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(54) **ENERGY-DELIVERY DEVICE INCLUDING
ULTRASOUND TRANSDUCER ARRAY AND
PHASED ANTENNA ARRAY**

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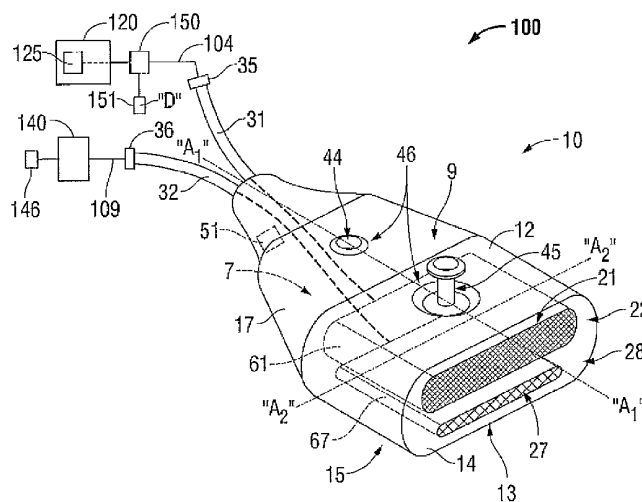
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(57) **ABSTRACT**

A medical device suitable for delivery of energy to tissue includes a housing, a phased antenna array disposed within the housing, and a user-interface coupled to the housing. The user-interface is adapted to enable a user to selectively adjust the radiation pattern of electromagnetic energy delivered into a tissue region by the phased antenna array. The medical device also includes an ultrasound transducer array disposed within the housing. The ultrasound transducer array is configured to acquire data representative of the tissue region during energy delivery into the tissue region by the phased antenna array.

19 Claims, 4 Drawing Sheets



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ENERGY-DELIVERY DEVICE INCLUDING ULTRASOUND TRANSDUCER ARRAY AND PHASED ANTENNA ARRAY

BACKGROUND

1. Technical Field

The present disclosure relates to electrosurgical devices suitable for tissue ablation applications and, more particularly, to an energy-delivery device including an ultrasound transducer array and a phased antenna array and systems including the same.

2. Discussion of Related Art

Treatment of certain diseases requires the destruction of malignant tissue growths, e.g., tumors. Electromagnetic radiation can be used to heat and destroy tumor cells. Treatment may involve inserting ablation probes into tissues where cancerous tumors have been identified. Once the probes are positioned, electromagnetic energy is passed through the probes into surrounding tissue.

In the treatment of diseases such as cancer, certain types of tumor cells have been found to denature at elevated temperatures that are slightly lower than temperatures normally injurious to healthy cells. Known treatment methods, such as hyperthermia therapy, heat diseased cells to temperatures above 41° C. while maintaining adjacent healthy cells below the temperature at which irreversible cell destruction occurs. These methods involve applying electromagnetic radiation to heat, ablate and/or coagulate tissue. Microwave energy is sometimes utilized to perform these methods. Other procedures utilizing electromagnetic radiation to heat tissue also include coagulation, cutting and/or ablation of tissue.

Electrosurgical devices utilizing electromagnetic radiation have been developed for a variety of uses and applications. A number of devices are available that can be used to provide high bursts of energy for short periods of time to achieve cutting and coagulative effects on various tissues. There are a number of different types of apparatus that can be used to perform ablation procedures. Typically, microwave apparatus for use in ablation procedures include a microwave generator that functions as an energy source, and a microwave surgical instrument (e.g., microwave ablation probe) having an antenna assembly for directing energy to the target tissue. The microwave generator and surgical instrument are typically operatively coupled by a cable assembly having a plurality of conductors for transmitting microwave energy from the generator to the instrument, and for communicating control, feedback and identification signals between the instrument and the generator.

There are several types of microwave probes in use, e.g., monopole, dipole and helical, which may be used in tissue ablation applications. In monopole and dipole antenna assemblies, microwave energy generally radiates perpendicularly away from the axis of the conductor. Monopole antenna assemblies typically include a single, elongated conductor. A typical dipole antenna assembly includes two elongated conductors that are linearly-aligned and positioned end-to-end relative to one another with an electrical insulator placed therebetween. Helical antenna assemblies include helically-shaped conductor configurations of various dimensions, e.g., diameter and length. The main modes of operation of a helical antenna assembly are normal mode (broadside), in which the field radiated by the helix is maximum in a perpendicular plane to the helix axis, and axial mode (end fire), in which maximum radiation is along the helix axis.

During certain procedures, a probe may be inserted directly into tissue, inserted through a lumen, e.g., a vein, needle or

catheter, or placed into the body using surgical techniques. Ultrasound or computed tomography (CT) guidance may be used prior to ablation treatments for aiding probe placement. Multiple probes may be used to synergistically create a large ablation or to ablate separate sites simultaneously.

The particular type of tissue ablation procedure may dictate a particular ablation volume in order to achieve a desired surgical outcome. Ablation volume is correlated with antenna design, antenna performance, antenna impedance, number of energy applicators used simultaneously, ablation time and wattage, and tissue characteristics, e.g., tissue impedance. During certain procedures, it can be difficult to assess the extent to which the microwave energy will radiate into the surrounding tissue, making it difficult to determine the area or volume of surrounding tissue that will be ablated.

Because of the small temperature difference between the temperature required for denaturing malignant cells and the temperature normally injurious to healthy cells, a known heating pattern and precise temperature control is needed to lead to more predictable temperature distribution to eradicate abnormal tissue structures, such as tumors, while minimizing the damage to surrounding normal tissue.

SUMMARY

The present disclosure relates to a medical device suitable for delivery of energy to tissue including a housing, a phased antenna array disposed within the housing, and a user-interface coupled to the housing. The user-interface is adapted to enable a user to selectively adjust the radiation pattern of electromagnetic energy delivered into a tissue region by the phased antenna array. The medical device also includes an ultrasound transducer array disposed within the housing. The ultrasound transducer array is configured to acquire data representative of the tissue region during energy delivery into the tissue region by the phased antenna array.

The present disclosure also relates to a system including an electrosurgical power generating source and a hand-holdable device operably associated with the electrosurgical power generating source. The hand-holdable device includes a phased antenna array, a user-interface coupled adapted to enable a user to selectively adjust the radiation pattern of electromagnetic energy delivered into a tissue region by the phased antenna array, and an ultrasound transducer array configured to acquire data representative of the tissue region during energy delivery into the tissue region by the phased antenna array.

