

USER MANUAL

MNPG362-02 Edition 04/07/2022

F3S2000 and F2S700 Band applicators with solenoids

I.A.C.E.R. Srl

www.itechmedicaldivision.com



Introduction

F3S2000 and **F2S700** devices are applicators with three and two solenoids for low-frequency magnetotherapy.

They can easily adapt to any area of the body thanks to the presence of the therapeutic elastic band.

The devices are classified as class I, in accordance with rule 1 (4.1) referred to in Annex VIII of Regulation (EU) 2017/745 of the European Parliament and of the Council dated 5 April 2017, relating to medical devices.

Basic UDI-DI: 8019781PEMFLFFASNS

F3S2000 UDI-DI: 08019781472052

F2S700 UDI-DI: 08019781472212

Technical data

Technical specifications:

- Applicator size: 38x9 cm + elastic band of 60 cm;
- Cable length: 158 cm;
- 3 solenoids or 2 solenoids;
- 1 connection cable;
- F3S2000 model compatible with MAG2000 models and LaMagneto models.
- F2S700 model is compatible only with the MAG700 model

Labelling

Label 1





Label 1



Model F3S2000



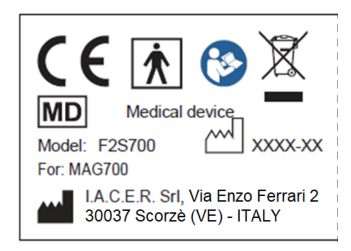
Model F2S700

Cable label:

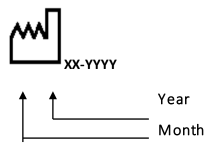
Modello / Model: F3S200



Modello / Model: F2S700







Symbols:

Symbols: Symbol	Meaning
I-TECH MEDICAL DIVISION	Manufacturer's logo.
CE	In compliance with regulation (EU) no. 2017/745
	Manufacturer data.
سا	Date of manufacture (MM-YYYY).
MD	Medical device.
	Follow the instructions for use.
	WEEE directive for the disposal of electronic waste.
*	Type BF applied part.
	Allowed temperatures (storage temperatures, on packaging).



Symbol	Meaning
<u></u>	Relative humidity (relative humidity of storage, on packaging).

Transport and storage

There are no special care instructions for transport. For storage, the equipment is protected up to the following environmental conditions:

room temperature from +5 to $+40^{\circ}$ C relative humidity from 15 to 93%

Instructions for use

Instructions for using F3S2000 and F2S700

Follow the instructions in the User Manual supplied with the device in your possession.

<u>Follow the Instructions given in the user manuals of the MAG 700, MAG 2000 series or the LaMagneto family.</u>

How to take care of F3S2000

Cleaning the F3S2000 and F2S700

To clean the F3S2000 and F2S700 devices, it is recommended to disconnect the three/2 solenoid applicator from the device before performing any operation. Clean the fabric with a cloth moistened with simple water and neutral soap and wait for it to dry completely before reusing the three solenoid applicator for a therapy session.

Information for disposal

The product is subject to the WEEE legislation (presence on the label of the symbol) relating to separate waste collection: to dispose of the product, use special areas equipped for the collection of electronic material by contacting the competent authorities of your country or the manufacturer directly.



Maintenance and Support

The device does not require any particular routine maintenance. Please contact the manufacturer to provide for a check of the functionality of the device, every two years.

The manufacturer is the sole entity authorized to provide technical support for the F3S2000 applicator. Should you need technical support, please contact:

I.A.C.E.R. S.r.l.

Via Enzo Ferrari, 2 • 30037 Scorzè (VE)

Ph. 041.5401356 • Fax 041.5402684

Warranty

F3S2000 is covered by a 2-year warranty starting from the date of purchase. The warranty shall lapse in case of tampering, neglect and inappropriate use of the applicator and in case of any intervention carried out by personnel who is not authorized by the manufacturer or by the authorized dealer.

Incident reporting

In compliance with the provisions of Regulation (EU) 2017/745, the manufacturer places the need to report any serious accident in relation to the device to the attention of the user.

The report must be addressed:

• to the device manufacturer:

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