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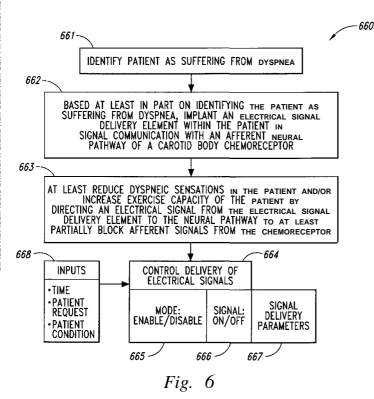
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[Continued on next page]

(54) Title: SYSTEMS AND METHODS FOR TREATING DYSPNEA, INCLUDING VIA ELECTRICAL AFFERENT SIGNAL **BLOCKING** 



(57) Abstract: Systems and methods for treating a patient with dyspnea are disclosed. A method in accordance with a particular embodiment includes identifying the patient as suffering from dyspnea, and, based at least in part on identifying the patient as suffering from dyspnea, implanting an electrical signal delivery element within the patient in signal communication with an afferent neural pathway of a carotid body chemoreceptor. The method can further include at least reducing dyspneic sensations in the patient by directing an electrical signal from the electrical signal delivery element to the neural pathway to at least partially block afferent signals from the chemoreceptor.

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# SYSTEMS AND METHODS FOR TREATING DYSPNEA, INCLUDING VIA ELECTRICAL AFFERENT SIGNAL BLOCKING

#### CROSS-REFERENCE TO RELATED APPLICATION

**[0001]** The present application claims priority to U.S. Provisional Application 61/087,945, filed on August 11, 2008 and incorporated herein by reference.

#### TECHNICAL FIELD

**[0002]** The present disclosure is directed generally to systems and methods for treating dyspnea, including via electrical signals that block or inhibit afferent neural signals from a patient's carotid bodies.

#### **BACKGROUND**

[0003] Dyspnea is the chief patient complaint in a variety of diseases of the pulmonary system. These diseases include chronic bronchitis (12.5 million US patients), emphysema (1.7 million US patients), and asthma (18 million US patients), collectively referred to as Chronic Obstructive Pulmonary Diseases, or COPD. Dyspnea is also reported by patients suffering from combinations of the foregoing diseases, and/or other pulmonary diseases, and non-pulmonary diseases (notably in heart failure). Dyspnea, while a common medical term, is actually poorly defined and ultimately subjective since it is generally the perception of difficulty breathing or difficulty catching one's breath, and more generally, an uncomfortable sensation of breathing.

**[0004]** The severity of pulmonary diseases can typically be measured using objective techniques, such as FEV1 (the patient's forced expiratory volume in the first second of exhalation), minute ventilation (the volume inhaled or exhaled by the patient in one minute), arterial blood gas levels (e.g., of oxygen or carbon dioxide), among others. By contrast, the patient's dyspnea experience can be simply one of difficulty breathing,

ultimately leading to a reduction or elimination of physical activity due to this discomfort. That is, the patient complaint is of dyspnea and a loss of mobility or physical function, not of a decreased FEV1.

[0005] In many ways dyspnea can be analogous to the perception of pain. While an organic source of the pain may be present (a broken bone, for example), the pain itself can be a problem and may require palliative treatment. Furthermore, in the same way that an individual can suffer from chronic pain for which an organic cause is either absent or inadequate to cause the pain, some patients can suffer from severe dyspnea despite relatively normal objective measures of pulmonary performance.

[0006] The origins of dyspnea remain unclear. Studies and experience have yielded confusing and often seemingly contradictory results. Treatments for dyspnea range from supplemental oxygen therapy to sitting in front of a fan to systemic opiates. Furthermore, dyspnea can be experimentally induced by vigorous exercise, breath-holding, breathing through a restrictive mouthpiece, or breathing carbon dioxide in symptomatic pulmonary disease patients. A common, though unproven theory, is that dyspnea derives from a mismatch between outgoing motor signals to the respiratory muscles and incoming afferent information. In one example, under a give set of conditions, the brain can expect a certain pattern of ventilation and associated afferent feedback. Deviations from this pattern can cause or intensify the sensation of dyspnea.

[0007] While dyspnea is often the chief complaint of a patient, there is currently no pharmacologic agent that primarily treats dyspnea. That is, a variety of bronchodilators are used to treat asthma and other COPD, and while they demonstrably increase FEV1, their effects on dyspnea can be modest and can fall below that of clinical significance. Accordingly, there remains a need for methods and devices that effectively treat dyspnea.

#### BRIEF DESCRIPTION OF THE DRAWINGS

[0008] Figure 1 is a partially cutaway, partially schematic illustration of the vascular and neural structure of a patient's neck, based on plate 124 of "Atlas of Human Anatomy", 2nd Edition, by Frank Netter (Icon Learning Systems, 2001).

**[0009]** Figure 2 is a partially schematic illustration of a system for delivering inhibitory signals to a patient's afferent chemoreceptor neural pathways in accordance with an embodiment of the disclosure.

**[0010]** Figure 3 is a partially schematic, generally ventral view of a patient's carotid arteries and carotid branch, based on Figure 1 of an article titled "Bilateral Carotid Body Resection for Asthma and Emphysema" (Winter, *International Surgery*, Volume 57, No. 6, June 1972, hereinafter "Winter").

**[0011]** Figure 4 is a partially schematic lateral view of the patient's carotid artery region, based on Figure 4 of Winter.

**[0012]** Figure 5 is a partially schematic illustration of a lead suitable for providing signals in accordance with an embodiment of the disclosure.

**[0013]** Figure 6 is a block diagram illustrating a method for treating a patient in accordance with an embodiment of the disclosure.

**[0014]** Figure 7 is a block diagram illustrating a method for diagnosing and treating a patient in accordance with another embodiment of the disclosure.

#### DETAILED DESCRIPTION

[0015] The present disclosure is directed generally to systems and methods for treating dyspnea, including via electrical afferent signal inhibition. Specific details of several embodiments of the disclosure are described below with reference to particular implementations to provide a thorough understanding of these embodiments, but in other embodiments, the systems and methods may have different features. Several details describing structures or processes that are well-known and often associated with related systems and techniques, but that may unnecessarily obscure some significant aspects of the present disclosure, are not set forth in the following description for purposes of clarity. Moreover, although the following disclosure sets forth several embodiments of different aspects of the invention, several other embodiments can have different configurations or different components than those described in this section. As such, the disclosure may

include other embodiments with additional elements, or without several of the elements described below with reference to Figures 1-7.

[0016] The present disclosure incorporates several documents by reference. In the event of inconsistent usages between this document and those documents so incorporated by reference, the usage in the incorporated reference(s) should be considered supplementary to that of this document. For irreconcilable inconsistencies, the usage in this document controls.

#### Overview

[0017] Several embodiments of the present disclosure are directed to using electrical signals to block, partially block, or otherwise inhibit afferent chemoreceptor signals in a patient suffering from dyspnea. The chemoreceptors provide signals to the patient's brain indicating a low level of oxygen or a high level of carbon dioxide in the patients' blood. Electrical signals provided in accordance with the technology described herein can interrupt the neuronal signals otherwise transmitted to the patient's brain by the patient's chemoreceptors, thus alleviating or eliminating the patient's sensation of breathlessness. As will be discussed in further detail later, this approach can have significant advantages over existing surgical techniques, which are irreversible and not adjustable. These and other advantages are described further below.

