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(54) Title: CARDIAC CYCLE - SYNCHRONIZED NEURAL STIMULATOR

(57) Abstract: A neural stimulator senses a reference signal indicative of cardiac cycles each including a predetermined type timing reference event using a sensor external to the heart and blood vessels. The delivery of the neural stimulation pulses are synchronized to that timing reference event. Examples of the timing reference event include a predetermined cardiac event such as a P-wave or an R-wave detected from a subcutaneous ECG signal, a predetermined type heart sound detected from an acoustic signal, and a peak detected from a hemodynamic signal related to blood flow or pressure.
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CARDBIAC CYCLE - SYNCHRONIZED NEURAL STIMULATOR

CLAIM OF PRIORITY

Benefit of priority is hereby claimed to U.S. Patent Application Serial Number 11/099,141, filed on April 5, 2005, which application is herein incorporated by reference.

FIELD OF THE INVENTION

This document generally relates to neural stimulation systems and particularly to a system providing for synchronization of neural stimulation to cardiac cycles.

BACKGROUND

The heart is the center of a person’s circulatory system. The left portions of the heart draw oxygenated blood from the lungs and pump it to the organs of the body to provide the organs with their metabolic needs for oxygen. The right portions of the heart draw deoxygenated blood from the body organs and pump it to the lungs where the blood gets oxygenated. These pumping functions are accomplished by cyclic contractions of the myocardium (heart muscles). Each cycle, known as the cardiac cycle, includes systole and diastole. During systole, the heart ejects blood. During diastole, the heart is filled with blood for the next ejection (systolic) phase, and the myocardial tissue is perfused. In a normal heart, the sinoatrial node generates electrical impulses called action potentials. The electrical impulses propagate through an electrical conduction system to various regions of the heart to excite the myocardial tissue of these regions. Coordinated delays in the propagations of the action potentials in a normal electrical conduction system cause the various portions of the heart to contract in synchrony to result in efficient pumping functions indicated by a normal hemodynamic performance. A blocked or otherwise abnormal electrical conduction and/or deteriorated myocardial tissue result in systolic dysfunction – because the myocytes do not contract in unison – and diastolic dysfunction – because the myocytes do not relax in unison. Decreased systolic and diastolic performance each contribute to a poor overall hemodynamic performance, including a diminished blood supply to the heart and the rest of the body.
The hemodynamic performance is modulated by neural signals in portions of the autonomic nervous system. For example, the myocardium is innervated with sympathetic and parasympathetic nerves. Activities in these nerves, including artificially applied electrical stimuli, modulate the heart rate and contractility (strength of the myocardial contractions). Electrical stimulation applied to the sympathetic nerves is known to increase the heart rate and the contractility, shortening the systolic phase of a cardiac cycle, and lengthening the diastolic phase of the cardiac cycle. Electrical stimulation applied to the parasympathetic nerves is known to have essentially the opposite effects.

The ability of the electrical stimulation of the autonomic nerves in modulating the heart rate and contractility is utilized to treat abnormal cardiac conditions, such as to control myocardial remodeling and to prevent arrhythmias following myocardial infarction. It is observed that the effects of such electrical stimulation are dependent on timing of the delivery of electrical stimuli in relation to the cardiac cycle. Thus, it is desirable to synchronize the delivery of the electrical stimuli to the cardiac cycle. Because the electrical stimuli are delivered to portions of nerves external to the heart, there is a need for detecting a timing reference signal for synchronizing the delivery of the electrical stimuli to the cardiac cycle without intracardiac sensing.

**SUMMARY**

A neural stimulator senses a reference signal indicative of cardiac cycles each including a predetermined type timing reference event using a sensor external to the heart and blood vessels. The delivery of the neural stimulation pulses is synchronized to that timing reference event.

In one embodiment, a neural stimulation system includes a stimulation output circuit, a reference signal sensor, a reference event detection circuit, and a stimulation control circuit. The stimulation output circuit delivers neural stimulation pulses. The reference signal sensor senses a reference signal indicative of cardiac cycles each including a predetermined type timing reference event. The reference signal sensor may be placed in a site external to the circulatory system. The reference event detection circuit detects the predetermined type timing reference event. The stimulation control circuit controls the delivery of the neural stimulation pulses and includes a
synchronization module. The synchronization module synchronizes the delivery of the neural stimulation pulses to the predetermined type timing reference event.

In one specific embodiment, the neural stimulation system includes a stimulation output circuit, one or more electrodes, a cardiac event detection circuit, and a stimulation control circuit. The stimulation output circuit delivers neural stimulation pulses. The one or more electrodes sense an electrocardiographic (ECG) signal. The cardiac event detection circuit detects predetermined type cardiac events from the ECG signal. The stimulation control circuit controls the delivery of the neural stimulation pulses and includes a synchronization module. The synchronization module synchronizes the delivery of the neural stimulation pulses to the predetermined type cardiac events.

In another specific embodiment, the neural stimulation system includes a stimulation output circuit, an acoustic sensor, a heart sound detection circuit, and a stimulation control circuit. The stimulation output circuit delivers neural stimulation pulses. The implantable acoustic sensor senses an acoustic signal indicative of heart sounds. The heart sound detection circuit detects predetermined type heart sounds using the acoustic signal. The stimulation control circuit controls the delivery of the neural stimulation pulses and includes a synchronization module. The synchronization module synchronizes the delivery of the neural stimulation pulses to the predetermined type heart sounds.

In another specific embodiment, the neural stimulation system includes a stimulation output circuit, a hemodynamic sensor, a hemodynamic event detection circuit, and a stimulation control circuit. The stimulation output circuit delivers neural stimulation pulses. The hemodynamic sensor senses a hemodynamic signal. The hemodynamic event detection circuit detects a predetermined type hemodynamic event using the hemodynamic signal. The stimulation control circuit controls the delivery of the neural stimulation pulses and includes a synchronization module. The synchronization module synchronizes the delivery of the neural stimulation pulses to the predetermined type hemodynamic event.

In one embodiment, a method for neural stimulation is provided. A timing reference signal is sensed using a reference signal sensor placed external to the circulatory system. The timing reference signal is indicative of cardiac
cycles each including a predetermined type timing reference event. The predetermined type timing reference event is detected from the reference signal. A delivery of neural stimulation pulses is synchronized to the detected timing reference event.

This Summary is an overview of some of the teachings of the present application and not intended to be an exclusive or exhaustive treatment of the present subject matter. Further details about the present subject matter are found in the detailed description and appended claims. Other aspects of the invention will be apparent to persons skilled in the art upon reading and understanding the following detailed description and viewing the drawings that form a part thereof. The scope of the present invention is defined by the appended claims and their legal equivalents.

