



(11) **EP 2 544 590 B1**

(12) **EUROPEAN PATENT SPECIFICATION**

(45) Date of publication and mention
of the grant of the patent:
14.02.2018 Bulletin 2018/07

(21) Application number: **11710603.9**

(22) Date of filing: **10.03.2011**

(51) Int Cl.:
A61B 5/00 (2006.01)

(86) International application number:
PCT/US2011/027956

(87) International publication number:
WO 2011/112843 (15.09.2011 Gazette 2011/37)

(54) **SYSTEM FOR IDENTIFYING A LOCATION FOR NERVE STIMULATION**

SYSTEM ZUR IDENTIFIZIERUNG EINER STELLE ZUR NERVENSTIMULATION

SYSTÈME D'IDENTIFICATION D'UN EMPLACEMENT POUR STIMULATION NERVEUSE

(84) Designated Contracting States:
**AL AT BE BG CH CY CZ DE DK EE ES FI FR GB
GR HR HU IE IS IT LI LT LU LV MC MK MT NL NO
PL PT RO RS SE SI SK SM TR**

(30) Priority: **12.03.2010 US 313406 P**

(43) Date of publication of application:
16.01.2013 Bulletin 2013/03

(60) Divisional application:
18150813.6

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SU-A1- 925 349 US-A1- 2009 187 124

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Description

BACKGROUND

[0001] Advanced imaging techniques, such as magnetic resonance imaging (MRI), have revolutionized diagnosis and treatment of various patient maladies. In particular, such techniques permit observation of internal structures, such as soft tissues, that were previously unidentifiable in traditional radiographic techniques (such as X-ray). As such, a wide variety of internal organs, tissues, connective tissue is now viewable along with the previously viewable bony structures. Using the MRI or other advanced imaging techniques (such as computer tomography, CT), physicians can readily diagnosis or exclude many conditions. US 2009/0187124 A1 discloses low levels of sensory stimulation to the throat area in a patient in order to stimulate the nerves underneath. Video fluoroscopy is thereafter turned on to obtain time series plots of the x and y points of the hyoid bone and the y coordinate of the larynx.

[0002] WO 2010/014260 A1 discloses electrical stimulation in the head-and-neck region.

[0003] WO 2005/013805 A2 discloses the assessment of muscle activity using an electromyogram (EMG) following nerve stimulation.

BRIEF DESCRIPTION OF THE DRAWINGS

[0004] The invention is defined in independent claim 1. Preferred embodiments are provided in the dependent claims. Aspects and features of the present invention will be appreciated as the same becomes better understood by reference to the following detailed description of the embodiments of the invention when considered in connection with the accompanying drawings, wherein:

Figure 1 is a schematic illustration of a system and method for navigation via images and via neuromuscular responses from stimulation;

Figure 2 is a flow diagram of a method of navigating placement of a stimulation element within a body;

Figure 3 is a block diagram of a system for navigation;

Figures 4-5 are a side plan view schematically illustrating an interface of a method and system for graphical differentiation between a target muscle and an activated muscle;

Figure 6A is a flow diagram of a method of identifying a stimulation site via an array of sensor probes;

Figure 6B is a block diagram of a nerve-muscle response index;

Figure 7 is a side view schematically illustrating a method of sensing neuromuscular responses via sensor probes in a head region of a patient;

Figure 8 is a top plan view of a sensor probe;

Figure 9 is a side plan view of a sensor probe;

Figure 10 schematically illustrates a system and user interface for navigating a pathway via images in relation to neuromuscular responses; and

Figure 11 schematically illustrates a system and user interface for navigating and tracking a pathway of a stimulation element.

Figure 12 is a schematic illustration of a system and user interface for navigating a path to implant a stimulation element and simultaneously evaluate, via graphic displays, in real-time whether the path or stimulation site is efficacious.

Figure 13 is a schematic illustration of a multi-balloon probe and display of pressure sensed via the probe.

Figure 14 is a schematic illustration an illumination system for illuminating anatomical features of a patient.

DETAILED DESCRIPTION OF THE EMBODIMENTS

[0005] The following detailed description is merely exemplary in nature and is not intended to limit the present general inventive concept or the application and uses of the present general inventive concept. Furthermore, there is no intention to be bound by any expressed or implied theory presented in the preceding technical field, background, or the following detailed description.

[0006] Embodiments of the present general inventive concept are directed to enhancing surgical navigation via combining, in a graphical user interface, image-based visualization of the soft and hard tissues of a relevant body region simultaneous with graphical representation (within the image-based visualization) of muscle activation from a stimulation signal applied at a target site. In some embodiments, in concert with visualization of a stimulation element within the image-based navigation interface, a stimulation path analyzer makes a quantitative evaluation of muscle responses from nerve stimulation to determine a precise location of the stimulation element relative to pertinent soft tissues (such as muscles, nerves, and major circulatory vessels) and also relative to bony structures, which serve as reference points. With this combination, one can determine and visually represent a location of a stimulation element in a body region in real time using image-navigation tools simultaneous with functional evaluation of the electrode placement (e.g. does the tongue protrude upon stimulation

from the stimulation element? is the proper muscle group activated ?) and simultaneous with quantitative evaluation of the stimulation location, such as nerve conduction and muscle response information (e.g. electromyography and/or compound muscle action potential).

[0007] In some embodiments, the quantitative evaluation information of which muscle is activated is expressed within the image-based navigation interface via highlighting a graphical representation of the activated muscle among other muscles and anatomical structures. In one embodiment, the system differentiates between target muscles and non-target muscles in the image-based navigation interface via color, shading, and/or other patterns. Accordingly, the system greatly simplifies the role of an operator by providing real-time representations of the effect of a particular location of a stimulation element. In one embodiment, this arrangement is of particular benefit to facilitate percutaneous placement of an implantable nerve stimulation electrode usable in a method of treating obstructive sleep apnea or other sleep-breathing disorders.

[0008] These embodiments, and additional embodiments, are described in more detail in association with Figures 1-14.

[0009] Figure 1 is diagram schematically illustrating a system and method for navigation via images and via neuromuscular responses from electrical stimulation. In one embodiment, system 20 has particular application to facilitate a minimally invasive, percutaneous or transvenous placement of an implantable nerve stimulation electrode and thereby avoid a traditional, highly invasive cut-down access procedure.

[0010] As illustrated by Figure 1, system 20 includes interface 30, image-based navigation module 50, and neuromuscular response evaluator 52. In general terms, interface 30 includes a graphical user interface configured to simultaneously display images and neuromuscular responses in an integrated manner. User controls 34 enable a user to operate and control at least some functions of image-based navigation module 50, parameters of interface 30, and/or neuromuscular response evaluator 52. In some embodiments, on screen controls would be supplemented by one or more additional control mechanisms, such as foot pedals, voice recognition commands, a sterilized navi-pad.

[0011] In one embodiment, user controls 34 include laser projection onto the patient and/or onto tissues for identifying anatomical features, landmarks, and distances. In some embodiments, multiple lasers (each having a different color) are used to help differentiate between multiple anatomical features, landmarks, and distances, as further described later in association with at least Figures 10 and 14.

[0012] As illustrated by Figure 1, interface 30 displays an image 40 of relevant anatomical features of a body portion of patient 22. In addition, as further described below, superimposed on the image 40 are muscle identifiers 42, 44 and a nerve identifier 46, which together

indicate which nerve is being stimulated, where is it being stimulated, and which muscles are activated as a result. In one aspect, nerve identifier 46 is operable to separately identify multiple nerves to differentiate within interface 30 each respective nerve, as well as differentiating identification of relevant branches of a nerve. In some embodiments, interface 30 additionally indicates a target nerve and a target muscle innervated by the target nerve, such that interface 30 enables confirmation of whether the target muscle or a different muscle has been activated via stimulation of a nerve, as will be further described later in association with Figures 4-11.

[0013] In one embodiment, the image-based navigation module 50 includes a tracking function 60, an imager function 62, a records function 64, and an anatomy map function 66.

[0014] In general terms, the imager function 62 obtains and/provides images of a body portion of patient 22. In some embodiments, the image is obtained via an imaging system 80 (not shown to scale for illustrative purposes) including a transmitter 82 and receiver 84, which function together to capture internal images of patient 22. The imaging system 80 includes, but is not limited to, any one of a magnetic resonance imager (MRI), computer tomography (CT) unit, fluoroscopic imager, endoscopy, etc., or combinations thereof, as is further illustrated in Figure 3. In one example, endoscopy is used to observe evoked responses.

[0015] Records function 64 of image-based navigation module 50 enables access to one or more stored image records of patient 22, via an electronic medical records (EMR) system or other resources as further noted in association with Figure 3. Anatomy map function 66 enables access to general anatomical maps and images, suitably scaled, to overlay or superimpose relative to obtained images of patient 22. In some embodiments, a patient-specific image (such as CT or MRI), is overlaid onto, and synchronized relative to, a generalized anatomy map using commercially available image analysis techniques, such as those provided via Vital Images, Inc. of Minnetonka, Minnesota. The patient-specific images are obtained via known systems such as electronic medical records (EMR) and/or a picture archiving communication system (PACS). In some embodiments, the images are obtained and/or communicated according to a Health Level 7 standard for electronic medical records.

[0016] In this way, a generally comprehensive picture of relevant anatomical features of a body portion of patient 22 are provided in interface 30 (FIG. 1) to facilitate identifying target nerves and muscles when locating a target stimulation site for an electrode (or electrode array) and when mapping a percutaneous access pathway to that target stimulation site.

[0017] By utilizing imager function 62, records function 64 and/or anatomy map function 66, the tracking function 60 enables visually tracking (via interface 30) a location of stimulation element 90 and/or support instruments used to guide and place stimulation element 90. In some

embodiments, the stimulation element 90 is a stimulation test tool while in other embodiments, the stimulation element 90 is an implantable electrode.

[0018] In one embodiment, as shown in Figure 1, the relevant body region of patient 22 displayed in interface 30 includes an upper respiratory system including the mouth to enable visualizing the tongue, its underlying and surrounding muscle groups, related nerve pathways, and supporting bony structures. In some embodiments, visualization of these detailed structures (and their inter-relationship) is used to treat sleep-disordered breathing, such as obstructive sleep apnea.

[0019] In general terms, the neuromuscular response evaluator 52 of system 20 (Figure 1) is configured to apply a stimulation signal through stimulation element 90 via a stimulation engine 70 according to stimulation parameters 72. The produced neuromuscular response is sensed with probes 92 (for example, probes shown in Figures 7-9) and fed to a stimulation path analyzer 74. Via an electromyography function 76 and a compound muscle action potential function (CMAP) 78, the stimulation path analyzer 74 differentiates whether the activation of the muscle occurred because of stimulation of a nerve innervating that activated muscle or because of direct electrical stimulation of the activated muscle. In some embodiments, at least some of the functions of the response evaluator 52 are provided via a nerve integrity monitor. In this regard, in one embodiment, the nerve integrity monitor comprises at least substantially the same features and attributes as the nerve integrity monitor described in U.S. Patent 6,334,068, entitled INTRA-OPERATIVE NEUROELECTROPHYSIOLOGICAL MONITOR, issued on December 25, 2001. In other embodiments, other commercially available nerve integrity monitors or an equivalent array of instruments (e.g., a stimulation probe and electromyography system) are used to apply the stimulation signal and evaluate the response of the muscle innervated by the target nerve.

[0020] In some embodiments, a plurality of sensor probes is placed at and near muscles to be activated. In one embodiment, the sensor probes comprise fine wire needle electrode typically used in nerve integrity monitoring. With the sensor probes in place and using interface 30, one can observe which muscles are activated upon stimulation of the different sites and via path analyzer 74, one can systematically reveal which stimulation site will result in activation of the desired muscle. This arrangement is described further in association with Figures 6A-6B. Accordingly, in some embodiments, system 20 includes a probe docking module 79 to establish electrical communication between the sensor probes 92, neuromuscular response evaluator 52, and interface 30. Via programming, the probe docking module 79 associates each separate probe 92 with a specific muscle of the patient 22 so that when the muscle associated with a particular probe is activated, the interface 30 can visually highlight that muscle on the displayed image 40. For example, highlighted region 42 of image 40 is associated

with probe P2 while highlighted region 44 of image 40 is associated with probe P1. In this way, the interface 30 correlates the location of the sensor probes 92 with the anatomical features of the patient 22 in the displayed image 40. In some embodiments, this arrangement is facilitated via a nerve-muscle index 294, which is further described later in association with Figures 3 and 6B.

[0021] In some embodiments, the stimulation element 90 includes a stimulation test tool configured to be inserted percutaneously into a body of patient 22 to releasably engage a potential stimulation site and enable a stimulation signal to be applied at that site, such as a target nerve. In other embodiments, the stimulation element 90 includes a cuff electrode (or other electrode configuration) configured to be implanted in a secured relationship relative to a target stimulation site of a nerve in order to provide long-term implantable stimulation therapy via activation of an innervated muscle. For example, in one embodiment, the target nerve is a hypoglossal nerve and the innervated muscle is the genioglossus muscle. In some embodiments, the target nerve is a particular branch (or branches) of the hypoglossal nerve. For example, in one example the target nerve includes one or more particular lateral branches or medial branch of the hypoglossal nerve, such that the innervated muscle includes the styloglossus muscle, the hyoglossus muscles, the geniohyoid muscle, and/or the genioglossus muscle.

[0022] In one instance, the cuff electrode is delivered to the target stimulation site of the nerve via minimally invasive, percutaneous access (which avoids the more traditional, highly invasive cut-down access procedure). As more fully described herein, such percutaneous access for cuff electrode is guided via both the image-based navigation module 50 and neuromuscular response evaluator 52. In one embodiment, one such percutaneous access method is described and illustrated in application serial number 61/165,110, filed March 31, 2009, and titled PERCUTANEOUS ACCESS METHOD AND SURGICAL NAVIGATION TECHNIQUES. It will be understood that in some embodiments, other minimally invasive methods, such as microendoscopic delivery techniques, are used to place the implantable cuff electrode in cooperation with the guidance of the image-based navigation module 50 and neuromuscular response evaluator 52.

[0023] In yet other embodiments, transvenous access delivery methods of a stimulation electrode are used to place the stimulation electrode at a target stimulation site relative to target nerve, and that even this transvenous delivery is aided or guided via image-based navigation module 50 and neuromuscular response evaluator 52 in accordance with principles of the present disclosure, or in accordance with other imaging techniques. In one embodiment, one such transvenous access method is described and illustrated in PCT application serial number PCT/US2009/059060, filed September 30, 2009, and titled TRANSVENOUS METHOD OF TREATING SLEEP APNEA.

[0024] With this arrangement in mind, the produced neuromuscular responses are visually represented on interface 30 via a highlighted portion of image 40, such as via color highlighting or shading. As shown in Figure 1, the highlighted portions can take particular shapes and sizes, such as rectangular shapes, that may or may not approximate the general shape and size of a subject's anatomical features. In some embodiments, the location, size, and/or shape of the highlightable portions of interface 40 (such as the colored or shaded portions corresponding to a particular anatomical feature) is established at the time that sensor probes are placed relative to the patient's anatomy.

