An infrasonic stethoscope for monitoring physiological processes of a patient includes a microphone capable of detecting acoustic signals in the audible frequency bandwidth and in the infrasonic bandwidth (0.03 to 1000 Hertz), a body coupler attached to the body at a first opening in the microphone, a flexible tube attached to the body at a second opening in the microphone, and an earpiece attached to the flexible tube. The body coupler is capable of engagement with a patient to transmit sounds from the person, to the microphone and then to the earpiece.

20 Claims, 17 Drawing Sheets
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FIG. 3

FIG. 4
FIG. 6

Infrascope signal 20

A/D

Wireless or by cable

Laptop or PC

Time History

Short Term Fourier Transform (STFT)

Spectrogram

Internet

Laptop, PC, Tablet, or Smart Phone

FIG. 7
Location A

Time analysis showing ECG and infrascope results

FIG. 8
Location A

Average of 3 minutes of signal collected from ECG and the infrascope at 500 Hz (Gain 1x). All units are in volts.

FIG. 9
Location P

Time analysis showing ECG and the infrascope results

FIG. 10
Average of 3 minutes of signal collected from ECG and the infrascope at 500 Hz (Gain 1x). All units are in volts.

**FIG. 11**
Location T

Time analysis showing ECG and the infrascope results

FIG. 12
Location T

Average of 3 minutes of signal collected from ECG and the infrascope at 500 Hz (Gain 1x). All units are in Volts.

FIG. 13
Location M

Time analysis showing ECG and the Infrascope results

FIG. 14
Location M

Average of 3 minutes of signal collected from ECG and the infrascope at 500 Hz (Gain 1x). All units are in Volts.

FIG. 15
Location T
Time analysis showing ECG and the Infrascope results

**FIG. 16**
Average of 3 mins of signal collected from ECG and the Infrascope at 500Hz (Gain 1x). All units are in Volts.
Location T, Subject 1

Average of 3 minutes of signal collected from ECG and the infrascope at 1000 Hz [Gain 1x]. All units are in Volts.

FIG. 18
Location T, Subject 2

Average of 3 minutes of signal collected from ECG and the Infrascope at 1000 Hz (Gain 1x). All units are in Volts.

FIG. 19
1

INFRASONIC STETHOSCOPE FOR MONITORING PHYSIOLOGICAL PROCESSES

CROSS-REFERENCE TO RELATED PATENT APPLICATION(S)

This application claims the benefit of and priority to U.S. Provisional Application Ser. No. 62/058,794, filed on Oct. 2, 2014, the contents of which are incorporated by reference herein in their entirety.

STATEMENT REGARDING FEDERALLY SPONSORED RESEARCH OR DEVELOPMENT

The invention described herein was made in the performance of work under a NASA contract and by employees of the United States Government and is subject to the provisions of Public Law 96-517 (35 U.S.C. §202) and may be manufactured and used by or for the Government for governmental purposes without the payment of any royalties thereon or therefore. In accordance with 35 U.S.C. §202, the contractor elected not to retain title.

FIELD OF THE INVENTION

The present disclosure relates to an infrasonic stethoscope (or “infrascope”) for monitoring physiological processes and is particularly related to a wireless infrasonic stethoscope.

BACKGROUND OF THE INVENTION

Sound at frequencies below 20 Hertz is termed “infrasound.” A particularly favorable property of infrasound is its propagation over long distances with little attenuation. Infrasound has this property because atmospheric absorption is practically negligible at infrasonic frequencies, and because there is an acoustic “ceiling” in the stratosphere where a positive gradient of the speed of sound versus altitude causes reflections of infrasonic rays back to Earth. Infrasound propagation over long distances (e.g., thousands of kilometers) is predominantly due to refractive ducting from the upper layers in the atmosphere, while propagation over short distances is completed by direct path.

The density, acoustic impedance, and speed of sound through different human and animal tissues varies depending upon location of the auscultation. When an acoustic signal travels through tissue layers, the amplitude of the original signal becomes more attenuated with depth of the acoustic signal source. Attenuation (i.e., energy loss) could be due to absorption, reflection, and scattering at interfaces of different tissues. The degree of attenuation also depend upon frequency of the sound wave and the distance it travels. Generally speaking, a high frequency acoustic signal is associated with high attenuation thus limiting tissue penetration, but lower frequencies do not have attenuation issue thus providing physicians better understanding of the heart performance. More than 60% power spectral density of heart signals fall in infrasonic bandwidth. Low frequency acoustic signals detected from different human organs, such as the heart, are valuable to physicians for monitoring heart and lungs.

