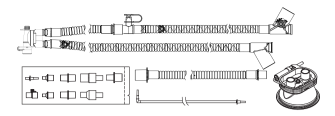
### **For Neonatal**

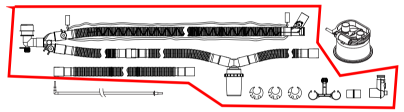


### 51051031

Model No.	Part No.	Tube length	Spare limb	Pressure line	Chamber VHC10	Adaptor kits	Box Qty
510-009	51005918	1.5m	0.6m				10
	51051033	1.5m	0.6m		$\sqrt{}$		10
510-009	51005928	1.5m	0.6m	$\sqrt{}$			10
	51005923	1.5m	0.6m	$\sqrt{}$	$\sqrt{}$		10
	51051031	1.5m	0.6m	$\sqrt{}$	$\checkmark$	$\sqrt{}$	10

### **Single Heated Breathing Circuits**

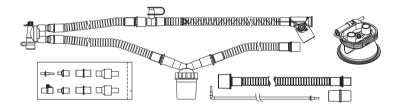
### **For Adult**



51050941

Model No.	Part No.	Tube length	Spare limb	Pressure line	Chamber VHC20	Catheter Mount	Box Qty
510-008	51048400	1.5m	0.45m				10
	51050941	1.5m	0.45m	$\sqrt{}$	$\sqrt{}$		10
	51006113	1.5m	0.45m		$\sqrt{}$	$\checkmark$	10

### **For Neonatal**



### 51051030

Model No.	Part No.	Tube length	Spare limb	Pressure line	Chamber VHC10	Adaptor kits	Catheter Mount	Box Qty
510-009	51005924	1.5m	0.6m	$\sqrt{}$				20
	51005921	1.5m	0.6m		$\sqrt{}$			10
	51005922	1.5m	0.6m	$\sqrt{}$	$\sqrt{}$			10
510-009	51005933	1.5m	0.6m	$\sqrt{}$		$\sqrt{}$		10
	51051030	1.5m	0.6m	$\sqrt{}$	$\sqrt{}$	$\sqrt{}$		10
	* 51006114	1.5m	0.6m		$\sqrt{}$	$\sqrt{}$	$\sqrt{}$	10
510-009	* 51051064	1.5m	0.6m			$\sqrt{}$		10

<sup>\*</sup>PN 51006114 & 51051064 adaptor kit contains 10 adaptors.

### **Humidification Chambers**









VHC10

VHC20

VHC25

VHC50

Model No.	VHC10	VHC20	VHC25	VHC50
Part No.	51003667	51002881	51002689	51007529
Туре	Disposable	Disposable	Reusable	Disposable
Application	Infant	Adult/Pediatric	Adult	Adult
Fill Water Type	Automatic	Automatic	Manual	Manual
Interface Connections	ISO 5356-1	ISO 5356-1	ISO 5356-1	ISO 5356-1
interface confidentions	(22 mm Male)	(22mm Male)	(22mm Male)	(22mm Male)
Compressible Volume (Empty)	300 mL	440 mL	640 mL	690 mL
Compressible Volume (Full)	120 mL	260 mL	140 mL	190 mL
Maximum Water Capacity	180 mL	180 mL	500 mL	500 mL
Box Qty	10	10	10	10
			J	





### nCPAP Patient Interface

Model No.	Part No.	Configuration	Box Qty
504-NCG	51005209	nCPAP Generator	50
504-NCK(4)	51005286	nCPAP Generator+ S/M/L Nasal prong	50

### **nCPAP Nasal Prongs**



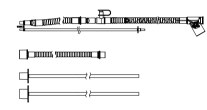
Model No.	Part No.	Description	Box Qty
504-NP(XS)	51005454	nCPAP nasal prong XS	100
504-NP(S)	51005467	nCPAP nasal prong S	100
504-NP(M)	51005468	nCPAP nasal prong M	100
504-NP(L)	51005469	nCPAP nasal prong L	100

### **nCPAP Nasal Masks**



Model No.	Part No.	Description		Box Qty
504-NM(S)	51005455	nCPAP nasal mask S		100
504-NM(M)	51005456	nCPAP nasal mask M		100
504-NM(L)	51005457	nCPAP nasal mask L		100
504-NM(XL)	51005458	nCPAP nasal mask XL	$\bigcirc$	100

### Neonatal Heated Breathing Circuits for nCPAP



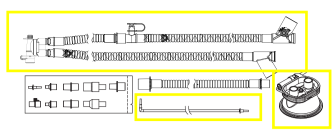
Model No.	Part No.	Description	Box Qty
510-038	51005856	Single limb, 2pcs PVC tubing, generator connector	20
510-038	51005858	Single limb, generator connector	20
511-038-C10	51007960	Single limb, 2pcs PVC tubing, generator connector, VHC10	10

### Infant Bonnets



Model No.	Part No.	Size	Colour	Box Qty
504-B(000)	51005300	16-18cm	$\bigcirc$	100
504-B(00)	51005302	18-20cm		100
504-B(0)	51005303	20-22cm		100
504-B(1)	51005304	22-24cm		100
504-B(2)	51005305	24-26cm		100
504-B(3)	51005306	26-28cm		100
504-B(4)	51005307	28-30cm		100
504-B(5)	51005308	30-32cm		100
504-B(6)	51005309	32-34cm		100
504-B(7)	51005310	34-36cm		100
504-B(8)	51005311	36-38cm		100
504-B(9)	51005312	38-40cm		100

### **For Neonatal**

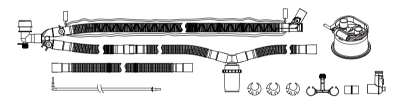


51051031

Model No.	Part No.	Tube length	Spare limb	Pressure line	Chamber VHC10	Adaptor kits	Box Qty
510-009	51005918	1.5m	0.6m				10
	51051033	1.5m	0.6m		$\sqrt{}$		10
510-009	51005928	1.5m	0.6m	$\sqrt{}$			10
	51005923	1.5m	0.6m	$\sqrt{}$	$\sqrt{}$		10
	51051031	1.5m	0.6m	$\checkmark$	$\checkmark$	$\sqrt{}$	10

### **Single Heated Breathing Circuits**

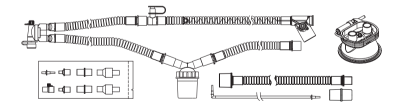
### **For Adult**



51050941

Model No.	Part No.	Tube length	Spare limb	Pressure line	Chamber VHC20	Catheter Mount	Box Qty
510-008	51048400	1.5m	0.45m				10
	51050941	1.5m	0.45m	$\sqrt{}$	$\sqrt{}$		10
	51006113	1.5m	0.45m		$\sqrt{}$	$\sqrt{}$	10

