EUROPEAN PATENT SPECIFICATION

AEROSOL DELIVERY APPARATUS WITH POSITIVE EXPIRATORY PRESSURE CAPACITY

AEROSOLSPENDER MIT EINER MÖGLICHKEIT FÜR POSITIVEN AUSATEMDRUCK

APPAREIL AEROSOL AVEC UNE CAPACITE DE PRESSION EXPIRATOIRE POSITIVE


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Description

FIELD OF THE INVENTION

[0001] The invention relates to an apparatus and method for performing Positive Expiratory Pressure (PEP) therapy. More particularly, this invention relates to a method and apparatus for performing PEP therapy alone or in conjunction with an aerosol delivery apparatus.

BACKGROUND

[0002] PEP therapy is used primarily in pulmonary secretion removal. Devices used to perform PEP therapy provide positive pressure during expiration. The patient or user exhales against a fixed orifice resistor and generates a pressure ranging approximately from 10-20 cm H2O. The resistance orifice is an important consideration and is typically set by a physician, veterinarian, or a skilled practitioner in the art. An orifice that is too large may result in a short exhalation that will not produce proper expiratory pressure. An orifice that is too small may result in a longer expiratory phase that raises the pressure above approximately 20 cm H2O and ultimately increases the work of breathing.

[0003] During the exhalation phase of PEP therapy, the airway is splinted open by the pressure. This causes the movement of secretions from the peripheral airways into the larger airways where they can be expelled. PEP therapy usually lasts for about 10-20 minutes and is performed as required, generally 1-4 times per day. Typically, the patient performs 10-20 PEP breaths, removes the device from their mouth and follows this with a forceful exhalation. This final exhalation triggers a cough that loosens secretions.

[0004] Studies indicate that PEP therapy dilates the airways and improves the distribution of ventilation, resulting in a better deposition of an inhaled substance, such as, but not limited to, a medicine or remedy. As used herein, the term "aerosol delivery apparatus" means any apparatus capable of producing and/or delivering a substance, such as, but not limited to, a medicament, may be provided a positive respiratory pressure apparatus having the features of claim 1.

[0005] Prior art document US-A-4267832 shows an exploration valve apparatus for use with a respirator. The device includes a valve means directing exhaled air through an oval opening adjustable by the overlap of two apertures which are moveable relative to each other. The valve means does not exclude entrainment of air through the exhalation opening into the stream of air flowing through the body of the valve.

SUMMARY OF THE INVENTION

[0006] According to the present invention there is provided a positive respiratory pressure apparatus having the features of claim 1.

[0007] The apparatus of the invention is capable of performing positive expiratory pressure (PEP) therapy alone or in combination with providing a substance, generally in aerosol form. As used herein, the term respiratory is intended to encompass both inhalation and exhalation. Whether inhalation resistance or exhalation resistance is called for will be known to one skilled in the art. The valve may be located at or near the output end of an aerosol delivery apparatus. The aerosol delivery apparatus with the PP apparatus may be used alone or in combination with a mask or mouthpiece.

[0008] In one embodiment, the PP apparatus is associated with a mask. The mask with the PP apparatus may be used alone or in combination with an aerosol delivery apparatus. In another embodiment, the PP apparatus is associated with a mouthpiece. The mouthpiece with the PP apparatus may be used alone or in combination with an aerosol delivery apparatus. In a further embodiment, the PP apparatus is associated with a nebulizer. The nebulizer with the PP apparatus may be used alone or in combination with a patient respiratory system interface, such as a mask or mouthpiece.

[0009] A method of performing positive expiratory pressure therapy includes providing a PP apparatus with a valve that is capable of providing a continuously variable expiratory window. The method further includes performing a series of breaths. When exhalation is performed, the exhalant is directed through the continuously variable expiratory window. Performance of a therapeutic cough triggers the loosening of secretions. Upon loosening of the secretions, a substance, such as a medicament, may be provided for inhalation into the respiratory system. In an alternative the PP valve may be positioned so as to provide positive inspiratory pressure upon inhalation into the apparatus.

[0010] A further aspect of another embodiment includes association of a PP apparatus associateable with a mask or mouthpiece engageable with a backpiece device. The backpiece device includes a plastic or an elastomeric adapter suited to receive the mouthpiece of a pressurized metered dose inhaler.

[0011] For performing positive respiratory pressure therapy in combination with providing an aerosolized medicament includes the apparatus an aerosol holding chamber having an input end and an output end.

[0012] The invention will best be understood by reference to the following detailed description of the preferred embodiment, taken in conjunction with the accompanying drawings. The discussion below is descriptive, illustrative and exemplary and is not to be taken as limiting the scope defined by any appended claims.
BRIEF DESCRIPTION OF SEVERAL VIEWS OF THE DRAWINGS

[0013]

Figure 1 is a cross sectional view of a mouthpiece associable with a chamber in conjunction with a PP apparatus.

Figure 2 is a perspective view of a mouthpiece associable with a chamber in conjunction with the PP apparatus.

Figure 3 is an exploded view of the preferred embodiment.

Figure 4a is a front view of one embodiment of the PP apparatus.

Figure 4b is a cross section drawn along line A-A of Figure 4a.

Figure 4c is a back view of one embodiment of the PP apparatus.

Figure 4d is a cross section drawn along line B-B of Figure 4a.

Figure 4e is a sectional cross section drawn along line C-C of Figure 4a.

Figure 4f is a rear perspective view of the embodiment of Figure 4a.

Figure 5 is a rear perspective of a mouthpiece according to a preferred embodiment.

Figure 6 is a front perspective view of a mouthpiece with one embodiment of the PP apparatus.

Figure 7a is a cross section drawn along line A-A of Figure 7a of the fitting and port.

Figure 7b is a cross section drawn along line A-A of Figure 7a of the fitting and port.

Figure 8a is a top view of one embodiment of the slide control.

Figure 8b is a side view of one embodiment of the slide control.

Figure 8c is a perspective view of one embodiment of the slide control.

Figure 8d is a cross section drawn along line A-A of Figure 8a.

Figure 9 is a top perspective view of one embodiment of the PP apparatus and a mouthpiece.

Figure 10 is a perspective view of an alternative embodiment of the PP apparatus of Figures 1-3 showing detent notches in conjunction with a mouthpiece.

Figure 11 is an exploded view of one embodiment showing a slide control having a port.

Figure 12 is one embodiment of the slide control having a port.

Figure 13 is one embodiment of the mouthpiece showing the annular sealing ring.

Figure 14 is one embodiment showing a plurality of detent notches.

Figure 15 is a side view of one embodiment of the control valve showing the port.

Figure 16 is a concave perspective view of the slide control showing the port.

Figure 17 is a concave bottom view of the slide control showing the port.

Figure 18 is an exploded view of an alternative embodiment of the PP apparatus of Figures 8-10.

Figure 19 is a cross-sectional view of the PP apparatus of Figure 18.

Figure 20 is a perspective view of a duck-bill valve used in the PP apparatus of Figures 18-19.

Figure 21 is a cross-sectional view of the duck-bill valve of Figure 20.

Figure 22a is a front exploded view of one embodiment of the PP apparatus in conjunction with a mouthpiece and associable with a spacer.

Figure 22b is a rear exploded view of one embodiment of the PP apparatus in conjunction with a mouthpiece and associable with a spacer.

Figure 22c is a front view of one embodiment of a valve showing the baffle.

Figure 23 is a perspective view of one embodiment of the PP apparatus in association with a nebulizer.

Figure 24 is an exploded view of one embodiment of the PP apparatus and a mouthpiece.

Figure 25 is a perspective view of one embodiment of the PP apparatus in an open position and a mouthpiece.

Figure 26 is a perspective view of one embodiment of the PP apparatus in a semi-open position and a mouthpiece.

Figure 27 is an exploded view of one embodiment of the PP apparatus showing the disc and a mouthpiece.

Figure 28a is a top view of one embodiment of the PP apparatus and a mouthpiece.

Figure 28b is a cross section of one embodiment of the PP apparatus and a mouthpiece showing a plurality of prongs holding the PP apparatus.

Figure 29 is an exploded view of a PP apparatus associated with a mouthpiece and having an inhalation valve.

