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MANUALE D'USO – USER MANUAL – MODE D'EMPLOI – MANUAL DE USO - BENUTZERHANDBUCH - MANUAL DO USUÁRIO - ΕΓΧΕΙΡΙΔΙΟ ΧΡΗΣΗΣ
 - РЪКОВОДСТВО ЗА ПОТРЕБИТЕЛ - UŽIVATELSKÁ PŘÍRUČKA - BRUGERVEJLEDNING - KASUTUSJUHEND - KÄYTTÖOHJE - PRIRUČNIK ZA UPOTREBU - FELHASZNÁLÓI ÚTMUTATÓ - NAUDOJIMO VADOVAS - LIETOJĀJA ROKASGRĀMATA - GEBRUIKERSHANDLEIDING - INSTRUKCJA OBSŁUGI - MANUAL DE UTILIZARE - POUŽIVATELSKÁ PŘÍRUČKA - PRIROČNIK ZA ANVÄNDARHANDBOK UPORABO - دليل المستخدم

- È necessario segnalare qualsiasi incidente grave verificatosi in relazione al dispositivo medico da noi fornito al fabbricante e all'autorità competente dello Stato membro in cui si ha sede.
- All serious accidents concerning the medical device supplied by us must be reported to the manufacturer and competent authority of the member state where your registered office is located.
- Il est nécessaire de signaler tout accident grave survenu et lié au dispositif médical que nous avons livré au fabricant et à l'autorité compétente de l'état membre où on a le siège social.
- Es necesario informar al fabricante y a la autoridad competente del Estado miembro en el que se encuentra la sede sobre cualquier incidente grave que haya ocurrido en relación con el producto sanitario que le hemos suministrado.
- Jeder schwere Unfall im Zusammenhang mit dem von uns gelieferten medizinischen Gerät muss unbedingt dem Hersteller und der zuständigen Behörde des Mitgliedsstaats, in dem das Gerät verwendet wird, gemeldet werden.
- É necessário notificar ao fabricante e às autoridades competentes do Estado-membro onde ele está sediado qualquer acidente grave verificado em relação ao dispositivo médico fornecido por nós.
- Σε περίπτωση που διαπιστώσετε οποιοδήποτε σοβαρό περιστατικό σε σχέση με την ιατρική συσκευή που σας παρέχουμε θα πρέπει να το αναφέρετε στον κατασκευαστή και στην αρμόδια αρχή του κράτους μέλους στο οποίο βρίσκεται.
- Всички сериозни инциденти, които са настъпили във връзка с доставеното от нас медицинско изделие, трябва да се сигнализират на производителя и на компетентния орган на държавата членка, в която производителят е установен.
- Je třeba nahlásit jakoukoli vážnou nehodu, ke které došlo v souvislosti s námi dodávaným zdravotnickým prostředkem, výrobci a příslušnému orgánu členského státu, ve kterém máte sídlo.
- Du skal rapportere enhver alvorlig hændelse, der opstår i forbindelse med det medicinske udstyr leveret af os, til producenten og til den kompetente myndighed i den medlemsstat, hvor du befinder dig.
- Peate teatama kõigist meie poolt tarnitud meditsiiniseadmega seotud tõsistest vahejuhtumitest tootjale ja selle liikmesriigi pädevale asutusele, kus te asute.
- Kaikista vakavista tapaturmista, jotka liittyvät toimittamamme lääkinnällisen laitteen käyttöön, on ilmoitettava valmistajalle sekä oman asuinpaikan jäsenmaan toimivaltaiselle viranomaiselle.
- Potrebno je prijaviti svaku ozbiljnu nezgodu koja se dogodila u vezi s isporučenim medicinskim proizvodaču i nadležnom tijelu države članice u kojoj se nalazi.
- A gyártónak, illetve a székhely szerinti tagállam illetékes hatóságának jelezni kell bármilyen olyan súlyos balesetet, amely az általunk szállított orvostechnikai eszközzel kapcsolatban történt.
- Privalote pranešti gamintojui ir valstybės narės, kurioje esate įsisteigęs, kompetentingai institucijai apie bet koki rimtą incidentą, įvykusį dėl mūsų tiekiamų medicinios prietaisų.
- Par nopietnu negadījumu, kas notiek saistībā ar mūsu piegādāto medicīnisko ierīci, jāziņo ražotājam un tās dalībvalsts kompetentā iestādei, kurā negadījums ir radies.
- Alle ernstige ongelukken die zich in verband met het door ons geleverde medische hulpmiddel voordoen, moeten gemeld worden aan de fabrikant en aan de bevoegde instantie van de lidstaat waar u gevestigd bent.
- Należy poinformować producenta i kompetentne władze danego Kraju członkowskiego o każdym poważnym wypadku związanym z wyrobem medycznym naszej produkcji.
- Orice accident grav produs, privitor la dispozitivul medical fabricat de firma noastră, trebuie semnalat producătorului și autorității competente în statul membru pe teritoriul căruia își are sediul utilizatorul.
- Každú vážnu udalosť, ktorá sa vyskytla v súvislosti s nami dodanou zdravotníckou pomôckou, je potrebné nahlásiť výrobcovi a príslušnému orgánu členského štátu, v ktorom máte sídlo.
- O vsakršni hujši nesreči, do katere bi prišlo v povezavi z medicinskim pripomočkom, ki smo vam ga dobavili, je treba obvestiti pristojne oblasti v državi v kateri imate sedež.
- Det är nödvändigt att meddela tillverkaren och de behöriga myndigheterna i den berörda medlemsstaten, om alla allvariga olyckor som inträffat i samband med den medicintekniska utrustning som levererats av oss.