Microphones and stethoscopes are regularly used by physicians in detecting sounds for monitoring physiological conditions. Phonocardiography has been in use for more than 75 years to monitor heart beats as well as to detect the audible sound of the blood flowing through the heart. These physiological condition monitors are coupled directly to a person’s body and processes are measured either by listening or by recording the signals in certain bandwidth. The physiological processes such as respiration and cardiac activity are reflected in a different frequency bandwidth from 1/20 Hertz to 500 Hertz. Other stethoscopes are capable of monitoring only audible frequency bandwidth, and are not capable of monitoring infrasonic frequencies below 20 Hz. Low frequency acoustic signals below 20 Hertz are not audible, but can provide useful information to physicians.

Inside of a normal heart, there are four chambers namely; the right atrium, the left atrium, the right ventricle, and the left ventricle. The function of a heart is to keep blood flowing in one-way direction. When a valve opens, the valve lets the right amount of blood through, and then closes to keep blood from flowing backwards in between beats. An easy and relatively inexpensive assessment of any patient’s cardiac status can be determined by sounds in the chest. The key to good auscultation lies in low and high pitched sounds. As the heart beats, blood flows from right atrium into the right ventricle through the tricuspid valve.

Blood then flows to the lungs through pulmonary valve (sometimes also called semilunar valve) to pick up right amount of oxygen. The blood flows from the lungs back into the left atrium and enters into the left ventricle through the mitral valve. Blood then is pumped to the aorta through the aortic valve and goes out to rest of the body providing oxygen and nutrients to the body cells. All four chambers (right atrium, right ventricle, left atrium, and left ventricle) must contract at just the right time for normal heart to functioning properly. The proper timing is coordinated by heart’s electrical pathways. The electrical signals are produced by the sinoatrial node (SA node) and atrioventricular node (AV node).

The SA node is a group of cells located in the right atrium that initiates contraction of both atria to push blood into their corresponding ventricles. Due to insulation between the atria and ventricles, the SA node signals do not continue directly to the ventricles but pass through the AV node, which is another group of cells located in the floor of right atrium between the atria and ventricles. The AV node regulates the signal to ensure that the atria are empty and closed before the ventricles contract to push the blood out of the heart. The SA node sends signals to stimulate the heart to beat between 60-100 times per minute.

The cardiovascular system is complex and numerous problems could take place inside anywhere from the electrical system of the heart to the large or small blood vessels. There are over 60 different types of cardiovascular disease, all of which somehow affect the cardio or vascular systems. The heart sounds generated by the beating of heart and the resultant flow of blood can provide important information about the condition of the heart. In healthy adults, two normal heart sounds occur in sequence with the heartbeat. A first sound is produced based on the closure of the atrioventricular valves (i.e. mitral and tricuspid valves) located between the atria and ventricles, and is referred to as S1. A second sound is produced as a result of closure of the semilunar valves (i.e. pulmonary and aortic valves), which control the flow of blood as it leaves the heart via the arteries, and is referred to as S2.

The first heart sound S1 consists of four sequential components: 1. Small low frequency vibrations that coincide with the beginning of left ventricular contraction. 2. High frequency vibration, easily audible related to mitral valve closure (M1). 3. A second high frequency component related
frequency ultrasound signals become attenuated due to filtering may be useful to record low and high frequency infrasonic microphone to acquire useful information that indicates valve malfunctions, such as stenosis or regurgitation. Other high frequency murmurs can occur between the two major heart sounds during systole or diastole. The cardiac sounds S1, S2, S3, and S4 can be attributed to specific cardiac activity. S1 is attributed to the onset of the ventricular contraction (10-140 Hertz bands). S2 is attributed to closure of the semilunar valves (10-400 Hertz bands). S3 may be attributed to ventricular gallop, which may be heard during rapid filling (i.e. diastole) of the ventricles. S4 may be attributed to atrial gallop, which may be heard in late diastole, during atrial contraction. S3 and S4 are of very low intensity and can be heard externally when amplified.

Other sounds may be heard from opening snaps of the mitral valve or ejection sound of the blood in the aorta which indicates valve malfunctions, such as stenosis or regurgitation. Other high frequency murmurs can occur between the two major heart sounds during systole or diastole. The murmurs can be innocent but can also indicate certain cardiovascular defects.

Continuous fetal heart monitoring is an important step to evaluate the well-being of a fetus. The fetal heart rate may indicate if the fetus is getting enough oxygen. Most of the time ultrasound transducers are used for monitoring fetal heart rate as conventional stethoscopes undesirably pick up signals from maternal abdominal vessels. Due to abdominal fat of the mother or fetal positioning, it may be difficult to monitor fetal heart passively, so most of the time ultrasound transducers are used where ultrasound pulses are radiated towards the fetus and reflective pulses are used for monitoring. If enough reflective signals are not received, the penetration depth of ultrasound pulses are increased which may decrease quality and signal-to-noise ratio. The low frequency ultrasound signals become attenuated due to absorption, reflection, and scattering due to abdominal fat. The insonified signals have relatively very low attenuation coefficient hence the signals are expected to be of high quality with better signal to noise ratio and helpful to gynecologists.