Figure 30 is an exploded view of a PP apparatus associated with a mouthpiece and having an exhalation valve.

Figure 31 is a further perspective view of one embodiment of the PP apparatus in conjunction with a mask having an opening for association with a chamber.

Figure 32a is a close up of one embodiment of the PP apparatus in a fully open position in conjunction with a mask.

Figure 32b is a cross section of one embodiment of the PP apparatus in a fully open position in conjunction with a mask.

Figure 33a is a close up of one embodiment of the PP apparatus in a partially open position in conjunction with a mask.

Figure 33b is a cross section of one embodiment of the PP apparatus in a partially open position in con-
juncture with a mask.

Figure 34a is a close up of one embodiment of the PP apparatus having a plurality of variable sized flow ports, in conjunction with a mask.

Figure 34b is a cross section of one embodiment of the PP apparatus having a plurality of variable sized flow ports, in conjunction with a mask.

Figure 35a is a front exploded view of a close up of one embodiment of the PP apparatus having a plurality of variable sized flow ports, in conjunction with a mask.

Figure 35b is a rear exploded view of a close up of one embodiment of the PP apparatus having a plurality of variable sized flow ports, in conjunction with a mask.

Figure 36 is a front view of one embodiment of the PP apparatus showing resistance setting indicia.

Figure 37 shows a perspective view of a spacer for the aerosol delivery apparatus or the patient respiratory system interface. The assembly 10 combines the aerosol delivery apparatus, mouthpiece, and PP valve into a tool for use in both positive expiratory therapy and asthma management. The assembly 10 includes an aerosol delivery apparatus, such as, but not limited to, an aerosol holding chamber 14, and a patient respiratory system interface, such as a mouthpiece 16 and/or mask attachable to the output end of the aerosol delivery apparatus. The PP valve 12 may be located anywhere on the mouthpiece 16. Although, it is understood that the aerosol holding chamber 14, but prevents exhalation back through the chamber. As illustrated in Figures 1-3, the aerosol holding chamber 14 is provided with an annular valve 24 located at its outlet end 22. The annular valve 24 allows the user to inhale medicament from the aerosol holding chamber 14, but prevents exhalation back through the chamber. As illustrated in Figures 1-3, and in more detail in Figures 4a-4f, 5 and 7, a PP valve 12 may be formed in a mouthpiece 16. The PP valve 12 includes a slide control 26 that is movably positioned relative to a resistance window 28. The slide control 26 is variably maneuverable to cover or uncover the resistance window 28 in a continuous manner. Further, the movement of the slide control 26 includes, but is not limited to, covering or uncovering, and/or opening or closing, the resistance window 28 or any variations thereof.

[0017] The PP valve 12 may be located on or in conjunction with a mouthpiece 16. An exemplary embodiment of the mouthpiece 16 shown in Figures 4a-4f has a distal end 30 and a proximal end 32. Commonly, the proximal end 32 of the mouthpiece is inserted or associated with the mouth or nostrils of the user. Additionally, the distal end 30 of a mouthpiece may or may not be associated with an aerosol delivery apparatus and the mouthpiece alone may be configured to constitute a PEP device.

[0018] Generally, in one exemplary embodiment, the PP valve 12 may be located at or near the distal end 30 of the mouthpiece 16. Although, it is understood that the PP valve 14 may be located anywhere on the mouthpiece 1 and its location is not to be limited. In an alternative embodiment, the PP valve 12 may be located at or near the outlet end 20 of the aerosol delivery apparatus, such as, but not limited to, the aerosol holding chamber 14 of Figures 1-3. Generally, the direction of travel of any fluid, particularly an aerosol or nebulizer medicament, is in the direction from the input end 20, through the channel or chamber body, and to or out the output end 22. This direction of travel from input end 20 to output end 22 is referred to as travel from downstream to upstream.

[0019] In a preferred embodiment the mouthpiece 16 is formed of plastic. The plastic may be either rigid or soft. Other materials that can also be used for the mouthpiece 16 include metal or other materials known to one in the art. In the embodiment depicted in Figures 1-3, 4a, 4b, 4c, and 4f, a tab 34 is provided allowing for the connection of a cap 36, as shown in Figure 2, to cover the proximal end 32 of the mouthpiece 16. In a preferred embodiment, the mouthpiece 16 may include indicia 35, or setting indications, representing the resistance setting. The indicia 35 may be in the form of num-
bers, bars, colors, a series of dots or the like.

[0020] In a further embodiment, as shown in Figures 7a and 7b, the PP apparatus 10 may include a fitting 39 sized for placement over the proximal end of the mouthpiece. The fitting 39 includes a manometer port 41 extending from the fitting over which a manometer can be attached. In a preferred embodiment the fitting 39 is formed of a plastic. The plastic may be either rigid or soft. Other materials that can also be used to form the fitting include metal.

[0021] Figures 3, 8a-8d and 9 show an embodiment of the slide control 26 of the PP apparatus 10. As shown in Figure 9, in the illustrated embodiment, the slide control 26 is of a semi-circular, quarter moon shape. The slide control 26 has a first lateral side 38 concave in shape and a second lateral side 40 opposite the first side 38. The slide control 26 also has a top 42 and a bottom 44 surface. From the top surface 42 of the slide control 26 extends a tab setting 46.

[0022] In the illustrated embodiment, the tab setting 46 is a uniformly molded projection from the slide control 26. In a preferred embodiment, the tab setting 46 has smooth edges for easy engagement with the finger, thumb or appendix of the user. The tab setting 46 may also have a serrated edge or any other edge known in the art. When assembled with the mouthpiece 16, the tab setting 46 projects through the mouthpiece from the tab window 48. The user of the device manipulates the tab setting 46 in such a manner as to cause, either directly or indirectly, the movement of the slide control 26 thereby varying the opening of the resistance window 28.

[0023] In the embodiment of Figures 1-3, 4a-4f, 5 and 7, the tab window 48 is arcuate in shape, parallel to the contour of the circumference of the mouthpiece 16. Referring to Figure 4f, the slide control 26 is preferably seated in a channel 50 located on the mouthpiece 1. The slide control 26 is held within the channel 50 by at least one tooth 52. Located on one or both of the walls of the channel 50 is a stepped surface 54 as shown in Figure 5.

[0024] In the embodiment shown in Figures 8a and 6, the control arm 56 of the slide control 26 is shown having a finger projection 58 from one end of the slide control 26. In another embodiment, the slide control 26 may have a control arm 56 located on both ends of the slide control 26. The finger projection 58 is capable of engagement with the stepped surface 54 inside the mouthpiece shown in Figures 4c and 5. In the illustrated embodiment, the stepped surface 54 includes a series of ribs extending a variable length of the internal diameter of the mouthpiece 1. In another embodiment, as shown in Figure 30, the stepped surface 54 may also be located along the tab window 48. Therefore, the location of the stepped surface 54 may vary while remaining engageable by the control arm 56. Additionally, the stepped surface 54 may be located on either or both of the internal walls of the channel 46. In the embodiment of Figures 4a-4f and 8a-8d, the slide control 26 is of a flexible material so that the control arm 56 can slide across the uppermost surface of the ribs projecting from the stepped surface 54. When the desired opening of the resistance window 28 is obtained, the control arm 56 engages in a semi-locked manner the ribs projecting from the stepped surface 54.

[0025] The end of the slide control 26 opposite the control arm 56 may either be provided with a finger projection 58 or may be smooth. The length of the slide control 26 extending from the tab setting 46 to the end of the control arm 56 opposite the projection 58 is generally the length of the resistance window 28. This resistance control length 60 is at least the length that the resistance window 28 can be opened allowing for exhalant to exit the window 28. In a preferred embodiment, the slide control 26 is manufactured of a plastic. The plastic may be either rigid or soft. Other materials that can also be used for the slide control 26 include metal or other materials known in the art.

