Note: Within nine months from the publication of the mention of the grant of the European patent, any person may give notice to the European Patent Office of opposition to the European patent granted. Notice of opposition shall be filed in a written reasoned statement. It shall not be deemed to have been filed until the opposition fee has been paid. (Art. 99(1) European Patent Convention).
Description

Field of the Invention

[0001] This invention relates to the detection of the occurrence of an apnea (i.e. the complete cessation of breathing) and to the determination of airway patency. The condition of patency of the airway is the converse of a total obstruction of the airway. The invention relates in particular to the detection of partial obstruction of the airway (i.e. obstructed breathing). The detection and monitoring of apneas, airway patency and obstruction is advantageous in the diagnosis and treatment of respiratory conditions that have adverse effects on a person’s wellbeing.

[0002] The expression "airway" as used herein is to be understood as the anatomical portion of the respiratory system between the nares and the bronchii, including the trachea. The expression "respiration" is to be understood as the continually repeating events of inspiration (inhaling) followed by expiration (exhaling).

Background of the Invention

[0003] In the Sleep Apnea syndrome a person stops breathing during sleep. Cessation of airflow for more than 10 seconds is called an "apnea". Apneas lead to decreased blood oxygenation and thus to disruption of sleep. Apneas are traditionally (but confusingly) categorized as either central, where there is no respiratory effort, or obstructive, where there is respiratory effort. With some central apneas, the airway is patent, and the subject is merely not attempting to breathe. Conversely, with other central apneas and all obstructive apneas, the airway is not patent (i.e. occluded). The occlusion is usually at the level of the tongue or soft palate.

[0004] The airway may also be partially obstructed (i.e. narrowed or partially patent). This also leads to decreased ventilation (hypopnea), decreased blood oxygenation and disturbed sleep.

[0005] The dangers of obstructed breathing during sleep are well known in relation to the Obstructive Sleep Apnea (OSA) syndrome. Apnea, hypopnea and heavy snoring are recognised as causes of sleep disruption and risk factors in certain types of heart disease. More recently it has been found that increased upper airway resistance (Upper Airway Resistance syndrome) during sleep without snoring or sleep apnea also can cause sleep fragmentation and daytime sleepiness. It is possible there is an evolution from upper airway resistance syndrome to sleep apnea, accompanied by a worsening of clinical symptoms and damage to the cardiovascular system.

[0006] The common form of treatment of these syndromes is the administering of Continuous Positive Airway Pressure (CPAP). The procedure for administering CPAP treatment has been well documented in both the technical and patent literature. Briefly stated, CPAP treatment acts as a pneumatic splint of the airway by the provision of a positive pressure, usually in the range 4 - 20 cm H2O (1 cm H2O = 98 Pa). The air is supplied to the airway by a motor driven blower whose outlet passes via an air delivery hose to a nose (or nose and/or mouth) mask sealingly engaged to a patient's face. An exhaust port is provided in the delivery tube proximate to the mask. More sophisticated forms of CPAP, such as bi-level CPAP and autosetting CPAP, are described in U.S. Patents No. 5,148,802 and 5,245,995 respectively.

[0007] Various techniques are known for sensing and detecting abnormal breathing patterns indicative of obstructed breathing. U.S. Patent No. 5,245,995, for example, describes how snoring and abnormal breathing patterns can be detected by inspiration and expiration pressure measurements while sleeping, thereby leading to early indication of preobstructive episodes or other forms of breathing disorder. Particularly, patterns of respiratory parameters are monitored, and CPAP pressure is raised on the detection of pre-defined patterns to provide increased airway pressure to, ideally, subvert the occurrence of the obstructive episodes and the other forms of breathing disorder.

[0008] As noted above, central apneas need not involve an obstruction of the airway, and often occur during very light sleep and also in patients with various cardiac, cerebrovascular and endocrine conditions unrelated to the state of the upper airway. In those cases where the apnea is occurring without obstruction of the airway, there is little benefit in treating the condition by techniques such as CPAP. Also, known automated CPAP systems cannot distinguish central apneas with an open airway from apneas with a closed airway, and may inappropriately seek to increase the CPAP splinting air pressure unnecessarily. Such unnecessary increases in pressure reflexly inhibit breathing, further aggravating the breathing disorder.

[0009] Other limitations associated with the prior art include the inability to detect airway patency and the absence of progressive, hierarchic response to increasingly severe indicators of airway obstruction for which the mask pressure should be increased.

[0010] It would be useful, however, to even more sensitively and reliably detect the conditions of partial obstruction, as well as apnea and patency, as this would assist in the design of equipment to prevent these conditions from occurring. In a similar way, means for detecting and monitoring mildly obstructed breathing would be useful in diagnosing and treating Upper Airway Resistance syndrome and monitoring that treatment is optimal.

[0011] In WO-A-9 309 834 there is disclosed an apparatus for detecting partial obstruction of the airway of a patient...
having the features of the pre-characterizing part of claim 1 hereof.

Summary of the Invention

According to the invention there is provided an apparatus for detecting partial obstruction of the airway of a patient according to claim 1 hereof.

Preferred features of the invention are set forth in the subordinate claims.

Brief Description of the Drawings

Embodiments of the invention will now be described with reference to the accompanying drawings, in which:

Fig. 1 shows a flow diagram of the basic methodology of an embodiment;
Fig. 2 shows, in diagrammatic form, apparatus embodying the invention;
Fig. 3 shows an alternative arrangement of the apparatus of Fig. 2;
Fig. 4 shows a graph of air flow with time for normal and partially obstructed inspiration;
Fig. 5 shows a flow diagram of the determination of an apnea;
Figs. 6a and 6b show a flow diagram of the calculation of the shape factors;
Fig. 7 shows a flow diagram of an embodiment utilising both shape factor methodologies;
Figs. 8a and 8b show clinical data of CPAP treatment utilising the shape factor methodologies;
Figs. 9a-9c and 10a-10c show clinical respiratory air flow and frequency signals during an apnea;
Fig. 11 shows a flow diagram for the cardiogenic determination of patency;
Figs. 12a-12d and 13a-13d show graphs of clinical respiratory data demonstrating the detection of patency;
Fig. 14 shows a flow diagram of an applied modulated output in the determination of patency;
Fig. 15 shows a flow diagram of leak compensated patency determination; and
Fig. 16 shows, in schematic form, a preferred CPAP treatment system.

