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DESCRIPTION

Description

Technical Field

[0001] The invention relates to a medicament delivery device and an actuation mechanism for a medicament delivery device.

Background of the Invention

[0002] Conventional medicament delivery devices containing a selected dose of a medicament are well-known devices for administering the medicament to a patient. A conventional delivery device comprises a needle to administer the medicament. Safety devices for covering a needle of the delivery device before and after use are also well known. In a conventional safety device, a needle shield is moved either manually or automatically (i.e., by spring) to cover the needle.

[0003] A specific type of a medicament delivery device is an autoinjector, which is equipped with an actuation button to actuate automatic delivery of the medicament. To administer the medicament, the autoinjector is pressed against an injection site, which retracts the needle shield. When the actuation button is pressed, the needle is inserted into the injection site and the medicament is administered. The conventional delivery device, thus, requires two acts - pressing of the delivery device to injection site and pressing the actuation button. It may be difficult to perform either or both of these acts when the patient/operator has lessened dexterity, e.g., due to age, disability, illness, sensory deficiency, etc.

[0004] Other conventional delivery devices are actuated upon contact with the injection site. These devices are pressed against the injection site, which retracts the needle shield, and pressed with enhanced force to initiate delivery of the medicament. However, patients may be confused by these types of delivery devices, because there is no actuation button.

[0005] WO 94/21316 A1 discloses an injection device in which a charged capsule with a needle at its forward end and a plunger projecting from its rear end is housed in an elongate body with coil spring means which, when released from compression, act on the plunger to push that forwards and thereby carry out the injection, and a release mechanism for the spring means including a sleeve over a substantial part of the body initially retained in or urged into a rearward position but which has to be shifted forwards in relation to the body to allow the

spring means to be released.

[0006] WO 2006/111861 A2 discloses an injection assistance device for an injection device for injecting a product into an injection site. The injection device comprises at least one hollow body intended to receive a product that is to be injected, at least one hollow injection needle intended to penetrate the injection site, and at least one piston plunger housed in the said body, the said body and the said piston plunger being able to be moved in axial translation one with respect to the other.

[0007] DE 10 2004 053529 A1 discloses a device for blood collection which has a drive unit coupled with a piston, which is movable in a blood container, where the piston can be moved by the drive unit.

[0008] EP 1 949 928 A1 discloses a retracting safety injector pen needle having the needle mounted on a carrier which is movable with respect to the hub, so that the non-patient end of an injection pen needle device is retracted into the hub after an injection for safety.

[0009] US 2003/050606 A1 discloses a needle covering device with telescopically arranged sleeve sections, wherein retraction of the telescopic sleeve sections closes a contact for automatically delivering a fluid product.

Summary of the Invention

[0010] It is an object of the present invention to provide an actuation mechanism for a medicament delivery device for easy and safe medicament delivery.

[0011] This object is achieved by the subject-matter of independent claim 1.

[0012] In an exemplary embodiment, an actuation mechanism for a medicament delivery device has a needle with a distal tip. The actuation mechanism comprises an outer sleeve telescopically arranged relative to the delivery device and an inner sleeve telescopically arranged relative to the outer sleeve. The outer sleeve is axially translatable relative to the delivery device, and the inner sleeve is axially translatable relative to the outer sleeve. In a first state, the inner sleeve protrudes distally from the outer sleeve and the outer sleeve protrudes distally from the delivery device. In a second state, the inner sleeve is contained within the outer sleeve. Movement of the outer sleeve proximally relative to the delivery device in the second state initiates delivery of a medicament in the delivery device. The actuation mechanism further comprises a first spring element biasing the inner sleeve in a distal direction relative to the outer sleeve. The actuation mechanism further comprises a second spring element biasing the outer sleeve in a distal direction relative to the delivery device. The second spring element is a harder compression spring than the first spring element.

[0013] In an exemplary embodiment, the inner sleeve and the outer sleeve have different

colors or indicia.

[0014] In an exemplary embodiment, the outer sleeve is positionally fixed relative to the delivery device in the first state. The inner sleeve engages the outer sleeve in the second state. The inner sleeve includes a latch adapted to engage a recess or opening in the outer sleeve. The outer sleeve includes a latch adapted to engage a recess or opening in the inner sleeve.

