Spatial-Temporal Deep Brain Stimulation Methods and Systems

10-4-2011

Richard Gilson
University of Central Florida

Nizam Razack
Nizam Razack

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Methods, systems, and devices to reduce power demands substantially for current deep brain stimulation (DBS) using smart technology type applications. The invention uses miniaturized components that allow integration with the implanted probe(s) themselves, and includes a skull-sited housing having all the controls and battery power supply needed. This avoids implanting obtrusive card-deck size batteries in the chest area and the use of vulnerable wire leads under the skin from the chest area to connect with the implanted electrode(s) on the skull, improving comfort. The generating of non-continuous pulses on demand of conditions such as the occurrence of a tremor occurs, without having to continuously run pulses at all times, substantially increasing life spans over current techniques. Shaped electrodes and their methods further reduce power demands and efficacy by directing electric fields to focus towards specific areas and regions of the brain rather than inefficient 360-degree emission.

12 Claims, 12 Drawing Sheets
Fig. 9

SIGNAL LEADS 200

NOISE AMPLITUDE FILTER 210

IMPULSE AMPLITUDE FILTER 220

SENSING CIRCUIT 230

IMPULSE FIRING THRESHOLD CIRCUIT/TRIGGER SWITCH 240

STIMULATING GENERATOR 250

OFF

FIELD OF INVENTION

This invention relates to deep brain stimulation (DBS), in particular to methods, systems, and devices for enhancing both efficacy and efficiency of “in-vitro” electrical stimulation known to inhibit symptoms of neurological diseases, disorders, and the like, and utilizes: (a) methods that tailor the direction and shape of electrical fields towards the area of best effect; (b) systems that interactively trigger stimulation to terminate the onset of tremors, and such; and (c) devices that use the aforementioned efficiencies to reduce power and allow for self-contained miniaturization at the implant site, thereby improving patient comfort.

BACKGROUND AND PRIOR ART

A first line of defense considered for most neurological disorders is treatment with psychoactive medicines, prior to any surgical intervention. Psychoactive medicines or drugs are those capable of acting on the nervous system and affecting mental states and behavior. Many physiological mechanisms are aided by such drugs, but these medicines have unwanted side effects, drug interactions, and long-term physiological tolerances that render the drug less effective over time.

In the case of Parkinson’s disease that serves as the exemplar disorder herein, the underlying etiology is that neurons, located in the substantia nigra of the mid-brain, for some unknown reason begin to produce less dopamine. As these neurons progressively and relentlessly deteriorate over years, less dopamine dramatically affects the motor control of the basal ganglia and thus outward behavior and everyday living. An estimated 1.5 million Parkinson’s patients have a visible cluster of diagnostically significant and debilitating Parkinson’s symptoms, typically tremors, stiffness, slowness, and balance. Patients describe the internal feeling as frozen still in the “set” stage of the “ready, set, go” sequence that started at a race.

The “gold” standard treatment for Parkinson’s is frequent daily administration of an indirect dopamine replacement, L-Dopa, a psychoactive drug that crosses the blood-brain barrier, and then alters form to produce dopamine as a supplement to the brain’s own production. This therapy, while initially quite beneficial, typically can lose much of its effectiveness over about five years, wherein patients have to take progressively larger and more frequent doses until eventually the result becomes inadequate. Without alternative drug therapies, patients are left to suffer both the ravages of primary symptoms and the manifestations resulting from the prolonged use of the drug itself, principally the repetitive spasmodic motions of dyskinesia. More than one million people endure these symptoms, including such notables as the Pope, Muhammad Ali, and Billy Graham, while thousands of others including Michael J. Fox have turned to surgical approaches.

Past surgical techniques treated symptoms of these diseases by selectively and permanently destroying or ablating structural areas in the brain. The net effect is to “shut the door”, in a neurological fashion, before dysfunctional brain signals are sent to the muscles, thereby relieving many symptoms. The advantage of surgical ablation is that it reduces the reliance on drug therapy with its attendant side effects. The disadvantage is that the procedure is irreversible. This may render such patients as unacceptable candidates for newly discovered techniques/therapies, such as stem cell implantation or viral transport of genome-altered DNA (deoxyribonucleic acid), both of which show promise in helping to augment or even to regenerate the natural production of dopamine as well as a number of other substances involved in neurological disorders.

Certain neurological disorders that produce debilitating motor symptoms are now being treated with Deep Brain Stimulation (DBS) through-skull implanted electrodes see FIG. 1 below. DBS essentially reversibly alters the local neurological structure(s) around the tip of an electrode implanted on the brain with electrical pulses that reduce or stop disabling symptoms, such as, but not limited to, severe tremor and rigidity found in Parkinson’s disease.

