NobelParallel™ Conical Connection
Instructions for use

Important: Please read.

Disclaimer of liability:
This product is part of an overall concept and may only be used in conjunction with the associated original products according to the instructions and recommendation of Nobel Biocare. Non-recommended use of products made by third parties in conjunction with Nobel Biocare products will void any warranty or other obligation, express or implied, of Nobel Biocare. The user of Nobel Biocare products has the duty to determine whether or not any product is suitable for the particular patient and circumstances. Nobel Biocare disclaims any liability, express or implied, and shall have no responsibility for any direct, indirect, punitive or other damages, arising out of or in connection with any errors in professional judgment or practice in the use of Nobel Biocare products. The user is also obliged to study the latest developments in regard to this Nobel Biocare product and its applications regularly. In cases of doubt, the user has to contact Nobel Biocare. Since the utilization of this product is under the control of the user, they are his/her responsibility. Nobel Biocare does not assume any liability whatsoever for damage arising thereof.
Please note that some products detailed in this Instruction for Use may not be regulatory cleared, released or licensed for sale in all markets.

Description:
Implant:
NobelParallel™ Conical Connection (CC) dental implants are made from biocompatible commercially pure grade 4 titanium with TiUnite® surface. The implant comes with a co-packed Cover Screw made of titanium alloy Ti-6Al-4V.

Tooling:
Nobel Biocare Twist Drills, Twist Step Drills, Cortical Drills and Screw Taps are made of stainless steel with an amorphous diamond coating and should be used in conjunction with NobelParallel™ CC implants.

Intended use:
NobelParallel™ CC implants are intended to be used in the upper or lower jaw bone (osseointegration) and used for anchoring or supporting tooth replacements to restore chewing function.

Indications:
NobelParallel™ CC implant restorations range from single tooth to fixed-removable full dental arch overdenture applications to restore chewing function. This can be achieved by a 2-stage or 1-stage surgical technique in combination with immediate, early or delayed loading protocols, recognizing sufficient primary stability and appropriate occlusal loading for the selected technique. Implants allow also for bi-cortical anchorage in cases of reduced bone density to obtain high initial stability.

Contraindications:
It is contraindicated placing NobelParallel™ CC implants in patients:
- who are medically unfit for an oral surgical procedure.
- with inadequate bone volume unless an augmentation procedure can be considered.
- in whom adequate sizes, numbers or desirable positions of implants are not reachable to achieve safe support of functional or eventually parafunctional loads.
- allergic or hypersensitive to commercially pure titanium grade 4 or titanium alloy Ti-6Al-4V (titanium, aluminium, vanadium).

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 from lower jaw surgery may potentially result in permanent numbness 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 jaw bone, one must avoid damage the nerves and vessels by referring to anatomical knowledge and preoperative radiographs.

Cautions:
General:
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 NobelParallel™ CC implants are used only with Nobel Biocare surgical instruments and prosthetic components, 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. 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 www.nobelbiocare.com.

Before surgery:
Careful clinical and radiological examination of the patient has to be performed prior to surgery to determine the physiological and physical status of the patient. Special attention has to be given 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).

After the implant installation, the surgeon's evaluation of bone quality and initial 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.

Surgical procedure:
1. During drilling procedures bone quality should be considered (please see table A). recommended drill sequences are based on bone quality to ensure optimal primary stability when applying immediate function.

A NobelParallel™ CC Recommended drill sequence based on bone quality. Drill data are stated in mm and the drills within square brackets denote as optional.

<table>
<thead>
<tr>
<th>Platform</th>
<th>Implant diameter</th>
<th>Soft Bone Type IV</th>
<th>Medium Bone Type II-III</th>
<th>Dense Bone Type I</th>
</tr>
</thead>
<tbody>
<tr>
<td>NP</td>
<td>3.75</td>
<td>2.0</td>
<td>2.2/2.4</td>
<td>2.0</td>
</tr>
<tr>
<td></td>
<td></td>
<td></td>
<td>[3.2/3.6]</td>
<td>[2.4/2.8]</td>
</tr>
<tr>
<td></td>
<td></td>
<td></td>
<td>[Screw Tap 3.75]</td>
<td>Cortical Drill 3.76</td>
</tr>
<tr>
<td></td>
<td></td>
<td></td>
<td>[Screw Tap 5.0]</td>
<td>Cortical Drill 3.76</td>
</tr>
<tr>
<td>RP</td>
<td>4.3</td>
<td>2.0</td>
<td>2.0/2.4</td>
<td>2.0</td>
</tr>
<tr>
<td></td>
<td></td>
<td></td>
<td>[3.2/3.6]</td>
<td>2.2/2.4</td>
</tr>
<tr>
<td></td>
<td></td>
<td></td>
<td>[Cortical Drill 4.3]</td>
<td>2.2/2.4</td>
</tr>
<tr>
<td></td>
<td></td>
<td></td>
<td>[Screw Tap 4.3]</td>
<td>[Screw Tap 4.3]</td>
</tr>
<tr>
<td>RP</td>
<td>5.0</td>
<td>2.0</td>
<td>2.0/2.4</td>
<td>2.0</td>
</tr>
<tr>
<td></td>
<td></td>
<td></td>
<td>[3.2/3.6]</td>
<td>2.2/2.4</td>
</tr>
<tr>
<td></td>
<td></td>
<td></td>
<td>[Cortical Drill 5.0]</td>
<td>2.2/2.4</td>
</tr>
<tr>
<td></td>
<td></td>
<td></td>
<td>[Screw Tap 5.0]</td>
<td>[Screw Tap 5.0]</td>
</tr>
<tr>
<td>WP</td>
<td>5.5</td>
<td>2.0</td>
<td>2.0/2.4</td>
<td>2.0</td>
</tr>
<tr>
<td></td>
<td></td>
<td></td>
<td>[3.2/3.6]</td>
<td>2.2/2.4</td>
</tr>
<tr>
<td></td>
<td></td>
<td></td>
<td>[Cortical Drill 5.5]</td>
<td>2.2/2.4</td>
</tr>
<tr>
<td></td>
<td></td>
<td></td>
<td>[Screw Tap 5.5]</td>
<td>[Screw Tap 5.5]</td>
</tr>
</tbody>
</table>

Note: all data is stated in mm.
Drilling must proceed at high speed (max. 2’000 rpm/min. for step/twist drills) under constant and profuse irrigation by sterile saline at room temperature. In dense bone situation drill with continuous back and forth motion.

