### TEHNOMEDICA

str.Ciuflea, 38/1 MD-2001, mun. Chişinău, Moldova tel./fax: (022)601 102, 601 087 e-mail <<u>tehnomedica\_md@yahoo.com</u>> <<u>tehnomedicamd@gmail.com</u>>

> Anexa nr. 7 la Documentația standard nr.115 din 15.09.2021

### **CERERE DE PARTICIPARE**

### Către IMSP Spitalul Clinic Republican "Timofei Moșneaga"

### Stimați domni,

Ca urmare a anunțului/invitației de participare/de preselecție apărut în Buletinul achizițiilor publice și/sau Jurnalul Oficial al Uniunii Europene, nr. <u>ocds-b3wdp1-</u><u>MD-1648195071000, ID:21053907 din 15.04.2022</u> privind aplicarea procedurii pentru atribuirea contractului privind <u>achiziționarea consumabilelor medicale</u> <u>pentru anul 2022 Repetat</u>, noi, <u>Tehnomedica SRL</u>, am luat cunoștință de condițiile și de cerințele expuse în documentația de atribuire și exprimăm prin prezenta interesul de a participa, în calitate de ofertant/candidat, neavînd obiecții la documentația de atribuire.

Data completării: 14.04.2022 Cu stimă, Tehnomedica SRL Director Tatiana Roibu (semnătura autorizată)

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> Anexa nr. 8 la Documentația standard nr.115 din 15.09.2021

### DECLARAȚIE privind valabilitatea ofertei

### Către IMSP Spitalul Clinic Republican "Timofei Moșneaga"

### Stimați domni,

Ne angajăm să menținem oferta valabilă, privind <u>achiziționarea consumabilelor</u> <u>medicale pentru anul 2022 REPETAT</u> prin procedura de achiziție licitație deschisă, pentru o durată de 90 zile, (nouăzeci zile), începând cu data de 15.04.2022, respectiv până la data de 15.07.2022 (ziua/luna/anul), și ea va rămâne obligatorie pentru noi și poate fi acceptată oricând înainte de expirarea perioadei de valabilitate.

Data completării: 14.04.2022

Cu stimă,

Tehnomedica SRL

Director Tatiana Roibu

(semnătura autorizată)



## CERTIFICAT DE ÍNREGISTRARE

### SOCIETATEA CU RĂSPUNDERE LIMITATĂ "TEHNOMEDICA" ESTE ÎNREGISTRATĂ LA CAMERA ÎNREGISTRĂRII DE STAT

### Numărul de indentificare de stat - codul fiscal 1002600053256

Data înregistrării

17.04.2002

MOLDOVA

Data eliberării

16.02.2005

Bolboceanu Adela, registrator de stat Funcția, numele, prenumele persoanei care a eliberat certificatul





MD 0027040

### **BARRIER**<sup>®</sup>

### SURGICAL CAP ANNIE



Ref. no	Description	RET/TRP
621500	Surgical Cap L (elastic), mixed colours	150/600
621715	Surgical Cap M (elastic), blue	150/600
621725	Surgical Cap M (elastic), green	150/600
621735	Surgical Cap M (elastic), white	150/600
621815	Surgical Cap L (elastic), blue	150/600
621825	Surgical Cap L (elastic), green	150/600
621835	Surgical Cap L (elastic), white	150/600
621925	Surgical Cap XL (elastic), green	150/600

#### SURGICAL CAP CHARLOTTE



Ref. no	Description	RET/TRP
620900	Surgical Cap M (elastic), white	-/200

#### SURGICAL CAP EUROCAP



Ref. no	Description	RET/TRP
620800	Surgical Cap one-size (elastic), blue	100/500
620810	Surgical Cap one-size (elastic), green	100/500

### **BARRIER**<sup>®</sup>

Headwear / Basic

### SURGICAL CAP KOSACK

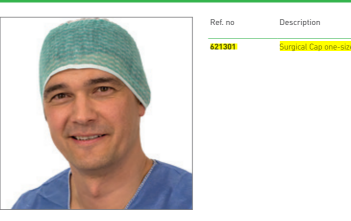


Ref. no	Description	RET/TRP
621000	Surgical Cap one-size (elastic), green	100/500
621001	Surgical Cap one-size (elastic), purple	100/500
621005	Surgical Cap one-size (elastic), blue	100/500

