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(54) **EXTENDED WEAR ELECTROCARDIOGRAPHY PATCH USING INTERLACED WIRE ELECTRODES**

ELEKTROKARDIOGRAFIE-PFLASTER MIT VERLÄNGERTER TRAGEZEIT UND MIT VERFLOCHTENEN DRAHTELEKTRODEN

TIMBRE D'ÉLECTROCARDIOGRAPHIE À UTILISATION PROLONGÉE UTILISANT DES FILS-ÉLECTRODES ENTRELACÉS

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- **WEI PINGHUNG ET AL: "A stretchable and flexible system for skin-mounted measurement of motion tracking and physiological signals", 2014 36TH ANNUAL INTERNATIONAL CONFERENCE OF THE IEEE ENGINEERING IN MEDICINE AND BIOLOGY SOCIETY, IEEE, 26 August 2014 (2014-08-26), pages 5772-5775, XP032674410, DOI: 10.1109/EMBC.2014.6944939 [retrieved on 2014-11-02]**
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Description

TECHNICAL FIELD

[0001] This application relates in general to electrocardiographic monitoring and manufacture and, in particular, to an extended wear electrocardiography patch using interlaced wire electrodes.

BACKGROUND ART

[0002] An electrocardiogram (ECG) is a tool used by physicians to diagnose heart problems and other potential health concerns. A full 12-lead ECG provides a multi-vector snapshot of heart function, typically recorded over 12 seconds, that can help diagnose rate and regularity of heartbeats, effect of drugs or cardiac devices, including pacemakers and implantable cardioverter-defibrillators (ICDs), and whether a patient has heart disease of any sort. Full 12-lead ECGs are used in-clinics or hospitals and, as a result, are limited to recording only those heart-related aspects present at the time of recording. Sporadic conditions that may not show up during a 12-second ECG recording require other means to diagnose them. These sporadic conditions include fainting or syncope; rhythm disorders, such as tachyarrhythmias and bradyarrhythmias; apneic episodes; and other cardiac and related disorders. Thus, a 12-lead ECG only provides a partial picture and can be insufficient for complete patient diagnosis of many cardiac disorders.

[0003] Diagnostic efficacy of problems, like syncope or cardiac arrhythmias, can be improved through the use of long-term extended wear ECG monitoring. Recording sufficient ECG and related physiological data over an extended period of time remains a significant challenge to healthcare providers, despite over a 40-year history of such efforts. Extended period monitoring essentially enables a physician to identify cardiac conditions, specifically, rhythm disorders, and other physiological events of potential concern. A 30-day observation day period is considered the "gold standard" of ECG monitoring, yet achieving a 30-day observation day period has heretofore proven unworkable because such ECG monitoring systems are arduous to employ, cumbersome to the patient, and excessively costly to manufacture and deploy. Nevertheless, if a patient's ECG could be recorded in an ambulatory setting over prolonged time periods, thereby allowing the patient to engage in activities of daily living, the chances of acquiring meaningful medical information and capturing an abnormal event while the patient is engaged in normal activities becomes more likely to be achieved.

[0004] Conventionally, maintaining continual contact between ECG electrodes and the skin after a day or two has been a problem. Time, dirt, moisture, and other environmental contaminants, as well as perspiration, skin oil, and dead skin cells from the patient's body, can get between an ECG electrode's non-conductive adhesive

and the skin's surface. All of these factors adversely affect electrode adhesion and the quality of cardiac signal recordings. Furthermore, the physical movements of the patient and their clothing impart various compressional, tensile, and torsional forces on the contact point of an ECG electrode, especially over long recording times, and an inflexibly fastened ECG electrode will be prone to becoming dislodged. Moreover, dislodgment may occur unbeknownst to the patient, making the ECG recordings worthless. Further, some patients may have skin that is susceptible to itching or irritation, and the wearing of ECG electrodes can aggravate such skin conditions. Thus, a patient may want or need to periodically remove or replace ECG electrodes during a long-term ECG monitoring period, whether to replace a dislodged electrode, re-establish better adhesion, alleviate itching or irritation, allow for cleansing of the skin, allow for showering and exercise, or for other purpose. Such replacement or slight alteration in electrode location actually facilitates the goal of recording the ECG signal for long periods of time.

[0005] In addition, the high cost of the patient-wearable components used to provide long-term extended ECG monitoring can negatively influence the availability and use of monitors. Ideally, disposable, single-use components, such as adhesive electrodes, should be low cost, while other components of higher complexity, particularly the electronics hardware that detects and records ECG and related physiology, may be of unavoidably higher cost. To a degree, costs can be balanced by designing higher complexity components to be re-usable, but when the total cost of a full ECG monitoring ensemble remains high, despite the utilization of re-usable parts, the number of monitors available for use by healthcare providers can be inhibited. Cost, then, becomes a barrier to entry, which, in turn, can hinder or prevent healthcare providers from obtaining the means with which to efficaciously identify the physiology underlying sporadic cardiac arrhythmic conditions and can ultimately contribute to a failure to make a proper and timely medical diagnosis.

[0006] Conventionally, Holter monitors are widely used for long-term extended ECG monitoring. Typically, they are often used for only 24-48 hours. A typical Holter monitor is a wearable and portable version of an ECG that include cables for each electrode placed on the skin and a separate battery-powered ECG recorder. The cable and electrode combination (or leads) are placed in the anterior thoracic region in a manner similar to what is done with an in-clinic standard ECG machine. The duration of a Holter monitoring recording depends on the sensing and storage capabilities of the monitor, as well as battery life. A "looping" Holter (or event) monitor can operate for a longer period of time by overwriting older ECG tracings, thence "recycling" storage in favor of extended operation, yet at the risk of losing event data. Although capable of extended ECG monitoring, Holter monitors are cumbersome, expensive and typically only available by medical prescription, which limits their usability. Further, the skill required to properly place the elec-

trodes on the patient's chest hinders or precludes a patient from replacing or removing the precordial leads and usually involves moving the patient from the physician office to a specialized center within the hospital or clinic.

[0007] The ZIO XT Patch and ZIO Event Card devices, manufactured by iRhythm Tech., Inc., San Francisco, CA, are wearable stick-on monitoring devices that are typically worn on the upper left pectoral region to respectively provide continuous and looping ECG recording. The location is used to simulate surgically implanted monitors. Both of these devices are prescription-only and for single patient use. The ZIO XT Patch device is limited to a 14-day monitoring period, while the electrodes only of the ZIO Event Card device can be worn for up to 30 days. The ZIO XT Patch device combines both electronic recordation components and physical electrodes into a unitary assembly that adheres to the patient's skin. The ZIO XT Patch device uses adhesive sufficiently strong to support the weight of both the monitor and the electrodes over an extended period of time and to resist disadherence from the patient's body, albeit at the cost of disallowing removal or relocation during the monitoring period. The ZIO Event Card device is a form of downsized Holter monitor with a recorder component that must be removed temporarily during baths or other activities that could damage the non-waterproof electronics. Both devices represent compromises between length of wear and quality of ECG monitoring, especially with respect to ease of long term use, female-friendly fit, and quality of cardiac electrical potential signals, especially atrial (P-wave) signals.

[0008] Therefore, a need remains for a low cost extended wear continuously recording ECG monitor practically capable of being worn for a long period of time, especially in patient's whose breast anatomy can interfere with signal quality in both men and women and that is capable of recording atrial action potential signals reliably.

[0009] Examples of relevant prior art include US 5511553 and US 2004/073127.

DISCLOSURE OF THE INVENTION

[0010] The present application provides an extended wear electrocardiography patch as claimed in claim 1. Advantageous features are provided in the dependent claims.

[0011] Physiological monitoring can be provided through a lightweight wearable monitor that includes two components, a flexible extended wear electrode patch and a reusable monitor recorder that removably snaps into a receptacle on the electrode patch. The wearable monitor sits centrally (in the midline) on the patient's chest along the sternum oriented top-to-bottom. The placement of the wearable monitor in a location at the sternal midline (or immediately to either side of the sternum), with its unique narrow "hourglass"-like shape, significantly improves the ability of the wearable monitor to cuta-

neously sense cardiac electrical potential signals, particularly the P-wave (or atrial activity) and, to a lesser extent, the QRS interval signals indicating ventricular activity in the ECG waveforms.

[0012] The electrode patch is shaped to fit comfortably and conformal to the contours of the patient's chest approximately centered on the sternal midline. To counter the dislodgment due to compressional and torsional forces, a layer of non-irritating adhesive, such as hydrocolloid, is provided at least partially on the underside, or contact, surface of the electrode patch, but only on the electrode patch's distal and proximal ends, where the electrode signal pickups are located. The unadhesed narrowed midsection rides freely over the skin. To counter dislodgment due to tensile and torsional forces, a flexible backing is reinforced with a flexile wire interlaced longitudinally through the narrowed midsection, with the curvature of the flexile wire providing both structural support and malleability. Each of these components are distinctive and allow for comfortable and extended wear, especially for women, where breast mobility would otherwise interfere with monitor use and wearer comfort.

[0013] Moreover, the interlacing of flexile wire simplifies manufacturing and reduces costs. A simple pair of flexile wires are used, instead of custom point-to-point circuit traces, to connect each electrode signal pickup to the receptacle. One end of each flexile wire can be sewn into the receptacle's circuit board, thereby obviating the need for conductive adhesive, soldered or electromechanical connection, and the other end of each flexile wire, when stripped of insulation, can act as an electrode signal pickup, which lowers component count.

[0014] One example provides an extended wear electrocardiography patch using interlaced wire electrodes. A flexible backing is formed of an elongated strip of stretchable material with a narrow longitudinal midsection evenly tapering inward from a distal end and a proximal end, the elongated strip adherable only to a contact surface defined on each of the ends. A pair of flexile wires make up a pair of electrodes, one such electrode comprising one of the flexile wires interlaced into the distal end of the elongated strip and continuing back along an axial path through the narrow longitudinal midsection, another such electrode comprising the other of the flexile wires interlaced into the proximal end of the elongated strip, each of the electrodes further comprising an electrically conductive area only exposed on the contact surface. A circuit board is affixed on the proximal end of the elongated strip, comprising a battery compartment operable to hold a battery for powering an electrocardiography monitor, and an electrical pad operable to electrically couple the electrocardiography monitor with the pair of the electrodes. Alternatively, the circuit board is replaced by a housing that is affixed on the proximal end of the elongated strip, comprising a battery compartment operable to hold a battery for powering an electrocardiography monitor and a plurality of electrical pads operable to electrically couple the electrocardiography monitor with

the pair of the electrodes. The housing can be affixed to the proximal end through the interlacing of the flexile wire. A non-conductive receptacle is securely adhered on the one end of the elongated strip opposite the contact surface and formed to removably receive the electrocardiography monitor, the non-conductive receptacle comprising electrode terminals aligned to electrically interface the pair of the flexile wires to the electrocardiography monitor.

[0015] A further example provides an extended wear electrocardiography patch using non-metal electrodes. A flexible backing is formed of an elongated strip of stretchable material with a narrow longitudinal midsection evenly tapering inward from a distal end and a proximal end, the elongated strip adherable only to a contact surface defined on each of the ends. A pair of flexile wires makes up a pair of electrodes, one such electrode comprising one of the flexile wires positioned on the distal end of the elongated strip on the contact surface and continuing back along an axial path through the narrow longitudinal midsection, another such electrode comprising the other of the flexile wires positioned on the proximal end of the elongated strip on the contact surface, each of the electrodes being embedded in an electrically conductive non-metal adhesive. A circuit board is affixed on the proximal end of the elongated strip, comprising a battery compartment operable to hold a battery for powering an electrocardiography monitor, and an electrical pad operable to electrically couple the electrocardiography monitor with the pair of the electrodes. Alternatively, the circuit board is replaced by a housing that is affixed on the proximal end of the elongated strip, comprising a battery compartment operable to hold a battery for powering an electrocardiography monitor and a plurality of electrical pads operable to electrically couple the electrocardiography monitor with the pair of the electrodes. The housing can be affixed to the proximal end through the interlacing of the flexile wires. A non-conductive receptacle is securely adhered on the one end of the elongated strip opposite the contact surface and formed to removably receive the electrocardiography monitor, the non-conductive receptacle includes electrode terminals aligned to electrically interface the pair of the flexile wires to the electrocardiography monitor.

[0016] A still further embodiment provides a method for constructing a stress-pliant physiological electrode assembly. An electrode backing is formed from a stretchable woven textile material compatible to contact the skin on at least one surface. A length of flexile wire is obtained, the flexile wire having a cross-sectional girth greater than the interstices formed between individual threads comprised in the stretchable woven textile material. An electrical contact area on the at least one surface area is defined, and an electrically-contacting end of the flexile wire is positioned substantially within the electrical contact area. The flexile wire is interlaced through the stretchable woven textile material from the electrical contact area to a surface of the stretchable woven textile material

opposite to the electrical contact area and is operable to receive an electrical connection.

[0017] A still further embodiment provides an extended wear electrocardiography patch. A flexible backing is formed of an elongated strip of stretchable material with a narrow longitudinal midsection evenly tapering inward from both ends. The elongated strip is adherable only on each end of a contact surface to serve as a crimp relief to facilitate compression of the narrow longitudinal midsection in response to compressional and torsional forces. A pair of electrocardiographic electrodes is respectively affixed to and conductively exposed on the contact surface of each end of the elongated strip. A flexible circuit is affixed on each end to the elongated strip. The flexible circuit includes a pair of circuit traces both originating within one of the ends of the elongated strip and which are electrically coupled to each electrocardiographic electrode. A laterally-extendable strain relief is defined in the flexible circuit and formed to facilitate extension and rotation of the flexible circuit in response to tensile and torsional forces. A non-conductive receptacle is securely adhered on the one end of the elongated strip opposite the contact surface and is formed to removably receive an electrocardiography monitor. The non-conductive receptacle includes electrode terminals aligned to electrically interface the pair of circuit traces to the electrocardiography monitor.