DBS functionally has the advantage of emulating ablation by changing the firing characteristics of nearby neurons, but it does so only while the pulsed stimulation persists. Since the structures remain intact and undamaged, when DBS is turned “off” these structures reactivate and symptoms return, unless otherwise treated. Thus, DBS overcomes the chief disadvantage of ablation in that it allows implanted stimulators to be withdrawn later for new techniques, with minimal residual effects. In 1998, the Federal Drug Administration (FDA) approved DBS as an alternative for, or as an adjunct to powerful psychoactive drugs (neuro-medicines) burdened with strong and often unacceptable side effects.

The only FDA approved DBS apparatus is currently being sold by Medtronic, Inc., 710 Medtronic Parkway, Minneapolis, Minn. 55432-560, see FIGS. 2 and 3 below, although other companies have substantial interests in implantable neurostimulation devices for a variety of neurological disorders, such as Advanced Bionics, Corp., 12740 San Fernando Road, Sylmar, Calif. 91342; see “http://www.advancedbionics.com.” However, there are problems with the approved apparatus.

At the very least, the presently approved apparatus is cumbersome and uncomfortable for many in its present form. As an outgrowth of legacy components from “pacemakers,” it consists of one or two remote stimulator/battery packs embedded under muscle tissue in the upper chest area, and requires subcutaneous leads up along the neck to the skull entry point. All these components are subject to corrosion and breakage, as well as to resistance or attacks by the body itself attempting by encapsulate it or dissolve it, leading to infection.

Medtronic currently uses a product called a Soletra™ Neurostimulator (see FIG. 4 below) to generate a continuous series of electrical pulses, typically at about 2-4 volts, to electrode(s) implanted in specific brain areas. Neurons, normally operating in the tens of milli-volts range, are massively over-stimulated by such voltage and temporally altered, thereby inhibiting the expression of certain motor dysfunctions. Pulse voltage is usually adjusted somewhat depending on the proximity of the electrode(s) to the targeted area, e.g., more voltage is needed for target variations. Typical pulse rates range around 130-185 pulses/second.

This combination of large amounts of voltage, amperage, and duty cycle creates a power drain that normally requires a non-renewable battery replacement in about 3-5 years, at a price of about $10,000. Many patients turn “off” the stimulation when going to sleep while tremor is quelled, in order to
conserves power. Current DBS stimulator(s) are designed to be turned “off” or “on” using a magnetic-switch placed briefly near the associated electronics.

Thus, without considering the remote apparatus itself, or the power requirements of the present design, a goal in this field should be to reduce stimulation-related side effects and complications caused by stimulating in the vicinity of the target. Two key objectives to meeting that goal are for the surgical implant to hit the center of a targeted cellular region, often not more than 2 mm across in any direction, and for the current field to stimulate the appropriate cells for symptom reduction or cessation, without triggering side effects in adjacent structures.


A key to a successful outcome is the precise positioning and placement of the electrode(s), aided pre-operatively by vast improvements in brain imaging techniques, but nevertheless requiring considerable surgical skill and judgment. Beside normal variations in brain structures themselves adding to the difficulty of targeting, during surgery the brain moves with each heartbeat, while its size, and position vary somewhat as a result of the surgical probe itself altering internal pressure. During the procedure, the use of fluoro-scope imaging assists y-z axis positioning, and awakening of the patient while in the operating room for testing of clinical signs can reveal and help avoid untoward side effects. However, the final outcome can only be assessed post-operatively. Results can vary by patient and over time. Some post-surgical adjustment is normally done with electrical parameters and z-axis programming of one or more of the electrode contacts available on the current DBS electrode, as is described below, but not to the extent that many physicians would like and not with respect to the x-y axis.

Current DBS electrodes have four (4) circumferential contacts that radiate current in a 360 degree configuration. This means that with any degree of off-target, or unnecessary stimulation even on target adjacent cells are exposed to current and potential side effects. While some of these side effects may be adjustable when electrical parameters are altered, or even reversible when the stimulation is shut down, the cost is decreased efficacy of stimulation on the symptoms. For example, in thalamic nucleus DBS, stimulation-induced side effects may include increased dyskinesias, blepharospasm or so-called “eyelid-opening apraxia,” confusion/memory disturbances, personality changes, mood changes, apathy, cognitive changes, dysphoria/dysarthria, and such.

Medtronic Inc. has proposed a “Directional Brain Stimulation and Recording Leads, title, in U.S. Published Patent Application 2002/0183817 to Van Venrooij et al., which is incorporated by reference. The proposed technique uses a “controller” as shown and referenced to FIG. 32 for recording “brain activity signals” to activate electrodes. However, this technique requires continuously generating pulsed type signals once the electrodes are activated whether or not a brain type tremor has ended, which would result in needless, unwanted and potentially-excessive electrical current being continuously generated inside the brain. The more unnecessary pulse type signals, the more undesirable side effects to the patient, for example, in thalamic DBS, stimulation-induced side effects may include paresthesias, muscular cramps, dystonia, dizziness, dysarthria, gait and balance disturbances, limb ataxia, impaired proprioception, and decreased fine motor movements.