Detailed Description of Preferred Embodiments and Best Mode

The methodology represented in Fig. 1 is of a clinical embodiment, where patient CPAP pressure is controlled over time as appropriate. A purely diagnostic embodiment operates in the same manner except it omits the CPAP pressure increase and pressure decrease actions of step 18 and step 17 respectively.
The pressure loss along tube 32 is calculated in step 74 from the flow through the tube \( f_{g}(t) \), and a knowledge of the pressure-flow characteristic of the tubing, for example by table lookup.

The pressure loss along tube 32 is then added to the desired set pressure at the mask \( p_{m} \) in summation step 78 to yield the desired instantaneous pressure at the pressure generator 34. Preferably, controller of the pressure generator 34 has a negative feedback input from the pressure transducer 70, so that the desired pressure from step 78 is achieved more accurately.

The flow through the exhaust 42 is calculated from the pressure at the mask (calculated in step 76) from the pressure-flow characteristic of the exhaust step 80, for example by table lookup.

Finally, the mask flow is calculated by subtracting the flow through the exhaust 42 from the flow through the tubing 32, in subtraction step 82.

The methodology put into place by the controller 62 will now be described with reference to the apparatus of Fig. 2.

A. Determination of Apnea

This section generally corresponds to steps 10, 12, and 14 as shown in Fig. 1.

Partial upper airway obstruction in untreated or partially treated Obstructive Sleep Apnea syndrome, and the related High Airway Resistance syndrome, leads to mid-inspiratory flow limitation, as shown in Fig. 4, which shows typical inspiratory waveforms respectively for normal and partially obstructed breaths.

As discussed previously, the respiratory air flow is determined by means of the differential pressure transducer 48, and a signal representing the air flow is continuously digitized and passed to the controller 62. If necessary, the controller 62 can linearise the flow signal, for example, by a table lookup. Occasionally, complete obstruction of the airway can occur unexpectedly, for example in a previously untreated patient, without a period of preceding partial obstruction. Consequently, the processing steps 12, 14 shown in Fig. 1 also detect the presence of complete or near-complete cessation of air flow, or apnea, using the measure of the Breathing Index in step 14.

This is achieved, for example as shown in Fig. 5, by low-pass filtering of the mask air flow signal \( f_{n} \) by low-pass filter element 125, typically with a 1 Hz cutoff, and calculating the moving average variance by the computational.
**B. Determination of Airway Obstruction**

[0045] The Obstruction Index is calculated in step 12. In accordance with the present invention, an Obstruction index referred to as shape factor 1 is calculated. In addition a further Obstruction index, referred to as shape factor 2, can be calculated.

[0046] The Obstruction Index is then compared with a threshold in step 16 of Fig. 1. If the obstruction Index is less than the threshold value, CPAP treatment pressure is increased in step 18. Otherwise, the CPAP pressure may be reduced in optional step 17.

[0047] As shown in Fig. 6a, the digitized airflow signal, \( f_n \), has any components below 0.1 Hz due to leaks of the mask 30 subtracted by a high-pass filter 90. The inspiratory and expiratory portions of each breath are then identified by a zero-crossing detector 92. A number of evenly spaced points (typically sixty-five), representing points in time, are interpolated by an interpolator 94 along the inspiratory flow-time curve for each breath. The curve described by the points is then scaled by a scaler 96 to have unity length (duration/period) and unity area to remove the effects of changing respiratory rate and depth.

[0048] Conveniently, the scaled breaths are compared in a comparator 98 with a pre-stored template representing a normal unobstructed breath. The template is very similar to the curve for a normal inspiratory event as shown in Fig. 4. Breaths deviating by more than a specified threshold (typically 1 scaled unit) at any time during the inspiration from this template, such as those due to coughs, sighs, swallows and hiccups, as determined by the test element 100, are rejected.

[0049] For data for which the test is satisfied, a moving average of the first such scaled point is calculated by the arithmetic processor 102 for the preceding several inspiratory events. This is repeated over the same inspiratory events for the second such point, and so on. Thus, sixty five scaled data points are generated by the arithmetic processor 102, and represent a moving average of the preceding several inspiratory events. The moving average of continuously updated values of the sixty five points are hereinafter called the "scaled flow", designated as \( f_S(t) \). Equally, a single inspiratory event can be utilised rather than a moving average.

[0050] From the scaled flow two shape factors that directly relate to the determination of partial obstruction are calculated. Each shape factor equates to the Obstruction Index discussed above.

[0051] Shape factor 1 is the ratio of the mean of the middle thirty-two scaled flow points to the mean overall sixty-five scaled flow points. This is thus a determination of the reduction of the magnitude (depression) of the mid-portion of the scaled inspiratory event(s). Since the mean for all sixty five points is unity, the division need not actually be
performed.

Mathematically, it is expressed as:

\[
\text{shape factor } 1 = \frac{1}{33} \sum_{t=16}^{48} f_s(t) - \frac{1}{65} \sum_{t=1}^{65} f_s(t)
\]

which reduces simply to

\[
\frac{1}{33} \sum_{t=16}^{48} f_s(t).
\]

For a normal inspiratory event this ratio will have an average value in excess of unity, because a normal such inspiratory event is of higher flow in the middle than elsewhere, as can be seen from Fig. 4. Conversely, for a severely flow-limited breath, the ratio will be unity or less, because flow limitation occurs particularly during the middle half of the breath when the upper airway suction collapsing pressure is maximal. A ratio of 1.17 is taken as the Threshold value (step 16 of Fig. 1) between partially obstructed and unobstructed breathing, and equates to a degree of obstruction that would permit maintenance of adequate oxygenation in a typical user.

In other embodiments the number of sampled points, number of breaths and number of middle points can be varied, and still achieve a meaningful determination of whether partial obstruction is occurring. The Threshold value similarly can be a value other than 1.17.

Alternatively, the second shape factor is calculated as the RMS deviation from unit scaled flow, taken over the middle thirty three points. This is essentially a measure of the flatness of the mid-portion of the scaled respiratory event(s). Expressed mathematically, this is:

\[
\text{shape factor } 2 = \sqrt{\frac{1}{32} \sum_{t=16}^{48} (f_s(t) - 1)^2}
\]

For a totally flow-limited breath, the flow amplitude vs. time curve would be a square wave and the RMS deviation would be zero. For a normal breath, the RMS deviation is approximately 0.2 units, and this deviation decreases as the flow limitation becomes more severe. A threshold value of 0.15 units is used in step 16 of Fig. 1.

While shape factor 1 can be utilised independently in implementing the methodology carried by the apparatus of Fig. 2, and result in the sensitive and reliable detection of partially obstructed breathing, better performance again is obtained by implementing both shape factors executed by the controller 62 so that both shape parameters act together. In this case, shape factor 2 is preferred for use to detect all but the most severe obstructions, and shape factor 1 therefore is preferred for detecting only the most severe obstructions, achieved by reducing the critical threshold from 1.17 to 1.0.

