Surgical Guideline

Mi1200 SYNCHRONY ABI
Mi1200 SYNCHRONY PIN ABI
Introduction

The MED-EL SYNCHRONY/SYNCHRONY PIN Auditory Brainstem Implant System serves to restore some sense of hearing by electrical stimulation of the cochlear nucleus (CN). It is the result of many years of research and is intended for use in patients with non-functional cochlear nerves.

This Surgical Guideline describes proper techniques for implanting the Mi1200 SYNCHRONY/SYNCHRONY PIN Auditory Brainstem Implant (ABI) (hereafter referred to as the SYNCHRONY/SYNCHRONY PIN ABI). It serves as additional information for professionals and should not be used as an "Instructions for Use".

MED-EL implants are manufactured to the highest quality standards in order to ensure long term reliability. All materials used in the implant have been rigorously tested for biocompatibility, durability and reliability. MED-EL applies a quality management system that meets all EN ISO 13485:2003 requirements and complies with US Quality System Regulations and Canadian Medical Device regulations (CAN/CSA ISO 13485-2003). Components of the MED-EL Cochlear Implant System meet the requirements for AIMD 90/385/EEC and MDD 93/42/EEC.

The information in this Surgical Guideline is believed to be true and correct, however, some specifications may be subject to change without notice.

For country specific information please see the applicable “Instruction for Use” delivered with the implant system.
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I. Patient selection and evaluation

Indication

The SYNCHRONY/SYNCHRONY PIN ABI is used for electrical stimulation of the cochlear nucleus (CN) via an implanted stimulator and a specially designed electrode array to evoke auditory sensations in patients with non-functional cochlear nerves. Likely cause of damage may include trauma, inflammation or in most cases the degeneration of the cochlear nerve, e.g. related to a bilateral acoustic neurinoma (Neurofibromatosis 2 - NF2).

Selection and Evaluation

If patients suffering from NF2, it is the responsibility of the surgeon to determine for each patient if implantation of the device during tumor removal surgery is appropriate. Most NF2 patients also suffer from multiple tumor formations in the spine. The personal situation of these patients may therefore be very difficult and critical.

Patients who are selected for an ABI implantation should have a stable psychological status and be willing to learn to use the ABI system. To obtain the optimal benefit from the implant, candidates must be sufficiently motivated and must understand the importance of returning to the implant center for regular audio processor programming, training, and assessment sessions. Preoperative psychological tests shall be performed. In addition to the mental condition of the patient, the physical condition shall also be assessed for an implantation. Tumor size and cochlear nerve condition should be assessed preoperatively either by performing MRI and/or CT scanning and electrophysiological measurements (ABR, EABR, eCochG). Also a preoperative spinal MRI is essential. Experienced Neuro- and ENT-surgeons shall decide in accordance with the test results and their own personal experience if a patient is a suitable candidate for an ABI implantation.

If previous irradiation of the tumor has taken place, the patient's evaluation should include possible structural deformation caused by such irradiation. Large structural deformation may prevent proper ABI placement and should be evaluated by the surgeon. Before surgery, the patient must be informed about the function of the implant, the risks of surgery and implantation, realistic expectations and rehabilitation plans after surgery. A personal conversation between patient and surgeon should help the patient to gain a general idea about the device and to estimate risks and benefits of the Implant SYNCHRONY/SYNCHRONY PIN ABI.

This device is to be used by surgeons who have been trained in the surgical placement of an Auditory Brainstem Implant.
II. Technical description of the ABI System

The SYNCHRONY/SYNCHRONY PIN ABI is the implantable part of the MED-EL Cochlear Implant System and can only be used together with compatible MED-EL external components.

The device consists of a stimulator, a coil with a removable magnet within its centre, a reference electrode, an EAP reference electrode and an active electrode permanently attached to the stimulator. This device is intended to be implanted by adequately trained and experienced surgeons only.

The SYNCHRONY/SYNCHRONY PIN ABI has been designed according to the highest safety and reliability standards. All materials used in the construction of the SYNCHRONY/SYNCHRONY PIN ABI have been extensively tested for biological compatibility and durability. The power required by the implant is transmitted from the external audio processor through the intact skin via an inductive link. The implant therefore contains no batteries or other components that require replacement.

The implant offers a stimulation mode and a telemetry mode. Stimulation sequences of biphasic and triphasic pulses can be delivered sequentially or simultaneously on two or more channels. In telemetry mode the device allows a functional check about the technical status of the implant including communication over the transcutaneous link as well as the assessment of the electrode impedances and recording of the electrically evoked compound action potential of the hearing nerve.

The electronics of the SYNCHRONY/SYNCHRONY PIN ABI contain a powerful custom-made circuit that is capable of processing large amounts of information at a very rapid rate. It can stimulate at 50,704 pulses per second. This capability makes the implant compatible with a wide range of pulsatile processing strategies and future developments in speech processing. A telemetry feature enables the clinic to verify the functional status of the implant within a matter of seconds. For added safety, each output has a capacitor to prevent any possible leakage of DC current to the auditory nerve.

Figure 1 Implant SYNCHRONY/SYNCHRONY PIN ABI (~ dimensions in mm, typical values)
Performance Characteristics

- Output characteristics of a stimulation signal on a 1 kOhm resistor:
  - Maximum current amplitude:
    - Median value = 1250 µA, range = 500 µA
  - Maximum pulse width:
    - Median value = 203.8 µs, range = 8.2 µs
- The impedance measurement accuracy is typically better than 5%.
- When keeping to the safety guidelines the implant is conditionally MRI safe for MR scanner field strengths of 0.2 Tesla, 1.0 Tesla and 1.5 Tesla.
- There are no default factory settings of the implant system.
- Proper functioning of the implantable part of the CI system can be checked by performing telemetry (refer to MED-EL application software user manual).
- The implant has 24 independent current sources stimulating 12 independent electrode channels in monopolar mode.
- The implant has a mass of 7.6 g (typical value).
- The volume of the implant without electrode is 3.7 cm³.
- The electrode is made of medical grade silicone, platinum (electrode contacts) and platinum/iridium (90/10) wires and nitinol.
- All electrode variants have a straight and flexible design. The electrode does not deliver any medicinal substances.
- Geometric surface area of the stimulation reference electrode = 50 mm².
- Following materials are in direct contact with human tissue: medical grade silicone, platinum, iridium, titanium and parylene c.

ABI Active Electrode

Order number SYNCHRONY ABI 31095
Order number SYNCHRONY PIN ABI 31096

The ABI Active Electrode has an oval shaped flat silicone paddle (electrode array) with 12 active contacts and 1 reference contact. A polyester mesh embedded in silicone exceeds the size of the silicone paddle.