### **For Neonatal**



### 51051030

Model No.	Part No.	Tube length	Spare limb	Pressure line	Chamber VHC10	Adaptor kits	Catheter Mount	Box Qty
510-009	51005924	1.5m	0.6m	$\checkmark$				20
	51005921	1.5m	0.6m		$\sqrt{}$			10
	51005922	1.5m	0.6m	$\sqrt{}$	$\sqrt{}$			10
510-009	51005933	1.5m	0.6m	$\sqrt{}$		$\sqrt{}$		10
	51051030	1.5m	0.6m	$\sqrt{}$	$\sqrt{}$	$\sqrt{}$		10
	* 51006114	1.5m	0.6m		$\sqrt{}$	$\sqrt{}$	$\checkmark$	10
510-009	* 51051064	1.5m	0.6m			$\sqrt{}$		10

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### **Humidification Chambers**









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Model No.	VHC10	VHC20	VHC25	VHC50
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Туре	Disposable	Disposable	Reusable	Disposable
Application	Infant	Adult/Pediatric	Adult	Adult
Fill Water Type	Automatic	Automatic	Manual	Manual
Interface Connections	ISO 5356-1	ISO 5356-1	ISO 5356-1	ISO 5356-1
interface definedations	(22 mm Male)	(22mm Male)	(22mm Male)	(22mm Male)
Compressible Volume (Empty)	300 mL	440 mL	640 mL	690 mL
Compressible Volume (Full)	120 mL	260 mL	140 mL	190 mL
Maximum Water Capacity	180 mL	180 mL	500 mL	500 mL
Box Qty	10	10	10	10





## Preventing healthcare associated infections with our filter portfolio





# High mortality caused by nosocomial infections and ventilation-induced lung damage

Minimising the risk of healthcare associated infections (HCAIs) and avoiding the extra workload, stress and costs caused by HCAIs is at the heart of improving your clinical outcomes and maintaining the hospital's reputation. To help address these concerns and decrease financial burdens, as Your Specialist in Acute Care, we support you in your fight against HCAIs and assist you in improving staff and patient safety—through the entire patient pathway.

### **Nosocomial infections**

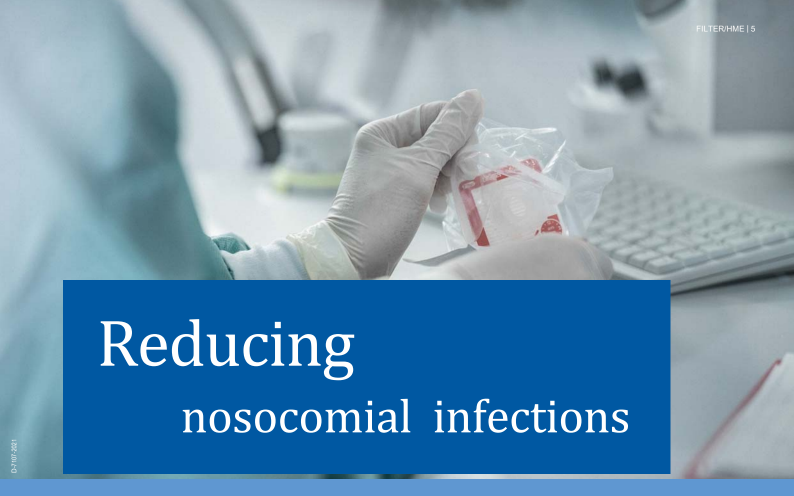
- Of every 100 hospitalised patients at any given time, 7 in developed and 10 in developing countries will acquire at least one health care-associated infection<sup>[1]</sup>
- 10 000-20 000 end fatally[2]
- -20-30% of nosocomial infections could be prevented by improved hygiene<sup>[3]</sup>
- Nosocomial infections prolong hospital stay by an average of 10 days + excess cost of \$15,275 for confirmed hospital-acquired infection [4]

### Ventilator-induced lung damages

- Ventilation-induced lung injury can contribute significantly to morbidity and mortality in critically ill patients [5]
- The lack of humidification of medically administered gases leads to ventilation-induced lung damage and increased risk of infection<sup>[6]</sup>









As a preventive measure for infection prophylaxis and avoiding the risk of crossinfection, various expert committees recommend the use of a breathing system filter

In order to avoid cross-contamination and microorganisms from entering the breathing circuit, it is advisable to place a barrier between the patient and the breathing circuit, especially when the device comes in contact with more than one patient. To protect you and your patients from getting in contact with contiguous bacteria and viruses, this barrier must be a filter which lets air pass but holds back microorganisms to the highest possible degree. Moreover, to ensure that your device is functioning at its most optimal against microorganisms, a filter is recommended on the device side whenever possible, thus protecting your staff at all times.

### Filtration Efficiency

In order to protect your patient and their surroundings, filtration efficiency is a significant parameter that ensures the avoidance of cross-contamination and infection prophylaxis. Filters have two main parameters: bacterial filtration efficiency (BFE) and viral filtration efficiency (VFE). Those two parameters are both decisive for the filtration efficiency as they indicate different things. BFE refers to how efficient the medium is in filtrating bacteria (larger in size), whereas VFE refers to how efficient the medium is in filtrating viruses (smaller in size).

### **Dead space**

When administering artificial ventilation, dead space is a vital parameter to monitor. This is because it represents the volume of ventilated air that does not participate in gas exchange. Therefore, the design of filters and HME (Heat and Moisture Exchanger) must ensure a small dead space while at the same time permit high filtration and HME performance with minimal resistance. We design our filters and HME with these requirements in mind so as to ensure a high-performance beneficial flow.



### Humidification to support lung-protective ventilation



Why is humidity important in ventilation therapies?