[0018] A still further embodiment provides an extended wear hourglass-shaped electrocardiography patch. A flexible backing is formed of an elongated strip of stretchable material with a narrow longitudinal midsection evenly tapering inward from both ends into an hourglass-shaped pattern when viewed from above. A contact surface of the flexible backing includes a non-irritating adhesive dressing at least partially coated on each end and lacking the non-irritating adhesive dressing on the narrow longitudinal midsection. A pair of electrocardiographic electrodes is respectively affixed to and conductively exposed on the contact surface of each end of the flexible backing. A flexible circuit is affixed to each end to the flexible backing. A pair of circuit traces both originate within one of the ends of the flexible backing. Each circuit trace is electrically coupled to one of the electrocardiographic electrodes. A pair of strain relief cutouts each face towards the other strain relief cutout. Each strain relief cutout partially extends transversely from and continuing longitudinally to a side of the flexible circuit different than the other strain relief cutout. One of the pair of circuit traces follows a path formed on the flexible circuit between the pair of strain relief cutouts. A non-conductive receptacle is securely adhered on the one end of the flexible backing opposite the contact surface and is formed to removably receive an electrocardiography monitor.

[0019] The monitoring patch is especially suited to the female anatomy, although also easily used over the male sternum. The narrow longitudinal midsection can fit nicely within the intermammary cleft of the breasts without in-

ducing discomfort, whereas conventional patch electrodes are wide and, if adhered between the breasts, would cause chafing, irritation, discomfort, and annoyance, leading to low patient compliance.

[0020] The foregoing aspects enhance ECG monitoring performance and quality by facilitating long-term ECG recording, which is critical to accurate arrhythmia and cardiac rhythm disorder diagnoses.

[0021] In addition, the foregoing aspects enhance comfort in women (and certain men), but not irritation of the breasts, by placing the monitoring patch in the best location possible for optimizing the recording of cardiac signals from the atrium, particularly P-waves, which is another feature critical to proper arrhythmia and cardiac rhythm disorder diagnoses.

[0022] Further, the interlaced flexile wires improve the dermal electrode's response to tensile, twisting, compressional, and torsional forces by providing a strain relief and tensile strength, while also diminish the cost and complexity of producing physiological electrode assemblies and other types of electrical circuits, where point-to-point interconnections are needed.

[0023] Still other configurations will become readily apparent to those skilled in the art from the following detailed description, wherein are described embodiments by way of illustrating the best mode contemplated. As will be realized, other and different configurations are possible and the embodiments' several details are capable of modifications in various obvious respects. Accordingly, the drawings and detailed description are to be regarded as illustrative in nature.

DESCRIPTION OF THE DRAWINGS

[0024]

FIGURES 1 and 2 are diagrams showing, by way of examples, an extended wear electrocardiography monitor, including an extended wear electrode patch in accordance with one embodiment, respectively fitted to the sternal region of a female patient and a male patient.

FIGURE 3 is a perspective view showing an extended wear electrode patch in accordance with one embodiment with a monitor recorder inserted.

FIGURE 4 is a perspective view showing the extended wear electrode patch of FIGURE 3 without a monitor recorder inserted.

FIGURE 5 is a top view showing the flexible circuit of the extended wear electrode patch of FIGURE 3. FIGURE 6 is a perspective view showing the extended wear electrode patch in accordance with a further configuration.

FIGURE 7 is an exploded view showing the component layers of the electrode patch of FIGURE 3.

FIGURE 8 is a bottom plan view of the extended wear electrode patch of FIGURE 3 with liner partially peeled back.

FIGURE 9 is a perspective view of an extended wear electrode patch with a flexile wire electrode assembly in accordance with a still further configuration.

FIGURE 10 is perspective view of the flexile wire electrode assembly from FIGURE 9, with a layer of insulating material shielding a bare distal wire around the midsection of the flexible backing.

FIGURE 11 is a bottom view of the flexile wire electrode assembly as shown in FIGURE 9.

FIGURE 12 is a bottom view of a flexile wire electrode assembly in accordance with a still yet further configuration.

FIGURE 13 is a perspective view showing the longitudinal midsection of the flexible backing of the electrode assembly from FIGURE 9.

FIGURE 14 is a longitudinal cross-sectional view of the midsection of the flexible backing of the electrode assembly of FIGURE 11.

FIGURES 15A-C are the electrode assembly from FIGURE 14 under compressional, tensile, and bending force, respectively.

FIGURE 16 is a flow diagram showing a method for constructing a stress-pliant physiological electrode assembly in accordance with a further configuration.

BEST MODE FOR CARRYING OUT THE INVENTION

[0025] Physiological monitoring can be provided through a wearable monitor that includes two components, a flexible extended wear electrode patch and a removable reusable monitor recorder. FIGURES 1 and 2 are diagrams showing, by way of examples, an extended wear electrocardiography monitor 12, including an extended wear electrode patch 15 in accordance with one embodiment, respectively fitted to the sternal region of a female patient 10 and a male patient 11. The wearable monitor 12 sits centrally (in the midline) on the patient's chest along the sternum 13 oriented top-to-bottom with the monitor recorder 14 preferably situated towards the patient's head. The electrode patch 15 is shaped to fit comfortably and conformal to the contours of the patient's chest approximately centered on the sternal midline 16 (or immediately to either side of the sternum 13). The distal end of the electrode patch 15 extends towards the Xiphoid process and lower sternum and, depending upon the patient's build, may straddle the region over the Xiphoid process and lower sternum. The proximal end of the electrode patch 15, located under the monitor recorder 14, is below the manubrium and, depending upon patient's build, may straddle the region over the manubrium.

[0026] The placement of the wearable monitor 12 in a location at the sternal midline 16 (or immediately to either side of the sternum 13) significantly improves the ability of the wearable monitor 12 to cutaneously sense cardiac electric signals, particularly the P-wave (or atrial activity) and, to a lesser extent, the QRS interval signals in the ECG waveforms that indicate ventricular activity. The sternum 13 overlies the right atrium of the heart and the

placement of the wearable monitor 12 in the region of the sternal midline 13 puts the ECG electrodes of the electrode patch 15 in a location better adapted to sensing and recording P-wave signals than other placement locations, say, the upper left pectoral region. In addition, placing the lower or inferior pole (ECG electrode) of the electrode patch 15 over (or near) the Xiphoid process and lower sternum facilitates sensing of right ventricular activity and provides superior recordation of the QRS interval.

[0027] During use, the electrode patch 15 is first adhered to the skin along the sternal midline 16 (or immediately to either side of the sternum 13). A monitor recorder 14 is then snapped into place on the electrode patch 15 to initiate ECG monitoring. FIGURE 3 is a perspective view showing an extended wear electrode patch 15 in accordance with one embodiment with a monitor recorder 14 inserted. The body of the electrode patch 15 is preferably constructed using a flexible backing 20 formed as an elongated strip 21 of wrap knit or similar stretchable material about 145mm long and 32mm at the widest point with a narrow longitudinal mid-section 23 evenly tapering inward from both sides. A pair of cut-outs 22 between the distal and proximal ends of the electrode patch 15 create a narrow longitudinal midsection 23 or "isthmus" and defines an elongated "hourglass"-like shape, when viewed from above, such as described in commonly-assigned U.S. Design Patent application, entitled "Extended Wear Electrode Patch," Serial No. 29/472,045, filed November 7, 2013, pending. The upper part of the "hourglass" is sized to allow an electrically non-conductive receptacle 25, sits on top of the outward-facing surface of the electrode patch 15, to be affixed to the electrode patch 15 with an ECG electrode placed underneath on the patient-facing underside, or contact, surface of the electrode patch 15; the upper part of the "hourglass" has a longer and wider profile than the lower part of the "hourglass," which is sized primarily to allow just the placement of an ECG electrode.

[0028] The electrode patch 15 incorporates features that significantly improve wearability, performance, and patient comfort throughout an extended monitoring period. The entire electrode patch 15 is lightweight in construction, which allows the patch to be resilient to disadhesing or falling off and, critically, to avoid creating distracting discomfort to the patient, even when the patient is asleep. In contrast, the weight of a heavy ECG monitor impedes patient mobility and will cause the monitor to constantly tug downwards and press on the patient's body; frequent adjustments by the patient are needed to maintain comfort.

[0029] During every day wear, the electrode patch 15 is subjected to pushing, pulling, and torsional movements, including compressional and torsional forces when the patient bends forward, and tensile and torsional forces when the patient leans backwards. To counter these stress forces, the electrode patch 15 incorporates crimp and strain reliefs, as further described *infra* respec-

tively with reference to FIGURES 4 and 5. In addition, the cut-outs 22 and longitudinal midsection 23 help minimize interference with and discomfort to breast tissue, particularly in women (and gynecomastic men). The cut-outs 22 and longitudinal midsection 23 allow better conformity of the electrode patch 15 to sternal bowing and to the narrow isthmus of flat skin that can occur along the bottom of the intermammary cleft between the breasts, especially in buxom women. The cut-outs 22 and longitudinal midsection 23 help the electrode patch 15 fit nicely between a pair of female breasts in the intermammary cleft. In one embodiment, the cut-outs 22 can be graduated to form the longitudinal midsection 23 as a narrow in-between stem or isthmus portion about 7mm wide. In a still further example, tabs 24 can respectively extend an additional 8mm to 12mm beyond the distal and proximal ends of the flexible backing 20 to facilitate purchase when adhering the electrode patch 15 to or removing the electrode patch 15 from the sternum 13. These tabs preferably lack adhesive on the underside, or contact, surface of the electrode patch 15. Still other shapes, cut-outs and conformities to the electrode patch 15 are possible.

[0030] The monitor recorder 14 removably and reusablely snaps into an electrically non-conductive receptacle 25 during use. The monitor recorder 14 contains electronic circuitry for recording and storing the patient's electrocardiography as sensed via a pair of ECG electrodes provided on the electrode patch 15, such as described in commonly-assigned U.S. Patent application, entitled "Extended Wear Ambulatory Electrocardiography and Physiological Sensor Monitor," Serial No. 14/080,725, filed November 14, 2013, pending. The circuitry includes a microcontroller, flash storage, ECG signal processing, analog-to-digital conversion (where applicable), and an external interface for coupling to the electrode patch 15 and to a download station for stored data download and device programming. The monitor recorder 14 also includes external patient-interfaceable controls, such as a push button to facilitate event marking and provide feedback. In a further example, the circuitry, with the assistance of the appropriate types of deployed electrodes or sensors, is capable of monitoring other types of physiology, in addition to ECGs. Still other types of monitor recorder components and functionality are possible.

[0031] The non-conductive receptacle 25 is provided on the top surface of the flexible backing 20 with a retention catch 26 and tension clip 27 molded into the non-conductive receptacle 25 to conformably receive and securely hold the monitor recorder 14 in place. The edges of the bottom surface of the non-conductive receptacle 25 are preferably rounded, and the monitor recorder 14 is nestled inside the interior of the non-conductive receptacle 25 to present a rounded (gentle) surface, rather than a sharp edge at the skin-to-device interface.

[0032] The electrode patch 15 is intended to be disposable. The monitor recorder 14, however, is reusable and can be transferred to successive electrode patches

15 to ensure continuity of monitoring. The placement of the wearable monitor 12 in a location at the sternal midline 16 (or immediately to either side of the sternum 13) benefits long-term extended wear by removing the requirement that ECG electrodes be continually placed in the same spots on the skin throughout the monitoring period. Instead, the patient is free to place an electrode patch 15 anywhere within the general region of the sternum 13.

[0033] As a result, at any point during ECG monitoring, the patient's skin is able to recover from the wearing of an electrode patch 15, which increases patient comfort and satisfaction, while the monitor recorder 14 ensures ECG monitoring continuity with minimal effort. A monitor recorder 14 is merely unsnapped from a worn out electrode patch 15, the worn out electrode patch 15 is removed from the skin, a new electrode patch 15 is adhered to the skin, possibly in a new spot immediately adjacent to the earlier location, and the same monitor recorder 14 is snapped into the new electrode patch 15 to reinitiate and continue the ECG monitoring.

[0034] During use, the electrode patch 15 is first adhered to the skin in the sternal region. FIGURE 4 is a perspective view showing the extended wear electrode patch 15 of FIGURE 3 without a monitor recorder 14 inserted. A flexible circuit 32 is adhered to each end of the flexible backing 20. A distal circuit trace 33 from the distal end 30 of the flexible backing 20 and a proximal circuit trace (not shown) from the proximal end 31 of the flexible backing 20 electrically couple ECG electrodes (not shown) with a pair of electrical pads 34. In a further example, the distal and proximal circuit traces are replaced with interlaced or sewn-in flexible wires, as further described *infra* beginning with reference to FIGURE 9. The electrical pads 34 are provided within a moisture-resistant seal 35 formed on the bottom surface of the non-conductive receptacle 25. When the monitor recorder 14 is securely received into the non-conductive receptacle 25, that is, snapped into place, the electrical pads 34 interface to electrical contacts (not shown) protruding from the bottom surface of the monitor recorder 14. The moisture-resistant seal 35 enables the monitor recorder 14 to be worn at all times, even during bathing or other activities that could expose the monitor recorder 14 to moisture or adverse conditions.

[0035] In addition, a battery compartment 36 is formed on the bottom surface of the non-conductive receptacle 25. A pair of battery leads (not shown) from the battery compartment 36 to another pair of the electrical pads 34 electrically interface the battery to the monitor recorder 14. The battery contained within the battery compartment 35 can be replaceable, rechargeable or disposable.