This paddle is slightly pre-shaped to fit onto the curved brain surface.

An additional contact is placed in the center of the silicone paddle to allow bipolar stimulation mode during intraoperative measurement. The diameter of the electrode lead increases from ≈ 0.7 mm at the silicone paddle to ≈ 1.3 mm over a length of ≈ 10 mm.

Designed Especially for Neurofibromatosis Type II

ABI is a solution for individuals with hearing loss due to a non-functioning auditory nerve (Neurofibromatosis Type II). Bypassing both the inner ear and the auditory nerve, the MED-EL ABI stimulates the cochlear nucleus (CN) and provides users with a variety of hearing sensations to assist with sound awareness and communication.

![ABI Electrode Array](Figure 2 ABI Electrode Array (~ dimensions in mm, typical values))
AB Placing System

CAUTION
For detailed information on the AB Placing System, MED-EL hardware interface system and MED-EL application software please refer to the applicable instruction for use and user manual.

The AB Placing System for the Implant SYNCHRONY/SYNCHRONY PIN AB is designed exclusively for transient intraoperative stimulation of the cochlear nucleus. During stimulation of the cochlear nucleus, Electrical Evoked Auditory Brainstem Responses (EABR) shall be recorded, to estimate the best position for the AB electrode array.

The AB Placing System consists of:
- AB Placing Electrode
- AB Connector Cable
- AB Stimulator Box

AB Placing Electrode
The electrode array and the lead of the AB Placing Electrode have the same dimensions as the SYNCHRONY/SYNCHRONY PIN AB electrode array. The paddle has only 4 electrode contacts instead of 12 + 1 and there is no polyester mesh. The Micro-D Plug is the connective part of the AB Placing Electrode and shall be connected to the AB Connector Cable (see section IV Set-up and recommended measurement for EABR recording).

Figure 3 AB Placing Electrode (~ dimensions in mm, typical values)
Technical description of the ABI Placing System

ABI Connector Cable
The ABI Connector Cable is designed to transfer stimulation pulses from the ABI Stimulator Box to the ABI Placing Electrode. The length of the cable is ≈2 m so that it can lead out of the patient’s environment. The connector on the left side of the ABI Connector Cable (see Figure 4) shall be connected to the ABI Placing Electrode (Micro-D Plug). The connector on the right side of the ABI Connector Cable (see Figure 4) shall be connected to the ABI Stimulator Box (see Figure 5).

ABI Stimulator Box
Pre-use check of the ABI Stimulator Box

CAUTION
Do not use an ABI Stimulator Box for intra-operative stimulation if the ABI Stimulator Box is not functioning correctly.

The ABI Stimulator Box generates biphasic stimulation pulses controlled by the MED-EL hardware interface system via the coil placed on the ABI Stimulator Box. The selector switch (see Figure 5) allows easy selection of the electrode configuration at the ABI Placing Electrode which is used for bipolar stimulation during EABR measurements.
Prior to the use of the ABI Placing System, a pre-use check shall be performed with the ABI Stimulator Box.

**Note:** Only personnel trained on the use of the MED-EL telemetry system shall perform the pre-use check.

The following steps shall be followed when performing the pre-use check:

**Setup of measurement**

Connect the ABI Stimulator Box Test Device to the ABI Stimulator Box.

**Telemetry**

Start the telemetry measurements at the selector switch configuration 1=>2 and proceed clockwise.

**Note:** The impedance measured at switch position 1=>2 shall be lower than the impedance measured at switch position 1=>3. All subsequently measured impedance values shall be lower than the values measured at the switch position before.

**Acceptance criteria**

The difference of the measured impedance values between each step shall be greater than 0.4 kΩ. All measured values shall lie between 1.5 kΩ and 9 kΩ. If the measured values fulfill the acceptance criteria the ABI Stimulator Box can be used for intra-operative stimulation together with ABI Placing Electrode and the ABI Connector Cable.

**Note:** Only the impedance value measured at Channel 1 shall be used as information for the pre use check. All other channels show HI (High impedance) as measured values.

If the measured values do not fulfill the requirements please return the ABI Stimulator Box to MED-EL for maintenance.
Set-up and recommended measurement for EABR recording

III. Set-up and recommended measurement for EABR recording

Set-up for ABI Placing System

Before starting the measurement, the ABI Placing Electrode shall be connected to the ABI Connector Cable via the Micro-D Plug (see Figure 9 for correct connection). **Note:** Make sure that the connection is as shown in Figure 10.

The ABI Connector Cable shall lead out of the immediate patient environment and shall be connected to the ABI Stimulator Box (used for ABI Placing Electrode only). The ABI Stimulator Box is driven via an inductive link by the MED-EL hardware interface system, which triggers the EABR measurement system. It is also possible that the EABR measurement system triggers the MED-EL hardware interface system.

**Note:** The integrity of the electrical pathway between the ABI Placing Electrode, ABI Connector Cable and the ABI Stimulator Box (and the functioning of the ABI Stimulator Box) shall be checked before starting the EABR measurement with a telemetry measurement of channel one. This measurement shall be done with the selector switch position 1-2 and 3-4. Make sure that the ABI Placing Electrode is inserted into the lateral recess.

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**Figure 9 Correct connection of the ABI Placing Electrode with the ABI Connector Cable**

**Figure 10 Setup for EABR recording with ABI Placing System**
Set-up for SYNCHRONY/SYNCHRONY PIN ABI System

**CAUTION**
Prior the EABR measurement impedance telemetry should be performed.

For the EABR set-up using SYNCHRONY/SYNCHRONY PIN ABI system, the coil from the MED-EL hardware interface system is directly placed on the SYNCHRONY/SYNCHRONY PIN ABI implant (Figure 11). Before starting the measurement, the ABI Electrode shall be placed on the brainstem.

**Recommended measurement for EABR recording**

**EEG – Electrode Placement**
Before surgery, EEG-Electrodes are placed on the contralateral mastoid (negative), on the vertex (positive) and on the lower forehead (ground). These electrodes are connected during surgery to the EABR recording system. Try to avoid placing any two electrodes too close together.

**Impedance of EEG-Electrodes**
The impedance of any recording electrode should be less than 2 kΩ. There should be no greater difference between the individual electrode impedanceless than 2 kΩ.

**General set-up of the EABR measurement system**
- **Window:** 10 ms
- **Points:** 256
- **Gain:** 20,000 - 50,000
- **Hi filter:** 3000 Hz
- **Lo filter:** 30 – 100 Hz
- **Sweep:** 200 – 1500
- **Trigger:** external (MED-EL hardware interface system triggers the EABR measurement system)

**Note:** It is also possible that the EABR measurement system triggers the MED-EL hardware interface systems (internal).