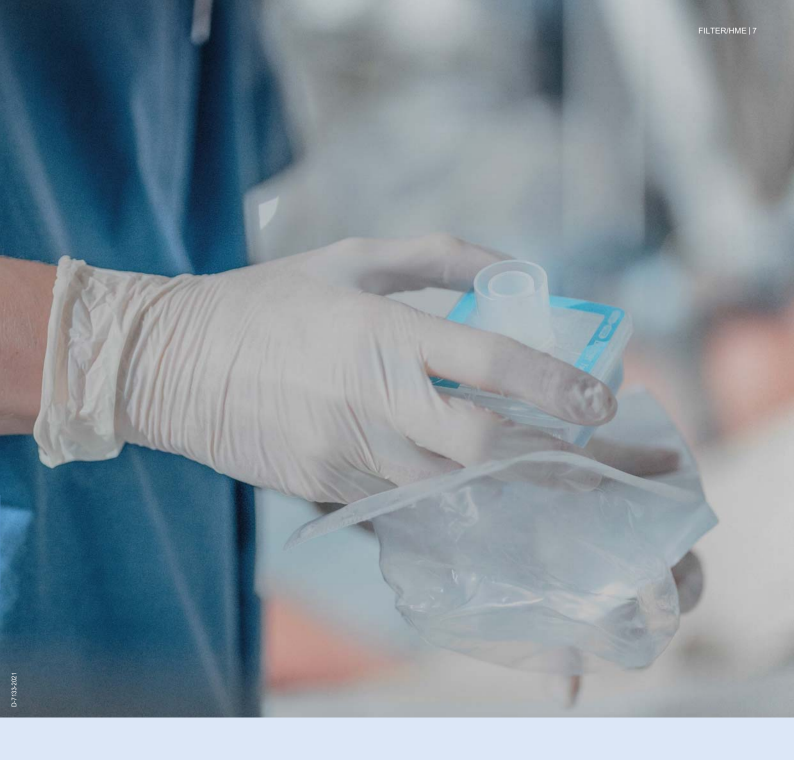
It's all about giving patients the most comfortable treatment to improve patient comfort and safety. Thus the right humidification of inspired gas in mechanical ventilation is an essential part in your daily routines. In patients receiving respiratory support therapy, the natural humidification process is often overwhelmed or even completely bypassed.



### Challenges possibly caused by dry inspired air

- Drying out of mucosa and hypothermia, resulting in viscous mucus
- Slowdown of the mucociliary transport system (contaminents aren't removed)
- 3 Higher infection risk
- /4 Impairment of surfactant activity
- Higher risk of air trapping, hyperinflation and atelectasis
- Possible degradation of gas exchange due to changes in lung
- Compliance and airway patency
  Increased airway workload

To improve outcomes in patients requiring ventilation therapy, all types of mechanical ventilation, artificial humidification, and warming of the inhaled air are recommended.



### Medical gas for ventilation has a low temperature and low humidity

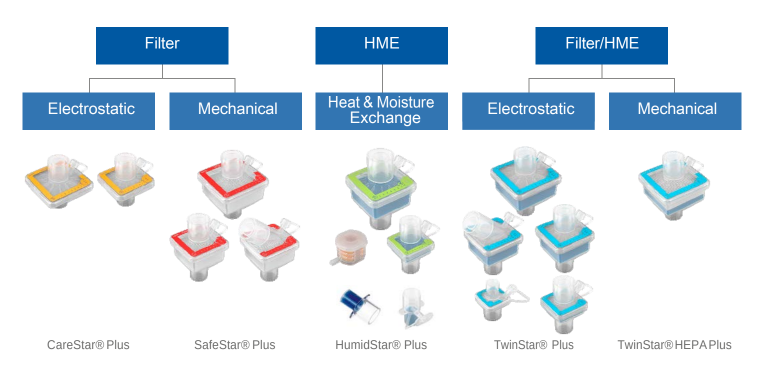
In some cases, patients receiving respiratory support therapy, the natural humidification process is often bypassed. During mechanical ventilation, the breathing gas used is much colder (~15°C) and dryer than the ambient atmosphere. When patients are intubated near the carina, this gas enters the lungs at a much lower temperature level. At this point, the gas can no longer be substantially humidified before it is distributed further into the lungs. This can have a negative effect on the pulmonary immune system: secretions tend to thicken, inhibiting their gas transport and clearance. Surfactant function is also negatively influenced, and the overall airway workload is increased. To address these concerns and improve outcomes, the sufficient humidification and warming of breathing gas can significantly help counteract these negative effects and reduce the rate of ventilator- associated lung injuries (VALI). [8]

With the utilisation of our technologies like the heat and moisture exchanger (HME), you can support your patient with a safe protective lung ventilation strategy. HMEs collect and store moisture from the expiratory phase of breathing and return a portion of both to the breathing gas during the inspiratory phase. This enables you to remarkably improve the conditions of inhaled gas (gas temperature: ~25-30°C, increased humidity) and protect the respiratory epithelium. [9]

## Our filters/HMEs for all your clinical applications and needs

### Filter/HME for all applications

Choosing the right filter application can have a significant impact on the success of patient ventilation therapies and recovery. As your Specialist in Acute Care, we offer you an extensive portfolio of high-quality breathing system solutions for all applications—supporting you with all your clinical needs.



### Manufacturing quality

- 1) Ensured quality thanks to fully automated production
- 2 Fully automated testing of every filter during the production processes
- 3 Clean room classified production (clean room class 8 acc. ISO 14644-1)
- Sustainable production thanks to optimised production and logistic processes to reduce emissions
- Production based in Lübeck, Germany



### Quality at every corner

Openianot quant

As a healthcare professional, you value quality when delivering ventilation therapy. This is why we test our products extensively for both quality and compatibility. As Your Specialist in Acute Care, we look back on more than 130 years of experience and expertise in the field of filtration. In order to protect your patients, staff and medical equipment from bacterial and viral contaminants, we know how important reliable filtration is in lung protection strategies.

### **Product quality**

Being able to rely on the quality of medical equipment is a needed prerequisite to fully concentrate on the application of therapies. We provide you with high quality filters for different fields of application.

- High bacterial -/ viral retention rates of up to >99.9999 %
- Standardised connectors provide proper and easy connection with other components of the ventilation circuit to simplify workflows
- Equipped with a 45° angled Luer-Lock connector for gas sampling to facilitate handling for clinical staff
- Transparent housing of the products allows for visual inspection at any time while in use
- Fast and easily identified due to their color coding and clear labeling
- Writable surface to easily document time of filter application and usage time to improve operating lives of system components and to ensure high filtration performance
- Lightweight product design to enhance patient comfort

- ortiono quanty

We make it possible for you to enable high quality therapy—all from a single source. Freeing you of routine tasks by streamlining and facilitating your hospital processes and workflows, reducing staff stress and unnecessary costs by offering you tailor-made solutions, at the end of the day it means giving you more time to care for your patients' wellbeing and recovery.

### Innovation quality

Our common goal is to optimise your workflows by equipping you with innovative medical equipment that facilitates clinical processes so as to save you time and simplify handling. That's why we put much effort into the innovation and design of our filters—to help you achieve these goals:

### Application safety



### Variety of applications



 Clear visibility of filter type and clear visibility of deadspace to ensure the right filter for each application

### Infection prevention



### Innovation



 Clear visibility of single use disposable product and writable surface to easily document time of filter application



### Reliable quality for every emergency



CareStar® Plus Electrostatic Filter Family



### Managing cost of high quality care with CareStar® Plus

Managing cost of care and at the same time delivering high quality therapies to improve patient outcomes are two main objectives in today's hospital environment. Together with you, we assist you in combining these objectives so that you can focus more on caring for your patients. Our CareStar Plus breathing system filters provide an excellent and cost-efficient solution. Due to its highperforming electrostatic filtration medium, CareStar Plus protects the patient from potentially present microorganisms in the inspired air as well as the ventilator breathing system from airborne microorganisms that the patient exhales. This reduces the risk of possible cross-infection and promotes patient and staff safety.

