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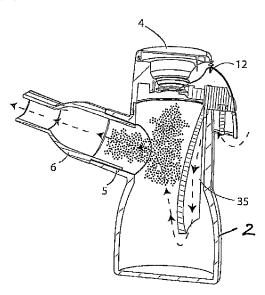
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(54) Title: A NEBULISER SYSTEM



(57) Abstract: A nebuliser 1 particularly for home use comprises a housing 2 having an aerosol generator 3 mounted therein. The housing is closed by a hinged cap 4 and has an outlet 5 through which aerosol generated by the generator 3 is delivered. A suitable connector such as a mouthpiece 6 may be connected to the outlet 5. The housing 2 has a receiver for receiving a nebule 8. The receiver comprises a slot 7 which is sized and/or shaped to accommodate only a nebule 8 of predetermined shape and/or size. The nebuliser cap 4 has a nebule opening means which may be in the form of a puncture pin 10 projecting therefrom to pierce the nebule 8 when the cap is moved to the closed position. The pin 10 is hollow or splined to regulate flow from the nebule. The nebuliser 1 also has a switch means for enabling operation of the aerosol generator 3 when the nebule 8 is opened. The switch means comprises a microswitch 12 on the housing which is engageable by a trigger 13 on the cap 4 when the cap is moved to the closed position. Aerosol generated by the aerosol generator 3 is delivered into a chamber defined by the nebuliser housing 2. Air passes into the chamber through air inlets 30. The air entrains the aerosolised medicament and the entrained aerosolised medicament is delivered from the nebuliser through the outlet 5.



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"A nebuliser system"

Introduction

Existing nebulisation systems for home use can be used with any drug.

In some instances this presents a risk that a patient or carer may inadvertently use an incorrect drug or an incorrect dosage of drug.

Statements of Invention

In one aspect the invention provides a nebuliser comprising:

a housing;

an aerosol generator mounted in the housing; and

a nebule receiver in the housing for receiving a nebule containing liquid to be aerosolised by the aerosol generator.

According to the invention there is provided a nebuliser comprising

a housing;

an aerosol generator mounted in the housing;

a receiver for receiving a nebule containing a liquid to be aerosolised by the aerosol generator, a nebule being opened on insertion into the nebule receiver to deliver liquid from the nebule to the aerosol generator; and

switch means for enabling operation of the aerosol generator on insertion and/or opening of a nebule.

The housing has an air inlet and an outlet for air with aerosol entrained therein. A mouthpiece can be attached to the housing outlet.

In one embodiment the nebuliser has an opening means for opening the nebule.

The opening means may comprise a nebule puncturing means.

The opening means may comprise means for cutting through a nebule.

The opening means may comprise means for tearing a nebule.

The opening means may comprise means for compressing a nebule.

The opening means may comprise means for squeezing a nebule.

In one case the nebuliser is closed by a cap.

The opening of the nebule may be triggered by the cap.

The activation of the switch means may be triggered by the cap.

In one embodiment the cap is movably mounted to the housing.

The cap may be hingedly mounted to the housing.

The cap may comprise a nebule opening means.

The nebule opening means may comprise a nebule puncturing means. The puncturing means may comprise a puncture pin projecting inwardly of the cap.

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The puncture system may consist of a splined dagger. The splines would facilitate the ingress of air to the nebule to facilitate a drug stream to flow into the nebulizing system.

The switch means may be activated on movement of the cap from an open to a closed position.

In one embodiment the receiver is configured to facilitate opening of a nebule.

The receiver can comprise an opening in the housing to receive a nebule.

The receiver can comprise a skirt extending from the housing opening.

The skirt can comprise the switch means which is activated on insertion of a nebule.

In one embodiment the receiver is adapted to allow insertion of only a pre-determined nebule.

In one embodiment the receiver comprises a slot for a nebule, the slot being sized and/or shaped to accommodate a nebule of predetermined shape and/or size.

In another embodiment the nebuliser comprises a baffle means to elongate the path travelled by aerosol. The elongation of the flow path is assisted by increasing the breathing resistance by adjusting the size of the openings of the air vents. The efficiency of drug deposition is thereby enhanced as the aerosol is entrained from the generator to the mouth piece.

In one embodiment the nebuliser has air vent means and an air vent restrictor means to adjust the size of the air vent means.

In another aspect the invention provides a nebulisation kit comprising a nebuliser of the invention and at least one nebule or ampoule for use with the nebuliser. The nebule is operable from a closed configuration to an open configuration.

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In one case the nebule comprises at least one zone of weakness which is fractured to open the nebule.

The nebule typically comprises a liquid outlet and a vent inlet. The liquid outlet and the vent inlet can be defined by zones of weakness in the nebule.

