(51) International Patent Classification: A61M 15/00

(21) International Application Number: PCT/EP2003/008432

(22) International Filing Date: 30 July 2003 (30.07.2003)

(25) Filing Language: English

(26) Publication Language: English


(72) Inventors: and

(75) Inventors/Applicants (for US only): PINON, John [BE/GB]; 6, rue des Frères Mourot, F-88390 Domeyne-sur-Avière (FR). SHIRGAONKAR, Sameer [GB/GB]; 15 Hazelbourne Road, London SW12 9NU (GB). SMIT, Christopher, James [GB/GB]; 37 Cage Hill, Swaffham Prior, Cambridge CB5 0JS (GB).

(74) Agent: ALBRECHT, Thomas; Kraus & Weisert, Thomas-Wimmer-Ring 15, 80539 Munich (DE).


(84) Designated States (regional): ARIPO patent (GH, GM, KE, LS, MW, MZ, SD, SL, SZ, TZ, UG, ZM, ZW), Eurasian patent (AM, AZ, BY, KG, KZ, MD, RU, TJ, TM),

[Continued on next page]

(54) Title: POWDER INHALER

(57) Abstract: A powder inhaler comprises a container (7) for storing a powdered medicament, a metering member (15) having a dosing recess (18) to be filled with a dose of the powdered medicament, and a mouthpiece (3) being in communication with an inhalation channel (27) of the powder inhaler. Furthermore, the powder inhaler comprises a protective member (19) which is slidingly moveable on the metering member (15) between a closed position, in which it at least covers the dosing recess (18) of the metering member (15) if the metering member (15) is in an inhalation position, and an open position, in which it exposes the dosing recess (18) thereby enabling inhalation of the dose of the powdered medicament contained in the dosing recess (18). The protective member (19) is preferably coupled to an inhalation actuated mechanism (21-23) in such a manner that the inhalation actuated mechanism (21-23) moves the protective member (19) from its closed position to its open position only if there is an inhalation suction force exerted by a user which exceeds a predetermined level. This prevents the dose contained in the dosing recess (18) from falling out of the dosing recess, unless there is no inhalation process initiated by the user. Therefore, the powder inhaler can also be operated reliably upside down. Furthermore, a deagglomerator arrangement (cyclose), which may be incorporated in such a powder inhaler, is proposed which comprises a vortex chamber (73) having a diameter between 6 mm and 8 mm, preferably 8 mm.
Published:

— with international search report
POWDER INHALER

The present invention relates to a powder inhaler, i.e. a device for dispensing a powdered medicament preparation by inhalation. The device is in particular a portable multiple-dose device without propellant gas, equipped with a metering member which dispenses doses from a medicament container. Moreover, the device is based on centripetal force for achieving a more effective pulverization of the powdered inhalation medicament in such a manner that the penetration of the medicament into the lungs of a patient is improved and the adhesion to the upper respiratory passages is reduced for alleviating possible side effects caused thereby.

The administering of a powdered medicament preparation by inhalation from an inhaler is commonly known. Multiple-dose type powder inhalers comprising a powder container and a metering member which measures and dispenses a unit dose are also known, for example from patent publications GB 2165159, EP 0 079 478, and EP 0 166 294. In these devices, a series of dosing recesses are notched into the surface of a cylindrical metering member, and the said member is disposed in a chamber of precisely the same shape. When the metering member is rotated, the dosing recesses in turn will move first to a position in alignment with the powder container for being filled and thereafter to a position in alignment with an inhalation channel, whereupon a unit dose will fall by gravity from the dosing recess into the inhalation channel. Thereafter the dose of medication is inhaled from the inhalation channel. These devices have the drawback that they make overdosing of the medicament possible by allowing the dispensing of a plurality of doses in succession into the
2

inhalation channel, whereby a multiple dose may be drawn by one
inhalation.

Inhalation devices having a metering plate movable between a
filling and a dispensing position are described e.g. in patent
EP 0 546 996, WO 94/04210 and US 5,161,524. A further example of
prior art devices is given in WO 93/03782. However, these de-
vices also suffer from a drawback that they make overdosing
possible by allowing the dispensing of a plurality of doses into
the inhalation channel.

Attempts have been made to solve this problem by using inhalers
or dispensing systems in which the dosing recess will not be
emptied into the inhalation channel by gravity but, instead, the
dose of medication is inhaled directly from the dosing recess,
such recesses having been notched into the surface of a metering
member. The metering member may have the shape of a cylinder, a
cone or a truncated cone, as disclosed in patent publications
WO 92/00771 and WO 92/09322. Furthermore, in these devices, the
metering member having the shape of a cylinder, a cone or a
truncated cone is disposed in a chamber having precisely the
same shape. When the metering member is rotated, the dosing re-
cesses will move first to a position in alignment with the me-
dicament container for filling, and then to the inhalation chan-
nel which is shaped so that the respective dosing recess will be
emptied under the effect of the air flow being inhaled, and
thereafter, having rotated through a full 360°, back to a posi-
tion in alignment with the medicament container. The lower sur-
face of the chamber wall may also have an emptying aperture from
which any powdered medicament possibly left in the dosing recess
will fall out during the said rotation.
In the rotating dispensing devices described above, the distance from the filling position to the inhalation position is less than 90° of a circle arc. Since the metering member is, for purposes of metering precision, disposed within a chamber of the same shape, and since it has to be rotated through 360°, of which at least 270° are useless for the actual function of the inhaler, in these devices particles will inevitably fall onto the slide surface between the metering member and the chamber. Thereby the rotation of the highly sensitive metering member will be disturbed and may even be completely obstructed. The metering member jamming inside the chamber will hinder the functioning of the whole device. Vigorous shaking or tapping will only increase the jamming as more powder flows into the gap between the chamber and the metering member.

An improvement of the powder inhalers of the prior art is suggested in EP 0 758 911. The described powder inhaler comprises a powder container, an air channel through which air is drawn via a mouthpiece, and a metering member equipped with a dosing recess, the metering member being movable in its longitudinal direction between a first position, in which the dosing recess is filled with powder coming from the container, and a second position, in which the filled dosing recess is brought into the air channel, the powder being maintained in the recess by the support of the recess bottom, and the air channel being directed to introduce the air flow into the bottom of the dosing recess during inhalation whereby the powder is released directly from the dosing recess. According to the powder inhaler of this prior art document, the metering member is a metering strip which is disposed on a flat surface and moves along said flat surface. When moving between the filling and the inhalation positions, the metering strip travels over an aperture for remnants, at which time any powder possibly remaining between the metering strip and the flat surface will fall out through the aperture. This
powder inhaler is still not completely satisfying for the following reasons, e.g. the inhaled air flow is directed longitudinally relative to the metering strip. In this condition both the deaggregation of the powder and the removal of the powdered medicament from the metering strip by the inhaled air flow is not efficient. Moreover, any powder possibly left after the inhalation will remain in the air channel until the metering strip again moves along the flat surface into the air channel during a subsequent inhalation process. This remaining powder could be accidentally inhaled by the patient. Furthermore, the powder remaining inside the air channel may deteriorate due to the friction between the surfaces.

As mentioned above, a further problem with respect to powder inhalers is that a sufficient deaggregation of the powder and the removal of the powdered medicament or drug from the metering member by the inhaled air flow is not satisfactory in the powder inhalers of the prior art.

It is generally known that the size of medicament particles should be 1-5 microns, preferably 2-3 microns, for the best possible penetration into their destination, i.e. deep into the lungs. The most common metering device is a so-called inhalation aerosol which is quite readily capable of reaching the optimal particle size. In addition to inhalation aerosols, an increasing number of powder inhalers are presently in use as these offer certain benefits, e.g. there is no need for ozone-destroying propellants. However, a drawback of powder inhalers is that a powdered medicament issuing from the powder inhaler has a too large particle size. Thus, most of the medicine dosage coming out of inhalers is retained in upper respiratory passages which, with certain medicines, can cause serious side effects. The medicine dosages required for different inhalation medicaments vary considerably, the smallest being approximately 0.01 mg and the largest 20 mg. When small amounts of medicine are metered in
powdered form, it is generally necessary to use some adjuvant or carrier, so that the sufficiently precise measuring of a dosage would be possible with the present technology. No matter if the dosage comprises just medicine or has a carrier admixed therein, the medicine dosage substantially comprises inter-adhered particles and most of these agglomerates are too large to penetrate into the lungs. As the agglomerates are released in a powder inhaler into an air flow directed into the lungs of a patient, there will occur some dispersal of these particle deposits, said dispersal resulting from the formulation of a powdered medicament and the construction of an inhaler. It is known that constructions creating a strong turbulence are capable of more effective pulverization.

In practice, however, no prior known powder inhalator structure and/or medicine formulation has produced results that would be equal to those achieved by an ordinary inhalation aerosol. It has been suggested as a partial solution that inhalation should be effected with as much force as possible, whereby the turbulence and pulverization of particles would accordingly be most effective. However, a quick inhalation is difficult for a person suffering e.g. from serious asthma and, on the other hand, a quick inhalation increases the residue in upper respiratory tracts. According to studies, pulverization of agglomerates is indeed intensified but the overall benefit is marginal. Excellent Pulmonary penetration in relation to the adherence of medicament to the upper respiratory tracts has been achieved by slow inhalation, corresponding to a flow rate of approximately 30 l/min or 0.5 l/sec.

Finnish patent application No. 87100 discloses a powder inhalator which has been designed in an effort to produce a clearly defined turbulence for pulverizing agglomerations of medicine. The centrally directed deflectors inside the device or the helical chute are explained to set the air flow in a spinning mo-
tion, whereby the medicine particles entrapped in the air abrade as a result of centrifugal force against the walls of the structure as well as collide with each other with resulting pulverization. The device described in the cited application has been marketed under the tradename Turbohaler® (Draco, Sweden), and the pulverizing structure therein is, as described above, a helical chute or groove. Laboratory tests indicated that this device had a relatively good pulverization of agglomerates of medicine which, however, could still be improved and intensified. In view of the pulverization of agglomerates or accumulations of medicine, there are a few defects in this device. The helical groove has in the centre thereof an open space having less air resistance than inside the groove. Accordingly, the flow rate of air and centrifugal force on the circumference of the groove are less than theoretically possible. Since the particles advance in the groove under a force caused by air resistance and centrifugal force tends to push the particles perpendicularly to the circumferential tangent, the actual force applied to the particles is a resultant of these forces and is applied diagonally relative to the circumferential tangent. Thus, the centrifugal force resulting from the spinning motion cannot be utilized in full extent for the pulverization of accumulations. In all deflector structures according to the cited application, the particles escape from the device within a few thousandths of a second when using conventional inhalation rates of 30-90 l/min, and that is a very short time for an effective pulverization. The residence time can be prolonged, e.g. by increasing the number of helices in groove portions or the number of separate deflector structures or the length of zigzagging air flow channels, but this would complicate manufacturing and cleaning and medicine residues in the actual device would increase. After all, cleaning of the structures disclosed in the cited application is difficult.
EP 0 407 028 shows in its Figs. 5 to 7 a vortex chamber into which an inlet tangentially directs air with a pulverized medicament. The dispersion air/medicament leaves the chamber axially at an outlet. Nothing is said about the diameter of the chamber.

FR-A-2352556 shows a cylindrical vortex chamber closed at one end, operated by the action of inhalation and having one tangential air-medicament inlet duct, an additional tangential air inlet duct and an axial outlet duct near the same end of the chamber as the inlet duct. The outlet is formed by a tube-like connection extending beyond the zone of the inlet duct and impeding the air flow. Nothing is said again about the diameter of the cylindrical chamber.