BRIEF DESCRIPTION OF THE DRAWINGS

In the drawings, which are not necessarily drawn to scale, like numerals describe similar components throughout the several views. The drawings illustrate generally, by way of example, but not by way of limitation, various embodiments discussed in the present document.

FIG. 1 is an illustration of an embodiment of a neural stimulation system and portions of an environment in which the neural stimulation system is used.

FIG. 2 is a block diagram illustrating an embodiment of a circuit of a cardiac cycle-synchronized neural stimulation system.

FIG. 3 is a block diagram illustrating a specific embodiment of the circuit of FIG. 2.

FIG. 4 is a block diagram illustrating an embodiment of a circuit using a wireless ECG to synchronize neural stimulation to cardiac cycles.

FIG. 5 is an illustration of an embodiment of an electrode system for subcutaneous ECG sensing.

FIG. 6 is a block diagram illustrating an embodiment of a circuit using heart sounds to synchronize neural stimulation to cardiac cycles.

FIG. 7 is a block diagram illustrating an embodiment of a circuit using a hemodynamic signal to synchronize neural stimulation to cardiac cycles.

FIG. 8 is a flow chart illustrating an embodiment of a method for synchronizing neural stimulation to cardiac cycles.
DETAILED DESCRIPTION

In the following detailed description, reference is made to the accompanying drawings which form a part hereof, and in which is shown by way of illustration specific embodiments in which the invention may be practiced. These embodiments are described in sufficient detail to enable those skilled in the art to practice the invention, and it is to be understood that the embodiments may be combined, or that other embodiments may be utilized and that structural, logical and electrical changes may be made without departing from the scope of the present invention. The following detailed description provides examples, and the scope of the present invention is defined by the appended claims and their legal equivalents.

It should be noted that references to “an”, “one”, or “various” embodiments in this disclosure are not necessarily to the same embodiment, and such references contemplate more than one embodiment.

This document discusses a neural stimulation system that synchronizes the delivery of neural stimulation pulses to cardiac cycles. The neural stimulation system includes an implantable neural stimulator that senses a reference signal indicative of cardiac cycles each including a predetermined type timing reference event using an implantable reference event sensor. The implantable reference event sensor is an extracardiac and extravascular sensor, i.e., a sensor that is placed external to a patient’s circulatory system including the heart and blood vessels. The delivery of the neural stimulation pulses are synchronized to the timing reference event. Examples of the reference signal include a wireless ECG, an acoustic signal indicative of heart sounds, and a hemodynamic signal.

In this document, “Surface ECG” refers to a cardiac electrical signal sensed with electrodes attached onto the exterior surface of the skin. “Wireless ECG” refers to a signal approximating the surface ECG, acquired without using surface (non-implantable, skin contact) electrodes. “Subcutaneous ECG” is a form of wireless ECG and includes a cardiac electrical signal sensed through electrodes implanted in subcutaneous tissue, such as through electrodes incorporated onto an implantable medical device that is subcutaneously implanted. As reflected in their corresponding morphologies, the surface ECG results from electrical activities of the entire heart. The wireless ECG, including
but not being limited to the subcutaneous ECG, has a morphology that
approximates that of the surface ECG and reflects electrical activities of a
substantial portion of the heart, up to the entire heart.

In this document, an “acoustic signal” includes any signal indicative of
heart sounds. “Heart sounds” include audible mechanical vibrations caused by
cardiac activity that can be sensed with a microphone and audible and inaudible
mechanical vibrations caused by cardiac activity that can be sensed with an
accelerometer. Known type heart sounds include the “first heart sound” or S1,
the “second heart sound” or S2, the “third heart sound” or S3, the “fourth heart
sound” or S4, and their various sub-components. S1 is known to be indicative
of, among other things, mitral valve closure, tricuspid valve closure, and aortic
valve opening. S2 is known to be indicative of, among other things, aortic valve
closure and pulmonary valve closure. S3 is known to be a ventricular diastolic
filling sound often indicative of certain pathological conditions including heart
failure. S4 is known to be a ventricular diastolic filling sound resulted from
atrial contraction and is usually indicative of pathological conditions. The term
“heart sound” hereinafter refers to any heart sound (e.g., S1) and any
components thereof (e.g., M1 component of S1, indicative of Mitral valve
closure).

In this document, a “hemodynamic signal” includes a signal providing
for monitoring, calculation, or estimation of one or more measures of
hemodynamic performance such as blood pressure or pressure-related
parameters, cardiac output, stroke volume, volume of blood flow, change in
(e.g., derivative of) the volume of blood flow, and/or velocity of blood flow.

FIG. 1 is an illustration of an embodiment of a neural stimulation system
100 and portions of an environment in which system 100 is used. System 100
includes implantable medical device 110 that delivers neural stimulation pulses
through leads 106 and 108, an external system 120, and a telemetry link 125
providing for communication between implantable medical device 110 and
external system 120. For illustrative purpose only, FIG. 1 shows that lead 106
includes an electrode 107 coupled to a nerve 102 of the sympathetic nervous
system, and lead 108 includes an electrode 109 coupled a nerve 104 of the
parasympathetic nervous system. Nerves 102 and 104 innervate a heart 101. In
various embodiments, implantable medical device 110 provides neural
stimulation to any one or more nerves through one or more leads for modulating one or more functions of the circulatory system including heart 101. Such leads include implantable neural leads each including at least one electrode for sensing neural activities and/or delivering neural stimulation pulses. One example of such an electrode includes a cuff electrode for placement around an aortic, carotid, or vagus nerve.

Implantable medical device 110 delivers the neural stimulation pulses and includes a cardiac cycle-synchronized neural stimulation circuit 130. Cardiac cycle-synchronized neural stimulation circuit 130 detects a predetermined type timing reference event from a cardiac cycle and synchronizes the delivery of neural stimulation pulses to that timing reference event. In one embodiment, cardiac cycle-synchronized neural stimulation circuit 110 starts a predetermined offset time interval upon detection of the timing reference event and delivers a burst of neural stimulation pulses when the offset time interval expires. In one embodiment, implantable medical device 110 is capable of monitoring physiologic signals and/or delivering therapies in addition to the neural stimulation. Examples of such additional therapies include cardiac pacing therapy, cardioversion/defibrillation therapy, cardiac resynchronization therapy, cardiac remodeling control therapy, drug therapy, cell therapy, and gene therapy. In various embodiments, implantable medical device 110 delivers the neural stimulation in coordination with one or more such additional therapies.