In one embodiment the nebule comprises a tab which, on insertion of the nebule into the receiver causes the liquid to be released from the nebule.

In one case the nebule comprises a top wall, a base and sidewall means extending between the base and the top, the tab forming an extension of the base.

The nebule may comprise an internal rib extending from the top to the base.

In one case the nebule comprises a first zone of weakness at a junction between the top wall and the internal rib. The nebule may also comprise a second zone of weakness at a junction between the base and the sidewall means.

In one case the tab is movable, on insertion of the nebule into the receiver to break the joint between the base wall and the sidewall. The movement of the tab may cause the joint between the internal rib and the top wall to break.

In a further aspect the invention provides a nebule comprising a top wall, a base wall, sidewall means extending between the base and a tab extending from the base wall, the tab being movable to fracture the joint between the base wall and the sidewall for release of liquid from the nebule.

In one case the nebule comprises an internal rib extending between the base wall and the top wall.

The nebule can have a zone of weakness in the top wall at a joint between the top wall and the internal rib, movement of the tab causing the joint between the rib and the top wall to fracture.

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In a further aspect the invention provides a nebuliser comprising:-

a housing;

an aerosol generator mounted in the housing; and

a baffle in the housing to elongate the path travelled by aerosol generated by the aerosol generator.

The housing has an air inlet and an outlet for air with aerosol entrained therein.

There may be a mouthpiece attached to the housing outlet.

In one case the baffle is located in the housing between the inlet and the outlet.

The elongation of the flow path may be assisted by increasing the breathing resistance by adjusting the size of the openings of the air vents. The efficiency of drug deposition is thereby enhanced as the aerosol is entrained from the generator to the mouth piece.

In another aspect the invention provides a nebuliser comprising:-

a housing;

an aerosol generator mounted in the housing;

the housing having an air vent means; and

an adjustable restrictor to adjust the size of the opening defined by the air vent means.

The housing has an outlet for air with aerosol entrained therein.

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There may be a mouthpiece attached to the housing outlet.

In one case the nebuliser comprises a baffle or auxiliary wall within the housing to elongate the aerosol flow path.

The auxiliary wall may be located between the air inlet and the aerosol outlet.

In one embodiment the nebuliser comprises a baffle or auxiliary wall to elongate the aerosol flow path. This increases the efficiency of drug deposition.

An advantageous feature is the matching of a particular drug contained within a pre specified nebule shape than can only be used in conjunction with the nebuliser design which is designed to nebulise the drug for a predetermined disease treatment. Putting the particular drug into a different nebuliser would not have the desired clinical effect. Putting a different drug into the same nebuliser would also not have the same clinical effect.

The nebuliser has opening means for opening the nebule to allow delivery of liquid from the nebule to the aerosol generator. The nebuliser also has switch means for enabling operation of the aerosol generator when the nebule is opened.

The closing cap may aid the egress of drug from the vial by a cutting, tearing, and/or a squeezing or compressing motion.

In one embodiment the switch means is activated on movement of the cap from an open to a closed position to facilitate activation of the aerosol generator to generate the aerosol

Preferably the receiver is adapted to allow insertion of only a pre-determined nebule. The receiver may comprise a slot for a nebule, the slot being sized and/or shaped to accommodate a nebule of predetermined geometrical shape and/or size.

In one embodiment the nebuliser comprises baffle means to elongate the path travelled by aerosol from the generator to the patient. This causes the aerosol to be entrained thus facilitating a greater amount of the aersolised drug to be delivered to the patient.

In another aspect the nebuliser has air vent means and an air vent restrictor means to adjust the size of the air vent means. This has the advantage of adjusting the breathing resistance of the nebulizer to suit different patient groups, for example a large elderly person would have a different breathing profile to a child.

Brief Description of the Drawings

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The invention will be more clearly understood from the following description thereof given by way of example only with reference to the accompanying drawings, in which:-

Fig. 1 is a cross sectional view of a nebuliser according to the invention in one configuration;

Fig. 2 is a cross sectional view of the nebuliser of Fig. 1 with a nebule in position;

Fig. 3 is a cross sectional view of the nebuliser in another configuration in which the nebule is punctured;

Fig. 4 is a cross sectional view of the nebuliser in the configuration of Fig. 3 with aerosol generated;

Fig. 5 is a cross sectional view similar to Fig. 4 illustrating the flow path;

Figs. 6 to 11 are views of part of the nebuliser of Figs. 1 to 4 in different configurations;

Figs. 12 and 13 are views similar to Figs. 7 and 8 of a nebuliser with an alternative nebule;