يجب الإبلاغ فوراً عن أي حادث خطير وقع فيما يتعلق بالجهاز الطبي الذي زودنا به إلى الجهة الصانعة والسلطة المختصة في الدولة العضو التي يقع فيها.



Jiangsu Saikang Medical Equipment Co., Ltd
 No. 35 Lehong Road, Modern Agriculture Demonstration Park,
 Zhangjiagang City, Jiangsu Province, China
 Made in China



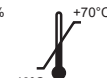
X36-1



SUNGO Europe B.V.
 Fascinatio Boulevard 522, Unit 1.7,
 2909VA Capelle aan den IJssel, The Netherlands



Gima S.p.A.
 Via Marconi, 1 - 20060 Gessate (MI) Italy
 gima@gimaitaly.com - export@gimaitaly.com
 www.gimaitaly.com



ENGLISH

INTENDED USE

An examination table is integral to patient comfort while suffering from injury, trauma or shock.

WARNING

Any serious incident that has occurred in relation to the device shall be reported to the manufacturer and the competent authority of the Member State in which the user and/or the patient is established.

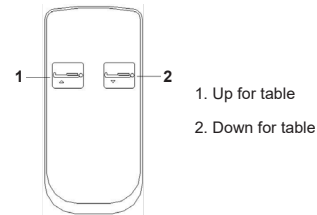
1. Use method



Table Size	Length	mm	1960±10
	Width		680±10
	The height of the table above the ground		470-910 ±10
Weight	Maximum patient weight	kg	180
Weight	Safe working load	kg	200

1.1 Height Adjustment

The whole examination table can be controlled by hand operator



2. Electric examination table-maintenance

1. In order to use the examination table safely, it is necessary to perform regular safety inspections on the examination table. It is recommended to check it every six months to ensure that the connections are not loose and all operations are normal.

2. When the examination table frame ages and it reaches a certain servicing time, which the metal part of the examination table and plastic parts of the stainless steel basin frame can be recycle.

3. Avoid scratching the panel with sharp-angle appliances or knives during use, and clean it frequently to keep it clean and dry.

4. If the panel is accidentally stained with stains, it is recommended to clean it. Do not use alkaline or corrosive chemicals to clean the examination table, which will cause the stainless steel surface to rust.

3. Examination table-cleaning

1. When cleaning, please wring out the cloth soaked in neutral detergent diluted with water and wipe it, then wring out the cloth soaked in water to wipe off the remaining detergent ingredients, and finally use a dry wipe dry with a cloth.

2. Do not use volatile substances (thinner, volatile agent, gasoline, etc.), which may cause chemical reactions and damage the examination table.

3. When using a disinfectant to clean, be sure to dilute it according to its specified concentration before use. Depending on the composition of the disinfectant, it may corrode metal parts, resin parts, etc., causing undesirable phenomena such as discoloration and deformation. Therefore, the content of the disinfectant is recommended as follows:

0.05 - 0.2% ammonium chloride disinfectant

0.05 - 0.2% Chlorinated phenyl disinfectant 0.05% diclofenac ethane solution 0.05 - 0.2% sodium hypochlorite disinfectant

Do not use smoking sterilizers, autoclaves, and do not use methyl (phenol) to clean the headboard and foot of the examination table. It may cause corrosion, discoloration and deterioration.

Note: When cleaning the examination table or changing the examination table clothes, please pay attention to the corners, edges and screws of the frame to prevent scratches.