#### Relevant Physiology

[0018] Figure 1 is a partially schematic, partially cutaway illustration of the neck region 101 of a patient 100. For purposes of clarity, many anatomical features in the neck region 101 have been eliminated in Figure 1. Figure 1 illustrates several relevant structures located on the right side of the patient's midline; however, the following discussion applies equally to contralateral structures located on the left side of the patient's midline. As will also be discussed further below, certain procedures may be conducted unilaterally (on either the left or right structure) or bilaterally (on both the left and right structures).

[0019] As shown in Figure 1, the common carotid artery 102 supplies blood to the patient's head, and splits to form the internal carotid artery 103 and the external carotid artery 104 in the neck region 101. The carotid sinus 105 is located in the region of the split between the internal carotid artery 103 and the external carotid artery 104. The carotid body 106 is a small sensory organ located high in the neck region 101, posterior to the patient's lower jaw and generally in the bifurcation region between the internal carotid artery 103 and the external carotid artery 104. The carotid body 106 is perfused by blood flow in the carotid arteries 102-104. The carotid body 106 generally includes peripheral chemoreceptors that can sense the oxygen level, carbon dioxide level, and possibly pH or other factors in the arterial blood flow. These chemoreceptors provide the large majority of the patient's ability to sense oxygen, and also provide approximately one-third of the patient's ability to sense carbon dioxide. Other chemoreceptors in the medulla provide approximately two-thirds of the patient's ability to sense carbon dioxide.

[0020] Afferent signals travel from the carotid body 106 to the patient's brain via the carotid branch 108 of the patient's glossopharyngeal nerve 109. The glossopharyngeal nerve 109 is the ninth cranial nerve, and descends alongside the vagus nerve or tenth cranial nerve 110. For purposes of simplicity, other cranial nerves are not specifically identified in Figure 1. The carotid body 106, the carotid branch 108, and the glossopharyngeal nerve 109 form an afferent neural pathway 107 along which afferent signals from the chemoreceptors of the carotid body 106 are transmitted to the patient's brain.

[0021] While the exact roles and interrelationships of the various chemoreceptors remain unknown, it is commonly accepted that they can react to low levels of oxygen or high levels of carbon dioxide, and can send afferent signals to the central respiratory center that can increase the patient's ventilatory drive in order to normalize the concentration of blood gases. Conversely, removing or debilitating the carotid body 106 can render it less able to support the body's response to changing levels of blood gases by eliminating an input to the central respiratory center. In certain examples, a complete

lack of response to low oxygen levels (hypoxia) or high levels of carbon dioxide (hypercapnia) can be life threatening.

#### Carotid Body Resection

[0022] A rare genetic mutation (often in Dutch patients) results in glomus cell tumors of the carotid body, requiring their removal via Carotid Body Resection or CBR. In some patients, bilateral tumors have required the removal of both carotid bodies. These patients typically exhibit increased exercise capacity, decreased dyspnea, increased breath-holding capacity, and a blunted ventilatory response to exercise. However, there is also evidence that these effects may be transitory and that the body may accommodate to the loss of the carotid bodies after some period of time.

[0023] Despite its current use for tumor resection, intentional CBR to treat pulmonary conditions is not currently in favor. Generally, CBR was used from the 1940's to the 1980's to treat dyspnea, including in asthmatic patients. The current disfavor derives from uncertainties about proper patient selection criteria, from concerns about inducing central sleep apnea, from possibly detrimental blood gas changes, and from the radical and irreversible nature of the surgery. Much of the disfavor derives from the history of treating asthmatic children, whose lack of response to hypoxia led them to delay treatment of acute status asthmaticus, resulting in several deaths.

[0024] Despite the foregoing drawbacks, CBR has also been associated with a significant decrease in dyspnea and a corresponding increase in exercise capacity. While blood gases are impacted, most authors appear to judge these changes to be relatively minor in many or most cases. For example, typical sequela to the chronic loss of carotid body function (in patients presumably not suffering from pulmonary disease) is a modest 6 mm Hg rise in PaCO<sub>2</sub>, where "PaCO<sub>2</sub>" refers to the partial pressure of carbon dioxide in the patient's arterial blood. In certain examples, patients selected for CBR have PaCO<sub>2</sub><45 mm Hg and PaO<sub>2</sub>>65 mm Hg (where "PaO<sub>2</sub>" refers to the partial pressure of oxygen in the patients' arterial blood), in order to ensure only a modest impact. In certain examples, many patients can have either improvements or minimal degradations in blood gases.

[0025] The literature and experience are mixed regarding the permanence of the effect of CBR. In certain examples, the effect of CBR decreases over time as the body compensates for the loss of signaling from the carotid bodies. In other examples, after CBR, other chemoreceptors can increase their responsiveness to carbon dioxide, raising the total body response back to normal levels, while having little to no compensatory change in the body's response to oxygen levels.

#### Inhibitory Electrical Stimulation

[0026] Aspects of the present disclosure are directed to replacing, in whole or in part, the foregoing resection procedure with an electrical stimulator implant procedure. The implanted stimulator can apply an inhibitory electrical signal to the afferent neural pathway 107, thus emulating at least in part the results of carotid body resection, but with additional control over the chemoreceptor inhibition and/or without several of the foregoing drawbacks.

[0027] Figure 2 is a partially schematic, isometric illustration of a system 120 suitable for delivering inhibitory electrical signals to the afferent neural pathway 107 described above with reference to Figure 1. Accordingly, by blocking, at least partially blocking, and/or otherwise inhibiting afferent neural signals from the chemoreceptors at the carotid body 106 (Figure 1), the patient's dyspneic symptoms can be reduced, alleviated and/or eliminated. As described further below, this can be done without unduly interfering with the body's ability to detect and respond to low blood oxygen levels and/or high blood carbon dioxide levels.

[0028] As shown in Figure 2, the system 120 can include a signal delivery device 121 coupled to a controller 122 with a communication link 126. In one aspect of this embodiment, the signal delivery device 121 can include a cuff electrode 123 that in turn includes a cuff or other support body 124 carrying one or more arcuate electrical contacts 125 (three are shown in Figure 2 as first, second, and third electrical contacts 125a, 125b, 125c). In this embodiment, the cuff electrode 123 can have a tripolar arrangement, with the first and third contacts 125a, 125c connected to an anodic potential, and the second contact 125b connected to a cathodic potential. Accordingly, the outer anodic contacts

125a, 125c can direct the field lines emanating from the central cathodic contact 125b to a target neural population. In other embodiments, the cuff electrode 123 can include other arrangements of contacts, e.g., to provide monopolar or bipolar signals. In any of these embodiments, the cuff electrode 123 can be positioned around or adjacent to any suitable portion of the afferent neural pathway 107 (Figure 1), including the carotid body 106 itself and/or the carotid branch 108. The particular location at which the cuff electrode 123 is placed can be selected based on factors including an individual patient's physiology, characteristic chemoreceptor responsiveness and/or characteristic baroreceptor responsiveness.

