinfection</th>
<th>Steam sterilization</th>
<th>Special reprocessing measures</th>
</tr>
</thead>
<tbody>
<tr>
<td>Babylog VN500 ventilation unit</td>
<td>Yes</td>
<td>No</td>
<td>No</td>
<td>No</td>
<td>No</td>
</tr>
<tr>
<td>Medical Cockpit display unit</td>
<td>Yes</td>
<td>No</td>
<td>No</td>
<td>No</td>
<td>Do not spray cleaning agent and disinfectant directly on the touch screen.</td>
</tr>
<tr>
<td>GS500 gas supply unit</td>
<td>Yes</td>
<td>No</td>
<td>No</td>
<td>No</td>
<td>No</td>
</tr>
<tr>
<td>PS500 power supply unit</td>
<td>Yes</td>
<td>No</td>
<td>No</td>
<td>No</td>
<td>No</td>
</tr>
<tr>
<td>Trolley with accessory mounts</td>
<td>Yes</td>
<td>No</td>
<td>No</td>
<td>No</td>
<td>No</td>
</tr>
<tr>
<td>System cable</td>
<td>Yes</td>
<td>No</td>
<td>No</td>
<td>No</td>
<td>No</td>
</tr>
<tr>
<td>Compressed gas hoses</td>
<td>Yes</td>
<td>No</td>
<td>No</td>
<td>No</td>
<td>No</td>
</tr>
<tr>
<td>CO2 sensor</td>
<td>Yes</td>
<td>No</td>
<td>No</td>
<td>No</td>
<td>Avoid residues on the test filter.</td>
</tr>
<tr>
<td>Connection cable for the neonatal flow sensor</td>
<td>Yes</td>
<td>No</td>
<td>No</td>
<td>No</td>
<td>No</td>
</tr>
<tr>
<td>Data link cable</td>
<td>Yes</td>
<td>No</td>
<td>No</td>
<td>No</td>
<td>No</td>
</tr>
<tr>
<td>Reusable cuvette for the CO2 sensor</td>
<td>No</td>
<td>Possible</td>
<td>Yes</td>
<td>Yes</td>
<td>Wipe off any contamination, particularly inside and outside the windows, using a soft disposable tissue and cotton swabs, under running water if necessary. Only cleaning agent, and no rinse aid, must be used for automatic cleaning of the cuvette. Otherwise, there is a risk of cracks developing.</td>
</tr>
</tbody>
</table>
### Reprocessing

<table>
<thead>
<tr>
<th>Components</th>
<th>Surface disinfection with cleaning</th>
<th>Manual cleaning followed by disinfection by immersion</th>
<th>Machine cleaning with thermal disinfection</th>
<th>Steam sterilization</th>
<th>Special reprocessing measures</th>
</tr>
</thead>
<tbody>
<tr>
<td>Neonatal flow sensor Y-piece or ISO 15</td>
<td>No</td>
<td>Yes</td>
<td>No</td>
<td>No</td>
<td>See &quot;Neonatal flow sensors&quot; on page 242.</td>
</tr>
<tr>
<td>Infinity ID neonatal expiratory valve</td>
<td>No</td>
<td>Possible</td>
<td>Yes</td>
<td>Yes(^1)</td>
<td>See &quot;Infinity ID neonatal expiratory valve&quot; on page 242.</td>
</tr>
<tr>
<td>Inspiratory unit</td>
<td>No</td>
<td>Possible</td>
<td>Yes</td>
<td>Yes(^1)</td>
<td>Only reprocess the inspiratory unit if breathing gas has passed through the safety valve.</td>
</tr>
</tbody>
</table>

\(^1\) For further information on sterilizing, see page 241.
Reprocessing

Reprocessing procedure

Validated reprocessing procedures

At the time of product-specific validation, the following reprocessing procedures showed good material compatibility and effectiveness:

<table>
<thead>
<tr>
<th>Procedure</th>
<th>Agent</th>
<th>Manufacturer</th>
<th>Concentration</th>
<th>Contact time</th>
<th>Temperature</th>
</tr>
</thead>
<tbody>
<tr>
<td>Surface disinfection with cleaning</td>
<td>Buraton 10F</td>
<td>Schülke &amp; Mayr</td>
<td>1 %</td>
<td>30 min</td>
<td>–</td>
</tr>
<tr>
<td></td>
<td>neoform Med AF¹</td>
<td>Dr. Weigert</td>
<td>According to manufacturer's data</td>
<td>According to manufacturer's data</td>
<td>–</td>
</tr>
<tr>
<td></td>
<td>Dismozon pur²</td>
<td>Bode Chemie</td>
<td>1.5 %</td>
<td>15 min</td>
<td>–</td>
</tr>
<tr>
<td>Manual cleaning</td>
<td>Neodisher LM2</td>
<td>Dr. Weigert</td>
<td>3 %</td>
<td>30 min</td>
<td>–</td>
</tr>
<tr>
<td></td>
<td>Sekusept Pulver Classic³</td>
<td>Ecolab</td>
<td>4 %</td>
<td>15 min</td>
<td>–</td>
</tr>
<tr>
<td>Disinfection by immersion⁴⁵³⁵</td>
<td>Sekusept Pulver Classic³</td>
<td>Ecolab</td>
<td>4 %</td>
<td>15 min</td>
<td>–</td>
</tr>
<tr>
<td></td>
<td>Korsolex Extra</td>
<td>Bode Chemie</td>
<td>3 %</td>
<td>15 min</td>
<td>30 min⁶</td>
</tr>
<tr>
<td>Machine cleaning</td>
<td>Neodisher Mediclean</td>
<td>Dr. Weigert</td>
<td>According to manufacturer’s data</td>
<td>–</td>
<td>–</td>
</tr>
<tr>
<td>Machine disinfection (thermal)</td>
<td>–</td>
<td>–</td>
<td>–</td>
<td>10 min</td>
<td>93 °C (199.4 °F)</td>
</tr>
<tr>
<td>Steam sterilization</td>
<td>–</td>
<td>–</td>
<td>–</td>
<td>5 min</td>
<td>134 °C (273.2 °F)</td>
</tr>
</tbody>
</table>

¹) For Dräger Infinity MCable – CO₂ mainstream sensor
²) For Medical Cockpit
³) For Infinity ID neonatal expiratory valve, neonatal flow sensor
⁴) Not for CO₂ sensor
⁵) For neonatal flow sensor
⁶) For neonatal flow sensor
⁷) For reusable cuvette for the CO₂ sensor

The effectiveness of the listed reprocessing procedures has been validated by independent laboratories that certified to the standard ISO 17025.
Reprocessing

Disinfectants

Use disinfectants that are nationally approved and are suitable for the particular reprocessing procedure.

Surface disinfectants

At the time of the test, the surface disinfectants listed in the following table showed good material compatibility. They can be used in addition to the surface disinfectants listed in the section “Validated reprocessing procedures”.

The manufacturers of the surface disinfectants have verified at least the following spectra of activity:

- Bactericidal
- Yeasticidal
- Virucidal or virucidal against enveloped viruses

Observe the specifications of the surface disinfectant manufacturers.

Other surface disinfectants are used at one's own risk.

<table>
<thead>
<tr>
<th>Class of active ingredient</th>
<th>Surface disinfectant</th>
<th>Manufacturer</th>
</tr>
</thead>
<tbody>
<tr>
<td>Chlorine-releasing agents</td>
<td>Actichlor plus</td>
<td>Ecolab</td>
</tr>
<tr>
<td></td>
<td>BruTab 6S</td>
<td>Brulin</td>
</tr>
<tr>
<td></td>
<td>Clorox Professional Disinfecting Bleach Cleaner</td>
<td>Clorox</td>
</tr>
<tr>
<td></td>
<td>Dispatch Hospital Cleaner Disinfectant Towels with Bleach</td>
<td></td>
</tr>
<tr>
<td></td>
<td>Klorsept 17</td>
<td>Medentech</td>
</tr>
<tr>
<td>Oxygen-releasing agents</td>
<td>Descogen Liquid</td>
<td>Antiseptica</td>
</tr>
<tr>
<td></td>
<td>Descogen Liquid r.f.u.</td>
<td>Bode Chemie</td>
</tr>
<tr>
<td></td>
<td>Dismozon plus</td>
<td>Ecolab USA</td>
</tr>
<tr>
<td></td>
<td>Dismozon pur</td>
<td></td>
</tr>
<tr>
<td></td>
<td>Oxyicide</td>
<td>Schülke &amp; Mayr</td>
</tr>
<tr>
<td></td>
<td>Perform</td>
<td>DuPont</td>
</tr>
<tr>
<td>Quaternary ammonium</td>
<td>Mikrozid sensitive liquid¹</td>
<td>Schülke &amp; Mayr</td>
</tr>
<tr>
<td>compounds</td>
<td>Mikrozid sensitive wipes¹</td>
<td>Schülke &amp; Mayr</td>
</tr>
<tr>
<td></td>
<td>Mikrozid alcohol free liquid¹</td>
<td>Schülke &amp; Mayr</td>
</tr>
<tr>
<td></td>
<td>Mikrozid alcohol free wipes¹</td>
<td>Schülke &amp; Mayr</td>
</tr>
<tr>
<td></td>
<td>acryl-des¹</td>
<td>Schülke &amp; Mayr</td>
</tr>
<tr>
<td>Aldehydes</td>
<td>Buraton 10 F</td>
<td>Schülke &amp; Mayr</td>
</tr>
</tbody>
</table>

¹ Virucidal against enveloped viruses

Dräger points out that oxygen-releasing agents and chlorine-releasing agents may cause color change in some materials. Color change does not indicate that the product is not functioning correctly.
Reprocessing

Surface disinfection with cleaning

<table>
<thead>
<tr>
<th>WARNING</th>
</tr>
</thead>
<tbody>
<tr>
<td>Risk due to penetrating liquid</td>
</tr>
<tr>
<td>Penetrating liquid may cause the following:</td>
</tr>
<tr>
<td>- Damage to the device</td>
</tr>
<tr>
<td>- Electric shock</td>
</tr>
<tr>
<td>- Device malfunction</td>
</tr>
<tr>
<td>Ensure that no liquid penetrates the device.</td>
</tr>
</tbody>
</table>

1. Remove soiling immediately. Use a cloth dampened with disinfectant to remove soiling.
2. Perform surface disinfection.
3. After the product has been exposed to the disinfectant for the specified contact time, remove residual disinfectant.
4. Wipe with a cloth dampened with water (preferably drinking-water quality). Allow the product to dry.
5. Check the product for visible soiling. Repeat steps 1 to 5 if necessary.
6. Check the product for visible damage and replace if necessary.

Manual cleaning followed by disinfection by immersion

Manual cleaning

The cleaning agent that is used must have a pH of between 9 and 12.
1. Wash off superficial soiling under running water.
2. Prepare the cleaning solution in accordance with the manufacturer's instructions.
3. Swirl the product backwards and forwards several times in the solution. Make sure that the solution reaches all surfaces and interior spaces.
4. Rinse the product under running water until residual cleaning agent is no longer discernible.
5. Check the product for visible soiling. Repeat steps 1 to 5 if necessary.
6. Check the product for visible damage and replace if necessary.

Disinfection by immersion

1. Prepare the disinfectant solution in accordance with the manufacturer's instructions.
2. Swirl the product backwards and forwards several times in the solution. Make sure that the solution reaches all surfaces and interior spaces.
3. After the contact time has elapsed, rinse the product under running water until residual disinfectant is no longer discernible.
4. Check the product for visible damage and replace if necessary.
5. Thoroughly shake out residual water. Allow the product to dry completely.
Reprocessing

**Machine cleaning with thermal disinfection**

Use a washer-disinfector that meets the requirements of the standard ISO 15883. Dräger recommends the use of a cart for anesthesia accessories and ventilation accessories.

1. Securely position the product in the basket. Ensure the following:
   - All surfaces and interior spaces can be flushed completely.
   - The water can drain off freely.
2. Use a suitable cleaning agent.
3. Select a suitable cycle.
4. Use demineralized water for the final rinsing.
5. After the cycle has ended, check the product for visible soiling. If necessary, repeat the cycle or perform manual cleaning and disinfection by immersion.
6. Check the product for visible damage and replace if necessary.
7. Allow the product to dry completely.

**Steam sterilization**

Use a steam sterilizer that meets the requirements of the standard ISO 17665. Dräger recommends steam sterilization with fractionated vacuum.

Prerequisite: The product has been cleaned and disinfected.

1. Sterilize the product.
2. Check the product for visible damage and replace if necessary.

**Additional information**

Sterilization of the expiratory valve or inspiratory valve may gradually impair the operation of RFID transmission. This may mean that Infinity ID breathing circuit functions may not work or may no longer work reliably. If the message **Infinity ID breathing circuit detected**, is not displayed when an Infinity ID breathing circuit is connected, use a different Infinity ID breathing circuit. If the message is still not displayed, replace the expiratory valve or inspiratory valve.
Reprocessing

Special reprocessing measures

Infinity ID neonatal expiratory valve

Carry out manual cleaning:

1. Immerse the neonatal expiratory valve in the solution and agitate it slightly so that the air can escape.

2. Before the contact time begins and after it has elapsed, fit a syringe (A) containing 20 mL of solution to the ejector channel (B). Inject and extract the solution several times with the syringe.

Perform manual disinfection in the same manner.

Before assembling the neonatal expiratory valve:

NOTE
To ensure the functional integrity of the device, check the ejector (A) for visible residues. If soiled, repeat manual cleaning.

Neonatal flow sensors

Manual cleaning followed by disinfection by immersion must be carried out for complete reprocessing of the flow sensor.

Remove the sensor insert of the neonatal flow sensor from the housing.

NOTE
- Do not use brushes for reprocessing the sensor insert and do not use a syringe on the sensor insert.
- For reprocessing the housing use lint-free brushes only.

Manual cleaning:

The cleaning agent that is used must have a pH of between 9 and 12.

1. Prepare the cleaning agent in accordance with the manufacturer's data in a container with a cover.

2. Place the housing and sensor insert in the solution, ensuring there are no bubbles. Swirl the parts back and forth for approx. 1 minute at the beginning and end of the contact time. Make sure that the solution reaches all surfaces and interior spaces.

3. At the beginning and end of the contact time, spray through each opening in the housing 3 times using a 20 mL syringe (D).
Reprocessing

4 Clean the housing and the Y-piece (B) with a lint-free brush (A): Insert and remove it vertically ten times in each of the two connection openings of the Y-piece (B) and then, at an angle, insert and remove it ten times in both corners of the opening for the sensor insert (C).

5 Rinse the housing and sensor insert in a water bath (preferably drinking-water quality) until cleaning agent residues are no longer discernible.

Disinfection by immersion:

1 Prepare the disinfectant solution in accordance with the manufacturer's data in a container with a cover.

2 Place the housing and sensor insert in the solution, ensuring there are no bubbles. Swirl the parts back and forth for approx. 1 minute at the beginning and end of the contact time. Make sure that the solution reaches all surfaces and interior spaces.

3 At the beginning and end of the contact time, spray through each opening in the housing 3 times using a 20 mL syringe (D).

4 Clean the housing and the Y-piece (B) with a lint-free brush (A): Insert and remove it vertically ten times in each of the two connection openings of the Y-piece (B) and then, at an angle, insert and remove it ten times in both corners of the opening for the sensor insert (C).

5 Rinse the housing and sensor insert in a water bath (preferably drinking-water quality) until disinfectant residues are no longer discernible.

6 Check parts for visible soiling or damage. Check the sensor insert for damage to the measuring wires and their pegs.

7 Thoroughly shake out residual water. Allow the parts to dry completely.

Machine cleaning with thermal disinfection:

**CAUTION**
Only carry out machine cleaning and disinfection on the housing.

Sterilization:
Only sterilize the flow sensor when it is assembled.

**CAUTION**
Risk of patient injury due to failure of the flow measurement
Improper sterilization may damage the flow sensor. Only use the specified sterilization procedures.
Reprocessing

After reprocessing

Assembling the components

Assembling expiratory valve

1. Make sure all parts of the expiratory valve are completely dry, otherwise this may impair proper functioning.

2. Fit the muffler (B) to the expiratory valve.

3. Fit the diaphragm (A) onto the edge of the expiratory valve housing.

4. Fit the water trap container (C).

Inserting the expiratory valve into Babylog VN500

Prerequisite: The flap on the front is pivoted upwards.

1. Turn the locking ring (D) as far as possible to the left.

2. Push the expiratory valve into the fitting.

3. Turn the locking ring (D) as far as it will go to the right until it clicks audibly into place.

4. Check that it is properly secured by gently pulling on the expiratory valve.

5. Close the flap.

Further information on the expiratory valve:
The expiratory valve can be reused as long as the test point in the device check is passed. Exchange the expiratory valve if signs of wear become visible, such as cracks in the plastic parts, deformation and hardening of the rubber parts. Discolorations of the metal insert do not impair its function.
Reprocessing

Assembling the inspiratory unit

1. Make sure the inspiratory unit and diaphragm are completely dry, otherwise this may impair proper functioning.

2. Insert the adapter (A) of the diaphragm into the opening of the fitting (B). The adapter must be able to slightly move up and down in the opening.

3. Position the diaphragm in such a way that it is in the recesses (C) of the fitting.

4. Fit the diaphragm onto the edge of the fitting (D).

Inserting the inspiratory unit into Babylog VN500

1. Insert the inspiratory unit (E) into the recesses of the fitting and push as far as it will go into the fitting.

2. Turn the inspiratory unit in clockwise direction until the lock clicks into place.

3. Check whether the inspiratory unit is properly engaged.

Assembling accessories

Assemble the medication nebulizer and breathing gas humidifier in accordance with the corresponding instructions for use.

- Fit the medication nebulizer into the breathing circuit, see page 104.
- Prepare the breathing gas humidifier, see page 53.

Preparations before reuse

1. Assemble and prepare the device so that it is ready for use, see chapter "Assembly and preparation".

2. Check the operational readiness, see chapter "Getting started".

Instructions for use Infinity Acute Care System – Babylog VN500 SW 2.n 245
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Maintenance

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Overview

This chapter describes all maintenance steps necessary to maintain the proper functioning of the device. These measures must be performed by the personnel responsible.

Only perform maintenance work when no patient is connected to the device.

**CAUTION**

Disinfect and clean device or device parts before any maintenance measures and also before returning the medical device for repair.

**WARNING**

Risk of electric shock

Current-carrying components are located under the cover. Do not remove the cover. Maintenance work must be performed by service personnel or by experts. Dräger recommends that DrägerService performs these tasks.

---

**Definition of maintenance concepts**

<table>
<thead>
<tr>
<th>Concept</th>
<th>Definition</th>
</tr>
</thead>
<tbody>
<tr>
<td>Maintenance</td>
<td>Appropriate measures intended to retain the specified condition of a medical device</td>
</tr>
<tr>
<td>Inspection</td>
<td>Measures intended to determine and assess the actual state of a medical device</td>
</tr>
<tr>
<td>Preventive maintenance</td>
<td>Repeated indicated measures intended to retain the specified condition of a medical device</td>
</tr>
<tr>
<td>Repair</td>
<td>Measures intended to restore the functional condition of a medical device after the failure of a device function</td>
</tr>
</tbody>
</table>

---

**Inspection**

Inspections must be carried out regularly according to the following specifications and in the specified intervals.

<table>
<thead>
<tr>
<th>Checks</th>
<th>Interval</th>
<th>Personnel responsible</th>
</tr>
</thead>
<tbody>
<tr>
<td>Inspection and safety checks</td>
<td>Every 12 months</td>
<td>Service personnel</td>
</tr>
</tbody>
</table>

1) Designation applies to the Federal Republic of Germany; corresponds to the “Recurring safety inspection” in the Republic of Austria
Safety checks

The safety checks are no substitute for the preventive maintenance measures (including preventive replacement of wear parts) indicated by the manufacturer.

CAUTION
Perform safety checks at the indicated intervals. Otherwise, the correct functioning of the medical device can be impaired.