4. Packaging, transport, storage

1. The packaging of examination table is carried out according to the contract or product standard.

2. Avoid rushing out, violent vibration, and protection from sunlight and rain during the transportation of the examination table.

3. The examination table should be stored:

a) Ambient temperature: -40°C ~ 70°C.

b) Relative humidity: ≤ 95%.

c) Atmospheric pressure: 500hPa ~ 1060 hPa.

5. After-sales service

1. Please keep the files that comes with the machine and invoices of this product properly, and you need to present these files when the company performs warranty and maintenance for the product.

2. If there is any problem in the process of using, please contact our company in time, so that our company can provide you with accurate and fast technical support and maintenance services in a timely manner.

3. From the date of sale, if the product is broken or damaged due to the correct installation and use according to the specification, the product will enjoy one-year free warranty and lifelong maintenance service with the "certificate" or invoice.

4. From the date of purchase, if it is indeed damaged or does not work normally due to quality problems within one year, the company will provide to the users a free repairing of the product.

5. Lifetime service from manufacturer: JIANGSU SAIKANG MEDICAL EQUIPMENT CO LTD.

EMC

1) This product needs special precautions regarding EMC and needs to be installed and put into service according to the EMC information provided, and this unit can be affected by portable and mobile RF communications equipment.

2) * Do not use a mobile phone or other devices that emit electromagnetic fields, near the unit. This may result in incorrect operation of the unit.

3) Caution: This unit has been thoroughly tested and inspected to assure proper performance and operation!

4) * Caution: This machine should not be used adjacent to or stacked with other equipment and that if adjacent or stacked use is necessary, this machine should be observed to verify normal operation in the configuration in which it will be used.

Guidance and Manufacturer's Declaration – Electromagnetic Immunity		
The V8v8c is intended for use in the following electromagnetic environment. The customer or user of the V8v8c should make sure it is used in such an environment		
Emission test	Compliance	Electromagnetic Environment Guidance

RF Emission to CISPR 11	Group 1	The bed uses RF energy for its internal function
RF Emission to CISPR 11	Class B	The bed is intended for use in all types of establishment including residential and similar user that are directly connected to a public supply network that also serves buildings used for residential purposes
Harmonics according to IEC 61000-3-2	Class A	
Voltage fluctuations/ flicker acc. to IEC 61000-3-3	Complies	

Guidance and Manufacturer's Declaration - Electromagnetic Immunity		
The V8v8c is intended for use within healthcare facility environment. Users or customers must use bed in this environment		
Immunity test	IEC 60601-1-2 EN 60601-1-2 Test level	Electromagnetic Environment Guidance
Electrostatic discharge (ESD)..... IEC 61000-4-2 /EN 61000-4-2	Contact discharge: +8kV Air discharge: +2kV, +4kV, +8kV, +15kV	Contact discharge: +8kV Air discharge: +2kV, +4kV, +8kV, +15kV Floor should be made of wood and concrete or be tiled with ceramic tiles. If the floor is covered with synthetic flooring material, the relative air humidity must be at least 30% Can be used when higher ESD levels are present
Electrical fast transient/burst IEC 61000-4-4 /EN 61000-4-4	+2kV Power supply line +1kV Input/Output lines	+2kV Power supply line The quality of the supply voltage should be equivalent to that of a typical business or hospital environment.
Surge IEC 61000-4-5 /EN 61000-4-5	Line - line: +0.5kV, +1.0kV Line - ground: +0.5kV, +1.0kV, +2.0kV	Line - line: +0.5kV, +1.0kV Line - ground: +0.5kV, +1.0kV, +2.0kV The quality of the supply voltage should be equivalent to that of a typical business or hospital environment.
Power Frequency Magnetic Fields IEC 61000-4-8 /EN 61000-4-8	30 A/m 50Hz/60Hz	30 A/m 50Hz/60Hz Magnetic fields with a network frequency should be equivalent to those to be found in a typical business or hospital environment.
Voltage Dips IEC 61000-4- 11 /EN 61000-4-11	0% UT: 0.5 cycles Phase: 00, 450, 900, 1350, 1800, 2250, 2700, 3150	0% UT: 0.5 cycles Phase: 00, 450, 900, 1350, 1800, 2250,
	0% UT: 1 cycle 70% UT: 25/30 cycles Single phase: 00	0% UT: 1 cycle 70% UT: 25/30 cycles Single phase: 00 0% UT ; 300 cycle The quality of the supply voltage should be equivalent to that of a typical business or hospital environment.
	0% UT ; 300 cycle	If the person using the bed requires that the bed functions must continue despite any interruptions in the energy supply, it is recommended that the bed be connected to an uninterruptible electricity supply or a battery
Short interruptions IEC 61000-4-11 /EN 61000-4-11	0% UT: 250/300 cycles	0% UT: 250/300 cycles The quality of the supply voltage should be equivalent to that of a typical business or hospital environment. If the person using the bed requires that the bed functions must continue despite any interruptions in the energy supply, it is recommended that the bed be connected to an uninterruptible electricity supply or a battery

