

# REVOIS® Dental Implantat System

## Instructions for Use (IFU) Non-sterile instruments and tools



### 1. Product description

These instructions for use apply to the under (11.) listed non-sterilely provided surgical instruments and tooling for the REVOIS® Dental Implantat System. Product specifics such as size are shown on the packaging labels. The full list of products of the REVOIS® Dental Implantat System, instructions for cleaning, disinfection and sterilization for those instruments that may be used more than one time, service materials and quality certificates can be found at [www.revois-dental.com](http://www.revois-dental.com).

Products are made of surgical steel, titanium grades 2 or 5, gold alloy (Ceramicor®, Elitor®), or Plastic (Rilsan®).

### 2. Indication

The REVOIS® Dental Implantat System is indicated for oral implantation into tooth spaces or the edentulous jawbones, serving as a fixation for prostheses (single tooth replacement, fixed or mobile pontics, overdentures or full dentures).

By this, REVOIS® supports the vital denture function and serves as well as base for a functional and highly aesthetic rehabilitation.

### 3. Field of use

The surgical instruments and prosthetic tools of the REVOIS® Dental Implantat System are used to prepare for insertion, insert, maintain & connect, or remove a REVOIS® implant. For specific indications for REVOIS® implants, please refer to the respective instructions for use that can be found on our website. If you would like to receive a printed copy, please contact our customer service.

Drills and bone taps are used to prepare the cavity for the implant. Use of conical drill and bone taps are mandatory for bone density D I, and optional for D II – IV (after Lekholm & Zarb).

Drill stops may be used with the precision pilot drills, but are not needed with conical drills or bone taps which are rounded at the tip.

### 4. Contraindications

Any disease questioning elective oral surgery. Further on: childhood (incomplete bone development); diseases and circumstances which may negatively impact the healing and performance of an implant such as insufficient oral hygiene, drug- or nicotine abuse; alcoholism; dysregulated metabolic disorders such as diabetes mellitus affecting bone metabolism or osteoporosis or therapy with bisphosphonates; allergy or hypersensitivity to Titanium; any kind of immune deprivation or deficiency, e.g. during therapy with cortisone or cytostatics; pregnancy or lactation; generalizing diseases of the connective tissue or the bones; rheumatological diseases; increased affinity to develop hemorrhage; mental disorders; diseases of heart, liver, kidney or the blood. Do not use instruments / tools from other implant systems which may not fully fit our products.

### 5. Side effects and interactions; Complications

As for any surgery, the patient should avoid physical stress after undergoing surgery / insertion of an implant. The Implantologist has to explain to the patient the possible risks, side effects, interactions, precautions and complications associated with the planned oral surgery and implantation.

This includes e.g. transient and –rarely- persistent paresthesia in the areas of lower lip and chin following mandibular surgery, and numbness of the perinasal tissue following maxillary surgery; further on pain, swelling, affection of speech and gingival inflammation.

In addition, bone height may reduce associated with loosening of the implant, local or secondary systemic infections may develop incl. the development of abscesses or fistula; Impairment of surrounding teeth, roots or nerves; fractures of the prostheses, of the implant or of the bone; aesthetic impairment; gingival hyperplasia.

Instruct your patient for a best possible oral hygiene.

### 6. Warnings and Precautions

#### General

All surgical and prosthetic components are relevant for osseointegration and implant survival and therefore must only be used by a trained specialist. If you would like to receive information about training opportunities, please contact us.

Each individual patient requires full anamnesis and examination with regards to his radiologic, mental and physical status, including teeth, soft- and hard tissue defects which could impair the result of an implantation. A close collaboration between Implantologist / Surgeon, Dentist and Dental technician is essential for an optimal functional and aesthetic result.

#### Other precautions

To grant the sterility of the implant make sure the blister is not damaged and the expiry date has not passed.

The trauma to the surrounding tissue and the implant site should be minimized as possible. In particular, overheating, multiple surgical trauma such as unnecessary drilling, and any risk for infections need to be strictly avoided.

Otherwise, the risk for a delayed or incomplete osseointegration and secondary periimplantitis will increase. For the different drills, please follow the recommended U/min (following table):

	U/min	REVOIS® PRO	REVOIS® compact
Gingival punch	25 – 30	X	X
3-sided drill or rose burr	800	X	X
Precision Pilot drills (optional: drill stop)	up to 1000	X	X
Conical drills	300 – 600	X	-
countersink	300-600	-	X
Bone taps	25-30	X	-
Implant insertion	-	Manually & torque wrench	Manually & torque wrench

Drilling and bone tapping require the use of appropriate, sharp and sterile instruments in combination with sufficient cooling (e.g. isotonic sodium chloride solution). Do not use instruments from other implant systems. Implants should be inserted always aiming for primary stability. All tools and instruments need to be inspected for proper function prior to their use and have to be used in accordance with the instructions for use of the manufacturer to avoid damage to implants, tissue, or other structures.