1 Check accompanying documents:
   – Instructions for use are available.

2 Perform a functional test of the following features:
   – Perform a device check and a breathing circuit check according to the instructions for use.
   – Perform a functional test of the airway pressure measurement.
   – Perform a functional test of the flow measurement.
   – Perform a functional test of the batteries (Babylog VN500 or PS500).

3 Verify that the device combination is in good condition:
   – Labels complete and legible.
   – No visible damage.
   – Fuses which are accessible from the outside are in compliance with the specified values.

4 Check that the equipment of the medical device is complete according to the instructions for use.

5 Check the electrical safety according to IEC 62353.

6 Check safety features:
   – Correct functioning of the emergency expiratory valve: Pressure rise 1.9 to 4.4 mbar (1.9 to 4.4 cmH2O) at a flow of 4.5 to 5.5 L/min.
   – Correct functioning of the non-return valves in the expiratory valve.
   – Correct functioning of the emergency breathing valve: Maximum pressure drop of 6.5 mbar (6.5 cmH2O) at a suction flow of 60 to 65 L/min.
   – Correct functioning of the alarm generator
   – Correct functioning of the non-return valves in the gas inlet for O2 and Air

Remote Service

From software release SW 2.20, Babylog VN500 supports the following Remote Service functionalities:
   – Help Ticket
   – Remote Device Check

Contact the responsible DrägerService representative for further information on the Remote Service function.

CAUTION
Perform safety checks at the indicated intervals. Otherwise, the correct functioning of the medical device can be impaired.
Maintenance

Preventive maintenance

**WARNING**
Risk due to defective components
Device failure is possible due to wear or material fatigue of the components. To maintain the function of all components, this device must be inspected and serviced at the intervals specified by the manufacturer.

<table>
<thead>
<tr>
<th>Component</th>
<th>Interval</th>
<th>Measure</th>
<th>Personnel responsible</th>
</tr>
</thead>
<tbody>
<tr>
<td>Ambient air filter</td>
<td>Every 4 weeks</td>
<td>Cleaning, replace if necessary, see page 251</td>
<td>User</td>
</tr>
<tr>
<td></td>
<td>Every 12 months</td>
<td>Replace, see page 251</td>
<td>User</td>
</tr>
<tr>
<td>Diaphragm of the expiratory valve</td>
<td>Every 12 months</td>
<td>Replace, see page 251</td>
<td>User</td>
</tr>
<tr>
<td>Expiratory valve</td>
<td>Every 2 years</td>
<td>Replace, see page 252</td>
<td>User</td>
</tr>
<tr>
<td>GS500: Breathing gas filter in the blower unit</td>
<td>Every 12 months</td>
<td>Replace, see page 254</td>
<td>Service personnel</td>
</tr>
<tr>
<td>GS500: Filter mat</td>
<td>Every 12 months</td>
<td>Replace, see page 253</td>
<td>Service personnel</td>
</tr>
<tr>
<td>Batteries</td>
<td>Every 3 months</td>
<td>Check capacity, see page 254</td>
<td>Service personnel</td>
</tr>
<tr>
<td></td>
<td></td>
<td>Replace if necessary</td>
<td>Experts</td>
</tr>
<tr>
<td></td>
<td>Every 2 years</td>
<td>Replace</td>
<td>Experts</td>
</tr>
<tr>
<td>Air filter (in the Air gas inlet)</td>
<td>Every 2 years</td>
<td>Replace</td>
<td>Experts</td>
</tr>
<tr>
<td>O2 filter (in the O2 gas inlet)</td>
<td>Every 6 years</td>
<td>Replace</td>
<td>Experts</td>
</tr>
</tbody>
</table>

Repair

Dräger recommends that all repairs are carried out by DrägerService and that only original Dräger parts are used.
Replacing the ambient air filter

**CAUTION**
Replace the ambient air filter at regular intervals. Otherwise operation of the device may be impaired.

Visually inspect the ambient air filter for contamination after 4 weeks; clean or replace if necessary. Replace after 12 months at the latest.

1. Unscrew the screw (A) on the cover of the ambient air filter.
2. Open the cover (B).
3. Remove the filter from the mount.
4. Fit a new filter or clean the old filter in warm soapy water and dry thoroughly.
5. Insert the filter into the mount without creasing.
6. Close the cover (B) and retighten the screw (A).
7. Dispose of used filter with domestic waste.

Replacing the diaphragm of the expiratory valve

Prerequisite: The expiratory valve has been removed, see "Removing the expiratory valve" on page 230.

1. Remove the diaphragm (A).
2. Fit the new diaphragm onto the edge of the expiratory valve housing. Make sure that the diaphragm is fitted properly.
3. Dispose of used diaphragm with domestic waste.
4. Fit the expiratory valve, see "Inserting the expiratory valve into the ventilation unit" on page 51.
Replacing the expiratory valve

1. Remove the expiratory valve, see page 230. Dispose of the expiratory valve in accordance with local waste disposal regulations.
2. Fit the expiratory valve, see page 244.
3. Insert the expiratory valve into the ventilation unit, see page 244.

Preventive maintenance on the GS500 gas supply unit

Replacing the breathing gas filter in the blower unit

The breathing gas filter is located behind the left-hand side panel of the GS500 gas supply unit.

1. Loosen the screws on the rear of the device sufficiently for the side panel to be removed. Remove the side panel. Ensure that the filter fleece attached to the side section is not loosened.
2. Take hold of the breathing gas filter by the handle and withdraw it from the GS500 gas supply unit. Dispose of the used breathing gas filter in accordance with local waste disposal regulations.
3. Insert the new breathing gas filter into the GS500 gas supply unit as far as it will go.
4. Fit the side panel and tighten the screws.
Replacing the filter mat

The filter mat is fastened to the inside of the left-hand side panel of the GS500 gas supply unit.

1. Loosen the screws on the rear of the device sufficiently for the side panel to be removed. Remove the side panel.
2. Remove the filter mat and dispose of it in accordance with local waste disposal regulations.
3. Fit the new filter mat with its side to the boundary. Carefully press the filter mat onto the pointed retaining elements. Check that the filter mat is secured.
4. Fit the side panel and tighten the screws.
Battery maintenance

Information on battery maintenance

The following actions are required to achieve the maximum life span of the batteries:

- Always fully charge the batteries.
- Connect the device to the mains power supply at the latest after 5 days to charge the batteries. Observe the required charging time.

If recharging is not possible after 5 days at the latest, do the following:

- Set the toggle switch to the position and then disconnect the power plug.

The device is then in the energy-saving mode and the discharge is reduced to the self-discharge of the batteries. Check that the capacity of the batteries is sufficient before use on a patient. The batteries may be exhausted or faulty as a result of excessively long storage.

Batteries are wear parts. The replacement intervals depend on the utilization. Observe the test intervals.

Storage at an increased ambient temperature reduces the life span of the batteries. The storage duration must not be exceeded. See chapter "Ambient conditions" on page 262.

The capacity of the batteries used must be checked regularly.

The batteries must have sufficient capacity. Replace the batteries if necessary.

Battery check

A battery check is required at regular intervals to determine the current state of the batteries. The battery check determines the approximate operating time.

The battery check consists of a charge-discharge-charge cycle. After the batteries have been fully charged, the device is operated in test mode with power supply from the batteries. The determined operating time is the approximate operating time to be expected in the next period of battery operation with typical ventilation without GS500.

Dräger recommends the following test intervals:

<table>
<thead>
<tr>
<th>Battery Type</th>
<th>Test Interval</th>
</tr>
</thead>
<tbody>
<tr>
<td>Internal battery (NiMH)</td>
<td>Every 3 months</td>
</tr>
<tr>
<td>PS500 power supply unit (VRLA)</td>
<td>Every 3 months</td>
</tr>
</tbody>
</table>

Prerequisites for the battery check

- The device is connected to the central gas supply.
- The device is connected to the mains power supply.
- The device is prepared and ready for use.
- The test lung is connected.
- A ventilation pattern is set, e.g.:
  - PC-AC
  - FiO2 = 21%
  - RR = 60/min
  - Pinsp = 20 mbar (or hPa or cmH2O)
  - PEEP = 5 mbar (or hPa or cmH2O)
The following table shows the typical operating time to be expected as a function of the ageing of a new battery. If the batteries do not correspond to the approximate operating time listed, replacement of the batteries is recommended.

<table>
<thead>
<tr>
<th>Age of the battery</th>
<th>Operating time of the internal battery (NiMH) with a full charge</th>
<th>Operating time of PS500 (VRLA) with a full charge</th>
</tr>
</thead>
<tbody>
<tr>
<td>3 months</td>
<td>29 min</td>
<td>225 min</td>
</tr>
<tr>
<td>6 months</td>
<td>28 min</td>
<td>210 min</td>
</tr>
<tr>
<td>9 months</td>
<td>27 min</td>
<td>195 min</td>
</tr>
<tr>
<td>12 months</td>
<td>26 min</td>
<td>180 min</td>
</tr>
<tr>
<td>15 months</td>
<td>25 min</td>
<td>165 min</td>
</tr>
<tr>
<td>18 months</td>
<td>24 min</td>
<td>150 min</td>
</tr>
<tr>
<td>21 months</td>
<td>23 min</td>
<td>135 min</td>
</tr>
<tr>
<td>24 months</td>
<td>22 min</td>
<td>120 min</td>
</tr>
</tbody>
</table>

NOTE
The operating time may be reduced due to the utilization of the battery. The data are approximate values and cannot be regarded as guaranteed for every battery.

NOTE
Replace the batteries if the operating time falls below the minimum value (see chapter "Battery ageing" on page 324) or after 24 months.
Battery check page

Prerequisite: The Start/Standby > System check dialog (A) window is opened.

1 Touch the Battery check tab (B).

The Battery check page contains the following:
- Battery check complete (C)
- Battery check PS500 (D)
- Battery check internal battery (E)

Information displayed in the (I) field for each battery:
- Date of the last battery check
- Determined operating time (value determined in the battery check during typical ventilation without GS500). See chapter "Battery check" on page 254.
- Next battery check due in xx days
- Battery replacement in xx months
- Current operating time
  This value is indicated to the nearest 5 or 10 minutes depending on the battery used and based on the present power consumption of the device.

Starting the battery check

The battery check can only be started if the device is connected to the mains power supply.

- Touch the Start button (F) and confirm. The appropriate battery check will be started. The result of the battery check is displayed after completion.

The duration of the battery check is decremented in hours and displayed in field (H).

If a battery check fails, the device will cancel the check. The canceled check is shown as a colorless dot.

Canceling the battery check

- Touch the Cancel button (G) and confirm. The appropriate battery check will be canceled. The canceled check is shown as a colorless dot.
Disposal

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Disposal of medical devices  . . . . . . . . . . . . . . . . 259
Safety information on disposal

**CAUTION**
The device and its components must be disinfect and cleaned before disposal!

For countries subject to EU Directive 2002/96/EC

This device is subject to EU Directive 2002/96/EC (WEEE). In order to comply with its registration according to this directive, this device may not be disposed of at municipal collection points for waste electrical equipment. Dräger has authorized a company to collect and dispose of this device. To initiate collection or for further information, visit Dräger on the Internet at www.draeger.com. Use the Search function with the keyword "WEEE" to find the relevant information. If access to Dräger's website is not possible, contact the local Dräger Organization.

Disposal of packaging material

Dispose of the packaging material of the device and the accessories listed in the list of accessories in accordance with the applicable laws and regulations.
Disposal

Disposal of batteries

The medical device contains batteries with toxic substances.
In the Federal Republic of Germany: The user is obliged by the ordinance on the return and disposal of used batteries to return batteries which contain toxic substances either to the manufacturer/sales outlet or to a collection center operated by public waste disposal corporations. The battery installed in the device must therefore be removed by experts before the device can be disposed of. Observe the applicable laws and regulations for battery disposal.

Disposal of neonatal flow sensor

The flow sensor must be disposed of as infectious waste. Low-emission combustion at over 800 °C (1472 °F).

Disposal of medical devices

At the end of its service life:

- Consult the relevant waste disposal company for appropriate disposal.
- Observe the applicable laws and regulations.
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## Technical data

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</tr>
</tbody>
</table>
Technical data

Ambient conditions

During operation

<table>
<thead>
<tr>
<th>Parameter</th>
<th>Requirement</th>
</tr>
</thead>
<tbody>
<tr>
<td>Temperature</td>
<td>10 to 40 °C (50 to 104 °F)</td>
</tr>
<tr>
<td>Pressure range</td>
<td>700 to 1060 hPa</td>
</tr>
<tr>
<td>Altitude</td>
<td>up to 3000 m (9842 ft)</td>
</tr>
<tr>
<td>Relative humidity</td>
<td>10 to 90 %, without condensation</td>
</tr>
</tbody>
</table>

During storage and transportation

<table>
<thead>
<tr>
<th>Parameter</th>
<th>Requirement</th>
</tr>
</thead>
<tbody>
<tr>
<td>Pressure range</td>
<td>500 to 1060 hPa</td>
</tr>
<tr>
<td>Relative humidity</td>
<td>5 to 95 %, non-condensing</td>
</tr>
</tbody>
</table>

Temperature

The technical specifications of the battery manufacturer regarding storage duration refer to a relative humidity of 45 to 85 %. Storage outside this range is possible. In this case, perform a battery test before using the device. Charging every 6 months at the latest is recommended. Several charge and discharge cycles may be required as a battery test in order to completely reactivate the electrochemical composition after long term storage.

Device without PS500 batteries for charging subsequent to storage

- For storage up to 6 months: −20 to <45 °C (−4 to <113 °F)
- For storage up to 1 month: −20 to <55 °C (−4 to <131 °F)
- For storage up to 1 week: −20 to 60 °C (−4 to 140 °F)

Device with PS500 batteries for charging subsequent to storage

- For storage up to 6 months: −15 to 25 °C (5 to 77 °F)
- For storage up to 3 months: −15 to 40 °C (5 to 104 °F)

Depending on the accessories used, more stringent ambient conditions can apply. Observe corresponding instructions for use.
Set values

The required parameters can be adjusted with the therapy controls of Babylog VN500 without any loss of accuracy. The controlled parameters pressure, flow, volume, and O2 concentration can only be applied with the accuracy of the associated measured values.

The accuracies indicated apply only under the following conditions:
- The device is ready for operation, see chapter “Getting started”.
- Any accessories being used are approved for the device, see the list of accessories.
- The type of humidification is selected correctly in the Start/Standby > Br. circuit/ Humidifier dialog window.

The tolerances do not include the measurement uncertainty of external test equipment. This information is available on request.

<table>
<thead>
<tr>
<th>Parameter</th>
<th>Value</th>
</tr>
</thead>
<tbody>
<tr>
<td>Respiratory rate $RR$</td>
<td>0.5 to 150/min</td>
</tr>
<tr>
<td>Inspiratory time $Ti$</td>
<td>0.1 to 3 s</td>
</tr>
<tr>
<td>Maximum inspiratory time for flow cycled breaths $Timax$</td>
<td></td>
</tr>
<tr>
<td>Pediatric patients</td>
<td>0.1 to 4 s</td>
</tr>
<tr>
<td>Neonates</td>
<td>0.1 to 1.5 s</td>
</tr>
<tr>
<td>Tidal volume $VT$</td>
<td></td>
</tr>
<tr>
<td>Pediatric patients</td>
<td>20 to 300 mL</td>
</tr>
<tr>
<td>Neonates</td>
<td>2 to 100 mL</td>
</tr>
<tr>
<td>Tidal volume for pressure support $VT$</td>
<td></td>
</tr>
<tr>
<td>Pediatric patients</td>
<td>20 to 300 mL</td>
</tr>
<tr>
<td>Neonates</td>
<td>2 to 100 mL</td>
</tr>
<tr>
<td>Upper limit of tidal volume for PPS $VT_{max}$</td>
<td></td>
</tr>
<tr>
<td>Pediatric patients</td>
<td>20 to 300 mL</td>
</tr>
<tr>
<td>Neonates</td>
<td>2 to 100 mL</td>
</tr>
<tr>
<td>Activation state of Apnea Ventilation</td>
<td>on, off</td>
</tr>
<tr>
<td>Status of the Automatic return from Apnea Ventilation function</td>
<td>on, off</td>
</tr>
</tbody>
</table>
### Set values (cont.)

<table>
<thead>
<tr>
<th>Parameter</th>
<th>Value</th>
</tr>
</thead>
<tbody>
<tr>
<td>Tidal volume during Apnea Ventilation</td>
<td><strong>VTapn</strong></td>
</tr>
<tr>
<td>Pediatric patients</td>
<td>20 to 300 mL</td>
</tr>
<tr>
<td>Neonates</td>
<td>2 to 100 mL</td>
</tr>
<tr>
<td>Respiratory rate during apnea ventilation <strong>RRapn</strong></td>
<td>2 to 150/min</td>
</tr>
<tr>
<td>Inspiratory flow</td>
<td><strong>Flow</strong></td>
</tr>
<tr>
<td>Pediatric patients</td>
<td>2 to 30 L/min</td>
</tr>
<tr>
<td>Neonates</td>
<td>2 to 30 L/min</td>
</tr>
<tr>
<td>Maximum inspiratory flow in NIV mode for neonates <strong>Flow max</strong></td>
<td>0 to 30 L/min</td>
</tr>
<tr>
<td>Inspiratory pressure <strong>Pinsp</strong></td>
<td>1 to 80 mbar (or hPa or cmH₂O)</td>
</tr>
<tr>
<td>Inspiratory pressure limit <strong>Pmax</strong></td>
<td>2 to 100 mbar (or hPa or cmH₂O)</td>
</tr>
<tr>
<td>O₂ concentration <strong>Fio₂</strong></td>
<td>21 to 100 Vol%</td>
</tr>
<tr>
<td>T₀...90</td>
<td>test conditions in accordance with ISO 80601-2-12:2011, Sec. 201.12.1.104</td>
</tr>
<tr>
<td>Time until the adjusted O₂ value reaches the patient connection</td>
<td>taking account of the airway-conducting accessories with the greatest internal volume; with flow monitoring switched on</td>
</tr>
<tr>
<td>Pediatric patients and neonates</td>
<td>&lt;20 s</td>
</tr>
<tr>
<td>Positive end-expiratory pressure <strong>PEEP</strong></td>
<td>0 to 35 mbar (or hPa or cmH₂O)</td>
</tr>
<tr>
<td>Trigger sensitivity <strong>Flow trigger</strong></td>
<td>0.2 to 5 L/min</td>
</tr>
<tr>
<td>Pressure support <strong>Psupp</strong></td>
<td>0 to 80 mbar (or hPa or cmH₂O)</td>
</tr>
<tr>
<td>Rise time for pressure support</td>
<td><strong>Slope</strong></td>
</tr>
<tr>
<td>Pediatric patients</td>
<td>0 to 2 s</td>
</tr>
<tr>
<td>Neonates</td>
<td>0 to 1.5 s</td>
</tr>
<tr>
<td>Airway Pressure Release Ventilation <strong>APRV</strong></td>
<td><strong>Thigh</strong></td>
</tr>
<tr>
<td>Inspiratory time</td>
<td>0.1 to 30 s</td>
</tr>
<tr>
<td>Expiratory time <strong>Tlow</strong></td>
<td>0.05 to 30 s</td>
</tr>
<tr>
<td>Maximum time of low pressure level in APRV/PEF <strong>Tlow max</strong></td>
<td>0.05 to 30 s</td>
</tr>
<tr>
<td>Inspiratory pressure <strong>Phigh</strong></td>
<td>1 to 80 mbar (or hPa or cmH₂O)</td>
</tr>
<tr>
<td>Expiratory pressure <strong>Plow</strong></td>
<td>0 to 35 mbar (or hPa or cmH₂O)</td>
</tr>
<tr>
<td>Termination criterion (peak expiratory flow) <strong>Exp. term.</strong></td>
<td>1 to 80 %<strong>PEF</strong></td>
</tr>
</tbody>
</table>
Technical data

Set values (cont.)

