

TEHNOMEDICA

str.Ciuflea, 38/1 MD-2001, mun. Chișinău, Moldova tel./fax: (022)601 102, 601 087
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Anexa nr. 7
la Documentația standard nr.115
din 15.09.2021

CERERE DE PARTICIPARE

Către **Centrul pentru Achiziții Publice Centralizate în Sănătate**

Stimați domni,

Ca urmare a anunțului/invitației de participare/de preselecție apărut în Buletinul achizițiilor publice și/sau Jurnalul Oficial al Uniunii Europene, BAP Nr.67 din 26.08.2022 privind aplicarea procedurii pentru atribuirea contractului privind achiziționarea dispozitivelor medicale (pansamente) conform Programului Național pentru combaterea maladiilor rare - "Epidermoliza buloasă", pentru anul 2023, noi, Tehnomedica SRL, 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: 01.10.2022

Cu stimă,

Tehnomedica SRL

Director Tatiana Roibu

(semnătura autorizată)

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din 15.09.2021

DECLARAȚIE privind valabilitatea ofertei

Către **Centrul pentru Achiziții Publice Centralizate în Sănătate**

Stimați domni,

Ne angajăm să menținem oferta valabilă, privind achiziționarea dispozitivelor medicale (pansamente) conform Programului Național pentru combaterea maladiilor rare - "Epidermoliza buloasă", pentru anul 2023 prin procedura de achiziție licitație deschisă, pentru o durată de 160 zile, (una sută șaiszeci zile), respectiv până la data de 14.03.2023 (ziua/luna/anul), și ea va rămâne obligatorie pentru noi și poate fi acceptată oricând înainte de expirarea perioadei de valabilitate.

Data completării: 01.10.2022

Cu stimă,

Tehnomedica SRL

Director Tatiana Roibu

(semnătura autorizată)

REPUBLICA



MOLDOVA

CERTIFICAT DE ÎNREGISTRARE

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

Data eliberării

16.02.2005

Bolboceanu Adela, registrator de stat

*Funcția, numele, prenumele persoanei
care a eliberat certificatul*

semnătura

MD 0027040



Nr. CIF26-842.2020
Data: 13 Februarie 2020

**CERTIFICAT
PRIVIND EXISTENTA CONTURILOR CURENTE**

Prin prezentul, **Mobiasbanca - OTP Group S.A.**, codul băncii (BIC): **MOBBMD22**, confirmă că compania **TEHNOMEDICA S.R.L.** cod fiscal (IDNO) **1002600053256**, detine următoarele conturi curente la Mobiasbanca - OTP Group S.A., Sucursala. 26 Negruzzi:

1. **MDL - MD65MO2224ASV98310887100**
2. **EUR - MD06MO2224ASV98311097100**


L.S.
Numele, Prenumele si Semnatura
Director sucursalei „Gheorghe Mocanu”



Executor :Eduard Cilcic
Tel: 022-812-150

LISTA FONDATORILOR SRL TEHNOMEDICA

Fondator unic: Roibu Tatiana

IDNP: date cu caracter personal

SITUAȚIILE FINANCIARE
pentru perioada 01.01.2021 - 31.12.2021

Entitatea: SRL "TEHNOMEDICA"
Cod CUIO: 37700778
Cod IDNO: 1002600053256

Sediul:
MD:
Raionul(municipiul): 102, DDF CENTRU
Cod CUATM: 0130, SEC.CENTRU
Strada:

Activitatea principală: G4646, Comert cu ridicata al produselor farmaceutice
Forma de proprietate: 16, Proprietate colectivă
Forma organizatorico-juridică: 530, Societăți cu răspundere limitată

Date de contact:
Telefon: 022601102
WEB:
E-mail:
Numele și coordonatele al contabilului-șef: DI (dna) Popescu Ecaterina Tel. 069153407

Numărul mediu al salariaților în perioada de gestiune: 5 persoane.

Persoanele responsabile de semnarea situațiilor financiare* Popescu Ecaterina

Unitatea de măsură: leu

BILANȚUL

Anexa 1

la 31.12.2021

Nr. cpt.	Indicatori	Cod rd.	Sold la	
			Începutul perioadei de gestiune	Sfârșitul perioadei de gestiune
1	2	3	4	5
	A C T I V			
	ACTIVE IMOBILIZATE			
	I. Imobilizări necorporale			
	1. Imobilizări necorporale în curs de execuție	010	0	0
	2. Imobilizări necorporale în exploatare, total	020	319	255
	din care:			
	2.1. concesiuni, licențe și mărci	021	0	0
	2.2. drepturi de autor și titluri de protecție	022	0	0
	2.3. programe informatice	023	0	0
	2.4. alte imobilizări necorporale	024	319	255
	3. Fond comercial	030	0	0
	4. Avansuri acordate pentru imobilizări necorporale	040	2861138	3906847
	Total imobilizări necorporale (rd.010 + rd.020 + rd.030 + rd.040)	050	2861457	3907102
	II. Imobilizări corporale			
	1. Imobilizări corporale în curs de execuție	060	0	0
	2. Terenuri	070	0	0
	3. Mijloace fixe, total	080	2174915	1775960
	din care:			
	3.1. clădiri	081	1038156	929187
	3.2. construcții speciale	082	0	0
	3.3. mașini, utilaje și instalații tehnice	083	34609	18080
	3.4. mijloace de transport	084	1042950	806787

A.	3.5. inventar și mobilier	085	0	0
	3.6. alte mijloace fixe	086	59200	21906
	4. Resurse minerale	090	0	0
	5. Active biologice imobilizate	100	0	0
	6. Investiții imobiliare	110	0	0
	7. Avansuri acordate pentru imobilizări corporale	120	0	0
	Total imobilizări corporale (rd.060 + rd.070 + rd.080 + rd.090 + rd.100 + rd.110 + rd.120)	130	2174915	1775960
	III. Investiții financiare pe termen lung			
	1. Investiții financiare pe termen lung în părți neafiliate	140	0	0
	2. Investiții financiare pe termen lung în părți afiliate, total	150	0	0
	din care:			
	2.1. acțiuni și cote de participație deținute în părțile afiliate	151	0	0
	2.2 împrumuturi acordate părților afiliate	152	0	0
	2.3 împrumuturi acordate aferente intereselor de participare	153	0	0
	2.4 alte investiții financiare	154	0	0
	Total investiții financiare pe termen lung (rd.140 + rd.150)	160	0	0
	IV. Creanțe pe termen lung și alte active imobilizate			
	1. Creanțe comerciale pe termen lung	170	0	0
	2. Creanțe ale părților afiliate pe termen lung	180	0	0
	inclusiv: creanțe aferente intereselor de participare	181	0	0
	3. Alte creanțe pe termen lung	190	0	0
	4. Cheltuieli anticipate pe termen lung	200	0	0
	5. Alte active imobilizate	210	0	0
	Total creanțe pe termen lung și alte active imobilizate (rd.170 + rd.180 + rd.190 + rd.200 + rd.210)	220	0	0
	TOTAL ACTIVE IMOBILIZATE (rd.050 + rd.130 + rd.160 + rd.220)	230	5036372	5683062
B.	ACTIVE CIRCULANTE			
	I. Stocuri			
	1. Materiale și obiecte de mică valoare și scurtă durată	240	4996	4996
	2. Active biologice circulante	250	0	0
	3. Producția în curs de execuție	260	0	0
	4. Produse și mărfuri	270	1326025	491017
	5. Avansuri acordate pentru stocuri	280	0	0
	Total stocuri (rd.240 + rd.250 + rd.260 + rd.270 + rd.280)	290	1331021	496013
	II. Creanțe curente și alte active circulante			
	1. Creanțe comerciale curente	300	1022910	2800290
	2. Creanțe ale părților afiliate curente	310	0	0
	inclusiv: creanțe aferente intereselor de participare	311	0	0
	3. Creanțe ale bugetului	320	6546	6546
	4. Creanțele ale personalului	330	0	0
	5. Alte creanțe curente	340	0	0
	6. Cheltuieli anticipate curente	350	13569	2114
	7. Alte active circulante	360	31297	
	Total creanțe curente și alte active circulante (rd.300 + rd.310 + rd.320 + rd.330 + rd.340 + rd.350 + rd.360)	370	1074322	2808950
	III. Investiții financiare curente			
	1. Investiții financiare curente în părți neafiliate	380	700000	0
	2. Investiții financiare curente în părți afiliate, total	390	0	0
	din care:			
	2.1. acțiuni și cote de participație deținute în părțile afiliate	391	0	0
	2.2. împrumuturi acordate părților afiliate	392	0	0
	2.3. împrumuturi acordate aferente intereselor de participare	393	0	0