Fig. 14 is an isometric partially cut-away view of another nebuliser according to the

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invention with a sealed nebule according to the invention positioned on top of the

nebuliser ready for opening;

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Fig. 15 is a cross sectional view of the nebuliser and nebule of Fig 14;

Fig. 16 is a cross sectional view of the nebuliser and nebule pushed into the nebuliser,

thus opening the nebule and allowing the escapement of the nebule contents (which may

include a drug) which egresses to the aerosol generator;

Fig. 17 is an isometric view of a nebule according to the invention with a top wall

removed;

Fig. 18 is a cross sectional view of the closed nebule;

Figs. 19 and 20 are enlarged cross sectional views of details of the nebule of Fig 18; and

Figs. 21 to 23 are cross sectional views illustrating the opening of the nebule.

Detailed Description

Referring to the drawings and initially to Figs. 1 to 11 thereof there is illustrated a nebuliser 1 according to the invention which comprises a housing 2 having an aerosol generator 3 mounted therein. The housing is closed by a hinged cap 4 and has an inlet and an outlet 5 through which aerosol generated by the generator 3 is delivered. A suitable connector such as a mouthpiece 6 may be connected to the outlet 5. The nebuliser may be used on an outpatient basis and on non ventilated patients and is particularly suited for home use.

The housing 2 has a receiver for receiving a nebule 8. The receiver comprises a slot 7 which is sized and/or shaped to accommodate only a nebule 8 of predetermined shape and/or size.

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The nebuliser cap 4 has a nebule opening means which in this case is in the form of a puncture pin 10 projecting therefrom to pierce the nebule 8 when the cap is moved to the closed position. The pin 10 is hollow or splined to regulate flow from the nebule. Making the pin hollow or splined facilitates air ingress to the nebule thus allowing the drug to be released into the nebuliser.

The cap, on closing may alternatively or additionally apply a compressive load on the nebule to aid the egress of drug from the nebule.

The nebuliser 1 also has a switch means for enabling operation of the aerosol generator 3 when the nebule 8 is opened. The switch means in this case comprises a micro-switch 12 on the housing which is engageable by a trigger 13 on the cap 4 when the cap is moved to the closed position.

In the invention the nebuliser has a hinged lid with a nebule-specific slot. Closing of the lid will enable drug release from the nebule. The closing of the lid will also electronically indicate to the nebuliser that the correct nebule is in correct position and that nebulisation can commence.

Aerosol generated by the aerosol generator 3 is delivered into a chamber defined by the nebuliser housing 2. Air passes into the chamber through air inlets 30. The air extrains the aerosolised medicament and the entrained aerosolised medicament is delivered from the nebuliser through the outlet 5. The vent openings 30 are adjustable to vary the breathing resistance. Different patients have different resistances and these can be adjusted to suit their breathing pattern. For example longer and slower breaths increases the drug intake.

A baffle system such as an internal or auxiliary wall 35 within the housing chamber 2 may be provided. The wall 35 is interposed between an air inlet 30 and the outlet 5 and forces air to flow towards the base of the housing 2 so that it entrains the aerosol generated. This elongates the distance travelled or uninterrupted pathway by aerosol from generation at 3 to the mouthpiece 6. This has been shown to increase the efficiency of the system by entraining more aerosol to be conveyed to the mouthpiece and patient.

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Referring to Figs. 12 and 13, there is illustrated part of another nebuliser according to the invention in which parts similar to those described with reference to Figs. 1 to 11 are assigned the same reference numerals. In this case a nebule 20 is of a different size and shape to the nebule 8 used in the nebuliser of Figs. 1 to 11.

Figs 14 to 16 illustrate a similar aerosolising system to those described above with like parts being assigned the same reference numerals. The nebuliser system uses a different nebule 50 which is illustrated in Figs 17 to 23.

In this case, the action of depressing the nebule 50 into the aerosol generator causes the nebule 50 to tear open in a controlled manner and release the contents of the nebule (which may include drug) {expect it always include a drug} to the aerosol generator and at the same time depressing a micro-switch 51 to close the electrical circuit to cause aersolisation to occur. The micro switch 51 may be engaged by the drug nebule 50. This switch 51 is activated by the nebule 50 and the switch 51 activates the generation of aerosol as the nebule is depressed into position.

The nebuliser in this case comprises a top opening 52 leading to a chamber 53 above the aerosol generator 3. A skirt 54 extends downwardly from the top opening 52. The switch 51 is provided on the skirt 54. The skirt 54 has an inturned lip 55 near the lower end thereof.

