TRYON® is the classic implant of S.I.N. The implant that has external hexagon, double acid etched and ensures high primary stability, effectiveness and safety for your surgeries.

**INDICATIONS OF USE**

S.I.N. Dental Implant System is intended for placement in the maxillary or mandibular arch to provide support for single-unit or multi-unit restorations. When a one-stage surgical approach is applied, the implant may be immediately loaded when good primary stability is achieved and the functional load is appropriate.

**PRODUCT DESCRIPTION**

TryOn implants are manufactured from unalloyed titanium conforming to ASTM F67, Grade 4, and are provided with an acid etched surface treatment. Provided STERILE. Sterilized by irradiation.

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<tr>
<th>Diameter of Implants (mm)</th>
<th>Platform Diameter (mm)</th>
<th>Length (mm)</th>
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<tbody>
<tr>
<td>3.25, 3.5, 3.75, 4.0, 5.0</td>
<td>4.1, 5.0</td>
<td>7.0, 8.5, 10, 11.5, 13, 15</td>
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Tryon implants are for conventional 1-stage and 2-stage surgical technique and immediate loading.

**CONTRAINDICATIONS**

S.I.N. Dental Implant System is contraindicated in the following conditions:
- Inability to build a functional prosthesis.
- Rehabilitation with dental implants is also contraindicated for children, pregnant women and during breastfeeding.

**WARNINGS**

The surgical technique of dental implant installation is highly specialized and the surgical procedure complex, it is recommended that the professionals be technically qualified so that the application of the S.I.N. implants is safe and efficient.

Product is for professional use only.

Product is sterilized by gamma radiation. Sterility is ensured except in cases where the package has been violated or damaged. Do not use if the package is damaged or after the expiration date.

Single use only. Do not re-sterilize.

The reuse or re-sterilization of this product can cause damage to health.

S.I.N. Dental Implant System has not been evaluated for safety and compatibility in the Magnetic Resonance (MR) Environment. It has not been tested for heating, migration, or image artifact in the MR environment. The safety of S.I.N. Dental Implant System in the MR environment is unknown. Scanning a patient who has this device may result in patient injury.

**PRECAUTION**

Before implant installation, to obtain a predictable long-term outcome, the professional must submit the patient to a detailed and careful medical history, examination, radiographs, laboratory tests, and study models for appropriate planning.
ADVERSE EFFECTS
Loss of the implant and prosthesis is possible due to a number of reasons, including implant contamination, inappropriate surgical technique, poor bone quality, inappropriate oral hygiene, and parafunctional habits (tooth grinding).

SURGICAL COMPLICATIONS
The implant installation surgical procedure may bring risks during and after the surgery, such as: pain, edema, hemorrhage, dehiscence, paresthesia, and infection.

SHIPMENT AND HANDLING
The S.I.N. implants are sent to professionals duly packaged, sealed and sterilized. Therefore, the package must be opened using sterile technique, and must be handled only with sterilized titanium instruments.
ATTENTION
In order to obtain technical support or additional information material about the product, contact: SIN - Sistema de Implante Nacional S.A. Contact details are provided at the end of these instructions.

INSTRUCTIONS FOR USE
Note: During all drilling to shape the implant site, avoid deflecting the drill sideways, and use continuous, copious irrigation.
S.I.N. implants were designed for a maximum torque of 80 Ncm. Higher torques may cause irreversible damages to the implants as well as surgical complications.
The torque for intermediary fixation (cemented abutment, conic abutment or mini-abutment) on the implant is 20 Ncm. The torque for component fixation on the intermediaries is 10 Ncm.
Do not install the protection screw (cover screw) with the ratchet wrench or torque meter since this may damage the implant; tighten it manually with a digital driver.

1. At the surgical site penetrate the cortical bone with the initial drill, Ø 2.0 mm (1500 RPM).
   Prepare the surgical site with the Ø 2.0 mm helical drill (1500 RPM).
   Use the pilot drill to enlarge the surgical site from Ø 2.0 mm to Ø 3.0 mm.
   Use the Ø 3.0 mm helical drill to the complete the surgical site preparation.
   Optionally, use the countersink drill according to the implant placement level (800 RPM).
   For the surgical sequence for each implant diameter, refer to the guide in the catalog.
2. Remove the adhesive part of the package and the inner tray containing the dental implant.
   Place the inner tray over a surgical tray or organizer.
   Remove the Tyvek label, exposing the implant.
3. Attach the wrench (CTIT 20) to the contra-angle.
   Attach the wrench CTIT 20 to the internal hex of the TryOn implant.
   Optionally, use the CTIT 24 to contra-angle and the CCIT 20 and CCIT 24 to ratchet wrench to the installation of the TryOn implant.
4. Take the assembled implant set to the previously prepared surgical site, and start the implant installation at a low speed (20 RPM).
5. If necessary, complete the installation with the ratchet wrench (TMECC or TMECC 02 attached to the CCIT 20 or CCIT 24).
6. For delayed loading procedures, after placing the implant, use the 0.9 mm hexagonal drive (CDH 0920 or CDH 0924) to place the cover screw. After installation of the cover screw, suture the gingiva.
7. For a single-stage surgical procedure or immediate loading, install the selected intermediary abutment and prosthetic components.
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<th>ATENÇÃO</th>
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<td>KEEP DRY</td>
<td>MANTÉNGALO SECO</td>
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<td>KEEP AWAY FROM SUNLIGHT</td>
<td>MANTÉNGALO LEJOS DE LA LUZ SOLAR</td>
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<td>DO NOT USE IF PACKAGE IS DAMAGED</td>
<td>NO LO UTILICE SI EL ENVOLTORIO ESTÁ DAÑADO</td>
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<td>DO NOT RESTERILIZE</td>
<td>NO LO REESTERILIZAR</td>
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<td>ATENÇÃO: A LEI FEDERAL (EUA) LIMITA A VENDA DESTE DISPOSITIVO POR OU POR ORDEM DE UM PROFISSIONAL DE SAÚDE LICENCIADO.</td>
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<tr>
<td>CAUTION: FEDERAL LAW (USA) RESTRICTS THIS DEVICE TO SALE BY OR ON THE ORDER OF A LICENSED HEALTHCARE PRACTITIONER.</td>
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<tr>
<td>PRECAUCIÓN: LAS LEYES FEDERALES (USA) RESTRINGEN LA VENTA DE ESTE DISPOSITIVO POR O EN EL ORDEN DE UN PROFESIONAL DE LA SALUD LICENCIADO.</td>
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**DEVELOPED AND MANUFACTURED BY:**
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**PRODUCT:**
Tryon Implant

**510 (k) FDA-US:**
170208