

EC Certificate - Full Quality Assurance System

Directive 93/42/EEC on Medical Devices, Annex II excluding Section 4

No. **CE 683722**
Issued To: **Breas Medical AB**
Företagsvägen 1
Mölnlycke
SE-435 33
Sweden

In respect of:

Design and Manufacture of Respiratory Therapy Systems, Respiratory Monitoring Devices, Sleep Apnea and Humidifier Systems and associated Stand Alone PC-Device Communication Software.

on the basis of our examination of the quality assurance system under the requirements of Council Directive 93/42/EEC, Annex II excluding section 4. The quality assurance system meets the requirements of the directive. For the placing on the market of class III products an Annex II section 4 certificate is required.

For and on behalf of BSI, a Notified Body for the above Directive (Notified Body Number 2797):



Gary E Slack, Senior Vice President - Medical Devices

First Issued: **2018-03-15**

Date: **2021-05-21**

Expiry Date: **2024-05-26**

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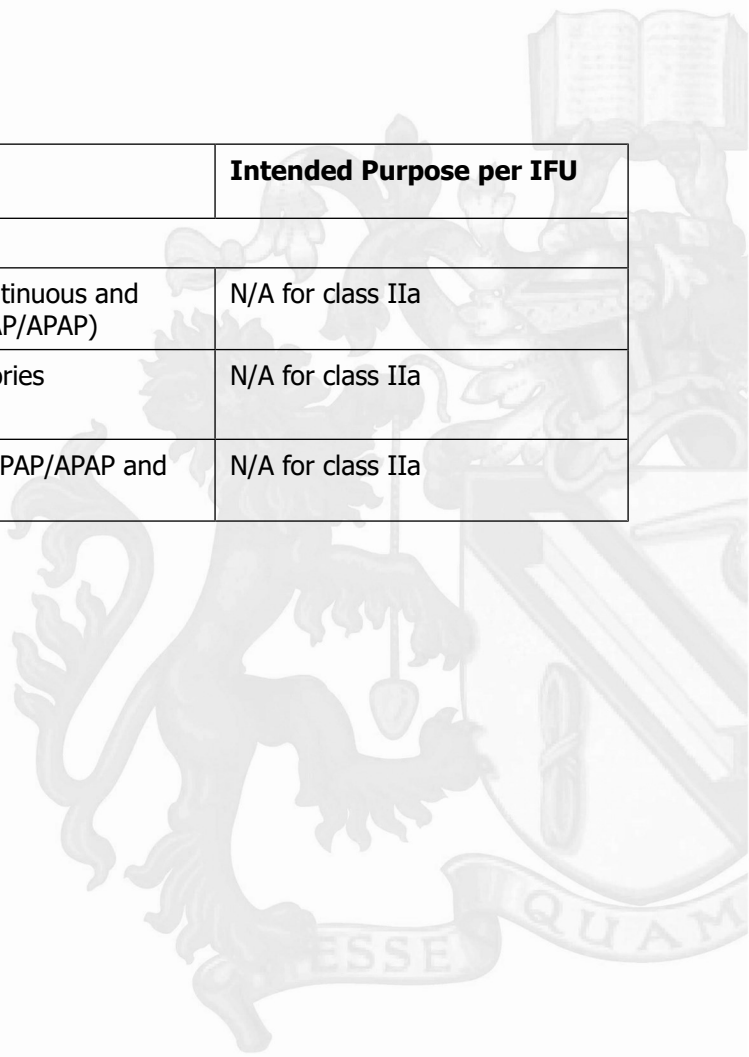
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Supplementary Information to CE 683722

Issued To: **Breas Medical AB**
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NBOG code(s)	Device Name	Intended Purpose per IFU
Class IIa		
MD 1102	Z1, Respiratory Therapy Devices for continuous and automatic positive airway pressure (CPAP/APAP)	N/A for class IIa
MD 0101 MD 1102	Breathing circuits and ventilator accessories	N/A for class IIa
MD 1111	Z1 Mobile App intended to control the CPAP/APAP and monitor sleep quality.	N/A for class IIa