Fig. 7 is a flow diagram illustrating the principle of the two shape factors operating in concert. The scaled flow signal \( f_s(t) \) is provided to a shape detector 112, such as has been described with reference to Figs. 6a and 6b. The shape detector 112 generates shape factor 1 and shape factor 2. Shape factor 1 is applied to a decision block 114 and compared against the Threshold value of 1.0. If the outcome of the comparison is "Yes", then it is determined that there should be an increase in the CPAP pressure setting, as indicated in block 116. The shape factor 2 is provided to the decision block 118, and a comparison made against the Threshold value of 0.15. If the answer is "Yes", then it also is appropriate for an increase in the CPAP pressure, as shown in block 120.

In either case, if the results of the comparison is "No", then those results are ANDed in the AND gate 122. That is, an output will only be achieved if both Threshold criteria are not satisfied. In this case, there is no partial obstruction, or partial obstruction has subsided, in which case, as indicated in block 124, it is appropriate to decrease the CPAP pressure.
The methodology to improve performance of the snore detector firstly involves a determination of the blower motor speed. This can be achieved by a tachometer located on the motor. Then follows a determination of an expected pressure, measured in cm H$_2$O. The results are shown in Figs. 8a and 8b.

In this patient there was an 83% correlation between shape factor 1 (Fig. 8a) and UAP, with low values of shape parameter one associated with a high pressure drop across the upper airway, indicating partial obstruction. Similarly, there was an 89% correlation between shape factor 2 (Fig. 8b) and UAP.

In summary, the shape factors provide an index of the state of the airway. They provide a sensitive warning of an airway becoming unstable, and allow early CPAP treatment to occur. Continuing calculation of the moving average shape, and thus the shape factors, provides an accurate on-going assessment of the degree of any such apnea that is not subverted by CPAP treatment in order that modified appropriate treatment or corrective action can be taken.

The shape factors discussed above provide the most sensitive indication of upper airway stability, and therefore in the smallest increase in the CPAP pressure that should restore stability to the airway, and similarly a correspondingly small decrease in the CPAP pressure when stability has so been restored. By being able to maintain the increases to such a small level, the patient is less likely to be woken, and will also benefit from avoiding apneas with their associated health risks.

For example, when shape factor 1 is below 1.0, the CPAP pressure is increased in proportion to the amount of the ratio being below 1.0. An increase of 1 cm H$_2$O per breath per unit below a ratio of 1.0 has been found particularly effective. Conversely, if the ratio is above 1.0, the CPAP pressure is gradually reduced with a time constant of 20 minutes. If shape factor 2 is below 0.2, the CPAP pressure is increased at a rate of 1 cm H$_2$O per breath per unit below 0.2. Conversely, if the shape factor is above 0.2 units, the pressure is gradually lowered with a time constant of 20 minutes.

An example of experimental validation involved a subject with severe Obstructive Sleep Apnea syndrome placed on nasal CPAP therapy. A catheter tip pressure transducer was placed in the hypopharyngeal space, below the site of upper airway obstruction, and the peak upper airway pressure gradient (UAP) from hypopharynx to mask calculated for each breath.

The CPAP pressure was intentionally reduced from time to time during stable sleep, in order to produce partial upper airway obstruction. For each breath taken during the night, the two shape factors were calculated, and plotted against the UAP., measured in cm H$_2$O. The results are shown in Figs. 8a and 8b.

In this patient there was an 89% correlation between shape factor 1 (Fig. 8a) and UAP, with low values of shape parameter one associated with a high pressure drop across the upper airway, indicating partial obstruction. Similarly, there was an 89% correlation between shape factor 2 (Fig. 8b) and UAP.

The function achieved by shape factor 1 also can be achieved by an improved methodology in the detection of snoring. Prior art U.S. Patent No. 5,245,995 describes signal processing of the mask flow signal to determine a snore characteristic, particularly as shown in Figs. 9 and 10 of that document. The respiratory air flow signal is bandpass filtered in the range 30 - 300 Hz. Snoring exhibits characteristic frequencies in this range, and as described in the prior art reference the sound intensity of snoring is indicative of almost complete obstruction of the airway. Thus CPAP pressure is increased if the snore signal is in excess of a snore threshold value. This then corresponds to the degree of obstruction otherwise detected by shape factor 1.

Although the snore detector and CPAP treatment effected in consequence of the occurrence of snoring operates satisfactorily, there is still scope for improvement. Once particular problem comes in that some CPAP apparatus caused wind noise occurs in the 30 - 300 Hz range, as does background noise due to the motor driving the blower.

As described herein, the digitized flow signal f$_n$ has been arrived at in a similar manner to that described in prior art U.S. Patent No. 5,245,995, and thus includes snore component frequencies.

The methodology to improve performance of the snore detector firstly involves a determination of the blower motor speed. This can be achieved by a tachometer located on the motor. Then follows a determination of an expected flow signal such as would occur in the absence of snoring. This is calculated as a function of motor speed and airflow by the following formula:

$$\text{predicted signal} = k_1 \omega + k_2 \omega^2 + k_3 f + k_4 \frac{df}{dt}.$$
results in an increase in the CPAP pressure.

C. Determination of Airway Patency

[0072] If the outcome of step 14 is “Yes”, then an apnea in progress. In accordance with the methodology of Fig. 1, a determination of airway patency (step 20) is made. Two methods are now described. The first is a measurement by cardiogenic airflow, and the second is an externally induced oscillation technique.

1. Cardiogenic Airflow

[0073] With each beat of the heart, of the order of 66 ml of blood is ejected from the chest over about 0.3 sec, producing a pulsatile blood flow out of the chest of the order of 0.22 l/sec peak flow. If the chest wall were rigid this would create a partial vacuum in the chest cavity, and, if the upper airway were open and of zero resistance, a similar quantity of air would be sucked in through the trachea.

[0074] In practice, the chest wall is not totally rigid, and the upper airway has a finite resistance. Consequently the observed airflow with each beat of the heart is of the order of 0.02 to 0.1 l/sec. If there is a central apnea with an open airway, there will be a very small pulsatile airflow of the order of 0.02 to 0.1 l/sec in time with the heart beat. Conversely, if the airway is closed, there will be no pulsatile airflow in time with the heart beat.