Note: UT is the AC Power voltage before applying in the test level

Guidance and Manufacturer's Declaration - Electromagnetic Immunity							
The V8v8c is intended for use in an electromagnetic environment that used wireless RF transceiver. Electromagnetic interference can be prevented by maintaining the minimum distance between RF transceiver and V8v8c depend on the maximum output and frequency of the communication device as recommended below.							
Immunity test	IEC 60601-1-2 /EN 60601-1-2 Test level	Compliance level (V/m)	Band (MHz)	Service	Modulation	Maximum power (W)	Distance (m)

Enclosure port to RF wireless communication equipment IEC 61000-4-3 /EN 61000-4-3	27V/m 385MHz	27	380-390	TETRA 400			
	28V/m 450MHz	28	430-470	GMRS 460, FRS 460			
	9V/m 710MHz	9	704-787	LTE Band 13, 17			
	9V/m 745MHz						
	9V/m 780MHz						
	28V/m 810MHz	28	800-960	GSM 800/900, TETRA 800, iDen 820, CDMA 850 LTE Band 5			
	28V/m 870MHz						
	28V/m 930MHz						
	28V/m 1720MHz	28	1700- 1990	GSM 1800; CDMA 1900; GSM 1900; DECT; LTE Band 1,3, 4,25; UMT S			
	28V/m 1845MHz						
	28V/m 1970MHz						
	28V/m 2450MHz	28	2400- 2570	Bluetooth, WLAN, 802.11 b/g/n RFID 2450, LTE band 7			
	9V/m 5240MHz	9	5100- 5800	WLAN 802.11 a/n			
	9V/m 5500MHz						
	9V/m 5785MHz						

Guidance and Manufacturer's Declaration - Electromagnetic Immunity			
The V8v8c is intended for use in the following electromagnetic environment. The customer or user of the V8v8c should make sure it is used in such an environment.			
Immunity test	IEC 60601-1-2 /EN 60601-1-2 Test level	Compliance level	Electromagnetic Environment - Guidance
RF Conductivity IEC 61000-4-6 EN 61000-4-6	3V 0.15MHz-80MHz	3V 0.15MHz-80MHz	Field intensity from the fixed RF transmitter determined by a field investigation of the magnetic field must be lower than the compliance level of the respective frequency band. Near devices with the following symbol, interference could occur. 
	6V in ISM bands between 0.15MHz- 80MHz 80%AM at 1kHz	6V in ISM bands between 0.15MHz- 80MHz 80%AM at 1kHz	
Radioactive RF electromagnetic field IEC 61000-4-3 /EN 61000-4-3	3V/M 80MHz-2.7GHz 80%AM at 1kHz	3V/M 80MHz-2.7GHz 80%AM at 1kHz	

a. The intensity of a magnetic field from fixed transmitters such as a radio (mobile/radio) telephone base station, land mobile radio, amateur radio, AM and FM radio broadcasting, and TV broadcasting cannot be theoretically predicted accurately. To determine the electromagnetic environment for a fixed RF transmitter, consider investigating the electromagnetic field in the location where the bed is used. If the intensity of the magnetic field in the location where the V8v8c is used exceeds the above RF compliance level, make sure to monitor if the V8v8c operates appropriately. If abnormal movement is found, take additional measures as needed, such as changing the direction or position of the V8v8c.

b. These guidelines may not apply in all situations. Electromagnetic propagation is affected by absorption and reflection from structures, objects, and people.

Disposal: The product must not be disposed of along with other domestic waste. The users must dispose of this equipment by bringing it to a specific recycling point for electric and electronic equipment.

GIMA WARRANTY TERMS

THE GIMA 12-MONTH STANDARD B2B WARRANTY APPLIES.