Additionally, this technique would require excessive power to operate, which is not only expensive since battery power supplies would need to be regularly replaced but also require large card-deck size batteries that must be mounted inside of the patient’s upper chest area. This proposed Medtronic technique would also be prone to circuit problems since the electrodes would be simultaneously operating as both transmitters and sensors, causing excessive and unnecessary power drain, shortening the lifespan of any batteries being used as well as increasing the costs for replacing the batteries.

According to the Movement Disorder Society© 2002, “Deep brain stimulation for the alleviation of movement disorders and pain is now an established therapy. However, very little has been published on the topic of hardware failure in the treatment of such conditions irrespective of clinical outcome. Such device-related problems lead to significant patient morbidity and increased cost of therapy in the form of prolonged antibiotics, in-patient hospitalization, repeat surgery, and device replacement [Joint, C., Nandi, D., Parkin, S., Gregory, R., and Aziz, T. Hardware-Related Problems of Deep Brain Stimulation. Movement Disorders, Vol. 17, Suppl. 3, 2002, pp. S175-S180].

Thus, the need exists for solutions to the problems encountered in the prior art.

SUMMARY OF THE INVENTION

A first objective of the present invention is to provide for deep brain stimulation (DBS) methods, systems and devices that avoid implanting large card-deck size batteries remotely in the chest area and the use of vulnerable wire leads under the skin from the chest area to connect with the implanted electrode on the skull.

A second objective of the present invention is to provide for deep brain stimulation (DBS) methods, systems and devices that allow for substantially all electronics including but not limited to an on-site battery pack to be located at or near the implanted electrodes on the skull.

A third objective of the present invention is to provide for deep brain stimulation (DBS) methods, systems, and devices that can use substantially less power than current DBS techniques and apparatus.

A fourth objective of the present invention is to provide for deep brain stimulation (DBS) methods, systems, and devices, that do not generate a continuous series of un-synchronized pulses that are continuously generated whether they are needed or not.

A fifth objective of this invention is to provide for deep brain stimulation (DBS) methods, systems, and devices, to generate pulses on demand. Specifically, inhibiting pulse(s) can be interactively triggered by the onset of each unwanted electro-physiologically signaled event, such as at the beginning of a tremor.

A sixth objective of the present invention is to provide for deep brain stimulation (DBS) methods, systems and devices that are more compact, easier to use and last longer than current DBS techniques, thereby improving patient comfort.

A seventh objective of the present invention is to provide for shaped electrodes and methods by using the electrode contacts in deep brain stimulation (DBS) methods, systems and devices, that orient configured electrical fields to specific points, preferentially directing current at targeted cellular region and facilitating the clinical effect. The best direction of
the shaped current can be determined intra-operatively through recordings and clinical testing along the "X-Y" plane.

An eighth objective of the present invention is to use shaped electrodes in deep brain stimulation (DBS) methods, systems and devices, in order to decrease off-target current stimulation, thereby minimizing or eliminating exposure to adjacent cellular regions that produce side effects.

A ninth objective of the present invention is to use the decreased electrical "aperture" of shaped electrodes in deep brain stimulation (DBS) methods, systems and devices, in order to maintain current density on the target region, but with less overall current flow and less battery drain over time.

Methods, systems and devices are provided for deep brain stimulation (DBS), that can include providing a pulse stimulator with battery supply and controls within one unit, imbedding the single unit in a head of a user, and generating a single treatment pulse from the stimulator to the brain of the user, the single pulse being generated on demand of a sensed initiation of an impulse burst from a condition. Another single pulse can be generated upon occurrence of a follow-up sensed impulse burst. The impulse bursts can come from a detected brain activity condition such as those associated with tremors, and the like.

The methods, systems and devices can also provide the battery power supply solely in a compact skull-mounted cylindrical disc cap without implanting of batteries in a chest area of the user and avoiding any use of wires under skin from the chest area to connect to the stimulator, mounted as a skull cap with the top cap disc cover and the bottom cylindrical disc housing underneath and covered by the scalp, with a single opening in the skull for the lead line, further improving patient comfort.

Controls can also include filtering sensed signals from electrode sensors to a noise amplitude filter, followed by counting impulses in the filtered signals with an impulse counter, and then triggering the single treatment pulse when a selected threshold of the counted impulses has been reached. Furthermore, controls can allow for shutting off the noise amplitude filter, and the impulse counter when the triggering of the single treatment pulse has been initiated.

Various types of directional components can be provided for directing the treatment signals to effected areas of the brain of the user.

Further objects and advantages of this invention will be apparent from the following detailed description of the presently preferred embodiments, which are illustrated schematically in the accompanying drawings.

**BRIEF DESCRIPTION OF THE FIGURES**

FIG. 1 shows a deep brain stimulation (DBS) apparatus of the prior art.

FIG. 2 is a front view of the current prior art DBS of FIG. 1.

FIG. 3 shows DBS head-mounting details of the prior, art of FIGS. 1-2.

FIG. 4 is another prior art product device.

