Guided (Pilot Drill) Sleeves, Guided Anchor Pin Sleeves and Mounting Tools Instructions for use







Guided Pilot Drill Sleeves Guided Anchor Pin Sleeves

Important: Please read.

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Description:

Guided (Pilot Drill) Sleeves and Guided Anchor Pin Sleeves:

The Guided (Pilot Drill) Sleeves are highly precise cylinders embedded in dental surgical templates to define the position, direction and height/depth of surgical sites. The surgical template is a plastic device that is to be fitted onto the soft tissue of the patient. In partial and single cases, it also fits onto the remaining teeth of the jaw. The Guided (Pilot Drill) Sleeves have the long axis identical to the planned long axis of the implant. The level of the outer shoulder of the Guided (Pilot Drill) Sleeve defines the depth of the surgical preparation and implant position since there is a pre-defined relationship between this level and the implant/abutment interface.

Also, embedded in the surgical template are Guided Anchor Pin Sleeves. These sleeves guide the preparation and installation of anchor pins. The anchor pins are thin rods of metal positioned close to horizontally into the jawbone to secure the surgical template in its intended position during implant surgery. Guided Anchor Pin Sleeves are optional.

Mounting Tools for Guided (Pilot Drill) Sleeves:

In decentralized/local production of surgical templates, the Mounting Tools for Guided (Pilot Drill) Sleeves can be used to glue the Guided (Pilot Drill) Sleeves into the printed/ milled/stereolithography (SLA) produced surgical template by the laboratory or dentist. The Mounting Tools for Guided (Pilot Drill) Sleeves will position the Guided (Pilot Drill) Sleeves accurately flush with the top shoulder of the surgical template's sleeve support material. For fixation of the sleeves a biocompatible glue / cement / adhesive agent is needed

The Mounting Tool Pins for Guided Pilot Drill Sleeves have either a 1.5 mm or 2.0 mm diameter and the Mounting Tool Base is designed to be used with either diameter. Due to their design, the Mounting Tools for Guided Pilot Drill Sleeves cannot be used in combination of pre-fabrication of master casts.

The Mounting Tools for Guided Sleeves (guided cylinder with pin, implant replica) can be used in a dental lab procedure when making a stone model based on the surgical template. The guided cylinders have an outer diameter that matches the internal dimensions of the Guided Sleeves (NP, RP, and 6.0/WP) and are inserted into the Guided Sleeves, embedded in the surgical template, and an implant replica from the Nobel Biocare standard assortment is screwed onto the pin that goes through the guided cylinder. Since there is a well-defined seating of the guided cylinder in the Guided Sleeve, the head of the implant replica is positioned in the same position in relationship to the surgical template, as the implant will be installed into the bone of the patient. This enables a stone model to be cast using the underside of the surgical template, corresponding to the soft tissue surface of the patient, and the attached implant replicas, corresponding to the pre-defined sites of the implants. The implant connection portion of the guided cylinder should match the available implant connections of tri-channel, external hex and conical connection.

In the analog workflow, the product is attached together with a Guided Sleeve to the implants installed in the die cast model. In doing so, the Guided Sleeves are positioned in a correct position and ready to be molded into a lab-made surgical template.

Intended use:

- The Guided (Pilot Drill) Sleeves are intended to be used in combination with surgical templates to facilitate implant installation by guiding the required instruments in the planned direction
- The Guided Pilot Drill Sleeves are intended to guide the first drill (either 1.5 mm or 2.0 mm)
- The Mounting Tools are intended to be used in the dental laboratory procedure to enable the correct positioning of the Guided (Pilot Drill) Sleeves in the surgical template.
- The Guided Anchor Pin Sleeves are intended to be used in surgical templates to establish secure fixation and stability of the surgical template, by guiding the preparation and installation of anchor pins.

Indications:

The components (Guided (Pilot Drill) Sleeves, Guided Anchor Pin Sleeves and Mounting Tools) are indicated for the assembly within a surgical template for the treatment of edentulous and partially edentulous jaws (including patients missing a single tooth) for placement of dental implants, if indicated in combination with immediate function to restore esthetics and functionality (e.g. masticatory, speech). The following prerequisites must be fulfilled:

- Adequate amount of jawbone volume (height and width).
- Quality of jawbone must be judged as adequate.
- Adequate mouth opening (minimum 40 mm) to accommodate guided surgery tooling.
- Exclusion of compromised diseases in conflict with dental implant treatment.

- Adequate compliance.

Contraindications:

It is contraindicated to use a surgical template in:

- Patients who are medically unfit for an oral surgical procedure.
- Patients in whom adequate sizes, numbers or desirable positions of implants are not reachable to achieve safe support of functional or eventually parafunctional loads.
- Patients who are allergic or hypersensitivity to commercially pure titanium grade 4, stainless steel, surgical template material or bonding material.