Many heart sounds are in a low-frequency band spectrum with low intensity level and may require extremely sensitive infrasonic microphone to acquire useful information that cannot be perceived by the physician’s ear. The passive filtering may be useful to record low and high frequency bands separately. The sounds are of short duration and highly non-stationary but enable to measuring systolic and diastolic time intervals, which may have diagnostic importance.

Accordingly, there is a need for a monitoring device that overcomes the disadvantages presented by the prior art.

BRIEF SUMMARY OF THE INVENTION

An infrasonic stethoscope (or infrascope) or monitoring physiological processes of a patient includes a microphone capable of detecting acoustic signals in the audible frequency bandwidth and in the infrasonic bandwidth. The microphone has a body, which includes first and second spaced apart openings. A body coupler is attached to the first opening of the body to form a substantially air-tight seal, wherein the body coupler is capable of engagement with the patient to monitor physiological processes. A flexible tube is attached to the body at the second opening in the microphone. An earpiece is attached to the flexible tube. The body coupler is capable of engagement with a patient to transmit sounds from the patient to the microphone, and then to the earpiece.

These and other features, advantages, and objects of the present invention will be further understood and appreciated by those skilled in the art by reference to the following specification, claims, and appended drawings.

BRIEF DESCRIPTION OF THE SEVERAL VIEWS OF THE DRAWINGS

The organization and manner of the structure and operation of the disclosed embodiments, together with further objects and advantages thereof, may best be understood by reference to the following description, taken in connection with the accompanying drawings, which are not necessarily drawn to scale, wherein like reference numerals identify like elements in which:

FIG. 1 is perspective view of an embodiment of an infrascope which can be used for external monitoring of a patient;

FIG. 2 is a perspective view of the infrascope attached to a catheter, for use in internal fetal monitoring of a patient;

FIG. 3 is a perspective view of a pair of infrascopes which can be used in Doppler phonocardiography;

FIG. 4 is a graph showing bandwidths of heart sound;

FIG. 5 is a cross-sectional view of a microphone which forms part of an infrascope of the present invention and of a body coupler according to a first embodiment attached to the microphone and which forms part of the infrascope of the present disclosure;

FIG. 5A is a cross-sectional view of a body coupler according to a second embodiment and which forms part of the infrascope of the present disclosure;

FIG. 6 is a schematic view of a skeleton of a patient;

FIG. 7 is a flow chart regarding the process of how signals from the infrascope are transmitted and analyzed;

FIGS. 8-17 are charts of infrascope signals collected at locations A, P, T and M of FIG. 6 with reference to electrocardiogram signals referred as ECG or EKG; and

FIGS. 18 and 19 show charts of infrascope data as compared to ECG or EKG on two different subjects from 1 Hz through 1000 Hz.

DETAILED DESCRIPTION OF THE INVENTION

While the disclosure may be susceptible to embodiment in different forms, there is shown in the drawings, and herein will be described in detail, a specific embodiment with the understanding that the present disclosure is to be considered an exemplification of the principles of the disclosure, and is not intended to limit the disclosure to that as illustrated and described herein. Therefore, unless otherwise noted, features disclosed herein may be combined together to form additional combinations that were not otherwise shown for purposes of brevity. It will be further appreciated that in some embodiments, one or more elements illustrated by way
of example in a drawing(s) may be eliminated and/or substituted with alternative elements within the scope of the disclosure.

As shown in FIG. 1, an infrascope 20 is provided to monitor physiological processes of a patient. The infrascope 20 detects signals in a bandwidth from 0.03 Hertz through 1000 Hertz, or alternatively 0.03 through 500 Hertz. These bandwidths contain signals which are audible and inaudible to the human ear. This infrascope 20 has multiple applications to measure a variety of human physiological processes, including, but not limited to, cardiac monitoring, external fetal monitoring, internal fetal monitoring, stress phonocardiography testing, Doppler phonocardiography, biometric identification and polygraphs. The bandwidth of audible and inaudible sounds produced by cardiac activity are shown in FIG. 4, which demonstrates energy distribution (dynes/cm²) as a function of frequency (Hz).

The infrascope 20 contains a microphone 22, a body coupler 24 or 24a attached to the microphone 22, a flexible tube 26 attached to the microphone 22 and earpiece 28 connected to the flexible tube 26. For internal fetal monitoring, as shown in FIG. 2 and as further described herein, the body coupler 24a is used and the microphone 22 is further connected to a catheter 23 via the body coupler 24a.

