Abstract Title: Electrical transcutaneous pain suppression and pain location device

The invention relates to a device for treating a patient for neurological or muscular pain by neurostimulation comprising a pair of electrodes to be applied externally, i.e. to the surface of the skin, in the region of the pain and to apply a current of between 0.2 and 12 mA at a frequency of between 1 and 50Hz, and preferably between 2 and 10 Hz. Optimal results appear to be achieved when the applied current is between 3 and 10 mA. In use at least one of the electrodes, a stimulating electrode is used to accurately locate the pain and during treatment the electrode is applied to the patient’s skin as located using mild pressure, whilst a stimulating pulse is applied as treatment. This action produces a remarkable and unexpected level of pain relief. The invention extends to a method for the treatment of neurological pain using the above parameters and procedure.
FIG. 3
IMPROVEMENTS IN AND RELATING TO
NEUROSTIMULATION

This invention relates to improvements in neurostimulation for the treatment of chronic pain.

Neurostimulation is gaining in popularity as a treatment of chronic pain. Traditionally such neurostimulation has relied on the implantation of a device in a position adjacent to an affected area. An electrical stimulation is applied to implanted electrodes to achieve a level of relief in the patient. However, such devices need to be accurately located and generally require a small surgical procedure in order to implant the device. The procedure is carried out by a trained medical practitioner or specialist.

In accordance with the invention a device for the treatment of neurological or muscular pain by electrical neurostimulation comprises a pair of electrodes and an electronic power supply arranged to supply a pulsed signal between the electrodes of which one electrode is a reference electrode for attachment to the skin of a patient, and the other electrode is a stimulating electrode preferably comprising a rigid stem with a substantially rounded end which is arranged, in use, to be pressed firmly onto a patient’s skin both to accurately locate the source of pain in response to a locating pulse and then to remain in place as located to treat the pain by applying an appropriate stimulating pulsed signal transcutaneously to treat the pain.

The fact that the electrodes are applied externally for non-invasive use permits the device to be used and, to a certain extent, controlled by a patient, although its initial set-up will be performed by a medical practitioner or specialist. The very low current which has been found to be effective allows the device to be manipulated safely by older people – the most frequent sufferers from chronic pain – or even by children or people who are handicapped.
The present invention must be distinguished from the known transcutaneous electrical nerve stimulation (TENS) in which silicon-rubber/carbon electrodes are stuck onto a patient’s skin in the region of an affected or painful area. In the present invention, a stimulating electrode is arranged to be accurately located with respect to an affected nerve. In addition the stimulating electrode has a rigid, rounded end, generally at the end of a rigid stem, which can be 1) applied firmly to the skin in the affected area as located either by the device in a location mode or by a separate nerve locator, and 2) moved in response to the patient’s reaction to the stimulating signal applied to the stimulating electrode further to fine-tune the treatment.

Once the stimulating electrode has been accurately located the current can be reduced in most cases and the frequency can be adjusted to produce the optimum level of relief. Accurate location of the electrode thus reduces the discomfort to the patient and the risk of burning. It also prolongs battery life for a portable device.

The energy required for the treatment may depend on the size of the nerves (large, heavily myelinated A motor fibres at one extreme versus smaller unmyelinated C fibres at the other). Thus the energy delivered \( E \) (energy in nC) = \( I \) (current in mA) \( \times t \) (duration in s). In order to obtain the necessary transcutaneous current a relatively high voltage – up to 85V, though more traditionally 65V - may be required. The pressure that can be applied by the stimulating electrode (and by the other, reference, electrode) to the skin allows the voltage to be reduced because of the lower resistance, making burning less likely. The rounded end of the electrode(s) also act to distribute the current more accurately and evenly than would be achieved by a traditional silicon carbon electrode.

The treatment is found to be effective in most cases in about five minutes, though it may be applied for less time or for longer according to the symptoms and level of relief achieved.