In more detail, the nebule or ampoule 50 comprises a base wall 60, a top wall 61 and side wall means extending between the base wall 60 and the top wall 61. In this case the nebule 50 is of generally rectilinear shape in transverse cross section and comprises side walls 62, 63 and end walls 64, 65. The nebule 50 also has an internal rib 66 which extends from the base wall 60 and is engagable in a receiver slot 71 on top of the nebuliser.

The nebule 50 has means to encourage localised fracturing or separation to facilitate opening of the nebule for release of the contents. In this embodiment the nebule has two zones of weakness which may be provided by a local thinning of the walls of the nebuliser. A first zone of weakness 75 is provided between the bottom of the side wall 64 and the base wall 60 opposite to the tab 70. A second zone of weakness 76 is provided between the top of the internal rib 66 and

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the top wall 61. In use, when a downwards force F is applied to the top of the nebule 50 the tab 70 is initially held in the slot 71. As further force is applied the zones of weakness 75, 76 cause parts of the nebule to fracture or separate. The second zone 76 causes the rib 66 to separate from the top wall 61 and thus define a hole 80 which defines an air inlet / vent. The first zone 75 causes the base wall 60 to fracture or separate from the side wall 64 thereby defining an outlet hole 81 through which the contents of the nebule 50 pass to be delivered into the chamber 53 which leads the liquid to the aerosol generator 3. Because the switch 51 is activated by the tab 70, as illustrated, a signal is provided to activate the aerosol generator.

It will be appreciated that at least some of the features described herein may be used in association with nebules and/or nebulisers for ventilator circuits. At least some of the features described herein may be used with nebules and/nebulisers for laparoscopy applications as described in our US2008/0243050A1, the entire contents of which are incorporated herein by reference.

The nebuliser (or aerosol generator) 3, has a vibratable member which is vibrated at ultrasonic frequencies to produce liquid droplets. Some specific, non-limiting examples of technologies for producing fine liquid droplets is by supplying liquid to an aperture plate having a plurality of tapered apertures extending between a first surface and a second surface thereof and vibrating the aperture plate to eject liquid droplets through the apertures. Such technologies are described generally in U.S. Pat. Nos. 5,164,740; 5,938,117; 5,586,550; 5,758,637; - 6755189, 6540154, 6926208, 7174888, 6546927, 6,085,740, and US2005/021766A, the complete disclosures of which are incorporated herein by reference. However, it should be appreciated that the present invention is not limited for use only with such devices.

In use, the liquid to be aerosolised is received at the first surface, and the aerosol generator 3 generates the aerosolised liquid at the second surface by ejecting droplets of the liquid upon vibration of the vibratable member. The apertures in the vibratable member are sized to aerosolise the liquid by ejecting droplets of the liquid such that the majority of the droplets by mass have a size of less than 5 micrometers.

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In one case the aerosol generator 3 comprises a vibratable member, a piezoelectric element and a washer, which are sealed within a silicone overmould and secured in place within the housing using a retaining ring. The vibratable member has a plurality of tapered apertures extending between a first surface and a second surface thereof.

The first surface of the vibratable member, which in use faces upwardly, receives the liquid from the nebule 50 and the aerosolised liquid, is generated at the second surface of the vibratable member by ejecting droplets of liquid upon vibration of the member. In use, the second surface faces downwardly. In one case, the apertures in the vibratable member may be sized to produce an aerosol in which the majority of the droplets by weight have a size of less than 5 micrometers. The vibratable member could be non-planar, and may be dome-shaped in geometry.

Any suitable medicament, therapeutic agent, active substance or pharmaceutically active compound than can be nebulised may be employed. {Comment – we do not expect this to be used to deliver a humidifying agent – only drug unless we are covering the potential for us ein the laparoscopy application} also act to deliver any agent presented in an aqueous drug solution. {These therapeutic agents could also act as humidifying substances in their own right.}

The system facilitates delivery in aerosol form of, for example, bronchodilators, including β-agonists, muscarinic antagonists, epinephrine;, surfactants; pain-relief medications including anaesthetics; migraine therapies; anti-infectives; anti-inflammatories, steroids, including corticostroids; chemotherapeutic agents;, mucolytics; vasodilators;,vaccines and hormones. In addition substances classified as anti-thrombogenic agents, anti-proliferative agents, monoclonal antibodies, anti-neoplastic agents, anti-mitotic agents, anti-sense agents, anti-microbial agents, nitric oxide donors, anti-coagulants, growth factors, translational promoter, inhibitors of heat shock proteins, biomoloecules including proteins, polypeptides and proteins, oligonucleotides, oligoproteins, siRNA, anti-sense DNA and RNA, ribozymes, genes, viral vectors, plasmids, liposomes, angiogenic factors, hormones, nucleotides, amino acids, sugars, lipids, serine proteases, anti-adhesion agents including but not limited to hyaluronic acid, biodegradable barrier agents may also be suitable.

