

Specificație tehnică completată

Model: Mașină de anestezie Carestation 650, cu monitor de pacient B155M

Producător: GE Medical Systems, Datex-Ohmeda Inc, Țara: USA

Specificarea tehnică deplină solicitată de către autoritatea contractantă	Specificarea tehnică deplină completată de către autoritatea ofertantă
<p>Mașină de anestezie (caracteristici avansate) Descriere Mașina de anestezie este destinată să livreze, să monitorizeze gazele anestezice și să asigure respirația artificială a pacientului în timpul actului chirurgical Parametru Specificația Prize de gaz O₂, Aer Display mașina de anestezie ≥15", color TFT sau LCD touch screen Debitmetre tipul electronice gaz O₂, Air gama, L/min ≥ 0 - 15</p> <p>Vaporizator tip vaporizator acceptate Izofluran da Sevofluran da Halothan da Enfluran da număr de vaporizatoare instalate la dipozitiv ≥ 2 unități da Izofluran da Sevofluran da interlock da sistem de absorbție da Mecanisme de siguranță siguranță O₂ acustică, vizuală siguranță de amestec hipoxic da Ventilator automat tip pacient Adult, Pediatric moduri de ventilație Manual/spontan, IMV, VCV, PCV, PSV, SIMV, PS mecanism electronic de amestec a gazelor (mixer) da volumul Tidal, ml 5-1500 frecvența respirației/minut 5 - 100 fluxul inspirator, L/min ≥ 3-40 raportul I:E minim 4:1 la 1:4 pauză de inspirație da limita de presiune, cmH₂O ajustabilă, ≥ 10-70 PEEP, cmH₂O ≥ 0-20</p>	<p>Mașină de anestezie (caracteristici avansate) Descriere Mașina de anestezie este destinată să livreze, să monitorizeze gazele anestezice și să asigure respirația artificială a pacientului în timpul actului chirurgical DA Parametru Specificația DA Prize de gaz O₂, Aer DA pag. 6 din Specificatia CS 650 Display mașina de anestezie ≥15", color TFT sau LCD touch screen DA 15 " pag. 4 din Specificatia CS 650 Debitmetre tipul electronice DA este present si flometru mechanic classic si cel electronic pe display. gaz O₂, Air DA gama, L/min ≥ 0 – 15 DA Maximul este de 120 L/min pag 3 din din Specificatia CS 650 Vaporizator tip vaporizator acceptate Izofluran DA Tec 820/850 Sevofluran DA Tec 820/850 Halothan DA Tec 7/7 plus Enfluran DA Tec 7 /7Plus Seria TEC 6 Plus, Tec 7 Plus Tec 820 si Tec 850 număr de vaporizatoare instalate la dipozitiv ≥ 2 unități DA Izofluran DA TEC 820 Sevofluran DA TEC 820 interlock DA pag. 4 din Specificatie CS 650 sistem de absorbție DA Suction pag. 16 (8-18) User Reference Manual Mecanisme de siguranță siguranță O₂ acustică, vizuală siguranță de amestec hipoxic DA Ventilator automat tip pacient Adult, Pediatric, neonatal DA pag. 1 Specificatie CS650 moduri de ventilație Manual/spontan, IMV, VCV, PCV, PSV, SIMV, PS DA mecanism electronic de amestec a gazelor (mixer) DA cu prezența pe ecran volumul Tidal, ml 5-1500 DA 1- 1500 ml pag. 3 Specificatie CS650 frecvența respirației/minut 4 - 100 DA pag. 3 Specificatie CS650 fluxul inspirator, L/min ≥ 3-40 DA raportul I:E minim 4:1 la 1:4 DA 2:1 la 1:8 pauză de inspirație da DA pag. 3 Specificatie CS650 limita de presiune, cmH₂O ajustabilă, 0-100 DA pag. 3 Specificatie CS650 PEEP, cmH₂O ≥ 0-30 DA pag. 3 Specificatie CS650</p>

<p>Sistem de autodiagnostic testare la scurgeri, testarea circuitelor respiratorii, compliantă, alimentarea cu gaz, verificarea tuturor sistemelor</p> <p>AGSS (sistem de evacuare a gazelor anestezice) da</p> <p>Sistem de management al consumului de agent anestezic da</p> <p>Poibilitate de a schimba tipul gazului principal din meniu da</p> <p>Circuitul pneumatic de ventilare a pacinetului cu funcție de incălzire a amestecului gazos da</p> <p>port auxiliar ieșire a amestecului gazos da</p> <p>Parametri monitorizați și afișați pe display Presiunea de aer Alarmă de înaltă presiune da</p> <p>Alarma presiune subatmosferică da</p> <p>Continuarea alarma presiune da</p> <p>Presiune scăzută / apnee da</p> <p>Alte alarme de presiune da</p> <p>Volumul expirator / flux da</p> <p>Volumul minut, l/min da</p> <p>Concentrația de O2 Alarmă apnea da</p> <p>Timp de răspuns, sec <30</p> <p>Concentrația de CO2 alarmă apnee da</p> <p>Monitorizare agent Tipul de agenți Halothan, isofluran, sevofluran, Enfluran</p> <p>Auto indentificarea gazelor anestezice da</p> <p>Alarmă concentrare agent da</p> <p>Determinarea și afișarea valorii MAC da</p> <p>spirometria da</p> <p>Modulul de gaze încorporat la mașina de anestezie da</p> <p>determină concentrațiile de gaze: O2, CO2, agenți anestezici da</p> <p>Celulă determinare O2 tip paramagnetic da</p> <p>Monitorul pentru afișarea funcțiilor vitale display ≥15", color TFT sau LCD da</p> <p>touch screen da</p> <p>monitor dedicat vizualizării funcțiilor vitale da</p> <p>braț de fixare a monitorului din laterală pe mașină de anestezie da</p> <p>imprimantă termică încorporată da</p> <p>baterie internă reîncărcabilă da</p> <p>interfață de comunicare cu altele da</p> <p>Modulele hemodinamice incluse Electro-cardio-grama (ECG) frecvența cardiacă da</p> <p>traseul ECG da</p> <p>analiza și măsurarea segmentui ST da</p> <p>determinarea cel puțin 20 de aritmii da</p> <p>Puls-oximetria (SpO2) fotopletismografia da</p> <p>valoarea SpO2 da</p>	<p>Sistem de autodiagnostic testare la scurgeri, testarea circuitelor respiratorii, compliantă, alimentarea cu gaz, verificarea tuturor sistemelor DA pag. 62 (3-4) User Reference Manual</p> <p>AGSS (sistem de evacuare a gazelor anestezice) DA pag. 7 din Specificatie CS650</p> <p>Sistem de management al consumului de agent anestezic DA ecoFLOW pag. 1 din Specificatie CS650,</p> <p>Poibilitate de a schimba tipul gazului principal din meniu DA doar in cazul incare se activeaza si al treile gaz N2O</p> <p>Circuitul pneumatic de ventilare a pacinetului cu funcție de incălzire a amestecului gazos DA</p> <p>port auxiliar ieșire a amestecului gazos DA</p> <p>Parametri monitorizați și afișați pe display Presiunea de aer Alarmă de înaltă presiune DA</p> <p>Alarma presiune subatmosferică DA</p> <p>Continuarea alarma presiune DA</p> <p>Presiune scăzută / apnee DA</p> <p>Alte alarme de presiune DA</p> <p>Volumul expirator / flux DA</p> <p>Volumul minut, l/min DA</p> <p>Concentrația de O2 Alarmă apnea DA</p> <p>Timp de răspuns, sec <30 DA</p> <p>Concentrația de CO2 alarmă apnee DA</p> <p>Monitorizare agent Tipul de agenți Halothan, isofluran, sevofluran, Enfluran, Desfluran DA</p> <p>Auto indentificarea gazelor anestezice DA</p> <p>Alarmă concentrare agent DA</p> <p>Determinarea și afișarea valorii MAC DA</p> <p>spirometria DA</p> <p>Modulul de gaze încorporat la mașina de anestezie DA</p> <p>determină concentrațiile de gaze: O2, CO2, agenți anestezici DA</p> <p>Celulă determinare O2 tip paramagnetic DA</p> <p>Monitorul pentru afișarea funcțiilor vitale display - 15.6", color TFT sau LCD DA pag.2 B155 specificatie B105M/125M/155M</p> <p>touch screen DA pag.2 B155 specificatie B105M/125M/155M</p> <p>monitor dedicat vizualizării funcțiilor vitale DA</p> <p>braț de fixare a monitorului din laterală pe mașină de anestezie DA</p> <p>imprimantă termică încorporată DA</p> <p>baterie internă reîncărcabilă DA</p> <p>interfață de comunicare cu altele DA</p> <p>Modulele hemodinamice incluse Electro-cardio-grama (ECG) frecvența cardiacă DA</p> <p>traseul ECG DA</p> <p>analiza și măsurarea segmentui ST DA</p> <p>determinarea cel puțin 20 de aritmii DA</p> <p>Puls-oximetria (SpO2) fotopletismografia DA</p> <p>valoarea SpO2 DA</p>
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Anexa 1

<p>indicile de perfuzie da Tensiune sanguină neinvazivă (NIBP) da Respirația (impedanța transtoracică) da Temperatura pe 2 canale da Tensiune sanguină invazivă (IBP) pe 2 canale da Modul de monitorizare BIS (bispectral index) sau modul de monitorizarea obiectiva a profunzimii blocului neuro-muscular intraanestezic (TOF/ NMT) da</p> <p>Alarme prioritare 3 Tensiune de alimentare 220 V, 50 Hz Prize auxiliare 220 v ≥ 3 buc da Baterie internă reîncărcabilă da autonomie de lucru ≥ 1.5h da Sertar pentru depozitare ≥ 3 buc da Frînă centralizată pentru fixarea aparatului da Presiune de alimentare cu gaze 3.0 - 6 bar Compresor Portabil pe rotile sau încorporat în mașina de anestezie da</p> <p>Punct de rouă sub presiune la max. și debit de ieșire la 3 bar, 5°C sub temperatura mediului ambiant Fluxul de ieșire la 3 bar ≥ 60 l/min. Filtrarea aerului 5 µm Mod de operare Continuu da Surgerea apei condensate Auto da Presiunea de ieșire 3.5 bar Volumul reciverului 5 l Accesori:</p> <p>Furtunul cu conector tip DIN, de conectare la sursa de aer comprimat 1 buc. Furtunul cu conector tip DIN de conectare la sursa de oxigen 1 buc. Circuit de ventilare Adult, reutilizabil ≥ 2 set. Plămîn de test Adult, reutilizabil ≥ 2 buc. Senzor de flux Reutilizabil ≥ 2 buc. Filtru antibacterial Adult, unică utilizare ≥ 100 buc. Accesori modul de gaz Adult ≥ 2 set. Cablu ECG Adult, reutilizabil 5 fire ≥ 2 buc. Senzor ECG Adult, unica utilizare ≥ 50 buc. Senzor SpO2 Adult, reutilizabil ≥ 2 Buc. Manșete NIBP Adult, reutilizabilă ≥ 2 buc. Adult mare, reutilizabilă ≥ 2 buc. Senzor de temperatură Adult, reutilizabil ≥ 1 buc. Canula pentru modulul de CO2 și capcană de condens (dacă este cazul) ≥ 12 buc. Accesori necesare de funcționare a modulului BIS sau TOF/ NMT Accesori pentru Adult ≥ 5 buc. Altele: Garanție ≥ 2 ani da Manuale de utilizare în limba de stat (Română), în cazul în care nu este prezent să fie prezentat manualul în limba de stat cu stampila biroului de traducere. da</p>	<p>indicile de perfuzie DA Tensiune sanguină neinvazivă (NIBP) DA Respirația (impedanța transtoracică) DA Temperatura pe 2 canale DA Tensiune sanguină invazivă (IBP) pe 2 canale DA Modul de monitorizare BIS (bispectral index) sau modul de monitorizarea obiectiva a profunzimii blocului neuro-muscular intraanestezic (TOF/ NMT) DA</p> <p>Alarme prioritare 3 DA Tensiune de alimentare 220 V, 50 Hz DA Prize auxiliare 220 v ≥ 3 buc DA Baterie internă reîncărcabilă DA autonomie de lucru ≥ 1.5h DA Sertar pentru depozitare ≥ 3 buc DA Frînă centralizată pentru fixarea aparatului DA Presiune de alimentare cu gaze 3.0 - 6 bar DA Compresor Portabil pe rotile sau încorporat în mașina de anestezie DA va fi inclus DC 50 Advance, EKOM Slovacia broshura este atasata. Punct de rouă sub presiune la max. și debit de ieșire la 3 bar, 5°C sub temperatura mediului ambient DA Fluxul de ieșire la 3 bar ≥ 60 l/min. DA maxim 3.5 bar. Filtrarea aerului 5 µm DA Mod de operare Continuu DA Surgerea apei condensate Auto DA Presiunea de ieșire 3.5 bar DA maxima Volumul reciverului 2 l DA Accesori Furtunul cu conector de conectare la sursa de aer comprimat 1 buc. DA Furtunul cu conector de conectare la sursa de oxigen 1 buc. DA Circuit de ventilare Adult, reutilizabil ≥ 2 set. DA Plămîn de test Adult, reutilizabil ≥ 2 buc. DA Senzor de flux Reutilizabil ≥ 2 buc. DA Filtru antibacterial Adult, unică utilizare ≥ 200 buc. DA Accesori modul de gaz Adult ≥ 2 set. DA Cablu ECG Adult, reutilizabil 5 fire ≥ 2 buc. DA Senzor ECG Adult, unica utilizare ≥ 100 buc. DA Senzor SpO2 Adult, reutilizabil ≥ 2 Buc. DA Manșete NIBP Adult, reutilizabilă ≥ 2 buc. DA Adult mare, reutilizabilă ≥ 2 buc. DA Senzor de temperatură Adult, reutilizabil ≥ 2 buc. DA Cablu de interconectare senzor IBP Adult, reutilizabil ≥ 1 buc. DA Accesori necesare de funcționare a modulului BIS sau TOF/ NMT Accesori pentru Adult ≥ 5 buc. DA Altele: Garanție - 2 anni DA Manuale de utilizare în limba de stat (Română), în cazul în care nu este prezent să fie prezentat manualul în limba de stat cu stampila biroului de traducere. da</p>
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Anexa 1

<p>Manual de service În limba de stat (română) sau în una din limbile de circulație internațională (rusă sau engleză) conform Legii nr. 102 din 09.06.2017, cap. 4, art. 14, al. (3) și art. 15, al. (6). da Transportare la locul necesar da Instalarea și testarea la locul necesar da Deservire în perioada de garanție și postgaranție da</p> <p>Training pentru utilizatori Pregătire de lucru, mod de utilizare a dispozitivului, întreținerea zilnică, etc. da Training pentru personal tehnic Ajustarea, calibrarea, înlăturarea defectiunilor minore, parole de acces, etc. da</p>	<p>În limba de stat cu stampila biroului de traducere. DA la momentul livrari. Manual de service În limba de stat (română) sau în una din limbile de circulație internațională (rusă sau engleză) conform Legii nr. 102 din 09.06.2017, cap. 4, art. 14, al. (3) și art. 15, al. (6). DA engleză Transportare la locul necesar DA Instalarea și testarea la locul necesar DA Deservire în perioada de garanție și postgaranție DA kiturile de mententa vor fi procurate de catre beneficiar. Training pentru utilizatori Pregătire de lucru, mod de utilizare a dispozitivului, întreținerea zilnică, etc. DA Training pentru personal tehnic Ajustarea, calibrarea, înlăturarea defectiunilor minore, parole de acces, etc. DA</p>
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B105M/B125M/B155M

Monitoare pentru pacienți

Vă alimentăm performanța.



