**Title:** DEVICE FOR CRANIAL IMPLANTATION, AND SYSTEM AND USE THEREOF

**Abstract:** A device for cranial implantation, comprising a ferrule and a plate for placement in the ferrule is provided. Furthermore, a neurological implant system comprising a probe and a device for cranial implantation is provided.
Device for cranial implantation, and system and use thereof

FIELD OF THE INVENTION

The present invention pertains to a device for cranial implantation. More specifically, the present invention pertains to a device for cranial implantation comprising a ferrule with a first hole.

BACKGROUND OF THE INVENTION

Within the field of neurotechnology, deep brain stimulation (DBS) is a surgical treatment involving the implantation of a medical device called a deep-brain stimulator, which sends electrical impulses to specific parts of the brain. DBS in certain brain regions has provided remarkable therapeutic benefits for otherwise treatment-resistant disorders such as chronic pain, Parkinson's disease, tremor and dystonia. Despite the long history of DBS, its underlying principles and mechanisms are still not clear. DBS directly changes brain activity in a controlled manner. Unlike lesioning techniques, its effects are reversible. Furthermore, DBS is one of only a few neurosurgical methods that allow blinded studies.

In principle, the deep brain stimulation system comprises two components: the implanted pulse generator (IPG), and the probe. The IPG is a battery-powered neurostimulator that sends electrical pulses to the brain to interfere with neural activity at the target site. The IPG is typically encased in e.g. a titanium housing. The probe consists of about 10-40 cm long wires and a plurality of electrodes. The wires connect the IPG to the electrodes, which are located at the distal end of the probe. The IPG may be calibrated by a neurologist, nurse or trained technician to optimize symptom suppression and control side effects.

DBS probes are placed in the brain according to the type of symptoms to be addressed. All components are surgically implanted inside the body. The typical procedure is performed under local anaesthesia, where a hole is drilled in the skull and the electrode is inserted with feedback from the patient for optimal placement. The right side of the brain is stimulated to address symptoms on the left side of the body and vice versa.
Commercially available DBS systems consist of a chest-implanted IPG, a connector cable of approximately 30 cm running subcutaneously from the IPG to the top of the patient's head, and a lead cable connected proximally to the connector cable by means of a connector, which may be approximately 20 mm long and 4-5 mm in diameter. The lead cable carries 4 electrodes distally, and is itself rather long, around 30 cm, so that a lot of excess length has to be left under the patient's scalp. This system has several disadvantages:

- the fixation of the lead cable is difficult and it is liable to lead displacement and inaccuracies;
- the long cables cause irritation and erosion of the skin;
- the cables are the source of lead break;
- infection can propagate subcutaneously along the cable into the brain; and
- the long and uncontrolled cable bundles give rise to potentially dangerous voltages when submitted to the electromagnetic fields of a Magnetic Resonance Imaging scanner (MRI).

US 2005/0,228,249 proposes solutions to the afore-mentioned by providing an IPG mounted in a ferrule, intracranially after a craniotomy is performed in the parietal bone, with the IPG device mounted into the ferrule, and further connected to seed electrodes by means of a seed electrode interface that can have the shape of a burrhole cover, or be simply a subcutaneous module.

However, this solution still needs cables running subcutaneously, to cranial positions, possibly quite remote from the parietal position of the ferrule-mounted IPG. This may give rise to unwanted motion of the wire, which lead to mechanical forces on the probe and thus potential damage to brain tissue. Also, the probe wire must have extra length to allow manipulation of the implantable device before the implantation. When this excess wire is stored subcutaneously after placement of the probe, it may lead to irritation, skin erosion, and potentially infection.

Hence, an improved way of mounting a DBS probe, allowing for increased flexibility, cost-effectiveness, safety and user friendliness would be advantageous.

SUMMARY OF THE INVENTION

Accordingly, the present invention preferably seeks to mitigate, alleviate or eliminate one or more of the above-identified deficiencies in the art and disadvantages singly or in any combination and solves at least one of the above mentioned problems e.g. by providing a cranial implant device.
In an aspect, a device for cranial implantation allowing for adjustment of positioning of a probe is provided. The device comprises a ferrule forming a first hole there through. The device also comprises a plate for placement in the ferrule. The plate is provided with a second hole, such that when the plate is placed in the ferrule, the first hole and the second hole at least partly coincides enabling accommodation of the probe. This provides for the advantage that the probe does not have to be placed in the middle of the first hole, but instead the positioning of the probe is adaptable, which is especially useful since the surgeon usually does not know exactly where the probe is to be placed prior to the surgery. The device provides for increased flexibility, cost-effectiveness, safety and user friendliness.