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The medicament may for example, comprise long-acting beta –adrenoceptor agonists such as salmeterol and formoterol or short-acting beta –adrenoceptor agonists such as albuterol.

The medicament may be a long-acting muscarinic antagonists such as tiotropium (Spiriva) or short-acting muscarinic antagonists such as ipratropium (Atrovent).

Typical anti-infectives include antibiotics such as an aminoglycoside, a tetracycline, a fluroquinolone; anti-microbials such as a cephalosporin; and anti-fungals. Examples of antibiotics include anti-gram-positive agents such as macrolides, e.g. erythromycin, clarithromycin, azithromycin, and glycopeptides, e.g. vancomycin and teicoplanin, as well as any other anti-gram-positive agent capable of being dissolved or suspended and employed as a suitable aerosol, e.g. oxazoldinone, quinupristin/dalfopristen, etc. Antibiotics useful as anti-gram-negative agents may include aminoglycosides, e.g. gentamicin, tobramycin, amikacin, streptomycin, netilmicin, quinolones, e.g. ciprofloxacin, ofloxacin, levofloxacin, tetracyclines, e.g. oxytetracycline, dioxycycline, minocycline, and cotrimoxazole, as well as any other anti-gram-negative agents capable of being dissolved or suspended and employed as a suitable aerosol.

Anti-inflammatories may be of the steroidal such as budesonide or ciclesonide, non-steroidal, such as sodium cromoglycate or biological type.

Typical local anaesthetics are, for example, Ropivacaine, Bupivacaine, levobupivacaine, and Lidocaine.

Chemotherapeutic agents may be alkylating agents, antimetabolites, anthracyclines, plant alkaloids, topoisomerase inhibitors, nitrosoureas, mitotic inhibitors, monoclonal antibodies, tyrosine kinase inhibitors, hormone therapies including corticosteroids, cancer vaccines, antiestrogens, aromatase inhibitors, anti-androgens, anti-angiogenic agents and other anti-tumour agents.

Surfactant medications (sometimes referred to herein as "surfactants") are protein-lipid compositions, e.g. phospholipids, that are produced naturally in the body and are essential to the

lungs' ability to absorb oxygen. They facilitate respiration by continually modifying surface tension of the fluid normally present within the air sacs, or alveoli, that tube the inside of the lungs. In the absence of sufficient surfactant, these air sacs tend to collapse, and, as a result, the lungs do not absorb sufficient oxygen. Insufficient surfactant in the lungs results in a variety of respiratory illnesses in both animals and humans. Since most of these surfactant medications are animal-based, the current supply is limited, and although synthetic surfactants are available, their manufacture is both inexact and expensive. In addition, the surfactant medications are typically high in viscosity and are difficult to deliver to the patient's respiratory system. The increased efficiency of the pressure-assisted breathing system of the present invention, and the smaller amount of medicament required for a treatment according to the present invention, can be a substantial advantage when such scarce and expensive medicaments are employed. The combination of surfactant with other medicaments to improve distribution in the lung and body is also possible. Surfactants also possess the capacity to act as anti-adhesion agents.

The invention provides a closed-cup home care nebuliser. This system ensures that only the required nebule (and therefore drug) can be used with this high-efficiency system. The nebuliser chamber is designed to maximise efficiency of drug delivery. In use, fewer steps are involved for a patient in using the system. This is particularly advantageous for older people.

An adjustable air vent restrictor (hydrophobic media, or a mechanical slider occlusion system may be incorporated into nebuliser vent holes 30 to vary the air resistance through the system to condition patients breathing. Patients of differing ages or ailment conditions would have a different breathing pattern need. The breathing pattern is influenced by the inhalation and exhalation resistance when using the nebuliser. This can be controlled by changing the area of the vent holes 30. Closing these more increases the resistance and opening these more reduces the resistance. The air vent restrictor can be used to condition a patients breathing pattern. This helps to create a longer and slower breathing pattern to maximise drug delivery.

The invention is not limited to the embodiments hereinbefore described which may be varied in detail.

Claims

1. A nebuliser comprising

a housing;

an aerosol generator mounted in the housing;

a receiver for receiving a nebule containing a liquid to be aerosolised by the aerosol generator a nebule being opened on insertion into the nebule receiver to deliver liquid from the nebule to the aerosol generator; and

switch means for enabling operation of the aerosol generator on insertion and/or opening of a nebule.