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Figure 11 Setup for EABR recording with SYNCHRONY/SYNCHRONY PIN ABI implant.
Set-up and recommended measurement for eABR recording

Using the MED-EL application software

EABR Task

Channel selection for ABI Placing System

Select channel 1 for stimulation during intraoperative measurement.

Note: The selection of the active contact pair at the ABI Placing Electrode is made with the selector switch of the ABI Stimulator Box. The selected stimulation configuration (1-2, 1-3, 1-4, 2-3, 2-4, 3-4) can be chosen via the selector switch. The selected stimulation configuration shall not be changed during stimulation in order not to influence the measured EABR curves.

Stimulation parameters

The stimulation parameters are set via the MED-EL application software.

Note: This setting shall be done by trained personnel only. The measurement typically starts with an electrode configuration of 1-4 selected with the ABI Stimulator Box. The stimulation parameters are set to pulse duration of 30 µs/phase and a stimulation current of 300 cu (current units).

Note: The following limits shall not be exceeded:
• 1000 cu
• 150 µs

Extended setup

For EABR recording, single stimulus pulses are used. Alternating pulses can be used to reduce the stimulus artifact. Leave the HF signal turned on. Stimulation rate: between 10 Hz and 35 Hz

EABR curve

Wave III generated at the cochlear nucleus is normally overlapped by the stimulus artifact during EABR recording. Waves IV and V become visible at a latency of 1 to 2 ms.

Note: Figure 12 shows a theoretical recording. During intraoperative EABR measurements only one wave (IV or V) is visible (at the recording monitor).

Figure 12 Theoretical EABR recording (stimulation of the cochlear nucleus – Waring 1993)
IV. Surgical tools

Note that the surgical tools supplied by MED-EL should not be modified in any way. Modification of any of the tools is done at the surgeon's own risk.

Detailed instruction of the reprocessing process and the individual preparation before cleaning the tools can be found in the appropriate Instruction for Use.

This is a collection of tools for implantation of the SYNCHRONY/SYNCHRONY PIN ABI Implant.

<table>
<thead>
<tr>
<th>Tool Description</th>
<th>Order number</th>
</tr>
</thead>
<tbody>
<tr>
<td>Mi1200 Implant Template</td>
<td>Shipped with the implant</td>
</tr>
<tr>
<td>Mi1200 Implant Template, PIN</td>
<td></td>
</tr>
<tr>
<td>PIN Drill Guide Si</td>
<td>09906</td>
</tr>
<tr>
<td>Processor Template</td>
<td>01557</td>
</tr>
<tr>
<td>Skin Flap Gauge 6</td>
<td>03543</td>
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<tr>
<td>Surgical Claw Angled</td>
<td>00284</td>
</tr>
<tr>
<td>Micro Forceps Angled</td>
<td>05761, 05777, 05778</td>
</tr>
</tbody>
</table>

**Mi1200 Implant Template**

**Mi1200 Implant Template, PIN**

Shipped with the implant

This template is used to assess the size and the position of the implant on the skull.

This tool is delivered in a sterile packaging and is a single-use device only.

Figure 13 Mi1200 Implant Template

Mi1200 Implant Template, PIN
PIN Drill Guide SI
Order Number: 09906

The PIN Drill Guide SI is a re-usable surgical instrument for creation of defined holes on the skull for the fixation pins of the SYNCHRONY PIN Cochlear Implant. It consists of a PIN Drilling Template made of titanium and a Clamping Handle made of stainless chromate steel. The device is especially recommended for surgeons using a small incision.

This tool is a re-usable surgical instrument for transient use. The device is delivered non-sterile.

Processor Template
Order number: 01557

The Processor Template (TEMPO+/OPUS template) shows the minimum spacing which must remain free behind the ear so that the external coil and the BTE Audio Processor do not interfere with each other when worn by the patient post-operatively. This tool is a re-usable surgical instrument for transient use made from medical grade stainless steel. The device is delivered non-sterile.

Skin Flap Gauge 6
Order number: 03543

Skin Flap Gauge 6 is used to evaluate the thickness of the skin flap in the area covering the cochlear implant. A skin flap thickness of 6 mm or less is recommended for a good magnetic hold and optimal signal transmission. Skin flaps thicker than 6 mm should be thinned out.

This tool is a re-usable surgical instrument for transient use made from medical grade stainless steel. The device is delivered non-sterile.
Micro Forceps Angled
Order number: 05761 right- & left-angled
05777 right-angled
05778 left-angled

The Micro Forceps left angled and the Micro Forceps right angled are used to grip, hold and insert the electrode without damaging it. Which kind of angled Micro Forceps is used to insert the electrode depends on the preference of the surgeon. In the closed position, the tips of the forceps are parallel to each other, separated by a distance of 0.25 mm.

This tool is a re-usable surgical instrument for transient use made from medical grade stainless steel. The device is delivered non-sterile.

Surgical Claw Angled
Order number: 00284

The claw can help to position and manoeuvre the electrode.

This tool is a re-usable surgical instrument for transient use made from medical grade stainless steel. The device is delivered non-sterile.
The following tools are additions to the surgical kit and may be ordered separately:

**Surgical Claw Straight**  
*Order number: 07711*

The Surgical Claw can help to position and insert the electrode array into the cochlea. The tip of this instrument is straight.

This tool is a re-usable surgical instrument made from medical grade stainless steel. The device is delivered non-sterile.

**Mi1000 PIN Drill Guide**  
*Order number: 07613*

The Mi1000 PIN Drill Guide is a re-usable surgical instrument for creation of defined holes in the skull for the pins of MED-EL hearing implant housing variants with pins. It consists of a PIN Stimulator Template made of titanium and a Clamping Handle made of stainless chromate steel. The device is delivered non-sterile.
Magnet Replacement Kit
Order number: 09693
Consisting of:

Non-Magnetic Spacer Ms010107
The Non-Magnetic Spacer (see Figure 21) is intended to be used as placeholder for the regular implant magnet of the Mi1200 Hearing Implant during MRI procedures, when a reduced image artifact is desirable.

Replacement Magnet Ms010108
The Replacement Magnet (see Figure 22) is intended to be used after an MRI, as replacement of the original implant magnet of the Mi1200 Hearing Implant and to restore full functionality of the Mi1200 Hearing Implant.

Magnet Tool Kit
Order number: 09734
Consisting of:

Magnet Removal Tool Ms050206
Magnet Insertion Tool Ms050205

The Magnet Removal Tool (see Figure 23) is for removal of the MED-EL removable implant magnet and the Non-Magnetic Spacer.