<table>
<thead>
<tr>
<th>Automatic Tube Compensation</th>
<th>ATC</th>
</tr>
</thead>
<tbody>
<tr>
<td>Inner diameter of the tube</td>
<td>Tube Ø</td>
</tr>
<tr>
<td>Endotracheal tube</td>
<td>ET</td>
</tr>
<tr>
<td>Pediatric patients</td>
<td>2 to 8 mm (0.08 to 0.31 in)</td>
</tr>
<tr>
<td>Neonates</td>
<td>2 to 5 mm (0.08 to 0.2 in)</td>
</tr>
<tr>
<td>Tracheostomy tube</td>
<td>Trach.</td>
</tr>
<tr>
<td>Pediatric patients</td>
<td>2.5 to 8 mm (0.1 to 0.31 in)</td>
</tr>
</tbody>
</table>

Degree of tube compensation **Compens.** 0 to 100 %

Status of ATC during mandatory inspiration **Inspiratory compensation** on/off

Status of ATC during expiratory phase **Expiratory compensation** on/off

Proportional Pressure Support **PPS**

Flow Assist **Flow Assist**

| Pediatric patients | 0 to 100 mbar/L/s (or hPa/L/s or cmH2O/L/s) |
| Neonates           | 0 to 300 mbar/L/s (or hPa/L/s or cmH2O/L/s) |

Volume Assist **Flow Assist**

| Pediatric patients | 0 to 1000 mbar/L (or hPa/L or cmH2O/L) |
| Neonates           | 0 to 4000 mbar/L (or hPa/L or cmH2O/L) |

Volume Assist corresponds to compliance compensation

| Pediatric patients | 10000 to 1 mL/mbar (or mL/hPa or mL/cmH2O) |
| Neonates           | 1000 to 0.3 mL/mbar (or mL/hPa or mL/cmH2O) |

O2 Therapy

Continuous Flow **Flow** (BTPS) 2 to 50 L/min

O2 concentration **FiO2** 21 to 100 Vol%

High Frequency Oscillation **HFO**

Mean airway pressure during HFO **MAPhf** 5 to 50 mbar (or hPa or cmH2O)

Frequency of oscillation in HFO **fhf** 5 to 20 Hz

I to E in HFO **I:Ehf** 1:1 to 1:3

Pressure amplitude in HFO **Ampl hf** 5 to 90 mbar (or hPa or cmH2O)

Maximum pressure amplitude in HFO (VG) **Ampl hf max** 5 to 90 mbar (or hPa or cmH2O)

Tidal volume in HFO **VThf** 0.2 to 40 mL

Sigh pressure in HFO **Psigh** 6 to 80 mbar (or hPa or cmH2O)

Respiratory rate of sigh in HFO **RRsigh** 0 to 30/min
Technical data

Set values (cont.)

<table>
<thead>
<tr>
<th>Sigh pressure rise time</th>
<th>Slopesigh</th>
</tr>
</thead>
<tbody>
<tr>
<td>Pediatric patients</td>
<td>0 to 2 s</td>
</tr>
<tr>
<td>Neonates</td>
<td>0 to 1.5 s</td>
</tr>
<tr>
<td>Sigh inspiratory time in HFO</td>
<td>0.1 to 3 s</td>
</tr>
</tbody>
</table>

Leakage compensation
- on/off
  - on: full compensation active
  - off: only trigger compensation active

Maneuver settings
- Sigh pressure \(\Delta p\text{PEEP}\)
  - 0 to 20 mbar (or hPa or cmH2O)
- Time interval between sighs \(\text{Interval sigh}\)
  - 20 s to 180 min
- Number of cycles for a sigh \(\text{Cycles sigh}\)
  - 1 to 20 exhalations
- Oxygen enrichment for suction maneuver
  - Factor for neonates: 1 to 2
  - Factor for pediatric patients: 1 to 2

Performance characteristics

<table>
<thead>
<tr>
<th>Control principle</th>
<th>time-cycled, volume-constant, pressure-controlled</th>
</tr>
</thead>
<tbody>
<tr>
<td>Intermittent PEEP duration</td>
<td>1 to 20 expiratory cycles</td>
</tr>
<tr>
<td>Medication nebulization</td>
<td>For 5, 10, 15, 30 minutes, continuously ((\times))</td>
</tr>
</tbody>
</table>

Endotracheal suction
- Disconnection detection: automatic
- Reconnection detection: automatic
- Initial oxygen enrichment: max. 3 minutes
- Active suction phase: max. 2 minutes
- Final oxygen enrichment: max. 2 minutes
## Performance characteristics (cont.)

<table>
<thead>
<tr>
<th>Supply system for spontaneous breathing and adaptive CPAP system with high initial flow</th>
</tr>
</thead>
<tbody>
<tr>
<td>Inspiratory flow (BTPS) for pediatric patients (if proximal flow measurement is used, the measuring range is restricted, see section on displayed flow values, page 270) max. 60 L/min</td>
</tr>
<tr>
<td>Inspiratory flow (BTPS) for neonates max. 30 L/min</td>
</tr>
<tr>
<td>Base flow, neonates 6 L/min</td>
</tr>
<tr>
<td>Base flow, pediatric patients 3 L/min</td>
</tr>
<tr>
<td>Base flow, pediatric patients, with active pneumatic nebulization 6 L/min</td>
</tr>
</tbody>
</table>

<table>
<thead>
<tr>
<th>Supply system for HFO Inspiratory flow (BTPS) max. 120 L/min</th>
</tr>
</thead>
<tbody>
<tr>
<td>Base flow 25 L/min</td>
</tr>
</tbody>
</table>

Minimum short-term negative pressure during HFO –40 mbar (or hPa or cmH₂O)

In a combination of proximal flow sensor with filters, HME, Ergostar and CO₂ cuvette, the airway resistance of the system can be more than 6 mbar (or hPa or cmH₂O) if there is a failure of the device and a flow of 15 L/min.

### Inspiratory resistance on device failure

<table>
<thead>
<tr>
<th>Pediatric patients, maximum value</th>
<th>&lt;6 mbar at 15 L/min</th>
</tr>
</thead>
<tbody>
<tr>
<td></td>
<td>&lt;6 hPa at 15 L/min</td>
</tr>
<tr>
<td></td>
<td>&lt;6 cmH₂O at 15 L/min</td>
</tr>
<tr>
<td>Neonates, maximum value</td>
<td>&lt;1.5 mbar at 2.5 L/min</td>
</tr>
<tr>
<td></td>
<td>&lt;1.5 hPa at 2.5 L/min</td>
</tr>
<tr>
<td></td>
<td>&lt;1.5 cmH₂O at 2.5 L/min</td>
</tr>
</tbody>
</table>

### Expiratory resistance on device failure

<table>
<thead>
<tr>
<th>Pediatric patients, maximum value</th>
<th>&lt;6.0 mbar at 15 L/min</th>
</tr>
</thead>
<tbody>
<tr>
<td></td>
<td>&lt;6.0 hPa at 15 L/min</td>
</tr>
<tr>
<td></td>
<td>&lt;6.0 cmH₂O at 15 L/min</td>
</tr>
<tr>
<td>Neonates, maximum value</td>
<td>&lt;1.0 mbar at 2.5 L/min</td>
</tr>
<tr>
<td></td>
<td>&lt;1.0 hPa at 2.5 L/min</td>
</tr>
<tr>
<td></td>
<td>&lt;1.0 cmH₂O at 2.5 L/min</td>
</tr>
</tbody>
</table>
**Performance characteristics (cont.)**

**Accuracy of measured values**
Depending on the patient category, the accuracies indicated for the measured values apply to the following performance characteristics of the breathing circuit.

<table>
<thead>
<tr>
<th>Breathing circuit for pediatric patients including additional components</th>
</tr>
</thead>
<tbody>
<tr>
<td>Compliance</td>
</tr>
<tr>
<td>Inspiratory resistance</td>
</tr>
<tr>
<td></td>
</tr>
<tr>
<td></td>
</tr>
<tr>
<td>Expiratory resistance</td>
</tr>
<tr>
<td></td>
</tr>
<tr>
<td></td>
</tr>
</tbody>
</table>

<table>
<thead>
<tr>
<th>Breathing circuit for neonates including additional components</th>
</tr>
</thead>
<tbody>
<tr>
<td>Compliance</td>
</tr>
<tr>
<td>Inspiratory resistance</td>
</tr>
<tr>
<td></td>
</tr>
<tr>
<td></td>
</tr>
<tr>
<td>Expiratory resistance</td>
</tr>
<tr>
<td></td>
</tr>
<tr>
<td></td>
</tr>
</tbody>
</table>

<table>
<thead>
<tr>
<th>Compliance of device incl. breathing circuit</th>
</tr>
</thead>
<tbody>
<tr>
<td>Neonates, maximum value</td>
</tr>
<tr>
<td></td>
</tr>
<tr>
<td></td>
</tr>
</tbody>
</table>

**Additional functions**

<table>
<thead>
<tr>
<th>Safety valve</th>
</tr>
</thead>
<tbody>
<tr>
<td>Opens if medical compressed air supply fails (supply gas flow is not sufficient to provide the inspiratory flow required), enables spontaneous breathing with ambient air.</td>
</tr>
</tbody>
</table>
### Technical data

#### Displayed measured values

Accuracy does only apply for the measurement range specified.

**Airway pressure measurement**
- Positive end-expiratory pressure \( P_{EEEP} \)
- Peak Inspiratory Pressure \( P_{IP} \)
- Mean airway pressure \( P_{mean} \)
- Minimum airway pressure \( P_{min} \)
- Lower pressure level in APRV \( P_{low} \)
- End-inspiratory pressure for mandatory breaths \( EIP \)
- Upper pressure level in APRV \( P_{high} \)

**Range**
- within the setting range of 0 to a maximum of 80 mbar (or hPa or cmH2O) (within the maximum sensor measuring range of –60 to 120 mbar (or hPa or cmH2O))

**Accuracy**
- in phases without flow:
  - ±6 % of measured value or ±0.5 mbar (or hPa or cmH2O), whichever is greater
- otherwise:
  - ±2 mbar (or hPa or cmH2O)

**T0...90 (for Pmean)**
- 20 s for pediatric patients,
- 10 s for neonates

**Airway pressure measurement during HFO**
- Pressure amplitude (peak-to-peak) in HFO \( \Delta P_{hf} \)
  - Range 0 to 90 mbar (or hPa or cmH2O)
  - Accuracy ±20 % of measured value, or ±5.5 mbar (or hPa or cmH2O), whichever is greater, after the breathing circuit has been calibrated

**Mean airway pressure during HFO**
- Range 0 to 50 mbar (or hPa or cmH2O)
  - Accuracy ±15 % of measured value, or ±1.9 mbar (or hPa or cmH2O), whichever is greater, with \( Ampl_{hf} <50 \) mbar (or hPa or cmH2O) after the breathing circuit has been calibrated
  - Accuracy ±20 % of measured value, or ±2.9 mbar (or hPa or cmH2O), whichever is greater, with \( Ampl_{hf} \geq 50 \) mbar (or hPa or cmH2O) after the breathing circuit has been calibrated
  - 20 ms
**Technical data**

**Displayed measured values (cont.)**

<table>
<thead>
<tr>
<th>O₂ measurement (inspiratory side)</th>
<th></th>
</tr>
</thead>
<tbody>
<tr>
<td>Inspiratory O₂ concentration (in dry air)</td>
<td>FiO₂</td>
</tr>
<tr>
<td>Range</td>
<td>18 to 100 Vol%</td>
</tr>
<tr>
<td>Accuracy</td>
<td>±3 Vol% at 20 °C (68 °F)</td>
</tr>
<tr>
<td>Drift of measurement accuracy</td>
<td>0.2 Vol% in 6 hours (corresponding to ISO 21647, ISO 80601-2-55).</td>
</tr>
<tr>
<td>The measured values of the O₂ measurement are barometrically pressure compensated.</td>
<td></td>
</tr>
<tr>
<td>Rise time T₀...₉₀</td>
<td>500 ms</td>
</tr>
<tr>
<td>Warm-up time</td>
<td>max. 3 minutes, typ. 1 minute</td>
</tr>
</tbody>
</table>

<table>
<thead>
<tr>
<th>Flow measurement (proximal)</th>
<th></th>
</tr>
</thead>
<tbody>
<tr>
<td>Minute volume measurement</td>
<td></td>
</tr>
<tr>
<td>Expiratory minute volume</td>
<td>MVₑ</td>
</tr>
<tr>
<td>Inspiratory minute volume</td>
<td>MVᵢ</td>
</tr>
<tr>
<td>Mandatory expiratory minute volume</td>
<td>MVₑₘₐⁿᵈ</td>
</tr>
<tr>
<td>Spontaneous expiratory minute volume</td>
<td>MVₑₛᵖₒⁿ</td>
</tr>
<tr>
<td>Minute volume, leakage-compensated</td>
<td>MV</td>
</tr>
<tr>
<td>Range</td>
<td>0 to 30 L/min, BTPS</td>
</tr>
<tr>
<td>Accuracy</td>
<td>Measured with neonatal flow sensor: ±10 % of measured value or ±0.6 mL * (RR + 2), whichever is greater under calibration conditions during device check (at 1013 mbar (1013 cmH₂O), gas with 50 % rel. humidity, 23 °C (73.4 °F)), no leakage and using a Dräger Y-piece, no HFO</td>
</tr>
<tr>
<td>T₀...₉₀</td>
<td></td>
</tr>
<tr>
<td>Pediatric patients</td>
<td>33 s</td>
</tr>
<tr>
<td>Neonates</td>
<td>20 s</td>
</tr>
</tbody>
</table>

<table>
<thead>
<tr>
<th>Tidal volume measurement</th>
<th></th>
</tr>
</thead>
<tbody>
<tr>
<td>Tidal volume</td>
<td>VT</td>
</tr>
<tr>
<td>Inspiratory tidal volume (not leakage-compensated) of mandatory breaths</td>
<td>VTₑₘₐⁿᵈ</td>
</tr>
<tr>
<td>Expiratory tidal volume (not leakage-compensated) of mandatory breaths</td>
<td>VTₑₘₐⁿᵈ</td>
</tr>
<tr>
<td>Inspiratory tidal volume (not leakage-compensated) of spontaneous breaths</td>
<td>VTᵢₛᵖₒⁿ</td>
</tr>
<tr>
<td>Range</td>
<td>0 to 1000 mL, BTPS</td>
</tr>
</tbody>
</table>
Displayed measured values (cont.)

<table>
<thead>
<tr>
<th>Parameter</th>
<th>Value</th>
</tr>
</thead>
<tbody>
<tr>
<td>Accuracy</td>
<td>Measured with neonatal flow sensor: ±10 % of measured value or ±0.6 mL, whichever is greater, under calibration conditions during device check (at 1013 mbar (1013 cmH2O), gas with 50 % rel. humidity, 23 °C (73.4 °F)), no leakage and using a Dräger Y-piece, no HFO</td>
</tr>
<tr>
<td>Device Flow (inspiratory, only with HFO)</td>
<td>Range 20 to 40 L/min, BTPS</td>
</tr>
<tr>
<td></td>
<td>Accuracy ±10 %</td>
</tr>
<tr>
<td>Respiratory rate measurement</td>
<td>RR</td>
</tr>
<tr>
<td>Mandatory respiratory rate</td>
<td>RRmand</td>
</tr>
<tr>
<td>Portion of mandatory triggered breaths</td>
<td>RRtrig</td>
</tr>
<tr>
<td>Spontaneous respiratory rate</td>
<td>RRspon</td>
</tr>
<tr>
<td>Range</td>
<td>0 to 300/min</td>
</tr>
<tr>
<td>Accuracy</td>
<td>±1/min for respiratory rates ≥2/min and ±2/min for respiratory rates &lt;2/min</td>
</tr>
<tr>
<td>T0...90</td>
<td>33 s</td>
</tr>
<tr>
<td>Effective inspiratory time during spontaneous breathing <strong>Tispon</strong></td>
<td>0 to 20 s</td>
</tr>
<tr>
<td>Effective expiratory time, only if additional setting <strong>AutoRelease</strong> is active <strong>Tlow</strong></td>
<td>0 to 20 s</td>
</tr>
<tr>
<td>Inspiratory time to expiratory time ratio for mandatory ventilation <strong>I:E</strong></td>
<td>1:300 to 600:1</td>
</tr>
<tr>
<td>Inspiratory time to expiratory time ratio for spontaneous breathing <strong>I:Espon</strong></td>
<td>1:300 to 600:1</td>
</tr>
<tr>
<td>CO2 measurement in mainstream</td>
<td>etCO2</td>
</tr>
<tr>
<td>Range</td>
<td>Range 0 to 100 mmHg or 0 to 13.2 Vol% (at 1013 mbar (1013 cmH2O)) or 0 to 13.3 kPa</td>
</tr>
<tr>
<td>Accuracy</td>
<td>±2.0 mmHg in the range 0 to 40 mmHg, ±5 % of the measured value in the range 41 to 100 mmHg ±0.27 kPa in the range 0 to 5.33 kPa, ±5 % of the measured value in the range 5.34 to 13.3 kPa ±0.26 Vol% in the range 0 to 5.26 Vol%, ±5 % of the measured value in the range 5.27 to 13.2 Vol%</td>
</tr>
</tbody>
</table>
**Technical data**

### Displayed measured values (cont.)

<table>
<thead>
<tr>
<th>Measurement conditions</th>
<th>Respiratory rate (pediatric patients):</th>
</tr>
</thead>
<tbody>
<tr>
<td></td>
<td>40 to 100/min</td>
</tr>
<tr>
<td></td>
<td>Inspiratory time: &gt;250 ms</td>
</tr>
<tr>
<td></td>
<td>Expiratory time: &gt;250 ms</td>
</tr>
</tbody>
</table>

**Drift of measurement accuracy**

The measured values of the CO₂ measurement are barometrically pressure compensated.

T₁₀...₉₀ < 35 ms

Total response time < 200 ms

Warm-up time, typical < 3 min (at 23 °C)

With reference to the displayed measured values, the following dead space volumes must be taken into account:

- **CO₂ cuvette, pediatric patients (6870280, MP01063)**: 1.9 mL
- **Neonatal flow sensor ISO 15 (8411130)**: 0.9 mL
- **Neonatal flow sensor Y-piece (8410185)**: 1.7 mL

### Displayed calculated values

- **Dynamic compliance** $C_{dyn}$
  - Range: 0 to 650 mL/mbar (mL/cm H₂O)
- **Resistance** $R$
  - Range: 0 to 1000 mbar/(L/s) (or hPa/(L/s) or cm H₂O/(L/s))
- **Patient resistance** $R_{pat}$
  - Range: 0 to 1000 mbar/(L/s) (or hPa/(L/s) or cm H₂O/(L/s))
- **Leakage minute volume** $MV_{leak}$
  - Range: 0 to 30 L/min, BTPS
  - Accuracy: ±10 % from measured value
    - T₀...₉₀: 20 s for pediatric patients,
    - 10 s for neonates
- **Leakage in % % leak**
  - Range: 0 to 100 %
- **Spontaneous breathing portion of minute volume in percent %MV_{spon}**
  - Range: 0 to 100 %
Technical data

Displayed calculated values (cont.)

