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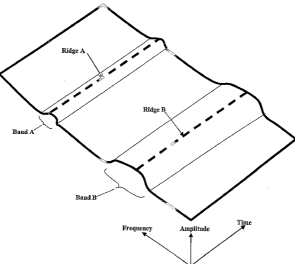
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[Continued on next page]

(54) Title: METHOD OF ANALYSING AND PROCESSING SIGNALS



(57) Abstract: A physiological measurement system is disclosed which can take a pulse oximetry signal such as a photoplethysmogram from a patient and then analyse the signal to measure physiological parameters including respiration, pulse, oxygen saturation and movement. The system can be used as a general monitor, or more specifically, to for infant or adult apnea, and to guard against sudden infant death syndrome. The system comprises a pulse oximeter which includes a light emitting device and a photodetector attachable to a subject to obtain a pulse oximetry signal; analogue to digital converter means arranged to convert said pulse oximetry signal into a digital pulse oximetry signal; signal processing means suitable to receive said digital pulse oximetry signal and arranged to decompose that signal by wavelet transform means; feature extraction means arranged to derive physiological information from the decomposed signal; an analyser component arranged to collect information from the feature extraction means; and data output means arranged in communication with the analyser component.



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Τ	METHOD OF ANALYSING AND PROCESSING SIGNALS
2	•
3	1. Introduction: Problem Domain / Field of Invention
4	
5	The present invention relates to a method of
6	analysing and processing signals. More specifically
7	the invention relates to the analysis and processing
8	of photoplethysmogram (PPG) signals. The invention
9	uses wavelet transform methods to derive clinically
10	useful information from the PPG including
1	information regarding the respiration, pulse, oxygen
.2	saturation, and patient movement. This information
L3	may be used within a device to monitor the patient
L 4	within a range of environments including the
L5	hospital and home environments. In one preferred
L6	embodiment the device may be used to detect
L7	irregularities in one or more of the derived
L8	signals: respiration, pulse, oxygen saturation and
L9	movement. The device allows output of this
20	information in a clinically useful form and
21	incorporates an alarm which is triggered when one or
2.2	a combination of signal irregularities are detected.

2

1	Of particular note is that the utility of current
2	pulse oximeter devices is greatly increased through
3	the provision of a robust measure of patient
4	respiration directly from the PPG signal.
5	
6	2. Background
7	
8	2.1 Blood Oxygen Saturation and its Measurement
9	Oximetry is an optical method for measuring oxygen
10	saturation in blood. Oximetry is based on the
11	ability of different forms of haemoglobin to absorb
12	light of different wavelengths. Oxygenated
13	haemoglobin (HbO_2) absorbs light in the red spectrum
14	and deoxygenated or reduced haemoglobin (RHb)
15	absorbs light in the near-infrared spectrum. When
16	red and infrared light is passed through a blood
17	vessel the transmission of each wavelength is
18	inversely proportional to the concentration of HbO_2
19	and RHb in the blood. Pulse oximeters can
20	differentiate the alternating light input from
21	arterial pulsing from the constant level
22	contribution of the veins and other non-pulsatile
23	elements. Only the alternating light input is
24	selected for analysis. Pulse oximetry has been shown
25	to be a highly accurate technique. Modern pulse
26	oximeter devices aim to measure the actual oxygen
27	saturation of the blood (SaO_2) by interrogating the
28	red and infrared PPG signals. This measurement is
29	denoted SpO_2 . The aim of modern device manufacturers
30	is to achieve the best correlation between the pulse

oximeter measurement given by the device and the

actual blood oxygen saturation of the patient. It is

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known to those skilled in the art that in current 1 2 devices a ratio derived from the photoplethysmogram 3 (PPG) signals acquired at the patients body is used to determine the oxygen saturation measurement using 4 a look up table containing a pluracy of 5 6 corresponding ratio and saturation values. Modern pulse oximeter devices also measure patient heart 7 8 rate. Current devices do not provide a measure of 9 respiration directly from the PPG signal. Additional expensive and obtrusive equipment is necessary to 10 11 obtain this measurement. 12 2.2 Time-Frequency Analysis in Wavelet Space 13 14 The wavelet transform of a signal x(t) is defined as 15 $T(a,b) = \frac{1}{\sqrt{a}} \int_{-\infty}^{+\infty} x(t) \psi^*(\frac{t-b}{a}) dt$ 16 [1] 17 18 where $\psi^*(t)$ is the complex conjugate of the wavelet 19 function $\psi(t)$, a is the dilation parameter of the wavelet and b is the location parameter of the 20 wavelet. The transform given by equation (1) can be 21 used to construct a representation of a signal on a 22 transform surface. The transform may be regarded as 23 a time-scale representation or a time-frequency 24 representation where the characteristic frequency 25

associated with the wavelet is inversely

proportional to the scale a. In the following

discussion 'time-scale' and 'time-frequency' may be

required for the implementation within a time-scale

interchanged. The underlying mathematical detail

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or time-frequency framework can be found in the 1 general literature, e.g. the text by Addison (2002). 2 3 4 The energy density function of the wavelet transform, the scalogram, is defined as 5 6 $S(a,b) = |T(a,b)|^2$ 7 [2] 8 where '| ' is the modulus operator. The scalogram 9 may be rescaled for useful purpose. One common 10 rescaling is defined as 11 12 $S_R(a,b) = \frac{\left|T(a,b)\right|^2}{a}$ 13 [3] 14 and is useful for defining ridges in wavelet space 15 when, for example, the Morlet wavelet is used. 16 17 Ridges are defined as the locus of points of local maxima in the plane. Any reasonable definition of a 18 ridge may be employed in the method. We also include 19 as a definition of a ridge herein paths displaced 20 from the locus of the local maxima. A ridge 21 associated with only the locus of points of local 22 maxima in the plane we label a 'maxima ridge'. For 23 practical implementation requiring fast numerical 24 computation the wavelet transform may be expressed 25 in Fourier space and the Fast Fourier Transform 26 (FFT) algorithm employed. However, for a real time 27 application the temporal domain convolution

expressed by equation (1) may be more appropriate.

In the discussion of the technology which follows

herein the 'scalogram' may be taken to the include

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1 all reasonable forms of rescaling including but not

- 2 limited to the original unscaled wavelet
- 3 representation, linear rescaling and any power of
- 4 the modulus of the wavelet transform may be used in

5 the definition.

6

7 As described above the time-scale representation of

- 8 equation (1) may be converted to a time-frequency
- 9 representation. To achieve this, we must convert
- 10 from the wavelet a scale (which can be interpreted
- 11 as a representative temporal period) to a
- 12 characteristic frequency of the wavelet function.
- 13 The characteristic frequency associated with a
- 14 wavelet of arbitrary a scale is given by

15

$$16 f = \frac{f_c}{a} [4]$$

17

where f_c , the characteristic frequency of the mother

- wavelet (i.e. at a=1), becomes a scaling constant
- 20 and f is the representative or characteristic
- 21 frequency for the wavelet at arbitrary scale a.

22

- 23 Any suitable wavelet function may be used in the
- 24 method described herein. One of the most commonly
- used complex wavelets, the Morlet wavelet, is
- 26 defined as:

27

28
$$\psi(t) = \pi^{-\frac{1}{4}} \left(e^{i2\pi f_0 t} - e^{-(2\pi f_0)^2/2} \right) e^{-t^2/2}$$
 [5]

- 30 where f_0 is the central frequency of the mother
- 31 wavelet. The second term in the brackets is known as

6

the correction term, as it corrects for the non-zero mean of the complex sinusoid within the Gaussian window. In practice it becomes negligible for values of $f_0 >> 0$ and can be ignored, in which case, the Morlet wavelet can be written in a simpler form as

6

7
$$\psi(t) = \frac{1}{\pi^{1/4}} e^{i2\pi f_0 t} e^{-t^2/2}$$
 [6]

8

This wavelet is simply a complex wave within a 9 10 Gaussian envelope. We include both definitions of the Morlet wavelet in our discussion here. However, 11 note that the function of equation (6) is not 12 strictly a wavelet as it has a non-zero mean, i.e. 13 the zero frequency term of its corresponding energy 14 spectrum is non-zero and hence it is inadmissible. 15 However, it will be recognised by those skilled in 16 the art that it can be used in practice with $f_0 >> 0$ 17 with minimal error and we include it and other 18 similar near wavelet functions in our definition of 19 a wavelet herein. A more detailed overview of the 20 underlying wavelet theory, including the definition 21 of a wavelet function, can be found in the general 22 literature, e.g. the text by Addison (2002). Herein 23 we show how wavelet transform features may be 24 extracted from the wavelet decomposition of pulse 25 oximeter signals and used to provide a range of 26 clinically useful information within a medical 27 device. 28

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31 3. Wavelet Feature Extraction

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In this section, methods are described for the 1 2 extraction and use of wavelet features from the PPG signals for use in the provision of clinically 3 useful information. These are incorporated within a 4 medical device and the information is output in a 5 range of formats for use in the monitoring of the 6 patient. The device comprises four key components 7 for the utilization of the wavelet transform 8 information, these are the Pulse Component, 9 Respiration Monitoring Component, Oxygen Saturation 10 Component and the Movement Component. The underlying 11 theory pertaining to these components is detailed 12 below. 13 14 3.1 Pulse Component 15 Pertinent repeating features in the signal gives 16 17 rise to a time-frequency band in wavelet space or a rescaled wavelet space. For example the pulse 18 component of a photoplethysmogram (PPG) signal 19 produces a dominant band in wavelet space at or 20 around the pulse frequency. Figure 1(a) and (b) 21 contains two views of a scalogram derived from a PPG 22 23 signal. The figures show an example of the band caused by the pulse component in such a signal. The 24 pulse band is located between the dashed lines in 25 the plot of figure 1(a). The band is formed from a 26 series of dominant coalescing features across the 27 scalogram. This can be clearly seen as a raised band 28 across the transform surface in figure 1(b) located 29 within a region at just over 1Hz, i.e. 60 breaths 30 per minute. The maxima of this band with respect to 31 frequency is the ridge. The locus of the ridge is 32

1	shown as a black curve on top of the band in figure
2	1(b). By employing a suitable rescaling of the
3	scalogram, such as that given in equation 3, we can
4	relate the ridges found in wavelet space to the
5	instantaneous frequency of the signal. In this way
6	the pulse frequency (pulse rate) may be obtained
7	from the PPG signal. Instead of rescaling the
8	scalogram, a suitable predefined relationship
9	between the frequency obtained from the ridge on the
LO	wavelet surface and the actual pulse frequency may
11	also be used to determine the pulse rate.
1.2	
13	By mapping the time-frequency coordinates of the
14	pulse ridge onto the wavelet phase information
15	gained through the wavelet transform, individual
16	pulses may be captured. In this way both times
1.7	between individual pulses and the timing of
18	components within each pulse can be monitored and
19	used to detect heart beat anomalies, measure
20	arterial system compliance, etc. Alternative
21	definitions of a ridge may be employed. Alternative
22	relationships between the ridge and the pulse
23	frequency may be employed.
24	
25	3.2 Respiration Monitoring Component
26	The respiration monitoring component uses wavelet
27	based methods for the monitoring of patient
28	respiration. This can include the measurement of
29	breathing rate and the identification of abnormal
30	breathing patterns including the cessation of
31	breathing. A key part of the respiration monitoring
32	component is the use of secondary wavelet feature

9

decoupling (SWFD) described below. The information 1 2 concerning respiration gained from the application of SWFD can then be compared and/or combined with 3 4 respiration information from other methods to provide a respiration measure output. 5 6 7 As stated above, pertinent repeating features in the signal give rise to a time-frequency band in wavelet 8 9 space or a rescaled wavelet space. For a periodic signal this band remains at a constant frequency 10 level in the time frequency plane. For many real 11 12 signals, especially biological signals, the band may be non-stationary; varying in characteristic 13 frequency and/or amplitude over time. Figure 2 shows 14 15 a schematic of a wavelet transform of a signal containing two pertinent components leading to two 16 17 bands in the transform space. These bands are 18 labeled band A and band B on the three-dimensional (3-D) schematic of the wavelet surface. We define 19 the band ridge as the locus of the peak values of 20 these bands with respect to frequency. For the 21 22 purposes of the discussion of the method we assume that band B contains the signal information of 23 24 interest. We will call this the 'primary band'. In 25 addition, we assume that the system from which the signal originates, and from which the transform is 26 27 subsequently derived, exhibits some form of coupling between the signal components in band A and band B. 28 29 30 When noise or other erroneous features are present in the signal with similar spectral characteristics 31 32 of the features of band B then the information

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within band B can become ambiguous, i.e. obscured, 1 2 fragmented or missing. In this case the ridge of band A can be followed in wavelet space and 3 4 extracted either as an amplitude signal or a frequency signal which we call the 'ridge amplitude 5 6 perturbation (RAP) signal' and the 'ridge frequency 7 perturbation (RFP) signal' respectively. The RAP and RFP signals are extracted by projecting the ridge 8 9 onto the time-amplitude or time-frequency planes respectively. The top plots of figure 3 shows a 10 schematic of the RAP and RFP signals associated with 11 ridge A in figure 2. Below these RAP and RFP signals 12 we can see schematics of a further wavelet 13 decomposition of these newly derived signals. This 14 15 secondary wavelet decomposition allows for information in the spectral region of band B in 16 figure 2 to be made available as band C and band D. 17 The ridges of bands C and D can serve as 18 instantaneous time-frequency characteristic measures 19 of the signal components causing bands C and D. This 20 21 method, which we call Secondary Wavelet Feature Decoupling (SWFD), therefore allows information 22 23 concerning the nature of the signal components associated with the underlying physical process 24 causing the primary band B (figure 2) to be 25 extracted when band B itself is obscured in the 26 presence of noise or other erroneous signal 27 28 features. 29 An example of the SWFD method used on a PPG signal 30 31 to detect patient breathing from the ridge

associated with patient pulse is shown in figures 4

11

and 5. During the experiment from which the signal 1 2 was taken the patient was breathing regularly at breaths of 6 seconds duration (= 0.167Hz). 3 4 Figure 4(a) contains the scalogram derived from the 5 6 PPG trace taken during the experiment. Two dominant bands appear in the plot: the pulse band and a band 7 associated with patient breathing. These are marked 8 P and B respectively in the plot. In this example we 9 are concerned with the detection of breathing 10 through time and hence here the breathing band is 11 the primary band. The pulse band appears at just 12 over 1Hz, or 60 beats per minute: the beat frequency 13 of the heart and the breathing band appears at 0.167 14 Hz corresponding to the respiration rate. However, 15 the identification of breathing features is often 16 masked by other low frequency artefact in these 17 signals. One such low frequency artefact feature, 18 'F', is indicated in the plot within the dotted 19 ellipse marked on the scalogram where it can be seen 20 to interfere with the breathing band. Figure 4(b) 21 contains a 3-D view of the scalogram plot shown in 22 figure 4(a). From the 3-D plot we can see that the 23 low frequency artefact feature causes a bifurcation 24 of the breathing band at the location shown by the 25 arrow in the plot. The pulse ridge is also shown on 26 figure 4(b), indicated by the black curve along the 27 pulse band. This is the locus of the maxima with 28 29 respect to frequency along the pulse band. 30 31 Figure 4(c) contains the RAP signal derived from the pulse ridge shown in figure 4(b) where the pulse 32

1	ridge is followed and its amplitude is plotted
2	against time. The top plot of figure 4(c) contains
3	the whole RAP signal. The lower plot of figure 4(c)
4	contains a blow up of the RAP signal over a 50
5	seconds interval. An obvious modulation with a
6	period of 6 seconds can be seen in this blow up. The
7	top plot of figure 4(d) contains the whole RFP
8	signal corresponding to the pulse ridge in figure
9	4(b). The lower plot of figure 4(d) contains a blow
LO	up of the RFP signal over 50 seconds. Again an
L1	obvious modulation (of 6 second period) can be seen
L2	in this blow up.
L3	
L4	A second wavelet transform was then performed on the
L5	RAP and RFP signals. The resulting scalograms
L6	corresponding to the RAP and RFP signals are shown
L7	in figures 5a and 5b respectively and the 3-D plots
18	of these scalograms are shown in figures 5c and 5d
L9	respectively. The breathing ridges derived from the
20	RAP and RFP scalograms are superimposed on the 3-D
21	scalograms. The RAP scalogram is the cleaner of the
22	two and can be seen not to contain interference from
23	the artefact feature 'F' found in the original
24	signal scalogram of figure 4(a). For this example
25	the RAP scalogram provides the best solution for the
26	removal of erroneous signal features and the
27	identification of the breathing band when compared
28	to the original scalogram and the RFP scalogram. In
29	practice all three scalograms are compared and the
30	optimal scalogram or combination of scalograms for
31	the extraction of the information required is
32	determined.