[0029] The signal delivery device 121 is coupled to the controller 122 via a communication link 126 having a first connector 127a that may be releasably engaged with a second connector 127b. The second connector 127b is electrically coupled to the controller 122, and both the second connector 127b and the controller 122 can be carried by an enclosed, hermetically sealed housing 128. The housing 128 can also enclose an internal power source 129 which provides power to a pulse generator 130. The pulse generator 130 generates pulses (e.g., square wave, biphasic, charge-balanced and/or other suitable pulses) that are transmitted to the signal delivery device 121 via the communication link 126, under the direction of the controller 122. The controller 122 can accordingly include a memory 131, a processor 132, and a receiver/transmitter 133. Instructions for delivering the electrical signal to the patient via the signal delivery device 121 can be stored in or on one or more computer readable media of the controller 122, e.g., the memory 131 and/or the processor 132. Accordingly, the controller 122 can include a specially programmed computer device.

[0030] In a particular embodiment, the receiver/transmitter 133 can receive inputs from devices within or outside the housing 128 to operate in an open loop manner and/or a closed loop manner. For example, the system 120 can include an external controller 134 that communicates instructions to the receiver/transmitter 133 via a wireless link. A physician can use an external controller 134 to change the instructions carried out by the controller 122. In another embodiment, the patient can use an external controller 134 to start and/or stop the signals directed by the controller 122 to the signal delivery device

121. The physician and the patient can each have separate external controllers 134, with the physician's external controller 134 able to carry out a broader range of control tasks than the patient's external controller 134. This arrangement can prevent the patient from inadvertently changing signal delivery parameters in an undesirable manner, while still allowing the patient to control certain tasks e.g., starting and stopping the electrical signals.

[0031] The receiver/transmitter 133 can also receive information from one or more sensors 136. The sensors 136 can be configured and positioned to provide information to the controller 122 useful for determining when and in what manner to provide electrical signals to the signal delivery device 121. For example, the sensor 136 can include an oxygen sensor (e.g., an implanted oxygen sensor, or an external fingertip-mounted oxygen sensor) that identifies the patient's blood oxygen levels. In another embodiment, the sensor 136 can include an accelerometer or other device that detects the patient's activity level. In still another embodiment, the sensor 136 can include a clock or timer that senses the passage of time, which, as described further below, may also be used to control the manner in which electrical signals are delivered to the patient.

As shown in Figure 2, the internal power source 129 can be included in the implantable housing 128. The housing 128 can be implanted at a subclavicular location in the patient's chest, or at another suitable location. In a particular embodiment (e.g., when the internal power source 129 includes a rechargeable battery), the system 120 can include an external power source 135 that is used to recharge the internal power source 129 within the implanted housing 128. For example, the external power source 135 can recharge the internal power source 129 via inductive coupling. In another embodiment, the internal power source 129 can be eliminated, and the external power source 135 alone can provide power to implanted controller 122. In still another embodiment, the controller 122 and the other components located in the housing 128 can be positioned outside the patient's body. For example, these components can be placed in an external housing and worn by the patient, e.g., beneath the patient's clothing. The controller 122 can be connected to the signal delivery device 121 via a hardwired transdermal communication link, or via a wireless transcutaneous link. Any one or combination of the foregoing

arrangements can be used to transmit suitable electrical signals to the signal delivery device 121.

Figure 3 is a partially schematic, ventral illustration of a portion of the anatomy [0033] shown in Figure 1, illustrating a technique for implanting a signal delivery device in accordance with an embodiment of the disclosure. Certain aspects of this technique are generally similar to those disclosed by Winter in an article titled "Bilateral Carotid Body Resection for Asthma and Emphysema," (International Surgery, Volume 7, No. 6, June 1972, hereinafter "Winter"), incorporated herein by reference. Figure 3 illustrates the region at which the common carotid artery 102 bifurcates into the internal carotid artery 103 and the external carotid artery 104. Arrows M and L identify medial and lateral directions, respectively. Most of the external carotid artery 104 has been cut away so as to illustrate the carotid branch 108 located between the internal carotid artery 103 and the external carotid artery 104. The carotid branch 108 includes baroreceptor neurons 112 and chemoreceptor neurons 111 that extend to the carotid sinus 104 and the carotid body 106 to transmit afferent neural signals from the patient's baroreceptors chemoreceptors, respectively. In a particular embodiment of the method disclosed herein, a probe 137 or other surgical implement can be used to separate or divide the chemoreceptor neurons 111 from the baroreceptor neurons 112, thus allowing a representative signal delivery device 121 (Figure 2) to be positioned in a manner that preferentially directs signals to the chemoreceptor neurons 111 over the baroreceptor neurons 112. For example, the cuff electrode 123 (Figure 2) can be positioned around the chemoreceptor neurons 111, with the electrode contacts 125a-c preferentially positioned to direct signals to the chemoreceptor neurons 111 over the baroreceptor neurons 112. While some electrical signals may still reach the baroreceptor neurons 112, this arrangement can more significantly inhibit or block afferent signals transmitted along the chemoreceptor neurons 111, without unduly interfering with afferent signals transmitted by the baroreceptor neurons 112.

[0034] Figure 4 is a partially schematic, left lateral view of the anatomy shown in Figure 3, also based on Winter, and illustrating the chemoreceptor neurons 111 after having been divided from the baroreceptor neurons 112. Arrows D and V identify dorsal

and ventral directions, respectively. As discussed above, a suitable signal delivery device 121 (Figure 2) can be positioned so as to preferentially direct electrical signals to the chemoreceptor neurons 111. In at least some embodiments, the characteristics of the signal delivery device 121 itself and the orientation of the signal delivery device 121 will preferentially direct signals to the chemoreceptor neurons 111. In other embodiments, the physician can implant an insulating shield 138 between the chemoreceptor neurons 111 and the baroreceptor neurons 112 so as to restrict or prevent electrical signals directed to the chemoreceptor neurons 111 from unduly affecting the baroreceptor neurons 112, and/or other nearby or neighboring structures. The shield 138 can be held in place with a suitable adhesive, suitable friction features (e.g., nubs) and/or other arrangements.

[0035] The signal delivery device 121 positioned in signal communication with the chemoreceptor neurons 111 can have an arrangement generally similar to that shown in Figure 2, or in other embodiments, it can have other arrangements. For example, Figure 5 illustrates a signal delivery device 521 that includes a lead 539 typically used for spinal cord stimulation. The lead 539 can carry a plurality of electrical contacts 525 (eight are shown in Figure 5 as contacts 525a, 525b... 525h). Each of the contacts 525 can have an annular ring shape with an outer surface exposed at the outer surface of the lead 539. Each contact 525 can be connected to an individual electrical conductor 540 (e.g., a wire) so as to receive a separately programmable electrical signal. The individual conductors 540 can form a communication link 526 via which the contacts 525 are connected to the controller 122 (Figure 2). The individual contacts 525a-525h can be selectively activated to provide suitable inhibitory signals, without unnecessarily stimulating adjacent structures. Individual active contacts may selectively be activated or deactivated to avoid habituation and/or tissue necrosis. The lead 539 can be delivered percutaneously and positioned alongside the chemoreceptor neurons 111 shown in Figure 4, and can be secured in place using suitable sutures or other securement techniques.