BRIEF DESCRIPTION OF THE DRAWINGS

Objects and features of the presently-disclosed energy-delivery device including a ultrasound transducer array and a phased antenna array and systems including the same will become apparent to those of ordinary skill in the art when descriptions of various embodiments thereof are read with reference to the accompanying drawings, of which:

FIG. 1 is a perspective and schematic view of a system including an energy-delivery (medical) device including an ultrasound transducer array and a phased antenna array in accordance with an embodiment of the present disclosure;

FIG. 2 is an enlarged, perspective view of a pointing device shown with two schematically-illustrated axis (shown by double arrowheaded lines) representative of indicative orientations of the pointing device in accordance with an embodiment of the present disclosure;

FIG. 3 is a schematic diagram of an energy-delivery system including a radiant electromagnetic energy transmissive

structure disposed at the distal end of a phased antenna array in accordance with an embodiment of the present disclosure;

FIG. 4 is a top, perspective view of the medical device of FIG. 1 showing the pointing device of FIG. 2 positioned in a first indicative orientation and showing a diagrammatic representation of a radiation pattern of electromagnetic energy delivered into tissue by the medical device responsive to the first indicative orientation of the pointing device in accordance with an embodiment of the present disclosure;

FIG. 5 is a top, perspective view of the medical device of FIG. 1 showing the pointing device of FIG. 2 positioned in a second indicative orientation and showing a diagrammatic representation of a radiation pattern of electromagnetic energy delivered into tissue by the medical device responsive to the second indicative orientation of the pointing device in accordance with an embodiment of the present disclosure;

FIG. 6 is a top, perspective view of the medical device of FIG. 1 showing the pointing device of FIG. 2 positioned in a third indicative orientation and showing a diagrammatic representation of a radiation pattern of electromagnetic energy delivered into tissue by the medical device responsive to the third indicative orientation of the pointing device in accordance with an embodiment of the present disclosure;

FIG. 7 is a top, perspective view of the medical device of FIG. 1 showing the pointing device of FIG. 2 positioned in a fourth indicative orientation and showing a diagrammatic representation of a radiation pattern of electromagnetic energy delivered into tissue by the medical device responsive to the fourth indicative orientation of the pointing device in accordance with an embodiment of the present disclosure; and

FIG. 8 is a schematic diagram of a control system in accordance with an embodiment of the present disclosure.

DETAILED DESCRIPTION

Hereinafter, embodiments of an energy-delivery device (also referred to herein as a “medical device” or a “handheld device”) including an ultrasound transducer array and a phased antenna array and systems including the same of the present disclosure are described with reference to the accompanying drawings. Like reference numerals may refer to similar or identical elements throughout the description of the figures. As shown in the drawings and as used in this description, and as is traditional when referring to relative positioning on an object, the term “proximal” refers to that portion of the apparatus, or component thereof, closer to the user and the term “distal” refers to that portion of the apparatus, or component thereof, farther from the user.

This description may use the phrases “in an embodiment,” “in embodiments,” “in some embodiments,” or “in other embodiments,” which may each refer to one or more of the same or different embodiments in accordance with the present disclosure. For the purposes of this description, a phrase in the form “A/B” means A or B. For the purposes of the description, a phrase in the form “A and/or B” means “(A), (B), or (A and B)”. For the purposes of this description, a phrase in the form “at least one of A, B, or C” means “(A), (B), (C), (A and B), (A and C), (B and C), or (A, B and C)”.

Electromagnetic energy is generally classified by increasing energy or decreasing wavelength into radio waves, microwaves, infrared, visible light, ultraviolet, X-rays and gamma-rays. As it is used in this description, “microwave” generally refers to electromagnetic waves in the frequency range of 300 megahertz (MHz) (3×10^8 cycles/second) to 300 gigahertz (GHz) (3×10^{11} cycles/second).

As it is used in this description, “ablation procedure” generally refers to any ablation procedure, such as, for example, microwave ablation, radiofrequency (RF) ablation, or microwave or RF ablation-assisted resection. As it is used in this description, “energy applicator” generally refers to any device that can be used to transfer energy from a power generating source, such as a microwave or RF electro-surgical generator, to tissue. For the purposes herein, the term “energy applicator” is interchangeable with the term “energy-delivery device”. As it is used in this description, “transmission line” generally refers to any transmission medium that can be used for the propagation of signals from one point to another.

As it is used in this description, “phased antenna array” generally refers to any multi-element antenna array capable of shifting the phase of the signal emitted from each radiating element, to provide constructive/destructive interference so as to steer the antenna beam in the desired direction. For the purposes herein, the term “radiating element” is interchangeable with the term “antenna element”. As it is used in this description, “electromagnetic window” generally refers to any and all types of radomes and windows through which electromagnetic signals are passed in use.

As it is used in this description, “length” may refer to electrical length or physical length. In general, electrical length is an expression of the length of a transmission medium in terms of the wavelength of a signal propagating within the medium. Electrical length is normally expressed in terms of wavelength, radians or degrees. For example, electrical length may be expressed as a multiple or sub-multiple of the wavelength of an electromagnetic wave or electrical signal propagating within a transmission medium. The wavelength may be expressed in radians or in artificial units of angular measure, such as degrees. The electric length of a transmission medium may be expressed as its physical length multiplied by the ratio of (a) the propagation time of an electrical or electromagnetic signal through the medium to (b) the propagation time of an electromagnetic wave in free space over a distance equal to the physical length of the medium. The electrical length is in general different from the physical length. By the addition of an appropriate reactive element (capacitive or inductive), the electrical length may be made significantly shorter or longer than the physical length.

As used in this description, the term “real-time” means generally with no observable latency between data processing and display. As used in this description, “near real-time” generally refers to a relatively short time span between the time of data acquisition and display.

Various embodiments of the present disclosure provide an ultrasound transducer array and a phased antenna array incorporated into one, direct-to-patient contact device capable of directing electromagnetic energy into tissue. The presently-disclosed energy-delivery devices including an ultrasound transducer array and a phased antenna array are adapted to enable user control of the radiation pattern of electromagnetic energy delivered into tissue, and may be suitable for use in a variety of procedures and operations. Various embodiments of the presently-disclosed energy-delivery device including an ultrasound transducer array and a phased antenna array are adapted to be hand-holdable and include an ergonomically located user-interface.

Various embodiments of the presently-disclosed energy-delivery device including an ultrasound transducer array and a phased antenna array are adapted to enable user-controllable focal location of electromagnetic energy delivery into tissue to depths ranging from about one centimeter (cm) to about three centimeters, e.g., in relation to a tissue surface, at an operational frequency between about 1 GHz and about 5

GHZ. Embodiments may enable user-controllable focal location of electromagnetic energy delivery into tissue to a variable predetermined depth or range of depths. In the case of a 3 cm ablation that is focally located 3 cm deep, for example, tissue 4.5 cm deep can be treated. By enlarging the device, decreasing operational frequency and/or increasing the number of array elements, deeper tissue may be treatable. In some embodiments, data acquired by the ultrasound transducer array may be outputted from the energy-delivery device to an ultrasound imaging system, and may be outputted from the imaging system to one or more display devices, which may be used by the clinician to visualize the targeted region in real-time and/or near real-time.

The presently-disclosed energy-delivery device including an ultrasound transducer array and a phased antenna array according to various embodiments is designed and configured to operate between about 300 MHz and about 10 GHz. Embodiments may be implemented using electromagnetic radiation at microwave frequencies, RF frequencies or at other frequencies.

Various embodiments of the presently-disclosed energy-delivery device including an ultrasound transducer array and a phased antenna array are suitable for microwave or RF ablation and for use to pre-coagulate tissue for microwave or RF ablation-assisted surgical resection. Although various methods described hereinbelow are targeted toward microwave ablation and the complete destruction of target tissue, it is to be understood that methods for directing electromagnetic radiation may be used with other therapies in which the target tissue is partially destroyed or damaged. In addition, although the following description describes the use of a microwave phased antenna array, the teachings of the present disclosure may also apply to other type of user-controllable phased antenna array.