Moreover, in the device described in EP 0 477 222 a powdered medicament intended for inhalation is pulverized on the basis of a sufficiently powerful centrifugal force prior to or during inhalation. The centrifugal force is produced by the action of inhalation. A powdered medicament is entrapped in a gas flow and forced in a substantially circular or rotationally symmetrical space to such a powerful rotating motion that an effective splitting of accumulations of medicine is obtained. This is effected in a rotationally symmetrical chamber whose largest internal diameter can be 20 mm. The optimum diameter of the vortex chamber operating by the action of inhalation is 10-20 mm. If the diameter is increased, the pulverization effect deteriorates in a manner that, with a diameter of more than 30 mm, the pulverization effect is no longer significant.

From EP 0 865 302, a further powder inhaler is known which comprises a medicament container for storing a dry powdered medicament, a mouthpiece being covered by a removable protective cap, and a movable dosing slide having a dosing cavity to be placed underneath a funnel outlet of the medicament container for fill-
ing. With opening of the protective cap, the movable dosing slide with the filled dosing cavity is pushed into a shutter, and the dosing cavity is closed thereby. Upon a sufficient suction force generated by an inhalation process, a valve shield pushes away the shutter, and the dosing cavity is released in order to enable inhalation of the medicament powder. Furthermore, means are provided for enabling the return of the dosing slide only after a correctly completed inhalation process. A recording unit records the number of correctly performed inhalations and blocks the powder inhaler after a predetermined number of inhalations. Between the inlet and the outlet of the mouthpiece a labyrinthine atomiser path for powder deagglomeration is provided.

The technical problem underlying the present invention is to eliminate the above drawbacks of hitherto known powder inhalers and to provide a powder inhaler with an improved functionality. In particular, it is an object underlying the present invention to provide a powder inhaler with an improved dosing ability, whereby unintended dosing can be avoided, and to provide a powder inhaler with an optimal pulverization of agglomerates of a medicament to be inhaled, respectively.

This technical problem is solved by a powder inhaler having the features defined in claim 1. Furthermore, a deagglomerator arrangement having the features of claims 59 or 75 is proposed, which may be incorporated into such a powder inhaler without, however, being limited to this preferred use. The dependent claims define preferred and advantageous embodiments of the present invention.

According to the present invention, a powder inhaler is provided comprising a container or a powder reservoir for storing a powdered medicament, a metering member having a dosing recess to be filled with one dose of the powdered medicament, a mouthpiece
being in communication with an inhalation channel, and a protective member being provided between the metering member and the inhalation channel. The metering member is moveable between a filling position, in which the dosing recess is in alignment with an opening of the container so as to be filled with one dose of the powdered medicament, and an inhalation position, in which the dosing recess is in alignment with the inhalation channel. The protective member is moveable between a closed position, in which it at least covers the dosing recess when the metering member is in the inhalation position, thereby preventing the powdered medicament contained in the dosing recess from entering into the inhalation channel, and an open position, in which the protective member does not cover the dosing recess, thereby exposing the dosing recess to the inhalation channel so as to enable inhalation of the dose of the powdered medicament contained in the dosing recess.

The protective member, which is preferably a thin plate sliding on the metering member between its closed position and its open position, prevents the drug or powdered medicament contained in the dosing recess from falling out of the dosing recess, thereby preventing an unintentional loss of the powdered medicament contained in the dosing recess. Although a manual movement of the protective member between its closed position and its open position is conceivable, the protective member is preferably automatically withdrawn and moved from its closed position into its open position upon an inhalation process, thereby enabling the powdered medicament contained in the dosing recess to be released into the inhalation channel. Thus, the powder inhaler can be used in a variety of orientations, even upside down when the user or patient is lying in a bed, for example. This is a distinctive advantage over prior art products where the dose can be lost by a poor orientation of the device.
In order to automatically withdraw the protective member from its closed position and move it into its open position upon an inhalation process, an inhalation actuated mechanism may be provided being coupled to the protective member such that, if the protective member is in its closed position, the inhalation actuated mechanism moves the protective member into its open position if the inhalation suction force of the respective user exceeds a predetermined value. Preferably, the inhalation or breath actuated mechanism is constructed such that it automatically returns the protective member into its closed position after the respective inhalation process has properly been completed.

The powder inhaler of the present invention may comprise a casing and an integral cover, the integral cover being rotatably or pivotably coupled to the casing so as to enable opening and closing of the integral cover. The casing may comprise a window or an opening for displaying the number of doses taken or the number of doses left in the container, this number being counted by a dose counting unit. If the cover is closed it covers the mouthpiece being located at the upper side of the casing. The casing may also comprise an opening for a mark, for example in the form of a flap, which shows if a dose is ready for inhalation or not. In particular, this flap may disappear upon completion of an inhalation process, thereby showing that the respective dose has been taken by the user.

The container storing the powdered medicament is preferably divided up into a medicament chamber storing the powdered medicament and a desiccant chamber for storing a desiccant provided for drying out the powdered medicament contained in the medicament chamber, the desiccant chamber being separated from the medicament chamber by a separate permeable membrane. The permeability of the membrane is different from, in particular greater than that between either the desiccant or the medicament and the
outside world. This can be achieved, for example, by making the membrane from a different material and/or a thinner material than the main body of the container. Both the medicament chamber and the desiccant chamber may be sealed by foils. The bottom of the medicament chamber may have a dosing opening so that the powdered medicament contained therein can be filled into the dosing recess of the metering member supported by gravity, if the metering member is in its filling position. The filling process is furthermore supported by an appropriate shape of the medicament chamber which should have a cross-section diameter gradually decreasing from its top to its bottom, thereby forming a funnel for the powdered medicament.

The metering member is preferably a slide or a shuttle which is provided within the casing slidingly moveable in the horizontal direction between the filling position and the inhalation position. In the filling position the dosing recess faces the dosing opening of the container, and in the inhalation position the dosing recess faces an inhalation opening of the inhalation channel being in communication with the mouthpiece. The slide is preferably coupled to the cover such that opening of the cover causes the slide to move from the filling position forward to the inhalation position, and closing of the cover causes the slide to move from the inhalation position backward to the filling position. Projections, for example in the form of bolts, may be formed at both longitudinal sides of the slide, these projections engaging with profiled cam tracks being formed at the respective sides of the cover. This allows that the fundamental operating sequence of the powder inhaler is just to open the cover, inhale the dose, and close the cover. It is the simplest possible operating sequence to reduce training time on the one hand and maximize patient compliance on the other hand.

The coupling between the cover and the metering member is preferably such that opening of the cover by a predetermined first
angle from its closed position, preferably by an angle up to about 30°, does not actuate the metering member at all. Within this range of movement of the cover is slack where no mechanism is driven. Furthermore, the coupling between the cover and the metering member is preferably such that the metering member is moved in its inhalation position already a predetermined second angle prior to the cover being fully open. For example, the metering member may be placed in its inhalation position already when the cover has been opened by about 90° from its closed position. Between an opening angle of about 90°-135°, for example, there is free play again. This ensures that, should the user attempt a discrete operation of the device, the drug or the powdered medicament is ready already prior to the mouthpiece becoming exposed to the user.

The dosing recess may be designed to maximize the accuracy of gravitationally filling the dosing recess with the powdered medicament and also to maximize the ease of airborne entrainment of the formulation upon inhalation. Therefore, the dosing recess may be a dosing cup which is circular in profile and has an elliptical cross-section, the diameter being preferably three times the depth thereof. This enables the inhalation airflow to scour the dosing cup effectively. The circular profile and the ratio of depth to top area also combines the lowest variability of filling (primarily associated with deep, narrow receptacles) and scraping upon leaving the container (primarily associated with shallow, wide receptacles).

The flat surface of the metering member may be provided with a slot so that, upon a backward movement of the metering member from its inhalation position to its filling position, any powdered medicament possibly left on the flat surface of the metering member outside the dosing recess is wasted through the slot and can fall into a waste bin being provided underneath the metering member so as to catch this resilient powdered medicament.
In this way, even when the dose was not completely inhaled by the user, there is no remaining medicament in the inhalation channel.

The inhalation actuated mechanism may comprise an inhalation actuated means/member, a resilient means/member, and a coupling means/member. The resilient member, which is preferably a spring, has a first end which holds the inhalation actuated member in a first position. In this condition, the aforesaid mark, which is preferably a flag, is visible through the respective opening of the casing, thereby indicating that a dose has not been taken and is ready for inhalation, respectively. The inhalation actuated member is preferably a flap. Upon forward movement of the metering member from its filling position to its inhalation position, the resilient member is charged up or tensioned, thereby releaseing the inhalation actuated member. The inhalation actuated member is arranged and constructed such that, in this condition, only a predetermined inhalation suction force of an user, however not blowing, can move the inhalation actuated member out from its first position into a second position. For example, in this case, the inhalation actuated member may pivot or rotate from its first position into its second position. Thereby, the mark of the inhalation actuated member disappears and is no longer visible through the respective opening in the casing, indicating to the user that a dose has been taken and, thus, currently no dose is ready for inhalation. The inhalation actuated member, the resilient member, and the coupling member are also arranged and constructed such that the inhalation actuated member holds the coupling member, which is preferably in the form of a yoke, in a first position. The coupling member is coupled to the protective member, and preferably also to the metering member.

When the inhalation actuated member is moved from its first position to its second position by a sufficient inhalation force
effected by a user, the coupling member is released and, by the
discharging effect of the resilient member, automatically moved
into its second position in which the coupling member automati-
cally moves the protective member from its closed position into
its open position, thereby releasing the dose contained in the
dosing recess. For example, the coupling member may have an arm
which is released upon movement of the inhalation actuated mem-
ber from its first position into its second position, and the
coupling member may also comprise a prolonged projection which,
on the one hand, engages an opening of the protective member
and, on the other hand, is slidingly moveable within a slid
formed in the metering member.

When the cover of the powder inhaler is closed again, the meter-
ing member is returned to its filling position, and the afore-
said dose counting unit is actuated and incremented. In particu-
lar, this is effected as follows:

The coupling member may comprise a further prolongation, pref-
erably in the form of a cantilever, which, when the metering
member causes the coupling member to move back from its second
position to its first position, actuates the dose counting unit.
In this regard, the dose counting unit may comprise a wheel ar-
 rangement having a plurality of wheels being numbered on one
 side facing the opening of the casing of the powder inhaler and
 being coupled to each other by a gear arrangement. In particu-
 lar, the wheel arrangement may comprise a plurality of wheels
 for displaying the numbers for a different order of magnitude,
 respectively. For example, the wheel arrangement may comprise a
 units wheel and a tens wheel being coupled by an idler wheel. On
 the other side of at least one wheel there may be arranged a
 plurality of drive teeth being arranged along the circumferen-
tial direction of the respective wheel. The above prolongation
 of the coupling member is moved over one of these drive teeth
 when the coupling member is moved from its first position to its
second position, thereby bringing the prolongation of the coupling member into engagement with the respective drive tooth. On the other hand, when the metering member is moved backward into its filling position (and the resilient member is thus allowed to discharge), the movement of the coupling member from its second position to its first position caused thereby results in the prolongation of the coupling member rotating the respective wheel by one step, thereby decrementing (or alternatively) the dose counting unit. At the same time, the coupling member also moves the protective member back into its closed position, and the resilient member returns the inhalation actuated member to its first position and holds it in this position, so that the mark attached to the inhalation actuated member is visible through the respective opening of the casing again. Furthermore by these movements the coupling member is again brought into engagement with the inhalation actuated member, and in particular in this condition the aforesaid arm of the coupling member is held by the inhalation actuated member again. Thereby, the initial condition of the inhalation actuated mechanism and of the device is resumed again, and the above described operation of the powder inhaler can be repeated.