[0015] In an exemplary embodiment, when in a third state, the inner sleeve is locked relative to the outer sleeve and the outer sleeve is locked relative to the delivery device.

[0016] In an exemplary embodiment, a drug delivery device comprises an actuation mechanism according to any one of the exemplary embodiments described above, and further includes a needle having a distal tip. In the first state, the inner sleeve and/or the outer sleeve cover the distal tip, and in the second state, the distal tip is adapted to protrude distally relative to the outer sleeve. In the third state, the inner sleeve and/or the outer sleeve cover the distal tip of the needle.

[0017] The term "drug" or "medicament", as used herein, means a pharmaceutical formulation containing at least one pharmaceutically active compound,

wherein in one embodiment the pharmaceutically active compound has a molecular weight up to 1500 Da and/or is a peptide, a proteine, a polysaccharide, a vaccine, a DNA, a RNA, an enzyme, an antibody or a fragment thereof, a hormone or an oligonucleotide, or a mixture of the above-mentioned pharmaceutically active compound,

wherein in a further embodiment the pharmaceutically active compound is useful for the treatment and/or prophylaxis of diabetes mellitus or complications associated with diabetes mellitus such as diabetic retinopathy, thromboembolism disorders such as deep vein or pulmonary thromboembolism, acute coronary syndrome (ACS), angina, myocardial infarction, cancer, macular degeneration, inflammation, hay fever, atherosclerosis and/or rheumatoid arthritis,

wherein in a further embodiment the pharmaceutically active compound comprises at least one peptide for the treatment and/or prophylaxis of diabetes mellitus or complications associated with diabetes mellitus such as diabetic retinopathy,

wherein in a further embodiment the pharmaceutically active compound comprises at least one human insulin or a human insulin analogue or derivative, glucagon-like peptide (GLP-1) or an analogue or derivative thereof, or exendin-3 or exendin-4 or an analogue or derivative of exendin-3 or exendin-4.

[0018] Insulin analogues are for example Gly(A21), Arg(B31), Arg(B32) human insulin; Lys(B3), Glu(B29) human insulin; Lys(B28), Pro(B29) human insulin; Asp(B28) human insulin;

human insulin, wherein proline in position B28 is replaced by Asp, Lys, Leu, Val or Ala and wherein in position B29 Lys may be replaced by Pro; Ala(B26) human insulin; Des(B28-B30) human insulin; Des(B27) human insulin and Des(B30) human insulin.

[0019] Insulin derivatives are for example B29-N-myristoyl-des(B30) human insulin; B29-N-palmitoyl-des(B30) human insulin; B29-N-myristoyl human insulin; B29-N-palmitoyl human insulin; B28-N-myristoyl LysB28ProB29 human insulin; B28-N-palmitoyl-LysB28ProB29 human insulin; B30-N-myristoyl-ThrB29LysB30 human insulin; B30-N-palmitoyl- ThrB29LysB30 human insulin; B29-N-(N-palmitoyl-Y-glutamyl)-des(B30) human insulin; B29-N-(N-lithocholyl-Y-glutamyl)-des(B30) human insulin; B29-N-(w-carboxyheptadecanoyl)-des(B30) human insulin and B29-N-(w-carboxyheptadecanoyl) human insulin.

[0020] Exendin-4 for example means Exendin-4(1-39), a peptide of the sequence H-His-Gly-Glu-Gly-Thr-Phe-Thr-Ser-Asp-Leu-Ser-Lys-Gln-Met-Glu-Glu-Glu-Ala-Val-Arg-Leu-Phe-Ile-Glu-Trp-Leu-Lys-Asn-Gly-Gly-Pro-Ser-Ser-Gly-Ala-Pro-Pro-Pro-Ser-NH₂.