### Emergency handling

- Blister packaging for quick application
- Lightweight to improve patient comfort
- Clear color coding
- Simple secure LuerLock port for quick connection of the Sample Line



### Economically attractive

- Cost-effective filter for protection
- Very good filtration performance
   Bacterial retention: ≥ 99.99%
   Viral filtration: ≥ 99.9%





SafeStar® Plus Mechanical Filter Family



### Improve patient outcomes by preventing HCAI

Preventing patients from acquiring a healthcare associated infection (HCAI), especially when they are that their most vulnerable, is a major undertaking in your daily work. It is also an issue that carries an immense financial burden for your institution. Our new SafeStar Plus mechanical HEPA breathing system filters meet high standards for infection prophylaxis in ventilation. The active medium here is a hydrophobic filter membrane of coated glass fibres developed specifically for this purpose. Due to its hydrophobicity, potentially contaminated fluids (e.g. blood, sputum and condensate) cannot pass through SafeStar Plus filters under the normal pressure conditions of mechanical ventilation. As a result, SafeStar Plus can inhibit the passage of fluid-borne microorganisms. Furthermore, SafeStar Plus' mechanical medium has high bacterial and viral filtration efficiency rates that considerably reduce the passage of airborne microorganisms. This significantly helps to decrease the risk of possible cross-infection. We aim to support you to achieve the goals of preventing HCAI and managing cost of care—at the same time.

### Performance

- Excellent bacterial filtration: ≥99.999 %
- Excellent virus filtration rate: ≥99.999%

### Application variety

- Wide range of applications depending on application to Y-piece, inspiratory port or expiratory port
- Extensive coverage of different tidal volumes



### Safety

- Outstanding product performance
- Cleanroom classificationISO8 (acc. ISO 14644-1)
- Safe, clean blister packaging
- Writable pad printing for safe application time



Findtherightfilterforyourindivivdualneeds

Which is the right filter for your specific needs? Choosing the appropriate filter can be an overwhelming decision. That is why we want to help you have a clear understanding of the differences between these filter varieties.

### Mechanical filters = high performance

- Irregular grid of fibrers
- No defined average pore size
- Rather tightly woven
- Typically glass/ceramic fibrers, resin bound
- -Thin filter paper, pleated to yield high surface area (often named "pleated" filter)



### Electrostatic filters = good performance

- Irregular grid of fibres
- No defined average pore size
- Rather loosely woven
- Polymeric fibrers
- -One "thick" layer
- Additionally: Polarisation of fibers resulting in an electrical charge







We designed our new filter portfolio to combine quality and sustainability. This is our commitment, this is the responsibility we have for a healthier environment.



New production in Germany with European suppliers leads to a 90% reduction in annual carbon dioxide equivalent ( $CO_2e$ ) transport emissions

All our filters are latex, PVC, BPA and DEHP free

Improved product design saves >13 tonnes of glue per year

Sustainable design results in 35% weight reduction per filter housing, saving >75 tonnes of plastic per year

### Humidification to protect the respiratory system





### Humidification prevents ventilator induced lung damages

Effectively prevent lung damage induced by mechanical ventilation that involves cold and dry gases. To protect patients' respiratory system from drying out and ensuing lung damage, our HumidStar Plus HME supports to passively humidificate the air they inhale. The HME medium of our HumidStar® Plus HMEs consists of a new microporous polymer foam that was specially developed for this application as it returns a high degree of moisture. In addition, we offer the HumidStar Trach Plus for tracheostomised patients to ensure a lung protective ventilation for all patients.

### Application comfort Easy-to-use alternative to active humidification

Cost effective alternative to active humidification



### Infection prevention

- Disposable product for the reduction of infection sources
- Passive humidification for lung-protective ventilation





TwinStar®PlusCombinedFilterFamily



### Improve your patient outcomes: Filtration and humidification at the same time

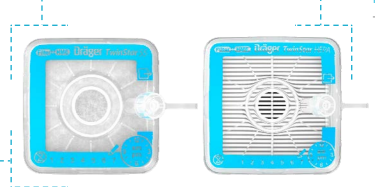
Offering patients a comfortable and quick recovery while managing cost of care are clinical goals you strive for every day. As Your Acute Care Specialist, we designed our TwinStar Plus breathing system filters/HMEs to combine all the advantages of our filter and HME portfolio—to help you save on costs and promote patient healing. They efficiently humidify and heat the inspired air of the ventilator-dependent patient. Additionally, with their high bacterial and viral filtration efficiency rates they exceptionally sustain infection prevention. Our TwinStar Plus portfolio supports the protection of your patient from potentially present microorganisms in the inspired air as well as safeguards the ventilator breathing system from airborne microorganisms that the patient exhales. To further increase patient safety, the TwinStar HEPA Plus is highlighted by a hydrophobic filter membrane of coated glass fibre.

### Optimal combination

- Economic solution through combination of HME and electrostatic/mechanical filter
- Excellent filtration and highly effective humidification performance combined

### Application variety

- Wide range of applications depending on application to Y-piece, inspiratory port or expiratory port
- Extensive coverage of different tidal volumes depending on patient group



### HEPA classification

 High-efficiency particulate filter with very good separation efficiency for increased safety