The above mark or flag has a very useful function. It shows a user if he has already taken a dose, thereby removing the possibility of double dosing. Furthermore, only inhaled doses are displayed by the dose counting unit. This reduces wastage and gives the user a true indication of what has been inhaled. The above described dose counting unit is directly driven by a closing operation of the cover. This is more reliable than using a stored energy. However, the driving of the dose counting unit may well be assisted by the stored energy of the resilient member.

The inhalation channel through which air is inhaled in use is preferably designed to comprise a deagglomerator arrangement
comprising a substantially tangential air inlet, a preferably rotationally symmetrical vortex chamber, and an air outlet which is axially aligned with the vortex chamber such that airflow within the vortex chamber leads to a strong velocity gradient. The vortex chamber has a diameter \( d \) of \( 6 \text{mm} \leq d \leq 10 \text{mm} \), preferably \( 6 \text{mm} \leq d \leq 8 \text{mm} \), in particular about 8 mm, since such a diameter dimension has proven to be most effective for the deagglomeration function. The air outlet of the deagglomerator arrangement has preferably a smaller diameter than the vortex chamber. The base section of the vortex chamber may have a substantially elliptical cross-section, while the air outlet (and the inhalation channel) is preferably circular in cross-section. In addition or alternatively, the outer walls of the vortex chamber have the shape of arcs which are non-concentric to the inner diameter of the vortex chamber so as to achieve an improved deagglomerator functionality.

Besides the above described features of the powder inhaler of the present invention, a couple of variants could be incorporated into the powder inhaler as well. For example, a manual override mechanism could be incorporated into the inhalation actuated mechanism for manually moving the protective member and manually actuating the inhalation actuated mechanism. This would allow users who could not generate the required flow rate for actuating the inhalation actuated mechanism to manually release the dose contained in the dosing recess and trigger the dose counting unit. Furthermore, an extra part could be added to override the inhalation actuated dose counting unit. This would especially be of benefit to those people who cannot achieve the required flow rate to operate the inhalation actuated mechanism. Furthermore, a one-way valve could be placed in the inhalation channel, preferably over the inlet to the deagglomerator arrangement (cyclone). This could reduce any moisture blown into the inhaler by about 50 \%.
Furthermore, another means of delivering a dose upon inhalation could be incorporated. Such a means could comprise a resilient member, in particular in the form of a spring, which is compressed by means of opening the cover. The resilient member would act on the metering member. The metering member would be free to move to a half-way position between the container and the inhalation channel. However, the metering member would be held at this half-way position until an inhalation actuated mechanism releases the metering member to complete its travel to the inhalation channel and, hence, present the dose contained in the dosing recess for inhalation. The half-way position (mid-point position) would have the combined effect of retaining the dose in the dosing recess and protecting it from moisture from user exhalation or discharge.

The device of the present invention delivers consistent respirable dose values to the patient across a wide range of pressure drops. For example, between 30 l/min and 90 l/min the fine particle fraction, a measure of Pulmonary penetration varies by less than 20%. Furthermore, this performance only requires a low work input from the patient with the device being classified as a low to medium resistance device.

In the following, the present invention will be described by means of a preferred embodiment with reference to the attached drawings.

Figure 1 shows a perspective outside view of a powder inhaler according to a preferred embodiment of the present invention,

Figure 2 shows a perspective view of the powder inhaler when a cover thereof is opened,

Figure 3 shows a top view of the powder inhaler when the cover is opened,
Figure 4 shows a perspective view of a dosing sub-assembly of the powder inhaler,

Figures 5A and 5B show perspective views of a dose counting sub-assembly of the powder inhaler,

Figures 6A and 6B show cross-sectional views of the dosing sub-assembly,

Figures 7A and 7B show cross-sectional side views of the powder inhaler when the cover is closed,

Figure 8 shows a perspective side view of the powder inhaler without side labels when the cover is closed,

Figure 9 shows a perspective view of an inhalation actuated mechanism and a dose counting unit of the powder inhaler,

Figure 10 shows a partial cross-sectional view of the inner construction of the powder inhaler upon inhalation,

Figure 11 shows a perspective view of the inhalation actuated mechanism and the dose counting unit of Figure 9 upon inhalation,

Figure 12 shows a perspective view of the inhalation actuated mechanism and the dose counting unit of Figure 9 after closing the cover of the powder inhaler,

Figure 13 shows a perspective view of a slide of the powder inhaler,

Figure 14 shows a perspective view of the cover of the powder inhaler,

Figure 15 shows a perspective view of a part of a casing of the powder inhaler,
Figure 16 shows a perspective view of a mouthpiece of the powder inhaler,

Figure 17 shows a top view of the dosing sub-assembly shown in Figures 6A and 6B,

Figure 18 shows a perspective view of a slide spring of the powder inhaler,

Figure 19 shows a perspective view, a cross-sectional view, and a front view of an inhalation actuated member of the inhalation actuated mechanism,

Figure 20 shows a perspective view of a protective member of the powder inhaler,

Figure 21 shows a perspective view of a resilient member of the inhalation actuated mechanism,

Figure 22 shows a front view, a perspective view, and a rear view of a units wheel of the dose counting unit,

Figure 23 shows a rear view, a perspective view, and a front view of a tens wheel of the dose counting unit,

Figure 24 shows a perspective view of an idler wheel of the dose counting unit,

Figure 25 shows a perspective view and a side view of a coupling member of the inhalation actuated mechanism and the dose counting unit,

Figure 26 shows a perspective and schematic bottom view of a deagglomerator arrangement (cyclone),

Figure 27 shows a cross-sectional view of the deagglomerator arrangement of Figure 26, and
Figures 28A, 28B and 28C show a perspective view, a bottom view and a top view, respectively, of a dosing sub-assembly of a powder inhaler according to a further embodiment of the invention.

The powder inhaler shown in Figure 1 comprises a casing with a lower shell 1 and an integral cover 2 being pivotably or rotatably coupled to the lower shell 1. In a side surface of the lower shell 1 a window 4 is formed for displaying numbers of a dose counting unit which will be described later.

As can be taken from Figure 2, the integral cover 2 can be opened to reveal a mouthpiece 3 with which a user can inhale a powdered medicament. At the upper front side of the mouthpiece 3 slots 6 are formed which allow air inlet. Furthermore, at the upper side of the mouthpiece 3 an opening or a hole 5 is formed which allows to view a visible mark or flag showing if a dose is ready. As will be described later, this flag disappears upon inhalation showing that the respective dose has been taken.

The structure of the lower shell 1, the integral cover 2 and the mouthpiece 3 can also be taken from Figure 3 which shows a top view of the powder inhaler. In Figure 3 (and in Figure 8), the integral cover 2 is shown without side labels which are depicted in Figures 1 and 2. These side labels prevent access to profiled cam tracks being described later so as to protect these cam tracks from dust etc.

Figure 14, Figure 15, and Figure 16 show perspective views of the integral cover 2, the lower shell 1, and the mouthpiece 3, respectively. The lower shell 1 and the mouthpiece 3 are constructed such that the mouthpiece 3 can be snap-fitted onto the lower shell 1. From both side surfaces of the lower shell 1 projections or bolts extend which engage with respective central openings at both side surfaces of the integral cover 2, thereby allowing rotational movement of the integral cover 2 relative to the lower shell 1. As can be seen from Figure 1 and Figure 2,
the integral cover 2 is closed when its lower surface rests on
the upper rim of the lower shell 1, and the integral cover 2 can
be opened until its rear edge abuts against the underside of the
lower shell 1 (see Figure 2). At both side surfaces of the inte-
gral cover 2, openings 31 having the shape of profiled cam
tracks are formed which are coupled to side projections 28 of a
shuttle or slide 15, a perspective view thereof being shown in
Figure 13. This kind of coupling between the integral cover 2
and the slide 15 will be described later in detail.

Within the casing and the lower shell 1, respectively, there are
two sub-assemblies arranged. The first sub-assembly is a dosing
sub-assembly 13 which in particular meters a powdered medica-
ment, while the second sub-assembly is a dose counting sub-
assembly 14 which comprises an inhalation actuated mechanism and
a dose counting unit for counting the number of doses taken by
the user.

Figure 4 shows a perspective view of the dosing sub-assembly 13.
As can be seen, the dosing sub-assembly 13 comprises a container
or a reservoir 7 for storing a powdered medicament, the above-
mentioned slide 15 shown in Figure 13, and a deagglomerator ar-
angement 16 to be coupled to an inhalation channel of the
mouthpiece 3. A spring 17 is clamped onto side projections of
the dosing sub-assembly 13 such that it holds the dosing sub-
assembly together.

Figure 18 shows a perspective view of the spring 17. As can be
easily seen, the spring 17 comprises four resilient side spring
members, two spring members being fixed to the rear side and two
spring members being fixed to the front side of the spring 17.
All four spring members extend in the longitudinal direction of
the spring 17 such that their free ends are arranged in a middle
portion of the spring 17. These spring members apply a force to
the slide 15 such that the slide 15 is continuously urged
against the underside of the dosing sub-assembly 13. From the rear side to the front side of the spring 17, there extends an additional resilient spring member which applies a separate force to the longitudinal middle region of the slide 15. As is shown in Figure 13, in this longitudinal middle region the slide 15 has a dosing recess 18 in the form of a dosing cup for metering a dose of the powdered medicament and for transporting the dose from a filling position underneath the container 7 to an inhalation position underneath the deagglomerator arrangement 16. The above-mentioned separate spring member extending along the longitudinal middle region of the spring 17 ensures that the dosing recess 18 is reliably pressed against the underside of the dosing sub-assembly 13 if the slide 15 is in its inhalation position so that the dosing recess 18 is properly located under the deagglomerator arrangement 16.

As already indicated above, the slide 15 serves as a metering member which can be moved in the horizontal direction from a filling position shown in Figure 6A to an inhalation position shown in Figure 6B. Thus, the slide 15 is slidingly moveable between the filling position, where the dosing recess 18 is located underneath a dosing opening of the container 7 and faces the dosing opening, and the inhalation position, where the dosing recess 18 is located underneath and faces an inhalation opening of the deagglomerator arrangement 16 which is in communication with an inhalation channel (to be described later) of the mouthpiece 3.

As is shown in Figure 6A, the container 7 is a container with integral desiccant. The container 7 comprises a medicament chamber 8 storing a powdered medicament and a desiccant chamber 9 storing a desiccant for absorbing moisture that may have entered the medicament chamber 8. The desiccant chamber 9 is separated from the medicament chamber 8 by a separate permeable membrane 10. This permeable membrane 10 is of a different permeability
than the permeability between either the desiccant or the medicament to the outside world. The permeability of the membrane 10 can be achieved, for example, by making it of a different material and/or a thinner section than the main body of the container 7. Foils 11, 12 are used to seal both the medicament chamber 8 and the desiccant chamber 9. As a matter of course, other suitable sealing means may be used for sealing both chambers 8, 9 as well.

The above described integral desiccant system has the following advantages. The desiccant has only to dry out the medicament chamber rather than the whole device. This requires less desiccant reducing product size and cost. Furthermore, the desiccant is always sealed. This means that the desiccant will still be effective even if the cover is left open. The desiccant is stored in the separate sealed desiccant chamber 9. This reduces the risk of incorrect assembly if the desiccant used the same closure as the medicament. Moreover, the integral container 7 comprising both the medicament chamber 8 and the desiccant chamber 9 can be manufactured using a 2-shot moulding. This ensures a good seal between the medicament chamber 8 and the desiccant chamber 9 at low product cost. Finally, the foil sealing provides a tamper-proof means of filling the device with the medicament or drug which has a very low permeability and requires only little product space.

