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[Continued on next page]

- (54) **Title:** DETERMINING ONSETS AND OFFSETS OF CARDIAC DEPOLARIZATION AND REPOLARIZATION WAVES

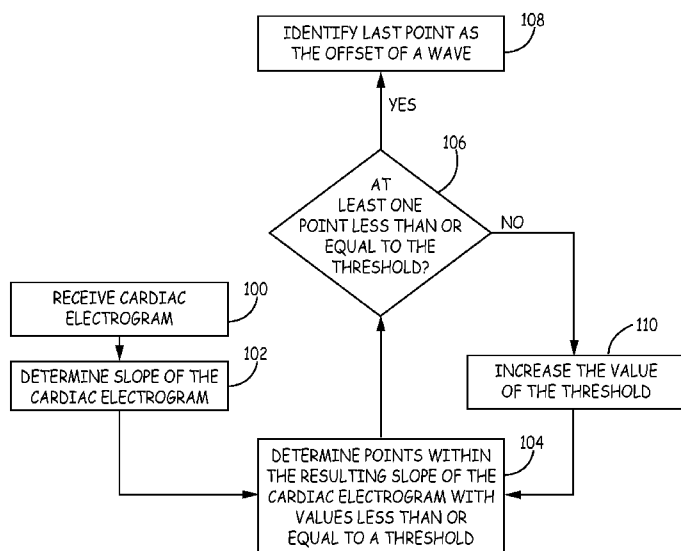


FIG. 3A

(57) **Abstract:** A method for detecting the onsets or offsets of heart depolarization or repolarization waves (e.g. QRS-complex or T-wave) includes determining a slope of a cardiac electrogram. A threshold value is determined based on a maximum of the slope of the cardiac electrogram, and a last point of the cardiac electrogram before the slope of the cardiac electrogram crosses the threshold is identified as one of an onset or an offset of a wave. The method may further include receiving an indication of local ventricular motion associated with a cardiac contraction, and determining an electromechanical delay between the identified onset and the local ventricular motion. Some examples include providing the electromechanical delay for configuration of cardiac resynchronization therapy (CRT), e.g. selection of pacing sites.



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DETERMINING ONSETS AND OFFSETS OF CARDIAC DEPOLARIZATION AND REPOLARIZATION WAVES

TECHNICAL FIELD

5 The invention relates to medical devices, and more particular, to medical devices for sensing, detection, and analysis of cardiac electrograms and electrocardiograms.

BACKGROUND

10 Implantable medical devices (IMDs), such as implantable pacemakers, cardioverters, defibrillators, or pacemaker-cardioverter-defibrillators, provide therapeutic electrical stimulation to the heart. IMDs may provide pacing to address bradycardia, or pacing or shocks in order to terminate tachyarrhythmia, such as tachycardia or fibrillation. In some cases, the medical device may sense intrinsic depolarizations of the heart, detect arrhythmia based on the intrinsic depolarizations (or absence thereof), and control delivery
15 of electrical stimulation to the heart if arrhythmia is detected based on the intrinsic depolarizations.

 IMDs may also provide cardiac resynchronization therapy (CRT), which is a form of pacing. CRT involves the delivery of pacing to the left ventricle, or both the left and right ventricles. The timing and location of the delivery of pacing pulses to the
20 ventricle(s) may be selected to improve the coordination and efficiency of ventricular contraction.

 IMDs sense signals and deliver therapeutic stimulation via electrodes. Implantable pacemakers, cardioverters, defibrillators, or pacemaker-cardioverter-defibrillators are typically coupled to one or more subcutaneous electrodes or intracardiac leads that carry
25 electrodes for cardiac sensing and delivery of therapeutic stimulation. The signals sensed via the electrodes may be referred to as a cardiac electrogram (EGM) and may include the depolarizations, repolarizations, and other intrinsic electrical activity of the heart.

 Systems for implanting medical devices may include workstations or other equipment in addition to the medical device itself. In some cases, these other pieces of
30 equipment assist the physician or other technician with placing the intracardiac leads at particular locations on the heart. In some cases, the equipment provides information to the

physician about the electrical activity of the heart and the location of the intracardiac lead. The equipment may perform similar functions as the medical device, including delivering electrical stimulation to the heart and sensing the depolarizations of the heart. In some cases, the equipment may include equipment for obtaining an electrocardiogram (ECG) via electrodes on the surface of the patient.

SUMMARY

In general, the disclosure describes techniques for determining when an electrical wave within a cardiac electrogram begins or ends. More particularly, the techniques for detecting the onsets and offsets of heart depolarization waves, such as the QRS complex and P-wave, and heart repolarization waves, such as the T-wave, are described. A variety of diagnostic metrics may be determined based on the determined onsets and/or offsets. As examples, QRS width, Q-T interval, or electromechanical delay, e.g., of ventricular contraction relative to the onset of the Q-wave, may be determined based on the determined onsets and/or offsets.

In some examples, the diagnostic metrics may be used to optimize or otherwise guide the configuration of therapy, such as CRT. For CRT, lead placement, pacing electrode configuration, or various atrio-ventricular or interventricular intervals may be configured based on metrics that are determined based on the identified onsets and/or offsets. In some examples, electromechanical delay may be used to configure CRT and, particularly, to select a lead placement, e.g. left-ventricular lead placement, during implantation of a CRT system.

In one example, a method includes determining a slope of a cardiac electrogram, determining a threshold value based on a maximum of the slope of the cardiac electrogram, and identifying a last point of the cardiac electrogram before the slope of the cardiac electrogram crosses the threshold as one of an onset or an offset of a wave.

In another example, a system includes a slope module that determines the slope of a cardiac electrogram and a wave detector module that determines a threshold value based on a maximum of the slope of the cardiac electrogram and identifies a last point of the cardiac electrogram before the slope of the cardiac electrogram crosses the threshold as one of an onset or and offset of a wave.

In another example, a system comprises means for determining a slope of a cardiac electrogram, means for determining a threshold value based on the maximum of the slope of the cardiac electrogram, and means for identifying a last point of the cardiac electrogram before the slope of the cardiac electrogram crosses the threshold as one of an onset or an offset of a wave.

In another example, a computer readable medium stores a set of instructions which, when implemented in a medical device system causes the system to determine a slope of a cardiac electrogram, determine a threshold value based on the maximum of the slope of the cardiac electrogram, and identify a last point of the cardiac electrogram before the slope of the cardiac electrogram crosses the threshold as one of an onset or an offset of a wave.

BRIEF DESCRIPTION OF DRAWINGS

FIG. 1 is a block diagram illustrating an example system that may determine the onsets and offsets of various heart repolarization and depolarization waves.

FIG. 2 is a block diagram illustrating an example configuration of a wave detection module.

FIG. 3A is a flow diagram illustrating an example technique for determining the onset of a wave.

FIG. 3B is a flow diagram illustrating an example technique for determining the offset of a wave.

FIG. 4A is a flow diagram illustrating an example technique for determining the onset of a wave.

FIG. 4B is a flow diagram illustrating an example technique for determining the offset of a wave.

FIG. 5 is a graph illustrating an example cardiac electrogram.

FIG. 6 is a graph illustrating an example cardiac electrogram including R-wave marker information.

FIG. 7 is a graph illustrating an example cardiac electrogram including R-wave marker information and example windows around each marker.

FIG. 8 is a graph illustrating an example cardiac electrogram including example determined wave onset points.

FIG. 9 is a graph illustrating both a cardiac contraction curve and a surface ECG curve with indicators marking the determined onset of depolarization of the heart and the detected timing of local heart mechanical contraction.

FIG. 10 is a graph illustrating both a cardiac contraction curve at a potential intracardiac lead implant site and a local EGM curve from the same potential implant site with indicators marking the determined onset of local depolarization of the heart and the detected timing of local heart mechanical contraction.

FIG. 11 is a flow diagram illustrating an example technique for selecting intracardiac lead implant sites.

FIG. 12 is a flow diagram illustrating an example technique for selecting intracardiac lead implant site.

FIG. 13 is a conceptual diagram illustrating an example system that determines the onset and offsets of heart depolarization and repolarization waves.

FIG. 14 is a conceptual diagram illustrating the implantable medical device (IMD) and leads of the system shown in FIG. 13 in greater detail.

FIG. 15 is a block diagram of an example implantable medical device that may determine the onset and offset of heart depolarization and repolarization waves.

FIG. 16 illustrates an example of the time-delay between the delivery of pacing and the onset of depolarization.

DETAILED DESCRIPTION

The techniques described in this disclosure allow a system to determine the onsets and offsets of heart repolarization and depolarization waves. A variety of diagnostic metrics may be determined based on the determined onsets and/or offsets. As examples, QRS width, Q-T interval, or electromechanical delay, e.g., of ventricular contraction relative to the onset of the Q-wave, may be determined based on the determined onsets and/or offsets. In some examples, the diagnostic metrics may be used to optimize or otherwise guide the configuration of therapy, such as CRT. For CRT, lead placement, pacing electrode configuration, or various atrio-ventricular or interventricular intervals

may be configured based on metrics that are determined based on the identified onsets and/or offsets. In some examples, electromechanical delay may be used to configure CRT and, particularly, to select a lead placement, e.g. left-ventricular lead placement, during implantation of a CRT system

5 In general, a heart produces a repetitive electrical signal which causes the heart to mechanically contract, thereby pumping blood throughout the body. Generally, the signal may be detected and displayed as a cardiac electrogram signal. Although the exact representation may differ depending on the placement of leads on or within the body to detect the heart signal, among other factors, a common cardiac electrogram includes
10 several recognizable features. The initial deflections of the signal represent the P-wave and the QRS complex. The P-wave represents the depolarization of the atria and the QRS complex represents the depolarization of the ventricles. The Q-wave of the QRS complex is the initial downward deflection of the signal during the complex. Following the Q-wave is the R-wave, which is an upward deflection of the signal. Finally, the S-wave is another
15 downward deflection. The next portion of the signal represents repolarization of the atria and ventricles. More specifically, what is generally called the T-wave represents the repolarization of the ventricles. There is no specific wave or feature of the signal that represents the repolarization of the atria because the generated signal is small in comparison to the T-wave. Together, the P-wave, the Q, R, and S waves, and the T-wave
20 represent the depolarization and repolarization waves of a heart electrical signal.

The described techniques may enhance the accuracy of determining the onsets and offsets of the various waves which comprise the repeated cardiac electrogram signal. In particular, knowing more accurately the timing of the onsets and offsets of the waves allows for a more accurate determination of the electrical-electrical delays and the
25 electromechanical delays. For example, the point of onset of ventricular depolarization on surface ECG forms a fiducial with respect to which local electrical activation or depolarization times and may be measured at different sites in the ventricle. During
30 implant of a heart lead for cardiac resynchronization therapy, the time-interval between this onset point and the time of local activation or depolarization at a candidate implant site within the ventricle may be evaluated and if it exceeds a certain threshold (e.g. 90 ms), that site may be selected for implant.

FIG. 1 is a block diagram illustrating an example configuration of a system for determining the onsets and offsets of heart depolarization and repolarizations waves. In the illustrated example, system 10 includes a device 60, a cardiac electrogram module 70, and a motion sensing module 80. Device 60 may further include a processor 72, a
5 memory 74, a peak detection module 76, and a wave detection module 78.

Processor 72 may include any one or more of a microprocessor, a controller, a digital signal processor (DSP), an application specific integrated circuit (ASIC), a field-programmable gate array (FPGA), or equivalent discrete or analog logic circuitry. In some examples, processor 72 may include multiple components, such as any combination
10 of one or more microprocessors, one or more controllers, one or more DSPs, one or more ASICs, or one or more FPGAs, as well as other discrete or integrated logic circuitry. The functions attributed to processor 72 herein may be embodied as software, firmware, hardware or any combination thereof. Generally, processor 72 controls cardiac
15 electrogram module 70, peak detection module 76, wave detection module 78, and motion sensing module 80 to determine timings of the onsets and offsets of heart depolarization and repolarizations waves.

Memory 74 includes computer-readable instructions that, when executed by processor 72, causes system 10 and processor 72 to perform various functions attributed to system 10 and processor 72 herein. Memory 74 may include any volatile, non-volatile,
20 magnetic, optical, or electrical media, such as a random access memory (RAM), read-only memory (ROM), non-volatile RAM (NVRAM), electrically-erasable programmable ROM (EEPROM), flash memory, or any other digital or analog media.

Generally, cardiac electrogram module 70 is configured sense electrical signals from a patient. Cardiac electrogram module 70 is electrically coupled to one or more
25 electrodes 2, 4...n by one or more leads. In some examples, the one or more electrodes 2, 4...n may be external electrodes, e.g., attached to the surface of a patient, or implanted at various locations within a patient, e.g., on or within a heart.

Peak detection module 76 may be configured to determine a maximum value of a particular signal. For example, peak detection module 76 may be configured to receive
30 the electrical signal from cardiac electrogram module 70 and determine the maximum value. In another example, as illustrated in FIG. 2, peak detection module 76 may be

configured to receive a signal from wave detection module 78 and determine a maximum value.

Wave detection module 78 determines the onsets and offsets on the heart depolarization and repolarizations waves. Wave detection module 78 may be configured to receive an electrical signal. For example, wave detection module 78 may be configured to receive an electrical signal sensed by cardiac electrogram module 70. Motion sensing module 80 may detect mechanical motion of a heart, e.g., during contraction of the heart. Motion sensing module 80 may comprise one or more sensors that generate a signal that varies based on cardiac contraction or motion generally, such as one or more accelerometers, pressure sensors, impedance sensors, or flow sensors. Motion sensing module 80 may provide an indication of the timing of motion, e.g., contraction, to device 60, e.g., to processor 72. The detected contraction may be contraction of cardiac tissue at a particular location, e.g., a particular portion of a ventricular wall.

In some examples, motion sensing module 80 may be configured to image the heart, or electrodes, catheters, wires, or other radio-opaque markers in or on the heart and identify motion associated with contraction based on images of the heart. In some examples, motion sensing module 80 may be configured to direct ultrasound energy toward a patient's heart. Motion sensing module 80 may also be configured to detect any ultrasonic energy deflected back toward motion sensing module 80 by the patient's heart. In this manner, motion sensing module 80 may capture information about the mechanical motion (i.e. the contracting and relaxing of the ventricles and/or atria) of the heart. Systems and methods for identifying heart mechanical contractions are described in U.S. Patent No. 7,587,074 to Zarkh et al., which issued on Sept. 8, 2009 and is entitled, "METHOD AND SYSTEM FOR IDENTIFYING OPTIMAL IMAGE WITHIN A SERIES OF IMAGES THAT DEPICT A MOVING ORGAN,".

Processor 72 may determine values of one or more metrics, such as cardiac intervals or cardiac electromechanical delay, based on the timing of onset and/or offset of a wave, as determined by wave detection module 78, and/or the timing of contraction, as determined by motion sensing module 80. For example, processor 72 may determine QRS width based on an onset and offset of the QRS complex as identified by wave detection module 78. As another example, processor 72 may determine a QT interval based on a QRS onset and T-wave onset identified by wave detection module 78. The processor may

also determine the interval between the onset of depolarization on a surface ECG lead and the time of local electrical activation as sensed by a lead or a mapping catheter or guidewire at a site within the heart. Furthermore, as described in greater detail below, processor 72 may determine electromechanical delay based on a QRS onset identified by wave detection module 78 and an indication of the timing of cardiac contraction received from motion sensing module 80.