An electrosurgical system including an energy-delivery device including an ultrasound transducer array and a phased antenna array according to various embodiments is capable of providing real-time and/or near real-time image feedback during electromagnetic energy-induced thermal therapy, e.g., to allow the clinician to better visualize and understand how to achieve more optimized results during thermal treatment of tissue.

FIG. 1 shows an electrosurgical system (shown generally as **100**) according to an embodiment of the present disclosure that includes an energy-delivery device **10** including an ultrasound transducer array **67** and a microwave phased antenna array **61**. Microwave phased antenna array **61** generally includes a plurality of radiating elements (e.g., "A₁", "A₂", "A₃", "A₄" through "A_N" shown in FIG. 3) positioned to form a desired number of rows and columns. In some embodiments, the radiating elements may be aperture (waveguide) or linear (dipole) antennas operating at S, L, or C band frequencies. In some embodiments, the radiating elements may be spiral, dipole, slot, or any type of microstrip antenna, e.g., a patch antenna (also known as a rectangular microstrip antenna), and may be formed on a substrate, such as a dielectric sheet material, e.g., using conventional printed circuit board (PCB) fabrication techniques.

Ultrasound transducer device **67** (also referred to herein as an "ultrasound transducer array") may be any suitable device capable of generating, transmitting and receiving ultrasound waves. Ultrasound transducer device **67** may include a one-dimensional or multi-dimensional array of transducer elements (not shown). Ultrasound transducer device **67** may be adapted for amplifying the reflected ultrasound signal received by the ultrasound transducer device **67**. In some embodiments, ultrasound transducer array **67** includes a plu-

ality of transducer elements that are individually controllable and operable to form a two-dimensional array, e.g., suitable for scanning a volumetric region in three dimensions. Individual transducer elements may be individually selectable and operable together to form a one-dimensional array, e.g., suitable for scanning a planar region in two dimensions. Ultrasound transducer array **67** may be adapted to produce an image over a wide field of view, such as a sector scan image produced by repeatedly transmitting and receiving ultrasound energy in radial directions from the medical device **10**. Ultrasound imaging may allow the clinician to observe the relationship between abnormal tissue structures, such as tumors, and normal tissue structures, such as vessels and tissue boundaries, during treatments.

Energy-delivery device **10** includes a housing **15** generally defining a first axis "A₁"-"A₁", e.g., a central longitudinal axis, and a second axis "A₂"-"A₂" disposed perpendicular to the first axis "A₁"-"A₁". In some embodiments, the housing **15** is formed from two housing halves (not shown). Each half of the housing **15** may include a series of mechanical interfacing components (not shown) configured to matingly engage with a corresponding series of mechanical interfaces (not shown) to align the two housing halves about the inner components and assemblies of the energy-delivery device **10**.

As shown in FIG. 1, the housing **15** includes a body member **17** including a distal end **13**. Body member **17** defines a tissue-contact surface **14** at the distal end **13**, a top surface **12** including a distal edge coupled to the tissue-contact surface **14**, and an internal chamber **7** configured to contain the ultrasound transducer device **67** and the microwave phased antenna array **61** therein. Tissue-contact surface **14** may have any suitable configuration, e.g., a flat, planar or curved configuration, and may be disposed generally perpendicular to the top surface **12**.

Tissue-contact surface **14** generally includes one or more regions defining one or more electromagnetic windows through which electromagnetic signals are passed in use. In some embodiments, the tissue-contact surface **14** includes a first region **28** defining an ultrasound transmissive window **27** and a second region **22** defining a microwave transmissive window **21**. As shown in FIG. 1, the first region **28** corresponds to a lower portion of the tissue-contact surface **14**, and the second region **22** corresponds to an upper portion of the tissue-contact surface **14**. Ultrasound transducer device **67** operations may involve directing ultrasound energy through the ultrasound transmissive window **27** and receiving ultrasound energy through the ultrasound transmissive window **27**.

Ultrasound transmissive window **27** and the microwave transmissive window **21** may be composed of low-loss dielectric materials. It will be appreciated that the ultrasound transmissive window **27** and the microwave transmissive window **21** may be disposed in any suitable relation to one another, such as one above (or below) the other, and may have any suitable shape, e.g., depending on the particular configuration of the ultrasound transducer device **67** and/or the microwave phased antenna array **61** housed within the body member **17**.

Body member **17**, or portion thereof, may be formed from metal, thermoplastic, e.g., polycarbonate, composites, e.g., plastic-metal or ceramic-metal composites, or other materials, and may be configured to be hand-holdable. The design and/or material of the ultrasound transmissive window **27** and the microwave transmissive window **21** may differ compared to one or more structural parts of the tissue-contact surface **14**, e.g., to achieve desired electrical performance. The size and shape of the housing **15** may be varied from the configuration depicted in FIG. 1.

As shown in FIG. 1, electro-surgical system 100 generally includes an electro-surgical power generating source 120, e.g., a microwave or RF electro-surgical generator, a user-interface 46 associated with the energy-delivery device 10, and a processor unit 150 communicatively coupled with the phased antenna array 61. User-interface 46 may be communicatively coupled with the processor unit 150 and/or other processor unit (not shown). Electro-surgical system 100 may include an ultrasonic imaging system 140 communicatively coupled with the ultrasound transducer array 67. Ultrasonic imaging system 140 may be connected to one or more display devices and/or screens 146 (e.g., LCD (liquid crystal display), plasma, OLED (organic light emitting diode), holographic, flat, and the like) for displaying output from the ultrasonic imaging system 140, which may allow clinicians to visualize the ablative process in real-time and/or near real-time.

User-interface 46 may be adapted to cooperatively operate with the processor unit 150 and/or other processor (not shown) to enable the user to selectively-control one or more parameters of electromagnetic energy delivery into tissue by the medical device 10. User-interface 46 may be disposed on, or otherwise associated with, the housing 15, e.g., ergonomically located on the top surface 12 of the body member 17. In some embodiments, the user-interface 46 includes a pointing device 45, e.g., a joystick, trackball, or the like, communicatively coupled to the processor unit 150.

In some embodiments, user-effected movement of the pointing device 45 is defined with respect to "X" and "Y" axes (schematically shown by double arrowheaded lines in FIG. 2), representative of indicative orientations of the pointing device 45. The axis "Y" may be oriented in a direction parallel to the first axis "A₁"-"A₁" of the housing 15, and the axis "X" may be oriented in a direction parallel to the second axis "A₂"-"A₂" of the housing 15. As described in more detail later in this description, one or more parameters of electromagnetic energy delivery into tissue by the medical device 10 may be correlated to the indicative orientations of the pointing device 45.

Pointing device 45 may be ergonomically located on the top surface 12 of the body member 17 such that the user can control the pointing device 45 easily with thumb, finger, or palm. As an alternative to (or in addition to) the pointing device 45, the user-interface 46 may include voice input technology, including, for example, hardware and/or software incorporated in the processor unit 150, or a separate digital module connected to the processor unit 150. The voice input technology may include voice recognition, voice activation, voice rectification, and/or embedded speech.

