Instructions for Use

External Fixator - Short Bones II

Subtitles of the symbols used on the packaging

REF	Catalogue Number
STERILE R	Sterile Product - Sterilized by Gamma Radiation
~~	Date of Manufacture
Ti	Consult instructions for use
	Do not use if package is damaged
淤	Keep out of the sun
*	Keep Dry

LOT	Batch Code
(E	Product certified in accordance with Directive 93/42/EEC). When applicable.
\square	Valid until
2	Single-Use Product
anality z	Do not re-sterilize
Ţ	Fragile, handle with care.
40 ° C	Temperature Limit (40°C);

Features and technical specifications of the product

Technical Name: External fixator for extremities of the skeleton - short bones

Trade Name: External Fixator - Short Bones II

Trade Models:

External Fixator MiniFixExternal Fixator Colles type

Raw Material:

• Stainless Steel Alloy (ASTM F899: AISI 304)

• Aluminum Alloy (ASTM B221 type 6351 and type 5052)

Validity: 05 years (after sterilization date)

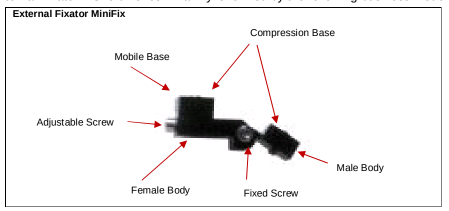
Sterile Product

Sterilization Method: Gamma Radiation (dosage of 25 kGy)

Description

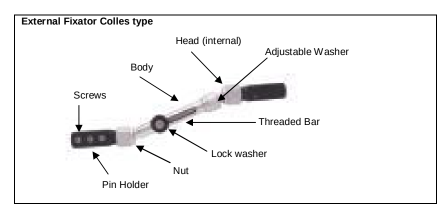
The **External fixator – Short Bones II** Family in general was designed to treat open fractures, with a degree of skin lesion and / or soft tissue that make unfeasible or counter indicate the use of gypsum or, internal fixation, due to the risk of infection or exposure of the product. With the passage of time, indications were extended to unstable fractures, bone transport, stretching and treatment of pseudarthrosis.

The External Fixator – Short Bones II Family is formed by the following business models:



The External Fixator MiniFix is unilateral with articulated body, for correction of simple deviations used in phalange, made up of the male body and female body, which are united through the fixed screw permitting the pivoting motion of the phalange. The adjustable screw located inside the female body determines the optimal position of the mobile base for fixing the Schanz pin through the compression base by the male body that also passes by the Schanz pin, which is fixed by the compression base.

The External Fixator MiniFix features a pivotal motion that benefits the patient in the treatment of fractures in this sector as it prevents hardening of the phalange due to inactivity.



The External Fixator Colles type is unilateral with bi-articulated body which enables the correction of angular and bone distraction. It is a product used in the distal radius extremity; its components have adjustment flexibility enabling greater precision when fixing the Schanz pin. The pin fixation is made through pin holder, located at the ends of the fixator, more precisely threaded to the nuts which are stuck to the head and the body. For adjusting the extension of the fixator it is necessary to unscrew the lock washer screw, and by rotating the adjustable washer allows the displacement of the threaded bar by changing the length of the fixator.

Composition

The materials selected for the composition have the properties required to achieve the desired performance for the product. This selection considered factors such as biocompatibility, physical, chemical and mechanical properties required for the product.

The trade models which make up the External Fixator – Short Bones II Family, are made of Stainless Steel Alloy (AISI 304), specified by the Norm ASTM F-899 - Standard Specification for Stainless Steel for Surgical Instruments and Aluminum Alloys (type 6351 and 5052) specified by the Norm ASTM B221.

Indication and Purpose

The **External Fixator - Short Bones II** Family, in general, was designed to treat open fractures, with an amount of skin lesion and/or soft tissue that make unfeasible or counter indicates the use of gypsum or, internal fixation, due to the risk of infection or exposure of the product. With the course of time, indications were extended to unstable fractures, bone transport, stretching and treatment of pseudarthrosis.

External Fixator MiniFix

Suitable for fracture stabilization in the phalange, it is widely used in:

- · Opened fractures with devitalized tissues;
- Complicated fractures due to soft tissues or bone losses;
- Comminuted fractures or those ones associated with burns or infections;
- · Poly fractures including multiple metacarpal fractures;
- Complex fractures with nerves, tendons and blood vessels injuries;
- In cases of explosion, avulsion or crush injury caused by firearms.

External Fixator Colles type

It is indicated for fracture fixation with posterior displacement characteristic of the fragment of the distal radius extremity going backwards and outwards, performing a typical aspect of the hand like the back of a fork, and its most frequent cause is a fall on the open hand.