Although in FIG. 1 device 60, module 70, and module 80 are depicted as separate, in other examples the modules and device may be combined into fewer separate components. For example, as illustrated in FIG. 15, all of the functionality of system 10 may be combined into a single device.

Furthermore, although processor 72, peak detection module 76 and wave detection module 78 are depicted as separate functional modules in the example of FIG. 1, their collective functionality may be provided by any number of physical or logical processing elements provided by one or more co-located or networked devices. In one example, peak detection module 76 and wave detection module 78 may be functional modules executed by processor 72. Similarly in FIG. 2, although the various modules 90, 92, 94, 96, 98, and 99 are depicted as separate modules in a single device, in other examples their functionality may be provided by any one or more devices.

FIG. 2 is a block diagram illustrating an example configuration of wave detection module 78. In the example of FIG. 2, wave detection module 78 comprises a low-pass filter 90, a window module 92, a slope module 94, a rectifier module 96, a smoothing module 98, and a threshold detection module 99.

Low pass filter 90 may generally be any low-pass filter designed to reduce or eliminate the high frequency components of electrical signals. Some examples of low-pass filters embodied in hardware include capacitive low-pass filters and inductive low-pass filters. Other low-pass filters may be embodied entirely within software. In some embodiments, low-pass filter 90 is embodied as a combination of hardware and software. Low-pass filter 90 may be a first, second, or higher order filter. In some examples, low-pass filter 90 comprises multiple filters placed in a succession in order to create a desired frequency response. In some examples, the low-pass filter 90 is a linear filter with a maximally flat group delay or maximally linear phase response. A constant group delay is a characteristic of phase response of an analog or a digital filter, which helps preserve the

shape of the signal in the pass-band. In at least one example, the low-pass filter is a Bessel filter with a cut-off frequency of 15Hz.

Window module 92 may generally window received signals. In some examples, window module 92 may receive cardiac electrograms, and in further examples, some of the cardiac electrograms may include a marker or markers indicating one or more points of interest. Some example of points of interest could be the R-wave, the P-wave, or any other wave of the cardiac electrogram. In some examples, peak detection module 76 may detect the locations of R-waves in a cardiac electrogram, according to techniques that are well known in the art, for example using a varying threshold. Peak detection module 76 may then place a marker within the cardiac electrogram identifying the location of the R-wave. In other examples, other modules or devices may detect and mark waves in the cardiac electrogram. In examples where the signal includes at least one marker, window module 92 may window the received electrical signal around the marker. For example, window module 92 may multiply the received electrical signal by zero outside of the area around the marker and by one inside of the area around the marker. In this way, window module 92 may modify the received electrical signal to only contain information in an area around the marker. In some examples, window module 92 may isolate the QRS complex. In some examples, window module 92 may window an area of interest of equal length around the marker, for instance 150ms before the marker and 150ms after the marker. In other examples, the area in front of the marker may be longer or shorter than the area behind the marker.

Slope module 94 may generally determine the slope of a received electrical signal. Some example techniques for determining a slope that slope module 94 may use are taking the simple difference between adjacent points on the received electrical signal, or determining the first derivative of the received electrical signal.

Rectifier module 96 may generally rectify a received electrical signal. For example, rectifier module 96 may half-wave rectify the received electrical signal to produce a resulting signal with information only where the original signal was above zero. In another example, rectifier module 96 may full-wave rectify the received electrical signal to produce a resulting signal where all the negative values of the original signal are now positive.

Smoothing module 98 may generally smooth a received electrical signal. For example, smoothing module 98 may be configured to increase the values of certain points and decrease the values of certain points so as to create a smoother signal. Some example smoothing algorithms include rectangular or un-weighted sliding average smoothing, triangular smoothing, and Savitzky-Golay smoothing. The smoothing algorithms may be implemented through one or more filters. In at least one example, the smoothing filter is 10-order median filter. In another example, the smoothing filter is a n-order median filter where n is an increasing linear function of the sampling frequency used to digitize the electrogram or electrocardiogram signals.

Threshold detection module 99 may generally be configured to determine a threshold and at what points a received electrical signal crosses a pre-determined threshold. The threshold may be determined based on a maximum value of the signal received from smoothing module 98. For example, as illustrated in FIG. 2, threshold detection module 99 may be configured to receive, from peak detection module 76, a maximum value of the signal received from smoothing module 98. Threshold detection module 99 may be configured to determine a threshold value based on the received maximum value. For example, threshold detection module 99 may determine the threshold to be ten percent, fifteen percent, twenty percent, or more of the maximum value. Ultimately, threshold detection module 98 may determine the onsets and offsets of heart depolarization and repolarizations waves based on at which points of a received electrical signal cross a threshold.

FIGS. 3A and 3B are flow diagrams illustrating example techniques for determining the onsets and offsets, respectively, of heart depolarization and repolarizations waves. The example techniques are described with respect to system 10. However, in other examples, the techniques may be practiced by a single device, such as the device illustrated in FIG. 12.

According to the illustrated example in FIG. 3A, wave detection module 78 receives a cardiac electrogram (100). In the example, wave detection module 78 may then determine a slope of the cardiac electrogram (102). Wave detection module 78 may determine a number of points within the resulting slope of the cardiac electrogram with values less than or equal to a threshold value (104).

As described above, the threshold value may be a percentage of the maximum value of the slope of the cardiac electrogram, with some examples including ten percent, fifteen percent, and twenty percent. In one example, the threshold value begins at ten percent of the maximum value of the resulting slope of the cardiac electrogram. Wave detection module 78 then determines if there is at least one one point that is less than or equal to the threshold value (106). If there is at least one point that is less than or equal to the threshold value (YES of 106), then wave detection module 78 determines that a last point before the slope of the cardiac electrogram crosses above the threshold value is an onset of a wave (108). If wave detection module 78 determines that there is not at least one point with a value less than or equal to the threshold value (NO of 106), then wave detection module 78 may increase the threshold value and return to step 104. For example, wave detection module 78 may increase the threshold value, e.g., to be fifteen percent of the maximum value of the slope of the cardiac electrogram. This process may repeat until wave detection module 78 finds at least one point with a value less than or equal to a threshold value.

According to the illustrated example in FIG. 3B, wave detection module 78 receives a cardiac electrogram (200). In the example, wave detection module 78 may then determine a slope of the cardiac electrogram (202). Wave detection module 78 may determine a number of points within the resulting slope of the cardiac electrogram with values greater than or equal to a threshold value (204). Wave detection module 78 may also determine that the last point before the slope of the cardiac electrogram crosses below the threshold value is the offset of a wave (208).

FIGS. 4A and 4B flow diagrams illustrating example techniques for determining the onsets and offsets of heart depolarization and repolarizations waves in greater detail. As with FIGS. 3A and 3B, the example techniques are described with respect to system 10. However, in other examples, the techniques may be practiced by a single device, such as the device illustrated in FIG. 15, or any device or combination of devices described herein.

According to the example illustrated in FIG. 4A, the wave detection module 78 receives a cardiac electrogram (300). Wave detection module 78 may then receive a wave marker. As described above, a wave marker may be a piece of information identifying a point of interest on the cardiac electrogram. For example, the wave marker might identify,

e.g., the R-wave, P-wave, or T-wave of the cardiac electrogram. In the example illustrated in FIG. 4A, once wave detection module 78 has received the cardiac electrogram and a wave marker, low-pass filter module 90 may low-pass filter the cardiac electrogram (304). As described previously, the low-pass filter may reduce or eliminate the high frequency components of the cardiac electrogram while preserving the shape of the waveform in the pass-band. Window module 92 may window the cardiac electrogram around the marker (306). In at least one example, the marker may identify an R-wave and the modules from FIG. 2 may window the cardiac electrogram around the marker to isolate a QRS complex. Slope module 94 may determine the slope of the windowed signal (308). In the example technique, rectifier module 96 may rectify the slope of the windowed signal (310). After the rectification of the signal, smoothing module 98, in the example technique, may smooth the signal (312).

After smoothing, wave detection module 78 may pass the signal to peak detection module 76, which may then determine the maximum value of the resulting slope signal (314). After the maximum value of the resulting slope signal is determined, threshold detection module 99 may determine a threshold value. For example, threshold detection module 99 may determine that the threshold value is ten percent of the maximum value of the resulting slope signal. Threshold detection module 99 may then determine if there is at least one point within the first portion of the resulting slope signal that is less than or equal to the threshold value of ten percent (316). In one example, the first portion of the resulting slope signal is the portion that comes before the wave marker. In another example, the first portion is the portion of the resulting slope signal that comes before the maximum value of the resulting slope signal. In still another example, the first portion is the portion of the resulting slope signal prior to a peak of the cardiac electrogram.

In the example technique, if threshold detection module 99 determines that there is at least one point in the first portion of the resulting slope signal that is less than or equal to the threshold value of ten percent (YES of 316), then the module determines that the last point before the resulting slope signal crosses above the threshold value is the onset of a wave. If threshold detection module 99 determines that there is not at least one point that is less than or equal to the threshold value of ten percent (NO of 316), then threshold detection module 99 may increase the threshold value. For example, threshold detection

module 99 may increase the threshold value to be fifteen percent of the maximum value of the resulting slope signal.

Then, threshold detection module 99 may determine if there are any points in the first portion of the resulting slope signal that are less than or equal to the threshold value of fifteen percent (318). In the example technique, if there is at least one point that is less than or equal to the threshold value of fifteen percent (YES of 318), then threshold detection module 99 determines that the last point before the resulting slope signal crosses above the threshold value is the onset of a wave. If threshold detection module 99 determines that there is not at least one point that less than or equal to the threshold value of fifteen percent (NO of 318), then threshold detection module 99 may increase the threshold value. For example, threshold detection module 99 may increase the threshold value to be twenty percent of the maximum value of the resulting slope signal.

Then, threshold detection module 99 may determine if there are any points in the first portion of the resulting slope signal that are less than or equal to the threshold value of twenty percent (320). In the example technique, if there is at least one point that is less than or equal to the threshold value of twenty percent (YES of 320), then threshold detection module 99 determines that the last point before the resulting slope signal crosses above the threshold value is the onset of a wave. If threshold detection module 99 determines that there is not at least one point that is less than or equal to the threshold value of twenty percent (NO of 320), then wave detection module 78 may abandon finding an onset point of a wave and return to step 300 to receive a new cardiac electrogram.

According to the example illustrated in FIG. 4B, wave detection module 78 receives a cardiac electrogram (400). At step 302, wave detection module 78 may receive a wave marker. In the example technique, once wave detection module 78 has received the cardiac electrogram and a wave marker, the low-pass filter module 90 may low-pass filter the cardiac electrogram (404). Window module 92 may also window the cardiac electrogram around the marker (406). In at least one example, the marker may identify an R-wave and the window module 92 may window the cardiac electrogram around the marker to isolate a QRS complex. Slope module 94 may further determine the slope of the windowed signal (408). In the example technique, rectifier module 96 may also rectify the slope of the windowed signal (410). After rectifying the signal, smoothing module 98, in the example technique, may smooth the signal (412).

After smoothing, wave detection module 78 may pass the signal to peak detection module 76. In the example method, peak detection module 76 may then determine the maximum value of the resulting signal (414). After determining the maximum value of the resulting slope signal, in the example technique, threshold detection module 99 may determine a threshold value. For example, threshold detection module 99 may determine that the threshold value is ten percent of the maximum value of the resulting slope signal. In the example technique, threshold detection module 99 may then determine all of the points within the second portion of the resulting slope signal that are greater than or equal to the threshold value of ten percent (416). In one example, the second portion of the resulting slope signal is the portion that comes after the wave marker. In another example, the second portion is the portion of the resulting slope signal that comes after the maximum value of the resulting slope signal. In still another example, the first portion is the portion of the resulting slope signal prior to a peak of the cardiac electrogram. In the example technique, wave detection module 78 determines that the last point before the resulting slope signal crosses below the threshold value is the offset of a wave.

FIGS. 5-8 illustrate example cardiac electrogram signals along with various aspects of the present disclosure. For example, FIG. 5 illustrates an example cardiac electrogram 502 that may be passed to wave detection module 78 at the beginning of the example techniques described in FIGS. 3-4. FIG. 6 illustrates a cardiac electrogram 502 along with wave markers 504. In the example of FIG. 6, wave markers 504 are R-wave markers. FIG. 7 also depicts an example cardiac electrogram 502 and R-wave markers 504. FIG. 7 also depicts example windows 506 that wave detection module 78, e.g. through window module 92, may generate about the wave markers 504. In the example of FIG. 7, windows 506 are configured to generally include the QRS complexes and isolate the QRS complexes from the whole cardiac electrogram, e.g. exclude other waves such as the P-wave and T-wave. In other examples, the windows 506 may be configured to generally include P or T-waves and isolate those waves from the whole cardiac electrogram. FIG. 8 further illustrates an example cardiac electrogram 502. FIG. 8 also illustrates the points that wave detection module 78 has determined are the onset of the waves (more specifically, in the example illustrated in FIG. 8, the onset of the QRS waves).

FIGS. 9 and 10 illustrate example cardiac electrograms combined with example heart mechanical contraction information. For example, FIG. 9 depicts an example

surface ECG signal 702 along with a cardiac contraction curve 704. Cardiac contraction curve 704 illustrates the contraction of a local point on the heart with respect to time. The cardiac contraction curve may be a signal generated by any of the sensors discussed above, for example accelerometers, pressure sensors, impedance sensors, or flow sensors associated with motion sensing module 80. In other examples, the cardiac contraction curve may be generated by the techniques described in the U.S. Patent No. 7,587,074 to Zarkh et al., which issued on Sept. 8, 2009 and is entitled, "METHOD AND SYSTEM FOR IDENTIFYING OPTIMAL IMAGE WITHIN A SERIES OF IMAGES THAT DEPICT A MOVING ORGAN,". In still other examples, the cardiac contraction curve may be generated by sensors within the tip of an intracardiac electrode. For example, the tip of a cardiac electrode may contain one or more motion sensors which generates a signal as the tip of the electrode moves in relation to the region of the heart in which it is implanted.

Also depicted in FIG. 9 are the timing of the detected onset of depolarization of the heart from the surface ECG 706 and the timing of the local cardiac contraction 708. The delay between the onset of depolarization 706 from the surface ECG and the timing of local cardiac contraction 708 may be described as the global electromechanical delay 710. As will be described in greater detail in FIGS. 11-12, measuring the global electromechanical delay 710 delay at various locations on the heart may help physicians to select potential intracardiac lead implant sites.

FIG. 10 depicts an example unipolar cardiac electrogram signal 752 from an electrode at a localized position on or within a heart, along with a cardiac contraction curve 754 from the same localized position on or within the heart. FIG. 10 also depicts the timing of the detected onset of depolarization of the heart at the localized position 756 and the timing of the local cardiac contraction 758. In contrast to FIG. 9, FIG. 10 displays the difference in timing between the local depolarization of the heart 756 and the timing of the local cardiac contraction 758 as the local electromechanical delay 760 or local electromechanical latency 760. The local electromechanical delay 760 differs from the global electromechanical delay 710 in that local electromechanical delay 760 measures the differences in timings both at the localized heart tissue and the global electromechanical delay 710 measures the difference in timings between the depolarization 706 detected at the surface ECG and the local mechanical contraction 708.

