

# Straumann Guarantee® Questionnaire

straumanngroup.ca/eshop

File Number (if known)

## CUSTOMER INFORMATION

### Customer Details

Facility Name \_\_\_\_\_  
Clinician Name \_\_\_\_\_  
Contact Phone \_\_\_\_\_  
Contact E-Mail \_\_\_\_\_

Sold to Account #: \_\_\_\_\_  
Address 1 \_\_\_\_\_  
Address 2 \_\_\_\_\_  
Address 3 \_\_\_\_\_  
City \_\_\_\_\_  
Prov \_\_\_\_\_ Postal Code \_\_\_\_\_

*Check if same as Sold To*

Ship to Account #: \_\_\_\_\_  
Address 1 \_\_\_\_\_  
Address 2 \_\_\_\_\_  
Address 3 \_\_\_\_\_  
City \_\_\_\_\_  
Prov \_\_\_\_\_ Postal Code \_\_\_\_\_

## PATIENT INFORMATION

### Patient Detail (for privacy DO NOT use patient's name)

Patient ID \_\_\_\_\_  
Patient Age \_\_\_\_\_  
Gender: Female Male  
Other \_\_\_\_\_  
Smoker? No Yes

### History

Psychological disorder      Blood coagulation disorder      Illness requiring steroids  
Lymphatic disorder      Untreated endocrine illness      Coincident chemotherapy  
Drug or alcohol abuse      Diabetes Mellitus      Xerostomia  
Compromised immunity      Radiation Tx (head/neck area)      No significant findings  
Relevant allergies: \_\_\_\_\_ Relevant diseases: \_\_\_\_\_

## PRODUCT INFORMATION \*Please list all involved Straumann products

*Lifetime Plus claims must be accompanied by the restoration and restoration details (include here) and any relevant radiographs.*

Article REF Number	Lot/Serial/Project #	Placement Date	Event/Removal Date	Site FDI
1 _____	_____	_____	_____	_____
2 _____	_____	_____	_____	_____
3 _____	_____	_____	_____	_____

Exchange with same Article#(s)? Yes No; specify Article (REF) No(s): 1 2 3  
Was the product used in a patient?: Yes No

## DESCRIPTION OF EVENT

## SURGERY INFORMATION - IMPLANT RELATED (required for implants)

Placement Method: Manually Handpiece adapter

If implant was placed and removed on same day, was another implant successfully placed at site during surgery?

Yes No – Why not? \_\_\_\_\_

If you experienced difficulty inserting an implant with a pre-mounted transfer piece when did this occur (check one)?

Implant removal from vial      Implant insertion into bone      Removal of device from implant

At the time of surgery, were any of the following conditions present (check all that apply)?

Periodontal disease      Local infection      Diseased mucous membrane      Complication in site prep

Bone quality (type):	I	II	III	IV
Was site tapped?	Yes	No		
Bone-level profile drill used?	Yes	No		
Tissue-level profile drill used?	Yes	No		
Was holding key used?	Yes	No		
Was primary stability achieved?	Yes	No		
Was osseointegration achieved?	Yes	No		
Was implant covered with bone?	Yes	No		
Was the implant immediately loaded?	Yes	No		

Was augmentation performed during surgery? Yes No  
Type of augmentation Sinus Ridge  
Material Used? \_\_\_\_\_  
Was a membrane used?  
Yes No Resorbable Non-Resorbable  
Material Used? \_\_\_\_\_

## EVENT INFORMATION (required for Implant and Biomaterial products)

Assessment of hygiene around implant:    Excellent    Good    Fair    Poor

Why do you believe the event occurred:

Were any of the following conditions involved in the event (check all that apply)?

Trauma/Accident	Implant fracture	Inadequate bone quality/quantity
Overheating of bone	Bruxism	Previous bone augmentation
Peri-implantitis	Nerve encroachment	Biomechanical overload
Sinus perforation	Tongue pressure	Adjacent to endodontic tooth
Infection	Immediate extraction site	Bone resorption

At the time of the event or implant failure/removal, was there (check all that apply)?