Monitoarele pentru pacienți B105M/B125M/B155M oferă performanțe clinice premium în toate zonele de îngrijire. Aceste monitoare scalabile, precise, integrate, cu design intuitiv, sunt disponibile având afișaje cu ecran tactil de 10, 12 sau 15 inch.

Capabilități avansate

Gama de monitoare B1x5M se poate implementa perfect într-o varietate de setări de îngrijire:

- ECG EK-Pro v14 cu 4 derivații pentru analiza simultană a aritmiei
- Măsurarea neinvasivă a tensiunii arteriale DINAMAP™ SuperSTAT
- Alegerea tehnologiilor de SpO₂: GE TruSignal™, Masimo SET® sau Nellcor™ OxiMax
- GE EtCO₂ pentru măsurarea fluxului lateral, a agentilor anestezici și a debitului cardiac
- Monitorizare* Entropy™ pentru monitorizarea stării creierului
- NMT pentru blocarea transmisiei neuromusculare și monitorizarea inversărilor
- Conectivitate la rețelele GE CARESCAPE™
- Flexibilitate pentru partajarea modulelor de parametri și a accesoriilor cu monitoarele CARESCAPE

*Pentru pacienți cu vârstă peste 2 ani.

gehealthcare.com

Model intuitiv. Flux de lucru neîntrerupt.

- 12 forme de undă pentru a vizualiza simultan toate formele de undă ale parametrilor necesari
- Comunicare Bed to Bed și funcționalitate Automatic view on alarm (AVOA) pentru a revizui datele de monitorizare la distanță ale pacientului
- Funcționalitate circulantă pentru tranziția perfectă a monitorului de la un pat la altul în cadrul rețelei CARESCAPE
- Platforma de serviciu la distanță InSite™ pentru depanare la distanță
- Scorul Național de Avertizare Timpurie (NEWS) pentru o intervenție la timp

Robust pentru sarcini solicitante.

Sigur într-o lume cibernetică.

- Respectă indicațiile FDA privind securitatea cibernetică a dispozitivelor medicale
- Un filtru ECG oferă o performanță sporită a semnalului în zonele zgomotoase
- Cu baterie de mare capacitate: >4 ore¹
- Testat cu standardul EMC Ediția a 4-a
- Rezistent la apă cu standardele IP22

¹ În funcție de configurație, cu configurație tipică ECG, timp ciclu NIBP 15 min, SpO₂, luminozitatea afișajului 70%.

Specificații tehnice

Afișaj

Dimensiune	B155M: 15,6 in (diagonală) B125M: 12,1 in (diagonală) B105M: 10,1 in (diagonală)	Alarme ventriculare Alarme atriale Alarmă PVC	VT>2, R pe T, bradicardie ventriculară, Cuplete, Bigeminie, Ventricular accelerat, Trigeminie, PVC-uri multifocale Fibrilație atrială, lipsă puls, pauză, neregulat, tahicardie SV PVC-uri frecvente, SVC-uri frecvente
Rezoluție	B155M: 1366x768 (HD) B125M / B105M: 1280x800 (WXGA)	Analiza segmentului ST	Interval numeric Acuratețe
Număr forme de undă	până la 12		-9 la +9 mm (-0,9 la +0,9 mV) $\pm 0,2$ mm sau $\pm 10\%$, oricare dintre acestea este mai mare, în intervalul de măsurare de la -8 la 8 mm
Afișarea și culorile ecranului	configurabile de utilizator		Rezoluție numerică Rezoluție numerică
Controale	Ecran tactil capacativ și Trim Knob™		0,1 mm (0,01 mV)

Parametri și module

Parametri	Module ²
ECG	Modul hemodinamic integrat
Resp	
SpO ₂	
NIBP	
Temp	
InvBP cu 2 canale	
CO ₂ în flux secundar	E-miniC ³
Entropie	E-Entropie ⁴
CO ₂ , O ₂ și N ₂ O în flux secundar	E-sCO
Sidestream CO ₂ , O ₂ , agenți și N ₂ O în flux secundar	E-sCAIO, N-CAIO
Debit cardiac + InvBP cu 1 canal	E-COP5
Transmisia neuromusculară	E-NMT

ECG

Derivații disponibile	configurație cu 3 derivații: I, II, III configurație cu 5 derivații: I, II, III, aVR, aVL, aVF și V configurație cu 10 derivații: I, II, III, aVR, aVL, aVF, V1, V2, V3, V4, V5 și V6
Viteza de baleaj	12,5, 25 sau 50 mm/s
Interval amplificare	0,5x, 1x, 2x și 4x
Precizia ritmului cardiac	20 la 300 bpm, $\pm 5\%$ sau ± 5 bpm, oricare este mai mare

Lățimea de bandă

Filtru ECG	Monitor: 0,5 la 40 Hz ST: 0,05 la 40 Hz Diagnostic: 0,05 la 145 Hz Moderat: 0,5-20 Hz
Detectare stimulator cardiac	Interval voltaj: 2 la 700 mV Lățime puls: 0,5 la 2 ms

Alarme de aritmii

Alarme letale	Asistolă, fibrilație ventriculară/tahicardie ventriculară, tahicardie ventriculară
Alarme de RC	Bradycardie, Tahicardie

Alarme

Alarme ventriculare	VT>2, R pe T, bradicardie ventriculară, Cuplete, Bigeminie, Ventricular accelerat, Trigeminie, PVC-uri multifocale
Alarme atriale	Fibrilație atrială, lipsă puls, pauză, neregulat, tahicardie SV
Alarmă PVC	PVC-uri frecvente, SVC-uri frecvente
Analiza segmentului ST	Interval numeric Acuratețe
	-9 la +9 mm (-0,9 la +0,9 mV) $\pm 0,2$ mm sau $\pm 10\%$, oricare dintre acestea este mai mare, în intervalul de măsurare de la -8 la 8 mm
Rezoluție numerică	0,1 mm (0,01 mV)
Impedanță respirație	Interval Acuratețe
	Adult/pediatric: 4 la 120 respirații/min Nou-născuți: 4 la 180 respirații/min $\pm 5\%$ sau ± 5 respirații/min, oricare este mai mare
Interval amplificare	0,1 la 5 cm/Ohm
SpO ₂	TruSignal SpO ₂ Interval măsurare Pulsoximetrie Puls IP (Index circulație sangvină)
	1 la 100% 30 la 250 bpm 0 la 32
Acuratețe măsurare	Saturație
	Fără mișcare-adult/pediatric Senzor de deget: 70 la 100% $\pm 2\%$ Fără mișcare-nou-născuți: 70 la 100% $\pm 3\%$ Cu mișcare-adult/pediatric/nou-născuți: 70 la 100% $\pm 3\%$ Circulație sangvină scăzută-adult/pediatric: 70 la 100% $\pm 3\%$ (<70% nespecificat)
	Puls
	fără mișcare: ± 2 bpm (Adult/Pediatric/Nou-născuți)
Nellcor OxiMax	Interval măsurare Pulsoximetrie Puls
	1 la 100% 20 la 250 bpm
Acuratețe măsurare	Saturație
	Adulți: 70 la 100% $\pm 2\%$ Nou-născuți: 70 la 100% $\pm 3\%$ Circulație sangvină scăzută: 70 la 100% $\pm 2\%$ <70% nespecificat
	Puls
	± 3 bpm
2 Consultați Manualul de utilizare B105M/B125M/B155M pentru mai multe informații.	
3 Măsurarea CO ₂ prin intermediu Modulului E-miniC este destinat utilizării numai la pacienții cu o greutate de peste 5 kg (11 lb).	
4 Modulul E-Entropy va fi utilizat doar la pacienții cu vârstă peste 2 ani.	
5 E-COP nu este destinat utilizării la pacienții nou-născuți.	

Masimo SET

Interval măsurare

Pulsoximetrie 1 la 100%

Puls 25 la 240 bpm

Acuratețe măsurare

Saturație Fără mișcare-adult/pediatric:
70 la 100% $\pm 2\%$
Fără mișcare-nou-născuți:
70 la 100% $\pm 3\%$
Cu mișcare-adult/pediatric/nou-născuți: 70 la 100% $\pm 3\%$
Circulație sangvină scăzută:
70 la 100% $\pm 2\%$
(<70% nespecificat)

Puls fără mișcare: ± 3 bpm
Cu mișcare: ± 5 bpm

IP (Index circulație sangvină) Da
Tehnologia APOD (Adaptive Probe Off Detection)

NIBP

Tehnică de măsurare Oscilometrică cu deflație în trepte
Moduri de măsurare Manuală, Automată (cu timpul ciclului de serie personalizat)
și STAT
Timpi automați ciclu Personalizat, 1, 2, 3, 4, 5, 10, 15, 20,
30 min, 1 oră, 1,5 ore și 2 ore

Intervale măsurare NIBP

Sistolic Adult/pediatric: 30 la 290 mmHg
Nou-născuți: 30 la 140 mmHg
MAP Adult/Pediatric: 20 la 260 mmHg
Nou-născuți: 20 la 125 mmHg
Diastolic Adult/pediatric: 10 la 220 mmHg
Nou-născuți: 10 la 110 mmHg

Acuratețe clinică

Diferență medie ± 5 mmHg
Deviație standard ≤ 8 mmHg
Standard raportare ANSI/AAMI ISO81060-2 și IEC 80601-2-30

Caracteristici de siguranță

Presiunea inițială de umflare implicită Adult/Pediatric: 135 ± 15 mmHg
Nou-născuți: 100 ± 15 mmHg
Timp maxim de determinare Adult/Pediatric: 2 min
Nou-născuți: 85 s
Monitor supra-presiune Adult/Pediatric: 300 ± 6 la 330 mmHg
Nou-născuți: 150 ± 3 la 165 mmHg

Puls din NIBP

Interval Măsurare 30 bpm la 250 bpm

Acuratețe $\pm 5\%$ sau ± 5 bpm
(oricare este mai mare)

Măsurarea invazivă a tensiunii arteriale

Din măsurătorile hemodinamice integrate

Interval măsurare -40 to 320 mmHg
(-5,3 la 42,7 kPa)
Acuratețe măsurare $\pm 4\%$ sau ± 2 mmHg, oricare este mai mare

Răspuns frecvență 4 la 22 Hz

Sensibilitate transductor $5\mu V/V/mmHg$

Interval puls (PR) 30 la 250

Din modulul E-COP

Interval măsurare -30 to 320 mmHg
(-4,0 la 42,7 kPa)

Acuratețe măsurare $\pm 4\%$ sau ± 4 mmHg, oricare este mai mare

Răspuns frecvență 4 la 22 Hz

Sensibilitate transductor $5\mu V/V/mmHg$

Interval puls (PR) 30 la 250

Calcule

SPV (Variația presiunii sistolice) $SBP_{max} - SBP_{min}$
(unde SBP este tensiunea arterială sistolică)

PPV (Variația presiunii pulsului) $(PP_{max} - PP_{min}) / [(PP_{max} + PP_{min})/2]$
 $\times 100$ (unde PP este presiunea pulsului)

Temperatură

Afișaj numeric T1, T2, Tsânge

Din măsurătorile hemodinamice integrate (T1, T2)

Interval măsurare 10 to 45 °C (50 to 113 °F)

Acuratețe măsurare $\pm 0,1$ °C fără sondă
 $\pm 0,2$ °C cu sondă de la 25 la 45 °C
 $\pm 0,3$ °C cu sondă de la 10 la 25 °C
(fără a include 25 °C)

Afișaj resolution 0,1 °C

Din modulul E-COP (Tsânge)

Interval măsurare 17,5 la 43 °C (63,5 la 109,4 °F)

Acuratețe măsurare $\pm 0,5$ °C (17,5 °C la 30,9 °C)
 $\pm 0,3$ °C (31°C la 43,0°C)

Afișaj resolution 0,1 °C

Arhitectura rețelei

Rețea fizică N/W 1000BaseT

Wireless Wi-Fi IEEE 802.11a/b/g/n,
roaming rapid

Servicii de networking

Outbound HL7®	Conecțivitate directă la EMR sau la terțe sisteme pentru trenduri numerice
CARESCAPE (Unity) CARESCAPE Gateway	Conecțivitate la CIS / HIS prin Alte aplicații de networking
Serviciu la distanță	Diagnoza la distanță a dispozitivului prin serverul InSite™ RSvP

Aplicații de networking CARESCAPE (Unity)

