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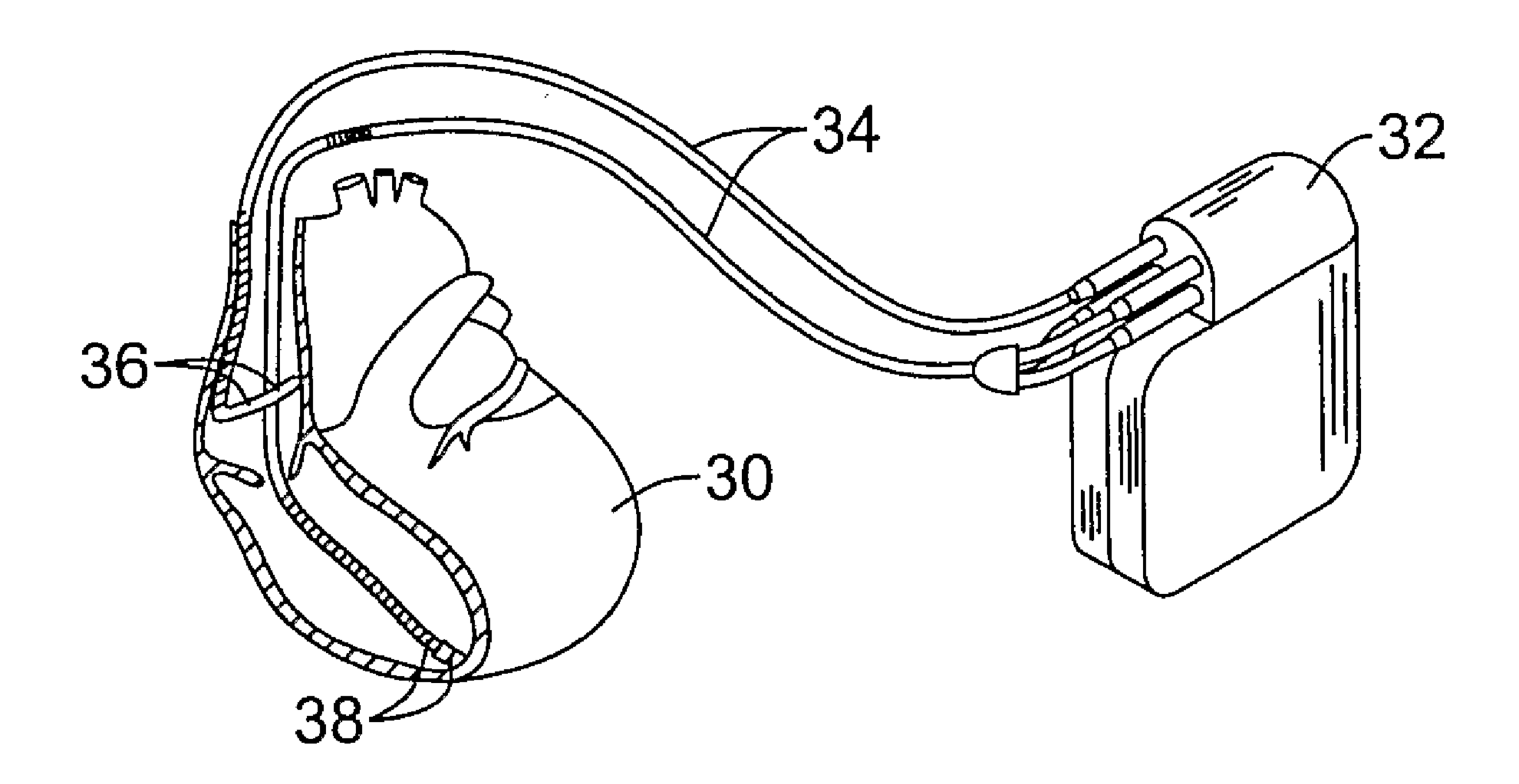
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(54) Title: SIGNAL TEMPLATES IN IMPLANTABLE CARDIAC DEVICES



(57) Abrégé/Abstract:

The present invention is directed toward template formation methods for use in cardiac rhythm management devices. The template formation methods of the present invention provide for creating a robust template to compare with sensed cardiac complexes.





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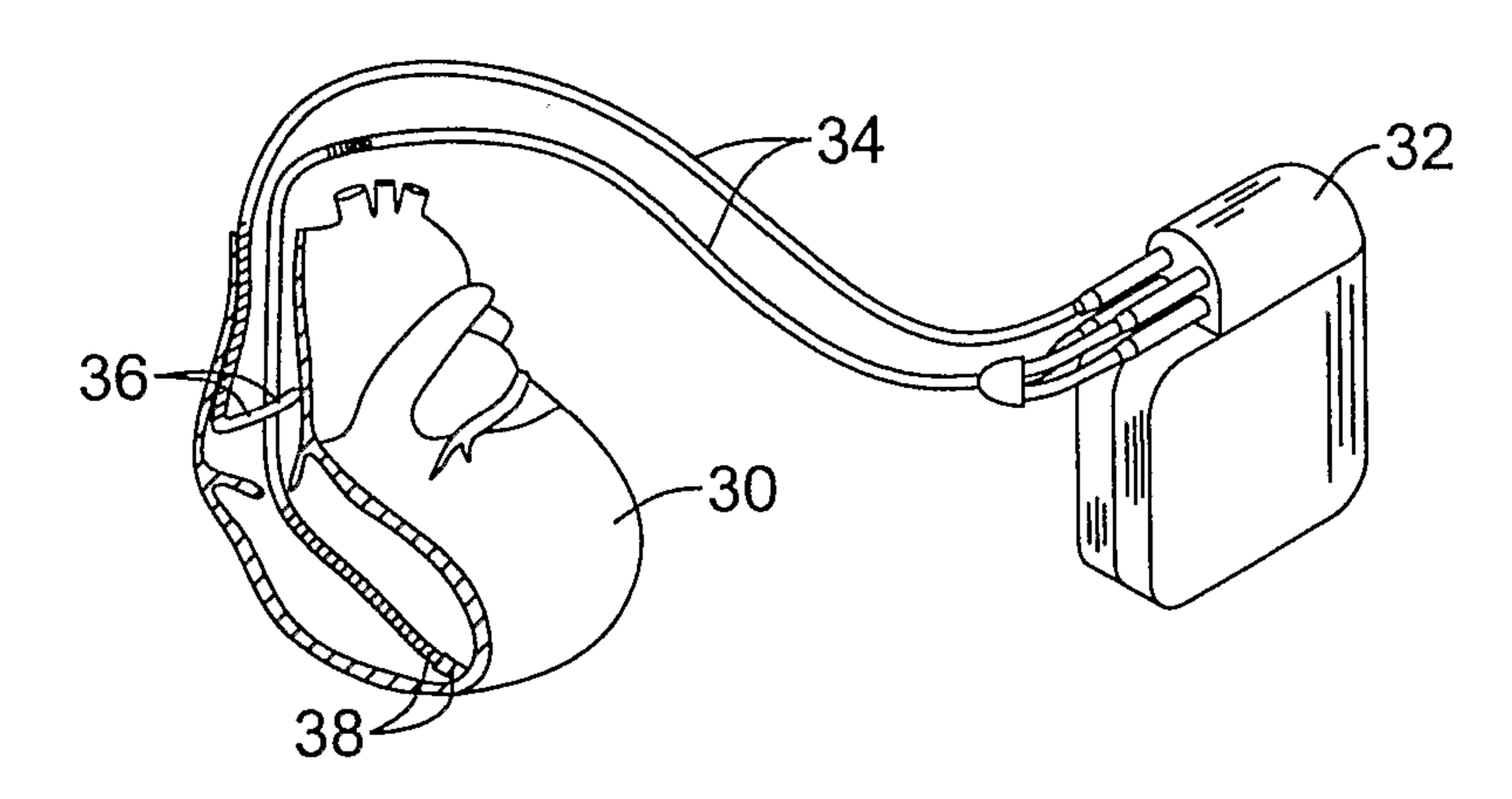
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(54) Title: SIGNAL TEMPLATES IN IMPLANTABLE CARDIAC DEVICES



(57) Abstract: The present invention is directed toward template formation methods for use in cardiac rhythm management devices. The template formation methods of the present invention provide for creating a robust template to compare with sensed cardiac complexes.

SIGNAL TEMPLATES IN IMPLANTABLE CARDIAC DEVICES

Field

The present invention relates generally to implantable cardiac systems that detect, sense and classify cardiac signals. More particularly, the present invention relates to implantable medical devices that generate a template from which the medical device can make comparisons to a patient's normal cardiac complex.

10 <u>Background</u>

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Implantable cardiac rhythm management devices are an effective treatment in managing irregular cardiac rhythms in particular patients. Implantable cardiac rhythm management devices are capable of recognizing and treating arrhythmias with a variety of therapies. To effectively deliver these therapies, however, cardiac rhythm management devices must first accurately sense and classify an episode.

In order to apply the proper therapy in responding to an episode, some cardiac rhythm management devices compare sensed cardiac signals to a previously stored "template" representing normal sinus rhythm (NSR) or other "template" frequently intended to represent the patient's NSR. This stored NSR template must accurately characterize a patient's true NSR in order to be used in a process that properly identifies potentially fatal deviations from normal cardiac activity.

Problems arise when the cardiac rhythm management device inaccurately compares a sensed cardiac complex to a stored NSR template, and as a result, misclassifies the sensed cardiac complex. The severity of this problem escalates if the cardiac rhythm management device inappropriately delivers therapy due to the misclassification. In illustration, when a particular group of sensed complexes are erroneously compared to a stored template because of an improper alignment to the

template, a cardiac rhythm management device may mistakenly classify these sensed complexes as a mismatch and even possibly as a tachyarrhythmia.

For the reasons stated above, and for other reasons stated below, which will become apparent to those skilled in the art upon reading and understanding the present specification, there is a need in the art for providing a reliable system to generate templates for comparison with sensed cardiac events to accurately classify and, if indicated, treat the cardiac rhythm a patient is experiencing.

Summary

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The present invention is directed toward template formation methods for use in cardiac rhythm management devices. The template formation methods of the present invention provide for creating a robust template to compare with sensed cardiac complexes. In an illustrative embodiment, the present invention is used to form templates having a template data set and template alignment parameters for use in aligning captured signals with the template data set prior to comparing the template data set to captured signals.

An illustrative embodiment includes a method of cardiac signal analysis comprising sensing a first cardiac event, configuring template parameters for analysis of the first cardiac event, defining a first sensed signal for the first cardiac event using the template parameters, sensing a second cardiac event, defining a second sensed signal for the second cardiac event using the template parameters, and comparing the second sensed signal to the first sensed signal to determine whether the first sensed signal and template parameters are suitable for defining a cardiac event template. In another embodiment, the illustrative method is performed such that the step of configuring template parameters includes selecting a rule for identifying a fiducial point, wherein the rule is selected from among a set of rules, the rule is selected in light of the characteristics of the first cardiac event, and the rule for identifying a fiducial point becomes one of the template parameters. In a further embodiment, the step of configuring template parameters further includes selecting a number of samples of the first sensed signal around the fiducial point, wherein the configuration of samples around the fiducial point becomes one of the template parameters. The step of selecting a number of samples may include identifying the start and end of a cardiac event. For some embodiments, the set of rules includes an amplitude rule

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related to the relative amplitudes of peaks in the sensed signal, and a location rule related to the location of a peak in the sensed signal.

Another illustrative embodiment includes a method of cardiac signal analysis including forming a template for cardiac event comparisons, the step of forming a template comprising sensing a first cardiac event, identifying a first fiducial point in the first cardiac event using a set of rules, sensing a second cardiac event, identifying a second fiducial point in the second cardiac event using the set of rules, determining whether the first fiducial point and second fiducial point were identified using the same rule, and, if not, discarding the first cardiac event.

In yet another embodiment, a method of cardiac signal analysis comprises sampling a signal using electrodes implanted in a patient's torso for capturing cardiac signals, defining a first sensing window around a first fiducial point to capture a QRS segment, observing the definition of the first sensing window to create template parameters, defining a second sensing window around a second fiducial point using the template parameters, and comparing data in the first sensing window to data in the second sensing window to verify whether to define a valid template using the template parameters.

Another embodiment includes a method of cardiac signal template formation comprising receiving a first cardiac signal from implanted electrodes, selecting a fiducial point in the first cardiac signal, forming a template around the fiducial point, and verifying the template by receiving a second cardiac signal and using the template to compare the second cardiac signal to the first cardiac signal.

Brief Description of the Drawings

Figures 1A-1B illustrate, respectively, representative subcutaneous and intravenous implantable cardiac treatment systems;

Figure 2 depicts a template formation system in accordance with an illustrative embodiment of the present invention;

Figure 3 shows selection of a positive peak of a cardiac complex based on an amplitude rule of a fiducial point selection process;

Figure 4 shows selection of a negative peak of a cardiac complex based on an amplitude rule of a fiducial point selection process;

Figure 5 shows selection of a positive peak of a cardiac complex based on a location rule of a fiducial point selection process;

Figure 6 shows selection of a negative peak of a cardiac complex based on a location rule of a fiducial point selection process;

Figure 7 depicts a cardiac signal possessing a notch in the QRS segment;

Figure 8 shows a pre-template template window;

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Figure 9 shows the pre-template template window depicted in Figure 8 after the monotonic segments are identified in the cardiac complex;

Figure 10 depicts a cardiac signal having a notch within the cardiac signal's ORS segment;

Figure 11 depicts the template window for a patient having a wide QRS;

Figure 12 shows a cardiac complex having a QRS segment that is capable of having its pre-template template window narrowed by masking;

Figure 13 depicts the observed template window after adjusting the template window's bounds;

Figure 14 shows a cardiac complex having a QRS segment width that is smaller than the acceptable minimum template window;

Figure 15 illustrates a QRS segment that was not properly captured through the pre-template template window formation process;

Figure 16 shows the result of an offset adjustment process to the QRS segment captured in Figure 15;

Figure 17 illustrates a template verification process;

Figures 18A-18C further illustrate a template verification step; and

Figure 19 is a block diagram for an illustrative template formation process.