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NBOG code(s)	Device Name	Intended Purpose per IFU
Class IIb		
MD 1102	Vivo 55	<p>The Vivo 55 (with or without the SpO₂ and CO₂ sensor) is intended to provide continuous or intermittent ventilatory support for the care of individuals who require mechanical ventilation. Specifically, the ventilator is applicable for paediatric through adult patients weighing more than 10kg.</p> <p>The Vivo 55 with the SpO₂ sensor is intended to measure functional oxygen saturation of arterial hemoglobin (%SpO₂) and pulse rate. The Vivo55 with the CO₂ sensor is intended to measure CO₂ in the inspiratory and expiratory gas.</p> <p>The device is intended to be used in home, institution, hospitals and portable applications such as wheelchairs and gurneys. It may be used for both invasive and non-invasive ventilation.</p>
MD 1102	Vivo 65	<p>The Vivo 65 (with or without the SpO₂ and CO₂ sensor) is intended to provide continuous or intermittent ventilatory support for the care of individuals who require mechanical ventilation. Specifically, the ventilator is applicable for paediatric through adult patients weighing more than 5 kg.</p> <p>The Vivo 65 with the SpO₂ sensor is intended to measure functional oxygen saturation of arterial haemoglobin (%SpO₂) and pulse rate. The Vivo 65 with the CO₂ sensor is intended to measure CO₂ in the inspiratory and expiratory gas.</p> <p>The device is intended to be used in home, institution, hospitals and portable applications such as wheelchairs and gurneys. It may be used for both invasive and non-invasive ventilation.</p>

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Information and Contact: BSI, Say Building, John M. Keynesplein 9, 1066 EP Amsterdam, The Netherlands Tel: + 31 20 346 0780

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NBOG code(s)	Device Name	Intended Purpose per IFU
Class IIb		
MD 1102	Vivo 45, Nippy 4	Vivo 45/Nippy 4 is intended to provide non-invasive or invasive ventilation for adult or paediatric patients weighing over 10 kg (22 lbs) who require long-term support or mechanical ventilation for respiratory insufficiency or respiratory failure, with or without obstructive sleep apnoea. Vivo 45/Nippy 4 is intended for spontaneously breathing patients.
MD 1102	Vivo 45LS, Nippy 4+	The Vivo 45 LS/Nippy 4+ ventilators (with or without the SpO2 and CO2 sensor) is intended to provide continuous or intermittent ventilatory support for the care of individuals who require mechanical ventilation. Specifically, the Vivo 45LS ventilator is applicable for paediatric through adult patients weighing more than 5 kg (11 lbs.) and the Nippy 4+ is applicable for paediatric through adult patients weighing more than 10 kg (22 lbs.) The Vivo 45LS/Nippy 4+ with the SpO2 is intended to measure functional oxygen saturation of arterial haemoglobin (%SpO2) and pulse rate. The Vivo 45LS/Nippy 4+ with the CO2 sensor is intended to measure CO2 in the inspiratory and expiratory gas. The device is intended to be used in home, institution, hospitals and portable applications such as wheelchairs and gurneys. It may be used for both invasive and non-invasive ventilation. The Vivo 45LS/Nippy 4+ are not intended to be used as an emergency transport or critical care ventilator.

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NBOG code(s)	Device Name	Intended Purpose per IFU
Class IIb		
MD 1102	Vivo 1, Vivo 2	Vivo 1 and 2 are intended to provide non-invasive ventilation for patients weighing over 10 kg (22 lbs) who require long-term support or mechanical ventilation for respiratory impairment, with or without obstructive sleep apnoea. Vivo 1 and 2 are intended for spontaneously breathing patients.
MD 1102	Vivo 3	Vivo 3 is intended to provide non-invasive or invasive ventilation for patients weighing over 10 kg (22 lbs) who require long-term support or mechanical ventilation for respiratory insufficiency or respiratory failure, with or without obstructive sleep apnoea. Vivo 3 is intended for spontaneously breathing patients.
MD 1111	Vivo 50/55/60/65 PC Software	The PC Software is intended to be used for follow-up on patient's ventilator treatment. The Software may indicate possible events that could require further clinical investigation. The Vivo PC Software is intended to be used in institutions, hospitals and clinics by trained clinical personnel, physicians, home care and service personnel. The Vivo PC Software can also be used for Remote Monitoring of ongoing treatment.

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NBOG code(s)	Device Name	Intended Purpose per IFU
Class IIb		
MD 1111	Vivo 30/40 PC Software	The Vivo 30/40 PC software setting values is designed to communicate with a Vivo 30 or Vivo 40 medical device, extracting data, or monitoring the device in real time. The PC software can also download device log files via cable or a CF card and manipulate patient information in patient archives.
MD 1111	Vivo 1/2/3/45/45LS/Nippy 4/4+ PC Software	The PC Software is intended to be used for follow-up on patient's ventilator treatment. The Software may indicate possible events that could require further clinical investigation. The Vivo PC Software is intended to be used in institutions, hospitals and clinics by trained clinical personnel, physicians, home care and service personnel. The Vivo PC Software can also be used for Remote Monitoring of ongoing treatment.