[0036] The monitor recorder 14 draws power externally from the battery provided in the non-conductive receptacle 25, thereby uniquely obviating the need for the monitor recorder 14 to carry a dedicated power source. The battery contained within the battery compartment 36 can be replaceable, rechargeable or disposable. In a further

example, the ECG sensing circuitry of the monitor recorder 14 can be supplemented with additional sensors, including an SpO₂ sensor, a blood pressure sensor, a temperature sensor, respiratory rate sensor, a glucose sensor, an air flow sensor, and a volumetric pressure sensor, which can be incorporated directly into the monitor recorder 14 or onto the non-conductive receptacle 25.

[0037] The placement of the flexible backing 20 on the sternal midline 16 (or immediately to either side of the sternum 13) also helps to minimize the side-to-side movement of the wearable monitor 12 in the left- and right-handed directions during wear. However, the wearable monitor 12 is still susceptible to pushing, pulling, and torqueing movements, including compressional and torsional forces when the patient bends forward, and tensile and torsional forces when the patient leans backwards. To counter the dislodgment of the flexible backing 20 due to compressional and torsional forces, a layer of non-irritating adhesive, such as hydrocolloid, is provided at least partially on the underside, or contact, surface of the flexible backing 20, but only on the distal end 30 and the proximal end 31. As a result, the underside, or contact surface of the longitudinal midsection 23 does not have an adhesive layer and remains free to move relative to the skin. Thus, the longitudinal midsection 23 forms a crimp relief that respectively facilitates compression and twisting of the flexible backing 20 in response to compressional and torsional forces. Other forms of flexible backing crimp reliefs are possible.

[0038] Unlike the flexible backing 20, the flexible circuit 32 is only able to bend and cannot stretch in a planar direction. FIGURE 5 is a top view showing the flexible circuit 32 of the extended wear electrode patch 15 of FIGURE 3. A distal ECG electrode 38 and proximal ECG electrode 39 are respectively coupled to the distal and proximal ends of the flexible circuit 32 to serve as electrode signal pickups. The flexible circuit 32 preferably does not extend to the outside edges of the flexible backing 20, thereby avoiding gouging or discomforting the patient's skin during extended wear, such as when sleeping on the side. During wear, the ECG electrodes 38, 39 must remain in continual contact with the skin. A strain relief 40 is defined in the flexible circuit 32 at a location that is partially underneath the battery compartment 36 when the flexible circuit 32 is affixed to the flexible backing 20. The strain relief 40 is laterally extendable to counter dislodgment of the ECG electrodes 38, 39 due to tensile and torsional forces. A pair of strain relief cutouts 41 partially extend transversely from each opposite side of the flexible circuit 32 and continue longitudinally towards each other to define in 'S'-shaped pattern, when viewed from above. The strain relief respectively facilitates longitudinal extension and twisting of the flexible circuit 32 in response to tensile and torsional forces. Other forms of circuit board strain relief are possible.

[0039] The flexible circuit 32 can be provided either above or below the flexible backing 20. FIGURE 6 is a perspective view showing the extended wear electrode

patch 15 in accordance with a further example. The flexible circuit (not shown) is provided on the underside, or contact, surface of the flexible backing 20 and is electrically interfaced to the set of electrical pads 34 on the bottom surface of the non-conductive receptacle 25 through electrical contacts (not shown) pierced through the flexible backing 20.

[0040] The electrode patch 15 is intended to be a disposable component, which enables a patient to replace the electrode patch 15 as needed throughout the monitoring period, while maintaining continuity of physiological sensing through reuse of the same monitor recorder 14. FIGURE 7 is an exploded view showing the component layers of the electrode patch 15 of FIGURE 3. The flexible backing 20 is constructed of a wearable gauze, latex, woven textile, or similar wrap knit or stretchable and wear-safe material 44, such as a Tricot-type linen with a pressure sensitive adhesive (PSA) on the underside, or contact, surface. The ends of the wearable material 44 are coated with a layer 43 of non-irritating adhesive, such as hydrocolloid, to facilitate long-term wear, while the unadhesed narrowed midsection rides freely over the skin. The hydrocolloid, for instance, is typically made of mineral oil, cellulose and water and lacks any chemical solvents, so should cause little itching or irritation. Moreover, hydrocolloid can be manufactured into an appropriate thickness and plasticity and provides cushioning between the relatively rigid and unyielding non-conductive receptacle 25 and the patient's skin. In a further example, the layer of non-irritating adhesive can be contoured, such as by forming the adhesive with a concave or convex cross-section; surfaced, such as through stripes or crosshatches of adhesive, or by forming dimples in the adhesive's surface; or applied discontinuously, such as with a formation of discrete dots of adhesive.

[0041] As described *supra* with reference to FIGURE 5, a flexible circuit can be adhered to either the outward facing surface or the underside, or contact, surface of the flexible backing 20. For convenience, a flexible circuit 47 is shown relative to the outward facing surface of the wearable material 44 and is adhered respectively on a distal end by a distal electrode seal 45 and on a proximal end by a proximal electrode seal 45. In a further embodiment, the flexible circuit 47 can be provided on the underside, or contact, surface of the wearable material 44. Through the electrode seals, only the distal and proximal ends of the flexible circuit 47 are attached to the wearable material 44, which enables the strain relief 40 (shown in FIGURE 5) to respectively longitudinally extend and twist in response to tensile and torsional forces during wear. Similarly, the layer 43 of non-irritating adhesive is provided on the underside, or contact, surface of the wearable material 44 only on the proximal and distal ends, which enables the longitudinal midsection 23 (shown in FIGURE 3) to respectively bow outward and away from the sternum 13 or twist in response to compressional and torsional forces during wear.

[0042] A pair of openings 46 is defined on the distal and proximal ends of the wearable material 44 and layer 43 of non-irritating adhesive for ECG electrodes 38, 39 (shown in FIGURE 5). The openings 46 serve as "gel" wells with a layer of hydrogel 41 being used to fill the bottom of each opening 46 as a conductive material that aids electrode signal capture. The entire underside, or contact, surface of the flexible backing 20 is protected prior to use by a liner layer 40 that is peeled away, as shown in FIGURE 8.

[0043] The non-conductive receptacle 25 includes a main body 54 that is molded out of polycarbonate, ABS, or an alloy of those two materials to provide a high surface energy to facilitate adhesion of an adhesive seal 53. The main body 54 is attached to a battery printed circuit board 52 by the adhesive seal 53 and, in turn, the battery printed circuit board 52 is adhered to the flexible circuit 47 with an upper flexible circuit seal 50. A pair of conductive transfer adhesive points 51 or, alternatively, soldered connections, or electromechanical connections, including metallic rivets or similar conductive and structurally unifying components, connect the circuit traces 33, 37 (shown in FIGURE 5) of the flexible circuit 47 to the battery printed circuit board 52. The main body 54 has a retention catch 26 and tension clip 27 (shown in FIGURE 3) that fixably and securely receive a monitor recorder 14 (not shown), and includes a recess within which to circumferentially receive a die cut gasket 55, either rubber, urethane foam, or similar suitable material, to provide a moisture resistant seal to the set of pads 34. Other types of design, arrangement, and permutation are possible.

[0044] In a still further example, the flexible circuit 32 (shown in FIGURE 4) and distal ECG electrode 38 and proximal ECG electrode 39 (shown in FIGURE 5) are replaced with a pair of interlaced flexile wires. The interlacing of flexile wires through the flexible backing 20 reduces both manufacturing costs and environmental impact, as further described *infra*. The flexible circuit and ECG electrodes are replaced with a pair of flexile wires that serve as both electrode circuit traces and electrode signal pickups. FIGURE 9 is a perspective view of an extended wear electrode patch 15 with a flexile wire electrode assembly in accordance with a still further embodiment. The flexible backing 20 maintains the unique narrow "hourglass"-like shape that aids long term extended wear, particularly in women, as described *supra* with reference to FIGURE 3. For clarity, the non-conductive receptacle 25 is omitted to show the exposed battery printed circuit board 62 that is adhered underneath the non-conductive receptacle 25 to the proximal end 31 of the flexible backing 20. Instead of employing flexible circuits, a pair of flexile wires are separately interlaced or sewn into the flexible backing 20 to serve as circuit connections for an anode electrode lead and for a cathode electrode lead.

[0045] To form a distal electrode assembly, a distal wire 61 is interlaced into the distal end 30 of the flexible

backing 20, continues along an axial path through the narrow longitudinal midsection of the elongated strip, and electrically connects to the battery printed circuit board 62 on the proximal end 31 of the flexible backing 20. The distal wire 61 is connected to the battery printed circuit board 62 by stripping the distal wire 61 of insulation, if applicable, and interlacing or sewing the uninsulated end of the distal wire 61 directly into an exposed circuit trace 63. The distal wire-to-battery printed circuit board connection can be made, for instance, by back stitching the distal wire 61 back and forth across the edge of the battery printed circuit board 62. Similarly, to form a proximal electrode assembly, a proximal wire (not shown) is interlaced into the proximal end 31 of the flexible backing 20. The proximal wire is connected to the battery printed circuit board 62 by stripping the proximal wire of insulation, if applicable, and interlacing or sewing the uninsulated end of the proximal wire directly into an exposed circuit trace 64. The resulting flexile wire connections both establish electrical connections and help to affix the battery printed circuit board 62 to the flexible backing 20.

[0046] The battery printed circuit board 62 is provided with a battery compartment 36. A set of electrical pads 34 are formed on the battery printed circuit board 62. The electrical pads 34 electrically interface the battery printed circuit board 62 with a monitor recorder 14 when fitted into the non-conductive receptacle 25. The battery compartment 36 contains a spring 65 and a clasp 66, or similar assembly, to hold a battery (not shown) in place and electrically interfaces the battery to the electrical pads 34 through a pair battery leads 67 for powering the electrocardiography monitor 14. Other types of battery compartment are possible. The battery contained within the battery compartment 36 can be replaceable, rechargeable, or disposable.

[0047] In a yet further embodiment, the circuit board and non-conductive receptacle 25 are replaced by a combined housing that includes a battery compartment and a plurality of electrical pads. The housing can be affixed to the proximal end of the elongated strip through the interlacing or sewing of the flexile wires or other wires or threads.

[0048] The core of the flexile wires may be made from a solid, stranded, or braided conductive metal or metal compounds. In general, a solid wire will be less flexible than a stranded wire with the same total cross-sectional area, but will provide more mechanical rigidity than the stranded wire. The conductive core may be copper, aluminum, silver, or other material. The pair of the flexile wires may be provided as insulated wire. In one example, the flexile wires are made from a magnet wire from Belden Cable, catalogue number 8051, with a solid core of AWG 22 with bare copper as conductor material and insulated by polyurethane or nylon. Still other types of flexile wires are possible. In a further example, conductive ink or graphene can be used to print electrical connections, either in combination with or in place of the flexile wires.

[0049] In a still further example, the flexile wires are uninsulated. FIGURE 10 is perspective view of the flexile wire electrode assembly from FIGURE 9, with a layer of insulating material 69 shielding a bare uninsulated distal wire 61 around the midsection on the contact side of the flexible backing. On the contact side of the proximal and distal ends of the flexible backing, only the portions of the flexile wires serving as electrode signal pickups are electrically exposed and the rest of the flexile wire on the contact side outside of the proximal and distal ends are shielded from electrical contact. The bare uninsulated distal wire 61 may be insulated using a layer of plastic, rubber-like polymers, or varnish, or by an additional layer of gauze or adhesive (or non-adhesive) gel. The bare uninsulated wire 61 on the non-contact side of the flexible backing may be insulated or can simply be left uninsulated.

[0050] Both end portions of the pair of flexile wires are typically placed uninsulated on the contact surface of the flexible backing 20 to form a pair of electrode signal pickups. FIGURE 11 is a bottom view of the flexile wire electrode assembly as shown in FIGURE 9. When adhered to the skin during use, the uninsulated end portions of the distal wire 61 and the proximal wire 71 enable the monitor recorder 14 to measure dermal electrical potential differentials. At the proximal and distal ends of the flexible backing 20, the uninsulated end portions of the flexile wires may be configured into an appropriate pattern to provide an electrode signal pickup, which would typically be a spiral shape formed by guiding the flexile wire along an inwardly spiraling pattern. The surface area of the electrode pickups can also be variable, such as by selectively removing some or all of the insulation on the contact surface. For example, an electrode signal pickup arranged by sewing insulated flexile wire in a spiral pattern could have a crescent-shaped cutout of uninsulated flexile wire facing towards the signal source.

[0051] In a still yet further example, the flexile wires are left freely riding on the contact surfaces on the distal and proximal ends of the flexible backing, rather than being interlaced into the ends of the flexible backing 20. FIGURE 12 is a bottom view of a flexile wire electrode assembly in accordance with a still yet further example. The distal wire 61 is interlaced onto the midsection and extends an exposed end portion 72 onto the distal end 30. The proximal wire 71 extends an exposed end portion 73 onto the proximal end 31. The exposed end portions 72 and 73, not shielded with insulation, are further embedded within an electrically conductive adhesive 81. The adhesive 81 makes contact to skin during use and conducts skin electrical potentials to the monitor recorder 14 (not shown) via the flexile wires. The adhesive 81 can be formed from electrically conductive, non-irritating adhesive, such as hydrocolloid.

