

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.

## **Contact Info**

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

failure

Kendrick Wroe

**Customer name** 

Kendrick Wroe

How did you learn about this issue? (select all that apply)?

I observed the issue

# Please provide any relevant details about your communication. Full complaint description will be captured on the following page:

TESTING ONLY. ALL FIELDS CHECK

## **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<br/>SI-BONE product.Tuesday, October 3, 2023Date of original surgery (if revision is<br/>being reported) or alleged productWednesday, June 7, 2023

Indicate affected device(s) (choose all that apply)

iFuse-3D

#### Part number(s) (please list the number of each part involved)(required)

TESTING ONLY. ALL FIELDS CHECK

Lot number(s) TESTING ONLY. ALL FIELDS CHECK

# **Product Complaint or Adverse Event?**

Decide what type of report you are submitting. Contact QA@si-bone.com if you have any questions.

# Did the product complaint result in a patient problem?

YES, potential or actual (Ex: required revision, patient adverse event)

# **Product Complaint Without Patient Problem**

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

Examples include:

- Damaged instrument/implant
- Broken/ bent/ cut pin
- Pin advancement but no patient injury
- Packaging issue

If patient injury occured, go back and click YES to report patient problem.

#### (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?

# Select Adverse Event Type

#### What problem did patient have?

Implant malposition (e.g. causing nerve irritation)

# **Implant Malposition Form**

Use this form if patient an implant malposition was detected.

On which side(s) were SI-BONE implants placed during initial surgery?	Left	
Which side shows implant malposition?	Left	
Did any SI-BONE staff attend initial surgery?	Yes	
Name of SI-BONE staff in attendance at initial surgery	Kendrick Wroe	
Imaging type used during initial surgery	C-arm	n/fluoro only

(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?

#### See summary of IFU steps. Did surgeon complete all steps as shown above (regardless of post-op symptoms)?

#### Please describe procedure steps not done properly or other pertinent information

TESTING ONLY. ALL FIELDS CHECK

Did patient have revision surgery?

Yes

## Continued, recurrent, or new pain

Use this form if pain did not improve, pain improved but then returned, or new onset pain

(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

### **Revision Procedure**

Complete this if patient underwent revision surgery.

Please indicate date of revision procedure

Monday, October 30, 2023

Please indicate all procedural steps that apply:

iFuse implant was repositioned

**Provide as much detail on the case as possible:** TESTING ONLY. ALL FIELDS CHECK

Was the patient's issue resolved after surgery? TESTING ONLY. ALL FIELDS CHECK

# You may be contacted for further information if your submission is lacking critical details. We appreciate your thoroughness.