Product name		Filter SafeStar® 55 Plus	Filter SafeStar® 60A Plus	Filter SafeStar® 90 Plus	Filter/HME TwinStar® 90 Plus	Filter/HME TwinStar® HEPA Plu
Part-no.		MP05790	MP05795	MP05785	MP05800	MP05801
Patient category		Adult	Adult	Adult	Adult	Adult
Recommended tidal vol	ume	300-1500 mL	300-1500 mL	300-1500 mL	300-1500 mL	300-1500 mL
PVC & DEHP free? Latex free? Lead (Pb) free? Polyester free?		Yes Yes Yes Yes	Yes Yes Yes Yes	Yes Yes Yes Yes	Yes Yes Yes Yes	Yes Yes Yes Yes
Reusable / Disposable?	?	Disposable	Disposable	Disposable	Disposable	Disposable
Reprocessing / Cleaning	ng	No	No	No	No	No
Maximum duration of us	se (hours)	24	24	24	24	24
	Deadspace (ml)	55	60	90	90	90
	Filtration Efficiency (%) (Non-Conditioned)*	≥99.709%	≥99.906 %	≥99.904 %	≥99.00 %	≥99.891%
	Bacterial retention (%)	≥99.999 %	≥99.999 %	≥99.9999 %	≥99.99 %	≥99.9999 %
	Viral retention (%)	≥99.999 %	≥99.9999 %	≥99.999 %	≥99.9 %	≥99.9999 %
	Moisture Loss (mg H2O/Lair)				≤5.9 at VT=500 mL	≤10.9 at VT=500 m
	Moisture Output (mg H2O/L air)				≥38.1 at VT=500 mL	≥33.1 at VT=500 m
	Filtration method	Mechanical	Mechanical	Mechanical	Electrostatic	Mechanical
Performance Data	Leakage @70mbar (ml/min)	≤5	≤5	≤5	≤5	≤5
enormance Data	Compliance @60mbar Compliance @30mbar	≤1 ≤1	≤1 ≤1	≤1 ≤1	≤1 ≤1	≤1 ≤1
	Resistance 2.5 L/min	≤0.3 mbar	≤0.3 mbar	≤0.3 mbar	≤0.3 mbar	≤0.3 mbar
	Resistance 5 L/min	≤0.4 mbar	≤0.4 mbar	≤0.3 mbar	≤0.3 mbar	≤0.4 mbar
	Resistance 15 L/min	≤1.1mbar	≤1.1mbar	≤0.7 mbar	≤0.6 mbar	≤0.8 mbar
	Resistance 30 L/min	≤2 mbar	≤2 mbar	≤1.3 mbar	≤1 mbar	≤1.6 mbar
	Resistance 60 L/min	≤4.2 mbar	≤4.2 mbar	≤2.8 mbar	≤2 mbar	≤3.3 mbar
	Resistance 90 L/min	≤6.7 mbar	≤6.7 mbar	≤4.6 mbar	≤3.5 mbar	≤5.2 mbar
	Sampling port	Luer-Lock with tethered cap	Luer-Lock with tethered cap	Luer-Lock with tethered cap	Luer-Lockwith tethered cap	Luer-Lockwith tethered cap
Connections towards de	evice	22F/15M	22F/15M	22F/15M	22F/15M	22F/15M
Connections towards pa	atient	22M/15F	22M/15F	22M/15F	22M/15F	22M/15F
General comment on co	nnections		angled connector			
ength (mm)		55	55	64	64	64
Width (mm)		55	55	64	64	64
Height (mm)		80.8	91.5	76.8	76.8	76.8
Weight (g)		20.8	22.8	27.3	22.2	26.8
during operation	Temperature range	5 to 40 °C (41 to 104 °F) 5 to 95%	5 to 40 °C (41 to 104 °F) 5 to 95 %	5 to 40 °C (41 to 104 °F) 5 to 95%	5 to 40 °C (41 to 104 °F) 5 to 95%	5 to 40 °C (41 to 104 °F) 5 to 95%
	Relative humidity range	(non-condensing) 570 to 1200 hPa	(non-condensing) 570 to 1200 hPa	(non-condensing) 570 to 1200 hPa	(non-condensing) 570 to 1200 hPa	(non-condensing 570 to 1200 hPa
	Air pressure range	(8.3 to 17.4 psi)	(8.3 to 17.4 psi)	(8.3 to 17.4 psi)	(8.3 to 17.4 psi)	(8.3 to 17.4 psi)
luring storage	Temperature range	–20 to 60 °C (–4 to 140 °F)	–20 to 60 °C (–4 to 140 °F)	−20 to 60 °C (−4 to 140 °F)	–20 to 60 °C (–4 to 140 °F)	–20 to 60 °C (–4 to 140 °F)
	Relative humidity range	5 to 95% (non-condensing)	5 to 95 % (non-condensing)	5 to 95 % (non-condensing)	5 to 95% (non-condensing)	5 to 95% (non-condensing
	Air pressure range	570 to 1200 hPa (8.3 to 17.4 psi)	570 to 1200 hPa (8.3 to 17.4 psi)	570 to 1200 hPa (8.3 to 17.4 psi)	570 to 1200 hPa (8.3 to 17.4 psi)	570 to 1200 hPa (8.3 to 17.4 psi)
during transport	Temperature range	-20 to 60 °C (-4 to 140 °F)	-20 to 60 °C (-4 to 140 °F)	-20 to 60 °C (-4 to 140 °F)	-20 to 60 °C (-4 to 140 °F)	-20 to 60 °C (-4 to 140 °F)
	Relative humidity range	5 to 95% (non-condensing) 570 to 1200 hPa	5 to 95 % (non-condensing) 570 to 1200 hPa	5 to 95 % (non-condensing) 570 to 1200 hPa	5 to 95% (non-condensing) 570 to 1200 hPa	5 to 95% (non-condensing 570 to 1200 hPa
e the nackaging materi	Air pressure range	(8.3 to 17.4 psi)	(8.3 to 17.4 psi)	(8.3 to 17.4 psi)	(8.3 to 17.4 psi)	(8.3 to 17.4 psi)
s the packaging materia s the packaging materia		Yes Yes	Yes Yes	Yes Yes	Yes Yes	Yes Yes
Sterile? Non-Sterile? Hygienic production and	d packaging conditions	non-sterile; assembled in clean environment**	non-sterile; assembled in clean environment**	non-sterile; assembled in clean environment**	non-sterile; assembled in clean environment**	non-sterile; assemble clean environment
Packing unit		100	100	100	100	100
Country of origin		Germany	Germany	Germany	Germany	Germany
Overall Shelf Life of the	product(inyears)	5	5	5	3	5

 $<sup>\</sup>hbox{\it *filters} tested in unused state ~~ \\ \hbox{\it |}^{\star\star} product is manufactured in clean room class ISO 8 acc. EN 14644-1:2015 \\$ 