- 2. A nebuliser as claimed in claim 1 wherein the housing has an air inlet and an outlet for air with aerosol entrained therein.
- 3. A nebuliser as claimed in claim 2 comprising a mouthpiece attached to the housing outlet.
- 4. A nebuliser as claimed in any of claims 1 to 3 wherein the nebuliser has an opening means for opening the nebule.
- 5. A nebuliser as claimed in claim 4 wherein the opening means comprises a nebule puncturing means.
- 6. A nebuliser as claimed in claim 4 or 5 wherein the opening means comprises means for cutting through a nebule.
- 7. A nebuliser as claimed in any of claims 4 to 6 wherein the opening means comprises means for tearing a nebule.

- 8. A nebuliser as claimed in any of claims 4 to 7 wherein the opening means comprises means for compressing a nebule.
- 9. A nebuliser as claimed in any of claims 4 to 8 wherein the opening means comprises means for squeezing a nebule.
- 10. A nebuliser as claimed in any of claims 1 to 9 wherein the nebuliser is closed by a cap.
- 11. A nebuliser as claimed in claim 10 wherein the opening of the nebule is triggered by the cap.
- 12. A nebuliser as claimed in claim 10 or 11 wherein the activation of the switch means is triggered by the cap.
- 13. A nebuliser as claimed in any of claims 10 to 12 wherein the cap is movably mounted to the housing.
- 14. A nebuliser as claimed in claim 13 wherein the cap is hingedly mounted to the housing.
- 15. A nebuliser as claimed in any of claims 10 to 14 wherein the cap comprises a nebule opening means.
- 16. A nebuliser as claimed in claim 15 wherein the nebule opening means comprises a nebule puncturing means.
- 17. A nebuliser as claimed in claim 16 wherein the puncturing means comprises a puncture pin projecting inwardly of the cap.
- 18. A nebuliser as claimed in any of claims 13 to 17 wherein the switch means is activated on movement of the cap from an open to a closed position.

- 19. A nebuliser as claimed in any of claims 1 to 3 wherein the receiver is configured to facilitate opening of a nebule.
- 20. A nebuliser as claimed in claim 19 wherein the receiver comprises an opening in the housing to receive a nebule.
- 21. A nebuliser as claimed in claim 20 wherein the receiver comprises a skirt extending from the housing opening.
- 22. A nebuliser as claimed in claim 21 wherein the skirt comprises the switch means which is activated on insertion of a nebule.
- 23. A nebuliser as claimed in any of claims 1 to 22 wherein the receiver is adapted to allow insertion of only a pre-determined nebule.
- 24. A nebuliser as claimed in any of claims 1 to 23 wherein the receiver comprises a slot for a nebule, the slot being sized and/or shaped to accommodate a nebule of predetermined shape and/or size.
- 25. A nebuliser as claimed in any of claims 1 to 24 comprising a baffle means to elongate the path travelled by aerosol.
- 26. A nebuliser as claimed in any of claims 1 to 25 wherein the nebuliser has air vent means and an air vent restrictor means to adjust the size of the air vent means.
- 27. A nebulisation kit comprising a nebuliser as claimed in any of claims 1 to 26 and at least one nebule for use with the nebuliser.
- 28. A kit as claimed in claim 27 wherein the nebule is operable from a closed configuration to an open configuration.

- 29. A kit as claimed in claim 28 wherein the nebule comprises at least one zone of weakness which is fractured to open the nebule.
- 30. A kit as claimed in claim 28 or 29 wherein the nebule comprises a liquid outlet and a vent inlet.
- 31. A kit as claimed in claim 30 wherein the liquid outlet and the vent inlet are defined by zones of weakness in the nebule.
- 32. A kit as claimed in any of claims 28 to 31 wherein the nebule comprises a tab which, on insertion of the nebule into the receiver causes the liquid to be released from the nebule.
- 33. A kit as claimed in claim 32 wherein the nebule comprises a top wall, a base and sidewall means extending between the base and the top, the tab forming an extension of the base.
- 34. A kit as claimed in claim 33 wherein the nebule comprises an internal rib extending from the top to the base.
- 35. A kit as claimed in claim 34 wherein the nebule comprises a first zone of weakness at a junction between the top wall and the internal rib.
- 36. A kit as claimed in claim 34 or 35 wherein the nebule comprises a second zone of weakness at a junction between the base and the sidewall means.
- 37. A kit as claimed in any of claims 34 to 36 wherein the tab is movable, on insertion of the nebule into the receiver to break the joint between the base wall and the sidewall.
- 38. A kit as claimed in claim 37 wherein the movement of the tab causes the joint between the internal rib and the top wall to break.