External system 120 provides for control of and communication with implantable medical device 110 by a physician or other caregiver. In one embodiment, external system 120 includes a programmer. In another embodiment, external system 120 is a patient management system including an external device communicating with implantable medical device 110 via telemetry link 125, a remote device in a relatively distant location, and a telecommunication network linking the external device and the remote device. The patient management system allows access to implantable medical device 110 from a remote location, for purposes such as monitoring patient status and adjusting therapies. In one embodiment, telemetry link 125 is an inductive telemetry link. In an alternative embodiment, telemetry link 125 is a far-field radio-frequency (RF) telemetry link. Telemetry link 125 provides for data transmission from implantable medical device 110 to external system 120. This
includes, for example, transmitting real-time physiological data acquired by implantable medical device 110, extracting physiological data acquired by and stored in implantable medical device 110, extracting patient history data such as occurrences of arrhythmias and therapy deliveries recorded in implantable medical device 110, and/or extracting data indicating an operational status of implantable medical device 110 (e.g., battery status and lead impedance). Telemetry link 125 also provides for data transmission from external system 120 to implantable medical device 110. This includes, for example, programming implantable medical device 110 to acquire physiological data, programming implantable medical device 110 to perform at least one self-diagnostic test (such as for a device operational status), and/or programming implantable medical device 110 to deliver one or more therapies and/or to adjust the delivery of one or more therapies.

FIG. 2 is a block diagram illustrating an embodiment of a circuit of a cardiac cycle-synchronized neural stimulation system 231. System 231 includes a reference signal sensor 215 and cardiac cycle-synchronized neural stimulation circuit 130.

Reference signal sensor 215 senses a reference signal indicative of cardiac cycles each including a predetermined type timing reference event. In one embodiment, reference signal sensor 215 is an implantable reference signal sensor. The timing reference event is a recurring feature of the cardiac cycle that is chosen to be a timing reference to which the neural stimulation is synchronized. In one embodiment, reference signal sensor 215 is configured for extracardiac and extravascular placement, i.e., placement external to the heart and blood vessels. Examples of reference signal sensor 215 include a set of electrodes for sensing a subcutaneous ECG signal, an acoustic sensor for sensing an acoustic signal indicative of heart sounds, and a hemodynamic sensor for sensing a hemodynamic signal indicative of hemodynamic performance. In one embodiment, implantable medical device 110 has an implantable housing that contains both a reference signal sensor 215 and cardiac cycle-synchronized neural stimulation circuit 130. In another embodiment, reference signal sensor 215 is incorporated onto the housing of implantable medical device 110. In another embodiment, reference signal sensor 215 is electrically connected to implantable medical device 110 through one or more leads. In another
embodiment, reference signal sensor 215 is communicatively coupled to implantable medical device 110 via an intra-body telemetry link.

Cardiac cycle-synchronized neural stimulation circuit 130 includes a stimulation output circuit 232, a reference event detection circuit 234, and a stimulation control circuit 236. Reference event detection circuit 234 receives the reference signal from reference signal sensor 215 and detects the timing reference event from the reference signal. Stimulation control circuit 236 controls the delivery of the neural stimulation pulses and includes a synchronization module 238. Synchronization module 238 receives a signal indicative of the detection of each timing reference event and synchronizes the delivery of the neural stimulation pulses to the detected timing reference event. Stimulation output circuit 232 delivers neural stimulation pulses upon receiving a pulse delivery signal from stimulation control circuit 236.

FIG. 3 is a block diagram illustrating an embodiment of a circuit of a cardiac cycle-synchronized neural stimulation system 331, which is a specific embodiment of system 231. System 331 includes reference signal sensor 215 and a cardiac cycle-synchronized neural stimulation circuit 330, which is a specific embodiment of cardiac cycle-synchronized neural stimulation circuit 130. Cardiac cycle-synchronized neural stimulation circuit 330 includes stimulation output circuit 232, a reference event detection circuit 334, and a stimulation control circuit 336.

Reference event detection circuit 334 is a specific embodiment of reference event detection 234 and includes a signal processor 342 and an event detector 344. Signal processor 342 receives the reference signal sensed by reference signal sensor 215 and processes the reference signal in preparation for the detection of the timing reference events by event detector 344. Event detector 344 includes a comparator having an input to receive the processed reference signal, another input to receive a detection threshold, and an output producing a detection signal indicating a detection of the timing reference signal.

In one embodiment, signal processor 342 processes the reference signal to provide for extraction of the timing reference event based on a single cardiac cycle. In one specific embodiment, signal processor 342 includes a filter having a pass-band corresponding to a frequency range of the timing reference event to prevent unwanted activities in the reference signal from being detected by event
detector 344. In another specific embodiment, signal processor 342 includes a blanking period generator to generate a blanking period that blanks the unwanted activities in the reference signal. This approach is applied when an approximate timing relationship between the timing reference event and the unwanted activities, or an approximate timing relationship between another detectable event and the unwanted activities, is predictable. In another specific embodiment, the blanking period generator generates a blanking period that blanks cardiac pacing artifacts in the reference signal, i.e., unwanted activities caused by delivery of cardiac pacing pulses. In another specific embodiment, signal processor 342 includes a timing interval generator to generate a timing interval between an intermediate event and the timing reference event. This approach is applied when the intermediate event is more easily detectable than the timing reference event and when an approximate timing relationship between the intermediate event and the timing reference event is predictable. In another embodiment, signal processor 342 processes the reference signal to provide for extraction of the timing reference event based on a plurality of cardiac cycles. In one specific embodiment, signal processor 342 includes a signal averaging circuit that averages the reference signal over a predetermined number of cardiac cycles before the detection of the timing reference event by event detector 344.

Stimulation control circuit 336 is a specific embodiment of stimulation control circuit 236 and includes a synchronization circuit 338, an offset interval generator 339, and a pulse delivery controller 340. Synchronization circuit 338 includes one or both of a continuous synchronization module 346 and a periodic synchronization module 348. Continuous synchronization module 346 synchronizes the delivery of the neural stimulation pulses to the timing reference event of consecutive cardiac cycles. Periodic synchronization module 348 synchronizes the delivery of the neural stimulation pulses to the timing reference event of selected cardiac cycles on a periodic basis. Offset interval generator 339 produces an offset interval starting with the detected timing reference event. Pulse delivery controller 340 sends the pulse delivery signal to start a delivery of a burst of a plurality of neural stimulation pulses when the offset interval expires. In one embodiment, pulse delivery controller 340 sends the pulse delivery signal after the detection of the timing reference event for each of consecutive cardiac cycles. In another embodiment, pulse delivery controller
340 sends the pulse delivery signal after the detection of the timing reference event for selected cardiac cycles according to a predetermined pattern or schedule, such as on a periodic basis.

FIG. 4 is a block diagram illustrating an embodiment of a cardiac cycle-synchronized neural stimulation system 431, which is a specific embodiment of system 231 and uses a wireless ECG to synchronize neural stimulation to cardiac cycles. System 431 includes ECG electrodes 415 and a cardiac cycle-synchronized neural stimulation circuit 430, which is a specific embodiment of cardiac cycle-synchronized neural stimulation circuit 230. Cardiac cycle-synchronized neural stimulation circuit 430 includes stimulation output circuit 232, a cardiac event detection circuit 434, an arrhythmia detection circuit 452, a cardiac parameter measurement circuit 454, and a stimulation control circuit 436.

