



TITANOX s.r.l. - via Canove 2/A - Loc. Canove dé
Biazzi -26038 Torre dé Picenardi (CR) - Italy -
www.titanox.it info@titanox.it MADE IN ITALY

Instruction manual “M600358/441”

All models are produced eliminating sources of potential risk to the user, should unintended collisions occur, such as cutting edges or sharp corners.

These instruments are intended to be used by medical practitioners who are specially trained on how to use and care of them. Use for purposes other than what it is intended for is not allowed. The incorrect use, poor or inappropriate maintenance can rapidly lead to deterioration of the instruments.

- Check the right tightening torque of all mobile parts before and during use.
- For cleaning, do not use abrasive substances, acids, alcohol, chlorine-based cleaners, disinfectants, and acetone: the manufacturer company cannot be held responsible for the damage that is caused by using materials that could damage the product surface or corrosive chemicals during cleaning.

WARNING: The patient must sit **ONLY** on the seat (the central section of the mattress). The backrest is not designed to support the full weight of the patient's body. Sitting on the backrest causes his damage.

TITANOX WARRANTY TERMS

Titanox 12-month warranty applies from final invoice date.










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.



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	IT Fabbricante ENG Manufacturer FR Fabricant
	IT Attenzione: Leggere e seguire attentamente le istruzioni (avvertenze) per l'uso ENG Caution: read instructions (warnings) carefully FR Attention: lisez attentivement les instructions (avertissements)
	IT Leggere le istruzioni per l'uso ENG Consult instructions for use FR Consulter les instructions d'utilisation
	IT Conservare al riparo dalla luce solare ENG Keep away from sunlight FR Á conserver à l'abri de la lumière du soleil
	IT Conservare in luogo fresco ed asciutto ENG Keep in a cool, dry place FR Á conserver dans un endroit frais et sec
	IT Codice prodotto ENG Product code FR Code produit
	IT Numero di lotto ENG Lot number FR Numéro de lot
	IT Dispositivo medico ENG Medical Device FR Dispositif médical
	IT Il dispositivo è conforme ai requisiti del regolamento sui dispositivi medici (UE) 2017/745 ed è classificato come dispositivo medico di classe I ENG The product complies with the requirements of the regulation on medical devices (EU) 2017/745 and is classified as a Class I medical device FR Le produit est conforme aux exigences du règlement sur les dispositifs médicaux (UE) 2017/745 et est classé comme dispositif médical de classe I

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