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	Through experimentation covering a variety of
3	patient groups (e.g. adult, child, neonate) we have
4	found that for certain signals the method can be
5	enhanced by incorporating paths displaced from the
6	band ridge in the SWFD method. In these cases the
	RAP signals derived from the displaced path exhibits
7	
8	much larger oscillations (compared to the low
9	frequency background waveform) than those of the
10	original ridge path. We find that this enhancement
11	allows us to better detect the breathing component
12	within the SWFD method. Hence we extend our
13	definition of a surface ridge as employed in the
14	method to include paths displaced from the locus of
15	the peak values, contours at a selected level of the
16	pulse band, and in general any reasonably
17	constructed path within the vicinity of the
18	pertinent feature under investigation, where the
19	vicinity is taken to be within the region of the
20	corresponding band.
21	
22	From the above example it can be seen how a
23	secondary wavelet transform of wavelet transform
24	ridge information derived from the pulse band ridge
25	may be used to provide a clearer manifestation of
26	the breathing features in wavelet space from which
27	pertinent breathing information may be derived.
28	
29	The SWFD method described above can form the basis
30	of completely new algorithms for incorporation
31	within devices which require the detection of
32	otherwise masked signal components. Herein, we show

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the application of the method to the detection of 1 breathing features from within the 2 photoplethysmogram, although it will be recognised 3 by those skilled in the art that the method may be 4 applied to other problematic signals. 5 6 In practice, both the original direct observation of 7 the primary band and the indirect observation 8 through perturbations to the secondary band may be 9 employed simultaneously and the optimal time-10 frequency information extracted. 11 12 Those skilled in the art will recognise that 13 modifications and improvements can be incorporated 14 within the methodology outlined herein without 15 departing from the scope of the invention. 16 17 Those skilled in the art will recognise that the 18 above methods may be performed using alternative 19 time-frequency representations of the signals where 20 the amplitude in the time-frequency transform space 21 can be related to the amplitude of pertinent 22 features within the signal. In addition the 23 decomposition of the original signal and the 24 subsequent decompositions of the RFP and RAP 25 scalograms may be performed, each with a different 26 time-frequency method. However, in the preferred 27 method the continuous wavelet transform is employed 28 in all decompositions, although different wavelet 29 functions may be employed in each of the wavelet 30 transforms employed in the method. 31 32

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The preferred method detailed herein departs from 1 2 alternate methods to probe the time-frequency information within wavelet space which follow paths 3 4 of constant frequency in wavelet space. The current 5 method involves following a selected path in wavelet space from which new signals are derived. This 6 7 allows signal components with non-stationary frequency characteristics to be followed and 8 9 analysed to provide information of other signal 10 components which may also exhibit non-stationary 11 behaviour. 12 13 It will be obvious to those skilled in the art that 14 the method relies on high resolution in wavelet 15 space hence the continuous wavelet transform is the preferred method. (The time-frequency discretisation 16 17 employed by the discrete wavelet transform and the stationary wavelet transform is, in general, too 18 coarse for the useful application of the method.) 19 20 The continuous wavelet transform is implemented in the method through a fine discretisation in both 21 22 time and frequency. 23 24 Although the method herein has been described in the 25 context of the detection of breathing features from the pulse band of the wavelet transform of the 26 photoplethysmogram, those skilled in the art will 27 recognise that the method has wide applicability to 28 29 other signals including, but not limited to: other 30 biosignals (e.g. the electrocardiogram, 31 electroencephalogram, electrogastrogram, 32 electromyogram, heart rate signals, pathological

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sounds, and ultrasound), dynamic signals, non-1 destructive testing signals, condition monitoring 2 signals, fluid signals, geophysical signals, 3 astronomical signals, electrical signals, financial 4 signals including financial indices, sound and 5 speech signals, chemical signals, and meteorological 6 signals including climate signals. 7 8 In summary a method for the decomposition of signals 9 using wavelet transforms has been described which 10 allows for underlying signal features which are 11 otherwise masked to be detected. The method is 12 described in the following steps 13 14 A wavelet transform decomposition of the (a) 15 signal is made. 16 (b) The transform surface is inspected in the 17 vicinity of the characteristic frequency of the 18 pertinent signal feature to detect the dominant 19 band (the primary band) associated with the 20 pertinent feature. This band is then 21 interrogated to reveal information 22 corresponding to the pertinent feature. This 23 interrogation may include ridge following 24 methods for identification of localised 25 frequencies in the time-frequency plane. 26 A secondary band is then identified (c) 27 outwith the region of the pertinent feature and 28 its ridge identified. 29 The time-frequency and time-amplitude 30 locus of points on the secondary ridge are then 31 extracted. These new signals are denoted the 32

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'ridge amplitude perturbation (RAP) signal' and 1 the 'ridge frequency perturbation (RFP) signal' 2 3 respectively. A wavelet transformation of the RAP and (e) 4 RFP signals is then carried out to give the RAP 5 and RFP scalograms respectively. 6 These secondary scalograms are then 7 (f) interrogated to reveal information in the 8 region of the primary band of the original 9 scalogram. This interrogation may include ridge 10 following methods for identification of 11 localised frequencies in the time-frequency 12 13 plane. (q) The information gained from step (b) and 14 step (f) are then used to provide the optimal 15 signal information pertaining to the signal 16 feature or features under investigation. 17 18 More than one secondary band may be present. These 19 additional secondary bands may be interrogated in 20 the same way, i.e. steps (c) to (g). 21 22 In the context of breathing detection from the 23 photoplethysmogram the 'primary band' referred to in 24 the above is the breathing band and the 'secondary 25 band' is the pulse band. In the method one or more 26 or a combination of PPG signals may be employed. 27 28 In an alternative methodology once the RAP and RFP 29 signals have been abstracted in step (d) these are 30 then interrogated over short segments using an 31 alternative time-frequency or frequency based method 32

18

(e.g. using a standard FFT routine to find a 1 dominant peak associated with the primary band 2 signal) or another method of signal repetition 3 including, but not limited to, turning points of the 4 signal. This may be employed to speed up the 5 computation of the characteristic frequency of the 6 RAP and RFP scalogram bands or to enhance the 7 8 technique. 9 In step (d) above a combination of the RAP and RFP 10 signals may also be used to generate a 11 representative signal for secondary wavelet 12 13 decomposition. 14 Patient respiration information from the secondary 15 wavelet feature decoupling incorporating the RAP and 16 17 RFP signals is used directly to monitor patient respiration. This can include the measurement of 18 breathing rate and the identification of abnormal 19 breathing patterns including the cessation of 20 breathing. Either the RAP-based SWFD or the RFP-21 based SWFD information may be chosen for patient 22 respiration monitoring. Alternatively a combination 23 of both may be employed where the respiration 24 information derived from each method may be graded 25 quantitatively according to a confidence measure. 26 27 Further the respiration information gained from the 28 RAP-based SWFD and the RFP-based SWFD may be 29 compared to and/or combined with respiration 30 information gained from other methods to provide an 31 optimal output for respiration measures including 32

1	respiration rate, breath timings, breathing
2	anomalies, etc. These other methods may include that
3	described in International Patent Application No
4	PCT/GB02/02843 , "Wavelet-based Analysis of Pulse
5	Oximetry Signals" by Addison and Watson. The chosen
6	respiration measure for output will be extracted
7	using a polling mechanism based on a quantitative
8	measure of the quality of the respiration
9	information derived by each method.
10	
11	Figures 6 to 10 illustrate the preferred embodiment
12	of the respiration monitoring methodology. The
13	wavelet transform of the PPG signal (figure 6(a)) is
14	computed. A plot of the resulting scalogram is shown
15	in figure 6(b). The 10 second PPG signal used in
16	this example was taken from a premature neonate. The
17	same methodology also works for adult and child
18	PPGs. The pulse ridge is shown plotted as a black
19	path across the scalogram in figure 6(b) at around
20	2.5Hz - typical for these young patients. The RAP
21	and RFP signals are then derived from the pulse
22	ridge of the wavelet transform. The RAP and RFP
23	signals are shown respectively in figure 6(c) and
24	figure 6(d). Also shown in figure 6(c) is the
25	patient switch signal which shows inspiration and
26	expiration of the patient as a high /low amplitude
27	square wave trace. The switch signal was activated
28	by an observer monitoring the movement of the chest
29	wall of the neonate during the experiment. The
30	turning points in the RAP and RFP signals may be
31	used as an initial detection mechanism for
32	individual breaths. The RFP and RAP signals are

20

assessed for quality using a confidence measure. 1 2 This measure may be based on any reasonable measure including but not limited to the entropy of the 3 4 signals. The signal with the highest confidence is used to extract information on individual breaths 5 and a breathing rate using the average duration of a 6 7 number of recently detected breaths. A second wavelet transform is performed on both signals. The 8 9 result of a second wavelet transform on the RAP signal of figure 6(c) is shown in figure 7(a) and 10 the ridges of this transform surface are extracted 11 12 as shown in figure 7(b). The result of a second wavelet transform on the RFP signal of figure 6(d) 13 is shown in figure 7(c) and the ridges of this 14 15 transform surface are extracted as shown in figure 16 7(d). 17 The extracted ridges from the RFP and RAP signal 18 transforms and the ridges found in the original 19 transform in the region of respiration, shown in 20 21 figures 8(a), (b) and (c) respectively, are then 22 analysed to determine a composite path which we call the 'selected respiration path' SRP. The analysis 23 may include, but is not limited to, the intensities 24 and locations of the ridges. The SRP represents the 25 most likely breathing components. The SRP derived 26 27 from the extracted ridges shown in figures 8(a), (b) and (c) is shown in figure 8(d). The SRP will 28 29 normally be determined within an initial predetermined "latch-on" time window and reassessed 30 within an updated time window. The ridge selection 31 procedure used to derive the SRP is based upon a 32

21

decision tree implementing a weighted branching 1 dependent upon, but not limited to, the following 2 local (i.e. relationship between ridge components 3 within a particular ridge set) and global (i.e. the 4 inter-relationship between ridge components across 5 ridge sets) criteria: start and end position, 6 length, average and peak strengths, various spatial 7 (i.e. movement range over the time-frequency 8 surface) statistical parameters including variance 9 and entropy, and a measurement of relative 10 switchback positions (i.e. degree of overlap with 11 other ridges). These criteria are based on results 12 of our in house experimentation across a range of 13 patient categories: adult, child and neonate. 14 15 A confidence metric for the accuracy of the SWFD 16 ridge obtained from the RAP signal can also be 17 acquired by comparing the resultant SWFD ridge 18 intensities derived from the RAP signal of the band 19 maxima ridge and ridges off-set from it. When 20 compared to RAP-SWFD derived from the band maxima 21 ridge, the off-ridge transform's ridges associated 22 with respiration have been observed to increase (to 23 a maximum) in intensity as the displacement of the 24 off-ridge from the maxima ridge is increased. Those 25 ridges associated with other features, however, 26 remain relatively static in amplitudes. In this way, 27 by interrogating the ridge amplitudes of a plurality 28 of RAP signals derived from the band maxima offsets, 29 the ridge or ridges associated with respiration can 30 be identified through a significant change in 31 amplitude relative to others. 32

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1	
2	The selected ridge path (SRP) is then used to
3	provide an overall confidence as to breathing rate
4	and/or provide individual breath monitoring and/or
5	prediction. By superimposing the SRP shown in figure
6	8(d) onto the phase information derived from the
7	original transform the phase along the SRP can be
8	determined as shown in figure 9. In this way
9	individual breaths may be identified through the
10	behaviour of the phase cycling. This phase
11	information along the SRP path may used to derive a
12	breathing signal either by displaying the phase
13	information as shown in figure 9 or by taking the
14	cosine, or similar function, of the phase
15	information to produce a sinusoidal waveform, or by
16	some other method to provide a waveform of choice
17	for visual display of the breathing signal. In an
18	alternative embodiment the phase information from
19	one of the secondary transforms or a combination of
20	the phase information from all transforms may be
21	used in the method. In addition, the phase
22	information used may be processed to remove
23	erroneous phase information for example caused by
24	movement artifact.
25	
26	Parts of the SPR may contain missing segments caused
27	by, for example, signal artefact. In these regions
28	the SRP may be inferred using the last available
29	path point and the next available path point as
30	shown schematically in figure 10. In the preferred
31	embodiment this is carried out using a linear fit
32	between the points. However, other methods may also

23

1 be used without departing from the scope of the

2 invention.

3 4

3.3 Oxygen Saturation Measurement

5 The amplitude of signal features scale with their

6 wavelet transform representation. Thus by dividing

7 components of the wavelet transform of the red PPG

8 signal by those of the infrared PPG signal we obtain

9 new wavelet-based representations which contain

10 useful information on the signal ratios for use in

11 the determination of oxygen saturation. If a complex

wavelet function is used this information may be

13 extracted using a suitable path defined on the ratio

of the moduli of the transforms or using a Lissajous

15 plot from the real or imaginary parts of the

16 transforms. If a wavelet function containing only a

17 real part is employed then this information should

18 be extracted using a Lissajous plot derived from the

19 transforms. Two complimentary methods for the

20 extraction of the wavelet-based ratio information

21 required for the determination of oxygen saturation

22 are given below.

23

24 Figure 11 shows the three dimensional plots of the

25 real-parts of the wavelet transforms of the

26 simultaneously collected red and infrared PPG

27 signals. A complex Morlet wavelet was used in the

28 transform. The dominant nature of the pulse band and

29 breathing band regions is evident in the figure.