Moreover, in use the device provides for less discomfort for the patient.

In an aspect, a neurological implant system is provided. The system comprises a probe and the device for cranial implantation is disclosed providing for less risk of potentially infection.

In another aspect, use of the device for cranial implantation for adjustment of positioning of a deep brain stimulation probe is provided.

Other embodiments and advantages will be disclosed in the following.

BRIEF DESCRIPTION OF THE DRAWINGS

These and other aspects, features and advantages of which the invention is capable of will be apparent and elucidated from the following description of embodiments of the present invention, reference being made to the accompanying drawings, in which;

Fig. 1 is a sectional view of a device according to an embodiment;
Fig. 2 is a top view of a device according to an embodiment;
Fig. 3A is a bottom view of the device according to an embodiment;
Fig. 3B is a top view of a device according to an embodiment; and
Fig. 4 is a sectional view of a device according to an embodiment.

DESCRIPTION FOR DETAILED EMBODIMENTS

Several embodiments of the present invention will be described in more detail below with reference to the accompanying drawings in order for those skilled in the art to be able to carry out the invention. The invention may, however, be embodied in many different forms and should not be construed as limited to the embodiments set forth herein. Rather, these embodiments are provided so that this disclosure will be thorough and complete, and will fully convey the scope of the invention to those skilled in the art. The embodiments do
not limit the invention, but the invention is only limited by the appended patent claims. Furthermore, the terminology used in the detailed description of the particular embodiments illustrated in the accompanying drawings is not intended to be limiting of the invention.

In an embodiment according to Fig. 1, a device 10 for cranial implantation is provided. The device 10 for cranial implantation is allowing for adjustment of positioning of a probe 16 comprises a ferrule 11, forming a hole there through, and a plate 12 for placement in the ferrule. The plate 12 is being provided with a hole, such that when the plate 12 is placed in the ferrule 11, the hole of the ferrule and the hole of the plate at least partly coincides enabling accommodation of the probe 16.

In use the ferrule 11 is positioned in a hole provided in the cranium 13 of a patient. The hole of the ferrule 11 is provided such that it faces towards the brain. The hole provided through the ferrule may have a cylindrical shape. The ferrule 11 has an edge 14, in use being located on the side facing away from the brain. The edge 14 is positioned to be in contact with the cranium 13 to stop the ferrule from moving towards the brain. The plate 12 has a hole positioned in such a way that the hole of the ferrule and the hole of the plate at least partly coincides enabling accommodation of the probe 16. In this way, a path enabled by the hole of the ferrule and the hole of the plate is created, through which a probe 16 may be inserted in the brain through the ferrule 11 and the plate 12.

In one embodiment, the plate 12 is configured with an outer structure enabling rotatable placement in the ferrule 11. In this embodiment, the ferrule 11 has an inverted, or corresponding, structure for receiving the outer structure of the plate 12. The outer structure of the plate 12 may e.g. comprise protrusions 23, such as cogs, which may be received by a corresponding structure 24 of the ferrule 11. When the hole is non-concentrically provided in the plate 12, by rotating of the plate 12 in relation to the ferrule 11, the position of the plate hole will change in relation to the ferrule 11. Accordingly, the position of the path being rendered by the coinciding plate hole and ferrule hole will also change by rotation of the plate 12 in relation to the ferrule 11.

In an embodiment, according to Fig. 2, a device 20 for cranial implantation is provided, wherein after placement of the plate 22 in the ferrule 21, due to the configuration of the structure of the plate 22, in Fig. 2 shown as cogs, further rotation of the plate 22 in relation to the ferrule 21 will be prevented. In Fig. 2 the plate is placed above the ferrule. The hole of the ferrule is indicated as a dashed line, indicating that the ferrule hole is located beneath the plate. An advantage with this is that the plate 22, and thus the position of the probe, by means of the hole 30 provided in the plate, is stabilized when the plate 22 has been
positioned in the ferrule 21. This means that the probe will be prevented from sliding in use. However, adjustment of the position of the probe may still be possible by lifting the plate 22 from the ferrule 21, and rotate the plate in relation to the ferrule, in order to change the position of the hole 30.