User-interface 46 may additionally, or alternatively, include a power on/off switch 44. The power on/off switch 44 may be disposed on, or otherwise associated with, the housing 15, e.g., ergonomically located on the top surface 12, and may have any suitable configuration, e.g., rotatable knobs, depressable buttons, toggle switches, slide switches, voice or sound actuated switches, or any other suitable device capable of turning off power to the medical device 10. The power on/off switch 44 may be implemented as a remotely operable device, such as a footswitch, a handswitch, or an orally-activated switch. User-interface 46 may additionally, or alternatively, include an indicator (not shown), such as an audible and/or visual indicator, e.g., an illuminated indicator (e.g., a single- or variably-colored LED indicator), to alert or signal the user that power is turned on/off.

User-interface 46 may be adapted to cooperatively operate with the processor unit 150 to enable the user to selectively-steer the focal point of energy delivery in tissue to various locations and/or to enable the user to the control the energy

deposition pattern, e.g., an ablation field radiating into tissue. One or more electrical signals outputted from the user-interface 46, e.g., responsive to a user-effected movement of the pointing device 45, received by the processor unit 150 may be used to determine and set the phasing of radiating elements of the microwave phased antenna array 61, e.g., to allow the focal point of energy delivery in tissue to be varied in position in real-time and/or near real-time.

Processor unit 150 may include any type of computing device, computational circuit, or any type of processor or processing circuit capable of executing a series of instructions that are stored in a memory, e.g., memory 151, associated with the processor unit 150. Processor unit 150 may be adapted to run an operating system platform and application programs. Although the processor unit 150 is illustrated as a standalone module in FIG. 1, it is to be understood that the processor unit 150 may be integrated fully or partially into the electro-surgical power generating source 120, or other component of the electro-surgical system 100. Medical device 10 may be configured with a memory 51 disposed within the body member 17 and communicatively coupled with the processor unit 150 and/or communicatively coupled with an internal processor (not shown).

Processor unit 150 may receive user inputs from the user-interface 46, such as an electric signal indicative of the position and/or a relative movement of the pointing device 45, e.g., a joystick or trackball, and/or other device communicatively coupled to the processor unit 150. In some embodiments, data "D" (representative of a mapping of the indicative orientations of the pointing device 45 to settings for properly phasing the phased antenna array 61 to achieve desired radiation patterns) is stored in a suitable memory for use by the processor 150, e.g., to enable steering of the beam and/or the focal point of energy delivery in the desired direction and/or to the desired location in tissue. Data "D" may be stored in any suitable data structure, such as a look-up table or other data structure. Data "D" may be stored in a memory 51 (internal to medical device 10) and/or stored in a memory 151 (external to medical device 10). In some embodiments, data "D" may be stored in a library (not shown) communicatively coupled to processor 150. As it is used in this description, "library" generally refers to any repository, databank, database, cache, storage unit and the like.

Electro-surgical power generating source 120 may be any generator suitable for use with electro-surgical devices, and may be configured to provide various frequencies of electromagnetic energy. In some embodiments, the electro-surgical power generating source 120 is configured to provide microwave energy at an operational frequency from about 300 MHz to about 10 GHz. An example of an electro-surgical generator that delivers 915 MHz, which may be suitable for use as a source of electro-surgical energy, is commercially available under the trademark EVIDENT™ Microwave Ablation Generator offered by Covidien.

Electro-surgical power generating source 120 may include a user-interface 125 in operable communication with processor unit 150. Electro-surgical power generating source 120 may include a database configured to store and retrieve energy applicator data, e.g., parameters associated with one or more energy-delivery devices. In use, the clinician may interact with the user-interface 125 to preview operational characteristics of an energy-delivery device, such as, for example, medical device 10. User-interface 125 may include a display device (not shown) adapted to visually display one or more user-interface elements. The display device may include touchscreen capability, e.g., the ability to receive user input through direct physical interaction with the display

device, e.g., by contacting the display panel of the display device with a stylus or fingertip.

Microwave phased antenna array **61** may be operably coupled to the processor unit **150** and/or the electrosurgical power generating source **120** by a cable connection or a wireless connection, e.g., a radiofrequency or infrared link. In some embodiments, energy-delivery device **10** includes a first cable assembly **31** operably coupled to a first connector **35**, which further operably connects the phased antenna array **61** via a first transmission line **104** to the processor unit **150**. First cable assembly **31** may have a proximal end suitable for connection to the electrosurgical energy source **120**.

Energy-delivery device **10** may additionally, or alternatively, include a second cable assembly **32** operably coupled to a second connector **36**, which further operably connects the ultrasound transducer device **61** via a second transmission line **109** to the ultrasonic imaging system **140**. Second cable assembly **32** may have a proximal end suitable for connection to the ultrasonic imaging system **140**.

In some embodiments, data acquired from the ultrasound transducer array **61** is outputted from the energy-delivery device **100** to the ultrasound imaging system **140**, e.g., for processing to provide an image format suitable for display, and may be outputted from the imaging system **140** to one or more display devices **146**, which may be used by the clinician to visualize the targeted region and/or the ablation isotherm volume in real-time or near real-time during a procedure. During activation of the ultrasound transducer array **61**, a bubble field or cloud of micro-fine bubbles may be generated in the targeted region, e.g., resulting from thermally-induced mass phase transition (e.g., liquid-gas phase transition), and may be visibly observable within the ultrasound imaging. Observation of the temporal evolution and spatial distribution of the bubble cloud generated in the target region may allow clinicians to better visualize and understand how to achieve more optimized results during thermal treatment of tissue, e.g., to allow clinicians to avoid ablating sensitive structures, such as large vessels, healthy organs or vital membrane barriers.

Electrosurgical system **100** may include a coolant supply system (e.g., **350** shown in FIG. **3**) coupled in fluid communication with one or more components of the medical device **10**. In some embodiments, the coolant supply system may be adapted to circulate coolant fluid (e.g., “F” shown in FIG. **3**) into and out of an electromagnetic window (e.g., **390** shown in FIG. **3**) disposed at the distal end **13** of the housing **15**.

During microwave ablation, e.g., using the electrosurgical system **100**, the medical device **10** is placed adjacent to tissue and microwave energy is supplied thereto. A clinician may pre-determine the length of time that microwave energy is to be applied. Application duration may depend on many factors such as tumor size and location and whether the tumor was a secondary or primary cancer. The duration of microwave energy application using the medical device **10** may depend on the progress of the heat distribution within the tissue area that is to be destroyed and/or the surrounding tissue. Treatment of certain tumors may involve probe repositioning during the ablation procedure, such as where the tumor is larger than the probe or has a shape that does not correspond with available probe geometry or radiation pattern.

User-interface **46** may include indicia thereon representative of one or more user-selectable parameters of electromagnetic energy delivery into tissue by the medical device **10**, e.g., a first scale “S₁” and a second scale “S₂”. As shown in FIG. **2**, the first scale “S₁” includes indicia graduation marks and angle in degrees (e.g., 45°, 0°, 45°), and the second scale “S₂” includes indicia graduation marks and a series of con-

secutive positive integers (e.g., 1, 2, 3) corresponding to increasing levels of energy intensity indicative of energy intensity levels. The indicia may be etched, stamped, formed or the like, e.g., on the upper surface **12** and neighboring the pointing device **45**. The design of the indicia may be varied from the configuration depicted in FIG. **2**.

