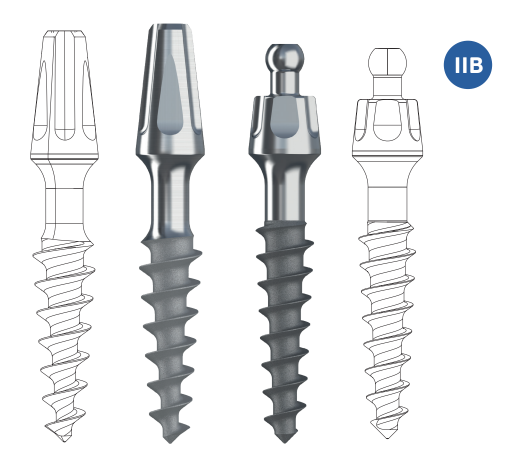
MAXIFIX COMPRESSIVE/P



MAXIFIX COMPRESSIVE is recommended for areas with very low bone density. Maxifix Compressive has been specifically designed and researched to avoid fenestration. The implant has a ground surface which can be bent up to 30°, a gradually expanding spiral and a groove which allows blood to drain with a new self-functioning, self-blocking tip.

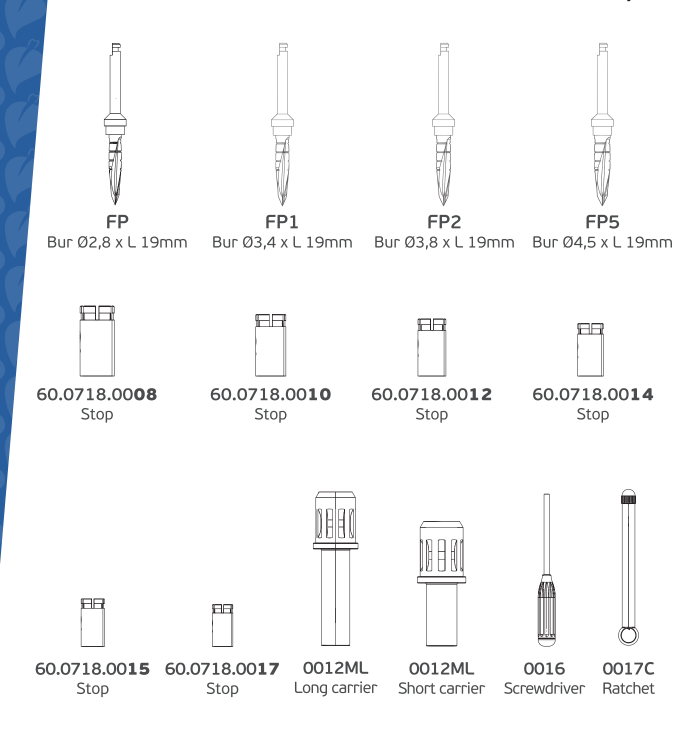
MAXIFIX COMPRESSIVE P is an easy-to-use-stage implant, with self-tapping spiral, made of Ti °5 titanium, which is perfectly biocompatible. Ideal for cases of dental agenesis and removable denture stabilisation.