The Magnet Insertion Tool (see Figure 24) is for insertion of the Non-Magnetic Spacer and the Replacement Magnet.

The instruments are made of surgical grade stainless steel. The devices are delivered non-sterile.
FENTEXmedical Forceps

FENTEXmedical GmbH is specialized in the development, manufacturing and marketing of surgical instruments and visualization systems for ENT, Head & Neck and Facial Surgery.

Basic description of the device:
CI Electrode Insertion Forceps L=155 mm, with longitudinal groove, for electrodes with a basal diameter in the range 0.8 – 1.3 mm

FENTEXmedical forceps have been successfully tested at headquarters with all MED-EL electrode arrays. This surgical tool is no MED-EL product and may therefore be ordered directly at your local FENTEXmedical distributor.

http://www.fentexmedical.com/

AESCULAP Micro Forceps, bayonet-shaped

It is recommended to use a micro forceps (bayonet-shaped, 200 mm long and with rounded edges at the tip with a width of 0.6 mm) during placement of the ABI electrode array and the ABI Placing Electrode to protect the leads of the electrodes against damage. This is a standard surgical tool for brain surgery.

This tool is a re-usable surgical instrument made from medical grade stainless steel. The device is delivered non-sterile.

Supplier:
AESCULAP AG & CO. KG
Am AESCULAP-Platz
D-78532 Tuttlingen
Germany

Technical Data:
Micro forceps, bayonet-shaped
Yasargil profile handles
200 mm long
Width at the tip 0.6 mm (rounded edges)
V. General remarks about the surgery

The implant is fixed in a bed drilled in the mastoid, and the ABI electrode array is inserted into the lateral recess [LR] of the IV. ventricle. The implant position should maximize the available electrode lead.

Surgical notes

- Sterility of the Implant SYNCHRONY/SYNCHRONY PIN ABI, the ABI Placing Electrode and the ABI Connector Cable must be ensured at all times.
- Prophylactic use of antibiotics is recommended for all patients unless medically contraindicated. The surgeon should prescribe adequate dosing for each patient’s condition.
- The SYNCHRONY/SYNCHRONY PIN ABI electrode array shall be inserted into the lateral recess without compressing the electrode lead or touching the electrode contacts.
- Only MED-EL approved and recommended surgical instruments should be used during the insertion process, other instruments (probes, hooks, forceps, tweezers, etc.) can damage the electrode or the other parts of the device. The implant contains a strong magnet. Never use magnetic surgical tools.
- The stimulator shall be securely anchored in a bed drilled in the mastoid bone and the electrode lead protected in a bony channel without sharp edges in such a way that there will be no postoperative movement. Movement results in fatigue and possible subsequent failure of electrical connections.
- Uni- or monopolar diathermy may not be used after the implant is introduced into the surgical field as it may cause damage to the implant.
- The implant must never be dropped onto a hard surface; damage to the stimulator or electrodes during the operation will invalidate the warranty.
- The serial number of the implant must be visible on the implant before fixing it in place.
- Other risks after surgery may be avoided by following the instructions in the applicable MED-EL SYNCHRONY/SYNCHRONY PIN ABI instruction for use.
- In order to achieve good magnetic holding power and optimal coupling the distance between the lateral side of the implant and the surface of the skin (with hair) shall not exceed 6mm.
- Do not place the sutures directly over the electrode lead.
VI. Surgical procedure

General information

In general, an ABI implantation is performed as a procedure for restoration of sound perception of NF2 patients immediately after resection of an acoustic neuroma. Patients with other lesions involving the cochlear nerve and causing massive hearing impairment up to deafness may also be candidates.

For the resection of an acoustic neuroma, which is the main indication for receiving an ABI, it is recommended to use the semi-sitting position of the patient as a result of clinical experience. The head is fixed in a Mayfield clamp and rotated approximately 30° to the affected side. This rotation is necessary for a direct approach to the dorsal side of the petrous bone, and, in addition, the opening of the lateral recess rotates towards the surgeon, so that access is facilitated.

The reported advantage of this semi-sitting position is an intraoperatively well-balanced blood circulation in the body of the patient. Therefore, the blood pressure in the venous vessels of the head decreases which is advantageous to the surgical procedure. It is also possible to perform this surgery with the patient in a horizontal position, but it is not recommended here.

The use of electrosurgery in ABI patients due to further tumor removal surgeries is likely. If an electroscalpel is used in close proximity of the SYNCHRONY/SYNCHRONY PIN ABI electrode, the electrode could be damaged by the electrical current and cause a system failure. The surgeon shall not use an electroscalpel while operating close to the SYNCHRONY/SYNCHRONY PIN ABI electrode. Electroscalpel use during operations not close the electrode poses no risk for the implant.

STEP 1: Placement of the EABR Recording Electrodes

After the patient is in a semi-sitting position, as recommended above, the EEG electrodes are placed on the contralateral mastoid (negative), on the vertex (positive) and on the lower forehead (ground). These electrodes shall be connected during surgery to the EABR recording system. (Also refer to the section III Set-Up and Recommended Measurement for EABR Recording.)
STEP 2: Mark Implant Position

Place the Processor Template behind the ear and position the Mi1200 Implant Template. There are various orientation options. A suggested orientation for each ear is shown in Figure 27 and Figure 28, but the orientation depends on various factors, like e.g. the curvature of the skull.

Make sure that the length of the electrode lead allows placement of the electrode paddle without compressing or extending the lead. The available length of the electrode lead is depending on the position of the implant bed. Rotating the implant clock or counterclockwise might increase the available length.

Position the implant template in such a way that the SYNCHRONY/SYNCHRONY PIN ABI will be in the hair bearing area. The lower part of the stimulator should be under or close to the temporal line, with an angle between 35° and 60°. The electrode exits on the lateral side of the implant. Therefore the electrode lead comes out superiorly for the left ear and inferiorly for the right ear.

Once the implant template is in place, surgical ink may be used to mark its position on the surface of the skin. Surgeons may choose to transpose the position of the implant template onto the surface of the bone by using a hypodermic needle inserted perpendicularly to the skin at points along the side of the implant template.

When implanting a patient bilaterally care should be taken of the placement of the implants. In particular the second side should be placed specifically to match the location of the first to give symmetric appearance of the external part. The skull curvature and pinna position needs to be taken into consideration when placing the second implant in a similar location as the contralateral side.

Figure 27 (left ear): Suggested orientation of the templates

Figure 28 (right ear): Suggested orientation of the templates
STEP 3: Plan Incision

Choose the line of incision so that a well vascularised skin flap results. Make the incision 1–2 cm from the implant to ensure that the scar will not lie directly over the body of the implant. Incise the tissue with a scalpel and use bipolar electrocoagulation for hemostasis.