FIG. 5a shows an open view of a miniaturized integrated battery/stimulator apparatus of the subject invention.

FIG. 5b shows a closed view of the apparatus of FIG. 5a.

FIG. 6 shows the novel invention apparatus of FIGS. 5a-5b utilized in a head mounted application.

FIG. 7a shows enlarged side views of four shaped electrodes with projected fields used in the apparatus of FIGS. 5a, 5b and 6.

FIG. 7b illustrates the axial views of the four shaped electrodes of FIG. 7a with respective projected fields.

FIG. 8a shows a recording trace graph of neuronal impulse bursts in millivolts verses time that can occur with an untreated patient.

FIG. 56 shows a graph of associated tremor displacement for FIG. 8a in millimeters verses time that can occur with the untreated patient.

FIG. 9 shows an exemplary flow chart of a triggering algorithm that can be used with the invention.

FIGS. 10, 10a and 10b shows a layout of the electronic components that can be used in the apparatus of FIGS. 5a, 5b, 6 along with the triggering algorithm of FIG. 9.

FIG. 11a shows a graph of the neuronal impulse bursts in milli-volts verses time of a patient being treated with a Medtronic type (prior art) technique.

FIG. 11b shows a graph of the associated tremor displacement for FIG. 11a verses time of the patient being treated with the Medtronic type (prior art) technique.

FIG. 11c shows the continuing pulse train that occurs with the Medtronic type (prior art) technique which causes the trace graphs of FIGS. 11a-11b.

FIG. 12a shows a graph of the neuronal impulse bursts in milli-volts verses time of a patient being treated from being treated by the subject invention.

FIG. 12b shows a graph of the associated tremor displacement for FIG. 12a verses time of the patient being treated by the subject invention.

FIG. 12c shows the single pulses that occur with the subject invention technique which results in the trace graphs of FIGS. 12a-12b.

**DESCRIPTION OF THE PREFERRED EMBODIMENTS**

Before explaining the disclosed embodiments of the present invention in detail it is to be understood that the invention is not limited in its applications to the details of the particular arrangements shown since the invention is capable of other embodiments. Also, the terminology used herein is for the purpose of description and not of limitation.

This invention provides methods, systems and devices to reduce power demands substantially for DBS (deep brain stimulation), allowing for miniaturized type components to be integrated with the implanted probe(s) themselves, see FIGS. 5a, 5b, 6 and 10. The invention avoids implanting large and cumbersome card-deck size batteries in the chest area and the use of vulnerable wire leads under the skin from the chest area to connect with the implanted electrode(s) on the skull. Notably, the remote stimulator/battery with its associated wiring are the chief source of complaints in the use of this technique. Other advantages of this invention are also apparent.

The invention substantially differs from that used in current techniques. Rather than generating a continuous series of un-synchronized pulses that act whether needed or not, pulses can be generated on demand. Specifically, inhibiting pulse(s) can be interactively triggered by the onset of an unwanted electo-physiologically signaled event, such as the beginning of a tremor. The same electrode(s) that now serves as a one-way input also can serve as a bi-directional conduit to sense the electrical beginnings of the event itself as well as delivering the pulse(s). Thus, with appropriately tailored, triggering algorithms, a preferred example of which is described below in reference to FIG. 9, an inhibiting pulse can be timed for delivery so as to suppress the event's full expression.
Using this approach, power drain reductions of conserva­

tively 85% or more are achievable, with the percentage
depending on the neurological events of interest. For
example, most tremors are less than approximately 10 tremor
cycles per second, and can be far less when the tremor is
interrupted by another intentional activity such as purposeful
movement, as in the case of Parkinson’s disease. Thus, the
constant power drain can be reduced from approximately 150
stimulating pulses/second using the current techniques
described in the prior art to now to as little as one stimulating
pulse countereacting each tremor onset using interactive trig­

Even greater power reductions are possible for “lower fre­
cquency” tremor, such as Essential tremor at approximately 5
to approximately 7 tremor cycles/second, or “low-frequency”
Parkinson tremor at approximately 3 to approximately 5 tremor
cycles/second.

With dramatically reduced power demands for DBS, this
invention then allows for miniaturized components to be inte­
grated with the skull cap attachment to the implanted probe(s)
sthemselves, as shown in FIGS. 5a, 5b, 6, and 10. F IG.
5a shows an open view of a miniaturized integrated battery/stimulator apparatus 1 of the subject invention. FIG.
5b shows a closed view of the apparatus 1 of FIG. 5a in a
cylindrical disc housing where the top cap covers the bottom
cylindrical disc of the disc housing. FIG. 6 shows the novel invention apparatus 1 of FIGS. 5a-5b utilized in a head
mounted application with the top cap disc cover and the
bottom cylindrical disc housing mounted as a skull cap under­
neath and covered by the scalp with the imbedded probe
comprising an exemplary triggering algorithm for selectively
activate electrodes and treat effected parts of the brain where
impulse bursts occur. FIG. 9 shows an exemplary flow chart of
a triggering algorithm steps that can be used with the inven­tion.
This nonconductive material can also be used to help direct
electrical field emissions to one side or to more specific
regions, and/or points, rather than to a 360-degree emission.
The invention can include a lead 30 can having a single