[0025] In the non-limiting example illustrated by Figure 1, a highlighted portion 42 of interface 30 corresponds to a deduced location of an activated muscle (i.e., the muscle actually activated, associated with probe P2) while a highlighted portion 44 (associated with probe P1) corresponds to a target muscle that is desired to be activated. Identifier 48 highlights the location at which stimulation takes place on nerve 46. Moreover, as the system 10 tracks the real-time location of the stimulation element 90, the location of the stimulation element 90 is dynamically represented via identifier 48 as part of displayed image 40.

[0026] With this arrangement, interface 30 simultaneously combines the image-based navigation information with the neuromuscular response information to provide a real-time indication of which muscles are activated via stimulation (and to what degree) and a real-time indication of the location of the stimulation element causing the muscle activation. Using this information, interface 30 assists a physician in maneuvering the stimulation element to a target stimulation site for an innervated muscle to be activated via stimulation.

[0027] Figure 2 is a flow diagram that schematically illustrates a method 100 of placing a stimulation element at a target stimulation site. In some embodiments, method 100 is performed using substantially the same the systems, components, modules, functions, etc. as described in association with Figures 1 and 3-11. In other embodiments, method 100 is performed using other systems, components, modules, and/or functions available to those skilled in the art.

[0028] As illustrated by Figure 2, method 100 includes advancing a stimulation element within a body, via an image-based navigation interface, to a target stimulation site, as at 102. In one embodiment, the stimulation element is advanced percutaneously while in other embodiments, the stimulation element is guided transvenously or through other access techniques. Each potential stimulation site is tested via observing neuromuscular responses produced from a stimulation signal (transmitted by the stimulation element), as at 104. In particular, a neuromuscular muscle response is determined separately for each potential target nerve stimulation site. In a superimposed combination with images of the location of the stimulation element among relevant anatomical

structures (soft and hard tissues), the produced neuromuscular responses are displayed as a graphic identification of which muscle or multiple muscles were activated from a respective stimulation site, as at 106. In some embodiments, these muscle responses are recorded for later comparison to subsequent muscle responses. Further details for performing aspects of method 100 are described below.

[0029] Figure 3 is a block diagram of a system 200 for providing image-based navigation of a stimulation element within a body while substantially simultaneously graphically identifying a target stimulation site (via the image-based navigation interface) via neuromuscular response evaluation. As illustrated by Figure 3, system 200 includes image-based navigation module 210, interface 212, and neuromuscular response evaluator 214, memory 216, and controller 218. In some embodiments, image-based navigation module 210, interface 212, and neuromuscular response evaluator 214 comprise at least substantially the same features and attributes as image-based navigation module 50, interface 30, and neuromuscular response evaluator 52, as previously described in association with Figure 1.

[0030] In one embodiment, the interface 212 comprises a graphical user interface 280 configured to display, and enable operation of, the various parameters, components, functions, and modules of system 200. Accordingly, via interface 212, system 200 as illustrated in Figure 3 represents the display of the respective parameters, components, functions, monitors, managers, and/or modules as well as representing the ability to activate or operate those respective parameters, components, functions, monitors, managers, and/or modules.

[0031] As illustrated by Figure 3, image-based navigation module 210 includes image module 230, records function 64, anatomy map function 66, and fiducials function 256. The image module 230 obtains and provides to interface 212 one or more of a MRI image 240, a CT image 242, a fluoroscopy image 244, and/or an endoscopic image 245 of a pertinent region of a patient. In some embodiments, aspects of different types of images are synthesized or combined to yield a single hybrid image for display at interface 212. In one aspect, anatomical structures (such as, but not limited to, nerves and muscles) are located manually, and highlighted to the user at interface 212. For example, an operator can review available images of a patient body region and then select or indicate a particular structure of interest. In one non-limiting example, an operator selects a mandible within several different images or different image types so that the images can be integrated in a manner that still visually displays the anatomical structure of interest, i.e., the mandible. In another non-limiting example, an operator selects a nerve of interest from the soft tissues rendered by an MRI or selects the tongue or other upper airway muscles of interest and associates those with placed sensor/monitors. Records function 64 and anatomy function 66 comprise substantially the same features and at-

tributes as previously described in association with Figure 1.

[0032] Fiducials function 256 provides for visually tracking one or more fiducial markers that are visibly distinct from tissues on a displayed image to provide references points that are independent of the tissues and anatomical structures of interest. In some embodiments, locations for placing fiducial markers include one or more of an anterior point of the mandible bone, the hyoid bone, the skull, and/or cervical vertebrae. Of course, other or additional anatomical structures can be used as a fiducial reference point by placing a marker there. These fiducial markers have shapes and/or sizes selected to provide an objective orientation within and relative to the anatomical structures of the displayed image (e.g., displayed image 40 in Figure 1) of interface 212. In one embodiment, during an imaging session to obtain images of relevant portions of patient anatomy, a radiopaque marker is placed at a desired implant site. Upon later use of fluoroscopy during placement of an implantable stimulation electrode, the radiopaque marker is easily identifiable and used as a general location marker to facilitate placement of the electrode.

[0033] Interface 212 of system 200 includes graphical user interface 280 (described above). In some embodiments, interface 212 includes target nerve identifier 282, target muscle identifier 284, actual nerve identifier 286, and actual muscle identifier 288, path marker function 290, target spot marker function 292, audio identifier module 300, and/or visual identifier module 320.

[0034] The actual nerve identifier 286 provides real-time visual identification of activated nerve(s) while the actual muscle identifier 288 provides real-time visual identification of activated muscle(s). Meanwhile, the target nerve identifier 282 and the target muscle identifier 284 provide time-independent display of a nerve intended to be stimulated and its innervated muscles that are intended to be activated. With these functions 282, 284 an operator designates on displayed image (e.g. displayed image 40 in Figure 1) the respective nerve and muscle that are targeted. This designation is made via touching the respective portions of displayed image 40 (when interface 30, 212 includes a touch screen) or otherwise selecting the desired nerve and muscle via user controls 34 (such as a cursor navigation/selection or keypad entry).

[0035] In some embodiments, activation of path marker function 290 causes interface 212 to display a path through which the stimulation element 90 is expected to travel percutaneously (or otherwise internally in the body) as mapped out by an operator on a displayed image (e.g. displayed image 40 in Figure 1) based on a probable target nerve stimulation site or a confirmed target nerve stimulation site. In some embodiments, target spot marker function 292 is provided to allow selective designation on a displayed image (e.g. displayed image 40 in Figure 1) of a general or specific intended target site for placement of an implantable electrode. The accuracy of the

target spot is confirmed or denied via the corroboration of neuromuscular response evaluation, and the target spot marker can be adjusted accordingly until an accurate target spot is located. Moreover, via target spot marker function 292, travel along the path is confirmed or evaluated by applying stimulation and observing which muscles are activated.

[0036] In some embodiments, the audio identifier module 300 comprises a hit function 302, a miss function 304, and/or an intensity function 306. In general terms, the audio identifier module 300 is configured to provide auditory identification of whether the intended target nerve is stimulated or not. In particular, feedback from which innervated muscle or muscles have been activated (via neuromuscular response evaluator 214) is communicated via an auditory signal, auditory words, or other easily recognizable auditory information. In some embodiments, the auditory alert occurs simultaneous with the period of activation of the muscle being stimulated. This auditory information communicates to the user whether the target nerve was activated or not. In some embodiments, this information is communicated as an audible alert by a spoken word, such as "hit", for activation of the target nerve (via hit function 302) and as a spoken word, such as "miss", for activation of a nerve other than the target nerve (via miss function 304).

[0037] The intensity function 306 of audio identifier module 300 provides an audio-based indication of an intensity of the muscle activation produced via nerve stimulation. In one aspect, this audio report is used to evaluate the intensity of the muscle response based on a given target nerve stimulation site. This audio-intensity information communicates to the user a relative degree of stimulation of the target nerve, and relative effectiveness of a particular stimulation site on the nerve given a nominal set of stimulation parameters. In one example, a relatively low volume audio sound indicates a relatively low-to- moderate muscle activation to the operator while a relatively higher volume audio sound indicates a relatively high or robust degree of muscle activation. In addition, it will be understood that, depending upon the stimulation parameters, the relative effectiveness of a particular stimulation site might vary. Accordingly, in some embodiments, at each particular potential stimulation site along a nerve, the stimulation parameters are varied in an organized manner to fully evaluate the effectiveness of that potential stimulation site in activating an innervated muscle. The intensity of muscle activation for each different combination of stimulation parameters and/or stimulation site is reported to the operator via audio cues by audio identifier module 300.

[0038] To maximize the available information for decision making during percutaneous delivery of an implantable electrode or during initial determination of a stimulation site, in some embodiments the audio identifier module 300 is used in concert with the visual identifier module 320. However, it is understood that in other embodiments, just one of these respective identifier mod-

ules 300, 320 can be used. In still other embodiments, the audio identifier module 300 and the visual identifier module 320 are used simultaneously, but independent from each other.

[0039] With further reference to Figure 3, in some embodiments, the visual identifier module 320 comprises a color function 322, a shading function 324, a numerical intensity function 326, a directional function 328, and/or a traced path function 330. In general terms, the visual identifier module 320 is configured to provide visual identification in interface 212 (such as interface 30 of Figure 1) of whether the intended target nerve is stimulated or not. In particular, feedback from which innervated muscle or muscles have been activated (via neuromuscular response evaluator 214) is communicated via visual signals, visual words or symbols, or other easily recognizable visual information. This visual information communicates to the user whether the target nerve was activated or not. While this visual information can include graphical display of a signal normally associated with electromyography, in some embodiments, the displayed visual information omits ordinary waveform signals in favor of words, symbols, and other more readily identifiable information about the status of a particular muscle. In other words, visual identifier module 320 converts or evaluates the quantitative data of the EMG response and/or CMAP response into a qualitative form that quickly communicates, in a visually intuitive manner, to the operator which muscle has been activated and/or the intensity of the muscle response.

[0040] In one example, as noted above in association with Figure 1, the visual identifier module 320 provides a graphical representation by which an activated muscle becomes highlighted on an image in a navigation interface such that the user can immediately recognize which muscle has been activated.

[0041] The color function 322 of visual identifier module 320 provides a color-based indication of which muscle is activated via nerve stimulation and/or a degree of muscle activation produced via nerve stimulation. This visual-intensity information communicates to the user a relative degree of stimulation of the target nerve, and relative effectiveness of a particular stimulation site on the nerve given a nominal set of stimulation parameters. In some embodiments, one color identifies a target nerve to be stimulated or a target muscle to be activated while a second color identifies which nerve is stimulated and/or which muscle is activated via a stimulation signal. In one embodiment, when the stimulated nerve matches the target nerve, a third color overlays the graphical representation of that target nerve to indicate that the target nerve was successfully located and was stimulated. Similarly, when the activated muscle matches the target muscle, the third color overlays the graphical representation of that target muscle to indicate that the target muscle was successfully located and was stimulated. It will be understood that the same color may be used for both nerves and muscles, or in other embodiments, that one set of

colors is used to exclusively represent nerves and another different set of colors is used to exclusively represent muscles. Of course, it will be understood that other designations of colors can be used to communicate on a displayed image (e.g. displayed image 40 in Figure 1) which muscles are actually activated via stimulation, which muscles are intended to be activated, and/or the degree of activation.

[0042] The shading function 324 of visual identifier module 320 provides a visual-based indication of which muscle is activated via nerve stimulation and/or a degree of muscle activation produced via nerve stimulation. A type of shading or a relative darkness of shading communicates to the user a relative degree of stimulation of the target nerve, and relative effectiveness of a particular stimulation site on the nerve (given a nominal set of stimulation parameters). In some embodiments, one type of shading is used to identify muscles and another type of shading is used to identify nerves.

[0043] In cooperation with the color function 322 and/or the shading function 324, the numerical intensity function 326 of visual identifier module 300 provides a visual-based numerical indication of an intensity of the muscle activation produced via nerve stimulation. This visual-intensity information communicates to the user a relative degree of stimulation of the target nerve, and relative effectiveness of a particular stimulation site on the nerve given a nominal set of stimulation parameters.

[0044] It will be understood that, depending upon the stimulation parameters, the relative effectiveness of a particular stimulation site might vary. Accordingly, in some embodiments, at each particular potential stimulation site along a nerve, the stimulation parameters are varied in an organized manner to fully evaluate the effectiveness of that potential stimulation site in activating an innervated muscle.

[0045] In one embodiment, the directional function 328 enables visual indication of directional movement of a stimulation element 90. However, in some embodiments, the directional function 328 provides a suggested direction in which to move the stimulation element. In this latter arrangement, the path analyzer 374 of neuromuscular response evaluator 214 evaluates the positive or negative outcome of the most recent stimulation sites (and the parameters of stimulation at those sites) and based on any recognized trends or patterns (from prior stimulation trials), then communicates a suggested direction of movement of the stimulation element. In one embodiment, the suggested direction is displayed graphically as a directional arrow on the displayed image (e.g. display image 40 in Figure 1).

[0046] The traced path function 330 enables visual indication of a path of different previously tested stimulation sites and/or display of a path through which a stimulation element 90 is maneuvered through a body portion on the way to or from a stimulation site (or other target location).

[0047] With further reference to Figure 3, in some embodiments, interface 212 includes a probe module 340

configured to track (based on a location of relevant EMG sensor probes and/or piezoelectric accelerometers) a target muscle, non-target nerves, and non-target muscles. In one embodiment, a target muscle parameter 346 of probe module 340 tracks, via a plurality of sensor probes configured to be removably coupled relative to a corresponding plurality of potentially activatable muscles, each potentially activatable muscle that is innervated by one of the potential target nerve stimulation sites. A non-target nerve parameter 344 of probe module 340 tracks, via a plurality of sensor probes configured to be removably coupled relative to a corresponding plurality of potentially activatable muscles, each potentially activatable muscle that is physically adjacent one of the potential target nerve stimulation sites and that is not targeted to be activated. Finally, a non-target muscle parameter 342 of probe module 340 configured to track, via a plurality of sensor probes configured to be removably coupled relative to a corresponding plurality of non-target muscles, each non-target muscle that is physically adjacent one of the target muscles.

[0048] In general terms, the neuromuscular response evaluator 214 of system 200 enables stimulation of nerves and muscles to identify a nerve stimulation site. In one embodiment, the neuromuscular response evaluator 214 includes a stimulation parameters module 350, a response module 352, and a stimulation element module 354.

[0049] In general terms, the stimulation parameters module 350 provides for selection of the various parameters of a nerve stimulation signal. In one embodiment, the stimulation parameters module 350 includes a pulse width parameter 360, frequency parameter 362, an amplitude parameter 364, a polarity parameter 366, and a duration parameter 368. Each of these parameters can be varied, as known by those skilled in the art, to achieve a desired stimulation signal on a nerve. The value of each of these parameters may vary from one stimulation site to another. It will also be understood, that in some embodiments, that a ground connection to the patient's body and a positive electrode will be used to enable, applying stimulation via a unipolar probe and identifying muscular responses.

[0050] The response module 352 provides for a mechanism to sense, record, and quantify neuromuscular responses of an activated muscle. In one embodiment, the feedback is sensed, recorded, and quantified via electromyography (EMG) via EMG function 380 while in some embodiments, the feedback is sensed, recorded, and quantified via compound muscle action potentials (CMAP) via CMAP function 382.