Fereastră Bed to Bed*

Date afișate	Forme de undă și valori numerice de la șase parametri, o alarmă la distanță și informații de la distanță despre pat
Paturi la distanță	Alarame de monitorizare pentru până la 40 de paturi
Monitorizat	Vizualizarea unui pat din 1023 paturi
AVOA (Auto View of Remote beds in alarm)*	
Informații despre mesajul de alarmă la distanță	Numele unității și al patului, mesaj de alarmă, alarmare mai mult de 1 pat
Notificare de alarmă configurabilă	Mesaj, Vizualizare automată, Vizualizare automată întotdeauna
Rotire	
Funcționalitate	Rotire între unități și paturi; Adăugarea de noi unități și paturi; Selectarea imprimantei

Periferice I/O

Conectori standard

Port Ethernet / WIFI	Suportă HL7 and CARESCAPE Unity N/W
Port USB 2.0	Jurnale serviciu descărcare
Setări de importare/exportare	
Trenduri numerice de export	Instalare software, firmware și e-manuale
Port HDMI	Suportă afișarea clonelor secundare B155M: 1366 x 768 pixeli B125M/B105M: 1280 x 800 pixeli
Port serial RS232	Exportă datele trendurilor o/p și alarmele către iCollect doar prin intermediul protocolului DRI

Conectori non-standard

Conector asistent medical	se conectează la sistemul de asistență medicală al spitalului
Conector sincronizare defibrilator	Ieșire sincronizare defibrilator Conector recorder Imprimantă termică autonomă B1X5-REC Recorder
Cadru B1X5-F2	al doilea cadru pentru conector suplimentar pentru module

Securitatea rețelelor și a datelor

Certificat Wi-Fi	CE, FCC
Autentificare Wi-Fi	Support WPA-Personal; WPA2-Personal; WPA-Enterprise; WPA2- Enterprise
Criptare date WIFI	Suport WPA/WPA2 cu TKIP și AES CCMP
Conexiune LAN / WLAN	Suportă IEEE 802.1X bazat pe port Control acces rețea (NAC)
Schimb de fișiere prin USB	Toate funcțiile USB sunt protejate prin parolă Export criptat de tendințe numerice, setări de utilizator și jurnale de servicii prin USB

Montare

Mâner de transport integrat compatibil cu GCX

Imprimantă termică locală

Metodă	Matrice de puncte termice
Rezoluții orizontale	24 puncte/mm (600 dpi)
Rezoluție verticală	8 puncte/mm (200 dpi)
Forme de undă	Selectabile 1, 2 sau 3 forme de undă
Tipărire trenduri numerice	HR, Pleth, NIBP, IBP1, IBP2, T1, T2, Et/FiCO ₂ , RR, Pleth, C.O., C.I., REF, SPV, PPV, IBP4, Tblood, RE, SE, BSR, NMT Count, O ₂ , N ₂ O, AA, BAL, MAC
Lățime hârtie	50 mm, lățime de imprimare 48 mm
Viteză hârtie	5, 10, 12,5 și 25mm/s, configurabilă de utilizator
Imprimantă la distanță	Suportă atât imprimantă laser, cât și imprimantă termică (cu stația centrală CARESCAPE)

Rack pentru module (integrat)

Slot pentru un singur modul



Cadru secundar B1X5-F2 (optional)

Al doilea cadru suplimentar pentru până la două module



Specificații de performanță

Alarme

Prioritate	Prioritate reglabilă: Ridicată, medie, scăzută și informare Control local și de la distanță din stația centrală
Detectare alarmă	Asistolă, fibrilație ventriculară/tahicardie ventriculară, tahicardie ventriculară, Bradicardie, FiO ₂ scăzut, EtO ₂ scăzut și FiN ₂ O ridicat
Configurabilitate alarmă	Definiți intervalul ratei pentru tahicardie ventriculară și criteriile de durată pentru o alarmă durabilă privind tahicardie ventriculară
Alarmă	tahicardie ventriculară
Notificare	Sonoră și vizuală
Ton alarmă	IEC, General, ISO, ISO2
Setare	Implicită și individuală
Notificare alarmă vizuală	roșu, galben, cyan
Reglarea limitei de alarmă	Mesaj audio silentios Mesaj general de alarmă
Temporizator pauză audio	Control local și de la distanță din stația centrală
Imprimare automată a alarmei până la 23 alarme	2 min

Trenduri

Grafice	Toți parametrii, scale de timp selectabile de la 20 min la 168h (7 zile)
Numerice	Toți parametrii, cu 168 de ore (7 zile) de eșantionare a datelor de trend în funcție de setarea timpului sau după determinarea NIBP, CO și PCWP
Instantaneu	Până la 200 instantanee Declanșat manual sau prin alarmă
Trend OxyCRG	Instantanee de evenimente cu formă de undă (pe stația centrală CARESCAPE) Doar mod nou-născuți Vizualizare în timp real sau instantaneu Stochează până la 70 instantanee OxyCRG Durata instantaneului cu 6 minute înainte și 2 minute după evenimentul OxyCRG
Cursor trend	În trendul grafic

Divulgare completă

Filă/pagină: toate ECG, Hemo

Toate vizualizările ECG	ECG I, II, III, aVR, aVL, aVF, forme de undă V1, V2, V3, V4, V5 și V6
Vizualizare Hemo	ECG II, IBP1, IBP2, IBP4, SpO ₂ și forme de undă Resp
Parametri suportați	ECG, SpO ₂ , IBP și RESP
Viteza de baleaj a revizuirii formelor de undă configurabilă	
Stocare	72 de ore cu toate datele despre forma de undă
Legătură integrată cu istoricul alarmelor	
Revizuirea divulgare completă cu privire la alarma specifică	
Revizuirea divulgare completă cu privire la timpul specific	

EWS (Scor de alarmare timpurie)

Protocol	Scorul național de avertizare timpurie (NEWS) 2
Parametri	HR/PR Puls, tensiunea arterială sistolică, LOC (nivel de conștiință), TEMP, SpO ₂ , Rată Resp și Aer sau Oxigen
	Istoric cu valori detaliate ale parametrilor și sub-scoruri
	Scorul total EWS pe ecranul principal cu codare color și marcaje de timp
	Răspunsul clinic și scorurile parametrilor individuali cu culori pe o fereastră dedicată
	Revizuire risc clinic EWS și îndrumările EWS

Specificații de mediu

Condiții de funcționare

Temperatură	5 la 40°C (41 la 104°F)
Umiditate relativă	15 la 90% fără condens
Presiune atmosferică	700 la 1060 hPa (525 la 795 mmHg)

Condiții de depozitare și transport

Temperatură	-20 la 60°C (-4 la 140°F)
Umiditate relativă	10 la 90% fără condens
Presiune atmosferică	700 la 1060 hPa (525 la 795 mmHg)

Specificații de putere

Intrare AC	100 la 240 V ±10%, 50/60 Hz
Consum de energie	Monitor ≤150 VA
	Cadru secundar B1X5-F2 ≤50 VA
Protecție	Clasa I
Acumulator	1, litiu-ion de mare capacitate < 4 ore la 90% din capacitate
Timp încărcare	>4,0 ore pentru B155M / B125M
Timp funcționare	>4,5 ore pentru B105M cu configurație tipică: ECG, timp ciclu NIBP 15 min, SpO ₂ , luminozitatea afișajului 70%



Specificații fizice

Monitor

Dimensiuni (î x l x A)	B155M: 305 x 405 x 175 mm B125M: 280 x 312 x 175 mm B105M: 275 x 265 x 175 mm
Greutate (cu acumulator și fără module)	B155M: ≤ 5,2 kg (11,5 lb) B125M: ≤ 4,2 kg (9,3 lb) B105M: ≤ 3,8 kg (8,4 lb)
Protecție infiltrare	IP22

Cadru secundar B1X5-F2

Dimensiuni (î x l x A)	160 x 132 X 266 mm cu plăcuță de montaj
Greutate	1,4 kg (30,9 lb) u plăcuță de montaj

Certificări

IEC 60601-1 admis

Marcaj CE conform Regulamentului UE privind dispozitivele medicale (UE) 2017/745

Marcaj UL

Certificări CB

Sistem

Sistem de operare	Linux®
Sistem de răcire	Convecție naturală, fără ventilator în interior pentru răcire

Este posibil ca produsul să nu fie disponibil în toate țările și regiunile. Specificațiile tehnice complete ale produsului sunt disponibile la cerere. Contactați un reprezentant GE Healthcare pentru mai multe informații. Vă rugăm intrați pe www.gehealthcare.com/promotional-locations. Datele pot fi modificate.

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B125M, B105M, B155M JB00262KE

2020-09-21

Versiunea de software VSP3.0 pentru monitoarele de pacienți B105M/B125M/B155M nu sunt disponibile pe toate piețele și nu sunt confirmare 510k.



GE Healthcare

Carestation™ 650

The Carestation 650 is a compact, versatile and easy to use anesthesia system designed to help clinicians deliver reliable anesthesia care to solve today's toughest challenges.

KEY FEATURES

- Elegant modern design in a slim, compact frame well suited for constrained environments
- Simple and easy to use 15" touchscreen ventilator display
- Intuitive CARESCAPE™ inspired user interface for the unified Carestation user experience
- Integrated CARESCAPE Respiratory Module
- Time saving tools to help streamline clinician workload
- Scalable software and hardware features: "build your own" Carestation
- ecoFLOW display option may help clinicians mitigate the risk of hypoxic mixtures while helping to reduce agent use by using low and minimum flows with continuous gas monitoring

VENTILATION

- Small, compact breathing system specifically designed for low flow anesthesia
- Fast gas kinetics for rapid wash-in and wash-out
- Digitally controlled flow valve ventilator supports all patient types from neonates to adults
- Advanced ventilation options including synchronized PCV-VG with pressure support (SIMV PCV-VG) and minimum rate ventilation (CPAP+PSV)
- Software enabled tools including Vital Capacity and Cycling Procedures to help automate repetitive tasks used during lung ventilation procedures
- Continual fresh gas flow with fresh gas flow compensation during mechanical ventilation



DESIGN

- Durable wheels, handles and central brake for mobility and stability
- Robust handles and mounting rails
- Easy to clean surfaces
- Movable display arm that rotates and tilts for ideal positioning
- Two vaporizer configuration
- Bi-level work surface illumination
- Absorbent canister designed for ease of use and long life
- Intelligent lighting that highlights active flow controls and auxiliary ports when in use

PHYSICAL SPECIFICATIONS

Product Description

Carestation 650 A1

Dimensions

Height:	135 cm/53.1 in
Width:	82.5 cm/32.4 in
Depth:	75 cm/29.5 in
Weight:	145 kg/320 lb*

Top shelf

Weight limit:	25 kg/55 lb
Width:	41.3 cm/16.3 in
Depth:	27.0 cm/10.6 in

Work surface

Height:	83.6 cm/32.9 in
Size:	1620 cm ² /251 in ²
Size: (with optional flip shelf)	2527 cm ² /392 in ²

Upper left Datex-Ohmeda (DO) dovetail

Dovetail length: 54 cm/21.3 in

Lower left Datex-Ohmeda (DO) dovetail

Dovetail length: 32 cm/12.6 in

Right Datex-Ohmeda (DO) dovetail

Dovetail length: 96.4 cm/38.0 in

Drawers (internal dimensions)

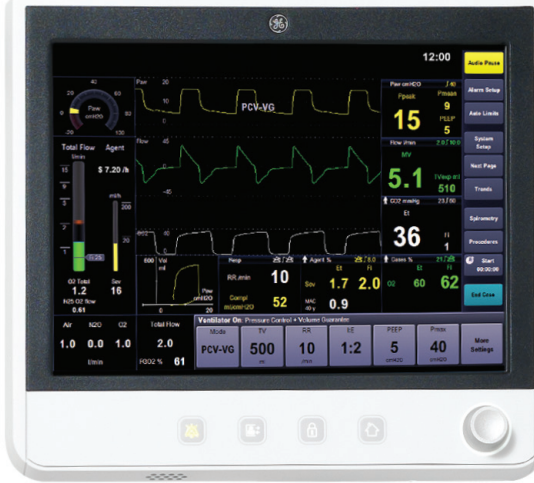
Height:	
Top and middle:	8.6 cm/3.4 in
Bottom:	13.3 cm/5.2 in
Width:	34 cm/13 in
Depth:	37 cm/14.6 in

Manual ventilation bag arm (optional)

Arm length:	39.8 cm/15.7 in
Bag arm height (adjustable):	53 cm/20.9 in 136 cm/53.5 in

Casters

Diameter:	12.5 cm/4.9 in
Brakes:	Central Brake



VENTILATOR OPERATING SPECIFICATIONS

Modes of ventilation – included

VCV (Volume Control) Mode with tidal volume compensation

Modes of ventilation – optional

PCV (Pressure Control Ventilation)

PCV-VG (Pressure Controlled Ventilation-Volume Guarantee)

SIMV (Synchronized Intermittent Mandatory Ventilation)
(volume and pressure)

PSVPro™ (Pressure Support with Apnea backup)

CPAP+PSV (Pressure support mode)

SIMV PCV-VG

Advanced software options

Spirometry (included)

Auto alarm limits (included)

ecoFLOW

Pause Gas

Vital capacity and cycling

VCV Cardiac Bypass

Ventilator parameter ranges

Tidal volume range:	5 to 1500 mL (PCV modes 5 to 1500 mL) (Volume Control, PCV-VG and SIMV volume 20 to 1500 mL)
Incremental settings:	20 to 50 mL (increments of 1 mL) 50 to 100 mL (increments of 5 mL) 100 to 300 mL (increments of 10 mL) 300 to 1000 mL (increments of 25 mL) 1000 to 1500 mL (increments of 50 mL)
Minute volume range:	Less than 0.1 to 99.9 L/min
Pressure (P_{inspired}) range:	5 to 60 cmH ₂ O (increments of 1 cmH ₂ O) above set PEEP
Pressure (P_{max}) range:	12 to 100 cmH ₂ O (increments of 1 cmH ₂ O)
Pressure (P_{support}) range:	Off, 2 to 40 cmH ₂ O (increments of 1 cmH ₂ O)
Respiratory Rate:	4 to 100 breaths per minute for Volume Control and Pressure Control; 2 to 60 breaths per minute for SIMV, PSVPro and SIMV PCV-VG; 4 to 60 bpm for CPAP+PSV (increments of 1 breath per minute)
Inspiratory/expiratory ratio:	2:1 to 1:8 (increments of 0.5) (VCV, PCV, PCV-VG)
Inspiratory time:	0.2 to 5.0 seconds (increments of 0.1 seconds) (SIMV, PSVPro and CPAP PSV)
Trigger window:	Off, 5 to 80% of T_{exp} (SIMV, PSVPro) (increments of 5%)
Flow trigger:	1 to 10 L/min (increments of 0.5 L/min) 0.2 to 1 L/min (increments of 0.2 L/min)
Inspiration termination level:	5 to 75% (increments of 5%)
Inspiratory Pause range:	Off, 5-60% of T_{insp}