One of the most important components of this novel approach to peripheral neuromodulation in treatment of neuropathic pain is the frequency of the
stimulation, which is contrary to established recommendations. The frequency is particularly effective in the slow range of stimulation 2 – 10Hz, but can occasionally be successful at higher levels 10 – 50Hz, but not usually in the 50 – 150 Hz range.

This external approach whereby the electrical impulse is applied externally over the nerves, plexuses and at non-specific areas in a non-segmental distribution produces results which are overwhelming. Pain relief can be compared to the percutaneous direct approach, with results of the same magnitude of 70 -100% pain relief achieved as measured on a VAS score.

The duration of the pain relief following a typical 5 min session where the electrodes have been placed correctly can vary between minutes or hours and days or even weeks.

The device is most effective when the stimulating electrode comprises a substantially hemispherical contact which projects in such a way that they can be applied to an affected area with sufficient pressure that it produces a small indentation in the skin. In one embodiment, one or both of the electrodes are mounted on a semi-rigid support or supports having straps or other means which when in use enable each contact to be pressed firmly against a patient’s skin in the affected region.

In other embodiments one of the electrodes, the reference electrode, comprises a silicon-carbon patch connected to the power supply and is arranged to be stuck to the skin close to the affected area. A gel may be used to enhance the conductivity between the reference electrode and the skin. The other electrode, the stimulating electrode, comprises a short rod or stem whose contact end is rounded or is provided with a small ball which is applied to a patient as described above. The stimulating electrode may either be rigidly attached (screwed in or permanently fixed, for example) to the power supply, or attached to it by a wire as mentioned above.
This combination of pressure together with the electrical stimulation provides a surprising level of relief comparable to that achieved by a percutaneous implant. In a recent trial, in more than half the patients on which the device was used the VAS pain score was reduced to 0 over a period of five minutes.

For maximum effectiveness the electrodes are accurately positioned using the device in its stimulator and location mode prior to treatment. Thus, the device is first used to position the treatment electrodes prior to fine-tuning the treatment current and frequency. The power supply may then be adjusted and used to provide the desired electrical output to the electrodes for the treatment.

As it may not be necessary to carry out the location procedure on each occasion, either a separate nerve stimulator may be used by a specialist practitioner, or the location function may be switched off so that the patient is not confused. The treatment parameters equally may be pre set or pre-limited to prevent the patient from harming himself by mistake.

The diameter of the contacts of the electrodes will depend on the muscle or nerve type to be treated. They may be between 1 and 12mm or in some cases up to 20mm, but generally 3 to 6mm. They may be mounted on the inner side (in use) of a semi-rigid strap which may be attached to a patient by means of a sticky plaster or strip of Velcro (® 3M Corp. Inc.).

As electronic devices become ever smaller and more compact, one embodiment of the device may be in the form of a pen. The stimulating electrode is located at the 'writing end' whilst the reference electrode is stored at the other end of the 'pen' and is arranged to be connected to the patient by means of a plaster or a patch. A gel may be required to improve the conductivity between the reference electrode and the skin.

The invention extends to a method of treating a patient for neurological or muscular pain by neurostimulation comprising applying a pair of electrodes externally, ie. to
the surface of the skin, in the region of the pain and locating the stimulating electrode accurately prior to applying a stimulating pulse of between 0.2 and 12 mA at a frequency of between 1 and 50Hz, and preferably between 2 and 10 Hz. Optimal results appear to be achieved when the applied current is between 3 and 10mA. As mentioned above, the electrodes are applied to the patient’s skin using mild but firm pressure which causes them to produce an indentation in the skin. This action appears to enhance the success of the treatment and indeed makes effective nerve stimulation possible by means only of externally-positioned electrodes.