These are marked 'B' and 'C' respectively in the

31 figure. A secondary band containing pulse components

32 can also be seen in the figure (marked 'A'). This

1	band is associated with the double humped morphology
2	of the PPG waveform. In the new wavelet-based
3	Lissajous method a number of frequency levels are
4	selected within a moving window. The moving window
5	is shown schematically on the plot in figure 12.
6	(Here we use a 4.56 second window for the purpose of
7	illustration although alternative window lengths may
8	be used as required.) The oscillatory nature of the
9	pulse band and breathing band regions is evident in
10	the plot. The wavelet transform values along each of
11	these frequency levels for the red and infrared
12	signals are plotted against each other to give a
13	Wavelet-Based Lissajous (WBL) plot. This results in
14	a multitude of WBL plots, one for each frequency
15	level selected. In the method, the selected
16	frequency levels lie in the range of expected pulse
17	frequencies which is, for the purposes of
18	illustration, herein defined as between 0.67 and
19	3.33 Hz. This range may be altered to reflect the
20	application. The multitude of WBL plots may be
21	displayed together to form a 3-D Lissajous figure,
22	as shown in figure 13(a).
23	
24	Note that, in the example shown here, a complex
25	wavelet function was used and hence both real or
26	both imaginary values of the transform can be
27	utilized in the method. Further, information from
28	the real WBL plots and imaginary WBL plots may be
29	combined to provide an optimal solution. If a real-
30	only wavelet function is used (i.e. a wavelet
31	function containing only a real part and no

imaginary part) then only one set of transforms

1

2 (real) are available to use. 3 Each Lissajous plot making up the 3-D Lissajous 4 figure is then probed to find its spread both along 5 its principle axis and that axis orthogonal to it. 6 To do this, any reasonable measure of spread may be 7 used. Here we employ the standard deviation (SD). 8 Figure 13(b) shows an end on view of the 3-D 9 Lissajous of figure 13(a). The region of the 3-D 10 Lissajous figures 13(a) and 13(b) in the vicinity of 11 the pulse frequency is marked by the letter 'B' in 12 the figures and higher frequencies are marked by the 13 letter 'A'. Figure 14 contains plots of the standard 14 deviation of data spread along the principle axis 15 (top plot) and minor axis (middle plot), and the 16 ratio of the standard deviations (lower plot) for 17 each Lissajous component making up the 3-D Lissajous 18 plot in figure 13(a). In the preferred embodiment 19 the Lissajous component with the maximum spread is 20 used in the determination of the oxygen saturation. 21 The location of this component is marked by the 22 arrow in the top plot of figure 14. This component, 23 with the maximum spread along the major principle 24 axis, is plotted in figure 13(c): the representative 25 slope of which is computed and used to determine the 26 local oxygen saturation value using a predefined 27 look-up table. This maximum spread is usually found 28 at or near the pulse frequency. A check is also made 29 on the SD ratios: defined as the SD of spread along 30 the major axis divided by the SD of spread along the 31 minor axis. A low SD ratio implies good correlation 32

26

between the two signals. The SD ratio for the 1 component with maximum spread is indicated by the 2 arrow in the lower plot of figure 14. We can see for 3 this case that a relatively low SD ratio occurs at 4 this location. The SD ratio check may be used to 5 pick a more appropriate wavelet-based Lissajous plot 6 and can form part of a noise identification and/or 7 reduction algorithm. Alternate methods of picking an 8 optimal wavelet-based Lissajous may also be employed 9 as appropriate. During periods of excessive noise, 10 the Lissajous components can become spread out in 11 shape, and in some cases the direction of the major 12 and minor principle axis can significantly change 13 from that of the relatively noise free portions of 14 the signals. A check can therefore be made to 15 determine if this has occurred by retaining a recent 16 history of the selected Lissajous components. This 17 can further be used as a confidence check on the 18 selected Lissajous figure used in the determination 19 of oxygen saturation. 20 21 Note that the ratio of the amplitudes of the 22 independent wavelet signals making up the selected 23 Lissajous component may also be used to determine 24 the oxygen saturation. Note also that the inverse 25 transform of these wavelet signals may be used in 26 the method to determine oxygen saturation. The 27 method described can be used to extract the 28 pertinent ratio information from wavelet transforms 29 computed using either complex or real-only wavelet 30 functions. 31 32

27

ㅗ	rigure is shows the oxygen saturation determined
2	using the 3-D Lissajous method (solid black line)
3	compared with the traditional signal amplitude
4	method (dotted) and signal Lissajous method
5	(dashed). All three methods employed a 4 second
6	smoothing window. It can be seen that for the
7	particular example signal interrogated here (the
8	signals taken from the finger of a healthy male
9	patient aged 42 sitting in an upright position at
LO	rest) the wavelet method produces a more consistent
L1	value.
L2	
L3	Figures 16 contains three-dimensional views of the
L 4	red and infrared scalograms corresponding to an
1.5	example PPG signal. Here the modulus of the complex
16	transform is used. The locations of the band
L7	associated with the pulse component are indicated in
18	the plots (denoted 'B' in the figures). We define
19	the collection of points corresponding to the path
20	of the maxima of the band projected onto the time
21	frequency plane as $P.$ A wavelet ratio surface (R_{WT})
22	can be constructed by dividing the wavelet transform
23	of the logarithm of red signal by the wavelet
24	transform of the logarithm of the infrared signal to
25	get a time-frequency distribution of the wavelet
26	ratio surface, i.e.
27	
28	
	T(a,b)

29
$$R_{WT} = \frac{|T(a,b)_R|}{|T(a,b)_m|}$$
 [7]

30

28

where where the subscripts R and IR identify the red 1 and infrared signals respectively. The wavelet ratio .2 surface derived from the two scalograms in figure 16 3 is shown schematically in figure 17. Note that as 4 described previously in our definition of scalogram 5 we include all reasonable forms of rescaling 6 including the original unscaled wavelet 7 representation, linear rescaling and any power of 8 the modulus of the wavelet transform may be used in 9 the definition. As the amplitude of the wavelet 10 components scale with the amplitude of the signal 11 components then for regions of the surface not 12 affected by erroneous signal components the wavelet 13 ratio surface will contain values which can be used 14 to determine the oxygen saturation using a pre-15 defined look-up table. 16 17 As can be seen in figure 17, the time frequency 18 wavelet ratio surface along, and in close proximity 19 to, the projection of the pulse ridge path P onto 20 the wavelet ratio surface are stable and hence may 21 be used in the robust determination of the oxygen 22 saturation. In the preferred embodiment the values 23 obtained along the projection of P onto R_{WT} are used 24 to determine oxygen saturation via a pre-defined 25 look-up table which correlates R_{WT} to oxygen 26 saturation. 27 28 A 2-D or 3-D view of the R_{WT} plot may be computed and 29 30 displayed in real time to provide a visual indication of the quality of the ratio of ratios 31

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obtained by the method, and hence the quality of the 1 measurement of oxygen saturation. 2 3 Figure 18 contains a plot of the end view of the 4 wavelet ratio surface shown in figure 17. From the 5 figure we see that a relatively stable, flat region 6 is also found at or near the respiration frequency 7 (R in the figure). It has been noted from 8 experimentation that for some cases the respiration 9 region of the wavelet ratio surface may lie at a 10 different level from the pulse band region. Hence, 11 for these cases, using R_{WT} obtained in the breathing 12 region would produce erroneous values of oxygen 13 saturation. By following a path in the region of the 14 pulse band our method automatically filters out 15 erroneous breathing components in the signal. 16 17 Figure 19 contains a plot of the oxygen saturation 18 determined by the wavelet ratio surface method as a 19 function of time as compared with two standard 20 methods: the traditional signal amplitude method and 21 the traditional Lissajous method. The PPG signals 22 were again taken from the finger of a healthy male 23 patient aged 42 sitting in an upright position at 24 rest. From visual inspection of the plot it can be 25 seen that, for this example, the wavelet-based 26 method produces a more consistent value of oxygen 27 saturation compared to contemporary methods. 28 29 30 It will be recognized by those skilled in the art that, in an alternative embodiment, the pulse band 31 ridge path P can also be projected onto the real or 32

1	imaginary transform components. From the values of
2	the transform components along this path over a
3	selected time interval a Lissajous figure may be
4	obtained and used in the determination of oxygen
5	saturation. It will also be recognized by those
6	skilled in the art that, in an alternative
7	embodiment, alternative paths may be projected onto
8	the wavelet ratio surface and used for the
9	determination of oxygen saturation. For example in
LO	regions where the pulse band exhibits noise causing
L1	the path of the ridge maxima to move far from the
12	actual pulse frequency a method for detecting such
13	noisy events and holding the path to the most
1.4	appropriate recent pulse frequency may be used until
15	the event has passed or until a preset period of
16	time whereby an alarm is triggered.
17	
18	The 3-D Lissajous and wavelet ratio surface
19	methodologies for the determination of oxygen
20	saturation, as described above, can form the basis
21	of an algorithm for incorporation within pulse
22	oximeter devices. Furthermore the ability of the
23	methodologies to restrict themselves to the optimal
24	wavelet transform values by picking the optimal
25	Lissajous or following the pulse band respectively,
26	allows for erroneous signal elements to be discarded
27	automatically; so leading to a more robust algorithm
28	for the determination of oxygen saturation.
	Tot the determination of oxygen bacaration.
29	TOT the determination of oxygen bacaracton.
29 30	Note that in both new methods the inverse transform

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1	
2	In the preferred embodiment, both the 3-D Lissajous
3	and wavelet ratio surface methods are employed
4	simultaneously and the optimal measured saturation
5	value determined. It is obvious from the above
6	description that the initial inputted signals and
7	wavelet transformation of these signals form common
8	elements to both methods.
9	
10	Those skilled in the art will recognise that
11	modifications and improvements can be incorporated
12	to the methodology outlined herein without departing
13	from the scope of the invention.
14	
15	Those skilled in the art will recognise that the
16	above methods may be performed using alternative
17	time-frequency representations of the signals where
18	the amplitude in the time-frequency transform space
19	can be related to the amplitude of pertinent
20	features within the signal. However, in the
21	preferred method the continuous wavelet transform is
22	employed.
23	
24	In summary a method for the decomposition of pulse
25	oximetry signals using wavelet transforms has been
26	described which allows for underlying
27	characteristics which are of clinical use to be
28	measured and displayed. These wavelet decompositions
29	can then be used to:
30	(a) provide, using information derived from the
31	signal wavelet transforms (i.e. from the original
32	transform, the rescaled wavelet transforms, the

ratio of derived wavelet transforms, the
scalograms, wavelet ridges, etc.) a method for
measuring oxygen saturation.

(b) construct, using information derived from the

wavelet transform (i.e. from the original transform, the rescaled wavelet transforms, the ratio of derived wavelet transforms, the scalograms, wavelet ridges, etc.), a plurality of wavelet-based Lissajous figures from which the optimum Lissajous representation is chosen using preset criteria and the slope of which is used to determine the oxygen saturation of the signal using a look-up table.

(c) construct, using information derived from the wavelet transform (i.e. from the original transform, the rescaled wavelet transforms, the ratio of derived wavelet transforms, the scalograms, wavelet ridges, etc.), a time-frequency equivalent of the ratio of ratios, the wavelet ratio surface, from which to determine the oxygen saturation of the signal by following a selected path through the time frequency plane. The preferred path through the time frequency

(d) provide an optimal oxygen saturation value from those derived in (b) and (c).

plane to be that corresponding to the pulse band.

3.4 The Monitoring of Patient Movement

- 29 Current devices are configured to remove detrimental
- 30 movement artifact from the signal in order to clean
- 31 it prior to determination of the clinical parameter
- of interest, e.g. the pulse rate or oxygen

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saturation. However, the method described herein as 1 embodied within a device monitors general patient 2 movement, including large scale body movements, 3 respiration and the beating heart. In this way the 4 absence of patient movement and/or irregularity of 5 movement can be detected and an alarm triggered. 6 7 Patient movement results in PPG signal artifact. The 8 manifestation of this artifact can be observed in 9 the wavelet transform of the signal. An example of a 10 movement artifact in the scalogram is shown in 11 figure 20(a). The PPG signal from which the wavelet 12 plot was derived was acquired from a premature baby 13 a few weeks after birth. The location of the 14 movement artifact is marked by the arrow in the 15 plot. The breathing band ridge has been superimposed 16 on the wavelet plot (marked R in the figure). The 17 pulse band is marked P in the figure. Notice that 18 the artifact causes a drop-out in the detected 19 breathing ridge (i.e. a missing fragment), and also 20 cuts through the pulse band where it can cause 21 similar drop outs to occur in the detection of the 22 pulse ridge. It has been the focus of pulse oximeter 23 device manufacturers to remove as much of the 24 movement artifact component from the signal while 25 leaving the information necessary to obtain accurate 26 oxygen saturation and pulse rate measurements. In a 27 preferred embodiment of the methods described herein 28 we extract a movement component from the PPG signals 29 for use in the monitoring of patient movement and, 30 in particular, for the monitoring of the movement of 31 infants. 32

1	
2	A three-dimensional view of the scalogram of figure
3	20(a) is plotted in figure 20(b). Here we see the
4	dominance of the movement artifact feature in
5	wavelet space. By identifying such features we can
6	monitor patient movement. It is common for young
7	babies to exhibit very variable respiration patterns
8	and to cease breathing for short periods of time,
9	especially when making a movement of the body. Hence
10	inspecting the derived movement signal when an
11	irregular respiration signal occurs, including
12	cessation of breathing, gives a further measure of
13	patient status.
14	
15	The modulus maxima of the wavelet surface is the
16	loci of the maxima of the wavelet surface with
17	respect to time. Figure 21(a) plots the modulus
18	maxima lines associated with figure 20(a). Figure
19	21(b) shows a three-dimensional view of the
20	transform surface with the modulus maxima lines
21	superimposed. Figure 22(a) shows an end view of the
22	maxima lines (without the surface shown)
23	corresponding to those shown in figures 21(a) and
24	21(b). We can see from the end view that the modulus
25	maxima line corresponding to the movement artifact
26	has a significantly different morphology to the
27	other maxima lines: it covers a large frequency
28	range and contains significantly more energy than
29	the other maxima, especially at low frequencies. By
30	setting amplitude threshold criteria at a frequency
31	or range of frequencies we can differentiate the

modulus maxima of the artifact from other features.

1	An example of this is shown schematically by the				
2	threshold level and frequency range depicted on				
3	figure 22(b), where maxima above the pre-defined				
4	amplitude threshold within a frequency range given				
5	by $f_{(1)} < f < f_{(2)}$ are identified as corresponding to				
6	movement artifact. In addition a check of local				
7	anomalies in the detected pulse and breathing ridges				
8	may also be made. For example modulus maxima which				
9	are at significantly higher amplitudes than the				
LO	pulse ridge mean value in their vicinity are deemed				
L1	to correspond to movement artifact. This is depicted				
L2	in figure 22(c). In addition, modulus maxima which				
13	are at a significantly higher amplitude than the				
L 4	respiration ridge mean value in their vicinity are				
L5	deemed to correspond to movement artifact. This is				
16	depicted in figure 22(d).				
17					
18	A region in the time frequency plane within the				
19	support of the wavelet is then deemed to contain				
20	artifact. The support of the wavelet is taken as a				
21	predefined measure of temporal 'width' of the				
22	wavelet. For wavelets with theoretical infinite				
23	width, such as the Morlet wavelet, the width is				
24	defined in terms of the standard deviation of				
25	temporal spread: for example we use three times the				
26	standard deviation of spread each side from the				
27	wavelet centre. Thus a cone of influence of the				
28	artifact may be defined in the transform plane.				
29					
30	Using the above method we can monitor patient				
31	movement by detecting modulus maxima corresponding				
32	to movement artifact. This information can be used				

to monitor patient movement and/or to provide a				
measure of confidence on the derived values of other				
measurements (e.g. oxygen saturation, pulse and				
respiration). These measurements may, for example be				
held at a previous value until the detected movement				
event has passed.				
Other artefact may exist in the signal which may				
originate from the drive and control electronics				
including, but not limited to, automatic gain				
adjustments. The occurrence of this type of artifact				
will be known and can be accounted for in the signal				
and hence differentiated from movement artifact.				

37

4. Device Configuration and Usage

- 2 The device may be used to monitor one or more of the
- 3 following signals: respiration, pulse, breathing and
- 4 movement. Useful information regarding these signals
- 5 would be displayed on the device or output in a
- 6 suitable format for use.

7

- 8 In one embodiment the device would be used to
- 9 continually monitor one or more of these signals.

10

- In another embodiment the device would be used to
- monitor one or more of these signals intermittently.

13

- 4.1 Device Configuration
- 15 Detailed block diagrams of the device are provided
- 16 in figures 23,24, 25 and 26.