[0021] Exendin-4 derivatives are for example selected from the following list of compounds:

H-(Lys)4-des Pro36, des Pro37 Exendin-4(1-39)-NH₂,

H-(Lys)5-des Pro36, des Pro37 Exendin-4(1-39)-NH₂,

des Pro36 Exendin-4(1-39),

des Pro36 [Asp28] Exendin-4(1-39),

des Pro36 [IsoAsp28] Exendin-4(1-39),

des Pro36 [Met(O)14, Asp28] Exendin-4(1-39),

des Pro36 [Met(O)14, IsoAsp28] Exendin-4(1-39),

des Pro36 [Trp(O₂)25, Asp28] Exendin-4(1-39),

des Pro36 [Trp(O₂)25, IsoAsp28] Exendin-4(1-39),

des Pro36 [Met(O)14 Trp(O₂)25, Asp28] Exendin-4(1-39),

des Pro36 [Met(O)14 Trp(O₂)25, IsoAsp28] Exendin-4(1-39); or

des Pro36 [Asp28] Exendin-4(1-39),

des Pro36 [IsoAsp28] Exendin-4(1-39),

des Pro36 [Met(O)14, Asp28] Exendin-4(1-39),

des Pro36 [Met(O)14, IsoAsp28] Exendin-4(1-39),

des Pro36 [Trp(O₂)25, Asp28] Exendin-4(1-39),

des Pro36 [Trp(O2)25, IsoAsp28] Exendin-4(1-39),

des Pro36 [Met(O)14 Trp(O2)25, Asp28] Exendin-4(1-39),

des Pro36 [Met(O)14 Trp(O2)25, IsoAsp28] Exendin-4(1-39),

wherein the group -Lys6-NH2 may be bound to the C-terminus of the Exendin-4 derivative;

or an Exendin-4 derivative of the sequence

des Pro36 Exendin-4(1-39)-Lys6-NH2 (AVE0010),

H-(Lys)6-des Pro36 [Asp28] Exendin-4(1-39)-Lys6-NH2,

des Asp28 Pro36, Pro37, Pro38 Exendin-4(1-39)-NH2,

H-(Lys)6-des Pro36, Pro38 [Asp28] Exendin-4(1-39)-NH2,

H-Asn-(Glu)5des Pro36, Pro37, Pro38 [Asp28] Exendin-4(1-39)-NH2,

des Pro36, Pro37, Pro38 [Asp28] Exendin-4(1-39)-(Lys)6-NH2,

H-(Lys)6-des Pro36, Pro37, Pro38 [Asp28] Exendin-4(1-39)-(Lys)6-NH2,

H-Asn-(Glu)5-des Pro36, Pro37, Pro38 [Asp28] Exendin-4(1-39)-(Lys)6-NH2,

H-(Lys)6-des Pro36 [Trp(O2)25, Asp28] Exendin-4(1-39)-Lys6-NH2,

H-des Asp28 Pro36, Pro37, Pro38 [Trp(O2)25] Exendin-4(1-39)-NH2,

H-(Lys)6-des Pro36, Pro37, Pro38 [Trp(O2)25, Asp28] Exendin-4(1-39)-NH2,

H-Asn-(Glu)5-des Pro36, Pro37, Pro38 [Trp(O2)25, Asp28] Exendin-4(1-39)-NH2,

des Pro36, Pro37, Pro38 [Trp(O2)25, Asp28] Exendin-4(1-39)-(Lys)6-NH2,

H-(Lys)6-des Pro36, Pro37, Pro38 [Trp(O2)25, Asp28] Exendin-4(1-39)-(Lys)6-NH2,

H-Asn-(Glu)5-des Pro36, Pro37, Pro38 [Trp(O2)25, Asp28] Exendin-4(1-39)-(Lys)6-NH2,

H-(Lys)6-des Pro36 [Met(O)14, Asp28] Exendin-4(1-39)-Lys6-NH2,

des Met(O)14 Asp28 Pro36, Pro37, Pro38 Exendin-4(1-39)-NH2,

H-(Lys)6-des Pro36, Pro37, Pro38 [Met(O)14, Asp28] Exendin-4(1-39)-NH2,

H-Asn-(Glu)5-des Pro36, Pro37, Pro38 [Met(O)14, Asp28] Exendin-4(1-39)-NH2,

des Pro36, Pro37, Pro38 [Met(O)14, Asp28] Exendin-4(1-39)-(Lys)6-NH2,

H-(Lys)6-des Pro36, Pro37, Pro38 [Met(O)14, Asp28] Exendin-4(1-39)-(Lys)6-NH2,

H-Asn-(Glu)5 des Pro36, Pro37, Pro38 [Met(O)14, Asp28] Exendin-4(1-39)-(Lys)6-NH2,