Example, an upper electrode can emit up to 360 degrees while
other electrodes are directed to emit in specific regions and/or

Alternatively, more than one electrode can be used on the
lead in series to one another. The nonconductive surface
region can be applied on the same side of the electrodes, or to
different side surface portions of the electrodes as needed.
Additionally, different combinations can be used. For

Additionally, circuitry can be added to control which elec­
trodes are being emitted at selected time periods, and the like.
For example, electrodes can be further programmed with a
microprocessor, to activate simultaneously and/or sequen­
tially, and/or staggered and/or over different combinations,
and the like. The invention can use directional type elec­
trodes, such as those shown and described in reference to U.S.
Published Patent Application 2002/0183817 to Van Venrooij
et al., which is incorporated by reference.

The miniaturized implantable electrode apparatus and
stimulation systems can also include those described in
United States Patent Application 20030023297, to Byers et
al. filed on Jan. 30, 2003, which is incorporated by reference.
This reference describes an eyelid stimulation system and
circuitry that causes a paralyzed eyelid to close or open by
passing an electrical stimulating current to a nerve.

The invention can be used to treat brain impulse bursts and
the resultant physical extremity tremors that result therefrom.
FIGS. 8a and 8b represent untreated brain impulse bursts and
resultant physical extremity tremor outcome that are
untreated. FIG. 8a shows the recording of impulse bursts
from the brain B, that results from the leads 30 that are
connected to the electrodes 40-70 (of the apparatus of FIG. 6).
The large amplitude bursts represent the impulses that result
in the tremors (impulses from brain to effected extremity).
FIG. 8b represents an extremity that is being affected by the
impulse (here for example, a single index right hand finger).
A laser measuring sensor was aimed at the index finger to
record the results shown in FIG. 8b. For example, a plus 5 mm
 apex reading would represent the finger moving forward from
a horizontal plane, and ~5 mm represents the finger moving
at the horizontal direction.

The invention can use a triggering algorithm to selectively
activate electrodes and treat effected parts of the brain where
impulse bursts occur. FIG. 9 shows an exemplary flow chart of
a triggering algorithm steps that can be used with the inven­tion.
The triggering algorithm can take a number of forms or
be adaptive, based on patient neuro-generative responses. In
a simple illustrative form, not intended to be limiting, trigger­ing
sensing can be based on exceeding a threshold-firing rate
measured in impulses/second.

FIG. 9 shows an exemplary triggering algorithm 200 for
using the electrodes 40-70 of the preferred embodiment
shown in the preceding figures that switches back and forth
between a sensing mode and transmitting mode. In the first
step 210, signals coming from leads 30 from the micro­
electrodes 40-70 in the implant receive a signal from the
electrodes 40-70 the first left trace signal from FIG. 8a) can be
passed through a noise amplitude filter 220 for example an
high pass amplitude filter cutting off a low frequency of for
example everything below approximately 20 milli-volts is
filtered out). Next during step 230, the filtered signal goes to
an impulse counter, such as a clock counter which can count
the number of impulses passing through, which can be rep­
resented in impulses per second. Next step 240, has an
impulse firing threshold type circuit that can be used that
filters out false alarms and allows for a selected threshold,
such as for example, 5 impulses within 50 milli-seconds,
would trigger the generator in step 250 to send a generated
pulse SP back down the signal leads 30 to the electrodes
40-70.

When the generator is triggered, a signal can also be sent
(along line Off) to simultaneously shut off the sensing mode
of the electrodes, allowing the electrodes to be in a transmitt­ing
turing mode state. Sensing mode parameters of the electrodes 40-70 can be in approximately 20 to approximately 60 millivolts, while the triggering/switching range of the electrodes 40-70 can be approximately 2 to approximately 4 volts. The novel triggering algorithm 200 allows the electrodes 40-70 to be switched back and forth between transmitting and sensing modes without overloading the circuit components, and thus maximizing power usage during operation.

Once triggered, stimulation blocking continues until firing impulse firing rate drops below the critical threshold value. The sensing and stimulation can use the same micro-electrode leads, alternatively, wherein the sensing circuitry first serves medical implants by emitting electrical micro impulses to/battery [about 20 by about 60 mm in a cutaneous wires of a neurological event onset, such as the beginning of a tremor. These signals, then in turn, can allow the triggering of the existing magnetic switch for turn “off” the sensor and turns “on” the stimulator for one or more pulses.

Various triggering algorithms can be developed for effective symptom suppression with animal studies. However, the safety of this invention’s procedure has already been established for humans with continuous and higher pulse rates through FDA (Food and Drug Administration) approval. Furthermore, this invention also exists for electronic retrofit to thousands of patients who have already been implanted previously, without further brain surgery since the existing DBS electrodes can serve as a two-way bi-directional conductor.