External neurostimulation in accordance with the invention results in improved relief of chronic neuropathic pain, an improvement in peripheral circulation, improved mobility and improved sensory perception, comparable with the results achieved by percutaneous treatment. It has the great advantage that it can be applied by the patient him/herself when required and as often as they wish with little or no adverse effects. As no surgical procedure is required, treatment can be easily be modified or stopped if it fails to produce the desired relief or if it causes an unwelcome response.

Additionally, it can be used in conjunction with traditional implants either to complement the relief or if the site of the pain moves. This is particularly useful in the treatment of non-specific or non-segmental pain.

The invention will now be further described by way of example with reference to the accompanying drawing in which:

Figure 1 is a diagrammatic view of a device in accordance with the invention,

Figure 2 is a diagrammatic side elevation of several electrode forms suitable for use with the device.

Figure 3 is a diagrammatic view of a device in accordance with the invention, similar to that shown in Figure 1,
Figure 4 is a diagrammatic view of a device in accordance with the invention, similar to that shown in Figure 3.

Figure 5 is a diagrammatic view of a device in accordance with the invention in the form of a pen. 5A shows the device as stored, and 5B shows it ready for use, and Figure 6 shows a typical wave form of a stimulating signal.

In Fig. 1 the device comprises a control unit which incorporates a power supply 10 which is controlled by a frequency button 12 and a power button 14. The output parameters are displayed on an LCD screen 16. An on/off switch is provided at 18.

The output from the device is supplied to two electrodes 20,22 by leads 24. The electrodes 20,22 each comprise a substantially hemispherical projection protruding from a semi-rigid strap 26. The strap shown is for attaching around a patient’s wrist and has a length of Velcro® 28 at one end which attaches to a corresponding patch 29 on the other side of the strap.

The electrodes each have a diameter of 4mm and project 8 to 10mm from the strap. They are held firmly on the strap so that when the latter is attached, for example, around a patient’s wrist, they cause an indentation in the patient’s flesh. They are spaced apart by about 20 to 30mm in most cases, though they may be adjustable as the exact required spacing will depend upon the diagnosis and the treatment required. In practice one of the electrodes acts to produce a contact with the patient’s skin as a reference electrode 22 whilst the other, or stimulating electrode 20, is positioned in respect of the point in the affected area, which produces the optimum level of relief.

Figure 2 shows elevations of several electrode forms suitable for use with the device. The electrode in Fig 2A has a stem 32 connected to a lead 24 from the power supply 10, and a rounded contact end 34 substantially in the form of a
sphere. This electrode form is used for larger sites and may measure up to about 20mm in diameter. It may be used with the higher currents prescribed.

Fig. 2B shows a flat elliptical electrode measuring up to 10mm by 20mm for use as the electrode in Fig. 2A. Fig. 2C shows a rectangular electrode for applications similar to those for which that in Fig. 2B is used.

Fig. 2D shows the preferred spherical electrode whose spherical contact end 34 is about 5 to 6mm in diameter. This electrode appears to produce good results in the majority of cases. However for specific applications it may be smaller, down to 1mm, or indeed larger as required.

Whilst a pair of similar electrodes may be used, a practitioner may decide that two dissimilar electrodes may produce a better response for a specific condition or may attach a reference electrode to the patient by means of a patch whilst using a solid metal electrode as shown in Figure 2 for the stimulating electrode in order to be able to apply it sufficiently firmly to the patient’s skin to achieve an improved level of relief.

In Figure 3 the device is similar to that in Figure 1 but that the reference electrode comprises a silicon-carbon patch 30 connected to the power supply 10 and arranged to be attached to the skin close to the affected area. A gel may be used to enhance the conductivity between the reference electrode and the skin. The other electrode, the stimulating electrode 20, comprises a short stem 32 whose contact end 34 is rounded or has a small ball formed at its end, or indeed any suitable form shown or described above in Figure 2. In use a stimulating pulse is applied to the stimulating electrode to find an accurate location of the affected nerve. Once this has been correctly located, the electrode is applied firmly to the patient’s skin whilst the stimulating signal is adjusted to provide the optimum level of relief without minimising any discomfort.
Figure 4 shows a device similar to that in Figure 3, but that the stimulating electrode 20 is rigidly attached for use by being screwed or clipped into the power supply 10. Alternatively it may be permanently fixed to it. In all other respects it is similar to that shown in Figure 3. The stimulating electrode must however be sufficiently stiff and suitably dimensioned that it can be applied firmly to a patient to produce a good electrical and physical contact with the skin.

