Multi-unit Abutment Plus and Multi-unit Abutment
Instructions for use

Important: Please read.
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Description:
A premanufactured dental implant abutment to be directly connected to an endosseous dental implant intended for use as an aid in prosthetic rehabilitation.

The Multi-unit Abutment Plus is made of titanium alloy.
The Multi-unit Abutment is made of pure titanium and/or titanium alloy.
Gold Coping Multi-unit may be used if a casted framework is preferred.

Multi-unit Abutment Plus, straight and angled 17° & 30°
Internal conical connection for: NobelActive®, NobelReplace® CC and NobelParallel™ CC.

Multi-unit Abutment, straight and angled 17° & 30°
Internal tri-channel connection for: NobelReplace®, Replace Select™, NobelSpeedy® Replace, NobelReplace® Platform Shift.

External hex connection for: Bräunemark System® and NobelSpeedy® Groovy.


Multi-unit Abutment Non-Engaging, angled 30°
The Multi-unit Abutment Non-Engaging angled 30° is available for use with the All-on-4® treatment concept with guided surgery only.

Internal tri-channel connection for: NobelReplace®, Replace Select™, NobelSpeedy® Replace, NobelReplace® Platform Shift.

External hex connection for: Bräunemark System® and NobelSpeedy® Groovy.

Multi-unit Abutment, straight
Other implant systems: Straumann® Octagon soft tissue level 4.8 and 6.5.
Ankylos® Implant System 3.5, 4.5, 5.5, 7.0 mm. Astra Tech Implant System™ 4.5ST, 5.0ST mm. Carex® Implant System 3.3, 3.8, 4.3, 5.0/6.0 mm.

Intended use:
Dental implant abutments are intended to be used in the upper or lower jaw and used for supporting tooth replacements to restore chewing function. Multi-unit Abutment/Plus in combination with endosseous implants are indicated for multiple unit reconstructions when screw retained prosthetics is preferred.

Indications:
Multi-unit Abutment/Plus is a premanufactured prosthetic component directly connected to the endosseous dental implant and is intended for use as an aid in prosthetic rehabilitation.

Contraindications:
It is contraindicated to use Multi-unit Abutment/Plus 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 hypersensitive to commercially pure titanium or titanium alloy Ti-6Al-4V (titanium, aluminium, vanadium), gold alloy (gold, platinum, palladium, iridium) polypropylene or PBT (Polybutylene terephthalate).

Cautions:
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 www.nobelbiocare.com.

Working the first time with a colleague, experienced with the new device/treatment method, avoids eventual complications. Nobel Biocare has a global network of mentors available for this purpose.

Special attention has to 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, pro-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 bruism or unfavorable jaw relation/appraisal 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 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. Because of the small sizes of the devices, care must be taken that they are not swallowed or aspirated by the patient. It is recommended to use a rubber dam in order to prevent inhalation of loose parts.

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.

Never exceed the recommended maximum prosthetic tightening torque for the abutment screw (see table 1). Tightening of abutment may lead to a screw fracture.

Warning:
Narrow platform implants are not recommended for the posterior region of the mouth due to the risk of prosthetic overload. In the posterior region, the Multi-unit Abutment Plus is to be used only in combination with regular platform (RP) and wide platform (WP) implants.

Caution: Never exceed recommended maximum tightening torque for the abutment screw. Tightening of abutment may lead to a screw fracture.

Handling instructions:

Clinical procedure:

1A. Straight Multi-unit Abutment/Plus:
1. Place appropriate abutment (A:1). Use plastic holder to facilitate the insertion (B). It is recommended to verify the final abutment seating using radiographic imaging.

1B. 17° and 30° Multi-unit Abutment/Plus:
1. Place appropriate abutment (A:2). Use holder to facilitate proper position, as there are several positions available (D). It is recommended to verify the final abutment seating using radiographic imaging.
2. Unscrew holder (D).
3. Tighten the abutment to 15 Ncm using Unigrip™ Screwdriver and Manual Torque Wrench prosthetic (E).

Caution: Never exceed recommended maximum tightening torque for the abutment screw. Tightening of abutment may lead to a screw fracture.
2. Take impression of abutments using open or closed impression tray technique (F). Note: Hand tighten only and close impression coping recess prior to impression taking.

Open tray

F:1

Closed tray

F:2

3. Provisionalize or attach healing caps.

Laboratory procedure:

4. Attach abutment replicas to impression copings.

5. Fabricate a working model with removable gingival material (G).

6. NobelProcera® Implant Bridge Wax-up:

   1. Create implant bridge framework using non-engaging temporary cylinders as a foundation and add pattern resin to fabricate desired framework design (H).
   2. Scan the acrylic framework using the NobelProcera® Scanner according to the tutorial found within the software.
   3. Once precision milled framework is delivered back to lab, veneering material is added (I).

6B. Laboratory – Casted Framework:

   1. Attach Gold Coping Multi-unit to the abutment replicas (I) and reduce the height of the plastic chimney.
   2. Wax up framework around gold copings (J).

Note: The Gold Coping Multi-unit is made from a non-oxidizing alloy. Cracking of porcelain may occur if it is applied directly to the gold coping. Make sure the wax covers the Gold Coping Multi-unit with a minimum wax thickness of 0.5 mm. A reduction to 0.3 mm can be made after casting.

3. Fabricate the restoration framework using standard techniques.

4. Complete framework with ceramic (if applicable).

Clinical procedure:

7. Remove temporary restoration if applicable.

8. Use the Screwdriver Machine Multi-unit and Manual Torque Wrench Prosthetic to verify the tightening of the straight Multi-unit Abutment/Plus according to table 1. Use the Unigrip™ Screwdriver and Manual Torque Wrench Prosthetic to verify tightening of the angulated Multi-unit Abutment/Plus to 15 Ncm.