In one embodiment, ECG electrodes 415 include surface ECG electrodes. In another embodiment, ECG electrodes 415 include electrodes for sensing a wireless ECG signal. In one embodiment, ECG electrodes 415 include subcutaneous electrodes for sensing a subcutaneous ECG signal. In one embodiment, the subcutaneous electrodes are incorporated onto the implantable medical device 110, which is to be subcutaneously implanted. Examples of such subcutaneous electrodes are discussed below with reference to FIG. 5. In one embodiment, at least one subcutaneous electrode is placed in a selected location in the body near the base of the heart to allow selective detection of atrial depolarizations (P-waves). In another embodiment, multiple subcutaneous electrodes are placed near base and apex of the heart to allow P-wave detection by subtracting out unwanted activities including ventricular depolarizations (R-waves). This approach applies when it is difficult to isolate P-waves by selecting electrode sites and filtering. At least one subcutaneous electrode is placed near the apex of the heart to allow detection of R-waves. The detected R-waves are then used to isolate, by subtraction, P-waves from a subcutaneous ECG signal that includes both P-waves and R-waves.

Cardiac event detection circuit 434 is a specific embodiment of reference event detection circuit 234. In one embodiment, cardiac event detection circuit 434 includes a signal processor such as signal processor 342 and an event detector such as event detector 344. The signal processor includes a wireless
ECG sensing circuit to amplify and filter the subcutaneous ECG signal sensed through ECG electrodes 415. An example of electrodes and a circuit for sensing wireless ECG signals including subcutaneous ECG signals is discussed in U.S. Patent Application Serial No. 10/795,126, entitled "WIRELESS ECG IN

IMPLANTABLE DEVICES," filed on March 5, 2004, assigned to Cardiac Pacemakers, Inc., which is incorporated herein by reference in its entirety. In one embodiment, as illustrated in FIG. 4, the timing reference event is a P-wave. Cardiac event detection circuit 434 includes a P-wave detector 450 to detect P-waves from the wireless ECG signal. In one specific embodiment, P-wave detector 450 includes a filter having a pass-band corresponding to a frequency range of P-waves. In another specific embodiment, P-wave detector 450 includes an R-wave detector to detect R-waves from one subcutaneous signal and a blanking period generator to generate blanking periods to blank unwanted activities including the R-waves in another wireless ECG signal. In another specific embodiment, P-wave detector 450 includes an R-wave detector to detect R-waves from the subcutaneous signal and a timing interval generator to generate a timing interval upon detection of each R-wave. A P-wave is estimated to occur at the end of the timing interval.

Arrhythmia detection circuit 452 and cardiac parameter measurement circuit 454 provide for control of neural stimulation based on cardiac conditions. Arrhythmia detection circuit 452 detects one or more types of arrhythmia from the wireless ECG signal. Cardiac parameter measurement module 454 measures one or more cardiac parameters such as a heart rate and an atrioventricular interval from the wireless ECG signal.

Stimulation control circuit 436 is a specific embodiment of stimulation control circuit 336 and includes a synchronization module 438. Synchronization module 438 synchronizes the delivery of the neural stimulation pulses to the detected cardiac events such as P-waves. In one embodiment, stimulation control circuit 436 includes elements corresponding to those of stimulation circuit 336, including offset interval generator 339 and pulse delivery controller 340. Synchronization circuit 438 includes one or both of a continuous synchronization module to synchronize the delivery of the neural stimulation pulses to the P-wave of each of consecutive cardiac cycles and a periodic synchronization module to synchronize the delivery of the neural stimulation.
pulses to the P-wave of each of selected cardiac cycles on a periodic basis. The offset interval generator produces an offset interval starting with each detected P-wave. The pulse delivery controller sends the pulse delivery signal to start a delivery of a burst of a plurality of neural stimulation pulses when the offset interval expires. In one embodiment, the pulse delivery controller sends the pulse delivery signal after the detection of the P-wave for each of consecutive cardiac cycles. In another embodiment, the pulse delivery controller sends the pulse delivery signal after the detection of the P-wave for each of selected cardiac cycles according to a predetermined pattern or schedule, such as on a periodic basis.

In one embodiment, stimulation control circuit 436 also controls the delivery of the neural stimulation pulses based on the cardiac rhythm detected by arrhythmia detection circuit 452 and/or the cardiac parameters measured by cardiac parameter measurement circuit 454. In one embodiment, stimulation control circuit 436 withholds or adjusts the delivery of the neural stimulation pulses when an arrhythmia is detected. In another embodiment, stimulation control circuit 436 starts, stops, or adjusts the delivery of the neural stimulation pulses based on the measured cardiac parameter, such as the heart rate and the atrioventricular interval.

FIG. 5 is an illustration of an embodiment of an electrode system for sensing one or more subcutaneous ECG signals. An electrode system for subcutaneous ECG sensing includes two or more implantable electrodes. These implantable electrodes are selected from the electrodes including, but not being limited to, those illustrated in FIG. 5. The electrodes are selected to allow for sensing electrical activities from a substantial portion of the heart, up to the entire heart. FIG. 5 shows an implantable medical device 510, which is a specific embodiment of implantable medical device 110, and electrodes incorporated onto that device. Implantable medical device 510 is to be subcutaneously implanted in a patient in need of neural stimulation to modulate cardiac functions. In one embodiment, ECG electrodes 415 include one or more electrodes shown in FIG. 5. In another embodiment, in addition to one or more electrodes shown in FIG. 5, ECG electrodes 415 include one or more electrodes each electrically connected to implantable medical device 510 through a lead.
Implantable medical device 510 includes a hermetically sealed can 511 to house its circuit. Can 511 has an outer surface subject to contact with body tissue. Can 511 includes or provides for a base of a can electrode 514 that is selectable as one of the electrodes for sensing a subcutaneous ECG signal. At least a portion of the outer surface of can 511 is made of electrically conductive material. In one embodiment, can 511 is used as can electrode 514. In one specific embodiment, can electrode 514 includes at least one conductive portion of can 511. In another embodiment, can electrode 514 is incorporated onto the outer surface of can 511 and is electrically insulated from any conductive portion of can 511 using a non-conductive layer. In one specific embodiment, a hermetically sealed feedthrough including a conductor provides for an electrical connection between can electrode 514 and the circuit housed in can 511.

A header 512 is attached to can 511 and includes connectors providing for electrical access to the circuit housed in can 511. In one embodiment, one or more of header electrodes 516A-B are incorporated into the header. Header electrodes 516A-B are each selectable as one of the electrodes for sensing a subcutaneous ECG signal.