In vitro testing of this approach also can be possible using current systems by sensing electro-magnetic forces from subcutaneous wires of a neurological event onset, such as the beginning of a tremor. These signals, then in turn, can allow the triggering of the existing magnetic switch for turn “off” and “on”.

Other types of triggering mechanisms can include those described in U.S. Pat. Nos. 6,539,263 to Schiff; 6,366,813 to DiLorenzo; 6,301,492 to Zourenshayn; 6,038,480 to Hrdlicka et al.; 5,833,709 to Rise et al.; 5,716,377 to Rise et al. and 5,707,396 to Benabid, which are all incorporated by reference. Additionally, U.S. Patent Application 2002/0183817 to Van Venrooij et al., which is incorporated by reference. FIG. 11a shows a trace graph of the neuronal impulse 100 of FIG. 9 and apparatus components of FIGS. 5a, 5b, 6, 10 and FIGS. 11a-11c, 12a-12c.

Benefits of using the novel triggering algorithm of FIG. 9 and apparatus components of FIGS. 5a, 5b, 6, 10 and FIGS. 11a-11c, 12a-12c:

FIG. 11a shows a trace graph of the neuronal impulse bursts in mill volts verses time of a patient being treated with a Medtronic type (prior art) technique such as the one shown and described in reference to U.S. Patent Application Publication 2002/0183817 to Van Venrooij et al., which is incorporated by reference. FIG. 11b shows a trace graph of the associated tremor displacement for FIG. 11a verses time of the patient being treated with the Medtronic type (prior art) technique. FIG. 11c shows the continuing pulse train that occurs with the Medtronic type (prior art) technique which causes the trace graphs of FIGS. 11a-11b.

Referring to FIGS. 11a-11c, results in a continuing pulse train (FIG. 11c) in order to cause the effects of no impulse bursts (FIG. 11a) and no resultant physical tremors (FIG. 11b). As described in the background section of this invention this technique requires continuously generating pulsed type signals once the electrodes are activated whether or not a brain type tremor has ended, which would result in needless and excessive, unwanted and potentially dangerous electrical current being continuously generated inside the brain. The more unnecessary pulse type signals, the more undesirable side effects to the patient.

Additionally, the technique proposed in this patent application publication would require excessive power to operate, which are not only expensive since battery power supplies would need to be constantly replaced but also require large card size batteries that must be externally mounted outside of a patient or mounted inside of the upper chest or be constantly connected to an external power supply. This proposed Medtronic technique would also be prone to short circuit since the electrodes would be simultaneously operating as both transmitters and sensors, which also causes excessive and unnecessary power drain, which also shortens the lifespan of any batteries being used as well as increase the costs for replacing the batteries.

FIG. 12a shows a trace graph of the neuronal impulse bursts in mill volts verses time of a patient being treated from being treated by the subject invention described above. FIG. 12b shows a trace graph of the associated tremor displacement for FIG. 12a verses time of the patient being treated by the subject invention. FIG. 12c shows the single pulses that occur with the subject invention technique which results in the trace graphs of FIGS. 12a-12c.
Referring to FIGS. 12a-12c, the invention causes single spaced apart pulses to occur over greater periods of time, where each pulse occurs on-demand, and not as a continuous series of pulses. As shown by FIG. 12a, the initial impulse burst and subsequent impulse bursts have long delay time periods there between, which can result in substantially reduced or eliminated tremors resulting therefrom, FIG. 11b.

The novel invention allows for less undesirable electrical current to be generated within the brain, less side effects that result therefrom, less power consumption as well as other apparent benefits. The novel invention can use small batteries that were not able to be used in the prior art systems and techniques, and also results in power supplies that require less replacements as well. The novel invention is able to eliminate the need for large and cumbersome card type batteries or external power supplies, and results in a better system for treatment.

Differences between the subject invention and that described in the Medtronic technique shown in U.S. Patent Application Publication 2002/0183817 to Van Venrooij et al. are compiled in Table 1.

<table>
<thead>
<tr>
<th>Attribute</th>
<th>Medtronic</th>
<th>Invention</th>
</tr>
</thead>
<tbody>
<tr>
<td>Sensing</td>
<td>Loss of Effectiveness</td>
<td>Single Event</td>
</tr>
<tr>
<td>Trigger</td>
<td>Brain &quot;Activity&quot;</td>
<td>Initial Impulse Burst</td>
</tr>
<tr>
<td>Objective</td>
<td>Select Electrodes</td>
<td>Stop Single Tremor</td>
</tr>
<tr>
<td>Stimulator</td>
<td>Normally-On</td>
<td>Normally-Off</td>
</tr>
<tr>
<td>Power</td>
<td>&quot;off&quot; (continuing pulses)</td>
<td>Only &quot;on&quot; as needed</td>
</tr>
<tr>
<td></td>
<td></td>
<td>(individual pulse)</td>
</tr>
<tr>
<td></td>
<td>Large Battery</td>
<td>Small Battery</td>
</tr>
<tr>
<td>Power Life span</td>
<td>Limited</td>
<td>Extended</td>
</tr>
<tr>
<td>Battery Mounting</td>
<td>&quot;Implantable&quot;</td>
<td>Skull—Mounted in Cap</td>
</tr>
<tr>
<td>Side Effects</td>
<td>Continuous electrical current</td>
<td>Limited Electrical Current</td>
</tr>
</tbody>
</table>