[0075] Figs. 9a-9c represent a central apnea with an open airway lasting approximately 30 seconds, determined from diaphragm electromyogram tracings (not shown). Conversely, Figs. 10a-10c represent an obstructive apnea with a closed airway. Figs. 9a and 10a respectively show a respiratory airflow signal, f(t), during which an apnea lasting approximately 25 seconds occurs, indicated by a near cessation of airflow.

[0076] Figs. 9b and 10b respectively show a ten second close-up (between t = 11.5 s to t = 21.5 s) of the airflow signal during the apnea. It can be noted that in Fig. 9b, where the airway is open, small rhythmic oscillations in the airflow are seen, with the expected peak flow of about 0.1 l/sec. Inspection of the corresponding electrocardiogram (not shown) confirms that these oscillations are of cardiac origin, with airflow either phase-locked with the heartbeat, or at exactly double the cardiac rate. Conversely, in Fig. 10b, there is either no airflow at all, or at least irregular airflow due to not quite complete obstruction.

[0077] Figs. 9c and 10c respectively show the discrete Fourier transform of Figs. 9b and 10b. In Fig. 9c (open airway), there are strong peaks in the frequency spectrum at around 1.25 Hz and/or 2.5 Hz corresponding to the heart rate and its first harmonic. The peaks reach an amplitude of at least 0.01 l/sec. Conversely, in Fig. 10c (closed airway), the discrete Fourier transform shows little or no activity between 0.75 and 3 Hz.

[0078] The methodology firstly records the airflow, f(t), using by the flow transducer 48 shown in Fig. 2 or Fig. 3. The signal is digitized, for example at 50 Hz, using the analog-to-digital converter (ADC) 54, and sampled by the controller 62. The subsequent processing steps are shown in Fig. 11.

[0079] If required, the flow signal, fn, is digitally bandpass filtered by the bandpass filter 130 between 0.1 and 6 Hz to remove low frequency components (leak) and high frequency components (noise) to yield a clean respiratory air flow signal.

[0080] The occurrence of an apnea will have previously been determined by, for example, the Breathing Index derived in Fig. 5. In that case the process continues.

[0081] A Discrete Fourier transform (DFT) is performed, by the processing element 132, of the airflow signal fn during the apnea. Only terms up to 6 Hz need to be calculated. In the case where the heart rate is not known, processing is as follows; if the amplitude of the DFT exceeds a Threshold value of 0.01 l/sec, as determined by the peak height detector 136 and the subsequent comparator element 138, the airway is declared open; otherwise it is declared closed. Patency Index 1 represents the output of the peak height detector 136.

[0082] If an electrocardiogram or other indicator of heartbeat, such as a pulse oximeter is available, then an appropriate method is to:

1. Use a digital or electronic trigger to trigger on each heart beat.
2. Accumulate the respiratory airflow signal at time nT after receipt of each trigger into element n of an array, summing with previous values at time nT for the duration of the apnea.
3. Divide by the number of heartbeats to obtain the average air flow as a function of time into the heartbeat.
4. Calculate the first two terms of the DFT of this signal (fundamental and first harmonic) and inspect for an amplitude of the order of 0.1 l/sec.

[0083] In such a case where the heart rate is known, then only the amplitudes at the heart rate and its first harmonic need be considered, leading to a more accurate estimation.
Instead of using the DFT, any suitable mathematical method of detecting a rhythmic oscillation with a frequency of the anticipated heart rate and its first harmonic (0.75 to 3 Hz) will suffice. Such methods could include measuring the regularity of peak heights and zero crossings, autocorrelation, or other digital filtering methods.

2. Externally Induced Oscillations

If the airway is open, but the respiratory muscles are relaxed (i.e. a central apnea with open airway), then small externally originating fluctuations in the mask pressure will induce a small respiratory airflow by inflating and deflating the lungs, and by compressing and decompressing the gas in the lungs. Conversely, if the airway is closed, no airflow will be induced.

Fig. 12a shows a respiratory airflow signal as a function of time during nasal CPAP therapy. In the first half of the tracing, there is a central apnea with open airway lasting approximately 22 seconds. Fig. 12b shows that the CPAP pressure is approximately 15.5 cm H₂O. The high frequency “noise” apparent through most of the pressure trace is largely due to cardiogenic airflow as previously discussed.

Approximately 5 seconds into the apnea a 2 Hz, 1 cm H₂O pressure oscillation is induced (applied) for 6 seconds (i.e. between t = 14 s to t = 20.5 s). It can be seen that this pressure modulation induces a corresponding 2 Hz modulation in the respiratory air flow signal. Figs. 12c-12d are an enlargement of the period of testing. The respiratory air flow signal has an amplitude of approximately + 0.2 l/sec.

Conversely, in Figs 13a-13d there is an obstructive apnea, with a closed airway. A similar tracing would be seen with a central apnea with a closed airway. It can be seen that in this case there is no obvious induced flow signal during the 6 second period of 2 Hz pressure oscillations. The mean induced signal was 0.01 l/sec.

The procedure is typically, at 4 - 6 seconds into the apnea, the CPAP pressure generator output pressure supplied to the motor-servo unit 40 is controlled to produce a modulated pressure output. As shown in Fig. 14, the output from the generation element 140 (controller 62) is a signal modulated with a low amplitude square wave, typically at 2 - 4 Hz. This produces a quasi-sinusoidal oscillation in the mask pressure, with a typical amplitude of 0.5 - 1 cm H₂O.

As further shown in Fig. 14, the air flow induced by the pressure modulation is separated from air flow induced by other factors (such as heartbeat), by demodulating the measured air flow signal, fᵣ, by a demodulator 142 with the 2 Hz driving signal. The components at 0 degrees and 90 degrees to the output signal are calculated, and their amplitudes are added vectorially to yield a mean induced air flow signal amplitude (Patency Index 2). The mean signal in this case is 0.12 l/sec.

Apneas are classified as "airway open" if the mean induced signal is more than 0.03 l/sec, and "airway closed" if the mean induced signal is less than 0.03 l/sec. Alternatively, the mean induced signal could be divided by the amplitude of the inducing pressure to yield the conductance (degree of openness) as a continuous variable.

When it is desired to determine the state of the airway in the presence of typical CPAP treatment, it is preferable to take into account the effect of mask leaks. A leak between the mask and the face can produce a false positive induced air flow signal. As shown in Fig. 15, the oscillator 140 induces the low-frequency, low amplitude pressure oscillations as previously described. The air flow signal fᵣ is high pass filtered by the high pass filter 148 (typically 0.1 Hz) to remove leak, and passed to the demodulator 146, which produces Patency Index 2 as previously described.