Due to the small size of the different components take special care that they must not be swallowed or aspirated by the patient. As technically possible, REVOIS® components have a whole to connect with floss that can be held by hand.

After implant insertion the Implantologist needs to decide the time point of first load based on bone quality and primary stability (defined as out-torque resistance of 20 Ncm).

With one or more of the following findings reconsider the risk / benefit ratio of a the intended therapy with a dental implant, and assure the patient also understands such: Osteopenia and insufficient bone quality for any other reason, in relation to intended implant diameter and length; metabolic disease such as diabetes mellitus or disease locally or generally affecting bone metabolism; drug- or nicotine abuse; alcoholism; bruxism; hematologic disease; HIV; autoimmune disease; therapy with a blood-thinning medication; past or planned irradiation therapy of head and neck; incapability of the patient for any reason to maintain the necessary oral hygiene.

The performance of a titanium dental implant may further be impaired by infections, in particular by periodontitis, chronic infections of the maxillary sinuses, untreated nasal airway abnormalities; facial neuralgia; a dehiscence requiring a bone

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transplant; further on formation of abscess or fistula; suppuration; perforation of maxillary sinus or the basis of the mandibular jaw, root of tongue or lower alveolar channel; bleedings, hematoma or edema; infections as well as local or systemic allergic reactions; neuralgic pain, paresthesia. In such cases we recommend to complete the therapy and healing process of such conditions first before the treatment with a dental implant.

### 7. Delivery and Preparation for Use; Reconditioning

The products are supplied cleaned and disinfected, but non-sterile. If they are supposed to be used sterilely, please follow the follow instructions:

Reconditioning and sterilization has to follow a validated process. All applicable laws have to be followed.

Reconditioning should be performed immediately following use of the instruments. By rinsing in a cleansing fluid such as Komet DC1 you can avoid protein fixation and ease the further cleaning process. It is recommended to use an automated washer for further cleaning and disinfection. For details of the reconditioning process validated for REVOIS please refer to our website [www.revois-dental.com](http://www.revois-dental.com).

After disinfection, all instruments and tools must be inspected for proper function and necessary calibration. Products with damage to their coating or cutting edges, deformations or oxidized surfaces and should not be used again, and have to be discharged in accordance with applicable laws.

Choose an appropriate packaging for sterilization. Single instrument packaging: Ensure the sealing is not tense. Set packaging: Put the instruments into a sufficiently large tray. Protect the tray with an appropriate sterilization bag.

Sterilisation: Vacuum steam sterilisation at 134°C in an autoclave with specifications fulfilling DIN EN 13060, following a validated process:

- Fractioned pre-vacuum
- Sterilisation temperatur 134 °C
- Time at temperature: 5 minutes min. (full cycling)
- Drying time: 10 minutes min.

To avoid instrument staining and oxidation, steam must be free of any oxidative substances.

Do not overload autoclave. Follow instructions for use of the manufacturer of the autoclave.

### 8. Storage

Please store the components of the REVOIS® Dental Implantat System in their original packaging, at room temperature, dry and protected from sunlight.

Any other form of storage may impact the sterile barrier and therefore endanger patient and the success of the procedure.

Use of the sterile product after the expiry date may lead to infections.

### 9. Product variants and compatibility

The REVOIS® Dental Implantat System consists of three different, and in-between each other not compatible product variants: **REVOIS® Classic, PRO und compact.**

All three product variants are made of Titanium and therefore have an excellent biocompatibility

For the implantation and prosthetic work only use the components of the respective product line chosen. For a complete overview of the various parts please refer to the User Manual available at [www.revois-dental.com](http://www.revois-dental.com) or contact us to receive your personal copy.

REVOIS® Classic consists of prosthetic accessories and tools for the between 2004 and 2011 commercialized REVOIS® implants.

REVOIS® PRO implants (Titanium grade 4) are delivered as a pre-assembled set consisting of implant, impression post and impression screw (open tray), and cover screw. REVOIS® PRO implants are available in the following sizes:

Length (mm)	Diameter (mm)	Color code
9, 11, 13, 15	3.8	Green
9, 11, 13, 15	4.3	Blue
9, 11, 13, 15	5.0	yellow

REVOIS® PRO implants, accessories, drills and tools are color-coded in accordance with the respective implant diameter. There is a broad range of prosthetic components available allowing the use of REVOIS® PRO for almost every patient.