The microphone 22 is substantially the same as the microphone described in U.S. Pat. No. 8,401,217, with the modifications described herein. The contents of U.S. Pat. No. 8,401,217 is incorporated by reference in its entirety. The microphone 22 is best shown in FIG. 5 and includes a cup-like body 30, a cup-like support plate 32, an insulating member 34, a conductor 36, a backplate 38, a membrane 40 and a low-noise preamplifier board 42.

The body 30 has a cylindrical side wall 44 having a proximal end and a distal end, an end wall 46 at the proximal end of the body 30, and a connection port 48 extending proximally from the end wall 46. The body 30 is formed of metal, such as stainless steel or aluminum. The side wall 44 and the end wall 46 define an internal cavity 50 within the body 30. The distal end of the body 30 is open such that an aperture 52 is defined in the body 30. A thread form 54 is provided on the exterior surface of the side wall 44 at the distal end. The end wall 46 substantially closes the proximal end of the body 30, with the exception of an aperture 56 therethrough, and may extend perpendicularly relative to the side wall 44. The aperture 56 may be centrally located in the end wall 46 and is communication with the connection port 48. The connection port 48 extends proximally from the end wall 46 and has a passageway 58 therethrough which is communication with the cavity 50 via the aperture 56. The exterior surface of the connection port 48 has a thread form 60 thereon. An aperture 62 is provided through the side wall 44 at a position spaced from the proximal end of the side wall 44.

The support plate 32 is attached to the internal surface of the side wall 44 and seats within the cavity 50. The support plate 32 is formed of metal, and has a circular base wall 64 which spans the diameter of the side wall 44 and is parallel to the end wall 46, and a depending side wall 66 which extends distally from the base wall 64. The side wall 66 terminates in a free end. The side wall 66 engages against the internal surface of the side wall 44 of the body 30, such that the free end of the side wall 66 is proximate to the distal end of the body 30, and the base wall 64 is spaced from the distal end of the body 30. The support plate 32 is affixed to the body 30 by suitable means, such as welding, in such a way that whole assembly can be connected to the ground of preamplifier board 42. As a result of this arrangement, a distal chamber 68 is formed between the base wall 64 and the distal end of the body 30, and a proximal chamber 70 is formed between the base wall 64 and the proximal end of the body 30. The base wall 64 has an aperture 72 therethrough, which may be centrally located. The base wall 64 also has at least one aperture 74 or slot therethrough to allow air to flow from the distal chamber 68 to the proximal chamber 70.

The insulating member 34, which may be formed of plastic, ceramic, wood or any suitable insulating material, seats within the aperture 72 in the support plate 32 and is used to electrically isolate the conductor 36, the backplate 38 and the preamplifier board 42 from the support plate 32. As shown, the insulating member 34 has a central portion 76 which extends through the aperture 72, a proximal portion 78 which extends radially outwardly from the central portion 76 on the distal side of the base wall 64, and a distal portion 80 which extends radially outwardly from the central portion 76 on the proximal side of the base wall 64. A passageway 82 extends through the central portion 76.

The backplate 38 is formed of a conducting material, and is formed from a base wall 88 and may further be formed of a proximal extending portion 90 which extends perpendicularly from the base wall 88. The backplate 38 may be formed of, for example, from conducting ceramics, brass, or stainless steel. A passageway 89 extends through the base wall 88, and extending portion 90 if provided, from its proximal surface to its distal surface. A permanently polarized thin polymer film 91 is coated on the distal surface of the backplate 38. The polarized thin polymer film 91 operates without the need for an external power supply. As described in U.S. Pat. No. 8,401,217, the backplate 38 has a plurality of spaced apart holes 92 therethrough (two holes are visible in FIG. 5). The extending portion 90 engages against the distal portion 80 of the insulating member 34, and is secured to a distal end of the conductor 36, such that the backplate 38 and the conductor 36 are in electrical communication. The base wall 88 of the backplate 38 is parallel to the base wall 64 of the support plate 32. A slot 94 is defined between the outer diameter of the backplate 38 and the side wall 44 of the body 30. The area between the backplate 38 and the proximal end of the body 30 defines a back chamber.

The conductor 36 extends through the passageways 82, 89 and extends into the proximal chamber 70. The conductor 36 is electrically connected to backplate 38. As shown, the conductor 36 is formed of a conducting rod or wire 84 which extends through the passageways 82, 89, and a conductive rod 86 extending proximally from the conducting rod or wire 84 and the insulating member 34. If formed of two components, the components are suitably connected to each other to form an electrical connection. The rod or wire 84 and rod 86 may be formed of brass, or may be formed of differing conductive materials. The proximal end of the conductor 46 is proximate, to but spaced from, the end wall 46 such that a gap is defined therebetween.