One or more parameters of electromagnetic energy delivery into tissue by the medical device **10** may be correlated to indicative orientations of the pointing device **45**. User-effected movement of the pointing device **45** may be defined in terms of movement in a first direction (e.g., an X-axis direction) and movement in a second direction (e.g., a Y-axis direction) perpendicular to the first direction. Signals outputted from the pointing device **45** representative of indicative orientations of the pointing device **45** may be correlated to one or more parameters of electromagnetic energy delivery into tissue.

In some embodiments, user-effected movement of the pointing device **45** in a first direction (e.g., an X-axis direction), a second direction (e.g., a Y-axis direction) and/or a third direction (e.g., a Z-axis direction) is correlated to a predetermined phasing of the phased antenna array **61**, to enable steering of the beam and/or steering of the focal point of energy delivery by the medical device **10** in the desired direction and/or to the desired location in tissue “T”.

In some embodiments, medical device **10** is configured to adjust power parameters (e.g., voltage, power and/or current intensity) and/or the power versus impedance curve shape to affect the perceived output intensity, responsive to user-effected movement of the pointing device **45** in a first direction (e.g., an X-axis direction). For example, the greater the lateral displacement of the pointing device **45** in a distal direction, the greater the level of the power parameters transmitted to the phased antenna array **61**. Intensity settings may be preset and selected from a look-up table, e.g., based on a configuration of the radiating elements of the phased antenna array **61**, desired surgical effect, surgical specialty and/or surgeon preference. The selection may be made automatically or selected manually by the user. The intensity values may be predetermined or adjusted by the user.

FIG. **3** is schematic diagram of an embodiment of an energy-delivery system (shown generally as **300**) that includes a signal source **310**, a phased antenna array **360** coupled to the signal source **310**, and a radiant electromagnetic energy transmissive structure **390** (also referred to herein as an “electromagnetic window”) disposed at the distal end of the phased antenna array **360**. Signal source **310** is generally configured to provide microwave frequency output signals.

Phased antenna array **360** includes a microwave amplifier unit **320** coupled to the signal source **310**, a microwave power splitter **330** coupled to the microwave amplifier unit **320**, a controller **340** coupled to the microwave power splitter **330**, and a plurality of radiating elements “A₁”, “A₂”, “A₃”, “A₄” through “A_N” coupled to the controller **340**. Microwave amplifier unit **320** may have any suitable input power and output power. Power splitter **330** may be implemented by a variety of components, including without limitation, coplanar striplines, coplanar waveguides, Wilkinson power dividers, and/or other suitable power dividers. In some embodiments, the power splitter **330** may be implemented by any suitable power divider that provides an equal or unequal power split at its output ports while substantially maintaining a predetermined phase relationship.

Controller **340** generally includes a plurality of phase shifters “S₁”, “S₂”, “S₃”, “S₄” through “S_N”. Controller **340** may include a number of processor units (not shown) coupled

to the phase shifters “S₁”, “S₂”, “S₃”, “S₄” through “S_N” for controlling output of one or more of the phase shifters “S₁” through “S_N” to provide a desired phase relationship of electrical signals in each channel of the phased antenna array 360. The processing units may include multiple processors and/or multicore CPUs and may include any type of processor capable of executing software, such as a microprocessor, digital signal processor, microcontroller, or the like.

Energy-delivery system 300 includes an electromagnetic window 390 disposed between the phased antenna array 360 and tissue “T”. Electromagnetic window 390 may include a water bolus, or other dielectric material. In some embodiments, the electromagnetic window 390 is coupled in fluid communication with a coolant supply system 350 including a coolant source 355.

Coolant source 355 may be any suitable housing containing a reservoir of coolant fluid “F”, and may maintain coolant fluid “F” at a predetermined temperature. For example, the coolant source 355 may include a cooling unit (not shown) capable of cooling the returning coolant fluid “F” from the electromagnetic window 390. Coolant fluid “F” may be any suitable fluid that can be used for cooling or buffering the electromagnetic window 390, e.g., deionized water, or other suitable cooling medium. Coolant fluid “F” may have dielectric properties and may provide dielectric impedance buffering for the phased antenna array 360. Various fluids may be used, e.g., liquids including, but not limited to, water, saline, perfluorocarbon, such as the commercially available Fluoriner® perfluorocarbon liquid offered by Minnesota Mining and Manufacturing Company (3M), liquid chlorodifluoromethane, etc. In other variations, gases (such as nitrous oxide, nitrogen, carbon dioxide, etc.) may also be utilized as the cooling fluid. In yet another variation, a combination of liquids and/or gases, including, for example, those mentioned above, may be utilized as the coolant fluid “F”.

FIGS. 4 through 7 show the medical device 10 positioned for delivery of electromagnetic energy into tissue “T” shown with the pointing device 45 positioned in varied indicative orientations and shown with diagrammatic representations of radiation patterns of electromagnetic energy delivered into tissue by the medical device 10 responsive to the indicative orientations of the pointing device 45. It is to be understood that the indicative orientations of the pointing device 45 and the radiation patterns of electromagnetic energy are provided for illustrative purposes only, and that medical device 10 embodiments of the present disclosure may be utilized with many different indicative orientations of the pointing device 45 and many different radiation patterns.

FIG. 4 shows the tissue-contact surface 14 of the medical device 10 disposed adjacent to tissue “T” during a procedure, e.g., an ablation procedure, wherein the pointing device 45 is positioned in a first indicative orientation “I₁”. For example, the first indicative orientation “I₁” may correlate with a 30° beam angle and an intensity level “1”, e.g., low-intensity level. FIG. 4 shows a diagrammatic representation of a radiation pattern “P₁” of electromagnetic energy delivered into tissue “T” by the medical device 10 responsive to the first indicative orientation “I₁” of the pointing device 45 in accordance with an embodiment of the present disclosure.

FIG. 5 shows the tissue-contact surface 14 of the medical device 10 disposed adjacent to tissue “T” during a procedure wherein the pointing device 45 is positioned in a second indicative orientation “I₂”. For example, the second indicative orientation “I₂” may correlate with a 0° beam angle and an intensity level “2”, e.g., medium-intensity level. FIG. 5 shows a diagrammatic representation of a radiation pattern “P₂” of electromagnetic energy delivered into tissue “T” by

the medical device responsive to the second indicative orientation “I₂” of the pointing device 45 in accordance with an embodiment of the present disclosure.

FIG. 6 shows the tissue-contact surface 14 of the medical device 10 disposed adjacent to tissue “T” during a procedure wherein the pointing device 45 is positioned in a third indicative orientation “I₃”. For example, the third indicative orientation “I₃” may correlate with a 0° beam angle and an intensity level “3”, e.g., high-intensity level. FIG. 6 shows a diagrammatic representation of a radiation pattern “P₃” of electromagnetic energy delivered into tissue “T” by the medical device responsive to the third indicative orientation “I₃” of the pointing device 45 in accordance with an embodiment of the present disclosure.