An example of a commonly used postaural incision is shown in Figure 29 and Figure 30. Postaural incisions start in the sulcus behind the pinna and extend posteriorly.

For greater mastoid bone exposure, each of these incisions can be extended posteriorly in the shape of an arc.
STEP 4-A: Open Skin Flap

The incision is made and the wound is held open by retractors. At all times care should be taken to ensure that the flap is kept moist with damp surgical gauze.

Either a single layer skin flap – all four layers, skin, subcutis, muscle and periosteum are incised in a single cut, or a double layer skin flap can be performed.

A double layer skin flap may:
• reduce the chance of infection because the incisions are at different locations and layers, and
• allow better healing so it is often used for re-implantations and when encountering postauricular scar formation.

Double layer skin flap (see Figure 31)
The four different tissue layers skin, subcutis, muscle, and periosteum are incised with two different incisions. First, the skin, subcutis and muscle are raised and retracted. Second, the periosteum is incised, the periosteum is freed from the surface of the bone and then retracted in another location.

Various methods may be used when incising the periosteum. Care should be taken to avoid incision over the implant later on.
STEP 4-B: Skin Flap Thickness

In order to achieve good magnetic hold and optimal signal transmission, the skin flap or the muscle may need to be thinned out so it does not exceed 6 mm.

Evaluate the portion of the flap over the magnet and receiving coil with the Skin Flap Gauge 6, as shown in Figure 32. If the flap does not fit in the gauge loosely, carefully thin the flap until it does. It is important to avoid over-thinning of the flap, which may result in wound complications. Care must be taken to avoid exposing hair follicles.

Figure 32 Using the Skin Flap Gauge 6 – right ear
STEP 5: Tumor Removal

Surgery

If necessary, the acoustic neuroma is removed in a standard tumor removal surgery. Either a suboccipital or a translabyrinthine approach can be used for this surgery. A suboccipital approach makes the preservation of the cochlear nerve during tumor removal possible due to a lateral perspective. By stimulating the tissue in the area of the cochlear nerve and measuring EABR potentials intraoperatively with special ball electrodes, the boundary between tumor and nerve tissue can be determined electrophysiologically. If the cochlear nerve is not yet destroyed by the tumor, it can be uncovered from tumor tissue and functionally preserved with this method. A suboccipital approach is recommended by the expert surgeons who shared their experience for these guidelines.

The opening in the skull normally has a diameter of 25 – 35 mm. The brain is retracted from the skull with a dissector for obtaining access to the region of the cochlear nerve. For ABI implantation via a suboccipital approach, a retrosigmoidal trepanation is used. If the opening is not located correctly (too high or too far back), the lead of the Auditory Brainstem Implant may be too short and the SYNCHRONY/SYNCHRONY PIN ABI electrode array not placeable in the lateral recess. A correct location of the trepanation is therefore of high importance.

STEP 6: Preparation of the Lateral Recess

After complete tumor resection, the exits of the nerves VII., VIII. and the caudal groups IX. and X. from the brainstem shall be identified. Then, the flocculus of the cerebellum is identified and smoothly retracted in a small amount. In most cases, the choroid plexus of the IV. ventricle is then exposed.

The arachnoid membrane covering these structures is then gently dissected. The preparation should be directed towards the space between plexus and brainstem.

All vessels supplying the brainstem must be preserved. Thereafter, a 45° inclined dissector can be inserted into the lateral recess. This can be facilitated by additional retraction of the cerebellar hemisphere. Care must be taken that there are no vessels inside the recess, running in cranio-caudal direction. Such vessels may be injured or impede implantation by narrowing or occluding the entrance to the IV ventricle.
STEP 7: EABR Recording

Prior to implantation of the Implant SYNCHRONY/SYNCHRONY PIN ABI, EABR measurements should be performed. Connect the ABI Connector Cable and the ABI Placing Electrode via the Micro-D Plug (also refer to III Set-up and recommended measurement for EABR recording). The transition of the ABI Connector Cable to the ABI Placing Electrode shall not be brought into contact with the surgery wound. Connect the ABI Connector Cable, located outside the patient environment, to the plug of the ABI Stimulator Box (used for ABI Placing Electrode only).

Prior to placement of the ABI Placing Electrode, the anatomical facts need to be established. The positioning of the ABI Placing Electrode shall be done in accordance with anatomical landmarks. The lateral recess is opened 4 mm and the ABI Placing Electrode is inserted with the recommended tools. The ABI Placing Electrode has the same shape and dimensions as the SYNCHRONY/SYNCHRONY PIN ABI electrode with a reduced number of contacts and no polyester mesh. Care shall be taken regarding the electrode orientation, the contacts shall face the brainstem surface.

By stimulating the brainstem with bipolar, biphasic current pulses, EABR potentials shall be assessed. If EABRs become identifiable, an ABI can be used. If no EABRs are measurable, a repositioning of the ABI Placing Electrode may be necessary. Again, EABRs shall be assessed. Placement of the SYNCHRONY/SYNCHRONY PIN ABI electrode in the absence of clear EABRs may only be done if the surgeon considers the anatomical landmarks to be sufficient and reliable.
STEP 8: Drilling the Implant Bed

STEP 8-A: SYNCHRONY

**CAUTION**
The implant must be immobilized in a flat stimulator bed drilled in the temporal bone. The electrode lead should be placed in a ramp-like bony channel without sharp edges to protect it against postoperative movement and excessive mechanical impact.

Care shall be taken that the platinum bridge and electrode leads are not compressed by sutures or other structures, i.e. bony rims.

The implant must be immobilized in a flat stimulator bed drilled in the temporal bone. In adults, it may not be necessary to expose the dura, but in small children with a thin skull, drilling to the dura may sometimes be required in order to ensure that the stimulator is well recessed in its bed. If drilling down to the dura is necessary a bony island should remain. Ideally, the stimulator is recessed approximately 2 mm.

Once again the Mi1200 Implant Template can be used to mark the flatness on the skull and the correct position for the implant bed (see Figure 33).

If, for example, the implant is fixed with sutures, a diamond burr should be used to drill the holes so that the implant can be immobilised later. The suture holes should be drilled such that the sutures do not cross the electrode, but rather only cross the silicone over-mold (see Figure 34).
STEP 8-B: SYNCHRONY PIN

**CAUTION**

- The area of the temporal bone on which the stimulator will be placed, shall be flattened in order to ensure that the implant is sufficiently immobilized.
- The two pins of the SYNCHRONY PIN ABI Cochlear Implant should be recessed into the skull to a depth of 1.5 mm.
- The electrode lead should be placed in a ramp-like bony channel without sharp edges to protect it against postoperative movement and excessive mechanical impact.
- All sharp edges of bone must be removed to avoid possible damage to the electrode lead. Drilling should be completed before the cochlea is opened to prevent any bone dust from entering.
- Protect the middle ear cavity from bone dust contamination by closing it with medical gauze.

