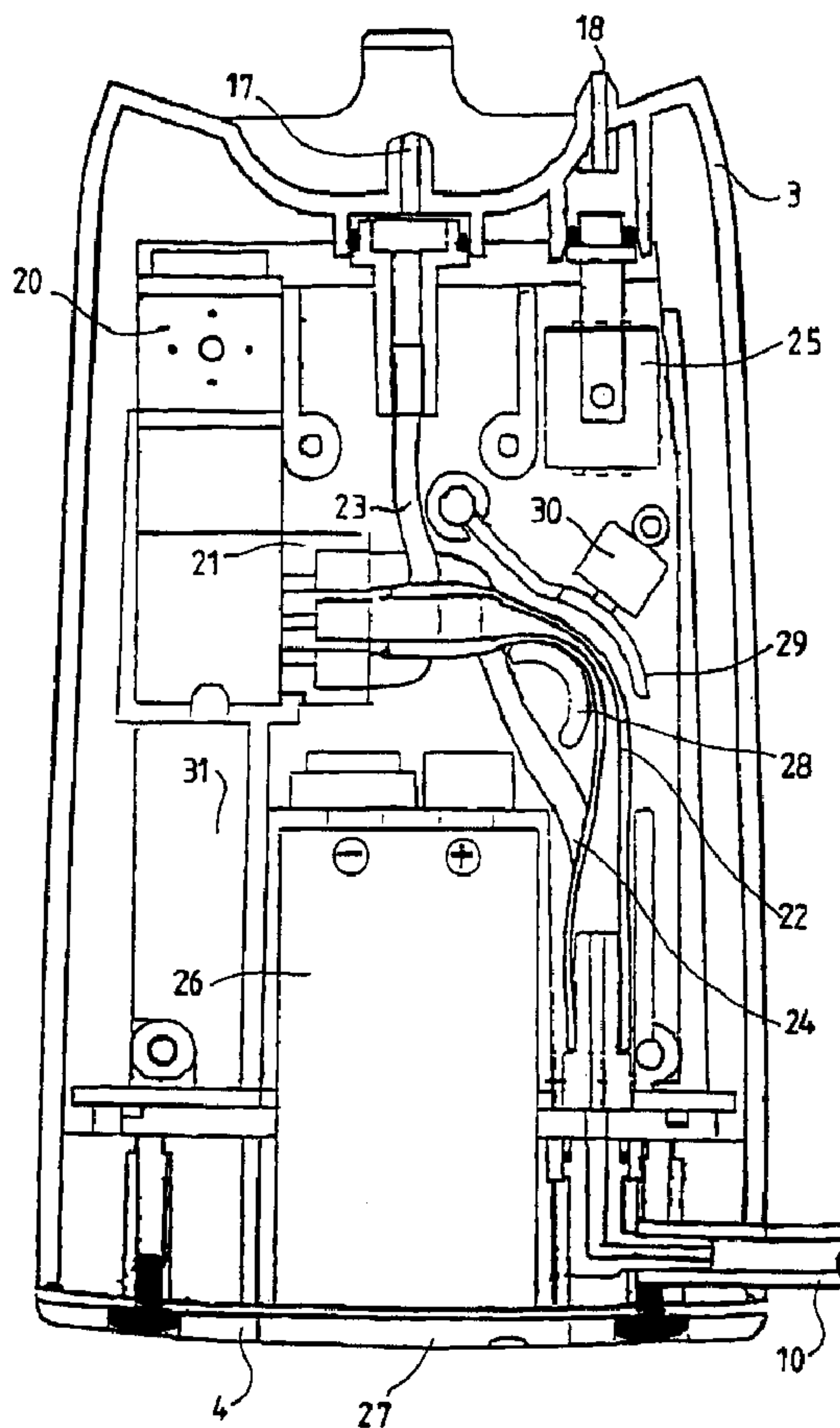




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 (54) Title: DISPENSING SYSTEM



(57) Abrégé/Abstract:

A product dispensing system comprises a calibrated nebulizer having a nebulizer jet, mouthpiece, means for sourcing compressed air, a manifold for distributing the source of compressed air in at least one direction, a nebulizer accommodation means which is

(57) **Abrégé(suite)/Abstract(continued):**

accessed by port, and a valve means for controlling the manifold and thereby the flow of compressed air to the port. The manifold and the port are linked by a length of tube, wherein the internal volume of the tube from the manifold outlet to the nebulizer jet is less than 0.7 ml.

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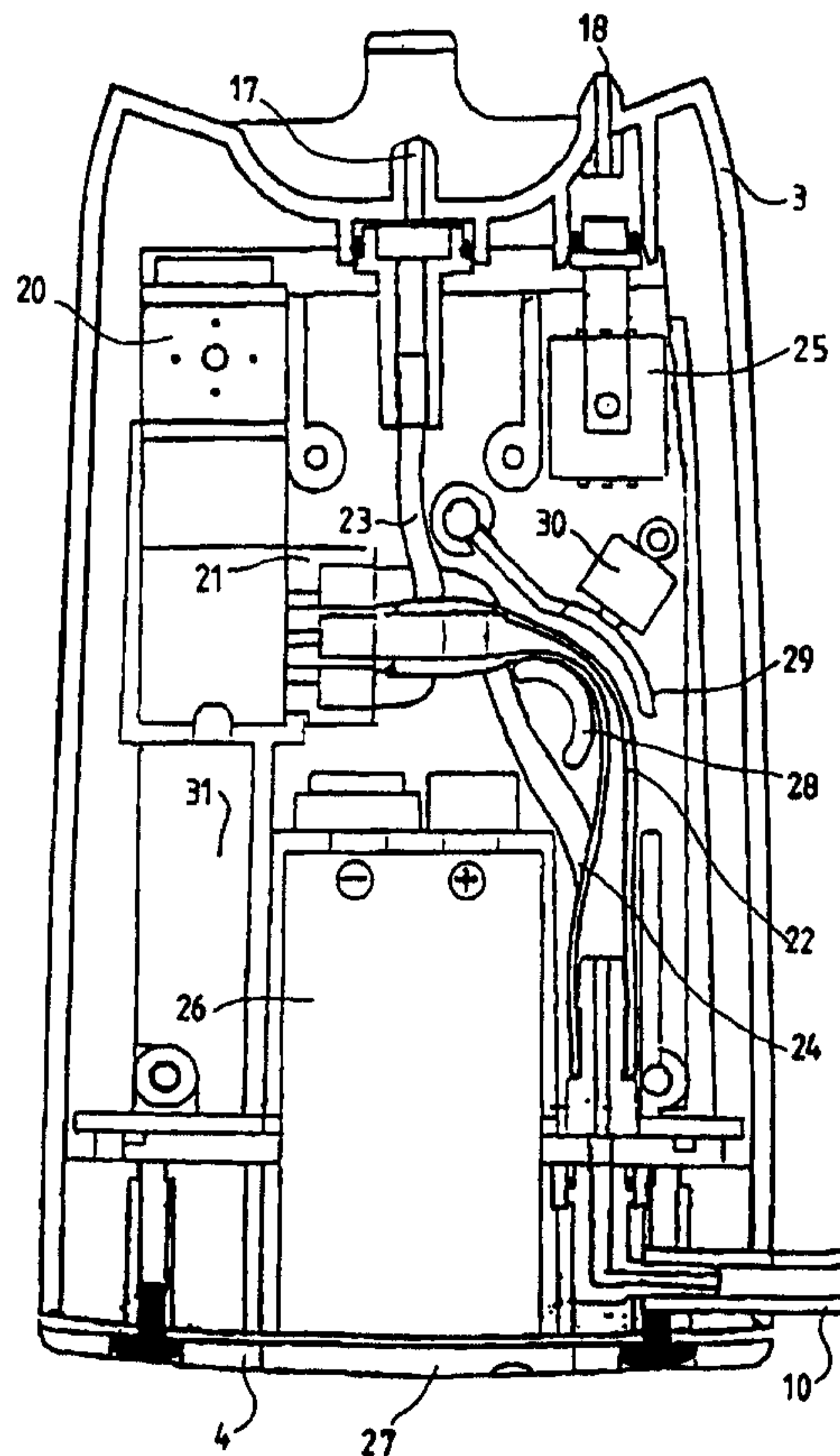
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(57) Abstract

A product dispensing system comprises a calibrated nebulizer having a nebulizer jet, mouthpiece, means for sourcing compressed air, a manifold for distributing the source of compressed air in at least one direction, a nebulizer accommodation means which is accessed by port, and a valve means for controlling the manifold and thereby the flow of compressed air to the port. The manifold and the port are linked by a length of tube, wherein the internal volume of the tube from the manifold outlet to the nebulizer jet is less than 0.7 ml.



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DISPENSING SYSTEM

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This invention relates to a dispensing system, in particular one which may be of use for example in the dispensing of small quantities of medicament, where the medicament is required to be dispensed in aerosol form.

15

It is known for dosing systems which supply a nebulized substance to containing a metering system. For example, in GB 1,568,808, (Rosenthal) there is described a metering system for supplying a nebulized substance for patient inhalation comprising a nebulizing means, a detecting means for detecting the initiation of the patient's inhalation, an adjustable timing means for adjusting a timed operation, and a valve means which is controlled by the timing means, in order to provide a controlled dosage of nebulized substance.

20

25

However, for this system to work and provide a precise dose of medicament, it is necessary that the system utilises a calibrated nebulizer which has a precise rate of output against time. Commercially available nebulizers have a wide range of outputs against time. In addition, the calibration must remain constant over the use of the nebulizer, and the nebulizer must be connected to the dosing device in such a way that the calibration of the nebulizer is valid and recognised when operated with the dispensing device. This is particularly important in relation to the length of tube between the valve and the nebulizer.

30

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5 Also, it is noted that this application describes no
teaching of a device which is in any way calibrated for
use with nominated drugs. This may have been due to the
fact that the apparatus embodied in this application is
10 designed for provocation testing, which is generally
carried out in a laboratory, and is used to deliver pulses
of aerosol into a patient's airway in order to determine
their reactivity to allergens. The apparatus is not
designed as one for a patient to take home, and to deliver
precise doses of medicament on a day to day basis.

15

 In addition to the above application, GB 2,164,569
(Etela-Hameen Kauhkovammayhdiatys RY (Finland)) describes
a similar system to that described above, which is also
designed for provocation testing, except that it has an
20 additional atomizing starting time control, which is used
for selecting the atomizing starting moment to coincide
with the beginning of the inspiration phase, which is
found by examination to be favourable for a particular
patient.

25

 EP 519,742 (DeVilbiss Healthcare Inc) describes a
medical nebulizer control system which has a three way
valve. In the embodiment, one part of the valve is
connected to a pressure sensor, and another is connected
30 to a compressed air supply. The control system within
this apparatus uses the supply tube to the nebulizer to
detect the patient's breathing pattern. When it detects
inhalation, it switches the valve over to the compressed
air supply, which then drives the nebulizer for a pre-
35 determined period of time, irrespective of whether the

patient has continued to inhale or not. This device also suffers from the problems of other nebulizers in that there is a long length of tube between the control box and the nebulizer. In addition the nebulizer can be
5 supplied from any manufacturer, and hence its calibration is not tied into the device, thereby causing variable amounts of medicament to be dispensed.

According to the present invention, a product dispensing
10 system comprises a calibrated nebulizer having a nebulizer jet, a mouthpiece, means for sourcing compressed air, a manifold for distributing the compressed air in at least one direction, a port, and a
15 valve means for controlling the manifold and thereby the flow of compressed air to the port, the manifold and the port being linked by a length of tube, characterised by a nebulizer accommodation means which is accessed by the
20 port, and in that the internal volume of the tube from the manifold outlet to the nebulizer jet is less than 0.7 ml.

Preferably, the internal volume of the tube from the manifold outlet to the nebulizer jet is less than 0.5 ml.

25 Conveniently, the dispensing system according to the invention is utilized in conjunction with a small air compressor as the source of compressed air, which has a flow rate in the region of 1-2 litres per minute

30 By "calibrated", in this instance, it is meant that the dosage rate for the nebulizer has already been established, and the dosimeter programmed accordingly, so as to provide a known dosage rate for a given medicament.