[0051] The stimulation path analyzer module 374 of response module 352 (of neuromuscular response evaluator 214) provides for differentiation between sources of stimulation of a sensed muscle response. In particular, a sensed muscle response can be caused by electrical stimulation of a nerve that innervates the muscle or by direct electrical stimulation of the muscle. Accordingly,

the stimulation path analyzer module 374 is configured to sort data from the electromyography function 370 according to the stimulation parameters and navigation data (from image-based navigation module 210) to make this differentiation between nerve stimulation and direct muscle stimulation. Upon trying various potential stimulation sites, the path analyzer module 374 provides a graphical summary of which locations cause nerve stimulation and which cause direct muscle stimulation. With this information, a target stimulation site is selected.

[0052] In particular, in cooperation with EMG function 370 and by observing data produced via the compound muscle action potential (CMAP) function 372, the analyzer module 374 automatically determines if there is a time delay between the electrical stimulation and the ensuing muscle activation. In particular, if direct electrical stimulation was performed on a target muscle, then no delay would be expected between that stimulation and the ensuing muscle activation. However, if electrical stimulation was performed on a target nerve, then a delay would be expected between that stimulation and the ensuing muscle activation. In one example, an approximately 100 microsecond delay would be expected for each 1 centimeter distance that the muscle is located away from the nerve. As described later in association with Figure 6B, this muscle response time is tracked via a nerve-muscle index 294.

[0053] The ability to automatically differentiate between electrical nerve stimulation and direct muscle stimulation via path analyzer 374 greatly facilitates percutaneous determination of a location of a target stimulation site of a target nerve because, in that situation, the user typically does not have direct physical sight of the nerves and/or muscles at which the stimulation tool is directed. However, via the analyzer module 384, system 200 is able to determine which nerve or muscle is being stimulated, and which muscle is activated as a result. Upon attempting stimulation at several different locations percutaneously with the stimulation element, the user determines which anatomical structure (e.g. target nerve) at which an implantable electrode should be located and at which location or position along that nerve the electrode should be secured.

[0054] Moreover, once the target stimulation site is determined, the user can further utilize system 200 to determine a pathway through which the electrode can be delivered percutaneously to arrive in the desired position at the target stimulation site.

[0055] In some embodiments, the analyzer module 374 determines a strength-distance curve for activation of target muscle groups. In one aspect, the strength-distance curve includes a graphic representation of the relationship between an intensity of the electric stimulation site and a distance between the actual stimulation site and the target nerve. Accordingly, using this strength-distance curve, a distance is estimated between the location of the stimulation element and the activated nerve (or activated muscle). This distance information is used

by the physician to determine how much further, and in which direction, to maneuver the stimulation element to arrive at the desired location or position relative to the target stimulation site. In some embodiments, this information is communicated via directional function 328 of interface 212. In addition, once a final placement of the stimulation element (e.g. cuff electrode) is determined, a strength-duration curve produced via analyzer module 374 (in association with CMAP function 372 and EMG function 370) facilitates a physician (or programming unit) in setting efficacious stimulation therapy settings to treat a physiologic condition, such as sleep-disordered breathing behaviors.

[0056] In some embodiments, system 200 includes a nerve-muscle index module 294 configured to correlate observed muscle response behavior with an associated nerve innervating the respective muscles. This index 294 can aid in placing sensor probes and/or in identifying a stimulation source upon observed muscle responses. In some embodiments, the interface stores in memory an array of nerve-muscles indices with a separate nerve-muscle index for each separate patient body portion. In addition, in some embodiments, a nerve-muscle index for a particular patient body portion is loaded into a memory of the system prior to performing a method of percutaneously advancing a stimulation element into and through that particular body portion. Further details regarding the nerve-muscle index 294 are described and illustrated later in association with Figure 6B.

[0057] In some embodiments, response evaluator 352 also comprises an ultrasound detection function 375 which is configured to detect muscle motion according to Doppler principles. For example, via an ultrasound sensor node placed underneath the chin, ultrasound detection function 375 detects motion of the genioglossus muscle in response to stimulation at or near a target stimulation site. In another example, via an ultrasound sensor node placed at the jaw, ultrasound detection function 375 detects motion of the styloglossus muscle in response to stimulation at or near a target stimulation site. Information gained via the ultrasound detection function 375 is deployed in at least muscle motion parameter 476 in nerve-muscle index 294, as later described in association with Figure 6B.

[0058] Stimulation element module 354 provides for applying electrical stimulation via a test tool 380 and/or an implantable electrode 382. In most instances, the test tool 380 is employed by the physician to first identify the target nerve(s) and target muscle(s) in cooperation with the other functions, modules of interface 212 and system 200. Thereafter, using the image-based navigation interface provided according to the present disclosure, the electrode 382 is delivered percutaneously to the target stimulation site with or without the assistance of test tool 380. It will be understood that in such a procedure, the patient is under anesthesia and without using muscle relaxants.

[0059] In either case, in some embodiments, the stim-

ulation element module 354 includes a handle control portion 384 for guiding test tool 380 or electrode 382. The handle control portion 384 includes an on/off function 386, increase function (+) 388, decrease function (-) 390, pause function 392, and mode function 394. The respective increase and decrease functions 388, 390 provide for a corresponding increase or decrease in the stimulation parameters (such as amplitude, pulse width, etc.) while the pause function 392 allows temporary suspension of the stimulation signal. The mode function 394 allows selecting various modes of stimulation, such as continuous, intermittent, or discrete (just when an input button is pressed). In some embodiments, the mode function 394 includes an automatic ramping function to determine the parameters 360-368 of module 350 of stimulation (such as amplitude, pulse width, frequency, polarity, duration) needed to recruit the targeted nerve and innervated muscles. Accordingly, this automatic ramping function provides information regarding the efficaciousness of the current stimulation location and the observed muscle responses. For example, a muscle response to a direct electrical stimulation of the muscle is generally less than a muscle response to electrical stimulation of the nerve that innervates the particular muscle. In another example, lower stimulation amplitudes required to produce a muscle response generally indicate that a stimulation element is close to the target nerve, whereas higher stimulation amplitudes required to produce a muscle response generally indicate that a stimulation element is further from the target nerve than desired.

[0060] Controller 218 comprises one or more processing units and associated memories 216 configured to generate control signals directing the operation of system 200 and its components. For purposes of this application, the term "processing unit" shall mean a presently developed or future developed processing unit that executes sequences of instructions contained in a memory, such as memory 216 or other memory. Execution of the sequences of instructions causes the processing unit to perform steps such as generating control signals. The instructions may be loaded in a random access memory (RAM) for execution by the processing unit from a read only memory (ROM), a mass storage device, or some other persistent storage. In other embodiments, hard wired circuitry may be used in place of or in combination with software instructions to implement the functions described. For example, controller 218 may be embodied as part of one or more application-specific integrated circuits (ASICs). Unless otherwise specifically noted, the controller is not limited to any specific combination of hardware circuitry and software, nor limited to any particular source for the instructions executed by the processing unit.

[0061] Figure 4 is side view schematically illustrating an image-based user interface that visually identifies muscle activation, according to an embodiment of the present invention, for use in percutaneous navigation and/or percutaneous placement of a stimulation element.

As illustrated by Figure 4, image 400 graphically depicts a body portion 402, such as a head-neck region of a patient. The body portion 402 includes bony structures 410 that support a group 420 of muscles and tissues. In the example shown, bony structures 410 include a lower jaw portion 414 that supports a tongue 412 and its underlying muscles, including (but not limited to) a styloglossus muscle 421, a genio-hyoideus muscle 422, a genioglossus muscle 423, a hyo-glossus muscle 426, and a stylo-hyoideus muscle 428.

[0062] In one embodiment, the genioglossus muscle 423 as shown in Figure 4 is a target muscle that is desired to be activated. In particular, activation of this genioglossus muscle 423 contributes to protrusion of tongue 412 forward, thereby increasing airway patency near the base of the tongue. In some instances, this protrusion produced via activating the genioglossus muscle 423 is used to treat sleep-disordered breathing behaviors, such as obstructive sleep apnea. In one embodiment, the designation of this muscle as a target muscle is graphically depicted via a pattern 424A.

[0063] As further shown in Figure 4, instead of target muscle 423 being activated, the hyo-glossus muscle 426 is activated, as graphically depicted via a pattern 427.

[0064] By displaying this juxtaposition of pattern 424A for target muscle 423 and pattern 427 for the actually activated muscle 426, the interface (30,212) provides an immediately recognizable differentiation between the activated muscle and the target muscle, and thereby indicates that the stimulation site did not result in activation of the target muscle and indicates which muscle (e.g. muscle 426) was activated. This differentiation informs the user that a different stimulation site should be selected.

[0065] In some embodiments, upon such differentiation, system 200 (FIG. 3) suggests graphically (e.g. via highlighting) which nerve to stimulate instead of the previously stimulated nerve or directly stimulated muscle. Moreover, in some embodiments, this suggestion is communicated via directional arrows, as provided via directional function 328 of visual identifier module 320 (Figure 3).

[0066] Figure 5 schematically illustrates substantially the same features and attributes as Figure 4, except for illustrating that the activated muscle (represented by dotted pattern 427) substantially coincides with the target muscle (represented by cross-shaped pattern 424A). Alternatively, an entirely different graphical pattern can identify the convergence of the activated muscle and the target muscle.

[0067] This arrangement provides easier and more effective indication to a user of when a target stimulation site has been located by visually identifying the corresponding innervated muscle upon its activation. Moreover, interface 212 graphically displays via highlighting which nerve was stimulated and, via a graphical marker or highlighting, the precise location along that nerve at which the stimulation was applied. Moreover, even after

the stimulation element is selectively deactivated, interface 212 allows the user to mark (or automatically marks) the stimulation site, such as via target spot marker function 292.

[0068] In some embodiments, the graphical patterns 424A, 427 are replaced with shading via shading function 324 and/or replaced with color via color function 322 of visual identifier module 320 (Figure 3).

[0069] Figure 6A is a flow diagram schematically illustrating a method 450 of identifying a target stimulation site. In one embodiment, method 450 is performed using the systems, components, and modules as previously described in association with Figures 1-5. In some embodiments, other systems and components are used to perform method 450.

[0070] As shown in Figure 6A, method 450 includes placing sensor probes onto various portions of a patient body portion in order to removably couple each sensor probe relative to a respective one of a plurality of muscles of the patient body portion. These muscles can include a target muscle innervated by a target nerve, non-target muscle(s) adjacent a target electrode site along the target nerve, and/or non-target muscles adjacent the target muscle(s), as shown at 452. Placement of sensor probes on these non-target muscles allows sensing the activation of the non-target muscles adjacent either the target nerve site or the target muscle, which is in turn, indicative of sub-optimal electrode positioning relative to a target nerve site. When graphically identified in an image-based navigation interface (or via direct observation of a functional result, such as tongue protrusion) sensing activation of the non-target muscles provides feedback to reposition the stimulation element toward the target nerve site.

[0071] Accordingly, sensor probes are placed at locations of muscles near the nerve to be stimulated and locations of muscles (or muscle groups) likely to result in muscle activation, whether or not those muscles are the target muscles. With the sensor probes arranged in this manner, upon implementing the nerve stimulation a displayed image (e.g. displayed image 40 in Figure 1) in interface 30, 212 will highlight which muscles are activated for each trial stimulation, as noted below.

[0072] In one embodiment, method 450 proceeds from the sensor probe placement at 452 to use a test tool (not an implantable electrode) for iterative insertion to stimulate potential sites while observing neuromuscular responses, as highlighted on displayed image (e.g. displayed image 40 of Figure 1) of interface 30, 212 to identify a probable target site and/or a probable percutaneous pathway to the target site. Thereafter, at 454, method 450 proceeds to advancing an implantable nerve stimulation electrode to a probable target site via image-based navigation while simultaneously provoking and observing neuromuscular responses. However, in performing method 450, one need not use the test tool (at 454) prior to advancing the implantable electrode within the pertinent body region (at 456).

[0073] In one embodiment, method 450 is performed in association with nerve-muscle index 294 (Figures 3 and 6B), among other resources of system 200. In one aspect, the nerve-muscle index 294 provides a basis to build a graphical user interface (e.g. user interface 30, 40 of Figure 1) by adding the monitored tissues to the skeletal outline. The nerve-muscle index 294 also provides an additional portion of interface 30 (as described later in association with Figure 12) or alternate view (such as Figure 6B) that supplements the visual representation of the current location of the stimulation electrode and sensor probes (in interface 30) by supplying a checklist of which muscle groups are activated or inactivated.

[0074] As illustrated by Figure 6B, in some embodiments, the nerve-muscle index 294 comprises an array 460 of primary parameters and an array 470 of corroborating parameters. The array 460 of primary parameters includes a nerve parameter 462, a muscle response parameter 464, and a probe parameter 466. The nerve parameter 462 maintains a list of nerves of the patient body portion while the muscle response parameter 464 maintains a list of muscles that respond to and that are innervated by a respective one of the listed nerves. In addition, the muscle response parameter 464 tracks an amplitude of muscle response for a particular muscle upon nerve stimulation. By comparing different stimulation sites and an amplitude of the muscle response (such as through electromyography), one can map the location of a target nerve. The probe parameter 466 maintains a listing of the various sensor probes that are removably coupled to the respective muscles with each sensor probe matched to a particular muscle. Accordingly, via index 294, each listed muscle is correlated to one of the respective sensor probes and relative to a respective one of the potential target nerves that innervate the respective listed muscle.

[0075] With further reference to Figure 6B, in one example, via index 294, a lateral branch of the hypoglossal nerve is correlated with a styloglossus muscle and sensor probe number 1. Accordingly, when a response is observed for sensor probe 1, the system automatically determines that the response is for the styloglossus muscle and an accompanying graphical identification is made on the user interface to visually highlight the activation of that muscle. In another example, via index 294, a medial branch of the hypoglossal nerve is correlated with a genioglossus muscle and sensor probe number 2. Accordingly, when a response is observed for sensor probe 2, the system automatically determines that the response is for the genioglossus muscle and an accompanying graphical identification is made on the user interface to visually highlight the activation of that muscle. As further illustrated in index 294, additional nerves or nerve branches are listed, such as the main trunk of the hypoglossal nerve, which activates several muscle groups, and which is associated with multiple sensor probes distributed across the multiple muscle groups. Accordingly, in cooperation with interface 40 (Figure 1), nerve-muscle index 294 enables an operator to differentiate which

nerve branch is being stimulated, among multiple potential target nerve branches, and therefore, responsible for activation of a particular muscle group or muscle groups.

[0076] The array 470 of corroborating parameters is used to confirm the identity of the muscle that was activated and/or whether the muscle activation resulted from nerve stimulation or direct electrical stimulation of the muscle. In some embodiments, the array 470 of corroborating parameters includes a muscle response time parameter 472, a visible muscle response parameter 474, a twitch response parameter 476, and a functional result parameter 478.

[0077] In some embodiments, the response module 352 of interface 212 (Figure 3) is configured to confirm that the target nerve identified via the nerve-muscle response index caused the activation of the respective listed muscle. In one embodiment, this confirmation is executed via a response time function and tracked as a response time parameter 472 in index 294. The response time parameter indicates the amount of time between a stimulation of a nerve and when the response is observed at the muscle (innervated by that stimulated nerve). This information is used to determine whether the activation of muscle was caused by electrical stimulation of the nerve or by direct electrical stimulation of the muscle, as the direct electrical stimulation of the muscle would result in a response time of near zero.