Figure 5 shows a stimulating device in the form of a pen whose body houses the power supply 10 and the necessary controls 18 (on/off and function button), increase button 40, decrease button 42 and an LCD screen 16. A stimulating electrode 24 in the form of a small ball is provided at the ‘writing end’ whilst a reference electrode 22 is stored at the other end of the ‘pen’ under a screw cap 44 (5A). In use, as shown in 5B, a reference electrode 22 is withdrawn from the upper end of the ‘pen’ and extended so that it can be attached to the skin of a patient by means of a plaster 30 as above. The ‘pen’ may be clipped into a patient’s pocket, ready for use, by means of a clip 48.

Other forms of the device are possible. For example, many sufferers of acute pain have muscular or other disabilities which make it difficult for them to hold the device in a way that applies the appropriate pressure in the desired location. In such cases it may be appropriate to provide a ‘pistol’ grip with the electrode, for example, at the end of the barrel.

In use, the combination of the pressure and the, albeit small, stimulating current appears to have a remarkably beneficial effect in terms of the pain relief achieved.

The power button 14 of the power supply is adjustable to provide a current of 0.2mA to 12mA at a frequency that can be varied by the frequency button 12 from 2 to 10 Hz. It has been found that for many patients the optimum stimulation occurs at a frequency of 2 to 3 Hz and a current of between 3 and 10mA. An example of the stimulating pulse is shown in Figure 6. In this case a square-wave with a frequency of 2Hz has a pulse with a duration of, say, 10% of the wave length. This appears to
produce an effective treatment whilst enhancing battery life in the case of a portable device. The duration of the pulse may vary from about 5% to 25% of the wavelength whilst remaining effective for the treatment. The shorter the pulse and the lower the current the longer the battery life in the case of a portable device.

In a trial, a device in accordance with the invention was used to treat patients with the following pain presentations: CRPS with scar areas on the pelvis, testes, abdomen, chest wall, neck, breast, as well as for phantom limbs and chest pain. The following Table summarises the results of the treatment

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<thead>
<tr>
<th>Patients</th>
<th>% pain relief</th>
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<tr>
<td></td>
<td>VAS score</td>
</tr>
<tr>
<td>17</td>
<td>100</td>
</tr>
<tr>
<td>4</td>
<td>90</td>
</tr>
<tr>
<td>8</td>
<td>50 – 90</td>
</tr>
<tr>
<td>3</td>
<td>&lt;50</td>
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Whilst such effective treatment would in all probability have been achieved using an implanted device, the success of this non-invasive, external treatment is astonishing. The flexibility and convenience of external stimulation is thus greatly appreciated by patients and specialists alike.
CLAIMS

1. A device for the treatment of chronic pain by means of electrical neuro-stimulation comprising a pair of electrodes and an electronic power supply arranged to supply a pulsed signal between the electrodes of which one electrode is a reference electrode for attachment to the skin of a patient, and the other electrode is a stimulating electrode comprising a rigid stem with a substantially rounded end which is arranged, in use, to be pressed firmly onto a patient’s skin both to accurately locate a source of pain in response to a locating pulse and then to remain in place as located to treat the pain by applying an appropriate stimulating pulsed signal transcutaneously to treat the pain.

2. A device as claimed in claim 1 in which the power supply is arranged to deliver a pulsed signal at between 1 and 50Hz, and preferably between 2 and 10Hz.