35

The dosage rate can be arranged to be fixed, regardless of the rate of inhalation of a patient, or variable depending on the rate of inhalation. What is important is that the dosage rate is known at a particular time.
5 It is envisaged that a given dosimeter will have one or more dedicated nebulizers.

In preferred embodiments of the invention, the dispensing system additionally comprises a switch means, which is
10 responsive as to whether compressed air is flowing in the system.

It is another preferred aspect of the invention that it may comprise a dispensing system comprising a dosimeter
15 which has been pre-programmed to dose one or more medicaments according to predetermined dosage profiles.

The dispensing system according to the invention is conveniently one which is capable of being pre-programmed
20 to dose one or more different types of drugs, which may need to be dosed according to different profiles. Methods of pre-programming the dispensing system in such a fashion will be familiar to those skilled in the art. In addition, the nebulizer and dosimeter according to the
25 invention are calibrated to determine the dosage of drug as described in EP 587,380 (Medic-Aid Limited).

The dispensing system according to the invention incorporates a nebulizer and a mouthpiece, by which
30 nebulized medicament can actually be delivered to the patient. The dispensing system can also additionally preferably comprise a pressure sensor, which is capable in

5 use of detecting a pressure drop in the apparatus in
response to the patient's breathing, and then delivering
a pulse of nebulized medicament into the mouthpiece. In
such instances, the dose of medicament can be calculated
10 by the known rate of output against time for the drug
selected, and the sum of all nebulizer pulses which the
system has delivered, in ways known to those skilled in
the art, as referred to above. Such ways may typically
include performing clinical trials on the drug to
determine appropriate dosages, and programming these
15 electronically into the dispensing system.

It is to be understood that the manifold in a
dispensing apparatus according to the invention can, in
fact, be an integral part of the valve means, and need not
20 be a separately discreet entity. In certain embodiments,
the manifold is a shaped block with internal galleys, for
example, made of plastics materials, and its presence
allows the tubes to be connected to the ports on the
valve. The ports on the valve may be too close together
25 to accommodate the connection of tubes directly, and a
manifold increases the space between them in other
embodiments it may be necessary for the manifold to
distribute compressed air in at least two different
directions.

30
The dispensing system according to the invention can
operate from a variety of different compressed air
sources, such as a continuous air supply at a rate of
around 6 litres per minute. This can typically be
35 provided either from a pressurized cylinder such as those

5 used in hospitals, or can be generated from a conventional
air compressor.

10 In an alternative and preferred configuration, the
dispensing system can be operated with a lightweight
compressor system, which typically generates a low flow
rate (in the order of 1-2 litres per minute, preferably
1.5 litres per minute), in conjunction with an
accumulator. Such a combination of low flow rate
15 compressor and accumulator allows the dispensing system to
produce pulses of compressed air to a nebulizer in the
dispensing system with the flow around 6 litres per
minute, where the flow from the compressor is matched to
the mean flow through the nebulizer (1.5 litres per
minute). In addition, such a combination of low flow rate
20 compressor with accumulator, in conjunction with the
dispensing system according to the invention, provides a
dosing apparatus which is more lightweight, compact and
readily portable than known dosing apparatuses.

25 With regard to the manifold, if the dispensing device
according to the invention is used in conjunction with a
low flow rate compressor, it is only necessary for the
valve to switch the compressed air on and off to the
nebulizer, in which case the manifold may only need to
30 direct the compressed air in one direction only. However,
if the dispensing system is used in conjunction with a
conventional air supply such as a six litre per minute
compressor or gas bottle supply, the manifold in
conjunction with the valve directs the flow to either the
35 nebulizer or the outlet orifice. In such circumstances,

5 the valve in conjunction with the manifold has either one
port or two outlet ports, depending on the application.

10 An important feature of the dispensing systems
according to the invention lies in the volume of the tube
linking the manifold and the port being less than 0.7 ml,
preferably less than 0.5 ml. It has been found that by
using lengths of tube which have a relatively small
volume, the nebulizer starts to work as quickly as
possible once the patient's inhalation has been detected.
15 Typically, it is possible that the nebulizer can work
within 50 milliseconds of detecting the patient's
inhalation. To facilitate both the rapid response time
and the low volume of the tube linking the manifold and
the port, it is preferred that the valve is physically
20 close to the nebulizer.

If a relatively long tube, or one with a large
internal volume is used between the manifold and the
nebulizer, this tube has to be pressurized before the
25 nebulizer starts to operate. This can have significant
effects on the performance of the system with regard to
the controlling the rate of output of the nebulizer, since
the nebulizer should preferably be provided with an
essentially constant rate ("square wave") of pressure over
30 time, so that its output is constant over time. The
nebulizer used in the dispensing system is preferably one
which delivers inspiratory patterns between 0.1 and 1.5
seconds duration.

The invention will now be described further by way of example only with reference to the accompanying drawings, in which:

5 Figure 1 shows a schematic view of a dispensing system according to the invention, complete with a nebulizer and mouthpiece fitted;

10 Figure 2 shows a schematic cross-section view of the nebulizer/mouthpiece end of the dispensing system of Figure 1;

15 Figure 3 shows a further schematic cross-section view of the nebulizer/mouthpiece end of the dispensing system of Figure 1 from an orthogonal position to that shown in Figure 2;

20 Figure 4 shows a schematic cross-section view of the control system end of the dispensing system of Figure 1;

 Figure 5 shows a schematic view of the dispensing system of Figure 1 attached to a low flow rate compressor;

25 Figure 6 shows a schematic representation of the combined dispensing system/compressor system of Figure 5;

30 Figures 7a-7d show a loading regime for loading an embodiment of dispensing system according to an aspect of the invention, wherein Figure 7a shows the dispensing system with a clean and empty nebulizer, Figure 7b shows removing the mouthpiece from the system; Figure 7c shows pouring the medication into the nebulizer; and Figure 7d shows placing the mouthpiece back onto the system to be ready for delivering the medication;

35

Figures 8a-8d show a cleaning regime for cleaning an embodiment of dispensing system according to an aspect of the invention wherein Figure 8a shows removing the mouthpiece; Figure 8b shows removing the funnel; Figure 8c shows the release of the nebulizer bowl assembly; and Figure 8d shows reassembling of the dispensing system after cleaning;

Figure 9 shows an accumulator consisting of a generally hemispherical elastic diaphragm; and

Figure 10 is a graph showing the nebulizer output rate of a typical nebulizer against the inspirator flow of a patient.

Referring to the figures, an embodiment of dispensing system according to the invention comprises a hand held dosimeter having a height of approximately 200 mm and a weight of approximately 200g. The dosimeter has a mouthpiece (1) and a nebulizer (2) attached, and a body (3) which contains a control valve, electronic circuitry and a battery. These components can be accessed through base (4).

The dosimeter is connected to a compressed air source via tube (10). To operate the dosimeter, the patient removes mouthpiece (1), and pours a liquid form of the medication (which may be a liquid or a powder in a fluidised form, or any other similar form) into the nebulizer (2). Then, the dosimeter is connected to the compressed air supply via tube (10). By means of a switch device to be described later, the presence of a positive pressure in the dosimeter activates the control circuitry in the dosimeter, and switches it on.

In this embodiment, although the nebulizer has only one well in which the medication sits, the dosimeter is nevertheless pre-programmed to deliver the correct dose of two different nominated drugs. The dosimeter could, however, be constructed and pre-programmed so as to deliver any desired number of nominated drugs. The drug which has been loaded into the nebulizer is selected using selector buttons (5) and (6). Once the drug type is selected, LED's (7) and (8) indicate the drug type selected by the patient. Button (9) is a re-set button, so that the user can correct any errors in selection.

The nebulizer used in the dispensing system according to the invention can be any suitable nebulizer design with a known calibration constant which is used to nebulize substances such as medicaments, which is suitably adapted to fit the dispensing system, and calibrated with the dosimeter. Suitable nebulizers include those which use a source of compressed air to nebulize the medicament, and are, for example, described in European Patent No. 672,266 (Medic-Aid Limited).

When the drug has been selected, the patient breaths in through a mouthpiece (1). A pressure sensor (to be further described later) within body (3) detects a pressure drop within the mouthpiece (1) due to the patient's inhalation, and then delivers a pre-programmed pulse of nebulized medicament into the first 50% of the inspiratory profile, until the dose regime programmed into the dispensing system has been delivered.

The dose is calculated from a known rate of output against time for the drug selected, and the sum of all the nebulizer pulses which the dosimeter has delivered. Further information on how the doses of drug may be derived and pre-programmed into the dosimeter may be obtained from GB 2,294,402 (Medic-Aid Limited et. al.). In this instance, clinical trials have been conducted to determine the dose of drug that must be delivered to achieve the correct therapeutic effect, and this dose has been programmed into the dosimeter for the two nominated drugs.

When the programmed dose has been delivered, the LED adjacent the button relating to the selected drug flashes rapidly and a buzzer sounds, indicating that the treatment is finished, and thereby ensuring that a precise dosage of drug is delivered to the patient on every occasion.

The nebulizer used in this embodiment is shown in more detail in Figures 2 and 3. Compressed air is supplied via port (17) to the nebulizer jet (11), and this works in conjunction with baffle (12) to aerosolize the liquid drug which has been placed in nebulizer bowl (19).

This nebulizer is venturi nebulizer which draws in air through the spout (13) into the centre of the baffle (12), and then up and out through the outlet of the spout (13) into the mouthpiece. As the nebulizer only generates aerosol during inspiration, the patients inhaled air is drawn through valve (14), and further through the

5 mouthpiece to the patient. A small proportion of this air
is drawn down through the centre of the nebulizer to
operate the venturi system. The flow through the
nebulizer is completely independent of the patient's flow,
and the nebulizer produces a constant rate of output.
10 Valve (14) creates a pressure drop within the mouthpiece,
and this is monitored through port (18). On exhalation
valve (14) will close, and the patient will exhale through
valve (15). Further information as the nebulizer on the
nebulizer is contained in EP 627,266 (Medic-Aid Limited).

15

The complete assembly is held in place by catch (16).
When the lower end of the catch is displaced the
mouthpiece can be removed. The top end of the catch can
then be tilted down to release the nebulizer bowl assembly
20 from the main dosimeter case (3). This allows the whole
nebulizer unit to be removed after the treatment has been
completed, and cleaned completely separate from the main
dosimeter case. This is further graphically illustrated
in Figures 7 and 8.

25

Figure 4 shows an internal section through the
dosimeter, in particular showing the electronic
configuration of the device. Air enters the dosimeter via
tube (10) and through flexible tube (22) to the manifold
30 (21) and onwards to valve (20). To determine whether
there is pressure in the tube (22), it is configured
around wall (28). When there is no pressure in the tube,
the tube collapses. However, under pressure it expands,
and moves lever (29) against switch (30), which switches
35 the dosimeter on.

5 The dosimeter can be operated from various different
compressed air sources; firstly it can use either a
continuous air supply of approximately 6 litres per
minute, generated from a conventional air compressor, or
10 a compressed bottled gas supply such as is used in
hospitals. With these compressed air sources, when the
compressed air is not being supplied to the nebulizer via
tube (23) and port (17), it must be vented externally
through tube (24) out of the base of the dosimeter via an
orifice which matches the size of the jet (11) diameter in
15 the nebulizer. This maintains a constant pressure in the
system irrespective of whether the valve is directing air
to the nebulizer or the vent.

 In an alternative and preferred embodiment, which can
20 be used in conjunction with other forms of dosimeter, the
dosimeter can be operated in conjunction with a low flow
rate compressor system having a flow rate of 1-2 litres
per minute, which has a fitted accumulator.

25 This compressor system generates a low flow of, for
example, 1.5 litres per minute in the accumulator, and
this allows the dosimeter to produce pulses of compressed
air to the nebulizer with an equivalent flow rate of
around 6 litres per minute, where the flow from the
30 compressor is matched to the mean flow through the
nebulizer (1.5 litres per minute). In this configuration
tube (24) is sealed, and when the valve (20) is closed the
pressure in the supply tube (10) is diverted into the
compressor's accumulator. Port (18) connects the pressure
35 sensor (25) to the mouthpiece. This is attached to the

5 printed circuit board (31) which is powered by battery
(26). When the electronic system detects the patient has
inhaled, valve (20) diverts the pressure flow into tube
(23) and out to a nebulizer via port (17). The battery is
inserted into the dosimeter via door (27) into base (4).