FIG. 11 is a flow diagram describing an example method for selecting an implant site for an intracardiac lead. In the example method, a system, such as system 10 of FIG. 1, may determine the intrinsic global electromechanical delay at a plurality of potential intracardiac lead implant sites. To determine the intrinsic global electromechanical delay, a system may determine the timing of a detected onset of depolarization from a surface ECG and a timing of cardiac contraction at various locations on the heart during an intrinsic heart rhythm. The difference between the timings is the global electromechanical delay (800). Then, in the example technique, a system, e.g., processor 72, or a user of the system, may select candidate intracardiac lead implant sites as sites that have a global electromechanical delay above a certain threshold (802). Finding particular candidate sites with a long global electromechanical delay may indicate that the particular location of the heart contracts late and may be the reason for dyssynchrony of cardiac wall motion. Selecting an implant site to provide pacing on or near that site may enhance the effectiveness of the electrical pacing therapy to restore cardiac synchrony.

In the example technique, processor 72 may then determine the local electromechanical delay of the selected candidate implant sites during an intrinsic heart rhythm (804). To determine the local electromechanical delay, processor 72 may determine the timing of a detected onset of depolarization from an electrogram taken at the potential implant site and a timing of the local cardiac contraction at the potential implant sites. The local electromechanical delay is the difference between the determined timings. Then, in the example technique, processor 72 may automatically select a subset of potential implant sites as sites that have a local electromechanical delay less than a threshold (806). In other examples, a user may manually select the potential implants based on information from processor 72. Candidate implant sites that have a long local electromechanical delay may indicate a region of scarred or non-viable or otherwise non-conductive tissue. Implanting an intracardiac lead at those locations may reduce the effectiveness of the electrical stimulation therapy.

Processor 72 may then deliver electrical pacing stimulus to the heart at the potential implant sites (808). Processor 72 may measure various metrics such as global or local electromechanical delay during pacing at each potential implant site. In the example technique, processor 72 may then automatically select one or more implant sites based on the reduction of a metric during pacing compared to the intrinsic baseline. In other

examples, a user may select one or more implant sites based on information from processor 72. For example, processor 72, or a user, may select one or more implant sites based on the reduction of local electromechanical, global electromechanical delay, or cardiac dyssynchrony, which is a measure of the distribution and dispersion (e.g. range, standard deviation, etc.) of global or local electromechanical delays at various cardiac sites. In other example techniques, processor 72 or a user may select other parameters based on the global and/or local electromechanical delay. For example, processor 72 may also vary parameters of the pacing, such as A-V delay, V-V delay, and pacing electrode configuration, and select a particular pacing electrode configuration or pacing intervals for cardiac resynchronization therapy based on the metric. In other examples, a user may ultimately select the pacing electrode configuration or pacing intervals based on the metric.

FIG. 12 is flow diagram illustrating another example technique for selecting an implant site for an intracardiac lead. In the example technique, a system, such as system 10, may determine a global electromechanical delay at a potential implant site during an intrinsic heart rhythm (850). In the example technique, a system may then determine local electromechanical delay at the same potential implant site during an intrinsic heart rhythm (852). A system may then determine a baseline electromechanical dyssynchrony index for an intrinsic heart rhythm (854). A system may determine the baseline electromechanical dyssynchrony index by comparing the differences between the global and local electromechanical delays during the intrinsic heart rhythm.

In the example technique, a system may then determine a global electromechanical delay at the same potential implant site during the delivery of electrical pacing therapy (856). In the example technique, a system may then determine local electromechanical delay at the same potential implant site during the delivery of electrical pacing therapy (858). A system may then determine an electromechanical dyssynchrony index during pacing therapy (860).

In the example technique, a system may then determine the percentage change between the baseline electromechanical dyssynchrony index and the electromechanical dyssynchrony index determined during pacing therapy (862). A system may then determine if there was a reduction from the baseline dyssynchrony index at the tested potential implant site above a threshold percentage (864). If the system determines that

the performance of the pacing at the potential implant site did produce a reduction in the baseline electromechanical dyssynchrony index above a threshold percentage (YES of 864), the system may select that implant site (866) for intracardiac lead implantation. In some examples, a processor of the system, such as processor 72, may automatically select the implant site according to the described techniques. In other examples, a user, based on information from a processor may manually select the implant site according to the described techniques. If the system determines that pacing at the potential site did not reduce the baseline electromechanical dyssynchrony index by a threshold percentage amount (NO of 864), the system may check to see if there are other potential lead implant sites to check (868). If there are still other implant sites to check (YES of 868), in the example technique, a system may change the potential lead implant site (870) and return to step 856. If there are no more potential lead implant sites to check (NO of 868), a system may select the implant site with the greatest reduction in the baseline electromechanical dyssynchrony index as the preferred intracardiac lead implant site.

In another example, the system may detect the time-interval from delivery of pace at a localized region within the heart to the onset of depolarization on a surface ECG lead, with the pace delivered at maximum pacing voltage (~ 6V to ensure that maximum energy is delivered during pacing) and at a short atrio-ventricular (A-V) interval (≤ 60 ms). If this time interval exceeds a certain threshold, then the system will indicate that particular area is not a suitable site for implanting the pacing lead. This is because a long time delay between delivery of local pace (at maximum energy) and onset of global depolarization on surface ECG indicates that the substrate of that area in the heart may be non-viable or otherwise not be a suitable site for implanting the pacing lead, e.g., comprised of scar tissue. FIG. 16 illustrates an example of the time-delay between the delivery of pacing and the onset of depolarization. FIG. 16 illustrates an example ECG signal 990. Arrow 992 points to the point in time when a patient received pacing at a localized region of the heart. Arrow 994 points to the determined point in time of the onset of the depolarization wave. The onset of the depolarization wave may be found, for example, according to the techniques described in this disclosure. Time period 996 is the difference in time between the beginning of pacing and the onset of the depolarization wave at a localized region of the heart. The time period 996 may also be termed the local electrical-electrical delay.

FIGS. 13-15 are conceptual diagrams illustrating an example system 910 that determines the onset and offset points of heart depolarization and repolarization waves of patient 914 according to the techniques described herein. As with the previously described system, in one example, system 910 may detect the onset and offset points of heart depolarization and repolarization waves in order to select a preferred location for implanting an intracardiac lead. In other examples, system 910 may detect the onset and offset points of heart depolarization and repolarization waves in order to select other parameters based on the electromechanical delay. For example, the system may select a particular pacing electrode configuration or pacing intervals for cardiac resynchronization therapy. In another example, with multipolar leads offering choices of more than one pacing electrode (cathode) in the ventricle, the system/device may automatically pace from each of the pacing electrodes at maximum pacing voltage and at a nominal atrio-ventricular delay (~100 ms) and measure onsets and offsets of the resulting depolarization waveforms on a far-field electrogram or on a leadless ECG or on a surface ECG lead, and choose the pacing electrode which produces the minimum difference between the offset of the local electrogram and the corresponding far-field onset, for delivery of cardiac resynchronization therapy. Alternatively, the pacing electrode which produces the narrowest far-field electrogram or surface ECG signal calculated as the difference between the offset and onset could be chosen. As illustrated in example diagram FIG. 13, a system 910 includes implantable medical device (IMD) 916, which is connected to leads 918, 920, and 922, and communicatively coupled to a programmer 924. IMD 916 senses electrical signals attendant to the depolarization and repolarization of heart 912, e.g., a cardiac EGM, via electrodes on one or more of leads 918, 920 and 922 or a housing of IMD 916. IMD 916 also delivers therapy in the form of electrical signals to heart 912 via electrodes located on one or more leads 918, 920, and 922 or a housing of IMD 916, such as pacing, cardioversion and/or defibrillation pulses. IMD 916 may include or be coupled to various sensors, such as one or more accelerometers, for detecting other physiological parameters of patient 14, such as activity or posture.

In some examples, programmer 924 takes the form of a handheld computing device, computer workstation, or networked computing device that includes a user interface for presenting information to and receiving input from a user. A user, such as a physician, technician, surgeon, electrophysiologist, or other clinician, may interact with

programmer 924 to communicate with IMD 916. For example, the user may interact with programmer 924 to retrieve physiological or diagnostic information from IMD 916. A user may also interact with programmer 924 to program IMD 916, e.g., select values for operational parameters of the IMD.

5 IMD 916 and programmer 924 may communicate via wireless communication using any techniques known in the art. Examples of communication techniques may include, for example, low frequency or radiofrequency (RF) telemetry, but other techniques are also contemplated. In some examples, programmer 924 may include a programming head that may be placed proximate to the patient's body near the IMD 916
10 implant site in order to improve the quality or security of communication between IMD 916 and programmer 924. In other examples, programmer 924 may be located remotely from IMD 916, and communicate with IMD 916 via a network.

 The techniques for identifying onsets and/or offsets of cardiac electrogram waves may be performed by IMD 916, e.g., by a processor of IMD 916, based on one or more
15 cardiac electrograms sensed by the IMD. In other examples, as described previously, some or all of the functions ascribed to IMD 916 or a processor thereof may be performed by one or more other devices, such as programmer 294 or a workstation (not shown), or a processor thereof. For example, programmer 924 may process EGM signals received from IMD 916 and/or cardiac mechanical contraction information to according to the
20 techniques described herein. Furthermore, although described herein with respect to an IMD, in other examples, the techniques described herein may be performed by or implemented in an external medical device, which may be coupled to a patient via percutaneous or transcutaneous leads.

 FIG. 14 is a conceptual diagram illustrating IMD 916 and leads 918, 920, and 922
25 of system 910 in greater detail. In the illustrated example, bipolar electrodes 940 and 942 are located adjacent to a distal end of lead 918. In addition, bipolar electrodes 944 and 946 are located adjacent to a distal end of lead 920, and bipolar electrodes 948 and 950 are located adjacent to a distal end of lead 922.

 In the illustrated example, electrodes 940, 944 and 948 take the form of ring
30 electrodes, and electrodes 942, 946 and 950 may take the form of extendable helix tip electrodes mounted retractably within insulative electrode heads 952, 954 and 956, respectively. Leads 918, 920, 922 also include elongated electrodes 962, 964, 966,

respectively, which may take the form of a coil. In some examples, each of the electrodes 940, 942, 944, 946, 948, 950, 962, 964 and 966 is electrically coupled to a respective conductor within the lead body of its associated lead 918, 920, 922, and thereby coupled circuitry within IMD 916.

5 In some examples, IMD 916 includes one or more housing electrodes, such as housing electrode 904 illustrated in FIG. 14, which may be formed integrally with an outer surface of hermetically-sealed housing 908 of IMD 916 or otherwise coupled to housing 908. In some examples, housing electrode 904 is defined by an uninsulated portion of an outward facing portion of housing 908 of IMD 916. Other division between insulated and
10 uninsulated portions of housing 908 may be employed to define two or more housing electrodes. In some examples, a housing electrode comprises substantially all of housing 908.

 As described in further detail with reference to FIG. 15, housing 908 encloses a signal generator that generates therapeutic stimulation, such as cardiac pacing,
15 cardioversion and defibrillation pulses, as well as a sensing module for sensing electrical signals attendant to the depolarization and repolarization of heart 912. Housing 908 may also enclose a wave detection module that detects the onsets and offsets of heart depolarization and repolarization waves. The wave detection module may be enclosed within housing 908. Alternatively, the wave detection module may be housed in a remote
20 piece of equipment, such as programmer 924 or a workstation (not shown) and communicate with the IMD 916 through wireless communication.

 IMD 916 senses electrical signals attendant to the depolarization and repolarization of heart 912 via electrodes 904, 940, 942, 944, 946, 948, 950, 962, 964 and 966. IMD 916 may sense such electrical signals via any bipolar combination of electrodes 940, 942, 944,
25 946, 948, 950, 962, 964 and 966. Furthermore, any of the electrodes 940, 942, 944, 946, 948, 950, 962, 964 and 966 may be used for unipolar sensing in combination with housing electrode 904.

 In some examples, IMD 916 delivers pacing pulses via bipolar combinations of electrodes 940, 942, 944, 946, 948 and 950 to produce depolarization of cardiac tissue of
30 heart 912. In some examples, IMD 16 delivers pacing pulses via any of electrodes 940, 942, 944, 946, 948 and 950 in combination with housing electrode 904 in a unipolar configuration. Furthermore, IMD 916 may deliver cardioversion or defibrillation pulses to

heart 12 via any combination of elongated electrodes 962, 964, 966, and housing electrode 904.

The illustrated numbers and configurations of leads 918, 920, and 922 and electrodes are merely examples. Other configurations, i.e., number and position of leads and electrodes, are possible. In some examples, system 910 may include an additional lead or lead segment having one or more electrodes positioned at different locations in the cardiovascular system for sensing and/or delivering therapy to patient 914. For example, instead of or in addition to intracardiac leads 918, 920 and 922, system 910 may include one or more epicardial or subcutaneous leads not positioned within the heart. In some examples, the subcutaneous leads may sense a subcutaneous cardiac electrogram, for example the far-field electrogram between the SVC coil and the can. The subcutaneous cardiac electrogram may substitute for the surface ECG in determining the global electromechanical delay.

FIG. 15 is a block diagram illustrating an example configuration of IMD 916. In the illustrated example, IMD 916 includes a processor 970, memory 972, signal generator 974, sensing module 976, telemetry module 978, motion sensing module 980, wave detection module 982 and peak detection module 984. Memory 972 includes computer-readable instructions that, when executed by processor 970, causes IMD 916 and processor 970 to perform various functions attributed to IMD 916 and processor 970 herein. Memory 972 may include any volatile, non-volatile, magnetic, optical, or electrical media, such as a random access memory (RAM), read-only memory (ROM), non-volatile RAM (NVRAM), electrically-erasable programmable ROM (EEPROM), flash memory, or any other digital or analog media.

Processor 970 may include any one or more of a microprocessor, a controller, a digital signal processor (DSP), an application specific integrated circuit (ASIC), a field-programmable gate array (FPGA), or equivalent discrete or analog logic circuitry. In some examples, processor 970 may include multiple components, such as any combination of one or more microprocessors, one or more controllers, one or more DSPs, one or more ASICs, or one or more FPGAs, as well as other discrete or integrated logic circuitry. The functions attributed to processor 970 herein may be embodied as software, firmware, hardware or any combination thereof. Generally, processor 970 controls signal generator 974 to deliver stimulation therapy to heart 912 of patient 914 according to a selected one

or more of therapy programs or parameters, which may be stored in memory 972. As an example, processor 970 may control signal generator 974 to deliver electrical pulses with the amplitudes, pulse widths, frequency, or electrode polarities specified by the selected one or more therapy programs.

5 Signal generator 974 is configured to generate and deliver electrical stimulation therapy to patient 912. As shown in FIG. 15, signal generator 974 is electrically coupled to electrodes 94, 940, 942, 944, 946, 948, 950, 962, 964, and 966, e.g., via conductors of the respective leads 918, 920, and 922 and, in the case of housing electrode 904, within housing 908. For example, signal generator 974 may deliver pacing, defibrillation or
10 cardioversion pulses to heart 912 via at least two of electrodes 94, 940, 942, 944, 946, 948, 950, 962, 964, and 966. In other examples, signal generator 974 delivers stimulation in the form of signals other than pulses, such as sine waves, square waves, or other substantially continuous time signals.