Simboli - Symbols - Symboles - Symbole - Símbolos - Símbolos - Σύμβολα - Символи - Symboly - Symboler - Symboldid - Symbolit - Simboli - Sziembólumok - Simboliai - Simboli - Symboler - Symbolen - Symbolika - Simboluri - Symboly - Simboli - Symboler

الرموز

	IT - Fabbricante GB - Manufacturer FR - Fabricant DE - Hersteller ES - Fabricante PT - Fabricante GR - Κατασκευαστής BG - производител CZ - Výrobce DA - Fabrikant EE - Tootja FI - Valmistaja HR - Proizvođač HU - Gyártó LT - Gamintojas LV - Ražotājs NO - Produsent NL - Fabrikant PL - Producent RO - Producător SK - Výrobca SL - Proizvajalec SV - Tillverkare
	IT - Codice prodotto GB - Product code FR - Code produit DE - Code produit ES - Código de producto PT - Código do produto GR - Κωδικός προϊόντος BG - Код на продукта CZ - Kód produktu DA - Produktkode EE - Tuotekoodi FI - Tuotekoodi HR - Šifra proizvoda HU - Termék kódja LT - Prekės kodas LV - Produkta kods NO - Produktkode NL - Productcode PL - Kod produktu RO - Cod produs SK - Kód produktu SL - Šifra izdelka SV - Produktkod
	IT - Rappresentante autorizzato nella Comunità europea GB - Authorized representative in the European Community FR - Représentant autorisé dans la Communauté Européenne DE - Autorisierter Vertreter in der Europäischen Gemeinschaft ES - Representante autorizado en la Comunidad Europea PT - Representante autorizado na Comunidade Europeia GR - Εξουσιοδοτημένος αντιπρόσωπος στην Ευρωπαϊκή Κοινότητα BG - Упълномощен представител в Европейската общност CZ - Autorizovaný zástupce v Evropském společenství DA - Autoriseret repræsentant i Det Europæiske Fællesskab EE - Volitatud esindaja Euroopa Ühenduses FI - Valtuutettu edustaja Euroopan yhteisössä HR - Ovlašteni predstavnik u Europskoj zajednici HU - Meghatalmazott képviselő az Európai Közösségből LT - Įgaliotas atstovas Europos bendrijoje LV - Pilnvarotais pārstāvis Eiropas Kopienā NO - Autorisert representant i Det europeiske fellesskap NL - Gemachtigde vertegenwoordiger in de Europese Gemeenschap PL - Autoryzowany przedstawiciel na terenie Wspólnoty Europejskiej RO - Reprezentant autorizat în Comunitatea Europeană SK - Autorizovaný zástupca v Európskom spoločenstve SL - Pooblaščen zastopnik v Evropski skupnosti SV - Auktoriserad representant i Europeiska gemenskapen
	IT - Importato da GB - Imported by FR - Importé par DE - Importiert von ES - Importado por PT - Importado por GR - Εισαγωγή από BG - Внесено от CZ - Dovezeno uživatelem DA - Importeret af EE - Importija FI - Tuoja HR - Uvezeno od strane HU - Importálta: LT - Importavo LV - Importēja NO - Importert av NL - Geïmporteerd door PL - Importowane przez RO - Importat de SK - Dovážal SL - Uvozil SV - Importerad av
	IT - Dispositivo medico GB - Medical device FR - Dispositif médical DE - Medizinisches Gerät ES - Dispositivo medico PT - Dispositivo médico GR - Ιατρική συσκευή BG - Медицинско изделие CZ - Lékařské zařízení DA - Medicinsk udstyr EE - Meditsiini-seade FI - Lääketieteellinen laite HR - Medicinski uređaj HU - Orvosi eszköz LT - Medicinos prietaisai LV - Medicīnas ierīce NO - Medisinsk utstyr NL - Medisch apparaat PL - Urządzenie medyczne RO - Dispozitiv medical SK - Lekárska pomôcka SL - Medicinski pripomoček SV - Medicinsk apparat
	IT - Smaltimento RAEE GB - WEEE disposal FR - Élimination des DEEE DE - WEEE-Entsorgung ES - Eliminación de RAEE PT - Eliminação de REEE GR - Απόρριψη ΑΗΗΕ BG - Изхвърляне на ОЕЕО CZ - likvidace OEEZ DA - Bortskaffelse af WEEE EE - WEEE kõrvaldamine FI - WEEE hävittäminen HR - odlaganje WEEE HU - WEEE ártalmatlanítása LT - EEJ atliekų šalinimas LV - EEJA utilizācija NO - EE-avhending NL - AEEA-verwijdering PL - Utylizacja WEEE RO - eliminarea DEEE SK - likvidácia OEEZ SL - Odstranjanje var OEEO SV - WEEE-avfallshantering
	IT - Attenzione: Leggere e seguire attentamente le istruzioni (avvertenze) per l'uso GB - Warning: Read and follow the instructions (warnings) for use carefully FR - Avertissement : Lisez et suivez attentivement les instructions (avertissements) d'utilisation. DE - Warnung: Lesen und befolgen Sie die Gebrauchsanweisungen (Warnhinweise) sorgfältig ES - Advertencia: Lea y siga atentamente las instrucciones (advertencias) de uso. PT - Aviso: Leia e siga as instruções (avisos) de uso com atenção GR - Προειδοποίηση: Διαβάστε και ακολουθήστε προσεκτικά τις οδηγίες (προειδοποιήσεις) χρήσης BG - Предупреждение: Прочетете и следвайте внимателно инструкциите (предупрежденията) за употреба CZ - Upozornění: Pečlivě si přečtěte a dodržujte pokyny (varování) k použití DA - Advarsel: Læs og følg brugsanvisningen (advarslerne) omhyggeligt EE - Hoiatus: Lugege kasutusjuhised (hoiatused) hoolikalt läbi ja järgige neid FI - Varoitus: Lue käyttöohjeet (varoitukset) huolellisesti ja noudata niitä HR - Upozorenje: Pažljivo pročitajte i slijedite upute (upozorenja) za uporabu HU - Figyelmeztetés: Olvassa el és kövesse figyelmesen a használati utasításokat (figyelmeztetéseket). LT - Įspėjimas: atidžiai perskaitykite ir laikykitės naudojimo instrukcijų (įspėjimų). LV - Brīdinājums: uzmanīgi izlasiet un ievērojiet lietošanas instrukcijas (brīdinājumus). NO - Advarsel: Les og følg bruksanvisningen (advarslerne) nøye NL - Waarschuwing: Lees en volg de instructies (waarschuwingen) voor gebruik zorgvuldig PL - Ostrzeżenie: Przeczytaj uważnie instrukcje (ostrzeżenia) dotyczące użytkowania i postępuj zgodnie z nimi RO - Avertisment: Citiți și urmați cu atenție instrucțiunile (avertismente) de utilizare SK - Upozornenie: Pozorne si prečítajte a dodržiavajte pokyny (upozornenia) na použitie SL - Opozorilo: natančno preberite in upoštevajte navodila (opozorila) za uporabo SV - Varning: Läs och följ instruktionerna (varningarna) för användning noggrant
	IT - Seguire le istruzioni per l'uso GB - Follow the instructions for use FR - Suivre les instructions d'utilisation DE - Befolgen Sie die Gebrauchsanweisung ES - Sigue las instrucciones de uso. PT - Siga as instruções de uso GR - Ακολουθήστε τις οδηγίες χρήσης BG - Следвайте инструкциите за употреба CZ - Postupujte podle návodu k použití DA - Følg brugsanvisningen EE - Järgige kasutusjuhendit FI - Noudata käyttöohjeita HR - Slijedite upute za uporabu HU - Kövesse a használati utasítást LT - Vadovaukitės naudojimo instrukcijomis LV - Izpildiet lietošanas instrukcijas NO - Følg bruksanvisningen NL - Volg de instructies voor gebruik PL - Postępuj zgodnie z instrukcją użycia RO - Urmăți instrucțiunile de utilizare SK - Postupujte podľa návodu na použitie SL - Sledite navodilom za uporabo SV - Följ bruksanvisningen