In one embodiment, two or more concentric electrodes 517A-C are incorporated onto the outer surface of can 511. Each of the concentric electrodes 517A-C is selectable as one of the electrodes for sensing a subcutaneous ECG signal. Concentric electrodes 517A-C are insulated from the conductive portion of can 511 with a non-conductive layer and connected to the circuit housed in can 511 via hermetically sealed feedthroughs. In one embodiment, two electrodes, including an inner electrode and an outer electrode, are selected from concentric electrodes 517A-C for the wireless ECG sensing. In one embodiment, the outer electrode has a ring shape. In another embodiment, the outer electrode has a shape approaching the contour of can 511.

In one embodiment, implantable medical device 510 includes an antenna 513 used for a far-field RF telemetry link providing for communication between implantable medical device 510 and external system 120. Antenna 513 is electrically connected to the circuit housed in can 511. In one embodiment, antenna 513 projects from header 512 and extends along one side of can 511. In one embodiment, antenna 513 includes a metal conductor with a distal portion.
exposed for functioning as an antenna electrode 518, which is selectable as one of the electrodes for sensing a subcutaneous ECG signal.

It is to be understood that the electrodes illustrated in FIG. 5 are intended to be examples but not limitations. Other electrode configurations are usable as long as they provide for sensing of surface ECG signals or signals that approximate the surface ECG or otherwise allows for detection of a timing reference signal for synchronizing the delivery of neural stimulation pulses to cardiac cycles. In various embodiments in which multiple subcutaneous ECG vectors are sensed, multiple pairs of electrodes are selected, simultaneously or one at a time, for a multi-channel (multi-vector) subcutaneous ECG sensing. In one specific embodiment, one or more of subcutaneous ECG vectors are sensed to approximate one or more vectors of a standard multi-lead surface ECG recording. In another specific embodiment, multiple subcutaneous ECG vectors are sensed based on needs of specific information for synchronizing the delivery of neural stimulation pulses to cardiac cycles. Such subcutaneous ECG vectors do not necessarily approximate standard surface ECG vectors. In one specific embodiment, implantable medical device 510 includes header electrodes 516A-B and can electrode 514 for the subcutaneous ECG sensing. Implantable medical device 510 is programmable for sensing subcutaneous ECG vectors between (1) header electrodes 516A and 516B, (2) header electrode 516A and can electrode 514, and/or (3) header electrode 516B and can electrode 514. In another specific embodiment, implantable medical device 510 includes one of header electrodes 516A-B, antenna electrode 518, and can electrode 514 for the subcutaneous ECG sensing. Implantable medical device 510 is programmable for sensing subcutaneous ECG vectors between (1) header electrode 516A or 516B and antenna electrode 518, (2) header electrode 516A or 516B and can electrode 514, and/or (3) antenna electrode 518 and can electrode 514. In another specific embodiment, implantable medical device 510 includes header electrodes 516A-B, antenna electrode 518, and can electrode 514 for the subcutaneous ECG sensing. Implantable medical device 510 is programmable for sensing subcutaneous ECG vectors between (1) header electrodes 516A and 518, (2) header electrode 516A and antenna electrode 518, (3) header electrode 516A and can electrode 514, (4) header electrode 516B and antenna electrode 518, (5) header electrode 516B and can electrode 514, and/or (6) antenna electrode 518.
and can electrode 514. Other specific embodiments involving any electrode combinations for the subcutaneous ECG sensing will be employed based on needs and consideration for synchronizing the delivery of neural stimulation pulses to cardiac cycles as well as needs and considerations for performing other diagnostic and/or therapeutic functions provided by implantable medical device 510.

The selection of subcutaneous ECG vectors depends on the purpose for the subcutaneous ECG sensing. When the subcutaneous ECG signal is sensed for detecting P-waves, the subcutaneous ECG vector that provide for a reliable P wave detection are selected. When the subcutaneous ECG signal is sensed for detecting R-waves, one or more subcutaneous ECG vectors that provide for a reliable R wave detection are selected. In one embodiment, when more than one subcutaneous ECG vector provides for a reliable sensing for a particular purpose, the subcutaneous ECG vector showing the highest signal-to-noise ratio (SNR) for that purpose is selected. For example, if the subcutaneous ECG is sensed for detecting P waves, the subcutaneous ECG vector showing the highest SNR with P waves being considered as the signal that is selected.

FIG. 6 is a block diagram illustrating an embodiment of a cardiac cycle-synchronized neural stimulation system 631, which is a specific embodiment of system 231 and uses heart sounds to synchronize neural stimulation to cardiac cycles. System 631 includes an acoustic sensor 615 and a cardiac cycle-synchronized neural stimulation circuit 630, which is a specific embodiment of cardiac cycle-synchronized neural stimulation circuit 230. Cardiac cycle-synchronized neural stimulation circuit 630 includes stimulation output circuit 232, a heart sound detection circuit 634, and a stimulation control circuit 636.

Acoustic sensor 615 senses an acoustic signal indicative heart sounds. In one embodiment, acoustic sensor 615 includes an implantable acoustic sensor. In one embodiment, acoustic sensor 615 includes an accelerometer. In another embodiment, acoustic sensor 615 includes a microphone. In one specific embodiment, acoustic sensor 615 is included in implantable medical device 110. In another specific embodiment, acoustic sensor 615 is incorporated onto a lead connected to implantable medical device 110.

Heart sound detection circuit 634 detects predetermined type heart sounds from the acoustic signal. Heart sound detection circuit 634 includes one
or more of a first heart sound (S1) detector to detect S1, a second heart sound (S2) detector to detect S2, a third heart sound (S3) detector to detect S3, and a fourth heart sound (S4) detector to detect S4. In one embodiment, the type of heart sounds to be detected is determined based on whether each particular type of heart sounds is consistently recurring and reliably detectable in an individual patient. In one embodiment, cardiac event detection circuit 634 includes a signal processor such as signal processor 342 and an event detector such as event detector 344. In one specific embodiment, heart sound detection circuit 634 includes a filter having a pass-band corresponding to a frequency range of the predetermined type heart sounds. In another specific embodiment, heart sound detection circuit 634 includes a signal averaging circuit to average the acoustic signal over a predetermined number of cardiac cycles before the detection of the predetermined type heart sounds. In another specific embodiment, heart sound detection circuit 634 receives an activity signal indicative of the patient’s gross physical activity level and stops detecting heart sounds while the activity signal exceeds a predetermined threshold activity level. In another embodiment, heart sound detection circuit 634 includes an S2 detector and/or an S3 detector such as those discussed in U.S. Patent Application Serial No. 10/746,853, "METHOD AND APPARATUS FOR THIRD HEART SOUND DETECTION," filed on December 24, 2004, assigned to Cardiac Pacemakers, Inc., which is incorporated by reference in its entirety.

