

For Zimvie Use Only Not to be Completed by the Reporter	CMP#:
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PRODUCT EXPERIENCE REPORT

The inclusion of as many details as possible greatly aids the investigation process, continuous improvement and is necessary **to comply with Medical Device Manufacturer Regulatory Requirements**. Missing information will delay processing. Required fields are identified with an asterisk(*).

Document if a Complaint # has been previously assigned	CMP #: _____
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A. EVENT INFORMATION	Placement Date*: _____ (dd/mmm/yyyy)	Event Date*: _____ (dd/mmm/yyyy)	Removal Date*: _____ (dd/mmm/yyyy)
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Discovered*: During receiving / unpacking During clinical procedure During Laboratory Procedure
 Other: _____

Description of the Event (Check all that apply)*

<input type="checkbox"/> Allergic Reaction	<input type="checkbox"/> Infection	<input type="checkbox"/> Nerve Injury	<input type="checkbox"/> Peri-implantitis
<input type="checkbox"/> Bone Loss	<input type="checkbox"/> Lack of Primary Stability	<input type="checkbox"/> Non-Integration (NI)	<input type="checkbox"/> Sinus Perforation
<input type="checkbox"/> Fracture	<input type="checkbox"/> Loss of Integration (LI)	<input type="checkbox"/> Other, please detail: _____	

Provide a detailed description of the reported problem (including procedure being performed, related products and settings used)*:

At the time of the event or implant failure/removal, was there ...? (Check all that apply)*:	<input type="checkbox"/> No Patient Impact <input type="checkbox"/> Abscess <input type="checkbox"/> Ingestion <input type="checkbox"/> Pain <input type="checkbox"/> Inflammation <input type="checkbox"/> Aspiration <input type="checkbox"/> Paresthesia <input type="checkbox"/> Edema <input type="checkbox"/> Other: _____
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Was surgical and/or medical intervention necessary to preclude permanent impairment?*	<input type="checkbox"/> Yes <input type="checkbox"/> No If Yes, please describe: _____
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Was there a delay during the procedure?*	<input type="checkbox"/> Yes <input type="checkbox"/> No If Yes, please describe: _____
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Will the patient have to return for an additional dental appointment to complete the procedure?*	<input type="checkbox"/> Yes <input type="checkbox"/> No If Yes, please describe: _____
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Was the procedure completed using another implant or another device?*	<input type="checkbox"/> Yes <input type="checkbox"/> No If Yes, please describe: _____
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Other Relevant Patient History (Check all that apply)*:	<input type="checkbox"/> Bruxism <input type="checkbox"/> Diabetes <input type="checkbox"/> Smoker / Tobacco use <input type="checkbox"/> Clenching <input type="checkbox"/> Osteoporosis <input type="checkbox"/> Inadequate Oral Hygiene <input type="checkbox"/> Other: _____
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Tooth Number*	_____ <input type="checkbox"/> Universal <input type="checkbox"/> FDI <input type="checkbox"/> Palmer	Bone Density Type*	<input type="checkbox"/> I <input type="checkbox"/> II <input type="checkbox"/> III <input type="checkbox"/> IV <input type="checkbox"/> Unknown
Tooth Number*	_____ <input type="checkbox"/> Universal <input type="checkbox"/> FDI <input type="checkbox"/> Palmer		

Additional Information:	<input type="checkbox"/> Grafted prior to implant placement <input type="checkbox"/> Grafted together with implant placement	<input type="checkbox"/> Site Grafted If Yes, Describe Material _____ Graft placement date: _____	<input type="checkbox"/> Allograft <input type="checkbox"/> Alloplast <input type="checkbox"/> Autogenous <input type="checkbox"/> Hybrid <input type="checkbox"/> Xenograft
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Note: This information is being gathered to assist in complying with regulatory requirements in the USA and other countries as applicable. Completion of this form does not constitute an admission that medical personnel, distributor, manufacturer, or product caused or contributed to the event.

B. PRODUCT INFORMATION: One form should be used per event and/or patient. If more than, one device is associated with a single event being reported, multiple Item numbers may be included below. Additional rows may be added, or additional information included as necessary.