H-Lys6-des Pro36 [Met(O)14, Trp(O2)25, Asp28] Exendin-4(1-39)-Lys6-NH₂,

H-des Asp28 Pro36, Pro37, Pro38 [Met(O)14, Trp(O2)25] Exendin-4(1-39)-NH₂,

H-(Lys)6-des Pro36, Pro37, Pro38 [Met(O)14, Asp28] Exendin-4(1-39)-NH₂,

H-Asn-(Glu)5-des Pro36, Pro37, Pro38 [Met(O)14, Trp(O2)25, Asp28] Exendin-4(1-39)-NH₂,
des Pro36, Pro37, Pro38 [Met(O)14, Trp(O2)25, Asp28] Exendin-4(1-39)-(Lys)6-NH₂,

H-(Lys)6-des Pro36, Pro37, Pro38 [Met(O)14, Trp(O2)25, Asp28] Exendin-4(S1-39)-(Lys)6-NH₂,

H-Asn-(Glu)5-des Pro36, Pro37, Pro38 [Met(O)14, Trp(O2)25, Asp28] Exendin-4(1-39)-(Lys)6-NH₂;

or a pharmaceutically acceptable salt or solvate of any one of the afore-mentioned Exendin-4 derivative.

[0022] Hormones are for example hypophysis hormones or hypothalamus hormones or regulatory active peptides and their antagonists as listed in Rote Liste, ed. 2008, Chapter 50, such as Gonadotropine (Follitropin, Lutropin, Choriogonadotropin, Menotropin), Somatotropine (Somatotropin), Desmopressin, Terlipressin, Gonadorelin, Triptorelin, Leuprorelin, Buserelin, Nafarelin, Goserelin.

[0023] A polysaccharide is for example a glucosaminoglycane, a hyaluronic acid, a heparin, a low molecular weight heparin or an ultra low molecular weight heparin or a derivative thereof, or a sulphated, e.g. a poly-sulphated form of the above-mentioned polysaccharides, and/or a pharmaceutically acceptable salt thereof. An example of a pharmaceutically acceptable salt of a poly-sulphated low molecular weight heparin is enoxaparin sodium.

[0024] Antibodies are globular plasma proteins (~150 kDa) that are also known as immunoglobulins, which share a basic structure. As they have sugar chains added to amino acid residues, they are glycoproteins. The basic functional unit of each antibody is an immunoglobulin (Ig) monomer (containing only one Ig unit); secreted antibodies can also be dimeric with two Ig units as with IgA, tetrameric with four Ig units like teleost fish IgM, or pentameric with five Ig units, like mammalian IgM.

[0025] The Ig monomer is a "Y"-shaped molecule that consists of four polypeptide chains; two identical heavy chains and two identical light chains connected by disulfide bonds between cysteine residues. Each heavy chain is about 440 amino acids long; each light chain is about 220 amino acids long. Heavy and light chains each contain intrachain disulfide bonds which stabilize their folding. Each chain is composed of structural domains called Ig domains. These domains contain about 70-110 amino acids and are classified into different categories (for example, variable or V, and constant or C) according to their size and function. They have a

characteristic immunoglobulin fold in which two β sheets create a "sandwich" shape, held together by interactions between conserved cysteines and other charged amino acids.

[0026] There are five types of mammalian Ig heavy chain denoted by α , δ , ϵ , γ , and μ . The type of heavy chain present defines the isotype of antibody; these chains are found in IgA, IgD, IgE, IgG, and IgM antibodies, respectively.

[0027] Distinct heavy chains differ in size and composition; α and γ contain approximately 450 amino acids and δ approximately 500 amino acids, while μ and ϵ have approximately 550 amino acids. Each heavy chain has two regions, the constant region (C_H) and the variable region (V_H). In one species, the constant region is essentially identical in all antibodies of the same isotype, but differs in antibodies of different isotypes. Heavy chains γ , α and δ have a constant region composed of three tandem Ig domains, and a hinge region for added flexibility; heavy chains μ and ϵ have a constant region composed of four immunoglobulin domains. The variable region of the heavy chain differs in antibodies produced by different B cells, but is the same for all antibodies produced by a single B cell or B cell clone. The variable region of each heavy chain is approximately 110 amino acids long and is composed of a single Ig domain.