Stimulation control circuit 636 is a specific embodiment and includes a synchronization module 638. Synchronization module 638 synchronizes the delivery of the neural stimulation pulses to the predetermined type heart sounds. In one embodiment, stimulation control circuit 636 includes elements corresponding to those of stimulation circuit 336, including offset interval generator 339 and pulse delivery controller 340. Synchronization circuit 638 includes one or both of a continuous synchronization module to synchronize the delivery of the neural stimulation pulses to the predetermined type heart sound of each of consecutive cardiac cycles and a periodic synchronization module to synchronize the delivery of the neural stimulation pulses to the predetermined type heart sound of each of selected cardiac cycles on a periodic basis. The offset interval generator produces an offset interval starting with the detected predetermined type heart sound. The pulse delivery controller sends the pulse
delivery signal to start a delivery of a burst of a plurality of neural stimulation pulses when the offset interval expires. In one embodiment, the pulse delivery controller sends the pulse delivery signal after the detection of the predetermined type heart sound for each of consecutive cardiac cycles. In another embodiment, the pulse delivery controller sends the pulse delivery signal after the detection of the predetermined type heart sound for each of selected cardiac cycles according to a predetermined pattern or schedule, such as on a periodic basis.

FIG. 7 is a block diagram illustrating an embodiment of a cardiac cycle-synchronized neural stimulation system 731, which is a specific embodiment of system 231 and uses a hemodynamic signal to synchronize neural stimulation to cardiac cycles. System 731 includes a hemodynamic sensor 715 and a cardiac cycle-synchronized neural stimulation circuit 730, which is a specific embodiment of cardiac cycle-synchronized neural stimulation circuit 230. Cardiac cycle-synchronized neural stimulation circuit 730 includes stimulation output circuit 232, a hemodynamic event detection circuit 734, and a stimulation control circuit 736.

Hemodynamic sensor 715 senses a hemodynamic signal indicative of hemodynamic performance, such as a signal indicative of blood pressure or flow. In one embodiment, hemodynamic sensor 715 is an implantable hemodynamic sensor. In one embodiment, hemodynamic sensor 715 includes a Doppler echocardiographic transducer to sense a peripheral blood flow. In another embodiment, hemodynamic sensor 715 includes a pressure sensor to sense a central or peripheral blood pressure. In another embodiment, hemodynamic sensor 715 includes a pulse oximeter to sense an oximetry signal, which is a plethysmographic signal indicative of blood flow.

Hemodynamic event detection circuit 734 detects predetermined type hemodynamic events from the hemodynamic signal. The hemodynamic events correspond to a recurring feature of the cardiac cycle that is chosen to be a timing reference to which the neural stimulation is synchronized. In one embodiment, hemodynamic event detection circuit 734 includes a peak detector that detects predetermined type peaks in the hemodynamic signal. In one specific embodiment, the peak detector is a pressure peak detector that detects predetermined type peaks in a blood pressure signal. In another specific embodiment, the peak detector includes a flow peak detector that detects
predetermined type peaks in a blood flow signal. The predetermined type peaks are peaks indicative of a characteristic event that occurs during each cardiac cycle. In another embodiment, cardiac cycle-synchronized neural stimulation circuit 730 includes a derivative calculator to produce a derivative hemodynamic signal by calculating a time derivative of the hemodynamic signal.

Hemodynamic event detection circuit 734 detects the predetermined type hemodynamic event from the derivative hemodynamic signal. In one embodiment, the peak detector detects predetermined type peaks in the derivative hemodynamic signal. In one specific embodiment, the peak detector is a pressure change peak detector that detects predetermined type peaks in a derivative hemodynamic signal indicative of changes in the blood pressure (e.g., dP/dt). In another specific embodiment, the peak detector includes a flow change peak detector that detects predetermined type peaks in a derivative hemodynamic signal indicative changes in the blood flow.

Stimulation control circuit 736 is a specific embodiment and includes a synchronization module 738. Synchronization module 738 synchronizes the delivery of the neural stimulation pulses to the predetermined type hemodynamic events. In one embodiment, stimulation control circuit 736 includes elements corresponding to those of stimulation circuit 336, including offset interval generator 339 and pulse delivery controller 340. Synchronization circuit 738 includes one or both of a continuous synchronization module to synchronize the delivery of the neural stimulation pulses to the predetermined type hemodynamic event of each of consecutive cardiac cycles and a periodic synchronization module to synchronize the delivery of the neural stimulation pulses to the predetermined type hemodynamic event of each of selected cardiac cycles on a periodic basis. The offset interval generator produces an offset interval starting with each detected predetermined type hemodynamic event. The pulse delivery controller sends the pulse delivery signal to start a delivery of a burst of a plurality of neural stimulation pulses when the offset interval expires. In one embodiment, the pulse delivery controller sends the pulse delivery signal after the detection of the predetermined type hemodynamic event for each of consecutive cardiac cycles. In another embodiment, the pulse delivery controller sends the pulse delivery signal after the detection of the predetermined type
hemodynamic event for each of selected cardiac cycles according to a predetermined pattern or schedule, such as on a periodic basis.

FIG. 8 is a flow chart illustrating an embodiment of a method for synchronizing neural stimulation to cardiac cycles. In one embodiment, the method is performed by cardiac cycle-synchronized neural stimulation system 231, including any of its specific embodiments or any combination of its specific embodiments discussed above.

A reference signal is sensed at 800. The reference signal is indicative of cardiac cycles each including a predetermined type timing reference event. In one embodiment, the reference signal is sensed using an implantable sensor placed external to the circulatory system. Examples of the reference signal include a cardiac signal such as a subcutaneous ECG signal, an acoustic signal indicative of heart sounds, and a hemodynamic signal such as a blood pressure or flow signal.

The predetermined type timing reference event is detected at 810. In one embodiment, the reference signal is processed to allow or to facilitate the detection of the predetermined type timing reference event. In one specific embodiment, the predetermined type timing reference event is detected based on the reference signal sensed over a single cardiac cycle. In another embodiment, the predetermined type timing reference event is detected based on the reference signal sensed over a plurality of cardiac cycles. Examples of such processing include filtering, blanking unwanted activities from the reference signal, detecting an intermediate event having an approximately predictable timing relationship with the predetermined type timing reference event, and averaging the reference signal over a plurality of cardiac cycles. Examples of the predetermined type timing reference event include P-wave and R-wave detected from the cardiac signal such as the subcutaneous ECG signal, a predetermined type heart sound from the acoustic signal, and a point of peak amplitude or any other morphologically distinctive point in the hemodynamic signal such as the pressure or flow signal.