As is shown in Figure 6A and Figure 6B, the medicament chamber 8 has a gradually decreasing cross-section diameter from its top to its bottom so that the medicament chamber 8 is shaped like a funnel supporting an easier filling of the dosing recess 18 through the dosing opening formed in the bottom of the medicament chamber 8.

The powder inhaler shown in the drawings solves many technical problems that may occur throughout the life cycle of the powder
The fundamental operating sequence of the powder inhaler is to open the integral cover 2, inhale the dose of the powdered medicament, and close the integral cover 2.

The cover 2 is gripped by the user and opened. As already described above, the projections 28 formed at both longitudinal sides of the slide 15 (see Figure 13) engage with the respective side openings 31 formed at both sides of the cover 2. In particular, these side openings 31 are profiled cam tracks. The coupling between the profiled cam tracks 31 and the projections 28 is such that opening of the cover 2 causes the slide 15 to move forward from its filling position (Figure 6A) to its inhalation position (Figure 6B). Likewise, closing of the cover 2 causes the slide to move from its inhalation position backward to its filling position again. That is to say, by opening/closing the cover 2, the slide 15 is moved substantially linearly with respect to the casing. In particular, the profiled cam tracks 31 are shaped such that opening of the cover 2 by a predetermined first angle, for example, by an angle of about 30°, from its closed position does not actuate the slide 15. That is the first 30° of the movement of the cover 2 is slack where no mechanism is driven. The industrial design of the powder inhaler is intended to convey the correct orientation of use. Furthermore, the coupling between the cover 2 and the slide 15 is such that the slide 15 is properly moved to its inhalation position already a predetermined second angle, prior to the cover 2 being fully opened. For example, the slide 15 may be moved to its inhalation position already when the cover 2 has been opened by 90°. In a range of 90°-135°, e.g., there is again free play. Therefore, the dose of the powdered medicament filled in the dosing recess 18 is correctly presented to the deagglomerator arrangement 16 as well as the respective inhalation channel coupled thereto, ready for inhaling, 90°-45° prior to the cover 2 being fully open (An opening angle of 180° is considered
25

as representing a fully open position of the cover). This ensures that the dose will be ready prior to the mouthpiece 3 becoming exposed to the user if the user should attempt a discrete operation of the powder inhaler, for example. There is an audible click indicating that the cover 2 is fully open.

When the cover 2 is closed, there are for example 45° of free play before a further closing of the cover 2 moves the slide 15 from the inhalation position to the filling position. Before the cover 2 is completely closed, there may be 15° of free play, for example. It should be noted that the profiled cam tracks 31 shown in the drawings are only exemplary.

As already mentioned before, the dosing recess 18 has the shape of a dosing cup which is designed to maximize the accuracy of gravitationally filling the dosing cup and maximize the ease of airborne entrainment of the formulation upon inhalation. The dosing cup is circular in profile (in top view) with an semi-elliptical cross-section (i.e. the cross-section has the shape of the half of an ellipse), the diameter being three times the depth. This enables the cyclonic airflow in the airway of the deagglomerator arrangement 16 to scour the dosing cup 18 effectively. The circular profile and the above-mentioned ratio of depth to top area also combines the lowest variability of filling and scraping upon leaving the container 7.

During opening the slide 15 is moved from the filling position to the inhalation position as well as after the slide 15 has reached its inhalation position, the dose of the powdered medicament filled in the dosing recess 18 of the slide 15 is prevented from falling out by a protective member, i.e. a dose protector 19. The dose protector 19 is arranged slidingly moveable on the slide 15 between a closed position and an open position. In its closed position, the dose protector 19 at least completely covers the dosing recess 18 when the slide 15 is in the
inhalation position, while in its open position the dose protector 19 exposes the dosing recess 18 to the deagglomerator arrangement 16 and the inhalation channel when the slide 15 is in its inhalation position. The dose protector 19 is held in its closed position by an inhalation or breath actuated mechanism which will be described later. This inhalation actuated mechanism is constructed such that the dose protector 19 is moved from its closed position to its open position only if the inhalation suction force exerted by the user in the inhalation channel exceeds a predetermined level. Furthermore, the inhalation actuated mechanism is arranged such that only an inhalation suction breath not a blowing breath, can actuate the inhalation actuated mechanism and cause a movement from the dose protector from its closed position to its open position.

In the following, the inhalation actuated mechanism in combination with the dose protector and the dose counting unit is described in detail.

Figure 5A and Figure 5B show perspective views of the dose counting sub-assembly 14 already mentioned above. The dose counting sub-assembly 14 consists of a sub-frame 20 which holds a flap 21 acting as an inhalation actuated member, a yoke 22 acting as a coupling member and a drive spring 23 acting as a resilient member. The drive spring 23 drives the dose counting unit which, in the present case, comprises an units wheel 24 and a tens wheel 26 being coupled by an idler wheel 25. Furthermore, the drive spring 23 drives the dose protector 19. The units wheel 24 and the tens wheel 26 display the number of doses remaining in the medicament chamber 8. As a matter of course, the drive spring 23 may be replaced with a resilient means being constituted by a plurality of spring elements or spring parts, for example.
In Figure 7A and Figure 7B partial cross-sectional views of the whole powder inhaler along different cross-sectional lines with the cover 2 being dosed are shown. In particular, from Figure 7A it can be seen that the mouthpiece 3 comprises the inhalation channel 27 extending from the upper side of the mouthpiece 3 downward so as to be coupled to the deagglomerator arrangement (cyclone) 16 of the dosing sub-assembly 13.

The functionality of the inhalation actuated mechanism as well as the dosing counting unit is as follows.

As shown in Figure 13, there are formed recesses 30 at both front corner portions of the slide 15. At one of these recesses 30, a prolonged end 34 of the drive spring 23 engages with the slide 15 if the slide 15 is moved forward. By the contact with the slide 15, the drive spring 23 of the inhalation actuated mechanism is tensioned and charged up. A first end 33 of the drive spring 23 rests at a portion 41 of the flap 21 when the drive spring 23 is in its discharged state. Therefore, by charging up the drive spring 23 this reset force exerted by the first end 33 of the drive spring 23 on the flap 21, normally holding the flap 21 in a first horizontal position shown in Figure 9, is released.

Figure 19 shows different perspective views of the flap 21. As can be seen, at the upper surface of the flap 21, a flag 38 is formed which acts as a mark being visible through the opening 5 in the mouthpiece 3 when the flap 21 is in its first horizontal position, whereby indicating that a dose is ready for inhalation. Furthermore, the flap 21 comprises a feature 40 for engagement with an arm 43 of the yoke 22. Finally, the flap 21 also comprises two projections 39 which act as a counterweight. This counterweight balances the flap 21 reducing not only the actuation force required but also the susceptibility of the mechanism to accidental triggering.
As shown in Figures 9 and 21, the drive spring 23 has a second end 32 which rests on a lateral side surface 48 of the yoke 22.

Figure 25 shows a perspective view and a side view of the yoke 22. The yoke 22 has a shaft-like portion 42 on which the drive spring 23 is mounted. Furthermore, in Figure 25 the arm 43 is depicted whose upper end 44 is retained and released, respectively, by the flap 21. At that lateral side of the yoke 22 which is opposite to the lateral side surface 48 on which the second end 32 of the drive spring 23 rests, there is formed a projection 45 having a thickening 46 at its end for operating the dose counting unit which will be described later. From the bottom of the yoke 22, there extents a prolongation 47 which engages, on the one hand, with an opening formed in the dose protector 19 and, on the other hand, with a slit 29 formed in the front end portion of the slide 15 (see Figure 13 and Figure 20).

As already described above, when the drive spring 23 is decompressed and discharged, its end 33 exerts a reset force on the portion 41 of the flap 21, thereby holding the flap 21 in its first horizontal position, as shown in Figure 9. In this condition, the dosing protector 19 prevents the powdered medicament contained in the dosing recess 18 from being displaced from the deagglomerator arrangement 16 (cyclone) if the user blows into the mouthpiece 3. Furthermore, the flap 21 provides a resistance if the user blows into the device giving positive feedback.

If, however, the slide 15 is pushed forward by opening the cover 2, thereby compressing and charging the drive spring 23, the reset force exerted by the end 33 of the drive spring on the flap 21 is released, and the flap 21 can be rotated from its first horizontal position shown in Figure 9 into a second position being pivoted relative to the first position if there is a sufficient high inhalation suction force effected by the user in the inhalation channel 27 of the powder inhaler.
In the latter case, the flap 21 is moved by this sufficient high inhalation force from its first position shown in Figure 9 into its second position shown in Figure 10. As can be also seen from Figure 10, by this movement of the flap 21 the arm 43 of the yoke 22 is released. This enables the drive spring 23, due to its compression, to move its second end 32, which is in engagement with the lateral side surface 48 of the yoke 22, and thus the yoke 22 slightly upward. By this rotational upward movement of the yoke 22 the prolongation 47 extending from the upper side of the yoke 22 is moved forward, thereby moving the dose protector 19 from its closed position to its open position. This situation is shown in Figure 10 as well as in Figure 11.

In Figure 20, a perspective view of the dose protector 19 is shown. In particular, in Figure 20 the opening 36 is shown which is in engagement with the prolongation 47 extending downwardly from the bottom of the yoke 22. The front end 35 of the dose protector 19 has a partial circular or semicircular shape so that it can form part of the wall of the deagglomerator arrangement or cyclone 16 when the dose protector 19 is in its closed position.

Since the dose protector 19 has been moved out from its closed position into its open position by the yoke 22, the dosing recess 18 of the slide 15 is exposed to the inside 50 of the cyclone, and the dose of the powdered medicament contained in the dosing recess 18 can be inhaled through the cyclone and the inhalation channel 27 as well as the mouthpiece 3. In the cyclone, the drug or the powdered medicament is entrained into a swirling airflow where the active part of the formulation is disaggregated from the carrier (see reference sign 49).

Furthermore, since the flap 21 has been moved from its first horizontal position (see Figure 9) to its second position rotated or pivoted relative to its first position (see Figures 10
and 11), the flag 38 formed at the upper surface of the flap 21 is no longer visible through the opening 5 in the upper side of the mouthpiece 3. That is the flag 38 has disappeared thereby indicating that a dose has been taken, and a new dose is not ready for inhalation again, yet. Figure 17 shows a top view of the dosing sub-assembly, depicting the inside 50 of the cyclone as well as portions 51 (corresponding to side walls 78 depicted in Figure 26) which allow the mouthpiece to be assembled, portions 52 which produce a cyclonic airflow within the cyclone, and projections 53 for mounting the dosing sub-assembly within the lower shell 1 of the powder inhaler. Furthermore, in Figure 17 there is also depicted the end stop 37 for the prolongation 47 of the yoke 22 and the dose protector 19, respectively.

In the following, the functionality of the dose counting unit is explained in detail. As already mentioned above, the dose counting unit, being provided for counting the number of doses taken (up counter) or, alternatively, the number of doses remaining in the container (down counter), comprises the units wheel 24 and the tens wheel 26 being coupled to one another by the idler wheel 25.