Positive End Expiratory Pressure (PEEP)

Type:	Integrated, electronically controlled
Range:	OFF, 4 to 30 cmH ₂ O (increments of 1 cmH ₂ O)

Ventilator performance

Peak gas flow:	120 L/min + fresh gas flow
Flow valve range:	1 to 120 L/min
Flow compensation range:	100 mL/min to 15 L/min

VENTILATOR ACCURACY

Delivery/monitoring accuracy

Volume delivery:	> 210 mL = better than 7% ≤ 210 mL = better than 15 mL < 60 mL = better than 10 mL
Pressure delivery:	±10% or ±3 cmH ₂ O (larger of)
PEEP delivery:	±1.5 cmH ₂ O
Volume monitoring:	> 210 mL = better than 9% ≤ 210 mL = better than 18 mL < 60 mL = better than 10 mL
Pressure monitoring:	±5% or ±2.4 cmH ₂ O (larger of)

Alarm settings

Tidal volume (V_{TE}):	Low: OFF, 1 to 1500 mL High: 20 to 1600 mL, OFF
Minute volume (V_{E}):	Low: OFF, 0.1 to 10 L/min High: 0.5 to 30 L/min, OFF
Inspired oxygen (FiO ₂):	Low: 18 to 99% High: 19 to 100%, OFF
Apnea alarm:	
Low airway pressure:	
High pressure:	4 cmH ₂ O above PEEP 12 to 100 cmH ₂ O (increments of 1 cmH ₂ O)
Sustained airway pressure:	

Mechanical ventilation ON:

$P_{\text{max}} < 30 \text{ cmH}_2\text{O}$,
the sustained limit is 6 cmH₂O
 $P_{\text{max}} 30 \text{ to } 60 \text{ cmH}_2\text{O}$,
the sustained limit is 20% of P_{max}
 $P_{\text{max}} > 60 \text{ cmH}_2\text{O}$,
the sustained limit is 12 cmH₂O

PEEP and mechanical ventilation ON:

Sustained limit increases by PEEP minus 2 cmH₂O

Mechanical ventilation OFF:

$P_{\text{max}} 12 \text{ to } 60 \text{ cmH}_2\text{O}$,
the sustained limit is 50% of P_{max}
 $P_{\text{max}} > 60 \text{ cmH}_2\text{O}$,
the sustained limit is 30 cmH₂O

Subatmospheric pressure: Paw < -10 cmH₂O

Audio pause countdown clock: 120 to 0 seconds

VENTILATOR COMPONENTS

Flow transducer

Type:	Variable orifice flow sensor (autoclavable)
Location:	Inspiratory outlet and expiratory inlet

Oxygen sensor

Type:	Optional galvanic fuel cell or paramagnetic with Airway Module option
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Ventilator screen

Display size:	15 inch
Pixel format:	1024 x 768

Battery backup

Backup power:	Battery time is 90 minutes when fully charged, which supports full system functionality and ventilation.
Battery type:	Internal rechargeable sealed lead acid

Communication ports

RS-232C compatible serial interface
Ethernet
Datex-Ohmeda device interface solutions port
USB port
VGA Output

ANESTHETIC AGENT DELIVERY

Delivery

Vaporizers:	Tec™ 6 Plus, Tec 7, Tec 820, Tec 850
Number of positions:	2
Mounting:	Tool-free installation Selectatec™ manifold interlocks and isolates vaporizers

AIRWAY MODULES

General

Supported modules:	E-sCAiO, E-sCAiOV, N-CAiO
Size (HxWxD), excluding water trap:	113 x 38 x 205 mm/4.4 x 1.5 x 8.1 in
Weight:	0.7 kg/1.5 lb
Sampling rate:	120 mL/min ±20 mL
Automatic compensation for atmospheric pressure variation (495 to 795 mmHg) temperature and CO ₂ /N ₂ O and CO ₂ /O ₂ collision broadening effect. Parameter display update interval typically breath-by-breath. Functional alarms for blocked sample line, D-fend check and D-fend replacement.	

Non-disturbing gases:

Ethanol, acetone, isopropanol, methane, nitrogen, nitric oxide, carbonmonoxide, water vapor, freon R134A (for CO ₂ , O ₂ and N ₂ O):	
Maximum effect on readings:	CO ₂ < 0.2 vol %; O ₂ , N ₂ O < 2 vol %, AA < 0.15 vol%

Carbon dioxide (CO₂)

EtCO ₂ :	End-tidal CO ₂ concentration
FiCO ₂ :	Inspired CO ₂ concentration

CO₂ waveform

Measurement range:	0 to 15%
	(0 to 15 kPa, 0 to 113 mmHg)
Accuracy:	±0.2 vol % + 2 % of reading
Datex-Ohmeda infrared sensor	

Adjustable low and high alarm limits for EtCO₂ and FiCO₂

Respiration rate (RR)

Measurement range:	4 to 100 breaths/min
Detection criteria:	1% variation in CO ₂
Adjustable low and high alarm limits for respiration rate; alarm for apnea	

Patient Oxygen (O₂)

FiO ₂ :	Inspired O ₂ concentration
EtO ₂ :	End-tidal O ₂ concentration
FiO ₂ -EtO ₂ :	Inspired-expired difference

O₂ Measurement

Measurement range:	0 to 100%
Accuracy:	±1 vol % + 2 % of reading
Datex-Ohmeda differential paramagnetic sensor	
Adjustable low and high alarm limits for FiO ₂ and EtO ₂ ; alarm for FiO ₂ < 18%	

Nitrous Oxide (N_2O)

Measurement range: 0 to 100%
Accuracy: ± 2 vol % +2 % of reading

Anesthetic Agent (AA)

Halothane, Isoflurane, Enflurane

Measurement range: 0 to 6%
Accuracy: $\pm (0.15$ vol% +5% of reading)

Sevoflurane

Measurement range: 0 to 8%
Accuracy: $\pm (0.15$ vol% +5% of reading)

Desflurane

Measurement range: 0 to 20%
Accuracy: $\pm (0.15$ vol% +5% of reading)

Waveform displayed

MAC value displayed (Airway Gas Option modules)

MACage value displayed (CARESCAPE modules)

Identification threshold: 0.15 vol%**

Agent mixture detection

Adjustable high and low alarm limits for EtAA, FiAA

Patient Spirometry™

Pressure-volume loop

Pressure-flow loop

Flow-volume loop

Airway pressure and flow waveforms

Adjustable low and high alarm limits for P_{peak} , PEEP_{tot} and MV_{exp}

Alarms for MV_{exp} << MV_{insp} and for MV_{exp} low. Detection through D-lite™ or Pedi-lite™ flow sensor and gas sampler with following specifications:

CARESCAPE Airway Modules

D-lite(+)

Respiration rate: 4 to 35 breaths/min

Pedi-lite(+)

4 to 70 breaths/min

Tidal volume

Measurement range: 150 to 2000 mL
Accuracy**: $\pm 6\%$ or 30 mL

Minute volume

Measurement range: 2 to 20 L/min 0.1 to 5 L/min

Airway pressure

Measurement range: -20 to +100 cmH₂O
Accuracy**: ± 1 cmH₂O
Display units: cmH₂O, mmHg, kPa, mbar, hPa

Flow

Measurement range: -100 to 100 L/min -25 to 25 L/min

I:E

Measurement range: 1:4.5 to 2:1

Compliance

Measurement range: 4 to 100 mL/cmH₂O 1 to 100 mL/cmH₂O

Airway resistance

Measurement range: 0 to 200 cmH₂O/L/s

Sensor specifications

D-lite/ D-lite(+)

Dead Space: 9.5 mL

Pedi-lite/ Pedi-lite(+)

2.5 mL

Resistance

at 30 L/min: 0.5 cmH₂O

at 10 L/min: 1.0 cmH₂O

ELECTRICAL SPECIFICATIONS

Current leakage

100/120 V: < 300µA

220/240 V: < 500µA

Power

Power input: 100-120 Vac, 50/60 Hz
220-240 Vac, 50/60 Hz
120/220-240 Vac ± 10%, 50-60 Hz

Power cord:

Length: 5 m/16.4 ft
Rating: 10A @ 220-240 Vac or
15A @ 100-120 Vac
10A @ 120/220-240 Vac

Inlet modules

100/120 V:
Without outlets: 2A
With outlets: 10A

220/240 V:
Without outlets: 2A
With outlets: 8A

Outlet modules (optional)

100/120 V:
3 outlets on side, 1-3A, 2-2A, individual breakers, isolation transformer (optional)

220/240 V:
3 outlets on side, 1-2A, 2-1A, individual breakers, isolation transformer (optional)

120/220-240 V:

No outlets

PNEUMATIC SPECIFICATIONS

Auxiliary O₂ (optional)

Connection:	7-10 mm hose barb port
O ₂ concentration range:	100% O ₂
Flow range:	0 to >10 L/min
Connection:	7-10 mm hose barb port
O ₂ concentration range:	100% O ₂ only, or 21% to 100% O ₂ with Air
Flow range:	0 and 100 mL/min to 15 L/min
for O ₂ and Air:	0 and 100 mL/min to 15 L/min

Auxiliary common gas outlet (optional)

Connector:	ISO 22 mm OD and 15 mm ID
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Gas supply

Pipeline input range:	280 kPa to 600 kPa (41 psig to 87 psig)
Pipeline connections:	DISS-male, DISS-female, AS4059, S90-116, or NIST All fittings available for O ₂ , N ₂ O, and Air, and contain pipeline filter and check valve.
Cylinder input:	Secondary O ₂ pipeline inlet available. Pin indexed in accordance with CGA V-1 or DIN-477 (nut and gland); con- tains input filter and check valve. Large cylinder kit available for O ₂ and N ₂ O (with DIN-477).

Note: Maximum 3 cylinders

Primary regulator diaphragm minimum burst pressure:	2758 kPa/400 psig
Primary regulator nominal output:	≤ 345 kPa/50 psig Pin indexed cylinder connections ≤ 414 kPa/60 psig DIN-477 cylinder connections

O₂ controls

Method:	N ₂ O shut off with loss of O ₂ pressure
Supply failure alarm:	< 252 kPa (36.55 psig)
O ₂ flush:	Range: 25 to 75 L/min

Fresh gas

Flow range: for O ₂ and Air:	0 and 100 mL/min to 15 L/min (minimal flow capable)
for N ₂ O:	0 and 100 mL/min to 10 L/min

Pneumatic Total Flow Tube:

1 to 10 L/min

Measurement accuracy

for O ₂ , Air and N ₂ O:	±6% of measured value, or ±25 mL/min (larger of)
for Total Flow tube:	±5% of full scale (larger of) at 100% O ₂
O ₂ concentration range:	21% to 100% when Air is available
O ₂ Cell accuracy:	±2.5% plus 2.5% of reading
Compensation:	Temperature and atmospheric pres- sure compensated to standard con- ditions of 20°C and 101.3 kPa
Hypoxic guard:	Mechanical Link-25: Provides a nominal minimum 25% concentration of oxygen in O ₂ /N ₂ O mixture.

Materials

All materials in contact with patient breathing gases are not made from natural rubber latex.

ENVIRONMENTAL SPECIFICATIONS

System operation

Temperature:	10° to 40°C (50° to 104°F)
Humidity:	15 to 95% relative humidity (non-condensing)
Altitude:	-440 to 3565 m (500 to 800 mmHg)
	-440 to 4000 m (without Airway Module) (475 to 800 mmHg)

System storage

Temperature:	-25° to 60°C (-13° to 140°F)
Humidity:	15 to 95% relative humidity (non-condensing)
Altitude:	-440 to 4880 m (425 to 800 mmHg)
Oxygen cell storage:	-15° to 50°C (5° to 122°F) 10 to 95% relative humidity 500 to 800 mmHg

Electromagnetic compatibility

Immunity:	Complies with all applicable requirements of EN 60601-1-2
Emissions:	CISPR 11 group 1 class A
Standard compliance:	AAMI ES60601-1, CSA C22.2 #601.1, EN/IEC 60601-1, ISO 80601-2-13
European Notified Body CE Mark:	CE0197

BREATHING CIRCUIT SPECIFICATIONS

Carbon dioxide absorbent canister

Absorbent capacity:	Reusable canister 1370 mL/1150 g Disposable canister 1440 mL/1200 g
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Ports and connectors

Exhalation:	22 mm OD ISO 15 mm ID taper
Inhalation:	22 mm OD ISO 15 mm ID taper
Bag port:	22 mm OD (15 mm ID), ROW 22 mm ID, Australia

Bag-to-Ventilator switch

Type:	Bi-stable
Control:	Controls ventilator and direction of breathing gas within the circuit

Integrated Adjustable Pressure Limiting (APL) valve

Range:	0.5 to 70 cmH ₂ O
Tactile knob indication at:	30 cmH ₂ O and above
Adjustment range of rotation:	0.5 to 30 cmH ₂ O (0 to 230°) 30 to 70 cmH ₂ O (230 to 330°)

Materials

All materials in contact with exhaled patient gases are autoclavable, except O₂ cell, and Airway Modules.
All materials in contact with patient gas are not made from natural rubber latex.