- 39. A nebule comprising a top wall, a base wall, sidewall means extending between the base and a tab extending from the base wall, the tab being movable to fracture the joint between the base wall and the sidewall for release of liquid from the nebule.
- 40. A nebule as claimed in claim 39 comprising an internal rib extending between the base wall and the top wall.
- 41. A nebule as claimed in claim 40 wherein the nebule has a zone of weakness in the top wall at a joint between the top wall and the internal rib, movement of the tab causing the joint between the rib and the top wall to fracture.
- 42. A nebuliser comprising:-

a housing;

an aerosol generator mounted in the housing; and

- a baffle in the housing to elongate the flow path travelled by aerosol generated by the aerosol generator.
- 43. A nebuliser as claimed in claim 41 wherein the housing has an air inlet and an outlet for air with aerosol entrained therein.
- 44. A nebuliser as claimed in claim 43 comprising a mouthpiece attached to the housing outlet.
- 45. A nebuliser as claimed in claim 43 or 44 wherein the baffle is located in the housing between the inlet and the outlet.
- 46. A nebulizer a claimed in any of claims 42 to 45 wherein the elongation of the flow path is assisted by increasing the breathing resistance by adjusting the size of the openings of the air vents.

- 47. A nebulizer as claimed in claim 46 wherein the efficiency of drug deposition is increased as the aerosol is entrained from the generator to the mouth piece.
- 48. A nebuliser comprising:-

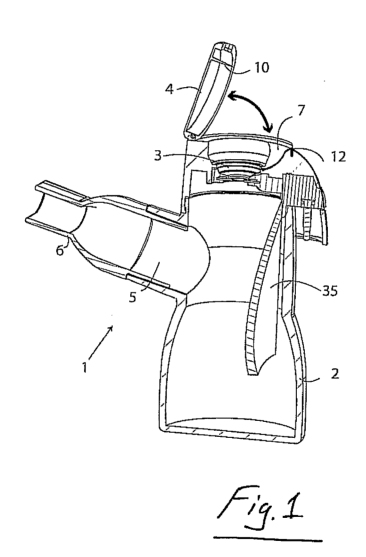
a housing;

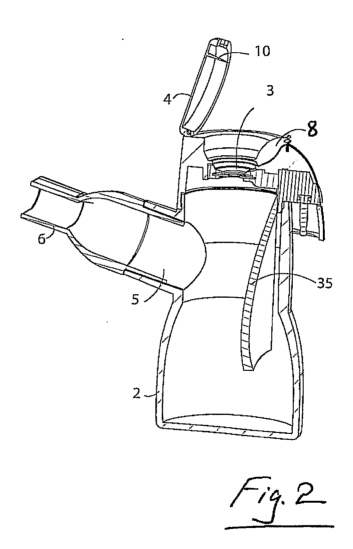
an aerosol generator mounted in the housing;

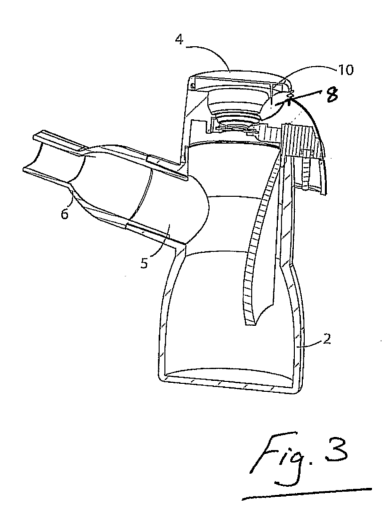
the housing having an air vent means; and

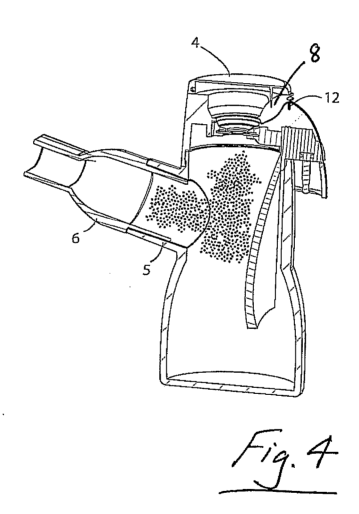
an adjustable restrictor to adjust the size of the opening defined by the air vent means.

- 49. A nebuliser as claimed in claim 48 wherein the housing has an outlet for air with aerosol entrained therein.
- 50. A nebuliser as claimed in claim 48 or 49 comprising a mouthpiece attached to the housing outlet.
- 51. A nebuliser as claimed in claim 49 or 50 comprising a baffle or auxiliary wall within the housing to elongate the aerosol flow path.
- 52. A nebuliser as claimed in claim 51 wherein the auxiliary wall is located between the air inlet and the aerosol outlet.
- 53. A nebuliser substantially as hereinbefore described with reference to the accompanying drawings.
- 54. A nebule substantially as hereinbefore described with reference to the accompanying drawings.