<table>
<thead>
<tr>
<th>Rapid Shallow Breathing</th>
<th>RSB</th>
</tr>
</thead>
<tbody>
<tr>
<td>Range</td>
<td></td>
</tr>
<tr>
<td>Pediatric patients</td>
<td>0 to 9999 (/min/L)</td>
</tr>
<tr>
<td>Neonates</td>
<td>0 to 300 (/min/mL)</td>
</tr>
<tr>
<td>For accuracy, see measurement of VT and RR</td>
<td></td>
</tr>
<tr>
<td>Time constant of expiration T&lt;sub&gt;ce&lt;/sub&gt;</td>
<td>0 to 20 s</td>
</tr>
</tbody>
</table>

Curve displays
- Airway pressure Paw (t): −30 to 100 mbar (or hPa or cmH<sub>2</sub>O)
- Flow (t): −40 to 40 L/min
- Volume V (t): 2 to 300 mL
- CO<sub>2</sub> (t): 0 to 100 mmHg or 0 to 13.2 Vol% (at 1013 mbar (1013 cmH<sub>2</sub>O)) or 0 to 13.3 kPa

Monitoring

Alarm sound pressure level L(A) at operator's position:
- Operator's position: at front of device at a distance of 1 m (39 in) and a height of 1.5 m (59 in).
- Free field measurement in accordance with ISO 3744 and IEC 60601-1-8:2003

Alarm tone sequence IEC/CEI
- Range for high-priority alarms: from about 52 dB(A) to 70 dB(A)
- Range for medium-priority alarms: from about 49 dB(A) to 67 dB(A)
- Range for low-priority alarms: from about 46 dB(A) to 65 dB(A)
- Incrementation: adjustable in 9 increments

Alarm tone sequence Dräger ventilation
- Range for high-priority alarms: from about 55 dB(A) to 73 dB(A)
- Range for medium-priority alarms: from about 51 dB(A) to 70 dB(A)
- Range for low-priority alarms: from about 47 dB(A) to 65 dB(A)
- Incrementation: adjustable in 9 increments

Alarm sound pressure level for power failure alarm and auxiliary alarm: from about 70 dB(A) to 75 dB(A)
### Sound pressure level LPA of alarm signals measured in accordance with IEC 60601-1-8 and A1:2012:

<table>
<thead>
<tr>
<th>Alarm tone sequence</th>
<th>Range for high-priority alarms according to volume setting</th>
<th>approx. 55 dB(A) to 72 dB(A)</th>
</tr>
</thead>
<tbody>
<tr>
<td></td>
<td>Range for medium-priority alarms according to volume setting</td>
<td>approx. 52 dB(A) to 69 dB(A)</td>
</tr>
<tr>
<td></td>
<td>Range for low-priority alarms according to volume setting</td>
<td>approx. 49 dB(A) to 67 dB(A)</td>
</tr>
<tr>
<td></td>
<td>Incrementation</td>
<td>adjustable in 9 increments</td>
</tr>
</tbody>
</table>

<table>
<thead>
<tr>
<th>Alarm tone sequence</th>
<th>Range for high-priority alarms according to volume setting</th>
<th>approx. 55 dB(A) to 72 dB(A)</th>
</tr>
</thead>
<tbody>
<tr>
<td></td>
<td>Range for medium-priority alarms according to volume setting</td>
<td>approx. 53 dB(A) to 70 dB(A)</td>
</tr>
<tr>
<td></td>
<td>Range for medium-priority alarms according to volume setting</td>
<td>approx. 45 dB(A) to 62 dB(A)</td>
</tr>
<tr>
<td></td>
<td>Incrementation</td>
<td>adjustable in 9 increments</td>
</tr>
<tr>
<td></td>
<td>Range for power supply failure alarm and auxiliary alarm</td>
<td>approx. 70 dB(A) to 75 dB(A)</td>
</tr>
</tbody>
</table>

| Delay time to sounding of auxiliary alarm if main alarm has failed | max. 18 s |

<table>
<thead>
<tr>
<th>Expiratory minute volume</th>
<th>MVe</th>
</tr>
</thead>
<tbody>
<tr>
<td>Upper alarm limit alarm</td>
<td>if the upper alarm limit has been exceeded</td>
</tr>
<tr>
<td>Setting range in invasive ventilation</td>
<td>0.03 to 41 L/min</td>
</tr>
<tr>
<td>Setting range in non-invasive ventilation</td>
<td>0.03 to 60 L/min</td>
</tr>
<tr>
<td>Alarm delay</td>
<td>MV delay</td>
</tr>
<tr>
<td>Pediatric patients</td>
<td>0 to 20 s</td>
</tr>
<tr>
<td>Neonates</td>
<td>0 to 15 s</td>
</tr>
</tbody>
</table>

| Lower alarm limit alarm | if the value has fallen below the lower alarm limit |
| Setting range | 0.02 to 40 L/min, Off (with NIV or neonates) |
| Alarm suppression | 2 minutes after leaving standby |
|                     | during and 2 minutes after suction maneuver |
|                     | 2 minutes after switching on flow monitoring |

<table>
<thead>
<tr>
<th>Alarm delay</th>
<th>MV delay</th>
</tr>
</thead>
<tbody>
<tr>
<td>Pediatric patients</td>
<td>0 to 20 s</td>
</tr>
<tr>
<td>Neonates</td>
<td>0 to 15 s</td>
</tr>
</tbody>
</table>
### Technical data

#### Monitoring (cont.)

<table>
<thead>
<tr>
<th>Parameter</th>
<th>Setting</th>
<th>Description</th>
</tr>
</thead>
<tbody>
<tr>
<td><strong>Airway pressure</strong></td>
<td></td>
<td></td>
</tr>
<tr>
<td>Upper alarm limit alarm</td>
<td></td>
<td>if the upper alarm limit has been exceeded</td>
</tr>
<tr>
<td>Setting range</td>
<td></td>
<td>7 to 105 mbar (or hPa or cmH2O)</td>
</tr>
<tr>
<td>Maximum airway pressure</td>
<td></td>
<td>120 mbar (or hPa or cmH2O)</td>
</tr>
<tr>
<td>Inspiratory O2 concentration</td>
<td></td>
<td></td>
</tr>
<tr>
<td>Upper alarm limit alarm</td>
<td></td>
<td>after 30 seconds at the latest, if the upper alarm limit has been continuously exceeded</td>
</tr>
<tr>
<td>Setting range</td>
<td></td>
<td></td>
</tr>
<tr>
<td>Lower alarm limit alarm</td>
<td></td>
<td>after 30 seconds at the latest, if the lower alarm limit has been continuously undershot</td>
</tr>
<tr>
<td></td>
<td></td>
<td>both alarm limits are automatically assigned to the set value: under 60 Vol% with ±4 Vol%, from 60 Vol% with ±6 Vol% (lower alarm limit 18 Vol% at 21 Vol%)</td>
</tr>
<tr>
<td><strong>End-expiratory CO2 concentration</strong></td>
<td></td>
<td></td>
</tr>
<tr>
<td>Upper alarm limit alarm</td>
<td></td>
<td>if the upper alarm limit has been exceeded</td>
</tr>
<tr>
<td>Setting range</td>
<td></td>
<td>1 to 98 mmHg (or 0.1 to 13.1 Vol% or 0.1 to 13.3 kPa)</td>
</tr>
<tr>
<td>Lower alarm limit alarm</td>
<td></td>
<td>if the value has fallen below the lower alarm limit</td>
</tr>
<tr>
<td>Setting range</td>
<td></td>
<td>0 to 97 mmHg (or 0 to 13.0 Vol% or 0 to 13.2 kPa)</td>
</tr>
<tr>
<td><strong>Respiratory rate</strong></td>
<td></td>
<td></td>
</tr>
<tr>
<td>Upper alarm limit alarm</td>
<td></td>
<td>if the respiratory rate (mandatory and spontaneous breaths) has been exceeded</td>
</tr>
<tr>
<td>Setting range</td>
<td></td>
<td>5 to 200/min, Off</td>
</tr>
<tr>
<td><strong>Volume monitoring</strong></td>
<td></td>
<td></td>
</tr>
<tr>
<td>Lower alarm limit alarm</td>
<td></td>
<td>if the set tidal volume has not been supplied</td>
</tr>
<tr>
<td>Setting range</td>
<td></td>
<td>90 % of VT set (only for modes with VG)</td>
</tr>
<tr>
<td>Alarm suppression</td>
<td></td>
<td></td>
</tr>
<tr>
<td>Pediatric patients</td>
<td></td>
<td>during the first five consecutive breaths where the applied inspiratory tidal volume has fallen below the lower alarm limit</td>
</tr>
<tr>
<td>Neonates</td>
<td></td>
<td>during the first eight consecutive breaths where the applied inspiratory tidal volume has fallen below the lower alarm limit</td>
</tr>
<tr>
<td><strong>Apnea alarm time</strong></td>
<td></td>
<td></td>
</tr>
<tr>
<td>Alarm</td>
<td></td>
<td>if no breathing activity is detected</td>
</tr>
<tr>
<td>Setting range</td>
<td></td>
<td>5 to 60 seconds, Off</td>
</tr>
<tr>
<td>Disconnect alarm delay time</td>
<td></td>
<td></td>
</tr>
<tr>
<td>Setting range</td>
<td></td>
<td>0 to 60 seconds</td>
</tr>
</tbody>
</table>
### Technical data

### Operating data

<table>
<thead>
<tr>
<th>Protection class</th>
<th></th>
</tr>
</thead>
<tbody>
<tr>
<td>Ventilation unit Babylog VN500</td>
<td>Class I</td>
</tr>
<tr>
<td>Medical Cockpit Infinity C500</td>
<td></td>
</tr>
<tr>
<td>Gas supply unit GS500</td>
<td></td>
</tr>
<tr>
<td>Power supply unit PS500</td>
<td></td>
</tr>
<tr>
<td>CO&lt;sub&gt;2&lt;/sub&gt; sensor (sensor connected)</td>
<td>Type BF</td>
</tr>
<tr>
<td>Proximal flow sensor (sensor connected)</td>
<td>Type BF</td>
</tr>
<tr>
<td>Degree of protection against ingress of liquids and particles</td>
<td>IP21</td>
</tr>
<tr>
<td></td>
<td>Protection against particles with a diameter of more than 12.5 mm (0.47 in)</td>
</tr>
<tr>
<td></td>
<td>Protection against vertically dripping water</td>
</tr>
</tbody>
</table>

<table>
<thead>
<tr>
<th>Mains power supply</th>
<th></th>
</tr>
</thead>
<tbody>
<tr>
<td>Mains power connection</td>
<td>100 V to 240 V</td>
</tr>
<tr>
<td></td>
<td>50/60 Hz</td>
</tr>
<tr>
<td>Current consumption</td>
<td></td>
</tr>
<tr>
<td>at 230 V</td>
<td>max. 1.4 A</td>
</tr>
<tr>
<td>at 100 V</td>
<td>max. 3.0 A</td>
</tr>
<tr>
<td>Inrush current</td>
<td>approx. 8 to 24 A peak</td>
</tr>
<tr>
<td></td>
<td>approx. 6 to 17 A quasi RMS</td>
</tr>
<tr>
<td>Power consumption</td>
<td></td>
</tr>
<tr>
<td>maximum</td>
<td>300 W</td>
</tr>
<tr>
<td>during ventilation, without charging the battery</td>
<td>approx. 100 W ventilation unit with Medical Cockpit</td>
</tr>
<tr>
<td></td>
<td>approx. 180 W with GS500</td>
</tr>
<tr>
<td>Device fuses</td>
<td></td>
</tr>
<tr>
<td>Range 100 V to 240 V</td>
<td>F6.3H 250V IEC 60127-2/V (2 pcs.) Ventilation unit</td>
</tr>
</tbody>
</table>

<table>
<thead>
<tr>
<th>Batteries</th>
<th></th>
</tr>
</thead>
<tbody>
<tr>
<td>The operating time applies when the batteries are fully charged and new and ventilation is typical.</td>
<td></td>
</tr>
<tr>
<td>Low temperatures may reduce the operating time.</td>
<td></td>
</tr>
<tr>
<td>The charging time applies to new and completely discharged batteries when ventilation is typical and GS500 is not used.</td>
<td></td>
</tr>
<tr>
<td>The actual charging time depends on the battery charge.</td>
<td></td>
</tr>
<tr>
<td>If GS500 is operating or the ambient temperature is high, the battery charging process may be restricted or interrupted.</td>
<td></td>
</tr>
</tbody>
</table>
Technical data

Operating data (cont.)

<table>
<thead>
<tr>
<th>Typical ventilation</th>
<th></th>
</tr>
</thead>
<tbody>
<tr>
<td>Ventilation mode</td>
<td>PC-AC</td>
</tr>
<tr>
<td><em>FIO₂</em></td>
<td>21 Vol%</td>
</tr>
<tr>
<td><strong>PEEP</strong></td>
<td>5 mbar (or hPa or cmH₂O)</td>
</tr>
<tr>
<td><strong>Pinsp</strong></td>
<td>20 mbar (or hPa or cmH₂O)</td>
</tr>
<tr>
<td><strong>RR</strong></td>
<td>60/min</td>
</tr>
<tr>
<td>Measured <strong>MV</strong></td>
<td>400 mL/min</td>
</tr>
<tr>
<td>Ambient temperature</td>
<td>22 °C (71.6 °F)</td>
</tr>
</tbody>
</table>

Internal battery of ventilation unit (without PS500)

<table>
<thead>
<tr>
<th>Type</th>
<th>NiMH battery, sealed</th>
</tr>
</thead>
<tbody>
<tr>
<td>Fuse</td>
<td>F15A 80V UL248</td>
</tr>
<tr>
<td>Capacity</td>
<td>2.5 Ah</td>
</tr>
<tr>
<td>Voltage</td>
<td>24 V</td>
</tr>
<tr>
<td>Current</td>
<td>0 to 15 A</td>
</tr>
</tbody>
</table>

Operating time if mains power supply is not available

<p>| | |</p>
<table>
<thead>
<tr>
<th></th>
<th></th>
</tr>
</thead>
<tbody>
<tr>
<td>without GS500</td>
<td>30 minutes</td>
</tr>
<tr>
<td>with GS500</td>
<td>15 minutes</td>
</tr>
</tbody>
</table>

Charging

<table>
<thead>
<tr>
<th>Charging</th>
<th></th>
</tr>
</thead>
<tbody>
<tr>
<td>Charging time (to charge battery fully)</td>
<td>&lt;4 hours (&lt;2 hours for 80 % charge)</td>
</tr>
</tbody>
</table>

Batteries of power supply unit PS500

<table>
<thead>
<tr>
<th>Type</th>
<th>VRLA batteries</th>
</tr>
</thead>
<tbody>
<tr>
<td>Fuse</td>
<td>triple F15A 80V UL248</td>
</tr>
<tr>
<td>Capacity</td>
<td>24 Ah</td>
</tr>
<tr>
<td>Voltage</td>
<td>24 V</td>
</tr>
<tr>
<td>Current</td>
<td>0 to 15 A</td>
</tr>
</tbody>
</table>

Operating time if mains power supply is not available

<p>| | |</p>
<table>
<thead>
<tr>
<th></th>
<th></th>
</tr>
</thead>
<tbody>
<tr>
<td>without GS500</td>
<td>240 minutes</td>
</tr>
<tr>
<td>with GS500</td>
<td>120 minutes</td>
</tr>
</tbody>
</table>

Charging

<table>
<thead>
<tr>
<th>Charging</th>
<th></th>
</tr>
</thead>
<tbody>
<tr>
<td>Charging time (to charge battery fully)</td>
<td>&lt;24 hours (&lt;20 hours for 80 % charge)</td>
</tr>
</tbody>
</table>
### Operating data (cont.)

**Gas supply**

<table>
<thead>
<tr>
<th>Description</th>
<th>Specification</th>
</tr>
</thead>
<tbody>
<tr>
<td>O₂ operating pressure</td>
<td>2.7 to 6.0 bar (or 270 to 600 kPa or 39 to 87 psi)</td>
</tr>
<tr>
<td>O₂ peak input flow</td>
<td>130 L/min (at 2.8 bar inlet pressure)</td>
</tr>
<tr>
<td></td>
<td>180 L/min (at 4.0 bar inlet pressure)</td>
</tr>
<tr>
<td>O₂ connection</td>
<td>depending on configuration: DIN, NIST, DISS, Air Liquide</td>
</tr>
<tr>
<td>Air operating pressure</td>
<td>2.7 to 6.0 bar (or 270 to 600 kPa or 39 to 87 psi)</td>
</tr>
<tr>
<td>Air peak input flow</td>
<td>130 L/min (at 2.8 bar inlet pressure)</td>
</tr>
<tr>
<td></td>
<td>180 L/min (at 4.0 bar inlet pressure)</td>
</tr>
<tr>
<td>Air connection</td>
<td>depending on configuration: DIN, NIST, DISS, Air Liquide</td>
</tr>
<tr>
<td>Dew point</td>
<td>at least 5 Kelvin or 5 °C or 9 °F below ambient temperature</td>
</tr>
<tr>
<td>Oil concentration</td>
<td>&lt;0.1 mg/m³</td>
</tr>
<tr>
<td>Particle size</td>
<td>dust-free air (filtered with pore size &lt;1 µm)</td>
</tr>
</tbody>
</table>

**Gas consumption**

- Consumption for ventilation: depends on ventilation settings
- Consumption for pneumatic medication nebulizer: compressed air or O₂, max. 2.1 bar (or 210 kPa or 30.5 psi), max. 11 L/min

**Automatic gas switch-over**

If one gas fails, the device switches to the other gas.


- A-class mean surface sound pressure level (LpA) with a radius of 2 m (79 in):
  - approx. 33.0 dB
  - approx. 43.5 dB with GS500
- A-class mean surface sound pressure level (LpA) with a radius of 2 m (79 in):
  - approx. 50.0 dB with HFO
- Uncertainty (k):
  - 3.5 dB
- A-class surface sound pressure level (LWA):
  - approx. 46.0 dB
  - approx. 57.5 dB with GS500
- A-class surface sound pressure level (LWA):
  - approx. 63.5 dB with HFO
- Uncertainty (k):
  - 3.5 dB
### Technical data

#### Operating data (cont.)

<table>
<thead>
<tr>
<th>Dimensions (W x H x D)</th>
<th>361 mm x 320 mm x 410 mm</th>
</tr>
</thead>
<tbody>
<tr>
<td>Babylog VN500 with lateral standard rail (without Infinity C500)</td>
<td>14.3 in x 12.6 in x 16.1 in</td>
</tr>
<tr>
<td>Babylog VN500 and Infinity C500 on the trolley, carrier frame without bar</td>
<td>577 mm x 1420 mm x 687 mm (22.7 in x 55.9 in x 27.1 in)</td>
</tr>
<tr>
<td>Babylog VN500 and Infinity C500 on the trolley, carrier frame with bar</td>
<td>577 mm x 1420 mm x 700 mm (22.7 in x 55.9 in x 27.6 in)</td>
</tr>
<tr>
<td>Weight</td>
<td>approx. 17 kg (37.5 lb)</td>
</tr>
<tr>
<td>Ventilation unit</td>
<td>approx. 8 kg (17.6 lb)</td>
</tr>
<tr>
<td>Medical Cockpit with holder</td>
<td>approx. 33 kg (72.8 lb)</td>
</tr>
<tr>
<td>Trolley</td>
<td>approx. 25 kg (55.1 lb)</td>
</tr>
<tr>
<td>Babylog VN500 and Infinity C500 on trolley</td>
<td>approx. 59 kg (130 lb)</td>
</tr>
<tr>
<td>PS500</td>
<td>approx. 27 kg (59.5 lb)</td>
</tr>
<tr>
<td>GS500</td>
<td>approx. 10.5 kg (23 lb)</td>
</tr>
<tr>
<td>Nominal weight (weight of ventilation unit and Medical Cockpit on trolley)</td>
<td>58 g (128 lb)</td>
</tr>
<tr>
<td>Maximum weight (permitted maximum total weight)</td>
<td>133 kg (293 lb)</td>
</tr>
<tr>
<td>Maximum load</td>
<td>100 kg (220.5 lb)</td>
</tr>
<tr>
<td>Trolley</td>
<td>10 kg (22 lb)</td>
</tr>
<tr>
<td>Universal holder with standard rail (G93140)</td>
<td>5 kg (11 lb)</td>
</tr>
<tr>
<td>Humidifier holder (8416325)</td>
<td>5 kg (11 lb)</td>
</tr>
<tr>
<td>Humidifier holder (G93111)</td>
<td>5 kg (11 lb)</td>
</tr>
<tr>
<td>If a hinged arm is attached to the lateral standard rails of the ventilation unit in addition to the humidifier holder (8416325), the maximum load of 5 kg (11 lb) per lateral standard rail must be observed. The humidifier holder can then only support 4 kg (8.8 lb).</td>
<td></td>
</tr>
<tr>
<td>Electromagnetic compatibility EMC</td>
<td>tested in accordance with IEC 60601-1-2</td>
</tr>
<tr>
<td>Classification according to Directive 93/42/EEC, Annex IX</td>
<td>II b</td>
</tr>
<tr>
<td>UMDNS code</td>
<td>17-429</td>
</tr>
<tr>
<td>Universal Medical Device Nomenclature System – Nomenclature for medical devices</td>
<td></td>
</tr>
</tbody>
</table>
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Operating data (cont.)