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<table>
<thead>
<tr>
<th>Abutment (clinical) Screw Tightening Torque</th>
<th>Angulated (17°, 30°)</th>
<th>Angulated (45°, 60°)</th>
</tr>
</thead>
<tbody>
<tr>
<td>Nobel Biocare implant systems</td>
<td>35 Ncm</td>
<td>15 Ncm</td>
</tr>
<tr>
<td>*Astra Tech Implant System™ Aqua</td>
<td>20 Ncm</td>
<td>15 Ncm</td>
</tr>
<tr>
<td>*Astra Tech Implant System™ Lilac</td>
<td>25 Ncm</td>
<td>15 Ncm</td>
</tr>
<tr>
<td>*Astra Tech Implant System™ 4.5ST, 5.0ST</td>
<td>25 Ncm</td>
<td>15 Ncm</td>
</tr>
<tr>
<td>*Straumann® Bone Level, Straumann® Octagon soft tissue level</td>
<td>35 Ncm</td>
<td>15 Ncm</td>
</tr>
<tr>
<td>*Ankylos® Implant System</td>
<td>25 Ncm</td>
<td>15 Ncm</td>
</tr>
<tr>
<td>*Camlog®</td>
<td>20 Ncm</td>
<td>15 Ncm</td>
</tr>
</tbody>
</table>

Note: Always refer to the original implant manufacturer’s instructions for use, with regards to the implant indications and contraindications, as well as tooling and tightening torque.

9. Insert fixed prosthesis and tighten the prosthetic screws by alternating left and right side (K, L). Finally tighten the prosthetic screws according to table 1 using Screwdriver Machine Multi-unit or Unigrip™ Screwdriver, as appropriate, and Manual Torque Wrench prosthetic (M).

10. Close screw access channel.

For additional information on restorative and dental laboratory procedures please consult treatment guidelines available at www.nobelbiocare.com or request latest printed version from a Nobel Biocare representative.

Gold Coping Multi-unit casting specifications: Melting range: 1400–1490° C/2550–2720° F. Coefficient of thermal expansion: 12 μm/m°C. K.

Recommended casting alloys: Conventional gold alloys: High gold content (min 75% Au + Pt metal) alloys, standard ISO 1602 type 4.

Ceramic bonding alloys: High gold content (min 75% Au) alloys, standard ISO/DIS 9693, NiOM type A. Soldering in the range of 800–890° C/1472–1634° F.

Gold Coping Bar: Soldering in the range of 800–890° C/1472–1634° F.

Materials:

- Straight Multi-unit Abutment/Plus for implants with external hex connection and internal tri-channel connection: commerically pure titanium.
- All other Multi-unit Abutment/Plus and Abutment/Prosthetic screws: Titanium alloy 90% Ti, 6% Al, 4% V.
- Holder for Multi-unit Abutment/Plus straight: PP (Polypropylen).
- Holder for Multi-unit Abutment/Plus angled: Titanium alloy 90% Ti, 6% Al, 4% V.
- Gold coping: Gold alloy 60% Au, 19% Pt, 20% Pt, 1% Ir.

Cleaning and sterilization instructions:

Multi-unit Abutment/Plus is delivered sterile for single use only prior to the labelled expiration date.

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

Caution: Multi-unit Abutment/Plus is a single use product and must not be reprocessed. Reprocessing could cause loss of mechanical, chemical and/or biological characteristics. Re-use could cause cross contamination.

Gold Coping Multi-unit Abutment and Multi-unit Abutment Non-Engaging 30° are delivered non-sterile for single use.

Final framework with the Gold Coping Multi-unit Abutment and the jig with the Multi-unit Abutment Non-Engaging 30° should be cleaned and disinfected, as applicable per manufacturer’s instructions, before intraoral use.

Warning: Use of non-sterile device may lead to infection of tissues or infectious diseases.

Caution: Gold Coping Multi-unit Abutment and Multi-unit Abutment Non-Engaging 30° 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.
**Magnetic Resonance (MR) safety information:**

**Note:** Only the Conical Connection Wide Platform abutments have been assessed as MR Conditional. The other platforms and sizes have not been evaluated for safety and compatibility in the MR environment and have not been tested for heating or migration in the MR environment.

Non-clinical testing has demonstrated that the product is MR conditional. A patient with this device can be safely scanned in a MR system meeting the following conditions:

- Static magnetic field of 1.5-Tesla and 3.0-Tesla only.
- Maximum spatial gradient magnetic field of 4000 Gauss/cm (40 T/m).
- Maximum MR system reported, whole body averaged specific absorption rate (SAR) of 4 W/kg (First Level Controlled Mode).

Under the scan conditions defined above, the product is expected to produce a maximum temperature rise of 4.1° C after 15 minutes of continuous scanning.

In non-clinical testing, the image artifact caused by the device extends approximately 30mm from the device. Removable restorations should be taken out prior to scanning, as is done for watches, jewelry etc. when imaged with a gradient echo pulse sequence and a 3.0 Tesla MRI system.

Should there be no MR symbol on the product label, 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 Magnetic Resonance Imaging, please consult the “Cleaning & Sterilization Guidelines including MRI Information of Nobel Biocare Products” 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 16 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.

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**Camlog®** is a trademark of Camlog Biotechnologies Group.

**Astra Tech Implant System™** is a trademark of Dentsply Group.

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