	2.4. alte investiții financiare în părți afiliate	394	0	0
	Total investiții financiare curente (rd.380 + rd.390)	400	700000	0
	IV. Numerar și documente bănești	410	6916759	6867356
	TOTAL ACTIVE CIRCULANTE (rd.290 + rd.370 + rd.400 + rd.410)	420	10022102	10172319
	TOTAL ACTIVE (rd.230 + rd.420)	430	15058474	15855381
	P A S I V			
C.	CAPITAL PROPRIU			
	I. Capital social și neînregistrat			
	1. Capital social	440	5400	5400
	2. Capital nevărsat	450	(0)	(0)
	3. Capital neînregistrat	460	0	0
	4. Capital retras	470	(0)	(0)
	5. Patrimoniul primit de la stat cu drept de proprietate	480	0	0
	Total capital social și neînregistrat (rd.440 + rd.450 + rd.460 + rd.470 + rd.480)	490	5400	5400
	II. Prime de capital	500	0	0
	III. Rezerve			
	1. Capital de rezervă	510	0	0
	2. Rezerve statutare	520	0	0
	3. Alte rezerve	530	0	0
	Total rezerve (rd.510 + rd.520 + rd.530)	540	0	0
	IV. Profit (pierdere)			
	1. Corecții ale rezultatelor anilor precedenți	550	X	0
	2. Profit nerepartizat (pierdere neacoperită) al anilor precedenți	560	14705486	12203358
	3. Profit net (pierdere netă) al perioadei de gestiune	570	X	2864668
	4. Profit utilizat al perioadei de gestiune	580	X	(0)
	Total profit (pierdere) (rd.550 + rd.560 + rd.570 + rd.580)	590	14705486	15068026
	V. Rezerve din reevaluare	600	0	0
	VI. Alte elemente de capital propriu	610	0	0
	TOTAL CAPITAL PROPRIU (rd.490 + rd.500 + rd.540 + rd.590 + rd.600 + rd.610)	620	14710886	15073426
D.	DATORII PE TERMEN LUNG			
	1. Credite bancare pe termen lung	630	0	0
	2. Împrumuturi pe termen lung	640	0	0
	din care:			
	2.1. împrumuturi din emisiunea de obligațiuni	641	0	0
	inclusiv: împrumuturi din emisiunea de obligațiuni convertibile	642	0	0
	2.2. alte împrumuturi pe termen lung	643	0	0
	3. Datorii comerciale pe termen lung	650	0	0
	4. Datorii față de părțile afiliate pe termen lung	660	0	0
	inclusiv: datorii aferente intereselor de participare	661	0	0
	5. Avansuri primite pe termen lung	670	0	0
	6. Venituri anticipate pe termen lung	680	0	0
	7. Alte datorii pe termen lung	690	0	0
	TOTAL DATORII PE TERMEN LUNG (rd.630 + rd.640 + rd.650 + rd.660 + rd.670 + rd.680 + rd.690)	700	0	0
	DATORII CURENTE			
	1. Credite bancare pe termen scurt	710	0	0

E.	2. Împrumuturi pe termen scurt, total	720	0	0
	din care:			
	2.1. Împrumuturi din emisiunea de obligațiuni	721	0	0
	inclusiv: împrumuturi din emisiunea de obligațiuni convertibile	722	0	0
	2.2. alte împrumuturi pe termen scurt	723	0	0
	3. Datorii comerciale curente	730	149510	288048
	4. Datorii față de părțile afiliate curente	740	0	0
	inclusiv: datorii aferente intereselor de participare	741	0	0
	5. Avansuri primite curente	750	0	0
	6. Datorii față de personal	760	977	100
	7. Datorii privind asigurările sociale și medicale	770	0	0
	8. Datorii față de buget	780	197101	493807
	9. Datorii față de proprietari	790	0	0
	10. Venituri anticipate curente	800	0	0
	11. Alte datorii curente	810	0	0
	TOTAL DATORII CURENTE (rd.710 + rd.720 + rd.730 + rd.740 + rd.750 + rd.760 + rd.770 + rd.780 + rd.790 + rd.800 + rd.810)	820	347588	781955
F.	PROVIZIOANE			
	1. Provizioane pentru beneficiile angajaților	830	0	0
	2. Provizioane pentru garanții acordate cumpărătorilor/clientilor	840	0	0
	3. Provizioane pentru impozite	850	0	0
	4. Alte provizioane	860	0	0
	TOTAL PROVIZIOANE (rd.830 + rd.840 + rd.850 + rd.860)	870	0	0
	TOTAL PASIVE (rd.620 + rd.700 + rd.820 + rd.870)	880	15058474	15855381

SITUAȚIA DE PROFIT ȘI PIERDERE

de la 01.01.2021 până la 31.12.2021

Anexa 2

Indicatori	Cod rd.	Perioada de gestiune	
		precedenta	curenta
1	2	3	4
Venituri din vânzări, total	010	16620028	21962759
din care:			
venituri din vânzarea produselor și mărfurilor	011	13778008	19347659
venituri din prestarea serviciilor și executarea lucrărilor	012	2842020	2615100
venituri din contracte de construcție	013	0	0
venituri din contracte de leasing	014	0	0
venituri din contracte de microfinanțare	015	0	0
alte venituri din vânzări	016	0	0
Costul vânzărilor, total	020	12527753	16833673
din care:			
valoarea contabilă a produselor și mărfurilor vândute	021	11595535	16054470
costul serviciilor prestate și lucrărilor executate terților	022	932218	779203
costuri aferente contractelor de construcție	023	0	0
costuri aferente contractelor de leasing	024	0	0
costuri aferente contractelor de microfinanțare	025	0	0
alte costuri aferente vânzărilor	026	0	0
Profit brut (pierdere brută) (rd.010 - rd.020)	030	4092275	5129086
Alte venituri din activitatea operațională	040	986	1238
Cheltuieli de distribuire	050	0	
Cheltuieli administrative	060	1569273	1518335
Alte cheltuieli din activitatea operațională	070	17430	79468

Rezultatul din activitatea operațională: profit (pierdere) (rd.030 + rd.040 - rd.050 - rd.060 - rd.070)	080	2506558	3532521
Venituri financiare, total	090	1257613	365919
din care:			
venituri din interese de participare	091	0	0
inclusiv: veniturile obținute de la părțile afiliate	092	0	0
venituri din dobânzi	093	0	0
inclusiv: veniturile obținute de la părțile afiliate	094	0	0
venituri din alte investiții financiare pe termen lung	095	0	0
inclusiv: veniturile obținute de la părțile afiliate	096	0	0
venituri aferente ajustărilor de valoare privind investițiile financiare pe termen lung și curente	097	0	0
venituri din ieșirea investițiilor financiare	098	0	0
venituri aferente diferențelor de curs valutar și de sumă	099	1257613	365919
Cheltuieli financiare, total	100	666851	603812
din care:			
cheltuieli privind dobânzile	101	0	0
inclusiv: cheltuielile aferente părților afiliate	102	0	0
cheltuieli aferente ajustărilor de valoare privind investițiile financiare pe termen lung și curente	103	0	0
cheltuieli aferente ieșirii investițiilor financiare	104	0	0
cheltuieli aferente diferențelor de curs valutar și de sumă	105	666851	603812
Rezultatul: profit (pierdere) financiar(ă) (rd.090 - rd.100)	110	590762	-237893
Venituri cu active imobilizate și excepționale	120	0	0
Cheltuieli cu active imobilizate și excepționale	130	0	0
Rezultatul din operațiuni cu active imobilizate și excepționale: profit (pierdere) (rd.120 - rd.130)	140	0	0
Rezultatul din alte activități: profit (pierdere) (rd.110 + rd.140)	150	590762	-237893
Profit (pierdere) pînă la impozitare (rd.080 + rd.150)	160	3097320	3294628
Cheltuieli privind impozitul pe venit	170	410288	429960
Profit net (pierdere netă) al perioadei de gestiune (rd.160 - rd.170)	180	2687032	2864668

SITUAȚIA MODIFICĂRILOR CAPITALULUI PROPRIU

de la 01.01.2021 pînă la 31.12.2021

Anexa 3

Nr. d/o	Indicatori	Cod rd	Sold la începutul perioadei de gestiune	Majorări	Diminuări	Sold la sfîrșitul perioadei de gestiune
1	2	3	4	5	6	7
I.	Capital social și neînregistrat					
	1. Capital social	010	5400	0	0	5400
	2. Capital nevărsat	020	(0)	(0)	(0)	(0)
	3. Capital neînregistrat	030	0	0	0	0
	4. Capital retras	040	(0)	(0)	(0)	(0)
	5. Patrimoniul primit de la stat cu drept de proprietate	050	0	0	0	0
	Total capital social și neînregistrat (rd.010 + rd.020 + rd.030 + rd.040 + rd.050)	060	5400	0	0	5400
II.	Prime de capital	070	0	0	0	0
III.	Rezerve					
	1. Capital de rezervă	080	0	0	0	0
	2. Rezerve statutare	090	0	0	0	0
	3. Alte rezerve	100	0	0	0	0
	Total rezerve (rd.080 + rd.090 + rd.100)	110	0	0	0	0

IV.	Profit (pierdere)					
	1. Corecții ale rezultatelor anilor precedenți	120	X	0	0	0
	2. Profit nerepartizat (pierdere neacoperită) al anilor precedenți	130	14705486	0	2502128	12203358
	3. Profit net (pierdere netă) al perioadei de gestiune	140	X	2864668	0	2864668
	4. Profit utilizat al perioadei de gestiune	150	X	(0)	(0)	(0)
	Total profit (pierdere) (rd.120 + rd.130 + rd.140 + rd.150)	160	14705486	2864668	2502128	15068026
V.	Rezerve din reevaluare	170	0	0	0	0
VI.	Alte elemente de capital propriu	180	0	0	0	0
	Total capital propriu (rd.060 + rd.070 + rd.110 + rd.160 + rd.170 + rd.180)	190	14710886	2864668	2502128	15073426