Materials used

<table>
<thead>
<tr>
<th>Material</th>
<th>Description</th>
</tr>
</thead>
<tbody>
<tr>
<td>Breathing hose (reusable)</td>
<td>Silicone rubber (milky, transparent)</td>
</tr>
<tr>
<td>Water trap (reusable)</td>
<td>Polysulphone (gray, transparent)</td>
</tr>
<tr>
<td>Y-piece (reusable)</td>
<td>Polysulphone (yellow, transparent)</td>
</tr>
<tr>
<td>Expiratory valve (reusable; housing, closure, nozzle)</td>
<td>Polyamide</td>
</tr>
<tr>
<td>Inspiratory unit (reusable; housing, nozzle)</td>
<td>Polyamide</td>
</tr>
<tr>
<td>Diaphragm</td>
<td>Silicone rubber and nickel (whitish and gray)</td>
</tr>
<tr>
<td>Reusable CO2 cuvette</td>
<td>Polysulphone with sapphire windows (gray violet, transparent: pediatric cuvette)</td>
</tr>
<tr>
<td>Disposable CO2 cuvette</td>
<td>Styrene-butadiene copolymer SBC (blue, transparent: pediatric cuvette)</td>
</tr>
<tr>
<td>CO2 sensor</td>
<td>Polysulphone (white)</td>
</tr>
<tr>
<td>CO2 sensor cable</td>
<td>Polyurethane (gray)</td>
</tr>
</tbody>
</table>

For Nurse call

<table>
<thead>
<tr>
<th>Connection</th>
<th>via cable 8417370 only</th>
</tr>
</thead>
<tbody>
<tr>
<td>Potential-free DC contact</td>
<td></td>
</tr>
<tr>
<td>Input voltage</td>
<td>24 V DC max.</td>
</tr>
<tr>
<td>Input current</td>
<td>1 A DC max.</td>
</tr>
<tr>
<td>Switching capacity</td>
<td>15 W max.</td>
</tr>
<tr>
<td>Contact pin assignment, see chapter</td>
<td>&quot;Assembly and preparation&quot;, &quot;Connecting the nurse call&quot;</td>
</tr>
</tbody>
</table>

Device ports

Outputs

<table>
<thead>
<tr>
<th>Port</th>
<th>Description</th>
</tr>
</thead>
<tbody>
<tr>
<td>V1</td>
<td>System cable</td>
</tr>
<tr>
<td>V2, V3</td>
<td>not used</td>
</tr>
<tr>
<td>V4</td>
<td>Nurse call</td>
</tr>
<tr>
<td>V5</td>
<td>Neonatal flow sensor</td>
</tr>
<tr>
<td>V6</td>
<td>not used</td>
</tr>
<tr>
<td>V7</td>
<td>CO2 sensor</td>
</tr>
<tr>
<td>V8</td>
<td>not used</td>
</tr>
<tr>
<td>V9</td>
<td>GS500</td>
</tr>
</tbody>
</table>
## Technical data

### Device ports (cont.)

<table>
<thead>
<tr>
<th>MEDIBUS or MEDIBUS.X protocol</th>
</tr>
</thead>
<tbody>
<tr>
<td>Baud rate</td>
</tr>
</tbody>
</table>

| Data bits | 8 |
| Parity    | even, odd, no |
| Stop bits | 1 or 2 |

### Pin assignment of COM1, COM2 and COM3

<table>
<thead>
<tr>
<th>Pin</th>
<th>Assignment</th>
</tr>
</thead>
<tbody>
<tr>
<td>1</td>
<td>DCD</td>
</tr>
<tr>
<td>2</td>
<td>RXD</td>
</tr>
<tr>
<td>3</td>
<td>TXD</td>
</tr>
<tr>
<td>4</td>
<td>DTR</td>
</tr>
<tr>
<td>5</td>
<td>GND</td>
</tr>
<tr>
<td>6</td>
<td>DSR</td>
</tr>
<tr>
<td>7, 8</td>
<td>RTS/CTS</td>
</tr>
<tr>
<td>9</td>
<td>RI</td>
</tr>
<tr>
<td>Housing</td>
<td>SHLD</td>
</tr>
</tbody>
</table>

### Galvanic isolation

<table>
<thead>
<tr>
<th>V</th>
<th>Description</th>
</tr>
</thead>
<tbody>
<tr>
<td>V1</td>
<td>The port is not electrically isolated from the device electronics.</td>
</tr>
<tr>
<td>V2, V3</td>
<td>not used</td>
</tr>
<tr>
<td>V4</td>
<td>The port is not electrically isolated from the device electronics.</td>
</tr>
<tr>
<td>V5</td>
<td>The port is electrically isolated from the device electronics (Type BF). The test voltage for electrical isolation is 1500 V.</td>
</tr>
<tr>
<td>V6</td>
<td>not used</td>
</tr>
<tr>
<td>V7</td>
<td>The port is not electrically isolated from the device electronics.</td>
</tr>
<tr>
<td>V8</td>
<td>not used</td>
</tr>
<tr>
<td>V9</td>
<td>The port is electrically isolated from the device electronics. The test voltage for electrical isolation is 500 V.</td>
</tr>
</tbody>
</table>
Automatic alarm limits

The following tables describe the alarm limits which cannot be set by the user.

Pressure monitoring

<table>
<thead>
<tr>
<th>Alarm message</th>
<th>Description/Detection</th>
</tr>
</thead>
<tbody>
<tr>
<td><strong>Airway pressure high</strong></td>
<td>The airway pressure is monitored to detect whether the upper alarm limit is exceeded. If the alarm limit indicating a too high airway pressure is linked to ventilation therapy controls, this limit is set 5 mbar (5 cmH₂O) above the highest pressure which is regularly applied during ventilation according to the user settings. This connection is switched on at the factory.</td>
</tr>
<tr>
<td><strong>VT low</strong></td>
<td>Under Volume Guarantee, 90 % of the set VT is not reached during eight consecutive breaths.</td>
</tr>
<tr>
<td><strong>Breathing hose kinked</strong></td>
<td>An excessive pressure during an O₂ therapy is monitored. The alarm limit is set at 30 mbar (30 cmH₂O).</td>
</tr>
<tr>
<td>(O₂ Therapy)</td>
<td></td>
</tr>
<tr>
<td><strong>Airway pressure negative</strong></td>
<td>Situations in which the pressure becomes negative are monitored. The alarm limit is set at –10 mbar (–10 cmH₂O).</td>
</tr>
<tr>
<td><strong>PEEP high/Plow high</strong></td>
<td>The alarm limit is 8 mbar (8 cmH₂O) above the set PEEP or Plow level. The alarm triggers a pressure release to ambient pressure. The alarm is not triggered below 11 mbar (11 cmH₂O). An alarm is triggered if this condition applies for 2 breaths or after a maximum of 15 seconds. To avoid false alarms, it is not monitored whether the lower pressure level has been reached if in APRV and if Tlow is smaller 1 s or AutoRelease is activated.</td>
</tr>
<tr>
<td>(!!!)</td>
<td></td>
</tr>
<tr>
<td><strong>PEEP high/Plow high</strong></td>
<td>The alarm limit is 4 mbar (4 cmH₂O) above the set PEEP. An alarm is triggered if this condition applies for 2 breaths or after a maximum of 15 seconds.</td>
</tr>
<tr>
<td>(!!)</td>
<td></td>
</tr>
<tr>
<td><strong>PEEP low/Plow low</strong></td>
<td>A too low PEEP or Plow value during ventilation is monitored. The alarm limit depends on the set value of the PEEP or Plow level. The alarm limit is 3 mbar (3 cmH₂O) below the set value. An alarm is triggered if this condition applies for 10 breaths.</td>
</tr>
</tbody>
</table>
### Technical data

#### Volume monitoring

The expiratory minute volume $\text{MVe}$ is monitored in the patient category *Ped. pat.* and in the case of invasive ventilation in the patient category *Neo.* within the set alarm limits.

The minimum tidal volume is monitored only when Volume Guarantee is activated. To accomplish this, with leakage compensation switched off the value $\text{VTi}$ is monitored in the patient category *Ped. pat.*, and the value $\text{VTe}$ is monitored in the patient category *Neo.* When leakage compensation is activated, $\text{VT}$ is generally used and the automatically set alarm limit $\text{VT low}$ is monitored, whereby the limit $\text{VT low}$ corresponds to 90% of the selected $\text{VT}$.

#### Alarm message

<table>
<thead>
<tr>
<th>Alarm message</th>
<th>Description/Detection</th>
</tr>
</thead>
<tbody>
<tr>
<td><strong>Pressure limited (ATC/PPS)</strong></td>
<td>The upper pressure limit is monitored to detect whether it is reached when using ATC or PPS. If the <strong>Paw high</strong> alarm limit is adjustable, the alarm limit is derived from this value and lies in the range $\text{Paw high} - 5 \text{ mbar}$ ($-5 \text{ cmH}_2\text{O}$) to $\text{Paw high} - 1 \text{ mbar}$ ($-1 \text{ cmH}_2\text{O}$), depending on how close the <strong>Paw high</strong> value comes to the currently applied ventilation. If the <strong>Paw high</strong> alarm limit is linked (<strong>Pmax/Paw high autoset</strong>), the pressure limit corresponds to the value of the <strong>Pmax</strong> therapy control.</td>
</tr>
<tr>
<td><strong>Airway pressure low</strong></td>
<td>An insufficient airway pressure is monitored by checking whether the integral of undercutting the measured pressure values of the lower pressure level exceeds $22.5 \text{ mbar} \times \text{s}$ ($22.5 \text{ cmH}_2\text{O} \times \text{s}$).</td>
</tr>
</tbody>
</table>

#### Airway pressure low

An insufficient airway pressure is monitored by checking whether the integral of undercutting the measured pressure values of the lower pressure level exceeds $22.5 \text{ mbar} \times \text{s}$ ($22.5 \text{ cmH}_2\text{O} \times \text{s}$).
Monitoring of the breathing circuit and the patient connection

<table>
<thead>
<tr>
<th>Alarm message</th>
<th>Description/Detection</th>
</tr>
</thead>
<tbody>
<tr>
<td><strong>Disconnection?</strong></td>
<td>Disconnection is monitored by checking that the mandatory breaths reach a minimum pressure level. The alarm limit is derived from the set points for ventilation.</td>
</tr>
<tr>
<td></td>
<td>During pressure-controlled ventilation, the alarm is triggered when the airway pressure is lower than the lower pressure level plus 50 % of the pressure difference between the upper and lower pressure levels.</td>
</tr>
<tr>
<td></td>
<td>During pressure-supported ventilation, the alarm is triggered when the airway pressure is lower than the lower pressure level plus 30 % of the pressure difference between the upper and lower pressure levels.</td>
</tr>
<tr>
<td></td>
<td>During ventilation with Volume Guarantee and volume support, the limit is 50 % of the pressure difference between the upper pressure level and the lower pressure level currently calculated by Babylog VN500.</td>
</tr>
<tr>
<td></td>
<td>During volume-controlled ventilation, the pressure level is 5 mbar (5 cmH₂O) above PEEP.</td>
</tr>
<tr>
<td></td>
<td>All pressure criteria become ineffective if a sufficient expiration has been detected.</td>
</tr>
<tr>
<td></td>
<td>In the event of an excessive inspiratory flow at the current airway pressure, a disconnection due to excessive inspiratory volume is detected. This volume depends on the patient category:</td>
</tr>
<tr>
<td></td>
<td>– 1.5 L in the <strong>Ped. pat.</strong> patient category</td>
</tr>
<tr>
<td></td>
<td>– 0.5 L in the <strong>Neo.</strong> patient category</td>
</tr>
<tr>
<td><strong>Leakage</strong></td>
<td>Leakages are monitored in the <strong>Ped. pat.</strong> patient category. The alarm limit is set at 55 % of relative Leakage. Leakages during NIV are not monitored.</td>
</tr>
<tr>
<td><strong>Airway obstructed?</strong></td>
<td>Obstructions in the breathing circuit are monitored by observing the flow delivered to the patient during a defined period.</td>
</tr>
</tbody>
</table>
FiO2 monitoring

<table>
<thead>
<tr>
<th>Alarm message</th>
<th>Description/Detection</th>
</tr>
</thead>
<tbody>
<tr>
<td><strong>FiO2 high</strong></td>
<td>An excessive O₂ concentration of the applied gas is monitored. The alarm limit is 4 Vol% above the set value if this is less than or equal to 60 Vol%. The alarm limit is 6 Vol% above the set value if this is greater than 60 Vol%.</td>
</tr>
<tr>
<td><strong>FiO2 low</strong></td>
<td>An insufficient O₂ concentration of the applied gas is monitored. For an FiO₂ concentration of 21 Vol% the alarm limit is 18 Vol%. The alarm limit is 4 Vol% below the set value if this is greater than 21 Vol% and less than or equal to 60 Vol%. The alarm limit is 6 Vol% below the set value if this is greater than 60 Vol%.</td>
</tr>
</tbody>
</table>

CO₂ monitoring

<table>
<thead>
<tr>
<th>Alarm message</th>
<th>Description/Detection</th>
</tr>
</thead>
<tbody>
<tr>
<td><strong>CO₂ sensor?</strong></td>
<td>The correct functioning of the CO₂ sensor is monitored. An alarm is immediately generated in the event of a technical defect or if a sensor is not connected. An alarm is generated after 60 s if the sensor is removed from the cuvette or the sensor does not detect any breathing activity.</td>
</tr>
</tbody>
</table>

Essential performance characteristics

The essential performance consists in a controlled and monitored patient ventilation with user-defined settings for the monitoring functions:

– minimum and maximum tidal volume,
– maximum airway pressure,
– minimum and maximum O₂ concentration in the breathing gas,

or, if a set limit is exceeded, an appropriate alarm.

Additionally, the integrated monitoring alarms in the following situations:

– Failure of the external power supply
– Battery discharge
– Failure of the gas supply
Connections to IT networks

In an IT network, data can be exchanged by means of wired or wireless technologies. An IT network can be any data interface (e.g., RS232, LAN, USB, printer interface) that is described in standards and conventions.

During operation, this device can exchange information with other devices by means of IT networks and supports the following functions:

- Display of waveforms and parameter data
- Signaling of alarms
- Transfer of device settings and patient data
- Service mode, access to logbooks

Connecting this device to a network that incorporates other devices or making subsequent changes to that network can lead to new risks for patients, users, and third parties. Before the device is connected to the network or the network is changed, these risks must be identified, analyzed, and evaluated, and appropriate measures taken.

Examples of subsequent changes to the network:

- Changing the network configuration
- Removing devices from the network
- Adding new devices to the network
- Performing upgrades or updates on devices that are connected to the network

Information on connecting to the network

Prerequisites

This device must only be connected to the network by service personnel. The IT representative of the hospital must be consulted in advance.

The following documents must be observed:

- Accompanying documents of this device
- Description of the network interface
- Description of the network-based alarm systems

Dräger recommends complying with IEC 80001-1 (risk management for IT networks with medical devices).

Serial interfaces

The following interfaces are supported:

- RS232 interfaces conforming to EIA RS232 (CCITT V.24/V.28) for the following applications:
  - MEDIBUS, MEDIBUS.X
  - Connection to medical devices from other manufacturers
Consequences of using an unsuitable network

If the network does not meet the requirements, dangerous situations can result. The following situations can occur with this device:

- Due to an insecure decentralized alarm system:
  - Alarms or data are transmitted at the wrong time.
  - Alarms are not transmitted.
- During an interruption of the network connection:
  - Suppressed alarms or alarm tones are not reactivated, but remain suppressed.
  - Alarms are not transmitted.
- Without firewall and antivirus software:
  - Data are not protected.
  - Device settings are changed.
  - The device generates false alarms or no alarms.
- Data are sent incomplete, sent to the wrong device, or not sent at all.
- Patient data are intercepted, falsified, or damaged.
- Data have incorrect time stamps.

Requirements for the electrical characteristics of connected devices and networks

The analog and digital ports are only appropriate for connecting devices or networks that have a nominal voltage on the network side of max. 24 V DC and meet the requirements of one of the following standards:

- IEC 60950-1: Ungrounded SELV circuits
- IEC 60601-1 (as of 2nd edition): Touchable secondary circuits

Open-source software

Dräger devices that use software may use open-source software, depending on their setup. Open-source software may be subject to different terms of license. Additional information regarding the open-source software used in this device is available at the following web page:

www.draeger.com/opensource
This page has been left blank intentionally.
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Principles of operation

Description of the ventilation modes

PC-CMV

Pressure Control-Continuous Mandatory Ventilation

Continuous pressure-controlled ventilation allowing spontaneous breathing (open system) during the entire respiratory cycle.

**Pressure-controlled ventilation**

The upper pressure level is determined by \( P_{\text{insp}} \). The duration of the mandatory breaths is determined by \( T_i \). As in all pressure-controlled ventilation modes, the tidal volume supplied depends on the difference in pressure \( P_{\text{insp}} - P_{\text{PEEP}} \), the lung mechanics (resistance and compliance) and the patient’s respiratory drive. The pressure rise from the lower pressure level \( P_{\text{PEEP}} \) to the upper pressure level \( P_{\text{insp}} \) is determined by the Slope or InsP. flow setting. The start setting can be configured on the System setup > Ventilation > Start settings > Pressures, O2, I:E page.

The mandatory breaths are time-cycled and are not triggered by the patient. The number of mandatory breaths is determined by the respiratory rate \( R_R \).
PC-SIMV

Pressure Control-Synchronized Intermittent Mandatory Ventilation

Intermittent, triggered, pressure-controlled ventilation allowing spontaneous breathing (open system) during the entire respiratory cycle.

Pressure-controlled ventilation

The upper pressure level is determined by PinSp. The duration of the mandatory breaths is determined by Ti. As in all pressure-controlled ventilation modes, the tidal volume supplied depends on the difference in pressure \( \text{P}_{\text{insp}} - \text{PEEP} \), the lung mechanics (resistance and compliance) and the patient’s respiratory drive. The pressure rise from the lower pressure level PEEP to the upper pressure level PinSp is determined by the \textbf{Slope} or \textbf{Insp. flow} setting. The start setting can be configured on the System setup > Ventilation > Start settings > Pressures, O2, I:E page.

In the Neo. patient category, this mode is not available with non-invasive ventilation.

Synchronization

The mandatory breaths can be triggered by the patient’s inspiratory effort from the PEEP level. By setting the trigger level, the mandatory breaths can be synchronized with the patient’s inspiratory efforts.

A mandatory breath can only be triggered within a “trigger window” by the flow trigger in synchrony with the patient’s spontaneous inspiratory effort. This prevents the mandatory breath being applied during expiration.

The trigger window is 1.5 seconds long. For expiratory times shorter than 1.5 seconds, the trigger window covers the entire expiratory time minus a refractory period for the previous expiration.
Principles of operation

Synchronization of mandatory breaths reduces the expiratory time. Babylog VN500 extends the subsequent expiratory time or spontaneous breathing time by the missing time. This prevents an increase of the mandatory respiratory rate.

The number of mandatory breaths is determined by the respiratory rate $RR$.

Pressure support

During spontaneous breathing from the PEEP level, the patient can be supported with $PS$. Every inspiratory effort of the patient that meets the trigger criteria triggers a pressure-supported breath. By setting the trigger level, the patient's inspiratory efforts are synchronized. The time, number, and duration of pressure-supported breaths is determined by the patient's spontaneous breathing.

As in all pressure-controlled ventilation modes, the tidal volume supplied depends on the difference in pressure "$Psupp - PEEP$", the lung mechanics (resistance and compliance) and the patient's respiratory drive. The pressure rise from the lower pressure level $PEEP$ to the upper pressure level $Psupp$ is determined by the $Slope$ or $Insp. flow$ setting. The start setting can be configured on the $System setup > Ventilation > Start settings > Pressures, O2, I:E$ page.

The pressure support is terminated as soon as the inspiratory flow falls below 15 % of the maximum inspiratory flow.