Detailed Description of the Drawings

The following detailed description should be read with reference to the drawings, in which like elements in different drawings are numbered identically. The drawings, which are not necessarily to scale, depict selected embodiments and are not intended to limit the scope of the invention. Those skilled in the art will recognize that many of the examples provided have suitable alternatives that may be utilized.

The present invention is generally related to implantable cardiac treatment systems that provide therapy for patients who are experiencing particular arrhythmias. The present invention is directed toward detection architectures for use in cardiac rhythm devices. In particular, the present invention is suited for implantable cardiac treatment systems capable of detecting and treating harmful arrhythmias. Although

the detection architecture is intended primarily for use in an implantable medical device that provides defibrillation therapy, the invention is also applicable to cardiac rhythm devices (including external devices) directed toward anti-tachyarrhythmia pacing (ATP) therapy, pacing or other cardiac stimulation techniques, and other cardiac rhythm devices capable of performing a combination of therapies to treat rhythm disorders.

To date, implantable cardiac treatment systems have been either epicardial systems or transvenous systems. For example, transvenous systems can be implanted generally as shown in Figure 1B. However, as further explained herein, the present invention is also adapted to function with a subcutaneous implantable cardiac treatment system as shown in Figure 1A.

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Figure 1A illustrates a subcutaneously placed implantable cardiac treatment system, in particular, an implantable cardioverter/defibrillator (ICD) system. In this illustrative embodiment, the heart 10 is monitored using a canister 12 coupled to a lead system 14. The canister 12 may include an electrode 16 thereon, while the lead system 14 connects to sensing electrodes 18, 20, and a coil electrode 22 that may serve as a shock or stimulus delivery electrode as well as a sensing electrode. The various electrodes define a number of sensing vectors V1, V2, V3, V4. It can be seen that each vector provides a different vector "view" of the heart's 10 electrical activity. The system may be implanted subcutaneously as illustrated, for example, in U.S. Patents No. 6,647,292 and 6,721,597. By subcutaneous placement, it is meant that electrode placement does not require insertion of an electrode into a heart chamber, in or on the heart muscle, or the patient's vasculature.

Figure 1B illustrates a transvenous ICD system. The heart 30 is monitored and treated by a system including a canister 32 coupled to a lead system 34 including atrial electrodes 36 and ventricular electrodes 38. A number of configurations for the electrodes may be used, including placement within the heart, adherence to the heart, or disposition within the patient's vasculature.

Figure 2 depicts a template formation system 40 in accordance with an illustrative embodiment of the present invention. The template formation system 40 can be used to create and store multiple static and/or dynamic templates. Static templates are cardiac complexes that are captured previously in time and stored for

reference by the device. Alternatively, dynamic templates are cardiac complexes that are continuously or periodically captured and/or updated.

The template formation system 40 of the present invention generally comprises a multi-stage data analysis – signal collection 42, fiducial point selection 44, pre-template formation 46, template optimization 48, and template verification 50. Sections of the multi-stage data analysis, however, may operate autonomously, as will be discussed in detail below. As such, a particular process in the template formation system 40 may be bypassed or may function independently in the device's overall detection architecture.

The system, in an illustrative embodiment, not only identifies NSR signals for comparison to sensed events, it also defines and re-defines the sensing parameters (for example, fiducial point selection, window size, and/or window/fiducial point alignment). These signals and parameters can then be used for making comparisons with a sensed cardiac signal to determine whether the signal is NSR.

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The processes within the template formation system 40 may additionally create or modify a template to accommodate for morphological changes in the patient's cardiac complex. For example, the ultimately formed template may be continually updated to adapt to certain morphological changes in the sensed cardiac complex. As such, the template formation system 40 of the present invention is adaptive and this adaptive characteristic may be automated.

The template formation system 40 is initiated by collecting a cardiac signal 42. The cardiac signal may be collected using any suitable capture method. This sensed cardiac complex is then processed for proper alignment. A method for repetitive and reliable alignment of a collected signal enhances the accuracy when comparing sensed signals to a stored template. In some embodiments, the step of collecting the cardiac signal 42 may include a signal certification process such as that illustrated in copending U.S. Patent No. 7,248,921, filed June 1, 2004 and entitled METHOD AND DEVICES FOR PERFORMING CARDIAC WAVEFORM APPRAISAL.

In several embodiments of the present invention, a fiducial point for alignment is generally established using a preferred peak of the sensed cardiac complex. The fiducial point may be selected manually for each patient, or alternatively, the fiducial point may be selected using a rule-based method. In preferred embodiments, the fiducial point is selected by analyzing the repetitive nature of peaks on 'n'

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consecutive complexes. In one embodiment of the present invention, the fiducial point selection process 44 is based on the results of the most recently sensed cardiac complex and the three (3) cardiac complexes previous to the most recently sensed complex. Alternative embodiments may base the fiducial point selection process 44 on the repetitive nature of as many as 20 consecutive complexes to as few as an ongoing beat to beat determination.

A preferred fiducial point selection process 44 implements a set of rules to choose the most appropriate peak for alignment in a cardiac complex. In preferred embodiments, the fiducial point selection process 44 is based on an amplitude rule and a location rule. Additionally, due to the nature of the fiducial point selection process 44 rules, while the R-wave will often be chosen since it is frequently associated as the most striking phase deflection observed in a cardiac complex, the R-wave is not necessarily selected as the fiducial point for alignment in any given cardiac complex.

The first rule used by the illustrative fiducial point selection process 44 is the amplitude rule. This rule sets the fiducial point on the peak (either positive or negative) of the QRS cardiac complex having the greatest relative amplitude. The amplitude rule is set forth as:

- If the positive peak amplitude > 2 times the negative peak amplitude, then fiducial point selection is on the peak of the positive phase deflection "positive amplitude";
- If the negative peak amplitude > 2 times the positive peak amplitude, then fiducial point selection is on the peak of the negative phase deflection "negative amplitude";
- If neither the positive peak nor the negative peak satisfies the amplitude rule, then the location rule, set forth below, controls.

The relative amplitudes for the positive and negative peaks are measured from the patient's isoelectric line 52 – illustrated in Figure 3. The isoelectric line represents a signal lacking significant detected phase deflection, i.e. a detected signal level that does not indicate cardiac activity and provides a baseline for signal analysis. The fiducial point selection process 44 then determines the largest positive and negative phase deflections from the isoelectric line 52. In the present example, the amplitude of the largest positive phase deflection is shown as 54. Similarly, the amplitude of the largest negative phase deflection is shown as 56. The relative

amplitudes of both the positive phase deflection 54 and negative phase deflection 56 are then assessed. If the relative amplitude of the positive phase deflection is greater than two times the relative amplitude of the negative phase deflection, the fiducial point selection is suggested to be on the positive amplitude peak.

In the present example, the fiducial point selection process is established by the repetitive nature of four consecutive cardiac complexes. Cardiac complexes 58, 60, 62 and 64 each demonstrate a positive peak amplitude greater than two times (2X) its corresponding negative peak amplitude. After the fourth consecutive cardiac complex 64, the fiducial point selection process establishes the positive peak as the fiducial point for alignment based on the amplitude rule. The triangles shown in Figure 3 represent points where the amplitude rule has been met in four consecutive complexes. Additionally, each triangle signifies an established fiducial point for template alignment.

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Figure 4 shows fiducial point selection of a negative peak based on the amplitude rule. In the present example, the fiducial point selection process is established by the repetitive nature of a sensed cardiac complex and the previous three sensed cardiac complexes (four consecutive cardiac complexes). Cardiac complexes 68, 70, 72 and 74 each demonstrate a negative peak amplitude greater than two times (2X) its corresponding positive peak amplitude. Specifically, the amplitude of the largest negative phase deflection 56 is assessed to be two times the relative amplitudes of the positive phase deflection 54. After the fourth consecutive cardiac complex 74, the fiducial point selection process establishes the negative peak as the fiducial point for alignment based on the amplitude rule. The triangles shown in Figure 4 represent a point where the amplitude rule has been met for four consecutive complexes. Additionally, each triangle signifies an established fiducial point for template alignment.

The second rule used by the illustrative peak alignment process is the location rule. This rule is premised on setting the fiducial point on the peak of the first significant phase deflection (either positive or negative) occurring in time within the ventricular cardiac complex. In certain embodiments, the location rule is considered when the amplitude rule cannot be established. Alternate embodiments utilize the location rule without deference to the amplitude rule. The location rule is set forth as:

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- If a significant positive phase deflection precedes a significant negative phase deflection in a cardiac complex, then fiducial point selection is on the peak of the positive phase deflection "positive location";
- If a significant negative phase deflection precedes a significant positive phase deflection in a cardiac complex, then fiducial point selection is on the peak of the negative phase deflection "negative location".

Figure 5 shows an illustrative fiducial point selection of a positive peak based on the location rule. In the present example, the fiducial point selection process is established by the repetitive nature of four consecutive cardiac complexes. Cardiac complexes 78, 80, 82 and 84 each show a significant positive phase deflection before a significant negative phase deflection in the cardiac complex. After the fourth consecutive cardiac complex 84, the fiducial point selection process establishes the peak of the positive phase deflection as the fiducial point for alignment based on the location rule. The triangles shown in Figure 5 represent points where the location rule has been met in four consecutive complexes. Additionally, each triangle signifies an established fiducial point for template alignment.

Figure 6 shows an illustrative fiducial point selection of a negative peak based on the location rule. The fiducial point selection process in Figure 6 is established by the repetitive nature of four consecutive cardiac complexes. Cardiac complexes 88, 90, 92 and 94 each show a significant negative phase deflection before a significant positive phase deflection in the cardiac complex. After the fourth consecutive cardiac complex 94, the fiducial point selection process establishes the peak of the negative phase deflection as the fiducial point for alignment based on the location rule. The triangles shown in Figure 6 represent a point where the location rule has been met for four consecutive complexes. Additionally, each triangle signifies an established fiducial point for template alignment.

In certain embodiments where the fiducial point selection process requires more than one cardiac complex to establish a fiducial point, the process may require each of the cardiac complexes assessed to adhere to the same rule (amplitude or location) before establishing a fiducial point for alignment. More particularly, each cardiac complex analyzed and used for establishing a fiducial point must adhere to the

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same one of the four possible rule bases: positive amplitude, negative amplitude, positive location, or negative amplitude.

In alternative embodiments, the fiducial point selection process may require all of the cardiac complexes assessed to establish the same fiducial point (i.e., the same positive peak) regardless of which rule was used. In an illustrative embodiment, the fiducial point selection process is established by the repetitive nature of three consecutive cardiac complexes. Two of the three cardiac complexes may establish the fiducial point on the positive peak using the positive amplitude rule base. The remaining cardiac complex may establish the same fiducial point on the complex's positive peak, however, using the positive location rule and not the amplitude rule. While not using the same rule, all three cardiac complexes indicate the same fiducial point, and as such, are indicated for submission to template verification, as referred to in Figure 2.

In certain circumstances, a notch is observed in the cardiac signal's QRS segment. Figure 7 depicts a cardiac signal possessing a notch 96 in the QRS segment. A notch in a cardiac signal normally fails to affect the fiducial point selection process. This is observed because a predominant peak usually exists amongst the peaks forming the notch. Thus, the fiducial point selection process will generally select the predominant peak. In instances where one peak does not stand out over the other peak (as is depicted in Figure 7), or when the predominant peak frequently changes from cardiac complex to cardiac complex, a potential for confusion in the fiducial point selection process may arise. In embodiments possessing such problematic notch segments, a notch analysis process may be used to assure the proper fiducial point selection for alignment.

In an illustrative notch analysis process, a notch is presumed to exist if the distance (in time) between the two peaks 98 is more than approximately 20 msec and/or if the difference in peak amplitudes 100 is less than approximately 115 μ V. These values may vary in several embodiments depending upon the placement and design of sensing electrodes, as well as the expected characteristics of notched QRS peaks for a given patient. If these conditions are not met, it is presumed that a predominant peak does exist, that the fiducial point selection process will identify the predominant peak, and so the illustrative notch analysis process is skipped. However, if these conditions are met, then the cardiac signal is presumed to possess a notch requiring further analysis for proper fiducial point selection.

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The illustrative notch analysis process identifies the peaks in the signal, and determines which peak has been initially identified as the fiducial point. If the first peak occurring in time is identified as the fiducial point, then the notch analysis is complete. If the second peak occurring in time is identified as the fiducial point, then the notch analysis process forces the fiducial point onto the first peak of the notch occurring in time.

Once the fiducial point is selected, the pre-template is then formed. Figure 8 is an illustrative embodiment of a pre-template 102. The pre-template 102 is populated with a number of samples taken at a sampling frequency which form a pre-template data set. In the illustrative embodiment, the disposition of the pre-template data set within the pre-template is determined by template alignment parameters including the fiducial point selection explained above and the placement and masking steps further discussed below.

In an illustrative embodiment, the fiducial point 104 is placed at the center of the pre-template 102. In preferred embodiments, a number of samples 'n' are established to the left of the fiducial point 104, and 'n' samples are also established to the right of the fiducial point 104. For example, some embodiments of the present invention utilize forty-one (41) samples sampled at 256 Hz, corresponding to approximately 160 msec. In an illustrative embodiment, twenty (20) samples are established to the left of the fiducial point 104 and another twenty (20) samples are established to the right of the fiducial point 104. The forty-one (41) samples form a pre-template window 106 in which the relevant portion of a cardiac signal will be analyzed. In alternative embodiments, the number of samples 'n' populated on either side of the center of the pre-template 102 may differ.

From this initially formed pre-template window 106, the boundaries of the cardiac complex's QRS segment are sought. Figure 8 shows a pre-template window 106 that includes a QRS segment as well as extraneous portions of a sensed cardiac signal. In this instance, it is desired to optimize the formed pre-template 102 by narrowing the pre-template window 106 to comprise mostly the QRS segment and reduce extraneous portions of the cardiac signal. The first step in this process is to identify the beginning and end of the QRS segment.