The Mi1200 Implant Template, PIN ABI can be used to mark the correct position of the stimulator and the electrode channel (see Figure 35).

The temporal bone area, in which the stimulator is placed, must be flattened to ensure sufficient stimulator immobilisation (see Figure 36). The flattening also ensures a flat stimulator position without a later rocking of the implant and enables a good positioning of the pins in the drilled holes. The immobilisation of the SYNCHRONY PIN ABI Cochlear Implant with the two pins stabilizes the stimulator against translational and rotational motion.

Furthermore, the electrode lead should be protected in a ramp-like bony channel without sharp edges which is drilled into the skull. It is important to ensure that the channel is deep and wide enough to comfortably accommodate the electrode. This protects the electrode lead against postoperative movement and excessive mechanical impact.
The two pins of the SYNCHRONY PIN ABI Cochlear Implant should be recessed into the skull with the PIN Drill Guide SI or the Mi1000 PIN ABI Drill Guide to a depth of 1.5 mm (see Figure 37).

For drilling the holes into the flat area, the following methods can be used:
PIN Drill Guide SI and a diamond burr of 1.0 mm

CAUTION

- Ensure that the dura is not inadvertently damaged when drilling the holes.
- Always use a slow turning drill, e.g. 2000 rpm.
- Stop drilling when a depth of 1.5 mm is achieved.

Use the diamond burr of 1.0 mm to mark both hole positions and drill through the PIN Drill Guide SI. Avoid recessing the pins deeper than 1.5 mm (see Figure 38).

PIN Drill Guide SI and a surgical pen (surgical ink)

CAUTION

- Ensure that the dura is not inadvertently damaged when drilling the holes.
- Use a diamond burr of 1.0 mm for drilling the holes.
- Always use a slow turning drill, e.g. 2000 rpm.
- Stop drilling when a depth of 1.5 mm is achieved.

When using a surgical pen or surgical ink for marking the position of the holes through the PIN Drill Guide SI, be aware that the irrigation of the drill can wash your markings away. Drying the bone with a sterile tissue helps to better mark the holes with the surgical pen or ink. Use the diamond burr of 1.0 mm to mark both hole positions. Then, drill the full depth of the pin hole. Avoid recessing the pins deeper than 1.5 mm.
STEP 9: Immobilize the Implant

STEP 9-A: SYNCHRONY

CAUTION

• If monopolar diathermy has been used, it must now be disconnected. High current levels may cause damage to the implant. If bipolar cautery must be used, the tips of the cautery should be kept at least 3 cm away from the implant.
• Additional immobilisation of the implant needs to be performed.
• If sutures are chosen for immobilisation of the implant do not place the sutures directly over the electrode lead.

Additional immobilisation of the implant needs to be performed (e.g. with sutures). It should be conducted in such a way that there will be no postoperative movement. Continuous movement may result in mechanical fatigue and subsequent premature failure of electrical connections.

When the implant is immobilised with sutures, the holes drilled in STEP 8 should be used to secure the implant in its bed and the electrode should be placed into the drilled channel leading into the mastoid. Make sure the electrode channel is deep enough to prevent the tie-down from exerting pressure and damaging the electrode. MED-EL recommends the use of the following techniques (details in Figure 40, Figure 41).
STEP 9-B: SYNCHRONY PIN

CAUTION

• If monopolar diathermy has been used, it must now be disconnected. High current levels may cause damage to the implant. If bipolar cautery must be used, the tips of the cautery should be kept at least 3 cm away from the implant.
• Recessing the pins and efficient immobilisation of the stimulator is important to prevent postoperative movement.
• Do not place the sutures directly over the electrode lead.

The area of the temporal bone on which the stimulator will be placed, shall be flattened in order to ensure that the implant is sufficiently immobilized. The two pins of the SYNCHRONY PIN Cochlear implant should be recessed into the skull with the PIN Drill Guide SI to a depth of 1.5mm. The electrode lead should be placed in a ramp-like bony channel without sharp edges to protect it against postoperative movement and excessive mechanical impact.

The two pins give additional stability against translational and rotational motion. Recessing the pins and efficient immobilization of the stimulator (e.g. with sutures) is important to prevent postoperative movement. Continuous movement may result in mechanical fatigue and subsequent premature failure of electrical connections.

When the implant is immobilised with sutures, holes drilled into the bone should be used to secure the implant in its flat area and the electrode should be placed into the drilled channel leading into the mastoid. Ensure that the electrode channel is deep enough to prevent the tie-down from exerting pressure and damaging the electrode. MED-EL recommends the use of different immobilization techniques (details on Figure 42, Figure 43).
STEP 10 Intra-operative Recordings

At this stage intra-operative recordings like Impedance Field Telemetry (IFT) and Electrically Evoked Brainstem Response (EABR) can be performed.

Intra-operative measurements are performed with the appropriate MED-EL application software and the MED-EL hardware interface system. For details please refer to the applicable User Manual. It is not possible to sterilise any component of the MED-EL clinical interface system. When used in a sterile environment, the coil and cable should be covered with sterile material (i.e. “sterile sleeve”).

Since the coil should not be placed directly on the implant, either sterile gauze drenched in saline solution or the skin flap should be placed between the coil and the implant. Moistening the underside of the skin flap with sterile saline or pooling saline over the ground electrode of the implant prior to performing intra-operative recordings may improve readings.

IFT (Impedance Field Telemetry)

After the implant is in place, a telemetry check allows:

- individual electrode impedance measurements
- verification of the absence of short and open circuits between electrodes
- determination of voltage distribution across the cochlear nucleus

As with any telemetry system, intra-operative impedance testing may not provide an accurate representation of later electrode function. “High” values observed intra-operatively may be caused by air bubbles on the electrode contact surface. These generally dissipate within a few hours or days after surgery.

STEP 11: Implantation Procedure with EABRs

After fixation of the stimulator, the ABI Active Electrode shall be placed. The polyester mesh which is embedded in the silicone of the ABI electrode array is cut to fit in the lateral recess, and the ABI electrode array is placed into the estimated correct location. Go on with the procedure in STEP 13: EABR Measurements during Stimulation via the Implant SYNCHRONY/SYNCHRONY PIN ABI.