SA - اتباع تعليمات الاستخدام

	IT - Dispositivo medico conforme al regolamento (UE) 2017/745 GB - Medical device compliant with Regulation (EU) 2017/745 FR - Dispositif médical conforme au Règlement (UE) 2017/745 DE - Medizinprodukt gemäß Verordnung (EU) 2017/745 ES - Dispositivo médico que cumple con el Reglamento (UE) 2017/745 PT - Dispositivo médico em conformidade com o Regulamento (UE) 2017/745 GR - Ιατροτεχνολογικό προϊόν συμμορφούμενο με τον Κανονισμό (ΕΕ) 2017/745 BG - Медицинско изделие, отговарящо на Регламент (ЕС) 2017/745 CZ - Zdravotnický prostředek v souladu s nařízením (EU) 2017/745 DA - Medicinsk udstyr i overensstemmelse med forordning (EU) 2017/745 EE - Määrusele (EL) 2017/745 vastav meditsiiniisese FI - Asetuksen (EU) 2017/745 mukainen lääkinnällinen laite HR - Medicinski uređaj uskladen s Uredbom (EU) 2017/745 HU - Az (EU) 2017/745 rendeletnek megfelelő orvostechnikai eszköz LT - Medicinos prietaisai, atitinkantis Reglamentą (ES) 2017/745 LV - Medicīnas ierīce, kas atbilst Regulai (ES) 2017/745 NO - Medisinsk utstyr i samsvar med forordning (EU) 2017/745 NL - Medisch hulpmiddel dat voldoet aan Verordening (EU) 2017/745 PL - Wyrób medyczny zgodny z Rozporządzeniem (UE) 2017/745 RO - Dispozitiv medical conform Regulamentului (UE) 2017/745 SK - Zdravotnícka pomôcka v súlade s nariadením (EÚ) 2017/745 SL - Medicinski pripomoček skladen z Uredbo (EU) 2017/745 SV - Medicinsk utrustning i enlighet med förordning (EU) 2017/745
	IT - Conservare in luogo fresco ed asciutto GB - Store in a cool, dry place FR - Conserver dans un endroit frais et sec DE - An einem kühlen, trockenen Ort aufbewahren ES - Conservar en un lugar fresco y seco. PT - Armazene em local fresco e seco GR - Φυλάσσετε σε δροσερό, ξηρό μέρος BG - Съхранявайте на хладно и сухо място CZ - Skladujte na chladném a suchém místě DA - Opbevares på et køligt, tørt sted EE - Hoida jahedas ja kuivas kohas FI - Säilytä viileässä, kuivassa paikassa HR - Čuvati na hladnom i suhom mjestu HU - Tárolja hűvös, száraz helyen LT - Laikyti vėsioje, sausioje vietoje LV - Uzglabāt vēsā, sausā vietā NO - Opbevares på et kjølig, tørt sted NL - Op een koele, droge plaats bewaren PL - Przechowywać w chłodnym, suchym miejscu RO - A se păstra într-un loc răcoros și uscat SK - Skladujte na chladnom a suchom mieste SL - Hraniti na hladnem in suhem mestu SV - Förvara på en sval, torr plats
	IT - Conservare al riparo dalla luce solare GB - Store away from sunlight FR - Conserver à l'abri du soleil DE - Vor Sonnenlicht schützen ES - Almacenar lejos de la luz solar PT - Armazene longe da luz solar GR - Φυλάσσεται μακριά από το φως του ήλιου BG - Да се съхранява далеч от слънчева светлина CZ - Skladujte mimo dosahu slunečního záření DA - Opbevares væk fra sollys EE - Hoida eemal päikesevalgusest FI - Säilytettävä poissa auringonvalolta HR - Čuvati dalje od sunčeve svetlosti HU - Tárolja napfénytől távol LT - Laikyti atokiau nuo saulės spindulių LV - Glabāt prom no saules gaismas NO - Opbevares unna sollys NL - Uit de buurt van zonlicht bewaren PL - Przechowywać z dala od światła słonecznego RO - A se păstra departe de lumina soarelui SK - Skladujte mimo dosahu slnečného žiarenia SL - Hraniti ločeno od sončne svetlobe SV - Förvaras åtskillt från solljus