SITUAȚIA FLUXURILOR DE NUMERAR
de la 01.01.2021 până la 31.12.2021

Anexa 4

Indicatori	Cod rd	Perioada de gestiune	
		precedentă	curentă
1	2	3	4
Fluxuri de numerar din activitatea operațională			
Încasări din vânzări	010	17211991	23444192
Plăți pentru stocuri și servicii procurate	020	13370873	18993827
Plăți către angajați și organe de asigurare socială și medicală	030	554000	456668
Dobânzi plătite	040	0	0
Plata impozitului pe venit	050	414542	331089
Alte încasări	060	2220519	1598942
Alte plăți	070	5025576	3293285
Fluxul net de numerar din activitatea operațională (rd.010 - rd.020 - rd.030 - rd.040 - rd.050 + rd.060 - rd.070)	080	67519	1968265
Fluxuri de numerar din activitatea de investiții			
Încasări din vânzarea activelor imobilizate	090	0	0
Plăți aferente intrărilor de active imobilizate	100	0	0
Dobânzi încasate	110	0	0
Dividende încasate	120	0	0
inclusiv: dividende încasate din străinătate	121		
Alte încasări (plăți)	130	0	0
Fluxul net de numerar din activitatea de investiții (rd.090 - rd.100 + rd.110 + rd.120 ± rd.130)	140	0	0
Fluxuri de numerar din activitatea financiară			
Încasări sub formă de credite și împrumuturi	150	630000	800000
Plăți aferente rambursării creditelor și împrumuturilor	160	830000	0
Dividende plătite	170	2019064	2512000
inclusiv: dividende plătite nerezidenților	171		
Încasări din operațiuni de capital	180	0	0
Alte încasări (plăți)	190	0	0
Fluxul net de numerar din activitatea financiară (rd.150 - rd.160 - rd.170 + rd.180 ± rd.190)	200	-2219064	-1712000
Fluxul net de numerar total (± rd.080 ± rd.140 ± rd.200)	210	-2151545	256265
Diferențe de curs valutar favorabile (nefavorabile)	220	401419	-305668
Sold de numerar la începutul perioadei de gestiune	230	8666885	6916759
Sold de numerar la sfârșitul perioadei de gestiune (± rd.210 ± rd.220 + rd.230)	240	6916759	6867356

CERTIFICAT
privind lipsa sau existența restanțelor față de bugetul public național

Nr.
№ **A2218603**

din
от **28.09.2022**

1. Destinația / Назначение

PENTRU PARTICIPAREA LA PROCEDURI PRIVIND ACHIZIȚIILE PUBLICE

2. Date despre contribuabil / Информация о налогоплательщике

Denumirea Наименование	Codul fiscal / Numărul de identificare Фискальный код / Идентификационный номер
TEHNOMEDICA S.R.L.	1002600053256
Adresa sediului de bază (strada, numărul) Адрес основного месторасположения (улица, номер)	Codul - Denumirea localității Код - Наименование населенного пункта
Ciuflea nr.38 bl.1	0130-SEC.CENTRU

3. Atestarea lipsei sau existenței restanțelor conform datelor Sistemului Informațional Automatizat /

Подтверждение отсутствия или наличия недоимки согласно данных Информационной автоматизированной системы

La data emiterii prezentului certificat restanța față de bugetul public național constituie/ На дату выдачи данной справки недоимка перед национальным публичным бюджетом составляет:
0,00 lei/лей.

4. Valabil pînă la / Действителен до 13.10.2022

5. Autentificarea Serviciului Fiscal de Stat / Подтверждение Государственной налоговой службы

Șef DDF CENTRU
Funcția/Должность

Digitally signed by Iasinschi Veronica
Date: 2022.09.29 08:38:02 EEST
Reason: MoldSign Signature
Location: Moldova Semnătura/Подпись



Veronica IASINSCHI
Numele și prenumele/Фамилия и имя

L.Ș/ М.П.

Executor: **Fodor V.**
Numele și prenumele/Фамилия и имя

Este extras din Sistemul Informațional al SFS SIA „Contul curent al contribuabilului”// 28.09.2022 ora 14:48:57
cu aplicarea prevederilor pct. 82-83 Ordin IFPS nr.400 din 14.03.2014 (Monitorul Oficial 72-77/399, 28.03.2014)

[NOTA \(304,54\)](#)

TEHNOMEDICA

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

**Către Centrul pentru Achiziții Publice
Centralizate în Sănătate**

În atenția Grupului de lucru
al Licităției Deschise nr. ocds-b3wdp1-MD-1662647101714,
ID: 21063206

Declarație privind disponibilitatea prezentării mostrelor

Prin prezenta, declarăm că vom prezenta mostre în decurs de 10 zile de la solicitarea autorității contractante pentru produsele oferite în cadrul licitației preonate privind achiziționarea dispozitivelor medicale (pansamente) conform Programului Național pentru combaterea maladiilor rare - "Epidermoliza buloasă", pentru anul 2023.

Cu respect,

Director

Tatiana Roibu

TEHNOMEDICA

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e-mail <tehnomedica_md@yahoo.com> <tehnomedicamd@gmail.com>

**Către Centrul pentru Achiziții Publice
Centralizate în Sănătate**

În atenția Grupului de lucru
al Licităției Deschise nr. ocds-b3wdp1-MD-1662536410242,
ID: 21063000

Declarație privind înregistrarea dispozitivelor medicale

Prin prezenta, declarăm că, produsele oferite în cadrul licitației deschise prenotate sunt înregistrate în Registrul de Stat al Dispozitivelor Medicale a Agenției Medicamentului și Dispozitivelor Medicale.

Dovada înregistrării dispozitivelor medicale se regăsește pe pagina web a Agenției Medicamentului și Dispozitivelor Medicale www.amdm.gov.md

Numerele de înregistrare se regăsesc în anexa prezentei declarații.

Pentru produsul oferit la lotul nr.7 Mesoft, cod 156329, a fost depusă notificarea privind înregistrarea dispozitivelor medicale. La solicitarea autorității contractante, numărul de înregistrare va fi prezentat în termen de 15 zile.

Cu respect,

Director

Tatiana Roibu

DM000374534	PANSAMENT ABSORBANT ADEZIV PE BAZĂ DE SPUMĂ DE POLIURETAN SI SILICON	MEPILEX®	TRANSFER	294502	Suedia	MÖLNLYCKE HEALTH CARE AB	TEHNOMEDICA S.R.L.	Rg04-000229	27-09-2022
DM000374528	PANSAMENT ABSORBANT ADEZIV PE BAZĂ DE SPUMĂ DE POLIURETAN SI SILICON	MEPILEX®	EM	284322	Suedia	MÖLNLYCKE HEALTH CARE AB	TEHNOMEDICA S.R.L.	Rg04-000229	27-09-2022
DM000374545	PANSAMENT ABSORBANT ADEZIV PE BAZĂ DE SPUMĂ DE POLIURETAN SI SILICON	MEPILEX®	XT	211100	Suedia	MÖLNLYCKE HEALTH CARE AB	TEHNOMEDICA S.R.L.	Rg04-000229	27-09-2022
DM000374561	PANSAMENT ABSORBANT ADEZIV PE BAZĂ DE SPUMĂ DE POLIURETAN SI SILICON, PENTRU CĂLCĂI	MEPILEX®	HEEL	288100	Suedia	MÖLNLYCKE HEALTH CARE AB	TEHNOMEDICA S.R.L.	Rg04-000229	27-09-2022
DM000374490	BANDAJ ELASTIC TUBULAR	TUBIFAST™	RED LINE, 10M, N20	2434	Suedia	MÖLNLYCKE HEALTH CARE AB	TEHNOMEDICA S.R.L.	Rg04-000229	27-09-2022
DM000374491	BANDAJ ELASTIC TUBULAR	TUBIFAST™	GREEN LINE, 10M, N24	2436	Suedia	MÖLNLYCKE HEALTH CARE AB	TEHNOMEDICA S.R.L.	Rg04-000229	27-09-2022
DM000374492	BANDAJ ELASTIC TUBULAR	TUBIFAST™	BLUE LINE, 10M, N30	2438	Suedia	MÖLNLYCKE HEALTH CARE AB	TEHNOMEDICA S.R.L.	Rg04-000229	27-09-2022
DM000374493	BANDAJ ELASTIC TUBULAR	TUBIFAST™	YELLOW LINE, 10M, N12	2440	Suedia	MÖLNLYCKE HEALTH CARE AB	TEHNOMEDICA S.R.L.	Rg04-000229	27-09-2022
DM000374494	BANDAJ ELASTIC TUBULAR	TUBIFAST™	PURPLE LINE, 10M, N15	2444	Suedia	MÖLNLYCKE HEALTH CARE AB	TEHNOMEDICA S.R.L.	Rg04-000229	27-09-2022

TEHNOMEDICA

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**Către Centrul pentru Achiziții Publice
Centralizate în Sănătate**

În atenția Grupului de lucru
al Licităției Deschise nr. ocds-b3wdp1-MD-1662647101714,
ID: 21063206

Declarație privind termenul de valabilitate

Prin prezenta, declarăm că termenul de valabilitate pentru produsele oferite în cadrul licitației preonate privind achiziționarea dispozitivelor medicale (pansamente) conform Programului Național pentru combaterea maladiilor rare - "Epidermoliza buloasă", pentru anul 2023 va constitui nu mai puțin de 60% din termenul inițial pentru bunurile cu o valabilitate de 2 ani și nu mai puțin de 80% din cel inițial pentru bunurile cu o valabilitate de până la 2 ani.