[0028] In mammals, there are two types of immunoglobulin light chain denoted by λ and κ . A light chain has two successive domains: one constant domain (C_L) and one variable domain (V_L). The approximate length of a light chain is 211 to 217 amino acids. Each antibody contains two light chains that are always identical; only one type of light chain, κ or λ , is present per antibody in mammals.

[0029] Although the general structure of all antibodies is very similar, the unique property of a given antibody is determined by the variable (V) regions, as detailed above. More specifically, variable loops, three each on the light (V_L) and three on the heavy (V_H) chain, are responsible for binding to the antigen, i.e. for its antigen specificity. These loops are referred to as the Complementarity Determining Regions (CDRs). Because CDRs from both V_H and V_L domains contribute to the antigen-binding site, it is the combination of the heavy and the light chains, and not either alone, that determines the final antigen specificity.

[0030] An "antibody fragment" contains at least one antigen binding fragment as defined above and exhibits essentially the same function and specificity as the complete antibody of which the fragment is derived from. Limited proteolytic digestion with papain cleaves the Ig prototype into three fragments. Two identical amino terminal fragments, each containing one entire L chain and about half an H chain, are the antigen binding fragments (Fab). The third fragment, similar in size but containing the carboxyl terminal half of both heavy chains with their interchain disulfide bond, is the crystalizable fragment (Fc). The Fc contains carbohydrates, complement-binding, and FcR-binding sites. Limited pepsin digestion yields a single $F(ab')_2$ fragment containing both Fab pieces and the hinge region, including the H-H interchain disulfide bond. $F(ab')_2$ is divalent for antigen binding. The disulfide bond of $F(ab')_2$ may be cleaved in order to obtain Fab'. Moreover, the variable regions of the heavy and light chains can be fused together to form a single chain variable fragment (scFv).

[0031] Pharmaceutically acceptable salts are for example acid addition salts and basic salts. Acid addition salts are e.g. HCl or HBr salts. Basic salts are e.g. salts having a cation selected from alkali or alkaline, e.g. Na⁺, or K⁺, or Ca²⁺, or an ammonium ion N⁺(R1)(R2)(R3)(R4), wherein R1 to R4 independently of each other mean: hydrogen, an optionally substituted C1-C6-alkyl group, an optionally substituted C2-C6-alkenyl group, an optionally substituted C6-C10-aryl group, or an optionally substituted C6-C10-heteroaryl group. Further examples of pharmaceutically acceptable salts are described in "Remington's Pharmaceutical Sciences" 17. ed. Alfonso R. Gennaro (Ed.), Mark Publishing Company, Easton, Pa., U.S.A., 1985 and in Encyclopedia of Pharmaceutical Technology.

[0032] Pharmaceutically acceptable solvates are for example hydrates.

[0033] Further scope of applicability of the present invention will become apparent from the detailed description given hereinafter. However, it should be understood that the detailed description and specific examples, while indicating preferred embodiments of the invention, are given by way of illustration only, since various changes and modifications within the scope of the invention will become apparent to those skilled in the art from this detailed description.

Brief Description of the Drawings

[0034] The present invention will become more fully understood from the detailed description given hereinbelow and the accompanying drawings, which are given by way of illustration only, and thus, are not limitive of the present invention, and wherein:

Figure 1

shows an exemplary embodiment of a medicament delivery device before use, and

Figure 2

shows an exemplary embodiment of a medicament delivery device during use.

[0035] Corresponding parts are marked with the same reference symbols in all figures.

Detailed Description

[0036] Figures 1 and 2 show an exemplary embodiment of a medicament delivery device 1 before and during administration of a medicament to a patient, respectively. Those of skill in the art will understand that the patient may be a human or animal. In the exemplary embodiment, the delivery device 1 is an autoinjector designed to automatically deliver a dose of a medicament by means of a needle 2 upon sleeve-driven actuation. Those of skill in the art will understand that the delivery device 1 may be a pen injector, a syringe, an infusion device,

etc.