	IT - Limite di pressione atmosferica GB - Atmospheric pressure limit FR - Limite de pression atmosphérique DE - Luftdruckgrenze ES - Limite de presión atmosférica PT - Limite de pressão atmosférica GR - Όριο ατμοσφαιρικής πίεσης BG - Граница на атмосферното налягане CZ - Mezní hodnota atmosférického tlaku DA - Atmosfærisk trykgrænse EE - Atmosfäärirõhu piir FI - Ilmanpaineen raja HR - Granica atmosferskog tlaka HU - Légköri nyomás határértéke LT - Atmosferos slėgio riba LV - Atmosfēras spiediena robeža NO - Atmosfærisk trykkgrense NL - Limiet atmosferische druk PL - Dopuszczalne ciśnienie atmosferyczne RO - Limita presiunii atmosferice SK - Hranica atmosférického tlaku SL - Omejitve atmosferskega tlaka SV - Atmosfäriskt tryckgräns
	IT - Limite di umidità GB - Humidity limit FR - Limite d'humidité DE - Luftfeuchtigkeitsgrenze ES - Limite de humedad PT - Limite de umidade GR - Όριο υγρασίας BG - Ограничение на влажността CZ - Limit vlhkosti DA - Fugtgrænse EE - Niiskuse piirang FI - Kosteusraja HR - Granica vlažnosti HU - Páratartalom határérték LT - Drėgmės riba LV - Mitruma robeža NO - Fuktighetsgrense NL - Vochtigheidslimiet PL - Limit wilgotności RO - Limita de umiditate SK - Hranica vlhkosti SL - Omejitve vlažnosti SV - Fuktighetsgräns
	IT - Limite di temperatura GB - Temperature limit FR - Limite de température DE - Temperaturgrenze ES - Limite de temperatura PT - Limite de temperatura GR - Όριο θερμοκρασίας BG - Температурна граница CZ - Teplotní limit DA - Temperaturgrænse EE - Temperatuuri piirang FI - Lämpötilaraja HR - Ograničenje temperature HU - Hőmérséklet korlát LT - Temperatūros riba LV - Temperatūras ierobežojums NO - Temperaturgrense NL - Temperatuurlimiet PL - Limit temperatury RO - Limită de temperatură SK - Teplotný limit SL - Temperaturna omejitve SV - Temperaturgräns
	IT - Identificatore unico dispositivo GB - Unique device identifier FR - Identifiant unique de l'appareil DE - Eindeutige Geräteerkennung ES - Identificador de dispositivo único PT - Identificador exclusivo do dispositivo GR - Μοναδικό αναγνωριστικό συσκευής BG - Уникален идентификатор на устройството CZ - Jedinečný identifikátor zařízení DA - Unik enhedsidentifikator EE - Seadme kordumatu identifikator FI - Ainutlaatuinen laitetunniste HR - Jedinstveni identifikator uređaja HU - Egyedi eszközazonosító LT - Unikalus įrenginio identifikatorius LV - Unikāls ierīces identifikators NO - Unik enhetsidentifikator NL - Unieke apparaat-ID PL - Unikalny identyfikator urządzenia RO - Identificator unic de dispozitiv SK - Jedinčný identifikátor zariadenia SL - Enolični identifikator naprave SV - Unik enhetsidentifierare
	IT - Numero di lotto GB - Batch number FR - Numéro de lot DE - Chargennummer ES - Número de lote PT - Número do lote GR - Αριθμός παρτίδας BG - Партиден номер CZ - Číslo šarže DA - Batchnummer EE - Partii number FI - Erän numero HR - Broj serije HU - Tételszám LT - Partijos numeris LV - Partijas numurs NO - Batchnummer NL - Batchnummer PL - Numer partii RO - Numărul lotului SK - Číslo šarže SL - Številka serije SV - Batchnummer
	IT - Data di fabbricazione GB - Manufacturing date FR - Date de fabrication DE - Herstellungsdatum ES - Fecha de fabricación PT - Data de fabricação GR - Ημερομηνία κατασκευής BG - Дата на производство CZ - Datum výroby DA - Fremstillingsdato EE - Tootmistuupäev FI - Valmistuspäivämäärä HR - Datum proizvodnje HU - Gyártási dátum LT - Pagaminimo data LV - Ražošanas datums NO - Produksjonsdato NL - Productiedatum PL - Data produkcji RO - Data fabricației SK - Dátum výroby SL - Datum izdelave SV - Tillverkningsdatum