Cu respect,

Director

Tatiana Roibu

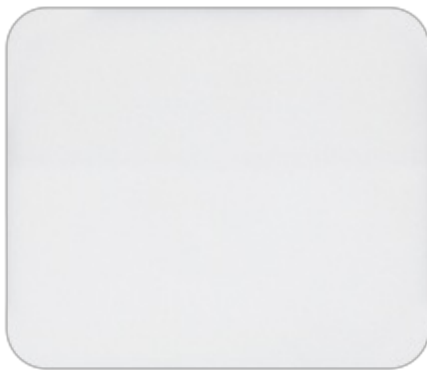
294502 Mepilex Transfer

Soft silicone exudate transfer dressing

Product details

Size : 20cm x 50cm
Descriptive feature : Exudate transfer, Foam, Non-border, Soft silicone
Sterile : Sterile

Images



Delivered items

294502-03

Sales released in: Algeria, Australia, Austria, Bahrain, Belarus, Belgium, Bosnia and Herzegovina, Bulgaria, Canada, China, Croatia, Czechia, Denmark, Estonia, Faroe Islands, Finland, France, Germany, Greece, Hong Kong, Hungary, Iceland, Iran (Islamic Republic of), Ireland, Israel, Italy, Japan, Kazakhstan, Kuwait, Latvia, Lithuania, Luxembourg, Macedonia (the former Yugoslav Republic of), Malaysia, Moldova (the Republic of), Netherlands, New Zealand, Norway, Poland, Portugal, Romania, Russian Federation, Saudi Arabia, Serbia, Singapore, Slovakia, Slovenia, South Africa, Spain, Sweden, Switzerland, Taiwan (Province of China), Thailand, Turkey, Ukraine, United Arab Emirates, United Kingdom of Great Britain and Northern Ireland

Country of origin: Finland

Shelf life: 3 years

Sterilization method: EtO

Packing information: First packaging layer is a peel-open sterile barrier, plastic/plastic. Once opened the sterile barrier cannot be closed again. Second layer is a cardboard dispenser box. Third layer is a corrugated board transport box.

Is suitable for Tray: No

Packing level	Quantity	GS1 code	WxLxH (mm)	Vol (dm ³)	Weight gross/net (kg)
Consumer pack	1	7332430003775			

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Packing level	Quantity	GS1 code	WxLxH (mm)	Vol (dm ³)	Weight gross/net (kg)
Dispenser box	4	7323190024032			
Transport box	24	7323190024025			
Pallet	2016	7323190024018			

Material

Natural rubber latex : No

Product Composition Wound Contact Layers

Product Component	Composition
Transferring layer	Polyurethane foam
Wound contact layer	Silicone
Protective release liner	Polyethylene film

Product Performance Wound Contact Layer Products

Characteristics	Test Method	Internal Test Method	Unit	Requirement	Product Performance
Free Swell Absorptive Capacity	EN 13726-1 part 3:2	T-1069	g/100 cm ²	Not specified	N/A
Free Swell Absorptive Capacity	EN 13726-1 part 3:2	T-1069	g/g	Not specified	N/A
Conformability-Extensibility, MD	EN 13726-4	T-1086	N/cm	Not specified	N/A
Conformability-Extensibility, CD	EN 13726-4	T-1086	N/cm	Not specified	N/A
Conformability-Permanent Set, MD	EN 13726-4	T-1086	%	Not specified	N/A
Conformability-Permanent Set, CD	EN 13726-4	T-1086	%	Not specified	N/A

Technical

Dimension

Dimension text	Dimension value
Product	20 cm x 50 cm

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Dimension text	Dimension value
Product	8 in x 20 in

Classifications

Regulation type(s)	MDD Class IIb	Locally Regulated	Unregulated
CE Certificate Number :	CE 01965		
Notified body medical devices/PPE :	BSI (2797)		
Intended use MDD :	Mepilex Transfer is designed for a wide range of exuding and difficult-to-dress wounds. Mepilex Transfer can also be used as a protective layer on non-exuding wounds and/or large areas of fragile skin. Mepilex Transfer can be used under compression.		
Sales released in :	Austria, Belgium, Bulgaria, Croatia, Czechia, Denmark, Estonia, Faroe Islands, Finland, France, Germany, Greece, Hungary, Iceland, Ireland, Italy, Latvia, Lithuania, Luxembourg, Netherlands, Norway, Poland, Portugal, Romania, Slovakia, Slovenia, Spain, Sweden, Switzerland, United Kingdom of Great Britain and Northern Ireland	Algeria, Australia, Bahrain, Canada, China, Israel, Japan, Kazakhstan, Kuwait, Macedonia (the former Yugoslav Republic of), Malaysia, Moldova (the Republic of), New Zealand, Russian Federation, Saudi Arabia, Serbia, Singapore, Taiwan (Province of China), Thailand, Turkey, Ukraine, United Arab Emirates	Belarus, Bosnia and Herzegovina, Hong Kong, Iran (Islamic Republic of), South Africa

Applied standards : The standards presented below is a selection of the most essential standards that are adhered to.

EN 1041, EN ISO 9001, EN ISO 13485, EN ISO 10993-1, EN ISO 10993-5, EN ISO 10993-7, EN ISO 11607-1, EN ISO 11607-2, EN ISO 15223-1, EN ISO 10993-11, EN ISO 10993-10, EN ISO 10993-18, ISO 14001

Removable label

No

GMDN Code (Global Medical Device Nomenclature)

46855 Wound - nonadherent dressing, permeable

Find out more at www.molnlycke.com

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284322 Mepilex EM

Absorbent soft silicone dressing

Product details

Size : 17.5cm x 17.5cm
Descriptive feature : Foam, Non-border, Soft silicone, Thin
Sterile : Sterile

Images



Delivered items

284322-01

Sales released in: Bosnia and Herzegovina, Bulgaria, Croatia, France, Greece, Hungary, Israel, Macedonia (the former Yugoslav Republic of), Martinique, Moldova (the Republic of), Pakistan, Poland, Portugal, Romania, Serbia, Slovakia, Slovenia, Spain, Turkey

Country of origin: Finland

Shelf life: 3 years

Sterilization method: EtO

Packing information: First packaging layer is a peel open sterile barrier, paper/plastic. Once opened the sterile barrier cannot be closed again. Second layer is a cardboard dispenser box. Third layer is a corrugated board transport box.

Is suitable for Tray: No

Packing level	Quantity	GS1 code	WxLxH (mm)	Vol (dm ³)	Weight gross/net (kg)
Consumer pack	1	7332430666642			
Dispenser box	5	7323190126606	26x220x236		
Transport box	35	7323190126590	234x263x215	13.2	1.3 / 0.5

Find out more at www.molnlycke.com

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Packing level	Quantity	GS1 code	WxLxH (mm)	Vol (dm ³)	Weight gross/net (kg)
Pallet	4200	7323190126583	800x1200x1884		

Material

Animal tissues : No
Natural rubber latex : No
Medicinal substances : No

Product Composition Non-bordered Foam Products

Product Component	Composition
Backing material	Polyurethane film
Wound pad	Polyurethane foam
Wound contact layer	Silicone
Protective release liner	Polyethylene film

Product Performance Non-bordered Foam Products

Characteristics	Test Method	Internal Test Method	Unit	Requirement	Product Performance
Free Swell Absorptive Capacity	EN 13726-1 part 3:2	T-1069	g/100 cm ²	Not specified	N/A
Free Swell Absorptive Capacity	EN 13726-1 part 3:2	T-1069	g/g	Not specified	N/A
Fluid Handling Capacity	EN 13726-1 part 3:3	T-1068	g/10 cm ² /24 h	Not specified	7.3
Absorbency	EN 13726-1 part 3:3	T-1068	g/10 cm ² /24 h	Not specified	1.82
Moisture Vapour Transmission Rate (MVTR)	EN 13726-1 part 3:3	T-1068	g/10 cm ² /24 h	Not specified	5.5
Moisture Vapour Transmission Rate (MVTR) of a wound dressing when in contact with water vapour	EN 13726-2 part 3:2	T-1070	g/m ² /24 h	Not specified	2282

Find out more at www.molnlycke.com

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Characteristics	Test Method	Internal Test Method	Unit	Requirement	Product Performance
Moisture Vapour Transmission Rate (MVTR) of a wound dressing when in contact with liquid	EN 13726-2 part 3:3	T-1075	g/m ² /24 h	Not specified	4617
Waterproofness	EN 13726-3	T-1083	Pass/Fail	>500 mm H ₂ O for 300 s	N/A
Conformability-Extensibility, MD	EN 13726-4	T-1086	N/cm	Not specified	0.6
Conformability-Extensibility, CD	EN 13726-4	T-1086	N/cm	Not specified	0.5
Conformability-Permanent Set, MD	EN 13726-4	T-1086	%	Not specified	0.3
Conformability-Permanent Set, CD	EN 13726-4	T-1086	%	Not specified	2.7
Resistance to microbial penetration - Wet	ISO 22610	T-1005	BI	6	N/A
Viral penetration	ASTM F 1671	N/A	Pass/Fail	29 out of 32 samples	N/A

Technical

Dimension

Dimension text	Dimension value
Product	17.5 cm x 17.5 cm
Product	6.9 in x 6.9 in

Find out more at www.molnlycke.com

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Classifications

Regulation type(s)	MDD Class IIb	Locally Regulated	Unregulated
MDD Classification Rule :	4		
CE Certificate Number :	CE 01965		
Notified body medical devices/PPE :	BSI (2797)		
Intended use MDD :	Mepilex Lite is designed for the management of a wide range of non to low exuding wounds, such as leg and foot ulcers, pressure ulcers, partial thickness burns, radiation skin reactions and Epidermolysis Bullosa. Mepilex Lite can also be used as protection of compromised and/or fragile skin.		
Sales released in :	Bulgaria, Croatia, France, Greece, Hungary, Martinique, Poland, Portugal, Romania, Slovakia, Slovenia, Spain	Bosnia and Herzegovina, Israel, Moldova (the Republic of), Pakistan, Serbia, Turkey	Macedonia (the former Yugoslav Republic of)

Applied standards : The standards presented below is a selection of the most essential standards that are adhered to.