### SURGICAL CAP JACK



#### SURGICAL CAP PHILIP



#### SURGICAL CAP FLORY



#### Headwear / Standard

	RET/TRP
ize (elastic), green	100/500

	RET/TRP
re (tie-band), green	100/500

	RET/TRP
ize (tie-band), green	100/500

### **BARRIER**<sup>®</sup>

### Headwear / Special / Extra comfort

#### SURGICAL HOOD GLENN PRO



Ref. no	Description	RET/TRP
620205	Surgical Hood with collar one-size (sweatband, tie-band), blue	35/175

#### SURGICAL HOOD ALBIN



Ref. no	Description	RET/TRP
620250	Surgical Hood with collar one-size (sweatband, tie-band), green	60/300

#### HEADWEAR / EXTRA COMFORT / SURGICAL CAP MISS

LATEX

LATEX



Ref. no	Description	RET/TRP
42022	Surgical Cap one-size (elastic), latex, blue	120/960
42023	Surgical Cap one-size (elastic), latex, green	120/960





Ref. no	Description	RET/TRP
42093	Surgical Cap one-size (elastic), latex, green	120/960

### SURGICAL HOOD ALL

**BARRIER®** 



SURGICAL HOOD ALL



Description

Surgical Hood one



### Ref. no 42074



#### SURGICAL HOOD ALL PRO



Ref. no Description 42076 Surgical Hood with co



114

#### Headwear / Extra comfort

	RET/TRP
size (tie-band), green	120/960

	RET/TRP
-size (sweatband, tie-band), green	70/560

ollar	one-size	(sweatband,	tie-band),	green

RET/TRP

50/400

### **BARRIER®**

### Medical face masks / Standard (Type II)

### **BARRIER**<sup>®</sup>

STANDARD (TYPE II)

SPECIAL (TYPE II)

Ref. no

42280

Ref. no

4231

Ref. no	Description	RET/TRP
4230	Medical face mask, blue	60/600



Ref. no	Description	RET/TRP
302	Medical face mask, green	50/600
JEC		
TIE-BAND		



Ref. no	Description	RET/TRP
4312	Medical face mask, blue	50/600
TIE-BAND		



Ref. no	Description	RET/TRI
657000	Medical face mask, blue	60/360
J		







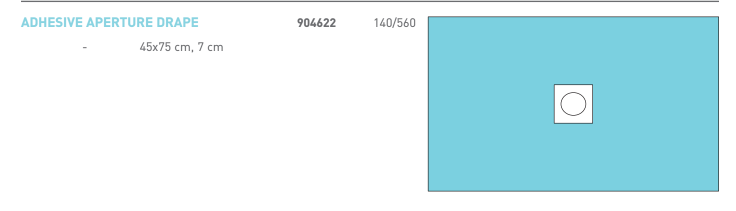


TIE-BAND

### Medical face masks / Standard (Type II) / Special (Type II)

Description	RET/TRP
Medical face mask, blue	60/600
Description	RET/TRP
ledical face mask, blue	60/600
50	
ANTI-F06	
ANTL-FOG	
ANTI-F06	
	RET/TRP
escription	RET/TRP 60/600
escription	
escription	
escription	
lescription	
escription	
escription	
escription	
ledical face mask, green	
ledical face mask, green	
escription fedical face mask, green	60/600
Description Aedical face mask, green	

#### **APERTURE DRAPES**



ADHESIVE APERTURE DRAPE	904624	80/320	
- 75x90 cm, 7 cm			
			$\bigcirc$

ADHESIVE APERTUR	E DRAPE	906540	50/300		
904741 6	0x75 cm, 6x9.5 cm				
				$\bigcirc$	

ADHESIVE APERTURE DRAPE	<mark>906542</mark>	40/160	
904758 <mark>75x90 cm, 6x8 cm</mark>			