Depth measurement system: The parallel drills have a true depth measurement system. All drills and components are marked to prepare the site to the correct depth and obtain a secure and predictable position.

Caution: Twist/Step Drills extend up to 1mm longer than the implant when seated. Allow for this additional length when drilling near vital anatomical structures (please see image B for drill reference lines).

2. Prepare implant site. When using a flapless approach add-on soft tissue height to drill depth.

3. Measure the final depth of implant site for applicable implant length using depth probe with same measurements as Twist/Step Drills.

4. Open the implant package and pick up the implant from inner casing with implant driver (please see image B). The implants are ideally installed with low speed, max. 25 rpm, using a drilling device.

5. Place and tighten the implant using max. 45 Ncm insertion torque.

Warning: Do not use device if the packaging has been damaged or previously opened.

Caution: Implants, Twist Drills, Twist Step Drills, Cortical Drills and Screw Taps are single use products not intended to be reprocessed. Reprocessing could cause loss of mechanical, chemical and / or biological characteristics. Reuse could cause cross contamination.

Note: Twist/Step Drills indicate actual millimeter lengths and correspond to the implant collar. Final vertical positioning depends on several clinical parameters, including esthetics, tissue thickness and available vertical space.

In situations where adjacent natural teeth interfere with the contra-angle head preventing the drill from reaching the desired depth, a drill extension shaft may be used.

6. Medium and dense bone protocol:

a. In cases of a thick cortical layer or dense bone a Cortical Drill and/or a Screw Tap is mandatory to be able to get the implant fully seated and to release pressure around the implant neck.

b. Select the Cortical Drill and/or use Screw Tap matching the diameter of the implant:

- If Cortical Drill is used: proceed with drilling at high speed max. 2’000 rpm/min and drill to appropriate depth (see image B).
- If Screw Tap is used: place the screw tap into prepared implant site using low speed 25 rpm/min. and drill to appropriate depth (see image B). Switch the drill device with handpiece to reverse mode and remove the Screw Tap.

c. Continue with implant installation until desired position is achieved using max. 45 Ncm of insertion torque.

7. For Immediate Function, the implant should be able to withstand a final torque between 35–45 Ncm.

8. Depending on surgical protocol of choice, place a cover screw or an abutment and suture. See table D for implant specifications.

D Implant specifications

<table>
<thead>
<tr>
<th>Platform</th>
<th>Platform diameter</th>
<th>Implant diameter</th>
<th>Lengths</th>
</tr>
</thead>
<tbody>
<tr>
<td>NP</td>
<td>Ø 3.5</td>
<td>Ø 3.75</td>
<td>7, 8.5, 10, 11.5, 13, 15, 18</td>
</tr>
<tr>
<td>RP</td>
<td>Ø 3.9</td>
<td>Ø 4.3</td>
<td>7, 8.5, 10, 11.5, 13, 15, 18</td>
</tr>
<tr>
<td>WP</td>
<td>Ø 5.1</td>
<td>Ø 5.5</td>
<td>7, 8.5, 10, 11.5, 13, 15, 18</td>
</tr>
</tbody>
</table>

Note: all data is stated in mm.

For additional information on surgical procedures please consult the NobelParallel™ CC “Procedures & products” treatment guidelines available at www.nobelbiocare.com or request latest printed version from a Nobel Biocare representative.


Cleaning and sterilization: NobelParallel™ CC Implants, Twist/Step Drills, Cortical Drills and Screw Taps are delivered sterile and for single use only prior to the labeled expiration date.

Warning: Do not use device if the packaging has been damaged or previously opened.

Caution: Implants, Twist Drills, Twist Step Drills, Cortical Drills and Screw Taps are single use products not intended to be reprocessed. Reprocessing could cause loss of mechanical, chemical and / or biological characteristics. Reuse could cause cross contamination.

MR safety information:

Please note that the product has not been evaluated for safety and compatibility in the MR environment. The product has not been tested for heating or migration in the MR environment.

For additional information on Cleaning and Sterilization and Magnetic Resonance Imaging, please consult the “Cleaning & Sterilization Guidelines for Nobel Biocare Products including MRI Information” available at www.nobelbiocare.com/sterilization or request latest printed version from a Nobel Biocare representative.

Storage and handling:
The product must be stored in a dry place in the original packaging at room temperature and not exposed to direct sunlight. Incorrect storage 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.

Manufacturer: Nobel Biocare AB, Box 5190, 402 26 Västra Hamngatan 1, 411 17 Göteborg, Sweden.
Phone: +46 31 81 88 00. Fax: +46 31 19 31 52. www.nobelbiocare.com

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.

EN All rights reserved.
Nobel Biocare, the Nobel Biocare logotype and all other trademarks used in this document are, if nothing else is stated or is evident from the context in a certain case, trademarks of Nobel Biocare. Product images in this folder are not necessarily to scale.