 Signal generator 974 may include a switch module (not shown) and processor 970
15 may use the switch module to select, e.g., via a data/address bus, which of the available electrodes are used to deliver the electrical stimulation. The switch module may include a switch array, switch matrix, multiplexer, or any other type of switching device suitable to selectively couple stimulation energy to selected electrodes. Electrical sensing module 976 monitors electrical cardiac signals from any combination of electrodes 904, 940, 942,
20 944, 946, 948, 950, 962, 964, and 966. Sensing module 976 may also include a switch module which processor 970 controls to select which of the available electrodes are used to sense the heart activity, depending upon which electrode combination is used in the current sensing configuration.

 Sensing module 976 may include one or more detection channels, each of which
25 may comprise an amplifier. The detection channels may be used to sense the cardiac signals. Some detection channels may detect events, such as R- waves or P-waves, and provide indications, such as wave markers, of the occurrences of such events to processor 970. One or more other detection channels may provide the signals to an analog-to-digital converter, for conversion into a digital signal for processing or analysis by processor 970.

30 For example, sensing module 976 may comprise one or more narrow band channels, each of which may include a narrow band filtered sense-amplifier that compares the detected signal to a threshold. If the filtered and amplified signal is greater than the

threshold, the narrow band channel indicates that a certain electrical cardiac event, e.g., depolarization, has occurred. Processor 970 then uses that detection in measuring frequencies of the sensed events.

5 In one example, at least one narrow band channel may include an R-wave or P-wave amplifier. In some examples, the R-wave and P-wave amplifiers may take the form of an automatic gain controlled amplifier that provides an adjustable sensing threshold as a function of the measured R-wave or P-wave amplitude. Examples of R-wave and P-wave amplifiers are described in U.S. Patent No. 5,117,824 to Keimel et al., which issued on June 2, 1992 and is entitled, "APPARATUS FOR MONITORING ELECTRICAL
10 PHYSIOLOGIC SIGNALS,".

In some examples, sensing module 976 includes a wide band channel which may comprise an amplifier with a relatively wider pass band than the narrow band channels. Signals from the electrodes that are selected for coupling to this wide-band amplifier may be converted to multi-bit digital signals by an analog-to-digital converter (ADC) provided
15 by, for example, sensing module 976 or processor 970. Processor 970 may analyze the digitized versions of signals from the wide band channel. Processor 970 may employ digital signal analysis techniques to characterize the digitized signals from the wide band channel to, for example, detect and classify the patient's heart rhythm.

Processor 970 may detect and classify the patient's heart rhythm based on the
20 cardiac electrical signals sensed by sensing module 976 employing any of the numerous signal processing methodologies known in the art. For example, processor 970 may maintain escape interval counters that may be reset upon sensing of R-waves by sensing module 976. The value of the count present in the escape interval counters when reset by sensed depolarizations may be used by processor 970 to measure the durations of R-R
25 intervals, which are measurements that may be stored in memory 972. Processor 970 may use the count in the interval counters to detect a tachyarrhythmia, such as ventricular fibrillation or ventricular tachycardia. A portion of memory 972 may be configured as a plurality of recirculating buffers, capable of holding series of measured intervals, which may be analyzed by processor 970 to determine whether the patient's heart 912 is presently
30 exhibiting atrial or ventricular tachyarrhythmia.

In some examples, processor 970 may determine that tachyarrhythmia has occurred by identification of shortened R-R interval lengths. Generally, processor 970 detects

tachycardia when the interval length falls below 360 milliseconds (ms) and fibrillation when the interval length falls below 320 ms. These interval lengths are merely examples, and a user may define the interval lengths as desired, which may then be stored within memory 972. This interval length may need to be detected for a certain number of
5 consecutive cycles, for a certain percentage of cycles within a running window, or a running average for a certain number of cardiac cycles, as examples.

In some examples, an arrhythmia detection method may include any suitable tachyarrhythmia detection algorithms. In one example, processor 970 may utilize all or a subset of the rule-based detection methods described in U.S. Patent No. 5,545,186 to
10 Olson et al., entitled, "PRIORITIZED RULE BASED METHOD AND APPARATUS FOR DIAGNOSIS AND TREATMENT OF ARRHYTHMIAS," which issued on August 13, 1996, or in U.S. Patent No. 5,755,736 to Gillberg et al., entitled, "PRIORITIZED RULE BASED METHOD AND APPARATUS FOR DIAGNOSIS AND TREATMENT OF ARRHYTHMIAS," which issued on May 26, 1998. However, other arrhythmia
15 detection methodologies may also be employed by processor 970 in other examples. For example, EGM morphology may be considered in addition to or instead of interval length for detecting tachyarrhythmias.

Generally, processor 970 detects a treatable tachyarrhythmia, such as VF, based on the EGM, e.g., the R-R intervals and/or morphology of the EGM, and selects a therapy to
20 deliver to terminate the tachyarrhythmia, such as a defibrillation pulse of a specified magnitude. The detection of the tachyarrhythmia may include a number of phases or steps prior to delivery of the therapy, such as first phase, sometimes referred to as detection, in which a number of consecutive or proximate R-R intervals satisfies a first number of intervals to detect (NID) criterion, a second phase, sometimes referred to as confirmation,
25 in which a number of consecutive or proximate R-R intervals satisfies a second, more restrictive NID criterion. Tachyarrhythmia detection may also include confirmation based on EGM morphology or other sensors subsequent to or during the second phase.

In the illustrated example, IMD 916 also includes peak detection module 980, wave detection module 982, and motion sensing module 984. Peak detection module 980
30 and wave detection module 982 may be configured and provide the functionality ascribed to peak detection module 76 and wave detection module 78 herein. Peak detection module 980 may be configured to determine a maximum value of a particular signal. For

example, peak detection module 980 may be configured to receive an electrical signal from wave detection module 982 or processor 970 and determine a maximum value of the received signal. Peak detection module 980 may, in some examples, comprise a narrow-band channel of sensing module 976 that is configured to detect R-waves, P-waves, or T-waves in cardiac electrogram signals, e.g., using an amplifier with automatically adjusting threshold.

Generally, wave detection module 982 determines the onsets and offsets on the heart depolarization and repolarizations waves. Wave detection module 982 may be similar to the wave detection module described previously, e.g. wave detection module 78, and more accurately described in FIG. 2. For example, wave detection module may include a low-pass filter, a window module, a slope module, a rectifier module, a smoothing module, and a threshold detection module. Wave detection module 982 may perform substantially similar to the wave detection module described previously in this application.

Peak detection module 980 and wave detection module 982 may receive a cardiac electrogram from sensing module 976, e.g., from a wide-band channel of the sensing module. In some examples, the cardiac electrogram may be a far-field cardiac electrogram, e.g., between superior vena cava coil 766 and housing electrode 904. A far-field cardiac electrogram may be used in the manner described herein with respect to a surface ECG, e.g., to determine a global electromechanical delay. In some examples, the cardiac electrogram may be a unipolar cardiac electrogram between housing electrode 906 and any of electrodes 942, 944, 946 and 950. The unipolar cardiac electrogram may be received from sensing module 976, e.g., via a wide-band channel of the sensing module, and may be used in the manner described herein with respect to local cardiac electrogram signals, e.g., to determine local electromechanical delays.

Motion sensing module 984 may sense the mechanical contraction of the heart, e.g., at one or more cardiac sites. Motion sensing module 984 may be electrically coupled to one or more sensors that generate a signal that varies based on cardiac contraction or motion generally, such as one or more accelerometers, pressure sensors, impedance sensors, or flow sensors. The detected contraction may be contraction of cardiac tissue at a particular location, e.g., a particular portion of a ventricular wall.

Although processor 970 and wave detection module 982 are illustrated as separate modules in FIG. 15, processor 970 and wave detection module 982 may be incorporated in a single processing unit. Wave detection module 982, and any of its components, may be a component of or module executed by processor 970.

5 Telemetry module 978 includes any suitable hardware, firmware, software or any combination thereof for communicating with another device, such as programmer 924 (FIG. 13). Under the control of processor 970, telemetry module 978 may receive downlink telemetry from and send uplink telemetry to programmer 924 with the aid of an antenna, which may be internal and/or external. In some examples, processor 970 may
10 transmit cardiac signals produced by sensing module 976 and/or signals generated by heart sound sensor 982 to programmer 924. Processor 970 may also generate and store marker codes indicative of different cardiac events that sensing module 976 or heart sound analyzer 980 detects, and transmit the marker codes to programmer 924. An example
15 IMD with marker-channel capability is described in U.S. Patent No. 4,374,382 to Markowitz, entitled, "MARKER CHANNEL TELEMETRY SYSTEM FOR A MEDICAL DEVICE," which issued on February 15, 1983. Information which processor 970 may transmit to programmer 924 via telemetry module 978 may also include indications of treatable rhythms, and indications of non-treatable rhythms in which the EGM based indication indicated that the rhythm was treatable and the heart sound based
20 indication indicated that the rhythm was non-treatable. Such information may be included as part of a marker channel with an EGM.

The techniques described in this disclosure, including those attributed to IMD wave detection module 80, programmer 24, or various constituent components, may be implemented, at least in part, in hardware, software, firmware or any combination thereof.
25 For example, various aspects of the techniques may be implemented within one or more processors, including one or more microprocessors, digital signal processors (DSPs), application specific integrated circuits (ASICs), field programmable gate arrays (FPGAs), or any other equivalent integrated or discrete logic circuitry, as well as any combinations of such components, embodied in programmers, such as physician or patient
30 programmers, stimulators, image processing devices or other devices. The term "processor" or "processing circuitry" may generally refer to any of the foregoing logic

circuitry, alone or in combination with other logic circuitry, or any other equivalent circuitry.

Such hardware, software, firmware may be implemented within the same device or within separate devices to support the various operations and functions described in this disclosure. In addition, any of the described units, modules or components may be
5 implemented together or separately as discrete but interoperable logic devices. Depiction of different features as modules or units is intended to highlight different functional aspects and does not necessarily imply that such modules or units must be realized by separate hardware or software components. Rather, functionality associated with one or
10 more modules or units may be performed by separate hardware or software components, or integrated within common or separate hardware or software components.

When implemented in software, the functionality ascribed to the systems, devices and techniques described in this disclosure may be embodied as instructions on a computer-readable medium such as random access memory (RAM), read-only memory
15 (ROM), non-volatile random access memory (NVRAM), electrically erasable programmable read-only memory (EEPROM), FLASH memory, magnetic data storage media, optical data storage media, or the like. The instructions may be executed to support one or more aspects of the functionality described in this disclosure.

20

CLAIMS:

1. A method comprising:
 - determining a slope of a cardiac electrogram;
 - determining a threshold value based on a maximum of the slope of the cardiac electrogram; and
 - identifying a last point of the cardiac electrogram before the slope of the cardiac electrogram crosses the threshold as one of an onset or an offset of a wave.
2. The method of claim 1, wherein the cardiac electrogram comprises one of a surface electrocardiogram, a local cardiac electrogram, or a subcutaneous cardiac electrogram.
3. The method of claim 1, wherein the threshold comprises a percentage of the maximum value of the slope of the cardiac electrogram.
4. The method of claim 1, wherein identifying the last point comprises identifying the last point of the cardiac electrogram before the slope of the cardiac electrogram exceeds the threshold as the onset of the wave.
5. The method of claim 4, wherein identifying the last point comprises identifying the last point within a portion of the cardiac electrogram prior to at least one of the maximum of the slope or a peak of the cardiac electrogram.
6. The method of claim 1, wherein identifying the last point comprises identifying the last point of the cardiac electrogram before the slope of the cardiac electrogram falls below the threshold as the offset of the wave.
7. The method of claim 6, wherein identifying the last point comprises identifying the last point within a portion of the cardiac electrogram after at least one of the maximum of the slope or a peak of the cardiac electrogram.

8. The method of claim 1, wherein the wave comprises one of a QRS complex, a P-wave, or a T-wave of the cardiac electrogram.
9. The method of claim 1, further comprising determining at least one of a QRS width or a Q-T interval based on the identified onset or offset of the wave.
10. The method of claim 1, further comprising:
 - receiving an indication of local ventricular motion associated with a cardiac contraction;
 - determining an electromechanical delay between the identified onset and the local ventricular motion; and
 - providing the electromechanical delay for configuration of cardiac resynchronization therapy.
11. The method of claim 10, further comprising selecting at least one of a pacing site, pacing electrode configuration, or pacing interval for cardiac resynchronization therapy based on the determined electromechanical delay.
12. The method of claim 10,
 - wherein the cardiac electrogram comprises a surface electrocardiogram far-field cardiac electrogram, or subcutaneous electrocardiogram,
 - wherein determining an electromechanical delay comprises determining a global electromechanical delay for each of a plurality of potential pacing sites,
 - the method further comprising selecting a subset of the pacing sites based on the global electromechanical delays for the pacing sites exceeding a global electromechanical delay threshold.

13. The method of claim 12,
wherein the cardiac electrogram comprises a plurality of local cardiac electrograms from each of the subset of pacing sites,
wherein determining an electromechanical delay comprises determining a local electromechanical delay for each of the subset pacing sites,
the method further comprising identifying one or more of the subset of pacing sites as candidate pacing sites based on local electromechanical delays for the candidate pacing sites being less than a local electromechanical delay threshold.
14. The method of claim 13, further comprising:
pacing at the candidate pacing sites;
during pacing at each of the candidate pacing sites, determining an electromechanical delay for each of a plurality of ventricular locations and determining an electromechanical dyssynchrony index based on the electromechanical delays; and
selecting one or more of the candidate pacing sites for cardiac resynchronization therapy based on dyssynchrony indices for the candidate pacing sites.
15. A system comprising:
a slope module that determines the slope of a cardiac electrogram; and
a wave detection module that determines a threshold value based on a maximum of the slope of the cardiac electrogram and identifies a last point of the cardiac electrogram before the slope of the cardiac electrogram crosses the threshold as one of an onset or and offset of a wave.