The pressure support is also terminated as soon as the duration of the support has reached the maximum inspiratory time. For intubated patients, the maximum inspiratory time in the $Ped. pat.$ patient category is limited to 1.5 seconds. In the $Neo.$ patient category, the maximum inspiratory time is limited to 130 % of $Ti$, maximum of 1.5 seconds.
**PC-AC**

Pressure Control-Assist Control

Assist-controlled, pressure-controlled ventilation allowing spontaneous breathing during the entire respiratory cycle and backup respiratory rate

### Pressure-controlled ventilation

The upper pressure level is determined by $P_{insp}$. The duration of the mandatory breaths is determined by $T_i$. As in all pressure-controlled ventilation modes, the tidal volume supplied depends on the difference in pressure "$P_{insp} - P_{EEP}$", the lung mechanics (resistance and compliance) and the patient’s respiratory drive. The pressure rise from the lower pressure level $P_{EEP}$ to the upper pressure level $P_{insp}$ is determined by the *Slope* or *Insp. flow* setting. The start setting can be configured on the System setup > Ventilation > Start settings > Pressures, O2, I:E page.

In the Neo patient category, this mode is not available with non-invasive ventilation.

### Assisted-controlled ventilation

Every inspiratory effort of the patient from the PEEP level triggers a synchronized mandatory breath. Thus, the time and number of mandatory breaths are determined by the patient. The trigger window covers the expiratory time minus a refractory period for the previous expiration. The expiratory time is determined by the respiratory rate $RR$ and the inspiratory time $T_i$. A non-synchronized mandatory breath is triggered at the latest at the end of the expiratory time (back-up respiratory rate).

The minimum number of mandatory breaths is determined by the respiratory rate $RR$. 
PC-PSV

Pressure Control-Pressure Support Ventilation

Pressure-controlled ventilation with guaranteed minimum respiratory rate (backup respiratory rate)

Pressure support

During spontaneous breathing from the PEEP level, the patient can be supported with PS. The level of pressure support is determined by Pinsp. Every inspiratory effort of the patient that meets the trigger criteria triggers a pressure-supported breath. By setting the trigger level, the patient's inspiratory efforts are synchronized. The time, number, and duration of pressure-supported breaths is determined by the patient's spontaneous breathing. If the patient's respiratory rate is less than the set back-up respiratory rate RR or there is no spontaneous breathing present, the device administers pressure-supported breaths with the respiratory rate RR.

As in all pressure-controlled ventilation modes, the tidal volume supplied depends on the difference in pressure "Pinsp – PEEP", the lung mechanics (resistance and compliance) and the patient's respiratory drive. The pressure rise from the lower pressure level PEEP to the upper pressure level Pinsp is determined by the Slope or Insp. flow setting. The start setting can be configured on the System setup > Ventilation > Start settings > Pressures, O2, I:E page.

The pressure support is terminated as soon as the inspiratory flow falls below 15 % of the maximum inspiratory flow, see "Inspiratory termination" on page 307.

The pressure support is also terminated as soon as the duration of the support has reached the maximum inspiratory time.

For intubated patients, the maximum inspiratory time in the Ped. pat. patient category is limited to 1.5 seconds. For the Neo. patient category, this mode is not available with non-invasive ventilation.
**PC-MMV**

Pressure Control-Mandatory Minute Volume
Ventilation
Pressure-controlled ventilation to ensure minimum minute ventilation

![Diagram of PC-MMV](image)

**Pressure-controlled ventilation with volume guarantee**

The tidal volume of the mandatory breaths is determined by the volume \( VT \). The duration of the mandatory breaths is determined by \( Ti \). The pressure rise is determined by the **Slope** or **Insp. flow** setting. The maximum pressure that Babylog VN500 uses is set via the **Pmax** therapy control. If **Pmax** is not linked to the alarm limit **Paw high**, the pressure can be limited using **Paw high**.

In this case, the maximum applied pressure is limited to 5 mbar (5 cmH2O) below **Paw high**. If the maximum pressure allowed is not enough to deliver the set **VT**, Babylog VN500 generates an alarm.

**MMV** works similar to SIMV, however, the mandatory breaths are only provided if spontaneous breathing is not sufficient and below the prescribed minimum ventilation. Should spontaneous breathing increase, fewer mandatory breaths will be provided. The minimum ventilation is determined by the setting of the tidal volume **VT** and the respiratory rate **RR**.
Principles of operation

The maximum number of mandatory breaths is determined by the respiratory rate $RR$. However, this number is only provided when there is insufficient spontaneous breathing or an apnea is present.

Pressure support

During spontaneous breathing from the PEEP level, the patient can be supported with $PS$. Every inspiratory effort of the patient that meets the trigger criteria triggers a pressure-supported breath. By setting the trigger level, the patient’s inspiratory efforts are synchronized. The time, number, and duration of pressure-supported breaths is determined by the patient’s spontaneous breathing.

As in all pressure-controlled ventilation modes, the tidal volume supplied depends on the difference in pressure "$P_{supp} - P_{EEP}$", the lung mechanics (resistance and compliance) and the patient’s respiratory drive. The pressure rise from the lower pressure level $P_{EEP}$ to the upper pressure level $P_{supp}$ is determined by the $Slope$ or $Insp.\ flow$ setting. The start setting can be configured on the $System\ setup > Ventilation > Start\ settings > Pressures, O2, I:E$ page.

The pressure support is terminated as soon as the inspiratory flow falls below 15 % of the maximum inspiratory flow.

The pressure support is also terminated as soon as the duration of the support has reached the maximum inspiratory time. For intubated patients, the maximum inspiratory time in the Ped. pat. patient category is limited to 1.5 seconds. In the Neo. patient category, the maximum inspiratory time is limited to 130 % of $T_i$, maximum of 1.5 seconds.

In the Neo. patient category, this mode is not available with non-invasive ventilation.

PC-HFO

Pressure Control-High Frequency Oscillation

Continuous pressure-controlled ventilation with high-frequency oscillations at the mean pressure level

Ventilation with high-frequency pressure oscillations enables gas to be exchanged in the lungs despite very small tidal volumes (often in the dead space volume range). While pressure amplitudes may be considerable in the breathing circuit, only small fluctuations occur around the
mean pressure in the lungs. The mechanical load due to periodic expansion and relaxation of the lungs is low.

Depending on the breathing circuit used the set pressure amplitude may not be reached.

The mean pressure, around which the oscillations occur, is determined by $\text{MAP}_{hf}$. The pressure amplitude is set directly using the $\text{Ampl}_{hf}$ therapy control. Here, $\text{Ampl}_{hf}$ is the difference between the maximum and minimum pressure of the oscillation. The rate at which oscillations occur per second is set with $\text{fhf}$.

Depending on the oscillation frequency $\text{fhf}$, there are up to three different I:E ratios to choose from:

<table>
<thead>
<tr>
<th>I:Ehf</th>
<th>Oscillation frequency</th>
</tr>
</thead>
<tbody>
<tr>
<td>1:1</td>
<td>5 to 20 Hz</td>
</tr>
<tr>
<td>1:2</td>
<td>5 to 15 Hz</td>
</tr>
<tr>
<td>1:3</td>
<td>5 to 10 Hz</td>
</tr>
</tbody>
</table>

**Device flow**

The displayed device flow is the flow delivered by the inspiratory valve and is based on customized settings.

The displayed device flow may change if the settings are changed by the user or are changed automatically by the device (e.g. in the case of VG (HF)).

When using HFO, the device flow delivered to the patient may be influenced by the following factors:

- Tube leakage
- Change in resistance
- Change in hose compliance

The measured value **Device flow** indicates only the flow delivered by the ventilator. The flow from external flow sources delivered to the patient system by the user is not considered.
Principles of operation

PC-APRV

Pressure Control-Airway Pressure Release

Ventilation

Spontaneous breathing under continuous positive airway pressure with brief pressure releases

The patient breathes spontaneously at a high pressure level \( P_{\text{high}} \) for an adjustable length of time \( T_{\text{high}} \). For very short expiratory times \( T_{\text{low}} \), Babylog VN500 switches to a low pressure level \( P_{\text{low}} \). The normal lung areas are emptied, but the "slow" lung areas only change volume to a lesser extent*.

The number of pressure releases is determined by the \( T_{\text{high}} \) and \( T_{\text{low}} \) settings. The releases are time-cycled and are not triggered by the patient. The duration is determined by \( T_{\text{low}} \). The tidal volume exchanged during the release phases depends on the difference in pressure \( P_{\text{high}} - P_{\text{low}} \), the lung mechanics (resistance and compliance) and the length of pressure release \( T_{\text{low}} \). The pressure rise from the lower pressure level \( P_{\text{low}} \) to the upper pressure level \( P_{\text{high}} \) is determined by the slope or \( \text{Insp. flow} \) setting.

The start setting can be configured on the System setup > Ventilation > Start settings > Pressures, O2, I:E page.

When \( \text{AutoRelease} \) is activated, the duration of pressure releases is determined by the expiratory flow trace. The Exp. term. setting determines the percentage by which the expiratory flow must fall short in relation to the peak flow for the ventilation to return to the high pressure level.

When \( \text{AutoRelease} \) is switched on, the change-over from the upper pressure level \( P_{\text{high}} \) to the lower pressure level \( P_{\text{low}} \) is synchronized with the patient's spontaneous breathing.

Synchronization of the mandatory breath reduces the time on the upper pressure level. Babylog VN500 prolongs the subsequent ventilation time on the upper pressure level by the missing time. This prevents an increase in respiratory rate.

In the Neo. patient category, this mode is not available with non-invasive ventilation.

* References [1], [2], [3], [4], see page 336.
**SPN-CPAP/PS**

Spontaneous-Continuous Positive Airway Pressure/Pressure Support

Spontaneous breathing with continuous positive pressure level with or without pressure support

When the pressure support is not switched on, the patient's spontaneous breathing is merely supported by an increased PEEP.

During spontaneous breathing from the PEEP level, the patient can be supported with PS. Every inspiratory effort of the patient that meets the trigger criteria triggers a pressure-supported breath. The time, number, and duration of pressure-supported breaths is determined by the patient's spontaneous breathing.

As in all pressure-controlled ventilation modes, the tidal volume supplied depends on the difference in pressure "Psupp – PEEP", the lung mechanics (resistance and compliance) and the patient's respiratory drive. The pressure rise from the lower pressure level PEEP to the upper pressure level Psupp is determined by the Slope or Insp. flow setting. The start setting can be configured on the System setup > Ventilation > Start settings > Pressures, Oz, I:E page.

The pressure support is terminated as soon as the inspiratory flow falls below 15% of the maximum inspiratory flow.

The pressure support is also terminated as soon as the duration of the support has reached the maximum inspiratory time. For intubated patients, the maximum inspiratory time in the Ped. pat. patient category is limited to 1.5 seconds. For the Neo. patient category, the maximum inspiratory time can be set with Timax to a maximum of 1.5 seconds. In the case of non-invasive ventilation, the maximum duration of support for the Ped. pat. patient category can be set with Timax.

In the Neo. patient category, this mode is not available with non-invasive ventilation.
**SPN-CPAP/VS**

**Spontaneous-Continuous Positive Airway Pressure/Volume Support**

Spontaneous breathing with continuous positive pressure level with or without volume support

For volume support **VS**, every inspiratory effort by the patient on the PEEP level that meets the trigger criteria triggers a volume-supported breath. By setting the trigger level, the patient's inspiratory efforts are synchronized. The time, number and duration of volume-supported breaths is determined by the patient's spontaneous breathing. The pressure rise is determined by the **Slope** or **Insp. flow** setting. The start setting can be configured on the System setup > Ventilation > Start settings > Pressures, O2, I:E page.

The volume support is terminated as soon as the inspiratory flow falls below 15% of the maximum inspiratory flow.

The volume support is also terminated as soon as the duration of the support has reached the maximum inspiratory time. For intubated patients, the maximum inspiratory time in the **Ped. pat.** patient category is limited to 1.5 seconds. For the **Neo.** patient category, the maximum inspiratory time can be set with **Timax** to a maximum of 1.5 seconds. In the case of non-invasive ventilation, the maximum duration of support for the **Ped. pat.** patient category can be set with **Timax**.

In the **Neo.** patient category, this mode is not available with non-invasive ventilation.

The set tidal volume of the supported breaths is reached through the automatically controlled pressure level of the volume support. With volume support, the support pressure is automatically adjusted to changes in lung conditions (resistance and compliance) and to the spontaneous breathing demand of the patient.

If **Paw high** is linked to the **Pmax** therapy control, set the maximum pressure that can be applied with the **Pmax** setting!

If **Paw high** is not linked to the **Pmax** therapy control, always set the **Paw high** alarm limit so that Babylog VN500 generates an alarm in the event of an increase in airway pressure due to reduced compliance. The maximum pressure that can be applied is limited to 5 mbar (5 cmH2O) below the upper alarm limit.
SPN-CPAP

Spontaneous-Continuous Positive Airway Pressure

Spontaneous breathing with continuous positive pressure level in the application mode NIV

The SPN-CPAP ventilation mode is only available with non-invasive ventilation in the Neo patient category.

The patient’s spontaneous breathing is supported with an increased PEEP.

For the Manual inspiration/hold maneuver, the pressure of the breath is set with the PmanInsP therapy control and the duration of the breath is set with the TmanInsP therapy control.
SPN-PPS

Spontaneous-Proportional Pressure Support

Spontaneous breathing with flow- and volume-proportional pressure support

In ventilation mode SPN-PPS, Babylog VN500 supports the patient’s spontaneous breathing in proportion to the inspiratory effort. If the patient breathes strongly, Babylog VN500 supports this effort with high pressure support. If the patient has shallow breathing, Babylog VN500 reacts with low pressure support. Mechanical support is omitted altogether if there is no spontaneous breathing. Monitoring of apnea and minute volume must therefore be set appropriately.

The degree of support in PPS mode can be set separately according to the resistive and elastic components. Using the resistive proportion Flow Assist, the user defines how much of the resistive work of breathing is taken over by Babylog VN500. Using the elastic proportion Vol. Assist, the user defines how much of the elastic work of breathing is taken over by Babylog VN500. This support is only effective during inspiration.
The tidal volume is limited by the $VT_{max}$ setting in order to protect the patient from excessive tidal volumes in case of sudden leakages or incorrectly set PPS. If $VT_{max}$ is reached, the breath is stopped and an expiration is started. Babylog VN500 displays the alarm message VT limited. If leakage compensation is activated, the leakage-compensated tidal volume is used. If leakage compensation is deactivated, the tidal volume measured on the inspiratory side is used. The start-up value for $VT_{max}$ corresponds to 130 % of the preconfigured tidal volume.

The pressure support is terminated as soon as the inspiratory flow falls below 15 % of the maximum inspiratory flow.

The pressure support is also terminated as soon as the duration of the support has reached the maximum inspiratory time. For intubated patients, the maximum inspiratory time in the Ped. pat. patient category is limited to 1.5 seconds. For the Neo. patient category, the maximum inspiratory time can be set with Timax to a maximum of 1.5 seconds. In the case of non-invasive ventilation, the maximum duration of support for the Ped. pat. patient category can be set with Timax.

In the Neo. patient category, this mode is not available with non-invasive ventilation.

If Paw high is linked to the Pmax therapy control, set the maximum pressure that can be applied with the Pmax setting!

If Paw high is not linked to the Pmax therapy control, always set the Paw high alarm limit so that Babylog VN500 generates an alarm in the event of an increase in airway pressure due to reduced compliance. The maximum pressure that can be applied is limited to 5 mbar (5 cmH2O) below the upper alarm limit.
Additional settings for ventilation

Apnea Ventilation

For switching over automatically to volume-guaranteed mandatory ventilation in case of apnea.

For Babylog VN500 to be able to detect an apnea, flow measurement with the neonatal flow sensor must function and flow monitoring with the neonatal flow sensor must be activated.

Babylog VN500 detects an apnea when no expiratory flow is measured or insufficient inspiratory gas is delivered during the set apnea alarm time $Tapn$. If apnea ventilation is activated, the device starts volume-guaranteed ventilation with the ventilation parameters $RRapn$ and $VTapn$. The inspiratory time for apnea ventilation is determined from the set apnea respiratory rate $RRapn$ and a fixed I:E ratio of 1:2.

The patient can breathe spontaneously and the mandatory breaths are synchronized with the patient’s spontaneous breathing. The apnea ventilation respiratory rate $RRapn$ remains constant. Babylog VN500 provides synchronized intermittent mandatory ventilation.

Apnea ventilation is terminated by touching the Apn. Vent. reset button. Babylog VN500 continues ventilating in the previously set ventilation mode. Changing the ventilation mode or the additional settings, e.g. PS, also terminates apnea ventilation.

If an apnea situation generating an alarm occurs again during apnea ventilation, this indicates that the apnea ventilation respiratory rate $RRapn$ has been set too low in relation to apnea alarm time $Tapn$. 
Automatic return from apnea ventilation

If the **Automatic return from Apnea Ventilation** function is configured, then the device automatically switches to the previous ventilation mode when sufficient spontaneous breathing is resumed. The following conditions must be met:

- Apnea ventilation must have been active for at least 2 minutes.
- The alarm message MV low is not active.
- One of the following conditions must additionally be met:
  - The ratio of MVespon to MVe is greater than 25 % and the ratio of MVleak to MVe is less than 40 %.
  - 80 % of the mandatory breaths are triggered spontaneously.

If apnea reoccurs within 3 minutes following automatic termination of apnea ventilation in the *Ped. pat.* patient category, the **Automatic return from Apnea Ventilation** function is disabled until apnea ventilation is terminated manually or another ventilation mode is selected.

For configuration of the **Automatic return from Apnea Ventilation** function, see "Configuring general settings" on page 177.
Principles of operation

Flow trigger

The flow trigger is used to synchronize mandatory or pressure-supported breaths with spontaneous breathing. The flow trigger is also used to trigger breaths with SPN-CPAP/PS and SPN-CPAP/VS.

![Diagram of Paw, PEEP, Flow, Trigger threshold, and Spontaneous breathing over time]

With the Flow trigger threshold, the mandatory breaths are synchronized with the inspiratory efforts. The start setting of the flow trigger can be configured on the System setup > Ventilation > Start settings > VT, RR, Trigger page.

Spontaneous breathing activity by the patient is indicated on the screen by the brief appearance of the symbol.

The flow trigger is automatically leakage compensated.

In order to prevent a possible error when measuring the respiratory rate, e.g., caused by cardiogenic oscillations, only those spontaneous breaths are counted which meet the adjustable trigger criterion.
Inspiratory termination

For spontaneous breaths supported with *PS, VS* and PPS, the length of inspiration is determined by the inspiratory termination criterion. Inspiratory termination specifies at which percentage of the peak inspiratory flow expiration is to start.

This value is set at 15 % by default and is automatically leakage-compensated.
Atelectasis can be prevented by activating the sigh function and setting the sigh in the form of an intermittent PEEP. The purpose of expiratory sigh is to open collapsed areas of the lungs or to keep open "more dependent" areas of the lungs.

The sigh function can be activated in all ventilation modes with mandatory breaths, except for PC-APRV. When the sigh function is activated, the end-expiratory pressure PEEP increases by the set value of the intermittent PEEP.

The time between the two sigh phases can be set with the therapy control Interval sigh.

The therapy control Cycles sigh controls how many respiratory cycles are covered by the sigh phase. The average airway pressure is higher, and a longer filling time is normally available.

During pressure-controlled ventilation, the inspiratory pressures $P_{insp}$, $P_{supp}$ increase by the amount $\Delta \text{intPEEP}$. 
**Principles of operation**

**HFO sigh**

A sigh (RRsigh) can be set during high-frequency oscillation.

**RRsigh** determines how often per minute the sigh is to occur. A conventional pressure-controlled breath is applied with pressure Psigh. The duration of this breath is set with Tisigh. The high-frequency oscillations are discontinued for at least 150 ms prior to the sigh and are resumed 250 ms after the sigh. The high-frequency oscillations start with an expiration.

Rise time and rise form of the sigh are determined by the configuration Slope adjustment (Slope or Insp. flow). If Insp. flow is configured, a constant time slope of 0.1 second is set for Tisigh.