EABR (Electrically Evoked Brainstem Response)

For details regarding EABR please refer to the section III Set-up and recommended measurement for EABR recording.
STEP 12: Implantation Procedure in Case of Absence of EABRs

To determine the best stimulation area, the ABI Placing Electrode is placed into the lateral recess again. EABRs are recorded by stimulating the CN in any bipolar configuration mode possible with the four contacts. EABR measurements are repeated until the stimulation area has been found where all electrode combinations elicit EABRs. If no EABRs can be recorded, the surgeon has to decide if the anatomical landmarks are sufficiently reliable to proceed with the implantation.

The ABI Placing Electrode is then retracted and substituted by the ABI Active Electrode. The polyester mesh which is embedded in the array is cut to fit into the lateral recess, and the SYNCHRONY/SYNCHRONY PIN ABI electrode array is placed in the location where the ABI Placing Electrode evoked EABRs. Therefore, the insertion depth into the lateral recess shall be the same for both electrodes.

STEP 13: EABR Measurements during stimulation via the SYNCHRONY/SYNCHRONY PIN ABI

Before fixation of the ABI Active Electrode, a final EABR check via the implant is performed. The appropriate coil of the MED-EL hardware interface system is put into a sterile sleeve and placed on the implant. For correct recording of potentials, the MED-EL hardware interface system shall trigger the measuring device (see III Set-Up and Recommended Measurement for EABR Recording).

This measurement via the implant should be used as a final check. The results from the ABI Placing Electrode are primarily used for the determination of the stimulation site.

STEP 14: Fixation of the Electrode

CAUTION

Improper fixation or placement of the ABI electrode array may result in dislocation. This is true especially when a large lateral recess is present.

The ABI Active Electrode can be prefixed onto the brainstem surface with a piece of hemostytic material and fibrine glue to prevent postsurgical displacement. The main fixation is achieved by gluing the electrode lead to the rostral surface of the cerebellum. Useful for this procedure is a collagen foam covered with instant glue (TachoComp™).
STEP 15: Removal of the Platinium Bridge on the Implant Housing

The platinum bridge (Figure 44) on the implant housing shall be removed with forceps. The platinum bridge is fixed with silicone only on its edges. This allows the lifting of the platinum bridge (Figure 45) from the surface of the implant. Pull out the whole platinum bridge (Figure 46 and Figure 47). Make sure that there is no residual platinum foil on the implant housing (Figure 47).

STEP 16: Closing the wound

After implantation, the dura shall be closed in a watertight fashion. The entry of the electrode lead into the dura is sealed with fibrin glue or TachoComp™. The housing is covered in a double layer technique by the periosteum and the galea flap, which has in contrast to the normal skin incision in acoustic neurinoma surgery a mirrored question mark shape on the left and a regular question mark shape on the right side. The wound should be closed in layers. No drainage is used. The area of the wound is covered with a compress and sterile gauze applying even pressure.

Remark for initial fitting

CAUTION

The initial fitting of the audio processor is done typically six to eight weeks after surgery. The fitting shall be performed in the intensive care unit of the hospital or in the presence of an intensive care crash cart.
MRI Caution

Evidence has been provided for this implant type to pose no known hazard in specified MRI environments (without surgical removal of the internal magnet) when adhering to the conditions and Safety Guidelines listed below. The implant has a specially designed magnet which allows safe MRI scanning with the magnet in place, and there is no need to remove the implant magnet regardless of the scanner field strength. The implant magnet can be surgically removed if needed to avoid imaging artifacts. The physician/MRI operator should always be informed that a patient is a auditory brainstem implant user and that special safety guidelines have to be followed:

MRI scanning is possible in consideration of the Safety Guidelines if the following conditions are fulfilled:

- MRI scanners with static magnetic fields of 0.2 T, 1.0 T, or 1.5 T only. No other field strengths are allowed. When using other field strengths, injury to the patient and/or damage to the implant are possible.
- In case of additional implants, e.g. a hearing implant in the other ear: MRI safety guidelines for this additional implant need to be met as well.

Safety Guidelines:
- Before patients enter any MRI room all external components of the implant system (audio processor and accessories) must be removed from the head. For field strengths of 1.0 T and 1.5 T, a supportive head bandage must be placed over the implant. A supportive head bandage may be an elastic bandage wrapped tightly around the head at least three times (refer to Figure 48). The bandage shall fit tightly but should not cause pain. Performing an MRI without head bandage could result in pain in the implant area and in worst case can lead to migration of the implant and/or dislocation of the implant magnet.
- Head orientation: In case of 1.0 T and 1.5 T MRI systems, straight head orientation is required. The patient should not incline his/her head to the side; otherwise torque is exerted onto the implant magnet which could cause pain. In case of 0.2 T scanners, no specific head orientation is required.
- Sequences in “Normal Operating Mode” shall be used only.
- During the scan patients might perceive auditory sensations such as clicking or beeping. Adequate counseling of the patient is advised prior to performing the MRI. The likelihood and intensity of auditory sensations can be reduced by selecting sequences with lower specific absorption rate (SAR) and slower gradient slew rates.
- The magnet can be removed by pushing on the top side of the magnet so that it comes out at the bottom side of the implant to reduce image artifacts. If the magnet is not removed, image artifacts are to be expected (refer to Figure 49).
- The exchange of the magnets with the Non-Magnetic Spacer and vice versa has been tested for at least five repetitions.
The above instructions should also be followed if areas of the body other than the head are to be examined (e.g. knee, etc.). When lower extremities are to be examined, it is recommended that the patient’s legs are positioned in the scanner first.

If the conditions for MRI safety and the Safety Guidelines are not followed, injury to the patient and/or damage to the implant may result!

Figure 49 Image artifacts arising in a 1.5 T scanner. The upper picture shows the artifacts obtained with the implant magnet in place whereas the lower picture illustrates the image artifacts when the implant magnet is replaced with the Non-Magnetic Spacer.
Magnet Removal Procedure

The following instruments are required for the Magnet Removal Procedure:

**Magnet Replacement Kit**
Order number: 09693
Consisting of:

- **Non-Magnetic Spacer Ms010107**
  The Non-Magnetic Spacer (see Figure 50) is intended to be used as placeholder for the regular implant magnet of the Mi1200 Hearing Implant during MRI procedures, when a reduced image artifact is desirable.

- **Replacement Magnet Ms010108**
  The Replacement Magnet (see Figure 51) is intended to be used after an MRI, as replacement of the original implant magnet of the Mi1200 Hearing Implant and to restore full functionality of the Mi1200 Hearing Implant.