The flow signal is also low pass filtered (typically 0.1 Hz) by the low pass filter 150 to derive a measurement of leak. The value calculated in step 142 represents the sum of the induced signal due to modulation of respiratory air flow and the induced signal due to modulation of flow through the leak. The induced signal due to modulation of flow through the leak is then calculated by arithmetic element 154, as:

\[
\frac{0.5 \cdot \text{leak} \cdot \text{inducing oscillation amplitude}}{\text{mean mask pressure}}
\]

This is then subtracted by the subtractor 156 from the uncompensated Patency Index to produce a leak-compensated Patency Index. The leak-compensated Patency Index can optionally be divided by the inducing oscillation amplitude to yield airway conductance, as described previously.

In the case of either methodology utilised to determine patency, if the result of that determination (step 20) is “No”, then as was the case for a partial obstruction, the CPAP treatment pressure is increased. If the result is “Yes”, then a central apnea with an open airway is occurring, and it is inappropriate to increase CPAP pressure. Instead the event is only logged, and step 17 follows, whereby CPAP pressure is reduced, as has previously been discussed.
3. Extensions to the Methodology of Determining Patency

[0096]

(1) Instead of declaring the airway open or closed, the airway can be declared open to a certain degree. For example, if the peak amplitude of the DFT was 50% of the threshold, the airway is taken as being patent to degree 0.5. Similarly with the externally induced oscillation method.

(2) Instead of using the entire duration of the apnea, calculations can be performed on a moving window of appropriate duration, such as 10 seconds. In this way, mixed apneas, in which the airway is open for only part of the apnea, can be detected.

(3) Other methods of measuring or inferring respiratory airflow can be utilised. For example, instead of measuring mask airflow with a flow-resistive element and differential pressure transducer, mask airflow could be measured using an ultrasonic flow transducer, or inferred from mask pressure, using a single ended pressure transducer. Alternatively, measurements of chest wall and/or abdominal movement (such as magnetometers, inductance plethysmography, or strain gauges) could be used.

D. A Combined System for Automatic Adjustment of CPAP Pressure

[0097] Fig. 16 illustrates, in schematic block form, a particular preferred embodiment of CPAP treatment apparatus. The CPAP machine 164 represents the component element shown in Fig. 2 or Fig. 3 except for the elements bearing the reference numerals 54, 58, 60 and 62. All of the logic blocks 166-176 are processing steps implemented in a microcontroller, which, in Fig. 2, is referred to by the reference numeral 62. The embodiment implements a hierarchic methodology, based around the methodology of Fig. 1, that allows the progressive use of pre-obstructive and obstructive indications to trigger CPAP treatment pressure increases of magnitude and duration appropriate for the severity of the event.

[0098] The mask pressure is initially set to a low pressure, typically 4 cm H2O. Whenever the apnea detector 168 detects an apnea, the airway patency detector 170 determines whether the airway is open or closed by the forced oscillation method, and if closed, the mask pressure is increased, typically by 1 cm H2O per 15 seconds of apnea. If a central apnea is occurring, no increase in CPAP pressure is instructed.

[0099] If a snore is detected by the snore detector 172 (such as that disclosed in U.S. Patent No. 5,245,995) the mask pressure is also increased. If the snore index on the given breath exceeds a critical threshold value, the pressure is increased by 1 cm H2O per unit above the threshold value. The defaults threshold for the snore index is 0.2 units, corresponding approximately to a snore that can only just be reliably detected by a technician standing at the bedside. The rate of rise in pressure is limited to a maximum of 0.2 cm H2O per second, or 12 cm H2O per minute.

[0100] In some patients, it is not possible to prevent the occasional snore, even at maximum pressure. Consequently, above pressures of 10 cm H2O, a heuristic methodology is used to perform a trade-off between the possible advantage of increasing the pressure and the disadvantage of increased side effects. Thus the threshold is adjusted as follows:

<table>
<thead>
<tr>
<th>Pressure (cm H2O)</th>
<th>Threshold (snore units)</th>
<th>Description</th>
</tr>
</thead>
<tbody>
<tr>
<td>&lt; 10</td>
<td>0.2</td>
<td>very soft</td>
</tr>
<tr>
<td>10-12</td>
<td>0.25</td>
<td></td>
</tr>
<tr>
<td>12-14</td>
<td>0.3</td>
<td>soft</td>
</tr>
<tr>
<td>14-16</td>
<td>0.4</td>
<td></td>
</tr>
<tr>
<td>16-18</td>
<td>0.6</td>
<td>moderate</td>
</tr>
<tr>
<td>&gt; 18</td>
<td>1.8</td>
<td>loud</td>
</tr>
</tbody>
</table>

[0101] If the shape factor 2 is less than the threshold value, the mask pressure also is increased. The default threshold value is 0.15 units. The default rate of increase of pressure is such that a severely abnormal shape factor of 0.05 units will produce a rise in pressure of 0.3 cm H2O per breath, or approximately 4.5 cm H2O per minute.

[0102] The lips and tongue can sometimes act like a one-way valve, forming a seal during inspiration when the pharyngeal pressure is lowest but failing during early to mid-expiration when the pressure is highest. Large leaks, and particularly valve-like leaks, can cause the shape factor to read low, falsely implying flow limitation. To compensate for this, the default threshold is increased according to an empirical heuristic technique if there is a large leak, or if there is a valve-like leak. This is to avoid the treatment pressure being increased unnecessarily. Consequently, in the presence of a large leak, more reliance is placed on the snore and apnea detectors.
In some patients, the shape factor does not become normal even at maximum pressure. Consequently, a further heuristic trade-off is made between possible increases in patency within increasing pressure, versus increasing side effects. The heuristics used are as follows:

(i) If the leak exceeds 0.7 l/sec, the critical threshold for the shape factor is 0. In the range 0.3 - 0.7 l/sec, the threshold is decreased proportionately, so that as the leak increases more severe flattening is required before the pressure will rise.

(ii) An index of the presence of valve-like leaks is calculated as the ratio of the peak flow during the first 0.5 seconds of expiration to the mean flow during the second 0.5 seconds of expiration. If this ratio exceeds 5:1, the threshold is 0. In the range 4:1 to 5:1, the threshold is reduced proportionately.

(iii) If the mask pressure is 20 cm H₂O, the threshold is 0, and is reduced proportionately in the range 10 - 20 cm H₂O. For example, if the leak is 0.4 l/sec, and the mask pressure is 15 cm H₂O, the threshold is reduced by 25% because of the leak, and a further 50% because of the already high treatment pressure so that the new threshold is 0.056 units. Conversely, if no abnormality is detected on a particular breath (block 176), the mask pressure is reduced with an appropriate time constant, typically 10-20 minutes per cm H₂O for snore or shape factor changes, and preferably, about 40 minutes per cm H₂O following apneas.