**[0036]** In one embodiment, the practitioner (e.g., a surgeon or other physician) can apply the dividing technique and the signal delivery device implanting technique described above to both left and right carotid branches. Accordingly, the practitioner can implant two bilaterally positioned signal delivery devices, coupled to a common implanted or

externally-worn controller. In another embodiment, the chemoreceptor neurons and/or the carotid body on one side of the patient's midline can be resected, ligated or otherwise surgically disabled, and the contralateral structures can receive inhibitory signals from an implanted electrical signal delivery element. In a further aspect of this embodiment, the resection process can include resecting both the chemoreceptor neurons and the baroreceptor neurons by resecting the carotid branch 108 or the carotid body 106. Due to the body's other still-active baroreceptors, the effect of this procedure on the body's overall baroreceptor functioning is not expected to be significant for at least some classes of patients. Due to the combination of surgically disabling chemoreceptor pathways on one side of the patient's body, and applying inhibitory electrical stimulation to the chemoreceptor pathways on the opposite side of the patient's body, the patient's dyspneic effects can be controlled.

[0037] The process of resecting one chemoreceptor pathway (and optionally one baroreceptor pathway) may be advantageous because it can simplify the surgical procedure, and/or it can conserve battery power. For example, it may be simpler for the practitioner to implant a signal delivery device at the patient's right side chemoreceptor pathway and then resect the patient's left side chemoreceptor pathway than it is for the practitioner to implant a second signal delivery device and tunnel the associated communication link to an implanted controller. In addition, one signal delivery device is expected to consume less power than two. Conversely, some patients may benefit from the ability to reactivate and/or modulate chemoreceptor functioning on both sides of the midline. In such cases, the practitioner may implant bilateral signal delivery devices, as discussed above, rather than surgically disable one chemoreceptor afferent neural pathway.

[0038] Figure 6 is a block diagram illustrating a process or method 660 for treating a patient in accordance with a particular embodiment of the present disclosure. The process 660 includes identifying a patient as suffering from dyspnea (process portion 661). The process 660 can further include implanting an electrical signal delivery element within the patient to be in signal communication with an afferent neural pathway of a carotid body chemoreceptor, based at least in part on identifying the patient as suffering

from dyspnea (process portion 662). Accordingly, the process 600 can include deliberately tying a diagnosis of dyspnea to treatment of dyspneic symptoms via specifically directed inhibitory electrical signals provided by an electrical signal delivery element in signal communication with target neurons of the afferent neural pathway. As used in this context, signal communication means that electrical signals emanating from the electrical signal delivery element have a direct effect on the afferent neural pathway by virtue of interactions between the electrical signal and the neurons of the afferent neural pathway. For example, the electrical signal delivery element can include one or more electrical contacts positioned along the carotid branch 108 of the glossopharyngeal nerve 109 (Figure 1). In particular embodiments, the electrical contacts can be positioned at or up to about five centimeters superior to the carotid body 106. The electrical contacts can be in physical contact with the adjacent neural tissue, or otherwise close enough to the neural tissue to have the desired inhibitory effect on the afferent signals. The contacts can accordingly be positioned to preferentially direct signals to the chemoreceptor neurons, as described above with reference to Figures 4 and 5.

[0039] The process 660 can further include at least reducing dyspneic sensations in the patient and/or increasing the patient's capacity for exercise by directing an electrical signal from the electrical signal delivery element to the neural pathway to at least partially block afferent signals from the chemoreceptor (process portion 663). For example, the electrical signal delivery element can direct biphasic, square wave, charge-balanced pulses at a frequency from about 1,000 Hz to about 10,000 Hz, an amplitude of up to about 12 volts or up to about 10 milliamps, with individual pulses having a duration of from about 10 microseconds to about 1,000 microseconds. In a particular embodiment, the frequency is selected to be about 5,000 Hz, the pulse width is selected to be 30-50 microseconds, and the amplitude is selected to be from about 1 mA to about 5 mA, with the selected value chosen to avoid muscle capture. In a particular embodiment, the pulses can be applied unilaterally to the afferent neural pathway associated with either the left side or right side carotid body. In a particular aspect of this embodiment, the signals can be provided in a burst lasting about 120 seconds. The resulting block or partial block provided by the applied signals can have a persistence (e.g., an effective duration after

the end of the burst) that lasts for approximately 90 to 120 seconds. The signal can be applied with a duty cycle of about 50% in a particular embodiment, or other suitable values selected to provide efficacy while conserving power in other embodiments. The level of efficacy can be based at least in part on the persistence effect of the signal.

[0040] In another embodiment, the stimulation can be provided bilaterally, via one signal delivery device positioned along one afferent neural pathway, and another signal delivery device positioned along the contralateral afferent neural pathway. Each of the signal delivery devices can be connected to a common controller. In this arrangement, delivering bilateral electrical signals can produce the desired inhibitory effect on afferent signals transmitted from the chemoreceptors that lasts for hours rather than minutes.

[0041] The process 660 can further include controlling the delivery of the electrical signals provided to the signal delivery element (process portion 664). The signals can be controlled in a manner that is responsive to one or more inputs 668. In a particular embodiment, the electrical signals can be controlled in part by alternating between an enable mode and a disable mode (process portion 665). In the enable mode, signals may be selectively turned on and off (process portion 666) via a separate instruction. The signals are the delivered in accordance with suitable signal delivery parameters 667, including the frequencies, amplitudes and pulse widths described above. In the disable mode, signals may not be selectively turned on or off, despite the separate instruction. For example, the system can enter the enable mode during normal waking hours, allowing the patient to activate the signal delivery element on an as-needed basis via a separate, patient-directed input. During normal sleeping hours, the system can enter the disable mode, during which the system will not direct electrical signals, even if concurrently requested by the patient. This arrangement can prevent the patient from inadvertently inhibiting the carotid body chemoreceptors at night, so as to reduce or eliminate the possibility of inducing sleep apnea. During waking hours, with the system in the enable mode, the patient can selectively inhibit chemoreceptor afferent signals, for example, before and/or during exercise, by providing a separate instruction. The instruction can be provided when the patient presses a button or otherwise enters an input signal via the external patient controller 134 described above with reference to Figure 2.

The foregoing example is representative of one in which the system responds [0042] to a time input (e.g. normal waking hours and normal sleeping hours) and a patient request (e.g. a specific request for inhibitory electrical signals). In other embodiments, the system can respond to other time-based and/or patient-based inputs. For example, the system can remain on (e.g. actively delivering signals to the signal delivery element) for a period of 30 minutes or another suitable period, in response to receiving a patient request. The patient request can come in the form of an input from a simple electronic device (e.g., the patient controller 134 described above with reference to Figure 2), or a magnet that activates a reed switch. The patient request can arrive in anticipation of carrying out an activity (e.g. exercise), and/or can be provided during an activity. Accordingly, the system can automatically provide for a maximum active signal time per patient-initiated or otherwise-initiated activation. In another embodiment, the system can automatically provide for a minimum inactive time or off time between patient or otherwise initiated activations. In still a further embodiment, the system can automatically track a maximum amount of signal delivery time per suitable time interval. For example, the system can track a maximum number of hours of active signal time per day. If the patient requests more than this amount of time, the system can prevent further activations, or require a particular activation sequence or physician intervention before authorizing additional activations. Representative suitable activation times per day include 1, 2, 4, 6 or 8 hours.