In one embodiment of the present invention, the observation of monotonic segments is used to estimate the beginning and end of the QRS segment. A monotonic segment is a signal segment of consecutive samples in which the sensed

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amplitude changes in the same direction or stays the same. For example, a series of consecutive samples in which each successive sample is greater than or equal to (in amplitude) the previous sample would be an increasing monotonic segment. Similarly, a series of consecutive samples in which each successive sample is less than or equal to (in amplitude) the previous sample would be a decreasing monotonic segment. One method for observing monotonic segments is by determining the zero crossing points of the first derivative of the cardiac complex signal.

In this embodiment, an arithmetic operation is performed on the initial pretemplate 102 to identify the cardiac complex's monotonic segments — as indicated by the zero crossing points of the first derivative of the cardiac complex signal. Figure 9 shows the pre-template window 106 depicted in Figure 8 after all of the monotonic segments are identified in the cardiac complex. Each diamond indicates the beginning/end of a monotonic segment. An arithmetic operation then identifies the largest monotonic segment (in terms of change of amplitude) in the initial pre-template 102 before the fiducial point 104. This sample is noted as "QRS begin" 108. The arithmetic operation further identifies the largest monotonic segment (in terms of change of amplitude) in the initial pre-template 102 after the fiducial point 104. This sample is noted as "QRS end" 110. QRS begin and QRS end estimate the boundaries for the cardiac complex's QRS segment in this embodiment.

The use of monotonic segments is further useful for eliminating errors in calculating QRS segment length with cardiac complexes having a notch in their QRS segment. Figure 10 illustrates a cardiac complex possessing a notch. Since the arithmetic operation of the illustrative embodiment identifies the largest monotonic segment (in amplitude) in the initial pre-template 102 before the fiducial point 104 and after the fiducial point 104, most notches will not affect the algorithm's ability to find the desired QRS begin and QRS end. As depicted in Figure 10, the relative monotonic segment amplitudes within the notch are smaller than the amplitudes of the monotonic segments at either end of the QRS segment. Therefore, the notch generally does not affect the estimated measurement of the QRS segment.

Alternative methods known in the art may also be utilized to estimate the beginning and end of the cardiac complex's QRS segment. The use of monotonic segments to estimate the QRS segment is merely illustrative, and various embodiments of present invention are not limited to this particular aspect of the illustrative embodiment.

After the QRS segment has been identified, the pre-template 102 is then optimized for performance – process 46 in Figure 2. Optimization includes, but is not limited to, masking the pre-template window 106 to include the most relevant samples in the cardiac complex, as well as offset adjustment.

One method for template optimization is to narrow or mask the pre-template window 106 to include only those samples indicative of the QRS segment. In patients with wide QRS segments, optimization by removal of some samples is not indicated. For example, in the above illustrative embodiment, if a patient has a QRS segment longer than 160 msec (or 41 samples), the patient's QRS segment exceeds the initially formed pre-template window 106. Thus, the patient's identified QRS begin 108 is the first sample within the pre-template window 106 and the identified QRS end 110 is the last sample within the pre-template window 106, even though the patient's actual QRS segment may extend beyond the confines of the formed pre-template window 106. An example of a wide QRS segment that exceeds the size of the pre-template window 106 is shown in Figure 11. Masking the pre-template window 106 is not indicated in these instances.

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In contrast, the pre-template window 106 may be masked when the QRS segment is less than the pre-template's window 106. For example, and as depicted in Figure 12, suppose a patient's QRS begin 108 is at the fourth sample within the pretemplate window 106. Similarly, suppose the patient's QRS end 110 occurs on the thirty-fifth sample within the pre-template window 106. Thus, the patient's QRS segment is thirty-two (32) samples long. The other nine (9) samples included in the original pre-template window 106 are generally not useful for analysis, and may introduce undesired effects if included in the final template. Therefore, the bounds of the pre-template 102 may be masked to form a masked pre-template window 114 that only includes the actual QRS segment – between QRS begin 108 and QRS end 110. In this example, the pre-template window 106 would be masked to the 32 samples representing the estimated QRS segment. Specifically, the pre-template bounds are masked so that the masked pretemplate window 114 begins on sample 4 and ends on sample 35, thereby eliminating extraneous samples 112 from the masked pre-template window 114. Figure 13 depicts the observed masked pre-template window 114 after the masking process. Such narrowing or masking, while useful in some embodiments, is not required by the present invention.

If desired, a minimum duration for the masked pre-template window 114 may be defined. In one embodiment of the present invention, the minimum masked pre-template window 114 is approximately 100 msec (25 samples at 256 Hz). In patients having narrow QRS segments (less than approximately 100 msec), the allowable masked pre-template window 114 may still include some extraneous samples with the QRS segment for these patients. For example, as depicted in Figure 14, if the QRS begin 108 is on sample twelve (12) and the QRS end 110 occurs on sample twenty-nine (29), then the width of the QRS segment is eighteen (18) samples. This QRS segment width is smaller than the illustrative minimum for the masked pre-template window 114 of twenty-five (25) samples. To mask the QRS segment to the minimum boundary allowable, the difference is first calculated between the masked pre-template window's minimum (25 samples) and the estimated QRS segment width (eighteen (18) samples in this example). This difference is seven (7) samples. The difference is then split in half and added equally (or as equally as possible) to both sides of the estimated QRS segment length. Thus, the optimized masked pre-template

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following the QRS end 110.

There are other instances where the pre-template window 106 does not include the complete QRS segment. An example of such an instance is when the sample indicating the QRS begin 108 or QRS end 110 occurs on the first or last sample within the initially formed pre-template window 106. In some embodiments, this gives rise to an assumption that the actual QRS begin 108 or QRS end 110 is not accurately captured and that the actual QRS begin 108 or QRS end 110 occurs sometime outside the boundaries of the initially formed pre-template window 106. An example of a pre-template window 106 where the last sample within the pre-template window is also the QRS end 110 is depicted in Figure 15.

window 114 in this example would include the actual QRS segment 116 with three (3)

additional samples preceding the QRS begin 108 and four (4) additional samples

In Figure 15, a pre-template window 106 is populated with samples 1 through 41. The first sample appears near the vertical axis midpoint of the template window. In contrast, the last sample (sample 41) appears near the bottom of the template window's vertical axis. As the samples move along the horizontal axis from sample 1, the samples gradually increase in height until reaching the QRS begin 108 for the cardiac complex segment. The remainder of the pre-template window 106 contains most, but not all, of the QRS segment. The remainder of the QRS segment not

captured within the boundaries of the pre-template window 106 is shown as 116. In such an instance, the entire QRS segment was not properly captured through the pre-template window 106 formation process. Some embodiments of the present invention resolve this issue through an offset adjustment.

The offset adjustment process first identifies which side of the QRS segment was not properly captured. As described above and depicted in Figure 15, the QRS begin 108 is sample eight (8) and the perceived QRS end 111 is sample forty-one (41). This generally indicates that the true QRS end 110 actually occurs at a point later in time and was not captured using the initial settings for forming the pretemplate window 106. When it is indicated that the true QRS end 100 was not properly captured, a number of samples will precede the QRS begin 108. These leading samples are called the "residue" 118. In Figure 15, the residue consists of the first seven (7) samples preceding the QRS begin 108. Since the samples constituting the residue 118 relay little information regarding the QRS segment itself, these samples may be discarded and replaced by samples that do represent the QRS segment but which were omitted through the initial pre-template window formation process. The process for shifting the pre-template window 106 in one direction is called offset. The effect of the offset process, in the present example, is to allow the pre-template window 106 to start 'n' number of residue samples later to ensure that the true QRS end 110 is captured.

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In a preferred embodiment, the sample representing the QRS begin 108 plus the immediately preceding sample (QRS begin -1), or the QRS end 110 plus the immediately following sample (QRS end +1), along with the samples therebetween, are retained. The remaining samples comprise the residue 118. In alternative embodiments, the QRS begin 108 or the QRS end 110, plus some 'n' number of samples preceding or following, is retained and the remaining samples comprises the residue 118. In yet alternative embodiments, just the QRS begin 108 or QRS end 110 is kept and the remaining samples are considered residue.

Figure 16 illustrates the offset process on the cardiac complex depicted in Figure 15. Specifically, Figure 16 depicts the formation of an offset pre-template window 120 to recapture the cardiac complex's true QRS end 110. As described above, Figure 15 shows that there are eight (8) residue samples 118 leading the QRS begin 108. These residue samples 118 are eliminated and QRS begin 108 is forced to be the first sample in a newly formed offset pre-template window 120. This

adjustment is graphically depicted in Figure 16. Thus, the offset pre-template window 120 starts at the QRS begin 108 and now ends eight (8) samples later than it initially did when the pre-template window 106 was initially formed. The result of this shift permits the newly formed offset pre-template window 120 to recapture the cardiac complex's true QRS end 110. Thus, the offset pre-template window 120 comprises the entire QRS segment including both the true QRS begin 108 and the true QRS end 110.

In preferred embodiments, after the offset adjustment process, the corrected template window is further optimized by masking the bounds of the offset template — as described above.

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The parameters used in defining the optimized pre-template window are, in an illustrative example, described as the template parameters. The template parameters describe how the template data set is defined and aligned within the template. These parameters, including the manner of fiducial point selection, offset (if any) and masking (if any) provide template parameters indicating how the template can be used in making future comparisons. The template parameters may be used as described in copending U.S. Patent No. 7,477,935, filed on November 29, 2004, entitled METHOD AND APPARATUS FOR BEAT ALIGNMENT AND COMPARISON, which is filed on even date herewith, is assigned to the assignee of the present invention.

However, in the illustrative embodiment, prior to using the template (including its associated template parameters and template data set) for future comparisons to sensed signals, the template data set is verified for validity.

Once the pre-template is optimized by defining its sample window characteristics, including but not limited to masking and offset adjustment, the data in optimized pre-template is verified for its validity – process 50 in Figure 2. The verification of optimized pre-template validity provides a check on both the template parameters and the template data set. In preferred embodiments, validity must be established before the optimized pre-template is stored as the final template, or as one template among several for use in comparing to subsequently sensed cardiac signals. Figure 17 illustrates the template verification process 50 for an optimized pre-template.

The optimized pre-template 130 is initially stored in a buffer. The device then senses a subsequent cardiac complex 132 using the optimized parameters set for the

optimized pre-template 130. Cardiac complex 132 is then compared to the stored optimized pre-template 130. In a preferred embodiment, an arithmetic operation similar to correlation is performed to determine the similarity between 130 and 132. An illustrative arithmetic operation includes correlation waveform analysis, which returns a result between -1 and 1, and which can be scaled using a number of linear, non-linear, and hybrid scaling methods as noted in co-pending U.S. Patent No. 7,330,757 filed May 27, 2004 and entitled METHOD FOR DISCRIMINATING BETWEEN VENTRICULAR AND SUPRAVENTRICULAR ARRHYTHMIAS.

In an illustrative embodiment, a correlation waveform analysis is performed and then scaled to a percentage value between 0-100%, with negative correlations given a 0%, and positive scores linearly scaled between 0-100%. If the similarity score between the subsequent cardiac complex 132 and the optimized pre-template 130 is greater than a specified threshold, the subsequent cardiac complex 132 is averaged with the optimized pre-template 130. In certain embodiments of the present invention, the threshold for comparison is specified at 80%. Alternative threshold levels may be set without deviating from the spirit and scope of the invention. Additionally, in certain embodiments, the cardiac complex that is compared to the optimized pre-template 130 is not averaged after comparison. If the similarity score does not surpass the specified threshold, then the optimized pre-template 130 is discarded and the entire template formation process is restarted.

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In certain embodiments, if the comparison threshold value is exceeded, then the verification process is repeated with another incoming cardiac complex, for example cardiac complexes 134, 136 and 138. The device captures the cardiac complex 134 using the parameters set for the averaged optimized pre-template (130+132) and performs a further comparison between the cardiac complex 134 and the averaged optimized pre-template (130+132). Again, alternative embodiments may compare the newly sensed cardiac complex 134 to the initially stored optimized pre-template 130. In the present illustrative example, the comparison score between the cardiac complex 134 and the averaged optimized pre-template (130+132) is 85%. Since this score is greater than the comparison threshold of 80%, the verification process is continued.

The verification process is repeated at least this one additional time in some embodiments of the present invention. In a preferred embodiment, this process is

iterated until four (4) consecutive cardiac complexes exceed the threshold level for comparison with either the initially stored optimized pre-template 130, or the averaged optimized pre-template (130+132+134+136). If at any time during the process the similarity score does not surpass the specified threshold, then the optimized pre-template is discarded and the template formation process is restarted in its entirety until a verified template is created.

The template is verified after completing the specified number of iterations for the verification process. In the present illustrative embodiment, the comparison scores to the averaged optimized pre-template for cardiac complexes 132, 134, 136 and 138 were 85%, 89%, 84% and 84%, respectively. Each of these comparison scores exceeded the comparison threshold set for the present example. Thus, the optimized pre-template is verified and the pre-template is considered the final template, thereby completing the template formation process. The formed template can then be used to observe and characterize incoming sensed cardiac signals.

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Figures 18A-18C further illustrate a template verification step. As shown in Figure 18A, a sampled signal is placed into a pre-template template having a fiducial point which is defined using the amplitude rule. The fiducial point is placed as sample s21, with 20 samples on either side making up the pre-template template window. QRS start and end points are identified at s10 and s33, respectively. The signal and its parameters are referred to as an optimized pre-template. Next, the signal is masked using QRS +/- 1 rules, such that the optimized pre-template is as shown by the box, extending from sample s9 to s34. The optimized pre-template is then stored until verified.

Turning to Figure 18B, another sampled signal is captured and the optimized pre-template parameters from Figure 18A are used to define the signal window. In particular, the amplitude rule is used to select a fiducial point and place it at sample s21, and the sample is masked to only include samples s9 to s34. As can be seen, the captured QRS segment in Figure 19B is not accurately masked, as the QRS signal ends at s35, outside the signal window, and the QRS start occurs one sample later than would be desired. However, the overall shape generally resembles that of Figure 18A, and a correlation of the two signals could be calculated to yield a score above a defined threshold such as 0.8 or 80% correlation. Thus the signal in Figure 18B could be retained for averaging with that of Figure 18A to further characterize the optimized pre-template. Alternatively, the data may not be averaged and the signal in the

optimized pre-template of Figure 18A used in further analysis. In another alternative, the verification provided by the signal in Figure 18B could be defined as sufficient to store that of Figure 18A as a template for comparison.