- 18 The following is with reference to figure 23. In the
- 19 present invention signals are acquired at the
- 20 patient's body 10. These are sent for digitization
- 21 11. The links between components of the system may
- 22 be fixed physical or wireless links, for example
- 23 radiofrequency links. In particular, either or both
- of the links between 10 and 11 or 11 and 12, or the
- links between the analyser component and a visual
- 26 display may a wireless link enabled by a
- 27 radiofrequency transmitter. The digitised cardiac
- signals 11 are sent to 12 where in the preferred
- 29 embodiment the natural logarithm of the signals are
- 30 computed. These are then sent to 13 where the
- 31 wavelet transforms of the signals are performed. The
- 32 components of the wavelet transformed signals,

1	including modulus, phase, real part, imaginary part
2	are then sent to 14 where the pulse ridge is
3	identified. The information from 13 and 14 is then
4	used in the extraction of patient pulse information
5	15, oxygen saturation 16, patient movement
6	information 17 and respiration information 18. The
7	information regarding oxygen saturation, pulse,
8	respiration and patient movement is all sent to the
9	Analyser component 19 where it is collected and
10	collated ready for outputting at 20. The oxygen
11	saturation, respiration, pulse rate and movement
12	information is output from the device 20 through a
13	number of methods, which may include a printout, a
14	display screen or other visual device, an audable
15	tone, and electronically via a fixed or remote link.
16	The output information may be sent to a location
17	remote from the patient, for example sent via
18	telephone lines, satellite communication methods, or
19	other methods. Further, real-time wavelet-based
20	visualisations of the signal (including the original
21	transform and/or the wavelet ratio surface with
22	projected pulse ridge path) may be displayed on the
23	device 20. These visualisations will highlight
24	salient information concerning the quality of the
25	outputted measurements. Additional useful
26	information regarding movement artefact and
27	breathing information may be apparent from such a
28	real time display.
29	
30	The workings of components 15, 16, 17 and 18 shown
31	in figure 23 are described below in more detail.
32	

Pulse Component 15: With reference to figure 23, 1 2 pulse information including pulse rate and pulse irregularities are derived at 15 using the 3 instantaneous frequency of the pulse band ridge 4 determined at 14. The instantaneous frequency may 5 correspond directly with the instantaneous ridge 6 frequency or require a mapping from the 7 8 instantaneous ridge frequency and the true respiration rate. Further the method allows for a 9 10 smoothing of this value over a fixed time interval. 11 Further the method allows for erroneous values of the pulse rate derived in this way to be excluded 12 from the outputted values. This component 15 may 13 also be used to measure inter-beat intervals and 14 pertinent pulse wave timings. The pulse information 15 determined at 15 is then sent to the Analyser 16 17 Component 19. 18 19 The Oxygen Saturation Component 16: The following is with reference to figures 23 and 24. The oxygen 20 saturation component 16 shown in figure 23 comprises 21 the subcomponents 31, 32, 33, 34, 35, 36 and 37 as 22 shown in figure 24. The wavelet transform 23 information and pulse ridge information from 14 is 24 input into this module at the feature sorter 31 25 which sends the relevant information to the 26 Lissajous computation unit (components 32, 33 and 27 28 34) and the pulse ridge computational unit (components 35 and 36). A predetermined number of 29 30 wavelet-based Lissajous are computed over the pulse region 32. An automated procedure is employed for 31 the determination of the optimal Lissajous for use 32

30

31

32

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40

in the oxygen saturation calculation 33. In the 1 preferred embodiment this would be achieved by 2 3 comparing the standard deviations of the data spread along of the principle axes of the Lissajous plot. 4 The slope of the principle axis is then used to 5 determine the oxygen saturation using a suitable 6 look-up table which correlates the slope to oxygen 7 saturation 34. The oxygen saturation determined at 8 34 is denoted 'Oxygen Saturation Determination (1)'. 9 10 The information regarding the wavelet transforms of 11 the PPG signals and the path of the pulse ridge is 12 13 collected at the feature sorter 31 used to compute the wavelet ratio surface 35. The wavelet ratio 14 corresponding to the pulse path is determined by 15 projecting the pulse path onto the wavelet ratio 16 surface. This ratio is then used to determine the 17 oxygen saturation using a look-up table which 18 correlates the wavelet ratio to oxygen saturation 19 36. The oxygen saturation determined at 36 is 20 denoted 'Oxygen Saturation Determination (2)'. The 21 two oxygen saturation values (1) and (2) are then 22 used to determine the most appropriate value of 23 oxygen saturation 37. This value is then sent to the 24 25 Analyzer Component 19. 26 Movement Component 17: The following is with 27 reference to figures 23 and 26. The Movement 28 component 17 of figure 23 comprises the 29

subcomponents 51, 52, 53, 54, 55 as shown in figure

ridge information is sent from 14 to the modulus

The wavelet transform information and pulse

41

maxima component **51** where the modulus maxima of the wavelet surfaces are computed. The modulus maxima

3 information is then sent to be analysed for movement

4 artifact. The modulus maxima information is sent to

5 the components 52, 53 and 54. These are described as

6 follows. The Threshold component 52 detects maxima

7 above a preset threshold and within a preset

8 frequency range which are them defined as movement

9 artifact. The Pulse Check component 53 checks the

10 maxima corresponding to the pulse band to see if

anomalously large excursion from the local mean

12 level has occurred. If so movement artifact is

detected. The Respiration Check component 54 checks

14 the maxima in the vicinity of the selected

respiration path SRP obtained from 18 to determine

if anomalously large excursion from the local mean

17 level has occurred. If so movement artifact is

detected. The information from components 52, 53 and

19 54 are then collected and collated at the Movement

20 Signal component 55 where a movement signal is

21 generated. This is then sent to the Analyser

22 Component 19.

23

24 Respiration Component 18: The following is with

reference to figures 23 and 25. The respiration

26 component 17 of figure 23 comprises the

27 subcomponents 61, 62, 63 and 64 as shown in figure

28 25. The wavelet transform and pulse ridge

29 information from 14 are input into this module at

30 component 61 which uses the information to derive

31 the ridge amplitude perturbation (RAP) signal and

32 the ridge frequency perturbation (RFP) signals. The

RAP and RFP signals are derived using the path 1 defined by the projection of the maxima of the pulse 2 band or a locus of points displaced from this maxima 3 path. A secondary wavelet transform is performed on 4 these signals 62 and then passed to the respiration 5 detection component 63 where the respiration ridges 6 are detected for the wavelet transforms of the RFP 7 8 and RAP signals. These are then used within an algorithm which decides the selected respiration 9 path (SRP). This algorithm may also incorporate 10 respiration information using complementary methods 11 64. Note that in the method the original transform 12 obtained at 13 and the secondary transform 62 may be 13 computed using different wavelet functions. The 14 respiration information is then sent to the Analyzer 15 Component 19 and also to the Movement component 17. 16 17 The Analyser Component 19: With reference to Fig. 18 23, the Analyzer Component collects the information 19 from the pulse component 15, Oxygen Saturation 20 Component 16, Movement Component 17 and Respiration 21 Component 18. During periods of detected motion or 22 other signal artifact the analyzer makes a decision 23 to hold the most appropriate recent values of these 24 signals until the artifact event passes or until 25 predetermined interval has passed at which point an 26 alarm signal sent to the device output 20. Further 27 the analyzer checks the incoming signals for 28 anomalous behaviour including, but not limited to: 29 low and or high pulse rates, pulse irregularities, 30 low and high breathing rates, breathing 31 irregularities, low and high oxygen saturation 32

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rates, movement irregularities including excessive 1 2 movement and absence of movement. Detected anomalous behaviour or combination of behaviours will trigger 3 an alarm signal sent to the device output 20. 4 5 6 4.2 Physical Attachment of Probes and Transmission 7 of PPG Signals 8 Referring to figure 23, the acquisition of the signal 10 takes place at a suitable site on the 9 patient's body. This signal is then sent to 10 11 component 11 where the signals are digitized then to component 12 where their natural logarithm is 12 computed prior to the wavelet analysis at 13. The 13 patient signal may be taken using a standard probe 14 configuration. For example a finger or toe probe, 15 foot probe, forehead probe, ear probe and so on. 16 17 Further the probe may function in either transmittance or reflectance mode. 18 19 20 In one preferred embodiment for use with neonates a foot/ankle mounted device such as a cuff is employed 21 as depicted schematically in figure 27. The cuff is 22 used to house the probe electronics, radio frequency 23 transmitter modules and battery. Figure 27(a) shows 24 the patients lower leg 80 and foot with the 25 preferred embodiment of the cuff 83 attached to the 26 foot. The patients heel 81 and toes 82 protrude from 27 the cuff. Figure 27(b) shows two views, one from 28 each side of the foot showing the cuff with 29 compartments for housing the electronic equipment 30 required for signal acquisition and transmission. 31 The PPG signals may be taken directly through the 32

T	root using Light Emitting Drodes (LEDS) 86 and				
2	photodetector 88 located as shown or, in an				
3	alternative embodiment, they may be taken at the toe				
4	using a short length of cable attaching the pulse				
5	oximeter probe to the electronics contained in the				
6	cuff. In a further alternative embodiment				
7	reflectance mode photoplethysmography may be				
8	employed. In a further alternative embodiment more				
9	suitable for adult monitoring the electronic				
10	equipment is packaged within a soft housing which is				
11	wrapped and secured around the wrist as shown in				
12	figure 28. The electronic components for receiving				
13	processing and transmitting the PPGs are housed in a				
14	unit 90 secured by a band 91 to the patients wrist.				
15	The PPG signals are acquired at a site local to the				
16	wrist band. For example from a finger 93 via a lead				
17	92 from the wrist unit 90, or at the site of the				
18	wrist band and housing using, for example,				
19	reflectance mode photoplethysmography. In yet				
20	another alternative embodiment, the signal from the				
21	pulse oximeter probe would be sent to the monitor				
22	device using a physical lead instead of the wireless				
23	method described here.				
24					
25 .	Light transmitters other than LEDs may be used in				
26	the device without departing from the scope of the				
27	invention.				
28					
29	In an alternative embodiment, the digitised signal				
30	from 11 may input directly to the wavelet transform				
31	component 13 without taking the natural logarithm.				
32					

1	In an alternative embodiment, more than two			
2	wavelengths or combination of more than two			
3	wavelengths of light may be employed in the Oximetry			
4	method.			
5				
6				
7				
8				
9 ,				
10	4.3 Use of the Device			
11				
12	4.3.1 General Use			
13	The device may be used for general patient			
14	monitoring in the hospital, home, ambulatory or			
15	other environment. For example in a preferred			
16	embodiment for a device for use within a hospital			
17	setting it may be used to continually or			
18	intermittently monitor patient respiration together			
19	with oxygen saturation and pulse rate.			
20				
21	4.3.2 Embodiment as an Apnea Monitor			
22	In another preferred embodiment of the device it			
23	would be used as an apnea monitor. Apnea is the			
24	cessation of breathing usually occurring during			
25	sleep. There is increasing awareness of this sleep			
26	disorder as the cause of a number of serious medical			
27	conditions in adults and infants. Separate areas of			
28	use are envisaged for the device as an apnea			
29	monitor. Examples of this use include, but are not			
30	limited to: (1) adult monitoring, where it can be			
31	used as a home screening diagnostic tool for			
32	potential appea patients and (2) infant monitoring.			

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where it can be used as either an in hospital or 1 2 home monitoring tool to alert the child's carer to this potentially fatal respiration irregularity. 3 4 5 Apnea monitors monitor heart and respiratory signals to detect apnea episodes - usually defined as 6 cessation of breathing for >20 seconds. Apnea is 7 associated with slowing of the pulse (bradycardia) 8 or bluish discoloration of the skin due to lack of 9 oxygenated haemoglobin (cyanosis). Long term effects 10 11 of apnea in adults are quite serious and have been reported to include: heavy snoring, weariness and 12 obsessive drive to fall asleep, reduced physical and 13 14 mental fitness, strokes, nervousness, fall in concentration and headaches, psychic symptoms up to 15 depressions, sexual dysfunctions, impotence, 16 17 dizziness and nightly perspiration. In babies apnea may lead to death if suitable resuscitation measures 18 are not taken. 19 20 As it measures respiration and movement directly 21 from the pulse oximeter signal (in addition to 22 23 oxygen saturation and pulse), the device can be fitted remote from the head; e.g. the foot or arm of 24 the patient. This has the advantage over current 25 devices which comprise of probes located on the 26 patients head and face to measure breathing at the 27 patients nose and/or mouth. As such they are 28 uncomfortable for adult patients and are quite 29 impractical for fitting to babies for the obvious 30 reason of causing a potential choking hazard. The 31 preferred embodiment of our invention allows the PPG 32

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signal collected at the patient to be sent via a 1 wireless link to a remotely located device. 2 3 In summary, embodied as an apnea monitor, the device 4 provides a method for the acquisition analysis and 5 interpretation of pulse oximeter signals to provide 6 clinically useful information on patient pulse rate, 7 oxygen saturation, respiration and movement. From a 8 combination of some or all of this information 9 clinical decisions can be made with regard to the 10 11 patient's health. The patient respiration information is used to monitor the patient in order 12 to compute a respiration rate and to detect 13 breathing abnormalities, for example: apnea events, 14 cessation in breathing, sharp intakes of breaths, 15 coughing, excessively fast breathing, excessively 16 17 slow breathing, etc. Information derived from one or more of the respiration, movement, oxygen saturation 18 and pulse measurements may be used to trigger an 19 alarm to call for medical help or to initiate an 20 automated process for the administration of a 21 therapeutic intervention. A method may be employed 22 23 for the archiving of the derived signals during the analysis period of the patient which may be used at 24 a later date for analysis by the clinician. 25 26 The device may be used to monitor the patient both 27 during sleep and when awake. 28 29 The device may be used to detect the onset of sudden 30 infant death syndrome SIDS by detecting and 31 analysing abnormalities in the measurement of one or 32

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1 more of the following: oxygen saturation, 2 respiration, movement and pulse. 3 4 4.3.3 Alarm As described above, it is envisaged that the 5 gathered information is used to trigger an alarm at 6 the bedside and/or at a remote nursing station. This 7 alarm would be graded according to a classification 8 of patient information. For example a reduction in 9 oxygen saturation below a predefined threshold with 10 associated loss or irregularity of patient movement, 11 irregularity of pulse rate and loss or irregularity 12 of patient respiration could trigger the highest 13 level of alarm, whereas a reduction of oxygen 14 saturation below a predefined threshold with a 15 normal level of patient movement and/or a regular 16

respiration pattern could trigger a lower level of

17

18

alarm.

- 1 5. Brief Description of Drawings
- Figure 1(a): A wavelet transform surface showing the
- 3 pulse band (located between the dashed lines). (High
- 4 to Low energy is graded from white to black in the
- 5 grey scale plot.)
- 6 Figure 1(b): Three-dimensional view of the wavelet
- 7 transform surface of figure 1(a) showing the maxima
- 8 of the pulse band with respect to frequency (the
- 9 ridge) superimposed as a black path across the band
- 10 maxima. (High to Low energy is graded from white to
- 11 black in the grey scale plot.)
- 12 Figure 2: 3-D Schematic of a wavelet transform
- 13 surface containing two bands. The locus of the local
- maxima on the bands (the 'ridges') are shown by
- 15 dashed lines.
- 16 Figure 3: Schematics of the RAP (top left) and RFP
- 17 (top right) signals derived from ridge A in figure 1
- 18 together with their corresponding wavelet transforms
- 19 shown below each (in 2D).
- 20 Figure 4(a): The SWFD method as applied to a pulse
- 21 oximeter signal Scalogram of Original Signal. .
- 22 (High to Low energy is graded from white to black in
- 23 the grey scale plot.)
- Figure 4 (b): The SWFD method as applied to a pulse
- 25 oximeter signal 3-D view of scalogram in (a) with
- the path of the pulse band ridge superimposed. .
- 27 (High to Low energy is graded from white to black in
- 28 the grey scale plot.)
- Figure 4 (c): The SWFD method as applied to a pulse
- 30 oximeter signal RAP signal (Top: full signal.
- 31 Lower: blow up of selected region)

- 1 Figure 4 (d): The SWFD method as applied to a pulse
- 2 oximeter signal RFP signal (Top: full signal.
- 3 Lower: blow up of selected region)
- 4 Figure 5(a): The SWFD method as applied to a pulse
- 5 oximeter signal RAP scalogram. (High to Low energy
- is graded from white to black in the grey scale
- 7 plot.)
- Figure 5(b): The SWFD method as applied to a pulse
- 9 oximeter signal RFP scalogram. (High to Low energy
- is graded from white to black in the grey scale
- 11 plot.)
- 12 Figure 5(c): The SWFD method as applied to a pulse
- oximeter signal 3-D view of RAP scalogram with
- 14 breathing band ridge shown. (High to Low energy is
- graded from white to black in the grey scale plot.)
- 16 Figure 5(d): The SWFD method as applied to a pulse
- oximeter signal 3-D view of RFP scalogram with
- 18 ridge shown. (High to Low energy is graded from
- 19 white to black in the grey scale plot.)
- 20 Figure 6(a): PPG Signal
- 21 Figure 6(b): Pulse band and ridge corresponding to
- 22 signal (a) . (High to Low energy is graded from
- white to black in the grey scale plot.)
- 24 Figure 6(c): RAP signal derived from ridge in (b)
- with breathing switch (square waveform)
- 26 superimposed.
- 27 Figure 6(d): RFP signal derived from ridge in (b)
- 28 Figure 7(a): Wavelet Transform of RAP signal. (High
- 29 to Low energy is graded from white to black in the
- 30 grey scale plot.)