The sigh can also be triggered with the Man. insp./hold function. The duration of the sigh is determined by touching and holding the Man. insp./hold button.
Volume Guarantee

The mandatory breaths are volume controlled with the Volume Guarantee additional setting. To apply the set tidal volume, Babylog VN500 controls the inspiratory plateau pressure $P_{insp}$.

Changes in lung conditions (compliance, resistance) are compensated. The tidal volume of the mandatory breaths remains constant.

Volume Guarantee can be switched on in the PC-SIMV, PC-CMV, PC-AC and PC-PSV ventilation modes. In the PC-MMV and SPN-CPAP/VS ventilation modes, volume guarantee is always available.

The advantage in contrast to time-cycled, pressure-limited ventilation, is that changes in lung conditions (compliance, resistance) have no impact on the tidal volume. If, for example, compliance increases, the inspiratory pressure decreases automatically. If, for instance, compliance decreases, then pressure rises but only up to the set pressure limit $P_{max}$.

If $P_{max}$ is not linked to the alarm limit $P_{aw\, high}$, Babylog VN500 increases $P_{insp}$ up to a maximum of 5 mbar (5 cmH2O) below the set alarm limit $P_{aw\, high}$.

Fluctuations in spontaneous breathing are also compensated. The greater the patient’s inspiratory efforts are, the lower the pressure Babylog VN500 applies. Thus with Volume Guarantee, Babylog VN500 always ventilates with just the right pressure required for the tidal volume desired. The pressure load on the lungs is limited to the extent absolutely necessary.

Without Volume Guarantee, the user must adjust the inspiratory pressure to reach the tidal volume desired.

The control works in the range $PEEP + 0.1$ mbar (0.1 cmH2O) to $P_{max}$ (or $P_{aw\, high} - 5$ mbar (5 cmH2O)) for spontaneous breaths. For triggered mandatory breaths, the control works in the range $PEEP + 3$ mbar (3 cmH2O) to $P_{max}$ (or $P_{aw\, high} - 5$ mbar (5 cmH2O)).
Principles of operation

Using the setting for $P_{\text{max}}$ or the alarm limit $P_{\text{aw high}}$ – 5 mbar (5 cmH₂O), the user limits the maximum pressure of the device.

The set tidal volume cannot be applied under these conditions:
- $P_{\text{max}}$ is insufficient
- The inspiratory pressure pattern has no plateau because the flow is too low or the inspiratory time $T_i$ is too short.

A set inspiratory time $T_i$ shorter than the lung filling time can be recognized from the flow curve. The flow at the end of the inspiratory time has not dropped to zero. In this case, it must be decided whether the current condition of the patient permits prolongation of the inspiratory time $T_i$ in order to reduce peak pressure even further. This effect can also be caused during ventilation, e.g., due to a build-up of secretion. In this situation, the pressure is limited by Babylog VN500 as described.

If the tidal volume measured is below 90 % of the set tidal volume, Babylog VN500 generates an alarm.

The control occurs gradually from breath to breath. The tidal volume is measured, then compared to the set tidal volume and a new plateau pressure is calculated for the next breath. After a change to the set tidal volume, the inspiratory pressure required for this is reached after just a few breaths.

In the Neo. patient category, the tidal volume measured on the expiratory side is taken as a basis for the control. In the Ped. pat. patient category, the inspiratory tidal volume is used. If leakage compensation is activated, the leakage-compensated tidal volume is used for the control.

The minimum inspiratory pressure for mandatory non-triggered breaths is 3 mbar (3 cmH₂O) above PEEP; for triggered mandatory and pressure-supported spontaneous breaths it is 0.1 mbar (0.1 cmH₂O) above PEEP.

In case of major tube leakage, the actual tidal volume in the patient's lungs can (as in other ventilation modes also) be larger than the tidal volume measured on the expiratory side. Then the inspiratory and expiratory tidal volumes are different. If, in the course of an inspiration, the delivered and measured VT exceeds the set VT by an amount dependent on the actual leakage rate, Babylog VN500 terminates the inspiration and starts the expiration.

If the flow sensor fails, ventilation is continued with the pressure used last and Babylog VN500 generates an alarm.

Set the alarm limits $MV_{\text{high}}$ and $MV_{\text{low}}$ appropriately in order to avoid excessive or insufficient flow following rapid changes in compliance. When using Volume Guarantee, activate flow monitoring!

If the Volume Guarantee function in the Ped. pat. and Neo. categories is switched on and a manual inspiration (Man. insp./hold) is triggered, a breath is applied to the maximum pressure $P_{\text{max}}$.

Start-up procedure with volume guarantee

When the Volume Guarantee function is switched on, Babylog VN500 applies the set tidal volume VT by delivering a pressure-controlled breath with an inspiratory pressure of 5 mbar (5 cmH₂O) above the set PEEP. Babylog VN500 measures the applied volume in this case and calculates an initial target pressure for the set volume. The next mandatory breath is applied with an inspiratory pressure that corresponds to 75 % of this target pressure. Babylog VN500 measures the applied volume again and calculates a new target pressure for the set volume. The next mandatory breath is applied with this target pressure. As described above, the following mandatory breaths are changed in the inspiratory pressure so that the set volume is reached on average.
Principles of operation

Volume Guarantee (HFO)

With the volume guarantee of the high-frequency oscillation (VG (HF)), Babylog VN500 calculates the amplitudes required to reach the set tidal volume $V_{Thf}$. The Ampl hf therapy control is inactive when volume guarantee is switched on. If the set $V_{Thf}$ is not reached, Babylog VN500 generates an alarm.

ATC

Automatic Tube Compensation

Compensation of the tube resistance

$\text{ATC}$ regulates the airway pressure to the tracheal level. This function calculates and displays the tracheal pressure on the basis of a mathematical tube model, the set tube type and the inner diameter of the tube.

When tube compensation is activated, Babylog VN500 displays the calculated tracheal pressure in the pressure curve together with the pressure at the Y-piece as a line. Activated tube compensation is indicated by $\text{ATC}$ and the tube diameter in the page header bar.

When selecting loops, tracheal pressure can also be selected as a parameter. Tracheal pressure can also be displayed when tube compensation is deactivated if the calculation of the tracheal pressure was activated on the $\text{Start/Standby} > \text{Tube/NIV}$ page and the tube type and diameter were entered. Babylog VN500 uses this value for calculating leakage and determining the lung mechanics, but not for tube compensation.
The selected degree of compensation is not considered when displaying tracheal pressure or when determining leakage and lung mechanics.

**Calculating tracheal pressure**

Babylog VN500 calculates tracheal pressure on the basis of a square function of tube resistance and patient flow.

\[
P_{\text{trachea}} = P_{\text{aw}} - K_{\text{tube}} \times \text{Flow} \times |\text{Flow}|
\]

- **P_{\text{trachea}}**: Pressure in the trachea
- **P_{\text{aw}}**: Pressure at the Y-piece of the breathing circuit
- **K_{\text{tube}}**: Tube coefficient (see page 314)
- **Flow**: Patient flow
  - Inspiration: Flow >0
  - Expiration: Flow <0

The maximum value for the airway pressure can be set using the \( P_{\text{max}} \) therapy control. If \( P_{\text{max}} \) is not linked to the alarm limit \( P_{\text{aw high}} \), the maximum pressure is limited to 5 mbar (5 cmH2O) below the alarm limit \( P_{\text{aw high}} \). The pressure limitation message is displayed when the maximum permitted values are reached.

If the value selected for \( P_{\text{aw high}} \) or \( P_{\text{max}} \) is too low, it may impair the effectiveness of tube compensation. If the value selected for \( P_{\text{aw high}} \) or \( P_{\text{max}} \) is too high, it may result in unwanted high airway pressures. When setting \( P_{\text{max}} \), be aware that this value may actually be reached in contrast to the value for \( P_{\text{aw high}} \).

**Calculating the support**

The level of support \( \Delta P_{\text{aw}} \) applied during ATC is calculated on the basis of a square law function of tube resistance and patient flow.

\[
\Delta P_{\text{aw}} = \text{Comp.} \times K_{\text{tube}} \times \text{Flow} \times |\text{Flow}|
\]

- **\( \Delta P_{\text{aw}} \)**: Degree of compensation 0 to 100 %
- **K_{\text{tube}}**: Tube coefficient (see page 314)
- **Flow**: Patient flow

**Tube coefficient**

The tube coefficient \( K_{\text{tube}} \) is largely determined on the basis of the results obtained by Guttmann et al*.

The tube coefficient \( K_{\text{tube}} \) for the full-length tube is always taken as the basis. The effect of the shortened length is negligible.

---

* Literature reference [5], see page 336
The values for the tube coefficients are shown in the following tables.

### Table for endotracheal tube:

<table>
<thead>
<tr>
<th>Inner diameter of the tube (mm)</th>
<th>Tube coefficient $K_{\text{Tube}}$ (mbar/L²/s²)</th>
</tr>
</thead>
<tbody>
<tr>
<td>2.00</td>
<td>1834.00</td>
</tr>
<tr>
<td>2.50</td>
<td>600.00</td>
</tr>
<tr>
<td>3.00</td>
<td>340.00</td>
</tr>
<tr>
<td>3.50</td>
<td>170.00</td>
</tr>
<tr>
<td>4.00</td>
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</tr>
<tr>
<td>4.50</td>
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</tr>
<tr>
<td>5.00</td>
<td>30.96</td>
</tr>
<tr>
<td>5.50</td>
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</tr>
<tr>
<td>6.00</td>
<td>17.21</td>
</tr>
<tr>
<td>6.50</td>
<td>13.05</td>
</tr>
<tr>
<td>7.00</td>
<td>10.56</td>
</tr>
<tr>
<td>7.50</td>
<td>8.41</td>
</tr>
<tr>
<td>8.00</td>
<td>6.57</td>
</tr>
</tbody>
</table>

### Table for tracheostomy tube:

<table>
<thead>
<tr>
<th>Inner diameter of the tube (mm)</th>
<th>Tube coefficient $K_{\text{Tube}}$ (mbar/L²/s²)</th>
</tr>
</thead>
<tbody>
<tr>
<td>2.50</td>
<td>600.00</td>
</tr>
<tr>
<td>3.00</td>
<td>340.00</td>
</tr>
<tr>
<td>3.50</td>
<td>170.00</td>
</tr>
<tr>
<td>4.00</td>
<td>100.00</td>
</tr>
<tr>
<td>4.50</td>
<td>50.00</td>
</tr>
<tr>
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<td>30.96</td>
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</tr>
<tr>
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</tr>
<tr>
<td>6.50</td>
<td>7.90</td>
</tr>
<tr>
<td>7.00</td>
<td>6.38</td>
</tr>
<tr>
<td>7.50</td>
<td>5.20</td>
</tr>
<tr>
<td>8.00</td>
<td>4.50</td>
</tr>
</tbody>
</table>
In ventilation mode PC-APRV, the duration of pressure release is determined from the expiratory flow curve when AutoRelease is activated. The Exp. term. setting specifies when the ventilation returns to the pressure level $P_{high}$ dependent on the decline in percent of the peak expiratory flow. The therapy control $P_{high}$ limits the maximum duration of pressure release.

When AutoRelease is switched on, the change-over from the upper pressure level $P_{high}$ to the lower pressure level $P_{low}$ is synchronized with the patient's spontaneous breathing.

Synchronization of the mandatory breath reduces the effective time of the upper pressure level. Babylog VN500 prolongs the subsequent ventilation time on the upper pressure level by the missing time.
Special functions

Medication nebulization

Insp. O₂ concentration during medication nebulization

Only use medication nebulizer 8411030. If other medication nebulizers are used, considerable deviations may occur in the tidal volume and the inspiratory O₂ concentration!

To minimize the deviation from the set O₂ concentration, Babylog VN500 uses a gas mixture to drive the medication nebulizer. The gas mixture is generated by switching over between compressed air and O₂ in short time intervals.

The medication nebulizer nebulizes continuously. The aerosol generated during expiration does not reach the lungs, however. The medication nebulizer is supplied with compressed air, O₂ or a mixture of compressed air and O₂ by Babylog VN500, depending on the set O₂ concentration.

The graph shows the possible deviations of the applied O₂ concentration from the set FiO₂ concentration with respiratory rates above 12/min.
Air supply from the GS500 gas supply unit

If Babylog VN500 is supplied with Air from the GS500 gas supply unit and O2 is supplied from the central gas supply system, the medication nebulizer operates with O2 only. The measured value $\text{FiO}_2$ indicates the O2 concentration of the gas supplied at the inspiratory port and not the O2 concentration reaching the patient. Depending on the patient category, the following systematic deviations are possible:

<p>| | |</p>
<table>
<thead>
<tr>
<th></th>
<th></th>
</tr>
</thead>
<tbody>
<tr>
<td><strong>Ped. pat.</strong></td>
<td>up to +30 Vol%</td>
</tr>
<tr>
<td><strong>Neo.</strong></td>
<td>up to +30 Vol%</td>
</tr>
</tbody>
</table>

C20/C

The C20/C index is a calculation of the compliance of the last 20 % (C20) of a breath in relation to the compliance (C) of the entire breath.

During a breath, Babylog VN500 determines continuously the pressure applied and the resulting tidal volume. The compliance of the last 20 % of a breath determined in this manner is set in proportion to the total compliance.

From the ratio determined, the following information can be derived:

- C20/C < 1: A decrease of compliance at the end of the breath was detected. The lungs may be overinflated.
- C20/C > 1: An increase of compliance at the end of the breath was detected. Tidal recruitment may be occurring.
- C20/C = 1: No change in compliance at the end of a breath could be detected. The lungs may not be overinflated, or tidal recruitment may not be occurring.

The calculation of C20/C takes into account the effect of the resistance of the endotracheal tube used or the tracheostomy tube used. For this, the tube diameter is required. The correct tube diameter entry of the tube used determines the quality of the C20/C index calculated.

The C20/C index is always displayed as long as a correction delivers plausible results with regard to the resistance. If, for instance, a smaller tube diameter was entered than that of the tube actually used, a correction to the measured values may deliver an implausible result. In this case, no C20/C index is displayed. The parameter field remains empty.
**Smart Pulmonary View**

Graphic display of lung characteristics

Smart Pulmonary View is a graphic display of lung flexibility (compliance) and resistance of the airways (resistance).

The representation corresponds to the displayed measured values of the respective patient.

The display range of compliance is 0 to 400 mL/mbar (400 mL/cmH2O).

The display range of resistance is 0 to 300 mbar/L/s (300 cmH2O/L/s).

To detect an improvement or deterioration of the patient's condition with regard to compliance and resistance, it is possible to adapt the representation to the current values of the patient. One measuring range starts at 0 and goes to double the value of the current compliance; the other measuring range starts at 0 and goes to double the value of the current resistance. After the adaptation, the measuring values determined are displayed as reference values with the time and date. In the graph, the current values (calibration values) are displayed as an orange broken line. The scales for compliance and resistance are adapted.

The compliance and resistance measured respectively are displayed by thin or thick lines accordingly.

The point when the maximum value that is based on the last calibration is reached is represented with a red line as a boundary. This indicates that the measured values determined can no longer be represented graphically. The measured values are outside the display range. Babylog VN500 displays a request for a new calibration.

The diaphragm is displayed schematically underneath the representation of the lungs. The movement of the diaphragm indicates synchronized mandatory breaths, supported (triggered) breaths, or spontaneous breaths.

The ratio between spontaneous breathing and mandatory ventilation is displayed in a diagram:

- $RR^\text{spont}$ and $VT^\text{spont}$ represent the spontaneous minute volume as an area
- $RR^\text{mand}$ and $VT^\text{mand}$ represent the mandatory minute volume as an area

The display is a qualitative representation of the respective minute volume.

From this, the following information can be derived:

- The ratio between the spontaneous and mandatory minute volumes
- The quality and pattern of the spontaneous breathing, e.g., Rapid Shallow Breathing

Smart Pulmonary View is a qualitative representation of the ventilation situation. Local pathophysiological peculiarities, such as atelectasis or airway obstructions of the lungs, cannot be displayed.

Furthermore, individual patient situations cannot be displayed, such as the condition after a pneumectomy or a diaphragmatic hernia.
Description of the therapy types

**O2 therapy**

O2 therapy can be used for patients with independent breathing. The continuous flow is applied via an oxygen mask, a hood, or nasal cannula. The O2 concentration and the flow can be set.

**NIV – Non-invasive ventilation**

Non-invasive ventilation by prongs or mask for patients with spontaneous breathing.

Leakages are greater with non-invasive ventilation than with invasive ventilation. Babylog VN500 takes into account the leakages in the NIV application mode accordingly.

The **Tdisconnect** setting can be used to delay the **Airway pressure low** alarm.

In the **Neo.** patient category, only the SPN-CPAP or PC-CMV ventilation modes may be selected. When using prongs or a mask, the neonatal flow sensor must be removed from the breathing circuit. Babylog VN500 switches off flow monitoring with the neonatal flow sensor.
Automatic leakage compensation

**Mode of operation**

Babylog VN500 determines the difference between the delivered inspiratory flow and the measured expiratory flow. This difference provides a measure of the amount of leakage and is displayed by Babylog VN500 as the leakage minute volume $\text{MV_{leak}}$ and relatively as $\% \text{ leak}$ ($\text{MV_{leak}}$ to $\text{MVi}$).

The calculation of leakage compensation takes into account the airway pressures. A higher percentage of volume is lost on the inspiratory side than on the expiratory side because the pressure during inspiration is higher. The displayed leakage minute volume $\text{MV_{leak}}$ is based on the mean pressure $P_{\text{mean}}$. The leakage minute volume $\text{MV_{leak}}$ also takes the inspiratory leakages into account. Due to technical tolerances, a small leakage minute volume may be displayed even if the tube leakage is closed. If there is a rapid change in the leakage, e.g., due to the leak being opened or closed suddenly, Babylog VN500 needs a few breaths to identify the new leakage value. Babylog VN500 prevents any potentially dangerous rises in pressure which might occur as a result of this.

The inspiratory flow trigger threshold and the inspiratory termination criterion are applied to the leakage-compensated flow, with both settings being continuously optimized with regards to the leakage. This automatic adjustment also takes place if leakage compensation is deactivated.

If leakage compensation is activated, the values measured for volume and flow as well as the curves for flow and volume are displayed with leakage correction, with the exception of the minute volume measured during expiration and all measured values which are explicitly marked as inspiratory or expiratory, such as $\text{VTi}$ and $\text{VTe}$.

Activate or deactivate leakage compensation on the **System setup > Ventilation > General settings** page.

**Example of leakage compensation with flow trigger or inspiratory termination criterion**

The mode of operation is illustrated using a simplified example with the following values:

- Flow trigger setting 0.2 L/min
- Leakage increases from 0 % to 20 %

Mode of operation with leakage compensation: Babylog VN500 determines the leakage flow. The leakage flow is subtracted from the total flow in order to determine the patient flow. Only this flow is used for the flow trigger or the inspiratory termination criterion. After a few breaths Babylog VN500 "learns" the leakage and avoids auto-triggering. If the leakage is closed, the sensitivity of the flow trigger is automatically increased again. The same applies to the inspiratory termination criterion for breaths with pressure support or volume support.

**Leakage rate**

Babylog VN500 determines the mean leakage flow from the difference between inspiratory minute volume $\text{MVi}$ and expiratory minute volume $\text{MVe}$ (displayed as $\text{MV}$). Standardized as $\text{MVi}$, the result is the leakage rate displayed in percent:

$$\text{Leakage rate} = 100 \times \frac{\text{MVi} – \text{MVe}}{\text{MVi}}$$
Flow reduction *Anti Air Shower*

When the *Anti Air Shower* function is activated and a disconnection is detected during ventilation, the flow is reduced until reconnection is detected. The *Disconnection*? alarm is displayed at the same time. With non-invasive ventilation, the time before the alarm is triggered can be delayed with *Tdisconnect*. The minute ventilation can be reduced due to the already reduced flow.