The membrane 40 is formed of a flexible conductive material and is seated at the distal free end of the side wall 66 of the support plate 32 such that the membrane 40 is positioned within the distal chamber 68 and is proximate to, but spaced from, the distal end of the body 30. The diameter of the membrane 40 is selected so that the membrane 40 stays within side wall 66. The membrane 40 is parallel to the end wall 46 of the body 30 and to the base wall 64 of the support plate 32. As a result, the membrane 40 is in electrical communication with the support plate 32. The tension of the membrane 40 may be less than about 400 Newton per meter.

The backplate 38 is proximate to, but spaced from the membrane 40, such that an air gap 98 is formed between the
membrane 40 and the backplate 38 to create a capacitor in the microphone 22 as described in U.S. Pat. No. 8,401,217. As described in U.S. Pat. No. 8,401,217, the number, locations and sizes of the holes 92, the size of the slot 94, and the inner volume of the backchamber are selected such to allow enough air flow to provide proper damping of the motion of the membrane 40. As described in U.S. Pat. No. 8,401,217, the backchamber serves as a reservoir for the airflow through the holes 92 in the backplate.

In an exemplary embodiment, the membrane 40 has a diameter of approximately 1.05 inches (0.0268 meter). The membrane 40 may have the following characteristics/dimensions:

- radius = 0.0134 meter;
- thickness = 2.54x10^{-5} meter;
- density = 8000 kilogram/meter^3;
- tension = 400 N/meter;
- surface density = 0.1780 kilogram/meter^2; and
- stress = 47.4054 PSI.

The microphone 22 may comprise an air layer which may have the following characteristics/dimensions:

- air gap = 2.54x10^{-5} meter;
- density = 1.2050 kilogram/meter^3;
- viscosity = 1.8x10^{-5} Pascal-second;
- sound velocity through the air gap = 290.2 meters per second; and
- gamma = 1.4.

The microphone 22 may also comprise a slot 94 which may have the following characteristics/dimensions:

- distance from the center of the backplate = 0.0117 meter;
- width = 0.0035 meter;
- depth = 0.00114 meter; and
- area = 0.000258 meter^2.

The backplate 38 may define six holes 92, and each hole 92 may have the following characteristics/dimensions:

- distance from center of backplate to center of hole = 0.00526 meter;
- radius = 0.002 meter;
- depth = 0.045 meter;
- angle between two lines going from center of backplate to either side edge of hole = 43.5 degrees; and
- area = 1.26x10^{-5} meter^2.

The microphone 22 may also have the following further characteristics/dimensions:

- volume of the backchamber = 5x10^{-5} meter^3;
- membrane mass = 480 kilogram/meter^2;
- membrane compliance = 3.2x10^{-11} meter^3/Newton; and
- air gap compliance = 3.5x10^{-10} meter^3/Newton.

In one embodiment, the resonant frequency of the microphone 22 may be 3108.01 Hertz.

The preamplifier board 42 is planar and extends radially outwardly from the proximal end of the conductor 36. The preamplifier board 42 is connected to the proximal end of the conductor 36 by suitable means such that there is an electrical connection between the preamplifier board 42 and the conductor 36, such as a brass screw 99. The preamplifier board 42 is parallel to the end wall 36 of the body 30, and the base wall 64 of the support plate 32 and the base wall 88 of the backplate 38. The position of the preamplifier board 42 defines a first proximal chamber 100 which has a volume V1 between the preamplifier board 42 and the end wall 46 of the body 30, and a second distal chamber 102 which has a volume V2 between the preamplifier board 42 and the base wall 64 of the support plate 32. A slot 104 is defined between the outer diameter of the preamplifier board 42 and the side wall 44 of the body 30 to allow air to flow from the distal chamber 102 to the proximal chamber 100. In an embodiment, volume V1 is approximately 0.1287 cubic inch, and volume V2 is approximately 0.6 cubic inch. The air can only flow from the distal chamber 102 to the proximal chamber 100 through the slot 104. In an embodiment, this slot 104 has a clearance distance between the outer diameter of the preamplifier board 42 and the side wall 44 of approximately 0.025", which slot 104 extends around the preamplifier board 42.

An electrical connection 106 extends through the aperture 62 in the side wall 44 and is sealed to the side wall 44 by suitable means. The electrical connection 106 is electrically connected with the preamplifier board 42 via wires 108, 110. The preamplifier board 42 is also electrically connected to the body 30 via a wire 110, which provides a ground to the preamplifier board 42. The preamplifier board 42 contains known components for measuring the capacitance between the membrane 40 and the backplate 38, and converting this measured capacitance into voltage.

