

Incident- / Malfunction Report

Incident

(for Malfunction Report go to page 3)

All pages should be emailed to the local distributor within 48 hours of knowledge of the incident.

Local contact:	Physician's stamp (name + address)	
1. Patient Details		
Initials: Sex: Fen	nale Male	
Date of Birth:		
2. Indication		
Urinary Incontinence (UI): V	/esicoureteral Reflux (VUR):	
3. Treatment details		
Date of treatment:	Lot no.: Total volume injec	ted:mL
4A. Procedure details for VUR	4B. Procedure details fo	r UI
STING Procedure: HIT Procedure: Double	e-HIT Procedure: Site of injections (please r for UI only:	nark) 9 (3
5. Details of complications		
Description of complication:		6
Onset date of complication:	Hospitalization required: Yes	No
Severity of symptoms: Mild Moderate	Severe Serious	
Treatment:		
Antibiotics: Yes No		
Other treatment: Yes No		
If yes, please describe the treatment (drug, (dosage	e, duration of treatment), surgery, catheter etc.):	



Patient's initials:_____

6. Relevant Medical History

Prior to treatment

Did the patient receive pharmacological treatment for urinary incontinence?	Yes	No
Did the patient suffer from Urinary Tract Infection (positive Nephur* stix)?		No
Did the patient receive treatment for Urinary Tract Infection?	Yes	No
Is the patient known to be allergic or hypersensitive?	Yes	No
Is the patient known to suffer from any connective tissue disease?	Yes	No
Had the patient received previous treatments with radiation in the pelvic floor?	Yes	No
Had the patient undergone previous surgery for the treatment of UI?	Yes	No
Had the patient undergone any previous surgery in the pelvic floor? If so, please name:	Yes	No
	Yes	No No
If so, please name:		
If so, please name:	Yes	No
If so, please name:	Yes	No No

5. Details of follow-up

Resolution of complication	
Total recovery:	_ Comments:
(Day/Month/Year)	
Is any further medical follow-up required?	Yes No
If yes, please specify:	
Do you need any further medical advice from C	ontura? Yes No
If yes, please specify:	

Please attach any additional relevant information.

(Day/Month/Year)

Details of injecting doctor:	tor:
Name of doctor:	
Clinic:	
Address of clinic:	
Profession (speciality):	
Telephone number:	
E-mail:	

_____ Signature of doctor: ___



Malfunction Report

Treatment of VUR (Vesicoureteral Reflux):

Treatment of UI (Urinary Incontinence):

This malfunction concerns

Medical Device	Lot no. / Serial no.	Device should always be returned to enable root cause analysis		
Bulkamid [®] Hydrogel, 1 mL prefilled syringe		Device enclosed	Device sent separately	
Bulkamid [®] Needle		Device enclosed	Device sent separately	
Bulkamid [®] Rotatable Sheath		Device enclosed	Device sent separately	
Whole Bulkamid [®] Kit		Device enclosed	Device sent separately	
Bulkamid [®] Urethroscope		Device enclosed	Device sent separately	
If device is not returned, please clarify why and explain what happened:				
Malfunction description:				

Date of malfunction	Did malfunction happen	Before bulking procedure: During bulking procedure: After bulking procedure:
Were there any risk to patient due to malfunction?	No 📕 Yes 📕 If yes	s please describe
Was the device handled in accordance with instructions for use? (e.g. was accidently dropped on the floor)	Yes No If no	please describe

If an incident occurred due to the malfunction, please complete the section for incidents in this form, page 1.

(Day/Month/Year)



Procedure for the return of Malfunctions of Bulkamid® Devices

All products that have been used or unpacked in the OR and have not been re-sterilized, must be packed in special yellow bags or boxes for contaminated utensils with biohazard warning before returning in order to indicate possible infection risks. It is important to capture the batch number and always send the device including the Incident - / Malfunction Report Form otherwise it is not possible to find out the cause of the malfunction.

Malfunctions of products from the Bulkamid[®] Kit, e.g. needles, sheath and syringes must be returned in original kitbox.

Bulkamid[®] Devices:

BULKAMID[®] ROTATABLE SHEATH:

The sheath must be packed in material for contaminated products as mentioned above.

BULKAMID[®] NEEDLE:

The needle must be sent with the protection sheath in place.

BULKAMID[°] SYRINGE:

The syringe must be sent in original blisterpack.

Bulkamid[®] Urethroscope:

The optic must be packed and shipped with the protection sheath and preferably in the original shipping box or similar protection. This is to prevent further damage during the transportation. The Incident - / Malfunction Report Form should be included besides the optic. Do not return the metal sterilization container.

Once you have completed this form, please return it to Complaints@contura.com.