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Date	Reference Number	Action
15 March 2018	8855397	First issue. Transfer from another Notified Body.
12 February 2019	8900397	Traceable to NB 0086.
24 March 2020	9754972	Renewal. Change of name and address of additional site from Human Design Medical, LLC in Newton, Massachusetts to Breas Medical, Inc. North Billerica, MA in June 2018. Addition of subcontractors: Inission Boras AB Gränsvägen 6 518 40 Sjömarken, Sweden, GlobalMed Inc. 155 North Murray Street, Trenton, Ontario, K8V 5R5 Canada, Guangzhou Schauenburg-truplast Hose Technology Ltd, No. 6, Nanjiang 3rd Road, Nansha District, Guangzhou, Guangdong, 511462, P.R. China, Productos Urológos de Mexico S.A.de C.V. Cerrada Via de la Produccion No. 85 Parque Industrial Mexicali III Baja California CP 21397 Mexico, I3tex AB, Klippan 1 A, 414 51 Göteborg Sweden, Plastiflex Group NV, Beverlosesteenweg 99, 3583 Paal-Beringen, Belgium, NOTE TORSBY AB, Inova Park, 685 29 Torsby, Sweden and Breas Medical Ltd, Units A1-A2, The Bridge Business Centre, Timothy's Bridge Road, Stratford Enterprise Park, Stratford-upon-Avon, CV37 9HW. Addition of product table.

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Date	Reference Number	Action
05 April 2021	3414247	Removal of subcontractor: "Productos Urológicos de Mexico S.A.de C.V, Cerrada Via de la Produccion No. 85, Parque Industrial Mexicali III, Baja California, CP 21397, Mexico"
21 May 2021	3446719	Re-issuing of certificate. Changes in device table. Amended device table, Class IIb devices analytically listed and intended purpose amended per device/device groups and Class IIa device names provided. Withdraw of Class IIb humidifiers as they were an accessory to withdrawn devices Vivo 30/40 devices.
Non-significant changes approved after the 26th May 2021 as per the Transitional Provisions of MDR Article 120.3		
27 May 2022	3696967	Removal of subcontractors i3TEX AB and Inission Boras AB

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Date	Reference Number	Action
23 March 2026	30634028	Removal of "Sleep Apnea" from the certificate scope. Removal of the following devices from the Class IIa device table: Z1, Respiratory Therapy Devices for continuous and automatic positive airway pressure (CPAP/APAP). Z1 Mobile App intended to control the CPAP/APAP and monitor sleep quality. Removal of subcontractor page.

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23 March 2026

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To whom it may concern,

The transitional provisions specified in MDR Article 120(3) (as amended by (EU) 2023/607) prohibit Notified Bodies from issuing new certificates or amending, modifying, supplementing any existing MDD/AIMDD certificates from 26th May 2021.

The transitional provisions specified in IVDR Article 110(3) (as amended by (EU) 2024/1860) prohibit Notified Bodies from issuing new certificates or amending, modifying, supplementing any existing IVDD certificates from 26th May 2022.

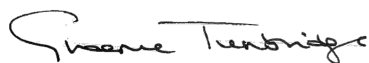
This letter is to confirm that BSI has reviewed and approved the change(s) detailed in the table below. These changes do not represent a significant change in design or intended purpose under MDR Article 120(3) or under IVDR Article 110(3), as applicable, and as per the guidance provided in MDCG 2020-3/MDCG 2022-6.

The related certificate specified below continues to remain valid and devices can be placed on the market based on this certificate as long as the manufacturer complies with the conditions specified in Section 3c of Article 120 of MDR or in Section 3c of Article 110 of IVDR, as applicable.

Certificate	Directive and Annex	Reference Number	Changes approved
CE 683722	93/42/EEC Annex II excluding Section 4	30634028	Removal of "Sleep Apnea" from the certificate scope. Removal of the following devices from the Class IIa device table: Z1, Respiratory Therapy Devices for continuous and automatic positive airway pressure (CPAP/APAP). Z1 Mobile App intended to control the CPAP/APAP and monitor sleep quality. Removal of subcontractor page.

Should you have any queries concerning your certification, or if we can be of further assistance to you, please contact your BSI Scheme Manager.

Yours sincerely,



Graeme Tunbridge
Senior Vice President, Medical Devices