The connection port 48 is connected to a distal end of the flexible tube 26, which may be formed of latex or rubber, which has an earpiece 28 at the proximal end of the tube 26. Such a flexible tube 26 and earpiece 28, like a typical stethoscope, are known in art for transmitting sound. The flexible tube 26 is attached to the connection port 48, such that there is no air exchange between the flexible tube 26 and the body 30, and such that the passageway through the tube 26 is in communication with the distal chamber 100 via the passageway 58 and aperture 56. When the earpiece 28 is inserted into the ears of the medical personnel, this allows substantially no air exchange between the cavity 50 of the microphone 22 and the outside the microphone 22. The length of the flexible tube 26 is adjusted so that maximum audible sound is received at the earpiece 28, which are used by medical personnel to hear the desired sounds in real time.

The combination of volumes V1 and V2 and the slot 104 around the preamplifier board 42 provide sufficient acoustic resistance for pressure equalization, and lowers the low frequency threshold. When the flexible tube 26 is connected to the earpiece 28, due to increased acoustic resistance and longer required period for pressure equalization, this lowers the low ~3 dB frequency to 0.03 Hertz.

As described herein, the microphone may differ from U.S. Pat. No. 8,401,217 in that the body 30 is not completely sealed in that a connection port 48 is provided for connecting the microphone 22 to the flexible tube 26 and the earpiece 28, in that the preamplifier board 42 is mounted horizontally in the body 30 to divide the backchamber into two lower chambers 100 and 102 and that the preamplifier board 42 is parallel to the membrane 40, rather than being positioned vertically that is perpendicular to the membrane 40. As is positioned in U.S. Pat. No. 8,401,217, and in that the grid of U.S. Pat. No. 8,401,217 is eliminated and instead body 30 includes threads 54 for connection of the body coupler 24 or 24a to the distal end of the body 30.

The body coupler 24, 24a threadedly attaches to the thread form 54 at the distal end of the body 30 such that there is no air exchange between the body coupler 24, 24a and the body 30. In one embodiment, as shown in Fig. 5, the body coupler 24 is formed of an outer ring 114 which has a flexible non-conductive diaphragm 116 attached thereto and which spans the diameter of the ring 114. The outer ring 114 may be formed either of thermoplastic polyurethane elastomers (TPU) or of closed cell polyurethane foam material which can be made of different densities, and has an internal thread form 118 for attachment of the outer ring 114 to the distal end of the body 30. The TPU material is used when full spectrum of acoustic signals are to be recorded from a
heart and closed cell polyurethane foam material is used only when infrasonic signals is to be recorded as this material acts as a passive filter and audible sound is shunted off. When attached, the membrane 40 of the microphone 22 and the diaphragm 116 of the body coupler is approximately 0.1 inch apart. The body coupler 24 is placed against the backplate 38, and the radius of the imaginary circle is about 0.0117 meter. The width of the slot 94 is about 0.00179 m². The volume of the backchamber is about 0.00005 m³. The holes 92 are evenly spaced along an imaginary circle on the backplate 38 and a center of each hole 92 is about 0.0342 meter. In this example, the backplate 38 defines six holes 92, each having a radius of about 0.000302 meter. The holes 92 are evenly spaced along an imaginary circle on the backplate 38 and a center of each 92 is aligned with the imaginary circle. The center of the imaginary circle is located coincident with a center of the backplate 38, and the radius of the imaginary circle is about 0.0105 meter. The width of the slot 94 is about 0.0144 meter and the area of the slot 94 is about 0.00179 m².

In an approximately 1.5 inch diameter infrasonic microphone 22, where the radius of the membrane 40 is about 0.0134 meter, the tension of the membrane 40 may be less than about 400 Newton per meter. Further, the resonance frequency of the microphone 22 may be less than about 1500 Hertz. Still further, the volume of the backchamber may be selected to produce a low-frequency air compliance that exceeds the compliance of the membrane 40 by a factor of at least 3. In one example, the radius of the membrane 40 is about 0.0342 meter. In this example, the backplate 38 defines six holes 92, each having a radius of about 0.000302 meter. The holes 92 are evenly spaced along an imaginary circle on the backplate 38 and a center of each hole 92 is aligned with the imaginary circle. The center of the imaginary circle is located coincident with a center of the backplate 38, and the radius of the imaginary circle is about 0.0105 meter. The width of the slot 94 is about 0.0144 meter and the area of the slot 94 is about 0.00179 m².

As discussed herein, the preamplifier board 42 is installed parallel to the base wall 54 and to the membrane 24. This slot 104 between the edge of the preamplifier board 42 and the side wall 44 is small, for example 0.025", to increase acoustic resistance. The combined volumes V₁ and V₂ and the volume in the flexible tube 26 is 5×10⁻⁵ meter³. Because of increased acoustic resistance, pressure equalization takes longer which aids in lower ~3 dB frequency to 0.03 Hertz.