[0078] Moreover, the response time of a particular muscle depends on the location of stimulation along the nerve and its respective branches. Conceptually speaking, one can divide the nerve into segments with each segment innervating a particular set of muscles. By using the measured reaction time and observing which muscle group is activated, one can determine the location of stimulation, and therefore which nerve or nerve branch was stimulated. In some embodiments, in accordance with general principles of the present disclosure, this process can be performed in whole, or in part, according to the methods and systems described and illustrated in association with Testerman U.S. Patent 5,591,216.

[0079] In some embodiments, a visible muscle response parameter 474 of index 294 is configured to indicate a visible muscle response that a user could observe, such as whether the tongue retracts which corresponds to activation of tongue protrusor muscles, such as a styloglossus muscle listed in the muscle response parameter 464. If the observed behavior upon application of a stimulation signal matches the expected response for a target nerve, then one gains assurance that the target stimulation site has been identified. On the other hand, if the observed behavior upon application of a stimulation signal does not match the expected response for a target nerve, then one learns that the target stimulation site has not been identified. In the latter case, upon repositioning a stimulation element, a subsequent stimulation signal is applied to activate a muscle and the response is observed to evaluate whether the target site has been identified.

[0080] In some embodiments, the muscle motion parameter 476 of index 294 is configured to track whether motion of the muscle can be detected, whether or not that also results in a highly visible response (such retraction or protrusion of the tongue). In one embodiment, the detectable muscle motion parameter 476 tracks muscle responses via ultrasound sensor nodes that detect muscle motion via Doppler principles. For example, in response to a stimulation of a nerve, motion of the genioglossus muscle is detectable via an ultrasound sensor node placed underneath the chin while motion of the styloglossus muscle is detectable via an ultrasound sensor node placed through the jaw. In cooperation with interface 40 and ultrasound detection function 375 of response evaluator 352 of system 200 (Figure 3), a corresponding anatomical feature (e.g. nerves and muscles) is highlighted in image 30 (Figure 1), in index 294 (Figure 6B), or combined interface 700 (Figure 12) upon ultrasound detection of motion for a particular muscle or nerve.

[0081] In another embodiment, the detectable muscle motion parameter 476 tracks whether a twitch response of a muscle is observed, which can further corroborate which nerve was stimulated by which muscle responds via the twitch.

[0082] Likewise, the functional result parameter 478 of index 294 is configured to identify what functional result (e.g. an increased or a decreased percentage of airway patency) would be observed upon activation of a particular muscle, which in turn helps to determine if the target nerve or portion of the target nerve was stimulated. For example, observation and/or measurement of the cross-sectional area of upper airway patency indicates whether the desired muscle response is achieved in response to a test stimulation at a target nerve site. In another example, a functional result tracked via parameter 478 includes a relative amount and location of air pressure within the upper airway 810 as indicated via a multi-sensor catheter probe 820, as illustrated in Figure 13. As shown in Figure 13, the multi-sensor probe 820 extends within upper airway 810 and includes an array of linearly arranged pressure-sensitive sensors 822 programmed to separately indicate a measured pressure at each sensor 822. Upon an obstruction occurring in the airway 810, the air pressure adjacent one or more of the sensors 822 will be significantly impacted, and thereby indicate a location of the obstruction along the airway. In one embodiment, each separate sensor 822 comprises a mini-lumen exposed to the air within the upper airway and in communication with a pressure transducer so that the air pressure is measurable at each location of the respective sensors 822 along the length of probe 820, thereby indicating the air pressure at different locations within the airway. In some other embodiments, each sensor 822 comprises a balloon-like structure adapted to measure the air pressure at each location of the respective sensors 822.

[0083] Figure 13 also illustrates an image 830 display-

able in a user interface, such as interface 30 in Figure 1 or in functional result display 710 of interface 700 in Figure 12, as will be described further. Image 830 displays an image 840 of airway 830, with the airway 830 schematically or graphically divided into segments 842 with each segment 842 generally corresponding to a location of a sensor 822. Based on the sensed pressures, image 840 indicates a location of obstruction from a sleep-related breathing disorder (such as obstructive sleep apnea) via highlighter indicator 845. This information is also used to evaluate whether stimulation of a target stimulation site on a nerve, results in a muscle response that opens the airway 830 in the vicinity of likely obstruction, such as marked visually via identifier 845.

[0084] In one embodiment, information available from nerve-muscle response index 294 (such as the corroborating parameters of array 470) is graphically displayed at the user interface at the time that a muscle is stimulated to aid the user in identifying corroborating information, such as a twitch or functional result, by reminding the user what behavior is to be observed.

[0085] In some embodiments, in addition to the image 40 being displayed in interface 30 (Figure 1), a nerve-muscle index (including at least some of the features of nerve-muscle index 294 of Figure 6A) is displayed in interface 30 along with image 40 and/or alternately with image 40. The displayed index lists a plurality of muscle groups for which a response is expected upon application of the stimulation signal, such as the listings in index 294. In some embodiments, the displayed index includes, but is not limited to, one or more of: (1) a description of an expected muscular response resulting from application of the stimulation signal at the target site; (2) highlighting, upon activation via the stimulation signal, at least one of the respective muscle groups listed in the index; and (3) data surrounding the muscular response including at least one of a response time and a response amplitude. In addition to supplementing the information displayed via image 40 in user interface 30, the nerve-muscle index aids in displaying or identifying to the user, which muscle group has been activated in the event that the graphical display of the activated muscle in image 40 is difficult to see.

[0086] In some embodiments, the nerve-muscle index 294 also lists one or more muscle groups for which stimulation is neither desired nor expected. To the extent that these respective muscle groups do not become highlighted in the nerve-muscle index 294, the physician receives assurance that undesirable neuromuscular stimulation is avoided. However, in the event that such undesirable neuromuscular stimulation does occur and that particular muscle group was highlighted, the nerve-muscle index 294 provides one warning mechanism to alert the physician of undesirable stimulation.

[0087] In some embodiments, the nerve-muscle index 294 is storable in memory and suitable for printing to provide a record of the navigation process and/or implantation procedure of an implantable stimulation electrode.

[0088] In one embodiment, an interface 700 that includes a combination of image 40 and nerve-muscle index 294 is described and illustrated in association with Figure 12. As shown in Figure 12, interface 700 includes the interface 30 (Figure 1), the interface 400 (Figure 4), nerve-muscle index 460 (Figure 6B), each having substantially the same features and attributes as previously described and illustrated in association with those respective Figures. In addition, in some embodiments, interface 700 further includes functional result display interface 710. As shown in Figure 12, interface 710 includes an image 720 of an anatomical feature (e.g., a cross-sectional view 722 of the upper airway viewable via endoscopy) pertinent to observing a functional result of stimulating a target nerve. Interface 710 also includes a size parameter 730, a shape parameter 732, a location parameter 731, a motion parameter 734, and a source function 740. The size and shape parameters 730, 732 track a change in the size or shape of an anatomical feature. For example, an upper airway can exhibit a general reduction in its cross-sectional area while maintaining a generally circular cross-sectional shape or alternatively could exhibit a significant reduction in its cross-sectional area as the generally circular shape disappears as the airway generally collapses. The location parameter 731 tracks a general anatomical location and/or a specific location on an anatomical feature at which a functional result can be observed or measured. The motion parameter 734 indicates whether or not motion has occurred (e.g. does tongue protrude?) upon stimulation at a target site.

[0089] The source function 740 tracks a source by which an image (or other graphical display) of a functional result is obtained or produced. In some embodiments, source function 740 includes an endoscope parameter 741, an air pressure parameter 742, an ultrasound parameter 743, an EMG/CMAP parameter 744, and an impedance parameter 745. Each respective parameter 741-745 provides information consistent with the previously described sensing or detecting functions. For example, ultrasound parameter 743 provides information available via ultrasound detection function 375 (Figure 3) while air pressure parameter 743 provides information available via multi-balloon catheter probe 820 (Figure 13).

[0090] In general terms, interface 710 offers a highly integrated display to facilitate navigation of a stimulation element and/or related tools by graphically displaying anatomical features (e.g., bones, tissues, etc.) and the instruments in the field of navigation in real time as the instruments are moved along a navigation path. Moreover, at the same time, interface 710 displays, in real-time, a functional visualization of whether a given path or nerve stimulation test site produces a desired functional result of the intended or target muscle. This functional visualization is coordinated with images of the navigation field so that the navigational information and the functional result information is displayed in a single interface to aid

the operator with real-time information that facilitates more accurate navigation and placement of a stimulation element or test tool. As further illustrated in Figure 12, nerve-muscle index 460 provides additional text information in real-time to aid the operator in quickly corroborate the visual/graphic information displayed at interfaces 40, 400, or 710 in interface 700. For example, index 460 in Figure 12 includes a highlighting function 750 in which a nerve or innervated muscle of interest (such as a nerve being stimulated) is highlighted in index 460 (represented by shading) and which corresponds to a highlighted nerve and/or innervated muscle in interface 40, 400, or 710. It will be understood that, in some embodiments, interface 700 is selectable controllable to display a subset of the interfaces 40, 400, 710 and index 460, at the discretion of the operator.

[0091] Figure 7 is a side view schematically illustrating an arrangement 500 of placed sensor probes (510, 512) for detecting impedance changes indicative of a functional response of a target muscle. In particular, Figure 7 illustrates a head region 502 of patient and in particular, an oral and neck region including tongue 522 and airway patency-related muscle 520. In one aspect, probe 510 is placed on tongue 522 while probe 512 is placed under the jaw adjacent airway patency-related muscles 520. With this arrangement, impedance is measurable with the two probes 510, 512 such that changes in impedance can be used to quantify movement of the tongue. In particular, increased impedances to the electrodes near the front of the tongue and the back of the airway would be indicative of a substantial forward movement of the tongue.

[0092] In general terms, correct placement of a probe can be confirmed by applying stimulation to one probe and then sensing the response from other probes or observation by a physician. While some probes may not be placed in ideal locations, it will be understood that whatever information is obtained from these probes is used to help determine or corroborate an intended navigation path and/or identification of a target stimulation site.

[0093] Figures 8-9 further illustrate a sensor probe 540 (that can act as probe 510 and/or 512). Figure 8 is a top view of probe 540, which includes a generally elongate flexible body 542 made of (or coated with) a biocompatible material. In some embodiments, probe 540 includes a plurality of anchors 544 that extend from opposite side portions 543 of body 542 and are spaced apart from each other longitudinally along a length of body 542. Each anchor 544 includes a pressure sensitive adhesive on its underside to allow secure direct adhesion relative to a surface body portion. However, in other embodiments, it will be understood that any number of different types of fasteners can be used with, or in place of, anchors 544. In some embodiments, additional fastening is achieved via coating the underside 547 of body 542 with adhesive.

[0094] Figures 8-9 further illustrate that probe 540 includes an array of fine gauge needles 546 arranged in a spaced apart relationship along the length of body 542,

and extend away from an underside 547 of body 542. In one embodiment, the needles 546 extend generally perpendicular to the underside 547 of body 542 although in other embodiments, needles 546 can extend at different angles relative to underside 547.

[0095] While Figure 7 provides one schematic illustration of probe placement, it will be understood that can have more than two sensors and that sensors can be placed in other locations. For example, as described previously, in some embodiments, some sensor probes are placed at the target muscle groups (i.e. the intended muscle to be activated) to confirm that the desired muscle group is activated and to obtain data re stimulation, muscle response, etc. In addition, in some embodiments, sensor probes are placed at muscle groups near the target nerve to facilitate optimal electrode placement. In particular, an observed activation of a muscle near the target nerve site would indicate a likely sub-optimal electrode placement at the target nerve, and thereby trigger the physician to adjust the placement of the electrode.

[0096] Figure 10 is a schematic illustration of an image-based visualization system 600 used in a method of identifying a target stimulation location, according to the present disclosure. As shown in Figure 10, system 600 includes an image module 602 and graphical user interface 604. In one embodiment, system 600 corresponds to a StealthStation™ type imaging system available from Medtronic of Fridley, Minnesota. As shown in Figure 10, image module 602 provides internal images of anatomical structures of a body region, with image module 602 including a front view 610, a side view 612, a top view 614, and a plan exterior view 616. The front view 610 and top view 614 illustrate a body region including a tongue portion 620 and a nerve 622. Marker 630 provides an independent reference point, such as a fiducial marker. The side view further illustrates the tongue portion 620 and nerve 622, although showing the various muscle groups (see also Figures 4-5) supporting tongue portion 620. Using the various views of the image module 602 to collectively provide a three-dimensional view, an operator can guide a stimulation element 635 relative to the displayed nerve 622 and tongue portion 620. Moreover, the different views of the image module 602 can be displayed in combination with a graphical display (such as image 40 in Figure 1 or the images of Figures 4-5) of activated muscles and stimulated nerves (as described in association with at least Figures 1-9), thereby providing real-time indication of the precise location of a stimulation element and the effectiveness of the stimulation element at that location for activating a target muscle.

[0097] In one embodiment, system 600 includes an array of reflectors placeable on various anatomical landmarks of a patient, on tools (and different portions of the tools), and the general environment in which the patient is situated. For example, as illustrated in Figure 14, placement of a marker 880 on a patient's chin provides one such anatomical landmark relevant for visualizing procedures in the area of the head and neck. Using an infrared

camera, the various locations of the reflectors are identified and compiled into a three-dimensional map suitable to guide navigation during an invasive procedure. In some embodiments, system 600 further includes a laser projection function 900, as illustrated in Figure 14. Laser projection function 900 projects different colors 902, 904, 906 directly onto a patient 910 (based on the infrared anatomical reference points, other interpolated anatomical positions, and internal imaging information, as described in association with Figure 10) to guide navigation of instruments (e.g. a test tool, a stimulation element, a percutaneous access system, a transvenous access system) on an intended navigational path 912, 914, actual navigational path, and/or intended target 916 of the navigational path, as generally illustrated in Figure 14. In one embodiment, the differently color illumination provided via the laser projection function 900 is displayed in combination with a graphical display (such as image 40 in Figure 1 or the images of Figures 4-5) of activated muscles and stimulated nerves (as described in association with at least Figures 1-9), thereby providing real-time indication of the precise location of a stimulation element and the effectiveness of the stimulation element at that location for activating a target muscle.

[0098] In some embodiment, the color laser projection functionality greatly facilitates a transvenous access method of delivery of the stimulation element and/or test tool, as this method does not include accessing the target nerve via surface cutting of tissue.

[0099] Figure 11 is a schematic illustration of an image-based visualization system 600 used in a method of identifying a target stimulation location and/or navigation pathway, according to the present disclosure. As shown in Figure 11, system 650 includes interface 652. In one embodiment, system 650 corresponds to a LocaLisa™ type non-fluoroscopic imaging system available from Medtronic of Fridley, Minnesota. Interface 652 provides a three-dimensional representation of a body region along three orthogonal axes 653 (x), 654 (y), 655 (z). Using surface mounted electrodes arranged about an outer surface of the body portion, movement of a stimulation element 635 (test tool or implantable electrode) is tracked along a path 660 of internal locations 662 using voltage and impedance information from the surface-mounted electrodes and the conductive stimulation element 635. A reference marker 665 provides an independent reference point. With this arrangement, an image is generated of a navigated path and the location of the stimulation element 635. In some embodiments, this generated image is superimposed with patient anatomy maps and more comprehensive images (e.g. MRI), as previously described in association with Figures 1-9, for display in interface 30, 212 (Figures 1, 3) to assist in placing the stimulation element.