# Certificate of Registration

### QUALITY MANAGEMENT SYSTEM - ISO 13485:2016 & EN ISO 13485:2016

This is to certify that:

Mölnlycke Health Care AB Gamlestadsvägen 3C Box 13080 SE-402 52 Göteborg Sweden

Holds Certificate Number:

MD 83345

and operates a Quality Management System which complies with the requirements of ISO 13485:2016 & EN ISO 13485:2016 for the following scope:

The design, development, manufacture, marketing, sales and distribution of sterile wound and scar dressings, open wound products, cavity dressings, polyurethane foam with and without additives for incorporation into medical devices, swabs, sponges, sterile alcohol wipes, skin care products, non-sterile textile bandages and support, sterile wound irrigation solutions, operation sets, surgical and equipment drapes, procedure packs, surgical gowns and medical staff clothing for use in the patient environment, sterile and non-sterile medical gloves and sterile surgical gloves, self-warming blankets, turning and positioning devices. The design, development, manufacture, marketing, sales and distribution of single patient use Negative Pressure wound therapy pumps and accessories. Distribution of laparoscopic instruments.

jang Conada

For and on behalf of BSI:

Gary E Slack, Senior Vice President - Medical Devices

Original Registration Date: 2004-07-21 Latest Revision Date: 2021-11-24 Effective Date: 2021-11-28 Expiry Date: 2024-11-27

Page: 1 of 1



...making excellence a habit."

This certificate was issued electronically and remains the property of BSI and is bound by the conditions of contract. An electronic certificate can be authenticated <u>online</u>. Printed copies can be validated at www.bsigroup.com/ClientDirectory

Information and Contact: BSI, Kitemark Court, Davy Avenue, Knowlhill, Milton Keynes MK5 8PP. Tel: + 44 345 080 9000 BSI Assurance UK Limited, registered in England under number 7805321 at 389 Chiswick High Road, London W4 4AL, UK. A Member of the BSI Group of Companies.





## EC Certificate - Production Quality Assurance

Directive 93/42/EEC on Medical Devices, Annex V

No. Issued To: CE 01966 Mölnlycke Health Care AB Box 13080 Gamlestadsvägen 3C SE-402 52 Göteborg Sweden

In respect of:

See certificate scope page.

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

For and on behalf of BSI, a Notified Body for the above Directive (Notified Body Number 0086):

Stewart Brain, Head of Compliance & Risk - Medical Devices

First Issued: 1998-06-29

Date: 2018-05-30

Expiry Date: 2023-06-28

...making excellence a habit.<sup>™</sup> Page 1 of 2

Validity of this certificate is conditional on the quality system being maintained to the requirements of the Directive as demonstrated through the required surveillance activities of the Notified Body. This approval excludes all products designed and/or manufactured by a third party on behalf of the company named on this certificate, unless specifically agreed with BSI.

This certificate was issued electronically and is bound by the conditions of the contract.

Information and Contact: BSI, Kitemark Court, Davy Avenue, Knowlhill, Milton Keynes MK5 8PP. Tel: + 44 345 080 9000 BSI Assurance UK Limited, registered in England under number 7805321 at 389 Chiswick High Road, London W4 4AL, UK. A member of BSI Group of Companies.





Certificate No: CE 01966

### Certificate Scope:

Those aspects of manufacture related to securing and maintaining sterility of absorbent tracheostomy dressing, sterile scar management dressing and transparent adhesive IV film dressing.

Those aspects of manufacture related to securing and maintaining sterility of negative pressure wound therapy (NPWT) accessories, surgical and equipment drapes and surgical gowns.

Those aspects of manufacturing relating to securing and maintaining sterility in the assembly of procedure packs in accordance with article 12 of the MDD.

First Issued: 1998-06-29

Date: 2018-05-30

Expiry Date: 2023-06-28

...making excellence a habit.<sup>™</sup> Page 2 of 2

Validity of this certificate is conditional on the quality system being maintained to the requirements of the Directive as demonstrated through the required surveillance activities of the Notified Body. This approval excludes all products designed and/or manufactured by a third party on behalf of the company named on this certificate, unless specifically agreed with BSI.

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