Contraindications

Relative contraindications for the use of the device are listed below, so that the surgeon in charge, after a thorough study of the case, will indicate the procedures:

- Patients with general or specific active infections that can lead the fixation to complications;
- Patients with impaired general state and/or immune compromised, unable to be submitted to a surgical procedure
- Patients with sensibility to foreign bodies, being that in these cases, tests should be performed;
- Patients with osteoporosis and/or bone affections that may compromise the fixation stability;
- Patients who use narcotic substance, alcohol or smoke.

Forms of Presentation

The business models that make up the **External Fixator – Short Bones II** Family are unitarily packed in a double primary packing system, blister type (PET) sealed with surgical grade paper (Tyvek® type) or in double envelope like package of surgical grade paper (Tyvek® type), which work as sterilization shield.

The product is available for commercialization in sterile condition, and the adopted sterilization method is gamma radiation (dosage of 25kGy). This procedure is performed by a suitably qualified outsourced company.

Once sterilized, the products packaged in their primary packaging properly labeled, are packed in a cardboard carton (secondary packaging), in which there is a leaflet with instructions for use of the product.

A label with information necessary for the product identification is glued on the top of the cardboard carton.

The **External Fixator – Short Bones II** Family are available in the trade models below, and each one are available for commercialization according to these codes and descriptions listed below:

Illustrative Image	Code	Description	Dimensions	Made of	Qtty. Packed
-	04.35.09.00000	External Fixator MiniFix	17x57 mm	Stainless Steel Alloy (ASTM F-899: AISI 304) + Aluminum Alloy (ASTM B221 types 5052 /6351)	01
	04.35.10.00000	External Fixator Colles type	14x188 mm	Stainless Steel Alloy (ASTM F-899: AISI 304) + Aluminum Alloy (ASTM B221 types 5052 /6351)	01

Ancillary Components:

The ancillary components to the External Fixator – Short Bones II are:

- Schanz Pin
- Conical Schanz Pin

Ancillary components are manufactured from Stainless steel Alloy (18Cr-14Ni-2.5Mo), which meets the requirements specified by ASTM F-138 – 'Standard Specification for Wrought 18 Chromium-14 Nickel-2.5 Molybdenum Stainless Steel Bar and Wire for Surgical Implants (UNS S31673)'.

The surgeon in charge of the adopted technique is also responsible for the right choice of the models and sizes of the External Fixator – Short Bones II and its ancillaries to be used. He must be familiar with the material, the application method and the adopted surgical procedures.

The procedure success is linked to the right choice, positioning, and the devices fixation. It is responsibility of the surgeon who evaluates the patient and decides which devices are to be used. Also, it is linked to the strict compliance with postoperative care recommended by the surgeon in charge.

The ancillaries below are not subject of this registration process, and should, therefore, be always purchased separately from the same manufacturer or from another one the company indicates.

List of ancillaries to the External Fixator – Short Bones II Family

Illustrative Image	Code	Description	Dimensions	Made of
	04.25.10.XXXXX	Schanz Pin	Ø 2,5 – 80, 100 mm Ø 3,0 – 10, 100 mm Ø 3,5 – 80, 100 mm Ø 4,0 – 90, 130, 150, 170 mm	Stainless Steel Alloy (18Cr-14Ni-2.5Mo) ASTM F138

Support Materials

The supporting materials for assembling the External Fixator – Short Bones II Family are the instruments designated for this purpose.

The instruments below are not subject of this registration process, and should, therefore, be always purchased separately from the same manufacturer or from another one the company indicates.

See below a list of the instruments available and provided by the manufacturer or their indication for using the External Fixator – Short Bones II:

0F.03 – Instrument – Colles External Fixator

The instruments are provided decontaminated, but not sterilized. Inappropriate sterilization of the surgical instrument might cause infection.

Surgical instruments are subject to wear and tear during their regular use. Therefore breaking may occur. The instruments should only be used for the purpose they were designed to and should be inspected regularly for possible wear and damage.

For further information concerning the instruments, please consult the dealer.