[0026] In general, as shown in Figure 5, the resistance window 28 may be an opening of any size or shape in the walls defining the channel 46 in the mouthpiece 16 which, in conjunction with the illustrated embodiment of the slide control 26, provides an opening in the mouthpiece 16 to produce sufficient pressure during exhalation of the patient performing PEP therapy. For example, the resistance window 28 may be formed with straight or slanted edges. If the edges are slanted, this provides a steeper effect to the resistance window 28. If the desired exhalation pressure is determined to range from 10-20 cm H2O, then the resistance window 28 in conjunction with the slide control 26, acting as a cover or closure mechanism for the resistance window 28, are sized in such a manner as to provide an appropriate opening for the desired exhalation pressure to be produced. As one example, if the resistance window 28 is generally narrow, then the length of the window may be of a longer length so as to provide a large enough opening through which PEP therapy is performed. Interdependent in the relationship is the resistance window length 60 of the control arm 26. In the above example, the resistance window length 60 of the control arm 26 is generally longer to cover the desired amount of the resistance window 28. The control arm 26 may provide a continuously adjustable variable resistance window between a first position where the control arm completely blocks the window 28, to a second position where the control arm leaves the window completely open.

[0027] An alternative embodiment of an assembly 100 for performing PEP therapy is shown in Figures 10-17. This embodiment is similar to the embodiment of Figures 11-13, but utilizes a variation of the resistance window and slide control in the PP valve 112. In the assembly 100 of Figures 10-17, the resistance window 128 and slide control 126 are positioned in the mouthpiece 116. The tab setting 146 of the slide control 126 is of a flexible material. A detent 158 protrudes from the first
tab setting face 156 located on the concave side 125 of the slide control 126. As shown in Figure 15, a position indicating rib 159 is located on the tab setting face 156 opposite the detent 158. In the illustrated embodiment, the rib 159 is shown as a generally rectangular protrusion from the surface of the tab setting 146. Yet, the rib 159 may be any shape protrusion, such as but not limited to circular, triangular. Further, the rib 159 may not be a protrusion at all but rather is a concave marking on the surface of the tab setting 146. The rib 159 has at least the function of indicating to the user the extent of the opening of the resistance window 128. Therefore, one skilled in the art can envision a variety of marking, shapes, indents or protrusions, or colors which serve at least the function of indicating the extent of the opening of the resistance window 28.

0028 The detent 158 located on the tab setting 146 is associated with at least one detent notch 154 as shown in Figures 10 and 14. The flexible tab setting 146 is movable within the tab window 148. The one or more detent notches 154 are preferably located along the boundary of the tab window 148 and indicia 135, representative of exhalation effort corresponding to the position of the slide control 126, are arranged adjacent the respective detent positions. Figures 10 and 14 show a plurality of detent notches 154. In operation, the flexible tab setting 146 is moved along the tab window 148. The detent 154 located on the first tab fitting face 125 of the tab setting 146 moves into and out of engagement with the detent notches 154. Movement along the boundary and engagement with the detent notches 154 removably fixes the slide control 126 in a variety of positions. Each varied position provides for a further opening or closing of the resistance window 128. Further, in operation, the detent 158 is not limited to being engaged with a detent notch but may be engaged or seated at any point along the boundary. Engagement with, or seating within, a detent notch 154 of a detent 158 provides for a variable securely fixed opening of the resistance window 128. Each detent notch 154 may correspond to a particular size opening or pre-set opening of the resistance window 128. Therefore, by engagement of the detent 158 within the detent notch 154, the user may be provided with a pre-set resistance window opening 128. Yet, the detent notch 154 may also be positioned anywhere along the boundary providing for a continuously variable resistance window opening 128.

0029 The slide control 126, as shown in Figures 15, 16 and 17 has located therein a port 143. The port 143 may be of any size or shape and in the illustrated embodiment is generally a rounded triangle having an elongated point. The port 143 operates in conjunction with the resistance window 128. In the embodiment illustrated in Figures 10-17, the size of the opening through which the exhalant passes is determined by how much of the port 143 is left uncovered or open and aligned with the resistance window 128. The slide control 126 is provided with a closure area 145, as shown in Figure 16. Further, the slide control 126 is provided additionally with a port area 160. When the resistance window 128 is closed or not open and thereby not allowing for the exiting of any exhalant, the closure area 145 of the slide control 126 is congruent with or aligned with the resistance window 128. As the slide control 126 is gradually moved, the port area 160 containing the port 143 is brought into alignment with the resistance window 128 in the mouthpiece 116. In this manner, a continuously variable opening is provided. For example, as the slide control 126 moves aligning a greater and greater amount of the port 143 with the resistance window 128, a greater opening or path for the exhalant is provided.

0030 As described above, and similar to the embodiment of Figures 1-3, the slide control 126 is seated in a channel 150 located on the mouthpiece 116. The slide control 126 is slidably moveable within the channel 150 in the manner described above so as to continuously variably align the port 160 with the resistance window 128. In the embodiment of the slide control 126 illustrated in Figures 13, 16, and 17, a retaining ridge 162 fits into the channel 150 thereby holding the slide control 126 in its desired position throughout its range of motion in the mouthpiece. The desired position of the slide control 126 is as close as possible to the annular sealing ring 164 shown in Figure 13. One function of the annular sealing ring 164 is to prevent leakage of the exhalant to ensure that the exhalant to the greatest extent exits from the resistance window 128. In a further embodiment, the PP valve 112 may optionally be provided with at least one locating ring or post 166 to help maintain the slide control 126 in alignment. As with the embodiment of Figures 1-3, a patient's exhalation effort is controlled by adjusting the slide control over the resistance window so that the patient's exhaled air, which is prevented from entering the aerosol chamber 114 by the annular valve 124 and a baffle (not shown), must exit through the resistance window and provide a desired exhalation resistance. More specifically, during inhalation the inner diameter 127 of the annular valve 124 (Figure 11) unseats from the baffle (not shown) on the output end 122 of the aerosol holding chamber and permits passage of fluid. Substantially simultaneously, the exhalation flange 186 of the valve 124 flexes to seal against the outer ridge 188 (Figure 13) formed inside the mouthpiece 116 to prevent ambient air from entering the mouthpiece. During exhalation, the process is reversed and the inner diameter 127 prevents exhalant from entering the aerosol holding chamber while the exhalation flange 186 flexes away from the outer ridge 188. Thus, the exhalant preferably passes through the resistance window 128 and may escape to the outside between the exhalation flange and the mouthpiece, and then through the gap between the mouthpiece and aerosol holding chamber.

0031 Referring to Figures 18-21, an alternative embodiment of an assembly 210 for PEP therapy combined with an aerosol delivery apparatus is shown. The aerosol delivery apparatus includes a chamber housing 214.
having an input end 220 and an output end 222. The chamber housing 214, input end 220, and output end 222 define an interior space 223. The apparatus may also include an elastomeric backpiece which may be similar to the backpiece in the embodiment shown in Figures 1-3. The output end 222 of the chamber housing 214 is shaped to receive the mouthpiece 216 and includes locking tabs 281 and a protrusion 283. The protrusion 283 is preferably annular in shape. The locking tabs 281 are spaced apart around the outside of the output end 222.

[0032] Referring to Figures 18-19, the mouthpiece assembly 216 is preferably substantially similar to that illustrated in Figures 10-15. The mouthpiece assembly includes an annular sealing ring 224, and a resistance window defined by a gap 225 in the sealing ring. A slide control 226 is slidably seated in a channel 250 between the annular sealing ring and support posts or a support ring. The tab setting 246 protrudes from the tab window 225 over the locking tabs 281. As with either embodiment of slide control and nozzle description above, using either embodiment of slide control and nozzle described above.

[0033] The mouthpiece assembly 216 is connected to the output end 222 of the chamber housing 214 by placing the apertures 284 over the locking tabs 281. As with the embodiment of Figures 1-3 and 10-17, in the embodiment of Figures 18-21 a containment baffle 275 may be integrally formed with the chamber housing 214 from a single piece of material and located near the output end 222. As shown in Figure 18, the containment baffle 275 includes connecting members 276 that extend from the edge of the containment baffle to the inner diameter of the output end 222 in the chamber housing. Vents 277 are defined between the outer perimeter of the containment baffle and the inner diameter of the outlet end of the chamber housing and are separated by the connecting members. The vents 277 are arcuate in shape and conform to the outer perimeter of the containment baffle 275. In a preferred embodiment, the containment baffle is dome-shaped where the concave end points towards the chamber 223. In alternative embodiments, the containment baffle may be any of a number of geometric shapes.