[0100] It will be understood that different combinations of the components of the image and navigation interfaces can be made, such as but not limited to, including or excluding a nerve-muscle index 294 (Figure 6B and 12)

with an image interface 40, including or excluding a functional result module, or even including or excluding a sensor probe interface.

[0101] Embodiments of the present invention provide for dynamic real-time identification of muscle activation from electrical stimulation in context with image-based navigation tools, thereby enhancing placement of stimulation element within a body portion of a patient.

[0102] While at least one exemplary embodiment has been presented in the foregoing detailed description, it should be appreciated that variations exist. It should also be appreciated that the exemplary embodiment or exemplary embodiments are only examples, and are not intended to limit the scope, applicability, or configuration of the present invention in any way. Rather, the foregoing detailed description will provide those skilled in the art with a convenient road map for implementing the exemplary embodiment or exemplary embodiments. It should be understood that various changes can be made in the function and arrangement of elements without departing from the scope of the invention as set forth in the appended claims.

Claims

1. A system for placing a stimulation element, the system comprising:

a neuromuscular response evaluation module including:

a stimulation engine configured to apply an electrical stimulation signal to at least one nerve within a patient head-and-neck portion via a stimulation element;

a muscle response module configured to determine a response of at least one muscle in the patient head-and-neck portion to application of the stimulation signal; and

a graphical user interface including:

an image display module configured to display at least one internal image of the patient head-and-neck portion;

a muscle identifier module configured to visually identify, superimposed on the at least one internal image, which muscles in the head-and-neck portion were activated for each respective application of the stimulation signal to at least one nerve in the head-and-neck portion.

2. The system of claim 1, wherein the neuromuscular response evaluation module comprises:

a probe interface including an array of ports; and

an array of sensor probes, each removably connectable to a respective one of the ports and removably attachable to a body portion corresponding to the muscle, the sensor probe configured to sense the muscle response.

3. The system of claim 2, wherein the user interface comprises:

a sensor designator configured to track and display a location of each sensor probe on the at least one internal image from the image display module, wherein the sensor designator is configured to determine the location based on at least one of:

a visualization of the sensor probe captured during creation of the at least one internal image; and

a user-positionable, graphic marker superimposed on the at least one internal image, wherein a location of the graphic marker represents the location of a respective one of the sensor probes.

4. The system of claim 2, wherein the sensor probes include at least one of:

an ultrasound detection node;

a multi-balloon catheter configured to sense, via air pressure, a location of obstruction within an airway; and

an endoscopic imaging device configured to detect at least one of a change in a cross-sectional area of an airway or a change in a cross-sectional shape of the airway.

5. The system of claim 1, wherein the user interface comprises:

a target nerve identifier configured to graphically identify on the at least one internal image a location of a target nerve; and

a target muscle identifier configured to graphically identify on the at least one internal image a location of a target muscle innervated by the target nerve, wherein the respective target nerve identifier and target muscle identifier are configured to be user-positionable on the at least one internal image.

6. The system of claim 1, wherein the user interface comprises:

an actually-stimulated nerve identifier configured to graphically identify on the at least one internal image a location of the nerve actually

- stimulated via the applied stimulation signal; and an actually-activated muscle identifier configured to graphically identify on the at least one internal image a location of the muscle actually activated via the actually stimulated nerve, wherein the respective locations of the actually-stimulated nerve identifier and actually-stimulated muscle identifier are automatically highlighted on the at least one internal image upon sensing the muscle activation via a sensor probe coupled to the muscle.
7. The system of claim 2, wherein the graphical user interface comprises:
- a nerve-muscle index including a first listing of potential target nerves and a second listing of muscles, wherein each listed muscle is correlated to one of the respective sensor probes and relative to a respective one of the potential target nerves that innervate the respective listed muscle.
8. The system of claim 7, wherein the muscle identifier module accesses the nerve-muscle index, so that upon detecting activation of one of the listed muscles, the corresponding target nerve responsible for activation of the muscle is highlighted within the index.
9. The system of claim 8, wherein the muscle identifier module is configured to confirm that the target nerve identified via the index caused the activation of the respective listed muscle via a response time function of the neuromuscular response evaluation module, wherein the response time function indicates whether the activation of muscle was caused by electrical stimulation of the nerve or by direct electrical stimulation of the muscle.
10. The system of claim 7, wherein the nerve-muscle index comprises an array of corroborating parameters including at least one of a muscle response time parameter, a visible muscle action parameter, a muscle motion parameter, or a functional result parameter.
11. The system of claim 10, wherein the functional result parameter is configured to identify a functional result of stimulation of the target nerve, wherein the identified functional result includes at least one of a retraction of a muscle, a protrusion of a muscle, an increase in a cross-sectional area of an airway, a decrease in the cross-sectional area of the airway, an at least partial collapse in the airway, or a maintained cross-sectional shape of the airway, wherein the graphical user interface includes a display of one or more of the identified functional results.
12. The system of claim 1, wherein the muscle identifier module includes:
- a visual response intensity function configured to visually indicate a degree of response by the activated muscles via at least one of:
- a color parameter configured to display a different color for each different degree of response of the activated muscles;
- a shading parameter configured to display a different degree of shading for each different degree of response of the activated muscles; or
- a numerical parameter configured to display a different numerical identifier for each different degree of response of the activated muscles.
13. The system of claim 1, wherein the muscle identifier module includes:
- a target muscle parameter configured to track, via a plurality of sensor probes configured to be removably coupled relative to a corresponding plurality of potentially activatable muscles, each potentially activatable muscle that is innervated by one of the potential target nerve stimulation sites;
- a non-target nerve parameter configured to track, via a plurality of sensor probes configured to be removably coupled relative to a corresponding plurality of potentially activatable muscles, each potentially activatable muscle that is physically adjacent one of the potential target nerve stimulation sites and that is not targeted to be activated; and
- a non-target muscle parameter configured to track, via a plurality of sensor probes configured to be removably coupled relative to a corresponding plurality of non-target muscles, each non-target muscle that is physically adjacent one of the target muscles.
14. The system of claim 1, wherein the graphical user interface includes at least one of:
- a path marker function configured to graphically identify an intended pathway of percutaneous access based on at least one of a probable target nerve stimulation site or a confirmed target nerve stimulation site;
- a target spot marker configured to graphically identify an intended target nerve stimulation site; and
- a directional function configured to visual indicate, via a graphical representation, a suggested direction in which to move the stimulation el-

ement toward a target nerve stimulation site.

15. The system of claim 1, wherein the image display module comprises:

an image source module configured to access the at least one internal image as at least one of a magnetic resonance imaging (MRI) image, a computer tomography (CT) image, a fluoroscopy image, and a non-radiographic image, wherein the image source module is configured to obtain the at least one internal image via at least one of electronic medical records or a live image; and wherein the image display module includes at least one of:

an anatomy map module configured to graphically superimpose a generalized patient anatomy map onto the displayed at least one internal image;

and

a fiducials marker module configured to locate fiducial markers onto the at least one internal image.

Patentansprüche

1. System (20) zum Anordnen eines Stimulationselements (90), das System aufweisend:

- ein Neuromuskuläre-Reaktion-Auswertungsmodul (52, 214), enthaltend:

* einen Stimulationsmotor (70), der zum Zuführen eines elektrischen Stimulationssignals zu mindestens einem Nerv (46) innerhalb eines Kopf/Hals-Abschnitts eines Patienten (22) über ein Stimulationselement (90) konfiguriert ist;

* ein Muskelreaktionsmodul (352), das zum Bestimmen einer Reaktion von mindestens einem Muskel im Kopf/Hals-Abschnitt des Patienten (22) auf Zuführung des Stimulationssignals konfiguriert ist; und

- eine grafische Benutzeroberfläche (30), enthaltend:

* ein Bildanzeigemodul, das zum Anzeigen von mindestens einem inneren Bild (40) des Kopf/Hals-Abschnitts des Patienten (22) konfiguriert ist;

* ein Muskelidentifiziermodul, das zum visuellen Identifizieren, überlagert über das mindestens eine innere Bild (40), konfiguriert ist, welche Muskeln im Kopf/Hals-Ab-

schnitt für jede jeweilige Zuführung des Stimulationssignals zu mindestens einem Nerv (46) im Kopf/Hals-Abschnitt aktiviert wurden.

2. System nach Anspruch 1, wobei das Neuromuskuläre-Reaktion-Auswertungsmodul (52, 214) Folgendes aufweist:

eine Sondenschnittstelle mit einer Gruppierung von Anschlüssen; und

eine Gruppierung von Sensorsonden, die jede abnehmbar mit einem jeweiligen der Anschlüsse verbindbar ist und abnehmbar an einem Körperabschnitt anbringbar ist, der dem Muskel entspricht, wobei die Sensorsonde zum Abfühlen der Muskelreaktion konfiguriert ist.

3. System nach Anspruch 2, wobei die Benutzeroberfläche (30) Folgendes aufweist:

einen Sensorbezeichner, der zum Verfolgen und Anzeigen einer Stelle von jeder Sensorsonde auf dem mindestens einen inneren Bild vom Bildanzeigemodul konfiguriert ist, wobei der Sensorbezeichner zum Bestimmen der Stelle auf Grundlage von mindestens einem des Folgenden konfiguriert ist:

einer Visualisierung der Sensorsonde, die während der Erzeugung des mindestens einen inneren Bilds erfasst wird; und einer vom Benutzer positionierbaren, grafischen Markierung, die auf dem mindestens einen inneren Bild überlagert ist, wobei eine Stelle der grafischen Markierung die Stelle einer jeweiligen der Sensorsonden darstellt.

4. System nach Anspruch 2, wobei die Sensorsonde mindestens eines von Folgendem enthält:

einen Ultraschallerkennungsknoten; einen Multiballonkatheter, der zum Abfühlen, über Luftdruck, einer Blockierungsstelle innerhalb eines Atemwegs konfiguriert ist; und ein endoskopisches Bildgebungsgerät, das zum Erkennen von mindestens einem von einer Änderung in einem Querschnittsbereich eines Luftwegs oder einer Änderung einer Querschnittsform des Atemwegs konfiguriert ist.

5. System nach Anspruch 1, wobei die Benutzeroberfläche (30) Folgendes aufweist:

einen Zielnervidentifizierer, der zum grafischen Identifizieren einer Stelle eines Zielnervs auf dem mindestens einen inneren Bild konfiguriert

- ist; und
einen Zielmuskelidentifizierer, der zum grafischen Identifizieren einer Stelle eines Zielmuskels, der durch den Zielnerv angeregt wird, auf dem mindestens einen inneren Bild konfiguriert ist, 5
wobei der jeweilige Zielnervidentifizierer und Zielmuskelidentifizierer dazu konfiguriert sind, vom Benutzer auf dem mindestens einen inneren Bild positionierbar zu sein. 10
6. System nach Anspruch 1, wobei die Benutzeroberfläche (30) Folgendes aufweist:
- einen Identifizierer eines tatsächlich stimulierten Nervs, der zum grafischen Identifizieren einer Stelle des Nervs, der tatsächlich über das zugeführte Stimulationssignal stimuliert wurde, auf dem mindestens einen inneren Bild konfiguriert ist; und 15
einen Identifizierer eines tatsächlich aktivierten Muskels, der zum grafischen Identifizieren einer Stelle des Muskels, der tatsächlich über den tatsächlich stimulierten Nerv aktiviert wurde, auf dem mindestens einen inneren Bild konfiguriert ist, 20
wobei die jeweiligen Stellen des Identifizierers des tatsächlich stimulierten Nervs und des Identifizierers des tatsächlich stimulierten Muskels nach dem Abfühlen der Muskelaktivierung über eine Sensorsonde, die an den Muskel gekoppelt ist, automatisch auf dem mindestens einen inneren Bild hervorgehoben werden. 25 30
7. System nach Anspruch 2, wobei die grafische Benutzeroberfläche (30) Folgendes aufweist: 35
einen Nerv/Muskel-Index, der eine erste Auflistung von potentiellen Zielnerven und eine zweite Auflistung von Muskeln enthält, wobei jeder aufgelistete Muskel mit einer der jeweiligen Sensorsonden und bezüglich eines jeweiligen der potentiellen Zielnerven, die den jeweiligen aufgelisteten Muskel anregen, korreliert ist. 40 45
8. System nach Anspruch 7, wobei das Muskelidentifizierermodul auf den Nerv/Muskel-Index zugreift, sodass nach der Erkennung einer Aktivierung von einem der aufgelisteten Muskeln der entsprechende Zielnerv, der für die Aktivierung des Muskels verantwortlich ist, innerhalb des Index hervorgehoben wird. 50
9. System nach Anspruch 8, wobei das Muskelidentifizierermodul zum Bestätigen, dass der Zielnerv, der über den Index identifiziert ist, die Aktivierung des jeweiligen aufgelisteten Muskels bewirkt hat, über eine Reaktionszeitfunktion des Neuromuskuläre-Reaktion-Auswertungsmoduls konfiguriert ist, wobei 55
- die Reaktionszeitfunktion anzeigt, ob die Aktivierung des Muskels durch elektrische Stimulation des Nervs oder durch direkte elektrische Stimulation des Muskels bewirkt wurde.
10. System nach Anspruch 7, wobei der Nerv/Muskel-Index eine Gruppierung von bekräftigenden Parametern aufweist, darunter mindestens ein Muskelreaktionszeitparameter, ein Sichtbare-Muskeltätigkeit-Parameter, ein Muskelbewegungsparameter oder ein Funktionsergebnisparameter.
11. System nach Anspruch 10, wobei der Muskelbewegungsparameter zum Identifizieren konfiguriert ist, ob Muskelbewegung über Ultraschallerkennung erkannt wird, und wobei der Funktionsergebnisparameter zum Identifizieren eines Funktionsergebnisses einer Stimulation des Zielnervs konfiguriert ist, wobei das identifizierte Funktionsergebnis mindestens eines von einer Retraktion eines Muskels, einer Protrusion eines Muskels, einer Zunahme in einem Querschnittsbereich eines Atemwegs, eine Abnahme im Querschnittsbereich des Atemwegs, einen mindestens teilweisen Kollaps im Atemweg oder eine beibehaltene Querschnittsform des Atemwegs enthält, 30
wobei die grafische Benutzeroberfläche eine Anzeige von einem oder mehreren der identifizierten Funktionsergebnisse enthält.
12. System nach Anspruch 1, wobei das Muskelidentifizierermodul Folgendes enthält:
- eine visuelle Reaktionsintensitätsfunktion, die zum visuellen Anzeigen eines Reaktionsgrads durch den aktivierten Muskel über mindestens eines von Folgendem konfiguriert ist:
- einen Farbparameter, der zum Anzeigen einer unterschiedlichen Farbe für jeden unterschiedlichen Reaktionsgrad der aktivierten Muskeln konfiguriert ist; 35
einen Schattierungsparameter, der zum Anzeigen eines unterschiedlichen Schattierungsgrads für jeden unterschiedlichen Reaktionsgrad der aktivierten Muskeln konfiguriert ist; oder
einen numerischen Parameter, der zum Anzeigen eines unterschiedlichen numerischen Identifizierers für jeden unterschiedlichen Reaktionsgrad der aktivierten Muskeln konfiguriert ist. 40 45 50
13. System nach Anspruch 1, wobei das Muskelidentifizierermodul Folgendes enthält:
- einen Zielmuskelparameter, der zum Verfolgen, über mehrere Sensorsonden, die dazu konfigu-

riert sind, bezüglich mehreren entsprechenden, potentiell aktivierbaren Muskeln abnehmbar angekoppelt zu sein, jedes potentiell aktivierbaren Muskels konfiguriert ist, welcher durch eine der potentiellen Zielnervstimulationsstellen ange- 5 regt wird;

einen Nichtzielnervparameter, der zum Verfolgen, über mehrere Sensorsonden, die dazu konfiguriert sind, bezüglich mehreren entsprechenden, potentiell aktivierbaren Muskeln ab- 10 nehmbar angekoppelt zu sein, jedes potentiell aktivierbaren Muskels konfiguriert ist, welcher physisch einer der potentiellen Zielnervstimulationsstellen benachbart ist und nicht zur Aktivierung zum Ziel genommen ist; und

einen Nichtzielmuskelparameter, der zum Verfolgen, über mehrere Sensorsonden, die dazu konfiguriert sind, bezüglich mehreren entsprechenden Nichtzielmuskeln abnehmbar ange- 20 koppelt zu sein, jedes Nichtzielmuskels konfiguriert ist, welcher physisch einem der Zielmuskeln benachbart ist.