Warnings:

- Failure to recognize actual lengths of drills relative to radiographic measurements can result in permanent injury to nerves or other vital structures. Drilling beyond the depth intended for lower jaw surgery may potentially result in permanent numbress to the lower lip and chin or lead to a hemorrhage in the floor of the mouth.
- Besides the mandatory precautions for any surgery such as asepsis, during drilling in the jawbone, one must avoid damaging the nerves and vessels by referring to anatomical knowledge and preoperative radiographs.
- Failure to use the right size of sleeve results in the tooling not fitting.

Cautions:

General:

It is strongly recommended that Nobel Biocare Guided (Pilot Drill) Sleeves and Mounting Tools are used only with Nobel Biocare surgical instruments, implants, prosthetic components, and with surgical templates designed using the 3D-planning software DTX Studio Implant as combining components that are not dimensioned for correct mating can lead to mechanical and/or instrumental failure, damage to tissue or unsatisfactory esthetic results.

One hundred percent implant success cannot be guaranteed. Especially, non-observance of the indicated limitations of use and working steps may result in failure.

Treatment by means of implants may lead to loss of bone, biologic or mechanical failures including fatigue fracture of implants.

Close cooperation between surgeon, restorative dentist and dental laboratory technician is essential for a successful implant treatment.

It is strongly recommended that clinicians, new as well as experienced implant users, always go through special training before undertaking a new treatment method. Nobel Biocare offers a wide range of courses for various levels of knowledge and experience. For more info please visit http://www.nobelbiocare.com.

When using a new device/treatment method for the first time, working with a colleague who is experienced with the new device/treatment method may help avoid possible complications. Nobel Biocare has a global network of mentors available for this purpose.

Before surgery:

Careful clinical and radiological examination and diagnosis of the patient must be performed prior to surgery to determine the psychological and physical status of the patient.

Special attention should be given to patients who have local or systemic factors that could interfere with the healing process of either bone or soft tissue or the osseointegration process (e.g., cigarette smoking, poor oral hygiene, uncontrolled diabetes, oro-facial radiotherapy, steroid therapy, infections in the neighboring bone). Special caution is advised in patients who receive bisphosphonate therapy.

In general, implant placement and prosthetic design must accommodate individual patient conditions. In case of bruxism or unfavorable jaw relationships reappraisal of the treatment option may be considered.

Pre-operative hard tissue or soft tissue deficits may yield a compromised esthetic result or unfavorable implant angulations.

All components, instruments and tooling used during procedure must be maintained in good condition and care must be taken that instrumentation does not damage implants or other components

At surgery:

Particular caution should be used when placing narrow platform implants in the posterior region due to risk of prosthetic overload. Care and maintenance of instruments are crucial for a successful treatment.

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Because of the small sizes of the devices, care must be taken that they are not swallowed or aspirated by the patient.

After the implant installation, the surgeon's evaluation of bone quality and primary stability will determine when implants may be loaded. Lack of adequate quantity and/or quality of remaining bone, infection and generalized diseases may be potential causes for failure of osseointegration both immediately after surgery, or after osseointegration is initially achieved.

After surgery:

To secure the long-term treatment outcome, it is advised to provide comprehensive regular patient follow up after implant treatment and to inform about appropriate oral hygiene.

Handling procedure:

1. Check the decentralized/locally produced surgical template.

- Check the sleeve seats for material residues and sharp, protruding edges. Remove or smoothen them if you find any.
- Check that the locally produced surgical template is manufactured from appropriate material: the material should be biocompatible and mechanically fit for purpose.
 Recommended material properties are listed in table 1.
- Verify optimal fit on stone model if applicable and/or in patient's mouth prior to surgery.

2a. Fixing the Guided Pilot Drill Sleeves into the surgical template.

Insert the Guided Pilot Drill Sleeve (1.5 mm – single pack article 300438, 20 pack article 300439; 2.0 mm – single pack article 300440, 20 pack article 300441) into the sleeve seat of the surgical template.

Note: Make sure the flat upper part of the sleeve is on the occlusal surface of the surgical template.

– For the correct fixation of the Guided Pilot Drill Sleeve; use the screw part of the Mounting Tool (Mounting Tool Pin for Guided Pilot Sleeve 1.5 mm – article 300442; Mounting Tool Pin for Guided Pilot Sleeve 2.0 mm – article 300443) from below (intaglio/inside of the surgical template) and the thread part (Mounting Tool Base for Guided Pilot Sleeve 1.5/2.0 mm – article 300444) from the top (occlusion/outer surface of the surgical template) (see picture 2). Tighten it by hand.