[0052] The distal wire 61 is interlaced or sewn through the longitudinal midsection of the flexible backing 20 and takes the place of the flexible circuit 32. FIGURE 13 is a perspective view showing the longitudinal midsection of

the flexible backing of the electrode assembly from FIGURE 9. Various stitching patterns may be adopted to provide a proper combination of rigidity and flexibility. In simplest form, the distal wire 61 can be manually threaded through a plurality of holes provided at regularly-spaced intervals along an axial path defined between the battery printed circuit board 62 (not shown) and the distal end 30 of the flexible backing 20. The distal wire 61 can be threaded through the plurality of holes by stitching the flexile wire as a single "thread." Other types of stitching patterns or stitching of multiple "threads" could also be used, as well as using a sewing machine or similar device to machine-stitch the distal wire 61 into place, as further described *infra*. Further, the path of the distal wire 61 need not be limited to a straight line from the distal to the proximal end of the flexible backing 20.

[0053] The distal wire 61 is flexile yet still retains a degree of rigidity that is influenced by wire gauge, composition, stranding, insulation, and stitching pattern. For example, rigidity decreases with wire gauge; and a solid core wire tends to be more rigid than a stranded core of the same gauge. The combination of the flexibility and the rigidity of the portion of the distal wire 61 located on or close to the midsection contributes to the overall strength and wearability of the patch. FIGURE 14 is a longitudinal cross-sectional view of the midsection of the flexible backing 20 of the electrode assembly of FIGURE 11. FIGURES 15A-C are the electrode assembly from FIGURE 14 under compressional, tensile, and bending force, respectively. The relative sizes of the distal wire 61 and flexible backing 20 are not to scale and are exaggerated for purposes of illustration.

[0054] The interlacing of the distal wire 61 through the narrow longitudinal midsection 22 of the flexible backing 20 bends the distal wire 61 into a line of rounded stitches that alternate top and bottom, which can be advantageous to long term wearability. First, the tension of the rounded stitches reinforces the planar structure of the narrow longitudinal midsection 22 and spreads a dislodging force impacting on one end of the flexible backing 20 to the other end of the flexible backing 20. Second, the rounded stitches leave room for stretching, compressing, bending, and twisting, thus increasing the wearability of the patch extended wear electrode patch 15 by facilitating extension, compression, bending, and twisting of the narrow longitudinal midsection 22 in response to tensile, compressional, bending, and torsional forces.

[0055] In a further example, the distal wire and the proximal wire may be stitched or sewn into the flexible backing 20. Depending upon the type of stitching used, the distal or proximal wire may use more than one individual wire. For instance, a conventional sewing machine used to stitch fabrics uses a spool of thread and a bobbin, which are both wound with thread that together allow the creation of various stitching patterns, such as the lock-stitch. Other type of stitching patterns are possible. Additionally, where more than one "threads" are used for stitching, the flexile wire may constitute all of the

"threads," thereby increasing redundancy of the circuit trace thus formed. Alternatively, just one (or fewer than all) of the threads may be conductive, with the non-conductive threads serving to reinforce the strength of the flexile wire connections and flexible backing 20. The additional threads can be made from line, threads, or fabrics of sufficient mechanical strength and do not need to be conductive; alternatively, the same flexile wires can be employed to serve as the additional threads.

[0056] Conventionally, flexible circuits, such as the flexible circuit 32 (shown in FIGURE 4) that connects the distal ECG electrode 38 and proximal ECG electrode 39 (shown in FIGURE 5) to the battery printed circuit board 62 (shown in FIGURE 9), are constructed using subtractive processes. In general, a flexible circuit interconnects electronic components with custom point-to-point circuit traces and is typically constructed by forming the conductive circuit traces on a thin film of insulating polymer. A flexible circuit is not an off-the-shelf component; rather, each flexible circuit is designed with a specific purpose in mind. Changes to a flexible circuit's design will generally require fabricating entirely new flexible circuits, as the physical circuit traces on the polymer film cannot be changed.

[0057] Manufacturing a flexible circuit typically requires the use of sophisticated and specialized tools, coupled with environmentally unfriendly processes, including depositing copper on a polyamide core, etching away unwanted copper with inline etching or an acid bath to retain only the desired conductive circuit traces, and applying a coverlay to the resulting flexible circuit. Significant amounts of hazardous waste are generated by these subtractive processes during the fabrication of each flexible circuit. Properly disposing of such hazardous waste is expensive and adds to the costs of the flexible circuit.

[0058] In the still further example described *supra* beginning with reference to FIGURE 9, the distal and proximal flexile wires replace the flexible circuit 32 and enables the electrode assembly to be constructed using additive processes with off-the-shelf, low cost components. The flexile wires serve the triple functions of an electrode signal pickup, electrical circuit trace, and support for structural integrity and malleability of the electrode assembly.

[0059] The general manner of constructing the electrode assembly can be applied to other forms of electronic components in which custom point-to-point circuit traces need to be affixed to a gauze or textile backing, as well as backings made from other materials. The circuit traces are replaced by the interlaced or sewn flexile wires, and the ends of each flexile wire are terminated, as appropriate to the application. The flexile wires may, by example, connect two circuit boards, or connect to an electrical terminal, power source, or electrical component. In addition, flexile wires may be used to replace a printed circuit board entirely, with each flexile wire serving as a form of sewn interconnect between two or more discrete com-

ponents, including resistors, capacitors, transistors, diodes, operational amplifiers (op amps) and other integrated circuits, and other electronic or electromechanical components.

[0060] By way of illustration, the flexile wires will be described as terminated for use in an electrode assembly, specifically, as terminated on one end to form an electrode signal pickup and on the other end to connect into a circuit board. Constructing the electrode assembly entails interlacing, including manually threading, or machine sewing the flexile, conductive wire through the flexible backing 20. FIGURE 16 is a flow diagram showing a method 90 for constructing a stress-pliant physiological electrode assembly in accordance with a further example. The method can be performed by a set of industrial machines, including a gauze cutting machine to cut the flexible backing 20 to form; a hole punch to cut a plurality of holes provided at regularly-spaced intervals; a stitching or sewing machine to interleave or sew the flexile wire through the flexible backing 20; a wire stripper or plasma jet to remove insulation from the flexile wire, when applicable; and a glue or adhesive dispenser to embed or coat electrode signal pickup in hydrocolloid gel or equivalent non-irritating adhesive. Other forms or combinations of industrial machines, including a single purpose-built industrial machine, could be used.

[0061] As an initial step, a backing is cut to shape and, if required, holes are cut at regularly-spaced intervals along an axial path (step 91) through which the flexile wire will be interlaced. Holes will need to be cut, for instance, if the flexile wire is to be hand-guided through the backing, or where the backing is cut from a material that is difficult to puncture with a threaded needle, such as used by a sewing machine. In one example, the backing is cut from wearable gauze, latex, woven textile, or similar wrap knit or stretchable and wear-safe material, such as a Tricot-type linen; the resulting backing is flexible and yielding. The backing is also cut into an elongated "hourglass"-like shape, when viewed from above, with a pair of cut-outs and a longitudinal midsection that together help minimize interference with and discomfort to breast tissue, particularly in women (and gynecomastic men), such as described *supra* with reference to FIGURE 3. The backing can be cut into other shapes as appropriate to need. In addition, depending upon the application, other materials could be substituted for the backing. For example, neoprene, such as used in wetsuits, could be used where a high degree of elasticity and ruggedness is desired.

[0062] The flexile wire is then interlaced or sewn into the backing (step 92). Interlacing can be performed by a machine that guides the flexile wire through the holes previously cut in the material in a crisscrossed, interwoven, or knitted fashion, as well as by hand. The flexile wire can also be guided through the backing without first cutting holes, provided that the weave of the material is sufficiently loose to allow passage of the flexile wire if the flexile wire is otherwise incapable of passing through the

backing without the assistance of a needle or other piercing instrument.

[0063] Alternatively, the flexile wire could be sewn into the backing by using the flexile wire as "thread" that is stitched into place using a needle or similar implement. If a single flexile wire is employed, the stitching will be a line of rounded stitches that alternate top and bottom, as described *supra*; however, if more than one flexile wire is used, or the stitching pattern requires the use of more than one thread, other forms of conventional machine-stitching patterns could be employed, such as a lock-stitch.

[0064] Once completed, the interlacing or sewing of the flexile wire into the backing creates an integrated point-to-point electrical path that takes the place of a custom circuit trace using an additive, rather than subtractive, manufacturing process. The flexile wire can be interlaced or sewn along a straight, curved, or arbitrary path. One flexile wire is required per point-to-point circuit trace. The strength and pliability of the flexile wire reinforces the backing and, in the still further example described *supra* beginning with reference to FIGURE 9, facilitates extension, compression, bending, and twisting of the narrow longitudinal midsection 22 in response to tensile, compressional, bending, and torsional forces. Thus, the path of the flexile wire along the backing can be mapped to take advantage of the strength and reinforcing properties of the flexile wire, which, when interlaced or sewn into the backing, help the backing counter the stresses to which the backing will be subjected when deployed.

[0065] The flexile wire itself may be insulated or bare (step 93). When one end of the flexile wire is connected to (or forms) an electrode, particularly a dermal physiology electrode that senses electrical potentials on the skin's surface, insulated flexile wire will ordinarily be used, with only a portion of the flexile wire incident to the electrode stripped of insulation. However, bare uninsulated flexile wire could alternatively be used throughout, so long as those portions of the uninsulated flexile wire that are exposed on the contact-facing surface of the backing are insulated and shielded from electrical contact (step 94), such as by applying a layer of plastic, rubber-like polymers, or varnish, or by an additional layer of gauze or adhesive (or non-adhesive) gel over the exposed wire. The uninsulated flexile wire exposed on other surfaces of the backing could also be insulated or simply be left bare.

[0066] One end of the flexile wire may be terminated as an electrode signal pickup (step 95). If insulated flexile wire is used, a portion of the end of the flexile wire is stripped of insulation (step 96) using, for instance, a wire stripper or plasma jet. The electrode signal pickup could either be formed by interlacing (or sewing) the flexile wire (step 97) into the backing in the shape of the desired electrode (step 98) or positioned over the contact-facing area of the backing designated to serve as an electrode signal pickup and embedded within an electrically con-

ductive adhesive (step 99). In a yet further example, the flexile wire could be terminated as a connection to a discrete electrode, such as by sewing an uninsulated portion of the end of the electrode wire into the discrete electrode to thereby establish an electrical contact and affix the discrete electrode to the backing. The Universal ECG EKG electrode, manufactured by Bio Protech Inc., Tustin, CA, is one example of a discrete electrode.

[0067] Finally, the other end of the flexile wire may be terminated as a connection to a circuit board (step 100). The flexile wire can be interlaced or sewn onto the circuit board, for instance, by back stitching the flexile wire back and forth across the edge of the circuit board to thereby establish an electrical contact and affix the discrete electrode to the backing.

[0068] In a further embodiment, flexile wire can be used to replace all or part of a printed circuit board, such as battery printed circuit board 62 used in constructing a stress-pliant physiological electrode assembly, as described supra, or for any other application that requires interconnection of electrical or electro mechanical components on a physical substrate or backing. Flexile wire in place of conductive circuit traces can work especially well with simple circuit board layouts, where ample space between components and relatively uncomplicated layouts are amenable to stitched-in interconnections. In addition, the use of flexile wire can simplify circuit layout design in multilayer circuits, as insulated flexile wires can be run across each other in situations that would otherwise require the use of a multilayer printed circuit board or similar solution.

[0069] Through such use of flexile wire, a printed circuit board can be omitted in whole or in part. Interconnects between and connections to the electronic and electro mechanical components formerly placed on the printed circuit board can instead be sewn from flexile wire. For instance, the battery printed circuit board 62 can be replaced by flexile wire interconnects that connect the electrodes to a sewn set of electrical pads formed by over-stitching the flexile wire into electrical contact surfaces of sufficient size to interface with a monitor recorder 14 when fitted into the non-conductive receptacle 25. Likewise, the spring 65 and clasp 66 can be sewn in place using flexile wire to hold a battery in place with flexile wire interconnects connecting the battery to a sewn set of electrical pads formed by over-stitching the flexile wire into electrical contact surfaces of sufficient size to interface with a monitor recorder 14 when fitted into the non-conductive receptacle 25. Still other approaches to replacing printed circuit boards with flexile wire interconnects are possible.

[0070] The resultant stress-pliant physiological electrode assembly may be electrically coupled to a broad range of physiological monitors not limited to electrocardiographic measurement. The foregoing method of constructing a stress-pliant electrode assembly is adaptable to manufacturing other forms of dermal electrodes, including electrodes for electrocardiography, electroen-

cephalography, and skin conductance measurements. Further, by adjusting the number of electrodes, the distances among the electrode signal pickups, and the thickness of the flexile wire, the method can be adapted to manufacturing at low cost an electrode assembly that is lightweight and resistant to tensile, compressional and torsional forces, thus contributing to long-term wear and versatility.

[0071] While the invention has been particularly shown and described as referenced to the embodiments thereof, those skilled in the art will understand that the foregoing and other changes in form and detail may be made therein without departing from the scope.

Claims

1. An extended wear electrocardiography patch (15), comprising:

a flexible backing (20) formed of an elongated strip (21) of stretchable material (44) with a narrow longitudinal midsection (23) evenly tapering inward from both ends (30, 31), the elongated strip (21) adherable only on a contact surface defined on each of the ends (30, 31), and operable to serve as a crimp relief to facilitate compression of the narrow longitudinal midsection (23) in response to compressional and torsional forces;

a flexible circuit (32) affixed on each end (30, 31) to the elongated strip (21) and comprising a pair of circuit traces (33, 37) each originating within one of the ends of the elongated strip (21), respectively;

a non-conductive receptacle (25) comprising a plurality of electrical pads and securely adhered on one end (31) of the elongated strip (21) opposite the contact surface and operable to removably receive an electrocardiography monitor (14), the non-conductive receptacle (25) aligned to electrically interface the pair of the circuit traces (33, 37) to the electrocardiography monitor (14);

a pair of electrocardiographic electrodes (38, 39) respectively affixed to and conductively exposed on the contact surface of each end (30, 31) of the elongated strip (21), and electrically coupled with a pair of the electrical pads (34) on the non-conductive receptacle and the pair of circuit traces;

a battery compartment formed on a bottom surface of the non-conductive receptacle; and
a battery located within the battery compartment of the non-conductive receptacle and electrically interfaced to a different pair of the electrical pads (34) of the non-conductive receptacle.