[0037] An exemplary embodiment of a sleeve-driven actuation mechanism comprises a housing 5, an inner sleeve 3 and an outer sleeve 4 telescopically arranged on the inner sleeve 3. The inner sleeve 3 and the outer sleeve 4 are axially translatable relative to each other and relative to the housing 5. The inner sleeve 3 covers the needle 2 before and after use of the delivery device 1 to prevent accidental needlestick injuries. The outer sleeve 4 serves to actuate a delivery mechanism in the delivery device 1. The sleeves 3, 4 may be arranged telescopically and substantially shaped as hollow cylinders with open proximal ends. The outer sleeve 4 has an open distal end 4.1 for accommodating the inner sleeve 3. A distal end 3.1 of the inner sleeve 3 may be opened or have a cover face with a central aperture for accommodating projection of the needle 2. In an exemplary embodiment, the distal end 3.1 of the inner sleeve 3 may be planar or curved.

[0038] Figure 1 shows an exemplary embodiment of the delivery device 1 in a first state, e.g., prior to use on an injection site 7. In the first state, the outer sleeve 4 projects distally out of the housing 5, and the inner sleeve 3 projects distally out of the outer sleeve 4. In the first state, the sleeves 3, 4 cover a distal needle tip 2.1 of the needle 2 and thus prevent accidental needlestick injuries. For example, in the first state, the distal needle tip 2.1 of the needle 2 may be proximal of the distal end 4.1 of the outer sleeve 4.

[0039] In an exemplary embodiment, in the first state, the inner sleeve 3 may be axially translatable relative to the outer sleeve 4, but the outer sleeve 4 may be locked relative to the housing 5. Thus, the inner sleeve 3 may be repeatedly retracted into the outer sleeve 4 a predetermined distance without triggering delivery of the medicament. This may prevent inadvertent triggering of the delivery device 1, allowing for realignment of the delivery device 1 on a different injection site.

[0040] In an exemplary embodiment, the inner sleeve 3 may be biased in the first state by a first spring element, and the outer sleeve 4 may be biased in the first state by a second spring element.

[0041] Figure 2 shows an exemplary embodiment of the delivery device 1 in a second state, e.g., during use. When the delivery device 1 is pressed against an injection site, the inner sleeve 3 may be pushed into an intermediate position in which it is fully contained inside the outer sleeve 4, and the distal end 4.1 of the outer sleeve 4 touches the injection site 7. When the distal end 3.1 of the inner sleeve 3 is in a same plane as the distal end 4.1 of the outer sleeve 4, the inner sleeve 3 and the outer sleeve 4 may be coupled together so that further pressing of the delivery device 1 against the injection site 7 causes the sleeves 3, 4 to move together proximally relative to the housing 5. For example, the inner sleeve 3 may engage the outer sleeve 4 when the inner sleeve 3 has attained a predetermined axial position relative to the outer sleeve 4.

[0042] In an exemplary embodiment, when the inner sleeve 3 engages the outer sleeve 4, the

needle 2 may be inserted into the injection site 7 and the medicament may be delivered. In another exemplary embodiment, when the outer sleeve 4 is pressed against the injection site 7, the needle 2 may be inserted into the injection site 7 and the medicament may be delivered.

[0043] A tactile feedback may be provided in the form of resistance. For example, the first spring element associated with the inner sleeve 3 may require less force to compress than the second spring element associated with the outer sleeve 4. Thus, an increased force is necessary to cause the outer sleeve 4 to move proximally, axially relative to the housing 5. This has the advantage that the patient can clearly distinguish the two steps of the process and thus removes a potential patient's feeling of insecurity concerning the injection process. A further advantage of the actuation mechanism according to the invention is that the different pressures for the two steps of the process can be realized more easily because they are induced automatically by coupling the sleeves to different compression springs. Of course, in alternative embodiments, the compression springs may be replaced by other tensioning members.

[0044] In an exemplary embodiment, the sleeves 3, 4 have different colors or indicia. For example, different colors emphasize the different functions of the sleeves 3, 4 and thus distinguish the two steps of the injection process even more clearly.

[0045] After the injection process, the delivery device 1 is withdrawn from the injection site 7. When force is removed from the sleeves 3,4, the compression springs relax and shift the sleeves 3, 4 distally toward the first state so that they again cover the needle 2. Thus, advantageously accidental needlestick injuries are prevented after use of the delivery device 1.

[0046] In a preferred extension of the invention, the delivery device 1 additionally comprises additionally a locking mechanism, which locks the position of the inner sleeve 3 and/or the position of the outer sleeve 4 relative to each other and/or the housing 5. The locking mechanism may ensure that the inner sleeve 3 and/or outer sleeve 4 cover the distal needle tip 2.1. This advantageously further reduces the risk of accidental needlestick injuries after using the delivery device 1.