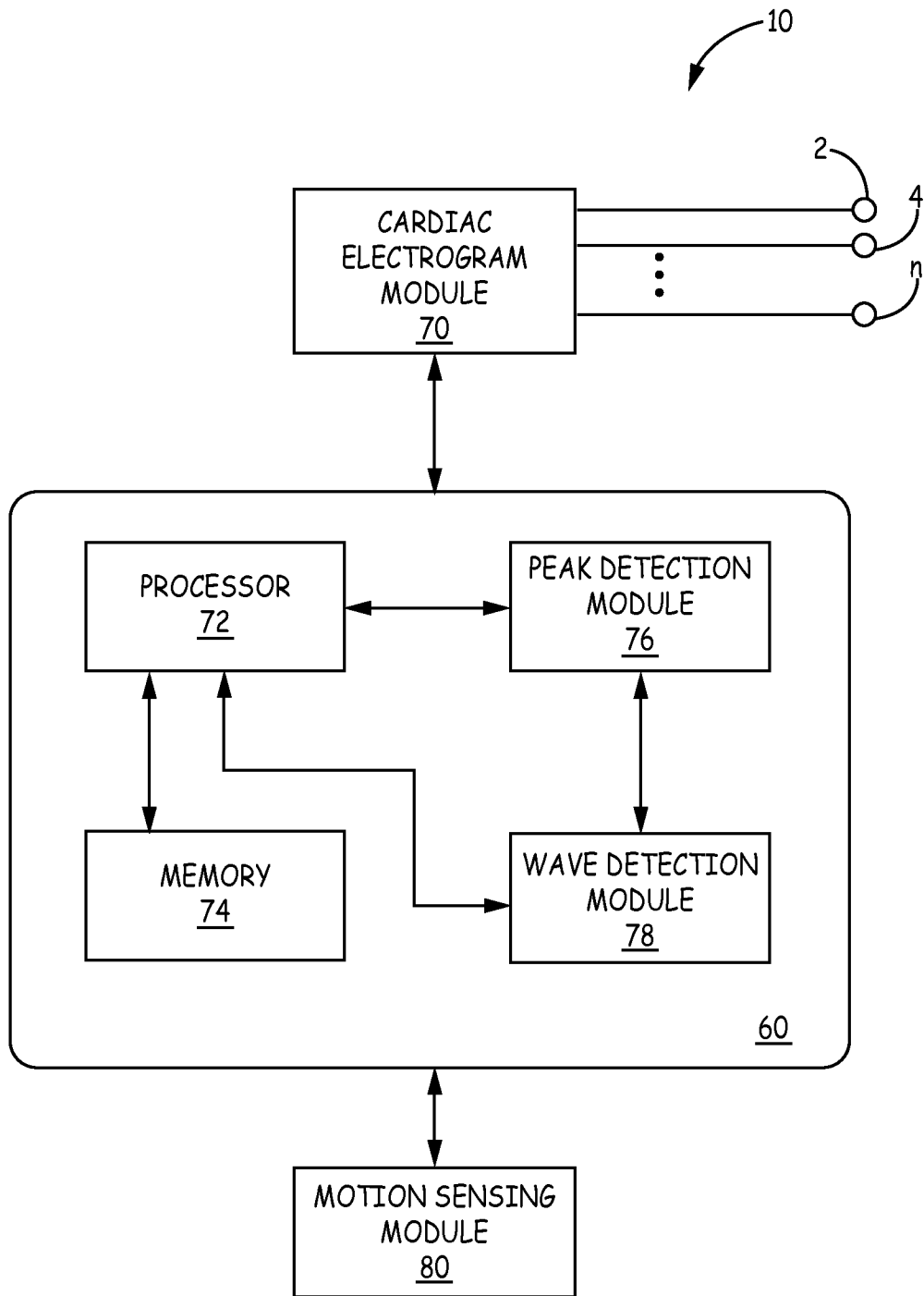


FIG. 1

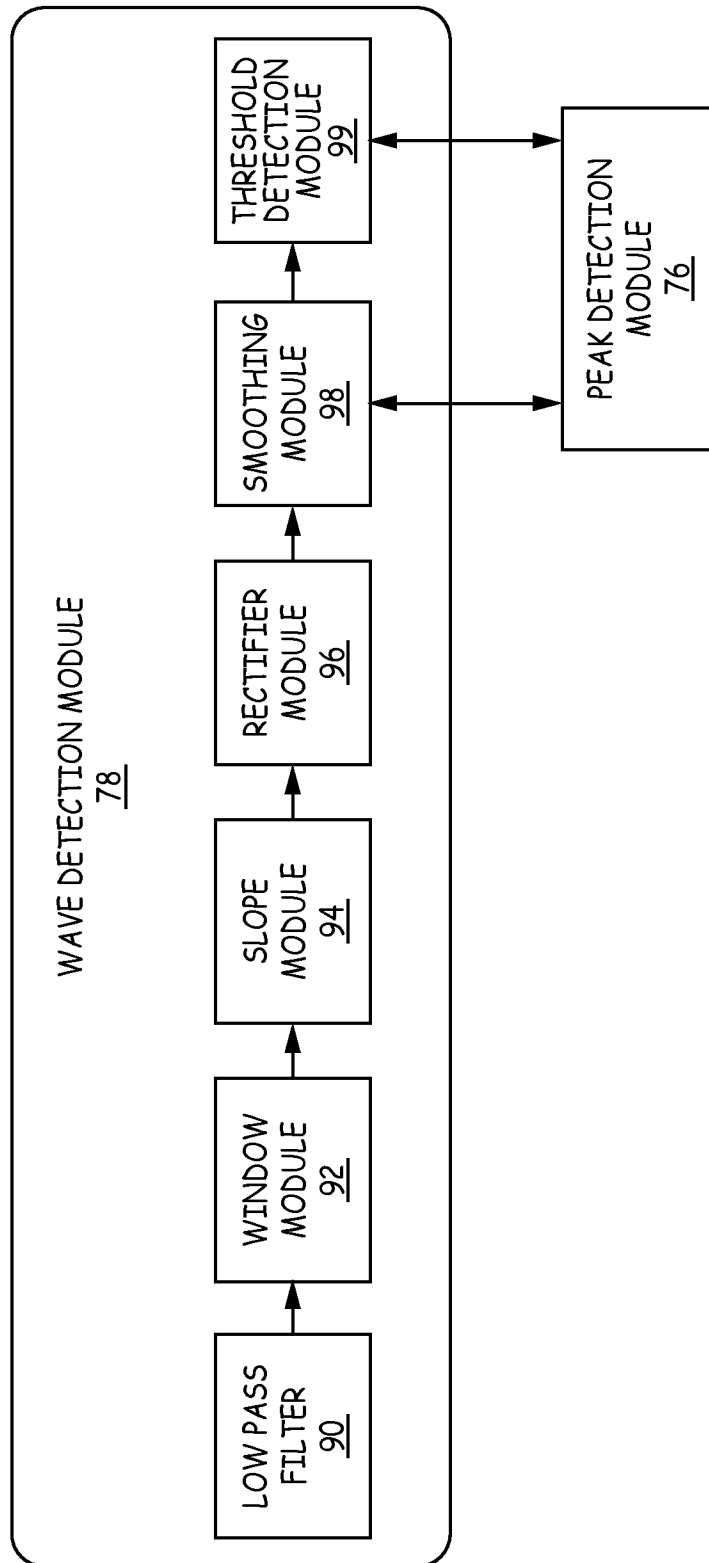


FIG. 2

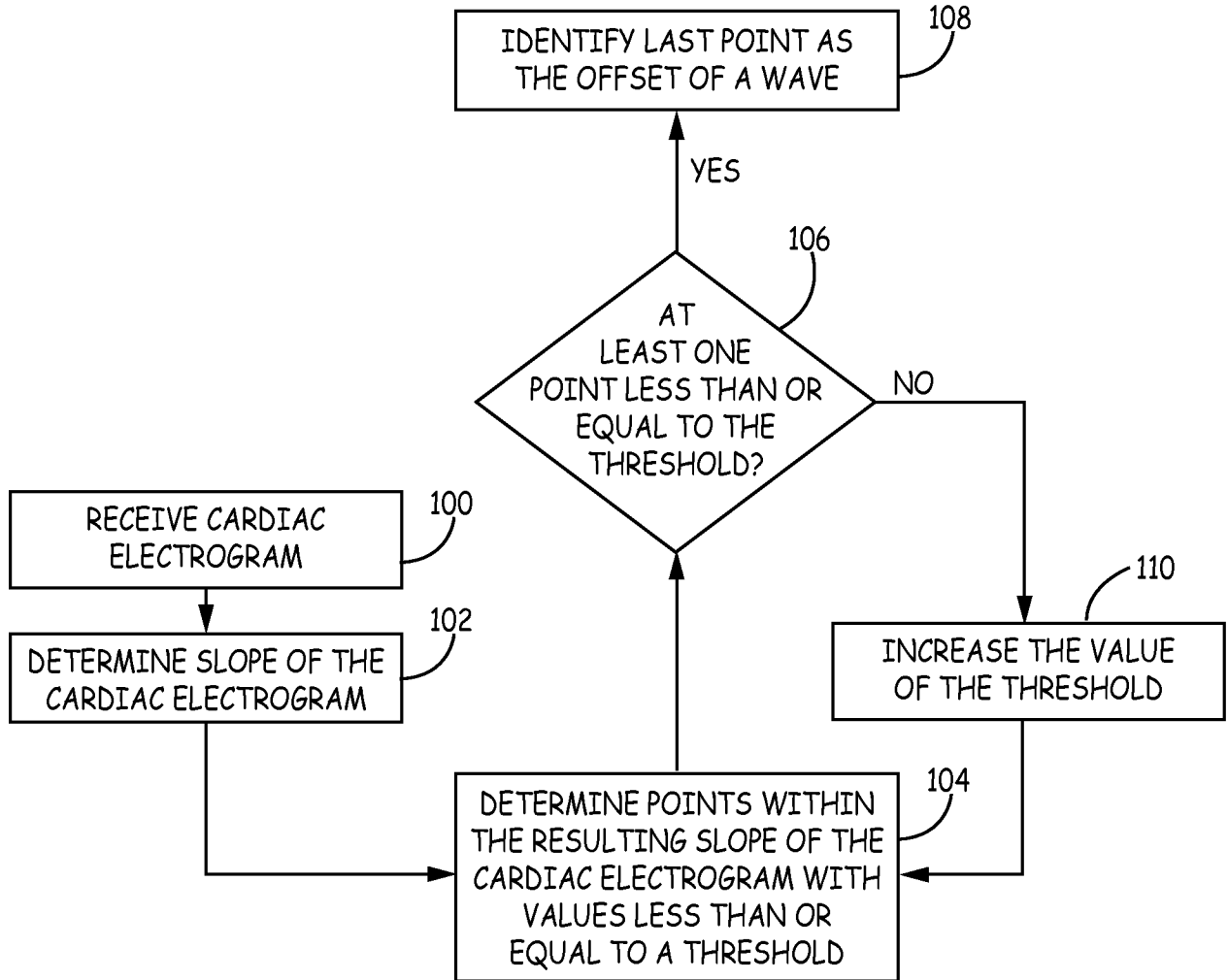


FIG. 3A

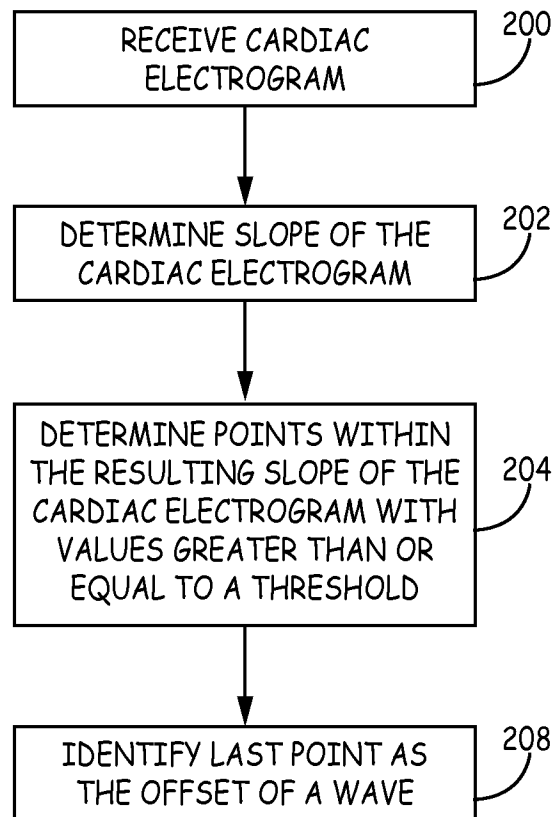


FIG. 3B

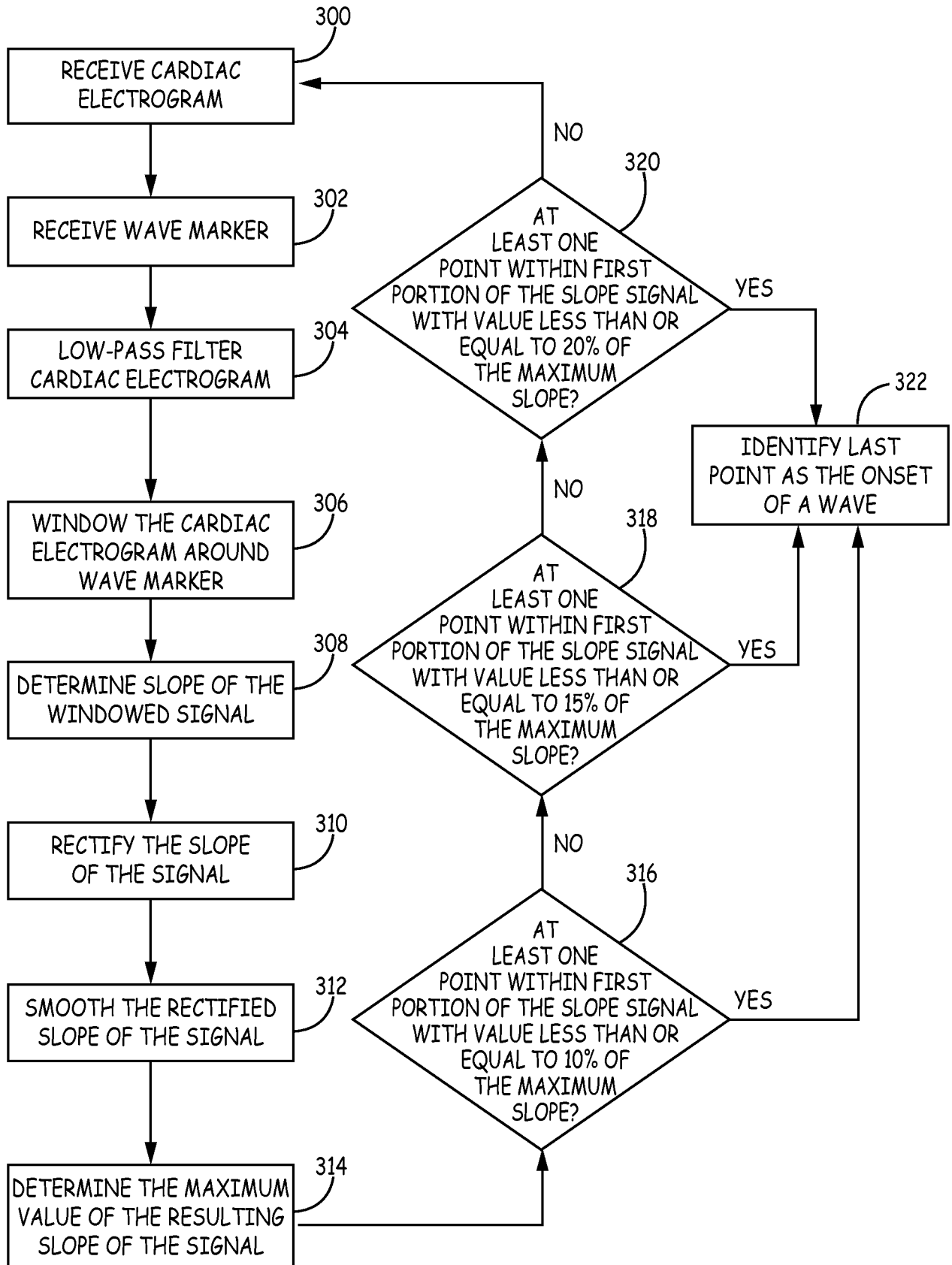


FIG. 4A

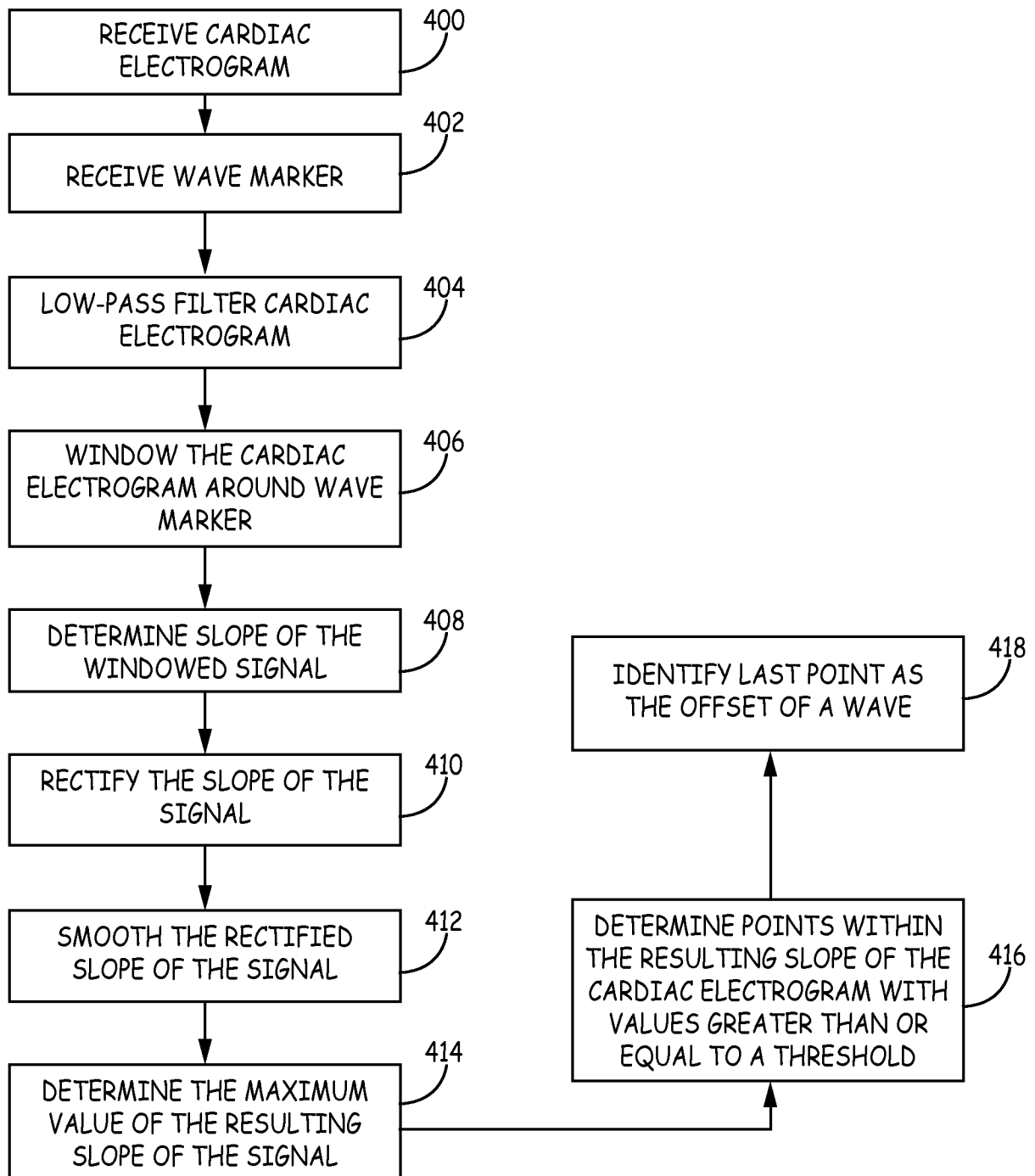


FIG. 4B

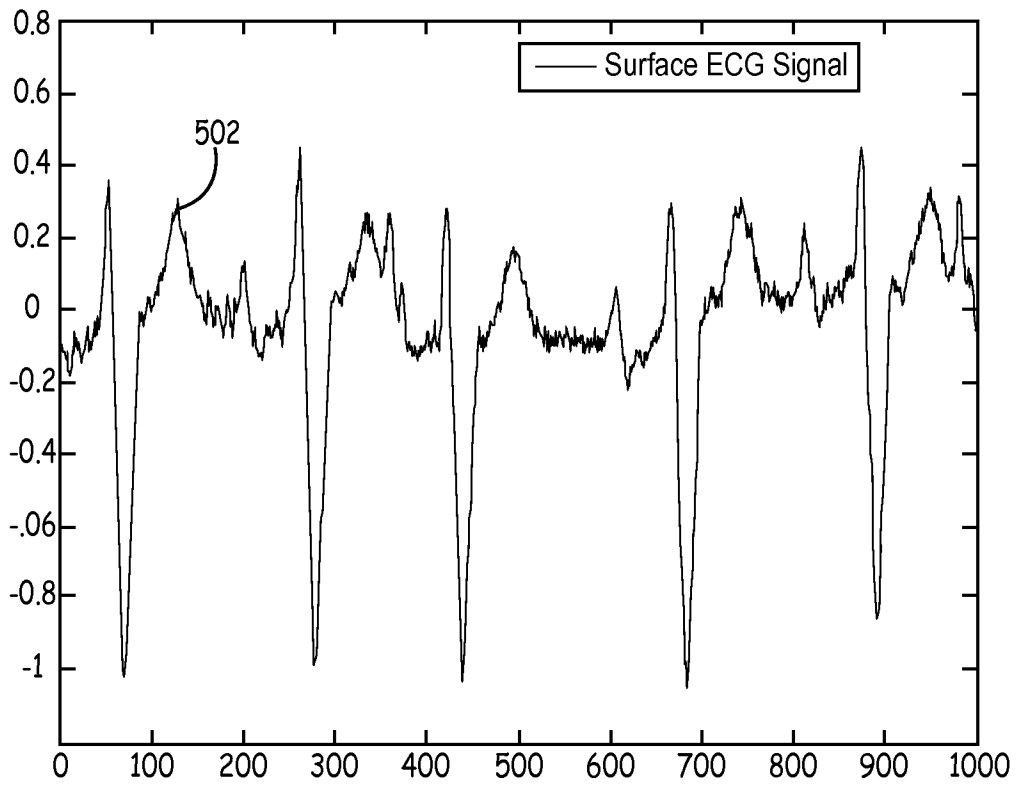


FIG. 5

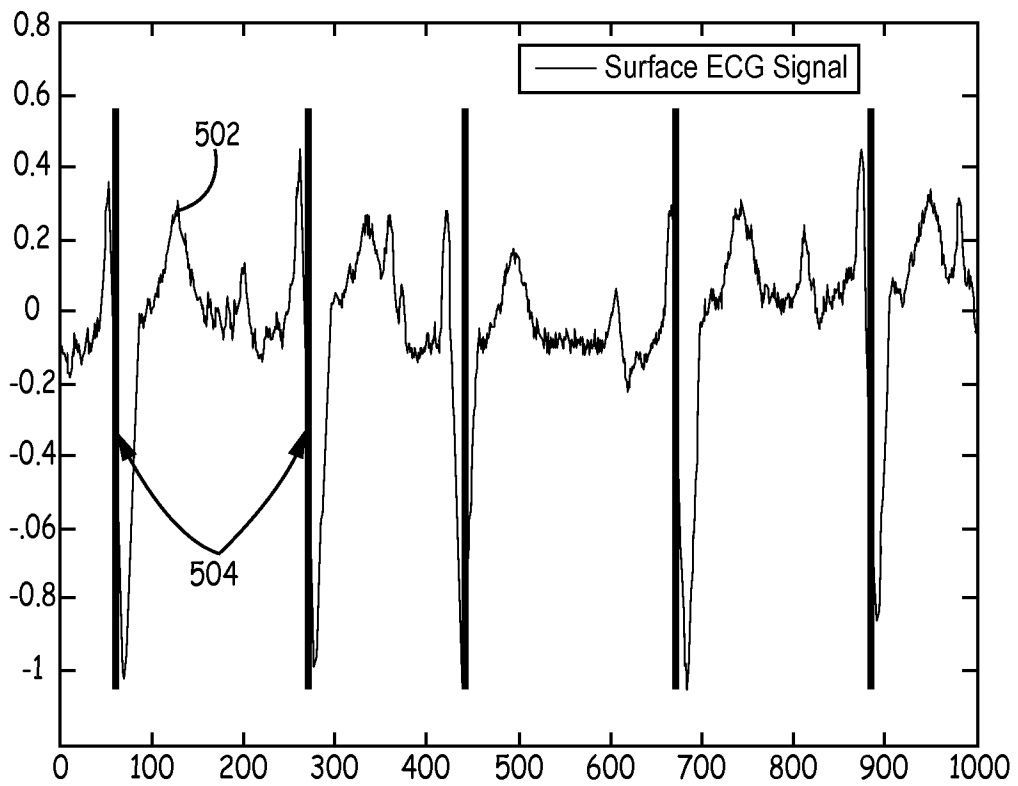


FIG. 6

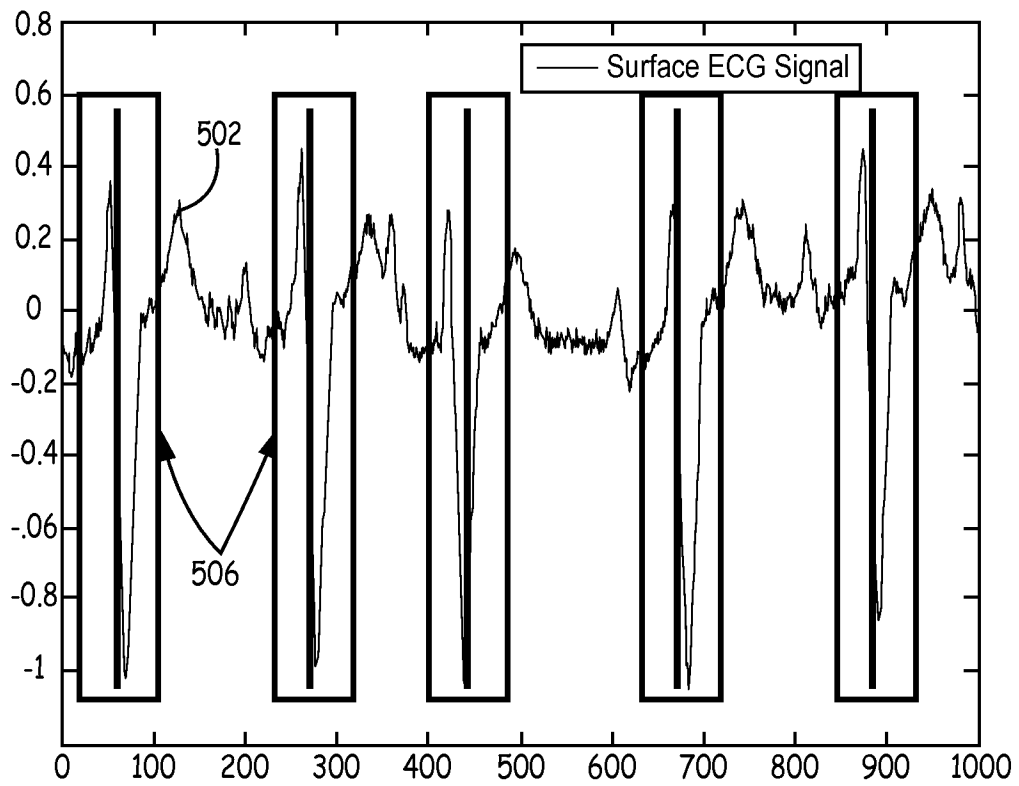


FIG. 7

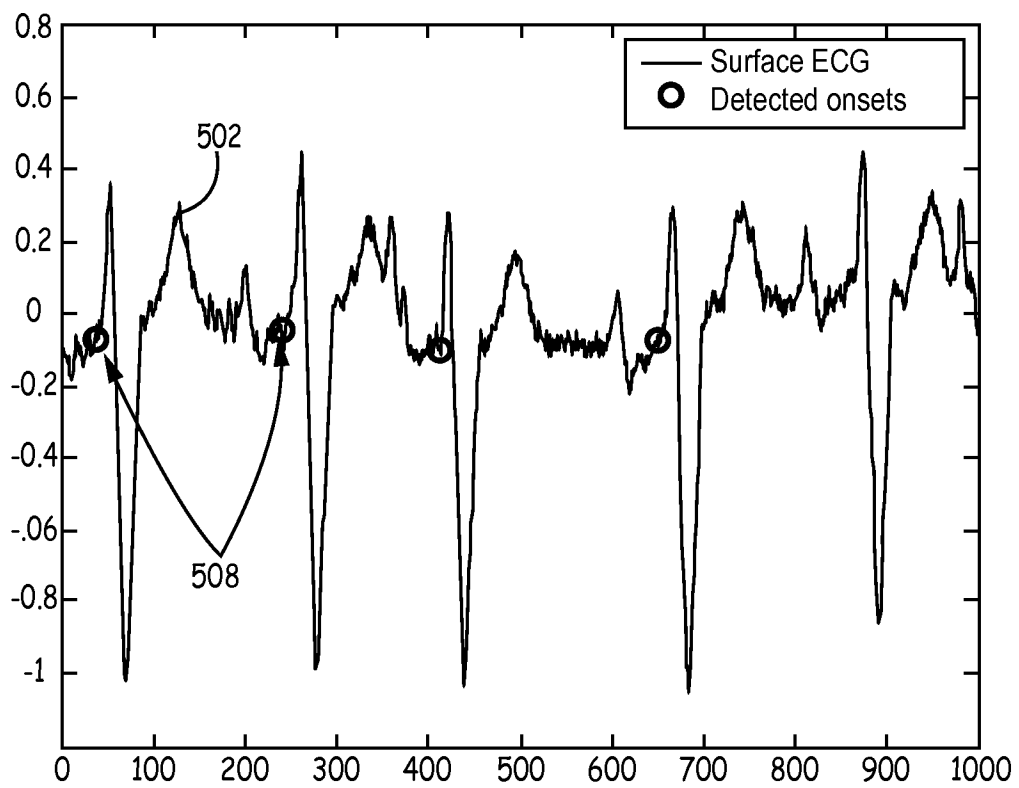


FIG. 8

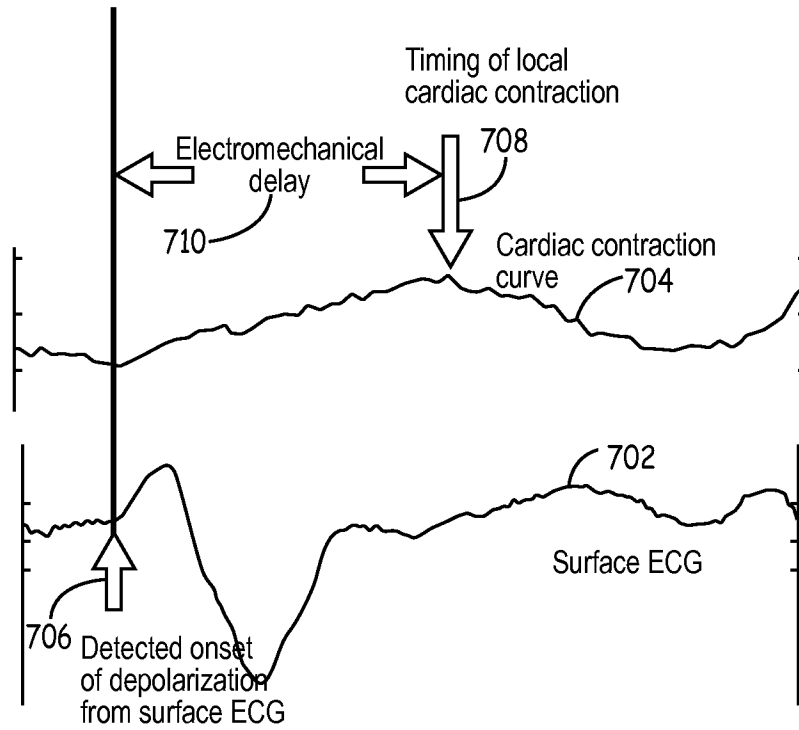


FIG. 9

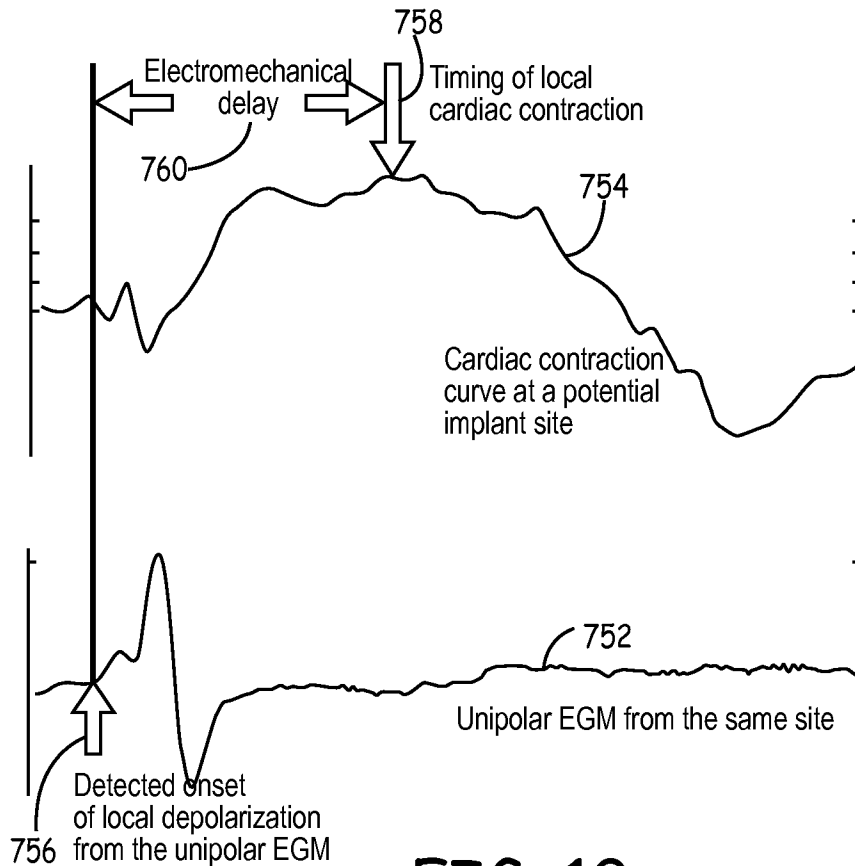


FIG. 10

10/15

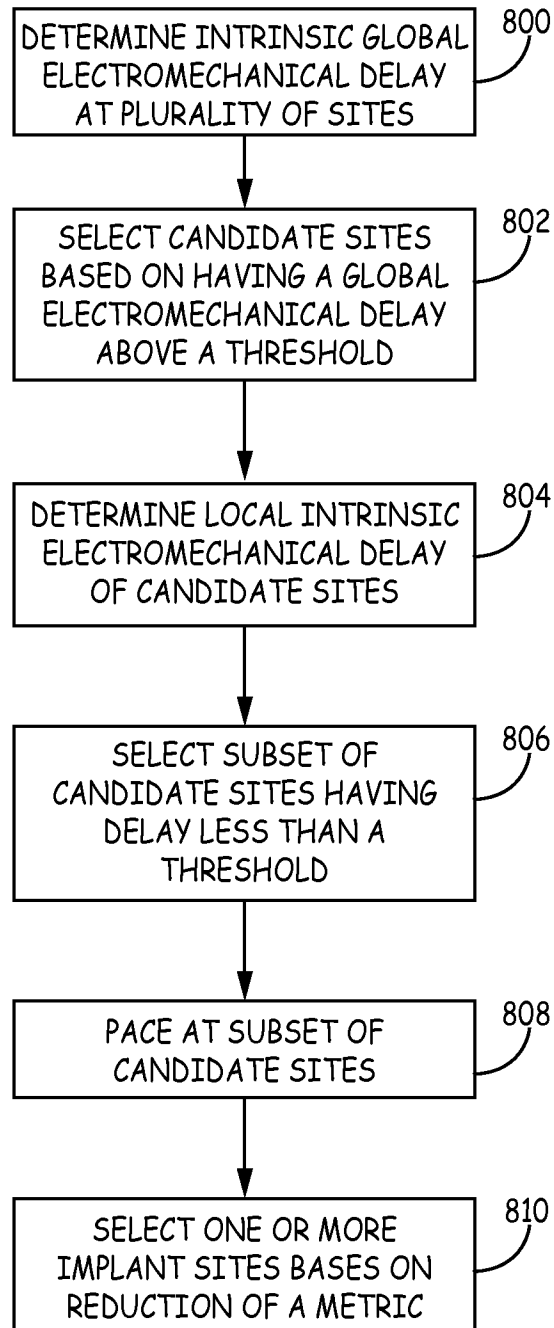


FIG. 11

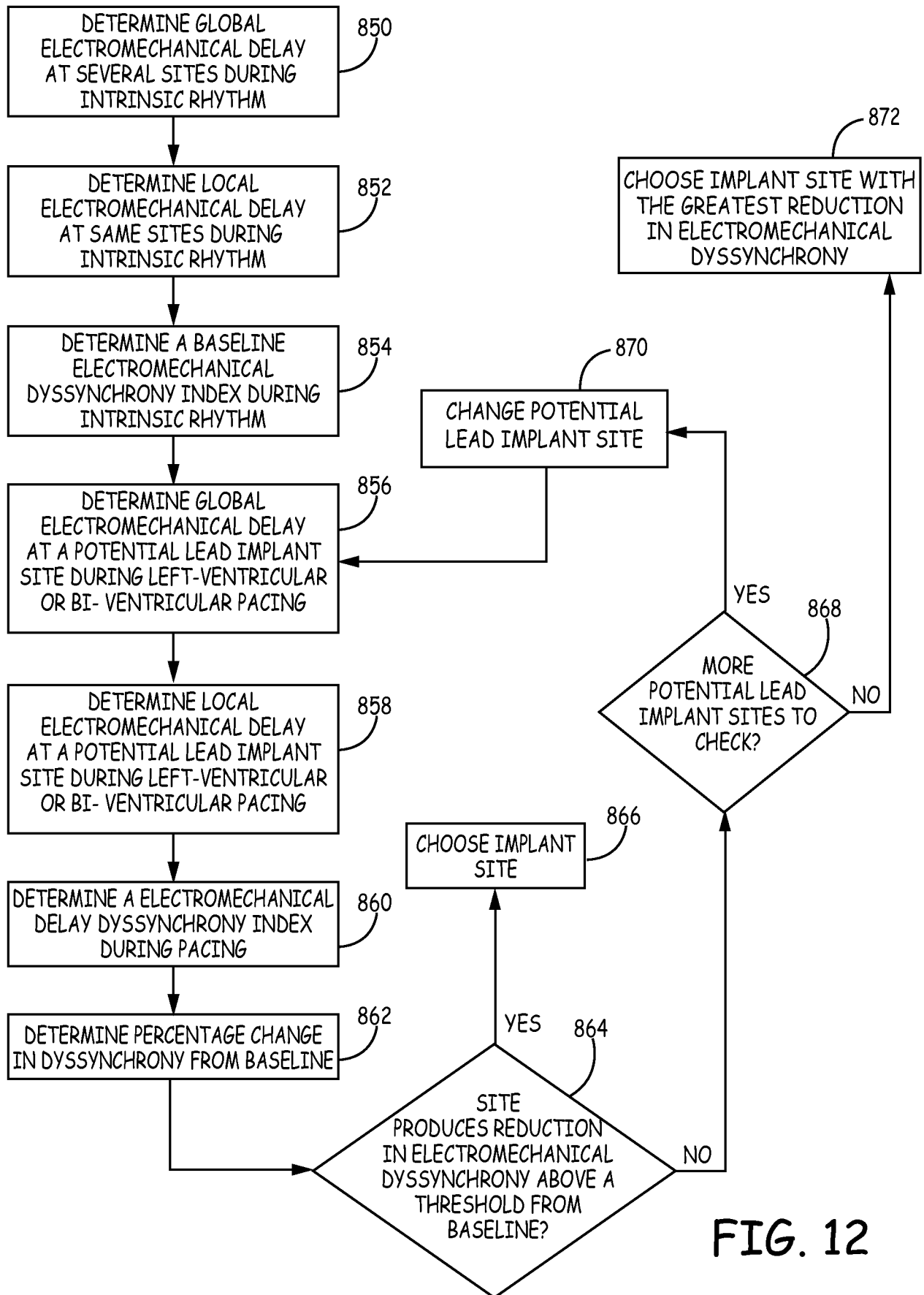


FIG. 12

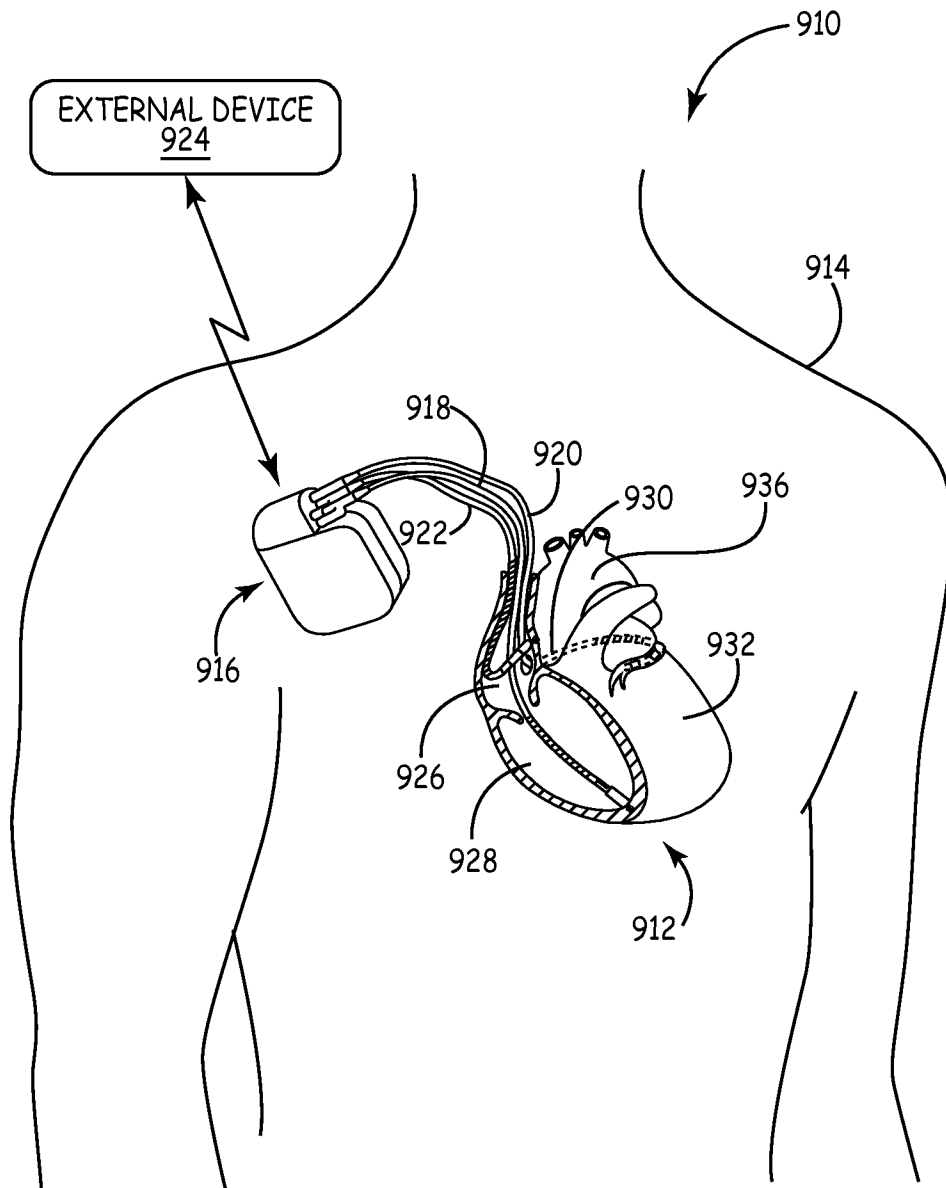


FIG. 13

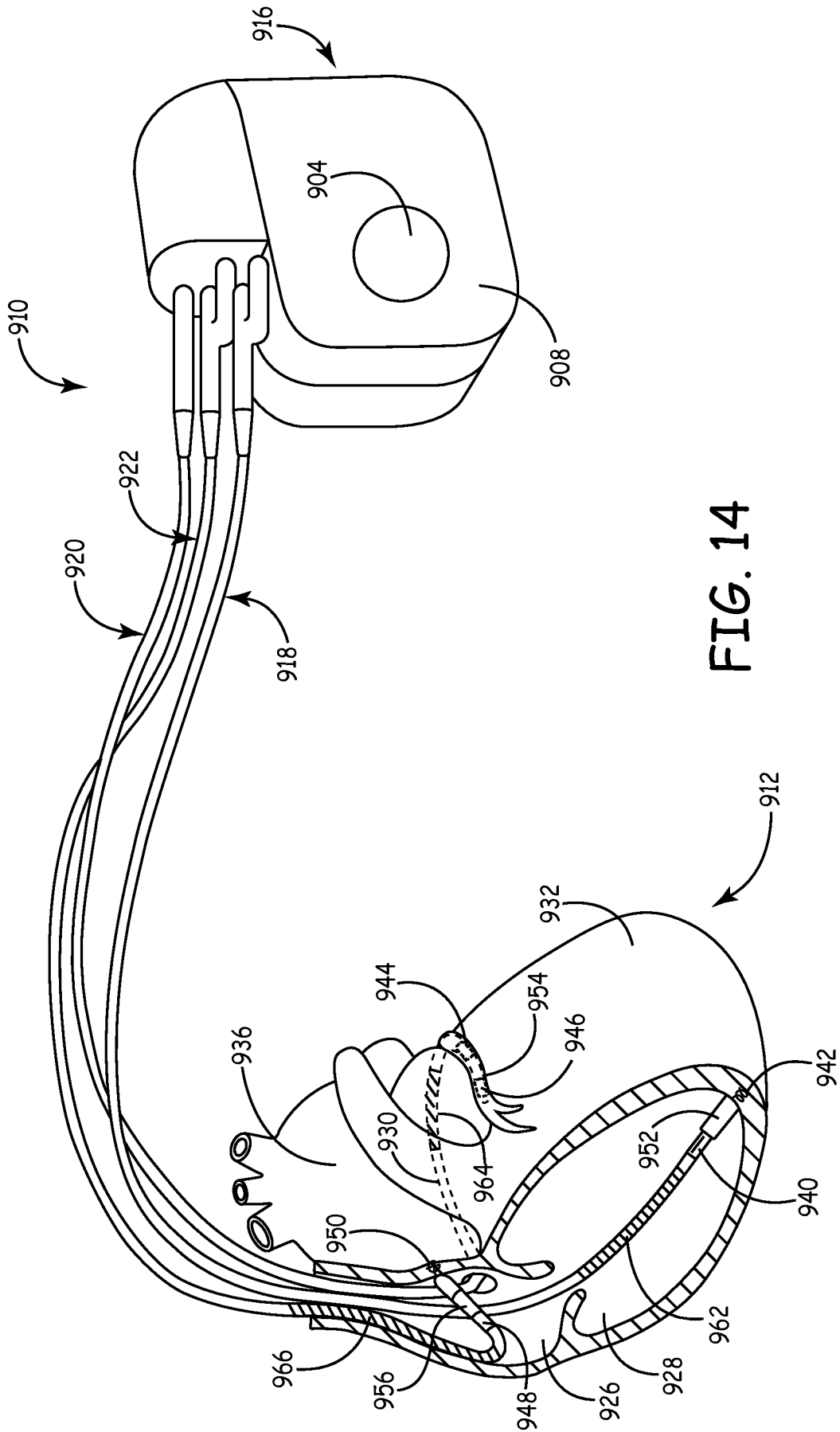


FIG. 14

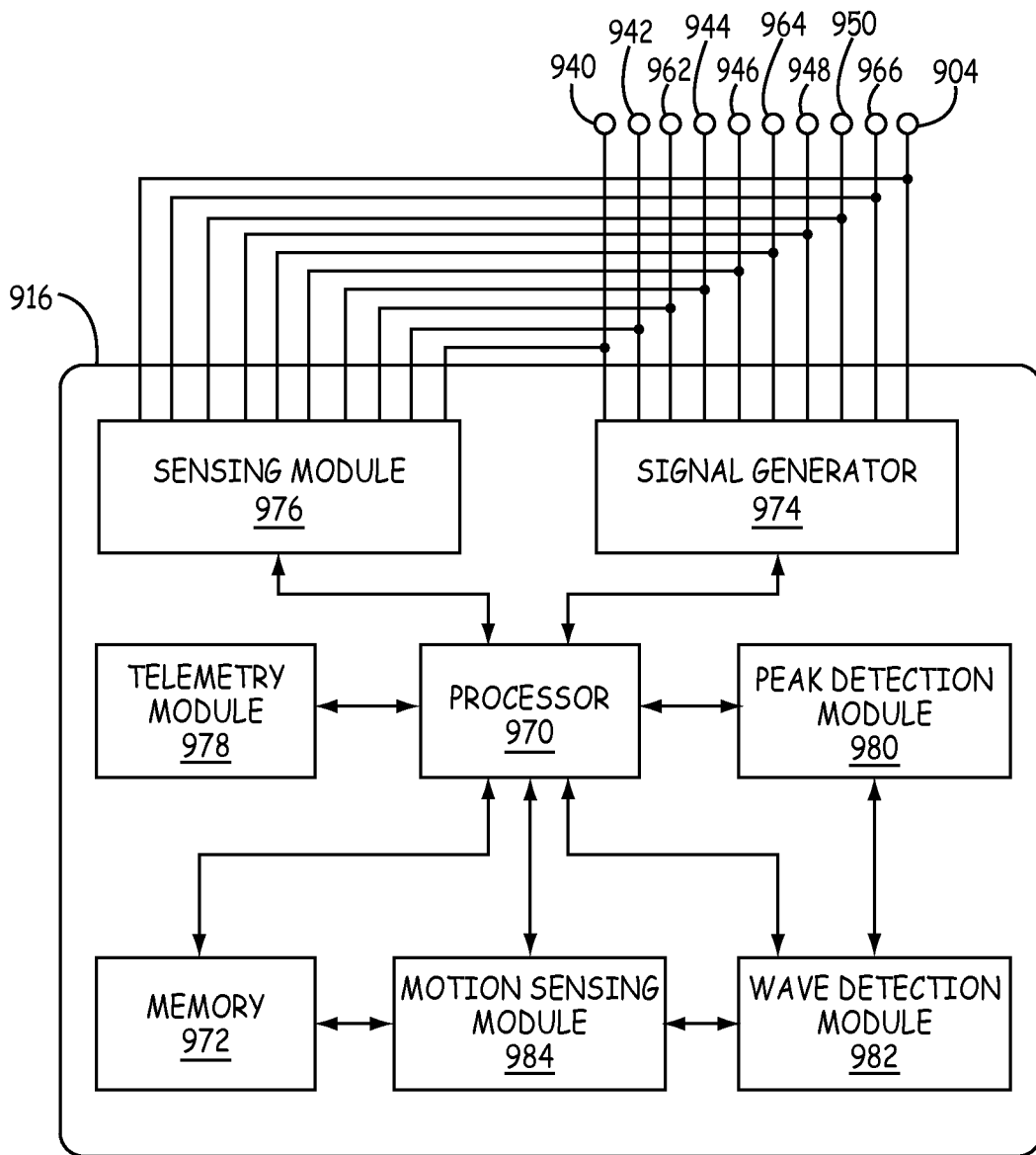


FIG. 15

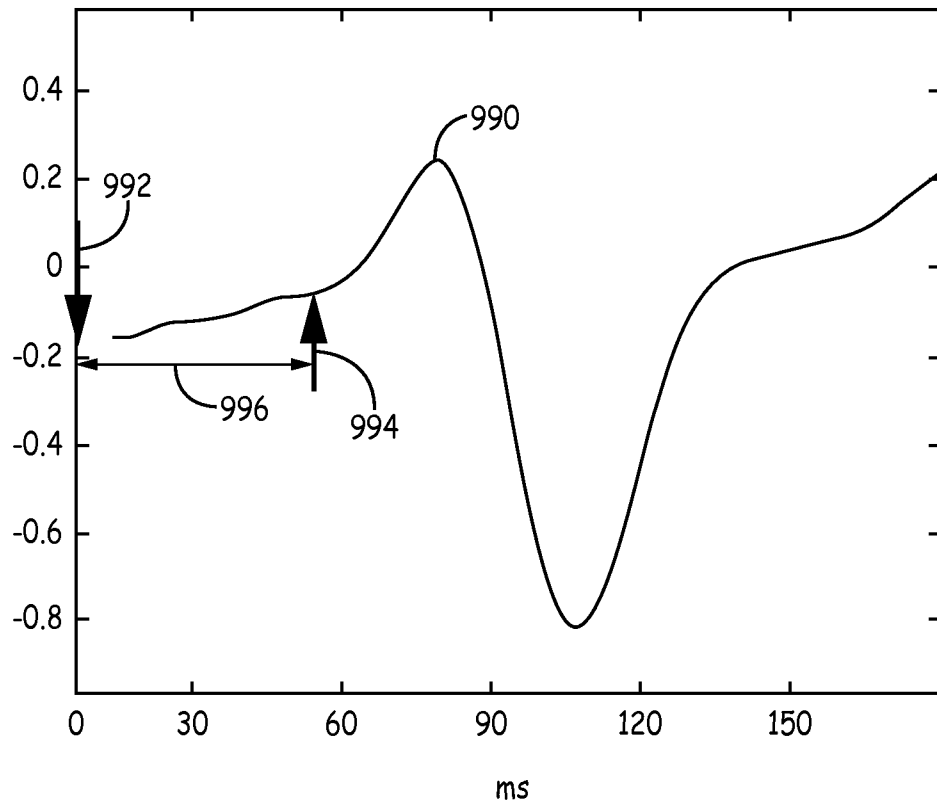


FIG. 16

INTERNATIONAL SEARCH REPORT

International application No
PCT/US2013/062949

A. CLASSIFICATION OF SUBJECT MATTER
INV. A61B5/0452 A61N1/368
ADD.
According to International Patent Classification (IPC) or to both national classification and IPC

B. FIELDS SEARCHED
Minimum documentation searched (classification system followed by classification symbols)
A61B A61N

Documentation searched other than minimum documentation to the extent that such documents are included in the fields searched

Electronic data base consulted during the international search (name of data base and, where practicable, search terms used)
EPO-Internal, WPI Data

C. DOCUMENTS CONSIDERED TO BE RELEVANT

Category*	Citation of document, with indication, where appropriate, of the relevant passages	Relevant to claim No.
X	LAGUNA P ET AL: "AUTOMATIC DETECTION OF WAVE BOUNDARIES IN MULTILEAD ECG SIGNALS: VALIDATION WITH THE CSE DATABASE", COMPUTERS AND BIOMEDICAL RESEARCH, ACADEMIC PRESS, LONDON, GB, vol. 27, 1 January 1994 (1994-01-01), pages 45-60, XP002046491, ISSN: 0010-4809, DOI: 10.1006/CBMR.1994.1006	1-9,15
Y	the whole document	10-13
Y	US 2010/268059 A1 (RYU KYUNGMOO [US] ET AL ROSENBERG STUART [US] ET AL) 21 October 2010 (2010-10-21) paragraphs [0023], [0037], [0075], [0136] - [0147]; figures ----- -/--	10-13