[0043] In still further embodiments, process portion 664 can include receiving inputs relating to patient state or condition. For example, the system can deliver or enable delivery of the electrical signals in response to an indication that the patient is physically active (a first state or condition), and disable or cease delivering the signals when the patient is resting or relaxing (a second state or condition). In a particular example, an accelerometer or other motion detection device can provide suitable inputs for this mode of operation. In another embodiment, an electrocardiogram can provide a generally similar function by providing an indication of the patient's heart rate. In still further embodiments, indications of the patient's condition can be used to provide alerts and/or to prevent the inhibitory signals from being directed to the chemoreceptors. For example, the input can include an indication of the patient's blood oxygen level, and can prevent the

system from delivering electrical signals if the blood oxygen level is below a particular threshold. In representative embodiments, the system can deliver signals when the blood oxygen level is at or above a first threshold of 89%, 90%, 91%, 92% or 93%, and can cease delivering signals when the blood oxygen level is at or below a second threshold of 93%, 92%, 91%, 90% or 89%. In other embodiments, these values can be based on partial pressures (e.g., 60, 65 or 70 mm PaO<sub>2</sub>). In still further embodiments, the foregoing activation determination can be based on blood carbon dioxide levels (e.g., less than 35, 40, 45 or 50 mm PaCO<sub>2</sub>). The foregoing actions can be accompanied by an alarm function, or the alarm function can be provided without the automatic shutdown feature, but with the patient taking the separate step of shutting the system down. This arrangement can make use of an external or internal pulse-oxymeter, such as a light-sensitive oxymeter worn on the patient's fingertip.

[0044] In other embodiments, the detector can detect breathing motion and/or other pulmonary indicators. Any of the foregoing detection techniques can be used to monitor the patient, alert the patient, and/or automatically or manually shut the system off or otherwise disable or deactivate the system, while the patient is awake and inactive, while the patient is awake and active, or while the patient is asleep. In any of these embodiments, the system can be subsequently reactivated or enabled, e.g. by a physician or other practioner, or by the patient.

Figure 7 is a block diagram illustrating a method 760 that includes diagnosing and/or screening patients in accordance with a particular embodiment of the disclosure. In general, the foregoing procedures for treating dyspnea may be applied to patients with severe or very severe COPD, as evidenced by FEVI values of between about 20% and about 40%. Such patients may also demonstrate exercise limited by dyspnea (e.g. a value of 3-4 on the MRC 1-5 point scale). Such patients may have similar blood chemistries to those selected for CBR, e.g.,  $PaCO_2 < 45 \text{ mm}$  Hg and  $PaO_2 > 65 \text{ mm}$  Hg.

[0046] Process portion 761 includes using a non-invasive, physiological-functioning screening procedure to assess the patient's suitability for a dyspnea treatment regimen. This process can include, for example, simulating chemoreceptor inhibition to determine or estimate the likelihood that the patient will respond to the subsequent inhibition provided

by an implanted signal delivery device and/or resection. The process can, in addition to or in lieu of the foregoing screen, include assessing the patient's baroreceptor functioning level, which can be used to aid the physician in determining whether some of the patient's baroreceptor functions can be reduced or eliminated in combination with inhibiting chemoreceptor functions. Each of these assessments is described in turn below.

The practitioner can employ one or more of several techniques for reversibly simulating the effects of chemoreceptor inhibition before undertaking a resection or implant procedure. For example, the patient can receive a locally injected anesthetic applied to the carotid bodies during or before an exercise test to determine the impact of carotid body blocking on exercise capacity and/or perceived dyspnea. The dyspneic relief and/or increase in exercise capacity can be compared to the impact on the patient's blood gasses to determine a suitable treatment. For example, if the patient reports a positive effect on the dyspneic symptoms and/or an increased capacity for exercise, and results indicate an acceptable effect on the patient's blood gasses (e.g. an acceptable reduction in blood oxygen), the patient may be identified as a suitable candidate for further treatment. A representative acceptable reduction in blood oxygen level is about 6 mm Hg PaO<sub>2</sub>, or less.

In another example, the carotid body's output can be artificially suppressed by the inhalation of high fraction oxygen (a gas mixture of greater than approximately 30% oxygen). Aspects of a representative process carried out on COPD patients are described by Somfay et al. in "Dose-reponse effect of oxygen on hyperinflation and exercise endurance in nonhypoxaemic COPD patients" (European Respiratory Journal, vol. 18, 77-88, 2001, incorporated herein by reference). These individuals exhibited significantly increased exercise capacity due to the elimination of carotid body nervous signaling. Accordingly, breathing such a hyperoxic gas mixture can be used both to quantify the magnitude of the effect of inhibitory electrical stimulation or combined electrical stimulation and resection. Suitable levels of oxygen for the foregoing screening procedure include at least 30%, at least 40%, at least 50% and about 100%.

[0049] Another aspect of process portion 761 is assessing the patient's baroreceptor functioning level. For example, known reversible techniques such as carotid sinus

massage or intraoperative anesthetic applied to the carotid sinus can be used to determine the functioning of the baroreceptors carried by the patient's carotid bodies. In particular, these techniques can be used to determine if disabling the carotid sinus baroreceptor function has an effect on the patient's overall barosensing function, and/or to determine the patient's sensitivity to loss of sinus cavity barosense function. determined that the patient's overall baroreceptor functioning is adequate without the specific feedback provided by the carotid body baroreceptors (e.g. if other baroreceptors located in the brain or elsewhere by themselves provide a suitable functioning level at or above a selected threshold), then the practitioner can undertake procedures that may inhibit the baroreceptor functioning of the carotid bodies in addition to the chemosensing functioning of the carotid bodies. For example, when resecting the carotid body or carotid nerve, the practitioner can resect both the chemoreceptor neurons and the baroreceptor neurons if doing so will not unacceptably affect the patient's baroreceptor functioning. Similarly, the practitioner can implant the signal delivery device in a manner that may inhibit afferent signals from the carotid body baroreceptors (e.g., without preference to inhibiting only chemoreceptor neurons) if doing so is not expected to unacceptably affect the overall baroreceptor functioning of the patient. If the patient's barosense function is significantly affected by temporarily reducing or eliminating the carotid sinus barosense function, the practitioner can take appropriate steps to preferentially direct inhibitory signals to the chemoreceptor neurons to avoid unnecessarily obstructing the functioning of the baroreceptor neurons.

[0050] After the chemoreceptor and/or baroreceptor screening functions have been conducted (process portion 761), the patient may be selected for dyspnea treatment (process portion 762). The dyspnea treatment can include, for example, the techniques described above with reference to Figures 1-6. In other embodiments, the screening techniques shown in Figure 7 can be used as a precursor to other dyspnea therapies.