[0034] A valve 278 having a valve member 279 and a valve seat 280 is shown. The valve seat 280 preferably comprises the rim of the valve and the corresponding raised lip 283 on the outlet end 222 of the chamber housing 214. The valve member has a sealing surface that preferably forms between two parallel portions, or lips, of the valve. In a preferred embodiment, the valve material seals against itself when fluid flows against a predetermined flow direction of the valve. In the embodiment shown, the valve is a duck-bill valve where the valve seat 280 is positioned axially away from the valve opening. The valve 278 defines a central open area toward the end having the valve seat. The valve member 279, shown as parallel sealing lips in Figures 18-21, acts to allow passage of fluid on inhalation, but upon exhalation, the lips of the valve member are held together by the force of fluid (e.g. exhaled air) pressing against the walls 285 of the valve member and collapsing the lips of the valve member 279 against each other in a closed position. The valve seat 280 also provides a seal against the chamber housing 214 during exhalation so that exhalant from a patient must be directed through the continuously variable resistance window in the mouthpiece.

[0035] In one preferred embodiment, the duck-bill valve 278 has a central open area 274 at its base that has a diameter of approximately 26.09 millimeters (mm). The width of the lips that form the valve member 279 is approximately 21.35 mm and the angle at which the walls 285 converge is approximately 72 degrees. Also, the height of the duck-bill valve 278 measured from the upper portion 282 of the valve seat to the valve member 279 is approximately 18.8 mm. The mouthpiece 216 for containing this valve 278 preferably includes a resistance window gap 225 having a length of 59 degrees of arcuate cut in the annular sealing ring 224, where the annular sealing ring is approximately 31.4 mm in diameter and has a height of 4.5 mm.

[0036] The operation of the apparatus will now be discussed generally with reference to the embodiments of Figures 1-3, 10-17 and 18-21. At rest, the valve is adjacent to the output end of the chamber housing. In the annular valve embodiment of Figures 1-3 and 10-17 the inner portion of the valve covers the vents at the outlet end of the chamber housing. In the duck-bill embodiment of Figures 18-21, the entire outlet end is covered by the valve. For either valve embodiment, the mouthpiece and outlet end of the chamber housing traps the valve in place. Inhalation by the patient causes the sealing portion of the annular valve to move, or alternatively the lips of the duckbill valve to separate, and permit fluid to pass. Fluid from the chamber housing may be inhaled into the patient’s respiratory system through the mouthpiece. The patient may then exhale into the mouthpiece.

[0037] Exhalation by the patient results in air traveling through the mouthpiece in a direction opposite the predetermined inhalation flow path of the valve. This air, which is blocked from passage through the valve along the inhalation path, then passes along a second path through the resistance window in the mouthpiece. Also, the force of the exhaled air causes the outer portion of the valve to move away from the mouthpiece in a direction towards the chamber housing. As a result, an exhalation pathway is created between the outer portion of the valve and the mouthpiece through which the exhaled air passes out to the atmosphere or some other predetermined location. As described above, the amount of effort that exhalation requires is set by the slide control, which may be set to block the appropriate amount of the resistance window to achieve the desired resistance.

[0038] Referring specifically to the duck-bill valve embodiment of Figures 18-21, the duck-bill valve 278 as-
consists both in preventing inhalation of ambient air through the resistance window 225 and providing an exhalation path for exhalant. When the apparatus 210 is assembled (Figure 19), the upper portion 282 of the valve seat seals against the annular sealing ring 224 except for the gap in the annular sealing ring 224 defining the resistance window 225. During inhalation, fluid flows through the central opening 274 and out the lips of the valve member 279. Also, the exhalation flange 286 flexes toward the proximal end of the mouthpiece and provides a secondary seal against the outer ridge 288 inside the mouthpiece. The outer ridge 288 is preferably continuously around the inner circumference of the mouthpiece. Upon exhalation, the lips of the valve member 279 close, the exhalant passes through the resistance window under the upper portion 282 of the valve seat positioned adjacent the resistance window opening, and the exhalation flange flexes away from the outer ridge 288 and out between the aerosol holding chamber 214 and mouthpiece 216.

[0039] As shown in Figures 22a-22c, an embodiment of an apparatus 310 for performing PEP therapy is disclosed that may be used with or without an aerosol delivery apparatus. In contrast to the embodiments discussed above, the PEP apparatus of Figures 22a-22c is a standalone PEP device in a mouthpiece incorporating the previously discussed continuous variable resistance window with slide control, and variations thereof, along with a self contained valve 378 that may be similar to that disclosed in the previous embodiments. Thus, the valve 378 need not be found on the outlet end of a separate chamber extension 314 but may be positioned on the distal end 330 of the mouthpiece. As best shown in Figures 22b-22c, the valve 378 may be an annular valve. The 378 is preferably retained toward the distal end 330 of the apparatus by a central baffle 385 supported by radial spokes 386. In one embodiment, the PEP apparatus 310 is formed of an attachable mouthpiece section 316 and a baffle section 317. The mouthpiece and baffle sections 316, 317 may be removably joined using snap-fit, threaded or other known attachment schemes. In another alternative embodiment, the mouthpiece and baffle sections 316, 317 may be integrally molded or welded shut to form a non-removable, unitary piece. An extended inlet 314, without any valves, may be used with the stand-alone PEP device 310 to enhance delivery of any medicine to the patient’s respiratory system. One function of the extended inlet is to provide a chamber for the dispersed particles from the pressurized metered dose inhaler. When desired, a pressurized metered dose inhaler may be coupled to the extended inlet with a backpiece and medicament supplied from the pressurized metered dose inhaler can be delivered directly to the user.

[0040] Although the embodiments of Figures 1-22 illustrate annular and duck-bill valves 24, 124, 278, 378, any of a number of other valve configurations may be used. A preferred valve is capable of passing a fluid moving in a first direction along a first path and also capable of passing a fluid moving in an opposite direction along a second path. In the example valves discussed above, inhalation draws fluid through a central opening in the valve while the perimeter of the valve prevents fluid flow. In the above examples, exhalation closes the path through the central opening and directs fluid along a second path around the perimeter of the valve. Other paths may also be used.

[0041] Figures 23-28 illustrate another embodiment of a PEP apparatus 410. As best shown in Figure 23, the PEP apparatus 410 has a patient respiratory system interface, such as a mouthpiece 416, on a proximal end 432 and may be connected with an aerosol delivery apparatus, in this example a nebulizer 414, at a distal end 430. The PEP apparatus includes a PP valve 412 positioned on top of the mouthpiece 416. The PP valve 412 preferably consists of a cover 417 that may be removable attached to a receiving area 419 (Figure 24) on the mouthpiece 416. The cover has a resistance window 428 and a tab window 448 extending through a top surface. A slide control 426 in the form of a disk with vents 460 extending through the thickness of the disk is movably positioned under the cover 417. A one-way valve 423 is positioned between the slide control 426 and the top of the mouthpiece 416 to allow air exhaled into the proximal end 432 to escape through the resistance window 428 while preventing any air from entering through the resistance window during inhalation.

[0042] In Figures 23-28, the resistance window 428 is shown generally as a pie-slice shaped cut-out with the point of the pie-slice removed so as to form a concave edge. The resistance window 428 may be any shape and should not be limited by the illustrated embodiment. Further, a plurality of resistance windows 428 may form the PP valve 12. The number of windows 428 is not intended to be limited by the illustrated embodiment. In the illustrated embodiment of Figures 23-28, and particularly Figure 27, the slide control 426 is shown as a circular disc having a pie-slice shaped cut-outs with the point of the pie-slice removed wherein which correspond to the openings of the resistance window 428. In operation, aligning the openings 460 of the disc with the resistance window 428 controls the opening of the continuously variable resistance window 428. When the resistance windows 428 are aligned with the disc openings 460, the resistance windows 428 are opened to their fullest extent allowing the resistance of the exhalant exiting the PP valve 12 to be lower. When only a small amount of the resistance windows 428 are aligned with the disc openings 460, the resistance of the exhalant exiting the PEP apparatus 410 is increased. By moving the tab setting 446 in the tab window 448, the vents may be adjusted in the disk 417 to any of a number of positions, thereby providing a continuously adjustable resistance. In this manner, positive expiratory pressure is controlled.