Note: The outer diameter of the Guided Pilot Drill Sleeves is the same. Please confirm that the correct sleeve size is used for each position by comparing the laser marking of the Guided Pilot Drill Sleeves with the planning overview before inserting it into the template (see picture 1).



Picture 1: Example of a comparison of laser marking on Guided Pilot Drill Sleeves with planning overview

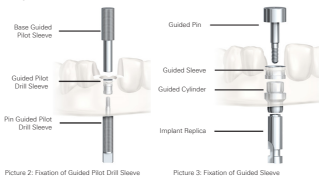
PID	Sleeve name	Article id
1	Guided Pilot Drill Sleeve 1.5 mm	300438
2	Guided Pilot Drill Sleeve 2.0 mm	300440

2b. Fixing the Guided Sleeve into the surgical template.

- Insert the Guided Sleeve (NP article 32754; RP article 32765 or WP article 32766) into the sleeve seat of the surgical template.
- Note: As the Guided Sleeve is symmetrical, there is no top or bottom.
- For the correct fixation of the Guided Sleeve; use the upper part of the Guided Cylinder with Pin (NP article 37172; RP article 37173; WP article 37950) and its lower part to fix the Guided Sleeve. Tighten the lower part of the Guided Cylinder with Pin to the appropriate Implant Replica (NP article 36697; RP article 36698; WP article 37879) (see picture 3). Fix all parts together and tighten them by hand or by using a screwdriver (e.g. Uniqrip™).

2c. Fixing the Guided Anchor Pin Sleeve.

- Insert the Guided Anchor Pin Sleeve (article 30908) into the sleeve seat of the surgical template.
- Note: As the Guided Anchor Pin Sleeve is symmetrical, there is no top or bottom.
- Make sure that the upper part of the sleeve is set flush into the seat socket of the surgical template (see picture 4)





Picture 4: Insertion of Anchor Pin Sleeve

3. Bonding process for the Guided (Pilot Drill) Sleeves and Guided Anchor Pin Sleeves

 Visually verify that the sleeves are flush with the top surrounding surface of the surgical template. If they are not flush, remove material as required.

For permanent fixation of the sleeves a biocompatible glue / cement / adhesive agent is needed. The user must use a biocompatible material and follow the manufacturer's instructions for use. Recommended material properties are listed in table 2.

Note: Once all the Guided (Pilot Drill) Sleeve(s) and Guided Anchor Pin Sleeve(s) are in place, glue them into the surgical template.

- Place the tip of the mixing cannula inside the glue hole located in the surgical template.
- Slowly squeeze the bonding material into the glue hole until it completely goes all the way around the Guided (Pilot Drill) Sleeve(s) or Guided Anchor Pin Sleeve(s).
- Repeat the application of bonding material to each Guided (Pilot Drill) Sleeve(s) and Guided Anchor Pin Sleeve(s).
- Once the bonding material has hardened, unscrew the Mounting Tools.
- Visually verify that there is no bonding material present on the top or bottom of the Guided (Pilot Drill) Sleeve(s) or Guided Anchor Pin Sleeve(s).

Dual curing is advised to control the timing of the curing process and ensure material is fully cured.

Caution: Introduce only as much bonding material to cover the outer diameter of the Guided (Pilot Drill) Sleeve(s) or Guided Anchor Pin Sleeve(s) in the glue channel. While introducing the bonding material, you should be able to see the progression of it. Avoid to introduce too much bonding material. Leaked bonding material should be removed immediately using a suitable instrument.

Warning: Do not introduce bonding material into the internal diameter of the Guided (Pilot Drill) Sleeve(s) or Guided Anchor Pin Sleeve(s). This could have a negative effect on the guided surgery.

For additional information on surgical procedures please consult the NobelGuide Concept Manual available at <u>www.nobelbiocare.com</u> or request the latest printed version from a Nobel Biocare representative.

Materials:

The Guided (Pilot Drill) Sleeves and Guided Anchor Pin Sleeves are made from stainless steel. The Mounting Tools for the Guided Pilot Drill Sleeves are made of stainless steel. The Mounting Tools for the Guided Sleeves are made of titanium.

Recommendation for materials:

For surgical template fabrication:

The user must use a certified material, intended for printing surgical guides and follow the manufacturer's instructions for use, as well as the recommended parameters and procedures.

The minimum mechanical properties for the surgical template material are provided in table 1. In addition, the material must be biocompatible.