10

An example of a suitable low flow rate compressor
which can be used as described above is shown in Figures
5 and 6, and comprises a small air compressor with a flow
of approximately 1.5 litres per minute, and an accumulator
15 with a volume of about 0.2 litres, which may be formed,
for example, in the tubing between the compressor and the
dosimeter, or using a bellows and spring arrangement
within the compressor casing. Below is described a rubber
hemisphere accumulator which works particularly well. The
20 compressor operates until the accumulator has reached its
capacity, and then switches off. The dosimeter receives
pulses of air from the accumulator until the accumulator
has been emptied to a specified volume (0.1 litre). The
compressor then cuts in and refills the accumulator. The
25 compressor produces a mean output flow rate which is known
to the dosimeter, and the dosimeter controls its supply of
air so that it does not exceed the mean flow rate from the
compressor, and any one pulse delivered by the dosimeter
does not exceed accumulator capacity. This system
30 therefore utilizes the compressor, which is approximately
one-quarter of the size of a conventional compressor
system, and one quarter of the energy requirement. Such
a compressor is also cheaper to
manufacture and provide.

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5 The accumulator in this embodiment should have a
linear pressure to volume relationship over the period of
a pulse delivery, which is typically 1.5 seconds long,
with a nebulizer jet flow of 6 litres per minute, the
accumulator should provide a volume flow of 150
10 millilitres at a constant pressure (1bar +/- 10%). It is
difficult to obtain a steady linear pressure to volume
relationship. Figure 9 shows a natural rubber
hemispherical accumulator. The rubber has a linear load
to extension between an extension of 100% and 500%. A
15 diaphragm 91 is a hemispherical component which is held
trapped between two rings 92 and 93. In Figure 9, the
diaphragm is shown in three positions labelled A, B, C.
Position A is the unloaded position from which the
diaphragm starts and as compressed air is supplied via a
20 port 94, the diaphragm expands to position C with a 200%
extension. The compressed air supply is then stopped
either by the actuation of a microswitch on the diaphragm
surface (not shown) which stops the compressors, or by a
pneumatic microswitch which then vents the compressor
25 output. The diaphragm 91 then stays in this position
until the valve sends a pulse of air to the nebulizer.
When the diaphragm delivers air, the pressure drop during
operation is less than 5%. The diaphragm can supply air
at a stable pressure until it reaches position B which is
30 a 120% extension. The microswitch would then restart the
compressor, or the pneumatic microswitch would be closed.
This hemisphere arrangement is relatively simple to
manufacture, but has great advantages over other
accumulator systems which might be used.

5 A suitable electrical configuration for such a
compressor is shown in Figure 6.

10 For the dispensing system according to the invention
to work effectively, the dosimeter valve (20) should be in
close proximity to the nebulizer, so that when the system
detects the patient's inhalation, the nebulizer starts to
work as quickly as possible, typically in less than 50
15 milliseconds. This means that the length of tube (23)
between the manifold outlet and the nebulizer jet must be
short, with an internal volume less than 0.7 ml,
preferably less than 0.5 ml, which represents 5% of the
shortest pulse the nebulizer delivers. If a long tube is
used between the valve and the nebulizer, this tube has to
be pressurized before the nebulizer starts to operate.
20 This significantly affects the performance of the system,
as to control the rate of output of the nebulizer it must
be supplied with a "square wave" of air pressure, so that
the output is constant over time, and is delivered at the
start of inhalation. The nebulizer delivers pulses to
25 inspiratory patterns between 0.1 and 1.5 seconds'
duration.

30 One of the simplest ways of determining the dose of
medicament which is received by the patient is, as
described as above, to multiply the nebulizer output rate
by the duration of each pulse. The doses then ascertained
by summing the amount of medicament which is received
during each pulse. This calculation relies on the fact
that the output rate of a nebulizer should be constant
35 regardless of the rate of inhalation of the patient.

5 Therefore, provided that the nebulizer output rate is the same for a person who inhales slowly as for a patient who inhales quickly, the calculation will be accurate. If the nebulizer output rate varies with the speed of inhalation, the precision of the dosimeter will vary.

10

Figure 10 is a graph showing the variation of aerosol output rate with the speed of inspiratory flow for a typical nebulizer. As will be seen, the output is not constant.

15

It is therefore proposed to use the pressure sensor to provide information on the patient flow rate so that the correct nebulizer calibration rate is determined during patient inhalation. This may be done in the form of a look-up table which is quite effective, and the look-up table can have two or more calibration points as required to provide the necessary accuracy. A satisfactory look-up table can be achieved by using an approximation of the look-up table which, in the case of the graph shown in Figure 10 can be made up of two straight lines, one line generally following the curve to the 30 lpm point and a second line which generally follows the curve above that. Such an approximation works well because, in reality, the breathing pattern of a patient is not at a fixed level, but is continually changing.

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Another way of using the correct calibration value is to use the computer which takes the inspiratory flow rate into account. One calibration value can be used above the 30 litres per minute level, and when the flow is below

5 about 15 litres per minute, then the calibration constant
is reduced to 60% of that value. Thus, the calibration
may be achieved in a number of different ways not just a
multi-point look-up table.

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CLAIMS

1. A product dispensing system comprising a calibrated nebulizer (2) having a nebulizer jet (11), a mouthpiece (1), means (10, 22) for sourcing compressed air, a manifold (21) for distributing the compressed air in at least one direction, a port (17), and a valve means (20) for controlling the manifold (21) and thereby the flow of compressed air to the port (17), the manifold (21) and the port (17) being linked by a length of tube (23), characterised by a nebulizer accommodation means which is accessed by the port (17), and in that the internal volume of the tube (23) from the manifold outlet to the nebulizer jet (11) is less than 0.7 ml.
2. The dispensing system as claimed in claim 1, pre-programmed to dose one or more medicaments according to predetermined dosage profiles.
3. The dispensing system as claimed in claim 1 or 2, additionally comprising a switch means (30) which is responsive as to whether compressed air is flowing in the system.
4. The dispensing system as claimed in any one of claims 1 to 3, further comprising a dosimeter which has been pre-programmed to dose one or more medicaments according to predetermined dosage profiles.
5. The dispensing system according to any one of claims 1 to 4, further comprising a pressure sensor (25) which is capable, in use, of detecting a pressure drop in the apparatus in response to the patient's breathing, whereby a pulse of nebulised medicament may be delivered into the mouthpiece (1).
6. The dispensing system according to claim 5, further comprising means for calculating a dosage of medicament on the basis of the known rate of medicament output against time for a selected drug.
7. The dispensing system according to any one of claims 1 to 4, further comprising a pressure sensor for detecting the rate of inhalation of air through the system and means for calculating a dosage of medicament delivered on the basis of the detected rate of inhalation over time.

8. The dispensing system according to claim 6 or claim 7, further comprising an indicator arranged to indicate when the dosage of medicament has been delivered.
- 5 9. The dispensing system according to claim 8, further comprising means for determining when a pre-set dose has been delivered and for controlling the indicator.
- 10 10. The dispensing system according to any one of claims 1 to 9, wherein the manifold (21) is an integral part of the valve means (20).
- 10 11. The dispensing system according to any one of claims 1 to 10, wherein the manifold (21) together with the valve means (20) serve to direct the flow of compressed air either to the nebuliser (2) or to an outlet orifice.
- 15 12. The dispensing system according to any one of claims 1 to 10, where the manifold (21) is a shaped block with integral galleys the presence of which allows the tubes to be connected to the ports on the valve.
- 20 13. The dispensing system according to any one of claims 1 to 10 and 12, further comprising a compressor and an accumulator (90).
- 25 14. The dispensing system according to claim 13, wherein the compressor is a lightweight compressor.
- 30 15. The dispensing system according to claim 14, wherein the lightweight compressor generates a low flow rate of the order of 1 to 2 litres per minute.
16. The dispensing system according to claim 14 or 15 wherein the combination of the low flow rate compressor and the accumulator (90) allows the dispensing system to produce pulses of compressed air to the nebuliser (2) with a flow of around of 6 litres per minute.
17. The dispensing system according to any one of claims 13 to 16, wherein the valve means (20) serves to switch the compressed air on and off.
- 35 18. The dispensing system according to any one of claims 13 to 17, wherein the accumulator (90) includes a resilient elastic body (91) having a generally linear pressure to volume relationship .

19. The dispensing system according to claim 18, wherein the resilient body (91) is a generally hemispherical resilient element.
- 5 20. The dispensing system according to claim 18 or claim 19, wherein the resilient body (91) is a diaphragm extendable to 500% of its unloaded volume.
21. The dispensing system according to any one of claims 1 to 20, wherein the volume of the tube (23) linking the manifold outlet to the nebuliser jet (11) is less than
10 0.5 ml.
22. The dispensing system according to any one of claims 1 to 21, wherein the nebuliser (2) serves to deliver inspiratory pulses between 0.1 and 1.5 seconds in duration.
- 15 23. A product dispensing system comprising a calibrated nebulizer (2) having a nebulizer jet (11), a mouthpiece (1), means (10,22) for sourcing compressed air, a manifold (21) for distributing the compressed air in at least one direction, a port (17), and a valve means (20) for controlling the manifold (21) and thereby the flow of
20 compressed air to the port (17), the manifold (21) and the port (17) being linked by a length of tube (23), characterised by:
- i) a nebulizer accommodation means which is accessed by the port (17),
 - ii) a compressor,
 - iii) an accumulator (90), wherein the accumulator (90) includes a resilient
25 elastic body (91) having a generally linear pressure to volume relationship and
 - iv) the tube (23) from the manifold outlet to the nebulizer jet (11) having an internal volume of less than 0.7ml.
- 30 24. A product dispensing system comprising a calibrated nebulizer (2) having a nebulizer jet (11), a mouthpiece (1), means (10,22) for sourcing compressed air, a manifold (21) for distributing the compressed air in at least one direction, a port (17), a valve means (20) for controlling the manifold (21) and thereby the flow of compressed air to the port (17), the manifold (21) and the port (17) being linked by a length of tube
35 (23), characterised by:
- i) a nebulizer accommodation means which is accessed by the port (17),

- 5
- ii) a pressure sensor (25) for detecting a drop in pressure in the system in response to a patient's breathing to indicate the rate of inhalation of the patient,
 - iii) the tube (23) from the manifold outlet to the nebulizer jet (11) having an internal volume of less than 0.7ml and
 - iii) means for determining the dose dispensed, the dose determined on the basis of the output rate of the medicament of the nebuliser (2), and the duration of the delivery of the medicament, the dose determining means serving to control the valve.