In use, the body coupler 24 or catheter tube 23 detects incident sound pressure from the heart, the uterus, or from any other location of the body where it is placed. For example as shown in FIG. 6, the body coupler(s) 24 may be placed at locations A, P, T, and/or M on the body of the patient. The sound pressure excites the motion of the membrane 40 within the microphone 22. The motion of the membrane 40 changes the capacitance between the membrane 40 and the backplate 38. This electrical signal travels from the backplate 38, through the conductor 36 and to the preamplifier board 42, thereby producing a proportional output voltage at the preamplifier board 42. The preamplifier board 42 is grounded via the conductor 36 to an electronics board which digitizes and provides control, analysis and display of the recorded data. MATLAB may also be used to process the data to generate the real-time spectrograms using short-time Fourier transform (STFT) spectrum of the corresponding data at 146 and 148. The time history and spectrogram of biological signals is transferred by the internet 150 to a remote workstation 152, if desired, for observation and analysis. Examples of such remote workstations 52 may be a remote computer monitor, smartphone or tablet. The signals may be sent via wired connection, or may be wirelessly transmitted, such as by using commercially available Bluetooth module, to PC or laptop for processing. The data is converted in useful visual format also called spectrogram, which may be helpful for physician to diagnose any abnormality. The display of short term spectra is performed in real time, in order to detect the presence of a short term event in the data.

FIGS. 8-17 show charts of infrascope signals collected at locations A, P, and M of FIG. 6 with reference to electrocardiogram signals usually referred as ECG or EKG. FIGS. 18 and 19 show charts of infrascope data as compared to ECG or EKG on two different subjects from 1 Hz through...
1000 Hz. The ECG signals of both subjects are quite different and infrascope signals also follow the trend of ECG.

The infrascope 20 can be used for a stress phonocardiography test. Some heart problems occur only during physical activity. Stress phonocardiography test can be accomplished using the signals from the infrascope 20 immediately before and after walking on a treadmill or riding a stationary bike.

The infrascope 20 may be used for fetal heart monitoring during pregnancy, labor, and delivery to keep track of the heart rate of a fetus and the strength and duration of the contractions of the uterus. External fetal heart monitoring which involves placing the body coupler 24 against the abdomen of the patient, keeps track of the baby’s heart rate while at rest and while moving; measures the number of contractions and how long contractions last during labor and delivery; determines if there is preterm labor. Internal fetal heart monitoring, as shown in FIG. 2 and which uses the catheter 23 as described herein, determines if the stress of labor is threatening the baby’s health; measures the strength and duration of the labor contraction.

The infrascope 20 can be used for Doppler phonocardiography as shown in FIG. 3. A Doppler phonocardiography can be used to measure blood flow within the heart without an invasive procedure. The left ventricular filling pressure and blood flow can be estimated by using two infrasopes 20. The infrasopes 20 can be placed at any desired location, for example locations A, P, T and M, by using a mounting structure 160 having adjustable rods 162 attached to the microphones 22 to determine two dimensional velocity estimation and imaging.

The infrascope 20 can be used for biometric identification. Fingerprints have been used for identification for more than 100 years, but using heartbeat for biometric identification has some advantages such as convenience and security. The heartbeat signatures can be extracted using either ECG/EKG or by using the infrascope 20 at remote locations. The security feature is preserved from the fact that a user’s ECG or acoustic signatures cannot be captured without a person’s consent. Another disadvantage of fingerprints are that these can be replicated by using samples left behind. The infrasonic bandwidth signals provide better and higher signal to noise ratio values and another tool for biometric identification.

The infrascope 20 can be used for polygraphs. Physiological processes measured by polygraphs are cardiovascular, electrodermal, and respiratory. The direction and extent of cardiovascular reactivity may be different across individuals in response to stimuli that may be considered arousing. Electrodermal activity in terms of skin resistance or conductance is measured by passing a current through the skin. In response to controlled and relevant questions, variations from basal levels are called electrodermal or EDR responses or electrodermal activity levels and is used for polygraph interpretation. Variations in respiration which also produce changes in heart rate and electrodermal activity is monitored to determine of responses to relevant and control questions are artifacts. Currently, the rate and depth of respiration during polygraph are measured by changes measured using strain gauges positioned on chest and abdomen. Extreme low frequency signal measurements can be made by positioning the infrascope 20 at a subject’s chest and abdomen is a relatively inexpensive tool to measure variation in respiratory and cardiovascular activity.

The infrascope 20 of the present disclosure enables medical personnel to look at the audible bandwidth as well as infrasonic bandwidth, thus providing medical personnel with another tool to analyze physiological processes. The infrascope 20 can be used for respiratory, cardiac, and for fetal heart monitoring. The infrascope 20 enables physiological process signals to be transferred to any place in the world in real time. Ambulances can be equipped with the infrascope 20 and medical personnel are able to obtain a patient’s physiological information in real time. The infrascope 20 is a relatively inexpensive tool to diagnose abnormality at early stage.