SA - تاريخ التصنيع

	IT - Parte applicata di tipo B GB - Type B applied part FR - Pièce appliquée de type B DE - Anwendungsteil vom Typ B ES - Pieza aplicada tipo B PT - Peça aplicada tipo B GR - Εφαρμοσμένο τμήμα τύπου Β BG - Приложна част тип B CZ - Použitý díl typu B DA - Type B anvendt del EE - B-tüüpi rakendatud osa FI - Tyypin B käytetty osa HR - Primijenjeni dio tipa B HU - B típusú alkalmazott alkatrész LT - B tipo pritaikyta dalis LV - B tipa lietotā daļa NO - Type B påført del NL - Type B toegepast onderdeel PL - Część aplikacyjna typu B RO - Piesa aplicata tip B SK - Aplikovaná část typu B SL - Uporabljeni del tipa B SV - Typ B applicerad del SA - الجزء المطبق من النوع B
IPX6	IT - Grado di protezione dell'involucro GB - Degree of protection of the envelope FR - Degré de protection de l'enveloppe DE - Schutzgrad des Gehäuses ES - Grado de protección de la carcasa PT - Grau de proteção da caixa GR - Βαθμός προστασίας του περιβλήματος BG - Степен на защита на корпуса CZ - Stupeň krytí pouzdra DA - Beskyttelsesgrad af kabinettet EE - Korpuse kaitseaste FI - Kotelon suojausluokka HR - Stupanj zaštite kućišta HU - A burkolat védeettségi fokja LT - Korpuso apsaugos laipsnis LV - Korpusa aizsardzības pakāpe NO - Beskyttelsesgrad for dekselet NL - Beschermingsgraad van de behuizing PL - Stopień ochrony obudowy RO - Gradul de protecție al carcasei SK - Stupeň ochrany krytu SL - Stopnja zaščite ohišja SV - Skyddsgrad för höljet SA - درجة حماية الغلاف