A delivery of neural stimulation pulses is synchronized to the predetermined type timing reference event at 820. In one embodiment, the delivery of the neural stimulation pulses is synchronized to the predetermined type timing reference event of each of consecutive cardiac cycles on a
continuous basis. In another embodiment, the delivery of the neural stimulation pulses is synchronized to the predetermined type timing reference event of each of selected cardiac cycles on a periodic basis. In one embodiment, a burst of neural stimulation pulses is delivered at the end of an offset interval starting with the predetermined type timing reference event. In one embodiment, the burst of neural stimulation pulses is delivered after the predetermined type timing reference event for each cardiac cycle of consecutive cardiac cycles. In another embodiment, the burst of neural stimulation pulses is delivered after the predetermined type timing reference event for each cardiac cycle of selected cardiac cycles according to a predetermined pattern or schedule, such as on a period basis.

In one embodiment, the delivery of the neural stimulation pulses is further controlled by the patient's cardiac condition and/or activity level. The patient's cardiac rhythm and one or more cardiac parameters indicative of the cardiac functions are monitored. In one embodiment, the delivery of the neural stimulation pulses is controlled based on the cardiac rhythm. In response to a detected arrhythmia, the delivery of the neural stimulation pulses is withheld or adjusted. In another embodiment, the delivery of the neural stimulation pulses is adjusted or optimized based on the one or more cardiac parameters. Examples of such cardiac parameters include heart rate, atrioventricular intervals, and interventricular intervals. The timing for the delivery of the neural stimulation pulses is adjusted, for example, for a desirable heart rate, an atrioventricular interval corresponding to a desirable hemodynamic performance, and/or a minimum interventricular interval.

It is to be understood that the above detailed description is intended to be illustrative, and not restrictive. Other embodiments will be apparent to those of skill in the art upon reading and understanding the above description. The scope of the invention should, therefore, be determined with reference to the appended claims, along with the full scope of legal equivalents to which such claims are entitled.
What is claimed is:

1. A neural stimulation system coupled to a living subject having a circulatory system, the neural stimulation system comprising:
   a stimulation output circuit to deliver neural stimulation pulses;
   an implantable reference signal sensor to sense a reference signal indicative of cardiac cycles each including a predetermined type timing reference event, the implantable reference signal sensor configured to be placed external to the circulatory system;
   a reference event detection circuit coupled to the implantable reference signal sensor, the reference event detection circuit adapted to detect the predetermined type timing reference events; and
   a stimulation control circuit coupled to the stimulation output circuit and the reference event detection circuit, the stimulation control circuit adapted to control the delivery of the neural stimulation pulses and including a synchronization module adapted to synchronize the delivery of the neural stimulation pulses to the predetermined type timing reference events.

2. The neural stimulation system according to claim 1, further comprising an implantable housing adapted to contain at least the stimulation output circuit, the reference event detection circuit, and the stimulation control circuit.

3. The neural stimulation system according to any of the preceding claims, wherein the reference signal comprises a subcutaneous electrocardiogram (ECG) signal, the predetermined type timing reference events comprise predetermined type cardiac events, the implantable reference signal sensor comprises one or more subcutaneous electrodes adapted to sense a subcutaneous ECG signal, and the reference event detection circuit comprises a cardiac event detection circuit adapted to detect predetermined type cardiac events from the subcutaneous ECG signal.

4. The neural stimulation system according to claim 3, wherein the cardiac event detection circuit comprises an R-wave detector to detect ventricular depolarizations (R-waves).
5. The neural stimulation system according to claim 3, wherein the cardiac event detection circuit comprises a P-wave detector to detect atrial depolarizations (P-waves).

6. The neural stimulation system according to claim 5, wherein the P-wave detector comprises a signal averaging circuit to average the subcutaneous ECG signal over a plurality of cardiac cycles.

7. The neural stimulation system according to any of claims 3 to 6, wherein the one or more subcutaneous electrodes are incorporated onto the implantable housing.

8. The neural stimulation system according to any of claims 3 to 7, further comprising an arrhythmia detection circuit to detect an arrhythmia from the subcutaneous ECG signal, and wherein the stimulation control circuit is adapted to withhold or adjust the delivery of the neural stimulation pulses when the arrhythmia is detected.

9. The neural stimulation system according to any of claims 3 to 8, further comprising a cardiac parameter measurement circuit to measure one or more cardiac parameters from the subcutaneous ECG signal, and wherein the stimulation control circuit is adapted to adjust the delivery of the neural stimulation pulses based on the measured one or more cardiac parameters.

10. The neural stimulation system according to any of claims 1 and 2, wherein the reference signal comprises an acoustic signal indicative of heart sounds, the predetermined type timing reference events comprise predetermined type heart sounds, the implantable reference signal sensor comprises an acoustic sensor to sense the acoustic signal, and the reference event detection circuit comprises a heart sound detection circuit adapted to detect predetermined type heart sounds using the acoustic signal.
11. The neural stimulation system according to claim 10, wherein the acoustic sensor comprises an implantable accelerometer.

12. The neural stimulation system according to any of claims 10 and 11, wherein the heart sound detection circuit comprises one or more of a first heart sound (S1) detector to detect S1, a second heart sound (S2) detector to detect S2, a third heart sound (S3) detector to detect S3, and a fourth heart sound (S4) detector to detect S4.

13. The neural stimulation system according to any of claims 1 and 2, wherein the reference signal comprises a hemodynamic signal indicative of hemodynamic performance, the predetermined type timing reference events comprise predetermined type hemodynamic events, the implantable reference signal sensor comprises a hemodynamic sensor to sense the hemodynamic signal, and the reference event detection circuit comprises a hemodynamic event detection circuit adapted to detect predetermined type hemodynamic events using the hemodynamic signal.

14. The neural stimulation system according to claim 13, further comprising a derivative calculator coupled to the hemodynamic event detection circuit, the derivative calculator adapted to produce a derivative hemodynamic signal by calculating a time derivative of the hemodynamic signal, and wherein the hemodynamic event detection circuit is adapted to detect the predetermined type hemodynamic events from the derivative hemodynamic signal.

15. The neural stimulation system according to claim 13, wherein the reference event detection circuit comprises a peak detector to detect predetermined type peaks in the hemodynamic signal.

16. The neural stimulation system according to any of claims 13 to 15, wherein the hemodynamic sensor comprises a Doppler echocardiographic sensor to sense a hemodynamic signal indicative of blood flow.
17. The neural stimulation system according to any of claims 13 to 15, wherein the hemodynamic sensor comprises a pressure sensor to sense a pressure signal indicative of blood pressure.

18. The neural stimulation system according to any of claims 13 to 15, wherein the hemodynamic sensor comprises an impedance sensor to sense an impedance signal indicative of blood flow.