NOTE: 1) Please make sure product listed below has been properly decontaminated. 2) For non-Patient Specific Products, return only the complaint product. 3) For ZFX products please indicate Order number if possible:

<i>Item Number*</i> <small>(If available, affix patient record label)</small>	<i>Lot / Serial Number*</i>	<i>Qty.*</i>	<i>Replacement Requested</i>	<i>Is Product Being Returned?*</i>	<i>If No, Why?*</i> <small>(i.e. retained by the hospital, scrapped, etc.)</small>
				<input type="checkbox"/> Yes <input type="checkbox"/> No	<input type="checkbox"/> Discarded <input type="checkbox"/> Used <input type="checkbox"/> Remains Implanted <input type="checkbox"/> Other:
				<input type="checkbox"/> Yes <input type="checkbox"/> No	<input type="checkbox"/> Discarded <input type="checkbox"/> Used <input type="checkbox"/> Remains Implanted <input type="checkbox"/> Other:
<i>Is destructive analysis permitted?*</i>				<input type="checkbox"/> Yes <input type="checkbox"/> No	

C. REPORTER INFORMATION	
<i>Reporter Name*</i>	
<i>Date of Report*</i>	
<i>Is the person submitting this report</i>	<input type="checkbox"/> Clinician <input type="checkbox"/> Lab <input type="checkbox"/> Distributor <input type="checkbox"/> Sales Representative
<i>Account Name</i>	
<i>Account #*</i>	
<i>Address</i>	
<i>City, State, Zip, Country</i>	
<i>Contact Name*</i>	
<i>Phone #*</i>	
<i>E-mail*</i>	

D. PATIENT INFORMATION	
<i>Patient Identifier*</i>	
<i>Gender*</i>	<input type="checkbox"/> Male <input type="checkbox"/> Female
<i>Age at the time of the event*</i>	
<i>Weight</i>	

Instructions for returning complaint product:

- (US, Canada, APAC and non-European Reporters)** Complete the Product Experience Report (PER) editable PDF, save and email to the appropriate ZimVie complaint handling contact email (see page 3). The complaint handling contact will reply with the complaint number (CMP #(s)) and the product return instruction.
- (All other reporters) Complete the Product Experience Report (PER) editable PDF, save and print. The printed form will be shipped along with the sterile product to the appropriate complaint handling site (see page 3).
- If a Serious Adverse Event related to Human Tissue occurs in the UK, the reporter has an obligation to notify Biomet3i UK, Ltd within 24 hours of event's discovery. Complaint Contact details are located on page 3 of this form.
- Contaminated product shall be sterilized and identified as **STERILE**.
- Return product labeled with the CMP # (if known) in an appropriate shipping container along with a copy of this completed PER form to the addresses provided and/or indicated on page 3 of this form.
- .Used and/or contaminated regenerative product shall **not** be returned to the Zimmer Biomet complaint handling contact site.

Note: This information is being gathered to assist in complying with regulatory requirements in the USA and other countries as applicable. Completion of this form does not constitute an admission that medical personnel, distributor, manufacturer, or product caused or contributed to the event.

Complaint Handling Contacts:

International (APAC & Non-European):

<p>US Biomet 3i & Zimmer Dental Attn: Complaints Handling 4555 Riverside Drive Palm Beach Gardens, FL 33410 Phone: 1.800.262.2702 Email: DentalComplaints@zimvie.com</p>	<p>Canada Biomet 3i & Zimmer Dental ZimVie – Zimmer Biomet Dental Canada Inc. 2345 Argentia Road Suite #106 Mississauga, Ontario L5N 8K4 Phone: 416-995-6664 Email: DentalComplaints@zimvie.com</p>	<p>International Biomet 3i & Zimmer Dental Attn: Complaints Handling 4555 Riverside Drive Palm Beach Gardens, FL 33410 Phone: 561.776.6918/ 1.800.262.2702 Email: DentalInternationalComplaints@zimvie.com</p>	<p>China Zimmer Dental Zimmer Dental (Shanghai) Medical Device Co Ltd Room 2001, Metro Plaza 555 Lou Shan Guan Road, Shanghai 200051 China Phone: 086 21 222 05180 Email: DentalInternationalComplaints@zimvie.com</p>
<p>Australia: Phone: +61 2 9855 4444 Mexico: Phone: +52 55 2282 0120</p>	<p>Chile Zimmer Dental Zimmer Dental Chile SPA Luis Thayer Ojeda 0130 Oficina 901/902 Providencia Santiago, Chile Email: DentalInternationalComplaints@zimvie.com</p>	<p>India Biomet 3i & Zimmer Dental ZB dental India Pvt. Ltd. Unit No. 904 & 905, A-Wing, Damji Shamji corporate Square, Off. Ghatkopar Andheri Link Road, Laxmi Nagar, Ghatkopar East, Mumbai, 400075, India. Phone: 18002669920 / + 91 022 6901 3700 Email: info.india@zimvie.com</p>	