[0043] Referring again to Figure 23, where the PEP
apparatus 410 is connected at its distal end to the nebulizer 414, the operation of this embodiment will be described. Upon inhalation, the nebulizer will provide an aerosol to the inhaling patient via the mouthpiece. A suitable nebulizer for use with the PEP apparatus 410 is a breath-actuated nebulizer such as disclosed in U.S. Patent No. 6,044,841 issued April 4, 2000 and entitled "Breath Actuated Nebulizer with Valve Assembly Having Relief Piston", the entirety of which is incorporated herein by reference. During inhalation from the nebulizer 414, a piston 452 is drawn down by negative pressure created by the inhalation in the nebulizer and ambient air is drawn through openings 454 in the lid 456 of the nebulizer 414. The one-way valve 459 in the PP valve assembly 412 remains shut during inhalation.

[0044] Upon exhalation into the proximal end 430 of the mouthpiece 416, a positive pressure builds in the nebulizer 414 and the piston acts as a one-way valve to close off the flow of air out of the nebulizer. Now, the exhalant must travel through the one-way valve in the PP valve assembly 412, through the slide control and out the resistance window. Preferably the slide control 426 under the resistance window 428 has been set to the appropriate position for the patient so that effective PEP therapy may be provided. Although the PEP apparatus of Figures 23-27 uses an aerosol delivery apparatus such as the nebulizer 414 to restrict air flow through any opening other than the PP valve assembly 412, other embodiments, such as shown in Figure 29 discussed below, are contemplated where a second one-way valve is associated with the distal end 432 of the PEP apparatus 410 so that the PEP apparatus may be used in a standalone fashion for PEP therapy. The illustrated embodiment of Figures 23-28 show an improved nebulizer 414 associated with a PEP apparatus 410 having a PP valve 410 and the disc 56 in order to allow the exhalation valve 402 to close off the flow of air out of the nebulizer. In this manner, the mouthpiece 416, the operation of this embodiment will be described. Upon inhalation, the nebulizer will provide an aerosol to the inhaling patient via the mouthpiece. A suitable nebulizer for use with the PEP apparatus 410 is a breath-actuated nebulizer such as disclosed in U.S. Patent No. 6,044,841 issued April 4, 2000 and entitled "Breath Actuated Nebulizer with Valve Assembly Having Relief Piston", the entirety of which is incorporated herein by reference. During inhalation from the nebulizer 414, a piston 452 is drawn down by negative pressure created by the inhalation in the nebulizer and ambient air is drawn through openings 454 in the lid 456 of the nebulizer 414. The one-way valve 459 in the PP valve assembly 412 remains shut during inhalation.

[0046] Although positive expiratory devices have been shown in detail, embodiments of positive inspiratory devices are also contemplated. Figure 30 shows one embodiment, similar in concept to the embodiment of Figure 29, but with the PP valve 475 attached in series with the one-way inhalation valve 476, rather than in series with the one-way exhalation valve, to provide for resistance upon inhalation only. The arrows drawn in Figure 30 depict the direction of travel from the downstream end of the mouthpiece 478 to the upstream end of the mouthpiece showing that all inhalation must pass through the continuous variable resistance window 480 and the port 482 of the slide control 484. If desired, in other embodiments PP valves may be placed in series with both the one-way input and one-way output valves to allow for simultaneous control of positive inspiratory and expiratory pressures at the same or different levels.

[0047] As discussed above, embodiments of patient respiratory system interfaces aside from the mouthpiece configurations already disclosed are contemplated. A PP apparatus 510 utilizing a mask 512 as the interface is illustrated in Figures 31-35. The mask may be a standard mask sized for adults or children and constructed of any of a number of materials such as silicon rubber. The mask 512 may have a frustoconical shaped main section 514 sized to cover the patient's mouth and a nose piece section 516 sized to cover a patient's nose. A central opening 518 in the mask 512 may be used to attach with an aerosol delivery apparatus such as the aerosol holding chamber 14 shown in Figures 1-3, and other aerosol delivery apparatus. Alternatively, the mask 512 may be fitted with a one-way valve in the central opening 518 for use as either a positive expiratory pressure device or a positive inspiratory pressure device. As with the embodiment of Figures 23-28 and 29-30, a PP valve assembly 520 is positioned on the device so that inspiration and exhalation paths travel off-axis from one another. The PP valve assembly 520 has an adjustable valve assembly cover 534 with tab window 522 and resistance window 528 openings positioned on it. The resistance window 528 is generally an oblong, tear-drop shape and the tab window 522 defines an arcuate opening in the PP valve cover 534. The tab window and resistance window may alternatively be rectangular, oval or any other shape. Although the above embodiments illustrate a tab window 522 located approximately on a top surface of a mouthpiece or on top of a mask positioned approximately adjacent the nose, the tab window 522 may be located anywhere on the mouthpiece or mask.

[0048] Referring to Figures 35a and 35b, the PP valve
assembly 520 has a fixed opening 530 and a set of detents 532 positioned on a disk-shaped platform 526 that connects to the nosepiece section of the mask through complementary tab 542 and slot 544 connectors. As best shown in Figure 35b, the PP valve cover 534 has a protrusion 536 sized to cooperate with the detents 532 on the platform 526 so that the valve cover may be moved to predetermined spots when the valve cover is rotated against the platform. An axle 538 on the valve cover fits into a central opening 540 in the platform 526 so that the resistance window 528 is rotatably positionable over the exhalation port 530 and the tab window 522 lines up with the tab extending from the platform 526.

This embodiment depicts the resistance window 528 as a curved tear-drop like shape. The platform 526 is shown as a circular disc having at least one port opening 530. The port opening 530 may vary in size and shape. The opening formed for the exhalant to pass through is related to the alignment of the resistance window 528 with the port opening 530. In this embodiment, the resistance window is moveably mounted relative to a fixed slide control portion attached to the mask. Tabs 542 on the platform 526 preferably mate with tab receiving regions 544 on the end of the nosepiece extension 516 to retain the platform in a fixed position relative to the mask. Moving the tear-drop shaped resistance window 528 past the port opening 530 varies the exhalant path. In other embodiments, a plurality of resistance window openings 528 may be moved past the port 530. Alternatively, there may be a plurality of ports in the slide control 526.

As shown in Figures 31 and 32a, one end of the tear-drop shaped resistance window 528 matches the size of the largest port opening 530 at a maximum flow position thereby providing a maximum flow and least resistance in that position. When the valve cover is rotated so that the resistance window 528 covers a greater portion of the port, as shown in Figures 33a-33b, a smaller exhalant path is created providing greater resistance. As shown in Figures 34a-34b, moving the valve cover until the tab reaches the opposite end of the tab window results in the smallest amount of the port being open, the highest airflow resistance and the least flow. It is envisioned that a plurality of size and shape port openings and resistance windows may be used and the disclosure is not to be limited to that depicted in the drawings. Referring to Figure 36, an embodiment is shown of a valve cover 550 having indicia 552 representative of a resistance setting. The indicia 552 are arranged to cooperate with the tab extension 554 on the platform to indicate the current resistance setting.

In alternative embodiments, PEP therapy may be performed with a mouthpiece or mask having the PP valve associated with a backpiece. The mask or mouthpiece may have an extended inlet for association with the backpiece.

Asthmatic medications are commonly supplied in metered dose inhalers, frequently referred to as pressurized metered dose inhalers. Pressurized metered dose inhalers are generally cylindrical canisters with axially extending vent tubes from internal valves. When the external tube or stem of a pressurized metered dose inhaler canister is depressed it operates the internal valve to dispense a measured dose of medicine from the stem. The medicine is commonly packed in the canister with a suitable compressed gas to propel the medicine and gas from the stem or tube when the later is depressed. The medicine may be in gas, liquid, or solid form. The manufacturer or distributor of the pressurized metered dose inhaler canister generally supplies it with a substantially L-shaped adapter which receives the canister in a substantially upright position, and has a substantial horizontal outlet portion for reception in the mouth of an asthmatic patient for inhalation of the medicine.

In order to address the problem of coordination and other problems known in the art with regard to pressurized metered dose inhalers, a spacer chamber with an integrated actuator, or an aerosol holding chamber, have been used in attempts to overcome inappropriate particle size. The aerosol holding chamber is generally provided at the upstream or entering end with a flexible, resilient adapter or backpiece made of rubber or the like material. A central aperture is provided for receipt of the horizontal outlet portion of the pressurized metered dose inhaler adapter.

