

Guarantee Form

PLEASE FILL-OUT IN ENGLISH

Customer Information							
Clinician's Name	Customer Account						
Adress	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						
	nonymised in the form (and additional attachments) and contains personal information, the patient has to give a written consent.						
Medical Record							
	vchological disorder Uncontrolled endocrine illness ostomia Compromised immuno resistance						
	Inphatic disorder Blood coagulation disorder						
Chemotherapy at time of implant placement Drug or alcohol abuse Immunologic disease Allergies Immunologic disease							
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 Adapt	er						
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							
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



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General Information (complete the	is section only, if ret	turning implant)					
Hygiene around implant	Excellent Go	ood Fair	Poor				
Other circumstances?							
Trauma/Accident Inadequate Bone Quality/Quantity							
Biomechanical Overload Overheating of bone				Previous Bone Augmentation			
Immediate Extraction Site				Nerve Encroachment			
Adjacent to Endotontic Tooth	Adjacent to Endotontic Tooth Infection			Sinus Perforation			
Tongue (Pressure)				Bone Resorption			
When the implant failed there had been (check all that apply)							
Pain	Bleeding		Swelling	Γ	Numbness		
Mobility	Fistula		Asymptomatic		Inflammation		
Increased Sensitivity	Hypertension		Abscess		Others		
The Prosthesis has been fitted?	No		Yes, please cor	nplete Section Prosthesis	 3		
If the implant isn't removed: Are there any indications of the following? (please tick the appropriate box)							
Expansion (mm) Bone loss	Dehiscence	Peri-Im	nplantitis	Fenestration	Others		
Please comment on why you think the impla	nnt failed / was removed						
Prosthesis Information (complete	this section only if	returning abutments	and restorations)				
	erapy	in use		. —	<i>n</i> .		
Type of restoration?	own	Bridge	RPD (uj	oper) F	RPD (lower)		
Full	(upper)	Full (lower)	Telesco	pe C	Others		
Abutment inserted (date)			Abut	ment removed (date)			
Torque control device used?	Yes	No Un	nknown	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	n only, if returning i	nstruments)					
Approximate number of uses	Initial use	2 - 5	6 - 10	11 - 15	over 15		
(Cutting Instruments only) Type of cleaning method	Manual	Ultrasonic	Thermodesinfe	ction Others			
Type of sterilization method	Autoclave	Dry heat	Chemiclav				
	, alcolave	Dry nour					
Short description of incident							
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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

* 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.

Date