Further documents are listed in the continuation of Box C.

See patent family annex.

* Special categories of cited documents :

- "A" document defining the general state of the art which is not considered to be of particular relevance
- "E" earlier application or patent but published on or after the international filing date
- "L" document which may throw doubts on priority claim(s) or which is cited to establish the publication date of another citation or other special reason (as specified)
- "O" document referring to an oral disclosure, use, exhibition or other means
- "P" document published prior to the international filing date but later than the priority date claimed

"T" later document published after the international filing date or priority date and not in conflict with the application but cited to understand the principle or theory underlying the invention

"X" document of particular relevance; the claimed invention cannot be considered novel or cannot be considered to involve an inventive step when the document is taken alone

"Y" document of particular relevance; the claimed invention cannot be considered to involve an inventive step when the document is combined with one or more other such documents, such combination being obvious to a person skilled in the art

"&" document member of the same patent family

Date of the actual completion of the international search 10 January 2014	Date of mailing of the international search report 20/01/2014
Name and mailing address of the ISA/ European Patent Office, P.B. 5818 Patentlaan 2 NL - 2280 HV Rijswijk Tel. (+31-70) 340-2040, Fax: (+31-70) 340-3016	Authorized officer Küster, Gunilla

INTERNATIONAL SEARCH REPORT

International application No
PCT/US2013/062949

C(Continuation). DOCUMENTS CONSIDERED TO BE RELEVANT		
Category*	Citation of document, with indication, where appropriate, of the relevant passages	Relevant to claim No.
X	<p>WENYU YE ET AL: "Detection of Multilead ECG Character Points and Assessment Based on a Reference Database", ENGINEERING IN MEDICINE AND BIOLOGY SOCIETY, 2005. IEEE-EMBS 2005. 27TH ANNUAL INTERNATIONAL CONFERENCE OF THE SHANGHAI, CHINA 01-04 SEPT. 2005, PISCATAWAY, NJ, USA, IEEE, 1 September 2005 (2005-09-01), pages 3895-3898, XP010906630, DOI: 10.1109/IEMBS.2005.1615312 ISBN: 978-0-7803-8741-6 see section II. D.</p> <p style="text-align: center;">-----</p>	1-8,15