Figure 22 shows a front view, a perspective view, and a rear view of the units wheel 24. The units wheel 24 comprises a central opening 54 at which it is rotatably mounted at the dose counting sub-assembly 14 inside the casing of the powder inhaler as shown in Figures 5A and 5B, for example. Reference sign 55 designates a feature which provides a thrust-bearing surface with the lower shell 1. Reference sign 56 designates numbers which are printed on the outer surface of the units wheel 24 along the circumferential direction thereof and with equal intervals therebetween. At the outer periphery of the units wheel 24, there are formed teeth 57 for driving the idler wheel 25. As
can be taken from the rear view of the units wheel 24, these teeth 57 are formed diametrically opposed to each other. Finally, on the back of the units wheel 24 there are drive teeth 58 which are brought into engagement with the projection or cantilever 45 of the yoke 22 so as to drive the units wheel 24 step by step upon completion of an inhalation process. As can be easily seen from Figure 22, the drive teeth 58 each are inclined in the circumferential direction of the units wheel 24. For example, the diameter 59 of the units wheel 24 may be about 20 mm.

Figure 23 shows a rear view, a perspective view and a front view of the tens wheel 26. On the back of the tens wheel 26, there is formed a plurality of teeth 62 in the circumferential direction of the tens wheel 26. These teeth 62 are driven by the idler wheel 25. Reference numeral 60 designates missing teeth which prevent a drive of the tens wheel 26 when the medicament chamber 8 is empty, that is the tens wheel 26 is constructed such that during one life cycle of the powder inhaler nearly one complete rotation of the tens wheel 26 is effected by the dose counting unit. Reference numeral 61 designates an end stop with the casing of the powder inhaler. The diameter 63 of the tens wheel 26, for example, may be about 25 mm. Reference numeral 64 designates an opening at which the tens wheel 26 is rotatably mounted at the dose counting sub-assembly 14, as shown in Figures 5A and 5B, for example. Reference numeral 65 designates a feature which provides a thrust-bearing surface with the casing of the powder inhaler. Furthermore, reference numeral 66 designates a feature which provides a thrust-bearing surface with the lower shell 1, and reference numeral 67 designates the periphery of the opening 64 which is located on the casing of the powder inhaler. On the outer surface of the tens wheel 26, there are formed two circumferential rows of numbers 68. These two rows of numbers display tens and hundreds numbers in correct orientation. In each case,
a combination of a units number of the units wheel 24 with a tens number and a hundreds number of the tens wheel 26 is visible through the opening 4 formed in the lower shell 1 of the powder inhaler (see Figure 1, for example). Each such combination of horizontally adjacent numbers of the units wheel 24 and the tens wheel 26 designates a corresponding number of doses remaining in the medicament chamber 8. Finally, at the outer periphery of the tens wheel 26, there is also formed a projection 69. Along the radial direction of this projection 69, there are no tens and hundreds numbers formed on the outer surface of the tens wheel 26, and this projection 69 covers the units wheel 24 if the medicament chamber 8 is empty such that no numbers are visible through the opening 4 of the lower shell 1, thereby indicating to the user that there is no dose remaining in the medicament chamber any more.

Figure 24 shows a perspective view of the idler wheel 25. The idler wheel 25 has a shaft 72 at which it is rotatably mounted on the sub-frame 20 of the dose counting sub-assembly 14 as shown in Figures 5A and 5B, for example. Furthermore, the idler wheel 25 has half-width teeth 70 which engage with the drive teeth 57 on the back of the units wheel 24. Furthermore, the idler wheel 25 comprises full width teeth 71 which lock against the units wheel 24. When the units wheel 24 is set to numbers "1"-"9" (reference numeral 56 in Figure 22), the teeth 57 on the back of the units wheel 24 fit between the full-width teeth 71 of the idler wheel 25. When the units wheel 24 is set to number "0", however, the teeth 57 engage with the half-width teeth 70.

As has been explained above, the coupling between the units wheel 24 and the tens wheel 26 is such that after each ten step-wise rotations of the units wheel 24 the tens wheel 26 is rotated by one step, thereby increasing the combination of tens and hundreds numbers on the outer surface of the units wheel 24. It should be noted that Figure 5A shows a situation in which no
numbers of the units wheel 24 and the tens wheel 26 are visible through the opening 4 of the lower shell 1, since the projection 69 of the tens wheel 26 covers the respective number of the units wheel 24 so as to indicate that the medicament chamber 8 is empty.

As described above, when the flap 21 is rotated from its horizontal first position to its second position upon an inhalation process initiated by the user (see Figure 11), the yoke 22 is slightly rotated clockwise (in Figure 11) so that the dose protector 19 is moved from its closed position to its open position. Furthermore, by this clockwise rotation of the yoke 22, the projection or cantilever 45 of the yoke 22 is also slightly moved clockwise along the inclination of the next drive tooth 58 of the units wheel 24 so as to bring the thickening 46 of the cantilever 45 into engagement with the respective drive tooth 58. Up to this point, no actuation of the units wheel 24 and the tens wheel 26 has taken place.

After inhalation, the user closes the cover 2 of the powder inhaler. With the closing of the cover 2, the slide 15 is moved backward from its inhalation position to its filling position by means of the coupling between the projections 28 of the slide 15 and the profiled cam tracks 31 of the cover 2.

As shown in Figure 12, this backward movement of the slide 15 causes a counter clockwise rotation (as regards the view depicted in Figure 12) of the yoke 22, since the prolongation 47 of the yoke 22 is moved together with the slide 15 backward. The counter clockwise rotation of the yoke 22 is supported by the drive spring 23 which is allowed to be discharged and decompressed upon backward movement of the slide 15. Due to this counter clockwise rotation of the yoke 22, the cantilever 45 is also rotated counter clockwise, thereby also rotating the units wheel 24 counter clockwise by one step (as regards the view de-
picted in Figure 12) which results in decreasing the number of doses left in the medicament chamber 8, which is visible through the opening fear of the lower shell 1. As a matter of course, the dose counting unit can also be arranged such that it does not display the number of doses remaining in the medicament chamber 8, but the number of doses already taken by the user.

Furthermore, since the yoke 22 and the drive spring 23 are moved back into their initial positions, the end 33 of the drive spring 23 urges the flap 21 back into its horizontal first position (as shown in Figure 12), thereby resetting the flag 38. Moreover, in this situation, the yoke 22 is again held by the engagement of its arm 43 with the feature 40 of the flap 21. Thus, the whole powder inhaler has been transferred into its initial position again.

Another advantage associated with the flag 38 is that it may be pushed down by fingers of a user so as to effect a manual override of the inhalation actuated mechanism. This would enable the user to take the dose if the user is not able to generate a sufficient force in order to actuate the inhalation actuated mechanism.

On completely closing the cover 2, there will be an audible click signalling that the cover is closed. Preferably, the powder inhaler will require the cover 2 to be completely closed to function correctly.

Finally, the deagglomerator arrangement 16 (cyclone) of the powder inhaler should be briefly discussed.

The purpose of this deagglomerator arrangement is to produce clearly defined turbulences within the inhalation channel 27 so as to pulverize agglomerations of the medicament. By the swirling airflow within the deagglomerator arrangement 16, the active
part of the formulation is disaggregated from the respective carrier.

In principle the deagglomerator arrangement of the present invention comprises a rotationally symmetrical vortex chamber, at least one substantially tangential air inlet, and an outlet or exit for outputting the air with the deagglomerated powdered medicament, the outlet being spaced from the air inlet in the longitudinal axial direction of the vortex chamber. The general structure of the deagglomerator arrangement, for example, can be taken from Figure 6A, Figure 6B, Figure 7A, and Figure 17.

In addition, the structure of the deagglomerator arrangement, which is generally not restricted to the usage in powder inhalers as described in connection with the above drawings, will be explained in detail by reference to Figure 26 and Figure 27, respectively.

Figure 26 shows a schematic perspective view of the deagglomerator arrangement (cyclone) according to a preferred embodiment, the deagglomerator arrangement being constructed to produce a very strong cyclonic flow within the deagglomerator arrangement resulting in a very strong velocity gradient. It is to be noted that Figure 26 is a schematic view to explain the design and the key features of the cyclone according to the preferred embodiment, but Figure 26 does not show the cyclone exactly as it is implemented in the above described powder inhaler. The implementation of the cyclone in the powder inhaler, for example, can be taken from Figure 4 or Figure 17.

As can be seen from Figures 26, the deagglomerator arrangement 16 has an opening in its bottom which disemboques into a vortex chamber 73. The vortex chamber 73 is designed substantially rotationally symmetrical. In addition, there are two air inlet conduits 75 which direct air substantially tangentially into the vortex chamber 73. As can be seen from Figure 26, air inlet win-
dows 76 are formed in the upper surface of the air inlet conduits 75 which for example cover 80° of the air inlet conduits 75 so as to enable entry of air from above into the air inlet conduits 75. The base section of the vortex chamber 73, in general, has an elliptical cross-section. As depicted in Figure 26, there is an outlet 74 formed in the upper end of the vortex chamber 73, the outlet 74 being spaced from the opening for the supply of the powdered medicament and both air inlets in the longitudinal axial direction of the vortex chamber 73. In particular, the outlet 74 is aligned coaxially to the longitudinal central axis of the vortex chamber 73 and extends along this longitudinal central axis. In particular, the outlet 74 has a substantially circular cross-section, the diameter of this circular cross-section being smaller than the diameter of the elliptical cross-section of the vortex chamber 73.

Figure 27 shows a cross-sectional view of the deagglomerator arrangement in accordance with a sectional plane which horizontally intersects the deagglomerator arrangement 16 and the vortex chamber 73, respectively. The view depicted in Figure 27 could be regarded as a bottom view of the deagglomerator arrangement 16 as well.

In Figure 27, the vortex chamber 73 and the circular outlet 74, which disemboques into the inhalation channel 27, are depicted. Although the horizontal cross-section of the base section of the vortex chamber 73 has a substantially elliptical shape, the cross-section of the base section of the vortex chamber 73 can also be defined by an imaginary circle which can be laid inside the base section of the vortex chamber 73 on a horizontal plane intersecting both air inlet conduits 75 such that the circumference or periphery of this circle just touches the inside surfaces of both side walls 78 at places which are diametrically opposed to one another without extending beyond the side walls of the vortex chamber 73. In particular, these diametrically op-
posed places of the side walls 78 are those places of the side-walls 78 where both air inlets of the air inlet conduits 75 dis-embogue into the vortex chamber 73. In Figure 27, the respective "base" circle is depicted with a dotted line and assigned with reference numeral 77. The air inlets are arranged along the circumferance of the "base" circle 77.

Studies have shown that the diameter $d$ of this "base" circle 77 has an important influence on the deagglomerator effect of the cyclone. For example, if the diameter $d$ is too small, the flow resistance is too high. On the other hand, if the diameter $d$ is too large, the flow resistance is too small resulting only in a minor improvement of the efficiency of the deagglomerator arrangement 16.

Comprehensive studies of the geometry of the deagglomerator arrangement 16 in terms of the flow resistance of the device and the quality of the cyclonic flow effected inside the vortex chamber 73 show that a very good efficiency of the device can be obtained if the diameter $d$ is between 6 mm and 10 mm, preferably $6 \text{ mm} \leq d \leq 8 \text{ mm}$.

For example, if $d$ is 6 mm, a flow rate of 30 l/min at a pressure of 4 kPa can be obtained. The cyclonic flow inside the vortex chamber 73 is of good quality. Nevertheless, the required increase in the width of the air inlets or in the outlet diameter to allow a higher target flow rate (for example 60 l/min at 4 kPa) would presumably result in a degradation of the quality of the cyclone. This is because the dimensions of the air inlets or of the outlet would be too large in comparison to the dimensions of the main section of the cyclone, that is the dimensions of the vortex chamber 73.

