# **ANSER Clavicle Pin Set**

#### BAAT Medical Products B.V.

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## INSTRUCTIONS FOR USE

## **Device description**

The ANSER Clavicle Pin is a device intended to be used for intramedullary fixation of mid-shaft clavicle fractures. The device consists of three parts. The main component is a nail, that is placed within the medullary canal of the fractured clavicle. It has a threaded medial end for fixation into the bone. At the lateral end an endcap is placed after insertion for lateral fixation of the pin. In this way the pin is secured both medially and laterally.

## Intended purpose

The Anser Clavicle Pin Set is an intramedullary fixation device, intended for fixation of midshaft clavicle fractures. The goal of the Anser Clavicle Pin Set is to restore the native length and alignment of the fractured midshaft clavicle in a minimally invasive manner.

## Indications for use

Indications for the ANSER Clavicle Pin include skeletally mature patients suffering from mid-shaft clavicular fractures.

## Relative contra-indications <sup>1</sup>

- · Patients suffering from (severe) osteoporosis
- Patients who are not fit for surgery
- Patients who are not skeletally mature
- · Possible non-compliant patients (e.g. due to alcohol and drug addiction, dementia)
- Additional neurovascular injury
- Pathologic fractures
- <sup>1</sup> Relative contraindications are contraindications for circumstances in which the patient is at higher risk of complications from treatment, but these risks may be outweighed by other considerations or mitigated by other measures. For example, a pregnant woman should normally avoid getting X-rays, but the risk may be outweighed by the benefit of diagnosing (and then treating) a serious condition such as tuberculosis. Relative contraindications may also be referred to as cautions, such as in the British National Formulary.

## Warnings and pre-cautions

#### Warnings

- Avoid, if possible, using multiple insertion attempts during placement of the Anser Clavicle Pin in the medial fragment
  and placement of the Anser Lateral Fixation Device in order to prevent weakening/destruction of the (self-) tapped
  threaded interfaces.
- If the extracortical portion of the Anser Clavicle Pin is plastically deformed, unintentional advancement of the Anser Clavicle Pin may occur when using the Anser tap. Adjust or replace the Anser Clavicle Pin if plastically deformed or damaged.
- In case of incorrect positioning of the Anser Lateral Fixation Device it is not possible to place the Anser End Cap. Clean
  the implant implant interface and/or adjust the Anser Clavicle Pin or Anser Lateral Fixation Device to position the
  Anser Lateral Fixation Device correctly. Ascertain the six resilient legs of the Anser Lateral Fixation Device have positioned themselves in one of the indentations of the Anser Clavicle Pin.
- Equivalent intramedullary implants have a reported risk of hardware failure. The risk exists that the pin may plastically
  deform or break due to peak loading.

Do not use the Anser Clavicle Pin as a lever or "joystick" during reduction. It may deform or break.

- Proper alignment of the fracture elements before traversing the fracture site with the Anser Clavicle Pin is of utmost
  importance. Failure to align the fragments may lead to complications such as damage to the subclavian artery or the
  lung.
- The surgeon should strictly follow the recommendations in the surgical technique. All staff involved should be familiar
  with the surgical procedures associated with intramedullary fixation of mid-shaft clavicle fractures to avoid adversely
  affecting device performance or surgical outcome.
- The use of a powered instrument for drilling or driving the Clavicle pin in position can damage the anatomical structures or damage the implant or instruments.
- Only use instruments and accessories listed in the surgical technique to avoid adversely affecting device performance
  or surgical outcome. The surgical team must verify that the instruments are in good condition and in operating order
  prior to use during surgery.

## Pre-operative pre-cautions

- Please consult the applicable surgical technique for intended use, indications, contraindications, selection and use of
  a device and check the full labeling for other necessary information.
- Surgical technique brochures may be by requested from the distributor or from BAAT Medical Products directly.
- Those using brochures published more than two years before the surgical intervention are advised to obtain an
  updated version.
- BAAT Medical Products devices can only be used by surgeons who are fully familiar with the surgical technique required and who have been trained to this end.

- The operating surgeon must take care not to use the specialized instruments to exert inappropriate stress on the
  patient or the implants and must scrupulously comply with any operating procedure described in the surgical technique provided. For example, the forces exerted when repositioning an instrument in-situ must not be excessive as
  this is likely to cause injury to the patient.
- To reduce the risks of breakage, care must be taken not to distort the implants or nick, hit or score them with when the devices are used near vital organs, nerves or vessels.
- In the presence of flaws on the packaging or expiration of shelf life, the device is considered non-sterile, and must be discarded and disposed.