FIG. 7 shows the tissue-contact surface 14 of the medical device 10 disposed adjacent to tissue “T” during a procedure wherein the pointing device 45 is positioned in a fourth indicative orientation “I₄”. For example, the fourth indicative orientation “I₄” may correlate with a -30° beam angle and an intensity level “1”, e.g., low-intensity level. FIG. 7 shows a diagrammatic representation of a radiation pattern “P₄” of electromagnetic energy delivered into tissue “T” by the medical device 10 responsive to the fourth indicative orientation “I₄” of the pointing device 45 in accordance with an embodiment of the present disclosure.

FIG. 8 is a schematic diagram of an embodiment of a control system 800 according to the present disclosure that is communicatively coupled with an on/off button 810 and configured to utilize a joystick position signal 820 indicative of intensity and angle of beam. As schematically-illustrated in FIG. 8, the control system 800 utilizes the joystick position signal 820 to determine whether to adjust antenna and/or amplifier gain 830 and/or to determine the phasing of the radiating elements (1 through N) of a phased antenna array 861, e.g., to allow the focal point of energy delivery in tissue to be varied in position in real-time and/or near real-time.

Control system 800 is configured such that when the on/off button 810 is in the “ON” state, adjustment of antenna and/or amplifier gain 830 is permitted, and when the on/off button 810 is in the “OFF” state, adjustment of antenna and/or amplifier gain 830 is not permitted. Joystick position signal 820 may be used in conjunction with a lookup table 840 to enable selective steering of the radiated beam of the phased antenna array 861. Lookup table 840 includes data representative of a mapping of the joystick positions to the phasing of the phased antenna array 861. As schematically-illustrated in FIG. 8, the control system 800 utilizes the lookup table 840 to determine the phasing of the radiating elements (1 through N) of the phased antenna array 861.

The above-described energy-delivery devices including an ultrasound transducer array and a phased antenna array are capable of directing energy into tissue, and may be suitable for use in a variety of procedures and operations. The presently-disclosed energy-delivery device including an ultrasound transducer array and a phased antenna array may be implemented using electromagnetic radiation at microwave frequencies, RF frequencies or at other frequencies.

The above-described energy-delivery device including an ultrasound transducer array and a phased antenna array according to embodiments of the present disclosure are adapted to be hand-holdable and include an ergonomically located user-interface.

The above-described electrosurgical systems provide clinicians the ability to visualize a tissue region during energy delivery into the tissue region. In the above-described electrosurgical systems, data acquired by the ultrasound transducer array may be outputted from the above-described

energy-delivery device to an ultrasound imaging system, and may be outputted from the imaging system to one or more display devices and/or screens, which may be used by the clinician to visualize the targeted region in real-time and/or near real-time.

Although embodiments have been described in detail with reference to the accompanying drawings for the purpose of illustration and description, it is to be understood that the inventive processes and apparatus are not to be construed as limited thereby. It will be apparent to those of ordinary skill in the art that various modifications to the foregoing embodiments may be made without departing from the scope of the disclosure.

What is claimed is:

1. A medical probe device suitable for delivery of energy to tissue, comprising:
 - a housing;
 - a phased antenna array disposed within the housing, the phased antenna array emitting therapeutic electromagnetic energy;
 - a user-interface disposed on the housing, the user-interface adapted to enable a user to selectively adjust the radiation pattern of electromagnetic energy delivered into a tissue region by the phased antenna array; and
 - an ultrasound transducer array disposed within the housing, the ultrasound transducer array configured to acquire data representative of the tissue region during energy delivery into the tissue region by the phased antenna array.
2. The medical device of claim 1, wherein the housing includes:
 - a body member defining an internal chamber configured to contain the ultrasound transducer device and the phased antenna array therein.
3. The medical device of claim 2, wherein the phased antenna array is operably coupled to an electrosurgical power generating source.
4. The medical device of claim 2, wherein the ultrasound transducer array is communicatively coupled with an ultrasound imaging system.
5. The medical device of claim 2, wherein the body member includes a distal end and defines a tissue-contact surface at the distal end.
6. The medical device of claim 5, wherein the tissue-contact surface includes a first region defining an ultrasound transmissive window.
7. The medical device of claim 6, wherein the ultrasound transducer array is operatively associated with the ultrasound transmissive window.

8. The medical device of claim 6, wherein the tissue-contact surface further includes a second region defining a microwave transmissive window.

9. The medical device of claim 8, wherein the phased antenna array is operatively associated with the microwave transmissive window.

10. The medical device of claim 1, wherein user-interface includes indicia thereon representative of at least one user-selectable parameter of electromagnetic energy delivery into tissue by the medical device.

11. The medical device of claim 10, wherein the indicia includes a first scale including indicia graduation marks and angle in degrees.

12. The medical device of claim 11, wherein the indicia further includes a second scale including indicia graduation marks and a series of consecutive positive integers indicative of energy intensity levels.

13. The medical device of claim 1, wherein the user-interface includes a pointing device.

14. The medical device of claim 13, wherein at least one parameter of electromagnetic energy delivery into tissue by the medical device is correlated to indicative orientations of the pointing device.

15. The medical device of claim 13, wherein the phased antenna array includes a plurality of radiating elements.

16. The medical device of claim 15, wherein at least one electrical signal outputted from the user-interface is responsive to a user-effected movement of the pointing device, and wherein the at least one electrical signal is used to determine phasing of the plurality of radiating elements.

17. A system, comprising:
 an electrosurgical power generating source; and
 a hand-holdable device operably associated with the electrosurgical power generating source, the hand-holdable device including:
 a phased antenna array emitting therapeutic electromagnetic energy;
 a user-interface disposed on the hand-holder device and adapted to enable a user to selectively adjust the radiation pattern of electromagnetic energy delivered into a tissue region by the phased antenna array; and
 an ultrasound transducer array configured to acquire data representative of the tissue region during energy delivery into the tissue region by the phased antenna array.

18. The system of claim 17, wherein the hand-holdable device further includes an electromagnetic window.

19. The system of claim 18, further comprising:
 a coolant supply system in fluid communication with the electromagnetic window.

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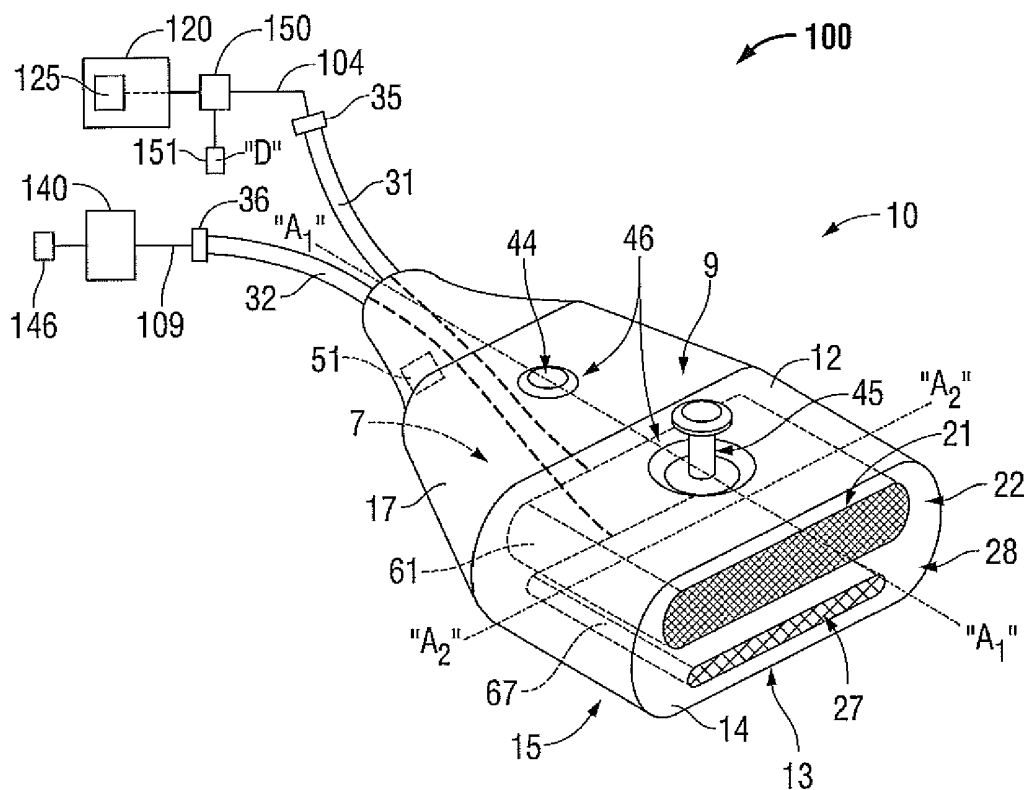