2. An electrocardiography patch (15) according to Claim 1, further comprising:

one of the electrocardiographic electrodes (38) being disposed for being adhered to a region overlying the lower sternum on a patient's chest; and
an other of the electrocardiographic electrodes (39) being disposed for being adhered to the region near the manubrium on the patient's chest oriented centrally (in the midline (16)) along the sternum (13) upwards from the one electrocardiographic electrode (38).

3. An electrocardiography patch (15) according to Claim 1, further comprising:
a non-irritating adhesive dressing (43) at least partially coated on each end of the elongated strip (21) on the contact surface.

4. An electrocardiography patch (15) according to Claim 1, further comprising at least one of:

a laterally-extendable strain relief (40) defined in the flexible circuit (32) and operable to facilitate extension and rotation of the flexible circuit (32) in response to tensile and torsional forces; and

a pair of strain relief cutouts (41) comprised in the strain relief (40) that each face towards the other strain relief cutout (41), each strain relief cutout (41) partially extending transversely from and continuing longitudinally to a side of the flexible circuit (32) different than the other strain relief cutout (41), one of the pair of the circuit traces (33, 37) following a path formed on the flexible circuit (32) between the pair of the strain relief cutouts (41).

5. An electrocardiography patch (15) according to Claim 4, wherein the pair of the strain relief cutouts (41) define an 'S'-shaped pattern in the flexible circuit (32) when viewed from above.

6. An electrocardiography patch (15) to Claim 1, wherein the narrow longitudinal midsection (23) defines an hourglass-shaped pattern in the flexible backing (20) when viewed from above conformal to fit comfortably within the intermammary cleft of a patient's pair of breasts.

7. An electrocardiography patch (15) according to Claim 1, further comprising:
a moisture-resistant seal (35) formed on the non-conductive receptacle (25) circumferentially about the electrode terminals.

8. An electrocardiography patch (15) according to

Claim 3, further comprising:

a contoured surface comprised on the non-irritating adhesive dressing (43) on at least one end of the elongated strip (21).

9. An electrocardiography patch (15) according to Claim 1, further comprising:

a plurality of fasteners comprised on the non-conductive receptacle (25) and oriented to securely receive and to hold captive the electrocardiography monitor (14) into the non-conductive receptacle (25).

10. An electrocardiography patch according to Claim 1, further comprising at least one of:

rounded edges defined about a bottom surface of the non-conductive receptacle (25);

a gel well (46) circumferentially defined about at least one of the electrocardiographic electrodes (38, 39) on the contact surface of the end (30, 31) of the flexible backing (20) on which the at least one electrocardiographic electrode (38, 39) is affixed; and

placement and removal tabs (24) comprised as part of the flexible backing (20) and extending longitudinally beyond at least one end (30, 31) of the flexible backing (20).

11. An electrocardiography patch (15) according to Claim 1 wherein the patch is an hourglass shaped pattern when viewed from above;

an upper part of the hourglass-shaped pattern sized to seat the non-conductive receptacle (25) on an outward-facing surface of the flexible backing (20); and
a lower part of the hourglass-shaped pattern comprising a size smaller than the upper part of the hourglass-shaped pattern.

12. An electrocardiography patch (15) according to Claim 1, further comprising:

the non-conductive receptacle (25) comprising electrode terminals aligned to electrically interface the electrocardiographic electrodes (38, 39) via the pair of the circuit traces (33, 37) to the electrocardiography monitor (14).

13. An electrocardiography patch (15) according to Claim 1, further comprising:

a pair of concave cut-outs (22) defined along the longitudinal edges of the elongated strip (21), each concave cut-out (22) contoured conformal to substantially clear an intermammary cleft.

14. An electrocardiography patch (15) according to Claim 1, wherein the stretchable material (44) comprises at least one of wearable gauze, wearable latex, wrap knit fabric, and Tricot linen.

15. A system comprising an electrocardiography patch (15) according to Claim 1 and an electrocardiography monitor, wherein a patient feedback button is provided on the electrocardiography monitor (14).

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Patentansprüche

1. Elektrokardiografiepflaster für längeres Tragen (15), umfassend:

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einen flexiblen Träger (20), der aus einem länglichen Streifen (21) aus dehnbarem Material (44) mit einem engen Längsmittelteil (23), der sich gleichmäßig von beiden Enden (30, 31) nach innen verjüngt, gebildet ist, wobei der längliche Streifen (21) nur auf einer an jedem der Enden (30, 31) definierten Kontaktfläche anhaftbar ist und funktionsfähig ist, um als eine Klemmentlastung zu dienen, um als Reaktion auf Kompressions- und Drehkräfte eine Komprimierung des engen Längsmittelteils (23) zu ermöglichen;

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einen flexiblen Schaltkreis (32), der an jedem Ende (30, 31) am länglichen Streifen (21) befestigt ist und ein Paar von Schaltkreisspuren (33, 37) umfasst, die jeweils innerhalb der Enden des länglichen Streifens (21) ausgehen; einen nichtleitenden Behälter (25), der eine Vielzahl von elektrischen Anschlussfeldern umfasst und sicher an einem Ende (31) des länglichen Streifens (21) gegenüber der Kontaktfläche angeheftet ist und funktionsfähig ist, um einen Elektrokardiografiewatch (14) entfernt aufzunehmen, wobei der nichtleitende Behälter (25) ausgerichtet ist, das Schaltkreisspurenpaar (33, 37) elektrisch an den Elektrokardiografiewatch (14) zu koppeln;

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ein Paar von Elektrokardiografieelektroden (38, 39), die jeweils an der Kontaktfläche jedes Endes (30, 31) des länglichen Streifens (21) befestigt und leitend freigesetzt sind und elektrisch an ein Paar der elektrischen Anschlussfelder (34) am nichtleitenden Behälter und das Paar von Schaltkreisspuren gekoppelt sind;

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ein Batteriefach, das an einer Unterseite des nichtleitenden Behälters gebildet ist; und eine Batterie, die sich im Inneren des Batteriefachs des nichtleitenden Behälters befindet und elektrisch an ein anderes Paar der elektrischen Anschlussfelder (34) des nichtleitenden Behälters gekoppelt ist.

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2. Elektrokardiografiepflaster (15) nach Anspruch 1, ferner umfassend:

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dass eine der Elektrokardiografieelektroden (38) angeordnet ist, um an einen Bereich ange-

heftet zu werden, der über dem unteren Sternum auf der Brust eines Patienten liegt; und dass eine andere der Elektrokardiografieelektroden (39) angeordnet ist, um am Bereich in der Nähe des Manubriums auf der Brust des Patienten, zentral (auf der Mittellinie (16)) entlang des Sternums (13) ausgerichtet nach oben von der einen Elektrokardiografieelektrode (38) angeheftet zu werden.

3. Elektrokardiografiepflaster (15) nach Anspruch 1, ferner umfassend:

einen nicht reizenden Klebeverband (43), der zumindest teilweise an jedem Ende des länglichen Streifens (21) an der Kontaktfläche beschichtet ist.

4. Elektrokardiografiepflaster (15) nach Anspruch 1, ferner umfassend mindestens eines von:

eine lateral streckbare Zugentlastung (40), die im flexiblen Schaltkreis (32) definiert ist und funktionsfähig ist, eine Streckung und Drehung des flexiblen Schaltkreises (32) als Reaktion auf Zug- und Drehkräfte zu ermöglichen; und ein Paar von Zugentlastungsaussparungen (41), die in der Zugentlastung (40) enthalten sind und jeweils zur anderen Zugentlastungsaussparung (41) zeigen, wobei jede Zugentlastungsaussparung (41) teilweise transversal von und in Längsrichtung zu einer jeweils anderen Seite des flexiblen Schaltkreises (32) als die andere Zugentlastungsaussparung (41) verläuft, wobei eine des Paares der Schaltkreisspuren (33, 37) einem Pfad folgt, der auf dem flexiblen Schaltkreis (32) zwischen dem Paar der Zugentlastungsaussparungen (41) gebildet ist.

5. Elektrokardiografiepflaster (15) nach Anspruch 4, wobei das Paar von Zugentlastungsaussparungen (41) in der Draufsicht eine 'S'-Form im flexiblen Schaltkreis (32) definiert.

6. Elektrokardiografiepflaster (15) nach Anspruch 1, wobei der enge Längsmittelteil (23) in der Draufsicht eine Sanduhrform im flexiblen Träger (20) definiert, die so geformt ist, dass er bequem in den intermamären Spalt eines Brustpaares eines Patienten passt.

7. Elektrokardiografiepflaster (15) nach Anspruch 1, ferner umfassend:

eine feuchtigkeitsbeständige Dichtung (35), die auf dem nichtleitenden Behälter (25) um die Elektrodenanschlüsse umlaufend gebildet ist.

8. Elektrokardiografiepflaster (15) nach Anspruch 3, ferner umfassend:

eine konturierte Oberfläche, die auf dem nicht reizenden Klebeverband (43) an mindestens einem

Ende des länglichen Streifens (21) enthalten ist.

9. Elektrokardiografiepflaster (15) nach Anspruch 1, ferner umfassend:
eine Vielzahl von Befestigungselementen, die auf dem nichtleitenden Behälter (25) enthalten sind und so ausgerichtet sind, den Elektrokardiografiemonitor (14) sicher im nichtleitenden Behälter (25) aufzunehmen und dort festzuhalten.

10. Elektrokardiografiepflaster nach Anspruch 1, ferner umfassend mindestens eines von:

abgerundete Kanten, die um eine Unterseite des nichtleitenden Behälters (25) herum definiert sind;
eine Gelmulde (46), die um mindestens eine der Elektrokardiografieelektroden (38, 39) auf der Kontaktfläche des Endes (30, 31) des flexiblen Trägers (20) herum umlaufend definiert ist, an dem die mindestens eine Elektrokardiografieelektrode (38, 39) befestigt ist; und
Platzierungs- und Entfernungsglaschen (24), die als Teil des flexiblen Trägers (20) enthalten sind und in Längsrichtung über das mindestens eine Ende (30, 31) des flexiblen Trägers (20) hinausreichen.

11. Elektrokardiografiepflaster (15) nach Anspruch 1, wobei das Pflaster in der Draufsicht eine Sanduhrform bildet;
ein oberer Teil der Sanduhrform bemessen ist, so dass der nichtleitenden Behälter (25) auf einer nach außen zeigenden Oberfläche des flexiblen Trägers (20) sitzt; und
ein unterer Teil der Sanduhrform eine Größe umfasst, die kleiner als der obere Teil der Sanduhrform ist.

12. Elektrokardiografiepflaster (15) nach Anspruch 1, ferner umfassend:
dass der nichtleitende Behälter (25) Elektrodenanschlüsse umfasst, die ausgerichtet sind, um die Elektrokardiografieelektroden (38, 39) über das Paar der Schaltkreisspuren (33, 37) elektrisch an den Elektrokardiografiemonitor (14) zu koppeln.

13. Elektrokardiografiepflaster (15) nach Anspruch 1, ferner umfassend:
ein Paar von konkaven Aussparungen (22), die entlang der Längskanten des verlängerten Streifens (21) definiert sind, wobei jede konkave Aussparung (22) konturiert ist, sodass er im Wesentlichen einen intermammären Spalt freigibt.

14. Elektrokardiografiepflaster (15) nach Anspruch 1, wobei das dehnbare Material (44) mindestens eines von tragbarem Verbandmull, tragbarem Latex, Wi-

ckelgewirke und Trikotleinen umfasst.

15. System, das ein Elektrokardiografiepflaster (15) nach Anspruch 1 und einen Elektrokardiografiemonitor umfasst, wobei eine Patientenrückmeldetaste auf dem Elektrokardiografiemonitor (14) vorgesehen ist.

10 Revendications

1. Timbre d'électrocardiographie à utilisation prolongée (15), comprenant :

un support souple (20) formé d'une bande allongée (21) de matériau extensible (44) avec une section intermédiaire longitudinale étroite (23) s'effilant uniformément vers l'intérieur à partir des deux extrémités (30, 31), la bande allongée (21) étant apte à adhérer uniquement sur une surface de contact définie sur chacune des extrémités (30, 31), et apte à servir de réducteur de frisure pour faciliter une compression de la section intermédiaire longitudinale étroite (23) en réponse à des forces de compression et de torsion ;

un circuit souple (32) fixé sur chaque extrémité (30, 31) à la bande allongée (21) et comprenant une paire de pistes de circuit (33, 37) démarquant chacune respectivement dans l'une des extrémités de la bande allongée (21) ;

un réceptacle non conducteur (25) comprenant une pluralité de plots électriques et amené à adhérer solidement sur une extrémité (31) de la bande allongée (21) opposée à la surface de contact, et actionnable pour recevoir de manière amovible un moniteur d'électrocardiographie (14), le réceptacle non conducteur (25) étant aligné pour servir d'interface électrique entre la paire de pistes de circuit (33, 37) et le moniteur d'électrocardiographie (14) ;

une paire d'électrodes d'électrocardiographie (38, 39) respectivement fixées à et exposées de manière conductrice sur la surface de contact de chaque extrémité (30, 31) de la bande allongée (21), et couplées électriquement à une paire de plots électriques (34) sur le réceptacle non conducteur et la paire de pistes de circuit ;

un compartiment de batterie formé sur une surface inférieure du réceptacle non conducteur ; et une batterie située à l'intérieur du compartiment de batterie du réceptacle non conducteur et en interface électrique avec une paire différente de plots électriques (34) du réceptacle non conducteur.