Postcured Data	Metric	Method
Tensile Strength	≥41 MPa	ASTM D 638
Tensile Modulus	≥2030 MPa	ASTM D 638
Elongation at Break	4 - 7 %	ASTM D 638
Heat Deflection Temperature		ASTM D 638
at 66Psi	46°C	
at 264Psi	41°C	
Flexural Strength	≥50 MPa	ISO 20795-1/ ASTM D 790
Flexural Modulus	≥1500 MPa	ISO 20795-1/ ASTM D 790
Hardness Shore	≥80 D	ASTM D2240

Table 1. Minimum mechanical properties of surgical template material

For sleeves bonding into the surgical template:

The minimum requirements for the mechanical properties of the glue to be used for bonding the sleeves into the surgical template are provided in table 2.

Mechanical property	Accepted range/level
Compressive Strength	≥ 200 MPa
Flexural Strength	> 2000 MPa
Water Sorption	≤ 200 µg/mm³

Table 2. Minimum mechanical properties of bonding material

Additional requirements for the bonding material:

- The bonding material should be biocompatible and suitable for dental applications.
- The bonding material should be able to bond metals to polymers.

Sterility and Reusability Information:

Delivered non-sterile for single use:

Guided (Pilot Drill) Sleeves and Guided Anchor Pin Sleeves are delivered non-sterile for single use only and do not need to be sterilized prior to use.

Caution: Guided (Pilot Drill) Sleeves and Guided Anchor Pin Sleeves are single use products and must not be reprocessed. Reprocessing could cause loss of mechanical, chemical and / or biological characteristics. Re-use could cause cross contamination.

Cleaning and sterilization instructions:

Surgical Template construct cleaning:

Following fixation of the Guided (Pilot Drill) Sleeves and (optionally) Guided Anchor Pin Sleeves into the surgical template, the surgical template construct should be cleaned and disinfected, as applicable per the guide material manufacturer's instructions, before intraoral use. The cleaning protocol selected should be appropriate for the surgical guide material, according to the material manufacturer's instructions, Guided (Pilot Drill) Sleeves and Guided Anchor Pin Sleeves are resistant to mild pH enzymatic detergents (e.g Cidezyme/ Enzol), high-level disinfectants (e.g. Cidex OPA), 0.5% of alkaline cleaning agents (e.g. neodisher Mediclean), and temperatures up to and including 90°C. For further information please contact a Nobel Biocare representative.

Cleaning of mounting tools:

Delivered non-sterile for multiple use to be used extra orally only.

Mounting Tools are used only in the dental laboratory (no intraoral use) and have no cleaning and sterilization requirements.

Magnetic Resonance (MR) safety information:

The Guided (Pilot Drill) Sleeves and Guided Anchor Pin Sleeves have not been evaluated for safety and compatibility in the MR environment. It has not been tested for heating, migration, or image artifact in the MR environment. The safety of Guided (Pilot Drill) Sleeves and Guided Anchor Pin Sleeves in the MR environment is unknown. Scanning a patient who has this device may result in patient injury.

For additional Information on Magnetic Resonance Imaging, please consult the "Cleaning & Sterilization Guidelines including MRI information of Nobel Biocare Products" available at www.nobelbicoare.com or request latest printed version from a Nobel Biocare representative.

Storage and handling:

The device must be stored and transported in dry conditions in the original packaging at room temperature and not exposed to direct sunlight. Incorrect storage and transportation may influence device characteristics leading to failure.

Disposal:

Disposal of the device shall follow local regulations and environmental requirements, taking different contamination levels into account.

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Canada license exemption: Please note that not all products may have been licensed in accordance with Canadian law.

Prescription device: Rx only

Caution: Federal law restricts this device to sale by or on the order of a licensed physician or dentist.







SN



Serial number



barrier system

Patient

identification



-

website

3/3

Single sterile barrier system barrier system with protective with protective packaging inside packaging outside



irradiation



UDI

Tooth number



PHT

Contains or

presence of

Do not re-use

phthalate

DEHP



ethylene oxide

STERILE



Upper limit of temperature

Sterilized using Unique Device steam or dry heat Identifier

Use-by date

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Double sterile

barrier system

Keep drv

Manufacturer



Health care centre or doctor



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symbol.glossary.nobelbiocare.com
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Link to Online Symbols Glossary and IFU Portal







Non-pyrogenic





Patient information Patient number



IFU2009 016 01





Symbols Glossary:

the applicable symbols.

representative in the

European Community

EC REP

Authorized

CE marking

Date

6

is damaged

Keep away from

Magnetic resonance

sunlight

MR

conditional

Do not use if package

The following symbols may be present on the device labeling or in information

LO

Consult

instructions for use

Batch code

accompanying the device. Refer to the device labeling or accompanying information for

REF

Contains

hazardous

substances

Do not resterilize

Rx Only

use only

Catalogue number