[0047] For instance, the locking mechanism may comprise at least one latch member of the inner sleeve 3 or the outer sleeve 4 and a corresponding groove located in the housing 5 of the drug delivery device 1, the groove being adapted to receive the latch member. Alternatively, the latch member may be part of the housing 5 and the groove may be located in a sleeve 3, 4.

REFERENCES CITED IN THE DESCRIPTION

Cited references

This list of references cited by the applicant is for the reader's convenience only. It does not form part of the European patent document. Even though great care has been taken in compiling the references, errors or omissions cannot be excluded and the EPO disclaims all liability in this regard.

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Patentkrav

1. Lægemiddelfremføringsanordning (1), der har en kanyle (2) med en distal spids (2.1), hvilken anordning omfatter:

5 et hus (5),
en aktiveringsmekanisme, der omfatter en ydre bøsning (4) og en indre bøsning (3),
med den ydre bøsning (4) teleskopisk indrettet i forhold til lægemiddelfremføringsanordningen (1), idet den ydre bøsning (4)
10 er aksialt forskydelig i forhold til huset (5),
med den indre bøsning (3) teleskopisk indrettet i forhold til den ydre bøsning (4), idet den indre bøsning (3) er aksialt forskydelig i forhold til den ydre bøsning (4),
hvor den indre bøsning (3) i en første tilstand rager distalt
15 ud af den ydre bøsning (4), og den ydre bøsning (4) rager distalt ud af huset (5),
hvor den indre bøsning (3) og den ydre bøsning (4) dækker kanylens (2) distale spids (2.1) i den første tilstand,
hvor den indre bøsning (3) i en anden tilstand er indeholdt i
20 den ydre bøsning (4), og den distale spids (2.1) rager distalt ud i forhold til den ydre bøsning (4),
hvor den indre bøsning er forspændt af et første fjederelement eller et andet spændingselement i den første tilstand, og den ydre bøsning er forspændt af et andet fjederelement eller et
25 andet spændingselement i den første tilstand,
hvor den ydre bøsning (4) bevægelse proksimalt i forhold til lægemiddelfremføringsanordningen (1) i den anden tilstand igangsætter fremføringen af et lægemiddel i lægemiddelfremføringsanordningen (1), **kendetegnet ved, at**
30 det for at bevirke, at den ydre bøsning (4) bevæger sig proksimalt i forhold til huset, er nødvendigt med en øget kraft sammenlignet med at bevæge den indre bøsning (3).

2. Lægemiddelfremføringsanordning (1) ifølge krav 1, hvor den
35 indre bøsning (3) og den ydre bøsning (4) har forskellige farver eller kendetegn.

3. Lægemiddelfremføringsanordning (1) ifølge et hvilket som helst af de foregående krav, hvor den ydre bøsning (4) position er fast i forhold til lægemiddelfremføringsanordningen (1) i den første tilstand.

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4. Lægemiddelfremføringsanordning (1) ifølge et hvilket som helst af de foregående krav, hvor den indre bøsning (3) er i indgreb med den ydre bøsning (4) i den anden tilstand.

10 5. Lægemiddelfremføringsanordning (1) ifølge krav 4, hvor den indre bøsning (3) indbefatter en klinke, der er egnet til at gå i indgreb med en fordybning eller en åbning i den ydre bøsning (4).

15 6. Lægemiddelfremføringsanordning (1) ifølge krav 4, hvor den ydre bøsning (4) indbefatter en klinke, der er egnet til at gå i indgreb med en fordybning eller en åbning i den indre bøsning (3).

20 7. Lægemiddelfremføringsanordning (1) ifølge et hvilket som helst af de foregående krav, hvor den indre bøsning (3) i en tredje tilstand er låst i forhold til den ydre bøsning (4), og den ydre bøsning (4) er låst i forhold til lægemiddelfremføringsanordningen (1).

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8. Lægemiddelfremføringsanordning (1) ifølge krav 7, hvor den indre bøsning (3) og/eller den ydre bøsning (4) dækker kanylens (2) distale spids (2.1) i den tredje tilstand.

DRAWINGS

Drawing