## Post-operative pre-cautions

Torque, Displacement and Image Artifacts according to ASTM F 2213-06, ASTM F 2052-06e1 and ASTM F2119-07

Non-clinical testing of worst case scenario in a 3 Tesla (T) MRI system did not reveal any relevant torque or displacement of the construct for an experimentally measured local spatial gradient of the magnetic field of 3.69 T/m. The largest image artifact extended approximately 169 mm from the construct when scanned using the Gradient Echo (GE). Testing was conducted on a 3 T MRI system.

## Radio-Frequency-(RF-)induced heating according to ASTM F2182-11a

Non-clinical electromagnetic and thermal testing of worst case scenario lead to peak temperature rise of 6.9.5 °C with an average temperature rise of 6.6 °C (1.5 T) and a peak temperature rise of 5.9 °C (3 T) under MRI Conditions using RF Coils (whole body averaged specific absorption rate (SAR) of 2 W/kg for 6 minutes [1.5 T] and for 15 minutes [3 T]).

Precautions: The above mentioned test relies on non-clinical testing. The actual temperature rise in the patient will depend on a variety of factors beyond the SAR and time of RF application. Thus, it is recommended to pay particular attention to the following points:

- It is recommended to thoroughly monitor patients undergoing MR scanning for perceived temperature and/or pain sensations.
- Patients with impaired thermoregulation or temperature sensation should be excluded from MRI scanning procedures.
- Generally, it is recommended to use a MR system with low field strength in the presence of conductive implants. The
  employed specific absorption rate (SAR) should be reduced as far as possible.
- Using the ventilation system may further contribute to reduce temperature increase in the body.

## Potential adverse effects

As with any major surgical procedure, there are risks involved in orthopedic surgery. Potential risks identified with the use of this system include, but are not limited to:

#### General surgery / procedure related

- Infection (superficial or deep infection)
- Neurological complications (e.g. brachial plexus palsy, nerve palsy, (temporary) paresthesia)
- Cardio/vascular complications (e.g. thromboembolic events)
- · Skin/wound related complications (e.g. skin irritation, skin perforation, keloid formation)
- (persistent) Pain
   Poor cosmetic appr
  - Poor cosmetic appearance
- · Other general surgery / procedure related complications (e.g. frozen shoulder)

## Device related

- Hardware irritation
- Hardware failure (e.g. breakage)
- Hardware migration (migration, protrusion, telescoping)
- Loss of fixation
- Delayed union
   Nonunion/malui
- Nonunion/malunion
   Befracture
- Reoperation/revision
- Other device-related complications (e.g. plastic deformation)

#### Disposal

The disposal of this medical product requires no special measures. Be sure to observe all national/local regulations and guidelines when disposing of the packaging material and potentially infectious items.

## Material specification

The ANSER Clavicle Pin Set design is fabricated from a titanium alloy (Ti-6AI-4V).

#### Packaging

Packages for each of the components should be intact upon receipt. To reduce the risk of infection the packaging of all sterile devices must be inspected for flaws in the sterile barrier or expiration of shelf life before opening. In the presence of such a flaw or expiration of shelf life, the device is considered non-sterile, and must be discarded and disposed.

## Recommended storage and handling conditions

The products shall be stored and handled with care. The primary, secondary and eventual tertiary packaging shall remain intact at all times.

## Sterilization

The ANSER Clavicle Pin Set components are packed and delivered together in one sterile package. Devices which are delivered sterile have been exposed to a minimum of 25kGy of gamma radiation. Only sterile products should be placed in the operative field.

## Product complaints

Any Health Care Professional (e.g., customer or user of this system of products), who has any complaints or who has experienced any dissatisfaction in the product quality, identity, durability, reliability, safety, effectiveness and/or performance, should notify the distributor or BAAT Medical Products BJ. Further, if any of the implanted devices ever malfunctions, (i.e., does not meet any of its performance specifications or otherwise does not perform as intended), or is suspected of doing so, the distributor should be notified immediately. If any ANSE product ever "malfunctioned" and/or may have caused or contributed to the death or serious injury of a patient, the manufacturer should be notified immediately. If telephone, fax or written correspondence. When filing a complaint, please provide the component(s) name and number, lot number(s), your name and address, the nature of the complaint and notification of whether a written report from the manufacturer is requested.

## End-user information

Further copies of the surgical technique, instruction for use, can be requested at BAAT MEDICAL Products B.V.

## Explanation symbols used in end-user information

LOT	Batch code
REF	Catalogue number
$\sim$	Date of manufacture
	Manufacturer
) III)	Consult Instructions for Use
$\mathbb{N}$	Use-by date
$\otimes$	Do not re-use
8	Do not use if package is damaged
R	Do not resterilize
NON	Non-sterile
STERILE R	Sterilized using irradiation