Turning to Figure 18C, a third sampled signal is captured for comparison to the signal in 18A. The first step here is to identify the fiducial point. However, it can be seen that there are two positive peaks X and Y which are near one another. Neither peak qualifies for the amplitude rule, as each has nearly the same amplitude. Therefore the location rule would have to be used to select the fiducial point. In an illustrative embodiment, this fact alone would be enough to discard the signal and/or discard the template formed using the signal of Figure 18A, as the same rule sets could not be used to define the fiducial point.

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In other embodiments, the sampled signal of Figure 18C may still be used for template verification even though a different fiducial point rule is used. Under such an embodiment, the signal from Figure 18C may still cause rejection of the template formed using the signal shown in Figure 18A. More particularly the signals in Figure 18A and Figure 18C are poorly correlated, as it can be seen that the signal to the left of the fiducial point s21 is lower, while the signal to the right of the fiducial point s21 is higher in Figure 19C than in Figure 18A. If the correlation falls below a defined level, then the template is discarded. In a further embodiment, a beat validation process may be used to assure that a sensed noisy cardiac event, or simply a noise signal, does not reach the template formation steps, preventing template verification due to the likely low correlation of such a non-validated signal. Some example beat validation processes are shown in co-pending U.S. Patent No. 7,248,921 filed June 1, 2004 and entitled METHOD AND DEVICES FOR PERFORMING CARDIAC WAVEFORM APPRAISAL.

Figure 19 is a block diagram for an illustrative template formation process. The process 200 begins by defining a number of sensing parameters, as shown at 202. The sensing parameters may include sampling, window and fiducial point characteristics. Next, a template is filled with data using the sensing parameters, as shown at 204. A validation step follows, as noted at 206. The validation step 206 may include, for example, comparison to successive samples. If validated, the template and its associated sensing parameters are retained as shown at 208. If the

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template and its associated sensing parameters cannot be validated, then they are discarded as shown at 210.

The present invention, in some embodiments, is also embodied in devices using operational circuitry including select electrical components provided within the canister 12 (Figure 1A) or canister 32 (Figure 1B). In such embodiments, the operational circuitry may be configured to enable the above methods to be performed. In some similar embodiments, the present invention may be embodied in readable instruction sets such as a program encoded in machine or controller readable media, wherein the readable instruction sets are provided to enable the operational circuitry to perform the analysis discussed in the above embodiments. Further embodiments may include a controller or microcontroller adapted to read and execute the above methods. These various embodiments may incorporate the illustrative methods shown above, for example.

The following illustrative embodiments are explained in terms of operational circuitry. The operational circuitry may be configured to include such controllers, microcontrollers, logic devices, memory, and the like, as selected, needed, or desired, for performing the method steps of which each is adapted and configured.

The present invention, in an illustrative apparatus embodiment, includes an implantable cardioverter/defibrillator comprising a lead electrode assembly including a number of electrodes, and a canister housing operational circuitry. The illustrative apparatus embodiment may be configured wherein the lead electrode assembly is coupled to the canister, and the operational circuitry is configured to perform steps of discriminating between cardiac rhythms of a patient's heart which are appropriate for therapy, the steps including: sensing a first cardiac event; configuring template parameters for analysis of the first cardiac event; defining a first sensed signal for the first cardiac event using the template parameters; sensing a second cardiac event; defining a second sensed signal for the second cardiac event using the template parameters; and comparing the second sensed signal to the first sensed signal to determine whether the first sensed signal and template parameters are suitable for defining a cardiac event template.

The operational circuitry may, in another embodiment, be configured such that the step of configuring template parameters includes selecting a rule for identifying a fiducial point, and the rule is selected from among a set of rules, the rule is selected in light of the characteristics of the first cardiac event, and the rule for identifying a

fiducial point becomes one of the template parameters. In yet another embodiment, the step of configuring template parameters further includes selecting a number of samples of the first sensed signal around the fiducial point, and the configuration of samples around the fiducial point becomes one of the template parameters. In another embodiment, the operational circuitry is configured such that the step of selecting a number of samples includes identifying the start and end of a cardiac event. In on embodiment, the operational circuitry is configured such that the cardiac event is a QRS complex. In some embodiments, the operational circuitry is configured such that the set of rules includes an amplitude rule related to the relative amplitudes of peaks in the sensed signal. The set of rules may include a location rule related to the location of a peak in the sensed signal. In yet another embodiment, the operational circuitry is configured such that the set of rules includes a location rule related to the location of a peak in the sensed signal.

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In yet another embodiment, the operational circuitry is configured such that the set of rules includes a notch rule related to identifying a notched cardiac signal, wherein the notch rule includes analysis of whether there are multiple peaks within a predefined range of one another in the cardiac signal. The operational circuitry may be configured such that the notch rule selects the first peak in time if there are multiple peaks within the predefined range. In another embodiment, the operational circuitry may be configured such that the step of configuring template parameters further includes selecting a number of samples of the first sensed signal around a fiducial point in the first sensed signal; wherein the configuration of samples around the fiducial point becomes one of the template parameters. The operational circuitry, in an illustrative embodiment, is configured such that samples are selected using the following steps: first, a number of samples are observed on either side of the fiducial point; next, it is determined whether a desired QRS segment begins and ends within the number of samples; and the number of samples on either side of the fiducial point is adjusted to capture the QRS segment and exclude at least some samples not corresponding to the desired QRS segment. Another illustrative embodiment includes one wherein the operational circuitry is configured such that the step of configuring template parameters includes observing whether a notched QRS complex is likely, and, if so, adjusting the template parameters to assure that a repeatably detectable fiducial point is chosen.

Another embodiment includes an implantable cardioverter/defibrillator comprising a lead electrode assembly including a number of electrodes and a canister housing operational circuitry; wherein: the lead electrode assembly is coupled to the canister; and the operational circuitry is configured to perform steps of discriminating between cardiac rhythms of a patient's heart which are appropriate for therapy. The steps may include sampling a signal using the lead electrode assembly while implanted in a patient's torso in locations chosen for capturing cardiac signals; defining a first sensing window around a first fiducial point to capture a QRS segment; observing the definition of the first sensing window to create template parameters; defining a second sensing window around a second fiducial point using the template parameters; and comparing data in the first sensing window to data in the second sensing window to verify whether to define a valid template using the template parameters. The operational circuitry may be configured such that the step of defining a first sensing window includes identifying a fiducial point by selecting a rule from among a set of rules in light of the characteristics of the QRS segment in the first sensing window, wherein the rule selected for identifying a fiducial point becomes one of the template parameters. Further, the operational circuitry may be configured such that the step defining a first sensing window around a first fiducial point includes identifying the start and end of a cardiac event. If desired, the cardiac event may be a QRS complex.

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In another embodiment, the operational circuitry is configured such that the set of rules includes an amplitude rule related to the relative amplitudes of peaks in the sampled signal, and a location rule related to the location of a peak in the sampled signal. In yet another embodiment, the operational circuitry is configured such that the step of defining a first sensing window includes selecting a number of samples around a fiducial point, wherein the configuration of samples around the fiducial point becomes one of the template parameters. The operational circuitry may be configured such that the samples are selected using the following steps: a fiducial point is selected; then a number of samples are observed on either side of the fiducial point; then it is determined whether a desired QRS segment begins and ends within the number of samples; and the number of samples on either side of the fiducial point is adjusted to capture the QRS segment and exclude at least some samples not corresponding to the desired QRS segment.

In yet another embodiment, the operational circuitry is configured such that the step of defining a first sensing window includes observing whether a notched QRS complex is likely, and, if so, adjusting the template parameters to assure that a repeatably detectable fiducial point is chosen.

implantable include an embodiment may illustrative An cardioverter/defibrillator comprising a lead electrode assembly including a number of electrodes and a canister housing operational circuitry, wherein: the lead electrode assembly is coupled to the canister; and the operational circuitry is configured to perform steps of discriminating between cardiac rhythms of a patient's heart which are appropriate for therapy. The steps of discriminating may include forming a template using at least the steps of: sensing a first cardiac event; identifying a first fiducial point in the first cardiac event using a set of rules; sensing a second cardiac event; identifying a second fiducial point in the second cardiac event using the set of rules; determining whether the first fiducial point and second fiducial point were identified using the same rule; and, if not, discarding the first cardiac event.

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implantable includes embodiment an illustrative Another cardioverter/defibrillator comprising a lead electrode assembly including a number of electrodes and a canister housing operational circuitry, wherein the lead electrode assembly is coupled to the canister and the operational circuitry is configured to perform steps of discriminating between cardiac rhythms of a patient's heart which are appropriate for therapy. The discriminating steps may include forming a template using at least the steps of receiving a first cardiac signal from the lead electrode assembly, selecting a fiducial point in the first cardiac signal, forming a template around the fiducial point, and verifying the template by receiving additional cardiac signals and using the template to compare the additional cardiac signals to the first cardiac signal. In another embodiment, the operational circuitry is configured such that the step of selecting a fiducial point includes identifying a fiducial point by selecting a rule from among a set of rules in light of the characteristics of first cardiac signal, wherein the rule selected for identifying a fiducial point becomes one of the template parameters. The operational circuitry may be configured such that the step forming a template around the fiducial point includes identifying the start and end of a cardiac event. In another embodiment, the operational circuitry may be configured such that the set of rules includes an amplitude rule related to the relative amplitudes of peaks in the cardiac signal. The set of rules may further include a location rule related to the location of a peak in the cardiac signal. In another embodiment, the operational circuitry is configured such that the set of rules includes a location rule related to the location of a peak in the cardiac signal.

In another embodiment, the operational circuitry is configured such that the step of forming a template includes selecting a number of samples around the fiducial point, wherein the configuration of samples around the fiducial point becomes one of the template parameters. The operational circuitry may be configured such that the samples are selected using the following steps: a number of samples are observed on either side of the fiducial point; it is determined whether a desired QRS segment begins and ends within the number of samples; and the number of samples on either side of the fiducial point is adjusted to capture the QRS segment and exclude at least some samples not corresponding to the desired QRS segment. In yet another embodiment, the operational circuitry is configured such that the step of selecting a fiducial point includes observing whether a notched QRS complex is likely, and, if so, adjusting the template parameters to assure that a repeatably detectable fiducial point is chosen.

Numerous characteristics and advantages of the invention covered by this document have been set forth in the foregoing description. It will be understood, however, that this disclosure is, in many aspects, only illustrative.

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What is claimed is:

1. A method of cardiac signal analysis performed as a diagnostic method performed by an implantable cardiac device which comprises a plurality of electrodes and a canister housing operational circuitry with the electrodes coupled to the operational circuitry, the method comprising:

sensing a first cardiac event;

configuring template parameters for analysis of the first cardiac event;

defining a first sensed signal for the first cardiac event using the template parameters;

sensing a second cardiac event;

defining a second sensed signal for the second cardiac event using the template parameters;

comparing the second sensed signal to the first sensed signal to determine whether the first sensed signal and template parameters are suitable for defining a cardiac event template; and

if the first sensed signal and template parameters are suitable, retaining the first sensed signal and template parameters for using to define a cardiac event template;

or, if the first sensed signal and template parameters are not suitable, discarding the first sensed signal and template parameters from use in defining a cardiac event template; and

wherein the step of defining the second sensed signal includes using the template parameters configured for analysis of the first sensed signal to define the second sensed signal.

2. The method of claim 1, wherein the step of configuring template parameters includes selecting a rule for identifying a fiducial point, wherein:

the rule is selected from among a set of rules;

the rule is selected in light of the characteristics of the first cardiac event; and the rule for identifying a fiducial point becomes one of the template parameters.

- 3. The method of claim 2, wherein the step of configuring template parameters further includes selecting a number of samples of the first sensed signal around the fiducial point; wherein the configuration of samples around the fiducial point becomes one of the template parameters.
- 4. The method of claim 3, wherein the step of selecting a number of samples includes identifying the start and end of a cardiac event.
- 5. The method of claim 4, wherein the cardiac event is a QRS complex.
- 6. The method of claim 2, wherein the set of rules includes an amplitude rule related to the relative amplitudes of peaks in the sensed signal.
- 7. The method of claim 6, wherein the set of rules includes a location rule related to the location of a peak in the sensed signal.
- 8. The method of claim 2 wherein the set of rules includes a location rule related to the location of a peak in the sensed signal.
- 9. The method of claim 2 wherein the set of rules includes a notch rule related to identifying a notched cardiac signal, wherein the notch rule includes analysis of whether there are multiple peaks within a predefined range of one another in the cardiac signal.
- 10. The method of claim 9 wherein the notch rule selects the first peak in time if there are multiple peaks within the predefined range.
- 11. The method of claim 1, wherein the step of configuring template parameters further includes selecting a number of samples of the first sensed signal around a fiducial point in the first sensed signal; wherein the configuration of samples around the fiducial point becomes one of the template parameters.

12. The method of claim 11, wherein the samples are selected using the following steps:

first, a number of samples are observed on either side of the fiducial point;

next, it is determined whether a desired QRS segment begins and ends within the number of samples; and

the number of samples on either side of the fiducial point is adjusted to capture the QRS segment and exclude at least some samples not corresponding to the desired QRS segment.

