# SI-BONE Complaint - Revision - Post Market - Feedback Reporting

Use this form to submit information about a complaint related to an SI-BONE product to Quality Affairs. Contact <u>QA@si-bone.com</u> if you have any questions.

# Select the type of problem you are reporting

Problem with individual device or individual patient

## **General Feedback about SI-BONE Product**

This is only for non complaint information. Examples: surgeon suggestions, meeting discussions, etc. For complaints, go back and select "Problem..."

#### **Contact Info**

Use this to record your attempts to contact and gather information from the surgeon.

Your name	Amanda Adams
Your email	amanda.adams@si-bone.com
Your phone number	(904) 466-2392
Surgeon's name	Robert Hurford
Contact #1: How did you attempt to contact the surgeon?	His distributor was my point of contact
Was this attempt successful?	Yes
contact the surgeon?	

#### Comments about attempts to contact surgeon (optional)

Requested visit with surgeon in clinic to discuss plan was unsuccessful. Attempted to visit while at St. Vincent's Southside, unsuccessful.

#### **Complaint Overview**

Enter information about surgery that resulted in a problem with any SI-BONE implant, instrument or accessory (e.g., drill bit, pin).

Date you first heard of problem with	Thursday, January 5, 2023
SI-BONE product.	

Which device(s) was/were affected?

iFuse-TORQ

#### Part number(s), if available

11.5x40 iFuse TORQ fully threaded 11.5x35 iFuse TORQ fully threaded



#### Lot number(s), if available

N/A, will have to get RPO from Nicole Hollinghead TR.

### **Product Complaint or Adverse Event?**

Decide what type of report you are submitting.

Use this form to submit information about a complaint related to an SI-BONE product to Quality Affairs. Contact <u>QA@si-bone.com</u> if you have any questions.

# What is the nature of the problem you are reporting?

Product complaint that caused or may have caused an adverse event or revision procedure

# **Product Complaint Without Adverse Event**

Use this form to report an SI-BONE product problem that was NOT associated with a patient adverse event.

Examples include:

- Damaged instrument
- Broken pin
- Bent pin
- Pin advancement but no patient injury
- Problem with packaging

If a damaged part injured a patient, go back and click on "Product complaint that caused or may have caused an adverse event."

# Select Adverse Event Type

Select from the choices below. Your choice guides which section to complete.

If the first 3 do not apply, select "Other".

If a product problem occurred that did NOT cause a patient problem, go back and complete the "Product Complaint Without Adverse Event" section.

What problem did patient have?

Other problem

## **Symptomatic Implant Malposition Form**

Use this form if patient experienced symptomatic implant malposition resulting in nerve irritation.

(Information should be from SI-BONE staff who attended initial surgery)

The surgeon completed all steps in the IFU, including:

- Patient setup (table, spine neutral position, prone or supine position)
- Inspection of instruments and implants prior to use
- Proper views (lateral, inlet and outlet) obtained during use of instruments and placement of implants that clearly showed recommended anatomic landmarks
- Appropriate implant starting position
- (If performed) proper use of neuromonitoring guide pin and equipment Use of length gauge for implant length selection
- Were standard wound closure techniques used?

For above question, be specific and include all steps of the revision surgery and any complications.

- Was the implant adjusted, removed or added?
- Were the explant holes filled with bone graft?
- Were osteotomes and the/or the removal adapter used?



• Was any non-iFuse hardware installed?

## Pain Did Not Improve or Recurred

Use this form if pain did not improve OR pain improved but then recurred

(Information should be from SI-BONE staff who attended initial surgery) The surgeon completed all steps in the IFU, including:

- Patient setup (table, spine neutral position, prone or supine position)
- Inspection of instruments and implants prior to use
- Proper views (lateral, inlet and outlet) obtained during use of instruments and placement of implants that clearly showed recommended anatomic landmarks
- Appropriate implant starting position
- (If performed) proper use of neuromonitoring guide pin and equipment Use of length gauge for implant length selection
- Were standard wound closure techniques used?

### **Surgical Wound Problem**

Use this form if patient developed postoperative surgical wound problem (infection, dehiscence).

# **Other Problem**

Use this section ONLY if the patient problem is NOT:

- Implant malposition causing nerve irritation
- Pain did not improve or recurred
- Surgical wound problem

#### Describe problem in detail

The middle and inferior iFuse TORQ implants did not fuse, and patient complained of pain.

### **Potential Warranty Case**

Please help evaluate whether this case could qualify for SI-BONE's warranty policy.

Did patient's situation meet all of the following?

Did patient undergo a revision procedure that met all of the following:

- 1. The initial procedure involved placement of 3 SI-BONE implants on the affected side
- 2. The revision procedure:
  - Was completed within 1 year of the initial procedure
  - Was completed at the same hospital or within the same hospital system as the initial procedure
  - Was completed during a separate hospitalization from the initial procedure
- 3. The patient followed all physician instructions after the initial procedure

# Did patient's situation meet warranty criteria shown above?

No, not all of the above apply