[0051] One aspect of several of the embodiments described above is that the foregoing systems and methods can relieve the patient's dyspnea while the patient is exercising and/or engaging in other waking activities, with an expected modest degradation of blood gasses. An advantage of this arrangement is that by reducing

dyspneic symptoms, the patient can be more relaxed and more likely to engage in exercise, and/or engage in exercise more often, and/or engage in more strenuous exercise, which can facilitate the patient's participation in pulmonary rehabilitation. The patient may have a brief reduction in blood oxygen (e.g. from 95% to 92%), but the short-term effect of this reduction may be more than offset by the long-term effects of increased exercise. This can halt or even reverse what has been viewed as a dyspneic spiral in which (a) the patient suffers from COPD and dyspnea, (b) the dyspnea discourages the patient from engaging in exercise, which (c) exacerbates the COPD and dyspnea. Thus, in a manner analogous to pain treatment, reducing the patient's dyspnea can allow the patient to engage in activity despite the existence of an underlying condition, and in at least some instances, to an extent that treats the underlying condition.

[0052] In addition to or in lieu of increasing the patient's level of exercise, embodiments of the foregoing systems and methods can increase the patient's level of other activities, e.g., activities of daily living. In addition to, or in lieu of the foregoing advantages, aspects of the foregoing systems and methods can improve the patient's quality of life by eliminating or reducing the patient's use of and/or reliance on supplemental oxygen, which is typically carried by the patient in a heavy, awkward tank. Eliminating or reducing the use of supplemental oxygen is expected to further increase the likelihood and/or frequency with which the patient exercises.

[0053] Another feature of several of the foregoing embodiments is that the effect of the electrical signal on the chemoreceptor afferent neural pathway is controllable and reversible. Accordingly, the signal can be halted while the patient is sleeping, resting, relaxing, or otherwise not engaged in strenuous activity. This can allow the afferent neurons to return to a normal state and, correspondingly allow the patient's blood gasses to return to normal (or at least normal for that patient) levels. In addition, if inhibiting the chemoreceptor afferent neural pathway later becomes undesirable, the system can be deactivated. The system can subsequently be reactivated if inhibiting the chemoreceptors again becomes part of a suitable therapy. This is unlike resection, which is generally irreversible.

[0054] Still another feature of at least some of the foregoing embodiments is that the system can automatically monitor and respond to changes, and can control the delivery of electrical signals accordingly. For example, the system can automatically disable electrical signal delivery during normal sleeping hours to avoid apnea. The system can automatically disable electrical signal delivery if blood gasses degrade below desired levels. This level of automation takes advantage of and builds on the fact that the effect of the electrical signals is reversible and haltable.

Still another aspect of several of the foregoing embodiments is that the system can be selectively activated and deactivated as part of an overall treatment regimen that includes exercise. For example, the system can be activated during high intensity exercise experienced during pulmonary rehabilitation or post rehabilitation maintenance. At other times, the system can be deactivated. In still a further embodiment, the degree to which the system inhibits the afferent signals from the chemoreceptors can be controlled. For example, the amplitude and/or duty cycle of the signal applied to the neural pathways can be increased for additional inhibition and decreased for less inhibition. In another approach that can be used in addition to or in lieu of the foregoing approach, the carotid body chemoreceptors on one side of the patient's midline can be inhibited while those on the other side are not. In yet a further embodiment, the system can alternate between inhibiting chemoreceptors on one side of the midline and those on the other. arrangement can be used to forestall or prevent patient habituation to the electrical signals. In addition to or in lieu of this arrangement, the signal delivery parameters (e.g., signal amplitude, frequency, pulse width, and/or parameters) can be varied to forestall or prevent habituation. In still another embodiment, the system can be activated only at selected times (e.g., during exercise) to reduce habituation. Accordingly, the foregoing processes can extend the effectiveness of the therapy for a longer, controlled period of time.

[0056] From the foregoing, it will be appreciated that specific embodiments of the disclosure have been described herein for purposes of illustration, but that various modifications may be made without deviating from the disclosure. For example, the particular signal delivery parameters described above can have other values in other

embodiments. The particular electrodes described above can have other configurations and other embodiments. The system can include sensors other than those specifically identified above. Certain aspects of the disclosure described in the context of particular embodiments may be combined or eliminated in other embodiments. For example the patient may receive an electrode device of the type shown in Figure 2 on one side of the body, and an electrode device of the type shown in Figure 5 on the other side. Further, while advantages associated with certain embodiments have been described herein in the context of those embodiments, other embodiments may also exhibit such advantages, and not all embodiments need necessarily exhibit such advantages to fall within the scope of the present invention. Accordingly, the disclosure can include other embodiments not explicitly shown or described above.

#### **CLAIMS**

#### I/We claim:

1. A method for treating a patient, comprising:

identifying the patient as suffering from dyspnea;

- based at least in part on identifying the patient as suffering from dyspnea, implanting an electrical signal delivery element within the patient in signal communication with an afferent neural pathway of a carotid body chemoreceptor; and
- at least reducing dyspneic sensations in the patient, or increasing an exercise capacity of the patient, or both, by directing an electrical signal from the electrical signal delivery element to the neural pathway to at least partially block afferent signals from the chemoreceptor.
- 2. The method of claim 1, further comprising;

hours;

delivery; and

- automatically enabling delivery of the electrical signal during normal waking hours; automatically disabling delivery of the electrical signal during normal sleeping
- directing the electrical signal only when both (a) the delivery of the electrical signal is enabled, and (b) the patient initiates a concurrent request for signal
- preventing delivery of the electrical signal when (c) the delivery of the electrical signal is enabled, even if (d) the patient initiates a concurrent request for signal delivery.
- 3. The method of claim 1 wherein directing the electrical signal includes directing the electrical signal while the patient is exercising.

4. The method of claim 3, further comprising increasing the patient's level of exercise while the electrical signal is directed to the neural pathway, relative to the patient's exercise level when the electrical signal is not directed to the neural pathway.

- 5. The method of claim 1, further comprising increasing the patient's activity level while the electrical signal is directed to the neural pathway, relative to the patient's activity level when the electrical signal is not directed to the neural pathway.
- 6. The method of claim 1, further comprising controlling delivery of the electrical signal to the neural pathway in response to receiving an input.
  - 7. The method of claim 6 wherein the input includes a time input.
- 8. The method of claim 7 wherein controlling delivery of the electrical signal includes directing the electrical signal during normal waking hours and halting the electrical signal during normal sleeping hours.
- 9. The method of claim 7 wherein controlling delivery of the electrical signal includes directing the electrical signal during a first predetermined period of time and halting the electrical signal during a second predetermined period of time.
- 10. The method of claim 7 wherein controlling delivery of the electrical signal includes directing the electrical signal for a predetermined cumulative maximum period of time per day.
- 11. The method of claim 7 wherein controlling delivery of the electrical signal includes preventing the initiation of the electrical signal for a predetermined period of time after the previous initiation of the electrical signal.

12. The method of claim 6 wherein receiving an input includes receiving an input corresponding to a patient activity level.