Product name		Filter/HME TwinStar® 55 Plus	Filter/HME TwinStar® 60A Plus	Filter/HME TwinStar® 25 Plus	Filter/HME TwinStar® 9 Plus	Filter CareStar® 20 Plus
					*	
Part-no.		MP05805	MP05810	MP05815	MP05820	MP05770
Patient category		Adult	Adult	Pediatric	Pediatric/Neonatal	Adult/Pediatric
	Recommended tidal volume		300-1500 mL	100-500 mL	30-150 mL	100-500 mL
PVC & DEHP free?		Yes	Yes	Yes	Yes	Yes
Latex free? Lead (Pb) free? Polyester free?		Yes Yes Yes	Yes Yes Yes	Yes Yes Yes	Yes Yes Yes	Yes Yes Yes
Reusable / Disposable	?	Disposable	Disposable	Disposable	Disposable	Disposable
Reprocessing / Cleaning	ng	No	No	No	No	No
Maximum duration of us	se (hours)	24	24	24	24	24
	Deadspace (ml)	55	60	25	9	20
	Filtration Efficiency (%) (Non-Conditioned)*	≥98.46 %	≥98.80 %	≥98.74%	≥97.07 %	≥99.551 %
	Bacterial retention (%)	≥99.99 %	≥99.99 %	≥99.98 %	≥99.99 %	≥99.99 %
	Viral retention (%)	≥99.9 %	≥99.9 %	≥99.9 %	≥99.9 %	≥99.9%
	Moisture Loss (mg H2O/L air)	≤9.4 at VT=500 mL	≤6.3 at VT=500 mL	≤11.8atVT=250mL	≤10.3 at VT=50 mL	
	Moisture Output (mg H2O/L air)	≥34.6 at VT=500 mL	≥37.7 at VT=500 mL	≥32.2 at VT=250 mL	≥33.7 at VT=50 mL	
	Filtration method	Electrostatic	Electrostatic	Electrostatic	Electrostatic	Electrostatic
Performance Data	Leakage @70mbar (ml/min)	≤5	≤5	≤5	≤5	≤5
Chomanoc Bata	Compliance @60mbar Compliance @30mbar	≤1 ≤1	≤1 ≤1	≤1 ≤1	≤1 ≤1	≤1 ≤1
	Resistance 2.5 L/min	≤0.3 mbar	≤0.3 mbar	≤0.3 mbar	≤0.3 mbar	≤0.3 mbar
	Resistance 5 L/min	≤0.3 mbar	≤0.3 mbar	≤0.4 mbar	≤0.6 mbar	≤0.3 mbar
	Resistance 15 L/min	≤0.7 mbar	≤0.7 mbar	≤1.1 mbar	≤1.5 mbar	≤0.7 mbar
	Resistance 30 L/min	≤1.3 mbar	≤1.3 mbar	≤1.8 mbar	≤3.3 mbar	≤1.3 mbar
	Resistance 60 L/min	≤3 mbar	≤3 mbar	≤3.8 mbar	≤7.2mbar	≤2.8 mbar
	Resistance 90 L/min	≤4.9 mbar	≤4.9 mbar	≤6.2 mbar	≤12.3 mbar	≤4.8 mbar
	Sampling port	Luer-Lockwith tethered cap	Luer-Lockwith tethered cap	Luer-Lock with tethered cap	Luer-Lock with tethered cap	Luer-Lockwith tethered cap
Connections towards de	evice	22F/15M	22F/15M	22F/15M	22F/15M	22F/15M
Connections towards pa	atient	22M/15F	22M/15F	22M/15F	22M/15F	22M/15F
General comment on co	onnections		angled connector			
Length (mm)		55	55	44	34	55
Width (mm)		55	55	44	34	55
Height (mm)		62	87.5	76.8	43.8	62
Weight (g)		17.6	19.3	12.4	7.1	14
during operation	Temperature range	5 to 40 °C (41 to 104 °F)	5 to 40 °C (41 to 104 °F)	5 to 40 °C (41 to 104 °F)	5 to 40 °C (41 to 104 °F)"	5 to 40 °C (41 to 104 °F)
	Relative humidity range	5 to 95 % (non-condensing)	5 to 95% (non-condensing)			
Ai	Air pressure range	570 to 1200 hPa (8.3 to 17.4 psi)	570 to 1200 hPa (8.3 to 17.4 psi)			
during storage	Temperature range	–20 to 60 °C (–4 to 140 °F)	–20 to 60 °C (–4 to 140 °F)			
	Relative humidity range	5 to 95 % (non-condensing)	5 to 95% (non-condensing)	5 to 95 % (non-condensing)	5 to 95 % (non-condensing)	5 to 95% (non-condensing)
	Air pressure range	570 to 1200 hPa (8.3 to 17.4 psi)	570 to 1200 hPa (8.3 to 17.4 psi)			
during transport	Temperature range	–20 to 60 °C (–4 to 140 °F)	–20 to 60 °C (–4 to 140 °F)	-20 to 60 °C (-4 to 140 °F)	-20 to 60 °C (-4 to 140 °F)	–20 to 60 °C (–4 to 140 °F)
	Relative humidity range	5 to 95 % (non-condensing)	5 to 95% (non-condensing)	5 to 95 % (non-condensing)	5 to 95 % (non-condensing)	5 to 95% (non-condensing)
	Air pressure range	570 to 1200 hPa (8.3 to 17.4 psi)	570 to 1200 hPa (8.3 to 17.4 psi)			
Is the packaging materials the packaging materials		Yes Yes	Yes Yes	Yes Yes	Yes Yes	Yes Yes
Sterile? Non-Sterile? Hygienic production and	d packaging conditions	non-sterile; assembled in clean environment**	non-sterile; assemble clean environment*			
Packingunit	·	100	100	100	100	100
Country of origin		Germany	Germany	Germany	Germany	Germany
				l .	1	l .