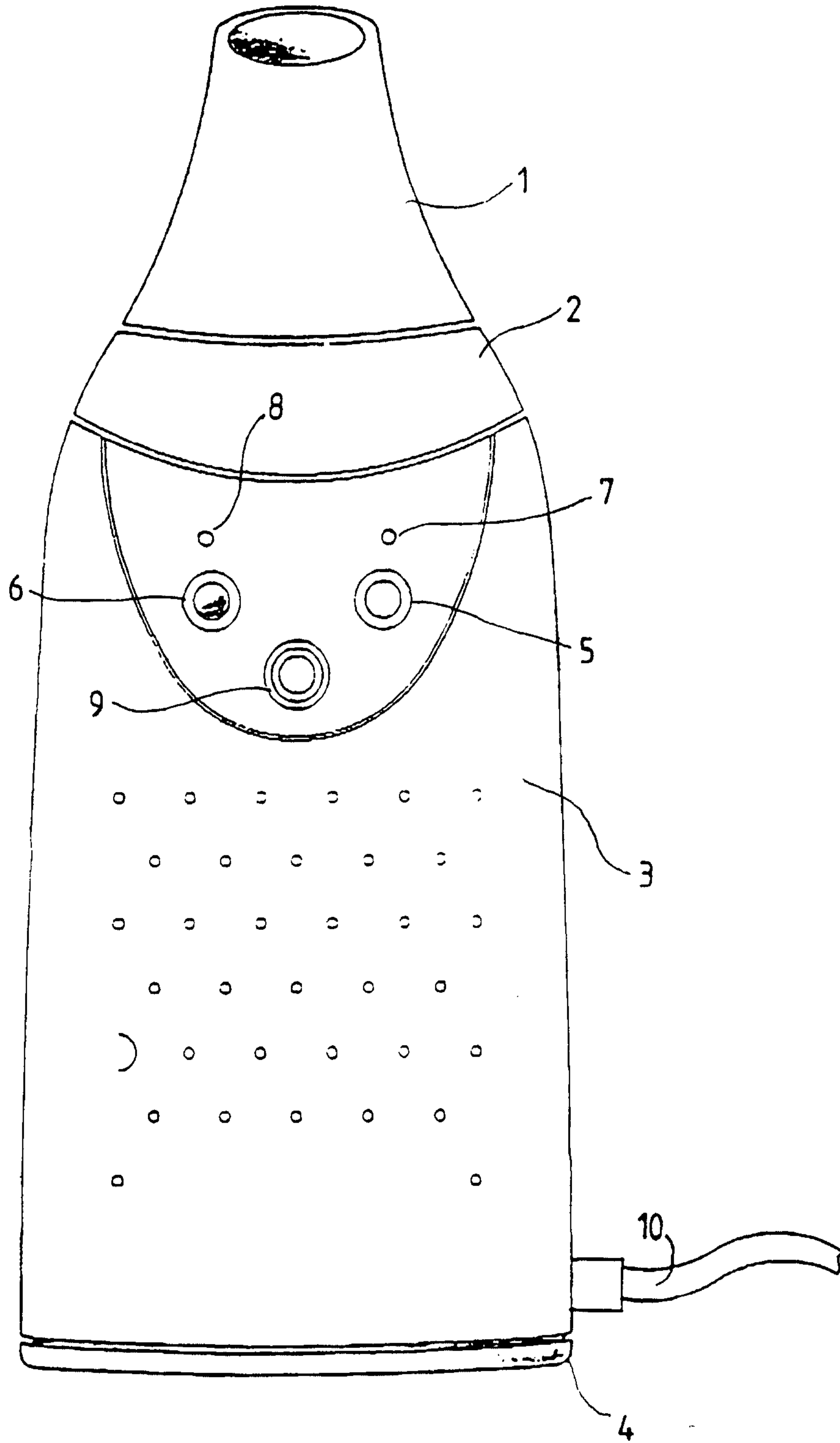


FIG.1

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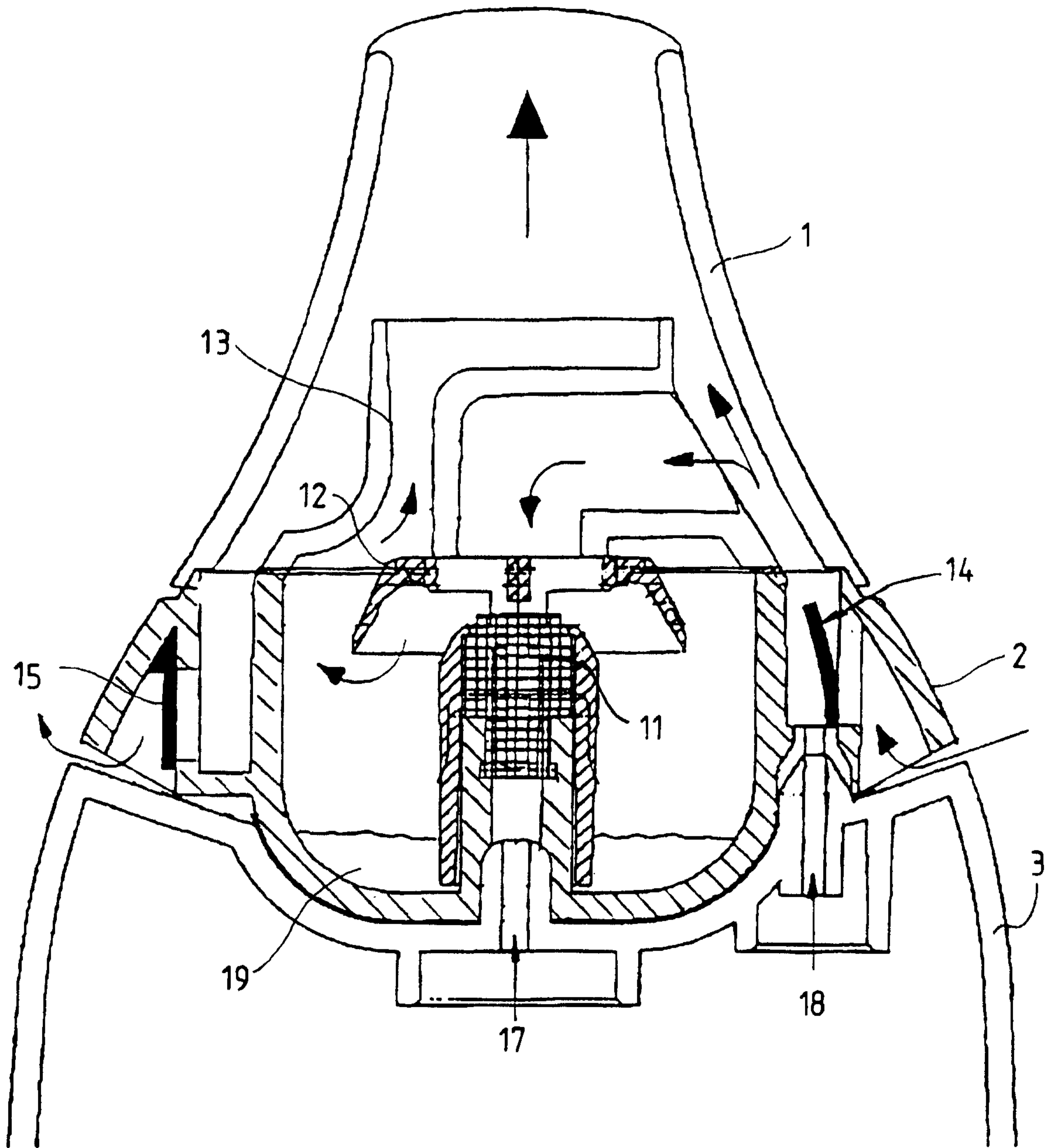