EN 1041, EN ISO 9001, EN ISO 13485, EN ISO 10993-1, EN ISO 10993-5, EN ISO 10993-7, EN ISO 11607-1, EN ISO 11607-2, EN ISO 15223-1, EN ISO 10993-11, EN ISO 10993-10, EN ISO 10993-18, ISO 14001

Removable label

No

GMDN Code (Global Medical Device Nomenclature)

46854 Wound - nonadherent dressing, absorbent, sterile

UNSPSC

42311510 Foam dressings

Commodity Code

3005100000 Wadding, Gauze, Dressings, drapes singlepacked - adhesive articles, Sets mainly consisting thereof

Find out more at www.molnlycke.com

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Mesoft®

Absorberende, usteril og steril kompres til rensning af hud og sår, som beskyttelse samt absorption af ekssudat



Absorption af
sårekssudat^{1,2}

Mesoft® er et alsidigt, usterilt produkt, der kan anvendes til at rense huden inden kirurgiske procedurer, sårrensning og beskyttelse, samt til absorption af ekssudat, når den anvendes som sekundær bandage^{1,2}. Mesoft findes også som sterilt¹ produkt.



1. Mesoft® kompres
4 lag, steril 40g



2. Mesoft slidskompres
4 lag, steril 40g



3. Mesoft kompres,
usteril 40g



4. Mesoft kompres,
usteril 30g



5. Mesoft tampon,
steril 30g



6. Mesoft, usteril 30g

Anvendelsesområde

Mesoft kompresser og tamponer kan anvendes til en række områder. De kan anvendes til absorption, sårrensning, beskyttelse, polstring eller som sekundær bandage.

Fordele ved Mesoft

- Alsidige anvendelsesområder^{1,2}: rensning, beskyttelse, ekssudat håndtering
- Ikke-sensibiliserende og ikke-toksisk¹
- Steril¹

Mesoft kompres 4 lag, steril 40g

Varenr	Str. cm	Stk./pk.	Stk./inderkrt.	Stk./trp.krt.
156029	5 x 5	2	20	720
156040		2	150	3000
156065		5	150	3000
156129	7,5 x 7,5	2	20	520
156102		2	250	3000
156140		2	150	4200
156105		5	250	5500
156165		5	150	4200
156329	10 x 10	2	20	400
156340		2	150	3450
156365		5	150	3450
156310		10	150	2250
156440	10 x 20	2	120	1200
156465		5	120	1200

Mesoft slidskompres 4 lag, steril 40g

Varenr.	Str. cm	Stk./pk.	Stk./inderkrt.	Stk./trp.krt.
155030	10 x 10	1	130	1950

Mesoft tampon, steril 30g

Varenr.	Str.	Diam. mm	Stk./pk.	Stk./inderkrt.	Stk./trp.krt.
156760	Small	28	5	100	2800
156860	Medium	39	5	100	1500
156960	Large	45	5	70	1050
156961	Large	45	10	120	1200

Mesoft kompres, usteril 40g

Varenr.	Str. cm	Stk./inderkrt.	Stk./trp.krt.
156015	5 x 5	* 100	3200
156000		** 300	2400
156115	7,5 x 7,5	* 100	2000
156100		** 250	3750
156315	10 x 10	* 100	3000
156300		** 250	2500
156415	10 x 20	* 100	2800
156400		** 250	1250

* Papirpose

** Karton

Mesoft kompres, usteril 30g

Varenr.	Str. cm	Stk./inderkrt.	Stk./trp.krt.
157000	5 x 5	100	8000
156056		* 150	3150
157100	7,5 x 7,5	100	6000
156156		* 150	4200
157300	10 x 10	100	6000
156356		* 120	2160
157400	10 x 20	100	2200

* "Cleanbox" muliggør dispensering og yder samtidig beskyttelse mod luftbåren kontamination. Alle andre varenumre leveres i papirposer.

Mesoft tampon, usteril 30g

Varenr.	Str.	Diam. mm	Stk./inderkrt.	Stk./trp.krt.
156700	Small	28	100	2200
156800	Medium	39	100	1300
156900	Large	45	70	900

Referencer: 1. Mölnlycke Health Care. Data on file. 2019. 2. Mölnlycke Health Care. Data on file, 2016.

Læs mere på www.molnlycke.dk

Mölnlycke Health Care ApS, Gydevang 39, 3450 Allerød. Tlf.: +45 80 88 68 10. info.dk@molnlycke.com
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Mepilex® XT

The highly conformable foam dressing with Safetac® technology suitable for all wound healing stages

Safetac® layer

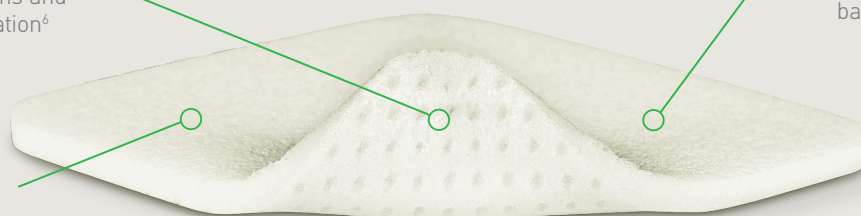
- Reduces pain and trauma at dressing change^{3,4}
- Does not adhere to the moist wound bed but to dry skin only
- Seals the wound margins and reduces risk of maceration⁶

Polyurethane backing film

- Breathable
- Waterproof
- Barrier to viruses and bacteria > 25nm⁸

Polyurethane foam pad with exudate channels

- Enhanced exudate management capacity
- Absorbs viscous exudate²
- Works well under compression^{1,2}
- Conformable and flexible foam with a pattern



Safetac®
TECHNOLOGY

- Effectively manages a wider range of exudate^{1,2} - absorbs viscous exudate²
- Non-traumatic to the wound and surrounding skin³ - fewer disturbances to the wound
- Less painful during dressing changes⁴ - less stress for patients⁵
- Enhanced exudate management¹ - reduced risk of leakage and maceration⁶
- Excellent conformity⁷ - Promotes patient comfort during wear

Proven choice for a better outcome

Safetac®, pioneered by Mölnlycke, delivers above and beyond the ordinary. Proven to help optimize the wound healing journey and even prevent wounds, dressings with Safetac are the safe choice for patients and a champion for higher standards in wound care.

In fact, we have a wealth of evidence that supports the clinical and economic benefits of dressings with Safetac, including Mepilex®, Mepitel®, Mepiform® and Mepitac®. To date, these dressings have helped millions of patients worldwide⁷⁻⁹.

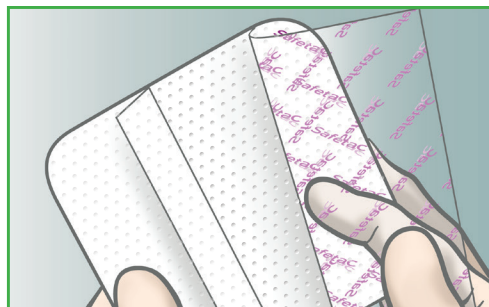
* A unique proprietary technology exclusive to Mölnlycke Health Care

Safetac®
TECHNOLOGY

Mepilex® XT


Mölnlycke®

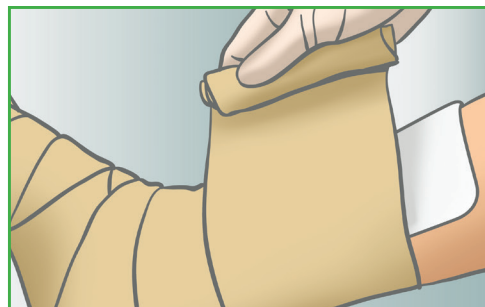
Directions for use



1. Cleanse the wound in accordance with normal procedures. Dry the surrounding skin thoroughly. Remove the release films. For best results, Mepilex® XT should overlap the dry surrounding skin by at least 1-2 cm for the smaller sizes (sizes up to 12.5 x 12.5 cm), and 5 cm for the larger sizes in order to protect the surrounding skin from maceration and excoriation.



2. Apply the dressing securely. If required, Mepilex® XT can be cut to size. Do not stretch. Mepilex® XT does not need to be cut to the size of the wound when compression is used.



3. When necessary, secure Mepilex® XT with a bandage or other secondary fixation product. Mepilex® XT may be left in place for several days depending on the condition of the wound and surrounding skin.

How Mepilex®XT works

Mepilex® XT is a soft and highly conformable foam dressing with a pattern on the wound contact side that can effectively manage a wider range of exudate and maintain a moist wound environment.

The Safetac® layer seals wound edges, preventing the exudate from leaking onto surrounding skin, thus minimizing the risk of maceration. The Safetac® layer ensures the dressing can be changed without damaging the wound or surrounding skin, or exposing the patient to additional pain.