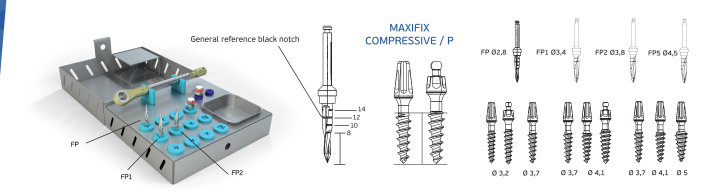




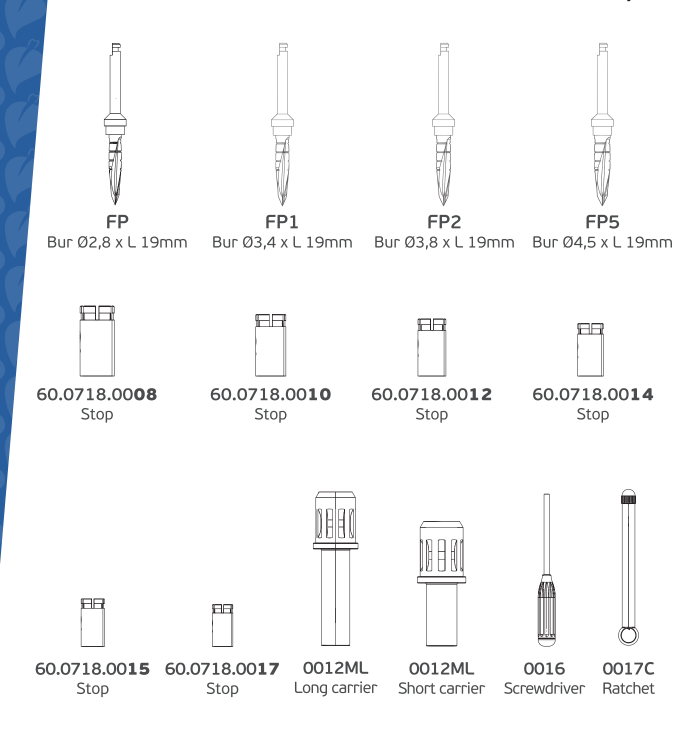


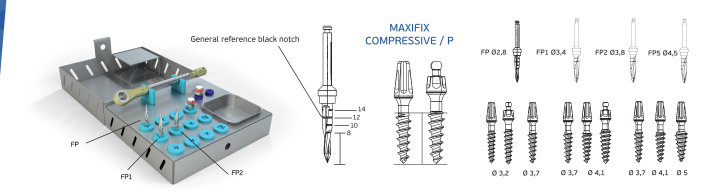
KIT MAXIFIX COMPRESSIVE/P

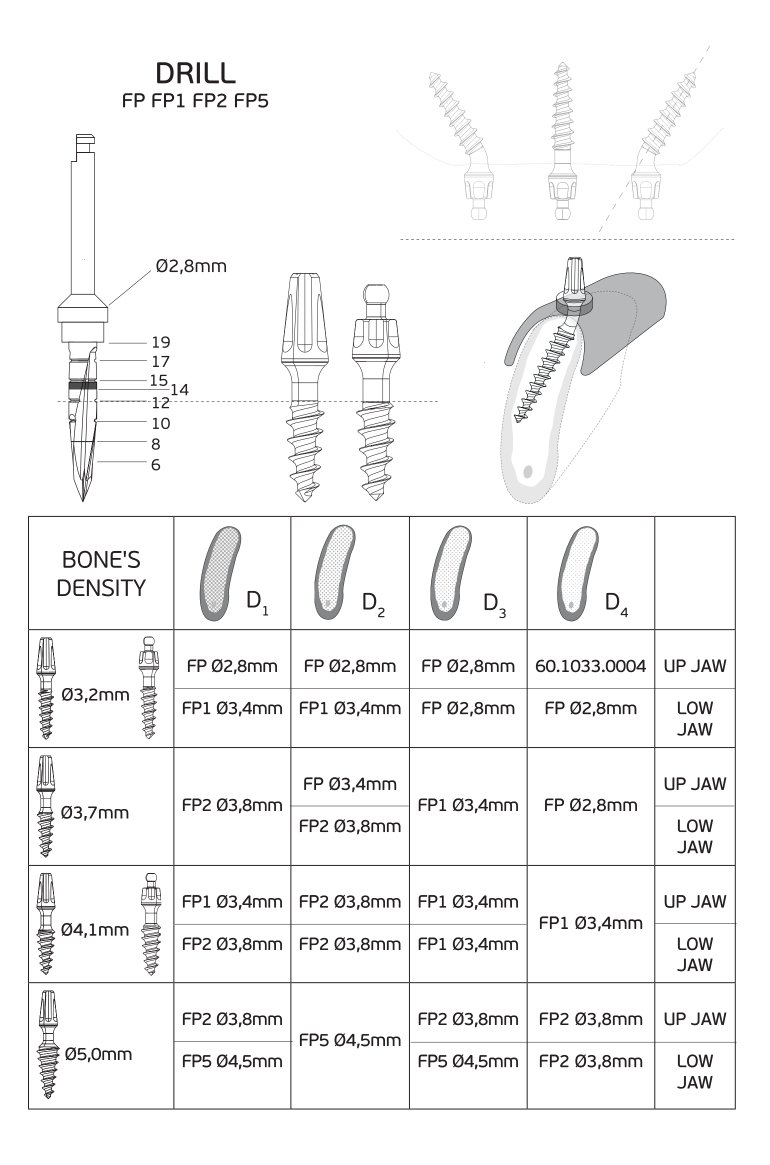




KIT MAXIFIX COMPRESSIVE/P







- The decision to use an NSI implant or prosthetic component cannot be made without an adequate diagnostic, clinical and radiological evaluation.
- NSI implants must be used exclusively with the surgical instruments designed by NSI and with the relative prosthetic components. The use of incorrect instruments or prosthetic parts could cause serious damage to the patient and the professional and in this case NSI does not guarantee the total success of the surgery both in the operative period and in the postoperative period.