Therefore, it may be recommendable to use a larger diameter $d$ of the "base" circle 77. A very compact design of the cyclone 16 and, in addition, the above-mentioned target flow rate in the
range of 60 l/min at 4 kPa can be obtained if the diameter d of the “base” circle 77 is in the range of 8 mm. Larger diameters d of the “base” circle 77 do not lead to further significant improvements of the flow pattern inside the vortex chamber 73 over that which can be achieved with a diameter d = 8 mm. Therefore, the diameter d = 8 mm is considered as a preferred embodiment of the cyclone 16.

As depicted in Figure 27, the deagglomerator arrangement or cyclone 16 according to this embodiment has two air inlets assigned with the respective air inlet conduits 75. Both air inlets disemboguing into the vortex chamber 73 are arranged diametrically opposed to one another and preferably have a width or a diameter of d/5. As shown in Figures 26 and 27, the side walls 78 separating the air inlet conduits 75 from the vortex chamber 73 end in a fillet which may have an end radius of 0.3 mm, for example. The side walls 78 may have a thickness of 0.6 mm, for example.

The cross-sectional structure of the deagglomerator arrangement 16 is such that a respective semicircle or arc 79, whose centre is on the diameter connecting both air inlets or both end portions of the side walls 78, connects the outer wall of one air inlet 75 with the inner surface of the opposing side wall 78 of the vortex chamber 73 at a place where this side wall 78 intersects with the respective other air inlet. As can be seen from Figure 27, this applies to both outer peripheral portions of the vortex chamber 73 and to both air inlets/air inlet conduits 75, respectively. The side walls of the air inlet conduits 75 are concentric to the side walls 78 of the vortex chamber 73. However, the semicircular or arched (curved) wall portions 79 of the vortex chamber 73 are non-concentric relative to the interior of the vortex chamber 73, i.e. the “base” circle 77. This contributes to a very effective deagglomeration within the vortex chamber 73.
The structure shown in Figure 27 of the deagglomerator arrangement or cyclone 16 may be extruded over a height of 7.7 mm, for example. The inhalation channel 27, that is the outlet cylinder extending from the circular outlet 74, preferably has a circular diameter of 0.75d and may be extruded on top of the vortex chamber 73 over a height of 37 mm, for example. The above-mentioned top inlet window 76, covering 80° of the air inlet conduits 75, may be extruded over 2 mm to provide the channels through which the air may enter into the device from the top of these windows.

Figures 28A, 28B and 28C show a perspective view, a bottom view and a top view, respectively, of another embodiment of a dosing sub-assembly 13 of a powder inhaler according to the invention. As can be seen, the container 7 comprises a medicament chamber 8 having an elliptical cross-section. Inside the medicament chamber 8, side walls 80 tapering or being slanted downward are provided, thereby facilitating the filling of the dosing recess of the slide, when it is in its filling position, by gravity. Again, the desiccant chamber 9 is separated from the medicament chamber 8 by a permeable membrane 10.

The dosing sub-assembly 13 of this embodiment comprises a deagglomerator arrangement (cyclone) 16 similar to that described above. From Figure 28C, the wall portions 79 being non-concentric to the interior diameter of the vortex chamber 73 are evident. Furthermore, Figure 28C also shows the tangential air inlet conduits 75.

As regards the dosing sub-assembly of Figure 28, generally reference can be made to the above description of the foregoing drawings.
Claims

1. Powder inhaler, comprising:
   - a container (7) for storing a powdered medicament,
   - a metering member (15) having a dosing recess (18), the metering member (15) being moveable between a filling position, in which the dosing recess (18) is in alignment with an opening of the container (7) so as to be filled with a dose of the powdered medicament, and an inhalation position, in which the dosing recess (18) is in alignment with an inhalation channel (27), and
   - a mouthpiece (3) being in communication with the inhalation channel (27) for enabling inhalation of the dose of the powdered medicament contained in the dosing recess (18) of the metering member (15) when the metering member (15) is in the inhalation position, characterized in that
     - a protective member (19) is provided between the metering member (15) and the inhalation channel (27), the protective member (19) being moveable between a closed position, in which the protective member (19) at least covers the dosing recess (18) of the metering member (15) when the metering member (15) is in the inhalation position, thereby preventing the powdered medicament contained in the dosing recess (18) from entering into the inhalation channel (27) and an open position, in which the protective member (19) does not cover the dosing recess (18), thereby exposing the dosing recess (18) to the inhalation channel (27) so as to enable inhalation of the dose of the powdered medicament contained in the dosing recess (18).

2. Powder inhaler according to claim 1, characterized in that
the powder inhaler comprises a casing (1) and a cover (2) being rotatably coupled to the casing (1) so that the cover (2) is moveable between a closed position, in which it covers the mouthpiece (3), and an open position, in which it exposes the mouthpiece (3).

3. Powder inhaler according to claim 2, characterized in that
the casing (1) comprises a window (4) for displaying a number of doses of the powdered medicament left in the container (7) or having been inhaled, the number of doses of the powdered medicament being counted by a dose counting unit (24–26).

4. Powder inhaler according to claim 2 or claim 3, characterized in that
the casing (1) comprises an opening (5) for displaying a mark (38) showing if the dose of the powdered medicament contained in the dosing recess (18) of the metering member (15) is ready for inhalation, or has already been inhaled.

5. Powder inhaler according to any one of the preceding claims, characterized in that
the container (7) comprises a medicament chamber (8) for storing the powdered medicament and an integral desiccant chamber (9) for storing a desiccant, the desiccant chamber (9) being separated from the medicament chamber (8) by a permeable membrane (10).

6. Powder inhaler according to claim 5, characterized in that
the permeable membrane (10) has a permeability which is different from that of a wall of the container (7) between
either the desiccant chamber (9) or the medicament chamber (8) and the outside of the powder inhaler.

7. Powder inhaler according to claim 6, characterized in that the material of the permeable membrane (10) is different from that of a wall of the container (7) between either the desiccant chamber (9) or the medicament chamber (8) and the outside of the powder inhaler.

8. Powder inhaler according to claim 6 or claim 7, characterized in that the thickness of the permeable membrane (10) is smaller than a wall of the container (7) between either the desiccant chamber (9) or the medicament chamber (8) and the outside of the powder inhaler.

9. Powder inhaler according to any one of claims 5-8, characterized in that both the medicament chamber (8) and the desiccant chamber (9) are sealed by a sealing means (11, 12).

10. Powder inhaler according to any one of claims 5-9, characterized in that the medicament chamber (8) has in its bottom a dosing opening through which the dose of the powdered medicament is filled into the dosing recess (18) of the metering member (15) by gravity if the metering member (15) is in the filling position.

11. Powder inhaler according to any one of claims 5-10, characterized in that
the medicament chamber (8) has a cross-section diameter which gradually decreases from the top of the medicament chamber to the bottom of the medicament chamber.

12. Powder inhaler according to any one of the preceding claims, characterized in that 
the metering member (15) is a slide being slidingly moveable between the filling position, in which the dosing recess (18) faces a dosing opening of the container (7), and the inhalation position, in which the dosing recess (18) faces an inhalation opening of the inhalation channel (27).

13. Powder inhaler according to claim 12, characterized in that 
the slide (15) is slidingly moveable in a horizontal direction such that the dosing recess (18) is located under the dosing opening of the container (7) if the slide (15) is in the filling position, while the dosing recess (18) is located under the inhalation opening of the inhalation channel (27) if the slide (15) is in the inhalation position.

14. Powder inhaler according to any one of claims 2-4 and any one of the preceding claims, characterized in that 
the metering member (15) is coupled to the cover (2) such that opening the cover (2) causes the metering member (15) to move from the filling position to the inhalation position, while closing the cover (2) causes the metering member (15) to move from the inhalation position to the filling position.

15. Powder inhaler according to claim 14, characterized in that
the coupling between the metering member (15) and the cover (2) comprises projections (28) engaging with recesses (31).

16. Powder inhaler according to claim 15, characterized in that the recesses (31) are formed at side surfaces of the cover (2), while the projections (28) are formed at sides of the metering member (15).

17. Powder inhaler according to claim 15 or claim 16, characterized in that the recesses (31) are shaped like profiled cam tracks.

18. Powder inhaler according to any one of claims 14-17, characterized in that the coupling between the metering member (15) and the cover (2) is such that opening of the cover (2) within a predetermined range of rotational movement of the cover (2) from the closed position of the cover (2) does not cause the metering member (15) to move from the filling position to the inhalation position.

19. Powder inhaler according to claim 18, characterized in that the predetermined range of rotational movement of the cover (2) corresponds to an angle of rotation up to about 30° from the closed position of the cover (2).

20. Powder inhaler according to any one of claims 14-19, characterized in that the coupling between the metering member (15) and the cover (2) is such that the metering member (15) reaches the inhalation position already a predetermined rotational movement of the cover (2) prior to the cover (2) being fully open.
21. Powder inhaler according to claim 20, characterized in that
the predetermined rotational movement of the cover (2), at
which the metering member (15) reaches the inhalation posi-
tion, corresponds to an angle of rotation of the cover (2)
of about 90°-135° from the closed position of the cover
(2).

22. Powder inhaler according to any one of the preceding claims, characterized in that
the dosing recess (18) is a dosing cup having a circular profile.

23. Powder inhaler according to claim 22, characterized in that
the dosing cup (18) has a semi-elliptical cross-section.

24. Powder inhaler according to claim 22 or claim 23, characterized in that
the diameter of the dosing cup (18) is about three times the depth of the dosing cup (18).

25. Powder inhaler according to any one of the preceding claims, characterized in that
the powder inhaler comprises a waste bin, and the metering member (15) comprises an opening so as to enable excess or residuary powdered medicament in the inhalation channel (27) or on the metering member (15) to fall through the opening into the waste bin.

26. Powder inhaler according to any one of the preceding claims, characterized in that
the powder inhaler comprises an inhalation actuated mechanism (21-23) being coupled to the protective member (19) such that, if the protective member (19) is in the closed position, the inhalation actuated mechanism (21, 23) causes the protective member (19) to move into the open position if an inhalation suction force being effected by a user exceeds a predetermined value.

27. Powder inhaler according to claim 26, characterized in that
the inhalation actuated mechanism comprises an inhalation actuated member (21) being moveable between a first position and a second position, the inhalation actuated member (21) being coupled to the protective member (19) such that, if there is an inhalation suction force exceeding the predetermined value, the inhalation actuated member (21) is moved from the first position to the second position, thereby causing the protective member (19) to move from the closed position to the open position.

28. Powder inhaler according to claim 27, characterized in that
the inhalation actuated member (21) is a flap being pivotable between the first position and the second position.

29. Powder inhaler according to claim 28, characterized in that
the first position is a horizontal position of the flap (21) while the second position is a position pivoted relative to the horizontal position by a rotational movement of the flap around an axis of rotation.

30. Powder inhaler according to any one of claims 27-29, characterized in that
the inhalation actuated mechanism (21-23) comprises a resilient means (23) being tensioned by a movement of the metering member (15) from the filling position to the inhalation position and being allowed to discharge upon a movement of the metering member (15) from the inhalation position to the filling position, the resilient means (23) being arranged such that the resilient means (23) holds the inhalation actuated member (21) in its first position if the resilient means (23) is discharged, while the resilient means (23) releases the inhalation actuated member (21) if the resilient means is tensioned, so as to allow the inhalation actuated member (21) to be moved from its first position to its second position by an inhalation suction force exceeding the predetermined value.

31. Powder inhaler according to claim 30, characterized in that
the inhalation actuated mechanism (21-23) comprises a coupling member (22) coupling the inhalation actuated member (21) to the protective member (19), the resilient means (23) being arranged on the coupling member (22).