19. The neural stimulation system according to any of the preceding claims, wherein the reference event detection circuit comprises a signal processor adapted to extract the predetermined type timing reference events from the reference signal for each of the cardiac cycles.

20. The neural stimulation system according to claim 19, wherein the synchronization module comprises a continuous synchronization module adapted to synchronize the delivery of the neural stimulation pulses to the predetermined type timing reference event consecutively for each of the cardiac cycles.

21. The neural stimulation system according to any of claims 1 to 18, wherein the reference event detection circuit comprises a signal processor adapted to extract each of the predetermined type timing reference events from a segment of the reference signal associated with a plurality of cardiac cycles.

22. The neural stimulation system according to any of the preceding claims, wherein the synchronization module comprises a periodic synchronization module adapted to synchronize the delivery of the neural stimulation pulses periodically for a cardiac cycle of a predetermined number of the cardiac cycles.

23. The neural stimulation system according to any of the preceding claims, wherein the stimulation control circuit comprises an offset interval generator adapted to produce an offset interval starting with one of the predetermined type timing reference events and a pulse delivery controller adapted to start a delivery of a burst of a plurality of neural stimulation pulses when the offset interval expires.
24. A method for operating a neural stimulation system coupled to a living subject having a circulatory system, the method comprising:
   sensing a reference signal indicative of cardiac cycles each including a predetermined type timing reference event using an implantable reference signal sensor placed external to the circulatory system;
   detecting the predetermined type timing reference events from the reference signal; and
   synchronizing delivery of neural stimulation pulses to the detected predetermined type timing reference events.

25. The method according to claim 24, wherein sensing the reference signal comprises sensing a subcutaneous ECG signal using implantable electrodes.

26. The method according to claim 25, wherein detecting the predetermined type timing reference events from the reference signal comprises detecting an atrial depolarizations (P-waves) from the subcutaneous ECG signal.

27. The method according to claim 24, wherein sensing the reference signal comprises sensing an acoustic signal indicative of heart sounds, and detecting the predetermined type timing reference events from the reference signal comprises detecting predetermined type heart sounds from the acoustic signal.

28. The method according to claim 24, wherein sensing the reference signal comprises sensing a hemodynamic signal indicative of blood flow or pressure.

29. The method according to claim 28, wherein detecting the predetermined type timing reference events from the reference signal comprises detecting predetermined type peaks from the hemodynamic signal.

30. The method according to any of claims 24 to 29, further comprising starting an offset interval with one of the detected timing reference event, and wherein synchronizing the delivery of neural stimulation pulses to the detected timing reference events comprises starting a delivery of a burst of a plurality of neural stimulation pulses when the offset interval expires.
31. The method according to any of claims 24 to 30, wherein synchronizing the delivery of neural stimulation pulses comprises synchronizing the delivery of the neural stimulation pulses to the timing reference events of consecutive heart beats on a continuous basis.

32. The method according to any of claims 24 to 30, wherein synchronizing the delivery of neural stimulation pulses comprises synchronizing the delivery of the neural stimulation pulses to the timing reference events of selected heart beats on a periodic basis.

33. The method according to any of claims 24 to 32, wherein detecting the timing reference events from the reference signal comprises extracting each of the timing reference events from a segment of the reference signal recorded during a plurality of cardiac cycles.
CARDIAC CYCLE-SYNCHRONIZED NEURAL STIMULATION CIRCUIT

STIMULATION OUTPUT CIRCUIT

REFERENCE EVENT DETECTION CIRCUIT

REFERENCE SIGNAL SENSOR

STIMULATION CONTROL CIRCUIT

PULSE DELIVERY CONTROLLER

OFFSET INTERVAL GENERATOR

SYNCHRONIZATION CIRCUIT

CONTINUOUS SYNCHRONIZATION MODULE

PERIODIC SYNCHRONIZATION MODULE

TO 106/108

Fig. 3
Fig. 4
Fig. 7

CARCIAL CYCLE-SYCHRONIZED NEURAL STIMULATION CIRCUIT

HEMODYNAMIC SENSOR

TO 106/108

STIMULATION OUTPUT CIRCUIT

HEMODYNAMIC EVENT DETECTION CIRCUIT

STIMULATION CONTROL CIRCUIT

SYNCHRONIZATION MODULE

Fig. 8

SENSING A REFERENCE SIGNAL INDICATIVE OF CARDIAC CYCLES EACH INCLUDING A PREDETERMINED TYPE TIMING REFERENCE EVENT

DETECTING THE PREDETERMINED TYPE TIMING REFERENCE EVENT

SYNCHRONIZING A DELIVERY OF NEURAL STIMULATION PULSES TO THE PREDETERMINED TYPE TIMING REFERENCE EVENT
A. CLASSIFICATION OF SUBJECT MATTER

INV. A61N1/36

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

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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Further documents are listed in the continuation of Box C. See patent family annex.

Date of the actual completion of the international search

21 August 2006

Date of mailing of the international search report

25/08/2006

Name and mailing address of the ISA/
European Patent Office, P.B. 5818 Patentlaan 2 NL – 2280 HV Rijswijk
Tel. (+31-70) 340-2040, Tx. 31 651 epo nl, Fax (+31-70) 340-3018

Authorized officer

Petter, E
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**Box II  Observations where certain claims were found unsearchable (Continuation of item 2 of first sheet)**

This International Search Report has not been established in respect of certain claims under Article 17(2)(a) for the following reasons:

1. **X** Claims Nos.: 24–33 because they relate to subject matter not required to be searched by this Authority, namely:
   
   Rule 39.1(iv) PCT - Method for treatment of the human or animal body by therapy

2. ☐ Claims Nos.: because they relate to parts of the International Application that do not comply with the prescribed requirements to such an extent that no meaningful International Search can be carried out, specifically:

3. ☐ Claims Nos.: because they are dependent claims and are not drafted in accordance with the second and third sentences of Rule 6.4(a).

**Box III  Observations where unity of invention is lacking (Continuation of item 3 of first sheet)**

This International Searching Authority found multiple inventions in this international application, as follows:

1. ☐ As all required additional search fees were timely paid by the applicant, this International Search Report covers all searchable claims.

2. ☐ As all searchable claims could be searched without effort justifying an additional fee, this Authority did not invite payment of any additional fee.

3. ☐ As only some of the required additional search fees were timely paid by the applicant, this International Search Report covers only those claims for which fees were paid, specifically claims Nos.:

4. ☐ No required additional search fees were timely paid by the applicant. Consequently, this International Search Report is restricted to the invention first mentioned in the claims; it is covered by claims Nos.:

**Remark on Protest**

☐ The additional search fees were accompanied by the applicant’s protest.

☐ No protest accompanied the payment of additional search fees.

Form PCT/ISA/210 (continuation of first sheet (2)) (January 2004)
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