Packaging

NSI prosthetic components are packaged as follows:

- Prosthetic component;
- Initial plastic ampoule with closure label and product label;
- Instructions for use Box with removable product label to be placed on the patient file.

The prosthetic components have an expected lifetime of about 10 years; appropriate laboratory tests have been conducted for this purpose. This period may decrease if the patient does not adopt the adequate daily hygiene and cleaning measures specified in the following paragraphs and in the case of abnormal chewing loads or undesirable impacts to the prosthetic components or to the entire dental arch.

Attention: The product is not sterilized but is packaged after a decontamination treatment in a controlled environment. This treatment ensures that the product will remain in hygienic conditions and clean if the package remains intact and is properly stored at a temperature between -5°C and 50°C. The manufacturer declines all liability for any product sterilization carried out by external personnel.

Directions for use

The success of the coupling between the endosseous NSI implants and the relative prosthetic components depends both on the correct application of the phases indicated in the Surgical Technique and on an adequate preoperative evaluation carried out by the professional. It is important that the dentist perform a thorough screening before surgery to obtain adequate information about the patient and the type of implant and prosthetic component to be used. In particular:

- General medical history, as indicated below.
- Local history regarding peri-implant tissues and evaluation of possible pre-implant therapies.
- Bone characteristics by means of specific radiological examinations.
- Evaluation of the growth curve.
- Evaluation of the possible simultaneous treatment of the two dental arches.

During surgery, it is essential that the conditions of the operating field ensure an adequate level of hygiene and cleanliness and that the product is handled with particular attention so as not to damage it and thus compromise the success of the intervention.

Preoperative and intraoperative procedures

Meticulous sterility, careful disinfection, accurate application of the NSI Surgical Technique.

Uncover the surgical screw, insert the healing screw, careful disinfection of the oral cavity with suitable solutions. Before the patient's discharge, it may be necessary to administer an analgesic and anti-inflammatory drug and check hemostasis.

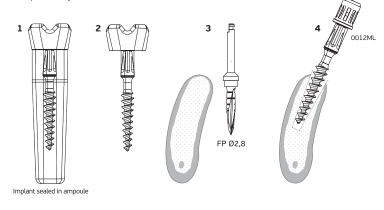
After 10 - 15 days, take the impression with a pick-up impression. Once the plaster model is obtained, choose the abutment.

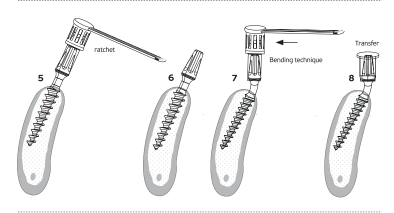
TIGHTENING THE THROUGH SCREW with a dynamometric ratchet to 30 to 35 Ncm, let rest for 2 minutes, unscrew and re-tighten at the sa me intensity. This will allow the passivation of the internal tensions guaranteeing stable tightening and coupling.

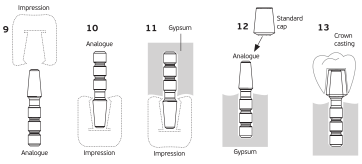
Rev. 7 del 10.12.2019

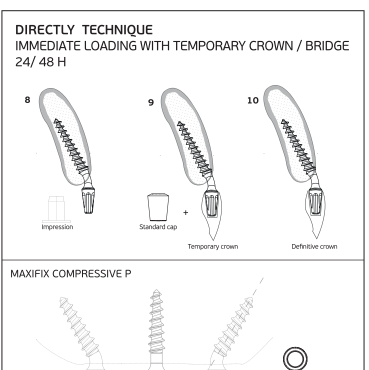
NSI's surgical protocol & prosthetic dental implant

NSI Srl patented system













IMPLANT

NSI srl Via Vittorio Emanuele II,1 25122 BRESCIA ITALIA	
Labeling Identification	
•••	Symbol for Legal Manufacturer
REF	Symbol for Catalogue number
LOT	Symbol for Batch code
SIZE	Symbol for Use by Prescription Only
Rx only	Symbol for Use by Prescription Only
<u>i</u>	Consult Instructions for Use www.ifu.biomet3i.com
	Symbol for Do not re-use
<u></u>	Symbol for Instruction inside
<i>₩</i>	Symbol for Date of Manufacture
	Symbol for Do not use if package is damaged
C€	CEE Approuved
MD Medical Device	Medical Device
Ť	Keep dry
类	Keep away from sunlight
STERILE R	Sterized using irradiation
STERNLIZE	Do not resterilize
UDI	Unique Device Identification
	QR code
	SERIAL NUMBER