32. Powder inhaler according to claim 31, characterized in that
the resilient means (23) has a prolonged end (34) which comes into contact with the metering member (15) upon movement of the metering member (15) from the filling position to the inhalation position, thereby tensioning the resilient means (23).

33. Powder inhaler according to claim 32, characterized in that
the metering member (15) comprises at least one recess (30) formed at a front end of the metering member (15), the pro-
longed end (34) of the resilient means (23) coming into contact with the recess (30) of the metering member (15) upon movement of the metering member (15) from the filling position to the inhalation position.

34. Powder inhaler according to any one of claims 31-33, characterized in that the resilient means (23) has an end portion (33) which holds the inhalation actuated member (21) in its first position when the resilient means (23) is discharged and releases the inhalation actuated member (21) when the resilient means (23) is tensioned by the metering member (15).

35. Powder inhaler according to any one of claims 31-34, characterized in that the coupling member (22) is a yoke.

36. Powder inhaler according to any one of claims 31-35, characterized in that the coupling member (22) has an arm (43) being held by the inhalation actuated member (21), if the inhalation actuated member (21) is in its first position, and being released by a movement of the inhalation actuated member (21) from its first position to its second position.

37. Powder inhaler according to any one of claims 31-36, characterized in that the resilient means (23) has an end portion (32) resting at the coupling member (22) such that upon tensioning of the resilient means (23) the coupling member (22) is biased to move from an initial position, in which the protective member (19) is in the closed position, to an end position, in which the protective member (19) is caused to move to the open position.
38. Powder inhaler according to claim 36 and 37, characterized in that the inhalation actuated mechanism (21-23) is arranged such that a movement of the coupling member (22) from the initial position to the end position is only enabled if the arm (43) of the coupling member (22) is released by a movement of the inhalation actuated member (21) from its first position to its second position.

39. Powder inhaler according to any one of claims 31-38, characterized in that the coupling member (22) comprises a prolongation (47) engaging with an opening (39) formed in the protective member (19).

40. Powder inhaler according to claim 39 and claim 37 or claim 38, characterized in that the prolongation (47) of the coupling member (22) in addition is moveably arranged in a longitudinal opening (29), which is formed in the metering member (15) along its longitudinal direction, such that the prolongation (47) of the coupling member (22) can freely move in the longitudinal opening (29) of the metering member (15) from its initial position to its end position, while a movement of the metering member (15) from the inhalation position to the filling position causes the prolongation (47) of the coupling member (22) to abut against an edge of the longitudinal opening (29) thereby moving the coupling member (22) back into its initial position.

41. Powder inhaler according to any one of claims 27-40, characterized in that
the inhalation actuated member (21) comprises a mark (38) which is visible through an opening (5) of a casing (1) of the powder inhaler if the inhalation actuated member (21) is in its first position, while the mark (38) is not visible through the opening (5) if the inhalation actuated member (21) is in its second position, the mark (38) thereby indicating whether the dose of the powdered medicament contained in the dosing recess (18) of the metering member (15) is ready for inhalation.

42. Powder inhaler according to any one of claims 27-41, characterized in that the inhalation actuated mechanism (21-23) is arranged such that the inhalation actuated mechanism (21-23) blocks a movement of the protective member (19) from the closed position to the open position if the user blows into the mouthpiece (3).

43. Powder inhaler according to any one of the preceding claims, characterized in that the powder inhaler comprises a dose counting mechanism (24-26) counting the number of doses of the powdered medicament remaining in the container (7) or having been inhaled, the dose counting mechanism (24-26) being arranged such that it is activated after completion of an inhalation process.

44. Powder inhaler according to claim 43, characterized in that the dose counting mechanism (24-26) comprises a wheel (24) being numbered on one side facing a window (4) of a casing (1) of the powder inhaler and being stepwise rotated with each inhalation process.

45. Powder inhaler according to claim 44,
characterized in that
the dose counting mechanism (24–26) comprises a further
wheel (26) being coupled to the wheel by a gear arrangement
(25), the further wheel being also numbered on one side
facing the window (4) of the casing (1).

46. Powder inhaler according to claim 45,
characterized in that
the wheels (24, 26) are provided for displaying different
orders of magnitude of the number of doses, the wheels (24,
26) being arranged such that in each case the numbers of
the wheels (24, 26) being located adjacent to one another
are visible through the window (4) of the casing (1).

47. Powder inhaler according to claim 45 or claim 46,
characterized in that
the further wheel (26) comprises two rows of numbers (68)
along its circumferential direction, both rows of numbers
(68) being provided for displaying a different order of
magnitude of the number of doses.

48. Powder inhaler according to any one of claims 45–47,
characterized in that
the further wheel has a projection (69) extending from its
outer periphery, the projection (69) being arranged such
that it covers a portion of the wheel (24) at a position
facing the window (4) of the casing (1) in such a manner
that no numbers of the wheel (24) and no numbers of the
further wheel (26) are visible through the window (4), the
further wheel (26) being not provided with numbers (68) in
a radial direction of the projection (69).

49. Powder inhaler according to any one of claims 44–48,
characterized in that
the dose counting mechanism (24-26) is arranged such that a further activation of the dose counting mechanism (24-26) is blocked if the dose counting mechanism (24-26) has counted that all doses of the powdered medicament contained in the container (7) have already been taken.

50. Powder inhaler according to any one of claims 44-49, characterized in that the dose counting mechanism (24-26) is coupled to the metering member (15) such that it is activated by a movement of the metering member (15) from the inhalation position to the filling position.

51. Powder inhaler according to any one of claims 44-50 and any one of claims 31-40, characterized in that the wheel (24) is provided with a plurality of drive teeth (58) along a circumferential direction thereof, and the coupling member (22) of the inhalation activated mechanism (21-23) comprises a projection (45) which, with each inhalation process, is brought into engagement with one of the drive teeth (58) so as to rotate the wheel (24) by one step.

52. Powder inhaler according to claim 51 and any one of claims 37, 38 or 40, characterized in that the projection (45) is moved to a next drive tooth (58) of the wheel (24) and brought into engagement therewith upon movement of the coupling member (22) from its initial position to its end position, the projection (45) of the coupling member (22) rotating the wheel (24) by one step upon movement of the coupling member (22) from its end position back to its initial position.
53. Powder inhaler according to claim 51 or claim 52, characterized in that each of the drive teeth (58) has an inclination in the circumferential direction of the wheel (24) so as to facilitate movement of the projection (45) of the coupling member (22) over the drive teeth (58).

54. Powder inhaler according to any one of the preceding claims, characterized in that the protective member (19) is arranged on the metering member slidingly moveable between the closed position and the open position.

55. Powder inhaler according to any one of the preceding claims, characterized in that a one-way valve is placed in the inhalation channel (27).

56. Powder inhaler according to any one of the preceding claims, characterized in that a manual override mechanism (38) is provided for manually actuating the protective member (19) so as to move the protective member (19) from the closed position to the open position.

57. Powder inhaler according to claim 56 and any one of claims 26-42 or 51-53, characterized in that the manual override mechanism (38) is provided for manually actuating the inhalation actuated mechanism (21-23).
58. Powder inhaler according to any one of claims 56, 57 and claim 41, characterized in that the manual override mechanism comprises the mark (38) for manually pushing down the inhalation actuated member (21) so as to manually actuate the inhalation actuated mechanism (21-23) and the protective member (19).

59. Deagglomerator arrangement (16) for deagglomerating a powdered medicament, comprising:
   a vortex chamber (73) having an opening for the supply of the powdered medicament,
   at least one air inlet (75) for directing air tangentially into the vortex chamber (73), and
   an outlet (74) for outputting air with the deagglomerated powdered medicament, the outlet (74) being spaced from the at least one air inlet (75) in an axial direction of the deagglomerator arrangement (16), characterized in that the vortex chamber (73) has a diameter \( d \) of \( 6 \text{ mm} \leq d \leq 10 \text{ mm} \).

60. Deagglomerator arrangement according to claim 59, characterized in that the diameter of the vortex chamber (73) is the largest possible diameter of a circle (77) which can be laid inside the vortex chamber (73) on a horizontal plane intersecting the at least one air inlet (75) without extending beyond the vortex chamber (73).

61. Deagglomerator arrangement according to claim 60, characterized in that the vortex chamber (73) has in the horizontal plane a substantially elliptical cross-section.
62. Deagglomerator arrangement according to any one of claims 59-61,
characterized in that
the vortex chamber (73) is designed rotationally symmetrical relative to its longitudinal central axis.

63. Deagglomerator arrangement according to any one of claims 59-62,
characterized in that
the vortex chamber (73) has a diameter d of about 8 mm.

64. Deagglomerator arrangement according to any one of claims 59-63,
characterized in that
two diametrically opposed air inlets (75) are provided.

65. Deagglomerator arrangement according to any one of claims 59-64,
characterized in that
the at least one air inlet (75) has a width of d/5.

66. Deagglomerator arrangement according to any one of claims 59-65,
characterized in that
the outlet (74) is associated with a respective outlet channel (27) having a circular cross-section.

67. Deagglomerator arrangement according to claim 66,
characterized in that
the outlet channel (27) extends along and coaxially to the longitudinal central axis of the vortex chamber (73).

68. Deagglomerator arrangement according to claim 66 or 67,
characterized in that
the outlet channel (27) has a diameter of 0.75d.

69. Deagglomerator arrangement according to any one of claims 59-68,
characterized in that
the at least one air inlet (75) is associated with a re-
pective air inlet conduit, the vortex chamber (73) being
separated from the at least one air inlet conduit by a side
wall (78) being concentric to the air inlet conduit.

70. Deagglomerator arrangement according to claim 69,
characterized in that
the side wall (78) has a rounded end portion at a place ad-
acent to the air inlet (75).

71. Deagglomerator arrangement according to any one of claims
59-70 and claim 64,
characterized in that
an outer wall of each air inlet (75) is connected to the
respective other air inlet by a substantially semicircular
wall portion (79) of the vortex chamber (73), the centre of
the respective semicircle of the wall portion (79) being
positioned on the diameter of the vortex chamber (73) con-
necting both air inlets (75) with one another.

72. Deagglomerator arrangement according to claim 71,
characterized in that
each substantially semicircular wall portion (79) is posi-
tioned non-concentric to a horizontal circle (77) defining
the diameter of the vortex chamber (73).

73. Deagglomerator arrangement according to any one of claims
59-72,
characterized in that
the at least one air inlet (75) is associated with a respective air inlet conduit, a top surface of the air inlet conduit being exposed so as to allow entry of air into the air inlet conduit.

74. Deagglomerator arrangement according to claim 73, characterized in that
the top surface of the air inlet conduit is exposed over an angle of about 80° along a circumferential direction of the air inlet conduit.

75. Deagglomerator arrangement (16) for deagglomerating a powdered medicament, comprising:
a vortex chamber (73) having an opening for the supply of the powdered medicament,
at least two air inlets (75) for directing air tangentially into the vortex chamber (73), and
an outlet (74) for outputting air with the deagglomerated powdered medicament, the outlet (74) being spaced from the air inlets (75) in an axial direction of the deagglomerator arrangement (16), characterized in that
an outer wall of each air inlet (75) is connected to the other air inlet (75) by an arched wall portion (79) of the vortex chamber (73), each arched wall portion (79) being positioned non-concentric to a horizontal circle (77) defining a diameter (d) of the vortex chamber (73).

76. Deagglomerator arrangement according to claim 75, characterized in that
the air inlets (75) are arranged along the circle (77) defining the diameter (d) of the vortex chamber (73).
77. Deagglomerator arrangement according to claim 75 or claim 76,
characterized in that
the deagglomerator arrangement (16) is a deagglomerator ar-
rangement according to any one of claims 59-74.