FIG. 2

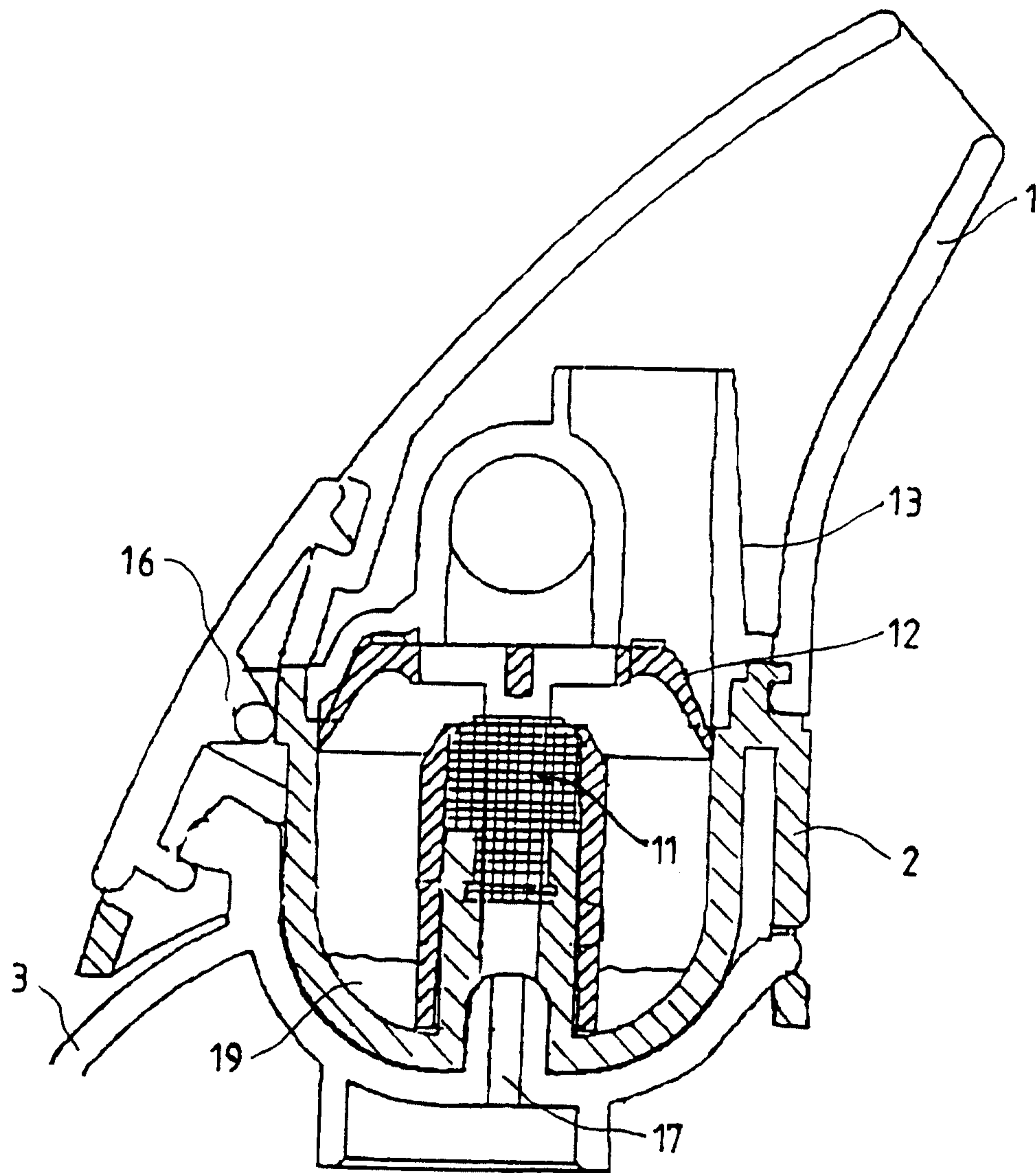


FIG. 3

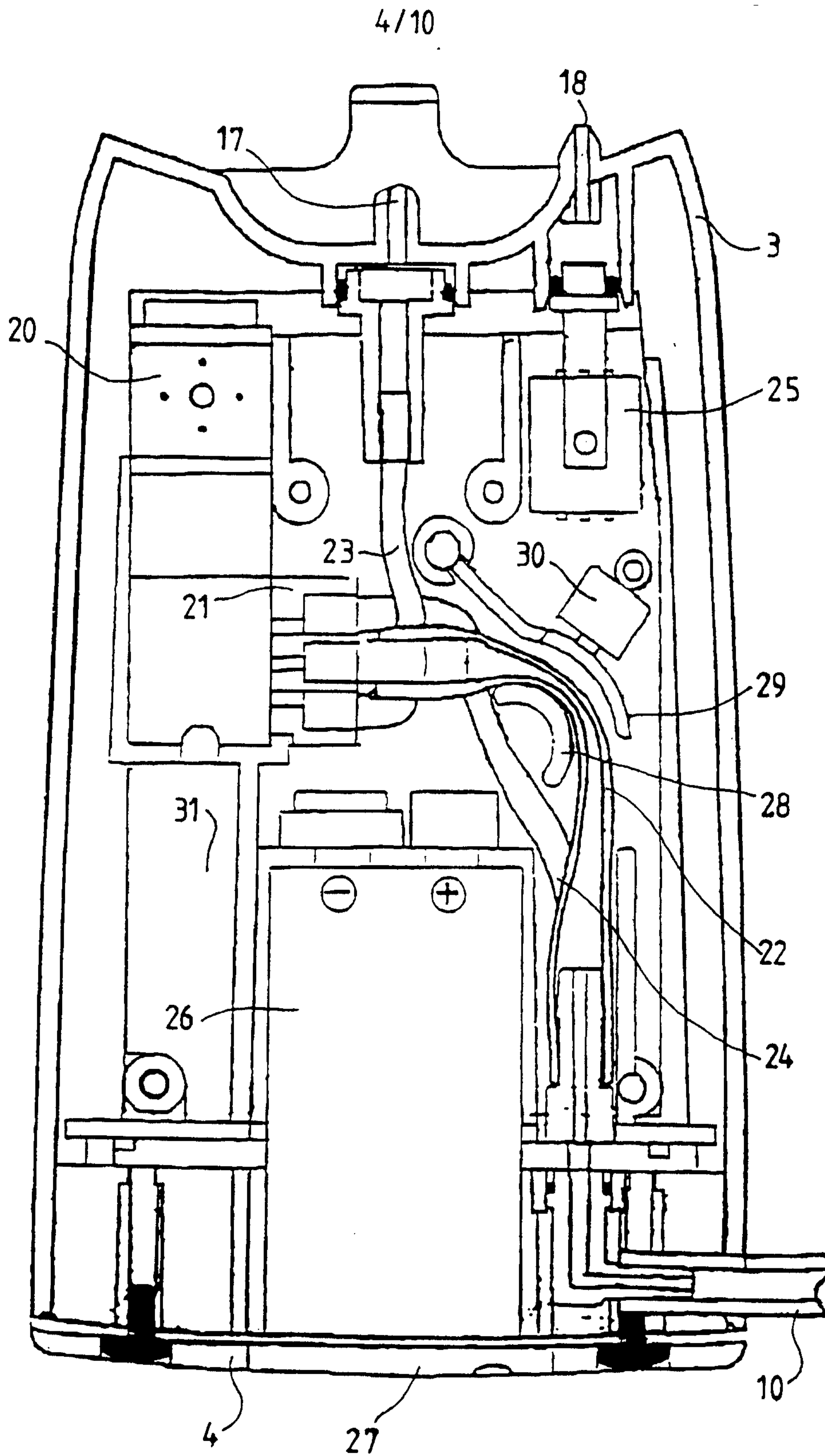


FIG. 4

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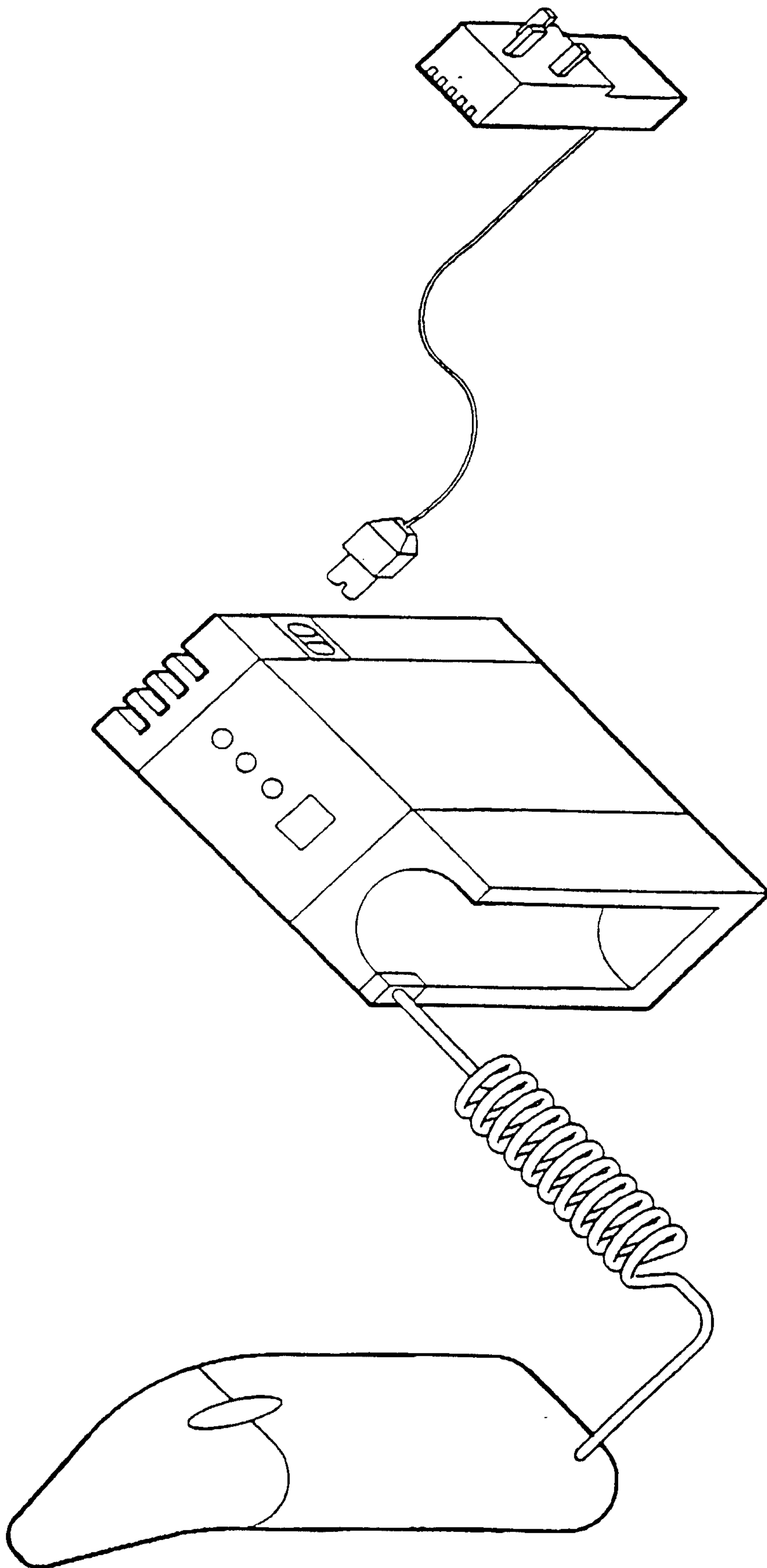


FIG. 5

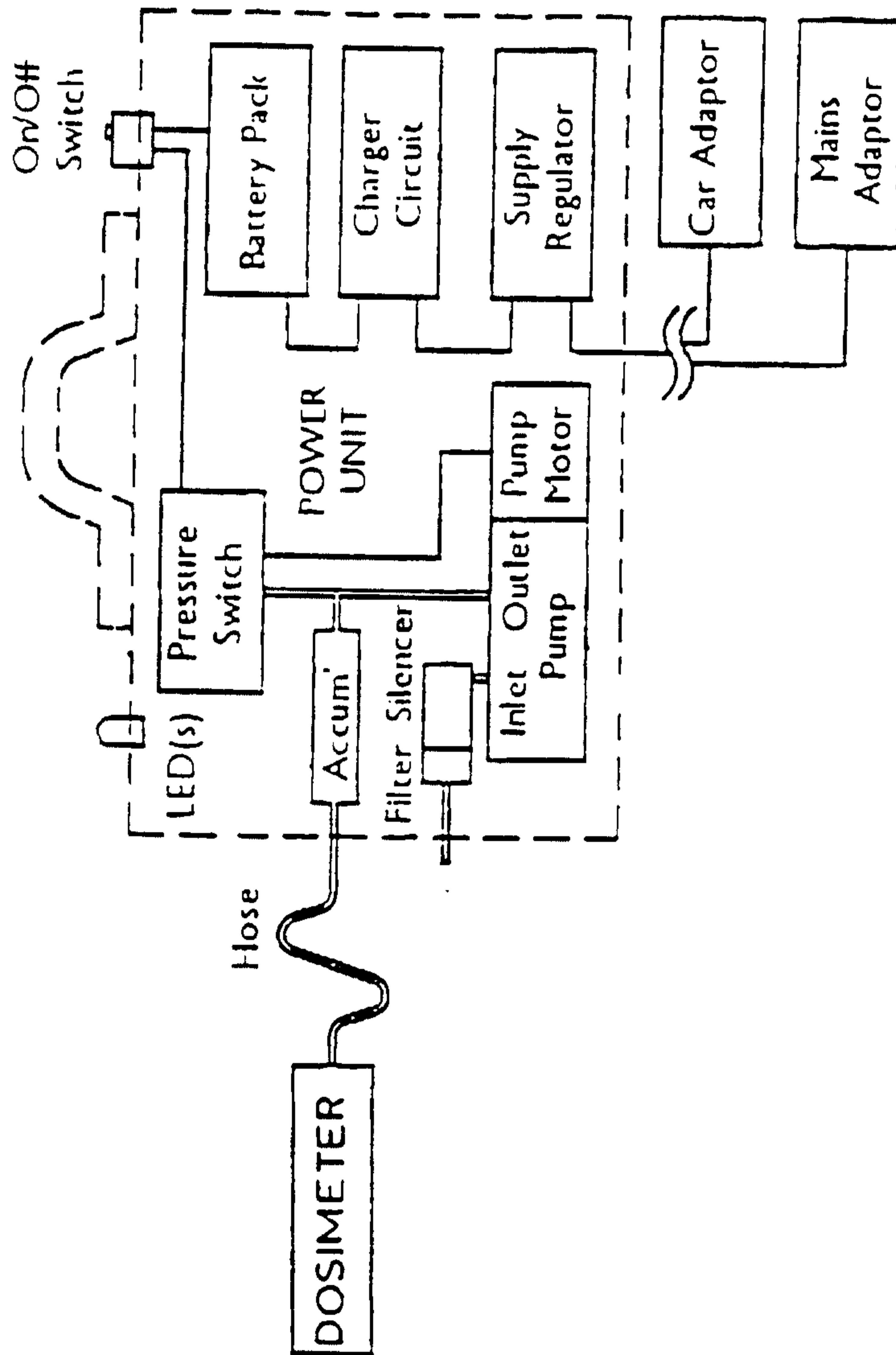
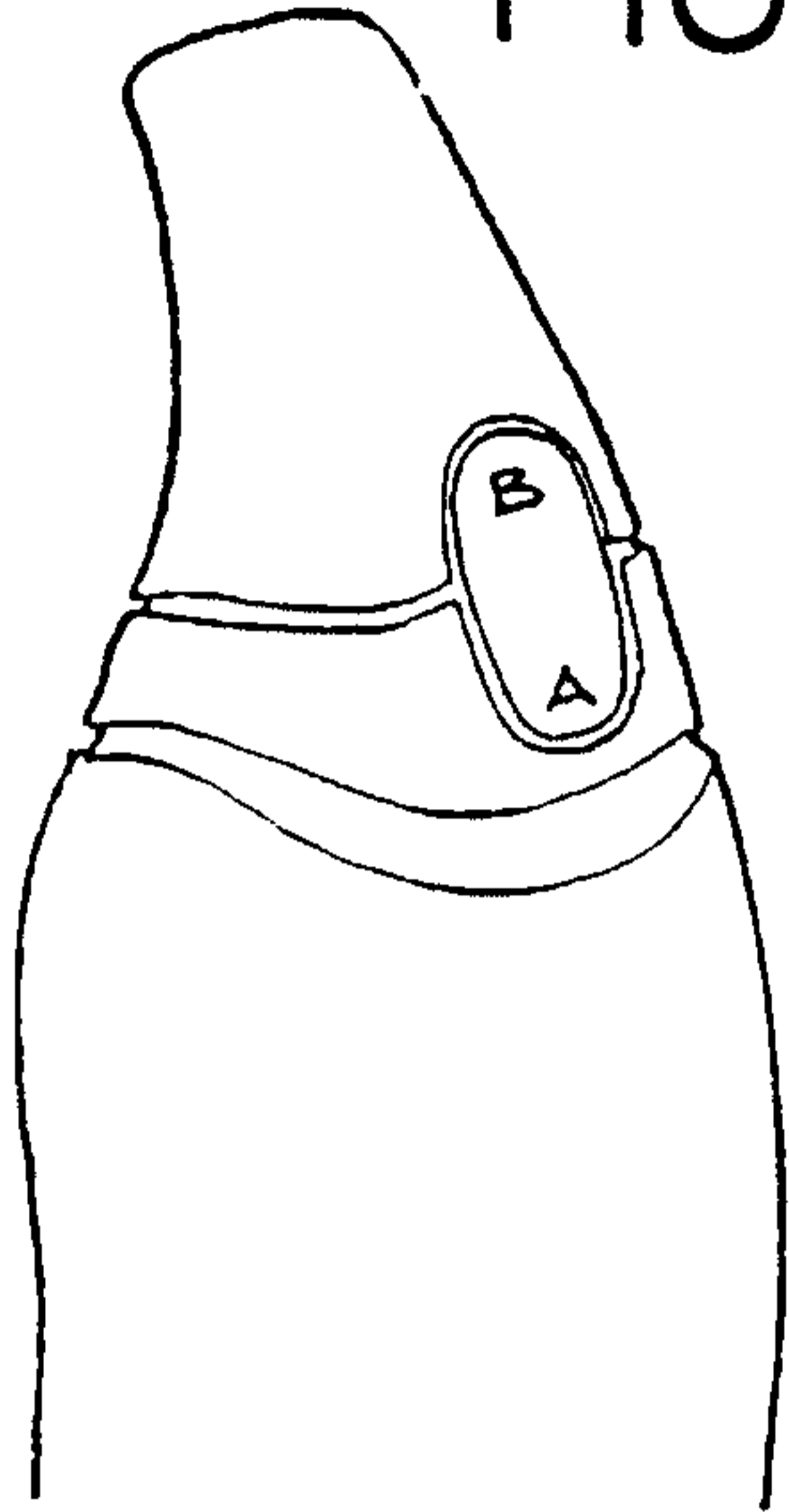


FIG. 6

TO LOAD

FIG 7a.