3. A device as claimed in claim 1 or 2 in which the power supply is arranged to operate at a voltage of 50 to 85 volts, and preferably at approximately 65 volts.

4. A device as claimed in any preceding claim which is capable of supplying a current of between 0.2mA and 12mA to the electrodes when applied externally to the skin of a patient.

5. A device as claimed in any preceding claim in which the power supply is arranged to be capable of supplying a current of between 3 and 10mA at a frequency of between 2 and 10Hz.

6. A device as claimed in any preceding claim in which the electrodes comprise substantially hemispherical contacts which project from a support or straps which enable each electrode when in use to be pressed and held firmly against a patient’s skin in the desired region.

7. A device as claimed in any preceding claim in which the stimulating electrode has a contact area with a diameter of 3 to 20mm.

8. A device as claimed in claim 6 or 7 in which the stimulating electrode has a diameter of between 3 and 12mm and preferably between 3 and 5mm.
9. A device as claimed in any preceding claim in which the reference electrode comprises a flexible silicone-rubber/carbon electrode for attachment to the skin of a patient.

10. A device as claimed in any preceding claim in which the stimulating electrode is rigidly attached to or forms part of the power supply.

11. A device as claimed in claim 10 in which the power supply is in the form of a pen whose ‘writing end’ comprises the stimulating electrode and a flexible reference electrode is connected to or near the other end.

12. A device as claimed in claim 10 in which the power supply is in the form of a pistol grip to allow a person to hold it firmly in order to apply pressure to the stimulating electrode at its extremity.

13. A method for treating chronic or acute pain by neurostimulation comprising accurately locating the affected nerve by means of electrical neurostimulation and positioning a stimulating electrode externally over the nerve and applying a low frequency electrical stimulating pulse to it.

14. A method as claimed in claim 13 in which a current of between 0.2 and 12 mA is applied to the stimulating electrode at a frequency of between 1 and 50Hz.

15. A method as claimed in claim 13 or 14 where the applied frequency is between 2 and 10 Hz.

16. A method as claimed in any of claims 13 to 15 in which the applied current is between 3 and 10mA.

17. A method as claimed in any of claims 13 to 16 in which the current is applied for 5 minutes.

18. A device as claimed in any of claims 1 to 12 and substantially as hereinbefore described with reference to the accompanying drawings.

19. A method as claimed in any of claims 13 to 17 and substantially as hereinbefore described.
Patents Act 1977: Search Report under Section 17

Documents considered to be relevant:

<table>
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<th>Category</th>
<th>Relevant to claims</th>
<th>Identity of document and passage or figure of particular relevance</th>
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<tr>
<td>Y</td>
<td>1 at least</td>
<td>US4769881 A (PEDIGO) See figure 1, 5 and columns 4, 9, 10</td>
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<tr>
<td>Y</td>
<td>1 at least</td>
<td>US4962766 A (HERZON) See especially figure 1 and the abstract</td>
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<tr>
<td>Y</td>
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<td>US4233986 A (AGAR) See figures, abstract and columns 2 to 3</td>
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<tr>
<td>Y</td>
<td>2-5</td>
<td>US4155366 A (ULTRA AIDS) See whole document, especially abstract</td>
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<tr>
<td>A</td>
<td>-</td>
<td>US3830226 A (CONCEPT) See abstract and figures</td>
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Categories:

- X Document indicating lack of novelty or inventive step
- Y Document indicating lack of inventive step if combined with one or more other documents of same category.
- & Member of the same patent family
- A Document indicating technological background and/or state of the art.
- P Document published on or after the declared priority date but before the filing date of this invention.
- E Patent document published on or after, but with priority date earlier than, the filing date of this application.

Field of Search:

Search of GB, EP, WO & US patent documents classified in the following areas of the UKC:

- A5R
- Worldwide search of patent documents classified in the following areas of the IPC:
- A61N

The following online and other databases have been used in the preparation of this search report:

- Online: OPTICS WPI EPODOC