Breathing circuit parameters

Compliance:

Bag mode:	1.81 mL/cmH ₂ O (filled disposable absorber canister)
	1.74 mL/cmH ₂ O (filled reusable absorber canister)
Mechanical mode:	Automatically compensates for compression losses within the absorber and bellows assembly
Volume:	2006 mL Ventilator side 500 mL Bag side 1000 mL Reusable canister 1000 mL Disposable canister

Expiratory resistance in bag mode:

<i>P_{exp}</i> Flow rate	<i>P_{exp}</i> Absorber canister Installed	<i>P_{exp}</i> Absorber canister Removed
5 L/min	0.57 cmH ₂ O	0.57 cmH ₂ O
30 L/min	2.47 cmH ₂ O	2.47 cmH ₂ O
60 L/min	5.60 cmH ₂ O	5.60 cmH ₂ O

Note: Values include patient circuit tubing and wye piece (0.65 cmH₂O at 60 L/min)

Anesthetic gas scavenging

AGSS Type	Hospital extract system required	Machine connection
High vacuum, low flow:	High vacuum 36 +/- 3 L/min @ 12 inHg (305 mmHg)	SIS evac
High vacuum, low flow:	High vacuum 25- 30 L/min @ 12 inHg (305 mmHg)	DISS evac
Low vacuum, high flow:	Low vacuum 50 to 80 L/min ISO 1H	BSI 30 mm threaded
Low vacuum, low flow:	Low vacuum 25 to 50 L/min ISO 1L	12.7 mm hose barb, 25 mm hose barb, or 30 mm ISO taper
Passive:	Passive system with air break	30 mm/1.2 in M ISO taper



Product may not be available in all countries and regions.

Contact a GE Healthcare Representative for more information.
Please visit www.gehealthcare.com

GE Healthcare
PO Box 7550
Madison, WI 53707-7550
USA

Always refer to complete instruction manual before use.

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This document applies to Carestation 650 A1.

DOC1649438 Rev7

Non-USA



Tec 820 Vaporizer

Combining clinical performance with ergonomic design

The agent specific Tec™ 820 Vaporizer from GE Healthcare delivers isoflurane and sevoflurane effectively and efficiently. Each Tec 820 presents an improved ergonomic design, with consistent agent delivery and reliability while reducing total cost of ownership

Features

- Tec 820 agent specific vaporizers are designed to deliver isoflurane and sevoflurane anesthetic agents
- Provides output consistent with the dial setting throughout the clinical flow range
- Easy to turn dial and fine graduations help control agent delivery
- Enhanced design with consistent, reliable operation over the product life
- Tec 820 vaporizer supports the Easy-Fil agent filling mechanism
- Improved ergonomics and contemporary design complements GE Healthcare's anesthesia systems
- Attaches to GE Healthcare interlocking Selectatec™ manifold
- No planned factory service needed
- Three-year warranty



Clinical performance

- Tec 820 Vaporizers are designed to provide consistent output throughout the clinical flow range from 200 ml/min to 15 l/min. Models are available for isoflurane and sevoflurane
- Tec 820 Vaporizers are equipped with a large diameter control dial with graduations up to 5% V/V (isoflurane models) or 8% V/V (sevoflurane models)
- The easy turning dials and small graduations on the Tec 820 Vaporizer help you fine tune anesthetic delivery over the full range of dial settings and flow rates

Enhanced ergonomics

- The ergonomic dial release on the Tec 820 Vaporizer allows either left or right hand operation
- Wide, centered liquid level indicator provides a clear indication of the fill level of the vaporizer
- Easy to turn dial with enhanced readability

Lower overall ownership costs

- The Tec 820 Vaporizer has no planned factory service, which helps lower the total cost of ownership and eliminate the logistical challenges associated with the return of vaporizers to the factory. For complete user maintenance requirements, refer to the User Reference Manual
- The Tec 820 Vaporizer comes with a three-year warranty

Reliability, convenience, and operating excellence

- Tec 820 is compatible with modern agent filling systems that simplify filling and help minimize agent leaks while filling
- To minimize filling frequency, the Tec 820 Vaporizer can hold up to 300 ml of anesthetic agent
- The Tec 820 Vaporizer can be mounted on GE Healthcare anesthesia systems equipped with the Selectatec manifold without tools and without taking the anesthesia system out of service
- The Tec 820 Vaporizer also interlocks in series with GE Healthcare Tec 7 and Tec 6 Plus Vaporizers
- The non-spill system in the Tec 820 Vaporizer limits movement of liquid agent if the vaporizer is tilted or inverted, protecting internal components and helping maintain output within clinically acceptable limits



Easy-Fil helps simplify agent filling



Tec 820 agent specific vaporizers

Physical specifications

Dimensions

Height:	25 cm
Depth:	22 cm
Width:	11 cm
Weight:	7 kg dry

Agent capacity

Total capacity:	300 ml
Capacity between min/max fill marks:	170 ml
Wick system capacity:	75 ml approximately

Pneumatics specifications

Calibration and flow resistance

Calibration:	Calibration of all models is done at 21°C using O ₂ as carrier gas, 5 l/min fresh gas flow, at sea-level ambient pressure
Isoflurane models:	Control dial calibrated up to 5% V/V. The control dial is marked in steps of 0.25% up to 1% and in steps of 0.5% between 1% and 5%.
Sevoflurane models:	Control dial calibrated up to 8% V/V. Control dial is marked in steps of 0.25% up to 1% and in steps of 0.5% between 1% and 8%.
Flow resistance:	OFF: Isolated, no resistance from vaporizer ON: 5 l/min O ₂ : <20 cmH ₂ O

Accuracy

Isoflurane models:	±0.25% of delivered agent or ±20% of control dial setting (whichever is greater)
Sevoflurane models:	+0.6/-0.4% of delivered agent or ±20% of control dial setting (whichever is greater)
Fresh gas flow:	200 ml/min to 15 l/min

Environmental specifications

Operation

Temperature:	15°C to 35°C
Humidity:	15% to 95% relative humidity (non-condensing)
Ambient pressure:	500-800 mmHg

Storage and Transport

Temperature:	-25°C to 60°C
Humidity:	15% to 95% relative humidity (non-condensing)
Ambient pressure:	400-800 mmHg

Service

Planned factory service free

MRI Safety Information

MR Conditional

Agent filling systems

Isoflurane models: Easy-Fil™
Sevoflurane models: Easy-Fil

Note: may not be available in all regions

Warranty

Warranted to be free from functional defects in materials and workmanship for a period of three years from the date of original delivery.



Agent color coded Easy-Fil adapters



GE Healthcare provides transformational medical technologies and services to meet the demand for increased access, enhanced quality and more affordable healthcare around the world. GE (NYSE: GE) works on things that matter – great people and technologies taking on tough challenges.

From medical imaging, software & IT, patient monitoring and diagnostics to drug discovery, biopharmaceutical manufacturing technologies and performance improvement solutions, GE Healthcare helps medical professionals deliver great healthcare to their patients.

Imagination at work

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DOC2018946



Tec 850 Vaporizer

Combining clinical performance with ergonomic design

The agent specific Tec™ 850 Vaporizer from GE Healthcare delivers isoflurane and sevoflurane effectively and efficiently. Each Tec 850 presents an improved ergonomic design, with consistent agent delivery and reliability while reducing total cost of ownership.

Features

- Tec 850 agent specific vaporizers are designed to deliver isoflurane and sevoflurane anesthetic agents
- Provides output consistent with the dial setting throughout the clinical flow range
- Easy to turn dial and fine graduations help control agent delivery
- Enhanced design with consistent, reliable operation over the product life
- Tec 850 vaporizer supports a variety of contemporary agent filling mechanisms
- Improved ergonomics and contemporary design complements GE Healthcare's anesthesia systems
- Attaches to GE Healthcare interlocking Selectatec™ manifold
- No planned factory service needed
- Five-year warranty period



Clinical performance

- Tec 850 Vaporizers are designed to provide consistent output throughout the clinical flow range from 200 ml/min to 15 l/min. Models are available for isoflurane and sevoflurane
- Tec 850 Vaporizers are equipped with a large diameter control dial with graduations up to 5% V/V (isoflurane models) or 8% V/V (sevoflurane models)
- The easy turning dials and small graduations on the Tec 850 Vaporizer help you fine tune anesthetic delivery over the full range of dial settings and flow rates

Enhanced ergonomics

- The ergonomic dial release on the Tec 850 Vaporizer allows either left or right hand operation
- Wide, centered liquid level indicator provides a clear indication of the fill level of the vaporizer
- Easy to turn dial with enhanced readability

Lower overall ownership costs

- The Tec 850 Vaporizer has no planned factory service, which helps lower the total cost of ownership and eliminate the logistical challenges associated with the return of vaporizers to the factory. For complete user maintenance requirements, refer to the User Reference Manual
- The Tec 850 Vaporizer comes with a five-year warranty

Reliability, convenience, and operating excellence

- Tec 850 is compatible with modern agent filling systems that simplify filling and help minimize agent leaks while filling
- To minimize filling frequency, the Tec 850 Vaporizer can hold up to 300 ml of anesthetic agent
- The Tec 850 Vaporizer can be mounted on GE Healthcare anesthesia systems equipped with the Selectatec manifold without tools and without taking the anesthesia system out of service
- The Tec 850 Vaporizer also interlocks in series with GE Healthcare Tec 7 and Tec 6 Plus Vaporizers
- The non-spill system in the Tec 850 Vaporizer limits movement of liquid agent if the vaporizer is tilted or inverted, protecting internal components and helping maintain output within clinically acceptable limits



Easy-Fil helps simplify agent filling



Tec 850 agent specific vaporizers

Physical specifications

Dimensions

Height:	25 cm
Depth:	22 cm
Width:	11 cm
Weight:	7 kg dry

Agent capacity

Total capacity:	300 ml
Capacity between min/max fill marks:	170 ml
Wick system capacity:	75 ml approximately

Pneumatics specifications

Calibration and flow resistance

Calibration:	Calibration of all models is done at 21°C using O ₂ as carrier gas, 5 l/min fresh gas flow, at sea-level ambient pressure
Isoflurane models:	Control dial calibrated up to 5% V/V. The control dial is marked in steps of 0.25% up to 1% and in steps of 0.5% between 1% and 5%.
Sevoflurane models:	Control dial calibrated up to 8% V/V. Control dial is marked in steps of 0.25% up to 1% and in steps of 0.5% between 1% and 8%.
Flow resistance:	OFF: Isolated, no resistance from vaporizer ON: 5 l/min O ₂ : <20 cmH ₂ O

Accuracy

Isoflurane models:	±0.25% of delivered agent or ±15% of control dial setting (whichever is greater)
Sevoflurane models:	±0.4% of delivered agent or ±15% of control dial setting (whichever is greater)
Fresh gas flow:	200 ml/min to 15 l/min

Environmental specifications

Operation

Temperature:	15°C to 35°C
Humidity:	15% to 95% relative humidity (non-condensing)
Ambient pressure:	500-800 mmHg

Storage and Transport

Temperature:	-25°C to 60°C
Humidity:	15% to 95% relative humidity (non-condensing)
Ambient pressure:	400-800 mmHg

Service

Planned factory service free

MRI Safety Information

MR Conditional

Agent filling systems

Isoflurane models: Easy-Fil™

Sevoflurane models: Easy-Fil, Quik-Fil™, SAFE-T-SEAL™, Piramal Fill

Note: may not be available in all regions

Warranty

Warranted to be free from functional defects in materials and workmanship for a period of five years from the date of original delivery.



Agent color coded Easy-Fil adapters



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DOC2018947



Tec 850 Vaporizer

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Features

- Tec 850 agent specific vaporizers are designed to deliver isoflurane and sevoflurane anesthetic agents
- Provides output consistent with the dial setting throughout the clinical flow range
- Easy to turn dial and fine graduations help control agent delivery
- Enhanced design with consistent, reliable operation over the product life
- Tec 850 vaporizer supports a variety of contemporary agent filling mechanisms
- Improved ergonomics and contemporary design complements GE Healthcare's anesthesia systems
- Attaches to GE Healthcare interlocking Selectatec™ manifold
- No planned factory service needed
- Five-year warranty period



Clinical performance

- Tec 850 Vaporizers are designed to provide consistent output throughout the clinical flow range from 200 ml/min to 15 l/min. Models are available for isoflurane and sevoflurane
- Tec 850 Vaporizers are equipped with a large diameter control dial with graduations up to 5% V/V (isoflurane models) or 8% V/V (sevoflurane models)
- The easy turning dials and small graduations on the Tec 850 Vaporizer help you fine tune anesthetic delivery over the full range of dial settings and flow rates

Enhanced ergonomics

- The ergonomic dial release on the Tec 850 Vaporizer allows either left or right hand operation
- Wide, centered liquid level indicator provides a clear indication of the fill level of the vaporizer
- Easy to turn dial with enhanced readability

Lower overall ownership costs

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Reliability, convenience, and operating excellence

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- The non-spill system in the Tec 850 Vaporizer limits movement of liquid agent if the vaporizer is tilted or inverted, protecting internal components and helping maintain output within clinically acceptable limits



Easy-Fil helps simplify agent filling



Tec 850 agent specific vaporizers

Physical specifications

Dimensions

Height:	25 cm
Depth:	22 cm
Width:	11 cm
Weight:	7 kg dry

Agent capacity

Total capacity:	300 ml
Capacity between min/max fill marks:	170 ml
Wick system capacity:	75 ml approximately

Pneumatics specifications

Calibration and flow resistance

Calibration:	Calibration of all models is done at 21°C using O ₂ as carrier gas, 5 l/min fresh gas flow, at sea-level ambient pressure
Isoflurane models:	Control dial calibrated up to 5% V/V. The control dial is marked in steps of 0.25% up to 1% and in steps of 0.5% between 1% and 5%.
Sevoflurane models:	Control dial calibrated up to 8% V/V. Control dial is marked in steps of 0.25% up to 1% and in steps of 0.5% between 1% and 8%.
Flow resistance:	OFF: Isolated, no resistance from vaporizer ON: 5 l/min O ₂ : <20 cmH ₂ O

Accuracy

Isoflurane models:	±0.25% of delivered agent or ±15% of control dial setting (whichever is greater)
Sevoflurane models:	±0.4% of delivered agent or ±15% of control dial setting (whichever is greater)
Fresh gas flow:	200 ml/min to 15 l/min

Environmental specifications

Operation

Temperature:	15°C to 35°C
Humidity:	15% to 95% relative humidity (non-condensing)
Ambient pressure:	500-800 mmHg

Storage and Transport

Temperature:	-25°C to 60°C
Humidity:	15% to 95% relative humidity (non-condensing)
Ambient pressure:	400-800 mmHg

Service

Planned factory service free

MRI Safety Information

MR Conditional

Agent filling systems

Isoflurane models: Easy-Fil™

Sevoflurane models: Easy-Fil, Quik-Fil™, SAFE-T-SEAL™, Piramal Fill

Note: may not be available in all regions

Warranty

Warranted to be free from functional defects in materials and workmanship for a period of five years from the date of original delivery.