2. Timbre d'électrocardiographie (15) selon la revendication 1, comprenant en outre :

- l'une des électrodes d'électrocardiographie (38) disposée de façon à adhérer à une région sous le sternum inférieur de la poitrine d'un patient ; et une autre des électrodes d'électrocardiographie (39) disposée de façon à adhérer à la région près du manubrium sur la poitrine du patient orientée de manière centrale (au milieu (16)) le long du sternum (13) vers le haut à partir de l'électrode d'électrocardiographie (38).
3. Timbre d'électrocardiographie (15) selon la revendication 1, comprenant en outre :
un pansement adhésif non irritant (43) appliqué en revêtement au moins partiellement sur chaque extrémité de la bande allongée (21) sur la surface de contact.
4. Timbre d'électrocardiographie (15) selon la revendication 1, comprenant en outre au moins un parmi :
un réducteur de tension extensible latéralement (40) défini dans le circuit souple (32) et actionnable pour faciliter l'extension et la rotation du circuit souple (32) en réponse à des forces de traction et de torsion ; et
une paire de découpes de réduction de tension (41) incluses dans le réducteur de tension (40), qui font chacune face vers l'autre découpe de réduction de tension (41), chaque découpe de réduction de tension (41) s'étendant partiellement transversalement à partir de et se prolongeant longitudinalement vers un côté du circuit souple (32) différent de l'autre découpe de réduction de tension (41), l'une de la paire de pistes de circuit (33, 37) suivant un trajet formé sur le circuit souple (32) entre la paire de découpes de réduction de tension (41).
5. Timbre d'électrocardiographie (15) selon la revendication 4, dans lequel la paire de découpes de réduction de tension (41) définissent un motif en forme de « S » dans le circuit souple (32) lorsque vues de dessus.
6. Timbre d'électrocardiographie (15) selon la revendication 1, dans lequel la section intermédiaire longitudinale étroite (23) définit un motif en forme de sablier dans le support souple (20) lorsque vue de dessus, conforme pour s'adapter au sillon intermammaire de la poitrine d'une patiente de manière confortable.
7. Timbre d'électrocardiographie (15) selon la revendication 1, comprenant en outre :
un joint résistant à l'humidité (35) formé sur le réceptacle non conducteur (25) de manière circconférentielle autour des bornes d'électrode.
8. Timbre d'électrocardiographie (15) selon la revendication 3, comprenant en outre :
une surface profilée incluse sur le pansement adhésif non-irritant (43) sur au moins une extrémité de la bande allongée (21).
9. Timbre d'électrocardiographie (15) selon la revendication 1, comprenant en outre :
une pluralité d'attaches incluses sur le réceptacle non conducteur (25) et orientées pour recevoir solidement et maintenir captif le moniteur d'électrocardiographie (14) dans le réceptacle non conducteur (25).
10. Timbre d'électrocardiographie (15) selon la revendication 1, comprenant en outre au moins un parmi :
des bords arrondis définis autour d'une surface inférieure du réceptacle non conducteur (25) ;
un puits de gel (46) défini de manière circconférentielle autour d'au moins une des électrodes d'électrocardiographie (38, 39) sur la surface de contact de l'extrémité (30, 31) du support souple (20) sur laquelle l'au moins une électrode d'électrocardiographie (38, 39) est fixée ; et
des languettes de mise en place et de retrait (24) faisant partie du support souple (20) et s'étendant longitudinalement au-delà d'au moins une extrémité (30, 31) du support souple (20).
11. Timbre d'électrocardiographie (15) selon la revendication 1, dans lequel le timbre est un motif en forme de sablier lorsque vu de dessus ;
une partie supérieure du motif en forme de sablier est dimensionnée pour placer le réceptacle non conducteur (25) sur une surface orientée vers l'extérieur du support souple (20) ; et
une partie inférieure du motif en forme de sablier a une taille plus petite que la partie supérieure du motif en forme de sablier.
12. Timbre d'électrocardiographie (15) selon la revendication 1, comprenant en outre :
le réceptacle non conducteur (25) comprenant des bornes d'électrode alignées pour servir d'interface électrique entre les électrodes d'électrocardiographie (38, 39), par l'intermédiaire de la paire de pistes de circuit (33, 37), et le moniteur d'électrocardiographie (14).
13. Timbre d'électrocardiographie (15) selon la revendication 1, comprenant en outre :
une paire de découpes concaves (22) définies le long des bords longitudinaux de la bande allongée (21), chaque découpe concave (22) étant profilée conforme pour sensiblement dégager un sillon intermammaire.

14. Timbre d'électrocardiographie (15) selon la revendication 1, dans lequel le matériau extensible (44) comprend au moins un parmi une compresse portable, un latex portable, un tricot à chaîne et un linge Tricot. 5
15. Système comprenant un timbre d'électrocardiographie (15) selon la revendication 1 et un moniteur d'électrocardiographie, un bouton de rétroaction de patient étant prévu sur le moniteur d'électrocardiographie (14). 10

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Fig. 1.

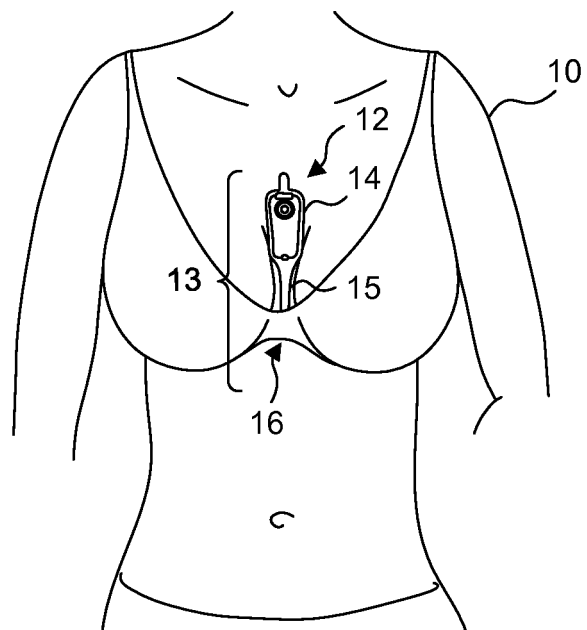


Fig. 2.

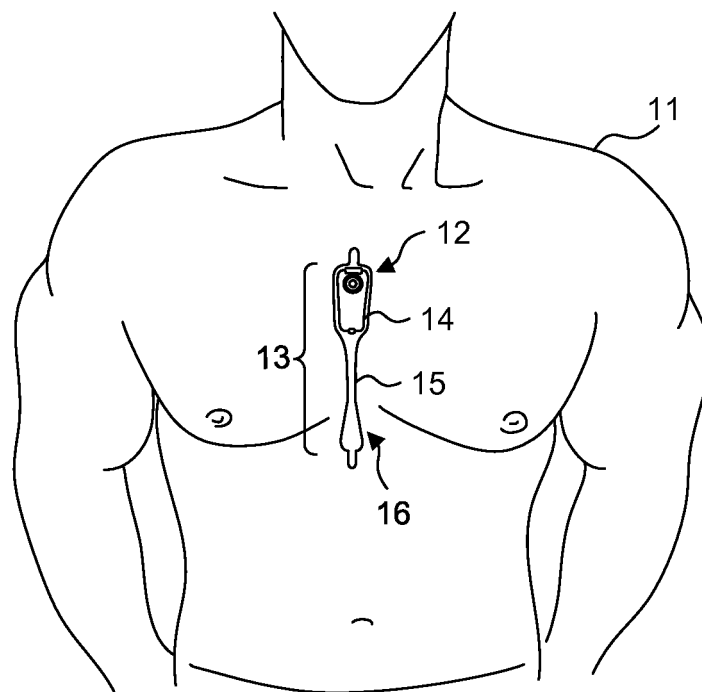


Fig. 3.

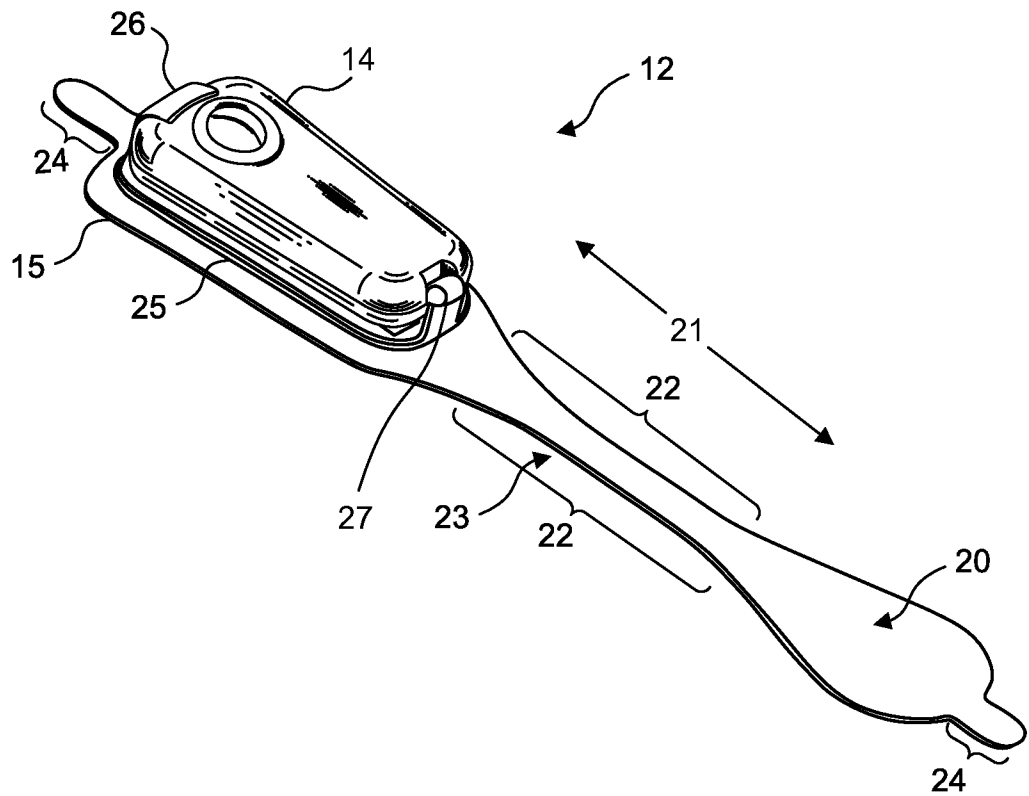


Fig. 4.

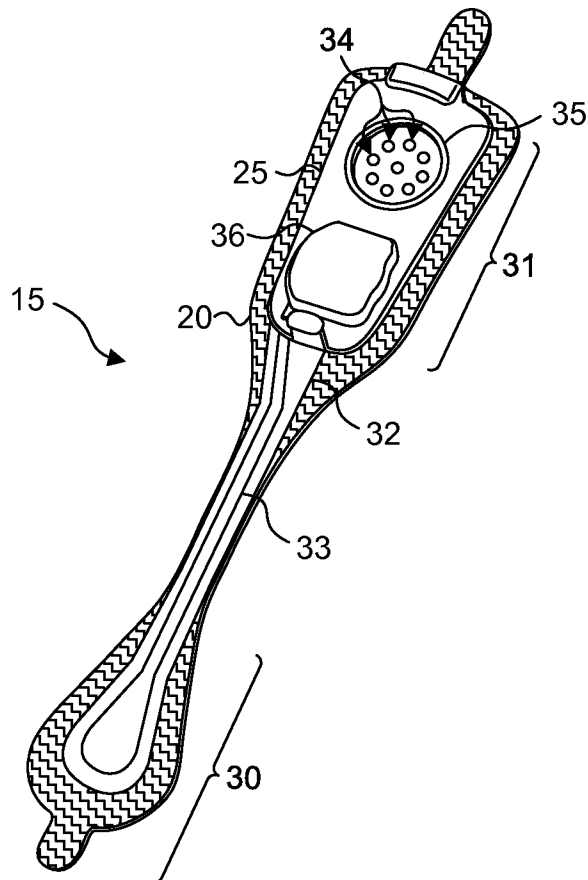


Fig. 5.

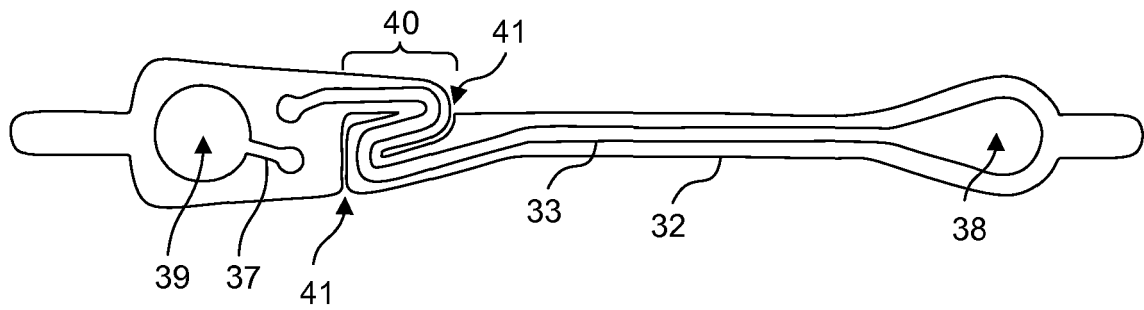


Fig. 6.