NEBULIZER CLEAN & EMPTY

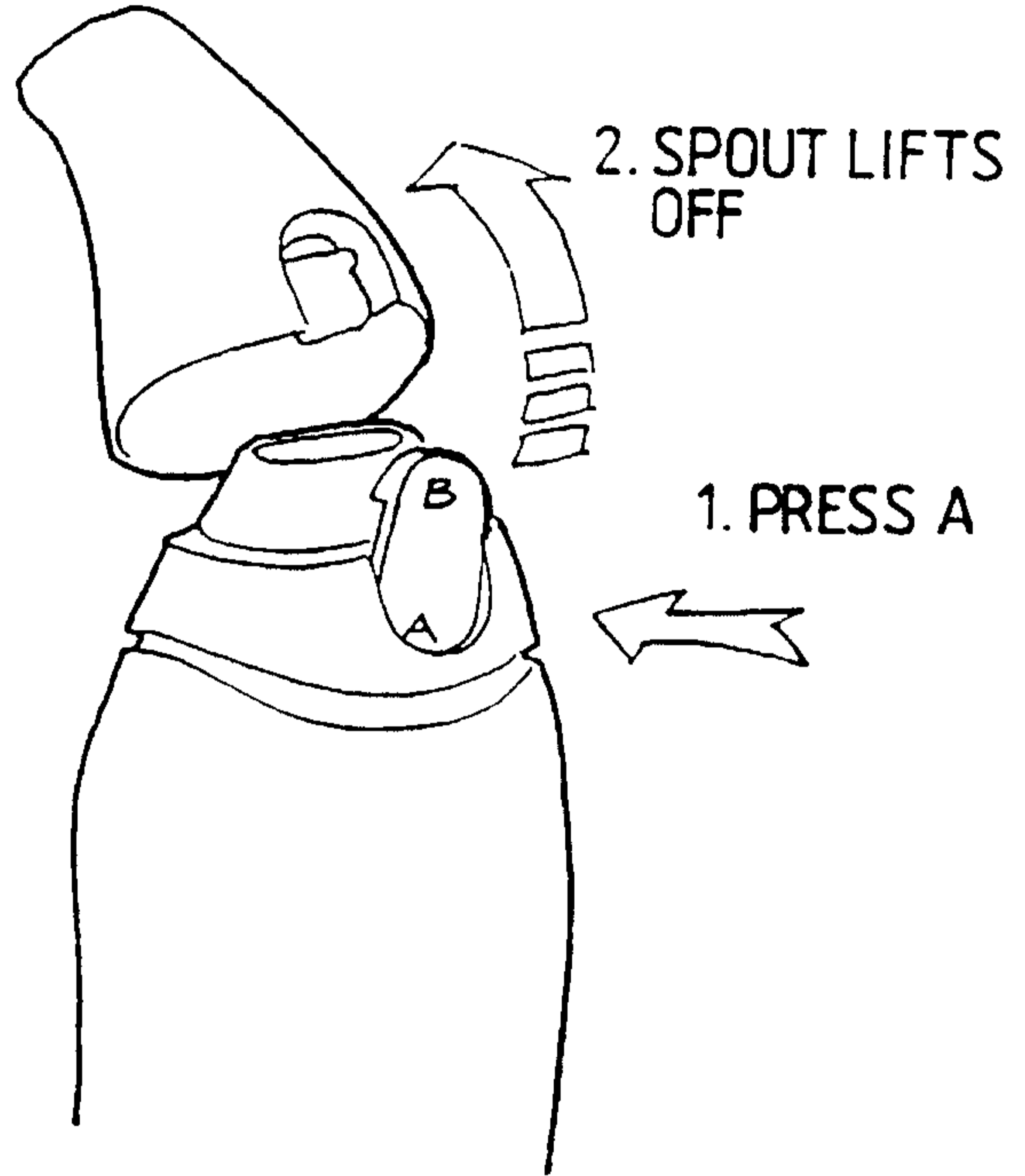
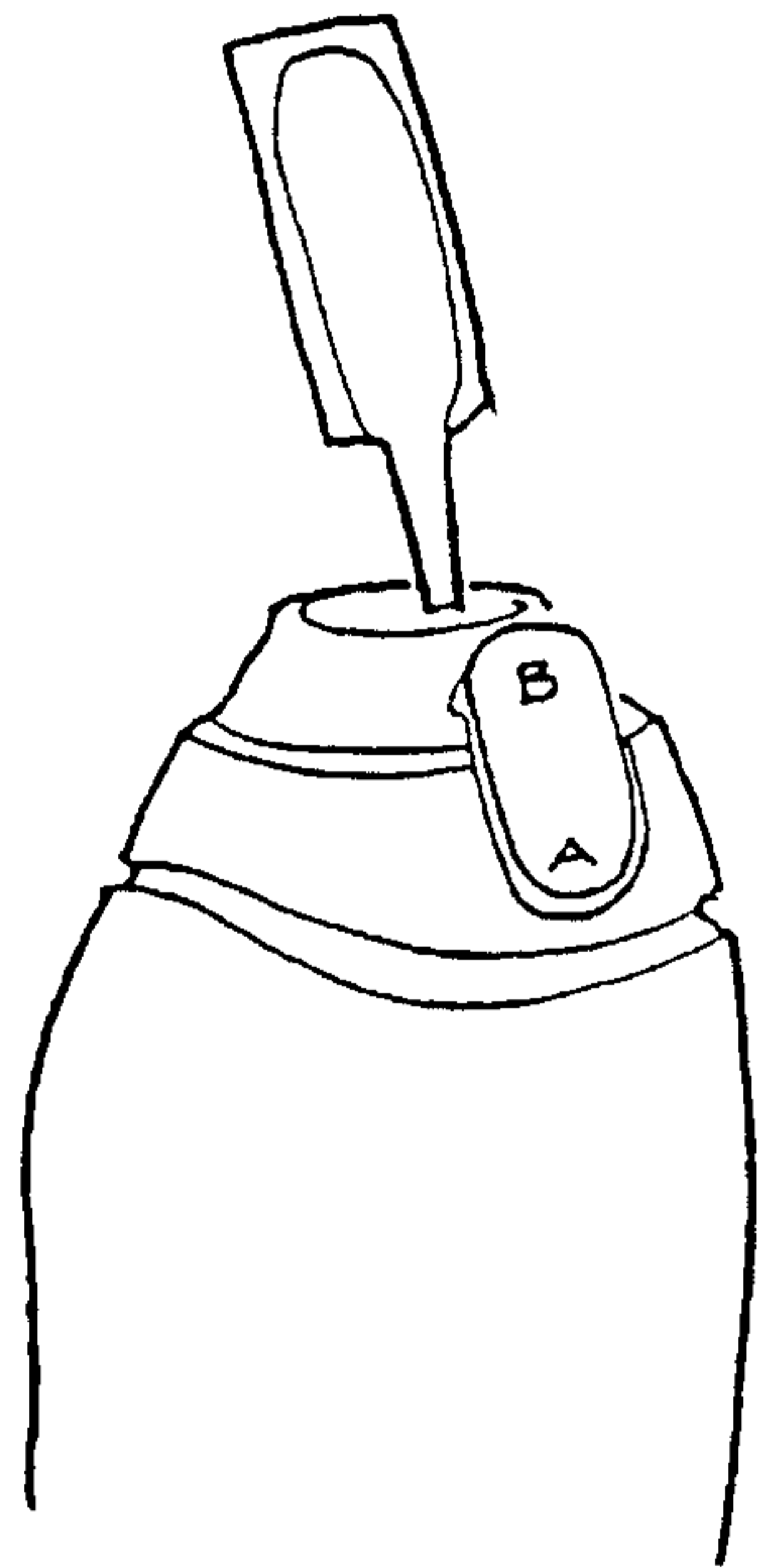
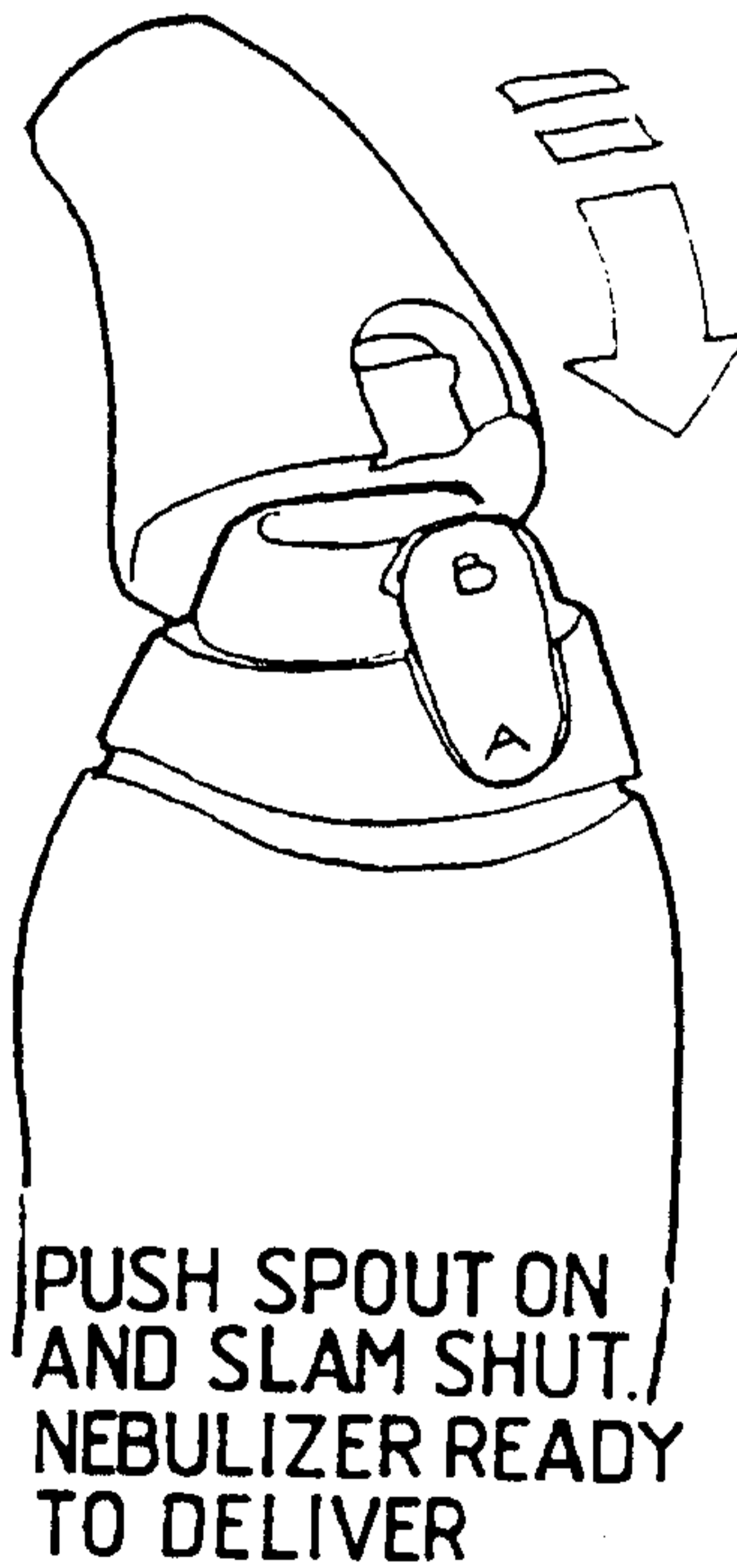


FIG.7b.