- 13. The method of claim 12 wherein the input includes an accelerometer input.
- 14. The method of claim 12 wherein the input includes a heart rate input.
- 15. The method of claim 12 wherein controlling delivery of the electrical signal includes:
  - directing the signal or increasing an inhibitory effect of the signal on the afferent neural pathway in response to an increase in patent activity level; and halting the signal or decreasing an inhibitory effect of the signal on the afferent neural pathway in response to a decrease in patent activity level
  - 16. The method of claim 6 wherein the input includes a manual patient input.
- 17. The method of claim 1, further comprising providing a patient alert in response to a detected condition.
- 18. The method of claim 17 wherein providing a patient alert includes providing a patient alert in response to a detected pulmonary condition.
- 19. The method of claim 18 wherein providing a patient alert includes providing a patient alert in response to a detected patient breathing rate.
- 20. The method of claim 18 wherein providing a patient alert includes providing a patient alert in response to a detected patient blood oxygen level.
- 21. The method of claim 17, further comprising controlling delivery of the electrical signal in response to the detected condition.

- 22. The method of claim 1, further comprising;
- automatically enabling delivery of the electrical signal when the patient is in a first state; and
- automatically disabling delivery of the electrical signal when the patient is in a second state different than the first state.
- 23. The method of claim 22 wherein the first state corresponds to a first patient blood oxygen level, and wherein the second state corresponds to a second blood oxygen level lower than the first.
- 24. The method of claim 22, further comprising automatically directing the electrical signal in response to an indication that delivery of the electrical signal is enabled.
- 25. The method of claim 22, further comprising only directing the electrical signal in response to (a) an indication that delivery of the electrical signal is enabled, and (b) a concurrent patient request for signal delivery.
- 26. The method of claim 1 wherein the carotid body chemoreceptor is one of two carotid body chemoreceptors spaced apart laterally on opposite sides of the patient's midline, and wherein implanting the signal delivery device includes implanting the signal delivery device unilaterally to be in signal communication with the afferent neural pathway of one of the carotid body chemoreceptors.
- 27. The method of claim 26, further comprising resecting the afferent neural pathway of the contralateral chemoreceptor.
- 28. The method of claim 1 wherein the carotid body chemoreceptor is one of two carotid body chemoreceptors spaced apart laterally on opposite sides of the patient's midline, and wherein implanting the signal delivery device includes implanting one or more

signal delivery devices to be in signal communication with afferent neural pathways of both the carotid chemoreceptors.

- 29. The method of claim 1, further comprising positioning the electrical signal delivery device to direct signals preferentially to chemoreceptor neurons over baroreceptor neurons.
- 30. The method of claim 29, further comprising electrically insulating structures neighboring the afferent neural pathway from effects of the signal.
  - 31. The method of claim 1, further comprising: assessing a baroreceptor function level of the patient;
  - if the patient's sensitivity to loss of barosense function has a first level, positioning the electrical signal delivery device to direct signals preferentially to chemoreceptor neurons over baroreceptor neurons; and
  - if the patient's sensitivity to loss of barosense function has a second level less than the first, positioning the electrical signal delivery device to direct signals without preference to affecting chemoreceptor neurons over baroreceptor neurons.
- 32. The method of claim 1 wherein directing an electrical signal from the electrical signal delivery element to the neural pathway includes directing the signal to a portion of the pathway at or superior to the patient's carotid body.
- 33. The method of claim 32 wherein directing an electrical signal from the electrical signal delivery element to the neural pathway includes directing the signal to a portion of the pathway no greater than 5 centimeters superior to the patient's carotid body
- 34. The method of claim 1 wherein directing an electrical signal includes directing a varying electrical signal at a frequency of from about 1KHz to about 10KHz, a

pulse width of from about 10 microseconds to about 100 microseconds, and a pulse amplitude of up to about 12 volts or up to about 10 milliamps.

- 35. The method of claim 1, further comprising varying a degree to which the afferent signals are blocked by varying an amplitude of the electrical signal.
  - 36. The method of claim 1, further comprising:

halting the signal;

- identifying a persistence period occurring after the signal is halted and during which dyspneic symptoms remain at least reduced; and
- based at least in part on identifying the persistence period, directing additional signals in accordance with a duty cycle having quiescent periods selected to be less than the persistence period.
- 37. A method for treating a patient, comprising:
- using a non-invasive, physiological-functioning screening procedure, assessing the patient's suitability for a dyspnea treatment regimen; and
- based at least in part on a positive result from the screening procedure, engaging the patient in the treatment regimen.
- 38. The method of claim 37 wherein assessing the patient's suitability for a dyspnea treatment regimen includes:
  - directing the patient to breathe a gas having an elevated oxygen content compared to standard air; and
  - receiving an indication from the patient identifying a change or lack of change in dyspneic symptoms, exercise capacity, or both.
  - 39. The method of claim 38 wherein the gas is at least 30% oxygen.
  - 40. The method of claim 38 wherein the gas is at least 50% oxygen.

41. The method of claim 38 wherein the gas is approximately 100% oxygen.

- 42. The method of claim 37 wherein engaging the patient in a treatment regimen includes at least reducing dyspneic sensations in the patient by directing an electrical signal from the electrical signal delivery element to the neural pathway to at least partially block afferent signals from the chemoreceptor.
- 43. The method of claim 42 wherein using a non-invasive, physiological-functioning screening procedure includes assessing a baroreceptor function level of the patient, and wherein the method further comprises:
  - if the patient's sensitivity to loss of barosense function has a first level, positioning the electrical signal delivery device to direct signals preferentially to chemoreceptor neurons over baroreceptor neurons; and
  - if the patient's sensitivity to loss of barosense function has a second level less than the first, positioning the electrical signal delivery device to direct signals without preference to affecting chemoreceptor neurons over baroreceptor neurons.
  - 44. A system for treating a patient, comprising:
  - an implantable electrical signal delivery element configured to direct electrical signals to a patient's neural pathway;
  - a power source coupled to the signal delivery element;
  - an oxygen sensor configured to transmit a signal corresponding to the patient's blood oxygen level; and
  - a controller coupled to the power source and the oxygen sensor, the controller being programmed with instructions that, when executed:
    - direct an inhibitory electrical signal to the signal delivery element when the oxygen sensor indicates a blood oxygen level above a first threshold; and
    - cease directing the inhibitory signal when the oxygen sensor indicates a blood oxygen level below a second threshold.

45. The system of claim 44 wherein the first and second thresholds are the same.

- 46. The system of claim 44 wherein the first threshold has a value in the range of from about 89% to about 93%.
- 47. The system of claim 44 wherein the second threshold has a value in the range of from about 93% to about 89%.
- 48. The system of claim 44 wherein the power source is implantable within the patient.
- 49. The system of claim 44 wherein the power source is an external power source.
- 50. The system of claim 44 wherein the signal delivery element includes a cuff electrode.
- 51. The system of claim 44 wherein the signal delivery element includes an elongated lead having a plurality of axially spaced-apart ring contacts.