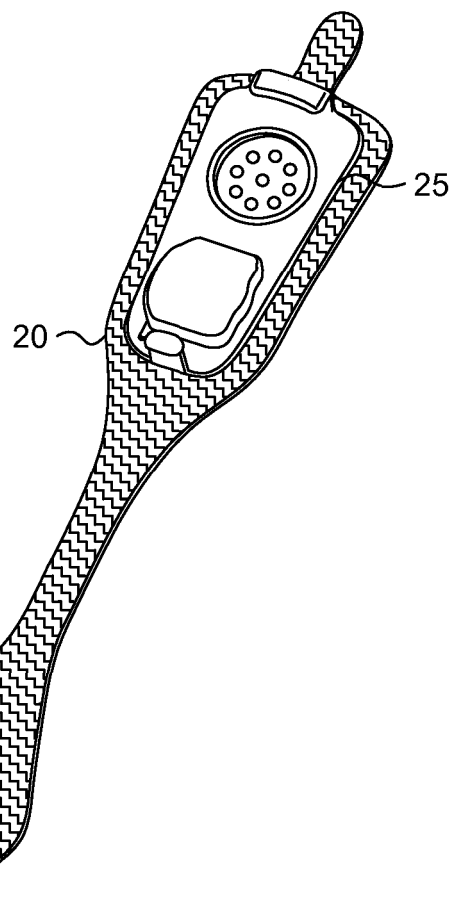


Fig. 7.

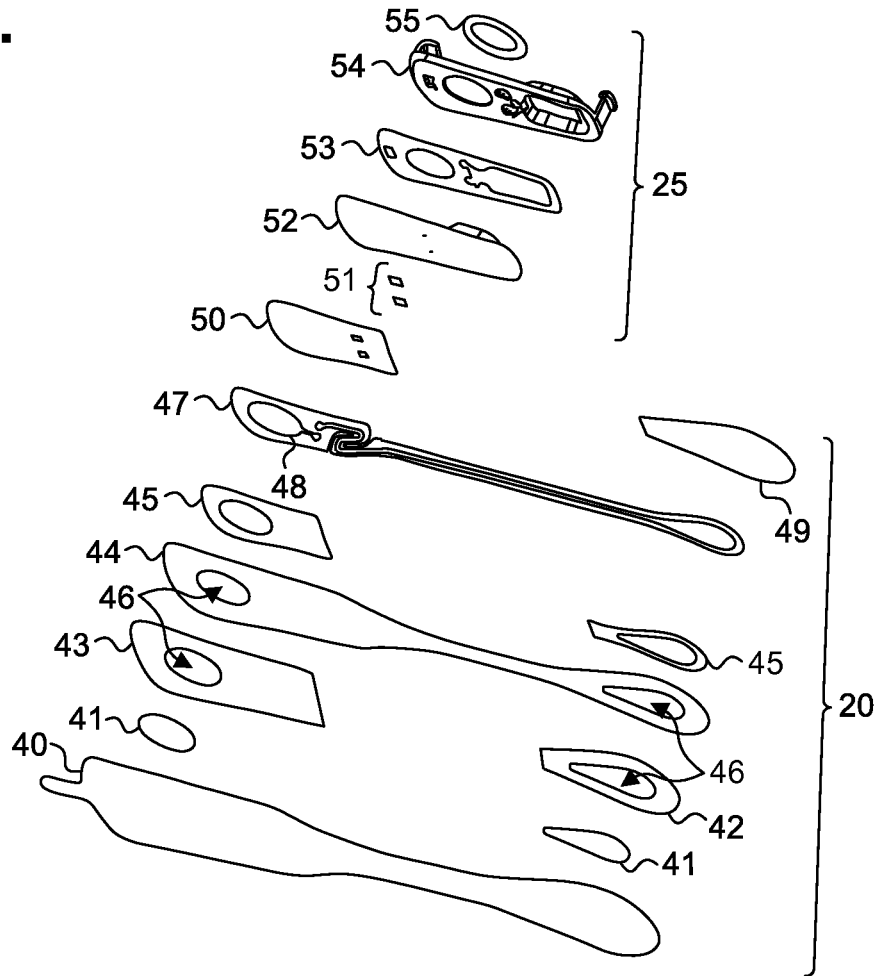


Fig. 8.

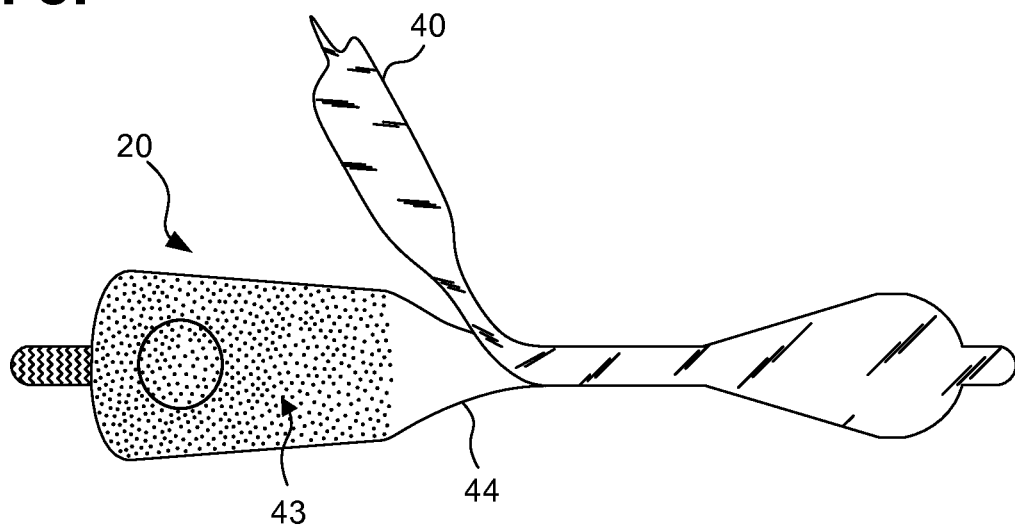


Fig. 9.
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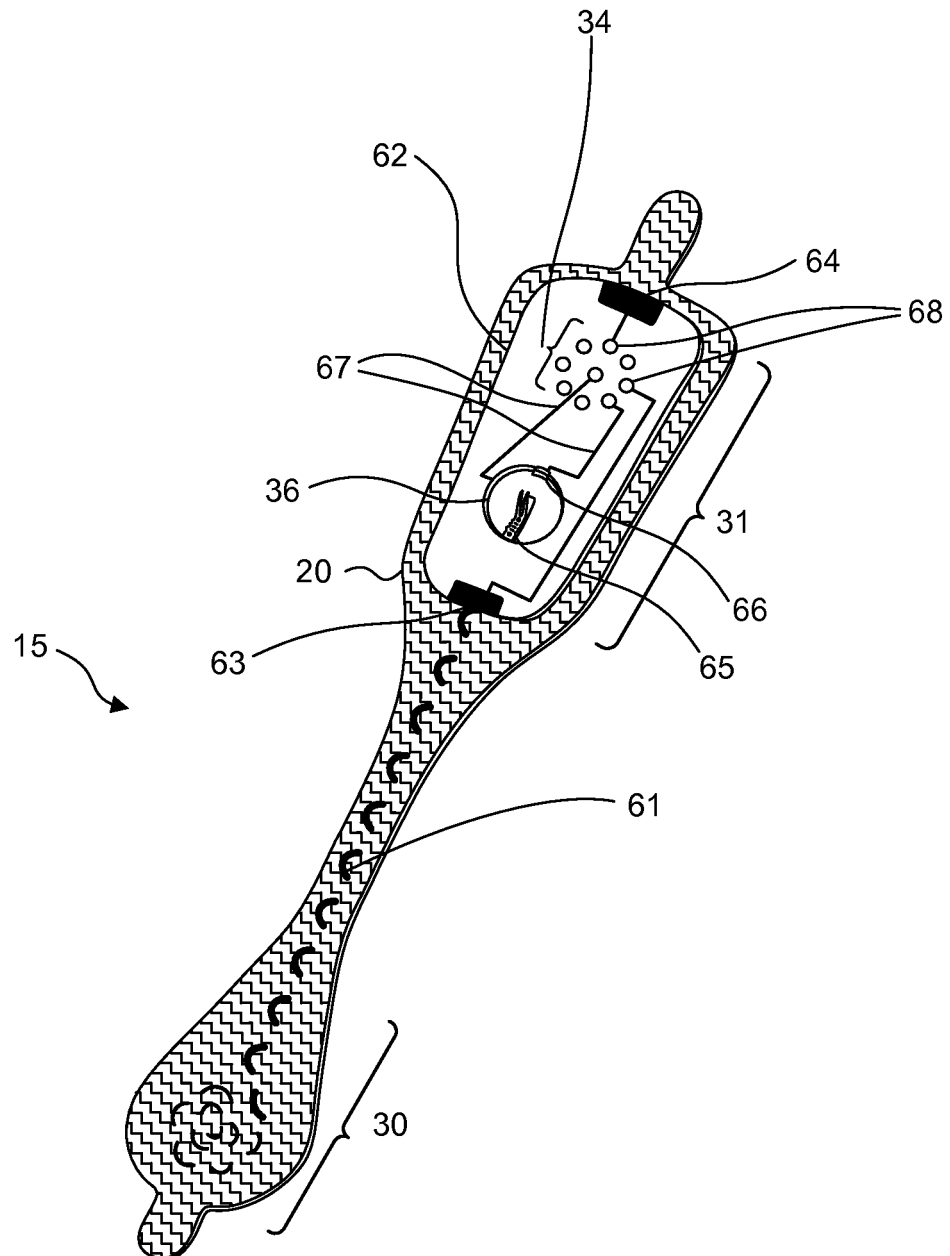


Fig. 10.

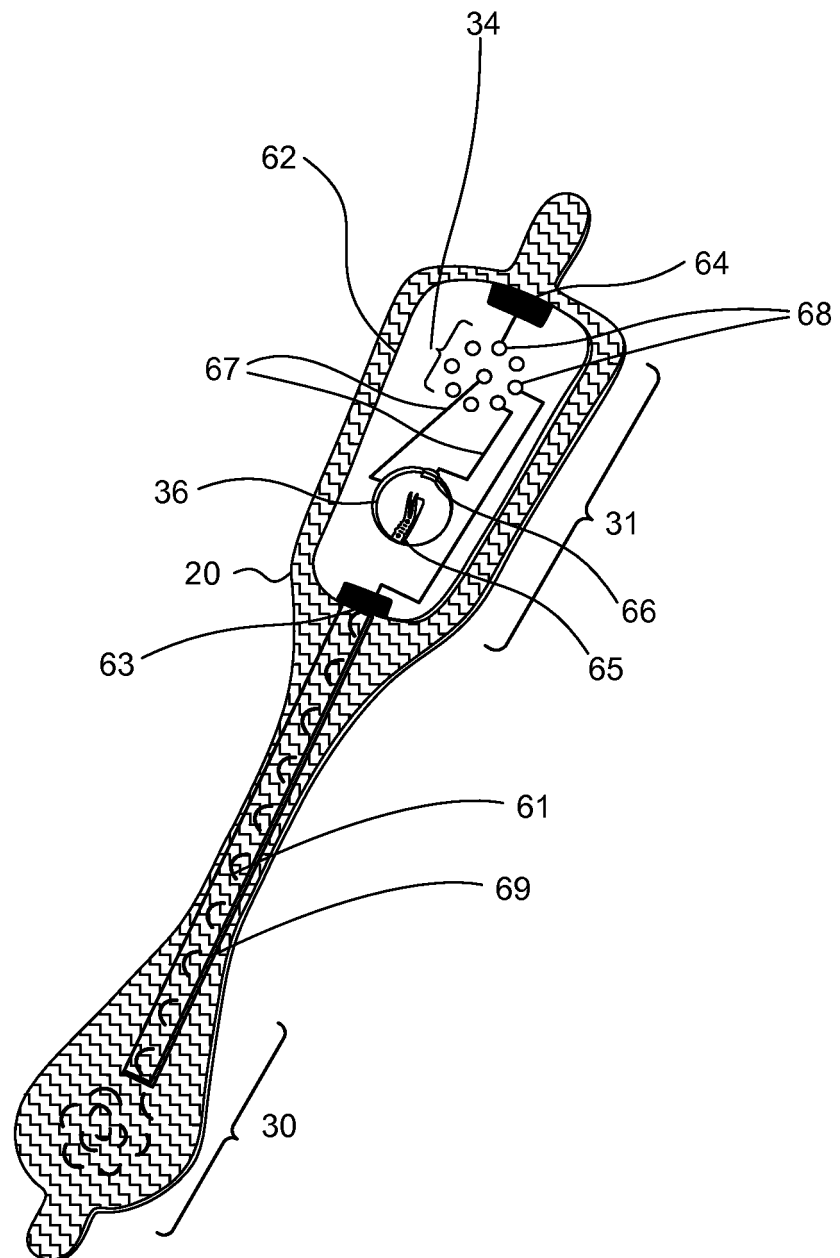


Fig. 11.