14. System nach Anspruch 1, wobei die grafische Benutzeroberfläche (30) mindestens eines von Folgendem enthält: 25

eine Pfadmarkierungsfunktion, die zum grafischen Identifizieren eines beabsichtigten Pfads eines perkutanen Zugangs auf Grundlage von mindestens einer einer wahrscheinlichen Zielnervstimulationsstelle oder einer bestätigten Zielnervstimulationsstelle konfiguriert ist; 30

eine Zielpunktmarkierung, die zum grafischen Identifizieren einer beabsichtigten Zielnervstimulationsstelle konfiguriert ist; und 35

eine Richtungsfunktion, die zum visuellen Anzeigen, über eine grafische Darstellung, einer vorgeschlagenen Richtung konfiguriert ist, in der das Stimulationselement zu einer Zielnervstimulationsstelle hin zu bewegen ist. 40

15. System nach Anspruch 1, wobei das Bildanzeigemodul Folgendes aufweist:

ein Bildquellenmodul, das zum Zugreifen auf das mindestens eine innere Bild als mindestens eines von einem Magnetresonanzbildgebungs- (MRI-) Bild, einem Computertomografie- (CT-) Bild, einem Fluoroskopiebild und einem nicht radiografischen Bild konfiguriert ist, wobei das Bildquellenmodul zum Erhalten des mindestens einen inneren Bilds über mindestens eines von elektronischen medizinischen Aufzeichnungen oder einem Live-Bild konfiguriert ist; und 50

wobei das Bildanzeigemodul mindestens eines von Folgendem enthält: 55

ein Anatomiekartenmodul, das zum grafischen Überlagern einer generalisierten Patientenanatomiekarte auf das angezeigte, mindestens eine innere Bild konfiguriert ist; und

ein Messmarkierungsmodul, das zum Lokalisieren von Messmarkierungen auf dem mindestens einen inneren Bild konfiguriert ist.

Revendications

1. Système (20) permettant de placer un élément de stimulation (90), le système comprenant :

- un module d'évaluation de réponse neuromusculaire (52, 214) comprenant :

* un moteur de stimulation (70) conçu pour appliquer un signal de stimulation électrique à au moins un nerf (46) dans une partie tête et cou d'un patient (22) par l'intermédiaire d'un élément de stimulation (90) ;

* un module de réponse musculaire (352) conçu pour déterminer une réponse d'au moins un muscle dans la partie tête et cou du patient (22) à l'application du signal de stimulation ; et

- une interface graphique utilisateur (30) comprenant :

* un module d'affichage d'images conçu pour afficher au moins une image interne (40) de la partie tête et cou du patient (22) ;

* un module d'identification de muscles conçu pour identifier visuellement, par superposition sur l'au moins une image interne (40), les muscles de la partie tête et cou qui ont été activés pour chaque application respective du signal de stimulation à au moins un nerf (46) dans la partie tête et cou.

2. Système selon la revendication 1, dans lequel le module d'évaluation de réponse neuromusculaire (52, 214) comprend :

une interface de sonde comprenant un réseau de ports ; et

un réseau de sondes de détection, chacune pouvant être connectée amovible à un port respectif des ports et pouvant être fixée amovible à une partie corps correspondant au muscle, la sonde de détection étant conçue pour détecter la réponse musculaire.

3. Système selon la revendication 2, dans lequel l'in-

terface utilisateur (30) comprend :

un élément de désignation de capteur conçu pour suivre et afficher un emplacement de chaque sonde de détection sur l'au moins une image interne à partir du module d'affichage d'images, dans lequel l'élément de désignation de capteur est conçu pour déterminer l'emplacement sur la base d'au moins un parmi :

une visualisation de la sonde de détection capturée pendant la création de l'au moins une image interne ; et

un repère graphique pouvant être positionné par l'utilisateur, superposé sur l'au moins une image interne, dans lequel un emplacement du repère graphique représente l'emplacement d'une sonde respective des sondes de détection.

4. Système selon la revendication 2, dans lequel les sondes de détection comprennent au moins un parmi :

un noeud de détection à ultrasons ;

un cathéter à ballonnets multiples conçu pour détecter, par l'intermédiaire de la pression de l'air, un emplacement d'obstruction dans des voies aériennes ; et

un dispositif d'imagerie endoscopique conçu pour détecter un changement de région de coupe transversale des voies aériennes et/ou un changement d'une forme de coupe transversale des voies aériennes.

5. Système selon la revendication 1, dans lequel l'interface utilisateur (30) comprend :

un élément d'identification de nerf cible conçu pour identifier graphiquement sur l'au moins une image interne un emplacement d'un nerf cible ; et

un élément d'identification de muscle cible conçu pour identifier graphiquement sur l'au moins une image interne un emplacement d'un muscle cible innervé par le nerf cible, dans lequel l'élément d'identification de nerf cible respectif et l'élément d'identification de muscle cible respectif sont conçus pour pouvoir être positionnés par l'utilisateur sur l'au moins une image interne.

6. Système selon la revendication 1, dans lequel l'interface utilisateur (30) comprend :

un élément d'identification de nerf en cours de stimulation conçu pour identifier graphiquement sur l'au moins une image interne un emplace-

ment du nerf en cours de stimulation par l'intermédiaire du signal de stimulation appliqué ; et un élément d'identification de muscle en cours d'activation conçu pour identifier graphiquement sur l'au moins une image interne un emplacement du muscle en cours d'activation par l'intermédiaire du nerf en cours de stimulation, dans lequel les emplacements respectifs de l'élément d'identification de nerf en cours de stimulation et de l'élément d'identification de muscle en cours de stimulation sont automatiquement mis en évidence sur l'au moins une image interne lors de la détection de l'activation du muscle par l'intermédiaire d'une sonde de détection couplée au muscle.

7. Système selon la revendication 2, dans lequel l'interface utilisateur graphique (30) comprend :

un index nerf-muscle comprenant une première liste de nerfs cibles potentiels et une seconde liste de muscles, chaque muscle répertorié étant corrélé avec l'une des sondes de détection respectives et par rapport à un nerf respectif des nerfs cibles potentiels qui innervent le muscle répertorié respectif.

8. Système selon la revendication 7, dans lequel le module d'identification de muscles accède à l'index nerf-muscle, de sorte que lors de la détection de l'activation de l'un des muscles répertoriés, le nerf cible correspondant responsable de l'activation du muscle soit mis en évidence dans l'index.

9. Système selon la revendication 8, dans lequel le module d'identification de muscles est conçu pour confirmer que le nerf cible identifié par l'intermédiaire de l'index a provoqué l'activation du muscle répertorié respectif par l'intermédiaire d'une fonction de temps de réponse du module d'évaluation de réponse neuromusculaire, la fonction de temps de réponse indiquant si l'activation du muscle a été provoquée par une stimulation électrique du nerf ou par une stimulation électrique directe du muscle.

10. Système selon la revendication 7, dans lequel l'index nerf-muscle comprend un réseau de paramètres de confirmation comprenant un paramètre de temps de réponse musculaire et/ou un paramètre d'action musculaire visible et/ou un paramètre de mouvement musculaire et/ou un paramètre de résultat fonctionnel.

11. Système selon la revendication 10, dans lequel le paramètre de mouvement musculaire est conçu pour identifier si le mouvement musculaire est détecté par l'intermédiaire d'une détection à ultrasons, et dans lequel le paramètre de résultat fonctionnel

est conçu pour identifier un résultat fonctionnel d'une stimulation du nerf cible, dans lequel le résultat fonctionnel identifié comprend une rétraction d'un muscle et/ou une saillie d'un muscle et/ou une augmentation de la région de coupe transversale des voies aériennes et/ou une diminution de la région de coupe transversale des voies aériennes et/ou un affaissement au moins partiel des voies aériennes et/ou une forme de coupe transversale maintenue des voies aériennes,

dans lequel l'interface graphique utilisateur comprend un affichage d'un ou de plusieurs résultats fonctionnels identifiés.

12. Système selon la revendication 1, dans lequel le module d'identification de muscles comprend :

une fonction d'intensité de réponse visuelle conçue pour indiquer visuellement un degré de réponse par les muscles activés par l'intermédiaire d'au moins un parmi :

un paramètre de couleur conçu pour afficher une couleur différente pour chaque degré différent de réponse des muscles activés ;

un paramètre d'ombrage conçu pour afficher un degré différent d'ombrage pour chaque degré différent de réponse des muscles activés ; ou

un paramètre numérique conçu pour afficher un identifiant numérique différent pour chaque degré différent de réponse des muscles activés.

13. Système selon la revendication 1, dans lequel le module d'identification de muscles comprend :

un paramètre de muscle cible conçu pour suivre, par l'intermédiaire d'une pluralité de sondes de détection conçues pour être couplées amovibles à une pluralité correspondante de muscles potentiellement activables, chaque muscle potentiellement activable qui est innervé par l'un des sites de stimulation nerveuse cible potentiels ;

un paramètre de nerf non cible conçu pour suivre, par l'intermédiaire d'une pluralité de sondes de détection conçues pour être couplées amovibles à une pluralité correspondante de muscles potentiellement activables, chaque muscle potentiellement activable qui est physiquement adjacent à l'un des sites de stimulation nerveuse cible potentiels et qui n'est pas ciblé pour être activé ; et

un paramètre de muscle non cible conçu pour suivre, par l'intermédiaire d'une pluralité de sondes de détection conçues pour être couplées

amovibles à une pluralité correspondante de muscles non cibles, chaque muscle non cible qui est physiquement adjacent à l'un des muscles cibles.

14. Système selon la revendication 1, dans lequel l'interface utilisateur graphique (30) comprend au moins un parmi :

une fonction de repère de trajet conçue pour identifier graphiquement un trajet prévu d'un accès percutané sur la base d'un site de stimulation nerveuse cible probable et/ou d'un site de stimulation nerveuse cible confirmé ;

un repère de point cible conçu pour identifier graphiquement un site de stimulation nerveuse cible prévu ; et

une fonction directionnelle conçue pour indiquer visuellement, par l'intermédiaire d'une représentation graphique, une direction suggérée dans laquelle déplacer l'élément de stimulation vers un site de stimulation nerveuse cible.

15. Système selon la revendication 1, dans lequel le module d'affichage d'images comprend :

un module de source d'images conçu pour accéder à l'au moins une image interne sous la forme d'une image d'imagerie par résonance magnétique (IRM) et/ou d'une image de tomodensitométrie (TDM) et/ou d'une image de fluoroscopie et/ou d'une image non radiographique, dans lequel le module de source d'images est conçu pour obtenir l'au moins une image interne par l'intermédiaire de dossiers médicaux électroniques et/ou d'une image en direct ; et dans lequel le module d'affichage d'images comprend au moins un parmi :

un module de carte anatomique conçu pour superposer graphiquement une carte anatomique de patient généralisée sur l'au moins une image interne affichée ; et un module de repère d'alignement conçu pour localiser des repères d'alignement sur l'au moins une image interne.