CHARGE THROUGH FUNNEL

FIG.7c.



PUSH SPOUT ON AND SLAM SHUT. NEBULIZER READY TO DELIVER

FIG.7d.

TO CLEAN

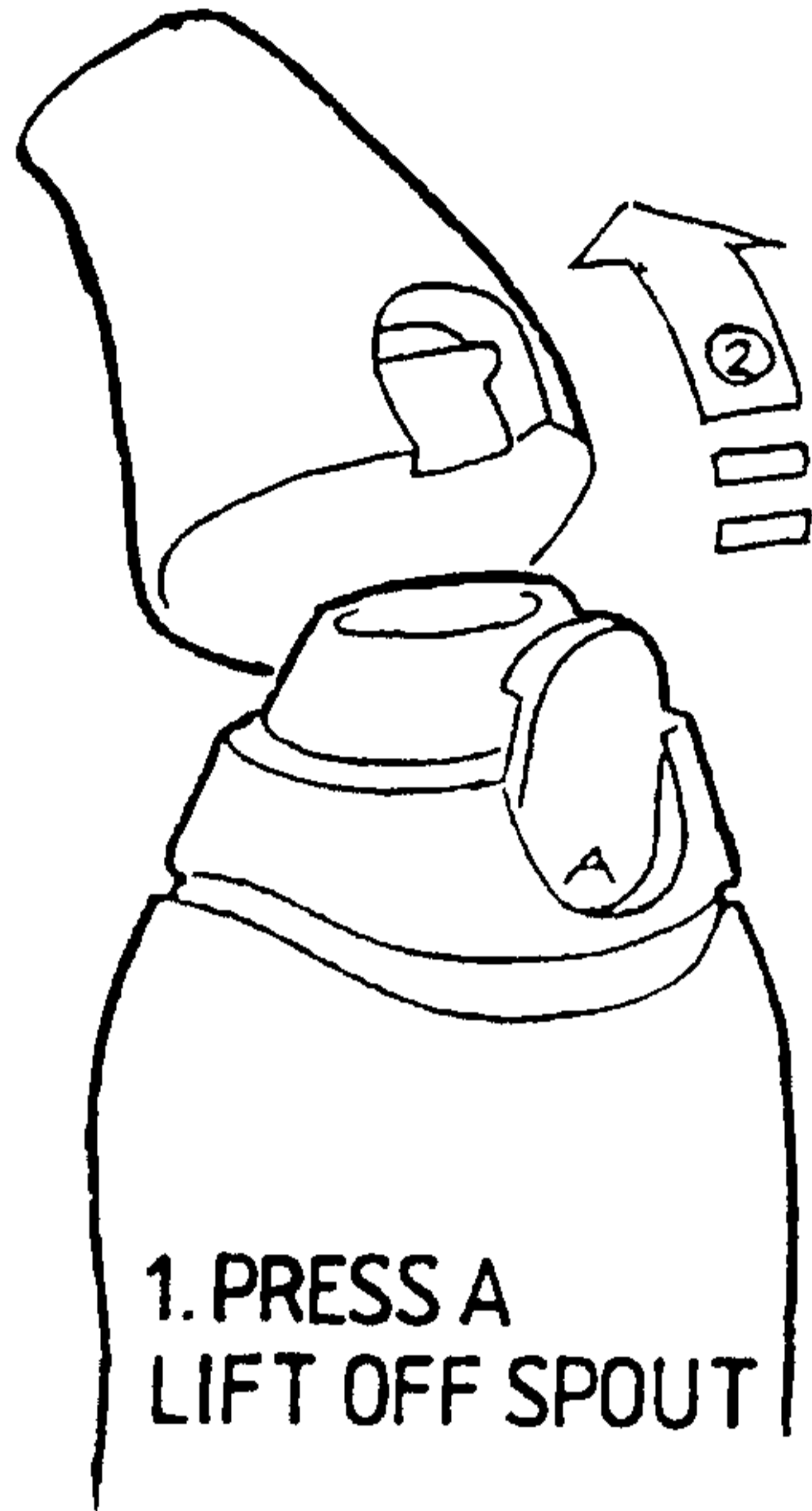


FIG. 8a.

LIFT OUT FUNNEL

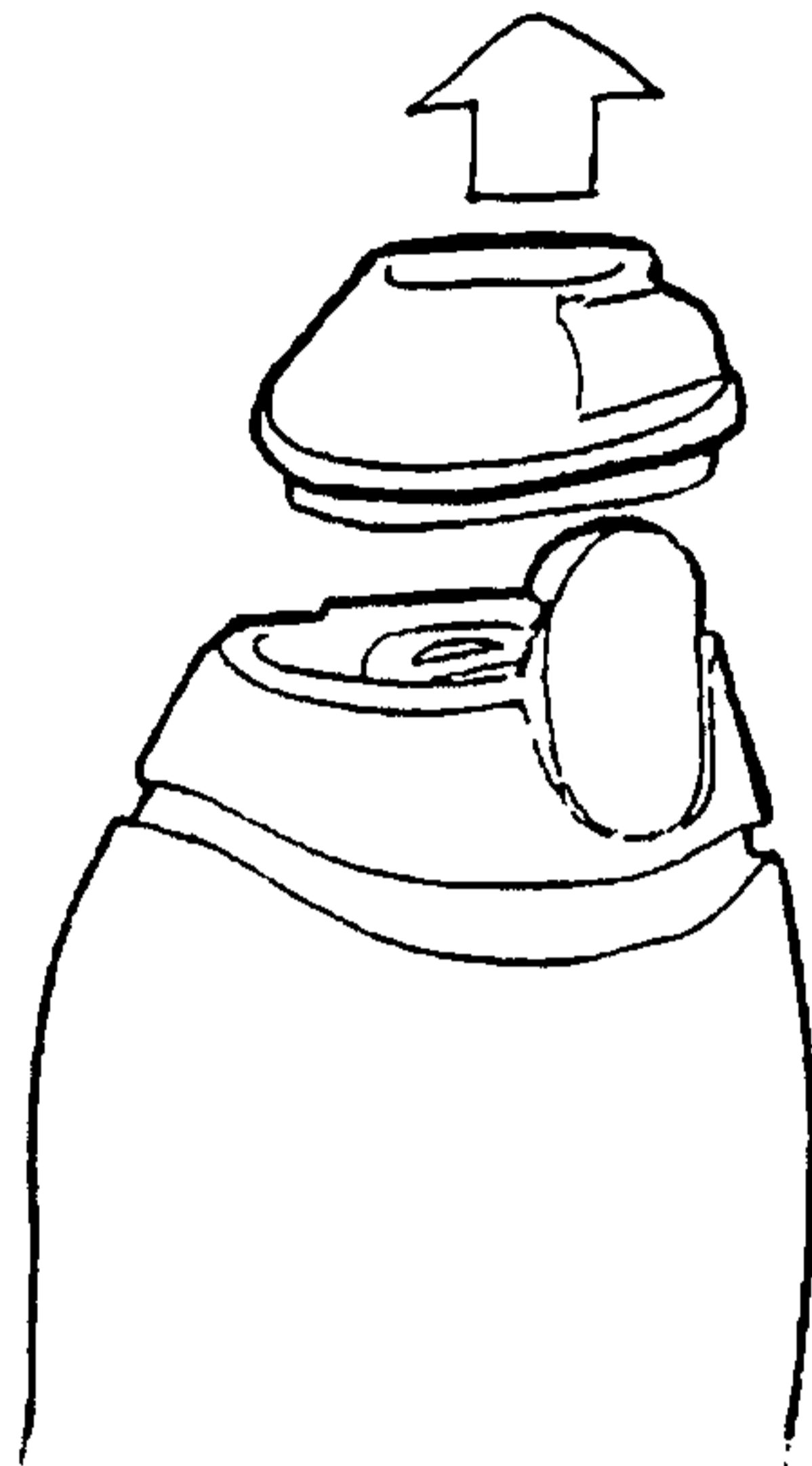


FIG. 8b.