Europe

Non- Patient Specific Product

<p>Austria Biomet 3i & Zimmer Dental Zimvie Austria GmbH Wienerbergstrasse 11/12a 1100 Wien, Austria Phone: +43 (0) 8000 700 17 Fax: +43 (0) 8000 700 18 Email: EMEAComplaints@zimvie.com</p>	<p>Belgium and Luxembourg Biomet 3i Biomet 3i Belgium Building MC Square Schaliënhoedreef 20T 2800 Mechelen, Belgium Phone: +32 80050311 Email: EMEAComplaints@zimvie.com</p>	<p>France and Luxembourg Biomet 3i & Zimmer Dental Zimmer Dental S.A.S. 19 rue d'Arcueil 94150 Rungis, France Phone: +33(0) 800 91 67 86 Email: EMEAComplaints@zimvie.com</p>	<p>Germany Biomet 3i & Zimmer Dental Zimmer Dental GmbH Wilhelm-Wagenfeld-Straße 88 80807 München, Germany Phone: +49 (0) 800 184 0271 / +49 (0) 800 101 6420 Fax: +49 (0)800 313 11 11 Email: EMEAComplaints@zimvie.com</p>
<p>Israel Zimmer Dental Zimmer Dental Ltd 13 Haamal St.Afeq Industrial Park Building A, 3rd Floor, Rosh Haayin 4809280, Israel Email: ZBI-CS@zimmerbiomet.com</p>	<p>Italy Zimmer Dental Zimmer Dental Italy S.R.L Viale Italia 205/D 31015 Conegliano (TV), Italy Phone: +39 0438 37681 Email: zimmerdental.italy@zimvie.com</p>	<p>Netherlands Biomet 3i Biomet 3i Netherlands B.V Marten Meesweg 25-G 3068 AV Rotterdam, Netherlands Phone: +31 078 62 92 800 Email: EMEAComplaints@zimvie.com</p>	<p>Spain and Portugal Biomet 3i & Zimmer Dental Biomet 3i Dental Ibérica, S.L.U WTC Almeda Park, Ed.4, Planta 2 C/Tirso de Molina, 40 08940 Cornellà de Llobregat (Barcelona) Spain Spain Phone: 900 800 303 Portugal Phone: 800 827 836 Email: EMEAComplaints@zimvie.com</p>
<p>Switzerland Biomet 3i & Zimmer Dental Biomet 3i Schweiz GmbH Grüzefeldstrasse 41 CH-8404 Winterthur, Switzerland Phone: +41 (0)800 24 66 38 Fax: +41 (0)800 24 66 39 Email: EMEAComplaints@zimvie.com</p>	<p>Biomet 3i (Biomax) Biomax SPA Via Zamenhof, 615 Vicenza, Italy Phone: +39 0444 913 410 Email: info@biomax.it</p>	<p>UK and Ireland Biomet 3i & Zimmer Dental Biomet 3i UK, Ltd Reading Business Centre, Suite 807, 8th Floor Fountain House 2 Queens Walk, Reading, Berks, RG1 7QF, United Kingdom UK Phone: +44 (0) 800 652 1233 Ireland Phone: +353 1800 552752 Email: EMEAComplaints@zimvie.com</p>	

Patient Specific Product (PSP)

Biomet 3i Dental Ibérica
 BellaTek Dept.
 Islas Baleares 50,
 Polígono Fuente del Jarro
 46988 Valencia Spain
Phone: +34 961379536 / 38
Fax: +34 961379505
Email: es.3ipsp@biomet.com

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