- 13. The method of claim 1, wherein the step of configuring template parameters includes observing whether a notched QRS complex is likely, and, if so, adjusting the template parameters to assure that a repeatably detectable fiducial point is chosen.
- 14. An implantable cardioverter/defibrillator comprising:

a lead electrode assembly including a number of electrodes; and

a canister that includes operational circuitry housed therein; wherein:

the lead electrode assembly is coupled to the canister; and

the operational circuitry is configured to perform steps of discriminating between cardiac rhythms of a patient's heart which are appropriate for therapy, the steps including:

sensing a first cardiac event;

configuring template parameters for analysis of the first cardiac event;

defining a first sensed signal for the first cardiac event using the template parameters;

sensing a second cardiac event;

defining a second sensed signal for the second cardiac event using the template parameters;

comparing the second sensed signal to the first sensed signal to determine whether the first sensed signal and template parameters are suitable for defining a cardiac event template; and

if the first sensed signal and template parameters are suitable, retaining the first sensed signal and template parameters for using to define a cardiac event template;

or, if the first sensed signal and template parameters are not suitable, discarding the first sensed signal and template parameters from use in defining a cardiac event template;

wherein the operational circuitry is configured to define the second sensed signal by using the template parameters configured for analysis of the first sensed signal to define the second sensed signal.

15. The implantable cardioverter/defibrillator of claim 14, wherein the operational circuitry is configured such that the step of configuring template parameters includes selecting a rule for identifying a fiducial point, and:

the rule is selected from among a set of rules;

the rule is selected in light of the characteristics of the first cardiac event; and the rule for identifying a fiducial point becomes one of the template parameters.

- 16. The implantable cardioverter/defibrillator of claim 15, wherein the operational circuitry is configured such that the step of configuring template parameters further includes selecting a number of samples of the first sensed signal around the fiducial point, and the configuration of samples around the fiducial point becomes one of the template parameters.
- 17. The implantable cardioverter/defibrillator of claim 16, wherein the operational circuitry is configured such that the step of selecting a number of samples includes identifying the start and end of a cardiac event.
- 18. The implantable cardioverter/defibrillator of claim 17, wherein the operational circuitry is configured such that the cardiac event is a QRS complex.

- 19. The implantable cardioverter/defibrillator of claim 15, wherein the operational circuitry is configured such that the set of rules includes an amplitude rule related to the relative amplitudes of peaks in the sensed signal.
- 20. The implantable cardioverter/defibrillator of claim 19, wherein the operational circuitry is configured such that the set of rules includes a location rule related to the location of a peak in the sensed signal.
- 21. The implantable cardioverter/defibrillator of claim 15, wherein the operational circuitry is configured such that the set of rules includes a location rule related to the location of a peak in the sensed signal.
- 22. The implantable cardioverter/defibrillator of claim 15, wherein the operational circuitry is configured such that the set of rules includes a notch rule related to identifying a notched cardiac signal, wherein the notch rule includes analysis of whether there are multiple peaks within a predefined range of one another in the cardiac signal.
- 23. The implantable cardioverter/defibrillator of claim 22, wherein the operational circuitry is configured such that the notch rule selects the first peak in time if there are multiple peaks within the predefined range.
- 24. The implantable cardioverter/defibrillator of claim 14, wherein the operational circuitry is configured such that the step of configuring template parameters further includes selecting a number of samples of the first sensed signal around a fiducial point in the first sensed signal; wherein the configuration of samples around the fiducial point becomes one of the template parameters.
- 25. The implantable cardioverter/defibrillator of claim 24, wherein the operational circuitry is configured such that the samples are selected using the following steps:

first, a number of samples are observed on either side of the fiducial point;

next, it is determined whether a desired QRS segment begins and ends within the number of samples; and

the number of samples on either side of the fiducial point is adjusted to capture the QRS segment and exclude at least some samples not corresponding to the desired QRS segment.

- 26. The implantable cardioverter/defibrillator of claim 14, wherein the operational circuitry is configured such that the step of configuring template parameters includes observing whether a notched QRS complex is likely, and, if so, adjusting the template parameters to assure that a repeatably detectable fiducial point is chosen.
- 27. The implantable cardioverter/defibrillator of claim 14, wherein the operational circuitry comprises a readable medium including an instruction set for performing the steps of discriminating.