- 1 Figure 7(b): Extracted ridges from wavelet transform
- 2 in (a). (High to Low energy is graded from white to
- 3 black in the grey scale plot.)
- 4 Figure 7(c): Wavelet Transform of RFP signal. (High
- 5 to Low energy is graded from white to black in the
- 6 grey scale plot.)
- 7 Figure 7(d): Extracted ridges from wavelet transform
- 8 in (c). (High to Low energy is graded from white to
- 9 black in the grey scale plot.)
- 10 Figure 8(a): Breathing ridges extracted from the
- 11 original wavelet transform
- 12 Figure 8(b): Breathing ridges extracted from the
- 13 secondary wavelet transform of the RAP signal
- 14 Figure 8(c): Breathing ridges extracted from the
- 15 secondary wavelet transform of the RFP signal
- 16 Figure 8(d): Selected respiration path (SRP).
- 17 Figure 9: Transform Phase along the SRP
- 18 Figure 10: Filling in missing segments of the SRP
- 19 Figure 11: Wavelet Representations of the Red PPG
- 20 (top) and Infrared PPG (bottom)
- 21 Figure 12: Schematic of the Sliding Window used to
- Obtain the Wavelet Components for the 3-D Lissajous
- Figure 13(a): Wavelet-based 3-D Lissajous: 3-D View.
- Figure 13(b): Wavelet-based 3-D Lissajous: End on
- 25 View of (a).
- Figure 13(c): Wavelet-based 3-D Lissajous: End on
- 27 View of Selected Component.
- 28 Figure 14: Standard Deviation of Lissajous
- 29 Components in Figure 3. Top plot: SD of principle
- 30 component; Middle plot: SD of minor component; Lower
- 31 plot: Ratio of SD components. All three plots
- 32 plotted against frequency in Hz.

- 1 Figure 15: Computed Oxygen Saturation Curves. Dotted
- 2 line: Signal Amplitude Method; Dashed Line
- 3 traditional Signal Lissajous Method; Solid Line:
- 4 Wavelet-based 3-D Lissajous Method.
- 5 Figure 16: The red and infrared wavelet modulus
- 6 surfaces corresponding to a 45 second segment of PPG
- 7 signals . (High to Low energy is graded from white
- 8 to black in the grey scale plot.)
- 9 Figure 17: The wavelet ratio surface derived from
- 10 the division of the red by the infrared wavelet
- 11 representations shown in Figure 16.
- 12 Figure 18: An end view of the wavelet ratio surface
- 13 shown in figure 17.
- 14 Figure 19: Computed Oxygen Saturation curves. Dotted
- 15 line: Oxygen Saturation from Traditional Signal
- 16 Amplitude Method; Dashed Line: Oxygen Saturation
- 17 from Traditional Signal Lissajous Method; Solid
- 18 Line: Oxygen Saturation from Traditional Wavelet-
- 19 Ratio Surface Method
- 20 Figure 20(a): Wavelet transform plot of a PPG signal
- 21 taken from a young baby showing a corresponding to
- 22 patient movement. Low to high energy is depicted
- 23 from black to white in the greyscale plot.
- Figure 20(b): Three-dimensional view of (a). Low to
- 25 high energy is depicted from black to white in the
- 26 greyscale plot.
- Figure 21(a): Transform plot of figure 20(a) with
- 28 modulus maxima superimposed. Low to high energy is
- depicted from black to white in the greyscale plot.
- 30 Figure 21(b): Three-dimensional view of figure
- 31 21(a). Low to high energy is depicted from black to
- 32 white in the greyscale plot.

- 1 Figure 22(a): End view of modulus maxima lines in
- 2 figure 21(b).
- 3 Figure 22(b): Amplitude threshold method of
- 4 identifying modulus maxima associated with movement
- 5 artefact
- 6 Figure 22(c): Pulse ridge-based method of
- 7 identifying modulus maxima associated with movement
- 8 artefact
- 9 Figure 22(d): Respiration ridge-based method of
- 10 identifying modulus maxima associated with movement
- 11 artefact
- 12 Figure 23: Block diagram of device configuration
- 13 Figure 24: Block diagram of subcomponents of oxygen
- 14 saturation component (16) shown in figure 23
- 15 Figure 25: Block diagram of subcomponents of
- 16 respiration component (18) shown in figure 23
- 17 Figure 26: Block diagram of subcomponents of
- 18 movement component (17) shown in figure 23
- 19 Figure 27(a): Schematic of foot cuff mounting: soft
- 20 housing surrounding foot used to hold monitoring
- 21 apparatus.
- 22 80 patient leg; 81 patient heel; 82 patient toes; 83
- 23 soft housing surrounding foot
- 24 Figure 27(b): View from both sides of the envisaged
- device: preferred embodiment for neonatal monitor.
- 26 84 connection cabling; 85 RF components attached to
- 27 housing; 86 LEDs; 87 pulse oximeter components
- 28 attached to housing; 88 photodetector. (Note LEDs
- and photodetector may also be located on toe using
- 30 short cable length from cuff.)

1	Figure 28: Schematic of wrist cuff mounting: 90				
2	electronic component housing; 91 wrist band; 92				
3	connector cable; 93 finger probe				
4					
5	6. General				
6	The invention has been described and shown with				
7	specific reference to specific embodiments. However				
8	it will be understood by those skilled in the art				
9	that changes to the form and details of the				
10	disclosed embodiments may be made without departing				
11	from the spirit and scope of the invention. For				
12	example signal transforms other than the wavelet				
13	transform may be used. Other variations may include				
14	using a multiplexed arrangement which alternates				
15	measurements for pulse, oxygen saturation,				
16	respiration and movement artefact using variations				
17	of the acquisition equipment and transmission				
18	electronics. These variations may include but are				
19	not limited to the use of more than two wavelengths				
20	of light and variable power and/or variable duty				
21	cycle to the light transmitters.				
22					
23	7. Reference				
24	Addison P.S., 'The Illustrated Wavelet Transform				
25	Handbook', Institute of Physics Publishing, 2002,				
26	Bristol, UK.				
27					

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1	CLAIMS				
2					
3	1. A method of measuring physiological parameters,				
4	comprising:				
5	using a signal acquisition means to obtain a				
6	<pre>pulse oximetry signal;</pre>				
7	decomposing the pulse oximetry signal by				
8	wavelet transform analysis;				
9	identifying a primary band and a secondary band				
1.0	on a transform surface constructed by the wavelet				
11	transform analysis; and				
12	interpreting the secondary band to reveal				
13	information pertaining to the physiological				
14	parameters causing the primary band.				
15					
16	2. The method of claim 1, wherein the step of				
17	deriving information from the secondary band				
18	comprises:				
19	defining a selected path along the secondary				
20	band;				
21	extracting a time-frequency locus of points on				
22	the selected path;				
23	extracting a time-amplitude locus of points on				
24	the selected path; and				
25	decomposing the time-frequency and time-				
26	amplitude loci by wavelet transform analysis.				
27					
28	3. The method of claim 1 or claim 2, wherein the				
29	selected path is in the vicinity of a ridge.				
30					
31	4. The method of claim 3, wherein the selected				

32 path is a ridge.

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1						
2	5. The method of any preceding claim, wherein the					
3	pulse oximetry signal is a photoplethysmogram (PPG).					
4						
5	6. The method of any preceding claim, wherein the					
6	wavelet transform analysis employs a continuous					
7	wavelet transform.					
8						
9	7. The method of any preceding claim, further					
LO	comprising the steps of;					
L1	defining a second selected path along the					
L2	primary band;					
L3	deriving primary band information from the					
14	second selected path;					
15	comparing the primary band information with the					
16	information derived from the secondary band; and					
17	choosing the optimum data set from the					
18	information, said optimum data set being the data					
19	set that most accurately represents the					
20	physiological parameters causing the primary band.					
21						
22	8. The method of claim 7 when dependent from any					
23	of claims 2-6, wherein the optimum data set is					
24	chosen from the group comprising information derived					
25	from the primary band, information derived from the					
26	decomposed time-frequency locus, and the information					
27	derived from the time-amplitude locus.					
28						
29	9. The method of any preceding claim, wherein the					

primary band is a breathing band and the secondary band is a pulse band.

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The method of any preceding claim, in which the 1 derived information is patient respiration 2 information. 3 4 The method of claim 10, wherein the information 5 is used to determine the respiration rate of a 6 7 patient. 8 The method of claim 10 or claim 11, wherein the 9 12. information is used to identify individual breaths 10 11 of the patient. 12 The method of any of claims 10-12, used to 13 reveal breathing irregularities. 14 15 A physiological measurement system comprising: 16 14. a signal acquisition means which includes a 17 light emitting device and a photodetector attachable 18 to a subject to obtain a pulse oximetry signal; 19 analogue to digital converter means arranged to 20 convert said pulse oximetry signal into a digital 21 pulse oximetry signal; 22 signal processing means suitable to receive 23 said digital pulse oximetry signal and arranged to 24 decompose that signal by wavelet transform means; 25 a respiration component arranged to identify a 26 primary band and a secondary band on a transform 27 surface constructed by the wavelet transform 28 analysis; and to interpret the secondary band to 29

reveal information pertaining to the physiological

parameters causing the primary band.

30

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1 15. The system of claim 14, in which the signal

- 2 processing means and respiration component are
- arranged to process the pulse oximetry signal by the

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4 method of any of claims 2-13.

5

- 6 16. The system of claim 14 or claim 15, further
- 7 comprising an analyser component arranged to collect
- 8 information from the respiration component, and a
- 9 device output arranged in communication with the
- 10 analyser component.

11

- 12 17. The system of claim 16 wherein the analyser
- component is arranged to generate an alarm signal
- upon detection of a predetermined set of conditions.

1.5

- 16 18. The system of claim 17, wherein the
- 17 predetermined set of conditions includes the
- 18 existence of a respiration artefact for at least a
- 19 given time.

20

- 21 19. The system of any of claims 16-18, wherein the
- 22 device output comprises visual display means
- operable to display the pulse oximetry signal and
- information derived therefrom in real time; and
- 25 alarm means operable to receive the alarm signal
- from the analyser component and to generate an
- 27 alarm.

- 29 20. A method of measuring physiological parameters,
- 30 comprising:
- using a signal acquisition means to obtain a
- red pulse oximetry signal in the red light spectrum

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and an infra-red pulse oximetry signal in the infra-1 2 red spectrum; decomposing each pulse oximetry signal by 3 wavelet transform analysis; and 4 combining the decomposed signals to obtain a 5 measure of a physiological parameter. 6 7 The method of claim 20, wherein the step of 21. 8 combining the decomposed signals to obtain a measure 9 of a physiological parameter comprises; 10 for each decomposed signal, selecting a 11 plurality of frequencies and a first set time 12 period; and 13 plotting the transform values over the set time 14 period at each frequency of the decomposed red pulse 15 oximetry signal against those of the decomposed 16 infra-red pulse oximetry signal as a plurality of 17 Lissajous figures. 18 19 The method of claim 21, wherein the Lissajous 20 21 figures are three-dimensional. 22 The method of claim 21 or claim 22, further 23 24 comprising plotting Lissajous figures for the selected frequencies over a second set time period, 25 the start of which is later than the start of the 26 first set time period. 27 28 The method of claim 23, wherein a plurality of 29 24. successive later time periods are selected such that 30

a frequency window moves across the time-frequency

plane of each decomposed signal.

31

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1					
2	25. The method of any of claims 21-24 further				
3	comprising the steps of analysing a characteristic				
4	parameter of each Lissajous figure; using the value				
5	of the characteristic parameter to select a				
6	Lissajous figure from the plurality of Lissajous				
7	figures; and				
8	using the selected Lissajous figure to				
9	determine a physiological parameter.				
10					
11	26. The method of claim 25, wherein the				
12	characteristic parameter of a Lissajous figure is				
13	derived from its spread along its principle				
14	components, and wherein the selected Lissajous				
15	figure is the one having the maximum spread.				
16					
17	27. The method of claim 26, wherein the spread of a				
18	Lissajous figure is represented by the standard				
19	deviation along the principle component or				
20	components.				
21					
22	28. The method of claim 27, wherein the ratio of				
23	standard deviations along two orthogonal principle				
24	components is calculated to give a further aid to				
25	the selection of the Lissajous figure for use in				
26	determining the physiological parameter.				
27					
28	29. The method of any of claims 21-28, wherein the				
20	glone of the gelected Lissaious figure is used as a				

slope of the selected Lissajous figure is used as a measure of the patient oxygen saturation.

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1	30. The method of claim 29, wherein a predefined					
2	look-up table is used to deduce the relationship					
3	between the slope and the oxygen saturation.					
4						
5	31. The method of claim 20, wherein the step of					
6	combining the decomposed signals to obtain a measure					
7	of a physiological parameter comprises the steps of:					
8	constructing a wavelet ratio surface based					
9	on the ratio of one of the decomposed signals to the					
10	other of the decomposed signals; and					
11	deriving one or more physiological parameters					
12	from the wavelet ratio surface.					
13						
14	32. The method of claim 31, wherein the step of					
15	deriving one or more physiological parameters from					
16	the wavelet ratio surface comprises the steps of:					
17	selecting a region of the wavelet ratio surface					
18	for use in determining the physiological parameter.					
19						
20	33. The method of claim 32, wherein the selected					
21	region is derived from a path in the wavelet ratio					
22	surface formed from a collection of points in the					
23	vicinity of the maxima of a pulse band.					
24						
25	34. The method of claim 33, wherein the selected					
26	region is derived from a path in the wavelet ratio					
27	surface formed from a collection of points					
28	corresponding to the maxima of a pulse band.					
29						
30	35. The method of claim 33 or claim 34, further					
31	comprising the step of inspecting a predefined look-					

1 up table to determine the correlation between the

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2 path and the oxygen saturation.

3

4 36. The method of any of claims 31-35, further

5 comprising the step of, when a local disturbance

occurs in the pulse band maxima, holding the

7 previously determined oxygen saturation value for a

8 specified duration or until the local disturbance

9 ceases.

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10

11 37. The method of any of claims 20-36, wherein the

pulse oximetry signal is a photoplethysmogram (PPG)

13 signal.

14

15 38. The method of any of claims 20-37, wherein the

16 wavelet transform analysis employs a continuous

17 wavelet transform.