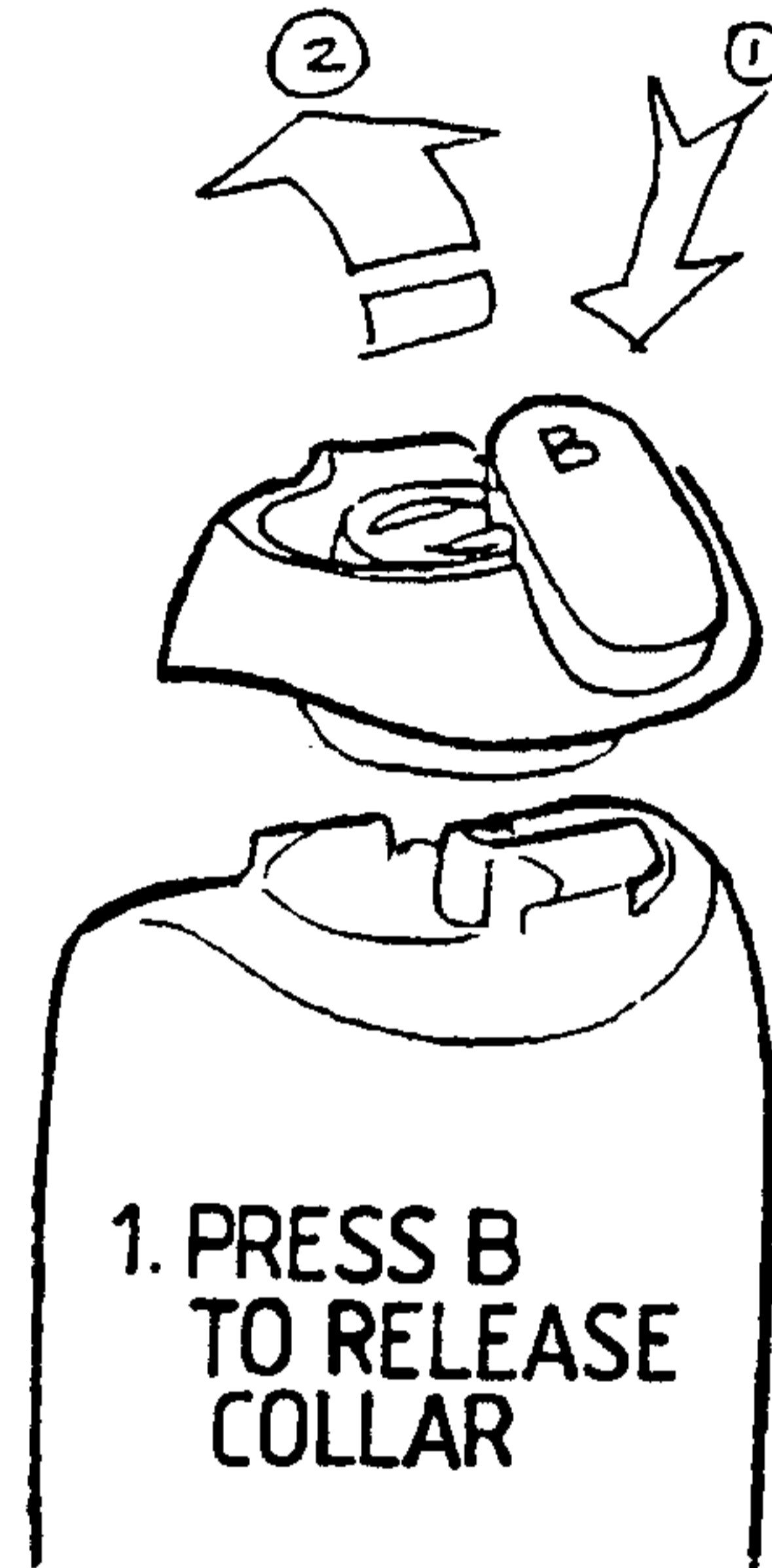
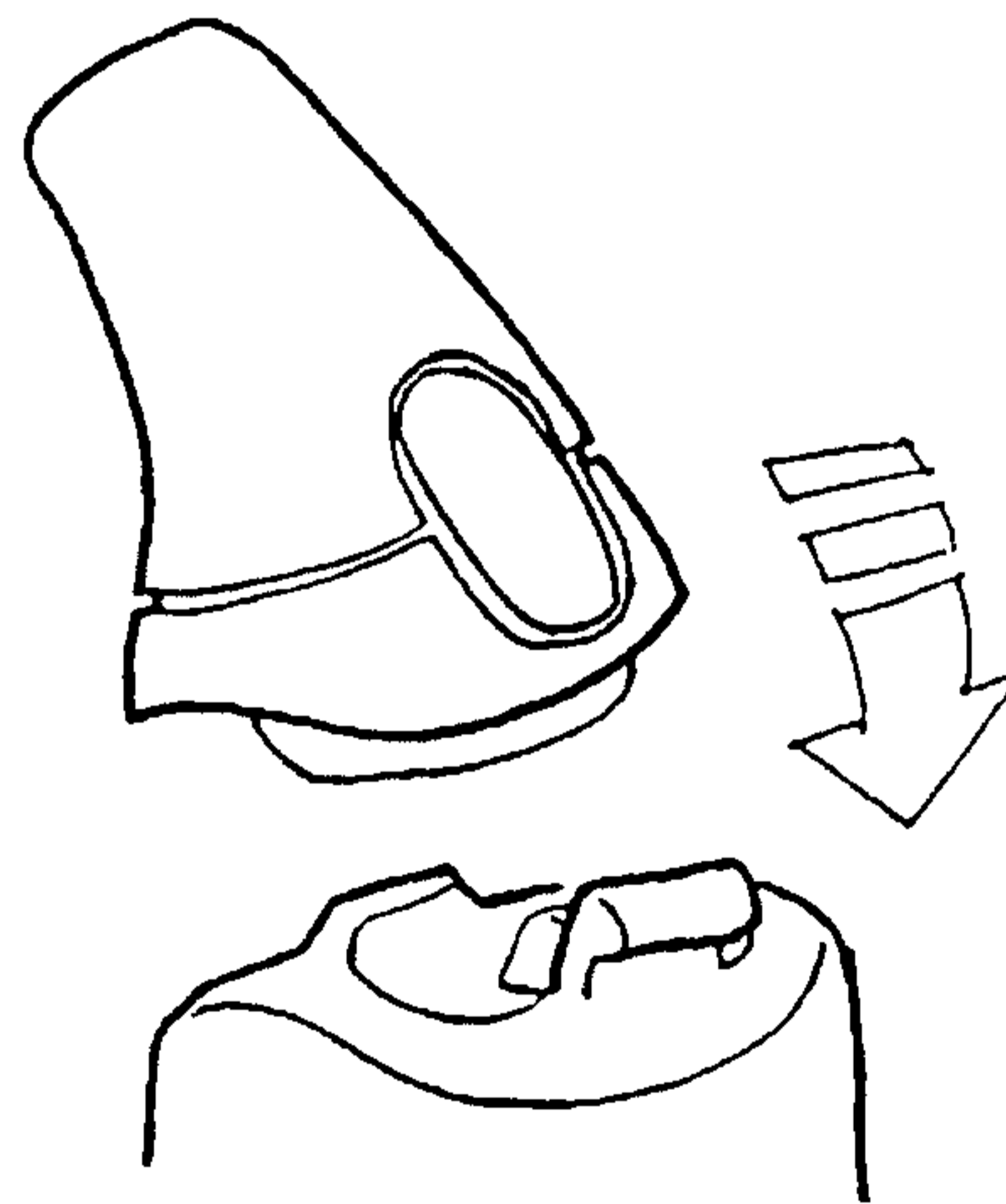


FIG. 8c

CLEAN & DRY
ALL PARTS
THEN EITHER...

REASSEMBLE BY STEPS
SHOWN IN FIGURES 8c,
8b,8a IN REVERSE ORDER



B. SNAP COLLAR,
FUNNEL & SPOUT
TOGETHER
AS A UNIT
THEN SNAP UNIT
ONTO BODY

FIG. 8d.

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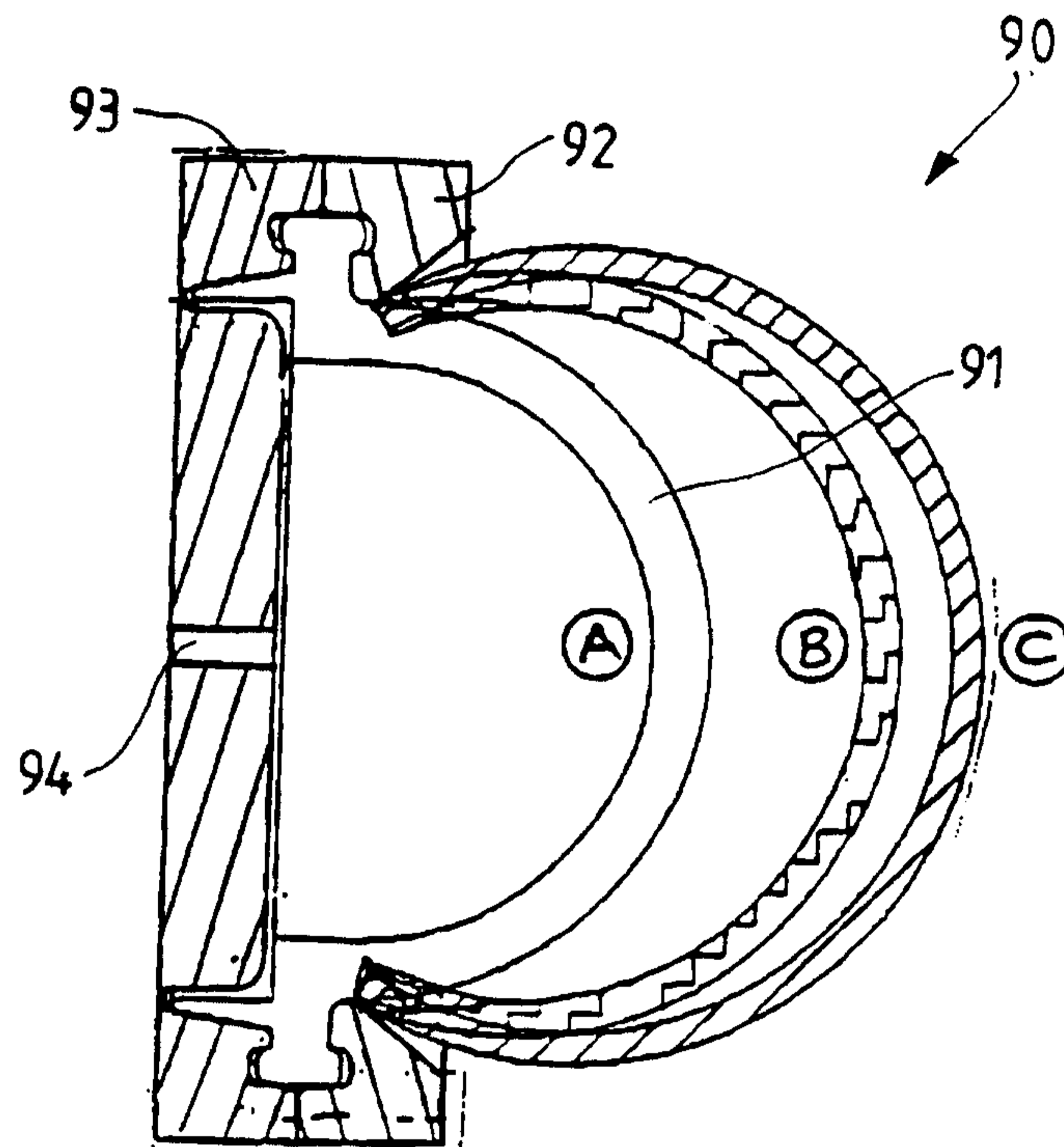


FIG. 9

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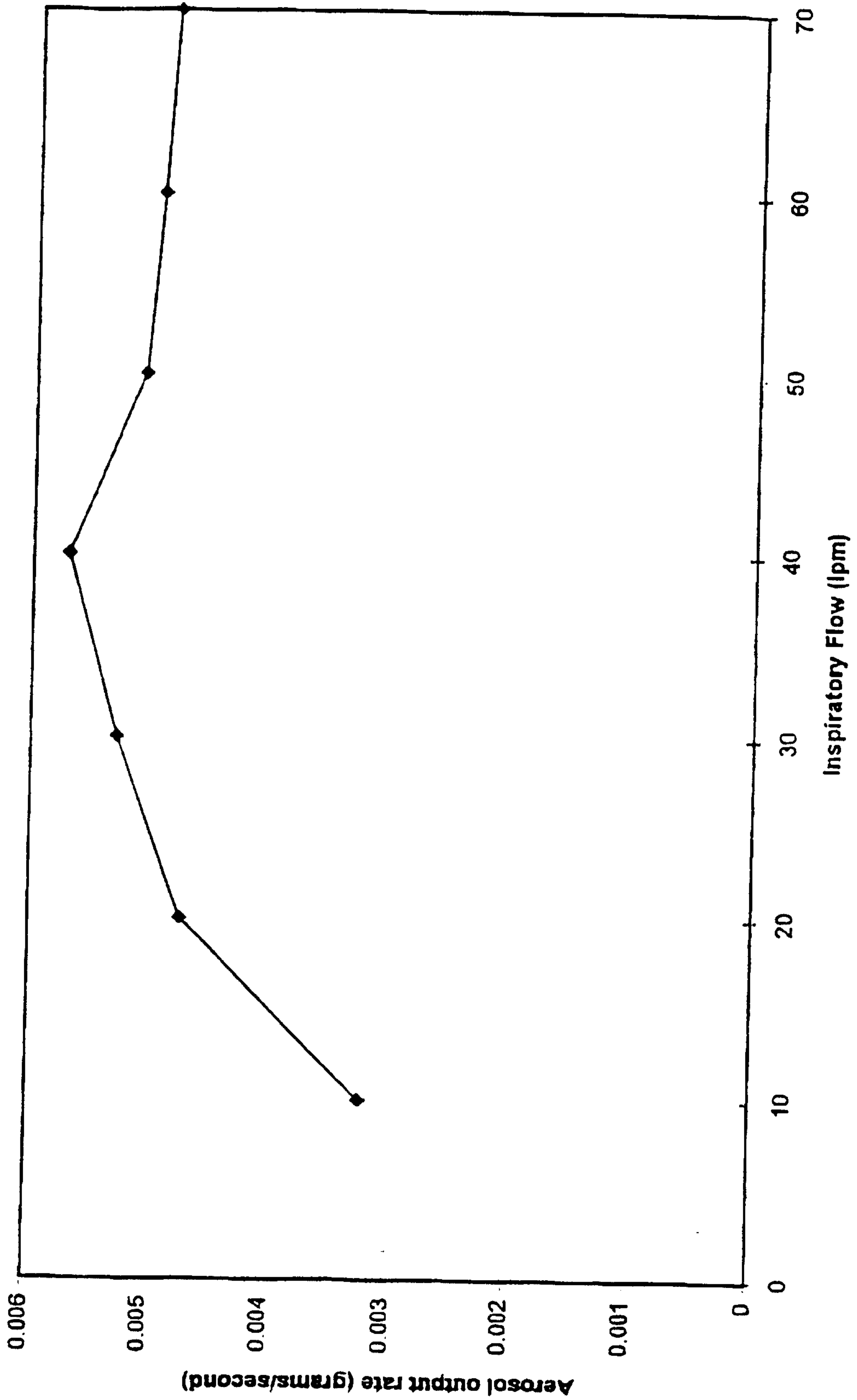


FIG.10