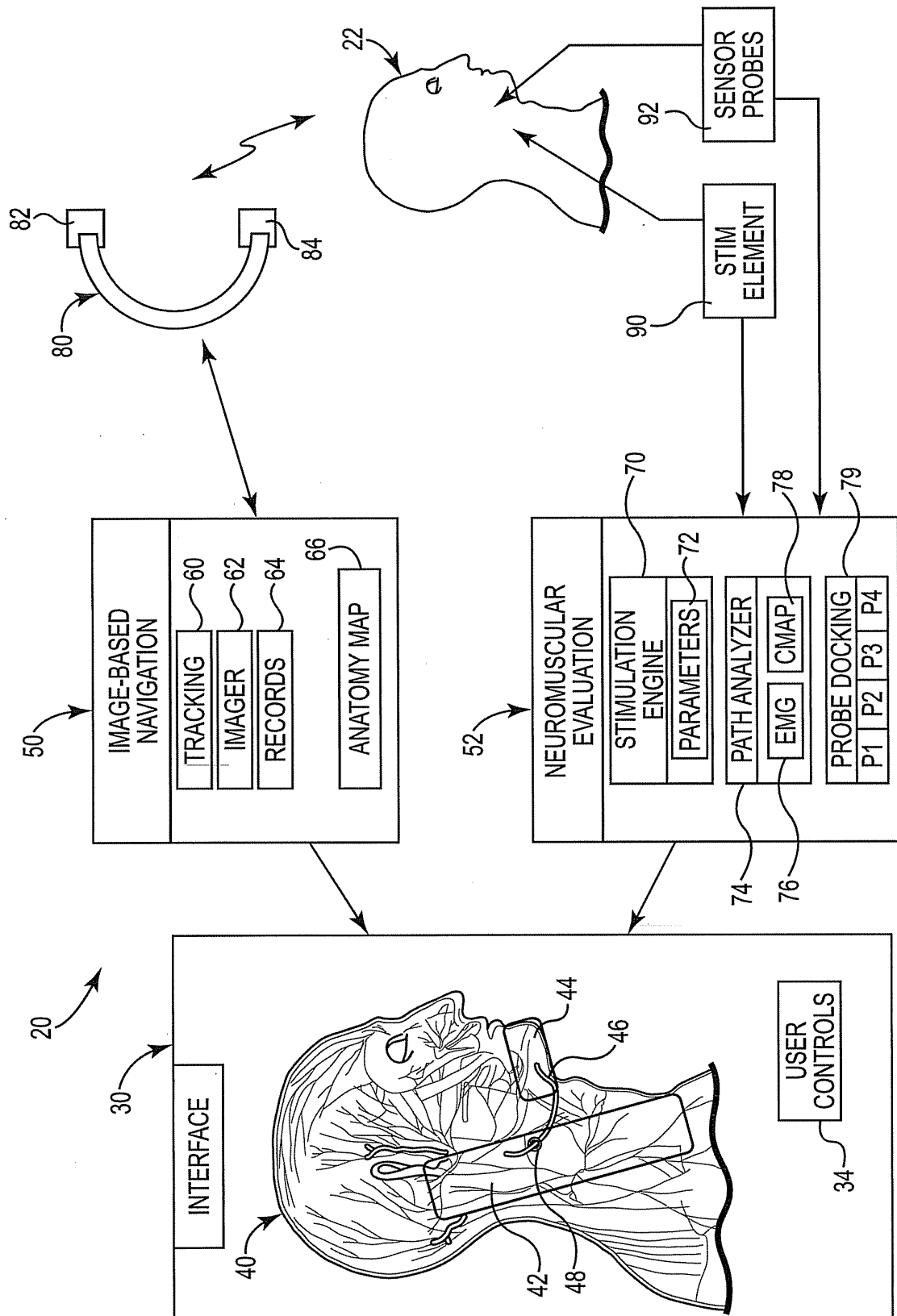
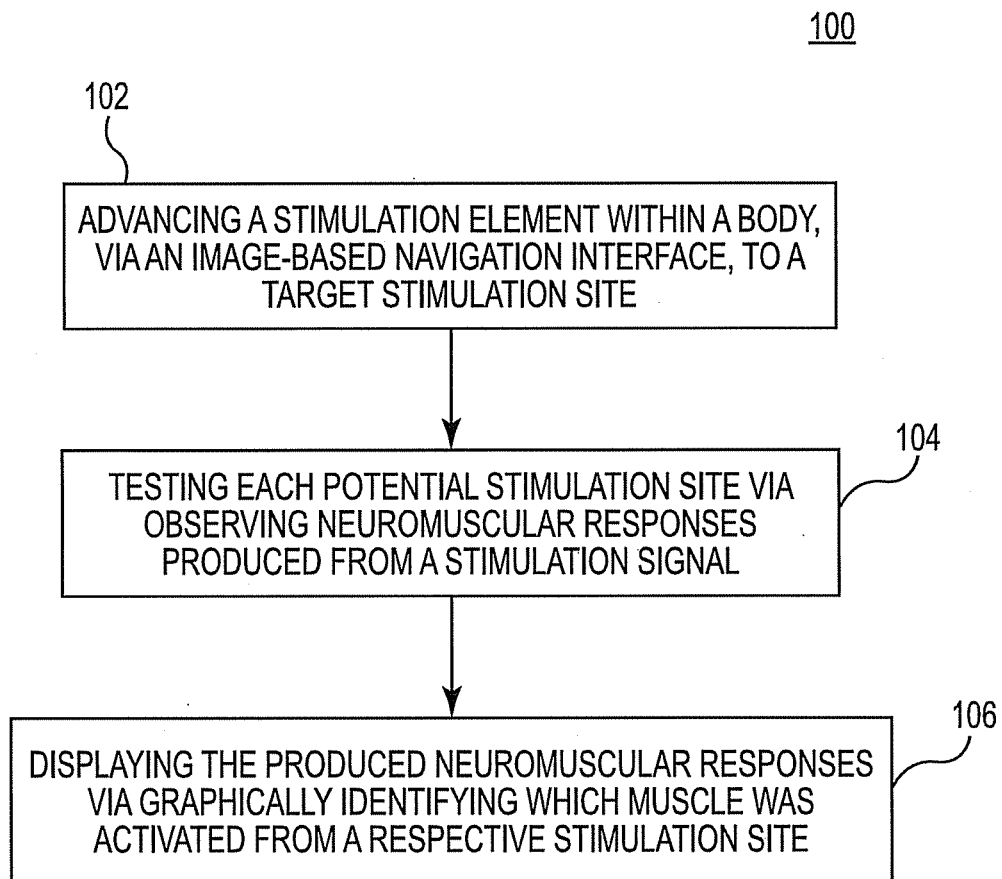


Fig. 1

**Fig. 2**

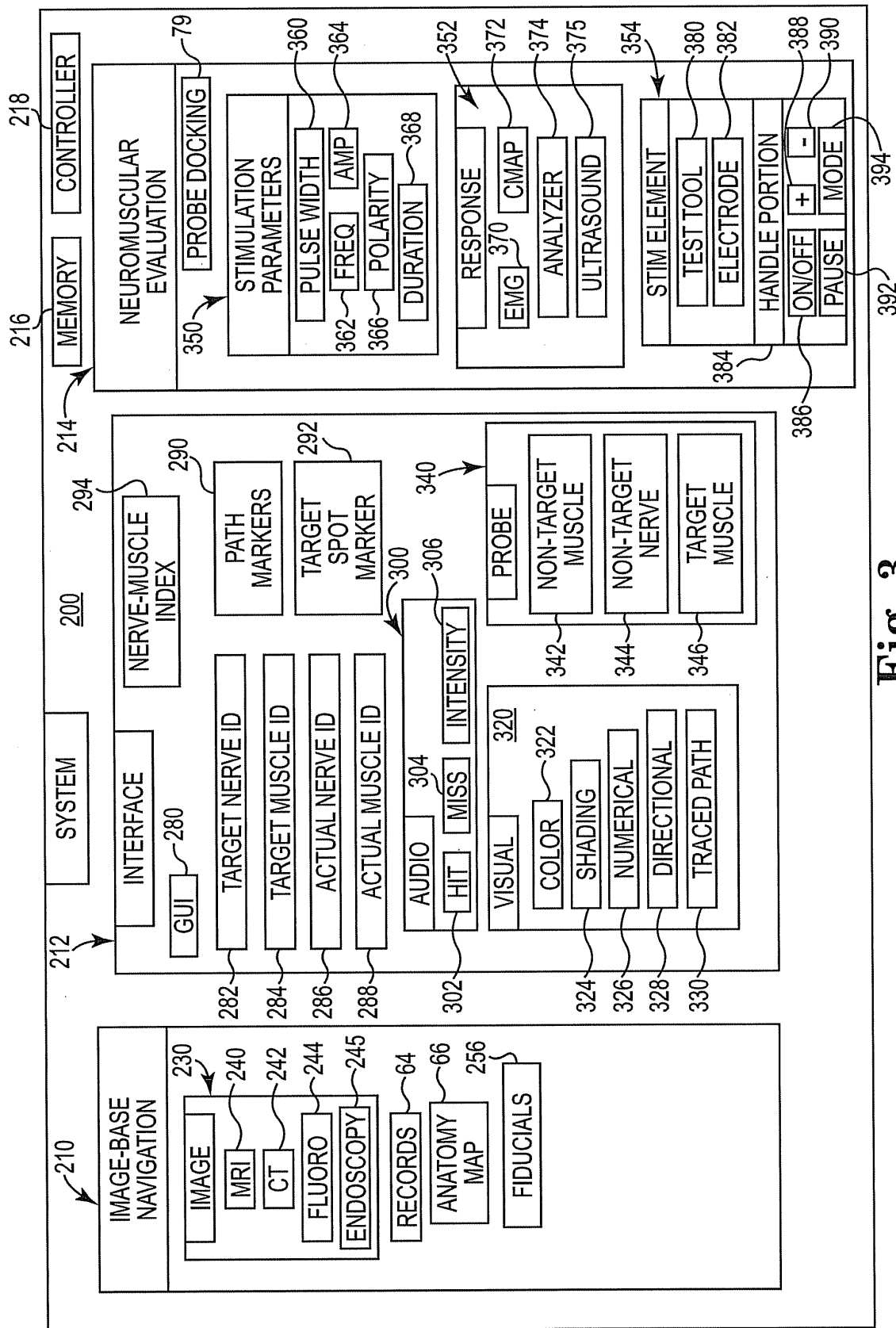


Fig. 3

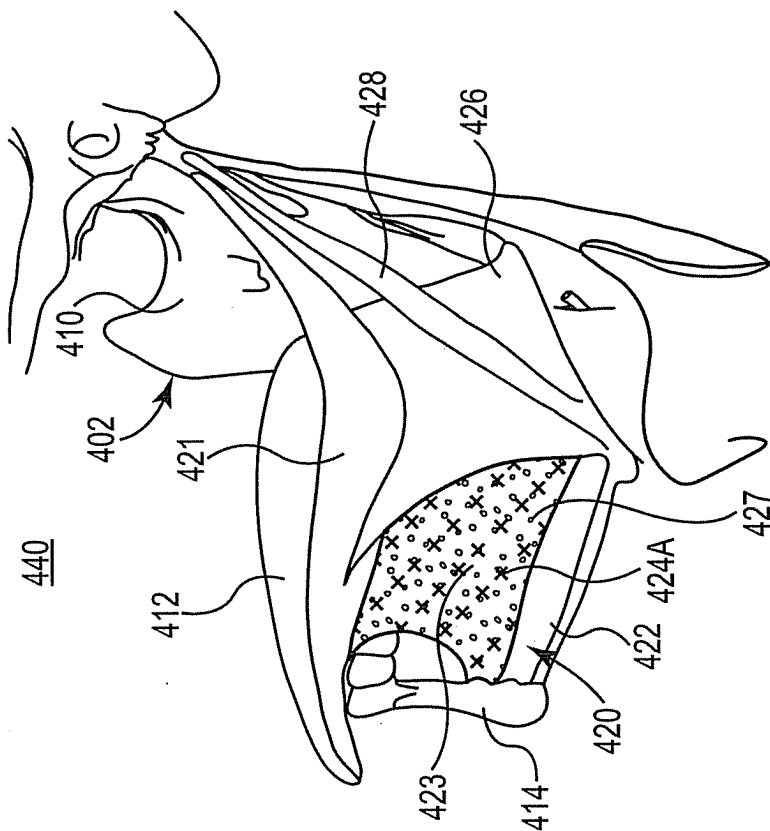


Fig. 5

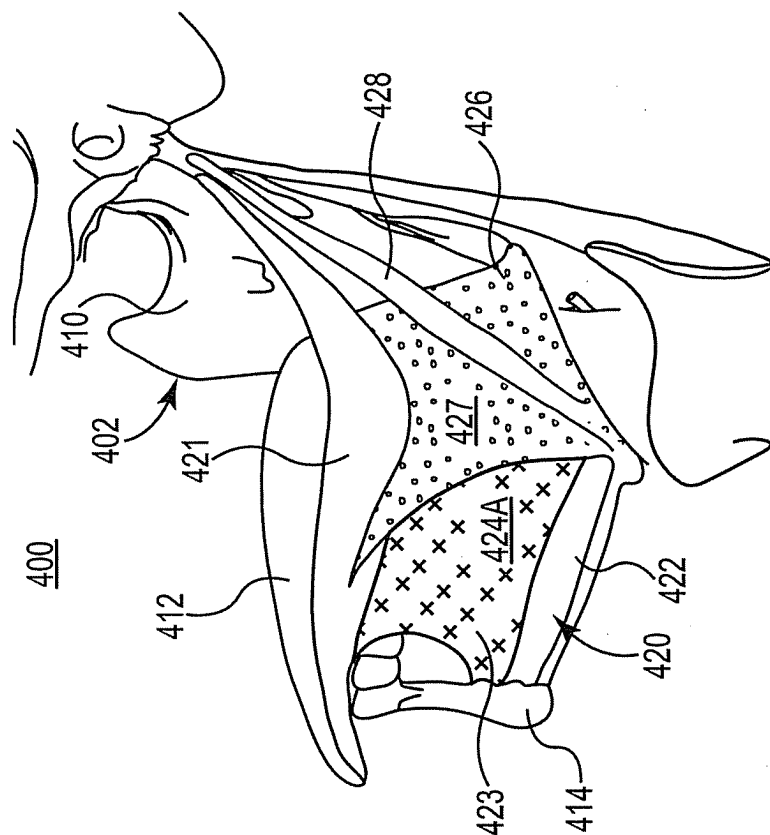
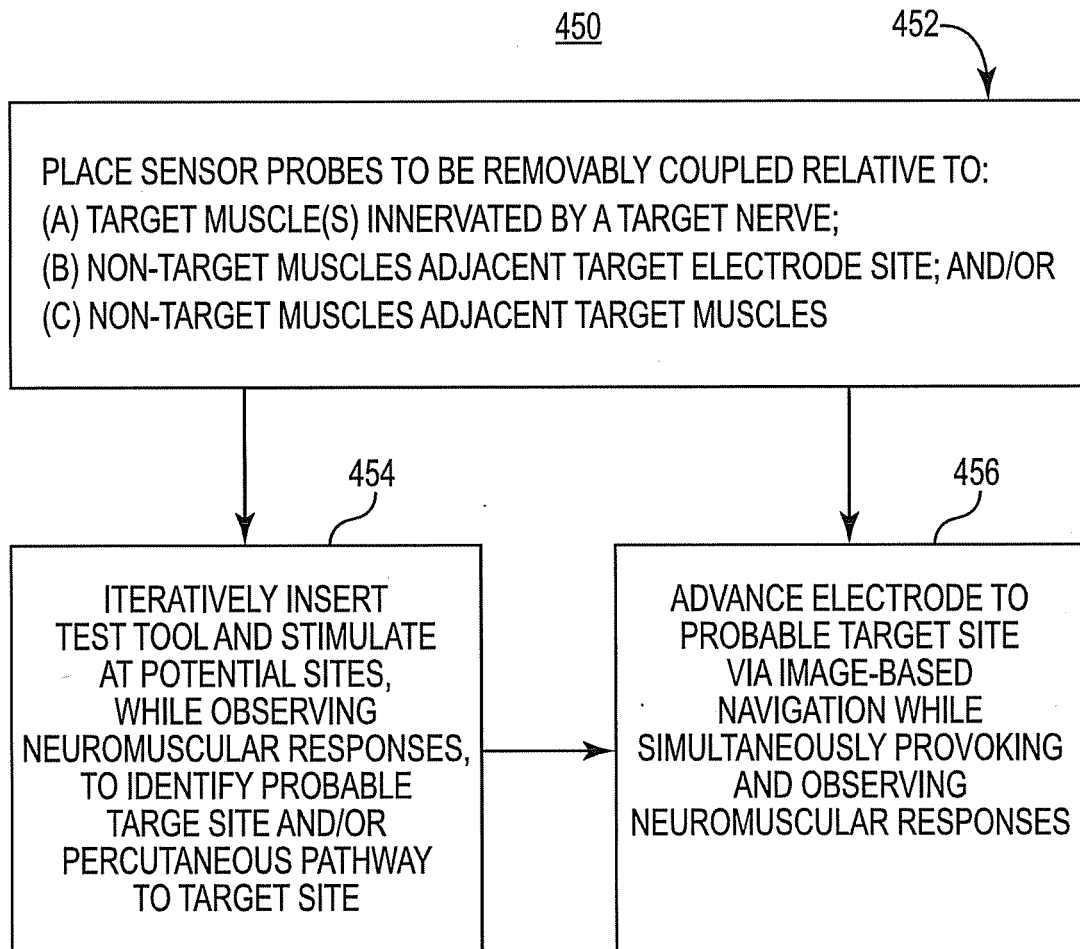


Fig. 4

**Fig. 6A**

[illegible]