Benefits of Mepilex® XT

- Can be used in all healing stages
- Minimizes pain and trauma during dressing changes
- Minimizes stress caused by pain during dressing changes, to reduce delays in healing
- Minimizes maceration, and effectively manages a wider range of exudate for an optimized wound healing environment
- Stays in place allowing for “hands-free” application to facilitate use of compression or retention bandages²
- Well suited for use under compression bandages
- Can be cut to size if required
- Promotes patient comfort during wear
- Can remain in place for several days depending on the condition of the wound^{4,9}
- Can be lifted and adjusted without losing its adherent properties
- Non-sensitizing¹⁰

References:

1. Fluid handling and retention properties Mepilex XT : Report no. 20130123-006 (SMTL). 2. Fluid handling and retention properties with Viscous test Fluid Mepilex XT, Report No. 20130104-004/ 20121012-004/20130104-004 (MHC). 3. White R. et al. Evidence for atraumatic soft silicone wound dressing use. Wounds UK, 2005. 4. White R. A Multinational survey of the assessment of pain when removing dressings. Wounds UK, 2008. 5. Upton D. et al. The Impact of Atraumatic Vs Conventional Dressings on Pain and Stress in Patients with Chronic Wounds. Journal of Wound Care, 2012. 6. Wiberg A.B. et al. Preventing maceration with a soft silicone dressing: in-vitro evaluations. Poster presented at the 3rd Congress of the WUWHs, Toronto, Canada, 2008. 7. Meuleneire F and Foster A. Local treatment of heel pressure ulcers with a silicone foam dressing. Poster presentation. WUWHs, 2008. 8. External Test Lab Report no. 413098 (Nelson Laboratories). 9. Eager CA. Comparison of two foams through the measurement of healing time, frequency of dressing changes and peri wound status. Poster presentation. Advanced Wound Care and Medical Research Forum on Wound Repair, 2001. 10. Biocompatibility Evaluation, Filed in MHC.

Indications for use

Mepilex® XT is designed for a wide range of exuding acute and chronic wounds in all healing stages.

Precautions

- Do not use on patients with known sensitivity to the dressing or its components.
- In case of signs of clinical infection, consult a health care professional for adequate infection treatment.
- Do not use Mepilex® XT together with oxidizing agents such as hypochlorite solutions or hydrogen peroxide.



Mepilex® XT Assortment (Sterile packed)

Art. no	Size cm	Pcs/Box	Pcs/Case
211015	5 x 5	5	40
211100	10 x 10	5	70
211200	10 x 20	5	45
211300	15 x 15	5	25
211500	20 x 50	2	12

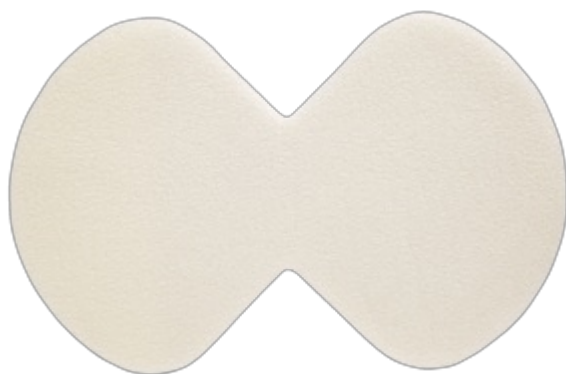
288100 Mepilex Heel

Soft silicone foam dressings

Product details

Size : 13cm x 20cm
Descriptive feature : Foam, Non-border, Soft silicone
Sterile : Sterile
Brand : Mepilex

Images



Delivered items

288100-02

Sales released in: Argentina, Australia, Austria, Belgium, Bolivia (Plurinational State of), Brazil, Canada, Chile, China, Colombia, Czechia, Denmark, Ecuador, Faroe Islands, Finland, Germany, Hong Kong, Hungary, India, Ireland, Italy, Japan, Kazakhstan, Korea (the Republic of), Luxembourg, Malaysia, Mexico, Netherlands, New Zealand, Norway, Peru, Poland, Portugal, Puerto Rico, Russian Federation, Singapore, South Africa, Spain, Sweden, Switzerland, Thailand, United Kingdom of Great Britain and Northern Ireland, United States of America

Country of origin: Finland

Shelf life: 3 years

Sterilization method: EtO

Production Responsibility: Mölnlycke Health Care Oy, PO Box 76, Saimaankatu 6, FI-50101 Mikkeli 10, Finland

Packing information: First packaging layer is a peel open sterile barrier, paper/plastic. Once opened the sterile barrier cannot be closed again. Second layer is a cardboard dispenser box. Third layer is a corrugated board transport box.

Is suitable for Tray: No

Packing level	Quantity	GS1 Code / UDI-DI	Width x Length x Height	Vol	Weight gross / net
Piece	1	7332430680822			

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Packing level	Quantity	GS1 Code / UDI-DI	Width x Length x Height	Vol	Weight gross / net
Consumer pack	1	7332430718785			
Dispenser box	5	7323190271405			
Transport box	25	7323190271399	228x387x230 mm	20.3 dm3	1.4 / 0.4 kg
Pallet	1750	7323190271382	800x1200x1872 mm		

Material

Animal tissues :	No
Human blood derivatives :	No
Natural rubber latex :	No
Medicinal substances :	No
Phthalates :	No
Polyvinyl chloride :	No

Product Composition Non-bordered Foam Products

Product Component	Composition
Backing material	Polyurethane film
Wound pad	Polyurethane foam
Wound contact layer	Silicone
Protective release liner	Polyethylene film

Product Performance Non-bordered Foam Products

Characteristics	Test Method	Internal Test Method	Unit	Requirement	Product Performance
Free Swell Absorptive Capacity	EN 13726-1 part 3:2	T-1069	g/100 cm ²	Not specified	77.9
Free Swell Absorptive Capacity	EN 13726-1 part 3:2	T-1069	g/g	Not specified	12.5
Fluid Handling Capacity	EN 13726-1 part 3:3	T-1068	g/10 cm ² /24 h	Not specified	33.07
Absorbency	EN 13726-1 part 3:3	T-1068	g/10 cm ² /24 h	Not specified	6.81
Moisture Vapour Transmission Rate (MVTR)	EN 13726-1 part 3:3	T-1068	g/10 cm ² /24 h	Not specified	26.26

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Characteristics	Test Method	Internal Test Method	Unit	Requirement	Product Performance
Waterproofness	EN 13726-3	T-1083	Pass/Fail	>500 mm H ₂ O for 300 s	Pass
Conformability-Extensibility, MD	EN 13726-4	T-1086	N/cm	Not specified	1.3
Conformability-Extensibility, CD	EN 13726-4	T-1086	N/cm	Not specified	1.0
Conformability-Permanent Set, MD	EN 13726-4	T-1086	%	Not specified	0.4
Conformability-Permanent Set, CD	EN 13726-4	T-1086	%	Not specified	0.3

Technical

Dimension

Dimension text	Dimension value
Product	13 cm x 20 cm
Product	5 in x 8 in

Classifications

Regulation type(s)	MDD Class IIb	CFR Class I	Locally Regulated	Locally Regulated
CE Certificate Number :	CE 01965			

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Regulation type(s)	MDD Class IIb	CFR Class I	Locally Regulated	Locally Regulated
Intended use MDD :	Mepilex is intended for a wide range of exuding wounds such as leg and foot ulcers, pressure ulcers and traumatic wounds, e.g. skin tears and secondary healing wounds. Mepilex can be used as a protection of compromised and/or fragile skin and may also be used as part of a prophylactic therapy to help prevent skin damage, e.g. pressure ulcers.			
Intended Purpose :			Mepilex is intended for a wide range of exuding wounds such as leg and foot ulcers, pressure ulcers and traumatic wounds, e.g. skin tears and secondary healing wounds. Mepilex can be used as a protection of compromised and/or fragile skin and may also be used as part of a prophylactic therapy to help prevent skin damage, e.g. pressure ulcers.	Mepilex is intended for a wide range of exuding wounds such as leg and foot ulcers, pressure ulcers and traumatic wounds, e.g. skin tears and secondary healing wounds. Mepilex can be used as a protection of compromised and/or fragile skin and may also be used as part of a prophylactic therapy to help prevent skin damage, e.g. pressure ulcers.
Conformity Annexes :	II			
Measuring Function :	No			

Find out more at www.molnlycke.com

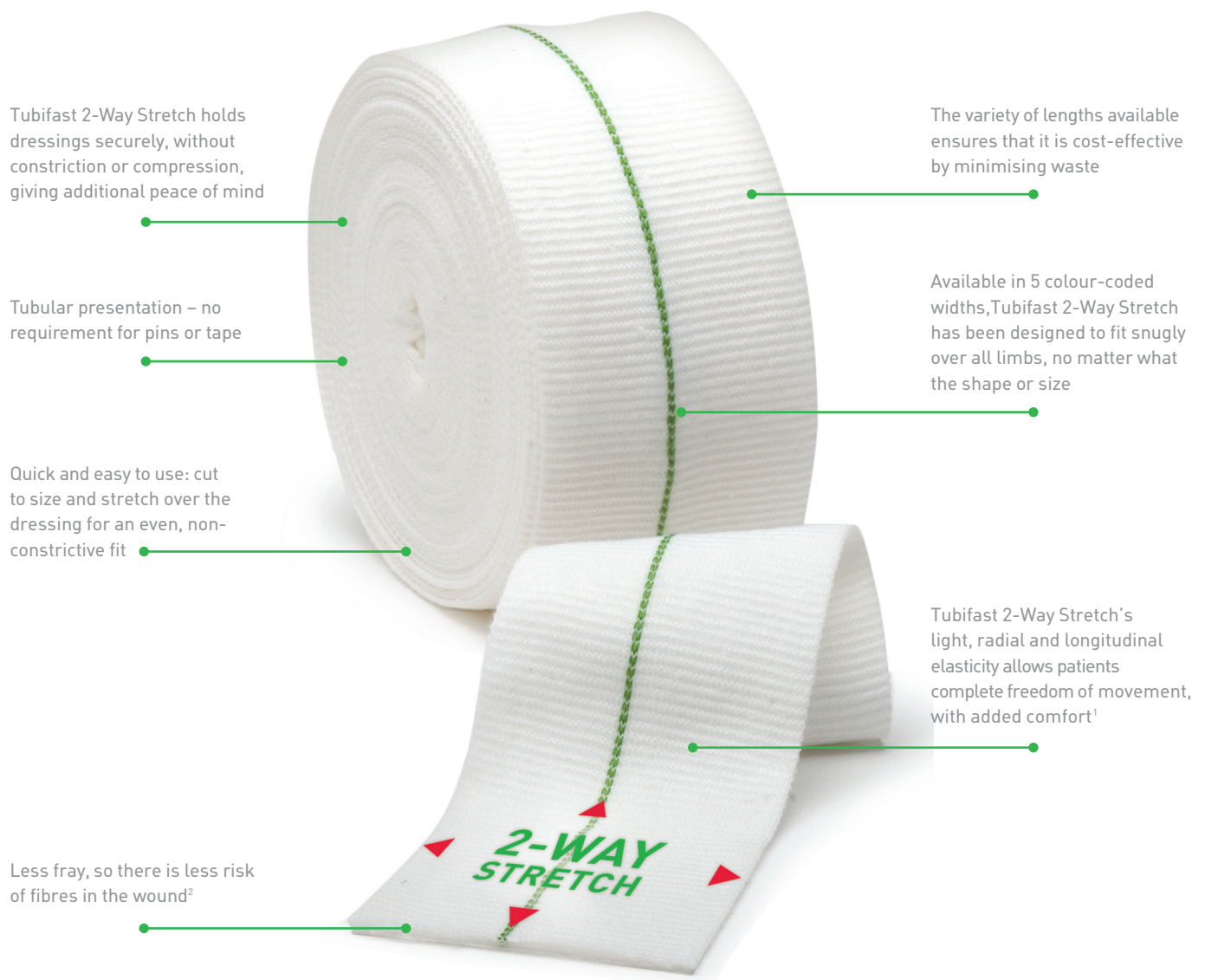
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Tubifast[®] WITH **2-WAY STRETCH**[™] TECHNOLOGY