The terms “patient” is used throughout the disclosure, which includes humans and animals, as it is anticipated that the present invention would also be capable of monitoring physiological processes for veterinary practices.

All references disclosed herein are hereby incorporated by reference in their entirety.

While particular embodiments are illustrated in and described with respect to the drawings, it is envisioned that those skilled in the art may devise various modifications without departing from the spirit and scope of the appended claims. It will therefore be appreciated that the scope of the disclosure and the appended claims is not limited to the specific embodiment illustrated and discussed with respect to the drawings and that modifications and other embodiments are intended to be included within the scope of the disclosure and appended drawings. Moreover, although the foregoing descriptions and the associated drawings describe example embodiments in the context of certain example combinations of elements and/or functions, it should be appreciated that different combinations of elements and/or functions may be provided by alternative embodiments without departing from the scope of the disclosure and the appended claims.

What is claimed is:

1. An infrasonic stethoscope for monitoring physiological processes of a patient, comprising:
   a microphone capable of detecting acoustic signals in an entire frequency range of 0.03 Hertz to 1000 Hertz, the microphone comprising a body having first and second spaced apart openings;
   a body coupler attached to the first opening of the body to form a substantially air-tight seal, wherein the body coupler is capable of engagement with the patient;
   a flexible tube attached to the body at the second opening; and
   an earpiece attached to the flexible tube.

2. The infrasonic stethoscope of claim 1, wherein the body forms a cavity, and further comprising a conductive backplate within the cavity and defining a backchamber between the backplate and the body, a conductive membrane within the cavity, the backplate and the membrane spaced apart from each other to form a capacitor, and further including a preamplifier board in electrical connection with the backplate, the preamplifier capable of measuring the capacitance between the membrane and the backplate and converting this measured capacitance into voltage.

3. The infrasonic stethoscope of claim 2, wherein the preamplifier board, the backplate and the membrane are parallel to each other.

4. The infrasonic stethoscope of claim 2, wherein the backplate defines a plurality of holes, wherein a slot is defined between an outer diameter of the backplate and an inner wall of the body, and wherein locations and sizes of the holes and a size of the slot are selected such that membrane motion is substantially critically damped.

5. The infrasonic stethoscope of claim 2, further comprising a conductive support plate mounted to said body, the support plate having a passageway and at least one aperture.
therethrough; an insulating member extending through the passageway in said support plate; a conductive member extending through the insulating member and extending therefrom, the conductive member is electrically connected to the backplate and to the preamplifier board, wherein the backplate is on one side of the support plate and the preamplifier board is on an opposite side of the support plate.

6. The infrasonic stethoscope of claim 5, wherein the backplate is seated on the insulating member.

7. The infrasonic stethoscope of claim 2, wherein a slot is defined between the preamplifier board and the body having a length of approximately 0.025 inch.

8. The infrasonic stethoscope of claim 7, wherein the preamplifier board defines a first chamber in the body having a volume of approximately 0.1287 cubic inch, and a second chamber in the body having a volume of approximately 0.6 cubic inch.

9. The infrasonic stethoscope of claim 2, wherein the preamplifier board defines a first chamber in the body having a volume of approximately 0.1287 cubic inch, and a second chamber in the body having a volume of approximately 0.6 cubic inch.

10. The infrasonic stethoscope of claim 2, further comprising a digitizer board which is remote from the microphone.

11. The infrasonic stethoscope of claim 2, wherein the signals from the preamplifier board are digitized and electronically transmitted to a remote location.

12. The infrasonic stethoscope of claim 11, wherein the signals are electronically transmitted by wire.

13. The infrasonic stethoscope of claim 11, wherein the signals are electronically transmitted wirelessly.

14. The infrasonic stethoscope of claim 1, wherein the body coupler is formed of a ring having a flexible, non-conductive diaphragm attached thereto, and the ring is attached to the body.

15. The infrasonic stethoscope of claim 14, wherein the ring and body are threadedly connected.

16. The infrasonic stethoscope of claim 1, wherein the body coupler is formed of a wall having an outlet port, and a catheter attached to the outlet port.

17. The infrasonic stethoscope of claim 1, wherein the body has an outlet port surrounding the second opening and the flexible tube is threadedly attached to the outlet port.

18. The infrasonic stethoscope of claim 1, further comprising a digitizer board which is remote from the microphone.

19. The infrasonic stethoscope of claim 1, wherein the signals from the preamplifier board are digitized and electronically transmitted to a remote location.

20. The infrasonic stethoscope of claim 19, wherein the signals are electronically transmitted in real time.

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