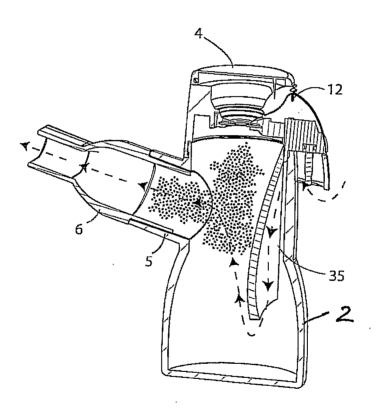




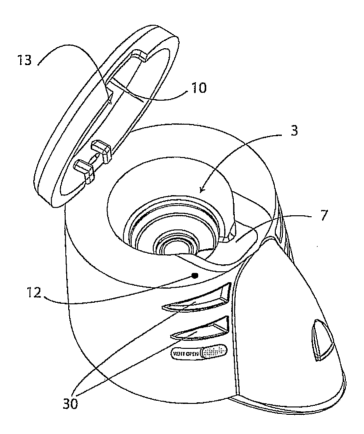




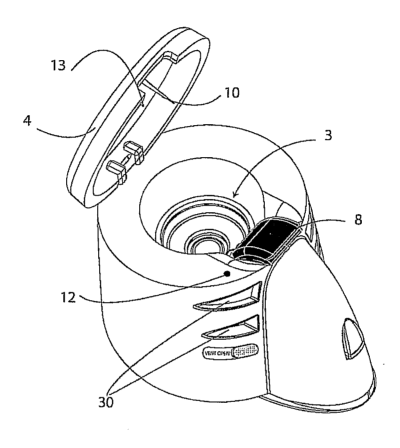
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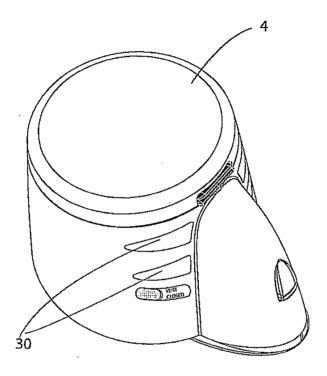
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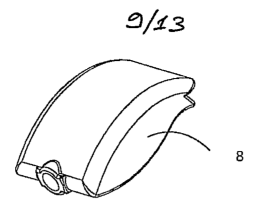


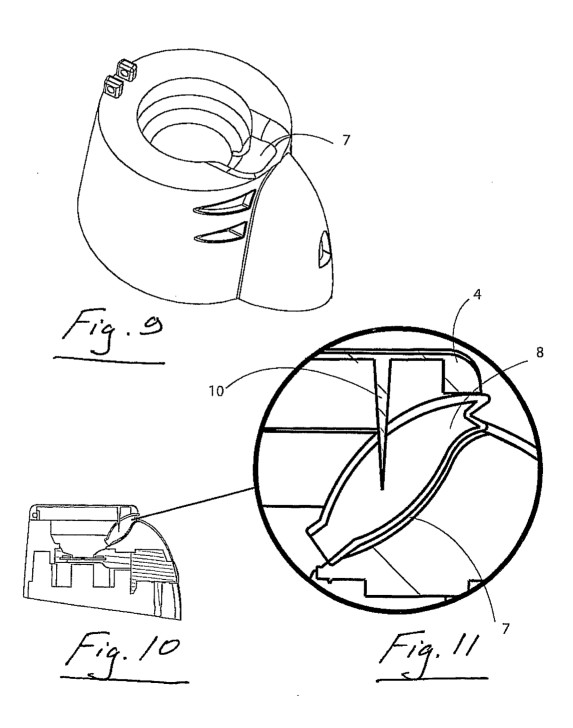
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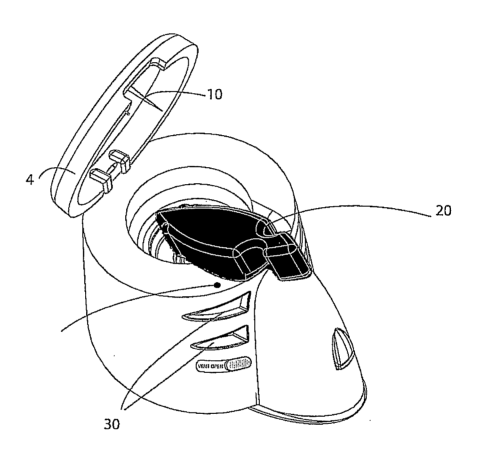


Fig. 12

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