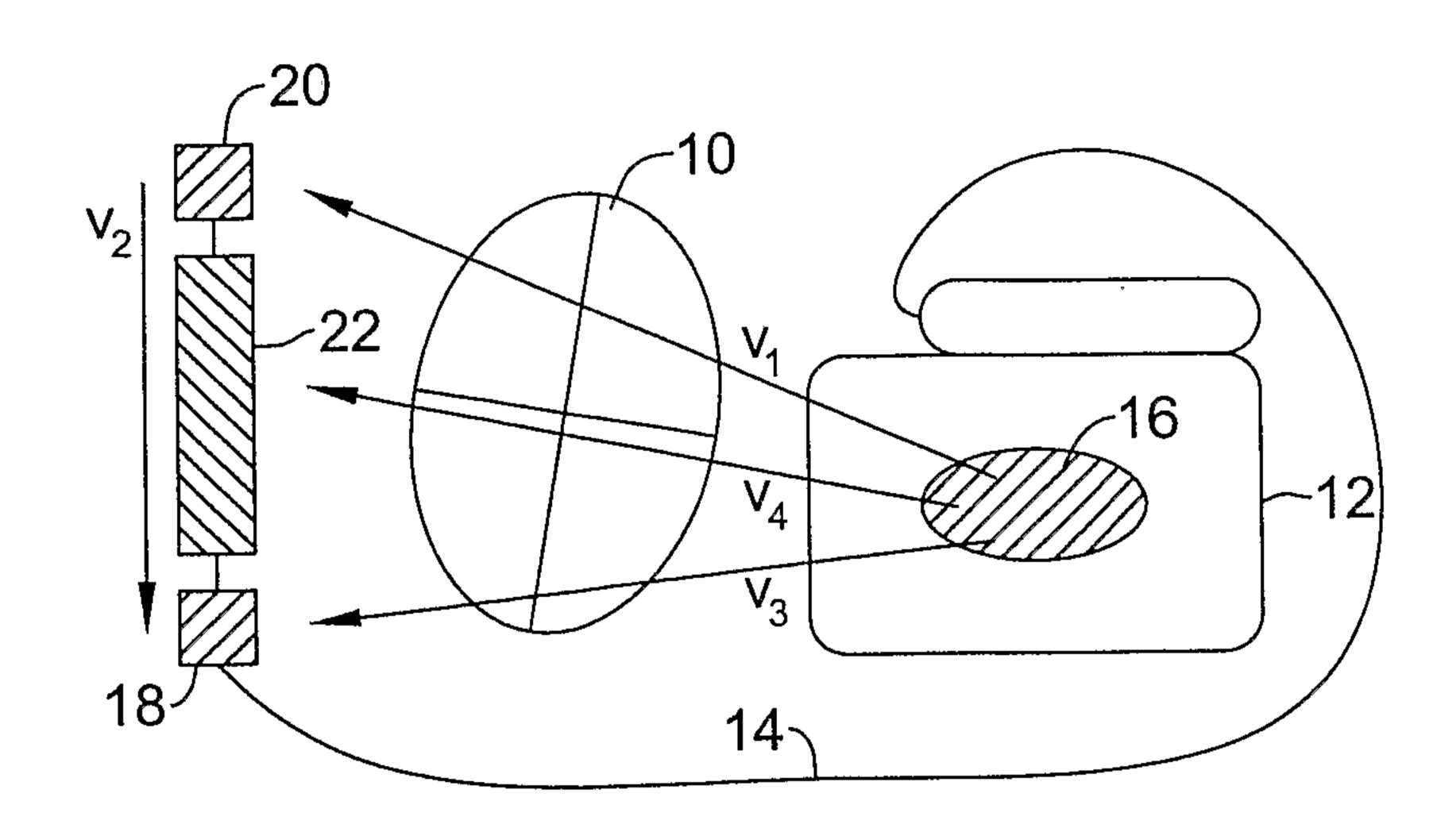


Fig. 1A

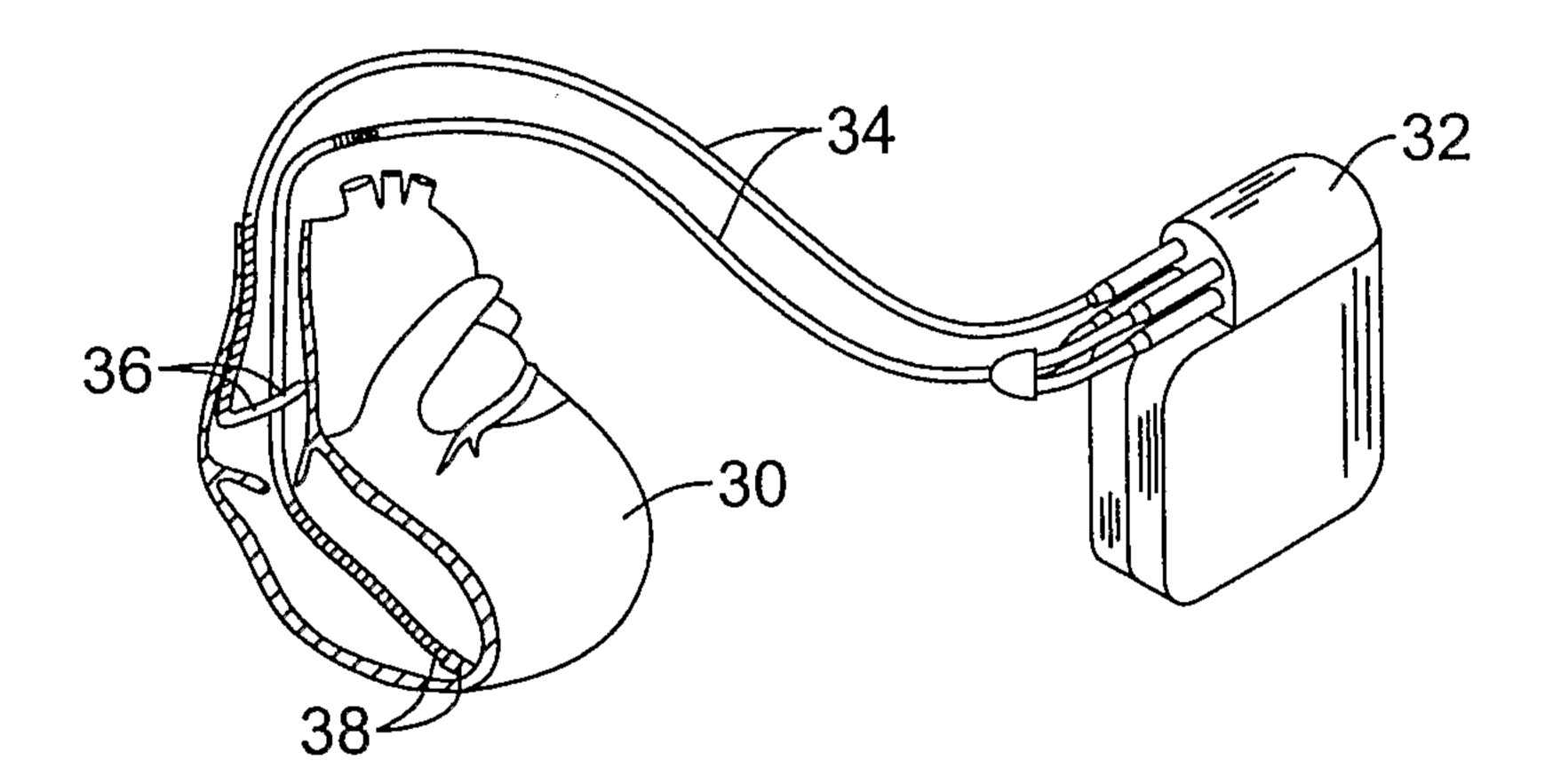


Fig. 1B

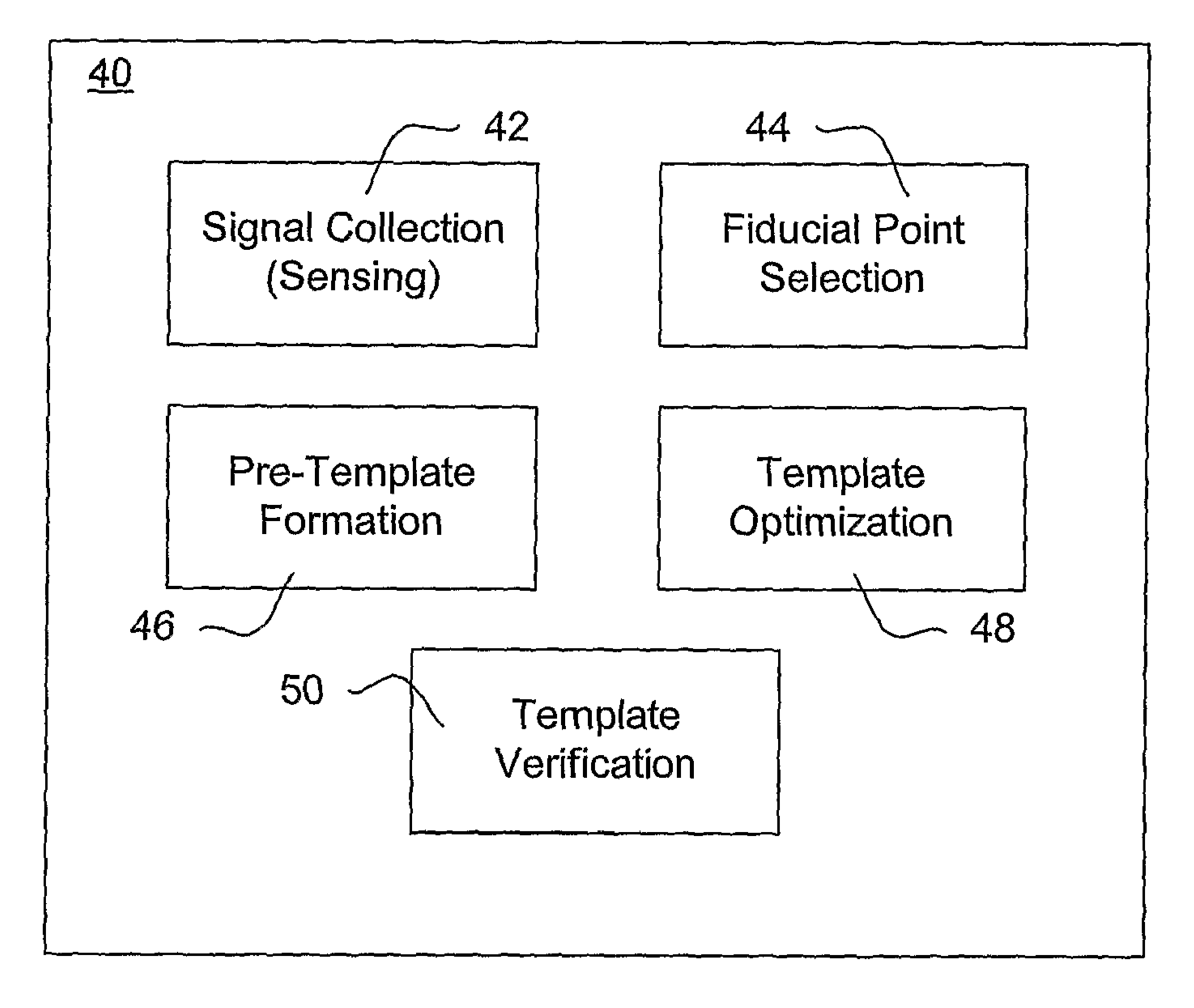
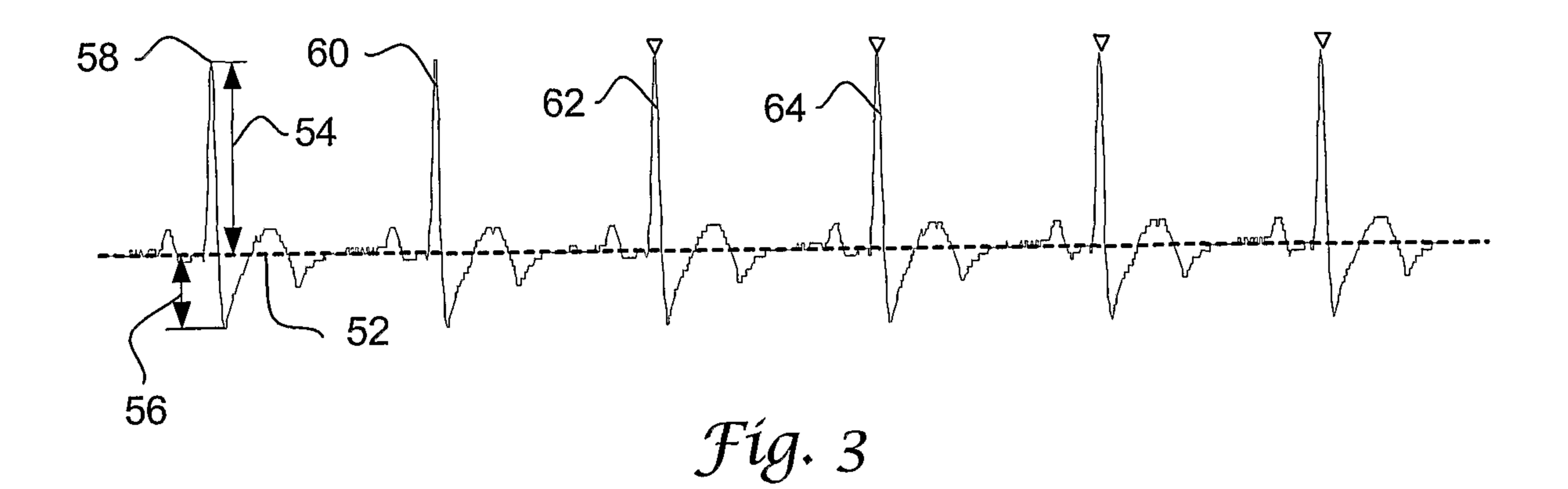
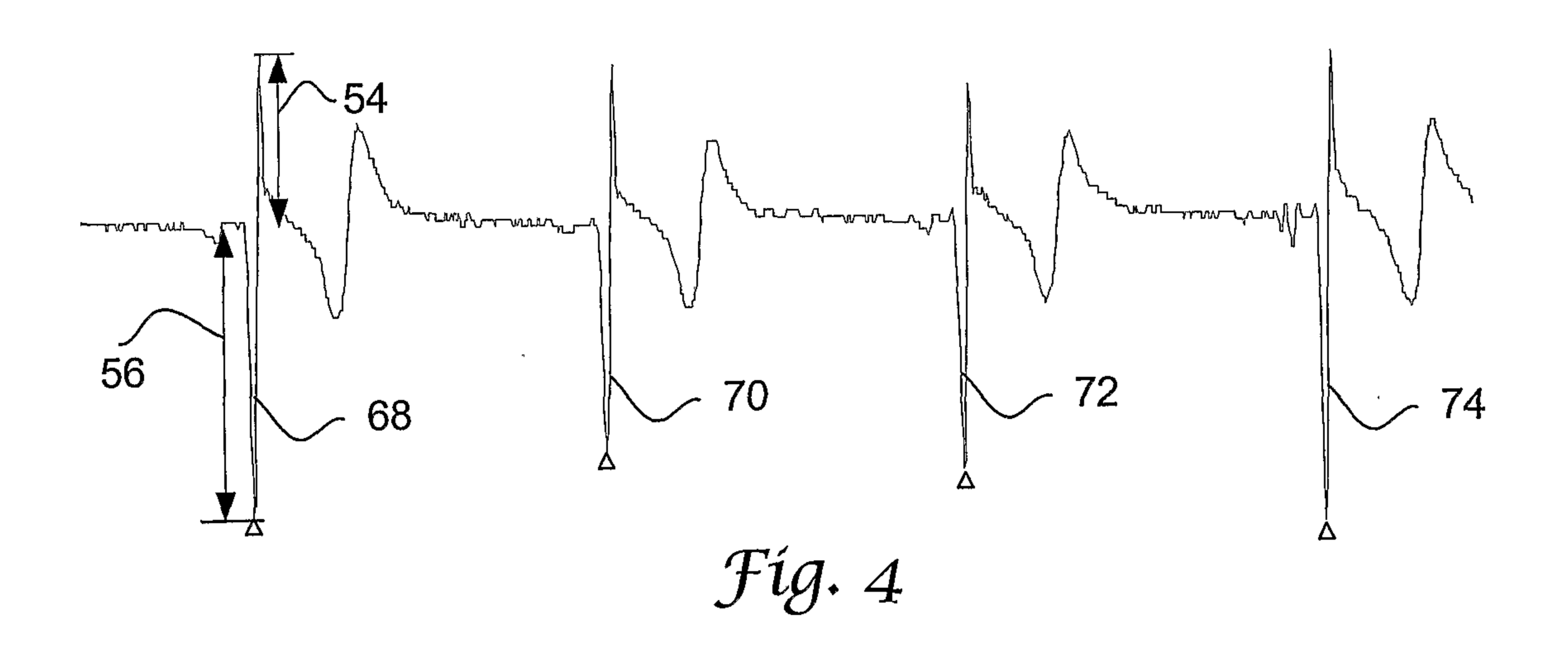
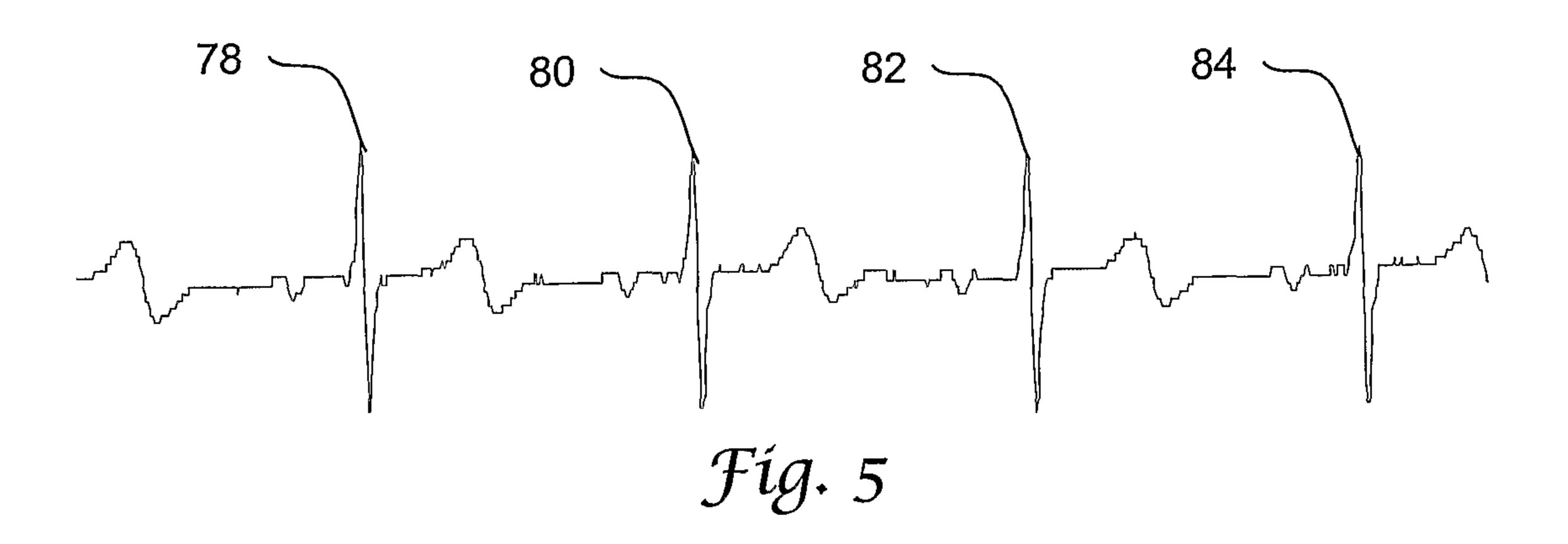


Fig. 2







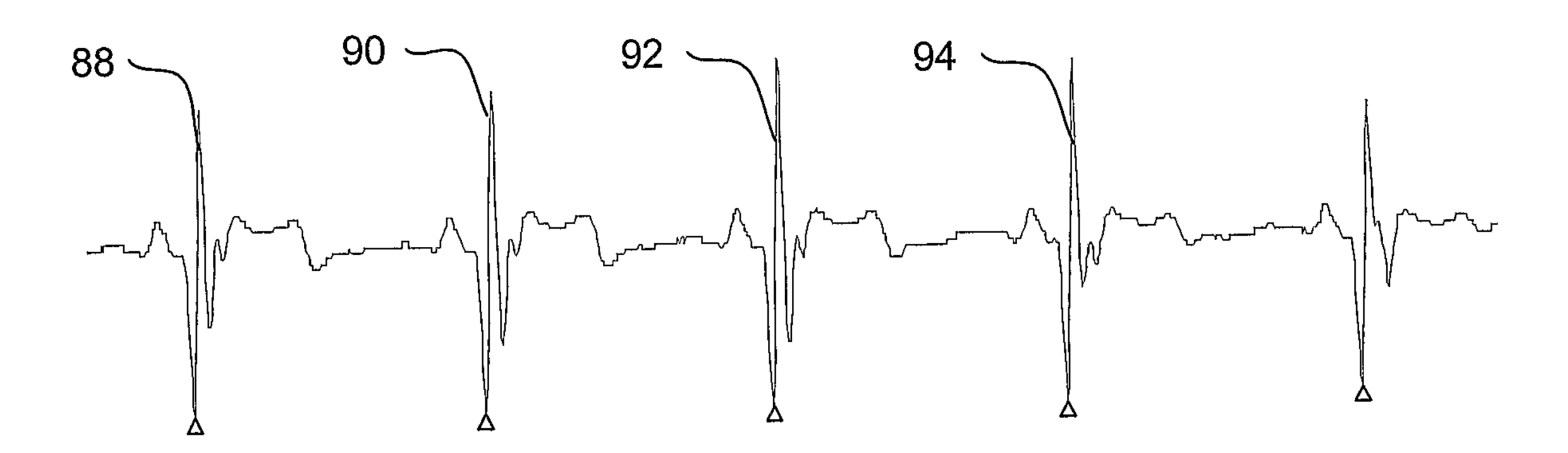
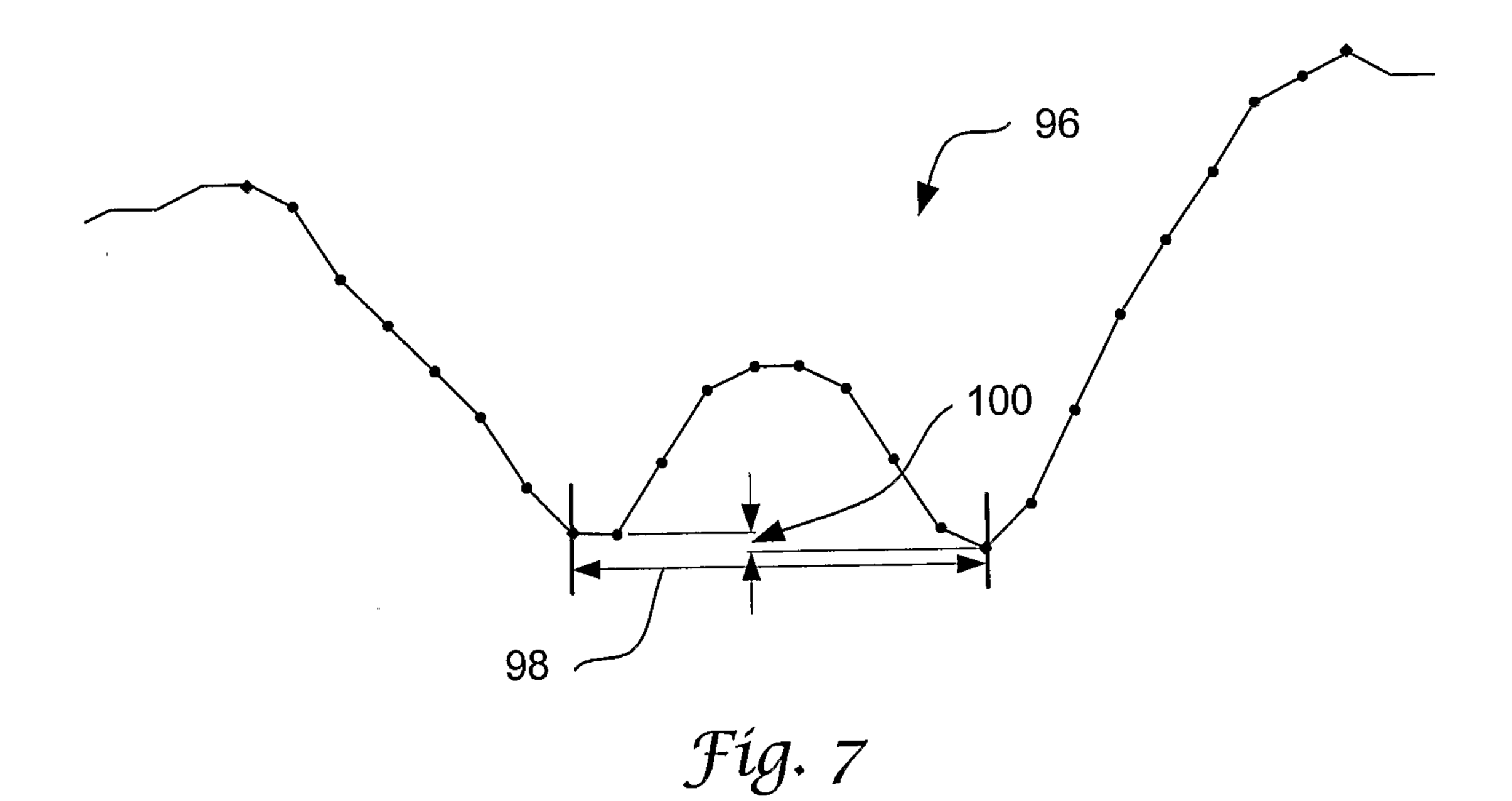