In another embodiment, the plate 12 is configured with an outer structure enabling rotation in relation to the ferrule 11 while placed in the ferrule 11. In this embodiment the outer structure of the plate 12 may be a smooth surface, e.g. enabling a mutual sliding motion against the ferrule 11.

An advantage rendered by the above-mentioned embodiments is that the probe does not have to be placed in the middle of the hole of the ferrule 11, since the positioning of the probe is determined by the placement of the hole of the plate 12, the plate's rotation in relation to the ferrule 11, and/or the location at which the hole of the ferrule 11 and the hole of the plate 12 coincides. This is especially useful since the surgeon usually does not know exactly where the probe is to be placed from the beginning of the surgery. In this way the positioning of the path enabling insertion of the probe is made adaptable.

The ferrule 11 may be mounted in the hole in the cranium 13 and fixed to the cranium 13 with methods well known to a person skilled in the art, such as glue, bio cement or thread.

After the ferrule 11 has been placed in the cranium, the brain is still accessible and visible to the physician. The plate 12 is then positioned over the probe 16. The probe 16 may then be placed in the brain after an extra electro-physiological analysis of the target tissue. After the probe 16 has been correctly positioned in the brain, the probe 16 is mechanically fixated by positioning the plate 12 so that it may be fixated to the ferrule, e.g. by glue. The ferrule is thus completely closed.

In Fig. 3, the positioning of the plate according to an embodiment is visualized. Fig. 3A is showing the ferrule 11 from inside the cranium. The second hole 30 in the plate 12 may be positioned anywhere in the plate 12, if it is placed on a suitable distance from the centre of the plate 12. The ferrule 11 may thus have a circular shape with a concentric circular first hole. The plate 12 may have a circular shape with an eccentric circular hole. If slided and turned correctly, the plate 12 will then always cover the first hole in the ferrule 1. However, the plate 12 may also be placed eccentric with respect to the ferrule 11.

However, the shape of the ferrule 11, plate 12, the first hole and the second hole 30 may be in any form such as elliptic, rectangular, triangular etc.
In an embodiment, the first hole in the plate 12 is a slot. This increases flexibility, since the probe location may be moved along the length of the slot and thereby adjusted. The slot space, which is not occupied by the probe, may then be closed with glue or bio cement.

The suitable distance may be calculated with the following formulas:

\[
\begin{align*}
D3+D4+2*E &= 2*D2+1 \\
D3-2*E &= D2+1 \\
D1-D2-D3+D4+2*E &= 0
\end{align*}
\]

Wherein D1 is the inner diameter of the ferrule, D2 is the diameter of the first hole, D3 is the diameter of the plate 12 and D4 is the diameter of the hole 30 in the plate 12. E is the eccentricity, or the distance between the centre of the plate to the centre of the hole 30 in the plate 12.

In an embodiment, where the outer diameter of the ferrule is 20 mm, D1 is 19.2 mm, D2 is 9.7 mm, D3 is 15 mm, and D4 is 1.27 mm, E will be 2.1 mm.

Fig. 3B is showing the ferrule 11 from outside the cranium. After the plate 12 has been fixated to the ferrule 11, the probe 16 may also be fixated to the plate, e.g. with glue.

In an embodiment according to Fig. 4, a device for cranial implantation 40 is provided. The device allowing for adjustment of positioning of a probe 16 comprises a ferrule 11, forming a first hole there through, and a plate 12 for placement in the ferrule. The plate 12 is being provided with a second hole, such that when the plate 12 is placed in the ferrule 11, the first hole and the second hole at least partly coincides enabling accommodation of the probe 16. Furthermore, the device for cranial implantation 40 comprises a lid 41 for fixing the lid 41 to the ferrule 11, thus closing the ferrule, e.g. by glue or bio cement. When the device for cranial implantation 40 is closed with the lid 41, the scalp 15 may be closed. In an embodiment, the device for cranial implantation 40 comprises electronics 42 and a power source 43, such as a battery, for driving the electronics 42. The electronics 42 may be connected to the probe 16. The probe 16 may be curled up inside the ferrule 11 so that the excess probe length may be stored in the ferrule 11. Since the curling of the excess length takes place after the plate 12 has fixated the probe 16, the brain is already protected from these mechanical actions.
In an embodiment, the electronics comprised in the device for cranial implantation comprises a battery, a pulse generator, a multiplexer, a board such as a printed circuit board or a flexible board, and a connector. However, the electronics may be any kind of electronics for regulating the probe, known to a person skilled in the art. Since the probe 16, electronics 42 and power source 43 is collected in the ferrule 11 and sealed on one side, towards the brain, with the plate 12 and on the other side, towards the scalp 15, with the lid 41, the device for cranial implantation 40 is safer from infection compared to prior art methods. Furthermore, the risk of displacement of parts is lower, as well as the risk of wear and tear.