18

19 39. A physiological measurement system comprising:

a signal acquisition means which includes a

21 light emitting device and a photodetector attachable

to a subject to obtain a red pulse oximetry signal

in the red light spectrum and an infra-red pulse

24 oximetry signal in the infra-red spectrum;

analogue to digital converter means arranged to

26 convert said pulse oximetry signals into digital

27 pulse oximetry signals;

28 signal processing means suitable to receive

29 said digital pulse oximetry signals and arranged to

decompose those signals by wavelet transform means;

31 and

	63				
1	an oxygen saturation component arranged to				
2	combine the decomposed signals to obtain a measure				
3	of a physiological parameter.				
4					
5	40. The system of claim 39, wherein the oxygen				
6	saturation component comprises a Lissajous				
7	computation unit arranged to receive the decomposed				
8	signals from the signal processing means, and, for				
9	each decomposed signal, to:				
10	select a plurality of frequencies and a first				
11	set time period; and				
12	plot the transform values over the first set				
13	time period at each frequency of the decomposed red				
14	pulse oximetry signal against those of the				
15	decomposed infra-red pulse oximetry signal as a				
16	plurality of Lissajous figures.				
17					
18	41. The system of claim 40, in which the signal				
19	processing means and the Lissajous computation unit				
20	are arranged to perform the method of any of claims				
21	21-30.				
22					
23	42. The system of claim 39, wherein the oxygen				
24	saturation component comprises a pulse ridge				
25	computational unit arranged to:				
26	receive the decomposed signals from the signal				
27	processing means;				
28	to construct a wavelet ratio surface based on				
29	the ratio of one of the decomposed signals to the				
30	other of the decomposed signals; and				
31	to derive one or more physiological parameters				

32 from the wavelet ratio surface.

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64 1 2 The system of claim 42, in which the signal processing means and the pulse ridge computational 3 unit are arranged to perform the method of any of 4 claims 31-38. 5 6 The system of claim 39, wherein the oxygen 7 saturation unit comprises the Lissajous computation 8 unit of claim 40 and the pulse ridge computational 9 unit of claim 42, said system further comprising a 10 signal sorting component arranged to receive a 11 signal from the signal processing means and to 12 allocate that signal to both the Lissajous 13 computation unit and the pulse ridge computational 14 15 unit. 16 The system of claim 44, further comprising 17 comparison means arranged to receive a signal from 18 each of the Lissajous computation unit and the pulse 19 ridge computational unit and to select a signal to 20 be representative of the physiological parameter. 21 22 The system of claim 45, in which the signal 23 24 processing means and the oxygen saturation component are arranged to perform the method of any of claims 25 20-38. 26 27 The system of any of claims 39-46, further 28 comprising an analyser component arranged to collect

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comprising an analyser component arranged to collect information from the respiration component, and a device output arranged in communication with the analyser component.

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2 48. The system of claim 47 wherein the analyser

3 component is arranged to generate an alarm signal

4 upon detection of a predetermined set of conditions.

5

- 6 49. The system of claim 48, wherein the
- 7 predetermined set of conditions includes the
- 8 existence of an abnormal oxygen saturation for at
- 9 least a given time.

10

- 11 50. The system of any of claims 47-49, wherein the
- 12 device output comprises visual display means
- operable to display the pulse oximetry signal and
- information derived therefrom in real time; and
- 15 alarm means operable to receive the alarm signal
- 16 from the analyser component and to generate an
- 17 alarm.

18

- 19 51. A method of measuring a movement artefact
- 20 occurring in a pulse oximetry signal, comprising:
- using a signal acquisition means to obtain a
- 22 pulse oximetry signal;
- 23 decomposing the pulse oximetry signal by
- 24 wavelet transform analysis;
- finding the modulus maxima of a transform
- 26 surface constructed by the wavelet transform
- 27 analysis;
- determining a set of modulus maxima that lie
- 29 within a predetermined amplitude threshold; and
- 30 associating this set of modulus maxima with the
- 31 movement artefact.

66

The method of claim 51, wherein the 1 predetermined threshold is defined as a given 2 quantitative excursion of the modulus maxima from a 3 localised mean value of a pulse ridge, said 4 excursion being encountered as the ridge is 5 followed. 6 7 The method of claim 51, wherein the 8 predetermined threshold is defined as a given 9 quantitative excursion of the modulus maxima from a 10 11 localised mean value of a breathing ridge, said excursion being encountered as the ridge is 12 followed. 13 14 The method of any of claims 51-53, wherein the 15 amplitude threshold is defined over a specific 16 frequency range. 17 18 55. A movement measurement system comprising: 19 a signal acquisition means which includes a 20 light emitting device and a photodetector attachable 21 to a subject to obtain pulse oximetry signal; 22 analogue to digital converter means arranged to 23 convert said pulse oximetry signals into digital 24 pulse oximetry signals; 25 signal processing means suitable to receive 26 said digital pulse oximetry signals and arranged to 27 decompose those signals by wavelet transform means; 28 and 29 movement measurement means arranged to receive 30 the decomposed signal; find the modulus maxima of a 31 transform surface constructed by the wavelet 32

67

transform analysis; determine a set of modulus
maxima that lie within a predetermined amplitude

3 threshold; and associate this set of modulus maxima

4 with a movement artefact in the decomposed signal.

5

6 56. The system of claim 55, wherein the movement

7 measurement means further comprises pulse check

8 means arranged to determine if the set of modulus

9 maxima that lie within the predetermined amplitude

threshold represents an excursion from a pulse band.

11

12 57. The system of claim 55 or claim 56, wherein the

13 movement measurement means further comprises

14 respiration check means arranged to determine if the

15 set of modulus maxima that lie within the

16 predetermined amplitude threshold represents an

17 excursion from a breathing band.

18

19 58. The system of any of claims 55-57, further

20 comprising an analyser component arranged to collect

21 information from the movement measurement means, and

22 a device output arranged in communication with the

23 analyser component.

24

25 59. The system of claim 58 wherein the analyser

26 component is arranged to generate an alarm signal

27 upon detection of a predetermined set of conditions.

28

29 60. The system of claim 59, wherein the

30 predetermined set of conditions includes the

31 existence of excessive or erratic movement, or the

lack of movement, for at least a given time.

68

1 61. The system of any of claims 58-60, wherein the 2 device output comprises visual display means 3 operable to display the pulse oximetry signal and 4 information derived therefrom in real time; and 5 alarm means operable to receive the alarm signal 6 7 from the analyser component and to generate an 8 alarm. 9 A method of measuring physiological parameters, 10 comprising: 11 using a signal acquisition means to obtain a 12 pulse oximetry signal; 13 decomposing the pulse oximetry signal by 14 wavelet transform analysis; and 15 identifying a dominant band in a transform 16 17 surface constructed by the wavelet transform analysis, the band being associated with pulse 18 19 components in the pulse oximetry signal. 20 The method of claim 62, further comprising the 21 step of deriving a selected time-frequency path 22 along the dominant band. 23 24 The method of claim 63, wherein the selected 25 time-frequency path is the ridge of the band. 26 27 The method of any of claims 62-64, wherein the 28 65. measured physiological parameter is a pulse rate. 29

32 comprising the step of using transform information

The method of any of claims 63-65, further

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66.

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along the selected path to provide information on 1 2 the times between individual pulses. 3 4 67. The method of claim 66, further comprising the 5 step of using transform information along the 6 selected path to provide information on the timings 7 of features within the pulse. 8 9 68. The method of claim 67, further comprising the 10 step of using the information on the timings of features within the pulse as a measure of arterial 11 compliance. 12 13 The method of any of claims 62-68, wherein the 14 pulse oximetry signal is a photoplethysmogram (PPG). 15 16 A physiological measurement system comprising: 17 a signal acquisition means which includes a 18 light emitting device and a photodetector attachable 19 to a subject to obtain a pulse oximetry signal; 20 analogue to digital converter means arranged to 21 convert said pulse oximetry signal into a digital 22 23 pulse oximetry signal; 24 signal processing means suitable to receive said digital pulse oximetry signal and arranged to 25 decompose that signal by wavelet transform means; 26 27 and 28 a pulse component arranged to identify a dominant band in a transform surface constructed by 29 30 the wavelet transform analysis, said band being

associated with pulse components in the pulse

3132

oximetry signal.

70

1

- 2 71. The system of claim 70, arranged to process the
- 3 pulse oximetry signal by the method of any of claims
- 4 58-65.

5

- 6 72. The system of claim 70 or claim 71, further
- 7 comprising an analyser component arranged to collect
- 8 information from the pulse component, and a device
- 9 output arranged in communication with the analyser
- 10 component.

11

- 12 73. The system of claim 72 wherein the analyser
- component is arranged to generate an alarm signal
- upon detection of a predetermined set of conditions.

15

- 16 74. The system of claim 73, wherein the
- 17 predetermined set of conditions includes the
- 18 existence of a pulse abnormality for at least a
- 19 given time.

20

- 21 75. The system of any of claims 72-74, wherein the
- 22 device output comprises visual display means
- operable to display the pulse oximetry signal and
- information derived therefrom in real time; and
- 25 alarm means operable to receive the alarm signal
- from the analyser component and to generate an
- 27 alarm.

- 29 76. A physiological measurement system comprising:
- a signal acquisition means which includes a
- 31 light emitting device and a photodetector attachable
- to a subject to obtain a pulse oximetry signal;

1 analogue to digital converter means arranged to convert said pulse oximetry signal into a digital 2 pulse oximetry signal; 3 signal processing means suitable to receive 4 said digital pulse oximetry signal and arranged to 5 6 decompose that signal by wavelet transform means; feature extraction means arranged to derive 7 8 physiological information from the decomposed 9 signal; 10 an analyser component arranged to collect 11 information from the feature extraction means; and data output means arranged in communication 12 with the analyser component. 13 14 The system of claim 76, wherein the feature 15 extraction means comprises one or more of: 16 17 (a) a respiration component arranged to 18 identify a primary band and a secondary band on a 19 transform surface constructed by the wavelet 20 transform analysis; and to interpret the secondary band to reveal information pertaining to the 21 physiological parameters causing the primary band; 22 23 (b) an oxygen saturation component arranged to combine the decomposed signals to obtain a measure 24 25 of a physiological parameter; (c) movement measurement means arranged to 26 receive the decomposed signal, find the modulus 27 maxima of a transform surface constructed by the 28 wavelet transform analysis; determine a set of 29 modulus maxima that lie within a predetermined 30 31 amplitude threshold; and associate this set of

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modulus maxima with a movement artefact in the 1 decomposed signal; and 2 (d) a pulse component arranged to identify a 3 dominant band in a transform surface constructed by 4 the wavelet transform analysis associated with pulse 5 components in the pulse oximetry signal. 6 7 The system of claim 77, wherein the respiration 8 78. component is arranged to process the pulse oximetry 9 signal by the method of any of claims 1-13. 10 11 79. The system of claim 77, wherein the oxygen 12 saturation component is arranged to process the 13 pulse oximetry signal by the method of any of claims 14 20-38. 15 16 The system of claim 77, wherein the movement 17 measurement means is arranged to process the pulse 18 oximetry signal by the method of any of claims 51-19 54. 20 21 The system of claim 80, wherein the output from 22 a respiration component is used as an input of the 23 movement measurement means. 24 25 The system of claim 77, wherein the pulse 82. 26 component is arranged to process the pulse oximetry 27 signal by the method of any of claims 62-69. 28 29 The system of any of claims 76-82, wherein the 30 83. analyser component is arranged to generate an alarm 31

signal upon detection of a predetermined set of

32

73

conditions exhibited by the information collected 1 from the feature extraction means. 2 3 The system of claim 83, further comprising 4 alarm means arranged to receive the alarm signal and 5 to generate an alarm. 6 7 The system of any of claims 76-84, wherein the 8 data output means comprises visual display means 9 operable to display the pulse oximetry signal and 10 information derived therefrom in real time. 11 12 The system of any of claims 76-85, wherein any 13 86. one or more of the analogue to digital converter 14 means, signal processing means, feature extraction 15 means, analyser component, and data output means are 16 arranged to receive and/or transmit signals via a 17 wireless communications link. 18 19 The system of any of claims 76-85, wherein the 20 87. signal acquisition means is arranged to transmit 21 signals via a wireless communications link. 22 23 The system of claim 87, wherein the signal 24 acquisition means is arranged to transmit signals to 25 the analogue to digital converter means via a 26 wireless communications link. 27 28

The system of any of claims 86-88, wherein the 29 analogue to digital converter means is arranged to 30 transmit signals to the signal processing means via 31

a wireless communications link. 32

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2 90. The system of any of claims 86-89, wherein the

3 analyser component is arranged to transmit signals

4 to the data output means via a wireless

5 communications link.

6

7 91. The system of any of claims 86-90, wherein the

8 wireless communications link is a radiofrequency

9 (RF) link.

10

11 92. The system of any of claims 85-91, wherein any

one or more of the signal acquisition means,

analogue to digital converter means, signal

14 processing means, feature extraction means, analyser

15 component, and data output means are embodied in a

16 form suitable to be worn on the body of a patient.

17

18 93. The system of claim 92, wherein any one or more

19 of the signal acquisition means, analogue to digital

20 converter means, signal processing means, feature

21 extraction means, analyser component, and data

output means are embodied in a cuff for attachment

23 to a patient's ankle.

24

25 94. The system of claim 92, wherein any one or more

of the signal acquisition means, analogue to digital

27 converter means, signal processing means, feature

28 extraction means, analyser component, and data

29 output means are embodied in a wrist band and

30 thimble for attachment to a patient's wrist and

31 finger respectively.

32

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95. The system of any of claims 76-94, embodied in a format suitable for home use.

The system of any of claims 77-95, where one or more of the respiration component, oxygen saturation

7 component are used as an apnea monitor.

8

6

9 97. The system of any of claims 77-95, where one or

component, movement measurement means and pulse

10 more of the respiration component, oxygen saturation

11 component, movement measurement means and pulse

component are used as a monitor for sudden infant

13 death syndrome.

14

18

98. A method of measuring a characteristic

parameter, comprising:

obtaining a signal which varies according to a

set of parameters including the characteristic

19 parameter;

decomposing the signal by wavelet transform

21 analysis;

identifying a primary band representative of

the characteristic parameter, and a secondary band

24 representative of a secondary parameter, on a

25 transform surface constructed by the wavelet

26 transform analysis; and

interpreting the secondary band to reveal

information pertaining to the parameters causing the

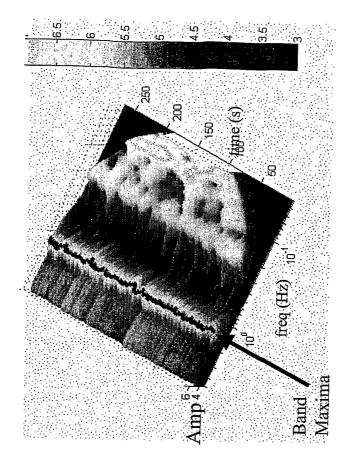
29 primary band.

30

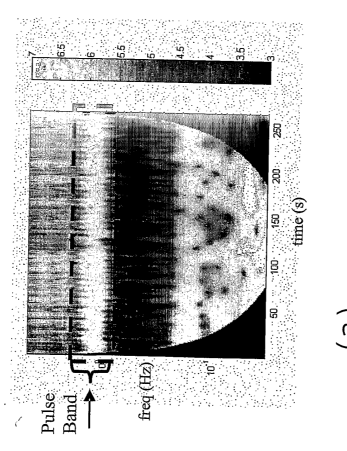
31 99. The method of claim 98, wherein the signal is

one of the group of signals comprising an

1	electrocardiogram, electroencephalogram,
т.	erectiocardrogram, erectioencepharogram,
2	electrogastrogram or electromyogram, heart rate
3	signals, pathological sounds, and ultrasound,
4	dynamic signals, non-destructive testing signals,
5	condition monitoring signals, fluid signals,
6	geophysical signals, astronomical signals,
7	electrical signals, financial signals including
8	financial indices, sound and speech signals,
9	chemical signals, and meteorological signals
10	including climate signals.
11	
12	100. The method of claim 97 or claim 98, as applied
13	in conjunction with any of claims 1-96.