Agent color coded Easy-Fil adapters



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DOC2018947

Airway Gas Option, N-CAiO

Essential anesthesia monitoring for adult, pediatric and neonatal anesthesia applications.



With the B40 Patient Monitor, the airway gas module, N-CAiO, supports respiratory monitoring in the Operating Room.

Features

- Airway gases measured by the sidestream method
- Et and Fi values updated breath by breath
- Fast oxygen measurement for accurate EtO₂ and FiO₂ values
- Automatic identification of the anesthetic agent in use
- Light, compact size with low power consumption
- Variety of GE-validated gas sampling accessories for monitoring application-specific needs

Clinical measurements

- CO₂ and N₂O – GE infrared technology:
Inspired and end-tidal values, CO₂ waveform and respiration rate
- Respiration rate – calculated from the CO₂ waveform
- Anesthetic agents – GE infrared technology
 - Measures and identifies all five agents halothane, enflurane, isoflurane, sevoflurane and desflurane
 - MAC (Minimum Alveolar Concentration)
- Patient oxygen – GE paramagnetic oxygen (O₂) technology:
Inspired, end-tidal and Fi-Et difference, waveform



Technical specifications

General

When monitoring neonatal or other patients that have high respiration rate or low tidal volume these modules shall be used within the limits of respiration rates and tidal volumes to ensure specified measurement accuracy.

Sampling flow $120 \pm 20 \text{ ml/min}$

Size and fit of gas sampling accessories may impact measured gas concentration values at low tidal volumes. Always ensure use of appropriate accessories according to patient and application.

Automatic compensation for atmospheric pressure variation (660-1060 mbar), temperature and CO_2 , O_2 , N_2O , agent cross effect compensation. Parameter display update interval typically breath-by-breath.

Functional alarms for

- Low gas sampling flow
- Blocked sample line
- Blocked sample gas outflow
- Disconnected water trap
- Blocked water trap

Letters in the module name stand for

C = CO_2 and N_2O

Ai = Anesthetic agents with single agent identification

O = Patient O_2

Non-disturbing gases

- Ethanol, acetone, isopropanol, methane, nitrogen, nitric oxide, carbon monoxide, water vapor and freon R134A (for CO_2 , O_2 and N_2O).
- Maximum effect of non-disturbing gases on readings:
 $\text{O}_2 & \text{N}_2\text{O} < 2\text{ vol\%}$, $\text{CO}_2 < 0.2 \text{ vol\%}$, AA < 0.15 vol%.

Carbon dioxide (CO_2)

GE infrared absorption sensor technology

CO_2 waveform

EtCO_2 End-tidal CO_2 concentration

FiCO_2 Inspired CO_2 concentration

Measurement range 0 to 15 vol%
(0 to 15 kPa, 0 to 113 mmHg)

Accuracy $\pm(0.2 \text{ vol\%} + 2\% \text{ of reading})$

Rise time <260 ms

Adjustable low and high alarm limits for EtCO_2 or FiCO_2

Respiration rate (RR)

Measurement range 4 to 100 breaths/min

Detection criteria 1 vol% change in CO_2 level

Alarm note sent to host device if no breath detected in 20 seconds

Patient oxygen (O_2)

GE differential paramagnetic sensor

O_2 waveform

FiO_2 Inspired O_2 concentration

EtO_2 End-tidal O_2 concentration

$\text{FiO}_2\text{-EtO}_2$ Inspired-expired difference

Measurement range 0 to 100 vol%

Accuracy $\pm(1 \text{ vol\%} + 2\% \text{ of reading})$

Rise time <260 ms

Nitrous oxide (N_2O)

GE infrared absorption sensor

FiN_2O Inspired N_2O concentration

EtN_2O End-tidal N_2O concentration

Measurement range 0 to 100 vol%

Accuracy $\pm(2 \text{ vol\%} + 2\% \text{ of reading})$
 $\text{N}_2\text{O} \leq 85\%$

Anesthetic agent (AA)

GE infrared absorption sensor

Anesthetic agent waveform, if requested by host device

FiAA Inspired anesthetic agent concentration

EtAA End-tidal anesthetic agent concentration

MAC value options for hosts

Measurement range

Sevoflurane 0 to 8 vol%

Desflurane 0 to 20 vol%

Isoflurane, enflurane,
halothane 0 to 6 vol%

Accuracy $\pm(0.15 \text{ vol\%} + 5\% \text{ of reading})$

Agent identification

Identification threshold 0.15 vol%

Detection time <20 sec

System compatibility

- B40 Patient Monitor, (2060600-002)

Environmental specifications

Operating conditions

Temperature	10 to 40°C (50 to 104°F)
Relative humidity	10 to 98% non-condensing
Ambient pressure	660 to 1060 mbar

Storage conditions

Temperature	-25 to 60°C (-13 to 140°F)
Relative humidity	10 to 90% non-condensing

Physical specifications

Dimensions (H x W x D), excluding water trap	11.3 x 3.8 x 20.5 cm (4.4 x 1.5 x 8.1 in)
Weight	0.7 kg (1.5 lb)

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Always refer to the user manual that accompanies the monitor/module.

GE Healthcare Finland Oy, a General Electric company, doing business as GE Healthcare.

GE Healthcare, a division of General Electric Company

About GE Healthcare

GE Healthcare provides transformational medical technologies and services that are shaping a new age of patient care. Our broad expertise in medical imaging and information technologies, medical diagnostics, patient monitoring systems, drug discovery, biopharmaceutical manufacturing technologies, performance improvement and performance solutions services help our customers to deliver better care to more people around the world at a lower cost. In addition, we partner with healthcare leaders, striving to leverage the global policy change necessary to implement a successful shift to sustainable healthcare systems.

Our “healthymagination” vision for the future invites the world to join us on our journey as we continuously develop innovations focused on reducing costs, increasing access and improving quality around the world. Headquartered in the United Kingdom, GE Healthcare is a unit of General Electric Company (NYSE: GE). Worldwide, GE Healthcare employees are committed to serving healthcare professionals and their patients in more than 100 countries. For more information about GE Healthcare, visit our website at www.gehealthcare.com.

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imagination at work

DOC1379111 5/13

Neuromuscular Transmission Module, E-NMT

For integrated measurement of the level of neuromuscular block



The neuromuscular transmission module, E-NMT, uniquely integrates the measurement of the level of neuromuscular block into a single-width, plug-in module.

Features

- Provides a quantitative, automatic measurement of muscle response to an electrical stimulus
- Offers all common stimulation modes: train of four (TOF), single twitch (ST), double burst stimulation (DBS) and tetanic stimulation
- Covers the whole range of neuromuscular blockade
- Provides two measurement sensor options: the easy-to-apply MechanoSensor (KMG) for routine clinical use, and ElectroSensor (EMG) for research use

- Offers two MechanoSensor options: for adults and for pediatric patients
- Provides automatic setting of supramaximal current
- Provides automatic measurement at user-selectable interval
- Recovery Note can be set to notify when the block is wearing off
- Recall function enables moving the module with the patient without losing the reference values and the measured supramaximal current
- In addition to the integrated neuromuscular block measurement, the module enables nerve location for regional block by utilizing a comfortable 40 µs pulse



Technical specifications

Direct function keys		Regional Block mode	
Start-up	Automatically initiates the measurement by setting supramaximal current, reference value and starting cycle	Stimulation mode	Single twitch
Stop/Continue	Stops / continues measurement with same patient	Stimulation intervals	1, 2, 3 s

NMT

Stimulation modes	Train of four, TOF Double burst, DBS (3,3) Single twitch, ST 50 Hz tetanic & post tetanic count, PTC	Monitor compatibility
Numeric display	TOF%/DBS%, Count, T1%, PTC	CARESCAPE modular monitors with OR, PACU, ED and/or Critical Care software S/5 modular monitors using software L-(C)ANE03(A) or later versions

Measurement intervals for TOF/DBS

CARESCAPE™ modular monitors with OR and PACU software:
Manual, 10 s, 12 s, 15 s, 20 s, 1 min, 5 min or 15 min.

CARESCAPE modular monitors with Critical Care and ED software: Manual, 20 s, 1 min, 5 min, 15 min, 30 min, 60 min or 120 min.

S/5 modular monitors with anesthesia software: Manual, 10 s, 12 s, 15 s, 20 s, 1 min, 5 min or 15 min.

S/5 modular monitors with critical care software: Manual, 20 s, 1 min, 5 min, 15 min, 30 min, 60 min or 120 min.

Measurement intervals for ST

Manual, 1 s, 10 s, 20 s

Stimulus pulse	Square wave, constant current
Pulse width	100, 200 or 300 µs
Stimulus current range	Supramax 10 to 70 mA Manual 10 to 70 mA with 5 mA steps
Stimulus current accuracy	10% or ±3 mA, whichever is greater
Load for 70mA	3 kΩ (max)
Voltage	300 V (max)

Environmental specifications

Operating conditions

Temperature	10 to 40°C (50 to 104°F)
Relative humidity	10 to 90% non-condensing

Storage conditions

Temperature	-25 to 70°C (-13 to 158°F)
Relative humidity	10 to 90% non-condensing

Physical specifications

Dimensions (H x W x D)	11.2 x 3.7 x 18.6 cm (4.4 x 1.5 x 7.3 in)
Weight	0.35 kg (0.8 lb)

Warranty

One year

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GE Healthcare Finland Oy, a General Electric company, doing business as GE Healthcare.

GE Healthcare, a division of General Electric Company.

About GE Healthcare

GE Healthcare provides transformational medical technologies and services that are shaping a new age of patient care. Our broad expertise in medical imaging and information technologies, medical diagnostics, patient monitoring systems, drug discovery, biopharmaceutical manufacturing technologies, performance improvement and performance solutions services help our customers to deliver better care to more people around the world at a lower cost. In addition, we partner with healthcare leaders, striving to leverage the global policy change necessary to implement a successful shift to sustainable healthcare systems.

Our “healthymagination” vision for the future invites the world to join us on our journey as we continuously develop innovations focused on reducing costs, increasing access and improving quality and efficiency around the world.

GE Healthcare
P.O. Box 900, FIN-00031 GE, Finland
Tel. +358 10 394 11
Fax +358 9 146 3310

www.gehealthcare.com



GE imagination at work



EC DECLARATION OF CONFORMITY

(Following the provisions of the medical devices directive 93/42/EEC, Annex II
and of the directive 2011/65/EU)

We

Manufacturer
Datex-Ohmeda, Inc.
3030 Ohmeda Drive
PO Box 7550
Madison, WI 53707-7550 USA

EU Authorized Representative
GE Healthcare Finland Oy
Kuortaneenkatu 2
Helsinki, Finland
FI-00510

Manufacturing Site
GE Medical Systems (China) Co., Ltd.
19 Changjiang Road
Wuxi, Jiangsu, PR China 214028

Declare under our sole responsibility that the class IIb devices:

Carestation 650

Version: A1 REF: 1012-9650-000

Carestation 650c

Version: A1 REF: 1012-9655-000

Carestation 620

Version: A1 REF: 1012-9620-000

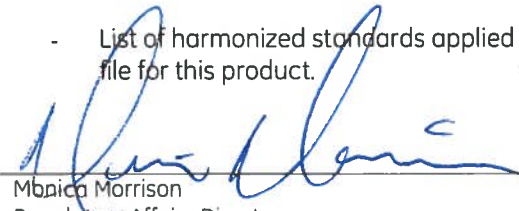
GMDN Code: 37710, UMDNS Code: 10-134

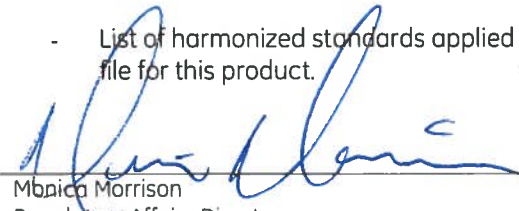
Classification rule (93/42/EC Annex IX): Class IIb, Rule 11

To which this declaration relates is in conformity with the requirements of the medical devices directive 93/42/EEC which apply to it. In addition, the product is in conformity with the requirements of the directive 2011/65/EU on the restriction of the use of certain hazardous substances in electrical and electronic equipment (as assessed by the manufacturer).

This conformity is based on the following elements:

- Information included in the documents:
Technical Documentation Ref: DOC1659791, of the product to which this declaration relates.
- EC certificate: approval of full quality assurance system (Annex II of the medical devices directive 93/42 EEC) delivered by TÜV Rheinland LGA Products GmbH, Tillystraße 2, 90431 Nuremberg, Germany, Notified Body # 0197, Certificate N° HD 60098566 0001 valid until 9 June 2016.
- List of harmonized standards applied for CE marking is located in the Technical Documentation file for this product.


Monica Morrison
Regulatory Affairs Director


16 JUN 2015
Madison, USA, Day Month -Year



EC DECLARATION OF CONFORMITY

(Following the provisions of the medical devices directive 93/42/EEC, Annex II
and of the directive 2011/65/EU)

We

Manufacturer
Datex-Ohmeda, Inc.
3030 Ohmeda Drive
PO Box 7550
Madison, WI 53707-7550 USA

EU Authorized Representative
GE Healthcare Finland Oy
Kuortaneenkatu 2
Helsinki, Finland
FI-00510

Manufacturing Site
GE Medical Systems (China) Co., Ltd.
19 Changjiang Road
Wuxi, Jiangsu, PR China 214028

Manufacturing Site
Datex-Ohmeda, Inc.
3030 Ohmeda Drive
PO Box 7550
Madison, WI 53707-7550 USA

Declare under our sole responsibility that the class IIb devices:

Carestation 650

Version: A1 REF: 1012-9650-000

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Carestation 620

Version: A1 REF: 1012-9620-000

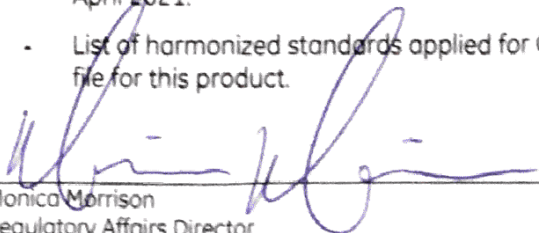
GMDN Code: 37710, UMDNS Code: 10-134

Classification rule (93/42/EC Annex IX): Class IIb, Rule 11

To which this declaration relates is in conformity with the requirements of the medical devices directive 93/42/EEC which apply to it. In addition, the product is in conformity with the requirements of the directive 2011/65/EU on the restriction of the use of certain hazardous substances in electrical and electronic equipment (as assessed by the manufacturer).