Besides circuitry miniaturization, by triggering pulse generation only when needed, fewer pulses per second also can reduce some of the mild, but known side effects of DBS, and the timed delivery also can delay the onset of the next event by physiological feedback, that is also known to be effective against a known clinical phenomena Parkinson’s Disease.

Other advantages can also be realized. The timed pulse feedback speculatively could induce the brain to adaptively alter actual neural circuitry, allowing a further reduction of pulse rate and even more efficiency or even better help reduce symptoms naturally, just as brain circuitry is known to be altered by experience.

While the invention has been described for use with tremors, the invention can have applicability to other medical applications, such as but not limited to movement disorders and cardiac dysfunctions, but extends to include other neurological diseases and disorders such as epilepsy, psychiatric and behavioral dysfunctions such as schizophrenia and drug-induced symptoms and the like.

While the invention has been described, disclosed, illustrated and shown in various terms of certain embodiments or modifications which it has presumed in practice, the scope of invention is not intended to be, nor should it be deemed to be, limited thereby and such other modifications or embodiments as may be suggested by the teachings herein are particularly reserved especially as they fall within the breadth and scope of the claims here appended.

The invention claimed is:

1. A system for both sensing a single brain tremor and for providing deep brain stimulation (DBS) by a single treatment pulse, consisting of:

   a single cylindrical disc housing consisting of a single bottom and side edges extending from the bottom, and a single removable cylindrical top cover with perimeter sides which overlap the side edges of the housing, so that the cylindrical top cover covers a single compartment inside of the cylindrical disc housing, the single cylindrical disc housing having a height of approximately 15 millimeters, the single cylindrical disc housing adapted to be mounted as a cap on a skull of a user, and not being implanted beneath the skull of the user;

   a single elongated lead line extending beneath a center portion of the bottom of the single cylindrical disc housing having a plurality of electrodes thereon, the electrodes each being used to both sense a single brain tremor from an effected part of a brain and generate pulses therefrom, only the single elongated lead line being adapted to be implanted into only a single opening in a skull of a head of a user while the single cylindrical disc housing remains mounted as a cap on top of the skull of the user;

   a pulse stimulator inside of the compartment of the cylindrical disc housing, the pulse stimulator only becoming activated upon sensing of the single brain tremor from one of the electrodes on the single lead line, the pulse stimulator not generating any pulses when a tremor is not being sensed, the pulse stimulator includes a pulse generator for generating single spaced apart non-continuous pulses upon initial occurrence of the single brain tumor tremor sensed from the one electrode on the single implanted lead line;

   sensing detection circuitry solely located inside of the compartment of the cylindrical disc housing for sensing the single brain tremor from the one electrode on the single lead line, the one electrode sensing the brain tremor produced from the effected part of the brain is used for treating the effected part of the brain that produces the brain tremor;

   controls for the pulse stimulator that are all solely located inside of the compartment of the cylindrical disc housing, the controls for activating the pulse stimulator only upon the sensing of the single brain tremor from the one electrode on the single lead line, the pulse stimulator for generating a single treatment pulse only upon the occurrence of the sensed single tremor, the controls for deactivating the pulse stimulator so that no pulses are being generated when a tremor is not being sensed from the one electrode, the electrodes on the single lead line being connected to the controls for both sensing brain activity and for generating the single treatment pulse upon activation of the pulse stimulator, wherein the single treatment pulse is generated after the single brain tremor is detected by the one electrode; and

   a battery power supply for the stimulator solely located inside the cylindrical disc housing for providing all power for the pulse stimulator and all power for the controls, wherein the controls control the battery power supply to supply power to the pulse stimulator only upon the sensing of the single tremor, and wherein the controls turn off the pulse stimulator after sensing detection circuitry has been turned on, wherein the cylindrical disc shaped housing eliminates implanting of any batteries in the chest area and the use of any other wires under the skin from the chest area to connect with the implanted...
electrodes, and wherein no other lead lines and no other wires are connected to the single cylindrical disc housing and the single implanted lead line extending below the single cylindrical disc housing, and only the single cylindrical disc housing and the single implantable lead line are used for both sensing and for providing deep brain stimulation (DBS), the single cylindrical disc housing being positioned as the cap on top of the skull of the user and adapted to cover the cylindrical disc housing with the scalp.