Q



Figu

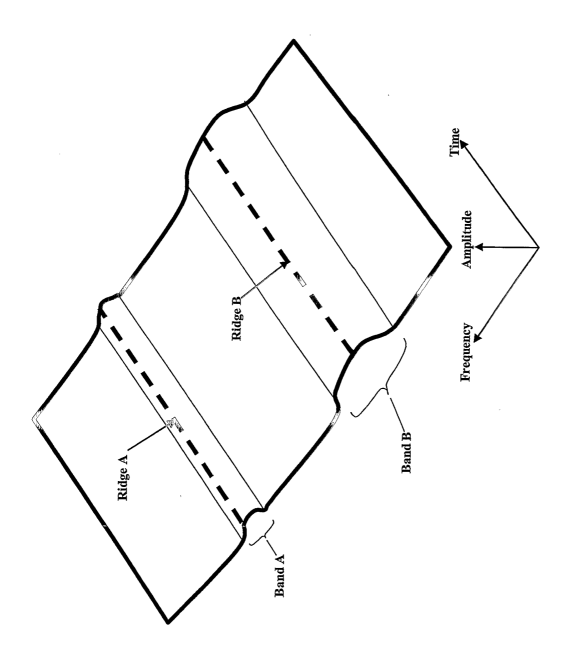


Figure 2

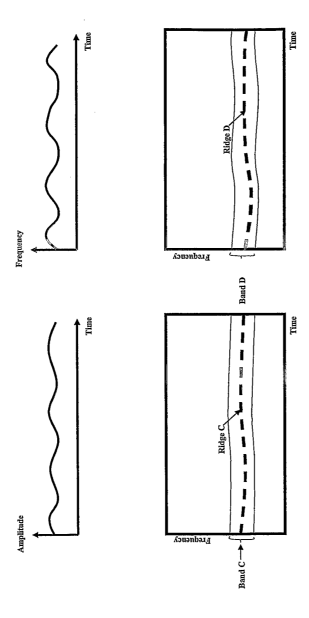


Figure 3

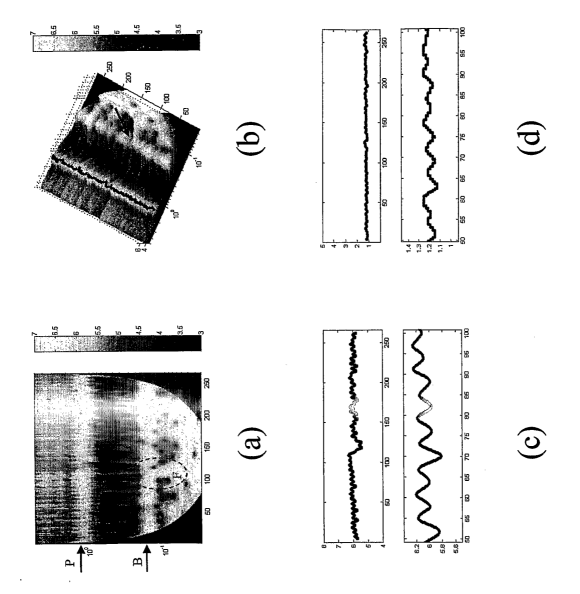
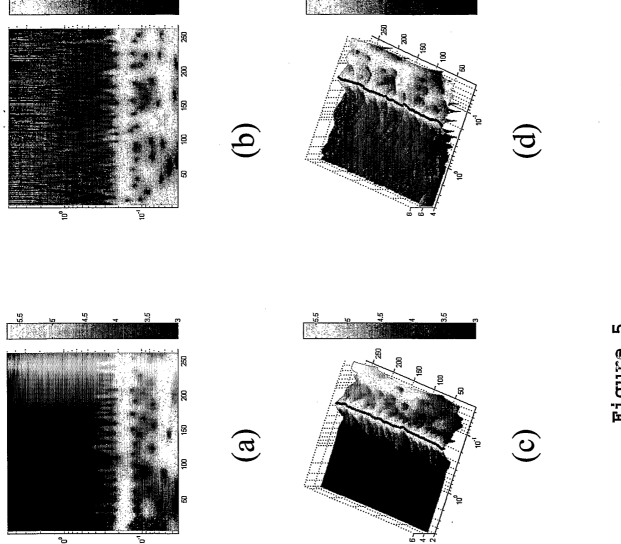
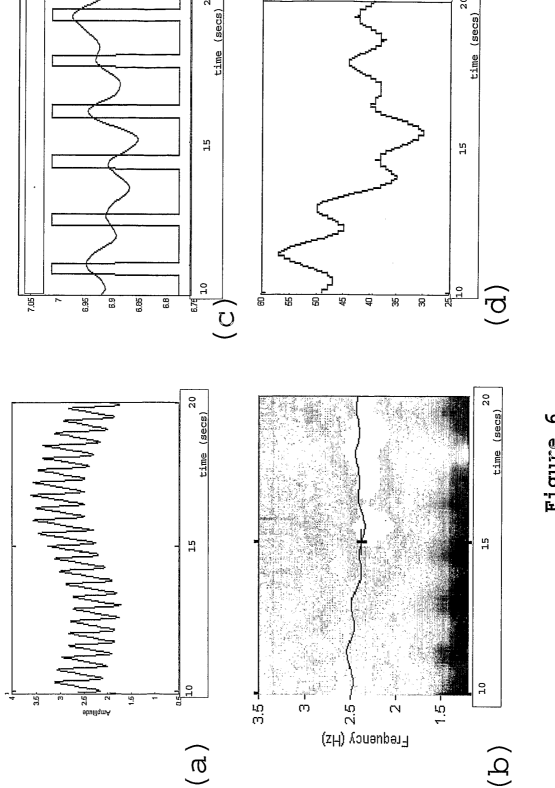
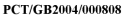
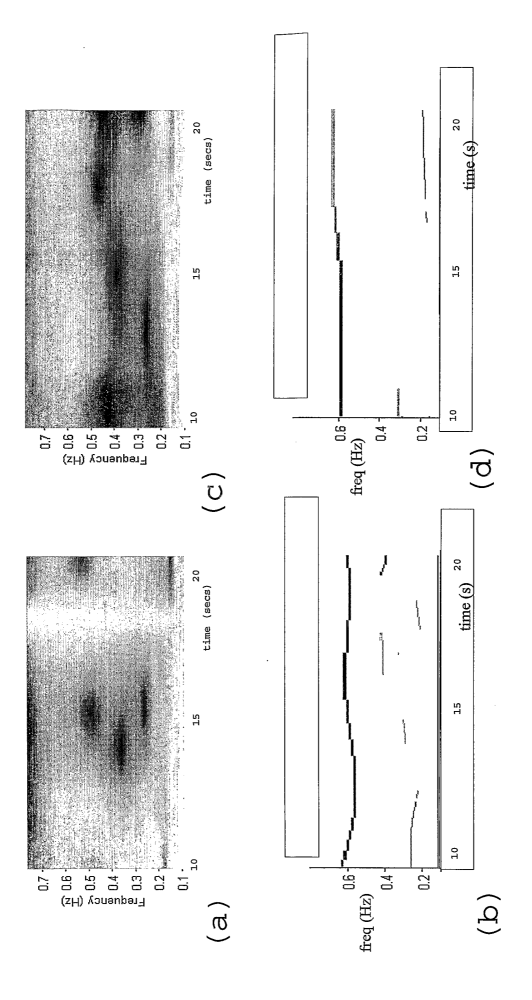


Figure 4

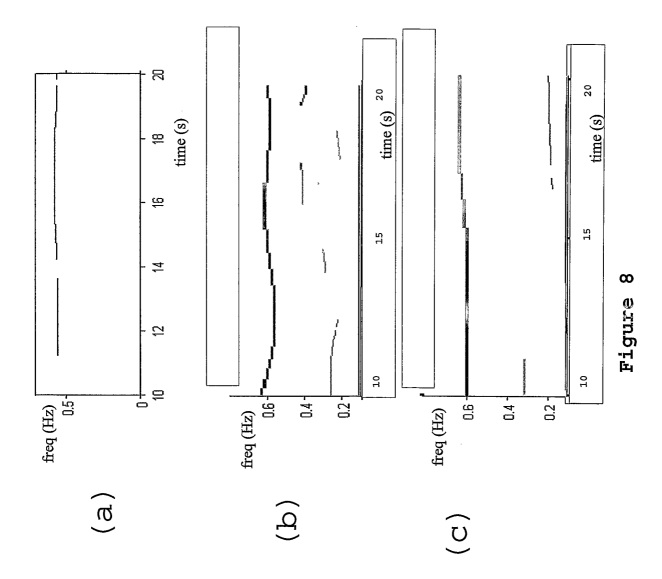








Figure



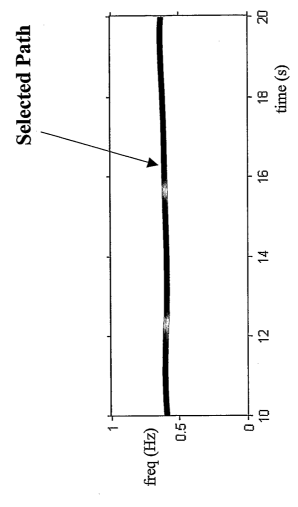


Figure 8 (continued)

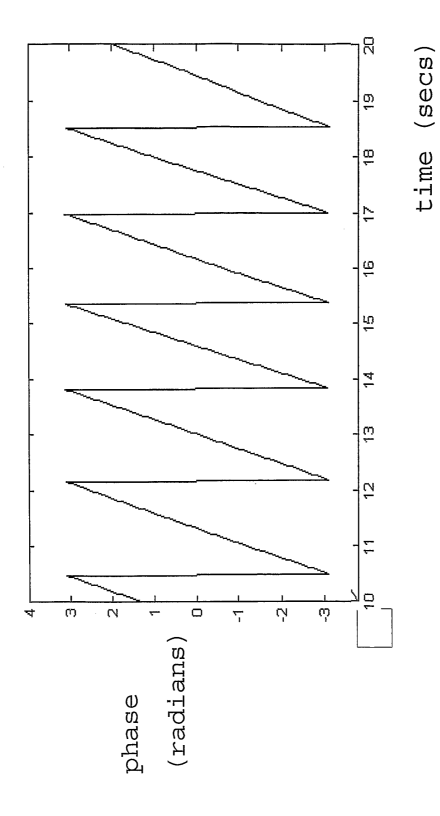


Figure 9

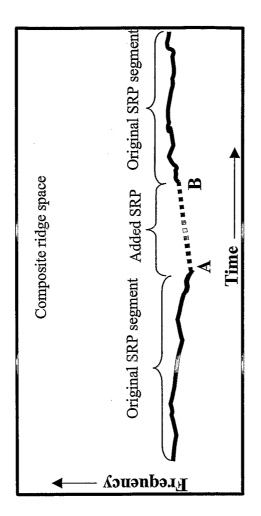


Figure 10

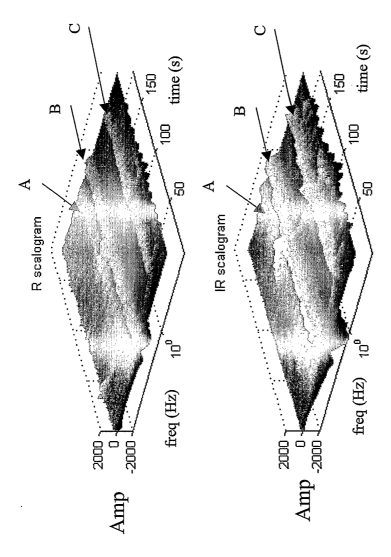


Figure 11

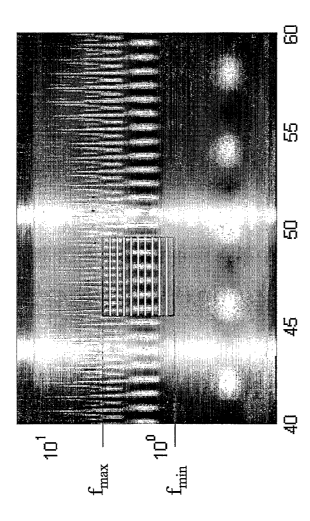
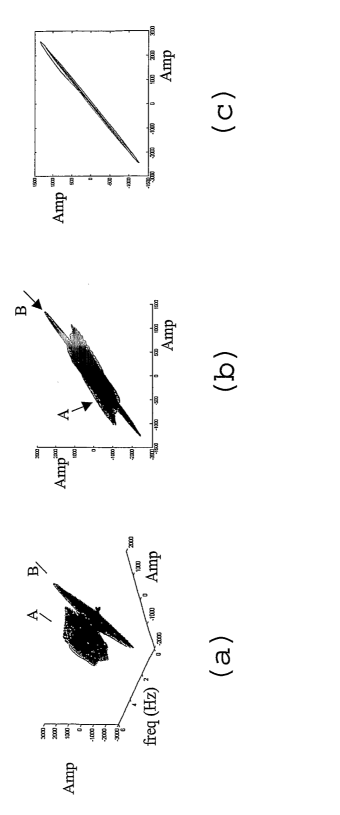


Figure 12



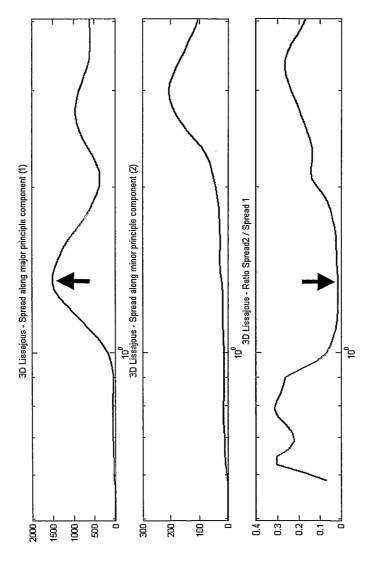


Figure 14

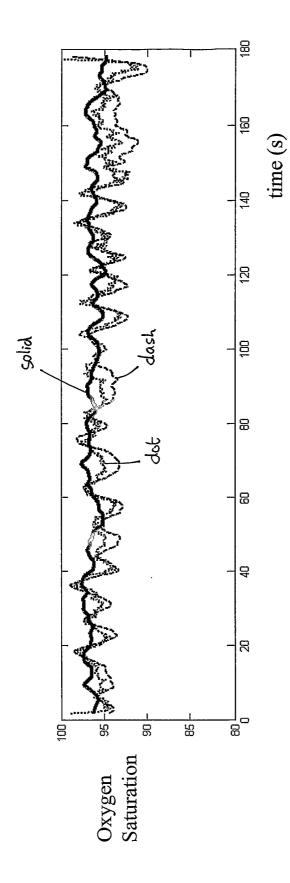


Figure 15

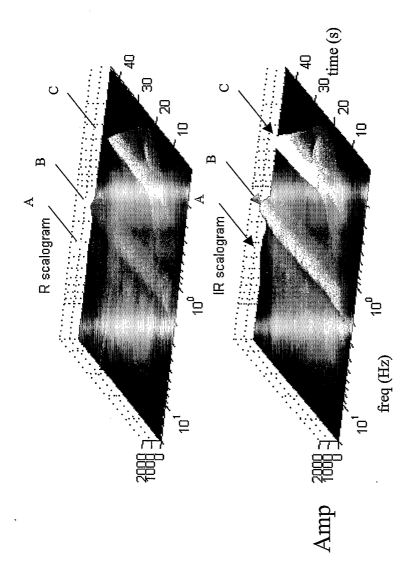


Figure 16

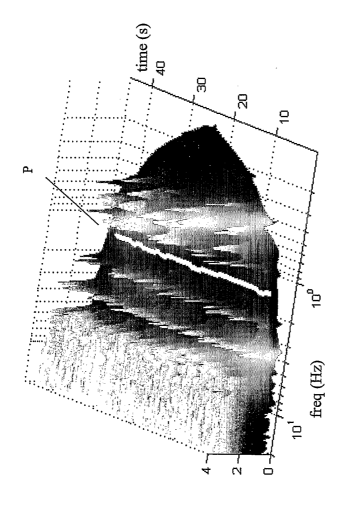
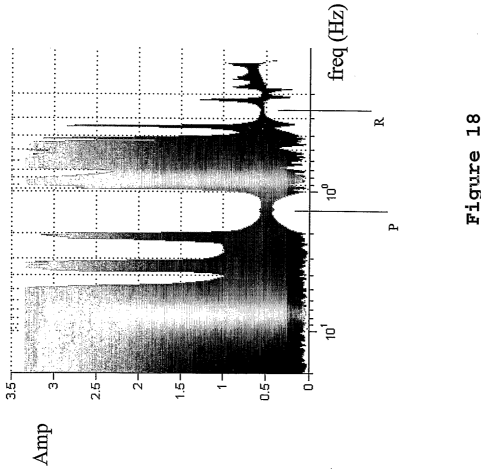