One embodiment provides for an improved pressurized metered dose inhaler or pressurized metered dose inhaler with an aerosol holding chamber. As shown in Figure 37, a PP apparatus 600 may be associated with the pressurized metered dose inhaler or the pressurized metered dose inhaler with an aerosol holding chamber. In the PP apparatus 600 of Figures 37-39, an L-shaped adapter portion 602 holds the pressurized canister and a horizontal outlet section 604 receives the medicament released in aerosol form. A one-way valve 606, which may be a flexible membrane, a rigid membrane, hinged door, or other commonly known valve mechanism is positioned at the proximal end 608 of the horizontal outlet section 604. To provide the positive expiratory pressure, the one-way valve 606 permits inhalation and blocks exhalation so that substantially all exhalation is routed through the variable resistance window 610 adjacent the one-way valve 606. A slide control 612 is movable in the resistance window 610 by a tab 614 to close off or open up as much of the resistance window as necessary to provide the desired expiratory pressure. Figures 38a and 39a illustrate the slide control in a completely open position and Figures 38b and 39b illustrate the slide control closing off the resistance window. The slide control may maintain its position in the resistance window through friction, detents or other known mechanisms for mechanically retaining one of multiple desired positions. The proximal end 608 of the metered dose inhaler 600 with PP functionality may be
used by a patient directly or fitted to an adapter on an aerosol chamber such as shown in Figure 3. Figure 40 illustrates another embodiment of a pressurized metered does inhaler 620 with a round proximal end 628 that may be used without the need for special mouthpieces or aerosol holding chambers. As with the embodiment of Figures 37-39, the alternative PEP enabled pressurized metered does inhaler 620 has a one-way valve 626 that shunts exhalant through a resistance window 622 that is continuously adjustable with a slide control 624 that can adjust the aperture of the resistance window.

[0055] Generally, a mouthpiece or mask may be associated the PP apparatus. In one configuration, an aerosol holding chamber may be attached to the mouthpiece or mask end and a metered dose inhaler may be positioned on a generally opposite end of the chamber via a backpiece. The user of the device may insert the mouthpiece into the mouth to obtain a dose of medication. Further, the user may place the mask over the mouth and/or nose to inspire a dose of the medicament. In either situation, the mask or mouthpiece aids in the delivery of the medicament to the user.

[0056] As has been described, a method and apparatus from providing positive expiration or inhalation therapy, with or without separate aerosol generating devices, has been disclosed. In the embodiment where the positive expiratory pressure valve is located at or near the output end of the aerosol delivery apparatus, a one way inhalation valve can be located further downstream from the positive expiratory pressure valve. A mouthpiece and or mask can be affixed at or near the output end of the aerosol delivery apparatus. The positioning of the inhalation valve either upstream or downstream in respect to the positive expiratory pressure valve is well known to one skilled in the art. Further, it is envisioned that PEP therapy may be performed nasally with the positive expiratory pressure apparatus.

[0057] When the mouthpiece having the PP apparatus associated therewith is used alone to perform PEP therapy, and not in conjunction with a mechanism for the delivery of a substance, a one way inhalation valve is engageable with the mouthpiece. The inhalation valve functions so as to allow for inhalation by the patient into the mouthpiece. The exhalant of the patient is prevented from exiting via the inhalation valve and is directed to exit through the PP valve. Generally, an inhalation valve opens upon inhalation to allow a fluid, such as an aerosol, to enter a chamber or channel or the like but that closes upon exhalation to prevent exhaled fluids to enter into the chamber of the like. The drawings depict an exemplary embodiment of the one-way inhalation valve but are not to be limiting to the embodiments shown.

[0058] One aspect of the method of use of the PP apparatus can be understood by the following disclosure and reference to Figures 1-3, 5 and 9. Particularly, the arrow 2 in Figure 1 indicates the direction of flow of the exhalant. The one-way valve shunts exhalant out between the mouthpiece and the aerosol chamber via the continuously variable resistance window. In carrying out the method, a physician may initially determine the proper resistance setting of the PP apparatus according to the patient's requirements. One manner in which the PP apparatus may be properly set is by attaching a fitting 39 to the mouthpiece. A manometer is then attached to the fitting port 41 and serves to measure the expiratory pressure. A patient will exhale into the mouthpiece and the pressure can be read from the manometer. The physician can then move the tab to one of the desired settings indicated on the mouthpiece. Once the proper resistance has been determined the fitting 39 can be removed from the mouthpiece. This fitting 39 will not be used again unless it is determined that the resistance should be adjusted.

[0059] The method of performing PEP therapy using the PP apparatus includes performing a series of breaths. When exhalation is performed, the exhalant is directed through the continuously variable expiratory window. Performance of a therapeutic cough triggers the loosening of secretions. Upon loosening of the secretions, a medicament may be provided for inhalation into the respiratory system. In one embodiment of PEP therapy, the user will exhale into the mouthpiece and/or mask, against the desired resistance. This is done either prior to or in combination with inhalation of the medicament. The exhaled gases exit through the resistance window. This process may be repeated as many times as prescribed by the patient's physician.

[0060] As has been described, a method and apparatus for providing positive expiration, or inhalation, pressure therapy, with or without separate aerosol generating devices, has been disclosed. The aerosol delivery apparatus with the PP apparatus may be used alone or in combination with a mask or mouthpiece. Also, an improved aerosol delivery apparatus with an integrated actuator has been shown, wherein the improvement comprises a PP valve. The discussion above is descriptive, illustrative and exemplary and is not to be taken as limiting the scope defined by any appended claims.

Claims

1. A positive respiratory pressure apparatus (10, 100, 210, 310, 410, 510, 600, 610) comprising:

   a patient respiratory system interface (16, 116, 216, 316, 416, 462, 512); and

   a valve assembly in fluid communication with the patient respiratory system interface, the valve assembly comprising:

   a valve (12, 112, 278, 378, 412, 467, 475, 520, 606, 626) configured to pass a fluid traveling in a predetermined direction from
a first side to a second side of the valve;
a variable resistance bypass window (28, 128, 225, 428, 471, 480, 528, 610, 622) positioned adjacent the valve and having a resistance to a fluid traveling in a direction opposed to the predetermined direction, wherein the variable resistance bypass window is continuously adjustable between a first fluid resistance and a second fluid resistance; and characterised by

a one-way exhalation valve means (286, 423) providing an exhalation path and preventing inhalation along the exhalation path, wherein the exhalation path is different than the inhalation path.

2. The apparatus of claim 1, wherein the valve comprises a one-way inhalation valve portion (127, 279, 463, 476), the one-way inhalation valve portion providing an inhalation path and preventing exhalation along the inhalation path.

3. The apparatus of claim 2, wherein the one-way exhalation valve means is comprised in the valve.

4. The apparatus of claim 3, wherein the valve (278) comprises a duck-bill valve comprising an open central region and a central valve member, wherein the open central region and the central valve member define the inhalation path, and wherein the duck-bill valve further comprises a peripheral exhalation flange (286) defining the exhalation path.

5. The apparatus of claim 1, wherein the valve comprises a one-way valve.

6. The apparatus of claim 1, wherein the valve comprises a one-way inhalation valve.

7. The apparatus of claim 1, wherein the valve comprises a one-way exhalation valve.

8. The apparatus of claim 1, wherein the fluid traveling in the predetermined direction comprises inhaled gas.

9. The apparatus of claim 1, wherein the fluid traveling in the predetermined direction comprises exhaled gas.

10. The apparatus of claim 1, wherein the patient respiratory system interface comprises a mouthpiece.

11. The apparatus of claim 1, wherein the patient respiratory interface comprises a mask.

12. The apparatus of claim 1, wherein the valve comprises an annular valve.

13. The apparatus of claim 1, wherein the valve comprises a duck-bill valve.

14. The apparatus of claim 1, wherein the variable resistance bypass window comprises first and second overlapping apertures (528, 530) continuously positionable relative to one another between a first position, where the first and second overlapping apertures align to provide a maximum opening between the patient respiratory system interface and ambient air outside of the patient respiratory system interface, and a second position, where the first and second apertures cooperate to provide a minimum opening between the patient respiratory system interface and ambient air outside of the patient respiratory system interface.