Warning and Precautions

For the product use, the medical team in charge must consider the following warning and precautions:

- The External Fixator Short Bones II must only be used after a thorough analysis of the surgical procedure to be adopted and complete reading of these instructions for use;
- The product should only be handled by specialized surgical teams with specific knowledge
 and capacity building concerning osteosynthesis techniques. The choice and dominance of
 the adopted technique to be applied are under the responsibility of the surgeon in charge;
- Inappropriate choice and selection of the products to be used, as well as mistakes concerning
 the indication, handling and application technique might cause excessive stress and tractions
 on the device leading to failure due to fatigue, fracture and even looseness;
- Clinical results and the durability of the device are totally dependent upon a precise surgical technique;
- The surgeon must have extensive knowledge of the local anatomy. It is extremely important the anatomic references for defining the insertion point;
- The use of bone graft may be needed, but it is under medical criteria;
- A greater risk of the device failure is its use in patients who are predisposed to disobey medical guidelines and postoperative restrictions, such as children, elderly, individuals with neurological changes, or addicted;
- Device failure risks are greater in patients who practice physical exertion activities or those
 who practice sports during the postoperative period, contradicting the medical restrictions;
- The patient must be submitted to periodic medical monitoring to check the device, the bone and the adjacent tissues conditions;
- The External Fixator Short Bones II and its respective ancillaries should not be used whether there is not an appropriate osseous support that can guarantee the device stability;
- The pre and perioperative prophylactic antibiotic therapy as well as antibiotic therapy in cases there is a local and/or systemic predisposition or infections occur – are under medical criteria;

- The correct matching of the External Fixator Short Bones II and its respective ancillary
 components is indicated in the "Ancillary Components" topic, and due to the possibility of
 dimensional and functional incompatibility, it must not be used with other components than the
 ones indicated by the manufacturer.
- Do not use the product if the shelf life is expired or packaging is violated;
- Falls or crushing on hard surfaces might damage the product. So, it is necessary the handler
 to perform inspection of the product to check its integrity while it is unpacked and if there is
 any abnormality, the product should not be used;
- SINGLE-USE product Do not Reuse
- Sterile Product Do Not Re-sterilize:
- Manufacturing date, expiry term and batch number: see label;

Adverse Effects

Every surgical procedure presents some common risks and complication possibilities such as infections, bleeding, allergic drug reactions and anesthetic risks, among others. The following complications and adverse effects can still be associated with the implantation of the product:

- Injuries to the nerves with sensitive or motor harm;
- Absence or delay of bone fusion resulting in the device breaking;
- Loosening, dismemberment, displacement, twisting or break of the device;
- · Deformation or fracture of the device;
- · Pains, discomfort or abnormal sensations due to the product;
- · Reaction to foreign body;
- · Bone necrosis or adjacent soft tissues;

The decision for the removal due to one of the above adverse effects is made by the surgeon in charge.

Use Instructions

For the correct use of product, the following instructions should be adopted:

- The care of this material is responsibility of the skilled staff, which should follow the standards and/or other local regulations applied;
- The product should be handled with appropriate care in adequate locations (materials center and operating rooms);
- The product should only be used by specialized surgical teams, with specific knowledge and training on techniques for osteosynthesis, and the surgeon in charge is responsible for the choice and dominance of the surgical technique to be performed;
- The insertion should be made from the anatomical site of greatest risk to the opposite direction;
- Palpate the closest artery and insert the pin at a minimum distance of 2 cm from it.
- The limb must be extended for drilling the flexor muscle and it must be flexed for drilling the extensor muscle.
- The device useful life is characterized by the time required for effective bone healing. After this
 period, in case of absence or problems with bone healing, these conditions can represent a
 risk of device failure by excessive mechanic stress;
- A revision surgery may be necessary in the case mentioned right above or if loosening of the components is observed.
- It is necessary to use specific instruments for applying the External Fixator Short Bones II as
 indicated in the "Supporting Material". Due to the possibility of dimensional and/or functional
 incompatibility it MUST NOT be used with any other instruments different from the ones
 indicated by the manufacturer;
- The External Fixator Short Bones II must not be used with other components than the ones
 indicated by the manufacturer due to the possibility of dimensional and functional
 incompatibility.

The responsible surgical team should guide the patient or his legal representative about:

- The suitable care and restrictions during the postoperative period. The capacity and willingness of patient to follow these guidelines are one of the most important aspects in a surgical procedure;
- The fact that the risks are greater when using in patients with predisposition to disobey the medical guidelines, care and postoperative restrictions, such as children, elderly, individuals with neurological disorders or addicted;
- The fact that the product does not substitute nor does have the same performance of normal bone and therefore can break, deform or loosening due to excessive effort or activities of early load and other situations:
- The need to restrict the effort activities or sportive practice during the postoperative period, whose extension is defined by the surgeon in charge;
- The necessity of use of external supports, aid to ambulate and orthopedic appliances, designed to limit the movement and/or load is under exclusive medical criteria;
- The necessity of periodic medical monitoring to check the conditions of the device, the bone and the adjacent tissues;
- The fact that the non-performing of the revision surgery, in cases in which bone fusion does not occur, can lead the mechanical failure of the device;
- · The need of review surgery in cases of components loosening;
- The fact that implants can interfere with results of imaging examinations. So, implant users should report this fact when submitted to such examinations;
- The listed information in this topic "Guidance to the patient and/or the Legal Representative" and in the topic "Adverse Effects.