The preferred embodiment of the combined system for automatic adjustment of CPAP treatment pressure described above was used to treat 28 patients with previously untreated obstructive sleep apnea syndrome. CPAP pressure commenced at 4 cm H₂O, and increased automatically in response to closed airway apneas, snoring, and inspiratory air flow limitation. The following table compares results with those obtained in the same subjects without treatment:

<table>
<thead>
<tr>
<th></th>
<th>Untreated (mean ± SEM)</th>
<th>Treated (mean ± SEM)</th>
</tr>
</thead>
<tbody>
<tr>
<td>Apnea Index (events/hr)</td>
<td>35.5 ± 5.9</td>
<td>1.5 ± 0.32</td>
</tr>
<tr>
<td>Time in Apnea (Percent of night)</td>
<td>24.5 ± 4.7</td>
<td>1.0 ± 0.37</td>
</tr>
<tr>
<td>Slow Wave Sleep (Percent of night)</td>
<td>7.0 ± 1.6</td>
<td>20.0 ± 2.2</td>
</tr>
<tr>
<td>REM Sleep (Percent of night)</td>
<td>9.4 ± 1.4</td>
<td>20.3 ± 2.1</td>
</tr>
<tr>
<td>Arousal Index (Events/hr)</td>
<td>55.9 ± 5.3</td>
<td>10.8 ± 1.9</td>
</tr>
<tr>
<td>Respiratory Arousals (Events/hr)</td>
<td>51.5 ± 5.4</td>
<td>4.2 ± 1.5</td>
</tr>
</tbody>
</table>

There was a dramatic reduction in the number of apneas per hour, and the percentage of time in apnea. There was a large increase in the percentage of deep restorative sleep (slow wave and REM sleep). There was a dramatic reduction in the number of arousals from sleep, particularly those of a respiratory origin. These results confirm that the combined system produces excellent results in treating obstructive sleep apnea syndrome.

The system described can also be utilised in a diagnostic mode, typically where nasal cannulae are utilized in the place of a mask arrangement sealed to the patient's face. In this mode, measurements of apneas, patency, and partial obstruction are logged, but no CPAP treatment is effected. The nasal cannulae are connected to one side of the flow sensor 50 in Fig. 2. Only elements 50, 54, 56, 58, 60 and 62 are required in this mode. Since with nasal cannulae, the signal from the flow transducer 50 is not linear with flow, there is an additional step in which the signal from the flow transducer is linearized, preferably by use of a lookup table in the microcontroller 62. The data collected provides the physician with the ability to diagnose conditions such as Obstructive Sleep Apnea syndrome and Upper Airway Resistance syndrome.

Numerous alterations and modification, as would be apparent to one skilled in the art, can be made without departing from the basic inventive concept.

More complex variants of CPAP therapy, such as bi-level CPAP therapy or therapy in which the mask pressure is modulated within a breath, can also be monitored and/or controlled using the methods described herein.

The moving average variance apnea detector, as described, can be extended to include a hypopnea detector by adding a second comparator set at a higher threshold, so that it will respond to partial reductions in ventilation.
Claims

1. An apparatus for detecting partial obstruction of the airway of a patient, the apparatus comprising:
   (a) air flow measurement means (50, 56) for deriving a patient respiratory air flow signal; and
   (b) processing means (54, 62) coupled to said air flow measurement means (50, 56) to receive said air flow
   signal, and having a plurality of processing elements comprising:
      (i) a sampling element (54) for sampling said airflow signal at multiple times during a breath; and
      (ii) a detection element (92) for identifying the inspiratory samples from said sampling element;
   characterized by
   (iii) a computational element (94, 96, 98, 100) to calculate a measure of partial obstruction which is a ratio
   of the mean of a mid-portion of said inspiratory samples to the mean of said inspiratory samples.

2. The apparatus of claim 1 wherein said computational element (94, 96, 98, 100) determines a second measure of
   partial obstruction as the root mean square deviation of a mid-portion of the inspiratory values to the mean of said
   inspiratory values.

3. The apparatus of claim 2 wherein said computational element (94, 96, 98, 100) scales the inspiratory samples so
   that the inspiratory samples represent an inspiratory flow versus time curve having unit duration and unit area.

4. The apparatus of claim 3 wherein said computational element (94, 96, 98, 100) rejects inspiratory samples deviating
   by more than a threshold value.

5. The apparatus of claim 2 wherein said computational element (94, 96, 98, 100) further compares said measure to
   a threshold value and said second measure to a threshold value.

6. The apparatus of claim 5 wherein said computational element (94, 96, 98, 100) determines the existence of partial
   obstruction if either of said measure or said second measure is less than its compared to threshold.

7. The apparatus of claim 5 wherein said computational element (94, 96, 98, 100) determines the absence of partial
   obstruction if neither of said measure or said second measure is less than its compared to threshold.

8. Apparatus as claimed in any one of the preceding claims, further for determining the occurrence of an apnea, and
   wherein said computational element (125, 126) is further adapted to determine the variance of said measured air
   flow, and said data processing means (54, 62) further includes a comparing element (127) adapted to compare
   said variance with a predetermined threshold value, and if said variance falls below said threshold value then it is
   determined that an apnea is occurring.

9. Apparatus as claimed in claim 8, wherein said computational element (125, 126) is adapted to determine said
   variance as a moving average over a time window.

10. Apparatus as claimed in any one of the preceding claims, further for determining patency of said airway, said data
    processing means (54, 62) further including detection means (132, 134, 136) adapted to detect the presence of
    cardiogenic components in said air flow to determine that said airway is patent.

11. Apparatus as claimed in any one of the preceding claims, further for controlling the administration of CPAP appa-
    ratus, the apparatus further comprising:

       a mask (30) to be worn by a patient to provide breathable gas at a pressure elevated above atmospheric
       pressure continuously to a patient's airway;
       a conduit (32) coupled to said mask (30);
       a controllable turbine (34, 38) coupled to said conduit to supply at its outlet said pressurized breathable gas
       to said mask via said conduit;
       control means (40) to control turbine outlet pressure in response to a control signal;
and wherein said respiratory airflow signal is derived from said mask (30), said conduit (32) or said turbine
(34, 38), and said data processing means (54, 62) is adapted to generate said control signal such that, if said
device exceeds a threshold, then the control signal causes CPAP pressure to be decreased, and if said
deviation is less than a threshold, then the control signal causes CPAP pressure to be increased.