REVOIS® compact consists of small one-piece implants (Titanium grade 5) delivered as pre-assembled sets consisting of implant, impression post and impression screw (open tray). REVOIS® compact implants are available in the following sizes:

Length (mm)	Diameter (mm)
7	3.0
9	3.0
11	3.0
13	2.2

The 2.2 mm REVOIS® compact implant is designed as a temporary implant where the need for provisional restauration is high but the loading of the final implant site is still not possible. REVOIS® compact is of particular benefit for single tooth replacement with small geometries (e.g. the enosseal replacement of the teeth 12/22/32/31/41/42) as well as for easy fixation of individual or standardized bar systems such as the SFI-BAR®, pontics, or overdentures.

**Designed as an open system, the REVOIS Dental Implantat System allows to connect with SFI-Bar®, Dalbo®, Locator®, and CM LOC®, to meet patient needs and implantologists' preferences in almost every situation.**

For an overview on all components and their use, please refer to the REVOIS® Dental Implantat System User Manual available at [www.revois-dental.com](http://www.revois-dental.com), or contact us for your personal printed copy.

### 10. Use of Product / Procedure

The tools for the preparation of the implant site, implant cavity, insertion and removal of an implant are the gingival punch, the rose burr or 3-sided drill; the measuring device / open-ended wrench, depth gauge, parallelizing pins, precision pilot drills with optional drill stops, conical drills and bone taps, hex tool, handle, angle, thumb wheel, removal tool, and a torque wrench (20-70 Ncm).

All recommended REVOIS® drills and tools are designed or selected for their precise fitting to the respective REVOIS® implants and product variant. Please be aware that the use of other tools or drills, screws or abutments may impact the intactness of the products and prohibit us to take liability for any resulting functional or material defects.

Tools have to be regularly maintained (and as appropriate calibrated), and in accordance with the instructions of the respective manufacturer, to be cleaned, disinfected and sterilized.

Implants and some articles as depicted on their packaging label, are for single use, only.

Prior to surgery, the implantologist has to verify through an appropriate method, e.g. radiography, that the bone quality at

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the intended implant size is sufficient relative to the intended implant size. As control for a correct implant placement, a subsequent radiography is recommended, as well.

If primary stability cannot be achieved, the time required for osseointegration and to the first load of the implant varies between individuals. It is at the discretion of the implantologist to define such time point.

The REVOIS® Dental Implantat System may be used in one-time technique immediately loading the implant after insertion (if primary stability is reached), or staged technique, after osseointegration.

### Insertion:

After preparation of the gingiva, roughen the implant site by using the 3-edged drill or the rose burr to ensure the safe use of the precision drills with or without drill stops. With these precision drills –use the ones appropriate for the respective implant size- prepare the cavity in direction and depth.

The REVOIS® precision drills are labeled with marks coding for the drill depth, and carry adapters to be used with the REVOIS® drill stops.

If using REVOIS® PRO, the cavity may then further be prepared in form and width with the conical drills, and subject to the bone density, finally with the bone taps mimicking the form of the implants.

The REVOIS® PRO conical drills and bone taps have a flat tip avoiding the risk of unnecessary depth drilling and subsequent injury of the underlying tissue. Their use is mandatory for hard bone (D I), and optional for D II to D IV. The implant-size specific REVOIS® PRO bone taps are color coded in line with the color code of the implant size.

**REVOIS® PRO:** Remove the insert blister by using the recessed grips. Touching the impression screw only, remove the premounted set and insert the implant into the prepared implant cavity screwing manually.

Then remove the impression screw and post (you may keep both for re-use for the open-tray impression procedure later on). Using insertion piece and torque wrench screw (turn right) the implant into its final position not exceeding forces of 30 Ncm. Ensure the insertion piece is fully fitting onto the implant. The typical final position of the REVOIS® PRO implant upper edge is 0.5 - 1 mm below the bone crest level. For the open-tray impression (re-)place impression post and screw onto the finally inserted implant. The impression post functions as impression negative and remains in the impression mass. You may hand over the impression mass together with a laboratory analogue to a dental lab to develop the chosen prosthetic solution. CAD/CAM data are available at [www.revois-dental.com](http://www.revois-dental.com).

If for any reason, the implant has to be removed again from the cavity, insert the insertion angle and put on the flipped-over torque wrench to turn out (leftwards) the implant not exceeding forces of 70 Ncm.