78. Powder inhaler according to any one of claims 1-58,
characterized in that
the inhalation channel (27) comprises a deagglomerator ar-
rangement (16) according to any one of claims 59-77.
INTERNATIONAL SEARCH REPORT

A. CLASSIFICATION OF SUBJECT MATTER

IPC 7 A61M15/00

According to International Patent Classification (IPC) or to both national classification and IPC

B. FIELDS SEARCHED

Minimum documentation searched (classification system followed by classification symbols)

IPC 7 A61M

Documentation searched other than minimum documentation to the extent that such documents are included in the fields searched

Electronic data base consulted during the international search (name of data base and, where practical, search terms used)

EPO-Internal, PAJ, WPI Data

C. DOCUMENTS CONSIDERED TO BE RELEVANT

<table>
<thead>
<tr>
<th>Category</th>
<th>Citation of document, with indication, where appropriate, of the relevant passages</th>
<th>Relevant to claim No.</th>
</tr>
</thead>
<tbody>
<tr>
<td>A</td>
<td>FR 2 701 653 A (VALOIS) 26 August 1994 (1994-08-26) page 1, line 1 - line 5 page 5, line 11 - page 6, line 14; figure 1</td>
<td>1</td>
</tr>
<tr>
<td>A</td>
<td>FR 2 667 509 A (VALOIS) 10 April 1992 (1992-04-10) page 21, line 3 - page 27, line 30; figures 9-12</td>
<td>1</td>
</tr>
<tr>
<td>X</td>
<td>WO 01 00262 A (CAMBRIDGE CONSULTANTS; EASON STEPHEN WILLIAM (GB); HARMER QUENTIN) 4 January 2001 (2001-01-04)</td>
<td>59, 60, 62, 63, 66, 67, 69, 70, 73</td>
</tr>
<tr>
<td>Y</td>
<td>page 7, line 1 - line 22; claims 1-4; figure 1</td>
<td>61, 64, 65, 71</td>
</tr>
<tr>
<td>A</td>
<td>----</td>
<td>75</td>
</tr>
</tbody>
</table>

Further documents are listed in the continuation of box C. Patent family members are listed in annex.

* Special categories of cited documents:

A* document defining the general state of the art which is not considered to be of particular relevance
E* earlier document but published on or after the international filing date
L* document which may throw doubts on priority claim(s) or which is cited to establish the publication date of another document, or for other reasons as specified
O* document relating to an oral disclosure, use, exhibition or other means
P* document published prior to the international filing date but later than the priority date claimed

*Y* Inter document published after the international filing date or priority date and not in conflict with the application but cited to understand the principles or theory underlying the invention
*X* document of particular relevance; the claimed invention cannot be considered novel or cannot be considered to involve an inventive step when the document is taken alone
*V* document of particular relevance; the claimed invention cannot be considered to involve an inventive step when the document is combined with one or more other such documents, such combination being obvious to a person skilled in the art.

Date of the actual completion of the international search: 31 October 2003

Date of mailing of the international search report: 10/11/2003

Name and mailing address of the ISA:
European Patent Office, P.B. 5818 Patentlaan 2 NL-2280 HV Rijswijk Tel. (+31-70) 340-2000, Fax: (+31-70) 340-3016

Authorized officer: Zeinstra, H
## INTERNATIONAL SEARCH REPORT

**C (Continuation) DOCUMENTS CONSIDERED TO BE RELEVANT**

<table>
<thead>
<tr>
<th>Category</th>
<th>Citation of document, with indication, where appropriate, of the relevant passages</th>
<th>Relevant to claim No.</th>
</tr>
</thead>
<tbody>
<tr>
<td>X</td>
<td>WO 02 07805 A (CAMBRIDGE CONSULTANTS; EASON STEPHEN WILLIAM (GB); HARMER QUENTIN) 31 January 2002 (2002-01-31) page 7, line 6 - line 16 page 10, line 12 - line 31; figures 1,2</td>
<td>59,60, 62,63, 66-70,73</td>
</tr>
<tr>
<td>X</td>
<td>US 5 476 093 A (LANKINEN TAPIO) 19 December 1995 (1995-12-19) column 5, line 7 - line 39 column 6, line 41 - column 8, line 2; figures</td>
<td>59,60, 62,63, 66,67, 69,70,73</td>
</tr>
<tr>
<td>Y</td>
<td>US 6 073 629 A (MAO LEI ET AL) 13 June 2000 (2000-06-13) column 5, line 8 - column 6, line 37; figures 1,4; table 2</td>
<td>61,65</td>
</tr>
<tr>
<td>Y</td>
<td>US 6 418 926 B1 (CHAWLA BRINDRA PAUL SINGH) 16 July 2002 (2002-07-16) column 8, line 6 - column 9, line 26; figures 6-10</td>
<td>64,71, 75</td>
</tr>
</tbody>
</table>
# INTERNATIONAL SEARCH REPORT

## Box I  Observations where certain claims were found unsearchable (Continuation of item 1 of first sheet)

This International Search Report has not been established in respect of certain claims under Article 17(2)(a) for the following reasons:

1. [ ] Claims Nos., because they relate to subject matter not required to be searched by this Authority, namely:

2. [ ] Claims Nos., because they relate to parts of the International Application that do not comply with the prescribed requirements to such an extent that no meaningful International Search can be carried out, specifically:

3. [ ] Claims Nos., because they are dependent claims and are not drafted in accordance with the second and third sentences of Rule 6.4(a).

## Box II  Observations where unity of invention is lacking (Continuation of item 2 of first sheet)

This International Searching Authority found multiple inventions in this International application, as follows:

```
see additional sheet
```

1. [ ] As all required additional search fees were timely paid by the applicant, this International Search Report covers all searchable claims.

2. [x] As all searchable claims could be searched without effort justifying an additional fee, this Authority did not invite payment of any additional fee.

3. [ ] As only some of the required additional search fees were timely paid by the applicant, this International Search Report covers only those claims for which fees were paid, specifically claims Nos.:

4. [ ] No required additional search fees were timely paid by the applicant. Consequently, this International Search Report is restricted to the invention first mentioned in the claims; it is covered by claims Nos.:

**Remark on Protest**

[ ] The additional search fees were accompanied by the applicant's protest.

[ ] No protest accompanied the payment of additional search fees.
This International Searching Authority found multiple (groups of) inventions in this international application, as follows:

1. Claims: 1-58, 78

   Powder inhaler
   (Problem: to provide an inhaler with an improved dosing ability whereby unintended dosing can be avoided (see the description at page 8, lines 17 – 19)).

2. Claims: 59-77

   Deagglomerator arrangement
   (Problem: to provide an optimal pulverization of agglomerate of a medicament to be inhaled (see the description at page 8, lines 20 – 21)).
<table>
<thead>
<tr>
<th>Patent document cited in search report</th>
<th>Publication date</th>
<th>Patent family member(s)</th>
<th>Publication date</th>
</tr>
</thead>
<tbody>
<tr>
<td></td>
<td></td>
<td>WO 9205823 A1</td>
<td>16-04-1992</td>
</tr>
<tr>
<td></td>
<td></td>
<td>WO 9205824 A1</td>
<td>16-04-1992</td>
</tr>
<tr>
<td></td>
<td></td>
<td>FR 2667790 A1</td>
<td>17-04-1992</td>
</tr>
<tr>
<td></td>
<td></td>
<td>US 5447151 A</td>
<td>05-09-1995</td>
</tr>
<tr>
<td>WO 0100262</td>
<td>04-01-2001</td>
<td>GB 2353222 A</td>
<td>21-02-2001</td>
</tr>
<tr>
<td></td>
<td></td>
<td>AU 5534600 A</td>
<td>31-01-2001</td>
</tr>
<tr>
<td></td>
<td></td>
<td>CA 2376868 A</td>
<td>04-01-2001</td>
</tr>
<tr>
<td></td>
<td></td>
<td>WO 0100262 A1</td>
<td>04-01-2001</td>
</tr>
<tr>
<td></td>
<td></td>
<td>EP 1191966 A1</td>
<td>03-04-2002</td>
</tr>
<tr>
<td></td>
<td></td>
<td>JP 2003503116 T</td>
<td>28-01-2003</td>
</tr>
<tr>
<td>WO 0207805</td>
<td>31-01-2002</td>
<td>GB 2364919 A</td>
<td>13-02-2002</td>
</tr>
<tr>
<td></td>
<td></td>
<td>AU 8971001 A</td>
<td>05-02-2002</td>
</tr>
<tr>
<td></td>
<td></td>
<td>WO 0207805 A2</td>
<td>31-01-2002</td>
</tr>
<tr>
<td></td>
<td></td>
<td>EP 1301231 A2</td>
<td>16-04-2003</td>
</tr>
<tr>
<td></td>
<td></td>
<td>US 2003172926 A1</td>
<td>18-09-2003</td>
</tr>
<tr>
<td>US 5476093</td>
<td>19-12-1995</td>
<td>none</td>
<td>none</td>
</tr>
<tr>
<td></td>
<td></td>
<td>WO 9915217 A1</td>
<td>01-04-1999</td>
</tr>
<tr>
<td></td>
<td></td>
<td>US 6394085 B1</td>
<td>28-05-2002</td>
</tr>
<tr>
<td></td>
<td></td>
<td>AU 734730 B2</td>
<td>21-06-2001</td>
</tr>
<tr>
<td></td>
<td></td>
<td>CA 2304792 A1</td>
<td>01-04-1999</td>
</tr>
<tr>
<td></td>
<td></td>
<td>JP 2001517499 T</td>
<td>09-10-2001</td>
</tr>
<tr>
<td></td>
<td></td>
<td>NZ 503874 A</td>
<td>25-10-2002</td>
</tr>
<tr>
<td></td>
<td></td>
<td>ZA 9808791 A</td>
<td>13-09-2000</td>
</tr>
<tr>
<td></td>
<td></td>
<td>AU 737921 B2</td>
<td>06-09-2001</td>
</tr>
<tr>
<td></td>
<td></td>
<td>AU 5329198 A</td>
<td>15-07-1998</td>
</tr>
<tr>
<td></td>
<td></td>
<td>BR 9713726 A</td>
<td>25-01-2000</td>
</tr>
<tr>
<td></td>
<td></td>
<td>CA 2322434 A1</td>
<td>25-06-1998</td>
</tr>
<tr>
<td></td>
<td></td>
<td>CN 1235136 A</td>
<td>26-04-2000</td>
</tr>
<tr>
<td></td>
<td></td>
<td>DE 69722257 D1</td>
<td>26-06-2003</td>
</tr>
<tr>
<td></td>
<td></td>
<td>EP 0969889 A2</td>
<td>12-01-2000</td>
</tr>
<tr>
<td></td>
<td></td>
<td>HU 002492 A2</td>
<td>28-11-2000</td>
</tr>
<tr>
<td></td>
<td></td>
<td>JP 2001506152 T</td>
<td>15-05-2001</td>
</tr>
<tr>
<td></td>
<td></td>
<td>NO 992941 A</td>
<td>18-08-1999</td>
</tr>
<tr>
<td></td>
<td></td>
<td>NZ 336686 A</td>
<td>01-02-2002</td>
</tr>
<tr>
<td></td>
<td></td>
<td>PL 334199 A1</td>
<td>14-02-2000</td>
</tr>
<tr>
<td></td>
<td></td>
<td>SK 81799 A3</td>
<td>11-07-2000</td>
</tr>
</tbody>
</table>