This conformity is based on the following elements:

- Information included in the documents:
Technical Documentation Ref: DOC1659791, of the product to which this declaration relates.
- EC certificate: approval of full quality assurance system (Annex II of the medical devices directive 93/42 EEC) delivered by TÜV Rheinland LGA Products GmbH, Tillystraße 2, 90431 Nuremberg, Germany, Notified Body # 0197, Certificate N° HD 60109676 0001 valid until 19 April 2021.
- List of harmonized standards applied for CE marking is located in the Technical Documentation file for this product.


Monica Morrison
Regulatory Affairs Director

6 MAY 2016
Madison, USA, Day Month -Year



EU DECLARATION OF CONFORMITY

Following the provisions of the medical devices regulation 2017/745,

ROHS directive 2011/65/EU and Radio Equipment Directive 2014/53/EU.

We:

Manufacturer	EU Authorized Representative
GE Medical Systems Information Technologies, Inc. 9900 Innovation Drive Wauwatosa, WI 53226, USA	GE Medical Systems SCS 283 rue de la Minière 78530 BUC, France

Manufacturing Site

GE Medical systems (China) Co., Ltd
No. 19, ChangJiang Road, WuXi National Hi-tech Development Zone
Jiangsu, P.R. China 214028

Declare under our sole responsibility that the device:

B125P/B105P/B125M/B105M/B155M Patient Monitor

Basic UDI-DI: 8406821BUG00102GM

Identification number:

B105P 6160000-001

B125P 6160000-002

B105M 6160000-003

B125M 6160000-004

B155M 6160000-005

SIGNATURE:

Monica Morrison

Date

Executive - Regulatory Affairs
Washington, DC USA

18 NOV 2020



Intended Purpose: Vital Signs Monitoring Instrument

GMDN Code and Description: 33586 Patient monitor, multiparameter

EMDN Code and Description: Z120302 Vital Signs Monitoring Instruments

Class: IIb

Classification rule (Annex VIII): Rule 10

To which this declaration relates is in conformity with the requirements of the medical devices regulation 2017/745 that apply to it and with the requirements of the RoHS directive 2011/65/EU on the restriction of the use of certain hazardous substances in electrical and electronic equipment.

This conformity is based on the following elements:

- Technical Documentation reference: DOC2430158, of the product to which this declaration relates.
- EC certificate No. HZ 2214580-1:
 - Conformity assessment procedure followed: Annex IX, Chapters I, III
 - Delivered by TÜV Rheinland LGA Products GmbH (0197)

We, manufacturer, declare under our sole responsibility that:

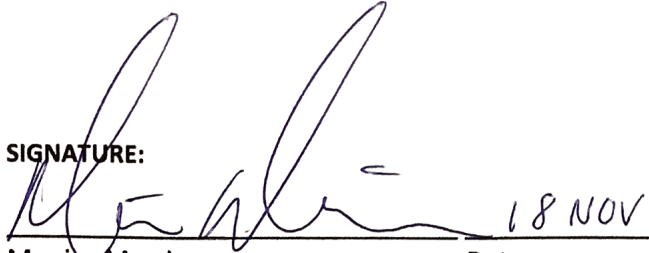
B125P/B105P/B125M/B105M/B155M Patient Monitor equipped with B1x5-01 WLAN module

To which this declaration relates is in conformity with the requirements of the Radio Equipment Directive 2014/53/EU which applies to it.

This conformity is based on the following elements:

- The device conforms to the Directive 2014/53/EU through Annex II-Internal Production Control
- The list of harmonized standards applied is located in the Technical File for this product.

SIGNATURE:



18 NOV 2020

Monica Morrison

Date

Executive - Regulatory Affairs
Washington, DC USA

EC Certificate

EU Quality Management System

REGULATION (EU) 2017/745 on Medical Devices, Annex IX Chapter I,
Section 2 and 3 and Chapter III



Registration No.: HZ 2214580-1

Manufacturer: GE Medical Systems Information Technologies, Inc.
9900 Innovation Drive
Wauwatosa, WI 53226
USA

EUDAMED Single Registration No.: N/A

Products: Class IIa- Z120503 ELECTROCARDIOGRAPHS
Class IIb -Z120302 VITAL SIGNS MONITORING INSTRUMENTS

Authorised representative(s): GE Medical Systems SCS
283 Rue de la Miniere, 78530 BUC
France

Certificate history		
Revision:	Description:	Issue date:
0	Initial	2020-11-17

The Notified Body hereby declares that the requirements of Annex IX, Chapter I, Section 2 and 3 of the REGULATION (EU) 2017/745 have been met for the listed products. The above named manufacturer has established and applies a quality management system, which is subject to periodic surveillance, defined by Annex IX, Chapter I, Section 3 of the aforementioned regulation. The requirements of Annex IX, Chapter III are fulfilled. If class III devices or class IIb implantable devices referred to in the second subparagraph of Article 52(4) are covered by this certificate an EU technical documentation assessment certificate according to Chapter II, Section 4.9 is required before placing them on the market.

Report No.: 234158038-30

Effective date: 2020-11-17

Expiry date: 2025-10-30

Issue date: 2020-11-17



Benannt durch/Designated by
Zentrale Stelle der Länder für Gesundheitsschutz
bei Arzneimitteln und Medizinprodukten
www.zsl.de
BS-MDR-091



TÜV Rheinland LGA Products GmbH
Tillystraße 2 · 90431 Nürnberg · Germany

TÜV Rheinland LGA Products GmbH is a Notified Body according to REGULATION (EU) 2017/745 concerning medical devices with the identification number 0197.

1 of 1

**DECLARATION OF CONFORMITY**

Following the provisions of the medical devices directive 93/42/EEC, Annex II
and of the directive 2011/65/EU

We

Manufacturer

Datex-Ohmeda, Inc.
3030 Ohmeda Drive
PO Box 7550
Madison, WI 53707-7550 USA

EU Authorized Representative

GE Medical Systems SCS
283 rue de la Minière
78530 BUC, France

Declare under our sole responsibility that the device:

Tec 820, Tec 850

Ref: 1177-9820-ISO, 1177-9820-SEV, 1177-9850-ISO, 1177-9850-SEV

GMDN Code: Isoflurane vaporizer - 36890; Sevoflurane vaporizer - 36980

GMDN Description: A device used to vaporize the anaesthetic agent isoflurane or sevoflurane and deliver a controlled amount of the agent to a patient being prepared for surgery. The device is usually mounted to an anaesthesia system or ventilator.

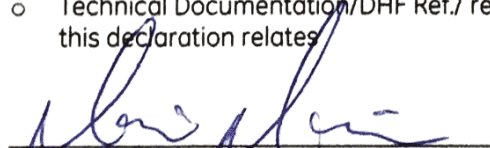
UDI-DI (GTIN) code: 1177-9820-ISO: 00840682124744; 1177-9820-SEV: 00840682124720;
1177-9850-ISO: 00840682124751; 1177-9850-SEV: 00840682124737

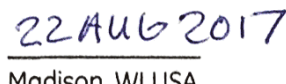
Classification rule (93/42/EEC Annex IX): 11 Class IIb

To which this declaration relates, is in conformity with the requirements of the medical devices directive 93/42/EEC which apply to it and with the requirements of the directive 2011/65/EU on the restriction of the use of certain hazardous substances in electrical and electronic equipment.

This conformity is based on the following elements:

- For the directive 93/42/EEC (MDD)
 - Technical Documentation/DHF Ref./ réf: DOC2019683, of the product to which this declaration relates
 - EC certificate: approval of full quality assurance system (Annex II of the directive 93/42 EEC) delivered by TÜV Rheinland LGA Products GmbH, Tillystraße 2, 90431, Neuremberg, Germany, Notified Body #0197, Certificate N° HD 60109676 0001, valid until 19 April 2021
 - harmonized standards applied on the product to which this declaration relates
- For the directive 2011/65/EU (RoHS)
 - Technical Documentation/DHF Ref./ réf: DOC1977840 and DOC1979675, of the product to which this declaration relates


Monica Morrison
Regulatory Affairs Director


22 Aug 2017
Madison, WI USA

This EC declaration of conformity is the first issue.

Tec 7 Vaporizer

Combining clinical performance
with ergonomic design

The Tec® 7 Vaporizer from GE Healthcare delivers isoflurane, sevoflurane, enflurane and halothane effectively and efficiently. Each agent specific Tec 7 incorporates significant performance, convenience and ergonomic benefits with important safety features to help improve agent delivery and reliability while reducing overall operating costs.



Features

- Tec 7 agent specific vaporizers are designed to deliver isoflurane, sevoflurane, enflurane and halothane anesthetic agents
- Provides output consistent with the dial setting throughout the clinical flow range
- Easy to turn dial and fine graduations help control agent delivery
- Three filling options: Easy-fil®, funnel fill, or Quik-Fil™ (sevoflurane only)
- Enhanced design helps improve reliability, operating excellence and product life
- Easy-fil filler is designed to simplify agent filling and help minimize agent leaks while filling. Tec 7 is also available in funnel fill and Quik-Fil (sevoflurane only) variants
- Improved ergonomics and contemporary design complements GE Healthcare's anesthesia systems
- Attaches to GE Healthcare interlocking Selectatec® manifold
- Planned factory service free
- Three year warranty period

Clinical performance

- Tec 7 Vaporizers are designed to provide consistent output throughout the clinical flow range from 200 mL/min to 15 L/min. Models are available for isoflurane, sevoflurane, enflurane and halothane.
- Tec 7 Vaporizers are equipped with a large diameter control dial which incorporates fine graduations of 0.2% between 0 and 1%, and 0.5% from 1% to 8%. The dial for sevoflurane is marked in steps of 0.2% up to 1% v/v, and in steps of 1% between 1% and 8%.
- The easy turning dials and small graduations on the Tec 7 Vaporizer help you fine tune anesthetic delivery over the full range of dial settings and flow rates.



Tec 7 agent specific vaporizers

Enhanced ergonomics

- The ergonomic agent dial release on the Tec 7 Vaporizer allows either left or right hand operation.
- A prismatic site glass on the front panel of the Tec 7 Vaporizer provides clear indication of liquid agent level.

Lower overall ownership costs

- The Tec 7 Vaporizer is planned factory service free, which helps lower ownership costs and eliminate the logistical challenges associated with the return of vaporizers to the factory. For complete user maintenance requirements, refer to the User Reference Manual.
- The Tec 7 Vaporizer comes with a three year warranty.



Easy-fil helps simplify agent filling

Reliability, convenience and operating excellence

- Easy-fil filler is designed to simplify agent filling and help minimize agent leaks while filling. The Tec 7 Vaporizer is also available with funnel fill and Quik-Fil (sevoflurane only) filling mechanisms.
- To minimize filling frequency, the Tec 7 Vaporizer accommodates 225 mL of anesthetic agent.
- The Tec 7 Vaporizer can be mounted on Datex-Ohmeda anesthesia systems equipped with the Selectatec manifold without tools and without taking the anesthesia system out of service.
- The Tec 7 Vaporizer also interlocks in series with Datex-Ohmeda Tec 4, Tec 5, Tec 6 and Tec 6 Plus Vaporizers.
- An innovative non-spill system in the Tec 7 Vaporizer limits movement of liquid agent if the vaporizer is tilted or inverted, protecting internal components and helping maintain output within clinically acceptable limits.

Physical specifications

Dimensions

Height:	25 cm/9.9 in
Depth:	21 cm/8.3 in
Width:	11.4 cm/4.5 in
Weight:	7 kg/15.4 lb dry

Agent capacity

Charged:	225 mL of free volatile anesthetic agent
Charging:	300 mL (nominal) to charge dry vaporizers
Retention:	75 mL (nominal) retained by wick system

Pneumatics specifications

Calibration and resistance

Calibration: The dials of the Tec 7 Vaporizers for enflurane, isoflurane and halothane are calibrated to 5% v/v in steps of 0.2% up to 1%, and in steps of 0.5% from 1% to 5% at 21°C using O₂ at 5 L/min as the carrier gas

Calibration of sevoflurane

sevoflurane: The dial of the Tec 7 Vaporizer for sevoflurane is calibrated to 8% v/v

Dial is marked in steps of 0.2% up to 1% v/v, and in steps of 1% between 1% and 8%

Resistance to gas flow

"OFF": Isolated, no resistance from vaporizer
"ON": 5 L/min O₂: 10 to 15 cm H₂O at 21°C ±2°C

Accuracy: 5 L/min O₂ @ 21°C ± 2°C:

Vaporizer 5% ±0.25% of delivered agent or ±15% of dial setting (whichever is greater)

Vaporizer 8% ±0.4% of delivered agent or ±15% of dial setting (whichever is greater)

Flowrate: Vaporizers are designed to provide consistent output throughout the clinical flow range from 200 mL/min to 15 L/min



Agent color coded Easy-fil bottle adapters

Environmental specifications

Operation

Temperature: 18° to 35°C/64° to 95°F

Humidity: 30% to 75% relative humidity (non-condensing)

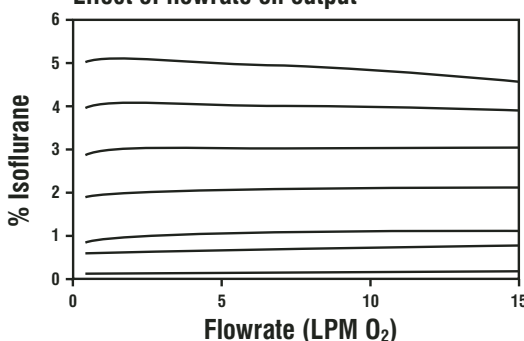
Service

Planned factory service free.

Warranty

Warranted to be free from functional defects in materials and workmanship for a period of three years from the date of original delivery.