Lightweight, tubular dressing retention

New Tubifast 2-Way Stretch is the world's first and only lightweight elasticated tubular bandage with radial and longitudinal stretch.¹ This unique mode of action allows patients complete freedom of movement, with added comfort. Ideal for dressing retention and skin covering for any part of the body. It can also be used for patch wrapping and as an under-cast stockinette. Tubifast 2-Way Stretch holds dressings securely in place without the need for tapes, pins or ties. It is quick and easy to use, with less risk of fraying² than traditional tubular bandages.



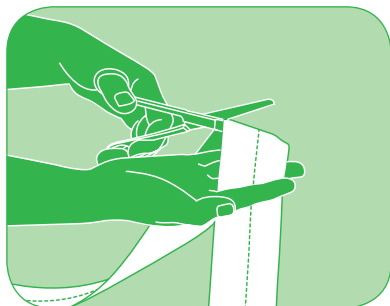
How Tubifast 2-Way Stretch works

New Tubifast 2-Way Stretch is manufactured from viscose with very fine elastane threads knitted into the fabric radially and longitudinally, to provide light elasticity.

Tubifast 2-Way Stretch holds dressings securely, without constriction or compression. No pins or tapes are necessary. Its two-way stretch allows patients complete freedom of movement and added comfort.

Tubifast 2-Way Stretch is available in a range of five widths, and has been designed to fit snugly over all limbs, no matter what the shape or size.

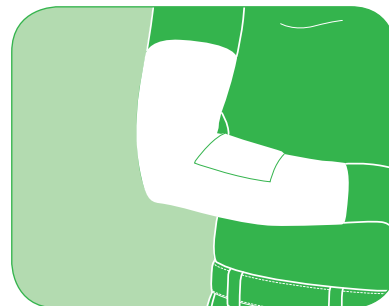
How to use Tubifast 2-Way Stretch



1. Choose the correct width bandage, using the quick reference colour coding, and cut to length.



2. Stretch Tubifast 2-Way Stretch over the affected area.



3. Position Tubifast 2-Way Stretch to cover the dressing. Illustrated, step-by-step instructions are available for more advanced applications.

Benefits of Tubifast 2-Way Stretch

- Tubifast 2-Way Stretch holds dressings securely, without constriction or compression
- Its radial and longitudinal stretch allows patients complete freedom of movement, with added comfort¹
- A variety of lengths are available, ensuring cost- effectiveness by minimising waste
- Quick and easy to use: simply cut to size and stretch over the dressing for an even, non-constrictive fit
- Available in a range of quick reference, colour-coded sizes, to fit everything from small limbs to adult trunks
- It has less fray so there is less risk of fibres in the wound²

Tubifast 2-Way Stretch Ordering information

Product	Width x length	Limb	Art. No.
RED LINE small limbs	3.5 cm x 10 m	8-15 cm	2434
GREEN LINE small and medium limbs	5 cm x 10 m	10-25 cm	2436
BLUE LINE large limbs	7.5 cm x 10 m	20-45 cm	2438
YELLOW LINE extra large limbs, heads, children's trunks	10.75 cm x 10 m	35-65 cm	2440
PURPLE LINE adult trunks	20 cm x 10 m	60-130 cm	2444

Fibre content

Viscose, Elastane and Polyamide

Washing instructions



Natural rubber latex is not a constituent of Tubifast 2-Way Stretch or its packaging



¹ Published data shows that Tubifast 2-Way Stretch exceeds the minimum requirement of longitudinal stretch for the actions of sitting, bending and flexing. Physical testing of textiles, B. P. Saville, Woodhead Publishing, 1999.

² Data on file.

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NSW 2086, Tel. 1800 005 231. www.molnlycke.com.au
New Zealand Orders & Enquiries 0800 005 231, www.molnlycke.co.nz

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AU012370903

2434 Tubifast

Elasticated viscose tubular bandage

Product details

Size :	Red
Descriptive feature :	Tubular bandage
Sterile :	Non-sterile
Brand :	Tubifast garments

Images



Delivered items

2434-03

Sales released in: Algeria, Argentina, Australia, Austria, Azerbaijan, Belgium, Bosnia and Herzegovina, Brazil, Bulgaria, Canada, Chile, China, Colombia, Croatia, Cuba, Cyprus, Czechia, Denmark, Estonia, Faroe Islands, Finland, France, French Guiana, Germany, Greece, Guadeloupe, Hong Kong, Hungary, Iceland, Indonesia, Ireland, Israel, Italy, Japan, Kazakhstan, Korea (the Republic of), Kuwait, Latvia, Lithuania, Luxembourg, Malaysia, Martinique, Mexico, Netherlands, New Zealand, Norway, Oman, Panama, Poland, Portugal, Puerto Rico, Qatar, Romania, Russian Federation, Saudi Arabia, Serbia, Singapore, Slovenia, Spain, Sweden, Switzerland, Taiwan (Province of China), Thailand, Turkey, Ukraine, United Arab Emirates, United Kingdom of Great Britain and Northern Ireland, United States of America, Uzbekistan, Viet Nam

Country of origin: United Kingdom of Great Britain and Northern Ireland

Shelf life: 5 years

Sterilization method: Non-sterile

Production Responsibility: Mölnlycke Health Care Ltd, Tubiton House, Medlock Street, Oldham, OL1 3HS, United Kingdom

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Packing information: First packaging layer is a cardboard dispenser box. Second layer is a corrugated board transport box.

Is suitable for Tray: No

Packing level	Quantity	GS1 Code / UDI-DI	Width x Length x Height	Vol	Weight gross / net
Piece	1	7332430888129			
Consumer pack	1	5055158008649			
Dispenser box	1	5055158008656			
Transport box	20	7332551927158	195x395x238 mm	18.3 dm3	2.7 / 2.1 kg
Pallet	1680	7332551927141	800x1200x1816 mm		

Material

Animal tissues :	No
Human blood derivatives :	No
Natural rubber latex :	No
Medicinal substances :	No
Phthalates :	No
Polyvinyl chloride :	No

Product Composition Compression and Retention Products, Tubigrip and TSSB

Product Component	Composition
Fabric	Viscose, Elastane

Product Performance Compression and Retention Products

Characteristics	Internal Test Method	Unit	Requirement	Product Performance
Extensibility Length	SM83	%	Not specified	35<x<65
Extensibility Width	SM2244	%	Not specified	x>300

Technical

Dimension

Dimension text	Dimension value
Limb measurement	9 cm - 18 cm

Find out more at www.molnlycke.com

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2436 Tubifast

Elasticated viscose tubular bandage

Product details

Size :	Green
Descriptive feature :	Tubular bandage
Sterile :	Non-sterile
Brand :	Tubifast garments

Images



Delivered items

2436-03

Sales released in: Algeria, Andorra, Argentina, Australia, Austria, Azerbaijan, Belgium, Bosnia and Herzegovina, Brazil, Bulgaria, Canada, Chile, China, Colombia, Croatia, Cyprus, Czechia, Denmark, Estonia, Faroe Islands, Finland, France, French Guiana, Germany, Greece, Hong Kong, Hungary, Iceland, Indonesia, Ireland, Israel, Italy, Japan, Kazakhstan, Korea (the Republic of), Kuwait, Latvia, Lithuania, Luxembourg, Malaysia, Mexico, Morocco, Netherlands, New Zealand, Norway, Oman, Panama, Poland, Portugal, Qatar, Romania, Russian Federation, Saudi Arabia, Serbia, Singapore, Slovenia, Spain, Sweden, Switzerland, Taiwan (Province of China), Thailand, Tunisia, Turkey, Ukraine, United Arab Emirates, United Kingdom of Great Britain and Northern Ireland, United States of America, Uzbekistan, Viet Nam

Country of origin: United Kingdom of Great Britain and Northern Ireland

Shelf life: 5 years

Sterilization method: Non-sterile

Production Responsibility: Mölnlycke Health Care Ltd, Tubiton House, Medlock Street, Oldham, OL1 3HS, United Kingdom

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Packing information: First packaging layer is a cardboard dispenser box. Second layer is a corrugated board transport box.