Figure 17



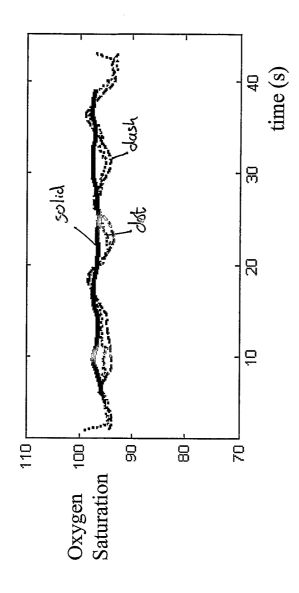
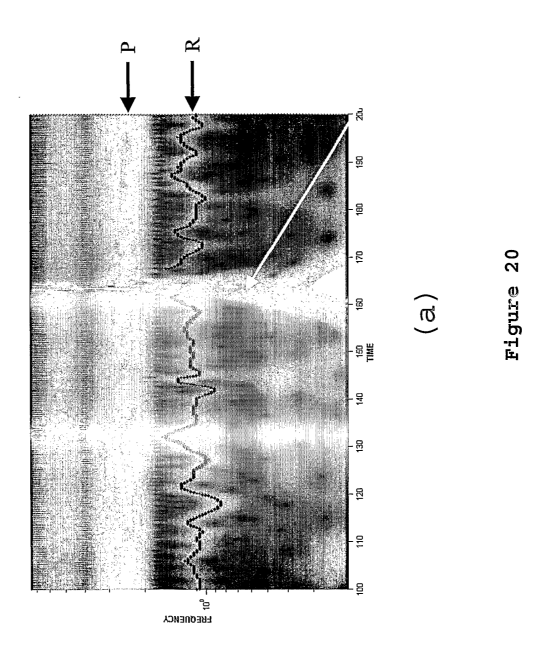


Figure 19

Movement artefact



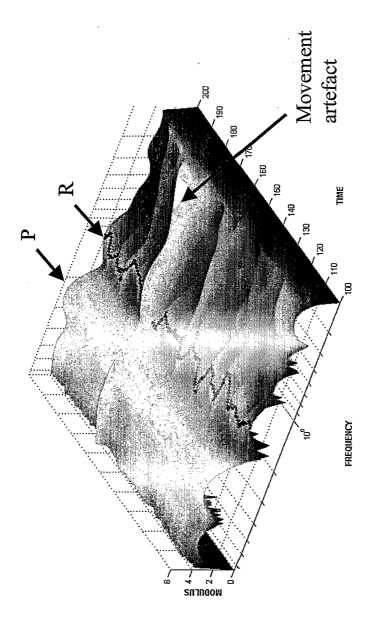
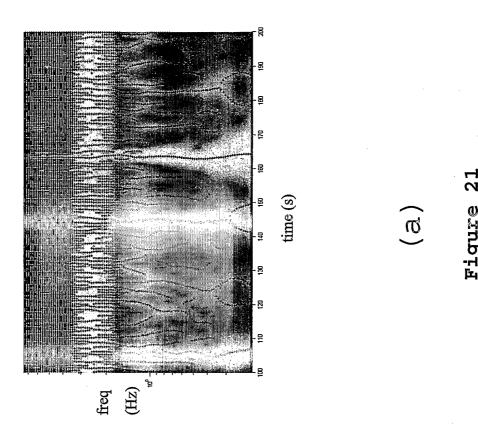


Figure 20 (continued)



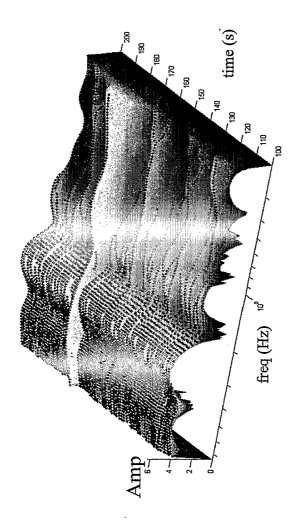


Figure 21 (continued)

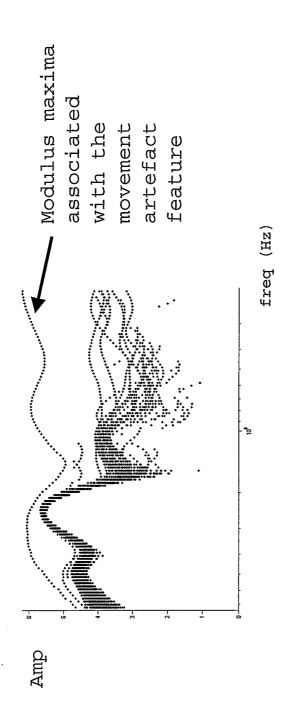
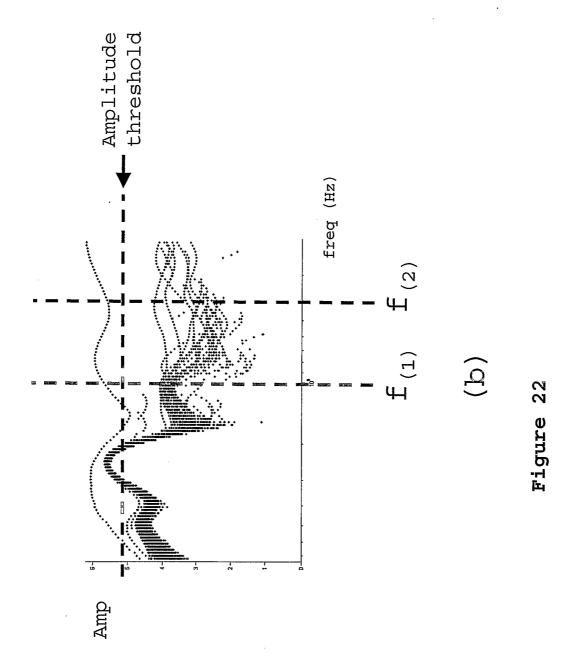
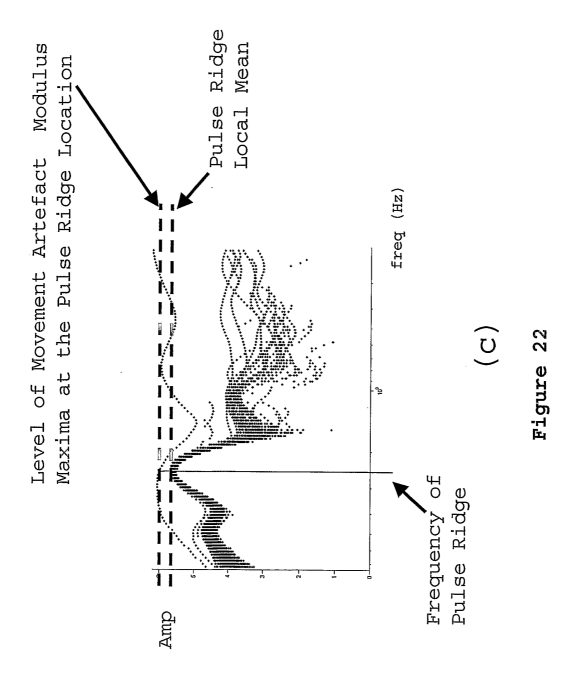
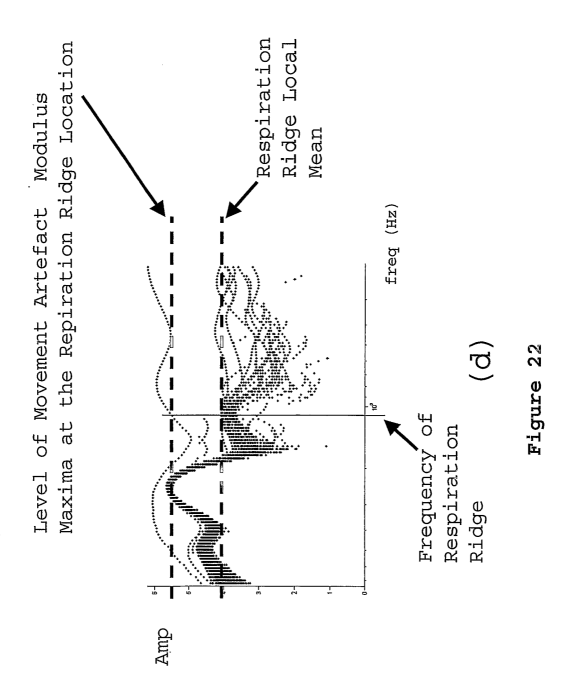


Figure 22







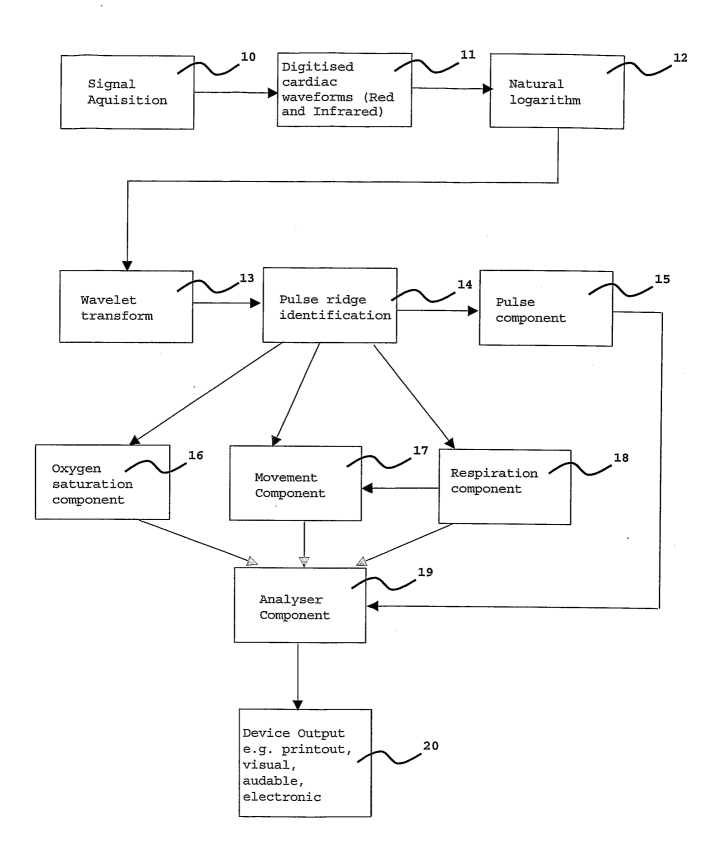


Figure 23

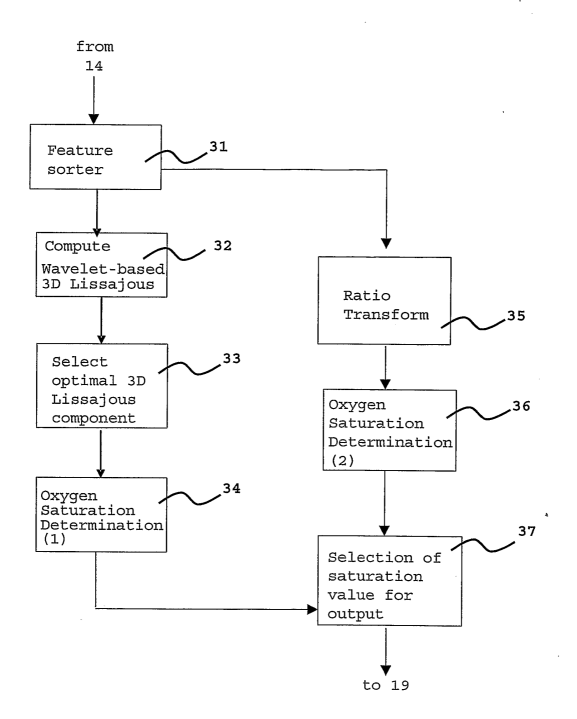


Figure 24

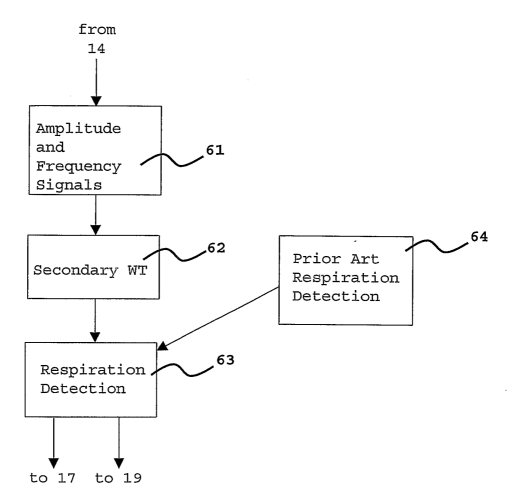


Figure 25

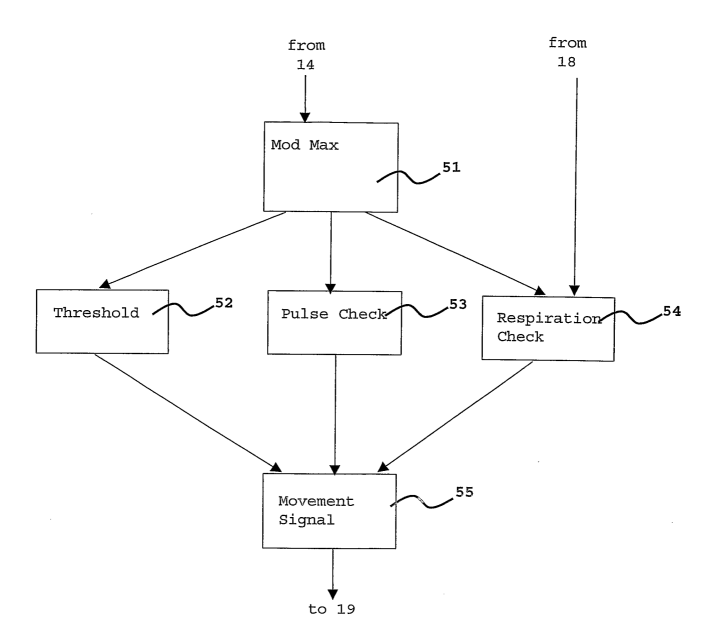
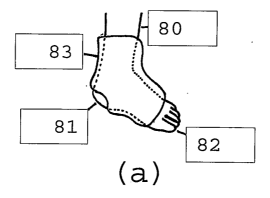


Figure 26



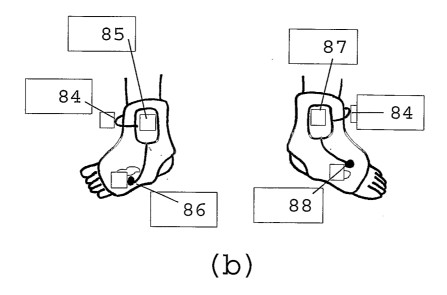


Figure 27

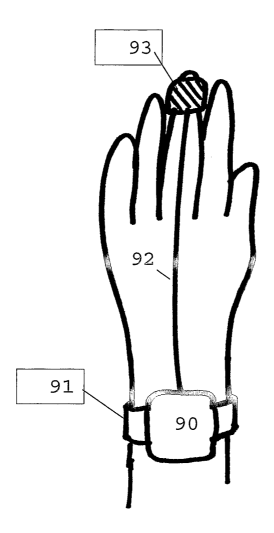


Figure 28