Fig. 6B

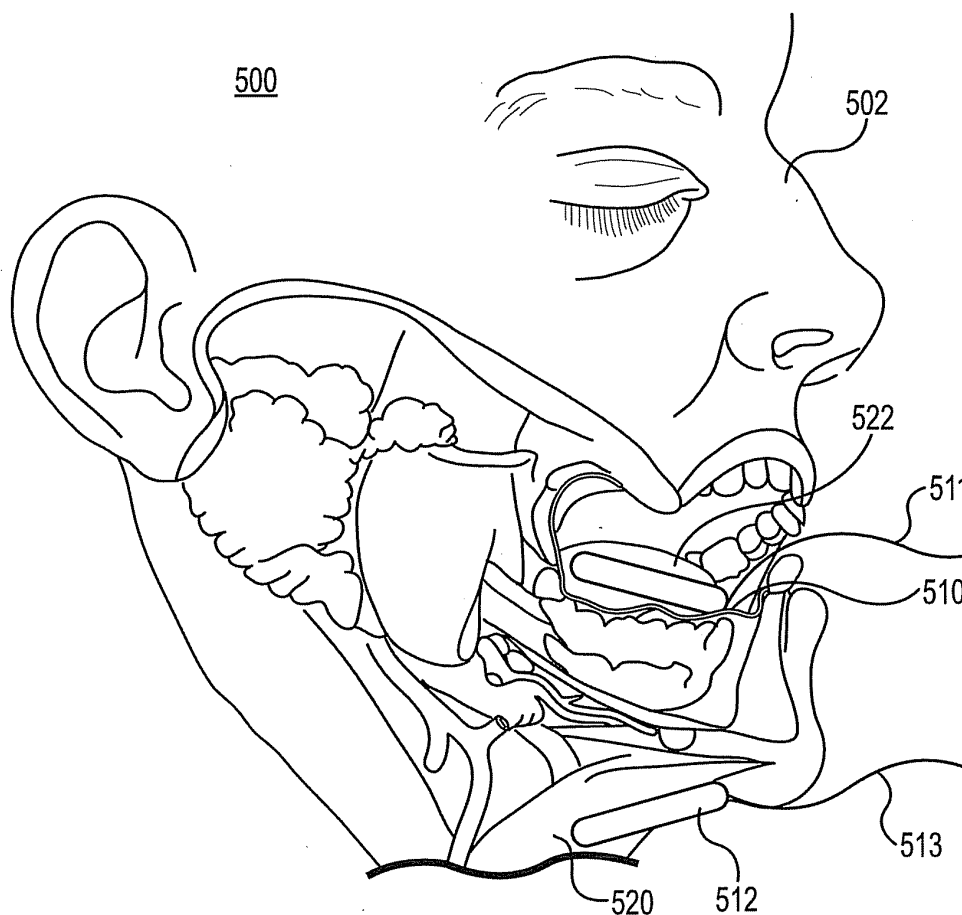


Fig. 7

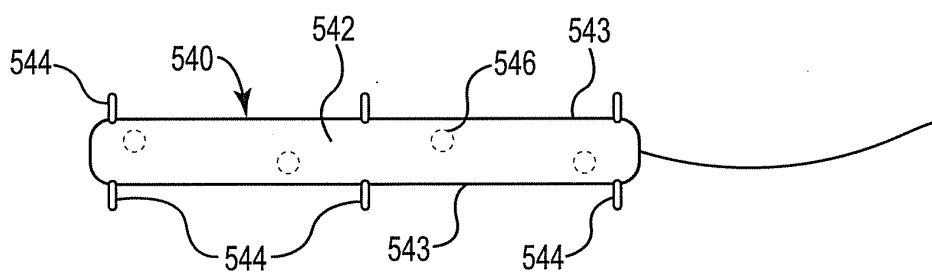


Fig. 8

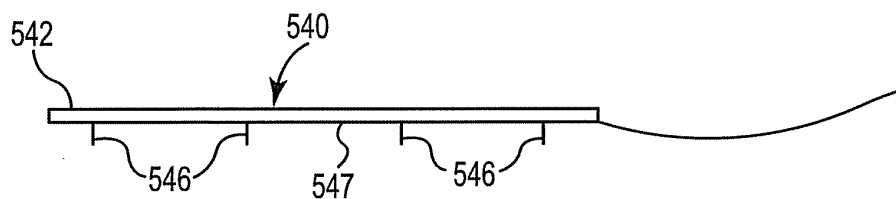


Fig. 9

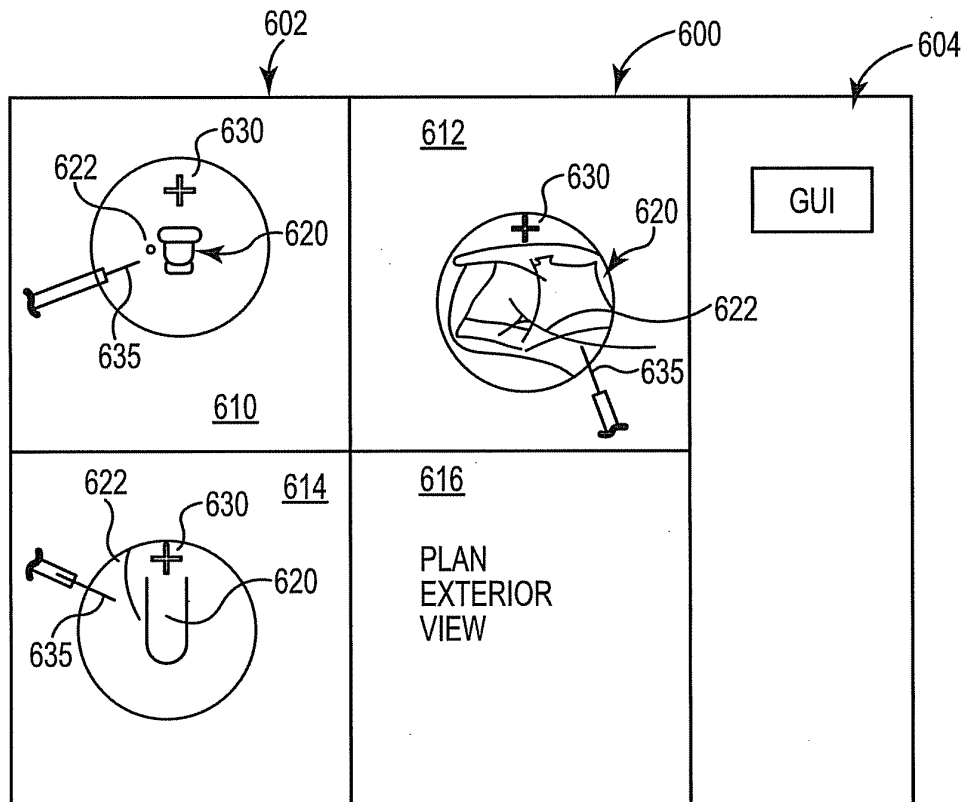


Fig. 10

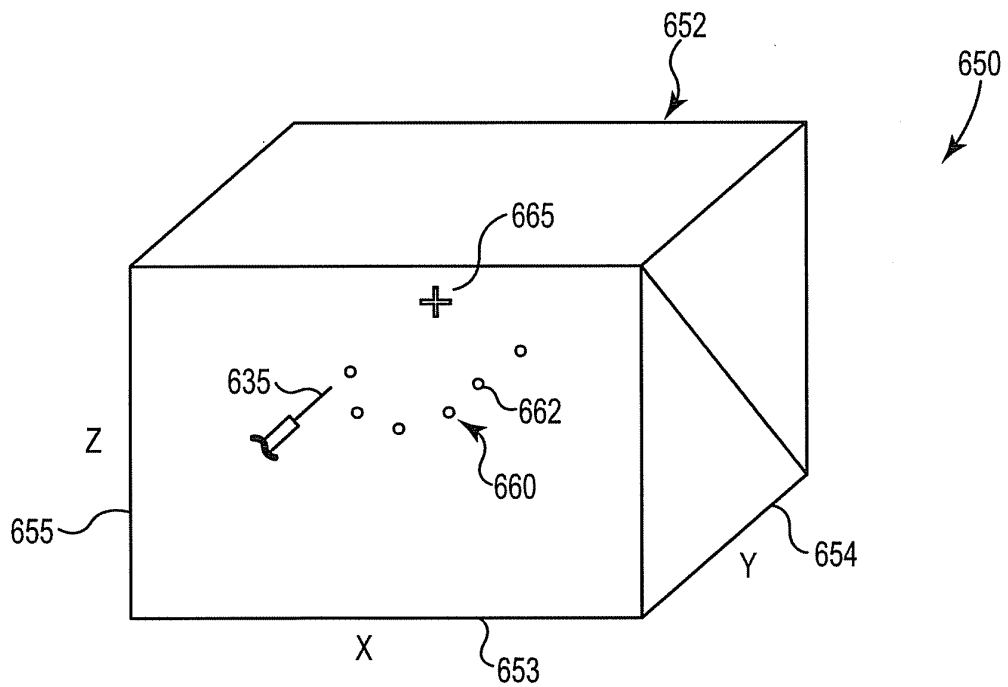


Fig. 11

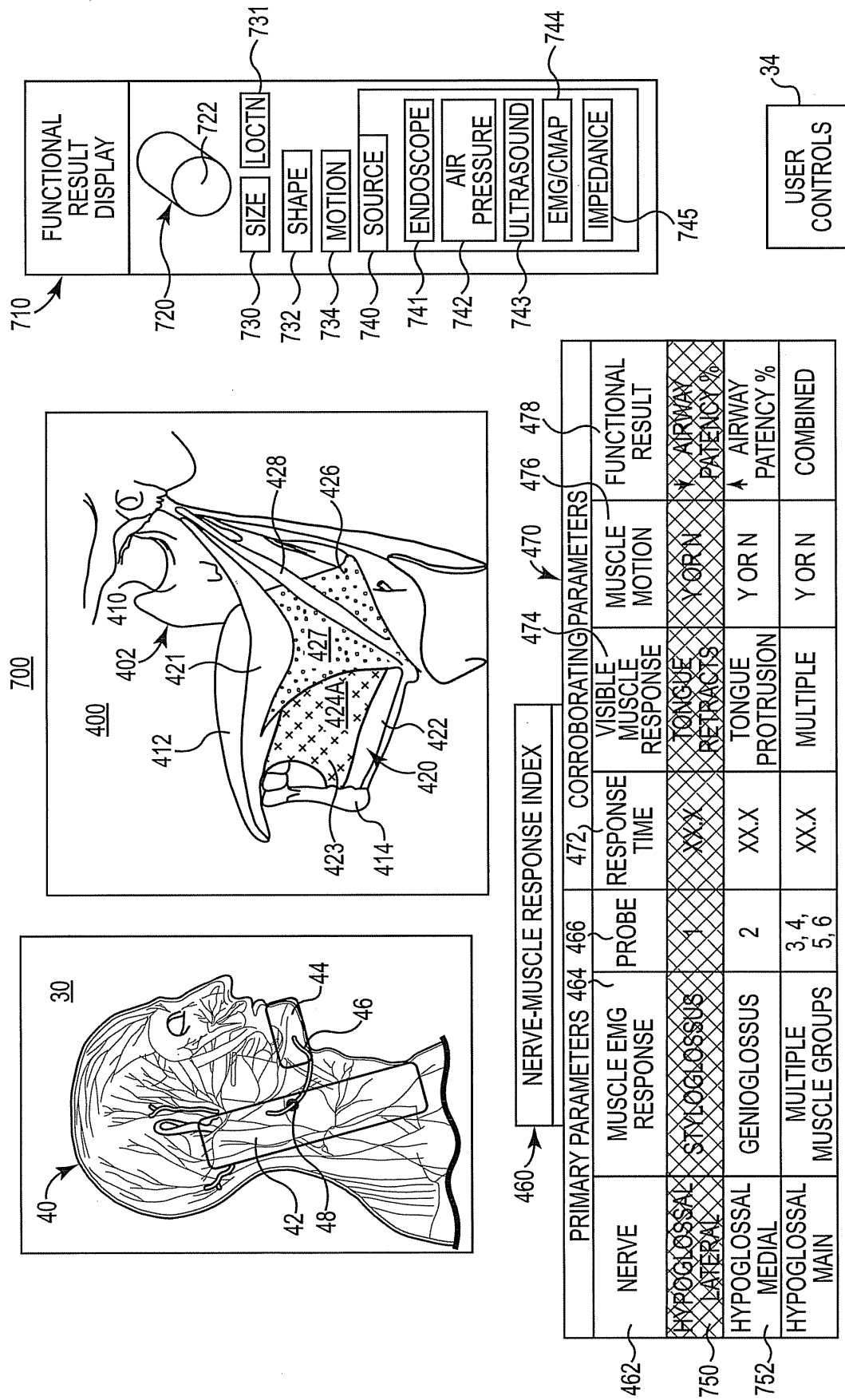


Fig. 12

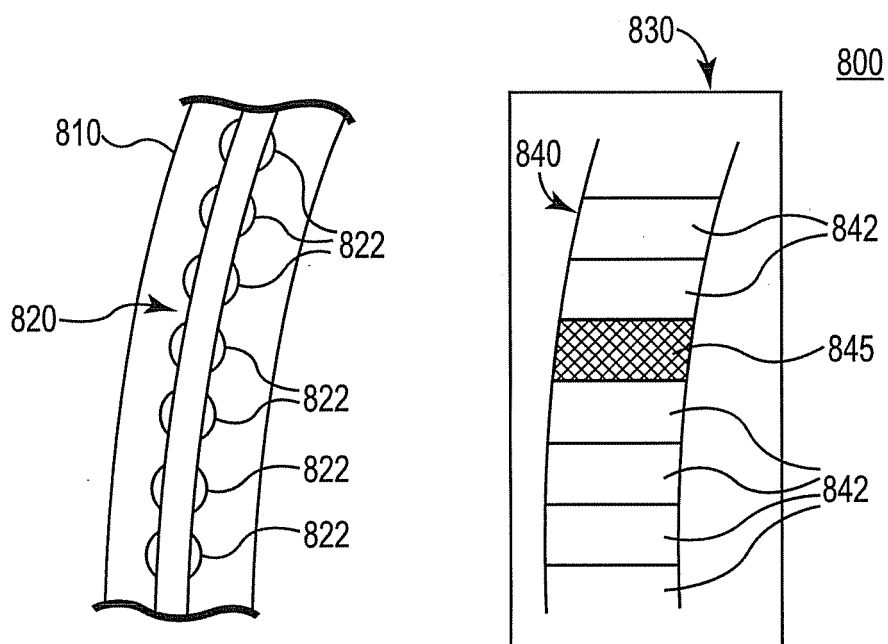


Fig. 13

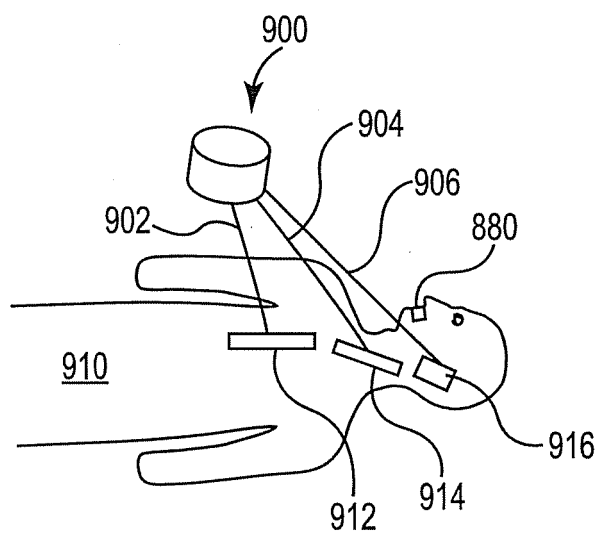


Fig. 14

REFERENCES CITED IN THE DESCRIPTION

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