15. The apparatus of claim 14 wherein the first overlapping aperture comprises a fixed opening in the patient respiratory system interface that is fixed relative to the patient respiratory system interface and the second overlapping aperture comprises a movable control portion defining a movable opening that is continuously adjustable to overlap with the fixed opening.

16. The apparatus of claim 15, wherein the fixed opening is positioned over the movable opening.

17. The apparatus of claim 15, wherein the fixed opening is positioned beneath the movable opening.

18. The apparatus of claim 14, wherein the valve is positioned to permit inhalation and restrict exhalation, wherein an exhalation flows through the variable resistance bypass window.

19. The apparatus of claim 16, wherein the fixed opening comprises an opening in a circular wall formed in the patient respiratory interface, and the control portion comprises a strip of material movably positioned adjacent the opening in the circular wall.

20. The apparatus of claim 19, wherein the movable opening defined by the strip of material comprises a variable height portion of the strip of material, the variable height portion having a continuously sloping height ranging from a first height that is at least as high as a height of the circular wall, to a second height that is less than the height of the circular wall.

21. The apparatus of claim 20, wherein the strip of material is an arcuate strip of material.

22. The apparatus of claim 19, wherein the movable
opening defined by the strip of material comprises a variable width opening in at least a portion of the strip of material, the variable width opening having a continuously changing width from a first position to a second position along a length of the strip.

23. The apparatus of claim 22, wherein the strip of material is an arcuate strip of material.

24. The apparatus of claim 1, wherein the valve comprises a one-way inhalation valve, and wherein the apparatus further comprises a discrete one-way exhalation valve mounted independently of the one-way inhalation valve.

25. The apparatus of claim 1 comprising:

an aerosol delivery apparatus in fluid communication with the patient respiratory system interface.

26. The apparatus of claim 25, wherein the valve assembly is in fluid communication with both the patient respiratory system interface and the aerosol delivery apparatus.

27. The apparatus of claim 25, wherein the aerosol delivery apparatus comprises a nebulizer.

28. The apparatus of claim 25, wherein the aerosol delivery apparatus comprises a metered dose inhaler.

29. The apparatus of claim 25, wherein the aerosol delivery apparatus comprises an aerosol holding chamber.

Patentansprüche

1. Vorrichtung mit positivem Atmungsdruck (10, 100, 210, 310, 410, 510, 600, 610) mit:

   einem Patienten-Beatmungssystemanschluss (16, 116, 216, 316, 416, 462, 512); und

   einer Ventilanordnung in Fluidverbindung mit dem Patienten-Beatmungssystemanschluss, wobei die Ventilanordnung umfasst:

   ein Ventil (12, 112, 278, 378, 412, 467, 475, 520, 606, 626), das ausgebildet ist, ein Fluid, das in einer vorbestimmten Richtung strömend, von einer ersten Seite zu einer zweiten Seite des Ventils zu leiten;

   ein Bypass-Fenster mit variablen Widerstand (28, 128, 225, 428, 471, 480, 528, 610, 622), das benachbart zu dem Ventil angeordnet und einen Widerstand für ein Fluid, das in einer Richtung umgekehrt zu der vorbestimmten Richtung strömend, aufweist, wobei das Bypass-Fenster mit variablen Widerstand kontinuierlich zwischen einem ersten Fluidwiderstand und einem zweiten Fluidwiderstand einstellbar ist: gekennzeichnet durch eine Einwege-Ausatemventileinrichtung (286, 423), die einen Ausatemweg bereitstellt und ein Einatem entlang des Ausatemweges verhindert, wobei der Ausatemweg unterschiedlich ist zu dem Einatemweg.

2. Vorrichtung nach Anspruch 1, wobei das Ventil einen Einwege-Einatemventilbereich (127, 279, 463, 476) aufweist, wobei der Einwege-Einatemventilbereich einen Einatemweg bereitstellt und ein Ausatem entlang des Einatemweges verhindert.

3. Vorrichtung nach Anspruch 2, wobei die Einwege-Ausatemventileinrichtung in dem Ventil enthalten ist.

4. Vorrichtung nach Anspruch 3, wobei das Ventil (278) ein Entenschabelventil mit einem offenen zentralen Gebiet und einem zentralen Venteilelement aufweist, wobei das zentrale offene Gebiet und das zentrale Venteilelement den Einatemweg definieren, und wobei das Entenschabelventil ferner einen peripheren Ausatemflansch (286) umfasst, der den Ausatemweg definiert.

5. Vorrichtung nach Anspruch 1, wobei das Ventil ein Einwegeventil umfasst.

6. Vorrichtung nach Anspruch 1, wobei das Ventil ein Einwege-Einatemventil umfasst.

7. Vorrichtung nach Anspruch 1, wobei das Ventil ein Einwege-Ausatemventil umfasst.

8. Vorrichtung nach Anspruch 1, wobei das in der vorbestimmten Richtung strömende Fluid ein eingemietetes Gas aufweist.

9. Vorrichtung nach Anspruch 1, wobei das in der vorbestimmten Richtung strömende Fluid ausgemaaktetes Gas umfasst.

10. Vorrichtung nach Anspruch 1, wobei der Patienten-Beatmungssystemanschluss ein Mundstück umfasst.

11. Vorrichtung nach Anspruch 1, wobei der Patienten-
Beatmungssystemanschluss eine Maske umfasst.

12. Vorrichtung nach Anspruch 1, wobei das Ventil ein ringförmiges Ventil umfasst.

13. Vorrichtung nach Anspruch 1, wobei das Ventil ein Entenschnabelventil umfasst.

14. Vorrichtung nach Anspruch 1, wobei das Bypass-Fenster mit variablen Widerstand eine erste Öffnung und eine zweite Öffnung (528, 530) aufweist, die sich überlappen und die in Bezug zueinander zwischen einer ersten Position, in der die erste und die zweite überlappende Öffnung zueinander ausgerichtet sind, um eine maximale Öffnung zwischen dem Patienten-Beatmungssystemanschluss und einer Umgebungsluft außerhalb des Patienten-Beatmungssystemanschlusses bereitzustellen und einer zweiten Position, in der die erste und die zweite Öffnung zusammenwirken, um eine minimale Öffnung zwischen dem Patienten-Beatmungssystemanschluss und der Umgebungsluft außerhalb des Patienten-Beatmungssystemanschlusses bereitzustellen, kontinuierlich positionierbar sind.

15. Vorrichtung nach Anspruch 14, wobei die erste überlappende Öffnung eine fixierte Öffnung in dem Patienten-Beatmungssystemanschluss aufweist, die relativ zu dem Patienten-Beatmungssystemanschluss fixiert ist, und wobei die zweite überlappende Öffnung einen bewegbaren Steuerbereich aufweist, der eine bewegliche Öffnung definiert, die kontinuierlich einstellbar ist, um mit der fixierten Öffnung zu überlappen.

16. Vorrichtung nach Anspruch 15, wobei die fixierte Öffnung über der bewegbaren Öffnung positioniert ist.

17. Vorrichtung nach Anspruch 15, wobei die fixierte Öffnung unter der bewegbaren Öffnung positioniert ist.

18. Vorrichtung nach Anspruch 14, wobei das Ventil so positioniert ist, dass Einatmung möglich und Ausatmung beschränkt ist, wobei Einatmungsgas durch das Bypass-Fenster mit variablen Widerstand strömt.

19. Vorrichtung nach Anspruch 16, wobei die fixierte Öffnung eine Öffnung in einer kreisförmigen Wand umfasst, die in dem Patienten-Beatmungssystemanschluss ausgebildet ist, und wobei der Steuerbereich einen Materialstreifen aufweist, der bewegbar benachbart zu der Öffnung in der kreisförmigen Wand positioniert ist.

20. Vorrichtung nach Anspruch 19, wobei die bewegbare Öffnung, die von dem Materialstreifen definiert ist, einen Bereich mit variabler Höhe des Materialstreifens umfasst, wobei der Bereich mit variabler Höhe eine kontinuierlich ansteigende Höhe im Bereich von einer ersten Höhe, die mindestens so groß ist wie eine Höhe der kreisförmigen Wand, bis zu einer zweiten Höhe aufweist, die kleiner ist als die Höhe der kreisförmigen Wand.