Pain	Increased Sensitivity	Numbness
Mobility	Abscess	Inflammation
Bleeding	Swelling	Hypersensitivity
Fistula	Asymptomatic	Other: _____

Was the prosthesis fitted?    Yes    No    For multiple unit restorations (i.e., bridges and dentures), how many implants supported the restoration: \_\_\_\_\_

If the implant is not being removed, is there evidence of the following (check all that apply)?

Bone loss; Extent (mm): \_\_\_\_\_    Dehiscence    Fenestration    Peri-implantitis    Other: \_\_\_\_\_

## PROSTHESIS INFORMATION (required for Abutment, Straumann CARES® Digital Solutions restoration, Roxolid and PURE Ceramic fracture)

Type of prosthesis?    Crown    Bridge    Inlay/Onlay    Veneer    Full (upper)    Full (lower)    Other: \_\_\_\_\_

Date abutment was installed \_\_\_\_\_    Date temporary restoration installed \_\_\_\_\_

Date abutment was removed \_\_\_\_\_    Date final restoration installed \_\_\_\_\_

Torque Control Device used    Unknown    No    Yes --- Torque Applied (Ncm): \_\_\_\_\_

Was the recall appointment schedule followed?    Yes    No

## INSTRUMENT INFORMATION (required for Surgical Instruments)

Be sure to *thoroughly clean* instruments and *reassess* prior to returning; most instances of poor instrument performance are due to retained contamination.

Approximate number of uses (cutting tools)?    Initial use    2-5    6-10    10-15    More than 15

Type of cleaning method used?    Manual    Ultrasonic    Thermodisinfection    Other: \_\_\_\_\_

Type of sterilization method used?    Autoclave    Dry heat    Chemiclave    Other: \_\_\_\_\_

Reason for return?    Rust    Other: \_\_\_\_\_

Straumann Use Only
Product Returned?
Product Lost?
Product Sterile?

## SUBMISSION INFORMATION

Products must be returned within 90 days of the date of the event in *protective packaging* (padded mailer) using a method that allows for shipment *tracking*:

- The complete product must be returned for investigation (i.e., package for a labeling issue)
- For products contaminated with bodily fluids, metal or ceramic items must be *autoclaved and marked sterile* by either an autoclave indicator or hand written; plastic items must be *cold sterilized*.
- Only one replacement implant per day per tooth site qualifies for replacement under the Straumann Guarantee.
- Relevant radiographs (these will not be returned unless specifically requested, please send copies).

Send shipment to: Straumann Canada Ltd.  
ATTN: Regulatory Affairs  
1109 Clay Avenue, Unit 8  
Burlington, ON L7L 0A1

Questions?  
Phone: 800/363 4024  
Fax: 978/747 0023  
E-Mail: reg\_complaint@straumann.com

Upon receipt, Straumann will review your feedback, assess the returned product and determine whether the product meets the conditions for replacement under the Straumann Guarantee. When all necessary information and product is received, replacement product can be provided in a timely manner.

Straumann Internal Use Only	
CSN - back office activity	
Regulatory Product Complaint	
PSO    Information incomplete	
ASR    Standard / No Report	
Straumann RA Signature	Date

## SIGNATURE (required - may be electronic)

By signing below I am acknowledging that I understand the terms and conditions of the Straumann Guarantee® and that the information being provided is truthful and accurate.

Clinician Name (print): \_\_\_\_\_ Signature: \_\_\_\_\_ Date: \_\_\_\_\_



Straumann Canada Ltd.  
**ATTN: Regulatory Affairs**  
1109 Clay Avenue, Unit 8  
Burlington, ON L7L 0A1



Package Address - Clip and Tape to Package

### Did you remember to...

- Verify the terms and conditions
- Complete the Straumann Guarantee Questionnaire as completely as possible
- Include your Straumann Account Number(s) on the Questionnaire
- Sterilize the product and mark it as STERILE
- Attach the Product to the Questionnaire or write the Patient ID on the container
- Have the Clinician sign and date Page 2
- Send Product and Questionnaire in protective packaging via a traceable method
- Keep Page 3 for your records