


# Guarantee form

 As specified in paragraph 8 of the Thommen Medical Guarantee, all applicable data protection regulations must be complied with and all patient data must be anonymized. A patient ID number, which does not allow any conclusions to be drawn about patient data, must be used for each patient.

## 1. CUSTOMER INFORMATION

Attending physician's name and address (use capital letters or stamp)

_____	Telephone	_____
_____	Country	_____
_____	Contact at the practice	_____
_____		

## 2. PRODUCT INFORMATION (please list all Thommen Medical products involved)

Art. no.	Lot no.	Placement Date (D/M/Y)	Removal Date (D/M/Y)	Region
_____	_____	____ / ____ / _____	____ / ____ / _____	_____
_____	_____	____ / ____ / _____	____ / ____ / _____	_____
_____	_____	____ / ____ / _____	____ / ____ / _____	_____
_____	_____	____ / ____ / _____	____ / ____ / _____	_____

**Date of the event:** \_\_\_\_\_ / \_\_\_\_\_ / \_\_\_\_\_

## 3. GENERAL PATIENT INFORMATION (complete this section only if returning implants)

Patient ID no. \_\_\_\_\_ Age: \_\_\_\_\_  Female  Male

### Medical Record

<input type="checkbox"/> Radiation Tx-head/neck area	<input type="checkbox"/> Blood coagulation disorder	<input type="checkbox"/> Psychological disorder
<input type="checkbox"/> Bisphosphonate treatment	<input type="checkbox"/> Compromised immune resistance	<input type="checkbox"/> Drug or alcohol abuse
<input type="checkbox"/> Illness requiring steroids	<input type="checkbox"/> Diabetes mellitus	<input type="checkbox"/> Xerostomia
<input type="checkbox"/> Chemotherapy around time of implant placement	<input type="checkbox"/> Uncontrolled endocrine illness	Smoker: <input type="checkbox"/> Yes <input type="checkbox"/> No

Other local or systemic diseases which may be significant: \_\_\_\_\_

Allergies: \_\_\_\_\_  No significant findings

## 4. INFORMATION (complete this section only if returning implants)

Manual placement  Mechanical insertion

**Comments** (use capital letters): \_\_\_\_\_

If implant was placed and removed the same day, was another implant successfully placed in the site during surgery?  Yes  No

**At the time of surgery, were any of the following present:**

<input type="checkbox"/> Periodontal disease	Bone quality	<input type="checkbox"/> Type I	<input type="checkbox"/> Type II	<input type="checkbox"/> Type III	<input type="checkbox"/> Type IV
<input type="checkbox"/> Diseased mucous membrane	Use thread tap?	<input type="checkbox"/> Yes	<input type="checkbox"/> No		
<input type="checkbox"/> Local infection/ subacute chronic osteitis	Was primary stability achieved?	<input type="checkbox"/> Yes	<input type="checkbox"/> No		
<input type="checkbox"/> Complication in site preparation	Did implant achieve osseointegration?	<input type="checkbox"/> Yes	<input type="checkbox"/> No		
Whichever: _____	Was the implant surface completely covered with bone?	<input type="checkbox"/> Yes	<input type="checkbox"/> No		

**Was augmentation performed at the time of surgery?**

No  Sinus  Ridge

Material used: \_\_\_\_\_

**Was GTR membrane used?**

No  Yes  Resorbable  Non-resorbable

Material used: \_\_\_\_\_

**5. EVENT INFORMATION** (complete this section only if returning implants)

Hygiene around implant  Excellent  Good  Fair  Poor

**Were any of the following involved in the event?**

- |   |   |   |
|---|---|---|
| <input type="checkbox"/> Trauma/Accident        | <input type="checkbox"/> Undersized implant bed           | <input type="checkbox"/> Preceding/simultaneous bone augmentation |
| <input type="checkbox"/> Biomechanical overload | <input type="checkbox"/> Overheating of bone              | <input type="checkbox"/> Bone resorption                          |
| <input type="checkbox"/> Bruxism                | <input type="checkbox"/> Nerve encroachment               | <input type="checkbox"/> Peri-Implantitis                         |
| <input type="checkbox"/> Implant fracture       | <input type="checkbox"/> Sinus perforation                | <input type="checkbox"/> Infection                                |
| <input type="checkbox"/> Immediate implantation | <input type="checkbox"/> Inadequate bone quality/quantity |   |

Other (please write in capital letters): \_\_\_\_\_

**At the time of implant failure, there was** (check all that apply):

- |  |                                   |                                       |                                   |
|--|-----------------------------------|---------------------------------------|-----------------------------------|
| <input type="checkbox"/> Pain                  | <input type="checkbox"/> Swelling | <input type="checkbox"/> Bleeding     | <input type="checkbox"/> Mobility |
| <input type="checkbox"/> Numbness              | <input type="checkbox"/> Fistula  | <input type="checkbox"/> Inflammation | Other: _____                      |
| <input type="checkbox"/> Increased sensitivity | <input type="checkbox"/> Abscess  | <input type="checkbox"/> Asymptomatic |                                   |

**Was the prosthesis fitted?**

- Yes  No

**If yes, please complete section 6.**

Please comment on why you think the implant failed/was removed (please write in capital letters):

\_\_\_\_\_

\_\_\_\_\_

**6. PROSTHESIS INFORMATION** (complete this section only if returning abutments and restorations)

- Model  Insertion  In use
- Type of restoration?  Crown  Bridge RPD:  Upper  Lower  
Full:  Upper  Lower
- Date abutment was installed? \_\_\_\_\_ / \_\_\_\_\_ / \_\_\_\_\_ Date of abutment removal: \_\_\_\_\_ / \_\_\_\_\_ / \_\_\_\_\_  
(D/M/Y) (D/M/Y)
- Was the MONO torque ratchet used?  Yes  No  Unknown Torque applied: \_\_\_\_\_ Ncm
- Date of temporary restoration installation: \_\_\_\_\_ / \_\_\_\_\_ / \_\_\_\_\_ Date of final restoration installation: \_\_\_\_\_ / \_\_\_\_\_ / \_\_\_\_\_  
(D/M/Y) (D/M/Y)
- Was a check-up performed?  Yes  No

**Description of event** (please write in capital letters):

\_\_\_\_\_

\_\_\_\_\_

**7. INSTRUMENTS** (complete this section only if returning instruments)

- Which drills were used:  VECTOdrill steel  VECTOdrill ceramic
- Other Whichever: \_\_\_\_\_
- Approximate number of uses (cutting instruments only):  Initial use  2-5 x  6-10 x  10-20 x  More than 20 x
- Type of cleaning method used:  Manual  Ultrasonic  Thermodesinfection Other: \_\_\_\_\_
- Type of sterilization method used:  Autoclave  Dry heat  Chemiclave

**Short description of incident** (please write in capital letters):

\_\_\_\_\_

\_\_\_\_\_

Please return questionnaire, autoclaved product and include X-rays (as appropriate) to your distribution partner.

**Use a padded pouch to return items – failure to do so could result in items lost during shipment and void guarantee program.**

**Autoclave** all products (do not clean) and label them as **sterile**.

Based on the Thommen Medical Guarantee Terms and Conditions, please consider replacing the above listed products.

Doctor's Signature: \_\_\_\_\_ Date: \_\_\_\_\_