12. Apparatus as claimed in claim 11, wherein the data processing means (54, 62) includes a detector (172) that is adapted to determine the loudness of snoring, and if the loudness of snoring exceeds a threshold the data processing means (54, 62) causes the control signal to adjust the CPAP pressure as a function of both the measure of partial obstruction and the loudness of snoring.

13. Apparatus as claimed in claim 12, wherein if either the measure of obstruction exceeds said deviation threshold or the measure of loudness of snoring exceeds said snoring threshold, the data processing means (54, 62) causes the control signal to increase the CPAP pressure, but to decrease it otherwise.

14. Apparatus as claimed in claim 13, wherein the extent of the increase of CPAP pressure is determined by the data processing means (54, 62) to be: the extent by which the measured degree of airway obstruction exceeds the obstruction threshold plus a second amount proportional to the extent by which the measured degree of snoring exceeds the snoring threshold.

15. Apparatus as claimed in claim 14, wherein the snoring threshold is an increasing function of at least one of the current pressure applied by the CPAP apparatus and the extent of any leak from the CPAP apparatus.

Patentansprüche

1. Vorrichtung zum Selektieren einer teilweisen Obstruktion bzw. eines Hindernisses in dem Luftweg eines Patienten, wobei die Vorrichtung umfaßt:

   (a) Luftstrommeßmittel (50, 56), um ein Atemluftstromsignal eines Patienten abzuleiten; und
   (b) Bearbeitungsmittel (54, 62), die mit den Luftstrommeßmitteln (50, 56) gekoppelt sind, um das Luftstromsignal zu erhalten, und die eine Mehrzahl von Ver- bzw. Bearbeitungselementen enthalten:

   (i) ein Probensammelelement (54) zum Sammeln bzw. Auswerten des Luftstromsignals zu mehreren Zeiten während eines Atemhubs; und
   (ii) ein Detektionselement (92) zum Identifizieren der inspiratorischen Proben von dem Sammelelement; gekennzeichnet dadurch, daß die Vorrichtung weiters umfaßt:

   (iii) ein Berechnungs- bzw. Computerelement (94, 96, 98, 100), um ein Maß einer teilweisen Obstruktion bzw. eines Hindernisses zu berechnen, welches ein Verhältnis des Mittelwerts eines Mittelbereichs der inspiratorischen Proben zu dem Mittelwert der inspiratorischen Proben ist.

2. Vorrichtung nach Anspruch 1, worin das Berechnungselement (94, 96, 98, 100) ein zweites Maß einer teilweisen Obstruktion als die mittlere quadratische Abweichung bzw. RMS-Abweichung eines Mittelbereichs der inspiratorischen Werte zu dem Mittelwert der inspiratorischen Werte bestimmt.

3. Vorrichtung nach Anspruch 2, worin das Berechnungselement (94, 96, 98, 100) die inspiratorischen Proben so skaliert, daß die inspiratorischen Proben einen inspiratorischen Fluß bzw. Strom gegenüber einer Zeitkurve, die eine Einheitsdauer und eine Einheitsfläche aufweist, darstellen.

4. Vorrichtung nach Anspruch 3, worin das Berechnungselement (94, 96, 98, 100) inspiratorische Proben, die um mehr als um einen Schwellwert abweichen, zurückweist.

5. Vorrichtung nach Anspruch 2, worin das Berechnungselement (94, 96, 98, 100) weiters dieses Maß mit einem Schwellwert und dieses zweite Maß mit einem Schwellwert vergleicht.

6. Vorrichtung nach Anspruch 5, worin das Berechnungselement (94, 96, 98, 100) die Existenz einer teilweisen Obstruktion bestimmt, wenn eines von diesem Maß oder diesem zweiten Maß kleiner als der verglichene Schwellwert ist.

7. Vorrichtung nach Anspruch 5, worin das Berechnungselement (94, 96, 98, 100) die Abwesenheit einer teilweisen Obstruktion bestimmt, wenn keines von diesem Maß oder diesem zweiten Maß kleiner als der verglichene Schwellwert ist.
8. Vorrichtung nach einem der vorhergehenden Ansprüche, weiters zur Bestimmung des Auftretens von Apnoe, und worin das Berechnungselement (125, 126) weiters adaptiert ist, um die Varianz des gemessenen Luftstroms zu bestimmen, und die Datenverarbeitungsmittel (54, 62) weiters ein vergleichendes bzw. Vergleichselement (127) umfassen, das adaptiert ist, um die Varianz mit einem vorbestimmten Schwellwert zu vergleichen, und wobei, wenn die Varianz unter den Schwellwert fällt, dann bestimmt wird, daß eine Apnoe auftritt.

9. Vorrichtung nach Anspruch 8, worin das Berechnungselement (125, 126) adaptiert ist, um die Varianz als sich bewegender Durchschnitt über ein Zeitfenster zu bestimmen.

10. Vorrichtung nach einem der vorhergehenden Ansprüche, weiters zur Bestimmung einer Durchgängigkeit des Luftwegs, wobei die Datenverarbeitungsmittel (54, 62) weiters Detektionsmittel (132, 134, 136) umfassen, die adaptiert sind, um die Anwesenheit von kardiogenen Komponenten in dem Luftstrom zu detektieren, um zu bestimmen, daß der Luftweg durchgängig ist.

11. Vorrichtung nach einem der vorhergehenden Ansprüche, weiters zur Steuerung bzw. Regelung der Verabreichung einer CPAP-Vorrichtung, wobei die Vorrichtung weiters umfaßt:
   eine Maske (30), die durch einen Patienten zu tragen ist, um atembares bzw. Atemgas mit einem Druck höher als Atmosphärendruck kontinuierlich einem Luftweg eines Patienten zuzuführen;
   eine Leitung (32), die mit der Maske (30) gekoppelt ist;
   eine steuer-bzw. regelbare Turbine (34, 38), die mit der Leitung verbunden ist, um an ihrem Auslaß das unter Druck gesetzte Atemgas zu der Maske über die Leitung zuzuführen;
   Steuer-bzw. Regelmittel (40), um den Turbinenauslaßdruck in Antwort auf ein Steuer-bzw. Regelsignal zu steuern bzw. zu regeln;


15. Vorrichtung nach Anspruch 14, worin der Schnarchschwellwert eine ansteigende Funktion von wenigstens einem aus dem gegenwärtigen Druck, der durch die CPAP-Vorrichtung aufgebracht bzw. angelegt ist, und dem Ausmaß von jeglichem Leck aus der CPAP-Vorrichtung ist.