Is suitable for Tray: No

Packing level	Quantity	GS1 Code / UDI-DI	Width x Length x Height	Vol	Weight gross / net
Piece	1	7332430888112			
Consumer pack	1	5055158008687			
Dispenser box	1	5055158008694			
Transport box	24	7332551933821	295x395x246 mm	28.7 dm3	5.2 / 4.2 kg
Pallet	1344	7332551933814	800x1200x1872 mm		

Material

Animal tissues :	No
Human blood derivatives :	No
Natural rubber latex :	No
Medicinal substances :	No
Phthalates :	No
Polyvinyl chloride :	No

Product Composition Compression and Retention Products, Tubigrip and TSSB

Product Component	Composition
Fabric	Viscose, Elastane

Product Performance Compression and Retention Products

Characteristics	Internal Test Method	Unit	Requirement	Product Performance
Extensibility Length	SM83	%	Not specified	35<x<65
Extensibility Width	SM2244	%	Not specified	x>300

Technical

Dimension

Dimension text	Dimension value
Limb measurement	14 cm - 24 cm

Find out more at www.molnlycke.com

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2438 Tubifast

Elasticated viscose tubular bandage

Product details

Size :	Blue
Descriptive feature :	Tubular bandage
Sterile :	Non-sterile
Brand :	Tubifast garments

Images



Delivered items

2438-03

Sales released in: Algeria, Andorra, Argentina, Australia, Austria, Azerbaijan, Belgium, Bosnia and Herzegovina, Brazil, Brunei Darussalam, Bulgaria, Canada, Chile, China, Colombia, Croatia, Cuba, Cyprus, Czechia, Denmark, Estonia, Faroe Islands, Finland, France, Germany, Greece, Hong Kong, Hungary, Iceland, Indonesia, Iraq, Ireland, Israel, Italy, Japan, Kazakhstan, Korea (the Republic of), Kuwait, Latvia, Lithuania, Luxembourg, Malaysia, Mexico, Netherlands, New Zealand, Norway, Oman, Panama, Poland, Portugal, Qatar, Romania, Russian Federation, Saudi Arabia, Serbia, Singapore, Slovenia, Spain, Sweden, Switzerland, Taiwan (Province of China), Thailand, Turkey, Ukraine, United Arab Emirates, United Kingdom of Great Britain and Northern Ireland, United States of America, Uzbekistan, Viet Nam

Country of origin: United Kingdom of Great Britain and Northern Ireland

Shelf life: 5 years

Sterilization method: Non-sterile

Production Responsibility: Mölnlycke Health Care Ltd, Tubiton House, Medlock Street, Oldham, OL1 3HS, United Kingdom

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Packing information: First packaging layer is a cardboard dispenser box. Second layer is a corrugated board transport box.

Is suitable for Tray: No

Packing level	Quantity	GS1 Code / UDI-DI	Width x Length x Height	Vol	Weight gross / net
Piece	1	7332430888105			
Consumer pack	1	5055158008724			
Dispenser box	1	5055158008731			
Transport box	30	7332551935511	266x528x356 mm	50.0 dm3	9.1 / 7.5 kg
Pallet	720	7332551935504	800x1200x1574 mm		

Material

Animal tissues :	No
Human blood derivatives :	No
Natural rubber latex :	No
Medicinal substances :	No
Phthalates :	No
Polyvinyl chloride :	No

Product Composition Compression and Retention Products, Tubigrip and TSSB

Product Component	Composition
Fabric	Viscose, Elastane

Product Performance Compression and Retention Products

Characteristics	Internal Test Method	Unit	Requirement	Product Performance
Extensibility Length	SM83	%	Not specified	35<x<65
Extensibility Width	SM2244	%	Not specified	x>300

Technical

Dimension

Dimension text	Dimension value
Limb measurement	24 cm - 40 cm

Find out more at www.molnlycke.com

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2440 Tubifast

Elasticated viscose tubular bandage

Product details

Size :	Yellow
Descriptive feature :	Tubular bandage
Sterile :	Non-sterile
Brand :	Tubifast garments

Images



Delivered items

2440-03

Sales released in: Algeria, Andorra, Argentina, Australia, Austria, Azerbaijan, Belgium, Bosnia and Herzegovina, Brazil, Bulgaria, Canada, Chile, China, Colombia, Croatia, Cuba, Cyprus, Czechia, Denmark, Estonia, Faroe Islands, Finland, France, French Guiana, Germany, Greece, Hong Kong, Hungary, Iceland, Indonesia, Iraq, Ireland, Israel, Italy, Japan, Kazakhstan, Korea (the Republic of), Latvia, Lithuania, Luxembourg, Malaysia, Mexico, Netherlands, New Zealand, Norway, Oman, Panama, Poland, Portugal, Qatar, Romania, Russian Federation, Saudi Arabia, Serbia, Singapore, Slovenia, Spain, Sweden, Switzerland, Taiwan (Province of China), Thailand, Turkey, Ukraine, United Arab Emirates, United Kingdom of Great Britain and Northern Ireland, United States of America, Uzbekistan, Viet Nam

Country of origin: United Kingdom of Great Britain and Northern Ireland

Shelf life: 5 years

Sterilization method: Non-sterile

Production Responsibility: Mölnlycke Health Care Ltd, Tubiton House, Medlock Street, Oldham, OL1 3HS, United Kingdom

Packing information: First packaging layer is a cardboard dispenser box. Second layer is a corrugated board

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transport box.

Is suitable for Tray: No

Packing level	Quantity	GS1 Code / UDI-DI	Width x Length x Height	Vol	Weight gross / net
Piece	1	7332430888099			
Consumer pack	1	5055158008762			
Dispenser box	1	5055158008779			
Transport box	12	7332551935542	295x395x246 mm	28.7 dm3	5.4 / 4.5 kg
Pallet	672	7332551935535	800x1200x1872 mm		

Material

Animal tissues :	No
Human blood derivatives :	No
Natural rubber latex :	No
Medicinal substances :	No
Phthalates :	No
Polyvinyl chloride :	No

Product Composition Compression and Retention Products, Tubigrip and TSSB

Product Component	Composition
Fabric	Viscose, Elastane

Product Performance Compression and Retention Products

Characteristics	Internal Test Method	Unit	Requirement	Product Performance
Extensibility Length	SM83	%	Not specified	35<x<65
Extensibility Width	SM2244	%	Not specified	x>300

Technical

Dimension

Dimension text	Dimension value
Limb measurement	35 cm - 64 cm
Limb measurement	14 in - 25 in

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2444 Tubifast

Elasticated viscose tubular bandage

Product details

Size :	Purple
Descriptive feature :	Tubular bandage
Sterile :	Non-sterile
Brand :	Tubifast garments

Images



Delivered items

2444-03

Sales released in: Andorra, Argentina, Australia, Austria, Azerbaijan, Belgium, Bosnia and Herzegovina, Brazil, Bulgaria, Canada, Chile, China, Colombia, Croatia, Cyprus, Czechia, Denmark, Estonia, Faroe Islands, Finland, France, Germany, Greece, Hong Kong, Hungary, Iceland, Indonesia, Ireland, Israel, Italy, Kazakhstan, Korea (the Republic of), Latvia, Lithuania, Luxembourg, Malaysia, Mexico, Netherlands, New Zealand, Norway, Oman, Panama, Poland, Portugal, Qatar, Romania, Russian Federation, Saudi Arabia, Serbia, Singapore, Slovenia, Spain, Sweden, Switzerland, Taiwan (Province of China), Thailand, Turkey, Ukraine, United Arab Emirates, United Kingdom of Great Britain and Northern Ireland, United States of America, Uzbekistan, Viet Nam

Country of origin: United Kingdom of Great Britain and Northern Ireland

Shelf life: 5 years

Sterilization method: Non-sterile

Production Responsibility: Mölnlycke Health Care Ltd, Tubiton House, Medlock Street, Oldham, OL1 3HS, United Kingdom

Packing information: First packaging layer is a cardboard dispenser box. Second layer is a corrugated board

Find out more at www.molnlycke.com

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transport box.

Is suitable for Tray: No

Packing level	Quantity	GS1 Code / UDI-DI	Width x Length x Height	Vol	Weight gross / net
Piece	1	7332430888075			
Consumer pack	1	5055158008847			
Dispenser box	1	5055158008854			
Transport box	15	7332551935573	395x590x268 mm	62.5 dm3	10.5 / 8.8 kg
Pallet	360	7332551935566	800x1200x1758 mm		

Material

Animal tissues :	No
Human blood derivatives :	No
Natural rubber latex :	No
Medicinal substances :	No
Phthalates :	No
Polyvinyl chloride :	No

Product Composition Compression and Retention Products, Tubigrip and TSSB

Product Component	Composition
Fabric	Viscose, Elastane

Product Performance Compression and Retention Products

Characteristics	Internal Test Method	Unit	Requirement	Product Performance
Extensibility Length	SM83	%	Not specified	35<x<65
Extensibility Width	SM2244	%	Not specified	x>300

Technical

Dimension

Dimension text	Dimension value
Limb measurement	64 cm - 130 cm
Limb measurement	25 in - 51 in

Find out more at www.molnlycke.com

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