Sterilization

The External Fixator – Short Bones II is provided in the sterile product condition. The adopted sterilization method is by Gamma Radiation (dosage of 25 kGy).

The External Fixator – Short Bones II manufacturing process is done with great care in order to meet the desirable performance of the product. So, the surgical team and all the staff involved must handle all the devices properly in order to reduce the risk of infection.

Sterile Product - Do Not Re-sterilize

Do not use the product if the packaging is violated.

Contamination Risk

As this is a surgically invasive product, in cases in which there is need of removal the External Fixator, there are risks of biologic contamination and viral disease transmission.

For minimizing these risks, the explanted External Fixator – Short Bones II components should be treated as potentially contaminant material and should be adopted the standardization and/or other local regulations applied.

Product Discard

The devices removed or regarded as inappropriate for use must be discarded. It's highly recommended that before discarding, the product is mischaracterized, and so its parts can be cut, bent or sanded.

The components should be discarded in proper locals, to avoid the environmental contamination and other individuals. It is recommended adoption of local legal regulations for potentially contaminant products.

Single use product – do not reuse.

Traceability

To ensure the traceability of the product, and comply with the requirements for sanitary surveillance, the surgeon or his team must register in the patient's medical record the information about the product. Furthermore, such information must also be passed to the distributor of the product and to the patient in order to complete the cycle of the product traceability established. The information necessary for traceability are relatives to the product used, surgery and patient, as below:

· Name of patient who received the product;

- Surgeon's name;
- Hospital's name;
- Manufacturer's name;
- Supplier's name;
- Surgery date;
- Code of product;
- · Number of batch of the product;
- Quantity used;
- · Registration product at ANVISA.

The following Information needed for the product traceability is engraved on the piece or may be acquired from the label of the product package:

- · Company logotype;
- · Manufacturing Batch;
- Code of the piece.

Traceability information is required for notifying the Sanitary Surveillance Agency ANVISA, either by the health service or by the patient him/herself, when serious adverse events occur, so that it helps to drive appropriate investigations.

Storage and Transport

For the storage it is recommended dry and airy place, without exposure to light incidence, humidity or contaminants substances.

The devices cannot be stored directly on the floor. It is recommended the use of shelves with minimal height of 20 cm.

As it is a sterile product, the storage place temperature and humidity must be monitored and kept under 40° C.

The product should be kept in its original packaging until the moment of its use, being that the surgical packaging opening and handling should be done by trained personal for this procedure.

The product should be transported properly, avoiding falls and friction that can damage the structure and surface of the part.

For information about manufacturing date and batch number: see label.

Further Information

Manufactured and distributed by:

MDT – Indústria Comércio Importação e Exportação de Implantes SA

Address: Av. Brasil, nº. 2983 - Distrito Industrial - Rio Claro/SP - Brasil

CEP: 13505-600

Phone/ Fax: (55-19) 2111-6500 **CNPJ:** 01.025.974/0001-92

Technician Responsible: Miguel Lopes Monte Júnior – CREA 0601150192 **CE 0434** (in accordance with EC Directive 93/42/EEC). When applicable.

ANVISA Registration No: 10417940089

Review: 00

Issue: December 23rd, 2013.



ALERT INSTRUCTIONS FOR USE

These INSTRUCTIONS FOR USE are not available in printed form, they are available at the website of the manufacturer www.mdt.com.br.

The INSTRUCTIONS FOR USE on the website are indexed by REGISTRATION/ CADASTRE ANVISA's NUMBER and its TRADE NAME of the product, informed at the label of the product purchased.

All the INSTRUCTIONS FOR USE provided on the website have the identification of the revision and date of emission of the document. The user must pay attention to the correct version (revision and date of emission) of the document regarding the MANUFACTURING DATE informed at the label of the product purchased.

If it is interest to the user, the INSTRUCTIONS FOR USE may be provided in printed form, with no extra cost and the request of the same shall be held by the CAP (Customer Service Department) manufacturer, as following:

Customer Service Department - CAP:

Telephone: +55 19 2111.6500

FAX: +55 19 2111.6500

http://www.mdt.com.br/contato

Avenida Brasil, 2983 - Distrito Industrial CEP: 13505-600 | Rio Claro - São Paulo - Brasil

Opening Hours: 8 AM to 5 PM, from Monday to Friday, except holidays.