**Magnet Tool Kit**
Order number: 09734
Consisting of:

- **Magnet Removal Tool Ms050206**
- **Magnet Insertion Tool Ms050205**

The Magnet Removal Tool (see Figure 52) is for removal of the MED-EL removable implant magnet and the Non-Magnetic Spacer.

The Magnet Insertion Tool (see Figure 53) is for insertion of the Non-Magnetic Spacer and the Replacement Magnet.

The instruments are made of surgical grade stainless steel. The devices are delivered non-sterile.
Surgical Procedure

STEP 1: Opening the skin flap
When opening the skin flap, keep an adequate distance between the incision and the coil. This will prevent damage to the implant under the skin. For marking the incision either the patient’s audio processor coil or the MAX Coil S can be used. When used in a sterile environment, the Coil should be covered with sterile material (i.e. “sterile sleeve”). MED-EL recommends a distance of 5 to 15 mm from the coil and an opening angle between 160° and 200°. Carefully dissect the fibrous tissue to locate the coil part of the implant and expose the magnet. The wound should be opened in layers.

STEP 2: Removing the Implant Magnet or Non-Magnetic Spacer

CAUTION To avoid movement of the implant it is recommended to fix the stimulator by pressing it against the bone with one hand.
1. Place the Magnet Removal Tool in front of the implant coil.

2. Lift the coil part of the implant by sliding the tip of the Magnet Removal Tool under the implant coil.

3. Centre the implant coil in the tip part of the Magnet Removal Tool.

4. Push the Implant Magnet or Non-Magnetic Spacer out of the implant coil by pressing together the two handles of the Magnet Removal Tool.

5. MED-EL recommends checking that the two handles of the Magnet Removal Tool are completely re-opened before pulling out the instrument.

6. Remove the Magnet Removal Tool by slowly pulling out the instrument from the implant coil.

7. After pulling out the instrument, the Implant Magnet or Non-Magnetic Spacer can be removed from the tip of the Magnet Removal Tool by lifting the upper handle. The removed Implant Magnet or Non-Magnetic Spacer can be disposed of.
STEP 3: Inserting the Non-Magnetic Spacer or Replacement Magnet

CAUTION
To avoid movement of the implant it is recommended to fix the stimulator by pressing it against the bone with one hand.

1. Open the upper handle of the Magnet Insertion Tool by unlocking the small locking mechanism and lifting the counter blade.

2. Place the Non-Magnetic Spacer or Replacement Magnet in the front part of the Magnet Insertion Tool. The Non-Magnetic Spacer or Replacement Magnet is correctly placed into the tip when the serial number labelling is not readable from the top.

3. Close the counter blade and lock the locking mechanism.

4. Place the Magnet Insertion Tool in front of the implant coil.

5. Lift the coil part of the implant by sliding the tip of the Magnet Insertion Tool under the implant coil.
6. Centre the implant coil in the tip part of the Magnet Insertion Tool so the Non-Magnetic Spacer or Replacement Magnet is completely visible through the hole in the implant coil.

7. For complete insertion of the Non-Magnetic Spacer or Replacement Magnet into the implant coil, insert the Non-Magnetic Spacer or Replacement Magnet into the implant coil by pressing the two handles of the instrument together until the two handles are touching.

8. Re-open the two handles of the Magnet Insertion Tool.

9. MED-EL recommends checking that the two blades of the Magnet Insertion Tool are completely re-opened before pulling out the instrument.

10. Remove the Magnet Insertion Tool by slowly pulling out the instrument from the implant coil.

11. Check the correct magnet position.
STEP 4: Close wound

Before closing the wound visually confirm that the blue Replacement Magnet (Ms010108) or the purple Non-Magnetic Spacer (Ms010107) was inserted as appropriate. When closing the wound, care should be taken not to damage the implant. The wound should be cleaned and closed in layers with staples or absorbable subcutaneous sutures. The area of the wound should then be covered with a compress and sterile gauze, and even pressure should be applied.

X-rays

The SYNCHRONY/SYNCHRONY PIN ABI can be identified by x-ray post surgery. Right is an example for the devices.

Explanting the Device

- The implant may become non-functional, either by accident or due to medical or technical reasons. In this case, it is strongly recommended to replace the device.
- If for any reason the device is not used anymore, it is strongly recommended to explant the device. If an explantation is not performed, functional checks of the implant on a regular basis are strongly recommended.
- If possible, the device should be removed without damaging or cutting it. Damage to the device during or after explantation may prevent or reduce the manufacturer’s ability to determine the root cause of failure.
- Staff should follow common universal precautions and handle the explanted device as potentially contaminated biohazardous material.
- After explantation, the implant should be appropriately cleaned and disinfected. During cleaning, extraneous tissue should be removed, but only to such an extent that damage to the implant is not risked.

- An explanted device should be placed in a leak-proof, disinfected (or sterile) container filled with saline and returned to MED-EL Headquarters. The device should be accompanied by written information including the reason for explantation.
MED-EL Surgical Videos

MED-EL Surgical (SONATA, PULSAR)
Order number: MKT 20265

EAS Surgical Video
Order number: MKT 20542

Small Incision Surgical Video
Order number: MKT 21627

Frankfurt, Germany Live Surgery DVD*
Order number: MKT 22987

MED-EL Advanced Surgical Techniques
3D BluRay English
Order Number: MKT 23518

For additional videos please visit the MED-EL Professional Webpage (www.medel.com/professionals)

Literature


* Note that the featured surgery is performed in Germany and the techniques / variants used may differ from other markets.
The MED‑EL SYNCHRONY/SYNCHRONY PIN Auditory Brainstem Implant System serves to restore some sense of hearing by electrical stimulation of the cochlear nucleus (CN). It is the result of many years of research and is intended for use in patients with non-functional cochlear nerves.

This Surgical Guideline describes proper techniques for implanting the Mi1200 SYNCHRONY/SYNCHRONY PIN Auditory Brainstem Implant (ABI) (hereafter referred to as the SYNCHRONY/SYNCHRONY PIN ABI). It serves as additional information for professionals and should not be used as an "Instructions for Use".

MED‑EL implants are manufactured to the highest quality standards in order to ensure long term reliability. All materials used in the implant have been rigorously tested for biocompatibility, durability and reliability. MED‑EL applies a quality management system that meets all EN ISO 13485:2003 requirements and complies with US Quality System Regulations and Canadian Medical Device regulations (CAN/CSA ISO 13485‑2003). Components of the MED‑EL Cochlear Implant System meet the requirements for AIMD 90/385/EEC and MDD 93/42/EEC.

The information in this Surgical Guideline is believed to be true and correct, however, some specifications may be subject to change without notice.

For country specific information please see the applicable "Instruction for Use" delivered with the implant system.