Product name		Filter CareStar®35Plus	HME HumidStar®55Plus	HME HumidStar®25Plus	HME HumidStar® 2 Plus	HME HumidStar® 2 Plus Luer-Lock
					*	
Part-no.		MP05755	MP05730	MP05735	MP05845	MP05840
Patient category		Adult	Adult	Pediatric	Neonatal	Neonatal
Recommended tidal volu	ume	300-1500 mL	300-1500 mL	100-500 mL	10-50 mL	10-50 mL
PVC & DEHP free? Latex free? Lead (Pb) free? Polyester free?		Yes Yes Yes Yes	Yes Yes Yes Yes	Yes Yes Yes Yes	Yes Yes Yes Yes	Yes Yes Yes Yes
Reusable / Disposable?	?	Disposable	Disposable	Disposable	Disposable	Disposable
Reprocessing / Cleanin	g	No	No	No	No	No
Maximum duration of us	e (hours)	24	24	24	24	24
	Deadspace (ml) Filtration Efficiency (%)	35 ≥99.217 %	55 	25		2
	(Non-Conditioned)*  Bacterial retention (%)	≥99.99 %				
	Viral retention (%)	≥99.9 %				
	Moisture Loss (mg H2O/Lair)		≤7.8 at VT=500 mL	≤9.3 at VT=250 mL	≤11.5 at VT = 45 mL	≤11.5 at VT = 45 m
	Moisture Output (mg H2O/L air)		≥36.2 at VT=500 mL	≥34.7 at VT=250 mL	≥32.5atVT=45mL	≥32.5atVT=45m
	Filtration method	Electrostatic	none	none	none	none
	Leakage @70mbar (ml/min)	≤5	≤5	≤5	≤1	≤2
erformance Data	Compliance @60mbar Compliance @30mbar	≤1 ≤1	≤1 ≤1	≤1 ≤1	≤1 ≤1	≤1 ≤1
	Resistance 2.5 L/min	≤0.3 mbar	≤0.3 mbar	≤0.3 mbar	≤0.3 mbar	≤0.3 mbar
	Resistance 5 L/min	≤0.3 mbar	≤0.3 mbar	≤0.3 mbar	≤1 mbar	≤1 mbar
	Resistance 15 L/min	≤0.6 mbar	≤0.3 mbar	≤0.3 mbar	≤1.2 mbar	≤3.5 mbar
	Resistance 30 L/min	≤0.9 mbar	≤0.6 mbar	≤0.3 mbar	≤3.2 mbar	≤3.5 mbar
	Resistance 60 L/min	≤2 mbar	≤1 mbar	≤0.9 mbar	≤11.5 mbar	≤12 mbar
	Resistance 90 L/min	≤3.5 mbar	≤2 mbar	≤1.5 mbar	≤25 mbar	≤27 mbar
	Sampling port	Luer-Lockwith tethered cap	Luer-Lock with tethered cap	Luer-Lock with tethered cap		Luer-Lock with tethered cap
Connections towards de	evice	22F/15M	22F/15M	22F/15M	15M	15M
Connections towards pa	atient	22M/15F	22M/15F	22M/15F	15F	15F
General comment on co	nnections					
ength (mm)		64	55	44		
Vidth (mm)		64	55	44		
leight (mm)		62	80.8	76.8		
Veight (g)		16.8	17	12.2	2.8	3.2
luring operation	Temperature range	5 to 40 °C (41 to 104 °F) 5 to 95%	5 to 40 °C (41 to 104 °F) 5 to 95%	5 to 40 °C (41 to 104 °F) 5 to 95%	5 to 40 °C (41 to 104 °F) 5 to 95%	5 to 40 °C (41 to 104 °F) 5 to 95 %
	Relative humidity range  Air pressure range	(non-condensing) 570 to 1200 hPa	(non-condensing) 570 to 1200 hPa	(non-condensing) 570 to 1200 hPa	(non-condensing) 570 to 1200 hPa	(non-condensing 570 to 1200 hPa
	All pressure range	(8.3 to 7.4 psi) -20 to 60 °C	(8.3 to 7.4 psi) -20 to 60 °C	(8.3 to 7.4 psi) -20 to 60 °C	(8.3 to 7.4 psi) -20 to 60 °C	(8.3 to 7.4 psi) -20 to 60 °C
luring storage	Temperature range	(-4 to 140 °F) 5 to 95%	(-4 to 140 °F) 5 to 95%	(-4 to 140 °F) 5 to 95 %	(–4 to 140 °F) 5 to 95%	(-4 to 140 °F) 5 to 95%
	Relative humidity range	(non-condensing) 570 to 1200 hPa	(non-condensing) 570 to 1200 hPa	(non-condensing) 570 to 1200 hPa	(non-condensing) 570 to 1200 hPa	(non-condensing
luda a ka	Air pressure range	(8.3 to 17.4 psi) -20 to 60 °C	(8.3 to 17.4 psi) -20 to 60 °C	(8.3 to 17.4 psi) -20 to 60 °C	(8.3 to 17.4 psi) -20 to 60 °C	(8.3 to 17.4 psi) -20 to 60 °C
luring transport	Temperature range	-20 to 60 °C (-4 to 140 °F) 5 to 95%	-20 to 60 °C (-4 to 140 °F) 5 to 95%	-20 to 60 °C (-4 to 140 °F) 5 to 95 %	-20 to 60 °C (-4 to 140 °F) 5 to 95%	-20 to 60°C (-4 to 140°F) 5 to 95%
	Relative humidity range	(non-condensing) 570 to 1200 hPa	(non-condensing) 570 to 1200 hPa	(non-condensing) 570 to 1200 hPa	(non-condensing) 570 to 1200 hPa	(non-condensing 570 to 1200 hPa
s the packaging materia	Air pressure range	(8.3 to 17.4 psi)	(8.3 to 17.4 psi) Yes	(8.3 to 17.4 psi) Yes	(8.3 to 17.4 psi) Yes	(8.3 to 17.4 psi) Yes
s the packaging materia Sterile? Non-Sterile?		Yes	Yes	Yes	Yes	Yes
	d packaging conditions	non-sterile; assembled in clean environment**	non-sterile; assembled in clean environment**	non-sterile; assembled in clean environment**		non-sterile; assemble clean environment
			4.0.0		4.0.0	
Packing unit Country of origin		100 Germany	100 Germany	100 Germany	100 Sweden	100 Sweden

 $<sup>\</sup>label{thm:complex} {}^*\text{filters} \, \text{tested} \, \text{in unused} \, \text{state} \, \\ I \, {}^{**}\text{product} \, \text{is} \, \text{manufactured} \, \text{in clean room class} \, \\ ISO \, 8 \, \text{acc.} \, EN \, 14644-1:2015 \, \\ \text{For more details} \, \text{see} \, \\ IFU \, \text{of the products} \, - \, \text{Not all articles} \, \text{are available worldwide} \, \\ \text{The product} \, - \, \text{Not all articles} \, \text{are available} \, \text{worldwide} \, \\ \text{The product} \, - \, \text{Not all articles} \, \text{are available} \, \text{worldwide} \, \\ \text{The product} \, - \, \text{Not all articles} \, \text{are available} \, \text{worldwide} \, \\ \text{The product} \, - \, \text{Not all articles} \, \text{are available} \, \text{worldwide} \, \\ \text{The product} \, - \, \text{Not all articles} \, \text{are available} \, \text{worldwide} \, \\ \text{The product} \, - \, \text{Not all articles} \, \text{The product} \, - \, \text{Not all articles} \, \text{The product} \, \\ \text{The product} \, - \, \text{Not all articles} \, \text{The product} \, - \, \text{Not all articles} \, \\ \text{The product} \, - \, \text{The pro$ 