X	<p>LAGUNA P ET AL: "NEW ALGORITHM FOR QT INTERVAL ANALYSIS IN 24-HOUR HOLTER ECG: PERFORMANCE AND APPLICATIONS", MEDICAL AND BIOLOGICAL ENGINEERING AND COMPUTING, SPRINGER, HEILDELBERG, DE, vol. 28, no. 1, 1 January 1990 (1990-01-01), pages 67-73, XP000297223, ISSN: 0140-0118, DOI: 10.1007/BF02441680 the whole document</p> <p style="text-align: center;">-----</p>	1-3,6-9,15
X	<p>FRIESEN G M ET AL: "A COMPARISON OF THE NOISE SENSITIVITY OF NINE QRS DETECTION ALGORITHMS", IEEE TRANSACTIONS ON BIOMEDICAL ENGINEERING, IEEE SERVICE CENTER, PISCATAWAY, NJ, USA, vol. 37, no. 1, 1 January 1990 (1990-01-01), pages 85-98, XP000101197, ISSN: 0018-9294, DOI: 10.1109/10.43620 see algorithm FD1 in section V. D.</p> <p style="text-align: center;">-----</p>	1-5,8,15
X	<p>SO H H ET AL: "Development of QRS detection method for real-time ambulatory cardiac monitor", ENGINEERING IN MEDICINE AND BIOLOGY SOCIETY, 1997. PROCEEDINGS OF THE 19TH ANNUAL INTERNATIONAL CONFERENCE OF THE IEEE CHICAGO, IL, USA 30 OCT.-2 NOV. 1997, PISCATAWAY, NJ, USA, IEEE, US, 30 October 1997 (1997-10-30), pages 289-292, XP032263702, DOI: 10.1109/IEMBS.1997.754529 ISBN: 978-0-7803-4262-0 see section III.</p> <p style="text-align: center;">-----</p>	1-5,8,15
A	<p>US 2003/050565 A1 (COUDERC JEAN-PHILIPPE [US] ET AL) 13 March 2003 (2003-03-13) paragraphs [0005], [0044], [0047], [0057]; figures</p> <p style="text-align: center;">-----</p> <p style="text-align: center;">-/--</p>	1,15

INTERNATIONAL SEARCH REPORT

International application No
PCT/US2013/062949

C(Continuation). DOCUMENTS CONSIDERED TO BE RELEVANT		
Category*	Citation of document, with indication, where appropriate, of the relevant passages	Relevant to claim No.
A	US 2011/196246 A1 (JOHNSON STEVEN R [US]) 11 August 2011 (2011-08-11) paragraph [0048]; figures -----	1,15

INTERNATIONAL SEARCH REPORT

International application No.
PCT/US2013/062949

Box No. II Observations where certain claims were found unsearchable (Continuation of item 2 of first sheet)

This international search report has not been established in respect of certain claims under Article 17(2)(a) for the following reasons:

1. Claims Nos.: 14