FIG. 1

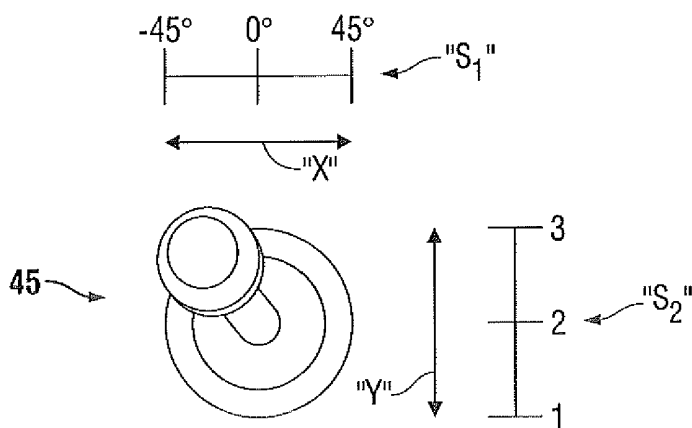


FIG. 2

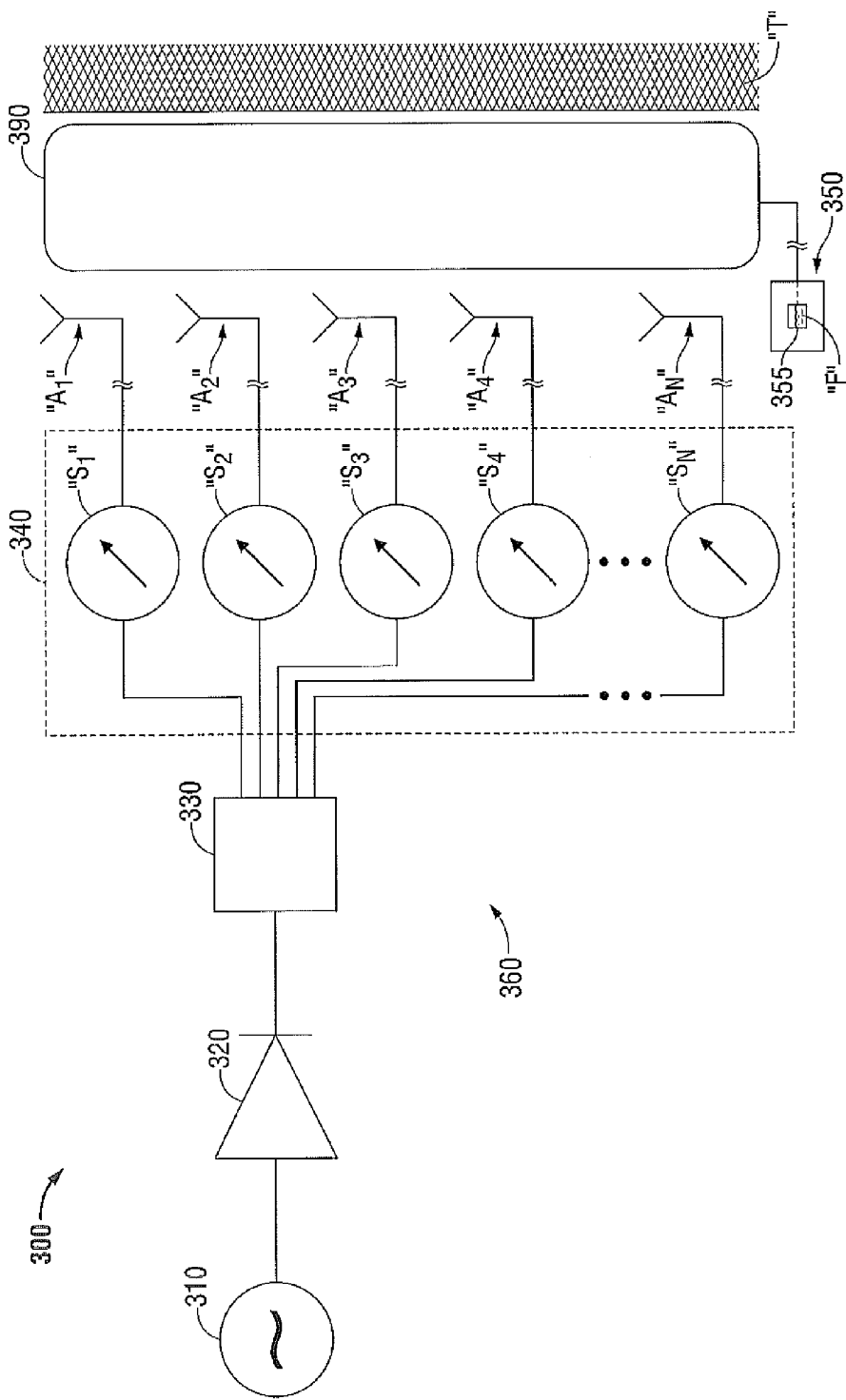


FIG. 3

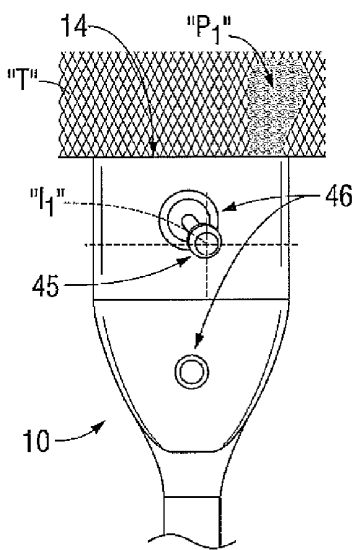


FIG. 4

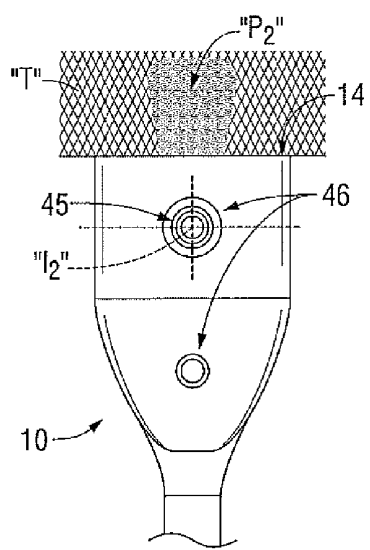


FIG. 5

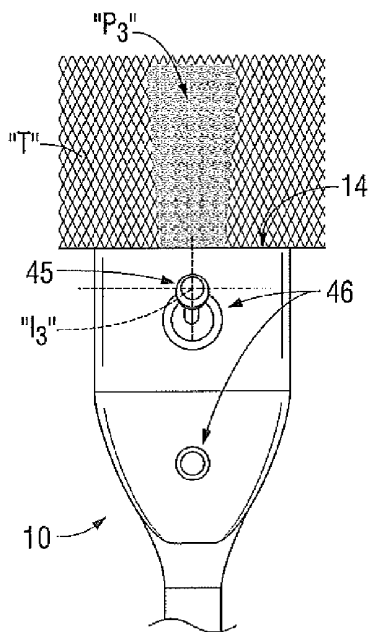


FIG. 6

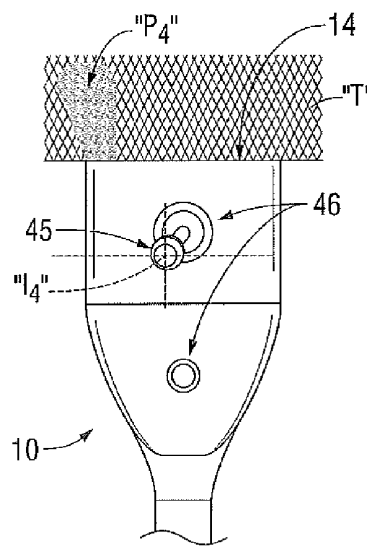


FIG. 7

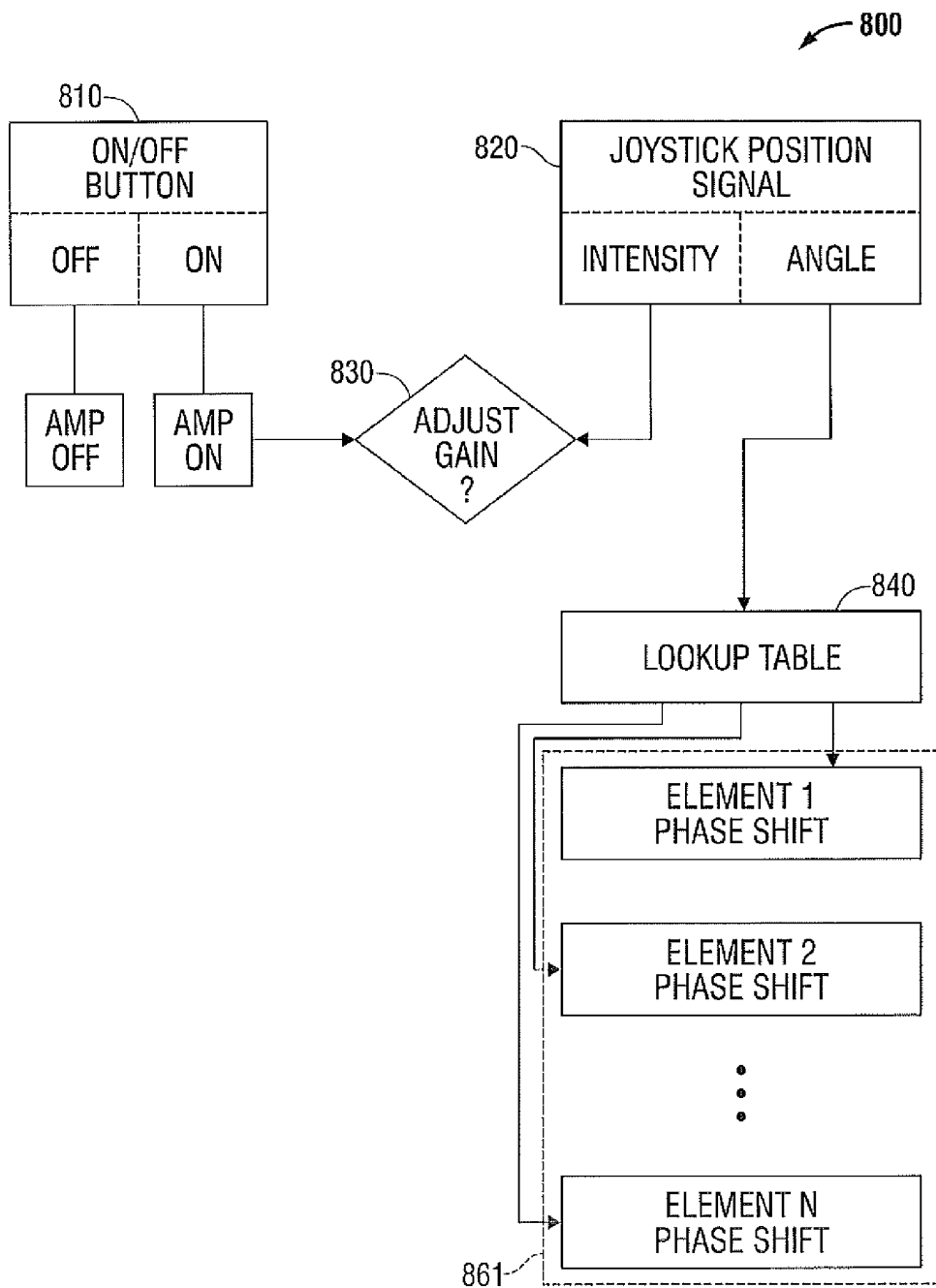


FIG. 8

专利名称(译)	能量输送装置包括超声换能器阵列和相控天线阵列		
公开(公告)号	US8376948	公开(公告)日	2013-02-19
申请号	US13/029521	申请日	2011-02-17
[标]申请(专利权)人(译)	VIVANT医疗		
申请(专利权)人(译)	VIVANT MEDICAL , INC.		
当前申请(专利权)人(译)	VIVANT MEDICAL , INC.		
[标]发明人	BRANNAN JOSEPH D		
发明人	BRANNAN, JOSEPH D.		
IPC分类号	A61B8/00		
CPC分类号	A61B18/1815 A61B8/44 A61B8/00 A61B2018/00023 A61B2018/00577 A61B2018/1846 A61B2019/5276 A61B2018/1838 A61B8/4416 A61B8/4444 A61B2090/378		
其他公开文献	US20120215103A1		
外部链接	Espacenet	USPTO	

摘要(译)

适于向组织输送能量的医疗装置包括壳体，设置在壳体内部的相控天线阵列，以及连接到壳体的用户界面。用户界面适于使用户能够选择性地调整由相控天线阵列传递到组织区域中的电磁能量的辐射图案。医疗设备还包括设置在壳体内部的超声换能器阵列。超声换能器阵列被配置为在通过相控天线阵列将能量输送到组织区域期间获取表示组织区域的数据。