In an embodiment, the ferrule 11, the lid 41 and the plate 12 are made of titanium. However, other materials may be possible, as will be appreciated by a person skilled in the art.

In an embodiment, a neurological implant system comprising a probe 16 and a device 10 for cranial implantation is provided.

In an embodiment, use of the device 10 for cranial implantation for adjustment of positioning of a deep brain stimulation probe is provided.

Although the present invention has been described above with reference to specific embodiments, it is not intended to be limited to the specific form set forth herein. Rather, the invention is limited only by the accompanying claims and, other embodiments than the specific above are equally possible within the scope of these appended claims.

In the claims, the term "comprises/comprising" does not exclude the presence of other elements or steps. Furthermore, although individual features may be included in different claims, these may possibly advantageously be combined, and the inclusion in different claims does not imply that a combination of features is not feasible and/or advantageous. In addition, singular references do not exclude a plurality. The terms "a", "an", "first", "second" etc do not preclude a plurality. Reference signs in the claims are provided merely as a clarifying example and shall not be construed as limiting the scope of the claims in any way.
CLAIMS:

1. A device (10) for cranial implantation allowing for adjustment of positioning of a probe, comprising
   - a ferrule (11, 21) forming a first hole there through,
   - a plate (12, 22) for placement in the ferrule (11, 21), said plate (12, 22) being provided with a second hole, such that said first hole and said second hole at least partly coincide enabling accommodation of said probe when said plate (12, 22) is placed in said ferrule (11, 21).

2. The device (20) according to claim 1, wherein said plate (22) is configured with an outer structure comprising protrusions (23), and the ferrule (21) is configured with a structure (24) for receiving the outer structure of the plate (22), such that when said plate (22) is placed in said ferrule (21) the plate (22) is prevented from rotating in relation to said ferrule (21).

3. The device (10) according to claim 1, wherein said plate (12) is configured with an outer structure with a smooth surface, and the ferrule (11) is configured with a structure for receiving the outer structure of the plate (12), such that when said plate (12) is placed in said ferrule (11), said plate is slidably rotatable in relation to said ferrule (11).

4. The device (10) according to claim 1, wherein the second hole is circular.

5. The device (10) according to claim 1, wherein the second hole is a slot.

6. The device (10, 20) according to any of claims 1 to 3, wherein the ferrule (11, 21) is fixed to the cranium (13).

7. The device (40) according to claim 1, comprising electronics (42) and a power source (43) inside the ferrule and a lid (41) for closing the ferrule towards the scalp.
8. The device (40) according to claim 7, wherein the lid (41) is fixed to the ferrule (11).

9. The device (40) for cranial implantation according to claim 7, wherein the ferrule (11), the plate (12) and the lid (41) are made by Titanium.

10. A neurological implant system comprising a probe (16) and a device (10, 20, 30) for cranial implantation according to claim 1.

11. The neurological implant system according to claim 10, wherein the probe (16) is a deep brain stimulation probe.

12. Use of a device (10, 20, 30) for cranial implantation according to claim 1, for adjustment of positioning of a deep brain stimulation probe.
# INTERNATIONAL SEARCH REPORT

**A. CLASSIFICATION OF SUBJECT MATTER**

**INV. A61N1/05 A61B5/03**

According to International Patent Classification (IPC) or to both national classification and IPC.

**B. FIELDS SEARCHED**

Minimum documentation searched (classification system followed by classification symbols)

A61N A61B

Documentation searched other than minimum documentation to the extent that such documents are included in the fields searched

Electronic data base consulted during the international search (name of data base and, where practical, search terms used)

EPO-Internal

**C. DOCUMENTS CONSIDERED TO BE RELEVANT**

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<td>column 5, lines 28-30,43-46 figures 1-5</td>
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Further documents are listed in the continuation of Box C

See patent family annex

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Name and mailing address of the ISA:

European Patent Office, P B 5818 Patentlaan 2 NL - 2280 HV Rijswijk

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Authorized officer: Worms, Georg
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