When choosing a staged technique (e.g. with bone type D III or IV), do not immediately proceed with the impression procedure, but cover the implant after the insertion into its final position with the cover screw avoiding strain on the implant.

After the healing period (osseointegration with or without augmentation) proceed with the impression procedure as explained before.

For alternative impression options (e.g. snap-on wings, titanium posts, multifunctional abutments) please refer to our manual.

Subsequent prosthetics work: With reaching primary stability or osseointegration, respectively, you can start structuring the oral mucosa by using a gingiva former available in different widths and heights.

After impression taking, one of the various straight or angled standard abutments, or the sprue-able gold-ceramic abutment (Ceramicor®) may be placed on the abutment with the fixation screw using the torque wrench at 30 Ncm. Make sure the screw angle is fully and straight placed onto the screw. Afterwards, the prosthesis can be glued using appropriate media.

In the rare case that a fixation screw has to be removed again, place the screw angle and put on the flipped-over torque wrench to turn out (leftwards) the screw not exceeding forces of 40 Ncm. When using conical abutments make sure to use the appropriate hulls, as well.

When using male/female housing systems such as Dalbo®, Locator®, CM LOC® or CM LOC® Flex, ensure to use the specific tools and accessories, only – all available from AUROSAN.

**REVOIS® compact:** Remove the insert blister by using the recessed grips. Touching the impression screw only, remove the premounted set and insert the implant into the prepared implant cavity screwing manually.

Then remove the impression screw and post (you may keep both for re-use for the open-tray impression procedure later on). Using insertion piece and torque wrench screw (turn right) the implant into its final position not exceeding forces of 30 Ncm. Ensure the insertion piece is fully fitting onto the implant. Typically, insert the REVOIS® compact implant until the upper end of the thread is just covered by the bone crest. For the open-tray impression (re-)place impression post and screw onto the finally inserted implant. The impression post functions as impression negative and remains in the impression mass. You may hand over the impression mass together with a laboratory analogue to a dental lab to develop the chosen prosthetic solution. CAD/CAM data are available at [www.revois-dental.com](http://www.revois-dental.com).

If for any reason, the implant has to be removed again from the cavity, insert the insertion angle and put on the flipped-over torque wrench to turn out (leftwards) the implant not exceeding forces of 70 Ncm.

When choosing a staged technique (e.g. with bone type D III or IV), do not immediately proceed with the impression procedure, but cover the implant after the insertion into its final position with the plastic cap avoiding strain on the implant.

After the healing period (osseointegration with or without augmentation) proceed with the impression procedure as explained before.

For alternative closed tray impression options (e.g. titanium or gold jackets) please refer to our manual.

*The use of the surgical instruments, tools & accessories is described in detail in the REVOIS® User Manual that can be downloaded at [www.revois-dental.com](http://www.revois-dental.com). Please contact us if you would like to receive a personal print copy.*

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11. These Instructions for use apply to the following REVOIS articles. The complete sets of the instructions can be found at [www.revois-dental.com](http://www.revois-dental.com). Please contact us if you wish to receive a printed copy.

REVOIS® Dental Implantat System Non-sterile surgical instruments and prosthetic tools		
Ref-No.	Product-variant	Article description
MD6000411	Classic	Snap-on Tool post for closed-tray impression)
MD6000412	Classic	Screw for Snap-on Tool, long
MD6000422	Classic	Laboratory analogue
MD6000427	Classic	Multifunctional precision abutment length 5.5 mm
MD6000429	Classic	Individual aesthetics abutment (sprue-able; Ceramicor®)
MD6000431	Classic	Standard abutment, 0°, long
MD6000432	Classic	Standard abutment, 18°
MD6000435	Classic	Abutment fixation screw
MD6243300	compact	ISO-adapter
MD6248000	compact	Bending tool, temporary implant
MD6001025	PRO	Screw driver, hex, short , Ø 1.25mm, 23mm
MD6001026	PRO	Screw driver, hex, long, Ø 1.25mm, 30mm
MD6001027	PRO	Implant insertion driver, short, Ø2.5mm, 23mm
MD6001028	PRO	Implant insertion driver, long, Ø2.5mm, 30mm
MD6001029	PRO	Removal tool Ø 2.3mm
MD6001030	PRO	Screw driver Dalbo®
MD6001040	PRO	Thumb wheel
MD6001050	PRO	Handle
MD6001077	PRO	Depth-gauge
MD6001520	PRO	Laboratory analogue
MD6001525	PRO	Training implant
MD6001774	PRO	Laboratory analogue, Dalbo®
MD6001197		Drill stop REVOIS® 7.0 mm
MD6001198		Drill stop REVOIS® 8.0 mm
MD6001201		Drill stop REVOIS® 9.0 mm
MD6001202		Drill stop REVOIS® 10.0 mm
MD6001203		Drill stop REVOIS® 11.0 mm
MD6001204		Drill stop REVOIS® 12.0 mm
MD6001205		Drill stop REVOIS® 13.0 mm
MD6001206		Drill stop REVOIS® 14.0 mm
MD6001207		Drill stop REVOIS® 15.0 mm
MD6001208		Drill stop REVOIS® 16.0 mm
MD6001250		Mandrin
MD6001700	PRO	Abutment fixation screw
MD6001705	PRO	Standard abutment, 0°, Ø 4.4 H5