because they relate to subject matter not required to be searched by this Authority, namely:
Rule 39.1(iv) PCT - Method for treatment of the human or animal body by therapy
2. Claims Nos.:
because they relate to parts of the international application that do not comply with the prescribed requirements to such an extent that no meaningful international search can be carried out, specifically:
3. Claims Nos.:
because they are dependent claims and are not drafted in accordance with the second and third sentences of Rule 6.4(a).

Box No. III Observations where unity of invention is lacking (Continuation of item 3 of first sheet)

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

1. As all required additional search fees were timely paid by the applicant, this international search report covers all searchable claims.
2. As all searchable claims could be searched without effort justifying an additional fees, this Authority did not invite payment of additional fees.
3. As only some of the required additional search fees were timely paid by the applicant, this international search report covers only those claims for which fees were paid, specifically claims Nos.:
4. No required additional search fees were timely paid by the applicant. Consequently, this international search report is restricted to the invention first mentioned in the claims; it is covered by claims Nos.:

Remark on Protest

- The additional search fees were accompanied by the applicant's protest and, where applicable, the payment of a protest fee.
- The additional search fees were accompanied by the applicant's protest but the applicable protest fee was not paid within the time limit specified in the invitation.
- No protest accompanied the payment of additional search fees.

INTERNATIONAL SEARCH REPORT

Information on patent family members

International application No

PCT/US2013/062949

Patent document cited in search report	Publication date	Patent family member(s)	Publication date	
US 2010268059	A1	21-10-2010	US 2010268059 A1	21-10-2010
			US 2011092809 A1	21-04-2011

US 2003050565	A1	13-03-2003	AU 2002324691 A1	03-03-2003
			US 2003050565 A1	13-03-2003
			US 2006189876 A1	24-08-2006
			WO 03015607 A2	27-02-2003

US 2011196246	A1	11-08-2011	NONE	