Revendications

1. Appareil destiné à détecter une obstruction partielle des voies respiratoires d'un patient, l'appareil comportant :
   (a) des moyens de mesure d'écoulement d'air (50, 56) destinés à obtenir un signal d'écoulement d'air respiratoire de patient; et
   (b) des moyens de traitement (54, 62) reliés aux dits moyens de mesure d'écoulement d'air (50, 56) afin de
recevoir ledit signal d'écoulement d'air, et ayant une multiplicité d'éléments de traitement comportant:

(i) un élément d'échantillonnage (54) pour l'échantillonnage dudit signal d'écoulement d'air à plusieurs instants pendant une respiration; et
(ii) un élément de détection (92) destiné à identifier les échantillons inspiratoires provenant dudit élément d'échantillonnage;

 caractérisé en ce que l'appareil comporte en outre:
(iii) un élément de calcul (94, 96, 98, 100) destiné à calculer une mesure de l'obstruction partielle qui est un rapport de la moyenne d'une partie médiane désdits échantillons inspiratoires sur la moyenne désdits échantillons inspiratoires.

2. Appareil selon la revendication 1, dans lequel ledit élément de calcul (94, 96, 98, 100) détermine une deuxième mesure d'obstruction partielle sous la forme de l'écart quadratique moyen d'une partie médiane des valeurs inspiratoires par rapport à la moyenne désdits valeurs inspiratoires.

3. Appareil selon la revendication 2, dans lequel ledit élément de calcul (94, 96, 98, 100) mesure les échantillons inspiratoires de telle sorte que les échantillons inspiratoires représentent une courbe de l'écoulement inspiratoire par rapport au temps ayant unité de durée et une unité de surface.

4. Appareil selon la revendication 3, dans lequel ledit élément de calcul (94, 96, 98, 100) rejette les échantillons inspiratoires s'écartant de plus d'une valeur de seuil.

5. Appareil selon la revendication 2, dans lequel ledit élément de calcul (94, 96, 98, 100) compare en outre ladite mesure à une valeur de seuil et ladite deuxième mesure à une valeur de seuil.

6. Appareil selon la revendication 5, dans lequel ledit élément de calcul (94, 96, 98, 100) détermine l'existence de l'obstruction partielle si ladite mesure ou ladite deuxième mesure est inférieure à son seuil comparé.

7. Appareil selon la revendication 5, dans lequel ledit élément de calcul (94, 96, 98, 100) détermine l'absence d'obstruction partielle si ni ladite mesure ni ladite deuxième mesure n'est inférieure à son seuil comparé.

8. Appareil selon l'une quelconque des revendications précédentes, destiné à déterminer en outre l'apparition d'une apnée, et, dans lequel ledit élément de calcul (125, 126) est en outre prévu pour déterminer la variance dudit écoulement d'air mesuré, et lesdits moyens de traitement de données (54, 62) comprennent en outre un élément de comparaison (127) prévu pour comparer ladite variance à une valeur de seuil prédéterminée, et si ladite variance tombe en dessous de ladite valeur de seuil, on détermine alors qu'une apnée se produit.

9. Appareil selon la revendication 8, dans lequel ledit élément de calcul (125, 126) est prévu pour déterminer ladite variance sous forme d'une moyenne mobile sur une fenêtre de temps.

10. Appareil selon l'une quelconque des revendications précédentes, destiné en outre à déterminer l'état ouvert desdites voies respiratoires, lesdits moyens de traitement de données (54, 62) comprenant en outre des moyens de détection (132, 134, 136) prévus pour détecter la présence de composants cardiogéniques dans ledit écoulement d'air afin de déterminer que lesdites voies respiratoires sont ouvertes.

11. Appareil selon l'une quelconque des revendications précédentes, destiné en outre à commander l'administration d'un appareil CPAP, l'appareil comportant en outre:

un masque (30) devant être porté par un patient afin de délivrer en continu aux voies respiratoires d'un patient un gaz respirable à une pression élevée au-dessus de la pression atmosphérique;
un conduit (32) relié audit masque (30);
ume turbine pouvant être commandée (34, 38) reliée audit conduit afin de délivrer à sa sortie ledit gaz respirable pressurisé audit masque par l'intermédiaire dudit conduit;
des moyens de commande (40) destinés à commander la pression de sortie de turbine en réponse à un signal de commande;
et dans lequel ledit signal d'écoulement d'air respiratoire est obtenu à partir dudit masque (30), dudit conduit (32) ou de ladite turbine (34, 38), et lesdits moyens de calcul (54, 62) sont prévus pour générer ledit signal de commande de telle sorte que, si ledit écart dépasse un seuil, le signal de commande amène alors la pression
CPAP à être diminuée, et si ledit écart est inférieur à un seuil, le signal de commande amène alors la pression CPAP à être augmentée.

12. Appareil selon la revendication 11, dans lequel les moyens de traitement de données (54, 62) comprennent un détecteur (172) qui est prévu pour déterminer le volume du ronflement, et si le volume du ronflement dépasse un seuil, les moyens de traitement de données (54, 62) amènent le signal de commande à ajuster la pression CPAP en fonction à la fois de la mesure de l'obstruction partielle et du volume du ronflement.

13. Appareil selon la revendication 12, dans lequel, si la mesure de l'obstruction dépasse ledit seuil d'écart ou si la mesure de volume du ronflement dépasse ledit seuil de ronflement, les moyens de traitement de données (54, 62) amènent le signal de commande à ajuster la pression CPAP, mais à la diminuer autrement.

14. Appareil selon la revendication 13, dans lequel la valeur de l'augmentation de la pression CPAP est déterminée par les moyens de traitement de données (54, 62) afin d'être : la valeur selon laquelle le degré mesuré d'obstruction des voies respiratoires dépasse le seuil d'obstruction plus une deuxième quantité proportionnelle à la valeur selon laquelle le degré mesuré de ronflement dépasse le seuil de ronflement.

15. Appareil selon la revendication 14, dans lequel le seuil de ronflement est une fonction croissante de la valeur la plus petite parmi la pression courante appliquée par l'appareil CPAP ou la valeur d'une fuite quelconque de l'appareil CPAP.
FIG. 5

125

126

Breathing Index

127

Trigger

128

Threshold

129

Ceased for more than 10 seconds?

NO: No Apnea

YES: Apnea
FIG. 12c

FLOW (L/sec)

14 16 18 20

FIG. 12d

PRESSURE (cm H2O)

12 14 16 18