To configure the *Anti Air Shower* function, see "Configuring general settings" on page 177.
Principles of operation

Measurements

Measurement principles

Measurement principle of the flow measurement

The measurement principle used for flow measurement is based on hot-wire anemometry. Hot-wire anemometry is a thermal measurement procedure in which the measuring wires of the flow sensor are kept at a constant excess temperature. The higher the flow, the more current is required to maintain a constant excess temperature. The flow rate is calculated based on the magnitude of the heating current.

To ensure correct function, check for visible damage, soiling, and particles before inserting the flow sensor. Repeat this check regularly. Replace flow sensors when damaged, soiled, or not particle-free. If the measurement wires of the flow sensor glow continuously during operation, this is an indication of contamination. Immediately reprocess or replace the flow sensor.

Flow measurement with neonatal flow sensor

The flow is measured with a hot wire anemometer between the Y-piece and the tube. The flow direction is detected by the use of two hot wires, one of which is shielded on one side.

The amount of energy required to maintain the wire at a temperature of 400 °C (752 °F) is used as a measure of the flow passing through the sensor, cooling the hot wire in the process.

The lowest flow at which detection functions reliably is 0.2 L/min. Lower flow values are therefore suppressed and displayed as zero.

Two different sensor types are available:
- Y-sensor, integrated in the Y-piece
- ISO sensor to insert between Y-piece and tube connector

Both sensor types use the same sensor insert. Despite this, the sensor properties are not identical. The sensor type is set in the Sensors/Parameters > Neonatal flow sensor dialog window in order to adapt the measurement for this type of sensor optimally.

O2 measurement

A heating and a temperature sensor are positioned in a homogeneous magnetic field which is periodically activated and deactivated. The thermal conductivity of O2 changes due to the magnetic field. The change in thermal conductivity is a measure of the O2 concentration.

CO2 measurement

CO2 is measured via a mainstream system based on absorption measurement. A light source generates a spectrum. Two light detectors record the characteristic absorption spectrum and supply electrical signals that change with the CO2 concentration.

These signals are then evaluated and displayed. Heating the CO2 sensor probe prevents condensation.
Airway pressure measurement

Babylog VN500 measures the airway pressure indirectly by means of two internal pressure sensors. The sensors are installed in the inspiratory and expiratory lines, thereby eliminating the need for an external pressure measuring line between the Y-piece and the device. As long as one side is without flow, the measured value of the flowless pressure sensor corresponds to the airway pressure at the Y-piece.

During ventilation, there is a constant base flow. However, due to this constant base flow, the zero-flow condition is never attained either on the inspiratory or expiratory side. The pressure measured by the inspiratory pressure sensor varies with the variations in airway pressure but is increased by the pressure drop in the inspiratory line of the breathing circuit. The pressure measured by the expiratory pressure sensor is reduced by the pressure drop in the expiratory line of the breathing circuit. These pressure differences are caused by the flow resistance of the breathing circuit.

During expiration, the value measured at the inspiratory pressure sensor ($P_{\text{insp}}$) is reduced by the pressure drop caused by the base flow ($\text{Flow}_{\text{bf}}$) in the inspiratory line of the breathing circuit ($R_{\text{insp}}$):

$$P_{\text{aw}} = P_{\text{insp}} - R_{\text{insp}} \times \text{Flow}_{\text{bf}}$$

During inspiration, the value measured by the expiratory pressure sensor ($P_{\text{exp}}$) is raised relative to the airway pressure by the pressure drop caused by the flow (normally $\text{Flow}_{\text{out}} \leq \text{Flow}_{\text{bf}}$) in the expiratory line of the breathing circuit ($R_{\text{exp}}$):

$$P_{\text{aw}} = P_{\text{exp}} + R_{\text{exp}} \times \text{Flow}_{\text{out}}$$

Adaptation to ambient conditions

The volume of a gas depends on the ambient conditions with regard to temperature, pressure, and humidity. In lung physiology, reference is made to the conditions inside the lungs for the values of minute volume and tidal volume: 37 °C (99 °F) body temperature, pressure inside the lungs, 100 % relative humidity.

Measured values for flow and volume under these conditions are characterized as BTPS. Medical gases from cylinders or from a central supply are dry (approximately 0 % relative humidity) and are delivered from the ventilator at 20 °C (68 °F) and 1013 mbar (1013 cmH2O). Measured values for flow and volume under these conditions are characterized as NTPD.

The difference between NTPD and BTPS measured values is approximately 12 % at a pressure of 1013 mbar (1013 cmH2O).

Example: 250 mL tidal volume NTPD become 282 mL BTPS when warmed to 37 °C (99 °F) and humidified to 100 % relative humidity.

Babylog VN500 controls tidal volume in such a way that the set tidal volume value is applied under BTPS conditions in the lungs.
Battery concept

General information
At the time of manufacture and delivery, batteries have a typical capacity which is in accordance with the information specified in the battery manufacturer's data sheet. The electrochemical composition of the battery is the determining factor for its total capacity. The operating time of the batteries derived from these specifications can be found in the "Technical data" chapter.

NOTE
The capacity of batteries changes with increasing age and utilization.

All the following information and specifications refer to perfectly functioning batteries. If batteries are defective or faulty, the functional integrity, e.g., battery charge indication or alarms, may be impaired. See chapter "Battery check" on page 254.

Display of battery charge
The battery charge indication shows the available battery charge determined by the electrochemical processes. When the batteries are, e.g., fully charged, this state is indicated by a corresponding symbol.

### Symbol Battery charge

<table>
<thead>
<tr>
<th>Symbol</th>
<th>Battery charge</th>
</tr>
</thead>
<tbody>
<tr>
<td>🌟🌟🌟🌟</td>
<td>90 to 100 %</td>
</tr>
<tr>
<td>🌟🌟🌟</td>
<td>60 to &lt;90 %</td>
</tr>
<tr>
<td>🌟🌟</td>
<td>40 to &lt;60 %</td>
</tr>
<tr>
<td>🌟</td>
<td>20 to &lt;40 %</td>
</tr>
<tr>
<td>🌟</td>
<td>&lt;20 %, flashes light and dark red in 1-second pulses.</td>
</tr>
<tr>
<td>🌟</td>
<td>Batteries defective or no information available on the battery charge.</td>
</tr>
</tbody>
</table>

The battery charge indication is a relative indication which is based on the electrochemical properties of the battery. The battery charge indication is evaluated on the basis of a battery model.

The use of a model-based indication is a state-of-the-art technique which finds application in many fields, e.g., computers, mobile phones, etc.

The model-based indication of battery charge takes account of the following information, among other things:
- Type of battery (e.g., NiMH or VRLA)
- Maximum capacity on delivery (e.g., 12 Ah)
- Age of the battery (e.g., new or 2 years)
- Capacity spent (irreversibly lost) over the utilization time (e.g., 1000 Ah)
- Present power requirement of the device (power consumption, e.g., 2.5 A)
- Discharge mode
- Charging mode

If the power consumption changes, e.g., due to switching to Standby, operation of a GS500, or adjustment of the screen brightness, the remaining available operating time of the device also changes. The battery charge indication is updated to take account of the present power requirement (power consumption).

In accordance with the specification, the battery charge indication is only displayed and updated after the device has been completely started up. This procedure may take a few minutes.

Battery ageing
The electrochemical composition of a battery alters as a result of ageing and utilization. Consequently, every battery loses a proportion of its maximum capacity in comparison with its new condition. This loss of capacity is typically irreversible.

As a result of ageing and utilization of the battery, there is a change in the actual maximum operating time which is displayed by the percentage values in
the battery charge indication. The percentage value refers to the currently available operating time of batteries which were fully charged before use.

New batteries
The following data for minimum operating time apply to new and fully charged batteries. The symbol for a fully charged battery is displayed. See also the "Display of battery charge" chapter and the "Technical Data" chapter. Owing to production fluctuations during the manufacture of batteries, the operating time can be considerably longer.

<table>
<thead>
<tr>
<th>Battery used (Battery type)</th>
<th>Minimum operating time without operation of a GS500</th>
<th>Minimum operating time with operation of a GS500</th>
</tr>
</thead>
<tbody>
<tr>
<td>Internal battery</td>
<td>30 min</td>
<td>15 min</td>
</tr>
<tr>
<td>PS500</td>
<td>240 min</td>
<td>120 min</td>
</tr>
</tbody>
</table>

Aged batteries, e.g., 2 years old
The following data for minimum operating time apply, e.g., to 2-year old and fully charged batteries. The data are approximate values and cannot be regarded as guaranteed for every battery. The symbol for a fully charged battery is displayed. See also the "Display of battery charge" chapter and the "Technical Data" chapter.

<table>
<thead>
<tr>
<th>Battery used (Battery type)</th>
<th>Minimum operating time without operation of a GS500</th>
<th>Minimum operating time with operation of a GS500</th>
</tr>
</thead>
<tbody>
<tr>
<td>Internal battery</td>
<td>22 min</td>
<td>11 min</td>
</tr>
<tr>
<td>PS500</td>
<td>120 min</td>
<td>60 min</td>
</tr>
</tbody>
</table>

NOTE
A reduction in the capacity of batteries due to ageing and utilization is normal. As a greatly simplified approximation, an average linear reduction in capacity can be assumed. The current individual capacity of a battery depends on the following factors, among others:
- Age
- Utilization (frequency, duration, and power consumption)
- Battery charge
- Ambient temperature

Spent batteries
When the internal battery and the batteries in the PS500 show less than the following residual operating times, they are considered to be spent.

<table>
<thead>
<tr>
<th>Battery used (Battery type)</th>
<th>Spent with remaining residual operating time without operation of a GS500</th>
</tr>
</thead>
<tbody>
<tr>
<td>Internal battery</td>
<td>&lt;22 min</td>
</tr>
<tr>
<td>PS500</td>
<td>&lt;120 min</td>
</tr>
</tbody>
</table>

Alarm behavior in battery operation
The switch-over of the device to battery operation is indicated by the Battery activated alarm (see chapter "Alarm – Cause – Remedy"). The alarm can be acknowledged by the user. Consequently the Battery activated alarm will no longer be displayed until the mains power supply is re-established.

When the device is equipped with a PS500 power supply unit, in battery operation the PS500 is discharged first. If the mains power supply has not been re-established, the device switches over to the internal battery after the operating time of the PS500 has elapsed. The switch-over is indicated by the Internal battery activated alarm.
Principles of operation

At the end of the operating time of the internal battery, the device generates the **Battery low** alarm. The **Battery discharged** alarm follows after that.

These alarms appear at the time specified by the model-based calculation for the particular battery.

### Schematic representation of the sequence of alarms

![Diagram showing the sequence of alarms]

The schematic representation of the sequence of alarms with respect to battery utilization is shown in an example with a PS500 but without the use of a GS500. The representation corresponds to the operating time of new and fully charged batteries.

When the voltage drop of the internal battery reaches an operationally critical value, a shut-down of the device due to an inadequate supply is immediately imminent. In this case, the power supply failure alarm sounds.

**NOTE**

If the device displays the **Battery low** alarm message or the **Battery discharged** alarm message, connect the device to the mains power supply.

**NOTE**

When the remaining calculated operating time is less than 10 minutes, the model-based **Battery low** alarm appears. When the remaining calculated operating time is less than 5 minutes, the model-based **Battery discharged** alarm appears.
NOTE
The operating time remaining after the corresponding alarms can be considerably longer than the specified minimum operating time.

NOTE
When the device is fitted with the GS500 gas supply unit, the device calculates the time for the Battery discharged alarm allowing for the power consumption of a GS500, regardless of whether the GS500 is activated or not.
Principles of operation

Pneumatic functional description

Pneumatic circuit diagram of Babylog VN500

1 Air gas inlet
2 O2 gas inlet
3 Air non-return valve
4 O2 non-return valve
5 Air metering valve
6 O2 metering valve
7 Tank
8 Mixed gas metering valve
9 Safety valve
10 Emergency expiratory valve
11 Emergency breathing valve
12 Patient's lungs
13 Expiratory valve
14 Non-return valve
15 Neonatal flow sensor
16 Barometric pressure sensor
17 Calibration valve for inspiratory pressure sensor
18 Inspiratory pressure sensor
19 Calibration valve for expiratory pressure sensor
20 Expiratory pressure sensor
21 O2 sensor
22 Nebulizer outlet
23 Air pressure regulator
24 O2 pressure regulator
25 Nebulizer mixer valve
26 Nebulizer changeover valve
27 Ejector changeover valve
28 Ejector
29 Muffler
30 CO2 sensor
Principles of operation

Babylog VN500 consists of 9 pneumatic assemblies.

**A** Gas mixture and gas metering assembly
**B** Inspiratory unit assembly
**C** Expiratory unit assembly
**D** Muffler
**E** Barometric pressure sensor
**F** Pressure measurement assembly
**G** Calibration assembly
**H** O2 sensor
**I** Medication nebulization assembly/Ejector drive

**Description of the pneumatic mode of operation**

Babylog VN500 delivers the time-variable flow of a gas mixture with adjustable proportions of O2 and air. Gas from the central gas supply system enters the device via the gas inlet connections for O2 and air (1, 2). Two non-return valves (3, 4) prevent one gas from returning to the supply line of the other gas. The mixing of the gases takes place in the tank (7) and is controlled via two control valves (5, 6). The supplied inspiratory flow is controlled via a third control valve (8).

The **inspiratory unit assembly** (B) consists of the safety valve (9) and two non-return valves (10, 11).

In normal operation, the safety valve is closed so that the inspiratory flow is supplied to the patient (12) from the gas mixture and gas metering assembly. During other operating states, e.g., when Babylog VN500 is in standby, the safety valve is open and enables spontaneous inspiration through the emergency breathing valve (11). The emergency expiratory valve (10) provides a second channel for expiration when the expiratory valve (13) is blocked.

The **expiratory valve assembly** (C) consists of the expiratory valve (13) and a non-return valve (14).

The expiratory valve is a proportional valve and is used to adjust the pressure in the breathing system. In conjunction with the spring-loaded valve of the emergency air outlet (10), the non-return valve (14) prevents pendulum breathing during spontaneous breathing. The neonatal flow sensor (15) measures the inspiratory flow and expiratory flow in accordance with the hot-wire anemometry measurement principle. Therefore the measured flow is a mass flow (NTPD).

The ejector (28) generates the negative pressure required for the HFO mode. For this purpose, the ejector valve (27) supplies the driving gas (medication nebulization assembly/ejector drive (I)).

To reduce disruptive noises, the flow is passed behind the expiratory valve (13) via the muffler (D, 29) into the surrounding area.

The inspiratory unit, the expiratory valve, and the muffler assemblies can be detached from Babylog VN500 for cleaning purposes.

The mass flow to volume flow conversion (BTPS) requires knowledge of the ambient pressure. The ambient pressure is measured with the **barometric pressure sensor** (E, 16).

The pressure in the breathing system is measured with two independent pressure sensors (18, 20) that form the **pressure measurement assembly** (F). The pressure sensors are regularly zero calibrated. For this, the pressure sensors are connected to ambient pressure via the two calibration valves (17, 19). The calibration valves form the **calibration assembly** (G).

The **O2 sensor** (H, 21) measures the inspiratory O2 concentration based on a sidestream measurement principle. For calibration by the user during the device check, the O2 sensor can be flushed with pure O2 from the tank (7).
A pneumatic medication nebulizer can be connected to the nebulizer gas outlet (22) for medication nebulization. Babylog VN500 provides an intermittent gas flow consisting of O2 and air to drive the medication nebulizer. This ensures that the deviation of the set O2 concentration remains within the specified limits. For this, the gas from the two gas inlet connections (1, 2) is throttled by the pressure regulators (23, 24). The intermittent gas delivery is done by nebulizer mixer valve (25). The nebulizer changeover valve (26) closes the nebulizer gas outlet when the nebulizer function is not switched on.

The nebulizer mixer valve, the nebulizer changeover valve, the ejector changeover valve, the nebulizer outlet and the two pressure regulators form the medication nebulization/ejector drive assembly (I).

The CO2 concentration of the breathing gas can be measured using the CO2 sensor (30). CO2 is measured according to an optical measurement principle in the mainstream.

An active breathing gas humidifier and a pneumatic medication nebulizer can also be connected.

Additional information can be found in the chapters "Assembly and preparation" and "Operation".
**Main menu bar structure**

The following table lists the buttons of the main menu bar with the resulting dialog windows of the same name and the tabs. Touching a tab opens the corresponding page. The dark gray buttons are always contained in the main menu bar. The white buttons are freely configurable and are assigned to the respective group. The freely configurable buttons open the corresponding page in the dialog window or activate a function.

<table>
<thead>
<tr>
<th>Group symbol</th>
<th>Button in main menu bar</th>
<th>Horizontal tab</th>
<th>Vertical tab</th>
<th>Additional tabs</th>
</tr>
</thead>
<tbody>
<tr>
<td>![Alarm icon]</td>
<td>Alarms...</td>
<td>Limits</td>
<td>Current alarms</td>
<td></td>
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<tr>
<td></td>
<td></td>
<td></td>
<td>Alarm history</td>
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<td></td>
<td>Settings</td>
<td></td>
</tr>
<tr>
<td>![Volume icon]</td>
<td>Alarm volume</td>
<td>Modes 1, 2, 3, 4</td>
<td></td>
<td></td>
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<tr>
<td>![Ventilation icon]</td>
<td>Ventilation settings...</td>
<td>General settings</td>
<td>Overview</td>
<td></td>
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<td>Additional settings</td>
<td>Apnea Ventilation</td>
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<td>Trigger</td>
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<td>Sigh</td>
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<td>Volume Guarantee</td>
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<td>ATC</td>
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<td>Auto Release</td>
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<td>Mode 5 (e.g. PC-HFO)</td>
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<tr>
<td>![Trigger icon]</td>
<td>Trigger</td>
<td>Other modes</td>
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<td></td>
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<tr>
<td>![Apnea Ventilation icon]</td>
<td>Apnea Ventilation</td>
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</table>

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### Principles of operation

<table>
<thead>
<tr>
<th>Group symbol</th>
<th>Button in main menu bar</th>
<th>Horizontal tab</th>
<th>Vertical tab</th>
<th>Additional tabs</th>
</tr>
</thead>
<tbody>
<tr>
<td>Views...</td>
<td></td>
<td>Trends</td>
<td></td>
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<tr>
<td>Day/Night</td>
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<tr>
<td>Freeze waveforms</td>
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<tr>
<td>Export screenshot</td>
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<tr>
<td>Main screen</td>
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<tr>
<td>Trends/Data...</td>
<td>Trends</td>
<td></td>
<td>Values</td>
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<td>Trends table</td>
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<td>Values</td>
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<td>Logbook</td>
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<td>Customized data</td>
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<td>Special maneuvers...</td>
<td>Manoevers</td>
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<tr>
<td>Nebulization</td>
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<tr>
<td>O2 suction</td>
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<tr>
<td>Man. insp./hold</td>
<td></td>
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<tr>
<td>Sensors/ Parameters...</td>
<td>Neonatal flow sensor</td>
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<tr>
<td>Neonatal flow sensor</td>
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<tr>
<td>O2 sensor</td>
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<tr>
<td>CO2 sensor</td>
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<tr>
<td>Zero calib. on/off</td>
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Password for Babylog VN500 SW 2.n

Cut out from the Babylog VN500 instructions for use SW 2.n

To prevent unauthorized adjustments, the following pages are password-protected:

– **System setup > Screen layout > Views**
– **System setup > Alarms**
– **System setup > Ventilation**
– **System setup > Applications**
– **System setup > Exchange intervals**

Information on the password

To prevent unauthorized adjustments, the following pages in the **System setup** dialog window are password-protected:

– **Screen layout > Views**
– **Alarms**
– **Ventilation**
– **Applications**
– **Exchange intervals**

The password appears on this page of the instructions for use. Cut out the area with the password and keep in a place which is safe from access by unauthorized persons.

If the area with the password has been removed, ask the person responsible for your device about making adjustments to the pages specified.
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This page has been left blank intentionally.
These instructions for use only apply to Babylog VN500 SW 2.n with the Serial No.: If no Serial No. has been filled in by Dräger, these instructions for use are provided for general information only and are not intended for use with any specific medical device. These instructions for use are provided for customer information only and will only be updated or exchanged upon customer request.

Directive 93/42/EWG concerning medical devices

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