2. The system of claim 1, wherein the controls include: a noise amplitude filter for filtering sensed signals from the electrodes that are below approximately 20 millivolts; an impulse counter for counting impulses in the filtered signals in impulses per second; and a trigger switch for triggering a single treatment pulse when a selected threshold of the counted impulses has been reached, wherein the controls shut off the noise amplitude filter and the impulse counter when the triggering of the single treatment pulse has been initiated.

3. The system of claim 1, wherein the cylindrical disc housing includes:

- dimensions of 60 millimeters in diameter.

- miniature batteries that are each approximately 1/36 of an AA battery.

5. The system of claim 4, wherein the plurality of electrodes includes:

- only four electrodes on the single implantable lead line.

6. The system of claim 5, wherein the only four electrodes includes:

- one electrode that emits in an approximately 360 degree field about the single implantable lead line; and

- three electrodes that each emit in limited selected directional regions.

7. The system of claim 2, wherein the controls include:

- sensing mode of the electrodes are approximately 20 to approximately 60 millivolts; and

- triggering range of the electrodes are approximately 2 to approximately 4 volts.

8. The system of claim 1, wherein the battery is wirelessly recharged by an external electric field.

9. A system for both sensing a single brain tremor and for providing deep brain stimulation (DBS) by a single treatment pulse, consisting of:

- a single cylindrical disc housing consisting of a single bottom and side edges extending from the bottom, and a single removable cylindrical top cover with perimeter sides which overlap the side edges of the housing, so that the cylindrical top cover covers a single compartment inside of the cylindrical disc housing, the single cylindrical disc housing having a height up to approximately 15 millimeters and a diameter of approximately 60 millimeters, the single cylindrical disc housing adapted to be mounted as a cap on a skull of a user, and not being implanted below the skull of the user;

- a single elongated lead line extending beneath a center portion of the bottom of the single cylindrical disc housing having a plurality of electrodes thereon, the electrodes each being used to both sense a single brain tremor and generate pulses therefrom, only the single elongated lead line being adapted to be implanted into only a single opening in a skull of a head of a user while the single cylindrical disc housing remains mounted as a cap on top of the skull of the user;

- a pulse stimulator inside of the compartment of the cylindrical disc housing, the pulse stimulator only becoming activated upon sensing of the single brain tremor from one of the electrodes on the single lead line, the pulse stimulator not generating any pulses when a tremor is not being sensed, the pulse stimulator includes a pulse generator for generating single spaced apart non-continuous pulses upon initial occurrence of the single brain tumor tremor sensed from the one electrode on the single implanted lead line;

- sensing detection circuitry solely located inside of the compartment of the cylindrical housing for sensing the single brain tremor from the one electrode on the single lead line;

- controls for the pulse stimulator that are all solely located inside of the compartment of the cylindrical disc housing, the controls for deactivating the pulse stimulator not generating any pulses when a tremor is not being sensed from the one electrode, the electrodes on the single lead line being connected to the controls for both sensing brain activity and for generating the single treatment pulse upon activation of the pulse stimulator, wherein the single treatment pulse is generated after the single brain tremor is detected by the one electrode, the one electrode sensing the brain tremor produced from the affected part of the brain is used for treating the affected part of the brain that produces the brain tremor, the controls include:

- a noise amplitude filter for filtering sensed signals from the electrodes,

- an impulse counter for counting impulses in the filtered signals in impulses per second;

- a trigger switch for triggering a single treatment pulse when a selected threshold of the counted impulses has been reached, wherein the controls shut off the noise amplitude filter and the impulse counter when the triggering of the single treatment pulse has been initiated, and

- an impulse firing threshold circuit for filtering out false alarms and allowing for a selected threshold value; and

- a battery power supply for the stimulator solely located inside the cylindrical disc housing for providing all power for the pulse stimulator and all power for the controls, wherein the controls control the battery power supply to supply power to the pulse stimulator only upon the sensing of the single tremor, and wherein the controls stop supplying power to the pulse stimulator after sensing detection circuitry has been turned on, wherein the cylindrical disc shaped housing eliminates implanting of any batteries in the chest area and the use of any other wires under the skin from the chest area to connect with the implanted electrodes, and wherein no other lead lines and no other wires are connected to the single cylindrical disc housing and the single implanted lead line extending below the single cylindrical disc housing, and the single cylindrical disc housing and the single implantable lead line extending below the single cylindrical disc housing, and only the single cylindrical disc housing and the single implantable lead line are used for both sensing and for providing deep brain stimulation (DBS), the single cylindrical disc housing being positioned as the
15 cap on top of the skull of the user and adapted to cover the cylindrical disc housing with the scalp.

10. The system of claim 9, wherein the noise amplitude filter is filtering sensed signals from the electrodes that are below approximately 20 milli-volts.

11. The system of claim 10, wherein the controls include:
   a sensing mode of the electrodes being approximately 20 to approximately 60 milli-volts; and
   a triggering range of the electrodes being approximately 2 to approximately 4 volts.

12. The system of claim 9, wherein the threshold value is approximately 5 impulses within approximately 50 milliseconds.

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