

EN INSTALLATION AND OPERATING INSTRUCTIONS

1. CLEAN AND DRY UP THE SURFACE WHERE THE MAT IS GOING TO BE PLACED

2. REMOVE THE LOWER PROTECTIVE SHEET

CAUTION: *WHILE REMOVING THE LOWER PROTECTIVE SHEET MAKE SURE NOT TO DETACH THE ADHESIVE SHEET THAT HAS TO STICK THE MAT TO THE GROUND*








3. MAKE THE MAT SMOOTHLY ADHERE TO THE FLOOR







4. REMOVE THE UPPER PROTECTIVE SHEET

5. PROGRESSIVELY REMOVE EACH OF THE 30 DISPOSABLE ADHESIVE SHEETS WHEN DIRTY

CAUTIONS

- FOR INDOOR USE ONLY
- INSTALL ON SMOOTH AND FLAT SURFACES ONLY
- AVOID CONTACT WITH WATER AND TRANSIT WITH WET SOLES: EXTREMELY SLIPPERY WHEN WET
- CARRY AND STORE IN THE ORIGINAL PACKAGE
- DO NOT FOLD THE PACKAGE
- STORE IN A DRY, COOL PLACE AWAY FROM STRONG HEAT SOURCES PROTECT FROM RAIN AND DIRECT SUNBEAMS
- DISPOSE IN RESPECT OF LOCAL LAWS AND REGULATIONS CONCERNING PLASTIC MATERIALS (LOW DENSITY POLYTHE-NE)

	IT Codice prodotto GB Product code FR Code produit ES Código producto PT Código produto FI Tuotekoodi
	IT Numero di lotto GB Lot number FR Numéro de lot ES Número de lote PT Número de lote FI Eränumero
	IT Conservare in luogo fresco ed asciutto GB Keep in a cool, dry place FR À conserver dans un endroit frais et sec ES Conservar en un lugar fresco y seco PT Armazenar em local fresco e seco FI Säilytä kuivassa ja viileässä
	IT Conservare al riparo dalla luce solare GB Keep away from sunlight FR À conserver à l'abri de la lumière du soleil ES Conservar al amparo de la luz solar PT Guardar ao abrigo da luz solar FI Säilytä auringonvaltalta suojassa
	IT Fabbricante GB Manufacturer FR Fabricant ES Fabricante PT Fabricante FI Valmistaja
	IT Data di fabbricazione GB Date of manufacture FR Date de fabrication ES Fecha de fabricación PT Data de fabrico FI Valmistuspäivämäärä
	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 ES Producto sanitario conforme con el reglamento (UE) 2017/745 PT Dispositivo médico em conformidade com a regulamento (UE) 2017/745 FI Lääkinnällinen laite, joka vastaa asetusta (EU) 2017/745

	IT Attenzione: Leggere e seguire attentamente le istruzioni (avvertenze) per l'uso GB Caution: read instructions (warnings) carefully FR Attention: lisez attentivement les instructions (avertissements) ES Precaución: lea las instrucciones (advertencias) cuidadosamente PT Cuidado: leia as instruções (avisos) cuidadosamente FI Huomio: Lue käyttöohjeet (varoitukset) ja noudata niitä huolellisesti
	IT Dispositivo monouso, non riutilizzare GB Disposable device, do not re-use FR Dispositif pour usage unique, ne pas réutiliser ES Dispositivo monouso, no reutilizable PT Dispositivo descartável, não reutilizar FI Kertakäyttöinen laite, ei saa käyttää uudelleen
	IT Data di scadenza GB Expiration date FR Date d'échéance ES Fecha de caducidad PT Data de validade FI Viimeinen voimassaolopäivä
	IT Dispositivo medico GB Medical Device FR Dispositif médical ES Producto sanitario PT Dispositivo médico FI Lääkinnällinen laite
	IT Non sterile GB Non-sterile FR Pas stérile ES No estéril PT Não estéril FI Ei-steriili
	IT Identificativo unico GB Unique identifier FR Identifiant unique ES Identificador único PT Identificador exclusivo FI Ainutlaatuinen tunniste

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.

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.

Kaikista vakavista tapaturmista, jotka liittyvät toimittamamme lääkinällisen laitteen käyttöön, on ilmoitettava valmistajalle sekä oman asuinpaikan jäsenmaan toimivaltaiselle viranomaiselle.