	Product name		HME HumidStar® Trach Plus	CombiStar Filter HME	CombiStar Filter HME flex	CombiStar F-HME HEPA flex	CombiStar mechanical Filterflex
ormation in General				7			<b>\(\frac{1}{2}\)</b>
format	Part-no.		MP05750	MP12060	MP12061	MP12062	MP12063
luct In	Patient category		Adult/Pediatric	Adult	Adult	Adult	Adult
Prod	Recommended tidal volum	ne	100 - 1500 mL				
	PVC & DEHP free? Latex free?		Yes Yes	Yes Yes	Yes Yes	Yes Yes	Yes Yes
	Lead (Pb) free? Polyester free?		Yes Yes	Yes Yes	Yes Yes	Yes Yes	Yes Yes
	Reusable / Disposable?		Disposable	Disposable	Disposable	Disposable	Disposable
Use	Reprocessing / Cleaning		No	No	No	No	No
	Maximum duration of use	(hours)	24	24	24	24	24
		Deadspace (ml)	6	Check individual components	Check individual components	Check individual components	Check individual components
		Filtration Efficiency (%) (Non-Conditioned)*		≥98.46%	≥98.46%	≥99.891%	≥99.709%
		Bacterial retention (%)		≥99.99 %	≥99.99%	≥99.9999 %	≥99.999 %
		Viral retention (%)		≥99.9 %	≥99.9 %	≥99.9999 %	≥99.999 %
		Moisture Loss (mg H2O/L air)	≤10.8at Vt=250 mL ≤14.4at Vt=500 mL	≤9.4 at VT=500 mL	≤9.4 at VT=250 mL	≤10.9 at VT=50 mL	
natior		Moisture Output (mg H2O/L air)	≥29.6 at VT=500 mL	≥34.6 at VT=500 mL	≥34.6 at VT=250 mL	≥33.1 at VT=50 mL	
nforn		Filtration method	none	Electrostatic	Electrostatic	Mechanical	Mechanical
cifici	Performance Data	Leakage @70mbar (ml/min)	n/a	-			
Product specific information		Compliance @60mbar Compliance @30mbar	n/a n/a				
roduc		Resistance 2.5 L/min	n/a				
Ф		Resistance 5 L/min	n/a	Check individual components	Check individual components	Checkindividual components	Check individual components
		Resistance 15 L/min	n/a	Components	Components	components	components
		Resistance 30 L/min	≤0.1				
		Resistance 60 L/min	≤0.3	=			
		Resistance 90 L/min	≤0.6				
		Sampling port		Luer-Lockwith tethered cap	Luer-Lock with tethered cap	Luer-Lock with tethered cap	Luer-Lockwith tethered cap
	Connections towards devi	ice		22F/15M	22F/15M	22F/15M	22F/15M
	Connections towards patie	ent	15F	22M/15F	22M/15F	22M/15F	22M/15F
S	General comment on conr	nections					
	Length (mm)						
uctS	Width (mm)						
Prod	Height (mm) Weight (g)		6				
	during operation	<u> </u>	5 to 40 °C	 10to 40 °C	10 to 40 °C	 10 to 40 °C	 10to40°C
	daming operation	Temperature range	(41 to 104 °F)	(50 to 104 °F)	(50 to 104 °F)	(50 to 104 °F)	(50 to 104 °F)
		Relative humidity range	5 to 95% (non-condensing)	5 to 95 % (non-condensing)			
			570 to 1200 hPa	570 to 1200 hPa	570 to 1200 hPa	570 to 1200 hPa	570 to 1200 hPa
sus		Air pressure range	(8.3 to 7.4 psi) -20 to 60 °C	(8.3 to 17.4 psi) -20 to 60 °C	(8.3 to 17.4 psi) -20 to 60 °C	(8.3 to 17.4 psi)	(8.3 to 17.4 psi)
conditio	during storage	Temperature range	–20 to 60 °C (–4 to 140 °F)	-20 to 60 °C (-4 to 140 °F)	-20 to 60 °C (-4 to 140 °F)	–20 to 60 °C (–4 to 140 °F)	–20 to 60 °C (–4 to 140 °F)
		Relative humidity range	5 to 95%	5 to 95%	5 to 95 %	5 to 95 %	5 to 95%
ment			(non-condensing) 570 to 1200 hPa	(non-condensing) 570 to 1200 hPa	(non-condensing) 570 to 1200 hPa	(non-condensing) 570 to 1200 hPa	(non-condensing) 570 to 1200 hPa
Environmental		Air pressure range	(8.3 to 17.4 psi)	(8.3 to 17.4 psi)	(8.3 to 17.4 psi)	(8.3 to 17.4 psi)	(8.3 to 17.4 psi)
Ш	during transport	Temperature range	–20 to 60 °C (–4 to 140 °F)	–20 to 60 °C (–4 to 140 °F)	–20 to 60 °C (–4 to 140 °F)	−20 to 60 °C (−4 to 140 °F)	–20 to 60 °C (–4 to 140 °F)
		Relative humidity range	5 to 95%	5 to 95%	5 to 95 %	5 to 95 %	5 to 95%
			(non-condensing)	(non-condensing)	(non-condensing)	(non-condensing)	(non-condensing)
		Air pressure range	570 to 1200 hPa (8.3 to 17.4 psi)	570 to 1200 hPa (8.3 to 17.4 psi)	570 to 1200 hPa (8.3 to 17.4 psi)	570 to 1200 hPa (8.3 to 17.4 psi)	570 to 1200 hPa (8.3 to 17.4 psi)
S	Is the packaging material I Is the packaging material I		Yes Yes	Yes Yes	Yes Yes	Yes Yes	Yes Yes
gistic	Sterile? Non-Sterile? Hygienic production and p	nackaging conditions		non-sterile; assembled in clean environment**			
og/Lo	Packing unit	rackaging continues	100	25	25	25	25
kagir	Country of origin		Sweden	Checkindividual	Checkindividual	Checkindividual	Checkindividual
Pac				components	components	components	components
	Overall Shelf Life of the pr	roduct (inyears)	5	2	2	2	2

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### Notes

### Notes

### Notes

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CORPORATE HEADQUARTERS Drägerwerk AG & Co. KGaA Moislinger Allee 53-55 23558 Lübeck, Germany

www.draeger.com

### REGION EUROPE

Drägerwerk AG & Co. KGaA Moislinger Allee 53-55 23558 Lübeck, Germany

- **4** +49 451 882 0
- **+49 451 882 2080**
- info@draeger.com

REGION MIDDLE EAST, AFRICA Drägerwerk AG & Co. KGaA Branch Office P.O.Box 505108

Dubai, United Arab Emirates \$\sim +971 4 4294 600

- **♣** +971 4 4294 699

### REGION ASIA PACIFIC

Draeger Singapore Pte. Ltd. The Galen #04-01 Singapore 117525

- **4** +65 6872 9288
- **♣** +65 6259 0398

### REGION NORTH AMERICA

Draeger, Inc. 3135 Quarry Road Telford, PA 18969, USA **♦** +18004DRAGER

- **♣** +12157235935
- info.usa@draeger.com

### REGION CENTRAL AND SOUTH AMERICA

Dräger Indústria e Comércio Ltda. Al. Pucurui - 51 - Tamboré 06460-100 - Barueri - São Paulo

- ✓ relacionamento@draeger.com

**Locate your Regional Sales** Representative at: www.draeger.com/contact