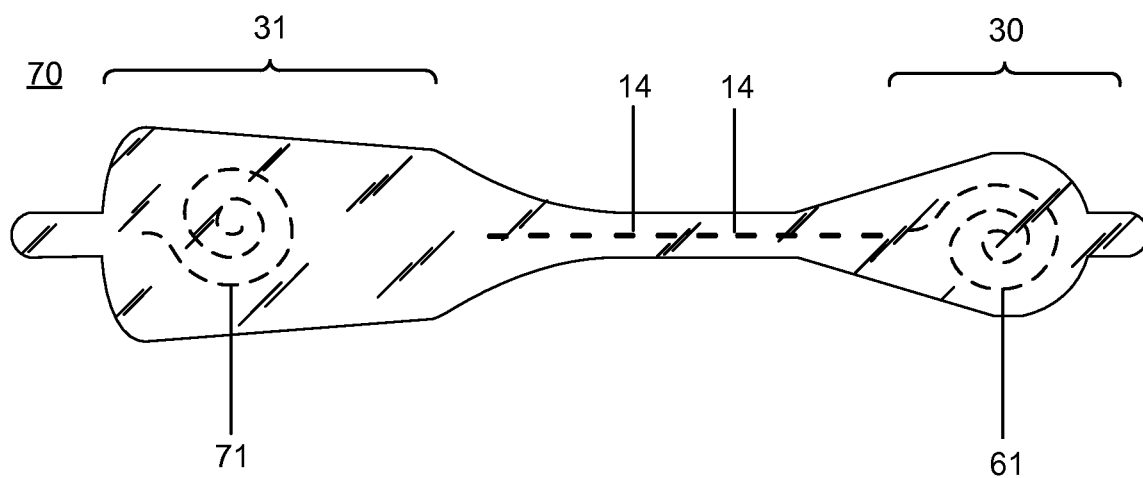


Fig. 12.

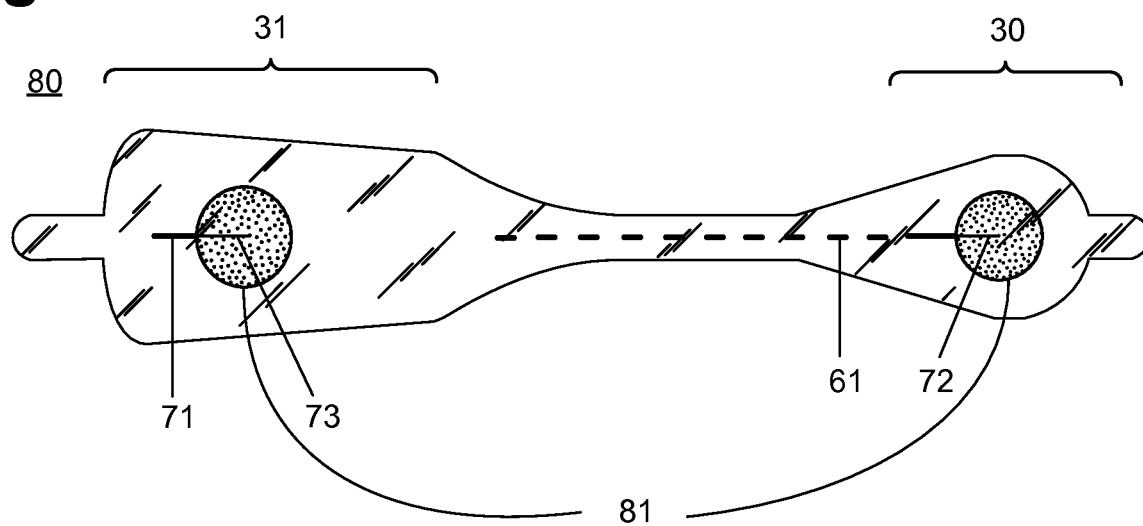


Fig. 13

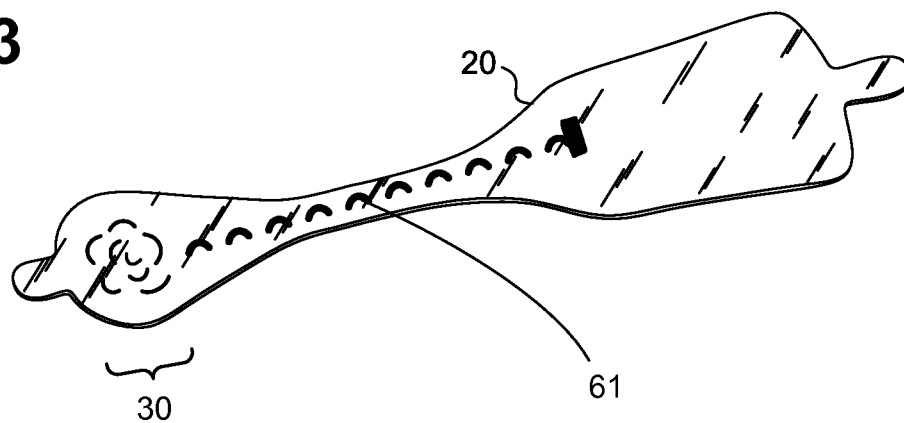


Fig. 14

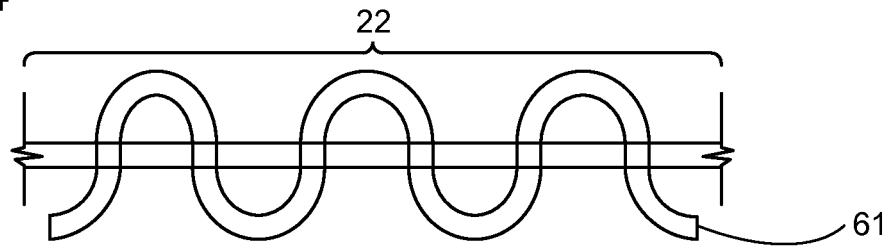


Fig. 15A.

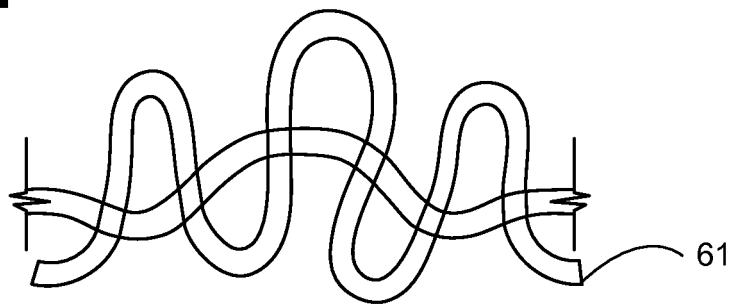


Fig. 15B.

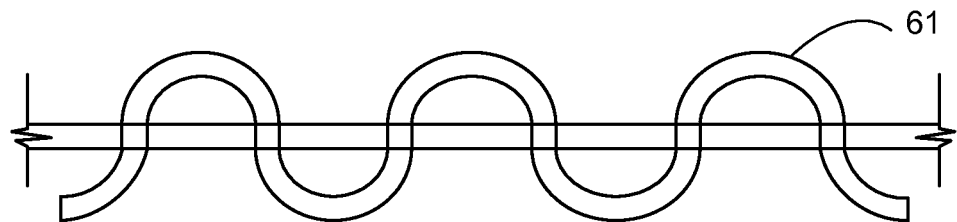


Fig. 15C.

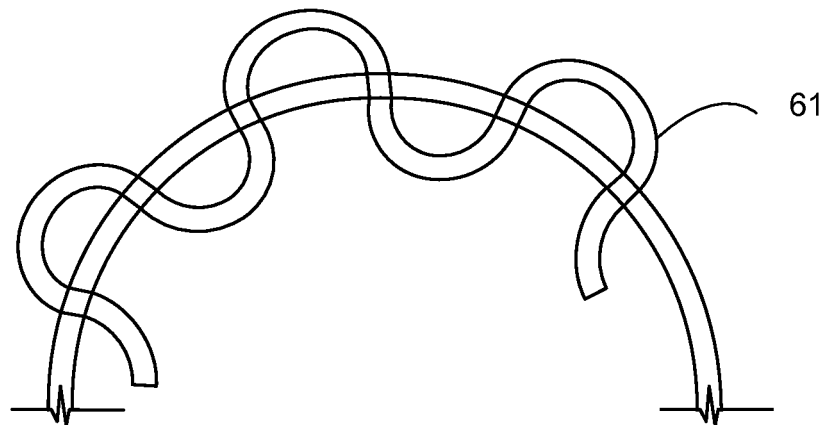
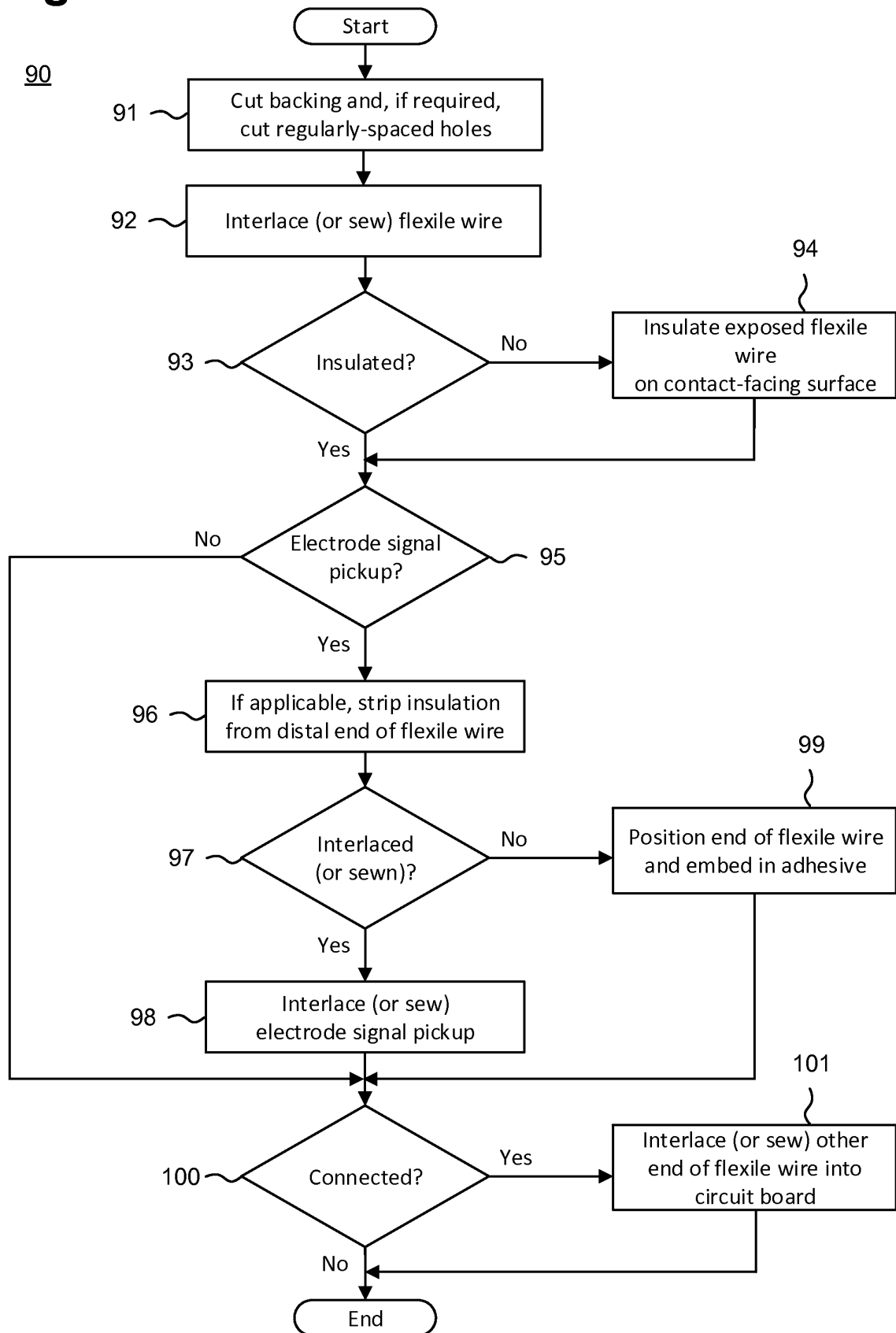


Fig. 16.

REFERENCES CITED IN THE DESCRIPTION

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Patent documents cited in the description

- US 5511553 A [0009]
- US 2004073127 A [0009]
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专利名称(译)	使用交错线电极的延长磨损心电图贴片		
公开(公告)号	EP3048963B1	公开(公告)日	2019-02-20
申请号	EP2014781025	申请日	2014-09-25
[标]申请(专利权)人(译)	BARDY诊断		
申请(专利权)人(译)	BARDY诊断, INC.		
当前申请(专利权)人(译)	BARDY诊断, INC.		
[标]发明人	FELIX JASON BISHAY JON MIKALSON BARDY GUST H		
发明人	FELIX, JASON BISHAY, JON MIKALSON BARDY, GUST H.		
IPC分类号	H01M2/10 H01M4/74 A61B5/0402 A61B5/0408 A61B5/00 A61B5/0428		
CPC分类号	A61B5/0402 A61B5/0408 A61B5/04085 A61B5/04087 A61B5/04286 A61B5/055 A61B5/6833 A61B2560/0214 A61B2562/12 A61B2562/164 A61B2562/166 A61B2562/187 A61B2562/247 H01M2/ /1044 H01M4/747		
代理机构(译)	HANNA MOORE + CURLEY		
优先权	14/080717 2013-11-14 US 61/882403 2013-09-25 US 14/463585 2014-08-19 US 14/463571 2014-08-19 US		
其他公开文献	EP3048963A2		
外部链接	Espacenet		

摘要(译)

可以通过可穿戴监视器 (12) 提供生理监测, 该可穿戴监视器包括柔性的延伸耐磨电极贴片 (15) 和可拆卸的可重复使用的监视器记录器 (14)。一对柔性线 (61,71) 交织或缝合到柔性背衬 (20) 中, 用作电极信号拾取器和电极电路迹线。监视器 (12) 沿胸骨 (13) 位于患者胸部的中央, 这显着提高了感知皮肤电信号的能力, 尤其是心房产生的心电信号。为了抵消由于压缩力和扭转力引起的移位, 在电极贴片 (15) 的下侧或接触表面上提供无刺激性粘合剂 (43), 但仅在远端 (30) 和近端 (31) 上提供。将柔性线 (61,71) 交织到柔性背衬 (20) 中还提供抵抗压缩力, 拉伸力和扭转力的结构支撑和延展性。

