



Guarantee Form

PLEASE FILL-OUT IN ENGLISH

Customer Information

Clinician's Name _____ Customer Account _____

Address _____ Telephone _____

_____ Country _____

Reported by _____

Product Information (Please list all involved MEISINGER IMPLANTS Products)

Article Number	LOT Number	Placement Day (DD/MM/YYYY)	Removal Day (DD/MM/YYYY)	Position

General Patient Information (complete this section only, if returning implants)

Patient ID No. * _____ Age _____ F M

* For privacy reasons do NOT enter the name of the patient, when the patient ID is not anonymised in the form (and additional attachments) and contains personal information, the patient has to give a written consent.

Medical Record

- Diabetes mellitus
- Radiation Tx-head/neck area
- Illness requiring steroids
- Chemotherapy at time of implant placement
- Psychological disorder
- Xerostomia
- Lymphatic disorder
- Drug or alcohol abuse
- Uncontrolled endocrine illness
- Compromised immuno resistance
- Blood coagulation disorder
- Immunologic disease

Allergies _____

Other local or systemic diseases which may be significant? _____

Smoker Yes _____ cigarettes/day No No significant findings

Implant Failure - Surgical Information (complete this section only, if returning implants)

Manuel Placement with Handpiece Adapter

If implant was placed and removed the same day, has another implant successfully been placed in the site during surgery?

Yes No

If you experienced difficulty with inserting device/pre-mounted transfer part this occurred upon?

Implant insertion Removal of device from implant Removal of implant from vial Other _____

Have there been present any diseases when placing the implants?

Periodontal disease Diseased mucous membrane Local Infection/Subacute Chronic Osteitis Complication in site preparation

Quality of bone Type D1 Type D2 Type D3 Type D4

Site tapped? Yes No Not applicable

Profile drill used? Yes No Not applicable

Holding key used? Yes No Not applicable

Primary stability achieved? Yes No

Osseointegration of implant achieved? Yes No

Implant surface completely covered with bone? Yes No

Augmentation at the time of surgery?

No Sinus Ridge Material used _____

GTR membrane?

No Yes Resorbable Non-resorbable

Material used _____



Guarantee Form

General Information (complete this section only, if returning implant)

Hygiene around implant Excellent Good Fair Poor

Other circumstances?

- | | | |
|---|--|---|
| <input type="checkbox"/> Trauma/Accident | <input type="checkbox"/> Implant Fracture | <input type="checkbox"/> Inadequate Bone Quality/Quantity |
| <input type="checkbox"/> Biomechanical Overload | <input type="checkbox"/> Overheating of bone | <input type="checkbox"/> Previous Bone Augmentation |
| <input type="checkbox"/> Immediate Extraction Site | <input type="checkbox"/> Peri-Implantitis | <input type="checkbox"/> Nerve Encroachment |
| <input type="checkbox"/> Adjacent to Endodontic Tooth | <input type="checkbox"/> Infection | <input type="checkbox"/> Sinus Perforation |
| <input type="checkbox"/> Tongue (Pressure) | <input type="checkbox"/> Bruxism | <input type="checkbox"/> Bone Resorption |

When the implant failed there had been (check all that apply)

- | | | | |
|--|---------------------------------------|---------------------------------------|---------------------------------------|
| <input type="checkbox"/> Pain | <input type="checkbox"/> Bleeding | <input type="checkbox"/> Swelling | <input type="checkbox"/> Numbness |
| <input type="checkbox"/> Mobility | <input type="checkbox"/> Fistula | <input type="checkbox"/> Asymptomatic | <input type="checkbox"/> Inflammation |
| <input type="checkbox"/> Increased Sensitivity | <input type="checkbox"/> Hypertension | <input type="checkbox"/> Abscess | <input type="checkbox"/> Others _____ |
- The Prosthesis has been fitted? No Yes, please complete Section Prosthesis

If the implant isn't removed: Are there any indications of the following? (please tick the appropriate box)

Expansion (mm) _____ Bone loss _____ Dehiscence _____ Peri-Implantitis _____ Fenestration _____ Others _____

Please comment on why you think the implant failed / was removed

Prosthesis Information (complete this section only, if returning abutments and restorations)

- Model Therapy in use
- Type of restoration? Crown Bridge RPD (upper) RPD (lower)
 Full (upper) Full (lower) Telescope Others _____
- Abutment inserted (date) Abutment removed (date)
- Torque control device used? Yes No Unknown Torque applied Ncm
- Temporary restoration (date of insertion) Final restoration (date of insertion)
- Did the patient follow recall instructions? Yes No

Comment

Instruments (complete this section only, if returning instruments)

- Approximate number of uses (Cutting Instruments only) Initial use 2 - 5 6 - 10 11 - 15 over 15
- Type of cleaning method Manual Ultrasonic Thermodesinfection Others _____
- Type of sterilization method Autoclave Dry heat Chemiclav

Short description of incident

Autoclave all return products and label them as sterile.

Please complete all necessary details of the products to be complained about in this warranty form, observing the warranty conditions of Meisinger Implants GmbH and return this form, including the sterilized products and X-ray images, to Meisinger Implants GmbH.
Use padded shipping bag for return - loss of individual items during shipping will void the warranty.

Signature _____ Date _____

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