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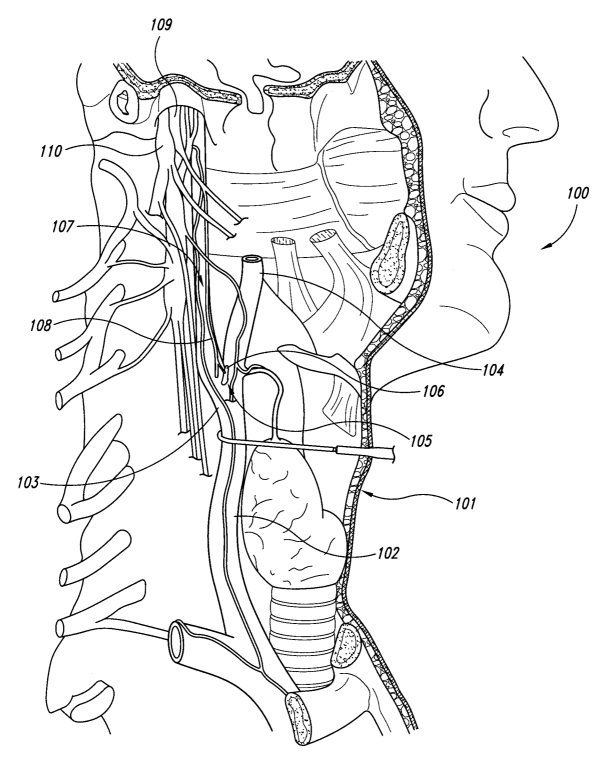


Fig. 1

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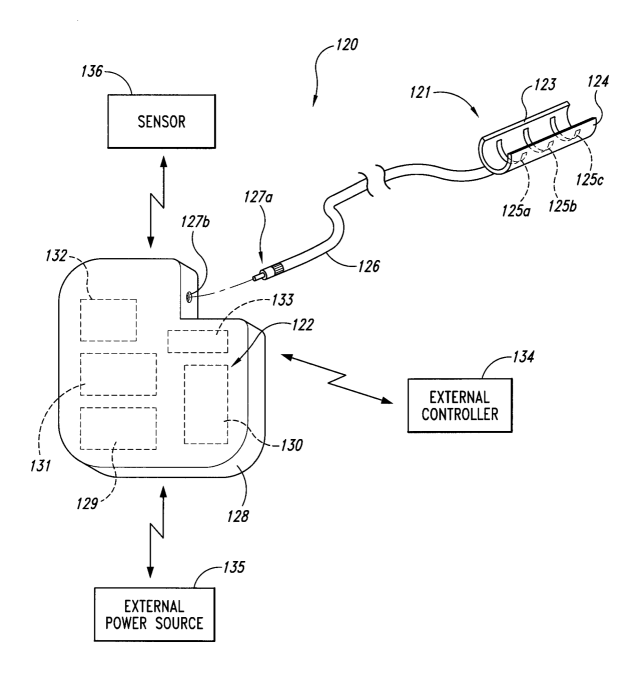
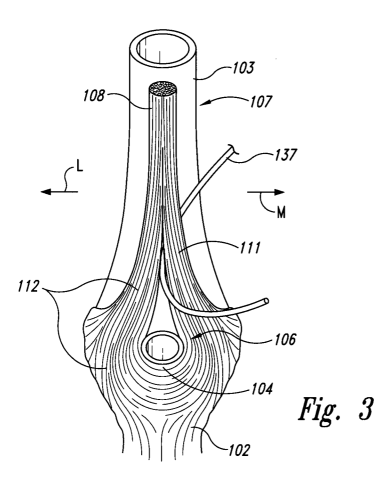
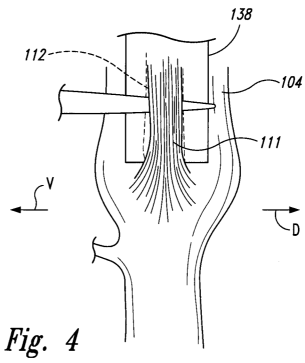


Fig. 2

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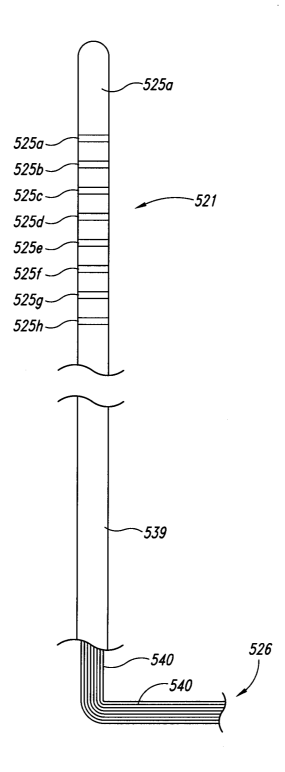


Fig. 5

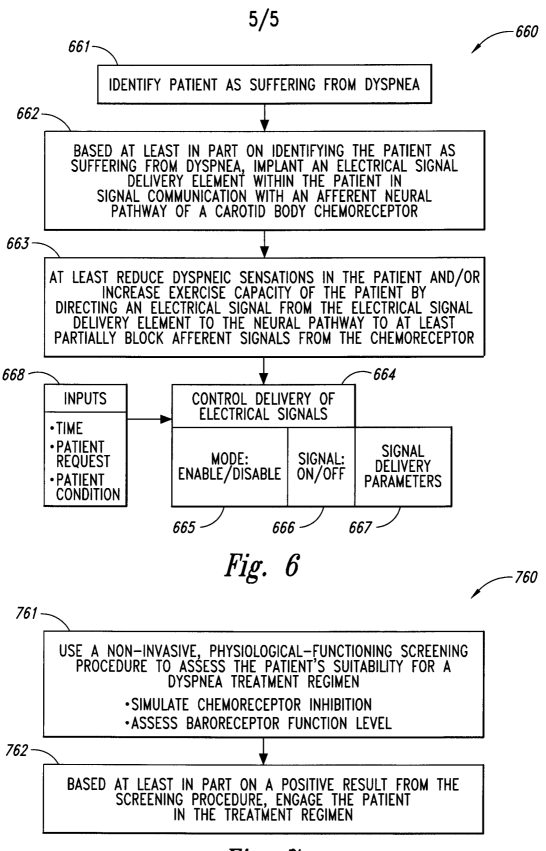


Fig. 7

#### INTERNATIONAL SEARCH REPORT

International application No PCT/US2009/053190

A CLASSIFICATION OF SUBJECT MATTER IPC(8) - A61N 1/372 (2009.01) USPC - 607/42 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)  IPC(8) - A61N 1/36, 1/372 (2009 01)  USPC - 607/2, 607/42				
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 practicable, search terms used)  PatBase, Google Patents, Google				
C DOCUMENTS CONSIDERED TO BE RELEVANT				
Category*	Citation of document, with indication, where ap	propriate, of the relevant passages	Relevant to claim No	
x -	US 2006/0282131 A1 (CAPARSO et al) 14 December 3	2006 (14 12 2006) entire document	1, 6, 16, 26, 28-30, 32, 33, 35	
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Y US 4,960,133 A (HEWSON) 02 October 1990 (02 10 1990) entire document		51		
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