MD6001707	PRO	Standard abutment, 0°, Ø 4.4 H7
MD6001718	PRO	Standard abutment, 18°, incl. Fixation screw
MD6001725	PRO	Standard abutment, 25°, incl. fixation screw
MD6001730	PRO	Indiv. aesthetics abutment, 0°, incl. fixation screw
MD6001735	PRO	Standard abutment, 0°, Ø 5.2 H5
MD6001740	PRO	Standard abutment, 0°, Ø 5.2 H7
MD6001745	PRO	c-base, CEREC compatible
MD6001741	PRO	Scan abutment, titanium, hex lock
MD6001742	PRO	Scan abutment, titanium, no lock
MD6001747	PRO	impression Set, closed tray
MD6001748	PRO	Snap-on wings, titanium, for closed tray,
MD6001749	PRO	Impression post, multi
MD6001750	PRO	Impression post, open tray
MD6001751	PRO	Impression screw, direct
MD6001752	PRO	Conical abutment, H2, Hull, screw
MD6001754	PRO	Conical abutment, H4, Hull, screw
MD6001756	PRO	Hull for conical abutment
MD6001772	PRO	Dalbo® ball attachment, H3
MD6001773	PRO	Dalbo® ball attachment, H5
MD6001775	PRO	SFI-Bar® attachment Ø 4.4 mm, H2
MD6001776	PRO	SFI-Bar® attachment Ø 4.4 mm, H3
MD6001777	PRO	SFI-Bar® attachment Ø 4.4 mm, H4
MD6001778	PRO	SFI-Bar® attachment Ø 4.4 mm, H5
MD6001871	PRO	CM LOC® Abutment GH1 PRO
MD6001872	PRO	CM LOC® Abutment GH2 PRO
MD6001873	PRO	CM LOC® Abutment GH3 PRO
MD6001874	PRO	CM LOC® Abutment GH4 PRO
MD6001875	PRO	CM LOC® Abutment GH5 PRO
MD6001876	PRO	CM LOC® Case Guide 20° PRO
MD6001881	PRO	CM LOC®FLEX Abut. PRO GH1
MD6001882	PRO	CM LOC®FLEX Abut. PRO GH2
MD6001883	PRO	CM LOC®FLEX Abut. PRO GH3
MD6001884	PRO	CM LOC®FLEX Abut. PRO GH4
MD6001885	PRO	CM LOC®FLEX Abut. PRO GH5
MD6001886	PRO	CM LOC®FLEX Case Guide 30° PRO
MD6213118	compact	Scan abutment, titanium, hex lock
MD6213119	compact	Scan abutment, titanium, no lock
MD6221130	compact	Titanium-made hull, no lock
MD6221131	compact	Plastic cap
MD6221132	compact	Titanium-made hull, hex lock
MD6225130	compact	Gold alloy hull, no lock
MD6225131	compact	Gold alloy hull, hex lock
MD6237120	compact	Inbus screw 0.9 mm
MD6001244		Drill extension

**12. Contact details of the manufacturer**



Aurosan GmbH  
 Frankenstrasse 231  
 D-45134 Essen, Germany  
 Fon: +49 (0)201 506 58151  
 Fax: +49 (0)201 506 58152  
 service@aurosan.de  
 www.revois-dental.com



For USA: Federal law permits the sale of this medical device only through or after prescription by a licensed dentist or medical doctor.

Pictogram	Meaning
	Manufacturer
	The REVOIS® Dental Implantat System is in conformity with the medical device guideline 93/42 EWG
	Read instructions for use, available at <a href="http://www.revois-dental.com/gba">www.revois-dental.com/gba</a>
<b>REF</b>	Catalog number
	Lot number
	Use before
	Sterilised by radiation
	Single use only
<b>QTY</b>	Quantity
	Keep dry
	Protect from sunlight
<b>Rx only</b>	Prescription only