Fig. 6



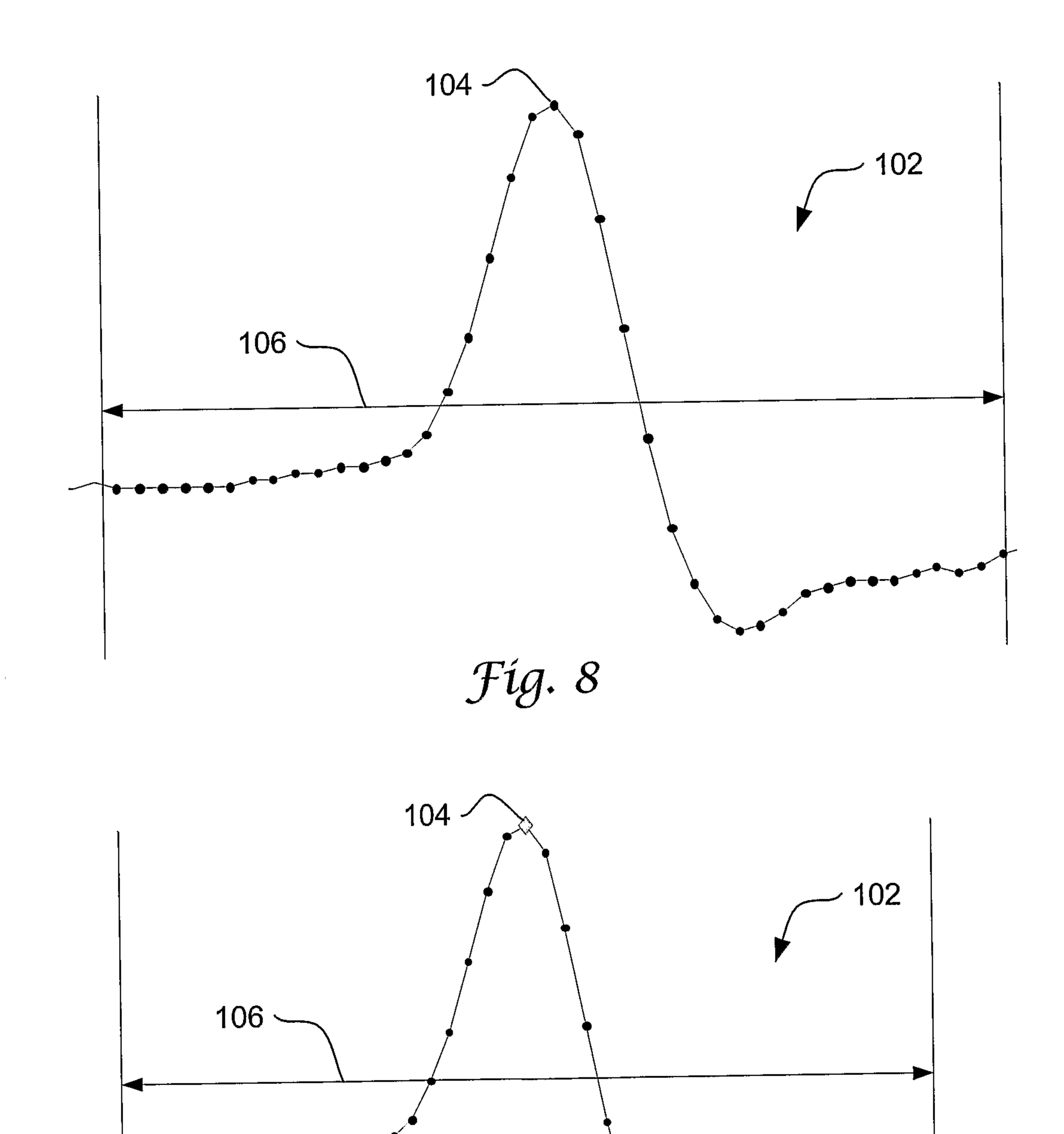


Fig. 9

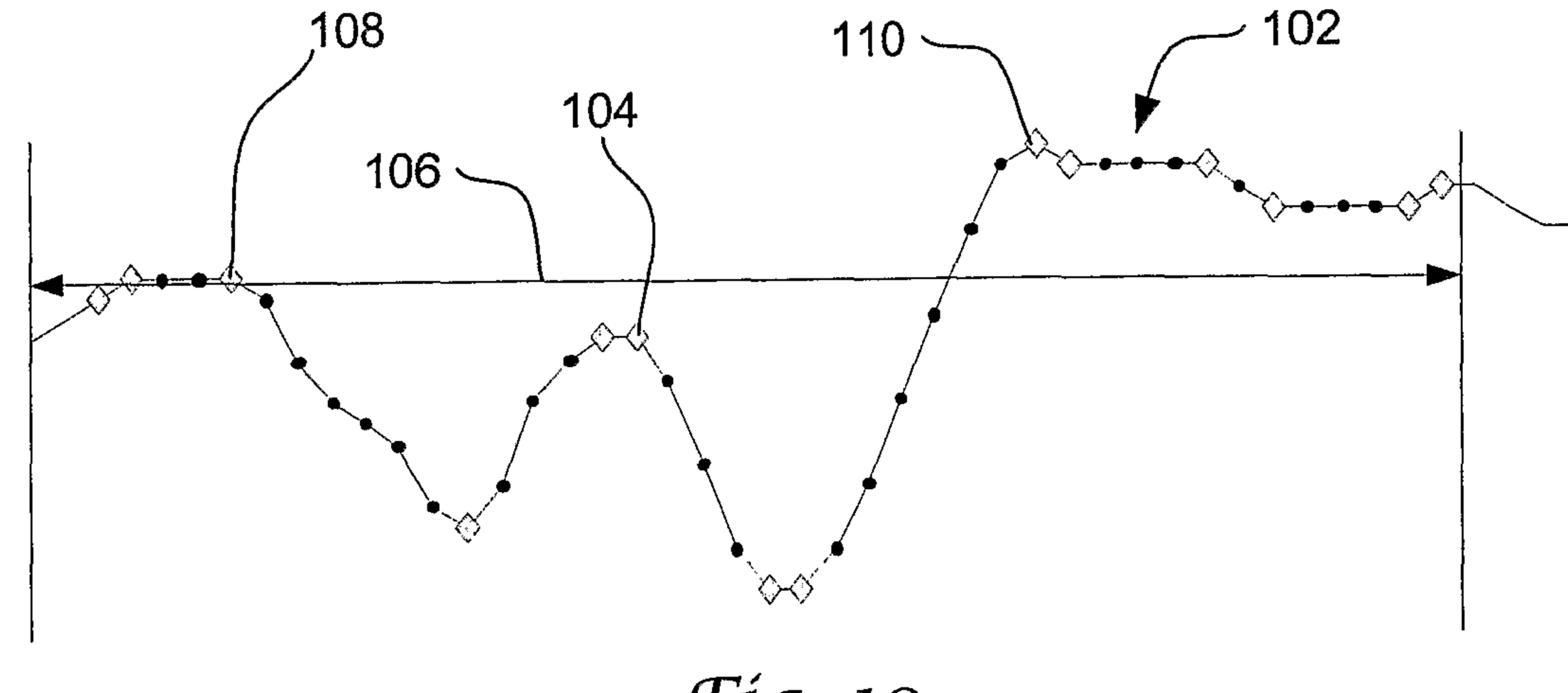


Fig. 10

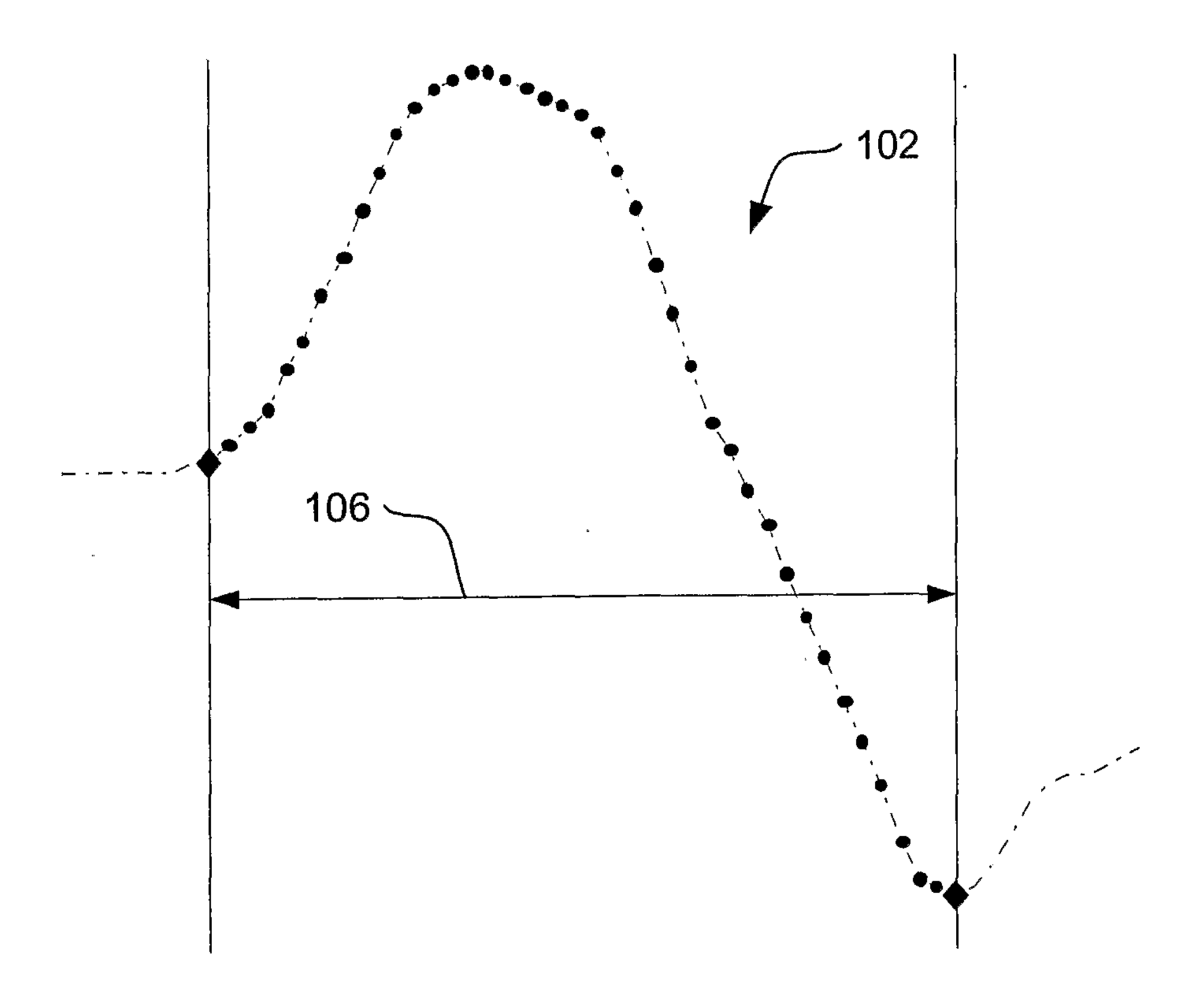
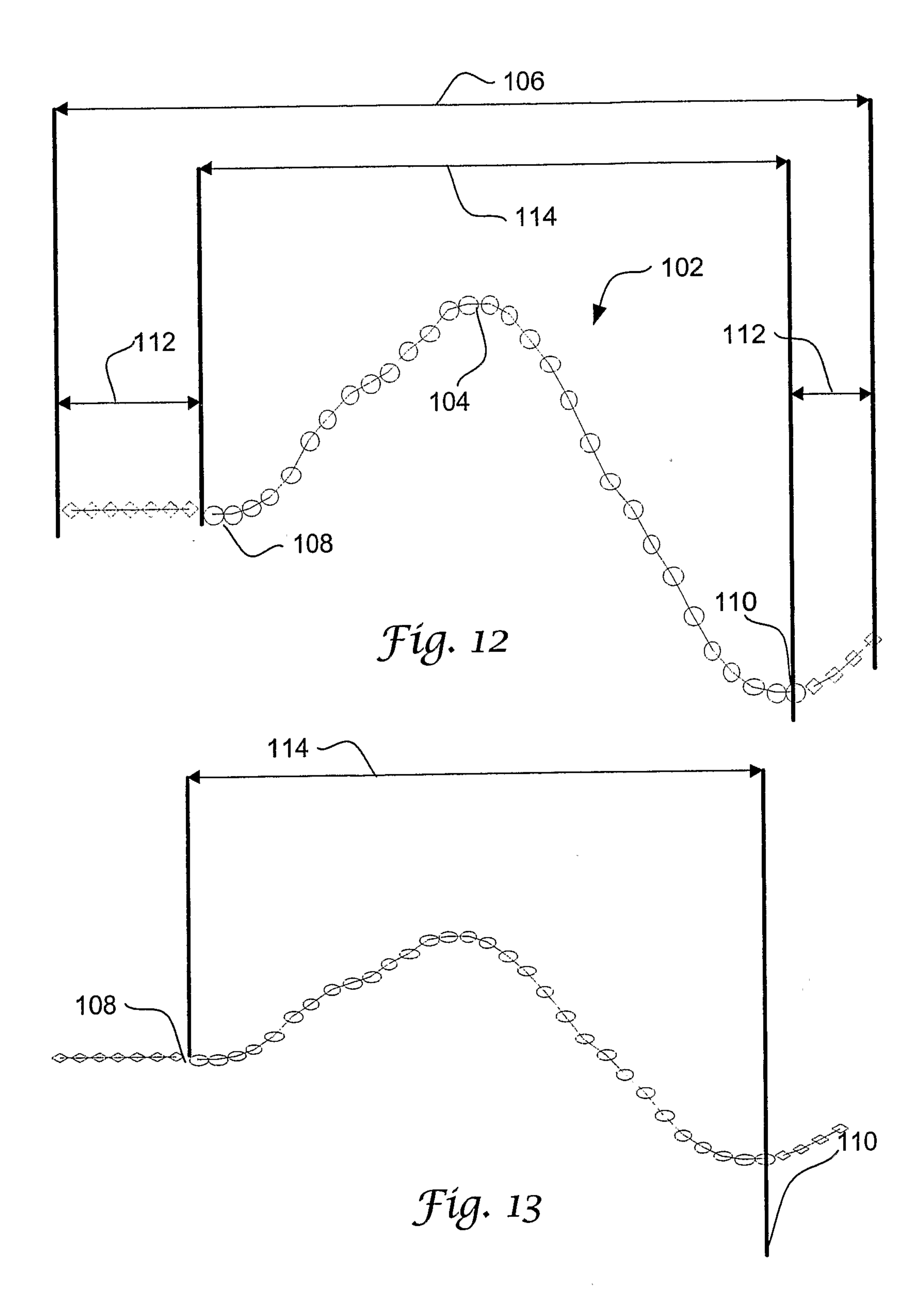