Effect of flowrate on output



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Quik-Fil is a registered trademark of Abbott Laboratories
and is currently not available in the United States.

Easy-fil, Tec and Selectatec are trademarks of
Datex-Ohmeda, Inc.

Datex-Ohmeda, Inc., a General Electric company,
going to market as GE Healthcare.

For more than 100 years, healthcare providers
worldwide have relied on GE Healthcare for medical
technology, services, and productivity solutions. So
no matter what challenges your healthcare system
faces, you can always count on GE to help you
deliver the highest quality healthcare. For details,
please contact your GE representative today.

GE Healthcare
P.O. Box 7550
Madison, WI 53707-7550
USA

www.gehealthcare.com



imagination at work



Neuromuscular Transmission

What is neuromuscular transmission?

Neuromuscular transmission (NMT) is the transfer of an impulse between a nerve and a muscle in the neuromuscular junction. NMT can be blocked by neuromuscular blocking agents – drugs which cause transient muscle relaxation and prevent the patient from moving and breathing spontaneously.

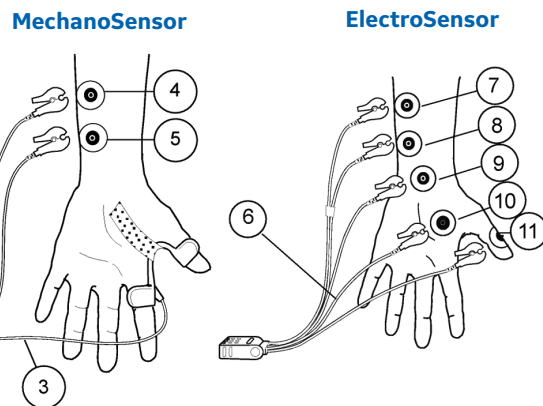
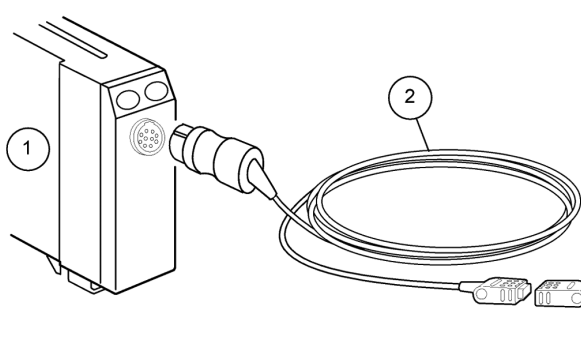
Muscle relaxation is used during general anesthesia to enable endotracheal intubation and to provide the surgeon with optimal working conditions. In critical care, muscle relaxation is used during mechanical ventilation to minimize the patient's work of breathing and to improve oxygenation.

How is the NMT block measured?

The level of neuromuscular block is routinely measured by stimulating a peripheral nerve, usually in the hand and by subjectively evaluating the muscle response either visually or tactiley with a device. Evidence suggests that despite use of subjective neuromuscular monitoring techniques, a large percentage of patients arrive to the PACU with residual paralysis.¹

By comparison, GEHC offers the NMT module, which provides quantitative, automatic measurements of muscle response to stimulus and consequentially, the level of block. This objective measurement can be used to safely time extubation and avoid the occurrence of residual paralysis.

GE Healthcare offers two different sensor types for the NMT module. The innovative MechanoSensor measures the motion of the thumb with a piezoelectric sensor, which converts the physical motion to an electrical signal and quantifies the evoked mechanical response. The MechanoSensor sensor is available in adult and pediatric sizes. The ElectroSensor directly measures the electrical activity of the muscle with recording electrodes, quantifying the response to nerve stimulation. The ElectroSensor can be used on the patient's hand or foot in both adult and pediatric patients.



NMT measurement setup with MechanoSensor based on kinemyography (KMG) for routine clinical NMT monitoring. Use a narrow tape to secure MechanoSensor securely on the patient's hand. Traditional electromyography (EMG) measurement with ElectroSensor.

- | | |
|--|--|
| 1. Module with NMT measurement capability | 7. White stimulating electrode |
| 2. NMT sensor cable | 8. Brown stimulating electrode |
| 3. MechanoSensor or Pediatric MechanoSensor lead wire set | 9. Electrode, black lead connection site, ground |
| 4. Electrode, white lead connection site for nerve stimulation | 10. Electrode, green lead connection site, recording muscle-contraction effect |
| 5. Electrode, brown lead connection site for nerve stimulation | 11. Electrode, red lead connection site, recording muscle-contraction effect |
| 6. ElectroSensor leadwire set | |

¹ Murphy GS, Brull SJ. Residual neuromuscular block: Lessons unlearned. Part 1: Definitions, incidence, adverse psychological effects of residual neuromuscular block. Anesth Analg 2010;111:120-128

Starting nerve stimulus

Start the NMT measurement by defining a patient specific reference with a supramaximal stimulus current for the unrelaxed patient before administering neuromuscular blocking agents. This will ensure reliable measurements are achieved during all levels of neuromuscular blockade. Thereafter module automatically maintains the optimum current throughout the procedure.

For reliable NMT measurement use GE validated NMT electrodes to provide large conductive surface.

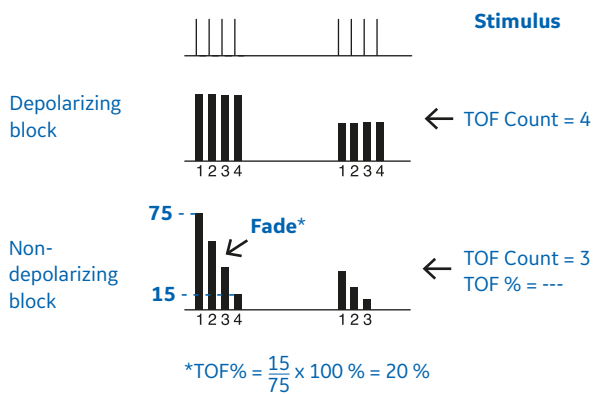
Train-of-four (TOF) is used as the default stimulation mode with four stimuli at 0.5 second intervals causing muscles to contract.

Quantitative muscle response

The muscle response can be quantified with different parameters depending on the type and the level of neuromuscular block.

TOF Count is the number of detected muscle responses. **Train-of-four ratio (TOF%)** is the ratio of the fourth muscle response to the first one. TOF% indicates fade in non-depolarizing block. Once the TOF Count drops below four responses or T1% is less than 10%, the TOF% is not shown.

When depolarizing agents are used, no fade occurs, and the height of the four responses indicates the level of block.

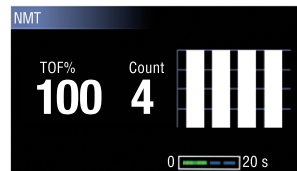


When no responses are detected to TOF stimulation, the **post tetanic count** (PTC) is the only way to measure the neuromuscular block. A tetanic stimulation (50 Hz) is generated for five seconds and post-tetanic responses to single twitch stimulation are counted. The larger the PTC, i.e., the number of detected responses, the sooner the normal TOF responses return. PTC is rarely used in critical care.

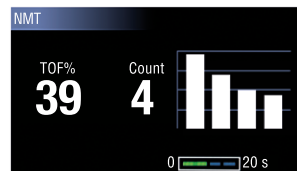
Light		Medium		Deep	
100	TOF%	20	4	Count	0
10	PTC	0	10	PTC	0

Relaxometer illustrates the level of neuromuscular blockade

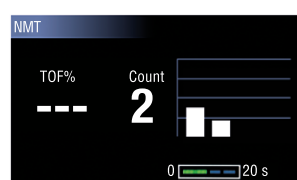
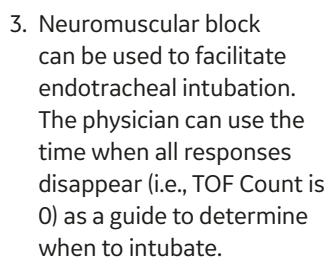
Monitoring of neuromuscular block in five steps



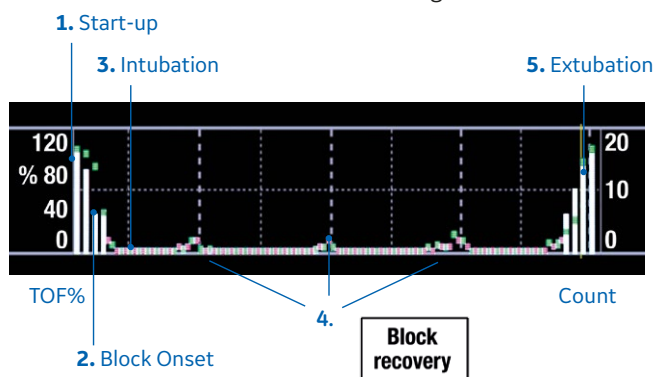
1. Properly secure the sensor of choice (as depicted on page 1). Press measurement start-up. The monitor will start the measurement by setting the stimulus current automatically and by performing a reference measurement. Depolarizing relaxants result in an equal drop in all four responses, without fade.



2. Non-depolarizing relaxants cause a fade in the responses, indicated by a lower TOF% and a slope in the bar graph.



3. Neuromuscular block can be used to facilitate endotracheal intubation. The physician can use the time when all responses disappear (i.e., TOF Count is 0) as a guide to determine when to intubate.
4. During surgery and in critical care, TOF Count is used to maintain steady optimal level of neuromuscular block. When TOF Count exceeds a level set by the user, the GE monitor will give a "Block recovery" message.



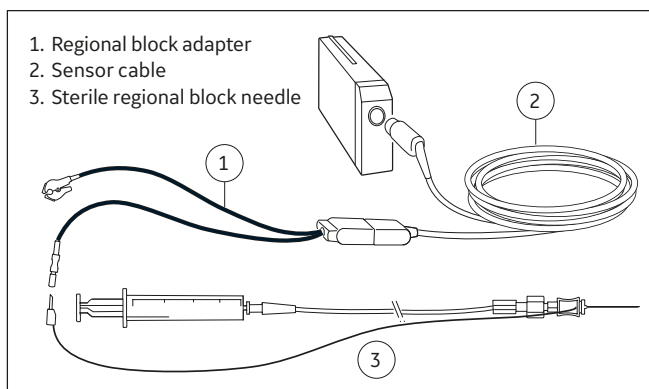
5. Antagonists, if used, should not be given before TOF count rises to 4. For safer extubation, TOF% should be higher than 90.

Nerve location for regional block

The NMT module can also be used to locate the nerve to be blocked in regional anesthesia.

A regional block needle is stimulated with small, repeated 2.0 mA stimuli while the nerve is approached. Each nerve stimulus should result in muscle contraction. The closer the needle is to the motor nerve, the lower the current is needed to give a response. When even a small stimulus current (e.g., < 0.5mA) results in a visible muscle contraction, the optimum site has been located and local anesthetic can be injected.

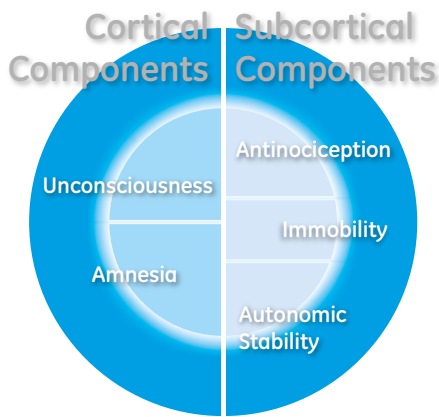
This method helps clinicians find the optimum site of the regional nerve to be blocked and, thus, optimizes the bolus of the anesthetic to be injected. The correct location also protects the patient against mechanical nerve and vessel lesions.



Adequacy of Anesthesia

Adequacy of Anesthesia consists of several interrelated components as depicted below.

One of the objectives of general anesthesia is **immobility**, i.e., the assurance that the patient does not move. Often neuromuscular blocking agents are used to achieve this goal. Anesthetic agents prolong and amplify the effects of neuromuscular blocking agents, therefore, it is recommended to use quantitative NMT monitoring when anesthetic agents are used together with neuromuscular blockade. NMT is an important part of adequacy of anesthesia monitoring and, when combined with other parameters such as the Entropy and hemodynamic measurements, it helps to achieve a more complete picture of the patient's status.



Why use the NMT module?

Automatic and hands-free

Neuromuscular block measurements with the NMT module are easy. Simply attach the sensor and push the Start-up key.

The module will set the supramaximal current and automatically cycle according to the user-defined measurement interval.

Optimal dosage during anesthesia and in critical care

Quantitative NMT monitoring gives a clear picture of the individual dosage needs of the patient and facilitates optimal and cost-effective administration of neuromuscular blocking drugs.

Optimized recovery

Monitoring the level of neuromuscular block enables follow-up and prediction of recovery and helps in correct timing of the antagonists, which may decrease the incidence of residual paralysis.

Enhanced patient safety

Objective, quantitative monitoring is the only means to safely indicate recovery of neuromuscular block (TOF%>90%).^{1,2} Therefore, utilizing the NMT module can help decrease the incidence of residual paralysis and the associated respiratory complications.

Fast patient throughput

Patients that arrive to the PACU with residual paralysis (TOF%<90) stay on average 90 minutes longer.³ Using NMT to guide extubation times can support shorter length of stay.

Integrated information

When the NMT measurement is integrated in a monitoring system, the measured values are displayed, trended and automatically documented together with all the other monitored parameters.

Additional resources

For white papers, guides and other instructive materials about our clinical measurements, technologies and applications, please visit <http://clinicalview.gehealthcare.com/>

¹ ATOTW 290 – Residual Neuromuscular Blockade, 26/08/2013

² Lars I. Eriksson, M.D., Ph.D., Evidence-based Practice and Neuromuscular Monitoring: It's Time for Routine Quantitative Assessment. Anesthesiology 5 2003, Vol.98, 1037-1039.

³ Butterly A, Bitner EA, George E, Sandberg WS, Eikermann M, Schmidt U. Postoperative residual curarization from intermediate-acting neuromuscular blocking agents delays recovery room discharge. Br J Anaesth. 2010; 105: 304-309.



Imagination at work

Product may not be available in all countries and regions. Full product technical specification is available upon request. Contact a GE Healthcare Representative for more information. Please visit www.gehealthcare.com/promotional-locations.

Data subject to change.

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