21. Vorrichtung nach Anspruch 20, wobei der Materialstreifen ein bogenförmiger Materialstreifen ist.

22. Vorrichtung nach Anspruch 19, wobei die bewegbare Öffnung, die von dem Materialstreifen definiert ist, eine Öffnung mit variabler Breite in mindestens einem Bereich des Materialstreifens aufweist, wobei die Öffnung mit variabler Breite eine Breite aufweist, die sich kontinuierlich von einer ersten Position zu einer zweiten Position entlang der Länge des Streifens ändert.

23. Vorrichtung nach Anspruch 22, wobei der Materialstreifen ein bogenförmiger Materialstreifen ist.

24. Vorrichtung nach Anspruch 1, wobei das Ventil ein Einwege-Einatmungsventil umfasst, und wobei die Vorrichtung ferner ein separates Einwege-Ausatmungsventil aufweist, das unabhängig von dem Einwege-Einatmungsventil montiert ist.

25. Vorrichtung nach Anspruch 1, mit:

einer Aerosol-Spendervorrichtung, die mit dem Patienten-Beatmungssystemanschluss in Fluidverbindung steht.


27. Vorrichtung nach Anspruch 25, wobei die Aerosol-Spendervorrichtung einen Vernebler umfasst.

28. Vorrichtung nach Anspruch 25, wobei die Aerosol-Spendervorrichtung eine Dossier-Inhaliervorrichtung umfasst.

29. Vorrichtung nach Anspruch 25, wobei die Aerosol-Spendervorrichtung eine Aerosol-Vorratskammer umfasst.

Revendications

1. Appareil de pression respiratoire positive (10, 100, 210, 310, 410, 510, 600, 610) comprenant:
une interface de système respiratoire de patient (16, 116, 216, 316, 416, 462, 512) ; et un ensemble de soupape en communication fluide avec l'interface de système respiratoire de patient, l'ensemble de soupape comprenant :

une soupape (12, 112, 278, 378, 412, 467, 475, 520, 606, 626) configurée pour faire passer un fluide se déplaçant dans un sens prédéterminé d'un premier côté à un second côté de la soupape ; une fenêtre de dérivation à résistance variable (28, 128, 225, 428, 471, 480, 528, 610, 622) positionnée à proximité de la soupape et ayant une résistance à un fluide se déplaçant dans un sens opposé au sens prédéterminé, dans lequel la fenêtre de dérivation à résistance variable est réglable en continu entre une première résistance au fluide et une seconde résistance au fluide ; et caractérisé par un moyen de soupape d'expiration à sens unique (286, 423) fournissant un passage d'expiration et empêchant l'inhalation le long du passage d'expiration, dans lequel le passage d'expiration est différent du passage d'inhalation.

2. Appareil selon la revendication 1, dans lequel la soupape comprend une partie de soupape d'inhalation à sens unique (127, 279, 463, 476), la partie de soupape d'inhalation à sens unique fournissant un passage d'inhalation et empêchant l'expiration le long du passage d'inhalation.

3. Appareil selon la revendication 2, dans lequel le moyen de soupape d'expiration à sens unique est compris dans la soupape.

4. Appareil selon la revendication 3, dans lequel la soupape (278) comprend une soupape en bec de canard comprenant une région centrale ouverte et un élément de soupape central, dans lequel la région centrale ouverte et l'élément de soupape central définissent le passage d'inhalation, et dans lequel la soupape en bec de canard comprend en outre une bride d'expiration périphérique (286) définissant le passage d'expiration.

5. Appareil selon la revendication 1, dans lequel la soupape comprend une soupape à sens unique.

6. Appareil selon la revendication 1, dans lequel la soupape comprend une soupape d'inhalation à sens unique.

7. Appareil selon la revendication 1, dans lequel la soupape comprend une soupape d'expiration à sens unique.

8. Appareil selon la revendication 1, dans lequel le fluide se déplaçant dans le sens prédéterminé comprend du gaz inhalé.

9. Appareil selon la revendication 1, dans lequel le fluide se déplaçant dans le sens prédéterminé comprend du gaz expiré.

10. Appareil selon la revendication 1, dans lequel l'interface de système respiratoire de patient comprend un embout buccal.

11. Appareil selon la revendication 1, dans lequel l'interface respiratoire de patient comprend un masque.

12. Appareil selon la revendication 1, dans lequel la soupape comprend une soupape annulaire.

13. Appareil selon la revendication 1, dans lequel la soupape comprend une soupape en bec de canard.

14. Appareil selon la revendication 1, dans lequel la fenêtre de dérivation à résistance variable comprend des première et seconde ouvertures en chevauchement (528, 530) pouvant être positionnées en continu l'une par rapport à l'autre entre une première position, où les première et seconde ouvertures en chevauchement s'alignent pour fournir une ouverture maximum entre l'interface de système respiratoire de patient et l'air ambiant à l'extérieur de l'interface de système respiratoire de patient, et une seconde position, où les première et seconde ouvertures coopèrent pour fournir une ouverture minimum entre l'interface de système respiratoire de patient et l'air ambiant à l'extérieur de l'interface de système respiratoire de patient.

15. Appareil selon la revendication 14, dans lequel la première ouverture en chevauchement comprend une ouverture fixe dans l'interface de système respiratoire de patient qui est fixe par rapport à l'interface de système respiratoire de patient et la seconde ouverture en chevauchement comprend une partie de commande mobile définissant une ouverture mobile qui est réglable dans un continu pour chevauchement avec l'ouverture fixe.

16. Appareil selon la revendication 15, dans lequel l'ouverture fixe est positionnée par-dessus l'ouverture mobile.

17. Appareil selon la revendication 15, dans lequel l'ouverture fixe est positionnée en dessous de l'ouverture mobile.
18. Appareil selon la revendication 14, dans lequel la soupape est positionnée pour permettre l'inhalation et limiter l'expiration, dans lequel une expiration s'écoule à travers la fenêtre de dérivation à résistance variable.

19. Appareil selon la revendication 16, dans lequel l'ouverture fixe comprend une ouverture dans une paroi circulaire formée dans l'interface respiratoire de patient, et la partie de commande comprend une bande de matériau positionnée de façon mobile à côté de l'ouverture dans la paroi circulaire.

20. Appareil selon la revendication 19, dans lequel l'ouverture mobile définie par la bande de matériau comprend une partie de hauteur variable de la bande de matériau, la partie de hauteur variable ayant une hauteur en pente continue allant d'une première hauteur qui est au moins aussi élevée qu'une hauteur de la paroi circulaire, à une seconde hauteur qui est inférieure à la hauteur de la paroi circulaire.

21. Appareil selon la revendication 20, dans lequel la bande de matériau est une bande de matériau arquée.

22. Appareil selon la revendication 19, dans lequel l'ouverture mobile définie par la bande de matériau comprend une ouverture de largeur variable dans au moins une partie de la bande de matériau, l'ouverture de largeur variable ayant une largeur changeant en continu d'une première position à une seconde position le long d'une longueur de la bande.

23. Appareil selon la revendication 22, dans lequel la bande de matériau est une bande de matériau arquée.

24. Appareil selon la revendication 1, dans lequel la soupape comprend une soupape d'inhalation à sens unique, et dans lequel l'appareil comprend en outre une soupape d'expiration à sens unique distincte montée indépendamment de la soupape d'inhalation à sens unique.

25. Appareil selon la revendication 1 comprenant :

un appareil de distribution d'aérosol en communication fluide avec l'interface de système respiratoire de patient.

26. Appareil selon la revendication 25, dans lequel l'ensemble de soupape est en communication fluide avec à la fois l'interface de système respiratoire de patient et l'appareil de distribution d'aérosols.

27. Appareil selon la revendication 25, dans lequel l'appareil de distribution d'aérosols comprend un nébuliseur.

28. Appareil selon la revendication 25, dans lequel l'appareil de distribution d'aérosols comprend un inhalateur-doseur.

29. Appareil selon la revendication 25, dans lequel l'appareil de distribution d'aérosol comprend une chambre de réception d'aérosol.