Fig. 11



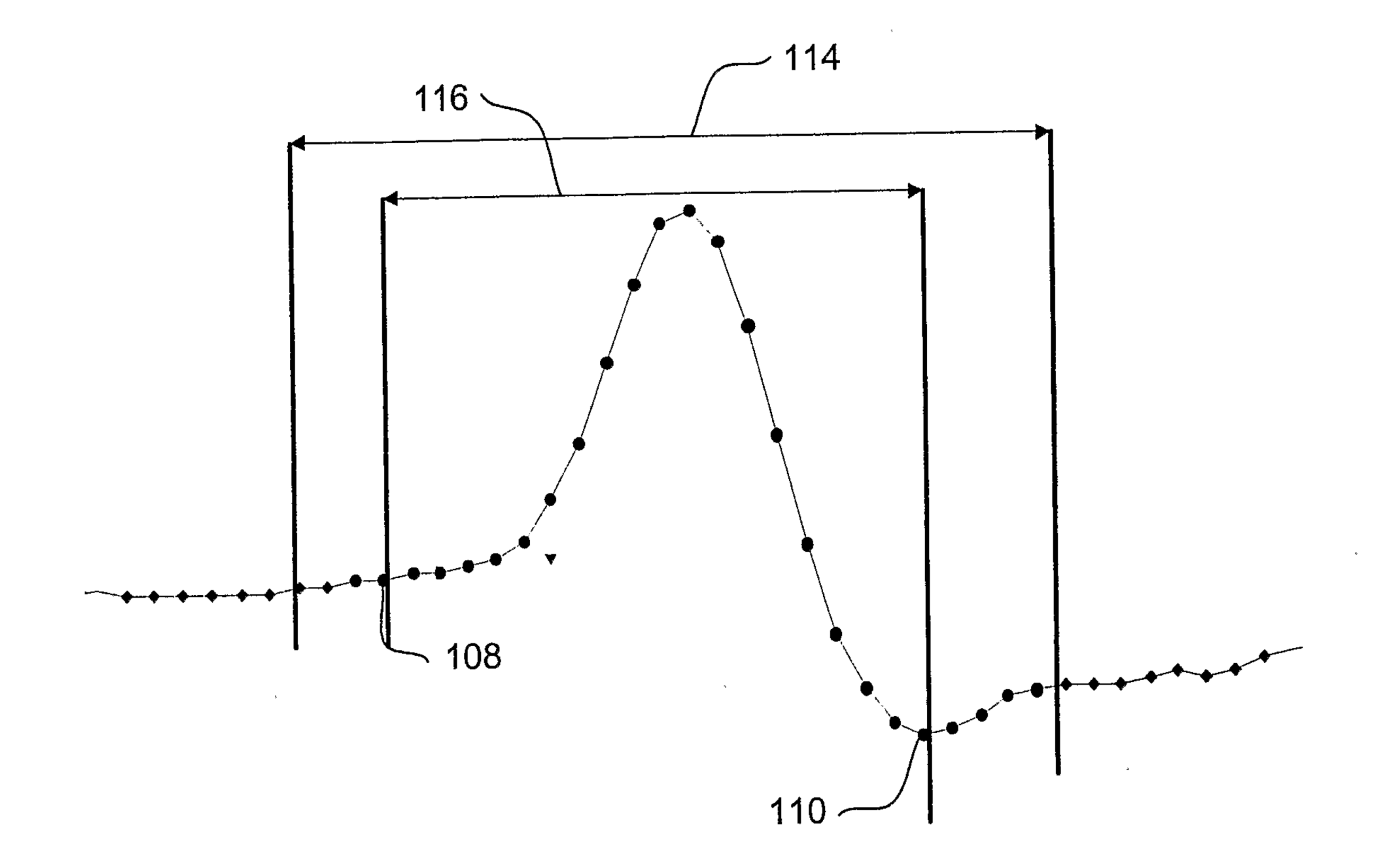
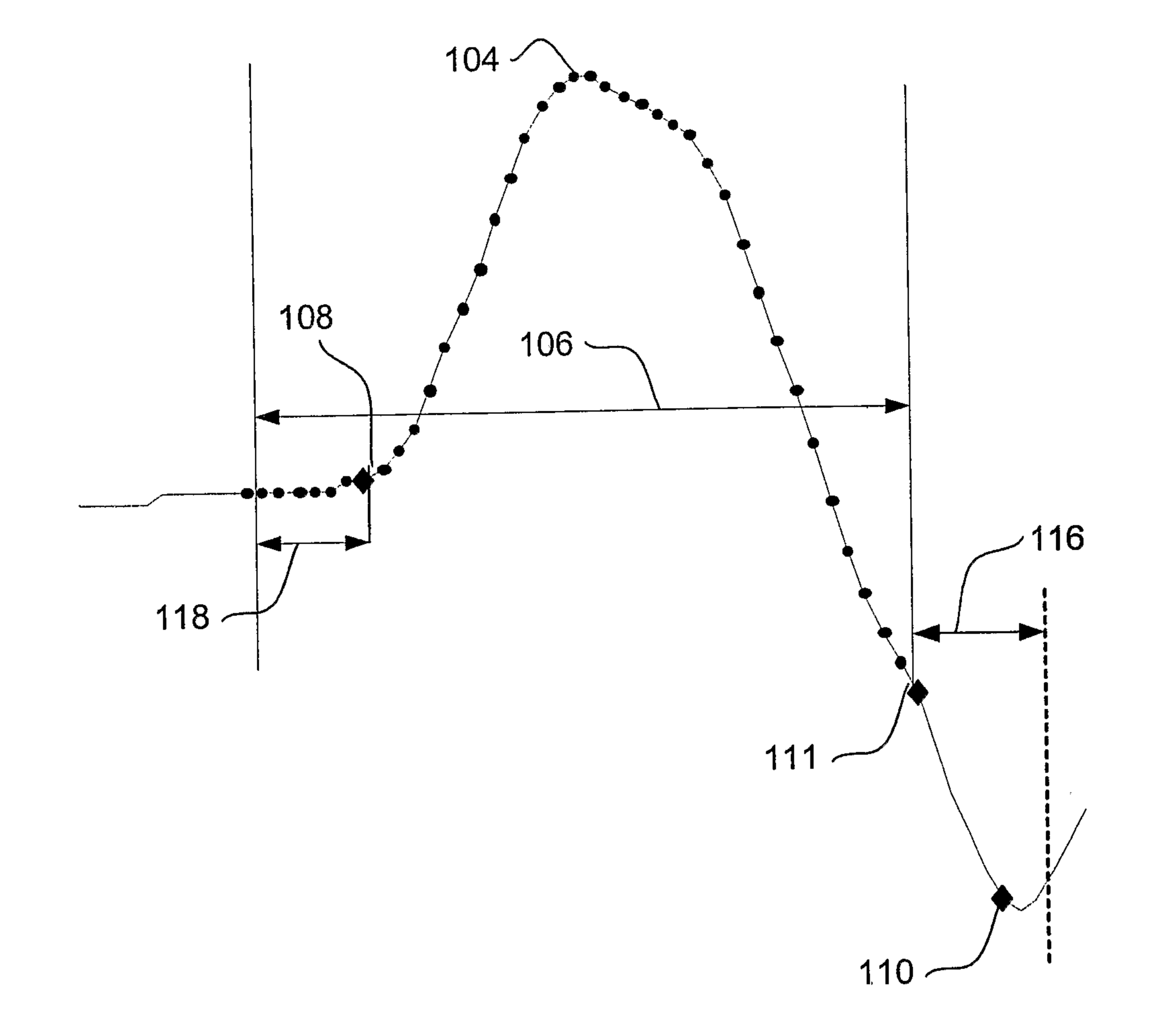


Fig. 14



Fíg. 15

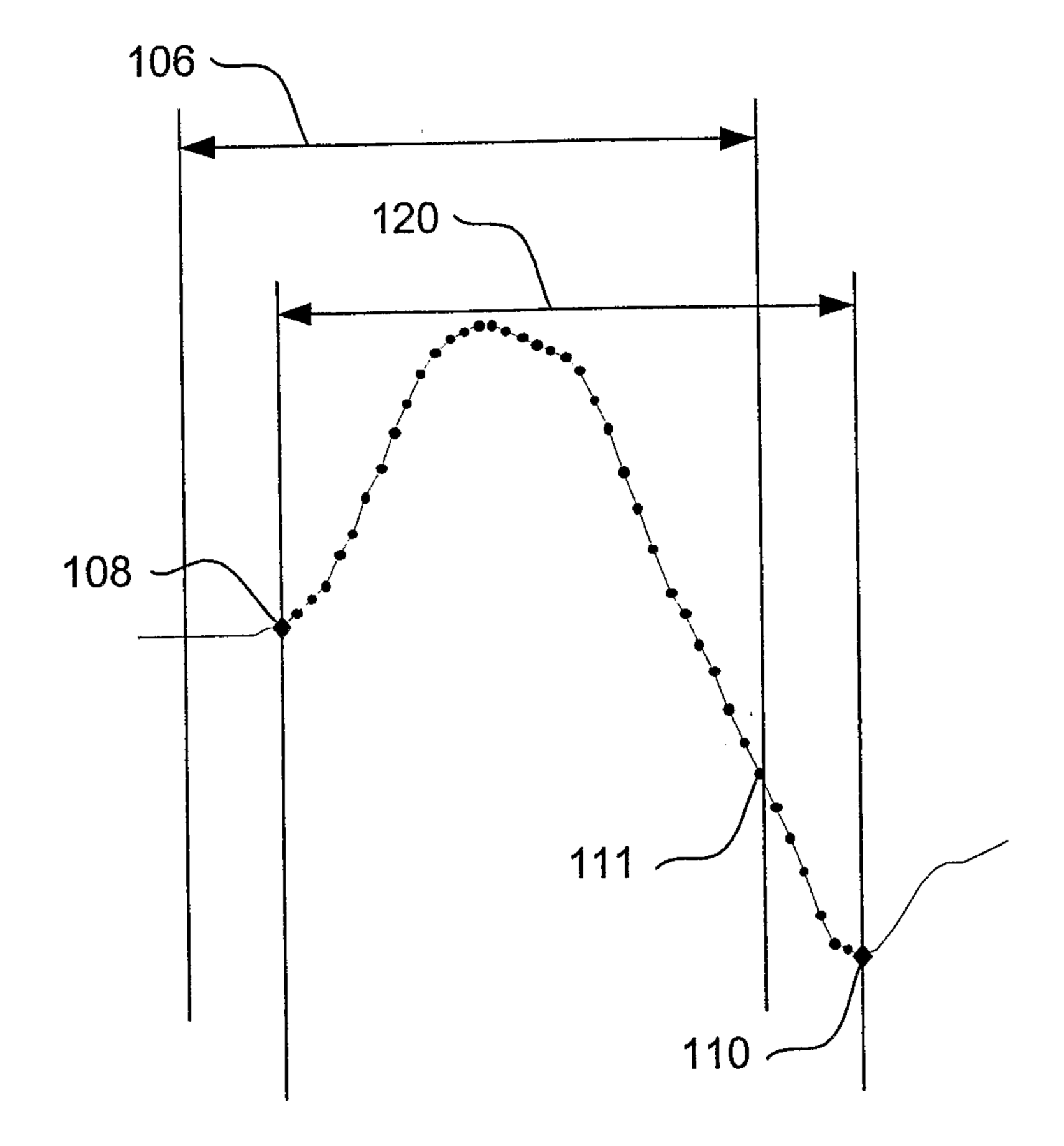


Fig. 16

